Sub-wavelength terahertz imaging through optical rectification

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We record a sub-wavelength terahertz image of a caster sugar grain thanks to optical rectification in the sample excited with a femtosecond laser beam. The lateral spatial resolution of this technique is given by the laser spot size at the sample and here its measured value is 50 μm, i.e. ~λ/12. We give an estimation of the ultimate resolution that could be achieved with this method.

Production of images in the terahertz (THz) spectral range with a sub-wavelength resolution is nowadays commonly achieved through near-field techniques. Most of the setups employ time-domain methods with femtosecond laser excitation of the THz emitting devices, but images have also been obtained with continuous THz sources like QCL or even with synchrotron radiation, allowing the study of biological tissues. Such techniques are extremely efficient and permit to observe nanometer detail. On the other hand, they require dedicated instrumentation and know-how, which are not available in every laboratory.

Another solution is to generate, by optical means, the THz signal at the surface of the studied sample. This has been greatly demonstrated for years by Japanese teams, who have produced THz pulses by exciting electronic integrated circuits with a femtosecond laser. Carriers are photo-excited in the semiconductor substrate by the laser pulses and then are accelerated by any electric field along interconnections and components of the integrated circuit. These accelerated free carriers radiate a THz signal, which allows one to map the electric field at the studied device and for example, to detect any failure in the electronic circuit. Here the spatial resolution of this mapping is limited either by the distance over which carriers accelerate or by the laser spot size. Typically this technique, called laser THz emission microscopy (LTEM), exhibits a micron spatial resolution. It allows also evaluating the surface state of semiconductors or the electrical properties of semiconductor components. A related technique was employed to produce THz sub-wavelength images by Lecaque et al. Here the sample is put on the surface of a nonlinear crystal. The laser pump beam is focused in the nonlinear crystal in which it produces a THz signal by optical rectification (OR) and the OR-generated THz pulse goes through the sample before reaching the receiver. Thus the recorded THz intensity is modulated by the near-field transmittance of the sample. The image is obtained by scanning the laser beam over the sample and a λ/10 (30 μm) lateral resolution is demonstrated after a numerical deconvolution of the image. A method inverse to that of Lecaque et al. was published by Blanchard et al.; here the sample is put over an electro-optic crystal that is illuminated by a wide THz beam. The THz field, which is scattered and diffracted by the sample, is read by the probe laser beam that is tightly focused at the surface of the crystal. Thus, the THz transmission of the sample is measured in the near field and the authors report a λ/30 (14 μm) lateral resolution. Thanks to a very powerful THz beam generated by tilted-pulse-front excitation in LiNbO₃, images of 370 × 740 μm² samples are captured at a rate of 35 frames per second.

These two techniques, i.e. photo-carriers acceleration in semiconductors and nonlinear effects in crystals, differ notably because, in the first one, the sample serves as a source of THz radiation, while in the second one, the sample perturbs the emission or detection of THz waves in a nonlinear crystal. Here we propose to benefit from the advantages of both techniques, i.e. to employ optical rectification (OR) through or at the surface of the sample in view of getting a THz image with a sub-wavelength resolution. Thus the technique is not limited to semiconductor devices and it can be applied to the characterization of dielectric samples, as soon as they show a nonlinear optical response. As compared to the techniques by Lecaque et al. or by Blanchard et al., which deliver a signal proportional to the transmission of the sample at THz frequencies, here the recorded signal depends on both the sample absorption at visible and THz frequencies and on sample crystalinity through the nonlinear tensor. Indeed, when the sample material is not centro-symmetric, focusing a powerful beam at its surface induces second order optical nonlinear phenomena in the enlightened region, i.e. second harmonic generation (SHG) and OR. The SHG technique is widely spread in laboratory and hospitals and it delivers superb images that render for

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the sample crystalline inhomogeneity15. For example, membranes of biological cells are highly symmetrical due to the preferred orientation of their molecules and thus membranes produce a rather strong SHG signal that also depend on the membrane potential16, while some inner parts of the cells are liquid and thus are almost amorphous, leading to no SHG signal or at least to a weak one. Thus the proposed method is complementary to SHG imaging, since a THz photon is produced any time a SHG photon is. However, the efficiencies of OR and SHG are different, as recorded signals depend not only on the nonlinear tensor, but also on the susceptibility of the material respectively in the THz and visible ranges. Therefore, the two techniques should bring complementary information on the sample. It should be also noted that, even if the number of generated THz and SHG photons at the molecular level are the same and even without taking into account propagation and absorption effects, the THz beam should be weaker than the SHG one, because the energy of THz photons is about 1000 smaller than those of visible ones. In fact, OR was already used to study biological materials (bacteriorhodopsin)17: the intense impulse polarization excitation leads to a strong resonant infrared (IR) radiation from the molecules. Such IR light, whose wavelength is about 10 μm, is easier to detect than the THz one, because the IR photons are about 25 times more energetic than the THz ones. However, this result by Groma and colleagues17 demonstrates that OR could deliver information on the molecular response of the samples, even if the imaging purpose was not addressed in their work.

In this paper, we demonstrate the proof of concept of this technique by recording for first time a THz–OR image of a grain of caster sugar, taken from a packet of caster cane sugar (brown sugar) bought in a grocery. Sugar presents the advantage of being an efficient and very cheap nonlinear material, which has already been used for SHG18–21. Moreover, caster cane sugar grain are highly crystalline, with almost cleaved faces, even if they are not single crystals and contain impurities. Furthermore, we give the ultimate performance in terms of spatial resolution of this technique.

To our knowledge, the idea of THz imaging through OR was only published by a Japanese team22–24, in view of observing ferroelectric domains and especially domain walls in supramolecular ferroelectrics. Impressive images were obtained, but microscopy was not the main goal of this study. As compared to their work, we address here for first time any kind of material with the aim of mapping the crystalline structure of the sample through its nonlinearity and we discuss about the limitations (especially the spatial resolution) of this technique.

Results

Figure 1 shows a typical THz waveform and its spectrum obtained with a rather big grain (thickness ~0.8 mm). The mean laser power is \( P_{\text{laser}} = 3.65 \text{ mW} \) (3.65 μJ per pulse), corresponding to a peak power density \( D_{\text{laser}} = 2.8 \times 10^{12} \text{ W/cm}^2 \). The spectrum is maximum around 0.495 THz and spreads up to 5–6 THz where it reaches the noise level (about −50 dB). Some strong dips are seen around 0.9 and 1.7 THz. As sugar does not present absorption peaks at these frequencies, they may be due to water vapor absorption lines25,26 or to tiny rebounds of the laser pulse in the rubber tape where the sugar grain was stick.

Let us notice that the sample is almost transparent at the pumping laser wavelength and that we didn’t observe any two-photon absorption effect: therefore THz emission by photo-generated carriers can be ignored. However, to definitively demonstrate that the observed THz signal is originating from OR, we performed measurements of the polarization state of the generated THz beam. As expected, we found an angle dependency between the relative positions of the sugar crystal with regards to the pump laser polarization angle, which is the signature of OR.

The recorded spectra obtained with grains of different thicknesses or for different pump powers present similar results. This is shown in Fig. 2 where the THz power spectral density (log scale, with a dynamic range normalized to 1), is plotted for the largest and weakest recorded signals, generated with \( P_{\text{laser}} = 3.65 \text{ mW} \) (\( D_{\text{laser}} \sim 2.8 \times 10^{12} \text{ W/cm}^2 \)) and \( P_{\text{laser}} = 0.056 \text{ mW} \) (\( D_{\text{laser}} \sim 0.043 \times 10^{12} \text{ W/cm}^2 \)) respectively.

The weakest THz pulse energy is thus about 4200 smaller than the largest one. The spectra are rather similar, but the weakest one is of course noisier.

This allows us to record only the peak amplitude of the THz waveforms, which makes the image recording time much shorter. Therefore, the effective experimental frequency is the one of the maximum of the THz spectrum, i.e. ~0.5 THz. The OR-THz image of a sugar grain is depicted on Fig. 3. Points are recorded every 10 μm in both directions. At the center of the grain, the OR-THz signal magnitude is almost constant, while it decreases strongly at the crystal border.

Figure 4 presents a photography of the grain (left) and the superposition of the sugar crystal outline from photography (black dashed line) with the OR-THz 2D plot extracted from Fig. 3 (right). The THz signal is generated...
Figure 2. Normalized power spectra generated by OR in a sugar grain (red continuous line $D_{\text{laser}} \sim 2.8 \times 10^{12} \text{W/cm}^2$, blue dashed line $D_{\text{laser}} \sim 43 \times 10^5 \text{W/cm}^2$). Both spectra are normalized to a dynamic range equal to 1.

Figure 3. 3-D surface plot of the recorded THz OR image.

Figure 4. Image (left side): Photography of the studied caster sugar grain. Image (right side): THz OR image. The border of the grain sugar deduced from the photography, is shown as a black dashed line.
The radiated THz power $P_{\text{THz}} \propto \frac{\mu E_{\text{THz}}^2}{S_{\text{THz}}}$ of the THz field is proportional to the pump laser field power $P_{\text{laser}}$:

$$E_{\text{THz}} \propto \left| \chi \right| \propto \frac{\mu E_{\text{laser}}^2}{S_{\text{laser}}} \propto \chi E_{\text{laser}}^2 \propto \frac{P_{\text{laser}}}{S_{\text{laser}}}.$$  

(1)

Here $E_{\text{laser}}$ is the laser field magnitude (note that we use here only peak values of either the laser or THz pulses and not averaged values) and $S_{\text{laser}} \propto w_{\text{laser}}^2$ is the laser spot surface at the sample surface. The radiated THz power $P_{\text{THz}}$ is given by:

$$P_{\text{THz}} \propto S_{\text{THz}} E_{\text{THz}}^2 \propto S_{\text{laser}} E_{\text{THz}}^2 \propto S_{\text{laser}} E_{\text{THz}}^2 \propto \chi^2 \frac{P_{\text{laser}}^2}{S_{\text{laser}}^2}.$$  

(2)

To derive this expression, we state that, at the sample surface, the THz spot size is the same as the laser one, i.e. $S_{\text{THz}} = S_{\text{laser}}$ since the THz signal is generated by the illuminated area of the sample. Because this illuminated area

\[ \frac{P_{\text{laser}}}{S_{\text{laser}}} \]

is the laser spot surface at the sample surface. The radiated THz power

\[ \frac{P_{\text{THz}}}{P_{\text{laser}}} \]

is almost superimposed with the laser ones, of the sugar grain and thus to a reduction of the grain thickness nearby its borders.

The decrease of the THz signal at the grain border depends on the shape of the grain side, but also on the waist of the laser beam and on the diffraction of both laser and THz beams by the sample edge and thus it reveals the lateral spatial resolution of the measurement. To determine this spatial resolution, we have measured the waist of the laser beam at the grain surface location using a razor blade technique. The measured data (Fig. 5 left) are well fitted with a complementary error function, which attests that the laser beam is Gaussian and permits us to determine the laser beam waist (radius) equal to $w_{\text{laser}} = 30 \mu m$ (the Gaussian shape of the laser beam is plotted as a dashed line in Fig. 5 left). We have also applied the razor blade technique to the generated THz signal. The data are almost superimposed with the laser ones, i.e. $w_{\text{THz}} = w_{\text{laser}}$ which proves that the THz signal is only generated by the illuminated part of the sugar grain and that the THz beam is also Gaussian. This is confirmed by similar data (see Fig. 5 left) recorded using a perfectly flat ZnTe wafer, for which scattering and diffraction effects do not perturb the THz beam shape. Therefore, following the Rayleigh criterion, the lateral resolution of our record is $b_{\text{THz}} = 2w_{\text{THz}} \sqrt{i(2)} = 50 \mu m$.

Figure 5 right depicts the profile of the OR-THz signal versus the position along a line that crosses the grain s(a horizontal line roughly at the center of the grain shown in Fig. 5). As already seen in Fig. 3, the THz signal is practically constant at the center of the grain, which reveals that the grain thickness as well as the grain homogeneity are also regular. The decrease of the OR signal at the grain border occurs over a distance of 300–400 $\mu m$. This is not due to the spatial resolution of the experiment, since both laser and THz beams are much narrower (see continuous red line on Fig. 5 right): this is validated by performing the deconvolution of the OR profile, which does not show noticeable differences with the profile plotted here. This smooth decrease is thus attributed to the shape of the sugar grain and thus to a reduction of the grain thickness nearby its borders.

**Discussion**

Let us evaluate the limit of the spatial resolution in the OR imaging technique. We suppose that OR occurs at the surface of the sample and thus phase-matching effect may be neglected (in the present case, the sample thickness is more or less equal to the THz wavelengths, which justifies that propagation effects - phase-matching - may be ignored). As well, because we only derive order of magnitude, thus we forget about the tensorial behavior of the nonlinear phenomenon (the nonlinear susceptibility tensor $\chi$ is simply written as a scalar $\chi$). It follows that the magnitude $E_{\text{THz}}$ of the THz field is proportional to the pump laser field power $P_{\text{laser}}$:

$$E_{\text{THz}} \propto \left| \chi \right| \propto \frac{\mu E_{\text{laser}}^2}{S_{\text{laser}}} \propto \chi E_{\text{laser}}^2 \propto \frac{P_{\text{laser}}}{S_{\text{laser}}}.$$  

(1)

Here $E_{\text{laser}}$ is the laser field magnitude (note that we use here only peak values of either the laser or THz pulses and not averaged values) and $S_{\text{laser}} \propto w_{\text{laser}}^2$ is the laser spot surface at the sample surface. The radiated THz power $P_{\text{THz}}$ is given by:

$$P_{\text{THz}} \propto \frac{P_{\text{laser}}^2}{S_{\text{laser}}^2} \propto S_{\text{THz}} E_{\text{THz}}^2 \propto S_{\text{laser}} E_{\text{THz}}^2 \propto \chi^2 \frac{P_{\text{laser}}^2}{S_{\text{laser}}^2}.$$  

(2)

To derive this expression, we state that, at the sample surface, the THz spot size is the same as the laser one, i.e. $S_{\text{THz}} = S_{\text{laser}}$ since the THz signal is generated by the illuminated area of the sample. Because this illuminated area
is much smaller than the THz wavelength, the radiated THz field is almost a spherical wave. This wave is collected by a lens of section $S_{\text{lens}}$, located at distance $r$ from the sample. The collected THz is then focused onto the receiver. The THz power $P_{\text{THz,d}}$ impinging the receiver is:

$$P_{\text{THz,d}} \propto S_{\text{lens}} \frac{P_{\text{THz}}}{r^2} \propto S_{\text{laser}} \frac{P_{\text{laser}}}{r^2} \propto \chi^2 \Omega_{\text{lens}} S_{\text{laser}} D_{\text{laser}}.$$

(3)

$\Omega_{\text{lens}}$ is the solid angle subtended at the sample source by the entrance aperture of the THz optical system and $D_{\text{laser}}$ is the laser power density at the sample surface. It follows that the laser waist $w_{\text{laser}}$ at the sample can be expressed as a function of $P_{\text{THz,d}}$ and the nonlinearity $\chi$ and $D_{\text{laser}}$:

$$w_{\text{laser}} \propto \sqrt{\frac{P_{\text{THz,d}}}{\Omega_{\text{lens}}}} \frac{1}{\chi D_{\text{laser}}}.$$

(4)

Two limits are imposed in the OR experiment: (1) the power $P_{\text{THz,d}}$ at the receiver must be larger than the noise equivalent power NEP and (2) the power density $D_{\text{laser}}$ must be smaller than the sample material damage threshold $D_{\text{damage}}$. Therefore, in this OR experiment, the laser beam can be focused to a minimum waist given by the following expression:

$$w_{\text{laser,min}} \propto \sqrt{\frac{\text{NEP}}{\Omega_{\text{lens}}}} \frac{1}{\chi D_{\text{damage}}}.$$

(5)

Because the THz light is generated in the sample area illuminated by the laser, the THz lateral resolution $\delta_{\text{THz}}$ is proportional to the minimum laser waist $w_{\text{laser,min}}$. As expected, when $D_{\text{damage}}$ is high, the laser beam can be strongly focused while keeping the laser power constant, resulting in a better THz resolution ($\delta_{\text{THz}}$ decreases). As well, using highly sensitive THz detectors (NEP small) like superconducting devices (hot electron bolometers, kinetic inductance detectors, etc.) may improve the THz signal detection threshold by 2 ~ 4 orders of magnitude27 and thus it would lead either to a sub-micron resolution, or to the study of fragile samples, like biologic tissues. However, because of the square root in Eq. (5), the resolution depends more on the sample damage threshold. Also, if the sample is highly nonlinear (large $\chi$), the resolution is better. It follows from relation (5) that the spatial resolution depends on both the sample ($D_{\text{damage}}$, $\chi$) and the setup (NEP, $\Omega_{\text{lens}}$). In the present experiment, we observe damage in the sugar sample for $D_{\text{laser}} \sim 3.2 \times 10^{-12} \text{W/cm}^2$ and we measure both the OR profile and the THz beam waist with $D_{\text{laser}} \sim 2.8 \times 10^{-12} \text{W/cm}^2$. We selected this pump power density in order to have a large measurement dynamics and actually the generated THz signal (power) was about 1600 times larger than NEP. Therefore, the reported 50-$\mu$m resolution could have been decreased by a factor $\sqrt{1600} = 40$, i.e. down to a few $\mu$m, at the expense of a strong lack of dynamics.

About the depth resolution of the technique for a bulk sample ($d \gg \lambda$), we can simply estimate it from Gaussian optics. Typically, when using a focused pump beam, the nonlinear signal is generated in a material region whose thickness is equal to the Rayleigh length $Z_R$ ($Z_R = \pi w_{\text{laser}}^2 n_{\text{laser}} / \lambda$), where $n_{\text{laser}}$ is the refractive index of the nonlinear material. For thin samples, the depth resolution is the minimum among $Z_R$ and the crystal thickness. In the present work, $Z_R \sim 5 \text{mm}$, therefore the depth resolution is equal to the crystal thickness. It is why the profile given in Fig. 5 right is directly proportional to the sugar grain thickness profile, assuming the material is homogeneous.

In conclusion, we have demonstrated that sub wavelength THz images can be obtained by OR in the sample. Here, while testing a sugar grain, we achieve a $\lambda/12$ lateral resolution. We also give an estimation of the limits of this technique, in terms of spatial resolution which is determined by both the sample (damage threshold and nonlinearity) and the setup (collecting optics and detector sensitivity).

Methods

The studied grains are taken in a packet of commercial caster cane sugar (brown sugar). Caster cane sugar is made at 85–90% of sucrose and the rest (10–15%) includes fructose, glucose and impurities30. Sucrose belongs to the space group P21 (monoclinic)30. The size of the sugar grains is typically of the order of millimeters. We chose sugar grains that exhibit as flat as possible upper and lower faces, which appear as if they have been cleaved. The studied sugar grains are stick on a transparent rubber tape (Fig. 6). The setup includes an amplified femtosecond laser system (Coherent Libra) that delivers 5 mJ pulses of 50 fs duration at a repetition rate of 1 kHz. The central wavelength of the laser pulses is $\lambda = 800 \text{nm}$. The power of the pumping beam is adjusted with a half-wave plate (HWP) and a polarizer. In front of the sample, a second HWP permits to rotate the laser polarization. The detection of the THz signal produced by the sample is recorded with an electro-optic antenna, which consists of a 200-$\mu$m thick [111] ZnTe crystal, a HWP, a Wollaston prism and a balanced photodiode system. The parabolic mirrors used to collect the THz wave have a $2^\circ (\sim 5.1 \text{cm})$ diameter. The one close to the sample has a reflected focal length of 6° (15.2 cm) and the other ones 4° (10.2 cm). The THz signal was recorded with a lock-in amplifier (time constant: 1 s). We performed the measurements with spatial steps of 10 microns, taking in average 1 minute for each point. The total recording time of the whole image was around 3.5 hours.
Since all the sugar crystal axes directions are unknown, we changed the pump beam polarization angle by rotating the HWP situated in front of the sample, in order to maximize the detected signal. Then, we optimized the signal by rotating the HWP close to the balanced photodiodes. This allows to put the detection axis in the same direction of the THz electric field. Because the detecting ZnTe crystal is [111] cut, there is no need to rotate it to maximize the detected signal[31].

Data Availability
The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

References
1. Von Ribbeck, H. G. et al. Spectroscopic THz near-field microscope. Opt. Exp. 16, 3430–3438 (2008).
2. Adam, A. J. L. Review of Near-Field Terahertz Measurement Methods and Their Applications. J. Infrared Milli Terahz Waves 32, 976–1019 (2011).
3. Mitrofanov, O. et al. Near-field terahertz probes with room-temperature nanodetectors for subwavelength resolution imaging. Scien. Rep. 7, 42420 (2017).
4. Schade, U., Holldack, K., Kuske, P., Wüstefeld, G. & Hübers, H. W. Near-field imaging employing synchrotron radiation. Appl. Phys. Lett. 84, 1422–1424 (2004).
5. Schade, U., Holldack, K., Martin, M. C. & Fried, D. THz near-field imaging of biological tissues employing synchrotron radiation. In: Ultrafast Phenomena in Semiconductors and Nanostructure Materials IX. International Society for Optics and Photonics, 46–53 (2005).
6. Chen, H. T., Kersting, R. & Cho, G. C. Terahertz imaging with nanometer resolution. Appl. Phys. Lett. 83, 3009–3011 (2003).
7. Klarskov, P., Kim, H., Colvin, V. & Mittleman, D. M. Nanoscale Laser Terahertz Emission Microscopy. ACS Photonics 11, 2676–2680 (2017).
8. Kiwa, T., Tonouchi, M., Yamashita, M. & Kawase, K. Laser terahertz-emission microscope for inspecting electrical faults in integrated circuits. Opt. Lett. 28, 2058–2060 (2003).
9. Yamashita, M., Kawase, K., Otani, C., Kiwa, T. & Tonouchi, M. Imaging of large-scale integrated circuits using laser terahertz emission microscopy. Opt. Exp. 13, 115–120 (2005).
10. Yamashita, M. et al. THz emission characteristics from p/n junctions with metal lines under non-bias conditions for LSI failure analysis. Opt. Exp. 19, 10864–10873 (2011).
11. Nakanishi, H. et al. Imaging of a Polycrystalline Silicon Solar Cell Using a Laser Terahertz Emission Microscope. Appl. Phys. Exp. 5, 112301 (2012).
12. Murakami, H., Fujiwara, S., Kawayana, I. & Tonouchi, M. Study of photoexcited-carrier dynamics in GaAs photoconductive switches using dynamic terahertz emission microscopy. Photon. Res. 4, A9–A15 (2016).
13. Lecaque, R., Grésillon, S. & Boccara, C. THz emission Microscopy with sub-wavelength broadband source. Opt. Exp. 16, 4731–4738 (2008).
14. Blanchard, F. et al. Real-time terahertz near-field microscope. Opt. Exp. 19, 8277–8284 (2011).
15. Pavone, F. S. & Campagnola, P. J. Second Harmonic Generation Imaging (ed. Paul, J.) (CRC Press, 2014).
16. Thier, P., Denk, W., Sheves, M., Lewis, A. & Detwiler, P. B. Second-harmonic generation imaging of membrane potential with retinal analogues. Biophys. J. 100, 232–242 (2011).
17. Groma, G. I. et al. Resonant optical rectification in bacteriorhodopsin. Proceedings of the National Academy of Sciences 101, 7971–7975 (2004).
18. Kurtz, S. K. & Perry, T. T. A powder technique for the evaluation of nonlinear optical materials. Appl. Phys. 39, 3798–3813 (1968).
19. Rosker, M. & Tang, C. Evaluation of second-harmonic generation of various sugars in the powder form. IEEE J. Quant. Electron 20, 334–336 (1984).
20. Halbout, J. M. & Tang, C. Phase-matched second-harmonic generation in sucrose. IEEE J. Quant. Electron 18, 410–415 (1982).
21. Bourhill, G. et al. Powder second harmonic generation efficiencies of saccharide materials. Chem. Mater. 5, 802–808 (1993).
22. Sotome, M., Kida, N., Horiuchi, S. & Okamoto, H. Terahertz radiation imaging of ferroelectric domain topography in room-temperature hydrogen-bonded supramolecular ferroelectrics. ACS Photonics 2, 1373–1383 (2015).
23. Sotome, M., Kida, N., Horiuchi, S. & Okamoto, H. Visualization of ferroelectric domains in a hydrogen-bonded molecular crystal using emission of terahertz radiation. Appl. Phys. Lett. 105, 041101 (2014).
24. Kinoshita, T. et al. Visualization of ferroelectric domains in boracite using emission of terahertz radiation. Japanese J. of Appl. Phys. 53, 09PD08 (2014).
25. Van Exter, M., Fattinger, C. H. & Grischkowsky, D. Terahertz time-domain spectroscopy of water vapor. Opt. Lett. 14, 1128–1130 (1989).
26. Cheville, R. A. & Grischkowsky, D. Far-infrared foreign and selfbroadened rotational linewidths of high-temperature water vapor. J. Opt. Soc. Am. B 16, 317–322 (1999).
27. Soyu, G. et al. Detection of terahertz time-domain signals using kinetic inductance detectors. Submitted to Journal of infrared, millimeter and terahertz waves.
28. Yokota, H., Kaneshiro, J. & Uesu, Y. Optical Second Harmonic Generation Microscopy as a Tool of Material Diagnosis. Phys. Res. Intern. 5, 704634 (2012).

Figure 6. Experimental setup.
29. White, J. S. Sucrose, HFCS and Fructose: History, Manufacture, Composition, Applications and Production in Fructose, High Fructose Corn Syrup, Sucrose and Health (ed. by Rippe, J. M.) 13–33, (Humana Press, 2014).
30. Beevers, C. A., McDonald, T. R. R., Robertson, J. H. & Stern, F. The crystal structure of sucrose. Acta Crystallogr. 5, 689–690 (1952).
31. Gaborit, G., Sanjuan, F. & Coutaz, J. L. Second order nonlinear optical processes in [111] cubic crystals for terahertz optoelectronics. Lithuanian J. of Phys. 58, 24–37 (2018).

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Author Contributions
J.-L.C. proposed this study, F.S. performed the experimental work and F.S., J.-L.C. and G.G. analyzed the results and wrote the manuscript. All authors reviewed the manuscript.

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