# Linkage Disequilibrium (LD)

 Alleles at one locus are correlated with alleles at a second locus on a population level.

AAAAAAAaaaaBBBBBBbbbbbbb

If you sample a haplotype at random from the population, does knowing the allele at the first locus of this haplotype give you information about the allele at the second locus?

# Linkage Disequilibrium

• One usual measure of LD is:

$$\mathbf{D}_{\mathbf{A}\mathbf{B}} = \mathbf{P}_{\mathbf{A}\mathbf{B}} - \mathbf{p}_{\mathbf{A}} \mathbf{p}_{\mathbf{B}}$$

AAAAAAAaaaaBBBBBBBbbbbbb

• 
$$P_A = 8/12$$

- $P_B = 6/12$
- $P_{AB} = 6/12$
- $D_{AB} = 1/6$

# Linkage Disequilibrium

• One usual measure of LD is:

$$\mathbf{D}_{\mathbf{A}\mathbf{B}} = \mathbf{P}_{\mathbf{A}\mathbf{B}} - \mathbf{p}_{\mathbf{A}} \mathbf{p}_{\mathbf{B}}$$

AAAAAAAaaaaBBBBBBBbbbbbb

- LD is a measure of extant haplotypes
  - Estimates do not rely on or measure inheritance
  - LD does not measure how often alleles are transmitted together

## Properties of LD

• One usual measure of LD is:

$$\mathbf{D}_{AB} = \mathbf{P}_{AB} - \mathbf{p}_{A} \mathbf{p}_{B}$$
  
•  $\mathbf{D}_{AB} = -\mathbf{D}_{Ab} = -\mathbf{D}_{aB} = \mathbf{D}_{ab} \quad \leftarrow \text{ can denote as } \mathbf{D}$ 



## Bounds on LD

- $0 \leq p_A p_B + D_{AB} \leq \min(p_A, p_B)$
- $0 \le p_A p_b D_{AB} \le \min(p_A, p_b)$
- $0 \le p_a p_B D_{AB} \le \min(p_a, p_B)$
- $0 \leq p_a p_b + D_{AB} \leq \min(p_a, p_b)$
- max  $(-p_A p_B, -p_a p_b) \leq D_{AB} \leq \min(p_A p_b, p_a p_B)$

#### Other Measures of LD

 D' ("D-prime") normalize D by the maximum value it can obtain:

$$D'_{AB} = \begin{cases} D_{AB} / [max(-p_A p_B, -p_a p_b)] \text{ if } D_{AB} < 0 \\ D_{AB} / [min (p_A p_b, p_a p_B)] \text{ if } D_{AB} > 0 \end{cases}$$

• 
$$r_{AB}^2 = \frac{D_{AB}^2}{p_A p_B p_a p_b}$$

(squared correlation coefficient, r)

# LD can be created by

- Mixing of populations.
- Population substructure (non-random mating within populations).
- Mutations creating new haplotypes.
- Selection favoring certain alleles.
- Founder effects.
- Genetic drift.

## LD decays over time

- LD decays because of recombination
- $D_{AB}(g) = (1 r)^g \times D_{AB}(0)$ 
  - r = recombination rate, g = number of generations
- Although this predicts the expected value of LD over time, there is a large variance around the mean.
- Populations with similar starting values can be quite different after time.

Reminder: meiosis & recombination (extremely simplified)



meiosis I

meiosis II

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# **Consider Two Populations**

Population 1 haplotypes:

<b>A</b>	<b>A</b>	A	A	A	A	A
B	B	B	B	B	B	B

Population 2 haplotypes:

a	a	a	a	a	a	a
b	b	b	b	b	b	b

# If These Populations Mix

Some individuals directly after mixing:



Their possible children:







# Haplotype Possibilities for the Grandchildren



children

Make sure you don't make unfounded assumptions about the probability of seeing any given haplotype! (e.g. 0.5)



# Possible full-chromosome haplotypes





# After a few generations ...



# After many generations (many recombination events)



## Linkage Disequilibrium versus Generations Since its Creation



#### Linkage versus LD

#### Linkage is *defined by* recombination.

- Recombination occurs during meiosis and thus, in turn, is observed via inheritance.
- Loci whose recombination rates are < 0.5 are linked.
- Loci whose recombination rates are = 0.5 are unlinked.
- Linkage is measured via correlated *transmission* of alleles.

#### LD is affected by recombination (over time).

- LD measures correlation between alleles in a population.
- LD is based on extant haplotypes and *estimates do not rely on or measure inheritance*.
- LD breaks down over time via recombination.
- LD does *not* measure correlated *transmission* of alleles.