One-step fabrication of high refractive index inorganic nanostructures

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Abstract
Direct printing of spin-on functional films is probably the most efficient method to develop low-cost novel photonic nanodevices, such as diffraction gratings, planar waveguides, nano- lasers, and antireflective coatings. For these applications high refractive index transparent materials are demanded; however, this class of materials generally requires inorganic oxides, well known for their hardness, typical of ceramic materials, and so incompatible with a soft character of printable resins. Herein, inorganic high refractive index TiO2 micro- and nano- structures, with unusual depth up to 600 nm and aspect ratio larger than 5, are obtained by combining thermal nanoimprint lithography (NIL) with UV curing. To achieve printed patterns, a hybrid organic-inorganic spin-on film is deposited at low-temperature by sol–gel processing. Two distinct bottom-up synthetic approaches are used, called in situ and ex situ, using titanium isopropoxide (90%) or TiO2 anatase nanoparticles (70%), respectively, and adding a silica sol modified by organic moieties. The two syntheses were optimized to obtain, after patterning by thermal imprint, amorphous or crystalline titania crack-free micro- and nano- patterns for in situ and ex situ, respectively. The further UV irradiation converts imprinted films to totally inorganic patterns, through the titania photocatalytic effect, allowing refractive indexes up to 1.82 at 632 nm to be achieved. This novel strategy of combining thermal imprint with UV exposure allows inorganic deep patterns to be fabricated without a calcination step, which is generally needed for inorganic resists processing. Eventually, a thermal treatment only at 300 °C can be applied to achieve a final refractive index of 2 at 632 nm.

Graphical Abstract
Direct printing of spin-on TiO2 high refractive index (1.82 @632 nm) transparent films, with depth up to 600 nm, aspect ratio larger than 5 and without lateral shrinkage are obtained combining thermal Nanoimprint Lithography (NIL) with UV curing.
Keywords  Nano-imprinting · Resist · Sol–gel · Hybrid organic-inorganic · TiO$_2$

Highlights
- A simple straightforward one-step surface patterning process by NIL technology of high refractive index TiO$_2$ systems is proposed.
- The hybrid nature of films combines the required intrinsic properties of inorganic titania and the simple processing of organic resists.
- Micro- and nano-inorganic structures up to 600 nm deep and refractive index up to 2 at 632 nm are achieved.

1 Introduction

Optical nanostructures, from diffraction gratings and waveguides to advanced geometries for light manipulation such as photonic crystals, have increased demands for development and applications. In particular, high refractive index materials and their patterning have acquired more and more importance in application ranging from ophthalmic lenses, filters, optical adhesives, highly reflective and anti-reflection coatings as well as advanced optoelectronic fabrications [1–9].

Titania, TiO$_2$, is certainly one of the most studied materials. However, it is conventionally patterned by lift-off [10, 11], a process that does not work well when thick films (some hundreds of nm) or complex patterns are needed. Other methods make use of different material templates or SAMs [12–17], allowing titania structures to be obtained by means of a sacrificial pattern acting as template that needs to be eliminated through high-temperature thermal treatment.

Some direct patterning techniques, such as electron beam, AFM, two-photon lithography and electro-hydrodynamic patterning, have then been used to obtain titanium oxides patterns [18–22]. However, these are mostly non-scalable imprinting processes.

Recently, much effort has been directed towards nanoimprint lithography (NIL) to obtain inorganic structures, since NIL is a low-cost, high throughput and high-resolution processes, presenting a potential industrial impact in the production of optical micro- and nano-systems [23–26], as was recently demonstrated for example by ultrafast nanoimprinting and roll-to-roll imprinting techniques. Few resists are generally available for NIL, but sol–gel methods enable spin-on inorganic film compositions suitable for NIL to be developed if properly designed and engineered. In fact, the few examples in the literature of direct imprint of inorganic TiO$_2$ systems reveal that some major challenges have to be faced when working with sol–gel compounds: (1) sol–gel films have high surface energy, causing mold release problems; (2) the solvent or organic moieties (chelating or coordinating agents) that help to ‘soften’ the material generally get trapped in the imprinted structures; and (3) sol–gel materials do not have a workable softening point, generally exhibiting a thermosetting behavior. As a consequence, inorganic titania patterned films obtained either by thermal (T-NIL) or UV assisted (UV-NIL) imprint, suffer from the following limits: (1) only shallow patterns, less than 100 nm [27, 28], can be obtained without crack formation; (2) only films with low titania content can be patterned [23, 29, 30]; (3) high shrinkages occur during processing up to 75% [31–33]; and (4) refractive index up to 1.98 at 632 nm [34] can be achieved only after annealing at 500 °C.

Soft lithography [35] helped to overcome these problems, since PDMS [36–38] and PFPE [36] soft molds, in contrast to the rigid stamps used in NIL, have low surface energies and a permeable nature that absorbs solvents typically used in sol–gel processing; nevertheless, this technique presents mold deformation problems, such as high shrinkages, and shallow structures, in particular, when sub-100-nm features are desired [39].

Herein, we propose a simple straightforward one-step patterning process of two transparent spin-on hybrid TiO$_2$ resists from wet chemistry (named in situ and ex situ) using T-NIL followed by UV curing. Both materials and processes benefit from several possible advantages. First, two different syntheses strategies (schematized in Fig. 1) are followed to obtain inorganic imprinted structures, i.e., using a Ti-precursor (in situ) or commercial anatase nanoparticles (ex situ), giving different final crystalline structures. Second, the hybrid nature of both films combines the required intrinsic properties of inorganic titania (e.g., high refractive index, up to 1.82 at 632 nm) and the simple processing of organic resists, which allows initially soft films to be easily imprinted and produces up to 600 nm deep micro- and nano- inorganic structures. Third, the composition and final refractive index of the patterned features can be easily tailored at room temperature through UV exposure, exploiting the photocatalytic activity of TiO$_2$. The overall imprinting process is schematized in Fig. 2. Finally, refractive index up to 2 at 632 nm can be eventually obtained with a 300 °C treatment if required for specific applications.
2 Results and discussion

2.1 Materials study and films behavior

2.1.1 Thermal behavior of in situ and ex situ films: rheology and FTIR analysis

The in situ and ex situ films were prepared by two different synthesis strategies: the first uses a Ti-precursor, Titanium (IV) isopropanoxide (TiOPr), and the second a pre-synthesized dense anatase nanoparticles in a titania sol. The specific compositions were selected after a preliminary set of NIL tests.

To control the overall process and achieve good quality patterns, we followed one of the most important parameters that influences the imprinting process: the viscosity of the resist and its variation with temperature. Both in situ and ex situ titania sols show a general increase of viscosity with temperature due to the crosslinking of precursors, with a slight decrease around 80 °C for the ex situ system just before a rapid increase at $T > 90$ °C (Fig. 3). It can be seen that the viscosities of the two sols are low with respect to the characteristic values of thermoplastic resists at typical imprinting temperatures. However, since the viscosity was evaluated on a thick sol film (around 500 µm), only the viscosity trends and not the absolute values reported in Fig. 3 are significant.

The most striking feature of Fig. 3 was the slow but continuous increase of viscosity versus temperature of the in situ sol. This fact indicates that crosslinking reactions take place just after the film deposition at room temperature, continue at increasing temperature and accelerate their kinetics between 80 and 100 °C. However, the
network formation was not completed even for $T > 140 \, ^{\circ}\mathrm{C}$, because viscosity is still increasing at the end of the measurement. On the contrary, the ex situ film viscosity was constant until $70 \, ^{\circ}\mathrm{C}$, slightly decreasing at $80 \, ^{\circ}\mathrm{C}$, followed by a rapid growth and stabilization at $T > 80–90 \, ^{\circ}\mathrm{C}$. At these temperatures, crosslinking reactions become very effective, suggesting a complete hardening at around $140 \, ^{\circ}\mathrm{C}$.

These diverse behaviors and the difference of one order of magnitude of viscosity values between the ex situ and in situ films, can be explained by analyzing the sol compositions: the in situ is a sol rich in pre polymerized TiOPr, that undergoes continuous crosslinking reactions with temperature; instead, the negligible amount of silicon and titanium alkoxides in the ex situ sol dissolved in a comparable amount of solvent prevents hardening of the network and requires higher temperatures in order to crosslink the alkoxides around anatase nanoparticles.

These observations were supported by FTIR spectra of films as-deposited and treated at increasing temperatures (Fig. 4a, b). The evaporation of the solvents (methoxyethanol, MEK and propanol) present in the films was the first feature clearly visible in both spectra: the doublets at 2930 and 2870 cm$^{-1}$ ($\nu_{as}(\mathrm{CH}_2)$ and $\nu_{s}(\mathrm{CH}_2)$, respectively) and the peak at 1100 cm$^{-1}$ due to --O--C-- bond decreased with temperature. Moreover, the peak at 2830 cm$^{-1}$ assigned to --OCH$_3$ of methoxyethanol and the peaks at 1470 and 1370 due to methyl C--H asym./sym. bend of MEK disappeared in both the in situ and ex situ spectra after $85 \, ^{\circ}\mathrm{C}$-treatment. The reduction of alcoholic solvents content in the films upon thermal treatment was finally confirmed by the decreasing of OH absorption at 3400 cm$^{-1}$, particularly evident for the ex situ after curing at $85 \, ^{\circ}\mathrm{C}$.

The abrupt decrease of the --OH band as well as all solvent absorbance peaks in the ex situ spectra at $85 \, ^{\circ}\mathrm{C}$ can be correlated to the sharp increase of viscosity around $90 \, ^{\circ}\mathrm{C}$ visible in Fig. 3, confirming it was a consequence of the sol increasing in concentration and crosslinking. Likewise, the continuous viscosity increase during curing of the in situ system is accompanied by a gradual reduction of --OH and --CH$_2$ (2930 and 2870 cm$^{-1}$) absorbances. In this latter case, condensation occurred also through hydrolysis of residual alkoxyl species (TiOPr and GPTMS), promoted by thermal curing, evident by the disappearance of the 2970 cm$^{-1}$ and 2830 cm$^{-1}$ absorbances. The former was due to the C--H bond of isopropoxide species in the titanium alkoxide precursor and the latter to unreacted Si--OCH$_3$ groups.

This viscosity analysis allowed the material’s structural evolution to be controlled during NIL and the correct
temperature to be chosen to avoid incomplete filling of the stamp, caused by premature or insufficient crosslinking of the film or the absence of solvents helping material flow during imprinting. Then, the time and pressure schedule were optimized for a good quality depth pattern and, in particular, to obtain a complete crosslink of the resist once the cavities were filled and to maintain the patterned shape after pressure removal.

Therefore, optimized schedules were selected to be: 20 min at 100–130 °C with a pressure of 30 MPa for the in situ system and temperatures ranging between room temperature to 80 °C for the ex situ system.

2.1.2 UV curing of in situ and ex situ

Resists are generally photo-cured to initiate polymerization reactions of organic monomers or degradation of UV absorbing species often coordinated to the inorganic network by chelating agents [40, 41]. The peculiarity of these TiO2 based resists is the possibility to convert patterned structures into inorganic oxide by photo-curing, determined from the photoactivity of TiO2 network that locally induce the degradation of organic species present in the material under UV illumination.

As can be seen by FTIR spectroscopy (Fig. 4a, b, black lines), after UV exposure the two sharp bands at 2930 and 2870 cm⁻¹, arising from the presence of residual organic component, progressively decreased and almost disappeared. The degradation of organic species was confirmed by the appearance of a peak at 1725 cm⁻¹ and by the reduction of the absorbance at 1100 cm⁻¹. The former peak is related to the absorption region of C=O bonds that can arise from the degradation of more labile organic species, such as the CH2–O–CH2 bond of the etheric group in the propylic chain of GPTMS or surfactants/dispersants used in the ex situ commercial dispersion [42]. The latter absorption was due to the degradation of the C–O bond of the ether group. The peaks at 1244 and 1200 cm⁻¹, related to the antisymmetric ring stretching of the epoxides and to the symmetric stretching vibration of CH2 groups in the propylic chain, respectively, and the aromatic ring vibration modes at 1613–1585 and 1510–1470 cm⁻¹ in the in situ spectrum (para-substituted phenyl groups stretching vibrations) decreased with UV exposure dose, indicating that ring opening and degradation of the organic chains occurred. Simultaneously, the growth of the band at 3400–3200 cm⁻¹ deriving from formation of H2O and –OH species and the appearance of the 1640 cm⁻¹ peak due to absorption of water molecules through an H bond with C=O groups, that are known to be very hydrophilic, were observed in both spectra. Finally, the slight increase of the 950 cm⁻¹ band absorbance with UV exposure can be ascribed to both condensation of the Ti–O–Si network and to Si–OH stretching vibrations [43–46]. Both absorbances may be present at this low treatment temperature.

All in all, these observations indicated that the UV photo-curing in the presence of the photocatalytic TiO2 network can initiate a marked densification, which is not achievable with a mild thermal curing, as is confirmed by the refractive index and thickness changes with UV dose reported in the following.

2.1.3 Refractive index change with temperature and UV curing

The refractive index and thickness of un-patterned films are reported in Fig. 4c, d. Only in situ films thickness showed an almost linear decrease accompanied by a slight refractive index growth, while the ex situ refractive index and thickness remained constant with thermal treatment. A comparison between in situ and ex situ thickness change showed that the shrinkage of in situ films after treatment at 130 °C is lower than 8%, while the ex situ film did not contract. At this temperature, the organic component (apart from the solvent) was still present in both films, but the densification of in situ sample was more efficient, as confirmed by the refractive index of films treated at 130 °C: 1.75 and 1.60 for in situ and ex situ, respectively. Moreover, even if in the ex situ film, the presence of anatase nanoparticles should further contribute to increasing the film’s refractive index, a poor compaction efficiency of the nanoparticles clearly took place with a consequent high final porosity of the film.

A 28% and 40% shrinkage, for in situ and ex situ films, respectively, was observed after UV exposure of films cured at the selected imprinting temperatures (130 °C for in situ and 80 °C for ex situ), accompanied by a refractive index rise up to 1.82 and 1.66 at 632 nm, respectively. Eventually, a refractive index up to 2 at 632 nm was reached with a thermal treatment at 300 °C for the in situ film.

However, these refractive index values compared with those of both amorphous and crystalline dense TiO2 (n around 2.4–2.6 [47]), suggested that the films were still quite porous, even taking into consideration the presence of the silica phase present in both in situ and ex situ films. The porosity of in situ films increased due to the removal of volatile organic species degraded under UV and from the presence of uncondensed TiOH bonds, while in the ex situ film it was the result of spaces between dense anatase nanoparticles that did not sinter at these low temperatures.

2.1.4 Crystalline structures of ex situ and in situ films

The crystallinity of the ex situ and in situ films, both as-deposited and UV exposed, was measured by XRD (Fig. 5). As expected, the (101) peak of the anatase phase at 20 = 25.3°
before and after UV irradiation was present in ex situ films (the patterns are compared with that of anatase particles dispersion used in the synthesis). On the contrary, no diffraction peaks were observed for the as-deposited in situ film and for the same film after 108 J/cm² UV dose exposure, indicating their amorphous structure.

It is well-known that the anatase titania phase shows the highest photoactivity compared to brookite or rutile phases, while amorphous titania generally exhibits negligible photocatalytic activity. It is in fact commonly accepted that amorphous metal oxides contain high concentrations of defects that will invariably function as rapid electron–hole recombination centers to render them inactive [48–50]. However, although in situ films were supposed to be totally amorphous, they probably displayed a relevant photocatalytic effect: a possible explanation is given by TEM analysis that revealed the presence of small quantities of a crystalline phase made of small crystals undetectable by XRD.

TEM analysis of the in situ films after 44 J/cm² UV exposure dose is reported in Fig. 5b; the occasional presence of crystalline phases dispersed in an amorphous background was detected and identified from diffraction analysis as anatase and rutile.

2.2 NIL of in situ and ex situ films

Due to their slow hardening, in situ films could be imprinted, down to sub-100 nm (Fig. 6b) at a range of temperatures from RT to the highest tested (130 °C), preserving sufficiently soft texture. NIL of in situ films confirmed the expectations from FTIR analysis, refractive index and rheological measurements. The network hardening and a complete filling of the cavities was reached in 20 min at 100–130 °C with a pressure of 30 MPa as confirmed by the crack-free flat profile of the structures, particularly evident in Fig. 6a.

In the case of the stamp with >0.5 um deep cavities, the difference between stamp depth and the resulting AFM profile of the structures was 30% (Fig. 7): in fact, with a 730 nm stamp depth, lines 510 nm high were obtained. Since the shrinkage of in situ flat films after treatment at 100 °C was lower than 5% (Fig. 3), a possible contribution of an incomplete filling of the structures has to be considered.

In the ex situ system, as expected from rheological measurements and FTIR spectra, the best imprint results, in terms of structure sharpness and uniformity of filling, were achieved with temperatures ranging between room
temperature to 80 °C and down to 100 nm lines (Fig. 6d). The possibility of obtaining imprinting at low temperatures was due to the low content of alkoxides in the ex situ system. The filling of the cavities was mainly achievable due to the presence of the solvent in the films up to 80 °C: the rheological curve showed, in fact, an abrupt increase in viscosity at temperature above 80–100 °C, mainly due to solvent evaporation with a minor contribution from alkoxide crosslinking. Structures with depth of up to 630 nm were produced with stamp cavity depths of 730 nm as can be seen from the corresponding AFM profile in Fig. 7: the measured difference between stamp depth and AFM profile of the structures is 15%, probably due to incomplete filling of cavities rather than shrinkage (which is negligible in flat ex situ films as already shown in Fig. 4).

Figure 8b shows an example of the application of these nanopatterned films as antireflecting coating. In the figure is shown an AFM image of a TiO₂ in situ hexagonal array of dense nanostructures achieved by imprinting a flat film, deposited on glass, with a Ni master. In Fig. 8c also the transparency of the films can be appreciated, as the flat films has a transmittance between 85 and 90%. The anti-reflective effect is also evident from the figure, as the transmittance increases after patterning, confirming the fidelity of master replication.

**2.3 UV curing of in situ and ex situ patterns**

After imprinting, patterned surfaces were UV exposed, with the minimum dose needed to eliminate organic component and transform the film in a totally inorganic material. The exposure was expected to lead to a further vertical and eventually lateral shrinkage, based on the data of Fig. 3. However, the shrinkage exhibited by in situ films was very low, as demonstrated by the AFM images of Fig. 7 after UV exposure. The lateral profile of simple gratings (period 1.5 µm and stamp depth 730 nm) revealed that the mean height of the gratings changed from 510 nm after imprinting to 479 nm with a resulting vertical shrinkage around 7%, much lower than a value of 28% measured on flat films treated at 130 °C and then exposed to the same dose, as shown in Fig. 3.

This effect is also more evident in ex situ films, that did not undergo vertical shrinkage after irradiation, as shown by the grating of Fig. 7 after UV exposure, with respect to a measured shrinkage of 40% shown in Fig. 3 after curing at 80 °C and UV exposure at the same dose.

Moreover, it is noteworthy that both in situ and ex situ patterns showed a relatively low vertical shrinkage, not accompanied by a lateral shrinkage that can be explained by the constraint deriving from the adhesion to the substrate. Therefore, in the imprinted structures, all the deformation necessary to compensate for the degradation of the organic part of the film was probably “absorbed” by vertical shrinkage. This behavior represents an important advantage simplifying the prediction of the final geometry of the structures: only vertical shrinkage must be considered during the design of the initial thickness. This behavior can be due to simultaneous application of pressure and temperature during the imprinting process, in which the thermal step is...
Fig. 7 SEM and AFM images of (left) TiO$_2$ in situ imprinted with a 730 nm depth stamp at 100 °C before and after exposure to 86 J/cm$^2$ UV dose; and (right) TiO$_2$ ex situ imprinted with a 730 nm depth stamp at room temperature and then exposed to 43 J/cm$^2$ UV dose.

Fig. 8 AFM images of TiO$_2$ in situ hexagonal array of dense nanostructures (b), imprinted with a Ni master (a), and antireflective effect of a patterned film deposited on glass (c). Master features: depth 140 nm, period 300 nm.
the HOI material, while in the ex situ one TiO2 NPs are synthesized of the HOI material, by hydrolysis and the titania network is formed simultaneously with the synthesis (Fig. 1).

4.1.1 In situ synthesis

The two methods used in our experiments to fabricate photosensitive hybrid organic-inorganic (HOI) sol–gel based spin-on TiO2 resists are named in situ and ex situ. In the in situ approach, the TiO2 is formed simultaneously with the synthesis of the HOI material, while in the ex situ one TiO2 NPs are added to the material already synthesized. The two systems were optimized in order to obtain, after patterning by thermal imprint, in the first case amorphous and in the second case crystalline titania crack-free micro- and nano-features with smallest sizes of 70 nm width and up to 630 nm deep. The further UV irradiation converted the imprinted films to totally inorganic patterns, without lateral shrinkage, loss of integrity or aspect ratio. Refractive indexes up to 1.82 at 632 nm and up to 2 at 632 nm were measured, without and with an eventual thermal treatment at 300 °C, respectively.

3 Conclusions

Direct printing of spin-on high refractive index transparent materials was demonstrated. Unusual depth up to 600 nm and aspect ratio larger than 5 were obtained by combining thermal NIL with UV curing and hybrid organic-inorganic spin-on films.

Two distinct bottom-up synthetic approaches, called in situ and ex situ were prepared. In the in situ approach, the TiO2 is formed simultaneously with the synthesis of the HOI material, while in the ex situ one TiO2 NPs are added to the material already synthesized. The two systems were optimized in order to obtain, after patterning by thermal imprint, in the first case amorphous and in the second case crystalline titania crack-free micro- and nano-features with smallest sizes of 70 nm width and up to 630 nm deep. The further UV irradiation converted the imprinted films to totally inorganic patterns, without lateral shrinkage, loss of integrity or aspect ratio. Refractive indexes up to 1.82 at 632 nm and up to 2 at 632 nm were measured, without and with an eventual thermal treatment at 300 °C, respectively.

4 Experimental section

4.1 Materials and synthesis

Titanium(IV) isopropoxide (TiOPr, purum, ≥97.0%), 3-glycidoxypropyltrimethoxysilane GPTMS and Bis[4-(glycidyloxy)phenyl)methane BFGDE were all purchased from Aldrich and used as received (Fig. 2). The dispersion of TiO2–anatase 15 nm particles in 2-Propanol (20wt%) was purchased from MKnano.

The two methods used in our experiments to fabricate photosensitive hybrid organic-inorganic (HOI) sol–gel based spin-on TiO2 resists are named in situ and ex situ synthesis (Fig. 1).

4.1.1 In situ synthesis

the titania network is formed simultaneously with the synthesis of the HOI material, by hydrolysis and condensation of TiOPr (90 mol%) and minor amounts (10 mol%) of silica precursor (GPTMS). GPTMS (molar ratio H2O:GPTMS=3:1) was first hydrolyzed in basic conditions for 1h at RT and 2 h at 80 °C under stirring in ambient air. 2-methoxyethanol, MetOEtOH, and NaOH (0.3mol% in GPTMS) were then added to the GPTMS solution in this order. Separately, TiOPr was added to MetOEtOH at a 1:1 volume ratio, and the solution was stirred for about 10 min. The two prepared solutions were then mixed and the monomer bisphenol F diglycidyl ether (BFDGE) (5% molar with respect to TiO2 content) was added to the solution to increase the processability and the final film thickness. The synthesis was carried out in MetOEtOH to obtain a final sol concentration of 100 g/l TiO2 content.

4.1.2 Ex situ synthesis

TiOPr was added to a dispersion of TiO2–anatase (15 nm particles mean size) in 2-Propanol (20wt%); this solution was loaded with 30 wt% of a previously prepared silica sol based on GPTMS, called GB,1 which improves film quality, increases layer thickness (up to 1.5 μm thickness, depending on the spin-coating speed) and makes possible film deformation during NIL. GB sol is obtained by GPTMS hydrolysis in basic conditions for 1 h at 80 °C (GPTMS: H2O: NaOH = 1:3:0.003, sol in MetOEtOH 150 g l−1 SiO2). The final amount of titanium isopropoxide is 10 mol% with respect to GPTMS.

The sol was left stirring for 20 min at RT followed by 10 min at 70 °C. The solvent (MEK) was added to obtain the final sol concentration of 90 g/l (TiO2), and a SiO2: TiO2 molar ratio of 1:3.

The sols were filtered using a microporous membrane (0.2 μm Millipore) to remove large particles and improve the resist spin-coating quality and deposited by spin-coating technique at different speeds to obtain films with the desired thickness.

Glass or silicon substrates were used for the following characterization or processes. The substrate was cleaned using plasma procedures in order to remove contaminants which may lead to poor adhesion or defects formation in the resist layer. Films were deposited by spin-coating the fresh sols immediately after preparation. By using different speeds, variable thickness was obtained, up to 1.5 μm for the ex situ film.

4.1.3 Characterization

the compositions of the films deposited on silicon wafers were examined by a Fourier transform infrared spectrometer (Jasco FT-IR-620) in the range of 400–4500 cm−1 with a resolution of 4 cm−1, on the basis of literature data [43–46].
peaks are labeled as follow: 1- two sharp bands at 2930 and 2870 cm$^{-1}$, 2- peak at 1725 cm$^{-1}$, 2' peak at 1100 cm$^{-1}$, 3- peaks at 1244 and 1200 cm$^{-1}$, 3' - set of bands of aromatic ring vibrations at 1613–1585 and 1510–1470 cm$^{-1}$ for the in situ film, 4- band at 3400–3200 cm$^{-1}$, 5- peak at 1640 cm$^{-1}$ (partially overlapped with peak at 1725 cm$^{-1}$), 6- 950 cm$^{-1}$ absorbance band.

The UV–Vis absorption of the films, deposited on glass substrates, was characterized using a UV–Vis spectrophotometer (JASCO V-570) in the wavelength range 200–600 nm.

The film thickness was examined using a profilometer and the quality of the gratings was characterized by Atomic Force Microscopy (NT-MDT) and SEM (LEO 1550).

The effects of thermal treatment and UV exposure on the refractive index and thickness of the hybrid titania sol–gel films were recorded by ellipsometric measurements (V-Vase J.A. Woollam Ellipsometer).

The film structure and titania crystalline cluster size were investigated through TEM (Jeol 3010 apparatus operated at 300 kV with a high-resolution pole) and X-ray diffraction (XRD) using an X-ray diffractometer (Philips PW 1729) with Cu Kα irradiation.

4.1.4 Imprint lithography conditions

the synthesized HOI titania films up to 1.5 μm were spin coated on silicon substrates and imprinted at temperatures ranging from room temperature (RT) to 130 °C with pressures between 20 and 40 MPa. Stamp materials, silicon (purchased from Silicon materials) or ORMOSTAMP (Micro resist technology), possessing multi micro- to nano- metric feature patterns and feature depths from 150 to 730 nm, and Ni stamp (NIL-....) with antireflecting features (depth 140 nm, period 300 nm) were used. The anti-adhesion layer was obtained using stamp silanization with 1H,1H,2H,2H-Perfluorooctyl-trichlorosilane 97% (Aldrich) in the vapor phase under vacuum for 3 h.

After thermal imprinting (T-NIL) the patterns were UV exposed at doses of 86 or 43 J/cm$^2$ (for in situ and ex situ respectively) selected to obtain totally inorganic features and eventually treated at temperatures of up to 300 °C for 30 min to further increase the refractive index.

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Compliance with ethical standards

Conflict of Interest The authors declare no competing interests.

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