Photolithography in the vacuum ultraviolet (172 nm) with sub-400 nm resolution: photoablative patterning of nanostructures and optical components in bulk polymers and thin films on semiconductors

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Precision photoablation of bulk polymers or films with incoherent vacuum ultraviolet (VUV) radiation from flat, microplasma array-powered lamps has led to the realization of a photolithographic process in which an acrylic, polycarbonate, or other polymer serves as a dry photoresist. Patterning of the surface of commercial-grade, bulk polymers (or films spun onto Si substrates) such as poly-methyl methacrylate (PMMA) and acrylonitrile butadiene styrene (ABS) with 172 nm lamp intensities as low as ~10 mW cm\(^{-2}\) and a fused silica contact mask yields trenches, as well as arbitrarily-complex 3D structures, with depths reproducible to ~10 nm. For 172 nm intensities of 10 mW cm\(^{-2}\) at the substrate, linearized PMMA photoablation rates of ~4 nm s\(^{-1}\) are measured for exposure times \(t \leq 70\) s but a gradual decline is observed thereafter. Beyond \(t \sim 300\) s, the polymer removal rate gradually saturates at ~0.2 nm s\(^{-1}\). Intricate patterns are readily produced in bulk acrylics or 40–200 nm thick acrylic films on Si with two or more exposures and overall process times of typically 10–300 s. The photoablation process is sufficiently precise that the smallest lateral feature size fabricated reproducibly to date, ~350 nm, appears to be limited primarily by the photomask itself. Examples of the versatility and precision of this photolithographic process include the fabrication of arrays of aluminum nanomirrors, each atop a 350 nm or 1 \(\mu\)m-diameter Si post, as well as optical components such as transmission gratings or Fresnel lenses photoablated into PMMA.

1. Introduction

Rapid progress in the advancement of photolithography over the past four decades is largely responsible for the success of the semiconductor device industry in keeping pace with Moore’s Law.1–3 Early optical exposure systems relied on the g- and i-line radiation of atomic Hg (436 nm and 365 nm, respectively) generated in arc lamps, and patterning processes for a broad range of electronic and photonic devices continue to be based on this incoherent source to this day. However, it was the transition to the shorter wavelengths afforded by the ArF (193 nm) and KrF (248 nm) excimer lasers, introduced to photolithography by Jain et al.4 in 1982, that accelerated the drive to realizing sub-100 nm features and, specifically, has resulted in the remarkable achievement of reaching the 14 nm node with 193 nm exposure systems.5,6 Although pulsed, deep-ultraviolet lasers (having wavelengths of ~200–250 nm) and, more recently, laser-produced tin plasma sources at 13.5 nm (in the extreme ultraviolet (EUV) spectral region) can be credited with the relentless shrinkage in feature sizes, this progress has come at a substantial cost. Increasingly sophisticated steppers and the expense associated with their acquisition and maintenance have progressively limited access to state-of-the-art photolithographic tools. For a significant fraction of the patterning processes required for research or the manufacturing of photonic and electronic devices and systems, however, the photolithographic resolution offered by EUV exposure systems is not required and the development of affordable, short wavelength processes and exposure tools is desirable.

We report here the realization of a scalable, precise, and rapid photolithographic process based on the photoablation of a wide range of polymer films with incoherent, vacuum ultra-
violet (VUV) radiation provided by newly-developed flat lamps. Specifically, the observed photoablation of a wide range of polymers, including acrylics and polycarbonates, at 172 nm ($h\nu = 7.2\ eV$) in the VUV spectral region with material removal rates up to 4 nm s$^{-1}$ for lamp intensities as low as 10 mW cm$^{-2}$ provides the foundation for a group of contact or projection photolithographic processes, operating in one atmosphere of nitrogen and requiring no homogenization or alteration of the lamp intensity spatial profile. Complex nanostructures have been fabricated by contact lithography in polymer films and bulk material with one or more exposures and overall exposure times of typically 10–300 s, and the depths of trenches and other features are routinely controlled to a precision of 10 nm. Currently available photomasks have been fabricated by electron beam writing but the lateral resolution (minimum feature size) is presently limited to ~350 nm. With fused silica masks of higher quality, sub-200 nm features are expected to be accessible. Consequently, the demonstrated resolution is superior to that of commercial i-line tools typically offering line-and-space resolutions of ~0.8–2 μm. The pattern resolution offered at present by polymer photoablation at 172 nm (~350 nm) is $\lambda /4$ at optical communications wavelengths (1450–1650 nm), and is $\lambda /2$ when considering the deep-red and near-infrared spectral regions. Therefore, the photolithographic capability described here presents a compact, inexpensive technology for the fabrication of Bragg gratings as well as other optical components and structures for fiber and semiconductor lasers operating in the 700–1600 nm interval. Regardless of the application, adopting 172 nm lamps as the exposure source allows for the elimination of wavelength-specific, photoactive molecules, such as diazonaphthoquinone (DNQ), in organic photoresists because the absorption of 7.2 eV photons modifies the polymer resin directly by bond scission in the primary and side chains. Consequently, one may dispense with photosensitizers and polymer resins of large molecular weight, while substituting alcohols and ketones (isopropanol and acetone) for developers of higher toxicity. The single-component photoresist chemistry reported here lowers significantly the cost and complexity of the VUV exposure tool while reducing the volume and toxicity of the chemical waste stream, in comparison with conventional photosist processing.

In an effort to benchmark the characteristics and potential of this photolithographic process, several discrete optical components and arrays have been fabricated and tested. These include large arrays of Al nanomirrors, each of which is mounted atop a 350 nm–1 μm diameter Si pillar, as well as Fresnel lenses and both reflection and transmission optical gratings. Fresnel lenses providing intensity concentration factors of 380 for a 50 mm focal length (at 632.8 nm), and transmission gratings exhibiting 40% efficiency for the diffraction of incident power into first order, have been realized because of the precision of the 172 nm photoablation of polymer films. Plasmonic arrays have also been fabricated by electro-deposition of gold into patterns defined by this VUV photolithographic process.

It has been known since the early 1980s that certain organics, as well as animal and human tissue, are non-thermally photoablated when exposed to VUV laser radiation. However, although precision machining of human hair was reported in ref. 11, and Trokel et al. demonstrated non-thermal ablation of cow corneas in 1983, for example, these and subsequent studies were conducted at 193 nm with pulsed ArF lasers generating peak intensities of ~10 MW cm$^{-2}$. One unexpected aspect of the results reported here is that the photoablation of polymers at 172 nm with pulsed lamps occurs despite peak radiated powers of only a few hundred watts, at least four orders of magnitude lower than those characteristic of the 193 nm laser.

The potential for photolithography in the 150–200 nm segment of the VUV spectral region was recognized 2–3 decades ago but the leading candidate for the optical source – the F$_2$ laser at 157 nm – later proved to be unsuitable because of its multiline spectral output and complex optical requirements. Motivated by quite different applications, several biomedical groups subsequently reported low resolution (50 μm) photopatterning of polydimethylsiloxane (PDMS) with a Xe$_2$ (172 nm) lamp and wire mesh serving as a mask. These efforts were directed primarily to the development of processes relevant to patterning cells, immobilizing biomolecules, and tissue engineering. It was also in this context that Hozumi et al. reported the spatially-selective hydrophobilization of poly-methyl methacrylate (PMMA) through the interaction of the surface with excited molecules and radicals produced in air by the 172 nm lamp. In 2006, however, Asakura et al. reported a photolithographic process in which a low-intensity, 172 nm lamp irradiated PMMA films immersed in room air at various pressures. Both positive and negative-tone lithography were demonstrated, but material removal was dependent upon the interaction of photochemically-generated ozone with the polymer. As in the experiments reported in ref. 16, the presence of atmospheric air was required for the photetching process to proceed. In addition to these experiments, studies of photoresist pattern freezing, the cleaning of mirror surfaces by the photochemical removal of organics, and extensive investigations of the photochemical effects of 172 nm radiation on organic films have been reported.

A transformation in the VUV source landscape occurred in 2017 when Park et al. announced the realization of flat, thin lamps emitting intensities at 172 nm up to, and beyond, 300 mW cm$^{-2}$ with peak emission powers of 600–800 W in a bandwidth of 9 nm. Lamps having an active surface area of ~100 cm$^2$ are now capable of generating average powers at 172 nm exceeding 25 W. To put such unprecedented VUV lamp average powers and peak intensities into perspective, we note that they are more than five orders of magnitude larger than solar exposure in this wavelength region (outside earth’s atmosphere and assuming the same bandwidth). Furthermore, the star in the Cygnus constellation known as 2 Cygni is one of the few that emits a blackbody spectrum peaking at 175 nm, which corresponds to a “color” (blackbody) temperature of 16 500 ± 200 K. Because such intensities are
also considerably larger than those commercially-available previously, the electrical ("wall plug") efficiency is >20%, and the lamp cost-of-ownership is at least 2–3 orders of magnitude lower than the corresponding value for a 193 nm excimer laser, it is now feasible to pursue broadly the photochemical processing of materials because the precipitous decline in the cost of a mole (i.e., Avogadro’s number) of VUV photons suggests that commercial applications are now feasible. Of equal significance is the potential for returning a fraction of the cost of a mole (i.e., Avogadro’s number) of VUV photons, which are useful for high-speed processing of materials because the precipitous decline in the cost of a mole (i.e., Avogadro’s number) of VUV photons suggests that commercial applications are now feasible.

2. Experimental procedures and results

All of the experiments reported here were conducted with the 172 nm contact mask lithography system (π−Cygni) manufactured by Cygnus Photonics. Incorporating a 50 mm × 50 mm lamp situated 2 cm above the substrate, this exposure tool produces measured 172 nm intensities of 10 ± 2 mW cm⁻² at the desired surface (typically either a Si wafer or a sample of commercial-grade or optical-grade polymer) and the emission intensity over the active area of the lamp’s output window is uniform to within 3%. The intensity of the lamp was monitored periodically by a calibrated energy detector, designed by Hamamatsu specifically for the 172 ± 15 nm spectral region. For those tests involving a Si substrate, the 50 mm diameter wafer was first cleaned by the standard rinse sequence of acetone, isopropyl alcohol (IPA), and deionized (DI) water which was followed by spin-coating a 40–200 nm thick polymer film onto the wafer. A fused silica photomask having the desired geometric pattern (but no proximity corrections or other resolution-enhancing features) was placed onto the substrate, and research-grade nitrogen was flowed through the lamp situated 2 cm above the substrate, this exposure tool produces exposure times (typically 5–10%) at a pulse repetition frequency of tens of kHz is 3–4 orders of magnitude higher than the corresponding value for an ArF laser operating at 100 Hz–1 kHz, exposure times can be maintained at or below tens of seconds while lowering the cost of the tool.

As discussed later, the rinsing of deep structures photoablated into polymer surfaces was found to not degrade the spatial resolution of the process, but rather improve it. It must also be emphasized that this photolithographic process required no other liquid processing, such as that normally required for conventional photoresists. In addition to polymer film-coated substrates, extensive testing of a wide variety of other organic materials and polymers, including ABS, polycarbonate, and allyl diglycol carbonate (ADC, also known as CR39) have been examined. Each of these was found to yield precisely-photopatterned structures in the polymer, regardless of whether the surface was of optical quality or not. For several experiments, gold was electro-deposited or aluminum was sputtered into windows opened in a thin polymer film by 172 nm photoablation.

Measurements of the photoablation rate for bulk PMMA for exposure times (t) up to ~900 s are summarized in Fig. 1. For these experiments, ~3 mm-thick PMMA sheets (non-optical grade) divided into ~76 mm × 76 mm squares served as the substrates, and the protective paper backing was removed immediately prior to inserting the samples into the VUV source/mask system. After a single exposure to the lamp with the intensity at the substrate surface fixed at 10 mW cm⁻², the samples were rinsed with IPA for 2 s and deionized water for 2 s, and this rinsing procedure was repeated for 5 cycles. Literally scores of measurements yielded results showing surprising reproducibility, within 10 nm and limited by the instrument (optical profilometer) resolution, for a given lamp intensity and sample exposure time. Therefore, the uncertainty in the measurements of Fig. 1 is smaller than the diameter of the symbols. Note from Fig. 1 that the initial, linearized photoablation rate is ~4 nm s⁻¹, but above t ~ 70 s the onset of saturation appears. For exposure times beyond ~300 s, the polymer removal rate is constant at ~0.2 nm s⁻¹. The upper abscissa in Fig. 1 shows that the fabrication of trenches 300 nm in depth in PMMA requires a cumulative exposure dosage of ~1 J cm⁻². Extensive observations of the wall and floor profiles of ablated trenches or cylinders demonstrate that the slopes of the vertical side walls are in the range of 25–35°.
Fig. 1 Dependence of the depth of trenches, photoablated in bulk acrylic sheets, on the 172 nm exposure time. All of these data were recorded for PMMA as the substrate, and the lamp intensity at the substrate fixed at 10 mW cm$^{-2}$. Also, each depth measurement was made after an IPA rinse of the photoablated surface. The upper abscissa shows the cumulative exposure dosage, expressed in J cm$^{-2}$, and the uncertainty in each measurement is smaller than the diameter of the data symbols (open circles).

Fig. 2 Optical micrograph of a panoramic view of a standard (U.S. Air Force) resolution pattern, reproduced in a 3 mm thick, bulk PMMA sheet by exposing the substrate to 172 nm radiation through a fused silica mask. This image required an exposure time of 60 s and 5 cycles of a 2 s IPA rinse, followed by a 2 s DI rinse. Note the presence of small bubbles located beneath the surface of this commercial-grade polymer sample.

(relative to a vertical plane orthogonal to the substrate surface) and are dictated by several factors: the flatness of the surface, the mask-sample contact, and the collimation of the incoming radiation. As noted earlier, the removal of the particulate debris with a post-exposure rinse does not degrade the resolution of the process. Specifically, direct photoablation accounts for only 25% of the final trench depth. The particulates observed in photoexposed trenches, channels, and cylinders are clearly the result of VUV photochemical modification of the polymer, and its subsequent degradation into hydrocarbon chains of lower molecular weight. We attribute this sequence of events generated by the photolysis of various polymers and detected by a residual gas analyzer (RGA) are in progress, but initial results indicate that the cleavage of both the C–C and C–O bonds of the ester carbonyl in the side chains is also playing a significant role.Regardless of the specific processes responsible for fragmenting PMMA and other polymers studied in these experiments, profilometry measurements demonstrate that the wall slopes of photopatterned trenches, for example, remain essentially unchanged in the portion of the trench occupied by debris (i.e., prior to rinsing). That is, following an exposure, debris occupies 75% of the trench volume (regardless of exposure duration), but the trench cross-section decreases monotonically and continuously with depth. Analysis of trench width measurements and wall profiles found that the maximum achievable resolution (a) is given by 

$$d = \frac{w}{2.5 \times d},$$

where $d$ is the depth of a feature. This is equivalent to stating that the maximum aspect ratio achievable at present is 0.4. Consequently, 700 nm deep trenches having lateral feature sizes of $\sim 1.75 \mu m$ are fabricated readily with the processes reported here. Shallower features, such as $\sim 150$ nm-deep trenches, allow for a lateral spatial resolution of $\sim 370$ nm. These comments are valid only for commercial-grade polymer samples, and it must be emphasized that the resolution achieved despite the poor optical quality of the substrates is surprising.

Fig. 2 is an optical microscope image of a representative, low resolution pattern photoablated into bulk PMMA. This U.S. Air Force pattern has observable features as small as 5 μm, and we note that several bubbles are visible within the commercial-grade polymer slab. It should be mentioned that the floor of the channels and other features precisely reproduce the original surface of the photomachined substrate. A series of SEMs of several patterns and specific features fabricated in a 55 nm – thick PMMA film spin coated onto a Si substrate is given in Fig. 3. In panels (a)–(c) of the figure, the lighter regions represent channels photoablated into the polymer, and Fig. 3(c) is a magnified view of a line having a width of $\sim 1.2 \mu m$. The exposure time required for each of these three patterns was 40 s. Panel (d) of Fig. 3 demonstrates the capability for fabricating both bright and dark field patterns (lower and upper portions of Fig. 3(d), respectively).

Experiments conducted over the past two years with a wide range of polymers, including PMMA, ABS, and CR-39, have demonstrated consistently that sub-400 nm features in the...
Fused silica photomasks are replicated in both bulk polymers and thin films on Si, and the lateral spatial resolution (∼350 nm) is limited primarily by the photomasks available at present. As an example, Fig. 4 is an SEM image of one tip of a circular tip array pattern such as that shown in Fig. 3(b). Surface topographical maps of other patterns photoablated into PMMA by a single exposure are shown in Fig. 5 and 6. Recorded by a 3D laser scanning confocal profilometer, the image of Fig. 5 is that of a pattern of ∼3 μm-wide trenches having depths of 500 nm. The dashed black line in the upper portion of the figure indicates the path along which the lineout (bottom half of the figure) was recorded. The cusping (narrowing) of linewidths evident at right angles and the intersection at the center of the image are intentional, and were designed into the photomask so as to illustrate one aspect of the achievable resolution. Fig. 6 is a laser confocal microscope image of a “Space Invaders” pattern having a depth of 240 nm, and fabricated by a 60 s exposure. Complex 3D patterns and structures can be fabricated quickly and precisely into the surface of a variety of polymers through multiple exposures of the substrate, and Fig. 7 provides several images illustrating multilayer patterns produced to date in bulk acrylic substrates.
Fig. 7 False color images of several examples of nanostructures and geometries readily obtainable through multiple exposures of a polymer film with the same or different photomasks: (a) a simple planar structure fabricated in PMMA by a double exposure. Color coding distinguishes three separate layers in the structure, each of which is 240 nm in height; (b) 3D structure exhibiting periodicity in two-dimensions and fabricated by two exposures and rotation of the mask by 90°; (c) “Spider’s Web” pattern fabricated by a double exposure and translation of the mask in a plane parallel to the substrate surface. The radial and azimuthal symmetry of the pattern is evident; (d) a four-level pattern in which various structures (mesas, vertically-oriented plates, etc.) were formed intentionally by a triple exposure. All exposures were one minute in duration, and the field-of-view for this image is 97 μm × 73 μm. In all of these examples, the exposures were 1 minute in duration, and the field-of-view for the images of panels (a) and (b) is 97 μm × 73 μm, whereas the corresponding value for Fig. 7(d) is 280 μm × 212 μm.

In fabricating the pattern of Fig. 7(c), a different procedure was adopted. In this case, the substrate was exposed twice with different photomasks. The first of these has a circular (azimuthally-symmetric) pattern similar to that of Fig. 3(b) with equally-spaced triangular points directed radially inwards. The second mask was designed for fabricating diffraction gratings and comprises a one-dimensional array of 5 μm lines and spaces. After the process was completed, the substrate was rinsed in IPA and DI. The result is a “Spider’s Web” array which is equivalent to a diffraction grating having a variable periodicity (i.e., a chirped grating). Deferring the rinsing of the substrate until after the exposures are completed has the effect of rounding and smoothing the array features, owing to the loss of spatial resolution arising from a reduction in the steepness of all vertical walls. Rounding sharp features, such as corners, is expected to be of value in microfluidics, for example, by reducing the liquid flow velocity gradient at channel intersections.

The precision of the 172 nm photoablation process opens the door to the fabrication of optical and microfluidic components and networks. Optics is of particular interest because of the potential for all-polymer networks in which polymer optical fiber, lenses, gratings, and waveguides are integrated. Fig. 8 summarizes measurements of the diffraction efficiency for the PMMA grating (mentioned earlier) having 5 μm wide lines (trenches) and spaces. Recorded for an incident laser wavelength of 632.8 nm, these data confirm that, with increasing exposure time (i.e., trench ablation depth), diffracted power is gradually transferred from the central order (m = 0) into higher grating orders (m = ±1, ±2, and ±3). Considering the first-order data, one notes that the diffracted power is −4 dBm when the trench depth is ~600 nm (Fig. 1). This value corresponds to a diffraction efficiency for the conversion of incident power into first-order of ~40%. An optical micrograph
Fig. 8 Diffraction efficiency for a transmission grating fabricated on a nonoptical-grade PMMA flat. Measurements of the optical power diffracted into several grating orders (central order \( m = 0 \) and \( m = \pm 1, \pm 2 \) and \( \pm 3 \)) are shown for 172 nm exposure times ranging from 60 s to 600 s. Solid and non-solid symbols represent positive and negative order signs, respectively. These were recorded at a wavelength of 632.8 nm with a He–Ne laser operating within the fundamental transverse mode, and were acquired by measuring the grating characteristics after each exposure to the lamp. Note that the power diffracted into first-order (\( \pm 1 \)) surpasses that for the central order when the exposure time reaches 600 s. The efficiency for diffracting power into 1st order is \( \sim 40\% \).

Fig. 9 (a) Optical micrograph (in plan view) of a Fresnel lens fabricated on a non-optical grade acrylic by a 3 minute exposure, and (b) laser confocal profilometer image of a section of the lens. The overall diameter of the lens structure is 1 mm, its focal length is 50 mm, and the wall roughness evident in the Fresnel rings is entirely due to the resolution of the photomask. The depth of the circular trenches is 390 nm, and this lens produces a 28 \( \mu m \) (FWHM) focal spot when illuminated by a 632.8 nm laser (He–Ne) having a beam diameter of 1.0 mm.

Combining 172 nm photolithography with well-developed semiconductor device fabrication processes allows for arrays such as those shown in Fig. 10 to be realized. Circular or square silicon posts were fabricated by first spin-coating a 100 nm Si(100) wafer with Microchem 950 PMMA A2 photoresist which has a molecular weight of 950 000. The thickness of the film was measured to be \( 55 \pm 3 \) nm, and was subsequently hard-baked at 150 °C for 60 s. Immediately thereafter, the PMMA film was exposed through a photomask with 172 nm intensities of either 10 mW cm\(^{-2}\) or 70 mW cm\(^{-2}\). Exposure times of 10–40 seconds and 4 cycles of rinsing the wafer in IPA and DI (for 2 s each) opened windows in the PMMA film, into which aluminum was sputtered. The remaining PMMA film was then removed by lift-off, and the Al pillars of Fig. 10 were produced by deep reactive ion etching (DRIE) of the substrate for 10 minutes. Using the Al discs (or squares) as a mask, Si posts 11 \( \mu m \) in length were etched. The top two panels of Fig. 10 are SEMs of portions of a large array of 1 \( \mu m \) diameter Si pillars, and atop each of these is an Al mirror. Arrays comprising Si posts of smaller diameter were also fabricated, and the lower portion of Fig. 10 shows SEMs of 350 nm – diameter pillars. At lower right, the periodic variations in the diameter of the post, characteristic of the Bosch etching process, are evident as is an out-of-plane annulus at the perimeter of the aluminum mirrors that is a result of the lift-off process. Undercutting of the Al discs by the etching process leads to cusping of each mirror.

In an effort to demonstrate the applicability of 172 nm photoablation of polymers to other semiconductor device fabrication processes, selective electro-deposition of gold onto exposed areas of a Si substrate (i.e., windows opened by photolithography) has also been realized. Prior to placing a Si wafer in the lamp chamber, the native oxide was immersed in a
0.8% HF solution for 20 seconds. After exposure, gold was electro-deposited onto the wafer at a temperature of 60 °C and a DC current of 10 mA. Results representative of those obtained in these experiments are presented in Fig. 11 by an SEM of gold lines, 2 μm in width, which were electro-deposited through windows opened in the PMMA photoresist. Notice that the edges are sharp, and the variation in the width of the gold lines was determined to be 150 nm. Gold is also of considerable interest for plasmonic devices and, to that end, arrays of gold dots and discs have been electro-deposited onto Si by the process described earlier. A final example of the versatility and resolution of the lithographic processes reported here is the fabrication of arrays of aluminum discs. Fig. 12 shows electron micrographs of segments of two such arrays, fabricated on Si(100) by sputtering Al through windows opened in a 55 nm-thick PMMA film by 172 nm photoablation. Panel (a) of the figure is an SEM of a portion of an array comprising 1 μm squares, whereas Fig. 12(b) is a magnified image of nine, ∼500 nm diameter discs from another array. A similar fabrication process was adopted for a preliminary examination of the dependence of feature resolution on VUV exposure of the polymer. Specifically, 55 nm thick films of Microchem 950 PMMA A2 resist were spun onto Si and baked at 150 °C for 60 seconds. After exposing these films with a VUV dosage of ∼400 mJ through an 8 mm-thick mask having no proximity corrections and rinsing with IPA and water as described earlier, Al was sputtered through square windows opened in the film. Afterwards, the remaining polymer resist was removed by lift-off. With this process sequence, the radius of curvature of the corners of 1 μm Al squares was observed to be ∼240 nm. When the VUV dosage is increased by 50% (to ∼600 mJ), the radius of curvature rose to 350 nm and the Al squares increased in surface area to 1.1 × 1.1 μm². It must, however, be emphasized that these results are tentative and this process, as well as all those presented earlier, have yet to be optimized.

3. Conclusions

A photolithographic process based upon flat 172 nm lamps and one of several polymers acting as a photoresist has been demonstrated. Data presented here show the resolution of 172 nm photolithography to be superior to that of Hg i-line lithography, at a cost orders of magnitude lower than that characteristic of 193 nm exposure tools. Intricate, multilevel
structures and patterns are formed on bulk acrylates such as PMMA, ABS, or CR-39 or in a polymer film spin-coated onto a semiconductor substrate. Photoablation of polymers at 172 nm produces nanoscale features of precise dimensions, and the maximum resolution (minimum lateral feature size) has been demonstrated to lie below ~350 nm. It is anticipated that feature sizes below 200 nm can be patterned with the appropriate photomasks and attention given to optimizing the polymer film thickness and other parameters of the exposure and rinsing sequence. The speed and precision of the processes reported here suggest that hybrid VUV/e-beam lithography patterning, both of which accommodate PMMA photoresists, will enable the rapid patterning of nanoscale features on the same wafer without the need to strip the polymer resist when transitioning between the VUV and e-beam exposure steps.\(^{23,26}\) It is our expectation that such processes will be of value in rapidly and inexpensively fabricating networks and patterns for optical wave front manipulation, micro- and nano-fluidics, electronic and optical components and networks, and biomedicine and pharmacology. Not only is 172 nm photolithography appealing for advancing research in micro- and nanotechnology and physics but the affordability, precision, and speed of the exposure process expands access of academic and industrial laboratories to a lithography capable of <200–400 nm feature fabrication.

Data availability

The data that support the findings of this study are available from the corresponding author upon request.

Conflicts of interest

The authors declare no conflicts.

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References

1. R. P. Seisyan, *Tech. Phys.*, 2011, **56**, 1061.
2. S.-S. Kim, R. Chalykh, H. Kim, S. Lee, C. Park, M. Hwang, J.-O. Park, J. Park, H. Kim, J. Jeon, I. Kim, D. Lee, J. Na, J. Kim, S. Lee, H. Kim and S.-W. Nam, *Extreme Ultraviolet (EUV) Lithography VIII*, 2017, pp. 1–10.
3. R. A. Hughes, E. Menumerov and S. Neretina, *Nanotechnology*, 2017, **28**, 282002.
4. K. Jain, C. G. Willson and B. J. Lin, *IEEE Electron Device Lett.*, 1982, **3**, 53–55.
5. P. Alagna, O. Zurita, V. Timoshkov, P. Wong, G. Rechtsteiner, J. Baselmans and J. Malfert, *Optical Microlithography XXVIII*, 2015, pp. 51–63.
6. S. Weichselbaum, F. Bornbroek, T. de Kort, R. Droste, R. F. de Graaf, R. van Ballegoij, H. Botter, M. G. McLaren and W. P. de Boeij, *Optical Microlithography XXVII*, 2015, pp. 277–287.
7. S.-J. Park, C. M. Herring, A. E. Mironov, J. H. Cho and J. G. Eden, *APL Photonics*, 2017, **2**, 041302.
8. Y. Mimura, T. Ohkubo, T. Takeuchi and K. Sekikawa, *Jpn. J. Appl. Phys.*, 1978, **17**, 541–550.
9. R. Srinivasan and V. Mayne-Banton, *Appl. Phys. Lett.*, 1982, **41**, 576–578.
10. R. J. Lane, R. Linsker, J. J. Wynne, A. Torres and R. G. Geronemus, *Arch. Dermatol.*, 1985, **121**, 609–617.
11. M. Witt, *Hyperfine Interact.*, 1987, **37**, 415–421.
12. S. L. Trokel, R. Srinivasan and B. Braren, *Am. J. Ophthalmol.*, 1983, **96**, 710–715.
13. H. G. Craighead, J. C. White, R. E. Howard, L. D. Jackel, R. E. Behringer, J. E. Sweeney and R. W. Epworth, *J. Vac. Sci. Technol., B: Microelectron. Process. Phenom.*, 1983, **1**, 1186–1189.
14. D. Henderson, J. C. White, H. G. Craighead and I. Adesida, *Appl. Phys. Lett.*, 1985, **46**, 900–902.
15. J. Gan, H. Chen, F. Zhou, H. Huang, J. Zheng, W. Song, L. Yuan and Z. Wu, *Colloids Surf., B*, 2010, **76**, 381–385.
16. A. Hozumi, T. Sai\(\text{\'}o\), N. Shirahata, Y. Yokogawa and T. Kameyama, *J. Vac. Sci. Technol., A*, 2004, **22**, 1836–1841.
17. S. Asakura, A. Hozumi, T. Yamaguchi and A. Fuwa, *Thin Solid Films*, 2006, **500**, 237–240.
18. M. Yamaguchi, T. Wallow, Y. Yamada, R.-H. Kim, J. Kye and H. J. Levinson, *J. Photopolym. Sci. Technol.*, 2008, **21**, 697–704.
19. K. Hamamoto, Y. Tanaka, T. Watanabe, N. Sakaya, M. Hosoya, T. Shoki, H. Hada, N. Hishinuma, H. Sugahara and H. Kinoshita, *J. Vac. Sci. Technol., B: Microelectron. Nanometer Struct.–Process., Meas., Phenom.*, 2005, **23**, 247–251.
20. F.-E. Truica-Marasescu and M. R. Wertheimer, *Macromol. Chem. Phys.*, 2005, **206**, 744–757.
21. *Solar Constant and Zero Air Mass Solar Spectral Irradiance Tables*, 2006, https://www.astm.org/DATABASE.CART/HISTORICAL/E490-00AR06.htm.
22. H. Okabe, *Photochemistry of small molecules*, Wiley, New York, 1978.
23. O. Sonoiki, Ph.D. thesis, University of Illinois, 2019.
24. M. Rezem, A. Gunther, B. Roth, E. Reithmeier and M. Rahlves, *J. Lightwave Technol.*, 2017, **35**, 299–308.
25. M. Gersborg-Hansen, L. Thamdrup, A. Mironov and A. Kristensen, *Microelectron. Eng.*, 2007, **84**, 1058–1061.
26. D. J. Carbaugh, S. G. Pandya, J. T. Wright, S. Kaya and F. Rahman, *Nanotechnology*, 2017, **28**, 455301.
27. G. Calafiore, A. Koshelev, F. I. Allen, S. Dhuey, S. Sassolini, E. Wong, P. Lum, K. Munecika and S. Cabrini, *Nanotechnology*, 2016, **27**, 375301.