# Linkage Introduction

#### Law of Independent Assortment

- Mendel's Second Law (Law of Independent Assortment) :
  - The segregation of the genes for one trait is independent of the segregation of genes for another trait, i.e., when genes segregate, they do so independently
- This law essentially states that during gamete formation, the segregation of one gene is independent of the other gene
- This "law" is frequently violated and is only true for loci/genes that are unlinked.

#### Recombination

- When a gamete is passed down, the chromosome inherited by an offspring from a parent is actually a mosaic of the parent's two chromosomes.
- Suppose we have two loci on the same chromosome, locus 1 and locus 2, where locus 1 has alleles A1 and A2, and locus 2 has alleles B1 and B2.
- In the example below, phase is known and is (A1,B1) and (A2,B2).
- If the genes are closely linked, a gamete is much more likely to contain (A1,B1) or (A2,B2), which are "non-recombinants."
- If there is recombination, a gamete will contain (A1, B2) or (A2,B1), but this is less likely if the loci are linked.

#### **Recombination Fraction**

- Two loci that are unlinked follow Mendel's Second Law, and all possible gametes for a parent are produced with equal frequency.
- When loci are physically located close to one another on a chromosome, there is a deviation from this relationship. This deviation is summarized by the recombination fraction.
- The recombination fraction is often denoted by  $\theta$  where  $0 \leqslant \theta \leqslant \frac{1}{2}$
- P(recombinant gamete)=  $\theta$
- If  $\theta < \frac{1}{2}$ , the loci are said to be linked or in genetic linkage
- When loci are completely linked,  $\theta = 0$
- Two loci are said to be unlinked if  $\theta = \frac{1}{2}$ .
- Note that if two loci are on different chromosome, then  $\theta = \frac{1}{2}$ .



#### Linkage in a simple genetic cross

- In the early 1900's, Bateson and Punnet conducted genetic studies using sweet peas. They studied two characters:
  - Petal color which has two alleles: P (purple) and p (red), where P is dominant.
  - Pollen grain shape has two alleles: L (elongated) and I (disc-shaped), where L is dominant

$$\begin{array}{c} \mathsf{PPLL} \times \mathsf{ppll} \\ \downarrow \\ \mathsf{PpLl} \end{array} \qquad \mathsf{F1}$$

- Plants in the F1 generation were intercrossed: PpLl X PpLl.
- According to Mendel's Second Law, during gamete formation, the segregation of one gene pair is independent of another gene pair.

# **Sweet Peas Linkage Example**

F2	PL	PI	pL	pl
PL	Purple/Long	Purple/Long	Purple/Long	Purple/Long
PI	Purple/Long	Purple/Disc	Purple/Long	Purple/Disc
pL	Purple/Long	Purple/Long	Red/Long	Red/Long
pl	Purple/Long	Purple/Disc	Red/Long	Red/Disc

# **Sweet Peas Linkage Example**

 The expected relative frequencies in the F2 generation if the genes segregated independently are

	Elongated	Disc-Shaped
Purple	9	3
Red	3	1

 The observed frequencies in 381 plants in the F2 generation where

	Elongated	Disc-Shaped
Purple	284	21
Red	21	55

- The observed data clearly do not fit what is expected under the model.
- The explanation: the petal color gene and the gene for pollen grain shape are linked.
- Let  $\theta$  be the recombination fraction between the two genes. What is the probability of each possible plant type?

# Sweet Peas Linkage Example

		$\frac{1}{2}(1- heta)$	$\frac{1}{2}\theta$	$\frac{1}{2}\theta$	$\frac{1}{2}(1- heta)$
		PL	PI	pL	pl
$\frac{1}{2}(1-\theta)$	PL	Purple/Long	Purple/Long	Purple/Long	Purple/Long
$\frac{1}{2}\theta$	PI	Purple/Long	Purple/Disc	Purple/Long	Purple/Disc
$\frac{1}{2}\theta$	pL	Purple/Long	Purple/Long	Red/Long	Red/Long
$\frac{1}{2}(1-\theta)$	pl	Purple/Long	Purple/Disc	Red/Long	Red/Disc

- P(red, disc-shaped)=  $\frac{1}{4}(1-\theta)^2$
- P(red,elongated)=  $\left(\frac{1}{2}\theta\right)\left(\frac{1}{2}\theta\right) + \left(\frac{1}{2}\theta\right)\left(\frac{1}{2}(1-\theta)\right) + \left(\frac{1}{2}(1-\theta)\right)\left(\frac{1}{2}(\theta)\right)$
- P(purple, disc-shaped) and P(purple, elongated) are calculated similarly.
- We can form a likelihood for the data that is a function of the recombination fraction θ. We can find the value of θ that maximizes this likelihood.
- Likelihood will follow a multinomial distribution.



# Parametric Linkage Analysis

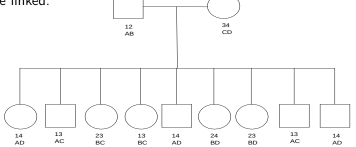
# Linkage Analysis

- Once aggregation and/or segregation studies established a genetic component for a phenotype of interest, parametric linkage analysis was the traditional approach used for Mendelian disease gene mapping since the 1970's
- Linkage analysis requires genetic marker data on pedigree.
- To illustrate linkage analysis, we will consider examples given by Suarez, B.K. and Cox, N.J. (1985)

#### **Nuclear Family Example**

- The figure below shows a large nuclear family segregating alleles from two loci: alleles at one of the loci are denoted by numbers while the alleles of the other are denoted by letters.
- Both of the parents are heterozygous at each locus and share no alleles in common, so the co-segregation of the alleles at the two loci can be unambiguously followed.

We are interested in determining whether or not the two loci are linked.



#### **Lod Scores**

- ullet LOD scores are calculated for recombination fraction heta values to determine if there is significant evidence for linkage
- ullet For a given value of heta, the lod score is

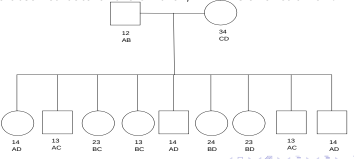
 $log_{10} \frac{P(\text{observed data assuming recombination fraction is } \theta)}{P(\text{observed data assuming recombination fraction is } .5)}$ 

- LOD stands for Log of ODds
- ullet Find the the value of heta that gives the maximum lod score
- Lod scores greater than 3 give evidence of linkage, and the null hypothesis of no linkage is rejected.
- How do you interpret a lod score equal to 3?
- Lod scores less than −2 give evidence that the loci are unlinked.



# **Nuclear Family Linkage Example**

- Can calculate a lod score for the large nuclear family. We only observe the genotypes at the two loci so the phase is unknown. Possible phase for the parents:
  - 1A 2B 3C 4D
  - 1A 2B 3D 4C
  - 1B 2A 3C 4D
  - 1B 2A 3D 4C
- Given each parental phase type, can obtain the probability of the observed data of the children, which is a function of  $\theta$



#### **Nuclear Family Linkage Example**

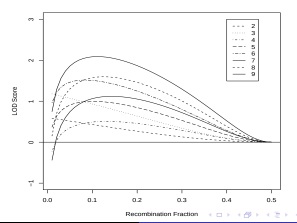
Phase	1A,2B,3C,4D	1A,2B,3D,4C	1B,2A,3C,4D	1B,2A,3D,4C
Phase Probability	.25	.25	.25	.25
Offspring Probability	$\left(\frac{1}{2}\right)^{18} (1-\theta)^{16} \theta^2$	$\left(\frac{1}{2}\right)^{18} \left(1-\theta\right)^9 \theta^9$	$\left(\frac{1}{2}\right)^{18} \left(1-\theta\right)^9 \theta^9$	$\left(\frac{1}{2}\right)^{18} (1-\theta)^2 \theta^{16}$

So, for  $\theta = .1$  the lod score is

$$\frac{.25(.9)^{16}(.1)^2 + .5(.9)^9(.1)^9 + .25(.9)^2(.1)^{16}}{.25(.5)^{18} + (.5)^{19} + .25(.5)^{18}}$$
$$= 2.08$$

# **Nuclear Family Linkage Example LOD Score Graph**

- For linkage analysis with nuclear families, data must be available on at least 2 offspring
- The figure below gives the lod score curves obtained for the large nuclear family according to the number of children included in the calculation.



# **Nuclear Family Linkage Example**

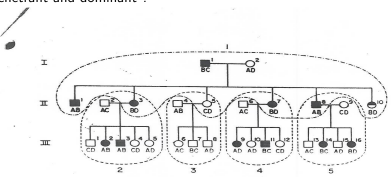
- The previous figure illustrates how the lod score curve changes as more information becomes available
- The lod score is always 0 at  $heta=rac{1}{2}$  since the odds ratio is 1
- The lod score calculated using the first 2 children and using the first 3 children steadily increases as  $\theta \longrightarrow 0$
- With the addition of the fourth child, the lod score curve changes from its monotonically increasing from as  $\theta \longrightarrow 0$  to one that increases as  $\theta$  moves away from  $\frac{1}{2}$
- Evidence for linkage becomes a little stronger with the addition of the fifth and sixth children, and decreases with the seventh child (due to an apparent maternal recombinant), and then increases with the remaining 2 children

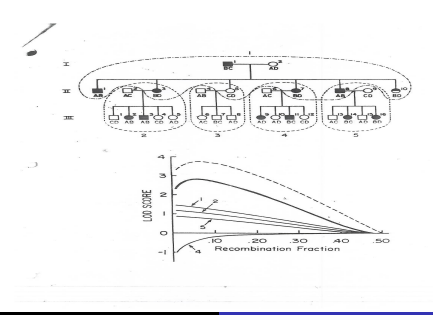
# **Nuclear Family Linkage Example**

- This nuclear family provides moderate evidence that the 2 loci are linked. At  $\hat{\theta}=0.11$ , the lod score curve reaches its maximum value of 2.09, indicating that the hypothesis of linkage with 11% recombination is about 123 times more likely than the hypothesis of no linkage
- Since the maximum lod score is in the range of -2 to 3, more families need to be sampled before a decision of  $\theta=\frac{1}{2}$  or  $\theta<\frac{1}{2}$  can be accepted or rejected.

- Linkage analysis for co-dominant loci in straightforward and a decision in favor or against the hypothesis of linkage can usually be reached with a few informative families.
- In general, however, nuclear families are less efficient than extended 3-generation pedigrees because extended pedigrees provide more information regarding phase

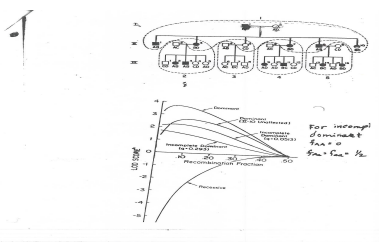
- Consider the 28 member 3-generation pedigree below
- We would like to determine if the locus with available genotype data is linked to a disease locus for which we do not know the location.
- What are the possible genotypes for the individuals in the pedigree if the disease is caused by a single locus that is fully penetrant and dominant?





- Nuclear families 1, 2, and 5 provide evidence for linkage. The lod score curves monotonically increasing as  $\theta \longrightarrow 0$  suggest that these families do not contain any recombinants. The different height of the lod score curves reflects that fact that larger nuclear families are more informative than smaller ones.
- Nuclear family 3 provides no information regarding linkage since neither partent is affected and at least one parent must be a double heterozygote to be informative.
- Nuclear family 4 provides slight evidence against the hypothesis of linkage.
- If the nuclear families were truly independent, then the lod scores could be summed, giving a maximum lod score of 2.81 at  $\hat{\theta}=0.05$ .
- When analyzing the pedigree as a whole, the maximum is also at  $\hat{\theta} = 0.05$  but with a lod score of 3.72.

 The plot below illustrates that misspecification of the mode of transmission of the disease affects the linkage analysis results.



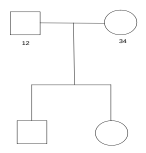
# Nonparametric Linkage Analysis

#### **Limitations of Parametric Linkage Analysis**

- We previously discussed parametric linkage analysis
- Genetic model for the disease must be specified: allele frequency parameters and penetrance parameters
- Lod scores results are highly sensitive to the assumed mode of transmission of the disease, which will generally be uknown
- Nonparametric linkage analysis methods does not make any assumptions about the disease model

#### **Sib Pair IBD Sharing Distribution**

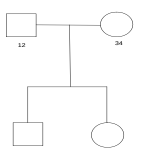
- Consider the nuclear family below with 2 siblings segregating alleles for a locus
- What is the probability of the siblings sharing 2, 1, or 0 alleles identical by descent (IBD)?



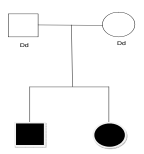
# Sib Pair IBD Sharing Distribution

Expected IBD Sharing

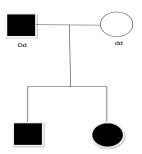
2 : 1 : 0 0.25 : 0.5 : 0.25



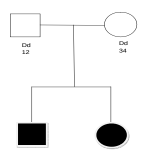
- Now consider a disease that is caused by a single locus.
- What would the allele sharing probabilities be for a sib pair at the disease locus?
- This depends on the mode of transmission of the disease.
   Assume for now that disease is caused by the D allele and D is recessive.

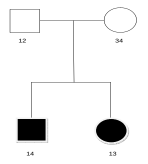


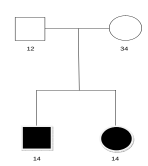
- Now assume that disease is caused by the D allele, and D is dominant.
- What would the allele sharing probabilities be for a sib pair at the disease locus?



- The location of the disease gene is unknown and we would like to determine if the locus is linked to the disease gene.
- If the locus is linked to the disease gene, then the expected IBD probabilities of sharing 2, 1, and 0 alleles IBD for sibs at the disease gene will not be .25, .5, and .25, respectively, regardless of the mode of inheritance of the disease.







- The null hypothesis: locus is transmitted independently of the disease locus D/d.
- Under the null, the expected IBD sharing for sibs is

2 : 1 : 0 0.25 : 0.5 : 0.25

- Under the alternative, the locus is linked to the disease locus, and as a result, the IBD sharing probabilities do not follow the distribution specified under the null hypothesis.
- If the null is false, then you should see an increase in affected sibs sharing either 1 or 2 alleles IBD.
- For example if disease is caused by a rare dominant allele and the locus is tightly linked to the disease gene, then expected IBD sharing for sibs might be around

2 : 1 : 0 0.5 : 0.5 : 0

 More realistic scenario: marker is very close to locus which influences risk of disease in a more subtle manner (heterogeneity, epistasis, gene-environment interaction)

```
2 : 1 : 0
0.35 : 0.45 : 0.2
```

#### **Model-Free Linkage Test**

- The Pearson chi-squared goodness of fit test is a sample way
  of comparing the observed counts of sib pairs sharing 0, 1 and
  2 alleles IBD with that expected under the null of no linkage.
- Let N be the number of affected sib pairs.
- Let  $n_i$  be the number of sib pairs that share i alleles IBD, where i = 0, 1, or 2.
- Under the null, what is the expected value of  $n_i$  for each i?
- Let the expected value of  $n_i$  under the null be  $E_{n_i}$ . The test statistic is:

$$X^{2} = \sum_{i=0}^{2} \frac{(n_{i} - E_{n_{i}})^{2}}{E_{n_{i}}}$$

• Under  $H_0$ , the  $X^2$  test statistic has an approximate  $\chi^2$  distribution with 2 degrees of freedom



#### **Extended Pedigrees**

- Nonparametric linkage analysis can also be used for extended pedigrees, not just nuclear families with affected sib pairs
- Can calculate the expected IBD sharing for more distant relatives
- What is the expected IBD sharing probabilities for first cousins under the null?

2 : 1 : 0 0 : 0.25 : 0.75

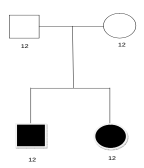
• What is the expected IBD sharing probabilities for second cousins under the null?

2 : 1 : 0

0 : 0.0625 : 0.9375

#### **IBD Allele Sharing Uncertainty**

- It may not be possible to determine exactly how many alleles a pair share IBD.
- In the example below, the affected sib pair could be sharing 2 or 0 alleles IBD, with each possibility having a probability of .5?



# **IBD Allele Sharing Uncertainty**

- Methods to allow for this uncertainty developed, e.g., Kruglyak et al. (1996), Kong and Cox (1997).
- Multi-point method that incorporates the genotypes of nearby loci
- Obtain a probability distribution of IBD sharing at the locus being tested for linkage

# **Allele Sharing Statistics**

• Allele sharing statistics *S* are often used for nonparametric linkage analysis. The general form of the statistics are

$$Z = \frac{S - \mu_0}{\sigma_0}$$

where  $\mu_0$  and  $\sigma_0$  are the expected value and variance of S, respectively, calculated under the null hypothesis. If a locus is not linked to a disease, Z will follow a standard Normal distribution.

- There are various types of allele sharing statistics
- S<sub>pairs</sub> counts, for each pair of affected relatives, the number of alleles shared IBD, and then sums that counts over all pairs of affected relatives.
- If all affected individuals in a pedigree have a common ancestor in the pedigree,  $S_{all}$  is the number of alleles shared IBD by all affected relatives.

#### Allele Sharing Statistics

- $S_{max}$  is the size of the largest group of related cases who all inherit the same allele IBD (high power for dominant disease alleles)
- McPeek (1999) showed that the optimal sharing statistic depends on the disease model