Mapping Quantitative Trait Loci in Crosses Between Outbred Lines Using Least Squares

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ABSTRACT

The use of genetic maps based upon molecular markers has allowed the dissection of some of the factors underlying quantitative variation in crosses between inbred lines. For many species crossing inbred lines is not a practical proposition, although crosses between genetically very different outbred lines are possible. Here we develop a least squares method for the analysis of crosses between outbred lines which simultaneously uses information from multiple linked markers. The method is suitable for crosses where the lines may be segregating at marker loci but can be assumed to be fixed for alternative alleles at the major quantitative trait loci (QTLs) affecting the traits under analysis (e.g., crosses between divergent selection lines or breeds with different selection histories). The simultaneous use of multiple markers from a linkage group increases the sensitivity of the test statistic, and thus the power for the detection of QTLs, compared to the use of single markers or markers flanking an interval. The gain is greater for more closely spaced markers and for markers of lower information content. Use of multiple markers can also remove the bias in the estimated position and effect of a QTL which may result when different markers in a linkage group vary in their heterozygosity in the F_1 (and thus in their information content) and are considered only singly or a pair at a time. The method is relatively simple to apply so that more complex models can be fitted than is currently possible by maximum likelihood. Thus fixed effects and effects of background genotype can be fitted simultaneously with the exploration of a single linkage group which will increase the power to detect QTLs by reducing the residual variance. More complex models with several QTLs in the same linkage group and two-locus interactions between QTLs can similarly be examined. Thus least squares provides a powerful tool to extend the range of crosses from which QTLs can be dissected whilst at the same time allowing flexible and realistic models to be explored.

UR ability to study gene action underlying quantitative variation has been greatly enhanced by the rapid development of genetic maps based on DNA markers combined with the development of statistical methods which allow the mapping of some of the loci responsible for quantitative variation (quantitative trait loci or QTLs). Among the statistical methodologies, interval mapping (LANDER and BOTSTEIN 1989) has been shown to be a powerful tool for the analysis of populations derived from crosses between inbred lines [e.g.,PATERSON et al. (1988, 1991), JACOB et al. (1991), and STUBER et al. (1992)]. In the method of interval mapping the intervals between pairs of flanking markers are explored in turn for evidence of the presence of a QTL at various positions between the markers. The methods were originally implemented using maximum likelihood (LANDER and BOTSTEIN 1989), in which information on the presence of a QTL is derived from both the mean differences between the flanking marker genotype classes and from the distribution of the trait within each marker genotype class. Compared to methods

which consider only a single marker at a time, interval mapping methods have been shown to provide some additional power and much more accurate estimates of QTL effect and position and to be relatively robust to failure of normality assumptions (LANDER and BOTSTEIN 1989; KNOTT and HALEY 1992a).

The disadvantage of maximum likelihood based methods for interval mapping is their computational complexity, which makes them relatively difficult to extend to allow the simultaneous analysis of several linked QTLs, interactions between QTLs, effects of unlinked QTLs and fixed effects (e.g., treatment and sex). The advantage of such simultaneous analyses is their potential to remove bias and to increase the power (by reducing the residual "noise" variance) of the analyses performed. We have recently demonstrated that ordinary least squares can be used for interval mapping and provides very similar estimates and test statistics to those obtained from maximum likelihood (HALEY and KNOTT 1992). This allows relatively complex (and potentially more realistic) models to be used without placing severe demands on computational resources (and incidentally demonstrates that the great majority of information extracted using maximum likelihood derives from mean differences between marker genotype classes, rather

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than from the distribution within the marker genotype class).

In a cross between two inbred lines the markers selected for mapping have heterozygosities of unity in the F_1 , as do any QTLs segregating in the cross. This greatly simplifies the analysis and means that, for co-dominant markers under the assumption of no interference, it is only the pair of markers flanking an interval that provide information on the transmission of a QTL within that interval. Thus markers can be considered a pair at a time without loss of information. In many cases, however, it is desirable to map QTLs in crosses between lines which are genetically divergent but are outbred. Often it may be reasonable to assume that the lines are fixed, or nearly so, for QTLs of moderate or large effect (i.e., those that it is feasible to consider mapping) even though some or all of the markers which are informative in the cross are segregating within each outbred line. Examples of such a situation would include experimental lines which have undergone divergent selection or long established breeds of plants or animals which have very different selection histories. Crosses in the latter category would include that between the Chinese Meishan pig and European commercial breeds, which differ for many traits (HALEY and ARCHIBALD 1992) or between Ndama and Boran cattle, which differ in their resistance to tick-borne disease (SOLLER 1990). In such cases it is impractical to produce inbred lines from the original outbred populations. Even for lines of experimental organisms which can have several generations per year, developing inbred lines may be time consuming and costly.

BECKMANN and SOLLER (1988) presented a method for the analysis of crosses between outbred lines based on tracing marker alleles through the three generations $(e.g., parents, F_1 and F_2)$ of the cross. A potential problem in the analysis is that the markers are not all completely informative and will vary in their heterozygosity in the F₁ cross. To overcome this problem BECKMANN and SOLLER (1988) suggested screening a number of markers in each chromosomal region and for each individual \mathbf{F}_1 cross selecting a marker that would be informative in the F_{2} in the region. This would be potentially wasteful of information, for to obtain at least one informative marker, several would need to be scored and rejected if not required. An alternative to this approach would be to develop interval mapping so that it could be applied to this data structure. However, using only flanking markers would lead to the same situation observed in analyses within outbred populations, that is that information, and thus power to detect a QTL, varies from interval to interval depending upon the markers flanking that interval. This can lead to biases in the estimated position and effect of a QTL (KNOTT and HALEY 1992b). To make most efficient use of marker data and thus to maximize experimental power and to minimize the risk of biased estimates it is necessary to take into account information from all of the informative markers in a linkage group. In this paper we develop a simple method which allows least squares to be applied to the mapping of QTLs in a cross between outbred lines using data from all markers in a linkage group simultaneously. The relative efficacy of using all markers in a linkage group compared to using only those flanking an interval is demonstrated by the analysis of simulated data.

METHOD

In the least squares method of mapping QTLs phenotypic values are regressed onto genetic coefficients calculated for a putative QTL at a fixed position. In the analysis of the generations derived from a cross between inbred lines the probability of an F2 individual, for example, being each of the three possible genotypes at a QTL in a given position in an interval can be calculated conditional solely upon the genotypes at the markers flanking that interval and the estimated recombination fraction between the markers and the QTL. The additive coefficient for the QTL in that individual is then the difference between the conditional probabilities of the two homozygous QTL genotypes and the dominance coefficient is equal to the conditional probability of the individual being the QTL heterozygote. (In this parameterization the additive and dominance coefficients, a and d, respectively, are defined as deviations from the mean of the two homozygotes for the QTL, i.e., the difference between the homozygotes is 2a.) For each putative QTL position, ordinary linear least squares can be used to regress the trait value for each individual onto their calculated additive and dominance coefficients. This provides estimates of a and d for that position. The procedure is repeated for chosen fixed positions (e.g., at 1-cM intervals) through a linkage group and the best estimate of the QTL effects and position are obtained at the position at which the residual sum of squares is minimized. Multiple QTL effects can be fitted by regression onto the coefficients for several QTLs in different positions (in the same or different linkage groups) simultaneously. We have previously described the method for inbred line crosses in more detail and shown that it gives very similar results to those produced by maximum likelihood (HALEY and KNOTT 1992). The method we develop here for analysing outbred line crosses is very similar in conception. The key to applying this method is developing a simple means of calculating the coefficients of a and d for each individual for a QTL in each putative position conditional upon multiple markers in a linkage group.

We develop the method throughout using an F_2 cross as an example, but the same method could be applied to the analysis of other types of cross. We consider that genotypic data is available on the F_2 individuals and their parents and grandparents and phenotypic data is avail-

QTL Mapping in Outbred Line Crosses

TABLE 1

	Line 1:		Line 2:	 I	ine 1:		Li	ne 2:		
sire of the sire (SS) $A_1A_1B_2B_2C_1C_2D_1D_1$		dam	of the sire (DS)	sire of t	sire of the dam (SD)		dam of the dam (DD)			
			$A_2A_2B_1B_2C_2C_3D_2D_2$		$A_1A_1B_2B_2C_2C_4D_1D_2$		$A_2A_2B_2B_2C_1C_4D_1D_2$			
1 1		Sire (S)				Dam (D)		2		
	A_1A	$_{2}B_{1}B_{2}C_{1}C_{3}D_{1}D_{2}$			$A_1 A_2$	$B_2 B_2 C_1 C_2 D$	$B_2 B_2 C_1 C_2 D_1 D_2$			
	Line 2:	Line 1:	A Line 2:	$_{2}A_{2}B_{1}B_{2}C_{2}C_{3}D_{2}D_{2}$			Ma	rker		
Line 1:	Line 2.									
sire of	dam of the sire	sire of the dam	dam of the dam	Line origin combination	Putative QTL	A	В	С		
sire of	dam of			0	QTL	A	В	С		
sire of he sire	dam of	the dam		combination 11 12	QTL	A	В			
	dam of	the dam	the dam	combination 11		A	<i>B</i> *	C *		

Example F_2 pedigree from a cross between outbred lines with four markers (A, B, C and D) and possible line origin combinations of marker alleles

Possible line origins of alleles are indicated by \times . For each line origin combination the putative QTL genotype at a locus fixed for allele Q in line 1 and allele q in line 2 is shown. The possible line origin combinations for the F_2 individual in the pedigree shown above are shown by asterisks in the right-hand side of the table.

able on the F_2 individuals. For any locus (marker or QTL) an F_2 individual must receive one allele from either the sire or dam of its sire and one allele from either the sire or dam of its dam. There are thus four possible combinations of alleles in terms of the outbred line from which they came. These line origin combinations of alleles are shown in Table 1. As an example, consider the three generation pedigree from a cross between two outbred populations and marker genotypes for an individual and its parents and grandparents which is shown in Table 1. For the first marker (A) it is clear that the F, individual has inherited two alleles from line 2, one from the dam of the sire and one from the dam of the dam (line origin combination 22 in Table 1). For the second marker (B), one allele in the F_2 individual has been inherited from line 2 (from the dam of the sire) but the inheritance of the second allele is equivocal because the dam is homozygous. Thus for marker B both of line origin combinations 21 and 22 are possible. For the third marker (C) only line origin combination 21 is possible (one allele from the dam of the sire and one from the sire of the dam). For a QTL fixed for alternative alleles in the two grandparental lines each line origin combination will correspond to one QTL genotype as shown in Table 1.

For each marker the potential line origin combinations can be derived for each individual in turn. For some markers in some individuals all four line origin combinations may be possible (*e.g.*, if all grandparents are homozygous for the same marker allele or if marker data is missing for that individual) and thus these markers are uninformative. For dominant markers for which the two lines are fixed for alternative alleles, F_2 individuals either have a single line origin combination possible (if they are homozygous for the recessive allele) or three are possible (if they are homozygous for the dominant allele or heterozygous). Note that for both codominant and dominant loci, in any case where two or more line origin combinations are possible in an F_2 individual, those that are possible are equally likely.

This simple method of assigning line origin to the markers considering each F₂ individual in turn is easily implemented and rapid. It should be noted, however, that it does not necessarily use all of the information. Consider marker D in Table 1. Both the sire of the dam and the dam of the dam are heterozygous for the same alleles as is the dam. The dam is thus an uninformative heterozygote because, considering only a single F, individual and this marker, line origin combinations 21 and 22 are both possible and they have equal probability (*i.e.*, it is not possible to infer whether offspring of this dam have inherited their allele from the sire of the dam or from the dam of the dam at this marker). As in most studies F2 individuals would be from groups of full and half sibs, the joint use of information from sibs and flanking markers would indicate the likely linkage phase in the dam and thus allow the relative probabilities of line origin combinations 21 and 22 to be calculated. However, the maximum expected frequency of such uninformative heterozygous parents is 0.125 (for markers at which both lines have two alleles at equal frequencies) and it will usually be much less, as many markers are multi-allelic and allele frequencies at segregating markers are often markedly different in different lines. Thus in many cases the benefits of the rapid and simple analysis possible treating each F, individual in turn will often outweigh the small amount of extra information that might be recovered by considering sibs jointly.

The probability of the line origin combination of a locus conditional upon the line origin combination of a linked locus and the recombination fractions between the two loci

Line origin	Line origin combination					
combina- tion	11	12	21	22		
11	$(1-r_m)(1-r_f)$	$(1-r_m)r_f$ $(1-r_m)(1-r_f)$	$r_m(1-r_f)$	$r_m r_f$		
12 21	$(1-r_m)(1-r_f) (1-r_m)r_f r_m(1-r_f)$	$(1-r_m)(1-r_f)$ $r_m r_f$ $r_m (1-r_f)$	$r_m r_f$ (1- r_m) (1- r_f) (1- r_m) r_f	$ \begin{array}{c} r_{m}r_{f} \\ r_{m}(1-r_{f}) \\ (1-r_{m})r_{f} \\ (1-r_{m})(1-r_{m}) \end{array} $		
22	$r_m r_f$	$r_m(1-r_f)$	$(1-r_m)r_f$	$(1-r_m)(1-r_m)$		

Conditional probabilities for the line origin combinations given in Table 1. The recombination frequencies between the pair of loci in the male and female parent are r_m and r_p respectively.

Once the possible line origin combinations of the markers have been derived, the probabilities of each of the four line origin combinations for a QTL at a given position in an F₂ individual can be calculated conditional upon the possible line origin combinations of the markers, the previously estimated recombination fractions between the markers and the recombination fraction between the assumed position of the QTL and the markers. Table 2 gives the probability of the line origin combination of a locus conditional upon the line origin combination of a linked locus and the recombination fraction between the two loci. The probabilities in Table 2 have been written allowing for different recombination rates in the two sexes; when these are the same Table 2 can be simplified. For each individual, each section of chromosome between pairs of markers which both have only a single possible line origin combination can be considered separately (as, in the absence of interference, markers outside this section provide no information about the line origin of positions within the section). The probability of each of the four line origin combinations for any point between these markers (i.e., the position of a putative QTL) conditional upon the observed marker genotypes can be calculated as a product of the probabilities shown in Table 2 for any possible combination of line origins of the markers scaled so that the total over all possible combinations sums to one. The formal derivation of this method is shown in the APPENDIX.

To clarify the calculation of these conditional probabilities, consider the example in Table 3. Marker loci A, B and C are used (these correspond to markers A, Band C for the pedigree of the F_2 individual shown in Table 1). Note that as markers A and C are fully informative, with only a single line origin combination each, markers outside this region add no further information for positions within the region for this individual. Consider there to be a QTL (Q) at the midpoint between markers B and C. The first possible combination of line origins shown in Table 3 for the three markers and the QTL is 22, 21, 11 and 21 for A, B, Q and C, respectively. As shown in Table 2, the conditional probability of a 21 line origin combination at marker B given a 22 line origin combination at marker A is $(1 - r_{ABm}) r_{ABP}$ where r_{ABm} and r_{ABt} are the recombination fractions between loci A and B in males and females, respectively. (This combination of line origins at markers A and B requires there to have been no recombination between markers A and B in the first (male) F_1 parent and a single recombination between the markers in the second (female) F_1 parent). Similarly, the conditional probability of a 11 line origin combination at the QTL given a 21 line origin combination at marker B is $r_{BOm}(1 - r_{BOf})$ and the conditional probability of a 21 line origin combination at marker C given a 11 line origin combination at the QTL is $r_{QCm}(1 - r_{QCf})$. The probability of the line origin combinations 21, 11 and 21 at B, Q and C conditional on a 22 line origin combination at A is thus the product of these probabilities:

or:

$$(1 - r_{AB})r_{AB} r_{BQ}(1 - r_{BQ})r_{QC}(1 - r_{QC})$$

 $(1 - r_{ABm})r_{ABf}r_{BQm}(1 - r_{BQf})r_{QCm}(1 - r_{QCf})$

on the assumption of equal recombination frequencies in males and females. Division by the sum of these probabilities for the possible line origin combinations (eight are possible in the example in Table 3) gives the probability conditional on the possible line origin combinations (and thus on the observed marker genotypes).

Once the conditional probabilities for the line origin combinations have been calculated the coefficients for a and d for a putative QTL in this position can be determined as:

- a: probability of line origin 11 conditional on the marker genotypes minus probability of line origin 22 conditional on the marker genotypes
- d: probability of line origin 12 conditional on the marker genotypes plus probability of line origin 21 conditional on the marker genotypes

or in the notation used in the APPENDIX:

$$a: \operatorname{prob}(\omega_{11} | \mathbf{P}) - \operatorname{prob}(\omega_{22} | \mathbf{P})$$

and

$$d: \operatorname{prob}(\omega_{12} | \mathbf{P}) + \operatorname{prob}(\omega_{21} | \mathbf{P})$$

where $\text{prob}(\omega_i | \mathbf{P})$ is the probability of line origin combination *i* for a QTL at a given position conditional on the observed marker genotypes in the individual and its parents and grandparents.

After calculation of the predicted coefficients for a putative QTL in a given position for all individuals, a and d can be estimated for that position by ordinary least squares, regressing the phenotypic values on to these coefficients. Several (or many) putative QTLs in a number of positions (linked or unlinked) can be fitted simultaneously and covariates or fixed effects can also be included in the model. For a fixed position of QTL, the ratio of the regression mean square to the

0.328

combinations of flanking markers					
Line origin combination of alleles (A-B-Q-C)	Conditional probability	Example: $r_{AB} = r_{BQ} = r_{QC} = 0.$			
22-21-11-21	$r_{AB}(1-r_{AB})r_{BQ}(1-r_{BQ})r_{QC}(1-r_{QC})/P$	0.004			
22-22-11-21	$(1-r_{AB})^2 r_{BQ}^2 r_{QC} (1-r_{QC})/P$	0.004			
22-21-12-21	$r_{AB}(1-r_{AB})r_{BQ}^2r_{QC}^2/P$	0.000			
22-22-12-21	$(1-r_{AB})^2 r_{BQ} (1-r_{BQ}) r_{QC}^2 / P$	0.004			
22-21-21-21	$r_{AB}(1-r_{AB})(1-r_{BQ})^2(1-r_{QC})^2/P$	0.328			
22-22-21-21	$(1-r_{AB})^2 r_{BQ} (1-r_{BQ}) (1-r_{QC})^2 / P$	0.328			
22-21-22-21	$r_{AB}(1-r_{AB}) r_{BO}(1-r_{BO}) r_{OC}(1-r_{OC})/P$	0.004			

Example calculation of the probabilities of line origin combinations of a putative QTL conditional upon the possible line origin combinations of flanking markers

For this individual the line origin combination (as defined in Table 1) of the first marker (A) is 22, that of the second marker (B) may be either 21 or 22 and that of the third marker (C) is 21. The putative QTL (Q) is placed between B and C. The recombination frequencies are assumed to be the same in both sexes and that between A and B is r_{AB} , that between B and the putative position of the QTL is r_{BQ} and that between the QTL and C is r_{QC} . P is the sum of the numerators of the conditional probabilities. An example is given for $r_{AB} = r_{BQ} = r_{QC} = 0.1$. For this individual the predicted coefficient for a would be -0.324 (= 0.004 + 0.004 - 0.004 - 0.328) and for d would be 0.660 (= 0.000 + 0.004 + 0.328 + 0.328).

 $(1-r_{AB})^2(1-r_{BQ})^2r_{QC}(1-r_{QC})/P$

residual mean square provides the usual variance (F) ratio test statistic.

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An alternative approximate log-likelihood ratio test statistic is provided by:

$$n \log_{e} \left(\frac{\text{residual sum of squares reduced model}}{\text{residual sum of squares full model}} \right)$$

where *n* is the number of observations. This test statistic is distributed approximately as a chi-square with degrees of freedom equal to the number of parameters included in the full model (*i.e.*, estimating the QTL effects) but omitted from the reduced model (*i.e.*, omitting QTL) (AITKEN *et al.* 1989). Dividing this test statistic by ($2\log_e 10$) would approximately give the LOD score. The use of LOD is of little relevance, however, for tests such as this which have more than a single degree of freedom.

When fitting a single QTL any of these test statistics can be plotted against position to give a curve or when fitting two QTLs this can be visualized as a surface (HALEY and KNOTT 1992). The maximum point of the curve or surface indicates the most likely position of the QTL and this point will be at the same position for any of the test statistics. We use the approximate loglikelihood ratio test statistic throughout this paper for consistency and to facilitate comparison with our previous work (*e.g.*, HALEY and KNOTT 1992; KNOTT and HALEY 1992a, b).

SIMULATIONS

General: The analysis of simulated data was used to explore the characteristics of the method. Each set of data included 500 F_2 individuals in 50 full-sib families of size 10 with their parents and grandparents. The genotype of each individual comprised a pair of chromosomes 100 cM in length. Depending upon the simulation there were either three markers at 50 cM spacing,

six markers at 20 cM spacing or eleven markers at 10 cM spacing. Markers of three types were generated, either fixed for alternative alleles in the two grandparental lines (*i.e.*, as in a cross between inbred lines), or segregating with the same two alleles at equal frequency in both grandparental lines or segregating with the same four alleles at equal frequency in both grandparental lines. QTLs of various effect and position were simulated (see below). In the analyses the additive and dominance effects (a and d, respectively) of a single QTL were estimated sequentially at each 1-cM point along the chromosome, with the distance between the markers set at that used to generate the data. The point along the chromosome at which the test statistic was highest was used to provide the estimates of the QTL position and effect for that analysis. Unless otherwise stated, 100 replicates were simulated and analyzed for each combination of parameters. The data were generated and analyzed using programs written in FORTRAN 77, supplemented with routines from the NAG library (Numerical Algorithms Group 1990) for random number generation and for ordinary least squares analysis (routine G02DAF).

All markers vs. flanking markers and size of QTL: To explore the general properties of the method and the advantage of using all markers on a chromosome, its behavior was compared to ordinary least squares in which only the pair of markers flanking an interval was used to predict the probabilities of each QTL genotype at a given point within the interval. For each replicate in these analyses a single set of phenotypic data (including effect of QTL and residual variance) was generated with a QTL of additive effect (*i.e.*, half the difference between the homozygotes) of either 0.25, 0.5 or 1.0 residual (*i.e.*, within QTL genotype) standard deviations 30 cM from one end of the chromosome. QTLs of these

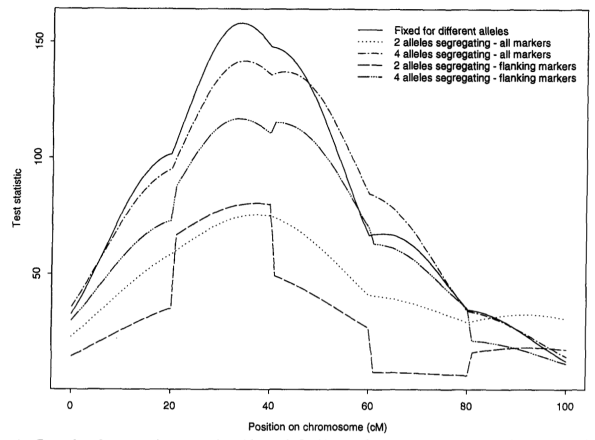


FIGURE 1.—Examples of test statistic curves using either only flanking markers or all markers. For each analysis markers were all of one of three levels of information content (fixed for alternative alleles in the two lines, or segregating with either two or four alleles at equal frequency in both lines) and were spaced at 20-cM intervals starting at 0 cM. The simulated QTL was additive in effect with two residual standard deviations between homozygotes and was located at the 30-cM position on the chromosome. The phenotypic data were the same for all analyses.

sizes would account for 3.3%, 11.1% or 33.3%, respectively, of the variance in the F₂ population. Each set of phenotypic data was analyzed with markers at 20-cM spacing which were all of one of the three levels of information content (*i.e.*, fixed for alternative alleles in the two grandparental lines or segregating with either the same two alleles at equal frequency or with the same four alleles at equal frequency in the two grandparental lines).

Marker density: Data were generated with a QTL with an additive effect of 0.5 residual standard deviation at 25 cM from one end of the chromosome. Each set of data had markers of the three levels of information content at either 10-cM spacing or at 50-cM spacing and were analyzed using information from all markers simultaneously.

Position of QTL: Data generated with a QTL with an additive effect of 0.5 residual standard deviations at either 10 or 50 cM from one end of the chromosome. Each set of data had markers of the three levels of information content at 20-cM spacing and were analyzed using information from all markers simultaneously.

Markers of varying information content: In these analyses markers varied in their information content along the chromosome. Data were generated with a QTL with an additive effect of 0.5 residual standard deviations at 30 cM from one end of the chromosome and markers at 20-cM spacing. For each set of data the first three markers (at positions 0, 20 and 40 cM) were of low information content (*i.e.*, the same two alleles at equal frequency segregating in each grandparental line) and the last three markers (at 60, 80 and 100 cM) were of higher information content (all three either having four alleles at equal frequency segregating in the grandparental lines or being fixed for alternative alleles in the grandparental lines). The data were analysed using either information only from flanking markers or using information from all markers simultaneously.

Null hypothesis: Data were generated with no QTL but with markers at either 10-, 20- or 50-cM spacing. For each marker density, markers of the three levels of information content were used to analyze the data. The data were analyzed using information from all markers simultaneously. For each combination of parameters 1000 replicates were generated and analyzed.

RESULTS

All markers vs. flanking markers: Examples of the curves produced by plotting the values of the test statistic

Relative mean test statistics and empirical standard deviations of parameter estimates using either only flanking markers or all
linked markers

		QTL effect	Test statistic		sD of test statistic		sp of position		SD of additive effect	
Marker type		simulated (a)	F	A	F	A	F	A	F	A
Fixed, alternative alleles in two lines	20	1.0	(155.5	155.5	20.2	20.2	1.9	1.9	0.066	0.066)*
	20	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
	10	0.5		0.35		0.72		2.11		1.09
	20	0.5	0.32	0.32	0.60	0.60	2.15	2.15	0.96	0.96
	50	0.5		0.21		0.49		5.37		1.30
	20	0.25	0.10	0.10	0.34	0.34	8.16	8.16	1.17	1.17
Segregating, 4 alleles in each line	20	1.0	0.81	0.88	0.87	0.91	1.74	1.16	1.09	1.06
	10	0.5		0.33		0.70		2.16		1.12
	20	0.5	0.27	0.29	0.54	0.59	4.00	3.42	1.08	1.05
	50	0.5		0.16		0.46		6.63		1.59
	20	0.25	0.09	0.09	0.32	0.33	9.58	9.32	1.21	1.14
Segregating, 2 alleles in each line	20	1.0	0.41	0.53	0.62	0.81	3.58	2.37	1.78	1.53
0 0 0	10	0.5		0.26		0.56		4.26		1.15
	20	0.5	0.15	0.18	0.40	0.47	8.11	6.79	1.56	1.39
	50	0.5		0.09		0.34		11.18		2.20
	20	0.25	0.07	0.07	0.24	0.26	13.37	13.32	1.83	1.45

F, A, analyses using flanking or all markers, respectively. Each value is based upon 100 replicate simulations. Simulated QTLs were located 30 cM from one end of the 100-cM chromosome with 20-cM spaced markers or 25 cM from one end of the 100-cM chromosome with 10- and 50-cM spaced markers. Simulated QTLs were additive in effect. The size of the additive effect (a) of the QTL is given as half the difference between the homozygotes in terms of the residual (*i.e.*, within QTL genotype) standard deviation.

* Absolute values this line only, all other values are given relative to these.

against the chromosomal position are shown in Figure 1. These curves were produced from a single set of phenotypic data analysed using markers of one of the three levels of information content at 20-cM spacing and either predicting the QTL genotype using just flanking markers or using all markers on the chromosome. The two curves (using just flanking markers or using all markers) produced when the markers were fixed for alternative alleles in the two lines are exactly the same, confirming that the flanking markers contain all the information on the interval between them for this type of marker under the assumption of no interference. For the less informative markers the use of just flanking markers produces steps in the curve between intervals and the maximum value of the test statistic reduces with the information content of the markers. The steps result because the different markers, and hence intervals, vary by chance in the information they contain. The use of multiple markers to predict the QTL genotype removes these steps and also increases the test statistic, although the maximum test statistic still increases with increasing marker information content.

For data in which the markers in a linkage group were all of the same type, estimates of position and effect of the QTL for the analyses using either all markers or just flanking markers were very close to those simulated and are not shown. The relative values of the mean (over replicate simulations) of the maximum test statistic on the chromosome and the empirical (over replicate simulations) standard deviations of estimates of position and additive effect for the analyses using either all markers or just flanking markers are shown in Table 4 for these analyses. Trends for the dominance effect were similar to those for the additive effect and are not shown. When the markers were fixed for alternative alleles in the two grandparental breeds, the results from analyses using all or flanking markers were, as expected, identical. For the less informative markers using all markers in the analysis generally increased the maximum test statistic. The largest increase (approximately 30%) was found for the least informative markers and for the QTL of largest effect, whereas for the QTL of smallest effect, no increase in the maximum test statistic was observed when using all markers rather than just flanking markers. The empirical standard deviation of the estimates of position and effect was decreased for markers which were not completely informative by using all markers in the analysis. For position, the magnitude of the decrease in the empirical standard deviation was greatest for the QTL of largest effect, but for the additive effect the magnitude of the decrease in the empirical standard deviation was greatest for the QTL of smallest effect.

Marker density: The relative mean of the maximum test statistic on a chromosome and empirical standard deviations of estimates of position and effect for the analyses using all markers at 10 and 50 cM spacing and the QTL at 25 cM are shown in Table 4. The relative increase in the maximum test statistic as the marker density increases is greatest for the least informative markers. Moving from 50-cM to 10-cM spaced markers increased the maximum test statistic by 2.7-fold for the least informative markers (two alleles segregating in each line) compared to a 1.6 fold increase for the most informative markers (fixed for alternative alleles in the

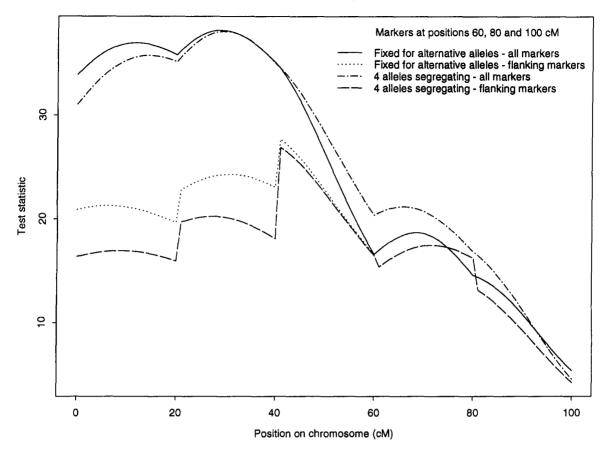


FIGURE 2.—Examples of test statistic curves using either flanking markers only or all markers and markers which vary in information content along the chromosome. In both cases markers at positions 0, 20 and 40 cM were of relatively low information content (two alleles at equal frequency in the two lines) and those at positions 60, 80 and 100 cM were of relatively high information content (case 1: with four alleles at equal frequency segregating in the two lines or case 2: fixed for alternative alleles in the two lines). The simulated QTL was additive in effect with one residual standard deviation between homozygotes and was located at the 30 cM position on the chromosome. The phenotypic data were the same for all analyses.

two lines). This difference can be explained by the fact that an increase in marker density when markers are not completely informative increases the probability that a QTL is flanked by informative markers, whereas this is not the case if the markers are already completely informative.

Position of QTL: The mean of the maximum test statistic on a chromosome and estimates of position and effect for the analyses using all markers at 20-cM spacing were little affected by the position of the QTL no matter what the information content of the markers. For a QTL at 10 and 50 cM, respectively, the mean maximum test statistics over 100 replicates were 50.3 and 48.2 for markers fixed for alternative alleles in the two lines, 27.9 and 29.2 for markers with the same two alleles at equal frequency in the two lines and 44.2 and 43.4 for markers with the same four alleles at equal frequency in the two lines. Thus for this marker spacing, position of the QTL has little effect on the power of its detection, despite a QTL at the centre of the chromosome having a greater chance of being flanked by two informative markers.

Markers of varying information content: Examples of the curves produced by plotting the values of the test

statistic against the chromosomal position when markers vary in information content along the chromosome are shown in Figure 2. These curves were produced from the analysis of data in which there was a QTL at 30 cM and the first three markers had the same two alleles segregating at equal frequency in the two grandparental breeds and the last three markers were of higher information content (either four alleles segregating in both breeds or fixed for alternative alleles in the two breeds). In these analyses the use of just flanking markers results in the highest test statistic being in the third interval, which is the first interval to be flanked by a marker of relatively high information content. Analyzing the same data using all markers results in the highest test statistic being in the second interval, which contained the simulated QTL.

The mean maximum test statistics and estimates of position and effect for the analyses using either all or just flanking markers at 20-cM spacing are shown in Table 5. The use of just flanking markers results in a significant bias in the estimated position of the QTL toward the more informative markers, with the bias being greater for the most informative markers. The use of all markers in the analysis both removes the bias in the estimated

QTL Mapping in Outbred Line Crosses

TABLE	5
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Mean test statistics and parameter estimates with markers varying in information content along the chromosome

M 1	OTI effect		Markers used			
Marker type (last 3 markers)	QTL effect simulated (a)	Parameter	Flanking	All		
Fixed, alternative alleles in two lines	0.5	Test statistic Position (cM) Additive effect (a)	26.9 (8.7) 36.8 (12.9) 0.473 (0.103)	$\begin{array}{ccc} 32.2 & (10.1) \\ 29.6 & (8.9) \\ 0.493 & (0.085) \end{array}$		
Segregating, 4 alleles in each line	0.5	Test statistic Position (cM) Additive effect (a)	26.0 (8.2) 34.5 (15.2) 0.498 (0.113)	33.1 (10.2) 29.0 (8.9) 0.511 (0.088)		

Each mean is based upon 100 replicate simulations and analyses and is shown with its empirical standard deviation over the replicates in parentheses. All simulated QTLs were located 30 cM from one end of a 100-cM chromosome with 20-cM spaced markers and were additive in effect. In each case the first three markers on the chromosome (at 0, 20 and 40 cM) were relatively low in information content, with the same two alleles at equal frequency segregating in both grandparental lines, and the last three markers (at 60, 80 and 100 cM) were more informative. The size of the additive effect (a) of the QTL is given as half the difference between the homozygotes in terms of the residual (*i.e.*, within QTL genotype) standard deviation.

position of the QTL and increases the mean maximum test statistic.

Null hypothesis: Results of the analyses of data generated with no QTL but with markers at either 10-, 20or 50-cM spacing are shown in Table 6. For all marker densities and markers of different information content a test performed at a fixed position on the chromosome has a mean and a standard deviation of close to 2, as expected for a test statistic distributed as a chi-square with 2 d.f. (2 d.f. as both an additive and a dominance effect have been estimated). The mean over replicates of the highest test statistic on the chromosome increases both with increasing marker density and with marker information content. The approximate empirical 5% threshold calculated from these simulations as the mean of the 50th and 51st highest test statistics over replicates is also shown in Table 6; this value increases with marker density but shows no consistent trend with marker information content.

DISCUSSION

The explosion in the availability of molecular genetic markers and the rapid development of linkage maps based on these markers is providing the geneticist with new tools to explore the genome (SOLLER and BECKMANN 1988; TANKSLEY et al. 1989). Interval mapping (LANDER and BOTSTEIN 1989) has proven to be a powerful tool for the dissection of some of the genetic factors underlying quantitative genetic variation in crosses between inbred lines (e.g., PATERSON et al. 1988, 1991; JACOB et al. 1991; STUBER et al. 1992). As more species become amenable to this form of probing, however, statistical tools are needed to analyze a wider range of types of population. We have shown here that, as is the case for crosses between inbred lines (HALEY and KNOTT 1992), a simple ordinary least squares method can be applied to the analysis of populations resulting from crosses between outbred populations which are fixed for alternative QTL alleles.

The least squares method is relatively simple to apply and can extract most of the information contained in multiple linked markers. The use of all the markers in a linkage group simultaneously increases the test statistic, and thus the power for the detection of QTLs. It also removes bias in the estimated position and effect of a QTL which can result when markers vary in information content and are only considered a pair at a time [Table 5 and KNOTT and HALEY (1992b)].

The prediction of the QTL genotype using all markers in a linkage group is more difficult than the use of just the flanking markers, which is all that is required for crosses between inbred lines. Thus programming the entire analysis in the language of a single statistical package, as is possible for inbred lines (HALEY and KNOTT 1992), becomes intractable. Once the predicted QTL genotypes have been calculated using a custom written computer program, however, these can be stored and the remainder of the analysis performed using a general statistical package.

The least squares analysis is very rapid and the time taken for computation does not increase greatly with the number of parameters estimated, as it does in many maximum likelihood analyses. Thus the great advantage of least squares methods, other than their simplicity, is that many parameters can be fitted simultaneously. This first allows the inclusion of fixed effects such as treatment or sex in the model. Second, when exploring one chromosome, background genetic noise attributable to the other chromosomes can be reduced by the inclusion of QTLs at reasonable (say 30-50 cM) intervals down the remaining chromosomes in the model [e.g.,]ANSEN (1992)]. Both of these strategies should increase the power to detect QTLs on the chromosome under study by minimizing the residual variance. Third, several linked QTLs can be fitted simultaneously to the chromosome under study (HALEY and KNOTT 1992). (It is unlikely, however, that most studies carried out at present will be of sufficient scale to detect more than two

TABLE	6
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Test statistic distribution under the null hypothesis

		Marker spacing				
Marker type	Statistic	10 cM	20 cM	50 cM		
Fixed, alternative alleles in two lines	Mean test statistic	2.09 (2.09)	2.06 (2.03)	1.95 (1.97)		
	Highest test statistic	5.49 (2.85)	4.80 (2.65)	3.92 (2.56)		
	Empirical 5% threshold	10.96	9.67	8.51		
Segregating, 4 alleles in each line	Mean test statistic	2.10 (2.05)	2.09 (2.05)	1.98 (1.93)		
	Highest test statistic	5.23 (2.76)	4.66 (2.61)	3.88 (2.41)		
	Empirical 5% threshold	10.85	9.85	8.34		
Segregating, 2 alleles in each line	Mean test statistic	2.08 (2.09)	2.08 (2.11)	2.04 (2.06)		
	Highest test statistic	4.72 (2.80)	4.22 (2.72)	3.79 (2.58)		
	Empirical 5% threshold	10.07	9.12	8.81		

The mean test statistic is based upon the mean of 101 positions (1-cM intervals on a 100-cM chromosome) over 1000 replicate simulations and analyses. The empirical standard deviation of the test statistic over the 1000 replicates averaged over the 101 positions is shown in parentheses. The highest test statistic represents the mean over 1000 replicates and its empirical standard deviation over the replicates is given in parentheses. The empirical 5% threshold is calculated as the mean of the 50th and 51st highest test statistics over the 1000 replicates.

or three linked QTLs.) This can remove bias introduced when linked QTLs are present but only a single QTL is fitted in the analysis (HALEY and KNOTT 1992; KNOTT and HALEY 1992a; MARTINEZ and CURNOW 1992). Fourth, more complex models of QTL gene action can be explored relatively easily, for example two locus epistasis (HALEY and KNOTT 1992). Finally, using the same predicted probabilities of QTL genotype the data could be analyzed using a generalized linear model, which is again possible in a number of statistical packages [*e.g.*, AITKEN *et al.* (1989)]. This would allow QTLs underlying non-continuously distributed traits, such as binomial threshold traits, to be detected and mapped.

The need for the outbred populations to be fixed, or nearly so, for alternative QTL alleles may be considered restrictive, but in fact many populations may be of this type for some traits. Such populations would include divergently selected experimental lines and breeds with very different selection histories. When the populations crossed are not fixed for alternative QTL alleles, the power to detect a QTL will be increasingly reduced and its effect will be increasingly underestimated as the QTL allele frequencies in the two populations become more similar. The ability to detect only QTLs which differ in allele frequency between two populations may not be a great disadvantage in some circumstances, particularly when it is desired to detect favorable alleles found in one breed for introgression into a second. In fact the least squares method could be modified to detect QTLs which were segregating at intermediate frequencies in the populations which were crossed. Such QTLs would result in there being an interaction between the estimated effect of the QTL and F, family and such an interaction could be included in a least squares analysis. The detection of such an interaction, however, would require the use of F_2 families of reasonable size.

The simple method we have used here to predict probabilities of QTL genotypes does not extract all possible information from the markers. There is a potential loss of information when both an F_1 parent and its parents are heterozygous for the same alleles and thus it is not possible to simply infer from which of these grandparents an F_2 individual inherits an allele. The proportion of these noninformative heterozygous parents is not expected to be greater than 0.125 (for markers at which the same two alleles are segregating at equal frequency in the grandparental lines) and will often be much less than this. In a recent analysis of data from a cross between two outbred pig breeds (the European Wild Boar and the Large White), across 70 markers (approximately equal numbers of protein polymorphisms, RFLPs and mini or microsatellites) the average heterozygosity in the F_1 animals was 0.60 with less than 0.01 of these being non-informative heterozygotes (L. ANDERSSON, personal communication).

Some of the information lost from non-informative heterozygotes by the simple method used here to infer genotype could be retrieved by the use of maximum likelihood methods to infer parental phase if there is data available on contemporary F_2 individuals (*e.g.*, full or half-sibs). However, the extra information gained is likely to be slight, in part because non-informative heterozygotes will often be relatively rare and also because the use of all markers means that some of the information lost is retrieved from informative flanking markers.

LANDER and BOTSTEIN (1989) suggested an approximation for the predicted size of the test statistic from interval mapping with a QTL responsible for a proportion p of the total variance, midway between two markers a recombination fraction of θ apart in a sample of size N. This approximation is equivalent to:

$$[(1-2\theta)/(1-\theta)]N\log_{e}[1/(1-p)]$$

for the test statistic we use in an F_2 cross between inbred lines, and we have previously found that this provides a good prediction for the mean of replicated simulations (KNOTT and HALEY 1992a). Thus for markers fixed for alternative alleles in the grandparental lines the test statistic is expected to scale approximately with the proportion of the variance due to the QTL and Table 4 shows that this relationship appears to hold for markers which are less informative. The above formula predicts that reducing the distance between markers from 20 to 10 cM will increase the test statistic by around 12%, which is about the increase observed for the most informative markers. When the markers are less informative, increasing their density has greater effect, the increase in the test statistic for the least informative markers being around 38% for the same change in marker spacing.

For the sake of consistency with previous work, we have chosen to use the approximate log-likelihood ratio test statistic rather than the F-ratio test statistic in this study. In practice familiarity might lead to the use of the F-ratio test statistic, but for neither of these test statistics is the distribution under the null hypothesis well understood when multiple correlated tests are being performed. Thus which ever test statistic is chosen it will probably be necessary to probe the null hypothesis distribution using simulation. The limited simulations of the null hypothesis situation which we have performed bear out the results of LANDER and BOTSTEIN (1989) in that the mean highest test statistic on a chromosome and the 5% significance threshold increases with marker density. The mean highest test statistic on a chromosome also increases with marker information content but the trend in the 5% significance threshold value is not so clear cut. In practice both marker density and marker information content will vary from experiment to experiment and even between different regions of the genome. Thus it will probably be necessary to use Monte-Carlo methods to derive an approximate genome wide threshold for a chosen level of false-positives for each experiment. The least squares method lends itself to this approach because for a given set of marker data the probabilities of QTL genotypes at each position in each individual need only be derived once and stored. Then the phenotypic data can be simulated repeatedly and analysed by least squares. Nonetheless, even the modest number of 1000 replicate simulations and analyses for the 1500 1-cM spaced analyses in a 15-Morgan genome would be quite time consuming.

For slow breeding plant or animal species, especially those that suffer severely from inbreeding depression, the production of inbred lines prior to the establishment of a QTL mapping study is not an option. For other species, such as mice, it is an option, albeit a relatively slow and costly one. The advantage of inbreeding is that any markers that are useful in the cross between the inbred lines will be fixed for alternative alleles in the two lines and thus fully informative as will any segregating QTLs. However, some markers that would have been partially informative in the cross between outbred lines may be fixed for the same allele and thus become noninformative in the inbred line cross. To take an extreme example, consider markers at 10-CM intervals with two alleles at equal frequencies in each of the two outbred lines. Inbreeding these lines would result in one fully informative marker on average every 20 cM (as half of the markers are expected to be fixed for different alleles and half for the same allele). In this case our results (Table 4) suggest that producing inbred lines would result in a more powerful test for the QTL. Most markers, however, are likely to be more informative in the cross between the outbred lines than in the example given above (as drift alone is likely to have made the allele frequencies differ between the lines), and so the choice based upon power alone is not likely to be clear-cut. Often other considerations, such as time and cost, will preclude the use of inbreeding making a cross between outbred lines the only viable solution.

C.S.H. and S.A.K. acknowledge support under the Agricultural and Food Research Council (AFRC)-Institut National de la Recherche Agronomique (INRA) fellowship scheme for their stay in Toulouse. This work was also supported by the AFRC and the Ministry of Agriculture, Fisheries and Food in the United Kingdom and is part of the Pig Gene Mapping Project which is supported by the BRIDGE programme of the Commission of the European Communities.

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Communicating editor: B. S. WEIR

APPENDIX

We consider a three generation family (Table 1) with four grandparents from two outbred lines (sire of sire, SS; dam of sire, DS; sire of dam, SD and dam of dam, DD), two F_1 parents (sire, S and dam, D) and one F_2 offspring (O). The seven members of the family are all typed for codominant markers at I loci which have been already mapped. Let **P** be the vector of marker phenotypes. **P** comprises $7 \times I$ terms.

We require the posterior probability, given **P**, of the grandparental origin ω of the two alleles at any position in the genome for the offspring, prob $(\omega | \mathbf{P})$.

Notation: The vector of marker phenotypes, **P**, is subdivided into seven sub-vectors,

$$\mathbf{P} = (\mathbf{P}_{\mathbf{O}} \, \mathbf{P}_{\mathbf{S}} \, \mathbf{P}_{\mathbf{D}} \, \mathbf{P}_{\mathbf{SS}} \, \mathbf{P}_{\mathbf{DS}} \, \mathbf{P}_{\mathbf{SD}} \, \mathbf{P}_{\mathbf{DD}})$$

with $\mathbf{P}_{p} = (\mathbf{P}_{S} \mathbf{P}_{D})$ and $\mathbf{P}_{gp} = (\mathbf{P}_{SS} \mathbf{P}_{DS} \mathbf{P}_{SD} \mathbf{P}_{DD})$. At a particular locus, the genotype of an individual is defined by an ordered couplet of digits with, in the first position the allele received from the sire and in the second position the allele received from the dam. A vector of genotypes **G**, with $7 \times I$ couplets, underlies the vector **P**. Hence for an observation **P** with a total (over all marker loci in all seven individuals in the pedigree) of *H* heterozygous loci there are 2^{H} corresponding different possible vectors **G**.

Let Ω be a vector of grandparental origins of the *I* marker loci in the offspring, $\Omega = (\Omega_1 \ \Omega_2 \ \dots \ \Omega_I)$. $\Omega_i \ (i = 1 \ \dots \ I)$ represents the line origin combination of the marker alleles and takes the value 11 when the offspring received the *SS* and *SD* alleles, 12 when *SS* and *DD*, 21 when *DS* and *SD* and 22 when *DS* and *DD*.

Result: If the markers are codominant and without missing data, assuming linkage equilibrium in the grandparents between the loci and no interference in recombination, the probability of the grandparental origin at any position in the offspring genome, given the phenotypes **P** is:

 $\operatorname{prob}(\omega | \mathbf{P}) = \operatorname{prob}(\omega | \Omega_i) \operatorname{prob}(\Omega_{i+1} | \omega)$

$$\times \frac{\prod_{i=2}^{j} \operatorname{prob}(\Omega_{i} | \Omega_{i-1}) \prod_{i=j+2}^{I} \operatorname{prob}(\Omega_{i} | \Omega_{i-1})}{\sum_{\Omega \in \Gamma \mathbf{p}} \prod_{i=2}^{I} \operatorname{prob}(\Omega_{i} | \Omega_{i-1})}$$

with

and

$$\prod_{i=2}^{j} \operatorname{prob}(\Omega_{i} | \Omega_{i-1}) = 1 \quad \text{when} \quad j = 1$$

$$\prod_{i=j+2}^{I} \operatorname{prob}(\Omega_i | \Omega_{i-1}) = 1 \quad \text{when} \quad j = I - 1$$

 $\Gamma_{\mathbf{P}}$ being the whole set of consistent vectors Ω considering the observations \mathbf{P} and markers j $(1 \le j \le I - 1)$ and j+1 bound the interval containing the putative QTL.

This is equivalent to:

 $\operatorname{prob}(\omega | \mathbf{P})$

$$= \sum_{\Omega \in \Gamma_{\mathbf{p}}} \operatorname{prob}(\omega \mid \Omega_{j}, \Omega_{j+1}) \frac{\prod_{i=2}^{l} \operatorname{prob}(\Omega_{i} \mid \Omega_{i-1})}{\sum_{\Omega \in \Gamma_{\mathbf{p}}} \prod_{i=2}^{l} \operatorname{prob}(\Omega_{i} \mid \Omega_{i-1})}$$

That is, the required probability can be written in terms requiring only the recombination rate between adjacent markers $(\operatorname{prob}(\Omega_i | \Omega_{i-1}))$ as given in Table 2) and between the putative QTL and its flanking markers.

Proof:

$$\operatorname{prob}(\omega | \mathbf{P}) = \sum_{\Omega} \operatorname{prob}(\omega | \Omega \mathbf{P}) \operatorname{prob}(\Omega | \mathbf{P}).$$

We will now consider the two components separately. First,

$$\operatorname{prob}(\omega \mid \Omega \mathbf{P}) = \operatorname{prob}(\omega \mid \Omega).$$

In the absence of interference in recombination, we have

$$\operatorname{prob}(\omega \mid \Omega) = \operatorname{prob}(\omega \mid \Omega_i, \Omega_{i+1})$$
(1)

where j and j + 1 are the markers flanking the considered position. Second,

 $\operatorname{prob}(\Omega | \mathbf{P})$

Rewriting in terms that are conditional on only parental and grandparental phenotypes gives:

$$\operatorname{prob}(\Omega | \mathbf{P}) = \frac{\operatorname{prob}(\Omega | \mathbf{P}_{gp}, \mathbf{P}_{p})\operatorname{prob}(\mathbf{P}_{O} | \Omega, \mathbf{P}_{gp}, \mathbf{P}_{p})}{\sum_{\Omega} \operatorname{prob}(\Omega | \mathbf{P}_{gp}, \mathbf{P}_{p})\operatorname{prob}(\mathbf{P}_{O} | \Omega, \mathbf{P}_{gp}, \mathbf{P}_{p})}$$

The two component probabilities will now be considered separately. First,

$$prob(\Omega | \mathbf{P}_{gp}, \mathbf{P}_{p})$$

= prob(\Omega)
= prob(\Omega_{1})prob(\Omega_{2} | \Omega_{1})prob(\Omega_{3} | \Omega_{1}, \Omega_{2}) \dots ...

In the absence of interference, we have $\operatorname{prob}(\Omega_i | \Omega_1,$

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 $\Omega_2 \ldots \Omega_{i-1}$ = prob $(\Omega_i | \Omega_{i-1})$. Thus

$$\operatorname{prob}(\Omega \mid \mathbf{P}_{gp}, \mathbf{P}_{p}) = \operatorname{prob}(\Omega_{1}) \prod_{i=2}^{I} \operatorname{prob}(\Omega_{i} \mid \Omega_{i-1}). \quad (2)$$

The first term, $\operatorname{prob}(\Omega_1)$, is simply $\frac{1}{4}$. Second,

$$\operatorname{prob}(\mathbf{P_{O}} \mid \Omega, \mathbf{P_{gp}}, \mathbf{P_{p}})$$

Rewriting this probability to be conditional on the parental genotypes gives:

$$\begin{aligned} \operatorname{prob}(\mathbf{P}_{\mathbf{O}} \mid \Omega, \, \mathbf{P}_{\mathsf{gp}}, \, \mathbf{P}_{\mathsf{p}}) \\ &= \sum_{\mathbf{G}_{\mathsf{p}}} \operatorname{prob}(\mathbf{P}_{\mathbf{O}} \mid \Omega, \, \mathbf{P}_{\mathsf{gp}}, \, \mathbf{P}_{\mathsf{p}}, \, \mathbf{G}_{\mathsf{p}}) \operatorname{prob}(\mathbf{G}_{\mathsf{p}} \mid \Omega, \, \mathbf{P}_{\mathsf{gp}}, \, \mathbf{P}_{\mathsf{p}}). \end{aligned}$$

Again, we will consider the two components separately. First,

$$prob(\mathbf{P_{o}} \mid \Omega, \mathbf{P_{gp}}, \mathbf{P_{p}}, \mathbf{G_{p}}) = prob(\mathbf{P_{o}} \mid \Omega, \mathbf{G_{p}})$$

$$= \prod_{i=1}^{I} prob(P_{Oi} \mid \Omega_{i}, \mathbf{G_{pi}}).$$
(3)

Due to the one-to-one correspondence between $(\Omega_{\nu} \mathbf{G}_{\mu})$ and G_{0i} , we have $\operatorname{prob}(P_{0i} | \Omega_{\nu} \mathbf{G}_{\mu}) = \operatorname{prob}(P_{0i} | G_{0i})$ which is 1 if phenotype and genotype are consistent and 0 otherwise. Second,

$$\begin{aligned} & \operatorname{prob}(\mathbf{G}_{p} \mid \Omega, \mathbf{P}_{gp}, \mathbf{P}_{p}) \\ &= \operatorname{prob}(\mathbf{G}_{p} \mid \mathbf{P}_{gp}, \mathbf{P}_{p}) \\ &= \operatorname{prob}(\mathbf{G}_{s} \mid \mathbf{P}_{ss}, \mathbf{P}_{Ds}, \mathbf{P}_{s}) \operatorname{prob}(\mathbf{G}_{D} \mid \mathbf{P}_{sD}, \mathbf{P}_{DD}, \mathbf{P}_{D}). \end{aligned}$$

Consider the sire probability. Assuming linkage equilibrium in the grandparents and with a single progeny (S), the events at each locus are independent, hence

$$\operatorname{prob}(\mathbf{G}_{\mathbf{S}} | \mathbf{P}_{\mathbf{SS}}, \mathbf{P}_{\mathbf{DS}}, \mathbf{P}_{\mathbf{S}}) = \prod_{i=1}^{I} \operatorname{prob}(G_{Si} | P_{SSi}, P_{DSi}, P_{Si}).$$
(4)

Considering the *i*th locus, three situations are possible, *viz*:

(S1) G_{Si} and P_{Si} are not consistent, giving prob $(G_{Si}|P_{SSi}, P_{DSi}, P_{Si}) = 0.$

(S2) G_{Si} and P_{Si} are consistent; P_{SSi} , P_{DSi} , P_{Si} are not all heterozygous for the same alleles. The probability is 1.

(S3) G_{Si} and P_{Si} are consistent; $P_{SSi} = P_{DSi} = P_{Si}$ are all heterozygous (say AB).

$$prob(G_{Si} = AB | P_{SSi} = P_{DSi} = P_{Si} = AB)$$
$$= prob(G_{Si} = BA | P_{SSi} = P_{DSi} = P_{Si} = AB) = \frac{1}{2}$$

Hence, combining the results from (3) and (4), we have

$$prob(\mathbf{P_{O}} \mid \Omega, \mathbf{P_{gp}}, \mathbf{P_{p}})$$

$$= \sum_{\mathbf{G}_{p}} \prod_{i} (prob(P_{Oi} \mid \Omega_{i}, \mathbf{G_{pi}}) prob(\mathbf{G_{pi}} \mid \mathbf{P_{gpi}}, \mathbf{P_{pi}}))$$

$$= \prod_{i} \left(\sum_{\mathbf{G_{pi}}} prob(P_{Oi} \mid \Omega_{i}, \mathbf{G_{pi}}) prob(\mathbf{G_{pi}} \mid \mathbf{P_{gpi}}, \mathbf{P_{pi}}) \right).$$

For any consistent Ω_i and \mathbf{G}_{pi} , the product $(\operatorname{prob}(P_{Oi}|\Omega_i, \mathbf{G}_{pi})\operatorname{prob}(\mathbf{G}_{pi}|\mathbf{P}_{gpi}, \mathbf{P}_{pi}))$ is a constant with a value of $\frac{1}{4}$, $\frac{1}{2}$ or 1, depending on the number of grandparent and parent trios that are heterozygous for the same genotype (i.e., 2, 1 or 0).

Additionally, there is only one possible G_{pi} in the summation. This is obvious in situations S1 and S2, above. In the situation S3, two G_{Si} are consistent with P_{S} , P_{SS} and P_{DS} but one only belongs to Γ_{Ω} . If, for example, $\Omega_i = 11$ and $G_{Di} = XY$, the genotype $G_{Si} = AB$ gives an offspring phenotype AX and the genotype $G_{Si} = BA$ gives an offspring genotype BX. If the observed phenotype P_{Oi} was AA, $G_{Si} = BA$ cannot be consistent; if it is AB, allele X must be A or B and only one of the G_{Si} (depending on the G_{Di}) is possible.

Hence $\operatorname{prob}(\mathbf{P}_0 | \Omega, \mathbf{P}_{gp}, \mathbf{P}_p)$ is constant over all line origin combinations (Ω) consistent with \mathbf{P} and it is equal to $(\frac{1}{2})^h$, h being the number of grandparental and parental trios that are heterozygous for the same genotype over all loci, and for all other combinations is 0.

Combining this result with (2) we have the following expression for any possible Ω ,

$$\operatorname{prob}(\Omega | \mathbf{P}) = \frac{\prod_{i=2}^{l} \operatorname{prob}(\Omega_{i} | \Omega_{i-1}) (\frac{1}{2})^{h}}{\sum_{\Omega \in \Gamma_{\mathbf{P}}} \prod_{i=2}^{l} \operatorname{prob}(\Omega_{i} | \Omega_{i-1}) (\frac{1}{2})^{h}}$$

and, incorporating the result from (1), the probability we require can be written as follows:

 $prob(\boldsymbol{\omega} | \mathbf{P})$

$$= \sum_{\Omega \in \Gamma_{\mathbf{p}}} \operatorname{prob}(\omega \mid \Omega_{j}, \Omega_{j+1}) \frac{\prod_{i=2}^{l} \operatorname{prob}(\Omega_{i} \mid \Omega_{i-1})}{\sum_{\Omega \in \Gamma_{\mathbf{p}}} \prod_{i=2}^{l} \operatorname{prob}(\Omega_{i} \mid \Omega_{i-1})}$$

This proof holds for all situations where the alleles in the parents and grandparents are known. So marker phenotypes may be missing in the offspring or dominant loci can be used if the parental and grandparental alleles are known. The assumption of linkage equilibrium is required only if there are loci at which both grandparents and a parent are heterozygous for the same alleles (*i.e.*, uninformative heterozygous parents). Note also that if grandparents or parents have more than one offspring these could be used to infer phase in uninformative heterozygous parents and the formula we give is then an approximation (in the sense that it does not use all available information).