3-24-2006

POOR PERFORMANCE OF BOOTSTRAP CONFIDENCE INTERVALS FOR THE LOCATION OF A QUANTITATIVE TRAIT LOUCS

アニ・マニチャクル
Department of Biostatistics, Johns Hopkins Bloomberg School of Public Health

ヨシエ・デプイユ
Department of Biostatistics, Boston University School of Public Health

サナック・セン
Department of Epidemiology and Biostatistics, University of California

カルル・W・ブロマン
Department of Biostatistics, The Johns Hopkins Bloomberg School of Public Health, kbroman@biostat.wisc.edu

Suggested Citation
http://biostats.bepress.com/jhubiostat/paper105

This working paper is hosted by The Berkeley Electronic Press (bepress) and may not be commercially reproduced without the permission of the copyright holder.
Copyright © 2011 by the authors
Poor performance of bootstrap confidence intervals for the location of a quantitative trait locus

Ani Manichaikul*, Josée Dupuis†, Šaunak Sen‡, Karl W. Broman*,1

*Department of Biostatistics, Johns Hopkins University, Baltimore, Maryland 21205
†Department of Biostatistics, Boston University School of Public Health, Boston, Massachusetts 02118
‡Department of Epidemiology and Biostatistics, University of California, San Francisco, California 94107

24 March 2006
Running head: QTL confidence intervals

Key words: QTL, LOD support intervals, confidence intervals, bootstrap, Bayes credible intervals

1Corresponding author:

Karl W Broman
Department of Biostatistics
Johns Hopkins University
615 North Wolfe Street
Baltimore MD 21205–2179
USA

Phone: 410–614–9408
Fax: 410–955–0958
Email: kbroman@jhsph.edu
ABSTRACT

The aim of many genetic studies is to locate the genomic regions (called quantitative trait loci, QTLs) that contribute to variation in a quantitative trait (such as body weight). Confidence intervals for the locations of QTLs are particularly important for the design of further experiments to identify the gene or genes responsible for the effect. Likelihood support intervals are the most widely used method to obtain confidence intervals for QTL location, but the non-parametric bootstrap has also been recommended. Through extensive computer simulation, we show that bootstrap confidence intervals are poorly behaved and so should not be used in this context. The profile likelihood (or LOD curve) for QTL location has a tendency to peak at genetic markers, and so the distribution of the maximum likelihood estimate (MLE) of QTL location has the unusual feature of point masses at genetic markers; this contributes to the poor behavior of the bootstrap. Likelihood support intervals and approximate Bayes credible intervals, on the other hand, are shown to behave appropriately.
INTRODUCTION

There is much interest in mapping the genetic loci (called quantitative trait loci, QTLs) that contribute to variation in a quantitative trait. Once such a QTL has been identified, interest turns to the calculation of a confidence interval for its location, as such an interval estimate can be a useful guide for the design of further experiments, such as the generation of congenic lines.

LOD support intervals are the most commonly used interval estimates for the location of a QTL. A LOD support interval is defined as the interval in which the LOD score is within some value of its maximum. As an illustration, Figure 1A displays the LOD curve for chromosome 4 for the data of SUGIYAMA et al. (2001), concerning salt-induced hypertension in 250 backcross mice. Assuming that there is a single QTL on this chromosome, the maximum likelihood estimate (MLE) of the location of the QTL is the position at which the LOD curve achieves its maximum, in this case at marker D4Mit164 (at 30 cM). The 1.5-LOD support interval for the location of the QTL is the region in which the LOD score is within 1.5 of its maximum; here, the interval extends from 19 to 31 cM. (When the relevant region is disconnected, we generally take the conservative approach of forming the longest contiguous interval.)

LANDER and BOTSTEIN (1989) recommended the use of 1- and 2-LOD support intervals. DUPUIS and SIEGMUND (1999) found that 1.5-LOD support intervals provide approximately 95% coverage in the case of a dense marker map. However, it has often been observed (see, e.g., MANGIN et al. 1994), that the coverage of LOD support intervals depends upon the effect of the QTL, and so they do not behave as true confidence intervals.

VISSCHER et al. (1996) recommended the use of a nonparametric bootstrap to derive a
confidence interval for the location of a QTL. For experimental cross data on \( n \) individuals, one makes \( n \) draws, with replacement, from the observed individuals to form a new data set in which some individuals are omitted and some appear multiple times. An estimate of QTL location is calculated with these new data, and the process is repeated many times. An approximate 95% confidence interval for the location of the QTL is obtained as the interval containing 95% of the estimated locations from the bootstrap replicates.

As an illustration, Figure 1B contains a histogram of the results of 10,000 bootstrap replicates using the chromosome 4 data of Sugiyama et al. (2001). The 95% bootstrap confidence interval extends from 14 to 32 cM. A striking feature of these results is that approximately 79% of the bootstrap replicates gave an estimated QTL location precisely at one of the 20 genetic markers on the chromosome. (Note that the calculations were performed at the markers and at 1 cM steps along the chromosome.) This is due to an unusual feature of the MLE of QTL location (previously observed by Walling et al. 1998): it has a great tendency to occur precisely at a marker.

Walling et al. (1998, 2002) investigated the performance of bootstrap confidence intervals for QTL location and concluded that they provide appropriate coverage. However, the unusual character of the distributions obtained in applications of the bootstrap for this problem, well illustrated in Figure 1B, led us to suspect that the performance of the bootstrap may be less than ideal, and that the bootstrap may be inappropriate for the construction of confidence intervals for QTL location. Thus, we conducted a large-scale computer simulation study to investigate the performance of bootstrap confidence intervals for QTL location.

We considered the case of a backcross with a single segregating QTL, normally distributed residual variation, and equally spaced genetic markers exhibiting complete genotype data.
While our simulation study is similar to those of WALLING et al. (1998, 2002), and differs largely in scale and thus precision, our conclusions are quite different. We find that the coverage of bootstrap confidence intervals for QTL location shows great variation as a function of the location of the QTL relative to the available genetic markers, and so we recommend against the use of the bootstrap for this problem.

One cannot reasonably recommend against the use of a method without providing some alternative, and so we further investigated the performance of LOD support intervals, as well as an approximate Bayes credible interval initially suggested by SEN and CHURCHILL (2001). Both of these types of intervals were found to display relatively stable coverage. On the basis of extensive simulations of backcrosses and intercrosses with varying marker densities and varying sizes of the effect of the QTL, we provide estimates of the appropriate amount to drop for LOD support intervals and the appropriate nominal fraction for the Bayes credible intervals in order to attain an actual coverage of 95%. The Bayes credible intervals are particularly attractive, as a nominal Bayes fraction of 96.5% in a backcross (and 97% in an intercross) is found to provide quite consistent coverage, irrespective of the size of the QTL effect, marker density, number of individuals.
METHODS

We consider the case of a backcross or intercross with a single segregating QTL. We focus on the single chromosome (taken to have length 100 cM) harboring the QTL, and assume equally-spaced markers with complete genotype data. The residual variation is assumed to follow a normal distribution, and QTL mapping was performed by standard interval mapping (LANDER and BOTSTEIN 1989), which we briefly describe.

One assumes the presence of a single QTL, and considers each position on the chromosome, one at a time, as the putative location of the QTL. (Our analyses were conducted at 1 cM steps along the chromosome.) While the QTL genotype, \( q \), of an individual is generally not known, its distribution, conditional on the available marker data, may be calculated. Under the assumption of no crossover interference and with complete marker genotype data, the distribution of \( q \) depends only on the genotypes at the flanking markers. Given the QTL genotype, the phenotype is assumed to follow a normal distribution with mean \( \mu_q \) and common standard deviation \( \sigma \). Given the available marker data, the phenotype follows a mixture of these normal distributions with known mixing proportions (the QTL genotype probabilities, conditional on the marker data). The nuisance parameters (the \( \mu_q \) and \( \sigma \)) are estimated by maximum likelihood via the EM algorithm (DEMPSTER et al. 1977), and a LOD score is calculated, comparing the hypothesis that there is a single QTL precisely at that location to the null hypothesis of no QTL anywhere (in which case the phenotypes are assumed to follow a single normal distribution, independent of genotype).

Let \( \theta \) denote the true location of the QTL. The result of interval mapping is a LOD curve, \( \text{LOD}(\theta) \), for the position of the QTL along the chromosome, \( \theta \). This LOD curve is equivalent to a profile log likelihood for the position of the QTL. The MLE of the location of the QTL, \( \hat{\theta} \),
is the position at which the LOD curve achieves its maximum. While analysis at one 1 cM steps along the chromosome results in a discrete distribution for $\hat{\theta}$, analysis on a finer grid would greatly increase the computational effort, and would provide similar results.

LOD support intervals were calculated as the longest contiguous interval in which the LOD score was within some chosen value of its maximum. Bootstrap confidence intervals were constructed via the percentile method, as described by VISSCHER et al. (1996). For each of 1000 bootstrap replicates, a sample of the same size as the available data were drawn with replacement from the available individuals, a new estimate of QTL location ($\hat{\theta}^*$) was obtained by application of standard interval mapping to the resampled data. The endpoints of the 95% bootstrap confidence interval were taken to be the 2.5 and 97.5 percentiles of the $\hat{\theta}^*$. Finally, an approximate Bayes credible interval was calculated: we treated the profile likelihood for QTL location as if it were a real likelihood, assigned a uniform prior on the location of the QTL, and so derived an approximate posterior distribution for QTL location,

$$f(\theta \mid \text{data}) = 10^{\text{LOD}(\theta)} / \sum_{\theta=0}^{100} 10^{\text{LOD}(\theta)}.$$  From this approximate posterior, a 95% Bayes credible interval was defined to be the interval, $I$, for which $f(\theta \mid \text{data})$ exceeded some threshold and for which $\sum_{\theta \in I} f(\theta \mid \text{data}) \geq 0.95$.

Effect of QTL location relative to markers: In our first simulation study, to investigate the coverage of bootstrap confidence intervals, we considered a backcross of 200 individuals and a single QTL whose position was allowed to vary at 1 cM steps along a chromosome of length 100 cM. Complete genotype data were available at 11 equally spaced markers (thus at a 10 cM spacing). The heritability due to the QTL (the proportion of the phenotypic variance due to the QTL) was taken to be 10%.

For each of the 101 possible QTL positions (at 0, 1, 2, ..., 100 cM), we performed 10,000
simulation replicates. At each replicate, we calculated the LOD curve by standard interval mapping at 1 cM steps along the chromosome and derived the 1-LOD support interval and approximate 95% Bayes credible interval. (We used 1-LOD support intervals here, as they were found to be somewhat conservative in this sparse-map case.) In addition, at each simulation replicate we constructed a 95% bootstrap confidence interval on the basis of 1000 bootstrap replicates, as described above. Great computational effort was expended in this investigation: at each of 1,010,000 simulation replicates (10,000 replicates for each of 101 QTL positions), 1000 bootstrap replicates were performed.

The simulations were performed using the R statistical software (Ihaka and Gentleman 1996) and R/qtl (Broman et al. 2003), an add-on package to R. For some aspects of our simulation studies, we used C code adapted from the R code in R/qtl in order to improve computational speed.

Effect of cross type, sample size, marker density and QTL effect: Based on the results of the first simulation study, we performed a second simulation study to more completely characterize the coverage of the LOD support and Bayes credible intervals. We varied the type of cross (backcross or intercross) the sample size (200 or 500), the marker density (1, 2, 10 or 20 cM spacing), and the effect of the QTL. We hypothesized that interval coverage might be more clearly expressed as a function of the power to detect the QTL rather than the heritability due to the QTL, and so heritabilities were chosen to give estimated power of 0.3, 0.4, . . . , 0.9, where power was defined as the probability of achieving a LOD score of at least 3. These heritabilities were estimated via R/qtlDesign (Sen et al. 2005), an add-on package to the R statistical software (Ihaka and Gentleman 1996). Our targeted values for power, calculated with R/qtlDesign, differed somewhat from the power estimated from our simulation results, as
we defined power to be the chance of a LOD score $\geq 3$ somewhere on the chromosome, whereas R/qtlDesign defines it to be the chance of a LOD score $\geq 3$ at the QTL. The results will be presented below using the power seen in our simulations.

The position of the QTL was fixed at a position equidistant between two markers and near the center of the chromosome. For the marker spacings of 1, 2, 10, and 20 cM, the QTL was placed at 50, 49, 45 and 50 cM, respectively. For each setting (of cross type, sample size, marker density, and QTL effect), 100,000 simulation replicates were performed.

For each simulation replicate, standard interval mapping was performed to obtain the LOD curve at 1 cM steps along the chromosome. Rather than investigate the coverage of the LOD support and Bayes credible intervals for particular choices of the drop in LOD and the nominal Bayes fraction, we chose to estimate the drop in LOD and the nominal Bayes fraction for which the two types of intervals would attain 95% coverage. These values could be obtained with little additional effort. At each simulation replicate, we kept track of the difference in the LOD score at the MLE and at the true location of the QTL. The 95th percentile of these differences is the value to drop in a LOD support interval in order to attain 95% coverage. A similar trick applies for the Bayes credible intervals. Note that here we are using a definition for the confidence interval that can lead to a set of disjoint intervals, rather than a single contiguous interval, and so the results are somewhat conservative.
RESULTS

**Distribution of the MLE of QTL location:** Our initial simulation study, comprising 10,000 replicates with a QTL at each of 0, 1, 2, . . . , 100 cM on a chromosome of length 100 cM, allows us to inspect the distribution of the MLE of QTL location and the dependence of this distribution on the location of the QTL relative to the markers. The simulations used a backcross of 200 individuals, 11 equally spaced markers (10 cM spacing), and heritability due to the single QTL at 10%.

Figure 2 displays the distribution of the MLE of QTL position, \( \hat{\theta} \), as a function of the true location of the QTL, \( \theta \), for \( \theta = 45, 46, \ldots , 50 \). The most striking feature of these distributions is the clear tendency for the \( \hat{\theta} \) to occur exactly at the marker loci. For example, in the case that the QTL is at 49 cM, immediately adjacent to a marker, there is a far greater chance that the QTL is estimated to be at the marker rather than at the true location of the QTL. A similar pattern was seen for other values of \( \theta \). The standard error (SE) of \( \hat{\theta} \) is smallest when the QTL is at a marker, and is \( \sim 25\% \) larger when the QTL is in the center of the interval between markers. When the QTL is near one of the ends of the chromosome, \( \hat{\theta} \) exhibits considerable bias, as we do not examine positions beyond the terminal markers on the chromosome. We calculated the LOD score at 1 cM steps along the chromosome, and so estimated QTL position only to within 1 cM. If calculations were performed on a more dense grid, the tendency for the MLE to occur precisely at the markers would be even more striking.

The dependence of the distribution of \( \hat{\theta} \) on the position of the QTL relative to the markers, both with respect to the large mass placed at the markers and the variation in the SE of \( \hat{\theta} \), will be seen to cause a breakdown in the performance of the bootstrap for this problem.

Coverage vs. true QTL location: Performance of a confidence interval is generally
assessed by its coverage (the probability that it contains the true parameter value) as a function of the true parameter. Ideally, a 95% confidence interval shows constant 95% coverage, regardless of the true parameter value. In Figure 3, coverage of the 95% bootstrap confidence interval (in black), 1-LOD support interval (in red) and the approximate 95% Bayes credible interval (in blue) are displayed as a function of the true location of the QTL. The bootstrap confidence intervals shows extremely high coverage (∼99%) when the true QTL is at a marker, low coverage (∼92.5%) when the QTL is right next to a marker, and above nominal coverage when the QTL is exactly between markers. Coverage of the 1-LOD support and approximate 95% Bayes credible intervals does not fluctuate as widely, though it is highest when the QTL is at a marker. Note that the SEs of our estimates of coverage are ∼0.3%.

Coverage vs. estimated QTL location: It is also of interest to consider coverage as a function of the estimated QTL location, \( \hat{\theta} \). In our simulations, we performed 10,000 replicates for each of the 101 possible positions of the QTL; here we consider the portion of those 1,010,000 simulation replicates in which the MLE was attained, for example, at 50 cM, and calculate the proportion of those replicates in which each type of confidence interval contained the true parameter value. This is an unorthodox mixture of Bayes and frequentist statistics. The coverage of a confidence interval is a quantity of interest only to frequentists; here we are taking the location of the QTL to be uniformly distributed on the positions 0, 1, 2, . . . , 100 cM, and inspecting the posterior probability, given the observed estimate of the QTL location, that the interval covers its true parameter value. Note that, across the 1,010,000 simulation replicates, each possible value of \( \hat{\theta} \) was observed at least 6700 times, and so the SEs of our estimates of coverage as a function of \( \hat{\theta} \) are ∼0.4%.

Coverage as a function of the estimated location of the QTL is displayed in Figure 4. These
results provide a qualitatively different perspective from those of coverage vs. $\theta$, shown in Figure 3. While coverage of the bootstrap confidence intervals is high when the QTL is at a marker (see Figure 3), coverage is low (~92%) when the QTL is estimated to be at a marker. Coverage of the 1-LOD support interval and approximate 95% Bayes credible interval is less variable as a function of $\hat{\theta}$, and is entirely above the nominal level, 95%, with the Bayes credible interval exhibiting slightly less variability than the LOD support interval.

We view this perspective (coverage of a confidence interval conditional on the observed estimate, $\hat{\theta}$) as the more relevant one for the user of a confidence interval. One does not know the true location of the QTL, but does know one’s estimate of that location, and so coverage as a function of the observed estimate is of greatest interest. But it is from this perspective that coverage of the bootstrap confidence intervals looks worst. While coverage is low only when the estimated location of the QTL is at a marker, it is quite low in that case, and, as we’ve seen, that is often the case.

**Interval widths:** Another important feature of a confidence interval is its width: one prefers intervals to be as small as possible, while maintaining the appropriate level of coverage. Averaging over all possible values of $\theta$, the 95% bootstrap confidence interval, 1-LOD support intervals and 95% Bayes credible intervals had average widths of 45, 24, and 29 cM, respectively. When the QTL was not close to the end of the chromosome, the 1-LOD support intervals were more than 40% smaller than the bootstrap confidence intervals about half of the time. The approximate 95% Bayes credible intervals and the 1-LOD support intervals were quite similar in width. The 1-LOD support and 95% Bayes credible intervals show not just better coverage probabilities than the 95% bootstrap intervals (see Figures 3 and 4), but are also generally smaller.
Coverage with varying cross type, sample size, marker density and QTL effect: As described above, coverage of the 95% bootstrap confidence intervals varied greatly according to the position of the QTL relative to the genetic markers; the 1-LOD support and 95% Bayes credible intervals, on the other hand, exhibited relatively stable coverage across the chromosome. We thus omitted the bootstrap confidence intervals from further consideration, but sought a more complete characterization of the performance of the LOD support intervals and the approximate Bayes credible intervals, as a function of sample size, marker density, and QTL effect, and considering both a backcross and an intercross.

Rather than study the coverage of the intervals for a fixed drop in LOD or nominal Bayes fraction, we sought the values that would give 95% coverage at different settings of the parameters of interest. The value to drop in LOD in order for the coverage of the LOD support interval to have 95% coverage in a backcross is displayed in Figure 5A. The results are displayed as a function of the size of the effect of the QTL, which has been reparameterized as the power to give a LOD score of at least 3. (The displayed values for the power were estimated from 100,000 simulation replicates at each point, and so have standard error <0.2%.) The black and red curves correspond to sample sizes of 200 and 500, respectively. As seen in the figure, sample size has little effect on the appropriate drop in LOD to give 95% coverage, for a given power to detect the QTL. Of course, the heritability due to the QTL that corresponds to a particular power is quite different for the two sample sizes. The biggest effect seen concerns the spacing of markers: one must drop $\sim 1.5$ in LOD to attain 95% coverage when markers are at a 1 cM spacing, but need drop only $\sim 1.2$ in LOD if the markers are at a 10 cM spacing. A slightly smaller drop is required when the QTL has a larger effect.

The nominal Bayes fractions at which the approximate Bayes credible intervals had 95%
coverage in a backcross are displayed in Figure 5B. Again, sample size has little effect, except in the case of very dense markers. The effect of marker spacing and of the size of the QTL effect is seen to be in the opposite direction for the Bayes intervals versus the LOD support intervals. A greater nominal Bayes fraction is needed for sparse markers and for a larger QTL effect.

Figures 5C and 5D show the corresponding results for an intercross. A greater drop in LOD is required in order for the LOD support interval to have 95% coverage in the intercross, and the QTL effect appears to have a somewhat greater influence on the appropriate value to drop. Sample size is again seen to have little effect, and the greatest influence comes from the marker spacing, with a greater drop in LOD required in the case of more densely spaced markers. There is remarkably little variation in the appropriate nominal Bayes fraction so that the approximate Bayes credible interval has 95% coverage in an intercross; for all sample sizes, marker spacings, and QTL effects, the appropriate nominal Bayes fraction was 96–97%.

These results suggest that, for the Bayes intervals, the use of 96.5% for a backcross and 97% for an intercross will provide greater than 95% coverage for all possible cases. For the LOD support intervals, if one drops by 1.5 for a backcross and 1.8 for an intercross, coverage will be maintained at greater than 95%. The actual coverage obtained with these choices are shown in Figure 6. The Bayes intervals are seen to be particularly attractive, as they exhibit quite stable coverage with sample size, marker density, and QTL effect.
DISCUSSION

We have shown that coverage of bootstrap confidence intervals for QTL location depends critically upon the location of the QTL relative to the typed genetic markers. Coverage is high when the QTL is at a marker but can be low when the QTL is immediately adjacent to a marker (see Figure 3). Especially interesting results were observed in the consideration of coverage as a function of the estimated location of the QTL, taking the true location of the QTL to be uniformly distributed along the chromosome. This perspective is most relevant for the user of such confidence intervals, and indicates poor performance of the bootstrap confidence intervals: coverage is quite far below the nominal level when the QTL is estimated to be at a marker (see Figure 4). The bootstrap confidence intervals were also seen to be much wider than the LOD support and approximate Bayes credible intervals.

Our results are similar to those of WALLING et al. (1998, 2002), but our conclusions are markedly different. It is important to point out that WALLING et al. (1998, 2002) used Haley-Knott regression (HALEY and KNOTT 1992), whereas we have focused on standard interval mapping (LANDER and BOTSTEIN 1989), using maximum likelihood via the EM algorithm. While it was not mentioned above, we did include the use of Haley-Knott regression in our initial simulation study, and found similar results by the two methods (data not shown).

Bootstrap methods have desirable properties in a wide variety of statistical problems. However, modifications to the bootstrap are necessary for problems that are not classically regular (BERAN 2003), and the QTL mapping problem is not regular (KONG and WRIGHT 1994, SIEGMUND 2004). Thus, our finding of inadequate bootstrap performance in QTL mapping is consistent with theory.

The poor performance of bootstrap confidence intervals for QTL location derives from the
unusual behavior of the MLE of QTL location: the MLE has a tendency to coincide with a marker position (see Figure 2 and Kong and Wright 1994), and its SE varies greatly according to the location of the QTL relative to the markers. Appropriate performance of the percentile-based nonparametric bootstrap confidence intervals (proposed, for this context, by Viisscher et al. 1996, and studied herein) generally requires the existence of some monotone transformation \( h(\cdot) \) such that \( h(\hat{\theta}) - h(\theta) \) has the same symmetric continuous distribution for all \( \theta \) (Shao and Tu 1995, pg 132). The tendency of the MLE to occur at a marker indicates that no such transformation exists for this problem.

An alternative heuristic for understanding the breakdown of the bootstrap in this problem is as follows: we hope to approximate the sampling distribution, \( f(\hat{\theta} \mid \theta) \), by the bootstrap distribution, \( g(\hat{\theta}^* \mid \hat{\theta}) \). But the bootstrap distribution better reflects the sampling distribution evaluated at the observed estimate, \( f(\cdot \mid \hat{\theta}) \), than it does the target, \( f(\cdot \mid \theta) \). That the MLE is most precise when the QTL is at a marker, and is less precise when the QTL is between markers, indicates that the bootstrap distribution will provide an overly optimistic view of our understanding of the location of the QTL in those cases in which we have estimated the QTL to be at a marker.

Walling et al. (1998) also assessed the performance of the parametric bootstrap for this problem; rather than resampling from the observed data, one simulates new data taking one’s estimate of the QTL location to be the true location. They obtained the surprising result that the parametric bootstrap performed more poorly than the nonparametric bootstrap; the result is surprising, because when one’s model is correct (as it was in their simulation study), the parametric bootstrap would be expected to give better performance than the nonparametric bootstrap. This result can now be clearly understood. In the parametric bootstrap, the bootstrap
distribution on which the confidence interval is based is simply the sampling distribution of the estimate in the case that the QTL is located at the observed estimate. Thus, when the QTL is estimated to be at a marker, the parametric bootstrap will provide an overly optimistic view of the precision of that estimate.

The tendency of the MLE for QTL location to occur precisely at a genetic marker (see Figure 2) is a major contributor to the failure of the bootstrap in this context. Our explanation of the cause of this behavior is as follows. The profile likelihood exhibits cusps at the markers. (Its first derivative is not continuous at the markers.) This is the result of the fact that, in the case of complete genotype data at the markers, and with the assumption of no crossover interference, the likelihood to the left of the marker incorporates data on the marker to the left but not that for the marker to the right, while the likelihood to the right of a marker incorporates data on the marker to the right but not that for the marker to the left. The abrupt change in the first derivative of the profile likelihood at the markers appears to lead to a greater chance of a change in the direction of the profile likelihood, and so a greater chance that the MLE occurs precisely at a marker.

It should be emphasized that these results were obtained in a single setting: a backcross of 200 individuals, equally spaced markers at a 10 cM spacing, and heritability due to the QTL at 10%. The behavior of the bootstrap seen here may not hold generally. In fact, for a cross with very dense markers and a QTL of not too strong effect, the bootstrap would likely behave reasonably. However, the setting in which our simulations were conducted is not unreasonable, and that the bootstrap performed so poorly here supports the general conclusion that it should not be used.

It should also be emphasized that we have considered only percentile-based nonparametric
bootstrap confidence intervals, as that was the approach recommended by Visscher et al. (1996). Other forms of bootstrap might be found to work in this context. For example one might use a bootstrap to calibrate the LOD support or approximate Bayes credible intervals. However, the good performance of the approximate Bayes credible interval suggests that the computational effort that must be expended in any bootstrap may not be necessary.

We have focused on the simplest possible QTL model: a single QTL with normally distributed residual variation. This simple model is not likely to hold in practice. An especially important departure concerns the presence of multiple linked QTLs. A confidence interval for QTL location derived from the results of analysis using single-QTL models has little meaning if there exist multiple QTLs on the chromosome. The LOD support and Bayes credible intervals have obvious extensions for the case of multiple QTLs; their performance, especially in the case of multiple linked QTLs, deserves further study.

While we have shown that bootstrap confidence intervals for QTL location perform poorly and so should not be used in this context, the LOD support and approximate Bayes credible intervals were seen to behave appropriately. This is in broad agreement with Dupuis and Siegmund (1999). They studied the performance of LOD support and Bayes credible intervals, focusing on the widths of the intervals. They found that when LOD support and Bayes credible intervals had similar coverage, their widths were generally comparable. For LOD intervals to have the target coverage properties, the LOD drop has to be adjusted, while the Bayes intervals give consistent coverage for a range of marker densities and QTL effects. Thus, the approximate Bayes credible intervals are particularly attractive; a nominal 96.5 or 97% Bayes credible interval was seen to exhibit coverage near 95% for different sample sizes, marker densities, and sizes of QTL effect.
Finally, we wish to emphasize that 95% is not a magic number, and investigators may wish to be more conservative (seeking, for example, 99% coverage), so that, for example, the formation of a congenic line does not miss the true location of the QTL.

ACKNOWLEDGMENTS

This work was supported in part by NIH grant GM074244 (to KWB) and a NSF Graduate Research Fellowship (to AM).
LITERATURE CITED


FIGURE LEGENDS

**Figure 1.** Results for the chromosome 4 data of Sugiyama et al (2001). **A.** The LOD curve and the 1.5-LOD support interval. Tick marks at the bottom of the figure indicate the locations of the genetic markers. **B.** A histogram of the estimated QTL locations in 10,000 bootstrap replicates, and the 95% bootstrap confidence interval, calculated by the method of Visscher et al. (1996).

**Figure 2.** Estimated distribution of the MLE of QTL location, \( \hat{\theta} \), as a function of the true location of the QTL, \( \theta \), for \( \theta \) varying from 45 to 50. The results are based on 10,000 simulation replicates of a backcross with 200 individuals for a chromosome of length 100 cM and having 11 equally spaced markers, and with the heritability due to the QTL at 10%.

**Figure 3.** Coverage of 95% bootstrap confidence intervals (black), 1-LOD support intervals (red) and approximate 95% Bayes credible intervals (blue) and as a function of the true QTL position, \( \theta \). The dashed vertical gray lines denote marker positions on the chromosome.

**Figure 4.** Coverage of 95% bootstrap confidence intervals (black), 1-LOD support intervals (red) and approximate 95% Bayes credible intervals (blue) as a function of the MLE of QTL position, \( \hat{\theta} \). The dashed vertical gray lines denote marker positions on the chromosome.

**Figure 5.** Estimated amount to drop in a LOD support interval (A and C) and the nominal Bayes coverage for the approximate Bayes credible interval (B and D) to give 95% coverage, on the basis of 100,000 simulation replicates. Backcross (A and B) and intercross (C and D) experiments with either 200 (black curves) or 500 (red curves) individuals were considered. The line types indicate different possible marker spacings. Values are plotted as a function of
the effect of the QTL, scaled according to the power to detect the QTL.

**Figure 6.** Coverage of the 1.5-LOD support interval in a backcross (A), the 96.5% Bayes interval in a backcross (B), the 1.8-LOD support interval in an intercross (C), and the 97% Bayes interval in an intercross (D), on the basis of 100,000 simulation replicates. The black curves are for 200 individuals; the red curves are for 500 individuals. The line types indicate different possible marker spacings. Values are plotted as a function of the effect of the QTL, scaled according to the power to detect the QTL.
Figure 1. Results for the chromosome 4 data of SUGIYAMA et al (2001). A. The LOD curve and the 1.5-LOD support interval. Tick marks at the bottom of the figure indicate the locations of the genetic markers. B. A histogram of the estimated QTL locations in 10,000 bootstrap replicates, and the 95% bootstrap confidence interval, calculated by the method of VISSCHER et al. (1996).
Figure 2. Estimated distribution of the MLE of QTL location, $\hat{\theta}$, as a function of the true location of the QTL, $\theta$, for $\theta$ varying from 45 to 50. The results are based on 10,000 simulation replicates of a backcross with 200 individuals for a chromosome of length 100 cM and having 11 equally spaced markers, and with the heritability due to the QTL at 10%.
Figure 3. Coverage of 95% bootstrap confidence intervals (black), 1-LOD support intervals (red) and approximate 95% Bayes credible intervals (blue) and as a function of the true QTL position, $\theta$. The dashed vertical gray lines denote marker positions on the chromosome.
Figure 4. Coverage of 95% bootstrap confidence intervals (black), 1-LOD support intervals (red) and approximate 95% Bayes credible intervals (blue) as a function of the MLE of QTL position, $\hat{\theta}$. The dashed vertical gray lines denote marker positions on the chromosome.
Figure 5. Estimated amount to drop in a LOD support interval (A and C) and the nominal Bayes coverage for the approximate Bayes credible interval (B and D) to give 95% coverage, on the basis of 100,000 simulation replicates. Backcross (A and B) and intercross (C and D) experiments with either 200 (black curves) or 500 (red curves) individuals were considered. The line types indicate different possible marker spacings. Values are plotted as a function of the effect of the QTL, scaled according to the power to detect the QTL.
Figure 6. Coverage of the 1.5-LOD support interval in a backcross (A), the 96.5% Bayes interval in a backcross (B), the 1.8-LOD support interval in an intercross (C), and the 97% Bayes interval in an intercross (D), on the basis of 100,000 simulation replicates. The black curves are for 200 individuals; the red curves are for 500 individuals. The line types indicate different possible marker spacings. Values are plotted as a function of the effect of the QTL, scaled according to the power to detect the QTL.