Safety and effectiveness of a synthetic hemostatic patch for intraoperative soft tissue bleeding

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Introduction

Uncontrolled intraoperative bleeding is a dangerous complication that can hinder both the surgeon’s ability to complete the procedure and the patient’s ability to recover.¹ Blood that remains intra-abdominally after surgery not only increases the extent and severity of postoperative intestinal adhesions, but also serves as a source of intra-abdominal infections.² Blood transfusions may become necessary in some cases; these increase the risk of postoperative morbidity and mortality.³,⁴ Furthermore, inadequate hemostasis increases operative time, recovery, and length of hospital stay, and constitutes a considerable economic burden.⁵ Conventional methods of obtaining hemostasis at continuously bleeding sites include repetitive direct cauterization, placement of sutures or clips, and prolonged direct compression, but these methods can impair tissue healing and recovery through the generation of tissue damage, char, and necrosis.⁶,⁷ Conventional methods might also be impractical or ineffective in regions that are hard to access or in organs and tissues located nearby that can be easily damaged (eg, nerves or the respiratory tract). Therefore, the development of more effective hemostatic agents is critical in order to keep surgical costs down and provide the best safety for patients.

Background: Continuous bleeding after using conventional hemostatic methods involving energy, sutures, or clips, is a serious and costly surgical complication. Many topical agents have been developed to promote intraoperative hemostasis, but improvement is needed in both decreasing time to hemostasis and increasing ease of use. Veriset™ hemostatic patch is CE-marked for controlling bleeding on the liver and in soft tissue. In the current study, we aimed to gather further evidence for the safety and effectiveness of Veriset™ hemostatic patch in soft tissue bleeding during a variety of surgical procedures.

Methods: Thirty patients scheduled for nonemergency surgery, each with an intraoperative soft tissue bleeding site, were treated with Veriset™ hemostatic patch. Time to hemostasis was monitored, and adverse events were assessed during the 90 days after surgery.

Results: When Veriset™ hemostatic patch was used, hemostasis occurred within 5 minutes in 29/30 (96.7%) subjects and within 1 minute in 21/30 (70.0%) subjects. No device-related serious adverse events were recorded during the 30 days after surgery, and no reoperations for device-related bleeding complications were performed during the 5 days after surgery.

Conclusions: Veriset™ hemostatic patch is a safe and effective hemostat for controlling soft tissue bleeding during a variety of surgical procedures.

Keywords: Veriset™ hemostatic patch, hemostasis, topical hemostat
Many of the commercially available hemostats have been associated with adverse safety.9 Hemostats that are based on human- or animal-derived components can lead to virus and disease transmission or elicit anaphylaxis.9-12 Additionally, multiple reports have shown granuloma formation and neurological effects associated with the use of some hemostatic agents if not removed.13-19 These observations support the need to develop products that effectively achieve hemostasis in a diverse patient population and introduce minimal safety concerns. An ideal hemostatic agent has the potential to decrease the use of blood products in elective and emergency surgery and demonstrates the ability to manage bleeding from both parenchymal and small or inaccessible arterial and venous structures. Such a product would also require adhesive and mechanical strength to withstand the pressure of bleeding.20

Veriset™ hemostatic patch (Covidien, Mansfield, MA, USA) is a recent addition to the plethora of available hemostatic agents. It is composed of an absorbable backing material, oxidized cellulose, and hydrogel components. Veriset™ hemostatic patch is provided as a ready-to-use patch that is applied polyethylene glycol-side down to the bleeding site. Unlike many other hemostatic agents, which require a dry field prior to application, Veriset™ hemostatic patch works with physiological fluids to promote hemostasis. Minimal preparation is required for Veriset™ hemostatic patch, and the device has been shown in preclinical studies to be fully absorbable (Howk, unpublished data, 2015). The device lacks any human- or animal-derived components; thus, it is unlikely to transmit viruses or elicit an immune response.

Veriset™ hemostatic patch is CE-marked for use in laparoscopic and open procedures involving solid organ or soft tissue bleeding. A previous randomized, controlled clinical study compared Veriset™ hemostatic patch to TachoSil® (Nycomed Austria GmbH, Linz, Austria) in hepatic procedures.21 The number of subjects who achieved hemostasis within 3 minutes was significantly greater among those treated with Veriset™ hemostatic patch than among those treated with Tachosil®. In addition, Veriset™ hemostatic patch was determined to be both safe and easy to use. The aim of the present study is to provide further evidence for the safety and effectiveness of Veriset™ hemostatic patch for controlling intraoperative bleeding in soft tissue. The clinical trial registration number for this study is NCT01719172, and it can be found at http://www.clinicaltrials.gov.

Materials and methods

Study design

This study was a prospective, multicenter, single-arm study of Veriset™ hemostatic patch to assess its safety and effectiveness in obtaining hemostasis in patients undergoing visceral surgical procedures in soft tissue that typically result in considerable sizes of intra-abdominal or intrathoracic connective tissue wound areas. In this study, surgical procedures were included that are likely to lead to hemorrhage in soft tissue areas, which are defined as areas containing tissue that connects, supports, or surrounds other structures of the body or parts of organs. From the perspective of the surgical problem addressed (bleeding from connective tissue in an otherwise highly vulnerable field), inclusion of a variety of procedures was expected to result in a tissue surface that was quite similar across procedures.

During surgery, Veriset™ hemostatic patch was applied to the target bleeding site (TBS), and its effectiveness was analyzed by capturing the time required for hemostasis to be achieved. Safety was determined by the number of device-related serious adverse events (SAEs) during the 30 days after surgery and the incidence of reoperation for device-related bleeding complications during the 5 days after surgery. Subjects were assessed during the procedure and then 24 hours, 7 days, 30 days, and 90 days after the operation. The clinical investigation was conducted according to Good Clinical Practice guidelines and the Declaration of Helsinki, and abided by European and national regulations.

Study participants

A total of 30 subjects participated in the study. They were selected from three European sites on the basis of protocol-defined inclusion and exclusion criteria. Preoperative inclusion criteria required subjects be ≥18 years of age and already scheduled for nonemergency surgery via an open approach, in which a topical hemostatic agent can be used to control bleeding if conventional methods using energy, sutures, or clips are impractical or unsuccessful. Subjects who were pregnant, breast-feeding, undergoing emergency surgery, scheduled for another planned surgery that could jeopardize study treatment, had a life expectancy of less than 6 months, or had participated in an investigational drug or device study within 30 days of enrollment that would interfere with this study were excluded from eligibility. All eligible subjects provided informed consent and agreed to comply with the treatment and evaluation schedule. Enrolled subjects who did not meet intraoperative criteria were considered screen failures;
these included subjects who exhibited a local infection at the TBS, whose safety or welfare was determined to be at risk by the investigator, or who did not have a bleeding site of Type 2 (oozing/mild) or Type 3 (moderate) severity (a rating system modeled after classification established by the Bleeding Academic Research Consortium).22

Materials
Veriset™ hemostatic patch was provided in 5×10 cm² sheets that could be cut to the appropriate size. Radiopaque gauze (Covidien) was used to assess relative intraoperative bleeding severity prior to the application of Veriset™ hemostatic patch.

Surgical procedure
The surgical procedure was performed in accordance with the appropriate practices of the institution at which the procedure occurred. After observing an active bleeding site in soft tissue and determining that conventional methods of obtaining hemostasis were either impractical (an assessment based on the investigator’s discretion) or ineffective (an assessment based on a failed attempt to achieve hemostasis with one or more of these methods), the site was selected as the TBS. Veriset™ hemostatic patch could be cut as necessary to cover the bleeding site with 1–2 cm margins. Multiple patches could be used, if required, with a maximum dosage of 6.43 cm²/kg. After Veriset™ hemostatic patch was placed over the TBS, direct pressure was applied. Hemostasis was assessed every 30 seconds until the 5-minute time point, then at 1-minute intervals for minutes 5–10 until hemostasis was achieved. Pressure was maintained in between assessments. Once bleeding had appeared to stop, the TBS was monitored for 1 minute with no pressure to confirm hemostasis. If rebleeding occurred, the assessment was continued at 30-second intervals until 5 minutes, and at 1-minute intervals for minutes 5–10. Additional Veriset™ hemostatic patch devices were applied if it was noticed that the initial application did not completely cover the TBS, and the timing of the procedure was continued during the subsequent applications. If bleeding persisted after 2 minutes, either continued pressure was maintained or an additional Veriset™ hemostatic patch was applied.

Assessing intraoperative bleeding with gauze
Intraoperative bleeding was assessed with gauze prior to the application of Veriset™ hemostatic patch. Once identified, extraneous fluids were removed. Then, ten gauze pads (40 layers total) were placed on the TBS, and gentle pressure was applied for 3 seconds. After 3 seconds, the gauze was removed, and the number of layers penetrated with blood was recorded. Eligibility was based solely on the discretion of the investigator, regardless of results from the gauze method of assessment, and the TBS was assigned a bleeding severity of Type 2 or Type 3.

Outcome measures
The primary effectiveness endpoint was the percent success in obtaining hemostasis within 5 minutes after the application of Veriset™ hemostatic patch. Secondary effectiveness endpoints included the percentage of subjects who achieved hemostasis within 1 minute and the median time required to achieve hemostasis. The primary safety endpoint was the number of device-related SAEs per subject during the 30 days after surgery. The secondary safety endpoint was the incidence of reoperation for device-related bleeding complications during the 5 days after surgery.

Postoperative visits
To determine adverse events (AEs), subjects were assessed at 24 hours after skin closure by laboratory tests, by testing of vital signs, and by surgical site/infection analyses. A phone call was made 7 days after the surgery to follow up on AEs. Subjects were assessed on the same characteristics 30 days and 90 days after surgery as during the 24-hour postoperative follow-up.

Statistical analysis
Sample size determination was based on the primary effectiveness endpoint, success in obtaining hemostasis within 5 minutes of Veriset™ hemostatic patch application. The sample size was calculated by using a one-sided exact test of a binomial proportion versus an objective performance criterion value for 0.5 (50%) to be the proportion of subjects obtaining hemostasis within 5 minutes. The test was performed with a true success proportion of 0.75, an alpha of 0.025, and a power of 80%. The required sample size based on these specifications is 30 subjects.

All statistical tests performed were one-sided at the 2.5% significance level. To analyze the effectiveness of Veriset™ hemostatic patch in achieving hemostasis, an exact (Clopper–Pearson) 95% confidence interval for the true success percentage was calculated. A one-sided exact test based on the binomial distribution was performed to test the null hypothesis that the true percentage of subjects who achieved
hemostasis is less than or equal to 50%; the alternative hypothesis was that the true percentage is greater than 50%. Time to hemostasis was analyzed using the Kaplan–Meier method to estimate the median time to hemostasis. A 95% Brookmeyer–Crowley confidence interval for the median was computed on the basis of the sign test.

To analyze the safety of Veriset™ hemostatic patch, a 95% confidence interval for the mean number of device-related SAEs per subject was calculated on the basis of the t-distribution. An exact (Clopper–Pearson) 95% confidence interval for the incidence of reoperation for device-related bleeding complications was calculated. Paired t-tests were used to analyze changes in vital signs and other laboratory tests from baseline values. Statistical analyses of data were performed by using SAS Version 9.1 or higher (SAS Institute, Inc., Cary, NC, USA).

Results

Surgery

A total of 37 subjects consented to and were enrolled in the study, but seven did not meet preoperative or intraoperative criteria and were considered screen failures (Figure 1).

The remaining 30 subjects were treated with Veriset™ hemostatic patch, but two of the 30 subjects did not complete the study. Of those two subjects, one was lost to follow-up and the other died during the study from embolic complications of a pre-existing atrial thrombus that was unrelated to the test device.

Subjects underwent various types of surgical procedures, of which the most common were lymphadenectomy, esophagectomy, colectomy, and pancreatic bed surgery. The mean operation time was 297.2 minutes (range 111–544 minutes). Characterization of the TBS is shown in Table 1. Bleeding occurred in several tissue types and from multiple sources, and the TBS ranged in size from 0.1–200 cm². Concerning bleeding severity, 22 (73.3%) subjects had a TBS categorized as Type 2, and the remaining eight (26.7%) were Type 3. There was a positive correlation between the layers of gauze penetrated with blood and the categorization of bleeding severity determined by the investigators (Table 2).

Effectiveness of Veriset™ hemostatic patch

In 26/30 (86.7%) subjects, conventional methods of obtaining hemostasis were attempted first but were deemed ineffective.

Table 1 Characterization of TBS

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Type of statistic</th>
<th>Veriset™ hemostatic patch (N=30)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source(s) of bleeding</td>
<td>Arterial</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Venous</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Resection/dissection field</td>
<td>n (%)</td>
</tr>
<tr>
<td>Tissue type(s)</td>
<td>Tumor bed</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Organ bed</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Other</td>
<td>n (%)</td>
</tr>
<tr>
<td>Bleeding severity</td>
<td>Type 1</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Type 2</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Type 3</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Type 4</td>
<td>n (%)</td>
</tr>
<tr>
<td>Approximate area of TBS (cm²)</td>
<td>N</td>
<td>30</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>20.26</td>
</tr>
<tr>
<td></td>
<td>Median</td>
<td>6.75</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>38.48</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>0.1–200.0</td>
</tr>
<tr>
<td>Conventional methods used</td>
<td>None</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Cautery</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Clips</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Staples</td>
<td>n (%)</td>
</tr>
<tr>
<td></td>
<td>Suture/ligature</td>
<td>n (%)</td>
</tr>
</tbody>
</table>

Note: Percentages may sum to more than 100% since more than one category may apply.

Abbreviations: TBS, target bleeding site; N, total number of subjects in study; n, number of subjects; SD, standard deviation.
Table 2  Bleeding severity and number of layers of gauze penetrated by blood

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Bleeding severity</th>
<th>Type of statistic</th>
<th>Veriset™ hemostatic patch (N=30)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Layers of gauze that blood penetrated</td>
<td>Type 1</td>
<td>n</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Type 2</td>
<td>n</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>8.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Median</td>
<td>8.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>4.4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>3–21</td>
<td></td>
</tr>
<tr>
<td>Type 3</td>
<td>n</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>20.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Median</td>
<td>16.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>12.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>8–39</td>
<td></td>
</tr>
<tr>
<td>Type 4</td>
<td>n</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: N, total number of subjects in study; n, number of subjects; SD, standard deviation.

and, in the remaining four subjects, the investigator determined that conventional methods were impractical (Table 1). In all of these subjects, Veriset™ hemostatic patch was applied to the TBS. Within 1 minute, 70.0% (21/30) of subjects had achieved hemostasis and, within 5 minutes, 96.7% (29/30) of subjects had achieved hemostasis (Table 3). The median time to hemostasis was 1.0 minute; only three subjects were still bleeding 2.0 minutes after application of the device (Figure 2). The maximum amount of the device required to obtain hemostasis in a single subject was four 5×10 cm² patches. In 27/30 (90.0%) subjects, Veriset™ hemostatic patch remained in place for the duration of the procedure. In one subject, the investigator opted to remove Veriset™ hemostatic patch after 2 minutes because hemostasis had not been achieved, and an alternative topical agent was administered. In another subject, hemostasis was achieved with Veriset™ hemostatic patch but, later in the procedure, it became detached during a gastric pull-up maneuver. No tissue damage was observed in these two subjects. In a third subject, Veriset™ hemostatic patch was later removed when the initial TBS was extracted along with an organ because of a procedural complications unrelated to the use of the device.

Safety of Veriset™ hemostatic patch

AEs were monitored at each follow-up, and device-relatedness was adjudicated by an independent medical monitor. A total of 136 AEs were observed in 23/30 (76.7%) subjects, and all were typical of the types of surgical procedures performed (Table 4). One subject experienced 25 AEs, and two subjects each experienced 13 AEs; however, most of these were only mild or moderate in severity. Anemia, nausea, impaired healing, and pleural effusion were the only AEs to occur in more than 10% of subjects. All but one (99.3%) of the AEs were determined to have no relationship to treatment with Veriset™ hemostatic patch. In one subject with anemia, the AE was adjudicated by the independent medical monitor as having an unknown or impossible to determine relationship with the device. There was a total of 20 SAEs in 11/30 (36.7%) subjects. One subject died before the 7-day follow-up, but the cause, embolism from an atrial thrombus diagnosed prior to surgery, was unrelated to the use of Veriset™ hemostatic patch. No device-related SAEs were observed for 30 days after the surgery, and no reoperations for device-related bleeding complications were performed within 5 postoperative days. The Data Safety Monitoring Board found no safety issues at the conclusion of the study.

Discussion

Many different approaches to achieving hemostasis are currently available; these include conventional methods as well as topical agents. Available methods or devices have advantages and disadvantages that are related to their modes of action or the underlying materials used. Classical surgical techniques, such as sutures, require the presence of mobile adjacent tissue, which will compress the bleeding site by mechanical force but will cause additional trauma to the target area. Electrocautery is effective but causes substantial tissue degradation of an area of the bleeding site and around the focus of application. Topical hemostats have been approved that are either based on coated patch application or that use blood clotting components in a semisolid, fluid-like texture. In certain situations, these existing products exhibit a supportive effect in reaching sufficient hemostasis, but there is still room for improvement in their efficacy, ease of
use, and adherence to anatomical tissue. Oxidized cellulose, a stand-alone hemostat, has been used for several decades in its regenerated form; however, a recent study showed significant improvement using nonregenerated oxidized cellulose in terms of hemostatic efficiency. In contrast to these and other available products, Veriset™ hemostatic patch promotes hemostasis through a dual mode of action that combines the features of two clinically used components: oxidized cellulose and polyethylene glycol hydrogel. Veriset™ hemostatic patch promotes hemostasis by serving as a tamponade to physically stem blood flow while concentrating platelets and other clotting factors at the bleeding site to accelerate the coagulation cascade.

In this study, a variety of procedure types were performed that were accompanied by soft tissue bleeding from multiple sources. Veriset™ hemostatic patch stopped blood loss in almost all of these situations, a finding that supports its potential broad range of use. Veriset™ hemostatic patch was deemed ineffective in only one subject whose bleeding site was determined to have an unusual geometry that did not allow sufficient contact with the device. Multiple applications of Veriset™ hemostatic patch are feasible to accommodate either especially large bleeding sites or more difficult to treat bleeding sites. Several subjects in this study required multiple applications, but this did not correlate with adverse safety events.

Figure 2 Kaplan–Meier analysis of time to hemostasis.

Notes: After the application of Veriset™ hemostatic patch, hemostasis was assessed every 30 seconds for the first 5 minutes and every 1 minute for minutes 5–10. All subjects achieved hemostasis within 6.0 minutes, and only three subjects required longer than 2.0 minutes to achieve hemostasis.

Table 4 Adverse events

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Type of statistic</th>
<th>Veriset™ hemostatic patch (N=30)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adverse events</td>
<td>n (%)</td>
<td>23 (76.7)</td>
</tr>
<tr>
<td>Serious adverse events</td>
<td>n (%)</td>
<td>11 (36.7)</td>
</tr>
<tr>
<td>Device-related adverse events</td>
<td>n (%)</td>
<td>1 (3.3)</td>
</tr>
<tr>
<td>Unanticipated adverse device effects</td>
<td>n (%)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>Device-related serious adverse events per subject during the 30 days after surgery</td>
<td>n/N (%)</td>
<td>0/30 (0.0)</td>
</tr>
<tr>
<td>Reoperation for device-related bleeding complications during the 5 days after surgery</td>
<td>n/N (%)</td>
<td>0/30 (0.0)</td>
</tr>
</tbody>
</table>

Note: Device-related adverse events are events with a definite, probable, possible, unknown/impossible to determine, or missing relationship to the device.

Abbreviations: n, number of subjects; N, total number of subjects in study.
to the surgical procedure and minimal risk of complications. In the two cases in which Veriset™ hemostatic patch either did not adhere or became detached after application, there was no visible tissue damage. This observation suggests that, unlike some conventional hemostatic methods that can cause tissue damage (ie, electrocautery), the use of Veriset™ hemostatic patch can be attempted with little risk to the patient.

Conclusion
The results shown here support the use of Veriset™ hemostatic patch across a variety of surgical procedures that can result in soft tissue bleeding. Additionally, the median time to hemostasis in this study was 1 minute, the same as that observed in the randomized liver study; this similarity indicates that Veriset™ hemostatic patch might offer consistent efficacy across multiple tissues. On a practical level, the device exhibits properties that render it convenient and widely applicable. Further studies, particularly in the form of randomized, controlled trials, should be performed in the future to assess such characteristics as safety, efficacy, usability (eg, adhesiveness to tissue, flexibility to underlying structures), and cost-effectiveness, compared to other topical hemostats on the market. While these data would be beneficial to the surgical community, our results combined with the previously published liver results, suggest that Veriset™ hemostatic patch offers a safe hemostatic method that is effective in many surgical situations.

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Disclosure
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References