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# Following In Situ the Degradation of Mesoporous Silica in Bio-relevant Conditions: at Last, a Good Comprehension of the Structure Influence.

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protein

ABSTRACT: Mesoporous silica nanoparticles (MSNs) have seen a fast development as drug delivery carriers thanks to their tunable porosity and high loading capacity. The employ of MSNs in biomedical applications requires a good understanding of their degradation behavior both to control drug release and to assess possible toxicity issues on human health. In this work, we study mesoporous silica degradation in biologically relevant conditions through in situ ellipsometry on model mesoporous nanoparticular or continuous thin films, in buffer solution and in media containing proteins. In order to shed light on the structure/dissolution relationship, we performed dissolution experiments far

from soluble silicate species saturation. Via a complete decorrelation of dissolution and

diffusion contributions, we proved unambiguously that surface area of silica vectors is the main parameter influencing dissolution kinetics, while thermal treatment and open mesoporous network architecture have a minor impact. As a logical consequence of our dissolution model, we proved that dissolution lag-time can be promoted by selective blocking of the mesopores that limits the access to the mesoporous internal surface. This study was broadened by studying the impact of the organosilanes in the silica structure, of the presence of residual structuring agents, and of the chemical composition of the dissolution medium. The presence of albumin at blood concentration was found affecting drastically the dissolution kinetics of the mesoporous structure, acting as a diffusion barrier. At the global, we could identify the main factors affecting mesoporous silica materials degradation and proved that we can tune their structure and composition for adjusting dissolution kinetics, in order to achieve efficient drug delivery.

#### INTRODUCTION

In the field of nanomedicine, research on therapeutic vectors for drug delivery led to many synthesis strategies aiming at creating functional nanovectors with various properties. Three of them are mandatory for a successful application: (i) the targeting ability (usually for a tumor or a given organ), (ii) the ability to deliver drug in a controlled manner, and (iii) the ability to be cleared via nontoxic pathways (usually by biological degradation such as enzymatic lysis or by dissolution). From a pharmacokinetic point of view, an optimal effect is obtained when these three properties happen sequentially. So far, if materials scientists have spent much effort in designing efficient nanovectors architectures and delivery strategies, very little literature can be found on the study of the in vivo fate of the carrier after drug release. The risk associated with uncontrolled degradation is either a non-targeted drug delivery or the in vivo accumulation of NPs leading to a limitation of administration frequency and/or toxic side effects. Yet, degradation studies in vivo are indeed very difficult to perform and require complex analytical approaches for recovering nanovectors from living tissues and/or the ability to analyze and discriminate within an organ between nanovectors and nanovectors degradation products. From a physico-chemical point of view, one key parameter to master is the control of nanoparticles degradation kinetics in the various environments in which vectors circulate (blood, extracellular matrix, endosomes, etc.). Consequently, much effort is still needed for creating nanovectors architectures and compositions allowing a controlled degradation kinetic of the carrier concomitant with, or following the drug release.

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In this aspect, mesoporous silica nanocarriers (MSN) have attracted considerable attention as drug delivery nanocarriers.<sup>[1,2]</sup> Thanks to its excellent biocompatibility and high loading capacity, mesoporous silica is frequently employed to realize nanoparticles for drug delivery or in combination with other materials to form multifunctional core-shell structures.<sup>[3–6]</sup> Additionally, mesoporous silica materials provide easy functionalization with silane agents and adjustable pore sizes to host a very large amount of active molecules. As for other nanovectors, their use for biomedical applications is raising the need to address their dissolution kinetics in vivo.

Silica is not chemically stable in water media as it undergoes hydrolysis forming orthosilicic acid and silicate oligomers. The hydrolysis of Si-O-Si bonds is the rate-limiting step of the reaction. The driving force of dissolution is the undersaturation in silicates of the medium at the vicinity of the vector's surface. As soon as saturation is reached (usually in confined media), silica can randomly nucleate and re-precipitate, getting to an equilibrium that hinders the complete dissolution of solid silica particles.<sup>[7]</sup> Besides, with its own porous architecture, this behavior can lead to random mesopores occlusion due to precipitation, further inhibiting dissolution by reduction of the accessible reactive surface.<sup>[8]</sup> The direct consequence of this aspect is that we must distinguish dissolution rates in a medium that reaches saturation and dissolution rates far from saturation conditions. If in the first case degradation can last several days (and eventually never be completed), in an undersaturated medium mesoporous silica is degraded within a few hours.<sup>[7, 9–</sup> <sup>11]</sup> Thus, for better control of silica dissolution rate, as well as to obtain unhindered drug release, it is more appropriate to perform experiments well below saturation limits, in conditions known as "sink conditions".

Silica solubility strongly depends also on two main classes of parameters: on its composition and on the properties of surrounding media. The degradation of mesoporous silica particles and films under physiological conditions has been studied in vitro using simulated body fluids, physiological buffers and biological media<sup>[7,9–19]</sup> finding dissolution rates going from several hours to several days depending on experimental conditions and silica structure.

Silica bulk composition influences dissolution kinetics and so does the material structure, since the Si-O bond length and energy vary in different silica materials causing hydrolysis to happen at different rates.<sup>[20]</sup> Introducing heteroatoms such as Zr or Al results in an effective way to increase silica hydrolytic stability <sup>[7,10]</sup> and grafting the silica surface with polymers or ligands also alters its dissolution kinetics.<sup>[13,21]</sup> Moreover, the structural organization of silica at a small scale can influence its surface reactivity, as discovered by Zhang et al. <sup>[22]</sup> and, consequently, its dissolution rate. Different methods of synthesis can produce silicas with different bulk structures (e.g. presence or absence of strained cyclic species or predominance of highly connected species <sup>[23]</sup>), that do not react with the same kinetics. This could explain some of the conflicting data found in the literature about silica dissolution kinetics. The surrounding medium characteristics strongly influence dissolution rate as well, in particular temperature and pH are known to be important parameters to tune silica solubility, which starts to increase abruptly above pH 9<sup>[24]</sup> and depends on temperature.<sup>[25]</sup> The presence of ions and molecules alters silica dissolution kinetics, for example in the presence of Ca<sup>2+</sup> and Mg<sup>2+</sup> cations, calcium/magnesium silicates are formed at the silica surface, protecting underneath silica from further dissolution.<sup>[9]</sup> Icenhower et al. showed that the presence of NaCl in the solution can also increase dissolution rates up to a factor of 20x. <sup>[20]</sup> When silica NPs are engineered with surface ligands or polymers the interactions with molecules or ions in the surrounding medium can differ sensibly with respect to bare silica NPs, leading to important changes in dissolution rates. Overall, although understanding silica degradation is a decisive step towards its use in drug delivery, existing reports on silica stability

are inconsistent and very difficult to compare, for environmental conditions and vector concentration, size and architecture differ a lot from one study to another. As a consequence, it is very difficult to address and rationalize the structure/dissolution kinetics relationship of such type of nanocarrier.

In this work, we present a normalized approach aiming at determining the relationship between MSNs structural parameters and their dissolution kinetics at pH 7.4 and temperature 37°C. In order to predict MSNs behavior in bloodstream, undersaturated environments where tested. Undersaturation allows observing the material evolution free from the saturation effect which slows down and eventually avoids complete dissolution. We aimed to compare the influence of some main regulating factors which control mesoporous silica degradation in biorelevant conditions, varying them one at a time, to identify the ones having the biggest impact on the degradation kinetics. The recently developed analytical approach used is based on in situ ellipsometric analysis of mesoporous thin films having a thickness similar to the diameter of a MSN.<sup>[7, 10]</sup> Such a strategy has the advantage over NPs to expose a plane interface, allowing ellipsometric analysis with a good time resolution of a few seconds.<sup>[26]</sup> With this method, structural evolutions of the silica-based nanostructure (porous volume, size) can be monitored during the degradation process, leading to a better understanding of the overall behavior (on the contrary to NPs studies which dissolution kinetics are usually evaluated through silica titration, losing the most of information on the material structural changes). Between the main structural factors, we considered surface area, porosity, silica condensation degree, and porous network morphology. Here, we separated the contribution of these parameters, analyzing their influence on the overall degradation kinetics. We demonstrated that pore-blocking due to molecule adsorption can change

the degradation kinetics, introducing a lag-time period, which is a very interesting behavior for drug carriers as discussed in this manuscript.

In parallel with this study, we compared the degradation kinetics of mesoporous silica and usual hybrid organosilica (known for increasing hydrolytic stability),<sup>[7,27,28]</sup> containing covalently linked amino-propyl moieties. We confirmed the role of organosilanes in slowing down the silica degradation, shedding some light on the degradation mechanisms of hybrid organosilica materials. We also addressed the influence of the degradation media and, particularly, of protein-rich media. Remarkably, we showed that mesoporous silica dissolution kinetics is strictly proportional to the accessible surface area while mesoporous structure ordering and thermal treatment do not play a major role. We also demonstrated that the surrounding media composition (in particular, the presence of proteins) can control silica degradation speed in biological media. Moreover, this case of study is representative of the first, important stage of drug delivery: after injection in the bloodstream, NPs are in an undersaturated medium until they are internalized by cells.

#### EXPERIMENTAL SECTION

**Mesoporous silica thin films synthesis and characterization;** Mesoporous thin films are synthesized with the evaporation induced self-assembly (EISA) approach, employing hexadecyltrimethylammonium bromide (CTAB) and Pluronic F127 as templating agents <sup>[29]</sup>, respectively indicated with letters CT and F. Briefly, templating surfactants are dissolved in H<sub>2</sub>O/EtOH/HCl together with inorganic precursors such as tetraethoxysilane (TEOS) and 3-aminopropyl triethoxysilane (APTES). They are added in determined proportions to obtain solutions described in **Table 1** (silica) and **Table 2** (hybrid organosilica). Solutions are stirred for 72 h at room temperature. Resulting sols are coated on silicon substrates by dip-coating (ACEdip

from SolGelWay, Paris, France) at withdrawal speed  $u = 2 \text{ mm} \cdot \text{s}^{-1}$ , 26 °C and 30-35% relative humidity (RH), obtaining film thickness of 100-150 nm. Thin films are then stabilized for 16 h at 130 °C. Surfactants were removed by extraction in ethanol (3x5 min.). Some references of calcined silica films were treated 30 min at 200 °C and 10 min at 450 °C and are identified as CT450 when templated with CTAB and F450 when templated with Pluronic F-127.

Hybrid organosilica films (NCT) were obtained by co-condensation of TEOS and APTES and templated with CTAB; they have propyl-amine groups anchored to the silica matrix.

#### Nanoparticles (NPs) thin film synthesis and characterization

Mesoporous silica nanoparticles were synthesized adding 3 mL of a solution 0.88 M of TEOS in EtOH to a solution of CTAB (0.29 g) in water (125mL) and ammonia 0.512 M (25mL). The water/ammonia solution is kept at 50 °C under stirring (750 rpm) during the addition of TEOS solution. The nucleation happens in the first 3 minutes at 50 °C and the solution is then kept under stirring at r.t. for 2 hours. The sol containing the silica particles was dip-coated on a silicon substrate at  $u = 0.01 \text{ mm} \cdot \text{s}^{-1}$  and 50 °C to obtain a homogeneous layer of nanoparticles. The film was then heated at 130 °C for 16 h and washed in EtOH to remove CTAB (3x10min). NPs and NPs films have been imaged by scanning electron microscopy (SEM) operated on a Hitachi SU-700.

#### **GI-SAXS**

The mesostructure of thin films was analyzed by Grazing Incidence Small Angle X-Ray Scattering (GI-SAXS). Patterns were recorded on a home-made set-up; It consists of a rotating anode generator (copper anode; small focus 0.1 x 0.1 mm<sup>2</sup>, operated at 40 kV, 20 mA) equipped with a

multilayer collimating optic. The sample was placed on a rotational stage and was allowed to oscillate by a few degrees at grazing incidence during data collection. The pattern was recorded on a photo-stimulable imaging plate and a vacuum pipe was inserted between the sample and the detector to minimize air scattering. The sample-to-detector distance is 600 mm. Typical exposure time was 20'.

#### Spectroscopic Ellipsometry

Ellipsometry and environmental ellipsometric porosimetry (EEP) analyses were recorded with a UV-IR (193-1690 nm) variable angle spectroscopic ellipsometer (VASE) M2000DI from Woollam. Ellipsometry measurements were performed at an incidence angle of 75°, and the data analysis was performed with the CompleteEASE software, modeling the deposited sol–gel layers with a Cauchy dispersion model. EEP analyses were performed to evaluate the accessible porous volume and pore size distribution of the mesostructured films, employing a controlled atmosphere cell in which relative humidity (RH) was set by mass flow controllers and varied from 0 to 100%.<sup>[30,31]</sup> A Bruggemann Effective Medium Approximation (BEMA) model was used to obtain porosity values. Pore size distribution was calculated from water physisorption isotherms through a modified Kelvin equation at the relative humidity value of capillary condensation. Surface area values are obtained from EEP data using a t-plot analysis as described in reference 26.

#### X-Ray photoelectron spectroscopy (XPS)

XPS experiments have been performed on a spectrometer by Omicron Scienta. Kinetic energies of electrons are measured through an Argus hemispheric analyzer. The XPS analyses were carried out using a monochromatic K Al ( $\alpha$ ) source (1486.6 eV, 300 W). The instrument was calibrated to give a binding energy (BE) of 103.3 eV for the Si 2p corresponding to Si in SiO2 samples. An

electron gun (1 eV, 5 mA) was used on all specimens as charge neutralizer system. Peak fitting of XPS spectra was performed with CasaXPS.

#### **Dissolution in PBS**

10 mM (millimol/L) phosphate buffer was prepared from phosphate-buffered saline (PBS) tablets (Sigma–Aldrich) dissolved in MilliQ water. This buffer solution was used as a medium for dissolution experiments and the preparation of the protein solution. For silica degradation studies in PBS (pH 7.4) we employed a thermostatic liquid cell of 5 mL volume from Woollam. Experiments were performed at 37 °C in static media, collecting data every 60 seconds. We worked in non-saturated conditions, being the silica mass involved in every experiment around 0.015 mg/mL, far below the silica saturation limit at 37 °C, which is 147 µg/mL.<sup>[25]</sup> The setup employed is represented in **Figure 1**. For dissolution, thin films were modeled with a two-components BEMA employing silica or hybrid organosilica as the first component and PBS as second component (optical properties of inorganic matrix and PBS were previously measured by ellipsometry, the ones of silica were evaluated on a dense film of similar composition). Bruggemann effective medium approximation (BEMA) model allows determining the relative volumetric fractions *fa* and *fb* of two materials A and B of known dielectric constants  $\varepsilon_a$  and  $\varepsilon_b$  within a volume unit of measured dielectric constant  $\varepsilon$ :

$$f_{a}\frac{\varepsilon_{a}-\varepsilon}{\varepsilon_{a}+2\varepsilon}+f_{b}\frac{\varepsilon_{b}-\varepsilon}{\varepsilon_{b}+2\varepsilon}=0$$

More details are given in SI2. With this method, we could calculate the volumetric fraction of silica  $f_s$  present in the film at every time point. Silica mass M was calculated from the silica

volumetric fraction and the thickness of the film, assuming the walls as chemically homogeneous with a density of amorphous silica fixed at 2.2 g  $\cdot$  cm<sup>-2</sup>. For hybrid organosilica film, we assumed a similar density. The dissolved silica mass M<sub>t</sub> is then equal to the difference between initial mass M<sub>0</sub> and the silica fraction present in the film M(t).

#### **Dissolution in BSA solution**

To reproduce an environment mimicking more closely biologic media, we performed dissolution experiments of mesoporous silica layers in a PBS solution of Bovine Serum Albumine (BSA at 37 g/L, pH = 7.4). The chosen concentration is in the range of albumin concentration in human blood. In these conditions, BSA will be in its N form and exhibits an average negative surface charge.<sup>[57]</sup> Experiments were performed at 37 °C, employing the setup described in **Figure 1** and the same protocol used for dissolution studies in PBS. The BSA solution remains transparent in the visible-NIR range and it is thus possible to perform ellipsometry analysis "through" it.

#### **Surface Plasmon Resonance (SPR)**

SPR experiments have been performed on a SPR-Navi 210A from Bionavis, working at a fixed wavelength of 785 nm, in scanning angle mode. Analysis took place in two parallel channels of 1  $\mu$ L of volume, at 37 °C. Mesoporous silica was deposed by spin coating (4000 rpm, 30 sec) on BK7 Glass slides (20x12x0.55 mm) coated with gold (SPR102-AU from Bionavis), then cured at 130 °C overnight, obtaining a silica layer of 70 nm. CTAB was removed by washing in EtOH.

#### RESULTS AND DISCUSSION

#### **Materials characterizations**

The mesostructuration of thin films obtained through EISA depends on several chemical and processing parameters and was characterized by grazing-incidence small-angle X-ray scattering.<sup>[32]</sup> Mesoporous silica films templated with CTAB have a Pm3n cubic structure, while films templated with Pluronic F-127 form a p6m 2D-hexagonal structure (**Figure 2a and 2c**). Hybrid organosilica films self-organize in wormlike structures (**Figure 2b**).

Porous volume  $V_p$ , specific surface area *S* and pore size of mesoporous silica films were obtained from EEP data and are reported in **Table 3** (isotherms reported in SI, **Figure S2**). Pore size was around 3 nm in diameter and it was calculated assuming elliptical pores for Pm3n structures, semimajor axis ( $r_l$ ) and semi-minor axis ( $r_s$ ) of the ellipse are reported. For p6m and wormlike structures a pore size distribution has been obtained assuming cylindrical pores with an elliptical base. From GI-SAXS experiments the cell parameters of materials were obtained and, comparing them with pore sizes acquired through EEP, we calculated the silica wall thickness  $w_t$ .

#### Silica dissolution in PBS, theoretical approach

Monitoring the degradation of mesoporous silica and hybrid organosilica thin films through *in situ* ellipsometry we could assess the influence of some structural and exterior parameters on dissolution kinetics. We performed experiments in static undersaturated conditions, at constant pH (7.4) and temperature (37 °C).

For describing the dissolution of a solid in a liquid, Noyes and Whitney elaborated an equation <sup>[33]</sup>, in which dissolution is driven by the concentration gradient between the surface of the solid material (where  $C_s$  is assumed to equal solubility at saturation of the solid) and concentration C in the bulk solution:

With  $k = \frac{D}{Vh}$ 

$$\frac{\partial C}{\partial t} = k(C_s - C) \tag{1}$$

Here, k is a constant factor gathering the diffusion coefficient of the dissolved species D, the volume of the solution V and the width of the diffusion layer h. The concentration gradient is controlled by the diffusion process between the surface and the bulk solution. This model assumes that saturation is rapidly achieved at the solid-liquid interface (fast dissolution of the solid) and then diffusion takes place across a layer of stagnant solution, called diffusion layer, towards the bulk solution <sup>[34]</sup>. In the case of silica, which cannot be considered as a solid with fast dissolution kinetic,  $C_s$  will be assumed as the concentration of dissolved silica species at the external surface of the film which area is constant at 5 cm<sup>2</sup> (imposed by the liquid cell used for ellipsometry analysis). After integration, **equation 2** gives the value of concentration of the solution at any given time  $t \ge 0$ :

$$C(t) = C_s(1 - e^{-kt})$$
(2)

This model applies well in many cases of study where a dense solid dissolves in a liquid, and it has been adapted to several systems. <sup>[35-37]</sup> In that case, the concentration raise in solution is the factor limiting the dissolution kinetic.

Nevertheless, when the solid is mesoporous, other factors may influence its dissolution behavior. In fact, the dissolution process would take place in the mesopores, following first order kinetics, and then the dissolved species would have to diffuse until the film interface with the bulk solution. From this point we can assume that they will follow Noyes-Whitney kinetics, but the overall dissolution kinetics has also to take into account the processes happening inside the porous network. This implies that  $C_s$  will not be rapidly equal to the solubility value, as assumed in the Noyes-Whitney model, but it will probably gradually increase towards this value, depending on both the dissolved amount and diffusion process inside the mesoporous structure. The dissolution process of a porous thin film can thus be divided into two phases: (I) Dissolution and diffusion of the dissolved species from inside the porous network towards the film interface with the bulk solution, and (II) the diffusion of dissolved species from the interface to the bulk solution, following the Noyes-Whitney kinetics, driven by the concentration gradient and limited by saturation (**Figure 3**).

We performed dissolution experiments in a static environment, and in this case of figure the diffusion layer thickness *h* is not constant, but it grows with  $(D \cdot t)^{1/2}$ . The concentration profile due to diffusion of soluble species can be described with an *erf c* (*t*, *x*) function where *t* is the diffusion time and *x* is the distance from the dissolution interface:

$$\frac{C(t,x)}{C_s} = \operatorname{erf} c \left( \frac{x}{2\sqrt{Dt}} \right)$$
(3)

By reversing **equation 3** we obtain the value of the diffusion layer thickness *h*, assuming it is equivalent to a certain value of x(t), for our calculations we chose arbitrarily the value of x(t)corresponding to a  $C(x) = 0.1 \cdot C_s$ , that is, we assumed that the diffusion layer thickness is the distance at which the solution concentration is 10% of C<sub>s</sub>. To do this inversion easily we can approximate the erfc function as follows (comparison is shown in SI7):

$$\operatorname{erf} c(x,t) = \exp\left[-1.9\left(\frac{x}{2\sqrt{Dt}}\right)^{1.3}\right]$$
(4)

obtaining equation (5) for x(t) when C(x) has reached 10% of the value of  $C_s$ .

$$x(t) = h(t) = 2.3186 \sqrt{Dt}$$
(5)

Inserting the expression (5) in equation (2) we can calculate the concentration  $C_s$  of dissolved silica at the external surface of the film as a function of time:

$$Cs(t) = \frac{C(t)}{\left[1 - exp\left(-\frac{Dt}{Vh}\right)\right]} = \frac{C(t)}{\left[1 - exp\left(-\frac{\sqrt{Dt}}{2.3186V}\right)\right]}$$
(6)

The values of  $C_s(t)$  and C(t) are reported in **Figure 4** normalized for 1 cm<sup>2</sup> of silica films templated with CTAB and with F-127, using values of 1 mL for the dissolution volume V and of  $1.3 \cdot 10^{-9}$  m<sup>2</sup> s<sup>-1</sup> for the diffusion coefficient of silicic acid in water at 37 °C.

As it can be seen, upon dissolution,  $C_s$  increases gradually and never reaches the saturation value of 147 µg/mL. We can observe that the maximum value of  $C_s$  is reached at different times for the two silica films, which have the same porous volume (62%) but very different pores size and surface area. We can infer that the architecture of the mesoporous material affects the concentration at the interface  $C_s$ , more precisely, that the maximum value of the concentration at the solid-liquid interface  $C_s$  is reached faster when the specific surface area of the film is higher. If we report  $C_s$  at two different times versus the exact mesoporous surface per film area unit (taking into account film thicknesses), we find a very good linear correlation reported in **Figure 5**. At that stage, the fact that CTAB template films and F127 templated films have different pore connectivity does not seem to affect the correlation.

If we look at the thickness profile of the mesoporous films during dissolution, we observe that a huge swelling of the films takes place when  $C_s$  gets close to its maximum value; after this time

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point,  $C_s$  decreases and dissolution is slowed down (**Figure 6**), showing that either the dissolved species production decreases and/or its diffusion through the layer is slowed down.

We assume that such an effect is likely to be due to a solid to gel transformation of the layers which is a documented behavior for siliceous materials.<sup>[38–42]</sup> If we extrapolate these results, the fact that saturation concentration value is or isn't reached by C<sub>s</sub> depends very probably from the amount of silica in the film, suggesting that thick and thin films could behave differently and show different dissolution kinetics. Once the gel is formed, the diffusion of silicates species changes, affecting the dissolution kinetics of the material. For this reason, to compare the dissolution of different mesoporous silica thin films we focused the following part of our work on the first part of the dissolution process, before the material transformation in a swollen gel. This describes the silica dissolution of about 80 wt% of the material deposited. Such swelling is an unexpected behavior that, as far as we know, was never reported for the dissolution of mesoporous silica nanoparticles in simulated biological media. If one assumes that a similar swelling behavior takes place for nanoparticles, one has to keep in mind that films swell perpendicularly to the substrate only while nanoparticles would swell in 3D. As an example, 100% volume film swelling would mean a 26% increase of NPs radius. Yet this effect is only observed after more than 80% of silica is dissolved, which means that the objects in such a state exhibits a poor electronic density contrast, making them hard to spot for usual nanoparticle analysis technics in liquid state. This could explain why this effect has never been reported. Another possibility is that the rigid substrate of the film maintains the swollen gel in place, while with nanoparticles it could shatter in smaller pieces.

To summarize these results, we learnt that we can indeed separate the degradation process in two phases: (I) dissolution of silica and diffusion of the dissolved species through the porous network and (II) diffusion from the external surface of the film towards the bulk solution. In the first part,

the structure of the porous material affects dissolution kinetics, as already reported by Higuchi<sup>[43]</sup>, in that case, the diffusion can be considered limited by the accessible volume and network tortuosity. Higuchi described the diffusion in a porous solid by defining a diffusion coefficient  $D_{eff}$ =  $D \cdot V_p / \tau$  where D is the diffusion coefficient when diffusion is not limited,  $V_p$  is the porous volume and  $\tau$  is the tortuosity. A full description of this part is limited by the lack of theory and normalized method for determining  $\tau$  (which can be affected by mesopore size and connectivity). As we have seen above, in this phase, degradation kinetics is linearly correlated with the surface area. In the second phase, outside the porous film, dissolved species follow Fick's diffusion laws towards the bulk, driven by the concentration gradient, sticking to the Noyes-Whitney model.

#### Silica dissolution in PBS, normalized approach

In order to compare more easily different samples degradation kinetics, the following part of this work will study dissolution kinetics by using the amount of dissolved silica  $M_b$  normalized on starting silica mass  $M_0$  as a function of dissolution time.

#### Continuous films versus nanoparticles films

As a first experiment, we verified the relevance of film models to explain nanoparticle degradation behavior. To do so, we compared the dissolution of a plane interface film of mesoporous silica and that of a film made of mesoporous silica NPs of similar mesostructure. For this experiment, we used mesoporous silica nanoparticles having diameters around 40 nm and 2D-hexagonal mesostructure (p6m) with pores of 2 nm. These have been prepared via usual silica-based therapeutic vectors synthesis methods. **Figure 7** shows images of NPs (**Figure 7a**) and of the multilayer film obtained dip-coating the NPs suspension on a silicon wafer (**Figure 7c-d**). The film refractive index at 632 nm was 1.102 meaning that the layer had a very high porosity, as

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expected for packed mesoporous nanoparticles. EEP data, reported in SI **Figure S9**, shows two adsorption steps which correspond to the capillary condensation into the mesopores of the NPs (lower P/P<sub>0</sub>) and to the filling of inter-particles pores (higher P/P<sub>0</sub>). Due to the interparticle porosity, huge surface roughness ( $\approx$  45 nm from ellipsometric measurements) and a non-flat surface morphology, as can be seen from SEM image (**Figure 7c**), we could not calculate the total specific surface area with the t-plot method using the film external surface area as a reference (experimental and theoretical details are provided in reference 30).

These nanoparticles have usually surface area ranging from 900 to 1100 m<sup>2</sup>/g but when packed in a multilayer film they most probably lose some external surface area due to their compact spatial arrangement. Comparing the dissolution of a plane film and of a NPs film having similar mesopore size distribution (Figure 7b) we observed similar degradation kinetics. The silica release for NPs films is slower than for plain films. We estimated surface area of the film made of NPs at 690  $m^2/$  $cm^3$ , assuming that NPs compacity in the film is 60%, and their surface area is 1000  $m^2 g^{-1}$ . A comparison of dissolved amounts after 1000 seconds, normalized with the surface area of the samples, shows that both NPs based film and continuous film have dissolution rates per square meter differing by only 18%. Even though we cannot rule out a different surface area evolution between a layer of nanoparticles and a plain layer (the former one having an higher external surface area than the latter), the very similar dissolution rate observed is encouraging. Yet, a comparison with experiments performed with suspended nanoparticles would be useful. One of the only articles working in similar concentration conditions (very far from saturation, same temperature and dissolution medium) is reported in ref. 9. In this work, they observed that the complete dissolution of the mesoporous structure takes about two hours (very close to our results). Both results obtained for deposited and suspended nanoparticles strongly support the fact that kinetical

dissolution data obtained with mesoporous films are relevant for studying the dissolution rate of mesoporous nanoparticles.

#### Effect of the thermal stabilization and the mesoordering of silica films

By comparing calcined and washed CTAB template mesoporous films, we observed that hightemperature thermal treatment doesn't improve silica stability towards hydrolysis to a great extent. In fact, dissolution curves of samples heated at 130°C and 450°C presenting similar surface area, mesostructure and composition don't show much difference (**Figure 8a**). The dissolution of CT450 is only slightly slower than the one of CT130. This result is consistent with the fact that very little difference exists in condensation degree of silica networks condensed with HCl catalyst and treated at 150°C and 450°C (about 89% of Si-O-Si bond are formed in both cases), as already reported <sup>[44]</sup> in previous studies.

To complete our structural investigation, we wanted to verify if the mesoporous structure can play a role in dissolution kinetics. To do so, we prepared CT450 samples with different relative humidity (RH) conditions during film deposition, to obtain a different arrangement of the mesopores, keeping constant all the other parameters.<sup>[29]</sup> We compared dissolution kinetics of samples having similar porosity (56-62 %) and surface area values (800-850 m<sup>2</sup>/cm<sup>3</sup>). We could observe that silica presenting a Pm3n cubic structure dissolves at the very same rate than silica having a wormlike structure for more than 70% of the dissolution process, that is, before the solid to gel transformation (**Figure 8b**), time after which we can't argue any further about its structure. Thus, we can infer that the 3D organization of a mesoporous network doesn't significantly affect silica dissolution kinetics when other parameters are kept constant (mesopore size, porous volume and high pores connectivity). Anyway, it has to be mentioned that this is true for these kinds of

structures having large porous volume and many interconnections between mesopores, for which we can assume diffusion isn't particularly hindered. Analyzing porous networks less opened, the mesostructure could have a more important influence on diffusion and on the overall dissolution behavior.

#### Investigation of pore-blocking effects

A classical problem met by experimentalists with mesoporous materials is that they exhibit high surface area and get quickly contaminated in air by volatile organic compounds (VOC), or in solution by small soluble molecules that adsorb into the mesopores (see SI, **Figure S4**). In such cases, when analyzed through EEP, their porous volume is reduced by a considerable amount, passing from 61-62% after the synthesis to 43% after 5 days of rest in a standard plastic box onto a laboratory bench (we assumed the refractive index of adsorbed pollutant to be 1.45 at

700 nm). If the pores are blocked, not all the surface is available to react with water. Moreover,

diffusion of the dissolved species may be hindered, slowing down the overall dissolution process. Indeed, performing dissolution experiments on these films, we noticed a slower dissolution rate and also a different shape of the dissolution curve which becomes more sigmoidal (**Figure 9**). The reduced surface area and the hindered diffusion due to pore-blocking are likely to make dissolution kinetics slower at the beginning, but, during the degradation process, adsorbates slowly diffuse out of the pores, leaving more surface available to react with water and a more open diffusive path. As a consequence, a pore-blocking effect must be investigated. To verify this hypothesis, we artificially blocked some pores of freshly made mesoporous samples by adsorption of preformed gold clusters. Mesoporous films were soaked 1 minute in an ethanol-based solution containing 4aminothiophenol stabilized gold NPs with size distribution ranging from 0.5 to 3.5 nm and we

obtained sigmoidal dissolution curves (data are reported in SI5), confirming that blocking some mesopores induces a decrease in dissolution rate due to a reduction of accessible reactive surface and/or a hindered diffusion of silicate species. So, basically, we can consider adsorbed molecules or particles as means to temporarily modify the accessible surface area of mesoporous materials, tuning their dissolution kinetics. This is a very interesting behavior for drug delivery applications because it means that (i) the dissolution kinetic of silica-based vectors is highly correlated with its drug loading, and (ii) that mesoporous particles hosting hydrophobic drugs will dissolve with a lag-time due to their cargo acting as a pore-blocking agent for water. Yet, if water can diffuse through the drug, intermediate effects can be expected. These results are in good accordance with previous works of drug release found for drug-templated silica materials.<sup>[45-48]</sup>

Globally, if one analyses the kinetic sequence of a high surface area mesoporous silica carrier loaded with a poorly water soluble drug, we see that it will consist in (i) a slow dissolution of the silica carrier before drug release (just after injection for example), (ii) a progressive drug release (ideally at the target position) and (iii) concomitantly with drug release, an acceleration of the silica carrier dissolution. This is close to the ideal kinetic sequence envisioned at the beginning of this manuscript.

#### Dissolution of amino-functionalized mesoporous silica.

In practical applications, silica is often employed as hybrid silica, containing organic functions such as methyl or amine anchored on the surface or in the silica matrix. Silica carrying amine moieties is particularly interesting for drug delivery applications because of its easy functionalization chemistry which allows binding proteins, dyes or antibodies to the surface.<sup>[49,50]</sup>

Anyway, the presence of organic groups is known to decrease sensibly silica hydrolysis due to the inductive effect of alkyl chains which gives electron density to the silicon centers.<sup>[7,49]</sup> Hydrophobic organic chains have an additional effect on silica dissolution by modifying the water hydrogen-bonding network close to the hybrid material surface, decreasing the affinity of water molecules for the surface.

We thus investigated the dissolution kinetics in physiological conditions of hybrid aminofunctionalized silica, mesostructured from CTAB template. Comparing silica and hybrid organosilica with similar porosity (0.42 vol. fraction) and surface area (600-650 m<sup>2</sup>/cm<sup>3</sup>) we noticed a very similar dissolution rate in the first 25 minutes, then hybrid organosilica dissolution rate slowed down sensibly and after 60 minutes of dissolution (50% of total mass released) dissolution rate changed to a much slower one (Figure 10a) for the following fifteen hours. This behavior suggests a dissolution mechanism in two steps, where in a first time the hydrolysis concerns those Si-O bonds which are far enough from the alkylammonium moieties not to be influenced by the inductive effect of the organic group, and behave as if they belong to a pure silica structure. This is supported by the fact that the starting dissolution rate is coherent with the one of pure silica layers having the same characteristics. In a second step, the hybrid network dissolves and the dissolution rate changes into a much slower one, in agreement with the kinetics previously reported for organosilica materials.<sup>[27,28,51,52]</sup> Besides, nucleophilic amino groups are excellent catalysts for silica hydrolysis, but this is not true for protonated aminosilanes. Thus, the starting rate of dissolution observed for hybrid organosilica could be due to a catalytic effect of nucleophilic amines, compensating the more hydrophobic nature of the material compared to silica. On the second part of the dissolution, inhibition due to both wetting and inductive effects could dominate, giving a slower dissolution rate. As a consequence, it is indeed very interesting to

determine the number of nucleophilic amine moieties in the samples. For this reason, XPS analyses were performed on samples before and after 60 minutes of dissolution. The results confirmed our hypothesis, finding an increase in N/Si ratio which passes from 0.13 before dissolution to 0.38 after 60 minutes of dissolution (data calculated from survey spectra (reported in SI, **Figure S10**). Moreover, observing the nitrogen peak, we noticed a decrease in the peak at 402.5 eV, which is usually attributed to nucleophilic amines interacting through hydrogen bonding with silanols on the silica surface (**Figure 10b-c**). The peak decrease observed after dissolution can be due to reduced availability of silanols at the mesopore surface. As a conclusion, the slowing down of the dissolution rate after 60 minutes is likely to be due to a concomitant effect of amino-silane enrichment and progressive protonation of surface amines (inducing a progressive disappearance of nucleophilic centers able to catalyze silica hydrolysis). The contributions of the inductive effect, hydrophobic effect and catalytic effect cannot be easily decorrelated at this point.

These results demonstrate that playing on the composition of hybrid organosilica makes possible to tune the dissolution kinetics easily. The influence of the structuring agent onto the homogeneity of the dispersion of organic functions, thus on its dissolution behavior, seems to be a crucial point since an excess in amine functions may destabilize the hybrid silica network, making in some cases the overall material less stable, as already observed.<sup>[7]</sup> More generally, the reported behavior is interesting for drug delivery applications because the amine group can interact with drugs loaded in the mesopores, retaining them and releasing them more slowly, while molecules bonded on the silica part could be released faster. Indeed, the behavior of drug-loaded hybrid nanoparticles will strongly depend on the matrix/drug interaction and on the solubility of the drug, but the two-step dissolution behavior observed for hybrid organosilica could be used to control drug release from mesoporous nanoparticles.

#### Degradation in protein solutions.

Silica dissolution experiments in PBS allow relating some silica structural properties with its degradation kinetics. These data provide useful information that can be used for optimizing the material synthesis in order to modify its degradation rate in biological media. However, silica dissolution in biological environments is strongly influenced also by the dissolving media which usually contains electrolytes and biomolecules. For example, it is nowadays established that a layer of proteins (corona) forms onto the surface of nanoparticles when they are in contact with biological fluids.<sup>[53–56]</sup> The protein corona mediates the interactions of nanomaterials with cells and it determines nanoparticle biocompatibility and biodistribution.

To mimic biological environments more closely we performed degradation experiments in a solution containing proteins with concentration comparable to the one present in the blood. We used 37.5 mg/mL of BSA in PBS. In these conditions, BSA will be in its N form and will have a global negative surface charge.<sup>[57]</sup> Silica has also a negative surface charge at neutral pH, but protein adsorption takes place anyway<sup>[58]</sup>, mainly for entropic reasons, via hydrogen bonding interactions between the silica surface silanols and the amino-acid chains of BSA. At such concentration, we assume a uniform adsorbed layer on the silica surface. In our degradation experiments (**Figure 11**), we observed that at an early time (less than 50 minutes) the presence of proteins slowed down silica dissolution by a 5x factor. Moreover, a close look at **Figure 11a** showed that the initial dissolution rate decreases progressively in the first 60 minutes for reaching another slower regime. This observation is fully consistent with the time needed for the complete formation of a tighter BSA hard corona onto silica nanoparticles.<sup>[59,60]</sup>

Removing samples after 3 hours of soaking at 37 °C and measuring them by ellipsometry after a quick rinsing in water, we found almost no silica residual layer (1-5 nm) for samples soaked in PBS, while samples soaked in BSA solution still presented a thick layer of silica (about 70% of the starting thickness), as shown in **Figure 11b**.

Silica dissolution happens at a slower rate in the presence of proteins, likely due to a barrier effect to diffusion caused by protein adsorption onto the surface, as already reported.<sup>[61]</sup> This barrier effect is likely to be due to the capping of mesopores. This behavior is consistent with the retarded drug release observed by Shahabi<sup>[53]</sup> for porous silica nanoparticles in the presence of proteins. Hybrid organosilica presents also a similar behavior, with a consistent slowdown in dissolution rate in protein enriched-media, as shown in **Figure 11c**, and a dissolution rate which decreases in the first 20 minutes until it reaches a constant rate. Qualitatively, the same effect of protein surface layer on the dissolution rate has been observed for every sample analyzed, independently from its porosity, structure, composition and thermal treatment. This strengthens the hypothesis that the main effect of proteins is the hindrance to the diffusion process of dissolved species from the porous network to the bulk solution. The dissolution rate of the hybrid film is similar to that of pure silica film after 60 minutes, as evident from **Figure 12**. This suggests that the hard corona of BSA is forming faster onto hybrid films than onto silica film probably due to the presence of the additional ammonium groups of the organosilica.

At that point, it is important to highlight that if protein corona controls particles' bioactivity, on the other hand, particles may have reverse effects on biomolecules which can be harmful. In fact, binding to nanoparticles can affect the structure and function of proteins (they can unfold and be denatured by the contact with the particle's surface, losing their function such as enzymatic activity), with catastrophic effects on cellular metabolism, and they can also undergo fibrillation

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due to the contact with nanomaterials.<sup>[62,63]</sup> The modification of protein structure is one of the main mechanisms of toxicity associated with nanoparticles in vivo. Residual molecules from particle's synthesis can also react with proteins and denature them, causing damages to cellular metabolism. While we were performing silica dissolution experiments in protein solutions, we could observe sometimes an effect of residual CTAB surfactant on surface adsorbed BSA (cf. **Figure 13**).

In fact, a large majority of mesostructured silica-based therapeutic vectors are prepared by using CTAB surfactant. This CTAB is then removed by solvent extraction. In our samples whose surfactant was removed by thermal treatment at 450°C, the refractive index of the mesoporous layer decreases as described here above during dissolution (**Figure 13b**). In solvent-extracted samples, the refractive index decrease is interrupted after 40 minutes of dissolution and then, its value starts increasing sensibly, until it reaches the starting value in about 40 min. After 80 minutes, it decreases again constantly (**Figure 13a**).

In order to determine if this behavior comes from the presence of residual CTAB, we reabsorbed CTAB in the pores of a calcined clean sample (CT450) by soaking it in an ethanolic solution of CTAB and observed that its dissolution presents a similar transient "sinusoidal" evolution of refractive index value. This confirms the role of CTAB in the observed phenomenon. From literature, we know that CTAB forms a protein-surfactant complex with BSA, not soluble in water, unfolding the protein and inducing its aggregation with other BSA molecules.<sup>[64,65]</sup> The positively charged head group of CTAB interacts electrostatically with carboxylic groups of aspartic acid and glutamic acid on BSA, which are deprotonated at pH = 7.4. This process reduces the  $\zeta$ -potential of BSA, triggering aggregation after an initial lag phase which was found to last around 20 minutes.<sup>[64]</sup> The presence of this CTAB-BSA complex and/or the BSA aggregates on the surface of mesoporous silica films could explain the optical response of the washed mesoporous silica

layers. Following the dissolution of the same films by surface plasmon resonance (SPR) we could observe a sudden decrease in the material refractive index when injecting the protein solution with the dissolution medium. This behavior was not observed in PBS, sustaining the hypothesis of a CTAB-BSA complex formation. In SPR experiments we could flush both PBS and BSA solution on the same sample in two different channels and compare the two dynamics being sure that there was no difference due to material synthesis. In **Figure 14a** (washed film with remaining CTAB), BSA injection promotes a decrease of the SPR angle (related to the refractive index of the film) which goes below the value of the film maintained in PBS. This behavior can be explained with a decrease in the refractive index due to the diffusion of CTAB molecules outside the silica matrix. Later, the SPR angle increases, responding to a higher refractive index due to protein adsorption onto the film's surface. This is the expected behavior, which has been observed on dense silica (not shown) as well as on films where CTAB was efficiently removed (**Figure 14b**).

By analyzing washed samples of silica films prepared with CTAB, we found that the residual amount of CTAB is very small because it was not possible to detect it by ellipsometry in air or by XPS, but it is enough to form some CTAB-BSA complex and adsorb it on the surface of the film. The amount of these aggregates is probably very low and doesn't seem to influence the overall dissolution kinetics (considering the residual silica after 3h, measured ex situ) but it is enough to be detected by in situ ellipsometry because it modifies sensibly the optical response of the sample. More generally, the observation of this phenomenon points out how important is the washing step in the synthesis protocol of nanoparticles made for drug delivery. This effect may explain some discrepancies in literature onto the colloidal stability of some silica-based therapeutic vectors dispersed in protein-rich media. As importantly, if the conformation of adsorbed protein may be not a major issue for dissolution experiments, it is indeed very important in a biological

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environment. The formation of protein-ligand complexes provokes the unfolding of the native structure of proteins and can be a starting point for aggregation, which compromises their biological function. It has been demonstrated in the case of CTAB, in which the reduced ζ-potential of the complex CTAB-BSA promotes aggregation.<sup>[64]</sup> Complexes with BSA have also been observed for other surfactants such as anionic sodium dodecyl sulfate (SDS) and nonionic polyoxyethylene-8-lauryl ether.<sup>[65]</sup> As therapeutic vectors usually cannot be calcined (nanoparticles can aggregate during thermal treatments and they contain usually some organic ligand), solvent extraction is usually the chosen method to eliminate surfactants from nanocarriers. These results highlight how important it is to use a very optimized washing protocol for preparing nanoparticles allowing to assure the complete removal of the templating surfactant.

Finally, we didn't observe the formation of CTAB-BSA complexes onto the surface of hybrid organosilica films, although they have been washed in ethanol to remove the surfactant with the same protocol used for pure silica films. We can infer that the washing was more efficient in the case of hybrid organosilica, probably because of the electrostatic repulsions between the cationic head of CTAB and the protonated amines on the silica surface. Nevertheless, hybrid silica materials carrying amine moieties have a modified surface charge compared to silica, depending on their aminosilanes content. These materials can reach neutral to positive surface charges, depending on their synthesis conditions, strongly influencing adsorption and denaturation of biomolecules. We were not able to detect visible aggregation and denaturation due to surface charge on the hybrid organosilica films nor the effect of CTAB residual presence on BSA conformation. Anyway, the dissolution of silica and hybrid organosilica films in protein media could present some differences due to the altered surface charge of the two materials. For

determining the exact effect of a moderate surface amine content on protein-surface interactions, a dedicated study would be necessary, which is not the objective of this work.

#### CONCLUSIONS

This study illustrates the set of chemical and physico-chemical parameters that can be used to tune the degradation of therapeutic silica-based carriers in simulated biological media. In summary, the influence of several synthesis and structural parameters on mesoporous silica degradation was investigated by a standard and well-controlled *in situ* ellipsometry analysis platform. We demonstrated that the main factor controlling silica dissolution rate is its accessible surface area (with a linear relationship) and that its mesostructure arrangement and thermal treatment have a minor impact on the overall kinetics. Anyway, a structure with high tortuosity can dissolve slower, as already evidenced by Braun et al. <sup>[16]</sup>, because the dissolution relies on molecular diffusion. The method employed in this work allowed us to split the process of silica dissolution into two parts: inside and outside the porous network. We found that in the first phase, the concentration of dissolved silica reaching the external surface of the porous material is depending linearly on the surface area. This result explains the key role of the surface area on the mesoporous silica degradation in aqueous media, already identified by Kuroda<sup>[46]</sup> and Shi.<sup>[9]</sup> Kuroda and co-workers demonstrated that silica particles with similar surface area have nearly identical degradability rates regardless of their diameter. On the other hand, Shi and co-workers <sup>[9]</sup> have studied the hydrolytic degradation of mesoporous silica with increasing surface area and found faster kinetics for higher surface areas. Moreover, we studied silica films having the same porous volume but different surface areas, separating the contributions of these two physico-chemical parameters, which are

often correlated, and we confirmed that the key feature controlling silica degradation in aqueous media is its surface area. We also observed that pore blocking can introduce a lag-time in the dissolution kinetics, useful to tune drug carrier degradation and release. This effect is most probably due to a decrease in the available surface area caused by pore blocking and/or an increase of tortuosity in the porous structure. The obtained results on silica degradation rate were consistent with the ones reported by He et al. <sup>[9]</sup> for undersaturated media, and comparable with data from Braun et al. <sup>[16]</sup>, considering differences in saturation conditions and surface evolution. Mesoporous silica films of 100-120 nm of thickness dissolve completely in PBS at 37 °C within a few hours. We observed that hybrid organosilica mesostructured from CTAB and carrying propylamine moieties degrade in two steps and with an overall slower rate than pure silica. We verified that the observed kinetics are respected even in the case of films made of nanoparticles, meaning that degradation data obtained from thin films can be applied to predict NPs behavior.

The degradation of mesoporous silica and hybrid organosilica in media containing high concentrations of proteins is slowed down. It was demonstrated that surface adsorbed molecules decrease the dissolution rate forming a barrier to diffusion.

Additionally, we demonstrated that residual surfactant molecules in the mesopores can influence the interactions of the material with biomolecules and reshape the interface, even if their amount is very low. They can unfold proteins forming stable complexes and induce protein aggregation, which can be harmful to cellular processes. This underlines the importance of efficient protocols to remove surfactants and other synthesis reagents from NPs employed for drug delivery purposes.

Considering the different physicochemical properties and degradation kinetics of each MSN system, researchers can select the best nanoplatform to fulfill a specific application. If long-term

circulation times are needed, hybrid silica-containing organic functions or Zr-doped silica would be probably the best candidates, because of their intrinsic higher hydrolytic stability. Hybrid organosilica can also be interesting to release drugs with two different kinetics, due to its domain structure: a burst release in the first hours can be followed by a much slower release over several days. Whether the nanoparticles are used as imaging or delivery platforms for long-term applications, their surface area should be kept as lowest as possible without compromising their function. On the contrary, if a fast degradation is desired, the surface area should be high. For delivery carriers, it must be remembered than the loading of cargoes also tunes the nanoparticles degradation kinetics, thus a cargo which has strong interaction with the host matrix can be capable to introduce a non-negligible lag-time in the degradation and in the drug release, providing good timing for targeted delivery without the need of gate blockers. One main issue when designing MSN as drug delivery platforms is to avoid their aggregation, remembering that surface functionalization stabilizes them also towards degradation, prolonging their lifetimes.

As a future perspective, we feel that it would be interesting to study mesoporous silica degradation behavior in dynamic conditions, in contact with flowing biological fluids, mimicking more closely the in vivo environment and addressing flow effects on silica dissolution.

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# **Author Contributions**

The manuscript was written through contributions of all authors. All authors have given

approval to the final version of the manuscript.

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## **Supporting Information**

Solution preparation procedures; GI-SAXS patterns of mesoporous films; Environmental Ellipsometric Porosimetry (EEP) curves; Derivation of silica dissolution from ellipsometry data; IR spectrum of VOC-contaminated mesoporous silica film; Gold clusters synthesis and adsorption in mesopores of hybrid silica films: protocols, TEM, UV-vis spectrum, Fluorescence spectrum; Approximation of the erfc (x) function; XPS analysis of amino-functionalized mesoporous silica films before and after dissolution.

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**Table 1:** Molar composition of solution used to prepare mesoporous silica thin films templatedwith CTAB surfactant (CT samples) and F-127 surfactant (F samples).

Sample	TEOS	EtOH	HCl	H <sub>2</sub> O	Pluronic F-127	СТАВ
СТ	1	40	0.09	5	-	0.14
F	1	40	0.09	5	0.005	-

Table 2: Molar composition of solution used to prepare hybrid organosilica mesoporous thin films

Sample	APTES	TEOS	EtOH	HCl	H <sub>2</sub> O	СТАВ
NCT	0.15	0.85	40	0.24	5	0.14

**Table 3**: Structural parameters of the studied mesoporous silica films, samples are identified with the templating surfactant (CT for CTAB and F for Pluronic F-127) and with the temperature at which they were stabilized (130°C or 450°C). We reported porous volume Vp and wall thickness  $w_t$ . Pore size were calculated assuming elliptical pores for Pm3n structures, semi-major axis ( $r_1$ ) and semi-minor axis ( $r_s$ ) of the ellipse are reported. For p6m and wormlike structures a pore size distribution has been obtained assuming cylindrical pores with an elliptical base, which semi-axes are reported as semi-major axis ( $r_1$ ) and semi-minor axis ( $r_s$ ).

Sample	$r_l$ (nm)	<i>r</i> <sub>s</sub> (nm)	$w_t$ (nm)	$S (m^2/cm^3)$	Vp (%)	Structure	Surfactant
CT130-1	1.6	1.5	1.2	700	45.5	Pm3n	СТАВ
CT130-2	-	-	-	850	60.1	Pm3n	СТАВ
CT130-3	-	-	-	650	42.0	Pm3n	CTAB
CT450-1	1.6	1.4	1.0	750	51.0	Pm3n	CTAB
CT450-2 <sup>a</sup>	1.3	1.1	-	800	56.2	Wormlike	CTAB
CT450-3	1.6	1.4	1.1	850	62.0	Pm3n	СТАВ
CT450-4 <sup>b</sup>	-	-	-	600	43.0	Pm3n	СТАВ
CT450-5	-	-	-	600	43.1	Pm3n	СТАВ
F450	2.9	2.0	5.7	370	61.4	p6m	Pluronic F- 127
NCT130-1	-	-	-	600	42.0	Wormlike	CTAB
NCT130-2	-	-	-	600	42.2	Wormlike	CTAB

<sup>a</sup>: just after the film deposition, sample has been dried at relative humidity RH 75% and 37°C for 20min, to obtain a wormlike mesostructure. <sup>b</sup>: Sample left 5 days on laboratory bench before taking these data. Immediately after synthesis it had a  $V_p$  of 61%.



**Figure 1**: Experimental setup to investigate mesoporous silica thin film degradation in PBS by in situ ellipsometry. Both the liquid cell and the liquid reservoir are kept to a constant temperature of 37 °C.



**Figure 2**: GI- SAXS patterns and the related mesostructures of mesoporous thin films obtained by dip-coating from solution at 26 °C and 30-35 % RH: **a**) Silica templated with CTAB, Pm3n structure **b**) Hybrid organosilica templated with CTAB, wormlike structure **c**) Silica, templated with Pluronic F-127, p6m structure. More detail with scale bar is provided in SI 1.





**Figure 3**: Scheme of the dissolution process of a mesoporous silica film with (red arrows) dissolved silicate species diffusion pathways, and black and gray curve are silicate diffusion layer concentration profiles represented at two different dissolution times t1 and t2.



**Figure 4**: Bulk concentration (black line) and surface concentration (blue line) of dissolved silica for 1 cm<sup>2</sup> of mesoporous films having equivalent porous volume and template with **a**) CTAB, sample CT450-3 and **b**) Pluronic F-127, sample F450.



**Figure 5**: Concentration of dissolved silica at the solid-liquid interface after 100 seconds (squares) and 1000 seconds (circles) is reported versus surface area of the mesoporous silica films. Linear fit is showed. The point at the lowest surface value refers to dense silica while the porous samples have porosity values close to 62%. The surface area has been evaluated from EEP measurements with t-plot method, the error bars reported corresponds to an error on the surface value of 20%.



**Figure 6**: Thickness and  $C_s$  versus time for a) mesoporous silica CT450-3 film b) mesoporous silica F450 film.



**Figure** 7: *a*) TEM image of mesoporous silica nanoparticles templated with CTAB, particles have size of 40 nm and a p6m structure (see blow-up) *b*) Pore size distribution of NPs film and mesoporous silica plane film CT130-1, obtained from EEP measurements *c*) SEM image of NPs film deposed on silicon *d*) Dissolution curves of NPs film and plane film in PBS at 37°C.



**Figure 8**: Dissolved silica mass  $M_t$  normalized on initial silica mass  $M_0$  during dissolution in PBS at 37 °C of **a**) mesoporous silica layers templated with CTAB and exposed to different thermal treatment (130 °C and 450°C). Samples are CT130-2 (red line) and CT450-3 (black line). **B**) mesoporous silica layers templated with CTAB, heated at 450 °C, having different mesostructures; they have similar porosity (56% for the wormlike structure and 62% for the Pm3n) and surface area (800-850 m<sup>2</sup>/cm<sup>3</sup>), Samples are CT450-2 (black line) and CT450-3 (red line).



**Figure 9**: Dissolved silica mass  $M_t$  normalized on initial silica mass  $M_0$  during dissolution in PBS at 37 °C for mesoporous silica films (CTAB templated, Pm3n structure) treated at 450°C, before (black curve, sample CT450-3) and after (red curve, sample CT450-4) contamination from environmental VOC adsorbate. Porous volume is reduced from 62% to 43% and accessible surface area from 850 m<sup>2</sup>/cm<sup>3</sup> to 600 m<sup>2</sup>/cm<sup>3</sup>. The red curve presents two different slopes at the beginning of dissolution and after 15 minutes, showing an acceleration in dissolution rate due to pore opening.



*Figure 10: a)* Dissolution curve of mesoporous hybrid organosilica (solid line, sample NCT130-1) and silica (dashed line, sample CT130-3) in PBS at 37°C. *b)* XPS peak of nitrogen from hybrid organosilica sample before dissolution, at 400 eV we observe the peak of free amine and at 402.5 eV the peak of amine interacting with silanols *c)* XPS peak of nitrogen from hybrid organosilica sample after 3h of dissolution, the peak at 402.5 decreased.



**Figure 11**: *a)* Dissolution curves of mesoporous silica at 37°C in PBS (dashed line, CT130-3) and in BSA solution (solid line, CT450-5) *b)* picture of mesoporous silica films after 3h of dissolution in PBS (left) and BSA solution (right), after 3h in PBS there is no more silica left and the silicon substrate is visible, while after 3h in BSA solution a thick layer of silica is still present. *c)* Dissolution curves of hybrid mesoporous organosilica at 37°C in PBS (dashed line, NCT130-1) and in BSA solution (solid line, NCT130-2).

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**Figure 13**: Refractive index at 632 nm of a mesoporous silica layer during dissolution in BSA solution at 37°C for **a**) sample washed to remove CTAB **b**) sample heated at 450°C to remove CTAB.



**Figure 14**: SPR angle shift of mesoporous silica layer **a**) containing CTAB **b**) without CTAB. On the same sample were used parallel cells of  $1\mu L$  volume, in the first was flushed PBS buffer, in the second, PBS buffer and BSA solution were flushed alternatively. Arrows indicate the BSA solution injections, while the vertical lines indicate the buffer injections. The flow speed was  $10 \mu L/min$ . In figure b sample was washed with EtOH/HCl 100:1(5x 5min), kept overnight in EtOH and washed again (3x10min) before the dissolution experiment took place, to be sure to extract all CTAB from the mesoporous film.