Ultraviolet laser-induced poling inhibition produces bulk domains in MgO-doped lithium niobate crystals

Andreas Boes, Hendrik Steigerwald, Didit Yudistira, Vijay Sivan, Scott Wade, Sakellaris Mailis, Elisabeth Soergel, and Arnan Mitchell

Citation: Applied Physics Letters 105, 092904 (2014); doi: 10.1063/1.4895387
View online: http://dx.doi.org/10.1063/1.4895387
View Table of Contents: http://scitation.aip.org/content/aip/journal/apl/105/9?ver=pdfcov
Published by the AIP Publishing

Articles you may be interested in
Polarization reversal induced by heating-cooling cycles in MgO doped lithium niobate crystals

Precision nanoscale domain engineering of lithium niobate via UV laser induced inhibition of poling

Annihilation of the OH absorption due to domain inversion in MgO-doped lithium niobate crystals

Light-induced order-of-magnitude decrease in the electric field for domain nucleation in MgO-doped lithium niobate crystals
Appl. Phys. Lett. 86, 212901 (2005); 10.1063/1.1929099

Second-harmonic generation of green light in periodically poled stoichiometric Li Ta O 3 doped with MgO
J. Appl. Phys. 96, 7445 (2004); 10.1063/1.1804616
Ultraviolet laser-induced poling inhibition produces bulk domains in MgO-doped lithium niobate crystals

Andreas Boes,1,2,a) Hendrik Steigerwald,1,2 Didit Yudistira,1 Vijay Sivan,1,2 Scott Wade,3 Sakellaris Mailis,4 Elisabeth Soergel,5 and Arnan Mitchell1,2
1School of Electrical and Computer Engineering, RMIT University, Melbourne, Victoria 3001, Australia
2ARC Center for Ultra-high Bandwidth Devices for Optical Systems (CUDOS), RMIT University, Melbourne, Victoria 3001, Australia
3Faculty of Science, Engineering and Technology, Swinburne University of Technology, Hawthorn, Victoria 3122, Australia
4Optoelectronics Research Centre, University of Southampton, Highfield, Southampton SO17 1BJ, United Kingdom
5Institute of Physics, University of Bonn, Wegelerstr. 8, 53115 Bonn, Germany

(Received 23 June 2014; accepted 29 August 2014; published online 5 September 2014)

We report the realization of high-resolution bulk domains achieved using a shallow, structured, domain inverted surface template obtained by UV laser-induced poling inhibition in MgO-doped lithium niobate. The quality of the obtained bulk domains is compared to those of the template and their application for second harmonic generation is demonstrated. The present method enables domain structures with a period length as small as 3 μm to be achieved. Furthermore, we propose a potential physical mechanism that leads to the transformation of the surface template into bulk domains. © 2014 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4895387]

Lithium niobate (LiNbO3) crystals have been widely used in many optical applications, in particular nonlinear optics, due to their excellent nonlinear optical properties.1 Nonlinear optical processes such as frequency conversion are based on quasi-phase matching, which requires periodic domain inversion of the crystal’s spontaneous polarization P_s at sub-micron to few-micron periods depending on the wavelength involved in the nonlinear process.2 As is the case for many nonlinear processes, frequency conversion also requires high intensity laser light, which may induce optical damage to the crystal that results in the distortion of the optical beam. It is known that adequate magnesium oxide (MgO) doping in LiNbO3 can significantly reduce the optical damage.3 However, while it is possible to find crystals suitable for high optical intensity applications, fine period domain inversion of MgO-doped crystals is known to be particularly challenging due to random domain growth and high leakage currents.4

A common method used for domain patterning of LiNbO3 is electric field poling (EFP),5 where a spatially modulated electric field is applied along the polar axis of the crystal. Domain inversion occurs in areas where the local electric field exceeds the coercive field (E_c). While this method has been shown as an effective way to achieve domain patterning in congruent LiNbO3, it is not always successful when applied to MgO-doped LiNbO3, due to a pronounced sideways domain spreading which occurs during EFP limiting the smallest achievable domain period.4 However, the ability to realize short domain periods on such crystals can widen the scope of its applications, as they are required for the generation of counter propagating photons in nonlinear optical applications.5

Recently, it has been shown that a two-step voltage application during UV illumination can produce good quality domain patterns in MgO-doped LiNbO3 with a period of 19 μm.6 This technique first generates surface domains by applying an electric field, which is smaller than the E_c while simultaneously illuminating the crystal with weakly absorbed UV light. The weakly absorbed UV light reduces the E_c,9 therefore allowing the nucleation of surface domains. By applying a second EFP step without UV light illumination, the surface domains behave as a template, and bulk domains are created. The major drawback of using this technique, especially when realizing small domains, is the need to simultaneously irradiate the crystal with UV light during EFP. Especially, as the experimental setup required for EFP can potentially limit the focal spot size of the UV laser light on the crystal surface and thus affecting the attainable minimum period lengths of the domain pattern. Separating the two steps (UV illumination and electric field application) could overcome this limitation.

In this paper, we report on the fabrication of bulk domain structures in MgO-doped LiNbO3, which are pre-defined by a spatial modulation of the lithium (Li) concentration induced by strongly absorbed UV laser light irradiation. The bulk domain distribution is subsequently produced by an even number of EFP steps. In this way, the laser irradiation and EFP steps can be separated removing the restrictions imposed in Refs. 7 and 8. We report the fabrication of periodic bulk domain structures with a period as short as 3 μm.

The methodology for generating and visualizing the domains is presented below. In our experiments, we used 500-μm-thick 5 mol. % of MgO doped Z-cut congruent LiNbO3 crystals (MgCLN), which were provided by “Yamaju Ceramics co. Ltd.” Focused UV laser light from a frequency doubled argon ion laser (λ = 244 nm) was scanned along the crystallographic Y direction on the +Z-face of the crystal. The laser beam was focused using a fused silica lens (f = 40 mm) to a focal beam diameter of ~6 μm and an intensity of ~3 × 10^5 W/cm^2. The laser beam was scanned to irradiate linear track sections with a period of 18.8 μm at a velocity of 0.5 mm/s.

---

a)Electronic mail: s3363819@student.rmit.edu.au
Local irradiation of the +Z-face with strongly absorbed UV laser light results in a Li deficiency enabling surface domain formation via poling inhibition (PI) during a subsequent EFP step. For the EFP process, we used a setup similar to the one introduced by Sones et al., which allowed us to visualize the domain growth during the poling process. The voltage applied to the crystal was ramped up at a rate of 5 V/(mm-s) until a non-ohmic behavior of the measured current was observed indicating domain reversal. After the completion of the poling process, the applied voltage was ramped down at a rate of 100 V/(mm-s). To avoid unintended domain inversion on specific parts of the crystal, areas of the crystal surface were covered with Kapton tape after each poling step. This step of process allowed us to study the domain formation after each poling step, namely, after the first forward poling, after the first reverse poling (completing the first poling cycle), after the second forward poling and after the second reverse poling (completing the second poling cycle). The resulting domain patterns were investigated using scanning electron microscopy (SEM) and optical microscopy imaging after domain selective etching of the polished crystal cross-sections (Y-face) using hydrofluoric acid (HF).

An investigation to explore the smallest achievable domain period with this method was also conducted. For this purpose, we used a fused silica lens with a shorter focal length ($f = 10 \text{ mm}$) to achieve a focal beam diameter of $\sim 2 \text{ \mu m}$. The focused laser beam was then scanned in periodic track patterns (similar to above) with periods of 4 \mu m and 3 \mu m at a velocity of 0.5 mm/s. As previously described, the same methodology was adopted for the generation and visualization of domains.

The quality and thermal stability (at working temperatures) of the periodic domain structure after the application of two poling cycles was investigated by performing a second harmonic generation (SHG) experiment. A beam from a tunable infrared laser was focused into a heated, periodically poled crystal ($L = 9.4 \text{ mm}$, $A = 18.8 \mu \text{m}$) and the SHG power was detected as a function of the fundamental wavelength. The SHG power was monitored over a time period of 2 h at crystal temperatures of 100, 150, 200, and 260°C to provide an in-situ measure of the thermal stability of the domains.

The results of the UV irradiation and subsequent EFP are shown in Fig. 1. Fig. 1(a) shows the SEM image of the crystal after the first forward poling step, where a surface domain can be recognized from the ridge that is formed on the Z-face of the crystal and the differentially etched pattern of the Y-face cross-section. This shows that the polarization of the whole crystal is reversed except a shallow area underneath the UV-irradiated track, which maintains the original polarization of the crystal (PI domain). The Y-face cross-section also reveals that the tail-to-tail boundary of the PI domain is very irregular (spiky). The measured width of the surface domain is $\sim 5 \mu \text{m}$, which is comparable to the focal diameter of the irradiating UV-laser beam. The measured depth of the PI domain is $\sim 1.5 \mu \text{m}$. Higher UV laser light intensities were found to generate both, wider and deeper PI domains, which corresponds to the findings in Refs. 13 and 14. After a subsequent reverse poling step, the volume underneath the PI domain maintains its polarization while domain reversal occurs everywhere else as shown in Fig. 1(b). As a result, a well-defined bulk domain is formed below the PI domain. From the visualization of the domain growth during reverse poling, which is not shown here, we observed that the domain nucleation started within the UV irradiated area.

The domain depth is qualitatively analyzed on the cross-section using optical microscope imaging after HF treatment. The images of the sample taken after the first and the second poling cycle are shown in Figs. 2(a) and 2(b), respectively. The needle-like periodic structures in Fig. 2 correspond to the UV laser irradiation as shown previously in Fig. 1(b). We find that the domain width becomes narrower as the crystal depth is increased and they do not terminate the opposite Z-face of the crystal, which is unlike the poling process with standard EFP. As shown in Fig. 2(a), we also find that after the first poling cycle, the depth of the individual domains varies dramatically and their alignment does not nicely follow the crystallographic Z-axis. This however is improved after applying the second poling cycle, as shown in Fig. 2(b). Much deeper domains are achieved, which, in addition, are now more uniformly aligned along the Z-axis. The improvement of the domain uniformity in depth and parallelism after the second poling cycle can be attributed to a smoother and more regular domain wall motion, achieved when poling the crystal multiple times. However, we observed that applying further poling cycles did not improve the quality of the domain in terms of the domain depth. The importance of the domain wall motion can be further supported by fact that this domain engineering method was not suitable for congruent LiNbO$_3$, where the domain wall motion is faster compared to MgO-doped LiNbO$_3$.16

![Fig. 1. SEM images of the sample crossed-sections after being treated with HF, taken after (a) forward poling and (b) reverse poling of the crystal.](image1)

![Fig. 2. Optical microscope images of the HF-etched cross-sections of the crystal after (a) the first poling cycle and (b) the second poling cycle. The periodic, needle-like structure are the inverted domains (correspond to the bulk domain in Fig. 1(b), which start at the +Z face and become thinner as they grow in depth. The domain depth uniformity improves after the second poling cycle.](image2)
In Figs. 3(a) and 3(b), we show fabricated samples with smaller domain periods, namely, 4 μm and 3 μm, achieved using a lens with shorter focal length. We can see that the domain periods match nicely with the scanning periods of the beam and that the left domain in Fig. 3(a) is smaller than the other domains in the SEM images. These results demonstrate that the present technique is also applicable to obtain short poling periods. However, good uniformity of the periodic domain structure over large areas was found to be difficult to achieve, which requires further investigation but it is beyond the scope of this paper. A smoother and slower domain wall motion, which could be beneficial for reaching this goal, might be achievable by using higher crystal temperatures during EFP or by conditioning the crystal by subjecting it to repetitive poling cycles prior to UV irradiation and the final poling steps.

The measured SHG power as a function of the fundamental pump wavelength for a crystal temperature of 100, 150, 200, and 260 °C is shown in Fig. 4. The SHG powers follow approximately a sinc-function, which is an indication of overall good domain uniformity. The maximum of the curves are at wavelengths of 1537.95, 1545.55, 1553.95, and 1565.8 nm for the corresponding crystal temperatures. The theoretical prediction for the temperatures that correspond to the measured maxima of the sinc-curves are 99, 151, 203, and 268 °C, when using the temperature dependent Sellmeier equation from Ref. 18, which is in good agreement with the chosen temperatures. The spectral bandwidth of the curves is ~1.25 nm, which is close to the expected theoretical spectral bandwidth of ~1.2 nm. The difference in the spectral bandwidth may be due to the variations in the duty cycle or the variations of the domain width itself. The graph also shows that the sinc-functions are not symmetrical, which is most apparent from the side-lobes. This may originate from a variation of the temperature profile along the sample. The monitored SHG power did not diminish over a time period of 2 h for all the temperatures, indicating that the domains are stable in this temperature range, which makes these poling patterns suitable for high power applications.

A phenomenological model that describes the formation mechanism of the bulk domains that have been observed in our experiments is schematically illustrated in Fig. 5. The wavelength of the focused UV laser light is 244 nm, which is well below the band edge of CLN, (around 350 nm). Thus,
the UV light is strongly absorbed, propagating only tens of nanometers into the surface of the MgCLN crystal before being dissipated, creating a localized heat profile with temperatures up to \( \sim 1000 \, ^\circ\text{C} \) (Fig. 5(a)). The Li ions are mobile at this temperature and diffuse into the adjacent cooler part of the crystal, resulting in Li deficient region below the irradiated tracks and a Li enriched region around it (Fig. 5(b)). As the coercive field \( E_c \) depends on the Li concentration of the crystal, it can be expected that the \( E_c \) would increase in the Li deficient region, while a decreased \( E_c \) is expected at the Li enriched region (Fig. 5(c)). When the forward poling step is applied (Fig. 5(d)), domains nucleate preferentially at the surface of the Li enriched regions next to the UV-irradiated tracks due to the reduced \( E_c \) (Fig. 5(e)). Under the influence of the applied field, the nucleated domains grow into the depth of the crystal while spreading laterally to merge with its adjacent nucleated domains. However, the Li deficient area at the surface is poling inhibited (due to the increased \( E_c \)), and thus the domain inversion will not spread into these Li deficient regions, resulting in a surface domain remaining just beneath the UV irradiated region at the end of the EFP step. This is illustrated schematically in Fig. 5(f) and corresponds to the observations in Fig. 1(a).

When the reverse poling step is applied (illustrated in Fig. 5(g)), the domains nucleate again at the Li enriched area next to the irradiated UV tracks (Fig. 5(h)), spreading laterally and merging with its adjacent nucleated domains. However, directly underneath the poling inhibited regions, the crystal polarization does not invert, as was experimentally found (Figs. 1(b) and 2) and illustrated in Fig. 5(i). This poling behavior can be explained by considering the surface domains acting as local electrical insulating barriers (similar to the properties of photoresist in standard EFP). The insulating nature of the surface domains is given, because they are already oriented along the applied electric field, therefore the internal polarization switching current cannot reach the crystal surfaces and be compensated by an external current. This is similar to the observation that a uniform surface domain layer, which is formed when fabricating titanium in-diffused poling of the crystal, can prevent electric field propagation into the Li deficient area and therefore they are even more resistive to domain inversion. Since the surface domains act as insulating barriers and the top domain wall is pinned in the Li deficient area of the PI domain, the polarization of the crystals below the surface domains prefers to have the same polarization orientation as the pinned domain wall and therefore generating the observed bulk domain, which propagates in a manner that is consistent with normal EFP.

In summary, a method for the production of bulk domains based on the UV laser pre-irradiation of the +Z surface of MgCLN crystals followed by a sequence of EFP steps have been presented. This method does not require simultaneous irradiation and application of an electric field, thus enabling the fabrication of fine periodic domain structures with periods as short as 3 \( \mu \text{m} \). A model is proposed suggesting that the bulk domains are generated because the PI surface domains, which are produced in the first poling step, act as electrical insulating barriers in subsequent poling steps. It was also determined that the uniformity in depth and parallelism of the bulk domains improved after a second full poling cycle. Although the top domain wall of the bulk domains might be pinned at defect sites, which may be influenced by elevated crystal temperatures, it was found that the bulk domains are stable at elevated temperatures while the good overall quality of the bulk domain distribution was verified by SHG experiments.

The authors acknowledge the facilities, and the scientific and technical assistance, of the Australian Microscopy & Microanalysis Research Facility at RMIT University and the Faculty of Science, Engineering and Technology at Swinburne University of Technology. This work was made possible through the generous support of the ARC Centre of Excellence CUDOS.

---