New bipolar tissue ligator combines constant tissue compression and temperature guidance: histologic study and implications for treatment of hemorrhoids

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Background: Several minimally invasive technologies are available to treat common soft tissue lesions including symptomatic hemorrhoids. The use of energy to deliver heat and coagulate target lesions is commonly practiced. This study compares the histologic effects produced on intestinal tissues by two energy-based systems which employ different approaches of heat delivery.

Methods: Two heat delivery systems were evaluated in vivo in a single porcine subject: infrared coagulator and bipolar tissue ligator utilizing constant tissue compression and temperature guidance. Eighteen treatment sites divided into three groups of six were assessed. Treatment site temperature was measured and the effects of thermal treatment in the mucosa, submucosa, submucosal vessels, and muscularis layer were scored. Lateral thermal spread beyond the energy application site was also assessed.

Results: Treatment site temperatures were much lower in the bipolar ligator group than in the infrared coagulator group. The mucosal and submucosal tissue changes observed in tissues treated with infrared energy and bipolar energy at 55°C were similar. Both the mucosal and submucosal tissue changes with bipolar energy at 50°C were significantly less.

Conclusion: Both devices achieved similar histologic results. However, the unique design of the bipolar ligator, which allows consistent capture, constant compression, and temperature monitoring of target tissue, accomplished the desired histologic changes with less muscular damage at much lower temperatures than the infrared coagulator. The use of bipolar ligation could offer clinical advantages such as reduced patient pain and a minimized chance of heat-related collateral tissue damage.

Keywords: bipolar ligator, internal hemorrhoids, tissue manipulation, ligation

Introduction
Symptomatic Grade 1 and 2 hemorrhoids rarely require surgical intervention and are usually treated with less invasive technologies – most frequently with rubber band ligation or infrared coagulation. Rubber band ligation is associated with frequent post-procedural pain and discomfort,1–6 as well as other, sometimes very serious, complications.7–14 Infrared coagulation causes much fewer complications than rubber band ligation, but produces inconsistent clinical results and is frequently ineffective.2,6,15–19 The purpose of this study was to compare histologic findings between the Infrared Coagulator (IRC 2100, Redfield Corporation, Rochelle Park, NJ, USA) and the HET™ Bipolar Ligator (HET Systems, Northvale, NJ, USA) – a new bipolar device for tissue manipulation and coagulation.
Technology and methods

The HET™ Bipolar Ligator System is comprised of an innovative tissue ligator and an associated temperature monitor. The HET™ Bipolar Ligator has a unique, constant tissue compression mechanism, a temperature sensor adjacent to the bipolar electrodes, and integral LED-based illumination. The HET™ Bipolar Ligator’s unique tissue clamp and bipolar radio-frequency-based tissue coagulation allow compression and ligation of the target tissue with a constant force and predictable energy delivery. When this operator-independent force is applied parallel to the rectal wall, the associated superior hemorrhoid vasculature in the formed tissue fold is predictably occluded and ligated (Figure 1). Continuous tissue temperature monitoring during use of the HET™ Bipolar Ligator provides the operator with objective, real-time feedback about the treated tissue temperature.

The Redfield IRC 2100 includes a halogen lamp-based heat generator and an elongated probe, which interfaces with the target tissue and delivers heat by regulating the amount of time the device is activated. The manufacturer provides predetermined recommended time intervals for treatment. This approach does not control the amount of heat applied to the tissue; however, it controls the length of time the tissue will be exposed to varying temperatures. The IRC system does not provide a mechanism for controlling or gauging tissue compression, or a method for measuring the temperature of the treated tissue. Additionally, the pressure applied to the treatment site by the IRC device is operator dependent.

An in vivo single animal, multiple treatment sites study was conducted to compare the histologic effects of the two systems when used on the recto-sigmoid swine colon.

The objective of this study was to assess histopathologic changes following treatment with heat delivered to normal colonic tissues by the two devices. Since both the Redfield IRC 2000 and the HET™ Bipolar Ligator are applied to the normal colonic mucosa of humans during the treatment of internal hemorrhoids, and since the hemorrhoidal tissue is not directly affected by thermal energy, only normal animal colonic tissues were studied. The effects of thermal treatment in the mucosa, submucosa, submucosal vessels, and muscularis layer were evaluated. Lateral thermal spread beyond the energy application site was also assessed.

Following receipt of University Institutional Animal Care and Use Committee approval, the study was carried out on a single female Sus scrofa pig weighing approximately 54.5 kg. Preanesthesia of intramuscular ketamine (14.7 mg/kg) and intravenous acepromazine (1.5 mg/kg) was given. Surgical plane was maintained by inhalation of a 3% isoflurane solution. The pig was connected to a heart rate monitor, oxygen monitor, and ventilator. A midline abdominal incision was performed, and the sigmoid colon was exposed. A longitudinal incision was made on the antimesenteric side of the recto-sigmoid colon and wide access to the colonic mucosa was obtained.

Three similar segments of the colon were used to study three treatment groups with six areas per group. The HET™ Bipolar Ligator was used to treat two groups at two different temperatures: 55°C (HET55 group) and 50°C (HET50 group). The IRC device was used to treat one group (IRC group).

Both the IRC 2100 (IRC) and HET™ Bipolar Ligator (HET) devices were used according to the manufacturer’s instructions for use. The HET™ Bipolar Ligator was connected to a Conmed Hyfrecator 2000 bipolar radiofrequency generator (Conmed, Utica, NY, USA) with an output coagulation setting at 10 watts.

The tissue in the HET™ Bipolar Ligator was compressed parallel to the bowel wall, and bipolar radiofrequency energy was then delivered until the temperature reached 50°C (across six areas) or 55°C (across six areas).

The IRC device delivers pulses of infrared light through a small contact tip applicator that is applied to the tissue.

Figure 1 Gross comparative mucosal changes in three rows of applications: IRC (IR) and HET with final tissue temperatures of 50°C and 55°C.

Abbreviations: IRC, infrared coagulator; HET, HET Bipolar Ligator.
The light causes thermal coagulation which results in tissue necrosis. The amount of energy delivered is regulated by the amount of time the device is activated. As recommended by the manufacturer, the IRC timer was set in between 1.0–1.5 seconds for the treatment of hemorrhoids. The IRC heat pulse was delivered perpendicular to the mucosal surface to each of the six treatment mucosal areas. The temperature of the IRC probe was continuously measured during the heat delivery using a digital thermometer (Tektronix DTM920, Tektronix, Inc, Beaverton, Oregon, USA). The maximum temperature, observed in all cases at 1.5 seconds, was documented.

When all treatment groups were completed, the pig was euthanized, and the recto-sigmoid colon segment removed. The excised segment was placed in 10% buffered formalin for 72 hours.

All samples were sectioned perpendicular to the mucosa into two pieces and embedded in paraffin blocks. The blocks were sectioned in 4 µ cuts, and an additional step was performed approximately 150 µ from the first cut. All slides were stained with hematoxylin and eosin.

The thermal damage in the mucosa, submucosa, and muscularis layers was measured as follows: layer thinning was measured by an ocular micrometer comparing the treated area to an adjacent normal area, and was graded on a scale of 0 to +4 with 0 representing 0% thinning, +1 representing 25% thinning, +2 representing 50% thinning, +3 representing 75% thinning, and +4 representing total loss of the layer. Multiple areas were analyzed for each of the coagulation areas. Absolute dimensions were not used as the layer thickness can vary from place to place, and such measurements would not reflect the importance of the percent of layer damage. The percentage measurements used are a standard technique in assessing histological damage, including thermal damage.

The mucosa was graded based on the layer thinning, loss of glands and loss of basophilic staining. The submucosa was graded based on the layer thinning and the degree of the vascular injury. The muscularis layer was graded based on the increased eosinophilia, vacuolization and nuclear changes. Averages and standard error of the mean scores were calculated. The values were compared with the standard Students t-test. Collagen damaged was accessed by polarized light microscopy.

In the HET50 and HET55 groups, the treated tissues were also examined at different distances from the bipolar electrodes. The lateral thermal spread outside of the energy application site was evaluated as well.

**Results**

The treatment time and treatment site temperature readings are summarized by study group in Table 1. The treated tissue reached 50°C in 4–7 sec (HET50 group) and 55°C in 6–9 sec (HET55 group). The compressed tissue in both HET groups was 1–2 mm.

In the IRC group, after 1.5 seconds of treatment, the temperature at the tip of the probe varied from 127.8°C to 155.7°C (149.9°C ± 11.1°C); with the final tissue temperature being highly dependent on the pressure applied by the device to the tissue. The gross, comparative changes between the IRC, HET55, and HET50 groups are shown in Figure 1.

Table 2 summarizes the comparative mucosal and submucosal changes between the IRC, HET55, and HET50 groups at sites immediately adjacent to or in contact with the treatment elements. The mucosal and submucosal tissue changes seen in the IRC and the HET55 groups were similar. Both the mucosal and submucosal tissue changes in the HET50 group were significantly milder than IRC and HET55 groups ($P < 0.05$). None of the samples displayed loss of tissue architecture or thermal damage of the full thickness of the bowel wall. The lateral thermal spread from the application site into the normal tissue in all cases was minimal.

**Mucosa layer**

The treated mucosa displayed thinning caused by thermal desiccation and the loss of crypt glands (Figures 2 and 3). Some thinning in the HET specimens was also a result of compression from the tissue clamp of the device. The basophilic staining of the layer was also decreased (Figure 4). These effects, with the exception of glandular loss, were significantly less pronounced in the HET50 group. Some vascular congestion and small hemorrhaging were seen but were not prevalent.

**Submucosa layer**

Submucosal injury was characterized by thinning and vascular injury. The submucosal thinning (Figure 5) was

<table>
<thead>
<tr>
<th>Group</th>
<th># treatment areas</th>
<th>Treatment site temperature (°C)</th>
<th>Treatment time (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HET50</td>
<td>6</td>
<td>50</td>
<td>4–7 (4.8 ± 0.8 sec)</td>
</tr>
<tr>
<td>HET55</td>
<td>6</td>
<td>55</td>
<td>6–9 (6.2 ± 1.8 sec)</td>
</tr>
<tr>
<td>IRC</td>
<td>6</td>
<td>127.8–155.7 (149.9 ± 11.1)</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Abbreviations: HET50, HET Bipolar Ligator at 50°C; HET55, HET Bipolar Ligator at 55°C; IRC, infrared coagulation.
similar between the IRC and the HET55 groups, while the HET50 group demonstrated significantly less submucosal thinning ($P < 0.05$). In the HET55 group, 50% of samples displayed a full thickness submucosal injury without muscle involvement, while none of the HET50 samples showed full thickness submucosal thermal injury.

Vascular injury included congestion and small amounts of hemorrhage, and in some cases, the vessel endothelial cells were affected (Figure 6). Small amounts of reactive neutrophilic infiltration were seen in most samples. Vascular changes were similar between the HET55 and HET50 groups.

There were signs of collagen denaturation present in all IRC, HET55, and HET50 specimens with all samples showing reduced collagen damage in the areas that were more remote from the source of the heat. This was demonstrated by the polarized photomicrograph showing the normal collagen as bright areas and the denatured as darkened areas (Figure 7).

The maximal tissue damage was observed immediately adjacent to the ligation electrodes in the HET55 and HET50 groups. While hemorrhage and occasional thinning were observed in the mucosa, the thermal effects were prevalent in the submucosa and presented as vascular injury including congestion and hemorrhage. Evaluation of folds of tissue between the ligation electrodes demonstrated no difference in vascular damage between the two HET groups (Table 3).

### Muscularis layer

All samples from the IRC group displayed similar thermal damage to the muscularis layer, but this damage was scant

**Table 2** Comparison between mucosal and submucosal changes at treatment sites

<table>
<thead>
<tr>
<th>N (# samples)</th>
<th>IRC</th>
<th>HET55</th>
<th>HET50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mucosa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Layer thinning</td>
<td>$1.70 \pm 0.14$</td>
<td>$2.00 \pm 0.18$</td>
<td>$0.56 \pm 0.18$</td>
</tr>
<tr>
<td>Loss of glands</td>
<td>$2.20 \pm 0.09$</td>
<td>$1.50 \pm 0.13$</td>
<td>$1.88 \pm 0.09$</td>
</tr>
<tr>
<td>Loss of basophilic staining</td>
<td>$1.60 \pm 0.11$</td>
<td>$1.62 \pm 0.11$</td>
<td>$1.12 \pm 0.09$</td>
</tr>
<tr>
<td>Submucosa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Layer thinning</td>
<td>$1.50 \pm 0.10$</td>
<td>$2.00 \pm 0.26$</td>
<td>$0.25 \pm 0.17$</td>
</tr>
<tr>
<td>Degree of vascular injury</td>
<td>$2.40 \pm 0.18$</td>
<td>$2.00 \pm 0.26$</td>
<td>$1.25 \pm 0.11$</td>
</tr>
</tbody>
</table>

**Note:** Tissue damage scores graded 0 to 4 (avg ± SD).

**Abbreviations:** IRC, infrared coagulator; HET55, HET Bipolar Ligat or at 55°C; HET50, HET Bipolar Ligat or at 50°C; SD, standard deviation.
and isolated. In the HET50 and HET55 groups, 75% of samples had no muscular injury, while 25% of the samples presented only with superficial muscularis injury (Figure 8).

Lateral thermal spread from the application site into normal tissue was minimal in all samples.

**Discussion**

The goal of therapeutic intervention is to effectively target the pathologic lesion while minimizing injury to collateral tissues. Using energy to treat gastrointestinal lesions, including hemorrhoids, is not new. Electrococagulation as a treatment modality was utilized as early as 1867 and explained by Dr. WE Kessey in 1934. Over the past several decades, new developments and techniques to treat hemorrhoids in a less invasive manner have targeted the superior hemorrhoidal vasculature and the associated submucosa above the dentate line in order to minimize patient pain, discomfort, and tenesmuses.

Both the IRC device and the HET™ Bipolar Ligator intend to produce histologic changes proximal to the internal hemorrhoid, which would lead to an occlusion of the superior hemorrhoid blood supply and formation of moderate scarring in the associated submucosa. To minimize potential collateral tissue damage, a precise capture of the target tissue proximal to the internal hemorrhoid, and predictable energy delivery to both the submucosal vessels and the surrounding connective tissue, are thus highly desirable.

In this evaluation there was greater variability in the temperature at the tissue treatment site in the IRC group than in the HET groups, and the temperature reached in the IRC treatment group was significantly higher. With the IRC device, the procedure was driven by the manufacturer recommended time

**Table 3** Comparison of submucosal changes between HET55 and HET50 groups in the treated tissue folds beyond immediate contact with the ligation electrodes

<table>
<thead>
<tr>
<th></th>
<th>HET55 (Avg ± SE)</th>
<th>HET50 (Avg ± SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of samples</td>
<td>16</td>
<td>16</td>
</tr>
<tr>
<td>Submucosa vascular damage</td>
<td>1.75 ± 0.11</td>
<td>1.5 ± 0.13</td>
</tr>
</tbody>
</table>

**Note:** Graded from 0 to 4; avg ± SD.

**Abbreviations:** HET55, HET Bipolar Ligator at 55°C; HET50, HET Bipolar Ligator at 50°C; SE, standard error.
setting and operator variability in compression pressure. Both inevitably impacted the amount of energy delivered to the target site. In contrast, when the HET Bipolar Ligator was used, temperature at the treatment site was monitored continuously, and energy was delivered consistently until the target tissue temperature of 50°C–55°C was achieved. The time of energy delivery during tissue ligation using the HET Bipolar Ligator is a derivative of the targeted tissue temperature. Additionally, the device is designed to provide constant consistent compression, which further minimized variability in both the energy delivery to the tissue and consequent tissue changes.

Tissue compression

Tissue compression is a known factor affecting energy delivery to targeted tissue. Variability in the tissue compression, therefore, would lead to variability in the energy delivery to the tissue. This was demonstrated in this investigation given that the consistently compressed tissue (among the HET groups) required much less temperature than non-compressed tissue (among the IRC group) to cause similar histologic effects.

The IRC device does not provide a mechanism for predictable or measurable tissue compression, meaning that the tissue compression was operator-dependent. Contrary and advantageous, the minimal variability observed within the HET treatment groups was a function of the unique design of the HET Bipolar Ligator device.

Tissue temperature

The amount of energy delivered to the tissue correlates with the histological changes. The tissue temperature monitoring, therefore, may serve as a useful guideline during the energy delivery to the tissue.

The IRC device did not offer a mechanism for temperature monitoring and relied strictly on the time of energy delivery for tissue treatment. As information on temperature generated by the IRC probe was not found in the literature, it was measured directly in this study. The IRC generated 149.9°C ± 11.1°C during 1.5 seconds of activation in this hemorrhoid treatment model. In contrast, the HET Bipolar Ligator was designed to allow temperature monitoring of the treated tissue. This affords standardization of the treatment by specifically exposing the target tissue to a constant 55°C temperature to achieve the desired therapeutic effect.

Both devices created similar histological changes in the submucosal vessels and submucosa layer of the target tissues. The IRC achieved these changes in 1.5 seconds at much higher temperature, while the HET Bipolar Ligator achieved the desired tissue changes using temperatures that were 3.0 and 2.7 times lower in HET50 and HET55 groups, respectively. It appears that the constant tissue compression accompanying the tissue ligation using the HET Bipolar Ligator reduced the amount of energy required to produce similar histological effects. Recent discovery of targeted metal nanoparticles in selective tissue hyperthermia and cytotoxicity may play an interesting role in facilitating a minimally-invasive chemical ligation of the targeted small vessels.

Tissue desiccation

In the IRC and both HET groups, signs of tissue desiccation were observed, including some loss of intercellular water and associated pyknotic and hyperchromic changes in the nuclei. Changes such as loss of normal tissue architecture and complete cellular protein denaturation were not observed in any of the samples.

Histologically, cells showed lighter hematoxylin and eosin staining due to the loss of cell nuclei and intracellular protein; such cells are sometimes referred to as ghost cells. White cell infiltration and capillary growth into the damaged area will occur from the surrounding normal tissue. This regenerative process will also lead to fibrosis. In contrast, a more significant heat-related injury typically produces more significant volume of tissue damage, and the adjacent tissue would be unable to regenerate, usually leading to ulceration.

Comparing HET bipolar ligator at 55°C and 50°C

While the overall histological changes created by the tissue ligation using the HET Bipolar Ligator at 50°C (HET50)
appeared to be milder than the ligation at 55°C (HET55), the submucosal changes in the treated folds of tissue between the ligation electrodes were substantially equivalent between the groups.

The observed vascular injury was consistent with vascular occlusion in both groups. This finding suggests that the efficacy of the HET™ device operating at 50°C and 55°C is similar. However, from a clinical standpoint, patients exposed to a lower temperature (50°C) may experience less discomfort.

The thermal injury in all HET™ specimens was limited to the areas compressed between the bipolar electrodes with no or minimal collateral tissue damage outside of the tissue that was in contact with the electrodes. Since the operator controls the position of the electrodes, the tissue compression within the unique HET™ clamp is constant, and therefore the tissue is treated to the defined temperature. The energy delivery to the target tissue by the HET™ Bipolar Ligator is highly consistent and predictable, minimizing collateral tissue damage. The demonstrated recognizable architecture, the lack of full thickness damage, the intact tunica muscularis, and the minimally damaged blood supply suggest that little ulceration would occur in tissues treated with the HET™ Bipolar Ligator at 55°C or 50°C. The intact residual blood supply would thus promote routine fibroblast infiltration and active fibrosis in the area of the thermal desiccation.36–38

**Conclusion**

The combination of constant tissue compression and temperature-guided energy delivery achieves desirable histological changes between 50°C and 55°C.

The unique design of the HET™ Bipolar Ligator provides precise delivery of energy to the target tissue, as well as real-time temperature monitoring and well-gauged, constant tissue compression. These mechanisms facilitate accurate therapeutic energy delivery to the target tissues between 50°C and 55°C while minimizing the collateral muscular damage.

In this comparative histological study, while both devices achieved similar results, the HET™ Bipolar Ligator achieved the desired histologic changes with less muscular damage, using focused delivery of radio frequency energy at a much lower temperature than IRC. Therefore, the use of the HET™ Bipolar Ligator could provide significant clinical advantages, potentially enhance patients’ tolerance of the ligation procedure, and minimize the chance of heat-related complications and collateral tissue damage. The conclusions of this study need to be considered with caution due to the anatomical differences between humans and pigs. Specifically, the differences in colonic layer thickness, tissue metabolism, and reactions to heat between human and pig tissue could potentially influence the study results. To make clinically meaningful conclusions, it is important to continue studying these new devices in a well-designed, outcome-based clinical study. Comparative clinical evaluations of the devices are planned to confirm these outcomes.

**Disclosure**

Gregory Piskun, MD is an inventor of HET™ Bipolar Ligator and founder of HET Systems. Robert Tucker reports no conflicts of interest in this work.

**References**