

3-D Hollow Fiber Bioreactors: “A better way to grow cells”

By John J. S. Cadwell

MOVE-2025, Tartu, Estonia October 7-10

“2-D, 3-D and Histocentric
Bioreactors, Recapitulating the
in vivo Microenvironment”



About FiberCell Systems Inc.

- Founded in 2000 by John J.S. Cadwell, President & CEO
- Over **25 years** advancing hollow fiber bioreactor technology
 - Supported by **500+ peer-reviewed publications** from scientists worldwide
- Trusted leader in laboratory-scale hollow fiber bioreactors

Presentation Outline

- The Evolution of Cell Culture
- Concepts of Histocentric Hollow Fiber Bioreactors (HFBR)
- 2D Vs. 3D Vs. 4D Cell Culture
- How Hollow Fiber Bioreactors Work
- Key Applications
- Case Studies & Results (3D & EV)
- Summary & Conclusion

The Evolution of Cell Culture

- * 1665 Hooke first observes “cells”
- * 1876 Charles Chamberland invents autoclave (Pasteur)
- * 1907 Harrison frog neuronal explants on plasma clots
- * 1935 Carrel and Lindberg develop Pyrex for organ perfusion pump
- * 1951 HELA Cells (roller tube culture)
- * Late 50’s Tissue dissociation enzymes from Worthington
- * 1954 Connaught Labs makes polio vaccine in 5 liter Pyrex flasks
- * 1958 Theodore Puck first uses fetal bovine serum
- * Mid 1970’s Plastic is now more common than glass for cell culture
- * Early 1970’s Laminar flow hoods
- * 1978 Milstein Hybridoma Nobel Prize

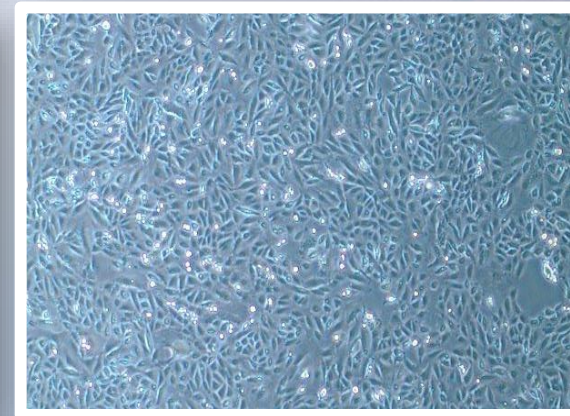
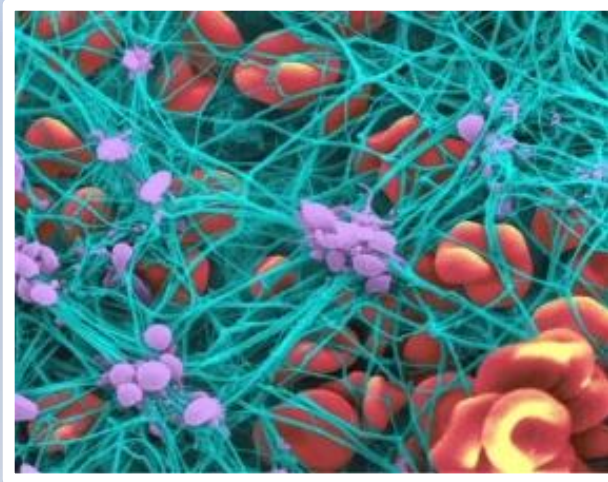
Science October 1972 Hollow Fiber Bioreactors Richard Knazek

The Evolution of Cell Culture:

1935 Carrel and Lindberg develop Pyrex for organ perfusion pump



The Evolution of Cell Culture: Cell Culture Through the Ages



2-D Vs. 3-D Cell Culture

2-D Cell Culture:

- **Environment:** Cells are grown on flat surfaces (e.g., Petri dishes), forming a monolayer.
- **Cell Interactions:** Limited cell-cell and cell-matrix interactions, which can lead to altered cell behavior.
- **Morphology:** Cells appear flat and stretched, often not reflecting their natural shape.
- **Gene Expression:** Often differs from *in vivo* conditions, leading to less accurate biological responses.

3-D Cell Culture:

- **Environment:** Cells grow in a three-dimensional space, allowing for more natural interactions.
- **Cell Interactions:** Enhanced cell-cell and cell-matrix interactions, mimicking *in vivo* conditions more closely.
- **Morphology:** Cells can form spheroids or organoids, resembling natural tissue structures.
- **Gene Expression:** More similar to *in vivo* models, providing better insights into cellular functions and drug responses.

Types of 3-D Cultures

- Spheroids (one cell type, no scaffold)
- Organoids (multiple cell types, self-organize, scaffold)
- Bio-Printed Tissues
- Hydrogels
- Organ on a chip
- Hanging drop
- Transwell dishes
- Hollow fiber bioreactors

Cell Growing as a monolayer in 2D Culture

Not very physiologic!



Scale-up Options in 2-D Cell Culture

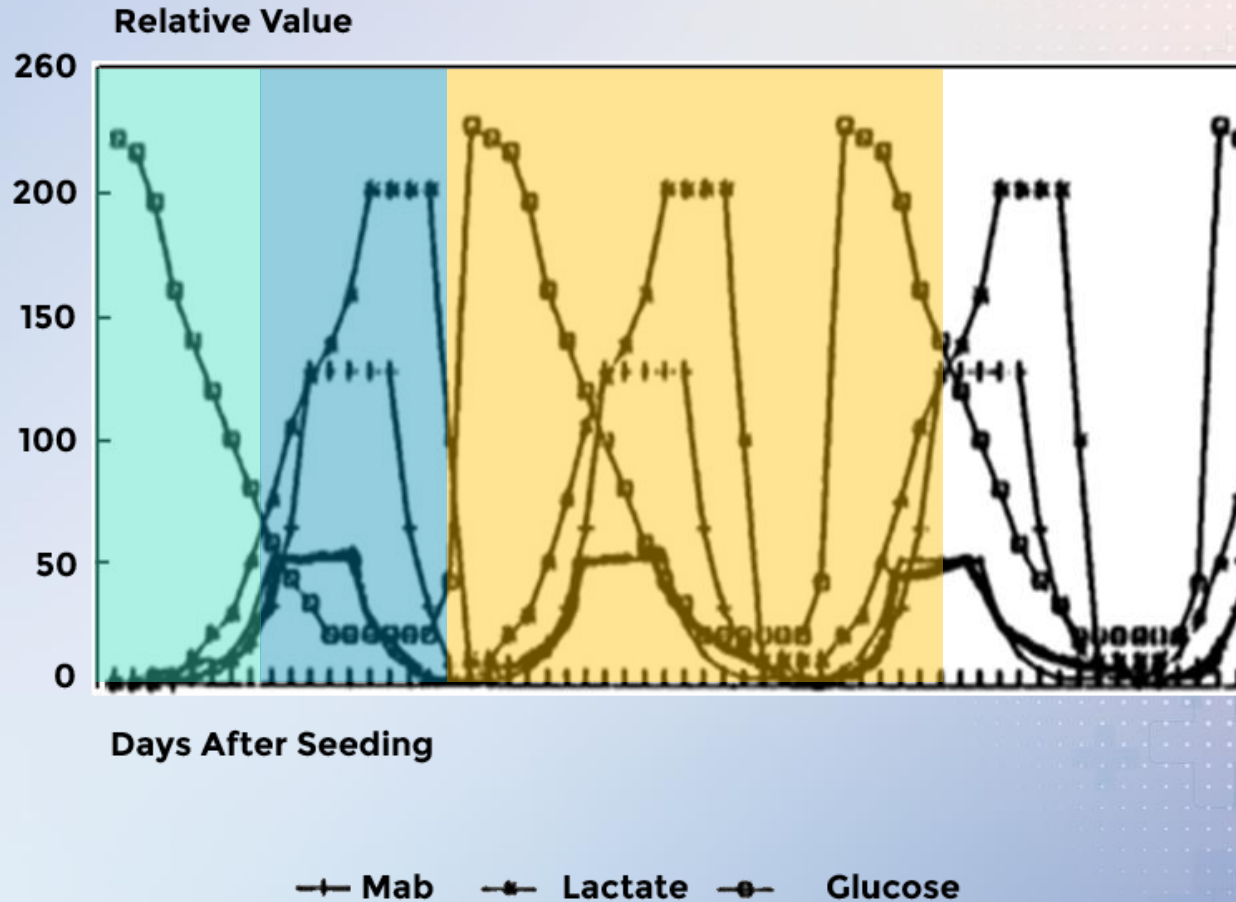
- Roller Bottles
- Cell Factory
- Cell Cube
- Cell Culture Bags
- Spinner Flasks
- Bioreactors
- Microcarriers

“Check the patents!”

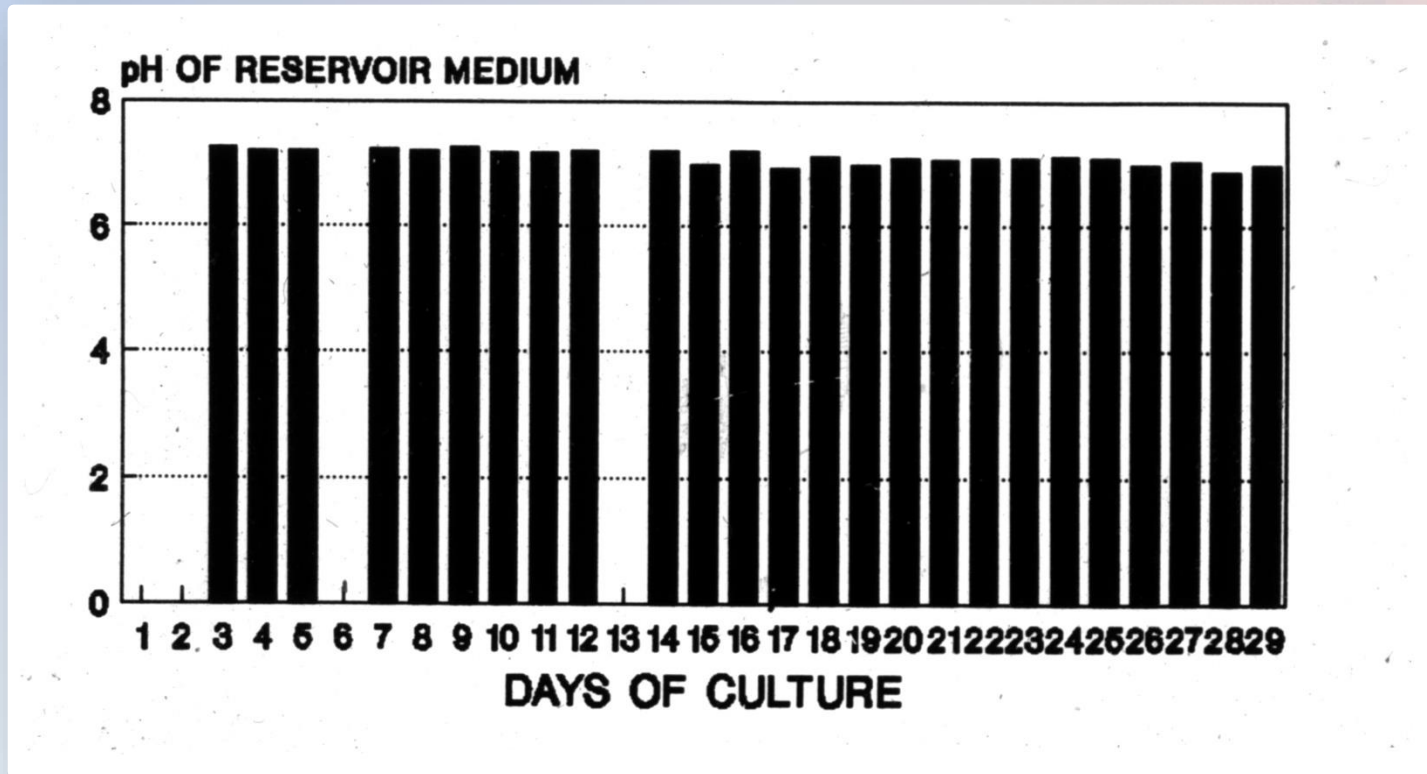
How About Scale-up in 3-D Culture?



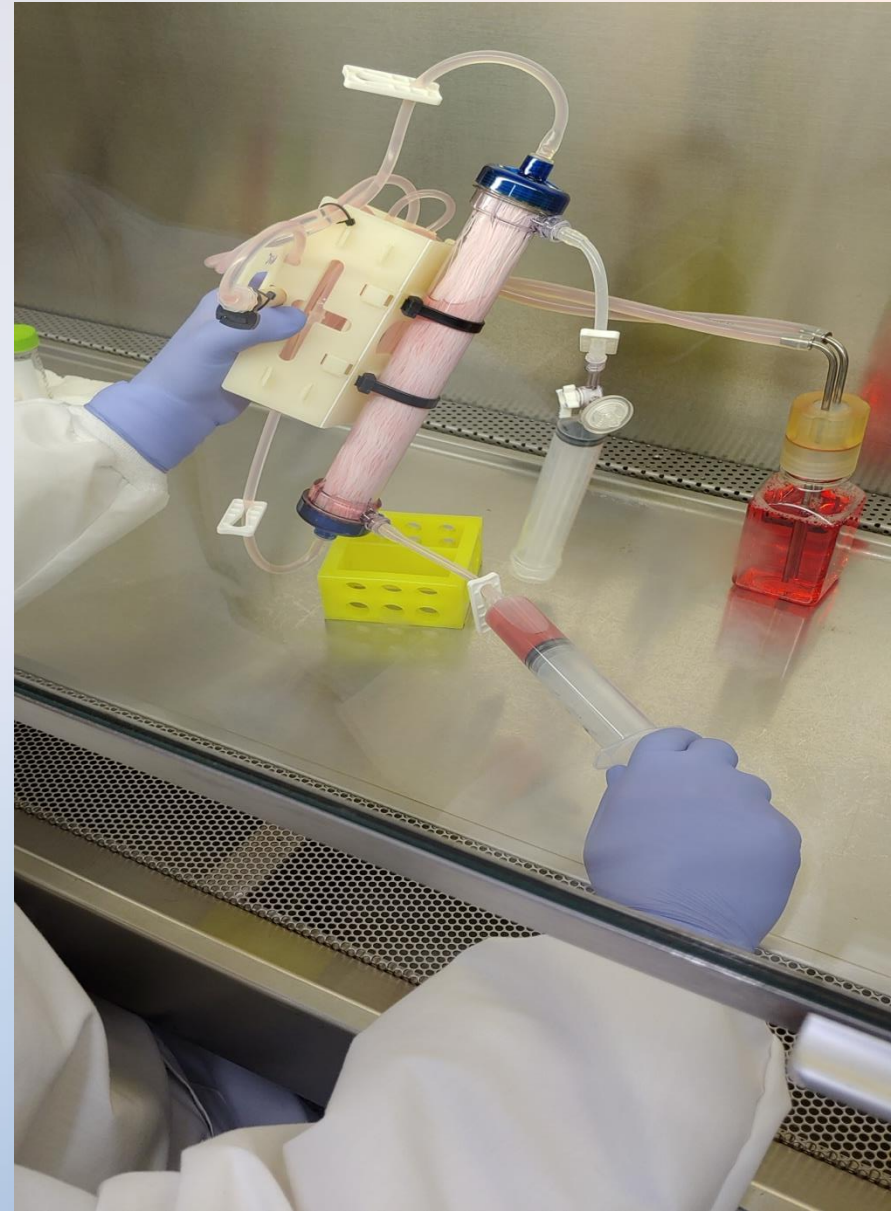
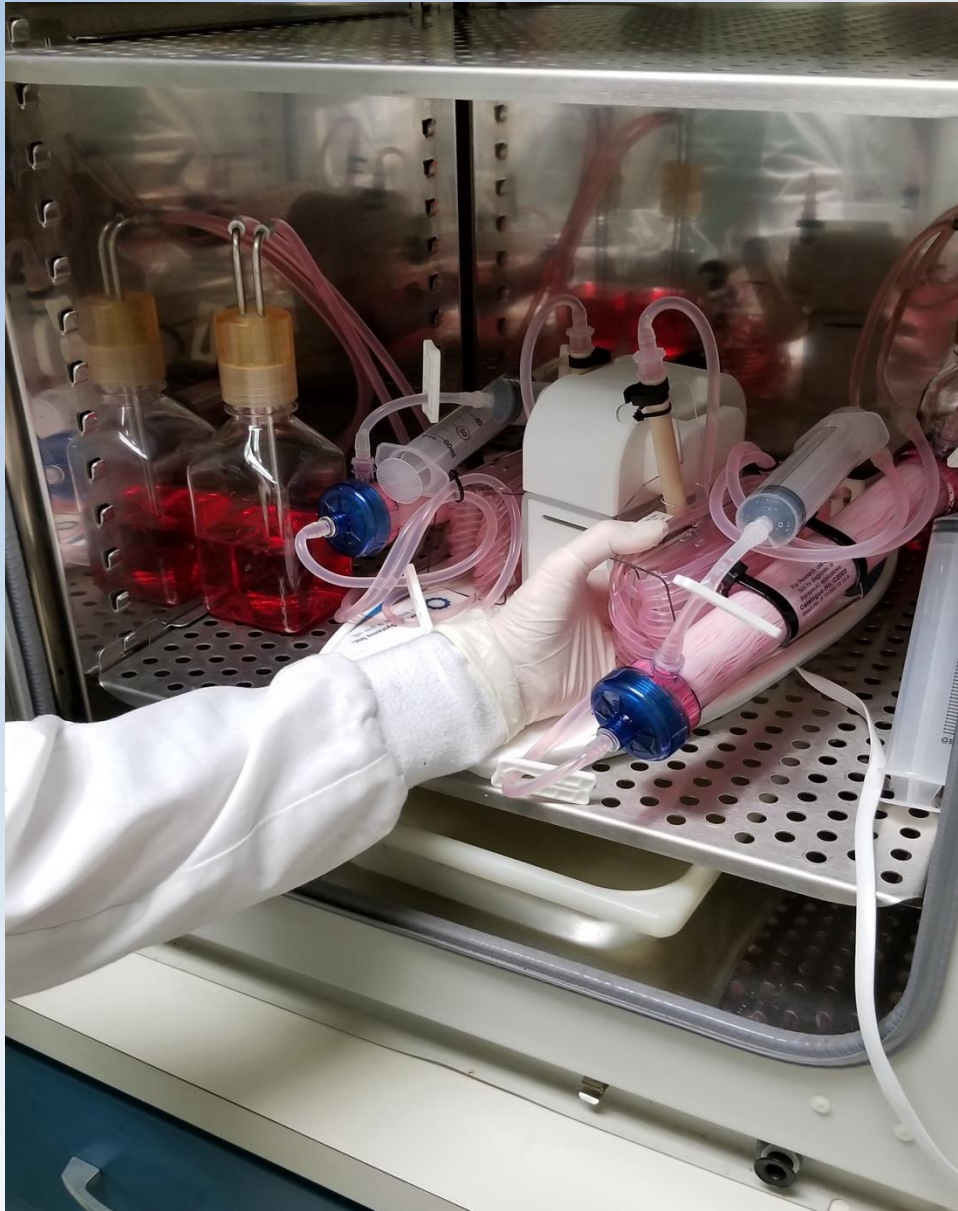
Feast or Famine in 2D Culture



CHO Cells, pH Changes in Hollow Fiber 3D Culture



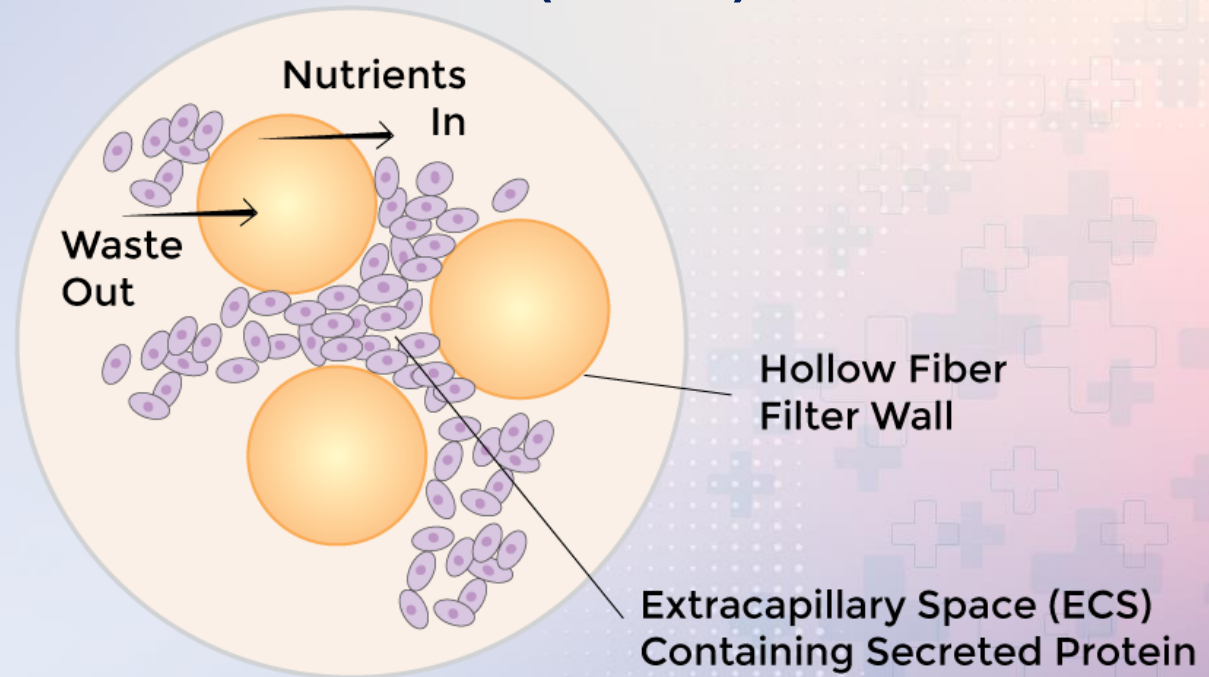
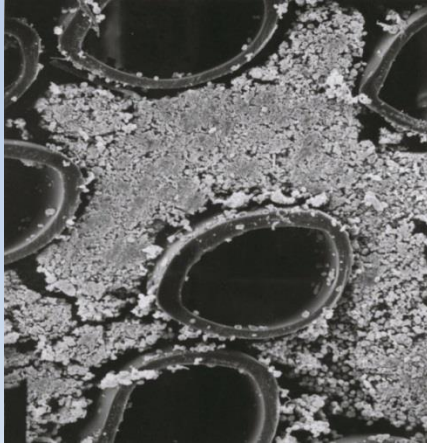
Scale-up Options in 3-D Cell Culture



Plastic Waste Generated by 1×10^9 Cells



What is a Hollow Fiber Bioreactor (HFBR)?



A 3-D Bioreactor for High-Density Cell Culture

Composition: Thousands of porous, semi-permeable capillary fibers in a cartridge.

Cell Residence: Cells grow in the *extracapillary space (ECS)*, bound to the porous fibers.

Media Flow: Culture medium is pumped through the *lumen* of the fibers.

Exchange: Nutrients diffuse out; waste products diffuse in.

Product Retention: Secreted products (proteins, exosomes) are concentrated in the ECS.

***Hollow fiber was 3-D
before 3-D was cool!***

Principles of Histocentric Bioreactors & How it Applies to Hollow Fiber Bioreactors

Histocentric Bioreactors recreate the *in vivo* microenvironment while allowing cell physiology to develop over time. 2-D, 3-D and now 4-D cell culture.

- Biological Mimicry
- Controlled Environment
- Dynamic Culture
- Cell Derived Microenvironment
- Compatibility With Various Cell Types
- Long Term Culture

Key Advantages of Histocentric HFBR

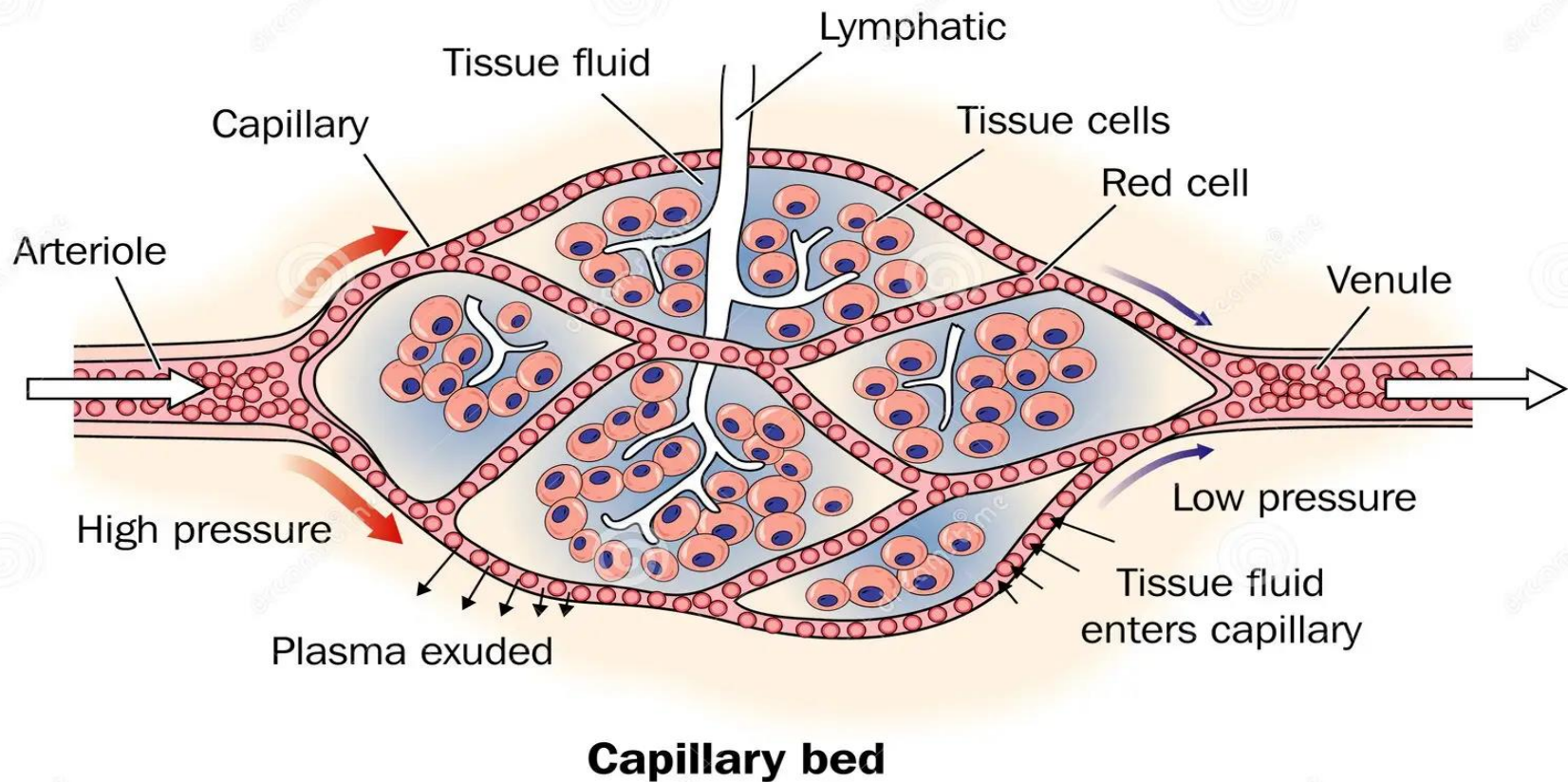
- Extremely high surface area/volume permits high density cell culture.
- Cells are bound to a porous support, not a non-porous 2-D flask.
- The molecular weight cut off (MWCO) of the fibers retains and concentrates secreted products.



Illustration of how cells grows *in vivo* compared to HFBR

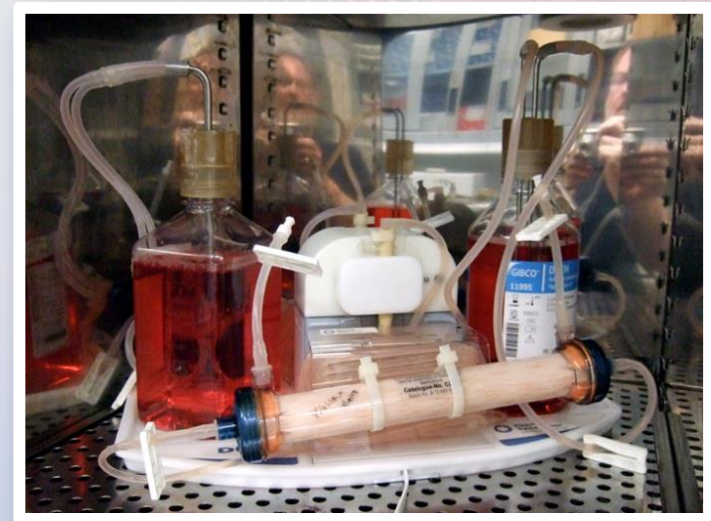
Histocentric Bioreactor attempts to recapitulate the *in-vivo* microenvironment.

Q: How many different medias are there in the human body?



Other Advantages of Hollow Fiber Cell Culture

- + Concentrated product
- + Uniform and complete post-translational modifications
- + Low apoptosis, less contamination with intracellular proteins and DNA
- + Consistency of production over many months
- + Protein-free medium (CDM-HD)
- + Histocentricity

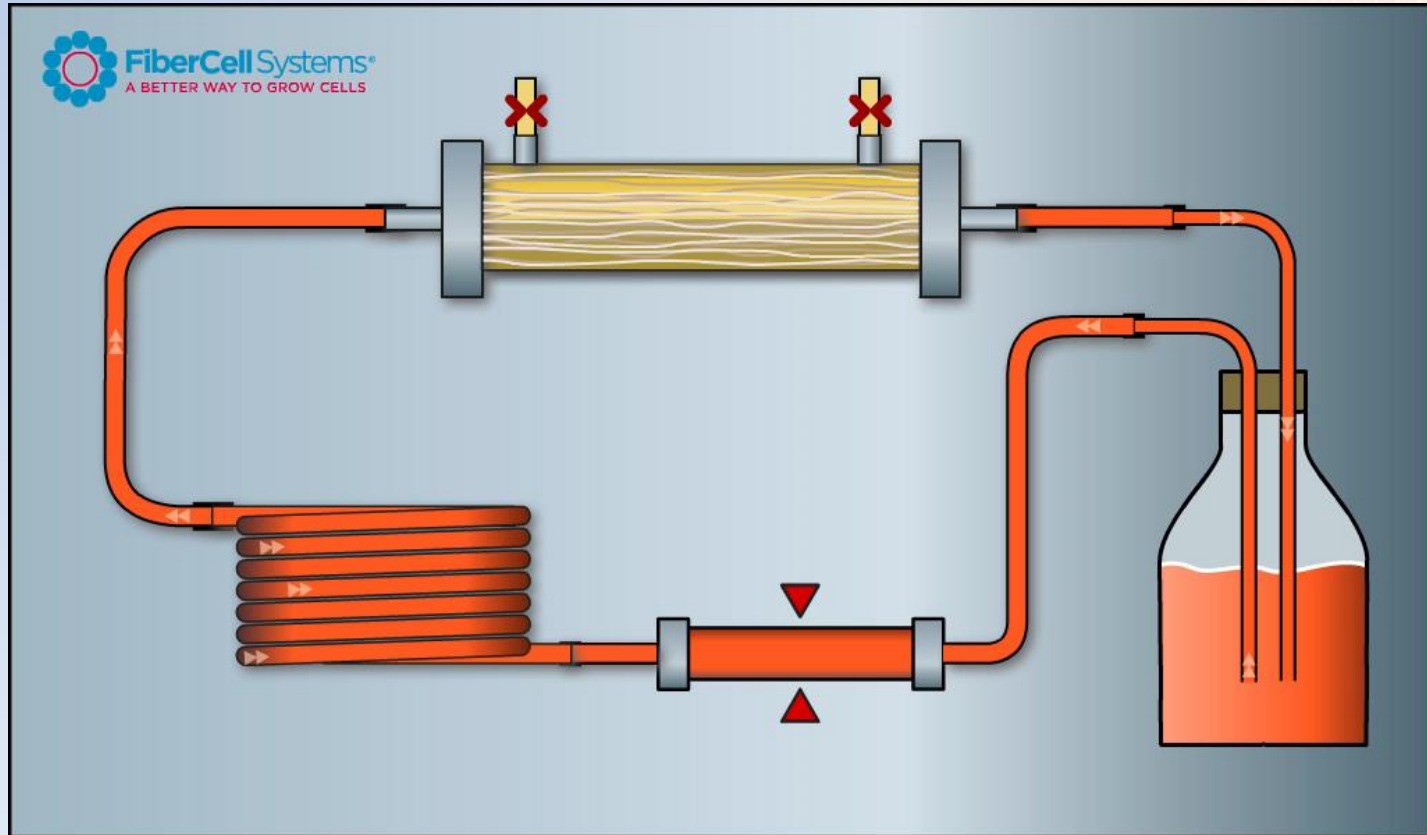


HFBR Applications

- + Monoclonal antibody production
- + Recombinant protein production
- + Conditioned medium
- + Exosome production
- + Endothelial cell culture under shear stress
- + Cell co-cultivation
- + Antibiotic Pk/Pd



Schematic of FiberCell Systems' HFBR



- + Positive pressure displacement pump
- + Silicone tubing for gas exchange
- + Closed, bio-safe system

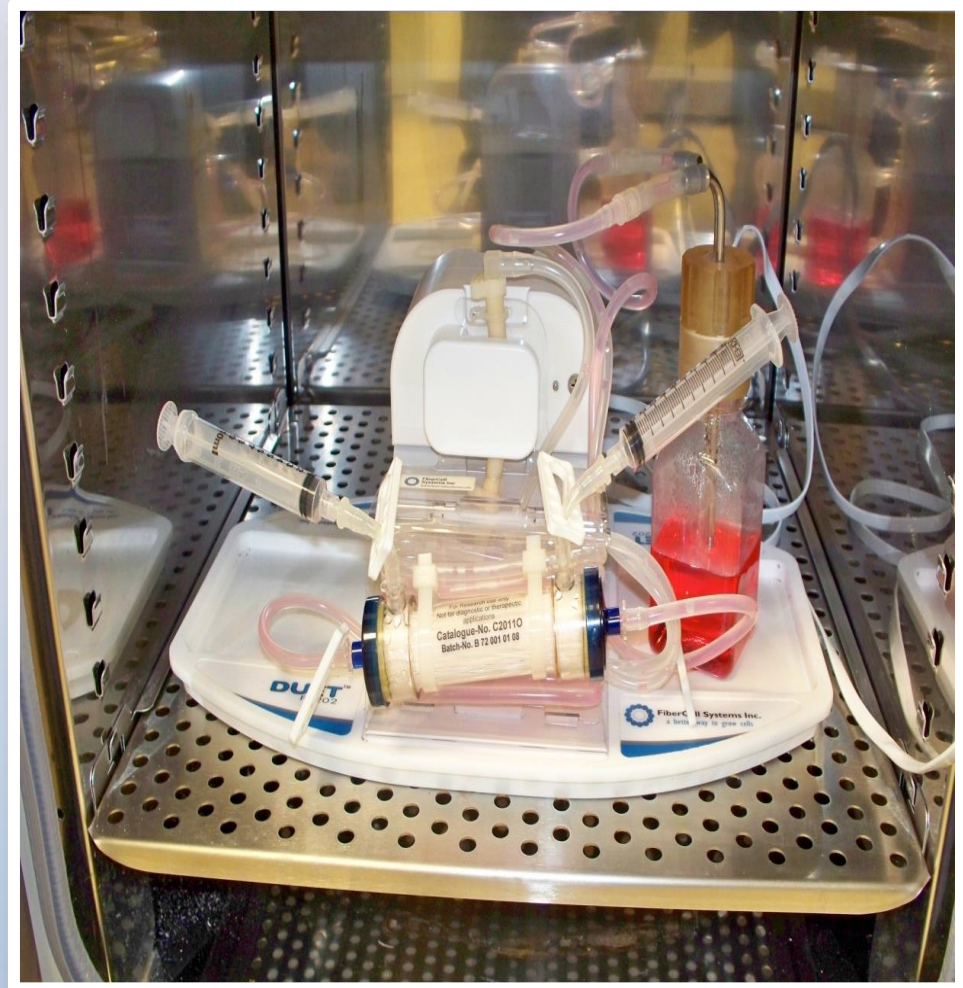
Illustration of how cell grow at high density



Best for collecting secreted products!

HFBR In the Laboratory

- + Fits in any standard sized incubator
- + Gas controlled by incubator
- + Temperature controlled by incubator
- + Thin cord for power



Working with HFBR In the Laboratory

- + Moves easily into hood
- + Good sterile technique always a plus
- + Maintenance only 15 minutes per day
- + Harvest product and measure glucose consumption



Good Sterile Technique is Always a Plus!



Glucose in Cell Culture: The Warburg Effect

The Warburg effect, also known as aerobic glycolysis, is a metabolic phenomenon observed in cancer cells where they convert glucose into lactate even in the presence of oxygen. (primary vs. transformed cells)

- Physiologic glucose: 80-90 mg/dl (.8 to .9 grams per liter)
- Cell culture medium 1-5 grams per liter
- Glucose has 4 calories per gram
- 100 kg person, 2,000 calories or 500 grams of glucose per day
- 5 grams per kilo of tissue
- Bioreactors for transformed cells consume 1 gram of glucose per 1×10^9 cells (1 gram) per day

Our CDM-HD Serum Replacement Media

- + Optimized and simplified for HFBR
- + Contains no surfactants
- + Chemically defined, protein-free
- + cGMP compliant
- + Lot-to-lot consistency
- + Ship at ambient, store at 4°C
- + Does not support cells at low density



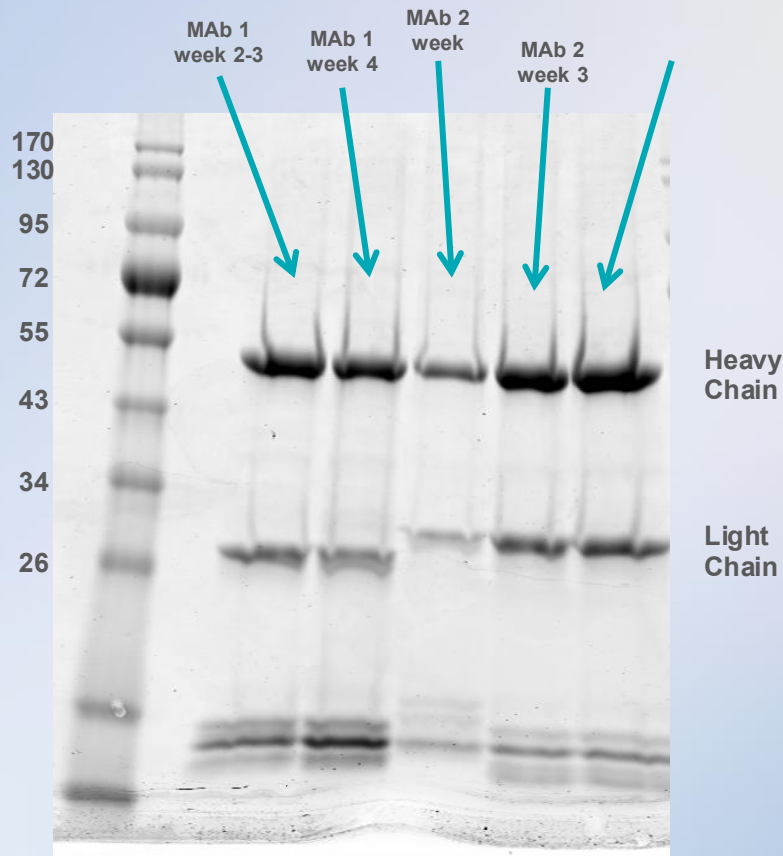
Mab Production using CDM-HD

Mab 1:

- 168 mg in 60 mL volume, 2.8 mg/mL.
- 9 L of medium consumed, 3 weeks culture.

Mab 2:

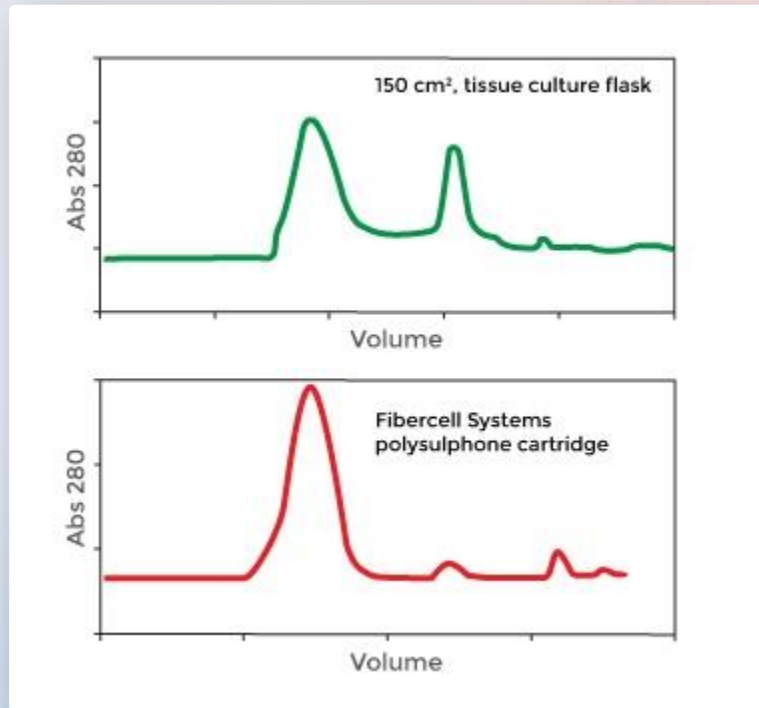
- 159 mg in 70 mL volume, 2.3 mg/mL.
- 11 L of medium consumed, 3 weeks culture.



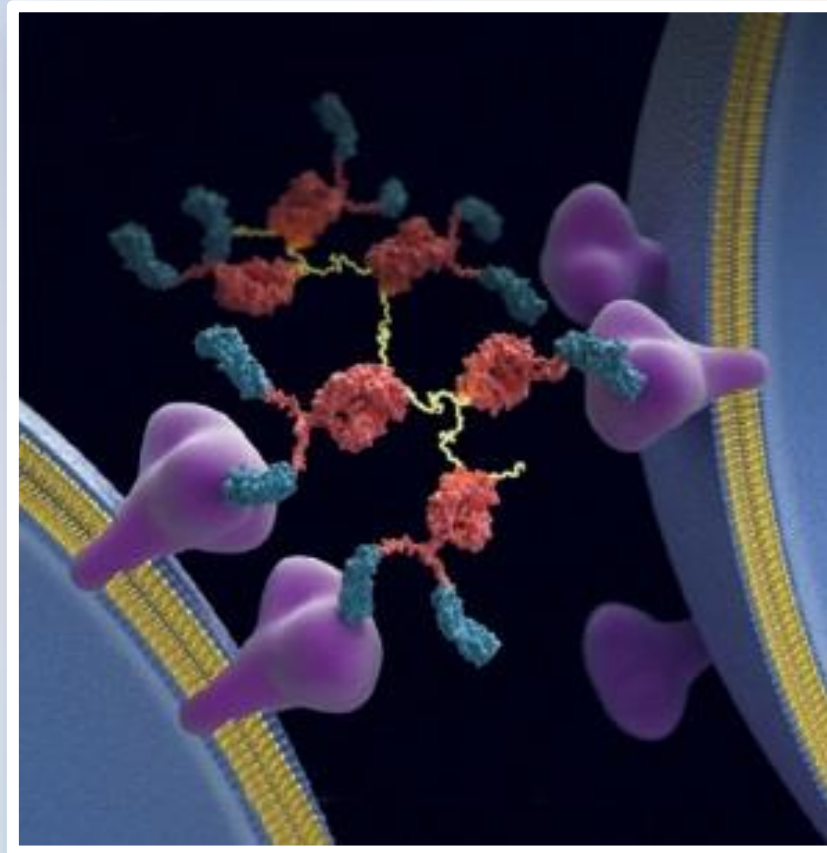
- TGF Beta diffuses out
- Mab trapped in ECS
- Easily adapt to SFM/CDM-HD
- Lower endotoxin
- .5 to 5 mg/mL conc.
- 5-100 mg per harvest
- Continuous production for over 6 months

Recombinant Protein Production

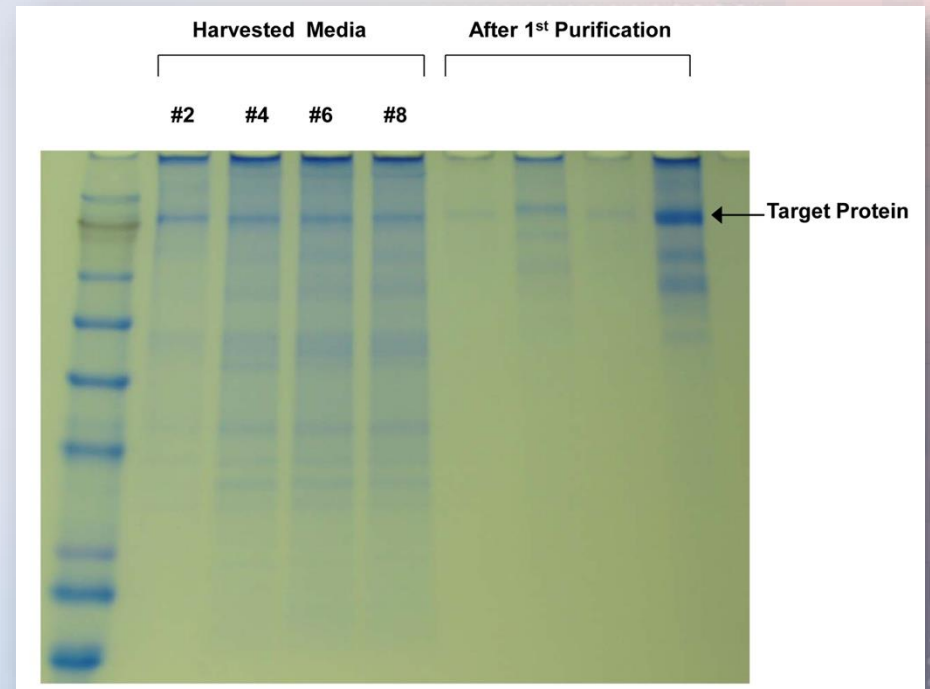
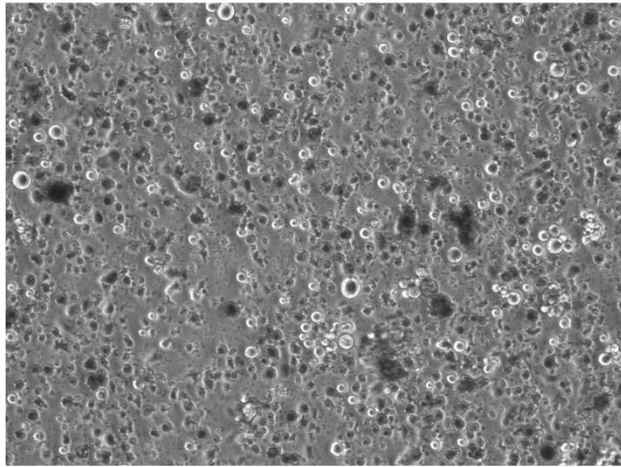
- + Both suspension and adherent cell types
- + 100x+ higher concentration
- + Easily adapt to SFM
- + Can provide improved protein folding



Journal of Biological Chemistry, Sept 2007



Raw Harvests from DG44 CHO Cell Line

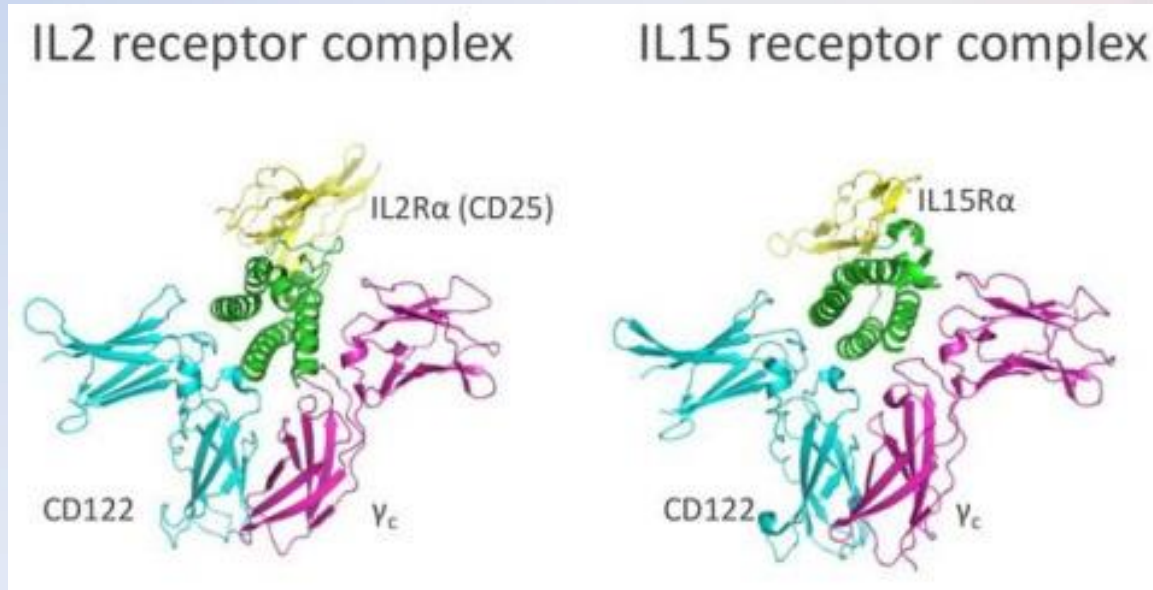


Difficult to Express Proteins

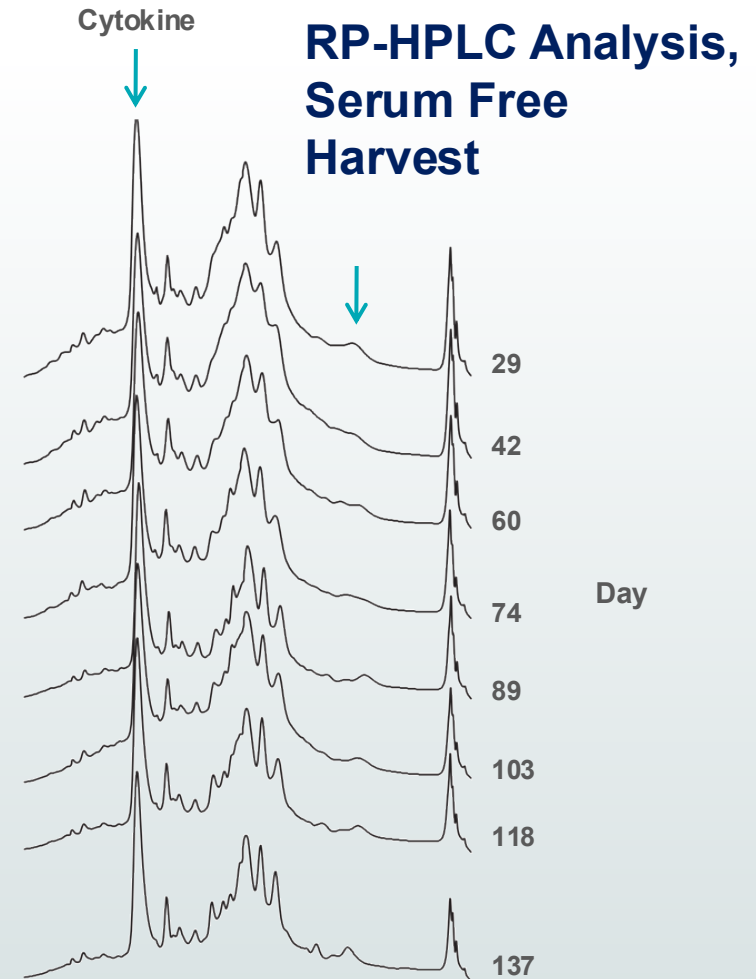
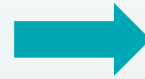
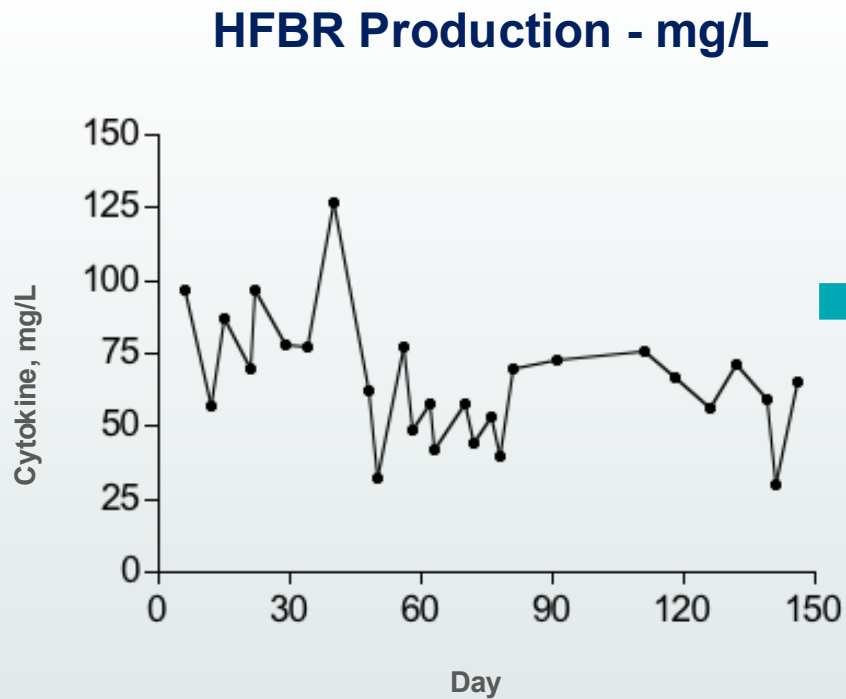
- Highly Glycosylated
- Large and/or Unstable
- Low Titers
- BITE and TRIKE, structures not found in nature

(Mammalian expression provides better solubility, longer serum $\frac{1}{2}$ life, better antigenicity and functionality)

IL15 RC is a Difficult to Express Protein

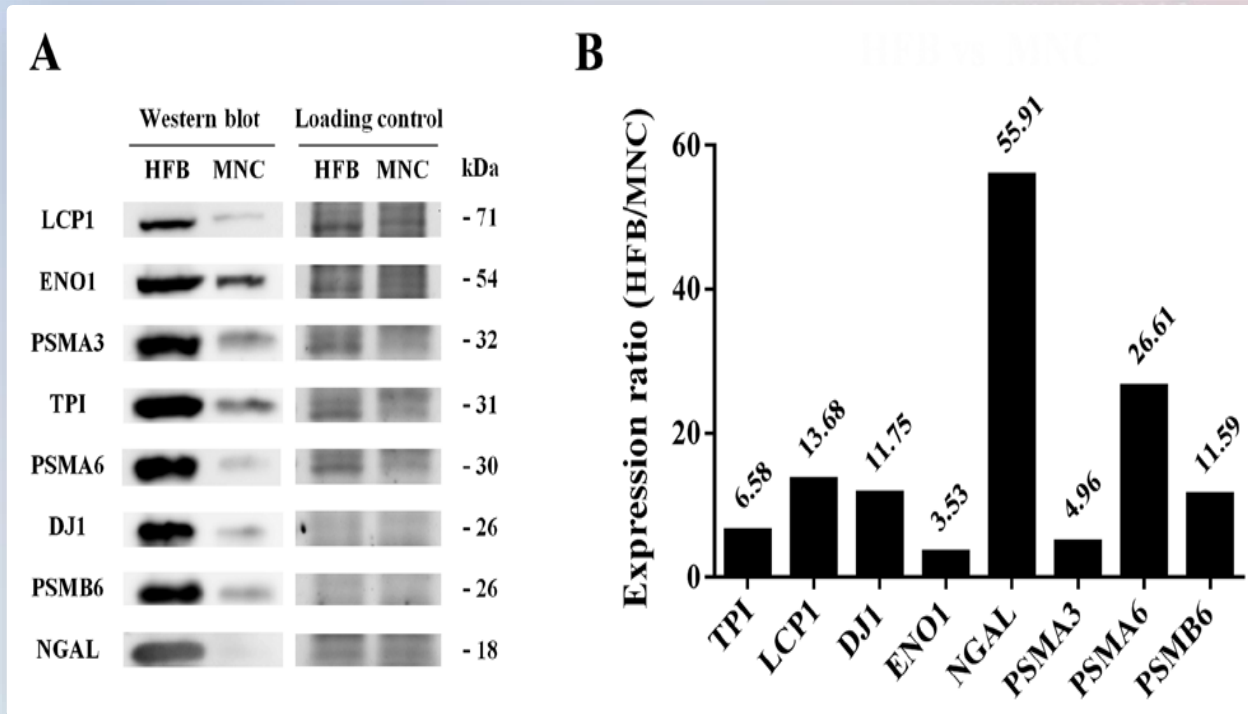


Stable IL15 RC Production in HFBR Over 5 Months

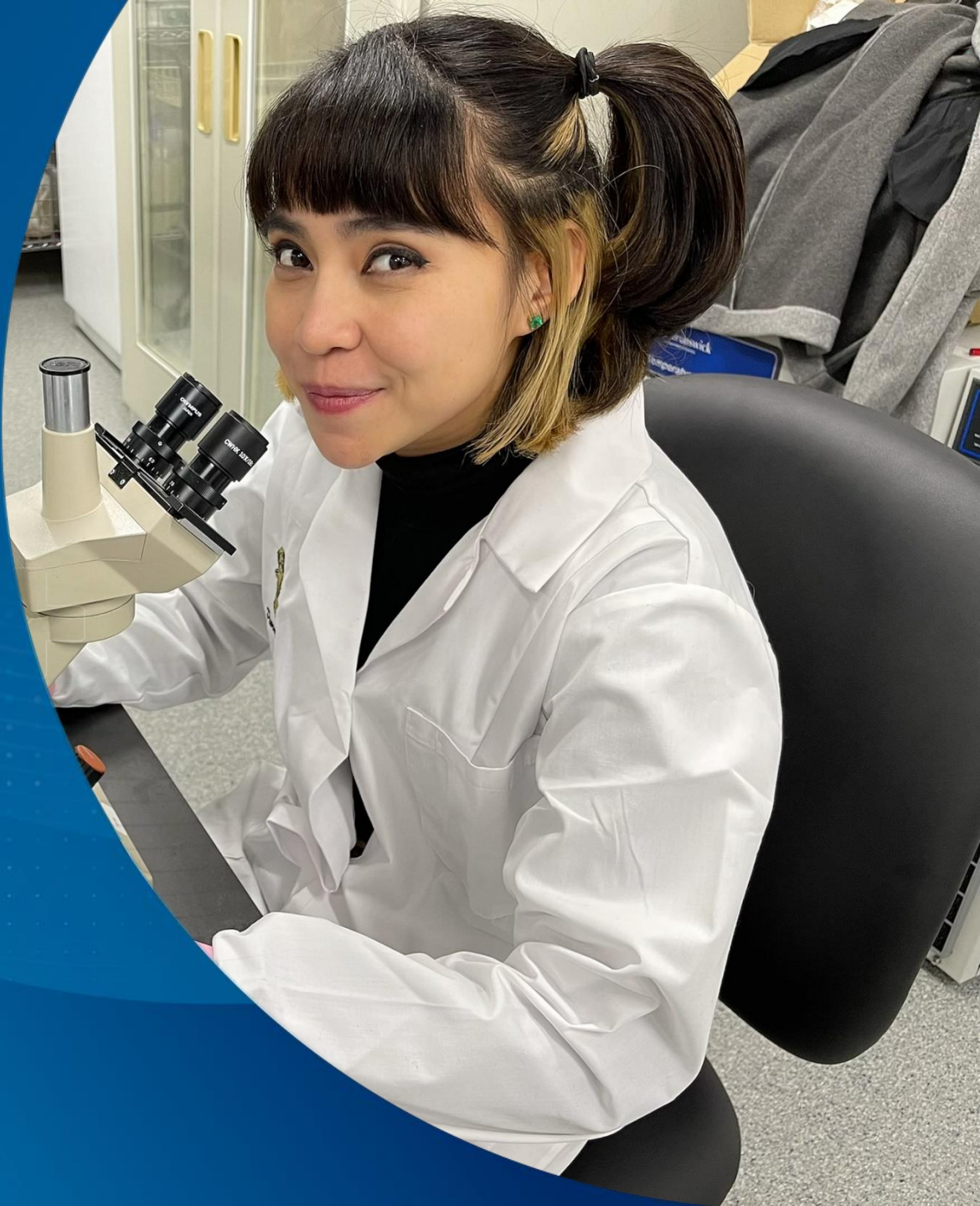


Secretome Analysis for Cancer Biomarker Discovery

Comparison of Flask vs. HFBR

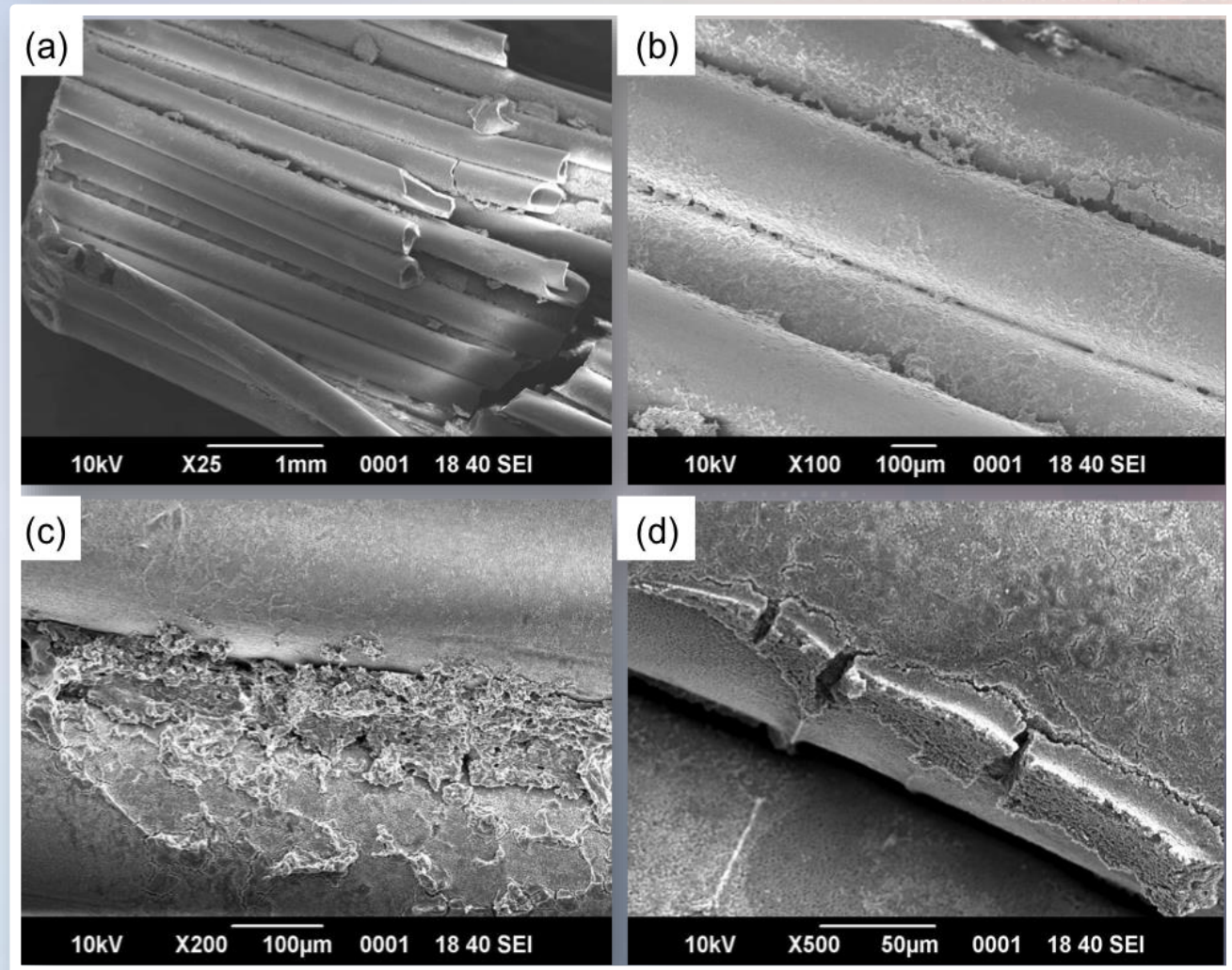


3-D
Models.....



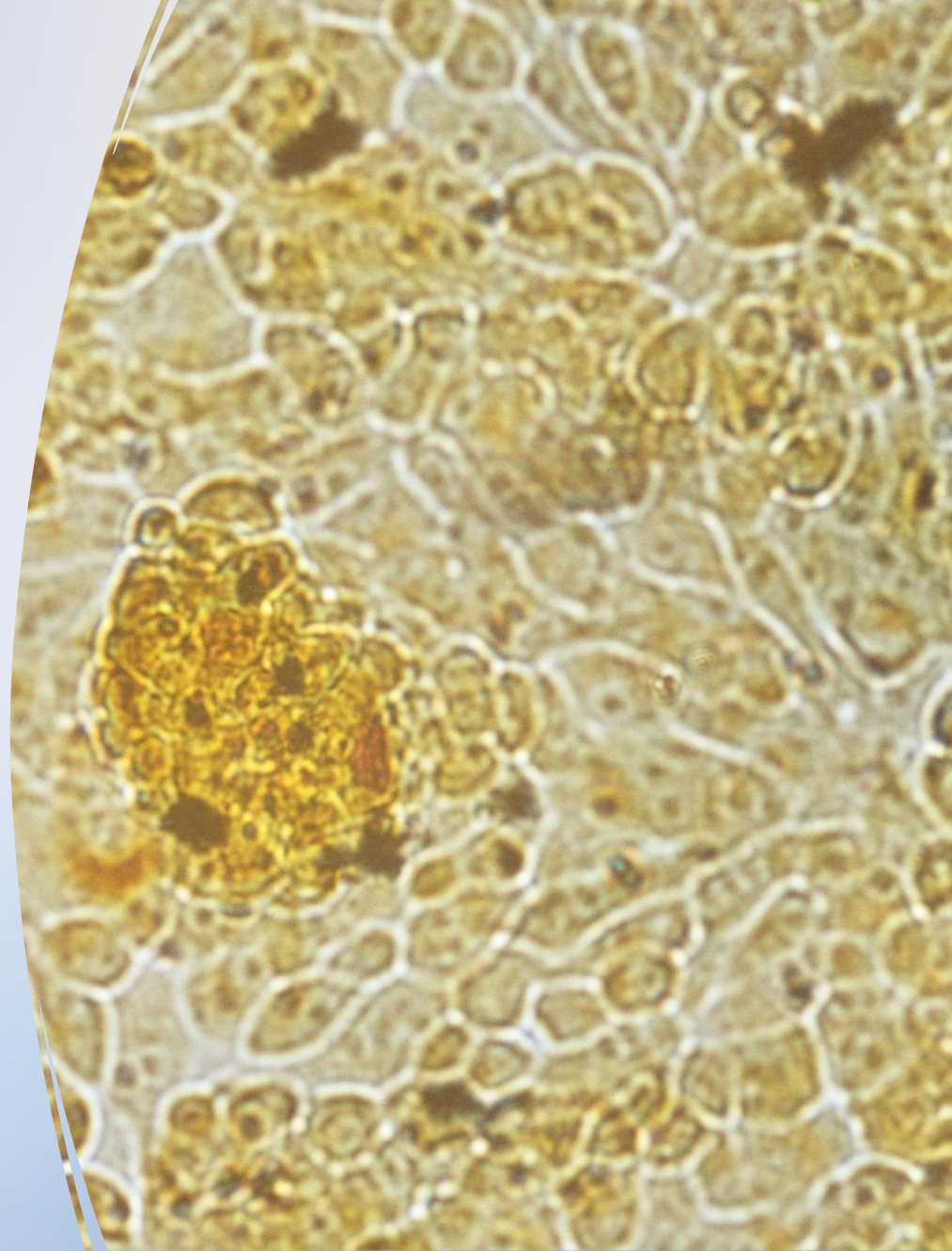
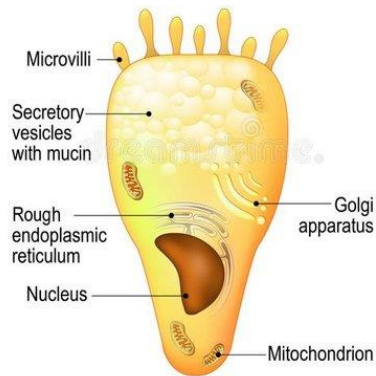
Bone Marrow Stroma/HSC Co-Culture

3-D CD 34 HSC engrafted more rapidly in nude mice compared to 2-D produced HSC.



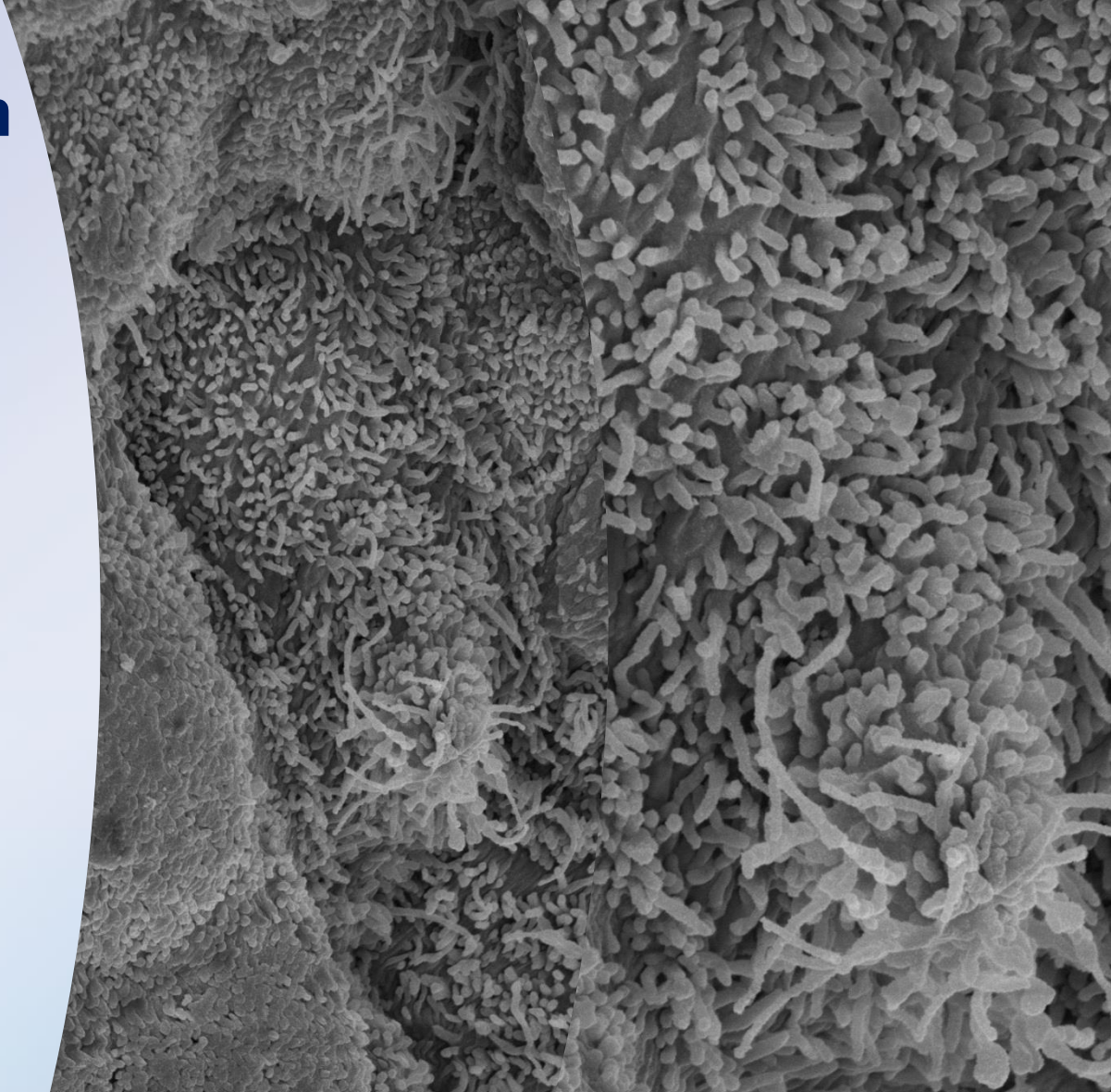
Mixed culture of 85% HCT-8 human intestinal epithelial cells and 15% LS174T goblet cells. Stained with bismark brown to show patches of goblet cells.

Goblet cell



Cryptosporidium

Intestinal Epithelial cells
cultured in 3-D perfusion
hollow fiber bioreactor
demonstrating polarity
and microvilli.



 **EINSTEIN**

Albert Einstein College of Medicine
OF YESHIVA UNIVERSITY

Mag = **EIN**

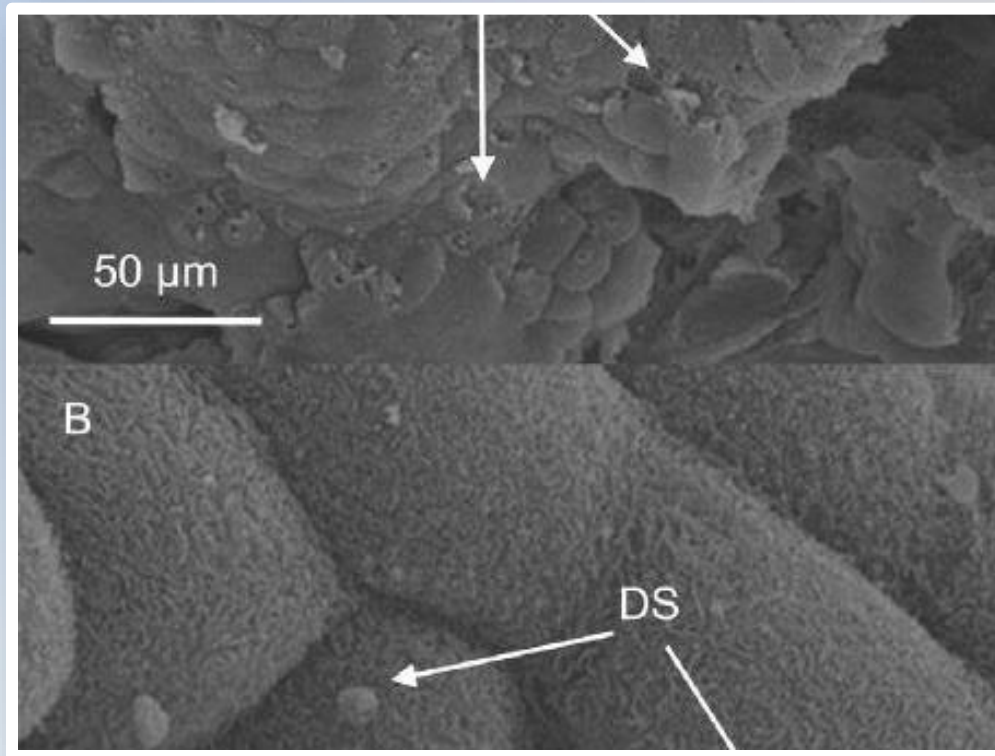
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OF YESHIVA UNIVERSITY

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Cryptosporidium Culture in an Artificial Gut Model

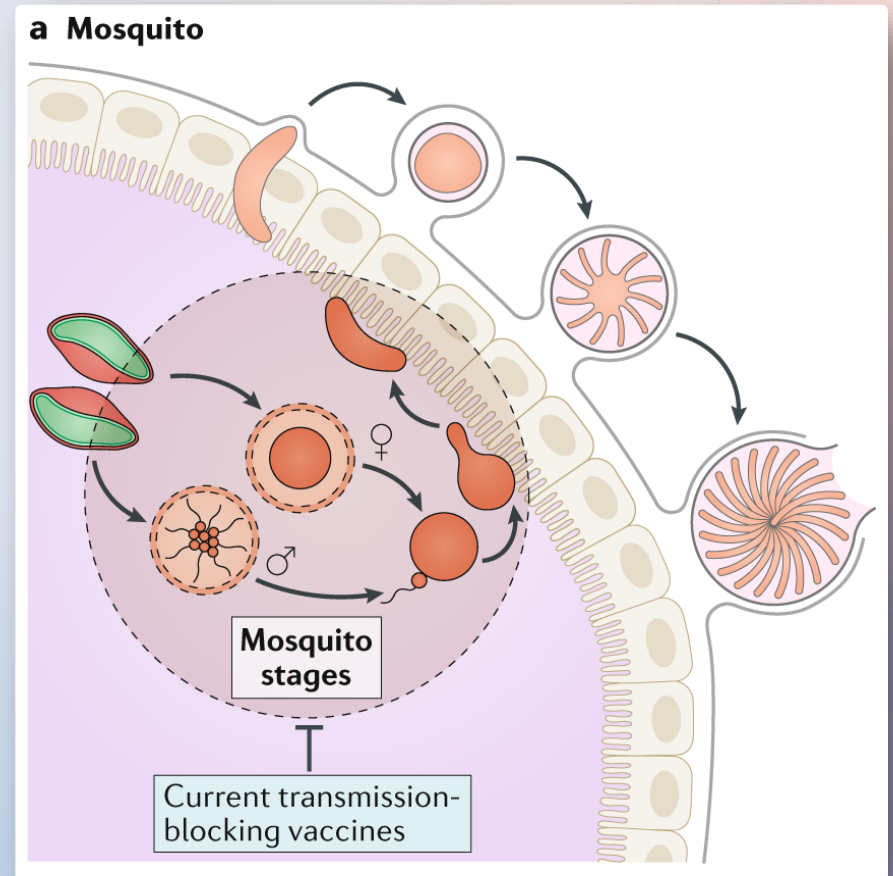


DS: Development stage.

Intracellular but extra-cytoplasmic stage (meront), the enlarged parasitophorous vesicle filled with 8 merozoites which are released when the meront bursts

Malaria Vaccine

Malaria parasites exhibit a complex lifecycle, requiring extensive asexual replication in the liver and blood of the vertebrate host, and in the haemocoel of the insect vector. Yet, they must also undergo a single round of sexual reproduction, which occurs in the vector's midgut upon uptake of a blood meal.




Malaria sporozoites are produced in nature in mosquitoes. In December 2022, a team from Sanaria Inc., a malaria vaccine company, published their groundbreaking working reporting that they could produce infectious *Plasmodium falciparum* (Pf) malaria sporozoites (SPZ) *in vitro* (i). The goal is to use these iPfSPZ in a vaccine.

Article

***In vitro* production of infectious *Plasmodium falciparum* sporozoites**

<https://doi.org/10.1038/s41586-022-05466-7>
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Accepted: 20 October 2022
Published online: 07 December 2022

 Check for updates

Abraham G. Eappen¹, Tao Li¹, Meghan Marquette¹, Sumana Chakravarty¹, Natasha KC^{1,2}, Gigliola Zanghi³, Benjamin U. Hoffman^{4,5}, Hashani Hettiarachchi^{1,6,7}, Asha Patil¹, Yonas Abebe⁸, Christiane Tran¹, Alemtaye A. Yossef¹, Ian McWilliams¹, Robert D. Morrison⁹, Ayyappan Rathakrishnan¹, Ehud Inbar³, Ahmed S. I. Aly¹, Patricia De La Vega¹, Maria Belmonte¹⁰, Martha Sedegah¹, Tint Wai¹¹, Joseph J. Campo¹, Harley King⁸, Stefan H. I. Kappe^{8,9,10}, MingLin Li^{1,2}, Peter F. Billingsley¹, B. Kim Lee Sim^{1,2}, & Stephen L. Hoffman^{1,2}

An effective vaccine is needed for the prevention and elimination of malaria. The only immunogens that have been shown to have a protective efficacy of more than 90% against human malaria are *Plasmodium falciparum* (Pf) sporozoites (PfSPZ) manufactured in mosquitoes (mPfSPZ)^{1–7}. The ability to produce PfSPZ *in vitro* (iPfSPZ) without mosquitoes would substantially enhance the production of PfSPZ vaccines and mosquito-stage malaria research, but this ability is lacking. Here we report the production of hundreds of millions of iPfSPZ. iPfSPZ invaded human hepatocytes in culture and developed to mature liver-stage schizonts expressing *P. falciparum* merozoite surface protein 1 (PfMSP1) in numbers comparable to mPfSPZ. When injected into FRGhuHep mice containing humanized livers, iPfSPZ invaded the human hepatocytes and developed to PfMSP1-expressing late liver stage parasites at 45% the quantity of cryopreserved mPfSPZ. Human blood from FRGhuHep mice infected with iPfSPZ produced asexual and sexual erythrocytic-stage parasites in culture, and gametocytes developed to PfSPZ when fed to mosquitoes, completing the *P. falciparum* life cycle from infectious gametocyte to infectious gametocyte without mosquitoes or primates.

Source: www.nature.com

Primary lung tissue inoculated as a single cell suspension and supported by CDM HD.

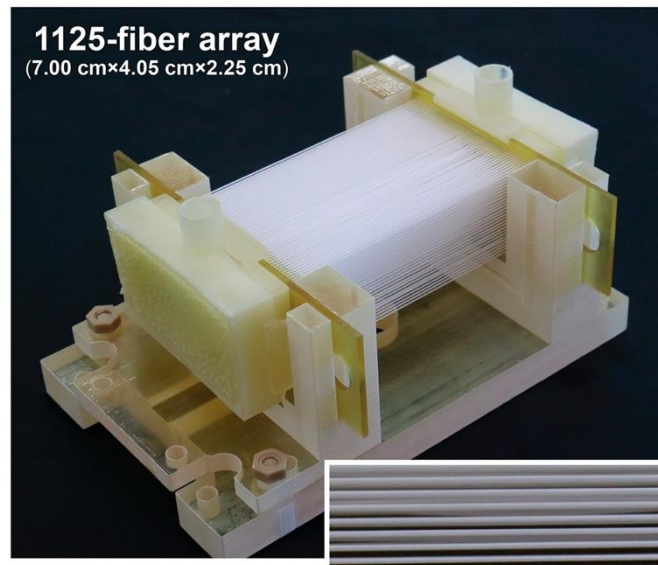
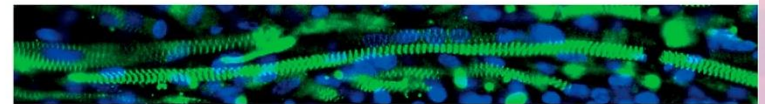
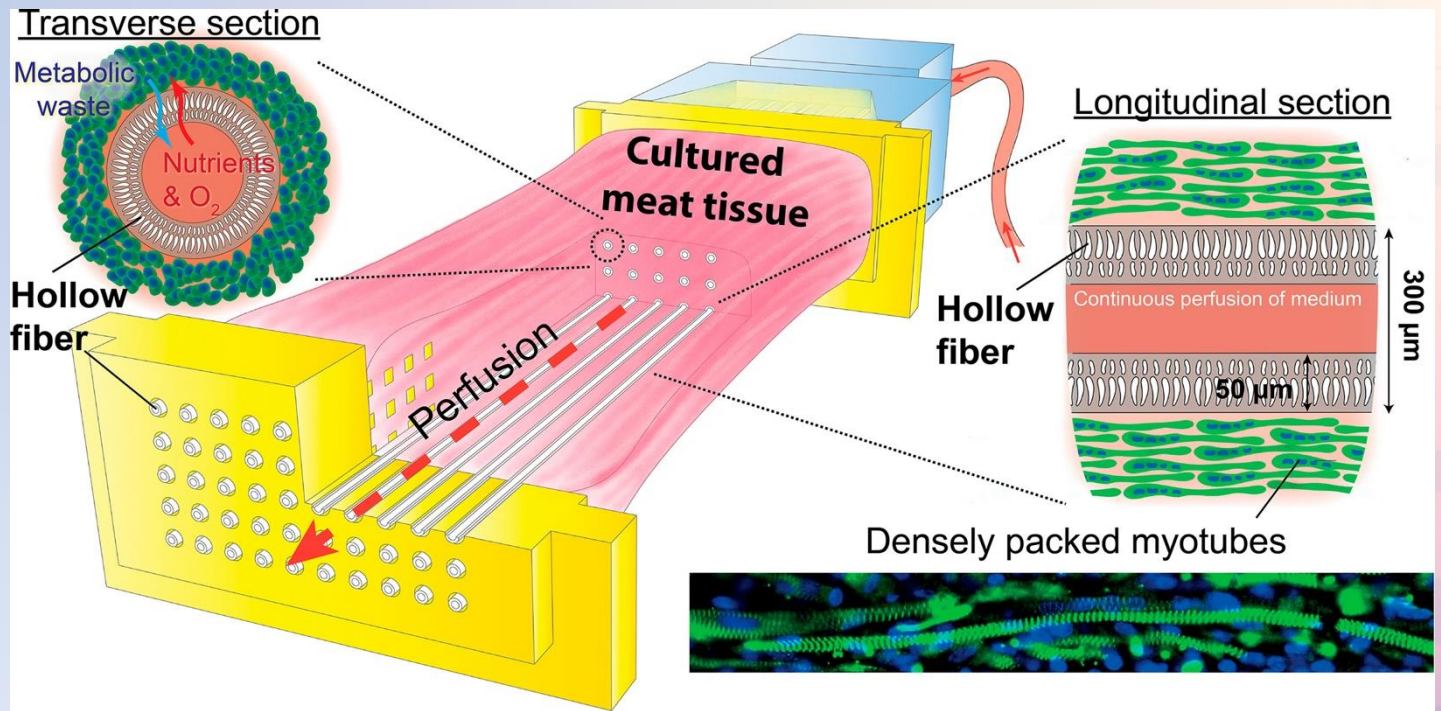
Dr. Akhil Srivastava, U. Missouri



Meat and Milk Production



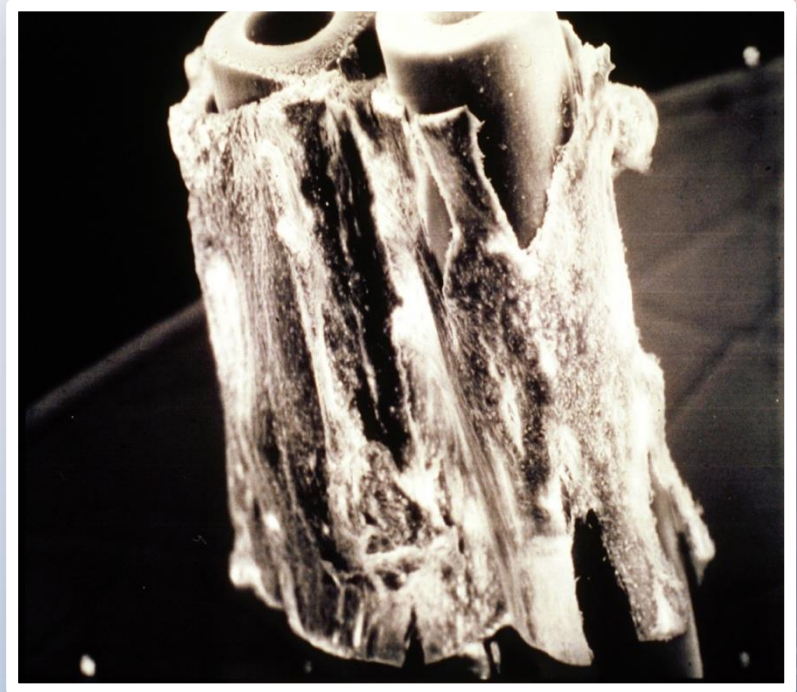
Making artificial meats in HFBR



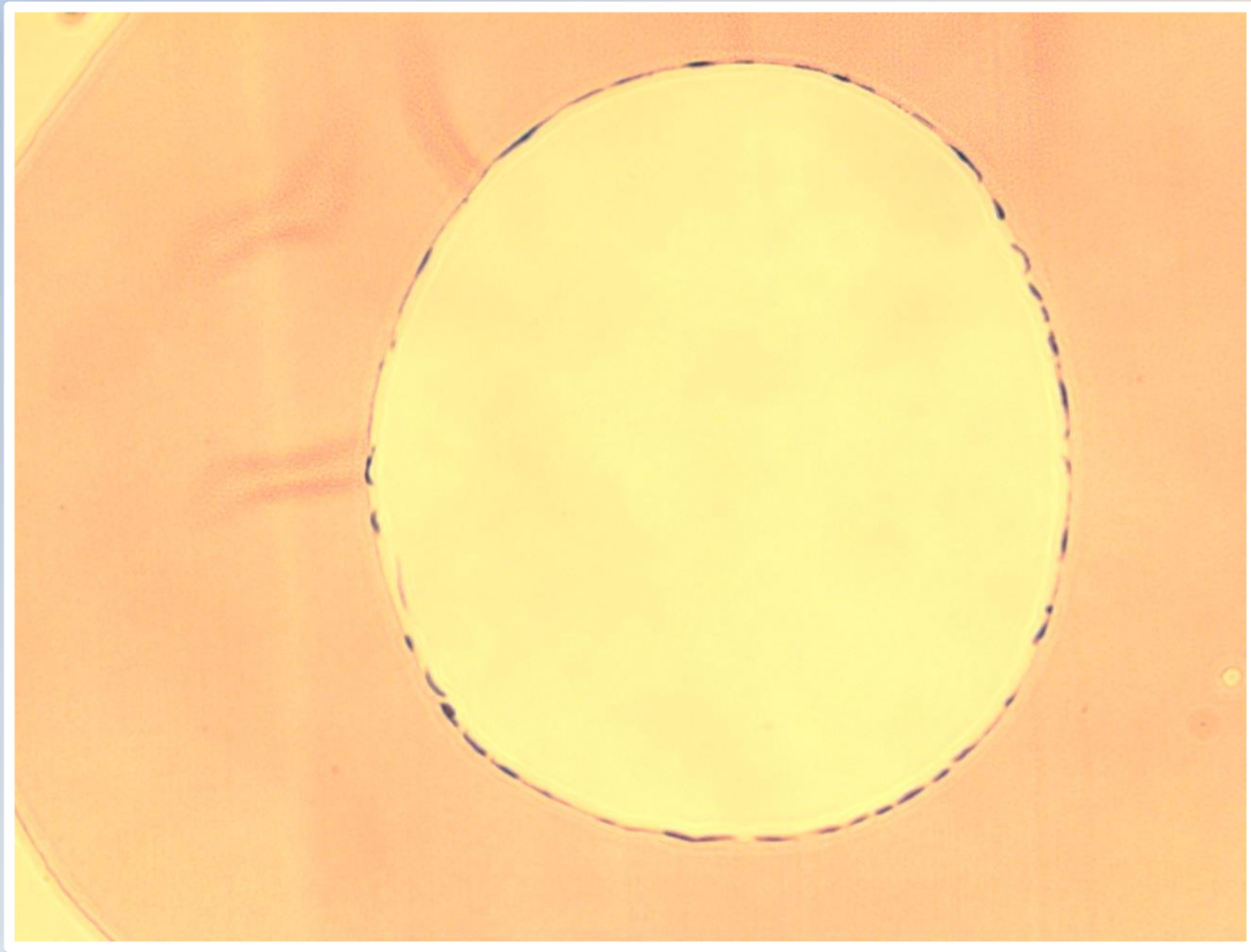
Trends in
Biotechnology
Volume 43, Issue 8
P1938-1960 August
2025

Asymmetric Cell Co-cultivation

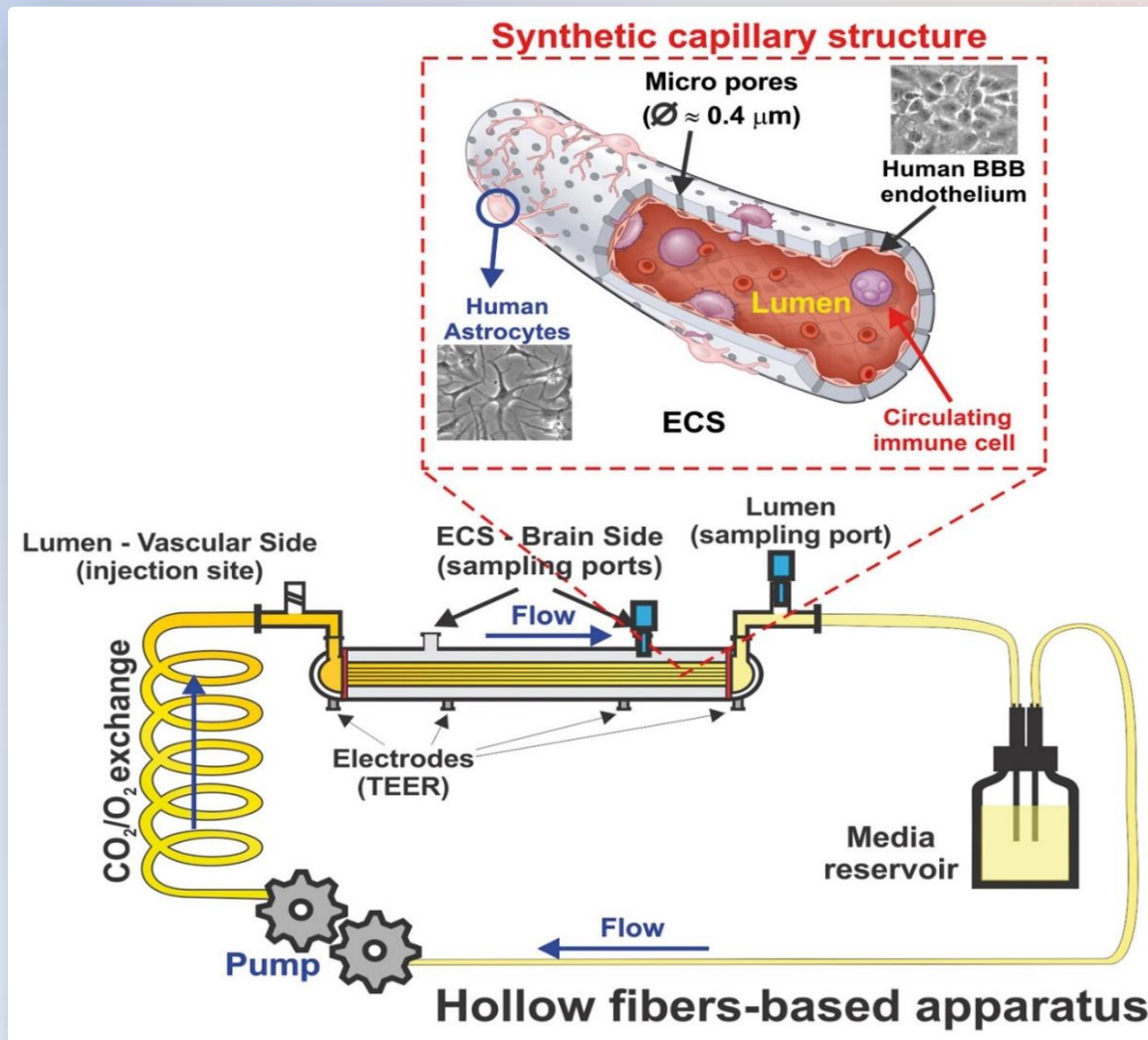
- + Co-cultivation of endothelial cells (inside) and vascular smooth muscle (outside)
- + Brain endothelial and astroglial cells to form in vitro blood brain barrier



Endothelial cells in HFBR

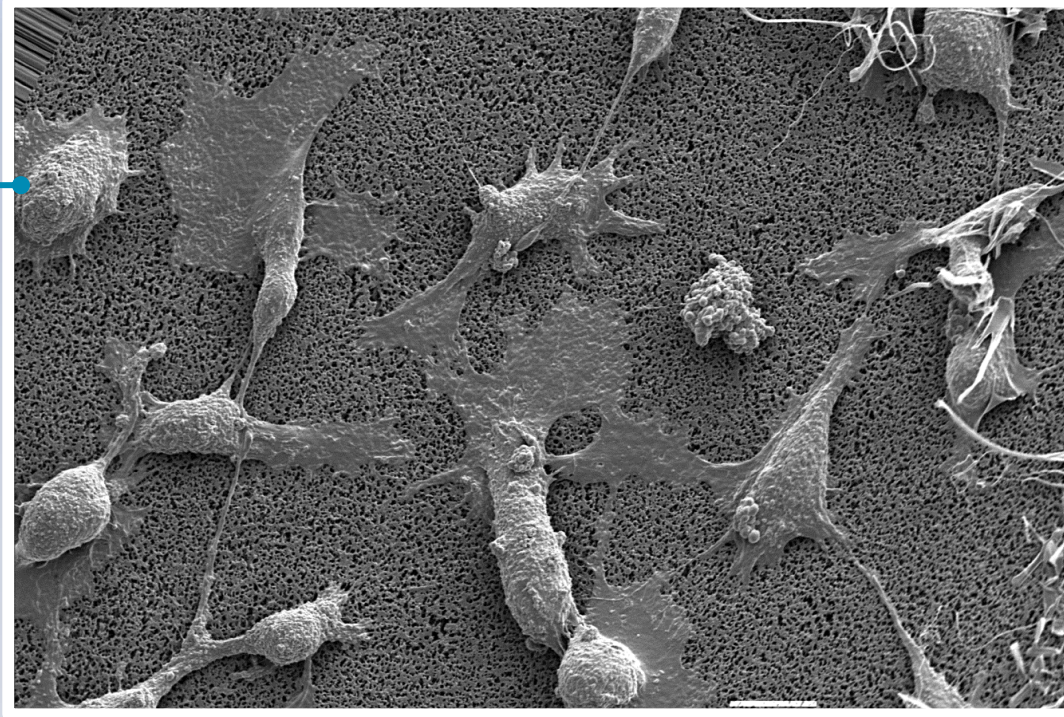


Blood-Brain Barrier Model



Endothelial cells grown in HFBR

Pores of fiber



Endothelial cells on the inside on a fiber- these have been adhered to the wall then subjected to very low shear force overnight followed by a few hours at 5 dynes/cm².

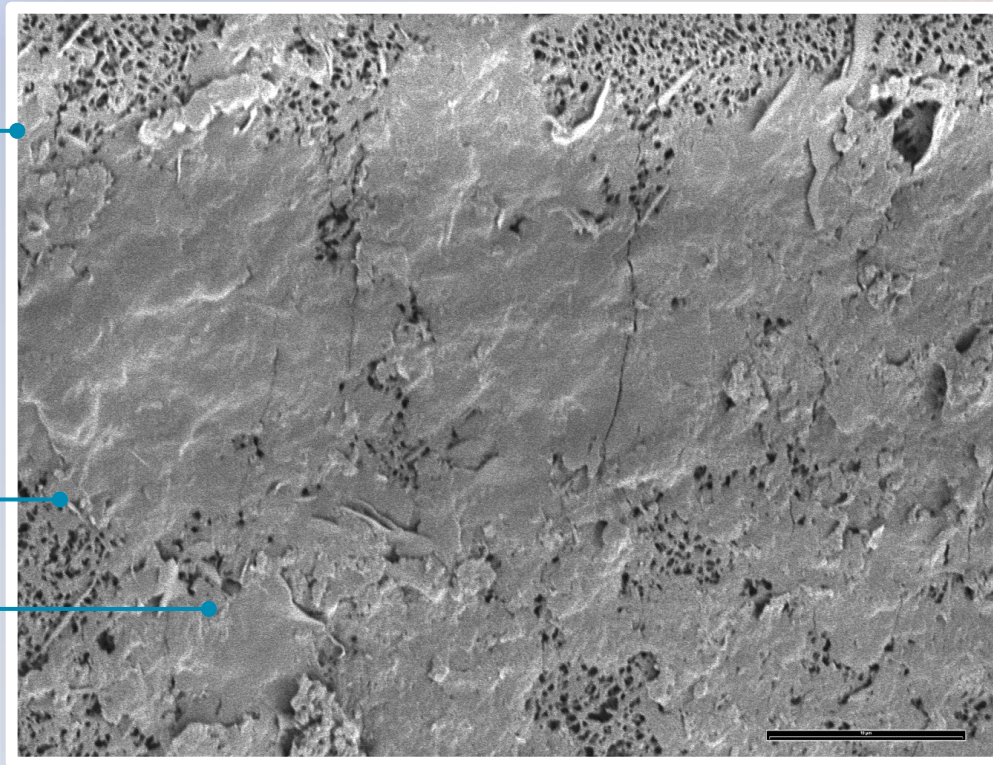
While the majority of the cells here are still bulky it is possible to see them begin to flattened down onto the wall of the fiber and really stretch out.

Endothelial cells grown in HFBR

Inside of
hollow fiber

Pore

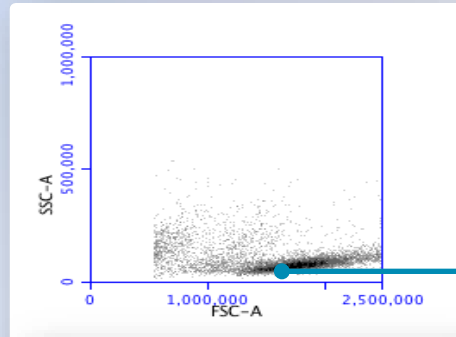
Endothelia
I cell



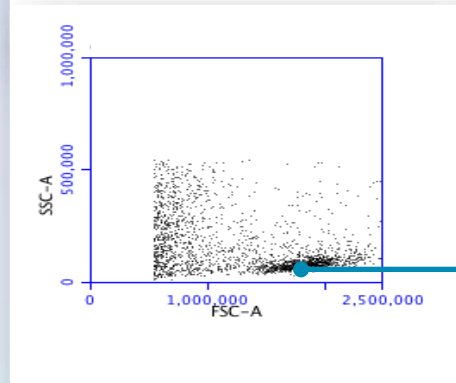
Flattened endothelial cells on inside of fiber. These endothelial cells had been adhered to the inside of the fiber and subjected to minimal shear force over night followed by a minimum of 5 hours at 10 dynes/cm².

CLL cells actively migrate into the extra-capillary space

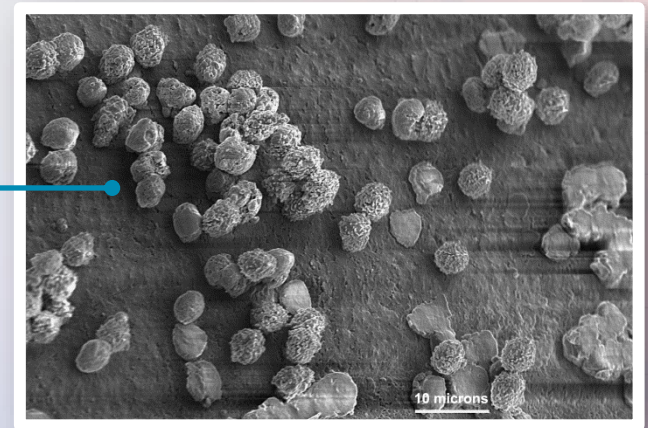
Circulating
Compartment



Extra-capillary
space (migrated
compartment)



CLL cells



Scanning electron
micrograph of the outside
of a hollow fibre after
circulation of CLL cells
around the system

Cardiff CLL Research Group

Exosome Cell Culture Conditions Affect EV Composition

Laminar Flow Alters EV Composition in HUVECs: A Study of Culture Medium Optimization and Molecular Profiling of Vesicle Cargo

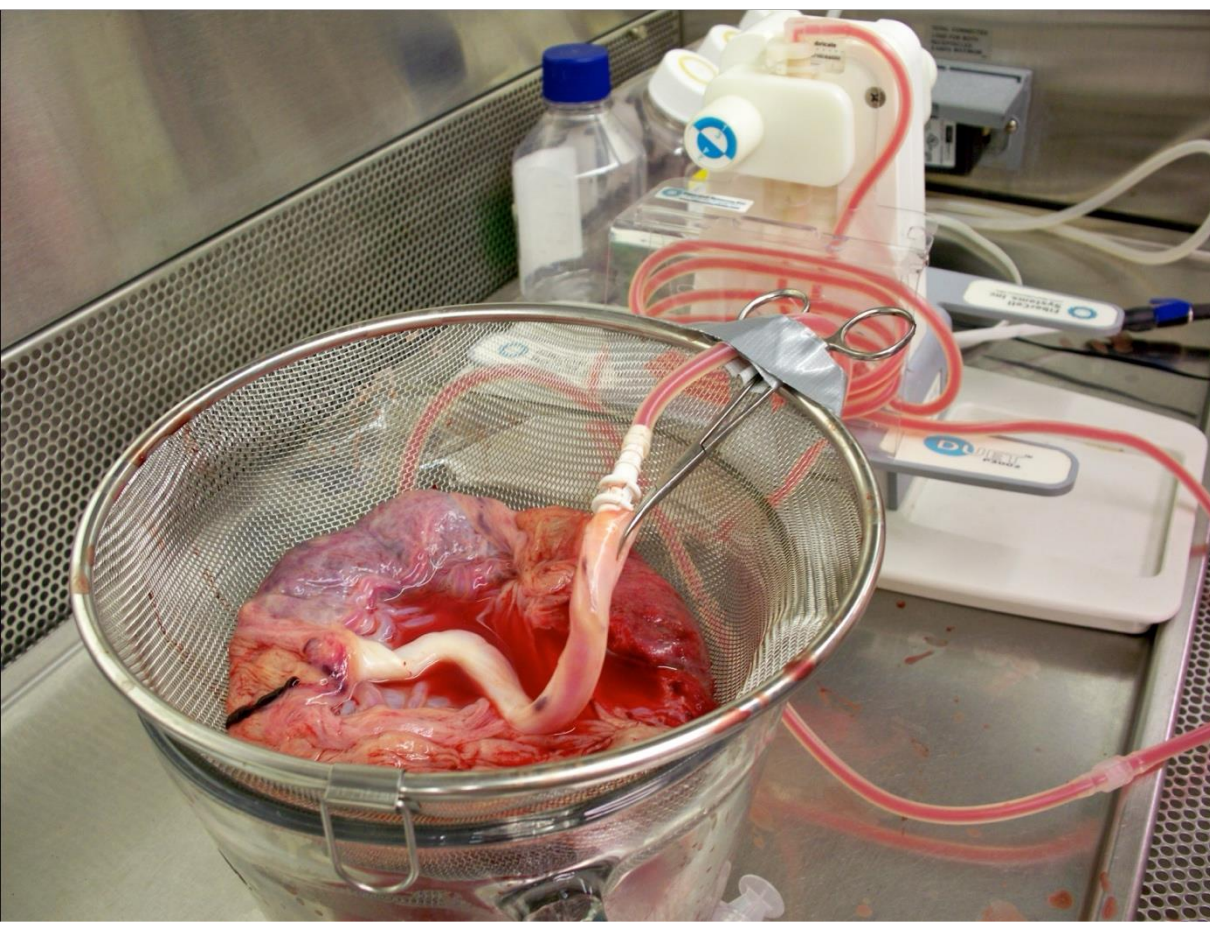
Arefeh Kardani, Jan Hemmer, Britta Diesel, Vida Mashayekhi, Annika Schomisch, Marcus Koch, Claudia Fecher-Trost, Markus R Meyer, Nicole Ludwig, Shusruto Rishik, Andreas Keller, Jessica Hoppstädter, Gregor Fuhrmann, and Alexandra K. Kiemer**

Endothelial cells (ECs) experience shear stress associated with blood flow. Such shear stress regulates endothelial function by altering cell physiology. Since most cell culture protocols and media compositions are designed for static cultures and experiments with ECs are predominantly conducted under these non-physiological conditions, a model for culturing ECs under flow conditions is developed, which more closely mimics their physiological environment. This approach also enables the isolation of EVs while minimizing FCS-derived contaminants. In this study, a comprehensive assessment of how physiologically relevant cultivation conditions influence the vesicle composition and function of ECs is provided. A detailed investigation is conducted for the effect of different cell culture media on morphology and marker expression of human umbilical cord endothelial cells (HUVECs) and EVs, and optimize the conditions to culture ECs under flow, tailoring them specifically to facilitate the efficient isolation of EVs using a hollow-fiber system model. These EVs are then characterized and compared to those isolated from traditional static culture conditions. Overall, this study presents a model on isolating EC-derived EVs under conditions that closely mimic physiological environments, and characterization at their proteome, gene expression, and microRNA profile.

1. Introduction

Extracellular vesicles (EVs) are nanosized membrane-bound structures released by almost all types of cells into their external environment. Eukaryotic EVs are usually classified into three main categories, based on their size and mode of production.^[1] Microvesicles are formed by the outward budding of membrane vesicles from the cell surface.^[2] Exosomes originate from the endocytic pathway through the 'outward' budding of the late endosomal membrane. Initially, they accumulate in structures known as multivesicular bodies (MVBs), which later fuse with the plasma membrane, releasing their contents as exosomes into the extracellular space.^[3] The third major type of eukaryotic EVs called apoptotic bodies are produced from cells undergoing programmed cell death by outward budding from the surface of apoptotic cell.^[4]

Pulsatile Perfusion of Placenta



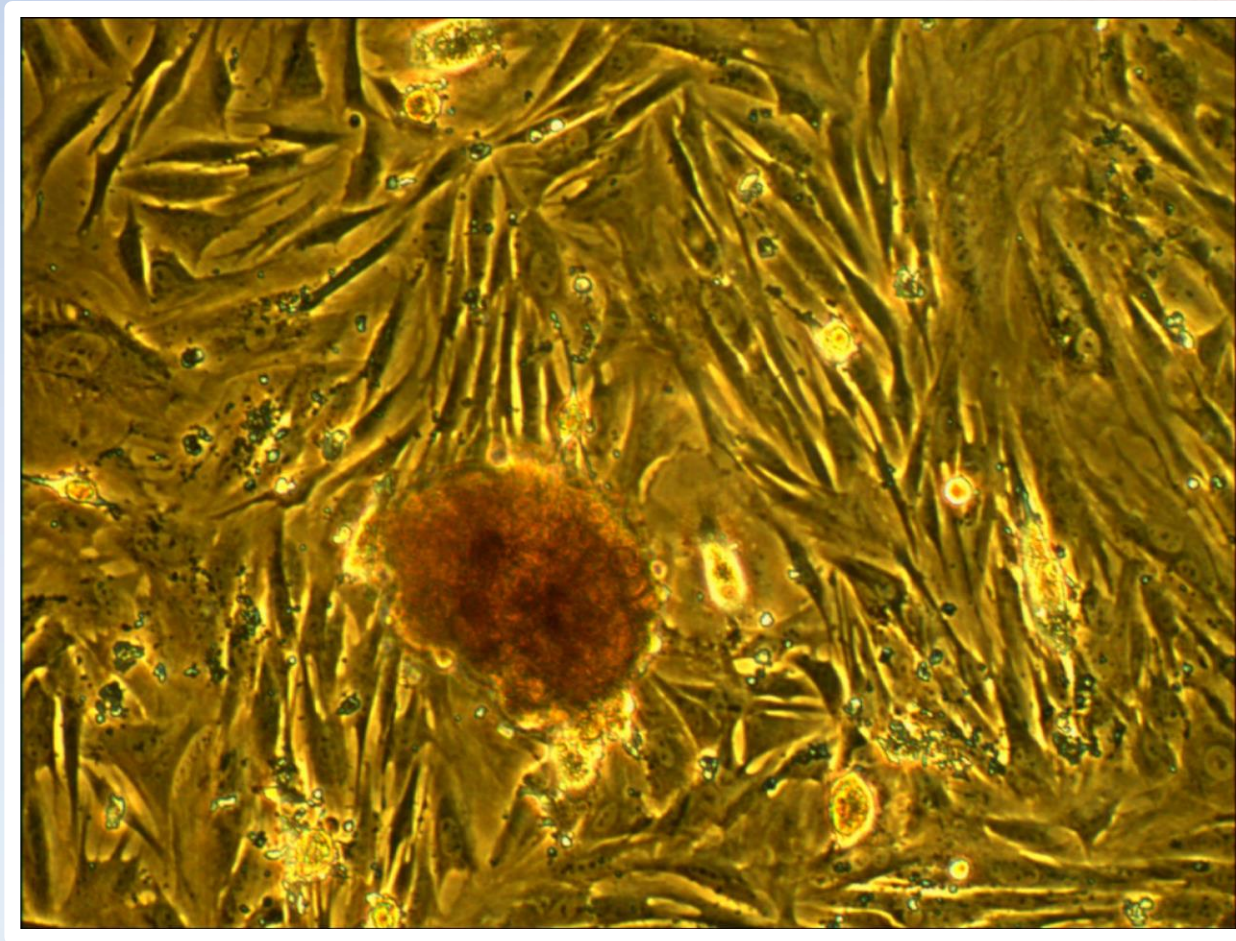
- 1) Flush with PBS
- 2) Digest with collagenase for one hour
- 3) Collect cells
- 4) Seed into hollow fiber bioreactor

Placental Co-Culture



- 1) Remove non-adherent cells after 3 days
- 2) Nodules form in 5-7 days
- 3) New population of non-adherent cells collected

Cells Harvested from HFBR & Placed in Flask



Harvest vs. Flask

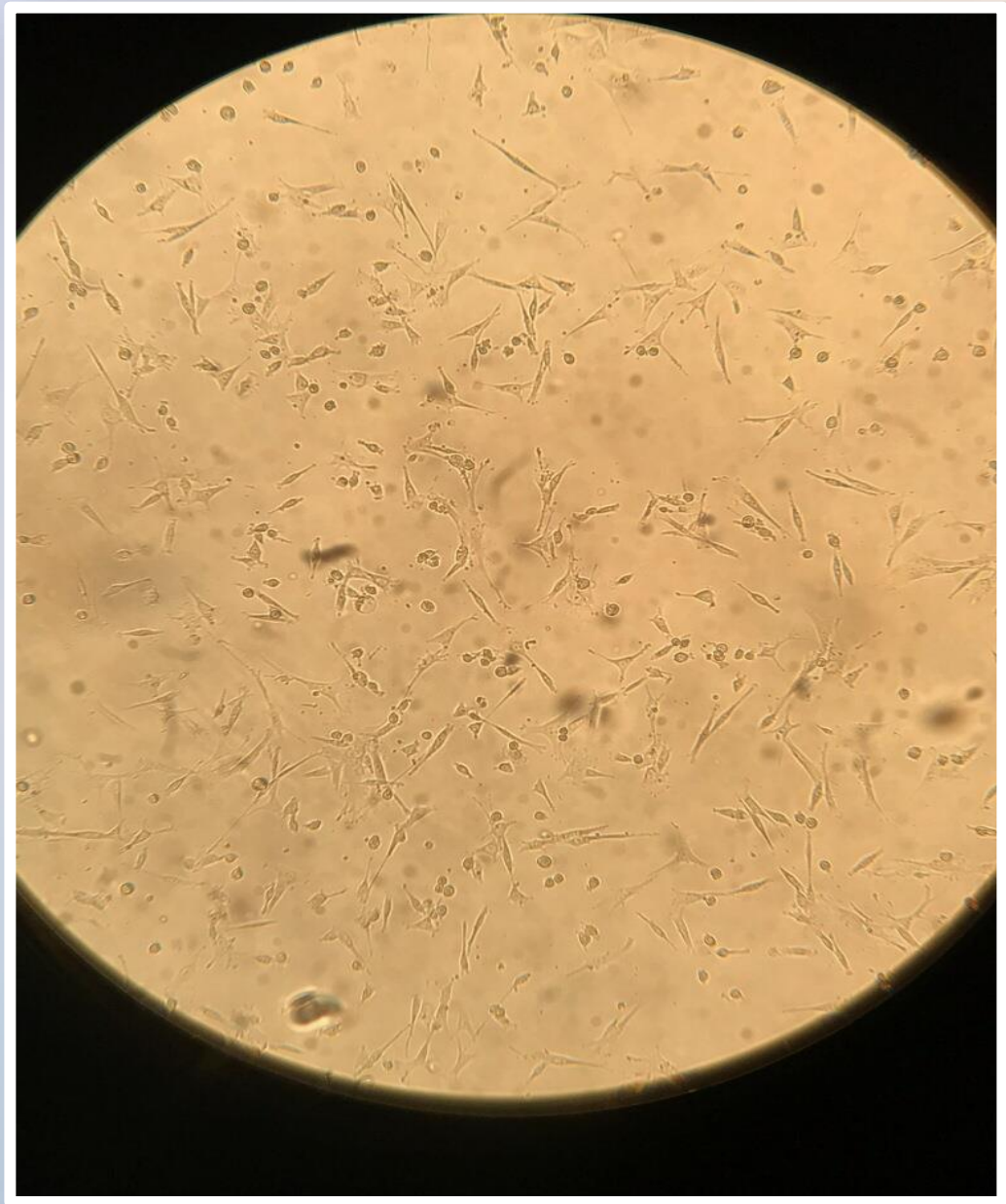
Phenotype	ECS Harvest	Flask
CD 45	4%	1%
CD 34	0%	0%
CD 133/2	2%	0%
CD 31	3%	48%
CD 13	6%	83%
CD 105	43%	99%
CD 73	18%	99%
CD 90	5%	96%
CD 14	23%	4%
NANOG	0%	0%

What is MSC?

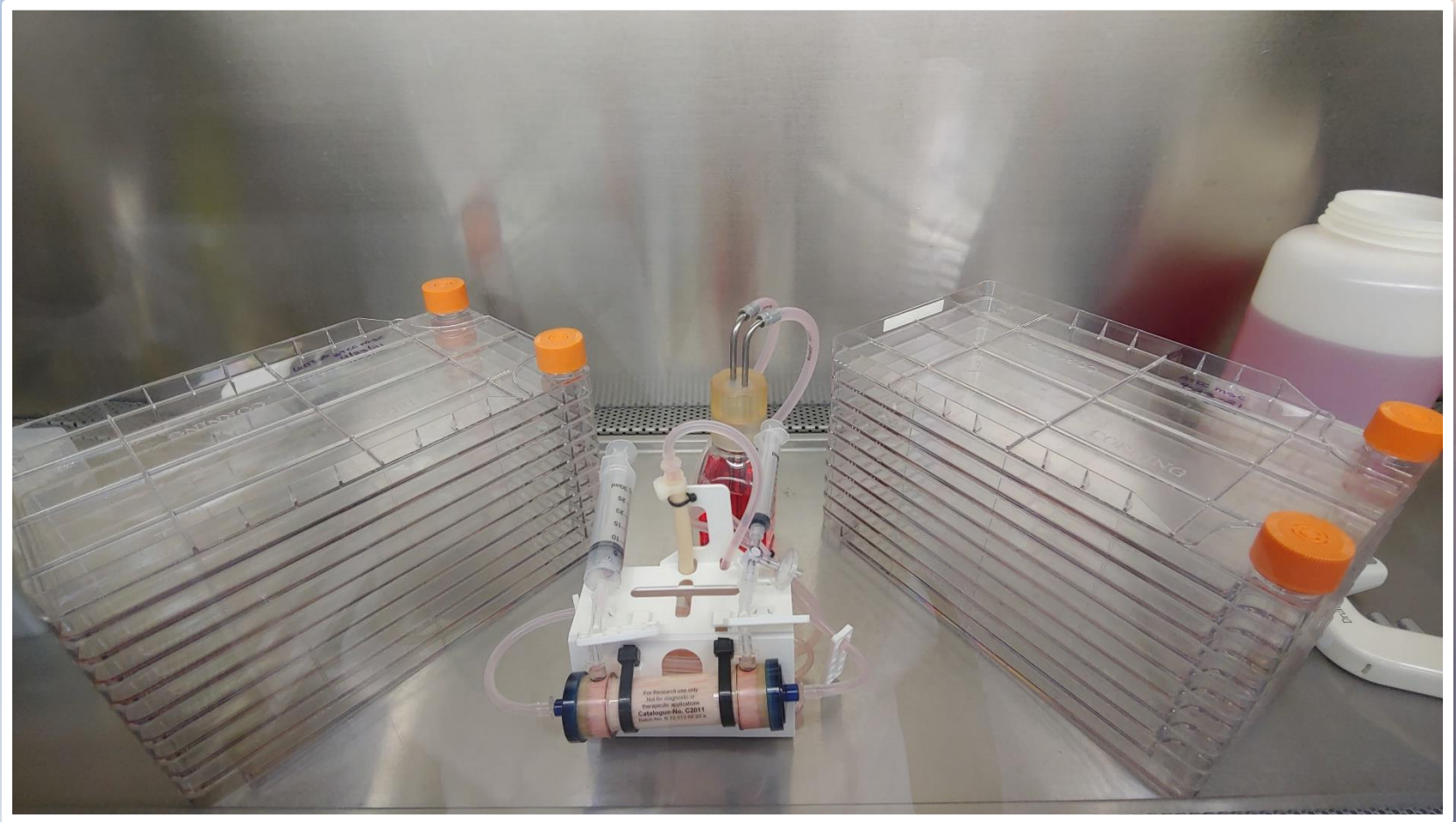
MSCs are plastic adherent fibroblastic cells with the “trilineage potential” of osteogenic, chondrogenic and adipogenic differentiation capabilities. Furthermore, they express the cell surface markers CD73, CD90, and CD105, and do not express haematopoietic and endothelial antigens (CD14 or CD11b, CD19 or CD79 α , CD34, CD45, HLA-DR)

Dominici M, Le Blanc K, Mueller I, et al. et al. Minimal criteria for defining multipotent mesenchymal stromal cells. The international society for cellular therapy position statement. *Cytotherapy*. 2006;8(4):315–317

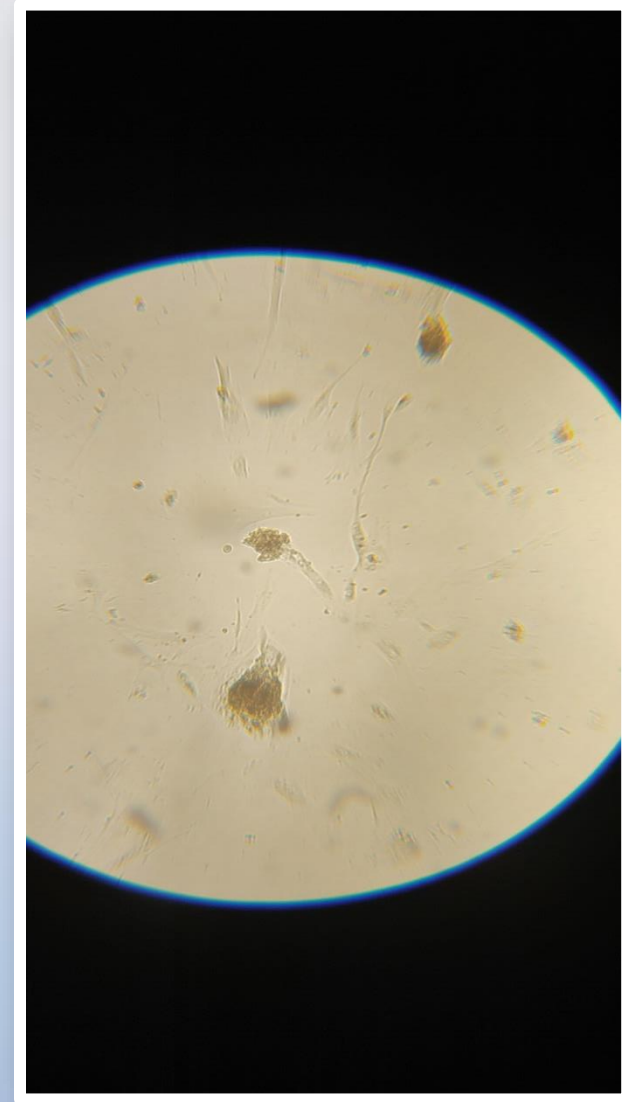
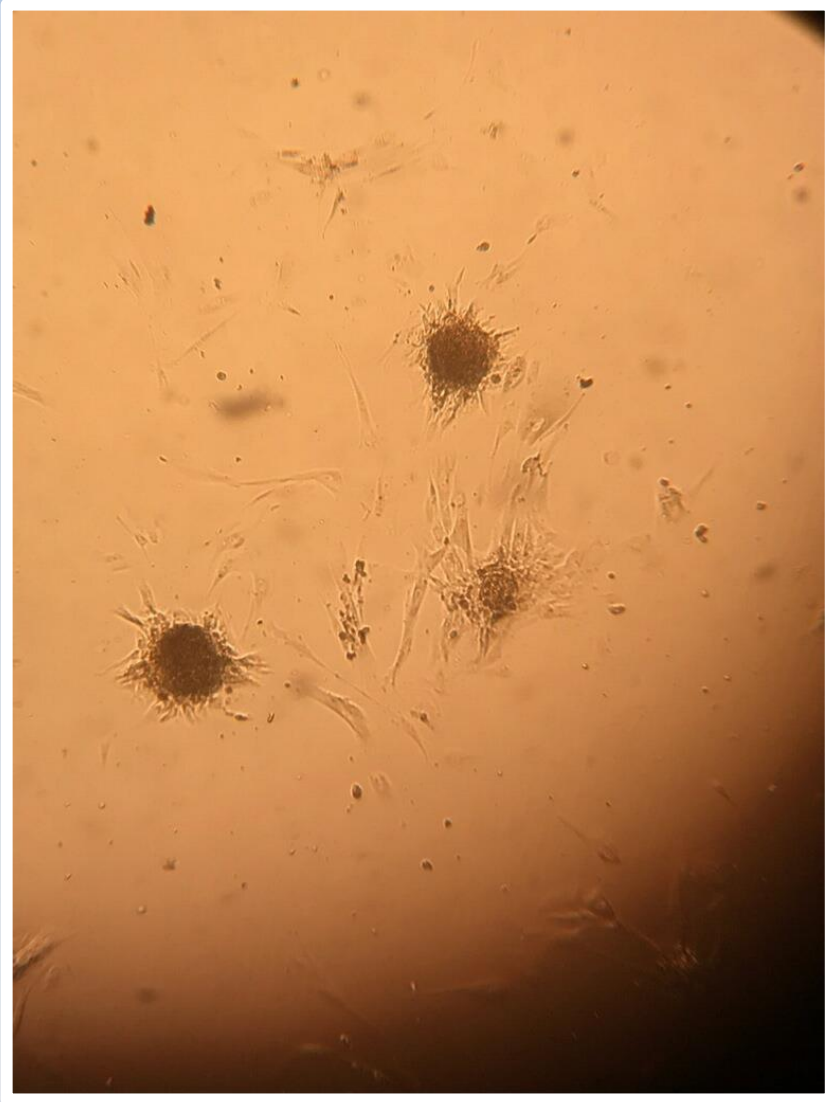
MSC in 2-D Culture



MSC in 3-D HFBR



MSC in 2-D After 3-D



Showing formation of Spheroids



Scalability of Production and Bio-Activity of Amniotic Fluid Stem Cell Extracellular Vesicles from 3-D Hollow Fiber Bioreactor and 2-D Culture.



FiberCell Systems
www.fibercellsystems.com/exosomes

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Introduction

EV clinical translation is constrained by limitations in scale-up of EVs production. Hollow fiber bioreactors (HFBR) support the culture of large numbers of cells, at high densities, producing significant numbers of EVs at high concentration. The high cell densities present in a HFBR can facilitate the use of xeno-free/chemically defined mediums, such as CDM-HD. Here we compare production, potency, identity and therapeutic potential of EVs collected from cells grown in culture dishes (2-D) vs. a HFBR (3-D).

Methods

Human clonal Amniotic Fluid Stem Cells, hAFSC, were derived from consented donor's amniotic fluid. 1×10^6 hAFSC were seeded in 2-D multi dishes (145 cm²) and 5.2×10^6 hAFSC were seeded into a 20 kD MWCO HFBR (FiberCell Systems C2011, 20 kD, 4,000 cm²) with fibronectin coating; both cultured in Chang's medium with 20% FBS. At confluence in the petri dish the medium was replaced with basal medium, stirred for 48 hr and EVs collected. After three days the medium in the ECS of the HFBR was replaced with Chang's medium alone, without 20% FBS, complete Chang's with 20% FBS remained in the central reservoir. The ECS was flushed with basal Chang's over the next 3 days and then harvesting of EVs every day was initiate. After two weeks of production serum in the reservoir was reduced stepwise to 5% and 5% CDM-HD introduced. After one more week serum was completely removed and replaced with 10% CDM-HD. The final weeks of EV production were produced using chemically defined medium, CDM-HD alone. Glucose consumption was monitored on a daily basis. 2-D EVs and 3-D EVs were compared by Nanosight, potency assay and by WB and therapeutic effect (in vivo injections in an animal model of chronic kidney disease, Aortt Syndrome).

Results

Control: 2-D EVs, Volume: 40ml, 3.0×10^{10} EV/ml, Total EV: 2×10^9

Figure 1: Daily Glucose Consumption - C2011 Hollow Fiber Bioreactor Culturing historical hAFSC
Day 17: 15% FBS
Day 19: 10% FBS
Day 21: 2.5% FBS, 2.5% CDM-HD
Day 22: 5% FBS, 5% CDM-HD
Day 26: 5% FBS, 10% CDM-HD
Day 34: 10% CDM-HD

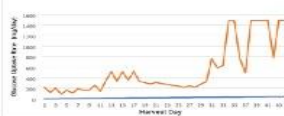


Figure 2: Daily EV Production - C2011 Hollow Fiber Bioreactor Culturing historical hAFSC

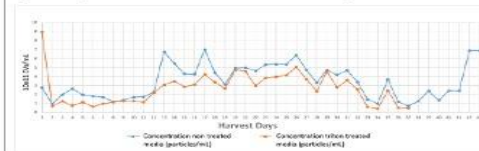


Figure 3 - hAFSC-EV size. Nanosight analysis EVs derived from 2-D (A) and 3-D (B) average size is 113 nm. Scale: Concentration not treated (white particles/ml); Concentration mice treated (red particles/ml).

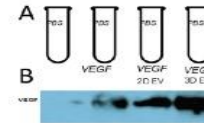
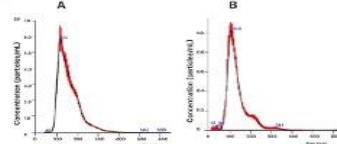


Figure 4: VEGF/VEGFR1 Co-immunoprecipitation. Immunoblot of VEGF (24 kDa, monomer), after co-IP with VEGFR1 in PBS+VEGF (100ng/ml) exposed-EVs from 2-D EVs (A) and 3-D EVs (B). Both EVs trap VEGF in a similar way. The weak band detected in the PBS/VEGF (second lane) represents a VEGF carryover due to incomplete removal of VEGF during washing.

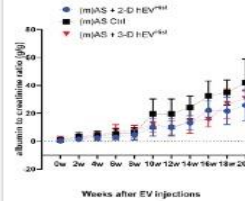
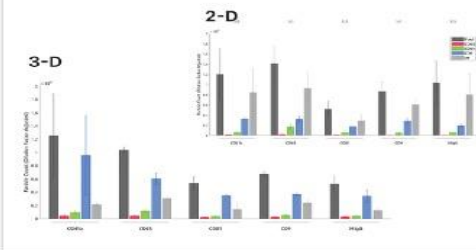


Figure 5: EVs ameliorate renal dysfunction. 2-D EVs and 3-D EVs, reduce proteinuria (measured by BSA) in treated mice vs untreated mice. AS injected with 3-D EVs (week 0) at injection and same collected every 2wks. WT and non-injected control mice were strain- and age-matched. *p<0.05** p<0.01. Mice are all in the study but the protective role of EVs is evident after injection.

Figure 6: 3-D hEVs present similar tetraspanin profile of 2-D hEVs as evaluated by ExoView



Discussion and Conclusion

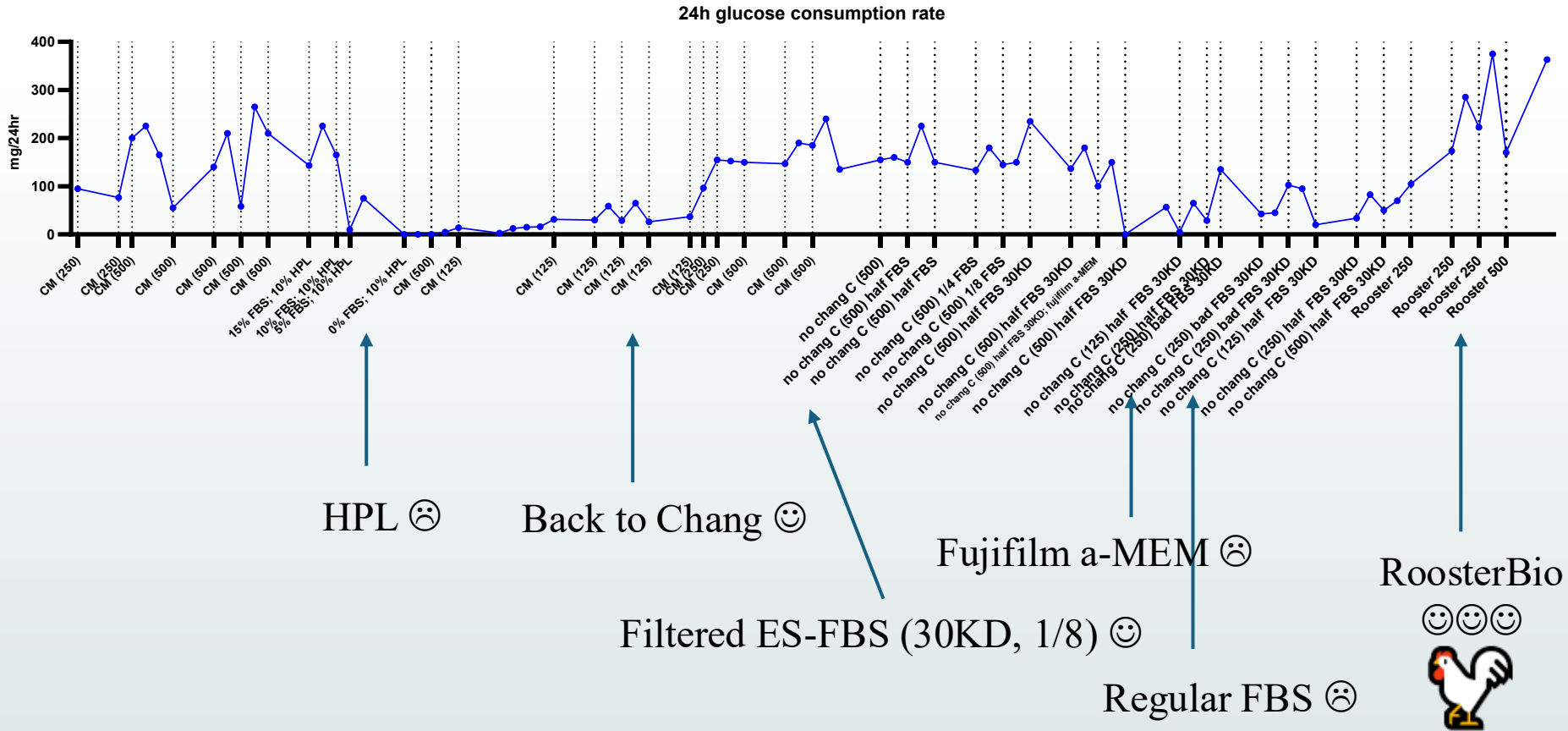
3-D EVs had comparable properties and bio-activity relative to 2-D EVs, but the HFBR produced 100x more concentrated EVs per mL. Each daily harvest produced more than 1×10^{10} EVs, 1×10^{10} would be an estimated human dose. The adaptation of these cells to a chemically defined medium and the demonstrated production range represents a significant step towards enabling therapeutic applications of hAFSC for treating kidney diseases in humans. The C2011 hollow fiber bioreactor module represents an additional 5x scale-up from the data presented here. The HFBR is a closed system that can be cGMP compliant. In conclusion, the HFBR can produce sufficient numbers of EV to support pre-clinical and clinical applications of EVs with at least similar properties to EVs produced by conventional 2-D methods.

Scaling up.

From
Impossible to
3,000 Doses a
Month.



110 Days Continuous Production of EVs from HAFSC



Data courtesy of Dr. Laura Perin and Paolo Neviani,
Children's Hospital, Los Angeles.



FiberCell Systems

www.fibercellsystems.com/exosomes



Retroviral Transduction and Production of Palm-GRET Labelled Extracellular Vesicles using Bone Marrow Derived MSC in a Hollow Fiber Bioreactor.

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2. Johns Hopkins Medical School, Baltimore, USA

Introduction

Various methods have been applied to the transfer of genes into mesenchymal stem cells (MSC). The generation of stable MSC transfectants is hampered by the limited number of passages MSC can undergo before they start to differentiate and difficulty in performing at clinical scale.

Methods

Bone marrow MSC from ATCC were expanded to 5X10^6 cells using DMEM/10% FBS and 10 T300 flasks. Retrovirus encoding for green fluorescent protein and Nanoluciferase protein was produced in culture.

Results

- 1. Isolated EVs from the first 16 days of harvest collection show strong Nuc signal and approximately 5-12% of all detected particles in those EV samples were GFP positive, indicating release of the GFP-Nanoluc fusion reporter proteins via EVs by MSC transfectants.
2. The particle counts of EVs produced by palmGRET MSC declined from 1.4X10^10 particles/mL in the first harvest to 3.9X10^9 particles/mL measured in the last harvest (day 27).
3. It was not really possible to directly determine transduction efficiency under these conditions.

Discussion

A hollow fiber bioreactor can reduce the volume required to perform transductions by 100X, and utilizes a closed, cGMP compatible format. Overall, these promising preliminary data warrant further optimization and refinement of the transduction protocol, particularly by modifying the viral titer, selection strategy, and length of the experiment.

Transient expression can be used with non-proliferating cells!

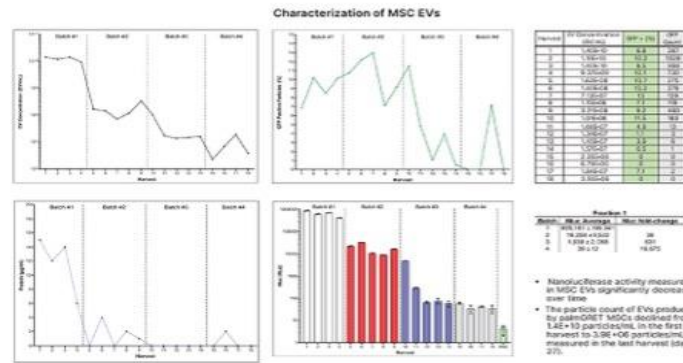
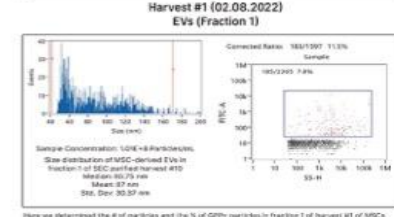
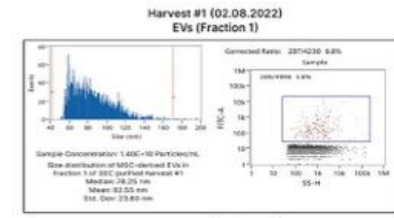


Table with 4 columns: Harvest, EV Concentration, GFP+ % of EVs, GFP+ % of MSCs. Rows 1-16 showing data trends.

Table with 3 columns: Harvest, EV Concentration, GFP+ % of EVs. Rows 1-16 showing data trends.



Here we determined the # of particles and the % of GFP+ particles in fraction 1 of harvest #1 of MSCs using Nanoluc.

**What is it?
See the Unseen!**



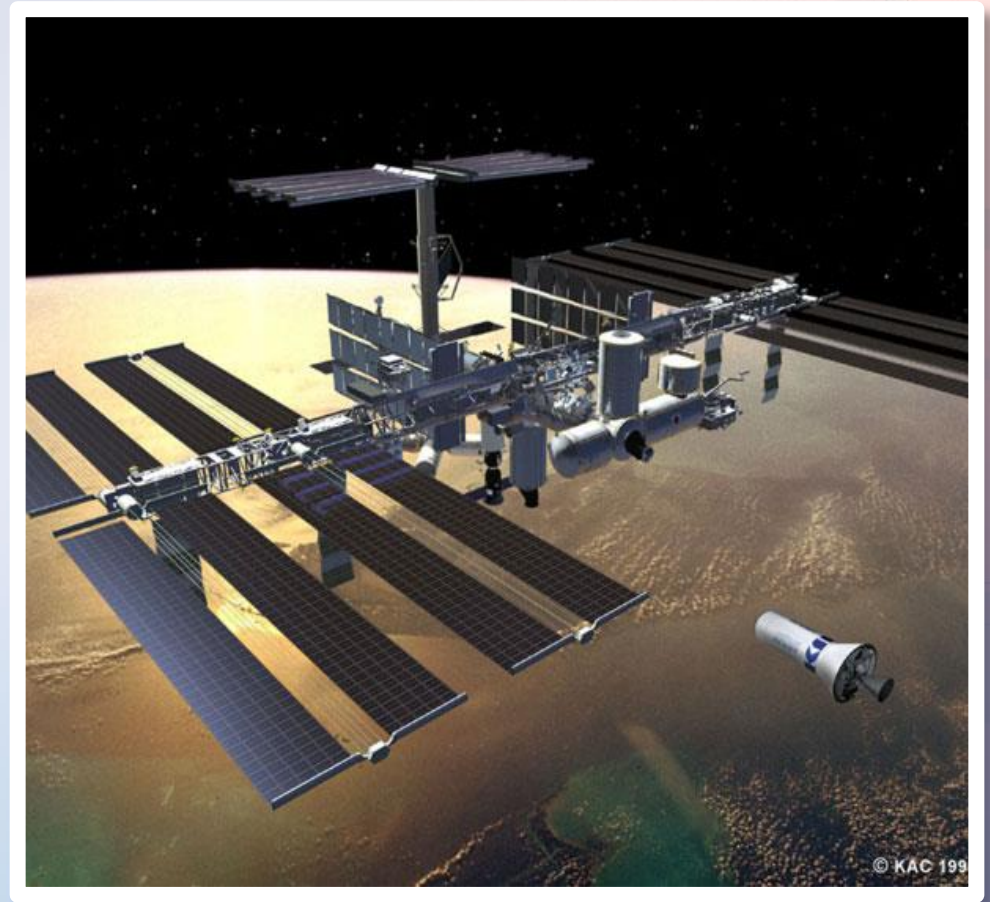
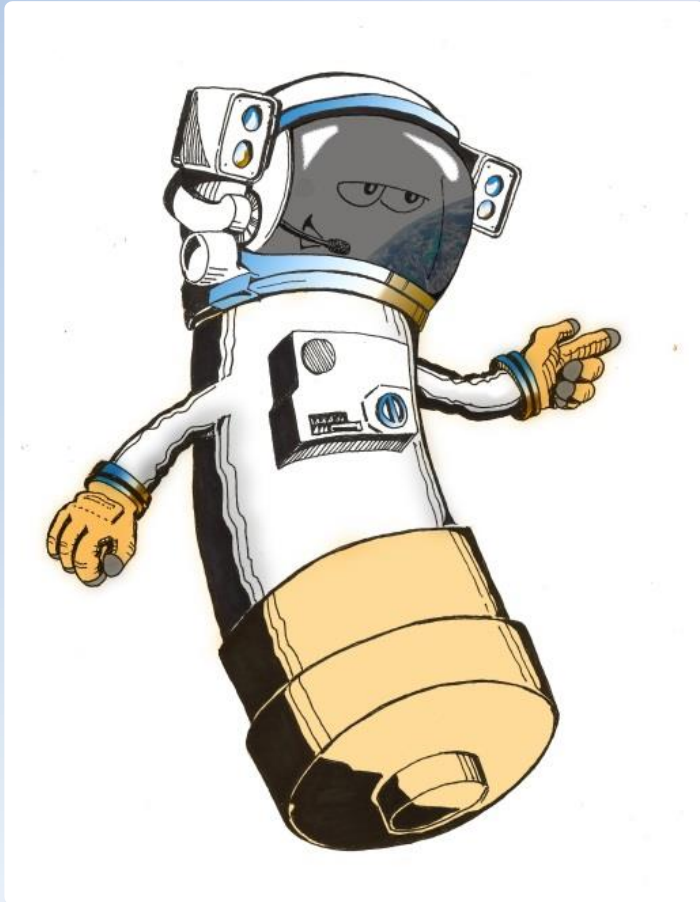
Summary

- The most *in vivo* method for culturing cells over long periods of time. Media and cells work together to generate their specific microenvironment.
- Hollow fiber was 3-D before the importance of 3-D culture was recognized
- Can be the only way to get two different cell types, in close enough proximity, at high enough density, for long enough time to observe interactions between the cells.
- 4-D culture, enough time for cells to self-organize, structures to form, and for cell-to-cell interactions to develop.

Summary

- Hollow fiber bioreactors are the method of choice for the culture of 10^9 to 10^{11} cells
- Can produce gram quantities of exosomes
- Concentration of 100x higher than with conventional methods
- The most *in vivo* method for culturing cells over long periods of time
- Suitable for cGMP production
- Permits use of FBS without endogenous EV contamination
- Enhanced bioactivity
- Saves time, space, purification costs

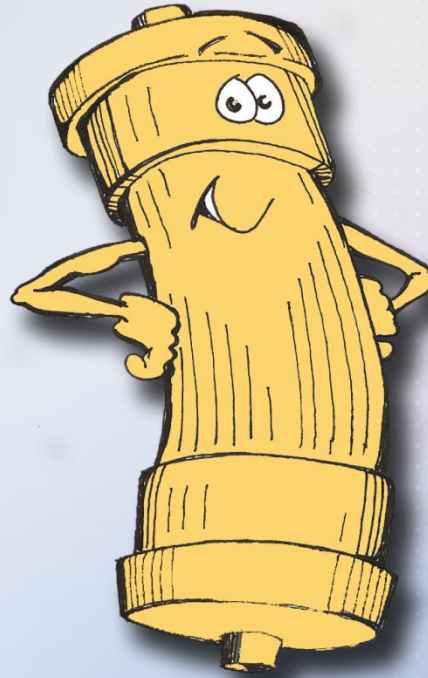
FiberCell Systems HFBR in Space



"...The evolution of cell culture"



Thank you.



Applications

