AVITA MEDICAL

Company Overview



January 2019

Disclaimer – Forward Looking Statements

This presentation may include forward-looking statements. You can identify these statements by the fact that they use words such as "anticipate", "estimate", "expect", "project", "intend", "plan", "believe", "target", "may", "assume" or similar expressions.

These forward looking statements speak only as at the date of this presentation and are based on management's expectations and beliefs concerning future events. Forward-looking statements are necessarily subject to risks, uncertainties and other factors, many of which are outside the control of Avita Medical that could cause actual results to differ materially from such statements.

Avita Medical makes no undertaking to subsequently update or revise the forward-looking statements made in this release to reflect events or circumstances after the date of this release.

This presentation is intended to provide background information only and does not constitute or form part of an offer of securities or a solicitation or invitation to buy or apply for securities, nor may it or any part of it form the basis of, or be relied on in any connection with any contract or commitment whatsoever.



Overview of Avita Medical



AVITA Medical – Regenerative Medicine Company

About AVITA

- Regenerative medicine company with platform technology providing innovative treatments derived from the regenerative properties of a patient's own skin
- Experienced leadership team
- Headquartered in California
- Traded on ASX, with ADRs in U.S.
- Substantial U.S. Government support under BARDA program

Lead Product Approved

FDA approved the RECELL System® PMA on 20 September 2018 as Class III device for treatment of acute thermal burns









Leadership Team with the Right Expertise



Dr. Michael S. Perry *CEO*>30 years experience



Dale Sander CFO>30 years experience

Affiliations:



CCO
16 years experience

Affiliations:



Tim Rooney CAO25 years experience



Andrew Quick Sr VP, Clinical Dev. 22 years experience



Donna Shiroma General Counsel 20 years experience

Affiliations:





BAY CITY CAPITAL





Allergan



Affiliations:

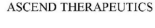






Affiliations:







Affiliations:

Johnson-Johnson









THERAPEUTICS







AVITA Medical Board and Capital Structure

A\$0.080 Share Price¹ 1.652 Billion Shares Outstanding

A\$132.2 Million Market Capitalization¹ A\$36.5 Million Cash²

A\$0.0 (Zero) Debt

DIRECTORS



Dr. Michael Perry CEO, AVITA Medical



Professor Suzanne Crowe
Multiple positions including
Associate Director of the
Burnet Institute



Louis Drapeau
Nektar Therapeutics, BioMarin
Pharmaceutical, Inc., and
Arthur Andersen LLP.



Karst Peak Capital Limited 17.3% Redmile Group 9.4% BioScience Managers Pty Ltd 5.0%



Jeremy Curnock Cook
Managing Director of
Bioscience Managers Pty
Ltd

Lou Panaccio, Chairman

Non-Executive Director

Sonic Healthcare Limited



Damien McDonaldChief Executive Officer of LivaNova

ANALYSTS

John Hester, Bell Potter (AUS) Brooks O'Neil, Lake Street (US)

^{2.} As of 30 September 2018, pro forma to include A\$22.4 million in net proceeds from equity placement received in December 2018. Excludes \$13.7 million in net proceeds scheduled to be received from 2nd Tranche of equity placement in January 2019.



^{1.} As of 4 January 2019

Strategic Path: Transforming Lives with Skin Regeneration



Leading the way in skin regenerative wound therapy Acute thermal burns, trauma, & chronic wounds



Expanding our footprint within regenerative dermatology
Hypopigmentation: Vitiligo



Advancing into Cell and Cell-Based Gene Therapy Aesthetics, Cell & Gene Therapy e.g., Dystrophic EB



RECELL Overview



Healthcare professionals should read the INSTRUCTIONS FOR USE - RECELL® Autologous Cell Harvesting Device for a full description of important safety information including contraindications, warnings and precautions.

RECELL System Skin Regeneration Platform

Regenerative Medicine Platform

 An Autologous Cell Harvesting Device that uses proprietary enzyme and buffer formulations to generate Spray-on Skin™ Cells within 30 minutes

Designed by Surgeons

 An elegant means to deliver skin regeneration to patients at point of care

Proven Safety and Effectiveness

- 7,000+ uses to date in multiple world markets with no observed safety signals
- Treatment area is 80X donor area (skin sample the size of credit card can be used to treat a patient's entire back)
- Compelling clinical results (RCTs) and robust health-economic data

>50 Peer-Reviewed Publications





Current Standard of Care Is Suboptimal and Expensive

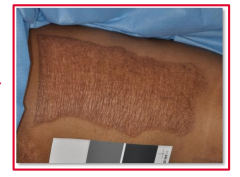
Autografts - Split-Thickness Skin Grafts (STSG) - Used in 75% of Cases



Harvesting skin from donor site for STSG



Donor site wound created while harvesting skin for autograft



Typical SOC donor site scar 52 weeks post procedure

KEY SHORTCOMINGS OF SOC

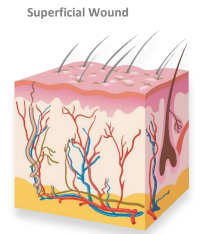
- Large donor area required
- Pain during and post procedure
- Extended hospitalization and costs
- Multiple complex, costly, surgical procedures
- Risk of infection

Healing burn injuries induces trauma of its own

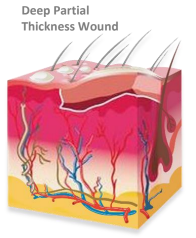
Under Current Standard of Care
Average USD \$792,000 cost and 59.4 days in hospital for 40% TBSA burns¹

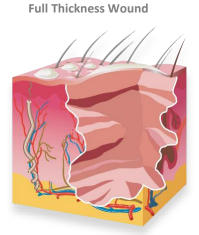


U.S. Clinical Trials Supporting RECELL Use in Burns









Clinical Support for RECELL System

- Two multicenter, randomized, controlled clinical trials consisting of 131 patients
- Additional 155+ patients treated in Compassionate Use and Continued Access programs
- Real-world experience in more than
 7,000 patients globally

Pivotal Trial #1 RECELL Versus SOC (STSG) in Second-Degree Burns Pivotal Trial #2 RECELL with widely spaced SG Versus SOC (STSG) in Third-Degree Burns

FDA Compassionate Use Investigational Device Exemption (IDE) Program (90+ Patients)

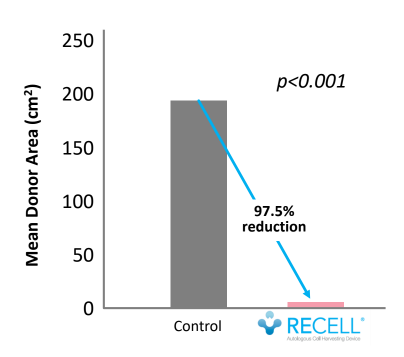
FDA Continued Access Investigational Device Exemption (IDE) Program (65+ Patients)



Pivotal Trial 1: RECELL System *Alone Versus* SoC (STSG)

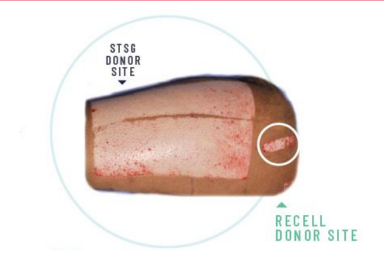
Deep-Partial Thickness (Second-Degree) Burns

Reduced Donor Skin Requirement



Equivalent healing of burn sites with significantly less donor skin required

Reduced Pain and Scarring



- Significantly less donor-site pain (p≤0.0025)
- Significantly better donor-site appearance (p≤0.0025)
- Significantly reduced donor-site scarring (p≤0.0025)
- Significantly greater incidence of donor-site healing at two weeks (p<0.001)

& Research

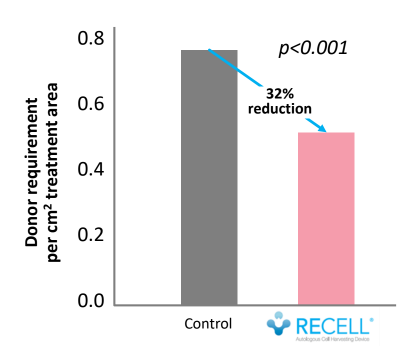
Published in JBCR and Presented at ABA



Pivotal Trial 2: RECELL System Combined With Widely-Spaced Skin Grafts <u>Versus</u> SoC (STSG)

Full-Thickness (Third-Degree) Burns

Reduced Donor Skin Requirement



Published in Burns and Presented at ABA

Positive Treatment Outcome

- RECELL System achieved definitive closure comparable to standard of care with significantly less donor skin
- At eight weeks post treatment,
 - <u>92 percent</u> of the burn sites treated with the RECELL System achieved complete healing versus
 - 85 percent for the sites treated with the standard of care





Compassionate Use Provides Additional Case Studies











Treatment Day

Day 7

Day 21

3 months

1 year

A CASE FROM A FACIAL BURN PATIENT Case Series Presented at ABA Meeting - APRIL 2018

- 12-year-old girl with 2nd-degree facial burn and widespread 3rd-degree burns
- 62% Total Body Surface Area (TBSA) burn injury
- Insufficient donor skin available for SoC (STSG)
- Reintroduction of melanocytes resulted in an excellent cosmetic outcome
- No facial contracture release surgery required
- Discharged in 24 days



RECELL is ideal for treatment of facial burns



Additional RECELL Clinical Results to be Featured at 2019 Burn Meetings (18 Abstracts Already Accepted)



American Burn Association Annual Meeting 2-5 April 2019

9 abstracts accepted for presentation

Presentations include:

- Pediatric patients
- Large burn injuries, TBSA 52% to 91%
- Donor sites
- Burn injuries of the hands
- Budget impact of RECELL use versus SOC (Arizona Burn Center)



John A Boswick Burn & Wound Symposium 2-7 February 2019

8 abstracts accepted for presentation

Presentations include:

- Pivotal trials in 2nd and 3rd degree burns
- Health economic model
- Necrotizing soft tissue infection
- Large TBSA burn injuries
- RECELL combined with dermal substitutes



LA-ACS/SAL Annual Meeting 18-20 January 2019

One abstract accepted for presentation

Presentation:

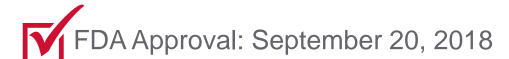
Reduction in hospital stay due to treatment with RECELL



Burn Market & RECELL Commercial Strategy



Commercial Plan was Well Prepared. FDA Approval Triggered a Cascade of Activities



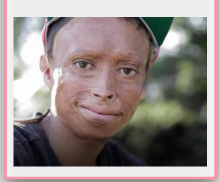
- Customer Service Line Active Within 2 Business Days
- American Burn Association (ABA) Issued Reimbursement Coding Guidelines Within 1 Week of Approval
- Commercial Product Availability Within 2 Weeks of Approval
- First Commercial Sale Within 2 Days of Product Availability
- Entire US Field Force in Place Within 8 Weeks of Approval
- National Sales Launch Commenced January 2019



Initial U.S. Target Market: In-Patient Burns of 10%+ **TBSA that Require Autografting**

486,000

Burn Patients Treated Annually in the US1



42,402

In-patient Burn Treatments²



75%

In-patient **Burns are Treated** in Burn Centers³



14,146

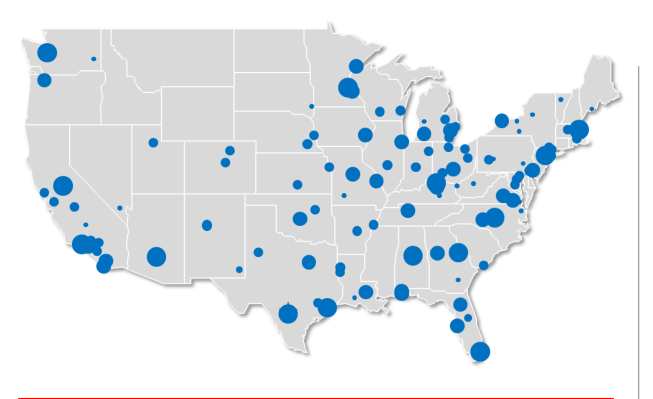
10%+ TBSA In-Patients at Burn Centers are **Target Candidates** for RECELL4



American Burn Association. National Burn Repository Report. 2016; Version 12.0 also http://ameriburn.org/who-we-are/media/burn-incidence-fact-sheet/

Calculated off inpatient population and triangulated from http://ameriburn.org/who-we-are/media/burn-incidence-fact-sheet/

U.S. Burn Market is Highly Concentrated Making It Easily Accessible



24 Burn Centers Already Have Experience with RECELL System⁴ Represent 30% of Patients Treated, Includes #1 and #2 Centers in U.S.

- 134 burn centers in the U.S.¹
- 300 burns surgeons in the U.S.²
- Burn centers see 65 times more burn hospitalizations than in general hospital setting³
- The ABA mandates that severe burns, meeting certain criteria, must be transferred to an ABA burn center



^{1.} American Burn Association. National Burn Repository Report. 2017; Version 12.0 ZS Associates Pricing Research 2018

^{2.} Calculated from: http://ameriburn.org/who-we-are/media/burn-incidence-fact-sheet/

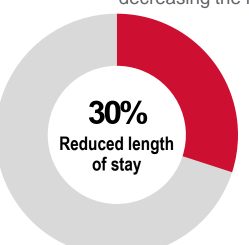
^{3.} Calculated off inpatient population and triangulated from http://ameriburn.org/who-we-are/media/burn-incidence-fact-shee

^{4.} Through date of approval from clinical trials and Compassionate Use and Continued Access programs

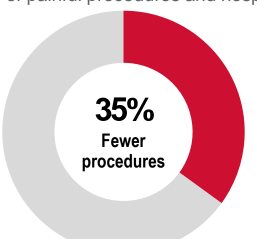
Health Economic Model Demonstrates that RECELL Can Reduce Overall Hospital Costs

Transforming Care

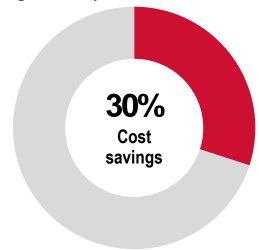
Reduces costs and accelerates recovery by decreasing the number of painful procedures and hospital length of stay



Fewer procedures and faster healing time gets patients home quicker



Reduced donor site size and greater meshing ratio enables permanent closure with fewer invasive autograft procedures



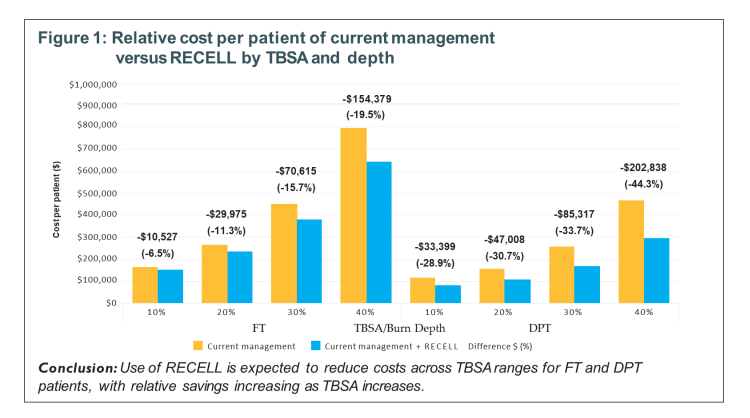
Shorter and fewer procedures, reduced length of stay, and reduced resource use translates into burn center savings

RECELL Saves the Hospital Money in All In-Patient Scenarios Where the Burn is 10% Total Body Surface Area (TBSA) or Larger



Health Economic Model Demonstrates RECELL Cost Savings Per-Patient Savings

- IQVIA (IMS)
 developed a Burn
 Care Pathway
 Health Economic
 model
 demonstrating
 RECELL savings
- Validated model provides VAC (Value & Analysis Committees) strong economic justification for adopting RECELL



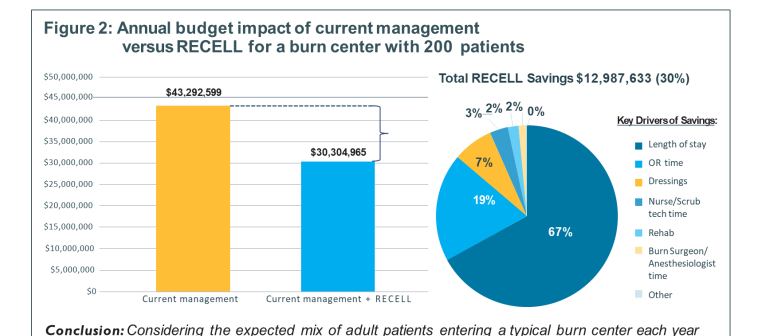
Use of the RECELL System could reduce the cost of treatment by 44% or greater in patients with large burns

Sets a New Standard of Validating Cost Effectiveness for Any New Product in Burns



Health Economic Model Demonstrates RECELL Cost Savings Annual Burn Center Savings

- Model can be tailored to patient populations relevant to individual hospitals, healthcare systems, etc.
- Robust publication and podium schedule



(as informed by NBR data), use of RECELL in burn management is expected to reduce costs overall.

For a burn center with 200 patients, the use of RECELL would reduce annual total treatment costs from \$43.3 million to \$30.3 million, saving 30% or \$13.0 million

Customized Projections Can be Created for Each Burn Center Showing Annual Savings



RECELL System is Priced Right for All Burn Sizes Pricing of Other Treatments Limit Them to Large Burns

Therapy Price/cm² (USD)



Assumptions

- Skin TE \$60/cm2
- Epicel ~\$50/cm2; 1%TBSA treatment with Epicel costs at \$6-10,000; Epicel Skin Grafts
- Integra \$28/cm2. Complementary product presented for pricing comparison
- RECELL® 1920 up to 10% TBSA. Complementary product presented for pricing comparison

RECELL is Priced for Broad Market Adoption



Creation of Best in Class Market Access Program Will Address Market Needs

Key Launch Need

Addressing the Need

Physician payment

CPT Codes





Ensure Hospital Payment

ICD-10 Code for procedural coding





Reimbursement Guidelines

Reimbursement and Coding Guides



Customers need quick, knowledgeable responses for reimbursement inquiries



Reimbursement Customer Service Line Director of Market Access



ABA Provided Recommended CPT Codes Within One Week of Approval

CPT = Current Procedural Terminology ICD = Internal Classification of Disease



US Commercial Field Team is in Place

25

Average Years of Industry Experience (Sales Leadership)

15.8

Average Years of Burn Care Experience (Entire Field) 100%

Have Burn Care Experience

12

Average Years of Surgical Selling & Case Support Experience 100%

Have Successful Launched a New Product

20 Field Positions Will Provide Deep Coverage to All 134 US Burn Centers



All Preliminary Indicators Point to Success

Tremendous excitement and interest from potential customers



36 burns centers have begun the purchase authorization process with their hospital administration



27 burn centers have already been trained and certified



12 accounts have already placed orders

All Accomplishments in Advance of Launch and Field Selling



All Requirements for Successful U.S. Launch in Place







✓ Compelling health economics (cost savings)



Reimbursement in place and well supported

Experienced field force in place

Credible and active communication plan

Robust field deployment model

✓ Customer service center



Pipeline and Milestones



AVITA Medical Pipeline

Development Programs

	Indication	Real-World Experience	POC Studies	Pivotal Trials	Approval U.S. OUS ¹
Current RECELL Platform	Thermal Burns Adults		$\boxed{\hspace{1.5cm}\checkmark\hspace{1.5cm}}$	lacksquare	
	Thermal Burns Pediatrics	✓	lacksquare	Randomized trials und	
	Chronic Wounds: VLU and DFU		✓	Partner and take into pivotal trials	
	Hypopigmentation: Vitiligo and Scars		✓	Advanced controlle	
	Trauma Wounds		Pivotal cli	nical trial	
and ne apy	Rejuvenation		Preclinical POC		
Cell and Gene Therapy	Skin diseases e.g. Epidermolysis		Preclinic	cal POC	



Europe: Burns, chronic wounds, scars and vitiligo China: Burns, acute wounds, scars and vitiligo Australia: Burns, acute wounds, scars and vitiligo



Financial Overview

(AUD in 000s)	Year Ended June 30,	
	2018	2017
Revenue	\$11,372	\$8,132
Operating Costs	28,571	20,186
Net Loss	(16,484)	(11,511)
Cash	14,826 ¹	3,790

Tickers: ASX:AVH and OTCQX:AVMXY





¹Excludes additional A\$3.0 million in net proceeds received in July 2018, A\$22.4 million in net proceeds received in December 2018, and \$13.7 million in net proceeds scheduled to be received in January 2019 from equity financings.

BARDA Program

- U.S. Biomedical Advanced Research and Development Authority
 - Mandate: disaster preparedness & response
- Providing sizable non-dilutive funding
- Total estimated contract value US\$80.1M
- Major programs supported:
 - o PMA
 - Health Economic Model
 - Pediatric clinical trials
 - Disaster preparedness stockpile









2018/2019 Value-Creating Milestones

2018 has been a Transformative Year for AVITA

- ✓ PMA approval by US Food & Drug Administration
- ✓ Accelerated launch preparation activities
- ✓ Robust manufacturing capabilities
- ✓ High impact of RECELL clinical data
- ✓ Continuing to advance pipeline

RECELL is Positioned for Successful Adoption in US Burns during 2019

Key milestones for 2019

- RECELL U.S. market launch / revenue growth
- Publication of 3rd degree burn pivotal trial results
- Publication of RECELL health economic model
- Presentations of RECELL compassionate use data at 2019 ABA meeting
- Commencement of traumatic wounds pivotal clinical trial
- Commencement of vitiligo clinical trial(s)
- BARDA procurement
- Listing of ADRs on NASDAQ



Risk Factors

There are numerous risk factors involved with the Company's business. Some of these risks can be mitigated by the use of safeguards and appropriate systems and controls, but some are outside the control of the Company and cannot be mitigated. Accordingly, an investment in the Company carries no guarantee with respect to the payment of dividends, return of capital or price at which securities will trade. The following is a summary of the more material matters to be considered. However, this summary is not exhaustive. Potential investor should consult their professional advisors before deciding whether to invest.

- Technological Change: Technological change presents the Company with significant opportunities for growth. However, the risk remains that any competitor may introduce new technology enabling it to gain a significant competitive advantage over the Company.
- Reliance on key personnel: The Company's success depends to a significant extent upon its key management personnel, as well as other
 management and technical personnel including sub-contractors. The loss of the services of any such personnel could have an adverse effect on
 the Company.
- Competition: The Company competes with other companies, including nationally in Australia and internationally. Some of these companies
 have greater financial and other resources than the Company and, as a result, may be in a better position to compete for future business
 opportunities. There can be no assurance that the Company can compete effectively with these companies.
- Patent Protection: The patent protection that the Company may obtain varies from product to product and country to country and may not be sufficient, including to maintain product exclusivity. Patent rights are also limited in time and do not always provide effective protection for products and services: competitors may successfully avoid patents through design innovation, the Company may not hold sufficient evidence of infringement to bring suit, or the infringement claim may not result in a decision that the rights are valid, enforceable or infringed.
 Legislation or regulatory actions subsequent to the filing date of a patent application may affect what an applicant is entitled to claim in a pending application and may also affect whether a granted patent can be enforced in certain circumstances. Laws relating to biotechnology remain the subject of ongoing political controversy in some countries. The risk of changed laws affecting patent rights is generally considered greater for the biotechnology field than in other longer established fields.
- Change in government policy and legislation: Any material adverse changes in relevant government policies or legislation of Australia / United States may affect the viability and profitability of the Company, and consequent returns to investors. The activities of the Company are subject to various federal, state and local laws governing prospecting, development, production, taxes, labor standards and occupational health and safety, and other matters."



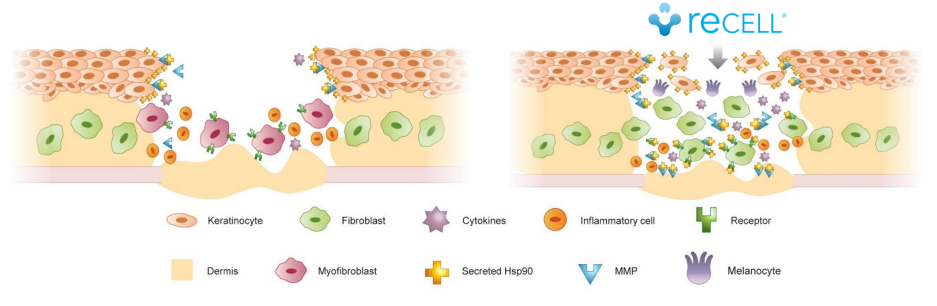
Appendix Mechanism and Additional Burn Case Studies



MOA: Disaggregated cells facilitate fast & effective skin regeneration

Healing Process Without RECELL

Healing Process With RECELL



- ReCell processes small samples of patients' own skin to create a cell suspension of disaggregated cells
- Disaggregated skin cells in suspension form new tissue across the entire area rather than waiting for cellular resources from the wound edge
- Cell suspension includes pigment-producing cells (melanocytes)
- Cell suspension facilitates re-epithelialization of areas of viable dermis (partial-thickness burns), and areas within the spaces of split-thickness autografts for full-thickness burns

Disaggregated Autologous Cells from RECELL Support Re-epithelialization



RECELL Achieved Healing and Pigmentation When Standard of Care Failed

Case Report: RECELL Treatment Outcome for Deep Partial-Thickness Burn

- 48-year-old victim of a gas boiler explosion
- Standard of care failed to heal the
 2nd degree facial burn wounds
- Use of RECELL achieved wound healing
- Reintroduction of melanocytes resulted in an excellent cosmetic outcome
- RECELL's unique advantages make it the ideal solution for facial burns and other visible burn sites

Treatment
Excision and ReCell®



Post-Operation





Restoration of Normal Pigment Critical For Patients



Pediatric Opportunity

High Percentage of Burns are in Pediatric Patients

- 32% of burns occur between ages 5 and 15.9
- Majority suffer from scald burns (65%)

Avita has Initiated studies with RECELL in Pediatrics

- Q3 2018 Commencement of US Paediatric Burns Clinical Trial
- Q3 2018 Commencement of Australian Paediatric Scald Study

Scalds Allows Expansion into another Site-of-Service – Outpatient Setting



Before treatment



3 weeks post RECELL treatment



post RECELL treatment



10 months post RECELL treatment

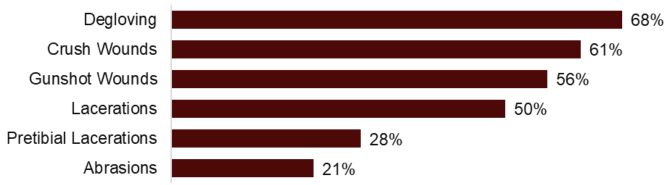


Appendix Follow-On Indications Beyond Burns



RECELL Presents a Strategic Opportunity in Traumatic Wounds

Percentage of Wounds Requiring Skin Grafts





- ~1/3 of all skin grafts are trauma related
- ~50% of Burn Surgeons also work in trauma centers
 - Synergistic with current commercial efforts in burns
- RECELL used by multiple international surgeons in Traumatic Wounds with positive outcomes



 ²⁰¹⁸ Internal Market Research

²⁰¹⁸ IMS Data

^{3.} US Skin Graft Market - Industry Trends Forecast to 2025

Regenerative Dermatology Opportunity in Vitiligo

High Market Value • Large Population • Focused to a Specialty • Clinical Data



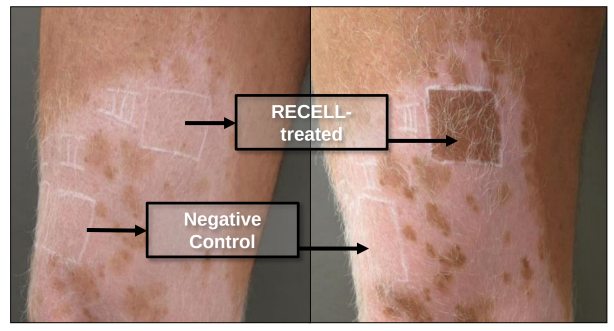


^{1.} Research & Markets: Vitiligo Therapeutics - Pipeline Assessment and Market Forecasts to 2019 2012

[.] Advances in Vitiligo: An Update on Medical and Surgical Treatments. A. Dillon, et al. J Clin Aesth Derm. 2017

^{3.} The Epidemiology and Treatment of Vitiligo: A Chinese Perspective Xiaolan Ding, Juan Du and Jianzhong Zhang* Journal of Pigmentary Disorders. 2014
4. Internal market research 2018

RECELL Was Able to Repigment 100% at 6-months (Vitiligo)



Baseline

6-months Follow-up

- After 6-months, the RECELL-treated area is 100% re-pigmented, with mild hyperpigmentation (UVA daily)
- Control area is 0% re-pigmented



Early Research Programs to Advance RECELL Platform

High Market Value • Focused to a Specialty • Expansion into other Disease States



Cell-based Skin Gene Therapy e.g., Epidermolysis Bullosa (EB)

EB: An incurable, group of genetic disorders characterized by skin fragility and blistering

25-50K/yr (US)1



Skin Aesthetics

\$22B Global Market²

>1MM aesthetic procedures/yr (US)²

Evolution of current RECELL platform required to incorporate cellular manipulation and/or genetically modified cells

Successful development of engineered (autologous) cell therapies will create a pathway to other applications