

Making multiple-locus association mapping on a genome-wide scale routine

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Since the earliest of genome-wide association studies, a key shortcoming in how their data have been analysed has persisted. The strength of association between a marker locus and trait is measured for each locus separately, on a locus-by-locus basis. Multiple-locus methods that map multiple locus-trait associations simultaneously have been available for some time. However, they have attracted little attention. They can be demanding, computationally, and their results are not always easy to interpret. Yet, it is widely accepted that multiple-locus methods are superior, statistically, to locus-by-locus methods. Here, we present our method that makes the multiple-locus analysis of data from genome wide association studies routine. It is formulated within a linear mixed model framework. We call our method AMplus. AMplus produces results faster than competing multiple-locus methods and often with greater statistical power. Also, it is just a little slower than the fastest single-locus linear mixed model implementations. AMplus is freely available as a fully documented R package.

Over the past decade, Genome wide association studies have changed considerably in both their analysis and design. Early gwas followed a case-control design. Association mapping methods were no more complicated than contingency table tests or simple linear regression. These

designs though had a tendency to yield spurious findings if there was unrecognised population stratification. This prompted a shift towards family-based designs and score tests, such as the tdt test and its variants (refs). Today, instead of by design, it is through statistical modelling that we account for the effects of population stratification. This has meant that data can be collected from general populations, even if these populations are highly structured. Analysis via sophisticated association mapping methods, such as linear mixed model based approaches, is now almost routine.

What has not changed over the past decade is that it remains common practice to analyse gwas data on a locus by locus basis. This is despite there being several significant problems with analysing data in this way. First, the aim of association mapping is to identify regions of the genome that house genes that are influencing a trait. The identification of these regions from these analyses is not always straightforward. Gwas results are reported, typically, via Manhattan plots that plot the $-\log_{10}$ of the p value for each locus against the map position of the locus. The location of peaks in this plot signify regions of interest. Inferring the exact number of regions of interest can be difficult if the peaks are not well separated. Second, when multiple statistical tests are performed, the probability of wrongly accepting a significant result (type 1 error) is inflated. This is known as the multiple testing problem (refs). Many different solutions have been offered (refs). Yet, there is still no well accepted way of correcting for multiple testing in the context of genome- wide association mapping. Third, many of the traits whose genetic secrets we are trying to discover are complex. There will be multiple loci in linkage equilibrium with genes that influence the trait. Yet, A locus by locus mapping approach only assesses the evidence for

association between a single marker locus and trait.

It is somewhat surprising then that multiple locus association mapping methods haven't attracted more attention. Methods based on regularisation techniques, such as ridge regression and lasso, measure all locus-trait associations simultaneously. Here, multiple testing is not an issue. These techniques though are computationally demanding. Also, their results can be difficult to interpret. The strength of association is not measured by a p-value but by the size of the regression coefficient for a locus in the model. More recently, associations have started to be mapped with random forests (refs). Similar to regularisation techniques though, it is not clear how to infer genomic regions of interest from their findings (refs). An multiple locus method that does show promise is the multi-locus linear mixed model method (ref). The best multiple locus model is built with simple variable selection. Results are immediately interpretable but here, computation can be a challenge for large data.

In this paper, we present our new multiple locus method for genome wide association mapping, which we are calling Eagle. Eagle combines the strength of regularisation techniques (being able to fit all locus-trait associations jointly), with simple variable selection (having easy to interpret results). Our method does not require a significance threshold to be set nor regularisation parameters to be fine tuned. Through a clever dimension reduction step, we are able to achieve a computational performance similar to the fastest single locus linear mixed model implementations. Eagle is statistically more powerful than single locus association mapping and is as and often more powerful than most multiple locus methods. Our aim is to make multiple locus association map-

ping on a genome wide scale routine. To this end, we have created a fully documented R package, that is easy to use, even for non R users. Our package accepts marker data of different formats and can handle data larger than a computer’s memory capacity. It includes detailed error checking and makes heavy use of distributed computing for computation when available. The purpose of this work was to make multiple-locus association mapping on a genome-wide scale, for large data sets, practical and we have built AMplus accordingly.

1 Results

Association Mapping Methods We compared AMplus, in terms of computational and statistical performance, against seven other association mapping methods. We chose methods that had been purpose built for genome-wide analysis, that could handle data from quantitative traits, and where the methods had been implemented in freely available computer software. Two of the methods are based on single locus (or locus-by-locus) models and five are based on multiple locus models. Of the many ways of performing single locus association mapping, we chose GEMMA and FaST-LMM because of their popularity and computational speed. For multiple locus association mapping, we chose bigRR, glmnet, LMM-Lasso, MLMM, and r2VIM. Each takes a different approach to mapping multiple locus-trait associations jointly.

From a practical perspective, how the methods were implemented, in terms of usability, varied greatly. In Table 1, we list seven important features for ensuring the usability of computer software for association mapping. We also identify which software has what features. In forming

this list of features, we assumed that the primary purpose of a genome-wide association study is to identify the set of marker loci in true and strongest association with a trait. The level of documentation varied across the implementations but only a few came with user manuals detailing the format of the input data and how to perform analyses. Most implementations could handle additional fixed effects which is important when accounting for hidden population structure. Few though could cope with marker data larger than the memory capacity of the computer. Similarly, few returned the best set of marker loci in strongest association with a trait as their result. Almost all the software returned results that required further processing, either by identifying the best set of marker loci via the setting of significance thresholds or having to do further analysis to calculate the significance of the results. Almost none of the association mapping software checked for errors in the input data. Only AMplus has all seven features.

Computational Performance: run times and memory usage To assess the computational performance of AMplus, we conducted a large simulation study. We were interested in the impact of study size on performance so we generated data under five different scenarios. These are a GWAS of size 150 individuals and 5000 snp (150 X 5K), 350 individuals and 400000 snp (350 X 400K), 1500 individuals and 50000 snp (1500 x 50K), 2000 individuals and 500000 snp (2000 x 500K), 4000 individuals and 1500000 snp (4000 x 1.5M), and 10000 individuals and 1500000 snp (10000 x 1.5M). We chose these scenarios to reflect some of the different sized GWAS being performed in animals, plants, and humans. For each scenario, we generated 100 replicates of data. A single replicate consists of snp genotype data and quantitative trait data. We obtained the snp data from the publicly available 1000 genome project (phase 3). The quantitative trait data we

generated from the snp data by selecting a set of snp loci, assigning allelic effects to these snp, and aggregating these effects for each individual along with random error. The number of snp selected per replicate follows a Poisson distribution with mean 30. The quantitative trait was generated to have a heritability of 50%. Analyses were performed on a high end desktop computer. It had dual 8-core Xeon processors, three Kepler Tesla GPUs, and 128 Giggabytes of RAM. All implementations except, GEMMA, made use of distributed computing, either explicitly or implicitly through multi-threaded BLAS/LAPACK libraries.

The run times for AMplus against the other software programs, across the five scenarios, is shown in figure 1 To help with interpretability of the results, in both plots, we have taken the x and y axes to be on a log scale. This means a unit change on the x or y axes is equivalent to a change in the order of magnitude. In the top plot, a point is the median of the ratios of elapse times of the multiple locus method to AMplus for a given scenario. The median is over the 100 replicates. Here, since the median of the ratios are all positive on the log scale, it means that AMplus had a shorter run time than than all the other multiple locus methods. In fact, in some cases, AMplus was over a hundred times, or over two orders of magnitude, faster. Unlike AMplus, the size of data under scenario 10000 X 1.5M was beyond the memory constraints of the other multiple locus implementations. This was also the case for LMM-Lasso, bigRR, and glmnet, for scenario 4000 x 1.5M.

In the bottom plot of figure 1, we compare the median run time of AMplus against the median run times of the single locus methods, FaST-LMM and GEMMA. FaST-LMM was run

in two ways. It was run where the genetic similarity matrix was built with all the marker data (FaST-LMM^{all}) or with data on every 500th snp (FaST-LMM^{few}). AMplus was also run in two ways. The default behavior for AMplus is to make use of CPUs for computing. However, AMplus also has the capacity to harness multiple GPUs (AMplus^{GPU}). From the bottom plot, all the implementations have short run times when analysing data from scenario 150x5K. However, for the other scenarios, AMplus and AMplus^{GPU} have significantly shorter run times than GEMMA and FaST-LMM^{all}. FaST-LMM^{few} has a shorter run time than AMplus and AMplus^{GPU} but only for scenarios 1500x50K and 10000x1.5M. Furthermore, for scenario 10000x1.5M, the median run time for GEMMA is 4071 minutes, for AMplus is 699 minutes, for AMplus^{GPU} is 447 minutes, and for FaST-LMM^{few} is 346 minutes. The very fast single-locus method FaST-LMM^{few} is only 29% faster than our multiple-locus method AMplus^{GPU}. It is worth noting though that there is a setup cost to accessing GPU computing, making AMplus^{GPU} most efficient on the larger data.

We also examined the memory usage of the different software programs (figure 1 supplementary materials). AMplus is comparable to the most efficient single locus implementation, FaST-LMM^{few}. AMplus is also the only multiple locus program able to analyse data under scenarios and AMplus can process data larger than the memory capacity of the computer.

Statistical Performance: power, FDR, and significance thresholds As part of assessing the performance of AMplus against the other association mapping methods, we calculated their statistical power and false discovery rates. We want a method to have high statistical power (probability of finding a true positive finding) but low false discovery rate (proportion of findings that are false

positives). All the methods, besides AMplus and MLMM, required a threshold to be set in order to identify significant findings. A conservative threshold guards against false discoveries but reduces power. A anti-conservative threshold increases power at the cost of also increasing the false discovery rate. Only AMplus and mlmm avoid thresholds by treating association mapping as a model selection problem where the best model is found by minimising the model selection statistic, extBIC.

Figures 1 and 1 show the relationship, calculated empirically, between the power and fdr for each method. The multiple-locus (Figure 1) and the single-locus (Figure 1) methods are plotted separately for clarity. AMplus features in both plots for comparison. By varying the significance thresholds for the different methods, we were able to calculate the change in power and fdr. Since AMplus and MLMM do not rely on thresholds, their power and fdr appear as single points in the plots. In practice, the true power and fdr is unknown. However, because we generated data where we knew which loci were acting as quantitative trait loci, we were able to calculate the true power and fdr for the methods.

From figures 1 and 1, the superior performance of AMplus is apparent. When the fdr is low, AMplus has the highest power with Mlmm also performing well. AMplus is noticeably more powerful than the single locus methods. It is possible to set a significance threshold for these other methods that results in higher power. However, this comes at the cost of an intolerably high fdr.

For the methods evaluated here that require a significance threshold, figure 1 shows how threshold size impacts fdr. Each plot corresponds to the results from a different method. A curve

within a plot is formed from analysing data with the method and using different threshold values to identify locus-trait associations. The true *fdr* is then calculated from these findings. We can see from figure 1 that the threshold impacts the *fdr* differently across methods. The relationship between threshold value and *fdr* even differs when data are analysed with the same method but for different scenarios. This demonstrates one of the challenges in using association mapping methods that are reliant upon thresholds. With AMplus, we do not have this challenge. For completeness, for the small sampled size scenarios ... And, the *fdr* for AMplus was 0.07 and 0.09, respectively. With the larger study sizes of ..., ..., and, the *fdr* drops to ..., ..., and respectively.

Mouse Study

2 Methods

Outbred Mice Study Phenotypic and genotypic data were collected on a large study population of commercially available outbred mice. The aim of the study was to map, through association, the genetic basis of complex traits in mice. The construction of the population and collection of data are described in detail in ¹. Briefly, the population consists of 2073 adult outbred mice from Charles River Laboratories. Phenotypic data were obtained from 200 traits. These traits fell into the categories of behaviour, tissue, and physiology. Genotype data were collected from 7, 073, 398 SNPs. A reduced set of 359, 559 SNP across the 19 autosomes and sex chromosome was then identified, upon which all genome-wide analyses were based. These SNP tagged all other SNP with $MAF > 0.1\%$ and $LD\ r^2 > 0.98$. Missing data?

Eagle Approach for Multiple-locus Association Mapping

2.0.1 Overview

Eagle is an approach that identifies genomic regions of interest for a trait via the building of a (linear mixed) model. The model is built iteratively, without need of significance levels or thresholds. The model has two parts, a fixed effects part and a random effects part. The random effects part contains the error and an effect for the snp-trait association across the entire genome. It is from this effect that we are able to identify which snp is in strongest association with the trait. Our model building process is as follows. First we fit the current model to the data. Second, from the random effect for association, we identify which snp is in strongest association with the trait. Third, we move this snp from the random effects to the fixed effects. We do this to stop the neighbours of the chosen snp from being selected in subsequent iterations and to give opportunity to snp in linkage disequilibrium with genes having lesser impact on the trait to be discovered. Fourth, we calculate if the new model is "better" than the current model. If so, the new model becomes the current model and we repeat the process. Once the model building process is complete, the findings from Eagle are the set of snp that have been moved into the fixed effects part of the model. These snp map the separate regions of the genome that house genes that are influencing the trait.

2.0.2 The specifics

The Eagle approach for multiple-locus association mapping is based on building a linear mixed model, iteratively, via forward selection. It is a simplification of a whole-genome method for mapping quantitative trait loci in classic linkage studies ². Suppose s iterations of our model building process have already been performed. This means s snp-trait associations have been identified. It also means that s separate genomic regions of interest have been found. To perform the $s + 1$ th iteration of our model building procedure, we do the following.

First, we fit the current model to the data. Let $S = \{S_1, S_2, \dots, S_s\}$ be a set of ordinal numbers where S_k is the S_k th ordered snp that was selected in the k th iteration. For example, if three iterations of our model building procedure have been performed and say the 500023rd, 15th, and 420th, snp were selected, then $S = \{500023, 15, 420\}$. Let $y^{(n \times 1)}$ be a vector containing the quantitative trait data. Let $M^{(n_g \times L)} = [m_1 m_2 \dots m_L]$ be a matrix containing the genotype data which have been collected from L loci that span the genome on n_g groups/lines/strains. In this paper, n and n_g are equal. It is common for the columns of this matrix to be in map order but this is not a requirement. The vector $m_j^{(n \times 1)}$ contains the genotypes, from the n individuals, for the j th snp. The genotypes are coded as -1, 0, and 1 corresponding to snp genotypes AA, AB, and BB, respectively.

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The (current) model is of the form

$$y = X\tau + Zu_g + e \quad (1)$$

where $X^{(n \times p)}$ and $Z^{(n \times n_g)}$ are known design matrices with X being of full rank and Z containing zeros and ones that assign the appropriate genetic effect to each observation. The vector $\tau^{(p \times 1)}$ has p fixed effects parameters including the intercept. The vector $u_g^{(n_g \times 1)}$ contains the genetic effects. The vector of residuals is $e^{(n \times 1)}$ whose distribution is assumed to follow $N(0, \sigma_e^2 I^{(n \times n)})$. So far, this model differs little from standard linear mixed models for association mapping (refs). However, it is how we specify u_g that distinguishes our model from others.

The genetic effects u_g are modelled as

$$u_g = \sum_{k=1}^s m_{S_k} a_{S_k} + M_{-S} a_{-S} \quad (2)$$

where $m_{S_k}^{(n_g \times 1)}$ is the vector of genotypes for the S_k th snp locus which is the k th selected snp, a_{S_k} is the additive effect of the S_k th snp locus, $M_{-S}^{(b \times L-s)}$ is the matrix of snp genotypes with the data for the selected snp in S removed, and $a_{-S}^{(L-s \times 1)}$ is a random effect whose distribution is $a_{-S} \sim N(0, \sigma_a^2 I^{(L-s \times L-s)})$. The terms in the summation on the left hand side are fixed effects. The other term is a random effect. The terms in the summation account for the additive effects of genes that are in linkage disequilibrium with the selected snp. The other term models snp-trait associations along the entire genome, except for those snp that have already been selected. This is a simple genetic model but it is effective for discovering snp-trait associations. It also reduces the computational cost of our model building procedure.

Second, we estimate the parameters of (1) and (2) via residual maximum likelihood estima-

tion.

Third, we identify the $(s + 1)$ th snp that is in strongest association with the trait, based on the maximum score statistic $t_j^2 = \frac{\tilde{a}_j^2}{\text{var}(\tilde{a}_j)}$ where \tilde{a}_j is the best linear unbiased predictor (BLUP), and $\text{var}(\tilde{a}_j)$ is its variance. This statistic is not only appealing intuitively, where we identify a snp based on its (random) effect size and accuracy, but is theoretically justified. It follows from outlier detection in linear and linear mixed models (ref).

Fourth, we determine the importance of the $(s + 1)$ th selected snp via a model selection strategy. We begin by reforming (2) where S now contains the $s + 1$ selected snp. We then fit this new model to the data via maximum likelihood and calculate its extended Bayesian information criteria (ExtBIC)³. The ExtBIC is a model selection measure that takes into account the number of unknown parameters and the complexity of the model space. It is especially well suited to the model selection problem in genome-wide association studies³. If this new model has a larger ExtBIC than the current model, then the $s + 1$ th selected snp is added to the current model and the above process is repeated. If this new model has a smaller ExtBIC than the current model, then the model building process is complete. The set of snp in strongest association with the trait is the s snp previously identified.

2.0.3 Reducing the dimension of the model

In practice, estimating the parameters of (2) can be demanding, computationally. The vector a_{-S} has $L - s$ random effects where in modern genome-wide association studies, L , the number of snp, can be extremely large. An alternative model is given by Verbyla ^{4,5}. They show how to reformulate (2) to be a model with a random effect with only n elements

$$u_g = \sum_{k=1}^s m_{S_k} a_{S_k} + (M_{-S} M_{-S}^T)^{1/2} a_{-S}^* \quad (3)$$

where $a^* \sim N(0, \sigma_a^2 I^{(n_g \times n_g)})$, and $(M_{-S} M_{-S}^T)^{1/2}$ can be calculated via single value decomposition (ref). Although it may not be obvious, the two models are equivalent, having identical variance structures. Yet, the computational cost of model (3) compared to model (2) is much less due to the random term having only n effects needing estimating.

Verbyla ^{4,5} go on to show how to recover \tilde{a} from estimates from model (3) with

$$\tilde{a} = [M_{-S}^T (M_{-S} M_{-S}^T)^{-1/2}] \tilde{a}^* \quad (4)$$

where its variance matrix is

$$\text{var}(\tilde{a}) = M_{-S}^T (M_{-S} M_{-S}^T)^{-1/2} \text{var}(\tilde{a}^*) (M_{-S} M_{-S}^T)^{-1/2} M_{-S} \quad (5)$$

These values are needed in order to calculate the score statistic t_j^2 for identifying the snp in strongest association with the trait. Fortunately, when calculating t_j^2 , only the diagonal elements of the variance matrix are needed which simplifies the calculation. of (5).

Comparison Methods

2.0.4 Multiple locus methods

We compared the computational and statistical performance of Eagle against five other multiple locus applications for association mapping. These were BigRR, LMM-Lasso, glmnet, MLMM and, r2VIM. BigRR, LMM-Lasso, and glmnet implement different regression-based regularisation methods. Regularisation methods have dominated the development of multiple locus approaches. They give opportunity for the effect of all the snp to be modelled simultaneously. By introducing a penalty function that balances bias and variance, regularisation methods can handle parameter estimation problems where the number of predictors is far greater than the number of samples. MLMM is similar in philosophy to Eagle in that it treats association mapping as a model building problem. Here, a linear mixed model is built through a forward-backward selected strategy. R2VIM is different to the others in that it implements random forests for association mapping. Random forests is a machine learning approach that forms an ensemble of decision trees for regression and classification. These five methods were selected because of their demonstrated value for analysing genetic data, they reflect a range of different statistical methodologies for association mapping, and they are available as either stand-alone computer programs or as packages. A summary of how the computer programs/packages differ in their features is given in Table X. Each method is now described in a little more detail.

BigRR: BigRR ⁶ is an R package that implements generalized ridge regression. In ridge regression, the coefficients of the predictors are shrunk to zero by a constant amount. In generalized ridge regression, the amount of shrinkage is allowed to vary, giving opportunity for a more realistic

model to be specified. The model used in BigRR can be formulated as a linear mixed model that is $y = X\beta + Mu + e$ where $u^{(L \times 1)}$ is a vector of snp effects, and each snp effect is normally distributed with its own variance such that $u_l \sim N(0, \sigma_l^2); l = 1, 2, \dots, L$. The parameters are estimated via a algorithm that makes clever use of the Cholesky decomposition of the genomic relationship matrix, and the equivalence between the above model and animal models ???. One of the challenges in using BigRR for association mapping is that while it has the capacity to estimate the effect sizes of all the snp simultaneously, obtaining their statistical significance is difficult. Permutation has been suggested as a way of calculating significance empirically ⁶ but we instead opted for stability selection.

LMM-Lasso: LMM-Lasso ⁷ is a stand-alone computer program that implements lasso within a linear mixed model framework. Unlike in ridge regression, lasso performs parameter estimation and variable selection, simultaneously. The coefficients of "unimportant" predictors are shrunk to exactly zero. The (full) model is $y = \sum_{l=1}^L m_l \theta_l + g + e$ where $g^{(n \times 1)}$ is a vector of random polygenic effects such that $g \sim N(0, \sigma_g^2 K)$, and $K^{(n \times n)}$ is a realised relationship matrix reconstructed from the snp information. It is challenging, computationally, to apply lasso directly to the full model for parameter estimation. Instead, a two step procedure is adopted. In the first step, the above model without any snp effects is fitted. In the second step, the unknown variance components in the full model are replaced by their estimates that were calculated in the first step. Lasso is then applied to this simpler model making lasso more tractable, computationally. As with BigRR, the statistical significance of the snp effects are not calculated. We use stability selection to obtain the significance empirically.

Glmnet: Glmnet ⁸, unlike BigRR and LMM-Lasso, was not purpose built for association mapping. It is a general R package for regression analysis. We have included it in our list of applications for comparison with Eagle because the regularization method, elastic net, upon which the package is based has been found to be superior to ridge regression and lasso (ref needed???). Elastic net uses a penalty function that is a mixture of ℓ_1 -norm (lasso) and ℓ_2 -norm (ridge regression) penalties ⁹. This allows elastic net to avoid the limitations of lasso while still being able to perform shrinkage and variable selection, simultaneously. Glmnet can handle a range of different models. For the analyses performed in this paper, the model is $y = X\beta + \sum_{l=1}^L m_l\theta_l + e$ where $\theta_l; l = (1, 2, \dots, L)$ is the coefficient for the l th snp. We used stability selection to obtain the significance of the coefficients (snp effects).

MLMM: Of the multiple-locus applications considered in this paper, MLMM ¹⁰ bares the greatest similarity to Eagle algorithmically. MLMM, like Eagle, performs association mapping by building the "best" linear mixed model. The model used for analysis is the same as (1) except that the genetic effect is

$$u_g = \sum_{k=1}^s m_{S_k} a_{S_k} + g \quad (6)$$

where $g^{(n \times 1)}$ is a vector of random polygenic effects with distribution $g \sim N(0, \sigma_g^2 K)$. The model is built iteratively, via forward-backward variable selection. In Eagle, at each iteration, the snp in strongest association with the trait is found by calculating the score statistic t_j^2 . In MLMM, at each iteration, a separate linear mixed model analysis is performed for each snp not already selected. The snp yielding the most significant result (lowest p-value) is selected for inclusion in the fixed effects part of (7). Two selection criteria are available for the model building process: the EBIC

and the multiple-Bonferroni criterion. We used only the EBIC.

r2VIM: R2VIM is an R package specifically designed for association mapping¹⁵. It implements random forests and a new way of measuring the importance of a snp. In random forests, the worth of a predictor is measured, empirically, by calculating its importance score. It is from these importance scores that snp can be ordered in terms of their strength of association with a trait. The challenge though is in knowing what proportion of the highest ordered snp are in true association. R2VIM addresses this by calculating a relative importance score for a snp. Briefly, m random forest analyses of the data are performed. Each analysis is reliant upon a different random number seed. From an analysis, a relative importance score for a snp is obtained by taking the ratio of its importance score to the absolute minimum of the importance scores across all the snp. Those snp with high relative scores across all m analyses are deemed to be of interest¹⁵. Applying r2VIM to the analysis of GWAS data is not without its challenges. First, it is not possible to force every decision tree to have important (fixed) effects without a change to the internal workings of the r2VIM package. So we adopted a two step strategy. In the first step, the fixed effects are regressed on to the trait and the residuals recorded. In the second step, r2VIM is employed where the trait data are the residuals obtained from the first step. This is not ideal (ref to problems with two step approaches). Second, there is no relationship between relative importance scores and significance levels. Consequently, setting an appropriate genome-wide threshold is problematic. We avoid this problem by focusing, in the simulation study, only on the relationship between threshold level and its false discovery rate and power.

2.0.5 Single locus methods

We were also interested in comparing Eagle to single locus methods to measure the difference in run times and statistical power. Even by limiting our focus to LMM-based methods, there were a number of efficient applications to choose from. We decided on GEMMA⁹ and FaST-LMM¹⁶. These two applications have the same computational complexity⁹, produce exact instead of approximate results, and are highly efficient computationally. They were developed at the same time, independently, but are very similar theoretically. Both perform a single spectral decomposition of the relationship matrix K . Both use the eigenvector matrix to rotate the data. Both reformulate the log likelihood and restricted/residual maximum-likelihood (REML) log likelihood into a sum of n terms that are easier to compute. The difference lies in their estimation procedure. FaST-LMM implement's the Brent's algorithm to optimise δ . GEMMA instead implement's the Newton-Raphson algorithm. Newton-Raphson is more complicated in that it also requires the first and second derivatives of a function to be calculated. However, it is superior in terms of its convergence properties to the Brent algorithm. Both applications are stand-alone computer programs, popular, and in common use.

3 Stability Selection

When using BigRR, LMM-Lasso, and glmnet, the results are affected by the amount of regularization (λ). BigRR has an approximate way of setting each snp's separate regularization parameter (ref). However, the optimal value of λ must be found when using LMM-Lasso and glmnet. Also,

all three applications yield the effect sizes of the snp across the entire genome but not their statistical significance. To address these issues, we made use of stability selection[?]. Stability selection is a resampling strategy. With stability selection, we were able to avoid having to optimise λ while still being able to calculate, empirically, the statistical significance of the snp effects.

We performed stability selection as follows. For LMM-Lasso and glmnet, we performed a preliminary analysis to find an appropriate value for the regularization parameter. We adjusted λ so that they yielded between 10 to 30 snp with non-zero effects. Fortunately, with stability selection, the setting of λ does not have to be exact but limiting LMM-Lasso and glmnet to only 10 to 30 non-zero effect sizes was straightforward. We then repeatedly subsampled, without replacement, the data. As recommended[?], we draw 100 subsamples of size $n/2$. We analysed the subsample, with λ fixed for LMM-Lasso and glmnet. To calculate, empirically, the statistical significance of each snp across the genome, we counted the number of times a snp had a non-zero effect size over all the replicates to the total number of replicates (which was 100).

For BigRR, we modified our stability selection procedure slightly. There was no need to find an appropriate value for the regularization parameters as the BigRR package has an internal procedure for their calculation. We draw subsamples as above and analyzed the data with BigRR. However, within an analysis, we ordered the snp according to the absolute size of their snp effects and recorded the top 20 snp. We then measured the significance of the snp across the entire genome as above.

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4 Simulation Study

To explore the computational and statistical performance of Eagle, we conducted a large simulation data. Genome-wide data were generated under five different scenarios. These scenarios were data where the sample size was X and the number of snp Y ($X \times Y$),

We generated data under five different scenarios.

We generated replicates via data perturbation (ref: Zhao et al. 2007).

These are a GWAS of size 150 individuals and 5000 snp (150 X 5K), 350 individuals and 400000 snp (350 X 400K), 1500 individuals and 50000 snp (1500 x 50K), 2000 individuals and 500000 snp (2000 x 500K), 4000 individuals and 1500000 snp (4000 x 1.5M), and 10000 individuals and 1500000 snp (10000 x 1.5M). We chose these scenarios to reflect some of the different sized GWAS being performed in animals, plants, and humans. For each scenario, we generated 100 replicates of data. A single replicate consists of snp genotype data and quantitative trait data. We obtained the snp data from the publicly available 1000 genome project (phase 3). The quantitative trait data we generated from the snp data by selecting a set of snp loci, assigning allelic effects to these snp, and aggregating these effects for each individual along with random error. The number of snp selected per replicate follows a Poisson distribution with mean 30. The quantitative trait was generated to have a heritability of 50%. Analyses were performed on a high end desktop computer. It had dual 8-core Xeon processors, three Kepler Tesla GPUs, and 128 Giggabytes of RAM. All implementations except, GEMMA, made use of distributed computing, either explicitly

or implicitly through multi-threaded BLAS/LAPACK libraries.

Replicates are generated with data perturbation (Zhao et al. 2007). Data perturbation makes use of the phenotypic and genotypic data observed in a real study to create a simulated (quantitative) trait. It affords us the opportunity to generate replicates whose analysis more closely mirrors the complexities of analyzing real data. For computational expediency, our simulation study is based on data observed from the Biloela site in 2006. In generating trait data for a replicate, we assume a broad based heritability of 0.6, there is a single major QTL, the polygenic component consists of 30 polygenes where the effect of a polygene is formed from an unmapped SNP which was chosen randomly, and the variance structure of the simulated trait closely follows that of the loaf volume trait. The marker data consists of those genotypes collected in the association study from the 3129 SNP across the 21 homologous chromosomes. Each replicate has the same marker data. It is the simulated trait data that varies across replicates. Our simulation study consists of three parts. First, we conduct a null study to investigate whether our Monte Carlo sampling approach controls correctly the genome-wide type I error rate. We also evaluate the impact of assuming a block diagonal structure for V . We do this by also calculating the genome-wide type I error rates when the variance matrix for the joint distribution of the test statistic is formed from loci pairings across the entire genome. Ten thousand replicates are generated where the QTL has no effect. Results are reported for QK-based association mapping implemented as a single-stage and weighted two-stage analysis. Second, a power study is performed to investigate if there is a difference in performance between implementing QK-based association mapping as a single-stage analysis versus a two-stage analysis. Data are generated under QTL of different sizes and with

varying percentages of missing marker data (1.1, 5, 9, 11). Third, a study to evaluate our heuristic procedure for choosing between single-stage and two-stage analysis is performed. Data are generated under a QTL of no, moderate, and large effect. One thousand replicates are generated for each differently sized QTL. The performance of our heuristic procedure is measured by calculating the proportion of replicates for which the analysis type is inferred correctly. To determine if a single-stage analysis is truly needed, it is necessary to analyze fully the genome-wide data via single-stage and weighted two-stage analysis. Then, for each marker locus j , perform the test — $-\log_{10} p_{adj} - \log_{10} q_{adj} - \frac{1}{2} - \log_{10} p_{adj} - 1.3$ —, where we have assumed a genome-wide significance level of 5.

5 Implementation

6 New section written on iPhone added 5 Feb 2018

7 Simulation Study

We conduct a large simulation study to examine the performance characteristics of Eagle. Genome-wide data are generated under five different scenarios. These are a GWAS of size 150 individuals and 5000 SNPs (150 x 5K), 350 individuals and 400,000 SNPs (350 x 400k), 1500 individuals and 50,000 SNPs (1500 x 50k), 200 individuals and We chose these five scenarios to reflect some of the different sized studies being run in animals, plants, and humans. For each scenario, 100 replicates are generated.

To generate a replacate, we used data perturbation (ref). Data perturbation makes use of real genotype data from which a (quantitative) trait is created. With data perturbation, we are able to preserve some of the challenges of analysing real data while still having trait data whose genetic architecture is known.

The genotype data comes from the 1000 genome project, version 3 (ref). The original data contains X individuals with XX XX snp. From these data, according to the specifications of the scenario, the individuals and the snp loci are identified at random. What about polymorphic affect numbers? Check).

To generate the quantitative trait data, first the number of snp that are to be in complete with an underlying qtl (x) is drawn from a poisson distribution with mean equal to 30. Second, x snp loci are select randomly from a scenario?s set of genotype data. Third, the snp values are summed across the selected loci for each individual to generate a vector of n genetic values. Here, each snp is having, roughly, the same effect on the trait, remembering that the maf of a snp also impacts a snps influence on a trait. Fourth, n deviates are drawn from a $N(0, \sigma_e^2)$ where σ_e^2 has been set to yield a trait with a heritability of 0.6. (Put in formula). Fifth, the quantitative trait data $= g + e$. In this way, the same genotype data are used, within a scenario, for all replicates. Only the quantitative

8 Implementation

Eagle has been implemented as an R package of the same name. However, much of the calculation is performed through c++ functions for improved performance. Eagle has been specially designed

to harness the multithreaded math libraries blast and lapack but it will run on a single processor computer. It can also deal with data larger than the memory capacity of a computer. Eagle can be run as a standard R package or via the web based GUI. The GUI relinquishes the need to know R to use Eagle.

Eagle can be downloaded from CRAN and installed in the usual way. A containerised version of Eagle is also available. It is built with singularity (ref). This container already has Eagle installed on a multi-threaded version of R. The container is downloaded as a single file from ccxx

A quick start guide, how to videos, installation instructions, and answers to frequently asked questions are available at the Eagle website (<http://...>)

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We compared the performance of Eagle against seven other association mapping applications. Five of these applications, BigRR, glmnet, FaST-LASSO, MMLM, r2VIM implement methods that have been applied to the multiple locus analysis of data from genome-wide association studies. The other two applications, GEMMA, and FaST-LMM implement single-locus association mapping methods. The packages differ in their features (summarized in Table X). The packages also differ in their underlying methods (summarized below).

8.0.1 Multiple locus methods

BigRR: In the BigRR package, generalised ridge regression is implemented for the analysis of genetic data ⁶. Generalised ridge regression is a regularisation method in that it produces regression solutions that have higher bias than ordinary least squares but reduced variance for improved prediction accuracy. The underlying model is $y = X\beta + Mu + e$ where $u^{(L \times 1)}$ is a vector of genetic effects, and each genetic effect is normally distributed with its own variance such that $u_l \sim N(0, \sigma_l^2); l = 1, 2, \dots, L$. The amount of shrinkage is allowed to vary across the genetic (snp) effects. This differs to standard ridge regression where the shrinkage is constant. Generalised ridge regression is apt at estimating the effect sizes of snp on a genome-wide scale but obtaining their statistical significance analytically is difficult. Shen ⁶ suggested permutation as a way of calculating significance empirically. We instead opted for stability selection (see below of details).

glmnet: Glmnet is a general purpose R package for regression analysis ⁸. The model used for analysis is $y = X\beta + \sum_{l=1}^L m_l \theta_l + e$ where $\theta_l; l = (1, 2, \dots, L)$ is the coefficient for the l th snp. All snp effects are modelled, simultaneously. Glmnet implements elastic net for the estimation of the parameters. Elastic net is a regularisation method where the penalty function is a mixture of ℓ_1 -norm (lasso) and ℓ_2 -norm (ridge regression) penalties ⁹. This allows elastic net to avoid the limitations of lasso while still being able to perform shrinkage and variable selection, simultaneously ⁹. Glmnet was not specifically built for association mapping. However, it is included in our list of comparison applications due to its solid performance for genomic prediction ^{11,12}. The glmnet

package implements a modification of the LARS algorithm ¹³ for finding the optimal regularisation (solution) path. From exploratory analyses of the simulated data, we found this algorithm yielded a high proportion of false positives. We instead fixed the shrinkage parameter to a suitable value and employed stability selection to estimate the significance of the snp coefficients, empirically (see below).

LMM-Lasso: LMM-Lasso is a stand-alone computer program for association mapping within a linear mixed model framework ⁷. The (full) linear mixed model used for analysis is $y = \sum_{l=1}^L m_l \theta_l + g + e$ where $g^{(n;\times 1)}$ is a vector of random polygenic effects such that $g \sim N(0, \sigma_g^2 K)$, and $K^{(n;\times n)}$ is a realised relationship matrix reconstructed from the snp information. All snp effects are modelled, simultaneously. However, a limitation of this model is its inability to accommodate additional fixed effects (there is no $X\beta$ term). The parameters are estimated via lasso ¹⁴. Lasso is a regularisation method with the capacity to perform parameter estimation and variable selection, simultaneously. Lasso is computationally intensive, especially when applied to linear mixed models. To avoid this problem, a two-step estimation procedure is adopted. In the first step, the above model but without any snp effects is fitted via maximum likelihood. In the second step, the unknown variance components in the full model are replaced by their estimates that were calculated in the first step. This turns the full model from a linear mixed model into a simpler linear model. Here, lasso is far more tractable, computationally. To assess the significance of the snp coefficients, as suggested ⁷, we implemented stability selection (see below).

MLMM: Of the multiple-locus applications considered in this paper, MLMM bares the

greatest similarity to Eagle algorithmically. MLM, like Eagle, performs association mapping by building the "best" linear mixed model. The model used for analysis is the same as (1) except that the genetic effect is

$$u_g = \sum_{k=1}^s m_{S_k} a_{S_k} + g \quad (7)$$

where $g^{(n \times 1)}$ is a vector of random polygenic effects with distribution $g \sim N(0, \sigma_g^2 K)$. The model is built iteratively, via forward-backward variable selection. In Eagle, at each iteration, the snp in strongest association with the trait is found by calculating the score statistic t_j^2 . In MLM, at each iteration, a separate linear mixed model analysis is performed for each snp not already selected. The snp yielding the most significant result (lowest p-value) is selected for inclusion in the fixed effects part of (7). Two selection criteria are available for the model building process: the EBIC and the multiple-Bonferroni criterion. We used only the EBIC.

r2VIM: R2RIM is an R package specifically designed for association mapping¹⁵. It implements random forests and a new way of measuring the importance of a snp. In random forests, the worth of a predictor is measured, empirically, by calculating its importance score. It is from these importance scores that snp can be ordered in terms of their strength of association with a trait. The challenge though is in knowing what proportion of the highest ordered snp are in true association. R2VIM addresses this by calculating a relative importance score for a snp. Briefly, m random forest analyses of the data are performed. Each analysis is reliant upon a different random number seed. From an analysis, a relative importance score for a snp is obtained by taking the ratio of its importance score to the absolute minimum of the importance scores across all the snp. Those snp with high relative scores across all m analyses are deemed to be of interest¹⁵.

8.0.2 Single locus methods

FaST-LMM: FaST-LMM is a highly efficient stand alone computer program for association mapping¹⁶. It is based on a standard single-locus model^{17,18}. Its computational efficiencies come through factorising K with spectral decomposition so that $K = U(S + \delta I)U^T$ where U contains the eigenvectors and δ is the ratio of the genetic variance to the residual variance. By rotating the data with U , the log likelihood of the linear mixed model can be expressed as the log likelihood for a simple linear model. This greatly reduces the computational cost of performing lmm-based association mapping. With FaST-LMM, analyses scale linearly with sample size n , both in terms of run-time and memory usage.

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Figure 1 Run times

Figure 2 Memory usage

Figure 3 Power curves for multiple locus methods

Figure 4 Power curves for single-locus methods – Put into sup methods

Figure 5 The impact of threshold on FDR