



Advanced Therapies for the Sports Medicine & Severe Burn Care Markets

CORPORATE PRESENTATION

MARCH 2024

Forward-Looking Statements and Legal Disclosure

Forward-Looking Statements

Vericel cautions you that all statements other than statements of historical fact included in this presentation that address activities, events or developments that we expect, believe or anticipate will or may occur in the future are forward-looking statements. Although we believe that we have a reasonable basis for the forward-looking statements contained herein, they are based on current expectations about future events affecting us and are subject to risks, assumptions, uncertainties and factors relating to our operations and business environment, all of which are difficult to predict and many of which are beyond our control. Our actual results may differ materially from those expressed or implied by the forward-looking statements in this presentation. These statements are often, but are not always, made through the use of words or phrases such as “anticipates,” “intends,” “estimates,” “plans,” “expects,” “continues,” “believe,” “guidance,” “outlook,” “target,” “future,” “potential,” “goals” and similar words or phrases, or future or conditional verbs such as “will,” “would,” “should,” “could,” “may,” or similar expressions.

Among the factors that could cause actual results to differ materially from those set forth in the forward-looking statements include, but are not limited to, uncertainties associated with our expectations regarding future revenue, growth in revenue, market penetration for MACI®, Epicel®, and NexoBrid®, growth in profit, gross margins and operating margins, the ability to continue to scale our manufacturing operations to meet the demand for our cell therapy products, including the timely completion of a new headquarters and manufacturing facility in Burlington, Massachusetts, the ability to achieve or sustain

profitability, contributions to adjusted EBITDA, the expected target surgeon audience, potential fluctuations in sales and volumes and our results of operations over the course of the year, timing and conduct of clinical trial and product development activities, timing and likelihood of the FDA’s potential approval of the arthroscopic delivery of MACI to the knee or the use of MACI to treat cartilage defects in the ankle, the estimate of the commercial growth potential of our products and product candidates, competitive developments, changes in third-party coverage and reimbursement, physician and burn center adoption of NexoBrid, supply chain disruptions or other events or factors affecting MediWound’s ability to manufacture and supply sufficient quantities of NexoBrid to meet customer demand, including but not limited to the ongoing Israel-Hamas war, negative impacts on the global economy and capital markets resulting from the conflict in Ukraine and the Israel-Hamas war, adverse developments affecting financial institutions, companies in the financial services industry or the financial services industry generally, global geopolitical tensions or record inflation and potential future impacts on our business or the economy generally stemming from a resurgence of COVID-19 or another similar public health emergency.

These and other significant factors are discussed in greater detail in Vericel’s Annual Report on Form 10-K for the year ended December 31, 2023, filed with the Securities and Exchange Commission (SEC) on February 29, 2024, and in other filings with the SEC. These forward-looking statements reflect our views as of the date hereof and Vericel does not assume and specifically disclaims any obligation to update any of these forward-looking

statements to reflect a change in its views or events or circumstances that occur after the date of this release except as required by law.

Discussion of Indications Currently Under Development

Additionally, portions of this presentation discuss the potential clinical advantages of the arthroscopic delivery of MACI to treat cartilage defects in the knee joint and the use of MACI in the ankle joint, as well as the potential effect the approval of those additional indications could have on MACI’s total addressable market. The reader is reminded that the implantation of MACI in the knee is currently approved to be performed via an arthrotomy. The arthroscopic delivery of MACI to the knee joint and the use of MACI in the ankle joint are currently under development and such uses have not been approved in the U.S.

Vericel is a Leader in Advanced Therapies in Sports Medicine and Burn Care, Combining Innovations in Biology with Medical Technologies



Our Vision

Every patient benefits from therapies as unique as they are



Our Mission

We provide precision therapies that repair injuries and restore lives

SPORTS MEDICINE



autologous cultured
chondrocytes
on porcine
collagen membrane

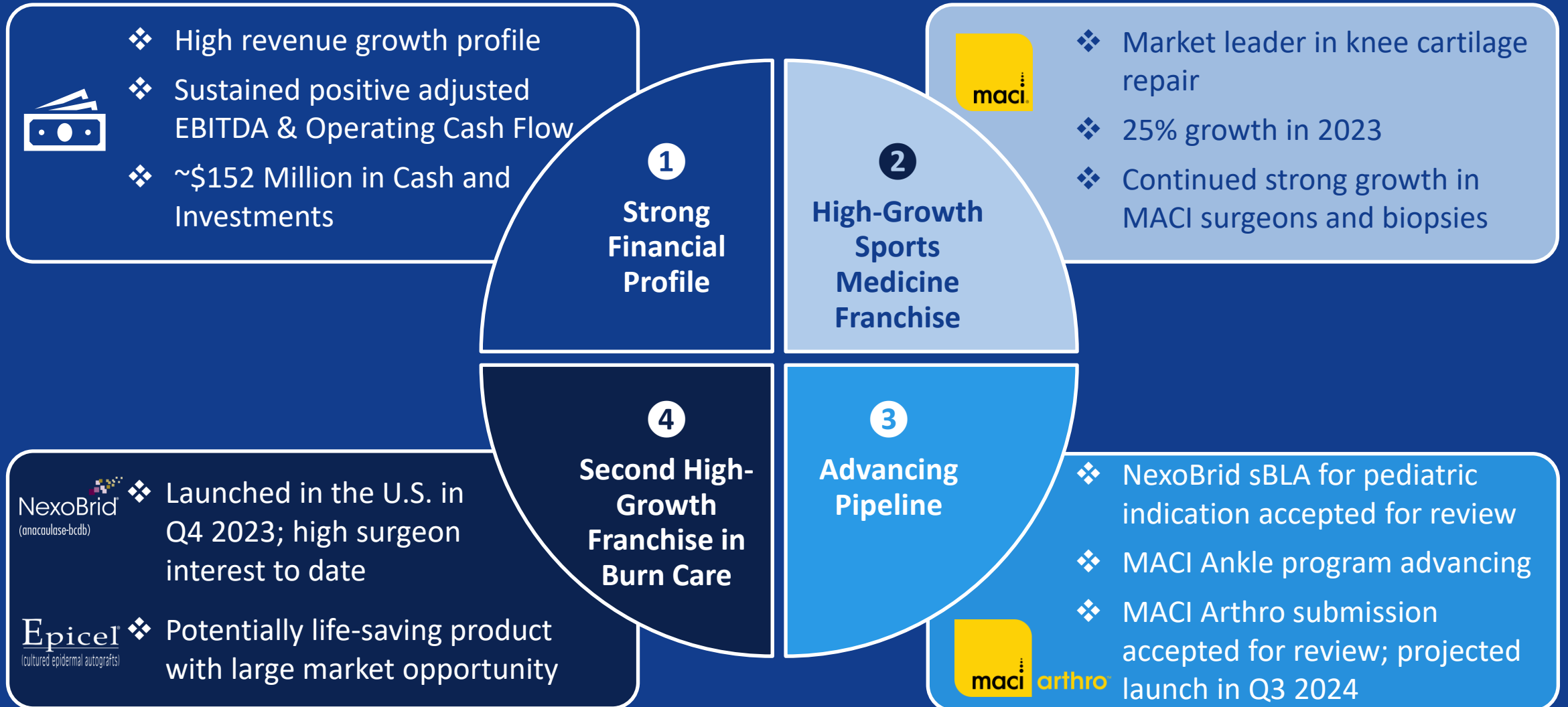
SEVERE BURNS

Epice^l
(cultured epidermal autografts)

NexoBrid[®]
(anacaulase-bcdb)

Portfolio of Innovative Cell Therapies and Specialty Biologics
with Significant Barriers to Entry

Vericel is Well-Positioned to Deliver Sustained Long-Term Growth



Large Underpenetrated Markets with Total Addressable Market Opportunity Expanding to Over \$4.5 Billion in the Years Ahead



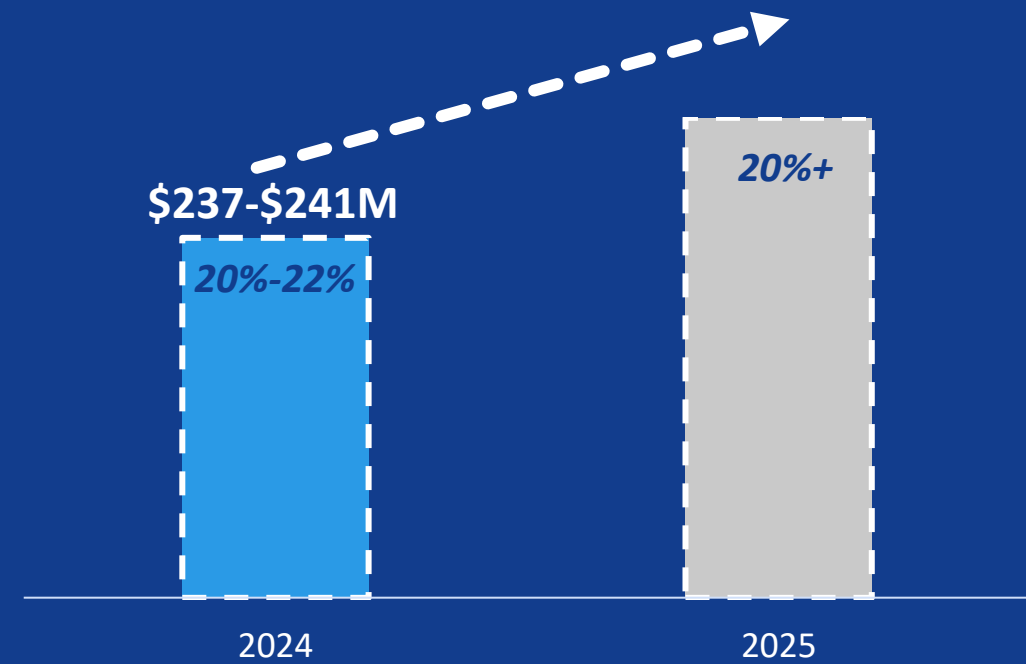
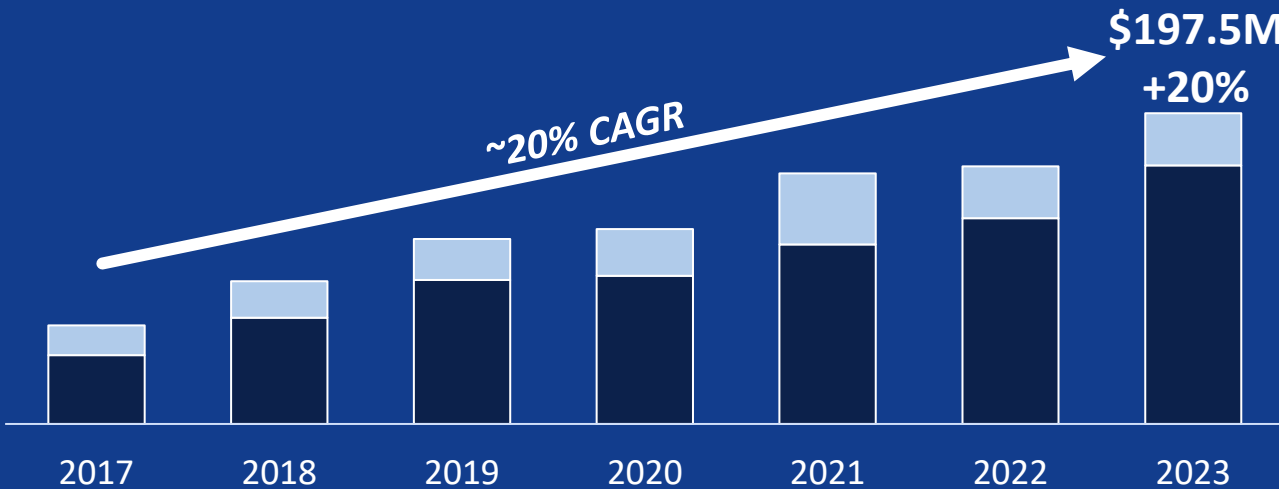
- ❖ NexoBrid launched in Q4 2023
- ❖ MACI Arthro targeting largest segment of current TAM and expected to launch in Q3 2024
- ❖ MACI Ankle trial anticipated to initiate in 2025

Core Portfolio Plus New Product Launches Expected to Drive Further Strong Revenue Growth in 2024 and Beyond



Top-Tier Revenue Growth

□ Sports Med ■ Burn Care



Durable Growth Platform

- ❖ Significantly underpenetrated markets
- ❖ Limited competition with strong barriers to entry
- ❖ Strong reimbursement profiles

With Additional Growth Drivers in 2024+

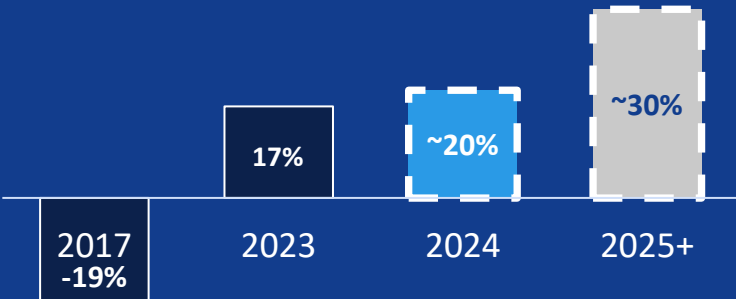
- ❖ First full year of NexoBrid revenue in 2024
- ❖ MACI Arthro launch in Q3 2024, with first full year of revenue in 2025

2024 and 2025+ estimates based on internal financial projections.

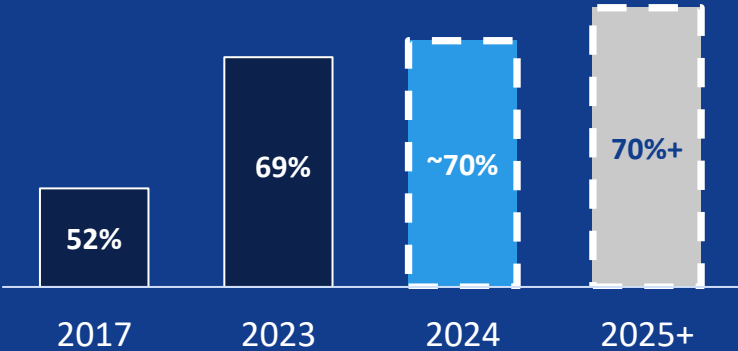
Driving High Revenue Growth with a Top-Tier Profitability Profile

20% Top Line Growth in 2023
Expect Continued 20%+ Growth in 2024+

Adjusted EBITDA Margin %



Gross Margin %



40% Adjusted EBITDA Growth in 2023
Expect Strong Adjusted EBITDA Growth in 2024+

2024 and 2025+ estimates are based on internal financial projections.
Adjusted EBITDA margin trending towards 30% in 2025 and approaching long-term goal of 30% or more beyond 2025

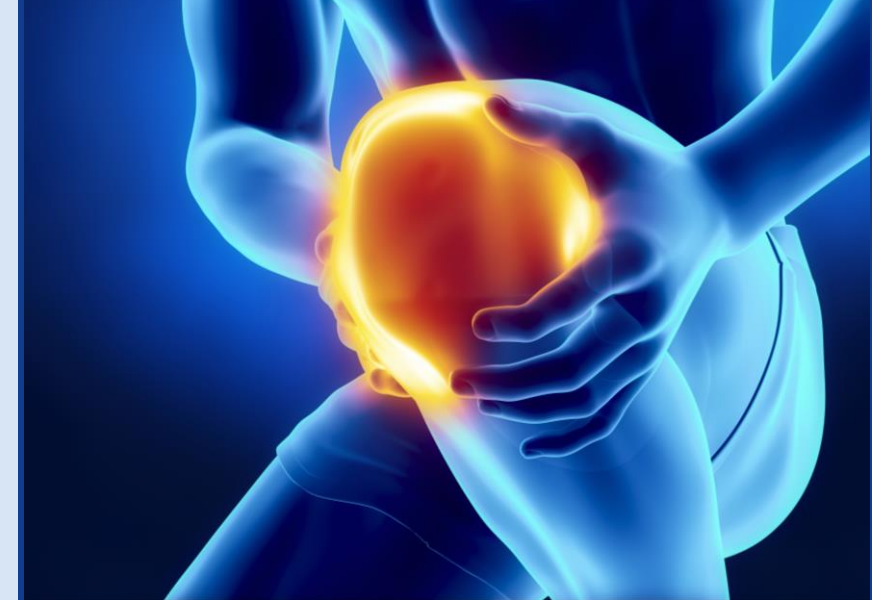
Knee Cartilage Injuries Represent a Significant Unmet Medical Need

Cartilage defects are found in ~60% of knee arthroscopies¹

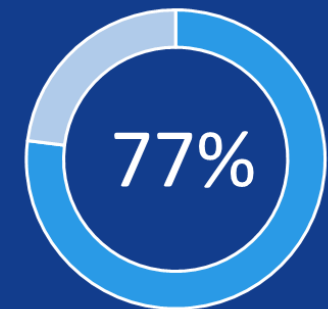
- ❖ Damage is caused by acute or repetitive trauma or degenerative conditions

Cartilage has limited capacity for intrinsic healing and repair

- ❖ Untreated cartilage defects may lead to debilitating joint pain, dysfunction, and osteoarthritis
- ❖ Defects can expand and new high-grade lesions can form over time



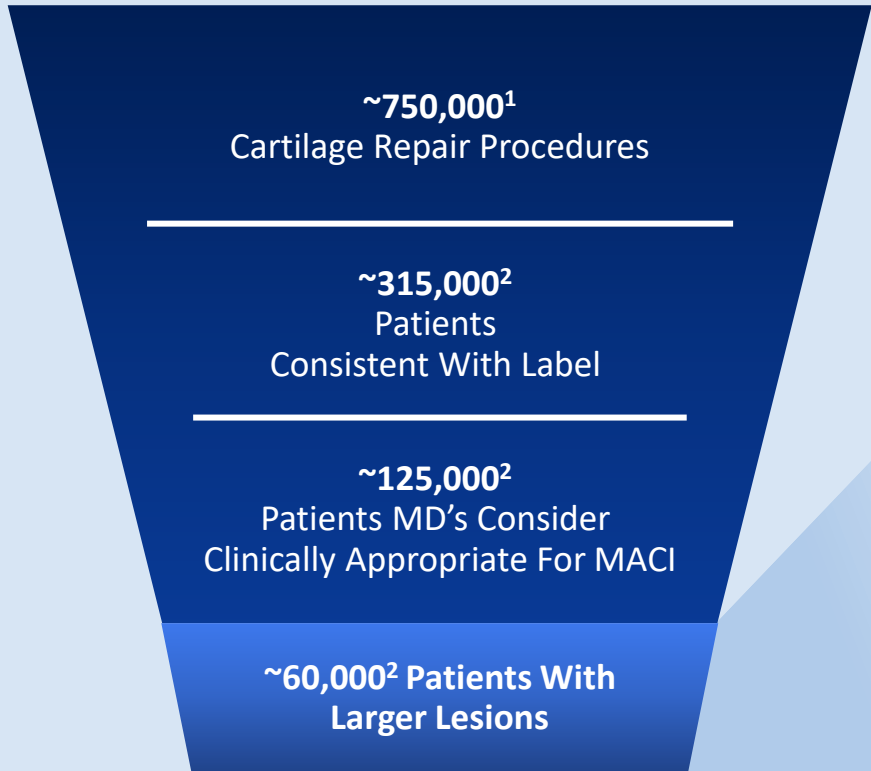
Impact of Knee Pain



Harris Poll found that 77% of knee pain sufferers can no longer participate in at least one activity they previously enjoyed because of knee pain²

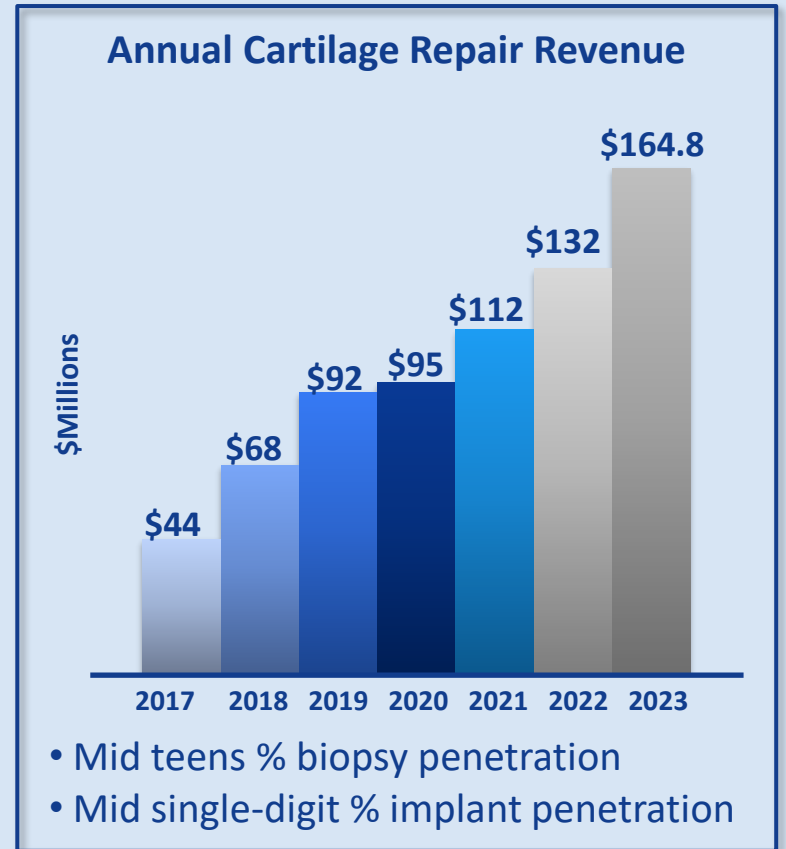
Large Addressable Knee Cartilage Repair Market for MACI

Estimated Annual Addressable Patient Population (U.S.)



\$3 Billion
Addressable Market
in the U.S.³

Annual Cartilage Repair Revenue



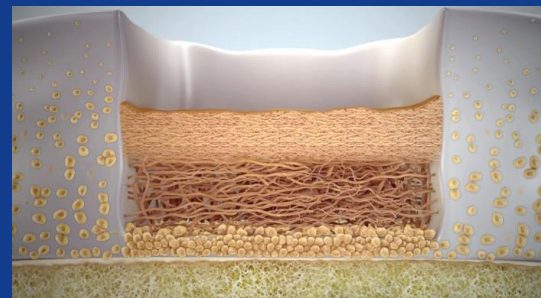
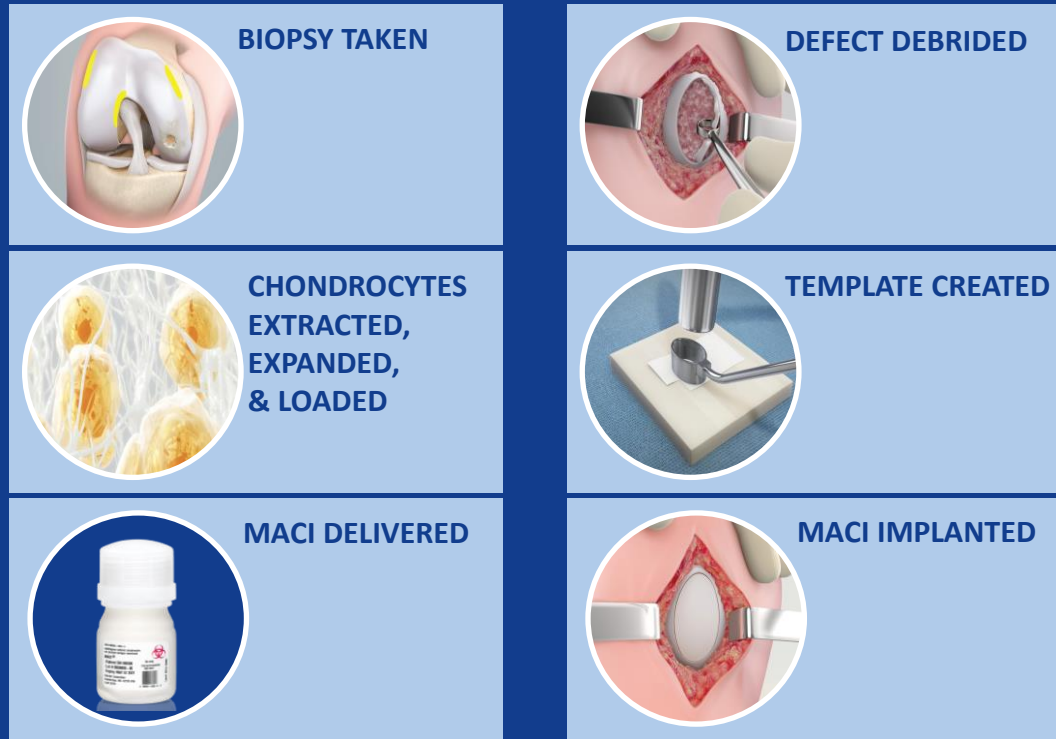
¹ Health Advances LLC MACI market assessment report (2018), Vericel data, LexisNexis, Medtech Insight, NY SASD, SmartTRAK, LSI, PSPS, McCormick, Frank et al. Arthroscopy, (2014) 30(2): 222-6, Montgomery, et al. Knee Surg Sports Traumatol Arthrosc (2014) 22: 2070.

² Health Advances LLC MACI market assessment report (2018).

³ Assumes MACI ASP of ~\$50,000+.



MACI is the Leading Restorative Cartilage Repair Product on the Market



MACI Product Attributes Driving Strong Growth Since Launch

Broad Label with Strong Clinical Data

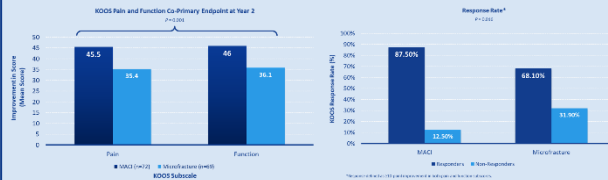
HIGHLIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use MACI safely and effectively. See full prescribing information for MACI.

MACI[®] (autologous cultured chondrocytes on porcine collagen membrane)
Cellular sheet for autologous implantation
Initial U.S. Approval: 2016

INDICATIONS AND USAGE

MACI[®] is an autologous cellularized scaffold product indicated for the repair of symptomatic, single or multiple full-thickness cartilage defects of the knee with or without bone involvement in adults. (1)
Limitations of Use

SUMMIT Clinical Study Results – Superiority of MACI Implant Versus Microfracture Treatment

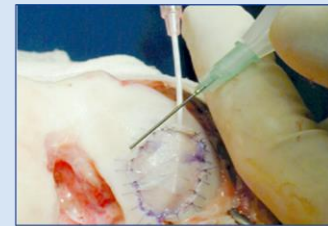


MACI demonstrated statistically significantly greater improvement in the co-primary endpoint of KOOS pain and function (SRA) scores compared to microfracture at year 2.

The proportion of patients responding to treatment was statistically significantly greater with MACI compared to microfracture at year 2.

*The American Journal of Sports Medicine 2016;43(10):1981-1994

Simpler, Less Invasive Procedure



Carticel

- ▷ Technically exacting procedure
- ▷ Required arthrotomy, periosteal patch harvest and sutures
- ▷ Extended surgical time



MACI

- ▷ Simpler, less invasive ACI procedure
- ▷ Eliminates periosteal harvest and sutures
- ▷ Significant reduction in surgical time
- ▷ Uniform distribution of cells
- ▷ Improved post-operative course

Shorter Rehab Protocols

ACHIEVE ROUTINE
0-3 months following surgery

BUILD STRENGTH
3-6 months following surgery

BE ACTIVE
6-9 months following surgery

Rehabilitation Timelines for ACI procedures: Time to Weight-Bearing¹



Published MACI rehabilitation protocols achieve full weight-bearing in 6-8 weeks compared to 10-12 weeks for published Carticel rehabilitation protocols

¹Libert J et al. Osteoarthritis & Cartilage 2008; Edwards PK et al. AISM 2013.

Strong Reimbursement Profile

MACI Insurance Approval Rates

89% of all MACI surgeries were approved by the insurer on initial submission

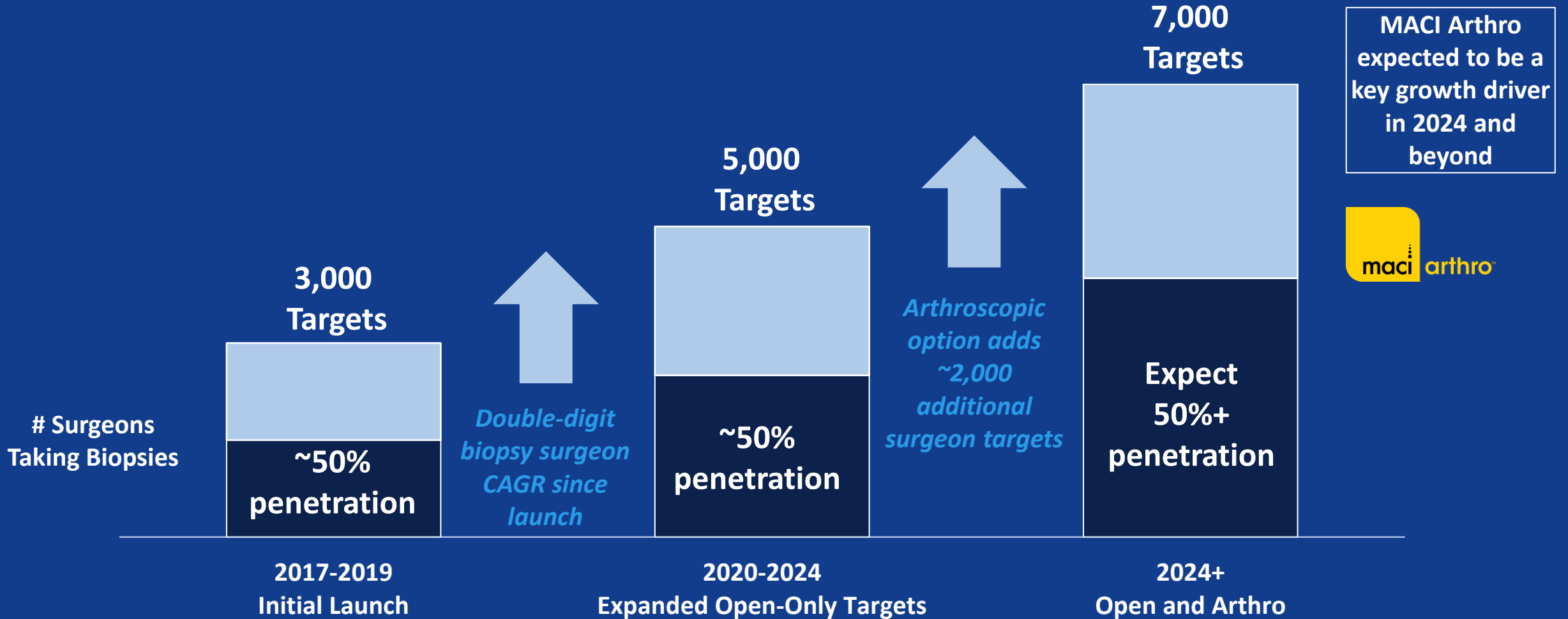
- 5% Approved on appeal
- 5% Not appealed
- 1% Denied after appeal

MACI Billing Codes

ACI CPT CODE	Code
Autologous Chondrocyte Implantation, Knee	27412

HCPCS CODES	Code
Autologous cultured chondrocytes, Implant	J7330

Surgeon Adoption Continues to be a Key MACI Growth Driver and Target Surgeons Will Increase With MACI Arthro Launch in 2024



Building a Robust and Innovative Pipeline Through Lifecycle Management and Business Development

PRODUCT	INDICATION/STUDY	IN DEVELOPMENT	PHASE I	PHASE II	PHASE III	REGISTRATION	APPROVAL
 MACI <small>autologous cultured chondrocytes on porcine collagen membrane</small>	Treatment of Symptomatic Cartilage Defects of the Knee in Adults	Commercialized					
	Pediatric (PEAK) Study – Knee	Currently Enrolling					
	Arthroscopic Delivery – Knee				Submission Accepted for Review		
	Treatment of Cartilage Defects – Ankle				Study Pending		
 Epiceel <small>(cultured epidermal autografts)</small>	Treatment of Large Deep Dermal and Full-Thickness Burns	Commercialized					
	Burn Eschar Removal in Adults	Commercialized					
 NexoBrid <small>(anacaulose-bctdb)</small>	Pediatric (CIDS) Study	sBLA Accepted for Review					
	Treatment of Acute Deep Partial and Full Thickness Burn Injuries (NEXT) Study	Expanded Access (Pediatrics)					

Key Highlights

MACI Arthroscopic Delivery

- ❖ MACI Arthro submission accepted for review; projected launch in Q3 2024

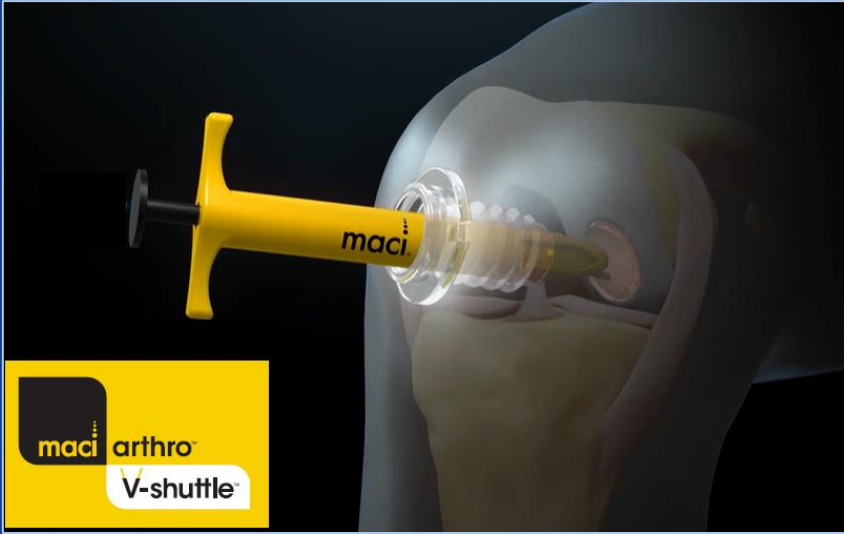
MACI Ankle Indication

- ❖ Trial anticipated to initiate in 2025

NexoBrid

- ❖ Launched in Q4 2023
- ❖ sBLA for pediatric indication accepted for review

MACI Arthro



Arthroscopic MACI Delivery Provides a Significant Growth Opportunity

High Surgeon Interest in MACI Arthro

~90% % of target surgeons expressed **Interest** in arthro MACI option¹

Potential for Increased MACI Volume

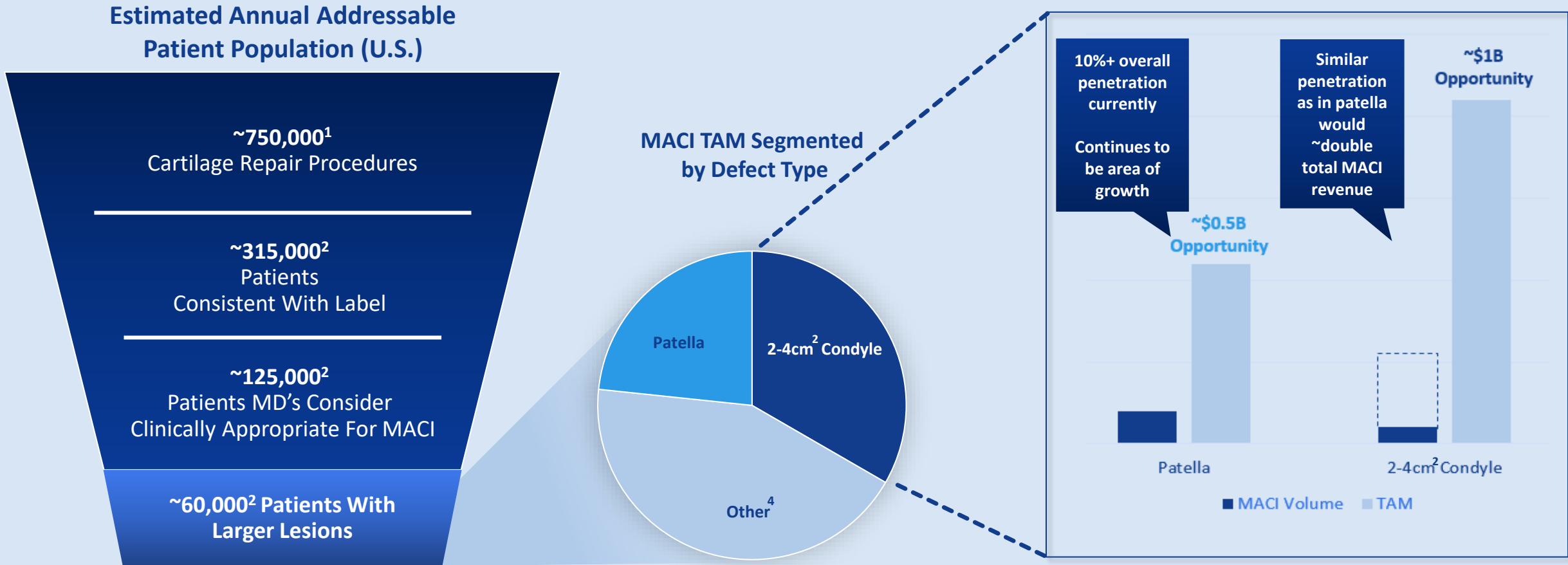
~90% % of current MACI users would expect to **Increase** MACI volume¹

Upon Launch MACI Arthro Will be the Only Restorative Cartilage Repair Product That Can be Administered Arthroscopically

[Click here to view an animation of the MACI arthroscopic delivery surgical technique](#)

¹Based on Health Advances, LLC MACI market assessment report (2018).

Arthroscopic MACI is Targeting 2-4cm² Femoral Condyle Defects, Which Represents the Largest Portion of the MACI Addressable Market



¹ Health Advances LLC MACI market assessment report (2018), Vericel data, LexisNexis, Medtech Insight, NY SASD, SmartTRAK, LSI, PSPS, McCormick, Frank et al. Arthroscopy, (2014) 30(2): 222-6, Montgomery, et al. Knee Surg Sports Traumatol Arthrosc (2014) 22: 2070.

² Health Advances LLC MACI market assessment report (2018).

³ Assumes MACI ASP of ~\$50,000+.

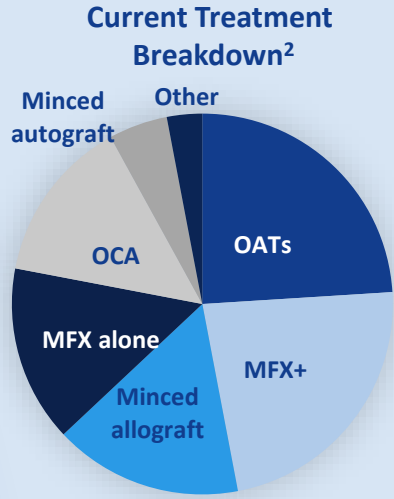
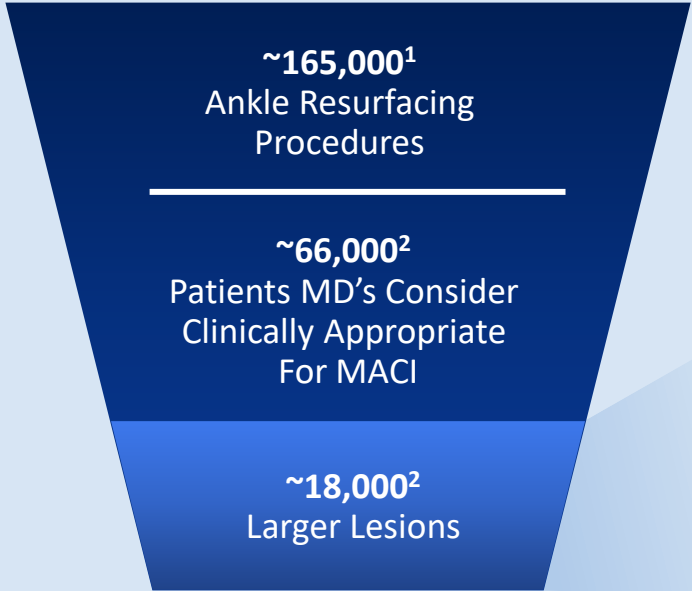
⁴ Includes defects on tibia, trochlea and other condyle defects.



Significant Ankle Cartilage Repair Opportunity



MACI Ankle Annual TAM Estimate (U.S.)

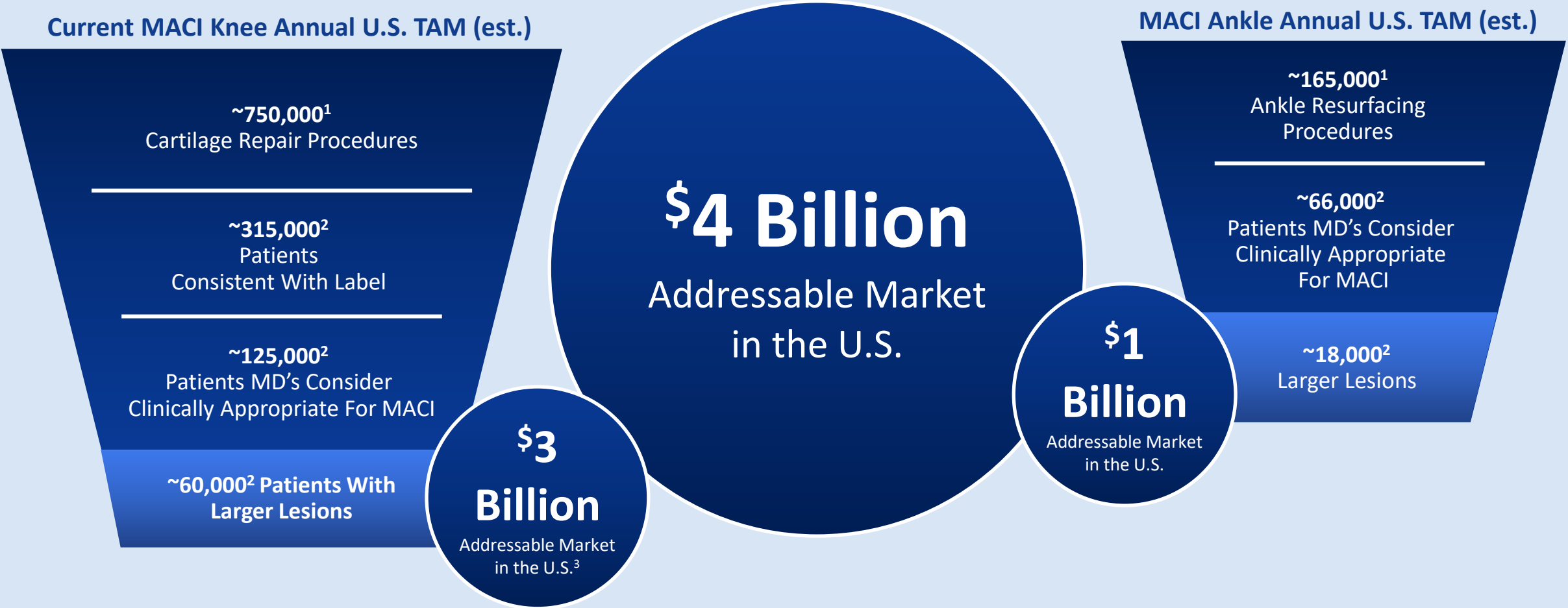


MACI for the treatment of cartilage defects in the ankle represents a \$1 billion³ market opportunity



¹ SmartTrak Cartilage Repair Procedures; resurfacing includes microfracture, OATs, OCA, etc. and does not include chondroplasty/debridement only.
² Cello Health MACI Ankle quantitative market research survey (2021).
³ Assumes MACI ASP of \$50,000+.

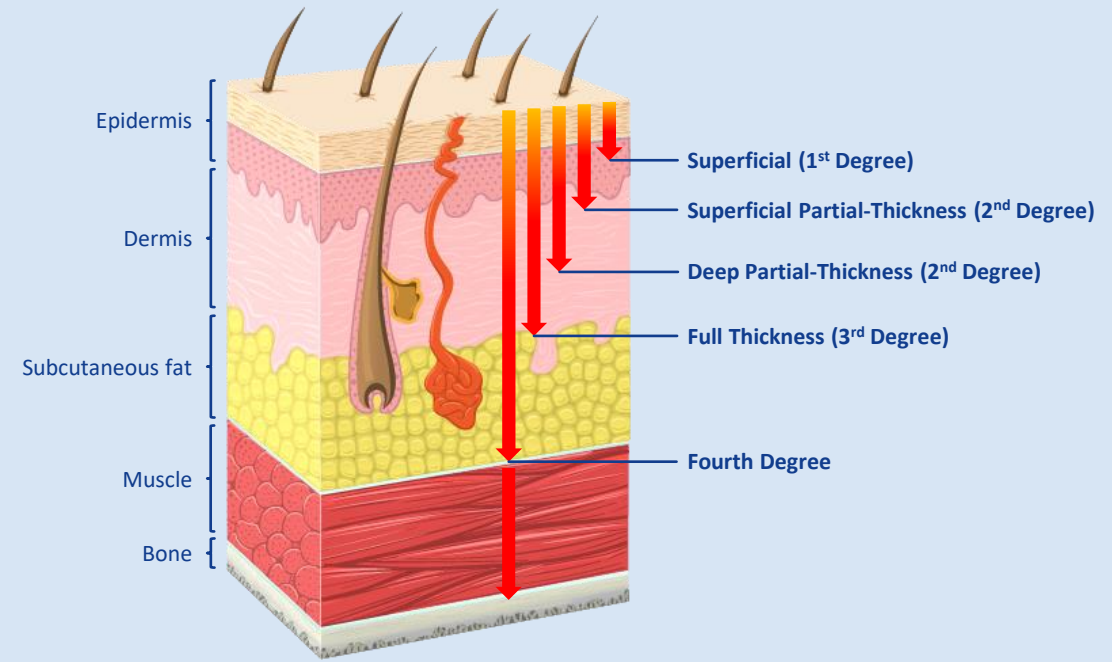
Potential MACI Ankle Indication Would Increase MACI Total Addressable Market to \$4 Billion



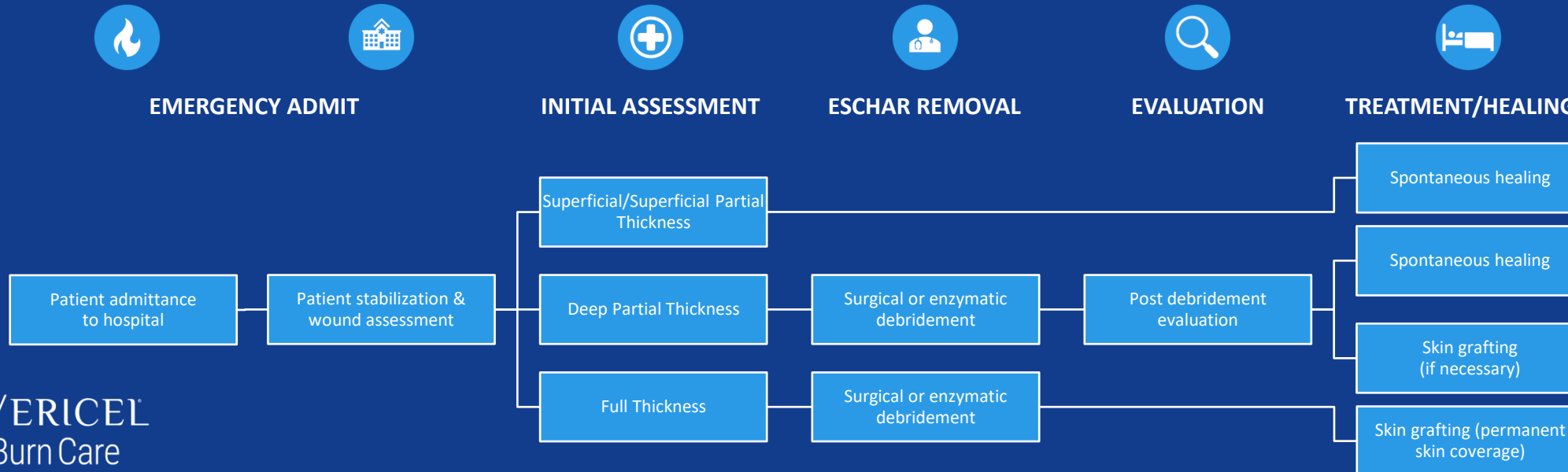
¹ Health Advances LLC MACI market assessment report (2018), Vericel data, LexisNexis, Medtech Insight, NY SASD, SmartTRAK, LSI, PSPS, McCormick, Frank et al. Arthroscopy, (2014) 30(2): 222-6, Montgomery, et al. Knee Surg Sports Traumatol Arthrosc (2014) 22: 2070. ² Health Advances LLC MACI market assessment report (2018) ³ Assumes MACI ASP of \$50,000+. ⁴ SmartTrak Cartilage Repair Procedures; resurfacing includes microfracture, OATs, OCA, etc. and does not include chondroplasty/debridement only. ⁵ Cello Health MACI Ankle quantitative market research survey (2021).

Burn Injury Size and Depth Determine Treatment Pathway

- ❖ Full thickness burn injuries of any size & partial thickness burn injuries >10% total body surface area (TBSA) are most often transferred to specialized burn centers
- ❖ Full thickness & deep partial-thickness burns **require eschar removal and grafting** to achieve wound closure

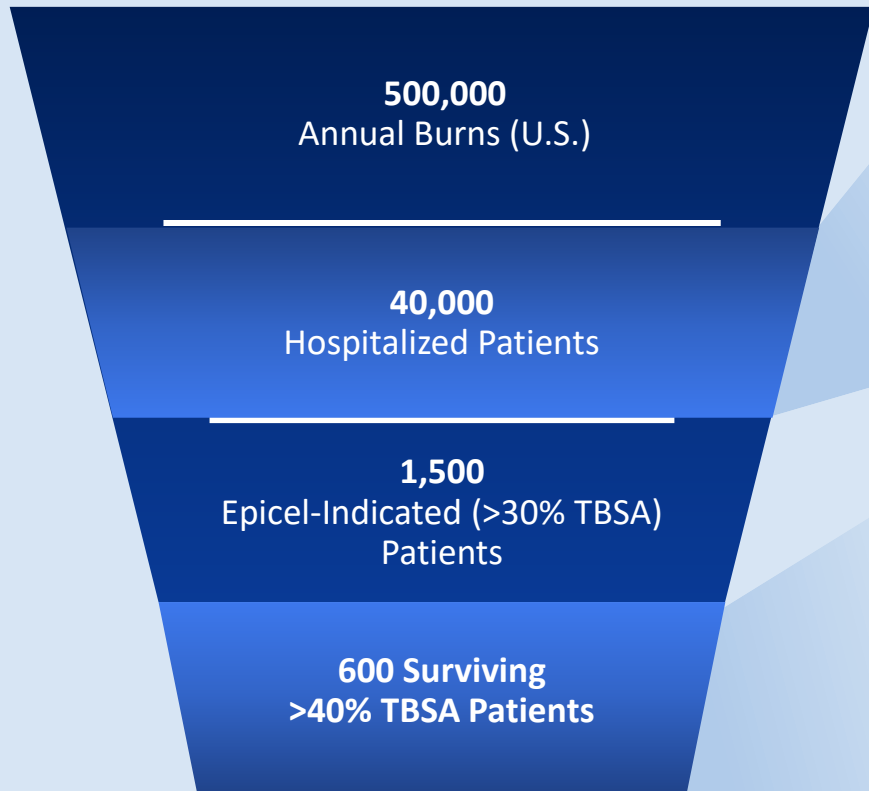


TREATMENT PATHWAY



Burn Care Franchise Addressable Market Opportunity

Estimated U.S. Burn Patients¹



NexoBrid commercialization significantly expands the total addressable market and establishes second high growth franchise for Vericel

¹ 2017 National Burn Repository Report Version 13.

² ~90% of hospitalized patients with thermal burns; ~90% of eligible patients require eschar removal (management estimate).

³ Assumes NexoBrid average price of ~\$9,000 per patient.

⁴ Assumes 600 patients x 120 grafts per patient x ~\$4,000+ per graft.

NexoBrid

Indications and Usage:

Contains proteolytic enzymes and is indicated for eschar removal in adults with deep partial-thickness and/or full-thickness thermal burns

NexoBrid can be applied to up to 20% body surface area in two applications



Significant Advancement in Burn Treatment Paradigm

- ❖ Concentrated mixture of proteolytic enzymes derived from the stem of the pineapple plant (*Ananas comosus*)
- ❖ Non-surgical topical agent that may be applied at the patient's bedside
- ❖ Selectively degrades eschar in four hours while preserving viable tissue



¹ NexoBrid Label. Cambridge, MA. Vericel Corporation; 2022.

² Krieger Y, Bogdanov-Berezovsky A, Gurfinkel R, et al. Efficacy of enzymatic debridement of deeply burned hands. Burns. 2012;38:108-112.

³ Palao R, Aguilera-Saez J, Collado JM, et al. Use of a selective enzymatic debridement agent (NexoBrid) for wound management: Learning curve. World J Dermatol. 2017;6(2):32-41.

NexoBrid Treatment Application

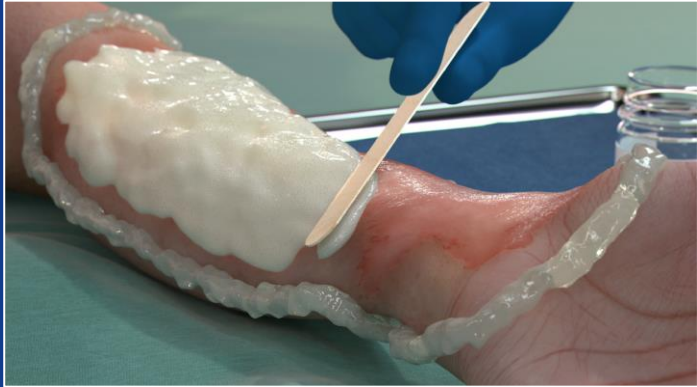
Clean Wound



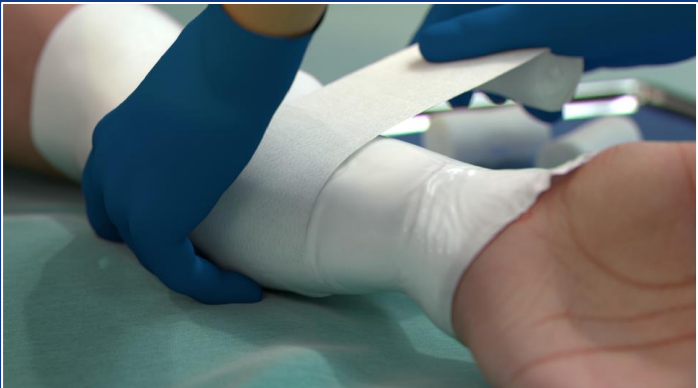
Antibacterial Pre-Soak



NexoBrid Application



Film Dressing (4 Hours)



Remove Eschar



NexoBrid Launch Progress

- ❖ NexoBrid launched in the U.S. in Q4 2023
- ❖ Key Performance Indicators*
 - **50+** Burn Centers have submitted packages to their P&T Committees
 - **25+** Burn Centers have P&T Committee approval
 - **~20** Burn Centers have placed initial orders

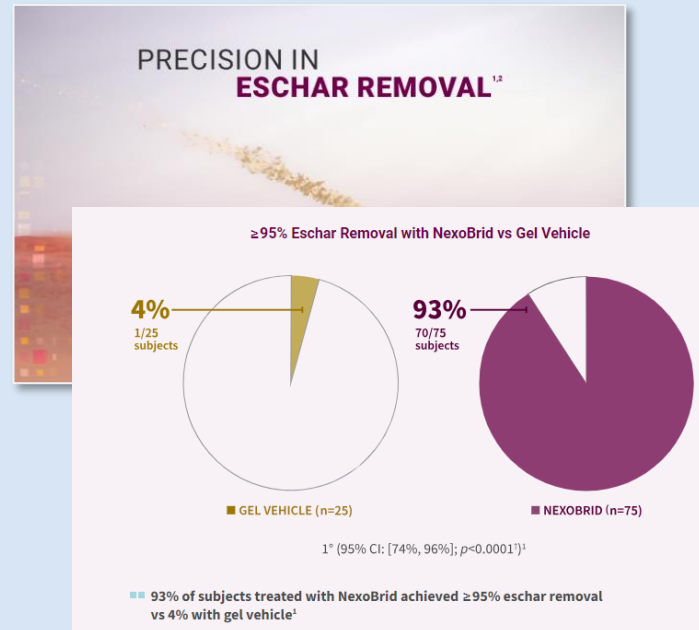


* As of December 31, 2023.

Strong Interest in NexoBrid by Treating Physicians and Burn Centers

NEXOBRID IS NOW COMMERCIALY AVAILABLE IN THE U.S.

Robust Clinical Efficacy



Application Demonstrations



Multi-Disciplinary Education & Clinical Application Training

NOW APPROVED!

NexoBrid
(anacaulase-bcdb)

You are invited to an educational program during the American Burn Association Annual Meeting 2023 at the Gaylord Texan Resort & Convention Center in Dallas, TX

Enzymatic Eschar Removal Practicum: Education and Clinical Application Training

Wednesday May 17, 2023
7:00-8:30pm
Texas Ballroom C

PRESENTED BY:

- Steven Kahn, MD**
Associate Professor
Chief of Burn Surgery
The South Carolina Burn Center
Medical University of South Carolina
- James Boron, MD**
Senior Medical Director,
Vericel Burn Care

REGISTER NOW

This program is sponsored by Vericel Corporation and is not eligible for CE credits

Topics presented will include:

- Presentation of NexoBrid® (anacaulase-bcdb) product background, clinical trial data, and application
- Case presentations and product use techniques
- NexoBrid® (anacaulase-bcdb) lab training demonstration

LAB DEMONSTRATION INSTRUCTORS:

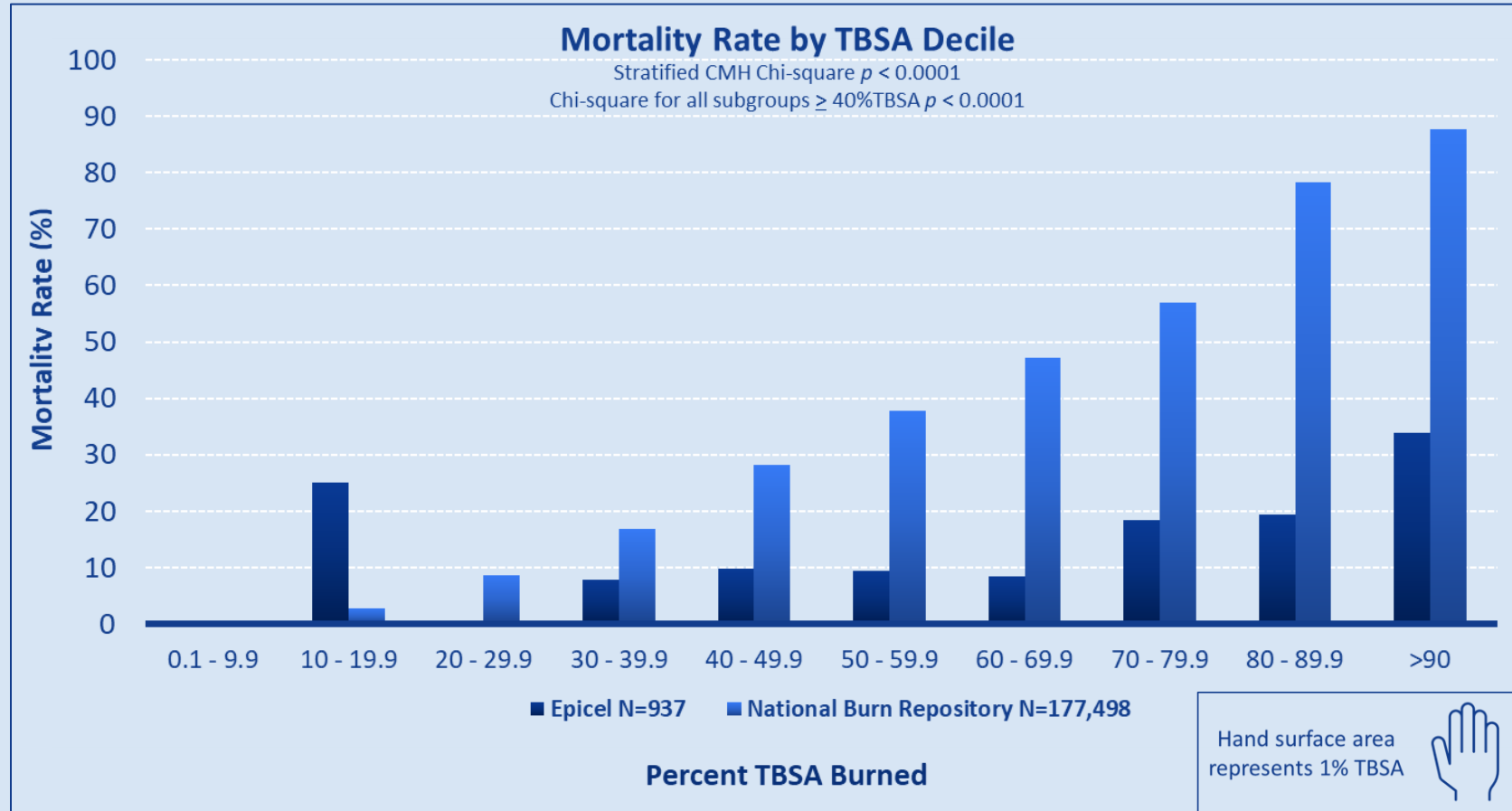
- Dr. Nicole Bernard, Ohio State, Wexner Medical Center
- Dr. Sigrid A. Howell-Baerem, Lehigh Valley Health Network
- Dr. Jeremy Goresman, Massachusetts General Hospital
- Dr. Tracee Shott, Baton Rouge General
- Dr. Adam Singer, Stony Brook University
- Dr. Lucy Wilkenberry, University of Iowa Health Care

Epichel

- ❖ Only FDA-approved permanent skin replacement for adult and pediatric patients with full-thickness burns $\geq 30\%$ of total body surface area
- ❖ Important treatment option for severe burn patients where little skin is available for autografts



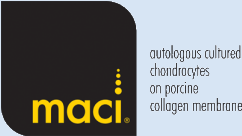
Comparison of Epichel Patient Database to National Burn Repository¹ Data Demonstrates Lower Mortality Rate



Vericel Remains Focused on Potential Strategic Transactions to Maximize Long-Term Value

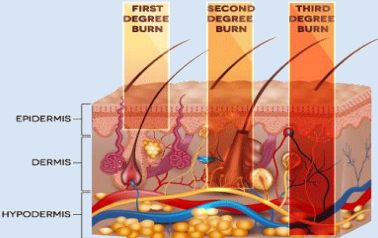
ADVANCED CELL THERAPY DEVELOPMENT & MANUFACTURING PLATFORM

Sports Medicine Franchise



autologous cultured chondrocytes on porcine collagen membrane

Severe Burn Care Franchise



Epicel®
(cultured epidermal autografts)

NexoBrid®
(anacaulase-bcldb)

New Advanced Cell Therapy Vertical(s)



Business development activities focused on opportunities having a strategic fit with current **franchises** or advanced cell therapy **platform**

Growth Strategy Leverages Near-Term & Long-Term Opportunities



Strong Financial Profile

- ❖ High revenue growth profile
- ❖ Sustained positive adjusted EBITDA and Operating Cash Flow
- ❖ ~\$152.6 million in cash and investments as of 12/31/2023



High-Growth Sports Medicine Franchise

- ❖ Market leader in knee cartilage repair
- ❖ 20%+ total revenue CAGR since 2017
- ❖ Focused on maximizing key growth drivers



Advancing Pipeline

- ❖ MACI Arthro submission accepted for review
- ❖ MACI Ankle program advancing
- ❖ NexoBrid sBLA for pediatric indication accepted for review



Second High-Growth Franchise in Burn Care

- ❖ NexoBrid launched in Q4 2023
- ❖ High surgeon interest to date