Multispectral Fluorescence Imaging

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Multispectral fluorescence imaging (MSFI) is a rapidly growing field with broad applications in both preclinical and clinical settings. Application of this novel technology in small-animal imaging and microscopy produces enhanced sensitivity and reliable quantification and resolves multiple simultaneous signals. MSFI flow cytometry can quantify multiple fluorescent parameters with morphologic or subcellular spatial details on millions of cells. MSFI has the potential to improve the accuracy of disease detection or differentiation and intrasurgical metastatic diagnosis, guide neurosurgeries, and monitor treatment response. Key Words: multispectral fluorescence imaging; multiplexing; spectral unmixing; small-animal imaging; microscopy; flow cytometry J Nucl Med 2009; 50:1563–1566 DOI: 10.2967/jnumed.109.063925

Multispectral imaging (MSI) is the synergistic combination of imaging and spectroscopy. Color is the appearance of a light most affected by wavelength (\(\lambda\)) and the observers’ visual system. Objects with similar colors are not necessarily the same. The simplest example is the spectrally pure yellow and the mixture of red and green, which have completely different spectral content but precisely the same red-green-blue (RGB) coordinates (1). Spectroscopy is the technique of breaking light down into its composite colors to identify the composition of the object. Distinguished from conventional RGB full-color imaging, MSI can obtain a high-resolution optical spectrum for each image pixel, resulting in a series of images of the same field of view that are acquired at different wavelengths. These can be stacked in 3-dimensional datasets or a cube (x, y, and \(j\)). After proper calibration, quantitative images of individual analytes can be generated.

Excellent reviews of MSI instrumentation and data analysis exist (1,2). Briefly, a camera is used to acquire the spatial information, and the spectral information is gained by scanning a dispersive element to record spectra for each image. Electronic tunable filters such as an acousto-optic tunable filter and liquid crystal tunable filters (LCTFs) are preferable to mechanically scanning dispersive devices (filter wheels, monochromators) because they are quiet, fast, compact, and stable and demonstrate increased spectral selectivity, spectral purity, and flexibility. The disadvantages of electronic tunable filters include low light throughput, photobleaching, and inappropriate capture if significant sample or camera movement occurs during the acquisition or if high temporal resolution is needed to capture certain events (e.g., calcium signaling transients) (3). The typical method for data analysis of MSI is a least-squares-fitting linear unmixing approach with additional constraints (such as nonnegativity), resulting component images, and a final composite image. This approach is limited to spectral analysis only, whereas combining the rich spatial information in the images with the spectral data is a more powerful and evolving field (4). Multispectral fluorescence imaging (MSFI) coupled with microscopy and flow cytometry are commercially available now (4,5).

MSI technologies have been widely used in the fields of astronomy, geology, agriculture, industry, and forensics (1,2). With the vast development of filters, detectors, data-analysis techniques, interdisciplinary approaches, and fluorescent dyes, MSFI is a rapidly growing field with applications in cell biology, preclinical drug development, and clinical pathology. Combining MSI with small-animal imaging and microscopy produces enhanced sensitivity, reliable quantitation, and resolved multiple simultaneous signals (3,6). MSFI is particularly useful for analyzing objects that have multiple fluorescent labels that may have similar RGB color or that may be localized in the same or spatially overlapping compartments; this modality is also effective for evaluating objects that have strong whole-animal autofluorescence (Fig. 1) (4,7,8). Here we review a selection of preclinical and clinical studies describing the use of MSFI applications.

PRECLINICAL APPLICATION OF MFSI

To Assess Tumor Vascularization and Monitor Antiangiogenic Therapeutic Response

Previous studies imaged tumor angiogenesis with fluorescent proteins (9,10). MSFI provides a powerful, noninvasive means by which to assess tumor vascularity and to monitor the response to antiangiogenic therapy. For example, in mice treated with sunitinib, a near-infrared dye–labeled monoclonal antibody (mAb)–based probe targeting vascular endothelial growth factor receptor-2 was imaged in vivo using an LCTF-based Maestro MSFI system (CRI Inc.). Tumor fluorescence intensity was quantified and normalized after autofluorescence removal. This study demonstrates that MSFI provides rapid, noninvasive, and longitudinal assessment of the efficacy of novel angiogenesis-directed therapies in preclinical models of cancer (11). To overcome constraints associated with exogenously administered vascular imaging probes such as high cost, toxicity, inconsistent delivery, differential biodegradation among animals, and high tumor accumulation resulting from vessel leakiness, a novel method to noninvasively image tumor vascularization in fluorescent tumors without exogenous imaging probes was introduced. LCTF-based Maestro and Nuance
systems (CRI Inc.) were used for in vivo MSFI to study the biology of tumor angiogenesis and to monitor the effects of antiangiogenic therapies with bevacizumab. In a mouse model bearing tumors expressing red fluorescent protein, the spectral fluorescence signatures of vascular and avascular components allow for the tumor vasculature to be imaged and quantified without contrast agents (Fig. 2). This technique provides real-time imaging of tumor vascularization and monitoring of the antiangiogenic therapeutic response in mice, without the concerns specific to an exogenously administered probe (12).

To Generate Region-Selective Brain Maps of Vascularized Brain Parenchyma

A fluorescein isothiocyanate–derivatized tomato lectin–based molecular imaging probe was combined with the Maestro system to generate region-selective brain maps of vascularized brain parenchyma ex vivo in rats and rabbits. Images were unmixed to get rid of brain tissue autofluorescence and resolve arterial blood and specific imaging probe fluorescence signal. After further refinement of imaging probes and metrics, this novel preoperative endovascular brain-mapping approach may facilitate neurosurgical guidance and therefore improve clinical outcomes after neurosurgical resections (13).

Automatable Analysis of Multiplexed Immunolabeled Samples

More than 6 markers can be quickly differentiated from each other and from autofluorescence using MSFI, thereby allowing for the localization of multiple probes and avoiding the need for serial sections. For example, Mansfield et al. performed 6-marker analyses in 1 sample with MSFI. Quantitative and marker-specific image data were presented for integrative system biologic studies. The drawbacks of this approach are that it is too much work to prepare a series of controls of all of the combinations of markers and the method is less optimal for membrane or cytoplasmic marker analyses (4).

To Differentiate Tissue Type and Provide Cellular-Level Diagnostic Information

The slit-scan confocal microendoscope uses a catheter based on a fiber-optic imaging bundle. This multispectral system uses a prism as a dispersive element to collect a full multispectral image with spectral resolution of 2.9 nm at 500 nm and 8.4 nm at 750 nm. Human ovary and esophagus and mouse peritoneal wall and liver stained with acridine orange and rat intestine tagged with 4-(4-(dihexadecylamino)-styryl)-N-methylpyridinium iodine (Invitrogen) were imaged ex vivo. This method of MSFI can distinguish tissue types and acquire high-quality images; it is minimally invasive and has the potential to provide cellular-level in vivo diagnostic information as a clinical diagnostic instrument (14).

Combining MSFI with Quantum Dots (MSFI-QDs)

Quantum dots (QDs) are novel nanocrystal fluorophores with constant broad excitation spectra, sharp and symmetric tunable emission spectra, improved brightness, superior photostability, and simultaneous excitation of multiple fluorescence colors. QDs have been successfully applied in various imaging applications for in vitro diagnostics and in vivo imaging because of these unique optical properties. MSFI-QDs can efficiently remove background and precisely delineate weak spectral signatures to present ultrasensitive and multiplexed imaging of molecular targets in vivo (15,16). Full reviews on MSFI-QDs in multiplexed immunohistochemistry and in situ hybridization (ISH) are available (17,18). Another use of MSFI-QDs is as a feasible substitute for microarray analysis. In this instance, multiplexed QD-labeled oligonucleotide probes can be used for ISH in human biopsies. This technique was tested in a study in which single and multiplexed QD-ISH was performed in samples with acute leukemia and follicular lymphoma. Spectral unmixing enables the separation of spatially colocalized signals. MSFI-QDs allow the quantitative characterization of multiple-gene expression, thereby making them useful for the analysis of processed human clinical tissues (19). MSFI-QDs were also tested using streptavidin-conjugated QDs to detect up to 7 signals in tonsil and lymphoid tissue. Slides were analyzed using confocal laser scanning microscopy. Five streptavidin-conjugated QDs were used on the same tissue section and could be analyzed simultaneously on routinely processed sections (Fig. 3). This multiplexing method has the potential to uncover the clinically relevant multidimensional cellular interactions that underlie diseases (20).
CLINICAL APPLICATION OF MSFI

Combining MSFI with 2-Photon Excitation (TPE) to Detect Cutaneous Tumors

TPE fluorescence microscopy offers high spatial resolution, deep penetration, and low photodamage. To detect cutaneous tumors, a combined TPE and MSFI system (MTPE) was used to identify different intrinsically fluorescent molecules (nictinamide adenine dinucleotide, tryptophan, keratins, melanin, elastin, cholecalciferol, and others). This technique provides functional information correlated with tissue structure in physiologic and pathologic states. MTPE is effective in differentiating healthy skin from basal cell carcinoma (21). As more samples are analyzed, it should be possible to differentiate between different types of malignancies as well.

Application of MSI Flow Cytometry

MSI flow cytometry can quantify multiple fluorescent parameters with morphologic or subcellular spatial details on millions of cells (5,22). Nuclear translocation of NFκB initiates the transcription of host defense-related critical genes. To quantitatively measure the translocation of signaling molecules from the cytoplasm to the nucleus in cells, cells were stained with fluorescein isothiocyanate–conjugated anti-NFκB and 7-AAD (DNA stain) and imaged by flow cytometry. A similarity score that quantifies the correlation of the spatial distribution of the stained, translocating signaling molecule with nuclear staining was generated for each cell using Pearson correlation coefficients to assess the nuclear translocation. This approach represents an advance over conventional microscopy or biochemical fractionation and Western blotting for high throughput single-cell or subpopulation analysis (23,24). MSI flow cytometry can not only distinguish live cells from cells in the early or late phases of apoptosis or from necrotic cells (25) but also accurately quantitate apoptosis based on morphologic characterization and allow a more complete and robust analysis of apoptosis than conventional microscopy (26). Covalent deposition of C3 fragments (C3b/C3b) on cells is closely associated with cell-bound chimeric humanized anti-CD20 mAb rituximab (RTX). To quantitatively analyze colocalization of C3 with RTX, Raji cells were stained with fluorescently labeled RTX and mAbs specific for C3b/C3b and then imaged in flow and analyzed for colocalization of fluorescent probes. Analysis verified that most deposited C3b/C3b is colocalized with bound RTX (27).

Autofluorescence Analysis in MSFI

MSFI of oral tissue autofluorescence in vivo with a 4-channel multispectral digital microscope can improve the detection of oral neoplasia. Decreased blue–green autofluorescence and increased red autofluorescence in lesions were observed in patients with histologically confirmed neoplasia (28). MSFI of the ocular fundus autofluorescence may differentiate age-related macular degeneration...
and diabetic retinopathy (RD). Autofluorescence in age-related macular degeneration patients was dominated by the red emission, whereas RD green-shifted. Lipofuscin is the dominant fluorophore in the red channel. The green shift in RD may be related to increased advanced glycation end product concentrations (29). MSFI of sentinel lymph node autofluorescence during surgery can provide a rapid method to verify whether a tumor has metastasized to help determine follow-up treatment. For example, to evaluate lymph nodes in colorectal and gastric tumors, autofluorescence microspectroscopy and MSI autofluorescence microscopy have been used to analyze the autofluorescence emission of metastatic lymph node sections. The autofluorescence pattern and the emission spectrum of metastatic lymph nodes can distinguish them from the normal ones (30). MSFI of autofluorescence has the potential for detecting oral neoplasia, discriminating fundus autofluorescence fluorophores and intrasurgical diagnosis of metastatic lymph nodes.

**Spatial–Spectral Analysis in MSFI**

Using exclusive spectral information from an LCTF-based MSFI system with image analysis software (Environment for Visualizing Images) can identify malignant urothelial cells in urine cytology specimens, with a sensitivity and specificity of 82% and 81%, respectively, or a combined sensitivity and specificity of 73% (31). To increase the accuracy and specificity of the diagnosis and follow-up of bladder cancer, the spatial–spectral features of multispectral image stacks acquired with the same MSFI system were analyzed using the GENieX Image Exploitation (GENIE) package (Los Alamos National Laboratory). GENIE showed a combined sensitivity and specificity of 85% and 95% in samples from 2 separate institutes. Furthermore, GENIE showed efficiency superior to a cytopathologist with respect to predicting the follow-up results (32). MSFI of fine-needle aspirations of follicular adenoma and parathyroid adenoma with the Nuance system and spatial–spectral analysis with CRI-MLS needle aspirations of follicular adenoma and parathyroid adenoma (33). MSFI of autofluorescence has the potential for detecting oral neoplasia, discriminating fundus autofluorescence fluorophores and intrasurgical diagnosis of metastatic lymph nodes.

**REFERENCES**