M⁴: an improved SNP calling algorithm for Illumina BeadArray data

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ABSTRACT

Summary: Genotype calling from high-throughput platforms such as Illumina and Affymetrix is a critical step in data processing, so that accurate information on genetic variants can be obtained for phenotype–genotype association studies. A number of algorithms have been developed to infer genotypes from data generated through the Illumina BeadStation platform, including GenCall, GenoSNP, Illuminus and CRLMM. Most of these algorithms are built on population-based statistical models to genotype every SNP in turn, such as GenCall with the GenTrain clustering algorithm, and require a large reference population to perform well. These approaches may not work well for rare variants where only a small proportion of the individuals carry the variant. A fundamentally different approach, implemented in GenoSNP, adopts a single nucleotide polymorphism (SNP)-based model to infer genotypes of all the SNPs in one individual, making it an appealing alternative to call rare variants. However, compared to the population-based strategies, more SNPs in GenoSNP may fail the Hardy–Weinberg Equilibrium test. To take advantage of both strategies, we propose a two-stage SNP calling procedure, named the modified mixture model (M⁴), to improve call accuracy for both common and rare variants. The effectiveness of our approach is demonstrated through applications to genotype calling on a set of HapMap samples used for quality control purpose in a large case–control study of cocaine dependence. The increase in power with M⁴ is greater for rare variants than for common variants depending on the model.

Availability: M⁴ algorithm: http://bioinformatics.med.yale.edu/group.

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Supplementary information: Supplementary data are available at Bioinformatics online.

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1 INTRODUCTION

Genome-wide association studies (GWAS) have resulted in the discovery of numerous genetic variants contributing to major human diseases (E SKAT et al 2009; SKAT et al 2009). The Wellcome Trust Case Control Consortium (2007). These studies benefit from the success of the International HapMap Project in cataloging and characterizing millions of single nucleotide polymorphisms (SNPs) for the purpose of GWAS (The International HapMap Consortium 2003). Another major contributing factor is the availability of high-density and low-cost SNP arrays, such as Illumina and Affymetrix, which allow researchers to perform genotyping platform, an accurate genotyping method: http://bioinformatics.med.yale.edu/group.

For any microarray platform, an accurate genotyping algorithm is needed to convert the observed probe intensities into genotypes, and the Illumina BeadStation platform is a critical step in data processing. So that probes for SNPs may be inferred from their expected positions (Giannoulatou et al 2008; Teo et al 2008, 2009), with green-red intensity to measure two alleles, A and a, at each SNP for every calling algorithm considers data from all SNPs at a time. We call these algorithms the population-based ones. Their basic premise is that the three possible genotypes of an SNP with alleles A and a, namely AA, Aa and aa, will form three distinct clusters from the minor allele frequency (MAF) of an SNP is low. In practice, three fraction of SNPs may be shifted away from their expected positions, which will lead to a genotyping error rate of ~1% from missing genotypes (Browning and Yd 2008). Another algorithm, GenoSNP, is distinguished from the population-based algorithms in that it genotypes all SNPs within one individual at a time under the assumption that probes for different SNPs have different response features across the genome. That is, instead of genotyping every individual in the Illumina BeadArray to both platforms, such as CRLMM (Carvalho et al 2009) and BEAGLECALL (Browning and Yd 2008). In this article, we focus on the Illumina platform.

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the Hardy–Weinberg Equilibrium (HWE) test using this algorithm, suggesting a possible violation of the assumption that all the SNPs behave similarly across the genome.

To take advantage of both the population-based and the SNP-based calling approaches, in this article, we propose a two-stage SNP calling procedure without a reference population for the Illumina ReadArray platform. We call this procedure the 'modified mixture model' (M2). Generally, this method integrates the population-based strategy, e.g. GenCall, with the SNP-based genotyping algorithm, e.g. GenoSNP, to improve call accuracy for both common and rare variants. M2 is evaluated through comparison with other genotyping algorithms for Illumina microarray data.

2 STATISTICAL METHODS

2.1 Illumina Chip data description

We first describe the features of the Illumina data before introducing our method. In the probe design, the Illumina array is composed of many beadpools that consist of hundreds of thousands of beadtypes. With the method. In its probe design, the Illumina array is composed of many 20mer probe sequence to hybridize near the beadtype site (Steemers et al. 2006), and 20 beads on average for each beadtype provide 20 pairs of allele-specific intensities for each DNA sample. Thus each beadtype assay two SNP alleles represents an SNP. In our study, the pair of raw intensities at each SNP for every individual are measured, and clusters of genotypes are inferred based on these measured intensities.

2.2 Model

Let \( x_{ji} \) denote the pair of raw intensities at the \( j \)-th SNP for the \( i \)-th individual, and its distribution is modeled as a four-component Gaussian mixture model (McLachlan and Peel 2000). In fact, each measurement \( x_{ji} \) can be considered as arising from one of these components with probability \( \pi_k \), where \( k = 1, 2, 3 \) or 4. While performing SNP calling, the first three components in the mixture model correspond to three genotypes (AA, Aa, and aa), and the last one is the null component with zero mean and large variance. The indicator variable \( z_{ji} \), where \( z_{ji} = 1, 2, 3, 4 \) or 0 denotes the latent genotype class for the \( j \)-th SNP of the \( i \)-th subject. Given the above notations, the complete likelihood function for the observed data is given by:

\[
\log L(\Theta; x) = \sum_{i=1}^{n_i} \Phi(x_{ji}; \pi_k; \Sigma_k) z_{ji} = \sum_{i=1}^{n_i} \Phi(x_{ji}; \pi_k; \Sigma_k) \mathbb{1}(z_{ji} = k)
\]

where \( k = 1, \ldots, 4 \), \( j = 1, \ldots, S \), \( n_i \) is the total number of individuals observed for the \( j \)-th SNP, and \( S \) is the total number of SNPs. Given the \( j \)-th SNP, \( x_{ji} \) follows a multivariate normal distribution. The unknown parameters of the Gaussian mixture model \( \pi_k, \Sigma_k, \Theta \) are estimated through the following Expectation Maximization (EM) algorithm (McLachlan and Peel 2000).

The E (Expectation) step calculates the conditional expectation of Equation (1) given the observed pair of raw intensities \( x_{ji} \). When the current estimates are \( \hat{\Theta} \) for \( \Theta \), in the \( t \)-th iteration, the conditional expectation of the log likelihood function is

\[
E(\Theta; x_{ji}) = \sum_{k=1}^{4} \pi_k \Phi(x_{ji}; \mu_k; \Sigma_k)
\]

Since the log likelihood function is a linear function of the unobservable indicator variable \( z_{ji} \), the E-step is replaced by the following expectation equation:

\[
f(x_{ji}; \hat{\Theta}) = \frac{\pi_k \Phi(x_{ji}; \mu_k; \Sigma_k)}{\sum_{k=1}^{4} \pi_k \Phi(x_{ji}; \mu_k; \Sigma_k)}
\]

The parameters of the normal components are estimated in the M (Maximization) step. The iterative estimates for the mean \( \mu_k \) and variance–covariance matrix \( \Sigma_k \) are

\[
\mu_k^{(t+1)} = \frac{1}{t} \sum_{j=1}^{t} \Phi(x_{ji}; \mu_k^{(t)}; \Sigma_k^{(t)})
\]

\[
\Sigma_k^{(t+1)} = \frac{1}{t} \sum_{j=1}^{t} (x_{ji} - \mu_k^{(t)}) (x_{ji} - \mu_k^{(t)})^T
\]

Note that the parameters can be estimated by gmm package in MatLab (MathWorks, 2009).

After the above EM algorithm converges, the conditional probability that the pair of intensities \( x_{ji} \) belong to the \( i \)-th cluster is inferred as the Posterior (PR) for the \( i \)-th SNP.

\[
P(x_{ji}; \hat{\Theta}) = \frac{1}{\sum_{i=1}^{4} P(x_{ji}; \hat{\Theta})}
\]

Note that the APR is one important criterion applied in the second stage of our two-stage SNP calling procedure to select high-quality SNPs. Here, the APR is defined as

\[
A_{i,j}(y) = \sum_{i=1}^{n_i} \Phi(x_{ji}; \mu_k^{(t)}; \Sigma_k^{(t)}) / \sum_{i=1}^{n_i} \Phi(x_{ji}; \hat{\Theta})
\]

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\]

The PR can be used to identify the observations with strong signals of clusters. Based on the PR, the average posterior rate (APR) for the \( j \)-th SNP \( y_j \) is defined by

\[
\rho_j = \frac{1}{\sum_{i=1}^{n_i} \Phi(x_{ji}; \hat{\Theta})}
\]

After the above EM algorithm converges, the conditional probability that the pair of intensities \( x_{ji} \) belong to the \( i \)-th cluster is inferred as the Posterior (PR) for the \( i \)-th SNP.

2.3 Two-stage SNP calling procedure

The two-stage genotyping procedure for SNP calling is designed to integrate the population-based strategy implemented in GenCall and the SNP-based approach implemented in GenoSNP. In the first stage, a Gaussian mixture model is used to call each SNP in turn across the whole genome. This step is a population-based calling method. In the second stage, a union set of SNPs with low MAFs (e.g. MAF < 0.05) and poor APR (e.g. APR < 0.9) are selected, and each selected SNP is re-called with the assistance
Although Ref-1 SNPs have a large APR, some of them may not have high-quality SNPs. We consider three possible criteria to select a reference SNP in this step. At the second step, Ref-1 SNPs with each cluster containing all subjects for one SNP. We have found that SNPs with a lower MAF tend to have their calling accuracy improved) and 'reference SNPs' to denote good-quality SNPs.

2.4 Reference SNP selection

A very important component of our proposed procedure is to find an appropriate reference SNP to improve SNP calling accuracy of SNPs that are difficult to call accurately on their own. To achieve this objective, a three-step selection procedure is proposed. Throughout this section, we use 'testing SNPs' to denote SNPs that need to be re-called (that is, to have their calling accuracy improved) and 'reference SNPs' to denote good-quality SNPs.

Step I: selecting SNPs with high APR as candidate reference SNPs. In the first stage, the APR of one SNP is defined as the average value of PR of good-quality SNPs.

Step II: selecting SNPs with good clustering properties from Ref-1 SNPs. Although Ref-1 SNPs have a large APR, some of them may not have high-quality clusters. The shapes, centers and boundaries of clusters for these Ref-1 SNPs may not provide precise distribution parameters of each cluster as a reference. At the second step, Ref-1 SNPs with each cluster containing at least 10% of samples are further selected, denoted as Ref-2 SNPs.

Step III: measuring the similarity between the testing SNP and each Ref-2 SNP. We consider three possible criteria to select a reference SNP in this step.

(a) Calculating the APR of each aggregated dataset formed by the testing SNP with each Ref-2 SNP in turn. For example, an aggregated dataset of size \( (n_0, n_{test}) = 1 \) is made up of the c-th testing SNP \( (n_{test} \times 1) \) and d-th reference SNP \( (n_0 \times 1) \). The relevant APR of this aggregated data is given by

\[
p_{tc}^d = \frac{\sum_{i=1}^{n_{test}} \sum_{j=1}^{n_0} P_{ij}}{n_{test} \times n_0}
\]

where \( P_{ij} \) is defined by Equation 8. We select the Ref-2 SNP that gives the largest APR \( p_{tc}^d \), where \( d = 1, \ldots, T \) is the total number of Ref-2 SNPs for the c-th testing SNP for this aggregated dataset.

(b) Calculating the Mahalanobis distance [4,6] between the testing SNP and each Ref-2 SNP, and the Ref-2 SNP with the minimum Mahalanobis distance value is selected. Generally, the Mahalanobis distance measures the overall similarity between two SNPs.

(c) To explore the detailed resemblance of each cluster between the testing SNP and each Ref-2 SNP, we further introduce a measure, Cluster Distance (\( S_c \)), in the following to compare the clusters between two SNPs. Both the testing SNP and Ref-2 SNPs are classified into three clusters corresponding to three genotypes in the following form,

\[
y_k \begin{cases} 
\text{if } y_k < u_1 \\
y_k \text{ if } u_1 \leq y_k < u_2 \\
y_k \text{ if } y_k \geq u_2 
\end{cases}
\]

where \( y_k \) is a simple projection function of \( x_k \). It transforms the 2D vector \( x_k \) into a univariate variable \( y_k \) without losing main clustering characteristics [4,6], and it is easy to group transformed intensity \( y_k \) into three clusters in terms of Equation 10. Note that \( r^* \) denotes the index of clusters where \( y_k \) falls inside the \( r^* \)-th cluster and \( r^*=1,2 \) or 3. \( u_1 \) and \( u_2 \) are used to classify the intensities \( y=(y_1, \ldots, y_5) \) of the c-th SNP into three genotype clusters with the following two steps:

(i) \( y_i \) is roughly classified into different clusters, and at least three observations are in each cluster. In the first step, \( u_1 \) and \( u_2 \) are given by

\[
\begin{align*}
u_1 &= -0.5 \\
u_2 &= 0.5
\end{align*}
\]

(ii) Update \( u_1 \) and \( u_2 \) to find the optimum boundaries to identify distinct clusters.

If \( y_i \) is classified into three clusters with means \( \mu_{r^*} \), \( r^*=1,2 \) or 3) corresponding to three genotypes in the first step, \( u_1 \) and \( u_2 \) are further defined by:

\[
\begin{align*}
u_1 &= \mu_{r^*} - \frac{\mu_{r^*-1} - \mu_{r^*}}{2} \\
u_2 &= \mu_{r^*} + \frac{\mu_{r^*} - \mu_{r^*+1}}{2}
\end{align*}
\]

If \( y_i \) is grouped into two clusters in the first step, \( u_1 \) and \( u_2 \) are given by:

\[
\begin{align*}
u_1 &= \mu_{r^*} - \frac{\mu_{r^*-1} - \mu_{r^*}}{2} \\
u_2 &= \mu_{r^*}
\end{align*}
\]

If only one cluster is generated for \( y_i \) in the first step, \( u_1 \) and \( u_2 \) are not defined.

Once \( y_i \) is grouped into different clusters in an appropriate way, we define a measure \( S_c \), which quantifies the similarity of clusters between the c-th testing SNP and each Ref-2 SNP.

\[
S_c = \min_{d=1, \ldots, T} \left\{ \sum_{r=1}^{3} (n_{test} - n_{ref}) \times \left( x_{ref}^{d,r} - x_{test}^{d,r} \right)^2 \right\}^{1/2}
\]

where \( d=1, \ldots, T \) and \( r \) is the total number of Ref-2 SNPs for the c-th testing SNP, \( n_{ref}^{d,r} \) is the transformed intensity vector of the \( r \)-th cluster at the c-th testing SNP, \( x_{ref}^{d,r} \) denotes the mean vector of the d-th Ref-2 SNP for the \( r \)-th cluster, \( \Sigma_{ref}^{d,r} \) and \( \Sigma_{test}^{d,r} \) are the variance–covariance matrices of the c-th testing SNP and d-th Ref-2 SNP, respectively. In general, \( S_c \) measures the minimum distance between the clusters of the c-th testing SNP and each Ref-2 SNP.

It is desirable to select one reference SNP so that the genotypes of the poor-quality SNP can be best called based on one of the three criteria discussed above. Figure 1 displays the comparison results between the three methods, and shows that the three methods are ranked in the order of Average Posterior Rate of the Mahalanobis Distance of Cluster Distance. The overall comparisons of three methods are evaluated in the Supplementary Materials. We note that this SNP is not a real testing SNP with low MAF, because the distinct performance of three methods, APR, Mahalanobis Distance and Cluster Distance, is difficult to show using low-frequency testing SNPs that may lack some clusters. Thus, we choose this common SNP to show the similarity between the testing SNP and the reference SNP.
3 RESULTS

3.1 Dataset and SNP calling methods

We consider an Illumina Omni 1M dataset that consists of 3258 samples, and 141 out of 3258 samples were from 38 distinct HapMap samples with some individuals genotyped multiple times. The overall performance is evaluated by comparing the SNP calls for these HapMap samples by different methods to those available from the International HapMap Project database (The International HapMap Consortium, 2005). In this article, we focus on the 942,313 SNPs in the whole genome with the exception of the sex chromosome. The X chromosome SNPs (24,717) are analyzed alone. The null genotypes in the HapMap project are ignored. The performance of our proposed method, M3, and other existing calling algorithms is evaluated based on this Illumina dataset.

It has been demonstrated that the call rate and call accuracy of both GenoSNP and CRLMM are better than those of Illuminus when a small number of samples are collected (Ritchie et al., 2011). As for CRLMM, its implementation depends on the reference population and its calculation strongly hinges on computer configuration (Ritchie et al., 2011; Zhang et al., 2010). Based on these considerations, we compare M3 with GenCall as a representative of the population-based method and GenoSNP as the SNP-based method. For the Illumina GenCall approach, genotypes with good GenCall scores (GC score ≥ 0.15) are used as the inferred genotypes. As discussed earlier, GenoSNP is built on a SNP-based mixture model without a reference population to genotype every individual in turn, and this model may be good at genotyping rare variants. It calculates the posterior probability of each sample at a specific SNP, and we use a cut-off value of 85% to select SNPs and samples with good quality. Samples and SNPs having poor clustering properties (low posterior probability) are treated as missing data. As discussed in the model section, the two-stage M3 approach aims to take the advantage of GenCall and GenoSNP. We use the PR of 0.85 as a cut-off to filter samples at a particular SNP in our analysis and a union set of SNPs with MAF < 0.05 and APR < 0.9 are selected to be re-called in the second stage of our proposed method.

3.2 Comparisons

We first evaluate the SNP calling results by evaluating the call rate of each method and the concordance among them. The call rate is defined as the ratio of genotypes passing the calling threshold to the total number of genotypes that need to be inferred. The concordance rate between two algorithms refers to the percentage of agreement of inferred genotypes between two algorithms. The relevant results are summarized in Table 1. In brief, there is high consistency among these three methods (M3, GenCall and GenoSNP) overall. But, there are some discrepancies among these three algorithms. We note that the major homozygote calls by GenCall are more frequently called heterozygote by M3 and M3 more likely genotypes null components by GenCall and GenoSNP (Supplementary Materials). This is partially due to the fact that M3 gives the largest call rate (99.64%), followed by GenoSNP (99.22%) and GenCall (98.16%) (Table 1).

Since the true genotypes are unknown, the above concordance comparisons do not reveal which method performs better. In the

Table 1. The comparisons of call rate and concordance rate among GenCall, GenoSNP and M3

<table>
<thead>
<tr>
<th>Algorithm 1</th>
<th>Algorithm 2</th>
<th>Call rate (%)</th>
<th>Concordance (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GenCall</td>
<td>M3</td>
<td>98.16</td>
<td>99.64</td>
</tr>
<tr>
<td>GenoSNP</td>
<td>M3</td>
<td>99.22</td>
<td>99.64</td>
</tr>
<tr>
<td>GenCall</td>
<td>GenoSNP</td>
<td>98.16</td>
<td>99.22</td>
</tr>
</tbody>
</table>

The unit of call rate and concordance rate is percentage %; M3: the modified mixture model.
Table 2. The comparisons of call rates and concordance on HapMap samples for overall SNPs

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Item</th>
<th>GenCall</th>
<th>GenoSNP</th>
<th>M3</th>
</tr>
</thead>
<tbody>
<tr>
<td>GenCall, GenoSNP and M3</td>
<td>&lt; 50 Call rate</td>
<td>98.03</td>
<td>99.13</td>
<td>99.75</td>
</tr>
<tr>
<td></td>
<td>&lt; 10 Call rate</td>
<td>98.03</td>
<td>99.20</td>
<td>99.76</td>
</tr>
<tr>
<td></td>
<td>&lt; 1 Call rate</td>
<td>98.03</td>
<td>99.21</td>
<td>99.77</td>
</tr>
<tr>
<td></td>
<td>Accuracy</td>
<td>97.85</td>
<td>98.45</td>
<td>99.11</td>
</tr>
<tr>
<td></td>
<td>Accuracy</td>
<td>97.87</td>
<td>98.68</td>
<td>99.20</td>
</tr>
<tr>
<td></td>
<td>Accuracy</td>
<td>97.87</td>
<td>98.74</td>
<td>99.23</td>
</tr>
</tbody>
</table>

M3: the modified mixture model; call rate: the percentage of valid genotypes; accuracy: the percentage of consistent genotype; criterion: which algorithm is selected to count E values between this algorithm and HapMap project due to the mis-assignment of major allele; E: the average error caused by the mis-assignment of the major allele under three criterions, followed by GenCall, GenoSNP and M3, and three E cutoffs, E < 50, 10 and 1 are set.

following, we focus on the genotype calls of 141 out of 3258 samples from 38 distinct HapMap samples using the genotypes of these individuals obtained from the HapMap project as a gold standard in our comparisons. One issue in using the SNP calls from the HapMap data is differentiating the major allele between two alleles at an SNP. In our comparisons, we count the discrepancy in homozygote calls, denoted by E (Error), between each of the three algorithms and HapMap data, and vary the cut-off level at 1, 10 and 50 to remove SNPs with different major allele assignments between each of the three algorithms and HapMap data. With a more stringent threshold, e.g. E < 1, fewer SNPs with inconsistent major allele assignment are compared. The results are summarized in Table 3.

Table 3. Comparisons of call rates and concordance on HapMap samples for rare variants

<table>
<thead>
<tr>
<th>SNPs</th>
<th>Item</th>
<th>GenCall</th>
<th>GenoSNP</th>
<th>M3</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAF &lt; 0.1</td>
<td>&lt; 50 Call rate</td>
<td>97.79</td>
<td>99.11</td>
<td>99.67</td>
</tr>
<tr>
<td></td>
<td>&lt; 10 Call rate</td>
<td>97.81</td>
<td>99.12</td>
<td>99.67</td>
</tr>
<tr>
<td></td>
<td>&lt; 1 Call rate</td>
<td>97.84</td>
<td>99.13</td>
<td>99.68</td>
</tr>
<tr>
<td></td>
<td>Accuracy</td>
<td>97.62</td>
<td>98.59</td>
<td>99.20</td>
</tr>
<tr>
<td>MAF &lt; 0.05</td>
<td>&lt; 50 Call rate</td>
<td>97.73</td>
<td>99.13</td>
<td>99.64</td>
</tr>
<tr>
<td></td>
<td>&lt; 10 Call rate</td>
<td>97.74</td>
<td>99.14</td>
<td>99.64</td>
</tr>
<tr>
<td></td>
<td>&lt; 1 Call rate</td>
<td>97.73</td>
<td>99.15</td>
<td>99.63</td>
</tr>
<tr>
<td></td>
<td>Accuracy</td>
<td>97.54</td>
<td>98.56</td>
<td>99.11</td>
</tr>
<tr>
<td>MAF &lt; 0.01</td>
<td>&lt; 50 Call rate</td>
<td>96.64</td>
<td>99.04</td>
<td>99.56</td>
</tr>
<tr>
<td></td>
<td>&lt; 10 Call rate</td>
<td>96.67</td>
<td>99.06</td>
<td>99.57</td>
</tr>
<tr>
<td></td>
<td>&lt; 1 Call rate</td>
<td>96.71</td>
<td>99.08</td>
<td>99.58</td>
</tr>
<tr>
<td></td>
<td>Accuracy</td>
<td>96.43</td>
<td>98.10</td>
<td>98.77</td>
</tr>
</tbody>
</table>

M3: the modified mixture model; call rate: the percentage of valid genotypes; accuracy: the percentage of consistent genotype; criterion: which algorithm is selected to count E values between this algorithm and HapMap project due to the mis-assignment of major allele; E: the average error caused by the mis-assignment of the major allele under three criterions, followed by GenCall, GenoSNP and M3, and three E cutoffs, E < 50, 10 and 1 are set. The different values in parentheses indicate the number of SNPs whose MAFs are < 0.1, 0.05 or 0.01, respectively.

Table 4. Comparisons of HWE test among GenCall, GenoSNP and M3

<table>
<thead>
<tr>
<th>Population</th>
<th>Num-Sample</th>
<th>Algorithm</th>
<th>No. of failed SNPs</th>
</tr>
</thead>
<tbody>
<tr>
<td>AA I</td>
<td>2005</td>
<td>GenCall</td>
<td>14 447</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GenoSNP</td>
<td>50 860</td>
</tr>
<tr>
<td></td>
<td></td>
<td>M3</td>
<td>27 288</td>
</tr>
<tr>
<td>AA II</td>
<td>83</td>
<td>GenCall</td>
<td>1450</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GenoSNP</td>
<td>14 432</td>
</tr>
<tr>
<td></td>
<td></td>
<td>M3</td>
<td>5155</td>
</tr>
<tr>
<td>EA I</td>
<td>867</td>
<td>GenCall</td>
<td>32 209</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GenoSNP</td>
<td>63 170</td>
</tr>
<tr>
<td></td>
<td></td>
<td>M3</td>
<td>44 123</td>
</tr>
<tr>
<td>EA II</td>
<td>158</td>
<td>GenCall</td>
<td>2801</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GenoSNP</td>
<td>20 342</td>
</tr>
<tr>
<td></td>
<td></td>
<td>M3</td>
<td>7631</td>
</tr>
</tbody>
</table>

AA I: African-Americans not of Hispanic origin; AA II: African-Americans of Hispanic origin; EA I: European Americans not of Hispanic origin; EA II: European Americans of Hispanic origin; Num-Sample: the number of subjects within each population; Algorithm: three algorithms in this table, that is, GenCall, GenoSNP and M3; No. of failed SNP: the number of SNPs fail the HWE test within each population.
A number of algorithms have been developed and are commonly used for SNP calling of Illumina genotyping arrays. One general strategy is the population-based approach where each SNP is analyzed individually and the data from all the study subjects at this SNP are used to define genotype clusters for calling. The higher call rate and more accurate genotype calls of M3 may help to increase statistical power to detect disease-associated variants, especially those with low MAFs. To quantify the power gain, we compare the average power to detect disease-associated variants based on three calling algorithms using the HapMap sample data [ASW (87) and CEU (174)]. We consider a case–control scenario for AA and EA separately and assume a similar genotyping characteristic for the collected data. We assume a disease prevalence of 0.05, and the relative risk $\lambda$ varies from 1.5 to 4 under the multiplicative model. The allele frequencies are based on empirical data from the HapMap project. We consider the statistical significance level by Bonferroni correction ($5 \times 10^{-8}$), and fix the number of cases and controls for both populations (AA: case:controls 1320:758; EA: case:control 749:239). The power to detect the risk alleles based on the correct calls for all individuals and all SNPs is measured as a standard. For the three calling algorithms, we assume that the observed allele frequencies at each SNP and the number of cases or controls are similar to those observed from the HapMap samples collected in our study. The power comparisons for the overall SNPs and rare SNPs are summarized in Figures 3 and 4 respectively. In brief, compared with GenCall and GenoSNP, M3 has the largest power for both common variants and rare variants in both populations. When the relative risk increases, we observe a larger improvement of M3 versus GenCall or GenoSNP. The ratio of $M^3$ versus GenCall or $M^3$ versus GenoSNP measures this improvement, and it has been found that the power of $M^3$ increases 1.46% and 1.03% compared with GenCall for AA and EA, respectively. Similarly, the increase in power of $M^3$ versus GenoSNP is ~0.12% and 0.16% for both populations. In particular, the improvement of $M^3$ versus GenCall or GenoSNP for rare variants is more noticeable than that for common variants. (The increase in power of $M^3$ versus GenCall: 1.84% and 1.12% for both populations; the increase in power of $M^3$ versus GenoSNP: 0.43% and 0.34% for both populations.) In general, the power achieved by $M^3$ is closer to the ideal power when the genotypes of all study subjects are correctly inferred.

We also evaluate the performance of three algorithms on the X chromosome SNP data. The average call accuracy is compared with the HapMap project calls under three $E(\text{Error})$ cutoffs, $E < 50$, 10 or 1. Table 5 summarizes the overall concordance result on the X chromosome among three algorithms. Again, $M^3$ provides the best call rate and accuracy on the sex chromosome SNPs, compared with GenCall and GenoSNP. It has been demonstrated that the model incorporating the gender information will perform better than methods (GenCall and GenoSNP) ignoring this gender information (Fasching et al., 2011). We further examine the gender-dependent model $M^3_{\text{dep}}$, denoted as $M^3_{\text{dep}}$, on the X chromosome. Compared with $M^3$ without incorporating the gender information, a higher call accuracy is achieved by the model $M^3_{\text{dep}}$ involving the gender information (Table 5). The higher call accuracy on the male subjects results in the great improvement of calls. We suggest that the X chromosome SNPs should be called separately using a gender-dependent model ($M^3_{\text{dep}}$) in practice.

4 DISCUSSION

A number of algorithms have been developed and are commonly used for SNP calling of Illumina genotyping arrays. One general strategy is the population-based approach where each SNP is analyzed individually and the data from all the study subjects at this SNP are used to define genotype clusters for calling.

![Fig. 3. The average power to detect risk alleles for overall SNPs by HapMap, GenCall, GenoSNP and $M^3$. Red solid line: the ideal power measured by the HapMap project; blue dash line: the average power measured by the GenCall; purple dot line: the average power measured by GenoSNP; green dot dash line: the average power measured by $M^3$; cutoff: $5 \times 10^{-8}$.

![Fig. 4. The average power to detect risk alleles for less common SNPs by HapMap data, GenCall, GenoSNP and $M^3$. Less common SNPs: the SNPs with MAF < 0.05; cutoff: $5 \times 10^{-8}$.

Different SNP calling algorithms based on the proportion of SNPs failing the HWE test. As mentioned above, our Illumina dataset includes 3258 individuals, consisting most of African-Americans (AA) and European-Americans (EA), who are either Hispanic or non-Hispanic. We perform the HWE test on these four populations separately. The results are summarized in Table 4. Since GenoSNP is a SNP-based calling algorithm that does not consider HWE in the HapMap project; blue dash line: the average power measured by the GenCall; purple dot line: the average power measured by GenoSNP; green dot dash line: the average power measured by $M^3$, respectively. The HWE test failed when the proportion of SNPs failing the HWE test is above 0.12% and 0.16% for both populations. In particular, the decrease in power of $M^3$ versus GenCall or GenoSNP for rare variants is more noticeable than that for common variants. The increase in power of $M^3$ versus GenCall: 1.84% and 1.12% for both populations; the increase in power of $M^3$ versus GenoSNP: 0.43% and 0.34% for both populations. In general, the power achieved by $M^3$ is closer to the ideal power when the genotypes of all study subjects are correctly inferred.

We also evaluate the performance of three algorithms on the X chromosome SNP data. The average call accuracy is compared with the HapMap project calls under three $E(\text{Error})$ error cutoffs, $E < 50$, 10 or 1. Table 5 summarizes the overall concordance result on the X chromosome among three algorithms. Again, $M^3$ provides the best call rate and accuracy on the sex chromosome SNPs, compared with GenCall and GenoSNP. It has been demonstrated that the model incorporating the gender information will perform better than methods (GenCall and GenoSNP) ignoring this gender information (Fasching et al., 2011). We further examine the gender-dependent model $M^3_{\text{dep}}$, denoted as $M^3_{\text{dep}}$, on the X chromosome. Compared with $M^3$ without incorporating the gender information, a higher call accuracy is achieved by the model $M^3_{\text{dep}}$, involving the gender information (Table 5). The higher call accuracy on the male subjects results in the great improvement of calls. We suggest that the X chromosome SNPs should be called separately using a gender-dependent model ($M^3_{\text{dep}}$) in practice.

4 DISCUSSION

A number of algorithms have been developed and are commonly used for SNP calling of Illumina genotyping arrays. One general strategy is the population-based approach where each SNP is analyzed individually and the data from all the study subjects at this SNP are used to define genotype clusters for calling.
Table 5. The comparisons of call rates and concordance on HapMap samples for X chromosome SNPs

<table>
<thead>
<tr>
<th>Criterion</th>
<th>E (Error)</th>
<th>Item</th>
<th>GenCall (%)</th>
<th>GenoSNP (%)</th>
<th>M3 (%)</th>
<th>M3 &lt; Error (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GenCall, GenoSNP</td>
<td>&lt; 50</td>
<td>Call rate</td>
<td>97.67</td>
<td>99.00</td>
<td>99.31</td>
<td>99.75</td>
</tr>
<tr>
<td>M3, M3 &lt; Error</td>
<td>&lt; 10</td>
<td>Accuracy</td>
<td>97.38</td>
<td>98.41</td>
<td>98.65</td>
<td>99.60</td>
</tr>
<tr>
<td>M3, M3 &lt; Error</td>
<td>&lt; 1</td>
<td>Call rate</td>
<td>97.77</td>
<td>99.01</td>
<td>99.32</td>
<td>99.76</td>
</tr>
<tr>
<td>M3, M3 &lt; Error</td>
<td>&lt; 1</td>
<td>Accuracy</td>
<td>97.48</td>
<td>98.45</td>
<td>98.69</td>
<td>99.63</td>
</tr>
</tbody>
</table>

M3 < Error: the gender-dependent modified mixture model; M3: the modified mixture model; call rate: the percentage of valid genotypes; accuracy: the percentage of consistent genotype; criterion: which algorithm is selected to count E values between this algorithm and HapMap project due to the mis-assignment of major allele. E: the average error caused by the mis-assignment of the major allele under four criterions. GenCall, GenoSNP, M3, and M3 < Error, and three E cutoffs, E < 50, 10 and 1 are set.

Since the performance of these population-based methods (such as GenCall) depends on well-defined genotype clusters, a large reference population is needed to achieve good accuracy. However, with an increasing number of less common SNPs on the arrays, the calling accuracy may not be as high due to the lack of information needed to define each of the three possible genotype clusters well. GenoSNP is an SNP-based approach that addresses this challenge, but it relies on the critical assumption that all the SNP probes perform similarly. This assumption is certain to be violated in practice. As a result, many more SNP calls than population-based calls fail the HWE test (P < 0.0001). To exploit the advantage of these two separate approaches, we have proposed a two-stage SNP calling procedure to improve the call accuracy of rare variants while retaining the accurate results for common SNPs. In the first stage of our procedure, we use a mixture model to call every SNP in turn, and then focus on a small fraction of poor-quality SNPs in the second stage by borrowing information from one reference SNP that matches well with the characteristics of the poor-quality SNPs in the genome.

This two-stage approach, named the modified mixture model (M3), was tested on an Illumina dataset with 3258 samples. In general, we observed good agreement between the M3 calls and those from GenCall and GenoSNP. Using 141 out of 3258 samples from 38 distinct HapMap samples in our data that have been genotyped, we were able to investigate the accuracy of different methods using the genotypes reported by the International HapMap Project. Our results show that M3 provides the highest call rate and the best genotyping accuracy. M3 performs better than GenCall on rare variants and generated fewer SNP calls that fail the HWE test than GenoSNP. In addition, M3 can increase statistical power to detect disease-associated variants compared with GenCall and GenoSNP, especially for rare variants.

The essence of M3 is to integrate the population-based statistical model with the SNP-based strategy (GenoSNP), to yield good call accuracy and a high call rate at each SNP without requiring a large reference population, especially for rare variants. An important aspect of M3 is that it searches for the appropriate SNP to assist in calling a poor-quality SNP. In practice, it is computationally prohibitive to search the reference SNP across the whole genome, thus we proposed to select a reference SNP near the testing SNP. Under the Illumina chip design, each bead accommodates a 50mer probe sequence that is made up of A, T, C and G near the SNP (Steemers et al., 2007), and it has been shown that the larger proportion of CG in this probe sequence of one SNP leads to the stronger intensity signal at this SNP (Carvalho et al., 2007). Thus, incorporating this probe sequence in the reference SNP selection procedure may help to find the most appropriate reference SNP. Additionally, empirical results suggest that our proposed selection procedure may be effective and the assumption that probes have similar response characteristics may hold at least for some of the probes. However, when some probes selected as the testing SNPs produce unusual variation in probe responses, the reference SNP cannot provide good cluster information to be used with these poor SNPs. In studies where certain samples with known genotypes, e.g. the HapMap samples in our dataset, the explicit consideration of these gold-standard samples may also help in the selection of reference SNPs. Our proposed method is a data-driven approach. In the future, we may borrow HapMap data information about genotype clusters to improve the quality of genotype calling of the illumina data. It remains to be determined whether there are better ways to identify a reference SNP or whether to include more than one reference SNP in the second-stage analysis.

The efficiency of M3 is strongly dependent on the assumption of the homogeneity of probe measurements. Since this condition may not be satisfied for some SNPs, it is desirable to develop a statistical model without the second-stage analysis to achieve more accurate genotyping results for both common and rare variants. Moreover, M3 fits the Gaussian mixture model on the raw intensities that may violate the assumptions of this mixture model. Thus, an additional normalization step (Illumina Inc., 2005; Teo et al., 2006) can be added to remove outlier SNPs and normalize intensities at each SNP before performing cluster analysis. This algorithm focuses on the analysis of Illumina arrays, and applicability of this idea to Affymetrix is worth investigating.

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M*: an improved SNP calling algorithm

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Conflict of Interest: none declared.

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