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Third harmonic generation microscopy of cells and tissue organization
Bettina Weigel1, Gert-Jan Bakker1 and Peter Friedl1,2,3,*

ABSTRACT
The interaction of cells within their microenvironmental niche is fundamental to cell migration, positioning, growth and differentiation in order to form and maintain complex tissue organization and function. Third harmonic generation (THG) microscopy is a label-free scatter process that is elicited by water-lipid and water-protein interfaces, including intra- and extracellular membranes, and extracellular matrix structures. In applied life sciences, THG delivers a versatile contrast modality to complement multi-parameter fluorescence, second harmonic generation and fluorescence lifetime microscopy, which allows detection of cellular and molecular cell functions in three-dimensional tissue culture and small animals. In this Commentary, we review the physical and technical basis of THG, and provide considerations for optimal excitation, detection and interpretation of THG signals. We further provide an overview on how THG has versatile applications in cell and tissue research, with a particular focus on analyzing tissue morphogenesis and homeostasis, immune cell function and cancer research, as well as the emerging applicability of THG in clinical practice.

KEY WORDS: Multiphoton microscopy, Third harmonic generation, Nonlinear imaging

Introduction
The visualization of living cells within their tissue environment is a central endeavor and challenge in life sciences. Besides chemical and molecular labels to generate contrast of defined molecular structures, and to allow visualization of cells in three dimensions and over time, nonlinear microscopy enables label-free visualization of cell and tissue structures (Alexander et al., 2013; Ellenbroek and van Rheenen, 2014; Helmchen and Denk, 2005). Originally used in material sciences, the development of high-power lasers, optimized objective design and high-sensitivity photomultipliers with a wide dynamic range have recently enabled the application of third harmonic generation (THG) microscopy to three-dimensional (3D) cell and tissue microscopy, which complements broadly used fluorescence and second harmonic generation (SHG) detection. We here review the principles and live-cell applications of THG microscopy, which together with SHG and fluorescence detection enables powerful 3D visualization of the extracellular matrix, cell morphology and subcellular organization for multi-parameter analysis of cell–tissue interaction and function.

Physical principles of THG
Higher harmonic generation is a nonlinear coherent scattering process, resulting from phase matching and summation of light waves that are induced by inorganic and organic structures that have specific physical properties, molecular arrangement and order. SHG results from the conversion of two incoming photons into one emitted photon with doubled energy and thus, the emission of light of half the wavelength (Fig. 1A,B). SHG occurs at ordered non-centrosymmetric structures that are found predominantly in metals and crystals (Chen et al., 2008; Chu et al., 2002). Likewise, biological materials contain large-scale repetitive non-centrosymmetric units – including coiled-coil structures and polymeric proteins, which facilitate SHG signals, for instance at striated muscle and fibrillar collagen (Campagnola et al., 2002; Friedl et al., 2007). THG results from the tripling of the frequency of the excitation wavelength, whereby the combined energy of three photons is converted into one emitted photon with one third of the excitation wavelength and tripled energy (Barad et al., 2002; Friedl et al., 2007). THG occurs at structural interfaces, such as local transitions of the refractive index or the third-order nonlinear susceptibility $\chi^{(3)}$ (Barad et al., 1997; Müller et al., 1998), which are both intrinsic physical properties of materials and liquids. Therefore, THG is more versatile compared to SHG because it does not require a specific asymmetry of the structure to be imaged. In tissues, THG excitation occurs predominantly at interfaces that are formed between aqueous interstitial fluids and lipid-rich structures, such as cellular membranes (Aptel et al., 2010; Rehberg et al., 2011), lipid droplets (Débarre et al., 2006; Tserevelakis et al., 2014; Watanabe et al., 2010), or inorganic structures, such as calcified bone or enamel in teeth (Chen et al., 2008; Oron et al., 2004). In addition, THG can also be found at interfaces between water and large protein aggregates, such as collagen bundles or muscle fibers (Rehberg et al., 2011; Weigel et al., 2012).

Technical and experimental considerations of THG application
Determinants of signal intensity
The signal intensity of THG crucially depends on the size and organization of the signal-eliciting structures at the micrometer scale (Fig. 1E), the excitation and detection approach (Fig. 1C,D), and both power and polarization of the incident light. Although no signal is obtained in homogeneous dispersed medium, the signal is elicited by interfaces (Fig. 1E, examples 5 and 6) and optical inhomogeneities of length scales that are comparable to the focal volume of the laser beam (Cheng and Xie, 2002; Débarre et al., 2005) (Fig. 1E, examples 1–4). Consequently, the THG signal intensity is strongly affected by the size of the structure and the resolution of the system, notably the numerical aperture of the objective. Tuning of the focal volume can thus be used to optimize THG intensity for specific structures of interest. In Drosophila embryos for example, lightening the focus enhances the signal from the external membrane (a scenario comparable to that in Fig. 1E, example 6), whereas, when
using a weakly focused beam, signals predominantly originate from densely packed internal micrometer-sized organelles (Débarre et al., 2005; Mahou et al., 2011) (a scenario comparable to that in Fig. 1E, example 4). THG signal strength and origin further depend on the orientation of the interface within the focal volume, with an optimal THG signal generated by an interface orientation that is parallel to the excitation beam (Cheng and Xie, 2002; Müller et al., 1998) (Fig. 1E, examples 5 and 6). Lastly, THG signals are sensitive to the polarization of the excitation light (Oron et al., 2004); in human cornea, using a circularly polarized excitation beam allows the specific detection of interfaces of the lamellae, whereas a linear polarized excitation beam reveals stromal keratocytes and epithelial cell boundaries (Aptel et al., 2010). Thus, signal intensity strongly depends on local imaging conditions.

Excitation wavelengths
In principle, THG is generated at any wavelength of light. Practically, the optimal excitation wavelength is determined by a trade-off between tissue scattering and absorption. THG detection in the UV range is precluded owing to a strong scattering and absorption of UV light by tissues, and the lack of UV transmission by most optical glasses. The optimal excitation window is determined by a trade-off between tissue scattering and water absorption of the excitation light. Longer wavelengths reduce scattering, but light with a wavelength above 1350 nm is more strongly absorbed by water, which limits tissue penetration. By contrast, in brain tissue for example, wavelengths in the spectral excitation window of 1700 nm provide optimal THG imaging depths of up to ~1 mm (stratum pyramidale) (Horton et al., 2013).

In practice, infrared excitation (wavelength range 1180–1350 nm) and detection within the lower blue spectrum (390–450 nm) is best suited to generate and detect THG using with the same objective lens. Detecting THG in the blue range further has the advantages of minimum spectral overlap and convenient combination with commonly used cyan, green and red fluorophores, as well as SHG in the green–yellow range (compare examples in Figs 1 and 2).

Laser-power range
THG as a third-order process is strongly reactive to alteration of excitation power density in the focus – i.e. a twofold increase of laser power results in an eightfold increase in THG signal intensity. Thus, a modest increase in excitation power will substantially improve the THG signal. Conversely, even only a small loss of photon density in the focus will result in a substantial loss of THG signal. Further, signal intensity strongly depends on excitation pulse length and laser repetition rate (with both inverse second-order dependence), excitation wavelength (third-order dependence) and the numerical aperture of the objective (second-order dependence) (Cheng et al., 2014). In deep-tissue imaging applications, THG signals are thus strongly affected by light scattering and wavelength dispersion that is caused by out-of-focus structures and by increasing distortions of the focus in depth (Dietzel et al., 2014). The detection of THG in 3D tissues can be improved by using adaptive optics in order to achieve a sharp focus in light-scattering samples, which maximizes the signal intensity of SHG and THG (Thayil et al., 2011). Further, THG signal strength benefits from pulse predispersion compensation, to counteract pulse dispersion by the optical components and the sample itself (McConnell, 2006). Because of this nonlinearity and sensitivity to even small changes of imaging conditions, conclusions that are based on THG intensity could be problematic.
Phototoxicity

In contrast to fluorescence, THG is an energy-conserving process whereby the emitted photon contains the total energy of three exciting photons, and no energy is absorbed by the signal-eliciting structures (Squier et al., 1998). Therefore, THG causes neither photobleaching nor generation of reactive oxygen.
species. In addition, owing to the lack of out-of-focus excitation, any photodamage caused by THG imaging in 3D tissue samples is confined to the scanned volume. However, the high illumination power required for THG can induce tissue damage. Independent from the THG process, phototoxic molecules can be generated by concurrent two- and three-photon (auto)fluorescence excitation, reabsorption of THG emission or thermal damage through tissue absorption of excitation light (Débarre et al., 2014). These indirect effects typically require careful choice of excitation wavelength (typically 1200 nm and higher), because phototoxic effects are significantly reduced with longer wavelengths (Andresen et al., 2009; Chen et al., 2002). For imaging of zebrafish embryos, wavelengths in the range of 1200–1300 nm do not seem to generate photodamage and have been proven to be suitable for the repeated scanning that is required for long-term time-lapse imaging (Sun et al., 2004; Chu et al., 2003).

**Forward and backward detection**

According to the momentum conservation law, the generated higher harmonic photons are mostly emitted in the same direction as the initiating photons, thus the emission light propagates in a forward direction, away from the objective (Cheng and Xie, 2002) (Fig. 1C). Therefore, THG signals from transparent samples are best detected with forward detectors mounted on the side of the sample that is opposite to the exciting laser beam. In non-transparent samples, scattering of the THG signal allows backward (epi-)detection of photons, which are backscattered by the tissue (Débarre et al., 2006). Consequently, the intensity of epi-THG signals improves in dense scattering tissue and media where more forward-generated THG photons are backscattered (Débarre et al., 2007; Rehberg et al., 2011) (Fig. 1C). In addition, theoretical studies and experimental evidence of nanoparticles in aqueous solution suggest that a direct backward component of THG exists, in addition to the photons that are indirectly scattered from forward THG (Chang et al., 2010a). Generally, backward THG appears as a weak forward THG signal with a decreased signal-to-noise ratio, and the same structures can be detected (Rehberg et al., 2011; Tai et al., 2005). There are only a few exceptions of differing forward:backward intensity ratios for specific structures, such as striated muscle, nuclear chromocenters (Rehberg et al., 2011) and collagen fiber cross sections (our own observation). Because of the weakness of THG, particularly in detection of backward scatter, high-sensitivity detectors, such as gallium arsenite phosphide (GaAsP) photomultipliers are required.

**THG in biological samples, in comparison to multiphoton and confocal fluorescence imaging**

For use in life sciences, THG has several unique technical and application advantages. In contrast to multiphoton and confocal fluorescence imaging, THG is readily detected without the need for tissue staining and delivers a broad range of structural information on cell and tissue architecture. Similar to multiphoton excitation, and in contrast to one-photon confocal imaging, THG is only generated in the focal volume; therefore, the imaging process and signals that are detected are intrinsically confocal and suited for scanning of 3D samples and for the reconstruction of their volume. Confocal imaging is limited to the superficial cell layers of tissues owing to scattering, absorption and phototoxicity limitations, whereas the infrared wavelengths that are used for multiphoton and THG excitation allow long-term time-lapse imaging (Fig. 2C,D,F; Sun et al., 2004) and facilitate deep-tissue penetration (Andresen et al., 2009). Similar to optical coherence tomography (OCT) and ultrasound, but in contrast to other label-free imaging modalities – including coherent anti-Stokes Raman scattering (CARS) and autofluorescence – THG provides structural but not molecular information (Box 1). Thus, THG is complementary to other imaging approaches and is best applied within multi-parameter acquisition schemes.

**Box 1. Label-free imaging modalities in life sciences**

Label-free imaging modalities generate image contrast by exploiting intrinsic material, structural or chemical properties of the sample. In optical coherence tomography and microscopy (OCT and OCM, respectively), a 3D image is reconstructed based on variations in the time-delay and scattering pattern of the excitation light, which is reflected by tissue structures and interfaces. OCT creates contrast in a similar manner to that of ultrasound; however, by using light instead of the long wavelengths of sound or radio frequency, it achieves higher resolution. OCT provides images of large fields of view in real-time with several millimeters of penetration depth while typically maintaining lateral and axial resolution in the micrometer range. Similar to THG, OCT provides structural but not molecular contrast. When combined with Doppler imaging for label-free micro-angiography, OCT delivers powerful structure–function insight into, for example, blood flow (Drexler et al., 2014). Photoacoustic imaging uses excitation by laser light, which is absorbed by tissue and partially converted into heat; this leads to ultrasonic emission, which is detected for image reconstruction (Beard, 2011). Thus, the high contrast of photo-absorption is combined with the low scattering properties of ultrasound (Zhang et al., 2015). Optical absorption properties of tissue components vary with physiological conditions, such as hemoglobin concentration and oxygen saturation, and this can be exploited, for instance, for quantitative measurements of blood oxygen saturation. The current resolution of photoacoustic imaging is in the range of approximately 100 μm, but can be improved to cell-level resolution by combining photoacoustic imaging with microscopy (PAM) (Zhang et al., 2015). Compared to photoacoustic imaging, the penetration depth of PAM is reduced from several millimeters to less than 1 mm. OCT and OCM, ultrasound and photoacoustic techniques provide fast image acquisition, large imaging depths, but (supra)cellular resolution. In contrast, optical modalities such as two photon autofluorescence (AF), SHG, THG and coherent anti-Stokes Raman scattering (CARS) provide sub-cellular resolution with added structural and molecular specificity. SHG and THG detect frequency changes of the tissue-scattered excitation light, such as frequency doubling (SHG) and tripling (THG), which occurs at structures of particular asymmetry (SHG) or optical interfaces (THG) using a single laser beam. For CARS, a pump beam and a Stokes beam interact with each other and the sample in a wave-mixing process, and thereby enable molecular vibrational spectroscopy of the sample. CARS can distinguish tissue structures based on their chemical composition, such as DNA, lipid, metabolite or protein content (Evans and Xie, 2008). Compared to CARS, THG produces high-contrast images that are rich in detail with a much simpler excitation scheme and minimal risk for aberration artifacts that are induced by the mixing of two excitation beams. Conversely, CARS provides information on the chemical properties of cell and tissue structures. Alternatively, label-free molecular contrast can originate from tissue autofluorescence. By analyzing the spectroscopic properties of the emission – e.g. by using fluorescence lifetime imaging (FLIM) or spectral imaging – multiple signals originating from, for example, NADH, flavins, retinol, keratin, FAD, melanin and elastin can be separated and quantified (Fereidouni et al., 2014; Skala et al., 2007; Zipfel et al., 2003). Compared to THG, autofluorescence signals require shorter near infra-red excitation wavelengths and usually have a low signal-to-noise ratio, which limits detection in scattering tissues. Two-photon-excited autofluorescence, SHG, THG and CARS can be easily combined with multiphoton fluorescence microscopy on the same platform, using the same laser source. Each modality adds distinct structural and/or chemical context to the fluorescence data.
Table 1. Overview of tissue structures that can be visualized with THG, and imaging applications

<table>
<thead>
<tr>
<th>Visualized structure</th>
<th>Tissue</th>
<th>Biological application and comments</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extracellular matrix and inorganic structures</td>
<td></td>
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<tr>
<td>Collagen fibers and bundles</td>
<td>Mouse skin; mouse cremaster muscle</td>
<td>3D reconstruction of tissue organization and interstitial space; cell–matrix interaction and guidance; signal originates from the fiber surface, complementary to SHG</td>
<td>Rehberg et al., 2011; Weigelin et al., 2012; Wu et al., 2015; Fig. 2A,B,G</td>
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<td>Elastic fibers</td>
<td>Human conjunctiva</td>
<td>Signal originates from the fiber surface, complementary to SHG; shows disease-related proliferation of elastic fibers</td>
<td>Sun et al., 2007</td>
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<td>Basement membranes</td>
<td>Mouse cremaster muscle, ear, dermis</td>
<td>Visualization of delineating vessels and muscle fibers, likely to represent the surface signal that results from the interface between the basement membrane and stroma or adherent cells</td>
<td>Dietzel et al., 2014; Rehberg et al., 2011; Weigelin et al., 2012</td>
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<tr>
<td>Mineralized bone matrix</td>
<td>Mouse embryonic bone</td>
<td>Surface signal; visualization of mineralized bone matrix</td>
<td>Oron et al., 2004</td>
</tr>
<tr>
<td>Calcified teeth</td>
<td>Human teeth; fossil vertebrate teeth</td>
<td>Detection of cracks or lesions resulting from mineral loss; classification of fossil vertebrate teeth</td>
<td>Chen et al., 2008; Chen et al., 2015b; Kao, 2004</td>
</tr>
<tr>
<td>Cells, cellular membranes and cell organelles</td>
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<tr>
<td>Cellular membranes</td>
<td>Human cornea and skin; zebrafish embryo</td>
<td>Visualization of tissue architecture and abnormalities during disease; cell lineage tracing during embryo development</td>
<td>Apter et al., 2010; Chen et al., 2009; Tsai et al., 2014; Olivier et al., 2010a,b</td>
</tr>
<tr>
<td>Nuclear membrane, nucleoli</td>
<td>Mouse cremaster muscle, human oral mucosa and skin</td>
<td>Visualization of abnormalities during disease; nuclear morphology changes during cell proliferation and death; analysis of the nucleocytoplasmic ratio to detect malignant lesions</td>
<td>Rehberg et al., 2011; Tsai et al., 2011; Fig. 2G</td>
</tr>
<tr>
<td>Adipocytes</td>
<td>Mouse cremaster muscle, dermis, human epicardial adipose tissue</td>
<td>Identification of the 3D organization and interfaces present in fat tissue, used to understand cell invasion into fat tissue, exploited for intraoperative assessment of cardiovascular risk factors based on adipocyte size and distribution</td>
<td>Tsai et al., 2013; Weigelin et al., 2012</td>
</tr>
<tr>
<td>Erythrocytes</td>
<td>Arterial and venous vessels, interstitial tissue after hemorrhage, whole-blood ex vivo</td>
<td>Hematological analysis of blood flow dynamics, erythrocyte:leukocyte ratios, stasis or turbulence; flow-based assessment of heart rate; detection of malaria infection</td>
<td>Bélisle et al., 2008; Chen and Liu, 2012; Dietzel et al., 2014</td>
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<td>Leukocytes</td>
<td>Mouse ear, cremaster muscle</td>
<td>Detection of leukocyte infiltration into tissues during disease; identification of leukocyte subsets; leukocyte polarity</td>
<td>Rehberg et al., 2011; Tsai et al., 2012; Fig. 2C</td>
</tr>
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<td>Microvesicles</td>
<td>Mouse melanoma</td>
<td>Visualization of oncogenic conditioning of the tumor microenvironment; kinetics of microvesicle distribution in tissue</td>
<td>Weigelin et al., 2012; Fig. 2G and 3F,I</td>
</tr>
<tr>
<td>Intracellular vesicles and lipid bodies</td>
<td>Mouse hepatocytes, C. elegans and mouse embryo</td>
<td>Monitoring of lipid metabolism and lipid body dynamics</td>
<td>Débarre et al., 2008; Tseresvelakias et al., 2014; Watanabe et al., 2010</td>
</tr>
<tr>
<td>Striated muscle</td>
<td>Mouse dermis, cremaster muscle</td>
<td>Detection of I-bands complementary to the SHG signature; identifies muscle as a 3D guiding structure for tumor cell migration</td>
<td>Rehberg et al., 2011; Weigelin et al., 2012</td>
</tr>
<tr>
<td>Axons and neurons</td>
<td>Mouse brain, spinal cord</td>
<td>Dynamic monitoring of myelin organization and degeneration; orientation during image-guided brain surgery</td>
<td>Farrar et al., 2011; Lim et al., 2014; Witte et al., 2011</td>
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**Higher harmonic generation in life sciences – from humble beginnings to state-of-the-art clinical applications**

Shortly after the first laser was developed, the principles and first applications of higher harmonic generation were established, and SHG was demonstrated by Peter A. Franken in 1961 by focusing the laser through crystalline quartz (Franken et al., 1961). In 1997, the first THG microscope was developed that demonstrated the applicability of THG generation by interfaces and inhomogeneities in three dimensions (Barad et al., 1997). THG was first used in applied physics to detect laser-induced breakdown in glass (Squier et al., 1998) and liquid crystal structures (Yelin et al., 1999). The first biological applications included 3D reconstructions of plant leaves, live neurons, organelles in yeast and statolith movement in rhizoid cells, which demonstrated that THG can be used to visualize structures and dynamics in live-cell culture and tissues without the need for any exogenous labeling (Müller et al., 1998; Squier et al., 1998; Yelin and Silberberg, 1999). Initial biological applications of THG prioritized the imaging of transparent objects and the detection of THG signals in the forward direction – e.g. to track single-cell and tissue movements in early-stage embryos (Débarre et al., 2004; Sun...
et al., 2004). Over the past decade, THG microscopy has evolved from proof-of-principle studies towards a valuable tool in life science research, tissue morphogenesis and disease progression, with the potential for clinical application in optical biopsies (Table 1).

**THG imaging of cells, cellular membranes and cell organelles**

**Cellular tissue architecture and composition**

Cellular membranes, including the cell membrane, nuclear membrane and intra- and extracellular vesicles, are an abundant source of THG signals (Zimmerley et al., 2013) (Fig. 2C–G; Movies 1–4). In intact epithelia, such as mouse and human skin, THG has been used to reveal the morphology, size and distribution of epidermal and dermal cells, such as keratinocytes, adipocytes (Fig. 3E) and infiltrating leukocytes with sub-micron spatial resolution (Tai et al., 2005; Weigelin et al., 2012). Likewise, in human cornea, THG has been used to visualize epithelial cell boundaries and corneal keratocyte networks, which has helped to monitor keratocyte migration and activation during haze formation and corneal allograft wound healing (Aptel et al., 2010; Olivier et al., 2010a). In tissue-engineered corneas, combined SHG and THG imaging has recently been used to provide noninvasive quality control measurements of cornea thickness, of collagen structure, and of the distribution and density of stromal fibroblasts and endothelial cells (Jay et al., 2015).

Similarly, the position and migration of leukocytes infiltrating the mouse cremaster muscle and the mouse ear have been readily detected by using THG microscopy (Rehberg et al., 2011; Tsai et al.,

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**Fig. 3. Examples of tissue structures visualized in vivo by using THG.** The diversity of the microanatomy of connective tissue (shown here, the mouse dermis) is visualized by combined THG, SHG and fluorescence imaging. Complementary to SHG and fluorescence, THG reveals collagen fibers and cell bodies in collagen-rich loose (A) and dense (B) interstitial tissue, basement membranes along myofibers and blood vessels (C), the multilayered myelination of peripheral nerves (D) and the surface of adipocytes (E). (F) In addition, THG-positive microparticles are detected, which drift through spaces between collagen fibers. Scale bars: 50 µm. Panels A–F are adapted from Alexander et al. (2013) with permission. (G) Erythrocyte flow in capillaries visualized by using THG. Shown here is a color-coded overlay of sequential images from a 1-min time-lapse sequence. Adapted from Weigelin et al. (2012). (H) Blood flow kinetics based on erythrocyte shape. The roundish shapes of immobile (lower vessel) or slowly moving erythrocytes (middle vessel) compared with a stripe-like THG pattern are the result of fast erythrocyte movement during image acquisition (upper vessel). The scanning direction was horizontal from left to right; this causes distortion of objects as they are moving during the scan. (I) B16F10 mouse melanoma cells invading into the dermis along muscle fibers (**m**). THG allows the visualization of the outer membrane of muscle fibers (asterisks, zoom 1) and perfused blood vessels, which serve as guidance structures for melanoma invasion (asterisks, zoom 2). In addition, THG-positive microvesicles are located in the interstitial space (arrowheads, zoom 1 and 2). Excitation conditions (all panels) – THG, SHG, dsRed, 1180 nm, 100 mW; GFP, Al750, 910 nm, 20 mW; backward detection.
functions and disorders of lipid biogenesis, and are associated with lipid bodies, which are ubiquitous structures involved in metabolic cellular processes and are present in a variety of healthy cells, including tumor cells, and have been found in tissues locally and systemically (Weigelin et al., 2012). Lipid bodies can also be shed by cells during apoptosis and fragmentation, generating vesicles of variable sizes (Fig. 2E,F), but microvesicles are also shed by cells in response to shear stress between erythrocytes and the vessel wall, and flow-dependent rates and rhythmicity of the heart beat (Dietzel et al., 2014). THG contrast in leukocytes primarily results from the plasma membrane and intracellular granules, which allows the identification of leukocyte subsets based on their differential granularity. Using this approach, it has been shown that neutrophils elicit a strong THG signal from intracellular granules, whereas lymphoid cells give rise to a preferential THG signal at cellular boundaries of hollow cores that resemble their large nuclei (Tsai et al., 2012). In addition, the polarity of moving leukocytes can be resolved easily owing to the preferential redistribution of granules to the rear uropod (Fig. 2C; Movie 4). Thus, known cell subsets can be discriminated by using THG based on morphology and kinetic criteria, without the need for a fluorescent transgenic reporter strain.

Complementary to the visualization of cells, the THG signal originating from nuclear membranes (Fig. 2G) allows the characterization of shape, size and distribution of nuclei in tissues (Chu et al., 2005; Rehberg et al., 2011). Furthermore, chromocenters and heterochromatin structures are revealed in native tissue because of their different refractive indexes, without the need for DNA-intercalating stains (Rehberg et al., 2011). Exogenous application of acetic acid, which reversibly crosslinks nucleoproteins, can further enhance THG contrast in intact nuclei by changing their refractive index (Lee et al., 2007). This can be exploited to improve the detection of (pre-)cancerous lesions in mucosal tissues with optical biopsies, based on the presence of abnormalities in nuclear morphology and nucleocytoplasmic ratios that are typical for malignant cells (Lee et al., 2007; Tai et al., 2006; Tsai et al., 2011).

Blood vessel structure and function
Vascular biology is an emerging field of application for THG. Erythrocytes elicit a particularly strong THG signal that originates from their cellular membrane (Millard et al., 1999). An additional signal, with variable efficiency, originates from basement membranes lining the vessels (Rehberg et al., 2011; Weigelin et al., 2012). The resulting time-lapse THG recordings of erythrocyte flow kinetics reflect hemodynamic parameters, including blood flow velocity, the shear stress between erythrocytes and the vessel wall, and flow-derived rates and rhythmicity of the heart beat (Dietzel et al., 2014) (Fig. 3G,H). In addition, resonance enhancement of the THG signal through distinct absorption maxima of oxy- and deoxyhemoglobin allows the simultaneous assessment of the level and pattern of blood oxygenation at cellular resolution (Chang et al., 2010b).

Cell-derived microvesicles
In addition to arising from intact cells, strong THG signals also originate from small cell-derived vesicles. In particular, cellular fragmentation during apoptosis generates cellular vesicles of variable sizes (Fig. 2E,F), but microvesicles are also shed by a variety of healthy cells, including tumor cells, and have been implicated in intercellular communication and oncogenic reprogramming of the local tumor microenvironment (Figs 2G and 3F,I) (Muralidharan-Chari et al., 2010; Zomer et al., 2015). In mouse melanoma lesions in vivo, extracellular tumor-associated microvesicles of variable size can be directly detected by using THG and followed in time (Weigelin et al., 2012). This has revealed at least three kinetic subsets of interstitial microvesicles, including stationary, slow diffusive and fast-moving vesicle subsets, and these properties might correlate with their function in conditioning cells and tissues locally and systemically (Weigelin et al., 2012).

Lipid bodies
Lipid bodies are ubiquitous structures involved in metabolic cellular functions and disorders of lipid biogenesis, and are associated with many human diseases, such as diabetes, cardiovascular diseases, obesity and cancer. Using THG, intracellular lipid body dynamics have revealed, for example, an altered metabolism that is defined by an increase in lipid body size and number in hepatocytes after partial hepatectomy (Débarre et al., 2006). Similarly, stage-dependent alterations of lipid deposits have been detected in Caenorhabditis elegans embryos and adult worms, including early-onset fat deposition during development followed by gradual reduction during aging, by using noninvasive long-term THG imaging (Tserêvelakis et al., 2014). Similarly, THG time-lapse recordings in early mouse embryos have made it possible to observe the dynamics and aggregation kinetics of lipid droplets, and their movement, which depends on microtubules and microfilaments (Watanabe et al., 2010). Thus, alongside CARS, spontaneous Raman spectroscopy and lipophilic fluorescent dyes (Yen et al., 2010), THG also allows us to monitor the kinetics of fat metabolism and lipid storage.

Myofibers
Although SHG signals are elicited from ordered myosin in the A-bands of the sarcomere in striated myofibers, THG provides an alternating and complementary signal originating from the actin-rich I-bands between sarcomeres (Rehberg et al., 2010, 2011; Weigelin et al., 2012). THG provides label-free confocal 3D analyses of contracting striated myofibers and cardiomyocytes. By using cross-correlation with SHG and fluorescence, the kinetics of macro- and nanocontractions, related NADH levels, mitochondrial positioning and energy state can be determined (Barzda et al., 2005).

Brain and nervous system
To facilitate fast conduction of action potentials, nerves and axons in the central and peripheral nervous systems are surrounded by fatty myelin sheets. The multi-layered, lipid-rich membranes of the myelin sheaths that surround peripheral axons are a particularly strong source for THG (Fig. 3D,I) (Rehberg et al., 2011; Weigelin et al., 2012). In ex vivo and in vivo preparations of sciatic nerves of mice and rats, THG can be used to further resolve subdomains of myelinated peripheral nerves, including nodes of Ranvier, Schmidt–Lantermann incisures and Cajal bands. By using THG, myelination characteristics are amenable to rapid acquisition and bias-free automated quantification over large imaging fields (Lim et al., 2014). Therefore, THG might be useful to trace myelin loss and recovery in live tissue; for example, the degeneration of myelin sheets in multiple sclerosis, which causes deficits in sensory and motor functions of the demyelinated nerves. In the brain, THG signals correspond to regions of white matter, such as the mossy fibers in the cerebellar lobes, the pons and the hindbrain, and axon bundles in the corpus callosum (Farrar et al., 2011). Dense gray matter also produces THG, with a less intense and more homogeneous signal compared to that produced by white matter (Witte et al., 2011). The structural information provided by THG signals in neuronal tissue, and the possibility of being able to simultaneously visualize glass pipettes or small electrodes based on their strong refractive index, can guide the targeting of micropipettes to specific neurons for patch-clamp experiments or image-guided microsurgery applications (Witte et al., 2011). Taken together, these examples indicate that THG is a particularly promising approach to visualize changes in myelin organization in vivo during development and neurodegenerative diseases.

Virtual optical biopsy
Live-tissue analysis is an emerging field in clinical diagnostics, whereby image contrast is based on the application of exogenous lipophilic fluorescent dyes (Yen et al., 2010), THG also allows us to visualize glass pipettes or small electrodes based on their strong refractive index, can guide the targeting of micropipettes to specific neurons for patch-clamp experiments or image-guided microsurgery applications (Witte et al., 2011). Taken together, these examples indicate that THG is a particularly promising approach to visualize changes in myelin organization in vivo during development and neurodegenerative diseases.
dyes, or absorption and endogenous fluorescence of human tissues. THG has the potential to complement these existing imaging approaches. The applicability of THG in optical biopsies has been demonstrated in hamster and human oral mucosa, as well as human skin and adipose tissue (Chen et al., 2009; Tai et al., 2005, 2006; Tsai et al., 2011, 2013). Furthermore, the image contrast generated using THG is suitable for discriminating melanoma and basal cell carcinoma lesions from normal skin and benign nevi (Chen et al., 2010). It has been shown that an abundant distribution of single cells or cell clusters with strongly enhanced THG intensity correlates with the morphology and melanin content of melanoma cells that have been identified in histological slices (Chen et al., 2010). THG could further be exploited to define the tumor margin during surgery or to verify the complete excision of the lesion in the removed tissue. Moreover, when THG has been used during heart surgery to assess the morphology, size and distribution of adipocytes, it has revealed a correlation between increased numbers of small-sized epicardial adipocytes and coronary artery disease (Tsai et al., 2013).

In recent clinical trials, the safety of THG for in vivo applications has been demonstrated. THG-based diagnosis of basal cell carcinoma in vivo depends upon the polymorphous variations in the size and shape of transformed cells, compared to normal tissue and the presence of peripheral palisading cells (Tsai et al., 2014). An increase in the size of cells and nuclei in intact skin are characteristics of aging tissue, which can be quantified in vivo based on THG analyses (Liao et al., 2013). By providing information on cell and tissue morphology in a label-free manner, THG thus generates diagnostic 3D information with a resolution comparable to that of traditional histopathology.

Abnormal vessel structures and altered hemodynamic factors are important parameters for diagnosing malignancies, and this has motivated the development of a miniaturized video-rate THG microscope for use in optical biopsies that has a sufficiently high spatial and temporal resolution to record blood flow in skin capillaries (Chia et al., 2010). Transcutaneous THG detection of blood cell flow in dermal capillaries of human volunteers is sensitive enough to capture blood flow dynamics, and the ratios of erythrocytes and white blood cells (Chen and Liu, 2012); this can be exploited for in vivo cytometry and hematological analysis without the need to draw blood. In malaria-infected blood cells, THG is particularly enhanced owing to the altered scatter properties that are caused by hemozoin, a parasite-derived pigment (Tripathy et al., 2010). The above studies clearly demonstrate the promise of THG to be used to detect interface changes at the microscale in the clinical diagnosis of various conditions.

**Extracellular matrix and inorganic structures**

**Collagen and basement membranes**

In addition to SHG, which originates from the core of fibrillar collagen, THG is excited from the surface of protein complexes, and delivers additional complementary detail and information regarding substructures. In thick collagen bundles and elastic fibers, THG is generated at the fiber surface and gives rise to a relief-like signal (Rehberg et al., 2011; Sun et al., 2007; Weigel et al., 2012) (Figs 2G and 3A,B). Similarly, the outer membranes of myofibers and vessels induce THG, which is likely to originate from basement membrane structures (Rehberg et al., 2011; Weigel et al., 2012) (Fig. 3C,H). Using combined SHG–THG imaging in late-stage melanoma that had been implanted in the mouse ear, it has been shown that significant collagen deposition and rearrangement of collagen structures occurs in parallel to expansive tumor growth (Wu et al., 2015). By generating a complementary signal to SHG (Fig. 2A,B), THG thus delivers consistent 3D information about healthy and diseased tissues at the subcellular level.

**Bone and teeth**

Inorganic crystals and mineralized complexes elicit strong THG signals owing to their high difference in refractive index compared to that of the surrounding tissue fluids. In mineralized bone, THG imaging allows the reconstruction of the outer and inner surfaces, which can be used to follow bone development in mouse embryos (Oron et al., 2004). In teeth, strong THG signals are produced by the calcified matrices of dentin and enamel, and have made it possible to highlight the dentinotubule structures and, to a lesser extent, the prismatic structures of enamel (Kao, 2004). By being able to detect dental discontinuities, such as cracks caused by thermal or mechanical stress, or white spot lesions as signs of mineral loss, THG is well suited for early diagnosis of enamel abnormalities (Chen et al., 2008). In paleontology research, teeth are the most abundant fossil material and contain key information about crucial biological functions, such as feeding and defense. Owing to its noninvasive nature, THG allows 3D reconstructions and morphometric analyses of dentinal structures without the need for histological preparation. These proof-of-concept results suggest that THG could contribute to classify the evolution of ancient vertebrate teeth without damaging rare fossils (Chen et al., 2015).

**Combining THG and multimodal microscopy**

To enhance current microscopy practice, THG can be combined with other multiphoton-excited fluorescence intensity signals or with fluorescence lifetime imaging (FLIM) within the same excitation and detection setup (Pelegati et al., 2012) (Box 1). For instance, multi-parameter approaches have combined label-free SHG and THG with fluorescence imaging (Fig. 3I). Furthermore, THG can also be used to correlate high-resolution electron microscopy with intravital imaging of molecules by detection with gold-labeled antibodies (Lippitz et al., 2005). Emerging key applications include the analysis of live cell–tissue interactions, cell dynamics in morphogenesis and cancer, vascular biology and organization of neuronal tissue.

**Tissue dynamics in morphogenesis**

The use of time-resolved 3D THG microscopy of transparent unstained zebrafish embryos until the 1000-cell stage has allowed reconstruction of a number of aspects of tissue kinetics with high detail, including imaging of the contour of the embryo, individual cell morphologies, yolk platelets and intracellular lipid vesicles (Olivier et al., 2010b). Combined with SHG – originating from mitotic spindles – and imaging of fluorescently labeled histone-2B of nuclear chromatin, the THG signal generated at cellular membranes allows measurement of the length of the cell cycle and of nucleocytoplasmic ratios, and enables lineage tracing over multiple cell divisions. Such tracing of cell cycle dynamics in time and space has shown that the cell cycle lengthens continuously, rather than abruptly, and that it is dependent on cell position and the embryo maturation stage (Olivier et al., 2010b). The THG signal originating from intracellular lipid bodies across the embryo can further be used for velocimetric analysis of morphogenetic movements, whereby the motion pattern of lipid bodies that are dispersed in the tissue are recorded to obtain a large-field map of...
motion hotspots over time. Combined with fluorescence detection and local laser ablation of embryonic tissue structures, THG-based tracking of tissue structures (velocimetry) in Drosophila has shown that there is a long-ranging mechanical coupling of distant tissues, and that tissue and cell deformation correlates with the expression of the twist gene, a central regulator of Drosophila morphogenesis (Débarre et al., 2004; Supatto et al., 2005). Thus, the use of multimodal microscopy combined with THG can provide insights into tissue mechanics in a spatio-temporal manner.

**Tumor–stroma interface**

With respect to the tumor microenvironment, combination of THG with SHG and fluorescence detection has provided unprecedented details regarding tissue organization, cell–tissue interactions, blood flow dynamics and the dissemination of microvesicles (Weigelin et al., 2012). For instance, multispectral 3D reconstructions of early-stage melanoma lesions in the mouse dermis have identified the invasion strategies of melanoma cells, which depend on the type of tissue being invaded, as well as on tissue density and geometry. Collectively invading cells preferentially infiltrate tissue using a mechanism that is guided by confinement – i.e. the movement through pre-existing spaces along the interfaces of collagen bundles, myofibers, nerves and vessels (Weigelin et al., 2012) (Fig. 31). In contrast, regions of tissue that comprise discontinuous fibrillar collagen-rich stroma support the dissemination of single cells or small clusters, suggesting that melanoma cells disperse in response to a loose organization of tissue. Quantification of tissue spaces before and after tumor invasion shows little tissue deformation and a lack of apparent structural degradation (Weigelin et al., 2012). This suggests a predominantly tissue-guided migration that is dictated by physical principles, including haptotaxis and contact guidance. By depicting the complexity of structural microenvironments in vivo, that study also provides the basis for bio-inspired engineered microdevices to probe cell migration in defined environments that simulate space restrictions, surface topography and geometry in the physiological range encountered by cells in tissues (Stroka et al., 2014; Yevick et al., 2015). Taken together, THG combined with SHG- or fluorescence-based microscopy can provide important anatomical context in three dimensions and so yield a framework of cause–consequence relationships in tumor progression, cell migration and the associated remodeling of the microenvironment.

**Conclusions and outlook**

By detecting cell and tissue interfaces in a noninvasive, label-free manner, THG fills the gap in resolution between mesoscale imaging, using OCT or high-resolution ultrasound, and nanoscopy that uses scanning electron microscopy to map the 3D tissue organization and function. Because of its ease of technical implementation in existing multiphoton imaging platforms and versatile application together with SHG and fluorescence imaging, THG shows great promise to emerge as a standard application of infrared multiphoton microscopy. Future in vitro applications in transparent samples will particularly benefit from collecting signals from forward THG and so help to address cell–matrix interactions in label-free samples, tissue remodeling and microvesicle biology in complex cell models. For intravitral microscopy in small-animal models, THG is particularly useful when combined with chemical or genetically encoded probes. Promising fields of application include neuroscience, to address normal and perturbed microanatomy of central and peripheral neuronal tissue; material sciences, to monitor the 3D organization and tissue integration of engineered microdevices and biomaterials; vascular biology, for label-free detection of vascular anatomy and cell flow kinetics; and immunobiology and cancer research, to reveal tissue structures that guide and determine the positioning of cancer and immune cells. Furthermore, with the development of simpler, portable and more affordable devices, THG could enter clinical applications and field research. Here, areas of application include the optical biopsy of lipid organization and interfaces of body surfaces, dental disease – including the surface structure of teeth and tooth–tissue interfaces – and the analysis of diseased erythrocytes, such as after infection with malaria parasites. As an emerging field, optical THG-based tissue analysis will benefit from the development of microendoscopes, which will allow us to reach inner body surfaces and cavities. Thus, by enabling kinetic 3D analysis of tissue organization and function, THG will enhance both integrative 3D cell biology and preclinical and clinical ‘dynamic histopathology’.

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**Competing interests**

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**Supplementary information**

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