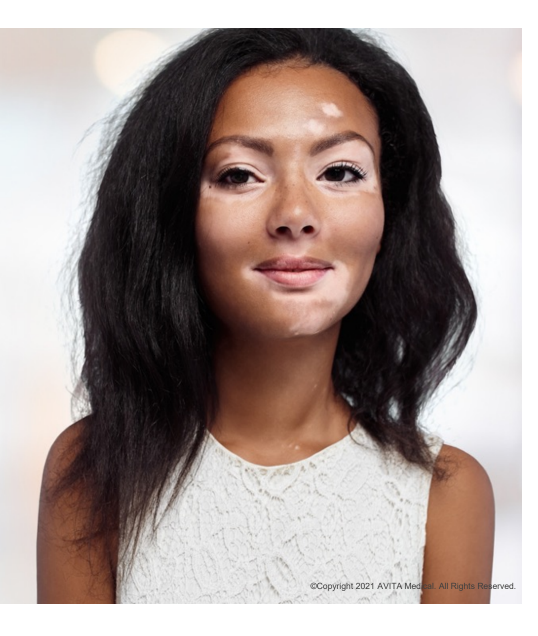
avita One Platform. Endless Possibilities.

November 2021

NASDAQ: RCEL

ASX: AVH



Legal Disclaimers

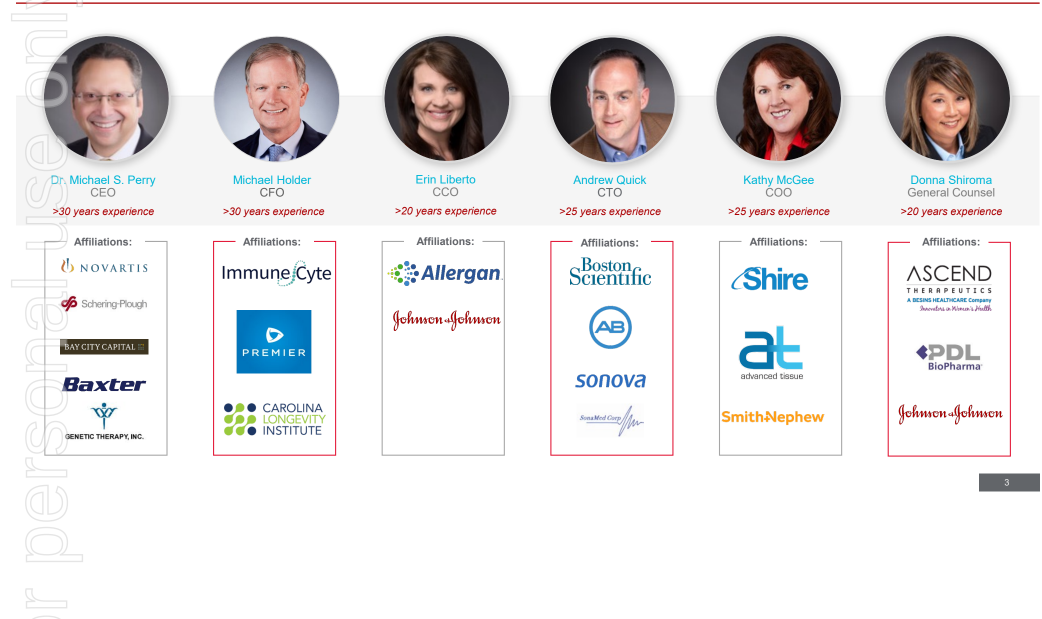
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Certain statements in this presentation and the accompanying oral commentary are "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995. All statements other than statements of historical facts contained in this presentation, including statements regarding our future financial condition, technology platform, development strategy, prospective products, pipeline and milestones, regulatory objectives, expected payments from and outcomes of collaborations, and likelihood of success, are forward-looking statements. Such statements are predictions only and involve known and unknown risks, uncertainties and other important factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements. These risks and uncertainties include, among others, the costs, timing and results of clinical trials and other development activities; the uncertainties inherent in the initiation and enrollment of clinical trials; the uncertainties associated with the COVD-19 pandemic; the unpredictability of the timing and results of regulatory submissions and reviews; market acceptance for approved products and innovative therapeutic treatments; competition; the possible mairment of, inability to obtain and costs of obtaining intellectual property rights; and possible safety or efficacy concerns, general business, financial and accounting risks and litigation. Because forward-looking statements are inherently subject to risks and uncertainties, some of which cannot be predicted or quantified and some of which are beyond our control, you should not rely on these forward-looking statements as predictions of future events. More information concerning us and such risks and uncertainties is available in our public filings with the U.S. Securities and Exchange Commission, including our most recent Quarterly Report on Form 10-K for the year ended June 30, 2021 and our

AVITA Medical's products are Rx only. Please reference the Instructions for Use for more information on indications, contraindications, warnings, precautions and adverse events.

In the United States, RECELL[®] is approved for use in patients suffering acute thermal burns. Use of RECELL in other patient populations is either prohibited by United States law or may be made available pursuant to a relevant investigational device exemption granted by the FDA (and likewise limited by United States law to investigational use only).

AVITA Leadership Team

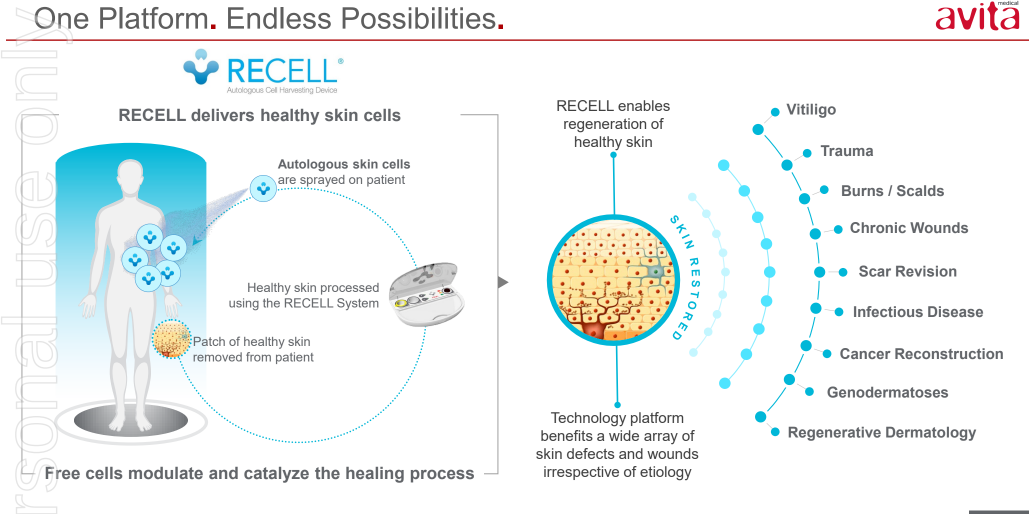


Value Creation

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	ecent Key Accomplishments	Projected Key Milestones
a 5	Soft Tissue Pivotal Trial: 89% Enrolled	Vitiligo Pivotal Trial Last Patient Enrolled / Q4 21 /
Y.	Vitiligo Pivotal Trial: Enrollment 70% Completed or	Vitiligo Commercial launch H2 23
α	Scheduled	Last patient enrolled in Soft Tissue Trial / Q1 22 / Soft Tissue Commercial Launch H2 23
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	Fiscal Q2'22 RECELL [®] revenue growth of +39% vs	PMDA Approval of Burns in Japan H1 22
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2	2018 FDA approval exceeding \$46M	EB: Initial proof of concept for delivery of genetically modified skin cells in suspension
-	FDA Approval of Pediatric label expansion	Telomerase/Rejuvenation: Initial proof of
	New Ease of Use RECELL Device Submitted to FDA for Review	concept on impact of telomerase on human skin in a mouse model

Quarters referenced in calendar year. As of January 1, 2022 Avita Medical will report on a calendar year basis.



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Focused Pipeline with Strong Growth Potential

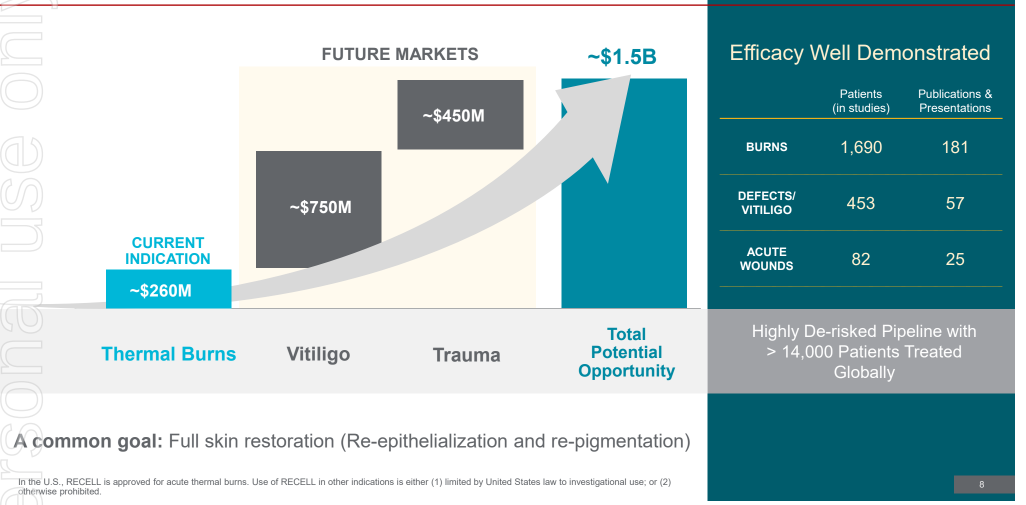
	DISCOVERY		FEASIBILITY		PIVOTAL		APPROVAL	-
Vounds &	Dermatology (C	urrent	Platform)					
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Innovation	CONCEPT	DESIGN	SUBMISSION	APPROVAL
New Device: Improved Ease of Use				
New Device: Fully Automated			:	

Focused Effort on Business Development to Supplement Pipeline

In the U.S., RECELL is approved for acute thermal burns. Use of RECELL in other indications is either (1) limited by United States law to investigational use; or (2) otherwise prohibited.

Current Platform Enables Access to a Large Serviceable Addressable Market (SAM)

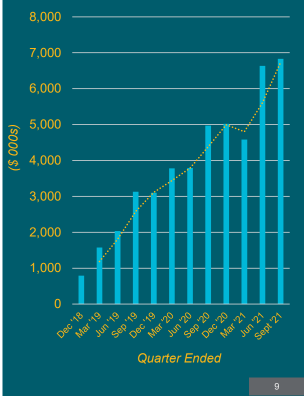


Thermal Burns: U.S. Target Market Expanded to Include Small Burns and Outpatient

Patient Funnel and Addressable Market ~486K PATIENTS Total Annual Burns in the U.S. *\$600 Million TAM ~80K PATIENTS Severe Burns (as defined as burns > 5% Body Surface Area that may require grafting) *\$260 Million SAM ~25K PATIENTS Target: Severe Burns Treated at Burn Centers (Both In and Outpatient)

Outpatient Pass Thru Code Opens Doors to Small Burns and Expands Serviceable Market Opportunity

U.S. RECELL Commercial Sales Since Approval





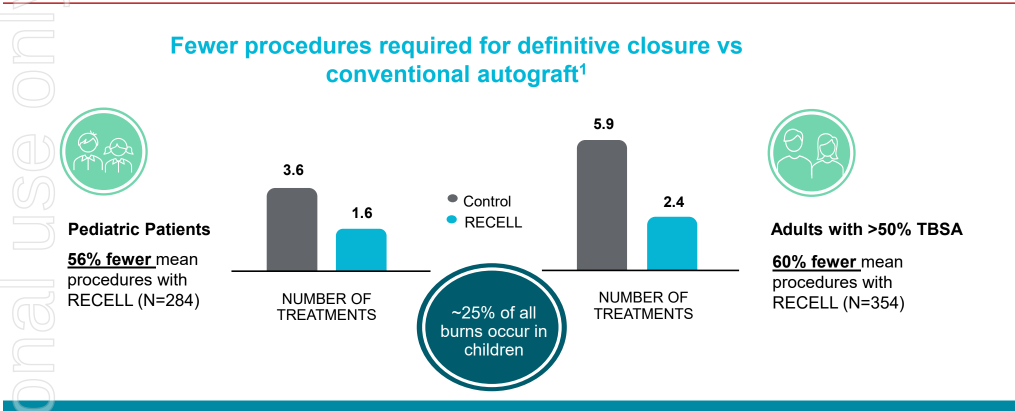
Strong Adoption of the RECELL System

avita

> \$46 Million in U.S. RECELL Revenue Since Approval

*Data is compiled based on information voluntarily provided by our customers and is subject to change.

Now Approved in Pediatric Full-Thickness Burns & Larger TBSAs avita



80% of RECELL Customers Stated that the New Label Enhancements Will Positively Impact Their Usage of RECELL

1. Instructions for Use. RECELL® Autologous Cell Harvesting Device * N = 41, "will significantly or somewhat impact RECELL usage"

11

Japan Is an Attractive Opportunity for AVITA Medical

Background

March 3, 2019, AVITA Medical and COSMOTEC Company, Ltd, an M3 Group company, announced agreement to market and distribute the RECELL System.

Based on feedback from the Japanese Health Authority (PMDA), the indication being pursued has been narrowed to focus on Burns given its robust randomized clinical data from the United States as well as local data in Japan.

RECELL System approval anticipated in Japan in H1 of CY 2022 followed by a reimbursement review with the Japanese Ministry of Health and Labour in June 2022. Commercial Launch will commence upon securing reimbursement.

Patient Funnel and Addressable Market

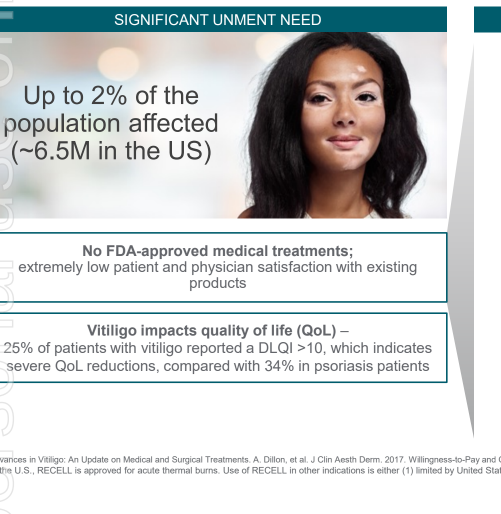


Approval Anticipated in H1 2022 with Commercial Launch mid-2022

Furue M, Yamazaki S, Jimbow K, Tsuchida T, Amagai M, Tanaka T et al. Prevalence of dermatological disorders in Japan: a nationwide, cross-sectional, seasonal, multi-center, hospital-based study. J Dermatol. 2011 April; 38(4):310-20, Japan Health System Review, 2018. Additional estimates based on data from 2016 JSBI National Burns Repository, https://injuryprevention.bmj.com/content/26/Suppl_2/i36#F2 and Costmotec estimates

Vitiligo: Unmet Need, No FDA-Approved Products

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LIMITED TREATMENT OPTIONS

Phototherapy

- 2-3 treatments / week for a few months to over a year
- Typically combined with a topical drug
- Not Durable

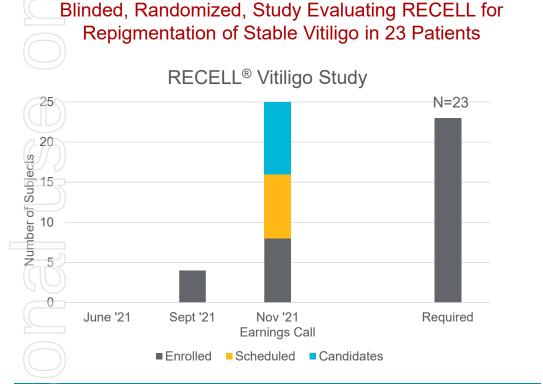
Melanocyte-Keratinocyte Transplantation

- For repigmentation of stable lesions
- Requires substantial laboratory equipment
- Performed rarely and only at 3 highly specialized academic centers in the United States

Advances in Vitiligo: An Update on Medical and Surgical Treatments. A. Dillon, et al. J Clin Aesth Derm. 2017. Willingness-to-Pay and Quality of Life in Patients with Vitiligo. Radtke, et al. BJD. 2009. In the U.S., RECELL is approved for acute thermal burns. Use of RECELL in other indications is either (1) limited by United States law to investigational use; or (2) otherwise prohibited.

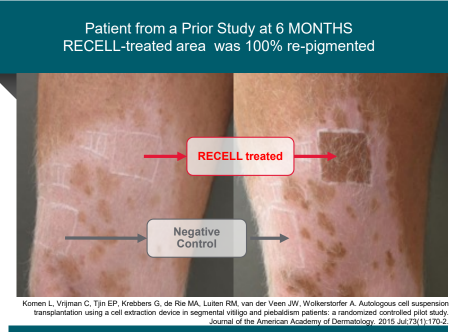
New Vitiligo Study Design Shortens Pathway to Completion

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U.S. Pivotal Study enrolling; last patient expected in H2 2021

In the United States, RECELL is not approved for treatment of vitiligo.



POTENTIAL RECELL BENEFITS

For Stable Vitiligo: Durable: One-time Segmental & Non-Segmental treatment

RECELL Case Study: Repigmentation of the Nipple-Areola Complex after Breast Treatment (B)

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Before RECELL®



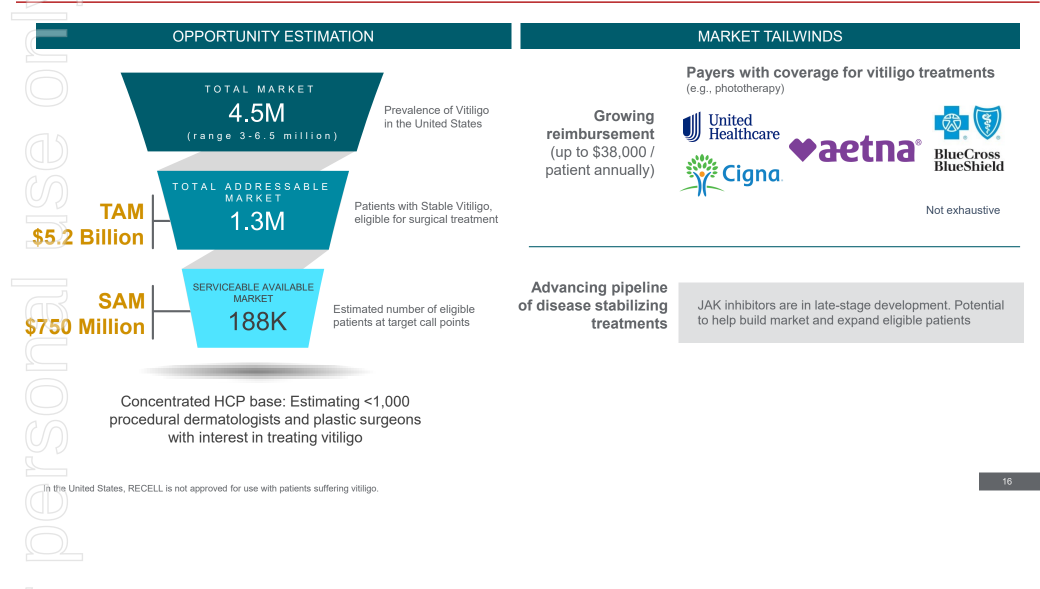
12 months After RECELL®

- 23 year old female with vitiligo.
- Donor skin was harvested from adjacent unaffected areas.
- Dermabrasion of the vitiligo patches was performed to the depth of the dermal-epidermal junction.
- The cellular suspension was then sprayed on both the recipient and donor areas (expansion ratio ranged from 1:20-1:40).

Established Track Record in Vitiligo: 1,000 patients treated internationally & 12 peer reviewed publications showing positive outcomes

Yu et al. Repigmentation of nipple-areola complex after RECELL® treatment on breast vitiligo. Journal of Cosmetic Dermatology, 2021 In the United States, RECELL is not approved for use with patients suffering vitiligo.

Significant Market Opportunity in Repigmenting Stable Vitiligo



Soft Tissue Reconstruction Trial Enrollment is Gaining Momentum

20

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Clinical trial demonstrates use of less donor skin without compromising healing outcomes relative to conventional autografting

N=65

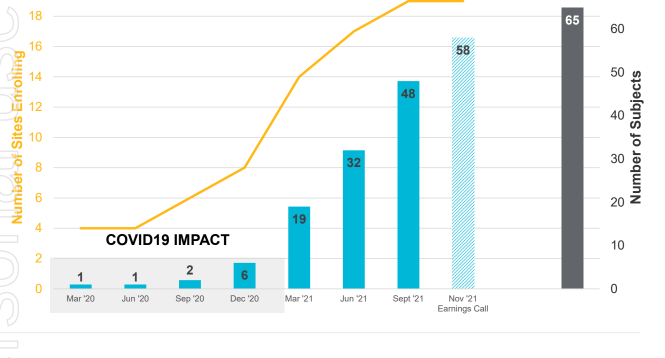
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Patient treated for necrotizing fasciitis.





Photos courtesy of Kevin Foster, Valleywise Health Medical Center

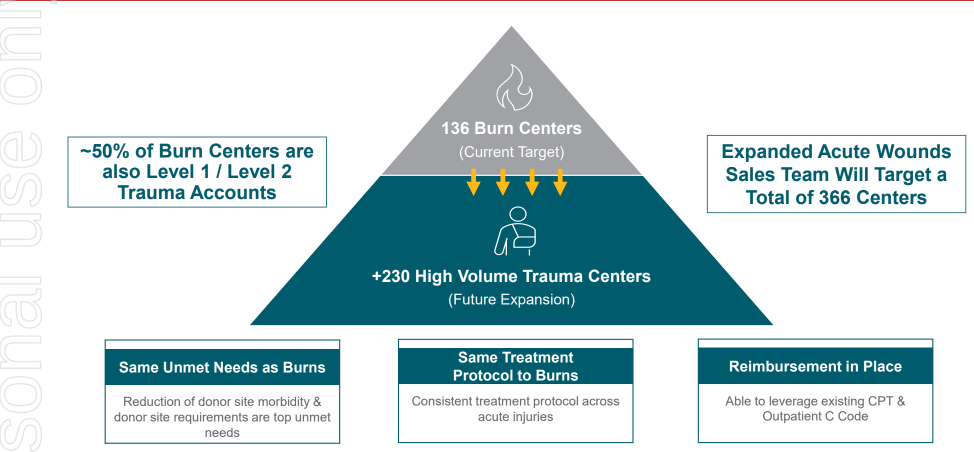


RECELL[®] Soft Tissue Study

In the U.S., RECELL is approved for acute thermal burns. Use of RECELL in other indications is either (1) limited by United States law to investigational use; or (2) otherwise prohibited.

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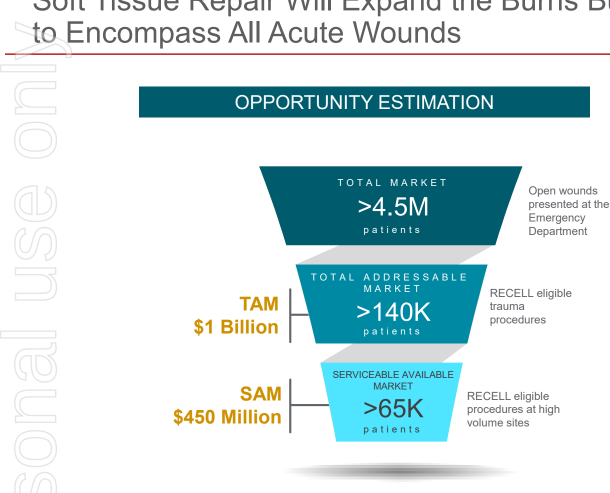




In the U.S., RECELL is approved for acute thermal burns. Use of RECELL in other indications is either (1) limited by United States law to investigational use; or (2) otherwise prohibited. In the United States, RECELL is not approved for use in pediatrics. Use of RECELL in this case was performed internationally where the indication is approved.

Soft Tissue Repair Will Expand the Burns Business

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Female, pregnant 28 year old who suffered from a de-gloving Injury

Post Debridement of Injury



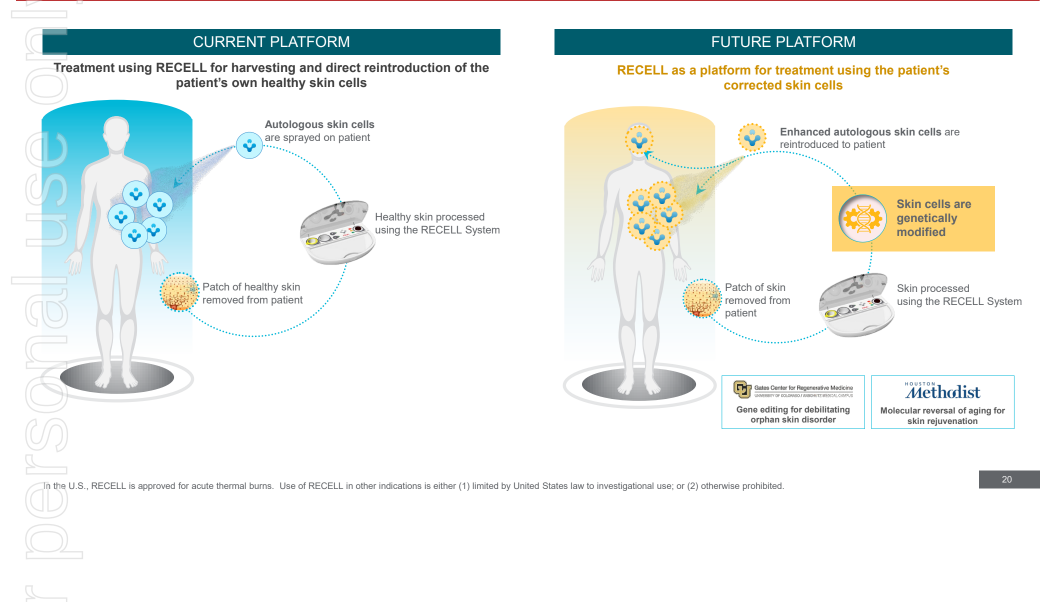
6 MONTH POST-RECELL TREATMENT



In the U.S., RECELL is approved for acute thermal burns. Use of RECELL in other indications is either (1) limited by United States law to investigational use; or (2) otherwise prohibited. In the United States, RECELL is not approved for use in pediatrics. Use of RECELL in this case was performed internationally where the indication is approved.

Poster: Use of regenerative suspension in the treatment of a complex de-gloving injury. Ian M Smith,

RECELL in Genetic Skin Defects and Rejuvenation



Exploring Cell-Based Gene Therapy for Epidermolysis Bullosa

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Preclinical research partnership underway with Gates Center for Regenerative Medicine (University of Colorado), exploring the combination of a novel gene correction approach with AVITA Medical's Spray-On Skin[™] Cells technology

THE CHALLENGE

DEBILITATING Skin fragility, disability, cancer

HIGH UNMET NEED

No FDA-approved treatment, only palliative measures

COST BURDEN Care of \$200K-\$500K per year per patient



THE OPPORTUNITY

CURATIVE: Technology for precise correction of genetic defect & banking for future use (vs ameliorating symptoms)



EFFICIENT: Suspension-based approach eliminates growth & transport of fragile skin sheets



CONVENIENT: Suspension-based product simplifies application onto patient wounds (vs surgical anchoring of epidermal sheets which can result in issues with "take rates"

Proof-of-concept for delivering genetically modified cells in suspension expected in 2021

Th the U.S., RECELL is approved for acute thermal burns. Use of RECELL in other indications is either (1) limited by United States law to investigational use; or (2) otherwise prohibited.

Exploring Novel RNA-Based Approach for Rejuvenation

Patented RNA technology for delivery of telomerase enzyme to aged cells
 Demonstrated reversal of aging and return of functionality in cells of progeria patients (human model of accelerated aging)

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- Patented and proprietary **Spray-On Skin Cells technology and device (RECELL)**
- Expertise in skin regeneration, including in preclinical models
- Strong track record and expertise in clinical development and commercialization

Multi-Billion Dollar Market Presents a Sizeable Opportunity

- >\$16.5B spent in aesthetic procedures per year (US)*
- >3M aesthetic procedures per year (US) aimed to improve skin tightness, texture & evenness in skin tone*
- Consumers desire superior results over current offerings
- Personalized, cellular-level approaches to skin rejuvenation, developed with robust evidence, is an area of significant interest

Sponsored research underway exploring use of telomerase for molecular reversal of skin cell aging

*American Society for Plastic Surgery Annual reports – 2018 and 2019. 2. Goddard et al. Aesthetic Surgery Journal, Volume 40, Issue 4, April 2020, Pages 460–465. In the U.S., RECELL is approved for acute thermal burns in patients > 18 years. Use of RECELL in other indications is either (1) limited by United States law to Investigational use; or (2) otherwise prohibited.





2019 29 5,474 	2020 14,263	2021 21,483	\$19.6 Share F		
	14,263	21 483			
	1	21,400			
	-	7,749	4 4 0 0 1		
29 5,474	14,263	29,232		\$480 Million Market Capitalization ¹	
83 4,203	11,290	23,283			
734 5,921	3,926	2,055	\$0.1	0	
.986 20,174	73,639	110,746		(Zero) Debt	
Analysts			Nasdaq ticker	ASX ticker	
Lyanne Harrison, BofA Global R	Research (AUS) • John H	lester, Bell Potter (AUS)	symbol: RCEL	symbol: AVH	
,	83 4,203 734 5,921 ,986 20,174 Analysts Brooks O'Neil, Lake Street (U.S Lyanne Harrison, BofA Global R	883 4,203 11,290 734 5,921 3,926 ,986 20,174 73,639 Analysts Brooks O'Neil, Lake Street (U.S.) Lyanne Harrison, BofA Global Research (AUS)	883 4,203 11,290 23,283 734 5,921 3,926 2,055 ,986 20,174 73,639 110,746 Analysts Brooks O'Neil, Lake Street (U.S.) Lyanne Harrison, BofA Global Research (AUS) • Chris Kallos, MST (AUS) • John Hester, Bell Potter (AUS) •	Analysts Analysts Brooks O'Neil, Lake Street (U.S.) Lyanne Harrison, BofA Global Research (AUS) • Chris Kallos, MST (AUS) • John Hester, Bell Potter (AUS)	

Financial Overview

A Global Total of 56 Granted Patents & 26 Pending Applications

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ROBUST PROTECTION ACROSS PATENT FAMILIES						
Cell Suspension Preparation Technique and Use	Commercial RECELL device, composition of matter, and associated methods of use					
Cell Suspension And Use Thereof	Method of preparing cell suspension with exogenous agent to promote wound healing					
Systems and Methods for Tissue Processing and Preparation of Cell Suspension Therefrom	Automated system for preparing cell suspension and method of production					
Devices, Methods, and Kits for Preparing a Cell Suspension	All-in-one RECELL kit, system, and associated method of use					
Methods for Identifying Cell Suspensions with Therapeutic Potential for Skin Regeneration	Method and system for validating the use of a cell suspension for administration to a patient					
Bioactive Therapeutic Suspensions with Cellular-Based Supernatant	Bioactive suspension derived from freshly disaggregated tissue, and associated methods of preparation and use					

EXPANDING PORTFOLIO TO SUPPORT CURRENT AND FUTURE INDICATIONS



Next Generation RECELL devices to improve ease of use in burns and pipeline indications



Potential to license patented technology for telomerase mRNA that has the potential to reverse aging of skin cells



Potential to license technologies for suspensionbased delivery of genetically modified cells, with applications to genetic skin disorders

Robust and Expanding Patent Estate: Expiration from 2022 to 2040

Note: AVITA Medical owns granted patents in Australia, Australia, Belgium, Brazil, France, Germany, Hong Kong, Italy, Japan, Netherlands, Portugal, Spain, Sweden, Turkey, United Kingdom and USA. AVITA Medical owns pending patent applications in Brazil, Canada, China, Europe, Hong Kong and USA. Patent count as of 6/30/2021

Value Creation

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	ecent Key Accomplishments	Projected Key Milestones
a 5	Soft Tissue Pivotal Trial: 89% Enrolled	Vitiligo Pivotal Trial Last Patient Enrolled / Q4 21 /
Y.	Vitiligo Pivotal Trial: Enrollment 70% Completed or	Vitiligo Commercial launch H2 23
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Quarters referenced in calendar year. As of January 1, 2022 Avita Medical will report on a calendar year basis.

Risk Factors and Disclosures

- 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 in the United States as well as 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.

Important Safety Information

- INDICATIONS FOR USE: The RECELL® Autologous Cell Harvesting Device is indicated for the treatment of acute thermal burn wounds. The RECELL device is used by an appropriately-licensed healthcare professional at the patient's point of care to prepare autologous RES® Regenerative Epidermal Suspension for direct application to acute partial-thickness thermal burn wounds in patients 18 years of age and older or application in combination with meshed autografting for acute full-thickness thermal burn wounds in pediatric and adult patients.
- CONTRAINDICATIONS: RECELL is contraindicated for: the treatment of wounds clinically diagnosed as infected or with necrotic tissue, the treatment of patients with a known hypersensitivity to trypsin or compound sodium lactate (Hartmann's) solution, patients having a known hypersensitivity to anesthetics, adrenaline/epinephrine, povidone-iodine, or chlorhexidine solutions.
- WARNINGS: Autologous use only. Wound beds treated with a cytotoxic agent (e.g., silver sulfadiazine) should be rinsed prior to application of the cell suspension. RECELL is provided sterile and is intended for single-use. Do not use if packaging is damaged or expired. Choose a donor site with no evidence of cellulitis or infection and process skin immediately. A skin sample should require between 15 and 30 minutes contact with Enzyme. Contact in excess of 60 minutes is not recommended. RECELL Enzyme is animal derived and freedom from infectious agents cannot be guaranteed.
- PRECAUTIONS: RECELL is not intended for use without meshed autograft for treatment of full-thickness burn wounds. The safety and effectiveness of RECELL without
 meshed autograft have not been established for treatment of partial-thickness burn wounds: on the hands and articulating joints, >320 cm2, in patients with wounds
 totaling >20% total body surface area (TBSA). The safety and effectiveness of RECELL with autografting have not been established for treatment of full-thickness burn
 wounds: on the hands and articulated joints, and in patients younger than 28 days of age (neonates).
 - SPECIAL PATIENT POPULATIONS: The safety and effectiveness of RECELL have not been established for treatment of acute thermal partial-thickness burn wounds in pediatric patients younger than 18 years of age.

Revolutionary treatment using a patient's own skin for life-changing outcomes avita

the U.S., RECELL is approved for acute thermal burns. Use of RECELL in other indications is either (1) limited by United States law to investigational use; or otherwise prohibited.

Zed, treated with the RECELL® System