### SUBE Annual Retreat

Saturday 9/23/23

### Agenda

### Le Meridien in Central Square

- Engaging workshops
- Inspiring talks
- Interactive activities
- Breakfast & lunch provided

## Action Kitchen in Seaport

Cook a chef-guided dinner as a group

# Recharge and grow community!

- Connect with peers in the BiologicalEngineering and Biology departments
- Expand your knowledge
- Make new friends





### **RSVP**

16 lucky participants chosen by lottery!

Enter lottery here: <a href="https://forms.gle/ka">https://forms.gle/ka</a>
<a href="b4TEPRNLxxpAVVJ6">b4TEPRNLxxpAVVJ6</a>



https://forms.gle/kab4TEPRNLxxpAWJ6

M1D3: Use immunofluorescence staining to assess repair foci experiment

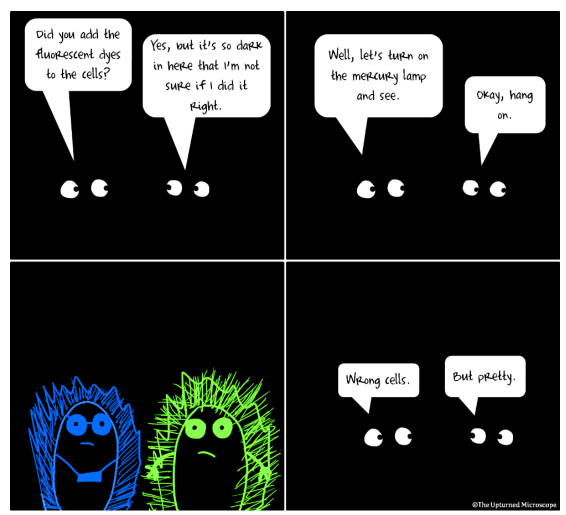
- 1. Prelab
- 2. Antibody staining for γH2AX assay
- 3. Image coverslips

### **Office Hours**

Jamie: 11a-12p Mondays

Noreen: 3-5pm Mondays

Becky: 10a-12p Wednesdays



### Mod1 Overview

### Last lab:

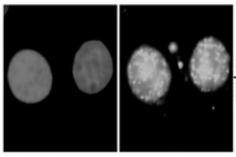
Treated MEF cells & fixed with PFA

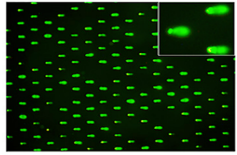
### This lab:

Stain cells with fluorescent antibodies & mount coverslips

#### Next lab:

Analyze yH2AX data & begin CometChip

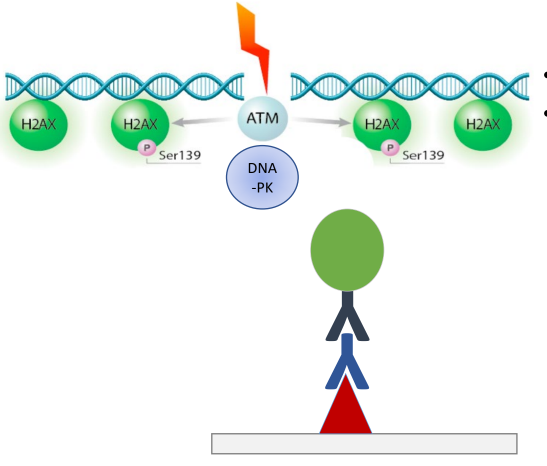




- 1. Use repair foci experiment to measure DNA breaks
- Examine effect of  $H_2O_2$  +/- As on double strand DNA breaks by measuring  $\gamma H2AX$  foci formation

- 2. Use high-throughput genome damage assay to measure DNA damage
- Measure effects of H<sub>2</sub>O<sub>2</sub> +/- As on DNA damage by measuring DNA migration in agarose matrix

## Using immunofluorescence: γH2AX assay to detect double-strand DNA breaks

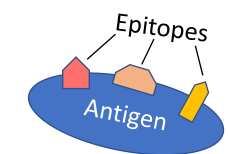


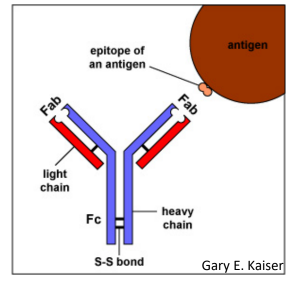
- Histone H2AX phosphorylated at Ser139 if DSB
- Antibodies against γH2AX (phosphorylated form)

protein of interest	A γH2AX	
primary antibody	k mouse anti-γH2AX	
secondary antibody	★ goat anti-mouse	
Fluorophore (conjugated to secondary antibody) exc./ em. wavelengths	488/525 nm	

## Considerations for using antibodies in the lab

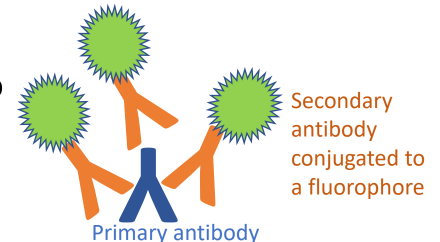
- Antibodies bind to specific epitopes on antigens
  - Antigens may have multiple epitopes





### Primary antibodies vs secondary antibodies

- Primary antibody recognizes the antigen
  - Specific protein sequence
  - Specific conformation of protein
  - Specific state of protein (i.e. phosphorylation)
- Secondary Ab recognizes the species of the primary Ab
  - Often conjugated to tag for visualization
    - Enzyme or fluorophore
  - Amplifies signal through multiple bindings
  - Consider sample species when choosing antibodies!



## Polyclonal vs. monoclonal antibodies

### **Polyclonal**

- How it's made: animal (often rabbit) immunized with antigen of interest then antibodies collected from blood sera and affinity purified
- Advantages:
  - Less expensive and faster to produce than monoclonal
  - Multiple antibodies in one polyclonal mixture can increase antigen recognition by binding multiple epitopes
    - Especially useful for proteins with low expression
- Disadvantages:
  - Variability from lot to lot

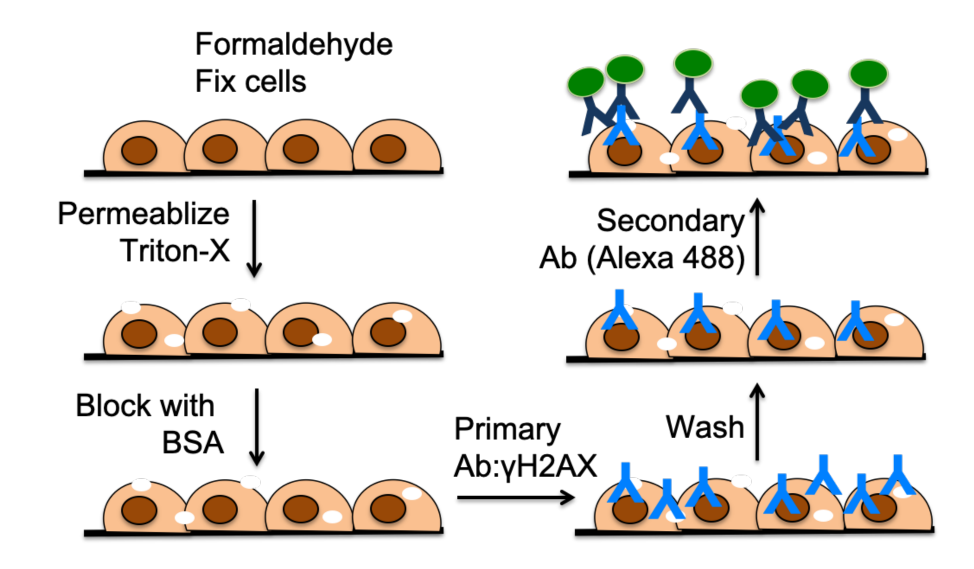
### **Monoclonal**

- How it's made: animal (usually mouse) immunized with antigen of interest then B cells from spleen are harvested and fused with myeloma cells to create hybridoma cell line that will continually produce single antibody clone
- Advantages:
  - Very consistent
  - Binds single epitope (can also be disadvantage)
- Disadvantages:
  - More expensive and requires animal sacrifice

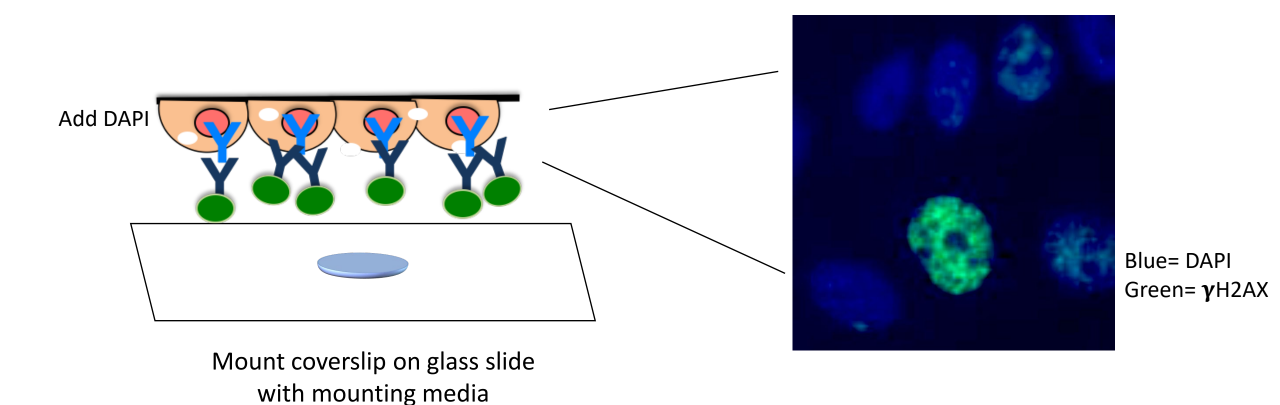


Antigen

## Using immunofluorescence (IF): steps in protocol



## Finish IF by adding DAPI, then mount slides for imaging



## Homework discussion

**Experimental Schematics** 

## Notes on experimental schematics

Research Question: How do I make the best baguette?







Add either 300g, 500g or 700g of warm water (37C)



Mix the dough in the mixing bowl, either by hand or using a rubber spatula



Combine 1000 g All purpose flour, 1 tsp

yeast, 1 tsp salt in a large mixing bowl



Turn the dough over onto a clean work surface and knead by pushing and folding the dough onto itself, creating gluten strands until smooth and elastic



Allow your dough to prove by returning it to the mixing bowl, covering with a wet teacloth in the fridge overnight

## What should be in the Title and Caption?

Title: State what is shown / represented in the schematic

### Caption:

- Explain the flow of information using concise / clear language
- Expand on text shown in figure labels to eliminate excess wordiness / clutter from the figure
- Define all abbreviations / jargon / labels / symbols

## Notes on experimental schematics

Research Question: How do I make the best baguette?

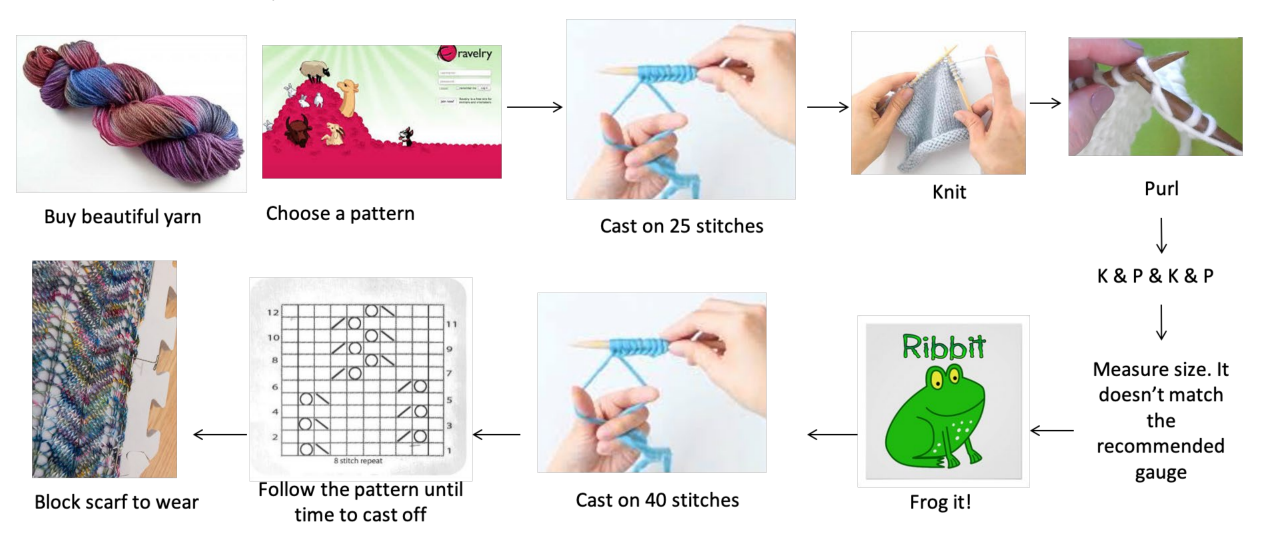
Score the dough



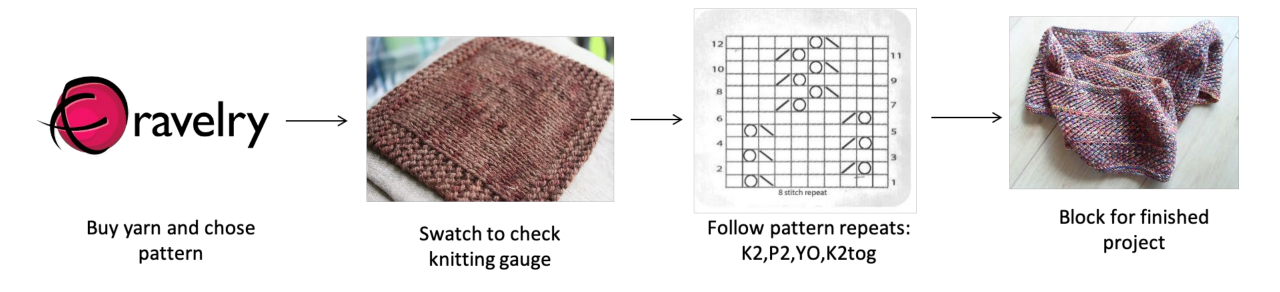
**Figure 1: Optimizing hydration for baguettes.** Baguettes were made with varying levels of water. Doughs were proofed at 4C overnight, then deflated, shaped, and proved at room temperature. Quality data was collected by subject reporting on a scale of 1-5. ON = overnight. RT = room temperature.

### Notes on experimental schematics...

How does Becky knit a scarf?



## Revised example:



**Figure 1: Becky's knitting process.** Becky follows a specific protocol to knit a scarf. She choses her yarn and checks the pattern before following the written pattern and blocking to complete the project. K2= knit two, P2= purl 2, YO= yarn over, K2tog= knit two together

## In lab today:

- 1. Complete IF staining for H2AX
  - 1. Blocking: Noreen lecture on Research Talk
  - 2. Primary & Secondary Ab: View H2AX images on 7<sup>th</sup> floor microscope

### HW due M1D4

Tues 9/26!

Thursday = Comm Lab lecture, no lab

### (group)

Create an experimental schematic for the H2AX staining process

### (individual)

- 1. Write outline for Research Talk
- 2. Read paper for discussion (linked on M1D4)
- 3. Visit Comm Lab before M1D5

## Research Talk due Saturday, Sept 30

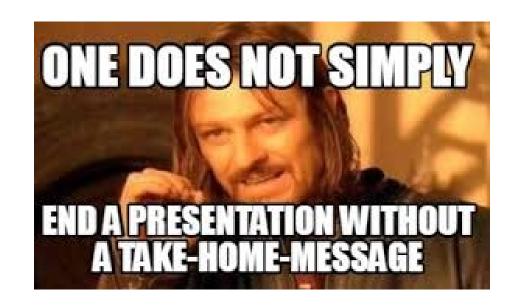
- Prepare a video of you verbally discussing your research
  - Use any device or Zoom
  - No visuals / slides
  - Do not edit / splice the video

#### Submit to Gmail account!

- bioeng20.109@gmail.com
- Remember to follow file name guidelines

## Presentation should be 3 min (+/- 15 sec)

- Introduce yourself
- Provide important background information
- Describe key results
  - Briefly describe critical methods used to generate important data
  - Use quantitative descriptions when discussing results
- Highlight the take-home message



## What data / results should be included?

Only include information on the gamma-H2AX experiment

How were the cells treated?

How were the cells stained?

How were the data analyzed?

• What are the results?

## Review assignment description on wiki

Category	Elements of a strong presentation	Weight
Introduction	<ul> <li>Introduce yourself and the research</li> <li>Summarize the background information necessary to understand the research</li> <li>State the research question</li> </ul>	25%
Methods & Data	<ul> <li>Provide ONLY the method information necessary to understand the results</li> <li>Give complete and concise explanations of the results</li> <li>Relate the results to the central question</li> </ul>	25%
Summary & Conclusions	Highlight the key finding(s) relevant to the central question / hypothesis	25%
Organization	Give a logical, easy-to-follow narrative     Include transition statements	15%
Delivery	<ul> <li>Show confidence / enthusiasm and speak clearly</li> <li>Use appropriate language (technical or informal, as appropriate)</li> <li>Be mindful of the time limit (3 minutes +/- 15 seconds!)</li> </ul>	10%

The Research talk will be graded by Dr. Noreen Lyell with input from Dr. Becky Meyer and Jamie Zhan.