Review Article

Neurosurgical confocal endomicroscopy: A review of contrast agents, confocal systems, and future imaging modalities

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Abstract

Background: The clinical application of fluorescent contrast agents (fluorescein, indocyanine green, and aminolevulinic acid) with intraoperative microscopy has led to advances in intraoperative brain tumor imaging. Their properties, mechanism of action, history of use, and safety are analyzed in this report along with a review of current laser scanning confocal endomicroscopy systems. Additional imaging modalities with potential neurosurgical utility are also analyzed.

Methods: A comprehensive literature search was performed utilizing PubMed and key words: \textit{In vivo} confocal microscopy, confocal endomicroscopy, fluorescence imaging, \textit{in vivo} diagnostics/neoplasm, \textit{in vivo} molecular imaging, and optical imaging. Articles were reviewed that discussed clinically available fluorophores in neurosurgery, confocal endomicroscopy instrumentation, confocal microscopy systems, and intraoperative cancer diagnostics.

Results: Current clinically available fluorescent contrast agents have specific properties that provide microscopic delineation of tumors when imaged with laser scanning confocal endomicroscopes. Other imaging modalities such as coherent anti-Stokes Raman scattering (CARS) microscopy, confocal reflectance microscopy, fluorescent lifetime imaging (FLIM), two-photon microscopy, and second harmonic generation may also have potential in neurological applications.

Conclusion: In addition to guiding tumor resection, intraoperative fluorescence and microscopy have the potential to facilitate tumor identification and complement frozen section analysis during surgery by providing real-time histological assessment. Further research, including clinical trials, is necessary to test the efficacy of fluorescent contrast agents and optical imaging instrumentation in order to establish their role in neurosurgery.

Key Words: Brain neoplasm, confocal endomicroscopy, fluorescent dyes, intraoperative imaging, neuronavigation, optical imaging
INTRODUCTION

Glioblastoma (GBM) is the most common primary malignant brain tumor with approximately 16,000 Americans diagnosed each year.\textsuperscript{[10]} Following surgical resection of this tumor, the incidence of recurrence at the primary tumor site is greater than 80%, and the median patient survival is only 12.2-18.2 months.\textsuperscript{[19,117]} GBM extent of resection is directly correlated with patient survival in some studies, with the maximum survival benefit achieved at greater than 98% tumor volume reduction.\textsuperscript{[156,35]} Aggressive treatment with maximal resection plus adjuvant radiation and chemotherapy can increase patient survival by a range of 3-4 to 7-12 months.\textsuperscript{[118]} While debate exists about the exact relationship between degree of resection and outcome, achieving maximal resection with minimal deficits remains a key neuro-oncological goal.

Such maximal resection is difficult to achieve due to the infiltrative nature of GBM and the need to preserve eloquent brain regions. Neurosurgeons use a variety of adjuncts such as intraoperative magnetic resonance imaging (MRI), neuronavigation, ultrasonography, and macroscopic fluorescence to help guide surgical resection, but none of these technologies provide detail at a cellular level.\textsuperscript{[98,107]} A means for examining neoplastic tissue at a cellular level could improve differentiation between tumor and normal brain and facilitate tumor cytoreduction. Successfully making this distinction could contribute to a higher deficit-free resection rate. Arguably, actual visualization of neoplastic cells could be preferable to resecting the tumor to the margins defined by gadolinium-enhancement on MRI, which are known to not reflect the true extent of tumor.

Advancing optical technologies, such as a laser scanning confocal endomicroscopy (LSCE), provide real-time histopathological information of GBM in vivo. This technology has been used to distinguish diseased and normal tissue in other systems, including the gastrointestinal (GI) tract,\textsuperscript{[19,23,34,37]} the cervix,\textsuperscript{[165]} and the bladder.\textsuperscript{[120]} LSCE allows microscopic tissue analysis with cellular and subcellular detail in vivo,\textsuperscript{[12,34]} and contrast of morphological details can be enhanced with the application of exogenous fluorescent dyes.\textsuperscript{[33]} This technology has recently been studied in the neurosurgical setting in an effort to improve tumor resection by providing immediate histological assessment of the brain-to-neoplasm interface.\textsuperscript{[28,64,88,89]}

In this review, we assess the properties and utility of each of the three fluorophores currently available for neurosurgical clinical use, as well as the specifications of the two currently available LSCE systems. Additionally, future optical technologies currently under investigation are discussed in relation to neurosurgical applications.

CLINICALLY AVAILABLE EXOGENOUS AGENTS

Exogenous dyes effective for neurosurgical applications require neurolocalization and selective contrast of neurohistopathological structures. Theoretically, effective tumor targeting occurs through enhanced permeability and retention, where vascular proliferation and anatomical abnormalities (such as limited lymphatic drainage and venous return) allow exogenous agents to concentrate at the tumor site.\textsuperscript{[14,36,46,167,93,103]} Thus, the breakdown of the blood-brain barrier in glioma allows exogenous dyes to concentrate in neoplastic regions and contrast abnormal tissue.\textsuperscript{[79]}

The Stokes shift describes the relationship between single photon excitation and emission of fluorophores, which is absorption of short-wavelength (higher energy) photons and emission of longer wavelength (lower energy) photons.\textsuperscript{[76]} Each exogenous dye has specific absorption and emission spectra, which can be filtered to visualize the fluorescent contrast agent in susceptible tissues.\textsuperscript{[108]}

Properties of effective exogenous fluorescent agents include minimal phototoxicity, scattering, and signal attenuation. Phototoxicity occurs with fluorophore excitation and resultant production of reactive oxygen species (ROS),\textsuperscript{[1,20,29,49,63,115,121]} which can damage cellular structures and reduce the signal of the fluorophore, leading to photobleaching.\textsuperscript{[6,94,95]} Ideal fluorophores have a high quantum yield (i.e. a high ratio of emitted fluorophores to absorbed fluorophores).\textsuperscript{[108]}

Scattering, signal attenuation, and native autofluorescence all limit the depth at which a fluorophore can be visualized within tissue.\textsuperscript{[31,59]} Scattering and native autofluorescence decrease as wavelength increases, with the infrared emission spectra (700-900 nm) being ideal for imaging deeper cellular structures.\textsuperscript{[50,108]}

FLUORESCEIN SODIUM

Optical and chemical properties

Fluorescein sodium is a small organic molecular salt that has an excitation maximum of 494 nm and an emission maximum of 521 nm.\textsuperscript{[79]} It readily crosses capillaries, provides fluorescent contrast in the extracellular matrix,\textsuperscript{[51]} and has a urine clearance of 24-36 h after intravenous injection.\textsuperscript{[79]} The amount of fluorescein delivered to a tumor site is increased by the breakdown of the normal blood-brain barrier, which becomes useful in neurosurgical applications.\textsuperscript{[5]}

Neurosurgical applications

Fluorescein sodium has been approved by the United States Food and Drug Administration (FDA) for ophthalmoscopic examinations of the retina since the 1960s.\textsuperscript{[152]} In the field of neurosurgery, fluorescein...
has been used in cerebral angiography to detect arteriovenous malformations, superficial temporal artery-middle cerebral artery anastomoses, and to aid in treating cerebral aneurysms.\[26\]

Fluorescein was first reported to contrast brain tumors in a 1948 study showing tumor visualization in 95.7% of patients. This demonstrated the ability of fluorescein to localize at the site of the tumor, mainly due to breakdown of the normal blood-brain barrier. Thus, fluorescein is not a tumor-specific agent, but it is excellent for visualizing regions of compromised neurovasculature. The presumption in its use for neuro-oncology is that these areas correspond to the enhancing regions, which also correspond to bulk tumor. It is believed that fluorescein does not differentially leak into and stain tumor cells that have infiltrated beyond the gross enhancing margin. However, resection of these cells within otherwise normal surrounding brain may also raise the risk of neurological deficit.

Several decades later, fluorescein was utilized to macroscopically demarcate brain tumors during resection under white light without surgical microscopes. One study reported that gross total resection (GTR) of GBM was achieved in 84.4% of patients who received fluorescein compared with only 30.1% of patients who did not. These results were reproduced in a prospective study, however, it should be noted that both studies failed to show a significant survival benefit despite higher rates of GTR in patients who received fluorescein. In a separate study of low-grade gliomas, use of fluorescein was shown to significantly increase the achievement of GTR, as well as to increase 6-month progression-free survival. This technique has been considered safe to use, easy to perform, and requires no extra surgical equipment for tumor resection. However, the extent of resection and visualization of the tumor border is dependent on the surgeon’s judgment of fluorescence, which may be limited in tumor areas with intact vasculature.

In 1998, a modified surgical microscope was developed equipped with dichroic mirrors specific for fluorescein-guided resection of malignant gliomas. This system allowed more accurate demarcation of tumor boundaries and reduced the amount of fluorescein given when compared with studies that visualized tumor macroscopically. The study reported 100% gross GTR resection in all patients, except those patients with tumors in eloquent areas, such as the basal ganglia and brain stem, which limited the resection.

LSCE was first used to image GBM in animal models in 2010. Using fluorescein delivered intravenously, researchers were able to distinguish between tumor and nontumor regions using a handheld confocal microscope. This technology identified infiltrating tumor margins at a cellular level in vivo for the first time. Another study conducted with fluorescein-guided LSCE demonstrated the ability of histological in vivo imaging of normal and neoplastic tissue in rat allograft models in vivo and human tissue ex vivo. In vivo fluorescein staining depicted tissue morphology and cellular structures as well as capillaries in normal tissue, while showing rapid tumor growth and massive extravasation in the region of the tumor. Fluorescein application ex vivo allowed the diagnosis of various brain tumor subtypes, not only GBM, using LSCE. However, there was limited depth perception, and it was noted that near-infrared probes may solve this issue.

Also in 2010, fluorescein-guided LSCE technology was utilized in a human trial, and it provided assessment of tumor grade, tumor histology, and tumor margins for a variety of tumor subtypes. Following this proof-of-principle study, a larger study imaging various brain tumors including meningiomas, schwannomas, gliomas of various grades, and a hemangioblastoma was performed. LSCE with fluorescein was able to correctly diagnose and correspond with traditional histopathological findings with an accuracy of 92.9% when the pathologist was given a list of clinically relevant diagnoses to choose from. Though interpretation was somewhat biased toward

Figure 1: Images obtained with intraoperative endomicroscopes of various clinically available fluorescent contrast agents. (a and b) Fluorescein-induced fluorescence of oligodendroglioma (Grade II), and corresponding H and E stain. (c and d) ICG-induced fluorescence of glioblastoma cells in a mouse model, and corresponding H and E stain. (e and f) 5-ALA induced fluorescence in low-grade glioma, and corresponding H and E stain. Figures a and b from Eschbacher et al.; used with permission from Journal of Neurosurgery. Figure c used with permission from Barrow Neurological Institute. Figure d from Martirosyan et al.; used with permission from Journal of Neurosurgery. Figures e and f from Sanai et al.; used with permission from Journal of Neurosurgery
a list of potential pathologies, this technology was able to give a good direction toward the final diagnosis and intraoperatively discern regions of tumor infiltration in both low- and high-grade gliomas [Figure 1a and b]. In this study, LSCE images were collected in grayscale, however, they can be pseudocolored for ease of interpretation by a clinician. The results from these two studies demonstrate the feasibility of this technology as a diagnostic and therapeutic tool, as it can help identify many of the pathognomonic cytoarchitectural features of various brain tumors and aid in the intraoperative diagnosis and resection of various central nervous system (CNS) tumors. Fluorescein in this case can both guide the operator grossly to areas to be viewed with LSCE and provide the contrast to allow cellular visualization. The next phase in validation of this technique is to provide optical “biopsies”, which can then be compared in blinded fashion against frozen section, to evaluate its overall diagnostic accuracy.

**Safety**

Fluorescein is FDA approved and is widely used in the field of ophthalmology as well as in GI studies. Most fluorescein clinical studies in neurosurgery report no serious adverse effects with use of the fluorophore, however, in an international study using fluorescein, 1.4% of patients developed minor complications such as nausea and vomiting. No patients suffered severe adverse events in this study. Deleterious effects can occur with rapid high dose administration (20 mg/kg) of fluorescein, but these high doses can be avoided when fluorescein is imaged using a surgical microscope that requires less dosage compared with macroscopic viewing.

**INDOCYANINE GREEN**

**Optical and chemical properties**

Indocyanine green (ICG) is a near-infrared fluorescent agent with maximal excitation at 778 nm and emission spectra range of 700-850 nm in serum. It is fairly water-soluble, which allows it to be given intravenously and cleared through renal and bile excretion. ICG concentration within a tumor site is enhanced by breakdown of the normal blood-brain barrier. ICG has a greater tissue penetrance than visible-wavelength fluorophores, such as 5-ALA and fluorescein. ICG is an anionic, amphiphilic, tricarbocyanine probe, which allows it to have a high affinity for proteins, such as albumin, and allows visualization of solid tumors, but may also cause higher levels of nonspecific binding. Similar to fluorescein, ICG provides nonspecific contrast in areas of permeable neurovasculature.

**Effectiveness in neurosurgical applications**

ICG has been given intravenously for blood vessel angiography, identifying extrahepatic bile ducts and detecting liver metastases. It can also be given subcutaneously for sentinel lymph node mapping for breast, anal, and GI cancer as well as assessing lymphatic drainage for lymphedema.

The first application of ICG for macroscopic demarcation of glioma tumor margins was investigated in 1993. The results demonstrated that ICG was able to contrast the fluorescent tumor tissue within 1 mm of the histological tumor margins in an animal model. However, this technique is not effective in distinguishing between malignant cells and other areas of the brain that may incidentally uptake the injected dye. Cellular visualization using ICG contrast may overcome these limitations.

With the development of an infrared LSCE, ICG was first investigated in vivo to diagnose liver steatosis and fibrosis. Recently, ICG has proven useful in neurosurgical applications. In animal glioma models, ICG imaging with a near-infrared wavelength confocal endomicroscope has been effective at detecting tumor, transitional zones, and normal brain. LSCE technology can confirm, at the cellular level, regions fluorescently labeled with ICG are truly representative of the tumor. In contrast, macroscopic detection is more subjective to the surgeon. Intravenously administered ICG is immediately localized to a tumor site, and the fluorescent signal remains in the tumor up to 1 h after injection, with constant imaging, demonstrating limited photobleaching and clearance. The delivery of the dye to the site of the tumor relies on binding to serum proteins and the damaged vasculature primarily located at the site of the tumor. However, with time the dye will diffuse into surrounding tissue. These properties allow real-time, in vivo assessment that differentiates glioma tissue and normal brain. In contrast to visible wavelength fluorophores, ICG provides visualization of deeper tissue structures due to its infrared excitation and emission spectra. However, depth of imaging still remains limited to a few hundred microns deep to the imaging surface.

**AMINOlevulinic acid (5-ALA)**

**Optical and chemical properties**

Produced in the mitochondria, 5-ALA is a natural precursor for the production of protoporphyrin IX (PpIX).
PpIX is a fluorescent molecule that binds membrane lipids and has an excitation range of 375-440 nm and emission range of 640-710 nm in vivo. Overloading this pathway with exogenous 5-ALA causes the collection of PpIX to fluorescently detectable levels in cells. As the production of PpIX occurs in situ in mitochondria, fluorescence is limited to cells. This decreases fluorescent signal in blood or edematous regions of the operative field. Furthermore, neoplastic cells demonstrate preferential uptake of exogenous 5-ALA and increased collection of PpIX, making the fluorescent signal more robust in these abnormal tissues and making intraoperative visualization feasible, especially in intracranial tumors of higher grades. Photobleaching occurs with PpIX fluorescence level dropping to 36% after 25 min under violet light or 87 min under white light. It can be limited by allowing excitation and white light to penetrate as small an area as possible in the surgical field. Phototoxicity is limited with 5-ALA, since fluorescent microscopes do not produce sufficient energy for significant ROS production.

Neurosurgical application

Due to its favorable optical and chemical properties, clinical applications of 5-ALA have been extensively studied in an array of medical specialties and a variety of tissues. Currently, the FDA has approved 5-ALA and its derivatives for research diagnostic applications in endoscopic, photodynamic detection of bladder cancer and residual glioma, as well as the treatment of basal cell carcinoma and actinic keratosis.

The first study of 5-ALA-induced fluorescence of human gliomas was reported in 1998. After oral administration of 5-ALA, PpIX fluorescence was exclusively observed in tumor regions using a surgical microscope modified for fluorescence. Histological analysis demonstrated 85% sensitivity, 100% specificity, and 90% accuracy for predicting the presence of malignant tissue from 5-ALA-induced macrofluorescence in this study. Photobleaching and spontaneous deterioration occurred in this study with excessive exposure to light. Necrotic and edematous regions had little fluorescence. Overall, the study revealed a high degree of congruity between macroscopic visualization of PpIX fluorescence and the presence of neoplastic tissue. However, the investigators did not determine the outcome of 5-ALA-guided tumor resection.

A follow-up study using similar operative methods was undertaken to determine the efficacy of fluorescence-guided resection with 5-ALA in 52 patients with GBM. This study showed that GTR was achieved in 63% of patients. Postoperative residual tumor was also assessed with contrast-enhanced MRI, which proved to be less sensitive in showing residual tumor compared with fluorescent enhancement of glioma in vivo. Finally, the level of residual solid tissue fluorescence was shown to correlate with patient survival: Patients with minimal to no fluorescence had higher survival rates than those with high levels of fluorescence postoperatively.

The results from this study led to a randomized controlled trial that further determined the efficacy and safety of 5-ALA-induced macrofluorescence-guided GBM resection. The phase III clinical trial with 322 patients demonstrated greater tumor resection using this technique compared with patients who underwent conventional microsurgery under white light (65% vs. 36% GTR). Overall progression-free survival time also increased by 1.5 months in groups administered 5-ALA, which included an increase of 6 months in the subgroup of patients over 55. Though survival time increased, quality of life was not measured and may be an important factor in deciding tumor resection strategy when using this technology.

While 5-ALA-induced fluorescence is successful for both diagnosis and resection of high-grade gliomas, there are many studies reporting that macroscopically detectable fluorescence does not occur in low-grade gliomas. Therefore, an investigation was conducted utilizing a hand held intraoperative confocal microscope to visualize low-grade gliomas after standard administration of 5-ALA. This technology allowed visualization of 5-ALA-induced fluorescence of low-grade glioma tumor and tumor margins in vivo that corresponded with standard histopathology. Overall, both pre- and postoperative MRI in 10 patients indicated a median of 95% extent of tumor resection using this imaging modality. This technology may permit intraoperative fluorescence contrast of low-grade gliomas if necessitated in the treatment plan of a patient. Some controversy still exists as to the biology of 5-ALA metabolism in low-grade gliomas and the need for complete resection by MRI. However, as the distinction between low-grade glioma and surrounding brain can be especially challenging, this remains an important area for future work.

Safety

Though 5-ALA administration is considered safe, minor systemic side effects have been reported, including nausea, vomiting, and hypotension as well as increased sensitivity to sunlight up to 48 h after administration. As with other fluorophores, these side effects are limited with lower dosing.

INTRAOPERATIVE ENDOMICROSCOPY SYSTEMS

Commercially introduced in the 1980s, confocal microscopy has been extensively used for molecular
imaging of thick tissues in the biomedical sciences. Miniaturization of this technology into an endoscope (LSCE) was fueled by a need to diagnose in vivo premalignant lesions in epithelial cancers.\cite{9} LSCE has been extensively tested for visualizing Barrett’s esophagus, urothelial bladder neoplasia, and cervical intraepithelial neoplasia.\cite{21,105,120}

LSCE provides noninvasive histological images in vivo through optical sectioning.\cite{17,44} Optical sectioning allows the microscope to selectively image a certain focal depth by mechanically filtering most out-of-focus photons above and below the focal plane with a pinhole.\cite{48} This enables imaging of thick tissue with greater contrast compared with conventional wide-field microscopy, which generates out-of-focus background light and scatter.\cite{13} Furthermore, endomicroscopy allows for three dimensional image reconstruction and greater imaging parameters.\cite{113} Most importantly, this technology permits real-time histopathological analysis with increased diagnostic yield without the traditional histopathological process of excision, fixation, and staining.

While neurosurgeons are equipped with macroscopic fluorescence-guided surgery, the distinction of healthy and tumor cells remains a challenge for immediate tumor distinction during tumor resection. Therefore, the application of this technology may provide novel cellular details of CNS tumor architecture. Confocal endomicroscopy using fluorescent dyes has successfully been researched in the imaging and resection of brain tumors in vivo in animal models and to some extent in human subjects.\cite{23,105,120} These results demonstrate in vivo visualization of brain tumors using fluorophores. Specifically, 5-ALA and LSCE systems can facilitate intraoperative diagnosis and real-time, fluorescence-guided resections of brain tumors. Further clinical trials are needed for nonspecific fluorescent agents, such as ICG and fluorescein that accumulate in the extracellular compartment. There are two clinically available confocal endomicroscopy systems, which are assessed in this review for their possible application for neurosurgical purposes: Optiscan/Pentax ISC-1000 and Mauna Kea Cellvizio.

**Optiscan**

The Optiscan is a conventional endoscope with a miniaturized confocal microscope at the tip designed mainly to image the lower GI tract, but has also been used for imaging of the stomach, duodenum, distal esophagus, and the cervix.\cite{48} This system provides excellent image clarity but requires image stabilization.\cite{23} The miniature confocal microscope scanner is within a rigid probe that is connected to an optical unit and PC unit via a flexible umbilicus [Figure 2a]. The probe uses a distal scanner with a single optical fiber, which contains both excitation and detection capability.\cite{48} A miniaturized objective lens comes into direct contact with tissue, creating a 475 × 475-μm field of view. The operator is able to control the focus depth through a foot pedal at 4-μm steps anywhere from 0 to 250 μm beyond the window surface.\cite{48} [Figure 2c]. Similar to the Cellvizio LSCE, both the excitation light and the emitted fluorescence are transduced and collected at the same point, which in this case is the single optical fiber. 488-nm excitation is emitted from the solid-state laser with a maximum power of 1 mW, while light at wavelengths outside the red-violet spectrum is filtered during detection, limiting perceived fluorescence to wavelengths of 505-705 nm.\cite{23,105} The Optiscan collects images at a rate of up to 0.7 frames per second. A customized rigid Optiscan probe was used in the fluorescein LSCE study by Eschbacher et al.\cite{24}

**Cellvizio**

The Cellvizio LSCE is a probe-based endomicroscopy system that has been used for in vivo imaging of the upper and lower GI tract, as well as the human bladder.\cite{22,54,96} Because of its compact size and maneuverability, this system is well-suited for microsurgical and experimental imaging techniques, such as examination of microvasculature, detection of dysplastic cells, and observation of neuroblast migration.\cite{17,44,57} The system also incorporates a range of miniprobes that can be inserted into an endoscope or used independently.\cite{48} [Figure 2b]. There are various miniprobes that differ in their field of view, fixed working distance, and lateral and axial resolution.
Each miniprobe has specific applications based on these technical parameters, one of which is designated for imaging the surface of the brain. The system as a whole can collect full frame images at 12 frames per second. The Cellvizio LSCE is composed of three main components: (i) a laser optoelectric unit, (ii) an array of optical fiber mini-probes linking the proximal scanning device and the micro-objective, and (iii) software that controls the system and manages the image data. A 488- or 660-nm laser source provides the excitation light. Real-time imaging is accomplished with a 4-kHz oscillating mirror for horizontal scanning and a galvanometric mirror for frame scanning, resulting in a frame rate of 12 Hz. The same fiber bundle that is used for illumination of the specimen collects fluorescence that is then diverted to the photodetector. The resultant raw image is first processed using an algorithm designed to eliminate inherent artifact resulting from the fiber optic bundles. Therefore, the system is first calibrated by measuring intrinsic autofluorescence of each fiber by imaging a nonfluorescent sample, in addition to the transmission and collection rate of each fiber. Distortions generated by the system can then be accounted for, thereby providing images capable of quantitative measurements.

**FUTURE DEVELOPMENTS**

**Coherent anti-Stokes Raman scattering microscopy**

Similar to MRI, coherent anti-Stokes Raman scattering (CARS) microscopy is a microscopy technique that produces an image based on intrinsic vibratory properties of the specimen. Using an oscillating laser system that produces molecular vibration in the specimen and a modified beam-scanning commercial microscope that detects the resonance of “anti-Stokes” oscillating molecules, detailed images of normal brain structures can be generated, as well as margins of intracranial gliomas, can be generated. These images have been shown to be both sensitive and specific for cancerous tissue at the cellular level. CARS depict the microenvironment and cellular morphology that can potentially provide the investigator with a means of diagnosis without the need for biopsy and hematoxylin and eosin staining. Therefore, in addition to the benefit of discerning normal versus pathologic CNS tissue in vivo for tumor resection, there is a diagnostic capability to CARS as well. However, further development of CARS for the in vivo neurosurgical setting will require its miniaturization for practical uses in the operating room.

**Confocal reflectance microscopy**

Using light scattering technology, confocal reflectance microscopy (CRM) is an imaging technique that noninvasively images a thin plane of tissue with high resolution. CRM generates contrast by utilizing the intrinsic differences in refractivity of targeted tissue. Images are generated with relatively lower refractive structures contributing fewer signals in the resultant image. Since CRM does not rely on a Stoke’s shift, it introduces a fraction of energy into tissue samples compared with fluorescence imaging techniques. Overall, this technique allows real-time analysis of tissue cellularity and cytoarchitecture with minimal production of ROS. Limitations of CRM technology include limited depth (200-300 μm) and sharpness to evaluate intracellular structures. Imaging depth can be increased with increasing wavelengths, but can compromise resolution and contrast.

CRM has been investigated to image cellular and subcellular tissue architecture, diagnose dermatological conditions noninvasively, identify cancerous tissue and margins intraoperatively, and assess cellular and subcellular detail of diseased and normal hepatic tissue. CRM has recently been used to determine the cellularity ex vivo of brain tumor specimens. Successful application of this imaging technique in the field of glioma surgery could provide faster throughput sampling from the surgical field and increase diagnostic yield.
Fluorescence lifetime imaging

Fluorescence lifetime imaging (FLIM) generates contrast by measuring the length of time a fluorophore remains in its excited state. By adding the extra dimension of time, FLIM provides a more specific means of identifying tumor cells based on the inherent amount of time endogenous fluorophores continue to emit photons. An alternative to measuring fluorescence intensity alone, FLIM capitalizes on the finite high-energy state by essentially averaging its lifetime in the tissues following excitation from the ground state, demonstrating relatively longer lifetime values in tumor cells.[62] [Figure 3e and f]. This is particularly useful in autofluorescence since many endogenous molecules possess overlapping spectra of fluorescence, thereby increasing the parameters needed for optimal contrast. Recently, FLIM has been used for identifying oral precancerous lesions, skin lesions due to UVB radiation, and in determining the function of retinal pigment cells.[47,66,97] The first use of FLIM for brain tumor image-guided surgery in humans showed that GBM had a significantly more irregular excitation lifetime distribution as compared to normal cortex.[102] While these results indicate the feasibility of such a time-dependent imaging modality in brain tumor resection, there exist several limitations for potential larger-scale in vivo human surgical trials, such as the lengthy time needed to acquire images, the complicated equipment that must be maneuvered and operated, and the cost of the procedure.[47]

Two-photon and second harmonic generation

While the depth of imaging limits confocal and other types of linear fluorescence microscopy, two-photon microscopy has the distinct advantage of visualizing deep cellular structures [Figure 3g and h]. With concentrated photons pulsed from a femtosecond laser, two photons, each containing approximately half of the energy required to linearly excite the fluorophore, simultaneously interact with the same molecule.[114] Intensity of these pulsed photons decreases circumferentially around the area of focus, resulting in a small volume of fluorescence emitting directly from the focal point. This allows spatial restriction of fluorescence to the focal point with reduced scattering. Additionally, two-photon microscopy uses infrared and far-red excitation wavelengths, which provide greater tissue penetration compared with the more superficial use of visible wavelength light.[114]

Another nonlinear imaging modality is second harmonic generation (SHG). This modality generates contrast based on the degree of photon scattering when incident photons interact with heterogeneous tissue. Although two photons with half the expected energy also interact at the focal point in SHG, they only do so after they are scattered by the same molecule rather than absorbed. This “emitting” photon with essentially twice the energy is then detected creating an image with potential for in vivo use during surgery. However, further development of such technologies is needed, including improvement of image processing time and spatial recognition issues.[83]

CONCLUSION

Currently, the frozen section provides intraoperative histopathological analysis of brain tumors. Though useful, this process is time consuming and requires the cutting, freezing, and staining of several biopsies. Frozen section sample preparation frequently damages tissue, alters cellular architecture, and introduces tissue artifacts, all of which hinder a proper diagnosis.[110,113] Application of exogenous fluorophores approved for neurosurgery and miniaturization of confocal microscopy may overcome the limitations of traditional intraoperative histopathology. Future operating rooms may see the routine use of fluorescence confocal endoscopes and clinically approved contrast agents, as well as the development of training programs to educate pathologists and surgeons to effectively apply this technology. The translation of this technology for neurosurgical use could improve patient care by providing the neurosurgeon with real-time histopathological information from more regions of interest than are possible with frozen section analysis.

Fluorophores approved for clinical neurosurgery include 5-ALA, fluorescein, and ICG. 5-ALA provides tumor-specific labeling for macroscopic detection of brain tumors. 5-ALA has been clinically used with fluorescence surgical microscopes to enhance the extent of resection of low-grade gliomas, which have limited uptake of 5-ALA. Confocal endomicroscopy with 5-ALA may provide visualization of tumors at the cellular level, which will allow real-time differentiation between normal and malignant cells. Fluorescein and ICG, provide less tumor-specific staining, and have demonstrated the ability to macroscopically demarcate brain tumors due to the permeability of tumor vasculature. Both of these fluorophores have been studied with fluorescence endomicroscopy. Using this technology, fluorescein and ICG generate contrast and allow the in vivo histopathologic visualization of tissue cellularity and intercellular architecture. Compared with fluorescein, ICG provides greater depth of imaging due to its excitation-emission wavelengths in the infrared spectra, thereby minimizing scatter and autofluorescence.

Optiscan and Cellvizio each produce commercially available confocal endomicroscopy systems for clinical use. Both utilize different imaging software and technical specifications to accomplish the goal of providing in vivo cellular detail during surgical procedures. These systems have shown utility in studies related to epithelial-derived structures. However, more rigorous studies such as human clinical trials with clinically approved fluorophores are required to establish the efficacy of these systems in...
the neurosurgical field. Novel imaging modalities are being studied in different organ systems. Investigation of future-imaging technologies in the field of neurosurgery is important for identifying new imaging tools available during neurosurgical procedures.

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