

Role of cerium in lithium niobate for holographic recording

Xuefeng Yue, A. Adibi, T. Hudson,^{a)} K. Buse, and D. Psaltis^{b)}
*California Institute of Technology, Department of Electrical Engineering, MS 136-93, Pasadena,
California 91125*

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Cerium-doped lithium niobate crystals are tested for holographic recording. A photochromic effect is observed in crystals doped with cerium and manganese. But two-center recording in the sample is not as effective as in iron and manganese doubly doped crystals. Photocurrent measurements in cerium and iron singly doped crystals indicate that the photovoltaic constant in the cerium-doped crystal is only one third of that of the iron-doped one. This is the main reason accounting for the low sensitivity of cerium-doped lithium niobate crystals. However, in the diffusion dominated case, i.e., for reflection geometry, cerium-doped lithium niobate may give a strong effect. © 2000 American Institute of Physics. [S0021-8979(00)08505-4]

I. INTRODUCTION

Holographic data storage is a promising candidate for the next-generation mass-storage system. It offers two substantial advantages over conventional storage technologies: (1) multiple pages of data can be stored in the same volume, (2) many bits are recalled in parallel. This enables a storage device that has potentially a high storage density and data transfer rate.¹ Inorganic photorefractive oxide crystals such as lithium niobate (LiNbO₃,LN) have attracted much attention in the past due to their reversibility, moderate sensitivity, and availability in large size and good quality.¹⁻⁴ The storage of volume phase holograms in LiNbO₃ relies on the presence of localized centers containing electrons that can be optically excited into the conduction band. Various transition metals have been introduced into LiNbO₃ for the improvement of sensitivity and dynamic range.^{5,6} Among various dopants, Fe is most effective in producing large improvements in both the sensitivity and the maximum diffraction efficiency. Recently two-center recording in Fe and Mn doubly doped LiNbO₃ has been realized to achieve persistent storage.⁷ Illumination with ultraviolet light excites electrons from deep center (Mn) to the conduction band and some of them may get trapped at the shallower center (Fe), which permits the storage of data by using red light (wavelength 633 nm), but no erasure during the subsequent readout in the absence of ultraviolet light. For the improvement of this technique, a more red-sensitive dopant, which can replace Fe, is very important.

Cerium is known to be an effective trap center, which can provide and capture charge carriers, in different kinds of photorefractive crystals including strontium-barium-niobate (SBN)⁸ and barium-titanate (BaTiO₃).^{9,10} Ce-doped SBN has a very high sensitivity at visible wavelength,⁸ and Ce-doped BaTiO₃ is sensitive to both visible and near infrared light.^{9,10} Some previous reports on Ce and Fe doubly doped LiNbO₃ suggest that this material has a wide spectral re-

sponse and shows higher sensitivity.^{11,12} Charge transport from and to Fe centers is already well known, but the impact of Ce doping on the buildup of space-charge fields in LiNbO₃ is still unclear. McMillen *et al.* have reported holographic recording in specially doped LiNbO₃ crystals including a Ce singly doped one,¹³ but no systematic consideration for the role of cerium has been carried out. In our present work, we have found that the Ce and Mn doubly doped LiNbO₃ has a strong photochromic effect, which may be very attractive for two-center persistent data storage. In this contribution, we investigate the photorefractive performance of Ce-doped LiNbO₃ crystals. Comparisons will be made among the nominally pure, Ce-, Fe-, and Mn-doped samples.

II. SAMPLES

Singly doped (Ce, Fe, and Mn) and doubly doped (Ce:Mn, Fe:Mn)LiNbO₃ samples as well as a nominally pure one are used in present experiments. Notations, doping levels, and dimensions of the samples are listed in Table I. All doping levels refer to the values introduced into the melt. It is known that even in nominally pure crystals there are usually some background impurities incorporated,¹⁴ but the concentrations of such background impurities are generally much smaller than the intentionally doped impurity levels.

All samples are *y* cut and polished to optical quality. Two kinds of thermal treatments have been conducted: oxidation and reduction. During oxidation, the samples are kept in an oven with oxygen atmosphere at 1000 °C for at least 12 h, while they are heated to 1000 °C in argon atmosphere for 12 h during reduction. In this way, the absorption of the doped samples can be significantly changed, i.e., the valence states of the trap centers can be varied.

III. EXPERIMENTS AND RESULTS

As mentioned above, the main purpose of this work is to test whether Ce is an effective trap center in lithium niobate crystals and to measure the performance of Ce-doped crystals for holographic recording, especially in the red. So we conducted measurements including conventional holographic

^{a)}Also with: U.S. Army Aviation and Missile Command, Redstone Arsenal, AL 35898-5248.

^{b)}Author to whom correspondence should be addressed.

TABLE I. Description of the samples used in the experiments. The doping concentrations are wt % of the oxide (Fe_2O_3 , Ce_2O_3 , and MnO) in the melt and dimensions are $a \times b$ (thickness) $\times c$ in mm^3 .

Notation	Dopant	Doping level	Dimensions
LN	normally pure	-	$5 \times 5 \times 7$
Ce:LN	Ce	0.01–0.02	$5 \times 5 \times 7$
Fe:LN	Fe	0.01–0.02	$5 \times 5 \times 7$
Mn:LN	Mn	0.05	$4 \times 1 \times 6$
Ce:Mn:LN	Ce	0.085	$10 \times 2 \times 10$
	Mn	0.01	
Fe:Mn:LN	Fe	0.085	$10 \times 2 \times 10$
	Mn	0.01	

recording in singly doped samples, sensitization by ultraviolet light, and two-center recording as well as bulk photovoltaic current measurements.

A. Photochromic effect

It is known that the crystals are often photochromic if two kinds of trap centers exist between the valence and conduction bands, which means that the absorption can be changed by illumination because of the redistribution of charge carriers between these different traps. Three samples, i.e., Mn:LN, Fe:Mn:LN, and Ce:Mn:LN are used in sensitizing measurements. All crystals are strongly oxidized before the sensitization measurements were started to ensure that most of electrons are removed from the shallower trap centers (Ce or Fe).

In our experiment, we use ultraviolet light (wavelength 404 nm and intensity 4 mW/cm^2) to illuminate the samples. A very weak red beam (633 nm) is used to probe the transmission as a function of illuminating time. Figure 1 shows the variation of the normalized transmission versus time for different samples. The relative change of transmission in Ce:Mn:LN is much larger than that in Fe:Mn:LN. We perform the sensitization measurement with the Mn singly doped sample, too. As it can be seen from Fig. 1, no change in absorption can be observed in this case, which means that the absorption variation in Ce:Mn:LN can be attributed to

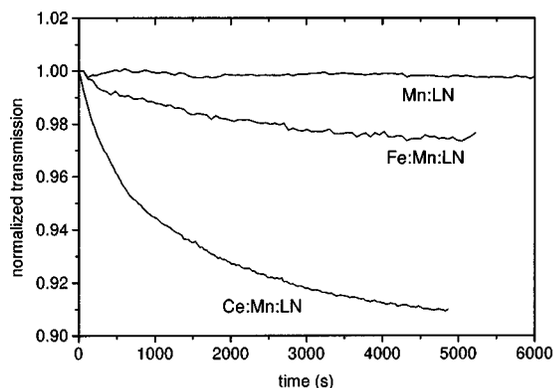


FIG. 1. Variation of normalized transmission at 633 nm for variously doped LiNbO_3 during illumination with ultraviolet light. Wavelength and intensity of the ultraviolet light are 404 nm and 4 mW/cm^2 , respectively. The samples were first strongly oxidized to ensure that most of the shallower centers are emptied.

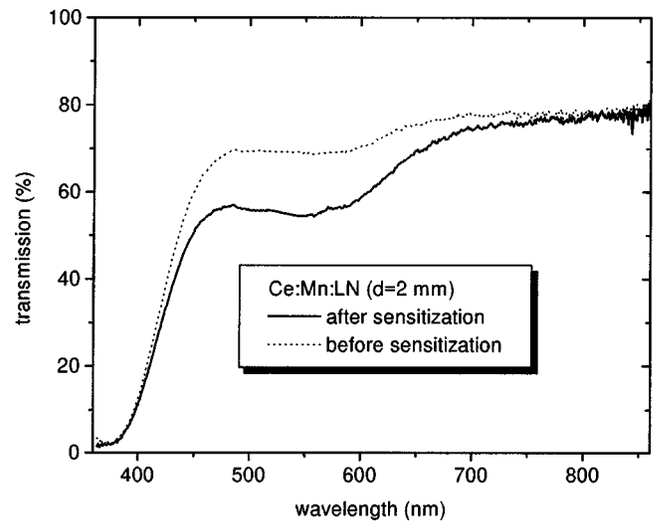


FIG. 2. Transmission spectra of the Ce and Mn doubly doped sample (Ce:Mn:LN) before (dotted curve) and after (solid curve) ultraviolet illumination. Wavelength and intensity of the ultraviolet light are 404 nm and 4 mW/cm^2 , respectively, and the illumination time is 2 h.

the filling of Ce traps with the help of ultraviolet light and not to some kinds of emptied background impurities.

The transmission spectra before and after ultraviolet illumination in Ce:Mn:LN are measured and the results are presented in Fig. 2. It can be seen that after 2 h illumination by ultraviolet, there is a broad induced absorption in the range from 450 to 650 nm. The large absorption caused by ultraviolet makes the material promising for two-center recording at red.

B. Sensitivity

A He–Ne laser operating at 633 nm is used for holographic recording. The total intensity I_0 of the writing beams is 26 mW/cm^2 . Both writing beams are ordinarily polarized and impinge on the crystal symmetrically at an incident angle of 23° in air. The grating vector of the interference pattern is aligned along the crystallographic c axis. The diffraction efficiency η is defined as the ratio between the diffracted and incident beam intensities, while the recording sensitivity is defined by $S = (\partial \sqrt{\eta} / \partial t) |_{t=0} / (I_0 d)$, where d is the thickness of the sample.

First we strongly reduce the samples including LN, Ce:LN, Ce:Mn:LN, and Fe:LN. In Fig. 3 we show the recording and erasure curves obtained in nominally pure, Ce-doped and Fe-doped samples. All of the samples have the same thickness. From these measurements, we get the sensitivity data: $S(\text{LN}) = 0.002 \text{ cm/J}$, $S(\text{Ce:LN}) = 0.007 \text{ cm/J}$, $S(\text{Ce:Mn:LN}) = 0.020 \text{ cm/J}$, $S(\text{Fe:LN}) = 0.033 \text{ cm/J}$. Clearly, the increase of Ce concentration leads to an increase of recording sensitivity. Mn traps are too deep to get involved in holographic recording in red. Therefore, holographic recording in Ce:Mn:LN is similar to a LiNbO_3 crystal doped with only 0.085 wt % Ce, corresponding to Ce concentration of $C_{\text{Ce}} = 14 \times 10^{24} \text{ m}^{-3}$. This is two times as large as the concentration in Fe:LN ($C_{\text{Fe}} \leq 7 \times 10^{24} \text{ m}^{-3}$). However, the sensitivity in Ce:Mn:LN is still smaller than that in Fe:LN.

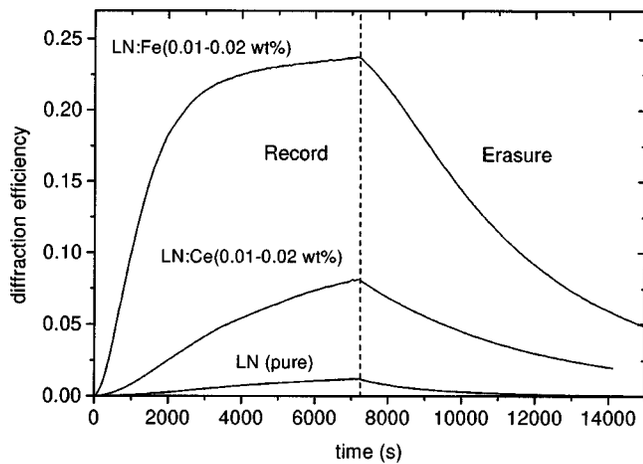


FIG. 3. Evolution of the diffraction efficiency during holographic recording and erasure in nominally pure, Ce-doped and Fe-doped LiNbO₃. The three samples have the same dimensions with the thickness $d=5$ mm (wavelength of writing beam 633 nm, total intensity 26 mW/cm², ordinary polarized, grating vector along c axis, and grating spacing 0.8 μ m).

C. Two-center recording

The large light-induced absorption at 633 nm caused by ultraviolet light in Ce:Mn:LN is a very promising aspect for two-center recording, but the smaller sensitivity of holographic recording in Ce-doped LiNbO₃ compared to that in the Fe-doped sample is a discouraging result. To be certain whether Ce:Mn:LN is a good alternative to Fe:Mn:LN, we conduct two-center recording in Ce:Mn:LN.

Two-center recording in Ce:Mn:LN is carried out by the following procedure: the strongly oxidized sample is first illuminated by incoherent ultraviolet light for about 2 h, then two ordinarily polarized red beams are turned on with the ultraviolet light still present. After 3 h recording, the ultraviolet light and one of the red beams are turned off. The diffracted beam intensity is monitored for about 10 h. As shown in Fig. 4, the erasure of the recorded grating consists of two parts: a fast decay and a relatively slow decay. These are typical characteristics of two-center recording.⁷ The same procedure is conducted for Fe:Mn:LN. A maximum diffraction efficiency of 25% is reached in this case, which is approximately 10 times as large as that obtained in Ce:Mn:LN.

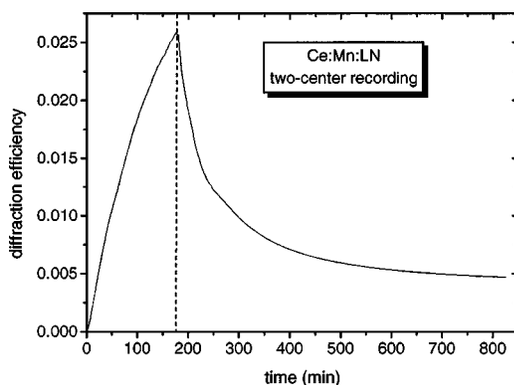


FIG. 4. Evolution of the diffraction efficiency during two-center holographic recording in Ce and Mn doubly doped LiNbO₃. During recording, ultraviolet light illuminates the sample, while it is blocked during readout.

D. Photovoltaic current

Recording sensitivity and saturated diffraction efficiency depend on the photovoltaic constant.^{6,15} During homogeneous illumination, the short-circuited photocurrent density is proportional to the absorbed power density, i.e.,

$$J = \kappa \alpha I, \quad (1)$$

where I is the incident intensity, α is the absorption coefficient, and κ is the photovoltaic constant depending mainly on absorption center and wavelength. For the thick samples, the light depletion due to absorption must be considered and an averaged light intensity $\bar{I} = I[1 - \exp(-\alpha d)](\alpha d)^{-1}$ must be used.

Measurements of the photocurrents are made with an electrometer having an input impedance much less than the crystal impedance. For illumination, we use a diode-pumped frequency-doubled Nd:YAG laser (wavelength 532 nm and power 400 mW). The ordinarily polarized beam is expanded to illuminate the whole sample homogeneously. All measurements are made at room temperature (25 °C). After the illumination beam illuminates the sample for at least 10 min, the steady-state photocurrent is detected. The delay is required to eliminate the influence of pyroelectric currents.¹⁶ The determined photovoltaic constant for reduced Fe:LN is $\kappa = 1.3 \times 10^{-9}$ cm/V at 532 nm, which is in good agreement with the results reported by Krätzig and Kurz.⁶ For reduced Ce:LN, $\kappa = 0.4 \times 10^{-9}$ cm/V at 532 nm is measured. Ordinarily polarized red light (633 nm) is also employed to illuminate the samples. In the case of Fe:LN $\kappa = 0.8 \times 10^{-9}$ cm/V is obtained, while the value of the photovoltaic constant in Ce:LN cannot be determined under the present experimental accuracy.

E. Reflection grating

In LiNbO₃ crystals, bulk photovoltaic currents as well as diffusion currents contribute to formation of the space-charge field during holographic recording. With a decrease of the grating spacing, the diffusion field becomes more important.¹⁷ The dynamics of the formation of the reflection grating can be used for detection of diffusion-related properties.

In our experiment, two ordinarily polarized beams impinge on the two opposite c faces of the Ce:LN crystal, with an angle of incidence of about 5° in air. The length of the sample along the c axis is 7 mm. The diffraction efficiency as a function of recording time is monitored. For the sake of comparison with the performance of transmission gratings, we convert the efficiency into the modulation of refractive index by taking reflection into account using the following relations:¹⁸

$$\eta = R \sin^2 \left(\frac{\pi \Delta n d}{\lambda \cos \theta_i} \right), \quad (2)$$

for the transmission grating and

$$\eta = R \tanh^2 \left(\frac{\pi \Delta n d}{\lambda \cos \theta_i} \right) \quad (3)$$

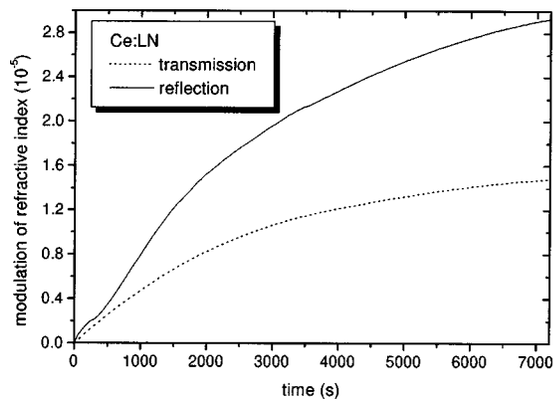


FIG. 5. Modulation of the refractive-index as a function of recording time for reflection geometry (solid curve) and transmission geometry (dotted curve) in Ce:LN. Wavelength and intensity of writing beam are 633 nm and 26 mW/cm², respectively. In transmission geometry both beams impinge on the *b* face with an angle of 23°, while in reflection geometry both beams impinge on the two opposite *c* faces with angle 5° in air. The grating vector of the interference pattern is in both cases aligned along the crystallographic *c* axis.

for the reflection grating. In the above equations, R is the factor caused by reflection loss, θ_i is the angle of incidence inside the crystal, d is the thickness along the grating vector, and the absorption is not considered due to the small value at red. As shown in Fig. 5, recording in the reflection geometry is faster than that in the transmission geometry. The corresponding sensitivity in the case of reflection geometry is $S = 0.009$ cm/J. The modulation of refractive index in the reflection geometry is twice as large as that in the transmission grating for Ce:LN. In Fe:LN, however, the reflection grating is only 1.4 times stronger than the transmission grating.

IV. DISCUSSION

The experimental results described above clearly show that Ce is a photorefractive dopant for LiNbO₃. An increase of the Ce concentration leads to a substantial increase of the sensitivity in the red spectral region. It is very interesting that the absorption change in Ce:Mn:LN with the aid of ultraviolet illumination is larger than that in the Fe:Mn:LN. Because red light essentially cannot excite electrons from Mn traps, the absorption of doubly doped LiNbO₃ for red light is proportional to the density of the filled shallower traps (Fe or Ce). Anyhow, the recording sensitivity in Ce singly doped LiNbO₃ is smaller than that in the Fe-doped one even through the concentration of Ce ions is twice as large as that of Fe ions. We know that in LiNbO₃ both the sensitivity and the maximum diffraction efficiency depend on the photovoltaic constant. So the smaller sensitivity in the Ce-doped sample can be attributed to the much smaller photovoltaic constant. Photovoltaic constant can be expressed as¹⁹

$$\kappa = (e/h\nu_0)\Phi L, \quad (4)$$

where $h\nu_0$ is the photon energy, Φ is the quantum efficiency that an absorbed photon will produce in a photoionized charge, and L is the migration length of an electron in the conduction band before it is recaptured. From the results

obtained with light of 532 nm, we get $\Phi L = 3 \times 10^{-11}$ m for the Fe-doped sample, while ΦL is approximately 1×10^{-11} m for the Ce-doped sample.

The amplitude of the steady-state space-charge field E_{sc} in photorefractive materials is determined by²⁰

$$E_{sc} = \left[\frac{E_p^2 + E_D^2}{(1 + E_D/E_q)^2 + (E_p/E'_q)^2} \right]^{1/2}, \quad (5)$$

where E_p is the photovoltaic field which is proportional to the ionized trap density, E_D is the diffusion field which depends linearly on the amplitude of the grating vector, and E_q and E'_q are limiting space-charge fields determined by effective trap density and filled trap density, respectively. In the case of oxidized Fe:LiNbO₃ where the density of emptied Fe centers is much larger than that of filled centers, the saturated refractive index is inversely proportional to the grating spacing. This means that a reflection grating should have a smaller amplitude compared to that of a transmission grating.²¹ But the sample we used in the reflection geometry is strongly reduced. If we suppose that no limitation of space-charge fields is present, Eq. (5) simplifies to

$$E_{sc} = (E_p^2 + E_D^2)^{1/2}. \quad (6)$$

We know that E_p is the same in both transmission and reflection geometry. However, E_D is larger in reflection geometry. This is due to the fact that E_D is inversely proportional to the grating spacing. Therefore, we expect to obtain a larger E_{sc} . The enhancement of performance for the Fe-doped sample in the reflection geometry is not so large. This can be attributed to a much higher photovoltaic field, which is independent of the amplitude of the grating wave vector.

In conclusion, Ce is an active red-sensitive photorefractive center in LiNbO₃ and can play an important role in charge transport during holographic recording. However, the photovoltaic constant in Ce-doped crystals is smaller than that in Fe-doped ones. This makes the recording in Ce-doped crystals not as sensitive as that in Fe-doped crystals. In reflection geometry, where diffusion field makes more contribution, the photorefractive performance of Ce-doped samples is enhanced.

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¹D. Psaltis and F. Mok, *Sci. Am.* **273**, 70 (1995).

²J. F. Heanue, M. Bashaw, and L. Hesselink, *Science* **265**, 749 (1994).

³M. P. Bernal *et al.* *Appl. Opt.* **35**, 2360 (1996).

⁴I. McMichael, W. Christian, D. Pletcher, T. Y. Chang, and J. H. Hong, *Appl. Opt.* **35**, 2375 (1996).

⁵W. Phillips and D. L. S. J. J. Amodei, *RCA Rev.* **33**, 94 (1972).

⁶E. Krätzig and H. Kurz, *Opt. Acta* **24**, 475 (1977).

⁷K. Buse, A. Adibi, and D. Psaltis, *Nature (London)* **393**, 665 (1998).

⁸K. Megumi, H. Kozuka, M. Kobayashi, and Y. Furuhta, *Appl. Phys. B: Photophys. Laser Chem.* **30**, 631 (1977).

⁹C. Yang, Y. Zhang, P. Yeh, and X. Wu, *Opt. Commun.* **113**, 416 (1995).

¹⁰S. Dou, Y. Lian, H. Gao, Y. Zhu, X. Wu, C. Yang, and P. Ye, *Appl. Opt.* **34**, 2024 (1995).

¹¹K. Xu, H. Xu, Y. Yuan, J. Hong, and Y. Xu, *Proc. SPIE* **1078**, 331 (1989).

- ¹²S. Yin and F. T. S. Yu, *IEEE Photonics Technol. Lett.* **5**, 581 (1993).
- ¹³D. K. McMillen, T. D. Hudson, J. Wagner, and J. Singleton, *Opt. Express* **2**, 491 (1998).
- ¹⁴K. Buse, *Appl. Phys. B: Lasers Opt.* **64**, 273 (1997).
- ¹⁵A. M. Glass, D. von der Linde, and T. J. Negran, *Appl. Phys. Lett.* **25**, 233 (1974).
- ¹⁶X. Yue, S. Mendricks, T. Nikolajsen, D. Kip, H. Hesse, and E. Krätzig, *J. Opt. Soc. Am. B* **16**, 389 (1999).
- ¹⁷A. Krumins, Z. Chen, and T. Shiosaki, *Opt. Commun.* **117**, 147 (1995).
- ¹⁸H. Kogelnik, *Bell Syst. Tech. J.* **48**, 2909 (1969).
- ¹⁹R. Grousson, M. Henry, S. Mallick, and S. L. Xu, *J. Appl. Phys.* **54**, 3012 (1983).
- ²⁰N. V. Kukhtarev, V. B. Markov, S. Odoulov, M. S. Soskin, and V. L. Vinetskii, *Ferroelectrics* **22**, 949 (1979).
- ²¹K. Peithmann, A. Wiebrock, and K. Buse, *Appl. Phys. B: Lasers Opt.* **68**, 777 (1999).