NEW DEVELOPMENTS AND APPLICATIONS IN SOL-GEL COATINGS

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ABSTRACT
In this overview we describe some recent developments of technological interest realized at INM in the processing of sol-gel coatings. Nanoscale crystalline particles, fully redispersible in aqueous solutions, have been developed to produce thin ceramic layers at low temperature. As an example, antistatic optical coatings and conducting membranes based on SnO\textsubscript{2}:Sb material are reported. New wet chemical processes have been developed to extend the sol-gel basic principles for the coating of oxide compounds on plastic (biomimetic process) and sulfide materials (MoS\textsubscript{2}). Also ink-jet and pad printing processes, well established in other fields, have been adapted for the development of micro structured coatings. As examples, the realization of single and arrays of spherical and cylindrical refractive microlenses is presented. Finally non conventional sintering process using high power CO\textsubscript{2} laser is exemplified for the obtention of optical quality transparent conducting coatings under high sintering rate(30 cm\textsuperscript{2}/s).

INTRODUCTION
Most of the interest in sol-gel synthesis for thin film deposition was concentrated in the past years on the hydrolysis of alkoxides in organic solvent. Many single or multi components oxide coatings have been obtained using essentially the dip and spin coating processes and characterized but, in spite of the great success obtained in the laboratories, only a few systems are presently used on a commercial basis as the industry still gives a preference to more conventional deposition techniques such as sputtering, physical and chemical vapor deposition, etc. Consequently several issues are of technological interest at present.

In this paper we briefly report on some recent developments realized at the Institut für Neue Materialien - INM to fulfill some of these issues. The use of alcoholic solvents is sometimes considered as a drawback of the method and alternative sol-gel processes using aqueous media can often be just as useful and we show that narrow sized nanocrystalline sol-gel powders fully redispersable in aqueous media can be used to obtain optical quality coatings. The case of SnO\textsubscript{2}:Sb is exemplified for the development of antistatic coating and conducting membranes for ultra filtration \cite{1,2,3}. The sol-gel process has been essentially developed for oxide materials and wet chemical processes allowing the use of the simple and inexpensive dip or spin coating techniques of non-oxide materials have not been systematically investigated so far. The development of wet chemical processes using single source non-oxide precursor is then reported. The technique has been successfully applied to obtain thin MoS\textsubscript{2} layers using the spin coating process \cite{4}. New coating deposition techniques are also discussed. The ink-jet and pad printing processes have been developed to deposit structural coatings and optical micro components using hybrid inorganic-organic sols \cite{5,6} while the biomimetic approach is shown to be of interest to deposit hydrated TiO\textsubscript{2} thin films on polycarbonate substrate \cite{7,8}. Finally the use of high power CO\textsubscript{2} laser is exemplified as a non conventional high rate sintering process to obtain transparent conduction SnO\textsubscript{2}:Sb coating of high optical quality \cite{9}.
COATING OBTAINED WITH REDISPERABLE NANOCRYSTALLINE PARTICLES [1-3]

The application of the sol-gel methods based on the well known hydrolysis-polycondensation processes using metal alkoxides or oxoalkoxides based sols results after deposition to wet and usually amorphous coatings whose composition reflects a complex structure where the metal ion is linked to anionic ligand (alkoxide, acetate, OH, Cl etc.). The organic content is then burnt by heating at 300 - 500°C and at the same time the coatings densify and in most cases crystallize. A complete densification is however never obtained (or obtained at very high temperature) and the sol-gel layers are usually more porous than those obtained by other deposition process (CVD, sputtering, PVD, etc.). The preparation of coatings using already dense crystalline nano particles with a controlled size is therefore an interesting issue which may lead to coatings presenting higher density and better crystallinity after treatment at a lower sintering temperature. This requires the availability of sols loaded with fully redispersable nanocrystalline particles which can then be processed by the now conventional spin or dip coating techniques. The nanosize of the precursor components will also be a warranty to obtain high optical quality coating presenting low light scattering.

As an example SnO2:Sb nanoscale powders have been prepared by a controlled growth technique [1,10, 11]. A solution of tin (IV) chloride in ethanol containing 0.1 - 10 mole % of SbCl3 (dopant) was added drop-wise to an aqueous ammonia solution containing 10 wt% of a surface modifying agent, β-alanine. The prepared suspensions were hydrothermally treated at 150°C under a 3 bar pressure for 3 h. The resulting powder was isolated by centrifuging and washed with water several times and then dried at 60°C. The powders consist of crystalline particles (cassiterite structure) with average particle size of 3-5 nm without evidence of aggregation. These particles can be fully redispersed in water at pH > 11 under ultrasound using tetramethylammoniumhydroxide as dispersing agent [1,2] with solid content up to 50 mass%.

Deposition by spin coating (2000 rpm) results to coating whose thickness increases exponentially with the concentration (typically from ~100 nm for a 2.5 mass% to 225 nm for a 26 mass%). The electrical specific resistivity of single SnO2:Sb layer heat treated at 550°C for 15 min. decreases with the particle concentration from 10 Ωcm (2.5 mass%) down to a minimum value of ~4.2 10⁻² Ωcm for a particulated sol with 26 mass%. Higher concentration leads to cracks in the coatings increasing the resistivity. These values are unfortunately still higher than those obtained with a conventional sol-gel process, typically 2 10⁻³ Ωcm [12] or prepared by spray pyrolysis (ρ ≈ 9 10⁻⁴ Ωcm) [13]. The coatings are highly transparent in the visible spectral range with transmission as high as 92% and present interesting antistatic and optical properties.

The BET surface and density (measured by He picnometry) of the powders processed in air at different temperatures and sintering times is shown in figure 1. Particles with the density close to the theoretical density (≈ 7g/cm³) can only be obtained at high temperature (T> 600°C) after long sintering time (300 min).

Fig. 1: Influence of temperature and sintering time on the BET surface and the density of SnO2:Sb powder
At temperature and time normally used to sinter coating on float or borosilicate glass the particle density is 10% smaller. The pore characteristics (volume, size) of these powders under the same treatments are shown in figure 2. The average pore diameter can be adjusted from ≤ 4 nm to ~ 20 nm by varying the sintering conditions. The pore distribution is very narrow (± 1 nm at 400°C, ± 2 nm at 800°C) and the maximum pore volume constant (~ 0.75 cm³/g). As these particles can be deposited into crack free layers by spin coating (200 nm for 1 layer, with the possibility to increase the thickness by repeating the coating process [14]), such a process turns out quite interesting for the development of conducting ultrafiltration membranes [3].

Fig. 2: Evolution of pore characteristics (volume, size) with sintering temperature and time

SPIN COATING OF THIN SULFIDE COATINGS [4]

Various deposition techniques have been used for the preparation of sulfide films among which the physical and chemical vapor deposition, chemical bath deposition, electric deposition are the most important. Wet chemical processes allowing for simple and inexpensive spin coating process have so far not been explored. First results for the deposition of sulfides form different precursor solutions have been however reported for amorphous As₂S₃ [15,16,17] and As₂Se₃ [18], but similar attempts for the preparation of Sb₂S₃ and GeS₂ have not been successful [17].

We have developed a different approach in which a sulfide salt dissolved in an organic solvent allows to obtain a spinnable solution which wets the substrate and where the solvent can be removed by a thermolysis process. MoS₂ thin films have been successfully obtained in this way by dissolving (NH₄)₂MoS₄ in an organic diamine [4]) such as 1,2-diaminoethane [en], 1,2-diaminopropane [12dap] or 1,3-diaminopropane [13dap]. During the reaction NH₃ is released briskly under formation of the corresponding diammonium salt [H₂N-C₆H₄-NH₃]MoS₄ leading to deep red solutions. Thin film deposited by spin coating technique at 2000 rpm during 1 min on fused silica substrates are orange in color and turn red-brown after a drying step at 250°C in air due to the thermolysis of the diammonium salt during the formation of amorphous MoS₂. Under further thermolysis at temperatures between 600 and 800°C during 1 h under N₂ atmosphere, MoS₂ slowly decomposes into MoS₂ and sulfur and deep brown highly reflective crystalline films, 90 to 150 nm thick, are obtained. X-ray diffraction of [enH₃]MoS₄ based films, heat treated at T>600°C shows only one peak at 2θ = 14° corresponding to the (002) plane (Fig.3). The coatings are poorly crystalline and a careful analysis indicates that they are made of MoS₂ layer fragments essentially oriented with their basal planes parallel to the surface corresponding to a type II coating [19]. A high resolution transmission electron microscopy analysis reveals that these fragments have an average length between 5 and 10 nm and an average number of layers in each stack between 1 and 3. Between the small crystallites more or less amorphous areas are visible (Fig.4). [12dapH₃]MoS₄ based films show a different structure where the 2θ (002) peak is slightly shifted to a lower angle and an additional broad 2θ peak which cannot be indexed to any of the known molybdenum sulfides appears at 10.5°. HRTEM images reveal that the fragments are mostly two layered stacks bent in direction of the c-axis with an
increased space between the basal planes due to the occurrence of fork or dislocation defects (insertion of additional layers) or the intercalation of foreign molecules.

Fig. 3: XRD of MoS2 films prepared from 0.5 M solutions of (NH4)2MoS4 in 1,2-diaminoethane after heat treatment at temperatures of 600 - 800°C for 1 h.

Fig. 4: HRTEM structure images of MoS2 films prepared from 0.5 M solutions of (NH4)2MoS4 in 1,2-diaminoethane after heat treatment at a temperature of 600°C for 1 h.

The optical spectra of films prepared from solutions of (NH4)2MoS4 in 1,2-diaminoethane, 1,2-diaminopropane, and 1,3-diaminopropane after heat treatment at a temperature of 600°C for 1 h are shown in figure 5. The coatings are brown in transmission with a reflection between 30 and 40% in the visible range of the spectrum.

Fig. 5: Optical spectra of films prepared from solutions of (NH4)2MoS4 in 1,2-diaminoethane (en), 1,2-diaminopropane (12dap) and 1,3-diaminopropane (13dap) after heat treatment at 600°C for 1 h in N2.
From the corrected transmission $T_{corr}$ the high absorption $A = 1 - T_{corr}$ in the range from 200 to 600 nm becomes evident. The absorption coefficient at 550 nm $\alpha_{550}$ is about $2 \times 10^5$ cm$^{-1}$. From a plot of $\alpha^2$ and $\alpha^{0.5}$ vs. the energy the direct and indirect band gap were determined as $1.80 \pm 0.02$ eV and $1.20 \pm 0.05$ eV, respectively. These values are in good agreement with band gaps obtained with other deposition techniques and for bulk MoS$_2$. Differences in the optical spectra for 1,2-diaminoethane and 1,3-diaminopropane on the one side and for 1,2-diaminopropane on the other side may arise from the different degree of crystallinity of the samples. The strong absorption at ca. 2700 nm can be attributed to Si-OH vibrational modes in the substrate.

The development of wet chemical preparation reported here for the obtention of high optical quality MoS$_2$ thin film by the spin coating technique opens new possibilities for the development of chalcogenide based coatings.

INK-JET AND PAD PRINTING OF Si BASED HYBRID COATINGS

Sol-gel coatings are usually obtained by dip and spin coating technique. Patterning of these coatings is possible and techniques such as laser ablation, UV (lamp or excimer laser) irradiation or embossing have been used. It exists however on the market printing equipments which allow to print patterns on substrates.

The ink-jet process is of common use in computer controlled printing [20]. To our knowledge sol-gel based inks have been only recently proposed to decorate ceramic tiles using a continuous process [21] and for the realization of spherical microlenses [22]. The pad printing is widely used in the industry to print small flat or curved substrates [23] but, to our knowledge, no sol-gel based ink have been yet developed. Such technologies have been used to manufacture microlenses on glass substrates using sols prepared Iron hybrid organic-inorganic materials.

Using a commercial drop-on-demand ink-jet-equipment (Microdrop SP-K 130), spherical droplets, typically 50 µm in diameter, have been deposited on clean glass substrates and glass substrates coated with a thin layer of a low surface energy perfluorated polymer (FTS/TEOS/MPTS/2-Propanol/Irgacure 184) [24] in order to vary the drop/substrate wetting conditions. Hybrid organic -inorganic sols have been prepared by hydrolysis of methacryloxypropyltrimethoxysilane (MPTS) mixed with an ethanolic solution of tetraethyleneglycoldimethacrylate (TEGDMA) and a 1 to 10 wt.-% UV photoinitiator (Irgacure 184). They have a typical kinematic viscosity $\eta = 3$ mm$^2$/s at 20°C. After deposition, the drops have been polymerized by a 17.5 J/cm$^2$ UV light irradiation (Kombistrahler Beltron) within 200 s [22]. The deposited material is practically transparent ($T \equiv 91\%$) in the 375 nm $< \lambda < 2.7 \mu$m region and therefore adequate for the preparation of refractive lenses. Sols having a too high ethanol amount (>40 wt.-%) wet the glass substrate and spreads on it. The inorganic-organic material is transported to the border of the drop and creates a ring shape lens due capillary effect [22]. Sols containing an ethanol fraction smaller than 30 wt.-% have a high viscosity $\eta > 5$ mm$^2$/s, and tend to block the piezoelectric nozzle.

Spherical lens formation was obtained on glass substrate with 30 wt.-% ethanol fraction with a mixture of 50 mol-% TEGDMA (fig. 6). This lens is spherical and has the following characteristics properties: radius $a = 25 \mu$m, height $h = 6.25 \mu$m , radius of curvature $r = 53.1 \mu$m, focal length (at 632.8 nm) $f = 100 \mu$m, lens power $P = 0.62$ and a surface roughness $R_s = 40$nm.

![Fig 6: AFM micrograph of a lens obtained on a glass substrate with optimized ethanol content (30 wt.%)](image)
In order to reduce the effect of the spreading of the sol having a higher ethanol content, lenses have been fabricated on a substrate whose interface has been coated with a thin layer (0.1 μm) of a low surface energy polymer. Figure 7 shows a fit of vertical cross section passing through the center of the lens by a semi circle, showing that the lens is practically spherical. This lens has the following properties: radius $a = 37.5$ μm, height $h = 9$ μm, radius of curvature $r = 82.6$ μm, focal length at 632.8 nm $f = 125$ μm, lens power $F = 0.72$, surface roughness $R_s = 40$ nm. 2-d intensity profiles of the light spot created at the focal plane by a collimated 632.8 nm laser beam traveling onto the substrate side of the lenses are shown in Fig. 8. Their shapes are practically Gaussian with a full width at half maximum (fwhm) equal to 1.8 μm and 6.2 μm respectively. These values are small and only slightly larger than those calculated for a diffraction limited lens of same dimensions (fwhm ∼0.7 μm).

Reproducible arrays of hard, homogeneous, plano-convex microlenses have been obtained on fluorine modified glass substrates (Fig. 9).

Pad printing experiments were done with a commercial pad printer (Massek, PP21) using silicone tampons for transferring the sol from a steel cliché to a glass substrate. For these experiments the surface of the substrate was not modified by a polymer. A similar sol as for the ink-jet-printing prepared by
hydrolysis of MPTS was used. For the pad printing experiments no TEGDMA or UV-photoinitiator was added to the sol.

The sol prepared with a mol ratio MPTS: H₂O = 1:1.5 showed a dynamic viscosity \( \eta = 0.3 \) Pa\( s \) after pre-hydrolyzing for 24 h at 40°C. With additions of different amounts of water and further hydrolyzing at 60°C for 4 h at every step, dynamic viscosities \( \eta \) up to 15 Pa\( s \) were obtained. Lines and dots with diameters from 0.1-0.4 mm, etched 30 mm deep into a steel cliché were filled with the sol. After the printing process the coated substrate was heated at 200°C for 15 min.

Cylindrical and spherical lenses with good reproducibility have been printed. Figure 10 shows a 3-d plot of an array of 3 x 3 printed spherical lenses resulting from etched dots with 0.1 mm diameter in the cliché obtained with a sol having a viscosity \( \eta = 15 \) Pa\( s \). The lenses have diameters of ~0.2 mm indicating that the pad printing process leads to a broadening effect. This effect is strong for small structures (~0.1 mm) and for sols having a low viscosity but small for larger structures (> 0.3 mm) and sols having a high viscosity (> 10 Pa\( s \)). Figure 11 shows a fit of a vertical cross section passing through the center of one of the printed lenses by a semi circle. The lens is practically spherical and has the following properties: radius \( a = 90 \) mm, height \( h = 4.2 \) mm, radius of curvature \( r = 0.95 \) mm, focal length \( f = 1.9 \) mm (assuming \( n = 1.5 \)) and lens power \( F = 5.2 \).

Figure 12 shows a 3-d profilometer plot of two printed cylindrical lenses with widths of about 0.4 mm resulting from pad printing two etched lines of nearly the same widths in the cliché. The result shows that for structures of this size the pad printing process is able to transfer the etched structure in nearly identical size to the substrate. The shape of the lines corresponds to that of plano-convex cylindrical lenses. Figure 13 shows a profilometer plot of a vertical cross section and a fit of the shape by a circle. The lens has the
following properties: radius $a = 220$ mm, height $h = 8.2$ mm, radius of curvature $r = 3$ mm, focal length $f = 6$ mm (assuming $n = 1.5$) and lens power $F = 6.3$.

**BIOMIMETIC PREPARATION OF HYDRATED TiO$_2$ THIN FILMS ON POLYCARBONATE**

Film deposition on plastic substrates requires temperatures lower than 150 to 200°C. The conventional sol-gel process will lead to the formation of very porous and mechanically unstable layers. Wet chemical processes using hybrid materials where organics are incorporated into inorganic networks, are however better suited. Another possibility is provided by biomimetic processes which try to imitate biological systems [25]. Such methods are characterized by low process temperatures and moderate chemical environments. One approach in this field is the surface functionalisation, a technique which has already led to thin film synthesis of FeOOH, Fe$_2$O$_3$, SnO$_2$, CdS, Al(OH)$_3$ and TiO$_2$ [26].

The surface of a polycarbonate substrate (Makrolon 281, Röhm) can be sulfonated with a solution of SO$_3$ in H$_2$SO$_4$ and then etched by the formation of water soluble compounds. Hydrated TiO$_2$ coatings were then deposited from 0.05 M aqueous solution of (NH$_4$)$_2$(OH)$_2$Ti(C$_2$H$_5$O$_3$)$_2$ (Tyzor LA®, Dupont) and HCl in a Ti: HCl molar ratio of 1:1.95 by dipping the pretreated substrates into the heated solution (70°C) for 1 to 30 min. Figure 14 shows the variation of the film thickness as a function of the coating time. After a rapid growth, the thickness increases at a rate of 10 nm/min. 500 nm thick film has been obtained after 30 min.

X-ray diffraction measurements indicate that the layers are amorphous. The coatings are homogeneous, without cracks or islands (fig. 15), scratch resistant according to the tape test DIN 58196-K1, have an optical transmission in the visible range of ~ 82% with an index of refraction of ~ 1.9 at 550 nm and a probable composition of TiO$_2$ x H$_2$O. This process is therefore quite promising to coat thin oxide layers on polycarbonate substrates.

![Figure 14: Thickness of the deposited TiO$_2$ x H$_2$O thin film on sulfonated polycarbonate vs. the coating time](image1)

![Figure 15: REM image of a TiO$_2$ x H$_2$O coating on polycarbonate (30 min sulfonated, 10 min deposited)](image2)

**FAST CO$_2$ LASER SINTERING OF SnO$_2$:Sb COATINGS**

The sol-gel coatings obtained by dip or spin coating technique are usually dried and fired in two separate processes in a furnace. The heating and cooling which should be repeated to produce thick coatings are by far the speed-limiting factor of the coating process. An interesting technique using high power CO$_2$ laser whose radiation is absorbed by the glass substrate has been developed. It allows fast surface firing speed (up to 30 cm$^2$/s) of sol-gel layers up to temperature several hundred degrees higher than the $T_g$ values of the glass substrates films.

The technique has been so far successfully tested for transparent conducting SnO$_2$:Sb coatings [9, 27]. Using a fast scan mode crack free coatings have been sintered at temperature as high as 1000°C obtained in a time as short as 0.1s (heating rate up to 4300 K/s). Laser firing was found to produce denser layers with larger crystallites than those fired in a furnace at the same temperature leading consequently to coatings showing lower resistivity. A comparison of the results obtained by both processes made for SnO$_2$:Sb coatings having an initial thickness after deposition of 158 nm and then
sintered at the same temperature (540°C) gives the following values: density 4.8 g cm\(^{-3}\) vs 3.2 g cm\(^{-3}\), crystallite size 12 nm vs 5 - 6 nm, sheet resistance 0.9 K\(\Omega\) vs. 4.4 K\(\Omega\), resistivity 8.5 \(10^{-3}\) \(\Omega\) cm vs. 60 \(10^{-3}\) \(\Omega\) cm. These correlated variations are thought to be due to a competition between nucleation at low temperature and the growth of the nuclei occurring at higher temperature. A low heating rate (0.2 K/s in a conventional furnace) leads to a low densification and favors a porous structure of small crystallites while a high heating rate (> 1000 K/s with the CO\(_2\) laser) favors a denser structure built with larger crystallites and fewer pores.

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