## **ALLELIC INDEPENDENCE**

## **Testing for Allelic Independence**

What is the probability a person has a particular DNA profile? What is the probability a person has a particular profile if it has already been seen once?

The first question is a little easier to think about, but difficult to answer in practice: it is very unlikely that a profile will be seen in any sample of profiles. Even for one STR locus with 10 alleles, there are 55 different genotypes and most of those will not occur in a sample of a few hundred profiles.

For locus D3S1358 in the African American population, the FBI frequency database shows that 31 of the 55 genotype counts are zero. Estimating the population frequencies for these 31 types as zero doesn't seem sensible.

## **D3S1358** Genotype Counts

Observed	<12	12	13	14	15	16	17	18	19	>19
<12	0									
12	O	0								
13	0	0	0							
14	0	0	0	2						
15	0	0	1	19	15					
16	1	1	1	15	39	19				
17	0	0	2	10	26	24	9			
18	1	0	1	2	6	10	3	0		
19	0	0	0	1	0	0	1	0	0	
>19	0	0	0	0	1	0	0	0	0	0

The number in row i and column j is the observed count of indivuals with alleles i and j.

## Hardy-Weinberg Law

A solution to the problem is to assume that the Hardy-Weinberg Law holds. For a random mating population, expect that genotype frequencies are products of allele frequencies.

For a locus with two alleles, A, a:

$$P_{AA} = (p_A)^2$$

$$P_{Aa} = 2p_A p_a$$

$$P_{aa} = (p_a)^2$$

For a locus with several alleles  $A_i$ :

$$P_{A_i A_i} = (p_{A_i})^2$$

$$P_{A_i A_i} = 2p_{A_i} p_{A_i}$$

## D3S1358 Hardy-Weinberg Calculations

The allele counts for D3S1358 in the African-American sample are:

											Total
Allele	<12	12	13	14	15	16	17	18	19	>19	
Count	2	1	5	51	122	129	84	23	2	1	420

If the Hardy-Weinberg Law holds, then we would expect to see  $n\tilde{p}_{13}^2=210\times(5/420)^2=0.03$  individuals of type 13,13 in a sample of 210 individuals.

Also, we would expect to see  $2n\tilde{p}_{13}\tilde{p}_{14}=420\times(5/420)\times(51/420)=0.61$  individuals of type 13,14 in a sample of 210 individuals.

Other values are shown on the next slide.

# **D3S1358** Observed and Expected Counts

		<12	12	13	14	15	16	17	18	19	>19
<12	Obs.	0									
	Exp.	0.0									
12	Obs.	0	0								
	Exp.	0.0	0.0								
13	Obs.	0	0	0							
	Exp.	0.0	0.0	0.0							
14	Obs.	0	0	0	2						
	Exp.	0.2	0.1	0.6	3.1						
15	Obs.	0	0	1	19	15					
	Exp.	0.6	0.3	1.5	14.8	17.7					
16	Obs.	1	1	1	15	39	19				
	Exp.	0.6	0.3	1.5	15.7	37.5	19.8				
17	Obs.	0	0	2	10	26	24	9			
	Exp.	0.4	0.2	1.0	10.2	24.4	25.8	8.4			
18	Obs.	1	0	1	2	6	10	3	0		
	Exp.	0.1	0.1	0.3	2.8	6.7	7.1	4.6	0.6		
19	Obs.	0	0	0	1	0	0	1	0	0	
	Exp.	0.0	0.0	0.0	0.2	0.6	0.6	0.4	0.1	0.0	
>19	Obs.	0	0	0	0	1	0	0	0	0	0
	Exp.	0.0	0.0	0.0	0.1	0.3	0.3	0.2	0.1	0.0	0.0

 $\mathsf{HWE}$ 

## Testing for Hardy-Weinberg Equilibrium

A test of the Hardy-Weinberg Law will somehow decide if the observed and expected numbers are sufficiently similar that we can proceed as though the law can be used.

In one of the first applications of Hardy-Weinberg testing in a US forensic setting:

"To justify applying the classical formulas of population genetics in the Castro case the Hispanic population must be in Hardy-Weinberg equilibrium. Applying this test to the Hispanic sample, one finds spectacular deviations from Hardy-Weinberg equilibrium."

Lander ES. 1989. DNA fingerprinting on trial. Nature 339: 501-505.

#### **VNTR** "Coalescence"

Forensic DNA profiling initially used minisatellites, or VNTR loci, with large numbers of alleles. Heterozygotes would be scored as homozygotes if the two alleles were so similar in length that they coalesced into one band on an autoradiogram. Small alleles often not detected at all, and this is a likely cause of Lander's finding (Devlin et al, Science 249:1416-1420.) .

Considerable debate in early 1990s on alternative "binning" strategies for reducing the number of alleles (Science 253:1037-1041, 1991).

Typing has moved to microsatellites with fewer and more easily distinguished alleles, but testing for Hardy-Weinberg equilibrium continues. There are still reasons why the law may not hold.

# Population Structure can Cause Departure from HWE

If a population consists of a number of subpopulations, each in HWE but with different allele frequencies, there will be a departure from HWE at the population level. This is the Wahlund effect.

Suppose there are two equal-sized subpopulations, each in HWE but with different allele frequencies, then

	Subpopn 1	Subpopn 2	Total Popn
$\overline{p_A}$	0.6	0.4	0.5
$p_a$	0.4	0.6	0.5
$P_{AA}$	0.36	0.16	$0.26 > (0.5)^2$
$P_{Aa}$	0.48	0.48	0.48 < 2(0.5)(0.5)
$P_{aa}$	0.16	0.36	$0.26 > (0.5)^2$

## **Population Structure**

Effect of population structure taken into account with the "theta-correction." Matching probabilities allow for a variance in allele frequencies among subpopulations.

$$\Pr(AA|AA) = \frac{[3\theta + (1-\theta)p_A][2\theta + (1-\theta)p_A]}{(1+\theta)(1+2\theta)}$$

where  $p_A$  is the average allele frequency over all subpopulations. We will come back to this expression.

## **Population Admixture**

A population might represent the recent admixture of two parental populations. With the same two populations as before but now with 1/4 of marriages within population 1, 1/2 of marriages between populations 1 and 2, and 1/4 of marriages within population 2. If children with one or two parents in population 1 are considered as belonging to population 1, there is an excess of heterozygosity in the offspring population.

If the proportions of marriages within populations 1 and 2 are both 25% and the proportion between populations 1 and 2 is 50%, the next generation has

	Population 1	Population 2
$\overline{P_{AA}}$	0.09 + 0.12 = 0.21	0.04
$P_{Aa}$	0.12 + 0.26 = 0.38	0.12
$P_{aa}$	0.04 + 0.12 = 0.16	0.09
	0.75	0.25

#### **Exact HWE Test**

The preferred test for HWE is an "exact" one. The test uses the conditional probability of the genotypic counts  $(n_{AA}.n_{Aa}, n_{aa})$  given the allelic counts  $(n_A, n_a)$  and given HWE:

$$Pr(n_{AA}, n_{Aa}, n_{aa} | n_A, n_a, HWE) = \frac{n!}{n_{AA}! n_{Aa}! n_{aa}!} \frac{2^{n_{Aa}} n_A! n_a!}{(2n)!}$$

Reject the Hardy-Weinberg hypothesis if this probability is unusually small.

## **Exact HWE Test Example**

Reject the HWE hypothesis if the probability of the genotypic array, conditional on the allelic array, is among the smallest probabilities for all the possible sets of genotypic counts for those allele counts.

As an example, consider  $(n_{AA}=1,n_{Aa}=0,n_{aa}=49)$ . The allele counts are  $(n_A=2,n_a=98)$  and there are only two possible genotype arrays:

AA	Aa	aa	$Pr(n_{AA}, n_{Aa}, n_{aa}   n_A, n_a, HWE)$
1	0	49	$\frac{50!}{1!0!49!} \frac{2^0 2!98!}{100!} = \frac{1}{99}$
0	2	48	$\frac{50!}{0!2!48!} \frac{2^2 2!98!}{100!} = \frac{98}{99}$

#### **Exact HWE Test**

The probability of the data on the previous slide, conditional on the allele frequencies and on HWE, is 1/99 = 0.01. This is less than the conventional 5% significance level.

In general, the p-value is the (conditional) probability of the data plus the probabilities of all the less-probable datasets. The probabilities are all calculated assuming HWE is true.

HWE Slide 14

For large sample sizes and many alleles per locus, there are too many genotypic arrays for a complete enumeration and a determination of which are the least probable 5% arrays.

A large number of the possible arrays is generated by permuting the alleles among genotypes, and calculating the proportion of these permuted genotypic arrays that have a smaller conditional probability than the original data. If this proportion is small, the Hardy-Weinberg hypothesis is rejected.

HWE Slide 15

Mark a set of five index cards to represent five genotypes:

Card 1: A A

Card 2: A A

Card 3: A A

Card 4: a a

Card 5: a a

Tear the cards in half to give a deck of 10 cards, each with one allele. Shuffle the deck and deal into 5 pairs, to give five genotypes.

The permuted set of genotypes fall into one of four types:

AA	Aa	aa	Number of times
3	0	2	
2	2	1	
1	4	0	

Check the following theoretical values for the proportions of each of the three types, from the expression:

$$\frac{n!}{n_{AA}!n_{Aa}!n_{aa}!} \times \frac{2^{n_{Aa}}n_{A}!n_{a}!}{(2n)!}$$

AA	Aa	aa	Conditional Probability
3	0	2	$\frac{1}{21} = 0.048$
2	2	1	$\frac{12}{21} = 0.571$
1	4	0	$\frac{8}{21} = 0.381$

These should match the proportions found by repeating shufflings of the deck of 10 allele cards.

#### Permutation Test for D3S1358

For an STR locus, where  $\{n_g\}$  are the genotype counts and  $n = \sum_g n_g$  is the sample size, and  $\{n_a\}$  are the alleles counts with  $2n = \sum_a n_a$ , the exact test statistic is

$$\Pr(\{n_g\}|\{n_a\}, \mathsf{HWE}) = \frac{n!2^H \prod_a n_a!}{\prod_g n_g!(2n)!}$$

where H is the count of heterozygotes.

This probability for the African American genotypic counts at D3S1358 is  $0.6163 \times 10^{-13}$ , which is a very small number. But it is not unusually small if HWE holds: a proportion 0.81 of 1000 permutations have an even smaller probability. We do not reject the HWE hypothesis in this case.

## Linkage Disequilibrium

This term is generally reserved for association between pairs of alleles — one at each of two loci. In the present context, it may simply mean some lack of independence of profile or match probabilities at different loci.

Unlinked loci are expected to be almost independent.

However, if two profiles match at several loci this may be because they are from the same, or related, people and so are likely to match at additional loci.

HWE Slide 20