The hydrogel sealant is naturally absorbed in approximately 4 to 8 weeks.

The DuraSeal Spine Sealant System is provided in two configurations. The 2 mL configuration consists of one 2 mL polymer kit and one MicroMyst Applicator (the MicroMyst Applicator requires the use of a compressed air source, such as the Confluent Surgical Flow Regulator or the Confluent Surgical Air Pump). The 5 mL configuration consists of one 5 mL polymer kit which includes the Dual Liquid Applicator (consisting of the Y-Applicator and three (3) Spray Tips). The polymer kits and applicators are provided sterile.

**INDICATION**
The DuraSeal Spine Sealant System is indicated for use as an adjunct to sutured dural repair during spinal surgery to provide watertight closure.

**CONTRAINDICATIONS**
Do not apply the DuraSeal hydrogel to confined bony structures where nerves are present since neural compression may result due to hydrogel swelling. The hydrogel may swell up to 50% of its size in any dimension.

**WARNINGS**
- Do not use if an active infection is present at the surgical site.
- Do not use the DuraSeal Spine Sealant as a hemostatic agent.
- The safety and effectiveness of the DuraSeal hydrogel has not been studied in:
  - Patients with a known allergy to FD&C Blue #1 dye.
  - Procedures involving non-autologous duraplasty.
  - Patients with severely altered renal or hepatic function.
  - Patients with a compromised immune system or autoimmune disease.
- Use only with the Confluent Surgical applicators.
- The polymer kits and applicators are provided sterile. Do not use if packaging or seal has been damaged or opened. Do not re-sterilize.
- The polymer kits and applicators are intended for single patient use only. Discard opened and unused product.
- Do not use if the PEG powder is not free flowing.
- Use within 1 hour of preparation.
- Do not use in combination with other sealants or hemostatic agents.
- Do not use in patients younger than 18 years of age, or in pregnant or breast feeding females.
- Prior to application of the hydrogel, ensure that adequate hemostasis has been achieved.
- Incidental application of hydrogel to tissue planes that will be subsequently approximated, such as muscle and skin, should be avoided.

**ADVERSE EVENTS**
The DuraSeal Spine Sealant System was evaluated in a pivotal clinical study, in which a total of 158 patients were enrolled (102 treated with DuraSeal Spine Sealant and 56 patients treated using Standard of Care methods). All Adverse Events were coded using the Medical Dictionary for Regulatory Activities (MedDRA) and are presented based on System Organ Class.

The DuraSeal Spine Sealant System consists of components for preparation of a synthetic absorbable sealant, and applicators for delivery of the sealant to the target site.

The sealant is composed of two solutions, a polyethylene glycol (PEG) ester solution and a trilysine amine solution (referred to as the "blue" and "clear" precursors, respectively). When mixed together, the precursors cross-link to form the hydrogel sealant. The mixing of the precursors is accomplished as the materials exit the tip of the applicator.
were treated with the DuraSeal Spine Sealant, and a total of 56 was conducted. Subjects that were scheduled for spinal procedures
requiring a dural incision and who met the preoperative eligibility criteria were considered for study participation. Subjects that met all of the intra-operative eligibility criteria were enrolled and randomized either to DuraSeal Spine Sealant or Control. The study involved 24 investigational sites within the United States. A total of 102 patients were treated using Standard of Care methods to obtain a watertight dural closure. Standard of care consisted of sutures, fibrin sealant, soft tissue patch/graft or a combination of the materials.

The primary endpoint for this study was the percent (%) success in obtaining a watertight closure following assigned treatment (DuraSeal Spine Sealant or Control), defined as a watertight closure of the dural repair intra-operatively confirmed by Valsalva maneuver at 20 - 25 cm H2O for 5 to 10 seconds.

Safety was assessed based on evaluation of the occurrence of postoperative CSF leaks and surgical site infection, the nature and severity of adverse events, protocol-specified laboratory tests, neurological assessments, and wound healing.

Inclusion/Exclusion criteria for the study included the following:

Pre-Operative Inclusion Criteria:
- Subject was between 18 and 75 years of age.
- Subject was scheduled for a spinal procedure that entails a dural incision.
- Subject required a procedure involving surgical wound classification Class IClean (per CDC criteria).
- Subject, or authorized representative, was informed of the nature of the study, and provided written informed consent, approved by the appropriate Institutional Review Board (IRB) of the respective clinical site.

Pre-Operative Exclusion Criteria:
- Subject had active spinal and/or systemic infection.
- Subject required additional spine surgery within the study time period.
- Subject had a previous spinal surgery involving dural exposure and/or entry at the same level(s) as the study procedure.
- Subject had pre-existing external lumbar CSF drain or internal CSF shunt.
- Subject participated in a clinical trial of another investigational device or drug.
- Subject with creatinine > 2.0 mg/dL.
- Subject with total bilirubin > 2.5 mg/dL.
- Pregnant or breast-feeding females or females who wished to become pregnant during the length of study participation.
- Subject treated with chronic steroid therapy unless discontinued more than 6 weeks prior to surgery (standard peri-operative steroids are permitted). For purposes of this protocol, chronic steroid therapy is defined as greater than 4 weeks.
- Subject had documented history of significant coagulopathy with a PT > 35 sec, PTT/INR > 1.2, receiving aspirin, or NSAIDS at the time of surgery. Note: Subjects who are receiving cardiovascular prophylaxis are not excluded.
- Subject received warfarin or heparin at the time of surgery (including anticoagulants).
- Subject diagnosed and documented compromise immune system and/or autoimmune disease.
- Subject had chemotherapy treatment within 6 months prior to, or planned during the study (until completion of last follow-up evaluation).
- Subject had prior radiation treatment to the surgical site or has planned radiation therapy within 30 days post procedure.
- Subject had a known malignancy or another condition with prognosis shorter than 8 months.
- Subjects with documented history of uncontrolled diabetes.
- The investigator determined that the subject should not be included in the study for reason(s) not already specified.

Intra-Operative Inclusion Criteria:
- Presence of non-watertight dural closure, either spontaneously or upon Valsalva maneuver to 20 - 25 cm H2O for 5-10 seconds.

| Potential, but not observed, risks and adverse events that could occur from the use of the hydrogel include, but are not limited to, renal compromise, inflammatory reaction, neurological compromise, allergic reaction and/or delayed healing. |

| CLINICAL EXPERIENCE |

A prospective, multi-center, randomized, two arm, single blind study designed to assess if the DuraSeal Spine Sealant System, when used as an adjunct to sutured dural repair, is more effective than Standard of Care methods for producing a watertight dural closure in subjects undergoing an intentional durotomy during spinal surgery was conducted. Subjects that were scheduled for spinal procedures requiring a dural incision and who met the preoperative eligibility criteria were considered for study participation. Subjects that met all of the intra-operative eligibility criteria were enrolled and randomized either to DuraSeal Spine Sealant or Control. The study involved 24 investigational sites within the United States. A total of 102 patients were treated with the DuraSeal Spine Sealant, and a total of 56 patients were treated using Standard of Care methods to obtain a watertight dural closure. Standard of care consisted of sutures, fibrin sealant, soft tissue patch/graft or a combination of the materials. |
Intra-Operative Exclusion Criteria:
- Incidental finding of any of the pre-operative exclusion criteria.
- Subject required use of a synthetic or non-autologous dural grafting material.
- Subject had a gap of greater than 2 mm following primary dural closure.
- Subject had undergone laminoplasty decompression.
- Subject had undergone a syringomyelia procedure where the shunt is not placed in the subarachnoid position.
- Subject had undergone a Chiari Malformation procedure that did not entail a dural incision at or below the C1 level.
- Investigator determined that participation in the study may jeopardize the safety or welfare of the subject.

Demographic information for patients treated in the study is shown in the table below:

<table>
<thead>
<tr>
<th>Demographic Information</th>
<th>Spinal Sealant (ITT)</th>
<th>Control (ITT)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>Mean (SD)</td>
<td>47.7 (13.65)</td>
</tr>
<tr>
<td></td>
<td>Range (min, max)</td>
<td>18.3 (74.5)</td>
</tr>
<tr>
<td>Gender, n(%)</td>
<td>Male</td>
<td>54 (52.9)</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>30 (29.6)</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>Mean(SD)</td>
<td>169.9 (11.74)</td>
</tr>
<tr>
<td></td>
<td>Range (min, max)</td>
<td>132.1 (153.0)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>Mean (SD)</td>
<td>60.0 (20.62)</td>
</tr>
<tr>
<td></td>
<td>Range (min, max)</td>
<td>45.7 (147.4)</td>
</tr>
<tr>
<td>BMI (Kg/m²)</td>
<td>Mean (SD)</td>
<td>27.6 (8.00)</td>
</tr>
<tr>
<td></td>
<td>Range (min, max)</td>
<td>17.9 (24.2)</td>
</tr>
<tr>
<td>Smoking Status, n(%)</td>
<td>Never</td>
<td>62 (60.8)</td>
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<tr>
<td></td>
<td>History</td>
<td>47 (44.2)</td>
</tr>
<tr>
<td></td>
<td>Current</td>
<td>21 (20.6)</td>
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<tr>
<td>ASA Score, n(%)</td>
<td>I</td>
<td>13 (12.7)</td>
</tr>
<tr>
<td></td>
<td>II</td>
<td>66 (64.7)</td>
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<tr>
<td></td>
<td>III</td>
<td>22 (21.6)</td>
</tr>
<tr>
<td></td>
<td>IV</td>
<td>1 (1.0)</td>
</tr>
<tr>
<td>History of peeled A-V</td>
<td>Malignancy</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td></td>
<td>Tethered cord</td>
<td>1 (1.0)</td>
</tr>
<tr>
<td></td>
<td>Tumor removal</td>
<td>64 (62.7)</td>
</tr>
</tbody>
</table>

Of the 158 subjects randomized, all 102 subjects (100.0%) treated with the DuraSeal Spine Sealant and 36 of the 56 subjects (64.3%) treated with Standard of Care methods displayed a watertight dural closure after assigned treatment, intra-operatively. Three (3) subjects randomized to Standard of Care were considered not evaluable for the per protocol analysis of the primary effectiveness endpoint, as the investigator chose not to use any of the Standard of Care methods per the protocol (i.e., devices designed to provide an intra-operative watertight closure).

Within the Control group, 35 subjects (62.5%) had a watertight closure upon Valsalva following the first Standard of Care application while 21 subjects had a non-watertight closure. For the ITT population, the majority of surgeons chose either sutures (37.5%) or Fibrin Sealant (44.6%) as the treatment method for first attempt.

A second attempt with Standard of Care methods was attempted in 4 of the 21 subjects, at which time only 1 subject achieved a watertight closure upon second post-treatment Valsalva. Per study protocol investigators were required to use the same standard of care technique for both attempts if a second attempt at dural closure was performed in the control cohort. Surgeons were allowed to use additional adjunctive or rescue therapy after the primary endpoint analysis to ensure that all subjects had a watertight dural closure.

The number and types of adverse events observed in both of the study treatment groups were anticipated, given the medical conditions of the enrolled subjects and nature of the complex neurosurgical procedures performed. There were no deaths or unanticipated adverse device effects observed in the study.

The incidence of protocol defined post-operative CSF leaks was comparable between the two treatment groups (7.8% vs. 5.4%). The number of adjunctive therapies used in the Control subjects, following determination that subjects were a primary effectiveness endpoint failure, was greater. In nineteen Control subjects the primary dural repair was reinforced with buttressing materials such as synthetic dura patches or direct dural overlay or an absorbable sponge. The number of adjunctive therapies used in the DuraSeal Spine Sealant vs. Control group, respectively, p=1.00. Overall, there were no clinically relevant differences in safety outcomes between the two treatment groups (DuraSeal Spine Sealant vs. Control) with respect to laboratory evaluations, neurological exams, vital signs, physical examination and wound healing.

In evaluation of the neurological assessment data and neurological complications, there is no indication of symptom complexes consistent with nerve root compression for subjects treated with the DuraSeal Spine Sealant, a potential concern when using hydrogel-based devices along the nerve roots. The data are consistent with the preclinical evaluation performed in a canine cauda equina discectomy model in which the DuraSeal Spine material (DuraSeal Sealant) was applied following lumbar discectomy, with exposure and abrasion of the lumbar nerve roots. In this severe model there were no significant neurological deficits noted and no adverse reactions were macroscopically observed for any of the dural sealant treated sites.
DIRECTIONS FOR USE

Device Preparation – 2 mL Configuration
REF 20-4300

A. Preparing the Blue Precursor

1. Remove the polymer kit tray and the MicroMyst Applicator from their respective outer pouches and introduce into the sterile field.
2. Remove lid from polymer kit tray.
3. Remove and discard syringe cap from Diluent Syringe (blue label).
4. Depress the threaded fitting of the vial cap (Figure 1).
5. Ensure red line is no longer visible (Figure 2).
6. Screw the Diluent Syringe (Blue Label) on to the Powder Vial and inject syringe contents into the vial (Figure 3).
7. Gently shake the vial/syringe assembly until the powder is completely dissolved (Figure 4).
8. Invert the vial/syringe assembly, and draw the vial contents back into the syringe (Figure 5).
9. Unscrew the syringe from the vial and discard the vial.
10. Remove syringe cap from Clear Precursor Syringe.
11. Remove excess air from both syringes
12. Ensure that the precursor volume in each syringe is equal.

B. Assembling the MicroMyst Applicator
Refer to Instructions for Use provided with the MicroMyst Applicator

Device Preparation – 5 mL Configuration
REF 20-4004

A. Preparing the Blue Precursor

13. Remove the polymer kit tray from its outer pouch and introduce into the sterile field.
14. Remove lid from polymer kit tray.
15. Remove and discard syringe cap from Diluent Syringe (blue label).
16. Attach the Diluent Syringe to the Powder Vial.
17. Without depressing the syringe plunger, pierce the vial seal until it is fully depressed (twisting is not required). The entire threaded portion of the vial cap should be depressed below the level of the surrounding plastic vial rim.
18. Inject syringe contents into the vial.
19. Gently shake the vial/syringe assembly until the powder is completely dissolved.
20. Invert the vial/syringe assembly, and draw the vial contents back into the syringe.
21. Unscrew the syringe from the vial and discard the vial.
22. Remove syringe cap from Clear Precursor Syringe.
23. Ensure that the precursor volume in each syringe is equal.

B. Assembling the Dual Liquid Applicator

1. Attach the Clear and Blue precursor syringes to the Y-Applicator.

2. Attach the Syringe Holder (A) to syringe barrels and the Plunger Cap (B) to syringe plungers.

Note:
Avoid touching the plunger cap before application to avoid inadvertent precursor injection and tip plugging.

3. Attach one Spray Tip to the Y-Applicator.

LCN - XXXX
Hydrogel Application
REF 20-4300, REF 20-4004

Note:

- Achieve hemostasis and minimize fluid (CSF, blood) outflow from the target site.
- Ensure that 2-3 mm margins around the defect edge are clear of blood clots, hemostatic reagents and/or loose connective tissue.
- Gel thickness should be limited to 1-2 mm. DuraSeal can swell after application, so it should not be used in areas where neural structures could be compressed.
- The blue color of the hydrogel aids in gauging thickness. As the thickness of the hydrogel increases to 2 mm, the fine epidural vasculature becomes less visible.

When using the MicroMyst Applicator:
1. Prime the Applicator by dispensing a small amount of hydrogel outside of the target site until both precursors flow evenly.
2. Paint the target site with a thin coating of hydrogel by gently pressing the Plunger Cap until a thin layer, approximately 1 - 2 mm in thickness, is formed (Figure 11).

When using the Dual Liquid Applicator
3. Position the applicator 2-4 cm from the target site. Apply firm, even pressure to the center of the plunger cap to dispense the precursors. Rapid initial spraying, followed by a slower controlled rate, is recommended.
4. Continue applying the hydrogel until a thin (1 - 2 mm) coating is formed.
   Note: If delivery is interrupted and the spray tip is plugged, remove the spray tip, wipe the applicator tip, attach a new spray tip and continue delivery.
5. Hydrogel application beyond the defect edges may be removed with scissors or mechanical disruption. Irrigation immediately after the sealant has solidified is permitted.

STORAGE
The DuraSeal Spine Sealant System should be stored at or below 77 °F (25°C).

SYMBOLS USED ON LABELING

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Do not reuse</td>
</tr>
<tr>
<td>Lot</td>
<td>Lot Number</td>
</tr>
<tr>
<td>REF</td>
<td>Catalog Number</td>
</tr>
<tr>
<td></td>
<td>Use by – year and month</td>
</tr>
<tr>
<td>Latex</td>
<td>Latex Free</td>
</tr>
<tr>
<td></td>
<td>See Instructions for Use</td>
</tr>
<tr>
<td>Sterile</td>
<td>Sterile unless the package is damaged or open. Method of sterilization – Radiation</td>
</tr>
</tbody>
</table>

RC only Caution: Federal law (USA) restricts this device to sale by or on the order of a physician.

For more information, or to obtain Covidien documents or references, contact:
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1-781-839-1700