Introduction

Thanks to the widespread implementation of screening protocols, colorectal malignant lesions are an increasingly detected pathology, being reported in up to 12% of resected polyps [1]. Complete endoscopic resection of rectal lesions can be achieved with snare-polypectomy, endoscopic mucosal resection (EMR) or endoscopic submucosal dissection (ESD). However, subsequent histopathological examination of the resected specimen may reveal signs of incomplete resection raising the need for additional treatments (Table 1) [2].

Limited by post-polypectomy submucosal fibrosis, a rescue endoscopic therapy is often challenging, especially for rectal lesions localized close to the dentate line [3]. On the other hand, major rectal surgery often results in temporary or definitive stoma and in frequent post-surgical complications. The Full-Thickness Resection Device (FTRD®) System (Ovesco Endoscopy, Tübingen, Germany) is a novel system that, besides having other indications, appears to be promising for wall-thickness excision of intestinal T1 carcinoma following incomplete endoscopic resection. However, follow-up data on patients treated with this device are scarce, particularly for ERC.
motic dehiscence, may arise from rectal surgery even in tertiary referral centers [4, 5].

To overcome these limitations, a novel endoscopic therapeutic tool, called the Full-Thickness Resection Device (FTRD®) System (Ovesco Endoscopy, Tübingen, Germany) has recently been introduced. This endoscopic approach is technically successful for intestinal T1 carcinoma following incomplete resection, non-lifting adenoma or adenoma arising in difficult positions (i.e. the neck of the diverticulum, appendix, and dentate line). In selected cases, the FTRD proved effective for endoscopic treatment of small submucosal tumors and for diagnostic purposes [6–12]. The main features of this technique lie in generating a pseudopolyp involving the intestinal wall-thickness within an endoscopic cap (diameter 13 mm, length 23 mm) followed by an en bloc resection using a hot snare technique (monofilament, 14-mm polypectomy snare preloaded in the tip of the cap), after the deployment of a modified over-the-scope clip (diameter 14 mm) to seal the likely underlying transmural defect [6–12]. The FTRD can be quickly pre-loaded on the tip of an endoscope with a tip diameter of 11.2–13.2 mm. Initial experiences using the FTRD have shown promising results, especially in high risk patients and in lesions located in peculiar anatomic sites where standard endoscopic or surgical approaches would carry considerable risks and require aggressive strategies [6–12].

The aim of this pilot study was to assess for the first time the feasibility and long-term clinical impact of endoscopic treatment with the new FTRD in selected patients with T1-early rectal cancer (ERC) following an incomplete endoscopic mucosal excision.

Patients and methods

Consecutive patients diagnosed with T1-ERC following incomplete endoscopic resection or showing submucosal involvement (R1) at histological examination were evaluated for endoscopic full-thickness resection (EFTR) after a complete oncologic work-up. The inclusion criteria were: (i) residual rectal lesion or rectal scar < 20 mm; (ii) increased probability of disease recurrence as defined by the histopathological evidence of > 1 criteria according to Ueno et al. [2]; (iii) no lymphatic or metastatic disease at computed tomography (CT) scan and rectal endoscopic ultrasound (EUS); (iv) patients defined as “unfit for surgery” according to their underlying general condition, or who had refused the surgical option despite having received exhaustive information about the natural history of the disease and the presence of a valid surgical option. The exclusion criteria were patients < 18 years old, pregnancy, severe uncontrolled coagulopathy, and inability to provide an informed consent. These patients were offered and accepted to undergo the EFTR procedure, providing their informed consent. This study was carried out in accordance with the Declaration of Helsinki adopted in 1964 incorporating all later amendments.

Antibiotic prophylaxis with iv cefalosporin was administered to all patients. Before EFTR, the target lesion was identified under white-light endoscopic imaging and marked using the FTRD marking probe. EFTR was performed in a standard technique [6–12] using a preloaded FTRD including a tissue anchor, a modified 14-mm over-the-scope clip and a monofilament, 14-mm polypectomy snare connected to a standard electrosurgical device (VIO; ERBE Elektromedizin GmbH, Tübingen, Germany; ENDO CUT® Q®, effect 3, cutting duration/interval 4/1). All procedures were conducted by expert endoscopists with initial experience in the use of the FTRD (less than 10 procedures) using high-resolution endoscopes with a tip diameter of 11.2–13.2 mm and digitally recorded. After endoscopic treatment, patient outcomes were strictly monitored performing a tailored oncologic work-up including endoscopy, CT scan, and rectal EUS. Two expert gastrointestinal pathologists performed the histopathological analysis on the en bloc specimens obtained by EFTR.

Results

From June 2015 to February 2016, six patients were consecutively treated at the IRCCS Policlinico San Donato, San Donato Milanese, Italy (5 men, mean age 63 years, range 51–78 years). All patients had previously received a rectal EMR within the previous 1–3 months, thereby receiving the histopathological diagnosis of high risk malignant polyp [2] (Table 2). Each patient had been treated with EFTR at hospital admission and discharged home within the following 24 hours. The endoscopic full-thickness resection was technically feasible in all cases within 8–15 minutes (15–30 minutes including lesion detection, demarcation, and FTRD assembly). No immediate or late complications occurred. No patient reported any symptom related to the endoscopic procedure. High resolution endoscopic images displaying the original rectal lesions, the EFTR procedures, and the follow-up are reported in Video 1. A full-length demonstrative video was also recorded.

The histopathologic analysis performed on the en bloc resected specimen demonstrated a complete endoscopic resection in all patients, with the achievement of a full-thickness excision in four patients. The resected specimen included the submucosal layer with an estimated depth ≥ 1000 µm (1250–3050 µm) but not the muscularis propria in the two remaining patients (Table 2). By revising these two endoscopic charts, no evident technical issue affecting the endoscopic procedures has been identified.

Table 1: Histopathological criteria for high risk malignant polyps according to Ueno et al. [2].

| Low tumor differentiation grade (G3) |
| Haggitt’s levels (pedunculated polyps): 3–4 |
| Kikuchi’s levels (sessile polyps): sm3 |
| Width of submucosal invasion: ≥ 4000 µm |
| Depth of submucosal invasion: ≥ 2000 µm |
| Positive tumor budding |
| Distance from the excision margin < 1 mm |
| Presence of vascular invasion |

Table 2: Histopathological criteria for high risk malignant polyps according to Ueno et al. [2].

| Presence of vascular invasion |
| Positive tumor budding |
| Depth of submucosal invasion: ≥ 4000 µm |
| Distance from the excision margin < 1 mm |
| Width of submucosal invasion: ≥ 2000 µm |
| Presence of vascular invasion |

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During follow-up, all patients underwent an oncologic work-up with endoscopy, CT scan, and rectal EUS every 6 months (▶Table 2). All patients were in oncologic remission after a median follow-up of 12 months (range 12–18 months) without any radio- or chemotherapy. One patient died from cardiac failure at the 8th month of follow-up after showing no sign of disease recurrence at the 6-month oncological work-up.

**Discussion**

Our series confirms that EFTR with the newly introduced FTRD System is feasible and safe in T1-ERC. This study also shows that EFTR is a valid option for intestinal tumor excision following incomplete endoscopic resection in patients without evidence of metastatic or lymphatic disease when the standard surgical op-
tion is contraindicated, refused by the patient or the patient is at high risk.

The proper management of high risk malignant polyps relies on a multidisciplinary decision-making process currently based on the estimated risk of residual disease, as well as on several patient-specific features such as age, general global assessment, underlying morbidities, long-term prognosis, and the patient’s wishes [13]. The proper management of T1 ERC is remarkably influenced by the risk of lymph node micrometastasis. Therefore, following the endoscopic removal of colorectal malignant polyps, the specific risk of disease recurrence depends on the histopathological identification of standardized microscopic criteria and the nodal status [2]. The precise impact of each single microscopic criterion and their interrela-

tionship is still unclear. Nonetheless, when several adverse risk factors are present and the risk of residual disease is substantial (> 20 %), the decision-making process to undergo further surgical treatment is usually straightforward [13]. Exceptions include those patients whose comorbidities outweigh the risk of surgery and those refusing major rectal surgery to avoid the risk of definitive stoma. Conversely, routine follow-up without further treatment is the best option after endoscopic removal of very low risk (< 3 %) malignant polyps [13].

Notably, margin positivity on its own does not appear to be an independent risk factor for lymph node metastasis, with the risk of nodal metastasis being similar in patients with and without margin involvement [14]. Previous studies have clearly shown that the risk of disease recurrence is often overestimated by histopathological assessment of involved margins within the resected specimens [15, 16]. Indeed, radical surgery for incomplete endoscopic resection of early colon-rectal cancers provides tumor-free specimens in up to 76 % of cases, thereby failing to improve the 2-year survival rates [17] while imposing significant risks of immediate morbidity and long-term complications on the patient.

Consequently, we performed a full thickness endoscopic resection following supposedly incomplete (R1) endoscopic excision with endoscopic mucosal resection of T1-ERC in those patients either refusing or unfit for the standard surgical options. In the present series, all but one of the specimens obtained by EFTR showed fibrotic submucosal tissue with no dysplastic residue. Within these cases, no residual or recurrent dysplastic tissue was observed when performing EFTR. In the remaining patient with incomplete endoscopic excision of T1-ERC, cancer recurrence was already evident at endoscopy.

As compared to established surgical curative treatments for rectal cancer (i.e. lower anterior resection with total mesorectal excision and abdominoperineal resection), less invasive trans-anal full-thickness excision techniques (e.g., conventional trans-anal excision, trans-anal endoscopic microsurgery (TEM), or trans-anal minimally invasive surgery (TAMIS)) have comparable 5- and 10-year survival rates [18, 19] but clear advantages in limiting either surgery-related mortality or morbidity, and the need for a permanent stoma [19]. However, by reducing both the resected specimens and the mesorectal lymph node assessment, these treatments hamper the exact disease staging, thereby implying an increased risk of local recurrence and missed micrometastasis [20]. In addition, following either TEM or TAMIS, major complications have been reported in 1.5 – 7 % of patients and conversion to laparotomy with or without total mesorectal excision or temporary stoma is sometimes necessary [21 – 23].

Within this context, our initial results indicate that the endoscopic approach with the FTRD is a valid alternative to trans-anal full-thickness excision techniques for non-pedunculated T1-ERC smaller than 20 mm (i.e. estimated maximal size referring to the luminal diameter of the rectal naïve lesion or to the scar following previous EMR or polypectomy), resulting in low comorbidity, fast operating time, and anesthesiology-free procedures. These results are consistent with other recent series showing positive outcomes when the FTRD has been used for
the treatment of adenomatous and early colonic cancer, including both right- and left-sided lesions [6–12]. According to these reports, the FTRD is technically feasible for lower gastrointestinal mucosal lesions up to 30 mm. However, we believe that lesions ≤ 20 mm represent an ideal target for stiff tissues (e.g., incomplete resections with tissue fibrotic retraction, nonlifting naïve lesions) and angulated positions (e.g., distal rectum, recto-sigmoid junction, colonic flexure).

When compared to ESD or EMR, EFTR has the potential to allow for en bloc radical endoscopic excision of T1-ERC involving all submucosal layers, with reduced risk of bleeding, perforation, and post-polypectomy syndrome, which appear comparable even in referral ESD centers [24, 25]. Furthermore, ESD is technically difficult, especially in fibrotic tissue due to previous excisions, it is time-consuming, and requires a prolonged learning curve for inexperienced endoscopists [24]. In fact, ESD outcomes from Western studies are substantially worse compared with Eastern studies, thereby limiting generalizability of the results [20]. However, ESD represents the only reasonable endoscopic approach for superficial (sm1) T1-ERC with a diameter exceeding 30 mm (i.e., large non-pedunculated colorectal polyps), since the use of the FTRD would not be feasible for technical reasons (cap diameter/length 13/23 mm) [6–8], while EMR often leads to piecemeal resection, challenging histopathological assessment of R0 resection, and increased risk of incomplete excision [24].

Our positive experience with EFTR has some inherent limitations. Despite being prospectively designed, the main drawback of this pilot study is the relatively short follow-up, which includes patients with a full negative oncologic work-up at 6 to 18 months since the time of EFTR. Secondly, six patients are not included patients with a full negative oncologic work-up at 6 to 18 months since the time of EFTR. Secondly, six patients are not included patients with a full negative oncologic work-up at 6 to 18 months since the time of EFTR. Lastly, the study was performed at two referral centers, and thus the data might not be representative of the general population. Finally, we identified a number of potential complications, such as the entrapment of other pelvic structures close to the rectal wall; to date, such complications have never been documented in the literature [6–12]. Finally, a complete full-thickness rectal excision was not feasible in one-third of our patients, where the deepest submucosal layer but not the muscularis propria was included in the resected specimens. A post-hoc revision of these endoscopic procedures was not able to identify any technical feature clearly affecting the successful full-thickness resections in those two patients. Post-polypectomy fibrotic changes following previous EMR can increase the stiffness of the rectal wall and thus impair the endoscopic suction of the deepest layers within the snare housing. In any case, the deepest layer of the suctioned tissue remains within the over-the-scope clasp, being thereby bound to ischemic injuries and fibrotic remodeling, thus decreasing the risk of local recurrence.

In conclusion, this study provides initial evidence in favor of EFTR with the newly introduced FTRD System for rectal malignant polyps featuring a medium risk of disease recurrence after endoscopic mucosal resection in patients either unfit for surgery or refusing the standard surgical approach.

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Competing interests

None

References


Soriani Paola et al. Endoscopic full-thickness resection... Endoscopy International Open 2017; 05: E1081–E1086
Over-the-scope clip-assisted full thickness resection after incomplete resection of rectal adenocarcinoma. Endoscopy 2016; 48: E59 – E60


Lymphatic vessels distribution in the mucosa and submucosa and potential implications for T1 colorectal tumors. Dis Colon Rectum 2011; 54: 35 – 40


Local resection compared with radical resection in the treatment of T1N0M0 rectal adenocarcinoma: a systematic review and meta-analysis. Dis Colon Rectum 2015; 58: 122 – 140

Management of colorectal T1 carcinoma treated by endoscopic resection from the Western perspective. Dig Endosc 2016; 28: 330 – 341


Transanal endoscopic microsurgery: indications and results after 100 cases. Colorectal Dis 2004; 6: 350 – 355
