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New insights into lipid and fatty acid metabolism from Raman spectroscopy.

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Abstract.

One of the challenges facing biology is to understand metabolic events at a single cellular level. While approaches to examine dynamics of protein distribution or report on spatiotemporal location of signalling molecules are well-established, tools for the dissection of metabolism in single living cells are less common. Advances in Raman spectroscopy, such as stimulated Raman scattering (SRS), are beginning to offer new insights into metabolic events in a range of experimental systems, including model organisms and clinical samples, and across a range of disciplines. Despite the power of Raman imaging, it remains a relatively under-used technique to approach biological problems, in part because of the specialised nature of the analysis. To raise the profile of this method, here we consider some key studies which illustrate how Raman spectroscopy has revealed new insights into fatty acid and lipid metabolism across a range of cellular systems. The powerful and non-invasive nature of this approach offers a new suite of tools for biomolecular scientists to address how metabolic events within cells informs on or underpins biological function. We illustrate potential biological applications, discuss some recent advances, and offer a direction of travel for metabolic research in this area.

Abbreviations:

ALS	Alternative least squares
A-PoD	Adaptive moment estimation optimization-based pointillism deconvolution
BMAds	Bone marrow-derived adipocytes
CARS	Coherent anti-Stokes Raman scattering
cisR	Cisplatin resistant (cells)
CRS	Coherent Raman scattering
hSRS	Hyperspectral-SRS
h ² SRS	High content, hyperspectral SRS
KCMA	K-means cluster analysis
LASSO	Least absolute shrinkage and selection operator
LD	Lipid droplet
MS	Mass Spectrometry
ODYA	17-octadecynoic acid
PCA	Principal component analysis
SILAC	Stable isotope labelling by amino acids in cell culture
SRG	Stimulated Raman gain
SRL	Stimulated Raman loss
SRS	Stimulated Raman scattering

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1. Introduction.

No two cells are the same. Driven initially by the application of single cell sequencing technologies, cellular heterogeneity is now a widely accepted feature of almost all cell types [1-5]. The study of multicellular tissue dynamics has reached a new frontier as it is recognised that individual cells possess unique characteristics indicative of distinct functional roles, and that this cellular variation has profound implications for understanding cell/tissue function and for disease development [2, 5-7]. Using stains, antibodies, or specific labels for (a) protein(s) of interest, fluorescence microscopy has provided a step-change in knowledge and understanding of biological complexity even within a tissue composed of supposedly similar cells. By contrast, our understanding of the metabolic processes localised to specific cells within a tissue remains ill-defined. This is largely because of difficulties in identifying specific chemical moieties in living cells by means other than using specific engineered reporter molecules (e.g. such as those described for NAD⁺ and ATP [8, 9]), and the limitation of population averaging effects in heterogeneous cell populations. Hence, key details of metabolic profiles within cells remain obscure. The ability to use Raman imaging, with or without non-invasive probes generating signals in the Raman spectrum, offers a way forward to resolve these issues. This has been a particularly fruitful avenue of research for those interested in fatty acid and lipid metabolism. Here we will consider how Raman has advanced this area and speculate on future developments (Figure 1).

Lipid and fatty acid metabolism. The storage, release and breakdown of fatty acids play key roles across many facets of physiology. Adipocytes represent the best understood and most extensively studied cell type involved in such regulation, accumulating, esterifying, and storing excess fatty acids and lipids as triacylglycerides in lipid droplets [10, 11]. Distinctions between adipose tissue subtypes (e.g. white and brown adipocytes) are well established, but recent work has revealed considerable heterogeneity even within a single adipose depot [2, 12-14]. These differences appear to be developmentally programmed and are associated with differential susceptibility to diseases, including obesity and diabetes (for recent reviews, see [11, 15, 16]). In addition to these depot-specific variations, adipocytes can respond to signals to prioritize lipid synthesis (lipogenic) or lipid breakdown (lipolytic) depending on the prevailing hormonal milieu [10, 11].

In addition to adipose tissue, other cell types can both utilise fatty acids for energy and exhibit the accumulation of excess lipid in the form of cytosolic lipid droplets [17-21]. The liver plays a major role in lipid metabolism and is known to carefully integrate the distribution of fatty acids to other tissues and balance fatty acid utilisation. Non-alcoholic fatty liver disease arises because of lipid accumulation in hepatocytes exceeding fatty acid oxidation or export, resulting in the formation of lipid droplets within hepatocytes and impaired metabolic control [20, 21]. Like the liver, skeletal muscle uses fatty acids for energy, but in obesity increased circulating fatty acid levels result in the accumulation of toxic lipid intermediates which drive oxidative stress. This is thought to underpin insulin resistance in muscle, an

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early stage in the development of type-2 diabetes [22-25]. Such toxic accumulation of lipid droplets/lipid intermediates is also observed in the pancreatic beta cells of people with type-2 diabetes, resulting in aberrant insulin biosynthesis and release [17]. Changes in lipid metabolism are also proposed to underlie changes in cancer cell function and may in turn modulate cell biology via changes in the plasma membrane lipidome [26, 27]. Hence, there is a need to quantify lipid/fatty acid metabolism in adipocytes and other cells, to understand how their metabolism changes in disease, and to develop approaches to allow the analysis of metabolic changes, ideally in real-time. Raman-based imaging has provided insight into adipose tissue structure and function, details of lipid composition, and has been shown to offer potential for either *in vivo* or intra-operative analysis (for recent reviews see [26, 28, 29]).

Arguably, the most important aspect of Raman analysis is the ability to non-destructively image cells and study cellular events without the need for labels, genetic reporters or antibodies, hence the challenge is to integrate this with enhanced resolution and specificity of analysis. In the sections which follow, we provide a bird's-eye view of Raman imaging, then delve into specific examples where Raman-based approaches have provided new insight into fatty acid and/or lipid metabolism, illustrating these with a range of biological systems from human cells, flies, worms, fungi and bacteria chosen from the past two to three years to reflect the breadth of approaches and molecular insight which Raman can provide. We end with a speculative look to the future of the technique, highlighting some recent advances which offer great potential. We believe these approaches will be of wide general interest to the cell biology community, as the approaches discussed here are applicable across metabolism and in all kingdoms of biology.

2. What is Raman and SRS imaging?

As an aide to understanding a glossary of key terms is provided in **Table 1**. Readers familiar with Raman and SRS can skip ahead to section 3.

2.1 A brief overview of Raman. Raman scattering is an optical technique that investigates the vibrational modes of molecules through inelastic scattering of the incident light source. As such, most Raman scattering experiments make use of a confocal Raman microscope coupled to a monochromatic light source, typically a laser at 405, 532, 633, 785 or 1064 nm. An objective lens is used to focus the laser onto the sample such that both elastic (referred to as Rayleigh scattering) and inelastic scattered photons (referred to as Raman scattering) are produced following the interaction with the molecules at the sample focus. Raman systems use objective lens focuses the laser onto the sample and collect the scattered (both Rayleigh and Raman scattered) photons. By employing a spectrograph, the Stokes Raman scattered photons can be separated based on wavelength. By determining the wavelength shift in nm, which is usually expressed as a Raman shift, (ν , wavenumber cm^{-1}), the resulting Raman spectrum can be used to identify molecular vibrations



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2 for molecular characterisation. As such, a Raman spectrum plots the wavelength shift (cm^{-1} , x-axis)
3 against the relative number of photon counts (usually counts/s, or equivalent, y-axis).

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5 Raman imaging of lipids is an active area of research in part due to the numerous spectroscopic
6 features for label-free characterisation of lipid type and composition. Perhaps the most characteristic
7 Raman mode for lipid samples is the abundant CH_2 stretches including the 2851 cm^{-1} (CH_2 symmetric
8 stretch) and 2881 cm^{-1} (CH_2 asymmetric stretch), together with scissoring modes detected in the
9 fingerprint region of the Raman spectrum (1440 cm^{-1}) [30]. Fatty acid esterification can be readily
10 detected in the Raman spectrum through C=O stretching at 1740 cm^{-1} [31], whilst unsaturation of
11 the alkyl chain is observed through C=C-H stretching at 3010 cm^{-1} , C=C stretching at 1655 cm^{-1} and
12 1262 cm^{-1} (C=C-H deformation). An important area of development is the use of ratiometric Raman
13 spectroscopy for molecular characterisation. As the Raman spectrum typically contains a high
14 density of peaks for lipid species, it is possible to take the ratio of several peaks in order to
15 characterise the lipid content in cellular lipid droplets, for example [32]. Jamieson *et al.* demonstrated
16 an increasing linear relationship between the intensity ratio $1262 \text{ cm}^{-1}/1440 \text{ cm}^{-1}$ across a series of
17 C18 fatty acids [33]. Alternative lipid classes are readily detectable in the Raman spectrum including
18 cholesterols, triacylglycerols and phospholipids.

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20 **2.2 CARS and SRS. CARS and SRS.** In practice, Raman scattering is an intrinsically weak
21 process, with approximately 1 in every 10^8 incident photons undergoing inelastic scattering [34].
22 Therefore, the integration time per spectrum is usually slow (seconds) such that Raman imaging
23 over an extended sample area is experimentally slow (mins to hours). In addition, it can be
24 challenging to separate the unique molecular signature of the Raman scattering from the
25 overwhelming Rayleigh scattered light or background fluorescence. Coherent Raman scattering
26 (CRS) techniques aim to improve the scattering efficiency, and therefore detection sensitivity and/or
27 imaging speed, through coherent excitation of the sample. The two most widely used CRS processes
28 for microscopy are coherent anti-Stokes Raman scattering (CARS) [35] and stimulated Raman
29 scattering (SRS), although coherent Stokes Raman scattering (CSRS) [36] has recently been
30 reported for microscopy applications. A comparison of the three techniques discussed here is
31 provided in **Table 2**.

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33 Coherent Raman scattering techniques require the use of two incident laser beams, termed the
34 pump and Stokes laser. In most cases, the pump laser is a tuneable laser across the range 700 -
35 990 nm generated by an optical parametric oscillator, whilst the Stokes beam is a fixed wavelength
36 laser, usually 1031, 1040 or 1064 nm. The two beams are spatially and temporally overlapped and
37 focussed onto the sample. When the frequency difference between the two lasers is tuned to match
38 a vibrational mode within the sample, coherent Raman scattering processes can occur.

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40 CARS was first described in 1965 [37] and generates an anti-Stokes scattering signal through four-
41 wave mixing of the incident pump and Stokes lasers (**Figure 1a**). When the difference in the

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2 frequencies of the pump and Stokes beams matches a molecular vibrational frequency (Ω_{vib}) in the
3 sample, a non-linear interaction between the incident photons causes vibrational resonance of the
4 chemical bond in the sample. The excited vibrational chemical bond further interacts with an
5 additional pump photon, resulting in the coherent generation of an anti-Stokes photon (ω_{as}). The
6 CARS signal is technically simple to detect using a photomultiplier tube because the signal is blue-
7 shifted relative to the incident laser wavelengths, with the use of bandpass filters enabling the
8 detection of the CARS signal. However, non-resonant background and resultant spectral distortions
9 can occur during image acquisition. The nonlinear electronic response of the molecules at laser
10 focus can generate a non-resonant background signal in the absence of a vibrational resonance.
11 The interference between non-resonant background and resonant vibrational signal can distort the
12 CARS image and CARS spectrum thereby limiting the detection sensitivity and specificity.

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SRS was first reported in 1962 [38] although it was not until 2008 that the first demonstration of SRS
microscopy for biological samples was realised [39]. When the frequency difference between the
pump and Stokes photons is matched to a vibrational frequency in the sample, stimulated Raman
loss (SRL) is observed in the pump beam whilst stimulated Raman gain (SRG) is detected in the
Stokes beam (**Figure 1b**). The detection of SRS is achieved by detecting the SRL signal or SRG
signal which is typically small (of the order $10^{-7} - 10^{-4}$) in the pump or Stokes beam [40] respectively
using a modulation transfer scheme via lock-in amplifier. Thus, when the frequency difference
between the pump and Stokes beam does not match a vibrational frequency in the sample, there is
no energy transfer and hence, SRL and SRG cannot occur. As such, SRS is free from non-resonant
background and SRS spectra are free from spectral distortion unlike CARS, which makes SRS an
attractive method for hyperspectral imaging applications for material characterisation (**Figure 1c**).

Whilst SRL and SRG signals are bond-specific, these signals are weak and potentially lower than
the fluctuation of laser noise, which posed a significant challenge for the implementation of SRS
microscopy. As laser noise primarily occurs at low frequencies (generally <1 MHz), SRS microscopy
is typically detected using high frequency modulation (>2 MHz). In most SRS systems, the Stokes
beam is modulated at a frequency greater than 2 MHz and the pump beam remains unmodulated.
Upon interaction with the target chemical bond in the sample, the SRL SRG signals occur at the
same frequency of modulation as the Stokes beam. Thus, it is possible to detect the SRL signal at
the modulation frequency of the Stokes beam which is de-modulated *via* lock-in amplifier (**Figure
1b**). Given that the Stokes beam is usually a fixed wavelength laser, in most cases, the SRL signal
in the pump beam is detected, as the Stokes beam can be easily removed using a filter. An important
advantage of SRS is that the SRL signal is directly proportional to the concentration of the target
molecule in the sample enabling straightforward quantification. In addition, the SRS spectrum
matches the Raman spectrum directly to enable direct sample characterisation. The detection
sensitivity of SRS microscopy is ~ 10 nM for endogenous biomolecules including C-H vibrations
indicative of protein and lipid species [41]. The development of bio-orthogonal Raman tagging has

1 pushed the detection sensitivity by employing chemical groups like alkynes and nitriles which have
2 a large Raman scattering cross section. Alternatively, the use of plasmonic materials including gold
3 nanoarchitectures to enhance the SRS signal have been reported [42], together with the use of
4 organic dyes to improve the detection sensitivity via electronic pre-resonance enhancement [43].
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9 **2.3 Unmixing the data.** Raman imaging methods typically acquire a full or partial Raman
10 spectrum at pixel locations from a pre-defined sample area of interest. In doing so, the result is a
11 three-dimensional dataset with coordinates of x and y for the spatial area and the spectral information
12 contained in the Z axis. The creation of a Raman image from the intensity of a Raman band or ratio
13 of multiple bands is easy to do. Alternatively, a wide variety of chemometric analysis techniques
14 have been reported that have the capability to cluster Raman spectra or classify pixels within the
15 image based on their spectral profile. Notable examples include principal component analysis
16 (PCA), k-means cluster analysis (KMCA) and hierarchical cluster analysis amongst others. For a
17 review of different spectral processing methods for vibrational spectroscopy, see [44].
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20 In the case of coherent Raman scattering techniques, vibrational spectroscopic information can be
21 acquired to provide molecular compositional information on the sample. Typically, images are
22 acquired across a portion of the spectrum by re-tuning the pump laser in between image frames.
23 The resulting series of images can be stacked together, and each pixel location will contain a pseudo-
24 Raman spectrum that can be used for molecular characterisation. This is referred to as hyperspectral
25 imaging and is a popular method for SRS microscopy of biological samples. As with Raman
26 scattering, several multivariate analysis methods have been reported to unmix hyperspectral SRS
27 datasets mainly because the C-H stretching region is highly overlapping and many biomolecules
28 have similar Raman spectra. Spectral unmixing using spectral phasor analysis, PCA, multivariate
29 curve resolution, and least absolute shrinkage and selection operator (LASSO) are popular methods
30 for unmixing hyperspectral SRS imaging datasets to enable the detection of discrete features within
31 a sample.
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Spectral phasor analysis uses the Fourier transform to project every pixel from a 3D hyperspectral
SRS dataset onto the 2D phasor plane. Spectral phasor is a convenient method because it requires
no prior knowledge of the sample, and clustering of the phasors based on spectral similarity can be
used to identify regions in the sample which have similar, or the same, Raman spectral profile. The
first example of spectral phasor analysis applied to hyperspectral SRS imaging data was reported in
2014 for label-free detection of cellular organelles [45], and since then it has been applied widely to
study cellular and tissue composition [46-50], bioorthogonal Raman probes [51, 52] and protein
aggregates [53].

PCA has been applied to SRS imaging datasets for dimensionality reduction by creating lower-
dimensional orthogonal projections that display maximum data variance. PCA was selected for
component identification in hyperspectral flow cytometry using SRS microscopy.

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2 K-means cluster analysis (KMCA) is a method that groups spectra based on similarity. Initially, the
3 operator selects the number of clusters into which the spectra will be assigned, whereupon each
4 spectrum is assigned based on their spectral similarity to the seed locations. Each spectrum in the
5 Raman or SRS hyperspectral stack is assigned to the cluster with the closest match to the mean
6 spectrum of the cluster. KMCA is routinely used to identify cellular organelles using Raman
7 spectroscopy [54, 55] and with SRS imaging for the localisation of cellular lipid droplets [27].

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12 Multivariate curve resolution is an alternative method which can decompose a measured spectral
13 dataset into concentration maps and spectral profiles of the principal components. It is often coupled
14 to alternative least squares (ALS) to unmix hyperspectral SRS imaging, including for cellular
15 segmentation and the analysis of lipid-rich regions, see section 4.5 below which describes this
16 approach in the model organism *C. elegans* [56].

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LASSO regression is a powerful technique in predictive analytics that introduces a penalisation factor
to reduce overfitting by selectively shrinking some spectral coefficients to zero. In doing so, it can
enhance the interpretation of large and complex hyperspectral datasets and has been successfully
applied to studying cholesterol and unsaturated fatty acids in living cancer cells using hyperspectral
SRS imaging (see 5.2 below and [57, 58]).

3. What does SRS/Raman tell us that other approaches do not?

As an optical technique, Raman microscopy is ideally suited for live-cell imaging. The use of near
infrared laser sources can help to minimise sample damage and improve tissue penetration depth
for imaging in complex sample types. A key advantage of Raman and SRS microscopy for cellular
imaging applications is the fact that the detection of cellular biomolecules including proteins, lipids
and DNA can be achieved based on the intrinsic Raman spectrum and without any additional
labelling. As such, this is a major advantage to other optical imaging techniques based on
fluorescence microscopy which require the use of reporters, stains or fluorescently labelled proteins
to generate image contrast. Additionally, the linewidth of Raman peaks is much narrower than
fluorescent counterparts ($\sim 20 \text{ cm}^{-1}$ vs. $\sim 1500 \text{ cm}^{-1}$) which enables a much greater multiplexing
capability for cellular imaging [59]. Recently, using engineered polyynes as Raman reporters in the
cell-silent region (see figure 1) of the Raman spectrum ($1800\text{-}2800 \text{ cm}^{-1}$), the detection of 26 discrete
species was achieved in a super-multiplex imaging experiment using SRS microscopy [60]. Given
the broad and featureless nature of fluorescence emission, there exists a colour barrier beyond
which the multiplexing of more than 5 spectral colours poses a significant challenge. A particularly
attractive feature of Raman reporters is that they can be extremely small by design (down to a single
chemical bond) which invokes a greatly reduced perturbation on the parent molecule. This has
opened up opportunities for imaging drugs and small molecules using miniaturised Raman reporters
based on alkynes and nitriles in a way that is not achievable using other optical imaging techniques



[61]. Furthermore, because the Raman stretching frequency is defined by the atoms which make up the molecular bond, the use of isotopic editing strategies for labelling and metabolic probing has been extremely fruitful. For example, the use of deuterium as a natural isotope of hydrogen results in a red shifting of the C-D stretching (2000-2300 cm⁻¹) compared to C-H stretching (2800-3100 cm⁻¹). As such, extremely minor modification of low-molecular weight metabolic precursors and drugs enables their visualisation in the cell-silent region of the Raman spectrum using deuterium labelling (e.g. see 4.2 and 4.4 below). Further expansion of the isotopic colour palette for Raman microscopy has demonstrated the impact of ¹³C labelling of alkynes and ¹⁵N probing of nitrile groups to increase the spectral colours available for imaging [62]. When compared to SILAC labelling for Mass Spectrometry (MS) and proteomics experiments, Raman imaging offers the spatial resolution for probing metabolism at the subcellular scale in a way that is unachievable using these techniques. In that way, coupling Raman microscopy with metabolomics studies can offer a unique insight into probing cellular metabolism in a complementary fashion.

Historically, the detection of Raman images was too slow to be competitive with fluorescence imaging modalities, although advances in SRS microscopy have enabled rapid acquisition and high-content imaging for biomolecular characterisation which is beginning to have a translational impact in biological and medical research. With the advent of commercially available SRS imaging systems, the potential for further integration of coherent Raman imaging in biological and medical imaging is anticipated. Some new advances in methodology and analysis are outlined in Section 5 below.

4. Raman-led insights into lipid and fatty acid metabolism.

In this section, we present some studies of interest which reveal how Raman-based imaging can be employed to gain insight into aspects of fatty acid or lipid metabolism. We recognise this is neither an exhaustive list, nor is it entirely systematic. Our approach is to present a few very recent studies to reveal the kinds of questions that Raman can answer, and to illustrate these using different biological systems. More exhaustive reviews of both lipid metabolism [25, 63], adipocyte biology [10, 11] and Raman [64-67] can be identified.

4.1 Studying lipid biology in adipocytes using Raman imaging. Molecularly specific information on lipid metabolism can reveal novel cell phenotypes in health and disease.

Lipid droplets are now recognised as highly dynamic organelles which play a major role in cellular and systemic homeostasis, and which contribute to disease aetiology, for example ectopic accumulation of lipid droplets in cells such as the liver or pancreatic beta cells are thought to be early drivers of insulin resistance [17, 19, 21]. Lipid droplets are usually studied or identified using fluorescent lipid-soluble dyes which accumulate in droplets and can thus be quantified. While useful, these studies cannot offer insight into droplet chemical composition and thus potential heterogeneity and may provide a flawed index of droplet size because of the need for the fluorescent dye to partition unfirmly



1 throughout the droplet. The former has been to some extent overcome using MS, but these
2 undoubtedly powerful molecular approaches lack capacity for high throughput, cannot inform on
3 spatial heterogeneity, are destructive and importantly are almost impossible to use for real-time
4 dynamics (see [2] for a recent review).
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8 The application of Raman to droplet biology exemplifies the power of Raman-based imaging. The
9 number and area of lipid droplets formed during differentiation in human adipocytes has been
10 quantified and mapped using label-free SRS imaging [68]. This is of interest because larger
11 (hypertrophic) adipocytes are associated with metabolic disease risk [69, 70] and may even act as
12 a reservoir for SARS-CoV2 [71], perhaps explaining the strong link between disease outcome and
13 obesity during the pandemic. Consistent with this notion, a recent study considered the structural
14 and functional changes in adipose tissue in rodent models of obesity and type-1 diabetes [72]. This
15 study analysed Raman spectra using direct band analysis, ratiometric analysis and chemometric
16 methods to identify significant spectral differences between adipocytes from control and Type-1
17 diabetes rodent models, and changes in unsaturation of lipid levels associated with obesity in
18 different fat depots [72]. These non-invasive methods provide a framework for future biological
19 experiments to understand how changing lipid droplet biology relates to function and/or disease risk
20 and begin to show how molecularly specific information can be gleaned from living cells.
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This approach has been further advanced by Tratwal *et al.*, who employed Raman micro-
spectroscopy to examine the subtleties of bone marrow-derived adipocyte biology [73]. Bone marrow
adipocytes (BMAds) are classically observed to be comprised of two distinct subtypes (constitutive
cBMAds and regulated rBMAds) which differ in the content of saturated fatty acids, the number and
size of lipid droplets and biological function [74, 75]. These different populations are notoriously
difficult to isolate and cell culture models hard to validate. Using lipid profiling at the cell population
level and Raman micro-spectroscopy on single cells, Tratwal and co-workers validated the
adipogenic potential of a bone marrow-derived stromal cell line, OP9 cells [73]. Previous studies
based on population analysis has suggested that the 16:1/16:0 fatty acid unsaturation ratio is a
discriminating index for c- and rBMAds. Raman micro-spectroscopy revealed a predominance of
unsaturated spectra after adipocyte induction; importantly, using the power of SRS they were able
to analyse the heterogeneity of *single lipid droplets* within a single cell and show that the lipid droplets
accumulated were not composed of uniformly saturated or unsaturated lipids, but rather were
characterised by different mixtures [73]. Using unguided hierarchical cluster analysis of the SRS
signal from 100s of individual lipid within so-called induced OP9 cells (i.e. cells exposed to
adipogenic induction media; iOP9) compared to spontaneously differentiating OP9 cells (sOP9), they
found that iOP9 adipocytes not only contained larger lipid droplets but that these contained a greater
frequency of unsaturated lipids than sOP9 adipocytes (**Figure 2**). Based on previous studies this
suggests that iOP9 adipocytes are similar in characteristics to rBMAds, and thus have opened the
way to the analysis of the cell biology and role of these different populations.

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2 These kinds of non-invasive approaches to study lipid or fatty acid metabolism are now gaining
3 considerable traction in adipocyte biology as scientists appreciate the molecularly specific insight
4 these approaches can deliver [26, 68, 72, 73, 76-78].

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7 **4.2 Single cell SRS provides a new insight into lipid metabolism associated with**
8 **phenotypic changes in a subset of tumour cells.** It has long been appreciated that cancer cells
9 exhibit distinct metabolic profiles to their non-transformed counterparts [79, 80]. Therapeutic
10 resistance is a major challenge to oncology treatment. Studies have suggested that the transition
11 from a drug-sensitive to a drug-resistant state is accompanied by changes in cellular metabolism,
12 but this has proven difficult to study as typically within a given population of cells only a small fraction
13 become drug-resistant [81, 82]. A good deal of recent work has focussed on understanding the
14 change from glycolytic to fatty acid uptake and oxidation and how this relates to cancer cell function
15 [79, 83-85].

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17 A recent paper has adapted stimulated Raman scattering and spectral phasor analysis to provide
18 single cell insight into metabolic reprogramming observed in cancer cells [7]. In a striking example
19 of the power of SRS, Tan *et al.* employed high throughput, large-area hyperspectral SRS imaging to
20 resolve changes in the metabolism of single cells within a population [7]. This approach involved
21 capture of a stack of large-area hyperspectral images (200-500 cells), extraction of the SRS
22 spectrum at each pixel and segmentation to generate maps of intracellular compartments.
23 CellProfiler® was then used to generate maps of the cell boundaries, allowing the lipid map of
24 individual cells to be identified. This in turn facilitated quantitation of the lipid metabolic signature on
25 a per-cell basis. Married to the use of specific glucose or fatty acid non-invasive Raman probes, this
26 approach allowed the authors to show that tumour cell drug-resistance (in this case to cisplatin) was
27 accompanied by a molecular switch – a shift from glucose/glycolysis-dependent metabolism to fatty
28 acid oxidation as a key indicator of a shift in drug-resistant cancer cells (**Figure 3**) [7]. As well as
29 providing new molecular understanding, this approach hints at a potential diagnostic strategy for
30 identifying and treating cisplatin-resistant tumours. It also powerfully illustrates the insight that
31 individual cell imaging within a population can provide and exemplifies the power of SRS since
32 images from dozens of cells can be acquired quickly and are data rich. Notably, this group also used
33 imaging approaches to quantify reactive oxidative species on a per cell basis [7]. The potential to
34 marry this with SRS analysis in the same experiment is worthy of further exploration. Such studies
35 signpost the power of SRS to study specific metabolic changes in a non-invasive system, allowing
36 the subsequent multiplexing of analysis with other signals, or indeed with other techniques such as
37 single cell metabolomics [86], lipidomics [87] or other MS-based methods [2].

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39 **4.3 Spectral resolution of specific metabolites in lipid droplets in melanoma.** Metastatic
40 melanoma is a deadly form of skin cancer, many of which contain a mutation in BRAF kinase [88].
41 A remarkable application of Raman imaging to understand cancer cell biology involved a series of
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1 melanoma cell lines, each encoding different BRAF mutations, corresponding to a different level of
2 cancer cell differentiation and accompanied by differential susceptibility to inhibitors and
3 immunotherapeutic interventions [6], each with a well annotated transcriptome [89]. Based on
4 spontaneous Raman spectra, it was found that the lipid/protein (CH₂/CH₃) ratio decreases with the
5 progression of de-differentiation. Based on this observation, SRS imaging in real-time was employed
6 to quantify single cells and thus capture cellular heterogeneity [6]. Targeting the lipid peak (2845cm⁻¹,
7 CH₂ vibrations) and the protein peak (2940cm⁻¹, CH₃ vibrations), the ratiometric images clearly
8 showed a decrease in lipid signals from melanocytic to mesenchymal cells, indicating that the more
9 differentiated cells are enriched in lipids. This observation strikingly correlated with transcriptomic
10 data, and therefore gives confidence that single-cell Raman imaging correlates with transcriptional
11 profiles. Using glucose-d7, the same group quantified *de novo* lipogenesis in the different cell lines
12 and observed that *de novo* fatty acid synthesis is most active in differentiated cell lines; thus, single-
13 cell Raman imaging has uncovered 'phenotype-specific druggable susceptibilities in cancer cells'
14 [6].

15 Not content with this remarkable set of observations, Du *et al.* hypothesised that the maintenance of
16 specific cancer cell characteristics could require specific examples of lipid biochemistry that could
17 offer insight for new therapies. By way of a proof of concept, they showed that analysis of individual
18 lipid droplets in one of the cell lines exhibiting a mesenchymal phenotype contained metabolic
19 activities associated with lipid unsaturation (**Figure 4**). They propose that lipid droplets in these cells
20 act as a reservoir of unsaturated fatty acids; depletion of this pool eventually drives apoptosis [6].
21 The data itself is exciting and provocative, but arguably just as important is their observation that
22 'the mechanism and applicability underlying reported susceptibilities in our work are distinctly
23 different from previous reports that relied on bulk analysis. This demonstration thus emphasises the
24 unique value of subcellular pharmaco-metabolomics as a revelatory tool for uncovering new cell
25 biology' [6]. We believe that Raman-based metabolomics is set to revolutionise biology as new tools
26 and image processing algorithms are developed.

27 **4.4 New insight into metabolic ageing in *Drosophila*.** *Drosophila* are a favoured model to
28 study the biology of ageing due to their conserved signalling pathways and short life span coupled
29 to excellent genetic tools, high throughput and low cost [90-92] thereby providing insight into how
30 lipid metabolism directly impacts the ageing process [93, 94]. Metabolism in ageing fruit flies has
31 been a point of interest due to the metabolic differences reported in ageing humans and how
32 differences in diet impact progression in ageing. Recent studies using Raman have added credence
33 to changes in lipid metabolism underscoring effects of ageing.

34 Comparison of the SRS signals of fat bodies from 7 and 35 day old flies revealed a significant
35 increase in retinoid presence (similar retinoid level increases have also been observed in *C. elegans*
36 [95]) and a decrease in unsaturated fatty acid levels in 35 day old flies compared to their younger
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1 counterparts [96]. While important, these observations are limited to a snapshot of levels at a given
2 time. Shi and colleagues therefore developed an approach to quantify changes in lipid turnover by
3 incorporating heavy water (D_2O) into the diet, allowing *de novo* lipogenesis to be measured (a
4 technique they christened DO-SRS) [96, 97]. As deuterium (D) incorporates into the newly
5 synthesized lipids, the SRS silent region can be utilized to quantify C-D bond formation. Alongside
6 reduced lipid droplet size, older (day 35) flies showed reduced lipogenesis across all diets when
7 subjected to ratio metric imaging of existing lipids (2850 cm^{-1}) and deuterated lipids (2140 cm^{-1}) in
8 comparison to younger flies (**Figure 5**). These data suggest that larger lipid droplets are more active
9 and may indicate that large and small lipid droplets play different roles in metabolism. This has
10 provided further impetus to the drive to study metabolic changes on single cells, rather than
11 populations, and clearly illustrates that all lipid droplets are not the same [96, 98].

12 The ability to 'track' deuterated molecules is a highly favourable aspect of SRS by allowing
13 visualisation of individual lipid droplet metabolic activity and appreciation of the heterogeneity of fat
14 depots [96, 97]. This has been strikingly applied to studies of how diet composition, caloric restriction,
15 and metabolic activity impact aging [96, 99] where a combination of Raman and SRS have provided
16 a wealth of insight allowing quantification of lipid droplet size and distribution. The new studies
17 alluded to above now extended this toolkit to allow quantification of metabolic activity at the level of
18 single cells and even single lipid droplets. These advances will likely open-up further insights when
19 applied across other metabolic and compartmentalised processes.

20 Two further studies along similar lines are worthy of highlight. In the first, harnessing the power of
21 *Drosophila* genetics and DO-SRS with deuterated probe molecules, in this case deuterated glucose
22 and deuterated acetate, Li *et al.* observed that caloric restriction and down-regulation of the insulin
23 signalling pathway was accompanied by a shift in brain metabolism to use acetate as a major carbon
24 source for lipid synthesis, and provided an elegant analysis of lipid turnover at the level of single lipid
25 droplets [100]. These studies will no doubt pave the way for more detailed metabolic studies across
26 a range of tissues and organisms by exemplifying the novelty and detail that Raman can provide. In
27 a second study, also in flies, Spratt and colleagues used deuterated methionine to study regional
28 and cellular specific uptake and incorporation of a specific amino acid during fly development [101].
29 This study is worthy of comment for two reasons. First, the weaker vibrational resonance of D_2O can
30 limit the potential signal, meaning that for studies of less abundant species, it may not be an ideal
31 choice as levels of incorporation may be too low for reliable quantification (this is often not the case
32 for lipids, as they accumulate in lipid droplets, concentrating the lipids into a defined location allowing
33 ready identification). Secondly, this group married SRS and fluorescence in an integrated
34 microscope, allowing the investigation of specific cells within a complex mixture that could be
35 identified using other labels [101]. This highlights a further potential advance in this rapidly moving
36 field.

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2 **4.5 Understanding drug resistance in *Candida albicans*.** Antifungal resistance is a significant
3 threat; fungal infections cause major diseases such as cryptococcosis and aspergillosis, chronic
4 conditions such as allergic bronchopulmonary aspergillosis and a range of less-threatening
5 conditions such as vaginitis or candidiasis [102, 103]. *Candida albicans* – a fungal pathogen – is a
6 contributing factor in hospital acquired infections and continued use of a first-line therapy
7 Fluconazole (the prototypical azole drug) has resulted in antifungal resistance within this strain [102].
8 Fluconazole disrupts the ergosterol biosynthesis pathway and results in accumulation of sterol
9 14,24-dimethylcholesta-8,24(28)-dien-3 β ,6 α -diol which is toxic to the fungus [104]. *C. albicans* can
10 form biofilms, a surface associated fungal community, which presents significant clinical issues in
11 hospitals, e.g. in catheters etc [105].
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SRS has provided insights into this antifungal resistance by identifying that lipid droplets are formed via *de novo* lipogenesis within these biofilms during the stationary phase [104]. Zhang *et al.* describe a novel approach to understanding the composition of these lipid droplets using pixel-wise least absolute shrinkage and selection operator (LASSO; see section 2.2) regression of the hyperspectral SRS image stack to 'decompose' this into specific chemical 'maps' [95]. Comparison of these chemical maps revealed differences between azole-sensitive and resistant strains of *C. albicans*, specifically upregulated levels of ergosteryl ester was detected in fluconazole resistant *C. albicans* strains. This was reflected in the sterol C=C peak (1603 cm⁻¹) and acyl C=C peaks (1655 cm⁻¹), absent in wild type strains. These were confirmed by referencing the spectra from pure ergosterol and glycerol trioleate and studying these chemical maps under conditions such as disruption of glycolysis. These data indicate that the accumulation of esterified ergosterol was a consequence of *de novo* lipogenesis [95], and suggests the hypothesis that inhibition of ergosterol esterification could be a viable therapeutic option. In an elegant set of experiments, Zhang *et al.* showed that treatment with oleic acid reduced the accumulation of ergosteryl ester in resistant *Candida* strains which in turn re-sensitised *Candida* to Fluconazole identifying a novel therapeutic strategy to deal with resistance [95]. The molecularly specific insight that Raman can provide underscores these exciting developments.

4.6 Fatty acid production in biotechnology. While many of us are familiar with the concept of using bacteria as a source of chemicals and drugs in biotech, what is seldom well-appreciated by those not directly in the field is the difficulty faced by companies because of both cell-to-cell and genetic variation that can occur during the fermentation processes that underpin these methods. Hence, a crucial requirement for the biotech field is an ability to quantify this variability, ideally in real-time. A recent study has shown that hyperspectral SRS offers a powerful technique for monitoring the fermentation process [106]. This is challenging in bacteria due to their small size which this shortens the axial signal integration length and so results in weaker SRS signals than are typically observed for larger cells. At the heart of the method developed by Tague *et al.*, hyperspectral SRS (section 2.2) was coupled to a novel computational analysis technique that can



1 generate predictions of chain length and saturation. After validating this using MS, they were able to
2 optimise imaging to capture information from living cells, and by marrying SRS and time-lapse phase
3 contrast imaging with automated image segmentation approaches, demonstrated a correlation
4 between fatty acid production and growth, revealing metabolic and temporal heterogeneity in actively
5 growing cultures [106]. Importantly this was achieved by recoding from the same cell over a period
6 of many hours. This work is sure to gain attention across fermentation biology-driven systems.
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14 **5. Examples of new Raman method developments in the metabolic toolkit.**

15 The approaches outlined above are characterised by the application of increasingly sophisticated
16 Raman-based approaches in different biological systems, chosen here to illustrate how these kinds
17 of approaches can be leveraged to gain new biological insight. Now, we turn our attention to one or
18 two novel examples of Raman technology development which we believe new advances offer the
19 scope for a paradigm shift, either in imaging, the biology it illuminates, or both.
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5.1 **Identification of specific lipid species in the absence of chemical labelling by pushing the spectral resolution limit.** Hyperspectral stimulated Raman scattering (SRS) microscopy has the potential to provide comprehensive lipid characterization, including carbon chain length quantification. However, the current spectral resolution of SRS microscopy is limited to about 10 cm⁻¹, which is not sufficient to resolve the subtle Raman shifts corresponding to chain lengths, e.g. between specific fatty acid species. This is important because population-level studies suggest that the degree of unsaturation and chain length differ in cancer cells, but how these relate to specific subsets of cells is unclear, largely because of an inability to identify chain lengths on a microscopic scale. Huang *et al.* have pushed the spectral resolution limit of hyperspectral SRS microscopy to 5.4 cm⁻¹ by employing a highly efficient spectral compressor, which enabled them to differentiate eight types of saturated lipids with carbon chain lengths from C8:0 to C22:0 [107]. Future applications of this approach would involve application to living cells where it has the potential to reveal the role of specific lipid/fatty acid molecules across a range of experimental systems.

5.2 **Single cell metabolism in cancer cells by high-content hyperspectral SRS-imaging.** As noted above and elsewhere, hyperspectral SRS (hSRS) can resolve multiple spectral metabolites. This has been reported at sub-second imaging speeds and with spectral coverage of >200 cm⁻¹ and a spectral resolution of less than 10 cm⁻¹ (see 5.1). Together with downstream spectral analysis methods, such as least-squares fitting and phasor segmentation, hSRS has revealed many new facets of biology, some of which were discussed above. However, many metabolites remain challenging to study, either because their signals are too weak or because their characteristic signals are either difficult to resolve or dominated by other biomolecules with strong Raman signatures. Recognising this limitation, a new approach has been developed and applied to cancer cells – high content, hyperspectral SRS (h²SRS)[57].



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2 Low sensitivity issues can be overcome in SRS approaches by focussing on the C-H high wave
3 number bands. However, this presents a problem, as all major metabolic species (proteins, lipids,
4 nucleic acids, glucose etc.) exhibit substantially overlapping peaks within this region which cannot
5 be resolved using existing hyperspectral data analytics approaches. This arises because of the
6 crosstalk between the C-H vibrations in the chemical maps. Tan *et al.* [57] reasoned that only a few
7 metabolic species make dominant contributions at any given point, a condition that can be
8 incorporated into the analysis of these signals as a sparsity constraint, a mathematical device in
9 which a model is designed to use only a small finite number of significant features at a time,
10 disregarding less relevant ones (this approach is gaining traction across many facets of biological
11 imaging [108-110]).

12
13 Using h^2 SRS, Tan *et al.* were able to identify metabolic alterations in cancer cells in response to
14 chemotherapy in single living cells. Specifically, increased intracellular fatty acid and carbohydrate
15 levels in brain cancer cells after cisplatin treatment and in the pancreatic cell model MIA PaCa-2
16 after gemcitabine treatment. This extraordinary study offers the potential to unravel many novel
17 facets of cellular metabolism on a single cell scale using non-destructive approaches. As they note,
18 because 'the unmixing algorithm is a supervised approach that requires prior knowledge of the
19 chemical composition, more detailed prior information (metabolite composition and abundance)
20 obtained through MS can effectively guide the selection and scaling of input references beyond the
21 current [five] components.' This powerful marriage of MS with live cell imaging using chemical
22 probes, coupled to high-throughput and single cell resolution, hints at a future in which multiple
23 metabolites can be studied non-invasively and with increased specificity in living cells, providing a
24 step-forward in our understanding of single-cell metabolism.

25
26 **5.3 Super-resolution meets SRS?** A further example of the power of SRS is provided by the
27 development of a novel deconvolution algorithm: adaptive moment estimation (Adam) optimization-
28 based pointillism deconvolution (A-PoD) which is illustrated conceptually in **Figure 6** [111].
29 Deconvolution is a computational approach, widely used in light microscopy, that can improve the
30 contrast and resolution of digital images by removing or minimising distortion present within the
31 images. Several deconvolution approaches are widely applied within fluorescence imaging, but
32 these approaches cannot be applied to single-frame images in low-sensitivity detectors, such as is
33 the case for SRS microscopy images. Building on previous attempts to overcome this issue for SRS
34 [112], Jang *et al.* developed a mathematical model to allow deconvolution and applied this to
35 mammalian cells in culture and to *Drosophila* brain. This approach provided molecularly specific
36 information at a spatial resolution of less than 59 nm on a single lipid droplet, allowing comparison
37 of nanoscopic distribution of lipids and proteins in cells and subcellular organelles which can be
38 applied to *in situ* samples; this approach also represents a step-change in processing time [111].
39 The potential application of this approach to wider aspects of biology offers a further driver for new
40 insight.

5.4 New methods for in situ analysis of fatty acids. Using deuterium isotopes in metabolic studies has offered substantial insight through the history of metabolic studies because the deuterium is eliminated slower than hydrogen: the kinetic isotope effect [113]. This has been elegantly exploited to reveal a tumour-selective cytotoxicity of γ -linoleic acid in VA-13 tumour cells [114]; coupled to the use of an alkyne reporter this offers the potential for distinguishing between lipids with very similar structures (see [65] for a recent review).

A further study coupled Raman microscopy with a novel spectral unmixing software (ImageCUBE) capable of separating overlapping spectra from the mixed measured spectrum within individual pixels [115]. Applied initially to an analysis of five labelled fatty acids (palmitic acid-Br, oleic acid-alkyne, α -linolenic acid-d₁₄, arachidonic acid-d₈, and eicosapentaenoic acid-d₅), these authors were able to show that fatty acids with higher degrees of unsaturation behaved differently and were concentrated in lipid droplets. While the biological mechanism underpinning this behaviour presently is not defined, the identification of this phenomenon suggests that further work on lipid droplet biology should focus on the substrate preferences of the enzymes involved in their biogenesis and hydrolysis. Uematsu *et al* raise the interesting speculation that because polyunsaturated fatty acids are readily oxidized, their presence within lipid droplets may protect from this [115]. Given the importance of fatty acid metabolism in tumour cell biology (see 4.2 and 5.2 above for illustrative examples), this could yet turn out to have far-reaching consequences.

5.5 New tags for multiplex imaging. The challenge facing those wishing to study complex metabolic events, such as those underpinning lipogenesis/fatty acid metabolism, involves unpicking the overlapping spectra and poor signal-to-noise ratio of some of the commonly used chemical reporters. One approach is to use bioorthogonal Raman groups, chemically functionalised to exhibit discrete functional vibration. Presently, these probes lie within the 2000-2300 cm⁻¹ range of the Raman spectrum. A notable recent advance has come from work which uses the B-H stretch of metallacarboranes, detected at 2570 cm⁻¹. The potential to marry these novel probes was evaluated using a metallacarborane B-H stretch in a triplex detection of a bis-alkyne and a deuterated fatty acid, all of which lie within the cell silent region of the Raman spectrum [116]. Murphy and colleagues used this approach coupled to imaging in the high wavenumber region of the cellular Raman spectrum and spectral unmixing of the hSRS dataset to enable 9-colour detection. This offers a remarkable step forward in live cellular imaging studies.

6. Conclusions and outlook.

It must be emphasised that many of the advances outlined here are relevant across many fields of biology. Molecularly specific insight obtained from living cells, organoids and intact tissues offers the potential for unrivalled insight into basic biological control mechanisms. The ability to use the approaches outlined here in model organisms offers huge potential for developmental biology to be

1 coupled firmly to metabolic changes in key defined cells, even within populations. This can offer new
2 understanding of the triggers for developmental events such as cellularisation in flies [117] or be
3 coupled to optogenetic controls in the study of embryogenesis [118]. The marriage of powerful
4 molecular tools to these real-time non-invasive techniques could drive considerable new insight.
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8 The capacity to understand how metabolism is sub-compartmentalised within cells [119, 120], and
9 how this changes in diseases such as obesity, diabetes and cancer, could offer new therapeutic
10 horizons. Driving the limit of SRS to even smaller scales is beginning to offer this kind of advance.
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13 Using real-time read-outs the impact of (for example) environmental factors or toxins on metabolic
14 events can reveal new understanding of cellular behaviour, particularly if married to high specificity
15 molecular analysis. The US National Research Council is driving toxicology screening away from
16 animal studies into cell systems, using metabolomics as a key platform. The potential to couple such
17 work to specific changes in key molecular signatures in real time in living cells offers one exciting
18 way in which SRS-metabolic studies could reveal new understanding.
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21 Advances in hyperspectral SRS imaging have also enabled a detailed molecular characterisation of
22 lipid species based on the Raman spectral signature, and when coupled to chemometric analysis
23 techniques, the potential to identify drug-induced effects and molecular features of disease has been
24 realised. Future development of the technique particularly in improving the spectral resolution (cm⁻¹
25 range) will help to advance studies in this direction.
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28 In the context of cellular imaging, SRS has the advantage over conventional Raman imaging speed
29 which is comparable to fluorescence imaging modalities. This feature has greatly facilitated SRS as
30 a modality for biological microscopy. However, Raman scattering has the clear advantage of
31 collecting a partial of full Raman spectrum at each pixel within an image. Given the favourable
32 spectral resolution associated with Raman spectroscopy, this feat enables the detection of subtle
33 changes that can be used to report on cellular metabolism or subtle biochemical changes that occur
34 within a cellular population. SRS imaging has yet to reach the spectral resolution of conventional
35 Raman microscopy, and this is anticipated to be an area under active investigation. Some early
36 progress towards this aim has been realised through the development of broadband SRS imaging
37 systems employing fast tuneable fibre lasers to enable rapid hyperspectral imaging.
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40 Many studies have demonstrated the live-cell imaging capability of SRS microscopy, however, very
41 few studies have demonstrated the long-term analysis of cellular metabolism during longitudinal live
42 cell imaging. The development of CARS and SRS imaging towards this aim would represent a
43 significant breakthrough for temporal monitoring of drug-cell interactions and cellular metabolism
44 during disease onset and progression. We recently showed it was possible to visualise drug uptake
45 over several hours in the same cellular population using bespoke perfusion chambers to do so [121].
46 As most SRS systems are inverted microscope set ups relying on the detection of transmitted light,
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2 the design of robust chambers that can enable favourable culture conditions of temperature, pH and
3 media exchange over hours-days would be a significant advantage.

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5 Several research groups are beginning to demonstrate the potential of super-resolution SRS
6 microscopy for live cell, label-free imaging at the sub-100 nm scale. Current approaches based on
7 expansion microscopy, deconvolution image analysis, photo-switchable Raman reporters and
8 optical engineering strategies have been developed over the course of the past 5 years. The
9 implementation of artificial intelligence and machine learning methodology to improve the spatial
10 resolution of SRS imaging will likely open new opportunities for studying intracellular lipid biology.
11 Finally, the integration of SRS microscopy with other analytical techniques including photothermal
12 imaging [52] and mass spectrometry [122] offer the potential for studying biological samples with
13 unprecedented detail and with a complimentary read out.

14
15 In sum, we believe that Raman spectroscopy offers an increasingly powerful toolkit for biologists,
16 providing molecularly specific insight at a single cell, and increasing sub-cellular level. Illustrated
17 here by considering aspects of fatty acid metabolism, we contend that these approaches are
18 beginning to rival the advances made by fluorescence microscopy and super-resolution imaging. As
19 increasingly novel probes are developed, together with data processing algorithms, we expect the
20 next decade to provide major advances in single cell metabolic analysis with Raman in the vanguard.

21 **7. Author Contributions.**

22 JCG Conceptualisation; Writing – original draft; Writing – review and editing; Visualization.

23 WJT Conceptualisation; Writing – original draft; Writing – review and editing; Visualization.

24 DG Conceptualisation; Project administration; Writing – review and editing.

25 KF Conceptualisation; Project administration; Writing – review and editing.

26 GWG Conceptualisation; Project administration; Writing – original draft. Visualization; Funding
27 acquisition. Supervision.

28 **8. Conflict of Interest.**

29 The authors declare no conflicts of interest.

30 **9. Acknowledgements.**

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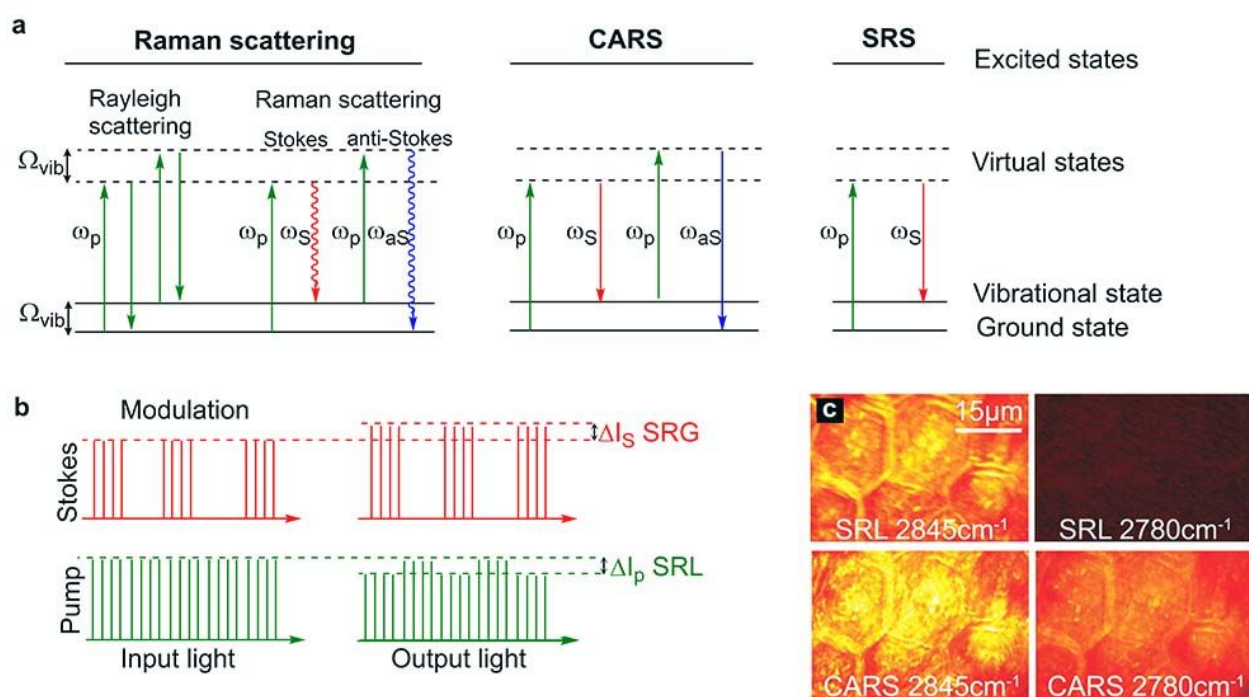
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10. Figures and Tables.

Figure 1. Energy level diagrams for Rayleigh scattering, Raman scattering, CARS and SRS.

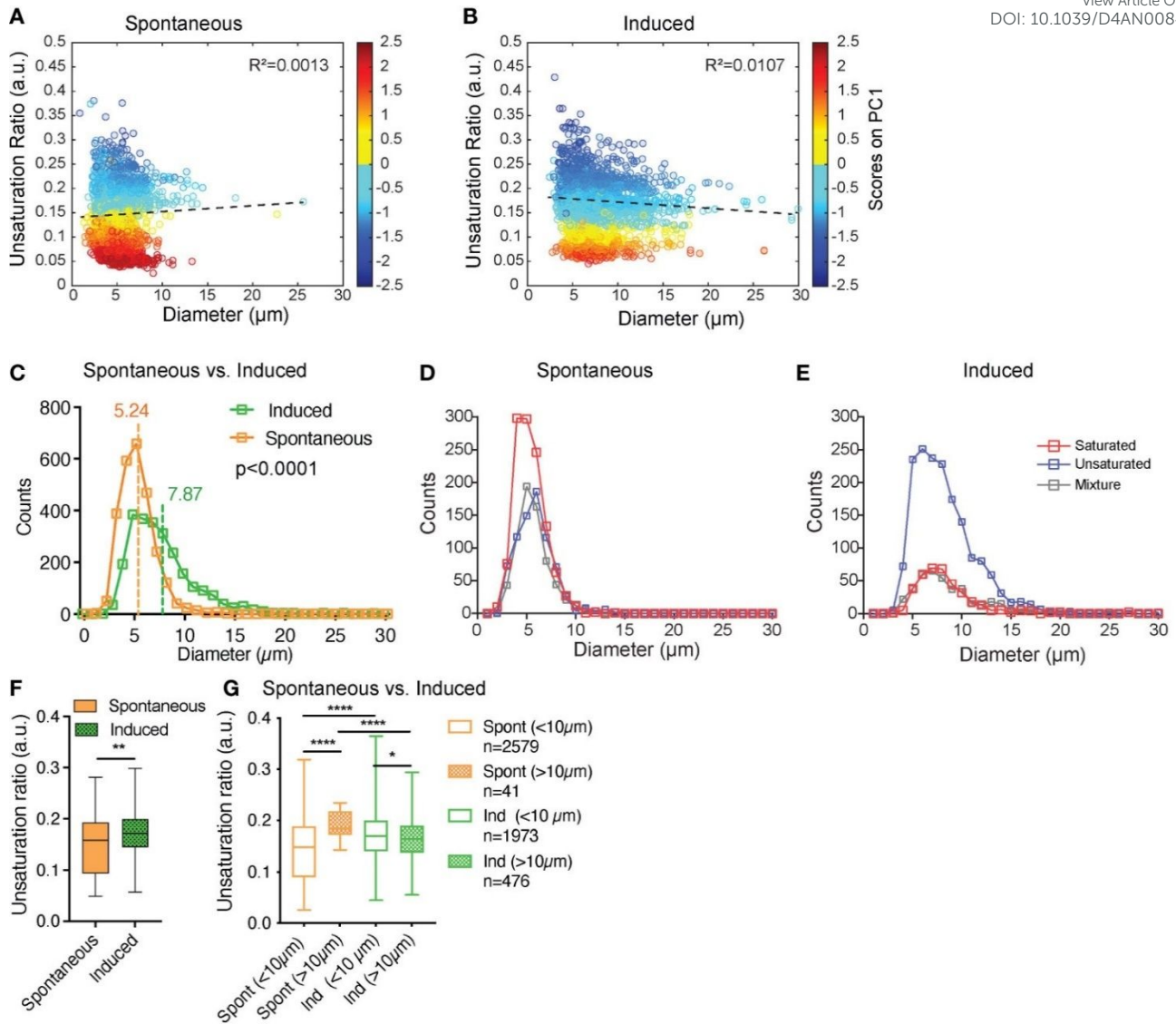


a Rayleigh scattering is an elastic process whereby photons are scattered with the same energy as the incident photons. Raman scattering is an inelastic process where photons are scattered with a lower energy (Stokes scattering) or higher energy (anti-Stokes scattering) relative to the incident photons. In CARS, two incident photons at the pump (ω_p) and Stokes (ω_S) frequencies simultaneously interact with the chemical bond which has a vibrational frequency that is equal to the difference frequency of the two beams. In doing so, the chemical bond is stimulated to a vibrational excited state. Upon interaction of the coherently excited chemical bond with a second pump photon results in the generation of an anti-Stokes photon (ω_{aS}) which is blue shifted (higher energy than the incident photons). In SRS, two photons interact with a chemical bond such that the frequency difference between the pump and Stokes beams matches the energy of the chemical bond. In doing so, the energy of the pump photon is transferred to the chemical bond resulting in the generation of a Stokes photon with a lower energy. **b** Modulation transfer detection scheme. The Stokes beam is modulated at a high frequency which results in a SRL signal that is modulated in the pump beam and a SRG signal that is detected at the modulation frequency in the Stokes beam. The SRL signal is usually extracted from the laser noise using a lock-in amplifier. **c** A comparison of SRS and CARS imaging of mouse skin. The CH_2 vibration of skin lipids is detected at 2845 cm^{-1} , whilst the off-resonance image at 2780 cm^{-1} is free from signal in SRS imaging, non-resonant background is detected in the CARS image. Image adapted and reproduced from Ref. 39 with permission the American Association for the Advancement of Science (2008).

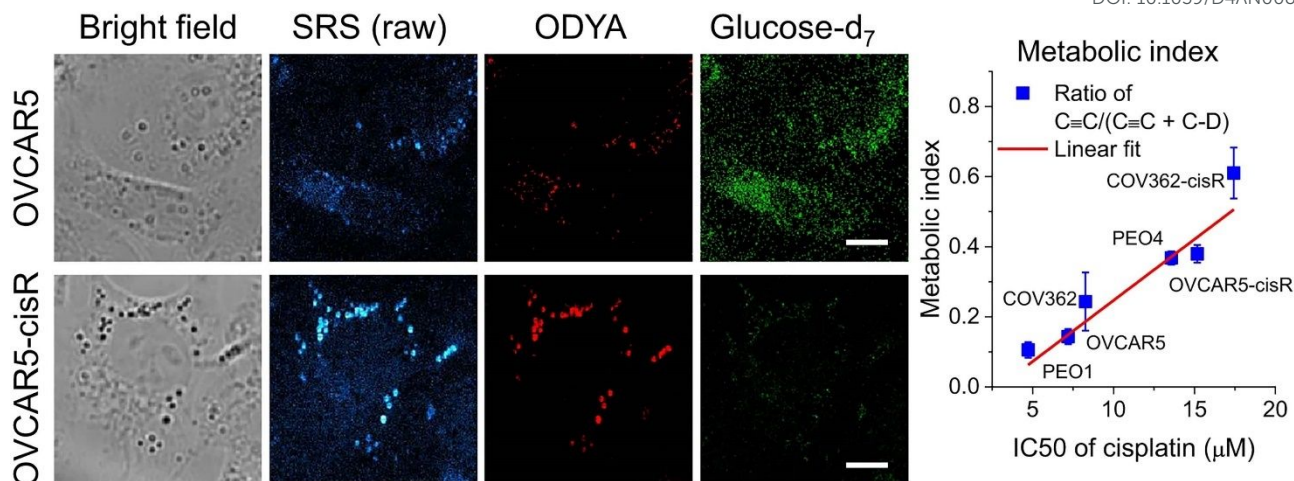
Figure 2 SRS provides insight into individual lipid droplet composition.

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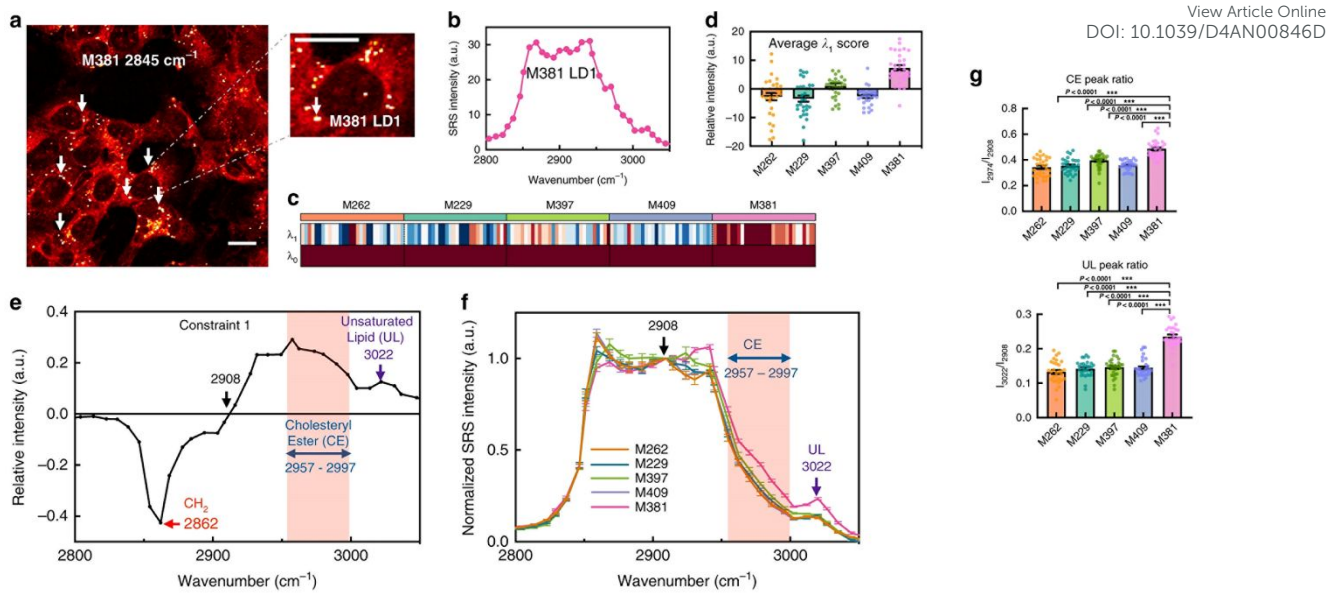
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Tratwal *et al.* examined the composition of individual lipid droplets in bone marrow adipocytes derived from OP9 bone marrow stromal cells, allowed to either differentiate spontaneously or in response to a defined induction cocktail (for details see [73]). **a,b** present the unsaturation ratio as a function of the diameter of the LDs for the two populations of cells. The circles in dark blue corresponds to LDs identified as unsaturated rich. The circles in dark red correspond to LDs identified as saturated-rich. The circles in yellow and light blue correspond to LDs identified as mixture. This Unsaturation Ratio was calculated from Raman microspectroscopy, defining the peak area assigned to unsaturated (C=C bonds) divided by the area of saturated bonds (CH₂), identified using unguided hierarchical cluster analysis. **c** separates LDs as a function of diameter showing that the LDs mean diameter of sOP9-adipocytes is significantly lower compared to iOP9-adipocytes. The power of this Raman based approach is revealed in **d,e** which shows that when separated according to LD type, a trend toward larger unsaturated lipid droplets in the induced condition is observed. **f** shows that the unsaturation ratios of sOP9 versus iOP9 at LD level. LDs from iOP9 adipocytes have a higher mean unsaturation ratio compared to LDs from sOP9 adipocytes. **g** illustrates the power of single droplet resolution: larger droplets (>10 μm) have a higher unsaturation ratio as compared to smaller lipid droplets, while smaller droplets (<10 μm) in induced OP9-adipocytes (n = 1973) have a higher unsaturation ratio than those of spontaneous OP9-adipocytes (n = 2579). Highest unsaturation ratio overall is seen in large droplets from spontaneous OP9-adipocytes (n = 41). Image from [73] via <http://creativecommons.org/licenses/by/4.0/>.

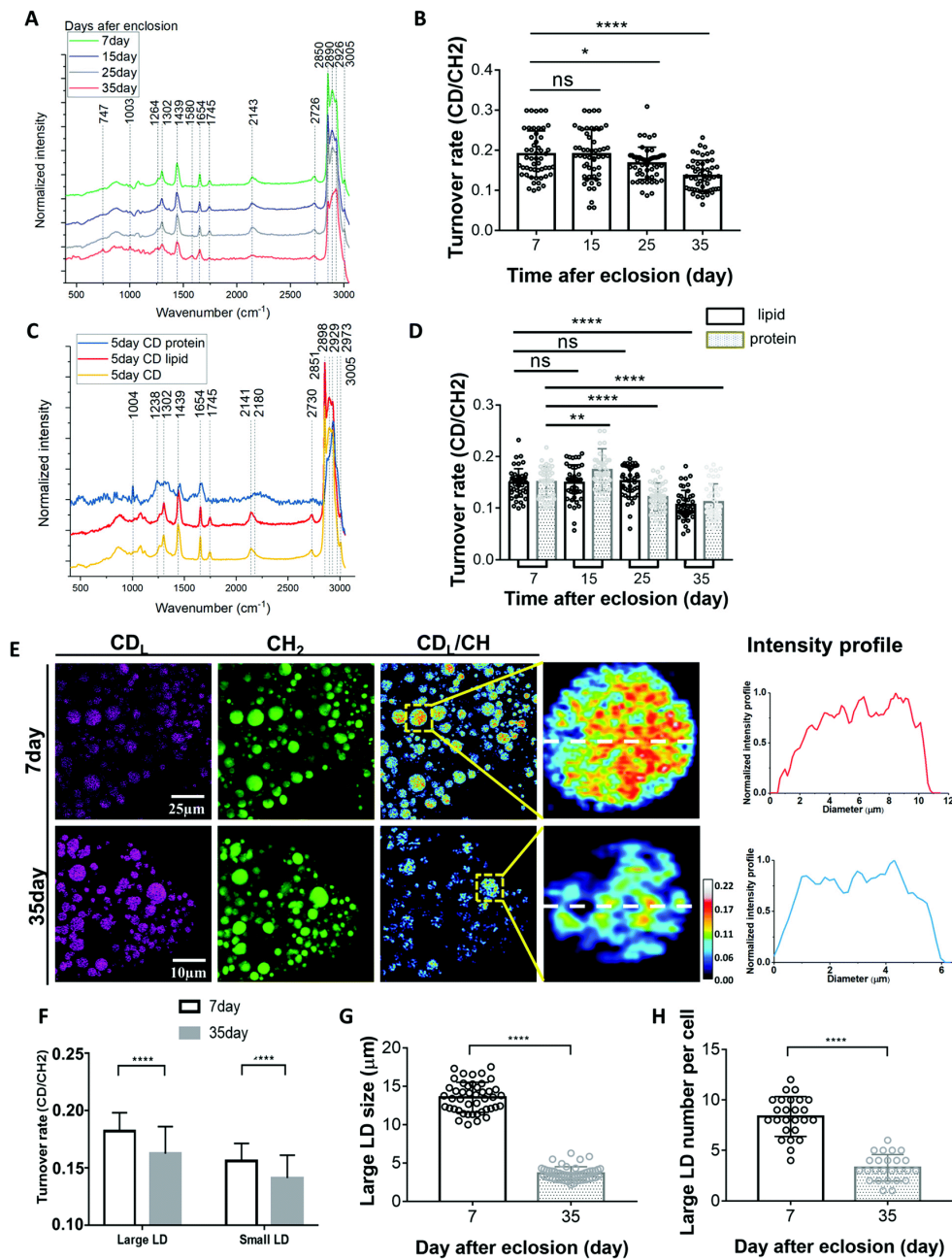
Figure 3 Metabolic phenotypic changes quantified by SRS.View Article Online
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The ability to study distinct aspects of cellular metabolism in living cells distinguished by different Raman probes is exemplified from studies of the consequences of cisplatin resistance in cancer cells. Shown here are data from OVCAR5 cells, a model of ovarian cancer. These cells are used to model metabolic changes, such as those which accompany resistance to cisplatin therapies. Cells from control or cisplatin resistant (cisR in the figure) cells were incubated with glucose-d₇ or 17-octadecynoic acid (ODYA), a fatty acid analogue with an endogenous alkyne allowing simultaneous analysis of glucose and fatty acid metabolism. Shown are representative bright field images, raw SRS images, and processed SRS images of ODYA and glucose-d₇ in OVCAR5 and -cisR cells. This clearly reveals a shift from glycolytic to fatty acid metabolism in the cisplatin resistant cells. By quantifying the metabolic index of $C \equiv C / (C \equiv C + C-D)$ to IC₅₀s of cisplatin in various ovarian cancer cell line pairs (COV362, PEO1 and 4 and OVCAR5), the authors show a shift from glycolytic to fatty acid metabolism which is correlated with the IC₅₀ of cisplatin in the cell type. Figure reproduced from [7] with permission <http://creativecommons.org/licenses/by/4.0/>.

Figure 4 Changes in the biochemistry in individual lipid droplets is associated with disease.

Du *et al.* hypothesised that the maintenance of specific cancer cell characteristics would likely be underpinned by changes in lipid biochemistry, and that hSRS could test this [6]. By way of a proof of concept, they showed that analysis of individual lipid droplets in a melanoma cell line which exhibits a pronounced mesenchymal phenotype contained metabolic activities associated with lipid unsaturation, and some of their data is reproduced here. **a** shows a representative SRS image of M381 cells imaged in the CH_2 (2845 cm^{-1}) channel, the LD signal can thus be clearly identified (arrows). A zoomed-in image highlights a single LD. The authors acquired Raman spectra on LDs from a range of melanoma cells with high spectral resolution (**b**) and extracted key data from these spectra using surprisal analysis. Panel B shows the hSRS spectrum of the zoomed-in LD indicated in panel A at the C–H stretch region ($2800\text{--}3050\text{ cm}^{-1}$). **c** shows a heatmap for scores of the top two constraints ($\lambda_0\text{--}\lambda_1$) by surprisal analysis of hSRS spectra on LDs across five cell lines (M numbers refer to different melanoma lines – see [6]). Each column represents an individual LD and each row represents the constraint scores, with the average score of λ_1 across five cell lines shown in **d**. Molecularly specific information can be obtained from the Raman peak assignments for constraint 1 (λ_1) (**e**). The pink shadowed range from 2957 to 2997 cm^{-1} is assigned to cholesteryl esters (CE), and the 3022 cm^{-1} peak (violet arrow) is assigned to unsaturated lipids (=C–H, UL). hSRS spectra (normalized at 2908 cm^{-1} , the zero point revealed in **e**) of LDs across each cell line is shown in **f**, and then used to compare across multiple cell lines (**g**), which shows quantification of relative CE ($2974\text{ cm}^{-1}/2908\text{ cm}^{-1}$, top panel) and UL ($3022\text{ cm}^{-1}/2908\text{ cm}^{-1}$, bottom panel) enrichment in LDs across different melanoma cell lines. Marrying this insight with transcriptomics data provides powerful new insight. Figure reproduced from [6] with permission <http://creativecommons.org/licenses/by/4.0/>.

Figure 5 D₂O-probed SRS to study metabolic adaption in organs from model organisms.

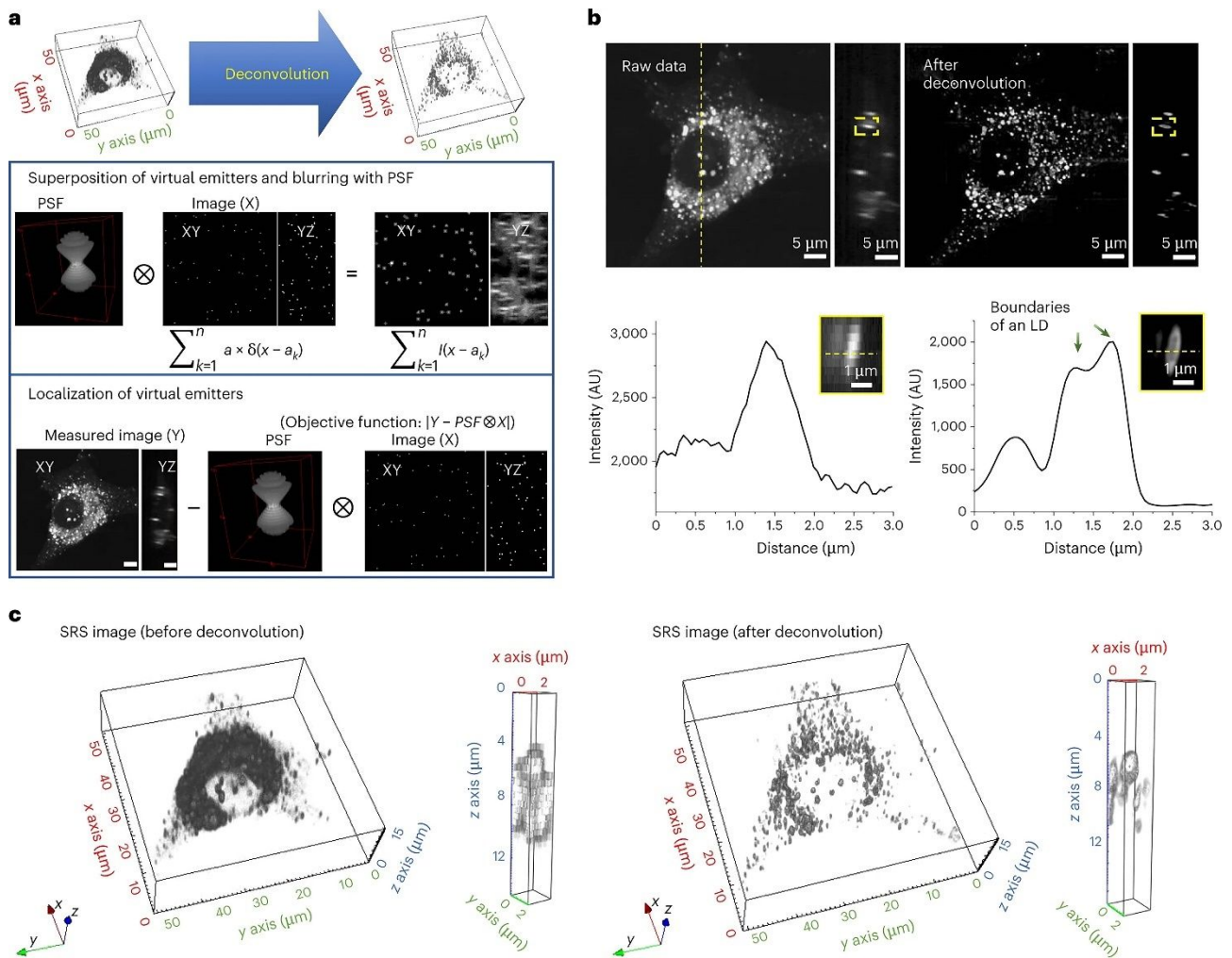
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Understanding how metabolism changes with age has far-reaching implications for human health. Marrying SRS with the power of model organisms led to the development of new approaches to define metabolic changes by incorporating heavy water (D₂O) into the diet [97, 98]. Shown here an example of the kind of insight this approach can provide. **a**, averaged Raman spectra were collected from flies of different ages labelled by 20% D₂O and then imaged. Quantification of the C–D turnover by dividing the intensity of 2143 cm⁻¹ (CD signal) to the peak at 2850 cm⁻¹ (lipid stretch) is shown in **b**. Unmixing of the C–D spectra into protein and lipid spectra is shown in **c**. The peak at 2141 cm⁻¹ corresponds to the newly synthesized D-labelled lipids, and the peak at 2180 cm⁻¹ is newly synthesized D-labelled protein. This allows quantification of lipids and protein turnover rates by comparing 2141 cm⁻¹ to 2850 cm⁻¹ and 2180 cm⁻¹ to 2929 cm⁻¹, respectively [98]. This methodology can then facilitate a direct comparison of metabolic activities in 7-day (young) and 35-day-old flies (representative SRS are shown in **e**). Old (35-day) flies showed reduced lipid metabolism compared with young flies. SRS imaging could visualize the lipids turnover inside single LDs at

the subcellular scale and subdivide this insight into small and large LDs (F-H). For further experimental details, see [98]. Figure reproduced from [98] with permission <http://creativecommons.org/licenses/by/4.0/>.

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Figure 6 Deconvolving SRS images using A-POD.



Deconvolution has played a key role in resolving light microscopy images. Jang *et al.* developed a mathematical model, A-POD (see 5.3), to allow deconvolution and applied this to mammalian cells in culture and to *Drosophila* brain. This approach provided molecularly specific information at a spatial resolution of less than 59 nm on a single lipid droplet, allowing comparison of nanoscopic distribution of lipids and proteins in cells and subcellular organelles which can be applied to *in situ* samples; this approach also represents a step-change in processing time [111]. The process is illustrated schematically, taken from Jang *et al.* [111] which shows the conversion of SRS images into super-resolution images (a). First, a specific number of virtual emitters proportional to the overall brightness of the image are placed on an image (X), and a blurred image (S) is created through convolution of X and the PSF. When the position of each virtual emitter is adjusted such that the difference between the blurred image S and the measured image (Y) is minimized, X becomes the image with the most optimal distribution of virtual emitters. They used a modified Adam solver for the optimization in A-PoD. In b, a three-dimensional deconvolution result of LDs ($2,850 \text{ cm}^{-1}$) in a live cell is shown. Following deconvolution, the membrane of an individual LD was clearly visualized in the intensity profile in the lower panel. AU, arbitrary units. Three-dimensional-rendering results of the SRS image before (left) and after (right) deconvolution are shown in c. After deconvolution, the shapes of $\sim 1\text{-}\mu\text{m}$ -sized LDs were clearly visible. Data reproduced from [111] with permission <http://creativecommons.org/licenses/by/4.0/>.

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Table 1. Glossary of key terms.

Rayleigh scattering	Light scattering from a sample at the same frequency, and hence energy, as the incident radiation.
Raman scattering	Light scattered from a sample at a different frequency, and hence energy, as the incident radiation.
Stokes and Anti-Stokes Raman scattering	Inelastic scattering can result in radiation that is either scattered with a higher wavelength (lower frequency) and is referred to as Stokes Raman scattering, or radiation scattered with a lower wavelength (higher frequency) which is referred to anti-Stokes Raman scattering.
Raman shift	Raman spectra are reported as a shift in frequency of the scattered radiation with respect to the incident radiation. The incident radiation is typically a laser which would have a Raman shift of 0 (i.e. elastic scattering).
Wavenumber	Wavenumber is a value proportional to the inverse of the wavelength in a harmonic wave. It is measured as the number of waves per unit length (centimetre), or cm^{-1} .
Spectral resolution	The ability to resolve two peaks in a spectrum which is defined by the minimum wavenumber difference that can be distinguished.
Spatial resolution	The smallest physical distance between two measured positions within the sample.
Chemometrics	The application of mathematics or statistics to chemical data.
Simulated Raman scattering (SRS)	A nonlinear Raman scattering technique that uses a pump (ω_p) and Stokes (ω_s) laser which are tuned to a defined frequency representative of a specific molecular vibration ($\omega_p - \omega_s$). When this occurs, ω_p observes a stimulated Raman loss in energy, and ω_s observes a stimulated Raman gain. The pump beam is retuned to enable a different molecular vibration to be visualised.
Coherent anti-Stokes Raman scattering (CARS)	A nonlinear Raman scattering technique that uses two lasers, a pump (ω_p) and Stokes (ω_s) beam, such that the frequency difference between the two lasers is matched to a vibrational frequency in the target sample generating a strong anti-Stokes Raman signal ($\omega_{as} = 2\omega_p - \omega_s$).
Hyperspectral imaging	A three-dimensional dataset formed of x, y, λ coordinates. In hyperspectral imaging each pixel contains a spectral profile that can be used for molecular characterisation.
Spectral phasor analysis	A Fourier transform-based technique that represents a 3D hyperspectral image stack, as a 2D density plot in which each point – or phasor – represents a single spectrum in the original 3D dataset.
Deconvolution (image) analysis	Image processing technique used to improve the contrast and resolution of images captured using an optical microscope. Blurring is a common type of image distortion that can be overcome using deconvolution analysis.
Multivariate analysis	A branch of mathematics that deals with data that change in multiple ways with respect to a single dependent variable (a reference analyte value).
Adam-based pointillism deconvolution (A-POD)	This approach mimics the concept of pointillism painting (small, distinctive dots of colour are applied in patterns) and describes images with multiple discontinuous spots (virtual emitters). The virtual emitters have the same unit intensity, and the total number of emitters is fixed across the image. This characteristic allows for suppressing of imaging artefacts and describes the real emitters' distribution. Therefore, from low-resolution images of high-density emitters, super-resolution images can be restored.

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Table 2 A comparison of spontaneous Raman scattering, CARS and SRS microscopy

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	Spontaneous Raman scattering	CARS	SRS
Quantitative	Yes	Semi-quantitative	Yes
Image acquisition rate	s/pixel	μ s/pixel	μ s/pixel
<i>In vivo</i> analysis and/or live cell imaging	Yes	Yes	Yes
Spatial resolution (nm)	~500	~300	~300
Spectral resolution (cm⁻¹)	<1	~10	~10
Advantages	Label-free, high spectral resolution, full or partial Raman spectrum at each pixel.	Label-free, high spatial resolution, fast, hyperspectral imaging is feasible	Label-free, high spatial resolution, fast, hyperspectral imaging is feasible, SRS spectrum matches Raman spectrum.
Disadvantages	Relatively slow imaging rate	Non-resonant background, indirect quantification, complex spectral retrieval	Some parasitic signals e.g. cross phase modulation can complicate imaging

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11. References.

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This is a review article and does not contain any new primary data, hence there is no relevant availability statement required.

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