Bayesian inference for genomic imprinting underlying developmental characteristics

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Abstract
The identification of imprinted genes is becoming a standard procedure in searching for quantitative trait loci (QTL) underlying complex traits. When a developmental characteristic such as growth or drug response is observed at multiple time points, understanding the dynamics of gene function governing the underlying feature should provide more biological information regarding the genetic control of an organism. Recognizing that differential imprinting can be development-specific, mapping imprinted genes considering the dynamic imprinting effect can provide additional biological insights into the epigenetic control of a complex trait. In this study, we proposed a Bayesian imprinted QTL (iQTL) mapping framework considering the dynamics of imprinting effects and model multiple iQTLs with an efficient Bayesian model selection procedure. The method overcomes the limitation of likelihood-based mapping procedure, and can simultaneously identify multiple iQTLs with different gene action modes across the whole genome with high computational efficiency. An inference procedure using Bayes factors to distinguish different imprinting patterns of iQTL was proposed. Monte Carlo simulations were conducted to evaluate the performance of the method. The utility of the approach was illustrated through an analysis of a body weight growth data set in an F2 family derived from LG/J and SM/J mouse stains. The proposed Bayesian mapping method provides an efficient and computationally feasible framework for genome-wide multiple iQTL inference with complex developmental traits.

Keywords: Bayes factor; Bayesian model selection; developmental traits; Markov chain Monte Carlo; imprinting pattern; quantitative trait loci

INTRODUCTION
Genomic imprinting is a genetic phenomenon in which the same genes express differently, depending on their parental origin [1]. On the molecular level, genomic imprinting may result from DNA methylation, histone modification, non-coding RNAs (ncRNA) and even long distance interchromosomal interactions [2]. As a ubiquitous phenomenon in nature, genomic imprinting has been broadly identified in plants [3], animals [4, 5] and humans [6, 7]. The role of genomic imprinting in shaping an organism’s development has been unanimously recognized [8–10]. The imprinting effect on traits of interest can be characterized by different types. When the paternal allele at a gene is expressed and the maternal allele is inactivated, this feature of imprinting is referred to as paternal imprinting. Maternal imprinting is defined similarly. Genomic imprinting has been traditionally viewed as a mono-allelic expression with complete maternal or
imprinting effect at specific developmental stages, such an approach can estimate and test time-specific mathematical function that best describes a developmental characteristic which incorporates a developmental iQTL mapping framework underlying development in mammals. As a highly complex process, genomic imprinting is involved in a number of growth axes operating coordinately at different developmental stages, and shows time-dependent effect during development. The unbalanced expression of an imprinted gene that occurs during a development stage challenges the traditional paradigm of inheritance and mapping methods. We argue that traditional methods, by treating a trait measured at a certain developmental stage as mapping subject, without considering the correlation information at different developmental stages, are less powerful in dissecting dynamic iQTL effects. Cui et al. recently proposed a functional iQTL mapping framework underlying developmental characteristics which incorporates a mathematical function that best describes a developmental feature into an iQTL mapping framework. Such an approach can estimate and test time-specific imprinting effect at specific developmental stages, and displays several merits over traditional iQTL mapping methods.

Current mapping procedures for iQTL inference are all based on single iQTL models, estimating and testing one locus at a time without considering the effects of other iQTLs. When multiple iQTLs are presented in the genome, such approaches are less efficient under the likelihood-based framework. For a dynamic trait, the number of parameters being estimated is several folds larger than those for a univariate trait. In our previous QTL mapping model, we demonstrated that a Bayesian mapping method can handle this issue well with high computational efficiency. In this study, we unify the two endeavors, Bayesian mapping of developmental traits and iQTL inference, into a unified framework called Bayesian functional multiple iQTL mapping (Bafmim). We propose an efficient Bayesian model selection strategy for multiple iQTL inference for developmental traits. The inference for the number, position and effect of multiple iQTLs as well as for different imprinting patterns is provided.

The statistical behavior of the proposed method is illustrated by simulation studies. The utility of the method is shown by applying to a real data set. A total of six iQTLs are identified with Bafmim, among which two were missed by the likelihood-based method. The proposed approach has great implications in understanding the function of imprinted genes governing developmental characteristics.

**STATISTICAL METHOD**

**The imprinting model**

In a mapping population, assume that there are four distinguishable genotypes, denoted by $Q_M Q_P$, $Q_M q_P$, $q_M Q_P$ and $q_M q_P$, at each locus where the subscripts $M$ and $P$ refer to an allele inherited from the mother and father, respectively. A set of codominant molecular markers can be genotyped and phenotypes for a developmental trait which is measured at $m$ time points on $n$ individuals. In general, the additive effect $a$ is defined as half of the phenotypic difference between two homozygotes; the dominance effect $d$, is defined as the difference between the joint mean of both heterozygotes and the mean of both homozygotes; and the imprinting effect $i$, is defined as the difference between both heterozygotes [19, 29]. Following the definitions for the genetic parameters, an imprinting model for a phenotype measured for...
where q is the number of potential (i)QTLs in the genome; \( \mu(t) \) is the population mean at time t; \( \alpha_j(t), d_j(t) \) and \( i_j(t) \) for \( j = 1, 2, \ldots, q \) are the additive, dominance and imprinting effects of the j-th iQTL at time point t; \( \xi_k(t) \) \( (k = 1, 2, \ldots, n) \) is an individual-specific time-dependent random environmental term modeled as a mean-zero Gaussian process, i.e. \( N(0, \sigma^2_\xi(t)) \) and \( \epsilon_k(t) \) is a random environmental error assumed to be normally distributed with mean zero and variance \( \sigma^2 \). Note that \( c_{kj}, z_{kj} \) and \( s_{kj} \) are genotype-specific indicator variables related to genetic effects \( \alpha_j(t), d_j(t) \) and \( i_j(t) \), which are defined by Mantey et al. [30] as:

\[
c_{kj} = \begin{cases} 
+1 & \text{for } Q_M Q_P \\
0 & \text{for } Q_M \bar{q} P \\
-1 & \text{for } \bar{q} M q P \\
0, z_{kj} = \begin{cases} 
+1 & \text{for } Q_M Q_P \\
0 & \text{for } Q_M \bar{q} P \\
-1 & \text{for } \bar{q} M q P \\
0, s_{kj} = \begin{cases} 
+1 & \text{for } Q_M Q_P \\
0 & \text{for } Q_M \bar{q} P \\
-1 & \text{for } \bar{q} M q P 
\end{cases}
\end{cases}
\]

Note that in practical mapping, unless QTLs are located exactly at the marker position (i.e. the markers and the QTLs overlap), these genotype-specific indicator variables are unknown and need to be inferred from flanking markers. We use Legendre polynomial of order r to fit changing trajectories of the population mean and the effects of each iQTL [31, 32]. Let \( \psi(t) = (\psi_0(t), \ldots, \psi_r(t))^T \) be the basis of the Legendre polynomial with order r and have that \( \mu(t) = \psi(t)^T \mu, a_j(t) = \psi(t)^T a_j, d_j(t) = \psi(t)^T d_j, i_j(t) = \psi(t)^T i_j \) and \( \xi_k(t) = \psi(t)^T \xi_k, \) where each one of \( \mu, a_j, d_j, i_j \) and \( \xi_k \) is a vector of r + 1 dimensions. Model (1) can then be rewritten as:

\[
y_k(t) = \psi^T(t) \mu + \sum_{j=1}^{q}[c_{kj}\psi^T(t)a_j + w_{kj}\psi^T(t)d_j + s_{kj}\psi^T(t)i_j] + \xi_k(t) + \epsilon_k(t) 
\]

Bayesian model selection for genetic parameters

Model (3) is a mixed-effect model where population mean and genetic effects are fixed and the time-dependent environmental effect is random. Also note that Model (3) is not a regular linear mixed-effect model, since the number of independent variables for the fixed effects and the associated indicator variables are unknown due to unknown number of QTLs. In principle, QTLs can be distributed anywhere in the genome, and hence any genomic positions can be potential QTL locations. Thus, we approximate them for all possible iQTLs by partitioning the entire genome into evenly spaced loci by 1 or 2 cM, covering all observed markers and additional loci between flanking markers. The expected values of the indicator variables at each locus can be calculated for an F_{2} mapping population based on the conditional probabilities of a QTL genotype given on two flanking markers as follows:

\[
E(z) = \pi_{QQ} - \pi_{qQ},
E(w) = \pi_{Qq} + \pi_{qq} \quad \text{and} \quad E(s) = \pi_{Qq} - \pi_{qq}
\]

with \( \pi_{QQ}, \pi_{Qq}, \pi_{qq} \) and \( \pi_{qQ} \) being the conditional probabilities of a tested QTL genotype \( Q_M Q_P, Q_M q P, q M q P \) or \( q M q P \) given on two flanking markers (see Table 1 in Cui et al. [21] for details).

For a supersaturated model where each genomic location could contain a potential iQTL, it is almost impossible to estimate a huge number of genetic effects. So we preset an upper bound on the number of iQTLs in the model [52]. The upper bound should be larger than the potential number of detectable iQTLs in a given data set. Given an upper bound on the number of iQTLs (L), these iQTLs can be drawn from densely spaced loci over the genome. Even with a moderate number of upper bound, there are many genetic effects being
estimated in Model (3). In order to infer the existence of these effects, we introduce a random binary variable $\gamma$ to indicate which genetic effects should be included in or excluded from the model, corresponding to $\gamma = 1$ or $\gamma = 0$ [33–35]. Model (3) then becomes

$$y_k = \psi^T \mu + \sum_{l=1}^{L} \left[ \gamma_{dl} \xi_l \psi^T a_l + \gamma_{dl} w_l \psi^T d_l + \gamma_{dl} \xi_l \psi^T i_l \right]$$

$$+ \psi^T \xi_k + \epsilon_k, \tag{4}$$

where $\gamma_{dl} (l = a, d, i \text{ or } i)$ is the indicator variable for genetic effects $a_l, d_l, i_l$. Within the framework of Bayesian model selection, Bayesian sampling for unknown parameters including $\mu, \gamma, a_l, d_l, i_l, \xi_k, \Sigma$ and QTL position $\lambda_j$, is implemented with the Markov chain Monte Carlo (MCMC) algorithm. Note that the released sampling value for binary variable $\gamma$ at a previous round determines which genetic effects and position of an iQTL should be drawn or estimated at the next round. This will save computing time significantly.

**Likelihood function**

Given unknown parameters $\theta$, the likelihood function of model (4) can be expressed as

$$L(\theta|y) = \prod_{k=1}^{n} p(y_k|\theta) \propto |V|^{(n+1)/2} \exp \left[ \sum_{k=1}^{n} (y_k - U_k)^T \psi^T \psi \right],$$

where $U_k = \psi^T \mu + \sum_{l=1}^{L} \left[ \gamma_{dl} \xi_l \psi^T a_l + \gamma_{dl} w_l \psi^T d_l + \gamma_{dl} \xi_l \psi^T i_l \right] + \psi^T \xi_k$ and $V = \psi^T \Sigma \psi + I \sigma^2$.

**Prior specification**

Following Yi et al., we take upper bound of iQTL number $L$ as $l_0 + 3\sqrt{l_0}$, where $l_0$ is the prior expected number of iQTL loci and is determined according to initial investigations with traditional methods [26]. The binary indicator $\gamma$ is assumed to have an independent prior $p(\gamma_k) = \prod_{k=1}^{n} u_{\gamma_k} (1 - u_{\gamma_k})^{(1-\gamma_k)}$, where $u_{\gamma_k} = 1 - \left[ 1 - \frac{1}{n} \right]^{(1-\gamma_k)}$ is the prior inclusion probability for certain QTL effect [52]. Priors on iQTL positions are assumed to be independent and uniformly distributed over the entire genome.

The prior for the population mean $\mu$ is assumed to be $N(\mu_0, \Sigma_0)$. We can empirically set $\mu_0 = \frac{1}{n} \sum_{k=1}^{n} \mu_k$ and $\Sigma_0 = \frac{1}{n-1} \sum_{k=1}^{n} (b_k - b)(b_k - b)^T$, where $b_k = (\psi^T \psi)^{-1} \psi^T y_k$ is a vector of regression coefficients obtained by fitting individual dynamic trajectory.

We propose the following hierarchical mixture prior for each additive genetic effect, $\beta_i \sim N_{p+1}(0, \Sigma_i)$ with $\Sigma_i = \gamma_i \mathbf{V}^{-1} \psi \sum_{k=1}^{n} (y_k - \epsilon_k)^{-1}$ and $\epsilon$ being taken to $n$ such that the prior variance of each fixed effect stays approximately the same as $n$ increases.

The random effects $\xi_k$ are assumed to have an independent multivariate normal distribution. That is, $\xi_k \sim N_{p+1}(0, \Sigma_{\xi})$, being an $(r + 1) \times (r + 1)$ matrix. An inverse Wishart prior is chosen for the covariance matrix of regression coefficients for random environmental effect, denoted by $\Sigma \sim IW_{p+1}(\nu_\xi, \nu_\xi S_{\xi})$ with $\nu_\xi$ being hyperparameter. The residual variance is assumed to have a scaled inverse $\chi^2$ distribution, i.e. $\sigma^2 \sim IC(\nu_e, \frac{1}{\nu_e})$ where $\nu_e$ and $s_e$ are hyperparameters. $p(\lambda_j) = \frac{1}{l_j}$ for $j = 1, 2, \ldots, L$, where $l_j$ is the distance between the two neighboring QTLs [36–38].

**MCMC sampling**

A joint posterior density can be formed by multiplying priors of all unknown parameters and the likelihood function above. The joint posterior distribution is analytically intractable, hence we need to derive the 'conditional posterior' for each unknown parameter from the joint posterior density. The details about the derivation can be found in the Appendix 1. A MCMC methods such as Gibbs sampler [39] and Metropolis–Hastings algorithm [40, 41] are applied to sample each parameter conditional on all other parameters. Prior to implementing MCMC sampling, we set an upper bound for the number of QTLs and distribute these QTLs on the genome evenly and initialize all variables with some initial values or values sampled from their prior distributions (see the Appendix 1 for details about prior specification). The MCMC sampling procedure for unknown parameters is summarized as follows:

(i) Update population mean $\mu$ by sampling from a normal distribution.

(ii) Update the genetic effects $\beta_i$ corresponding to $\gamma_i = 1$ by drawing from a normal distribution.

(iii) Update the binary indicators $\gamma_i$ by adopting an efficient Metropolis–Hastings algorithm [52].
(iv) Update individual-specific regression coefficients for random environmental effects $\xi_k$ by sampling from a normal distribution.
(v) Update the covariance matrix $\Sigma$ of $\xi_k$ by drawing from an inverse Wishart distribution.
(vi) Update the residual variance $\sigma^2$ by sampling from an inverse $\chi^2$ distribution.
(vii) Update joint QTL positions $\lambda$ by adopting the Metropolis–Hastings algorithm.
(viii) Repeat steps (i)–(viii) until the Markov chain reaches a desirable length.

**Post-MCMC analysis**

Post-MCMC analysis includes the monitoring of the mixing behavior and convergence rates of the MCMC algorithm, and the assessment of characteristics of the imprinting genetic architecture. The former can be checked by visually inspecting trace plots of the sample values of scalar quantities of interest or formal diagnostic methods provided in the package R/coda [42]. The latter can use model averaging which accounts for model uncertainty and average over possible models weighted by their posterior probabilities [43–45]. The posterior inclusion probability for each locus is estimated as its frequency in the posterior samples. Bayes factor (BF) is used as a measurement for inclusion against exclusion at each iQTL locus [46]. Generally, a threshold of BF is empirically determined as 3, or $2\ln BF = 2.1$, for declaring statistical significance of an (i)QTL.

**Bayesian inference for imprinting mode of action**

Generally speaking, (i)QTLs detected with the above Bayesian algorithm cannot be declared as iQTLs until we do further imprinting inference. After an (i)QTL is detected, we can adopt the idea of a Bayes factor to infer statistical significance for its imprinting effect with the form:

$$BF = \frac{p_g}{1 - p_g} \cdot \frac{1 - p}{p}$$

where $p$ is a prior probability and $p_g$ is a posterior probability for a certain genetic effect, which is calculated as the proportion of samples in which $\gamma_g = 1$ in MCMC sampling rounds. If the $BF$ is >3 (or $2\ln BF > 2.1$) for the imprinting effect $i$, then the detected iQTL can be claimed as a true iQTL, otherwise as a Mendelian QTL.

Following the definition of imprinting types and the corresponding null hypothesis [13, 47, 48], we classify imprinting patterns as parental imprinting, i.e. $a = \pm i$ and $d = 0$ which includes paternal ($a = i$) and maternal ($a = -i$) imprinting subtypes; and dominance imprinting with $a = 0$ but $i \neq 0$, which can be further distinguished as bipolar imprinting in which $d = 0$ and $i \neq 0$ and polar imprinting in which $d = \pm i$. Since the imprinting pattern for the detected QTL depends on whether genetic effect $a$ or $d$ equals imprinting effect $i$, it can also be statistically inferred through the Bayes factor statistic. The Bayes factor can be simplified as the ratio of posterior probabilities for the genetic effects being compared, due to same prior probability for each genetic effect. Imprinting patterns, hypotheses and corresponding statistical criteria for the iQTLs are detailed in [48].

**SIMULATION STUDIES**

We conducted simulation studies to evaluate the performance of the proposed Bayesian functional multiple iQTL mapping approach. A genome consisting of one single large chromosome of 600 cM was simulated covering 61 evenly spaced markers. The growth pattern of a dynamic trait was assumed to be controlled by one QTL inherited in a Mendelian fashion and four iQTLs with their imprinting patterns, positions and effects listed in Table 1. The order of the Legendre polynomial that generates the developmental trajectory was assumed to be $r = 3$. The trajectory for each (i)QTL genetic effect is plotted in Figure 1. We simulated a dynamic trait measured at eleven time points assuming different sample sizes ($n = 250, 500$) with inbred $F_2$ individuals. The marker and QTL genotypes in the $F_2$ family were generated by mimicking sex-specific recombination fractions (see Cui et al. [21] for more details). The population mean and the individual-specific environmental error covariance matrix were set the same as described in Yang and Xu [31] and the residual variance was set as four.

In Bayesian analyses for simulated data, we set the prior number of main–effect QTLs as 3. The upper bound of the number of QTLs was then equal to $L = 3 + 3\sqrt{3} = 8$. The actual values for the hyper-parameters used here mimic the results obtained in real data analyses (see the real data analysis section). The initial values of all variables were sampled from their prior distributions. The MCMC was run for
Table 1: The imprinting type and parameters (regression coefficients of Legendre polynomials) of iQTLs used in simulation studies

<table>
<thead>
<tr>
<th>QTL no.</th>
<th>Imprinting type</th>
<th>Position</th>
<th>$a_0$</th>
<th>$a_1$</th>
<th>$a_2$</th>
<th>$a_3$</th>
<th>$d_0$</th>
<th>$d_1$</th>
<th>$d_2$</th>
<th>$d_3$</th>
<th>$i_0$</th>
<th>$i_1$</th>
<th>$i_2$</th>
<th>$i_3$</th>
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<tbody>
<tr>
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<td>Mendelian</td>
<td>23</td>
<td>1.82</td>
<td>-0.80</td>
<td>-1.20</td>
<td>-0.80</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Paternal</td>
<td>148</td>
<td>0.00</td>
<td>1.65</td>
<td>2.52</td>
<td>1.20</td>
<td>0.00</td>
<td>1.65</td>
<td>2.52</td>
<td>1.20</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Maternal</td>
<td>256</td>
<td>2.55</td>
<td>1.36</td>
<td>-2.02</td>
<td>-1.27</td>
<td>-2.55</td>
<td>-1.36</td>
<td>2.02</td>
<td>1.27</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>4</td>
<td>Bipolar</td>
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<td></td>
<td></td>
<td></td>
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<td>2.94</td>
<td>0.00</td>
<td>1.08</td>
<td>1.72</td>
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<td>5</td>
<td>Polar</td>
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<td>-1.28</td>
<td></td>
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</tr>
</tbody>
</table>

Figure 1: The trajectories for genetic effects of the simulated (i)QTLs: (1) $d = i = 0$, (2) $a = i$, (3) $a = -i$, $d = 0$, (4) $a = d = 0$ and (5) $d = i$, $a = 0$. In each plot, time is the original time measurement ranging from 1 to 10 weeks as in real data.
10,000 cycles as a burn-in period (deleted) and then for an additional 150,000 cycles after the burn-in. The chain was then thinned to reduce serial correlation by saving one observation in every 50 cycles. The posterior sample contained 3,000 observations for post-MCMC analysis. Note that the length of the burn-in was judged by visually inspecting the plots of some posterior samples across rounds and was set to enough cycles to ensure MCMC convergence. The simulation experiment was replicated 100 times in order to evaluate the statistical power of our method.

Table 2 shows the mean estimates as well as their SD (in parenthesis) for the parameters given in Table 1. The relative statistical power to detect each QTL is also listed. Overall, the Bayesian mapping approach is able to estimate the regression effects of the iQTLs with reasonable precision. All four iQTL positions can be accurately estimated with high precision. As we expected, increasing sample size always leads to small bias, increased precision of parameter estimation, and high mapping power. For example, the mapping power for QTL 1 increased from 70 to 85% when sample size was increased from 250 to 500. Even with small sample size \( n = 250 \), the QTL position can also be estimated with high precision. In addition, we can also accurately infer the imprinting pattern of the detected locus using Bayes factor (data not shown). These indicate the power of Bayesian mapping for multiple iQTL inference. The simulated data sets were also analyzed by likelihood-based functional iQTL mapping [26]. We found that the likelihood-based functional mapping provides less accurate iQTL parameter estimates, cannot infer the given imprinting pattern well, and performs poorly with relatively low statistical power of iQTL detection than the proposed Bayesian method (Table 3). Moreover, the likelihood-based functional mapping has a high false positive rate (FPR) in iQTLs detection with 12% for sample size of 250 and 9% for sample size of 500, while the Bayesian method gives lower FPR. As the maximum likelihood-based method is not efficient in handling multiple iQTLs, this might explain the performance difference of the two approaches. The simulation results indicate the robustness of the proposed method in multiple iQTL detection for dynamic traits with moderate sample size.

Under the same simulation scenarios, replacing both time-dependent permanent environmental

<table>
<thead>
<tr>
<th>Sample size</th>
<th>QTL no.</th>
<th>a_0</th>
<th>a_1</th>
<th>a_2</th>
<th>a_3</th>
<th>d_0</th>
<th>d_1</th>
<th>d_2</th>
<th>d_3</th>
</tr>
</thead>
<tbody>
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<td>250</td>
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<td>1.77 (0.23)</td>
<td>-0.05 (0.25)</td>
<td>-0.97 (0.33)</td>
<td>-1.07 (0.40)</td>
<td>-1.70 (0.45)</td>
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<tr>
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<td>2</td>
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<td>1.18 (0.38)</td>
<td>1.59 (0.49)</td>
<td>1.50 (0.39)</td>
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<tr>
<td></td>
<td>3</td>
<td>2.38 (0.25)</td>
<td>1.84 (0.25)</td>
<td>1.98 (0.34)</td>
<td>1.98 (0.34)</td>
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<tr>
<td></td>
<td>4</td>
<td>1.66 (0.21)</td>
<td>0.85 (0.23)</td>
<td>1.18 (0.34)</td>
<td>2.49 (0.34)</td>
<td>2.04 (0.33)</td>
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<tr>
<td></td>
<td>5</td>
<td>2.56 (0.25)</td>
<td>0.02 (0.19)</td>
<td>1.02 (0.24)</td>
<td>1.40 (0.24)</td>
<td>0.72 (0.24)</td>
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<tr>
<td>500</td>
<td>1</td>
<td>1.96 (0.22)</td>
<td>0.85 (0.23)</td>
<td>1.18 (0.34)</td>
<td>2.49 (0.34)</td>
<td>2.04 (0.33)</td>
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<td></td>
<td>2</td>
<td>-0.02 (0.19)</td>
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<td>3</td>
<td>2.56 (0.25)</td>
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Table 2: Mean estimates and SDs (in parentheses) of iQTL regression effects with polynomial residuals.
effects and residual errors in model (1) with AR(1) structure where autoregressive coefficient was taken as 0.6, we simulated phenotypes and fit model (1) to the simulated data sets with the Bayesian method. Table 4 lists mean estimates and SDs (in parentheses) of the given (i)QTL regression effects. As expected, the Bayesian method can powerfully identify the given (i)QTLs and estimate the given (i)QTL regression effects reasonably well based on model (1), indicating the flexibility of our model in fitting different time-dependent residuals.

We also simulated data assuming the true dynamics of the repeated measurement following a logistic regression curve for genotypic effects of the five simulated QTLs. The simulated parameters for the logistic function are obtained by fitting changes in genotypic means generated from the Legendre polynomials (Supplementary Table 1). Residuals are modeled by AR(1) structure with autoregressive coefficient of 0.6. Estimates of QTL positions and powers to detect QTLs are listed in Supplementary Table 2. The results indicate that the proposed method performs better than the likelihood-based method.

### REAL DATA ANALYSIS

We illustrated the application of our proposed approach by reanalyzing a mouse body weight growth data set with an F2 mating population derived from two inbred strains, the Large (LG/J) and the Small (SM/J). A total of 502 F2 mice were genotyped for 96 microsatellite markers located on 19 autosomal chromosomes. A linkage map of a total length 1780 cM has been constructed [49]. The body mass was measured on each mouse at 10 weekly intervals starting at Day 7. The raw weights were adjusted for the effects of each covariate due to dam, litter size at birth and parity, and sex [49]. The data set has been analyzed by Cui et al. [26] with likelihood-based functional mapping.

By fitting the mean change of weight growth over age, we chose the Legendre polynomial of order 3 as the base model to describe the changing trajectory for each component except for residuals, described in Model (1). The female-to-male recombination rate of 1.25:1 is used to estimate conditional probabilities for the four QTL genotypes [21]. The expected number of main-effect QTLs was set as $l_0 = 4$ according to the results by Cui et al. [26] and the upper bound of the number of QTLs was then calculated as $L = 4 + 3\sqrt{4} \approx 10$. Thus, the prior inclusion probability for QTL effects was determined to be $1 - \left(1 - \frac{1}{4}\right)^L = 0.156$. The actual values for the hyperparameters were set as $S_b = S_e = 0.5I$, $v_b = r + 1$ and $v_e = 0$. The initial values of all variables were sampled from their prior distributions. The MCMC was run 200 000 cycles after the burn-in period of 10 000 cycles. The analysis of the real data set with our MATLAB code took ~3 h on an Intel Core 2 PC with a 2 GHz processor and 4 GB RAM.

Figure 2 plots the profiles of $2\log$BF obtained with the Bayesian model selection. The top panel shows that there are six peaks for $2\log$BFs exceeding the horizontal reference line with an empirical critical value 2.1, indicating that six iQTLs are detected on chromosomes 2, 4, 6, 7, 10 and 15. All iQTLs detected show significant imprinting effect, as their relative $2\log$BFs are $>2.1$ for imprinting effects (the bottom panel in Figure 2). Table 5 tabulates the position on each chromosome and the estimated effects (additive, dominance and imprinting) for the six detected iQTLs. The estimated regression coefficients for each iQTL have no biological meaning, but they can be used to predict the effects of an iQTL at a given time point by substituting time
into the Legendre polynomial with the regression coefficients. Figure 3 shows the dynamic trajectory of the genetic effects over time. Among the six iQTLs, the main effect (i.e. $a$ or $d$) shows a stronger effect on mouse growth than the imprinting effect, which makes biological sense as imprinting effect generally plays a modifying role.

We further evaluated the imprinting pattern of the six detected iQTLs. Two of them (on chromosomes 4 and 10) show polar imprinting, two (on chromosomes 2 and 6) show maternal imprinting and two (on chromosomes 7 and 15) show paternal imprinting, based on the results obtained from the significance analysis given in Table 6. Compared to the likelihood-based method [26], the Bayesian method identified two more QTLs (on chromosomes 2 and 4).

**DISCUSSION**

The epigenetic phenomenon in genomic imprinting has been constantly challenging and revising the traditional paradigm of inheritance. The inheritable property of imprinting provides clues for complicated genetic disorders [6]. In the meantime, it also brings challenges for statistical modeling and mapping. We developed a Bayesian model selection method to identifying multiple iQTLs for developmental traits illustrated in an F2 mating population. Extensions to other mating designs such as a reciprocal backcross design [50], is straightforward. The Bayesian method has shown its relative merits in handling multiple QTLs partly due to its flexibility to handle a large parameter space [28, 31, 51, 52]. Both simulation and real data analysis indicate the power and relative advantage of the proposed Bayesian multiple iQTL mapping for dynamics traits compared to the likelihood-based mapping method. In addition, we proposed several inference procedures to infer different imprinting types, which have not been discussed in the likelihood-based iQTL mapping study. The identified iQTLs as well as their imprinting property provide valuable information for further experimental verification.

The current method is developed specifically for longitudinal or functional traits. Recently, Hayashi and Awata [15] proposed a Bayesian mapping approach that can simultaneously map multiple QTLs, and further discriminating Mendelian and imprinting expressions of a QTL. Although the approach shows improvement in iQTL detection,
it is limited by a number of facts. For example, drawing number of QTLs with a reversible-jump MCMC procedure may have low convergence efficiency. Moreover, the method is developed for univariate traits and ignores the dynamics of gene effects. In an earlier article, Yang et al. [53] developed a Bayesian multiple iQTL mapping method for univariate traits. The current work is an extension of our previous work, but is more challenging in modeling the dynamics of genetic effects.

Our method assumes a maximum number of detectable iQTLs and introduces latent binary variables to indicate which main effects for a putative iQTL should be included or excluded from the model. Compared to the likelihood-based method [26], it allows MCMC sampling for iQTL parameters to carry out in the reduced model space, and thus enhances the computational efficiency of Bayesian multiple iQTL mapping with many parameters. Computationally, it also outperforms the likelihood-based method. In addition, it facilitates statistical inference for imprinting patterns of the detected iQTLs with appropriately defined Bayes factors. In the real data analysis, we took the dynamic course of a developmental trait as the mapping subject, and identified more iQTLs altering the developmental trajectory than separately performing iQTL mapping at each time point [21].

The key issue for iQTL mapping of developmental traits is to choose appropriate submodels for imprinting inference. In the likelihood-based mapping framework, Cui et al. [26] proposed to describe

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**Table 5:** Estimates of iQTL positions and regression coefficients for body weight growth in mice

<table>
<thead>
<tr>
<th>QTL</th>
<th>Position (Chr.-cM)</th>
<th>$a_0$</th>
<th>$a_1$</th>
<th>$a_2$</th>
<th>$a_3$</th>
<th>$d_0$</th>
<th>$d_1$</th>
<th>$d_2$</th>
<th>$d_3$</th>
<th>$i_0$</th>
<th>$i_1$</th>
<th>$i_2$</th>
<th>$i_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2–104.3</td>
<td>0.254</td>
<td>0.633</td>
<td>0.166</td>
<td>0.033</td>
<td>0.074</td>
<td>0.180</td>
<td>0.061</td>
<td>0.013</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>4–38.1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>6–73.2</td>
<td>0.425</td>
<td>0.525</td>
<td>0.075</td>
<td>0.055</td>
<td>0.162</td>
<td>0.210</td>
<td>0.036</td>
<td>0.032</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>7–63.1</td>
<td>0.787</td>
<td>0.851</td>
<td>0.044</td>
<td>0.103</td>
<td>0.185</td>
<td>0.210</td>
<td>0.016</td>
<td>0.141</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>10–72.7</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.027</td>
<td>0.165</td>
<td>0.018</td>
<td>0.018</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>15–12.7</td>
<td>0.896</td>
<td>0.311</td>
<td>0.314</td>
<td>0.062</td>
<td>0.023</td>
<td>0.061</td>
<td>0.004</td>
<td>0.006</td>
<td>0.017</td>
<td>0.071</td>
<td>0.035</td>
<td>0.034</td>
</tr>
</tbody>
</table>
the changes of iQTL genotypic effects by a logistic growth function. This parametric assumption is, however, difficult to implement in a multiple iQTL model for Bayesian mapping due to the non-linearity of different submodels. With the orthogonal polynomial function, it is flexible enough to capture the underlying trajectory for dynamic iQTL inference. One issue when fitting data with a polynomial function is to appropriately select optimal polynomial order. Cui et al. [32] proposed two methods to choose an optimal order: (i) assume the same order for different phenotypic

![Figure 3: The trajectories for different genetic effects (a, i and d) of the detected iQTLs for body weight growth in mice.](image)

<table>
<thead>
<tr>
<th>QTL</th>
<th>a</th>
<th>d</th>
<th>i</th>
<th>Imprinting type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.29</td>
<td>0.64</td>
<td>2.20</td>
<td>Additive-Paternal</td>
</tr>
<tr>
<td>2</td>
<td>1.01</td>
<td>3.34</td>
<td>4.47</td>
<td>Dominance-Over</td>
</tr>
<tr>
<td>3</td>
<td>4.72</td>
<td>1.83</td>
<td>3.24</td>
<td>Additive-Paternal</td>
</tr>
<tr>
<td>4</td>
<td>2.95</td>
<td>0.54</td>
<td>2.97</td>
<td>Additive-Paternal</td>
</tr>
<tr>
<td>5</td>
<td>0.87</td>
<td>4.32</td>
<td>3.68</td>
<td>Dominance-Over</td>
</tr>
<tr>
<td>6</td>
<td>4.46</td>
<td>1.70</td>
<td>4.23</td>
<td>Additive-Paternal</td>
</tr>
</tbody>
</table>

Table 6: List of 2lnBFs for genetic effects and imprinting types of the detected iQTLs for body weight growth in mice
trjectories which can be chosen under the null hypothesis or (ii) assume different orders for different phenotypic trajectories. The second method, however, is computationally intensive. Even though one can select the polynomial order for different QTL effects, the technical challenge lies on the selection of the order for the time-dependent permanent environmental effects [54]. Thus, we adopted the first method and assumed the same order to fit changes in QTL genetic effects and time-dependent environmental effects. The polynomial regression coefficients determine the shape of the population mean and each genetic effect. Certainly, an optimal mapping strategy is to choose polynomials of different orders to best model each component in the imprinting model, which we leave it for future investigation.

In functional mapping, how to model the correlation structure for repeated measurements is also a challenging problem. In most functional mapping studies, a parametric residual covariance structure such as the autoregressive model with order 1 [AR(1)] is often assumed [55]. However, it is difficult to implement a Bayesian method for this covariance function due to the difficulty to choose an appropriate prior for the autoregressive coefficient. In contrast, the covariance structure described by $\psi^T \sum \psi + I \sigma^2$ is more flexible than the parametric structure because we can actually choose different degrees of polynomial order to fit a covariance structure with a large degree of complexity [31]. Moreover, we can easily sample the covariance matrix $\sum$ from a closed form of marginal posterior distribution. Yap et al. [56] recently proposed a non-parametric method for covariance structure modeling in functional mapping. More work is needed to integrate these techniques into a Bayesian mapping framework to improve mapping power.

When unbalanced data are recorded, we can change time covariate $\psi(t)$ or $\psi$ to $\psi_k(t)$ or $\psi_k$ depending on the $k$th individual, to make the method more general. Also, the multiple iQTL model for developmental traits proposed herein could be treated as a general form of the model for analyzing genomic imprinting of a quantitative trait. For instance, let $\psi = I$ and $\xi_k = 0$ in scale, that is, only one measurement at a fixed time point is taken for each individual, leading to a multiple interacting iQTL model for a univariate quantitative trait; take $\psi$ to be an identity matrix of order $m$, and $\xi_k$ to be a zero vector, resulting in a multiple interacting iQTL model for a multiple quantitative trait. If $\xi_k$ is assigned to be non-zero in the two cases above, this leads to a multiple iQTL model for a single developmental trait or multiple developmental traits. Corresponding Bayesian model selection approaches can be likewise obtained by taking different values for $\psi$ and $\xi_k$. The computational code to implement the proposed method (termed Bafmim) is available upon request.

SUPPLEMENTARY DATA
Supplementary Data are available online at http://bib.oxfordjournals.org/.

Key Points
- We developed a Bayesian method for mapping iQTL underlying dynamic/developmental traits.
- A Bayesian model selection strategy for longitudinal traits was proposed to infer iQTLs with different imprinting patterns and mechanisms as discussed in Cheverud et al. [13].
- The method can do multiple iQTL inference and outperforms maximum likelihood-based single iQTL mapping methods for developmental traits in terms of power to detect iQTLs.
- The method was applied to identify iQTLs responsible for growth trajectory of body weight in mice. The pattern of imprinting for the identified iQTLs was further dissected.

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References


44. Ball RD. Bayesian methods for quantitative trait loci mapping based on model selection: approximate analysis using the Bayesian information criterion. *Genetics* 2001;159:1351–64.


APPENDIX 1

The derivation of conditional posteriors

The joint posterior density can be expressed as

\[ p(\theta | y) = p(y | \theta)p(\theta) = p(y | \mu)p(\mu)p(\gamma)p(\xi)p(\Sigma)p(\lambda), \]

where \( \theta = (\mu, \beta, \gamma, \xi, \Sigma, \lambda) \) with \( \beta = (\beta_j) \), \( \gamma = \{ \gamma_j, \lambda_j \} \) for \( j = 1, 2, \ldots, L \) and \( \xi = \{ \xi_k \} \) for \( k = 1, 2, \ldots, n \).

The full conditional posterior distributions of all parameters can be derived from the joint posterior density by fixing other parameters as constants.

Given the other parameters, the full conditional posterior distribution of \( \mu \) is derived as a multivariate normal with mean

\[ (n\Sigma^{-1}\psi^T)^{-1}\Sigma^{-1}\sum_{i=1}^{n}(y_i - U_i + \psi^T\mu) \]

and covariance matrix \( (n\Sigma^{-1}\psi^T)^{-1} \).

The full conditional posterior distribution of \( \beta_j \) is also normal, whose mean is

\[ \beta_j = \left(1 + \frac{1}{w}\right)\sum_{k=1}^{n}x_{kj}^2\psi V^{-1}\psi^T^{-1}\sum_{k=1}^{n}x_{kj}(y_k - U_k + \gamma_j x_k \psi^T\beta_j) \]

and covariance matrix \( \Sigma^{-1}\psi V^{-1}\psi^T^{-1} \).

The full conditional posterior distribution of \( \xi_k \) is normal, if \( \psi \) follows a normal distribution with mean

\[ \psi = \psi^{-1} \sum_{k=1}^{n}x_{kj}y_k \psi \]

and covariance matrix \( \Sigma = \Sigma^{-1}\psi V^{-1}\psi^T \).

The full conditional posterior distribution of \( \Sigma \) follows a normal Wishart distribution

\[ \Gamma \left( \nu_v + n, \sum_{k=1}^{n}z_{kj}^Tz_{kj} + \psi^T \psi \right). \]

The full conditional posterior distribution for residual variance \( \sigma^2 \) corresponds to a scaled inverse \( \chi^2 \) distribution with parameters \( \nu_v + n \) and \( (\nu_v + n)x_k + \sum_{k=1}^{n}e_k^T e_k \), where \( e_k = y_k - U_k - \psi^T\xi_k \).

The full conditional posterior for \( \gamma_j \) for \( j = 1, 2, \ldots, L \) can be constructed as a Bernoulli distribution. To improve sampling efficiency, the Metropolis–Hastings algorithm is applied with the acceptance rate given by

\[ \alpha = \frac{p(y | \gamma_j = 1, \gamma_{-j}, \beta_{-j})}{p(y | \gamma_j = 0, \gamma_{-j}, \beta_{-j})} \frac{1 - w}{w} \]

where \( \gamma_{-j} \) means all the elements of \( \gamma \) except for \( \gamma_j \) and \( \beta_{-j} \) represents all the elements of \( \beta \) except for \( \beta_j \). The full conditional posterior distribution for QTL positions does not have a closed form, and the Metropolis–Hastings algorithm will be used to sample QTL positions. Each locus is sampled from a variable interval size whose boundaries are the positions of adjoining QTLs. We restricted the minimal distance between two QTLs to be 10cM at least. A new position \( \lambda_j^* \) for \( j = 1, 2, \ldots, L \) is sampled from two placed loci on the left and right of existing position \( \lambda_j^0 \) with a uniform distribution, then the new position is accepted with a probability of \( \min(1, \alpha) \) with

\[ \alpha = \frac{\prod_{j=1}^{L} p(y_k | \lambda_j^*, \theta_{-j})}{\prod_{j=1}^{L} p(y_k | \lambda_j^0, \theta_{-j})} \]