Breakthrough Technologies

In Vivo Chemical and Structural Analysis of Plant Cuticular Waxes Using Stimulated Raman Scattering Microscopy

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The cuticle is a ubiquitous, predominantly waxy layer on the aerial parts of higher plants that fulfills a number of essential physiological roles, including regulating evapotranspiration, light reflection, and heat tolerance, control of development, and providing an essential barrier between the organism and environmental agents such as chemicals or some pathogens. The structure and composition of the cuticle are closely associated but are typically investigated separately using a combination of structural imaging and biochemical analysis of extracted waxes. Recently, techniques that combine stain-free imaging and biochemical analysis, including Fourier transform infrared spectroscopy microscopy and coherent anti-Stokes Raman spectroscopy microscopy, have been used to investigate the cuticle, but the detection sensitivity is severely limited by the background signals from plant pigments. We present a novel method for label-free, in vivo structural and biochemical analysis of plant cuticles based on stimulated Raman scattering (SRS) microscopy. As a proof of principle, we used SRS microscopy to analyze the cuticles from a variety of plants at different times in development. We demonstrate that the SRS virtually eliminates the background interference compared with coherent anti-Stokes Raman spectroscopy imaging and results in label-free, chemically specific confocal images of cuticle architecture with simultaneous characterization of cuticle composition. This innovative use of the SRS spectroscopy may find applications in agrochemical research and development or in studies of wax deposition during leaf development and, as such, represents an important step in the study of higher plant cuticles.

The majority of land plants possess an extracellular, waxy cuticle that covers the surface of their aerial parts and protects them against desiccation, external physical and chemical stresses, and a variety of biological agents (Grncarevic and Radler, 1967; Barthlott and Neinhuis, 1997; Krauss et al., 1997; Ristic and Jenks, 2002; Yeats and Rose, 2013). The cuticle is a composite layer composed mainly of cutin and overlaid by cuticular waxes. Cutin is a macromolecular structure consisting primarily of hexadecanoic (palmitic) and octadecenoic (vaccenic) acids that are covalently linked by ester bonds to generate a rigid, three-dimensional network that is embedded with polysaccharides. Cuticular waxes are composed of long-chain (C20-C40) aliphatic molecules derived from fatty acids (Samuels et al., 2008), and studies over the last several decades have identified structural and regulatory constituents of the biosynthetic pathways of cuticular components (Kolattukudy, 1981; Beisson et al., 2012). In addition to the physiochemical properties conferred by its lipid components, the architecture of the cuticle plays an essential role in physiological function. For example, through understanding the properties of the cuticular structure, the extraordinary superhydrophobicity of the Lotus spp. leaf has been mimicked in micro- and nanotechnology to generate self-cleaning surfaces (Bhushan and Jung, 2006; Bhushan et al., 2009; Koch et al., 2009).

As may be expected, given the diversity of plants, the habitats they inhabit, and individual life histories, the morphology and composition of plant cuticle varies extensively between and within species and includes plate-, needle-, and pillar-shaped wax crystals (Barthlott et al., 2008). In some species, cuticular wax composition is known to vary with depth, giving rise to chemically distinguishable layers (Yeats and Rose, 2013). Finally, the cuticle is increasingly shown to be important in development (Koornneef et al., 1989; Yeats and Rose, 2013).
and pathogenesis (Lee and Dean, 1994; Gilbert et al., 1996; Bessire et al., 2007; Delventhal et al., 2014). It is therefore unsurprising that interest in cuticle composition, structure, and physiology is increasing (Buschhaus et al., 2014; Hen-Avivi et al., 2014; Heredia-Guerrero et al., 2014; Xu et al., 2014). Moreover, a greater understanding of the relationship between structure and chemical composition of cuticle waxes is vital for enhancing agriculture yields, as it will further our knowledge of how plants regulate water balance and inform the application of nutrition (foliar feeds) and pesticides, leading to improved formulation strategies for agrochemicals.

The chemical composition and topological architecture of cuticular waxes are both critical for optimal physiological function. Analyses of these essential properties have typically been performed separately. Cuticle wax composition is normally determined using gas chromatography; cuticle ultrastructure may be analyzed using destructive imaging techniques such as scanning electron microscopy (SEM; Baker and Holloway, 1971; Jetter et al., 2000; Barthlott et al., 2008) and laser desorption/ionizing mass spectroscopy (Jun et al., 2010) or, in vivo, using nondestructive real-time techniques, including white-light scanning interferometry (Kim et al., 2011), atomic force microscopy (Koch et al., 2004), confocal microscopy in reflection mode (Veraverbeke et al., 2001), fluorescence microscopy of chemical stains (Pighin et al., 2004), coherent anti-Stokes Raman scattering (CARS) microscopy (Yu et al., 2008; Weissflog et al., 2010), and total internal reflection Raman spectroscopy (Greene and Bain, 2005). Despite the advances in our understanding of the cuticle that have been made with these techniques, there is a great need for techniques that combine chemical and structural information to provide in situ high-resolution chemical analysis of epicuticle waxes.

Techniques based on vibrational spectroscopy offer in situ chemical analysis derived from the vibrational frequencies of molecular bonds within a sample. However, due to water absorption and the intrinsically low spatial resolution associated with the long infrared (IR) wavelengths required to directly excite molecular vibrations, IR absorption techniques have limited value for bioimaging. Raman scattering, however, provides analysis of vibrational frequencies by examining the inelastic scattering of visible light. Raman scattered light is frequency shifted with respect to the incident light by discrete amounts that correspond to the vibrational frequencies of molecular bonds within the sample. The spectrum of Raman scattered light consists of a series of discrete peaks that each correspond to a molecular bond and can be regarded as a chemical fingerprint holding a wealth of information regarding chemical composition. Unfortunately, Raman scattering is an extremely weak effect, and typical signals from biological samples are at least six orders of magnitude weaker than those from fluorescent labels. This severely limits the application of Raman for studying living systems because long acquisition times (100 ms–1 s per pixel) and relatively high excitation powers (several hundred milliwatts) are required to image most biomolecules with sufficient sensitivity. Furthermore, the lack of sensitivity is compounded by autofluorescence, which in plant tissues completely overwhelms the Raman signal, prohibiting its application in planta.

Far stronger Raman signals can be obtained using coherent Raman scattering (CRS; Min et al., 2011). CRS achieves a Raman signal enhancement by focusing the excitation energy onto a specific molecular vibrational frequency (Fig. 1A). A pump and Stokes beam, with frequencies $\omega_p$ and $\omega_s$, respectively, are incident upon the sample, with their frequency difference $(\omega_p - \omega_s)$ tuned to match the molecular vibrational frequency of interest. Under this resonant condition, the excitation fields efficiently drive bonds to produce a strong nonlinear coherent Raman signal. When applied in microscopy format, the nonlinear nature of the CRS process confines the signal to a submicron focus that can be scanned in space, allowing three-dimensional confocal-like mapping.

![Figure 1](https://example.com/figure1.png)
of biomolecules. CRS microscopy has particular advantages for bioimaging: (1) Chemically specific contrast is derived from the vibrational signature of endogenous biomolecules within the sample, negating the need for extraneous labels/stains; (2) Low-energy, near-IR excitation wavelengths can be employed, which reduces photodamage and increases depth penetration into scattering tissues; and (3) The CRS process does not leave sample molecules in an excited state, does not suffer from photobleaching, and can be used for time course studies.

CRS microscopy may be achieved by detecting either CARS or stimulated Raman scattering (SRS).

**CARS MICROSCOPY**

CARS microscopy relies on detection of the anti-Stokes signal generated at frequency $\omega_\text{as} = 2\omega_p - \omega_S$, which, by using filters, is spectrally isolated from the pump and Stokes beams, and its intensity is used to map the location of biomolecules of interest (Zumbusch et al., 1999). The CARS signal is blue shifted with respect to the pump and Stokes wavelengths (Fig. 1B), making CARS more resilient to sample autofluorescence than spontaneous Raman. However, in highly autofluorescent samples such as plant tissues, the usually weak two-photon fluorescence (also blue shifted with respect to the excitation wavelengths) overwhelms the CARS signal. Consequently, CARS imaging in planta has only been applied to samples with reduced chlorophyll autofluorescence including dried tissues (Zeng et al., 2010; Ding et al., 2012), roots (Ly et al., 2007), and cuticle waxes after they have been stripped away from the leaf (Weissflog et al., 2010).

**SRS MICROSCOPY**

SRS relies on detecting subtle changes in the intensities of the excitation fields that occur by virtue of stimulated excitation (Freudiger et al., 2008). When the difference frequency, $\omega_p - \omega_S$, matches the frequency of a molecular vibration, the Stokes beam intensity experiences a gain, while the intensity of the pump beam experiences a loss, shown in Figure 1C. This transfer of intensity between the excitation beams only occurs when both beams are incident simultaneously on the sample and is measured by detecting the modulation that is transferred to the pump beam after it has passed through the sample when the intensity of the Stokes beam is modulated. The amplitude of the transferred intensity modulation is directly proportional to the concentration of target molecules and, by modulating at frequencies above laser noise (>1 MHz), can be detected with a lock-in amplifier with great sensitivity (Ye et al., 2009).

Because SRS is detected at the same wavelength as the excitation fields, it is not affected by fluorescent emission and, as recently demonstrated by Mansfield et al. (2013), may also be used in the presence of highly pigmented samples such as plant tissues.

We have recently shown that it is possible use phase-sensitive detection to separate the vibrational SRS signal from the electronic absorption processes (Mansfield et al., 2013). In this investigation, we used SRS imaging to investigate both the structure and chemical composition of plant cuticular waxes, simultaneously and in vivo. We compared SRS images of cuticle wax structures from a variety of plant species that display characteristic wax morphologies with SEM and CARS images of the leaf surface. We demonstrated that spectroscopic information can be obtained to provide chemically specific, in situ analysis of waxes on living leaves with submicron spatial resolution. Using SRS microscopy, we compared the cuticle structure and composition between wild-type Arabidopsis (Arabidopsis thaliana) and an nctf1 mutant (cer1) mutant with impaired cuticle deposition and have also shown that SRS imaging is sufficiently sensitive to track temperature-induced changes in cuticle formation in Thellungiella salsuginea. We conclude that, with the increasing availability of commercial Raman-based microscopes, SRS imaging has the potential to radically advance our understanding of plant cuticular structure and composition and their effects on physiology and development.

**RESULTS**

**Characterization of the Raman Spectra of Dissolved Cuticle Waxes**

Spontaneous Raman spectra of hexane-extracted wax samples over the CH vibrational region from 2,700 to 3,200 cm$^{-1}$ were acquired as a baseline reference for each plant species (Fig. 2). Gaussian peaks were fitted underneath each of the spectra to visualize the relative contribution of different molecular vibrations, based on peaks previously identified in the literature (Snyder et al., 1978; Ho and Pemberton, 1998; Greene and Bain, 2005). The wax extracts from Arabidopsis, Thellungiella parula, banana (Musa acuminate), and cheese plant (Monstera delicosa) had similar spectral profiles that matched those reported for other waxes and alkanes reported in the literature (Snyder et al., 1978; Greene and Bain, 2005) and deconvolved into seven identifiable peaks (Table I). The peaks, corresponding to the symmetrical and antisymmetrical CH$_2$ stretch, are prominent on all spectra. The ratio antisymmetric-to-symmetric peak height is known to give a strong indication of the alkyl chain conformational order with a ratio of 1.6 to 2 indicating a highly crystalline structure, and a ratio of 0.6 to 0.9 indicating a liquid structure (Snyder et al., 1978; Ho and Pemberton, 1998, Greene and Bain, 2005). For the wax samples analyzed here, the ratios were as follows: Arabidopsis, 1.13; T. parula, 1.58; banana, 1.34; and cheese plant, 1.54; indicating structures intermediate between a highly crystalline structure and a liquid. The curve fits to the spectra also included two Fermi resonances of the CH$_2$ bond at 2,930 cm$^{-1}$ and another very broad Fermi resonance at approximately 2,870 cm$^{-1}$ (Snyder et al., 1978). The contributions due to
the CH$_3$ symmetric and antisymmetric stretches were small, as proportionally there are much less of these groups due to the long chain lengths. The fitting parameters for each wax along with the goodness of fit are included in Supplemental Text S1.

Two of the species investigated, Dudleya anthonyi and silver dollar plant (Xerosicyos danguyi), showed dramatically different Raman spectra from Arabidopsis, T. parvula, banana, and cheese plant, characterized by additional peaks, which are indicative of a more complex chemical structure, with additional chemical bonds to those described in Table I. In D. anthonyi, the Raman spectrum changed depending on the number of the hexane wash and hence, we surmise, on the depth of penetration into the cuticle; the initial, most superficial hexane wash showed an unusual Raman spectrum, but subsequent washes that solubilize waxes deeper in the cuticle showed spectra that matched more closely with those of a typical alkane. These differences are most readily explained by both these species possessing a thick, glaucous cuticle compared with the other plant species studied, an adaptation enabling survival in xeric environments.

In Vivo Comparison of CARS and SRS Images of Plant Cuticle

Having characterized the Raman spectra of waxes extracted from the cuticle of a variety of plants, we imaged the cuticles of the same plants in vivo, using both CARS microscopy (Weissflog et al., 2010) and SRS microscopy (Fig. 3). SRS and CARS images of the epicuticular wax layer of T. parvula leaves were acquired simultaneously (Fig. 3) and are presented as the exemplar to compare the two techniques.

The images acquired using CARS are dominated by high levels of background autofluorescence from cell walls and chloroplasts (Fig. 3A). This autofluorescence is due to inevitable two-photon absorption by the compounds in these organelles and the resulting fluorescence emission at similar wavelengths to the CARS signal. Consequently, in the CARS images, only the largest wax crystals can be visualized.

Conversely, in the SRS image (Fig. 3B), there is no fluorescent background because the signal measured is

| Table I. Raman peaks used in curve fitting for cuticular waxes |
|-----------------|-----------------|
| Peak Wavelength | Assignment       |
| (Approximate)   |                 |
| 2,840 (crystalline) | CH$_3$ symmetric stretch |
| 2,850 (liquid)  | CH$_3$ antisymmetric stretch |
| 2,870 (crystalline) | CH$_3$ antisymmetric (out of plane) |
| 2,880 (liquid)  | CH$_3$ antisymmetric (in plane) |
| 2,930           | CH$_3$ Fermi resonance |
| 2,952           | CH$_3$ antisymmetric (out of plane) |
| 2,964           | CH$_3$ antisymmetric (in plane) |
| 2,880 to 2,901  | CH$_3$ symmetric |
| 2,870 (very broad) | CH$_3$ Fermi resonance |
not due to a change in wavelength (i.e. fluorescence relative to excitation) but a change in intensity between the excitation lasers (the pump and Stokes beams) following stimulated Raman excitation of the samples. The cuticular wax crystals are clearly visible, and the SRS signal spectrally matches that of the spontaneous Raman spectrum of purified *T. parvula* wax (Fig. 3C).

**Figure 3.** CARS and SRS images of the *T. parvula* leaf surface. The surface of a leaf from *T. parvula* was simultaneously imaged using CARS microscopy (A) and SRS microscopy (B). Both images were acquired at a 2,845 cm\(^{-1}\) CH\(_2\) symmetric stretch. The CARS image (A) is dominated by autofluorescence, and only the largest wax crystals are visible against the background. Conversely, the SRS image (B) is almost background free, enabling the cell walls and wax crystals to be clearly visualized. C shows the spectral scan of the SRS (red crosses) from the wax crystals in situ overlaid onto the spontaneous Raman spectra of purified *T. parvula* wax (blue line), purified cellulose (green line), and purified pectin (purple line).

**Figure 4.** SRS images and scanning electron micrographs of *Arabidopsis* cuticle. The surface of *Arabidopsis* stems were imaged using SRS microscopy (A, C, and D) and SEM (B, D, and E). A and B, The surface of the untreated stem of wild-type *Arabidopsis*, ecotype Landsberg *erecta*, with the structure of cuticle wax crystals clearly visible. C and D, The surface of the wild-type *Arabidopsis* stem following a wash in hexane that dissolved the cuticle waxes. E and F, The surface of the stem from a *cer1* *Arabidopsis* mutant. A, C, and E were generated from three-dimensional stacks taken of a 64\(\times\)64-\(\mu\)m field of view and are displayed in false color.
of the cuticular waxes and cellulose. SRS microscopy therefore, not only provides clearer images of plant cuticle structure than CARS microscopy, but, because of the low levels of background noise in the images, also has the capacity to yield information on specific compounds that compose the imaged structures.

Comparison of SRS and Scanning Electron Microscope Images of Plant Cuticle

To confirm the capacity of SRS microscopy for accurately resolving cuticular structures, the surface of Arabidopsis stems were imaged using SRS microscopy and the more conventional SEM (Fig. 4). SRS and SEM images of untreated, wild-type stems (Fig. 4, A and B) show an uneven surface composed of similarly proportioned globules, although the level of detail is higher in the SEM image. We ascertained that these globular structures were waxes by washing the stems in hexane prior to imaging (Fig. 4, B and C). In this case, both images show that the surface of the stem is smooth, with clearly visible cells and stomata. Finally, stems from Arabidopsis defective for CER1 gene expression were imaged using either technique (Fig. 4, E and F). cer1 mutants are impaired in cuticle wax biosynthesis (Aarts et al., 1995; Bernard et al., 2012), and the images show stems with markedly reduced surface structures compared with the wild type (Fig. 4, A and B).

Unlike SEM, which provides information only on the surface topology of imaged samples, SRS microscopy is a technique of multiphoton confocal microscopy that, like conventional confocal microscopy, can penetrate materials and provide information on their subsurface structure. To illustrate this technical capability, we imaged the cuticles of banana and silver dollar plant (Fig. 5). Banana was chosen, as it is a monocot, with a visibly waxy cuticle and regular files of elongated, epidermal cells. Silver dollar plant was chosen because of its thick, glaucous cuticle composed of different waxes with different Raman spectra.

Three-dimensional reconstructions of confocal SRS images of banana (Fig. 5, A and C) and silver dollar plant (Fig. 5, B and C) cuticles show very different structures. The banana cuticle is formed of a number of hair-like projections from the leaf surface. In the comparator SEM image (Fig. 5E), these projections clump together, most likely due to the process of fixation, and show a less even distribution. By contrast, the silver dollar plant cuticle has an amorphous surface, supported by a
A reticulate pattern (6–10 μm below the surface) of columnar structures emerging from the leaf epidermis. The amorphous surface of the silver dollar plant cuticle is clear on the comparator SEM image (Fig. 5F), but the intriguing cuticle substructure is not.

The comparison between SRS and SEM images therefore demonstrates that sufficient contrast can be produced using SRS microscopy to provide credible, structural images of waxes with submicron resolution that correlate well with images from SEM and can provide critical, structural information not previously available.

**Simultaneous Ultrastructural Imaging and Chemical Analysis of Cuticle by SRS Microscopy**

As previously noted, SRS imaging relies on the chemically specific Raman spectrum of the cuticle constituents. To verify the potential of the technique to provide structural and chemical information, simultaneously and in vivo, we first analyzed the cuticle of *D. anthonyi* (Fig. 6). *D. anthonyi* was selected for this aspect of the investigation because previous work had shown stratification of the composition of cuticular wax in the *D. anthonyi* cuticle (Jetter et al., 2000).

Both the SRS (Fig. 6A) and SEM (Fig. 6B) images of the *D. anthonyi* cuticle show a complex structure; the outer cuticle layer has a structure in which many small wax crystals are amassed together, with the deeper layers of the cuticle having a smoother, somewhat reticulated or wavelet-like appearance (Fig. 6D). Most importantly for this investigation, it is clear that the wax composition, as reported by the SRS spectra (Fig. 6C), was different depending on cuticle depth. The SRS spectrum acquired from deeper in the cuticle was most similar to an alkane, whereas the SRS spectrum from the surface components was markedly different, indicating a different chemical composition.

It is well documented that changes in wax composition and deposition during development and in response to abiotic stress or pathogen attack are important in the life history of many plant species (Raffaele et al., 2009; Bourdenx et al., 2011). *T. salsuginea* is particularly interesting because it is a close relative of Arabidopsis and *T. parvula*, and its cuticle composition and structure alters following exposure to prolonged cold treatment or vernalization (Teusink et al., 2002; Amtmann, 2009; Xu et al., 2014).

To determine whether SRS microscopy could, in principle, document the ultrastructural and chemical changes in cuticle that may occur during plant development, we imaged leaves from *T. salsuginea* plants that had been grown either in constant temperature or exposed to a 14-d-long cold shock using SRS microscopy and SEM (Fig. 7). Plants grown at a constant temperature of 20°C show a cuticle containing aggregates of waxes on the adaxial surface of the leaf (Fig. 7, A and B) and relatively little cuticular waxes on the abaxial surface (Fig. 7, C and D). However, following incubation at 4°C for 14 d, a marked alteration in cuticle structure can be observed on both the adaxial and abaxial surfaces (Fig. 7, C–D). Following exposure to a 14-d-long cold shock, a marked increase in waxy deposits in the cuticle was observed covering the entire leaf. Moreover, the ultrastructural aggregation of the cuticle waxes was smaller and more regular than that observed on the adaxial surface of leaves grown in a constant temperature of...
20°C (Fig. 7, A and B). SRS microscopy therefore precisely reported the anticipated changes in cuticle formation that occur following exposure to cold in *T. salsuginea* and, by extension, may prove a useful, in vivo tool for investigating other dynamic changes in cuticle architecture and composition during development and in response to environmental or biotic perturbation.

**DISCUSSION**

The cuticle is a composite, hydrophobic layer composed mainly of waxes and cutin secreted from the aerial epidermis of land plants. The cuticle performs a number of physiological roles, notably acting as a barrier to water loss, which, in higher plants, enables the exquisite control of evapotranspiration by stomatal guard cells, increasing light reflection and heat tolerance in xerophytes and providing an essential barrier to the ingress of toxins and pathogens into the plant. Cuticle topography and composition has mainly been investigated using destructive techniques that provide only a fragmented understanding of the complexity of this essential structure.

In this investigation, we have generated in vivo images of cuticle waxes at submicron resolution, using SRS microscopy. Like other forms of nonlinear Raman-based imaging techniques, SRS is label free and enables the simultaneous acquisition of chemically specific data and confocal quality images. Most importantly, SRS microscopy relies on a change in signal intensity rather than a change in wavelength.
than wavelength, so images can be acquired without the strong autofluorescent background from chloroplasts and cell walls that hampers the resolution and interpretation of CARS and spontaneous Raman images (Gierlinger et al., 2010, 2012).

The structures observed using SRS microscopy compared well to those seen under the SEM, albeit with a lesser degree of resolution due to the inherent limitations of optical microscopy relative to electron microscopy. However, unlike SEM, which generates images from the surface of fixed or cryogenically preserved samples, SRS imaging, like other forms of single- or multiphoton laser microscopy, enables information to be acquired in vivo from within the sample and three-dimensional reconstructions of internal structures to be reconstructed. This capacity is particularly advantageous for documenting dynamic changes of topography or composition that may occur and that are otherwise undetectable using fixed samples.

As demonstrated here, the chemical specificity of the SRS signal allows differentiation of different wax components from the main constituents of cell walls, cellulose, and pectins. In this study, we showed that the xerophytes *D. anthonyi* and silver dollar plant had different cuticular wax compositions from the other mesophytic plants investigated. Although this result is unsurprising, as these plants were selected for precisely their singularly thick and glutinous cuticles, it does demonstrate the potential of SRS for in vivo characterization of cuticle constituents. Moreover, although we did not attempt in this study to quantify, in situ, the different compounds for which we acquired SRS spectra, it may be possible to use this technique in a more quantifiable manner to characterize, in vivo, the relative abundance of cuticle or cell wall components and the changes that may occur in response to environmental or developmental stimuli. In this investigation, we have shown that SRS imaging is sufficiently sensitive enough to show developmental changes in wax production in *T. salsuginea*. Moreover, the technique may be useful for characterizing wax biosynthesis mutants, such as the exemplar used here, cer1, which has been shown to have a role in both water use and Arabidopsis interaction with fungal and bacterial pathogens (Raffaele et al., 2009; Bourdenx et al., 2011). SRS microscopy may therefore provide new insights into the physiological responses of plants to drought, temperature, chemicals, or pathogens and underpin the use of alternative model systems, such as *T. salsuginea* (Amtmann 2009), to investigate the control of cuticle deposition.

However promising, any new technology or application must be simplified to enable widespread use. Currently, SRS microscopy is somewhat niche and requires specialist knowledge. However, commercial, user-friendly CARS microscopes that can be converted to SRS imaging exist and, like laser confocal microscopy, will become increasingly common and add real-time, label-free Raman microscopy to the range of imaging techniques widely available to researchers in plant science.

CONCLUSION

The chemical composition and structure of the cuticle in mesophyllous plants are crucial elements for survival, physiology, and development. We have shown that SRS microscopy can be used to acquire in vivo, three-dimensional images of the cuticle in a variety of plant species, enabling simultaneous analysis of cuticle structure and chemical composition. In contrast to CARS microscopy, which is more common, SRS images of plant tissues contain very low autofluorescence from chloroplasts or cell walls and are therefore easier to interpret. Moreover, SRS images compare well with those acquired using SEM, albeit with the lower resolution that is due to the use of laser light rather than an electron beam. Moreover, we have shown that SRS can be used to track and quantify dynamic changes in cuticle structure and composition in response to environmental stimuli and can therefore increase our understanding of this essential and often-overlooked structure.

MATERIALS AND METHODS

Plants

Silver dollar plant (*Xerococcus danguyi*), Dudleya anthonyi, banana (*Musa acuminata*), and cheese plant (*Monstera deliciosa*) were grown in soil in the University of Exeter greenhouses and harvested between February and April 2012. The photoperiod was set to 16 h, from 5 AM to 9 PM. Supplemental lighting and shading were provided to ensure irradiance between 540 and 720 μmol s⁻¹ m⁻². The greenhouse temperature was set to 19.5°C with an SD of 3.7°C. Arabidopsis (*Arabidopsis thaliana*), *Thellungiella salsuginea*, and *Thellungiella parula* were grown in a 16-h-light/8-h-dark photoperiod at 20°C in growth rooms controlled for temperature and humidity. Four weeks after germination, the *Thellungiella* spp. were transferred to an ambient temperature of 4°C for 2 weeks and then returned to 20°C for an additional 2 weeks. The light regime remained at 16 h of light/8 h of dark during all experiments. Leaves were harvested for imaging and biochemical analysis.

Purification of Wax from Plant Cuticle

Following excision from plants, leaf surface areas were measured and logged. Leaf surfaces were washed for 30 s in 15 mL of high-performance liquid chromatography-grade hexane (Sigma). The hexane solvent and dissolved cuticular waxes were decanted into a glass vial that had been washed with acetone and dried. Leaves were washed a second time with 2 mL of hexane, which was added to the appropriate sample vial. Hexane was evaporated under a continuous stream of N₂ to dryness, and the cuticle wax was redissolved in 250 μL of hexane.

Spontaneous Raman Spectroscopy

Spontaneous Raman spectra of purified wax samples were acquired using a Renishaw RM100 Raman microscope equipped with a 785-nm diode laser and a 1,200 line mm⁻¹ spectral grating, which gave a spectral resolution of 1 cm⁻¹. Samples were mounted on aluminum-coated microscope slides.

SRS Microscopy

A detailed technical explanation of the materials and methods for SRS microscopy is provided to promote the implementation of the technique as widely as possible. For stimulated Raman gain microscopy, two laser systems were used: The Stokes beam was provided by the signal output from the picosecond Optical Parametric Oscillator (Levanté, Emerald APE) pumped by the frequency doubled output from a neodymium vanadate ps oscillator (picoTrain, HighQ laser). The pump beam was provided by a titanium sapphire laser (MIRA 900 D, 1,200 line mm⁻¹ spectral grating, which gave a spectral resolution of 1 cm⁻¹. Samples were mounted on aluminum-coated microscope slides.
Coherent) tuned to 770 nm and operating in ps mode, which was electronically synchronized to the neodymium vanadate laser (Coherent, Synchro-lock AP), thereby ensuring the laser pulses were temporally overlapped. The 770-nm pump beam was amplitude modulated at 1.7 MHz using an acousto-optical modulator (Crystal Technology, model 3800-122) and combined with the signal output from the optical parametric oscillator using an 850-nm short-pass dichroic mirror. Imaging was performed using a modified confocal laser scanning unit (Flouview 300, Olympus) and Olympus IX71 inverted microscope. The laser light was focused onto the sample using a 60×1.2 numerical aperture water immersion microscope objective (UP130N Apo, Olympus). The transmitted light from the sample was collected with a 60×1.0 numerical aperture water-dipping condenser (LUMFI, Olympus). The transmitted Stokes beam was detected using an silicon photodiode (Thorlabs, FDS1010) with a 70-V reverse bias. The pump beam was blocked from the photodiode using an 850-nm-long pass filter (HP850p, Chroma Technologies). The output from the photodiode was passed through a long-pass filter (minicircuits, BLF-1.9+) to remove modulations at the 76-laser repetition rate and terminated by a 50-Ω resistor. The filtered beam was then fed into a lock-in amplifier (Zurich Instruments, HF2LI Lock-In Amplifier) that separated out the modulated SRS signal at 1.7 MHz. SRS spectra were obtained by tuning wavelength of signal beam in 2-nm intervals and acquiring a series of images.

Stimulated Raman gain was measured in preference to stimulated Raman loss, as it allowed the chemically specific Raman signal to be separated from any additional signal due to two photon absorption (Ye et al., 2009) or photothermal lensing (Lu et al., 2010; Moger et al., 2012) by phase-sensitive lock-in detection (Mansfield et al., 2013).

Fresh, excised plant leaves were mounted in perfluorodecalin (Littlejohn et al., 2011) prior to imaging, which has been shown not to interfere with Raman-based imaging in plants (Mansfield et al., 2013; Littlejohn et al., 2014).

**SEM**

Leaf samples were imaged using cryogenic SEM. Samples were flash frozen in liquid N₂ slush, transferred to a vacuum, and coated in gold using the Catan Alto 2100 system. Images were acquired using a JEOL JSM-6390 LV scanning electron microscope operating at 5 kV with a working distance of 10 to 12 nm.

**Supplemental Data**

**Supplemental Text S1.** The Raman fitting parameters for each purified cuticular wax.

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