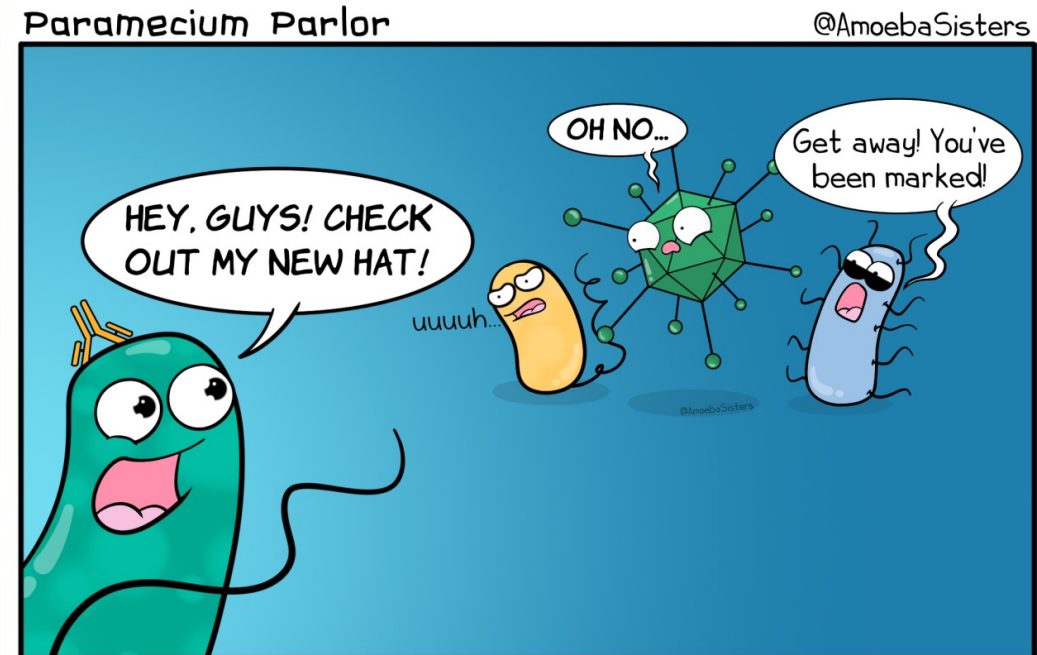


# M1D5:

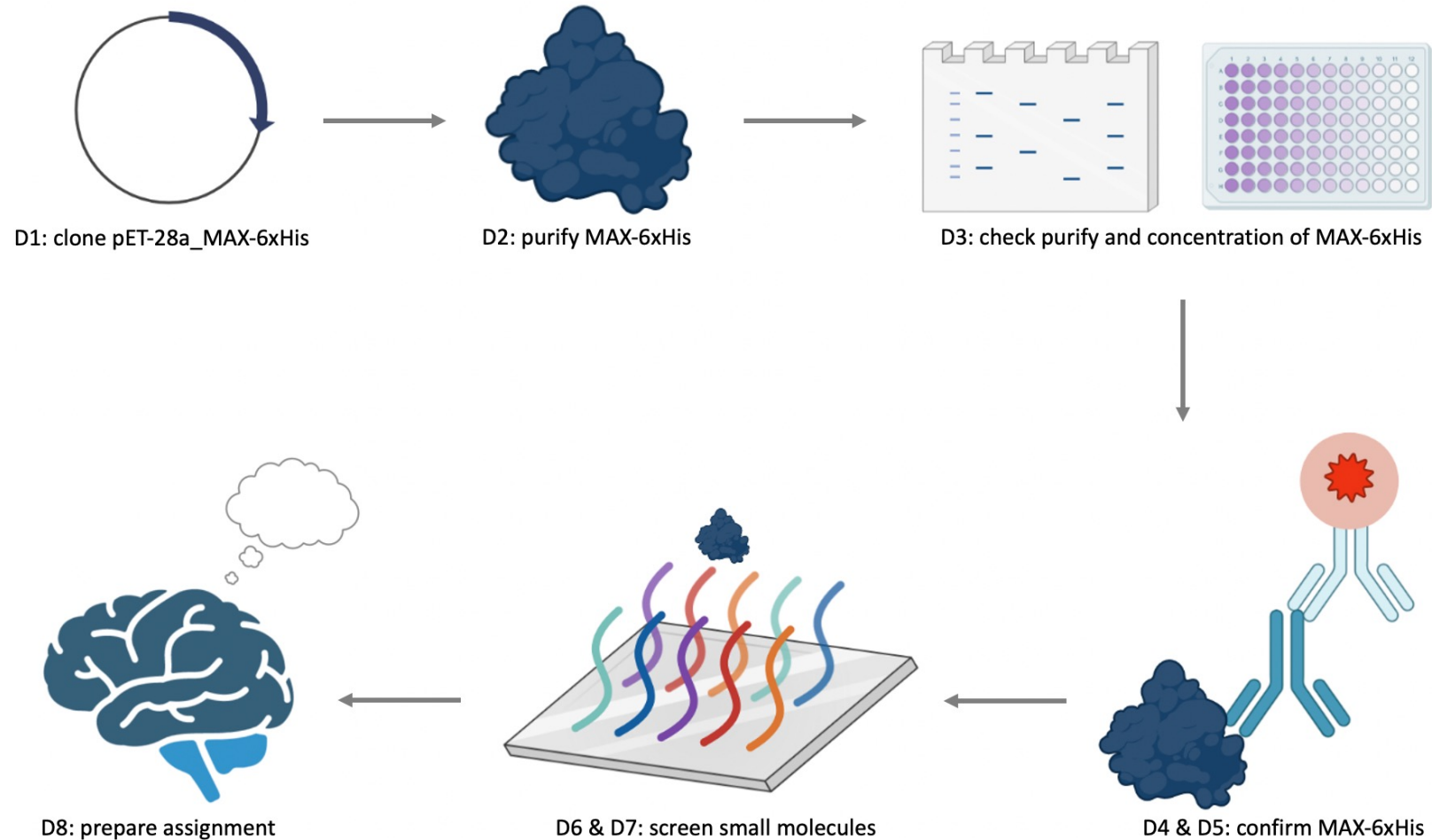
## Image Western blot of purified protein

1. Prelab discussion
2. *Complete western blot experiment – moved to Thursday!*
3. Prepare Data summary slide

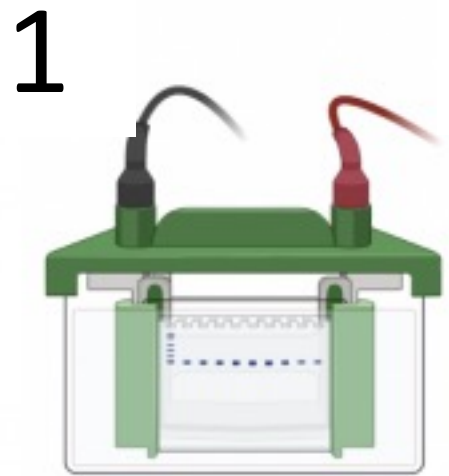


And, thus, Ned learned the dangers of accessorizing with antibodies.

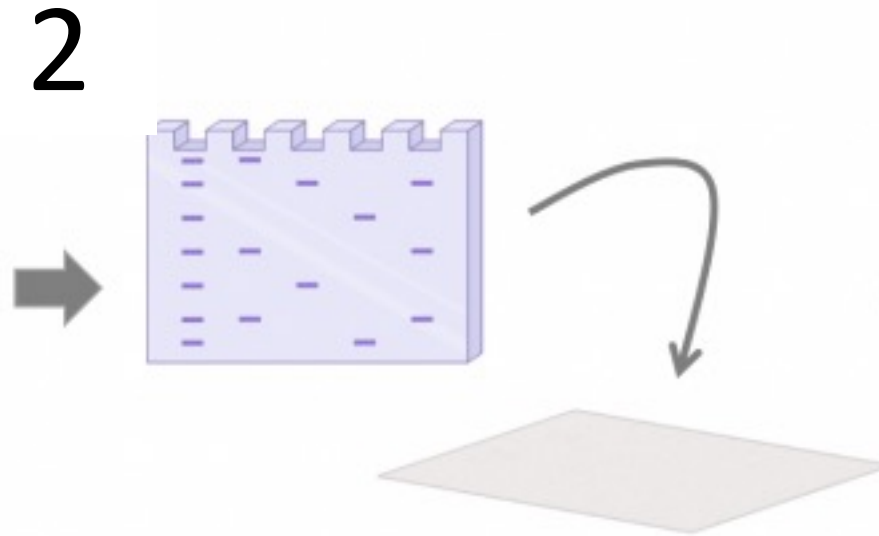
# Overview of Mod 1 experiments:



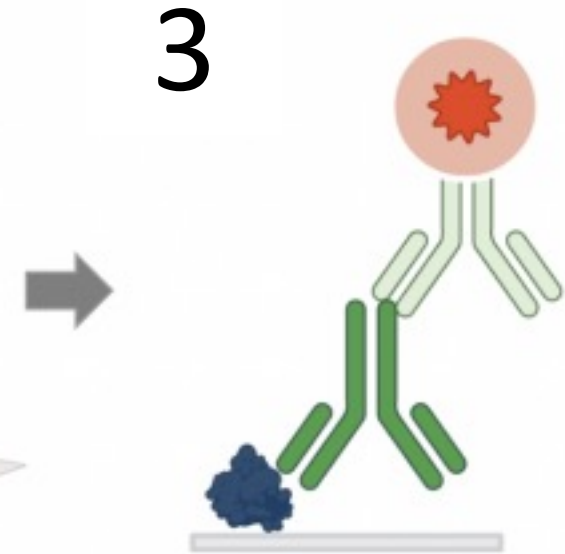
# Western blots probe for specific proteins



separate proteins using electrophoresis



transfer proteins onto nitrocellulose membrane



probe membrane using antibodies

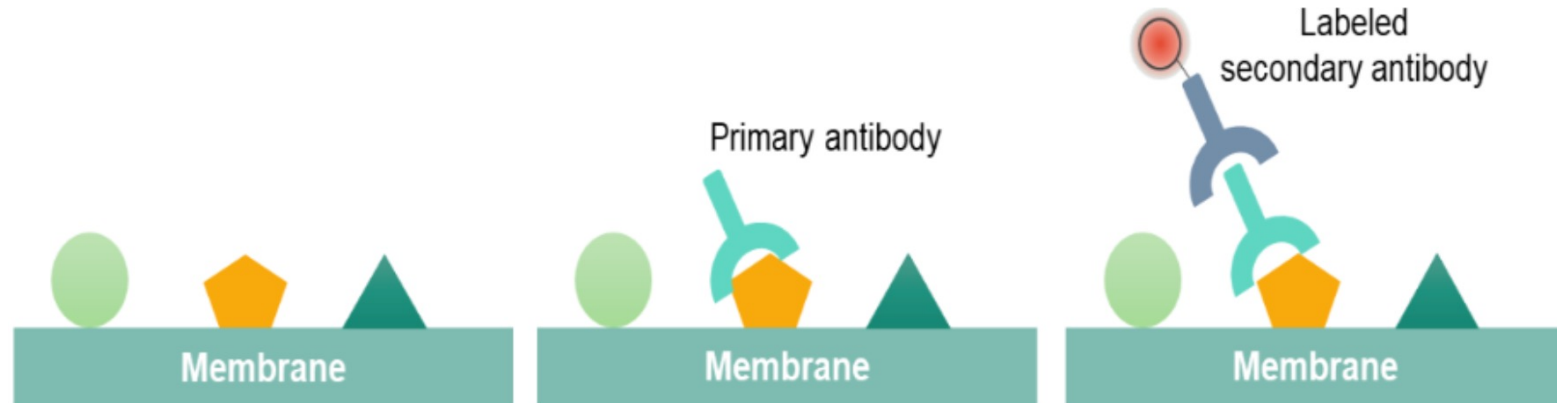
# Let's review the separation and transfer steps...

Step 1: SDS-PAGE used to separate proteins into distinct bands

Step 2: Protein bands from polyacrylamide gel transferred to a nitrocellulose membrane via applying a current

- Why is it necessary to transfer proteins onto a membrane?

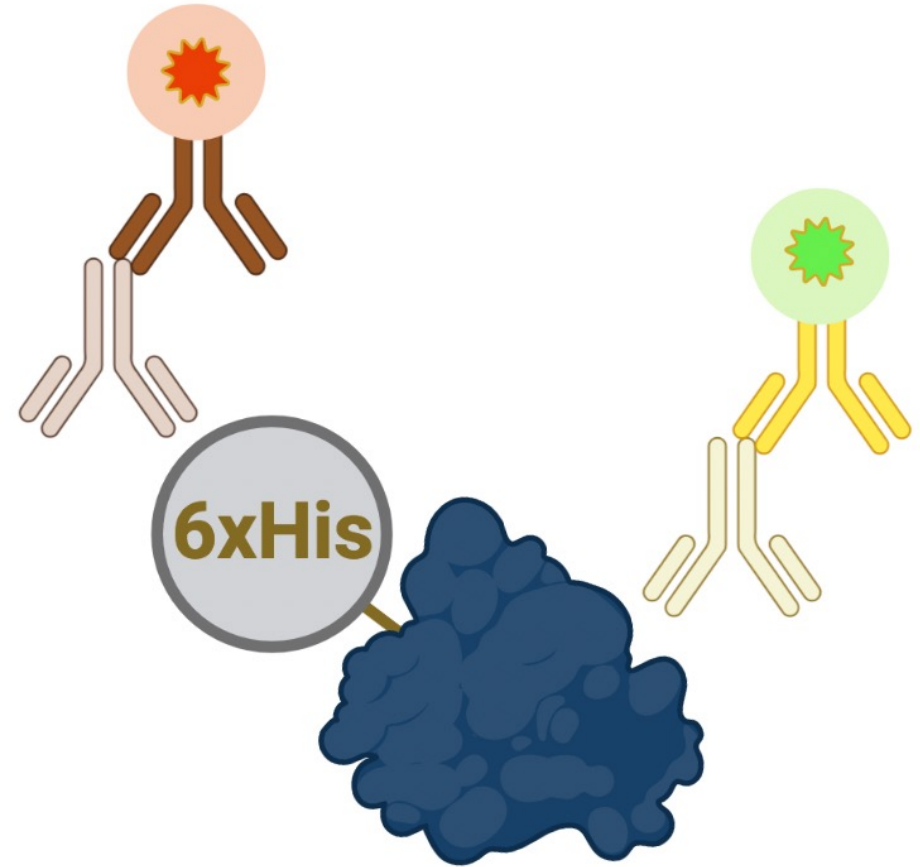
# Step 3: probe membrane using antibodies



- Primary antibody raised against protein of interest to identify band that corresponds to specific protein on the blot
- Secondary antibody raised against the species of the primary antibody to visualize band that corresponds to specific protein of interest

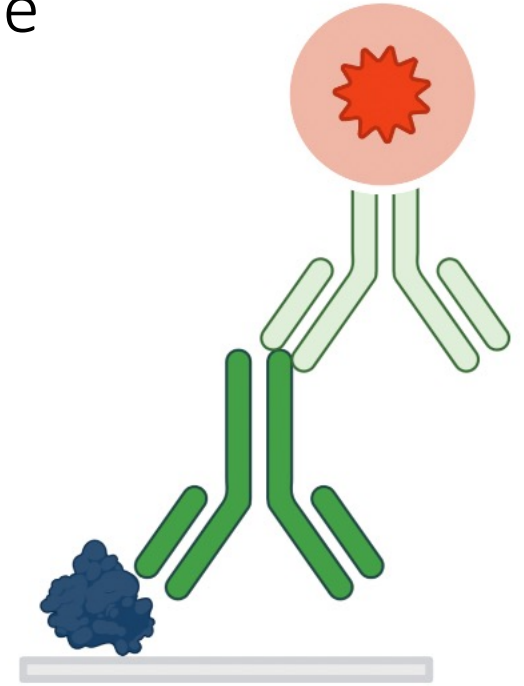
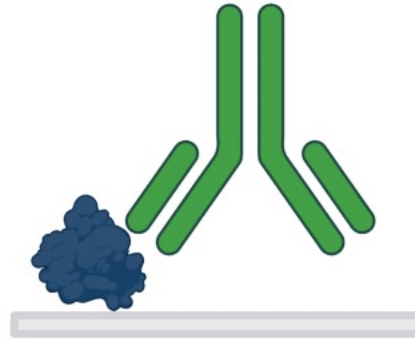
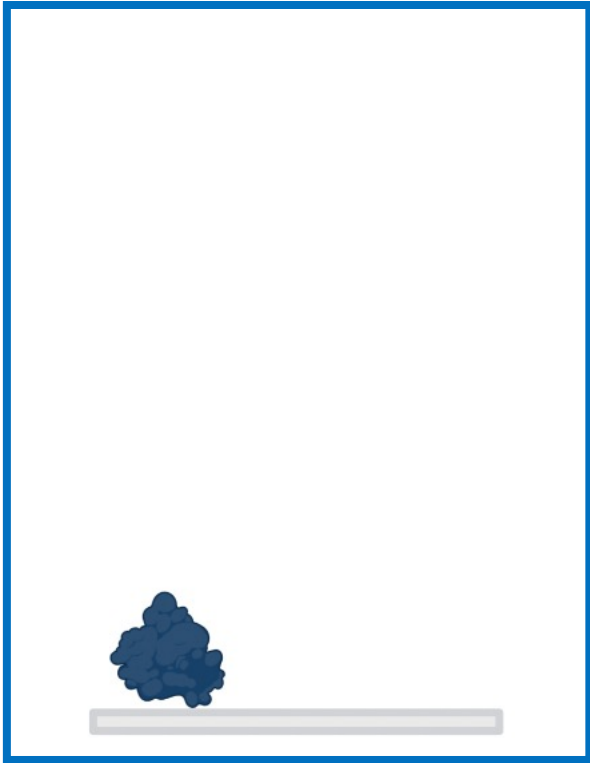
# What antibodies are we using?

- To probe for MAX...
  - 1° = rabbit  $\alpha$ -MAX
  - 2° = donkey  $\alpha$ -rabbit IR680
- To probe for 6xHis tag...
  - 1° = mouse  $\alpha$ -His
  - 2° = goat  $\alpha$ -mouse IR800



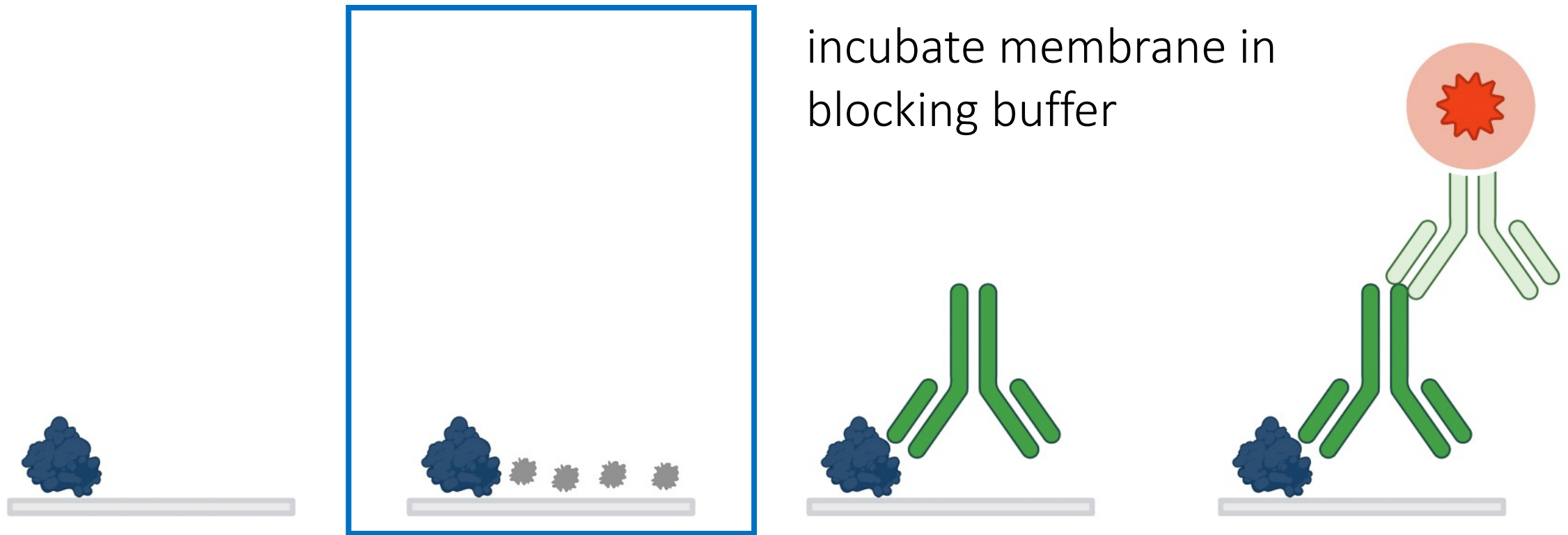
# Antibodies used to detect specific protein

transfer proteins from polyacrylamide gel onto nitrocellulose membrane



- Which proteins are transferred to the membrane?

# Antibodies used to detect specific protein

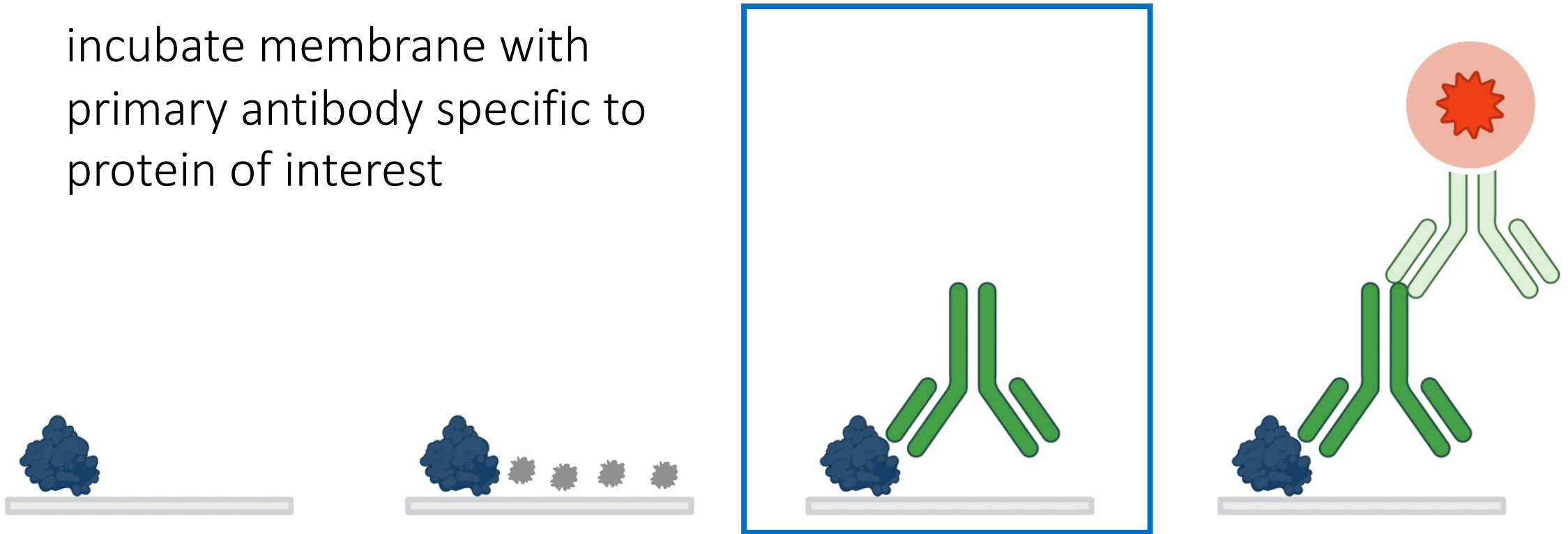


- How does blocking buffer prevent non-specific binding of antibodies?



# Antibodies used to detect specific protein

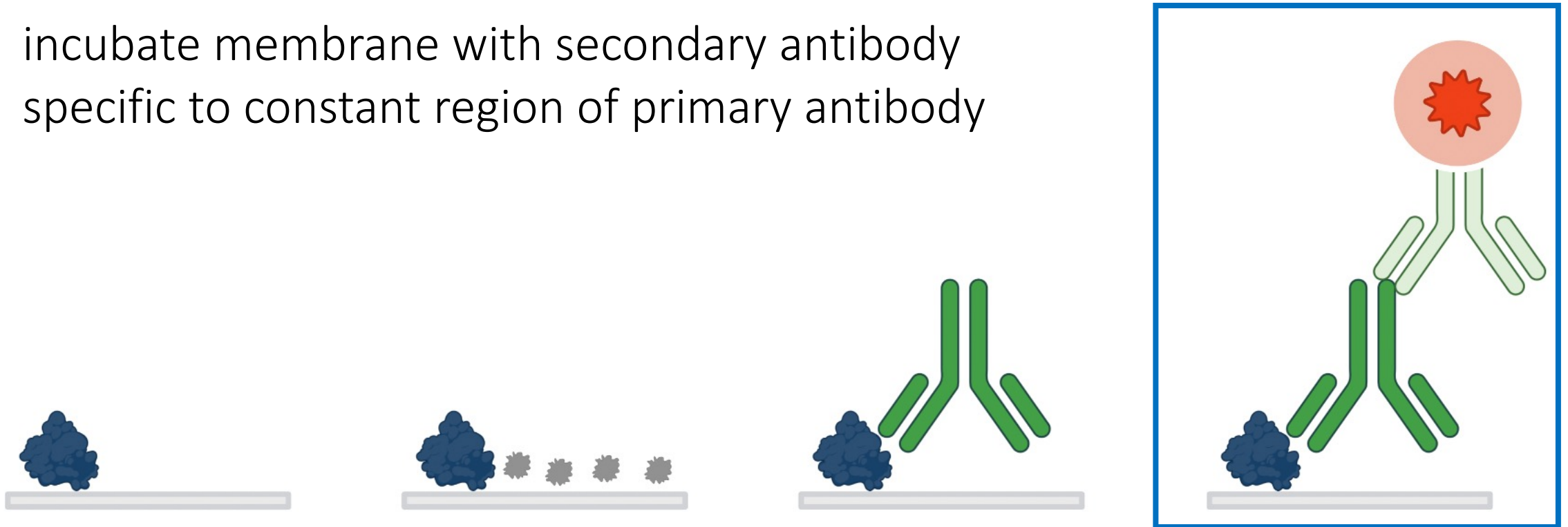
incubate membrane with primary antibody specific to protein of interest



- Why are we using two primary antibodies? For what does each probe?

# Antibodies used to detect specific protein

incubate membrane with secondary antibody  
specific to constant region of primary antibody



- Why use secondary antibodies (rather than labeled primary antibodies)?

## For today...

- Use laboratory time to complete Data summary slide for protein purify and concentration results with your partner in Breakout room

## For M1D6...

- **With your laboratory partner**, draft methods section
- Submit write-up from meeting with Comm Lab fellow

# Notes on methods section...

Include **enough information to replicate** the experiment

- Cite manufacturer for supplies / equipment (Company)
- Be concise and clear in your description

Use subsections with **descriptive titles**

- Put in logical order, rather than chronological order
- Begin with topic sentence to introduce purpose / goal of each experimental procedure

Use clear and concise **full sentences**

- NO tables or lists, all information should be provided in full sentences and paragraphs
- Write in passive voice and use past tense

Use the most **flexible units**

- Write concentrations (when known) rather than volumes

**Eliminate 20.109 specific language and obvious details**

- Example “labeled Row A, Row B...”
- Do not include details about tubes and water!
- Assume reader has some biology experience
- Include parts of the protocol that the teaching faculty completed, but do not say “completed by teaching faculty.”

# How can you improve this example?

DNA was cut to check insert. Enzymes were used for single and double

digest then run on gel made by adding 1 g of agar to 100 mL of water.

Gel was imaged on a gel box.