Functional Capillary Density for in Vivo Estimation of Intestinal Perfusion using Real-Time Confocal Endomicroscopy

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Abstract

Background and aim: Confocal Laser Endomicroscopy (CLE) has been successfully used to appreciate microcirculation changes of the digestive mucosa. Our aim was to evaluate CLE scanning complemented by functional capillary density area (FCD-A) estimation to define the micro-vessel status in a reiterate, long-lasting porcine model of bowel ischemia.

Materials and methods: A laparotomy was performed in 4 pigs, and a segmental (3–4 cm) ischemia of the sigmoid colon was induced with vascular clamps. Ischemic and perfused regions were clinically defined. After an injection of 5 ml of sodium fluorescein 10% (Fluocyn, SERB, Paris, France), the Cellvizio™ confocal probe (Mauna Kea Technologies, Paris, France) was directly applied onto the mucosa’s surface through a full-thickness enterotomy. Both ischemic area (IA) and control region-perfused area (PA) – were scanned and video sequences were recorded.

Results: Confocal evaluation of the ischemic area revealed a different aspect of the mucosal tissue when compared to the normal perfused area. Statistically, FCD-A at the perfused area was significantly higher when compared to the ischemic area, irrespective of the time point. After 1 hour, FCD-A was (0.189 ± 0.094 vs. 0.365 ± 0.030; p=0.0001), after 2 hours (0.252 ± 0.056 vs. 0.389 ± 0.024; p<0.0001), after 3 hours (0.252 ± 0.050 vs. 0.353 ± 0.030; p=0.0001) and after 4 hours (0.262 ± 0.044 vs. 0.358 ± 0.019; p<0.0001), at ischemic and perfused areas respectively.

Conclusions: Confocal imaging allows real-time discrimination between perfused and ischemic areas of the bowel using morphological clues, while the functional capillary density area adds a quantitative measurement.

Keywords: Confocal laser endomicroscopy; Functional capillary density area; Cellvizio™ system; Quantitative assessment of bowel perfusion

Introduction

Confocal laser endomicroscopy (CLE) is a high-resolution imaging modality allowing for real-time in vivo virtual biopsies with in vivo magnification up to 1000X [1] and spatial lateral resolution ranging from 1 to 3.5 μm [2].

CLE systems include through-the-scope probes (probe-based CLE, or pCLE), e.g. the Cellvizio™ system (Mauna Kea Technologies, Paris, France) (Figure 1) or endoscope-based CLE (eCLE), e.g. the Pentax Confocal Endomicroscope (EC-3870CIFK, Pentax, Tokyo, Japan) [2]. The main advantages of eCLE when compared to pCLE are the larger visual field (475 × 475 μm vs. 240-325 × 240-325), and the adjustable imaging plane depth [1]. These features improve the performance of eCLE in terms of accuracy [1]. However, pCLE is more versatile since various sized probes can be inserted through the operative channel of standard endoscopes, as well as through biopsy needles (needle-based CLE). Additionally, pCLE allows to perform a virtual biopsy under visual control [2].

Image acquisition in CLE requires a fluorophore, which can be administered using a systemic injection (mainly sodic fluorescein) or through a topic application (acrilavlin or violet cresyl). Those “contrast” agents may add some dynamic and functional data to morphological information, making the confocal method more than a mere digital version of histopathology.

Current clinical applications for pCLE in GI endoscopy include post-resection follow-up of colonic lesions [3], diagnosis of indeterminate biliary strictures [4], Barrett’s esophagus surveillance, and post-treatment pathologic assessment [5,6]. Evolving applications include inflammatory bowel disease [7], gastric diseases [8], differentiation of colorectal polyps [9], and pancreatic cysts [10]. In the experimental setting, CLE can provide in vivo analysis of microcirculation [11,12]. Recently, CLE was used to appreciate microcirculation changes of the digestive mucosa during septic shock in the experimental and clinical setting [13]. Our group demonstrated the ability of the Cellvizio™ system to accurately discriminate between perfused and ischemic areas in a 1-hour model of bowel ischemia [14]. The aim of this experimental study was to evaluate confocal scanning complemented by functional capillary density area (FCD-A) estimation to define the micro-vessel status in a reiterate, long-lasting porcine model of bowel ischemia.

Materials and Methods

Animals

Four female pigs (Sus scrofa domesticus, ssp. Large White), mean

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weight of 31.025 ± 3.34 kg, and were used in this non-survival study. The protocol (No. 38.2012.01.039) was approved by the local Ethical Committee on Animal Experimentation. Animals were managed according to ARRIVE guidelines [15] and in accordance with French laws for animal use and care and according to the directives of the European Community Council (2010/63/EU). Twenty-four hours before surgery, pigs were fasted with free access to water. Intramuscular injection of ketamine (20 mg/Kg) and azaperone (2 mg/Kg) (Stressnil, Janssen-Cilag, Belgium) were administered 10 minutes before surgery. Induction was achieved using intravenous propofol (3 mg/Kg) combined with rocuronium (0.8 mg/Kg). Anesthesia was maintained with 2% isoflurane. At the end of the procedure, animals were sacrificed with an intravenous injection of a lethal dose of potassium chloride.

Surgical procedure and Confocal Endomicroscopy assessment

A median laparotomy was performed in all pigs, and segmental (3–4 cm) ischemia of the sigmoid colon was induced by occluding terminal arterial branches with vascular clamps. Ischemic regions were clinically defined through observation and taking into account the vascular anatomy. After an injection of 5mL of sodium fluorescein (Fluocyn, SERB, Paris, France), the Cellvizio ™ pCLE Gastroflex probe-based Confocal Laser Endomicroscopy (pCLE) was applied directly onto the sigmoid mucosa through full-thickness enterotomy. UHD was used to perform a scan of the sigmoid mucosa (Figure 1).

The perfused area was invariably normal at standard histology (score 0) assessed with a standard pathology (1.71 ± 0.49 vs. 1 ± 0; p=0.0082). The score scale was as follows: 0= normal mucosa; 1= partial epithelial edema and necrosis; 2=diffuse swelling and necrosis of epithelium; 3=necrosis with submucosal neutrophil infiltration; 4=widespread necrosis and massive neutrophil infiltration and hemorrhage.

Statistics

Statistics were performed using the GraphPad Prism® software. A Student’s t-test was performed to calculate differences between ischemic and perfused areas. Differences were considered statistically significant for p values<0.01.

Results

Confocal evaluation of the ischemic area revealed a different aspect of the mucosal tissue when compared to the normal perfused area. CLE identifies the ischemic area with blurred images and swelling with an increased demarcation of the cryptal border, due to increased basolateral permeability. The presence of “target cells” is also characteristic, defined by the presence of a hyper-fluorescent crypt centre and by the distortion of the enterocyte silhouette. These typical features cannot be found in the normally perfused area (Figure 4 and 5).

Statistically, FCD-A at the perfused area was significantly higher when compared to the ischemic area, irrespective of the time point. After 1 hour, FCD-A was (0.189 ± 0.094 vs. 0.365 ± 0.030; p=0.0001), after 2 hours (0.252 ± 0.056 vs. 0.389 ± 0.024; p<0.0001), after 3 hours (0.252 ± 0.050 vs. 0.353 ± 0.030; p=0.0001), and after 4 hours (0.262 ± 0.044 vs. 0.358 ± 0.019; p<0.0001), at ischemic and perfused areas respectively.

Overall, the ischemia score was low, ranging from 0 to 2. The CLE-based score of the ischemic area was significantly higher than the one assessed with a standard pathology (1.71 ± 0.49 vs. 1 ± 0; p=0.0082). The perfused area was invariably normal at standard histology (score 0) and at confocal evaluation, except in one case.

Discussion

CLE is an emerging real-time diagnostic tool, which can

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**Figure 1:** Probe-based Confocal Laser Endomicroscopy (pCLE). A) Cellvizio ™ monitor showing perfused sigmoid mucosa. B) The Cellvizio ™ pCLE Gastroflex UHD was applied directly onto the sigmoid mucosa through full-thickness enterotomies.
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complement an endoscopic assessment by performing a digital biopsy. It is being progressively implemented in the management of neoplastic or inflammatory diseases including gastrointestinal, urinary or respiratory tract [2] diseases. In addition, CLE can provide information about the microcirculatory status of tissues [12].

More particularly, CLE might provide a quantitative real-time assessment of bowel perfusion, which could have a relevant impact on various clinical situations, including the management of intensive care unit patients or the intraoperative evaluation of stoma or bowel stump perfusion.

Yasumura et al. used a charge-coupled microscopic device to analyze perfusion from the serosal layer in a bowel ischemia model. Authors could calculate a ratio of circulating blood cells to the functional bowel vascular bed as a predictive index of intestinal survival [17]. This type of accurate quantification of intestinal perfusion, when it is not time-consuming or overwhelmingly expensive, might represent a paradigmatic shift towards micro-image-guided therapies, allowing for a Doppler-like real-time examination at a microscopic scale.

Schmidt et al. were able to describe sepsis-related changes in mucosal microcirculation in a porcine model of septic shock and also
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in patients using the Cellvizio system and a computer-based evaluation of functional capillary density area (FCD-A). In the future, such detectable changes could potentially help to adjust fluid resuscitation regimens, given the complexity of fluid management in critical patients presenting a shock of abdominal origin [13].

In a porcine laparoscopic model of bowel ischemia limited to 1 hour, we could observe that a real-time confocal morphological evaluation provides early and specific clues to identify the vascular status. CLE was more specifically used to confirm and validate the accuracy of a surgical navigation system to detect bowel perfusion, based on the dynamics of the fluorescent signal [18-20].

The aim was to perform assessments over time and to complement the morphological analysis with FCD-A computation. For that purpose, we designed the current open surgery and long-lasting model of sigmoid ischemia (up to 4 hours).

The vessel detection tool, which is included in the software provided by Mauna Kea Technologies (IC-Viewer 3.8.5), may provide the FCD-A index, which can estimate vascular flow. The FCD-A index is based on fluorescein signal detection. In healthy mucosa, fluorescein is confined

Figure 4: Comparative images: standard histology versus confocal analysis. A blinded pathologist assigned an ischemia score to histological and confocal images. A) 20X Hematoxylin-eosin (HE), and B) confocal imaging score 0 (normal mucosa), C) 20X HE, and D) confocal imaging score 1 (submucosal congestion and swelling), E) 20X HE, and F) confocal imaging score 2 (diffuse swelling and necrosis of epithelium).

Figure 5: Confocal description of ischemic area. Black arrows indicate an enhanced cryptal border in the colonic mucosa. White arrowheads show the fluorescein leakage inside the cryptal lumen.
to the vessels’ lumen. Ischemic injury produces leaks of fluorescein by increasing vascular and basolateral intestinal permeability. This fluorescent leakage is found primarily in the cryptal lumen, as an early indirect sign of ischemia. This creates the aspect of “target cells” and subsequently forms a “palisade” aspect due to an increased spacing of the cryptal border. A prolonged ischemic injury leads to a pooling phenomenon in homogeneously distributed on the mucosal surface, creating visible artefacts during the software vessel detection process.

In order to optimize vessel detection, a post-processing analysis is required to subtract fluorescein leaking areas around and within the intestinal crypts.

When comparing the ischemia score applied by a blinded pathologist (VL) to both techniques, it was clear that the morphological confocal analysis tends to overestimate injuries when compared to standard histology. However, when it comes to a standard pathology, the perfused area was consistently deemed to be strictly normal and only mild damage was found at the ischemic areas, irrespective of ischemic time. Such findings were different from those made with CLE. Findings made with CLE demonstrated more discriminant signs of ischemic injury, which matched the FCD-A analysis. This study model was probably limited by the short ischemic segment, in which some reperfusion might have occurred, protecting the mucosa from further degeneration.

Conclusions

Confocal imaging allows for real-time discrimination between the bowel’s perfused and ischemic areas using morphological clues, while the functional capillary density area adds a quantitative measurement. This micro-image quantitative analysis might be helpful in clinical conditions requiring an accurate assessment of bowel perfusion, such as in the presence of a stoma or in the management of a shock of abdominal origin.

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