Erasmus MC hurs

Genome-Wide Association analysis significance, power and coverage

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Outline

- Significance of GWA study
- Power of GWA study
- Coverage of GWA study
- Concluding remarks

Bonferroni for GW significance?

Bonferroni correction

- GW type 1 error rate of 0.05 corresponds to nominal P = 0.05/(# SNPs)
- Problems:
 - Bonferroni assume that tests are independent
 - <u>SNPs are not</u> (because of LD)
 - Therefore Bonferroni is conservative correction (meaning you loose power and can miss association when it is truly there)
 - 550K SNPs were typed, and imputations were done to 2.5M SNPs using HapMap panel. How many tests are done? 0.5M or 2.5M? ... or neither?

Empirical GW significance?

- Empirical estimation of GW (experiment-wise) significance gives exact answer, taking the LD structure and phenotype distribution into account
- Works very well for a single one-stage study
- Problems:
 - May be technically demanding (no problem for few dozens of traits, but is a problem for 100s)
 - More complex design: e.g. two-stage, or multiple independent studies
 - Knowledge accumulation (meta-analysis)



Multiple testing burden: fixed threshold

- Pe'er et al, Genetic Epi, 2008, 32: 381-385
- If we measured all common SNPs in the genome, what number of "independent" SNPs could mimic the null distribution of the test statistics?
 - ~1M tests \rightarrow GW 5% ~ nominal P = 0.05/1M = 5 10⁻⁸
 - To keep in mind:
 - Above is true for CEU (2M for Yoruba)
 - Estimated using 1/600th of the genome (ENCODE)

So is my *p*-value significant or not?!

- You (your referees) may be convinced (or not) by a *p*-value which pass
 - Permutation procedure
 - Bonferroni correction
 - $P < 5 \times 10^{-8}$

- Ultimate answer: replication
- This is a way to
 - Achieve "overwhelming significance"
 - Exclude possibility that the finding is "study-specific"

Example

- A genetic study estimates effect of the SNP rs724016*C allele on height as +4.6 mm (s.e. = 0.88)
 - Nominal *p*-value = 2×10^{-7}
 - Permutation-based *p*-value = 0.045
 - Bonferroni p-value = 0.06
 - Fixed threshold: $2 \times 10^{-7} > 5 \times 10^{-8}$
- Is that a true finding or not?
- Replicate!



Replication in three populations

Study Reference

Study	Effect	S.E.	P-value
Original	4.6	0.88	2 x 10 ⁻⁷
Rep 1	3.5	2.21	0.11
Rep 2	3.6	1.59	0.02
Rep 3	2.8	1.15	0.001
Total	4.14	0.62	2 x 10 ⁻¹¹





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Estimating power

Is study large enough to achieve statistical significance?

- Proportion of trait variance (V_{SNP}) explained by the SNP (this is the coefficient of determination, Rho2!)
- The non-centrality parameter (NCP)
 - Measures (under alternative) how much the $(\chi\eta\iota \sigma\theta\upsilon\alpha\rho\epsilon)$ test statistic is expected to deviate from it's expectation under the null
 - NCP = (no. samples) x V_{SNP}
- Power to achieve critical threshold X is $Pr(T^2_{NCP} > X)$ Can be computed in R using pchisq(X,df=1,ncp=*NCP*,low=FALSE)

Exact (not known!) model of the gene action is to be assumed -need to pick up some reasonable model



Power as function of NCP



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Power of GWA study

	Sample size	V_{SNP}	NCP	Power to achieve p < 5 x 10 ⁻⁸
"Biggest common loci":		3%	30	51%
	1,000	1%	10	1%
		0.5%	5	<1%
• HDL: <i>CETP</i> ~ 2.5%		0.1%	1	<1%
		3%	150	100%
• Total chol.: $APOE \sim 0.5\%$	5,000	1%	50	95%
		0.5%	25	33%
• Height $HMGA2 \sim 0.3\%$		0.1%	5 <1%	
1101 <u>5</u> 111. 11110712 0.570		3%	300	100%
	10,000	1%	100	100%
		0.5%	50	95%
		0.1%	10	1%



A note on adjustment for the covariates

- Consider *HMGA2* which explains 0.15% of height variation
- Expected power in a study of 14000 people is 20%
- Sex and age together explain ~50% of height variation
- Therefore in the adjusted data the QTL explains 0.3%
- The power to detect it GW is thus 84%



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How many SNPs we capture?



- Red: typed SNPs
- Green: SNPs with R2 ≥ 0.8 with a typed SNP (well-captured)
- Blue: SNPs with 0.8>R2≥0.5 with a typed SNP (captured)
- Black: SNPs with R2<0.5



Max R2 with a typed SNP depends on MAF



- Selected SNPs are likely to be common (if it is very rare, it is not likely to be known!)
- High R2 between two SNPs is possible only if their frequencies are similar



Genomic coverage by standard panels

What proportion of common SNPs (MAF \geq 0.05) are in the genotyped set or are in high LD (r²>0.8) with at least one genotyped SNP?

SNP panel	Туре	CEU	JPT+CHB	YRI
Affymetrix 111K	Random	31	31	15
Affymetrix 500K	Random	65	66	41
Affymetrix 1M	Combined	80		
Illumina 300K	Tag	75	63	28
Illumina 550K	Tag	87		
Illumina 1M	Tag	91		Not0 a mot

Barret & Cardon, NatGenet, 2006 Anderson et al., AJHG, 2008

HapMap population



Coverage pitfalls

- With 1,000K-2,000K SNP panels we may expect good coverage of common variants for any human population
- Some diseases/traits may be expected to be explained in large part by common variants
- For other disease multiple rare variants may play large role
- Coverage is poor for rare variants



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Painting everything black?!

- Study of height in a cohort of 10,000 people of European origin, using Illumina 300K
 - 25% of common variants are **not** captured
 - Vast majority of rare variants is **not** captured
 - Power to detect even the biggest effect using the directly typed SNP (rs724016 at HMGA2, explains 0.3%) is only 58%!

Power to detect height loci in a study of 10,000 people

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What is the chance to miss ALL loci?

- P(miss all) = P(miss locus 1) x P(miss locus 2) x ... P(miss locus N-1) x P(miss locus N)
- Chance to miss all 20 loci from the height paper of Weedon: only 8%
- Thus you will find at least one with power of 92%
- There are much more than 20 loci involved in height
- Your chances (to find >1 loci) are very good!