## Linkage Disequilibrium

**Biostatistics 666** 

#### Logistics: Office Hours

Office hours on Mondays at 4 pm.

Room 4614
 School of Public Health Tower

#### Previously ...

- Basic properties of a locus
  - Allele Frequencies
  - Genotype Frequencies
- Hardy-Weinberg Equilibrium
  - Relationship between allele and genotype frequencies that holds for most genetic markers
- Exact Tests for Hardy-Weinberg Equilibrium

#### Today ...

We'll consider properties of pairs of alleles

Haplotype frequencies

Linkage equilibrium

Linkage disequilibrium

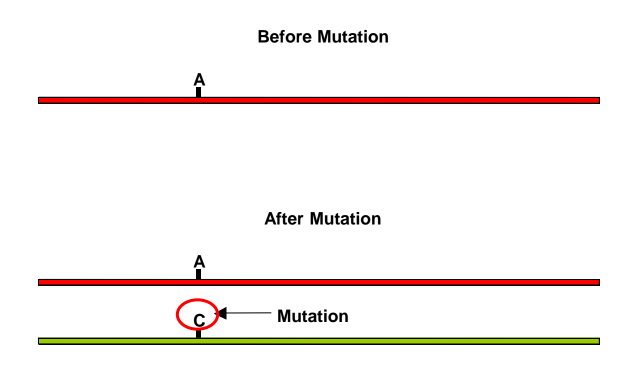
# Let's consider the history of two neighboring alleles...

## Alleles that exist today arose through ancient mutation events...

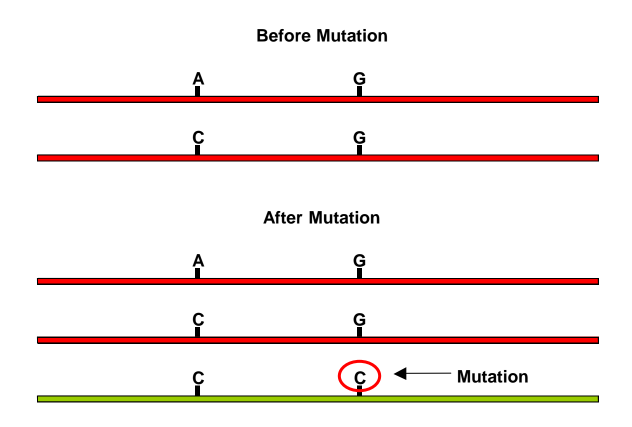
#### **Before Mutation**

#### **After Mutation**

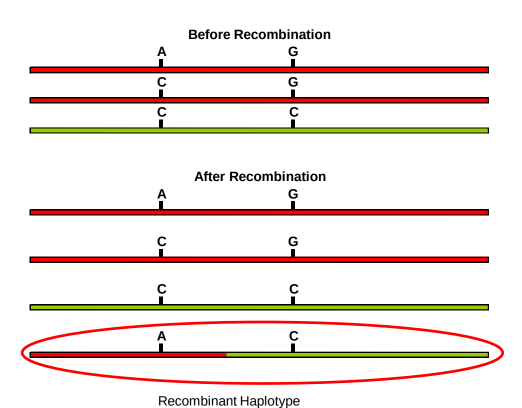
## Alleles that exist today arose through ancient mutation events...



## One allele arose first, and then the other...



## Recombination generates new arrangements for ancestral alleles



#### Linkage Disequilibrium

- Chromosomes are mosaics
- Extent and conservation of mosaic pieces depends on
  - Recombination rate
  - Mutation rate
  - Population size
  - Natural selection
- Combinations of alleles at very close markers reflect ancestral haplotypes



Why is linkage disequilibrium important for gene mapping?

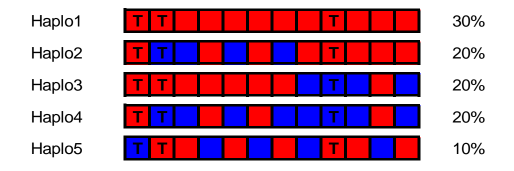
## Association Studies and Linkage Disequilibrium

 If all polymorphisms were independent at the population level, association studies would have to examine every one of them...

 Linkage disequilibrium makes tightly linked variants strongly correlated producing cost savings for association studies

### Tagging SNPs

- In a typical short chromosome segment, there are only a few distinct haplotypes
- Carefully selected SNPs can determine status of other SNPs



## Basic Descriptors of Linkage Disequilibrium

#### Commonly Used Descriptors

- Haplotype Frequencies
  - The frequency of each type of chromosome
  - Contain all the information provided by other summary measures
- Commonly used summaries
  - D
  - D'
  - $r^2$  or  $\Delta^2$

#### Haplotype Frequencies

### Linkage Equilibrium Expected for Distant Loci

$$p_{AB} = p_{A}p_{B}$$

$$p_{Ab} = p_{A}p_{b} = p_{A}(1 - p_{B})$$

$$p_{aB} = p_{a}p_{B} = (1 - p_{A})p_{B}$$

$$p_{ab} = p_{a}p_{b} = (1 - p_{A})(1 - p_{B})$$

## Linkage Disequilibrium Expected for Nearby Loci

$$p_{AB} \neq p_A p_B$$

$$p_{Ab} \neq p_A p_b = p_A (1 - p_B)$$

$$p_{aB} \neq p_a p_B = (1 - p_A) p_B$$

$$p_{ab} \neq p_a p_b = (1 - p_A) (1 - p_B)$$

### Disequilibrium Coefficient D<sub>AB</sub>

$$D_{AB} = p_{AB} - p_{A}p_{B}$$

$$p_{AB} = p_{A}p_{B} + D_{AB}$$

$$p_{AB} = p_{A}p_{B} - D_{AB}$$

$$p_{AB} = p_{A}p_{B} - D_{AB}$$

$$p_{AB} = p_{A}p_{B} - D_{AB}$$

$$p_{AB} = p_{A}p_{B} + D_{AB}$$

### D<sub>AB</sub> is hard to interpret

- Sign is arbitrary ...
  - A common convention is to set A, B to be the common allele and a, b to be the rare allele

- Range depends on allele frequencies
  - Hard to compare between markers

### What is the range of $D_{AB}$ ?

- What are the maximum and minimum possible values of D<sub>AB</sub> when
  - $p_A = 0.3$  and  $p_B = 0.3$
  - $p_A = 0.2$  and  $p_B = 0.1$
- Can you derive a general formula for this range?

#### D' - A scaled version of D

$$D'_{AB} = \begin{cases} \frac{D_{AB}}{\min(p_{A}p_{B}, p_{a}p_{b})} & D_{AB} < 0\\ \frac{D_{AB}}{\min(p_{A}p_{b}, p_{a}p_{b})} & D_{AB} > 0 \end{cases}$$

- Ranges between –1 and +1
  - More likely to take extreme values when allele frequencies are small
  - ±1 implies at least one of the observed haplotypes was not observed

#### More on D'

#### • Pluses:

- D' = 1 or D' = -1 means no evidence for recombination between the markers
- If allele frequencies are similar, high D' means the markers are good surrogates for each other

#### • Minuses:

- D' estimates inflated in small samples
- D' estimates inflated when one allele is rare

#### $\Delta^2$ (also called $r^2$ )

$$\Delta^{2} = \frac{D_{AB}^{2}}{p_{A}(1 - p_{A})p_{B}(1 - p_{B})}$$
$$= \frac{\chi^{2}}{2n}$$

- Ranges between 0 and 1
  - 1 when the two markers provide identical information
  - 0 when they are in perfect equilibrium
- Expected value is 1/2n

#### More on r<sup>2</sup>

- $r^2 = 1$  implies the markers provide exactly the same information
- The measure preferred by population geneticists
- Measures loss in efficiency when marker A is replaced with marker B in an association study
  - With some simplifying assumptions (e.g. see Pritchard and Przeworski, 2001)

# When does linkage equilibrium hold?

### Equilibrium or Disequilibrium?

- We will present simple argument for why linkage equilibrium holds for most loci
- Balance of factors
  - Genetic drift (a function of population size)
  - Random mating
  - Distance between markers
  - •

#### Why Equilibrium is Reached...

- Eventually, random mating and recombination should ensure that mutations spread from original haplotype to all haplotypes in the population...
- Simple argument:
  - Assume fixed allele frequencies over time

### Generation t, Initial Configuration

Assume arbitrary values for the allele frequencies and disequilibrium coefficient

#### Generation t+1, Without Recombination

Haplotype Frequencies Remain Stable Over Time Outcome has probability 1- R

#### Generation t+1, With Recombination

Haplotype Frequencies Are Function of Allele Frequencies Outcome has probability R

#### Generation t+1, Overall

$$B$$
  $b$   $A$   $p_{A}p_{B} + (1-r)D_{AB}$   $p_{A}p_{b} - (1-r)D_{AB}$   $p_{A}$   $p_{A}p_{b} - (1-r)D_{AB}$   $p_{a}$   $p_{a}p_{b} + (1-r)D_{AB}$   $p_{a}$   $p_{b}$ 

Disequilibrium Decreases...

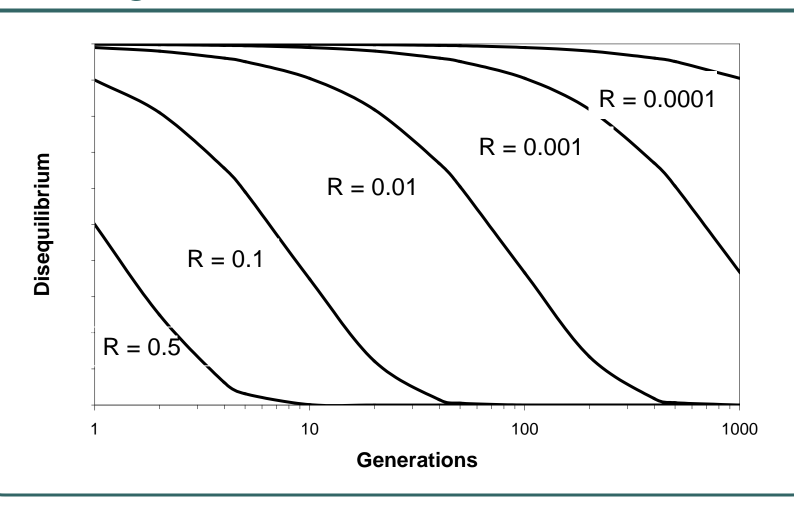
#### Recombination Rate (R)

 Probability of an odd number of crossovers between two loci

 Proportion of time alleles from two different grand-parents occur in the same gamete

 Increases with physical (base-pair) distance, but rate of increase varies across genome

### Decay of D with Time



#### Predictions

- Disequilibrium will decay each generation
  - In a large population
- After t generations...
  - $D_{AB}^{t} = (1-R)^{t}D_{AB}^{0}$
- A better model should allow for changes in allele frequencies over time...

#### Linkage Equilibrium

 In a large random mating population haplotype frequencies converge to a simple function of allele frequencies

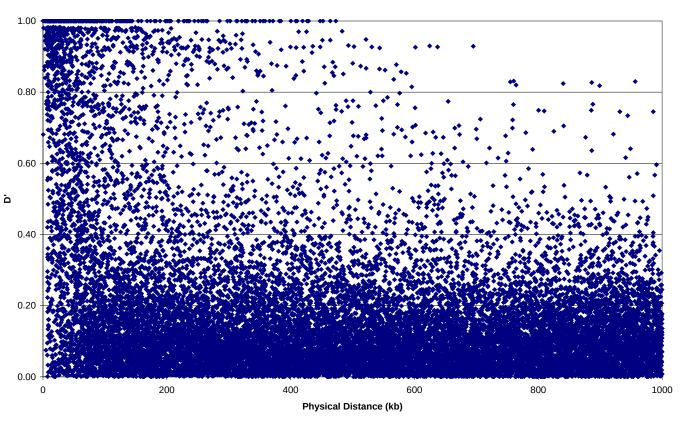
## Some Examples of Linkage Disequilibrium Data

## Summary of Disequilibrium in the Genome

• How much disequilibrium is there?

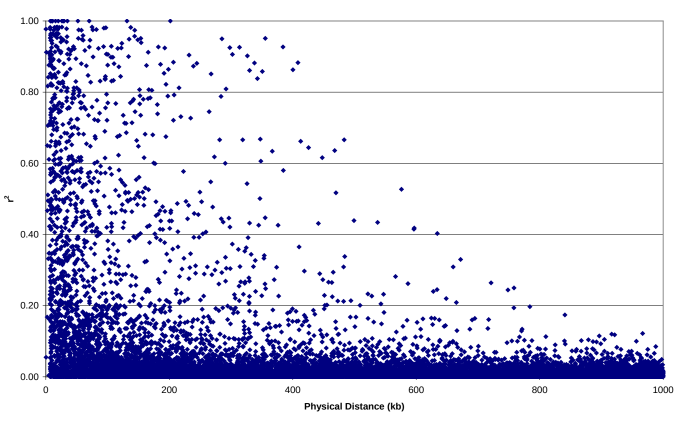
What are good predictors of disequilibrium?

## Raw |D'| data from Chr22



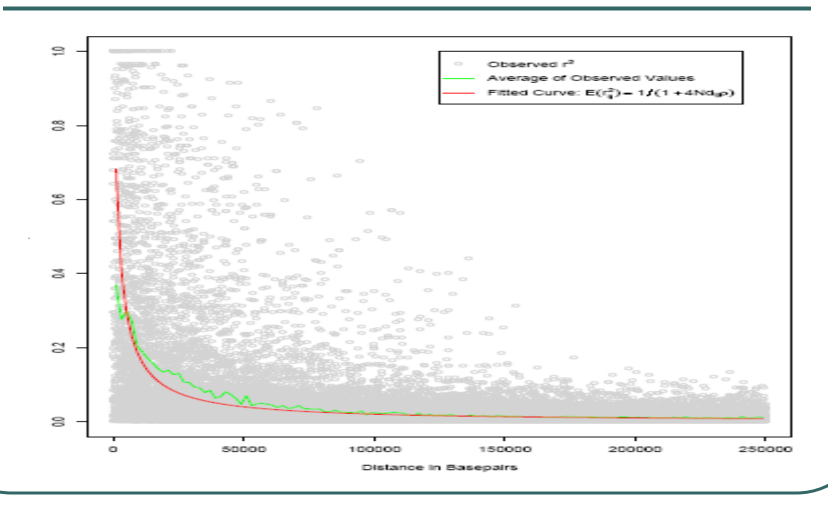
Dawson et al, Nature, 2002

### Raw $\Delta^2$ data from Chr22

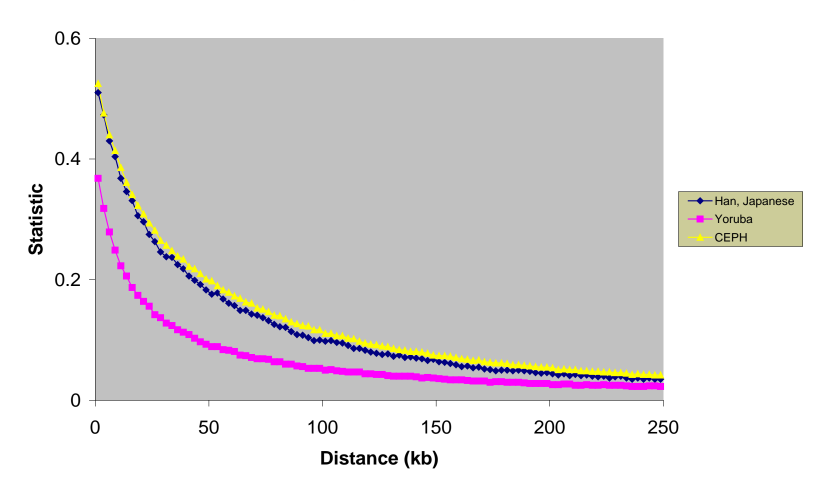


Dawson et al, Nature, 2002

## Summarizing Disequilibrium



### Comparing Populations ...



LD extends further in CEPH and the Han/Japanese than in the Yoruba

International HapMap Consortium, *Nature*, 2005

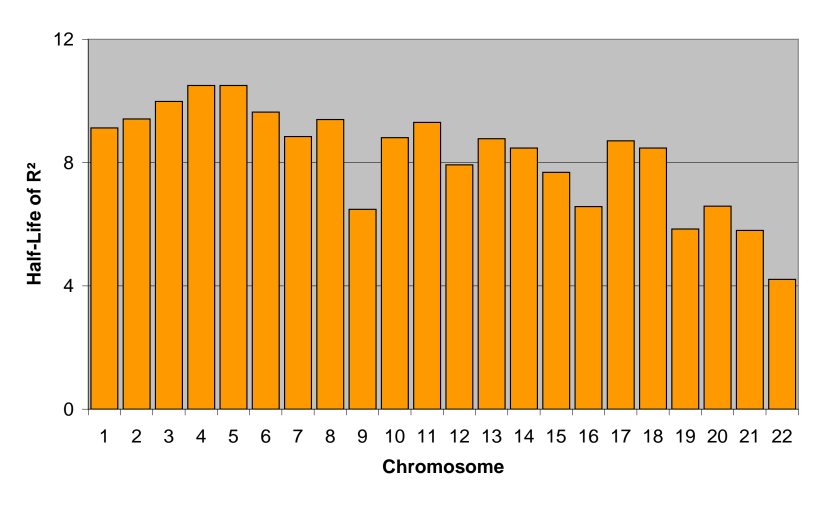
## Variation in Linkage Disequilibrium Along The Genome

## Comparing Genomic Regions ...

 Rather than compare curves directly, it is convenient to a pick a summary for the decay curves

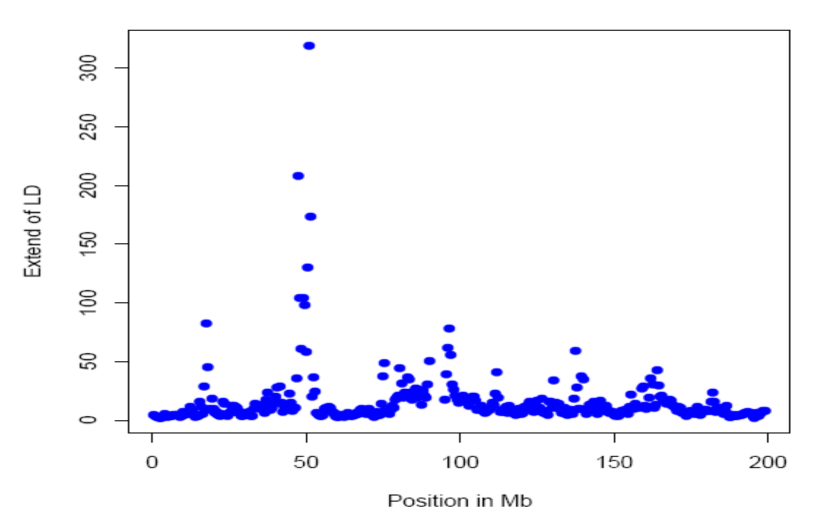
 One common summary is the distance at which the curve crosses a threshold of interest (say 0.50)

#### **Extent of Linkage Disequilibrium**



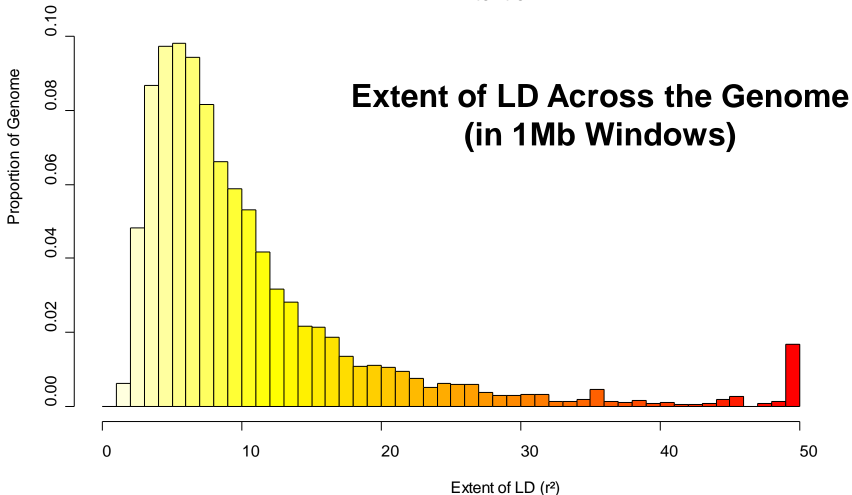
LD extends further in the larger chromosomes, which have lower recombination rates

#### Chromosome 3



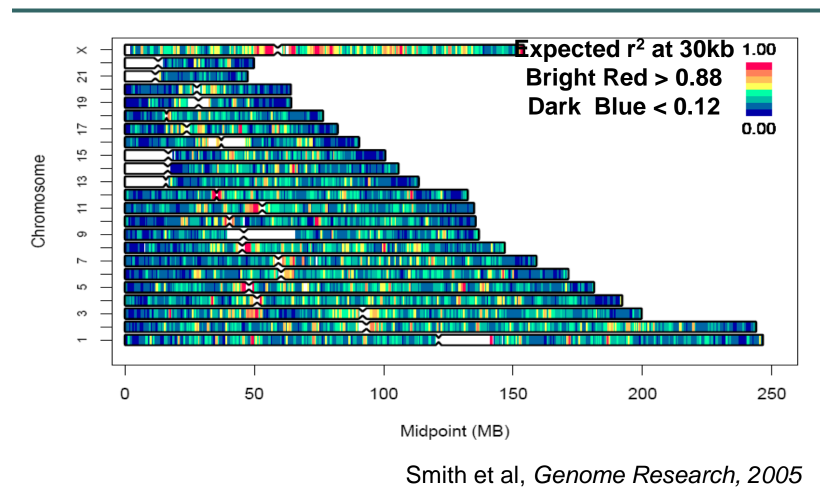
But within each chromosome, there is still huge variability!

#### **Extent of LD**

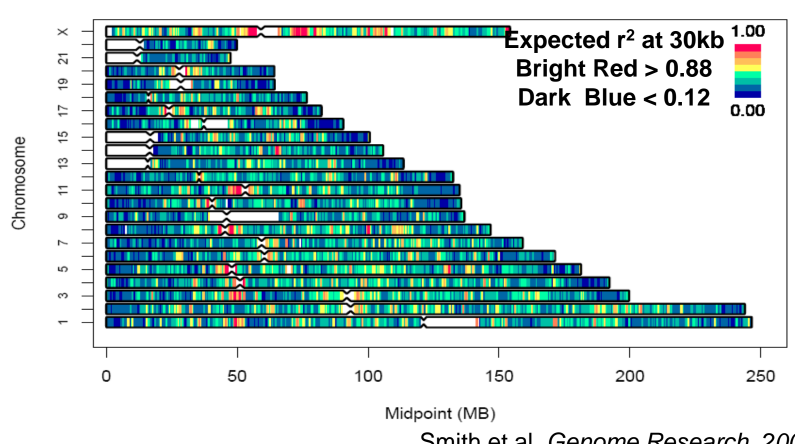


Average Extent: 11.9 kb
Median Extent: 7.8 kb
10<sup>th</sup> percentile: 3.5 kb
90<sup>th</sup> percentile: 20.9 kb

## Genomic Variation in LD (CEPH)

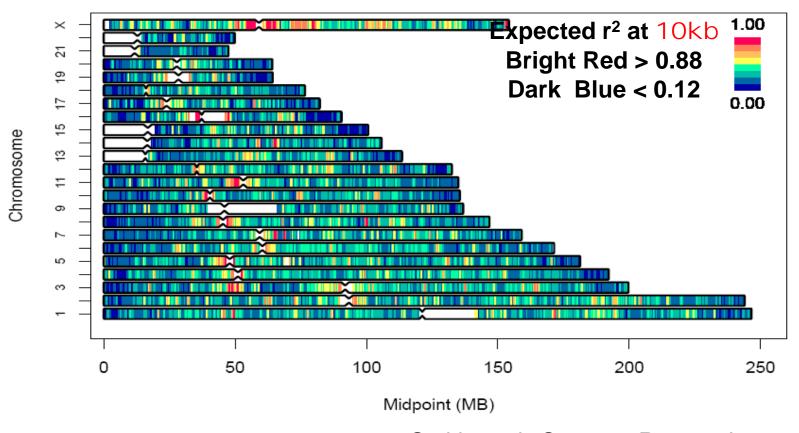


### Genomic Variation in LD (JPT + CHB)



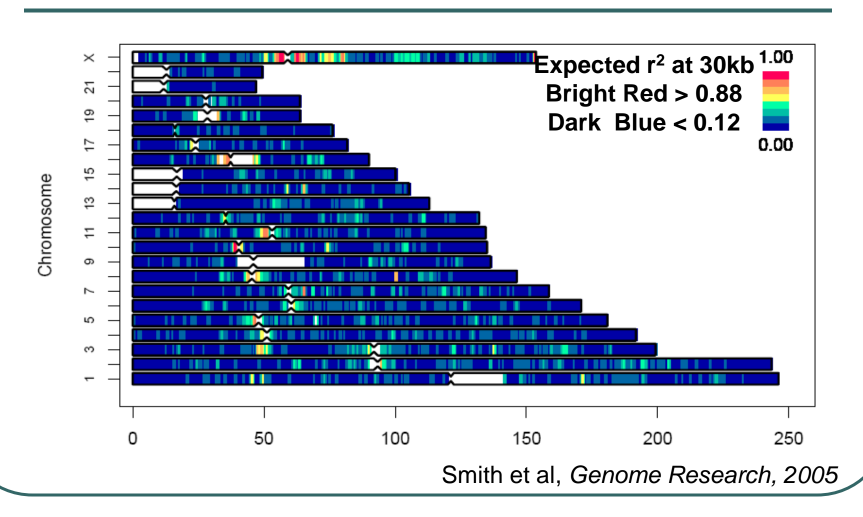
Smith et al, Genome Research, 2005

## Genomic Variation in LD (YRI)



Smith et al, Genome Research, 2005

## Genomic Distribution of LD (YRI)



## Sequence Composition vs. LD (some selected comparisons)

	Genome Quartiles, Defined Using LD						
	(Low LD)			<b>(</b> l	High LD)		
G	enome	Q1	Q2	Q3	Q4	Trend	
Basic Sequence Features							
GC Bases (%)	40.8	43.5	41.0	39.6	39.0	Decreases With LD	
Bases in CpG Islands (%)	0.7	0.9	0.7	0.6	0.6	Decreases With LD	
Polymorphism ( $\Pi * 10,000$ )	10.1	11.9	10.6	9.6	8.3	Decreases With LD	
Genes and Related Features							
Known Genes (per 1000 kb)	6.4	6.6	6.1	6.2	6.7	U shaped	
Genic Bases (Exon, Intron, UTR, %)	38.5	37.6	34.6	36.0	45.9	U shaped	
Coding Bases (%)	1.2	1.1	1.0	1.1	1.4	U shaped	
Conserved Non-Coding Sequence (%)	1.4	1.6	1.5	1.3	1.1	Decreases with LD	
Repeat Content							
Total Bases in Repeats (%)	47.9	44.2	46.4	48.6	52.3	Increases with LD	
Bases in LINE repeats (%)	20.9	16.5	19.9	22.4	24.9	Increases with LD	
Bases in SINE repeats (%)	13.6	14.7	13.1	12.6	14.0	U shaped	

Smith et al, Genome Research, 2005

# Gene Function in Regions of High and Low Disequilibrium

#### **Annotaated**

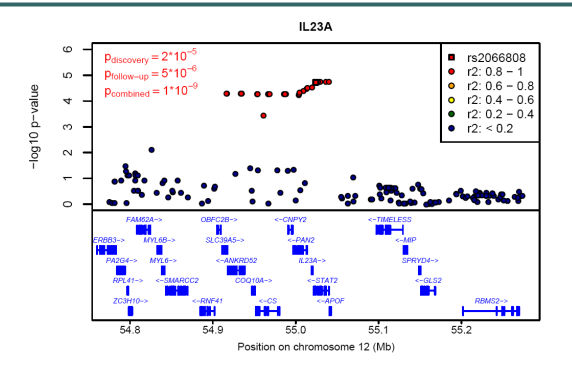
Gene Function (GO Term)	Genes	Low LD	High LD	χ²	P-value
All Swissprot Entries Examined	7520	2305	2045	-	_
DNA metabolism	366	74	139	35.37	<.0001
Immune Response	622	232	94	34.36	<.0001
Cell cycle	493	119	177	26.24	<.0001
Protein Metabolism	1193	318	375	23.33	<.0001
Organelle Organization and Biogenesis	444	107	152	19.65	<.0001
Intracellular Transport	263	56	95	19.61	<.0001
Organogenesis	805	294	162	16.49	0.00005
Cell Organization and Metabolism	545	138	178	16.43	0.00005
RNA Metabollism	208	41	71	15.33	0.00009

Results from a comparison of the distribution of 40 most common gene classifications in the GENE Ontology Database

# Implications for Association Studies

#### Linkage Disequilibrium in Association Studies:

### Psoriasis and IL23A Example



Multiple nearby SNPs show evidence for association with psoriasis. The SNPs are all in linkage disequilibrium, so it is hard to pinpoint causal SNP.

Nair et al, Nature Genetics, 2009

#### Linkage Disequilibrium in Association Studies:

## Tag SNP Picking

 Many nearby SNPs will typically provide similar evidence for association

- To decrease genotyping costs, most association studies will examine selected "tag SNPs" in each region
- The most common tagging strategy focuses on pairwise r<sup>2</sup> between SNPs

#### Linkage Disequilibrium in Association Studies:

## Pairwise Tagging Algorithm

- Select an r<sup>2</sup> threshold, typically 0.5 or 0.8
  - SNPs with r<sup>2</sup> above threshold can serve as proxies for each other
- For each marker being considered, count the number of SNPs with r<sup>2</sup> above threshold
- Genotype SNP with the largest number of pairwise "proxies"
- Remove SNP and all the SNPs it tags from consideration
- Repeat the previous three steps until there are no more SNPs to pick or genotyping budget is exhausted

Carlson et al, AJHG, 2004

## Potential Number of tag SNPs

Table 3 | Number of tag SNPs required to capture common (MAF  $\geq$  0.05) Phase II SNPs

Threshold	YRI	CEU	CHB+JPT
$r^2 \ge 0.5$ $r^2 \ge 0.8$ $r^2 = 1.0$	627,458	290,969	277,831
	1,093,422	552,853	520,111
	1,616,739	1,024,665	1,078,959

Current tag SNP panels typically examine 500,000 – 1,000,000 SNPs for a cost of \$200 - \$500 per sample.

It is anticipated that future panels will allow for as many as 5 million SNPs.

The International HapMap Consortium, Nature, 2007

### Today ...

Basic descriptors of linkage disequilibrium

 Learn when linkage disequilibrium is expected to hold (or not!)

## Additional Reading I

- Dawson E et al (2002). A first-generation linkage disequilibrium map of human chromosome 22.
   Nature 418:544-548
- The International HapMap Consortium. (2005). A haplotype map of the human genome. Nature 437:1299-320
- Carlson CS et al (2004). Selecting a maximally informative set of single-nucleotide polymorphisms for association analyses using linkage disequilibrium. Am J Hum Genet 74:106-120

## Additional Reading II

 Cardon and Bell (2001) Association study designs for complex diseases.
 Nature Reviews Genetics 2:91-99

- Surveys important issues in analyzing population data.
- Illustrates shift from focus on linkage to association mapping for complex traits.