Source: [KBBiologyMasterIndex]

# 1 | SNP

**Single Nucleotipe Polymorphism**: pairs of single-base pair variations of genes that are not uncommon (>1%), and hence not quite considered a mutation.

SNPs could be in any part of the genome: sometimes in a gene, but more often in a non-coding region.

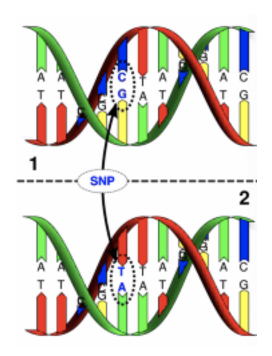


Figure 1: Pasted image 20210505134813.png

These could do absolutely nothing, but they could also create large trait differences (sickle-cell is a SNP). The DNA by mail kits are actually just looking for SNPs!

Multiple SNPs come together to form a "Haplotype"; they are SNPs that are fairly close together that frequently occur in sequence (like the chain of three SNPs A something something G something T would be a haplotype.)

People of the same haplotype are likely of same ancestry.

## 1.1 | SNP Guidelines

- · No medical SNPs; so avoid SNPs that could be associated with heavy medical information
- No SNPs that could potentially give you PTSD

#### 1.2 | the Reaction

- Each student's PCR reaction will only amplify 500-800 nucleotide region
- Find SNPs that have specific traits, but most SNPs are po

#### 1.3 | Polymerase Chain Reaction (PCR)

A reaction by which the DNA replication process is attempted to be replicated out of the cell in preparation to sequence one's DNA

PCR provides a way of creating lots of copies of a small strand of DNA

#### 1.3.1 | Reaction Need

- The DNA Portion (duh)
- · A buffer to host the liquid
- · Primers Bit of RNA to help DNA polymerase get going
- DNA Polymerase The Enzyme to Copy the DNA => Typically, the TAQ Polymerease is used because it is from a bacterial that's quite resistant to heat
- · DNA Nucleotides to build the replicated DNA

## 1.3.2 | Steps

- 1. Denaturation
  - 1. Cook the DNA ("denature it"/heat it) to seperate the two strands of the DNA
- 2. Annealing
  - 1. Cool the Cooked DNA, and dump in the primers
  - 2. The specific temperature in this step is important because it controlls where the primer binds, and hence which part of the DNA you want to amplify
- 3. DNA Synthesis => make more copies of the DNA
  - This needs a slightly higher temperature than annealing, but which one depends on what polymerase you used
- 4. Go to step 1

Note: the DNA polymerase may run over, so we just keep repeating this and it will slowly become more successful as the DNA polymerease repeats its work repeatedly.

BTW: if you read carefully, you will realize that you made TWO DNA double-strands out of a single double strand. So exponential growth baybee and then you could get lots of copies of that region.

# 1.3.3 | But, why?

In order to have enough copies for gel electroprorisis (like, the rRT-PCR COVID test) or, for things like crime forensics, you need many copies of the same DNA for a clearly detectable result for analysis that may have a high threshold.