Showcasing Raman detection of early stages of malaria infection from the Biophotonics Laboratory of Nicholas Smith at the Immunology Frontier Research Center, Osaka University, Japan.

Title: Raman spectroscopic analysis of malaria disease progression via blood and plasma samples

We show that Raman spectroscopy can discriminate between resonant heme and bio-crystallized hemozoin in the blood of malaria-infected patients. Both are present in blood and plasma following infection, and by multivariate analysis, their discrimination within blood samples allows early, quantitative and potentially automated detection of malaria, and its effects on the immune system.

As featured in:

See Nicholas I. Smith et al., Analyst, 2013, 138, 3927.
Illuminating disease and enlightening biomedicine: Raman spectroscopy as a diagnostic tool

David I. Ellis,*a David P. Cowcher,a Lorna Ashton,a Steve O’Hagana and Royston Goodacreab

The discovery of the Raman effect in 1928 not only aided fundamental understanding about the quantum nature of light and matter but also opened up a completely novel area of optics and spectroscopic research that is accelerating at a greater rate during the last decade than at any time since its inception. This introductory overview focuses on some of the most recent developments within this exciting field and how this has enabled and enhanced disease diagnosis and biomedical applications. We highlight a small number of stimulating high-impact studies in imaging, endoscopy, stem cell research, and other recent developments such as spatially offset Raman scattering amongst others. We hope this stimulates further interest in this already exciting field, by ‘illuminating’ some of the current research being undertaken by the latest in a very long line of dedicated experimentalists interested in the properties and potential beneficial applications of light.

Introduction

The properties of light have been of interest to experimentalists for millennia. From the publication of Ibn al-Haytham’s seven volume treatise the ‘Book of Optics’ (Kitab al-Manazir) in the early 11th century (leading to him being regarded as the father of modern optics1), through to one of the very first publications in a scientific journal, Isaac Newton’s paper on the theory of the properties of light2 itself followed some years later by his famous book, Opticks, in 1704.3 Whilst separated by seven centuries, these two polymaths and their respective bodies of work shared similarities, perhaps the most important being that both were the result of systematic and methodical experimentation. In the century following the publication of Newton’s

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David Cowcher obtained an undergraduate Masters degree in Chemistry and Forensic Science in 2009 from the University of Manchester, which included a year working as an analytical chemist for GlaxoSmithKline. He is currently studying for a PhD with Roy Goodacre at the Manchester Institute of Biotechnology, University of Manchester, on the development of enhanced Raman scattering methods for biochemical analysis.
Opticks, as early as the mid-1800s, the interaction of light within tissue was being used by physicians to assist in disease recognition.4

It was during a sea voyage from India to England in 1921 that the brilliant Indian physicist C.V. Raman, another renowned experimentalist, undertook some on-board experiments which were later submitted to *Nature* in a letter called the ‘The Colour of the Sea’.5 Unable to accept Lord Rayleigh’s explanation that the colour of the sea was just a reflection of the colour of the sky,6 Raman’s experiments showed that the colour of the sea was in fact a direct result of the molecular scattering of light and independent of absorption or the reflection of light from the sky.7 This was very closely followed by another letter to *Nature* concerning the molecular scattering of light in liquids and solids.8 These experiments opened up a deep interest in C.V. Raman and a new field of research on his return to Kolkata (Calcutta) on the scattering of light,9 as well as the publication of another article on the molecular diffraction and quantum structure of light in the following year.10

Raman and collaborators such as K.S. Krishnan began a series of seminal experiments concerning the scattering of light in a large number of liquids, as well as theories about the potential applications of their experiments, which culminated in their discovery of the inelastic scattering effect named after Raman in 1928 on 28 February,11 and his award of the Nobel Prize for Physics in 1930. This discovery was also independently observed by Landsberg and Mandelstam later in 1928 (see Fig. 1 for a timeline of events in Raman spectroscopy). There was a great deal of interest in the Raman effect and this not only aided the fundamental understanding about the quantum nature of light, and its interaction with matter at the molecular level, but also opened up a completely novel area of optics and spectroscopic research that is, particularly in terms of biological and biomedical applications,12-15 accelerating at a greater rate during the last decade than at any time since its inception.

Whereas infrared (IR) spectroscopies measure the absorption energy, Raman spectroscopy measures the exchange of energy with electromagnetic (EM) radiation of a particular wavelength, usually provided by a monochromatic light source such as a laser in the visible to near-IR portion of the EM; although it is also possible to conduct experiments in the near- and deep-UV. From the exchange in EM energy a measurable Raman shift in the wavelength of incident laser light is observed, this is also referred to as the inelastic light scattering effect.14-16 It is usually the Stokes shift which is measured, as

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Lorna Ashton studied for her BSc with the Open University before joining the University of Manchester where she obtained her PhD in Biomolecular Sciences and Raman Optical Activity. She has since worked as a post doctoral research associate specialising in Raman Spectroscopy and two-dimensional correlation analysis for the characterisation of conformational transitions in biomolecules. Lorna is currently working as part of a BBSRC funded project for rapid evolution of enzymes and synthetic microorganisms developing Raman spectroscopy as a high-throughput analytical technique for industrial bioprocesses.

Roy Goodacre is a PhD graduate from the University of Bristol (UK) where he studied mass spectrometry of microbial systems. After a postdoc, Wellcome Trust fellowship and lectureship in the University of Wales, Aberystwyth, he is now Professor of Biological Chemistry at the University of Manchester (UK). His group’s main areas of research (http://www.biospec.net/) are broadly within analytical biotechnology, metabolomics and systems biology. His expertise involves many forms of Raman spectroscopy (including deep UV resonance Raman and SERS), FT-IR spectroscopy, and mass spectrometry, as well as advanced chemometrics, machine learning and evolutionary computational methods. He is Editor-in-Chief of the journal Metabolomics, on the Editorial Advisory Boards of Analyst and Journal of Analytical and Applied Pyrolysis, a founding director of the Metabolomics Society and a director of the Metabolic Profiling Forum.
this has a higher probability of occurring than anti-Stokes.\(^\text{17}\) It is important to note that this shift is complementary to IR absorption and a spectroscopic ‘fingerprint’\(^\text{13,17,18}\) of the same sample can be analysed and constructed by both vibrational spectroscopies. Whilst mid-IR spectroscopy is known to be intensely sensitive to and a high absorber of water, this is generally not the case with Raman spectroscopy as water is a weak scatterer.\(^\text{14}\) For biomedical and routine clinical applications (and with low laser output power at the point of contact), this allows for the direct collection in vivo of Raman spectra. The utility of Raman spectroscopy has been demonstrated for a diverse and wide range of potential biological and biomedical applications, such as bacterial identification,\(^\text{19,20}\) chemical hazards and illicit substance detection,\(^\text{21,22}\) as well as food and product authentication,\(^\text{23,24}\) with a great deal of interest and research into its potential for disease diagnosis and use in biomedical applications seen during the last decade (Fig. 2).

Some of the previously documented limitations of Raman spectroscopy for biomedical applications have for example included issues such as weak scattering signals, subsequently...
long spectral acquisition times, fluorescence from biological samples, and interference from silica within fibre-optics. However, within this exciting and highly research active era of biomedical Raman spectroscopy, and with the hard work and application of experimentalists (in the long tradition of those already mentioned above), these so-called limitations are being overcome by groups of scientists and engineers around the globe who are not content to remain within the bounds of current knowledge, or the confines of commercially available optics for that matter, and who are constantly pushing the field forward. This introductory overview focuses on some of the recent developments within this exciting field, highlighting a small number of high-impact studies in imaging, endoscopy, stem cell research, and other recent developments such as spatially offset Raman scattering (SORS), coherent anti-Stokes Raman scattering (CARS) and stimulated Raman scattering (SRS), amongst others.

**Imaging**

Biomedical imaging is an extremely useful and important field which allows for the collection and processing of highly complex biochemical and physiological data, and the creation, manipulation, and in-depth analysis of three-dimensional colourised images to aid molecular sensing, drug transport, characterization of cells and tissues, and, perhaps the ultimate goal, the rapid diagnosis of disease (*vide infra*). Label-free optical imaging, particularly *in vivo*, would be highly advantageous as dyes or fluorescent labels, which are needed as contrast agents, can be toxic or perturbative to the cell/tissue. Optical techniques have also been said to have the potential to be complementary to existing techniques such as magnetic resonance imaging (MRI), and they offer superior sensitivity and high spatial resolution (compared to MRI, see Fig. 3). Whilst mass spectrometry (MS) has mass appeal for metabolomics for example and can be used for chemical imaging of tissues and cells including in 3D as well as being combined with vibrational spectroscopy, imaging MS is a destructive analysis and the practical spatial resolution for MALDI-MS is 50–200 μm and for SIMS is 1 μm, thus Raman spectroscopy has considerable opportunities to shed light on disease.

In any review of biomedical imaging, and particularly one focussed on Raman spectroscopy, it would be a heedless omission not to include any one of a number of studies by the Sunney Xie group. A major contributor to biomedical imaging, based at Harvard University (http://bernstein.harvard.edu), this group’s body of work includes both coherent Raman scattering techniques, comprising of coherent anti-Stokes Raman scattering (CARS) microscopy and stimulated Raman scattering (SRS) microscopy. Descriptions of these techniques can be found in Table 1 and it should also be noted that the signal from CARS is non-linearly proportional to species concentration whilst the signal from SRS is linearly proportional.
As briefly mentioned above, the signal from spontaneous Raman scattering is known to be weak, with potentially long integration times, which impacts by significantly reducing imaging speed (an important factor as in vivo samples are in constant motion at the micro scale). Although these nonlinear methods were first invented in the 1960s, coherent Raman scattering techniques can be said to have very recently surmounted any technical difficulties and imaging speed hurdles by enhancing the Raman signal level by up to $10^5$ to $10^6$ through the use of CARS microscopy in skin imaging of humans. Recent articles within the last half decade have explored SRS as an alternative imaging technology, which, unlike CARS, does not exhibit a nonresonant background and is a good technique for sensitive, high spatial resolution 3D imaging. Successful demonstration as a biomedical imaging method for a range of ex vivo biological samples such as brain structures, colon tissue, and arterial tissue for example, as well as in vivo studies including sciatic nerve tissue and atherosclerotic plaque deposits in animals, with a recent report also demonstrating the use of CARS microscopy in skin imaging of humans in vivo.

Recent articles within the last half decade have explored SRS as an alternative imaging technology, which, unlike CARS, does not exhibit a nonresonant background and is inherently more quantitative, though like other nonlinear multiphoton techniques (including CARS), does allow for sensitive, high spatial resolution 3D imaging. Whilst initial reports demonstrated stimulated Raman scattering’s advantages over CARS (e.g. a lack of nonresonant background complications), it was said to not be suitable for bioimaging due to sample photodamage from excessive laser power and had a limited spatial and spectral resolution as well as a slower image acquisition rate than CARS. These challenges were rapidly overcome by the Xie group the following year using a multifaceted approach which lowered peak rates by three orders of magnitude! (resulting in no photodamage), optimised spectral resolution, increased sensitivity by four orders of magnitude than the previous year’s report and surpassed the detection limit previously stated for CARS microscopy.

Table 1. Definition of some of the main approaches of Raman spectroscopy discussed in this review

<table>
<thead>
<tr>
<th>Technique</th>
<th>Acronym</th>
<th>Short definition</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface enhanced Raman spectroscopy</td>
<td>SERS</td>
<td>Requires close proximity/adsorption onto a roughened metal surface, a colloidal solution or a roughened electrode (usually Ag or Au). Enhancement explained by two processes; an electromagnetic enhancement effect (thought to dominate) and a charge transfer mechanism, known as chemical enhancement. Has a fluorescence quenching effect. Can tune to a specific chromophore for additional resonance enhancement (known as SERRS). Enhancements over normal Raman scattering of typically $10^5$ to $10^6$.</td>
<td></td>
</tr>
<tr>
<td>Coherent anti-Stokes Raman spectroscopy</td>
<td>CARS</td>
<td>A multiphoton form of Raman spectroscopy based on a non-linear conversion of two lasers into a coherent high intensity beam in the anti-Stokes region. The emission is usually many orders of magnitude greater than spontaneous Raman scattering. Useful for obtaining spectra of fluorescing samples. Nonresonant background can complicate spectral assignment, limit sensitivity, and affect quantitative interpretation</td>
<td></td>
</tr>
<tr>
<td>Stimulated Raman scattering</td>
<td>SRS</td>
<td>A multiphoton technique analogous to stimulated emission, where two lasers coincide on a sample. The sample is excited by colinear and tightly focused pump and Stokes beams. When the differences in known frequencies match a molecular vibration in the sample, the Stokes beam intensity increases and pump beam intensity decreases as a result of the coherent excitation of molecular vibration. Unlike CARS, it does not exhibit a nonresonant background and is a good technique for sensitive, high spatial resolution 3D imaging</td>
<td></td>
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<tr>
<td>Spatially offset Raman spectroscopy</td>
<td>SORS</td>
<td>Raman spectra are collected from locations spatially separated from the point of laser illumination on the sample surface. SORS allows for the isolation of chemically rich spectral information from distinct substructures or layers and through other barriers, not accessible via spontaneous Raman. Typical wavelengths used for biological tissue are 785 or 830 nm. Ideal for detecting disease in cells and tissue underlying other tissue types, such as bone through skin, cancer cells through muscle and lipid tissue for example</td>
<td></td>
</tr>
</tbody>
</table>

References

19, 122, 136 and 137
27, 38, 41, 45 and 138
25, 26, 39, 61, 63 and 139
25, 26, 39, 61, 63 and 139
Not content with these and other leaps forward, three bioimaging applications were then elegantly presented by Xie and co-workers. The first application monitored and imaged the uptake of polyunsaturated omega-3 fatty acids by living human cancer cells, specifically eicosapentaenoic acid (EPA). Using the Raman band at 3015 cm\(^{-1}\) (attributable to unsaturated fatty acids), and in contrast, the Raman band at 2920 cm\(^{-1}\) whose peak intensity is similar for saturated and unsaturated fatty acids, they were able to use SRS to follow the uptake of EPA by living cells, concluding that EPA is taken up by the cells and more strongly enriched in lipid droplets compared to other cellular organelles.\(^46\) The second application presented the potential for tissue imaging without the requirement for staining, highlighting the 3D sectioning capability and subcellular resolution of SRS, using the CH\(_2\) stretching vibration at 2845 cm\(^{-1}\) (see Table 2 for Raman band frequencies of biological interest).\(^46\)

This was demonstrated in a variety of mouse tissues; from neuron bundles in corpus callosum (highlighting myelin sheaths), thick brain tissue, and from depth profiles of ear tissue, in addition to comparative SRS and CARS images of stratum corneum (visually illustrating how the nonresonant background from CARS can complicate image interpretation). Finally, this seminal series of experiments showed the use of SRS to monitor drug delivery, namely deuterated dimethyl sulfoxide (DMSO), a skin penetration enhancer and retinoic acid (RA) used to treat a range of skin conditions and acute promyelocytic leukaemia. Drug delivery into fresh mouse skin was monitored by tuning into the vibrations for DMSO at 670 cm\(^{-1}\) and RA at 1570 cm\(^{-1}\) (as well as lipids of subcutaneous fats at 2845 cm\(^{-1}\)).\(^46\) This series of experiments elegantly demonstrated the potential for SRS as a new approach for studying pharmacokinetics \textit{in situ}, sensitive label-free imaging, and molecular sensing in 3D in living cells and tissue.

Whilst the potential of SRS had been so ably demonstrated, the series of experiments above had been undertaken \textit{ex vivo}. Well aware of the importance of \textit{in vivo} optical imaging in biomedicine, and perhaps acutely mindful of the challenges required to achieve this goal, as well as the fact that SRS imaging had not yet been accomplished \textit{in vivo} (in animals or humans), an article was published in 2010 demonstrating that \textit{in vivo} SRS was not only possible (both in mice and humans), but that it could also be performed at video-rate speeds.\(^29\) This high-speed imaging was achieved by modulating the intensity of the Stokes beam at 20 MHz and the group building their own all-analogue lock-in amplifier with a response time of \(\sim 100\) ns (whereas this was previously limited by commercially available 100 \(\mu\)s lock-in amplifiers).\(^29\) This new custom-built system raster scanned across a sample with a line rate of 8 \(\text{kHz}\) (100 ns per pixel at 512 \(\times\) 512 pixels with up to 25 frames per s), increasing imaging speed by three orders of magnitude. In addition, to significantly increase the collection of backscattered light from \textit{in vivo} samples further in-house modifications were undertaken, such as changing the geometry of the photodetector and microscope objective. This involved exciting light from the objective through an aperture in the detector placed between the microscope objective and the sample, and specially designing a filter to block the modulated Stokes beam whilst transmitting the pump beam.\(^29\)

<table>
<thead>
<tr>
<th>Band frequency (cm(^{-1}))</th>
<th>Vibration mode</th>
<th>Assignment</th>
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</thead>
<tbody>
<tr>
<td>930–950</td>
<td>N–C=C stretch</td>
<td>Skeletal stretch/(\alpha)-helix</td>
</tr>
<tr>
<td>1260</td>
<td>N–H and C–H band</td>
<td>Amide I/disordered</td>
</tr>
<tr>
<td>1300–1340</td>
<td>H-bonded C=O stretch</td>
<td>Amide I/(\beta)-helix</td>
</tr>
<tr>
<td>1650–1655</td>
<td>H-bonded C=O stretch</td>
<td>Amide I/(\beta)-sheet and (\beta)-barrel</td>
</tr>
</tbody>
</table>

**Amino acid residues**

| 508–545 | S–S stretch | Trans and gauche conformers |
| 655     | C–S stretch | Tyrosine Fermi doublet |
| 704     | C–S stretch | Tyrosine conformer |
| 835/857 | H-bonding of indole ring | Tryptophan orientation |
| 875–880 | H-bonding of indole ring | Phenylalanine |
| 1008/1034 | Phenyl ring | Tryptophan |
| 1551–1556 | Indole ring | Phenylalanine |
| 1605    | Indole ring | Tyrosine |
| 1615    |             | |

**RNA and DNA**

| 813–816 | O–P–O stretching | A-form helix |
| 914–925 | C–O and C–C stretching | Ribose–phosphate |
| 1095    | PO\(_4\) symmetric stretch | B-DNA and Z-DNA marker |
| 1135, 1235, 1395 | Ring stretch | Uracil |
| 1174, 1325, 1370 | Ring stretch | Guanine |
| 1245, 1275 | Ring stretch | Cytosine |
| 1256, 1514 | Ring stretch | Adenine |
| 1671    | C=O stretch | Thymine |

**Sugars**

| 1000–1200 | C–O and C–C stretch | \(\beta\)-d-Glucose, \(\alpha\)(+) dextrose |
| 1025, 1047, 1155 | C–O stretch from | Glycogen |
| 1267 | CH\(_2\)OH of carbohydrates | |
| 1300–1500 | NH\(_2\) rocking | GlcNAc and GalNAc |
|           | CH\(_3\) and CH\(_2\)OH deformations | \(\beta\)-d-Glucose, \(\alpha\)(+) dextrose |

**Lipids**

| 891, 908 | CH\(_2\) rocking | Fatty acid chain length |
| 1080    | PO\(_4\)-symmetric stretch | Phospholipids |
| 1259    | PO\(_4\)-asymmetric stretch | Phospholipids |
| 1296    | C–C stretch | Unbranched saturated fatty acids |
| 1660    | C=O stretch | Unsaturated lipid bonds |
| 2873, 2931, 2961 | C–H stretch | Ayl chains of lipids |
| 2888, 2926 | CH\(_3\) asymmetric stretch | Saturated lipid bonds |
| 3009    | H–C=C stretch | Unsaturated lipid bonds |

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**Table 2. A selection of Raman band frequency assignments of biological interest**

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The effectiveness of this new video-rate SRS imaging technique was then demonstrated through a series of experiments such as skin imaging in living mice. The CH₂ stretching vibration at 2950 cm⁻¹ was shown to mainly highlight proteins, as well as residual lipid structures and individual red blood cells in the viable epidermis, whilst the water signal at 3250 cm⁻¹ was coincident with sebaceous glands in both positive and negative contrast. In vivo imaging of drug penetration of trans-retinol was demonstrated in living mouse skin, visualising with SRS imaging and 3D depth projection that penetration of the topically applied drug occurs along the hair shaft and 3D depth projection that penetration of the topically applied drug occurs along the hair shaft. This pathway was not observed in previous experiments with excised fresh tissue,⁴⁸ which was said to further highlight the importance of in vivo imaging, as it can lead to insights into the transport mechanisms of small molecules in living organisms.

Finally, a series of in vivo SRS imaging experiments were performed on human skin of a volunteer’s forearm showing cell layers of the viable epidermis to a depth of 50 µm, and the boundary (via observation of varying nuclear sizes) between the viable epidermis and stratum corneum when tuned into the CH₃ stretching vibration at 2950 cm⁻¹. The penetration-enhancing small molecule DMSO (which was deuterated) was applied to the skin, and its accumulation in the hair shaft and lack of complete penetration into the hair itself was imaged using the C–D stretching vibration at 2125 cm⁻¹. With image acquisition times of 150 ms and 37 ms respectively, these blur-free high-speed in vivo vibrational imaging experiments demonstrated the diagnostic potential of this technology in humans.⁴⁹

### Stem cells

Stem cells,⁵⁰ stem cell therapy,⁵¹,⁵² and stem cell engineering⁵³ are extremely important and highly topical areas of scientific research with huge potential benefits for the treatment of a wide range of diseases and biomedical applications. The ability to identify the phenotypic purity of live cells is absolutely crucial (and a noninvasive optical method would be highly beneficial), as excessive proliferation of unwanted phenotypes (i.e., uncontrolled differentiation) following transplantation could result in undesirable consequences, such as tissue overgrowth and tumour formation for example.⁵²,⁵³ One of the recent studies by the Notingher group applied Raman spectroscopy to determine if it could be used as a label-free, noninvasive method to detect and image intrinsic chemical differences that could be used as molecular markers among highly heterogeneous stem cells populations; specifically, cardiomyocytes (CMs) derived from human embryonic stem cells (hESCs).⁵⁴

Using a custom-built instrument, Raman detection and imaging of molecular markers specific to hESC-derived CMs was carried out, along with retrospective phenotypic identification of all cells via immunofluorescence imaging integrated with the Raman microscope. Multivariate statistical analysis of Raman spectra and cross-validation methods were used to develop a model (from 50 CMs and 40 non-CMs within the same heterogeneous populations) to determine the true accuracy of phenotypic identification of CMs, the sensitivity and specificity parameters, and select discriminatory Raman bands. Raman spectral images corresponding to the Raman bands identified by both the multivariate model and immunostaining of the same cells allowed for the accurate assignment of Raman molecular markers. The conclusions drawn from the results were that spectral differences were mainly attributable to glycogen and myofibrils, with glycogen being responsible for discrimination of CMs (with a band assignment at 860 cm⁻¹), and myofibrils proteins providing a lesser contribution (with a band assignment at 938 cm⁻¹).⁵⁵ This study demonstrated the potential of Raman spectroscopy for noninvasive phenotypic identification of stem cells, though the authors themselves stated that it was not yet practical for medical applications due to their long spectral acquisition times.

However, a later study by the same group on the same hESC-derived cardiomyocytes reduced spectral acquisition times by a hundredfold, from minutes per cell to 5 s per cell, without the need for raster scanning. When incorporating high-powered commercial lasers, this could be further reduced to cell sorting speeds of approximately 10 cells per s.⁵⁶ In addition, a recent study by the same group has successfully applied Raman to identify, image and quantify the differentiation status of live neural stem cells in vitro, where this time the spectral differences were said to be related to cytoplasmic RNA.⁵⁷ A number of other interesting studies involving Raman analysis of stem cell populations have also appeared in the literature very recently⁵⁸–⁵⁹ and this area of interest, not surprisingly, continues to grow and flourish.

As well as cellular differentiation, the location of specific drugs inside cells is also of interest as this may allow the elucidation of the pharmaceutical’s site of action. A recent study addressing this enabled mapping the site of action of the HIV protease inhibitors indinavir and lopinavir in cervical carcinoma cells expressing the E6 oncogene from human papilloma virus (HPV).⁶⁰ This study demonstrated that indinavir undergoes enhanced nuclear accumulation in E6 expressing cells, indicating this as the site of action for this compound against the HPV. Further interesting studies showing the plethora of Raman imaging applications include a report on stimulated Raman photoacoustic imaging,⁶¹ surface enhanced Raman scattering (SERS) imaging using nanotags in live mice, as a potential multiplexed imaging detection method for multiple biomarkers in living subjects associated with a specific disease,⁶² quantitative multiplex SRS imaging,⁶³ noninvasive time-course imaging of apoptotic cells,⁶⁴ and multivariate image reconstruction methods for Raman hyperspectral datasets.⁶⁵ For those specifically interested in coherent nonlinear optical imaging, which includes stimulated Raman scattering, the reader is directed to a recent review by Min et al.,⁶⁶ others include a review of CARS microscopy,⁶⁷ as well as gold nanoparticles and imaging in medicine.⁶⁸

### Endoscopy

The clinical potential for in vivo Raman endoscopy has been the subject of research for over a decade, since the first published report of in vivo Raman spectra of human gastrointestinal tissue measured during routine clinical endoscopy in 2000.⁶⁹ In the same year work by the Stone group incorporated Raman spectroscopy and clinical endoscopy to discriminate between
normal, dysplastic and cancerous laryngeal tissue.\textsuperscript{69} Nick Stone’s group, based in the UK (http://www.exeter.ac.uk/) have played a major role in pioneering the field of clinical optical diagnostics with (in addition to other vibrational spectroscopies) well received studies using Raman spectroscopy/ endoscopy to analyse a range of diseases/disorders such as Barrett’s oesophagus\textsuperscript{70} and bladder and prostate cancer.\textsuperscript{71} One of the recent studies by the Stone group involved the evaluation of the suitability of a custom-built fibre-optic Raman probe for the potential \textit{in vivo} diagnosis of early onset oesophageal neoplasia. Whilst this involved \textit{ex vivo} sampling, the results clearly demonstrated the potential for the rapid and accurate differentiation between benign tissue, Barrett’s oesophagus and both dysplastic and malignant tissue.

The culmination of this study was a custom-built confocal Raman probe constructed by the group. This novel probe had been reported by the group previously,\textsuperscript{72} although by the time of the latest study, the probe had undergone some modifications which had improved its performance for spectral acquisition during oesophageal endoscopy. The 90 cm long, 2.7 mm diameter fibre-optic probe was designed to fit into the instrument channel of a standard clinical endoscope and to have direct contact with oesophageal epithelial tissue. The optics had been modified by incorporating a graded index lens at the tip (and the output power regulated to 60 mW), so that a sampling depth of 100 to 200 μm could be ensured. Collecting spectra from this depth meant that signals from deeper tissue structures would not obscure those collected from mucosal abnormalities such as early neoplastic changes,\textsuperscript{73} and that these abnormalities could be quickly classified in timescales suitable to a clinical setting. The performance of the probe \textit{ex vivo} had been evaluated for translational use for \textit{in vivo} sample collection, with low laser power at the probe tip and short spectral acquisition times said to enable its routine use for oesophageal endoscopy and paving the way for \textit{in vivo} clinical trials.\textsuperscript{73} This group have recently published a review on \textit{in vitro} and \textit{in vivo} Raman spectroscopy as a potential routine tool for the rapid, noninvasive, early diagnosis of lesions and preventing development of cancer in the oesophagus.\textsuperscript{74}

Another group who have made a significant contribution to the field of Raman endoscopy in recent years is headed by Zhiwei Huang and based at the Optical Imaging Laboratory in Singapore (http://www.bioeng.nus.edu.sg/optbioimaging/huang/). Several of these studies utilised image-guided Raman endoscopy, which the group reported for the first time in 2009.\textsuperscript{75} Whilst image-guided endoscopy is by no means novel,\textsuperscript{76,77} the integration of image-guided techniques with Raman endoscopy is relatively recent. The first report on this technique involved integrating Raman spectroscopy with trimodal imaging techniques (white-light reflectance, autofluorescence and narrow-band) and the development of a novel 1.8 mm Raman probe which filtered out interference from fluorescence as well as interference from silica from within optical fibres. This was demonstrated via the rapid collection of Raman spectra (<1 s) and the corresponding endoscopic images of different locations of the upper gastrointestinal tract of a healthy volunteer in real-time and \textit{in vivo}.\textsuperscript{75}

Since then this group have published several articles using image-guided Raman endoscopy, demonstrating its potential as an \textit{in vivo} real-time detection method for a variety of diagnostic applications. These studies have included, perhaps not surprisingly, the \textit{in vivo} diagnosis of oesophageal cancer using this technique in conjunction with biomolecular modelling.\textsuperscript{78} This involved collecting spectra from 75 oesophageal tissue sites from 27 patients of normal tissue (squamous mucosa) and malignant tumours. The cancerous tissue was said to show distinct Raman signals mainly associated with cell proliferation, lipid reduction, abnormal nuclear activity and neovascularisation. To estimate the biochemical composition of oesophageal tissue, biomolecular modelling was employed using six basis reference spectra from actin, collagen type I, DNA, histones, tr ilein and glycogen. This allowed for the construction of a linear discriminant analysis (LDA) model with a sensitivity of 97% and specificity of 95.2% for the \textit{in vivo} diagnosis of oesophageal cancer.\textsuperscript{79} These results have since been said to be extremely promising, but that image-guided Raman endoscopy is yet to be used to detect dysplasia or the early onset of cancer.\textsuperscript{73}

Nevertheless, the Huang group have previously demonstrated the potential of Raman spectroscopy for the detection of dysplasia, with a ball-lens fibre-optic probe, in the high wavenumber region (HW) (2800–3700 cm\textsuperscript{-1}) for the \textit{in vivo} detection of cervical dysplasia (a HW Raman probe was first presented by Gerwin Puppels and co-workers for \textit{in vitro} measurements of brain tissue in 2005\textsuperscript{79}). The perceived main advantages of HW Raman were said to be a significant reduction in fluorescence and background signal from optical fibres, more intense Raman signals (compared to the fingerprint region), and the possibility of an unfiltered single fibre Raman probe design for \textit{in vivo} clinical use.\textsuperscript{80} In addition to this study, several more recent reports have continued to investigate this area.\textsuperscript{81–83}

The \textit{in vivo} detection of epithelial neoplasia in the stomach using image-guided Raman endoscopy has also been demonstrated, and significant differences between normal and cancerous gastric tissue were reported.\textsuperscript{84} More recently, a study has been published showing the development of an online automated spectral diagnostics system integrated with image-guided Raman endoscopy for real-time \textit{in vivo} diagnosis during endoscopic examination.\textsuperscript{85} This system was built on a database of 2465 normal and 283 cancerous gastric tissue spectra acquired from 305 patients, with the system employing a variety of diagnostic algorithms. Other so-called \textit{in vivo} real-time applications recently reported have involved transnasal image-guided Raman endoscopy of the larynx and nasopharynx, which the authors hoped would pave the way for realizing early diagnosis and detection of cancers and precancers of the head and neck.\textsuperscript{86} All of these studies show promise and many have similarities in terms of rapid spectral acquisition times (absolutely crucial in \textit{in vivo} studies), and novel developments in optical engineering, but there remains more to accomplish, such as the ability not only to discriminate cancer, but to classify and grade both cancerous and precancerous cells and tissue.\textsuperscript{87,88}
Recent developments

In terms of noninvasive biomedical Raman other very exciting developments are spatially offset Raman spectroscopy (SORS) as well as transmission Raman, both techniques directly resulting from research involving depth profiling using Raman Kerr-gating methods. SORS was invented and developed in the Central Laser Facility of the Rutherford Appleton Laboratory (http://www.clf.rl.ac.uk) in 2005 by Pavel Matousek and readers are directed to an excellent overview of this fast moving field by Matousek and Stone.

The central difference in SORS is that Raman spectra are collected from different locations, spatially separated (offset) from the point of laser excitation on the sample (Fig. 4). As a consequence of photon diffusion processes within tissue the resultant Raman spectra contain different relative contributions from different depths within the sample and thus allows for the highly accurate chemical analysis of subsurface objects of interest. These spectra acquired from different spatial offsets are processed to reveal pure Raman spectra of subcomponents from separate depth locations within tissue. SORS can be applied in a number of Raman collection and beam delivery geometries including single point collection, ring, and other pattern illumination. In respect to biomedical imaging this approach is said to be particularly useful when configured as inverse SORS, where the sample is illuminated by an adjustable ring-shaped laser beam (generated by a conical (axicon) lens) and the Raman light collected via fibres in the centre of the ring. The radius of this ring is said to define the spatial offset and as it is adjustable, this can be optimised to suit both the scattering properties and dimensions of each sample and covers a wider illumination zone on a sample surface than conventional SORS. SORS is said to be effective at tissue depths in excess of 500 μm, which is well beyond the accessible range of conventional confocal Raman spectroscopy. Applications can be as diverse and wide-ranging as noninvasive detection of pharmaceuticals through packaging, detection of hidden explosives and drug precursors behind opaque plastics and garments, as well as agricultural and food product analysis.

From the outset, Matousek and collaborators immediately realised the diagnostic potential of SORS with its ability for noninvasive subsurface probing of for example bone through skin and dermatology studies. This foresight was quickly realised with the first SORS spectra of ex vivo bone collected in 2006 from animal and human cadavers, and in vivo SORS demonstrated by Matousek during the same year. There have been several studies since including accurate in vivo assessment of bone composition using the carbonate (1070 cm$^{-1}$)–to-phosphate (958 cm$^{-1}$) ratio through the skin of live mice, in vivo evaluation of bone grafts, transcutaneous in vivo monitoring of glucocorticoid induced osteoarthritis and in vivo measurement and evaluation of subtle changes in bone composition.

A significant body of pioneering work forwarding Raman spectroscopy as a tool for breast cancer diagnosis was undertaken in the George R. Harrison Spectroscopy Lab at MIT (http://web.mit.edu/spectroscopy/) under the directorship of Michael S. Feld. This included identifying chemical differences in microcalcifications from benign and malignant breast lesions, demonstrating the real time capabilities of an in vivo Raman system during femoral bypass and breast lumpectomy surgeries, and also involved testing of spectral diagnostic algorithms for breast cancer diagnosis. The results from the Raman research on the chemical composition and identification of the different types of microcalcifications has led directly to others investigating the potential of SORS as a possible clinical adjunct to mammography, for the noninvasive diagnosis of breast cancer. As the changing concentration of carbonate substitution for phosphate ions in the calcium hydroxide lattice in microcalcifications may relate to the process of tumour cell metastasis, and the ability to measure the magnitude of this (as well as soft tissue signals) by Raman, could indicate the potential progression of this cancer.

Research by Stone, Matousek, and collaborators, demonstrated the proof-of-principle of SORS for potential in vivo breast cancer diagnosis in a model system using three calcification standards overlaid with various preparations of animal tissue (i.e. chicken breast tissue, with and without skin). For this work they utilised a continuous wave 827 nm laser with a spatial offset for collection of 3 mm, which enabled the probing and collection of Raman spectra from calcifications through up to 10 mm of tissue. Previous results by this group using Kerr-gated Raman techniques had achieved penetration depths of 1 mm in comparison. This SORS study demonstrated the collection of high quality Raman spectra and biochemical information measured through 8.7 mm of tissue, identifying the difference between three calcification standards.

Subsequent experiments by the same group applied transmission Raman spectroscopy in combination with chemometrics in similar model systems collecting Raman calcification signals from depths of 20 mm, this time through porcine tissue, as a breast tissue ‘phantom’. This was said to reach the lower range for clinically relevant breast tissue thicknesses from

![Fig. 4](simplified_graphical_representation_of_A_spontaneous_Raman Compared_to_B spatially_offset_Raman_scattering_illustrating_the Spatial_offset_and_deeper_ subsurface_probing_in_skin_tissue. L_light,L_Raman_light.)
mammographic screening (1.9 cm), with further work said to be required to reach the top of this range (5 cm).\(^{118}\) Whilst these studies utilised the carbonate-to-phosphate band ratio (similar to the SORS bone studies above\(^{116}\)), a more recent study showed that the intense Raman phosphate band at 960 cm\(^{-1}\) broadens and shifts as carbonate concentration (from calcium hydroxyapatites) increases in the calcifications.\(^{119}\) This study was said to pave the way toward a new generation of noninvasive breast cancer screening methods based around SORS and transmission Raman spectroscopy.\(^{119}\) Some very recent studies by the Mahadevan-Jansen group have investigated SORS for the real-time, intraoperative assessment of breast cancer tumour margins.\(^{112}\) This same group then developed and tested a SORS probe with multiple source detector offset limits specific to this type of analysis, and published results acquired from 35 freeze-thaw breast cancer samples in vitro.\(^{113}\)

Variants of SORS have also recently been used to augment analysis and include in vivo transcutaneous glucose sensing in rats\(^{114}\) and further demonstrated the accuracy and functionality of this in a later study.\(^{115}\) These studies employed what they term surface enhanced SORS (SESORS), first published in 2010,\(^{116}\) and presented at the Federation of Analytical Chemistry and Spectroscopy Societies (FACSS) meeting in the USA during the same year. SESORS marry SERS techniques using nanoparticles and nanosurfaces with the subsurface probing of SORS, with a further study exploring this technique’s potential for Raman imaging.\(^{117}\)

On the subject of SERS, which has not been mentioned in any great detail here, many studies have appeared in the literature during the last few years related to disease detection and we would like to highlight just a modest selection of these. They of course include the multiplexed imaging of SERS nanotags in vivo already mentioned above,\(^{62}\) as well as other multiplexing studies, such as multiplex single nucleotide polymorphism (SNPs) genotyping coupled to SERS,\(^{118}\) multiplexed in vivo cancer detection using SERS NIR nanotags, demonstrating the excellent sensitivity, stability and tumour specificity of three bioconjugated nanotags,\(^{119}\) recent reviews on the area of SERS multiplexed detection for disease diagnostics,\(^{120}\) as well as SERS cancer detection and imaging and the potential of SERS agents for targeted drug delivery and photothermal therapy.\(^{121}\)

Several SERS studies involving immunoassays, including cancer detection,\(^{122}\) detection of a potential pancreatic cancer marker in serum,\(^{123}\) and on-chip immunoassays using hollow gold nanospheres.\(^{124}\) Fluorescent SERS gold co-functionalized nanorod probes for in vivo imaging of lymph node mapping and tumor targeting in mice,\(^{125}\) high sensitivity in vivo detection of inflammation using gold nanoclusters conjugated to monoclonal antibodies\(^{126}\) and the use of functionalised nanoparticles and SERS for the detection of DNA relating to disease.\(^{127}\) With the recent interest concerning the potential resurgence in microbial disease, as well as bioterrorism, an article demonstrating a portable quantitative SERS system for detection of Bacillus spores at levels significantly lower than those previously reported for SERS has been published.\(^{128}\) Fig. 5 shows SERS spectra of a biomarker from the spores of Bacillus, a species of which Bacillus anthracis is the cause of the acute, and mostly lethal, disease anthrax. The detection limit of this dipicolinic acid biomarker is estimated at 5 ppb (30 nM).\(^{128}\)

In addition to those primary articles and reviews already mentioned above, other recently published reviews and books on an array of Raman-based topics which may be of interest include those concerning in vivo and in vitro analysis,\(^{129}\) Raman scattering in pathology,\(^{130}\) optical tumour margin identification in the larynx,\(^{131}\) data-classification algorithms for spectral analysis,\(^{132}\) recent advances in gold nanoparticle based assays for detecting and identifying microbes,\(^{133}\) and emerging Raman applications and techniques in biomedical and pharmaceutical fields.\(^{134}\)

**Concluding remarks**

We hope that whilst only an introductory overview to some of the more recent work in this field, the range and scope of the studies shown here elegantly demonstrate the development and exploitation of Raman spectroscopy as a medical diagnostic tool. We believe this truly is an exciting field, where those at the forefront are propelling it forwards and are simply not content to remain within the confines of current knowledge or commercially available optics. These studies highlight the necessary interdisciplinary nature of this field, with inspirational contributions from a range of analytical scientists and technologists including clinicians, biologists, chemists, optical engineers, as well as statisticians and chemometrists. These multidisciplinary inputs and fresh approaches are invigorating this field, unlocking the doors to new insights,\(^{135}\) adding to knowledge, and opening up new dimensions and avenues of study, as well as potential clinical applications, for biomedical Raman spectroscopy.
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