Technical Note: First Use of Endonasal Confocal Laser Endomicroscopy – Feasibility and Proof of Concept

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Abstract

Introduction Probe-based confocal laser endomicroscopy (p-CLE) is a method for real-time in vivo visualization of mucosal changes on a cellular level. Due to the size of the endoscopes, it was mainly used in the gastrointestinal tract so far. First investigations on head and neck carcinoma described the oropharyngeal application. The further miniaturization of the laser probe now allows endonasal application and, thus, first experiences with the investigation of endonasal neoplasms.

Objectives The aim of the present investigation is to elucidate, based on the morphological criteria validated in the oropharynx, whether these criteria be transferred in a similar way to the endonasal mucosa.

Methods We conducted p-CLE (Cellvizio, Paris, France) with intravenous fluorescein staining in endoscopic sinus surgery in a patient with sinonasal inverted papilloma and a histologically confirmed squamous cell carcinoma. We compared the cellular visualization of pathological changes with those of healthy mucosa in the same specimen, and also with our former findings in the oropharynx.

Results Endonasal p-CLE proved to be quite feasible in the surgical setting, and the transfer of malignancy criteria in analogy to histological examination could be optically retraced. Furthermore, additional criteria for tissue dignity assessment were obtained.

Conclusion Our results suggest that endonasal application of p-CLE represents a valuable extension of the diagnostic repertoire available to date by an additional real-time analysis of the nasal mucosa. This is of particular value in surgically challenging anatomical areas such as the paranasal sinuses.

Further investigation and validation will be necessary.
considered a second tumor. For most authors, early recurrence implicates incomplete resection. Clear differentiation of healthy and neoplastic mucosa can rarely be made by endoscopic inspection alone, especially regarding the margins of a lesion.

Real-time information on cellular and subcellular details, especially of the resection margins at the time of the endoscopic examination, could facilitate immediate definitive surgical treatment and, additionally, yield more information on the surrounding tissue.

Confocal laser endomicroscopy (CLE) allows the assessment of changes in the vascular architecture, connective tissue and cellular components of the mucosa in vivo. This technology was established in the gastrointestinal (GI) tract by researchers of our group using a flexible confocal endomicroscope in order to detect different inflammatory and neoplastic abnormalities in the human colon, esophagus, and stomach.

In 2012, our group described the first oropharyngeal application of this technique and validated malignancy criteria comparing with the gold standard of histological examination. We present the first endonasal application of in vivo fluorescein-enhanced confocal laser endomicroscopy in IP and SCC to further investigate the use of this technology in the field of functional endoscopic endonasal surgery and tumor resection.

**Material and Methods**

The present study was approved by the local ethics committee of Hessen, Germany (No.: FF 146/2017); written informed consent was obtained from each patient before examination. We performed computer-navigated endoscopic endonasal removal of a right-sided sinonasal IP, as well as of a left sided endonasal squamous cell carcinoma (see Fig. 1). Probe-based CLE (pCLE) was conducted during these procedures, and fluorescein was used as contrast agent. We followed an algorithm comparing healthy mucosa of the opposite side with the respective lesion in both cases. After detailed documentation, the surgery was completed.

In CLE, a low-power laser is focused to a single point in a microscopic field of view, and the same lens is used as both condenser and objective folding the optical path, so the point of illumination coincides with the point of detection within the specimen. Light emanating from that point is focused through a pinhole to a detector, and light emanating from outside the illuminated spot is rejected from detection.

Illumination and detection systems are at the same focal plane and are termed “confocal”. All detected signals from the illuminated spot are captured and measured. The intensity of emitted light is translated into a gray-scale image, which represents microscopic tissue structures.

The evolution of this technology over the past several years has led to the integration of a miniaturized laser scanner into the tip of a conventional flexible video endoscope that can visualize the mucosal details at subcellular resolution during endoscopy.

**Results**

Confocal imaging provided instant real-time microscopic imaging during the ongoing surgical procedure. There

![Fig. 1](image-url)  
*Fig. 1* Intraoperative setting from left to right: AIDA system for functional endoscopic sinus surgery, computed navigation (Medtronic) and confocal laser endomicroscopy (Cellvizio).
were no adverse side effects from intravenous fluorescein application after our investigation.

Imaging of the mucosal lesion in the nose as well as in the paranasal sinuses was feasible, and images of good quality could be acquired. Artifacts were most frequently caused by unstable positioning of the probe or by mucus or blood adherent to the probe.

Confocal imaging rarely added > 10 minutes to the total examination time. Healthy mucosa showed homogenous configuration of the superficial layers, with regular and defined cellular structure. Capillaries displayed regular and longitudinal configuration (see Fig. 2).

In both entities, the IP and the carcinoma, we visualized an inhomogeneous configuration of the superficial layers with less defined cellular structures and extended, irregular configuration of the capillaries. In squamous cell carcinoma, loss of cellular borders could be seen even more distinctively when compared with IP (see Fig. 3). Table 1 displays the morphologic properties for basic screening examination as used in this first report.

Resecting with clear margins was supported by verifying the absence of any of these changes when moving the probe away from the defined lesions. Handling of the probe simultaneously to the endoscope in the narrow conditions of the nose and of the paranasal sinuses was demanding; however, they were quite feasible in the hand of an experienced endoscopic-sinus-surgeon. Most areas, even within the maxillary sinus, were accessible, the anterior wall being the most challenging.

**Discussion**

This is, to our knowledge, the first report on the intraoperative application of pCLE in neoplasms of the nose and of the paranasal sinuses. Probe-based CLE provides real-time microscopic imaging during ongoing endoscopy and surgical treatment. This advanced technique has been successfully established in many fields, including gastroenterology, urology, dermatology and otolaryngology.

The probe-based system (pCLE, CeliVizio Endomicroscopy System, Mauna Kea Technologies, Paris, France) consists of a 1.6 mm flexible miniprobe with a field of view of 240µm diameter, a confocal depth of 55–65 µm and a resolution of 1 µm.

These probes can be used solo or be fitted through the working channel of most endoscopes for clinical use. Confocal laser endomicroscopy is only possible using fluorescence contrast agents. Intravenously applied fluorescein sodium (5 ml, 10%) distributes throughout the entire

![Fig. 2](image1.png)

Left: capillaries with regular and longitudinal configuration and erythrocytes (arrow). Right: extended, “cork-screw-like”-configured capillaries (arrow).

![Fig. 3](image2.png)

Left: healthy mucosa: homogenous configuration of the superficial mucosal layer with regular and defined cellular structure and cross-section of two capillaries. Middle: inverted papilloma: superficial epithelial layer with irregular cellular configuration but preserved cell margins. Right: squamous cell carcinoma: inhomogeneous cell structure and extended capillaries.
mucosa with a strong contrast within the connective tissue and the capillary network. Fluorescein binds to serum albumin and the remaining unbound dye molecules pass across systemic capillaries and enter the tissue, highlighting the extracellular matrix. Confocal laser endomicroscopy is possible within seconds after injection. Intravenous fluorescein is a nontoxic agent that is commonly used in ophthalmology with a long and substantial history of clinical use.

The most common side effects related to the IV administration, which are vomiting and nausea, are transient and minor. More severe side effects, such as vasovagal response, cardiac or respiratory response are extremely rare. In a pilot study, our group investigated the feasibility of the method using the flexible endoscope in different regions of the human oral cavity and of the oropharynx after intravenous application of fluorescein. We were able to show for the first time, to our knowledge, that confocal laser endomicroscopy is suitable for the evaluation of epithelial linings of the buccal mucosa, tongue, and floor of the mouth.

Signs of malignancy (altered tissue architecture and increased density of blood vessels with irregular, elongated, and enlarged appearance) could be identified in vivo.

In 2012, our group elaborated a systematic intraoperative evaluation of confocal laser endomicroscopy in the oropharynx. Characteristics of healthy mucosa such as homogenous configuration of the superficial mucosal layer, clearly defined cellular structure, and longitudinal configuration of capillaries could be reproducibly demonstrated in different locations of the head and neck. In SCC, malignancy criteria such as inhomogenous mucosal configuration, blurry cellular borders, as well as extended and irregular configuration of capillaries could be applied. Based on the current investigation, these criteria appear to be applicable to our endonasal findings.

In 2014, Nathan et al. showed an overall sensitivity for the diagnosis of dysplasia versus nondysplasia of 80.0% using pCLE in the oral cavity; while the dorsal surface of the tongue was not well visualized, the remaining nonkeratinized subsites, including the buccal mucosa, the floor of mouth and the ventral tongue, were well visualized. Most recently, Wu et al. showed the application of pCLE in nasopharyngeal carcinoma. Goncalvez et al. demonstrated the application in vocal cord malignancies, which demonstrated similar characteristics.

The further miniaturization of the probe now allowed nasal passage and examination of the paranasal sinuses. The application of fluorescein showed to be quite feasible. Tissue was adequately assessable ~ 30 seconds after intravenous application. However, the handling of the flexible laser probe is a challenge for the surgeon in the variable conditions of the paranasal sinuses and is subject to an individual learning curve. The necessary mucosal contact pressure of the probe, as well as the contact angle for an adequate visualization, are variable regarding the irregular surfaces of the paranasal sinuses. The guidance of the probe through an instrument such as a straight or angled forceps proved practicable.

The limitations of visualization were particularly seen in increased bleeding, which makes handling and application in well vascularized tumors more demanding.

**Conclusion**

The novel application of using p-CLE with miniaturized flexible laser probes appears to be suitable for noninvasive real-time in vivo diagnostics of endonasal lesions. Cellular changes and the extent of resection lines of such lesions can be visualized. Further investigations have to evaluate and validate the significance of this technology in endonasal surgery for invasive malignancy, especially regarding critical resection margins.

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**Conflict of Interests**

The authors have no conflict of interests to declare.

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