Quantitative assessment of femtosecond laser-induced stress waves in fused silica

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(Received 15 February 2024; revised 6 August 2024; accepted 9 August 2024; published 26 August 2024)

Controlled processing of transparent materials with ultrashort laser pulses requires a detailed and precise knowledge of the various laser-matter interaction mechanisms from laser energy deposition and conversion in internal energy of the material to hydrodynamic relaxation and mechanical response. To address this issue, we first develop a multi-timescale experimental approach based on a femtosecond pump-and-probe microscopepolariscope. The pump is a 360-fs, 1-µJ infrared (1030-nm) laser pulse which is split to provide a 515-nm femtosecond probe with delay adjustable from femtoseconds to nanoseconds. The obtained time-resolved shadowgraphy images allow measurement of the transient probe transmission. Then, the carrier density is determined by use of the Beer-Lambert law and Drude model approach, demonstrating the ultrafast formation of a slightly overcritical plasma inside the bulk of fused silica. In parallel, quantitative birefringence images provide a measurement of stress by use of the photoelastic law revealing the partial dissipation of the absorbed laser energy through emission of a gigapascal stress wave, a few hundreds of picoseconds after the laser pulse. A multiscale-multiphysics model is then used to interpret the experimental observations, accounting for the electron dynamics, laser propagation, and hydrodynamic response. Following experimental validation, simulations allow determining the temporal evolution of local fundamental material properties (stress, density, and temperature). This combination of experimental and simulation results enables us to quantitatively discuss the importance of the different laser energy relaxation channels in the material, uncovering the full scenario of the interaction. Our model predicts a 20-GPa maximum initial stress loading with maximum lattice temperatures reaching. 3.5×10^4 K. We also show that a fraction of $\sim 2\%$ of the total absorbed laser energy is dissipated through emission of a weak shockwave. Our approach could be used in the future to account for mechanically driven transparent material structuring induced by ultrashort laser pulses.

DOI: 10.1103/PhysRevB.110.054112

I. INTRODUCTION

Femtosecond lasers allow reaching high optical intensities (in the TW/cm² range) with modest nanojoule laser pulse energies. At such intensities the interaction of the laser field with matter is highly nonlinear, allowing processing of transparent materials with unsurpassed quality [1,2]. This unique feature facilitates three-dimensional writing inside transparent materials, serving as a foundation for the microfabrication of monolithic microsystems. Applications include optofluidic biochips [3], optical memories [4] and other lab-on-a-chipbased devices [5]. Equally important are applications of fundamental interest, such as laser-driven polymorphic transitions of materials [6], generation of high-order harmonics in transparent solids [7], and laser amplification in photoexcited materials such as fused silica (SiO₂) and sapphire [8].

The above-mentioned applications result from a complex, multiscale, and multiphysical interaction of femtosecond laser pulses with matter [9,10]. Starting at the femtosecond timescale, the intense laser field is responsible for the photoionization of band-gap materials such as fused silica. Then, the generated free-carrier population linearly absorbs the laser energy during the laser pulse by inverse bremsstrahlung. At

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this stage, inelastic scattering of high-energy electrons may multiply the free-carrier density by impact ionization which in turn enhances optical absorption of the remaining laser pulse energy. At the same time, the laser propagation is modified due to variations of the dielectric permittivity induced by electron excitation. The laser-heated electron population further relaxes, progressively determining on the subpicosecond timescale a Fermi-Dirac distribution centered on a few-eV average temperature in such operating conditions. While electron transport is largely inhibited in dielectrics, the energy transfer from the electronic subsystem towards the lattice locally sets in, this process being complete at the picosecond timescale. Later, from subnanosecond to nanosecond timescales, hydrodynamic relaxation takes place including shock and/or stress wave emission and heat diffusion, with a characteristic time of heat energy fully diffused outside the focal volume at the microsecond timescale. Depending on laser irradiation conditions, the material ultimately finds its final state and may be irreversibly modified. At all stages, the dielectric permittivity of the medium is changed, enabling the use of optical diagnostic tools for observation and investigation of the interaction [11].

This phenomenological description demonstrates the difficulty to obtain detailed quantitative information about the interaction. The above-mentioned processes span and may occasionally overlap over multidecade spatial and temporal intervals. Optical experiments are often used for the investigation of these processes but they are limited intrinsically in terms of spatial (diffraction limit) and temporal resolution (pulse duration). Moreover, almost any modification of the electronic and lattice subsystems results in a change of its optical response, making it difficult to discern between individual mechanisms occurring simultaneously. To solve these issues, well-engineered experimental techniques must be employed to extract usable quantitative information. In parallel, numerical simulations based on modeling are used in conjunction with experiments to provide information on the hindered parts of the interaction. A validated model would be the cornerstone of a digital twin that would allow prediction of the interaction outcome and revolutionize laser-based manufacturing.

The pump-probe technique has been used successfully in transparent solids to capture ultrafast electron dynamics with femtosecond temporal resolution. Spatially resolved information may be obtained by combining a pump-probe scheme with microscopy tools. In this context, time-resolved pumpprobe shadowgraphy techniques have emerged as a solution for investigating fundamental questions in laser-matter interaction [12]. Several reports discussed transient photoinduced plasma inside the bulk of SiO₂ using optical transmission measurements [13,14], holography [15], or phase-contrast imaging [16,17]. Other studies explored the relaxation of the laser-deposited energy in terms of stress wave emission by the transient-lens method [18] or by polariscopic techniques [19]. In particular, important efforts were made to capture a continuum of experimental observations assessing both the laser energy deposition and material relaxation by Mermillod-Blondin et al. [20] and Hayasaki et al. [21].

Although many studies exist within a broad timescale, very few provide data sets with quantitative information on the evolution of the thermodynamic state of a material following irradiation by a femtosecond laser pulse. Recently, Nguyen et al. [22] provided quantitative measurements in the nanosecond-microsecond timescale, probing the thermomechanical relaxation of crystalline SiO₂ following irradiation by a femtosecond laser Bessel beam. In this study, we aim at bridging this gap by providing quantitative information on the interaction of an intense femtosecond laser Gaussian pulse with amorphous SiO₂ on a timescale ranging from a few hundred femtoseconds to several nanoseconds. This achievement allows us to uncover the time evolution of the physical quantities defining the thermodynamic state of the material upon laser processing with high precision from its initial laser excitation to its final transformation. By focusing a near-infrared 360-fs laser pulse inside the bulk of fused silica, we first achieve above-damage optical intensities, corresponding to \sim 10^{14} W/cm² in air. Different optical techniques are then combined to provide a continuum of direct experimental information in a large spatiotemporal domain. We perform time- and space-resolved optical transmission and birefringence measurements of the laser-excited material from the femtosecond to the nanosecond timescales in perpendicular geometry. By use of the Drude-Lorentz and photoelastic models, we deduce the laser-induced carrier density and stress amplitude, respectively. In parallel, we have developed a predictive model, accounting for the electron dynamics, laser propagation, and

hydrodynamic relaxation. The model has been experimentally validated and then used to address long-standing unanswered questions. These include, for instance, the transient pressuretemperature-density evolution of the irradiated material and the relative importance of the different energy relaxation channels, i.e., how the absorbed laser energy is transformed and dissipated into other kinds of energy in the material.

II. EXPERIMENT

A. Experimental setup

Figure 1 presents the time-resolved microscope-polariscope setup. A diode-pumped linearly polarized Yb-doped fiber laser (Satsuma, Amplitude Laser) delivers 20-µJ pulses with 360-fs duration [measured at full width at half maximum (FWHM)] and a central wavelength (λ) of 1030 nm. The laser beam is divided into two parts by use of a plate beam splitter (20% reflection, 80% transmission). The energy of the transmitted (pump) beam is varied by using a half-wave plate and a polarizing beam-splitter cube. Finally, the beam is focused with an aberration-corrected microscope objective (LCPLN50XIR, Olympus) of 0.65 numerical aperture (NA) inside the bulk of a fused silica sample. The sample is mounted on a three-axis motorized translation stage to allow precise positioning. The energy of the beam is set to 1 µJ after the microscope objective lens by using a calibrated power meter yielding a corresponding calculated peak focused intensity in air of $\sim 10^{14}$ W/cm² (waist $w_0 \sim 1 \mu m$).

The reflected beam on the plate beam splitter is converted to its second-order harmonic ($\lambda = 515$ nm) with the help of a type-I β -barium borate crystal. It is subsequently used for the illumination of the pump-irradiated volume (probe beam) in a Köhler-type perpendicular configuration. This choice enables the estimation of the optical integration length from direct experimental evidence. The pump-irradiated volume is then imaged by means of an objective lens (50× Mitutoyo Plan Apo NIR infinity corrected, NA = 0.42) and a CMOS camera (CS165MU Zelux, $3.45 - \mu m^2$ pixel size). The delay of the probe beam with respect to the pump is adjusted by changing the optical path of the probe with a motorized delay line, which is composed of four mirrors, within a range of about 5.5 ns. To acquire dynamic information over the whole range, single-shot irradiation is repeated each time with a different pump-probe delay and always on a fresh site of the sample at the same focusing depth. In overall, this configuration enables femtosecond temporal resolution (supposed to be equal to the probe pulse duration) and micrometric spatial resolution (0.6 µm according to the Abbe diffraction limit).

Birefringence measurements, being only sensitive to anisotropic changes, allow for reliable measurement of stress-induced modifications. Furthermore, the perpendicular configuration between the pump and probe pulses allows the measurement of the integration length, a crucial parameter in quantitatively determining stress-inducing birefringence. To perform birefringence measurements, a circular polariscope is implemented. The probe beam is first circularly polarized before the sample (polarizer P and quarter-wave plate QWPI in Fig. 1). It is analyzed after the sample using an elliptical analyzer consisting of a second quarter-wave plate (QWPII) and a polarizer (A) mounted on separate motorized rotation



FIG. 1. Experimental setup.

stages. Phase-shifting techniques can be used to retrieve the retardance of the sample [23]. We employ the Hecker-Morche arrangement [24], which yields a satisfactory compromise between accuracy and acquisition speed. To obtain a retardance map, five images are recorded with different analyzer settings corresponding to different orientations for QWPII and A. In between recordings, the sample is spatially shifted to always impinge on a pristine SiO₂ volume. One of the settings corresponds to the bright-field arrangement. It is simply the transmitted probe intensity in the absence of any analyzer. By dividing the transmitted probe intensity with and without the pump beam, we obtain the optical transmission. Finally, the signal-to-noise ratio is improved by systematically correcting for the background retardance of the sample and averaging over images. The accuracy of our setup has been calibrated by employing a variable Soleil-Babinet retarder.

B. Experimental results

The laser is operated in single-shot mode and the pump pulse is focused at a depth of 600 µm inside a polished SiO₂ cube (Heraeus Spectrosil2000). Pulse elongation due to chromatic dispersion is negligible due to low group velocity dispersion in SiO_2 at the pump wavelength. The intensity is maintained constant for all experiments and corresponding to a calculated peak optical intensity (in air) of $\sim 10^{14}$ W/cm². Using the above-mentioned pump-and-probe microscope-polariscope we record the isotropic (transmission) and anisotropic (birefringence) optical response of the sample upon laser irradiation. Time-resolved optical transmission and retardance measurements are presented in Fig. 2. In all of these images, the pump pulse is propagating from right to left (wavevector k_{pump}) while the probe pulse is propagating perpendicular to the screen towards the observer (wavector $k_{\rm probe}$).

The zero-time delay has been arbitrarily chosen as the beginning of the transmission drop due to the arrival of the pump pulse. Focusing on the transmission measurements at short delays [Fig. 2(a)], we observe a swift drop in the transmission at 200-fs time delay. This drop is attributed to free-carrier absorption previously generated by the pump beam due to photoionization. At this stage, we note that transmission measurements are sensitive to both attenuation and diffraction of the probe beam. When a phase object is created [Fig. 2(a)], diffraction generates a wavy modulation of the transmission. The transmission reaches its minimum approximately 600 fs after zero-time delay. Starting at 800 fs, we observe the rise of the transmission level again locally, indicating that the free-electron population begins to decrease in the focal volume with complete recombination occurring for nanosecond delays. At around 10 ps, a white spot appears locally at approximately the time when the transmission has been a minimum before (0.6 ps delay), indicating that transmission dynamics varies across the trace following variation of the deposited energy density. Transmission measurements within our conditions are affected by changes in both real and imaginary parts of the refractive index, making it difficult to discern between different contributions including the temperature rise. At ~ 1 ns, we have slightly modified the distance between the plasma and the imaging objective in order to better visualize the stress wave in transmission. This results in the sudden change of transmission measurement. After 2 ns from the arrival of the pump pulse we observe the formation and propagation of a stress wave that can be identified by the local change in the transmission [Fig. 2(b)). Postmortem transmission measurement on the irradiated sites [Fig. 2(e)] reveals a permanent modification of the material, demonstrating the use of an above-damage-threshold intensity in the experiment.



FIG. 2. Dynamic observation of the [(a), (b), (e)] optical transmission and [(c), (d)] retardance modifications induced by a 1- μ J laser pulse inside SiO₂. A free-carrier plasma is formed during the first hundreds of femtoseconds (a) followed by stress wave generation and propagation [(c), (d)] before an irreversible modification is observed at the end of the interaction [(e), (f)].

We obtain complementary results about the interaction by measuring the optical retardance. In Fig. 2(c), we present retardance maps for pump-probe delays from 0 to 1 ns. An increase of the birefringence signal is observed at around 400 fs inside the focal volume. Being more accurate than the transmission images to capture such information, the stress wave emission is detected earlier at 500 ps time delay and its outward propagation perpendicular to the optical axis is recorded until 6 ns. At late delays [Fig. 2(d)], this stress wave is propagating at a measured speed equal to the acoustic velocity of the material, approximately 5.72 µm/ns [25]. Experimental artifacts and limited spatial resolution hinder the determination of the exact shockwave position for short delays (<1 ns), thereby not excluding the presence of shockwave propagation at these delays. The optical retardance decays from 120 mrad to approximately 50 mrad within this time frame, revealing the decaying amplitude of the stress wave. Retardance measurements of the final modified structure reveals stress presence as illustrated in Fig. 2(f). A more detailed discussion on the precise nature of this wave will follow in Sec. IV.

III. MULTISCALE MODELING

As mentioned in the Introduction of this paper, a multiscale model describing the interaction of a femtosecond laser pulse with a dielectric material should include, first, the coupled microscopic electron dynamics and laser propagation and, second, the macroscopic hydrodynamic material response. Before presenting each part in detail, the synopsis and main assumptions of our multiscale modeling are briefly presented. First the laser energy deposition is determined by describing the coupled laser propagation and electron dynamics. The laser propagation is described by the nonlinear Schrödinger equation within the framework of the paraxial assumption. The laser reflection is thus neglected; the Kerr effect is also not included. The optical material response is evaluated with the standard Drude model which depends in particular on the free electron density. The latter is evaluated with a rate equation which accounts for photoionization, impact ionization, and electron recombination. Previous calculations provide the distribution in space of the energy density of free electrons just after the laser irradiation. Assuming this energy density is transferred into the lattice on a timescale much shorter than the material hydrodynamic response, the equivalent internal energy brought by the laser is used as an initial condition to carry out hydrodynamic simulations which also include the mechanical material response. Since hydrodynamics take place when there is no longer laser irradiation, a single temperature is assumed. Also, a one-dimensional cylindrical geometry is considered, which is close to experimental observations.

In our operating conditions using a 360-fs near-IR single laser pulse, the laser-induced electron dynamics is described with the standard rate equation providing the evolution of the free electron density, which has been shown to correctly account for various experimental observations [26]. In addition, such an approach is not computationally expensive (compared with *ab initio* calculations, for instance), allowing us to build a numerically efficient multiscale-multiphysics model well adapted for parametric studies. Within the present framework, the rate equation providing the evolution of the free electron density, ne, reads

$$\frac{dn_e}{dt} = \left(1 - \frac{n_e}{n_a}\right) \left(W_{PI} + \frac{\sigma_{col}I}{E_g}n_e\right) - \frac{n_e}{\tau_r},\tag{1}$$

where W_{PI} is the photoionization rate reading $\sigma_8 I^8$ with σ_8 the generalized eight-photon absorption cross section to bridge the band gap (value provided later). *I* is the laser intensity, E_g , the material band gap, is set to 9 eV [27], τ_r , the self-trapped exciton recombination time that should be accounted for in fused silica, is set to 150 fs [28], and n_a , the density of valence electrons that could be ionized, is set to 2×10^{22} cm⁻³ [29]. The term $(1 - \frac{n_e}{n_a})$ accounts for the depletion of the valence band. The absorption cross section, σ_{col} , obtained with the Drude model reads

$$\sigma_{col} = \frac{e^2 v_c}{\varepsilon_0 n_0 c m_e \left(\omega^2 + v_c^2\right)},\tag{2}$$

where ε_0 is the vacuum dielectric permittivity, e and m_e the electron charge and mass, respectively, n_0 the material refractive index set to 1.45, c the speed of light, ω the laser frequency, and v_c the electron collision frequency (value provided later). This description of the laser-induced free electron dynamics involves the following assumptions to build a computationally efficient multiscale approach. The contribution of self-trapped excitons to the ionization is not included and only multiphoton absorption is considered (no tunneling). Since there is an adjustable parameter for the photoionization process (σ_8), it can account implicitly for all processes. It is worth noting that, despite the nonlinear behavior of the process, we have observed a rather low sensitivity of simulation results with respect to this parameter due to a self-saturation mechanism (discussed in Sec. IV). This implies a low sensitivity to the precise nature of the ionization process. The electron transport is also neglected as usually done for dielectrics within the present physical conditions [30].

Experimental results shown in Fig. 2 demonstrate an elongated absorption area around the focal plane position, of length similar to the confocal length (calculated to be ~ 9 μm). This area appears rather homogeneous [see Fig. 2(a)], indicating that the absorption of laser energy develops in a large volume following beam propagation with negligible contribution from nonlinear propagation effects (not detected in our experiments). This smooth propagation of the laser beam is also supported by the use of (i) a small peak power in comparison with the critical power $(P_{\text{peak}}/P_{\text{critical}} = 0.56 <$ 1), (ii) a high focusing NA objective (limiting the length over which the nonlinear phase effects can develop), and (iii) the self-saturation intensity effect [31] that limits the intensity delivered in the bulk of a dielectric. We thus further assume that the spatiotemporal structure of the Gaussian laser beam is nearly unchanged in the course of interaction, which is in line with a free electron density lower than or close to the critical plasma density [32,33]. Under these conditions, the laser light reflection also remains low. The mechanical information further shows a moderate amplitude of stress waves (with moderate material damage), also suggesting a moderate laser energy deposition into the material. Within the scope of building a numerically efficient multiscale-multiphysics model, the previous conclusion from experimental observations leads us to model the laser propagation within the paraxial approximation instead of solving Maxwell equations. It is important to note that despite the moderate NA (0.65) used for the pump beam, refraction inside SiO₂ significantly lowers the vacuum equivalent NA down to ~0.45. This is smaller than the NA values leading to considerable nonparaxial and vectorial effects during the propagation [34]. This assumption will be also validated *a posteriori*. The axisymmetric equation driving the evolution of the laser electric field, *E*, then reads [26,32]

$$\frac{\partial E}{\partial z} = \left(\frac{i}{2k_0}\Delta_r - \frac{\sigma_{col}}{2}n_e - \frac{W_{pi}E_g}{2I}\right)E,\tag{3}$$

where z and r are the longitudinal (direction of laser propagation) and radial coordinates, respectively. Δ_r accounts for the diffraction within the cylindrical symmetry. The following terms are consistent with Eq. (1) and lead to both intensity attenuation in course of propagation and energy deposition into the material.

Figure 3 illustrates the coupled dynamics of electron population and laser field. Considering interaction in vacuum (without any excited material), Fig. 3(a) shows that the laser intensity distribution in space is symmetric as expected, accounting for the free propagation of a Gaussian field. The coupling with the electron dynamics generated in the excited material breaks this symmetry as depicted by Fig. 3(b) which shows the distribution of absorbed laser energy density in space after the interaction. As already described in Refs. [26,33], the first part of the propagation in front of the focal point has a Gaussian shape following the laser beam distribution. This is consistent with the low intensity ahead of the focal point inducing a weak absorption and a small number of free electrons. High intensity is reached nearly at the focal point, yielding strong absorption and a large population of free electrons. This further leads to a significant decrease in the laser intensity, which then is not able to further ionize the material beyond the focal point, leading to the observed asymmetry.

Because the characteristic timescales of energy deposition and hydrodynamics are significantly different (a few hundred femtoseconds and a few tens of picoseconds, respectively), the energy deposition previously evaluated from the electron dynamics and beam propagation modeling is used as an initial source term for the hydrodynamic simulations. Those processes and simulations developing at a later timescale ($\gg 1$ ps), electron-lattice relaxation is assumed to be finished and thermal equilibrium is reached. Therefore, no two-temperature model is used in our approach and in the corresponding equations only a single temperature is considered. Furthermore, cylindrical symmetry is considered because the initial energy deposition has a large enough aspect ratio (elongated in the direction of laser propagation as observed experimentally). Finally, these simulations are carried out in the radial direction with the Esther code which solves Euler equations (hydrodynamics) including mechanical material behavior (deviatoric component of stress) and heat diffusion [35]. SESAME table 7387 defining the equation of state (EOS) of fused silica including the various states of matter is used and the mechanical behavior is assumed to be elastoplastic [36,37]. The elastic limit and the shear modulus



FIG. 3. (a) Intensity profile without interaction; it is symmetric with respect to the propagation direction z. (b) Spatial profile of absorbed energy density in the material as predicted by the present model; the symmetry is broken (see text). Note that the horizontal axes are slightly different to exhibit the change in symmetry. The laser pulse is propagating from right to left towards negative z.

are set to 7.1 and 22.6 GPa, respectively. Both tend smoothly to zero at the softening point set to 2.15 kJ/g.

The coupled Eqs. (1) and (3) are numerically solved with a finite difference scheme. Note that both excited electron density and laser electric field depend on space and time. The electron density is initially set to zero everywhere, and the laser electric field is propagated from a Gaussian field as boundary condition (with a waist of 1 μ m at the focal point). We have checked that numerical convergence is achieved in terms of space and time discretization and domain size. The obtained two-dimensional energy deposition is averaged over the direction of laser propagation to provide the initial condition of hydrodynamic simulations, i.e., a radial profile of internal energy.

Considering the widely used value of photoionization cross section (order of magnitude of empirical expression of Agostini and Petite [38]) and the inverse of collision frequency in the standard femtosecond range $(1/\nu_c = 3.7 \text{ fs} \text{ for critical plasma [39]})$, we correctly reproduce the magnitude of absorption and amplitude of stress waves in the gigapascal range. More precisely, the microscopic modeling parameters have been set to precisely account for the experimentally measured maximum stress amplitude at t = 1 ns which is close to 1 GPa, determining $\sigma_8 = 10^{-270} \text{ m}^{16} \text{ s}^{-7}$ and $\nu_c^{-1} = 4 \text{ fs}$.

Simulations of the macroscopic material response can then be performed. Figure 4 shows the obtained spatial profile of the principal stress difference, $\Delta \sigma = \sigma_r - \sigma_z$, with σ_r and σ_z the principal stresses in cylindrical coordinates for suitable comparison to the experimental observation. The stress wave can be clearly observed in a nanosecond timescale consistent with experimental observations (after 500 ps, two symmetric structures appear). Profiles for a subnanosecond timescale are provided in Sec. IV. The propagation of a stress wave clearly appears. The positions of the wave are in a good agreement with experimental observations, with a velocity of \sim 5.5 µm/ns which corresponds to the sound speed, c_s . The compression wave (positive stress) is followed by a rarefaction wave (negative stress) which originates from the very early stress relaxation associated with the formation of a cavity in the interaction region [36,37]. The final important

characteristic of this dynamics is the attenuation of the wave in the course of propagation, decreasing from 1.5 to 0.25 GPa at the end of the observation, i.e., by a factor of 6. The processes at play are discussed in Sec. IV.

IV. ANALYSIS AND DISCUSSION

First, we compare the experimental results with simulations in order to validate our model in a large temporal interval. From the probe transmission results of Fig. 3(a), it is possible to calculate the average absorption coefficient $a = (2\omega/c) \times \text{Im}(n)$ at the center of the energy deposition volume, with *n* the refractive index of the excited material. We consider that the free carriers are homogeneously distributed in a cylinder. Then, to calculate the transmission perpendicular to the long axis of the cylinder we simply use the Beer-Lambert law, neglecting reflectivity:

$$T = e^{-al}, (4)$$



FIG. 4. Theoretical spatial evolution of the stress at various times. The wave propagation can be observed. The initial time corresponds to the end of the laser energy deposition into the material.

with *l* the traveled distance inside the cylinder equal to its diameter. At 600 fs, we extract the minimum transmission level at around 30%. The diameter of the cylinder can be estimated from a vertical transmission profile (perpendicular to the pump-pulse propagation axis). Taken as equal to the FWHM, we measure 1.6 µm. Finally, using the Drude-Lorentz model (see Sec. III), we calculate the carrier density $n_e = 3.80 \times 10^{21}$ cm⁻³. This value is approximately equal to 3.6 times the critical density at the pump wavelength $n_{\rm critical} = 1.05 \times 10^{21}$ cm⁻³, demonstrating the generation of a highly absorbing, slightly overcritical plasma, consistent with the observed permanent damage of the material.

Numerical simulations presented in Fig. 3 confirm the previous analysis. By averaging in time and space, we obtain an electron density of the same order of magnitude. It is worth mentioning that the paraxial approximation used to model the laser propagation is not strictly valid within such a condition but forms an acceptable approximation. As a demonstration, we recall that the theoretical shape of absorbed energy is in very good agreement with the plasma observation [Fig. 3(b) can be compared with Fig. 2(a) since absorption and transmission are mainly proportional to the free electron density]. By analyzing numerical results, the laser energy deposition into the material can be evaluated. We obtain roughly 250 nJ. Since the incident laser energy is 1 μ J, that corresponds to a total absorption of 25%.

On the other hand, birefringence measurements allow estimating the laser-induced stress inside the material. The photoelastic effect expresses that the dielectric permittivity is a function of stress and strain inside a transparent material. According to the stress-optic law, the optical retardance δ is linearly connected to the principal stress difference $\Delta \sigma$ with the relation $\delta = \frac{2\pi h}{\lambda} C \Delta \sigma$ [40], where λ is the probe wavelength, h the path length, and C the stress-optic coefficient equal to $2.5 \times 10^{-12} \text{ N}^{-1} \text{ m}^2$ for SiO₂ at $\lambda = 515 \text{ nm}$ [41]. Next, we consider that the shape of the laser-induced stress wave can be modeled by a hollow cylinder surrounding the focal volume with its height along the pump propagation direction. The difference between the outer and inner radii is equal to the optical thickness of the stress wave that can be accessed by plotting a vertical profile along the direction perpendicular to the optical axis of the pump beam [Fig. 2(d)]. Then, the path length is equal to twice the chord length that is tangent to the inner circle and can be easily retrieved geometrically. The expansion of the stress wave results in the increase of the path length in between different delays and it has to be accounted for to provide quantitative information about the generated stress. It is important to note that determination of the path length with a precision better than 10% is not possible in our conditions and that precise evaluation of the stress value should involve Abel transform algorithms that are out of the scope of this study. The retrieved principal stress difference as a function of the pump-probe delay is plotted in Fig. 5. The generated stress level is measured in the gigapascal range for subnanosecond delays (0.5–1 ns) before dropping down to the 100-MPa range at later delays (1-6 ns). At all times, the generated stress wave is measured to propagate at the speed of sound. This is in contrast with laser-induced shockwaves observed in air [42] or at the interface between



FIG. 5. Principal stress difference as a function of pump-probe delay. Experiment (black dashed line) and simulation (red dashed line) are shown. The decrease in the amplitude is due to geometric attenuation (see text).

 SiO_2 surface and air [43] where supersonic conditions are reached.

Modeling results in Sec. III indicate that the hydrodynamics proceeds in the acoustic regime within the present laser irradiation conditions for delays longer than roughly 500 ps. The maximum stress with respect to time can be extracted from these results. It is reported in Fig. 5 and directly compared with the experimental observation. There is good general agreement over the whole considered timescale. Since the stress wave evolves within the acoustic regime, no attenuation is expected (for shocks, the attenuation is mainly due to the rarefaction wave which meets the shock front). Viscothermal damping should also not contribute on this timescale [44]. The attenuation then is attributed to the spreading of acoustic wave energy in space. Within a cylindrical symmetry and propagation along the radial coordinate, the amplitude is expected to evolve as 1/r. With $r(t) = r_0 + c_s t$, a good fit of the maximum stress evolution is obtained (result not shown), confirming the geometrical attenuation. The possible main difference between experimental and theoretical results appears for the longest delays above 3-4 ns. For long delays, the wave propagation no longer obeys the theoretical cylindrical symmetry because its radial position is of the same order of magnitude as the length (along z) of the absorbing area, of the order of 15 μ m. The wave amplitude thus should decrease faster in this region, and as $1/r^2$ asymptotically (spherical expansion with a point source). That explains why modeling predicts larger amplitudes in this region.

Postmortem analysis indicates that within the observed discontinuous damage track [Fig. 2(e)] a dark scattering point appears, next a region with no transmission change, and again a region with lowered transmission. Similar morphologies have been observed by other authors [45,46] where dark points correlate with low-density regions revealed by phase contrast microscopy. The formation of voidlike structures has been the subject of several investigations, with their formation being attributed to either ultrafast formation from a



FIG. 6. Numerical simulations to exhibit physical processes on the short subnanosecond timescale. (a) Spatial profile of stress. The vertical arrow shows the position of the launched stress wave when it can be clearly distinguished from the stress background. (b) Both material density and temperature as a function of time on the central axis of laser propagation (r = 0).

rarefaction wave following microexplosion [47] or a much slower process in the presence of a phase transition where viscosity is considerably lowered, allowing cavity formation [48]. Furthermore, internal transformations towards densified structures have been already demonstrated [49].

The modeling is also expected to provide reliable information on physical quantities and processes in regions currently not accessible by our experiment, in particular how the absorbed laser energy transforms into other forms including mechanical deformations and hydrodynamics constraints on a subnanosecond timescale. Figure 6 provides such physical information, showing the evolution of stress, material density, and temperature as predicted by the present modeling. After the laser energy deposition and before hydrodynamic relaxation, Fig. 6(a) shows that the maximum stress is in excess of 20 GPa on the central axis of laser propagation (r = 0), i.e., more than one order of magnitude larger than the value observed at 500 ps. Discussing the calculated pressure of 20 GPa is essentially equivalent to discussing the calculated deposited energy density of approximately 150 kJ/cm³ [refer to Fig. 3(b)] as both are connected by the EOS. It is therefore important to highlight that this calculated energy accurately replicates the experimental stress measurements (see Fig. 5). While considering the potential experimental errors and the assumptions inherent in our model, an overestimation of this value cannot be ruled out, although its order of magnitude aligns with the expected physical processes. Specifically, this value is anticipated to be significantly greater than the melting enthalpy of fused silica ($\sim 5 \text{ kJ/cm}^3$) as indicated by the notable hydrodynamic response evidenced by the visualization of intense stress waves (see Fig. 2). Similar energy densities in the range of a few hundred kJ/cm³, with peak values around 300 kJ/cm³, have recently been reported by Rudenko *et al.* [50].

A strong and fast decrease in the pressure is observed in Fig. 6(a) and it is due to the formation of a cavity on a timescale of 100 ps as shown by Fig. 6(b), where the material density drops down to 0.2 g/cm³ (initial density is 2.2 g/cm³). The near-void formation is due to the central pressure which expels matter [36,47]. The pressure first drops because it is nearly proportional to the density (EOS) and because of energy transformation to kinetic energy: numerical simulations indicate that matter is expelled with a velocity in excess of hundreds of meters per second. Between 100 and 200 ps, the pressure profiles of Fig. 6(a) show that the short scale hydrodynamic relaxation stops and gives rise to the formation of a stress wave that can escape from the interaction region. Indeed, this timescale is the ratio between the size of the region in which significant laser energy deposition occurs, and the speed of sound, i.e., $\sim 1 \,\mu m/(5.5 \,\mu m/ns) = 180 \,ps$. The first instant where the stress wave can be clearly distinguished is indeed around 180 ps as shown by Fig. 6(a), in which the associated maximum stress is depicted by the vertical arrow. This delay is very similar to the one experimentally retrieved of a few hundreds of picoseconds [Fig. 2(c)]. It is interesting to evaluate whether an electronic contribution can be the main mechanism behind this pressure wave emission. However, in that case the wave would have been detected at earlier delays (~500 fs) when carrier density is at its maximum value. Furthermore, estimating the electronic pressure as $P_e = n_e k_B T_e$ with k_B the Boltzmann constant and T_e the average electronic temperature set to 5 eV, we obtain $P_e \sim 800$ MPa, which is negligible compared to the matrix pressure of ~ 20 GPa obtained from our model. Therefore, we attribute the origin of this wave to the matrix response following the laser energy deposition.

Figure 4 shows the propagation of this wave on a longer timescale. After a delay time of 2 ns [see Fig. 6(b)], the cavity closes because not enough energy was absorbed [36]. The origin of kinetic energy is first due to the matter movement associated with cavity formation and then is due to the stress wave. By analyzing numerical results, it appears that the kinetic energy associated with the cavity dynamics and the stress wave energy are rather similar, close to 5 nJ. Since the energy deposition is around 250 nJ, 2% of the absorbed laser energy is transformed into mechanical energy. The rest, 98%, is transformed into heat, with a maximum material temperature of about 35 000 K. Once again, the calculated temperature is linked to the calculated energy density of approximately 150 kJ/cm^3 via the EOS.

After 2 ns a long scale hydrodynamic relaxation takes place in the interaction region as shown by Fig. 6(b). The material density goes back to 2 g/cm³, i.e., not the initial density within these timescales. In the same time, the temperature drops due to diffusion and resolidification.

We finish this discussion on the possible formation of a laser-induced shock wave. When numerically measuring the speed of the wave after a few nanoseconds, we obtain $\sim 5.5 \,\mu$ m/ns which corresponds to the sound speed. Here the wave pertains to an acoustic wave. When measuring its speed right when it is launched from the interaction area (at t = 200 ps), we obtain 5.89 μ m/ns, i.e., slightly faster than the sound speed. This suggests the formation of a weak shock wave, which is consistent with a pressure of a few gigapascals corresponding to the upper limit of the acoustic regime. After a few microns of propagation, the pressure amplitude decreases, and the initial shock wave transforms into an acoustic wave.

V. CONCLUSIONS

In conclusion, we have investigated the interaction of a femtosecond laser pulse with SiO_2 from the early photoionization stage and energy deposition until its dissipation by means of stress wave emission. Our results bridge various physical processes taking place on different temporal and spatial scales. By focusing an infrared (1.03-µm) femtosecond (360-fs) laser pulse at above-damage-threshold intensity inside the bulk of SiO₂, we have generated an absorbing micrometric plasma. Time-resolved transmission microscopy was

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used to measure a maximum carrier density slightly in excess of the critical plasma density. Simultaneous time-resolved birefringence microscopy was used to directly measure gigapascal stress induced by the acoustic wave passage inside the material. A multiscale model with only two free microscopic parameters was developed to account for the laser pulse propagation, electron dynamics, and the macroscopic hydrodynamic response of the material. Following experimental validation, simulations provided additional insights into the interaction. Our numerical simulations suggest that the laser-irradiated volume is loaded with a maximum stress of 20 GPa and a temperature of 3.5×10^4 K. To obtain these results, we rely on the validation of the numerical simulations with experimental measurements therefore being potentially sensitive to experimental artifacts discussed in Sec. II B. A weak shock wave is emitted at around 200 ps that quickly degenerates into the experimentally observed acoustic wave. Terapascal generated pressures have been reported before to explain the formation of cavities by microexplosion [47]. However, recent theoretical investigations demonstrate that intensity clamping results in the achievement of more modest stress increases in the 100-GPa range [50]. Our approach could contribute to experimentally verifying such assertions in the future. Only 2% of the absorbed laser energy is transformed into a shock wave while the remaining energy is dissipated following a thermal pathway. In the future, our methodology can be used for further parametric studies such as laser pulse energy and number of pulses, providing a descriptive and reliable prediction tool for laser-based material processing.

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