Using Linkage Genome Scans to Improve Power of Association in Genome Scans

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Scanning the genome for association between markers and complex diseases typically requires testing hundreds of thousands of genetic polymorphisms. Testing such a large number of hypotheses exacerbates the trade-off between power to detect meaningful associations and the chance of making false discoveries. Even before the full genome is scanned, investigators often favor certain regions on the basis of the results of prior investigations, such as previous linkage scans. The remaining regions of the genome are investigated simultaneously because genotyping is relatively inexpensive compared with the cost of recruiting participants for a genetic study and because prior evidence is rarely sufficient to rule out these regions as harboring genes with variation of conferring liability (liability genes). However, the multiple testing inherent in broad genomic searches diminishes power to detect association, even for genes falling in regions of the genome favored a priori. Multiple testing problems of this nature are well suited for application of the false-discovery rate (FDR) principle, which can improve power. To enhance power further, a new FDR approach is proposed that involves weighting the hypotheses on the basis of prior data. We present a method for using linkage data to weight the association $P$ values. Our investigations reveal that if the linkage study is informative, the procedure improves power considerably. Remarkably, the loss in power is small, even when the linkage study is uninformative. For a class of genetic models, we calculate the sample size required to obtain useful prior information from a linkage study. This inquiry reveals that, among genetic models that are seemingly equal in genetic information, some are much more promising than others for this mode of analysis.

Methods to detect liability alleles for complex disease are at a crossroads. Previously, tests of association between disease status and alleles at specific markers targeted several markers within a handful of candidate genes. With the advent of relatively inexpensive molecular methods for genotyping, the trend is moving toward whole-genome association studies (Thomas et al. 2005). Such investigations might involve hundreds of thousands of markers and hypothesis tests. Current analytical tools for such massive investigations are limited. The ultimate success of whole-genome association studies will depend largely on the development of innovative analytic strategies.

Although the whole genome might be tested for association with the disorder of interest, typically, some regions of the genome are favored because of prior investigations or knowledge of the biological function of particular genes. Other regions are investigated simultaneously because genotyping is relatively inexpensive compared with the cost of recruiting study participants and the cost of designing large-scale study-specific assays. Moreover, for complex phenotypes, few regions can be excluded as not harboring liability genes.

For any well-calibrated statistical procedure, simultaneously looking for association across the whole genome leads to a loss in power to detect signals in a specified list of genes. Some statistical approaches, however, are better suited to large-scale testing than others. For instance, if we aim to control the false-discovery rate (FDR), defined as “the expected fraction of false rejections,” the loss in power will be less than if we aim to control the type I error rate (Genovese and Wasserman 2002; Benjamini and Hochberg 1995). For this reason, FDR procedures are widely used in multiple-testing problems that arise in genetic studies (Efron and Tibshirani 2002; Devlin et al. 2003a, 2003b; Sabatti et al. 2003; Storey and Tibshirani 2003).

To enhance the power to detect association in favored regions while simultaneously testing the whole genome, we considered an FDR procedure that facilitates weighting hypotheses on the basis of prior information (Genovese et al. [in press]; see also Benjamini and Hochberg [1997]). Specifically, our goal was to increase the power to detect signals for association in preselected regions by judiciously “up-weighting” the relevant $P$ values while “down-weighting” $P$ values in all other regions. Linkage data offer a natural choice for weighting the $P$ values for a whole-genome association study. In the present study, we investigated how linkage traces can be formulated as weights. If some of the loci under linkage peaks are indeed more likely to be associated with the phenotype than
the others, then the weighted procedure offers a chance to maintain good power even in a whole-genome association study. Remarkably, we will show that if the linkage trace is informative, then power is enhanced; yet, with uninformative linkage traces, the procedure experiences a fairly small loss in power. We also provide guidance on the type of genetic models and required sample sizes likely to yield informative weights. Although we focused our investigation on weighted FDR procedures, these ideas can be easily applied to other multiple-testing procedures, such as Bonferroni and Holm’s procedure (Holm 1979).

Methods

The proposed multiple-testing situation consists of m hypotheses \( H_1, \ldots, H_m \), for which \( H_i = 1 \) if the \( i \)th null hypothesis is false, and 0 otherwise. The evidence corresponding to the \( i \)th test for association is summarized in the \( P \) value \( P_i \). The ordered \( P \) values are indicated as \( P_{[1]} \leq P_{[2]} \leq \cdots \leq P_{[m]} \), with \( P_{[i]} \) defined to be 0 for convenience.

Many FDR-based procedures are based on the following pattern: reject each hypothesis for which \( P_i \leq T \) when the threshold \( T \) is selected on the basis of the observed \( P \) values so as to maintain FDR at level \( \alpha \). The FDR at threshold \( T \) is defined to be the expectation of

\[
\frac{\text{no. of false rejections}}{\text{no. of rejections}} = \frac{\sum_{i=1}^{m} I(P_i \leq T)(1 - H_i)}{\sum_{i=1}^{m} I(P_i \leq T)},
\]

where the ratio is defined to be 0 when the denominator is 0. The most common procedure for choosing \( T \) is that of Benjamini and Hochberg (1995) and is given by

\[
T = \max \left\{ P_i; P_i = \frac{\alpha i}{m} \right\}.
\]

Let \( w_1, \ldots, w_m \) be the chosen weights. Following the argument for the Benjamini and Hochberg (1995) FDR procedure, the “\( w \)BH” procedure of Genovese et al. (in press) finds the threshold for rejection (\( T \)) that controls the FDR at rate \( \alpha \) when the weighted values \( \{P_i/w_i\} \) play the role of the \( P \) values. In practice, the weights adjust the threshold for rejection individually for each \( P \) value, in that we reject the \( i \)th hypothesis if \( P_i \leq w_i T \).

To illustrate the effect of weights, we present a theoretical exploration of the power of weighted tests that would have had 50% power in the absence of weighting. Tests with considerably more power will not benefit substantially from weighting, and tests with less power are not likely to be powerful even with weights. We call these 50%-power alternatives “marginal” because they are on the margin of detectability. In this context, power is defined as the probability that \( P_i/w_i \leq T \).

To maintain FDR at a fixed level, a set of prior weights \( \{w_i\} \) must satisfy two criteria: \( w_i \geq 0 \) and

\[
\overline{w}_w = \frac{1}{m} \sum_{i=1}^{m} w_i = 1
\]

( Genovese et al., in press). Thus, candidates for linkage-based weights are numerous. For simplicity of exposition, we initially consider binary weights with a fraction \( \epsilon \) of weights \( w_1 = B/(B \epsilon + 1 - \epsilon) \)—that is, up-weighted—and the remainder \( w_2 = 1/(B \epsilon + 1 - \epsilon) \) down-weighted. In figure 1, we compare the achieved power over a grid of choices for the two free parameters, \( \epsilon = (0.001,0.01,0.1) \) and \( B = (2,6,50) \). Within each bar plot, the left (right) bar reveals the power if the value is up-weighted (down-weighted). Thus, the bars show the deviation from the marginal unweighted power (0.5) and the contrast in power if a test is correctly up-weighted or incorrectly down-weighted. For a given \( \epsilon \), the power gain for the up-weighted tests increases as \( B \) increases. Meanwhile, the down-weighted tests lose increasingly more power as \( B \) increases; however, the loss is disproportionately smaller than the gain attained by the up-weighted tests. The potential gains are more striking if \( \epsilon \) is small, whereas the potential losses are greater if \( \epsilon \) is large. In effect, as \( \epsilon \) increases, constraining the average weight to be 1 results in more-modest up-weights and more-severe down-weights. This reduces the power to detect a given signal, regardless of whether it is up- or down-weighted. This analysis reveals that sparse, dramatic weights can yield large increases in power with very little potential loss. There is a catch, though. This striking improvement in power is attainable only if the weights are correctly placed. With small
of the unknown quantity $h$, it is much more challenging to place the weights correctly. A causal variant is more likely to be up-weighted if $\epsilon$ is larger.

Next, we considered the effect of weights on the number of discoveries made. Consider a situation in which $H = 1$ for a fraction $0 < a < 1$ of the tests, and use the binary weights described above. It is convenient for this analysis to treat the weights as random variables. A measure of the informativeness of the betting is

$$
\eta = \frac{Pr(W = w_1 | H = 1)}{Pr(W = w_1 | H = 0)}.
$$

When $\eta = 1$, $W$ and $H$ are independent; for $\eta > 1$, there is greater likelihood of betting correctly, and, for $0 < \eta < 1$, incorrectly. With use of marginal alternatives, half of the tests with $H = 1$ are likely to be discovered. We measure our genomewide success rate by counting how many true discoveries are made in excess of expectation. Figure 2 considers a scenario of the number of excess discoveries increases smoothly with $\eta$ (short-dashed line). When $\eta = 2$, the number of excess discoveries increases smoothly with $\eta$ (dotted line). For $\eta$ close to 1, a choice of $\epsilon < a$ leads to more discoveries than obtained when $\eta = a$; however, this advantage is lost as $\eta$ increases (dot-dashed line = 0.2; long-dashed line = 0.4). From this analysis, we conclude that an optimal choice of $\epsilon$ is slightly $> a$.

For our application, the value of the linkage test at the position of the $i$th association test $z_i$ is the basis for the proposed weighting system. For linkage data, the test statistics along a chromosome can be summarized as a Gaussian process that, at any point on the chromosome, is approximately normally distributed. Under the null hypothesis, the distribution has mean 0 and variance 1. Unlike the binary scenario investigated thus far, the test statistics are strongly correlated within a chromosome. Even under the alternative hypothesis, a linkage peak for a complex disease tends to be broad and ill defined (fig. 3; blue) because of the physical process of recombination. It is desirable to up-weight the entire region under a linkage peak to capture the causal variant (or variants). Because linkage results are quantitative, and because a linkage peak is not a well-defined region, we consider continuous weights.

We use the following heuristic to motivate our exponential weighting scheme. Assume the model-free linkage statistics are distributed normally $(\beta,1)$, with mean $\beta = 0$ when unlinked and $\beta > 0$ when linked. A natural weighting candidate is the posterior odds that an observation is linked, which is proportional to $v_i = e^{\beta z_i}$. Because the weights are constrained to have mean 1, $w_i = v_i/\sum_i v_i$ is a valid choice. Thus, choosing the weights is tantamount to choosing a constant $B \geq 0$ to play the role of the unknown quantity $\beta$.

The exponential weighting scheme with $B = 1$ is depicted in figure 3 (green). The implicit meaning of $\epsilon$ in the binary weighting scheme is determined automatically by the number and length of the linkage peaks. If there are many peaks, then the up-weighting will be constrained more strongly by the requirement that $\bar{w} = 1$. Increasing $B$ decreases the length of the peak that is strongly up-weighted. Consequently, whereas a large value of $B$ will yield greater power in some instances, it might fail to up-weight a causal variant located away from the crest of the corresponding linkage peak.

Exponential weights have the disadvantage of being highly sensitive to large values, a feature that is particularly apparent in the top panel of figure 3. For this reason we considered cumulative weights, defined as $\phi(z - B)$, where $\phi(z)$ is the standard normal cumulative distribution function. This function has the desirable property of increasing approximately linearly for values of $z$ near $B$, but quickly reaching an asymptote for large values of $|z - B|$. Consequently, it gives approximately equal up-weighting (down-weighting) to any $z$ value 2 or more units above (below) $B$. See figure 3 (red) for a comparison between the exponential and cumulative weights; $B = 1$ for the former and $B = 2$ for the latter. Cumulative weighting has the advantage of providing smoother up-weights and broader peaks. Its disadvantage is that it strongly down-weights regions without linkage signals, perhaps making it an inferior choice when the linkage input is uninformative because of low power. This feature is most apparent in the bottom panel of figure 3.

**Results**

**Simulations**

To evaluate wBH, we simulate whole-genome linkage traces (22 autosomes and chromosome X) and corresponding association tests. We simulate linkage traces that approximate the information obtainable from an independent 10,000-SNP linkage study. Because of practical constraints on computation time, we simulate the linkage traces indirectly. As described by Bacanu (2005), we approximate linkage traces on chromosomes as independent realizations of an underlying time series process. The linkage trace is constructed by adding random
noise to a deterministic model of the linkage signal. The random noise of a linkage trace is distributed as a Gaussian process from an autoregressive moving average (ARMA 2,1) model with parameters $a_1 = 1.51$, $a_2 = -0.51$, and $ma_1 = 0.22$ (Venables and Ripley 2002). This model was selected on the basis of features expected in actual linkage traces (see appendix A for details). To introduce $L$ linkage signals, the position of the disease variant is randomly placed on $L$ chromosomes, one signal per chromosome. Each expected signal is centered at the designated location, with peak height $\mu$, which decays smoothly to zero as the distance $t$ from the disease variant increases. If the autocorrelation pattern determined by the ARMA model is $\rho(t)$, then a marker $t$ units from the disease variant has signal equal to $\rho(t)\mu$. Chromosomes with no disease variants have signal equal to zero at all points. The resulting traces resemble the outcome of a linkage study that relies on sib pairs, analyzed using a model-free linkage analysis, for a complex genetic disease.

Superimposed on the linkage traces are $m = 500,000$ (250,000) simulated association statistics, equally spaced in genetic distance. To realistically simulate a dense set of association data, detailed information about the linkage disequilibrium (LD) structure between SNPs would be required. Such a detailed simulation, however, would be impractical for this investigation. Consequently, we simulated independent association tests. The $P$ values for the noncausal association statistics are uniformly distributed. Each of the $P$ values obtained from one of the $L$ disease variants is derived from a normal $N(\mu, 1)$ test statistic.

Our simulations explore various levels of signals that might be encountered in practice under a wide variety of genetic models. The intensity of the signal is a statistical parameter, not immediately interpretable via the genetic model. The quantities $\mu$ and $\mu_\alpha$ represent the shift in the mean of the linkage and association test statistics, respectively, between the unlinked and linked hypotheses. The shift is determined by the informativeness of the genetic model for the designated type of study and the size of the sample. Theoretically, any desired combination of linkage and association shifts could be attained with appropriate sample sizes in each type of study. However, for some genetic models, the sample size required might be immense for a linkage and/or association study. Below, we discuss further the connection between the statistical parameters and the genetic models.

For each of $>50$ simulation conditions, which correspond to specific levels of $\mu$ and $\mu_\alpha$, we generated 100

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Figure 3  Plot of linkage traces for three chromosomes with weights. The green and red traces are weights with exponential ($B = 1$) and cumulative ($B = 2$) weights, respectively, that are based on the linkage trace (blue). The bottom panel has no signal.
linkage traces and 10 replicate association studies per linkage trace. Consequently, at each level, the performance of the inference procedures were estimated from 1,000 simulations. In addition, of \( m = 500,000 \) association tests, \( L = 10 \) disease variants were present. We also tried \( L = 5 \) and 15 of \( m = 250,000 \) and 500,000 tests, but the results for those conditions were qualitatively similar to those for \( L = 10 \) and \( m = 500,000 \), so those results are not shown.

Simulations displayed in figure 4 show that solid gains in power are possible with wBH (dashed lines) as compared with BH (solid line) and especially compared with the traditional Bonferroni correction for multiple testing (dotted line). For signals arising from weak, moderate, and strong linkage signals \( (\mu_i = 1.0, 2.5 \) and 3.5—dashed lines, from lowest to highest, respectively), the increase in power is steady across the range of association signals \( (\mu_p = 4.0–6.0) \). The increase when \( \mu_i = 1 \) is encouraging. With such a weak signal, the linkage trace is likely to appear to have no meaningful signal. It is also noteworthy that moving from good \( (\mu_i = 2.5) \) to excellent \( (\mu_i = 3.5) \) information does not result in a dramatic improvement in power. There appears to be a limit to how much power can be obtained from linkage-based weights.

The linkage shift of \( \mu_i = 0 \) is intended to approximate a linkage study that has essentially no power because of insufficient sample size. As noted by Risch and Merikangas (1996), this scenario is certainly plausible. The simulations reveal that a fairly small amount of power is lost because of weights that are based on an uninformative linkage study \( (\mu_i = 0) \) (fig. 4; dot-dashed line). Even with uninformative weights, the power of wBH is greater than that obtained using a Bonferroni correction. In the best-case scenario, increases in power over Bonferroni-corrected tests can be >35%. The wBH results were obtained using exponential weights with \( B = 1 \). Very similar results were obtained when cumulative weights with \( B = 2 \) were used. Exponential weights slightly outperformed cumulative weights when \( \mu_i = 0 \).

We also investigated other choices of constants for the weighting schemes. In practice, we found that good choices for the arbitrary parameters are \( B = 1 \) for exponential weighting and \( B = 2 \) for cumulative weighting, because they provide a substantial gain in power when the linkage trace is informative and a small loss in power when the linkage trace is uninformative.

Another interesting feature of the simulations is the behavior of the linkage signals for individual genome scans. Although there were \( L = 10 \) loci that could produce linkage signals, for any particular simulation, only a few of those loci were likely to produce a notable linkage signal, even when \( \mu_i = 3.5 \). In fact, for a commonly accepted bound for significant linkage—a LOD of 3.6 (Lander and Kruglyak 1995) and \( \mu_i = 1.0, 2.5 \), and 3.5—the expected numbers of loci exceeding the threshold are 0.01, 0.6, and 2.8 of 10; for suggestive linkage—requirement of a LOD of 2.2 and \( \mu_i = 1.0, 2.5 \), and 3.5—the expected numbers of loci exceeding the threshold are 0.1, 2.5, and 6.2 of 10. Naturally, when \( \mu_i = 0.0 \), the expected number of loci exceeding either bound is essentially 0 of 10.

The size of the weights varies as a function of many factors, including characteristics of the linkage trace (the number, width, and height of the peaks), the choice of weight function, and related parameter \( B \). To provide insight into to the size of weights in our simulations, we summarized the mean and variance of the weights derived from the various types of linkage traces evaluated at the 10 causal variants (table 1). When there was no notable signal in the linkage data, the weights were approximately equal to unity, on average, which is a desirable feature. As the linkage signal increased, so did the weights (table 1). The increase was more dramatic for exponential weights than for cumulative weights. Notably, the exponential weights were more variable than

![Figure 4](https://example.com/figure4.png)

**Figure 4**  Power as a function of \( \mu_i \). Power is defined as the number of true discoveries of \( L = 10 \) causal variants. Distinct lines correspond to different methods and/or linkage signals, coded as follows (from lowest to highest power): dotted is the Bonferroni method; dot-dashed is wBH through use of linkage data with no signal \( (\mu_i = 0) \); solid is the BH method (Storey’s 2002 version); dashed are wBH through use of linkage data with a weak \( (\mu_i = 1) \); moderate \( (\mu_i = 2.5) \), and strong \( (\mu_i = 3.5) \) signal.

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the cumulative weights, which suggests the cumulative weights could be a better choice.

Relating Genetic Models to Statistical Models

The intensity of the signals from linkage and association studies are determined by the underlying genetic model and the sample size. With some statistical manipulation, the expected signal resulting from the genetic model can be parameterized by the statistical shift parameters \( \mu_L \) and \( \mu_A \) used in our simulations. In this section, we develop a mapping between the statistical parameters and the genetic models. This mapping should be useful for evaluating the results of our simulations and for the design of whole-genome association studies, especially realistic evaluation of their power.

For a given sample size and genetic model, we can compute the resulting shift of the linkage and/or association test statistic by making use of two statistical relationships. For simplicity of exposition, we assume that an observation in a linkage study is a fully informative affected sib pair (ASP). In an association study, an observation is a single case and a single control. For a particular genetic model and study design, a certain amount of information is expected per observation, and the sum of this information over observations is called the “noncentrality parameter.” This quantity determines the shift parameter. Next, a regression model can summarize the functional relationship between a genetic model and the sample size required to achieve a particular size of shift for a particular study design.

We investigate a variety of genetic models that assume a prevalence of \( \kappa \) (0.01–0.03), a disease variant that occurs in the population with frequency \( \rho \) (0.05–0.5), and an additive penetrance model. Models vary in the levels of genetic effect assumed. To obtain small-to-moderate genetic effects, we consider models with an odds ratio \( (\phi) \) of 1.5–3.0. This quantity indexes the genetic effect of the disease variant independent of the prevalence \( \kappa \); however, as with all measures of genetic impact, the strength of a given value can be interpreted only within the context of the other parameters in the genetic model. Some measures of genetic effect are more useful for linkage analysis, whereas others are more useful for association analysis. For this reason, we also report the risk to relatives (\( \lambda \)) (Risch 1990) and the attributable fraction (\( \Delta \)) (Pfeiffer and Gail 2003). The latter, which is equivalent to Levin’s population-attributable risk, is defined to be one minus the ratio of the probability of affection, given that an individual has no copies of the detrimental allele, over the prevalence.

To see how the apparent genetic effect varies with the measure, consider two models, each with \( \kappa = 0.01 \) and \( \phi = 2 \). If the causal variant is uncommon (\( \rho = 0.05 \)), \( \Delta = 9\% \) and \( \lambda = 1.04 \), whereas, if the causal variant is common (\( \rho = 0.5 \)), the apparent genetic effect is much stronger: \( \Delta = 66\% \) and \( \lambda = 1.11 \). From \( \Delta \) and \( \lambda \), we see that, when \( \rho \) is small, it will be difficult to detect the signal if \( \phi \leq 2 \), because most of the risk will not be attributable to the locus.

The number of ASPs (\( n_L \)) necessary to achieve a signal of size \( \mu_L = 2.0 \) in a linkage analysis with fully informative markers is given for a variety of genetic models (table 2). For each model, the corresponding number of cases (\( n_A \)) necessary to achieve a signal of size \( \mu_A = 4.5 \) in a case-control association study with equal numbers of cases and controls is given, with the assumption that the disease variant is measured. Notice that some genetic models are favorable to association studies, whereas others are favorable to linkage studies. For instance, when \( \kappa = 0.01 \), \( \phi = 2 \), and \( \rho = 0.05 \), the required sample size for a linkage study is large (\( n_L = 5,664 \)), yet, for an association study, it is tenfold smaller (\( n_A = 648 \)).

Table 2 shows the range of the genetic model space we explored, but only a fraction of the specific settings. To obtain estimates of the sample sizes required for linkage and association studies for other genetic models, we provide a regression model that is a function of \( (\rho, \kappa, \phi) \). See appendix B for details. For both regression models, the fit was excellent (\( R^2 > 99\%) \).

For a given genetic model defined by \( (\rho, \kappa, \phi) \), let \( N_L \) and \( N_A \) denote the sample size required to obtain shift parameters of \( \mu_L = 2.0 \) and \( \mu_A = 4.5 \), respectively. We chose these shift parameters for illustrative purposes only. It is straightforward to extend to other desired shift parameters. If \( n_L \) is the sample size necessary to obtain a

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linkage shift of size $c$, then to obtain a shift of size $b$, we require a sample size of

$$n_b = n_c \left( \frac{b}{c} \right)^2.$$ 

From the regression model and for $c = 2.0$, $n_c$ is well approximated by $N_c$. Substituting this in the formula yields the desired sample-size estimate for the given genetic model.

For an association study, it is quite likely that the causal SNP is unmeasured. Let $r^2$ denote the Pearson squared correlation between the allele count at the causal SNP and the most highly correlated SNP that is measured. As in the linkage scenario, it follows that if $n_c$ is the sample size necessary to obtain an association shift of size $c$ when the causal SNP is measured, then to obtain a shift of size $b$ with an unmeasured causal SNP, we require a sample size of

$$n_b = n_c \left( \frac{b}{c} \right)^2 \left( \frac{1}{r^2} \right).$$

These formulas may be used in reverse as well. Suppose sample sizes are predetermined. Then, for a plausible genetic model, one can compute the expected shift, $b$, in the formula above. See appendix B for detailed numerical examples of these calculations.

**Discussion**

When planning a whole-genome study to discover genes associated with a complex phenotype, a host of issues arise, many of which were discussed in a recent review by Thomas et al. (2005). To detect association via LD in a typical population requires a marker density on the order of 1 every 6 kb, which results in $\sim 500,000$ SNP markers. Many investigators have recommended a two-stage association analysis to circumvent the challenges of multiple testing (e.g., Sobell et al. [1993] and Sata-gopan and Elston [2003]). The concept of this study design is to perform an analysis on an initial sample of subjects for a dense grid of markers and then to follow up with an independent sample, tested for only the markers yielding “promising” results in the first stage. Although this approach is helpful, testing such a large number of hypotheses makes it difficult to detect meaningful associations. In the present study, we propose a novel approach for improving the power of a whole-genome association study that is applicable to one- or two-stage designs.

Investigators typically favor certain regions of the genome on the basis of the results of prior investigations, such as linkage analysis. We propose using this information in the form of weights for the $P$ values. Specifically, we divide $P$ values by weights and then apply the traditional FDR procedure to the weighted $P$ values. We investigate how weights might sensibly be chosen on the basis of a prior linkage study. Our investigations reveal that if the prior weights are informative, the procedure improves power considerably. Remarkably, even if the weights were uninformative, the loss in power is small.

Theory imposes few constraints on the weights beyond a basic conservation principle: the weights must average to 1. There remain many open questions about how to best incorporate information from multiple prior linkage studies. Presumably, the basic principles of meta-analyses can be applied to combine information into a single, so-called meta-linkage trace. Another obvious candidate for weights is “gene-based” and involves up-weighting regions—including coding, splice-site, and regulatory regions—and conserved intronic regions (see Thomas et al. 2005). Nevertheless, although many different weighting schemes are permissible in principle, searching to find the weights that give the desired significant results is not acceptable. For this reason, we give some suggested weighting schemes and presume that any other choice of weights would require scientific motivation before it would be considered valid.

From simulations, it is clear that a few dramatic weights are ideal if the prior data are highly informative. Otherwise, modest weights provide a more robust choice. For linkage-based weights, such as the exponential weights, it might be ideal to have a way of choosing the constant $B$ to maximize the power of the analysis. A data-driven option to achieve this goal is to select the $B$ that maximizes the number of discoveries for a particular genome scan; call it $D(B)$. Simulations reveal that $D(B)$ is a well-behaved step function. In most simulations, it starts low, steps up quickly to a maximum value, stabilizes for a broad range of choices, and then declines as $B$ gets larger. The exception is when the linkage trace is uninformative. In this case, the $B$ that maximizes $D(B)$ typically occurs for a value near zero. Although this empirical approach is appealing, it does have one notable drawback. Because the rate of false discoveries is controlled in expectation at level $\alpha$ for each $B$, maximizing the number of discoveries over $B$ leads to an excess of false discoveries. Our simulations suggest, however, that this approach results in a minor increase in FDR of $\sim 0.04$ over $\alpha = 0.05$. This increased error rate was often accompanied by a more-than-compensatory increase in power (results not shown). This promising idea requires further study. Already, it can provide a useful tool for selecting genes meriting further investigation.

In consideration of the use of prior information to improve testing, a Bayesian approach comes to mind. Indeed, the Bayesian FDR method given by Genovese and Wasserman (2003) can easily be extended to incorporate distinct priors for each hypothesis. Storey (2002) and
Efron et al. (2001) have given Bayesian interpretations of FDR. Here, we have proposed using P value weighting as a frequentist method for including prior information about the hypothesis, while leaving the false discovery rate unchanged. The relationship between weighted FDR and a fully Bayesian procedure for evaluating linkage and association is not obvious to us but could be an interesting area of inquiry.

Risch and Merikangas (1996) compared the power of case-control association designs to linkage studies that used ASPs. For realistic sample sizes, they showed that some genetic models were much more likely to produce substantial association statistics than they were to produce substantial linkage statistics. We revisited this topic and calculated the sample size required to obtain useful prior information from a linkage study for the space of genetic models likely to underlie complex phenotypes (table 2). This inquiry reveals that, among genetic models that are seemingly equal in genetic information, as measured in terms of roughly equal sample sizes required to detect an association signal, some are much more informative for linkage than others. This observation reinforces the fact that loci “detectable” by linkage designs are not always the same as those detectable by association, and vice versa. Nonetheless, by exploring the power of the weighted procedures for a wide range of scenarios, we show that weighted FDR can improve power for scenarios quite likely to be relevant for some complex diseases/phenotypes. A caveat is also worth noting: our simulations assume the loci important for generating liability in multiplex families and those generating liability in population-based samples overlap. It is theoretically possible that there is little overlap for these two sets of loci; in that case, using linkage information to weight association statistics will not improve power as substantially as our results suggest.

Even when a dense set of markers has been genotyped, the choice of tag SNPs is critical to ensure that signals are detectable indirectly via LD (Rinaldo et al. 2005). The best tag SNPs to pick depends somewhat on the nature of the statistical analysis planned (Roeder et al. 2005). In our simulations, we required an accurate approximation of the underlying time-series process. Consequently, we researched the appropriateness of an ARMA process to model the autocorrelation structure of linkage traces for 10,000 scans. For this purpose, we simulated and analyzed two data sets under the null hypothesis of no linkage between disease and markers. The two data sets comprised 400 ASPs. Each family has genotyped parents and missing genotypes at 7% of the loci overall. The data sets were simulated using SIMULATE (Terwilliger et al. 1993), and the linkage statistics were output every 1/3 cM, by use of Allegro (Gudbjartsson et al. 2000), with an exponential model option. We used the ARMA function in the R software (Venables and Ripley 2002) to fit two data sets to each of 16 models in the family of ARMA models, with AR and MA orders between 0 and 3. By using Akaike’s information criterion (AIC), the best fits were obtained using ARMA(2,1) and ARMA(3,1) models. Time-series diagnoses showed that these models provided a good fit to the autocorrelation

In closing, we note that technology has a tremendous effect on scientific inquiry, not all of it positive. In the workshop on genomewide association studies, Alice Whittemore asked “Is the technology driving the science?” (Thomas et al. 2005, p. 342). In the case of genomewide association studies, it is important to recall that just because we can afford to measure hundreds of thousands of SNPs does not mean we should let this additional data diminish our chances of detecting association signals in a targeted set of genes. The methods we propose allow scientific intuition to coexist more comfortably with emerging technology. Yes, we can measure more, but we can also use our scientific prior insights to target promising regions through the use of weights.

Acknowledgments

This research was supported by National Institutes of Health grant MH057881. Free software for performing weighted FDR analysis can be obtained at the Web site of the Computational Genetics Lab (University of Pittsburgh Medical Center).

Appendix A

A linkage scan with 10,000 SNPs takes considerable computer time to simulate; on a 1.4-GHz Opteron processor, simulation of a single linkage scan required 25 min. To estimate power, thousands of simulations are needed for each experimental condition investigated. Thus, direct simulation and analysis of 10,000 SNP linkage scans are infeasible, even on a cluster of processors.

In our simulations, we required an accurate approximation of the underlying time-series process. Consequently, we researched the appropriateness of an ARMA process to model the autocorrelation structure of linkage traces for 10,000 scans. For this purpose, we simulated and analyzed two data sets under the null hypothesis of no linkage between disease and markers. The two data sets comprised 400 ASPs. Each family has genotyped parents and missing genotypes at 7% of the loci overall. The data sets were simulated using SIMULATE (Terwilliger et al. 1993), and the linkage statistics were output every 1/3 cM, by use of Allegro (Gudbjartsson et al. 2000), with an exponential model option. We used the ARMA function in the R software (Venables and Ripley 2002) to fit two data sets to each of 16 models in the family of ARMA models, with AR and MA orders between 0 and 3. By using Akaike’s information criterion (AIC), the best fits were obtained using ARMA(2,1) and ARMA(3,1) models. Time-series diagnoses showed that these models provided a good fit to the autocorrelation
function for statistics from map locations separated by <50–60 cM. Whereas the fit was not as good for statistics separated by >60 cM, both the empirical and fitted autocorrelations were close to zero at that distance and thus had little practical impact.

Appendix B

After computing \( n_i (n_p) \) for a broad range of values of \( \kappa, \rho, \phi \), we fit a generalized linear model with the log link to predict \( n_i (n_p) \) from \( \kappa, \rho, \phi \), the square root of each of these terms, and all pairwise interactions. The best models among this class were chosen using the stepwise AIC option. The resulting models, which had \( R^2 = 99.9\% \) for linkage and 99.4\% for association, are presented in table B1. These are the models to obtain \( \mu_i = 2 \) \((\mu_s = 4.5)\) for a linkage (association) study.

For instance, if the genetic model is specified by \( \rho = 0.25, \kappa = 0.05 \), and \( \phi = 3 \), we can use table B1 to compute the number of sib pairs necessary to obtain an expected shift of 2 in a linkage study. Simply calculate the number of sib pairs necessary to obtain an \( r_p \) from the numbers reported in table 2 were obtained for a subset of genetic models with use of this formula, but with greater numerical precision in the coefficients. We can also compute the number of sib pairs necessary to obtain a shift of size \( b \neq 2 \) with use of the adjustment formula presented in the “Results” section. For example, if \( b = 3 \), we calculate \( n_b = n_u (b/\sqrt{k})^2 = 125(3/2)^2 = 281 \).

To determine the sample size necessary to obtain a shift of size \( \mu_s = 4.5 \) in an association study, we calculate \( n_r = \exp [x^T \gamma_r] \), where \( \gamma_r \) is the vector of association coefficients from table B1. For this example, if the causal SNP is measured, then ~62 cases and controls will be required to obtain the desired results. On the other hand, if the causal SNP had not been measured but has maximum correlation \( r = 0.8 \) with a measured SNP, then an additional 35 cases and controls would be required to obtain the same expected shift. Finally, if we have 200 cases and controls available, we expect a shift of 6.5 = 4.5 \( \times \sqrt{200/(62 + 35)} \). As indicated in table 2, these values are highly dependent on the assumed genetic model.

Web Resource

The URL for software presented herein is as follows:

University of Pittsburgh Medical Center, http://wpic.wpic.pitt.edu/wpicompgen

References

——— (2003b) False discovery or missed discovery? Heredity 91:537–538


