Probability of Detecting Disease-Associated SNPs in Case-Control Genome-Wide Association Studies

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Outline

• Genetics background
• Case-control Genome-Wide Association Study (GWAS)
• Ranking and selection procedures
• Performance criteria
  – Detection probability
  – Positive proportion
• Analytic results & simulations
• Extensions: two stage designs
• Conclusions
The Human Genome

- Four DNA bases (nucleotides): A (adenine), T (thymine), G (guanine), C (cytosine)
- 3 billion base pairs
- 22+2 chromosomes
- > 99% of human DNA sequences identical
- ~8,000,000 single nucleotide polymorphisms (SNPs)
  - alterations in single nucleotide e.g. AAGGC -> ATGGC

For variation to be considered a SNP, it must occur in >= 1% of population. SNPs occur every 100-300 bases, in coding (gene) and noncoding regions.

Find disease genes: study variation in SNPs
Linkage Disequilibrium

Genotyping all loci is not possible (not yet!)
=> Utilization of correlation of alleles at two loci

Marker locus

a: 10% ($p_a$)

Disease locus

G: 1% ($p_G$)

A: 90%

g: 99%

“D”
Linkage Disequilibrium

Bi-allelic disease locus: disease allele G ($p_G$), wild type allele g ($p_g$); Bi-allelic marker: a, A ($p_a$, $p_A$)

**Linkage disequilibrium (LD) defined as**

$$D = P(A, G) - p_A p_G$$

$$D' = \frac{D}{D_{\text{max}}} = \frac{D}{\min(p_G p_a, p_g p_a)} \text{ for } D>0$$

$$r^2 = (D')^2 \frac{p_a p_G}{p_A p_g}$$

$D'$ is upper bound of $r^2$
Genome wide approaches

• Try to get closer to disease locus by high density SNP coverage: tag SNPs

• **tag SNP:** representative SNP in region of genome with high LD ($r^2 > 0.8$) with untyped SNPs in that region (HapMap project)

• Typically: **550,000-650,000** tag SNPs/subject

• *Still estimated that 25% of SNPs not captured adequately by tag SNPs*
Case-control Genome Wide Association Studies

- Retrospective sample of unrelated cases (diseased) & controls (non-diseased)
- Exposure: tagSNPs covering genome
- Assess correlation between SNP genotypes and disease status
- Common SNPs (minor allele frequencies > 5%)
- Complex diseases (e.g. cancer)
- Small/moderate genetic effects (OR ~ 1.3)
Genetic Association

Disease outcome
Y=1 or 0

Tag SNP
Alleles
A, a

a: 10% (p_a)

Disease Locus

G: 1% (p_G)
## Observed data for cases and controls

<table>
<thead>
<tr>
<th>SNP i Genotype</th>
<th>Cases</th>
<th>Controls</th>
<th>Total counts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genotype</td>
<td>aa</td>
<td>aA</td>
<td>AA</td>
</tr>
<tr>
<td>cases</td>
<td>(r_0)</td>
<td>(r_1)</td>
<td>(r_2)</td>
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<tr>
<td>controls</td>
<td>(s_0)</td>
<td>(s_1)</td>
<td>(s_2)</td>
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<tr>
<td>total counts</td>
<td>(n_0)</td>
<td>(n_1)</td>
<td>(n_2)</td>
</tr>
</tbody>
</table>
Assumed Penetrance Model

\[ Y = \begin{cases} 
1, \text{ diseased individual} \\
0, \text{ healthy individual} 
\end{cases} \]

\[ P(Y=1 \mid X_i) = \frac{\exp[\mu + \beta X_i]}{1 + \exp[\mu + \beta X_i]} \]

\( X_i \) is a score attached to genotype of SNP \( i \)

additive model: \( X_i = \# \) of A alleles (0,1,2)
dominant model: \( X_i = 1 \) if \{aA, AA\}; =0 if \{aa\}
recessive model: \( X_i = 1 \) if \{AA\}; =0 if \{aa,aA\}
Tests for Association based on tag SNP

Test $H_0: \beta = 0$ using

Wald Test: $W_i = \hat{\beta}_i^2 / \hat{\text{Var}}(\hat{\beta}_i)$

Score Test: $S_i = U_i^2 / \hat{\text{Var}}_0(U_i)$

where $U_i = \phi \sum_{\text{cases}} X_i - (1 - \phi) \sum_{\text{controls}} X_i$
Remark on the null hypothesis

H₀: ”no association between SNP i and disease”

is true if either one of the following

1. Disease has no genetic component
2. D=0 (true disease locus not in LD with observed SNP)
Factors causing false positive associations

• Population Stratification
  – Confounding by ethnicity (bias)

• Cryptic relatedness
  – Variance distortion
  – Genomic control methods (Devlin & Roeder, 1999)

• Differential genotyping error (Clayton et al, Nature Genetics, 2005)
  – Failure to call genotype not independent of case-control status

• Multiple comparison: test 550,000 hypotheses
Controlling for Multiplicity

• Control experiment-wise Type I error
  – Bonferroni correction: $\alpha/N=10^{-7}$
  – Refinements (e.g. Simes Test)
  – Adaptations for two-phase designs (Skol et al., 2006)

• Control false discovery rate (FDR)
  (Benjamini and Hochberg, 1995)

• Ranking and Selection
Motivating Example: Cancer Genetic Markers of Susceptibility (CGEMS)


• A three-year, $14 million initiative

• **Outcome**: Prostate cancer (finished), breast cancer (ongoing)

• Combine data from 5 large studies

• **550K chip**: measures N=550,000 tag SNPs

Replication Strategy for Prostate Cancer

Whole Genome Scan
1200 cases / 1200 controls

Replication Study 1
2000 cases / 2000 controls

Replication Study 2
2000 cases / 2000 controls

Replication Study 3
2000 cases / 2000 controls

Replication Study 4
1200 cases / 1200 controls

>500,000 Tag SNPs

~20,000 SNPs

~1,500 SNPs

200+ New ht-SNPs*

25-50 Loci

*htSNP = haplotype-tagging SNPs
GWAS Designs

• Single stage design: all markers measured on all samples

• Two stage design:

  **Stage 1:** Proportion of available samples genotyped on large number of markers

  **Stage 2:** Proportion of these markers are followed up by genotyping them on remaining samples
Ranking and Selection for Single Stage Designs

• Sample $n$ unrelated cases ($Y=1$) and $n$ controls ($Y=0$)

• For each subject, measure $X_i=0,1,2$, the number of minor alleles at SNP $i$ for $i=1,\ldots,N$. 
Models for Genetic Effects in Source Population

$X_i = 0, 1, 2$, - number of minor alleles at ith SNP

SNPs 1,2, . . ., M disease-associated
SNPs M+1, . . . , N non-disease associated

Probability of disease in source population given by

$$P(Y = 1 | X_1, ..., X_N) = \frac{\exp(\mu + \sum_{i=1}^{M} \beta_i X_i)}{1 + \exp(\mu + \sum_{i=1}^{M} \beta_i X_i)}$$
Models for the genetic effects

• Fixed Effects Model:
  \[ \beta_i = \beta \text{ for } i=1,2, \ldots, M \]
  \[ \beta_i = 0 \text{ for } i=M+1, \ldots, N \]

• Random Effects Model:
  \[ \beta_i \sim N(0,\tau^2) \text{ for } i=1,2, \ldots, M \]
  \[ \beta_i = 0 \text{ for } i=M+1, \ldots, N \]
  \[ \mathbb{E}|\beta_i| = \tau (2/\pi)^{1/2} \approx 0.798\tau \]
**Chi-Square Statistics for Detecting Disease Association**

**Wald Test**: \( W_i = \hat{\beta}_i^2 / \hat{\text{Var}}(\hat{\beta}_i) \)

**Score Test**: \( S_i = U_i^2 / \hat{\text{Var}}_0(U_i) \)

where \( U_i = 0.5(\sum_{\text{cases}} X_i - \sum_{\text{controls}} X_i) \)

Use 2-sided test with additive scores, \( X = 0, 1, 2 \), both tests unchanged by assignment of “minor” allele.
Definitions for Ranking and Selection in One Stage Design

- **SNP i “T-selected”** if $W_i$ in top $T$ values, i.e. $\text{rank}(W_i) > N - T$

- **Detection probability (DP):** probability that a disease SNP will be T-selected

- **Proportion positive (PP):** proportion of true disease SNPs among T selected SNPs
Show: use of marginal model in cases and controls appropriate

Assumption: tagSNPs independent in source population, i.e. $X_1$ independent of $X_2, \ldots, X_N$

For rare disease,

$$P(Y = 1 \mid X) \approx \exp(\mu + \sum_{i=1}^{M} \beta_i X_i)$$

For a single disease SNP, say SNP 1,

$$P(Y = 1 \mid X_1) \approx \exp(\mu^* + \beta_1 X_1)$$

where $\mu^* = \mu + \log\{E \exp(\sum_{i=2}^{M} \beta_i X_i)\}$
Marginal model in cases and controls

In case-control population,

$$\logit\{P(Y = 1 \mid X_1)\} = \mu^{**} + \beta_1 X_1$$

where $\mu^{**} = \mu^* + \log(\pi_1 / \pi_0)$

$$\pi_i = P(\text{sampled} \mid Y = i), \ i = 0,1$$
If tagSNPs independent in source population, then tagSNPs independent in cases and controls

Let \( \rho_{ki} = P(X_i = k) \)

\( g_{ki} \equiv P(X_i = k \mid Y = 0) \approx \rho_{ki} \)

\( g_{klih} \equiv P(X_i = k, X_h = l \mid Y = 0) \approx \rho_{ki} \rho_{lh} \)

\( f_{ki} \equiv P(X_i = k \mid Y = 1) = \rho_{ki} \exp(\beta_i k) \left\{ \sum_{l=0}^{2} \rho_{li} \exp(\beta_i l) \right\}^{-1} \)

\( f_{klih} \equiv P(X_i = k, X_h = l \mid Y = 1) = \rho_{ki} \rho_{lh} \exp(\beta_i k + \beta_h l) \left\{ \sum_{s_1=0}^{2} \sum_{s_2=0}^{2} \rho_{s_1 i} \rho_{s_2 h} \exp(\beta_i s_1 + \beta_h s_2) \right\}^{-1} = f_{ki} f_{lh} \)
Properties of test statistics $W_i$

Based on independence of genotypes in cases and controls, we compute expected values of prospective estimating equations and their cross products using the retrospective sampling distributions and show that scores are uncorrelated.

Estimating each $\beta_i$ using separate logistic models yields independent estimates of $\beta_i$, thus $W_i$ are independent.

Similar results for score test $S_i$. 
Analytic Calculation of DP and PP

**Special Case:** disease SNPs have same allele frequency and $\beta$

Then $W_i \sim G$ for $i=1,...,M$: non-central chi-square with
non-centrality $\beta^2/\text{Var}(\beta)$ for fixed effects model

$W_i \sim F$, central chi-square (1df), $i=M=1,...,N$

$$DP = \int_0^{\infty} \left[ \sum_{m=0}^{\min(M-1,T-1)} \binom{M-1}{m} g(c)G(c)^{M-1-m} \{1-G(c)\}^m \sum_{s=0}^{T-m-1} \binom{N-M}{s} \{1-F(c)\}^s \{F(c)\}^{N-M-s} \right] dc$$

$$PP = \frac{M}{T}(DP)$$

Generalizes to different $G_i$ for various disease SNPs with differing allele frequencies and $\beta_i$
Approximation for M=1 Disease SNP

\[ DP \approx 1 - G\{F^{-1}(1 - T/N)\} \]

In this case, DP equivalent to power of a test with same non-centrality but with type I error (alpha-level) T/N
Simulations

Brute force simulation not feasible; use analytic results including independence

• For each simulation (ISIM) NSIM=10,000
• For SNPs i=1,2, M obtain fixed or random $\beta_i$
• For SNPs i=M+1, . . . N let $\beta_i=0$
• For each SNP:
  • Draw random minor allele frequency (MAF) from distribution of MAFs in CGEMS controls (MAF≥0.05): mean=0.276, median=0.26, IQR: 0.15-0.38
  • Solve for $\mu^{**}$ such that $P(Y=1)=0.5$
  • Compute information matrix from case-control logistic model, using retrospective sampling distributions
  • Compute $\text{Var}(\hat{\beta}_i)$
  • Sample $\hat{\beta}_i$ from $N(\beta_i, \text{Var}(\hat{\beta}_i))$
  • Compute $W_i=\frac{\hat{\beta}_i^2}{\text{Var}(\hat{\beta}_i)}$
Simulated Estimates of DP and PP

Define \( I(m, ISIM, T) = 1 \) if \( \text{rank}(W_m) > N-T \), 0 otherwise.

\[
\hat{DP} = NSIM^{-1}M^{-1} \sum_{m=1}^{M} \sum_{ISIM=1}^{NSIM} I(m, ISIM, T)
\]

• Probability a given disease SNP is T-selected
• Proportion of disease SNPs selected

\[
\hat{PP} = (M)(\hat{DP})/T
\]
Results for Fixed Effects Model
DP for Fixed Effects Model. n=1000 black; n=8000 red. Odds ratios 1.1, 1.2, 1.3, 1.5, 2.0. T on log scale.
PP on log scale versus T on log scale. M=1 black; M=100 red. Odds ratios 1.2, 1.3, 1.5, 2.0.
Results for Random Effects Model
DP for Random Effects Model. $n=1000$ black; $n=8000$ red. $0.798\tau=\log$ of 1.1, 1.2, 1.3, 1.5, 2.0. $T$ on log scale.
Summary: single stage design

DP and PP are useful criteria for ranking
Related work: Zaykin & Zhivotovsky, 2005; Satagopan et al, 2004

For OR=1.2, large T required to assure adequate DP. T=25,000 yields DP=0.69 (fixed effects model)

DP larger for fixed effects than random effects model

DP is decreased by increasing M for M>T

PP decreases with T for T>M. Increasing T to increase DP decreases PP and is futile if n is too small.
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>500,000 Tag SNPs

~20,000 SNPs

~1,500 SNPs

200+ New ht-SNPs*

25-50 Loci

*htSNP= haplotype-tagging SNPs
Extension: two stage designs

Stage 1: \( \pi_{\text{samples}} \) genotyped on 550,000 SNPs
Stage 2: \( T_1 \) SNPs largest \( \chi^2 \)-square statistics followed up by genotyping on remaining samples.

Replication analysis: view stage 2 as a replication study; final selection of the SNPs depended on ranking only the \( \chi^2 \)-square statistics in stage 2.

Joint analysis: for each of the \( T_1 \) SNPs selected in stage 1, compute

\[
\lambda C_1 + (1-\lambda)C_2, \quad \lambda = 0.0, 0.05, 0.1, \ldots, 1.0
\]

\( C_i \) chi-square statistic observed in stage i
For each value of \( \lambda \) estimate DP, present maximal DP with corresponding \( \lambda \).

Skol et al, Nature genetics, 2006
Two stage designs, cont.

Compute probability that exactly $M_2$ disease SNPs selected after stage 1, given that $T_1$ SNPs were selected and there were $K_1$ non-disease SNPs and $M_1$ disease SNPs.

Use joint densities of the $(M_1-M_2+1)th$ and $(M_1-M_2)th$ order statistic of $M_1$ disease SNPs, and $(K_1-T_1-M_2+1)th$ and $(K_1-T_1-M_2)th$ order statistic of the $K_1$ non-disease SNPs.

$W_1^{(i)}$ - order statistics of disease SNPs

$W_0^{(i)}$ - order statistics of non-disease SNPs
Fixed effects model, same allele frequencies for all disease SNPs

\[
P(\text{exactly } M_2 \text{ disease SNPs chosen after stage 1}) = \\
P(W_1^{(M_1-M_2+1)} > W_0^{(M_1-M_2+1)}; W_0^{(K_1-T_1+M_2+1)} > W_1^{(M_1-M_2)}) = \\
\int_{-\infty}^{\infty} \int_{-\infty}^{w_1} \int_{\min(x_1,w_1)}^{\infty} \frac{K_1!}{(K_1-T_1+M_2-1)!(T_1-M_2-1)!} \frac{M_1!}{(M_1-M_2-1)!(M_2-1)!} \\
\{1-G(w_1)\}^{M_2-1} g(w_1) g(w_2) \{1-F(x_1)\}^{T_1-M_2-1} F(x_2)^{K_1-T_1+M_2-1} f(x_1) f(x_2) dx_2 dx_1 dw_2 dw_1
\]
Special cases

Special case: $M_2=0$

\[
P(\text{zero disease SNPs chosen after stage1}) = P(W_0^{(K_1-T_1+1)} > W_1^{(K_1-T_1+1)}) = \int_0^\infty \int_0^y \frac{K_1!}{(K_1-T_1)!(T_1-1)!} \{1 - F(y)\}^{T_1-1} F(y)^{K_1-T_1-1} f(y) M_1 G(x)^{M_1-1} g(x) \, dx \, dy
\]

Special cases: $M_2=M_1$

\[
P(\text{all disease SNPs chosen after stage1}) = P(W_1^{(1)} > W_0^{(K_1-T_1+M_1)}) = \int_0^\infty \int_y^\infty \frac{K_1!}{(K_1-T_1+M_1-1)!(T_1-M_1)!} \{1 - F(y)\}^{K_1-T_1+M_1-1} F(y)^{T_1-M_1} f(y) M_1 (1-G(x))^{M_1-1} g(x) \, dx \, dy
\]

Special case $M_2=T_1$

\[
P(\text{all disease SNPs chosen after stage1}) = P(W_1^{(M_1-M_2+1)} > W_0^{(K_1)}) = \int_0^\infty \int_y^\infty K_1 F(y)^{K_1-1} f(y) \frac{M_1!}{(M_1-M_2)!(M_2-1)!} G(x)^{M_1-M_2} [1-G(x)]^{M_2-1} g(x) \, dx \, dy
\]
Probability of detecting a disease SNP (DP) and optimal stage1 weight for fixed effects model with log odds ratio per allele $\beta = \log(1.2)$ for 8000 cases and 8000 controls

<table>
<thead>
<tr>
<th>$\pi_{\text{sample}}$</th>
<th>Analysis</th>
<th>Number of disease SNPs, $M_0 = 1$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T_1 = 1000$</td>
<td>$T_1 = 25,000$</td>
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<tr>
<td></td>
<td>$T_2 = 1$</td>
<td>$T_2 = 100$</td>
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<tr>
<td>0.125</td>
<td><strong>Replicate</strong></td>
<td>.266</td>
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<tr>
<td></td>
<td><strong>Joint</strong></td>
<td>.267</td>
</tr>
<tr>
<td></td>
<td>$\lambda_{\text{opt}}$</td>
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<tr>
<td>0.25</td>
<td><strong>Replicate</strong></td>
<td>.612</td>
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<tr>
<td></td>
<td><strong>Joint</strong></td>
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<tr>
<td>1.00</td>
<td><strong>One-stage$^a$</strong></td>
<td>.882</td>
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</tbody>
</table>
Summary, two stage design

• A small first stage can only partially be compensated for by selecting a large number of SNPs for further testing
• Even joint analysis does not improve DP much if $\pi_{\text{sample}} \leq 0.25$
• As the cost per genotype for stage1 drops, compared to later stages, economic incentives for multi-stage designs decrease
• Advantage of one-stage design: offers unbiased estimates of genetic effects that can be combined easily with those from other studies; multi-stage designs introduce selection biases that complicate meta-analyses.
References


Reich and Goldstein, 2001. Genetic Epidemiology, 20: 4-16
Platforms & SNP Chips

- Affymetrix 100K
- Affymetrix 500K
  
- Illumina 317K
- Illumina 550K
  
- Illumina 650Y (550K+100K YRI fill in)

essentially random set of SNPs
designed using Hapmap