EVERATION

Variation in the Genome



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Contents

Variation in the Genome Module Overview	
Introduction & Theme	3
Learning Outcomes	3
Lesson & Activity Descriptions Establishing Genotype-Phenotype Associations	4
Lessons & Activities	5
Establishing Genotype-Phenotype Associations: Paper-Based	5
Establishing Genotype-Phenotype Associations: Spreadsheets	27
Implementation Strategies	47
NGSS Alignments	49
Supporting Materials	50
Feedback	51

Variation in the Genome Module Overview

Introduction & Theme

This module aims to engage students in performing biologically meaningful calculations and basic statistical analysis. Through the broad lens of Genome Wide Association Studies (GWAS), students will learn how researchers use statistics to establish associations between genetic variants and specific traits. Students will then be introduced to a simplified dataset which will enable them to investigate the connection between bitter taste perception ability and genetic variants in the *TAS2R38* gene. Students will perform calculations and statistical analysis using the dataset population, interpret the outcomes, determine to what extent they can generalize their results, and finally consider why studies that include millions of genetic variants across larger populations require computational tools and automation.

Learning Outcomes

Essential Question

How can we determine associations between genetic variants and complex traits?

Enduring Understandings

- Genetic variations can impact sensory perception (a polygenic trait) in humans.
- Polygenic traits are influenced by multiple genes and variants, as well as the environment.
- Human genomic variation is something to celebrate.

Skills

Data analysis and interpretation; computation (math skills); basic statistical analysis; pattern recognition; constructing explanations

Lesson & Activity Descriptions

Establishing Genotype-Phenotype Associations

In this activity, students work through calculations and statistical analysis (chi-square) to establish an association between a specific genetic variant in the *TAS2R38* gene and bitter taste perception ability. Students are provided a real dataset with the *TAS2R38* genotypes and bitter taste perception phenotypes from 60 individuals.

In the first part of the activity, students count the number of alleles associated with each of the three bitter taste perception phenotypes: super taster, taster, and non-taster. Guided questions prompt students to identify any patterns in the data and consider what additional analysis might be useful.

In the second part, students calculate allele frequencies based on the allele count data. Students again identify patterns and consider which representation of the data is easiest for them to understand.

In the third and fourth part, students set up and perform a chi-square analysis. First, students calculate expected counts, which represent a random distribution of alleles across the phenotypes. Students then compare their actual counts to their calculated expected counts use a chi-square test to determine if there is a difference between their actual data and a random distribution of alleles.

Finally, students draw conclusions based on their calculations. Through guided questions, students consider whether they have established an association or causation and how broadly they could generalize their results. Students also connect their experience performing statistical analysis back to the concept of GWAS.

There are two versions of this activity: a paper-based version and a spreadsheet version. The two versions follow same general structure and cover the same biological and mathematical concepts. The primary difference is whether students perform calculations and document their results by hand or within a spreadsheet. You can either select one version for students to complete, or you can mix and match sections of each version. For example, students could complete Part 1 of the paper-based version then switch to the spreadsheet version for Parts 2-5.

Paper-Based

In the paper-based version, students perform all calculations by hand and write their results into provided printed tables. Students are provided the p-value result for the chi-square test in the final part of the activity.

Spreadsheets

In the spreadsheet version, students (or teachers) can opt to perform calculations by hand or using spreadsheet formulas. Students enter all results into tables within the spreadsheet. Students use a built-in function to obtain the p-value result for the chi-square test in the final part of the activity.

Lessons & Activities

Establishing Genotype-Phenotype Associations: Paper-Based

Introduction

What is a GWAS?

Did you know that most of our traits and many human diseases are not caused by a single genetic variant, but still are influenced by genetics?

For example, our heart health is influenced by both genetics and our environment. An individual with a variant in the *CDKN2A* gene might have an increased risk of developing heart disease, but whether that individual develops disease is also dependent on other genetic variants and on their environment. Even someone who does not have that variant in *CDKN2A* can develop heart disease, especially if factors in their behavior, such as smoking, or environment, such as exposure to air pollution, lead to increased risk.

For complex traits, like our heart health, how do scientists identify associations between genetic variants and a trait of interest when there are so many other variables? One tool that scientists use is called a Genome Wide Association study, or GWAS (pronounced *GEE-wahs*).

For a GWAS, scientists start by collecting genetic data from a large population with varied phenotypes for a particular trait or disease. Then the scientists look at millions of variants and determine if any of those variants are associated with a particular phenotype of that trait. Typically, scientists will look at single nucleotide polymorphisms, or SNPs (pronounced *snips*). SNPs are a type of variant where just one nucleotide varies from person to person.

For example, to try to find associations between genetic variants and heart disease, scientists would recruit a large group of people to participate in their study, some of whom have heart disease and others who do not. The scientists look at all of the participants' genetic variants and see if there are any variants that are more common in the group with heart disease than they are in the group without heart disease. Then, they perform statistical tests to determine if those variants are more common in the heart disease group than would be expected just by chance.



Figure 1. Representation of potential outcomes of a GWAS study. In this example, the variants SNP 1 and SNP 2 are found in approximately equal percentages of the disease and non-disease populations. SNP 3, however, is more common in individuals with disease. (From <u>"Genome-Wide Association Studies Fact Sheet"</u> by the <u>National Human Genome Research Institute</u>).

The TAS2R38 Gene

In this activity, you will look at an example of the types of calculations involved in a GWAS to demonstrate an association between bitter taste perception and variants in the *TAS2R38* gene.

The Taste 2 Receptor Member 38 (*TAS2R38*) gene produces the TAS2R38 protein, which functions as a receptor to perceive a wide range of bitter compounds. Bitter taste receptors, including TAS2R38, are proteins found on the taste cells of the tongue. Bitter taste perception is affected by multiple genes and by environmental factors, such as the foods we are exposed to and have regular access to.

There are three bitter taste perception phenotypes which you will investigate today: non-taster, taster, and super taster.

- Non-tasters do not perceive bitter taste.
- Tasters perceive bitter taste, although the taste may be mild.
- Super tasters perceive bitter taste, and the taste is typically strong.

While we group people into these three phenotypes, these phenotypes are subjective and correspond to a range of actual bitter taste sensing levels.

Part 1: Allele Counts

Background

For our investigation, you will start with one variant in *TAS2R38*. The *TAS2R38* variants are single nucleotide polymorphisms, or SNPs (pronounced *snips*). This means that just one nucleotide varies from person to person. There are three common SNPs in *TAS2R38*, which each occur at different locations in the gene.

- SNP 1: At this variant location, the nucleotide is usually either a guanine (G) or a cytosine (C).
- SNP 2: At this variant location, the nucleotide is usually either a thymine (T) or a cytosine (C).
- SNP 3: At this variant location, the nucleotide is usually either an adenine (A) or a guanine (G).

We have collected genetic data from a group of individuals, recording both their *TAS2R38* genotype and bitter taste perception phenotype. The samples were provided voluntarily, meaning that each person agreed to provide a sample. The samples were also anonymized, meaning they were labeled with random sample names so that they cannot be traced back to the person who provided the sample.

Recall that everyone has two copies, or alleles, of each gene. For example, for the first variant (SNP 1), individuals can have a C on both copies, a G on both copies, or one C and one G.

Activity

To start your analysis, you will count how many of each allele of your assigned SNP are associated with each tasting phenotype.

- Column 1 Column 2 Column 3 Column 4 **Sample Name** SNP 1 (G/C) Alleles Phenotype Row 1 Row 2 Row 3 Row 4 Row 5 * *****... *
- 1. Find **Table 1. Allele Data** at the end of this handout. **Table 1** has 4 data columns and 60 data rows.

Figure 2. Image of Table 1 with rows and columns numbered for reference. The full table has 60 rows—only 5 are shown here.

a. What type of data is in columns 2 and 3? What does that data represent?

b. What type of data is in column 4? What does that data represent?

c. What does each row represent?

d. Some of the table entries are blank. What do you think this means about that data?

e. How many individuals are included in the dataset? How many alleles are included in the dataset?

Now that you are familiar with the raw data, you can count the number of G and C alleles that are associated with each of the bitter taste perception phenotypes. You will enter this information into one of the analysis tables, which is labeled **Table 2. Allele Counts**.

	Column 1	Column 2	Column 3	Column 4
	Non-taster	Super taster	Taster	Total
G				
с —				
Total	•			

Table 2 has 3 data rows and 4 data columns, <u>not</u> including the row and column labels.

Figure 3. Image of Table 2 with rows and columns numbered for reference.

- 2. Find **Table 2. Allele Counts** and use the table to enter the following questions.
 - a. What do columns 1, 2, and 3 represent?
 - b. What does column 4 represent?
 - c. What do rows 1 and 2 represent?
 - d. What does row 3 represent?
 - e. Into which cell would you enter the data for the number of C alleles associated with the Taster phenotype? Put an X in the correct cell in the example table below.

Table 2. Allele Counts

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

f. Into which cell would you enter the data for the number of G alleles associated with the Non-taster phenotype? Put an X in the correct cell in the example table below.

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

Table 2. Allele Counts

Now that you are familiar with **Table 2**, you can count the alleles in each phenotype group and enter the counts data into **Table 2**.

- 3. For each phenotype, count the number of G alleles and enter the data into the correct column of the **G** row of **Table 2**. Be sure to include data from both alleles for each individual!
- 4. For each phenotype, count the number of C alleles and enter the data into the correct column of the **C** row of **Table 2**. Be sure to include data from both alleles for each individual!
- 5. Add the allele counts across each row and enter those values into the **Total** column.
- 6. Add the allele counts down each column and enter those values into the **Total** row.

After completing all the calculations, use your completed **Table 2** to answer the following questions.

- 7. How many G alleles are associated with each phenotype?
 - a. Non-taster:
 - b. Super taster:
 - c. Taster:
- 8. How many C alleles are associated with each phenotype?
 - a. Non-taster:
 - b. Super taster:
 - c. Taster:

9. What patterns do you notice in the data?

10. Based on this data alone, without performing additional calculations, do you think this variant in *TAS2R38* is associated with bitter tasting phenotype? Why or why not?

11. How could you use additional data collection, analysis, or representation to better support your hypothesis?

Part 2: Allele Frequency

Note: If your students are not yet familiar with statistics, this section provides an opportunity to do additional calculations and data analysis.

Background

Next, you can use the allele count data in **Table 2** to calculate allele frequencies. Allele frequency refers to the distribution of alleles across a population. For example, the allele frequency of the G allele in the total population is the fraction, or percentage, of the total alleles in the population that are G alleles. The equation we will use for allele frequency is:

Allele Frequency =
$$\frac{v}{2N}$$

Where the variable v is the number of each allele in the dataset, and the variable N is the total number of diploid individuals in the dataset.

In Part 1, you counted the number of each allele in the dataset, so **Table 2** contains the values for v. Additionally, since N represents the number of diploid individuals and each individual has two alleles, 2N is equal to the total allele count. So, we can also write the allele frequency equation as:

$$Allele Frequency = \frac{Allele Count of a specific allele in the population}{Total Allele Count in the population}$$

For example, in our population of 60 individuals, we have 106 known alleles in total (**Table 2**, **Total** row, **Total** column). Of those 106 alleles, 65 are G alleles (**Table 2**, **G** row, **Total** column). We can calculate the allele frequency of the G allele in our total population as follows:

Total G Allele Frequency =
$$\frac{Allele Count of G allele in the population}{Total Allele Count in the population} = \frac{65}{106} = 0.613$$

If we look at just the Taster population, there are 35 G alleles (**Table 2**, **G** row, **Taster** column). However, because we are only looking at the Taster population, our total number of alleles also changes. There are 30 Tasters, corresponding to 60 alleles (**Table 2**, **Total** row, **Taster** column). We can calculate the allele frequency of the G allele in the Taster population as follows:

$$Taster \ G \ Allele \ Frequency = \frac{Allele \ Count \ of \ G \ allele \ in \ Taster \ population}{Total \ Allele \ Count \ in \ the \ Taster \ population} = \frac{35}{60} = 0.583$$

Activity

First, practice calculating allele frequency using the data for the number of C alleles in the Taster population.

 Before you begin, find the number of C alleles associated with the Taster phenotype in Table 2. Allele Counts. What cell in Table 2 contains the number of C alleles associated with the Taster phenotype? Put an X in the correct cell in the example table below.

Table 2. Allele Counts

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

- 2. What is the allele count of C alleles associated with the Taster phenotype?
- 3. Now, use the allele count to complete the calculation below. Round your answer to three decimal places.

You will use **Table 3. Allele Frequencies** to record the results from your allele frequency calculations. Before completing the remaining calculations, get familiar with the format of **Table 3**.

	Colu	mn 1	Colu	mn 2	Colu	mn 3	Colu	mn /
	Colu		Colu		Colu		Colui	
	Non-tast	er	Super ta	ster	Taster		Total Po	pulation
G								
c —								
Total		7						

Table 3 has 3 data rows and 4 data columns, <u>not</u> including the row and column labels.

Figure 4. Image of Table 3 with rows and columns numbered for reference.

4. Find Table 3. Allele Frequencies and use the table to enter the following questions.a. What does column 4 represent?

b. What does row 3 represent? Predict the value of each cell in row 3. Hint: In allele frequency, we are calculating the frequency of each allele within the population. What should be the frequency of both alleles combined in the population?

c. Into which cell would you enter the data for the frequency of G alleles in the total population? Put an X in the correct cell in the example table below.

Table 3. Allele Frequency

	Non-taster	Super taster	Taster	Total Population
G				
С				
Total				

d. Into which cell would you enter the data for the frequency of C alleles associated with the Super taster phenotype? Put an X in the correct cell in the example table below.

Table 3. Allele Frequency

	Non-taster	Super taster	Taster	Total Population
G				
с				
Total				

Now you can calculate the allele frequencies for each phenotype and enter the data into Table 3.

- 5. For each phenotype and for the total population, calculate the allele frequency of the G allele and enter the data into the correct column of the **G** row of **Table 3**.
- 6. For each phenotype and for the total population, calculate the allele frequency of the C allele and enter the data into the correct column of the **C** row of **Table 3**.
- 7. Add the data down each column and enter those values into the **Total** row.

After completing all the calculations, use your completed **Table 3** to answer the following questions.

- 8. What is the G allele frequency for each of the following phenotype groups:
 - a. Non-taster:
 - b. Super taster:
 - c. Taster:
 - d. Total Population:
- 9. How is the C allele frequency for each of the following phenotype groups:
 - a. Non-taster:
 - b. Super taster:
 - c. Taster:
 - d. Total Population:
- 10. What patterns do you notice in the data?

11. The data in Table 2 and Table 3 come from the same raw data but show different ways of analyzing and representing that data. Which set of analyzed data, allele counts (**Table 2**) or allele frequency (**Table 3**), is easier for you to understand? Why?

Part 3: Expected Counts

Note: Part 3 and Part 4 introduce some basic statistics and are intended for AP students or for students with a statistics background. With some additional background information and support from the teacher, this portion of the activity could also be used to introduce some statistical concepts to students who are not as familiar.

Students can still complete Part 3 even if you do not intend for them to complete Part 4.

Background

In **Table 2. Allele Counts** and **Table 3. Allele Frequency**, we have data about the distribution of the different *TAS2R38* SNP 1 alleles across the different bitter tasting phenotypes. Now we can ask the question: is there an association between genetic variant and phenotype?

To answer this question, we can use a statistical test called a Pearson's chi-square test of independence. The Pearson's chi-square test of independence evaluates whether observations of measures on two variables (in our case, allele and phenotype) are independent of each other. The chi-square test requires us to compare our actual allele counts to **expected counts**.

Expected counts represent what the allele counts would be if there was no association between the variant and phenotype. In other words, **expected counts** represent a random distribution of alleles across the phenotypes, such that <u>each phenotype has the same allele frequencies as the total population</u>.

Example Calculation

For example, let's look at the expected count of G alleles associated with the Non-taster phenotype.

The frequency of G alleles in the total population is 0.613. Importantly, expected count calculations assume that the total number of alleles associated with each phenotype remains the same. In our population, there are 30 total alleles associated with the Non-taster phenotype.

1. The frequency of <u>G alleles</u> in the <u>total population</u> (0.613) is in **Table 3. Allele Frequency**. What cell in **Table 3** contains the frequency of G alleles in the total population? Put an X in the correct cell in the example table below.

	Non-taster	Super taster	Taster	Total Population
G				
С				
Total				

Table 3. Allele Frequency

 The total number of alleles associated with the <u>Non-taster</u> phenotype (30) is in **Table 2. Allele** Counts. What cell in **Table 2** contains the total number of alleles associated with the Non-taster phenotype? Put an X in the correct cell in the example table below.

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

The expected count of G alleles for the Non-taster population represents how many G alleles would be associated with the Non-taster phenotype if:

- the frequency of G alleles associated with the Non-taster phenotype was also 0.613
- the Non-taster phenotype was still associated with 30 alleles total

So, to calculate the expected count for a specific allele and phenotype, multiply:

- a) the Allele Frequency of the specific allele in the total population BY
- b) the total Allele Count for the specific phenotype.

The equation for the expected count of a specific allele associated with a specific phenotype is:

Expected Count =

Allele Frequency of specific allele in total population * total Allele Count for specific phenotype

Now, we can use this equation to calculate the expected count for G alleles associated with the Non-taster phenotype:

Expected Count =

Allele Frequency of G allele in total population * total Allele Count for Non-taster phenotype =

0.613 * 30 = 18.4

So, if this variant was not associated with bitter tasting perception, we would expect there to be 18.4 G alleles associated with the Non-taster phenotype.

It may seem strange to have an expected count of 18.4 alleles, since it is not possible to have just part of an allele in real life. However, since these are not real counts, it is okay to have decimals.

Activity

As another practice, calculate the expected count for C alleles associated with the Taster phenotype.

- 3. First, find the frequency of <u>C alleles</u> in the <u>total population</u> in **Table 3. Allele Frequency**.
 - a. What cell in **Table 3** contains the frequency of C alleles in the total population? Put an X in the correct cell in the example table below.

Table 3. Allele Frequency	Table	3. Alle	ele Fre	equency
---------------------------	-------	---------	---------	---------

	Non-taster	Super taster	Taster	Total Population
G				
С				
Total				

- b. What is the allele frequency?
- 4. Next, find the <u>total number of alleles</u> associated with the <u>Taster</u> phenotype in **Table 2. Allele Counts.**
 - a. What cell in **Table 2** contains the total number of alleles associated with the Taster phenotype? Put an X in the correct cell in the example table below.

	Non-taster	Super taster	Taster	Total
G				
с				
Total				

Table 2. Allele Count

- b. What is the total number of alleles associated with the Taster phenotype?
- 5. Now, use the numbers you just identified to calculate the expected count for C alleles associated with the Taster phenotype.

Expected Count =

Allele Frequency of C allele in total population * total Allele Count for Taster phenotype =

Now that you've had some practice calculating expected counts, you can proceed with the rest of the analysis. In this step of the analysis, you will calculate the expected counts for each combination of allele and phenotype. You will enter this information into **Table 4. Expected Counts**. Before you begin, familiarize yourself with **Table 4**.

Table 4 has 3 data rows and 4 data columns, <u>not</u> including the row and column labels.

- 6. Find **Table 4. Expected Counts** and use the table to enter the following questions.
 - a. What does column 4 represent? Predict the value of each cell in column 4. Hint: When calculating expected counts, we assume that the total number of each allele in the population remains the same as in the original dataset.

b. What does row 3 represent? Predict the value of each cell in row 3. *Hint: When calculating expected counts, we assume that the total number of individuals with each phenotype remains the same as in the original dataset.*

Following the same process as the example calculations, calculate the expected count for each allele and phenotype combination and enter the data into **Table 4. Expected Counts**.

- 7. For each phenotype, calculate the expected count for the G allele using the expected count equation and enter the data into the correct column of the **G** row of **Table 4**.
- 8. For each phenotype, calculate the expected count for the C allele using the expected count equation and enter the data into the correct column of the C row of **Table 4**.
- 9. Add the expected counts across each row and enter those values into the **Total** column.
- 10. Add the expected counts down each column and enter those values into the **Total** row.

After completing all the calculations, use your completed **Table 4** to answer the following questions.

11. How do the values in the **Total** row and **Total** column compare to the values in the same row and column in **Table 2**? Why?

12. What patterns do you notice in the data?

13. Are the actual counts similar to or different from the expected counts? What conclusions could you draw from this comparison, if any?

Part 4: Chi-Square Test

Note: Part 3 and Part 4 introduce some basic statistics and are intended for AP students or for students with a statistics background. With some additional background information and support from the teacher, this portion of the activity could also be used to introduce statistical concepts to students who are not as familiar.

Background

There is one final step to answering our question: is there an association between this *TAS2R38* variant and bitter taste perception phenotype?

Specifically, we will perform a Pearson's chi-square test of independence using our actual and expected allele counts. The Pearson's chi-square of independence evaluates whether observations of measures on two variables (in our case, allele and phenotype) are independent of each other.

Remember that our expected counts represent what the allele counts would be if the counts were randomly distributed and independent of phenotype. So, by using the chi-square test to compare our expected and actual allele counts, we are testing how likely it is that the difference between our actual counts and the expected counts is due to chance.

Our null and alternative hypotheses for this test are:

- Null hypothesis (H_o): the count of each allele is independent of phenotype (the count of each allele is the same or very similar for each phenotype)
- Alternative hypothesis (H_a): the count of each allele is dependent on phenotype (the count of each allele is different for each phenotype)

For statistical tests, it is important that we establish a level of significance before we perform the testing. The level of significance, also called the alpha value, represents the probability of obtaining your results due to chance. Your chosen alpha value can change based on your research question and how much potential error you are willing to tolerate in your results.

In this case, we will choose an alpha value of .05, representing a 5% probability that our results are due to chance. So, if the outcome of our test suggests that the probability that our results are due to chance is <u>less than 5%</u>, we can reject our null hypothesis and conclude that there is an association between this *TAS2R38* variant and bitter taste perception phenotype.

Activity

The formula for the Pearson's chi-square test is complex. Luckily there is a formula built into Excel and Google Sheets that can do this calculation for us!

The formula is called CHISQ.TEST(). It takes the actual and expected counts as inputs and returns a probability value, or **p value**. The **p value** represents the probability that the distribution of alleles in our actual counts table is due to chance. If you were to use this formula to perform the test on your actual and expected counts, you would get a **p value** of 2.16×10^{-10} , or <u>0.00000000216</u>.

- 1. Use the result from the Pearson's chi-square test of independence to answer the following questions:
 - a. Does the **p value** meet our established criteria for statistical significance (alpha = 0.05)? Explain.

b. What does this mean about association between this *TAS2R38* variant and bitter taste perception phenotype?

c. Does this outcome reflect your earlier conclusions from Parts 1-3?

d. How broadly can we generalize these results, given what we know (and don't know) about our sample population?

Part 5: Reflection

Note: If you perform this activity as a jigsaw with the three different SNPs, you can have students who worked on each different SNP share their spreadsheet results and present their conclusions. After each group member has had a chance to share, have students discuss the following questions.

Use the following questions to draw conclusions and reflect on the dataset and calculations.

1. What can we conclude about allele/phenotype association from this dataset?

In this study, **Table 1** represents genotype and phenotype data from 60 individuals. The genotype data was determined from DNA sequencing. It is important to note that, while **Table 1** separates the genotype into two **allele** columns, we only know the genotype (i.e., what combination of alleles is present) for each SNP location. We do not know which allele is on which chromosome.

2. Can we draw any conclusions about genotype/phenotype association based on these calculations? Why or why not?

- 3. Each individual has two total *TAS2R38* alleles. Each of those two alleles has one specific nucleotide at each variant location. This combination of the three variants on a specific allele is called a haplotype. For example, one possible haplotype is GTA, representing a G at the location of SNP 1, a T at the location of SNP 2, and an A at the location of SNP 3.
 - a. From this dataset, do we know which combination of variants were inherited together on a single chromosome in a single individual? Why or why not?

b. Can we draw any conclusions about haplotype/phenotype association from this dataset? Why or why not?

4. How broadly can we generalize our results, given what we know (and don't know) about our sample population?

5. Based <u>only on your results</u>, can you conclude that these variants in *TAS2R38* cause an individual's bitter taste perception phenotype? Explain.

Remember that in a GWAS, scientists look at millions of variants across a large population with varied phenotypes for a trait, or disease, of interest. Typically, GWAS are used to study complex diseases or traits that are impacted by multiple genetic variants and environmental factors.

6. Generally, a GWAS can establish correlations between genetic variants and a disease or phenotype. However, a GWAS cannot be used to prove that a certain variant *causes* a phenotype. Why might it be important to establish a correlation between a small number of variants and a phenotype even if we cannot yet prove that those variants cause the phenotype?

7. Based on your experience completing calculations for one variant, why might computational tools and automation be important for GWAS?

VARIATION IN THE GENOME TEACHER GUIDE

Appendix: Data and Analysis Tables

Table 1. Allele Data

Sample Name	SNP 1 (G	C) Alleles	Phenotype
AG07	G	G	Non-taster
AG12	G	G	Non-taster
AG24	G	G	Non-taster
AG36	G	G	Non-taster
KL10	G	G	Non-taster
KL17	G	G	Non-taster
KI 10	G	G	Non-taster
KL 20	C C	C	Non-taster
KL20	0	0	Non-laster
KLZ1	G	G	Non-taster
KL28	G	G	Non-taster
KL36	G	G	Non-taster
KL52	G	G	Non-taster
KL54	G	G	Non-taster
KL59	G	G	Non-taster
KL60	G	G	Non-taster
AG04	С	С	Super taster
AG10	С	С	Super taster
AG11	C	C	Super taster
AG19	C	C	Super taster
AG25	C	C	Super tester
KI 40	0	0	Super taster
NL18	0		Super taster
KL34	C	C	Super taster
KL38	С	С	Super taster
AG01	С	G	Taster
AG03	G	С	Taster
AG05	G	G	Taster
AG13	С	G	Taster
AG14			Taster
AG15	G	c	Taster
AG18	0		Taster
AG10	C	<u> </u>	Taster
AGZI	G	0	Taster
AGZZ	G	G	Taster
AG23	С	G	Taster
AG26	С	G	Taster
AG30	С	G	Taster
AG31	С	С	Taster
KL03	С	G	Taster
KL04	G	С	Taster
KL06	С	G	Taster
KL08	G	С	Taster
KI 11	C	G	Taster
KI 12	G	C	Taster
	6	G	Taster
NL13	0	0	Taster
NL14	G	C	Taster
KL15	G	G	Taster
KL16	G	G	Taster
KL26	G	C	Taster
KL29			Taster
KL30			Taster
KL32	С	G	Taster
KL33	С	G	Taster
KI 35	G	C	Taster
KL 27	G	C	Taster
KL 20	0	0	Taster
NL39	C	9	Taster
KL53	G	G	laster
KL55	С	G	Taster
KL56	G	С	Taster
KL57			Taster
KL61			Taster
AG02			

Table 2. Allele Counts

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

Table 3. Allele Frequency

	Non-taster	Super taster	Taster	Total Population
G				
С				
Total				

Table 4. Expected Counts

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

Establishing Genotype-Phenotype Associations: Spreadsheets

Introduction

What is a GWAS?

Did you know that most of our traits and many human diseases are not caused by a single genetic variant, but still are influenced by genetics?

For example, our heart health is influenced by both genetics and our environment. An individual with a variant in the *CDKN2A* gene might have an increased risk of developing heart disease, but whether that individual develops disease is also dependent on other genetic variants and on their environment. Even someone who does not have that variant in *CDKN2A* can develop heart disease, especially if factors in their behavior, such as smoking, or environment, such as exposure to air pollution, lead to increased risk.

For complex traits, like our heart health, how do scientists identify associations between genetic variants and a trait of interest when there are so many other variables? One tool that scientists use is called a Genome Wide Association study, or GWAS (pronounced *GEE-wahs*).

For a GWAS, scientists start by collecting genetic data from a large population with varied phenotypes for a particular trait or disease. Then the scientists look at millions of variants and determine if any of those variants are associated with a particular phenotype of that trait. Typically, scientists will look at single nucleotide polymorphisms, or SNPs (pronounced *snips*). SNPs are a type of variant where just one nucleotide varies from person to person.

For example, to try to find associations between genetic variants and heart disease, scientists would recruit a large group of people to participate in their study, some of whom have heart disease and others who do not. The scientists look at all of the participants' genetic variants and see if there are any variants that are more common in the group with heart disease than they are in the group without heart disease. Then, they perform statistical tests to determine if those variants are more common in the heart disease group than would be expected just by chance.



Figure 1. Representation of potential outcomes of a GWAS study. In this example, the variants SNP 1 and SNP 2 are found in approximately equal percentages of the disease and non-disease populations. SNP 3, however, is more common in individuals with disease. (From <u>"Genome-Wide Association Studies Fact Sheet</u>" by the <u>National Human Genome Research Institute</u>).

The TAS2R38 Gene

In this activity, you will look at an example of the types of calculations involved in a GWAS to demonstrate an association between bitter taste perception and variants in the *TAS2R38* gene.

The Taste 2 Receptor Member 38 (*TAS2R38*) gene produces the TAS2R38 protein, which functions as a receptor to perceive a wide range of bitter compounds. Bitter taste receptors, including TAS2R38, are proteins found on the taste cells of the tongue. Bitter taste perception is affected by multiple genes and by environmental factors, such as the foods we are exposed to and have regular access to.

There are three bitter taste perception phenotypes which you will investigate today: non-taster, taster, and super taster.

- Non-tasters do not perceive bitter taste.
- Tasters perceive bitter taste, although the taste may be mild.
- Super tasters perceive bitter taste, and the taste is typically strong.

While we group people into these three phenotypes, these phenotypes are subjective and correspond to a range of actual bitter taste sensing levels.

Part 1: Allele Counts

Background

For our investigation, you will start with one variant in *TAS2R38*. The *TAS2R38* variants are single nucleotide polymorphisms, or SNPs (pronounced *snips*). This means that just one nucleotide varies from person to person. There are three common SNPs in *TAS2R38*, which each occur at different locations in the gene.

- SNP 1: At this variant location, the nucleotide is usually either a guanine (G) or a cytosine (C).
- SNP 2: At this variant location, the nucleotide is usually either a thymine (T) or a cytosine (C).
- SNP 3: At this variant location, the nucleotide is usually either an adenine (A) or a guanine (G).

We have collected genetic data from a group of individuals, recording both their *TAS2R38* genotype and bitter taste perception phenotype. The samples were provided voluntarily, meaning that each person agreed to provide a sample. The samples were also anonymized, meaning they were labeled with random sample names so that they cannot be traced back to the person who provided the sample.

Recall that everyone has two copies, or alleles, of each gene. For example, for the first variant (SNP 1), individuals can have a C on both copies, a G on both copies, or one C and one G.

Activity

To start your analysis, you will count how many of each allele of your assigned SNP are associated with each tasting phenotype.

- 1. Create your own copy of the <u>TAS2R38 Google Sheet</u> or <u>TAS2R38 Excel file</u>.
 - a. Note: It is good practice in data analysis to keep a copy of your raw data separate from the data that you analyze, so that you can restart from the raw data if you make a mistake in your analysis. If at any point you feel like you need to start over, you can always make or download a new copy of the *TAS2R38* spreadsheet.
- 2. Open your *TAS2R38* spreadsheet and navigate to the **SNP 1** tab.
- 3. Find **Table 1. Allele Data** on the top left of the spreadsheet. **Table 1** has 4 data columns and 60 data rows.

	Column 1	Column 2	Column 3	Column 4
	Sample Name	SNP 1 (G/	C) Alleles	Phenotype
Row 1				
Row 2				
Row 3 —				
Row 4 —				
Row 5 —				
	•	•		·



a. What type of data is in columns 2 and 3? What does that data represent?

b. What type of data is in column 4? What does that data represent?

c. What does each row represent?

d. Some of the table entries are blank. What do you think this means about that data?

e. How many individuals are included in the dataset? How many alleles are included in the dataset?

As is, the data is organized alphabetically by sample name. However, to establish correlation between genotype and phenotype, you need to organize the data by phenotype. To do this, you will sort the data in the table.

4. Sort the data table by the Phenotype column using Sort range function in Google Sheets or the Sort & Filter function in Excel. Make sure that the range you select to sort contains only the data in Table 1. If you include the column names in your selected range, make sure that you indicate that your data has a header row.

Since the data is now sorted by phenotype, you can more easily count the number of G and C alleles that are associated with each of the bitter taste perception phenotypes. You will enter this information into one of the analysis tables, which is labeled **Table 2. Allele Counts**.

Table 2 has 3 data rows and 4 data columns, <u>not</u> including the row and column labels.

- Find Table 2. Allele Counts and use the table to enter the following questions.
 a. What do columns 1, 2, and 3 represent?
 - b. What does column 4 represent?
 - c. What do rows 1 and 2 represent?
 - d. What does row 3 represent?

Figure 3. Image of Table 2 with rows and columns numbered for reference.

e. Into which cell would you enter the data for the number of C alleles associated with the Taster phenotype? Put an X in the correct cell in the example table below.

Table	2. Al	lele	Counts
-------	-------	------	--------

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

f. Into which cell would you enter the data for the number of G alleles associated with the Non-taster phenotype? Put an X in the correct cell in the example table below.

Table 2. Allele Counts

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

Now that you are familiar with **Table 2**, you can count the alleles in each phenotype group and enter the counts data into **Table 2**. You can either count manually or count using a spreadsheet formula. As you enter the data, the **Total** column and **Total** row should update automatically.

Note: You can instruct students to count manually, use the formula, or complete both steps.

Manual counting:

- 6. For each phenotype, count the number of G alleles and enter the data into the correct column of the **G** row of **Table 2**. Be sure to include data from both alleles for each individual!
- 7. For each phenotype, count the number of C alleles and enter the data into the correct column of the **C** row of **Table 2**. Be sure to include data from both alleles for each individual!

Using a formula:

8. Use the countif() formula in Excel or Google Sheets to calculate count data in **Table 2**. For each cell in **Table 2**, type:

=countif(range, condition)

where range is a subset of columns 2 and 3 from **Table 1** representing the allele data for the phenotype corresponding to your current column in **Table 2**, and condition represents the allele in the corresponding row of **Table 2** (either "G" or "C").

After completing all the calculations, use your completed **Table 2** to answer the following questions.

- 9. How many G alleles are associated with each phenotype?
 - d. Non-taster:
 - e. Super taster:
 - f. Taster:
- 10. How many C alleles are associated with each phenotype?
 - d. Non-taster:
 - e. Super taster:
 - f. Taster:
- 11. What patterns do you notice in the data?

12. Based on this data alone, without performing additional calculations, do you think this variant in *TAS2R38* is associated with bitter tasting phenotype? Why or why not?

13. How could you use additional data collection, analysis, or representation to better support your hypothesis?

Part 2: Allele Frequency

Note: If your students are not yet familiar with statistics, this section provides an opportunity to do additional calculations and data analysis.

Background

Next, you can use the allele count data in **Table 2** to calculate allele frequencies. Allele frequency refers to the distribution of alleles across a population. For example, the allele frequency of the G allele in the total population is the fraction, or percentage, of the total alleles in the population that are G alleles. The equation we will use for allele frequency is:

Allele Frequency =
$$\frac{v}{2N}$$

Where the variable v is the number of each allele in the dataset, and the variable N is the total number of diploid individuals in the dataset.

In Part 1, you counted the number of each allele in the dataset, so **Table 2** contains the values for v. Additionally, since N represents the number of diploid individuals and each individual has two alleles, 2N is equal to the total allele count. So, we can also write the allele frequency equation as:

$$Allele Frequency = \frac{Allele Count of a specific allele in the population}{Total Allele Count in the population}$$

For example, in our population of 60 individuals, we have 106 known alleles in total (**Table 2**, **Total** row, **Total** column). Of those 106 alleles, 65 are G alleles (**Table 2**, **G** row, **Total** column). We can calculate the allele frequency of the G allele in our total population as follows:

Total G Allele Frequency =
$$\frac{Allele Count of G allele in the population}{Total Allele Count in the population} = \frac{65}{106} = 0.613$$

If we look at just the Taster population, there are 35 G alleles (**Table 2**, **G** row, **Taster** column). However, because we are only looking at the Taster population, our total number of alleles also changes. There are 30 Tasters, corresponding to 60 alleles (**Table 2**, **Total** row, **Taster** column). We can calculate the allele frequency of the G allele in the Taster population as follows:

$$Taster \ G \ Allele \ Frequency = \frac{Allele \ Count \ of \ G \ allele \ in \ Taster \ population}{Total \ Allele \ Count \ in \ the \ Taster \ population} = \frac{35}{60} = 0.583$$

Activity

First, practice calculating allele frequency using the data for the number of C alleles in the Taster population.

 Before you begin, find the number of C alleles associated with the Taster phenotype in Table 2. Allele Counts. What cell in Table 2 contains the number of C alleles associated with the Taster phenotype? Put an X in the correct cell in the example table below.

Table 2. Allele Counts

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

- 2. What is the allele count of C alleles associated with the Taster phenotype?
- 3. Now, use the allele count to complete the calculation below. Round your answer to three decimal places.

You will use **Table 3. Allele Frequencies** to record the results from your allele frequency calculations. Before completing the remaining calculations, get familiar with the format of **Table 3**.

		Colu	mn 1	Column 2		Column 3		Column 4	
		Non-tast	er	Super ta	ster	Taster		Total Po	pulation
ow 1 —	G								
ow 2 —	с								
ow 3 —	Total						7		

Table 3 has 3 data rows and 4 data columns, <u>not</u> including the row and column labels.

Figure 4. Image of Table 3 with rows and columns numbered for reference.

4. Find Table 3. Allele Frequencies and use the table to enter the following questions.e. What does column 4 represent?

f. What does row 3 represent? Predict the value of each cell in row 3. Hint: In allele frequency, we are calculating the frequency of each allele within the population. What should be the frequency of both alleles combined in the population?

g. Into which cell would you enter the data for the frequency of G alleles in the total population? Put an X in the correct cell in the example table below.

Table 3. Allele Frequency

	Non-taster	Super taster	Taster	Total Population
G				
С				
Total				

h. Into which cell would you enter the data for the frequency of C alleles associated with the Super taster phenotype? Put an X in the correct cell in the example table below.

Table 3. Allele	Frequency
-----------------	-----------

	Non-taster	Super taster	Taster	Total Population
G				
с				
Total				

Now you can calculate the allele frequencies for each phenotype and enter the data into **Table 3**. You can either calculate manually using the equation, or you can use a spreadsheet formula to complete the calculations. As you enter the data, the numbers in the **Total** row should update automatically.

Note: You can instruct students to count manually, use a formula, or complete both steps.

Manual calculation:

- 5. For each phenotype and for the total population, calculate the allele frequency of the G allele and enter the data into the correct column of the **G** row of **Table 3**.
- 6. For each phenotype and for the total population, calculate the allele frequency of the C allele and enter the data into the correct column of the **C** row of **Table 3**.

Using a formula:

7. There is no pre-existing formula for allele frequency, but you can set up your own. Create your own function in Excel or Google Sheets to calculate the allele frequencies for each phenotype and for the whole population.

Hint: For each cell in Table 3, divide the corresponding cell in Table 2 by the cell in the Total row of the same column of Table 2.

After completing all the calculations, use your completed **Table 3** to answer the following questions.

- 8. What is the G allele frequency for each of the following phenotype groups:
 - e. Non-taster:
 - f. Super taster:
 - g. Taster:
 - h. Total Population:
- 9. How is the C allele frequency for each of the following phenotype groups:
 - e. Non-taster:
 - f. Super taster:
 - g. Taster:
 - h. Total Population:
- 10. What patterns do you notice in the data?

11. The data in Table 2 and Table 3 come from the same raw data but show different ways of analyzing and representing that data. Which set of analyzed data, allele counts (**Table 2**) or allele frequency (**Table 3**), is easier for you to understand? Why?

Note: This question could be a small group or whole class discussion. Students can discuss why they selected a particular analysis method. With teacher scaffolding, students may also discuss how this relates to why data in science is often analyzed, represented, and displayed in different ways.

Part 3: Expected Counts

Note: Part 3 and Part 4 introduce some basic statistics and are intended for AP students or for students with a statistics background. With some additional background information and support from the teacher, this portion of the activity could also be used to introduce statistical concepts to students who are not as familiar.

Students can still complete Part 3 even if you do not intend for them to complete Part 4.

Background

In **Table 2. Allele Counts** and **Table 3. Allele Frequency**, we have data about the distribution of the different *TAS2R38* SNP 1 alleles across the different bitter tasting phenotypes. Now we can ask the question: is there an association between variant and phenotype?

To answer this question, we can use a statistical test called a Pearson's chi-square test of independence. The Pearson's chi-square test of independence evaluates whether observations of measures on two variables (in our case, allele and phenotype) are independent of each other. The chi-square test requires us to compare our actual allele counts to **expected counts**.

Expected counts represent what the allele counts would be if there was no association between the variant and phenotype. In other words, **expected counts** represent a random distribution of alleles across the phenotypes, such that <u>each phenotype has the same allele frequencies as the total population</u>.

Example Calculation

For example, let's look at the expected count of G alleles associated with the Non-taster phenotype.

The frequency of G alleles in the total population is 0.613. Importantly, expected count calculations assume that the total number of alleles associated with each phenotype remains the same. In our population, there are 30 total alleles associated with the Non-taster phenotype.

1. The frequency of <u>G alleles</u> in the <u>total population</u> (0.613) is in **Table 3. Allele Frequency**. What cell in **Table 3** contains the frequency of G alleles in the total population? Put an X in the correct cell in the example table below.

	Non-taster	Super taster	Taster	Total Population
G				
С				
Total				

Table 3. Allele Frequency

 The total number of alleles associated with the <u>Non-taster</u> phenotype (30) is in **Table 2. Allele** Counts. What cell in **Table 2** contains the total number of alleles associated with the Non-taster phenotype? Put an X in the correct cell in the example table below.

_	Non-taster	Super taster	Taster	Total
G				
С				
Total				

The expected count of G alleles for the Non-taster population represents how many G alleles would be associated with the Non-taster phenotype if:

- the frequency of G alleles associated with the Non-taster phenotype was also 0.613
- the Non-taster phenotype was still associated with 30 alleles total

So, to calculate the expected count for a specific allele and phenotype, multiply:

- c) the Allele Frequency of the specific allele in the total population BY
- d) the total Allele Count for the specific phenotype.

The equation for the expected count of a specific allele associated with a specific phenotype is:

Expected Count =

Allele Frequency of specific allele in total population * total Allele Count for specific phenotype

Now, we can use this equation to calculate the expected count for G alleles associated with the Non-taster phenotype:

Expected Count =

Allele Frequency of G allele in total population * total Allele Count for Non-taster phenotype =

0.613 * 30 = 18.4

So, if this variant was not associated with bitter tasting perception, we would expect there to be 18.4 G alleles associated with the Non-taster phenotype.

It may seem strange to have an expected count of 18.4 alleles, since it is not possible to have just part of an allele in real life. However, since these are not real counts, it is okay to have decimals.

Activity

As another practice, calculate the expected count for C alleles associated with the Taster phenotype.

- 3. First, find the frequency of <u>C alleles</u> in the <u>total population</u> in **Table 3. Allele Frequency**.
 - a. What cell in **Table 3** contains the frequency of C alleles in the total population? Put an X in the correct cell in the example table below.

Table	3.	Allele	Frequency	
10010		/ mere	i i equeiley	

	Non-taster	Super taster	Taster	Total Population
G				
С				
Total				

- b. What is the allele frequency?
- 4. Next, find the <u>total number of alleles</u> associated with the <u>Taster</u> phenotype in **Table 2. Allele Counts.**
 - a. What cell in **Table 2** contains the total number of alleles associated with the Taster phenotype? Put an X in the correct cell in the example table below.

	Non-taster	Super taster	Taster	Total
G				
С				
Total				

Table 2. Allele Count

- b. What is the total number of alleles associated with the Taster phenotype?
- 5. Now, use the numbers you just identified to calculate the expected count for C alleles associated with the Taster phenotype.

Expected Count =

Allele Frequency of C allele in total population * total Allele Count for Taster phenotype =

_____* _____ = _____

Now that you've had some practice calculating expected counts, you can proceed with the rest of the analysis. In this step of the analysis, you will calculate the expected counts for each combination of allele and phenotype. You will enter this information into **Table 4. Expected Counts**. Before you begin, familiarize yourself with **Table 4**.

Table 4 has 3 data rows and 4 data columns, <u>not</u> including the row and column labels.

- 6. Find **Table 4. Expected Counts** and use the table to enter the following questions.
 - c. What does column 4 represent? Predict the value of each cell in column 4. Hint: When calculating expected counts, we assume that the total number of each allele in the population remains the same as in the original dataset.

d. What does row 3 represent? Predict the value of each cell in row 3. Hint: When calculating expected counts, we assume that the total number of individuals with each phenotype remains the same as in the original dataset.

Following the same process as the example calculations, calculate the expected count for each allele and phenotype combination and enter the data into **Table 4. Expected Counts**. You can either calculate manually using the equation, or you can use a spreadsheet formula to complete the calculations.

Note: You can instruct students to count manually, use the formula, or complete both steps

Manual calculation:

- 7. For each phenotype, calculate the expected count for the G allele using the expected count equation and enter the data into the correct column of the **G** row of **Table 4**.
- 8. For each phenotype, calculate the expected count for the C allele using the expected count equation and enter the data into the correct column of the C row of **Table 4**.

Using a formula:

9. There is no pre-existing formula for expected counts, but you can still set up your own. Create your own function in Excel or Google Sheets to calculate the expected count for each allele and phenotype combination.

Hint: For each cell in Table 4, multiply the cell that corresponds to the same <u>phenotype</u> in the Table 2 Total row by the cell that corresponds to the same <u>allele</u> in the Table 3 Total Population column.

After completing all the calculations, use your completed **Table 4** to answer the following questions.

10. How do the values in the **Total** row and **Total** column compare to the values in the same row and column in **Table 2**? Why?

11. What patterns do you notice in the data?

12. Are the actual counts similar to or different from the expected counts? What conclusions could you draw from this comparison, if any?

Part 4: Chi-Square Test

Note: Part 3 and Part 4 introduce some basic statistics and are intended for AP students or for students with a statistics background. With some additional background information and support from the teacher, this portion of the activity could also be used to introduce statistical concepts to students who are not as familiar.

Background

There is one final step to answering our question: is there an association between this *TAS2R38* variant and bitter taste perception phenotype?

Specifically, we will perform a Pearson's chi-square test of independence using our actual and expected allele counts. The Pearson's chi-square test of independence evaluates whether observations of measures on two variables (in our case, allele and phenotype) are independent of each other.

Remember that our expected counts represent what the allele counts would be if the counts were randomly distributed and independent of phenotype. So, by using the chi-square test to compare our expected and actual allele counts, we are testing how likely it is that the difference between our actual counts and the expected counts is due to chance.

Our null and alternative hypotheses for this test are:

- Null hypothesis (H_o): the count of each allele is independent of phenotype (the count of each allele is the same or very similar for each phenotype)
- Alternative hypothesis (H_a): the count of each allele is dependent on phenotype (the count of each allele is different for each phenotype)

For statistical tests, it is important that we establish a level of significance before we perform the testing. The level of significance, also called the alpha value, represents the probability of obtaining your results due to chance. Your chosen alpha value can change based on your research question and how much potential error you are willing to tolerate in your results.

In this case, we will choose an alpha value of .05, representing a 5% probability that our results are due to chance. So, if the outcome of our test suggests that the probability that our results are due to chance is <u>less than 5%</u>, we can reject our null hypothesis and conclude that there is an association between this *TAS2R38* variant and bitter taste perception phenotype.

Activity

The formula for the Pearson's chi-square test is complex. Luckily there is a formula built into Excel and Google Sheets that can do this calculation for us!

The formula is called CHISQ.TEST(). It takes the actual and expected counts as inputs and returns a probability value, or **p value**. The **p value** represents the probability that the distribution of alleles in our actual counts table is due to chance.

1. Calculate the Pearson's chi-square test of independence **p value** using the CHISQ.TEST() formula and your actual and expected counts tables.

To use the CHISQ.TEST() function, choose a cell to store your results and type: =CHISQ.TEST(actual_range, expected_range)

where actual_range represents the cells containing your actual count data from **Table 2** and expected_range represents the cells containing your expected count data from **Table 4**. Be sure to only include the count data and <u>not</u> the **Total** rows/columns or the row/column headers.

Hint: If you have kept everything in your spreadsheet in its original location, your formula should read: =CHISQ.TEST(112:K13, 128:K29)

- 2. Use the result from the Pearson's chi-square test of independence to answer the following questions:
 - a. Does the **p value** meet our established criteria for statistical significance (alpha = 0.05)? Explain.

b. What does this mean about association between this *TAS2R38* variant and bitter taste perception phenotype?

c. Does this outcome reflect your earlier conclusions from Parts 1-3?

d. How broadly can we generalize these results, given what we know (and don't know) about our sample population?

Part 5: Reflection

Note: If you perform this activity as a jigsaw with the three different SNPs, you can have students who worked on each different SNP share their spreadsheet results and present their conclusions. After each group member has had a chance to share, have students discuss the following questions.

Use the following questions to draw conclusions and reflect on the dataset and calculations.

1. What can we conclude about allele/phenotype association from this dataset?

In this study, **Table 1** represents genotype and phenotype data from 60 individuals. The genotype data was determined from DNA sequencing. It is important to note that, while **Table 1** separates the genotype into two **allele** columns, we only know the genotype (i.e., what combination of alleles is present) for each SNP location. We do not know which allele is on which chromosome.

2. Can we draw any conclusions about genotype/phenotype association based on these calculations? Why or why not?

- 3. Each individual has two total *TAS2R38* alleles. Each of those two alleles has one specific nucleotide at each variant location. This combination of the three variants on a specific allele is called a haplotype. For example, one possible haplotype is GTA, representing a G at the location of SNP 1, a T at the location of SNP 2, and an A at the location of SNP 3.
 - a. From this dataset, do we know which combination of variants were inherited together on a single chromosome in a single individual? Why or why not?

b. Can we draw any conclusions about haplotype/phenotype association from this dataset? Why or why not?

4. How broadly can we generalize our results, given what we know (and don't know) about our sample population?

5. Based <u>only on your results</u>, can you conclude that these variants in *TAS2R38* cause an individual's bitter taste perception phenotype? Explain.

Remember that in a GWAS, scientists look at millions of variants across a large population with varied phenotypes for a trait, or disease, of interest. Typically, GWAS are used to study complex diseases or traits that are impacted by multiple genetic variants and environmental factors.

6. Generally, a GWAS can establish correlations between genetic variants and a disease or phenotype. However, a GWAS cannot be used to prove that a certain variant *causes* a phenotype. Why might it be important to establish a correlation between a small number of variants and a phenotype even if we cannot yet prove that those variants cause the phenotype?

7. Based on your experience completing calculations for one variant, why might computational tools and automation be important for GWAS?

Implementation Strategies

This module was built to be flexible and adaptable to different classroom environments. These strategies are provided as examples for how you might implement these activities in the classroom. If you have any questions about implementing this module, reach out to the TtGG team at <u>ttgg@jax.org</u>.

Strategy 1: Full Activity

Day 1

Pre-work:

- Complete the <u>Introduction</u> reading on <u>What is a GWAS?</u> and <u>The TAS2R38 Gene</u>
- Review page 1 of the <u>TAS2R38 Gene Sheet</u>
 - Note: The second page describes the association between TAS2R38 genotype and bitter taste perception phenotype, so you may want to hold off on sharing page 2 with students until after they complete the Establishing Genotype-Phenotype Associations activity.
- If using the <u>Spreadsheet</u> version, have students each make or download a copy of the TAS2R38 Student Data <u>Google Sheet</u> or <u>Excel file</u>.

In class:

- Address any student questions about the pre-work
- Complete Part 1: Allele Counts (<u>Paper-Based</u> version or <u>Spreadsheets</u> version)
 - o Individually
 - In pairs/groups, each group working on SNP 1
 - In pairs/groups, each group working on a different SNP (1, 2, or 3)
 - See <u>Supporting Materials</u> for links to student worksheets for each SNP
- Complete <u>Background</u> reading for Part 2: Allele Frequency and/or teacher presents on how to perform Allele Frequency calculations
- Complete Part 2: Allele Frequency (Paper-Based version or Spreadsheets version)
 - $\circ \quad \text{Individually} \\$
 - In pairs/groups, each group working on SNP 1
 - In pairs/groups, each group working on a different SNP (1, 2, or 3)
 - See <u>Supporting Materials</u> for links to student worksheets for each SNP

Day 2

Pre-work:

• Complete <u>Background</u> reading and <u>Example Calculation</u> for Part 3: Expected Counts In class:

- Address any student questions about the pre-work
- Complete Part 3: Expected Counts Activity (<u>Paper-Based</u> version or <u>Spreadsheets</u> version)
 - o Individually
 - In pairs/groups, each group working on SNP 1
 - In pairs/groups, each group working on a different SNP (1, 2, or 3)
 - See <u>Supporting Materials</u> for links to student worksheets for each SNP
- Complete Part 4: Chi-Square Test (Paper-Based version or Spreadsheets version)
 - $\circ \quad \text{Individually} \\$
 - In pairs/groups, each group working on SNP 1
 - In pairs/groups, each group working on a different SNP (1, 2, or 3)
 - See <u>Supporting Materials</u> for links to student worksheets for each SNP

- Complete Part 5: Reflection (<u>Paper-Based</u> version or <u>Spreadsheets</u> version)
 - o Students complete individually
 - Students complete in same small groups as whole activity
 - Students complete in new small groups, as jigsaw with one student who worked on each SNP in each new group
 - Group discussion as a whole class

Strategy 2: Allele Frequency Only (No Statistical Analysis)

Pre-work:

- Complete the <u>Introduction</u> reading on <u>What is a GWAS?</u> and <u>The TAS2R38 Gene</u>
- Review page 1 of the <u>TAS2R38 Gene Sheet</u>
 - Note: The second page describes the association between TAS2R38 genotype and bitter taste perception phenotype, so you may want to hold off on sharing page 2 with students until after they complete this activity.
- If using the spreadsheet version, have students each make or download a copy of the TAS2R38 Student Data <u>Google Sheet</u> or <u>Excel file</u>. You may wish to remove Table 4 from the spreadsheet.

In class:

- Address any student questions about the pre-work
- Complete Part 1: Allele Counts (<u>Paper-Based</u> version or <u>Spreadsheets</u> version)
 - \circ Individually
 - In pairs/groups, each group working on SNP 1
 - In pairs/groups, each group working on a different SNP (1, 2, or 3)
 - See <u>Supporting Materials</u> for links to student worksheets for each SNP
- Complete <u>Background</u> reading for Part 2: Allele Frequency and/or teacher presents on how to perform Allele Frequency calculations
- Complete Part 2: Allele Frequency (<u>Paper-Based</u> version or <u>Spreadsheets</u> version)
 - o Individually
 - In pairs/groups, each group working on SNP 1
 - In pairs/groups, each group working on a different SNP (1, 2, or 3)
 - See <u>Supporting Materials</u> for links to student worksheets for each SNP
- Complete Part 5: Reflection (<u>Paper-Based</u> version or <u>Spreadsheets</u> version)
 - Students complete individually
 - Students complete in same small groups as whole activity
 - Students complete in new small groups, as jigsaw with one student who worked on each SNP in each new group
 - Group discussion as a whole class

Post-work (Optional):

If all students completed SNP 1 in class, ask students to repeat the process for SNP 2 and/or SNP 3 as homework. You can provide the corresponding handouts (see <u>Supporting Materials</u>) or ask them to follow the same process without providing direct instructions.

NGSS Alignments

Activity or Lesson	Disciplinary Core Ideas	Cross Cutting Concepts	Science & Engineering Practices
<u>Establishinq</u> <u>Genotype-Phenotype</u> <u>Associations: Paper-</u> <u>Based</u>	LS3.B: Variation of Traits	Patterns	Analyzing and Interpreting Data Using Mathematics and Computational Thinking Constructing Explanations and Designing Solutions
<u>Establishinq</u> <u>Genotype-Phenotype</u> <u>Associations:</u> <u>Spreadsheets</u>	LS3.B : Variation of Traits	Patterns	Analyzing and Interpreting Data Using Mathematics and Computational Thinking Constructing Explanations and Designing Solutions

VARIATION IN THE GENOME TEACHER GUIDE

Supporting Materials

Spreadsheets

- TAS2R38 Student Data: Google Sheet
- TAS2R38 Student Data: Excel
- TAS2R38 Data Teacher Answer Key: Google Sheet
- TAS2R38 Data Teacher Answer Key: Excel

Student Worksheets

- Spreadsheets:
 - Establishing Genotype-Phenotype Associations: TAS2R38 SNP 1 Spreadsheets
 - o Establishing Genotype-Phenotype Associations: TAS2R38 SNP 2 Spreadsheets
 - <u>Establishing Genotype-Phenotype Associations: TAS2R38 SNP 3 Spreadsheets</u>
- Paper-Based:
 - o Establishing Genotype-Phenotype Associations: TAS2R38 SNP 1 Paper-Based
 - o Establishing Genotype-Phenotype Associations: TAS2R38 SNP 2 Paper-Based
 - o Establishing Genotype-Phenotype Associations: TAS2R38 SNP 3 Paper-Based

Documents

• TAS2R38 Gene Sheet

Tutorials

• Spreadsheet tutorials coming soon!

Minute to Understanding Video

• What are DNA variants?

Related Ethics Lessons from the Personal Genetics Education Project (pgEd)

- Snapshots
 - o <u>The Common Thread: DNA and Human Variation</u>
- Full lessons
 - o <u>Introduction to Personal Genetics</u>
 - o <u>Scientific Themes in Personal Genetics</u>

Related Interactives from LabXchange

- How Does Taste Work?
- The TAS2R38 Taste Receptor: Does That Taste Bitter to You?

Feedback

Did you use any of the Variation in the Genome content in your class(es)? If so, we'd love to hear how it went. Use the following form to provide the TtGG team with feedback.

Feedback Form