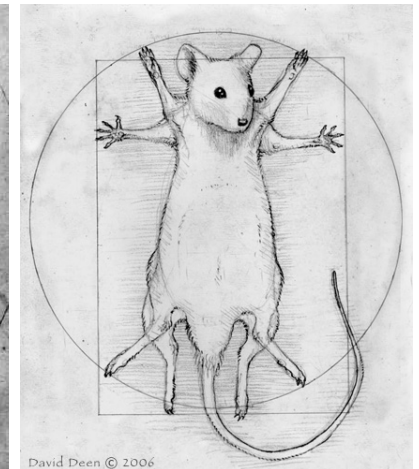
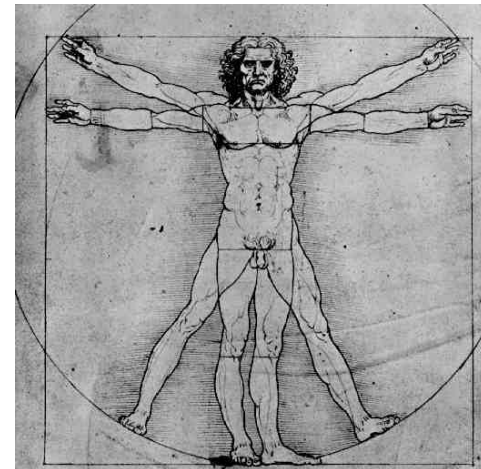


# Mapping multiple QTL in experimental crosses

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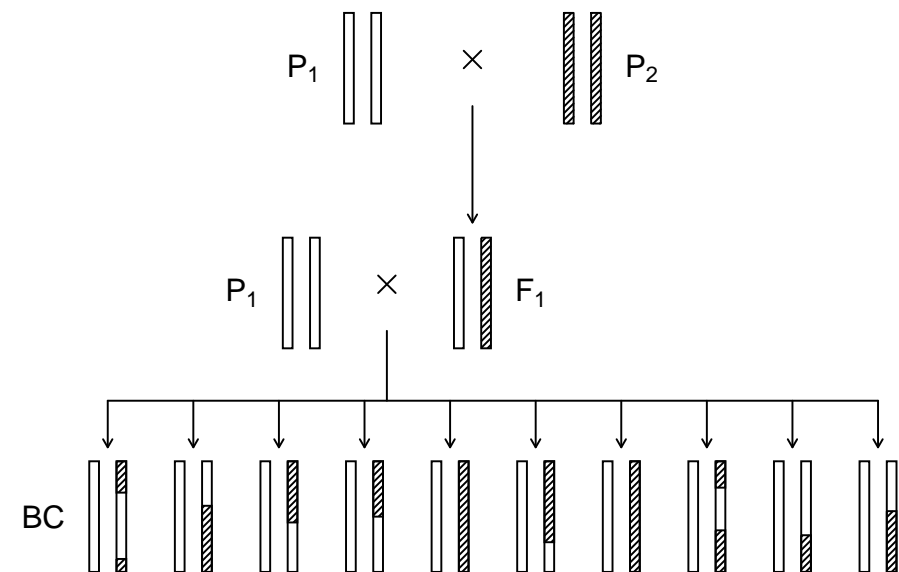


[www.daviddeen.com](http://www.daviddeen.com)

3

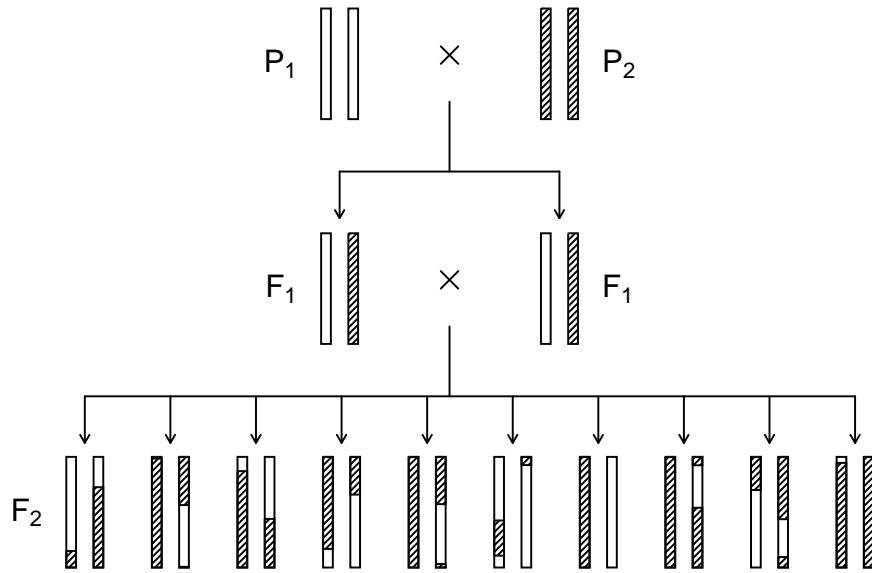


## Backcross



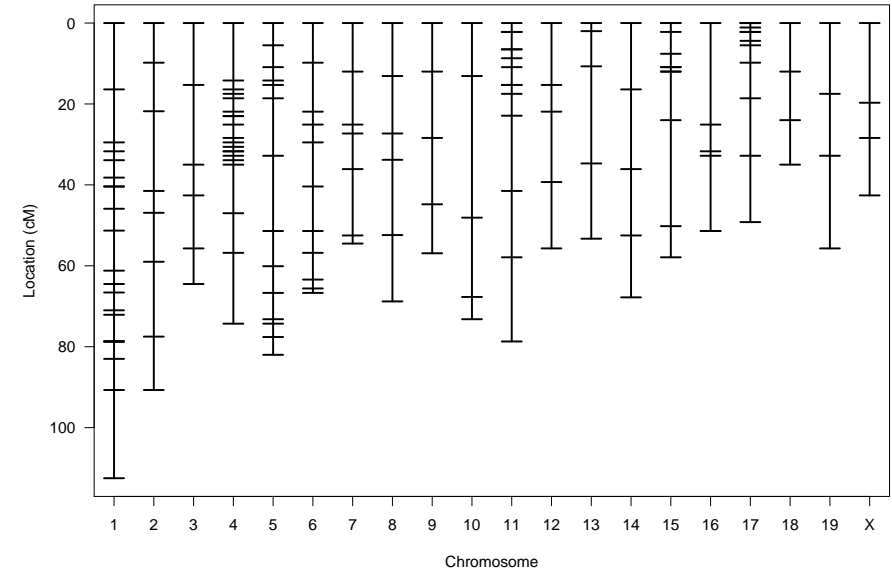
4

## Intercross



5

## Genetic map



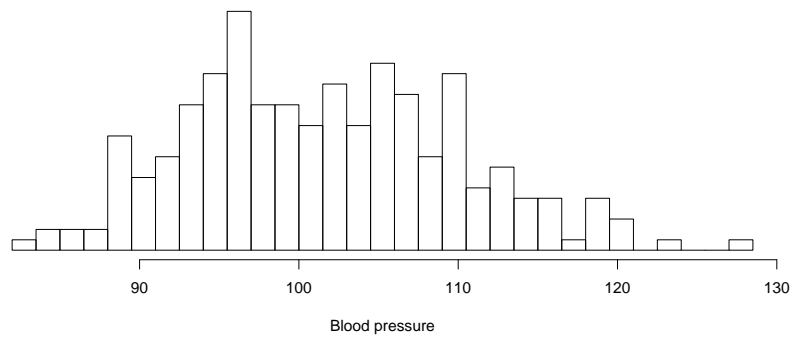
7

## Phenotype data

Sugiyama et al. Genomics 71:70-77, 2001

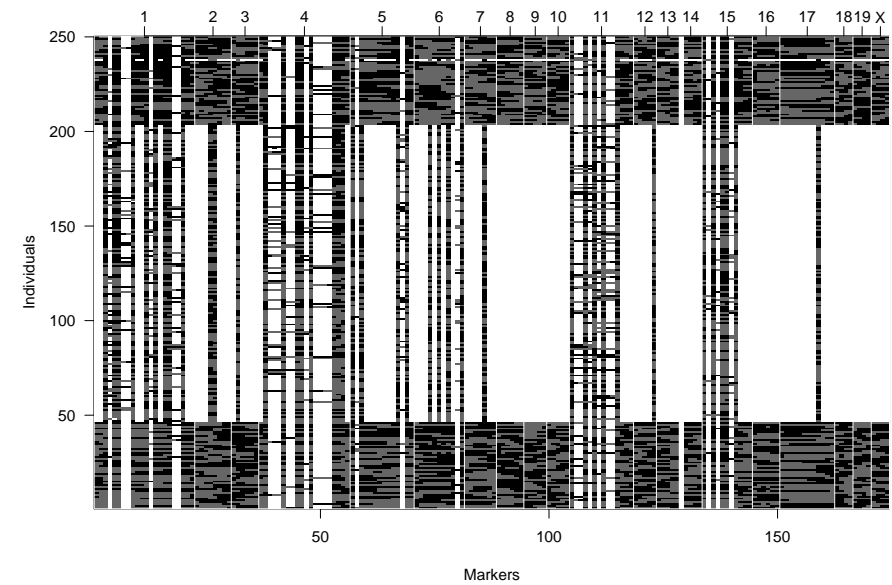
250 male mice from the backcross (A × B) × B

Blood pressure after two weeks drinking water with 1% NaCl



6

## Genotype data



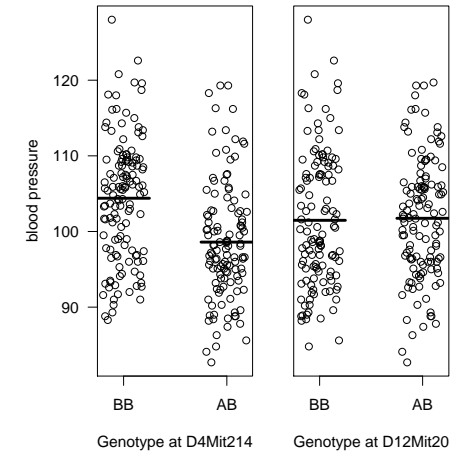
8

# Goals

- Identify quantitative trait loci (QTL) (and interactions among QTL)
- Interval estimates of QTL location
- Estimated QTL effects

# ANOVA at marker loci

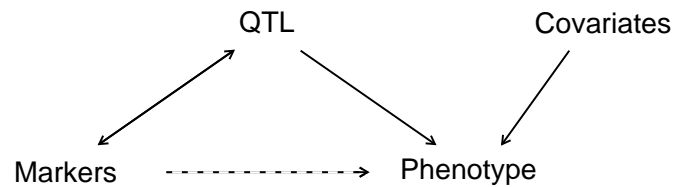
- Split mice into groups according to genotype at a marker.
- Do a t-test / ANOVA.
- Repeat for each marker.



9

11

# Statistical structure



# Interval mapping

## Lander & Botstein (1989)

- Assume a single QTL model.
- Consider each position in the genome, one at a time, as the location of the putative QTL.
- Let  $q = 0/1$  if the (unobserved) QTL genotype is BB/AB. (Or  $0/1/2$  if the QTL genotype is AA/AB/BB in an intercross.)

Assume  $y | q \sim N(\mu_q, \sigma)$

- Calculate  $p_q = \Pr(q | \text{marker data})$ .

$y | \text{marker data} \sim \sum_q p_q \phi(y | \mu_q, \sigma)$

The missing data problem:

Markers  $\longleftrightarrow$  QTL

The model selection problem:

QTL, covariates  $\longrightarrow$  phenotype

10

12

## LOD scores

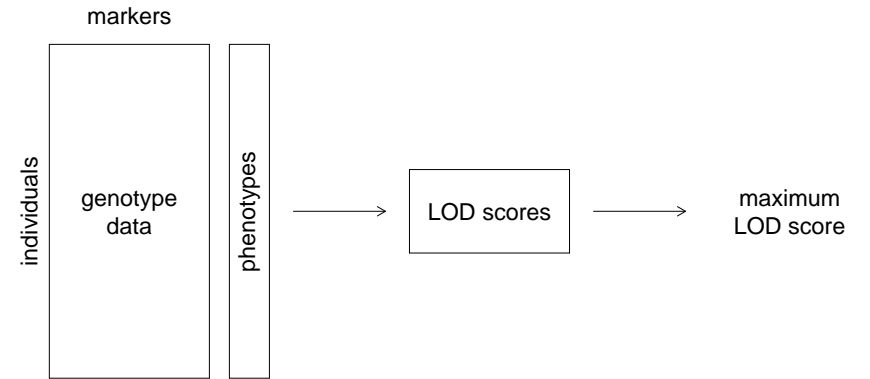
$\text{LOD}(\lambda) = \log_{10}$  likelihood ratio comparing the hypothesis of a QTL at position  $\lambda$  versus that of no QTL

$$= \log_{10} \left\{ \frac{\Pr(y|\text{QTL at } \lambda, \hat{\mu}_{q\lambda}, \hat{\sigma}_\lambda)}{\Pr(y|\text{no QTL}, \hat{\mu}, \hat{\sigma})} \right\}$$

$\hat{\mu}_{q\lambda}, \hat{\sigma}_\lambda$  are the MLEs, assuming a single QTL at position  $\lambda$ .

No QTL model: The phenotypes are iid  $N(\mu, \sigma^2)$ .

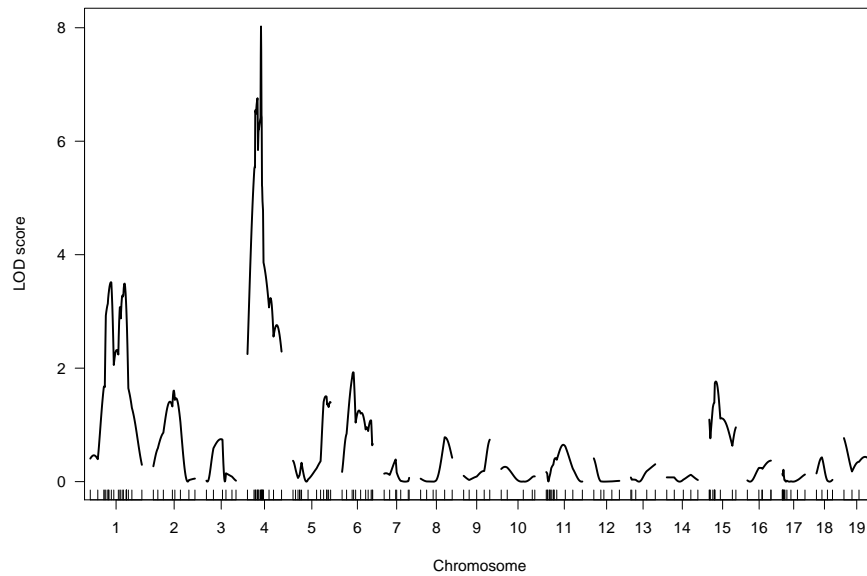
## Permutation test



13

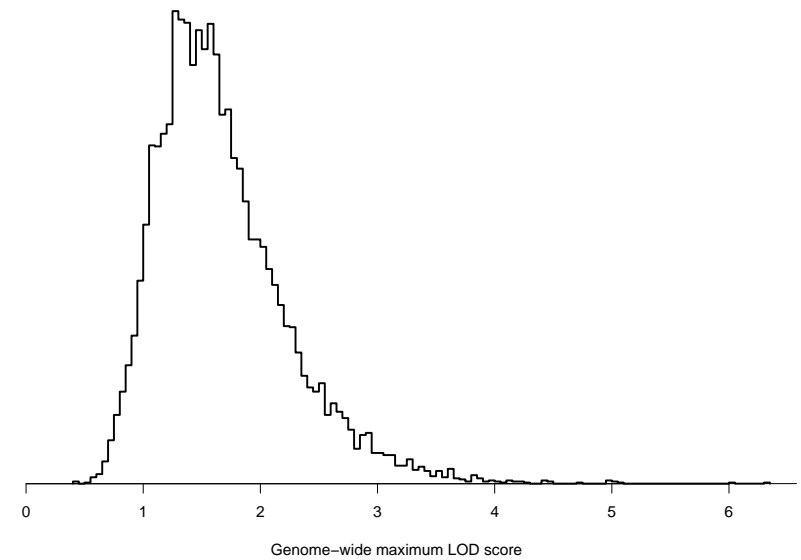
15

## LOD curves



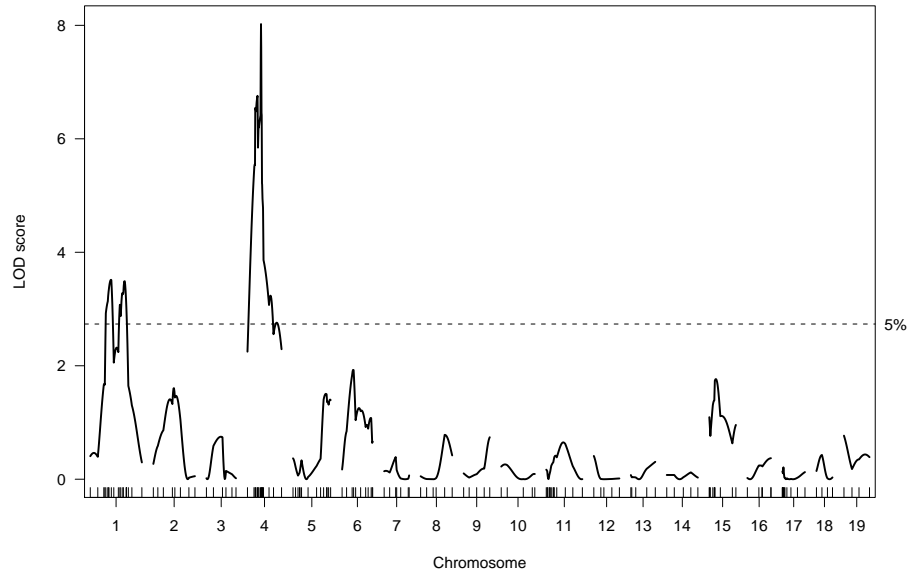
14

## Permutation results



16

# LOD curves



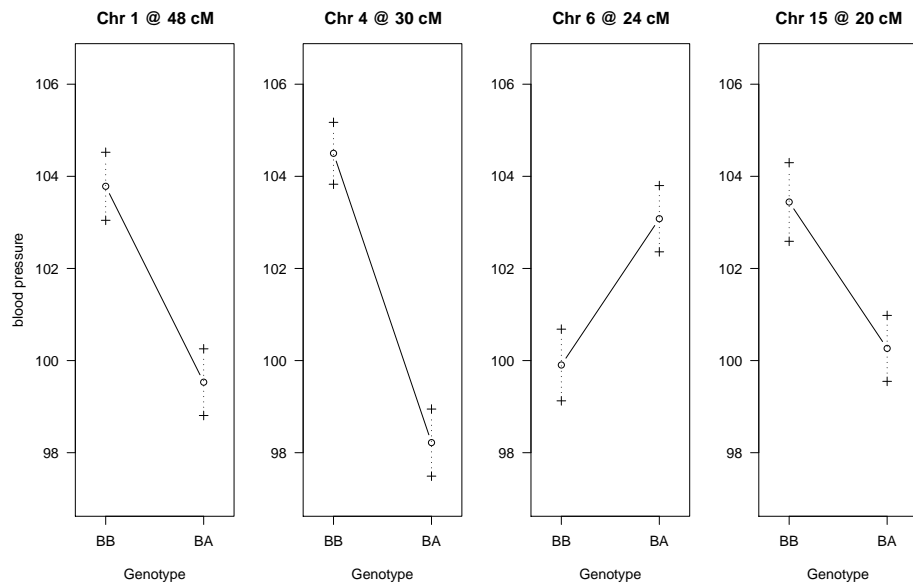
17

# Modeling multiple QTL

- Reduce residual variation → increased power
- Separate linked QTL
- Identify interactions among QTL (epistasis)

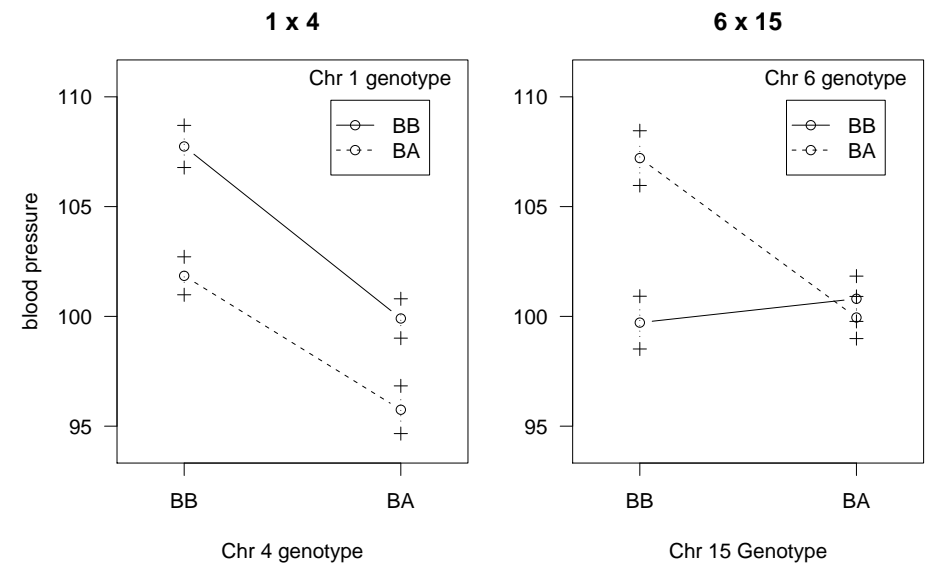
19

# Estimated effects



18

# Estimated effects



20

# Hypothesis testing?

- In the past, QTL mapping has been regarded as a task of hypothesis testing.

Is this a QTL?

Much of the focus has been on adjusting for test multiplicity.

- It is better to view the problem as one of model selection.

What set of QTL are well supported?

Is there evidence for QTL-QTL interactions?

Model = a defined set of QTL and QTL-QTL interactions  
(and possibly covariates and QTL-covariate interactions).

21

# Target

- Selection of a model includes two types of errors:
  - Miss important terms (QTLs or interactions)
  - Include extraneous terms
- Unlike in hypothesis testing, we can make both errors at the same time.
- Identify as many correct terms as possible, while controlling the rate of inclusion of extraneous terms.

23

# Model selection

- Class of models
  - Additive models
  - + pairwise interactions
  - + higher-order interactions
  - Regression trees
- Model fit
  - Maximum likelihood
  - Haley-Knott regression
  - extended Haley-Knott
  - Multiple imputation
  - MCMC
- Model comparison
  - Estimated prediction error
  - AIC, BIC, penalized likelihood
  - Bayes
- Model search
  - Forward selection
  - Backward elimination
  - Stepwise selection
  - Randomized algorithms

22

# What is special here?

- Goal: identify the major players
- A continuum of ordinal-valued covariates (the genetic loci)
- Association among the covariates
  - Loci on different chromosomes are independent
  - Along chromosome, a very simple (and known) correlation structure

24

# Automation

- Assistance to the masses
- Understanding performance
- Many phenotypes

# Experience

- Controls rate of inclusion of extraneous terms
- Forward selection over-selects
- Forward selection followed by backward elimination works as well as MCMC
- Need to define performance criteria
- Need large-scale simulations

Broman & Speed, JRSS B 64:641-656, 2002

25

32

# Additive QTL

Simple situation:

- Dense markers
- Complete genotype data
- No epistasis

$$y = \mu + \sum \beta_j q_j + \epsilon \quad \text{which } \beta_j \neq 0?$$

$$\text{LOD}_\delta(\gamma) = \text{LOD}(\gamma) - T |\gamma|$$

0 vs 1 QTL:  $\text{LOD}_\delta(\emptyset) = 0$

$$\text{LOD}_\delta(\{\lambda\}) = \text{LOD}(\{\lambda\}) - T$$

30

# Epistasis

$$y = \mu + \sum \beta_j q_j + \sum \gamma_{jk} q_j q_k + \epsilon$$

$$\text{LOD}_{\delta\epsilon}(\gamma) = \text{LOD}(\gamma) - T_m |\gamma|_m + T_i |\gamma|_i$$

$T_m =$  as chosen previously

$T_i = ?$

33

# Idea 1

Imagine there are two additive QTL and consider a 2d, 2-QTL scan.

$$T_i = 95\text{th percentile of the distribution of } \max \text{LOD}_f(s, t) - \max \text{LOD}_a(s, t)$$

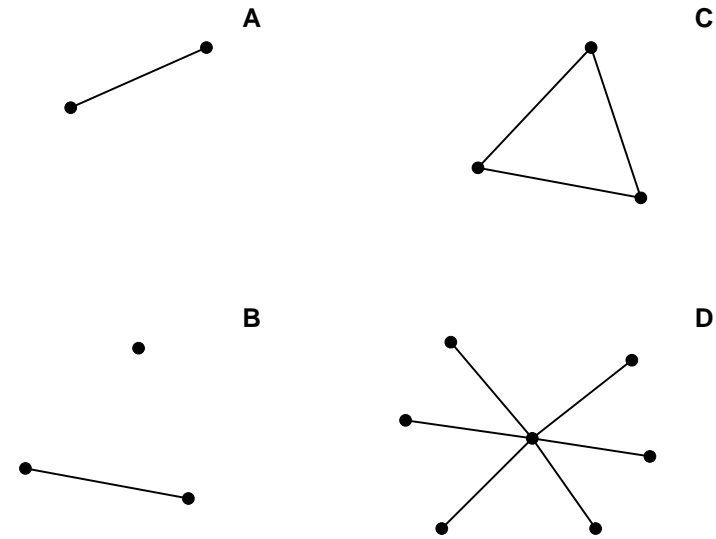
For the mouse genome:

$$T_m = 2.69 \text{ (BC) or } 3.52 \text{ (F}_2\text{)}$$

$$T_i^H = 2.62 \text{ (BC) or } 4.28 \text{ (F}_2\text{)}$$

35

# Models as graphs



38

# Idea 2

Imagine there is one QTL and consider a 2d, 2-QTL scan.

$$T_m + T_i = 95\text{th percentile of the distribution of } \max \text{LOD}_f(s, t) - \max \text{LOD}_1(s)$$

For the mouse genome:

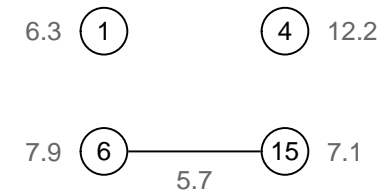
$$T_m = 2.69 \text{ (BC) or } 3.52 \text{ (F}_2\text{)}$$

$$T_i^H = 2.62 \text{ (BC) or } 4.28 \text{ (F}_2\text{)}$$

$$T_i^L = 1.19 \text{ (BC) or } 2.69 \text{ (F}_2\text{)}$$

37

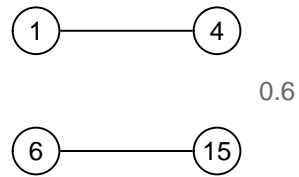
# Results



$$T_m = 2.69 \quad T_i^H = 2.62 \quad T_i^L = 1.19 \quad T_m + T_i^H = 5.31 \quad T_m + T_i^L = 3.88$$

41

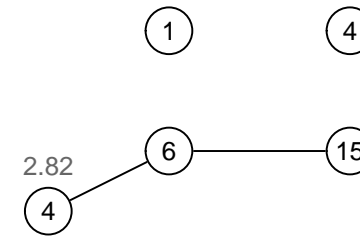
## Add an interaction?



$$T_m = 2.69 \quad T_i^H = 2.62 \quad T_i^L = 1.19 \quad T_m + T_i^H = 5.31 \quad T_m + T_i^L = 3.88$$

42

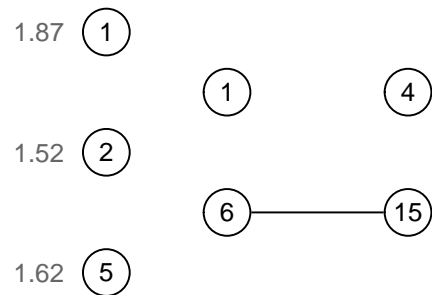
## Add another QTL?



$$T_m = 2.69 \quad T_i^H = 2.62 \quad T_i^L = 1.19 \quad T_m + T_i^H = 5.31 \quad T_m + T_i^L = 3.88$$

49

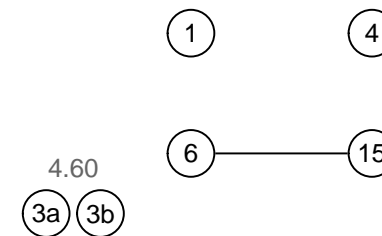
## Add another QTL?



$$T_m = 2.69 \quad T_i^H = 2.62 \quad T_i^L = 1.19 \quad T_m + T_i^H = 5.31 \quad T_m + T_i^L = 3.88$$

47

## Add a pair of QTL?



$$T_m = 2.69 \quad T_i^H = 2.62 \quad T_i^L = 1.19 \quad T_m + T_i^H = 5.31 \quad T_m + T_i^L = 3.88$$

50

## To do

- Improve search procedures
- Study performance  
(especially relative to other approaches)
- Measuring model uncertainty
- Measuring uncertainty in QTL location

51

## Summary

- QTL mapping is a model selection problem
- The criterion for comparing models is most important
- We're focusing on a penalized likelihood method  
and are close to a practiceable solution

52

## Acknowledgments

Ani Manichaikul	Johns Hopkins University
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53