A genomic investigation on hybrid sterility in house mouse using a two stage model

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Summary: Genome wide association studies (GWASs) commonly used to search for genetic variants associated with quantitative traits. Pleiotropic effect of genes may cause the observed correlations among different phenotypes. This study proposed a two stage multilocus model for pleiotropic GWAS using a Bayesian mixture model to take into account of both small and major gene effects. The objectives of this study were to investigate if the two-stage model was useful for detecting pleiotropic genes using a simulated data set and to investigate existence of pleiotropic genes for testis weight and testis gene expression levels in house mouse. The analyses included relative testis weights and testis gene expression traits. The results showed that two stage model had higher power to detect the pleiotropic QTL than the single marker model. It was also noted the possible economical impact of sampling informative individuals for the GWAS analyses by observing genomic trends in the simulated dataset. Two stage model detected 50 and 53 major SNP effects using first and the second principal components. Additive genetic variation explained by chromosome X was found to be 4% for the testis weight.

Keywords: Bayesian mixture model, genome wide association analyses, pleiotropy.

Introduction

Gene flow is one of the important sources of genetic variation by hybridization of two distinct populations. Genetic biodiversity increase at hybridization area of locally separated populations (21). This hybrid might have hybrid sterility phenotypes due to divergent alleles at various loci (25). Storchova et al. (20) investigated the role of X chromosome in association with hybrid sterility phenotype.

Genome wide association studies (GWASs) commonly used to search for genetic loci associated with quantitative phenotypes. A single gene might affects multiple phenotypes: that leads to construct of networks and pathways of genes (7,8). Pleiotropic effect of genes may cause the observed correlations among different phenotypes (23). Hartley et al (8) reported more than 8500 associations obtained from 350 human complex traits: indicates that pleiotropy exist (24). Shriner (18) reviewed the statistical models to detect pleiotropic effects. Currently multivariate approaches are commonly used to combine multiple phenotypes for pleiotropic GWAS analyses. Gao et al. (4) demonstrated that GWAS using residuals of linear combination of multiple phenotypes lead to higher statistical power and lower false discoveries over single trait analyses. However most of the pleiotropic models are employing single marker regression approaches using outcome of multivariate models. Assumptions for different gene effects lead to different models (hence results) in GWAS. Yang et al. (26) used a polygenic GWAS approach for human height and
explained higher proportion of missing heritability (explanatory variance was found to be 50 %) compared with the common practice of single marker regression GWAS (explanatory variance was found to be 5 %). Hill (9) suggested to incorporate small and pleiotropic effects into genomic studies. Gratten et al (7) also stressed the importance of both polygenic and mendelian pleiotropic effects for the proper interpretation of GWAS. Based on this suggestions and discussions this study proposed a two-stage model for pleiotropic GWAS using a Bayesian mixture model (16) to take into account of both small (polygenic) and major gene effects.

The objectives of this study were to 1) investigate if the two-stage model was useful for detecting pleiotropic genes using a simulated data set (22) and 2) to investigate existence of associated loci and pleiotropic genes for hybrid sterility (for testis weight and testis gene expression levels) for house mouse (21) using the two stage model.

**Materials and Methods**

The present work describes the two-stage model for detecting pleiotropic genes using both simulated QTLMAS data and real mouse datasets.

**QTLMAS data set:** The pedigree included four generations with 4100 individuals for the simulated quantitative traits (22). The genome consisted of 10000 Single Nucleotide Polymorphisms (SNPs) distributed over 5 chromosomes. The number of population founders were 1020 (1000 females and 20 males). Each male mated with 51 females to create families. Generations were forced to be nearly discrete hence overlapping. Phenotypes of 3080 individuals were available. Three correlated quantitative phenotypes were simulated to mimic milk yield (heritability =0.35), fat yield (heritability=0.35) and fat content (heritability=0.50) in the sheep. The detail of the simulated dataset could be found in Usai et al (22).

**Mice Data Set:** The mice data consists of 179 males from the house mouse hybrid zone (21). The analyses included relative testis weight and testis gene expression traits. Before statistical analyses, SNPs were removed from the data if the call rate are < 90%, minor allele frequencies are < 1 %. Finally, a total of 152342 SNPs were collected for the genomic analyses. The detail of the mouse dataset could be found in Turner and Harr (21).

**A two Stage Pleiotropic GWAS model:** This study proposed a two-stage model of pleiotropic GWAS as a means of increasing power of detecting both small and major gene effects using multivariate dimension reduction (3) and genomic prediction models (15). Following we give a general formulation of the two-stage model.

Factor and principal component analyses could be used to discover loadings that combines underlying relationships hence pleiotropy among different phenotypes in GWAS. Stage 1 can be based on linear and orthogonal combinations of the residuals of the correlated phenotypes ( y₁, y₂, ..., yₚ);

\[
\begin{align*}
y₁ &= a_{11}x₁ + a_{12}x₂ + \ldots + a_{1p}xₚ \\
y₂ &= a_{21}x₁ + a_{22}x₂ + \ldots + a_{2p}xₚ \\
&\vdots \\
yₚ &= a_{p1}x₁ + a_{p2}x₂ + \ldots + a_{pp}xₚ
\end{align*}
\]

with the coefficients (a) being chosen to convey highest explanatory variances compared with the original explanatory variables, x₁, x₂, ..., xₚ , (3).

The second stage employs a Bayesian mixture model (BayesR) (16) for predicting pleiotropic SNP effects based on outcome of the Stage 1. BayesR assumed a hierarchical models of mixtures using different SNP effects:

\[
p(β|p, σ^2_β) = \sum_{k=1}^{p} p_i f(x|θ_i)
\]

where β is the SNP effects, p is the mixture proportions (assumed to be 0.00001, 0.0001, 0.001, 0.01), σ^2_β is the genetic variance, f(x|θ_i) is normally distributed mixture densities with θ parameters vectors and observations, x_i. The present study sampled 50000 markov chains and discarded first 5000 as burn in period and recorded every 10th sample for thinning the chain. Uninformative priors were also used to obtain desired posteriors.

**A Single SNP Model:** Discordant sib pair experimental design was also employed as described in Karacaören (10) using Yu’s linear mixed model (27) as implemented in rrBLUP package (2,17). Pearson homogeneity statistic could be estimated from 2xm table via following formula;

\[
T^1 = \sum_{i=1}^{m} \frac{(n_{ij} - n_{ij})^2}{n_{ij} + n_{ij}}
\]

where ni,j stands for counted alleles among cases and controls, i = 1, 2 for sib 1 and sib 2, respectively and j = 1,…,m (number of alleles). Due to dependency of test statistics within siblings; permutation tests could be used for deciding of statistical thresholds. This study permuted the case and control states of the siblings with probability of 1/2 to detect threshold of significance level of the test statistics.

The present study used genomic kinship information in linear mixed model to take into account of the pedigree structure as was implemented in rrBLUP: (2) a linear mixed model (27) approach in R software (17)

The linear mixed model (single marker regression approach) was used as

\[
y = Xb + Za + e
\]
where y contains the observations, b is the sex effects, a is the additive genetic effect, matrices X and Z are incidence matrices, and e is a vector containing residuals.

\[
\text{Var}
\begin{pmatrix}
    a \\
    e
\end{pmatrix}
\sim
N
\begin{pmatrix}
    0 & \sigma^2_a \\
    0 & \sigma^2_e
\end{pmatrix}.
\]

For the random effects, it is assumed that A is the coefficient of coancestry obtained from genotype of animals; I is an identity matrix, \(\sigma^2_a\) is the additive genetic variance and \(\sigma^2_e\) is the residual variance.

**Results**

**QTLMAS dataset:** 301 SNP’s were excluded due to minor allele frequency <1%, leaving 9699 SNP’s in the analyses. Mean heterozygosity for the SNPs and the individuals were estimated as 0.35 with a standard error of 0.01. Normality was confirmed by Kolmogorov Smirnow test, \(p>0.150\) for each trait. Based on model (1) estimated genomic heritabilities were 0.31, 0.30, 0.47 for quantitative traits, respectively.

Different genetic architectures were assumed for the QTLMAS phenotypes. BayesR assumed that SNPs effects were obtained by different mixture proportions from the normal distribution. Since some phenotypes could be controlled by both polygenic and major genes such an approximation might be useful. The Markov chain algorithm was run 4 times and the trace plot of number of significant SNPs was investigated (Table 1). Visual inspection of the trace plots show convergence (results not shown) of Markov chains. The posterior mean number of major SNPs were 51, 73 and 200 for trait 1, trait 2 and trait 3 respectively.

Single marker regression model was used (2, 27) to detect putative genomic associations using different number of discordant sib pairs (DSP) (Table 1). DSP design leads to homogenized samples by controlling for population stratification. In DSP design, sib pairs from each family could be used as cases and controls. Boehnke and Langefeld (1) used one case and one control from each family. In this study different number of DSP was sampled to association mapping using QTLMAS simulated dataset similar to Karacaören et al. (11) and Karacaören(12). Table 1 showed that number of detected true SNPs and QTLs increased with larger discordant sib pair families.

Table 2 (a, b, c, d) presents genotype and allele counts for top two markers (1685 and 3585) by counting all alleles (scheme 1) or discordant alleles (scheme 2). Table 2 showed that both GWA using all individuals and samples of discordant sip pairs gave similar results. Contrasting alleles that are discordant between sib pairs (scheme 2) also increased the association test statistics compared with test statistics obtained by all alleles (scheme 1) (Table 3). The p values were obtained using Monte Carlo simulations (Figure 1) based on 100.000 permutations of the data. Permutated p-values show agreement with GWA results.
Table 1. Number of true / number of total significant SNPs (number of true QTLs) from genome wide association analyses (GWA) with all individuals (All) and different sampling schemes for discordant sib pairs (DSP) (n=2, 5, 10, 15, 20).

Table 2a. Genotype counts for markers 1685
Table 2b. Genotype counts for markers 3585
Table 2c. Allele counts for markers 1685
Table 2d. Allele counts for markers 3585

Table 3. Test statistics, T^2, for all alleles, DSP alleles and whole genome wide association analyses for different markers (Pearson correlation coefficient values and p values).

Table 4. Estimates of genomic heritabilities using first (1) and second (2) scores obtained from different rotation techniques.

Linear combinations of multivariate phenotypes were used (loadings) for detecting pleiotropy. Loadings were used as a response variable with the models to detect pleiotropic genes. Principal component analyses can be used to discover underlying pleiotropic patterns of genomic datasets. The method could also be used for reduction of dimension of datasets as well. Principal component rotation techniques were used to obtain linear combinations of multivariate phenotypes for detecting pleiotropy. Loadings were used to estimate heritabilities and to detect putative associations (Table 1 and Table 4).
30 (45) SNPs were detected. A total of 36 and 48 pleiotropic QTLs were detected by single marker and the two stage approaches. Therefore, the two stage model had higher power to detect the pleiotropic QTL than the single marker model.

**Mouse data set:** A hierarchical Bayesian mixture model was used (16) for detecting major and minor SNP effects for relative testis weight and genome wide testis gene expression patterns. Since most of the quantitative traits influenced by number of genes with various effect sizes: such an assumption might lead to useful findings in practice (6). Visual inspection of the trace plots show convergence (results not shown) of Markov chains. The posterior mean number of SNPs for the testis weight were 1123.

Table 5 shows that there are SNPs with major SNP effects. Additive genetic variation explained by chromosome X was found to be 4% for the testis weight. Over all 1123 SNPs explained 99% of the total genetic variance.

**Discussion and Conclusion**

**QTLMAS data set:** Table 1 showed that number of detected true SNPs and QTLs increased with larger discordant sib pair families. This was particularly true for the trait 1. Although Marker 1685 had highest –logP value (Table 3), it was not found to be statistically significant by counting Scheme of 1 (p =0.1009). The main reason was uncorrected inbreeding structure in the genotypic data set. However, when counting Scheme of 2 was used to incorporate the inbreeding structure, the genomic signal was detected by at the level of 100,000 permutations. There was a good agreement for marker 3585 using both allele counting schemes and whole genome wide association analyses (Table 3). The results showed that both GWA using all individuals and samples of discordant sip pairs gave similar results. Contrasting alleles that are discordant between sib pairs (scheme 2) also increased the association test statistics compared with test statistics obtained by all alleles (scheme 1) (Table 3). Hence evidence for association was much higher when using discordant alleles instead of using all alleles. The p values were evaluated using Monte Carlo simulations (Table 3) based on 100,000 permutations of the data. Permutated p-values show agreement with full genome wide association results. Stronger association were observed for marker 3585 compared with marker 1685. The possible economical impact of sampling informative individuals was also noted for the GWAS analyses by observing genomic trends in the Table 1. For example, usage of discordant sib pairs of 20 (detected 16 QTLs) had higher accuracy instead of using the whole individuals (detected 11 QTLs).
The two stage analyses were able to identify additional new positions missed by the single marker regression model. This was particularly true for the phenotype 2 (14 true QTLs) and phenotype 3 (28 true QTLs).

For pleiotropic analyses: simulated GWAS data of 9699 SNPs were employed with 3080 individuals for 3 quantitative phenotypes: the heritabilities were equal to 0.35, 0.35, and 0.50 respectively. The outcome of principal component analyses of the 3 quantitative phenotypes were used for detecting pleiotropic effects by the Bayesian hierarchical model (BayesR). The assumed SNPs effects were obtained with different effect sizes: the first containing 56 SNPs with major effects, the second containing 128 SNPs with smaller effect, and a large group of 1410 SNPs representing a polygenic component for the first principal component. The second principal component contained 64 major effects, 66 moderate effects and 545 polygenic effects. Since the QTLMASS data set simulated with high level of inbreeding structure: moderate and polygenic effects were essential for taking into account of the associated pedigree (and population) structure. The genomic signals from the major effects of the BayesR were compared with the organizers true results (22) by the threshold of 1Mb. BayesR was able to detect 20 and 30 true SNPs using first and second principal components respectively. A total of 48 (out of 50) QTLs were detected on both principal components. In general, the two-stage model showed the highest detection power, particularly when it was used with the second principal component. However, the two stage model also produced larger number of false positives: success rates were found to be 0.42 and 0.67 (for principal components 1 and 2, respectively): compared with the results of single marker regression approach as 0.85 and 0.61 (for principal components 1 and 2, respectively). However, the two-stage model was able to detect higher number of true QTLs (48 true QTLs) compared with the single marker regression approach (36 true QTLs). Similar to our findings, importance of genetic architecture was also stressed by Leong et al. (13), Liu et al. (14) and Usai et al. (22) by comparing the different genetic models and assumptions for detecting pleiotropic genes. However, Gianola et al. (5) pointed out potential problems for detecting pleiotropy using molecular markers due to unaccounted linkage disequilibrium structure in the GWAS. The proposed two-stage model may have limitations in terms of false positive signals, however it has an advantage over most other single marker regression models since it does take into account of linkage disequilibrium by modeling whole SNPs simultaneously.

Mouse data set: Turner and Harr (21) assumed major genes for the testis weight and detected genomic signals from 9 autosomes and X chromosome by using permissive false discovery rate (stringent threshold detected four SNPs). Since Turner and Harr (21) used threshold for p values to take into account of multiple hypothesis testing such results expectable. The proposed model that affects multiple SNPs simultaneously can resolve the multiple hypothesis testing problem in pleiotropic analyses of GWAS. BayesR model provides flexibility in terms of effect sizes for the SNPs. The analyses of this study showed that most of the SNPs (>0.98) had small effects for the testis weight, demonstrating importance of the polygenic component. By carrying out a pleiotropic GWAS in a male house mouse, up to 103 SNPs that are associated in top two principal components variation with joint effects of testis weight and testis gene expression levels were identified. Top two principal components explain 21% of the total variation.

A new two stage method to detect pleiotropic SNPs using a Bayesian mixture model (16) was presented to increase accuracy in this study. By definition, GWAS uses hypothesis free explanatory tools. In that regard two-stage model of the present study might be useful to investigate pleiotropic GWAS referring different genetical architectures for the phenotypes. Employing the Bayesian mixture model (BayesR) at the second stage corrects for both the linkage disequilibrium and the polygenic effects (16). Compared with the single marker regression pleiotropic models: the two stage model was also prune to well known multiple hypothesis testing problems in GWAS. Although the simulated data showed that two-stage model of the present study subject to false positive results, it still might be informative (and hypothesis producer) to use it for scanning the genome to detect major and small pleiotropic effects.

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