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Channel Waveguides in Lithium Niobate and Lithium Tantalate

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Abstract: Low-loss photonic waveguides in lithium niobate offer versatile functionality as nonlinear frequency converters, switches, and modulators for integrated optics. Combining the flexibility of laser processing with liquid phase epitaxy we have fabricated and characterized lithium niobate channel waveguides on lithium niobate and lithium tantalate. We used liquid phase epitaxy with K₂O flux on laser-machined lithium niobate and lithium tantalate substrates. The laser-driven rapid-prototyping technique can be programmed to give machined features of various sizes, and liquid phase epitaxy produces high quality single-crystal, lithium niobate channels. The surface roughness of the lithium niobate channels on a lithium tantalate substrate was measured to be 90 nm. The lithium niobate channel waveguides exhibit propagation losses of 0.26 ± 0.04 dB/mm at a wavelength of 633 nm. Second harmonic generation at 980 nm was demonstrated using the channel waveguides, indicating that these waveguides retain their nonlinear optical properties.

Keywords: Lithium niobate; laser processing; crystalline waveguide; liquid phase epitaxy.

1. Introduction

Integrated optics employs a wide range of miniaturized, high-speed, broad-band and reliable components suited for telecommunications, data processing, optical computing and other applications. Because of its excellent electro-optic and nonlinear properties, lithium niobate (LiNbO₃) is a key material for active optical waveguide applications [1-5].

Channel waveguides, the basic building blocks of optical circuits used in both active and passive devices of integrated optics, are fabricated by many methods. Proton exchange waveguides [6] offer high refractive index contrast in one polarization, and this technique has been employed for precision structures such as tapers [7]. Ion implantation of titanium or zinc into lithium niobate is a well-established technique for creating waveguides, but the dopant profile is sensitive to annealing [2, 8] and can be difficult to control. The dopant ions can also be substituted into the epitaxially-grown waveguides during the growth phase [9]. Direct laser treatment of ion-doped lithium niobate can induce gratings and channel optical waveguides within the surface layers [10]. Alternatively thin films of LiNbO₃ deposited on sapphire can be ablated by an excimer laser to create smooth ablated structures, with little damage to adjacent areas [11].

Ridge waveguide structures can be fabricated in planar lithium niobate devices by plasma dry etching [12, 13], wet etching [14], or ion beam etching [15], and also precision diamond cutting [16]. Alternatively femtosecond laser machining is increasingly favored for creation of ridge optical waveguides in various crystalline laser materials [17, 18]. The femtosecond laser direct writing technique has been demonstrated for the fabrication of Type I and Type II waveguides in lithium niobate, for efficient nonlinear frequency conversion [19, 20]. Laser direct writing can also facilitate
hybrid integrated structures [21]. Ridge waveguide lasers were fabricated in neodymium doped lithium niobate via femtosecond direct laser writing of gradient index planar waveguides, fabricated by Zn in-diffusion [22]. The ridge waveguide approach increases the index contrast, thus increasing the mode confinement for small bend radius waveguide features.

The crystal growth process chosen can ensure high quality crystallinity in the lithium niobate. Low-loss thin films of lithium niobate have been sputtered onto alternative substrates including sapphire and silica [23]. Liquid phase epitaxial growth on patterned substrates also offers flexible options for fabricating waveguides [24, 25]. Here we report liquid phase epitaxy growth of channel waveguides on laser-machined substrates to fabricate high optical quality lithium niobate channel waveguides with single crystalline properties on lithium niobate and lithium tantalate substrates. Liquid phase epitaxy is a versatile crystal growth technique which creates a step-index profile rather than a graded-index profile for improved mode confinement. A slight dissolution at the surface during the process leads to smoothing of the waveguide edges, thus reducing the losses. Furthermore, by using lithium tantalate substrates, high refractive index contrast can be realized for both optical polarization directions.

2. Results

2.1 Laser processing

Laser machining of the LiNbO$_3$ and LiTaO$_3$ substrates was carried out using a femtosecond pulsed Ti:sapphire laser. The laser-machined features were studied in detail with different sample feed-rates and numbers of passes. The shape of the machined features is also governed by the laser beam profile. The laser beam used in micro-machining has a typical Gaussian profile which can be expressed by:

$$I(r) = I(0)e^{-2r^2/w_0^2}$$

(1)

where $I(r)$ is the local irradiance, $I(0)$ is the peak irradiance, $w_0$ is the radius where the irradiance has decreased to 1/e$^2$ of the peak. The Gaussian irradiance profile needs to be considered when translating pulse energy to the irradiance or energy distribution on target. Laser ablation occurs when the irradiance exceeds the ablation threshold, but not all the beam profile lies above this threshold. The laser-machined feature diameter $D$ can be related to the Gaussian beam profile and irradiance threshold by:

$$D^2 = 2w_0^2 \ln \frac{I_{\text{peak}}}{I_{\text{threshold}}}$$

(2)

Experimentally this relationship is used to determine the beam waist on target and the pulse energy required for ablation from the slope $(2w_0^2)$ and intercept $(\ln(I_{\text{threshold}}))$ respectively, of Eqn. (2). Measurements of the ‘single shot’ crater diameters and depths for various pulse energies were carried out using optical profilometry (Veeco NT3300). A typical optical profile measurement is shown in Figure 1 with both a 3D visualisation and 2D cross-sectional representation of the data.
The evolution of the crater diameter as the pulse energy is varied near the ablation threshold is shown for both lithium niobate and lithium tantalate in Figure 2, with the square of the diameter plotted against the logarithm of the pulse energy plotted consistent with Eqn. (2). The experimental data suggest a beam waist of \( w_0 = 4.8\text{–}5 \mu \text{m} \) (1/e² diameter of 9.6–10 \( \mu \text{m} \)) and corresponding threshold fluence of 1.46 Jcm⁻² for lithium niobate and 1.80 Jcm⁻² for lithium tantalate, which is consistent with the results of Zhang et al [25].

![Figure 2](image_url)
Control over the trench depth was exercised by varying the sample translation speed through the laser beam and by varying the number of passes. The measured depths (from the microscope images of the trench cross-sections) for one to five passes over a range of translation speeds from 25 mm/min to 100 mm/min are shown in Figure 4. The trench depth generally shows a linear relationship with respect to the number of passes.

Microscope images of the machined substrates before the liquid phase epitaxy experiments were recorded to compare with later results after crystal growth. A typical sample is shown in Figure 5. The microscope images show the machined lithium niobate substrate samples with machined trenches of width about 13.2 μm and spacings from 5 μm up to 100 μm. All these features were pre-programmed and are adjustable.
2.2 Liquid phase epitaxial crystal growth

The channels were filled by liquid phase epitaxial growth with K$_2$O flux using established thin film growth conditions [26]. As indicated in Figure 6 (a), the growth initiated in both sidewalls of the V-groove and each nucleation site has the shape of a pyramid. Lithium niobate grown in K$_2$O flux has strong faceting effects along the (012) direction [27]. The pyramids inside the V groove were most likely caused by this faceting tendency. The pyramids continue growing at both sidewalls of the V
groove in Figure 6 (b) and join together at Figure 6 (c). In Figure 6 (d), isolated islands from different nucleation sites start to join together within the V groove. No apparent growth was observed on the planar surface outside the V groove. On the contrary, some degree of dissolution occurred along the edge of the V groove. Similar growth surface morphology was reported in an epitaxial growth model on patterned substrates of GaAs (001) surface and (111) V groove [28]. Figure 7 shows microscope images of an as-grown machined sample. For good control of the surface quality, the growth rate needs to be stable to avoid poly-crystallization, kinetic roughening and mis-orientation roughening. To achieve this, we set the growth temperature to be 0.5°C below the saturation temperature and set the growth time to be 10 minutes. As indicated in Figure 7 (LiNbO₃ channel waveguides grown on a machined LiTaO₃ substrate), the crystallization only took place inside the V grooves while leaving the plane surface untouched, which gave us much better surface quality.

Figure 7. Optical micrographs showing an example of liquid phase epitaxial growth of LiNbO₃ on a LiTaO₃ substrate at a temperature 0.5 °C below saturation temperature for ten minutes (a) top view; (b) end view.

The 3-D contour map of the liquid phase epitaxy grown sample surface is shown in Figure 8. The surface roughness inside the V groove was checked with an optical profiler (Veeco). The surface roughness of the plane surface of lithium tantalate substrate (8.6 nm) was unchanged after liquid phase epitaxial thin film growth, while the surface roughness inside the machined V groove was ~ 92 nm. The surface roughness of the as-grown lithium niobate channels was larger than that for waveguides fabricated by the titanium in-diffusion technique, which is 20-62 nm [29]. Further optimization of liquid phase epitaxial growth parameters is required to improve the surface roughness of the channel waveguides.
Figure 8. Three dimensional surface image of the as-grown liquid phase epitaxial layer on the machined lithium tantalate substrate from the optical profiler (Veeco).

2.3 Optical characterization

Optical micrographs of the channel waveguide sample used for optical characterization experiments are shown in Figure 9. Figure 9 (a) shows the top view of the machined lithium niobate substrate. A microscope image of the sample surface after liquid phase epitaxial thin film growth is shown in Figure 9 (b). The top surface of this sample was lightly polished after the liquid phase epitaxy. The end faces of the as-grown channel sample were polished, as shown in Figure 9 (c). The as-grown channels were observed using the microscope in difference interference contrast (DIC) mode. The average width of the V groove channels was about 20 µm while the average spacing between the channels was about 46 µm as determined by the initial laser machining.
To characterize the waveguiding properties within the channel waveguides, a probe beam was coupled into the guide and the near-field beam profiles were imaged by CCD camera. The waveguide sample was characterized using a linearly polarized Nd:YAB laser probe beam operating at 1064 nm. The input beam was focused down to 50 µm using a 10X microscope objective on the end face of the channel waveguide sample. The near-field of the output TE beam profile was collected by a 4X microscope objective and imaged using a TM-745 CCD camera. The near-field beam profiles were analyzed by an optical beam profiler (Spiricon) and are shown in Figure 10. The TM modes are not supported in this sample with lithium niobate as a substrate.
Figure 10 Near-field beam profiles for TE modes propagating at 1064 nm in channel waveguide samples obtained by imaging with a 10X microscope objective.

As shown in the image, the beam has two V shaped profiles with beam radius about 20 μm in the X direction and 14 μm in the Y direction. The beam shape and size are comparable with the machined features. The input laser beam was coupled into two channel waveguides. The spacing between the two beams is 42 μm which is also the same as the spacing between the machined V grooves.

The near-field beam profiles indicate well-confined waveguide structures. The near-field profiles from the channel waveguide sample were further characterized by end-fire coupling of 975 nm and 633 nm light from single mode optical fibre. The output near-field profiles imaged by a 40X microscope objective are shown in Figure 11.

Figure 11. Near-field beam profiles for (a) 633 nm and (b) 975 nm from the same channel with lithium tantalate as the substrate; imaging with a 40X microscope objective.
The near-field beam profiles indicate multimode propagation within the channel, [30] which is consistent with the refractive index contrast ($\Delta n_o \sim 0.1$) provided by the lithium tantalate substrate and lithium niobate channels. Numerical simulations were performed using BeamPROP (part of the RSoft Photonics CAD Suite) which is based on a finite difference beam propagation method assuming 10 nm step sizes in the $x$, $y$ and $z$ directions. The simulated TE mode profiles are plotted in Figure 12. The simulation is based on the channel size measured by the microscope images and the refractive index for bulk lithium tantalate and lithium niobate at a wavelength of 0.980 $\mu$m for TE mode propagation. The calculated mode profiles have similar size to the measured beam profiles and are also multimode.

Temperature-tuned second harmonic generation of a 975 nm diode laser was also measured from the lithium niobate channel waveguides. The second harmonic generation phase-matching temperature for stoichiometric lithium niobate is 42 °C for 979 nm fundamental light, and blue second harmonic light is detectable from the channel samples at room temperature, indicating the good crystallinity of the channel waveguide. The second harmonic output spectrum is plotted in Figure 13.
Figure 13. Second harmonic generation spectrum detected from lithium niobate channel waveguides with lithium tantalate substrate.

The propagation loss of the as-grown channel sample was measured by the back-reflection method at 633 nm. The operational principle of the back-reflection method is sketched in Figure 14. The propagation loss was extracted from the ratio between the output signal power ($P_{\text{out}}$) and the back-reflected signal power ($P_{\text{back}}$) [31]. The propagation loss was $0.26 \pm 0.04$ dB/mm. The relatively high propagation loss within the channel samples is probably due to the interaction of higher order modes with the surface roughness of the lithium niobate channel. We expect that thinner and smaller channels, operating in a single mode regime, would be less lossy.

Figure 14 Scheme of the back-reflection method.

3. Discussion

Liquid phase epitaxial growth on laser micro-machined lithium tantalate and lithium niobate substrates is a practical and promising way to fabricate channel waveguides, and allows rapid prototyping of the guide design to be implemented. Lithium niobate and lithium tantalate substrates were first patterned by laser micro-machining. The fabrication of desired features is programmable with prescribed parameters such as pulse energy, the number of passes on the same area, the feed-rate of the sample with respect to the laser spot and the laser beam profile. After liquid phase epitaxy growth of lithium niobate on the patterned substrate, the surface roughness was measured to be ~92 nm, compared with ~60 nm for channel waveguides fabricated by the ion in-diffusion technique. In the case of liquid phase epitaxial growth, the waveguide is inherently single crystalline.

The beam confinement of the channel waveguide sample was examined by propagating a linearly-polarized Nd:YAB probe beam. The output beam, analyzed by a beam profiler, showed coupling into one or two channels, with a spacing of 42 µm. The near-field beam profiles have typical
“V” shapes with beam width about 20 μm in the X direction and 14 μm in the Y direction. The near-field beam profile was further examined using a high magnification objective with a single mode fibre probe at 975 nm and 633 nm. The beam profile indicates multi-mode propagation within the channel sample. The beam profiles are consistent with the results of RSoft modelling, providing evidence for the high refractive index contrast (~0.1) in the as-grown channel samples. Second harmonic generation from the channel samples was demonstrated from a fundamental wavelength of 975 nm. The propagation loss was 0.26 ± 0.04 dB/mm, measured by the back-reflection method at 633 nm.

4. Materials and Methods

4.1 Femtosecond laser machining of substrates

Laser machining of the LiNbO₃ and LiTaO₃ substrates was carried out using a femtosecond laser micromachining facility (OptoFab, Australian National Fabrication Facility), which incorporates an infrared femtosecond pulsed laser (Spectra Physics Hurricane), 3D motion control stages (Aerotech) and diagnostic and alignment tools to create an integrated and completely automated experimental setup for fabricating optical devices. The experimental setup for laser machining is shown in Figure 15.

The Spectra-Physics Hurricane comprises the Mai-Tai Ti:Sapphire 80 MHz oscillator with a 1 kHz regenerative amplifier. The system produces ~100 fs pulse-width pulses with up to 1mJ energy with a default 1 kHz repetition rate pulse train which may be reduced or controlled externally. The power on target is controlled by a half-wave-plate-linear-polarizer arrangement with the half-wave plate held in a computer-controlled rotation mount for power control. The pulse duration can be controlled by tuning the compression stage gratings.

To fabricate lithium niobate channel waveguides, we used laser micro-machining to write a pattern on the surface of the substrates. The pattern described here was fabricated by moving the substrate through the focus of the laser beam. For example, laser-ablated trenches on the surface of the substrates were attained by moving the sample several passes through the laser focus spot. The depth of the laser-cut features was controlled by the pulse energy, the number of passes on the same area and the feed-rate of the sample with respect to the laser spot. The effect of the pulse energy was studied by comparing the crater morphologies produced by single shot ablation on the substrate materials for various laser parameters. The appropriate laser pulse energy generates smooth craters with little debris deposited on the surrounding surface. To determine the effects of number of passes and sample feeding rate, ideally, we assume the depth of the machined features linearly scales with
the number of passes and scales inversely with sample feeding rate. However deviations may arise from changing surface morphology and material properties in the laser-treated area.

4.2 Channel waveguides grown by liquid phase epitaxy on laser-machined substrates

Liquid phase epitaxy growth was carried out on the laser micro-machined substrates. The experimental setup and procedure were similar to that reported previously for planar samples [26]. Briefly, we used a K$_2$O (13.8 mol%) Li$_2$O (43.1 mol%) Nb$_2$O$_5$ (43.1 mol%) flux, at a temperature of 1100°C and with z-cut lithium niobate or lithium tantalate substrates. The laser-machined substrate was dipped into the preheated molten solution just below the saturation temperature for about 10 minutes. The growth rate was approximately 4.5 μm/min.

The changed surface morphology of the substrate due to laser machining requires more stringent growth controls. The features fabricated by laser machining induce an inhomogeneous morphology on the sample surface at the initial stage of thin film growth. For the z-cut lithium niobate substrate, the surface is the (001) plane while the sidewall inside the machined “V” groove has a different orientation. The initiation of growth during liquid-phase epitaxy takes place in isolated islands [32]. The islands grow and join up to give a smooth surface. For liquid phase epitaxy growth on a patterned surface, the site environment inside the V-groove is different from that of the planar surface, which results in different diffusion rates and growth rates. The isolated islands developing from different sites show steps at the boundaries of adjacent islands.

The roughness of the epitaxial layer grown on the machined samples arises from two causes. The first is kinetic roughness, which arises from a high deposition rate occurring at the low growth temperature. The second is substrate mis-orientation inside the machined sidewall. The sample was dipped vertically into the LiNbO$_3$ flux (see Figure 16 (b)) at the saturation temperature for ten minutes with the machined features aligned vertically inside the furnace. The temperature profile shown in Figure 16 (a) was obtained by suspending a thermocouple inside the empty crucible at different positions.

![Figure 11. (a) Temperature profile in the crucible as a function of position. (b) Schematic of the crystal growth furnace layout for liquid phase epitaxial growth of channel waveguides.](image-url)
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