Review Article

Shedding quantitative fluorescence light on novel regulatory mechanisms in skeletal biomedicine and biodentistry

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Summary  Digitalized fluorescence images contain numerical information such as color (wavelength), fluorescence intensity and spatial position. However, quantitative analyses of acquired data and their validation remained to be established. Our research group has applied quantitative fluorescence imaging on tissue sections and uncovered novel findings in skeletal biomedicine and biodentistry. This review paper includes a brief background of quantitative fluorescence imaging and discusses practical applications by introducing our previous research. Finally, the future perspectives of quantitative fluorescence imaging are discussed.

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1. Introduction

Undoubtedly, biologists and dental and medical scientists have long benefited from the microscope since the initial studies of biological structures reported by pioneers such as Robert Hook and Antoni Van Leeuwenhoek in the 17th century [1–3]. The imaging quality has much improved following advances in the development of technology and manufacturing. We are standing in a historical decade of microscopic development, with the awarding of the Nobel Prize in Chemistry to Osamu Shimomura, Martin Chalfie and Roger Tsien in 2008 for the discovery and application of green fluorescent protein (GFP), and the current awarding of the Nobel Prize in Chemistry to Eric Betzig, Stefan W. Hell and William E. Moerner in 2014 for the development of super-resolution fluorescence microscopy. In this review, the application of confocal imaging and image processing in hard tissue biology is discussed by introducing our research on skeletal biology.

2. Confocal microscope

Confocal laser scanning microscopy (CSLM) is designed to allow only light originating from the nominal focus to pass through a detector, such as a photomultiplier tube (PMT). To scan the specimen, a tightly focused laser is used, and the emitted light is eliminated by a pinhole placed just before the PMT, so that the image is constructed by spatially mapping the detected (well-focused) signal in accordance with the position of the scanning spot (confocality). This confocality of the scanning spot and the detected light enables the exclusion of out-of-focus blurring, thus improving the signal-to-noise ratio and image resolution.

Principaliy, CSLM can obtain a better resolution image than widefield fluorescence microscopy. However, to obtain a significant advantage by CSLM, the pinhole has to be closed to an extent where most of the emitted light is discarded, thus only a faint signal is detected. Therefore, for the practical use of CSLM, the pinhole must be opened even wider than the theoretically optimized size to obtain suitable images. In this case, it is extremely important to notice that the obtained signal may include some degree of out-of-focus signals.

3. Image processing and quantitative analyses

In the fluorescence processing system, charged-coupled devices (CCD) or photomultipliers (PMT) are used as detection apparatuses that capture visual information in a numerical form. Digitalized fluorescence images are composed of a number of pixels (or voxels in three-dimensional images) that contain numerical information such as color (wavelength), fluorescence intensity and the spatial position. By taking advantage of this digital information, reliable quantitative data can be extracted from acquired fluorescence images. Therefore, the quantitative analysis of labeled molecules or cellular functions in a spatial manner is possible, which also facilitates the comparison analysis of numerical data sets and subsequent statistical analyses. These technical advantages can be beneficial to understand a given biomedical phenomena in a compressive manner from the tissue to subcellular levels.

4. Fluorescence morphometry

Our research group has applied quantitative fluorescence imaging on histological skeletal sections by measuring the three-dimensional distributions and intensities of fluorescence signals, which is referred to as “surface rendering” [4–8]. Surface rendering is a common image processing method that provides the surface area, volume, signal intensity, fluorescence wavelength and spatial information of a given signal that is three-dimensionally distributed. Another image processing technique we have applied is filament tracing, which enables us to trace a network of cellular processes, such as neuronal dendrites and osteocytic cellular processes, in a three-dimensional manner [4]. Using this image processing method, the number, diameter and branching number of cellular processes can be numerically scored. Therefore, we can quantitatively estimate the cellular network pattern.

5. Visualization and quantification of the cellular input levels of TGF-β and BMP signaling in cartilage development

Transforming growth factor-β (TGF-β) and Bone morphogenetic proteins (BMP) have long been described as critical growth factors of cartilage development and chondrocyte
differentiation. A previous study has demonstrated that TGF-beta and BMP isoforms regulate immature chondrocyte proliferation and maturation (differentiation to hypertrophic chondrocyte), respectively [9–14]. However, the stage-specific cellular input levels of these growth factors in chondrocyte differentiation have not been well characterized.

To overcome this issue, we applied surface rendering of 3D fluorescence images to immunohistological sections stained with anti-phospho-Smad2 and anti-phospho-Smad1/5/8, which are major cellular effectors of TGF-β and BMP, respectively (Fig. 1) [6]. We measured the relative intensities of these signals in developing mouse vertebral cartilage. Consequently, phospho-Smad2 was increasingly detectable from the stage of flat proliferative chondrocytes to more mature chondrocytes, while phospho-Smad1/5/8 was highly up-regulated in prehypertrophic and hypertrophic chondrocytes.

This approach can successfully visualize when, where and how much the level of TGF-β and BMP signal is activated in differentiating chondrocytes. Furthermore, this approach facilitated the evaluation of slight developmental delay and disordered cell arrangement as a phenotype of Mesp2 knockout mice, which exhibit obvious morphological disorder of the axial skeleton [6].

6. Distinct patterns of the osteocyte network between flat and long bones

Osteocytes are embedded cells in the mineralized bone matrix [15,16]. Through cellular process running throughout the osteocytic lacunar—canaliculr system, each osteocyte maintains contact with other osteocytes, which comprise the cellular network throughout the bone tissue. This architectural structure also enables osteocytes to communicate with other bone cells, such as osteoclasts and osteoblasts located on the bone surface, therefore establishing a regulatory cellular network for bone metabolism.

Osteocytes are major mechanosensing cells. Therefore, it is postulated that the osteocyte network mentioned above is influenced by and adapted to distinct mechanical stimuli of attached muscle to each bone. To quantitatively address this hypothesis, we stained decalcified frozen bone sections
Figure 2  Three-dimensional (3D) reconstitution and surface rendering of osteocytes in the parietal bone (a, c, and e) and tibia (b, d, and f). The actin-rich cell body and cellular processes of osteocytes are visualized by a fluorescent dye-conjugated phallolidin (in green). Nuclei were counterstained by a fluorescence dye (in red). (a, b) Representative 3D-reconstituted images of the confocal z-series slices from the parietal bone (a) and tibia (b) show osteocytes and their cellular process networks. (c, d) Surface renderings of osteocyte cell bodies of the parietal bone (c) and tibia (d) from the 3D-reconstituted images (a and b, respectively) enable morphometric analyses. (e, f) Merged images of the surface renderings of cell bodies and nuclei are shown with 50% transparency of the cell bodies (in green) to visualize the relative position of the nucleus in the cell body (in red). (e, f) Dendritic tree models of the osteocytic cellular processes in the parietal bone (e) and tibia (f) from the 3D-reconstituted images (a and b, respectively). Scale bars (in white): 10 μm. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.) Modified from Himeno-Ando et al., 2013 [Elsevier] [4].

with fluorescence dyes for nuclear DNA and cellular f-actin and conducted three-dimensional quantitative analyses that enabled us to measure the volumes, surface areas and shape of nuclei and cells, and the numbers of osteocytic cellular process and its branching points (Fig. 2) [4].

These analyses revealed distinct shapes of the osteocytes and complexities of the osteocyte network between flat and long bones. Specifically, in the parietal bone, osteocytes were relatively round in shape compared to tibial osteocytes, which showed a spindle shape. The relative volume of the cytoplasm in parietal osteocytes was significantly smaller than that of tibial osteocytes, although the average volume of the nuclei was not significantly different between these two populations of osteocytes. Because it is well recognized that an augmented cellular volume, which is possibly due to an increased amount of organelles such as mitochondria and ribosomes, reflects the level of cellular activity (such as energy metabolism and protein synthesis), these observations suggest that osteocytes in the long bone exhibit a more active cellular function compared to that in the flat bone. When we measured the number and complexity of the osteocyte network composed by the osteocytic cellular process, long bone osteocytes consistently showed a higher score than flat bone osteocytes. Seeing that the tibial bone is endowed with a higher level of mechanical loading than that of the parietal bone, these findings suggest that the development, maintenance and function of osteocytes and their cellular networks are mechanically adapted and exert distinct sensitivity to loading [4].

7. Sclerostin: a potent marker of site-specific bone metabolism

Sclerostin is a secreted glycoprotein that negatively regulates the proliferation and differentiation of osteoblasts,
thus regulating bone formation through inhibiting canonical Wnt signaling [17–22]. The SOST gene encoding sclerostin is the responsible gene for osteosclerosis or Van Buchem disease, which are both associated with a high bone mass and dramatically thickened bone. Therefore, sclerostin is clearly a critical regulator of bone metabolism through regulating the osteoblast lineage. However, the involvement of sclerostin in pre- and postnatal bone development has remained unclear.

By obtaining whole views of bone sections with confocal tiling imaging and successive three-dimensional fluorescence morphometry for sclerostin distribution in osteocytic lacunae, we uncovered an increasing expression of sclerostin during postnatal bone development (Fig. 3) [8]. Furthermore, we observed that sclerostin is preferentially produced by cortical osteocytes, but not trabecular osteocytes. In vitro analyses of osteoblasts suggested that the regulatory expression of sclerostin is tightly associated with the bone formation ratio in distinct bone ages and sites. The sclerostin level was relatively low in osteocytes in bone that exert a high turnover, such as prenatal, immature postnatal bone and adult trabecular bone, while a high expression was preferentially observed in osteocytes that show a relatively low turnover, such as adult cortical bone. It is possible that highly expressed sclerostin inhibits the recruitment of osteoblast lineage cells to the bone surface, thus concomitantly and indirectly inhibiting the commitment of osteoclasts, while bone sites which express none or a small amount of sclerostin recruit and activate both osteoblast and osteocyte lineages. Therefore, sclerostin is a potential marker of site-specific bone metabolism [8].

8. Sclerostin: a potent trigger for alveolar bone remodeling in response to orthodontic forces

Osteocytes are translators of mechanical stimuli to the bone to biochemical signals that regulate the differentiation and function of other bone cells, such as osteoblasts and osteoclasts, thereby regulating bone remodeling. Sclerostin, a hallmark of osteocytes as mentioned above, is regulated by mechanical stimuli to the bone. This molecule is up- and down-regulated by unloading and loading to the bone, respectively; therefore, the expression level of this molecule may be a good marker of osteocyte responses to distinct site-specific mechanical stimuli.

Using osteocyte-ablation model mice, in which diphtheria toxin receptor-mediated cell ablation is driven by a osteocyte-specific promoter, our research group demonstrated that osteocytes played an essential role in bone resorption during orthodontic tooth movement [23]. Therefore, sclerostin was postulated to be involved in alveolar bone remodeling due to orthodontic forces.

We took advantage of multimodal confocal imaging combining zoom confocal microscopy and confocal-based...
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Figure 4 Magnified confocal images of immunofluorescence of sclerostin in the alveolar bone. Horizontal sections of alveolar bone were processed for immunofluorescence using anti-sclerostin antibodies (green) with nuclear counterstaining (red). (a, b) Optical slices obtained with zoom confocal microscopy on the control side (a) and orthodontic force side (b), respectively, are shown. The proximal-distal and palatal-buccal sides are indicated by horizontal and vertical black arrows, respectively. The red horizontal arrow indicates the direction of the orthodontic force. The white broken-lined rectangles highlight the distobuccal roots of the first molar, which are further quantitatively analyzed by three-dimensional fluorescence morphometry. Scale bars: 200 μm. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Modified from Nishiyama et al., 2015 [Elsevier] [7].

fluorescence morphometry to analyze spatial changes in the sclerostin expression in the osteocytic lacunar–canalicular system in a mouse orthodontic tooth movement model (Fig. 4) [7]. Zoom confocal microscopy enables us to acquire wide views in the order of a few centimeters at a high resolution. Therefore, this system is advantageous to map local, site-specific changes in whole tissue. We successively employed a three-dimensional fluorescence morphometry analysis as described above and observed significant down-regulation of sclerostin in the osteocytic lacunar–canalicular system specifically in tensile sites of the alveolar bone, although the sclerostin signals in compression sites were significantly higher than those in the control site. The down-regulation of sclerostin in tensile sites favors subsequent bone formation through activating osteoblasts as a result of mechanical loading induced by orthodontic forces (Fig. 5). In compression sites, the up-regulation of sclerostin is induced by unloading or physiological tension release by local periodontal ligaments, thus inhibiting bone formation and making the bone prone to resorption at this bone site. These observations suggest that spatial changes in the level and distribution of sclerostin in the alveolar lacunacanalicular system are a potent trigger for successive bone remodeling due to the orthodontic tooth. The multimodal confocal imaging analyses applied in this work enhances comprehensive understanding regarding the spatial regulation of molecules of interest from the tissue to the cellular level [7].

9. DMP1: a direct negative regulator of FGF23 production by osteocytes

FGF23, a member of fibroblast growth factor, is predominantly produced by osteocytes and hormonally regulates serum phosphate levels by targeting phosphate reabsorption in kidneys [24–29]. DMP1 is a non-collagenous extracellular matrix protein also produced by osteocytes and odontoblasts and regulates mineralization of the bone and dentin, respectively [30]. DMP1-deficient mice exhibited hyperphosphatemia through up-regulated FGF23 production, indicating that DMP1 is a negative regulator of FGF23 production [31]. The molecular mechanism underlying this phenotype, however, remains unclear.

To elucidate this issue, we first conducted double immunostaining of DMP1 and FGF23 in tissue sections of rat femurs and successive three-dimensional fluorescence morphometry (Fig. 6) [5]. The spatial distribution patterns and expression levels of DMP1 and FGF23 were observed to be inversely related, especially in cortical osteocytes. Specifically, in a single bone, DMP1-high or FGF23-high domains were designated. This finding, together with the genetic phenotype of DMP1-deficient bone, prompted us to postulate that DMP1 is a direct negative regulator of FGF23.

Inoculation of recombinant DMP1 protein to FGF23-producing osteoblasts showed significant down-regulation
of FGF23 production in a dose-dependent manner, while positively regulating the extension of cellular processes [5]. Furthermore, molecular and cellular in vitro analyses demonstrated that negative regulation of FGF23 production and positive regulation of the extension of cellular processes by DMP1 were achieved via the FAK-MAPK pathway and the FAK-ROCK pathway, respectively (Fig. 7). Taken together, these findings obtained from in vivo quantitative fluorescence imaging and in vitro signaling analyses demonstrated that extracellular DMP1 conveys intracellular signaling pathways that regulate FGF23 production and osteocyte morphology [5].

10. Morphological and functional heterogeneity of osteocytes

Osteocytes are conventionally defined as a population of cells embedded in the mineralized bone matrix. However, the findings of osteocytes obtained by our quantitative fluorescence imaging approaches have provided a distinct view of osteocytes as a morphologically and functionally heterogenic population in bones [5]. Furthermore, this heterogeneity of osteocytes is possibly regulated by local mechanical loading and bone metabolism.

11. Future perspective

In this short review, quantitative fluorescence imaging approaches applied in skeletal tissue sections were discussed. The application of quantitative fluorescence imaging has enabled our research group to successfully uncover unprecedented biological phenomena. As described above, magnified imaging modalities, such as confocal tiling and zoom confocal imaging, are an effective approach to excluding the subjective selection of regions of interest.
for further detailed image quantification. Therefore, multimodal imaging that combines several imaging modes at distinct magnifications and scale orders would be an extremely powerful tool to investigate molecular and cellular functions from the tissue to subcellular levels in a consistent manner. Now that super-resolution microscopy is being applied in biomedical science and provides imaging of unprecedented resolution (20–100 nm) [32], the application of multimodal imaging, including super-resolution microscopy, could become an attainable and powerful tool for biomedical and biodental science.

Needless to say, it is also important to understand the basic technology of fluorescence imaging and microscopy, in particular the technical limitations and biomedical application of the technique. Spectrum management of excitation wavelengths and characteristics of the fluorescence dye or fluorophore is a mandatory step to eliminate nonspecific signals from specific signals. It is also important to note that data acquired from quantitative fluorescence imaging should be fully supported or confirmed by other methods or previous literature to exclude misinterpretation of the acquired imaging data. The relevance of biological evaluations as well as optical knowledge is crucial to obtain novel insights by fluorescence imaging.

Image processing and mathematical analyses have remained ongoing subjects in the fluorescence imaging field. Live imaging-based embryonic body elongation and numerical analyses of tissue growth velocity have successfully unveiled novel regulatory mechanism by Hox genes for global body patterning [33]. Our current approach of fluorescence live imaging-based mathematical modeling is a powerful approach to unveil a tissue growth mode that is masked by a group of cells that exhibit stochastic functions [34].

Conflict of interest

The authors declare no conflicts of interest.

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