SUMMARY OF SAFETY AND EFFECTIVENESS DATA (SSED)

I. GENERAL INFORMATION

Device Generic Name: Mechanical and enzymatic autologous skin processor for preparing cell suspension, with applicator.

Device Trade Name: RECELL® Autologous Cell Harvesting Device

Device Procode: QCZ

Applicant’s Name and Address: Avita Medical Americas, LLC.
28159 Avenue Stanford, Suite 220
Valencia, CA 91355

Date(s) of Panel Recommendation: None

Premarket Approval Application (PMA) Number: BP170122

Date of FDA Notice of Approval: September 21, 2018

Priority Review: Granted October 27, 2017

Expedited Access Pathway (EAP): Granted EAP designation status on December 10, 2015 because the device provides for more effective treatment of a life-threatening condition and its availability is in the best interest of patients.

II. INDICATIONS FOR USE

The RECELL® Autologous Cell Harvesting Device is indicated for the treatment of acute thermal burn wounds in patients 18 years of age and older. The RECELL® Device is used by an appropriately-licensed healthcare professional at the patient’s point-of-care to prepare autologous Regenerative Epidermal Suspension (RES™) for direct application to acute partial-thickness thermal burn wounds or application in combination with meshed autografting for acute full-thickness thermal burn wounds.

III. CONTRAINDICATIONS

- RECELL® is contraindicated for use on a wound clinically diagnosed as infected or with necrotic tissue present in wound bed.
• RECELL® should not be used to prepare cell suspensions for application to patients with a known hypersensitivity to trypsin or compound sodium lactate solution (“Lactated Ringer’s”).

• The skin sample collection procedure specified for use of RECELL® should not be used with patients having a known hypersensitivity to anesthetics, adrenaline/epinephrine, povidone-iodine, or chlorhexidine solutions.

IV. WARNINGS AND PRECAUTIONS

The warnings and precautions can be found in the RECELL® Autologous Cell Harvesting Device labeling.

V. DEVICE DESCRIPTION

The RECELL® Autologous Cell Harvesting Device is a stand-alone, battery powered cell separation device operated by an appropriately-licensed healthcare professional at the patient’s point of care. The device enables the processing of a small, thin split-thickness skin sample 0.006-0.008 inch (0.15-0.20 mm) in depth to prepare a cell population in suspension for immediate delivery onto a prepared wound surface. Nonclinical performance testing demonstrates processing of harvested autologous skin samples ranging in size from 1 cm² to 6 cm² using the RECELL® device generates a cell suspension that consists of approximately viable cells/cm² tissue processed. By processing multiple (up to 4) samples of 6 cm² autologous thin split-thickness skin, a single RECELL® device can generate a volume of REST™ sufficient to cover an acute burn wound area of up to and including 1,920 cm². However, as stated in the Precautions listed in the RECELL® Autologous Cell Harvesting Device labeling, the RECELL® device should not be used for wounds > 320 cm² or in patients with wounds totaling >20% Total Body Surface Area (TBSA).

The device is a sterile, single use, stand-alone unit with a built-in heater, process indicators, and work surface (the RECELL® Processing Unit, RPU). The user can enzymatically and mechanically process a small skin sample to produce REST™. Processing tools provided with the device include off-the-shelf syringes, scalpels, and fill needles. The device also includes nozzles that attach to syringes and can be used to aerosolize the cell suspension onto the wound. The proprietary RECELL® Enzyme is reconstituted with sterile water (included) and used to facilitate disaggregation of cells from the harvested donor skin. A buffer solution is also provided to suspend the disaggregated cells for delivery to the prepared wound site. The device is designed for point of care use. No cell culturing processes are involved in the procedure. The resulting suspension of cells comprises a mixed population predominantly of keratinocytes and fibroblasts. The presence of viable melanocytes has also been demonstrated.

The product is packaged to facilitate the processing steps for the system components, which are assembled within three (3) boxes, denoted as “A”, “B” and
“C”. The component boxes are placed in a shelf box tray with the RECELL® RPU, a Procedure Guide, and Instructions for Use.

VI. ALTERNATIVE PRACTICES AND PROCEDURES

There are several alternative approaches that may be used for the treatment of serious burns. The most common approach used for burn injuries is excision of the burn wound and prompt closure to stabilize the patient. Currently, the standard of care for wound closure is the use of autologous skin grafting. A wide variety of techniques and methods have been applied for temporary and permanent coverage of an open burn wound.

A commonly used method for reducing the amount of healthy skin harvested, i.e. autograft sparing, involves the use of an autograft meshing device that perforates the harvested skin so that it can be physically expanded and spread across a larger area. Burn surgeons frequently use an autograft, meshed at a 2:1 expansion ratio and applied to the burn injury. For patients with extensive burns, such as greater than 40% total body surface area (TBSA), use of conventional meshing ratios for autograft may not be a feasible option due to lack of sufficient donor sites.

In addition to traditional meshing techniques, other epithelial coverage strategies have been developed to decrease the amount of tissue required for harvesting, including micrografting and suction blister grafting, multiple re-harvests from donor sites, and cultured epidermal autografts. Some of these approaches fall under the purview of ‘practice of medicine’ and are not regulated by FDA; others have limited clinical data and/or are currently being studied but have not undergone complete review by the FDA.

Among the approaches considered for autograft sparing, cultured epidermal autograft products represent a commercially available autologous skin substitute offering the benefits of using autologous tissue and maximal expansion. These products have been approved for use in patients who have deep partial-thickness or full-thickness burns comprising a TBSA of greater than or equal to 30%. It must be noted that the manufacture of these types of products may involve the use of murine (mouse) cells. In these circumstances, when during the manufacturing process human cells/tissues come into immediate direct contact with non-human cells/tissues, the FDA considers the product a xenotransplantation product. Recipients of xenotransplantation products are counseled to actively defer from donating whole blood, blood components, source plasma, source leukocytes, tissue, breast milk, ova, sperm or other body parts for use in humans due to the potential risk for transmission of a zoonotic infection acquired from mouse cells to recipients of donated fluids/tissues.

VII. MARKETING HISTORY
The RECELL® Autologous Cell Harvesting Device has not been marketed in the United States but has received a Conformité Européenne (“European Conformity”, CE) mark and was commercialized in the European Union (EU) in 2005. In addition, the RECELL® device also has been commercially marketed in Australia since 2006 and in China since 2008. The device has not been withdrawn from marketing for any reason related to its safety or effectiveness.

VIII. POTENTIAL ADVERSE EFFECTS OF THE DEVICE ON HEALTH

Most potential adverse effects associated with the RECELL® Autologous Cell Harvesting Device are those typical of care and treatment of burns including those related to the grafting procedure itself and subsequent wound care at both the recipient and donor sites. These include failure to heal or loss of part or all of the graft at the recipient site requiring subsequent surgical and or medical procedures, as well as edema, seroma, infection, pain, neuralgia, scarring (hypertrophy or discoloration), blistering, folliculitis, dermatitis, erythema, pruritus, delayed healing and/or reinjury at the recipient or donor site.

For the specific adverse events that occurred in the clinical studies, please see Section X below.

IX. SUMMARY OF NONCLINICAL STUDIES

A summary of non-clinical laboratory studies that were performed on the RECELL® Autologous Cell Harvesting Device is provided below:

A. Laboratory Studies

**Bench Testing – Design Verification:** Testing was performed on discrete components and assemblies of the device to verify that individual elements function and perform as specified. The purpose and results of the design verification testing performed are summarized in Table 1 below:

Table 1 – Design Verification Testing

<table>
<thead>
<tr>
<th>Test Description</th>
<th>Test Purpose</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>RECELL® Processing Unit (RPU) Functional Testing</td>
<td>Verify the integrated RPU assembly functionality met specified device</td>
<td>Results demonstrated that all RPU assemblies passed the acceptance criteria for the self-test, run sequence, and device lock-out</td>
</tr>
<tr>
<td></td>
<td>requirements. The function of all features of the integrated RPU assembly</td>
<td>functionality. Temperature logs obtained during monitoring of the heating cycle of the RPU, verified that all units met the criteria and</td>
</tr>
<tr>
<td></td>
<td>was tested to established criteria as defined in the RPU specification.</td>
<td>performed as specified.</td>
</tr>
<tr>
<td>Test Description</td>
<td>Test Purpose</td>
<td>Results</td>
</tr>
<tr>
<td>-----------------------------</td>
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<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| RPU Environmental Use Conditions Testing | Verify the integrated RPU assembly functionality met specified device requirements at environmental operating limits. | Results of the environmental conditions testing demonstrated that all RPU assemblies passed the acceptance criteria for all functional tests at the following environmental conditions:  
  - Temperature: 15 - 35°C  
  - Humidity: 10 – 90% relative humidity (RH)  
  Pressure: 65 – 106 kPa |
| Spray Nozzle – Cell Distribution | Demonstrate the cell distribution obtained using the RECELL device spray nozzle provides sufficient coverage during application. | Sprayed cell suspensions demonstrated adequate coverage. The results indicate that the sprayed cells are spread over the surface and provide coverage over nearly the entire surface. |
| Enzyme Activity Verification | Verify that the reconstituted Enzyme maintained sufficient activity to make viable cell suspensions when processing multiple skin samples. | Results show the enzyme activity remained above the acceptance criterion of (b) (4) for all test groups throughout the processing cycle. Use of the same Enzyme solution for multiple samples did not significantly impact cell disaggregation or viability of the cells. |

**Bench Testing – Integrated Device Performance:** Testing was performed using complete RECELL® devices to establish that the different steps of tissue processing with the device are capable of reproducibly processing tissue into viable cell suspensions. The conditions and methods were planned to simulate the intended skin sample processing procedure for the RECELL® device. The objective was to characterize and demonstrate the technical capability of the integrated device under controlled simulated conditions. The purpose and results of the design verification testing performed is summarized in Table 2 below:
<table>
<thead>
<tr>
<th>Test Description</th>
<th>Test Purpose</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell Suspension Characterization</td>
<td>To establish cell processing yields, to verify the viability of cells pre- and post-spray application, and to determine the proportion of cell types. The study also evaluated the</td>
<td>The following results were observed: • Verified that no significant differences in cell viability occurred between pre- and post-spray suspensions produced by the RECELL® device, nor between small and large skin graft sizes.</td>
</tr>
<tr>
<td>Cellular Characterization</td>
<td>cell suspension for the proportion of single cells to aggregates and to confirm that the tissue processing did not significantly increase apoptotic activity. The intent of this study was to demonstrate the ability of the device to produce viable cell suspensions.</td>
<td>• Results demonstrated that fibroblasts and keratinocytes represent the largest proportions of cells, followed by a small proportion of melanocytes. • Determined the absence of cell aggregates of any size in amounts of significance that could affect the application of cell suspensions to patient wounds. • Only approximately 2%, of cells in the suspension were apoptotic, and verified that pre- and post-spray suspensions did not have statistically different apoptotic activity.</td>
</tr>
<tr>
<td>Cell Suspension Reproducibility</td>
<td>Establish that when used by different operators the RECELL® device can generate viable cell yields similar to or greater than yields established in the cell characterization study.</td>
<td>In all cases, users were able to prepare suspensions with viable cell yields at the average or above cell yields established in the cell suspension characterization study. This confirmed that different users could consistently process tissue samples using the RECELL® device to produce cell suspension with viable cells.</td>
</tr>
<tr>
<td>Trypsin Carry-Over Testing</td>
<td>(b) (4)</td>
<td>(b) (4)</td>
</tr>
</tbody>
</table>
**Biocompatibility:** Biocompatibility testing was performed on the sterile RPU assembly, spray nozzles and Enzyme. Testing was performed by an independent testing facility on finished and sterilized product in accordance with International Organization for Standardization (ISO) 10993-1:2009 “Biological Evaluation of Medical Devices Part-1: Evaluation and Testing” as specified in the FDA guidance ‘Use of International Standard ISO 10993-1, Biological Evaluation of Medical Devices – Part 1: Evaluation and testing within a risk management process’ dated June 16, 2016. Biocompatibility tests were conducted in compliance with U.S. Food and Drug Administration Good Laboratory Practice (GLP) regulations set forth in 21 CFR Part 58. All biocompatibility tests passed their corresponding acceptance criteria. Table 3 and Table 4 summarize the results of biocompatibility tests that were performed.

Table 3 – Biocompatibility Testing for RPU Assembly and Spray Nozzles

<table>
<thead>
<tr>
<th>Biocompatibility Test</th>
<th>Results</th>
<th>PASS/FAIL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cytotoxicity (b) (4)</td>
<td>Non-Cytotoxic</td>
<td>PASS</td>
</tr>
<tr>
<td>Guinea Pig Maximization Sensitization Test</td>
<td>No evidence of sensitization</td>
<td>PASS</td>
</tr>
<tr>
<td>Intracutaneous Reactivity Test</td>
<td>Non-irritant</td>
<td>PASS</td>
</tr>
<tr>
<td>Acute Systemic Injection Test</td>
<td>Non-Toxic</td>
<td>PASS</td>
</tr>
<tr>
<td>Rabbit Pyrogen Test (Material Mediated)</td>
<td>Non-pyrogenic</td>
<td>PASS</td>
</tr>
</tbody>
</table>

Table 4 – Biocompatibility Testing for Enzyme

<table>
<thead>
<tr>
<th>Biocompatibility Test</th>
<th>Results</th>
<th>PASS/FAIL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genotoxicity: <em>Salmonella typhimurium</em> Reverse Mutation Assay – Ames Test</td>
<td>Non-mutagenic</td>
<td>PASS</td>
</tr>
<tr>
<td>Genotoxicity: In Vitro Mouse Lymphoma Assay</td>
<td>Non-mutagenic and non-clastogenic</td>
<td>PASS</td>
</tr>
<tr>
<td>Genotoxicity: In Vivo Mouse Micronucleus Assay</td>
<td>Non-mutagenic</td>
<td>PASS</td>
</tr>
</tbody>
</table>
Sterility Assurance: The applicable components and assemblies of the RECELL® device are sterilized by the following traditional methods:

- Enzyme: Gamma radiation sterilization cycle at a [1(4)] dose;
- Buffer: Moist heat sterilization at [1(4)];
- RPU: Ethylene Oxide gas sterilization [1(4)];
- Spray Nozzle Assembly: Gamma radiation sterilization cycle at a [1(4)] dose.

For the Enzyme and the spray nozzle assembly, sterilization validations were completed in accordance with the following standards:

- A Verification Dose Maximum (VD_{max}) method was used for the substantiation of the [1(4)] sterilization dose. All required testing as established by the above standards for sterilization validation was successfully performed and passed all acceptance criteria. The gamma sterilization dose has been successfully demonstrated to be effective in providing the sterility assurance level (SAL) of [1(4)] for the Enzyme and spray nozzle assembly of the RECELL device.

For the Buffer, a sterilization validation was completed in accordance with [1(4)]. An Overkill validation method was employed to assure an SAL [1(4)]. All required testing as established by the above standard for moist heat sterilization validation was successfully performed and passed all acceptance criteria.

For the RPU, a sterilization validation was completed in accordance with [1(4)]. An Overkill validation method was employed to assure an SAL [1(4)]. All required testing as established by the above standard for ethylene oxide sterilization validation was successfully performed and passed all acceptance criteria. Representative samples of the RPU were evaluated for Ethylene Oxide [1(4)] residuals in accordance with [1(4)]. Based on the results, ethylene oxide [1(4)] residuals for the RPU
were within specified limits for devices with limited exposure (b) (4).

**Packaging Integrity / Shipping Testing:** The RECELL® device is packaged in a single corrugated shelf box that has a sliding inner tray that contains a single sterile-packaged RPU and three (3) component set boxes. The capability of the device packaging to protect the device and maintain a sterile barrier has been validated in accordance with the following standards:

- Enzyme and Nozzles- Gamma radiation: Maximum dose (b) (4)
- Buffer- Moist heat: steam cycle
- RPU- EO: ethylene oxide cycle

Finished packaged RECELL® devices were subjected to the following conditioning prior to testing:

- Environmental conditioning (b) (4)
- Simulated transportation conditioning (in accordance with (b) (4))

Results of the device and package integrity testing demonstrate that all components of the RECELL® device met all defined acceptance criteria after being subjected to the environmental and shipping conditions. The integrity of all sterile barrier packages was demonstrated to be intact with (b) (4) detected and the seal strength was maintained within specification for all packages. Container closure integrity for all vials of Enzyme, Buffer, and water-for-injection (WFI) were demonstrated to remain intact with (b) (4) within the specified limits. All remaining tests for product integrity and functionality demonstrated that all components were unaffected by the conditioning and met their design specifications. The packaging configuration of the RECELL® device was qualified to provide a sterile barrier and sufficient protection for the device under expected storage, handling and distribution conditions.
Shelf Life of RECELL® device: Product stability and sterile package shelf life are being qualified under an ongoing stability test program. Finished packaged RECELL® devices were initially subjected to the following conditioning prior to testing:

- Sterilization:
  - Enzyme and Nozzles- Gamma radiation: Maximum dose
  - Buffer- Moist heat: steam cycle
  - RPU- EO: ethylene oxide cycle
- Environmental conditioning
- Simulated transportation conditioning (in accordance with)

RECELL® devices were then divided into two groups, the first group subjected to real-time aging at ambient conditions (25°C/60% RH) and the second group to accelerated-aging conditions. Real-time aging will be performed from 3 months to months, and accelerated aging performed from months representing an equivalent real time of months. Accelerated aging time points were determined in accordance with . At each time point the product integrity and functionality testing is performed.

Currently, product stability and packaging shelf life at 3, 6, months real-time and accelerated aging have been completed. Results of testing demonstrate that the components of the RECELL® device met all defined acceptance criteria for product stability and packaging integrity after being subjected to the aging conditions.

It is noted that the Enzyme stability is evaluated separately. The finished and packaged Enzyme component will be subjected to the following conditioning prior to testing:

- Sterilization - Gamma radiation: Maximum dose
- Environmental conditioning
- Simulated transportation conditioning (in accordance with)

Real time aging will be performed from 3 months to months under ambient temperature 25°C and 60% RH. At each time point the product stability and container integrity testing is performed. Determining stability of the RECELL® Enzyme device component includes measurement of the
Enzyme’s activity using a-based assay that has undergone prior validation.

To date, a retrospective study of Enzyme stability has been performed using Enzyme samples stored at 20-25°C. Results from the retrospective study provide sufficient initial evidence to support a 6-month shelf life for the RECELL® device (based on available stability data provided for the activity of the Enzyme component). A prospective study assessing Enzyme stability will be performed to support future extension of the current 6-month shelf-life.

**Electromagnetic Compatibility and Electrical Safety Testing:**
Electromagnetic Compatibility (EMC) testing was performed on a standalone RPU of the RECELL® device. Testing included compliance for both emission and immunity tests in accordance with . The RPU assembly of the RECELL device met all the applicable requirements of .

Electrical safety testing of the RECELL® device was performed in accordance with . In addition, safety testing performed in accordance with for usability of the RECELL® device. Overall testing results demonstrate the RECELL® meets all applicable requirements set forth in .

**Software Testing:** Testing was conducted to ensure the performance of the embedded RPU firmware (the only software containing component of the RECELL® device) met the software requirements specifications. Verification and validation activities were completed for the device firmware according to the FDA guidance ‘General Principles of Software Validation’ dated January 11, 2002. Testing of the firmware implementation was accomplished through unit, integrated system, and regression testing. The results of all software testing passed all test criteria and based on these results the RECELL® device firmware was verified and validated to meet its functional requirements.

**Human Factors and Usability:** The Human Factors Report provided in the PMA original submission contains summative validation results which demonstrate the RECELL® device user interface (including the device and accessories, Instructions for Use (IFU), Procedure Guide (PG), and representative training) allows for safe and effective operation for the intended use by intended users within the intended environment, without compromise to medical care or patient or user safety. The Report was in alignment with FDA/Center for Devices and Radiologic Health (CDRH) Human Factors guidance and FDA-recognized Human Factors standards:
• FDA CDRH’s final guidance: Applying Human Factors and Usability Engineering to Medical Devices that was issued on February 3, 2016
• IEC 62366-1:2015, titled Medical Devices – Part 1: Application of Usability Engineering to Medical Devices
• AAMI/ANSI HE75: 2009 Human Factors Engineering – Design of Medical Devices

The summative validation study was performed with a total of healthcare professionals who specialized or were experienced in burn care and with intended prior experience with aseptic technique. Each participant completed a series of simulated use scenarios in a representative operating room as they were observed and asked questions by independent moderators. During the collection of objective and subjective data, it was found that all users were able to prepare and apply a simulated cell suspension successfully. No user errors were observed that would lead to or result in death or a severe or permanent injury to the patient or user. There were no critical user errors that would be further mitigated via modifications of the device or user interface and therefore the device has been demonstrated to be suitably designed for its intended use.

B. Animal Studies

Three animal studies were conducted during the initial development of the RECELL® device to establish feasibility of applying autologous cell suspensions to a wound bed. The first two studies were performed early in the development life cycle of the RECELL® device to focus on the feasibility and potential of using an autologous cell suspension to improve wound healing with split thickness autografts. These studies occurred prior to completion of the finished device but used a similar process to the current RECELL® device to generate and apply a cell suspension. A third study was performed using RES from the RECELL® device in porcine tissue. Each of these three studies were designed to evaluate the ability of a cell suspension to facilitate wound re-epithelialization. All three studies were performed in a porcine model, as this has been identified as an acceptable wound healing model. These studies demonstrated that use of an autologous cell suspension combined with autograft can reduce the time required for wound re-epithelialization when compared to autograft. FDA has reviewed the findings in these publications and found them supportive of this approach for human studies.
C. Additional Studies

**Viral Clearance Studies:** The RECELL® device uses as the active ingredient of the RECELL® Enzyme in the cell suspension preparation process.

To establish that the RECELL® Enzyme poses a minimal risk of introducing adventitious viruses to a patient, the effectiveness of the current control measures has been verified for viral clearance in the manufacture of the Enzyme of the RECELL® device.

**Bacterial Endotoxin Testing:** Routine bacterial endotoxin testing (BET) is performed on every lot of the device. Testing was conducted in accordance with the recommendations described in Section V part A.4 of the FDA Guidance “Submission and Review of Sterility Information in Premarket Notification (510(k)) Submissions for Devices Labeled as Sterile” issued on January 21, 2016 to support this labeling. Due to the number of separate sterile components provided as part of the RECELL® device, routine testing is performed on separate components and then the total sum of endotoxin levels from all applicable components is used to determine if the lot has met the BET release criteria. The total combined endotoxin exposure attributable to all device components was demonstrated to be after considering a worst-case enzyme carryover scenario involving the final disaggregated skin cell suspension device output. Test methods performed per are used to detect and quantify bacterial endotoxin for all applicable components of the RECELL® device, with an exception made for the RECELL® Enzyme which requires using a modified
validated method for sample preparation followed by testing in accordance with (b) (4).

X. SUMMARY OF PRIMARY CLINICAL STUDIES

Two prospective randomized clinical studies, CTP001-5 and CTP001-6, were conducted under BB-IDE 13053 to evaluate the safety and effectiveness of the RECELL® device in the treatment of acute burn wounds in a total of 131 subjects. Collectively, these studies evaluated healing results with RECELL® compared to conventional autografting, as well as the differences in donor-site outcomes between the two approaches. In addition to healing, short-term outcomes such as pain, infection and graft loss were also evaluated. Long-term outcomes included evaluation of durability of healing, scarring and patient satisfaction. Study design, study subject demographics, safety and effectiveness endpoints for the two studies are summarized below.

Together, the clinical outcomes support the use of the RECELL® device for treatment of acute thermal burn injuries in adult patients 18 years of age and older and demonstrate that the RECELL® device can be used to facilitate definitive closure of full-thickness and deep partial-thickness burns in a way that spares autografting.

A. Study Design

Study CTP001-6:

In this randomized, multi-center, standard-of-care controlled study, RECELL® was used in combination with autografts that had a higher meshing ratio than control, for the treatment of deep (including full-thickness), extensive burn injuries. The study population included 30 subjects from 6 clinical sites. Subjects were older than 5 years with 5-50% TBSA burn injuries requiring skin grafts for closure. Each subject served as his/her own control, using two comparable contiguous or non-contiguous areas of at least 300 cm² in size. The areas were randomly assigned to receive autografting consistent with the investigator’s pre-specified graft plan (control) or application of the RES over autograft meshed more widely (one ratio higher) than identified in the pre-specified graft plan. Acute healing and pain outcomes were evaluated through 12 weeks; and pain, healing, durability, and scar outcomes were evaluated in the longer-term follow-up visits conducted at 24, 36, and 52 weeks.

1. Clinical Inclusion and Exclusion Criteria

   Enrollment was limited to subjects who were at least five years of age with an acute thermal burn injury of 5-50% total body surface area (TBSA) requiring skin grafting with two areas at least 300 cm², excluding hands, face and joints.
Patients were not permitted to enroll in this study if their burn wounds were due to chemical, electrical, or radioactive sources.

2. Follow-up
Schedule:
All patients were scheduled to return for follow-up examinations at Week 1 +/- 1 day, Week 2 +/- 3 days, Week 4 +/- 3 days, Week 6 +/- 3 days, Week 8 +/- 5 days, Week 10 +/- 5 days, Week 12 +/- 5 days, Week 24 +/- 14 days, Week 36 +/- 14 days, and Week 52 +/- 28 days postoperatively.

Assessments:
Preoperative digital photographs of the Control and RECELL® treated wound sites and the donor autograft sites were taken. Postoperative assessments included the following: 1) Visual Analog Scale (VAS) of study subjects’ satisfaction of the appearance of the donor, control, and RECELL® sites at all time points and 2) blinded evaluator, and study subject completion of the Patient and Observer Scar Assessment Scale (POSAS) to assess the characteristics of the wound scar at 12, 36, and 52 weeks. Clinical assessments were performed at all time points. Adverse events and complications were recorded at all visits.

3. Clinical Endpoints
The co-primary study endpoints were: 1) Non-inferiority (pre-specified non-inferiority margin of 10%) of the incidence of complete wound closure of burn injuries treated with the combination of RES and more widely meshed autografts compared to that for Control (conventional autograft) wounds at eight weeks post-treatment, as assessed by a blinded evaluator; and 2) Superiority in reduced donor area autograft requirements for RECELL® versus control treatment (to establish RECELL® as an autograft-sparing technology), as assessed by the Geometric Mean Ratio (GMR) of the RECELL®:Control autograft expansion ratios, at the time of the autograft procedure. Complete wound closure was defined as complete skin re-epithelialization without drainage, confirmed at two consecutive study visits at least two weeks apart. Safety assessment included evaluation of healing time based on the investigator’s assessment, infection, allergic response to trypsin, durability of wound-healing, scarring outcomes, and device-related adverse events and serious adverse events.

Regarding safety, delayed wound healing, infection and graft failure were adverse events of particular interest.

Regarding effectiveness, subject satisfaction and pain scores were of additional interest.
Regarding success/failure criteria, confirmed complete wound closure prior to eight weeks was defined as treatment success, with the pre-specified non-inferiority margin of 10% (Control minus RECELL®) constituting study success. The other success criterion (co-primary effectiveness outcome) was superiority of RECELL® at the donor site, relative to Control, based on reduction in amount of harvested donor skin.

**Study CTP001-5:**

In Study CTP001-5, the RECELL® Device was evaluated as a primary intervention for the treatment of acute burn injuries in a randomized, multi-center, standard of care controlled (standard split-thickness meshed skin graft) study. The study population included consenting subjects between the ages of 18 and 65 with 1-20% TBSA thermal burn injuries. Each subject served as his/her own control, using two comparable contiguous or non-contiguous areas of deep partial-thickness burns. One site was treated with RES produced by RECELL, and the other was treated with 2:1 meshed autograft. Subjects were evaluated at 1, 2, 3, 4, 8, 16, 24, and 52 weeks.

1. **Clinical Inclusion and Exclusion Criteria**

   Enrollment was limited to subjects who had acute thermal burn injuries requiring skin grafting for closure (deep, partial-thickness) over 1-20% TBSA and who were 18-65 years of age. Subjects were not permitted to enroll in the study if their burn wounds were due to chemical, electrical, or radioactive sources.

2. **Follow-up Schedule:**

   All patients were scheduled to return for follow-up examinations at 1, 2, 4, 6, 8, 10, 12, 24, 36, and 52 weeks after treatment.

   Assessments:
   Preoperative digital photographs of the Control and RECELL®-treated wound sites and the donor autograft sites were taken. Postoperative assessments included the following: Visual Analog Scale (VAS) of study subjects’ satisfaction of the appearance of the donor, control, and RECELL sites at all time points; also, the Vancouver Scar Scale (VSS) was evaluated starting at Week 16. Clinical assessments were performed at all time points. Adverse events and complications were recorded at all visits.

   The key time points are shown above in Table 5 which provides a summary of the study design and safety and effectiveness endpoints.
3. **Clinical Endpoints**

The co-primary effectiveness endpoints were: 1) Non-inferiority of the incidence of RECELL® recipient site (burn injury) wound closure (≥ 95% re-epithelialization) at 4 weeks compared to that of the Control. The pre-specified non-inferiority margin was -10% (RECELL® minus Control); and 2) Superiority of the incidence of donor site healing at 1 week (100% re-epithelialization) for the RECELL® donor site compared to that of the Control site. For both endpoints, healing was confirmed at two consecutive visits. Safety assessments included evaluation of delayed healing, infection, allergic response to trypsin, wound durability, scarring outcomes, device-related adverse events and serious adverse events.

Regarding safety, delayed wound healing, wound scarring, infection and graft failure were adverse events of particular interest.

Regarding effectiveness, patient satisfaction and pain scores were of additional interest.

Regarding success/failure criteria, confirmed complete wound closure prior to four weeks was defined as treatment success, with the non-inferiority of RECELL® relative to Control constituting study success. The other success criterion (co-primary effectiveness outcome) was superiority of RECELL® at the donor site, relative to Control, based on complete donor site healing at one week.

**B. Accountability of PMA Cohort**

**Study CTP001-6**

Thirty subjects were enrolled. All 30 of these comprise the safety population, and 26 were included in the Per-Protocol analysis of the primary non-inferiority effectiveness endpoint. Four of the enrolled 30 subjects were excluded from analyses due to major protocol deviations (two subjects missed the primary endpoint visit at week eight and two study subjects’ wounds included a joint, in violation of the enrollment criteria). The intention to treat (ITT) population for the assessment of superiority at the donor site consisted of 29 subjects.

**Study CTP001-5**

Of the 101 subjects enrolled in this study, 83 were available for assessment of the primary non-inferiority effectiveness evaluation (the modified Per-Protocol population), and 100 were available for evaluation of superiority at the donor site (the ITT population).
C. **Study Population Demographics and Baseline Parameters**

**Study CTP001-6**

Thirty subjects were enrolled in the study, and their wound sites were randomized to the control or the treatment group. The majority of subjects were male (25/30, 83.3%); 66.7% were Caucasian (20/30). The mean age was 39.1 years. Nine subjects had risk factors for impaired wound healing including smoking, drug and alcohol abuse, and inadequate nutrition. The majority of the burn injuries were the result of fire or flames (22/30, 73.3%). The mean percent TBSA affected by burn injuries was 21.2% (±12.8%).

**Study CTP-001-5**

A total of 101 subjects enrolled in the study at 12 US Centers. The mean age of the subjects was 39.5 years (range: 18.2-63.5), with the majority being male (85/101, 84.2%) and Caucasian (59/101, 58.4%). Most of the burn injuries were the result of fire or flames (78/101, 77.2%). Mean percent TBSA affected by burn injuries was 10% (±4.53%) with similar autografting areas for RECELL® and Control recipient (treatment) sites (168.2 ± 68.0 cm² vs. 165.0 cm² ± 66.5 cm², respectively; p=0.3656). On average, autografting was performed 7 days following the burn injury, demonstrating that these partial-thickness burns failed to heal with conservative measures, and therefore confirming that autografting was indicated.

D. **Safety and Effectiveness Results**

1. **Safety Results**

   **Study CTP001-6**

   No unanticipated adverse device effects or device-related events were reported. The number of subjects with any treatment-emergent adverse event (TEAE) at the RECELL® treatment site was the same as the number of subjects with any TEAE at the Control treatment area (17/30, 57%). Similar numbers of TEAEs were reported in areas that were not involved in the study treatments (63%), and 17% of reported TEAEs were not related to any burn. Most subjects experienced TEAEs that were mild (27%) or moderate (37%). There was no overt difference between RECELL® and Control in the incidence of TEAEs at the treatment area (impaired healing, pain, graft loss, skin abrasion, and skin graft failure). The most common TEAE at both the RECELL® and Control treatment areas was pruritus, experienced by 7 (23%) subjects. One or more severe TEAEs were experienced by 7 (23%) subjects; however, no TEAE were related to the RECELL® device. Twelve subjects had serious adverse
events (SAEs), and one died as a result of the SAEs (acute respiratory distress syndrome and subarachnoid hemorrhage, both of which were severe and not related to the study device). There was no difference in the incidence and types of SAEs at the RECELL® and Control treatment areas.

**Study CTP001-5**

Of the 101-subjects, 58% experienced an adverse event, with 36% having an adverse event at the RECELL® sites and 23% at the Control sites. Overall, adverse experiences reported for RECELL®-treated sites were typical for the type of injury sustained by subjects with burn wounds requiring skin grafting procedures. A greater number of subjects had adverse events at the RECELL® sites when compared with the Control sites; however most of these events were mild in nature, were not considered device-related, and were not serious. Additionally, the greater incidence of adverse events noted at RECELL® recipient sites is primarily attributed to events including events contributing to primary endpoint failures, re-injury at the recipient site, and other primarily self-limited skin and subcutaneous tissue disorders such as blisters and excessive granulation tissue. There was no overt difference in the incidence of adverse events between the RECELL® or Control donor sites (4.0% vs. 6.9%, respectively). The observed systemic AEs are consistent with a study population undergoing grafting, and there is no evidence of systemic toxicity associated with the application of RES. In ancillary burn injury areas not included in the randomized treatment areas, 27.7% of subjects experienced similar AEs to the study sites including hypertrophy, hypertrophic scarring, and additional injury (i.e., laceration, skin wound, and skin injury).

**Safety – Additional Endpoints:**

**Study CTP001-6**

Pre-specified safety events including delayed healing, scar necessitating surgical intervention, allergic response to trypsin, wound durability issue, infection, and pain, were evaluated during the study. There was no difference in the incidence of delayed healing and scar revision surgery at RECELL® and Control treatment areas. No patient had either an allergic response to trypsin or a wound durability issue. Infection was not observed at the RECELL® treatment areas but was observed at two Control treatment sites; however, the numbers were too small for any conclusion to be drawn regarding infection and treatment. No clinically meaningful difference was observed in the degree of pain associated with the two treatments.
Study CTP001-5

There was no overt difference in the incidence of graft loss and graft and donor site infection between the RECELL® and the Control sites at any time point. Recipient site scarring was measured by mean total Vancouver Scar Scale (VSS) scores, and there were no differences between the two recipient sites. The RECELL® donor sites had improved appearance at all time points, based on the numerical comparisons of VSS total score outcomes between treatment and Control. Long-term durable wound healing was achieved for both the RES treated and control wounds, as no events of late wound breakdown were reported.

Device Failures: A total of 138 devices were used under clinical study CTP001-5, with 125 used successfully. Root cause analysis identified four device malfunctions attributable to fluid ingress causing electrical shorting in either the heating circuit or the thermistor circuit. Three (3) device failures were determined to be the result of flex circuit crack associated with reported malfunctions of “failed during self-test” and “failed during use”. Investigation revealed cracks in the heating element (flex circuit) which resulted from the use of an incorrect metal during device manufacturing. One device that was returned to Avita under CTP001-5 due to expiration was incorrectly reported as a malfunction. For the remaining 4 devices returned due to reported malfunction, the root cause investigation was either inconclusive, unresolved, or no problem was found.

A total of 34 devices was used under Study CTP001-6, 32 used successfully with 2 device malfunctions reported. Root cause analysis performed in association with the 2 reported device failures was either inconclusive/unresolved or no problem was found.

With respect to all device failures reported for both Studies CTP001-5 and CTP001-6, clinical outcomes were unaffected given the device failures occurred either after the patient had been treated or at a stage when a second RECELL® device could be used to complete the procedure successfully.

2. Effectiveness Results

Study CTP001-6

The co-primary endpoints for this study were non-inferiority of RECELL® plus autograft relative to autograft control, at the recipient sites at Week 8 and superiority in sparing of donor site skin for RECELL® relative to control at the time of treatment.
Non-inferiority of RECELL® relative to Control for recipient site healing was established using the pre-specified non-inferiority margin of 10%. Confirmed treatment area closure by Week 8 was 92.3% for RECELL® vs. 84.6% for the Control treatment areas. The treatment difference was -7.7% (Control minus RECELL®, 95% CI upper bound of 9.55%).

Superiority of RECELL® was established with respect to relative reduction in donor site harvesting (p<0.001). The mean donor site areas for RECELL® and Control were 270.5 ± 123.7 cm² and 368.0 ± 150.1 cm², respectively. Secondary effectiveness outcomes (patient satisfaction, Week 24 observer overall opinion on POSAS, and Week 24 patient overall opinion on POSAS) were comparable between treatments.

**Study CTP001-5**

The co-primary endpoints for this study were non-inferiority of RECELL®, relative to autograft control, at the recipient sites at Week 4 and superiority of donor site healing for RECELL® relative to control at Week 1.

At Week 4, 94.3% of the RECELL® recipient sites achieved healing vs. 100% of the Control recipient sites. Although comparable numbers were achieved, the primary endpoint of non-inferiority of RECELL® relative to Control for recipient site healing was not established in the per protocol population using the pre-specified non-inferiority margin of -10% (RECELL® minus Control; difference -5.7%, 95% CI: -12.8% to -0.4%). When evaluating a Modified per protocol population (MPP) (post-hoc analysis), defined to exclude the subjects managed post-operatively with silver sulfadiazine (a cytotoxic agent), the difference in proportions for complete wound healing between the RECELL® and Control recipient areas established non-inferiority for RECELL® compared with Control treatment (RECELL® minus Control difference was -2.4%, 95% CI: -8.4 to 2.3%). The MPP analysis provides evidence that when RES-treated sites receive appropriate aftercare, healing is non-inferior to those sites treated with meshed autografts. Donor site healing was superior at Week 1 for the RECELL® donor sites versus the Control donor sites (p=0.0042). Subjects reported less pain at the RECELL® donor site compared to the Control donor site at every time point within the 8 weeks following treatment. Similarly, subjects expressed greater satisfaction with the visual appearance of the RECELL® donor site compared with the Control donor site at all longer-term follow-up visits. The mean donor site area for RECELL® was substantially less than that of the Control: 4.7 ± 3.19 cm² vs 194.1 ± 158.5 cm², respectively.

3. **Subgroup Analyses**

**Study CTP001-6**
The study population was mostly comprised of White adult males. Due to the small sample size and the relative homogeneity of the study population, no conclusions can be drawn regarding outcomes for subgroups based on sex, age, and ethnicity.

**Study CTP001-5**

The limited sample size and the relatively homogeneous study population limit the ability to draw conclusions regarding the safety or efficacy of RECELL® in subgroups based on age, sex, or race.

4. **Pediatric Extrapolation**

**Study CTP001-6**

Although this study did enroll pediatric subjects, the youngest study subject was nine years old, and only three study subjects were younger than 18 years. Thus, there is an insufficient amount of data to assess the safety and efficacy of RECELL® for treating full-thickness acute thermal burn wounds in pediatric subjects.

**Study CTP001-5**

Pediatric study subjects were excluded, and none were enrolled.

In this premarket application, existing clinical data were not able to be leveraged to support approval of a pediatric patient population. The data were derived from a compassionate use protocol that used a treatment methodology that differed substantially from that of the controlled clinical studies. In addition, the compassionate use protocol lacked a control, and there was insufficient follow-up for most of the patients. Finally, pediatric patients have skin that is still growing and developing, and therefore a separate clinical study is needed to establish safety and efficacy of RECELL® in the pediatric population.

**E. Financial Disclosure**

The Financial Disclosure by Clinical Investigators regulation (21 CFR 54) requires applicants who submit a marketing application to include certain information concerning the compensation to, and financial interests and arrangement of, any clinical investigator conducting clinical studies covered by the regulation. As certified in the FORM FDA 3454 submitted by the applicant, none of the 66 clinical investigators had disclosable financial interests/arrangements as defined in sections 54.2(a), (b), (c), and (f). The information provided does not raise any questions about the reliability of the data.
XI. SUMMARY OF SUPPLEMENTAL CLINICAL INFORMATION

No supplemental clinical information was included in this application.

XII. PANEL MEETING RECOMMENDATION AND FDA’S POST-PANEL ACTION

In accordance with the provisions of section 515(c)(3) of the act as amended by the Safe Medical Devices Act of 1990, this PMA was not referred to an FDA advisory committee for review and recommendation because it was judged that the expertise of the FDA PMA review staff was sufficient to determine there is a reasonable assurance the device is safe and effective based on the information provided in the PMA application when used in accordance with the indications for use.

XIII. CONCLUSIONS DRAWN FROM PRECLINICAL AND CLINICAL STUDIES

A. Effectiveness Conclusions

The effectiveness of RECELL® as an autograft-sparing technology capable of achieving definitive wound closure is substantiated through results derived from two separate Avita Medical-sponsored US multicenter, randomized, controlled clinical studies, collectively providing experience for 131 patients for a 52-week period following treatment. The use of RECELL® either as a primary or adjunct intervention to meshed skin grafts, results in wound closure that is non-inferior to that achieved with conventional autografts, with substantially less donor site harvesting requirements (97% reduction in donor skin area in CTP001-5 and approximately 30% in CTP001-6).

Together with the smaller donor site taken for the RECELL® process (compared to autografting), the donor site was determined to heal faster than the donor site for the conventional autograft and was associated with less pain, improved appearance, and patient satisfaction (CTP001-5). During the follow-up period of 52 weeks, there were no events of recurrent wound breakdown following closure, indicating durable wound healing.

Among the two studies presented, there were no differences between RECELL® and the control treatment with respect to the type of complications of concern for autografting procedures --- i.e., graft loss or graft failure, impaired healing, infection, or scarring. Within the CTP001-5 study, only 1 of the 101 subjects (1%) treated with RES required subsequent re-grafting for healing. Within the CTP001-6 protocol, 2 (7%) subjects treated with RECELL® required a secondary grafting procedure, whereas 3 (10%) of Control subjects required re-grafting.
B. **Safety Conclusions**

The risks of the device are based on non-clinical bench testing and animal studies as well as data collected in clinical studies conducted to support PMA approval as described above. Within randomized studies, adverse events documented are both anticipated and consistent with a patient population undergoing autografting for treatment of burn injuries.

C. **Benefit-Risk Determination**

Two adequate and well-controlled clinical studies were conducted to generate safety and effectiveness data to support PMA approval, as described above. The data provide a reasonable assurance of safety and effectiveness for RECELL® as an autograft-sparing technology indicated for use at the patient’s point-of-care for preparation of an autologous skin cell suspension to be applied to a prepared wound bed. When used by a licensed healthcare professional at point-of-care, the suspension can be used to achieve epithelial regeneration for definitive closure of acute thermal burn injuries. Furthermore, these data support a low rate of device-related adverse events.

Potential challenges with RECELL® system use and patient post-discharge wound care were identified during the two clinical trials. To minimize the possibility that adverse effects that could result from insufficient familiarity with the recommended procedures for device use and post-discharge wound care, the Applicant developed a health care practitioner training program, as well as post-discharge patient education material.

Clinical data demonstrate that the risks associated with RECELL® use are low and consistent with complications of standard autograft treatment such as potential for non-healing, graft loss, infection and scarring. The benefits derived from the use of the device, i.e., definitive wound closure of a larger burn surface with a smaller donor site defect outweigh the potential for harm.

Thus, when RECELL® is used in accordance with instructions for use and patient wound care is performed according the recommended methods, the benefits associated with use of the RECELL® device outweigh the risks.

D. **Overall Conclusions**

The data in this application provide a reasonable assurance of safety and effectiveness of this device when used in accordance with the indications for use.
XIV. CBER DECISION

CBER issued an approval order on September 21, 2018. The final nonclinical conditions of approval cited in the approval order are shown below.

(b) (4) Assay Validation

1. The applicant will perform Assay validation (specificity, accuracy, linearity, repeatability, intermediate precision, range, and ruggedness) at each laboratory that conducts the assay using a statistically meaningful number of samples of the RECELL® Enzyme obtained from multiple production lots. Results obtained by each participating laboratory should be comparable when testing the same sample using the validated assay. The applicant will submit the testing protocol(s) and complete assay validation test report(s) as a Post-Approval Study Report by February 18, 2019.

Enzyme Stability and RECELL Device Shelf Life

2. RECELL® Enzyme stability determines RECELL® device shelf life. The applicant will conduct a prospective RECELL® Enzyme stability study that involves assessment of multiple attributes (appearance, dissolution time, container closure/integrity, enzyme activity, and bacterial endotoxin). Samples of enzyme from separate production lots (individual samples per lot) that have been subjected to worst-case sterilization, atmospheric conditioning, simulated shipping, and real-time aging under ambient conditions will be evaluated according to the stability study protocol to re-affirm the current 6-month RECELL® device shelf life. The applicant will submit the results of the executed stability study re-affirming 6-month stability as a Post-Approval Study Report by September 20, 2019 and as additional 30-Day Notices for further shelf life extensions.

Manufacturing Facility Pre-Approval Inspection

During review of the PMA application, the determination was made to recommend waiver of the pre-approval inspection of the applicant’s manufacturing facilities. This decision was based on information provided in the PMA application in conjunction with prior inspection reports and related correspondence supporting the overall compliance of the applicant’s registered manufacturing facilities with the device Quality System (QS) regulation (21 CFR 820).

XV. APPROVAL SPECIFICATIONS

Directions for use: See device labeling.
Hazards to Health from Use of the Device: See Indications, Contraindications, Warnings, Precautions, and Adverse Events in the device labeling.

Post-approval Requirements and Restrictions: See approval order.

XVI. REFERENCES

None