A new device to expedite endoscopic submucosal dissection procedures: a randomized animal study of efficacy and safety (with videos)

Background and study aims: Endoscopic submucosal dissection (ESD) is a recognized method for the curative treatment of superficial neoplasia, but its use is limited by lengthy procedures and the lack of versatility of existing knives. We developed a prototype ESD device with the ability to work as a needle, hook, or scythe. This new device was compared to regular ESD knives in a randomized animal study.

Patients and methods: Eight pigs underwent two gastric ESD procedures each, similar in size and difficulty, one with a regular ESD device and the other with the new device. The order and location of each ESD, as well as the performing operator, were randomized. Primary judgment criterion was safety of procedures. Overall and submucosal dissection procedure times were measured. Time-to-surface ratios were measured and estimated for ESDs larger than those performed. Histopathology of the resected tissue and remaining stomach was done after each experiment.

Results: No complications were observed throughout the study and all resections were completed en-bloc and uneventfully. The submucosal extension of resections was similar with both the standard and the new devices. A comparison of time-consumption between groups did not show statistically significant differences, but a dramatic reduction of procedure duration was observed in some procedures with the new device; based on observed data, a potential timesaving of up to 66% was anticipated, with a relatively short learning curve.

Conclusions: This new versatile device proved to be as safe as regular ESD knives, and seems likely to help reduce the duration of the procedure.

Introduction

Endoscopic Submucosal Dissection (ESD) is a recently introduced technique to achieve en-bloc resections of digestive epithelial or subepithelial tumors, sparing the muscularis propria (MP). ESD was described and developed in the early 2000s by Japanese authors and has been widely accepted in Far Eastern countries as a carcinologic method of treating superficial tumors with good prognostic features [1-3]. A quick Medline search yields more than 1200 publications up to the end of 2012, including thousands of patients and some randomized studies, showing that ESD is safe, reliable, and effective. However, despite its clear advantages – a minimally invasive approach allowing en-bloc resections with clean deep and lateral margins for histopathologic analysis – the spread of ESD in western countries has been much slower and remains restricted to a small number of skilled endoscopists in a few referral centers [4,5]. Such a disappointing outcome more than 10 years after the introduction of the technique is probably due to some well-known shortcomings of ESD, such as a long and demanding training curve, a very high time consumption relative to the amount of resected tissue, as well as the lack of versatility of many ESD devices [4,6]. Because the development of ESD is desirable, given the increasing incidence of early/superficial carcinomas resulting from widespread cancer screening and improved diagnostic techniques, it seems necessary to make the procedure simultaneously technically easier, sufficiently versatile to be used in the most difficult anatomical conditions, less time-consuming, and more affordable [7]. Our aim was to develop a new device that would fulfill those objectives and this is the first assessment of such a device in a porcine model.
Patients and methods

ESD device

The device was designed in partnership with Medwork Medical Products and Services GmbH (Höchstadt an der Aisch, Germany) as a monopolar electrosurgical knife to be used by deploying the active wire out of the catheter sheath via a handle equipped with an electric connection port, as with most existing ESD knives. The major difference with this device is that the instrument can be used as a needle when only the very tip of the wire is deployed (2 mm in the current prototype) and also like a “scythe” when the wire is fully deployed. The needle position is typically suited to a peripheral incision, as with any other ESD device, whereas the “scythe” position offers a much longer electrical contact surface, which can make progression during the submucosal phase of the ESD procedure much easier and faster. In the “scythe” position, the 0.4-mm wire can be oriented easily over 360° by rotating the handle, so that the cutting plane remains always strictly parallel to the submucosal plane. When deployed halfway, the wire presents in the form of a hook, and can be used as such, to grab and cut fibers in the submucosa. Fig. 1 shows these different configurations.

Furthermore, the prototype used was designed with additional features, including a slightly enlarged tip that improves the safety of the incision when the catheter is placed bluntly on the muscular plane, and a flushing port that makes use of the same lumen as the cutting wire for removal of coagulation debris from the wire, cleaning of the operating field, and inflation of the submucosa during ESD. Pressure at the catheter outlet, however, is not sufficient to lift the mucosa during the early phase of ESD (i.e., before mucosal incision) unlike the high pressure achieved with devices such as the Erbejet™-Hybrid-knife™ system [9]. Therefore, an injection needle is mandatory for performing ESD with this instrument and the flushing port can be used for the same purpose as a Flush-knife™. Fig. 2 shows the prototype.

Video 1 show how to maneuver it.

Animals

Eight 35-kg male or female pigs from the same farm, were used for the study. The pigs were housed at our facility during the 48 hours before the procedure. Endoscopies were performed under general anesthesia. All animals were prepared for anesthesia with a 12-hour diet, and received intramuscular injection of 10 mg/kg ketamine and 2 mg/kg azaperone (Stresnil®, Janssen Cilag, Titusville, New Jersey) 30 minutes before induction. After induction with 8 mg/kg of intravenous (IV) propofol (1% Diprivan®, AstraZeneca, London, UK) and endotracheal intubation, ventilation was performed with a Siemens 900 C ventilator (Siemens, Solna, Sweden). Anesthesia was maintained with inhaled 1% to 2% isoflurane (Abbott, Abbott Park, IL). All animals received IV infusion of 10 mg/kg/h crystalloid solution. The experimental protocol received approval from the scientific committee of the Surgical School of Paris (École de Chirurgie de l’Assistance Publique – Hôpitaux de Paris, Paris, France). Experiments were performed according to the standard guidelines of the French Ministry of Agriculture, which regulates animal research in France.

Study design

The study was designed as a small-sample randomized trial whose main aim was to compare the efficacy, ease of use, and safety of the new device vs a high-quality conventional ESD knife. The conventional or “regular” ESD knife used was the 2-mm straight (I-type) Flush-Knife™ (Fujifilm Corporation). The two endoscopists performing ESDs (FP, SL) each had experience with 100 or more ESDs in patients at the time the study started and more than 1 year of experience with the Flush-knife™. Their only experience with the new device was in use of the first prototype devices on a few cases in live pigs a few weeks before randomization of animals for this study.

ESD description and group allocation

Each animal in the study underwent two gastric ESDs in the same locations, one in the anterior wall approximately 4 cm below the cardia and close to the lesser curvature, and the other one next to the first, also on the anterior wall but closer to the greater curvature and a bit more distant in the stomach. All procedures were performed in the forward position and none in retroflexion. These locations were chosen for their ease of reproduction from one animal to the next one. All ESDs were approximately the same size, between 3.5 and 4 cm “Lesion” diameter was measured with the help of an adult biopsy forceps with the jaws open, which was moved four times along a straight line in both horizontal and vertical axes. After marking four quadrants with a biopsy, a circular marking was done around these four points by coagulation dots. This method allowed a fairly reproducible “lesion” size.

In all animals, one ESD was performed with a “regular” ESD device (Flush-knife™), and the other one was performed with the new device. The order in which both ESDs were done was randomized. All procedures were performed by the same senior and ESD-trained endoscopists (FP and SL); in each animal, one ESD was done by FP and the other one by SL; at the end of each series, the same number of ESDs had been done by both endoscopists with each of the devices to be compared (regular and new), and the same number of ESDs had been done with each of the devices in both gastric locations, so as to eliminate both operator and anatomical biases.

The ESD technique was conventional, with peripheral marking using the device (regular or new), submucosal lifting with indigo-stained 9% saline, peripheral incision with a 1- to 2-mm margin beyond markings, and submucosal dissection with the help of a transparent distal attachment (model D-201-10704, Olympus, Germany) and an ERBE ICC 350 electrosurgical unit (ERBE, Tubingen, Germany). Electrical settings were Endocut and Forced Coagulation 40 W.

The animal was sacrificed with 100 mg/kg IV injection of 7 mg of pentobarbital (Dolethal®, Vétoquinol, Paris, France) after completion of both ESDs. A gastrectomy was done and the stomach was opened to check for any accidental perforation unrecognized during procedures, lesions of the serosal face of the stomach, and accurate measurement of the mucosal defects. After macroscopic examination and measurement with a calliper, specimens were fixed on cork for histopathology. All procedures were achieved with a Storz gastroscope (Karl Storz GmbH, Tuttingen, Germany) and an ERBE ICC 350 electrosurgical unit (ERBE, Tubingen, Germany). Electrical settings were Endocut and Forced Coagulation 40 W.
data were compared using the paired Student’s t-test for small samples.

**Judgment criteria**

Our primary judgment criterion was safety, as assessed by the rate of complications (essentially perforations) that occurred during the procedures or observed at necropsy. Another important outcome measure was efficacy, measured by the time needed to complete procedures. Because the statistically significant demonstration of a moderate improvement in procedure time would require a large number of experiments and animals, we did not tailor the sample to accommodate this criterion. However, procedure duration was analyzed by subdividing time into...
duration of circumferential incision and duration of submucosal dissection. We estimated a potential time saving as a function of the lesion area, based on observed minimum and maximum procedure time/area ratios with both devices. Because the dimensions of resected tissue could differ slightly from one animal to another as well as within each experiment, a ratio of time to area was calculated for each ESD. Accessory criteria were the tissue and knife visibility and the ease of use of the device, as perceived by both endoscopists.

**Results**

ESD procedures were carried out uneventfully in all cases. In particular, no perforation was observed both during procedures and after autopsy on macroscopic examination of the gastric wall in the two groups. The visibility of catheter tip was excellent in all cases, as was that of the needle tip. Switching from one shape to another (i.e., hook to scythe to needle and back to hook) and fine-tuning the orientation of the wire (like “scything” parallel to the MP or “hooking” away from MP) were feasible in all cases without any dysfunction. Flushing through the lumen was found to be efficient in cleaning the operating field, but not always enough in lifting the submucosa.

All three available knife shapes (needle, hook, scythe) were used in all procedures with the new device. The scythe shape was
mostly used during the median or central part of the ESD, when the submucosal plane offered full and large visibility and a feeling of safety, whereas the needle shape was used essentially for edge incision and the hook shape for initial and final submucosal cuttings. Macroscopic findings are summarized in Table 1. The lesions were similar in size in the two groups (8.97 cm² in the “regular” knife group and 8.76 cm² in the “new knife” group, respectively P = 0.85). Microscopically, histopathologic analysis of resected tissue and gastric wall after ESD revealed similar features with both the regular and the new devices. Most of the submucosa was resected (approximately 500μm) with the mucosa and muscularis mucosae, whereas only a tiny fraction of the submucosa remained attached to the muscularis propria (Fig. 3). Although microscopic measurements of incision depth or thickness were not done on all resected specimens, no difference was observed between the thickness of submucosa after dissection with the regular and with the new devices. No undesirable contacts of the wire tip with the mucosa during “scythe” resection were observed macroscopically or microscopically in resected tissues.

No significant differences in time needed to complete the procedures were observed between procedures with the new devices vs the regular one (mean ESD time of 29.6 ± 11.56 minutes for the new device vs 35.5 ± 7.95 minutes for the regular one, P = 0.11), but that was as expected, given the small numbers of animals involved and procedures performed. However, overall procedure times were never longer with the new device than with the regular one; on the contrary, procedure time with the new device was dramatically shorter in several procedures (Table 1). Findings were the same when overall ESD procedure time was calculated as a fraction of the lesion area, approximated as a disc with a trend to shorter procedures with the new device (a mean of 3.80 min/cm² for the new device vs 4.33 min/cm² for the regular one), but not statistically different (P = 0.22). Some procedures were remarkably shorter with the new device, particularly in the latest experiments (Fig. 4a). In addition, when the submucosal phase of the dissection was taken into account separately, as was done with the last four animals, the same trend was observed, with two experiments showing no or only a little time

| N° animal | Type of ESD device | Circular Incision Time (min) | Submucosal Dissection Time (min) | Overall Dissection Time (min) | Lesion Diameter (Max × min, mm) | Lesion Area (mm²) | Time to Surface Ratio (min/cm²) |
|-----------|--------------------|-----------------------------|---------------------------------|------------------------------|-------------------------------|------------------|-------------------------------|
| 1 R e g u l a r | – – | 42 | 35 × 20 | 593.65 | 7.07 |
| 1 N e w k n i f e | – – | 26 | 28 × 20 | 452.16 | 5.75 |
| 2 P r o t o t y p e | – – | 40 | 37 × 30 | 880.9 | 4.54 |
| 2 N e w k n i f e | – – | 35 | 45 × 25 | 961.6 | 3.64 |
| 3 R e g u l a r | – – | 50 | 28 × 22 | 490.62 | 10.18 |
| 3 N e w k n i f e | – – | 50 | 30 × 25 | 593.65 | 8.41 |
| 4 R e g u l a r | – – | 36 | 30 × 20 | 490.62 | 7.33 |
| 4 N e w k n i f e | – – | 36 | 30 × 20 | 490.62 | 7.33 |
| 5 P r o t o t y p e | 5 23 | 28 | 30 × 20 | 490.62 | 5.70 |
| 5 N e w k n i f e | 5 23 | 28 | 30 × 20 | 490.62 | 5.70 |
| 6 R e g u l a r | 6 31 | 37 | 36 × 25 | 730.24 | 5.07 |
| 6 N e w k n i f e | 4 15 | 19 | 43 × 35 | 1193.98 | 1.60 |
| 7 P r o t o t y p e | 10 21 | 31 | 35 × 30 | 829.15 | 3.74 |
| 7 N e w k n i f e | 10 21 | 31 | 35 × 30 | 829.15 | 3.74 |
| 8 R e g u l a r | 7 18 | 25 | 28 × 28 | 615.44 | 4.07 |
| 8 N e w k n i f e | 4 18 | 22 | 28 × 26 | 572.26 | 3.84 |

1 Regular device corresponding to Flush-Knife™ (2 mm).

2 Lesion area is approximated as a disc, using the formula 3.14 × [(Diam Max + Diam min)/2]^2.
saving, and the other two exhibiting a remarkable fall in sub-procedure time (Fig. 4b). Finally, a simulation of the potential time saving as a function of the lesion area, based on observed minimum and maximum procedure time/area ratios with both devices, showed that the new device could save up to 66% of the procedure time when comparing minimum time/area ratios for both devices (Fig. 5).

Discussion

The new device evaluated in this study is similar to existing ESD knives in that it presents as a catheter-based, handle-controlled monopolar electrosurgical device. More specifically, it shares with the Flush-Knife™ a sideport for intraprocedural flushing to help clean the device tip and the operating field. However, this new ESD device differs from conventional ones in its unique versatility and the extended range of cutting wire to be put in contact with tissue, from the very tip for marking and incision to

Fig. 4  Time/surface ratio for completing and ESD with the standard ESD knife vs the new device. a Overall results in all animals. b Results for the submucosal phase of ESD in the second series of 4 pigs.

Fig. 5  A model of time/surface required to complete an ESD with a standard device vs the new device. Min and max are based on data observed during the animal study.
nearly a half snare for quick submucosal cut. As shown in the figures and videos, this ESD instrument can be used as a standard needle, as a hook that can be easily oriented to grasp and cut submucosal collagen fibers, and more specifically as a “scythe” with lateral motion of the endoscope or by using gentle forward motion of the wire fully extended and oriented in a plane parallel to that of the muscularis propria.

It is worth observing that in several experiments, ESD was achieved more expeditiously with the new device than with conventional ones to which both operators were used before starting the study, and that this time saving was entirely due to improved performance in the second part of the study, exhibiting a learning-curve effect. Although a trend to a decreased operating time over the course of the study was also visible with the regular knife, probably as a consequence of the standardization of lesion size and location (Fig. 4a), this trend was more pronounced with the new knife despite having little or no experience with it, in contrast to the Flush-knife™. This learning curve seems relatively short, since fewer than five procedures were sufficient to observe some time savings that, while not statistically significant, was not detrimental to the safety of procedures. Previous similar studies of new devices failed to show a significant time-saving compared to earlier-generation ESD knives while exhibiting suboptimal versatility and possibly less accurate submucosal dissection [8]. Intra-procedural complications were a matter of concern because the device offers a long extension of bare wire that could harm and puncture the muscular layer. However, no complications were observed in either group and there were no more superficial muscular bruises with the new device ESDs than when the conventional device was used, on either macroscopic or microscopic examination. This is probably explained, in part, by the protective effect of the transparent hood.

A caveat for the use of the device in its “scythe” position is probably to avoid extending the catheter more than a few millimeters beyond the tip of the hood, as was done in this study. With that in mind, we consider the scythe shape to be the most innovative feature of this instrument, and one that can save the most time in the submucosa, as observed in the swiftest of our procedures. It is difficult to know if the device presents a safety profile different from regular devices with regard to risk of bleeding because the pig model is not a good one for intraoperative bleeding, but there is no particular reason for concern on that issue. The new device has the same ability to cut a vessel and trigger bleeding and to coagulate it as do other knives. As with any other ESD knife, it is certainly necessary to consider having coagulation graspers at one’s disposal when undertaking an ESD.

Another point this study cannot address completely is the ability of the device to achieve ESD in especially difficult locations. Because our goal was to perform two ESDs in reproducible conditions, we had to choose relatively “standard” locations and levels of difficulty. For example, we did not perform ESDs in retroflexed positions or in the distal part of the stomach, which in the pig model requires deep insertion of the endoscope with a long loop formation. We assume that the device’s ability to conform to many types of angulation and lengths of active wire would make dissection much easier in such difficult locations than with other knives, but we recognize the need to confirm that with further studies. We wondered whether the flushing lumen could inject fluid with pressure sufficient to lift the submucosa during ESD. As indicated previously, we found that function to be suboptimal, but significant improvement has been made in that regard in more recent prototypes, which were not tested in this study. The last caveat for use of this new device is that some training is necessary to correctly manipulate the handle because it is slightly more complex than the push-pull handling of a standard knife or resection snare. Therefore, we recommend that a trained and skilled nurse be present when ESD is performed. Improvements to the handle are under way.

Finally, this study has several limitations: 1. Gastric ESD in the pig differs from that in humans because of the pig’s distinctive gastric wall features (thick mucosa, few submucosal vessels, mucosal lifting easily sustained); 2. A small number of procedures were performed; and 3. Experience with the new device is limited. We consider this new device as of potentially significant help in making ESD both easier and less time-consuming without compromising the safety of procedures. Thus we are contemplating initiating a clinical study with this novel ESD knife.

Competing interests: The author (Frederic Prat) declare that a patent is pending with regard to the device investigated with co-ownership of Medwork GmbH and Assistance Publique-Hôpitaux de Paris (AP-HP).

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