Early clinical experience of the safety and efficacy of EndoClot in the management of non-variceal upper gastrointestinal bleeding

Sabina Beg, Ibrahim Al-Bakir, Meha Bhuvva, Jay Patel, Mark Fullard, Anthony Leahy

Background and study aims: EndoClot is a novel topical hemostatic powder approved for use in non-variceal upper gastrointestinal bleeding. This study examines its impact as rescue therapy in the management of gastrointestinal bleeding for which standard endoscopic therapy failed to achieve hemostasis.

Methods: This observational study covered a 24-month period. Data were collated from patients treated with EndoClot for comparison with a cohort of patients managed with standard endoscopic therapy. End points of this study included immediate hemostasis, 30-day rebleed rate, 30-day mortality rate, and adverse events.

Results: Between April 1, 2012, and March 31, 2014, gastroscope procedures were performed in 1009 patients, of whom 173 required endoscopic therapy. EndoClot was used in 21 patients, with immediate hemostasis achieved in all cases, a 30-day rebleed rate of 4.8 % (95 % confidence interval [95 %CI] 4.34% to 3.94%), and a 30-day mortality rate of 19.0 % (95 %CI 2.29% - 35.91 %). Despite higher risk bleeds in this cohort of patients, Fisher’s exact test demonstrated no significant difference between their 30-day mortality rate (P=0.51) and rebleed rate (P=0.31) and those of the patients treated with standard endoscopic hemostatic techniques.

Conclusions: This study demonstrates that EndoClot can be used both safely and effectively in the management of non-variceal upper gastrointestinal bleeding.
forces that cause the particles to slough off and enzymatic degradation by endogenous amylase and glucoamylase [8]. EndoClot is administered through a spray catheter, with particles dispersed over a large field of distribution, making precise deployment unnecessary. The catheter is not required to come directly into contact with the bleeding lesion during application, so that there is no risk of causing further tissue damage or exacerbating bleeding. As a consequence of these features, EndoClot requires minimal technical expertise to use, in comparison with conventional treatment modalities (8 – 10). The exact duration of adherence to the mucosa is unknown but is thought to range between 1 and 48 hours; patient factors and lesion characteristics are likely to be influential.

EndoClot is an addition to a growing variety of commercially available hemostatic powders, which include Hemospray (Cook Medical, Bloomington, Indiana, USA) and Ankaferd Blood Stopper (Ankaferd Health Products, Istanbul, Turkey). These powders have been adopted from military medicine and have been successfully used in emergency situations. Their characteristics and role in the context of upper gastrointestinal bleeding are currently under evaluation, with promising results [9].

 Patients and methods

This study evaluated the impact of EndoClot in the management of non-variceal upper gastrointestinal bleeding in a large district general hospital serving a population of more than 500,000 people. Although large randomized controlled trials will ultimately be required to advocate the widespread use of this product, at present there is paucity of even preliminary evidence to guide the effective use of EndoClot. Rather than a formal trial designed to demonstrate superiority, we present here our early clinical experience of EndoClot, comparing outcomes with those of patients treated with the more established techniques.

This was a single-center retrospective observational study conducted over a 24-month period. A pragmatic approach to the use of EndoClot was adopted because even in the event of proven benefit, the cost of this product in comparison with that of standard hemostatic techniques would likely preclude its routine use as first-line therapy. To reflect realistic clinical practice, EndoClot was applied at the endoscopist’s discretion only if conventional endoscopic treatment with dual or triple therapy had failed to accomplish complete hemostasis or was technically impossible. The product was not used as monotherapy, primary therapy, or prophylaxis. All procedures were performed or supervised by consultant gastroenterologists with appropriate experience in the management of gastrointestinal bleeding. Again, to reflect daily clinical practice, patients with non-variceal upper gastrointestinal bleeding of all etiologies were considered for EndoClot treatment and included in this analysis.

The EndoClot application system consists of 2 g of powder within a mixing chamber, which is applied through a 2300-mm delivery catheter fed through the working channel of an endoscope. An external air compressor creates the sustained force required to drive the powder from within the chamber through the catheter, resulting in the multidirectional distribution of the product onto the mucosa (Fig. 1, Fig. 2). Given the wide field of distribution, an en face view of the point of bleeding is not essential; this is particularly useful for lesions that are notoriously difficult to access, such as ulcers on the posterior duodenal wall. Both the endoscopists and the endoscopy nurses underwent a brief training session in the use of this system before its introduction.

The decision to use EndoClot was made at the time of endoscopy and so did not affect pre-endoscopy treatment, including appropriate resuscitation. There was no limit on the quantity that could be used to achieve cessation of bleeding. Failure of treatment was defined as the inability to achieve hemostasis on completion of the endoscopic procedure. Following treatment of the bleeding lesion, standard medical therapy was continued with inpatient observation and, for ulcer bleeds, a 72-hour infusion of a proton pump inhibitor. Patients did not routinely undergo a second-look endoscopy unless clinical or biochemical evidence of recurrent bleeding was noted.

We evaluated all patients with non-variceal upper gastrointestinal bleeding requiring endoscopic therapy over a 24-month period between April 1, 2012, and March 31, 2014. The demographic and clinical data of those treated with EndoClot were collated for comparison with the data of the cohort of patients managed with standard endoscopic therapy. The primary outcome measure of this study was successful hemostasis, defined as the cessation of active bleeding as visualized by the endoscopist at the time of the procedure. Secondary outcomes included 30-day rebleed rate,
Results

During the 24-month study period, 1009 patients underwent endoscopic procedures for non-variceal upper gastrointestinal bleeding, of whom 173 patients required endoscopic therapy (Fig. 3, Table 1). Those who received treatment had a median age of 77 years (range 18–101). The male-to-female ratio was 2:1. The majority of the endoscopic procedures were performed during the working day within 12 hours of presentation; however, 3 patients underwent endoscopy out of hours in an operating theatre, and 10 patients required stabilization in intensive care with intubation. The lesions accounting for active bleeding included the following: duodenal ulcer (n = 96), gastric ulcer (n = 23), esophageal ulcer (n = 16), malignancy (n = 11), Mallory–Weiss tear (n = 9), gastric antral vascular ectasia (n = 5), sphincterotomy bleeding after endoscopic retrograde cholangiopancreatography (n = 4), angiodysplasia (n = 3), Dieulafoy lesion (n = 3), non-specific oozing (n = 1), aneurysm (n = 1), and bleeding after biopsy (n = 1). There were 24 deaths within 30 days (13.9%; 95% CI 7.74%–17.66%) in patients with a median age of 81 years; duodenal ulcers were the underlying pathology in the majority of cases. A single endoscopic modality was used in 43 patients; this cohort consisted mainly of patients with low risk lesions, such as known malignant lesions with oozing, angiodysplasia, and gastric antral vascular ectasia. Given the more indolent course of these lesions, the patients in this group were deemed inadequate for comparison and therefore excluded from our analysis.

Dual endotherapy was required in 91 patients, the most common combination being epinephrine with gold probe. This resulted in a rebleed rate of 8.8% (95% confidence interval [95%CI] 2.98%–14.62%) at 30 days, with 1 patient requiring surgery to achieve definitive hemostasis. There was a 30-day mortality of 14.3% (95%CI 7.11%–21.49%). There were 18 patients who had bleeding lesions requiring triple therapy with gold probe, epinephrine injection, and clips. The associated 30-day rebleed rate was 11.1% (95%CI 3.41% to 25.61%), and the 30-day mortality was 11.1% (95%CI 3.41% to 25.61%). No patient receiving dual or triple therapy required surgery.

EndoClot was used in a total of 21 patients: this was as a second agent in 7 cases, a third agent in 9 cases, and the fourth form of endoscopic therapy in 5 cases. The lesions treated included duodenal ulcer (n = 14), Mallory–Weiss tear (n = 2), gastric ulcer (n = 2), malignancy (n = 1), esophageal ulcer (n = 1), and non-specific gastric oozing (n = 1). Of the treated lesions, 5 were characterized...
as 1a and 16 as 1b per the Forrest classification [11]. EndoClot was used by 7 different endoscopists, giving a median experience with this product of 3 applications per operator (range 1–5).

Immediate hemostasis was achieved in all patients treated with EndoClot. A single patient underwent endoscopy because of rebleeding; although the second gastroscopy demonstrated active bleeding, this was from a newly diagnosed Mallory-Weiss tear rather than the originally treated duodenal ulcer and therefore is not representative of a failure of EndoClot therapy. Another patient, in whom a large malignant duodenal ulcer was diagnosed, demonstrated biochemical and clinical evidence of rebleeding; however, it was deemed inappropriate to perform a second endoscopy. The cohort treated with EndoClot had a 30-day rebleed rate of 4.8% (95% CI – 4.3% to 13.9%). The 30-day mortality was 19.0% (95% CI 2.2% – 35.9%), although only 1 death was attributable to upper gastrointestinal bleeding (this was the patient with the aforementioned malignant ulcer, for whom surgery was not an option because of significant co-morbidities).

There was no significant difference between the 30-day mortality rate and rebleed rate of the group of patients treated with EndoClot and the rates of those successfully treated with dual or triple endoscopic therapy alone (P = 0.51 and P = 0.31, respectively, with Fisher’s exact test).

In no instance did the use of the EndoClot system complicate or potentiate bleeding. In two cases, the administration of treatment was hampered by occlusion of the spray catheter with activated particles. There were no reported side effects in association with the use of EndoClot. Two patients underwent another endoscopy following the use of EndoClot for suspected rebleeding; both procedures occurred within 24 hours of the initial gastroscopy. Residual EndoClot was not seen in either patient.

**Discussion**

Upper gastrointestinal bleeding remains a common medical emergency, with endoscopic hemostasis the treatment of choice in patients with high risk lesions. The annual incidence of upper gastrointestinal bleeding requiring endoscopic therapy in our population was 8.6 per 100,000 patients. Our institution offers a daily endoscopy list for upper gastrointestinal bleeding within the endoscopy department 7 days a week and an out-of-hours on-call service on weekends. During the week, there is an ad hoc arrangement, with out-of-hours endoscopy performed in the operating theatre with theatre nurses and anesthetic support if required. This setup is not atypical and makes the availability of an effective hemostatic agent desirable [1, 2, 9, 10]. Our data support the main advantages of EndoClot—namely, efficacy comparable with that of the more established therapeutic modalities, ease of use, ability to overcome challenging anatomy, and the lack of complications [8–10].

By definition, the cohort receiving EndoClot therapy in our study included many of the patients at highest risk because they were the patients in whom standard endotherapy failed to achieve hemostasis. It is therefore reassuring to note that the addition of EndoClot therapy resulted in no statistical difference between the 30-day mortality rate (P = 0.51) and rebleeding rate (P = 0.31) of these patients and the rates of the patients in whom hemostasis was achieved with standard dual or triple therapy alone. It was possible to use EndoClot despite limited experience, with a median of three applications per operator. There was no discernible difference in outcomes between those with the least experience and those with greater experience with this system. On two occasions, EndoClot was used outside the endoscopy department, where ease of use was demonstrated; these procedures were assisted by operating theatre nurses who had no specific training in the use of EndoClot.

A difficult anatomical position of the lesion was the indication for EndoClot use in the majority of cases, most commonly in the duodenum. Diffuse bleeding was seen in one patient; this is likely to be an increasing problem with the advent of novel oral anticoagulants, for which at present there is no clear consensus regarding endoscopic management.

In our study, no complications were associated with the use of EndoClot. Theoretical risks include allergic reaction, embolization, and intestinal obstruction. Limited data have been published on the use of EndoClot, so it is unclear how frequent these complications are likely to be; greater experience is required, in addition to vigilance for their occurrence.

Of the available hemostatic powders, Hemospray has perhaps the most evidence behind its use, with safety and ease of use demonstrated in case reports and several small case series. Our preliminary data compare favorably with the immediate success rate of Hemospray of 88.5% and rebleed rate of 16.2%, described in a recent review of 234 pooled patients [12–14]. Hemospray has been tested in a wide range of clinical scenarios, with growing evidence for its use in bleeding associated with portal hypertension or ischemic colitis, bleeding after endoscopic mucosal resection, and even diverticular bleeding [15–18]. Randomized controlled trials are awaited.

Ankaferd is licensed for use in Turkey but is not currently available in the United Kingdom. Several small case series have been published, with heterogeneity in the pathological conditions and populations studied. One of the larger series, including 27 patients, demonstrated immediate hemostasis in 73% with a rebleed rate of 20% [19]. No studies have compared the available hemostatic agents in regard to efficacy, although with increasing interest and their greater position in the marketplace, head-to-head studies are likely to follow.

The main limitations of this study are its observational nature and the relatively small number of patients who received EndoClot therapy. EndoClot was used only as rescue therapy in the study, so it is difficult to speculate on its potential efficacy as primary monotherapy or as part of conventional dual therapy. We are unable to compare usefully the efficacy of EndoClot with that of other commercially available hemostatic agents. However, the study has demonstrated the successful use of EndoClot in a range of pathological conditions, with promising results. Greater experience is required to fully understand the potential of topical hemostatic agents, such as EndoClot. It is unlikely that conventional endoscopic techniques will be discontinued in favor of these; however, EndoClot is clearly a useful tool in the armamentarium of treatments for non-variceal upper gastrointestinal bleeding.

**Competing interests:** None

**Acknowledgment**

Images were provided by Vitramed.
References


Beg Sabina et al. EndoClot in the management of non-iceal upper gastrointestinal bleeding ... Endoscopy International Open 2015; 03: E605–E609