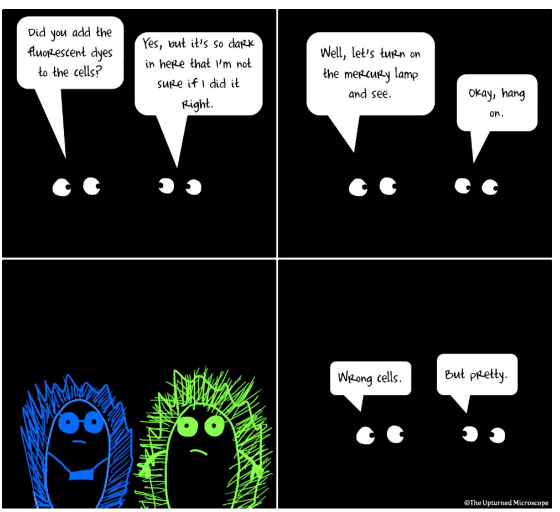
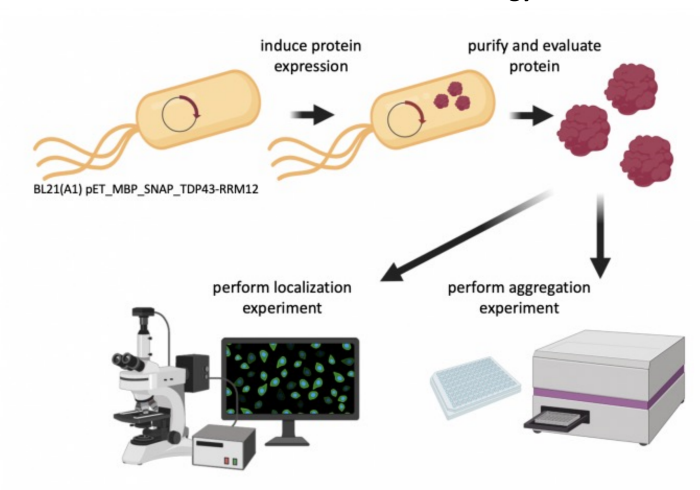
M1D3: Use immunofluorescence staining to assess repair foci experiment

- 1. Prelab
- 2. Antibody staining for TDP43 localization



## Overview of Mod 1 experiments

Research goal: Use functional assays to characterize ligands identified as binders to TDP43 from SMM technology

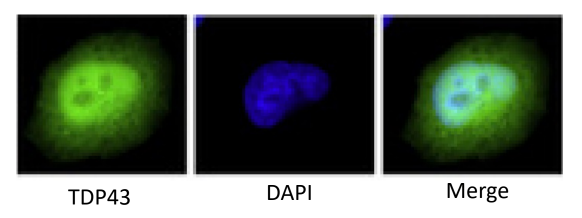


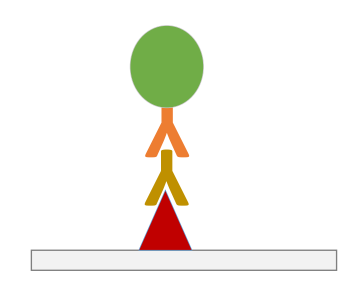
# Using immunofluorescence: Localization of TDP43 in CAD cells

 CAD cells expressing endogenous TDP43 are **DMSO** treated for 1 hour with small molecule 3 µM small molecule **EMPTY WELLS** 30 µM small molecule TDP43

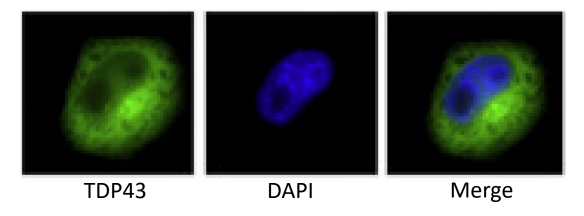
# Using immunofluorescence: Localization of TDP43 in CAD cells

#### **Condition 1:**





#### **Condition 2:**



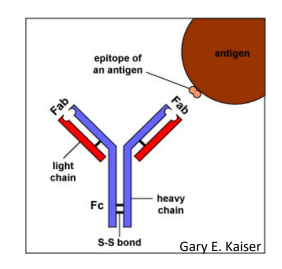
protein of interest	▲ TDP43
primary antibody	rabbit anti-mouse anti-TDP43
secondary antibody	description
Fluorophore (conjugated to secondary antibody) exc./ em. wavelengths	488/525 nm

## Considerations for using antibodies in the lab

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

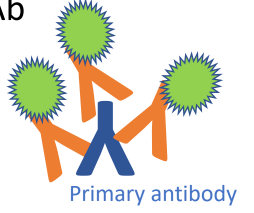


- Specific protein sequence
- Specific conformation of protein
- Specific state of protein (i.e. phosphorylation)



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!



Secondary antibody conjugated to a fluorophore

Epiţopes

Antigen

## 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

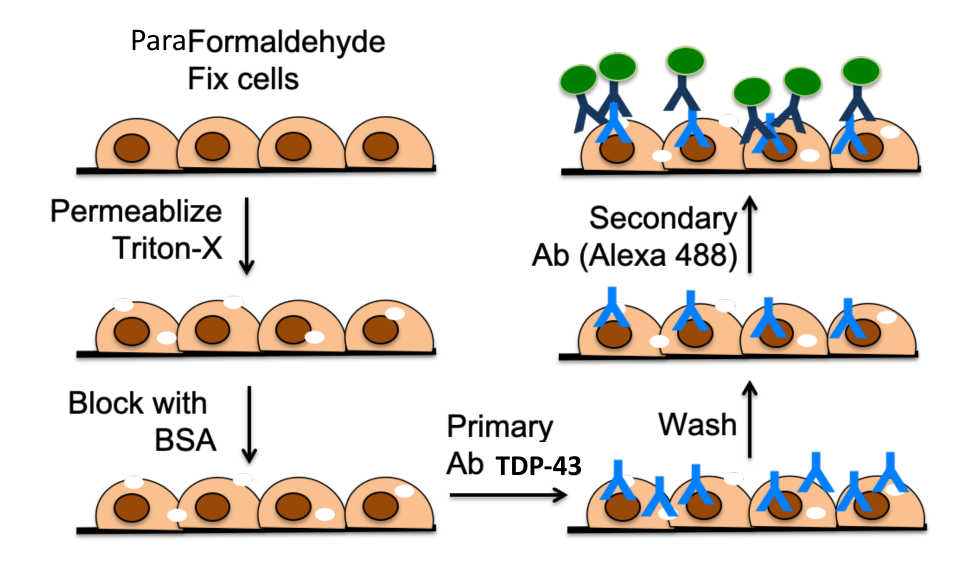
### **Monoclonal**

- 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

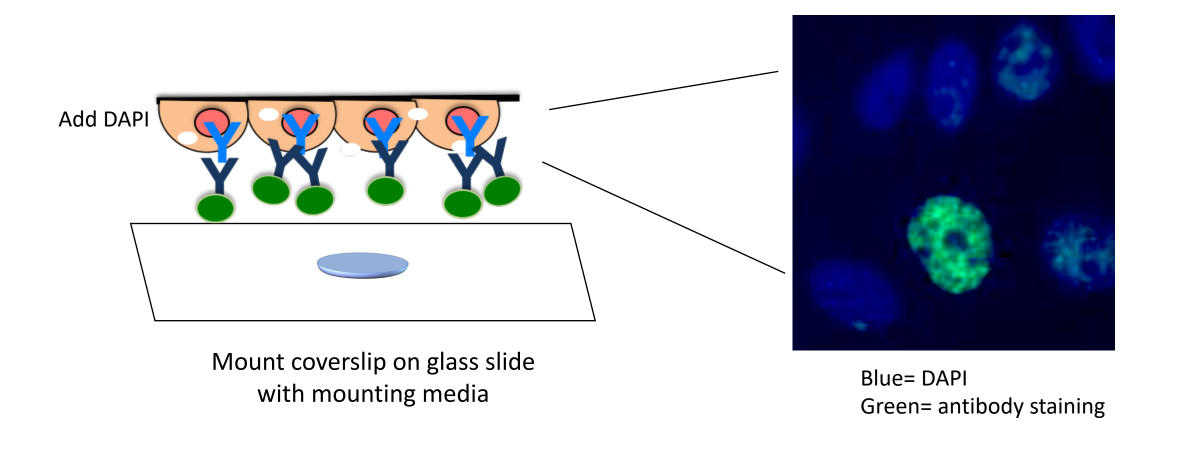


Antigen

## Using immunofluorescence (IF): steps in protocol



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



## For today:

- 1. Complete IF staining for TDP43 Localization
  - 1. Christine will demo staining chamber assembly
- 2. Work on Methods revision with partner

### For M1D8

- Individually, answer the question prompts for the Implications and Future works section of your Data Summary
- 2. With your lab partner, revise your methods homework and add M1D4-M1D5