# Hardy-Weinberg Equilibrium

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# Hardy-Weinberg equilibrium

- A biological population of *n* individuals.
- A bi-allelic genetic marker.
- One locus with alleles A and B, frequencies p and q.

Ternary plot representation

• Three genotypes AA, AB, BB frequencies  $f_{AA}$ ,  $f_{AB}$  and  $f_{BB}$ .

$f_{AA}$	$f_{AB}$	$f_{BB}$
$p^2$	2pq	$q^2$

- Equilibrium achieved in one generation.
- Note that the allele frequency of A in the new generation is  $p' = \frac{2p^2 + 2pq}{2} = p^2 + pq = p(p+q) = p.$

# Hardy-Weinberg equilibrium: a longer derivation

- Let P, Q and R be the frequencies of genotypes AA, AB and BB, with P+Q+R=1, and p, q the A and B allele frequencies.
- Note that  $p = P + \frac{1}{2}Q$  and that  $q = R + \frac{1}{2}Q$ .

Mating	Frequency	AA	AB	BB
$AA \times AA$	$P^2$	$P^2$	0	0
$AA \times AB$	2PQ	PQ	PQ	0
$AA \times BB$	2PR	0	2PR	0
$AB \times AB$	$Q^2$	$\frac{1}{4}Q^{2}$	$\frac{1}{2}Q^{2}$	$\frac{1}{4}Q^{2}$
$AB \times BB$	2 <i>QR</i>	0	QR	QR
$BB \times BB$	$R^2$	0	0	$R^2$

In the next generation

$$P' = P^{2} + PQ + \frac{1}{4}Q^{2} = (P + \frac{1}{2}Q)^{2} = p^{2}$$

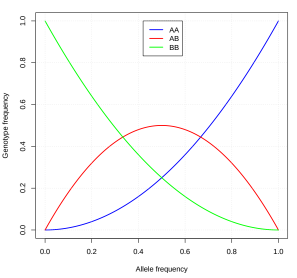
$$Q' = PQ + 2PR + \frac{1}{2}Q^{2} + QR = 2(P + \frac{1}{2}Q)(R + \frac{1}{2}Q) = 2pq$$

$$R' = \frac{1}{4}Q^{2} + QR + R^{2} = (R + \frac{1}{2}Q)^{2} = q^{2}$$

• Note we move from any arbitrary composition (P, Q, R) to  $(p^2, 2pq, q^2)$  in a single generation

# A classical genetic textbook figure

#### Genotype frequencies under HWE



# Hardy-Weinberg equilibrium

- **Equilibrium** refers to the fact that once the proportions  $p^2$ , 2pq and  $q^2$  are reached, allele frequencies and genotype frequencies will remain the same over the generations.
- Statistical tests for HWE test if the hypothesis  $f_{AA} = p^2$ ,  $f_{AB} = 2pq$ ,  $f_{BB} = q^2$  is tenable.

Ternary plot representation

 Strictly speaking, statistical tests for HWE do not assess equilibrium, but test for Hardy-Weinberg **proportions** (HWP).

# The history of Hardy-Weinberg equilibrium (1/4)



Hardy, G.H. (1908) Mendelian proportions in a mixed population. *Science* 28: 49-50.

"In a word, there is not the slightest foundation for the idea that a dominant character should show a tendency to spread over the whole population, or that a recessive should tend to die out."

JULY 10, 1908

SCIENCE

N. S. Vol. XXVIII: 49-50

DISCUSSION AND CORRESPONDENCE Mendelian Proportions in a Mixed Population

To The Billion of Science Lear relaxuate to intrude in a discussion concerning matters of which I have no expect knowledge, and I should whose these date the very simple point which I wish to make to have been familiar to biologists. However, some remarks of Mr. Udny Yale, to which Me. R. C. Pattert has called my niteration, suggest that it may still be weeth making. In the Proceedings of the Royal Society of

Medicine (Vol. L. p. 165) Mr. Yule is repersed to have suggessed, as a criticism of the Mendellan position, thus if brachydactyly is destriant. 'In the course of time one would expect, is the absence of coursewarding factors, to get these brachydacybus persons to one normal.'

It is not difficult to prove, however, thus such as

expectation would be quite groundless. Suppose that As is a pair of Mendelian characters, A being decrinant, and that in any given generation the numbers of pure dominants (Ads. heterotoguess of the control of the control of the theory of the Health, suppose that the numbers are fairly large. Health, suppose that the numbers are fairly large, which were the control of the control of the control of the secso are everly distributed among the three varieties, and that all are equally freitle. A fulle numbers will be a 'the man promotion to contribute the control of the sundipolarious table type is contributed in the transfer of the sundipolarious table to the numbers will be as 'the tree groundless' the contribute of markers will be a 'the free groundless' the contribute of the numbers will be a 'the free groundless' the production to contribute the contribute of the production of the contribute of the

 $(p+q)^2 : 2(p+q)(q+r) : (q+r)^2$ , or as  $p_1 2q_1 r_1$ , say.

The increasing question is — in what circumstances will this distribution be the same as that in the generation before? It is easy to see that the condition for this is  $q^2 = p \nu$ . And since  $q^2 = p \mu_{P_1}$ , whatever the values of p, q, and r may be the distribution will lin any case continue unchanged after this second generation.

where the second generalizes. Suppose the set of suppose the set of the set of second generalizes and second generalizes and second generalizes and second generalizes as set of the second generalizes as the second generalizes as 2000-1100,000,000, or particularly 24,0000, oncide that in the first generalizes and proportion will distribute the second generalizes proportion will distribute the second generalized second generalizes as the second generalizes are to increase if, one other bank of the second generalizes are to increase if, on the other bank cannot be second generalized as the second generalizes and better the second generalizes as well be 11,000,2001, or

practically 1:100,000,000, and this proportion would afterwards have no tendency to decrease. In a word, these is not the slightest tourshation for the idea that a dominant character should show a tendency to spend over a whole population, or that a recessive should tend to die out. I cough perhaps to add a few words on the effect

of the small deviations from the theoretical proportions which will, of course, occur in every generation. Such a distribution as p<sub>1</sub>:2q<sub>1</sub>:r<sub>1</sub>, which satisfies the condition  $q_1^2 = p_1 r_0$ , we may call a stable distribution. In actual fact we shall obtain in the second generation not  $p_1/2q_1x_1$  but a slightly different distribution p:2q:r, which is not "suble." This should, according to theory, give us in the third ecogration a "stable" distribution as 2as:rs. also differing from as 2acts; and so on. The sense in which the distribution  $p_1:2q_1:r_1$  is "stable" is this, that if we allow for the effects of casual deviations in any subsequent generation, we should, according to theory, obtain at the next generation a new "stable" distribution differing but slightly from the original distribution

Law, of ecums, consistent only the very starplent hyperhoan possible. Hyperhoans other that [ac] that of purely condens reading statistics, that of purely condens reading, will give different results, and, of course, if, as appears to be the case sometimes, the character is nor independent of that of sex, or has an inflaence on trelliny, the whole question may be grashy ascomplicated. But such complications seem to be inflaenced to the simple issue mixed by Mr. Yulkes

> Trinity College, Cambridge, April 5, 1906

P. S. I understand from Mr. Parnett that he has submitted the substance of what I have said above to Mr. Yuke, and that the lance would accept it as a satisfactory answer to the difficulty that he raised. The "stability" of the particular ratio 1:21 is recognized by Professor Karl Pearson (Phil. Trans. Roy, Sec. (A), vol. 200, p. 601.

Hardy, G. H. 1988. Mendellan proportions in a mixed

## Ternary plot representation The history of Hardy-Weinberg equilibrium (2/4)



Weinberg, W. (1908) ber den Nachweis der Vererbung beim Menschen. Jahreshefte des Vereins fr vaterIndische Naturkunde in Wrttemberg, 64:369-382.

"Thus we obtain under the influence of panmixis in each generation the same proportion of pure and hybrid types

#### Über den Nachweis der Vererbung beim Menschen\*. Von Dr. med, W. Weinberg in Stuttgart,

Unter Vererbung verstehen wir die Tatsache, daß bei der Befruchtung des Eies durch den Samen dem werdenden Individuum Art und individuelle Eigenschaften seiner Eltern erhalten werden. Dabei wird dem Kern und speziell den Chromosomen der Keimzellen der wesentlichste Anteil zugeschrieben, eine Anschauung, die allerdings gerade in der letzten Zeit wieder lebhaft bestritten ist. Die reife Geschlechtszelle macht vor der Kopulation eine doppelte Teilung durch, die sogenannte Reduktionsteilung. Durch diesen Vorgang wird nach einer verbreiteten Auffassung ein Teil der von beiden Eltern herstammenden Erbmasse ausgeschieden. Dieser Vorgang ist von der größten Wichtigkeit für die Beurteilung des Verhältnisses des Individuums zu seinen Ahnen und insbesondere für die Beurteilung der Vererbungsgesetze, die beim Menschen aufgestellt werden. Wenn wir die Ahnentafel eines Individuums konstruieren, d. h. eine schematische Übersicht seiner Vorfahren, so haben wir nur eine einflußt haben können. Aber von diesen theoretischen Möglichkeiten kommen nur wenige tatsächlich in Betracht, nicht für alle Ahnen besteht eine Kontinuität des Keimplasmas in bezug auf alle Rigenschaften, bei der Konkurrenz um die Bestimmung des Individuums in Beziehung auf iede einzelne Eigenschaft werden die meisten Ahnen ausgeschaltet. Wie viele Ahnen wirklich das Individuum in bezug auf eine bestimmte Eigenschaft determinieren, wissen wir nicht, wir können nur sagen, daß es mindestens zwei, einer väterlicher- und einer mütterlicherseits, sein müssen. Je mehr Ahnen tatsüchlich in Betracht kommen, desto größer werden wir uns die Zahl der Ah-

Vortrag am wissenschaftlichen Abend zu Stuttgart, am 13. Jan. 1908.

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# Hardy-Weinberg assumptions

- The organism under study is diploid.
- There is sexual reproduction.
- Non-overlapping generations.
- Random mating (w.r.t the trait under study).
- Population size is very large.
- Migration is negligible.
- Mutation can be ignored.
- Natural selection does not affect the trait under study.
- There is no genotyping error.

- Genetic markers are, in general, expected to follow the HW law.
- If they do not follow the law, one (or more) of the HWE assumptions is/are violated.
- The most likely cause for disequilibrium is genotyping error.
- Markers need to be checked for HWE as part of a quality control procedure.

# Hardy-Weinberg Equilibrium

$$f_{AA}$$
  $f_{AB}$   $f_{BB}$   $p^2$   $2pq$   $q^2$ 

Alternatively:

$$f_{AB}^2 = 4 f_{AA} f_{BB}$$

# Hardy-Weinberg for multiple alleles

Ternary plot representation

If a marker has three alleles (e.g. the bloodgroup system A, B and O), with frequencies  $p_1$ ,  $p_2$  and  $p_3$  with  $p_1 + p_2 + p_3 = 1$ , then under random mating we would obtain the genotype frequencies

				9	
			$p_1$	$p_2$	$p_3$
			Α	В	Ο
	$p_1$	Α	$p_1^2$	$p_1 p_2$	$p_1 p_3$
ď	$p_2$	В	$p_{2}p_{1}$	$p_{2}^{2}$	$p_2p_3$
	$p_3$	Ο	$p_{3}p_{1}$	$p_{3}p_{2}$	$p_3^2$

In general, for a k-alleles system, homozygotes  $A_iA_i$  will have frequency  $p_i^2$ , and heterozygotes  $A_iA_i$  will have frequency  $2p_ip_i$ .

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# Why is Hardy-Weinberg equilibrium important?

- It is a basic principle that, in the absence of disturbing forces, any genetic marker is expected to follow.
- Deviation from HWP is apparently most often due to genotyping error (confusion of homozygotes with heterozygotes)
- Deviation from HWP is expected (among cases) if the marker is related to disease.
- For other reasons, depending on the context of the study.

...

## Hardy-Weinberg equilibrium and disease (numerical example)

• Let A be a rare, disease-predisposing allele with  $p_A = 0.025$  (at birth, say).

	$f_{AA}$	$f_{AB}$	f <sub>BB</sub>	$p_A$
Initial	$p^2$	2pq	$q^2$	
Population	0.0006	0.0488	0.9506	0.0250

- Let P(D|AA) = 0.80, P(D|AB) = 0.40 and P(D|BB) = 0.02
- Then, potentially after many years:

	$f_{AA}$	$f_{AB}$	f <sub>BB</sub>	$p_A$
Diseased	0.0128	0.4998	0.4873	0.2627
Non-diseased	0.0001	0.0304	0.9694	0.0153

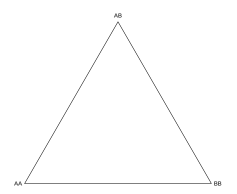
 $\bullet$  Sampling from these distributions (n = 1000), and testing for HWP with an exact test:

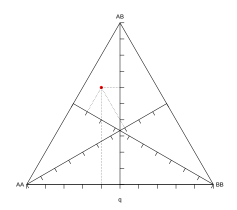
	AA	AB	BB	Exact p-value
Diseased	11	510	479	≈ 0
Non-diseased	0	19	981	$\approx 1$

Disequilibrium observed in cases, but not detected in controls.

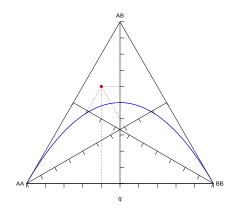
# Statistical Tests for Hardy-Weinberg Equilibrium

- Classical  $\chi^2$  test.
- Exact test (based on  $P(N_{AB} \mid N_A)$ ).
- Likelihood ratio test.
- Permutation test.
- Bayesians tests.
- ...

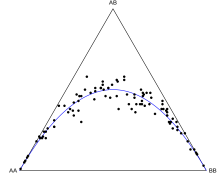




# Hardy-Weinberg Equilibrium and the Ternary Plot



# Hardy-Weinberg Equilibrium and the Ternary Plot



100 samples with n = 100,  $p \sim U(0,1)$ , simulated under HWE

# Classical $\chi^2$ test for Hardy-Weinberg equilibrium

- The counts  $n_{AA}$ ,  $n_{AB}$  and  $n_{BB}$  are regarded as a sample from a multinomial distribution.
- Expected counts under HWE are  $np^2$ , n2p(1-p) and  $n(1-p)^2$ .
- A chi-square statistic for goodness-of-fit can be used

$$X^2 = \sum_{genotypes} \frac{(observed - expected)^2}{expected}$$

- The reference distribution is a  $\chi_1^2$  distribution.
- If we define the deviation from independence  $D = \frac{1}{2}(n_{AB} e_{AB})$ , then

$$X^2 = \frac{D^2}{p^2(1-p)^2n}$$

# Example

• For an A/T polymorphism with counts AA=46, AT=39 and TT=15 we have

Ternary plot representation

$$\hat{p}_A = \frac{2 \cdot 46 + 39}{200} = 0.655$$

Expected counts under HWE

$$e_{AA} = n\hat{p}_A^2 = 100 \cdot (0.655)^2 = 42.9025$$
  
 $e_{AT} = 2n\hat{p}_A(1 - \hat{p}_A) = 2 \cdot 100 \cdot 0.655 \cdot 0.345 = 45.195$   
 $e_{TT} = n(1 - \hat{p})^2 = 100 \cdot (0.345)^2 = 11.9025$ 

•

$$X^{2} = \frac{(46 - 42.9025)^{2}}{42.9025} + \frac{(39 - 45.195)^{2}}{45.195} + \frac{(15 - 11.9025)^{2}}{11.9025} = 1.8789$$

•

$$p - \text{value} = P(\chi_1^2 > 1.8789) = 0.1704601$$

# Example in R

```
> library(HardyWeinberg)
> x <- c(46,39,15)
> names(x) <- c("AA","AT","TT")
> results <- HWChisq(x,cc=0,verbose=TRUE)
Chi-square test for Hardy-Weinberg equilibrium
Chi2 = 1.878892 p-value = 0.1704601 D = -3.0975
>
```

# Chi-square test with continuity correction

Ternary plot representation

• If the expected counts are small, a continuity correction can be applied.

$$X_c = \sum_{i=1}^{3} \frac{(|n_i - e_i| - c)^2}{e_i}$$
  $c = 0.5$ 

In R

```
> results <- HWChisq(x,verbose=TRUE)
Chi-square test with continuity correction for Hardy-Weinberg equilibrium
Chi2 = 1.441744 p-value = 0.2298573 D = -3.0975
```

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# The exact test for HWE (Levene, Haldane)

$$P(N_{AA} = n_{AA}, N_{AB} = n_{AB}, N_{BB} = n_{BB}) = \frac{n!}{n_{AA}! n_{AB}! n_{BB}!} (p_A^2)^{n_{AA}} (2p_A p_B)^{n_{AB}} (p_B^2)^{n_{BB}}$$

$$P(N_A = n_A) = \frac{2n!}{n_A! n_B!} (p_A)^{n_A} (p_B)^{n_{BB}}$$

$$P(N_{AA}, N_{AB}, N_{BB} | n_A, n_B) = \frac{n_A! n_B! n! 2^{n_{AB}}}{\frac{1}{2} (n_A - n_{AB})! n_{AB}! \frac{1}{2} (n_B - n_{AB})! (2n)!}$$

#### Notes:

- p-value: sum all probabilities of samples as extreme or more extreme as the one you observed (there are alternatives).
- It eats much more CPU than a  $\chi^2$  test (use recursion).
- It is conservative.

# Exact test computations

	Possible samples for $n = 100$ and $n_B = 14$									
	AA	AB	BB	$P(n_{AB} n_A)$	p — value	$\chi^2$	p — value	$\chi_c^2$	p — value	
1	93	0	7	0.0000	0.0000	100.00	0.0000	86.17	0.0000	
2	92	2	6	0.0000	0.0000	71.64	0.0000	60.01	0.0000	
3	91	4	5	0.0000	0.0000	47.99	0.0000	38.58	0.0000	
4	90	6	4	0.0002	0.0002	29.07	0.0000	21.86	0.0000	
5	89	8	3	0.0051	0.0053	14.87	0.0001	9.86	0.0017	
6	88	10	2	0.0602	0.0654	5.38	0.0204	2.58	0.1081	
7	87	12	1	0.3209	0.3864	0.61	0.4334	0.02	0.8849	
8	86	14	0	0.6136	1.0000	0.57	0.4516	0.02	0.8936	

# Example of the exact test

```
results <- HWExact(x,pvaluetype="selome",verbose=TRUE)
Haldane's Exact test for Hardy-Weinberg equilibrium
sample counts: nAA = 46 nAB = 39 nBB = 15
HO: HWE (D==0), H1: D <> 0
D = -3.0975 p = 0.1852682
```

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Computer exercise

The Hardy-Weinberg law essentially states that alleles combine at random into genotypes.

- Compute a test statistic (e.g.  $\chi^2$ ,  $n_{AB}$ , ...) for the observed data.
- Obtain the number of A and B alleles from the observed data.
- Permute the alleles and assemble pairs of alleles into genotypes.
- Compute the test statistic for the permuted data set (pseudo-statististic)
- Repeat this N times.
- Count the number of times the pseudo-statistic is as larger or larger than the value for the observed data (C)

• Calculate the *p*-value as C/N.

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# Example permutation test

```
> x <- c(46,39,15)
> names(x) <- c("AA","AT","TT")
> HWPerm(x)
Permutation test for Hardy-Weinberg equilibrium
Observed statistic: 1.878892 17000 permutations. p-value: 0.1864706
>
```

# Measures of (dis)equilibrium

Several statistics are being used as measures of the degree of disequilibrium:

- The  $X^2$  statistic of a test for HWE
- The p-value of an exact test for HWE
- The inbreeding coefficient  $(\hat{f})$

...

# The inbreeding coefficient (f)

$$P_{AA} = p_A^2 + p_A p_B f$$

$$P_{AB} = 2p_A p_B (1 - f)$$

$$P_{BB} = p_B^2 + p_A p_B f$$

Ternary plot representation

It can be shown that:

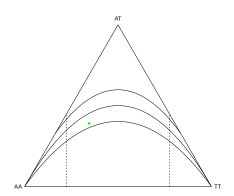
$$\frac{-p_m}{1-p_m} \le f \le 1 \text{ with } p_m = \min(p_A, p_B)$$

- f = 0: HWE
- f = 1: No heterozygotes
- f < 0: Heterozygote excess
- f > 0: Heterozygote dearth

For sample data, f is estimated by ML as:

$$\hat{f} = \frac{4n_{AA}n_{BB} - n_{AB}^2}{n_A n_B}.$$

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# Testing Equilibrium with multiple alleles

- With many alleles, some are common and many are rare.
- Asymptotic procedures do not work well with rare alleles (small counts).
- Exact procedures and permutation tests are preferable
- Computational cost increases
- Exact density for multiple alleles:

$$P(N_{ij} = n_{ij}|n_1, \dots, n_k) = \frac{n!2^h \prod_{i=1}^k n_i!}{(2n)! \prod_{i>j} n_{ij}!},$$
(1)

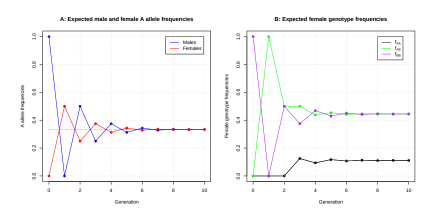
- where  $h = \sum_{i>j} n_{ij}$  is the total heterozygote frequency.
- P-value: sum of all probabilities equal or smaller than the observed sample

# Some special cases

How to test for equilibrium if...

- The variant has some recessive alleles (e.g. ABO blood groups)
- The variant is X-chromosomal
- The organism studied is tetraploid
- The variant studied has multiple copies
- ...

Ternary plot representation



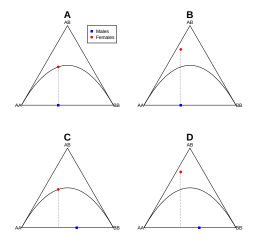
(Crow & Kimura, 1979)

# Hardy-Weinberg equilibrium and the X chromosome

Ternary plot representation

- For a marker on the X chromosome, it can take several generations before HWE is reached.
- A marker on the X chromosome is in HWE if and only if
  - Females occur in the HWP proportions (AA:  $p^2$ , AB: 2pq BB:  $q^2$ ).
  - 2 Male and female allele frequencies are equal.
- In practice, the second condition is ignored.
- If recognized, then four scenarios are possible.

# The four scenarios in ternary plots



Graffelman & Weir (Heredity, 2016)

- Let  $p_A$  be the relative frequency of the A allele.
- Let  $n_{AA}$ ,  $n_{AB}$  and  $n_{BB}$  be the numbers of the three possible genotypes, if the sexes are not distinguished.
- Let  $m_A$  and  $m_B$  be the number of males carrying the A and B allele respectively,
- Let  $f_{AA}$ ,  $f_{AB}$  and  $f_{BB}$  be the number of females of each of the three possible genotypes.
- Let  $n_m$  be the number of males, and  $n_f$  the number of females, and  $n = n_m + n_f$  the total sample size.
- The total number of alleles is given by  $n_t = 2n_f + n_m$ .

## Chi-square test for the X-chromosome

	Males Females				
Genotype	Α	В	AA	AB	BB
Probability	$\theta p_A$	$\theta(1-p_A)$	$(1-\theta)p_A^2$	$2(1-\theta)p_A(1-p_A)$	$(1-\theta)(1-p_A)^2$
Observed	$m_A$	$m_B$	$f_{AA}$	$f_{AB}$	$f_{BB}$
Expected	$n\hat{\theta}\hat{p}_A$	$n\hat{ heta}(1-\hat{p}_{A})$	$n(1-\hat{\theta})\hat{\rho}_A^2$	$2n(1-\hat{ heta})\hat{ ho}_A(1-\hat{ ho}_A)$	$n(1-\hat{ heta})\hat{ ho}_A^2$

Observed and expected genotype counts for a X-chromosomal marker under Hardy-Weinberg equilibrium.

ML estimators:

$$\hat{\theta} = \frac{n_m}{n}, \qquad \hat{p}_A = \frac{n_A}{2n_f + n_m}.$$

Chi-square statistic:

$$X^{2} = \frac{(m_{A} - e_{A})^{2}}{e_{A}} + \frac{(m_{B} - e_{B})^{2}}{e_{B}} + \frac{(f_{AA} - e_{AA})^{2}}{e_{AA}} + \frac{(f_{AB} - e_{AB})^{2}}{e_{AB}} + \frac{(f_{BB} - e_{BB})^{2}}{e_{BB}}$$

Reference distribution:

 $\chi_2^2$ 

# R Software for studying HWP

- Plink (Purcell, 2007)
- R-package HWEBayes (Wakefield, 2010)
- R-package HardyWeinberg (Graffelman, 2008)

Ternary plot representation

- R-package HWEintrinsic (Venturini, 2011)
- R-package hwde (Maindonald & Johnson, 2011)

...

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# Computer exercises

- Install the package HardyWeinberg.
- For a certain C/G polymorphism, the genotype counts n<sub>CC</sub> = 23, n<sub>CG</sub> = 48 and n<sub>GG</sub> = 29 are observed. Perform a χ<sup>2</sup> (without continuity correction) test for Hardy-Weinberg equilibrium. What is your conclusion? Repeat the test with continuity correction. Also perform the exact test for HWE. Are the results of the different tests consistent?
- For a certain C/T polymorphism, the genotype counts n<sub>CC</sub> = 0, n<sub>CT</sub> = 7 and n<sub>TT</sub> = 93 are observed. Perform a \( \chi^2 \) (without continuity correction) test for Hardy-Weinberg equilibrium. What is your conclusion? Repeat the test with continuity correction. Also perform the exact test for HWE. Are the results of the different tests consistent?
- Represent both polymorphisms in a ternary plot using the routine HWTernaryPlot.

Ternary plot representation

- Write an R function for carrying out a permutation test for HWE.
- Apply the permutation test to the two polymorphisms studied. Are the results consistent with the tests you already performed?
- Simulate 100 SNPs with a uniform allele frequency under HWE using routine HWData. Depict your results in a ternary plot. How many SNPs are out of equilibrium according to a χ<sup>2</sup> test? How many are out of equilibrium according to an exact test?
- Collect all chi-square statistics obtained in your simulation, and make a histogram. What distribution do
  they follow? Repeat your simulation with 1000 or more SNPs to get a more precise idea of the distribution.