# ENOME 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

Markers for labeling

Amplified DNA samples from the PCR PROTOCOL

Micropipettors & tips (size P20)

0.2 mL tubes in strips of 4

1.5 mL tubes

Tube holders/racks

Mini-microcentrifuge

Thermal cycler

Molecular biology grade water

(F)orward sequencing primer (4  $\mu$ M)

(R)everse sequencing primer (4  $\mu$ M)

Invitrogen ExoSAP-IT enzyme (on ice)

#### **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

**NOTES** 

## **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.

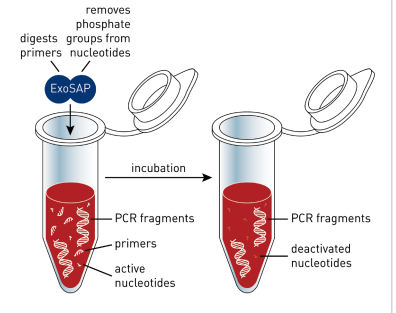
#### □ STEP 2

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** 

		_	
. C.		ט	1.
	ΙГΙ	Г	4
	-		

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

#### □ STEP 5

Spin the tubes briefly in the minimicrocentrifuge 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  $85\ ^{\circ}\text{C}$   $15\ \text{min}.$ 

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.

□ STEP 8	NOTES
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 $\mu$ 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:	
<ul><li>10 μL of molecular biology grade water</li><li>3 μL of ExoSAP-IT/DNA mix</li></ul>	

that has been incubated

 $\square$  2 µL of (F)orward sequencing primer

□ STEP 12	NOTES
Ising the P20 micropipettor, add the bllowing to the REVERSE reaction tube:	
10 μL of molecular biology grade water	
<ul> <li>3 μL of ExoSAP-IT/DNA mix</li> <li>that has been incubated</li> </ul>	
2 μL of (R)everse sequencing primer	
□ STEP 13	
ightly cap the tubes and flick the tube gently to mix.	
□ STEP 14	
pin the tubes briefly in the mini- nicrocentrifuge to collect the solution.	
eaction solution.	
□ STEP 15	
follow the final sample labeling, shipping and packaging directions dictated y the sequencing company.	
he DNA sample is now eady for sequencing.	