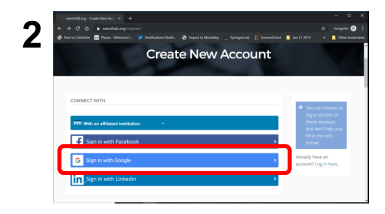
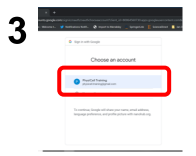
nanoHUB Account

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

• Steps:

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







Agent-based modeling of multicellular systems and cancer in PhysiCell

Part 2: Creating models

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August 13, 2020





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

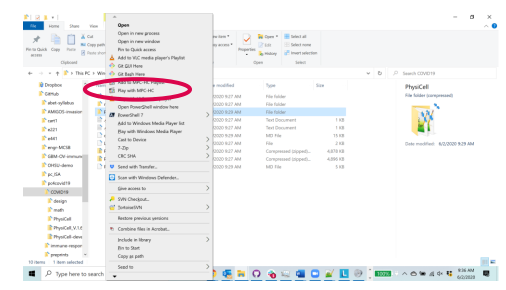


Of particular note, go to ./documentation and open the User_Guide.pdf

Open a terminal in your code directory

 Open the extracted code. Browse until you can see a PhysiCell directory.

- Open a terminal window in the PhysiCell directory
 - Windows:
 - ♦ <shift>-<right click>
 - Open power shell (or command) window here
 - Mac: Sorry, ask a Mac person.



Compiling, running, and visualizing your first project

Sample projects

- It's inefficient (and a little insane) to code new projects entirely from scratch.
- So, we provide sample projects:
 - 2D and 3D template projects
 - Cancer models
 - Synthetic multicellular systems
 - Viral dynamics in tissue
- 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)
- make list-projects: display all available sample projects

Documentation: User Guide Sections 6, 7.

PhysiCell Project Essentials (1)

- Each PhysiCell release includes sample projects. To list them:
 - make list-projects
- Your first step is to populate a project.
 - make project_name>
 - Let's use biorobots-sample:
 - ♦ make biorobots-sample
 - This copies source code, a tailored make file, and configuration files

```
C:\Physicell>make list-projects
Sample projects: template2b template3b biorobots-sample cancer-biorobots-sample heterogeneity-sample cancer-immune-sample virus-macrophage-sample

C:\Physicell>
C:\Physicell>make biorobots-sample virus-macrophage-sample projects/biorobots/custom_modules/* ./custom_modules/
touch main.cpp && cp main.cpp main-backup.cpp

cp ./sample_projects/biorobots/main-biorobots.cpp ./main.cpp

cp ./sample_projects/biorobots/main-biorobots.cpp

cp ./sample_projects/biorobots/Makefile

cp ./config/Physicell_settings.xml ./config/Physicell_settings-backup.xml

cp ./sample_projects/biorobots/config/* ./config/

c:\Physicell>
```

PhysiCell Project Essentials (2)

- Now, compile the project
 - make

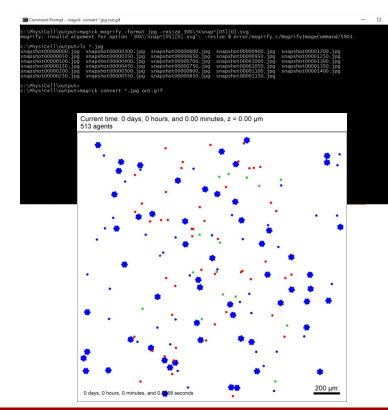
- Then, run the project
 - ./biorobots (Linux, OSX)
 - biorobots.exe (Windows)
- This should take about 5 minutes



PhysiCell Project Essentials (3)

Look at saved data

- Most projects save data to ./output
 - ♦ XML files give metadata, mesh, and substrate info
 - ♦ MAT file save (compressed) substrate and cell data
 - ♦ SVG files are visual quick snapshots
 - ♦ More on loading XML / MAT files in Python later
- Let's convert SVG to rescaled JPEG
 - magick mogrify -format jpg -resize 30% snap*.svg
 - ◆ Convert snapshot00000000.svg, snapshot00000001.svg, ...
 - magick mogrify -format jpg -resize 30% snap*[05][0].svg
 - ♦ Convert snapshot00000000.svg, snapshot00000050.svg, ...
- Now, let's create an animated GIF
 - magick convert *.jpg out.gif



Reference: working with the images

- To convert all the SVG files to PNG format magick mogrify -format png snap*.svg
- To convert every SVG file ending in 0 or 5 to JPG format magick mogrify -format jpg snap*[05].svg
- To convert the JPG files to an animated GIF magick convert *.jpg out.gif
- To create an mp4 movie:

```
ffmpeg -r 24 -f image2 -i snapshot%08d.jpg -vcodec libx264 -pix_fmt yuv420p -strict -2 -tune animation -crf 15 -acodec aac out.mp4
```

PhysiCell Project Essentials (4)

Data cleanup

- Clean up data to get ready for another run
- make data-cleanup

- Reset to a clean slate
 - De-populate the project
 - Get ready for another project
 - make reset

```
c:\Physicell>make data-cleanup
rm -f *.mat
rm -f *.xml
rm -f *.xy
rm -f .output/*
touch ./output/enpty.txt

c:\Physicell>make reset
rf * cpp
p ./sample_projects/Makefile-default Makefile
rm -f ./custom.modules/*
touch ./custom.modules/*
touch .ALL_CITATIONS.txt
touch .ALL_CITATIONS.txt
rm ALL_CITATIONS.txt
cp ./config/Physicell_settings-backup.xml ./config/Physicell_settings.xml
c:\PhysiCell>|
```

Changing settings in a project

First, populate the cancer heterogeneity project

List all available sample projects

 Populate the cancer heterogeneity project

Build the project

Change some settings (next slide)

```
PowerShell 7 (x64)
 C:\Users\PaulT\Downloads\PhysiCell-1.7.1\PhysiCell-1.7.1> make list-projects
ample projects: template2D template3D biorobots-sample cancer-biorobots-sample heterogeneity-sample
                cancer-immune-sample virus-macrophage-sample template
 C:\Users\PaulT\Downloads\PhysiCell-1.7.1\PhysiCell-1.7.1> make heterogeneity-sample
   sample projects/heterogeneity/custom_modules/* ./custom_modules//
    main.cpp && cp main.cpp main-backup.cpp
   /sample_projects/heterogeneity/main-heterogeneity.cpp ./main.cpp
   sample projects/heterogeneity/Makefile .
   config/PhysiCell settings.xml ./config/PhysiCell settings-backup.xml/
  /sample projects/heterogeneity/config/* ./config/
 C:\Users\PaulT\Downloads\PhysiCell-1.7.1\PhysiCell-1.7.1> make _
```

How to change settings in XML

- Open config/PhysiCell_settings.xml
- Major sections:
 - domain -- how big of a region to simulate
 - overall -- how long to simulate, time step sizes
 - parallel -- OpenMP settings
 - save -- how often to save SVG images and full data
 - microenvironment -- settings on diffusing substrates
 - user_parameters -- model-specific settings

Future versions:

cell_definitions -- set default cell properties

Exercise: change settings and run

Let's set the maximum simulation time to 2160 minutes

• Let's set the domain to [-500,500] x [-500,500] to speed it up

- Let's set the oncoprotein standard deviation to 3
- Let's set the max oncoprotein to 9 (3 standard deviations)
- Compile and run as before.

Let's set options and run (1)

- Open ./config/PhysiCell-settings.xml
- Let's set the domain size in the domain block
 - Switch to [-500,500] x [-500,500] x [-10,10] to speed it up

Let's set options and run (2)

- Let's also look at the user_parameters block
 - Let's change the oncoprotein standard deviation (oncoprotein_sd) to 3 (more variation)
 - Let's change the max oncoprotein (oncoprotein_max) to mean + 3 sds = 1 + 9 = 10

Let's set options and run (3)

- Let's look at the overall block
 - Set max time to 1.5 days = 1.5 x 24 x 60 = 2160 minutes

- Let's look at the save block
 - Set the full save interval to 6 hours = 360 minutes

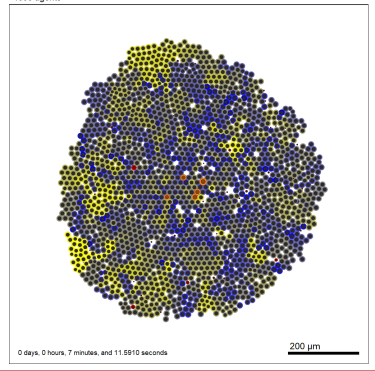
• Now, run! (./heterogeneity)

Let's do a quick visualization

- magick mogrify -format jpg *.svg
- magick convert *.svg out.gif

- We can see that the yellow cells eventually "win": they grow faster and form microcolonies within the tumor
- The effect is greatest on the outside edge: They have access to more O₂ here

Current time: 5 days, 0 hours, and 0.01 minutes, $z = 0.00 \mu m$ 1996 agents

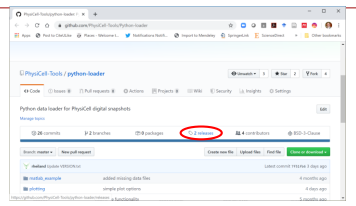


Let's load some data!



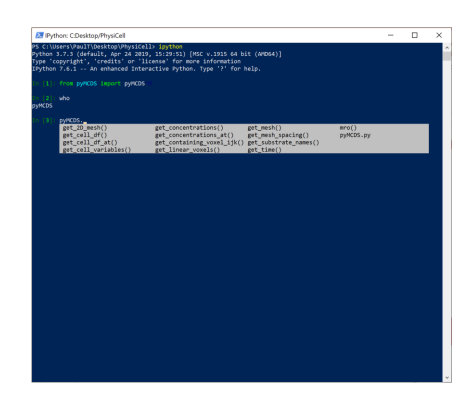
Let's get ready to load in Python

- We'll go to Python-loader and get the source:
 - https://github.com/PhysiCell-Tools/Python-loader
- Get the latest release:
 - Click on "releases" link
 - Click the green "clone or download" button
 - ♦ (For simplicity, I'm using "download ZIP" option)
- Copy the following Python file (end in .py) to the root of PhysiCell
 - pyMCDS



Let's get started in ipython

- Start ipython (interactive python)
 - ipython
- Import the python loader:
 - from pyMCDS import pyMCDS
- Import other useful things
 - import numpy as np
 - import matplotlib.pyplot as plt
- Let's see what is available.
 - Type pyMCDS.
 - Hit "tab" to autocomplete
- Historical note:
 - MCDS = MultiCellDS, our multicellular data standard



Let's load a single saved time

Syntax: result = pyMCDS(filename , directory):

```
mcds = pyMCDS('output0000000.xml', 'output')
```

• Let's get some basic info on the snapshot:

```
print(mcds.get_time()) # what simulation time is saved here?
print(mcds.get_cell_variables()) # what data are saved in the cells?
print(mcds.get_substrate_names()) # what diffusing substrates?
```

mcds.data is a dict. Let's see what's available:

```
mcds.data.keys()
Out[41]: dict_keys(['metadata', 'mesh', 'continuum_variables', 'discrete_cells'])
```



Let's access some cell data

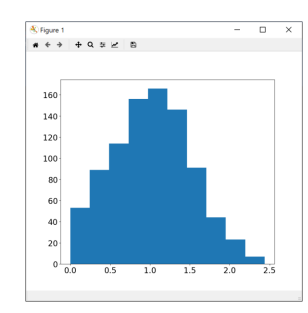
- First, let's find out the mean value of the oncoprotein
 - np.mean(mcds.data['discrete_cells']['oncoprotein'])

```
Out[61]: 1.0177198768372775
```

Let's make sure matplotlib doesn't use small fonts

```
import matplotlib
matplotlib.rc('xtick', labelsize=20)
matplotlib.rc('ytick', labelsize=20)
```

- · Now, let's plot a histogram
 - plt.hist(mcds.data['discrete cells']['oncoprotein'])



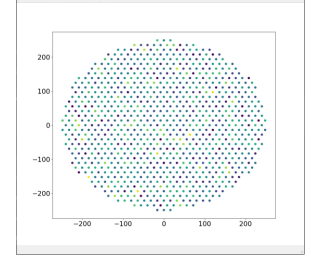


Let's plot the cells

- We'll do a scatter plot of the cells, and color by oncoprotein.
- First, let's grab the data to make our typing easier

```
cx = mcds.data['discrete_cells']['position_x']
cy = mcds.data['discrete_cells']['position_y']
op = mcds.data['discrete_cells']['oncoprotein']
```

- Now, a scatter plot.
 - Note: these are not plotting by the *physical* cell size plt.scatter(cx,cy,c=op)



• This plot is pretty ugly. let's improve it.

Improving the plot scatter plot

Let's replot with bigger dots

```
plt.clf()
plt.scatter( cx , cy, c=op, s=40 )
```

Make sure aspect ratio is right:

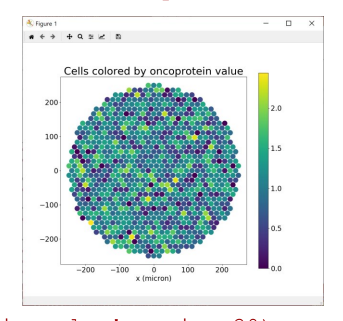
```
plt.axis( 'image' )
```

Now, let's add a colorbar

```
plt.colorbar()
```

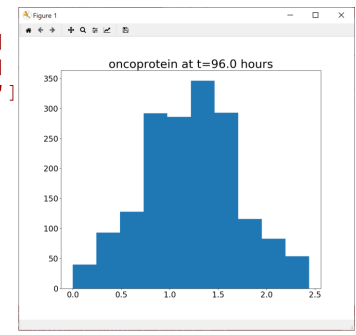
Now, let's add labels

```
plt.title( 'Cells colored by oncoprotein value', size=30)
plt.xlabel( 'x', size=20 )
plt.ylabel( 'y', size=20 )
```



Let's load another time

```
mcds = pyMCDS('output00000006.xml', 'output')
t=mcds.get_time()
cx = mcds.data['discrete_cells']['position_x']
cy = mcds.data['discrete_cells']['position_y']
op = mcds.data['discrete_cells']['oncoprotein']
plt.clf()
plt.hist( op )
plt.title( 'oncoprotein at t=' + \
str(t/60) + ' hours' , size=30)
```



Let's find live and dead cells

- Each cycle model has a unique code
 - Codes ≥ 100 denote death cycles

Let's get the cycle code of each cell, and convert to integers

```
cycle = mcds.data['discrete_cells']['cycle_model']
cycle = cycle.astype( int )
```

Let's find the live cells

```
live = np.argwhere( cycle < 100 ).flatten()
dead = np.argwhere( cycle >= 100 ).flatten()
```



Let's work with these

Live and dead cell counts

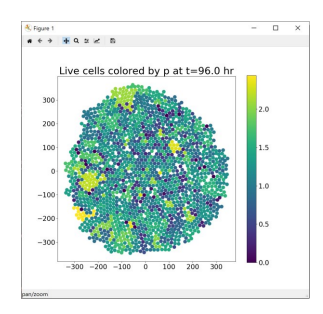
```
n_live = len( live )
n_dead = len( dead )
```

Mean oncoprotein in live cells only

```
np.mean( op[live] )
```

Let's scatter plot of only live cells

```
plt.clf()
plt.scatter( cx[live],cy[live],c=op[live],s=40);
plt.colorbar()
plt.axis('image')
plt.title( 'Live cells colored by p at t=' +str(t/60) + ' hr', size=30)
```

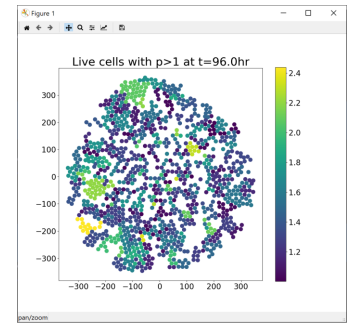


Let's do a fancier search

• Only plot live cells with *p* > 4:

```
ind = np.argwhere( (cycle<100) & (op>4) ) .flatten()
plt.clf()
plt.scatter( cx[ind], cy[ind], c=op[ind], s=40 )
plt.title( 'Live cells with p>4 at t='\
+str(t/60) + 'hr', size=30)
plt.axis('image')
plt.colorbar()
```

• **Note:** The circle size (s=40) will vary based on your desktop resolution and window size. You will need to experiment.



Now let's plot the oxygen

```
plt.clf()
mcds.get_substrate_names();

o2 = mcds.get_concentrations( 'oxygen');
X,Y = mcds.get_2D_mesh();

plt.clf()
plt.contourf(X,Y,o2[:,:,0]);
```

Now let's plot the oxygen with cells

```
circle size = 30
plt.clf()
mcds.get substrate names();
o2 = mcds.get concentrations( 'oxygen');
X,Y = mcds.get 2D mesh();
plt.contourf(X, Y, o2[:,:,0], cmap='spring');
plt.scatter( cx[live], cy[live], c=op[live], s=circle size);
plt.colorbar()
plt.axis('image')
plt.title('Live cells colored by p at t=' +str(t/60) + ' hr', size=30)
# let's plot dead cells as black
plt.scatter( cx[dead], cy[dead], c='k', s=circle size );
```

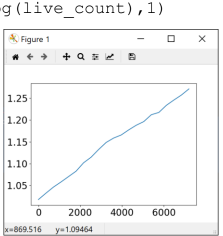
Now, let's do some time series analysis

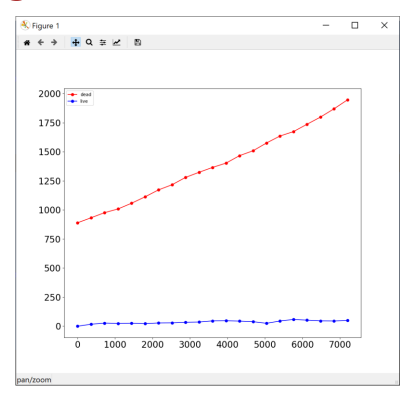
• Let's get live and dead cell counts, mean p (in live cells). We need to loop overall simulation times

```
last index = 6;
live count = np.zeros( last index+1 );
dead count = np.zeros( last index+1 );
mean p = np.zeros( last index+1 );
std p = np.zeros( last index+1 );
times = np.zeros( last index+1 );
for n in range( 0, last index+1 ):
     filename='output'+"%08i"%n+'.xml'
     mcds=pyMCDS(filename, 'output')
     times[n] = mcds.get time()
     cycle=mcds.data['discrete cells']['cycle model']
     p = mcds.data['discrete cells']['oncoprotein']
     live = np.argwhere(cycle<100).flatten()</pre>
     dead = np.argwhere(cycle>=100).flatten()
     live count[n] = len(live)
     dead count[n] = len(dead)
     mean p[n] = np.mean(p[live])
     std p[n] = np.std(p[live])
```

Let's plot and get growth rates

```
plt.clf()
plt.plot( times, live count , 'r-o' )
plt.plot( times, dead count , 'b-o' );
plt.legend( {'live', 'dead' } )
poly=np.polyfit( times, np.log(live count), 1)
# growth rate is 0th element
# in units of 1/min
                             1.25
plt.clf()
                             1.20
plt.plot(times, mean p);
```





More data loading (on your own)

Let's work on data with multiple types

Let's go and run the biorobots sample

```
make data-cleanup
make reset
make biorobots-sample
make
```

Edit the config file to only run to 1440 min, and save every 240 min

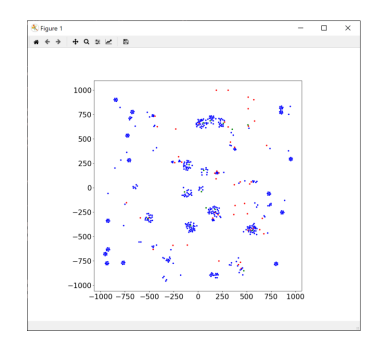
```
./biorobots
```

Let's load an intermediate time

```
n = 3
filename='output'+"%08i"%n+'.xml'
mcds=pyMCDS(filename, 'output')
t = mcds.get time()
cell type=mcds.data['discrete cells']['cell type']
cell type=cell type.astype(int)
ind0 = np.argwhere(cell type==0).flatten();
ind1 = np.argwhere(cell type==1).flatten();
ind3 = np.argwhere(cell type==3).flatten();
cx = mcds.data['discrete cells']['position x']
cy = mcds.data['discrete cells']['position y']
```

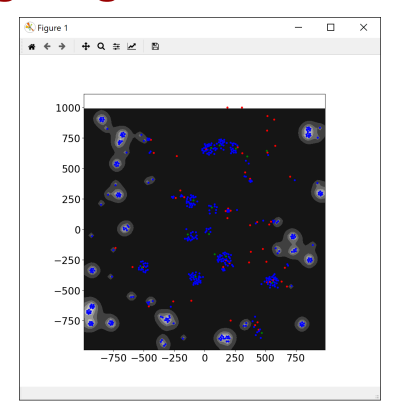
Let's plot each type a different color

```
plt.clf()
plt.scatter(cx[ind0],cy[ind0],c='r',s=circle_size)
plt.scatter(cx[ind1],cy[ind1],c='b',s=circle_size)
plt.scatter(cx[ind3],cy[ind3],c='g',s=circle_size)
plt.axis('image');
```



Overlay on cargo signal

```
mcds.get substrate names();
cs = mcds.get concentrations( 'cargo signal' );
X,Y = mcds.get 2D mesh();
plt.clf()
plt.contourf(X,Y,cs[:,:,0],cmap='gray');
plt.scatter(cx[ind0],cy[ind0],c='r',s=circle size)
plt.scatter(cx[ind1],cy[ind1],c='b',s=circle size)
plt.scatter(cx[ind3],cy[ind3],c='g',s=circle size)
plt.axis('image');
```



Let's build a new model!



Modeling task

- Iteratively build up a cancer model, starting simple and adding more
- Model 1:
 - Proliferating cancer cells with motility
 - Oxygen uptake
 - Can mostly be designed in XML
- Model 2:
 - Refine model 1 to add an aggressive subclone
 - more motile and proliferative, but higher O2 uptake rate
- Model 3:
 - Refine model 2 so that proliferation and death are oxygen-based
 - Add a small probability that



Model 1

- In the XML configuration file:
 - Define diffusing substrates
 - Create cell definition(s)
 - Set time, domain, other settings
 - Custom parameters as needed

- In the custom.cpp file:
 - Add cells in the initial simulation

Use the template project

clean up date (if any)

```
make data-cleanup
```

reset to blank slate

```
make reset
```

populate and compile the (new) template project

```
make template
make
```



XML file: substrates

• Let's do oxygen with standard diffusion coefficients, decay for a 1 mm length scale, and 5% (38 mmHg) Dirichlet boundary condition

$$L = \sqrt{\frac{D}{\lambda + U}}$$

XML file: default cell definition

- The default cell definition sets default parameters for all cell types.
- It also sets custom variables available for all other cell types.

- If you create a cell in a simulation without choosing a cell definition, it will use the default cell definition.
- I suggest mostly leaving it alone. But make sure the <secretion> section has the same variables as the microenvironment. (Some day we will write an XML validator. It would be a great community project!)
- But definitely add custom variables the default definition so that all cell types have the same custom variables.
- · We don't need any for now.

XML file: default cell definition - secretion

- Two gotcha's:
 - Need to make sure that secretion and chemotaxis only reference things that are defined in your microenvironment.

XML file: default cell definition - secretion

- Two gotcha's:
 - Need to make sure that secretion and chemotaxis only reference things that are defined in your microenvironment.

XML file: cancer cell definition

- We will create a cancer cell definition that "inherits" all the physical rate parameters from the default definition.
- We only need to define how it differs from default:
 - proliferation
 - death (set apoptosis to zero)
 - O2 uptake (no change from default)
 - motility
- Copy the default cell definition. Glve it a new ID (1)

XML file: cancer cell definition

- Copy the default cell definition.
 - Give it a name (cancer)
 - Give it a new ID (1)
 - Make sure it inherits from the default cell definition
- If it's already there, overwrite it. Delete any other cell defs.

XML file: cancer: cycling

- Let's leave all the cycle parameters alone, except that we want cells to spend (on average) 1 hour in the G0/G1 cycle phase.
- Let's edit the <cycle> part of <phenotype>

Notice that we only wrote the rate that differs from the default model.

XML file: cancer: cycling (alternative)

• Alternatively, you can set a transition rate from phase 0 (G0/G1) to phase 1 (S). For a mean holding time of 60 min, the rate is 1/60 min⁻¹

```
<phenotype>
  <cycle code="6" name="Flow cytometry model (separated)">
        <!-- phases are: G0/G1 , S, G2, M -->
        <!-- use phase_transition_rates OR phase_durations -->
        <phase_transition_rates units="1/min">
        <rate start_index="0" end_index="1" fixed_duration="false">0.016667</rate>
        </phase_transitions>
        </cycle>
```

• You can use durations **OR** rates, but not both within one cell def.

XML file: cancer: death

Let's set the apoptosis and necrosis rates to zero

XML file: cancer: motility

• Let's set biased random migration up oxygen gradients.

XML file: custom parameters

- Let's add a parameters for:
 - initial number of cells
 - max |x| and |y| coordinates

custom.cpp (1)

- Our cells are set up. Now we need to place some
- Open ./custom_modules/custom.cpp
- Look for a function called setup_tissue

 We'll read our parameter for number of cells, and place them randomly within the coordinate bounds

custom.cpp (2)

```
void setup tissue( void )
  double min = -parameters.doubles( "coordinate max" );
  double max = parameters.doubles( "coordinate max" );
  double range = max - min;
  Cell* pC;
  // place cancer cells
  for ( int n = 0 ; n < parameters.ints("number of cancer") ; <math>n++ )
      std::vector<double> position = {0,0,0};
     position[0] = min + UniformRandom()*range; // choose position
     position[1] = min + UniformRandom()*range;
     pC = create cell( get cell definition("cancer") ); // place a cancer cell
     pC->assign position( position ); // set its position
  return;
```

Modify the color function

```
std::vector<std::string> my coloring function( Cell* pCell )
  static int cancer type = get cell definition( "cancer" ).type;
  // start with flow cytometry coloring
  std::vector<std::string> output = false cell coloring cytometry(pCell);
  // if it's not a cancer cell, color it black
  if( pCell->phenotype.death.dead == false && pCell->type != cancer type )
     output[0] = "black";
     output[2] = "black";
  return output;
```

One last thing:

- Make sure there aren't any extra functions in the cell setup
- (Future versions of PhysiCell will have a cleaner template project!)
- Search for the function create_cell_types
- Make sure this section is empty:

```
/*
    Put any modifications to individual cell definitions here.

This is a good place to set custom functions.

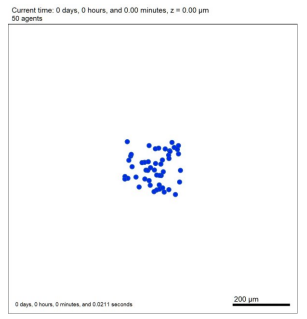
*/
// nothing here!
    /*
    This builds the map of cell definitions and summarizes the setup.
    */
```

Build and run!

Set the max time to 1 day (1440 minutes), and output every 15 minutes

make

./project



Next steps

• I'll add Models 2 and 3 as exercises, with code solutions.

 A great way to learn is to look at existing codes. Look at the sample projects. See how they work. Give them a try!

We are developing a detailed suite of training materials. Stay tuned!

• We are happy to meet one-on-one to consult on possible projects. Just give us a shout, and we'll help set up your project!

Some models to explore

On nanoHUB:

- pc4heterogen: heterogeneous cancer growth (https://nanohub.org/tools/pc4heterogen)
- **pc4cancerbots:** use the "biorobots" as a cell-based cancer therapy (https://nanohub.org/tools/pc4cancerbots)
- pc4livermedium: tumor-stroma biomechanical feedbacks (https://nanohub.org/tools/pc4livermedium)
- pc4cancerimmune: basic cancer immunotherapy model (https://nanohub.org/tools/pc4cancerimmune)
- pc4covid19: COVID-19 simulation model (https://nanohub.org/tools/pc4covid19)
- trmotility: training on biased random cell migration (https://nanohub.org/tools/trmotility)
- pc4thanos: Avengers Endgame battle using cell rules (https://nanohub.org/tools/pc4thanos)

Bundled in PhysiCell:

• biorobots, cancer biorobots, heterogeneity, cancer immunotherapy (3D version), virus-macrophage sample, project templates

Further reading (1)

BioFVM method paper (3-D diffusion)

A. Ghaffarizadeh, S.H. Friedman, and P. Macklin. BioFVM: an efficient, parallelized diffusive transport solver for 3-D biological simulations. Bioinformatics 32(8):1256-8, 2016. DOI: 10.1093/bioinformatics/btv730.

PhysiCell method paper (agent-based model)

A. Ghaffarizadeh, R. Heiland, S.H. Friedman, S.M. Mumenthaler, and P. Macklin. PhysiCell: an open source physics-based cell simulator for 3-D multicellular systems. PLoS Comput. Biol. 14(2):e1005991, 2018. DOI: 10.1371/journal.pcbi.1005991.

PhysiBoSS (PhysiCell + MaBoSS for Boolean networks)

G. Letort, A. Montagud, G. Stoll, R. Heiland, E. Barillot, P. Macklin, A. Zinovyev, and L. Calzone. PhysiBoSS: a multi-scale agent based modelling framework integrating physical dimension and cell signalling. *Bioinformatics* 35(7):1188-96, 2019. DOI: 10.1093/bioinformatics/bty766.

xml2jupyter paper (create GUIs for cloud-hosted models)

R. Heiland, D. Mishler, T. Zhang, E. Bower, and P. Macklin. xml2jupyter: Mapping parameters between XML and Jupyter widgets. *Journal of Open* Source Software 4(39):1408, 2019. DOI: 10.21105/joss.01408.

• PhysiCell+EMEWS (high-throughput 3D PhysiCell investigation)
J. Ozik, N. Collier, J. Wozniak, C. Macal, C. Cockrell, S.H. Friedman, A. Ghaffarizadeh, R. Heiland, G. An, and P. Macklin. High-throughput cancer hypothesis testing with an integrated PhysiCell-EMEWS workflow. BMC Bioinformatics 19:483, 2018. DOI: 10.1186/s12859-018-2510-x.

PhysiCell+EMEWS 2 (HPC accelerated by machine learning)

J. Ozik, N. Collier, R. Heiland, G. An, and P. Macklin. Learning-accelerated Discovery of Immune-Tumour Interactions. *Molec. Syst. Design Eng.* 4:747-60, 2019. DOI: 10.1039/c9me00036d.



Further reading (2)

A review of cell-based modeling (in cancer):

J. Metzcar, Y. Wang, R. Heiland, and P. Macklin. A review of cell-based computational modeling in cancer biology. *JCO Clinical Cancer Informatics* 3:1-13, 2019 (invited review). DOI: 10.1200/CCI.18.00069.

Progress on multicellular systems biology:

P. Macklin, H.B. Frieboes, J.L. Sparks, A. Ghaffarizadeh, S.H. Friedman, E.F. Juarez, E. Jockheere, and S.M. Mumenthaler. "Progress Towards Computational 3-D Multicellular Systems Biology". In: . Rejniak (ed.), *Systems Biology of Tumor Microenvironment*, chap. 12, pp. 225-46, Springer, 2016. ISBN: 978-3-319-42021-9. (invited author: P. Macklin). DOI: 10.1007/978-3-319-42023-3 12.

Challenges for data-driven multicellular systems biology

P. Macklin. Key challenges facing data-driven multicellular systems biology. *GigaScience* 8(10):giz127, 2019. DOI: 10.1093/gigascience/giz127

COVID-19 community preprint

Y. Wang et al., Rapid community-driven development of a SARS-CoV-2 tissue simulator. *bioRxiv* 2020.04.02.019075 (2020). DOI: 10.1101/2020.04.02.019075

Some links

- PhysiCell project & downloads: http://PhysiCell.org
- Twitter updates: https://twitter.com/PhysiCell
- Tutorials: http://www.mathcancer.org/blog/physicell-tutorials/
- Tools (in progress): https://github.com/PhysiCell-Tools
- Training (in progress): https://github.com/PhysiCell-Training
- Wiki (in progress): http://PhysiCell.org/wiki