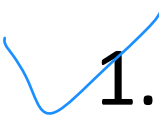


M1D2: Clean and cut DNA

9/18/15

- 
1. Visit from BE Communications Instructors
 2. Lab quizzes- Why?!
 3. Preview of M1D3 homework
 4. Discuss PCR product design and purification
 5. Review restriction enzyme digests

Quiz on M1D4

- Material from lectures (M1D1-M1D3): ~60%
- Material from prelab/wiki (M1D1-M1D3): ~40%

Why do we give quizzes?

1. To refresh your memory about the long-term experiment you are performing
2. To provide you with an opportunity to show your technical knowledge decoupled from your communication skills

Attending lecture is



Homework due M1D3

Due M1D3 [\[edit\]](#)

1. Sketch the expected product from the PCR you performed on M1D1.
 - You may work on paper or electronically. Either way, prepare a schematic rather than detailing each base.
 - Clearly indicate the 5' and 3' end of each DNA strand.
 - Be sure to reflect every new feature that you have introduced (e.g., restriction site) or deleted.
 - What is the expected size of the PCR product?
2. Following the directions in Part 4 of M1D2, prepare a plasmid map in ApE of the clone you are trying to create in lab. Print the graphic map with all singly present restriction sites shown.
 - Hint: You may choose to show fewer restriction sites in your Module 1 summary.
 - Using your map, calculate the fragment sizes expected for each double digest below. **Please show your work.**
 - *EcoRV* and *XbaI*
 - *BamHI* and *XhoI*
 - Use the skills you learned from the BE Communications instructors to write a title and caption for your plasmid map figure.
3. In Module 2, you will document your experiments in a written methods section that will be part of a larger report. To help you prepare, as well as give you feedback early on, you will draft portions of the Module 1 methods. For this assignment, write a draft of the Methods concerning PCR and DNA digestion. Be sure to read the Materials and Methods section [guidelines at this link](#) before you begin; doing so may save you some effort.

Methods section tips

- Use subsections with descriptive titles
 - Put in logical order
 - Begin with topic sentence
- Use clear and concise full sentences
 - Avoid tables and lists
- Use the most flexible units
 - Write concentration rather than volume

What no why

★ when possible ★

Some methods practice...

EGFP

more than 750h

PCX-EGFP

[x2] D32N (CAT TAG)

"A plasmid with ~~template DNA~~ (5ng) and primers were mixed with 20 ~~uL~~ of 2.5X ~~Master Mix~~ in a PCR tube. Water was added to 50 uL. A tube without template was prepared and labeled control."

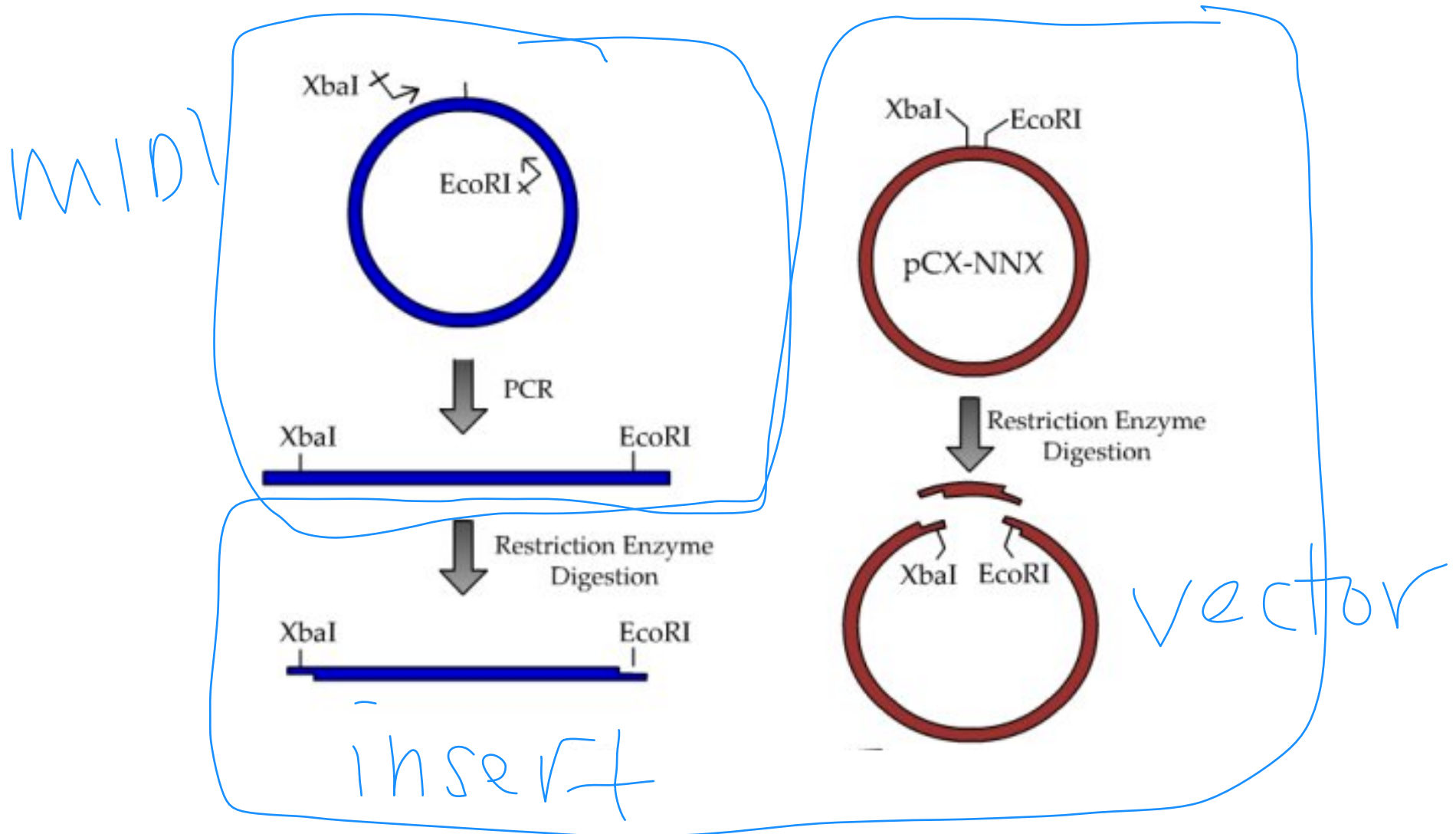
A NTC was included.

(SPrime, City, St)

MOD1 major assignments

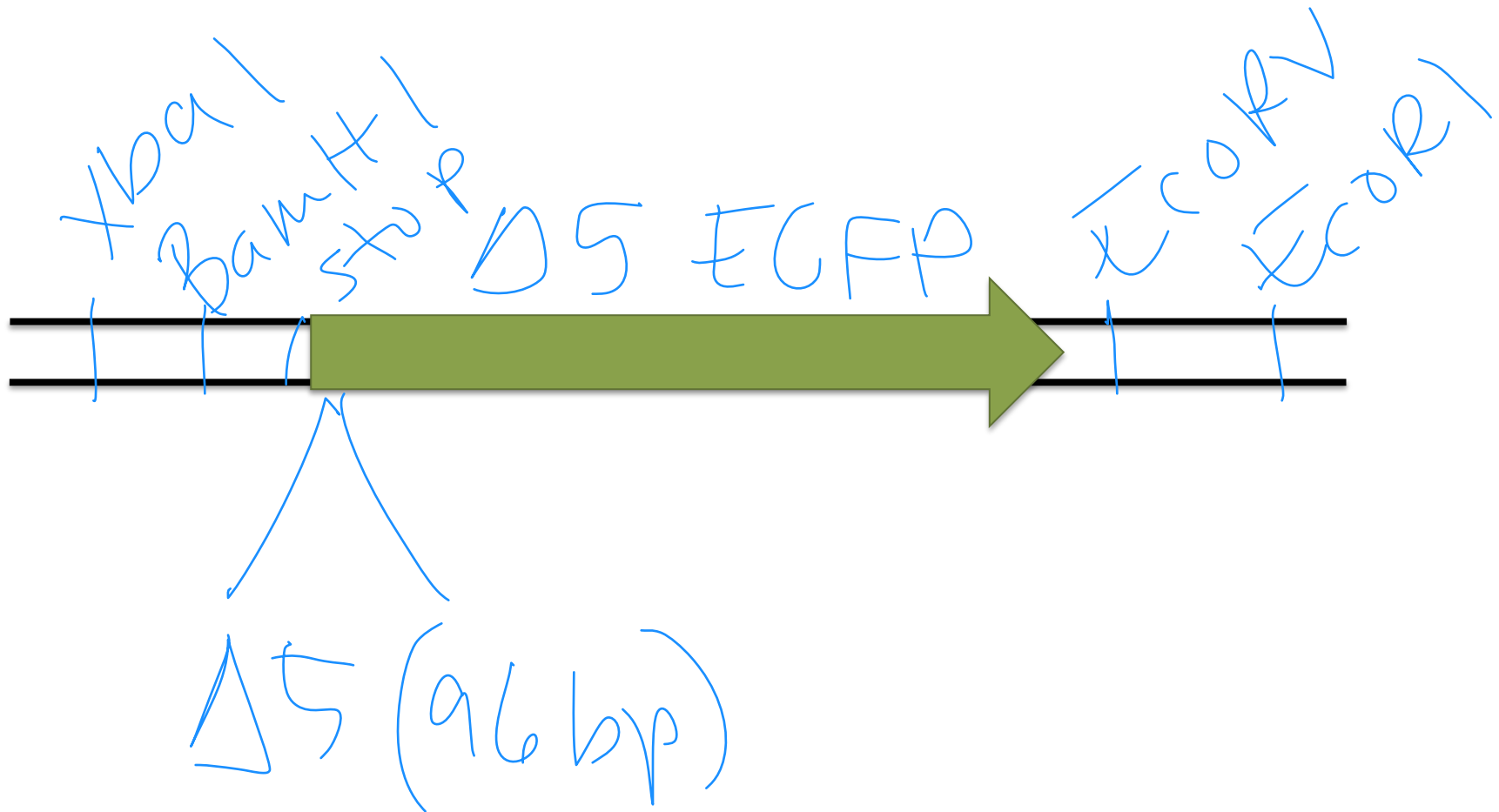
- DNA engineering summary (15%)
 - Title and abstract
 - Background/motivation
 - Results and interpretations (figures with captions)
 - Implications/future work
- DNA engineering mini-presentation (5%)
 - Abbreviated oral overview of summary
 - ‘elevator pitch’ or ‘flask talk’

Using PCR amplification and REs to build $\Delta 5$ EGFP construct



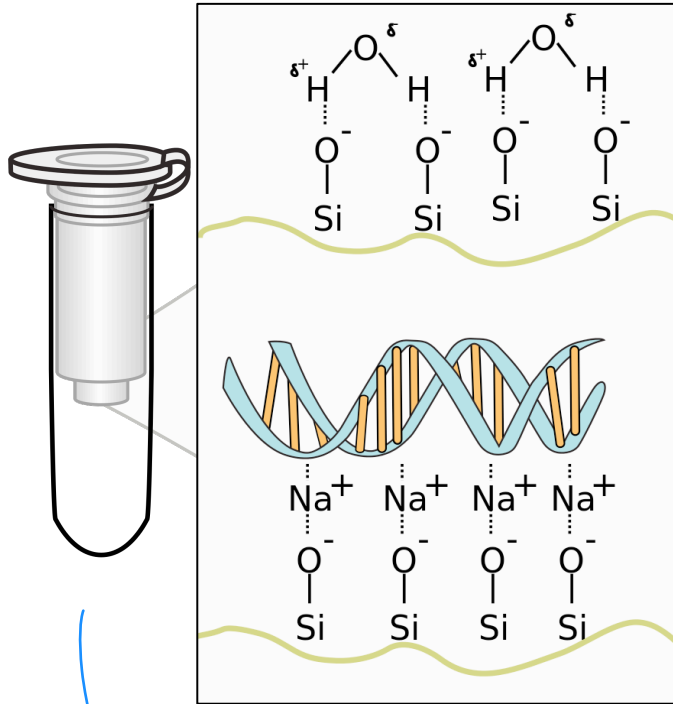
From last time...

Did your primer design meet all of the guidelines?



Purification of PCR product

Qiagen



> 100 bps

1) Bind: PB
chaotropic salts
guanidine HCl

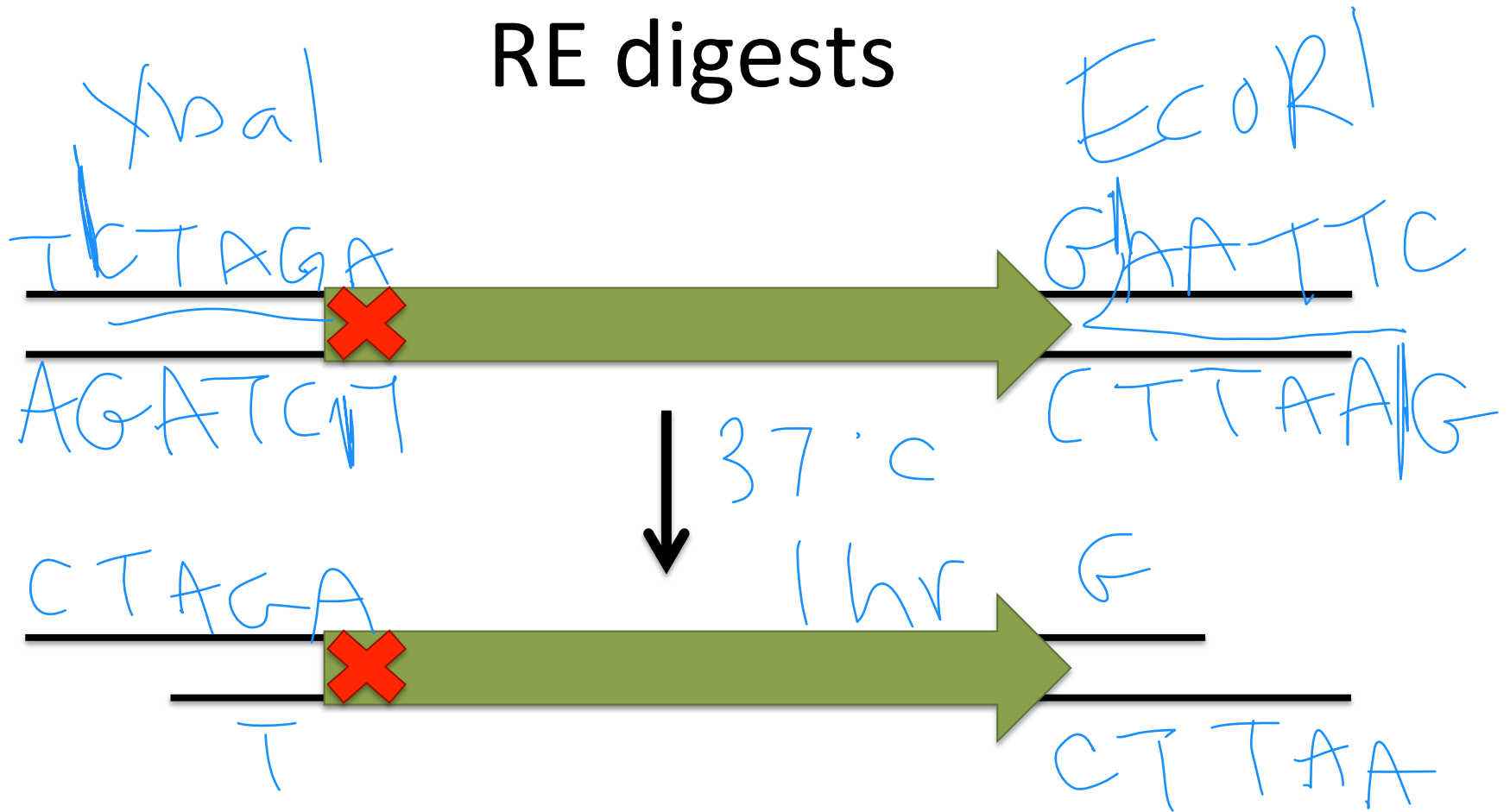
↓ pH

2) Wash: EtOH^{PE}

3) Elution: TE
low salt ↑ pH

Keep track of your DNA!!

RE digests



What about pCX-NNX digest product?

Today

- Clean and digest your PCR product
- Digest pCX-NNX vector
- ApE (plasmid editor) practice

- Networking event
 - Thursday, September 24 at 5:30-7p
 - First floor lobby of the Koch

