



Innovations in Vascular Tissue Engineering- A Promising New Potential for Patients

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CONFLICTS OF INTEREST & DISCLOSURES

Dr. Blum is Co-Founder & EVP, Corporate Development and a shareholder in Humacyte, Inc.

None of the data presented in this lecture is intended to be construed as claims for the clinical use, efficacy, or safety of any medical product. The human accellular vessel (HAV) is an investigational product that has not been approved by FDA for any indication.

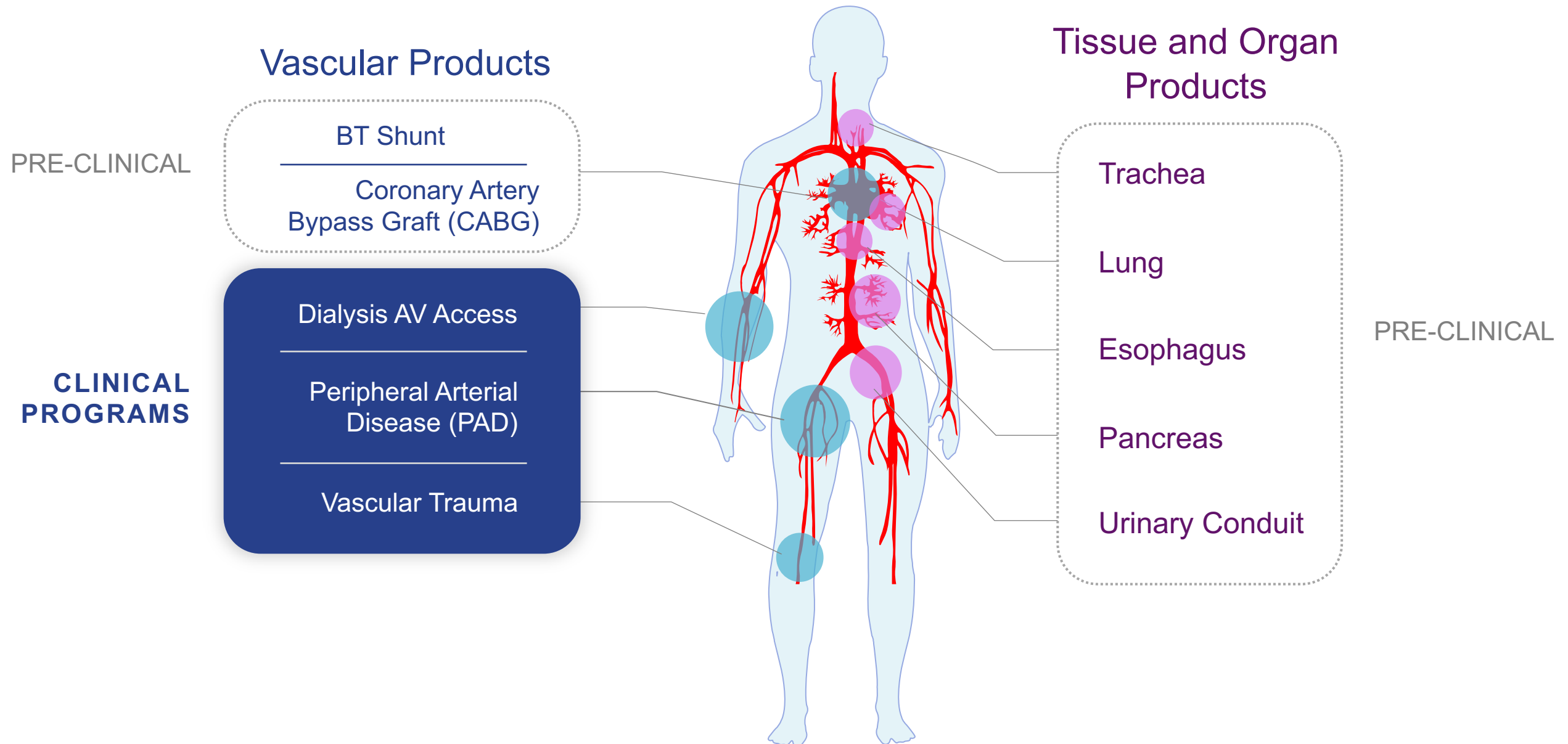
These slides and the accompanying oral presentation contain forward-looking statements. All statements, other than statements of historical fact, included in these slides and the accompanying oral presentation are forward-looking statements reflecting management's current beliefs and expectations. In some cases, you can identify forward-looking statements by terminology such as "will," "anticipate," "expect," "believe," "intend" and "should" or the negative of these terms or other comparable terminology. Forward-looking statements in these slides and the accompanying oral presentation include, but are not limited to, statements about the initiation, timing, progress and results of our clinical trials; the anticipated characteristics and performance of our human acellular vessels (HAVs), our ability to successfully complete, clinical trials for our HAVs; the anticipated benefits of our HAVs relative to existing alternatives; the commercialization of our HAVs and our ability to manufacture at commercial scale; the implementation of our business model, strategic plans for our business; the scope of protection we are able to establish and maintain for intellectual property rights covering our HAVs and related technology; estimates of our expenses, health economics, future revenues, capital requirements and our needs for additional financing; the timing or likelihood of regulatory filings and approvals; timing, scope and rate of reimbursement for our HAVs; our estimated available market opportunity; our ability to maintain and establish collaborations; our financial performance; developments relating to our competitors and our industry; statements regarding our markets, including the estimated size and anticipated growth in those markets; and statements related to our proposed business combination with a subsidiary of Alpha Healthcare Acquisition Corp. (AHAC), including the timing and structure of the transaction and our ability to recognize the anticipated benefits of the business combination. These statements relate to future events or to our future financial performance and involve known and unknown risks, uncertainties and other 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 these forward-looking statements. Except as required by law, we assume no obligation to update these forward-looking statements, even if new information becomes available in the future.

COMPANY & TECHNOLOGY OVERVIEW

Humacyte is pioneering the development and manufacture of off-the-shelf, universally implantable, bioengineered human tissues



HUMACYTE DEVELOPS BIOENGINEERED TISSUES TO TREAT DISEASES THROUGHOUT THE BODY



HISTORY DEFINING MILESTONES



2004

Founded by Drs. Laura Niklason, Juliana Blum, and Shannon Dahl, Durham, NC



2013

First human surgical implantation of HAV at Duke University



2014

Humacyte's HAV received Fast Track Designation for Vascular Access in Hemodialysis Program



2017

Humacyte Receives Regenerative Medicine Advanced Therapy Expedited Review Designation for HAV



2018

FMC & Humacyte Announce Global Partnership Supported by \$150M Equity Investment



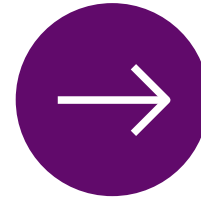
2019

Manufacturing facility validation qualification, Durham NC



2021

Humacyte goes public on Nasdaq as \$HUMA



FUTURE

Product Launch & Pipeline Development

MANUFACTURING BIOENGINEERED TISSUES

Modular Manufacturing System

Bioreactor bag



Each bioreactor bag contains a single polymer mesh scaffold, seeded with donated human cells



Growth drawer



10 bioreactor bags per growth drawer; tubing connects to shared nutritive media



LUNA200 system



With 20 growth drawers, each LUNA200 can produce 200 HAVs per batch (or ~1,000 HAVs annually) ¹

Commercial 83,000 sq ft Bioprocessing Facility



- Currently operating 8 LUNA200 systems
- Annual Capacity expected to exceed 40,000 HAVs
- Functionally closed system with state-of-the-art process automation

READILY AVAILABLE HUMAN ACELLULAR VESSELS (HAVs)

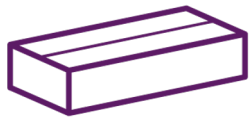


HAVs are shipped to hospitals for use in operating rooms.
During surgery, the HAV is removed from its packaging and then implanted into the patient.

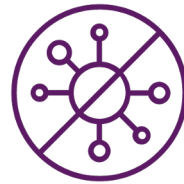
HUMAN ACELLULAR VESSELS (HAVs)- PLACEHOLDER FOR DEMO VIDEO

KEY FEATURES OF HUMACYTE TECHNOLOGY

Potential benefits of HAVs evaluated in completed and ongoing clinical trials across multiple indications



Off-the-shelf,
immediately available
with 18-month shelf life



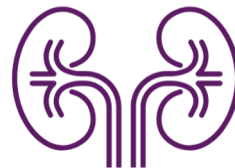
HAV is highly
resistant to infection



No evidence of
immunogenicity



Host cells repopulate
the HAV

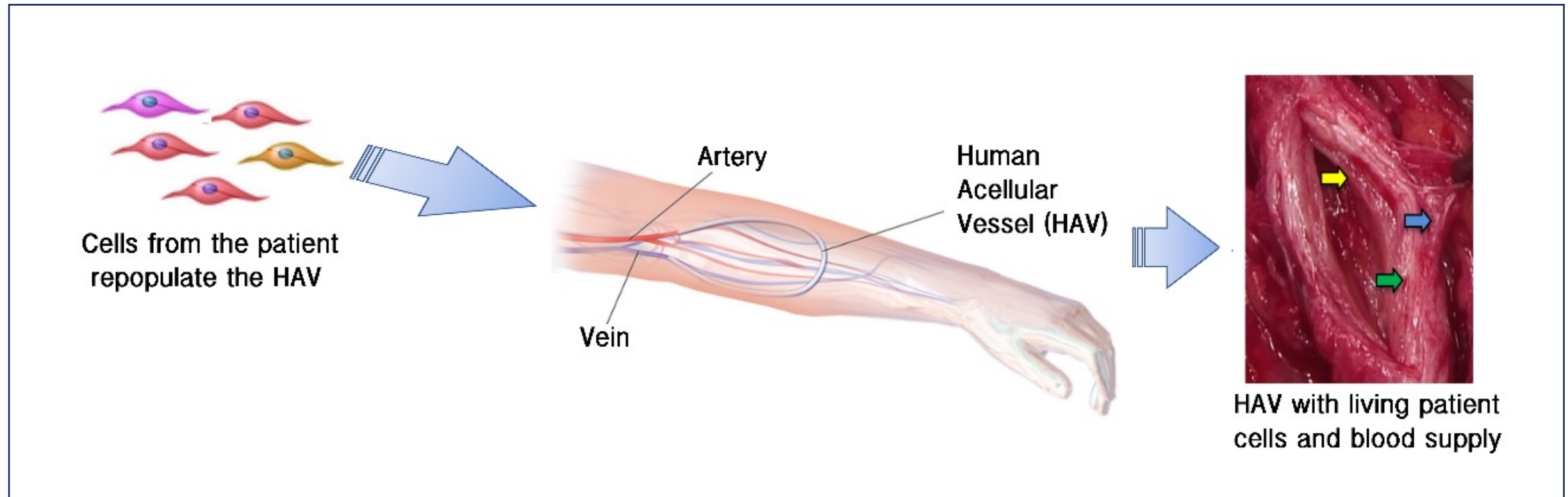


For ESRD patients,
accessible for dialysis
access within one
month of implantation



Long-term durability
is demonstrated in
ongoing studies

CLINICAL EVIDENCE THE HAV REGENERATES WITH PATIENT CELLS

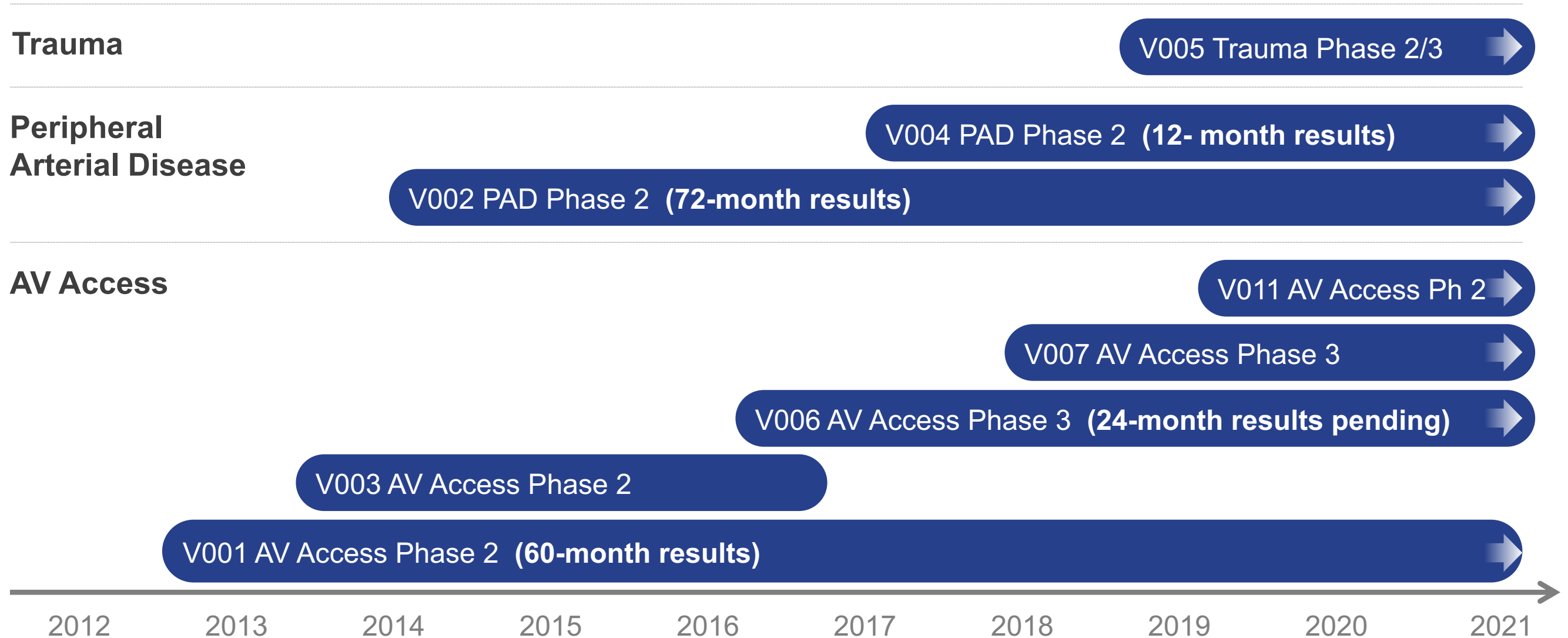


After implantation, clinical data has shown cells from the patient gradually repopulate the HAV, producing a tissue that has living cells and its own blood supply.

CLINICAL OVERVIEW - DIALYSIS



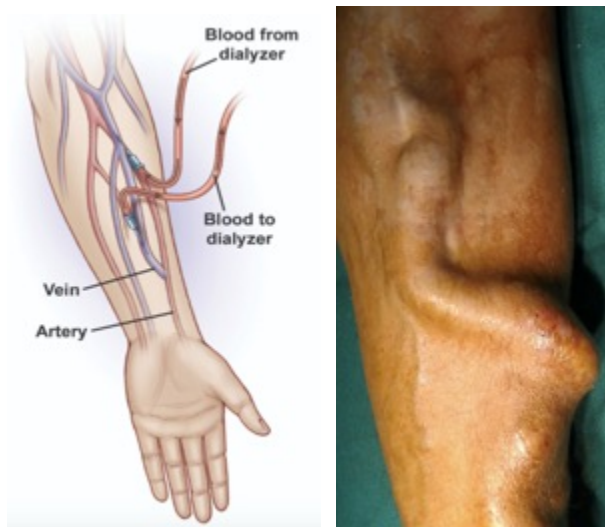
EXTENSIVE CLINICAL EXPERIENCE ACROSS MULTIPLE CLINICAL TRIALS



HAVs have been implanted into hundreds of patients over more than 8 years.

THE HAV in DIALYSIS: ADDRESSING RECURRENT INFECTIONS AND FISTULA FAILURE

AV Fistula



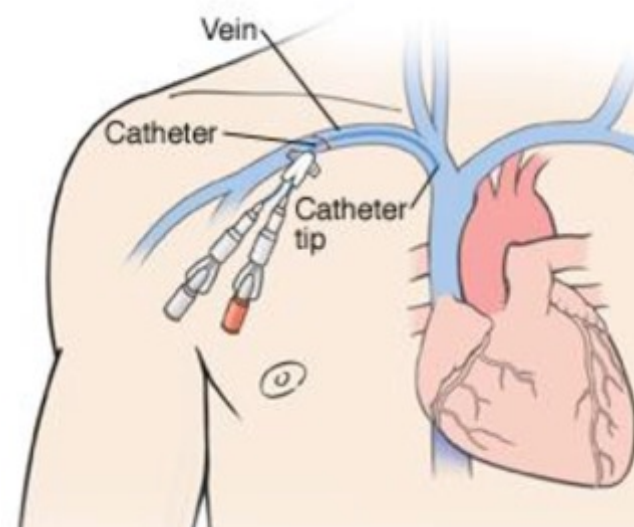
Standard of Care

- Major risks associated with catheter during wait for fistula maturation
- ~40% of fistulas fail

Humacyte HAV

- HAV usable within 1 month vs 3-6 months for fistulas
- Decreased catheter contact time in patients awaiting fistula maturation

Catheter



- High blood stream infection rates (up to 200% per patient-year)

Infection rate for:

- Catheters: up to 200% per patient year¹
- HAV: 1% per patient year²

Synthetic Graft



- 10-15% annual infection rate: sepsis, hospitalization, death
- Not durable: ~50% fail in 2 years¹

- 10-15x lower rate of infection versus ePTFE
- Excellent Durability: used for dialysis for ~7 years

¹Lawson, J.H., et al, The Lancet 2016; 387: 2026-2034.

²Halbert, R.J., et al, Kidney360 2020; doi: 10.34067/KID.003502020.

HAV IN HEMODIALYSIS ACCESS: PHASE 2 STUDY THROUGH 12 MONTHS

- **Methods:** Six centers in the US and Poland, HAV implanted in patients who were in need of dialysis access and who were suitable for arteriovenous grafting ¹.
- **Subjects:** 60 patients, mean follow-up 16 months
 - Age = 59 ± 10y;
 - 77% Caucasian;
 - 90% with hypertension;
 - 43% diabetic;
 - Prior AV accesses: 3.6 ± 2.1.
- **Safety Outcomes:**
 - No aneurysmal degeneration;
 - No clinical rejection;
 - Multiple subjects subsequently received successful kidney transplants.
- **Results:**
 - 12 month HAV outcomes published in *The Lancet* ¹

Volume 387, No. 10032, p2026-2034
Published in issue: May 14, 2016

Phase 2 HAV Results vs. Historical Fistula & ePTFE Data

Conduit	6-month Secondary Patency	12-month Secondary Patency	Infection Rate per patient-year
HAV Phase 2	97% (85-98%)	89% (74-93%)	1.3%
Historical Fistula ^{2,3,4}	61% ³ (useable for dialysis)	59.5% ⁴	4.0% ⁵
Historical ePTFE ⁵	80% (75-84%)	70%(64-75%)	9.0%

¹ Lawson, J.H. et al. The Lancet 2016; 387: 2026-2034.

² Halbert, R.J. et al. Kidney360 December 2020, 1 : 1437-1446

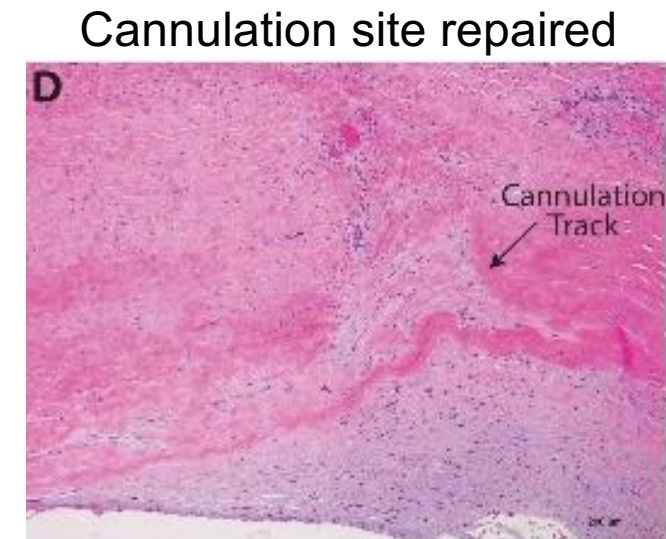
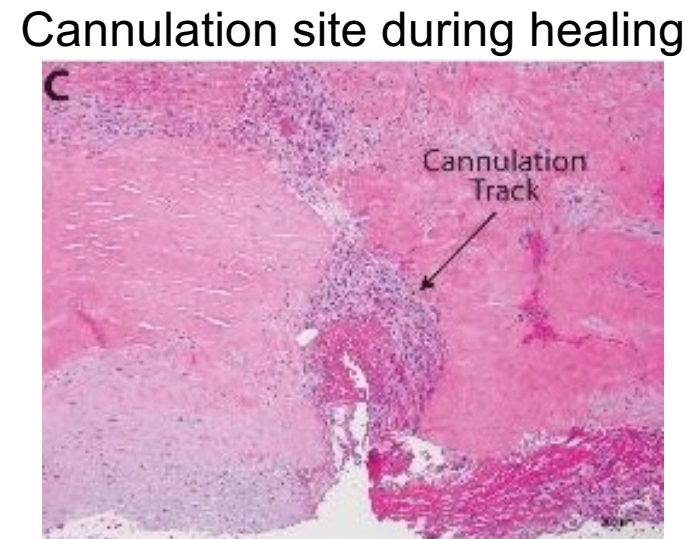
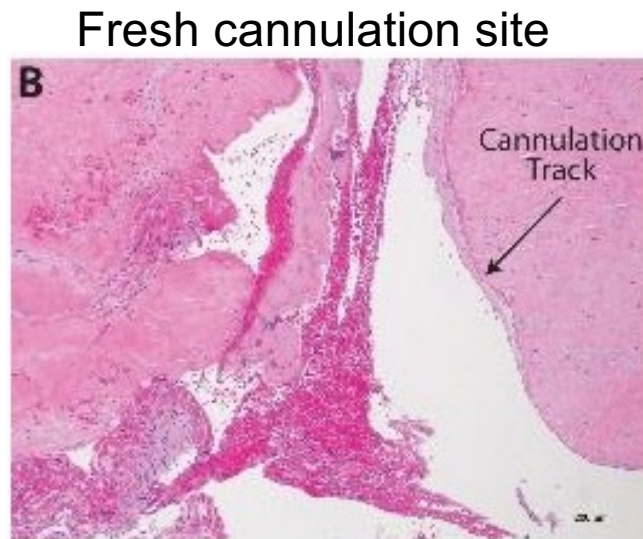
³ Allon, M., et al. American J Kidney Disease 2018; 71: 677-689.

⁴ Arhuidese, I.J., et al. Journal Vascular Surgery 2018; 68: 1166-1174

⁵ Al-Jaishi, A.A., et al. JASN 2017; 28: 1839-1850.

CLINICAL EVIDENCE OF HEALING

Low
magnification of
3 cannulation
sites



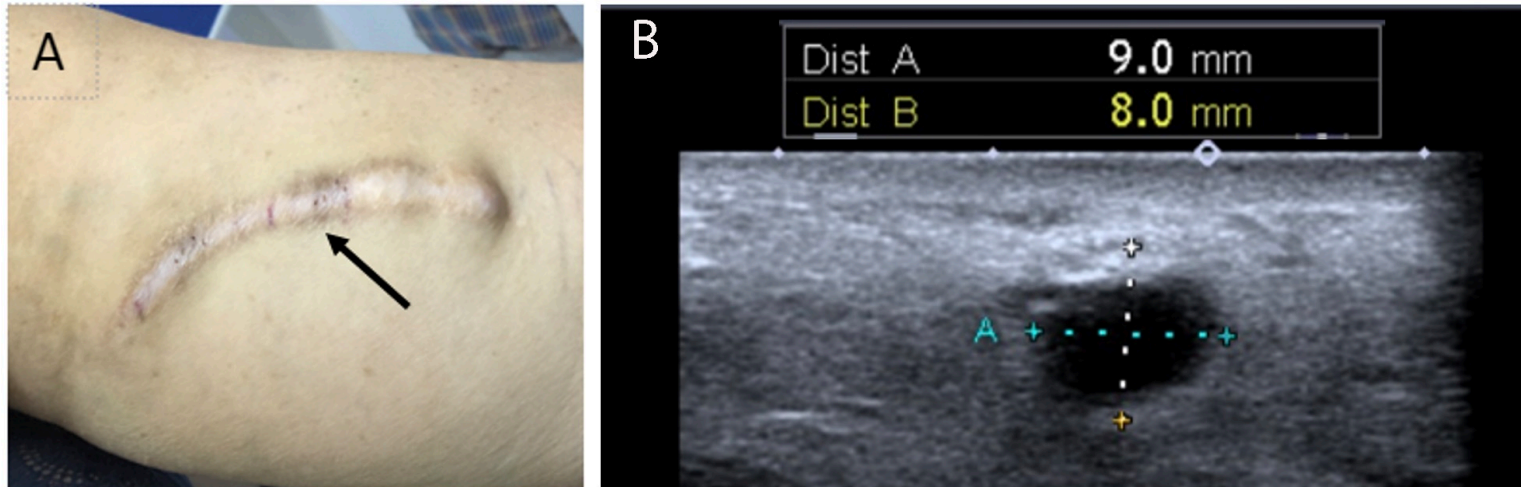
Repopulation with host vascular cells and angiogenesis enable healing



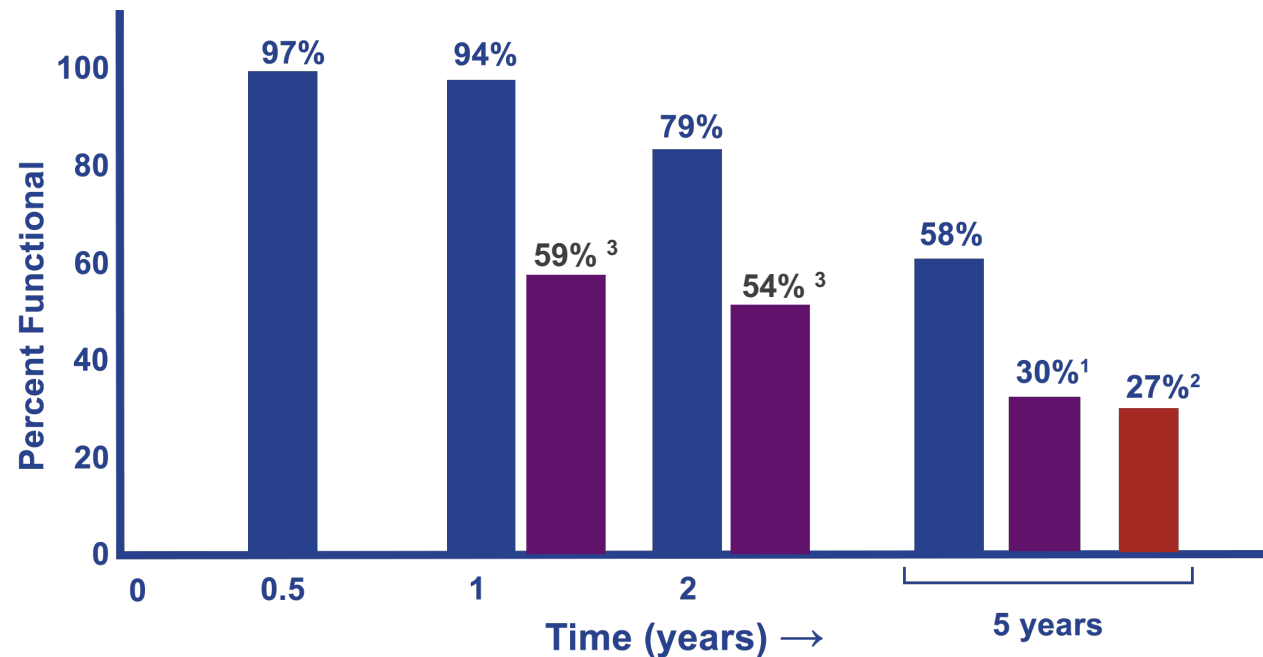
**Clinical data¹ suggests after cannulation, HAV potentially heals to close the cannulation injury site.
In contrast, PTFE has permanent cannulation injury with no healing.**

¹ HUMACYL® Patient CLN-PRO-V003 01-001 histology Phase II trial, 2014. unpublished data. A section of an implanted Humacyte graft removed at 11 months. All images are Hematoxylin & Eosin stain (H&E) (n=1)

HAV IN HEMODIALYSIS ACCESS: PHASE 2 STUDY ≥ 5 YEARS, LONG TERM DURABILITY



- A) Patient with access site utilized for 6 years (arrow).
- B) Ultrasound of HAV from same patient.



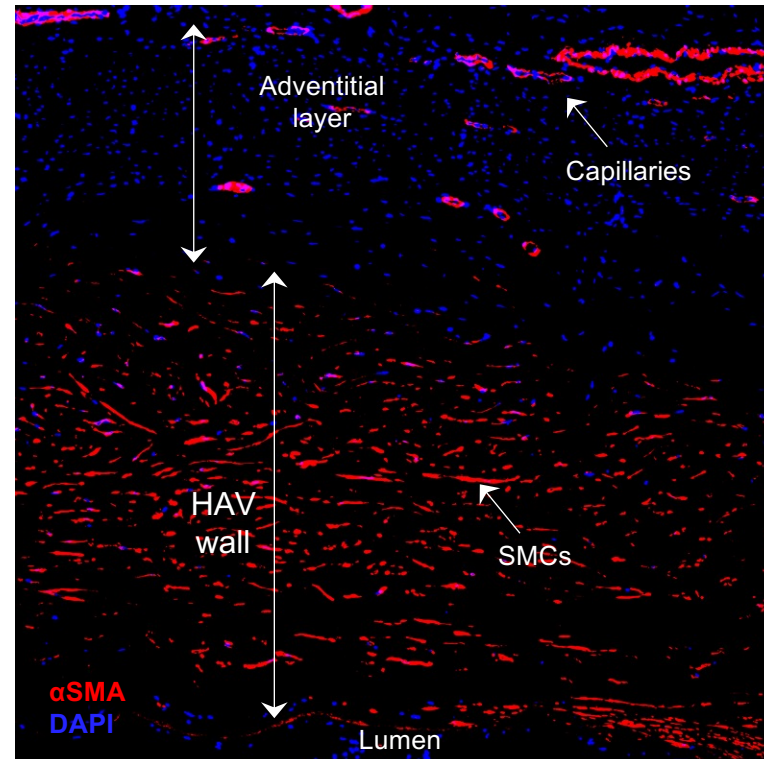
■ HAV ■ Fistula ■ ePTFE

58% secondary patency at 5 years compares well to historical ePTFE and arteriovenous fistulas.

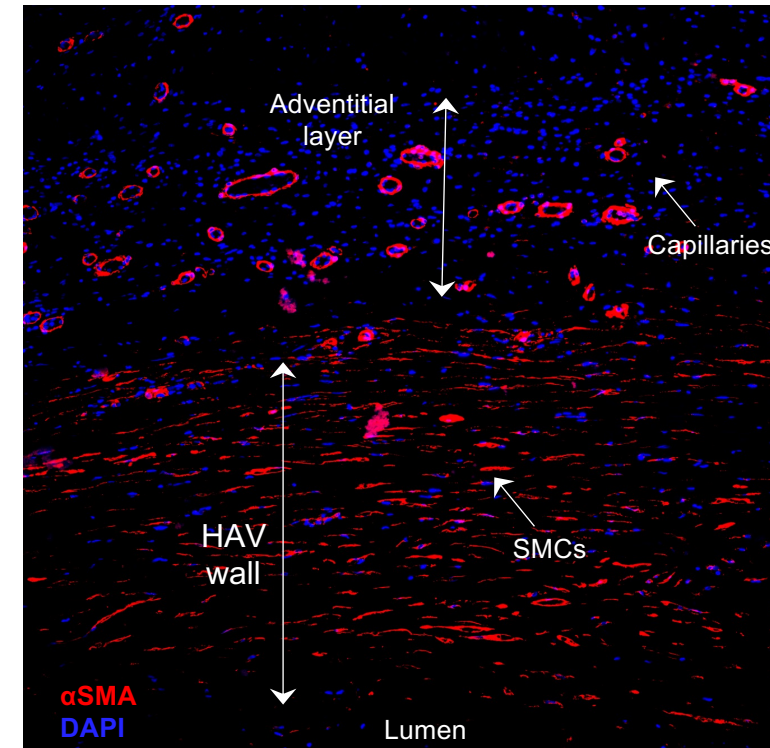
¹ Lok, et al; 2013 CJASN
² Kakisis et al; 2017, JVS
³ Arhuidese, et al, 2018; JVS.

CLINICAL EVIDENCE SHOWS REMODELING OF THE HAV IS CONSISTENT AND ANGIOGENIC

Subject 3079-012-V006, male, 26 years old.
67 weeks post-implantation of HAV



Subject 1006-001-V006, male, 83 years old.
66 weeks post-implantation of HAV



**Patient cells remodel the HAV across a wide range of patient ages.
Repopulation with vascular cells is combined with robust peri-HAV angiogenesis.**

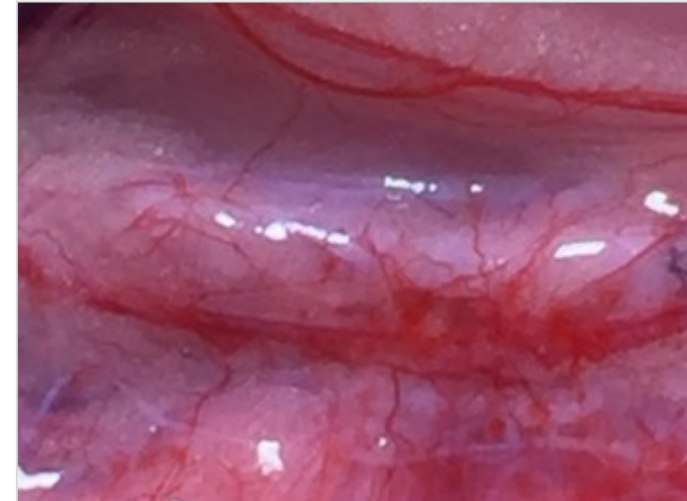
THE ERA OF READILY AVAILABLE, ENGINEERED HUMAN TISSUES HAS ARRIVED

Bioengineered Blood Vessel

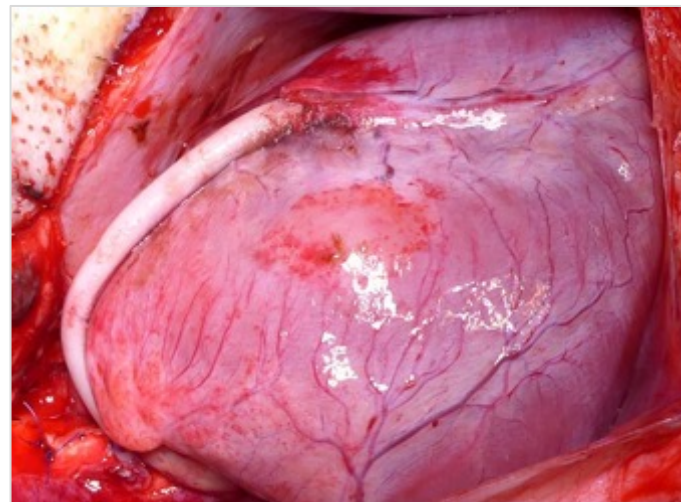
- Hemodialysis
- Vascular Trauma
- PAD



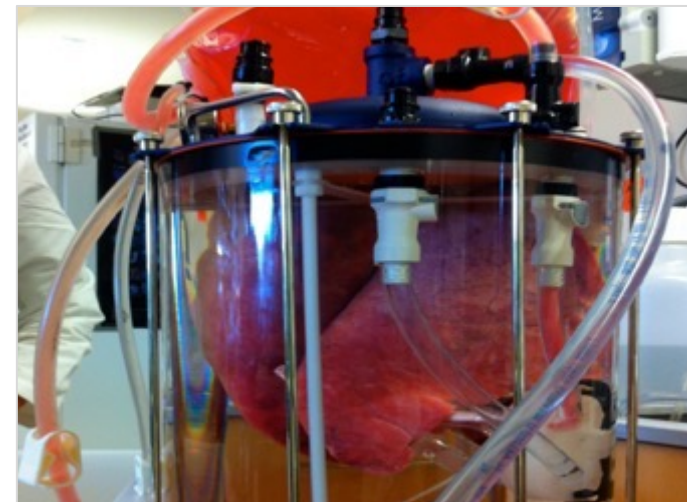
Bioengineered Pancreas



Bioengineered Human Coronary Artery



Bioengineered Human Lung



SUMMARY

- Breakthrough innovations in patient care- advancing the future of therapeutic opportunities today
- Committed to developing product opportunities across multiple therapeutic areas- addressing significant unmet medical needs
- Focused on patient need- bringing new opportunities to those seeking better options for their care
- Integrating the patient voice and perspective is critical to our mission





THANK YOU

We are committed to bringing first-in-class regenerative medicine products to the marketplace that will improve and save the lives of patients worldwide and transform the practice of medicine.





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