Precision Mapping of Quantitative Trait Loci

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ABSTRACT

Adequate separation of effects of possible multiple linked quantitative trait loci (QTLs) on mapping QTLs is the key to increasing the precision of QTL mapping. A new method of QTL mapping is proposed and analyzed in this paper by combining interval mapping with multiple regression. The basis of the proposed method is an interval test in which the test statistic on a marker interval is made to be unaffected by QTLs located outside a defined interval. This is achieved by fitting other genetic markers in the statistical model as a control when performing interval mapping. Compared with the current QTL mapping method (i.e., the interval mapping method which uses a pair or two pairs of markers for mapping QTLs), this method has several advantages. (1) By confining the test to one region at a time, it reduces a multiple dimensional search problem (for multiple QTLs) to a one dimensional search problem. (2) By conditioning linked markers in the test, the sensitivity of the test statistic to the position of individual QTLs is increased, and the precision of QTL mapping can be improved. (3) By selectively and simultaneously using other markers in the analysis, the efficiency of QTL mapping can be also improved. The behavior of the test statistic under the null hypothesis and appropriate critical value of the test statistic for an overall test in a genome are discussed and analyzed. A simulation study of QTL mapping is also presented which illustrates the utility, properties, advantages and disadvantages of the method.

MANY traits in plants and animals are quantitative in nature, influenced by many genes. It has been, for a long time, an important aim in genetics and breeding to identify those genes contributing significantly to the variation of traits within and between populations or species. With rapid advancement of molecular technology, it is now possible to use molecular marker information to map major quantitative trait loci (QTLs) on chromosomes (e.g., PATERSON et al. 1988, 1991; HILBERT et al. 1991; JACOB et al. 1991; STUBER et al. 1992).

There have been several statistical methods developed to analyze mapping data to search systematically for major QTLs in experimental organisms (e.g., Soller et al. 1976; Weller 1986; Lander and Botstein 1989), among which the interval mapping of Lander and Botstein (1989) has now become the current standard method used by many geneticists for mapping QTLs. Compared with the traditional method (Soller et al. 1976), the interval mapping method has a number of advantages. But it still has several problems, particularly in distinguishing multiple linked QTL effects. When there are two or more QTLs located on a chromosome, the mapping of QTLs can be seriously biased, and QTLs can be mapped to wrong positions (Knott and Haley 1992; Martinez and Curnow 1992).

To increase the reliability and accuracy of QTL mapping, the effects of possible multiple linked QTLs on a chromosome should be adequately separated in testing and estimation. In this paper, a mapping procedure is

elaborated with the aim to improve both the precision and efficiency of mapping multiple QTLs, using multiple markers. The basis of the method is an interval test in which the test statistic is constructed to be unaffected by QTLs located outside a defined interval. This is achieved by using the properties of multiple regression analysis (ZENG 1993). Although multiple regression analysis has been used for mapping QTLs (Cowen 1989; STAM 1991), the theoretical properties of multiple regression analysis in relation to QTL mapping were analyzed in detail only recently by ZENG (1993) and also independently by RODOLPHE and LEFORT (1993). Moreover, ZENG (1993) proposed to utilize these properties to construct a composite interval mapping method to improve the precision and efficiency of mapping QTLs. Recently, Jansen (1993) also proposed a procedure to combine interval mapping with multiple regression for mapping QTLs. There are some similarities and also differences between the method analyzed in ZENG (1993) and here and that proposed by Jansen (1993). These are discussed at the end of this report.

In this article, following a presentation of background and problems and a summary of properties of multiple regression analysis, the method is elaborated in detail for a backcross design. Then the statistical issue to determine an appropriate critical value for a test is discussed. Simulation examples of QTL mapping are also presented to illustrate various properties of the method.

BACKGROUND AND PROBLEMS

Data type: Data for mapping QTLs consist of marker information (marker types for a number of polymorphic markers) and quantitative trait values for a number of individuals. Marker types for each marker can be recorded in digital form, such as 1 and 0 for distinguishing the two marker types (homozygote and heterozygote) for a backcross population from two inbred lines, and 2, 1 and 0 for distinguishing the three marker types (homozygote, heterozygote and another homozygote) for an F_2 population from two inbred lines. Based on segregation analysis, these markers can usually be ordered in linkage groups or are located linearly on chromosomes

Traditional method-simple linear regression: The simplest method of associating markers with quantitative trait variation is to test for trait value differences between different marker groups of individuals for a particular marker (e.g., Soller et al. 1976). For example, if we let $\tilde{\mu}_{M_1/M_1}$ and $\tilde{\mu}_{M_1/M_2}$ be the observed trait means of the groups of individuals with marker genotypes M_1/M_1 and M_1/M_2 for a particular marker in a backcross population, we can test for significance between means $\tilde{\mu}_{M_1/M_1}$ and $\tilde{\mu}_{M_1/M_2}$ using the usual t test. The hypotheses under the test can be

$$H_0: \mu_{M_1/M_1} = \mu_{M_1/M_2}$$
 and $H_1: \mu_{M_1/M_1} \neq \mu_{M_1/M_2}$.

Statistically, this is equivalent to the simple regression analysis with a model

$$y_i = b_0 + bx_i + e_i \quad j = 1, 2, \dots, n$$
 (1)

where y_j is the trait value of the jth individual in a population, b_0 is the mean (a parameter) in the model, x_j is a dummy variable for the jth individual, taking a value of 1 for marker type M_1/M_1 and 0 for marker type M_1/M_2 , $b = \mu_{M_1/M_1} - \mu_{M_1/M_2}$ is the simple regression coefficient, and e_j is a random residual variable for the jth individual. A test can be performed in this model on the regression coefficient.

To understand the relevance of this test to QTL mapping, we need to know what exactly is tested in genetic terms. Suppose that there are m QTLs contributing to the genetic variation in a backcross population from two inbred lines. Genetically the expected difference between $\tilde{\mu}_{M_1/M_1}$ and $\tilde{\mu}_{M_1/M_2}$ is

$$\varepsilon(\tilde{\mu}_{M_1/M_1} - \tilde{\mu}_{M_1/M_2}) = \sum_{i=1}^{m} (1 - 2r_i)a_i$$
 (2)

ignoring epistasis, where ε denotes expectation, a_i is the effect of the *i*th QTL expressed as a difference between the recurrent parent homozygote and the heterozygote, and r_i is the recombination frequency between the *i*th QTL and the marker. This means that essentially we are testing a composite parameter that

constitutes gene effects and recombination frequencies for (potentially) a number of genes. Of course, many QTLs may not be linked to the marker and thus have 0.5 recombination frequency. The above hypotheses are then equivalent to

$$H_0$$
: all $r_i = 0.5$ and H_1 : at least one $r_i < 0.5$,

because a_i 's are assumed to be non-zero (*i.e.*, by experiment we know that there are some genes which are segregating in the population). If $\tilde{\mu}_{M_1/M_1}$ and $\tilde{\mu}_{M_1/M_2}$ are found to be significantly different, it is indicated that the marker is linked to one or possibly more QTLs.

Although simple, this analysis captures the basic ideas of QTL mapping. Clearly there are many problems with this simple approach (LANDER and BOTSTEIN 1989), such as: (i) the method cannot tell whether the markers are associated with one or more QTLs, (ii) the method does not estimate the likely positions of the QTLs, (iii) the effects of QTLs are likely to be underestimated because they are confounded with the recombination frequencies and (iv) because of the confounding effects, the method is not very powerful and many individuals are required for the test.

LANDER and BOTSTEIN's interval mapping: If there is only one QTL on a chromosome, LANDER and BOTSTEIN (1989) proposed the use of a pair of markers to disentangle r and a from the test statistic. Specifically, for a backcross design they proposed the following linear model to test for a QTL located on an interval of markers i and i+1

$$y_i = b_0 + b^* x_i^* + e_i$$
 $j = 1, 2, ..., n$ (3)

where b^* is the effect of the putative QTL expressed as a difference in effects between the homozygote and heterozygote, x_j^* is an indicator variable, taking a value 1 or 0 with probability depending on the genotypes of markers i and i+1 and the position being tested for the putative QTL (Table 1). Statistically this is a mixture model (Titterington et al. 1985; McLachlan and Basford 1988). By using the property and procedures of mixture model analysis, Lander and Botstein built up a likelihood ratio test based on the hypotheses

$$H_0: b^* = 0$$
 and $H_1: b^* \neq 0$

assuming that the putative QTL was located at the point of consideration. This test can be performed at any position covered by markers and thus the method creates a systematic strategy of searching for QTLs. The evidence of QTLs is measured by the likelihood ratio test statistic (the so-called likelihood profile) at any particular location in the genome. If the likelihood profile at a region exceeds a pre-defined critical threshold, a QTL is indicated at the neighborhood of the maximum of the likelihood profile with the width of the neighborhood defined by the so-called support interval (LANDER and

TABLE 1 Specification of indicator variable x^*

	Marke	genotype	Sample	_
Group	i	i + 1	size	x*
1	+	+	n_1	1
2	+	-	n_2 $\bigg\{$	1 with probability $1 - p$ 0 with probability p
3	-	+	n_3 $\bigg\{$	1 with probability p 0 with probability $1 - p$
4	-	-	n_4	0

⁺ denotes homozygote of the marker genotype and – denotes heterozygote. In analysis, p can be treated either as a parameter or as a constant with $p = r_{iq}/r_{i(i+1)}$, where r_{iq} is the recombination frequency between marker i and the position q being tested for a putative QTL and $r_{i(i+1)}$ is the recombination frequency between markers i and i+1. Double recombination within the marker interval is ignored.

BOTSTEIN 1989). By the property of the maximum likelihood analysis, the estimates of locations and effects of QTLs are asymptotically unbiased if the assumption that there is at most one QTL on a chromosome is true. They also suggested methods to increase the power of QTL mapping, notably the selective genotyping of the extreme progeny.

Compared with the traditional method, the interval mapping method has several advantages. These include: (i) the probable position of the QTL can be inferred by the support interval, (ii) the estimated locations and effects of QTLs tend to be asymptotically unbiased if there is only one segregating QTL on a chromosome and (iii) the method requires fewer individuals than the traditional approach for the detection of QTLs.

There are, however, still many problems with the interval mapping. These include the following. (i) The test is not an interval test (a test which could distinguish whether or not there is a QTL within a defined interval and should be independent of the effects of QTLs that are outside of a defined region). Even when there is no QTL within an interval, the likelihood profile on the interval can still exceed the threshold significantly if there is a QTL at some nearby region on the chromosome. If there is only one QTL on a chromosome, this effect, though undesirable, may not matter because the QTL is more likely to be located at the region which shows the maximum likelihood profile. However, the number of QTLs on a chromosome is unknown. (ii) If there is more than one QTL on a chromosome, the test statistic at the position being tested will be affected by all those QTLs and the estimated positions and effects of "QTLs" identified by this method are likely to be biased (KNOTT and HALEY 1992; MARTINEZ and CURNOW 1992; see also below). (iii) It is not efficient to use only two markers at a time to do the test, as the information from other

markers is not utilized. These problems also apply to many other comparable QTL mapping methods (e.g., KNAPP et al. 1990).

Recognizing these problems, Lander and Botstein (1989) proposed to extend the method to analyze multiple markers for multiple QTLs simultaneously by introducing more b^* and x_i^* in the model (3). With this extension, some of above stated problems can be alleviated. But as the search now becomes multidimensional (LANDER and BOTSTEIN 1989; KNOTT and HALEY 1992), there are some difficulties in parameter estimation and model identifiability. As genetic structures for many quantitative traits can be very complex, a search in a space with unknown true dimension can be a problem. Both effort and ambiguity can be multiplied. Also, since the true number of QTLs on a chromosome is unknown, estimates of locations and effects of QTLs can still be biased if wrong models are fitted and tested. In addition useful information from other markers is still not used simultaneously in analysis by this method.

PROPERTIES OF MULTIPLE REGRESSION ANALYSIS

Ideally, when we test an interval for a OTL, we would like our test statistic be independent of the effects of possible QTLs at other regions of the chromosome. If such a test can be formulated, we can simplify the process of mapping multiple QTLs from a multiple dimensional search problem to an one dimensional search problem, as the test for each interval is independent and for each marker interval we can consider the possibility of the presence of only a single QTL. This test can be constructed by using a combination of interval mapping with multiple regression analysis (ZENG 1993). Various properties of the multiple regression analysis in relation to QTL mapping have been analyzed by ZENG (1993) and are summarized here. STAM (1991) has previously shown Property 1 for a special case in an unpublished conference paper. Recently, RODOLPHE and LEFORT (1993) also independently established many of these properties.

Suppose that we have a sample of n individuals from a backcross population with observations on a quantitative trait and t ordered markers, and suppose further that we analyze the data by the following linear regression model:

$$y_j = b_0 + \sum_{i=1}^{t} b_i x_{ji} + e_j$$
 for $j = 1, 2, ..., n$ (4)

where x_{ji} is the type of the *i*th marker in the *j*th individual. Definitions of other model variables and parameters are the same as (1) except that here more markers are included in the model. Note that b_i (also denoted by b_{yi,s_i} where s_i denotes a set which includes all markers except the *i*th marker) is the partial regression coefficient of phenotype γ on the *i*th marker conditional on

all other markers. For this analysis, the following properties have been established.

Property 1: In the multiple regression analysis (4), assuming additivity of QTL effects between loci (i.e., ignoring epistasis), the expected partial regression coefficient of the trait on a marker depends only on those QTLs which are located on the interval bracketed by the two neighboring markers, and is unaffected by the effects of QTLs located on other intervals. This property essentially says that a conditional (interval) test can be constructed based on the partial regression coefficient and such a test would test the linkage effect of only those QTLs which are located within the defined interval.

Property 2: Conditioning on unlinked markers in the multiple regression analysis will reduce the sampling variance of the test statistic by controlling some residual genetic variation and thus will increase the power of QTL mapping. This means that even unlinked markers contain useful information which can be used to increase the statistical power of the test and the efficiency of the genetic mapping. This useful information has not been utilized in the current QTL mapping methods.

Property 3: Conditioning on linked markers in the multiple regression analysis will reduce the chance of interference of possible multiple linked QTLs on hypothesis testing and parameter estimation, but with a possible increase of sampling variance. The first part of the sentence restates Property 1, and the second part of the sentence says that an interval test may entail a loss in the statistical power of the test because the test is a conditional test (see Zeng (1993) for details). This summarizes the advantage and disadvantage of the interval test: that is, there is a trade-off between precision and efficiency of mapping by using an interval test. Effective balance on these two issues will be the major consideration in practical mapping of QTLs (see below).

Property 4: Two sample partial regression coefficients of the trait value on two markers in a multiple regression analysis are generally uncorrelated unless the two markers are adjacent markers. This is related to the correlation between two test statistics in two intervals for an interval test. It has been shown that, for an interval test, a test statistic on an interval is generally asymptotically uncorrelated to the test statistic on another interval unless two intervals are adjacent intervals [Equation 13 of Zeng (1993)]. Even when the two intervals are adjacent intervals, the correlation between two test statistics in two intervals is usually very small. This property is related to the issue of determining an appropriate critical value of a test statistic under a null hypothesis for an overall test covering a whole genome (see below).

COMPOSITE INTERVAL MAPPING

Direct use of the multiple regression analysis (4) is not an appropriate way for mapping QTLs, because a partial regression coefficient is generally a biased esti-

mate of the relevant QTL effect (ZENG 1993). As stated above, however, there are several distinctive features of a multiple regression analysis which can be used to design a more accurate and efficient mapping method. Based on the above properties of multiple regression analysis, an interval test procedure is developed which combines interval mapping with multiple regression analysis to fully utilize the information in mapping data. This is illustrated here. For simplicity, let us consider data from a backcross population. The population is assumed to be derived from two inbred lines which are fixed for different alleles at m QTLs and t genetic markers. The data consist of observations on a quantitative trait and t ordered markers of n individuals. Suppose that the t markers are more or less evenly distributed in a genome.

Suppose that we want to test for a QTL on a marker interval (i, i + 1). We can use markers i and i + 1 as an indicator for the genotype of the putative QTL within the interval, and write the statistical model as

$$y_{j} = b_{0} + b^{*}x_{j}^{*} + \sum_{k \neq i, i+1} b_{k}x_{jk} + e_{j}$$
for $j = 1, 2, ..., n$
(5)

where y_i is the trait value of the jth individual, b_0 is the mean of the model, b^* is the effect of the putative QTL expressed as a difference in effects between homozygote and heterozygote, x_i^* is an indicator variable, taking a value 1 or 0 with probability depending on the genotypes of markers i and j and the position being tested for the putative QTL (Table 1, ignoring double recombination within the marker interval), b_k is the partial regression coefficient of the phenotype y on the kth marker, x_{ik} is a known coefficient for the kth marker in the jth individual, taking a value 1 or 0 depending on whether the marker type is homozygote or heterozygote, and e_i is a random variable. The summation of other markers in the model depends on the balance of the trade-off of Property 3 and also on the consideration of degrees of freedom of the test (see below).

Assuming that e_j 's are identically and independently normally distributed with mean zero and variance σ^2 , the likelihood function is given by

$$L_1 = \prod_{j=1}^{n} [p_j(1)f_j(1) + p_j(0)f_j(0)]$$
 (6)

where $p_j(1)$ gives a prior probability of $x_j^*=1$ (Table 1), $p_j(0)=1-p_j(1)$, $f_j(1)$ and $f_j(0)$ specify a normal density function for the random variable y_j with a mean $b_0+b^*+\sum_{k\neq i,\,i+1}b_kx_{jk}$ and $b_0+\sum_{k\neq i,\,i+1}b_kx_{jk}$, respectively, and a variance σ^2 . By differentiating the likelihood function (6) with respect to individual parameters, setting the derivatives equal to zero and then solving the equations, the maximum likelihood (ML)

estimates of the parameters b^* , b_k 's and σ^2 are found to be the solutions of

$$\hat{b}^* = (\mathbf{Y} - \mathbf{X}\hat{\mathbf{B}})'\hat{\mathbf{P}}/\hat{c} \tag{7}$$

$$\hat{\mathbf{B}} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'(\mathbf{Y} - \hat{\mathbf{P}}\hat{b}^*)$$
 (8)

$$\hat{\sigma}^2 = [(\mathbf{Y} - \mathbf{X}\hat{\mathbf{B}})'(\mathbf{Y} - \mathbf{X}\hat{\mathbf{B}}) - \hat{c}\hat{b}^{*2}]/n \tag{9}$$

where **Y** is a $(n \times 1)$ vector of y_j 's, $\hat{\mathbf{B}}$ is a $((t-1) \times 1)$ vector of the ML estimates of b_k 's (including b_0 but excluding b^*), **X** is an $(n \times (t-1))$ matrix of x_{jk} 's, $\hat{\mathbf{P}}$ is a $(n \times 1)$ vector with elements \hat{P}_j specifying the ML estimate of the posterior probability of $x_i^* = 1$:

$$\hat{P}_i = p_i(1)\hat{f}_i(1)/[p_i(1)\hat{f}_j(1) + p_i(0)\hat{f}_i(0)]$$
 (10)

and

$$\hat{c} = \sum_{j=1}^{n} \hat{P}_{j}.$$
 (11)

The prime indicates transposition of a vector or matrix. Note that if p is treated as a parameter, the ML estimate of p is the solution of

$$\hat{p} = \frac{\sum_{j=1}^{n_2} (1 - \hat{P}_j) + \sum_{j=1}^{n_3} \hat{P}_j}{n_2 + n_3},$$
(12)

and the above equations, where the summations $\sum_{j=1}^{n_2}$ and $\sum_{j=1}^{n_2}$ indicate sums of those individuals belonging groups 2 and 3 of Table 1.

These estimates can be found by iteration of the above equations via the expectation/conditional maximization (ECM) algorithm (Meng and Rubin 1993) beginning with the initial estimate $\hat{b}^* = 0$ or the least squares estimates of b^* and **B** using $x_j^* = p_j(1)$. In each iteration, the algorithm consists of one E-step, Equation 10, and three CM-steps, Equations 7, 8 and 9. The convergence of the algorithm to ML estimates has been proven by Meng and Rubin (1993) as their condition (3.6) is satisfied in this case. The advantage of this algorithm over the full EM algorithm (maximizing \hat{b}^* and $\hat{\mathbf{B}}$ simultaneously in the M step) is that the inverse, $(\mathbf{X}'\mathbf{X})^{-1}$, does not need to be updated, and thus the efficiency of the numerical evaluation is improved substantially.

The hypotheses to be tested are H_0 : $b^* = 0$ and H_1 : $b^* \neq 0$. The likelihood function under the null hypothesis is

$$L_0 = \prod_{j=1}^n f_j(0)$$

with the ML estimates

$$\hat{\mathbf{B}} = (\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'$$

$$\hat{\mathbf{\sigma}}^2 = (\mathbf{Y} - \mathbf{X}\hat{\mathbf{B}})'(\mathbf{Y} - \mathbf{X}\hat{\mathbf{B}})/n.$$

The likelihood ratio (LR) test statistic is found to be

$$\begin{aligned} \text{LR} &= -2\ln(L_0/L_1) \\ &= n(\ln \hat{\sigma}^2 - \ln \hat{\sigma}^2) \\ &+ \sum_{j=1}^{n_2} \left[2\ln[(1-p)\exp(\hat{d}_j/2) + p] - \hat{P}_j \hat{d}_j \right] \\ &+ \sum_{j=1}^{n_3} \left[2\ln[p\exp(\hat{d}_j/2) + (1-p)] - \hat{P}_j \hat{d}_j \right], \end{aligned} \tag{13}$$

where

$$\hat{d}_i = [2\hat{b}^*(y_i - \hat{b}_0 - \sum_{k \neq i, i+1} \hat{b}_k x_{ik}) - \hat{b}^{*2}]/\hat{\sigma}^2.$$

If there is no epistasis, estimates of position, p, and effect, b^* , of a QTL by this method are unaffected by other linked QTLs if there are markers which separate those QTLs from the QTL under consideration and these markers are fitted in the model as a control [Property 1]. By the property of maximum likelihood, these estimates also tend to be asymptotically unbiased (see Table 4 below). [However, if we estimate the effects of QTLs conditional on that the test for QTLs is significant as we usually do in practice, the estimates of QTL effects can still be biased (Table 4).] Thus by combining interval mapping with multiple regression, this method creates a condition that individual QTLs can be separated for testing and estimation.

There will be, however, some interference on testing and estimation between those QTLs which are located in adjacent marker intervals when using this composite interval mapping method because two flanking markers are used for interval mapping. If there are indeed two major QTLs located in two adjacent marker intervals, depending on the positions of QTLs on the intervals and the sizes of the intervals the likelihood profile may be very significant for both intervals and some parts of their adjacent intervals, and in some cases may show bimodes. Under the interval test, this may indicate the possibility of the presence of two QTLs in two adjacent intervals. In this case, if necessary, two variables may be fitted on the intervals to test for two QTLs together with other markers, but it may need very large sample size to test the hypotheses.

BEHAVIOR OF THE TEST STATISTIC UNDER THE NULL HYPOTHESIS

Like interval mapping, this test can also be performed at any position in the genome covered by markers. Thus the method creates a systematic search for QTLs and reduces a multiple dimensional search problem for multiple QTLs to a one dimensional search problem. Since this is a multiple test situation (for multiple locations), a practical question arises of determining the critical

TABLE 2

Mean, variance (var.) and 95 percentile (95%) of the test statistics for a particular position, for a marker interval and for the whole genome under the null hypothesis

		Average	e for each p	osition	Average m	aximum for eac	h interval	Ov	erall maxim	um
M	n	Mean	Var.	95%	Mean	Var.	95%	Mean	Var.	95%
1	500	1.01	2.09	3.80	1.39	2.72	4.47	1.39	2.72	4.47
5	500	1.03	2.11	3.93	1.63	3.19	5.15	3.43	5.59	7,99
10	500	1.03	2.10	3.95	1.64	3.19	5.25	4.60	6.12	9.31
20	500	1.05	2.14	4.03	1.70	3.41	5.35	6.07	6.79	10.91
40	500	1.10	2.46	4.22	1.78	3.79	5.69	7.95	8.34	13.23
1	200	0.99	2.03	3.88	1.38	2.70	4.65	1.38	2.70	4.65
5	200	1.03	2.15	3.88	1.62	3.20	5.20	3.47	5.68	7.97
10	200	1.06	2.24	4.04	1.70	3.49	5.42	4.79	6.69	9.63
20	200	1.11	2.42	4.26	1.79	3.75	5.71	6.28	7.60	11.61
30	200	1.18	2.98	4.50	1.91	4.94	5.97	7.65	9.52	13.64
40	200	1.29	3.48	4.98	2.10	5.71	6.65	9.03	10.85	15.20

All markers are linked on one chromosome with 10 cM for each marker interval. Replicates of simulations are 1000 for sample size n = 500 and 2000 for n = 200.

value of the test statistic for the test performed in a genome. To determine an appropriate critical value of the test statistic for this testing method, we need to know the behavior of the test statistic under the null hypothesis, not only for a particular interval, but for a whole genome, because the entire genome is tested for the presence of QTLs. Lander and Botstein (1989) discussed the issue of the critical value of the test statistic (using the LOD score) for the simple interval mapping. The threshold of the test statistic for the composite interval mapping is, however, different. The difference is that with multiple regression the test statistic is more or less uncorrelated for different intervals [Property 4]. Thus to achieve an overall significance level α for the test with M intervals, a nominal significance level α/M may be used for the test in each interval, a situation corresponding to the sparse-map case of Lander and Botstein (1989).

However, the distribution of the maximum of LR over an interval (i.e., the distribution of the LR of (13) taking p as a parameter) under the null hypothesis is not clear. This distribution will depend greatly on sample size, the number of markers fitted in the model and the genetic size of each interval. To investigate the behavior of the test statistic, a series of simulations were performed under the null hypothesis. Although our null hypothesis is that there is no QTL on the relevant interval being tested, no QTL in the genome was simulated to examine the behavior of the test statistic in a genome. This does not make much difference under our model because segregating QTLs on other intervals are more or less controlled by conditioning markers in the model so that they would not greatly influence the hypothesis testing on the interval being tested (cf. Lander and Botstein 1989). Two extreme situations, linked and unlinked cases, were simulated. In the linked case, M equally spaced marker intervals (with M+1 markers) are located on one chromosome, and in the unlinked case, M equally spaced marker intervals (with $2\,M$ markers) are located on different chromosomes. For each replicate of simulations, n backcross individuals were simulated on M+1 (or $2\,M$) markers (with the recombination frequency r for each marker interval) and a quantitative trait (simulated simply as a normal random variable). Analysis was performed as stated above, and LR test statistic was calculated by (13) along the genome at every 1-cM position.

Results are presented in Table 2 and Figure 1. Table 2 gives the means, variances and 95 percentiles (over replicates) of LR for a particular position [i.e., LR] of (13) taking a fixed value for p], the maximum of LR for a marker interval [i.e., LR] of (13) taking p as a parameter] and the overall maximum of LR over a genome under the null hypothesis. There is generally little difference among statistics of LR from position to position and from interval to interval, so only averaged statistics (mean, variance and 95 percentile) of LR over all positions and over all marker intervals are presented. Figure 1 plots the 95 percentiles of the overall maximum of LR against the number of intervals for sample sizes (n) 500 and 200.

These results strongly suggest that, when the sample size is large and the number of markers fitted in the model is relatively small, the LR test statistic for fixed testing position (i.e., for fixed p) is approximately χ^2 distributed with 1 degree of freedom (with mean 1, variance 2 and 95 percentile 3.84), as might be expected [but see Goffinet et al. (1992) for a special case]. When the sample size is relatively small and the number of markers fitted in the model is large, the observed LR deviates from χ_1^2 distribution, most likely due to slow convergence of the statistic to χ_1^2 distribution.

The maximum of the LR over a marker interval for the cases considered generally fall between χ_1^2 and χ_2^2 (with mean 2, variance 4 and 95 percentile 5.99) distributions in the simulations (except for the cases for n = 200 and

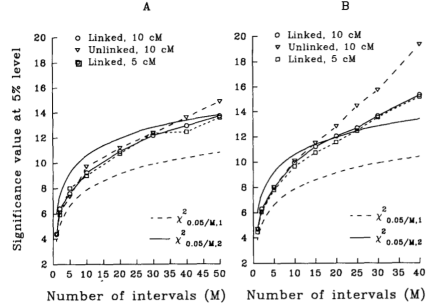


FIGURE 1.—The 95 percentile of the overall maximum of the test statistic under the null hypothesis is plotted against the number of intervals (M) from simulation. Three cases are plotted. In the unlinked case, all marker intervals (with 10 cM) are on different chromosomes. In the linked cases (one with a 10-cM interval and one with a 5-cM interval), all marker intervals are on one chromosome. The values of $\chi^2_{0.05/M,1}$ and $\chi^2_{0.05/M,2}$ are also plotted for reference. Sample size is 500 for graph A with 2000 replicates and 200 for graph B with 5000 replicates.

M = 30 and 40 in which the number of degrees of freedom for the test is reduced significantly). The distribution depends on the sample size, the number of markers fitted in the model and the genetic size of the interval. The correlations of maxima of LRs between intervals are generally very small (positive) [Property 4]. (The average correlations for adjacent intervals are generally less than 0.3 and the average correlations for non-adjacent intervals are effectively close to zero.) The effect of marker interval size (comparing the cases of 10-cM interval with 5-cM interval in Figure 1) tends to be minor, compared with other factors such as sample size, number of markers fitted in the model and number of intervals tested. Thus it appears that when the sample size is large and the number of markers fitted in the model is not too many, $\chi^2_{\alpha/M,2}$ can be used as an approximation for the $100\alpha\%$ critical value for an overall test with M intervals in a genome for the model (5) (Figure 1A). (Although only the 5% critical value is presented in Figure 1, other critical values such as 10% show similar patterns.)

However, in practice, sample sizes of mapping data are generally not very large, and markers are unlikely to be evenly spaced in the genome. In that case, it may not be appropriate and desirable to fit too many markers in the model even when they are unlinked to the interval being tested, because too many markers fitted in the model can substantially increase the critical value of the test statistic [for example, comparing the cases of linked (M+1) markers with unlinked (2M) markers in Figure 1B], and thus reduce the power of the test. Many markers in mapping data are also usually clustered in some regions. If that is the case, it would be more appropriate to use in the model as a control only those selected markers which are more or less evenly spaced in the genome or those preidentified markers (e.g., by stepwise regres-

sion) which explain most of the genetic variation in the genome. Other markers can still be used for testing QTLs at relevant intervals. For many data, the above criterion may be used as a guide (not necessarily for $\alpha = 0.05$). If necessary, computer simulations can be used to determine an appropriate critical value for the test for a given data set.

QTL MAPPING SIMULATION ANALYSIS

Methods: To illustrate the properties, utility, advantages and disadvantages of the method, a simulation study of mapping QTLs was performed. Four "chromosomes" each with 16 markers separated in 15 10-cM intervals were simulated for a backcross population. The trait is affected by 10 QTLs with positions and effects given in Table 3 and depicted in Figure 2. Together the QTLs account for 70% of the phenotypic variance in a backcross population. Sample size is 300. The trait value of an individual is determined by the sum of effects of the QTLs which the individual possesses, plus a random (environmental) variable which is normally distributed with mean zero and variance scaled to give the expected 0.7 heritability of the population. Both marker and QTL types were simulated for each individual with the linkage map specified above.

This data set can be fitted for analysis by numerous models. For the purpose of illustration, three simplified models were fitted in the analysis following the procedures outlined above:

- *Model I:* Composite interval mapping with *k* in model (5) summed over all other markers (solid curves of Figure 2);
- Model II: Semi-composite interval mapping with k
 in model (5) summed over all unlinked markers
 (long-dashed curves of Figure 2);

TABLE 3

Parameters and point estimates of positions and effects of QTLs from one replicate of simulation

	C	hromosom	e 1	C	Chromosom	e 2	C	hromosom	e 3	Chromosome
					Parameter	s				
Position (cM): Effect:	16 0.42	48 0.75	108 0.58	3 1.02	43 -1.23	77 -1.26	33 -0.46	68 1.61	129 0.88	$\begin{array}{c} 26 \\ 0.74 \end{array}$
Model I				P	oint estima	tes				
Position (cM)		48		6	43	70		CC	100	
Effect		1.02		1.29	-1.16	78 -1.18		$\frac{66}{1.68}$	130	
Model II		1.02		1.49	-1.16	-1.16		1.00	1.39	
Position (cM)		45			7	3		68	130	32
Effect		1.18				1.43		1.67	1.52	0.69
Model III						2.10		1.07	1.52	0.03
Position (cM)		45			6	5		69	125	
Effect		1.42				1.46		1.60	1.83	

Point estimates of QTL effects are made at the peaks of the likelihood profile in the regions where the presence of QTLs is indicated (see Figure 2).

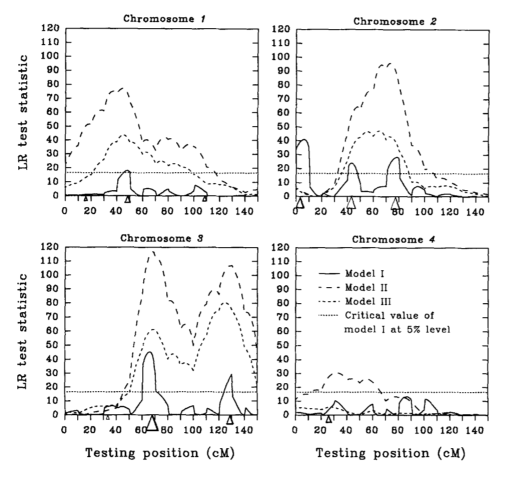


FIGURE 2.—A simulation ample of QTL mapping on an hypothetical backcross population. Likelihood ratio test statistic is calculated and plotted at every 1-cM position of the four "chromosomes" to give a likelihood profile. The genetic length of each "chromosome" is 150 cM with markers at every 10 cM. Ten OTLs were simulated with positions indicated by triangles. The size of each triangle is in proportion to the magnitude of the effect of the QTL (Table 3). Dark-sided triangles are used to indicate for OTLs with positive effects, and light sided triangles for QTLs with negative effects. Three curves (likelihood profiles) are plotted for the three models fitted for analysis: model I (solid curves) is the composite interval mapping, model II (longdashed curves) is the semicomposite interval mapping and model III (short-dashed curves) is the simple interval mapping. The dotted line is the simulated 5% critical value, 16.5, for the test under model I.

• *Model III:* Simple interval mapping with model (3) (short-dashed curves of Figure 2).

The dotted lines of Figure 2 give the simulated overall 5% critical value, 16.5, for the test of model I (based on a simulation with 5000 replicates under the null hypothesis), which is a little higher than $\chi^2_{0.05/60,2} = \chi^2_{0.00083,2} = 14.2$. The overall 5% critical values for models II and III are smaller (12.6 for model II and 10.7 for model III). Some point estimates of positions and effects of QTLs at

the peaks of the likelihood profile in the regions where the presence of QTLs is indicated are given in Table 3 for reference. Both Figure 2 and Table 3 depict results of mapping of a single replicate of simulation.

To show the general patterns of the results, 100 replicates of simulation were performed based on the same set of parameters and the same procedures of analysis. Results are given in Table 4. In this table, means and standard deviations of estimates of positions and effects

Parameters and summary statistics of estimates of positions and effects of OTLs from 100 replicates of simulation TABLE

Position (cM): 16 48 1 Effect: 0.42 0.75 Model I Powera Position (cM) ^b 496 (6.4) Position (cM) ^c 50.0 (5.2) Effect ^b 6.43 Model II 0.12 Model II 1.14 (0.12) Position (cM) ^b 48.2 (13.3)	108 3 0.58 1.02 0.87 3.4 (2.9) 3.3 (2.4)	Parameters 43 -1.23 Summary statistics 0.83 42.1 (3.4) 41.8 (3.0)	77 -1.26 .s 0.80 76.8 (2.9) 76.8 (2.6)	33 68 -0.46 1.61 0.99 68.8 (2.2	129	
cM): 16 48 0.42 $0.751 \text{ (cM)}^b 0.22 0.221 \text{ (cM)}^c 0.83 0.26)1.14$ 0.12) 1.00 1.00	58 1.02 0.87 3.4 3.3	43 -1.23 Summary statisti 0.83 42.1 (3.4) 41.8 (3.0)	77 -1.26 0.80 76.8 76.8	0.99	0.88	
$\begin{pmatrix} \text{cM} \end{pmatrix}^b \\ \text{1 (cM)}^c \\ \text{1 (cM)}^b \end{pmatrix}$	0.87 3.4 (2.9) 3.3 (2.4)	Summary statisti 0.83 42.1 (3.4) 41.8 (3.0)	0.80 76.8 76.8	0.99		26 0.74
$\begin{pmatrix} \operatorname{cM} \end{pmatrix}^b \\ \operatorname{cM} \rangle^c \\ \operatorname{cM} \rangle^b \\ \operatorname{cCM} \rangle^b $	0.87 3.4 (2.9) 3.3 (2.4)	0.83 42.1 (3.4) 41.8 (3.0)	0.80 76.8 (2.9) 76.8 (2.6)	68.8 (2.2		
$\begin{pmatrix} \text{(CM)}^b \\ \text{(CM)}^c \\ \text{(CM)}^b \end{pmatrix}$	3.4 (2.9) 3.3 (2.4)	42.1 (3.4) 41.8 (3.0)	76.8 (2.9) 76.8 (2.6)	68.8 (2.2	0.58	0.17
1 (cM) ^c 1 (cM) ^b	3.3 (2.4)	41.8 (3.0)	76.8 (2.6)	1 1 1 1		27.1 (6.2)
1 (CM) ^b	1 0 1 1			68.9 (2.1		27.9 (4.5)
1 (CM) ^b	1.02 (0.24) $1.08 (0.20)$	-1.24 (0.23) -1.29 (0.21)	-1.24 (0.25) -1.32 (0.20)	1.62 (0.26) $1.63 (0.24)$	0.94 (0.23) $0.94 (0.23)$	$0.78 (0.26) \\ 1.12 (0.19)$
$_{1}$ $(cM)^{b}$	(01:0)	5	1			
		1.00	7	1.00	1.00	0.99
		-1.67	0.15)	1.65 (0.13)		0.74 (0.13)
Model III Power ^a 1.00		1.00		1.00	1.00	0.58
(cM) ^b 5		71.3 (8.9)	3.9) 3.18)	70.3 (2.6)) 121.9 (10.9) 3) 1.34 (0.19)	27.6 (7.1) 0.76 (0.20)

estimates based on all 100 replicates. estimates based on the replicates which are significant in the test at the relevant regions. Empirical estimate of the power of the test for a QTL at the region indicated based on 100 replicates of simulation. Mean and standard deviation (in parentheses) of estimates based on all 100 replicates. Mean and standard deviation (in parentheses) of estimates based on the replicates which are significant in the test a of QTLs which are indicated in Figure 2 are reported. For each replicate, estimates were made at the distinctive peaks of the likelihood profile around the relevant regions. Although Figure 2 is an analysis based on a single replicate, the general patterns of mapping among 100 replicates conform to that of Figure 2. There are, however, a few replicates for which the mapping under models II and III indicates only one rather than two QTLs on "chromosome" 3. In these cases, to make the results comparable with other replicates, the results were still interpreted to indicate two QTLs and estimates of positions and effects of QTLs were made at relevant local maxima of the likelihood profile. Empirical estimates of the power of the test (i.e., the ratio of the replicates that are significant at the relevant regions over the total replicates) are also presented in Table 4.

Results: In this analysis, neither model II nor model III controls the effects of possible QTLs located at other regions of the chromosome when testing for a QTL at a particular chromosome position. Model II, however, effectively removes, from the residual variance of the model, most of the variation due to segregating QTLs located on unlinked chromosomes, so that the LR test statistic on most intervals under this model is significantly larger than those under the other two models [Property 2]. This model has the highest statistical power, among the three models, for detecting marker and QTL linkages. However, as the test under model II (as well as model III) is not an interval test, this model does not necessarily have high probability of locating individual QTLs accurately, because estimates of position and effect of a QTL can be influenced by other linked QTLs. Only model I provides an interval test in which the test statistic on an interval is unaffected by all those QTLs which are located outside the interval being tested and its two adjacent intervals. This is an advantage of the interval test. Model I also effectively removes from the model residual variance most of the variation due to segregating QTLs. However, because the test under model I is a conditional test [Property 3, see Zeng (1993) for more detailed discussion on the theoretical ground], the test statistic on many intervals is smaller than those under models II and III (Figure 2). This is a disadvantage of the interval test.

Nevertheless model I correctly identifies the six largest QTLs with relatively higher resolution in Figure 2, judged by the relatively accurate positions of the significant peaks of the likelihood profile and also by the slope of the likelihood profile (see also Table 4). This increase of precision of mapping is gained by making the test conditional on nearby markers so that the sensitivity of the test statistic to the position of a QTL is increased and the position effect of a QTL is emphasized in a short region of the interval conditioned. This is another advantage of the interval test.

Models II and III give two significant peaks of the likelihood profiles on "chromosome" 3 in Figure 2 which

correctly indicate the presence of two major QTLs (the third QTL is minor), because the two QTLs are separated by a sufficiently long distance. On "chromosome" 2, however, the likelihood profiles of models II and III seem to indicate the presence of a single OTL at wrong positions. This is due to the presence of three QTLs with comparable (and opposite) effects in a relatively close range. Should this occur, we could be deceived by using wrong models. Of course, these data can be fitted and analyzed for two or three QTLs simultaneously as suggested by Lander and Botstein (1989), but the statistical tests for two QTLs vs. one QTL, and three QTLs vs. two QTLs are not easy matters. In real applications, the number of QTLs on a chromosome (i.e., the true genetic model for testing) is unknown, and we can very easily be deceived by fitting and testing the wrong models. Also, even when the correct region of a OTL can be identified under models II and III, the likelihood profiles under models II and III tend to be flatter (i.e., the supporting intervals as defined by LANDER and BOTSTEIN (1989) are wider) (Figure 2) and the precision of mapping is still relatively lower (Table 4).

It seems that only model I is more likely to give unbiased estimates of position and effect of QTLs (Table 4). However, as discussed above, model II has the highest statistical power for identifying possible QTL regions, and the statistical power of model I for identifying QTLs is relatively lower (and can be very low), because of the fitting of the closely linked markers in the model [Property 3]. Only model II strongly indicates the presence of a QTL on "chromosome" 4 in Figure 2. In practice, with limited data, some combinations of models I and II (i.e., deleting or inserting some linked markers as a control in the model) may have to be used to maximize the probability of detecting QTLs while controlling the precision of mapping (i.e., balancing the type I and type II errors of the statistical test). This is illustrated, as an example, by a further analysis on "chromosome" 1 of the simulation example given in Figure 2. On this chromosome, both models II and III give a major peak of the likelihood profile in Figure 2, which would indicate the presence of a QTL somewhere on the fifth interval. However, model II also seems to suggest a second QTL roughly located between positions 70 and 110 cM for which the test under model I failed to show significance. To test this hypothesis, a test can be performed between 60 and 150 cM using a model combining model II with six more markers located between 0 and 50 cM for background control. This would eliminate the effect of the QTL mapped on the fifth interval (and also other possible QTLs on the left of 50 cM position) in the test and estimation. Indeed this test strongly indicates the presence of another QTL with effect estimated to be 0.52 at position 100 cM (Figure 3).

On the accuracy of estimates of QTL effects, we have to distinguish two issues affecting the relative accuracy

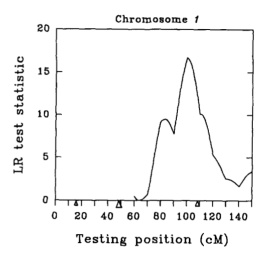


FIGURE 3.—A reanalysis of Figure 2 for mapping for a QTL on the region between 60 and 150 cM of "chromosome" *I*. The test for a QTL at each position is conditional on fitting 6 markers located between 0 and 50 cM of "chromosome" *I* and all other unlinked markers. There is strong evidence for the presence of one more QTL on this chromosome by this analysis. The effect of the QTL is estimated to be 0.52 at position 100 cM

of estimates: bias and sampling error. Estimates of QTL effects under model I tend to be less biased compared with those under models II and III, because the estimates under model I are unaffected by QTLs that are located at other marker intervals [Property 1] (Table 4). However, the estimates under model I will have larger sampling variance than those under model II [Property 3] (Table 4). If we know that there is at most one QTL on a chromosome, model II tends to have both higher statistical power for the test of QTL and smaller sampling variance for the estimate of QTL effect (see, for example, chromosome 4 of Table 4); otherwise the test and estimation under model II (and also model III) can be misleading and QTLs can be mapped to wrong positions (or intervals).

Admittedly, in this analysis, we may have used more markers in models I and II to control genetic background than necessary. There are, in general, three roles that a marker can play in mapping QTLs, depending on the chromosome position being tested. A marker can be used to construct the indicator variable x^* for interval mapping, can be used to provide a boundary for an interval test, and can also simply be used to control the residual genetic variation in test and estimation. For the third role, a marker is informative only when it is linked to QTLs. For each chromosome, there are probably only a few markers which are closely linked to QTLs. (We do not know, of course, how many QTLs are on a chromosome.) Given that a few markers that are closely linked to QTLs are fitted in the model, other markers will not be informative [Property 1] and tend to be redundant for fitting in the model. Redundant fitting of markers can reduce the power of the test and increase the sampling variance of estimates particularly when the sample size is small. In practice, we may need first to select a few markers for each chromosome or some chromosomes by, for example, stepwise regression and use those markers to control genetic background when mapping QTLs by interval test.

DISCUSSION

For mapping QTLs, unbiasedness and accuracy should be more important than other issues of mapping techniques, such as ease of computation. Because quantitative trait variation is generally influenced by multiple genes, we have to take into account effects of possible multiple (linked) genes when designing a mapping method. Effective control of individual gene effects is a key to increasing the precision of mapping. Every effort should be made in mapping to preserve the accuracy and quality of mapping and at the same time to maximize the chance of finding more QTLs.

A new QTL mapping framework is proposed in this paper. The key feature of the approach is the idea of an interval test which tries to separate and isolate individual QTL effects when testing and mapping for QTLs. Depending on data and underlying genetic mechanisms, the precision of mapping can be significantly improved. The method works better for traits with high heritability, because most of the genetic variation can be controlled and removed from the residual variation in the model by conditioning on multiple markers. For traits with low heritability, the gain of fitting multiple markers in the model as a control may not justify the cost in losing statistical power of finding a QTL due to both increased sampling variation of estimates if closely linked markers are fitted in the model for an interval test [Property 3], which would reduce the test statistic, and increased threshold of the test (this is due to both increased number of intervals tested and reduction of degrees of freedom of the test). In that case, the first task might be to increase the chance of finding anything in the genome. In practice, the best strategy would be to fit data with multiple models to identify an appropriate model which balances both type I and type II errors of the test.

In this study, only the backcross population design is analyzed. Other population designs, such as F₂ populations, and dominant genetic models can readily be implemented in this framework. Epistasis is, however, ignored here. With possible epistatic effects of QTLs, this mapping method can still be biased. Problems with analyzing epistatic effects of genes in mapping QTLs are that there could be many types of genetic epistasis and that the underlying genetic parameters for epistasis are not well defined. In principle, epistatic effects can be fitted in the model for mapping QTLs if the type of genetic epistasis is identifiable (e.g., HALEY and KNOTT 1992).

Regression analyses have been used for mapping

QTLs in various ways by HALEY and KNOTT (1992), MAR-TINEZ and CURNOW (1992), MORENO-GONZALEZ (1992), and Jansen (1992, 1993). Both Haley and Knott (1992) and MARTINEZ and CURNOW (192) used the regression analysis simply to approximate the simple interval mapping procedures, although they also suggested using a bivariate regression analysis to search the twodimensional space along the chromosome for mapping two QTLs. Moreno-Gonzalez (1992) proposed a stepwise multiple regression procedure to fit multiple markers (with multiple "QTLs" arbitrarily assumed to be located in the middle of each marker interval) in the model for mapping QTLs. It appears that this procedure is very arbitrary and imprecise. JANSEN (1992) described a mixture model analysis in the framework of the interval mapping. He, however, gave a simulation example of using a third marker as a covariable in the linear model as a control, which has some similarity to the method proposed here. Very recently, JANSEN (1993) also proposed a procedure for mapping QTLs which combines interval mapping with multiple regression. There are some similarities between the method of ZENG (1993) (and also here) and that of JANSEN (1993). But clearly, there are critical differences in concepts and procedures to map multiple linked QTLs. The method used in this paper is the interval test. The emphasis is to control the precision of mapping as much as possible. JANSEN (1993), however, used a procedure of testing multiple markers on a chromosome simultaneously to indicate possible multiple QTLs on a chromosome, which seems to be very imprecise for mapping multiple QTLs. Moreover, many theoretical and statistical properties and behaviors of their method were not analyzed and discussed by Jansen (1993).

In summary, there are four advantages of this mapping strategy. (i) First, by confining the test to one region at a time, it reduces a multidimensional search problem (for multiple QTLs) to a one-dimensional problem, and also estimates of locations and effects of individual QTLs are likely to be asymptotically unbiased. (ii) Second, by conditioning on linked markers in the test, the precision of QTL mapping can be greatly improved. (iii) Third, by selectively conditioning multiple markers in the test, the method simultaneously utilizes more information in the data to make inferences and should be more informative and efficient for mapping QTLs than the current methods. (iv) Fourth, it can still use the QTL likelihood map (the likelihood profile) to present the strength of the evidence for QTLs at various positions along the entire genome, and preserves the feature of interval mapping. These advantages are brought about by the realization that a complete linkage map can be used not only to provide an anchor to fix a position to test for a QTL anywhere in a genome covered by markers (interval mapping), but also to provide a boundary condition for the test, and at the same time to

control the residual genetic variation in the rest of the genome for the test (interval test).

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