

TEACHING THE  
GENOME  
GENERATION™

*PREP FOR SEQUENCING PROTOCOL*

# PREREQUISITES & GOALS

## PREREQUISITES

Prior to implementing this lab, you should understand:

- The central dogma of how DNA bases code for mRNA and then for proteins
- How DNA samples were collected and prepared for PCR
- The steps that occur during the process of polymerase change reaction (PCR), as well as the reagents used
- The variants of ACTN3, CYP2C19, and/or TAS2R38
- The connection between the bands on the gel electrophoresis and the sequential gene variants
- The purpose of using ExoSAP-IT
- The purpose of the PREP FOR SEQUENCING PROTOCOL is to prepare PCR-amplified DNA samples for Sanger sequencing

## LEARNING GOALS

Prepare amplified DNA products for Sanger sequencing.

## NOTES

The **ExoSAP** system uses two enzymes to prepare amplified PCR products for sequencing:

**Exonuclease I (Exo)** digests residual single-stranded DNA primers and any extraneous single-stranded DNA fragments produced during the PCR process.

**Shrimp Alkaline Phosphatase (SAP)** dephosphorylates remaining dNTPs (free nucleotides) from the PCR product so they do not interfere with the sequencing reaction

# MATERIALS

## REQUIRED LAB MATERIALS

Ice bath or crushed ice	Mini-microcentrifuge
Markers for labeling	Thermal cycler
Amplified DNA samples from the PCR PROTOCOL	Molecular biology grade water
Micropipettors & tips (size P20)	(F)orward sequencing primer (4 $\mu$ M)
0.2 mL tubes in strips of 4	(R)everse sequencing primer (4 $\mu$ M)
1.5 mL tubes	Invitrogen ExoSAP-IT enzyme (on ice)
Tube holders/racks	

## WORKSTATION NEEDS

*These materials should be at each workstation.*

Micropipettors and tips  
0.2 mL tubes in strips  
1.5 mL tubes  
Molecular biology grade water  
(F)orward and (R)everse primers  
Tube holders and markers for labeling  
Crushed ice/ice bath  
Exo-SAP IT enzyme (on ice)  
Amplified DNA samples

# PROCEDURE

## □ STEP 1

Obtain 0.2 mL strip tubes and label them with the DNA sample numbers/letters.

NOTE: Do not use the negative control sample in this procedure, only samples that demonstrated positive amplification during gel electrophoresis should be processed.

## □ STEP 2

Using the P20 micropipettor, transfer 5  $\mu$ L of the amplified DNA samples from the PCR PROTOCOL to the new tubes.

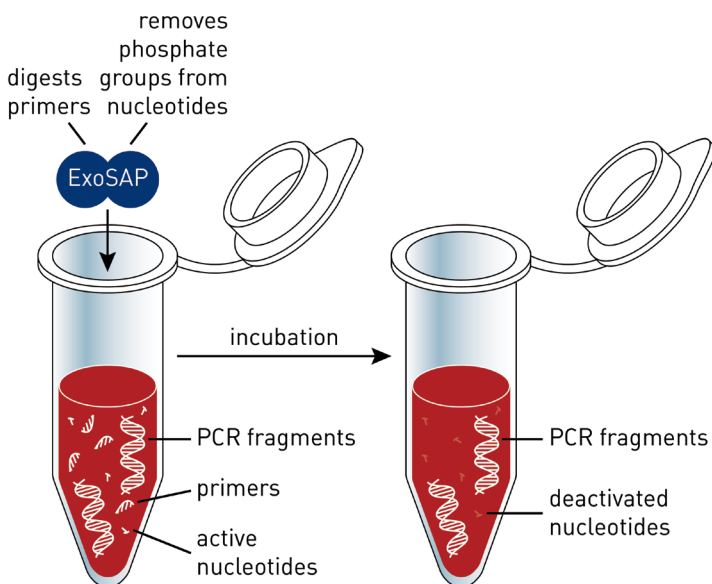
## □ STEP 3

Using the P20 micropipettor, add 2  $\mu$ L of ExoSAP-IT to the new tubes.

NOTE: Enzymes must be kept on ice at all times.

## NOTES

A large grid of 20 columns and 30 rows of small grey dots, intended for taking notes.



## STEP 4

Tightly cap the tubes and flick the tube gently to mix.

## STEP 5

Spin the tubes briefly in the mini-microcentrifuge to collect the solution.

## BREAK POINT IF NEEDED

Expected result is to have one tube per DNA sample with 7  $\mu$ L of reaction solution.

## STEP 6

The thermal cycler should be programmed with the digestion protocol.

### EXOSAP: Digests free nucleotides and primers

1. Digestion                    37 °C   15 min.
2. Protein degradation   85 °C   15 min.
3. Final hold                    4 °C   forever

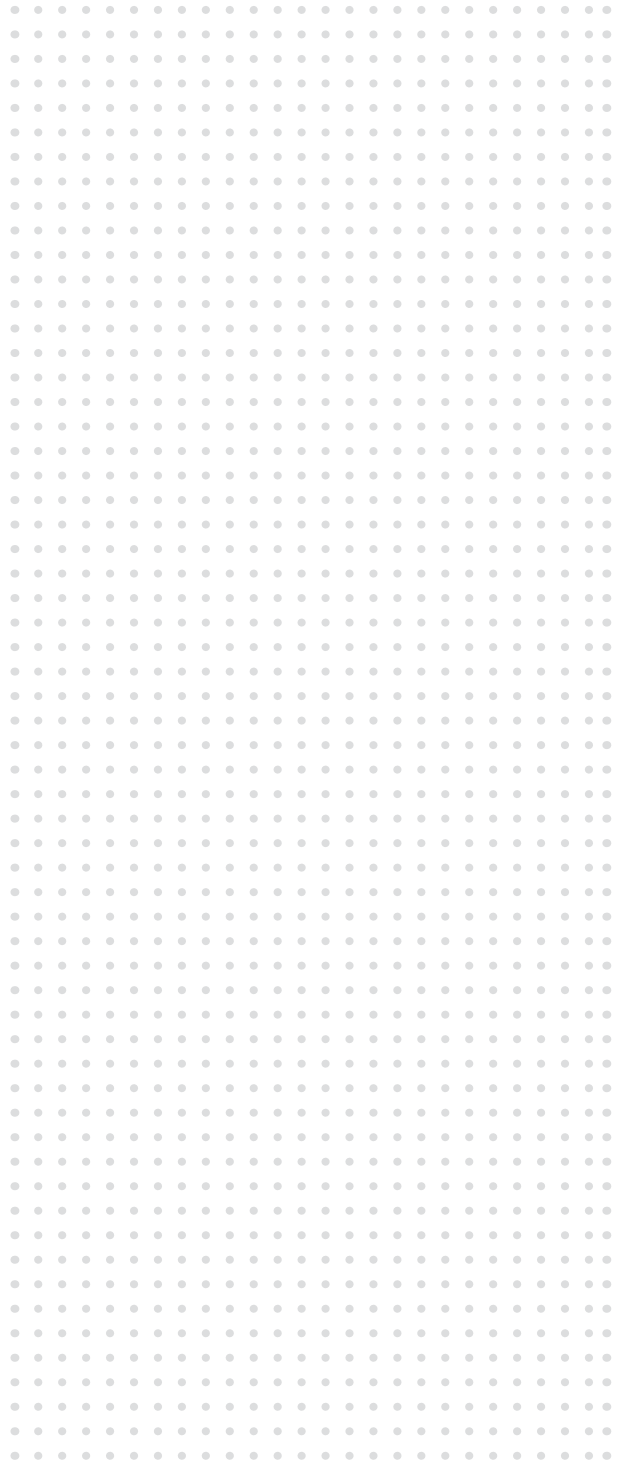
## STEP 7

Consult your teacher on proper use of the thermal cycler provided.

## BREAK POINT

The reaction will proceed for 30 minutes.

## NOTES



## STEP 8

Remove the samples after the protocol is complete, stop the program and turn the machine off.

### BREAK POINT IF NEEDED

Expected result is to have one tube per DNA sample with 7  $\mu\text{L}$  of reaction solution. Nothing should look different about the solution after the reaction.

## STEP 9 **CRITICAL STEP**

Obtain two new 0.2 mL strip tubes per DNA sample to be sequenced.

## STEP 10 **CRITICAL STEP**

Label one tube as the forward sequencing reaction and one as the reverse sequencing reaction for each sample.

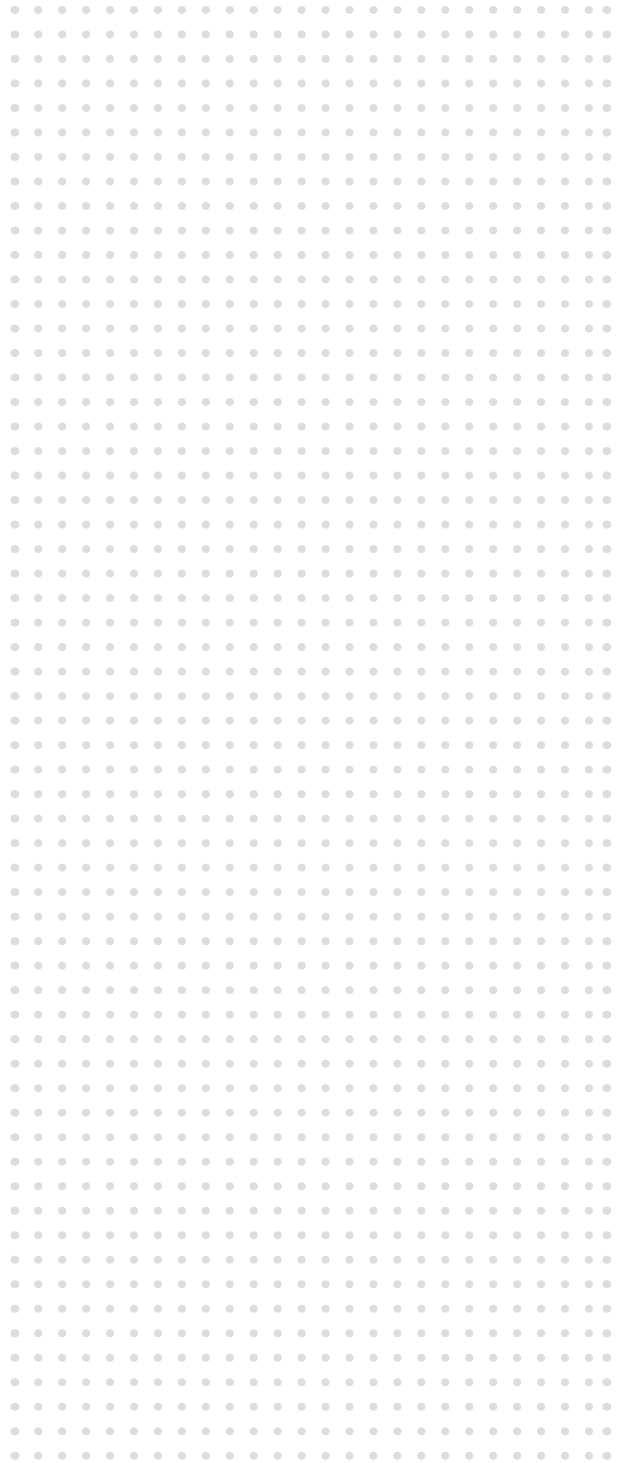
EXAMPLE: 4-F would be the label for DNA sample #4 to be sequenced with the Forward primer.

## STEP 11

Using the P20 micropipette, add the following to the FORWARD reaction tube:

- 10  $\mu\text{L}$  of molecular biology grade water
- 3  $\mu\text{L}$  of ExoSAP-IT/DNA mix that has been incubated
- 2  $\mu\text{L}$  of (F)orward sequencing primer

## NOTES



## STEP 12

Using the P20 micropipettor, add the following to the REVERSE reaction tube:

- 10  $\mu$ L of molecular biology grade water
- 3  $\mu$ L of ExoSAP-IT/DNA mix that has been incubated
- 2  $\mu$ L of (R)everse sequencing primer

## STEP 13

Tightly cap the tubes and flick the tube gently to mix.

## STEP 14

Spin the tubes briefly in the mini-microcentrifuge to collect the solution.

Expected result is to have two tubes per DNA sample with 15  $\mu$ L of reaction solution.

## STEP 15

Follow the final sample labeling, shipping and packaging directions dictated by the sequencing company.

The DNA sample is now ready for sequencing.

## NOTES

