Generation, control and erasure of dual LIPSS in germanium with fs and ns laser pulses

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Abstract
Laser-induced periodic surface structures (LIPSS) can readily be fabricated in virtually all types of materials and benefit from an efficient parallel patterning strategy that exploits self-organization. The wide range of different LIPSS types with different spatial scales and symmetries is continuously growing, addressing numerous of applications. Here, we report on the formation of two fundamentally different types of LIPSS on germanium upon exposure to femtosecond laser pulses (\( \lambda = 800 \text{ nm}, 130 \text{ fs} \)), featuring different periods and orthogonal orientations. On the one hand, the well-known low-spatial frequency LIPSS (LSFL) with a period \( \approx \lambda \) and perpendicular orientation to the laser polarization are formed, which can be extended homogeneously in 2D by sample scanning. Additionally, extremely smooth ripples with a period \( \approx \lambda/2 \) and parallel orientation were generated at lower pulse numbers. We show that this new kind of ripples, named parallel high-spatial frequency LIPSS (HSFL-∥), can be superimposed onto LSFL by increasing the pulse number, forming complex dual LIPSS with nanohill-like morphology. While exposure to multiple nanosecond laser pulses is found to trigger also the formation of LSFL, HSFL-∥ cannot be formed under these conditions, which points out the role of ultrafast excitation in the formation of the latter. By performing time-resolved reflectivity measurements, we are able to resolve the melting and solidification dynamics, revealing melting of a very shallow surface layer (<20 nm) and melt durations of a few ns for both pulse durations pulses at the fluences employed for LIPSS formation. Finally, we demonstrate erasure of both types of LIPSS by exposure to single nanosecond pulses at high fluences, which paves the way for erasable multi-level data storage.

Keywords: laser-induced periodic surface structures, femtosecond laser, nanosecond laser, ripples, high-spatial frequency laser-induced periodic surface structures, low-spatial frequency laser-induced periodic surface structures

(Some figures may appear in colour only in the online journal)

1. Introduction
Laser-induced periodic surface structures (LIPSS) have gained considerable attention from researchers and industry due to their surface functionalization potential [1]. They can be produced by a single-step method, which enables efficient and flexible fabrication for numerous applications. LIPSS can readily be formed in metals [2], semiconductors [3] and dielectrics [4] by exposure to multiple short or ultrashort laser pulses. Their morphology strongly depends on many parameters [5–8] related both to the irradiation conditions [9, 10] (laser fluence, wavelength, polarization, repetition rate, environment, and processing speed) and material properties. LIPSS are typically distinguished by their spatial period. Low-spatial
frequency LIPSS (LSFL), with a period close to laser wavelength, are fairly well understood in most materials [11, 12]. Their origin is generally interpreted as the result of the interaction between the incident light and an electromagnetic surface wave; the latter being either scattered light or surface plasmon polaritons (SPP) propagating at the air/material interface. This interaction causes a periodic modulation of the spatial intensity distribution, in which processes like melting, ablation, or even oxidation [13] are triggered preferentially at the local intensity maxima, imprinting a permanent ripple pattern onto the sample surface after multiple laser pulses irradiation.

In contrast, for high-spatial frequency LIPSS (HSFL), having periods much shorter than the laser wavelength, large discrepancies are often observed between existing models and the experimentally measured features like period and orientation. The possible formation mechanisms of these ripples include interference along with transient changes in the optical properties during laser irradiation [14], surface second harmonic generation [15], excitation of surface plasmon polaritons [11] and self-organization [16].

The fabrication of LIPSS in semiconductors is of particular interest due to their dominant role in the electronics and photovoltaic industries, in which flexible and rapid nanostructuring strategies are desirable. It has been shown that features as small as 12 nm (λ/66) on silicon surfaces can be fabricated via a LIPSS-based process, employing irradiation with orthogonally polarized double femtosecond laser beams [17] while most of the published work has been concentrated on silicon [5, 14, 17–27], several works have been focused on germanium (Ge) [15, 22, 28–36]. One of the early works reporting LIPSS generation on Ge was published by Young et al., who employed ns pulses [22] to generate LSFL at different angles of incidence, reporting a strong angle dependence of their period and orientation. Virtually all of the later works on LIPSS in Ge are performed with ultrashort laser pulses. This strong change in the excitation conditions, generating higher electron densities, faster melting and thus steeper initial temperature gradients, might be underlying reason for the different type of ripples observed in these works. Le Harzic et al. [15], for instance, reported HSFL ripples in Ge oriented perpendicular to the laser polarization, with an average period of 160 nm upon irradiation with femtosecond laser pulses at 750 nm wavelength. Similarly, using fs laser pulses at 800 nm, Lin et al. [33] reported HSFL in Ge with a period close to 95 nm, also oriented perpendicular to the polarization and superimposed on LSFL with period of 500 nm, also perpendicular to the polarization. In contrast, Austin et al. [34] reported HSFL in Ge aligned parallel to the polarization with a period of 850 nm using mid-IR laser irradiation at 2400 nm and normal incidence. Borowiec et al. [23] observed LSFL in Ge only at a very long laser wavelength (2100 nm) and reported no evidence of HSFL or LSFL at 800 nm or intermediate wavelengths. However, the same authors reported LSFL and HSFL in GaP at 800 nm, with respective periods of 680 nm and 170 nm and perpendicular orientation to the laser polarization. The broad spectrum of different results illustrates the still limited understanding of the mechanisms and main parameters that control the triggering HSFL formation in Ge.

In this work, we report on the formation of two fundamentally different types of LIPSS in Ge by exposure to fs laser irradiation, featuring different periods and orthogonal orientations. Additionally, we have investigated the differences in the LIPSS morphologies induced by femtosecond or nanosecond laser pulses. Lastly, we demonstrate a simple method for LIPSS erasure by exposure to single nanosecond pulses.

2. Materials and methods

The material for studying ripple formation was commercial polished crystalline germanium <100> wafers (Ga-doped, 160 μm thick). For sample irradiation, a Ti:Sapphire laser system consisting of a mode-locked oscillator and a regenerative amplifier was used. The system delivered pulses of 130 fs duration at a central wavelength of 800 nm, at 100 Hz repetition rate. This system could also provide pulses in the ns regime with a temporal duration of 8 ns. The pulse energy was externally adjusted by means of a variable attenuator formed by a half-waveplate and a polarizing beam splitter. A mechanical shutter was used to control the number of laser pulses incident on the sample while a half-waveplate allowed to select the orientation of the linearly polarized laser light. The beam was sent through a circular aperture of 3.5 mm in diameter, before being focused on the sample surface by a lens with 150 mm focal length. A three-dimensional translation stage was used to move the sample and a rotating platform to select the incidence angle of the beam. Experiments were carried out at 0º and at 52º. The laser spot diameter was determined experimentally, following the method used by Liu [37], to be Gaussian with radial symmetry at normal incidence, having a 1/e² diameter of d = 59.0 μm. The fluence values quoted in the paper correspond to the peak fluence F, calculated as F = 2E/A, with E being the pulse energy and A = πd²/4. All experiments were performed in air.

For analyzing the temporal evolution of the surface reflectivity upon laser excitation, a real-time reflectivity setup, with a temporal resolution of 2 ns and a measured rise-time of a reflectivity change of ≈600 ps, was used [26]. The setup is composed of a continuous wave probe laser at a wavelength of 532 nm focused by a microscope objective at normal incidence onto the center of the pump laser-excited region, the spot size of pump laser is ≈1 μm, coinciding with the peak pump fluence. The reflected probe beam is sent to an avalanche photodiode, protected by a short pass filter from pump light scatter. The signal of the photodiode is displayed by an oscilloscope.

The surface morphology was analyzed by means of an optical microscope (OM) supplied with monochromatic light illumination at 460 nm and a field emission scanning electron microscope (SEM). The period of the LIPSS was measured by performing a 2D-FFT (fast Fourier transform) of the recorded OM and SEM images, employing a custom Matlab code. The surface topography was measured with an atomic force microscope (AFM) in tapping mode.
Figure 1. Optical micrographs of fs laser irradiated regions at different laser fluences $F$ (rows) and number of pulses $N$ (columns). The red arrow indicates the laser polarization direction. Each frame contains a magnification of a representative region that contains ripples. Note the appearance of a new kind of LIPSS oriented parallel to the polarization (HSFL-∥).

3. Results and discussion

3.1. LIPSS formation upon fs laser pulse irradiation

The Ge surface was irradiated at normal incidence with different combinations of laser fluence and number of fs pulses, thus obtaining individual processed spots. Figure 1 shows the OM images of the different resulting spots for selected combinations of parameters. The fluence $F$ was varied from 106–139 mJ cm$^{-2}$ and the pulse number $N$ from 50 to 300. For $F = 106$ mJ cm$^{-2}$ and $N = 75$, the irradiated spot features fine ripples oriented parallel to the polarization of the incident laser. Increasing the pulse number, these ripples become more pronounced for $N = 100$ and $N = 200$. At $N = 300$, it can be seen that the ripple orientation has changed, being now perpendicular to the laser polarization and displaying a strong optical contrast and a coarser spacing compared to lower pulse numbers. For a slightly higher fluence $F = 128$ mJ cm$^{-2}$ and $N = 75$, fine ripples appear just in an external ring of the irradiated spot whereas the central disk appears featureless. This behavior shows that ripple formation is limited to a small fluence window, namely $\Delta F = 22$ mJ cm$^{-2}$, similar to the case of amorphous-crystalline LSFL observed in silicon under similar processing conditions [24].

For $N = 100$, this external ring shows two kinds of ripples, fine horizontal ripples and coarser ripples oriented perpendicular to laser polarization, while the central disk remains featureless. Further increasing the number of pulses to $N = 200$, the horizontal ripples disappear while the vertical ones remain. Interestingly, in this case the central disc shows weak signs of vertical ripples extended from the outer ring, but with reduced contrast. For $N = 300$, the entire irradiated spot is covered with vertical ripples. For the highest laser fluence studied, $F = 139$ mJ cm$^{-2}$, the ripple evolution follows the same behavior: for low pulse numbers, the external ring contains horizontal ripples, which evolve upon pulse number increase into vertical or dual type ripples covering most of the spot.

In order to obtain high-resolution images of the morphology of the generated ripples and study their evolution with pulse number, we have performed SEM on selected spots. Figure 2(a) shows the image of a spot generated at $F = 128$ mJ cm$^{-2}$ and $N = 100$, featuring the characteristic outer ring that contains ripples oriented parallel to the laser polarization. The magnification of a ring region shown next to it (figure 2(b)) reveals that the ripples are extraordinarily smooth, which suggests that the self-organization process was dominated by melting, material displacement and re-solidification, involving no (or only minimal) material removal. This conclusion is supported by the very sharp transition in the horizontal direction from a non-modified region to the ripple region, without signs of an ablation rim, as it would be expected for ablative LIPSS. The ripple period was determined to be $\Lambda_\parallel = 400 \pm 50$ nm, half of the laser wavelength ($\Lambda = 800$ nm). Their maximum modulation depth is $d < 30$ nm, as it can be seen in figure 2(c), displaying an AFM map of a small region of the outer ring.

The above features have not been reported before for any type of ripples in semiconductors. LSFL typically reported in Ge and Si are always oriented perpendicular to the laser polarization and have a period close to the laser wavelength [5, 33]. The observed behaviour is also fundamentally
Figure 2. Transition from HSFL-∥ to dual LIPSS: SEM and AFM images of fs laser irradiated regions at \( F = 128 \text{ mJ cm}^{-2} \) for different pulse numbers \( N = 100 \) (a–c) and \( N = 200 \) (d–f). The red arrow indicates the laser polarization direction valid for all images. (b), (d) are magnifications of the white squares marked in the SEM images (a), (d). The white lines in the AFM images (c), (f) indicate the positions of the topography profiles plotted next to the images.

different from the recently reported amorphous-crystalline LSFL-type in silicon, featuring perpendicular orientation and exact-wavelength periodicity [24]. In this context, it is worth noting that we have performed micro-Raman measurements (excitation at 532 nm, spatial resolution 1 \( \mu \text{m} \)) on the irradiated regions in order to investigate if the amorphous phase of Ge might be present. Neither in the annular ring structure containing HSFL-∥ and LSFL nor in the central disk we could detect Raman bands, except the one corresponding to the crystalline phase (results not shown). Moreover, no Raman bands corresponding to GeO\(_x\) were observed, confirming that no surface oxidation was induced by the laser irradiation even at high pulse numbers. The latter result is relevant for classifying the ripple type, since several works have reported the existence of so-called LSFL-∥, attributed to so-called by radiation remnants. This is the case for instance in Ref [22], who reported LSFL-∥ in Ge upon ns-laser irradiation but only at elevated angles of incidence and resulting in periods larger than the laser wavelength. Another work reported LSFL-∥ imprinted at the interface of steel coated with a 100 nm thick oxide layer, attributing it to the scattering of the laser radiation at the nanoscale surface roughness and the propagation through a superfi- cial oxide layer [38]. The orientation of such structures is parallel to the polarization and the period scales as \( \Lambda \approx \lambda/n \), with \( n \) being the refractive index of the irradiated material. However, the absence of Raman bands corresponding to GeO\(_x\) in our case supports our classification of the ripple structures as HSFL-∥.

The characteristics of the ripples reported here also differ strongly from those of HSFL reported in Ge and Si, which typically show much shorter periods (\( \Lambda_{\text{HSFL}} \approx 100 \text{ nm} \) for Ge and Si [33]), perpendicular orientation with respect to the laser polarization (same as LSFL) [33, 39], and appear at higher pulse numbers than LSFL [33] rather than at lower ones. These fundamental differences to existing ripple structures reported, together with their particularly smooth morphology with almost no ablation, suggests the existence of a new kind of LIPSS, named hereafter HSFL-∥, obeying a different formation mechanism, on which we will shed more light in the following sections.

The influence of pulse number is illustrated in figure 2(d), displaying the SEM image of a spot generated at \( F = 128 \text{ mJ cm}^{-2} \) and \( N = 200 \). Here, well-pronounced vertical ripples, with a larger period are observed. Both, period \( \Lambda_{\perp} = 790 \pm 30 \text{ nm} \) and perpendicular orientation to the laser polarization are consistent with the well-known LSFL reported in semiconductors [5, 33]. Interestingly, upon magnification (figure 2(e)) it can be seen that the structures formed are rather complex and still contain the signature of the HSFL-∥ structures, forming dual LIPSS. In particular, a splitting of each individual vertical ripple is observed, effectively forming an array of nanohills, with a vertical spacing equal to \( \Lambda_{\parallel} = 400 \pm 50 \text{ nm} \) and a horizontal spacing of \( \Lambda_{\perp} = 803 \pm 20 \text{ nm} \). As it can be seen in figure 2(f), the maximum modulation depth of these dual LIPSS structures is considerably larger than for HSFL-∥ structures, namely \( d < 90 \text{ nm} \), which is qualitatively consistent with the high pulse number used that cause an increase in surface roughness and modulation depth [40].

A powerful method to improve fringe homogeneity and alignment, as well as to extend fringes in 2D, is to scan the laser spot over the surface at a well-defined repetition rate and
speed \[24, 25\]. Figure 3 shows regions of a surface irradiated while the sample was scanned in the horizontal direction under the incident fs laser beam operating at 100 Hz repetition rate, giving rise to laterally extended LSFL. Figure 3(a) shows highly homogeneous LSFL formed upon irradiation with horizontal polarization, yielding a period \(\Lambda_{\perp} = 800 \pm 30\) nm. Upon rotation of the beam polarization, the ripple orientation rotates as well and the period remains unchanged (figure 3(b)), although the ripple homogeneity clearly degrades. This fact demonstrates that sample scanning parallel to the polarization vector enables the formation of homogeneous and periodic grating structures.

As already reported for amorphous crystalline ripples formed in Si \[25\], this scanning strategy incubates a fringe structure along the scan direction before the actual laser spot passes, and thus triggers melting of periodic ripples, which greatly facilitates the formation of homogeneous, large scale periodic structures.

3.2. LIPSS generation upon ns laser pulse irradiations

In order to investigate the influence of the pulse duration on LIPSS formation, a similar study using different laser fluences and pulse numbers was carried out using operating the laser system at 8 ns pulse duration, still at 800 nm wavelength and normal incidence. In this case, the fluence range explored was \(F_{ns} = [149\ \text{mJ cm}^{-2} - 298\ \text{mJ cm}^{-2}]\) and the pulse number range was \(N_{ns} = [100 - 300]\), the results being shown in figure 4. Globally, it can be stated that for nanosecond pulse irradiation, LSFL aligned perpendicular to the polarization direction are mainly observed. Moreover, as the fluence is increased, the LSFL appear in an outer ring surrounding the of the spot in a similar arrangement as in the results obtained with fs laser pulses. Yet, in this case, the central disk is less homogenous compared to fs pulses, and shows clear signs of thermal effects, ablation and corrugation due to the longer pulse duration.

The fact that the lowest fluence \(F_{ns} = 149\ \text{mJ cm}^{-2}\) required to produce rippled structures is only 40\% higher than the one used for fs pulses (\(F_{fs} = 106\ \text{mJ cm}^{-2}\)) illustrates the importance of linear absorption of the laser light by the material, even for excitation with ultrashort laser pulses. The underlying reason is the relatively short penetration depth of the laser light in crystalline Ge \((L_{opt}(800\ \text{nm}) = 198\) nm). Despite the similar fluence and pulse number range for ns and fs laser pulses to form periodic LSFL, considerable differences were though observed in other aspects. Most notably, the absence of HSFL-\(\parallel\) structures for ns pulses, even when extending the parameter range (pulse number and fluence). This suggests the importance of ultrafast excitation and/or related non-linear effects in the formation of HSFL-\(\parallel\). Whether an indirect effect of ultrashort laser pulse excitation, the generation of a very steep temperature gradient and resulting faster melting and solidification dynamics, might also contribute to the formation of HSFL-\(\parallel\) will be investigated in the next section.

3.3. Time-resolved reflectivity measurements

We have investigated the temporal evolution of laser-induced heating, melting and solidification processes at the germanium surface upon fs and ns laser irradiation, employing time-resolved reflectivity measurements. Of particular interest to us was to determine the presence and lifetime of the molten phase in the fluence regime used to generate LIPSS structures. It has to be noted, though, that the fluence values used in this experiment cannot be compared directly to the results shown in figures 2–4, since the irradiation angle is different (\(\theta = 52^\circ\) for time-resolved experiments, \(\theta = 0^\circ\) for figures 2–4), and thus the absorbed energy of the p-polarized pulses. In order to identify the correct laser fluence for the time-resolved experiments, we have first performed a systematic study of static irradiations at \(N = 200\) and \(\theta = 52^\circ\), varying the laser fluence. By inspection of the fabricated spots by optical microscopy, we identified the spots that most that closely resembled those shown in figure 2 for fs pulses \((F(0^\circ,fs) = 106\ \text{mJ cm}^{-2})\) and figure 4 for ns pulses \((F(0^\circ,ns) = 149\ \text{mJ cm}^{-2})\). While considerable differences in the LIPSS structures were observed for the two different irradiation angles (as expected from the Sipe theory and reported in literature \[22\]), this method allows an identification of the optimum laser fluence for LIPSS formation at 52\%. The values obtained were \(F_{fs}(52^\circ) = 74\ \text{mJ cm}^{-2}\) and \(F_{ns}(52^\circ) = 137\ \text{mJ cm}^{-2}\), which were used for the time-resolved experiments shown in the following.

Figure 5 shows the evolution of the surface reflectivity (normalized to the sample reflectivity before irradiation) upon irradiation with single pulses at the two different pulse durations. For fs laser excitation, a rapid and strong reflectivity increase, limited by the rise-time of the detection system (600 ps), is observed, followed by slightly slower decrease, reaching quickly a reflectivity value similar to the initial value. This behavior is consistent with melting and solidification of a thin surface layer, since the reflectivity of the molten phase of Ge is much higher than that of the solid phase \[35\]. It can be seen that the measured reflectivity maximum \(\Delta R_{\text{max}}(\text{fs}) = 1.37\) is smaller than the calculated reflectivity increase for deep melting of Ge \((\Delta R_{\text{molten}} = 1.46)\). We have ensured that the expected reflectivity level for complete melting can be achieved when increasing the fluence only slightly, to \(F_{fs}(52^\circ) = 78\ \text{mJ cm}^{-2}\) (data not shown). This comparison
Figure 4. Optical micrographs of ns laser irradiated regions at different laser fluences $F$ (rows) and number of pulses $N$ (columns). The red arrow indicates the laser polarization direction. Each frame contains a magnification of a representative region that contains ripples. Compared to the results obtained with fs pulses, the most notable differences are the absence of horizontal HSFL-∥ and the appearance of thermal effects, ablation and corrugation in the central disk.

This result is important for the understanding of the LIPSS formation mechanism, clarifying that LIPSS formed upon irradiation with fs laser pulses emerge from a very shallow pool of molten Ge. This result is consistent with time-resolved measurements in Ge upon ns laser irradiation reported by Ehrlich et al [30], who found that efficient ripple growth occurs when the melt depth is less than 20 nm.

In the case of the ns laser excitation, the reflectivity response also displays a fast increase and decrease, consistent with the melting of a surface layer that is thinner than $d_{OTL}(532 \text{ nm}) \approx 20 \text{ nm}$ [28]. However, in this case the melting process is delayed with respect to the onset of the excitation pulse (included in figure 5 as a blue shaded curve at the bottom), starting at the trailing edge of the ns pulse, when most of the pulse energy has already been delivered to the material. Interestingly, the reflectivity rise is preceded by a reflectivity decrease, which is caused by heating of the solid phase, since the reflectivity of the latter is a decreasing function with temperature [26].

The melt durations for both pulse durations can be determined by measuring the time window in which the reflectivity stays above the reflectivity of the solid phase, yielding $t = 3.6 \text{ ns}$ for the fs laser pulse and $t = 6.4 \text{ ns}$ for the ns laser pulse. While these values are different, they appear to be too short for considering LIPSS erasure through surface tension, smoothing ripples on a long-lived melt. This scenario was proposed by [30] who estimated threshold values of $t = 200 \text{ ns}$ for this process to occur. Our results suggest therefore that the absence of HSFL-∥ for ns laser pulses is not a consequence of a long-lived liquid phase, underlining the above expressed hypothesis of an ultrafast, non-linear origin of HSFL-∥.
Figure 6. SEM images of fs laser irradiated regions before and after an erasure operation with a single nanosecond pulse. (a) fs laser irradiation with \( F = 142 \text{ mJ cm}^{-2} \) and \( N = 200, \theta = 0^\circ \). (b) after exposure to a single nanosecond pulse with \( F = 587 \text{ mJ cm}^{-2} \) at \( \theta = 52^\circ \). The red arrow indicates the laser polarization direction.

3.4. Erasing fs-LIPSS upon ns laser pulse irradiation

In view of their subtle surface morphology, small modulation depth and absence of ablation in the induced LIPSS by fs-laser exposure, the possibility of erasing them by exposure to nanosecond laser pulses was explored. The approach consisted in using a single ns pulse to locally melt a thin surface layer over a sufficiently long time to level out any ripple structures by exploiting the high surface tension of the molten phase. For this experiment, a series of spots with LIPSS were generated with fs pulses \( F = 142 \text{ mJ cm}^{-2} \) and \( N = 200 \) at normal incidence, as shown in figure 6(a). These parameters were chosen to generate a variety of structures, including dual LIPSS in the outer ring and LSFL in the central disk. In order to avoid resonant enhancement of these LIPSS by the nanosecond laser pulse (also able to trigger formation of LSFL as shown in the previous section) the erasure experiment was performed at 52\(^\circ\) angle of incidence angle. The significantly different angle of incidence ensures that, if present, a periodic intensity modulation due to interference of the laser light with a surface wave will have a very different period [24] and will not enhance the existing ripples. As can be seen in figure 6(b), upon exposure to a single pulse at a high laser fluence the complete LIPSS spectrum consisting in the dual-LIPSS in the outer ring and the LSFL in the central disk could be completely erased, leaving behind a darker but homogenous surface. The laser fluence of the ns pulse was chosen to lie below the single pulse ablation threshold of Ge under that conditions. This suggests that homogeneous melting and solidification of all fs laser-induced structures has been induced, erasing the entire LIPSS landscape due to the high surface tension of the long-lived molten phase, whose lifetime we have measured with the real time reflectivity setup to be \( t = 83 \text{ ns} \) (results not shown). These results point out the possibility to perform write-read-erase operations for data storage using fs dual-LIPSS in Ge. Multi-level, eraseable storage can be envisaged by data encoding in form of four different states, namely HSFL-∥, dual (HSFL-∥ + LSFL), LSFL, and no LIPSS.

4. Conclusions

We report on the formation of a new kind of LIPSS in Ge upon exposure to multiple pulse (\( N > 50 \)) fs laser irradiation, so-called HSFL-∥, in addition to the well-known LSFL. The main characteristics of HSFL-∥ are a sub-wavelength ripple period of \( \Lambda∥ = 400 \pm 50 \text{ nm} \), half of the laser wavelength and parallel orientation to the laser polarization direction. HSFL-∥ feature an extraordinary smoothness with a shallow modulation depth \( d < 30 \text{ nm} \), which suggests that the formation process involves melting, material displacement and resolidification, without material removal. By increasing the pulse number to \( N = 200 \), orthogonal LSFL emerge, superimposing onto the HSFL-∥ ripples, effectively forming dual LIPSS. The latter consist of nanohills arranged in a cubic structure, separated by the corresponding periods of the two LIPSS types. We also demonstrate the viability of suppressing HSFL-∥ ripples and homogeneously extending LSFL structures in 2D by sample scanning. Interestingly, LSFL can also be generated by ns pulses, at similar fluences and number of pulses as for fs pulses. However, this is not possible for the case of HSFL-∥ ripples, which are absent after ns laser exposure, which underlines the importance of ultrafast excitation to the formation of HSFL-∥. Time-resolved reflectivity measurements reveal melting of a very shallow surface layer (<20 nm) and short melt durations for both pulse durations. This result suggests that the absence of short-period HSFL-∥ for ns laser pulses is not caused by relaxation/smoothing of high-spatial frequency surface features through a long-lived melt. Finally, we demonstrate the viability of erasing fs-laser-induced dual LIPSS by exposure to single nanosecond pulses, which points out the feasibility of performing write-read-erase operations in dual-LIPSS in Ge for information storage applications.

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