A fast thermal-curing nanoimprint resist based on cationic polymerizable epoxysiloxane

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Abstract

We synthesized a series of epoxysiloxane oligomers with controllable viscosity and polarity and developed upon them a thermal-curable nanoimprint resist that was cross-linked in air at 110°C within 30 s if preexposed to UV light. The oligomers were designed and synthesized via hydrosilylation of 4-vinyl-cyclohexane-1,2-epoxide with poly(methylhydrosiloxane) with tunable viscosity, polarity, and cross-linking density. The resist exhibits excellent chemical and physical properties such as insensitivity toward oxygen, strong mechanical strength, and high etching resistance. Using this resist, nanoscale patterns of different geometries with feature sizes as small as 30 nm were fabricated via a nanoimprint process based on UV-assisted thermal curing. The curing time for the resist was on the order of 10 s at a moderate temperature with the help of UV light preexposure. This fast thermal curing speed was attributed to the large number of active cations generated upon UV exposure that facilitated the thermal polymerization process.

Keywords: Nanoimprint, Epoxysiloxane, Transfer layer, UV-assisted thermal curing, Cationic polymerization

Background

Resist is one of the key components for nanoimprint lithography (NIL) in addition to the imprint mold, the imprint machine, and imprint processes [1,2]. Depending on the process, nanoimprint resists can be classified into two basic categories: thermoplastic polymers [3] or thermal curable materials [4,5] for thermal-NIL, and UV-curable monomers or oligomers for UV-NIL [6–9]. The most commonly used NIL resists in earlier years were thermal plastic polymers, which required high pressure and high temperature so that the mold can be pressed into the molten resist film. Curing of thermal curable resists, although it requires lower pressure, is a time-consuming process due to the low speed of thermal-initiated polymerizations. Recently, UV-NIL has been developed for device fabrication, and it allows for imprinting at a low pressure and room temperature [6,10,11]. The process involves pressing a transparent mold into a low viscous photo-curable liquid thin film on a substrate and then solidifying the liquid materials via a UV light irradiation. The liquid resist can automatically fill the recess portions of the features on the mold due to capillary effect. An ideal UV resist usually has low viscosity, low surface tension, good adhesion to the substrate, fast cross-linking speed, high mechanical strength, and high etching resistance after cross-linking.

The fundamental ingredients of a UV resist include a UV-curable matrix, a photo-initiator, and other additives such as plasticizer, curing accelerator, photo-sensitizer, flow and leveling agent, fluorinated surfactant, and so on. Acrylate-based resins have been widely used in various UV-curable material systems as the backbone because of their great reactivity with a wide choice of acrylated monomers and oligomers [12,13]. However, a significant drawback of acrylated materials is that free radical polymerization is strongly inhibited by oxygen scavenging of the free radicals. This prevents UV-curing imprint process from being operated in ambient air environment, increasing the cost of equipment. To address this issue, materials such as vinyl ethers [14] and epoxides [15] based on cationic polymerization were used for UV-curable nanoimprint resists, which effectively cross-link upon UV exposure in the presence of air. However, vinyl ether...
resins are costly because of a narrow choice of commercially available monomers and oligomers. The cationic polymerization rate of epoxy groups is low, \[16\] and the conversion rate is less than 10\% when the temperature is lower than 50°C \[17,18\]. Recently, new epoxy-based materials were developed for imprint resists \[19,20\]; the mechanism of the polymerization is not fully revealed. Furthermore, these resists can only be cross-linked by UV light, so a transparent mold or substrate is required. On the other hand, epoxy-based negative photoresist, such as SU-8, was used for combined UV and thermal NIL, but it required very high imprint pressure (as high as 30 bar) to press the mold into the molten resist \[21–23\].

Herein, we report on a new liquid NIL resist based on epoxysiloxane oligomers that can be thermally cured within a short period of time in air at a moderate temperature using a UV-assisted thermal curing process. We designed and synthesized these oligomers via hydrosilylation of 4-vinyl-cyclohexene-1,2-epoxide with Si-H group-functionalized siloxane precursor. The viscosity, cross-linking density, and polarity of epoxysiloxane were controlled on demand by employing siloxane precursors with varied molecular weights and Si-H functionality. We also developed a fast UV-assisted thermal imprint process that requires low pressure, moderate temperature without the demand for optically transparent mold and vacuum equipment. We fabricated nanostructures of different geometries and sizes with the resist and further demonstrated high aspect ratio pattern transfer by reactive ion etching (RIE).

**Methods**

**Materials**

Polymethylhydrosiloxane (PMHS) precursors with different Si-H functionality, (15\% to 18\% methylhydrosiloxane)-dimethylsiloxane copolymer, (50\% to 55\% methylhydrosiloxane)-dimethylsiloxane copolymer and polymethylhydrosiloxane terminated with trimethylsilyl, and p-(octyloxyphenyl)phenylidonium hexafluoroantimonate (95\%, photo-acid generator) were purchased from Gelest, Inc. (Morrisville, PA, USA). A 4-vinyl-1-cyclohexene 1,2-epoxide (VCHE) mixture of isomers, propylene glycol monomethyl ether acetate (PGMEA), and chlorotris(triphenylphosphine) rhodium(I) (Rh catalyst) were purchased from Sigma-Aldrich Corporation (St. Louis, MO, USA). Lift-off layer (LOL) 2000 was purchased from MicroChem Corp (Newton, MA, USA). All materials were used as purchased.
Synthesis and characterization

The hydrosilylation of olefinic functionalized compounds is one of the most reliable ways to introduce a desired functionality as a side- or an end-group on Si-H containing siloxane polymers [24]. The synthesis approach for the epoxysiloxane in our study is shown in Figure 1. We used the Rh catalyst instead of Pt-based catalyst because the latter can cause the ring-opening polymerization of epoxy groups and can lead to an undesirable gelled product [25]. PMHS (40 mmol of Si-H), VCHE (5.96 g, 48 mmol), toluene (30 ml), and chlorotris(triphenylphosphine)rhodium(I) (5.4 mg) were added to a 100-ml three-neck flask that was equipped with stirrer, thermometer, and reflux condenser. After chlorotris(triphenylphosphine)rhodium(I) was dissolved, the system was stirred at 35°C for 24 h. The temperature was then raised to 80°C, and the compound was constantly stirred until the end of the reaction (when the Si-H infrared absorption peak at about 2,150 cm$^{-1}$ disappeared). The resulting liquid was purified via flash column chromatography using petroleum ether-ethyl acetate (10:1 volume ratio) as the eluent to yield a light yellow-colored compound. Infrared spectra and solution state hydrogen-1 nuclear magnetic resonance (1H NMR) spectra were recorded via a Bruker Vector-22 (BRUKER AXS GMBH, Karlsruhe, Germany) Fourier transform infrared (FTIR) spectrometer and a Bruker DRX-500 spectrometer, respectively. In NMR characterization, all chemical shifts are reported in parts per million downfield from tetramethylsilane (TMS) using the residual protonated solvent as an internal standard (CDCl$_3$, $^1$H 7.29 ppm). The viscosity of the epoxysiloxanes and their mixtures was measured using HAAKE RheoStress 600 rotational rheometer from Thermo Fisher Scientific (Waltham, MA, USA). Young's modulus of cured epoxysiloxanes was examined by Series IX Automated Materials Testing System from Instron Corporation (Norwood, MA, USA).

Imprinting and pattern transfer

The synthesized epoxysiloxane was diluted in propylene glycolmethyl ether acetate (5 wt.%), and the solution was filtered before being used as a resist. The UV-assisted thermal curing NIL process is schematically illustrated in Figure 2. Uniform liquid resist films were spin-coated on bare silicon wafers with or without prebaked LOL as the transfer layers. The resist film was then exposed briefly to a UV light source (for example, the lamp used on a Karl Suss MA-6 mask aligner with 240 to 450-nm wavelengths and a cold light mirror) with an optimized UV dose of 300 mJ/cm$^2$ (Figure 2a). An imprint mold of quartz or silicon was then pressed into the thin resist film while the substrate was heated at 110°C for 30 s before mold separation. All imprint experiments were conducted in air. The etching of the epoxysiloxane resist and LOL was carried out in an Oxford RIE system. Typical etching recipes were 40 sccm O$_2$, 3.0-mTorr pressure, 50-W power, $-20°C$ (helium flow rate, 4.3 sccm) for LOL and 60 sccm CF$_4$, 2.0-mTorr pressure, 40-W power, 20°C for the cross-linked resist films.

Cross-linking degree characterization

Solubility measurements of the resist films under various curing conditions were taken to check if the resist film was fully cross-linked. The measurement was simplified by rinsing the resist films with acetone after curing. Once fully polymerized, the thin film is insoluble in acetone due to the formation of highly cross-linked polymer network. An uncured or partially cured film would be stripped or destroyed by acetone [26].

Figure 3 FTIR spectra of PMHS polysiloxanes reacted with VCHE at different stages. (a) Pure PMHS precursor; (b) reacted at 35°C for 24 h; (c) reacted at 35°C for 24 h and then at 80°C for 24 h.
Results and discussion
Figure 3 shows the FTIR spectra at different reaction stages for the hydrosilylation reaction of PMHS with VCHE. The adsorption peak at 2,150 cm$^{-1}$ was attributed to the Si-H group. The absorption intensity decreased with the extension of reaction time, while new adsorption peaks appeared at 2,920 and 2,850 cm$^{-1}$ attributed to the -CH$_2$- group, suggesting that VCHE had been grafted onto the PMHS backbone by hydrosilylation. When the temperature was raised to 80°C and reacted for 24 h, the peak at 2,150 cm$^{-1}$ disappeared, indicating the Si-H groups had been completely consumed by VCHE. Figure 4 represents the $^1$H NMR spectrum of the epoxy-siloxane oligomer synthesized from PMHS with a Si-H content of 50% (mole percentage). The characteristic groups on the chemical structure of the aimed product and their positions in the NMR spectrum were labeled. The spectrum illustrated a complete absence of Si-H proton peak at $\delta = 4.7$ ppm, and the presence of the characteristic epoxide, Si-CH$_2$- and Si-CH$_3$ proton resonances at $\delta = 3.1$ to 3.2, 0.47, and 0.08 ppm without peaks of methyl and methylene of oxyethyl group at $\delta = 1.28$ and 3.65 to 3.71 ppm, respectively. It was confirmed that the aimed epoxy-siloxane was successfully obtained, and the competing epoxide ring-opening reaction was avoided. Using this synthesis approach, a series of multifunctional epoxy-siloxane with varied viscosity and epoxy functionality were prepared. Various properties of different oligomers and their cross-linked films are listed in Table 1. Since the hydrosilylation proceeded to high conversion without the formation of by-product, desired epoxy functionality could be obtained from the starting PMHS with the same Si-H functionality. The viscosity and its growth range of the epoxy-siloxane increased with the increase of Si-H content of the PMHS. This was attributed to the link of the epoxy groups to PMHS that increased not only the molecular weight, but also the polarity of the epoxy-siloxane. The increase of epoxy content of the oligomers also strongly influenced the properties of their cured films. Higher cross-linking density hindered the movement of the chain segments and made the cross-linked matrix more rigid but brittle, which resulted in higher modulus and lower strain. The epoxide and its ring-opening product, ether, were polar groups, so the increase of epoxy content of the siloxane led to the increase of the polarity of its cured film and the decrease of the water contact angle. As shown in Table 1, the desired parameters of epoxy-siloxanes and their cured film, such as viscosity, polarity, and mechanical strength, could be further optimized via mixing the

| Epoxysiloxane | Si-H of PMHS (mol%) | Viscosity of PMHS (mPa s) | Viscosity of epoxy-siloxane (mPa s) | Water contact angle (degrees) | Strain at peak (%) | Young’s modulus (MPa) |
|---------------|---------------------|--------------------------|----------------------------------|-----------------------------|------------------|---------------------|
| I             | 100                 | 33                       | 257                              | 74.9                        | 1.085            | 381,930             |
| II            | 50 to 55            | 15                       | 60                               | 83.7                        | 1.813            | 114,553             |
| III           | 15                  | 28                       | 37                               | 98.9                        | 15.100           | 10,379              |
| IV$^a$        | -                   | -                        | 52                               | 91.3                        | 6.300            | 88,617              |
| V$^b$         | -                   | -                        | 112                              | 80.6                        | 1.232            | 225,063             |

$^a$IV = 60 wt.% II + 40 wt.% III; $^b$V = 20 wt.% I + 80 wt.% II.
epoxysiloxanes with different epoxy functionality and viscosity at different ratio.

Due to their insensitivity toward oxygen during curing, mechanical strength, and resistance to oxygen plasma after cross-linking, epoxy-containing silicone resins are highly attractive for making lithography resists [27,28]. We prepared NIL resists with the synthesized products, and the ingredients and their functions of a typical NIL resist formulation are listed in Table 2. In this resist, the multifunctional epoxy groups in the curable materials enable a high curing speed, a high cross-linking degree of the resist, and enhanced mechanical strength. Epoxysiloxane I bearing one epoxy group per repeating unit is a cross-linker to intensify the mechanical strength of the resist. Epoxysiloxane II with a moderate viscosity and epoxy functionality is used as a major ingredient to provide a proper mechanical strength and high oxygen RIE etching resistance. Epoxysiloxane III serves as a reactive plasticizer to reduce the brittleness of the cured resist film due to its high cross-linking density. In addition, the resist consists of a diaryliodonium photo-acid generator and an organic solvent as a diluent for spin coating. By adjusting the mixture ratio of different epoxysiloxanes, we fine-tuned the viscosity and polarity of the resist, which were critical to the film forming ability on a substrate. For example, it was difficult for a resist of too low viscosity and polarity such as Epoxysiloxane III to form a thin uniform film on a Si wafer by spin coating method (Figure 5a). Due to surface tension of the liquid film, the film ruptured into polygonal networks of liquid rims or sets of liquid droplets as have been observed in other resists [29–31]. However, a resist with optimized viscosity and polarity synthesized in this study formed a continuous and uniform film on the Si substrate after spin coating (Figure 5b). Nanoscale and microscale features with a variety of geometries have been faithfully imprinted into the resist films (Figure 6). First, groups of 50-nm half-pitch nanowires, 50-μm-wide contact pads, and the fan-out structure connecting them (1,000 times difference in lateral scale) were simultaneously imprinted into the resist using the same mold over a large area (Figure 6a, b). Second, complex nanostructures such as arrays of fourfold symmetric L-shaped features with sharp corners were imprinted with high uniformity (Figure 6c, d). The uniform color of the patterned region in the optical micrograph suggested minimal residual layer thickness variation over the 1 mm × 100 μm imprinted area (Figure 6c). The high fidelity duplication of the sharp corners (Figure 6d) without peel off proved that the resist has excellent cavity filling properties as well as easy release from the mold and strong adhesion to the substrate. Third, dot arrays of 100-nm pitch and 30-nm diameter were imprinted into the resist, demonstrating a 30-nm resolution. It is worth noting that the resolution was mainly limited by the feature size on the imprint mold, and it is possible to imprint features much smaller than 30 nm provided such a mold is available.

### Table 2 Ingredients of a typical NIL resist used in this study and their functions

| Ingredient          | wt.% | Function                                                                 |
|---------------------|------|--------------------------------------------------------------------------|
| Ia                  | 1    | Silicon-containing oligomer; increases the viscosity and polarity of resist and provides high cross-linking density |
| IIa                 | 4    | Silicon-containing oligomer; provides oxygen RIE etching resistance and moderate cross-linking density |
| IIIa                | 0.5  | Silicon-containing oligomer; relieves mechanical properties as plasticizer |
| PGMEA               | 94.2 | Low-viscosity diluents; improves resist flow for spin coating            |
| Diphenyliodonium salt | 0.3 | Photo-initiator; generates cationic acids upon exposure to UV radiation   |

*I, II, and III are the same epoxysiloxane samples listed in Table 1.*
The cured resist exhibits excellent etching properties. The silicon-containing materials in the resist provide high resistance to O2 RIE, opening the opportunities for high aspect ratio nanostructure fabrication provided that a proper underlying transfer layer is used. We started with an 80-nm-thick resist film (see Table 2 for composition) on a 200-nm-thick LOL on Si, followed by UV-assisted thermal NIL using a mold of 200-nm pitch gratings (60-nm-wide and 82-nm-deep trenches). After removing the residual epoxysiloxane layer using RIE with CF4 gas, the LOL was etched using O2 plasma with the remaining epoxysiloxane as an etching mask. Figure 7 shows the 200-nm pitch etched epoxysiloxane/LOL lines of 60-nm width and 280-nm height (an aspect ratio of 4.7) that have vertical sidewalls (Figure 7).

Most currently available thermo-curable resists require lengthy curing cycles (e.g., epoxy or (meth)acrylate-based resists). This is because the thermally catalyzed polymerization for these materials is very slow. Our UV-assisted thermal curing process significantly shortens the
curing time. In our current process, the UV exposure before thermal curing quickly generates a large amount of active cations in the resist through a mechanism as described in Figure 8. These cations facilitate the cross-linking of the resists through the ring-opening polymerization mechanism (Figure 8). Due to the long lifetime of the cations, the polymerization of the epoxy will continue effectively even when the UV light is removed. Furthermore, this process can be significantly accelerated by heating, leading to the full polymerization of the epoxy group in the resist within 30 s.

We have quantitatively studied the dependence of the thermal curing time on the temperature and preexposure UV dose for a 200-nm-thick resist, as shown in Figure 9. It is observed that a higher UV exposure dose and a higher temperature significantly reduce the time needed to fully cross-link the resist film. On one hand, without heating, the resist preexposed to UV radiation does not cross-link at all. This is because the conversion level of the cationic polymerization of epoxide was less than 10% as the reaction temperature below 50°C. Raising the polymerization temperature above 60°C caused a sharp increase in the conversion level [19,22]. On the other hand, without UV exposure, the resist only starts to polymerize at a temperature above 110°C, and the time needed to fully cross-link the resist is on the order of 10 min. The photo-acid generator, diaryliodonium salt, was thermally unstable at the temperature over 110°C and pyrolyzed into the same catalytic cation pieces to initiate the polymerization. While the pyrolysis of the photo-acid generators might be slower than the photolysis of it, thus, it took much longer time to fully cross-link the resist without UV radiation. Based on our experiments, we have determined the optimized dose to be the minimum dose (300 mJ/cm²) when enough photo-acid generators are turned into cations. If the dose is too low, there would not be enough cations to initiate the polymerization of epoxy groups, while a higher dose is not necessary, and it actually leads to longer processing time for the imprint.

Finally, in contrast to the photo-initiated radical polymerization, photo-initiated cationic polymerization can be implemented in the presence of oxygen. The resist also has long shelf time due to the fact that the propagating polymer cations are not reacting with themselves [12]. We have achieved similar imprint results with resists that were prepared a year earlier and have been kept in dark bottle at room temperature. This indicates that the shelf time of the resists is at least 1 year.

![Figure 7](image1.png)

**Figure 7** High aspect ratio pattern transfer using two-step RIE. The inset shows a high magnification SEM image of 60-nm-wide, 280-nm-high grating lines of 200-nm pitch in which the high aspect ratio (4.7) of the resist stacks is demonstrated. Samples were tilted 45° for the cross-section imaging.

![Figure 8](image2.png)

**Figure 8** Formation of cross-linked polymer network by photo-initiated cationic polymerization of multifunctional epoxysiloxane.

![Figure 9](image3.png)

**Figure 9** The correlation of curing time with UV-exposure dose and heating temperature. A higher UV dose for preexposure and a higher curing temperature greatly shorten the thermal curing time.
Conclusions
We have synthesized a series of epoxysiloxane oligomers with controllable viscosity and polarity and developed a fast curable nanoimprint resist based on them. The new imprint resist was thermally cured within 30 s at a moderate temperature when a preexposure to UV light was adopted. Various patterns with feature sizes ranging from 30 nm to 50 μm were faithfully duplicated onto the resist film during this UV-assisted thermal curing NIL that was carried out in ambient air environment. The short cross-linking time was attributed to the large amount of cations that accelerated the thermal polymerization process. The unique properties of the resist enabled a novel imprint process that combines the advantages of both thermal and UV-curable NIL such as low pressure, short processing time with a wider choice of imprint mold materials.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
JZ, XH, JZ, and YC carried out the synthesis and characterization of the epoxysiloxanes. QX carried out the nanoimprint and RI experiments. HG, QX, WW, CY, and YC conceived the study and participated in its design and coordination. HG and QX drafted the manuscript. All authors read and approved the final manuscript.

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