#### **UC Irvine**

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#### **Title**

CD Microfluidic Device for Capture and Isolation of Cancer Stem Cells

#### **Permalink**

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#### **Authors**

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#### **Publication Date**

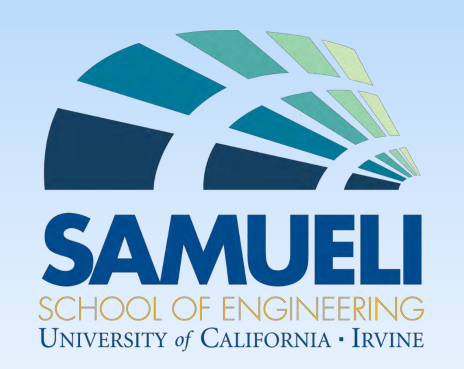
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# CD Microfluidic Device for Capture and Isolation of Cancer Stem Cells

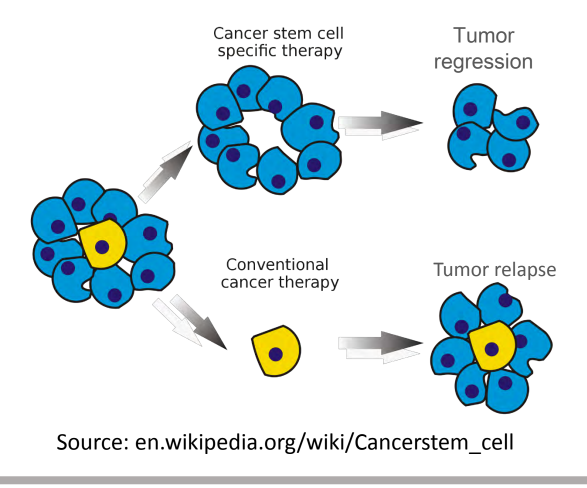
Lydia Ameri, Mechanical Engineering and Materials Science Engineering Gema Vera, Biomedical Engineering

Faculty Advisors: Dr. William Tang, Biomedical Engineering Department & Dr. Lawrence Kulinsky, Department of Mechanical and Aerospace Engineering



### Background

Cancer stem cells (CSC) are known to be the main culprits in the promotion and formation of tumors throughout an individual's body. Though they compose approximately 3% of cancer cells, their ability to differentiate into various cell types, self-renewal and divide indefinitely prevents the successful elimination of cancer. This devices aims to help researchers track CSC's in the body in order to successfully remove cancer. More research can also be done to study CSC's specially to understand them better and come up with treatments to destroy them.



### **Project Goals**

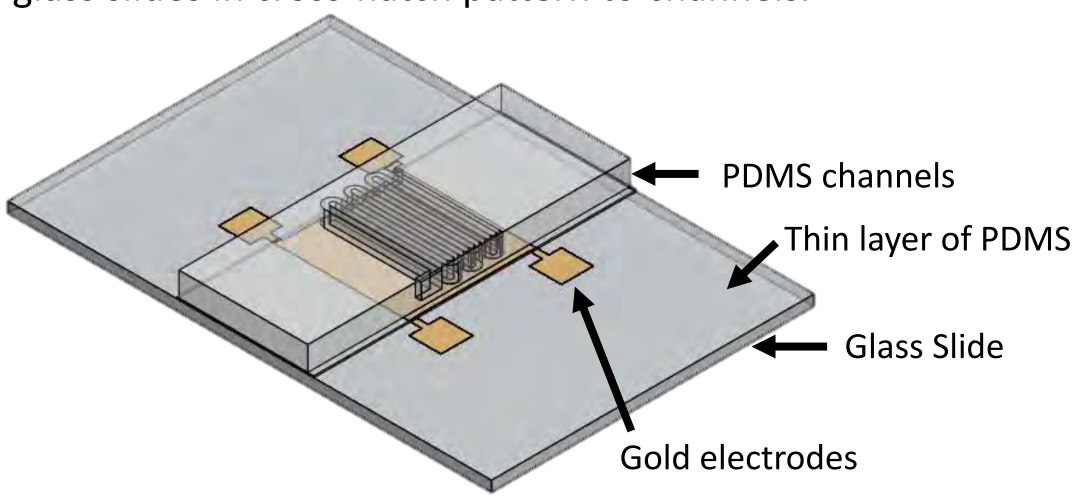
- Isolate majority of Cancer Stem Cells from non-tumorigenic cells using a process that that will separate the cells by creating a non-uniform electric field, called Dielectrophoresis (DEP).
- Separate cells based on temporal separation.
- Determine which design would provide for the best results in terms of cell separation.

### **Project Design**

PDMS will be the primary material used to create the fluid channels. If leakage occurs, then the second method, using CNC machined plastic, will be used. The device will be spun on a CD to separate the cells based on density and will be captured based on temporal separation.

## Polydimethylsiloxane (PDMS) Channels

A PDMS mold will be created for the channels. The glass slide and PDMS structure will be plasma bonded together to prevent leakages. Gold electrodes will be patterned on the glass slides in cross-hatch pattern to channels.

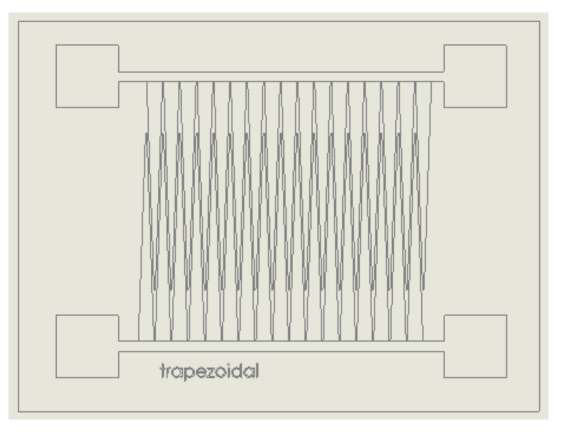


**CNC Machined Plastic** 

The channels will be created on the glass slide and a Computer Numeric Controlled (CNC) machined plastic for the channels. It will be attached to the glass using a pressure sensitive adhesive.

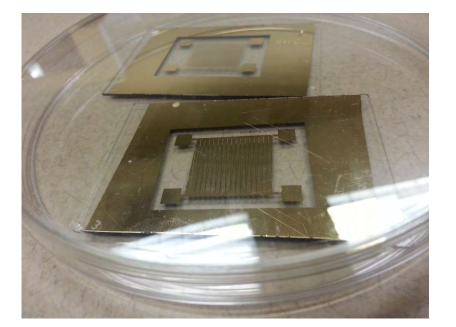
### **Electrodes**

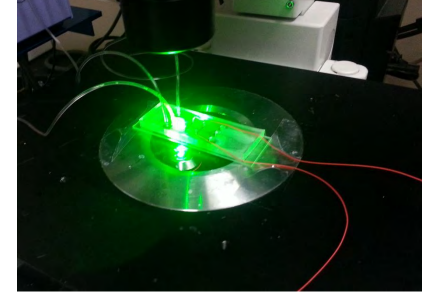
The electrodes will be created in trapezoidal or rectangular interdigitated arrays to determine which shape would provide for the strongest electric field.



### **Progress & Current Status**

- Created masks for rectangular and trapezoidal shaped electrodes.
- Observed DEP on test electrodes using polystyrene beads.
- Tested plasma bonding between two layers of PDMS. Fluid was able to flow through channels without leaking.





Gold electrodes on glass slides

DEP on sample electrode slide.

Test to determine effectiveness of plasma bonding 2 layers of PDMS. Green dyed water used to observe flow through the channels.

### **Spring Quarter Timeline**

Weeks 1-2: Create and test PDMS prototype

Weeks 2-3: Create and test CNC plastic prototype

Weeks 4-6: Make revisions to design as necessary

Week 6: Test prototypes with blood sample

Week 7: Present at UROP symposium

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