# Efficient Control of Population Structure in Model Organism Association Mapping

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#### ABSTRACT

Genomewide association mapping in model organisms such as inbred mouse strains is a promising approach for the identification of risk factors related to human diseases. However, genetic association studies in inbred model organisms are confronted by the problem of complex population structure among strains. This induces inflated false positive rates, which cannot be corrected using standard approaches applied in human association studies such as genomic control or structured association. Recent studies demonstrated that mixed models successfully correct for the genetic relatedness in association mapping in maize and Arabidopsis panel data sets. However, the currently available mixed-model methods suffer from computational inefficiency. In this article, we propose a new method, efficient mixed-model association (EMMA), which corrects for population structure and genetic relatedness in model organism association mapping. Our method takes advantage of the specific nature of the optimization problem in applying mixed models for association mapping, which allows us to substantially increase the computational speed and reliability of the results. We applied EMMA to in silico whole-genome association mapping of inbred mouse strains involving hundreds of thousands of SNPs, in addition to Arabidopsis and maize data sets. We also performed extensive simulation studies to estimate the statistical power of EMMA under various SNP effects, varying degrees of population structure, and differing numbers of multiple measurements per strain. Despite the limited power of inbred mouse association mapping due to the limited number of available inbred strains, we are able to identify significantly associated SNPs, which fall into known QTL or genes identified through previous studies while avoiding an inflation of false positives. An R package implementation and webserver of our EMMA method are publicly available.

TITH the recent development of high-throughput genotyping technologies, genetic variation in many model organisms such as mice, Arabidopsis, and maize is being discovered on a genomewide scale (JANDER et al. 2002; Pletcher et al. 2004; Flint-Garcia et al. 2005; Frazer et al. 2007). Genomewide association mapping in model organisms has great potential to identify risk factors for complex traits related to human diseases. Despite the disadvantage that direct inferences from model organisms are not always applicable to human traits, model organism association mapping is potentially more powerful than human association mapping because it is possible to reduce the effect of environmental factors by replicating phenotype measurements in genetically identical organisms (Belknap 1998). In addition, it is often easier and more cost effective to verify associated signals in model organisms than in

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human subjects. Moreover, many ongoing genotyping and phenotyping projects in model organisms such as the Mouse Phenome Database (MPD) (http://www.jax.org/phenome), the Mouse HapMap project (http://www.broad.mit.edu/personal/claire/MouseHapMap), and the Perlegen/NIEHS resequencing project (http://mouse.perlegen.com) (Frazer et al. 2007) provide publicly available resources to perform in silico mapping of complex traits in model organisms (Peter et al. 2007).

However, genetic association studies in inbred model organisms are confronted by the problem of inflated false positive rates due to population structure and genetic relatedness among inbred strains caused by the complex genealogical history of most model organism strains. Conventional statistical tests of independence between a genetic marker and a phenotype are prone to spurious associations because the marker and the phenotype are likely to be correlated due to population structure that violates the independence assumption under the null hypothesis. Recent association- or linkagemapping studies in model organisms attempt to avoid

inflated false positive rates by designing the studies using recombinant inbred lines generated from a small number of parental strains (Bystrykh *et al.* 2005; Zou *et al.* 2005). However, these studies are limited by the variation present in the parental strains and have long regions between recombinations due to relatively few generations between the recombinant inbred strains and the parental strains. Traditional QTL mapping using  $F_2$  or backcross suffers from the same problem in fine-resolution mapping in addition to expensive genotyping cost (Belknap 1998; Flint *et al.* 2005).

An alternative approach to reduce the inflation of false positives is to apply a statistical test that corrects for the bias due to population structure or genetic relatedness. The most widely used methods to reduce such bias in human association mapping are genomic control (DEVLIN and ROEDER 1999), structured association (Pritchard et al. 2000), and principal component analysis (Patterson et al. 2006; Price et al. 2006). However, these methods are inadequate in the case of model organism association mapping. Genomic control suffers from weak power when the effect of population structure is large as in model organisms (PRICE et al. 2006; Yu et al. 2006). Structured association or principal component analysis, which assumes a small number of ancestral populations and admixture, only partially captures the multiple levels of population structure and genetic relatedness in model organisms (ARANZANA et al. 2005; Yu et al. 2006; Zhao et al. 2007). Recently, it has been suggested that linear mixed models can effectively correct for population structure in the association mapping of quantitative traits (Yu et al. 2006). Linear mixed models incorporate pairwise genetic relatedness between every pair of individuals in the statistical model directly, reflecting that the phenotypes of two genetically similar individuals are more likely to be correlated than genetically dissimilar individuals. Applications of mixed models to association mapping in maize, Arabidopsis, and potato panels demonstrate that mixed models obtain fewer false positives and higher power than previous methods including genomic control, structured association, and principal component analysis (Yu et al. 2006; Malosetti et al. 2007; Zhao et al. 2007).

Although mixed models can effectively capture statistical confounding due to population structure, the currently available implementations have several limitations in the context of model organism association mapping. First, the variance components numerically estimated by various hill-climbing approaches such as the Nelder–Mead simplex algorithm (Nelder and Mead 1965; Graser *et al.* 1987; Meyer 1989), the EM algorithm (Smith 1990), and the Newton–Raphson algorithm (Lindstrom and Bates 1988; Gilmour *et al.* 1995; Johnson and Thompson 1995) provide only a locally optimal solution, which may cause the statistical inferences based on these estimates to be inaccurate.

Second, the computational cost of the numerical optimization procedure is substantial, requiring a large number of computationally expensive matrix operations at each iteration. Computational considerations are important when large data sets are to be tested. For example, the association mapping with maize panels consisting of hundreds of SNPs over hundreds of strains takes hours for a single run with currently available implementations such as TASSEL (Yu et al. 2006) or SAS (SAS INSTITUTE 2004). A microarray data set tested for genomewide association mapping between thousands of transcripts and tens of thousands of markers would take several years of CPU time. Third, when inferring the genetic variance component referred to as the kinship matrix, the importance of a mathematically correct form of kinship matrix estimation is often overlooked. For example, Yu et al. (2006) proposed to infer a kinship matrix using SPAGeDi software, setting negative kinship coefficients to zero. Such a kinship matrix may not be positive semidefinite and thus not be a valid form of variance component. Using a nonpositive semidefinite kinship matrix generates ill-defined likelihood for a subset of parameter space in the estimation of the variance component.

In this article, we propose a new method, efficient mixed-model association (EMMA), which corrects for population structure and genetic relatedness in model organism association mapping. Our method takes advantage of the specific nature of the optimization problem in applying mixed models for association mapping, which allows us to substantially increase computational speed by orders of magnitude and improve the reliability of results by achieving near global optimization. Our method improves the efficiency of the mixedmodel method by enabling us to perform statistical tests with single-dimensional optimization. Our method's efficiency is further increased by avoiding redundant computationally expensive matrix operation at each iteration in the computation of likelihood function by leveraging spectral decomposition, reducing the computational cost of each iteration from cubic to linear complexity. Due to a substantially decreased computational cost of each iteration, it is possible to converge the global optimum of the likelihood in variancecomponent estimation with high confidence by combining grid search and the Newton-Raphson algorithm even though the likelihood function may not be convex. Our method is related to a similar technique developed in a different context of simulating the null distribution of variance-component test statistics (CRAINICEANU and Ruppert 2004).

We show that a simple genetic similarity matrix can serve as a kinship matrix accounting for genetic relatedness as effectively as previously suggested methods while guaranteeing positive semidefiniteness. Our results are consistent with other studies (Zhao *et al.* 2007), which suggests that these simpler kinship matrices reduce the

false positive rate as effectively as or more effectively than the kinship matrices generated by previous methods (Yu et al. 2006). We propose an additional method called phylogenetic control based on the assumption that a phylogenetic tree is a good approximation of the genealogical history of an inbred model organism. In such cases, the phylogenetic tree may be used as a confounding factor, correcting for the complex genetic relations between strains. We show that phylogenetic control can be formulated as a linear mixed model and present an algorithm for inferring the phylogenetic kinship matrix. We show that the phylogenetic kinship matrix is always positive semidefinite and its optimal variance components are unique regardless of the choice of root.

One of the important questions in the design of model organism association-mapping studies is estimating the statistical power for any specific set of inbred strains. We performed a simulation study of the power of our EMMA method to identify causal SNPs both on a genomewide scale and within a smaller region such as a QTL interval. Our results show that with a limited number of genetically diverse strains, such as the currently available panel of inbred mice, it is possible to identify causal loci with a genomewide significance only if the locus explains a large portion of phenotypic variance. However, with more strains, the power of these association studies increases dramatically. Our analysis of statistical power in model organism association mapping demonstrates the dramatic increase in power using multiple measurements of phenotypes from multiple animals for each strain. Study designs that do not replicate phenotype measurements and analysis methods that do not take individual measurements into account suffer a significant decrease in statistical power.

We applied our EMMA method to association mappings of various inbred model organisms. First, we verified that EMMA gives almost identical results to other widely used implementations using the maize panel data sets (Yu et al. 2006). In terms of computational time, EMMA is shown to be orders of magnitude faster than the previous methods while performing near global optimization. Second, we performed a genomewide association mapping of Arabidopsis flowering-time phenotypes. Our results are consistent with the recently published results (ZHAO et al. 2007), reducing most of the inflated false positives. Finally, we used our EMMA method to perform a whole-genome association-mapping study of inbred mouse strains. We analyzed nearly 140,000 mouse HapMap SNPs over 48 strains and three quantitative phenotypes, liver weight, body weight, and saccharin preference, with QTL identified by previous studies. We identified significant associations for the three phenotypes while our results show a significant reduction in the inflation of false positives. Interestingly, many of the significantly associated SNPs fall into the known QTL, suggesting the results are likely to be true associations.

An R package implementation of EMMA and the webserver containing the mouse association results are publicly available online at http://mouse.cs.ucla.edu/emma.

## MATERIALS AND METHODS

Genotypes and phenotypes: Genotypes, phenotypes, SPAGeDi-based kinship matrix, and the STRUCTURE outputs from 277 maize strains across 553 SNPs as described in Yu et al. (2006) were downloaded from the Buckler lab web site (http://www.maizegenetics.net). The Arabidopsis genotypes and phenotypes and the output from STRUCTURE were obtained from the published data sets (Aranzana et al. 2005; Nordborg et al. 2005). The 13,416 nonsingleton Arabidopsis SNPs with no more than 10% of genotype calls missing were tested for association after imputing the missing alleles using HAP (Halperin and Eskin 2004). The flowering-time phenotypes over 95 strains were log transformed to fit to a normal distribution.

For inbred mouse association mapping, the Broad mouse HapMap SNP sets were obtained from the mouse HapMap web site. The 106,040 SNPs that have no more than 10% of genotype calls missing were tested after imputing the missing alleles. The initial body weight (MPD10305) and liver weight phenotypes (MPD2907) were downloaded from Jackson Laboratory MPD (Jackson Laboratory MPD (Jackson Laboratory 2004). They consist of 374 and 308 phenotype measurements over 38 and 34 strains, respectively. The saccharin preference phenotypes consist of 280 phenotype measurements in 24 strains (Reed et al. 2004).

**EMMA:** Suppose that n measurements of a phenotype are collected across t inbred strains. A linear mixed model in model organism association mapping is typically expressed as

$$\mathbf{y} = X\mathbf{\beta} + Z\mathbf{u} + \mathbf{e},\tag{1}$$

where  $\mathbf{y}$  is an  $n \times 1$  vector of observed phenotypes, and X is an  $n \times q$  matrix of fixed effects including mean, SNPs, and other confounding variables.  $\beta$  is a  $q \times 1$  vector representing coefficients of the fixed effects. Z is an  $n \times t$  incidence matrix mapping each observed phenotype to one of t inbred strains.  $\mathbf{u}$  is the random effect of the mixed model with  $\mathrm{Var}(\mathbf{u}) = \sigma_{\mathrm{g}}^2 K$ , where K is the  $t \times t$  kinship matrix inferred from genotypes as described in the following section, and  $\mathbf{e}$  is an  $n \times n$  matrix of residual effect such that  $\mathrm{Var}(\mathbf{e}) = \sigma_{\mathrm{e}}^2 I$ . The overall phenotypic variance—covariance matrix can be represented as  $V = \sigma_{\mathrm{g}}^2 ZKZ' + \sigma_{\mathrm{e}}^2 I$ .

Instead of solving mixed-model equations by obtaining the best linear unbiased prediction (BLUP) of random effects  ${\bf u}$  via Henderson's iterative procedure (Henderson 1984; Arbelbide et al. 2006), we directly estimate the variance components  $\sigma_{\rm g}$  and  $\sigma_{\rm e}$ , maximizing the full likelihood or restricted likelihood that is defined as full likelihood with the fixed effects integrated out (Dempster et al. 1981). The restricted likelihood avoids a downward bias of maximum-likelihood estimates of variance components by taking into account the loss in degrees of freedom associated with fixed effects. Under the null hypothesis, the full log-likelihood and restricted log-likelihood function can be formulated as

$$\begin{split} l_{\mathrm{F}}(\mathbf{y}; \boldsymbol{\beta}, \boldsymbol{\sigma}, \boldsymbol{\delta}) &= \frac{1}{2} \bigg[ -n \, \log(2\pi\sigma^2) - \log|H| \\ &\quad - \frac{1}{\sigma^2} (\mathbf{y} - X\boldsymbol{\beta})' H^{-1} (\mathbf{y} - X\boldsymbol{\beta}) \bigg] \end{aligned} \tag{2} \\ l_{\mathrm{R}}(\mathbf{y}; \boldsymbol{\sigma}, \boldsymbol{\delta}) &= l_{\mathrm{F}}(\mathbf{y}; \hat{\boldsymbol{\beta}}, \sigma^2, \boldsymbol{\delta}) \\ &\quad + \frac{1}{2} \big[ q \log(2\pi\sigma^2) + \log|X'X| - \log|X'H^{-1}X| \big] \end{aligned} \tag{3}$$

(Welham and Thompson 1997), where  $\sigma = \sigma_g$  and  $H = \sigma^{-1}V = ZKZ' + \delta I$  is a function of  $\delta$ , defined as  $\delta = \sigma_e^2/\sigma_g^2$ .

The full-likelihood function is maximized when  $\beta$  is  $\hat{\beta} = (X'H^{-1}X)^{-1}X'H^{-1}\mathbf{y}$ , and the optimal variance component is  $\hat{\sigma}_{\mathbf{r}}^2 = R/n$  for full likelihood and  $\hat{\sigma}_{\mathbf{r}}^2 = R/(n-q)$  for restricted likelihood, where  $R = (\mathbf{y} - X\hat{\beta})'H^{-1}(\mathbf{y} - X\hat{\beta})$  is a function of  $\delta$  as well.

Using spectral decomposition, it is possible to find  $\xi_{\it i}$  and  $\lambda_{\it s}$  such that

$$H = ZKZ' + \delta I = U_{\rm F} \operatorname{diag}(\xi_1 + \delta, \dots, \xi_n + \delta) U_{\rm F}'$$
(4)

$$SHS = S(ZKZ' + \delta I)S$$

$$= [U_R W_R] \operatorname{diag}(\lambda_1 + \delta, \dots, \lambda_{n-q} + \delta, 0, \dots, 0)[U_R W_R]'$$

$$= U_R \operatorname{diag}(\lambda_1 + \delta, \dots, \lambda_{n-q} + \delta) U_R',$$
(5)

where  $S = I - X(X'X)^{-1}X'$ ,  $U_{\rm F}$  is  $n \times n$ , and  $U_{\rm R}$  is an  $n \times (n-q)$  eigenvector matrix corresponding to the nonzero eigenvalues.  $W_{\rm R}$  is an  $n \times q$  eigenvector matrix corresponding to zero eigenvalues. As shown in the APPENDIX, our decomposition satisfies the properties of the decomposition suggested by previous studies (Patterson and Thompson 1971). It should be noted that  $U_{\rm F}$  and  $U_{\rm R}$  are independent of  $\delta$ . Let  $U'_{\rm R} \mathbf{y} = [\eta_1, \eta_2, \cdots, \eta_{n-q}]'$ ; then finding maximum-likelihood (ML) or restricted maximum-likelihood (REML) estimates is equivalent to optimizing the following functions with respect to  $\delta$ :

$$f_{F}(\delta) = l_{F}(\mathbf{y}; \hat{\boldsymbol{\beta}}, \hat{\boldsymbol{\sigma}}, \delta)$$

$$= \frac{1}{2} \left[ n \log \frac{n}{2\pi} - n - n \log \left( \sum_{s=1}^{n-q} \frac{\eta_{s}^{2}}{\lambda_{s} + \delta} \right) - \sum_{i=1}^{n} \log(\xi_{i} + \delta) \right]$$
(6)

$$f_{R}(\delta) = l_{R}(\mathbf{y}; \hat{\sigma}, \delta)$$

$$= \frac{1}{2} \left[ (n-q) \log \frac{n-q}{2\pi} - (n-q) - (n-q) \log \left( \sum_{s=1}^{n-q} \frac{\eta_{s}^{2}}{\lambda_{s} + \delta} \right) - \sum_{s=1}^{n-q} \log(\lambda_{s} + \delta) \right]$$
(7)

(see the APPENDIX for the mathematical details). The derivatives of these functions follow that

$$f_{\rm F}'(\delta) = \frac{n}{2} \cdot \frac{\sum_{s} \eta_s^2 / (\lambda_s + \delta)^2}{\sum_{s} \eta_s^2 / (\lambda_s + \delta)} - \frac{1}{2} \sum_{s} \frac{1}{\xi_s + \delta}$$
(8)

$$f_{R}'(\delta) = \frac{n-q}{2} \cdot \frac{\sum_{s} \eta_{s}^{2} / (\lambda_{s} + \delta)^{2}}{\sum_{s} \eta_{s}^{2} / (\lambda_{s} + \delta)} - \frac{1}{2} \sum_{s} \frac{1}{\lambda_{s} + \delta}.$$
 (9)

It should be noted that the likelihood functions are continuous for all  $\delta > 0$  if and only if all the eigenvalues  $\lambda_s$  are nonnegative. Otherwise, such as in the case of the nonpositive semidefinite kinship matrix, the likelihood would be ill defined for a certain range of  $\delta$ .

The suggested procedure in computing likelihood and its derivatives involves only a linear time vector operation at each iteration once the spectral decomposition is computed. The time complexity of the method is  $O(n^3 + rn)$ , where r is the number of iterations required. The time complexity of standard EM or Newton–Raphson algorithms is  $O(rn^3)$ , and the actual ratio of the running time is much bigger than r because the existing methods typically require a large number of matrix multiplications and inverses at each iteration while EMMA computes spectral decomposition only once. Since the computational cost of each iteration has decreased dramatically, instead of obtaining a locally optimal solution during the

numerical optimization, it is now computationally feasible to perform a grid search combining with the Newton–Raphson algorithm in the single-dimensional parameter space consisting of  $\delta$ , which is the ratio of the environmental random effect to the genetic background effect, to optimize the likelihood globally with high confidence.

Furthermore, when a large number of multiple measurements are phenotyped per strain, *i.e.*,  $n \gg t$ , the execution time can be further reduced using the fact that the nonnegative eigenvalues of ZKZ' and SZKZ'S are the same as those of KZ'Z and KZ'SZ, respectively. Combining this fact with a simple modification of the Gram–Schmidt process greatly reduces the execution time of eigenvalue decomposition, reducing the time complexity into  $O(n^2t + m)$ . When multiple phenotypes are tested such as in expression quantitative trait loci (eQTL) mapping, the spectral decomposition can be reused, and only a square-time matrix–vector multiplication is required for each phenotype. Thus, the time complexity with m different phenotypes is  $O(n^2t + n^2m + rmn)$ , which is much more efficient than  $O(rm^3m)$  achieved by previous approaches.

In the application of our EMMA method to the various data sets presented in this article, the  $\delta$ 's ranged from  $10^{-5}$  (almost pure population structure effect) to 105 (almost pure environmental or residual effect) and are divided evenly into 100 regions in logarithm scale to compute the derivatives of likelihood functions. The global ML or REML is searched for by applying the Newton-Raphson algorithm to all the intervals where the signs of derivatives change and taking the optimal  $\delta$  among all of the stationary points and endpoints. Since the derivatives of both the full- and the restrictedlikelihood function are continuous with nonnegative eigenvalues, such an optimization technique has guaranteed convergence properties as long as the kinship matrix is positive semidefinite. In the following two sections, we describe different methods to infer a kinship matrix K, based on either a genetic similarity matrix or a phylogenetic tree.

Similarity-based kinship matrix: A number of methods for inferring a kinship matrix from a large number of molecular markers have been suggested, including a simple identical-bystate (IBS) allele-sharing matrix, an allele-frequency weighted IBS matrix (Lynch and Ritland 1999), a maximum-likelihood kinship matrix (THOMAS and HILL 2000), and a Monte Carlo simulation-based matrix (WANG 2002). Comparisons of different kinship matrices for explaining genetic differentiation among populations show similar results with small quantitative differences (Nievergelt et al. 2007). Recent studies on the association mapping of Arabidopsis thaliana in a structured population show that a simple IBS allele-sharing matrix effectively corrects for confounding from population structure, even better than more sophisticated methods (Zhao et al. 2007). Although recently suggested estimators of pairwise relatedness have some desirable statistical properties over a simple IBS allele-sharing matrix (CASTEELE et al. 2001), they are not guaranteed to be positive semidefinite.

Here we show that a simple IBS allele-sharing matrix based on the assumption of each SNP or haplotype inducing the same level of small random changes on the phenotype guarantees positive semidefiniteness and convergence if missing alleles are handled appropriately.

Let  $l_{i,j,h} \in \{0, 1\}$  be a binary variable that has a value of 1 only when the genotype (or haplotype) allele at jth locus in the ith strain is  $h \in 1, \dots |\mathcal{H}_j|$ , where  $|\mathcal{H}_j|$  is the total number of alleles at the jth locus. Let  $x_{h,j}$  be random variables independently sampled from  $N(0, \sigma^2)$ ; then the genetic background effect  $u_i$  of strain i can be modeled as an accumulation of small random effects as follows, assuming that  $x_{h,j}$  denote the random genetic effect caused by allele h at the jth locus,

$$u_{i} = \sum_{j} \sum_{h=1}^{\mathcal{H}_{j}} l_{i,j,h} w_{j} x_{h,j}, \tag{10}$$

where  $w_j$  is the weight of each SNP's contribution to the genetic background effect. If each SNP is assumed to have the same level of random effect,  $w_j = 1$  can be assumed. Alternatively,  $w_j$  can be a function of allele frequency or a function depending on the genomic region of the SNP. Let  $|\mathcal{H}| = \max(|\mathcal{H}_j|)$ , and let  $L_h$  be the matrix whose element at (i, j) is  $l_{i,j,h}$ ; then the overall genetic background effect  $\mathbf{u}$  is expressed in the form

$$\mathbf{u} = \sum_{h=1}^{|\mathcal{H}|} L_h W \mathbf{x}_h, \tag{11}$$

where W is a diagonal square matrix with  $w_i$  at the ith diagonal element. Assuming that each  $x_{h,j}$  follows a normal distribution with zero mean and variance of  $\sigma^2$  independently, the variance-covariance matrix of **u** becomes  $Var(\mathbf{u}) = \sigma^2 \sum_h$  $L_h W^2 L'_h$ . Since its  $(i_0, i_1)$ th element  $\sum_h \sum_j w_j^2 l'_{i_0,j,h} l_{i_1,j,h}$  represents the number of shared IBS alleles between the  $i_0$ th and  $i_1$ th strains directly if  $w_i = 1$ ,  $Var(\mathbf{u})$  is equivalent to a weighted IBS allele sharing a kinship matrix with the scaling factor  $\sigma^2$ . It is obvious from Equation 11 that the kinship matrix is positive semidefinite. When missing genotypes exist, we estimate  $l_{i,j,h}$  to be the square root of the probability of the SNP or haplotype allele at the *j*th locus having the allele *h*. This is so the random effect for each allele is assigned probabilistically. We generated genotype similarity of maize, Arabidopsis, and mouse data sets using uniform weight. When a haplotype similarity matrix is used, the haplotype window size resulting in the largest ML estimates is selected as the optimal window size. In the Arabidopsis and mouse association-mapping results of this article, the optimal haplotype window size is set to five in both cases.

Phylogenetic control: Evolutionary biologists have modeled interspecific phenotype distribution using various phylogenetic comparative methods (PCMs) (MARTINS and HANSEN 1997). The correlation structure between phenotypes can be effectively captured with phylogenetic trees, and PCMs have been applied to evolutionary analysis of quantitative traits such as gene expression (Gu 2004; Oakley et al. 2005) or, very recently, to the association mapping of dichotomous phenotypes (Bhattacharya et al. 2007; Carlson et al. 2007). Felsentein's independent contrast (FIC) method (Felsenstein 1985) models the correlation between phenotypes under the assumption of Brownian motion of phenotypic change along the phylogeny. Since random phenotypic changes occur within a species as well, in cases where the phylogenetic tree is a good approximation of genealogical history, it is reasonable to apply PCMs such as the FIC method in modeling the phenotypic variation in model organisms.

We followed Felsenstein's assumption of Brownian phenotypic changes along the phylogeny. Although multiple fluctuating selection may lead to a Brownian motion model (Felsenstein 1981), here we assume a neutral model where phenotypic changes are explained by accumulated random pleiotropic effects by the genetic background to mathematically model Brownian phenotypic changes. Let T be a phylogenetic tree with t leafs and m edges, and let  $\mathbf{z} \in \mathbf{R}^m$  be random variables independently sampled from  $N(0, \sigma_g^2)$ . At each branch i whose length is  $b_i$ , we represent the amount of random phenotypic changes along the branch as  $\sqrt{b_i z_i}$ . Let  $\Psi_i$ denote the set of branches connecting to a leaf node i from the root. Then the accumulated phenotypic changes are equivalent to  $\sum_{e \in \Psi_i} \sqrt{b_e} z_e$ . If  $X\beta$  is the ancestral mean at an arbitrarily chosen root node, then the phenotype values at the leaf nodes are expressed in the form

$$\mathbf{y} = X\mathbf{\beta} + ZE\mathbf{z} + \mathbf{e},\tag{12}$$

where E is a  $t \times m$  matrix whose (i, j) th element is  $\sqrt{b_j}$  if branch j exists in the path from the root to the leaf node i and zero otherwise. The kinship matrix of random effect  $\mathbf{u} = E\mathbf{z}$  is K = EE' and is proportional to its covariance. If the root of the phylogenetic tree changes, E is changed into  $E + \mathbf{1}_t \mathbf{c}^T$ , with  $\mathbf{1}_t \mathbf{a}$  vector of ones and another vector  $\mathbf{c}$ . However, the restricted likelihood does not change because  $SZ\mathbf{1}_t = 0$  always holds.

In our results, we adjusted the genetic distance matrix using the F84 model (KISHINO and HASEGAWA 1989; FELSENSTEIN and CHURCHILL 1996) from the genomewide genotypes and inferred the phylogenetic tree with the Fitch–Margoliash and least-squares distance method (FITCH and MARGOLIASH 1967).

Statistical tests and multiple hypothesis testing: Once the ML or REML variance component  $\hat{V} = \hat{\sigma}_{g}^{2}K + \hat{\sigma}_{e}^{2}I$  is estimated, a general F-statistic testing the null hypothesis  $M\beta = 0$  for an arbitrary full-rank  $p \times q$  matrix M can be constructed as suggested in Kennedy et al. (1992) and Yu et al. (2006),

$$F = \frac{(M\hat{\beta})'(M(X'\hat{V}^{-1}X)^{-1}M')^{-1}(M\hat{\beta})}{p},$$
 (13)

with p numerator degrees of freedom and n-q denominator degrees of freedom. The Satterthwaite degrees of freedom may also be computed, avoiding computationally intensive matrix operations.

The likelihood-ratio test can also be performed on the basis of the estimated ML variance components under different fixed effects. The statistic asymptotically follows a  $\chi_p^2$  distribution unless the estimated variation component meets the boundary of parameter space.

When a large number of correlated SNPs are tested, Bonferroni correction may lead to too conservative type I error control. Alternatively, permutation tests or other multiple hypothesis-testing procedures can be used (Piepho 2001; Storey and Tibshirani 2003). If permutations of simulation-based approaches are applied, the computational cost is much larger but can be reduced by reusing the spectral decomposition in the same way described in the context of multiple phenotypes. For each permuted y, only  $U_R'y = [\eta_1, \eta_2, \cdots, \eta_{n-q}]$  has to be computed again to compute the full or the restricted likelihood in linear time at each iteration. Thus, the computational cost for a cubic-time spectral decomposition at each permutation can be substituted by a square-time matrix–vector multiplication, reducing the overall time complexity from  $O(n^2t+m)$  to  $O(n^2+m)$ .

The variance-component estimation is performed on the basis of REML for the *F*-test, and ML estimations are used for the likelihood-ratio test and the computation of the Bayesian information criterion (BIC). The *P*-values are computed from the asymptotic null distribution.

**Simulation studies:** We performed two simulation studies for analyzing the statistical power of EMMA. The first simulation is similar to those from other mixed-model studies (Yu et al. 2006; Zhao et al. 2007). A fixed effect based on a randomly chosen causal SNP from the genome with minor allele frequency >10% is added to the existing phenotypes, and the statistical power is computed at the causal SNP. At each fixed effect, the simulation study was performed 1000 times to estimate the average power. The variance explained by a SNP is computed assuming that average minor allele frequency of the causal SNP is 0.3.

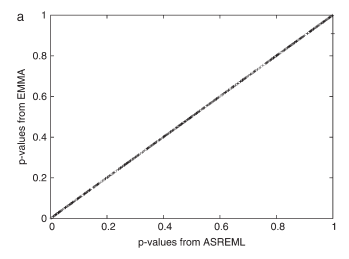
Next, we generated simulated phenotypes sampled from a multivariate normal distribution. A random noise vector is added according to the contribution of genetic background to phenotypes,  $h_{\rm g}^2$ . If  $h_{\rm g}^2$  is the fraction of variance due to genetic

background excluding the SNP effect, then the covariance of the simulated data is simulated as  $\mathrm{Var}(y) = ((n-1)h_\mathrm{g}^2/\mathrm{tr}\ (S_0ZKZ'S_0))K + (1-h_\mathrm{g}^2)I$ , where  $S_0 = I-11'/n$ , where 1 denotes a vector of ones. Similar to the first simulation study, a fixed effect based on a randomly chosen causal SNP is added to the simulated phenotypes and the average power is computed from 1000 independent simulations.

## RESULTS

Comparison with previous methods over maize and **Arabidopsis strains:** We applied our EMMA method to the same maize panel data consisting of 553 SNPs and three phenotypes across 277 diverse inbred lines (FLINT-GARCIA et al. 2005) analyzed with the current mixed-model implementations (Yu et al. 2006). We used the genotype similarity matrix defined in MATERIALS AND METHODS as an additional variance component. Both the SAS and the TASSEL implementations of a unified mixed model (Yu et al. 2006) take nearly 2 hr for a single run over the flowering-time phenotype data set with Intel 2.8-GHz Dual Core CPU, and ASREML, which is known to be more efficient than SAS, takes 20 min (1201 sec) of running time. The execution time of our mixed-model implementation is substantially faster, taking only 2.6 min (157 sec). The comparison of the P-values obtained from the ASREML package and EMMA for flowering-time phenotypes shown in Figure la shows perfect concordance between the methods, suggesting that both methods provide the same accuracy. It should be noted that EMMA is much more efficient in spite of using orders of magnitude more of iterations to find the near global REML estimate using a grid search over the entire parameter space. EMMA also shows high stability of the numerical optimization procedure. In our results, TASSEL and ASREML implementations failed to provide P-values in 4 and 1 SNPs, respectively, of 553 SNPs, possibly due to the instability of the numerical optimization procedure, while EMMA succeeds for all the SNPs over all the data sets covered in this article.

Since the kinship matrix based on SPAGeDi software as suggested by the unified mixed model is not guaranteed to be positive semidefinite, we explore other ways to estimate the variance components due to genetic background. We use a genotype similarity matrix and a phylogenetic control matrix that guarantee positive semidefiniteness. Haplotype similarity matrices are not applicable to this data set due to sparse genotype density. We compared the goodness-of-fit of these kinship matrices in addition to the SPAGeDi-based kinship matrix over three maize phenotypes using the BIC, which provides a measure of how well each model fits the data. Adjusting for the sample size and the number of free parameters, Table 1 shows that the goodness-of-fits of the three kinship matrices based on maximum-likelihood estimates are comparable, while all of them were significantly better than not using a mixed model.



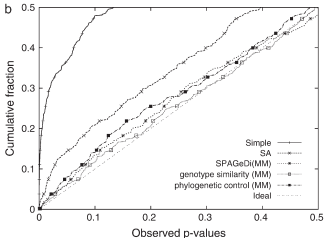


FIGURE 1.—(a) Direct comparison of *P*-values between ASREML and EMMA, computed from 553 SNPs of maize panel data and the flowering-time phenotype using a similarity-based kinship matrix. All *P*-values are almost identical, implying that two methods are almost identical in terms of accuracy. One SNP in ASREML failed to converge during the variance-component estimation while it succeeded in EMMA. (b) Cumulative distribution of *P*-values across different models. Under the assumption that the SNPs are unlinked and there few true SNP associations, the observed *P*-values are expected to be close to the cumulative *P*-values. A large deviation from the expectation implies that the statistical test may cause spurious associations. Simple, a simple *t*-test; SA, structured association; MM, an *F*-test with a mixed model with a specified kinship matrix.

The cumulative *P*-value distribution seen in Figure 1b shows that the three kinship matrices correct for the inflated false positives significantly better than the simple linear model. As illustrated by previous studies, the cumulative distribution of *P*-values is expected to follow that of a uniform distribution with no inflated false positives because only a tiny fraction of all SNPs are expected to be true positives (ARANZANA *et al.* 2005; YU *et al.* 2006; ZHAO *et al.* 2007). The genotype similarity matrix performs slightly better than the other two kinship matrices, especially for small *P*-values. Since

TABLE 1

Goodness-of-fit of different models and kinship matrices in explaining phenotypic variation of maize quantitative traits

		Flowering time		Ear height		Ear diameter	
Method	Kinship matrix	-2 (ML)	BIC	-2 (ML)	BIC	-2 (ML)	BIC
Simple	NA	1632.8	1643.9	2296.0	2307.1	1282.6	1293.5
MM	SPAGeDi	1524.3	1541.0	2237.7	2254.3	1254.2	1270.5
MM	Genotype similarity	1527.5	1544.2	2243.1	2259.8	1266.6	1282.9
MM	Phylogenetic control	1521.6	1538.6	2227.3	2243.9	1248.9	1265.2
SA	NÁ	1525.7	1547.9	2248.9	2271.1	1276.9	1298.7
SA+MM	SPAGeDi	1494.9	1522.7	2220.3	2248.1	1253.6	1280.8
SA+MM	Genotype similarity	1500.9	1528.7	2227.1	2254.9	1266.5	1293.7
SA+MM	Phylogenetic control	1491.6	1519.4	2213.2	2241.0	1248.2	1275.4

Comparison of the maximum likelihood (ML) and the Bayesian information criterion (BIC) of each model with different kinship matrices for maize quantitative traits. The model with the smaller BIC is preferred. Simple, the simple linear model without adjustment for population effect; SA, the model using the output from STRUCTURE as covariates; MM, the mixed model with different kinship matrices. The descriptions of kinship matrices are the same as in the Figure 1 legend.

the simpler kinship matrices show comparable or better goodness-of-fit and false positive reduction results while guaranteeing positive semidefiniteness, we apply only these simple kinship matrices in the following sections.

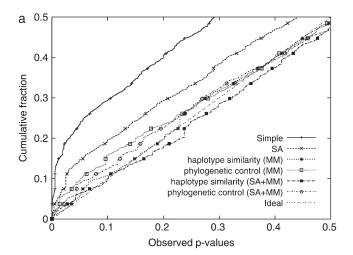
We also applied our EMMA method to perform genomewide association mapping of the flowering-time phenotype in which statistically significant associations are reported in previous studies. The cumulative distribution of P-values across 13,416 nonsingleton SNPs across 95 strains obtained from EMMA is shown in Figure 2a. The cumulative distribution of *P*-values with a haplotype similarity matrix nearly follows the expected distribution, implying that mixed models significantly outperform structured association in eliminating the inflation of false positives for this data set. Phylogenetic control reduces a large portion of inflated false positives, but residual inflation is still observed. Structured association and simple linear regression showed much larger inflation of false positives, consistent with the previous studies. After correction for genetic relatedness, the previously known FRI gene is still found to be significant at a nominal P-value of  $P = 10^{-5}$  across different kinship matrices. Our independent analyses are consistent with the more extensive results of Arabidopsis association mapping recently published (ZHAO et al. 2007).

High-resolution genomewide association mapping in inbred mouse strains: We performed a high-resolution genomewide association-mapping study using our mixed-model method over inbred mouse strains. We used the Broad mouse HapMap SNPs, containing nearly 140,000 SNPs expected to cover most of genetic variation among 48 inbred strains. For phenotypes, we used initial body weight and liver weight phenotypes downloaded from the Jackson Laboratory mouse phenome database (Jackson Laboratory 2004). In addition, we used a saccharin preference phenotype where statistically significant associations were identified in a previous study

(Reed et al. 2004). Among 48 genotyped strains, 38, 34, and 24 strains had phenotype values available for body weight, liver weight, and saccharin preference, respectively. Each phenotype has on average 10 multiple measurements across different individual mice per strain.

The cumulative distributions of observed *P*-values in Figure 2 show that, without correcting for population structure, the rate of false positives is very high. In particular, the body weight phenotype has a substantial inflation of false positives. When our mixed model is used, the inflation of the statistics is significantly reduced in all three phenotypes.

Figure 3 shows genomewide association signals for the three phenotypes. Comparing Figure 3a and 3b, it is obvious that, without correcting for population structure, many false positives are observed at a genomewide level of significance due to inflated P-values. Without correcting for population structure, we were able to identify nearly 6000 SNPs at a nominal P-value of 10<sup>-6</sup> and 283 SNPs with *P*-values  $<10^{-10}$ . However, none are significant after applying the mixed model. This strongly supports that most of the significant associations without correcting for population structure are indeed false positives. Interestingly, although the strongest signals for the body weight after applying the mixed model are not genomewide significant, they are concentrated in the region around 114 Mb in chromosome 8. This region almost exactly falls into the LOD peak of a previously known body weight QTL Bwq3 (Annuciado et al. 2001). The P-value of the most significant locus is  $3.8 \times 10^{-6}$  with the F-test, explaining 49% of the overall phenotypic variance and 39% of the phenotypic variation due to the genetic variance component. Although it is slightly below the genomewide significance threshold with a conservative Bonferroni correction, if utilizing the results from previous QTL studies, the locus can be declared as significant over the region of known body weight QTL.



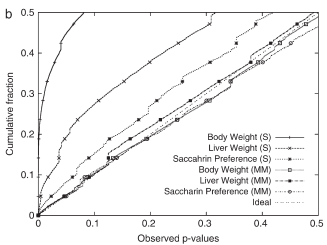


FIGURE 2.—Genomewide cumulative distribution of observed *P*-values between (a) 13,416 Arabidopsis SNPs and flowering-time phenotypes across 95 strains using various models and (b) 106,040 mouse HapMap SNPs and three phenotypes, body weight (374 measurements over 38 strains), liver weight (304 measurements over 34 strains), and saccharin preference (280 measurements across 24 strains). S or Simple, a simple *t*-test; SA, structured association; MM, an *F*-test with a mixed model with a haplotype similarity kinship matrix; SA+MM, the unified mixed model using the output of STRUCTURE as additional fixed effects.

For the liver weight phenotype, we identified a genomewide significant association around the region of 34.5 Mb in chromosome 2. This falls into a previously known liver weight QTL Lvrq1 (Rocha et~al.~2004). The region also contains many potentially relevant QTL such as organ weight (Orgwq2) (Leamy et~al.~2002), spleen weight (Sp1q1) (Rocha et~al.~2004), heart weight (Hrtq1) (Rocha et~al.~2004), and lean body mass (Lbm1) (Masinde et~al.~2002). The pointwise P-value of the most significant SNP was  $1.2 \times 10^{-9}$ , which explains 59% of the genetic variance component. When comparing the genomewide P-values between the simple t-test and mixed models in Figure 3, c and d, we observe that the inflation of t-values is reduced, but the signals are even more significant around the significant SNP at chromo-

some 2. This demonstrates that mixed-model association mapping can not only reduce the inflated false positives, but also reveal significant associations that have remained unidentified using conventional statistical methods in the case when the associated SNP is not highly correlated with population structure.

For the saccharin preference phenotype, we were able to identify a SNP 30 kb away from the Tas1r3 gene that is perfectly correlated with the SNP previously reported to have significant association with the phenotype (Reed et al. 2004). It explains 51% of the genetic variance component, with a P-value of  $1.0 \times 10^{-5}$ . The SNP is neither genomewide significant nor the most significant. We believe this is due to the limited power of the study with a small number of strains.

Power of inbred model organism association mapping: We evaluated the statistical power of association mapping of inbred model organisms in two different ways. First, we simulated an additive effect of a causal SNP over the existing phenotypes for mouse, Arabidopsis, and maize strains, similar to previous studies. Such simulation studies evaluate the SNP effect on the power maintaining the existing correlation structure of phenotypes. However, they do not change the effect of the genetic background or the number of multiple measurements, and no random variable other than the SNP is involved in the power simulation. As an alternative model-driven method for simulation studies, we generated simulated phenotypes randomly sampled from a multivariate normal distribution with various effects of population structure on the phenotypic variation. A SNP effect is simulated on the randomly generated samples, and the statistical power is evaluated. In this way, we can simulate not only the SNP effect but also different genetic backgrounds and different numbers of replicated measurements. We believe that our simulation analysis provide a more extensive understanding of the statistical power of association mapping with model organisms.

Figure 4 shows the statistical power with respect to the additive SNP effect on the Arabidopsis and maize flowering-time phenotypes and three inbred mouse phenotypes used in this article. The maize panel data set consisting of 277 strains has high statistical power, achieving 80% power with a SNP effect explaining 5% of phenotypic variation. Genomewide significance can also be achieved with high power with 10% of SNP effects. For the Arabidopsis data set consisting of 95 strains, the statistical power is decreased, and roughly twice the SNP effect would be needed compared to the maize panels to achieve the same statistical power. For the inbred mouse phenotypes, genomewide power is achievable only when the SNP explains a very large portion of phenotypic variance. In our results, the plausible significant associations explained >35% of the phenotypic variance. The power to achieve genomewide power is largely dependent on the number of

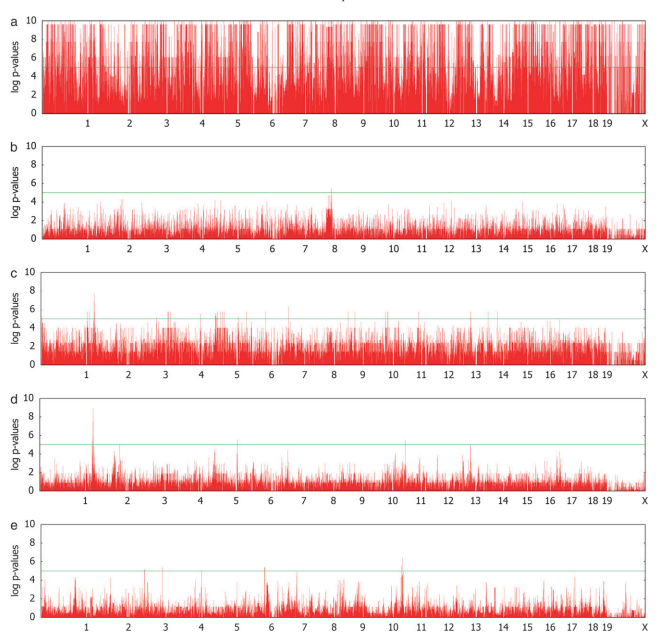


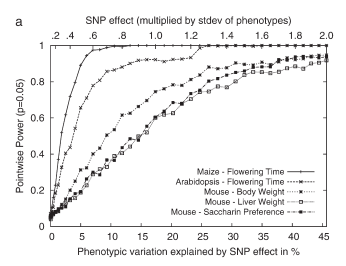
FIGURE 3.—Genomewide scans for association with initial body weight, liver weight, and saccharin preference, using simple *t*-tests and *F*-tests with mixed models, on the basis of a kinship inferred from haplotype similarities.

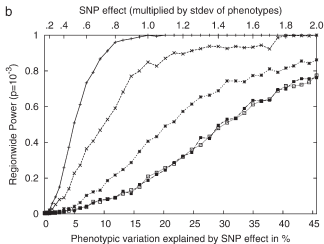
available strains. Table 2 summarizes the most plausible associations in these three phenotypes.

Next, we performed simulation studies by sampling phenotypes from multivariate normal distribution on the basis of the kinship matrix of 48 inbred mouse strains with different effects of genetic background due to population structure. We observed a significant increase of power when multiple measurements are used. Figure 5a shows the effect of multiple measurements on the statistical power when the variance from the genetic component and the residual component are the same. It suggests that using just a single measurement per strain may result in a significant decrease in power. Even though multiple measurements are used, if only the phe-

notypic mean is used in the analysis and the individual measurements are not taken into account, the statistical power would decrease significantly. Comparing Figure 5b with 5a clearly shows the advantage of using individual measurements over the phenotypic mean in the statistical analysis. It shows that the statistical power may differ by up to a factor of two between the two methods. Other mixed-model association-mapping studies use only the mean values in their analysis, not fully utilizing the potential of individual measurements.

Figure 5c shows that a large relative effect from genetic background reduces the statistical power. As the genetic background contributes a larger portion of phenotypic variance, the within-strain variance becomes smaller than





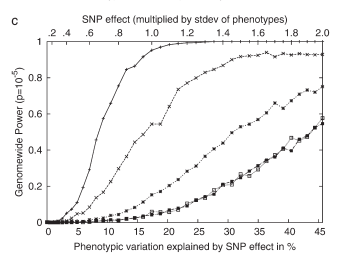


FIGURE 4.—Comparisons of the statistical power of the EMMA method across three different inbred mouse phenotypes and flowering time of Arabidopsis and maize, by randomly selecting causal SNPs across the genomewide SNPs. (a) Pointwise power denotes the power to identify causal SNPs at a nominal *P*-value of 0.05. (b) Regionwide power assumes 50 hypothetical tagSNPs in a genomic region. With 20 kb between tagSNPs, the genomic region covers up to 1 Mb. (c) Genomewide power is the power to achieve genomewide significance using the *P*-value threshold 10<sup>-5</sup>, which is conser-

the between-strain variance, and this limits the contribution of multiple measurements to the statistical power (Belknap 1998). For example, in an extreme case, when  $h_{\rm g}^2=1$ , the residual variance is zero and the replicated measurement does not increase the power since there is no variability of phenotype within strains.

Figure 5d shows more clearly the effect of genetic background and multiple measurements at a glance. When a SNP explains a fairly large fraction (17%) of phenotypic variance, the genomewide significance level can be achieved with high power only when the phenotype has very small population structure effect and the number of replicates is large. As the effect from genetic background becomes larger, the advantage of using multiple measurements decreases significantly.

### DISCUSSION

In this article, we proposed an efficient statistical method to perform association mapping with structured samples on the basis of a linear mixed model. Our results with maize and Arabidopsis panels show that EMMA robustly reduces the inflated false positives under a structured population similar to currently available mixed-model implementations. The accuracy and stability of the numerical optimization in EMMA is greater than others due to global optimization of likelihood function and guaranteed convergence properties with a smaller search space. Our presentation of the EMMA method is focused on a particular case of a mixed model where two variance components are involved because this is the typical model that previous studies assume, and it is straightforward to correct population structure via one kinship matrix inferred from genomewide markers.

The computational efficiency of EMMA is orders of magnitude greater than that of other widely used implementations. When multiple measurements per strain are used across different individuals, the relative efficiency is further increased. This is of a great importance when the computational cost may be a bottleneck in the statistical analysis of high-throughput data such as genomewide gene expressions. For example, the single run of genomewide association mapping of mouse body weight phenotypes with multiple measurements would take up to a month of CPU time with other implementations, while EMMA takes only a single CPU hour. When hundreds and thousands of phenotypes are available such as in the analysis of whole-genome expression data, the computational cost of previous implementations is prohibitive even with high-performance

vative compared to the permutation-based genomewide significance thresholds using the original phenotypes. The phenotypic variation explained by SNP effect is computed assuming a minor allele frequency (MAF) of 0.3.

TABLE 2
List of plausible associations in the mouse association mapping

			<i>P</i> -value		Variance explained (%)				
Phenotype	Chromosome	Position	F-test	LR test	Overall	Genetic	Alleles	MAF	Notes
Body weight	8	113,588,970	$3.9 \times 10^{-6}$	$1.9 \times 10^{-5}$	49.0	38.7	A/C	0.27	300 kb from the LOD peak of <i>Bwq3</i> QTL
Liver weight	2	34,499,435	$1.2 \times 10^{-9}$	$1.4 \times 10^{-7}$	39.1	58.6	G/C	0.50	Genomewide significant, within <i>Lvrq1</i> QTL
Saccharin preference	4	154,883,600	$1.0 \times 10^{-5}$	$7.5 \times 10^{-5}$	35.9	50.6	G/A	0.31	30 kb from <i>Tas1r3</i> gene

MAF, minor allele frequency.

computing. It should be noted that there are other techniques developed for improving computational efficiency of the numerical estimation in a more general context of linear mixed models such as average information REML (GILMOUR *et al.* 1995), but these techniques would not provide us with the same improvements on the efficiency of each iterative procedure.

Our results of inbred mouse association mapping show the potential and limitations of genomewide inbred mouse association studies. It is remarkable that we were able to identify significant associations at a genomewide level without inflation of false positives, under the limited statistical power of the method. Although there is a possibility that residual confounding still remains with mixed-model association, we believe that the most significant SNP associated with liver weight is likely to be a true positive because it explains a large portion of phenotypic variations between the strains beyond genetic background effect so that the conservative Bonferroniadjusted P-value still remains significant. The SNP associated with body weight looks also plausible, but it could possibly be due to residual confounding that is not completely captured by a kinship matrix. Likewise, other significant associations can possibly be due to residual confounding not captured by the kinship matrix, so the identified associations must be verified through independent analysis.

In a more general context of association mapping that requires the use of multiple variance components, the computational advantages of EMMA are not applicable since EMMA can effectively solve a model only with one correlated variance component. For example, when allowing heterozygous alleles for outbred individuals, the full model typically takes both additive and dominant variance components in the linear mixed model (Lynch and Ritland 1999; Arbelbide et al. 2006). Likewise, if strain-specific environmental random effects or other additional random effects are to be considered such as in plant association mapping, multiple variance components need to be used. In such cases where EMMA is not directly applicable, computational bottlenecks may be the biggest obstacles in analyzing large amounts of data such as genomewide

expression profiles. EMMA can still be applied in this case if a reasonable approximation is combined with other standard mixed-model methods taking multiple variance components. Under the null hypothesis, it is possible to estimate the ratio between multiple variance components using the full model, and EMMA can be applied under an alternative hypothesis assuming that the ratio between variance components is preserved. Since variance-component estimation under the null hypothesis needs to be done once across a larger number of alternative hypotheses for each marker, such an approximation procedure provides a large amount of computational efficiency essentially equivalent to EMMA with one variance component. Although the approximated test may lose statistical power slightly, the false positive rates would not be inflated.

There have been several genomewide associationmapping studies with inbred mouse strains. To the best of our knowledge, our results are the first wholegenome association mapping of inbred mice that takes the genetic relatedness into account via a statistical method supported by asymptotic theory. Previous studies either do not take the population structure into account (Cervino et al. 2007) or apply heuristics to reduce the confounding effect from population structure. For example, the weighted version of the *F*-statistic (PLETCHER et al. 2004) does not follow the asymptotic null distribution. Redefining the significance level on the basis of the empirical null distribution given the heritability parameter (Liu et al. 2006) or the weighted permutation (McClurg et al. 2007) rescales the P-values only similar to genomic control and will suffer from a lack of power as the genetic background effect becomes larger.

Our power simulation studies provide assistance to the design of the association study under the effect of population structure. Multiple factors are involved in determining the condition for identifying a locus, and it cannot be represented simply by a single value such as phenotypic variance explained by the SNP. Our results show the importance of multiple measurements of phenotypes from multiple animals for each strain and of directly using the individual measurements in the

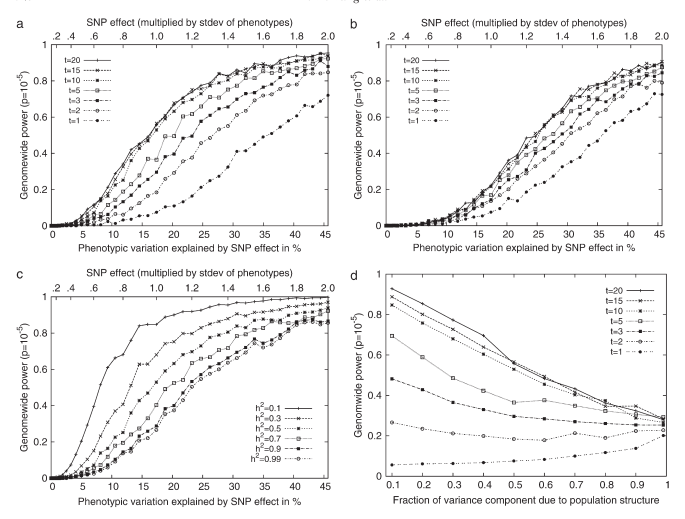


FIGURE 5.—Comparisons of the genomewide power of the EMMA method applied to inbred mouse associations for simulated phenotypes with various SNP effects, genetic background effects, and numbers of multiple measurements. The significance threshold is  $P=10^{-5}$ . t is the number of multiple measurements per strain, and  $h_{\rm g}^2$  is the fraction of the variance explained by genetic background among overall phenotypic variances when the SNP effect is not added. (a) With  $h_{\rm g}^2=0.5$ , varying  $\beta$  and t. (b) The same as a, using the mean phenotype value per strain instead of individual measurements. (c) With 10 multiple measurements per strain, varying  $\beta$  and  $h_{\rm g}^2$ . (d) With  $\beta=\sigma$ , varying t and t. The effect of population structure is varied by changing the ratio of two variance components, and the numbers of multiple measurements are simulated with (a) 10 measurements and (b) a single measurement per strain.

statistics for association mapping. Taking individual measurements into account within the association mapping is much more computationally intensive. EMMA provides a method for efficiently handling individual measurements. In addition, our results also demonstrate the effect of genetic background on the statistical power. As the population structure explains larger phenotypic variance, the power using multiple measurements becomes lower.

Our results show that phylogenetic control can control for population structure as effectively as the linear mixed model based on the genetic similarity matrix in some data sets despite the limited ability of the model to represent complex genetic relatedness. Since genetic similarity matrices are better models when accounting for recombination and hybridization, and also are easier to compute, phylogenetic control is not pre-

ferred in association mapping in model organisms. However, it is possible to compute the likelihood of the phylogenetic control model in linear time (Felsenstein 1985), and this may be useful when a very large number of individuals are to be tested.

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## APPENDIX: DERIVATION OF RESTRICTED LIKELIHOOD AND ITS DERIVATIVES

A derivation of Equations 6 and 7 from Equations 2 and 3 is presented in Patterson and Thompson (1971) and Harville (1974). However, their derivation is not straightforward, and it needs to be clarified how exactly it is related to spectral decomposition. Here we describe a more detailed description of obtaining Equations 6 and 7.

Plugging in the optimal parameters  $\hat{\beta}$  and  $\hat{\sigma}_F = R/n$  in Equation 2, it follows that

$$f_{\mathrm{F}}(\delta) = l_{\mathrm{F}}(\mathbf{y}; \hat{\boldsymbol{\beta}}, \hat{\boldsymbol{\sigma}}, \delta) = \frac{1}{2} \left[ -n \log \frac{2\pi R}{n} - \log |H| - n \right]. \tag{A1}$$

From Equation 4, it is straightforward that  $\log |H| = \sum_{i=1}^{n} \log(\xi_i + \delta)$ . And R can be rewritten as follows:

$$R = (\mathbf{y} - X\hat{\boldsymbol{\beta}})'H^{-1}(\mathbf{y} - X\hat{\boldsymbol{\beta}})$$
(A2)

$$= \mathbf{y}' (I - X(X'H^{-1}X)^{-1}X'H^{-1})'H^{-1}(I - X(X'H^{-1}X)^{-1}X'H^{-1})\mathbf{y}$$
 (A3)

$$= \mathbf{y}' P' H^{-1} P_{\mathbf{y}}, \tag{A4}$$

where  $P = I - X(X'H^{-1}X)^{-1}X'H^{-1}$ .

It is straightforward to show that

$$(SHS)(P'H^{-1}P)(SHS) = SHS$$
(A5)

$$(P'H^{-1}P)(SHS)(P'H^{-1}P) = P'H^{-1}P,$$
(A6)

using the fact PS = S and SP = S. Consequently,

$$P'H^{-1}P = (SHS)^{+} = U_{R}\operatorname{diag}\left[(\lambda_{s} + \delta)^{-1}\right]U_{R}', \tag{A7}$$

where  $(\cdot)^+$  denotes the pseudo-inverse of a matrix. Therefore, it follows that

$$R = \mathbf{y}'(P'H^{-1}P)\mathbf{y} \tag{A8}$$

$$= (U_{\mathsf{R}}'\mathbf{y})' \operatorname{diag}[(\lambda_{\mathsf{s}} + \delta)^{-1}](U_{\mathsf{R}}'\mathbf{y}) \tag{A9}$$

$$=\sum_{s=1}^{n-q} \frac{\eta_s^2}{\lambda_s + \delta}.\tag{A10}$$

From Equations A1 and A10, it follows that

$$f_{\rm F}(\delta) = \frac{1}{2} \left[ n \log \frac{n}{2\pi e} - n \log \left( \sum_{s=1}^{n-q} \frac{\eta_s^2}{\lambda_s + \delta} \right) - \sum_{i=1}^n \log(\xi_i + \delta) \right]. \tag{A11}$$

The restricted likelihood of y is equivalent to computing the likelihood of Ay where S = AA' and A'A = I:

$$(SHS)(SHS)^{+} = (SHS)(P'H^{-1}P) = SHP'H^{-1}P = SP = S$$
 (A12)

(PATTERSON and THOMPSON 1971; HARVILLE 1974). On the other hand,

$$(SHS)(SHS)^{+} = (U_{R}\operatorname{diag}(\lambda_{s} + \delta)U_{R}')(U_{R}\operatorname{diag}[(\lambda_{s} + \delta)^{-1}]U_{R}') = U_{R}U_{R}'. \tag{A13}$$

Accordingly,  $U_R U_R' = S$  and  $U_R' U_R = I$  hold, and the restricted likelihood of  $\mathbf{y}$  is equivalent to the likelihood of  $U_R' \mathbf{y} \sim N(0, \sigma^2 \operatorname{diag}(\lambda_s + \delta))$ . By plugging in  $\hat{\sigma}_R^2$  to  $\sigma^2$ , it immediately follows that

$$f_{R}(\delta) = \frac{1}{2} \left[ (n-q) \log \frac{n-q}{2\pi e} - (n-q) \log \left( \sum_{s=1}^{n-q} \frac{\eta_{s}^{2}}{\lambda_{s} + \delta} \right) - \sum_{s=1}^{n-q} \log(\lambda_{s} + \delta) \right]. \tag{A14}$$