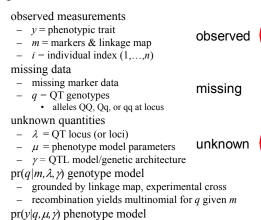
#### **Bayesian Interval Mapping**

1.	Bayesian strategy	3-19
2.	Markov chain sampling	20-27
3.	sampling genetic architectures	28-35
4.	criteria for model selection	36-44

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## QTL model selection: key players



distribution shape (assumed normal here)

- unknown parameters  $\mu$  (could be non-parametric)

m q  $\mu$ after

Sen Churchill (2001)

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#### 1. Bayesian strategy for QTL study

- augment data (y,m) with missing genotypes q
- study unknowns  $(\mu, \lambda, \gamma)$  given augmented data (y, m, q)
  - find better genetic architectures  $\gamma$
  - find most likely genomic regions = QTL =  $\lambda$
  - estimate phenotype parameters = genotype means =  $\mu$
- sample from posterior in some clever way
  - multiple imputation (Sen Churchill 2002)
  - Markov chain Monte Carlo (MCMC)
    - (Satagopan et al. 1996; Yi et al. 2005, 2007)

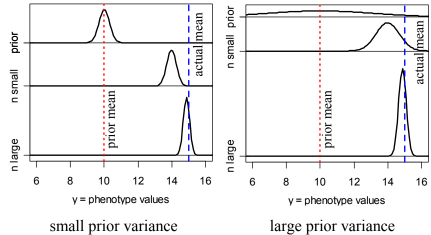
$$posterior = \frac{likelihood*prior}{constant}$$

posterior for 
$$q, \mu, \lambda, \gamma = \frac{\text{phenotype likelihood}*[\text{prior for } q, \mu, \lambda, \gamma]}{\text{constant}}$$

$$\operatorname{pr}(q, \mu, \lambda, \gamma \mid y, m) = \frac{\operatorname{pr}(y \mid q, \mu, \gamma) * [\operatorname{pr}(q \mid m, \lambda, \gamma) \operatorname{pr}(\mu \mid \gamma) \operatorname{pr}(\lambda \mid m, \gamma) \operatorname{pr}(\gamma)]}{\operatorname{pr}(y \mid m)}$$
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## Bayes posterior for normal data



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## Bayes posterior for normal data

model  $y_i = \mu + e_i$ 

environment  $e \sim N(0, \sigma^2), \sigma^2$  known

likelihood  $y \sim N(\mu, \sigma^2)$ 

prior  $\mu \sim N(\mu_0, \kappa \sigma^2)$ ,  $\kappa$  known

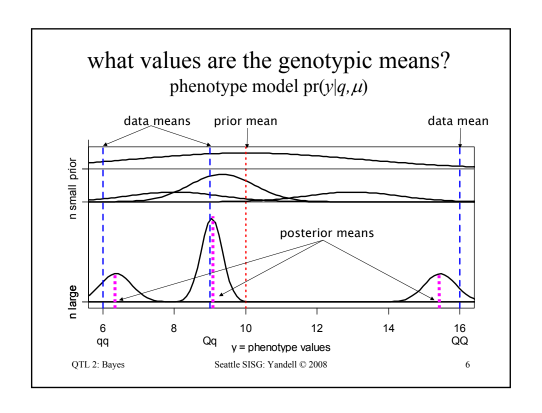
posterior: mean tends to sample mean single individual  $\mu \sim N(\mu_0 + b_1(y_1 - \mu_0), b_1\sigma^2)$ 

sample of *n* individuals  $\mu \sim N(b_n \overline{y}_{\bullet} + (1 - b_n)\mu_0, b_n \sigma^2 / n)$ 

with  $\overline{y}_{\bullet} = \sup_{\{i=1,\dots,n\}} y_i / n$ 

shrinkage factor (shrinks to 1)  $b_n = \frac{\kappa n}{\kappa n + 1} \rightarrow 1$ 

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### Bayes posterior QTL means

posterior centered on sample genotypic mean but shrunken slightly toward overall mean

phenotype mean: 
$$E(y | q) = \mu_q$$
  $V(y | q) = \sigma^2$ 

genotypic prior: 
$$E(\mu_q) = \bar{y}_{\bullet}$$
  $V(\mu_q) = \kappa \sigma^2$ 

posterior: 
$$E(\mu_q \mid y) = b_q \overline{y}_q + (1 - b_q) \overline{y}_{\bullet} \quad V(\mu_q \mid y) = b_q \sigma^2 / n_q$$

$$n_q = \operatorname{count}\{q_i = q\} \qquad \overline{y}_q = \sup_{\{q_i = q\}} y_i / n_q$$

shrinkage: 
$$b_q = \frac{\kappa n_q}{\kappa n_q + 1} \rightarrow 1$$

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# partition genotypic effects on phenotype

- phenotype depends on genotype
- genotypic value partitioned into
  - main effects of single QTL
  - epistasis (interaction) between pairs of QTL

$$\mu_q = \beta_0 + \beta_q = E(Y;q)$$
  
$$\beta_q = \beta(q_2) + \beta(q_2) + \beta(q_1,q_2)$$

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## partitition genotypic variance

- consider same 2 QTL + epistasis
- centering variance  $V(\beta_0) = \kappa_0 \sigma^2 = s^2$
- genotypic variance  $V(\beta_q) = \kappa_1 \sigma^2 = \sigma_q^2 = \sigma_1^2 + \sigma_2^2 + \sigma_{12}^2$
- heritability  $h_q^2=rac{\sigma_q^2}{\sigma_q^2+\sigma^2}=h_1^2+h_2^2+h_{12}^2$

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Q

## posterior mean $\approx$ LS estimate

$$\beta_q \mid y \sim N(b_q \hat{\beta}_q, b_q C_q \sigma^2)$$

$$\approx N(\hat{\beta}_q, C_q \sigma^2)$$

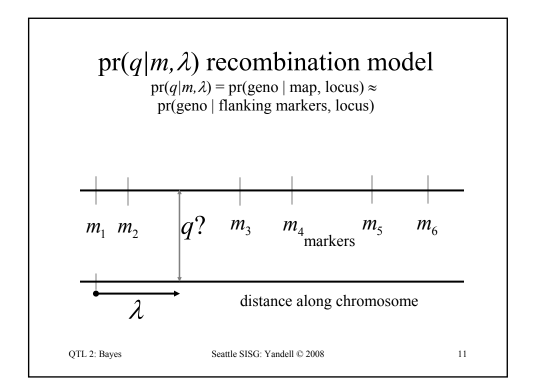
LS estimate  $\hat{\beta}_q = \text{sum}_i[\text{sum}_j \hat{\beta}(q_{ij})] = \text{sum}_i w_{qi} y_i$ 

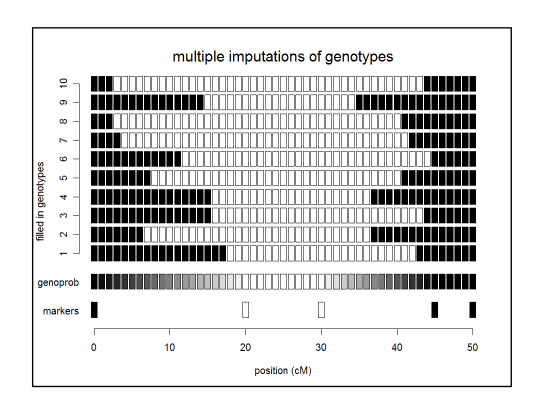
variance 
$$V(\hat{\beta}_q) = \text{sum}_i w_{qi}^2 \sigma^2 = C_q \sigma^2$$

shrinkage 
$$b_q = \kappa_1 / (\kappa_1 + C_q) \rightarrow 1$$

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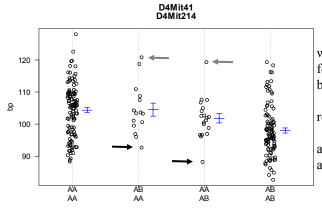
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## what are likely QTL genotypes q?

how does phenotype y improve guess?



Genotype

what are probabilities for genotype *q* between markers?

recombinants AA:AB

all 1:1 if ignore *y* and if we use *y*?

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## posterior on QTL genotypes q

- full conditional of q given data, parameters
  - proportional to prior  $pr(q \mid m, \lambda)$ 
    - weight toward q that agrees with flanking markers
  - proportional to likelihood pr( $y \mid q, \mu$ )
    - $\bullet$  weight toward q with similar phenotype values
  - posterior recombination model balances these two
- this is the E-step of EM computations

$$\operatorname{pr}(q \mid y, m, \mu, \lambda) = \frac{\operatorname{pr}(y \mid q, \mu) * \operatorname{pr}(q \mid m, \lambda)}{\operatorname{pr}(y \mid m, \mu, \lambda)}$$

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#### Where are the loci $\lambda$ on the genome?

- prior over genome for QTL positions
  - flat prior = no prior idea of loci
  - or use prior studies to give more weight to some regions
- posterior depends on QTL genotypes q

$$\operatorname{pr}(\lambda \mid m,q) = \operatorname{pr}(\lambda) \operatorname{pr}(q \mid m,\lambda) / \operatorname{constant}$$

- constant determined by averaging
  - over all possible genotypes q
  - over all possible loci  $\lambda$  on entire map
- no easy way to write down posterior

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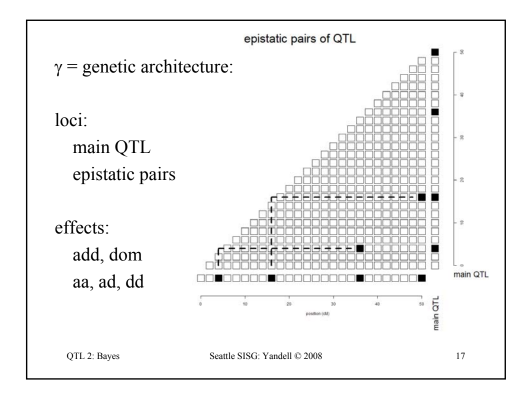
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#### what is the genetic architecture $\gamma$ ?

- which positions correspond to QTLs?
  - priors on loci (previous slide)
- which QTL have main effects?
  - priors for presence/absence of main effects
    - same prior for all QTL
    - can put prior on each d.f. (1 for BC, 2 for F2)
- which pairs of QTL have epistatic interactions?
  - prior for presence/absence of epistatic pairs
    - depends on whether 0,1,2 QTL have main effects
    - epistatic effects less probable than main effects

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## Bayesian priors & posteriors

- augmenting with missing genotypes q
  - prior is recombination model
  - posterior is (formally) E step of EM algorithm
- sampling phenotype model parameters  $\mu$ 
  - prior is "flat" normal at grand mean (no information)
  - posterior shrinks genotypic means toward grand mean
  - (details for unexplained variance omitted here)
- sampling QTL loci  $\lambda$ 
  - prior is flat across genome (all loci equally likely)
- sampling QTL genetic architecture model γ
  - number of QTL
    - · prior is Poisson with mean from previous IM study
  - genetic architecture of main effects and epistatic interactions
    - priors on epistasis depend on presence/absence of main effects

### 2. Markov chain sampling

- construct Markov chain around posterior
  - want posterior as stable distribution of Markov chain
  - in practice, the chain tends toward stable distribution
    - initial values may have low posterior probability
    - · burn-in period to get chain mixing well
- sample QTL model components from full conditionals
  - sample locus  $\lambda$  given  $q, \gamma$  (using Metropolis-Hastings step)
  - sample genotypes q given  $\lambda, \mu, y, \gamma$  (using Gibbs sampler)
  - sample effects  $\mu$  given  $q, y, \gamma$  (using Gibbs sampler)
  - sample QTL model  $\gamma$  given  $\lambda, \mu, y, q$  (using Gibbs or M-H)

$$(\lambda, q, \mu, \gamma) \sim \operatorname{pr}(\lambda, q, \mu, \gamma \mid y, m)$$

$$(\lambda, q, \mu, \gamma)_1 \rightarrow (\lambda, q, \mu, \gamma)_2 \rightarrow \cdots \rightarrow (\lambda, q, \mu, \gamma)_N$$

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## MCMC sampling of unknowns $(q, \mu, \lambda)$ for given genetic architecture $\gamma$

- Gibbs sampler
  - genotypes q
  - effects μ
  - not loci  $\lambda$

$$q \sim \operatorname{pr}(q \mid y_{i}, m_{i}, \mu, \lambda)$$

$$\mu \sim \frac{\operatorname{pr}(y \mid q, \mu)\operatorname{pr}(\mu)}{\operatorname{pr}(y \mid q)}$$

$$\lambda \sim \frac{\operatorname{pr}(q \mid m, \lambda)\operatorname{pr}(\lambda \mid m)}{\operatorname{pr}(q \mid m)}$$



- Metropolis-Hastings sampler
  - extension of Gibbs sampler
  - does not require normalization
    - $\operatorname{pr}(q \mid m) = \operatorname{sum}_{\lambda} \operatorname{pr}(q \mid m, \lambda) \operatorname{pr}(\lambda)$

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## Gibbs sampler for two genotypic means

- want to study two correlated effects
  - could sample directly from their bivariate distribution
  - assume correlation  $\rho$  is known
- instead use Gibbs sampler:
  - sample each effect from its full conditional given the other
  - pick order of sampling at random
  - repeat many times

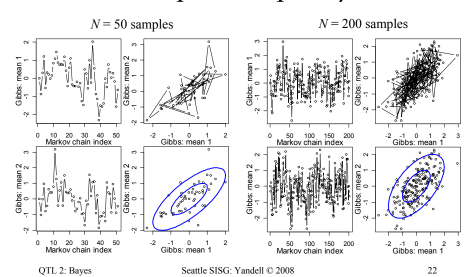
$$\begin{pmatrix} \mu_1 \\ \mu_2 \end{pmatrix} \sim N \begin{pmatrix} 0 \\ 0 \end{pmatrix}, \begin{pmatrix} 1 & \rho \\ \rho & 1 \end{pmatrix} \end{pmatrix}$$
$$\mu_1 \sim N \left( \rho \mu_2, 1 - \rho^2 \right)$$
$$\mu_2 \sim N \left( \rho \mu_1, 1 - \rho^2 \right)$$

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## Gibbs sampler samples: $\rho = 0.6$



#### full conditional for locus

- cannot easily sample from locus full conditional  $pr(\lambda | y, m, \mu, q) = pr(\lambda | m, q)$ =  $pr(q | m, \lambda) pr(\lambda) / constant$
- constant is very difficult to compute explicitly
  - must average over all possible loci  $\lambda$  over genome
  - must do this for every possible genotype q
- Gibbs sampler will not work in general
  - but can use method based on ratios of probabilities
  - Metropolis-Hastings is extension of Gibbs sampler

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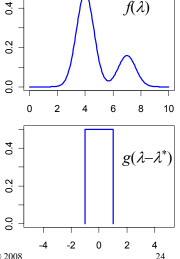
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## Metropolis-Hastings idea

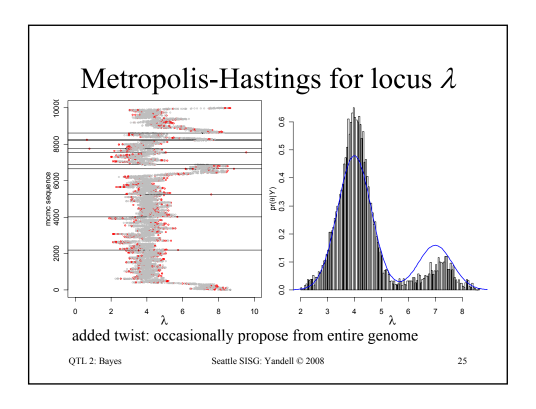
- want to study distribution  $f(\lambda)$ 
  - take Monte Carlo samples
    - unless too complicated
  - take samples using ratios of f
- Metropolis-Hastings samples:
  - propose new value  $\lambda^*$ 
    - near (?) current value  $\lambda$
    - from some distribution g
  - accept new value with prob a
    - Gibbs sampler: a = 1 always

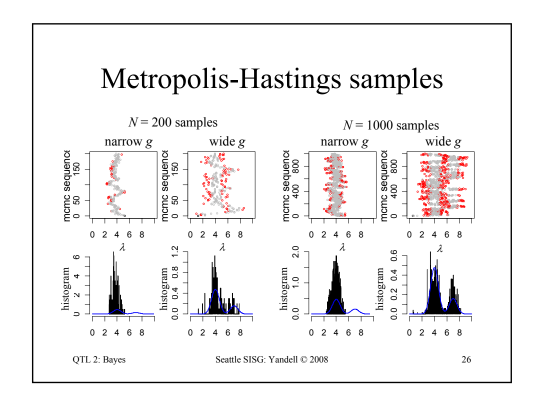
$$a = \min\left(1, \frac{f(\lambda^*)g(\lambda^* - \lambda)}{f(\lambda)g(\lambda - \lambda^*)}\right)$$



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#### 3. sampling genetic architectures

- search across genetic architectures A of various sizes
  - allow change in number of QTL
  - allow change in types of epistatic interactions
- methods for search
  - reversible jump MCMC
  - Gibbs sampler with loci indicators
- complexity of epistasis
  - Fisher-Cockerham effects model
  - general multi-QTL interaction & limits of inference

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#### reversible jump MCMC

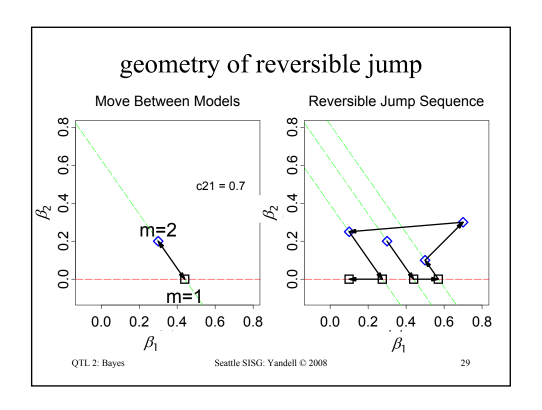
- consider known genotypes q at 2 known loci  $\lambda$ 
  - models with 1 or 2 QTL
- M-H step between 1-QTL and 2-QTL models
  - model changes dimension (via careful bookkeeping)
  - consider mixture over QTL models H

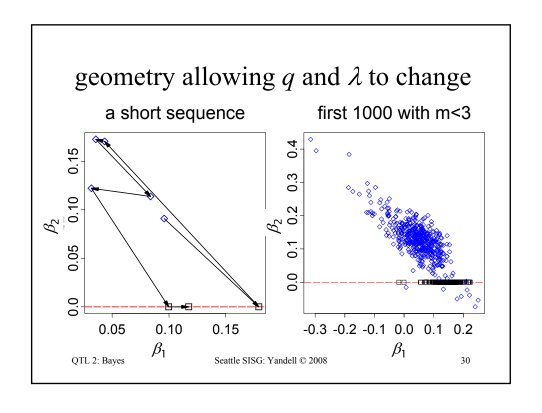
$$\gamma = 1 \text{ QTL} : Y = \beta_0 + \beta(q_1) + e$$

$$\gamma = 2 \text{ QTL} : Y = \beta_0 + \beta(q_1) + \beta(q_2) + e$$

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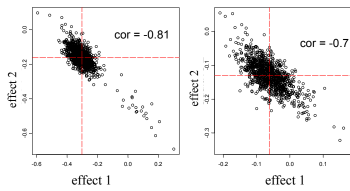




## collinear QTL = correlated effects



8-week



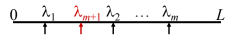
- linked QTL = collinear genotypes
  - > correlated estimates of effects (negative if in coupling phase)
  - > sum of linked effects usually fairly constant

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## sampling across QTL models $\gamma$



action steps: draw one of three choices

- update QTL model  $\gamma$  with probability 1- $b(\gamma)$ - $d(\gamma)$ 
  - update current model using full conditionals
  - sample QTL loci, effects, and genotypes
- add a locus with probability  $b(\gamma)$ 
  - propose a new locus along genome
  - innovate new genotypes at locus and phenotype effect
  - decide whether to accept the "birth" of new locus
- drop a locus with probability  $d(\gamma)$ 
  - propose dropping one of existing loci
  - decide whether to accept the "death" of locus

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### Gibbs sampler with loci indicators

- consider only QTL at pseudomarkers
  - every 1-2 cM
  - modest approximation with little bias
- use loci indicators in each pseudomarker
  - $\gamma = 1$  if QTL present
  - $\gamma = 0$  if no QTL present
- Gibbs sampler on loci indicators  $\gamma$ 
  - relatively easy to incorporate epistasis
  - Yi, Yandell, Churchill, Allison, Eisen, Pomp (2005 Genetics)
    - (see earlier work of Nengjun Yi and Ina Hoeschele)

$$\mu_q = \mu + \gamma_1 \beta(q_1) + \gamma_2 \beta(q_2), \ \gamma_k = 0.1$$

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## Bayesian shrinkage estimation

- soft loci indicators
  - strength of evidence for  $\lambda_i$  depends on  $\gamma$
  - 0 ≤  $\gamma$  ≤ 1 (grey scale)
  - shrink most ½s to zero
- Wang et al. (2005 Genetics)
  - Shizhong Xu group at U CA Riverside

$$\mu_{q} = \beta_{0} + \gamma_{1}\beta_{1}(q_{1}) + \gamma_{2}\beta_{2}(q_{1}), \ 0 \le \gamma_{k} \le 1$$

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## 4. criteria for model selection balance fit against complexity

- classical information criteria
  - penalize likelihood L by model size  $|\gamma|$
  - $-IC = -2 \log L(\gamma | y) + \text{penalty}(\gamma)$
  - maximize over unknowns
- Bayes factors
  - marginal posteriors  $pr(y \mid y)$
  - average over unknowns

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#### classical information criteria

- start with likelihood  $L(\gamma | y, m)$ 
  - measures fit of architecture ( $\gamma$ ) to phenotype (y)
    - given marker data (m)
  - genetic architecture  $(\gamma)$  depends on parameters
    - have to estimate loci ( $\mu$ ) and effects ( $\lambda$ )
- complexity related to number of parameters
  - $|\gamma| = \text{size of genetic architecture}$ 
    - BC:  $|\gamma| = 1 + n.qtl + n.qtl(n.qtl 1) = 1 + 4 + 12 = 17$
    - F2:  $|\gamma| = 1 + 2n.qtl + 4n.qtl(n.qtl 1) = 1 + 8 + 48 = 57$

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#### classical information criteria

- construct information criteria
  - balance fit to complexity
  - Akaike AIC =  $-2 \log(L) + 2 |\gamma|$
  - Bayes/Schwartz BIC =  $-2 \log(L) + |\gamma| \log(n)$
  - Broman BIC<sub> $\delta$ </sub> = -2 log(L) +  $\delta |\gamma| \log(n)$
  - general form: IC =  $-2 \log(L) + |\gamma| D(n)$
- compare models
  - hypothesis testing: designed for one comparison
    - $2 \log[LR(\gamma_1, \gamma_2)] = L(y|m, \gamma_2) L(y|m, \gamma_1)$
  - model selection: penalize complexity
    - $IC(\gamma_1, \gamma_2) = 2 log[LR(\gamma_1, \gamma_2)] + (|\gamma_2| |\gamma_1|) D(n)$

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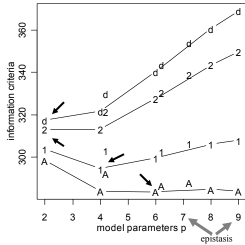
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#### information criteria vs. model size

- WinQTL 2.0
- SCD data on F2
- A=AIC
- 1=BIC(1)
- 2=BIC(2)
- d=BIC(δ)
- models
  - 1,2,3,4 QTL
    - 2+5+9+2
  - epistasis

• 2:2 AD



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#### Bayes factors

- ratio of model likelihoods
  - ratio of posterior to prior odds for architectures
  - averaged over unknowns

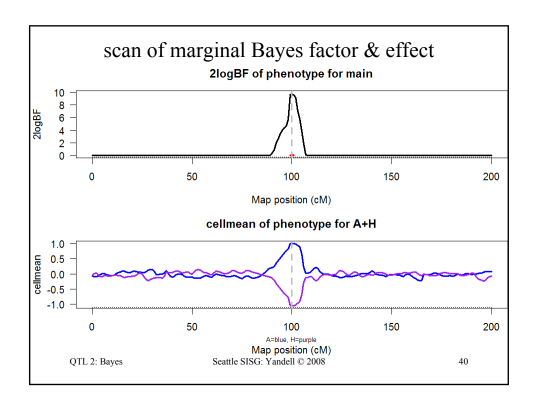
$$B_{12} = \frac{\text{pr}(\gamma_1 \mid y, m) / \text{pr}(\gamma_2 \mid y, m)}{\text{pr}(\gamma_1) / \text{pr}(\gamma_2)} = \frac{\text{pr}(y \mid m, \gamma_1)}{\text{pr}(y \mid m, \gamma_2)}$$

- roughly equivalent to BIC
  - BIC maximizes over unknowns
  - BF averages over unknowns

$$-2\log(B_{12}) = -2\log(LR) - (|\gamma_2| - |\gamma_1|)\log(n)$$

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#### issues in computing Bayes factors

- BF insensitive to shape of prior on  $\gamma$ 
  - geometric, Poisson, uniform
  - precision improves when prior mimics posterior
- BF sensitivity to prior variance on effects  $\theta$ 
  - prior variance should reflect data variability
  - resolved by using hyper-priors
    - automatic algorithm; no need for user tuning
- easy to compute Bayes factors from samples
  - sample posterior using MCMC
  - posterior pr(y | y, m) is marginal histogram

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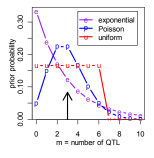
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#### Bayes factors & genetic architecture $\gamma$

- $|\gamma|$  = number of QTL
  - prior  $pr(\gamma)$  chosen by user
  - posterior  $pr(\gamma | y, m)$ 
    - sampled marginal histogram
    - shape affected by prior pr(A)

$$BF_{\gamma_1,\gamma_2} = \frac{\operatorname{pr}(\gamma_1|y,m)/\operatorname{pr}(\gamma_1)}{\operatorname{pr}(\gamma_2|y,m)/\operatorname{pr}(\gamma_2)}$$

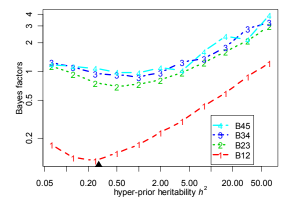


- pattern of QTL across genome
- gene action and epistasis

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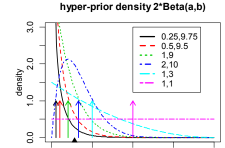
$$\beta_{qj} \sim N(0, \sigma_G^2/m), \sigma_G^2 = h^2 \sigma_{\text{total}}^2, h^2 \text{ fixed}$$

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#### BF insensitivity to random effects prior



1.0

hyper-parameter heritability  $h^2$ 

#### 

insensitivity to hyper-prior

$$\beta_{qj} \sim N(0, \sigma_G^2 / m), \sigma_G^2 = h^2 \sigma_{\text{total}}^2, \frac{1}{2} h^2 \sim \text{Beta}(a, b)$$

2.0

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0.0

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