Raman spectroscopy and coherent anti-Stokes Raman scattering imaging: prospective tools for monitoring skeletal cells and skeletal regeneration

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The use of skeletal stem cells (SSCs) for cell-based therapies is currently one of the most promising areas for skeletal disease treatment and skeletal tissue repair. The ability for controlled modification of SSCs could provide significant therapeutic potential in regenerative medicine, with the prospect to permanently repopulate a host with stem cells and their progeny. Currently, SSC differentiation into the stromal lineages of bone, fat and cartilage is assessed using different approaches that typically require cell fixation or lysis, which are invasive or even destructive. Raman spectroscopy and coherent anti-Stokes Raman scattering (CARS) microscopy present an exciting alternative for studying biological systems in their natural state, without any perturbation. Here we review the applications of Raman spectroscopy and CARS imaging in stem-cell research, and discuss the potential of these two techniques for evaluating SSCs, skeletal tissues and skeletal regeneration as an exemplar.

1. Introduction

Throughout the last century, medical breakthroughs have led to a tremendous increase in life expectancy. However, as a consequence, an increasing ageing population has resulted in an increase in age-related diseases, as well as associated reductions in quality of life, leading to a dramatic impact on healthcare. Indeed, skeletal tissue loss due to injury or disease results in a significantly reduced quality of life at significant socio-economic cost. Fractures alone cost the European economy €17 billion and the US economy $20 billion annually [1]. In the USA, there are around 8 million bone fractures per year, of which approximately 5–10% are associated with delayed healing or non-union [2]. Each year in the UK there are approximately 150,000 wrist, vertebral and hip fractures due to osteoporosis, with an estimated healthcare cost of £2.1 billion per annum. Thus, novel and effective medical approaches are essential to fulfil the current demographic challenges [3–5].

The use of stem cells for cell-based therapies is one of the most promising and exciting areas for tissue repair and disease treatment, including those affecting brain, skeletal muscle and heart [6,7]. In fact, the unique properties of stem cells, with their ability to self-renew and potential to differentiate into several different specialized cell types, present an ideal tool for reparative medicine [3,8].

The bone marrow, which is the major site of haematopoiesis (the process which leads to the formation of all blood cells), serves as a reservoir for a variety of cells, including haematopoietic cells as well as cells of the non-haematopoietic stroma [4,9–11]. The bone-marrow stroma constitutes the scaffold that supports the regulation of haematopoiesis, establishing and maintaining the haematopoietic microenvironment necessary for growth and blood-cell
maturation [9,12]. Within the bone-marrow stroma reside a rare multipotent stem-cell population called skeletal stem cells (SSCs). The term ‘skeletal stem cell’ is all too frequently confused with ‘mesenchymal stem cell’ (MSC) in the literature (reviewed in [12]). However, MSCs are developmentally distinct from skeletal lineages, and it is important to recognize that the various extra-skeletal tissues and organs noted to retain MSCs are developmentally distinct from skeletal lineages and, critically, are not generated by skeletal progenitors present in bone marrow. The term ‘skeletal stem cell’, as eloquently detailed by Bianco & Robey [13], denotes specifically the rare population of postnatal non-haematopoietic stromal cells found in the bone marrow with the capacity to regenerate bone and bone-marrow stroma (reviewed in [3]), and the term SSC will be used throughout this review to refer to this select population.

The ability to isolate SSCs from human bone marrow, together with their capability to differentiate into skeletal tissues, has attracted significant attention within the medical community, and promises new opportunities for the use of SSCs in clinical applications [3,10,14]. For example, SSCs could be applied for bone/cartilage tissue repair for a range of musculoskeletal conditions, or as key components of the haematopoietic microenvironment for microvessel assembly guidance, or even as conceptual/modelling instruments for stem-cell biology and mechanistic studies [12]. However, pivotal in the development of successful therapies for bone augmentation will be the design of well defined and reproducible protocols for SSC differentiation. SSCs can be induced to form bone, cartilage and fat depending on their microenvironment (figure 1) [4,15,16], and the differentiation into osteoblasts, chondrocytes and adipocytes, both in vitro and in vivo, is plastic (reversible) [11]. Controlled differentiation of stem cells offers significant therapeutic potential in skeletal regeneration, with the prospective to permanently repopulate a host with stem cells and their progeny, an objective that could be achieved by careful monitoring and characterization of the differentiation process of SSCs.

Currently, SSC differentiation can be evaluated through a number of assays including colorimetric assays [17], real-time quantitative polymerase chain reaction (qPCR) [17,18], histochekmical analysis [18] and immunohistochemical assays [19]. These methods reveal the detailed biochemical features of SSCs and provide important information regarding their state of differentiation. However, current approaches are invasive, require cell fixation or lysis, and/or are destructive [20–22], indicating current monitoring techniques are unsuitable for time-course studies, as well as unsuitable for SSC characterization prior to therapeutic use. For example, photobleaching of fluorophores/fluorescent dyes limits the temporal availability for imaging the sample of interest. A further important issue is the non-specific binding of dyes or labels, and the staining process may modify the functionality of the target molecule and affect cell biochemical phenomena [21,23].

The current limitations in the characterization of SSC differentiation have led to the need to find new alternatives. Prospective approaches seek to identify molecules at subcellular level without using any dye or label, i.e. ‘label-free’, but by using the intrinsic properties of the molecules. Label-free techniques which offer chemical or molecular selectivity are particularly attractive. Methods which rely on the measurement of vibrational information such as Raman spectroscopy and coherent anti-Stokes Raman scattering (CARS) microscopy are inherently non-invasive (as no introduction of labels is required), non-destructive, as well as chemically selective. It is noted that as with all optical techniques power and exposure to light need to be within a threshold to prevent any damage and phototoxic effects on cells. In this review, we present a detailed overview of the recent applications of Raman spectroscopy and CARS microscopy in stem-cell research, and discuss the potential of label-free techniques to study SSCs for skeletal regeneration.

### 2. Characterization of skeletal stem cells using label-free approaches

Raman spectroscopy provides information from the intrinsic vibrations of molecules in their native state, without the need for an exogenous label and has been explored as a label-free method for biological and biomedical applications [24–26]. Fourier transform infrared (FTIR) spectroscopy, a technique which provides complementary information to Raman spectroscopy, has also been applied to biological samples in order to study their vibrational profile. However, FTIR is not ideal for biomedical imaging because water strongly absorbs infrared light [21,23,27]. Raman spectroscopy is typically carried out with visible or near-infrared laser sources, and is ideal for measurement of biological samples in the natural state, as water is a very poor Raman scatterer. Both FTIR and Raman spectra can reveal a molecular fingerprint of SSCs, from single molecules to complex structures, providing broad chemical information on their content (lipids, proteins, nucleic acids, etc.). A comparison of the key features from different characterization methods for SSCs is presented in table 1 [23,25,28–31], and it can be seen that Raman spectroscopy and CARS microscopy offer some advantages over traditional SSC characterization techniques.

### 3. Raman spectroscopy

When a beam of light interacts with a molecule, the energy is mostly scattered elastically (no change in energy) with the same frequency as the incident light, known as Rayleigh scattering (figure 2a). A few incident photons interact and exchange energy with the molecular bond vibration, resulting in inelastically scattered light, also known as Raman scattering [31].
A molecule that absorbs a photon enters into a virtual excited state and, almost immediately, another photon is emitted at a slightly different wavelength. When the incident photon loses energy to a molecular bond vibration a red-shifted Stokes photon, i.e. a photon with longer wavelength, is generated and the molecule ends up in a higher vibrational state (figure 2b). A blue-shifted anti-Stokes scattering can also occur, if the molecule of interest is already in a higher vibrational state, resulting in the emission of a lower wavelength photon (figure 2c). The difference in frequency between incident and scattered photons corresponds to the vibrational energy level of the molecule [23,31–33], and is typically called Raman shift (as the shift in wavelength from the incident radiation is measured).

As each molecule is unique with its own set of characteristic bonds and therefore vibrational modes, Raman spectroscopy provides a molecular ‘fingerprint’. Given that cells and tissues are composed of different molecules, Raman spectroscopy offers, potentially, a powerful tool to generate a characteristic signature of SSCs and skeletal tissues.

Furthermore, Raman spectroscopy offers other advantages to study living cells including: (i) high spatial resolution, (ii) qualitative and quantitative spectral information, (iii) ability to detect at the subcellular level, and (iv) the ability to analyse cells in real time without altering cell function, as a laser operating at the visible/near-infrared region is applied to prevent significant damage to proteins, DNA, RNA and other present biomolecules [30,34–37].

The last decade has witnessed the use of Raman spectroscopy in the search for prospective spectral markers for characterizing stem cells, including murine embryonic stem cells [20,37–39], human embryonic stem cells [21,40–43], MSCs [21,26,35], SSCs [36,44,45], adipose-derived stem cells (ADSCs) [21,46], as well as the monitoring of stem-cell differentiation into skeletal tissues [21,35,36,45,46]. Table 2 summarizes the current applications of Raman spectroscopy for stem-cell research and the major outcomes. We discuss in detail those studies relevant for skeletal regeneration.

In 2009, Chiang et al. [35] studied osteogenic differentiation of MSCs applying Raman spectroscopy, with the purpose to monitor the production of hydroxyapatite throughout the osteogenic process. Chiang and colleagues found changes in the hydroxyapatite characteristic ‘chemical’ shift, over the period of 7–21 days following the commencement of differentiation. The state of differentiation of MSCs was confirmed by the use of alizarin red S staining for calcium. Chiang et al. also detailed a novel marker in MSC-derived osteoblasts by monitoring hydroxyapatite with Raman spectroscopy, providing the first indication that this technique could be a promising tool for the study of skeletal tissue development.

Downes et al. [21] also induced MSC osteogenic differentiation for 7 days, and observed characteristic peaks in the osteoblast spectra related to phosphate in hydroxyapatite, collagen and carbonate.

Table 1. Comparison of the main features of skeletal stem-cell characterization techniques.

<table>
<thead>
<tr>
<th></th>
<th>detection sensitivity</th>
<th>quantitative analysis</th>
<th>molecular specificity</th>
<th>sample preparation</th>
<th>non-invasive analysis</th>
<th>time consumption</th>
<th>in vivo relevance</th>
</tr>
</thead>
<tbody>
<tr>
<td>biochemical assays</td>
<td>low</td>
<td>semi</td>
<td>medium</td>
<td>yes</td>
<td>no</td>
<td>medium</td>
<td>medium</td>
</tr>
<tr>
<td>histological assays</td>
<td>low</td>
<td>no</td>
<td>medium</td>
<td>yes</td>
<td>no</td>
<td>high</td>
<td>high</td>
</tr>
<tr>
<td>qPCR</td>
<td>high</td>
<td>yes</td>
<td>high</td>
<td>yes</td>
<td>yes</td>
<td>low</td>
<td>low</td>
</tr>
<tr>
<td>IR spectroscopy</td>
<td>medium</td>
<td>yes</td>
<td>high</td>
<td>yes</td>
<td>yes</td>
<td>low</td>
<td>low</td>
</tr>
<tr>
<td>Raman spectroscopy</td>
<td>low</td>
<td>yes</td>
<td>high</td>
<td>no</td>
<td>yes</td>
<td>low</td>
<td>high</td>
</tr>
<tr>
<td>CARS</td>
<td>medium</td>
<td>semi</td>
<td>medium</td>
<td>no</td>
<td>yes</td>
<td>low</td>
<td>high</td>
</tr>
</tbody>
</table>

Figure 2. Schematic diagram of the energy transitions involved in Rayleigh scattering (a) and Raman scattering (b,c). Raman scattering occurs through the interaction of an incident photon with a molecular vibration mode, gaining (anti-Stokes scattering, blue-shifted) or losing (Stokes scattering, red-shifted) an amount of energy equal to that vibrational mode. (Online version in colour.)
Table 2. Raman spectroscopy applications in stem cells.

<table>
<thead>
<tr>
<th>reference (year)</th>
<th>cell type</th>
<th>study</th>
<th>main findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Notingher et al. (2004) [37,39,47]</td>
<td>murine embryonic stem cells</td>
<td>characterization of murine embryonic stem cells</td>
<td>changes in the Raman spectra in the RNA peak region can be used as a differentiation marker</td>
</tr>
<tr>
<td>Ichimura et al. (2014) [38]</td>
<td>murine embryonic stem cells</td>
<td>spontaneous differentiation of embryonic stem cells</td>
<td>differences between Raman spectra of embryonic stem cells before and after spontaneous differentiation</td>
</tr>
<tr>
<td>Downes et al. (2011) [21]</td>
<td>human embryonic stem cells</td>
<td>characterization of human embryonic stem cells</td>
<td>differences in the Raman spectra between nucleus (higher levels of RNA) and cytoplasm (higher levels of protein and glycogen)</td>
</tr>
<tr>
<td>Chan et al. (2009) [40]</td>
<td>human embryonic stem cells</td>
<td>embryonic stem-cell differentiation into cardiomyocytes</td>
<td>changes in the RNA and DNA Raman peaks, before and after differentiation</td>
</tr>
<tr>
<td>Schulze et al. (2010) [42]</td>
<td>human embryonic stem cells</td>
<td>differentiation status of human embryonic stem cells</td>
<td>identification of Raman bands and ratios (e.g. RNA/proteins) to indicate embryonic stem-cell state of differentiation</td>
</tr>
<tr>
<td>Pasut et al. (2013) [41]</td>
<td>human embryonic stem cells</td>
<td>embryonic stem-cell differentiation into cardiomyocytes</td>
<td>changes in the Raman spectra of carbohydrate and lipid chemical shifts, increasing during differentiation process</td>
</tr>
<tr>
<td>Tan et al. (2012) [43]</td>
<td>human embryonic stem cells and human-induced pluripotent stem cells</td>
<td>differences between embryonic stem cells and induced pluripotent stem cells</td>
<td>very similar Raman spectra, with small changes in the glycojen bands</td>
</tr>
<tr>
<td>Pijanka et al. (2010) [26]</td>
<td>human embryonic stem cells and human mesenchymal stem cells</td>
<td>differences between human embryonic stem cells and MSCs</td>
<td>Raman scattering allowed one to distinguish an increase in the DNA band when comparing the embryonic stem cells with the MSCs nuclei</td>
</tr>
<tr>
<td>Chiang et al. (2009) [35]</td>
<td>human mesenchymal stem cells</td>
<td>MSC differentiation into osteoblasts</td>
<td>changes in the Raman spectra in the hydroxyapatite characteristic peak region during the osteogenic differentiation</td>
</tr>
<tr>
<td>Downes et al. (2011) [21]</td>
<td>human mesenchymal stem cells</td>
<td>MSC differentiation into osteoblasts</td>
<td>changes in the Raman spectra in the hydroxyapatite, collagen and carbonate chemical shifts during the osteogenic differentiation</td>
</tr>
<tr>
<td>McManus et al. (2011) [45]</td>
<td>human skeletal stem cells</td>
<td>SSC differentiation into osteoblasts</td>
<td>changes in the spectra in the hydroxyapatite Raman shift during osteogenic differentiation; measurement of carbonate-to-phosphate and mineral-to-matrix ratios at different stages of development</td>
</tr>
<tr>
<td>Hung et al. (2013) [36]</td>
<td>human skeletal stem cells</td>
<td>SSC differentiation into osteoblasts</td>
<td>changes in the spectra in the octacalcium phosphate, β-tricalcium phosphate and hydroxyapatite Raman shifts, able to detect the extent of maturation during osteogenic differentiation</td>
</tr>
<tr>
<td>James et al. (2015) [44]</td>
<td>human skeletal stem cells</td>
<td>analysis of functional markers in SSCs using immortalized SSC clonal lines</td>
<td>different SSC clones were identified by Raman spectroscopy, presenting the same biomolecular profile as human SSC fractions</td>
</tr>
</tbody>
</table>

(Continued.)
Similar approaches were used, where SSCs derived from human bone marrow, and subsequent differentiation into osteoblasts, were characterized and monitored [36,45]. For example, McManus et al. [45] used Raman spectroscopy as a biochemical characterization tool for SSC differentiation into osteoblasts, and compared the results with immunocytochemistry and qPCR analysis (figure 3). McManus et al. determined carbonate-to-phosphate and mineral-to-matrix ratios using specific peaks in Raman spectra at different stages of osteogenic development.

Table 2. (Continued.)

<table>
<thead>
<tr>
<th>Reference (Year)</th>
<th>Cell Type</th>
<th>Study</th>
<th>Main Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Downes et al. (2011) [21]</td>
<td>Human adipose-derived stem cells</td>
<td>ADSC differentiation into osteoblasts and adipocytes</td>
<td>Changes in the Raman spectra in the hydroxyapatite, collagen and carbonate chemical shifts after osteogenic differentiation; Raman peaks from lipids/proteins are sharper after adipogenic differentiation</td>
</tr>
<tr>
<td>Ojansivu et al. (2015) [46]</td>
<td>Human adipose-derived stem cells</td>
<td>ADSC differentiation into osteoblasts, using different bioactive glasses</td>
<td>Similarities in the hydroxyapatite, octacalcium and β-tricalcium phosphate Raman chemical shifts between different cell-culture conditions</td>
</tr>
</tbody>
</table>

Figure 3. Raman spectra of pre-mineralized SSCs cultured in osteogenic media at day 7 (a) (b) and day 14 (c) (f). Immunocytochemical localization of type II collagen in SSCs cultured for 14 days in basal media (d) and osteogenic media (e). Relative expression of type II collagen gene in SSCs cultured for 14 days in basal and osteogenic media, determined by qPCR (g). Adapted from [45]. Copyright Royal Society of Chemistry. (Online version in colour.)
and observed an increase of the two ratios with time. Hung’s research group identified new spectral markers for osteogenic differentiation in SSCs [36]. In this case, the characteristic chemical shift of octacalcium phosphate was present before differentiation, and the peak decreased throughout the assay period. By contrast, the hydroxyapatite signal increased during SSC differentiation into osteoblasts, and, in addition, a new peak belonging to the β-tricalcium phosphate appeared following differentiation. Hung et al. further corroborated their results using histochemical and gene expression analyses.

Other groups have reported on ADSCs for skeletal regeneration and their characterization by Raman spectroscopy [21,46]. Downes et al. [49] differentiated ADSCs into osteoblasts and adipocytes, and characterized the different populations using Raman spectroscopy. Similar to Hung’s work [36], Ojansivu et al. [46] recently used octacalcium phosphate, hydroxyapatite and β-tricalcium phosphate as specific markers for osteogenic differentiation, in order to compare culture conditions of ADSCs with different bioactive glasses. More recently, Mitchell et al. [48] demonstrated that Raman spectroscopy can be used to detect biochemical changes associated with adipogenic differentiation of ADSCs in a non-invasive and aseptic manner. Mitchell and colleagues were able to monitor the adipogenic differentiation of live ADSCs during 14 days and found significant differences from day 7.

An interesting possible application of Raman spectroscopy in SSC research is the identification of different single-cell-derived clones, which could guide the search for new strategies to analyse the differentiation potential of SSCs, or even SSC isolation from human bone marrow. James et al. [44] studied distinct subtypes of human bone marrow stromal cells, and Raman spectroscopy was applied to identify the molecular fingerprint of the stromal cells subtypes together with the biomolecular profile of human bone-marrow CD317+ fractions. Peak intensity ratios were obtained, and the main difference in the Raman shift was found at 1088.6 cm−1, which is related to the symmetric phosphate stretch of the DNA backbone, indicating a fundamental difference in the DNA of the stromal cell subtypes.

Raman spectroscopic analysis is frequently applied in conjunction with multivariate statistical analysis, for example, principal component analysis and hierarchical clustering analysis [26,37,45], due to the huge amount of chemical information embedded in Raman spectra. These methods provide an approach and are often necessary to extract information on different constituents which exist in varying proportions in a heterogeneous sample such as bone. In addition, a large amount of spectral data can be generated from a tissue map or from different populations (healthy versus diseased, etc.). Identification of compositional changes, classification and quantification of concentrations require the use of such statistical methods as has been shown in several studies with stem cells [40,42,45], and other diseases such as cancer [50]. Indeed, multivariate statistical methods for spectra analysis have contributed to the increase in Raman spectroscopy applications in the examination of living cells, and could be a helpful and vital tool for the comparison of biological samples.

4. Coherent anti-Stokes Raman scattering imaging

While extremely powerful in terms of characterization ability, the efficiency of Raman scattering is extremely small (approximately 1 in 107 scattered photons [51]). Thus for imaging Raman scattering is rather limited as acquisition times per pixel can be up to several seconds. The use of alternative Raman-based techniques that can significantly boost and enhance Raman signal levels, such as CARS or surface-enhanced Raman scattering (SERS), offers exciting alternatives. SERS is highly sensitive but involves using an exogenous material to facilitate the read-out, usually functionalized metal nanoparticles, which may not be desirable. CARS provides the same label-free chemical contrast as Raman spectroscopy, although, and importantly from the current perspective, with improved sensitivity [52].

CARS occurs when a target molecule is irradiated using two laser beams simultaneously at different frequencies, a pump beam ωp and a Stokes beam ωs (figure 4) [53]. In essence, when the difference between the higher frequency (pump beam) and the lower frequency (Stokes beam) equals the vibrational frequency of the target bond of the molecule, a CARS signal is generated [32,33,52,54], with an angular frequency equal to ωCARS = 2ωp − ωs. By tuning the frequencies of the two beams to match a particular vibration, a coherent signal with much higher intensity than the signal from spontaneous Raman scattering (up to five orders of magnitude [21,55]) is emitted, providing vibrational contrast in the subsequent CARS image [32,52]. Acquisition times per pixel in CARS imaging are typically between 1 and 10 μs.

CARS is a multiphoton (four-wave mixing) process with no net deposition of energy within the molecule (as illustrated in figure 4), and is most often carried out with near-infrared laser sources. Besides being non-invasive, non-destructive and label-free, CARS provides relevant benefits over other imaging techniques: (i) the sample photo-damage is minimized, as no net energy is transferred to the sample [54], (ii) CARS provides inherent three-dimensional sectioning ability and video-rate imaging [23,54,56,57], due to the non-linear multiphoton nature of CARS process, which is of paramount importance for studying cells or tissues, and (iii) fluorescence does not interfere in CARS, as anti-Stokes Raman scattering occurs at a different (blue-shifted) wavelength from fluorescence [32,52].

CARS microscopy has been applied in the imaging of living cells and ex vivo tissues, using diverse vibrational contrasts such as for DNA, lipids and proteins [10,52,54,58]. Figure 5, as an example, shows the CARS imaging of a human squamous-cell carcinoma metastasis in the brain. Meyer et al. analysed the same tissue sample with CARS microscopy (targeting lipids) without any staining, and with brightfield microscopy after staining the sample with the gold standard haematoxylin and eosin (H&E).

The abundance and the essential role of lipids in the human body, combined with the fact that lipids have strong CH2 and CH3 stretching vibration signals at 2853 cm−1 and 2935 cm−1 [24,52,57], respectively, have made lipids the target of choice for demonstrating CARS applications. CARS applications in cell studies are being expanded—for instance, in order to develop an adipose-tumour epithelial cell co-culture system designed to reproduce the in vivo mammary environment, Salameh et al. [60] required a non-destructive and non-invasive technique to access the viability of adipocytes in co-culture. In this study, CARS analysis demonstrated the sustained viability of adipocytes, and an ex vivo co-culture model system to evaluate stromal—epithelial interactions in breast cancer was established. Another example is the application of CARS to study the effect of chemotherapeutic drugs in colon tumour
cells. As demonstrated by Steuwe et al. [24], CARS was a useful tool to rapidly image the subcellular accumulation of lipids in cancer cells undergoing cell death induced by different chemotherapeutic drugs.

The last few years have seen emergent data of in vitro cell studies for skeletal regeneration that have applied CARS imaging. The principal findings regarding CARS applications for stem-cell characterization are summarized in table 3. One of the first examples, shown in 2007 by Konorov et al. [56], used CARS microscopy to analyse murine embryonic stem cells. Although image quality did not allow the identification of individual cells, this approach provided a first step towards the use of CARS imaging in stem-cell research. The following year, Schie et al. induced and characterized MSC differentiation into adipocytes, which is commonly assessed by identifying the development of lipid droplets. As the CARS signal for lipids is particularly strong, Schie and colleagues successfully imaged the lipid droplets after 21 days of adipogenic differentiation [61]. Recently, Smus et al. [66] used CARS imaging to assess adipogenic differentiation of SSCs, in conjunction with molecular analysis of gene expression and conventional oil red O staining methodology. The authors demonstrated that CARS microscopy was a valuable technique to detect early stages of SSC differentiation (24 h and 72 h after adipogenesis induction), with enhanced resolution and definition of lipid droplets. Furthermore, CARS microscopy provided an alternative method to monitor changes in SSCs as a result of chemical modulation of adipogenic differentiation.

Figure 4. Schematic diagram of the energy transitions involved in coherent anti-Stokes Raman scattering (CARS). In CARS, molecular vibrational modes are coherently populated through optical pumping, by the combination of the pump and the Stokes. The vibration coherence is probed by the pump to generate the CARS signal. Thus when the energy difference between the pump beam (higher frequency) and the Stokes beam (lower frequency) equals a particular molecular vibration of the target, the CARS signal is maximized. (Online version in colour.)

Figure 5. Comparison of H&E staining (a) and CARS imaging (b) of a human squamous-cell carcinoma metastasis in the brain. The bright components indicate the intensity of the CARS signal from lipids. Adapted from [59]. Copyright Wiley-VCH Verlag GmbH & Co. KGaA. (Online version in colour.)
Different research groups have worked with ADSCs and monitored cell differentiation with CARS microscopy. Jo et al. [62] selectively imaged lipid droplets with high contrast in differentiated adipocytes, and it was possible to observe a chronological differentiation of ADSCs, comparing oil red O staining with CARS images. In turn, Downes et al. [21] not only used CARS microscopy to characterize adipocytes, but also tried to establish the use of CARS to image osteoblasts. By imaging at the Raman frequency of hydroxyapatite (960 cm⁻¹), Downes obtained preliminary results with some vibrational contrast although with limited resolution.

Undoubtedly, CARS imaging has emerged as an important alternative technique for monitoring the development of lipid droplets in SSCs. The combination of CARS microscopy with complementary imaging techniques, namely two-photon excitation fluorescence (TPEF) and second harmonic generation (SHG), can be highly advantageous and give an important insight on the development of SSCs from different angles. TPEF and SHG are also nonlinear imaging modalities which usually can be carried out with the same laser excitation sources thus allowing multimodal read-out. While TPEF allows mapping fluorophores (or auto-fluorophores) with near-infrared excitation, SHG is particularly sensitive to non-centrosymmetric structures such as collagen. CARS together with SHG and/or TPEF could give a more holistic insight into the development of SSCs. Figure 6 shows an exciting application of CARS and SHG microscopy as used by our research group to assess the state of differentiation of fetal femur-derived SSCs. Simultaneous imaging of lipids (CARS) and collagen fibres (SHG) in human fetal femur-derived SSCs cultured in adipogenic (figure 6a) and chondrogenic (figure 6b) media is shown.

CARS combined with SHG and TPEF imaging has been applied by Mouras et al. to assess ADSC differentiation into adipocytes and osteoblasts [63]. In the same year, Mortati et al. [64] followed collagen production in living SSCs within a fibrin hydrogel scaffold in a three-dimensional culture system using SHG and CARS, clearly demonstrating the potential of multimodal CARS imaging. Applications of multimodal CARS imaging in SSCs are anticipated to be an expanding field.

Most CARS imaging applications map cellular components at a single vibration frequency and do not have the same chemical selectivity as Raman spectroscopy. Broadband or hyperspectral CARS imaging overcome this limitation—the
A non-invasive probe has huge potential for evaluating bone tissue measurements for early detection of invasive cancer cells in the brain, based on Raman signal, and for monitoring the formation of new bone/cartilage. The exemplar developments discussed here and others are being followed by clinical trials to validate the use of non-invasive Raman spectroscopy probes in the clinic for objective diagnostics, with the aim to improve patient outcomes and extend patient survival time.

As previously discussed in this review, in spontaneous Raman scattering there are few inelastically scattered photons, resulting in a weak Raman signal [29,32,54,58] due to which it is used primarily for point measurements especially in clinical applications. In order to circumvent this, instruments for Raman spectroscopy systems could be improved by including more powerful detectors and advanced algorithms to enhance signal to noise. However, in CARS microscopy the Raman signal is enhanced, becoming more suitable for real-time imaging applications. For CARS imaging to be used as a routine technique several technological hurdles still need to be overcome. The majority of CARS applications involve imaging at a single vibrational frequency, and to date only a couple of biomolecules are typically visualized using CARS microscopy (including lipids and proteins) given the limitations related to the availability of broadband laser sources. To surmount this limitation, broadband CARS techniques are being developed, where multiple Raman frequencies are imaged simultaneously [51]. It should also be mentioned that in CARS the signal is proportional to the square of the concentration of vibrational oscillators, hence for low quantities of biomolecules, CARS sensitivity considerably decreases and imaging becomes more challenging [21,52]. Another issue is the non-resonant background observed in CARS imaging which can decrease signal-to-noise ratio. Potentially, these disadvantages can be addressed by using a similar coherent Raman technique named stimulated Raman scattering (reviewed in [72] and [73]), though it still faces many similar challenges as CARS [73,74]. Furthermore, CARS microscopy remains relatively expensive in comparison to other characterization techniques, and requires sophisticated instrumentation. CARS microscopy is now commercialized by Leica Microsystems [75], but further developments are still required to make the technique more accessible. High-quality imaging systems...
will need to be built using affordable components, to enable and enhance uptake and application of this technique. On a different note, while Raman probes are in advanced stage of translation, application of CARS through optical fibres is still in its infancy. The development of special fibres that can handle laser pulses used in CARS that have large bandwidth and low distortion in the required wavelength range is an active area of research [76].

In summary, while promising, CARS imaging still does not provide the simplicity offered by many established characterization techniques for skeletal regenerates for routine use. For adoption of Raman spectroscopy and CARS imaging in the clinic, in addition to overcoming the technological hurdles, it will also be necessary to: (i) standardize protocols (sample preparation, data analysis and presentation, etc.), (ii) perform multicentre studies, (iii) provide cost rationale/justification for the national health systems, and (iv) provide specific training to clinicians (reviewed in Sulé-Suso et al. [29]). Nevertheless, only by acknowledgment of the relative strengths/weaknesses and current challenges facing Raman spectroscopy and CARS imaging can a step change occur in methodologies to monitor the differentiation of SSCs in their natural state, and indeed other stem and progenitor populations in other tissues.

The research findings discussed in this review provide a strong case for the use of Raman-based techniques for SSC characterization. The label-free, non-destructive and non-invasive nature of Raman spectroscopy and CARS microscopy present an exciting prospective alternative to dynamically monitor SSC development for skeletal regeneration, with widespread potential in other hard and soft tissues.

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