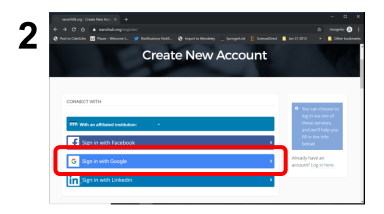
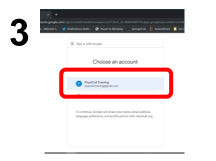
#### nanoHUB Account

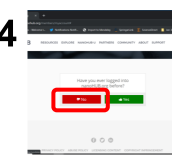
- These tutorials use cloud-hosted PhysiCell models on nanoHUB.org.
- nanoHUB is free, but it requires a onetime registration.

#### • Steps:

- Visit <a href="https://nanohub.org/register">https://nanohub.org/register</a>
- Choose "Sign in with Google"
- 3. Choose a Google account
- Click "No" (so it doesn't try to associate with some other nanoHUB account)
- 5. Finish filling in details, and you're done!
- 6. Use your google account for future sign-ins.







# A first course in PhysiCell

Get the lecture and codes here!



Paul Macklin, Ph.D.

Intelligent Systems Engineering Indiana University

August 21, 2019

MathCancer.org

#### **Thanks: Partners**

#### Colon cancer metabolism:

Stacey Finley (USC, U01 Contact PI)

#### Colon cancer organoids:

Shannon Mumenthaler (USC, U01 PI)

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- Gary An (University of Vermont)

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- alumni: S.H. Friedman (OKSI), A. Ghaffarizadeh (USC)

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- Yafei Wang (liver mets, nanotherapy)
- Furkan Kurtoglu (multicellular metabolism, hypoxia)
- Aneequa Sundus (cyanobacteria, synthetic multicellular systems)

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- Metastasis: B. Fischer, D. Murphy, K. Konstantinopoulos, B. Duggan
- Nanotherapy: T. Mahjan
- Jupyter GUIs: E. Bower, D. Mishler, T. Zhang
- PhysiCell technologies: E. Freeman, A. Connor

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  - projects with PIs Agus, Gilkes, Peyton, Ewald, Newton, Bader









### Cancer is a systems problem

#### Interconnected systems and processes:

- Single-cell behaviors
- Cell-cell communication
- Physics-imposed constraints (e.g., diffusion)
- Systems of systems (e.g., immune system)

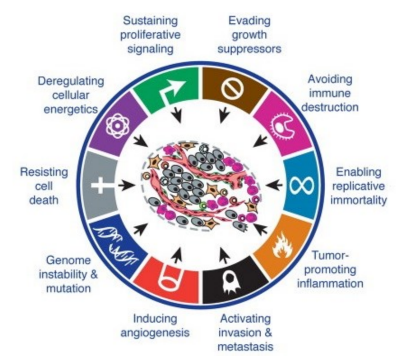
In cancer, these systems become dysregulated.

Treatments target *parts* of these systems.

Cancer is a **complex system**: changing one part can have **surprising effects**!

Modeling can help **understand** this system. This is **multicellular systems biology**.

If we can **control** these systems, we've arrived at **multicellular systems engineering**.



Source: Hanahan & Weinberg (2011)

DOI: 10.1016/j.cell.2011.02.013

## Design target for multicellular modeling

- Model multiple diffusing chemical factors
  - Growth substrates and metabolites
  - Signaling factors
  - Drugs
- Model many cells in these chemical environments
  - Environment-dependent behavior
  - Mechanical interactions
  - Heterogeneity:
    - ♦ individual states
    - individual parameter values
    - ♦ individual model rules
- Run many copies of the model in high throughput

## BioFVM: Simulating 3-D biotransport

<u>Design goal:</u> Simulate multiple diffusing substrates in 3D with desktops or single HTC/HPC nodes

Typical use: pO<sub>2</sub>, glucose, metabolic waste, signaling factors, and a drug, on 10 mm<sup>3</sup> at 20 μm resolution

#### **Features:**

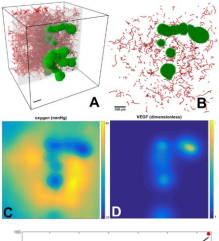
- Off-lattice cell secretion and uptake
- 2<sup>nd</sup>-order accurate (space), 1<sup>st</sup>-order accurate (time), numerically stable

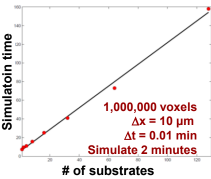
#### **Method:**

- Operator splitting, LOD, customized Thomas solvers, etc.
- Standard C++11, cross-platform
- OpenMP parallelization
- O(n) cost scaling in # substrates, # voxels
- Easy to simulate 5-10 substrates on 10<sup>6</sup> voxels

Reference: Ghaffarizadeh et al., Bioinformatics (2016)

DOI: 10.1093/bioinformatics/btv730





### PhysiCell: A multicellular framework

<u>Design goal</u>: Simulate 10<sup>6</sup> or more cells in 2D or 3D on desktops or single HPC nodes

#### Features:

- · Off-lattice cell positions
- Mechanics-based cell movement
- Cell processes (cycling, motility, ...)
- Signal-dependent phenotype
- Can dynamically attach custom data and functions on a cell-by-cell basis
- Deployed from Raspberry Pi to Crays

#### Method:

- Standard C++11, cross-platform
- OpenMP parallelization
- O(n) cost scaling in # cells

Reference: Ghaffarizadeh et al.,

PLoS Comput. Biol. (2018)

**DOI**: <u>10.1371/journal.pcbi.1005991</u>



Try this model yourself!

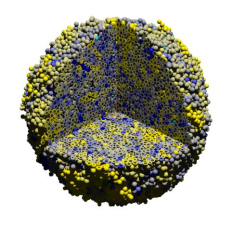
Computational Biology

Research Prize for

Public Impact

nanohub.org/tools/pc4heterogen

Current time: 7 days, 0 hours, and 0.00 minutes 53916 cells



Competition in a 3-D tumor

[View on YouTube (8K)]

MathCancer.org

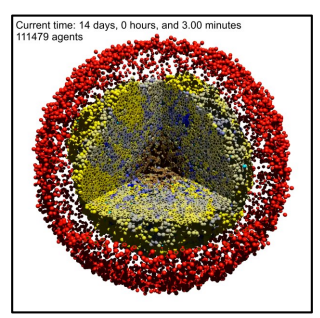
## Some examples

# Tumor-parenchyma mechanical feedbacks



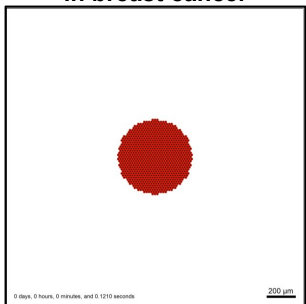
Y. Wang (IU), with Mumemthaler (USC), Sparks (Miami U), Frieboes (U Louisville)

#### Cancer immunotherapy



with G. An (U. Vermont), Ozik, Wozniak, and Collier (Argonne National Lab)

# Phenotypic persistence in breast cancer



with D. Gilkes (Johns Hopkins U.)

## Some key extensions

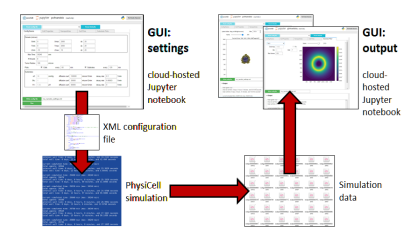
- Incorporated Boolean signaling networks by combining MaBoSS with PhysiCell to create *PhysiBoSS* (Letort et al., *Bioinformatics* 2019)
  - DOI: <u>10.1093/bioinformatics/bty766</u>.
- Performed high-throughput model investigations on supercomputers with Argonne National Lab (Ozik et al., BMC Bioinformatics 2018)
  - **DOI**: 10.1186/s12859-018-2510-x
- Extended work to use machine learning to accelerate parameter investigation and interpret model results. (Ozik et al., Molec. Syst. Des. Eng. 2019)
  - **DOI**: 10.1039/c9me00036d.
- Automated creation of Jupyter-based GUIs and cloud-hosted PhysiCell-powered models (Heiland et al., J. Open Source Soft. 2019)
  - DOI: 10.21105/joss.01408

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### Let's try one!

#### cancer biorobots:

- green:
  - ♦ cycle entry scales with O2
  - O2 depletion causes necrosis
  - cumulative drug exposure causes apoptosis
- blue:
  - drug-loaded "cargo"
- red:
  - worker cells that seek and haul cargo towards hypoxic zones



XML+Jupyter architecture

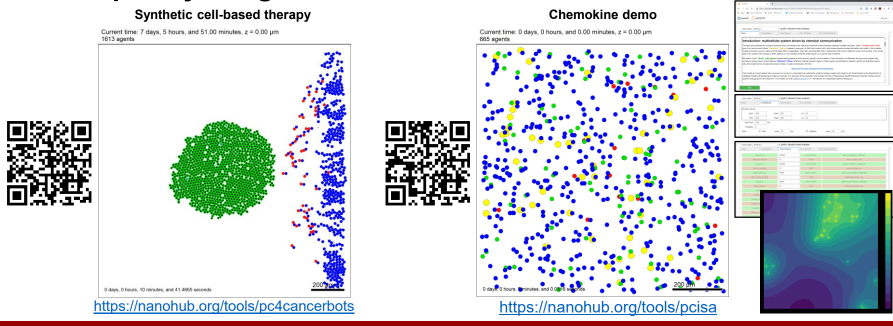


Try this model yourself!

https://nanohub.org/tools/pc4cancerbots

#### Use case: Research in the classroom

 Because it's easy to convert a research-grade model to a graphical app, we can quickly integrate new research into the classroom.



## Let's download PhysiCell

- Two download options to get the latest numbered release:
  - GitHub:
    - https://github.com/MathCancer/PhysiCell/releases/latest



- SourceForge:
  - https://sourceforge.net/projects/physicell/files/latest/download
- Unzip the download, and enter the PhysiCell root directory



Of particular note, go to ./documentation and open the User\_Guide.pdf

#### A note about time steps

 PhysiCell is designed to account for the multiple time scales inherent to these problems, and has 3 time scales:

| • | $\Delta t_{ m diffusion}$ | diffusion, secretion, and uptake | (default: 0.01 min) |
|---|---------------------------|----------------------------------|---------------------|
| • | $\Delta t_{ m mechanics}$ | cell movement                    | (default: 0.1 min)  |

•  $\Delta t_{\text{cell}}$  phenotype and volume changes (default: 6 min)

 This allows some efficiency improvements: not all functions need to be evaluated at each time step.

• See the PhysiCell method paper. (Oddly, not in the User Guide (yet).)

### PhysiCell agents

- Each PhysiCell agent has:
  - functions to sample the microenvironment
  - State elements (things that are measured from an external frame of reference)
    - ♦ ID, type, position, velocity, base-to-apex orientation, ...
  - Phenotype
    - ♦ hierarchically-ordered behavioral state and parameters (much more detail soon)
  - user-defined functions
    - functions that act on the cell's phenotype (or state)
    - ♦ can replace the built-in mechanics and other behaviors
  - user-defined custom data
    - ♦ used to track additional details specific to user models, if they aren't already

Documentation: User Guide, Sec. 9

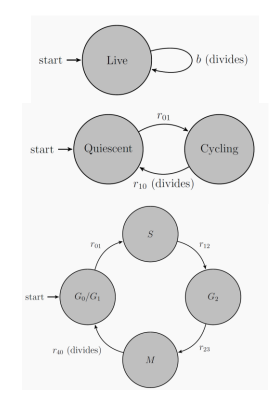
### **Cell phenotype**

- One of the most critical data elements in a PhysiCell Cell is phenotype
- Hierarchically organize key behavioral elements:
  - Phenotype
    - ◆ cycle: advancement through a cell cycle model
    - ♦ death: one or more types of cell death
    - ♦ volume: cell's volume regulation
    - ♦ geometry: cell's radius and surface area
    - ♦ mechanics: adhesion and resistance to deformation ("repulsion")
    - ◆ motility: active motion (other than "passive" mechanics)
    - ◆ secretion: both release and uptake of chemical substrates. Interfaces with BioFVM
    - ♦ molecular: a place to store internalized

**Documentation:** User Guide, Sec. 10

### **Phenotype: Cycle**

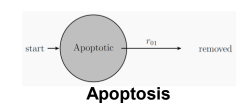
- Each agent's **phenotype** had a **cycle** with:
  - Cycle model
    - ♦ A directional graph: *nodes* are cycle **phases**  $\{P_i\}$  and *edges* are **transition rates**  $\{r_{ij}\}$
    - $r_{ij}$  is the transition rate from phase  $P_i$  to phase  $P_i$
    - One of the transitions must be marked as a division transition
    - Users can attach arrest condition functions to these transitions (e.g., size checks)
  - Cycle data
    - stores the cell's current transition rates
- Documentation: User Guide, Sec. 11.1

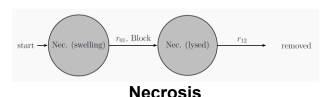


### **Phenotype: Death**

- Death has one or more death models:
  - A specialized cycle model with a removal transition rate
  - Extra parameters to help govern cell volume
  - Each death model has an associate death rate
  - Also stores an easy Boolean dead to easily check if the cell is alive.
- PhysiCell has built-in apoptosis and necrosis death models

**Documentation:** User Guide, Sec. 11.2





## Phenotype: Volume

- volume records the cell's sub-volumes:
  - nuclear and cytoplasmic
  - solid vs. fluid
  - calcified fraction
  - key parameters
- a very simple default model to regulate volume based on ODEs
  - Change the parameters, target values based on environment and cell state

$$\frac{dV_F}{dt} = r_F (V_F^* - V_F)$$

$$\frac{dV_{NS}}{dt} = r_N (V_{NS}^* - V_{NS})$$

$$\frac{dV_{CS}}{dt} = r_C (V_{CS}^* - V_{CS})$$

$$\frac{dV_{NS}}{dt} = r_N (V_{NS}^* - V_{NS})$$

$$\frac{dV_{CS}}{dt} = r_C (V_{CS}^* - V_{CS})$$

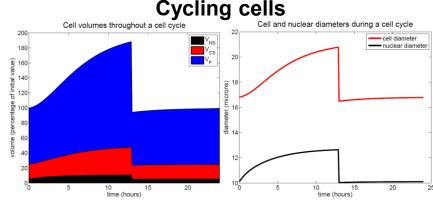
**Documentation:** User Guide 11.3

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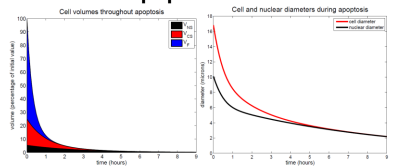
### **Phenotype: Geometry**

- geometry records:
  - surface area (not actively tracked)
  - equivalent spherical radius
  - equivalent nuclear radius

**Documentation:** User Guide 11.4



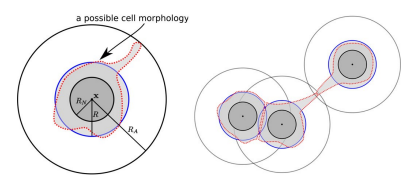


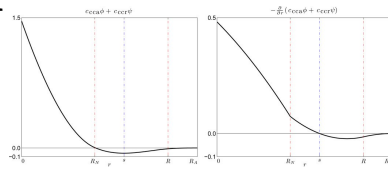


### **Phenotype: Mechanics**

- Mechanics keeps parameters for adhesion and "repulsion"
  - Key parameter: maximum adhesion distance
    - ♦ a multiple of the cell's radius
  - (as a multiple of the cell's radius)
- Default model uses potential functions, but this can be supplemented or replaced.

**Documentation:** User Guide 11.5

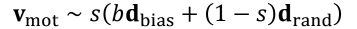


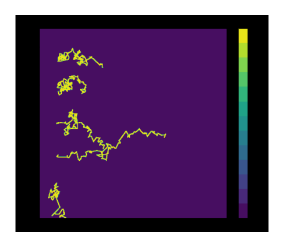


## **Phenotype: Motility**

- Motility controls biased random migration
  - Migration speed s
  - Bias direction d<sub>bias</sub>
  - Migration bias  $0 \le b \le 1$ 
    - If b = 1, deterministic motion
    - If b = 0, purely Brownian motion
  - Persistence time  $T_{per}$

**Documentation:** User Guide Sec. 11.6







Try this model yourself! (2D)

https://nanohub.org/tools/motility

## **Phenotype: Secretion**

 Secretion stores parameters for secretion and uptake of diffusing substrates

$$\frac{\partial \boldsymbol{\rho}}{\partial t} = \nabla \cdot (\boldsymbol{D} \nabla \boldsymbol{\rho}) - \boldsymbol{\lambda} \cdot \boldsymbol{\rho} + \sum_{i} \delta(\boldsymbol{x} - \boldsymbol{x}_{i}) V_{i} (\boldsymbol{S}_{i} \cdot (\boldsymbol{\rho}_{i}^{*} - \boldsymbol{\rho}) - \boldsymbol{U}_{i} \cdot \boldsymbol{\rho})$$

**Future**: Will add an additional "Export" term since the **S** form isn't right for all types of measurements and assays.

**Documentation:** User Guide Sec. 11.7

### Phenotype: Molecular

- Molecular is where total internalized substrates are tracked. (optional)
  - A fraction (or all) of substrates can be released at cell death
  - A fraction (or all) of substrates can be transferred when a cell is ingested
  - Internalized substrates are divided among daughter cells
- Eventual support for molecular-scale models will be attached here.

**Documentation:** User Guide Sec. 11.8

## Phenotype-centric programming

- The core cell behaviors are implemented:
  - Cell cycling (with user-selectable models)
  - Cell death
  - Cell adhesion / repulsion
  - Cell motility
  - Cell secretion / uptake
- Modelers can focus on writing functions that control these behaviors.
- This is phenotype-centric programming.

## Functions in PhysiCell

Almost all functions in PhysiCell have this form:

```
void my_function( Cell* pCell, Phenotype& phenotype, double dt );
```

#### All cells have the following key functions (in pCell->functions):

- volume update function
- update\_migration\_bias
- custom\_cell\_rule (default NULL, evaluated at each mechanics time step)
- update velocity
- set orientation

**Documentation:** User Guide Sections 9.4.3, 19.1

#### **Cell Definitions**

- A Cell Definition is a convenient way to set the parameters and functions for a whole class of cells
  - Users can instantiate cells of a specific type using create\_cell()
  - With no argument, new cells use the cell\_defaults definition
- Best practice: set up the cell\_defaults definition with the correct custom data and functions, then copy this to make new cell definitions
- <u>Tip</u>: Refer back to the phenotype in your agent's cell definition as a reference parameter set (i.e., to get the initial parameter values)

**Documentation:** User Guide Sec. 9.4.5

# Okay, let's try some models

### Sample projects

- It would be inefficient (insane) to code new projects from scratch.
- So, we provide sample projects:
  - 2D and 3D template projecgts
  - Sample cancer models
- make [project-name]: populate a sample project
  - Then use **make** to compile it
- make data-cleanup: clean up the output data
- make reset: return to a "clean slate" (depopulate the project)

**Documentation:** User Guide Sections 6, 7.

### Typical model implementation

- Plan!
- Create a new project (start with 2D or 3D template)
- Set up the XML ( ./config/PhysiCell\_settings.xml ).
  - Set up the chemical microenvironment
  - Set up user-defined parameters
- Define cells (create\_cell\_types())
  - Add custom data to the default cell definition
  - Create any needed custom function (for model rules)
  - Create and configure cell definitions
- Place cells ( setup\_tissue() )
  - Place different cell types in their initial positions
- Compile and run

### **Our first project**

#### Populate the biorobots sample:

make biorobots-sample

#### Compile the project:

make

#### run the project:

./biorobots

#### cleanup:

make data-cleanup
make reset.

### **Our Project**

- Modify heterogeneity sample:
  - Dimensionless oxygen, glucose, and metabolic waste
  - Energy-dependent cycle entry
  - Energy-dependent necrosis
  - Waste-dependent apoptosis
  - Cells have an initial distribution of parameters
- Maybe later, if we have time:
  - A new cell type (interloper) that;
    - ♦ use the "glycolytic" energy, secrete waste, resist waste-based death
    - ♦ chemotax towards low oxygen

### Plan

### **Example: tumor competition**

Cells can use two types of "metabolism"

$$\frac{dE}{dt} = \alpha\sigma g + \beta g - uE$$

• Cells consume supplies and make waste

$$U_{\sigma} = 10\alpha$$
 $U_{g} = 0.2(\alpha + \beta)$ 
 $S_{w} = 10r$ 

All cell behaviors consume energy:

$$u = \mu(\alpha + \frac{2}{3}\beta + \frac{2}{3}r)$$

 Let's give each cell independent "gene expressions" for α, β, r Suppose cell cycling increases with energy:

$$b = \begin{cases} 0 & E < 0.1 \\ b_M \frac{E - 0.1}{0.9 - 0.1} & 0.1 < E < 0.9 \\ b_M & 0.9 < E \end{cases}$$

Necrotic death increases below E = 0.1

$$d_N = d_{nec} \left( \frac{E - 0.1}{0.1} \right)^+$$

- Apoptotic death increases with waste:
  - But *r* gives tolerance (resistance) to the waste

$$d_A = d_{apop} (1 + 9w(1 - r))$$

#### What we'll need

- Modify the microenvironment
  - oxygen, glucose, waste
- Change the custom variables in cell defaults
  - alpha, beta, resistance, energy, use rate
  - Let's leave all the birth and death rates relatively untouched.
- A new phenotype function
  - void energy\_based\_cell\_phenotype( Cell\* , Phenotype& , dt )
- Modify the cell defaults with:
  - Custom data and the new phenotype function.
- Modify the setup\_tissue() to choose random values of alpha, beta, resistance for each cell.
- New coloring function (to tailor the SVGs)
  - energy coloring function( Cell\* )

# Create a new project

### Modify an existing project

```
make data-cleanup
```

make reset

make heterogeneity-sample

# Set up the XML

### Let's modify the microenvironment

- Open ./config/PhysiCell\_settings.xml
- Browse to **microenvironment\_setup** block
- Modify the first variable
- Now, copy it to make glucose and waste

**Documentation:** User Guide Sec. 13.4

**Tutorial:** 

http://www.mathcancer.org/blog/setting-up-the-physicell-microenvironment-with-xml/

### **Modifying the microenvironment (1)**

Modify oxygen to be dimensionless, set initial and boundary values

### **Modifying the microenvironment (2)**

Copy oxygen to make glucose and waste.

```
<variable name="glucose" units="dimensionless" ID="1">
  <physical parameter set>
     <diffusion coefficient units="micron^2/min">18000</diffusion coefficient>
     <decay rate units="1/min">0</decay rate>
  </physical parameter set>
  <initial condition units="mmHg">1</initial condition>
  <Dirichlet boundary condition units="mmHg" enabled="true">1</Dirichlet boundary condition>
</variable>
<variable name="waste" units="dimensionless" ID="2">
  <physical parameter set>
     <diffusion coefficient units="micron^2/min">100000.00</diffusion coefficient>
     <decay rate units="1/min">0</decay rate>
  </physical parameter set>
  <initial condition units="mmHg">0</initial condition>
  <Dirichlet boundary condition units="mmHg" enabled="true">0</Dirichlet boundary condition>
</variable>
```

### Modifying the microenvironment (3)



#### Checkpoint:

To catch up to the code at this point, unzip <a href="mailto:checkpoint1.zip">checkpoint1.zip</a>

MathCancer.org

### **Define cells**

### Let's add new custom variables

in create\_cell\_types() in heterogeneity.cpp (This is usually ./custom\_modules/custom.cpp in most projects.)

```
// add custom data

cell_defaults.custom_data.add_variable( "alpha" , "dimensionless", 1.0 );

cell_defaults.custom_data.add_variable( "beta" , "dimensionless", 1.0 );

cell_defaults.custom_data.add_variable( "resistance" , "dimensionless", 1.0 );

cell_defaults.custom_data.add_variable( "use_rate" , "dimensionless", 1.0 );

cell_defaults.custom_data.add_variable( "energy" , "dimensionless", 1.0 );
```

### Let's make a new phenotype function (1)

```
First, declare the new function in heterogeneity.h
void energy based cell phenotype (Cell* pCell, Phenotype phenotype, double dt);
Next, let's implement. In the function in heterogeneity.cpp
void energy based cell phenotype ( Čell* pCell, Phenotype& phenotype, double dt )
   // housekeeping: one-time searches for variables
       // for finding the right cycle phases
   static int cycle start index = live.find phase index( PhysiCell constants::live );
   static int cycle end index = live.find phase index( PhysiCell constants::live );
       // for finding the death model indices
   static int apoptosis i =
       cell defaults.phenotype.death.find death model index( PhysiCell constants::apoptosis death model );
   static int necrosis i =
       cell defaults.phenotype.death.find death model index( PhysiCell constants::necrosis death model );
       // for accessing the custom variables
   static int energy i = pCell->custom data.find variable index( "energy" );
   static int alpha i = pCell->custom data.find variable index( "alpha" );
   static int beta i = pCell->custom data.find variable index( "beta" );
   static int resistance i = pCell->custom data.find variable index( "resistance");
   static int use rate i = pCell->custom data.find variable index( "use rate" );
       // for sampling the microenvironment
   static int oxygen i = microenvironment.find density index( "oxygen" );
   static int glucose i = microenvironment.find density index( "glucose" );
   static int waste i = microenvironment.find density index( "waste" );
```

### Let's make a new phenotype function (2)

Next, a fun trick: check to see if the cell's dead. If so, set uptake / secretions rates to zero, and overwrite the function pointer so that we don't keep asking dead cells to do things. :-)

```
// if I'm dead, set secretoin rates to zero, and tell us not to
// bother checking ever again.

if( phenotype.death.dead == true )
{
    phenotype.secretion.set_all_secretion_to_zero();
    phenotype.secretion.set_all_uptake_to_zero();

    pCell->functions.update_phenotype = NULL;
    return;
}
```

### Let's make a new phenotype function (3)

#### Let's do the energy model

```
// do the basic energy model
  // use rate : u = alpha + beta + resistance
double alpha = pCell->custom data[alpha i];
double beta = pCell->custom data[beta i];
double resistance = pCell->custom data[resistance i];
pCell->custom data[use rate i] = alpha + beta + resistance;
  // sample the oxygen, glucose, and waste
double oxygen = pCell->nearest density vector()[oxygen i];
double glucose = pCell->nearest density vector()[glucose i];
double waste = pCell->nearest density vector()[waste i];
  // run the ODE. Let's use backwards Euler
double use rate = pCell->custom data[use rate i];
pCell->custom data[energy i] +=
  dt*( alpha*oxygen*glucose + beta*glucose );
pCell->custom data[energy i] /= ( 1.0 + dt*use rate );
```

### Let's make a new phenotype function (4)

```
Let's uptake secretion/ uptake
```

```
// set secretion parameters
phenotype.secretion.secretion_rates[waste_i] = beta * 10.0;
phenotype.secretion.saturation_densities[waste_i] = 1.0;
    // set uptake rates
phenotype.secretion.uptake_rates[oxygen_i] = alpha * 10.0;
phenotype.secretion.uptake_rates[glucose_i] = beta * 0.2;
```

#### Next, the cell cycle rate

```
// set cycle parameters
double energy = pCell->custom_data[energy_i];
double scale = ( energy - 0.1 )/( 0.9 - 0.1 );
if( scale > 1.0 )
{ scale = 1.0; }
if( scale < 0.0 )
{ scale = 0.0; }
phenotype.cycle.data.transition_rate( cycle_start_index ,cycle_end_index ) = 6.94e-4 * scale;
// 1/24 hr^-1 max birth rate, in units of min^-1</pre>
```

### Let's make a new phenotype function (5)

#### Lastly, let's set the death rates

```
// set necrotic death rate
scale = (0.1 - energy)/0.1;
if( scale < 0.0 )
{ scale = 0.0; }
if ( scale > 1.0 )
{ scale = 1.0; }
phenotype.death.rates[necrosis i] = scale * 0.01;
  // 100 minute survival time when zero energy;
  // set the apoptotic death rate
scale = 1.0 + 9.0*(1-pCell->custom data[resistance i])*waste;
phenotype.death.rates[apoptosis i] = scale * 6.94e-6;
// 1% of the max birth rate
                                                         Checkpoint:
return;
```

Now, in create\_cell\_types, let's switch to this rule:

To catch up to the code at this point, unzip checkpoint2.zip

cell defaults.functions.update phenotype = energy based cell phenotype;



## Place cells

### **Modify setup\_tissue()**

#### Some bookkeeping

```
// some bookkeeping
static int energy_i = cell_defaults.custom_data.find_variable_index( "energy" );
static int alpha_i = cell_defaults.custom_data.find_variable_index( "alpha" );
static int beta_i = cell_defaults.custom_data.find_variable_index( "beta" );
static int resistance_i = cell_defaults.custom_data.find_variable_index( "resistance");
static int use_rate_i = cell_defaults.custom_data.find_variable_index( "use_rate" );
```

#### and in four places

```
// choose random properties
pCell->custom_data[alpha_i] = UniformRandom();
pCell->custom_data[beta_i] = UniformRandom();
pCell->custom_data[resistance_i] = UniformRandom();
```

#### Then clear out unneeded junk

# Make a coloring function

### A custom coloring function for SVGs (1)

#### Declare the function in heterogeneity.h

```
std::vector<std::string> energy coloring function( Cell* );
Create it in heterogeneity.cpp
std::vector<std::string> energy coloring function( Cell* pCell )
  // color 0: cytoplasm fill
  // color 1: outer outline
  // color 2: nuclear fill
  // color 3: nuclear outline
  // some bookkeeping
  static int energy i = pCell->custom data.find variable index( "energy" );
  static int alpha \bar{i} = pCell - custom \bar{d}ata.find \bar{v}ariable \bar{i}ndex("alpha");
  static int beta \bar{i} = pCell - custom data.find variable index("beta");
  static int resistance i = pCell->custom data.find variable index( "resistance");
  static int use rate i = pCell - custom data.find variable index("use rate");
  // start black
  std::vector< std::string > output( 4, "black" );
```

### A custom coloring function for SVGs (2)

Sample cell properties and set colors (if not dead)

```
// red: alpha
// green: beta
// blue: resistance
// nucleus: energy (black = 0, white >= 1)
if( pCell->phenotype.death.dead == false )
  int red = (int) round( 255.0 * pCell->custom data[alpha i] );
  int green = (int) round( 255.0 * pCell->custom data[beta i] );
  int blue = (int) round( 255.0 * pCell->custom data[resistance i] );
  int grey = (int) round( 255.0 * pCell->custom data[energy i] );
  char szTempString [128];
  sprintf( szTempString , "rgb(%u,%u,%u)", red, green, blue );
  output[2].assign( szTempString ); // nucleus by alpha, beta, resistance "genes"
  sprintf( szTempString , "rgb(%u,%u,%u)", grey, grey, grey );
  output[0].assign( szTempString ); // cyto by energy
  return output;
```

### A custom coloring function for SVGs (3)

#### Standard dead colors

```
// if not, dead colors
if (pCell->phenotype.cycle.current phase().code == PhysiCell constants::apoptotic )
  // Apoptotic - Red
  output[0] = "rgb(255, 0, 0)";
  output[2] = "rgb(125,0,0)";
// Necrotic - Brown
if ( pCell->phenotype.cycle.current phase().code == PhysiCell constants::necrotic swelling ||
  pCell->phenotype.cycle.current phase().code == PhysiCell constants::necrotic lysed ||
  pCell->phenotype.cycle.current phase().code == PhysiCell constants::necrotic)
  output[0] = "rgb(250, 138, 38)";
  output[2] = "rgb(139,69,19)";
return output;
```

### A custom coloring function for SVGs (4)

#### Select this coloring function in main.cpp

```
std::vector<std::string> (*cell_coloring_function)(Cell*) = energy_coloring_function;
```

#### Now, compile and run!

make && ./heterogeneity



#### Checkpoint:

To catch up to the code at this point, unzip checkpoint3.zip

### Some postprocessing tips

#### convert the SVGs to JPEGs

```
magick mogrify -format jpg -resize 50% snap*.svg
```

#### stitch those JPEGs into an animated GIF

```
magick convert *.jpg out.gif
```

#### create a movie

```
mencoder "mf://snapshot*.jpg" -ovc lavc -lavcopts vcodec=mpeg4:vbitrate=10000:mbd=2:trell
-mf fps=24:type=jpg -nosound -o out.avi
```

Current time: 0 days, 0 hours, and 0.00 minutes, z = 0.00  $\mu m$  889 agents 200 µm 0 days, 0 hours, 0 minutes, and 0.1854 seconds

### Other work that should be done

- Remove hard-coded parameters
  - Use "user\_parmameters" in the XML to set the values
- Fix glucose uptake, energy use rate in model
- Double check that waste-based apoptosis is working

Use internalized substrates for the energy model

Create cloud-hosted version of the model

### **Opportunities**

- Winter 2019: All-new training materials (Thanks, NCI!)
  - A series of ~10-15 minute modules with webapps to illustrate key concepts
  - New website and wiki at PhysiCell.org
  - Basic online development environment on nanoHUB
- Spring 2020: funded hackathon at Indiana University (Thanks, NCI!)
  - Contribute to the PhysiCell ecosystem
- Summer 2020: two NCI-funded 6-week visitors to IU (Yay, NCI!)
  - Likely recruited from the hackathon participants
  - Build a model or new PhysiCell capability, learn to create nanoHUB apps



#### Postdoctoral Research Fellow Position Indiana University, Bloomington

Dr. Paul Macklin, the leader of a dynamic laboratory for open source computational biology and oncology, seeks an interdisciplinary postdoctoral fellow for a project in computational data-driven modeling of breast cancer metastasis, jointly with experimental collaborators at Johns Hopkins University. Work in the MathCancer lab includes agent-based modeling of multicellular systems, multi-substrate diffusion solutions, cancer modeling, high-throughput computing, machine learning, and digital pathology, with a heavy emphasis in building open source technologies that benefit the entire scientific community. Learn more at <a href="http://MathCancer.org">http://MathCancer.org</a>

The fellow will join a dynamic, interdisciplinary, and collaborative research environment with ample opportunities for career development and mentoring. IU SICE is first of its kind and among the largest in the country, with more than 125 full-time faculty and more than 1000 graduate students. SICE has received recognition as a "top-ten program to watch" (Computerworld) thanks to its excellence and leadership in academic programs, interdisciplinary research, placement, and outreach. IUB is also renowned for its top-ranked music school, performing and fine arts, and historic campus. Bloomington is located in the wooded rolling hills of southern Indiana, 50 miles south of Indianapolis. It is a culturally thriving college town with a moderate cost of living, cycling traditions, active lifestyle, and natural beauty.



