Safety analysis of endoscopic haemostasis using a high-frequency live tissue electric welding device – EKVZ300 PATONMED

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Abstract

Introduction: The method of a bipolar high-frequency welding (HFEW) of soft living tissues of animals and humans has been used in various areas of surgery. However, it has not been utilized in endoscopic gastrointestinal procedures yet. HFEW has strong potential to be used in gastrointestinal endoscopic procedures due to the competitive cost of generator devices and due to its proven ability to coagulate vessels of wide diameter as compared to standard electrocautery devices.

Aim: To investigate the effectiveness of the endoscopic haemostasis using HFEW generator device – 300 PATONMED – in a porcine model of arterial gastrointestinal bleeding.

Material and methods: A porcine model of arterial gastrointestinal bleeding was created. A 300 PATONMED set to the “welding” regime and a flexible 7 Fr bipolar coagulation probe with two electrodes on the tip fashioned spirally attached to convey energy were tested. Once bleeding from the artery had been initiated, the bipolar probe was applied to coagulate the bleeding site. Animals were observed for clinical evidence of recurrent bleeding and subsequently were euthanised for histological examination.

Results: A total of 10 experiments were successfully completed. An optimal haemostatic effect was achieved with durations of cautery of five to eight seconds in all animals. Continuous observation after haemostasis revealed no evidence of re-bleeding. No systemic side-effects of the technique were observed. Histological examination has shown that the peripheral thermal injury area that surrounded the coagulation zone did not spread beyond the mucosal layer in depth and 2 mm in width.

Conclusions: This animal study provided evidence for the safety of an HFEW in the treatment of gastrointestinal bleeding. The advantages of this technology are smokeless operative area, no tissue overheating, minimal necrosis and damage to surrounding gastric tissue, and the fact that the area of HFEW is confined to the area of the electrodes.

Introduction

Acute non-variceal upper gastrointestinal bleeding (NVUGIB) remains a common, potentially life-threatening condition. Endoscopic haemostatic therapy has been shown to improve outcomes of NVUGIB [1–3]. There are many safe and effective devices available for endoscopic haemostatic therapy in NVUGIB; for example, contact thermal devices, noncontact thermal devices, injection needles, and mechanical devices. A combination of any of the treatment modalities is commonly used to control NVUGIB. However, there are few compelling data favouring a particular device for treatment of NVUGIB, and selection of the optimal haemostatic device depends on the characteristics of the lesion, local expertise, equipment availability, and cost. The best combination for endoscopic haemostasis should obtain the highest rate of haemostasis while minimising the rates of re-bleeding and adverse events, it should be easy to use in all locations, and it should be inexpensive [4].

The method of a bipolar high-frequency welding (HFEW) of soft living tissues of animals and humans
has been used in various areas of surgery [5–9]. The method is based on measurement of tissue impedance to accurately control the degree of coagulation and, unlike conventional bipolar coagulation technique, allows avoidance of over-heating and burning of the exposure area, sticking of the electrodes to the tissue, and the production of smoke [10, 11]. However, HFEW has not been utilised in endoscopic gastrointestinal procedures yet. High-frequency welding has strong potential to be used in gastrointestinal endoscopic procedures due to the competitive cost of generator devices and its proven ability to coagulate vessels of wide diameter as compared to standard electrocautery devices.

Aim

The aim of this study was to investigate the safety and effectiveness of endoscopic haemostasis using a HFEW generator device – a 300 PATONMED (E.O. Paton Electric Welding Institute of the National Academy of Sciences of Ukraine, Kiev, Ukraine) – in a porcine model of massive arterial gastrointestinal bleeding.

Material and methods

Preparation and development of porcine bleeding gastric ulcer model

A porcine model of arterial gastrointestinal bleeding was created using standard technique with minor modification [12, 13]. Six healthy female domestic farm pigs (Sus scrofa domesticus) were observed for 1 week prior to study initiation, to ensure baseline health. Three days prior to the procedure, omeprazole was administered to all animals. One day prior to the procedure, omeprazole, famotidine, and antacids were given. All feeding was withheld for 12 h prior to the endoscopy. The baseline haematocrit, platelets, and activated clotting time (ACT) were measured in all animals prior to the procedure. They were observed for clinical evidence of reflux. Three days before the procedure, omeprazole was administered to all animals. One day prior to the procedure, omeprazole, famotidine, and antacids were given. All feeding was withheld for 12 h prior to the endoscopy. The baseline haematocrit, platelets, and activated clotting time (ACT) were measured in all animals prior to the procedure. They were observed for clinical evidence of reflux.

Endoscopic device and procedure

An HFEW generator device – 300 PATONMED (E.O. Paton Electric Welding Institute of the National Academy of Sciences of Ukraine, Kiev, Ukraine) – set to “automatic welding” regime with electric resistance 20 Ohms and power 200 W was connected to a flexible 7 Fr bipolar coagulation probe with two electrodes on the tip fashioned spirally. The “Automatic welding” regime identifies the optimal time required to achieve complete coagulation by measuring tissue impedance using the device’s software and makes a sound signal to stop coagulation. This complex was tested in the prepared animal models to coagulate arterial defects. Once bleeding from the LGA had been initiated by the means of “pure-cut” electrocautery, the bipolar probe was inserted through the channel of the gastroscope. The bipolar probe was applied perpendicularly or tangentially with mild to moderate pressure for 3 to 8 s to compress and coagulate the bleeding site. The bleeding spot was irrigated with normal saline to ensure completeness of haemostasis.

Follow-up and post-mortem examination

All animals were fed on the same day after the operation. They were observed for clinical evidence of recurrent bleeding. They were not put on proton pump inhibitors. All animals were euthanised 10 days after the procedure. Qualitative histologic analysis of each site of probe application was performed on formalin-fixed specimens by an independent board-certified veterinary pathologist.
Outcomes

The primary end point was the survival of the animals after thermal endoscopic haemostasis using HFEW technique in an ulcer bleeding model. The secondary end point included the time to achieve endoscopic haemostasis when using the HFEW-300 PATONMED device, the rate of recurrent bleeding, and the extent of local injury.

Results

A total of 10 experiments were successfully completed. Creation of an active arterial spurting haemorrhage model followed by endoscopic needle-knife injury was successful in all 10 animals. An optimal haemostatic effect was achieved with 200 W of power and durations of cauterity of 3 and 8 s. The process was smokeless and there was no sticking of the electrode to the coagulation zone. There was no overheating observed. Continuous observation after haemostasis revealed no evidence of re-bleeding. Repeat endoscopy on day 8 revealed no evidence of active bleeding and a healed but identifiable gastrotomy site in all animals. There was no adverse event in the survival period. All animals tolerated a full diet. Histopathological findings showed a precise demarcated coagulation zone at the LGA wall, which matched the size of the electrode. This zone was surrounded by peripheral tissue injury. This injury was manifested by destruction of epithelial cells of the outer zone of gastric glands and leukocytic mucosal infiltration reaching a maximum depth of 0.5 mm. Deep parts of glands were mostly not involved, and in all cases the maximum spread of injury laterally was 2 mm (Figure 1).

Discussion

This animal study provides evidence for the safety of a HFEW in the treatment of active, severe, NVUGIB. The size of an artery is one of the main determinants of the success of endoscopic haemostasis. Our model simulated bleeding ulcers with a large vessel at the base. Therefore, an ulcer model with haemorrhage commonly refractory to existing endoscopic devices was created. Initial haemostasis was achieved in all animals, and there were no systemic or local procedure-related complications. The procedure was smokeless, and there was no sticking of the electrode to the tissue – this was due to the device’s ability to control coagulation and to avoid overheating. In all cases the depth of the injury did not spread beyond 0.5 mm, due to the nature of the controlled thermal effect of the device, and hence low risk of gastric perforation.

The artery diameter that was successfully coagulated is comparable to the size of the gastroduodenal artery in humans. However, the chronicity of ulcer characterised by hard fibrotic tissue could not be achieved in this experiment. The technique is operator dependant like all contact thermal haemostatic techniques, hence potential obstacles related to operator experience and bleeding location in achieving haemostasis may arise in clinical practice. The learning curve of this technique was not specifically assessed in this study, but it may be comparable to the learning curve of all endoscopic contact thermal haemostatic devices. Cost effectiveness analysis of this technique in comparison to conventional techniques has not been performed, but based on the market price of the generator device at the time of publication, the 300 PATONMED has a more favourable price when comparing it to other known bipolar and argon-plasma cautery devices.

Conclusions

From the results of the study, we believe that the HFEW generator device, the 300 PATONMED, provides a feasible and reliable technique in achieving endoscopic haemostasis on bleeding peptic ulcers. There are several advantages of this technique over conventional methods. However, the data provide only limited conclusions. The next logical step would be to assess clinical application of the 300 PATONMED in humans.

Conflict of interest

The authors declare no conflict of interest.

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