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Citation: Applied Physics Letters 105, 191103 (2014); doi: 10.1063/1.4901528
View online: http://dx.doi.org/10.1063/1.4901528
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(Received 4 September 2014; accepted 31 October 2014; published online 10 November 2014)

Two-photon ionization by focused femtosecond laser pulses initiates the development of micrometer-scale plasmas in the bulk of silicon. Using pump-and-probe transmission microscopy with infrared light, we investigate the space-time characteristics of these plasmas for laser intensities up to $10^{12}$ W/cm². The measurements reveal a self-limitation of the excitation at a maximum free-carrier density of $\approx10^{19}$ cm⁻³, which is more than one order of magnitude below the threshold for permanent modification. The plasmas remain unchanged in the ~100 ps timescale revealing slow carrier kinetics. The results underline the limits in local control of silicon dielectric permittivity, which are inherent to the use of single near-infrared ultrashort Gaussian pulses. © 2014 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4901528]

In the past two decades, high¹ (up to ~MJ/cm³) and localised² (down to 0.008 μm³ volumes) energy concentrations inside non-conducting solids have been demonstrated. At these energy densities, sub-bandgap wavelength light can be used to locally generate free carriers by nonlinear interaction, thus transforming an otherwise transparent solid in an absorbing one. At this stage, the dielectric permittivity of the medium is changed, leading to an efficient transfer of energy from the laser to the lattice. Finally, if the deposited energy concentration exceeds the threshold for permanent modification, irreversible damage may take place. As a result, numerous applications appeared in dielectrics including optical storage³ and the direct writing of waveguides⁴,⁵ and other integrated optical functionalities.⁴

Silicon (Si) becoming an increasingly dominant material in the field of hybrid devices (optical and electronic), similar developments (e.g., waveguide writing) are highly desired for semiconductors. However, the development remains far behind and there is so far no demonstration of true 3D control of induced modifications inside Si. On the contrary, recent works report on a strong delocalization of intense light behind and there is so far no demonstration of true 3D control of induced modifications inside Si. Nevertheless, even below the permanent modification level, reversible changes, which are the result of dynamic control of the dielectric permittivity, are beneficial for a certain number of applications including electrical circuit defect analysis⁷ and cryptography⁸. All these processes underline the importance of quantitative information related to the transient evolution of the Si dielectric permittivity.

In this paper, we propose the study of the dielectric permittivity in bulk crystalline Si (c-Si) excited by near infrared (NIR) light. Ultrashort laser pulses at 1.3-μm wavelength (0.95 eV) have been focused inside the 1.1-eV bandgap Si. The nonlinear nature of the induced ionization⁹ allows for the generation of well localised carriers, at which we will be referring as microplasma, inside the crystal without observing any permanent modification. Pump-and-probe infrared imaging technique is used to study the spatio-temporal evolution of the plasmas. At this stage, let us mention that the necessary temperature for the bandgap close-up (down to 0.95 eV) with temperature is ~850 K (Ref. 10) and that we are far from reaching these temperatures in our conditions as we will see later. Furthermore, our samples have been tested for defects and no significant linear absorption has been measured at 1.3-μm wavelength.

As shown in Fig. 1, a Ti-Sa laser system (ASUR facility, Amplitude technologies) is delivering 30 fs pulses at 800 nm

![Image](463x179 to 469x224)

FIG. 1. Experimental setup using: (Ti:S) Ti:sapphire femtosecond laser; (S) mechanical shutter electrically commanded; (OPA) optical parametric amplifier for wavelength conversion at 1.3 μm; (BS) 10:90 beamsplitter; (HWP) half waveplate; (DL) delay line; (PD) photodiode; (OB) objective; (IS) integrating sphere; (PC) polarizing cube; and (ES) energy sensor. Inset: Intensity of the transmitted probe (a.u.) in presence of a pump pulse (10 nJ, 10 ps delay) as acquired with the sensor. The microplasma is visible (black stripe) at the focus of the pump beam.

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105, 191103-1 © 2014 AIP Publishing LLC
with a repetition rate of 100 Hz. A high energy optical parametric amplifier (OPA) system (TOPAS-HE, Light Conversion) is used to tune the wavelength to 1.3 μm yielding linearly polarised pulses with energy up to 1 mJ and duration τ ≃ 80 fs (FWHM) as measured using a single shot auto-correlator (TiPAA5C3, Light Conversion). A beamsplitter reflects 10% of the beam to create the probe pulse. The remaining part (pump) propagates along a half-wave plate and a polarising cube combination to control the incident energy, measured with an integrating sphere and a photodiode. Our measurements are reported in terms of energy delivered at the focus as if no sample were in place (reflection from the sample is not being taken into account).

The pump pulse is focused (NA = 0.3) with a NIR microscope objective (Olympus, LMPLN10XIR). The sample is an intrinsic c-Si wafer (Siltronix) with 200 Ω·cm resistivity, (100) orientation, and 1 mm thickness. The transmitted pump intensity is measured with a high-sensitivity photodiode energy sensor (OPHIR, PD10-IR-PJ). Then, using a so-called Z-scan transmission measurement procedure, we can position precisely the pump beam focus at a depth of 400 μm below the surface of the sample before the experiments.11 This ensures sufficiently low intensity on the surface to investigate only bulk effects. Motorised stages ensure sample translation between shots so that each single pulse interacts with a fresh material. Before the plasma investigations, we used a high resolution imaging setup (not shown) to fully characterize the laser intensity distribution in the focal region. We found a nearly Gaussian distribution with spot size and confocal parameter of 4 μm and 20 μm (FWHM) in air respectively (corresponding to 4 μm and 70 μm in Si). Finally, lateral imaging of the interaction region is made with a 20× magnification (LMPLN20XIR microscope objective, Olympus) on an InGaAs detector (XenicsXeva) yielding observations with ±1.4 μm spatial resolution.

According to the linear response of our detector, we display directly the probe transmission through the plasma by normalisation of the images using the signal outside the plasma (background signal). The plasma transmission retrieval assumes that reflection from the generated plasma is negligible, which is reasonable for N < 10^20 cm^-3 yielding K < 1%. To avoid any false considerations, we have also performed calibration measurements (not shown here) to verify the linearity of probe absorption by the plasma at the intensity used for the imaging experiments. Then, acquiring images for different pump-probe delays and pump energies allows us to investigate the microplasmas.

A time-resolved transmission measurement along the optical axis of the pump (longitudinal direction of the plasma) is shown in Fig. 2. One can see a significant drop over the first ~130 ps. Since thermal equilibrium between free-carriers and the lattice can be rather fast in Si (~500 fs), the first step was to estimate the influence of a potential bandgap close-up due to pump energy transfer in the focal volume. The difference between pump photon and bandgap energy being ~150 meV, linear absorption could arise from a temperature raise of ~550 K. With simple thermodynamic considerations, one can estimate the maximum local heating in the experiments as ΔT(C) = E/(V × ρ × C) with E as the total pump energy, V as the interaction volume (modelled by a cylinder 100 μm long over a 3 μm diameter, see inset of Fig. 1), ρ as the Si density (2.33 g·cm^-3), and C as the thermal capacity (0.712 J·g^-1·K^-1). This yields a potential local heating ≤ 1 K in the framework of our experiments (pump energy ≤ 100 nJ) and thus we can neglect any thermal contribution to the observed absorption.

Free from this consideration, we can now attribute the observed probe absorption to inverse Bremsstrahlung absorption by the free-carriers generated by two-photon absorption (TPA) of the pump pulse.15 According to the time-resolved response at the theoretical focus shown in Fig. 3, we measure a rise time of the absorption of ~300 fs. This compares favourably with the convolution product of the pump and probe pulses that are stretched before to reach the interaction region due to dispersion in Si. For comparison, we have measured that our pulses are stretched up to 120 fs

FIG. 2. (a) Probe transmission along the optical axis of the pump (depth) versus pump-probe delay inside Si. Depth axis is translated so that zero corresponds to 400 μm below the surface. Pump energy is 85 nJ. (b) The transmission measured at the center of the micro-plasma is plotted (dotted line on Fig. 2(a)), as a function of delay, for two different pump pulse energies: 5 nJ and 85 nJ.

FIG. 3. Probe transmission along the center of the micro-plasma as a function of the pump and probe delay. Pump pulse energy is ~18 nJ. For comparison, the solid line displays the temporal intensity profile of a typical 80 fs pulse (FWHM).
after propagation inside a 1 mm thick Si plate. Fig. 2(b) shows the time-evolution of transmission at longer pump-probe delays for low (5 nJ) and high (85 nJ) pump pulse energies. From these data, we estimate that the stability of our measurement is ±3% of the signal but no transient evolution of the absorption is observed within this time scale. Among the processes that can lead to a decrease of the excitation density and by extension a decay in absorption, carrier diffusion is believed to play an important role due to the high electron diffusion coefficient $D_e$ of semiconductors. Using a mobility $\mu_e = 1400 \text{ cm}^2/\text{V} \cdot \text{s}$ associated to intrinsic Si at room temperature, the Einstein relation gives $D_e \approx 36 \text{ cm}^2/\text{s}$. At the maximum probe delay $t$ of 130 ps, limited by the length of the delay line used, we can predict that free electrons will have diffused over a typical distance $L = \sqrt{D_e \times t}$ exceeding 0.7 $\mu$m that is comparable to the radial dimension of the plasmas (3 $\mu$m, see inset in Fig. 1). However, our measurements do not exhibit any significant decay showing that a simple free-carrier diffusion description does not hold for these microscale plasmas.

In Fig. 4(a), we show the dependence of the probe transmission on the pump energy. For the measurement, the pump-probe delay is fixed at 10 ps so that we ensure the plasma is fully established and we measure the transient absorption level that decays at much longer time scale (see Fig. 2(b)). Raising the energy, we see that the absorption front is shifted towards the pre-focal region (see arrow to guide the eyes). As a result, the interaction volume increases while the minimum transmission rapidly drops down to a saturation level of $T \approx 80\%$. This may be better appreciated on Fig. 4(b) where the plasma transmission at the theoretical focus of the pump is plotted as a function of the pump energy. For energies exceeding 30 nJ, the transmission remains constant within the range of stability of our measurements.

The measured transmission level in the time-resolved images allows us to estimate the generated free-carrier density $N$ in the solid. According to the Drude model, the complex refractive index $\tilde{n}$ can be expressed as $\tilde{n}^2 = n_0^2 - \frac{\omega_p^2}{\omega^2 + i\Gamma}$, with plasma frequency $\omega_p = \sqrt{\frac{Ne^2}{\varepsilon_0 m_e}}$ where $\omega$ is the laser frequency, $e$ is the electron charge, $\varepsilon_0$ is the vacuum permittivity, $m^*$ is the effective electron mass ($m^* = 0.18 m_e$ for low-density plasma in Si), and $\gamma$ is the collision frequency. We choose $\gamma = 0.3 \text{ fs}^{-1}$ corresponding to a rate dominated by electron-hole collisions, which is reasonable for intermediate to low excitation densities. A sensitivity analysis to this parameter gives less than 50% variation in the density estimation using $\gamma$ in the range 0.12–0.5 $\text{fs}^{-1}$ corresponding to its upper and lower limits. Then, assuming a homogeneous plasma, the transmission $T$ is described by a simple Beer-Lambert law, $T = \exp\left(\frac{-4\pi Im(\tilde{n})}{\lambda} d_{\text{plasma}}\right)$ where $Im(.)$ denotes the imaginary part, $\lambda$ is the probe wavelength (1.3 $\mu$m) and $d_{\text{plasma}}$ is the measured plasma diameter (3 $\mu$m).

Under these assumptions, the 80% minimum transmission $T_{\text{min}}$ measured (Fig. 4(b)) corresponds to an increase of the imaginary part of the refractive index from $<10^{-6}$ to $7.7 \times 10^{-3}$ at 1.3 $\mu$m and a free-carrier density $N_{\text{max}}$ of 3.1 $\times 10^{19}$ cm$^{-3}$. This reveals a limitation of the excitation to an underdense plasma with density one order of magnitude below the critical density $N_{\text{crit}}$ at 1.3 $\mu$m ($6.6 \times 10^{20}$ cm$^{-3}$) consistent with the impossibility to induce local breakdown inside Si with single ultrashort pulses.

One obvious contribution to this limitation lies in beam energy depletion by TPA before the focus. The sensitivity of our diagnostic allows us to observe that the plasma develops in a pre-focal region for pump pulse energies from ~40 nJ (see arrow in Fig. 4(a)). In addition, it is worth noting that the pump pulse energy has been in most of the measurements above the threshold for self-focusing. According to the nonlinear refractive index in Si$^{13}$ ($\kappa_{NL} = 3 \times 10^{-14}$ cm$^2$/W), the critical power for self-focusing$^{16}$ is only 24 kW corresponding to a pump energy $E_{\text{pump}} \approx 1$ nJ for 80 fs pulses. However, we did not observe any significant focus shift or deviation from Gaussian focusing in the images. Moreover, the spectrum of the transmitted pump did not exhibit any significant change. Taken together, these observations indicate that nonlinear propagation effects remain limited and the limitation of excitation is likely associated to intensity clamping by TPA.

To connect the plasma characteristics to the absence of permanent material modification, one can turn to energy considerations. Assuming that each free electron is generated by a single ultrashort pulse, one can turn to energy considerations. Assuming that each free electron is generated by a single ultrashort pulse, one can estimate the deposited energy $E_{\text{dep}, \text{tot}}$ as $E_{\text{dep}, \text{tot}} = E_{\text{pump}} \times (1 - T_{\text{pump}})$ as it is shown on the same graph. For

FIG. 4. (a) Probe transmission as a function of the pulse pump energy and the depth inside Si. Pump-probe delay is fixed at 10 ps. Dashed arrow follows the shift of the absorption front. Zero depth corresponds to 400 $\mu$m below the Si surface. (b) The transmission at the center of the nano-plasma is plotted as a function of the pump pulse energy (dotted line on Fig. 4(a)).
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FIG. 5. Normalized transmission of the pump pulse as a function of pulse energy (right axis) measured for the same experimental conditions as for the imaging experiments. The corresponding total deposited energy is plotted on the same graph (left axis).

In conclusion, space-time control of the dielectric permittivity is the first step towards 3D material processing in c-Si. Our experiments show local modification of the imaginary part of the refractive index. However, we observe considerably low absorption levels independently of the incoming pulse energy. Moreover, our time-resolved study of this absorption reveals very slow carrier kinetics and no significant recombination within the 130 ps after the laser excitation. This reveals strong limitations when considering bulk excitation of Si by the use of single NIR ultrafast Gaussian pulses. While we have concentrated on measuring the inherent characteristics of the microplasmas, we will focus a future paper on conditions for improved energy confinement. Next, natural step will be to explore how our observations translate using tighter focusing conditions. We envisage also that the formation of microplasmas, similar to that in the inset in Fig. 1, can serve as a strongly absorbing media for efficient energy deposition in Si with a second synchronized pulse as it is proposed for ultrafast dielectric interactions.20

This research has received financial support from the French National Research Agency (ANR 2010-JCJC-913-01), the French Carnot Star Institute (Via-LASER), and the CNRS PICS Program No. 45052. A.R. is grateful for partial support by the Australian Research Council’s (Discovery Project DP 120102980) and by the Air Force Office of Scientific Research (USA, Grant No. FA 9550-12-1-0482).

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