Parity violation in chiral structure creation under femtosecond laser irradiation in silica glass?

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The paper addresses the creation of circular optical properties from a femtosecond laser light beam with a linear polarization in an achiral material (glass) under an orthogonal incidence. In this situation, all aspects of the experiment are achiral and therefore should not give rise to chiral property creation. From that observation, we propose an interpretation that involves the action of a light-induced torque on the matter carrying a light-induced dielectric moment. We found that a direct current (DC) electric field could be produced in the lattice by the femtosecond laser in our conditions and that a non-collinear dielectric moment is created by a nonlinear effect between the DC electric field and the stress field due to the transformation of the material. We reveal that it is possible to break the chiral symmetry of glass using an intense, ultrashort laser light pulse.

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**INTRODUCTION**

Femtosecond laser pulses focused in silica are absorbed through nonlinear photionization mechanisms; such mechanisms induce various modifications in the glass volume, depending on the laser parameters, such as refractive index changes¹, the formation of three-dimensional (3D) nanostructures², or voids³. Applications are numerous in a large number of domains, including health⁴, electronics, optical data storage⁵–⁷, optofluorics⁸–⁹ and optical components (for example, 3D optical waveguides¹⁰ or polarizing optical devices)¹¹–¹⁴.

Considerable research efforts are underway in the laser processing of optical materials because it is possible to achieve strong refractive index increase (a few $10^{-3}$ to $10^{-2}$ in a large range of transparent glasses) localized in the bulk (due to a strong nonlinear effect) and because the spatial resolution coincides with the requirement for waveguide elaboration (a few µm). It is thus necessary to understand how to control the interaction of femtosecond laser light with optical materials, in particular, with optical glasses, the most important of which being silica. If one proves to be able to control the forces involved in these processes, one could exceed the current applications of the lasers and open new possibilities in materials sciences.

In the past decade, several new phenomena have been discovered: nanogratings for high birefringence application with polarization direction dependence¹⁵, ‘quill’ writing¹⁶ with dependence of modification with the direction of writing, silica decomposition¹⁷, shear stress-induced field¹⁸ and an efficient creation of oriented anisotropic defects¹⁹. All these findings show that asymmetric printing in a centrosymmetric medium could be controlled²⁰. This concept of symmetry breaking (the Pierre Curie principle stipulates that the symmetry of the effects is always higher than the symmetry of the causes²¹) from nonlinear interaction of the infrared (IR) light appeared with the development of nonlinear optics and optical fibers. A famous example is the spontaneous appearance of second harmonic generation under IR irradiation²², which is normally forbidden in centrosymmetric media. Alternatively, it is conceivable that chiral light action in an achiral material can produce chiral modification. However, the efficiency is usually low or undetectable if an amplification process is not used²³.

In 2003, our group proved that femtosecond laser interaction can shear matter in a manner similar to a scissor cutting material, moving matter on one side in one direction and matter on other side in the opposite direction, depending on the laser scanning direction and on the femtosecond light polarization¹⁸. It was surprising to discover that an achiral beam (axially symmetric distribution of intensity, linearly polarized) penetrating into an achiral material (namely, pure silica glass) in an achiral geometry (orthogonal incidence) can give rise to a chiral strain if we take into account the light propagation axis. This observation was performed in the domain of the strain field (especially regarding shearing) and this strain does not give rise to optical circular properties by itself like the torsion can do in matter. However, in the present paper, it is demonstrated that another phenomenon occurring in the laser–matter interaction can actually lead to imprinting of chiral properties within a glass material. This reveals a parity violation (the reciprocity of the light propagation in the glass is broken) by efficient chiral structure creation. This chiral structure leads to the introduction of a chiral force (similar to the creation of torque) with a tightly focused femtosecond laser light in transparent solid material via the combination of nonlinear effects.

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This new technology enables the creation and the control of the circular properties, namely, rotating power and circular dichroism (CD). Currently, it is not possible to obtain similar properties in solids other than in elaborating single crystals. Such optical activity has a crucial role in optoelectronic devices, biological sensing and analytical chemistry. This new technology could provide a platform for a versatile new class of photonic devices. We can imagine a switchable plasmonic material based on chiral plasmonic nanostructures on glass surfaces (for example, DNA nanostructures decorated with metal nanoparticles) and a plethora of chiral photonic devices. More generally, this approach paves the way towards control of the orientation of the crystallization of the compound of interest, for example, the materials for second harmonic generation or for piezoelectric or ferroelectric effects, that is, second-order nonlinear materials.

MATERIALS AND METHODS
Femtosecond laser writing
The laser radiation, in a Gaussian mode, was produced by a regenerative amplified mode-locked Ti:sapphire laser operating at 100 kHz repetition rate that produces 160-fs pulses of light at a wavelength of 800 nm or by a femtosecond laser system (Satsuma, Amplitude Systèmes Ltd, Allée de Canteranne, Pessac, France) operating at 100 s of kHz and delivering pulses of 300 fs in duration at a wavelength of 1030 nm. Considering that the propagation vector $\vec{k}$ is along the $z$ direction, the beam was focused from 100 to 200 $\mu$m below the entry surface using a 0.5 numerical aperture (NA) microscope objective. The writing configurations defined by writing and polarization directions are shown in Figure 1. Therefore, when the laser was moving along $X$ (or $Y$) and the polarization was lying along $x$, it is defined as ‘$Xx$ writing’ (or ‘$Yx$ writing’). Next, moving the sample along the $X$ or $Y$ axis at a scanning speed of 500 $\mu$m s$^{-1}$, either single lines or 10 mm squares made of a group of lines were photoinduced in silica plates with various pulse energies ranging within a few fractions of $\mu$J per pulse and at different polarization directions (along the $X$ or $Y$ axis). The lines were written while paying attention to their direction (referenced to the laboratory) and orientation (for example, left to right and right to left). The irradiated squares were written following a linear scanning trajectory with a 1-µm pitch to achieve uniformly scanned region and eliminate any diffraction effects. Subsequently, some single laser lines were observed in transmission using a polarized optical microscope for detecting neutral axes, determining the slow axis orientation, optical rotation power and for measuring the linear birefringence via the Sénarmont method. Square samples were used to perform the CD measurements using a JASCO J-810 spectro-polarimeter (Jasco Company, rue des cerisiers, Lisses, France) equipped with a 50 kHz photoelastic modulator. Measurements were performed in the 185–700 nm spectral range with a 1-nm spectral resolution and with a 2-s integration time, resulting in an error of $< 1$ mdeg.

Measurement of CD in a substance exhibiting heterogeneous linear dichroism in depth but with transverse homogeneity
Usually, CD is measured in homogeneous substances having a center of symmetry but no circular symmetry, as in the case of a liquid. CD is measured by recording the light attenuation through the sample alternating left and right circular polarization using a photoelastic modulator. The attenuation difference normalized by the sum is called the ellipticity $\theta$ (expressed in mdeg). The relation with the CD is $\theta = (\alpha_1 - \alpha_2)2\epsilon$, where $\alpha$ is the absorption and $\epsilon$ is the circular property thickness.

In addition, in the case of a sample mixing uniformly linear and circular properties in depth, the isotropy is broken and we can suspect a dependence of the properties on the azimuthal position of the sample. The calculation shows that such mixing is not the case and the angular position of the neutral axes of the sample can be ignored.

In the case of a sample exhibiting a non-uniform distribution of linear and circular properties, any series of rotator and dephasor matrices can be reduced to a simple combination of one rotator ($R$) and one dephasor ($D$) matrices in arbitrary order, containing the equivalent rotation angle, the equivalent slow axis angle and the equivalent dephasing angle. In addition, in the case of linear properties much larger than the circular one, the equivalent orientation of the slow axis of the equivalent dephaser can be ignored and set to

![Figure 1](https://via.placeholder.com/150)

**Figure 1** Experimental setup scheme of the configuration for writing (from the left side) and schematic of a sub-wavelength periodic structure formed in a cross-section of the irradiated region. $n_1$ and $n_2$: local refractive indices of the nanoplanes; $t_1$ and $t_2$ are their respective thicknesses. Position 1 and 2 refer to light polarization. Reproduced from Desmarchelier et al. Copyright © 2015 by Desmarchelier et al. and Scientific Research Publishing Inc.
zero. It is thus possible to simulate the sample action as $R(\phi) D(\phi)$ through one face and $D(\phi) R(\phi)$ through the other face, arbitrarily. $\phi$ is the complex equivalent circular angle containing the circular birefringence ($\text{Im}(\phi)$, imaginary part) and dichroism ($\text{Re}(\phi)$, real part); $\phi$ is the same for the linear properties. In this case, the ellipticity $\theta$ follows $\tanh(\theta) = \text{tanh}(2\text{Im}(\phi))$ when incident on one sample face and $\tan(\theta) = \text{tanh}(2\text{Im}(\phi))/\cos[\text{Re}(\phi)]$ when incident on the other sample face. This result shows that the ellipticity difference between faces arises from the linear optical properties and that a non-zero ellipticity indicates the presence of CD.

From a practical perspective, this result shows that the light propagation is likely to be non-reciprocal in our samples, and thus, ellipticity measurements must be performed from both sides of the sample. Fortunately, the measurement is still independent of the azimuthal rotation of the sample.

RESULTS AND DISCUSSION
Femtosecond laser writing
Usually, in pure silica, at low exposure energy ($E<0.31 \mu J$ per pulse for laser polarization parallel to the scanning direction and 0.5 NA, 160 fs), one detects only a very weak linear birefringence of $<2 \times 10^{-4}$. The femtosecond laser interaction with silica is nearly ‘isotropic’ and is mainly due to the density changes that we attribute to an increase in the glass fictive temperature. On the contrary, beyond the T2 threshold, a strong linear birefringence appears. Measured retardance is approximately a few hundreds of nanometers for an interaction length of a few 10’s of $\mu m$. Thus, this birefringence is on the order of $10^{-2}$, similar to quartz. This birefringence was shown by Kazansky et al. to be related to the formation of nanoplanes of low refractive index. Recently, we established that nanoplanes are formed by the decomposition of SiO$_2$ into nanoporous silica. In the following, most experiments were performed in the laser parameter domain of existence of porous nanoplanes domain.

Rotated neutral axes
For this specific observation, single lines of glass modifications, separated typically by 100 $\mu m$ at least, were written in glass plates, as described in the section Femtosecond laser writing. These lines were then placed under a polarized optical microscope for examination in transmission and birefringence quantitative measurements. The samples exhibited strong linear birefringence on the order of 250 nm for a thickness of 40 $\mu m$, that is, $6 \times 10^{-3}$. From the symmetry of the writing and when the polarization is aligned or perpendicular to the written line direction, it is expected that one neutral axis aligned with the written line direction and the other one aligned perpendicular to it. However, it was surprising to observe that complete optical extinction was not possible between crossed polarizers. It was necessary to slightly uncross the analyzer, that is, the neutral axis turned a few degrees around the laser propagation axis. This departure angle depends on the writing sense in such a way that it is possible to vary the contrast of the line written in one sense over the others written in the other sense just by rotating the sample between crossed polarizers. This observation leads us to consider the optical rotating power.

Optical rotation
One of the sample used in the section Rotated neutral axes was used for measuring the optical rotation. The maximal extinction between crossed polarizers was searched, and the analyzer was then uncrossed for tentatively improving the optical extinction. We found an uncrossing angle of 0.5° for a 50 $\mu m$ (or less) active thickness corresponding to an optical rotation of $10^8$ mm$^{-1}$ at 632 nm; such an optical rotation adequate for practical applications. For this sample, the irradiation parameters were as follows: 1.5 $\mu J$, 1030 nm, 100 kHz, 0.6 NA, and 300 fs. Note that the rotating power for quartz is $\approx 21.73^\circ$ mm$^{-1}$ at 589 nm (Ref. 28). The sign is not defined because it is determined by the crystal arrangement, which can be levogyre or dextrogrye. No rotating power was detected for pulse energies below the porous nanoplanes formation. This result is the first piece of evidence that the femtosecond laser can render a glass chiral at the macroscopic level. In solids, optical chirality can originate from chirality at the molecular level and/or from a chiral arrangement of achiral motifs, as in quartz itself (which is, by the way, a crystalline phase of silica). The irradiated glass is thus linearly and circularly birefringent. In that case, some additional CD can be expected.

Spectral measurement of the CD
Here due to the size of the beam (0.8 cm), a series of close lines (1-3 um pitch) were written following a Peano function for filling a square of 1×1 cm$^2$. Recall that the quantitative CD in a solid sample that is linearly and circularly birefringent at the same time is not directly measurable. The quantity actually measured by the instrument is the ellipticity of the electromagnetic field (mdeg), which is defined as follows:

$$\theta = 2\text{tanh}(\text{Im}(\alpha), \cos[\text{Re}(\phi)]) = -(\alpha_R - \alpha_L)$$

The ellipticity is proportional to the CD $\alpha_R - \alpha_L$, where $\alpha$ is the linear absorption and $l$ is the length of the absorptive layer, only when the sample is isotropic, that is, in samples that do not exhibit either linear birefringence or linear dichroism. Otherwise, it is preferable to use circular anisotropic degree of attenuation (CADA) instead of CD, as it will be called in the following. Moreover, before performing these experiments, we must take into account that the linear and circular properties are now mixed in the same sample. If the circular properties are negligible compared with the linear ones by approximately three orders of magnitude, then they do not perturb the measurement of the linear properties. It is quite the opposite situation when measuring the circular properties. The reverse situation is of course not true, which represents some of the difficulties to overcome. As explained in the experimental details section, when linear and circular properties are not interrelated (that is, separated within the sample depth), the light propagation is no longer reciprocal. Thus, CADA must be measured from both sides of the sample; in addition, its real amplitude cannot be deduced without fitting and without knowing the quantitative values of the linear optical properties, namely, linear dichroism and linear birefringence.

For the data reported in Figure 2, a 160-fs pulse duration laser was used because this pulse duration allows a clear splitting between T1 and T2 thresholds. As shown in Figure 2, below the energy threshold for the appearance of porous nanoplanes and related strong linear birefringence, there is no or negligible CADA. On the contrary, for pulse energy above T2, an ellipticity appears. The sign of the CADA appears to be reversible with laser polarization, that is, positive when the polarization is perpendicular to the writing direction; however, it is not generally true, as the linear properties can modify this sign. Notably, the measurements are indeed non-reciprocal, as expected due to the linear optical properties. In Figure 3, we reported samples irradiated in the following conditions: 1030 nm 300 fs, 500 kHz, 500 $\mu J$ s$^{-1}$, 0.6 NA and 1 $\mu J$ per pulse (that is, above the T2 threshold). The CADA varies with the direction of the writing.
direction and polarization. The difference from one face to the other face arises from the linear birefringence and its spectral dependence when the linear and circular properties are not located at the same place in depth. The difference reaches a magnitude that is quite high compared with some organic molecules31, θ can reach 0.1° in the blue or ultraviolet (UV) range; however, in most samples irradiated under various laser conditions, it is at the level of a few tens of mdeg. This observation should be compared with the linear dichroism that reaches the range of 20 mdeg32.

Figure 4 shows that CADA depends on the laser pulse energy as well. At low energy of ~ 0.2 μJ, CADA does not depend on the sample face order through which the measurement is performed. In other words, linear birefringence is too weak to have a role here if the linear and circular properties are dependent on depth or if the linear and circular properties are not dependent on depth and at the same place. CADA disappears above 1.8 μJ for the laser conditions used. The scanning electron microscope images show that nanoparticles form clearly at 0.3 ± 0.1 μJ and disappear at 1.8 μJ along with substantial formation of damage. This observation indicates that circular properties are likely appearing in the same conditions as those for nanoparticles formation, that is, the type II regime.

Tentative interpretation
The measured results above establish that linearly polarized femtosecond laser beam entering an isotropic sample, a glass, under an axially symmetric geometry, is able to break the chiral symmetry of the glass. Notice that because a mirror exists in the middle of the line containing the laser propagation axis and the direction of writing, writing a line breaks axial symmetry but does not break chiral symmetry. This apparently violates the Curie principle resulting in parity violation, as stated in the paper title. Indeed, we discover here a phenomenon in which a material is predicted to be achiral but actually gives rise to a chiral property, for example, optical rotation or CD.

In determining the mechanism of such an unusual process, it must be first mentioned that chiral forces (shearing named as the scissor effect) have been proved to be relevant18,33. In addition, linear birefringence axes are expected to be aligned with the written line direction or the laser polarization ones (when it is perpendicular to the written line), which are the apparent symmetry axes of the written object; however, very often it is not the case. This leads to consider that a torque is applied by the light on the matter in the course of the laser direct writing process. Therefore, we are seeking the chiral force exerted on the material that can imprint a twist in the atomic arrangement, it is the case in the crystalline quartz or produce chiral defects from oxygen vacancies, for example. Notice that our observations are not related to the rotation in space of the nanogratings that has been recently reported by Dai et al.34,35; indeed, there is no chiral arrangement reported in these two references, as the images are symmetric in a mirror.

Figure 2 Circular anisotropy degree of attenuation (CADA) according to the probe wavelength, measured below and above the porous nanoplanes energy threshold T2. Laser writing conditions: wavelength: 800 nm, pulse duration: 160 fs, repetition rate: 250 kHz, numerical aperture: 0.55, writing speed: 500 μm s⁻¹, focal depth: 192 μm, energy: 0.07 μJ (below T2) and 1.3 μJ (above T2).

Figure 3 Comparison of spectral circular polarization degree of attenuation (CADA) for four configurations of writing and polarization direction from the two faces of the samples (Xy, Yy, Xx and Yx). Laser writing conditions: wavelength: 1030 nm, pulse duration: 300 fs, repetition rate: 500 kHz, numerical aperture (NA): 0.6, energy: 1 μJ per pulse, writing speed: 500 μm s⁻¹, focal depth: 192 μm.

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Our experimental results are explained as follows. Intense Gaussian femtosecond laser pulses focused in the material gives rise to multiphoton ionization until saturation in our conditions in the focal area. However, the electron density is not homogeneous due to the ponderomotive force exerted on the free electrons, as it is described by Bethune36 and considered again in Ref. 37. The ponderomotive potential $U_p$ is $\sim 10$ eV in our conditions38. However, notice that the CD is created by a significant pulse accumulation (1000 pulses per μm). Within the approximation of linear polarization, the force is as given in the following Equation (2):

$$\vec{f}_{fe} = \left( \vec{P}_e \cdot \vec{\nabla} \right) \vec{E} + \frac{\partial \vec{P}_e}{\partial t} \lambda \vec{B}$$

(2)

where $\vec{P}_e$ is the polarization of the quasi-free electrons only in the Drude approximation ($\vec{P}_e = \varepsilon_0 \chi_{fe} \vec{E}$ with $\varepsilon_0 \chi_{fe} = -\frac{n_e e^2}{m_e}$, $e$ and $m$ are relative to the quasi-free electrons, and $n_e$ is its density). We obtain

$$\vec{f}_{fe} = \varepsilon_0 \chi_{fe} \left( \vec{\nabla} \varepsilon_0 \vec{E} + \frac{\varepsilon_0}{\varepsilon_0} \vec{E} \times \vec{B} + \frac{\varepsilon_0}{\varepsilon_0} \vec{E} \times \vec{B} \right).$$

This force pushes the electron from the high-intensity area towards the edge of the laser beam. In the framework of the approximation used here, the force is sensitive to the pulse front tilt. Indeed, it has been shown by several groups that unusual effects, for example, asymmetrical writing of nanostructures or linear birefringence39–41, are dependent on the pulse front tilt occurring from the time compression process in the ultrashort pulse laser system. Our laser system exhibits a pulse front tilt oriented $36^\circ$ anticlockwise in the $xy$ plane and $0.064^\circ$ out of the $z$ axis for a 5-mm-wide beam before focusing, resulting in an angle of $\sim 60^\circ$ from the $z$ axis at the focus. Therefore, the electrons are not only expelled from the center of the beam towards the side but also the pulse front pushes some electrons to only one beam side in the manner of a ’snow-plough’, an effect previously described in Ref. 16. The electron density is thus higher on one side of the beam defined by the pulse front tilt only.

However, the experiment shows that the laser polarization has an important role in the orientation of the effect. To achieve such an effect, it is necessary to relax some of the approximations. The first approximation is the linearity of the matter polarization. In the dipolar approximation, the first non-zero term at pump frequency is the third-order polarization; this is the Kerr effect, but this one does not introduce any force component sensitive to laser polarization.
orientation. The next term at the pump frequency is the third-order quadrupolar term, given as follows:

$$P_{3Q} = e_0 \epsilon_0 \langle \hbar q \rangle |E|^2 \left( \frac{\partial B}{\partial t} + \frac{e_0 \epsilon_0 \langle \hbar q \rangle}{2} \left( \nabla \nabla \nabla \right) |E|^2 \right)$$

(3)

This term alters the free-electron density according to the laser polarization and gives rise to a non-centrosymmetric density, depending on the pulse front tilt and the laser polarization orientation.

At the end of the pulse, electrons relax and become trapped in the material in <1 ps (Ref. 42) resulting in a direct current (DC) field because the counter-ions are much heavier than electrons and are not displaced significantly by the electromagnetic field. There is thus a DC electric field applied in the material in the focal area at the same time as the solid temperature increased due to electron energy loss by electron–phonon coupling during the relaxation process. The solid temperature at this time is high enough for allowing this DC field to distort the ionized atomic network$^{43}$ and, probably, to partly screen the DC field. A stress field balances the other part. Next, >1μs later, another pulse arrived that has moved slightly. This second pulse changes the charge distribution everywhere except in certain places where the material (especially in the back of the beam displacement) cannot relax because the temperature increase is not high enough and the glass network distortion thus becomes recorded within the medium. Therefore, a stress field remains along the writing trajectory. This gives rise to unusual surface relaxation topography, as has been noted in our previous publications.$^{18,33}$

After the first pulse, the material is under stress that is a non-local property extending largely out of the irradiated area and rendering anisotropic properties onto the matter. As the silica photosensitivity is dependent on the stress field$^{44}$, the coexistence of a DC field and a stress field leads to a possible coupling between them. In such conditions, the material polarization after pulses must be written as follows:

$$\overrightarrow{P} = e_0 \epsilon_0 \langle \hbar q \rangle \overrightarrow{E}_{DC} + e_0 \epsilon_0 \overrightarrow{\sigma} \overrightarrow{E}_{DC}$$

(4)

where $\overrightarrow{\epsilon}$ is the fourth-order piezoelectric tensor and $\overrightarrow{\sigma}$ is the mechanical stress tensor. In isotropic media, $\overrightarrow{\epsilon} = \frac{1}{3} \overrightarrow{\epsilon} \frac{\overrightarrow{\epsilon}}{I} + 2 \frac{\overrightarrow{\epsilon} \overrightarrow{\epsilon}}{I}$, where Voigt notation has been used for the susceptibility components, $\overrightarrow{Tr(\overrightarrow{\epsilon})}$ is the trace of the stress tensor and $I$ is the identity tensor. Thus, the polarization can be written:

$$\overrightarrow{P} = e_0 \epsilon_0 \langle \hbar q \rangle \overrightarrow{E}_{DC} + e_0 \epsilon_0 \overrightarrow{Tr(\overrightarrow{\epsilon})} \overrightarrow{E}_{DC} + 2 e_0 \epsilon_0 \epsilon_0 \overrightarrow{\sigma} \overrightarrow{E}_{DC}$$

(5)

The first and second terms in the above expression give rise to the polarization contribution in the same direction as the DC electric field. On the contrary, as the stress tensor cannot in general be reduced to a scalar term, the third term introduces a polarization contribution in a direction that is perpendicular to the DC electric field. The misalignment between $\overrightarrow{E}_{DC}$ and $\overrightarrow{P}$ generates a volumic torque, that is, $\overrightarrow{P} \times \overrightarrow{E}_{DC}$. However, what are its direction and the sense of rotation? As previously stated, the recorded DC field is oriented in the direction of the pulse front tilt (here 36° anticlockwise in the x,y plane) deviated by the action of the laser polarization. In addition, the stress field deduced from the surface topography analysis reveals a shear stress that can be written as below in the written line reference containing

**Figure 5** Scheme showing the combined action of $E_{DC}$ and the stress field giving rise to a volumic torque at the end of the written line. $\overrightarrow{P}_{\perp}$ is the component of $\overrightarrow{P}$ perpendicular to $E_{DC}$.

This volumic torque may be responsible for the chiral atomic arrangements thought to underlie the present observations by cumulative effect. Thus, the control of the pulse front tilt or of a simple intensity gradient$^{44}$ and of the laser polarization according to the displacement should lead to the control of circular properties in a matter (pure silica glass here but in principle in any glasses), where such effect is not currently possible before irradiation. Femtosecond direct laser writing conjugated to pulse front tilt engineering is thus a new functionalization process of the glass that may be further extended to other optical materials.

**CONCLUSIONS**

We showed that the irradiation of an achiral material (pure silica glass) with achiral light (linear polarized Gaussian beam) from a femtosecond laser system can render the material chiral. However, the pulse energy must overcome the energy threshold related to the production of nanogratings giving rise to form birefringence. The circular birefringence (optical rotation) appears to be ~5° mm$^{-1}$, which is a result that has been noted in our previous publications$^{18,33,19}$. An example could be harnessed for the creation of CD in the UV–visible range. Moreover, from the theory developed above, we propose that the breaking of symmetry arises from the chirality of the set, including the DC electric field (defined by the pulse front tilt and the laser polarization), the direction of light propagation and the direction of scanning.

This new functionalization process could provide a platform for a versatile new class of photonic devices. With regard to biomimetics, we can imagine production of a cholesteric liquid crystal analogous optical devices using tiny lengths of inorganic glass, that is, ‘twisted silica glass’ rather than a macroscopic assembly$^{14}$. An example could be the production of ‘chiral fiber gratings’ that can be harnessed for producing sensors, polarizers, optical isolators and filter/lasers. Femtosecond laser processing thus offers a new advantage, partly in a non-conventional way: it allows restructuring of our most important optical material to enable chirality and rotating power. The important scientific point is that the pulse duration is so short that direct action on the lattice is not possible. However, the creation of the electric field creates a memory effect staying between pulses that may act on the material. Other experiments are planned to study the level of CD according to the pulse number and its spatial distribution along the laser traces. Last, we think that the use of high peak power and
ultrashort pulses is a powerful tool for the development of new applications in materials science for optics.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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