

*The E-M Algorithm in  
Genetics*

**Biostatistics 666**

# Maximum Likelihood Estimation of Allele Frequencies

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- Find parameter estimates which make observed data most likely
- General approach, as long as tractable likelihood function exists
- Can use all available information
- Provides justification for natural estimators

# Today:

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- The Expectation–Maximization algorithm in Genetics
- Frequency estimates for...
  - Recessive alleles
  - A, B, O alleles
  - Haplotype frequencies

## Setting for the E-M Algorithm...

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- Specific type of incomplete data
  - More possible **categories** (genotypes) than can be **distinguished** (phenotypes)
- For example, consider disease locus with recessive alleles...
  - What are the possible genotypes?
  - What are the possible phenotypes?

## Setting for the E-M Algorithm...

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- Problem is simple with complete data ...
  - For example, estimating allele frequencies when all genotypes can be distinguished ...
- ... but available data can be “incomplete”.
  - For example, for recessive disease phenotypes, homozygotes and heterozygotes are hard to distinguish.

# The E-M Algorithm

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- Consider a set of starting parameters
- Use these to “estimate” the complete data
- Use estimated complete data to update parameters
- Repeat as necessary

## An Example ...

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- A random sample of 100 individuals
- 4 express a recessive phenotype
  - Assume the phenotype is controlled by a single gene
- Let's follow E-M algorithm steps ...

## Step 1:

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- Set starting values for parameters
- For allele frequency estimation...
  - Equal frequencies are a common choice
  - $p_{\text{rec}} = 0.5$
- Useful to repeat process using different starting point



## Step 2:

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- Estimate “complete data”
- Assign phenotypes to specific genotype categories
- Use Bayes’ Theorem

## Step 2 (continued):

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- Calculate probability of each genotype among the 96 “normal” individuals

$$\begin{aligned} P(+/+; Normal) &= \frac{P(+/+, Normal)}{P(Normal)} = \\ &= \frac{P(+/+, Normal)}{P(+/+, Normal) + P(+/-, Normal)} \\ &= \frac{P(+/+)}{P(+/+) + P(+/-)} \end{aligned}$$

## Step 2 (Finally!):

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- At the first iteration, the complete data would be filled in as:
  - 4 individuals with recessive genotype
  - 64 individuals with heterozygous genotype
  - 32 individuals with dominant genotype

## Step 3:

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- Estimate allele frequencies by counting...

$$P_{rec} = \frac{N_{het} + 2N_{rec/rec}}{2N}$$

- What would be the estimated allele frequencies?

# Repeat as necessary ...

Round	Estimate	E(+/+)	E(+/-)	E(-/-)	In L
1	0.50	32.00	64.00	4.00	-14.40240
2	0.36	45.18	50.82	4.00	-9.33657
3	0.29	52.36	43.64	4.00	-8.02405
4	0.26	56.60	39.40	4.00	-7.58067
5	0.24	59.21	36.79	4.00	-7.41213
6	0.22	60.87	35.13	4.00	-7.34396
7	0.22	61.94	34.06	4.00	-7.31540
8	0.21	62.64	33.36	4.00	-7.30317
9	0.21	63.10	32.90	4.00	-7.29787
10	0.20	63.40	32.60	4.00	-7.29555
11	0.20	63.60	32.40	4.00	-7.29453
12	0.20	63.73	32.27	4.00	-7.29408
13	0.20	63.82	32.18	4.00	-7.29388
14	0.20	63.88	32.12	4.00	-7.29379
15	0.20	63.92	32.08	4.00	-7.29375
16	0.20	63.95	32.05	4.00	-7.29374

# Alternatives

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- Analytical solutions
- Generic maximization strategies
- Calculating second derivatives a useful complement, whatever method we use...

# Other Applications of the E-M Algorithm in Genetics

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- **Classic example:**
  - ABO blood group
- **Most common application:**
  - Haplotype frequency estimates

# The ABO blood group

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- Determines compatibility for transfusions
- Controlled by alleles of ABO gene
- 3 alternative alleles
  - A, B and O
- 6 possible genotypes,  $n(n + 1) / 2$ 
  - A/A, A/B, A/O, B/B, B/O, O/O



# ABO Blood Group II

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Phenotype	Antigen		Antibody	
	A	B	A	B
A	+	-	-	+
B	-	+	+	-
O	-	-	+	+
AB	+	+	-	-

There are only 4 possible phenotypes for the ABO blood group.

# Genotypes and Phenotypes

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<b>Genotype</b>	<b>Phenotype</b>
A/A	A
A/B	AB
A/O	A
B/B	B
B/O	B
O/O	O

# ABO Example

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- Data of Clarke et al. (1959)
  - *British Med J* 1:603-607
  - Reported excess of gastric ulcers in individuals with blood type O
- $n_A = 186$ ,  $n_B = 38$ ,  $n_{AB} = 36$ ,  $n_O = 284$

## Quick Exercises!

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- Write out the likelihood for these data...
- What are complete data categories?
- Express the complete data “counts” as a function of allele frequency estimates and the observed data...

The iterations give ...

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<b>Iteration</b>	<b><math>p_A</math></b>	<b><math>p_B</math></b>	<b><math>p_o</math></b>
<b>1</b>	0.300	0.200	0.500
<b>2</b>	.243	.074	.683
<b>3</b>	.228	.070	.700
<b>4</b>	.228	.070	.702
<b>5</b>	.228	.070	.702

# Alternatives to E-M...

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- Analytical solutions are not known for the general case
- Generic maximization strategies could be employed
- Could derive solutions using part of the data...
  - Would this be a good idea?

# The E-M Haplotyping Algorithm

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- Excoffier and Slatkin (1995)
  - *Mol Biol Evol* **12**:921-927
  - Provide a clear outline of how the algorithm can be applied to genetic data
- Combination of two strategies
  - E-M statistical algorithm for missing data
  - Counting algorithm for allele frequencies

# Original Application of the E-M Algorithm to A Genetic Problem

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- Ceppellini R, Siniscalco M and Smith CAB (1955) The Estimation of Gene Frequencies in a Random-Mating Population. *Annals of Human Genetics* **20**:97-115
- This was ~20 years before the E-M algorithm was formally outlined in the statistical literature!



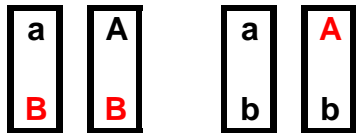
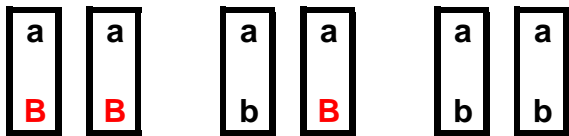
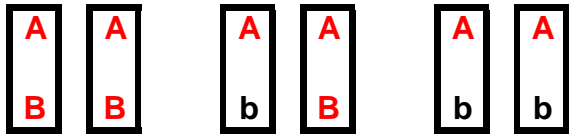
# Counting for Allele Frequencies

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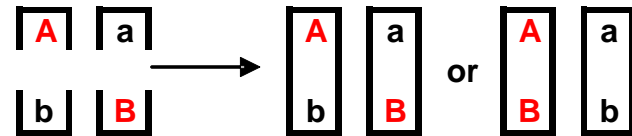
- For co-dominant markers, allele frequency typically carried out in very simple manner:
  - Count number of chromosomes (e.g.  $2N$ )
  - Count number of a alleles (e.g.  $n_a$ )
  - Allele frequency is simple proportion ( $n_a/2N$ )
- Haplotypes can't always be counted directly
  - Focusing on unambiguous genotypes introduces bias

# Counting Haplotypes for 2 SNPs

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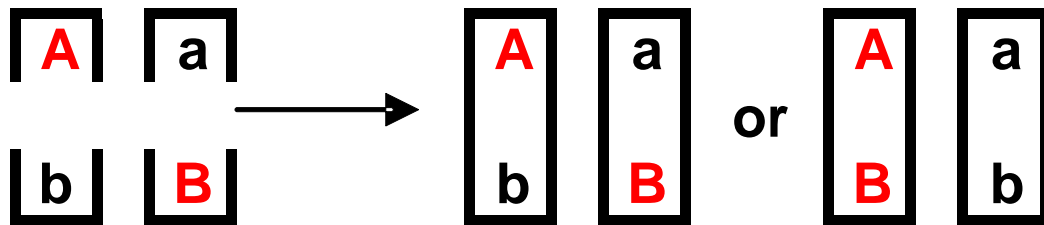
**Unambiguous Genotypes**  
Underlying Haplotype is Known



**Ambiguous Genotype**  
Multiple Underlying Genotypes Possible

# Probabilistic Interpretation

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Probability of first outcome:

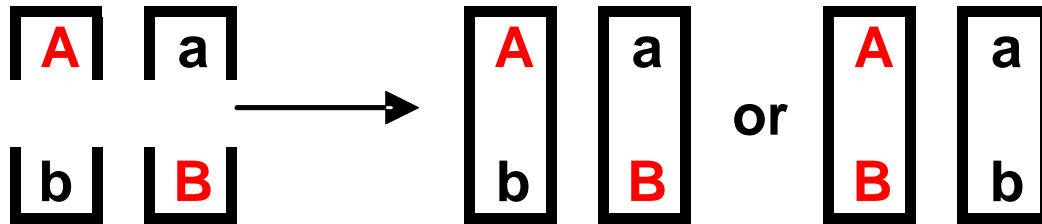
$$2 P_{Ab} P_{aB}$$

Probability of second outcome:

$$2 P_{AB} P_{ab}$$

# Probabilistic Interpretation

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For example, if:

$$P_{AB} = 0.3$$

$$P_{ab} = 0.3$$

$$P_{Ab} = 0.3$$

$$P_{aB} = 0.1$$

Probability of first outcome:

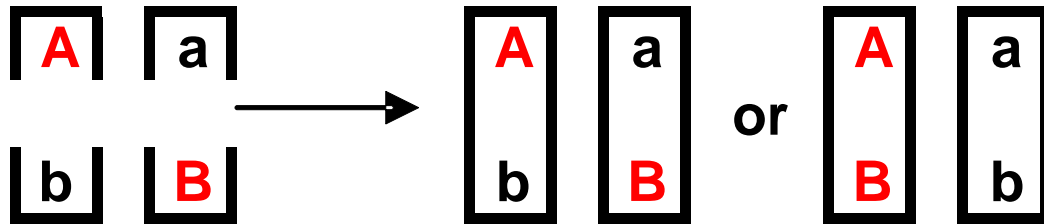
$$2 P_{Ab} P_{aB} = 0.06$$

Probability of second outcome:

$$2 P_{AB} P_{ab} = 0.18$$

# Probabilistic Interpretation II

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Conditional probability of first outcome:

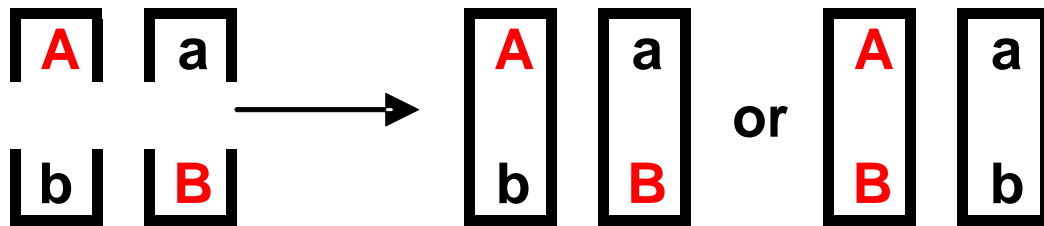
$$2 P_{Ab} P_{aB} / (2 P_{Ab} P_{aB} + 2 P_{AB} P_{ab})$$

Conditional probability of second outcome:

$$2 P_{AB} P_{ab} / (2 P_{Ab} P_{aB} + 2 P_{AB} P_{ab})$$

# Probabilistic Interpretation II

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For example, if:

$$P_{AB} = 0.3$$

$$P_{ab} = 0.3$$

$$P_{Ab} = 0.3$$

$$P_{aB} = 0.1$$

Conditional probability of first outcome:

$$2 P_{Ab} P_{aB} / (2 P_{Ab} P_{aB} + 2 P_{AB} P_{ab}) = 0.25$$

Conditional probability of second outcome:

$$2 P_{AB} P_{ab} / (2 P_{Ab} P_{aB} + 2 P_{AB} P_{ab}) = 0.75$$

# E-M Algorithm For Haplotyping

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1. “Guesstimate” haplotype frequencies
2. Use current frequency estimates to replace ambiguous genotypes with fractional counts of phased genotypes
3. Estimate frequency of each haplotype by counting
4. Repeat steps 2 and 3 until frequencies are stable

# Computational Cost (for SNPs)

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- Consider sets of  $m$  unphased genotypes
  - Markers  $1..m$
- If markers are bi-allelic
  - $2^m$  possible haplotypes
  - $2^{m-1} (2^m + 1)$  possible haplotype pairs
  - $3^m$  distinct observed genotypes
  - $2^{n-1}$  reconstructions for  $n$  heterozygous loci



# E-M Algorithm for Haplotyping

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- Cost grows rapidly with number of markers
- Typically appropriate for  $< 25$  SNPs
  - Fewer microsatellites
- Fully or partially phased individuals contribute most of the information

# Other Common Applications

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- E-M Algorithm also commonly used for:
  - Estimating recombination fractions
  - Defining genotype intensity clusters
  - Finding sub-populations and their allele frequencies

# Today:

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- The E-M algorithm in genetics
- Outline the approach
- Examined specific examples

## Next Lecture ...

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- E-M algorithm for Haplotyping
- Historical Alternatives
- Recent Enhancements and Alternatives
- Hypothesis testing

## Recommended Reading

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- Excoffier and Slatkin (1995)  
*Mol Biol Evol* **12**:921-927
- Introduces the E-M algorithm in the context of haplotyping