

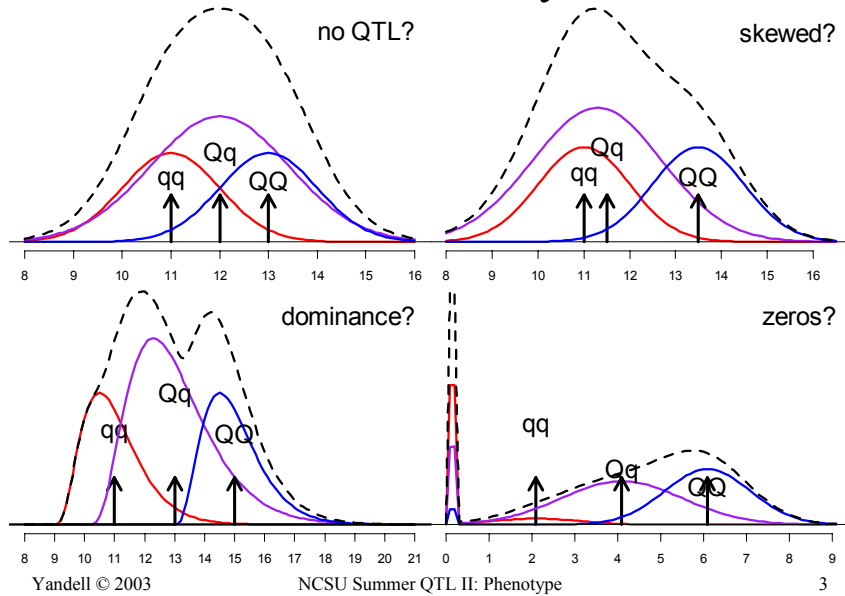
Extending the Phenotype Model

- limitations of parametric models
- diagnostic tools for QTL analysis
- QTL mapping with other parametric "families"
- quick fixes via data transformations
- semi-parametric approaches
- non-parametric approaches
- bottom line:
 - normal phenotype model works well to pick up loci, but may be poor at estimating effects if data not normal

limitations of parametric models

- measurements not normal
 - categorical traits: counts (*e.g.* number of tumors)
 - use methods specific for counts
 - binomial, Poisson, negative binomial
 - traits measured over time and/or space
 - survival time (*e.g.* days to flowering)
 - developmental process; signal transduction between cells
 - TP Speed (pers. comm.); Ma, Casella, Wu (2002)
- false positives due to miss-specified model
 - how to check model assumptions?
- want more robust estimates of effects
 - parametric: only center (mean), spread (SD)
 - shape of distribution may be important

what if data are far away from ideal?



diagnostic tools for QTL (Hackett 1997)

- illustrated with BC, adapt regression diagnostics
- normality & equal variance (fig. 1)
 - plot fitted values vs. residuals--football shaped?
 - normal scores plot of residuals--straight line?
- number of QTL: likelihood profile (fig. 2)
 - flat shoulders near LOD peak: evidence for 1 vs. 2 QTL
- genetic effects
 - effect estimate near QTL should be $(1-2r)a$
 - plot effect vs. location

marker density & sample size: 2 QTL

modest sample size
dense vs. sparse markers

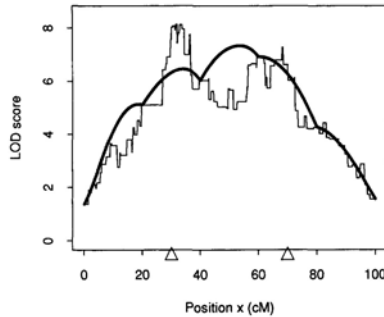


FIGURE 1.—The two-QTL true model with a QTL at 30 cM and a second QTL of somewhat smaller effect at 70 cM (true locations indicated by Δ). A normal single-QTL model is assumed and the LOD score for 100 simulated individuals is given for dense markers (thin curve) and markers at 20-cM intervals (bold curve).

Wright Kong (1997 *Genetics*)

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NCSU Summer QTL II: Phenotype

large sample size
dense vs. sparse markers

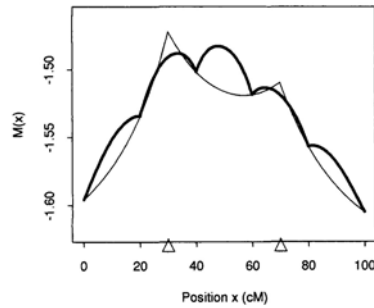


FIGURE 4.— $M(x)$ for a normal single-QTL assumed model under a two-QTL true model when both of the genes lie on the chromosome under study. This scenario was originally depicted in Figure 1. With dense markers (thin curve), $M(x)$ peaks at exactly 30 cM, the location of the QTL of stronger effect. With nondense markers at 20-cM intervals, $M(x)$ peaks at 47 cM in an incorrect interval (bold curve). Note the similarity in shape between the LODs in Figure 1 and the limiting forms depicted here.

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robust locus estimate for non-normal phenotype

large sample size &
dense marker map:
no need for normality

but what happens for
modest sample sizes?

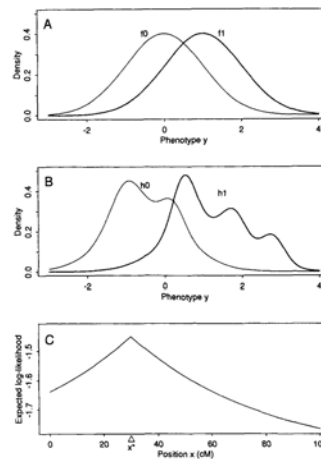


FIGURE 2.—Misspecification of the phenotype model. (A) The assumed distributions f_0 and f_1 . (B) The true distributions h_0 , h_1 . (C) The expected log-likelihood across the chromosome when the markers are dense. Despite the misspecification, the function is maximized at exactly the true location $x^* = 30$ cM (indicated by Δ).

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What shape is your histogram?

- histogram conditional on known QT genotype
 - $\text{pr}(Y|qq, \theta)$ model shape with genotype qq
 - $\text{pr}(Y|Qq, \theta)$ model shape with genotype Qq
 - $\text{pr}(Y|QQ, \theta)$ model shape with genotype QQ
- is the QTL at a given locus λ ?
 - no QTL $\text{pr}(Y|qq, \theta) = \text{pr}(Y|Qq, \theta) = \text{pr}(Y|QQ, \theta)$
 - QTL present mixture if genotype unknown
- mixture across possible genotypes
 - sum over $Q = qq, Qq, QQ$
 - $\text{pr}(Y|X, \lambda, \theta) = \sum_Q \text{pr}(Q|X, \lambda) \text{pr}(Y|Q, \theta)$

interval mapping likelihood

- likelihood: basis for scanning the genome
 - product over $i = 1, \dots, n$ individuals
 - $$L(\theta, \lambda | Y) = \text{product}_i \text{pr}(Y_i | X_i, \lambda)$$

$$= \text{product}_i \sum_Q \text{pr}(Q | X_i, \lambda) \text{pr}(Y_i | Q, \theta)$$
- problem: unknown phenotype model
 - parametric $\text{pr}(Y|Q, \theta) = f(Y | \mu, G_Q, \sigma^2)$
 - semi-parametric $\text{pr}(Y|Q, \theta) = f(Y) \exp(Y\beta_Q)$
 - non-parametric $\text{pr}(Y|Q, \theta) = F_Q(Y)$

useful models & transformations

- binary trait (yes/no, hi/lo, ...)
 - map directly as another marker
 - categorical: break into binary traits?
 - mixed binary/continuous: condition on $Y > 0$?
- known model for biological mechanism
 - counts Poisson
 - fractions binomial
 - clustered negative binomial
- transform to stabilize variance
 - counts $\sqrt{Y} = \text{sqrt}(Y)$
 - concentration $\log(Y)$ or $\log(Y+c)$
 - fractions $\arcsin(\sqrt{Y})$
- transform to symmetry (approx. normal)
 - fraction $\log(Y/(1-Y))$ or $\log((Y+c)/(1+c-Y))$
- empirical transform based on histogram
 - watch out: hard to do well even without mixture
 - probably better to map untransformed, then examine residuals

semi-parametric QTL

- phenotype model $\text{pr}(Y|Q, \theta) = f(Y)\exp(Y\beta_Q)$
 - unknown parameters $\theta = (f, \beta)$
 - $f(Y)$ is a (unknown) density if there is no QTL
 - $\beta = (\beta_{qq}, \beta_{Qq}, \beta_{QQ})$
 - $\exp(Y\beta_Q)$ 'tilts' f based on genotype Q and phenotype Y
- test for QTL at locus λ
 - $\beta_Q = 0$ for all Q , or $\text{pr}(Y|Q, \theta) = f(Y)$
- includes many standard phenotype models
 - normal $\text{pr}(Y|Q, \theta) = N(G_Q, \sigma^2)$
 - Poisson $\text{pr}(Y|Q, \theta) = \text{Poisson}(G_Q)$
 - exponential, binomial, ..., but not negative binomial

QTL for binomial data

- approximate methods: marker regression
 - Zeng (1993,1994); Visscher et al. (1996); McIntyre et al. (2001)
- interval mapping, CIM
 - Jansen (1992); Xu Atchley (1996); Yi Xu (2000)
 - $Y \sim \text{binomial}(1, \pi)$, π depends on genotype Q
 - $\text{pr}(Y|Q) = (\pi_Q)^Y (1 - \pi_Q)^{(1-Y)}$
 - substitute this phenotype model in EM iteration
- or just map it as another marker!
 - but may have complex

EM algorithm for binomial QTL

- E-step: posterior probability of genotype Q

$$\text{pr}(Q | Y_i, X_i, \lambda, \pi_Q) = \frac{\text{pr}(Q | X_i, \lambda) (\pi_Q)^{Y_i} (1 - \pi_Q)^{(1-Y_i)}}{\text{sum}_Q \text{ of numerator}}$$

- M-step: MLE of binomial probability π_Q

$$\pi_Q = \frac{\text{sum}_i Y_i \text{pr}(Q | Y_i, X_i, \lambda, \pi_Q)}{\text{sum}_i \text{pr}(Q | Y_i, X_i, \lambda, \pi_Q)}$$

threshold or latent variable idea

- "real", unobserved phenotype Z is continuous
- observed phenotype Y is ordinal value
 - no/yes; poor/fair/good/excellent
 - $\text{pr}(Y = j) = \text{pr}(\tau_{j-1} < Z \leq \tau_j)$
 - $\text{pr}(Y \leq j) = \text{pr}(Z \leq \tau_j)$
- use logistic regression idea (Hackett Weller 1995)
 - substitute new phenotype model in to EM algorithm
 - or use Bayesian posterior approach
 - extended to multiple QTL (papers in press)

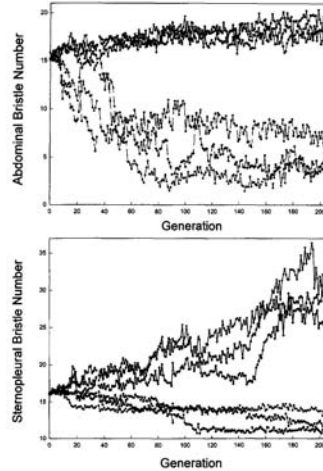
$$\text{pr}(Y \leq j | Q) = \text{pr}(Z \leq \tau_j | Q) = [1 + \exp(\mu + G_Q - \tau_j)]^{-1}$$

quantitative & qualitative traits

- Broman (2003): spike in phenotype
 - large fraction of phenotype has one value
 - map binary trait (is/is not that value)
 - map continuous trait given not that value
- multiple traits
 - Williams et al. (1999)
 - multiple binary & normal traits
 - variance component analysis
 - Corander Sillanpaa (2002)
 - multiple discrete & continuous traits
 - latent (unobserved) variables

other parametric approaches

- Poisson counts
 - Mackay Fry (1996)
 - trait = bristle number
 - Shepel et al (1998)
 - trait = tumor count
- negative binomial
 - Lan *et al.* (2001)
 - number of tumors



Mackay Fry (1996 *Genetics*)

semi-parametric empirical likelihood

- phenotype model $\text{pr}(Y|Q, \theta) = f(Y) \exp(Y\beta_Q)$
 - “point mass” at each measured phenotype Y_i
 - subject to distribution constraints for each Q :

$$1 = \sum_i f(Y_i) \exp(Y_i \beta_Q)$$
- non-parametric empirical likelihood (Owen 1988)

$$L(\theta, \lambda | Y, X) = \text{product}_i [\sum_Q \text{pr}(Q|X_i, \lambda) f(Y_i) \exp(Y_i \beta_Q)]$$

$$= \text{product}_i f(Y_i) [\sum_Q \text{pr}(Q|X_i, \lambda) \exp(Y_i \beta_Q)]$$

$$= \text{product}_i f(Y_i) w_i$$
 - weights $w_i = w(Y_i|X_i, \beta, \lambda)$ rely only on flanking markers
 - 4 possible values for BC, 9 for F2, etc.
- profile likelihood: $L(\lambda | Y, X) = \max_{\theta} L(\theta, \lambda | Y, X)$

semi-parametric formal tests

- clever trick: use partial empirical LOD
 - Zou, Fine, Yandell (2002 *Biometrika*)
 - Lange, Whittaker (2001 *Genetics*) GEE
- has same formal behavior as parametric LOD
 - single locus test: approximately χ^2 with 1 d.f.
 - genome-wide scan: can use same critical values
 - permutation test: possible with some work
- can estimate cumulative distributions
 - nice properties (converge to Gaussian processes)

log empirical likelihood details

$$\log(L(\theta, \lambda | Y, X)) = \sum_i \log(f(Y_i)) + \log(w_i)$$

now profile with respect to β, λ

$$\log(L(\beta, \lambda | Y, X)) = \sum_i \log(f_i) + \log(w_i) \\ + \sum_Q \alpha_Q (1 - \sum_i f_i \exp(Y_i \beta_Q))$$

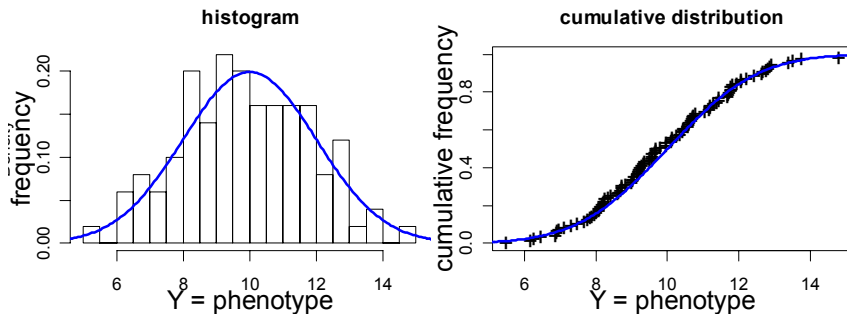
partial likelihood: set Lagrange multipliers α_Q to 0

point mass density estimates

$$f_i = \left[\sum_Q \exp(Y_i \beta_Q) p(Q | X, \lambda) \right]^{-1}$$

$$\text{with } p(Q | X, \lambda) = \sum_i \text{pr}(Q | X_i, \lambda)$$

histograms and CDFs

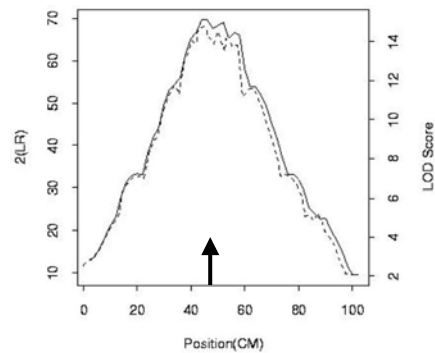


histograms capture shape
but are not very accurate

CDFs are more accurate
but not always intuitive

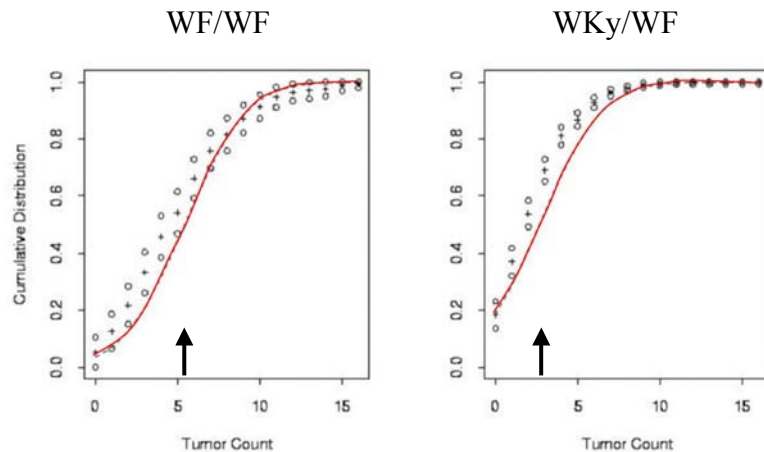
rat study of breast cancer Lan *et al.* (2001 *Genetics*)

- rat backcross
 - two inbred strains
 - Wistar-Furth susceptible
 - Wistar-Kyoto resistant
 - backcross to WF
 - 383 females
 - chromosome 5, 58 markers
- search for resistance genes
- $Y = \#$ mammary carcinomas
- where is the QTL?



dash = normal
solid = semi-parametric

what shape histograms by genotype?



line = normal, + = semi-parametric, o = confidence interval

non-parametric methods

- phenotype model $\text{pr}(Y|Q, \theta) = F_Q(Y)$
 - $\theta = F = (F_{qq}, F_{Qq}, F_{QQ})$ arbitrary distribution functions
- interval mapping Wilcoxon rank-sum test
 - replaced Y by $\text{rank}(Y)$
 - (Kruglyak Lander 1995; Poole Drinkwater 1996; Broman 2003)
 - claimed no estimator of QTL effects
- non-parametric shift estimator
 - semi-parametric shift (Hodges-Lehmann)
 - Zou (2001) thesis, Zou, Yandell, Fine (2002 in review)
 - non-parametric cumulative distribution
 - Fine, Zou, Yandell (2001 in review)
- stochastic ordering (Hoff et al. 2002)

rank-sum QTL methods

- phenotype model $\text{pr}(Y|Q, \theta) = F_Q(Y)$
- replace Y by $\text{rank}(Y)$ and perform IM
 - extension of Wilcoxon rank-sum test
 - fully non-parametric (Kruglyak Lander 1995; Poole Drinkwater 1996)
- Hodges-Lehmann estimator of shift β
 - most efficient if $\text{pr}(Y|Q, \theta) = F(Y+Q\beta)$
 - find β that matches medians
 - problem: genotypes Q unknown
 - resolution: Haley-Knott (1992) regression scan
 - works well in practice, but theory is elusive
 - Zou, Yandell Fine (*Genetics*, in review)

non-parametric QTL CDFs

- estimate non-parametric phenotype model
 - cumulative distributions $F_Q(y) = \text{pr}(Y \leq y | Q)$
 - can use to check parametric model validity
- basic idea:
$$\text{pr}(Y \leq y | X, \lambda) = \text{sum}_Q \text{pr}(Q|X, \lambda) F_Q(y)$$
 - depends on X only through flanking markers
 - few possible flanking marker genotypes
 - 4 for BC, 9 for F2, etc.

finding non-parametric QTL CDFs

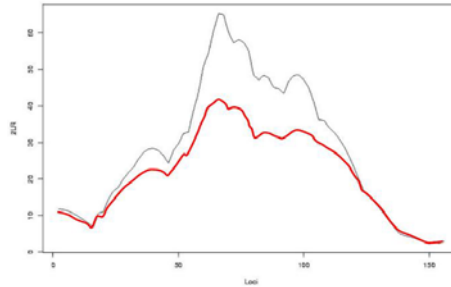
- cumulative distribution $F_Q(y) = \text{pr}(Y \leq y | Q)$
- $F = \{F_Q, \text{all possible QT genotypes } Q\}$
 - BC with 1 QTL: $F = \{F_{QQ}, F_{Qq}\}$
- find F to minimize over all phenotypes y
 $\text{sum}_i [I(Y_i \leq y) - \text{sum}_Q \text{pr}(Q|X, \lambda) F_Q(y)]^2$
- looks complicated, but simple to implement

non-parametric CDF properties

- readily extended to censored data
 - time to flowering for non-vernalized plants
- nice large sample properties
 - estimates of $F(y) = \{F_Q(y)\}$ jointly normal
 - point-wise, experiment-wise confidence bands
- more robust to heavy tails and outliers
- can use to assess parametric assumptions

what QTL influence flowering time? no vernalization: censored survival

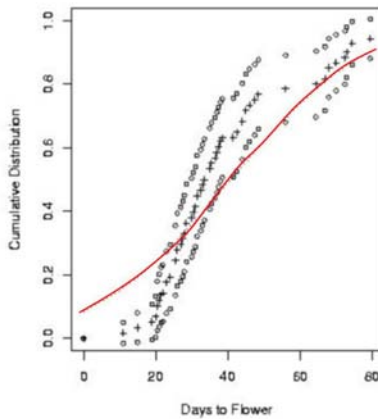
- *Brassica napus*
 - Major female
 - needs vernalization
 - Stellar male
 - insensitive
 - 99 double haploids
- $Y = \log(\text{days to flower})$
 - over 50% Major at QTL never flowered
 - log not fully effective



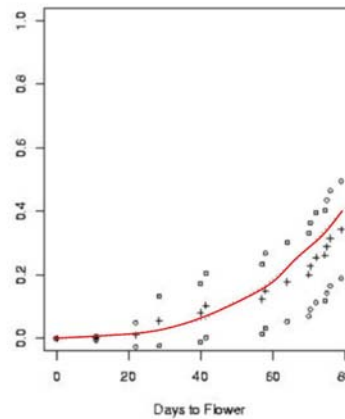
grey = normal, red = non-parametric

what shape is flowering distribution?

B. napus Stellar



B. napus Major



line = normal, + = non-parametric, o = confidence interval