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A new statistic and its power to infer membership and phenotype in a genome-wide association study using genotype frequencies

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Abstract

Aggregate results from Genome-Wide Association Studies (GWAS)1-3, such as genotype frequencies for cases and controls were available until recently on public websites4-5 because they were thought to reveal negligible information concerning an individual's participation in a study. Homer *et al.*6 suggested a method for forensic detection of an individual's contribution to an admixed DNA sample could be applied to aggregate GWAS data. Using a likelihood-based statistical framework, we develop an improved statistic that uses genotype frequencies and an individual's genotypes to infer whether the individual or a close relative participated in the GWAS and, if so, the participant's phenotype status. Our statistic compares the logarithm of genotype

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frequencies, in contrast to that of Homer *et al.*6, which is based on differences in SNP probe intensity or allele frequencies. We derive the theoretical power of the test statistics and explore the empirical performance in scenarios with a varying numbers of randomly chosen or top-associated SNPs.

A recent report by Homer *et al.*6 demonstrated a method to determine whether a given person contributed a trace amount (<0.1%) of DNA to a pool of DNA from 200 individuals based on allelic probe intensities from the same microarray genotyping technology commonly employed in GWAS. Three sets of probe data are required for Homer's test statistic, denoted here as T_{allele} : 1) from a pool to which the individual may or may not have contributed DNA; 2) from a reference pool of DNA sampled from members of the same genetic population; and 3) from DNA obtained from a single individual to be tested for membership in the first pool. Their primary applications were directed toward specific forensic challenges, such as determining whether a DNA sample contributed to a biospecimen of mixed origin. They speculated that their method could be applied to GWAS allele frequency data.

We explored that suggestion and have extended the approach of Homer *et al.*6 to propose an improved test statistic T_{geno} to detect whether an individual contributed genotypes to a GWAS, and, if so, determine their case/control phenotype. Conceptually, T_{geno} substitutes frequencies from two groups (e.g., cases and controls) and genotype states of the individual to be tested for pools of admixed DNA and measures of allelic probe intensities in the method of Homer *et al.*6. T_{geno} is a novel likelihood ratio statistic that can detect whether an individual contributed DNA to neither, one, or both groups given only genotypes for the individual to be tested and genotype frequencies from each group. Our statistic is similar to T_{allele} except that T_{geno} contrasts the logarithm of frequencies of the each observed genotype of the tested individual (as opposed to allelic probe intensity or allele frequency in T_{allele}) between two populations.

The predictive power of T_{geno} was derived approximately (see Methods), and evaluated by simulation under a range of scenarios consistent with the size and scope of current GWAS designs. In each scenario, we assumed two independent groups of individuals, denoted "test" and "reference", drawn from a homogeneous population with genotypes from independent biallelic single-nucleotide polymorphism (SNP) markers in Hardy-Weinberg equilibrium with fixed minor allele frequency (MAF). We varied the size of each group, the number and MAF of independent loci, and genotype error rate. Power was computed by simulation, randomly sampling test and reference groups as well as individuals to be tested and comparing T_{geno} to its theoretical null distribution using a two-sided test that assumes Z could be a member of either group.

Table 1 shows simulation results of the sensitivity and specificity of T_{geno} for significance levels ranging from 0.05 to 10^{-6} for scenarios labeled (a)-(e). We observed that the power to detect membership in either group increased as the test group size decreased due to the larger contribution of the individual relative to the size of the test group; power also increased as the reference group size increased due to the improved knowledge of underlying population frequencies. Scenario (c) demonstrated that substantial genotyping

error was required to attenuate the power of the method. Power was essentially unaffected with 1% genotyping error and only began to appreciably diminish with rates approaching 10%. Scenario (d) demonstrated that the power to detect membership does not depend on MAF. Scenario (e) emphasized the role of the size of the population that did not contain the individual being tested. Power was dramatically reduced when only 60 individuals were available in the reference group, reaching only 20% at the 0.05 significance level and no power at the 10^{-6} significance level; power increases to 100% at the 0.05 significance level and to 96% at the 10^{-6} significance level with 5,000 reference individuals. In each scenario, the predicted theoretical and empirical power matched very closely.

In Table 2, we explored the power to detect membership of a close relative of an individual in the test group using 200,000 independent SNPs simulated under the same assumptions as in Table 1. In (f) and (g) the genotype discordance rate due to the familial relationship was a function of MAF (see Methods) and reflects no additional discordance due to assay-related genotyping errors. Power to detect membership of relatives decreased as MAF increased (and consequently the genotype discordance rate increased), but remained high for MAFs lower than 30%. In populations of European descent, the average MAF for most whole-genome SNP panels is between 20-25%.

We applied our method to data combined from several GWAS conducted at the US National Institutes of Health using genotype data from individuals of European descent. The dataset contained 6,733 individuals affected with a disease (cases) and 6,871 unaffected individuals (controls) and genotype data for 557,005 SNPs from the Illumina HumanHap550 assay. Figure 1 shows a histogram of T_{geno} for a hypothetical GWAS with 1,000 cases (in red), 1,000 controls (in blue) and 11,604 subjects not in the virtual study (in gray), and with the theoretical null density curve shown in black. The groups were well-differentiated, indicating that Tgeno is capable of accurately inferring membership and case/control phenotype. Both the mean and variance of the empirical null distribution were shifted from those of the theoretical standard normal distribution. These differences arise from use of fixed test and reference samples, while the theoretical null distribution holds over repeated random samples of the two groups. In addition, LD between SNPs is expected to inflate the variance of the test-statistics relative to our derivation that assumes locus independence. Thus, making inferences about Tgeno when applied to particular GWAS data requires suitable calibration of the null distribution. Figure 2 shows histograms comparing T_{geno} and Tallele for a fixed subset of 1,000 cases and 1,000 controls for varying numbers of "top associated" SNPs.

Using these data, GWAS scenarios were explored by selecting subsets of cases and controls, estimating genotype frequencies of each group, and fitting logistic genotype-phenotype association models. In each scenario, all 13,604 individuals were tested for membership conditional on a fixed set of cases and controls chosen. We also attempted to infer the phenotype of the cases and controls selected in each scenario given the knowledge that they participated in the study. Individuals not selected as cases or controls for a given scenario were used to empirically estimate the null distribution. Figure 3 contains Receiver Operating Characteristic (ROC) curves showing empirical sensitivity and specificity for classifying individuals as participants and the determination of their phenotype given knowledge of

participation. GWAS scenarios with a fixed subset of 1,000 and 5,000 cases and an equal number of controls are shown for varying numbers of randomly chosen or top associated SNPs. These ROC curves focus on high values of specificity with 1-specificity in the range of 0.05 to 10^{-6} on a logarithmic scale. Supplementary Figure 1 shows ROC curves for additional GWAS scenarios with 1,000-5,000 cases and an equal number of controls. Supplementary Figure 2 is analogous to Supplementary Figure 1 except showing ROC curves for the non-log scale for the full range of specificity. Supplementary Figure 3 is analogous to Supplementary Figure 1, except with ROC curves for T_{allele} .

Figure 3(a) shows a hypothetical GWAS with 1,000 cases and 1,000 controls. If the top 1,000 associated SNPs are published, the power to detect whether an individual is included in the study is approximately 43%, when the chance of a false positive, incorrectly concluding the individual is in the study, is 1 in 20. Power approaches zero when the false-positive rate is less than 1 in 1,000. If frequencies of 5,000 of the top associated SNPs are published, approximately 90% power is attained at a false-positive rate of 1 in 20 and 41% power when the false-positive rate is 1 in 1,000. When 50,000 of the top associated SNP frequencies are published, near perfect power is attained at a false-positive rate of 1 in 20 and approximately 98% power when the false-positive rate is 1 in 1,000.

We have shown the robustness of a novel test statistic, T_{geno} , an extension of the T_{allele} approach by Homer *et al.*6 adapted for use on genotype frequency data rather than SNP probe intensity or allele frequency data. We found that T_{geno} was more powerful than T_{allele} since the T_{geno} statistic conditions on known genotype states and is a likelihood-ratio statistic, and has optimality properties for both hypothesis testing and classification (see Figure 2 and Supplementary Figure 2).

This method makes a critical assumption that individuals in both groups are sampled from the same population. Unless population structure or substructure is identical in both groups, incorrect inferences are likely to be made. Essentially, T_{geno} and T_{allele} test whether an individual's genotypes are more consistent with one group versus another (see Methods for a formal description). These statistics are valid for determining membership only when the differences between groups are due to sampling variation. Otherwise, the relatively small bias due to membership of an individual quickly becomes swamped by the systematic differences in the underlying populations. Although this characteristic may be problematic in forensic applications in which test groups are often of unknown and potentially varied population origin, GWAS case and control groups are usually drawn from genetically comparable populations.

In addition to the number and selection method of SNPs, the sensitivity of our method, that is the power to detect true membership, is affected by several parameters: the number of subjects in each group, the number of SNPs tested, genotyping error, and genetic relatedness. As the size of the group containing the individual decreases, it becomes easier to detect the contribution of a single individual; as the size of the group without the individual increases, a more precise knowledge of the background population frequencies improves power. Because this method does not rely on patterns of rare or unique variation, power is not greatly affected by MAF for relatively common SNPs (MAF>5%) (see

Methods for details). Although substantial genotype error can decrease power, modest error rates (<5%) have little effect (current commercial genotyping platforms typically have error rates <0.1%). One can detect membership of siblings, parents and offspring, whereas the presence of more distant relatives would be more challenging to detect. For some applications, the ability to detect relatives is not desirable and could decrease specificity.

The conditional power to detect membership of an individual given a test and a reference sample is affected by sampling variation in the *estimates* of genotype frequencies between the two groups. The genetic variants need not have different underlying frequencies in the two groups, as might be a consequence of association with disease; only a non-zero difference in estimates is required. SNPs with low association p-values or, equivalently, large differences in estimated genotype frequencies between cases and controls, result in higher power because the magnitude of the ratio of frequencies between the two groups is larger. Two-sided association p-values are inversely correlated with the magnitude of the differences or provide additional information useful for detection of membership when using T_{geno} .

This method is particularly well-suited to case-control GWAS designs since subjects are partitioned into disjoint sets that provide both groups of frequency data required by this method. Publication of genotype frequencies separately for cases and controls also increases power to detect membership, since each group is typically a large fraction of the size of the overall study. GWAS are also often composed of a combination of multiple study populations and results may be published for each subset, increasing the power of this method, particularly for rare diseases that necessitate the collaboration of multiple studies. Similarly, re-use of sets of cases and controls among several GWAS can allow the calculation of frequencies for smaller groups by removing the contribution of overlapping individuals.

Further work is required to address some issues raised by our analyses. A more thorough understanding of the test statistic is required to account for linkage disequilibrium. Aggregate statistics other than frequencies could reveal membership information. It is notable that our approach could be applied to other types of dense datasets, both genetic and non-genetic. The underlying principle is based on detecting small correlations in categorical or continuous data accumulated over a sufficient number of highly repeatable outcomes.

We anticipate that these findings will aid in understanding the implications of publishing and sharing aggregate genomic data in ways that protect the privacy of research subjects. This method could be used to determine if specific individuals participated in a clinical study and perhaps influence the decision to enroll or continue participating. Our results should be considered as a lower bound for the power to detect membership and phenotype in an aggregate genotype dataset, as more efficient methods may yet exist. In light of these developments, the policies and practices guiding genomic data sharing should continue to evolve in order to promote quality science, minimize duplicative research, and merit the ongoing trust of the research subjects who consent to participate in scientific studies.

METHODS

Let $X_{i,g}$ and $Y_{i,g}$ denote genotype frequencies genotype g at locus i=1..s for two independent sets X and Y of DNA samples from unrelated individuals of size n and m, respectively. Let Z_i denote the genotype of an individual at locus i. We assume all loci are in linkage equilibrium, that samples were drawn from the same population with genotype frequencies $f_{i,j}$ for j=1..g_i, the number of genotypes that may be observed at each locus. We wish to test the ratio of frequencies of genotypes Z_i in groups X and Y and define a distance metric based on the log transformed ratio,

$$d = log\left(\frac{P\left(Z|X\right)}{P\left(Z|Y\right)}\right) = log\left(\prod_{i=1}^{s} \frac{X_{i,Z_{i}}}{Y_{i,Z_{i}}}\right) = \sum_{i=1}^{s} \left(logX_{i,Z_{i}} - logY_{i,Z_{i}}\right)$$

d is the sum of the differences in the natural logarithm of the frequency of individual Z's genotypes between the two groups. If individual Z is a member of a group X or Y, then the observed genotype count at each locus for Z's genotype has a contribution of 1 instead of the population frequency to the frequency statistic, and the resulting frequency so will tend to be slightly higher than that of the overall population by a factor of 1/n or 1/m, respectively. Notice that the distance measure d can also be motivated as a log-likelihood-ratio statistic for classifying individual Z's genotypes as "randomly" drawn from group X versus group Y.

We assume that genotypes are drawn from the same population with frequencies $f_{i,j}$, so that $X_{i,g}$ is distributed as Bin(n, $f_{i,g}$)/n and $Y_{i,g}$ is distributed as Bin(m, $f_{i,g}$)/m, where Bin(n,f) denotes the binomial distribution with n trials with success probability f. Throughout the calculations below we assume that mean and variance of $log(X_{i,g})$ and $log(Y_{i,g})$ can be well approximated by first order Taylor's approximation, given that in large samples we can assume $X_{i,g}$ and $Y_{i,g}$ are approximately distributed as normal variates.

Under the listed ideal assumptions and in sufficiently large samples, a T statistic can be utilized to test the significance of d:

$$\mathrm{T_{geno}}{=}\frac{\mathrm{d}{-}\mathrm{E_{H0}}\left(\mathrm{d}|\mathrm{Z}\right)}{\sqrt{Var_{H0}}\left(\mathrm{d}|\mathrm{Z}\right)}$$

A two-sided test of T_{geno} is appropriate when individual Z's DNA may be in either group. Otherwise, a one-sided test would be more powerful.

Consider four scenarios $(H_0 - H_3)$:

 H_0 : Individual Z's DNA is in neither group. Under this null model, the expected value of d is zero

$$\begin{split} \mathbf{E}_{\mathrm{H0}}\left(\mathbf{d}\right) &= \sum_{\mathrm{i}=1}^{\mathrm{s}} \left[\mathrm{E}\left\{ log\left(\frac{1}{\mathrm{n}}Bin\left(\mathbf{n},\mathbf{f}_{\mathrm{Z}_{\mathrm{i}}}\right)\right) \right\} - \mathrm{E}\left\{ log\left(\frac{1}{\mathrm{m}}Bin\left(\mathbf{m},\mathbf{f}_{\mathrm{Z}_{\mathrm{i}}}\right)\right) \right\} \right] \\ &\approx \sum_{\mathrm{i}=1}^{\mathrm{s}} \left[log\left(\mathbf{f}_{\mathrm{Z}_{\mathrm{i}}}\right) - log\left(\mathbf{f}_{\mathrm{Z}_{\mathrm{i}}}\right) \right] = 0 \end{split}$$

with variance

$$Var_{\rm H0}\left(d|Z\right) \approx \left(\frac{1}{n} + \frac{1}{m}\right) \sum_{i=1}^{s} \frac{1 - f_{Z_i}}{f_{Z_i}}$$

*H*₁: *Individual Z's DNA is in X and not in Y*. X is composed of n-1 genotypes observed by chance, but Z's genotype is known to be in X. Thus $X_i \sim (Bin(n-1,f_{z_i})+1)/n$ and the expected value of d given Z is

$$\begin{split} \mathbf{E}_{\mathrm{H1}}\left(\mathbf{d}|\mathbf{Z}\right) &= \sum_{i=1}^{\mathrm{s}} \left[\mathrm{E}\left\{ \log\left(\frac{1}{\mathrm{n}}\left(Bin\left(\mathrm{n}-1,\mathbf{f}_{\mathbf{Z}_{i}}\right)+1\right)\right)\right\} - \mathrm{E}\left\{\log\left(\frac{1}{\mathrm{m}}Bin\left(\mathrm{m},\mathbf{f}_{\mathbf{Z}_{i}}\right)\right)\right\} \right] \\ &\approx \sum_{i=1}^{\mathrm{s}} \log\left(1 + \frac{1 - \mathbf{f}_{\mathbf{Z}_{i}}}{\mathrm{nf}_{\mathbf{Z}_{i}}}\right) \approx \frac{1}{\mathrm{n}} \sum_{i=1}^{\mathrm{s}} \frac{1 - \mathbf{f}_{\mathbf{Z}_{i}}}{\mathbf{f}_{\mathbf{Z}_{i}}} \end{split}$$

 H_2 : Individual Z's DNA is in Y, but not X: Using an analogous argument to H_1 , the expected value of d given Z is

$$\begin{split} \mathbf{E}_{\mathrm{H2}}\left(\mathbf{d}|\mathbf{Z}\right) &= \sum_{i=1}^{\mathrm{s}} \left[\mathrm{E}\left\{ \log\left(\frac{1}{\mathrm{n}}Bin\left(\mathbf{n},\mathbf{f}_{\mathbf{Z}_{i}}\right)\right) \right\} - \mathrm{E}\left\{ \log\left(\frac{1}{\mathrm{m}}\left(Bin\left(\mathbf{m}-1,\mathbf{f}_{\mathbf{Z}_{i}}\right)+1\right)\right) \right\} \right] \\ &\approx -\sum_{i=1}^{\mathrm{s}} \log\left(1 + \frac{1 - \mathbf{f}_{\mathbf{Z}_{i}}}{\mathrm{m}\mathbf{f}_{\mathbf{Z}_{i}}}\right) \approx -\frac{1}{\mathrm{m}}\sum_{i=1}^{\mathrm{s}} \frac{1 - \mathbf{f}_{\mathbf{Z}_{i}}}{\mathbf{f}_{\mathbf{Z}_{i}}} \end{split}$$

H₃: Individual Z's DNA is in both X and Y. Then n-1 genotypes from X and m-1 genotypes from Y observed by chance, but Z's genotype is known to be in both. The expected value of d given Z is

$$\begin{split} \mathbf{E}_{\mathrm{H3}}\left(\mathbf{d}|\mathbf{Z}\right) &= \sum_{i=1}^{\mathrm{s}} \left[\mathbf{E}\left\{ log\left(\frac{1}{\mathrm{n}}\left(Bin\left(\mathrm{n}-1,\mathbf{f}_{\mathbf{Z}_{i}}\right)+1\right)\right)\right\} - \mathbf{E}\left\{ log\left(\frac{1}{\mathrm{m}}\left(Bin\left(\mathrm{m}-1,\mathbf{f}_{\mathbf{Z}_{i}}\right)+1\right)\right)\right\} \right] \\ &\approx -\sum_{i=1}^{\mathrm{s}} log\left(\frac{\mathrm{m}(\mathrm{n}-1)\mathbf{f}_{\mathbf{Z}_{i}}+\mathrm{m}}{\mathrm{n}(\mathrm{m}-1)\mathbf{f}_{\mathbf{Z}_{i}}+\mathrm{m}}\right) \approx \left(\frac{1}{\mathrm{n}}-\frac{1}{\mathrm{m}}\right)\sum_{i=1}^{\mathrm{s}}\frac{1-\mathbf{f}_{\mathbf{Z}_{i}}}{\mathbf{f}_{\mathbf{Z}_{i}}} \end{split}$$

This scenario is particularly relevant when X or Y represents a large population sample or perhaps even the whole population of interest.

Genotyping error

It is useful to consider the effect of a class of genotyping error in these models though we restrict our model to genotyping errors in Z that result in discordant genotype calls between Z and any group that may contain Z. We denote this genotype discordance rate ε . We utilize this error term to model genotype discordance when Z is a close relative of a member of X or Y.

For hypothesis 1, unless an error in genotyping occurred resulting in discordant genotypes for Z with probability ε , in which case $X_i \sim Bin(n-1,f_{z_i})/n$. Incorporating this non-zero possibility of genotyping error, $X_i \sim (Bin(n-1,f_{z_i})+Bin(1,1,-\varepsilon))/n$, the expected value of d given Z

$$\begin{split} \mathbf{E}_{\mathbf{H}\mathbf{1}}\left(\mathbf{d}|\mathbf{Z}\right) &= \sum_{\mathbf{i}=1}^{\mathbf{s}} \left[\mathbf{E}\left\{ \log\left(\frac{1}{\mathbf{n}}\left(Bin\left(\mathbf{n}-1,\mathbf{f}_{\mathbf{Z}_{\mathbf{i}}}\right) + Bin\left(1,1-\varepsilon\right)\right)\right) \right\} - \mathbf{E}\left\{ \log\left(\frac{1}{\mathbf{m}}Bin\left(\mathbf{m},\mathbf{f}_{\mathbf{Z}_{\mathbf{i}}}\right)\right) \right\} \right] \\ &\approx \sum_{\mathbf{i}=1}^{\mathbf{s}} \log\left(1 + \frac{1 - \mathbf{f}_{\mathbf{Z}_{\mathbf{i}}} - \varepsilon}{\mathbf{n}\mathbf{f}_{\mathbf{Z}_{\mathbf{i}}}}\right) \approx \frac{1}{\mathbf{n}} \sum_{\mathbf{i}=1}^{\mathbf{s}} \frac{1 - \varepsilon - \mathbf{f}_{\mathbf{Z}_{\mathbf{i}}}}{\mathbf{f}_{\mathbf{Z}_{\mathbf{i}}}} \end{split}$$

The effect of genotyping error for other hypotheses is similar.

Statistical Power

For large s, the test-statistic T_{geno} will approximately follow a standard normal distribution under H_0 given Z. Moreover, given the conditional null distribution being independent of Z, T_{geno} will also follow standard normal distribution unconditionally on Z under H_0 .

The *conditional* power of the test-statistics given Z will depend on the "non-centrality" parameter

$$\mathbf{E}_{\mathrm{H}} \mathbf{T}_{\mathrm{geno}} = NC_{\mathrm{H}} \left(\mathbf{Z} \right) = \frac{\mathbf{E}_{\mathrm{H}} \left(\mathbf{d} | \mathbf{Z} \right)}{\sqrt{Var_{\mathrm{H0}} \left(\mathbf{d} | \mathbf{Z} \right)}}$$

where the formulae for $E_H(d|Z)$ under various hypotheses are shown above. The *unconditional* non-centrality parameter can be calculated using the double expectation formula and the fact that $E((\Sigma_{i=1..s} w(Z_i))^{1/2}) \approx (\Sigma_{i=1..s} E(w(Z_i))+o(s))^{1/2}$ for any function w, assuming large s. The non-centrality parameter under H_1 is

$$\begin{split} \mathbf{E}\left[NC_{\mathrm{H1}}\left(\mathbf{Z}\right)\right] &= \mathbf{E}\left[\frac{\mathbf{E}_{\mathrm{H}}(\mathbf{d}|\mathbf{Z})}{\sqrt{Var_{\mathrm{H0}}(\mathbf{d}|\mathbf{Z})}}\right] \\ &\approx \sqrt{\mathbf{E}\left[\frac{\mathbf{m}}{\mathbf{n}(\mathbf{n}+\mathbf{m})}\sum_{i=1}^{\mathbf{s}}\frac{1-\mathbf{f}_{\mathbf{Z}_{i}}-\varepsilon}{\mathbf{f}_{\mathbf{Z}_{i}}}\right]} \\ &\approx \sqrt{\frac{ms\left(\bar{\mathbf{g}}(1-\varepsilon)-1\right)}{\mathbf{n}(\mathbf{n}+\mathbf{m})}} \end{split}$$

where $\overline{g} = \frac{1}{s} \sum_{j=1...s} g_{j}$, the average number of possible genotype categories over all loci. There are three possible genotype categories per locus ($\overline{g} = 3$) when T_{geno} is computed from biallelic SNP data with MAF sufficiently high to ensure that all genotypes are observed. Assuming normality of the test-statistic and that the variance of under H_1 remains approximately the same as under H_0 , the unconditional power of a two-sided test of T_{geno} (which assumes Z may be a member of X or Y) under H_1 at significance level α is

$$\operatorname{power}_{_{\mathrm{H1}}}\left(\alpha\right) = Pr_{_{\mathrm{H1}}}\left(|\mathbf{T}_{\mathrm{geno}}| > \mathbf{T}_{\alpha/2}\right) \approx \Phi\left(\Phi^{-1}\left(\alpha/2\right) + \sqrt{\frac{ms\left(\bar{\mathbf{g}}\left(1-\varepsilon\right)-1\right)}{n\left(n+m\right)}}\right)$$

where Φ is the standard normal cumulative distribution function and Φ^{-1} its inverse. Power may be derived under the other alternative hypotheses in a similar fashion. The power of a one-sided test, one that assumes Z is not a member of Y, substitutes α for $\alpha/2$ in the righthand side of previous equation. Use of low frequency SNPs (e.g., MAF<5%) will reduce \overline{g} and, as a consequence, the power of the test.

Modeling Relatives

The genotype discordance rate ε can be used to represent other sources of discordance other than that due to genotyping error. Genotype concordance among a pair of relatives is equivalent to the probability of sharing two alleles identical by state (IBS), which are functions of the relationship type and minor allele frequencies for independent loci in Hardy-Weinberg equilibrium. The probability of not sharing two alleles identical by state for parent offspring pairs is

$$P(IBS \neq 2|Parent - Offsping) = 1 - p^4 - q^4 - pq(p^2 + q^2 + 1)$$

and for sibling pairs is

$$P(IBS \neq 2|sibling) = 1 - p^4 - q^4 - 2pq(p^2 + \frac{7}{4}pq + q^2)$$

where p is the minor allele frequency and q=1-p is the major allele frequency.

Relaxing assumptions

Independence of X and Y: Similar results hold when at least one member of X and one member of Y are the same person or close relatives. The expected values of d under each case are unchanged but the variances are reduced depending on the amount of overlap as the effective sizes of X and Y are reduced.

Independence of loci: The assumption of linkage equilibrium is currently necessary to derive analytical expectations of the moments and power of our test statistic. When applying this method to empirical data, many of the model assumptions needed to precisely determine the null distribution of the test statistic and determine significance by comparison to a theoretical distribution no longer hold. Thus it seems sensible to shift perspective from a hypothesis testing framework to one of classification, which examines the trade-off between sensitivity (true positive rate) and specificity (true negative rate) across a range of cut-off values for the statistic.

When applied to empirical GWAS data, the distribution of T_{geno} need not be centered at zero (μ_0 0) and the variance is greater than would be expected (see Figure 1). The lack of centrality is likely due to sampling variation, while the variance is inflated due primarily to correlation among loci from linkage disequilibrium.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Histogram of $T_{\mbox{geno}}$ for a GWAS with 1,000 cases and controls

The figure presents data using 1,000 cases (group 1 in red), 1,000 controls (group 2 in blue) and 1,000 subjects not in the study based on genotypes from Illumina HumanHap550 assay. The theoretical null density curve is shown in black.



Figure 2. Histograms of calibrated $\rm T_{geno}$ and Homer's $\rm T_{allele}$ with 1,000 cases and controls and varying numbers of SNPs

The figure presents theoretical null density curves (black) for a GWAS with 1,000 cases (group 1 in red), 1,000 controls (group 2 in blue) and 12,000 subjects not in the study (in gray) using genotypes for (a) 10,000, (b) 100,000, and (c) 550,000 top associated SNPs from the Illumina HumanHap550 assay. Statistics were calibrated so that the null distribution was centered at zero with unit variance.



Figure 3. Sensitivity and specificity of T_{geno} applied to GWAS data

Log-scale Receiver Operating Characteristic (ROC) curves of T_{geno} with Illumina HumanHap550 data from GWAS scenarios with 1000/1000 and 5000/5000 cases and controls of European descent.

Table 1

Test group.
the
in
individual
an
detect
5
power of T _{genc}
Theoretical

a, 1.0	00 samples/or	oun, number	of loci varied		Po	wer of	Lane at 9	Significs	ance Le	vel
Test group	Ref group	Ind. SNPs	Error rate	MAF	0.05	0.01	0.001	10-4	10^{-5}	10^{-6}
1,000	1,000	5,000	0.00	0.25	0.62	0.38	0.15	0.05	0.02	0.00
1,000	1,000	10,000	0.00	0.25	0.89	0.73	0.45	0.24	0.11	0.04
1,000	1,000	25,000	0.00	0.25	1.00	0.99	0.96	0.86	0.72	0.54
1,000	1,000	50,000	0.00	0.25	1.00	1.00	1.00	1.00	1.00	0.99
1,000	1,000	75,000	0.00	0.25	1.00	1.00	1.00	1.00	1.00	1.00
b. 5,0	00 samples/gr	oup, number.	of loci varied		Po	wer of .	F _{geno} at 9	Significs	ance Le	vel
Test group	Ref group	Ind. SNPs	Error rate	MAF	0.05	0.01	0.001	10^{-4}	10^{-5}	10^{-6}
5,000	5,000	10,000	0.00	0.25	0.23	0.09	0.02	0.00	0.00	0.00
5,000	5,000	25,000	0.00	0.25	0.55	0.31	0.12	0.04	0.01	0.00
5,000	5,000	50,000	0.00	0.25	0.92	0.78	0.53	0.30	0.15	0.07
5,000	5,000	75,000	0.00	0.25	0.98	0.91	0.74	0.52	0.32	0.17
5,000	5,000	100,000	0.00	0.25	1.00	0.98	06.0	0.75	0.56	0.38
5,000	5,000	150,000	0.00	0.25	1.00	1.00	0.98	0.93	0.83	0.69
5,000	5,000	200,000	0.00	0.25	1.00	1.00	1.00	0.99	0.97	0.92
c. 1,000	samples/grou	ıp, genotype e	rror rate vari	ed	Po	wer of 7	F _{geno} at 9	Significa	ance Le	vel
Test group	Ref group	Ind. SNPs	Error rate	MAF	0.05	0.01	0.001	10^{-4}	10^{-5}	10^{-6}
1,000	1,000	50,000	0.00	0.25	1.00	1.00	1.00	1.00	1.00	0.99
1,000	1,000	50,000	0.01	0.25	1.00	1.00	1.00	1.00	0.99	0.98
1,000	1,000	50,000	0.05	0.25	1.00	1.00	1.00	1.00	0.98	0.94
1,000	1,000	50,000	0.10	0.25	1.00	1.00	1.00	0.98	0.93	0.84
1,000	1,000	50,000	0.25	0.25	0.99	0.95	0.81	0.61	0.40	0.24
d. 1,000 s	amples/group), minor allele	frequency va	ried	Po	wer of 7	F _{geno} at 9	Significa	ance Le	vel
Test group	Ref group	Ind. SNPs	Error rate	MAF	0.05	0.01	0.001	10^{-4}	10^{-5}	10^{-6}
1,000	1,000	25,000	0.00	0.05	1.00	1.00	0.97	0.88	0.72	0.52
1,000	1,000	25,000	0.00	0.10	1.00	0.99	0.96	0.87	0.73	0.55

u. 1,000 34	mples/group	, minor allele	frequency va	riea	20	wer of .	I geno at 2	Signific	ance Le	vel
Test group	Ref group	Ind. SNPs	Error rate	MAF	0.05	0.01	0.001	10^{-4}	10^{-5}	10^{-6}
1,000	1,000	25,000	0.00	0.20	1.00	0.99	0.96	0.87	0.72	0.54
1,000	1,000	25,000	0.00	0.30	1.00	0.99	0.96	0.87	0.72	0.55
1,000	1,000	25,000	0.00	0.40	1.00	0.99	0.96	0.87	0.72	0.54
1,000	1,000	25,000	0.00	0.50	1.00	0.99	0.96	0.87	0.72	0.54

e. 1,000	samples/grou	p, reference g	roup size var	ied	Por	wer of	F _{geno} at S	Significa	ance Le	vel
Test group	Ref group	Ind. SNPs	Error rate	MAF	0.05	0.01	0.001	10^{-4}	10^{-5}	10^{-6}
1,000	60	25,000	0.00	0.10	0.20	0.07	0.02	0.00	0.00	0.00
1,000	100	25,000	0.00	0.10	0.46	0.23	0.07	0.02	0.00	0.00
1,000	200	25,000	0.00	0.10	0.74	0.51	0.24	0.09	0.03	0.01
1,000	300	25,000	0.00	0.10	0.92	0.78	0.51	0.27	0.12	0.05
1,000	400	25,000	0.00	0.10	0.97	0.88	0.68	0.43	0.23	0.11
1,000	500	25,000	0.00	0.10	0.98	0.93	0.78	0.56	0.35	0.19
1,000	750	25,000	0.00	0.10	1.00	0.98	0.91	0.77	0.58	0.39
1,000	1,000	25,000	0.00	0.10	1.00	0.99	0.96	0.87	0.73	0.55
1,000	2,000	25,000	0.00	0.10	1.00	1.00	0.99	0.97	0.92	0.82
1,000	5,000	25,000	0.00	0.10	1.00	1.00	1.00	1.00	0.98	0.95
1,000	10,000	25,000	0.00	0.10	1.00	1.00	1.00	1.00	0.99	0.97
scenarios exar	nined: (a) Hyp	othetical GWA	AS with 1,000	cases (tes	t group)	and 1,(000 conti	ols (ref	erence g	roup), no

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o genotyping error, and 5,000 to 200,000 independent SNPs with fixed MAF of 25%. The European descent. (b) As for (a) with 5,000 cases and 5,000 controls. (c) As for (a) with genotype discordance rates from 0% to 25% and 50,000 independent loci. (d) As for (a) with varying MAF from 5% to 50% for 25,000 independent loci. (e) For varying sizes of the reference group from 60 (the size of the HapMap CEU founder population) to 10,000 (near perfect estimation of the genotype frequencies) populations of European descent. A MAF of 25% was chosen based on a survey of several fixed-content assays which were found to have average MAFs ranging from 20% to 25% in populations of upper bound on the number of SNPs was chosen based on estimates (unpublished) that the Illumina HumanHap550 assay provides information equivalent to ~200,000-300,000 independent SNPs in for a fixed MAF of 10%. Author Manuscript

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Test group	Ref group	Ind. SNPs	Parent/ Offspring Discordance	MAF	0.05	0.01	0.001	10^{-4}	10^{-5}	10^{-6}
1,000	1,000	200,000	0.10	0.05	1.00	1.00	1.00	1.00	1.00	1.00
1,000	1,000	200,000	0.18	0.10	1.00	1.00	1.00	1.00	1.00	1.00
1,000	1,000	200,000	0.32	0.20	1.00	1.00	1.00	1.00	1.00	0.99
1,000	1,000	200,000	0.42	0.30	1.00	1.00	0.97	0.91	0.79	0.63
1,000	1,000	200,000	0.48	0.40	0.98	0.92	0.77	0.55	0.35	0.20
1,000	1,000	200,000	0.50	0.50	0.93	0.82	0.58	0.35	0.19	0.09
a	. 1,000 sampl	<u>es/group, det</u>	ection of a sibling		Pov	ver of T	geno at S	ignific	ince Le	vel
Test group	Ref group	Ind. SNPs	Parent/ Offspring Discordance	MAF	0.05	0.01	0.001	10-4	10 ⁻⁵	10^{-6}
1,000	1,000	200,000	0.092	0.05	1.00	1.00	1.00	1.00	1.00	1.00
1,000	1,000	200,000	0.168	0.10	1.00	1.00	1.00	1.00	1.00	1.00
1,000	1,000	200,000	0.282	0.20	1.00	1.00	1.00	1.00	1.00	1.00
1,000	1,000	200,000	0.354	0.30	1.00	1.00	1.00	1.00	0.99	0.96
1,000	1,000	200,000	0.394	0.40	1.00	1.00	0.99	0.97	0.92	0.83
1,000	1,000	200,000	0.406	0.50	1.00	1.00	0.99	0.95	0.87	0.74