

# Gene-gene (and gene-environment) interactions

Heather J. Cordell

Institute of Genetic Medicine  
Newcastle University, UK



## (Pairwise) interaction

- Statistical interaction most easily described in terms a of (logistic) regression framework
  - Suppose  $x_1$  and  $x_2$  are binary factors whose presence/absence (coded 1/0) may be associated with a disease outcome
  - Logistic regression models their effect on the log odds of disease as:

$$\log \frac{p}{1-p} = \beta_0 + \beta_1 x_1$$

Marginal effect of factor 1

$$\log \frac{p}{1-p} = \beta_0 + \beta_2 x_2$$

Marginal effect of factor 2

$$\log \frac{p}{1-p} = \beta_0 + \beta_1 x_1 + \beta_2 x_2$$

Main effects of factors 1 and 2

$$\log \frac{p}{1-p} = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2$$

Main effects and interaction term

- For quantitative traits, use linear regression (replace  $\log \frac{p}{1-p}$  with  $y$ )

# Gene-gene interaction

- Expected trait values (log odds of disease) take the form:

Factor 1	Factor 2	
	1	0
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 $\Rightarrow$  Implies that the overall effect of two variables is greater (or less) than the 'sum of the parts'
- Suppose no main effects ( $\beta_1 = \beta_2 = 0$ )
  - Then we have

Factor 1	Factor 2	
	1	0
1	$\beta_0 + \beta_{12}$	$\beta_0$
0	$\beta_0$	$\beta_0$

- Trait value only differs from baseline if both factors present

# Gene-gene interaction

- However genetic predictors e.g. SNPs are not binary, but rather take 3 levels according to the number of copies (0,1,2) of the susceptibility allele possessed
- Most general ‘saturated’ (9 parameter) genotype model allows all 9 penetrances to take different values
  - Via modelling log odds in terms of:
    - A baseline effect ( $\beta_0$ )
    - Main effects of locus  $G$  ( $\beta_{G_1}, \beta_{G_2}$ )
    - Main effects of locus  $H$  ( $\beta_{H_1}, \beta_{H_2}$ )
    - 4 interaction terms

Locus G	Locus H		
	2	1	0
2	$\beta_0 + \beta_{G_2} + \beta_{H_2} + \beta_{22}$	$\beta_0 + \beta_{G_2} + \beta_{H_1} + \beta_{21}$	$\beta_0 + \beta_{G_2}$
1	$\beta_0 + \beta_{G_1} + \beta_{H_2} + \beta_{12}$	$\beta_0 + \beta_{G_1} + \beta_{H_1} + \beta_{11}$	$\beta_0 + \beta_{G_1}$
0	$\beta_0 + \beta_{H_2}$	$\beta_0 + \beta_{H_1}$	$\beta_0$

- Corresponds in statistical analysis packages to coding  $x_1, x_2$  (0,1,2) as a “factor”



# Gene-gene interaction

- Alternatively we can assume additive effects of each allele at each locus:
  - Corresponds to fitting

$$\log \frac{p}{1-p} = \beta_0 + \beta_G x_1 + \beta_H x_2 + \beta_{GH} x_1 x_2$$

with  $x_1, x_2$  coded (0,1,2)

Locus G	Locus H		
	2	1	0
2	$\beta_0 + 2\beta_G + 2\beta_H + 4\beta_{GH}$	$\beta_0 + 2\beta_G + \beta_H + 2\beta_{GH}$	$\beta_0 + 2\beta_G$
1	$\beta_0 + \beta_G + 2\beta_H + 2\beta_{GH}$	$\beta_0 + \beta_G + \beta_H + \beta_{GH}$	$\beta_0 + \beta_G$
0	$\beta_0 + 2\beta_H$	$\beta_0 + \beta_H$	$\beta_0$

- Much discussion in the literature

- Siemiatycki and Thomas (1981) *Int J Epidemiol* 10:383-387
- Thompson (1991) *J Clin Epidemiol* 44:221-232
- Phillips (1998) *Genetics* 149:1167-1171
- Cordell (2002) *Hum Molec Genet* 11:2463-2468
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- Bottom line is, little direct correspondence between statistical interaction and biological interaction
  - In terms of whether, for example, gene products physically interact
- However, existence of statistical interaction does imply both loci are “involved” in disease in some way
  - Provides a good starting point for further investigation of their (joint) involvement
    - Can be informed by the estimated penetrance values
    - Best addressed through other types of experimental data

## Some references

- For more details on gene-gene ( $G \times G$ ) interactions (epistasis) see
  - Cordell HJ (2009) Nat Rev Genet 10(6): 392-404
  - Wei, Hemani and Haley (2014) Nat Rev Genet 15(11):722-33
- For more details on gene-environment ( $G \times E$ ) interactions see
  - Thomas D (2010) Nat Rev Genet 11(4): 259-272
- Conceptually many similar issues in terms of definition and mathematical modelling
  - However, many practical issues rather different

# $G \times G$ versus $G \times E$

- For  $G \times E$ , we generally have to decide which environment(s) to measure/test
  - For  $G \times G$ , assuming we have GWAS data, we have already measured the genetic factors of interest
- Measurement error and confounding are a bigger issue for environmental factors?
  - e.g. diet, smoking, pollution levels
  - Issues of recall bias?
  - Current genotyping platforms (though not necessarily sequencing platforms) have relatively low error rates, less prone to biases

# $G \times G$ versus $G \times E$

- Typically GWAS measure thousands if not millions of genetic variants
  - But only a few (tens or at most 100s) of environmental factors
- Feasible to consider all  $G \times E$  combinations
- All pairwise  $G \times G$  combinations possible, but much more time consuming
  - And leads to greater multiplicity of tests
  - Also, why stop at 2-way interactions?
    - Could look at all 3 way, 4 way etc. combinations
    - Scale of problem quickly gets out of hand
    - Less obvious reason to do this for  $G \times E$ ...



# G×G versus G×E

- Risk estimation more important for G×E (?)
  - Estimating genetic risks in particular environments
  - Estimating effect of environmental factor on particular genetic background
    - Important for treatment/screening strategies and public health interventions
- For G×G, focus of interest is more related to
  - Increasing power to detect an effect (by taking into account the effects of other genetic loci)
  - Modelling the biology, especially related to the joint action of the loci

# Testing association and/or interaction

- Go back to binary coding of genetic (and/or environmental) factors

$$\log \frac{p}{1-p} = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2$$

- 3df test of  $\beta_1 = \beta_2 = \beta_{12} = 0$  tests for association **at both loci** (or both variables), allowing for their possible interaction

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- Depending on circumstances, any of these tests may be a sensible option
- Most tests of interaction/joint action can be thought of as a version of one or other of these tests
  - Although different tests vary in their precise details
  - And their relationship to the logistic regression formulation not always clearly described

# Testing for interaction

- Case/control studies

- Measure risk factors (e.g. SNP genotypes)  $x_1$  and  $x_2$  in sample of affected individuals (cases) and unaffected individuals (controls)
- At each SNP each person has one of 3 possible genotypes

$$1|1, \quad 1|2 = 2|1, \quad 2|2$$

- Can code as 3 different levels, count alleles or make dominance/recessive assumptions  $\Rightarrow$  binary factor
- Analyse using logistic regression (e.g. in R, SAS, PLINK)
  - If use binary coding, end up with 1 interaction term
  - If genotypes coded as 3 levels, get 4 interaction terms.

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    - If genotypes coded as 3 levels, get 4 interaction terms.
- For analysis of quantitative traits in unrelated individuals, use linear rather than logistic regression
- Family studies: use extension of case/pseudo-control approach (Cordell et al. (2004) Genet Epidemiol 26:186-205) or else use linear mixed models



# Case-only analysis

- Piergorsh et al. 1994; Yang et al. 1999; Weinberg and Umbach 2000
- Several authors have shown that, for binary predictor variables, a test of the interaction term  $\beta_{12}$  in the logistic regression model

$$\log \frac{p}{1-p} = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2$$

can be obtained by testing for correlation (association) between the genotypes at two separate loci, within the sample of cases

- Gains power from making assumption that genotypes (alleles) at the two loci are uncorrelated in the population
  - So only really suitable for unlinked or loosely linked loci (since closely linked loci are likely to be in LD)
- Alternatively **contrast** the genotype correlations in cases with those in controls (`--fast-epistasis` in PLINK)

# PLINK's *fast-epistasis* statistics

- PLINK takes unphased genotype data

Locus G	Locus H		
	2	1	0
2	<i>a</i>	<i>b</i>	<i>c</i>
1	<i>d</i>	<i>e</i>	<i>f</i>
0	<i>g</i>	<i>h</i>	<i>i</i>

and expands it to 2×2 allelic table

Locus G	Locus H	
	<i>H</i> <sub>1</sub>	<i>H</i> <sub>2</sub>
<i>G</i> <sub>1</sub>	<i>A</i> = 4 <i>a</i> + 2 <i>b</i> + 2 <i>d</i> + <i>e</i>	<i>B</i> = 4 <i>c</i> + 2 <i>b</i> + 2 <i>f</i> + <i>e</i>
<i>G</i> <sub>2</sub>	<i>C</i> = 4 <i>g</i> + 2 <i>h</i> + 2 <i>d</i> + <i>e</i>	<i>D</i> = 4 <i>i</i> + 2 <i>h</i> + 2 <i>f</i> + <i>e</i>

- PLINK estimates the log OR ( $\lambda$ ) for association/correlation between the loci as as  $\log(AD/BC)$  with estimated variance ( $v$ ):

$$\frac{1}{A} + \frac{1}{B} + \frac{1}{C} + \frac{1}{D}$$

- A z-test of whether correlation **exists** (case-only) or is **different** in cases and controls (case/control) is:

- Case-only:  $T_{FE-co} = \frac{\lambda_A^2}{\hat{v}_A}$
- Case/control:  $T_{FE-cc} = \frac{[\lambda_A - \lambda_N]^2}{\hat{v}_A + \hat{v}_N}$



# Testing correlation between loci

- A similar idea is implemented in EPIBLASTER (Kam-Thong et al. 2011 EJHG 19:465-571)
- Wu et al. (2010) (PLoS Genet 6:e1001131) also proposed a similar approach
- All these methods test whether correlation exists (case-only) or is different in cases and controls (case/control) via testing a log OR for association between two loci
  - However, the log OR for association ( $\lambda$ ) encapsulates a slightly different quantity between the different methods

# Testing correlation between loci

- Unfortunately, both Wu et al. (2010) and PLINK calculate the variance of their log ORs incorrectly
  - Resulting in a severe inflation in type 1 error (false positive) rate for the Wu et al. method
  - PLINK's statistic remains approximately correct (and completely correct in PLINK 1.9)
- We demonstrated this problem, and used the results of Brown (1975) to calculate the correct variances
  - Resulting in adjusted versions of all four statistics
- We also proposed some alternative Joint Effects (JE) statistics that have some advantages over these previous methods
  - See Ueki and Cordell (2012) PLoS Genetics 8(4):e1002625
  - Implemented in CASSI
    - <http://www.staff.ncl.ac.uk/richard.howey/cassi/>

# Screening for interactions

- So far we have considered how to test for interaction between two specific factors
- In GWAS we typically test for (marginal) association at between 500,000 and 1 million SNPs across the genome
- Simplest way to search for interactions is to perform an **exhaustive search**, considering all pairwise combinations
  - If testing  $G \times E$  with 5 environmental variables (for example), we end up with  $5 \times 1 \text{ million} = 5 \times 10^6$  tests
  - If testing  $G \times G$ , we end up with [1 million choose 2]  $\approx 5 \times 10^{11}$  tests
    - Computationally possible, but time-consuming
    - And dramatically increases multiple testing burden
    - But may be outweighed by increased power  
(Marchini et al. (2005) Nat Genet 37:413-417)
- Also need to decide exactly which test to perform
  - Several 'methods' (programs) choose their test on the basis of convenience/speed, given their chosen search strategy

# Exhaustive testing

- Several recent publications have focussed on trying to speed up exhaustive search procedure
- E.g. by making use of data compression techniques and parallelization
  - Steffens et al. (2010) Hum Hered 69:268-284
- Or by using Graphical Processing Units (GPUs)
  - Sinnott-Armstrong et al. (2009) BMC Res Notes 2:149
  - Greene et al. (2010) Bioinformatics 26:694-695
  - Hu et al. (2010) Cell Res 20:854-857
  - Hemani et al. (2011) Bioinformatics 27:1462-1465
  - Kam-Thong et al. (2012) Hum Hered 73:220-236

# Exhaustive testing (cont.)

- Or by computing faster tests e.g. log linear models rather than logistic regression
  - INTERSNP (Herold et al. (2009) Bioinformatics 25:3275-3281)
  - BOOST (Wan et al. (2010) AJHG 87:325-340)
  - BiForce Toolbox (Gyenesei et al. (2012) PMID:22689639)
- Or by performing an 'approximately' complete search
  - SIXPAC (Prabhu and Pe'er (2012) Genome Res 22:2230-2240)


# Problems with exhaustive testing

- Problem of interpretation/noise
  - Will the effect be strong enough to withstand the multiple testing problem/lower prior probability that any effect is real?
- Why stop at pairwise combinations (why not 3-way, 4-way etc.)?
  - Most methods do not scale up to all 3-way, 4-way etc. combinations
  - Even if they did, problem of interpretation/multiple testing would be even worse
- May need to use 'filtering' approach where only consider a subset of loci chosen based on loose single-locus significance or other (biological or statistical) considerations






# Biological filtering

- Emily et al. (2009) reported four significant cases of epistasis between unlinked loci in the WTCCC (Crohn's, Bipolar, Hypertension and Rheumatoid Arthritis) data
  - When limiting search on the basis of experimental knowledge of biological networks
- Herold et al. (2009) used their INTERSNP program to identify two SNPs that predispose to male pattern baldness, lying in genes from a joint pathway 
- Results require replication (as has become gold standard in GWAS)
  - Problematic for interactions, owing to larger sample size required to give sufficient power to detect interactions (in comparison to main-effects)
    - Gauderman (2002) Am J Epid 155:478-484
    - Zuk et al. (2012) PNAS 109:1193-1198

# Biological filtering

- Test at the gene level rather than the SNP level?
- E.g. across genes ( $G \times G$ )
  - Wang et al. (2009) *Genet Epidemiol* 33:6-15
  - He et al. (2011) *EJHG* 19:164-172
  - Li and Cui (2012) *Annals of Applied Statistics* 6:1134-2261
  - Rajapakse et al. (2012) *Genet Epidemiol* 36:622-630
  - Ma et al. (2013) *PLoS Genet* 9(2): e1003321
    - Compared 4 different tests that combine  $P$  values from pairwise (SNP  $\times$  SNP) interaction tests: min  $P$ , extended Simes, truncated tail, truncated product
    - Showed that the truncated tests did best
    - Presented an application only considering gene pairs known to exhibit protein-protein interactions
- Or within genes 
  - Dinu et al. (2012) *PLOS ONE* 7:e43035
  - Wei et al. (2013) *PLOS ONE* 8:e71203

# Statistical filtering

- Only test for interactions between 'significant' loci from a single-locus scan
  - Strange et al. 2010 (Nat Genet 42:985-990) found interactions between *HLA-C* and *ERAP1* in psoriasis
  - Evans et al. 2011 (Nat Genet 43:761-767) found interactions between *HLA-B27* and *ERAP1* in ankylosing spondylitis
  - Castillejo-Lopez et al. (2012) (Ann Rheum Dis 71:136-142) found interactions between polymorphisms in *BANK1* and *BLK* in SLE
- Only test for interactions between top 20% (or similar) loci from a single-locus scan
  - Nothing found in WTCCC Crohn's data (Cordell 2009)
- Test for interactions between 'significant' loci and all other loci


- Two-stage procedures
  - Test all pairwise combinations at screening stage
  - Follow up with independent test of all pairs passing some threshold
  - Reduces multiple testing problem at second stage by constructing tests that are independent of 1st stage
    - Murcray et al. (2009) Am J Epidemiol 169:219-226
    - Lewinger et al. (2013) Genet Epid 37:440-451
    - Jiao et al. (2013) Genet Epid 37:452-464
    - Fråanberg et al. (2015) PLOS Genetics 11(9):e1005502  
“Discovering genetic interactions in large-scale association studies by stage-wise likelihood ratio tests”

# Data mining approaches

- Clever computational algorithms for searching (fast) through plausible space of models
  - Including models that involve multi-way (not just pairwise) interactions
- Most methods cross-validation to avoid over-fitting
  - E.g. fit a model using 9/10 of data, use remaining 1/10 to assess performance, and repeat many times
  - Choose final model (set of predictors) that performs well
- Often use permutation approaches to assess final significance
  - And final model often re-fit by logistic regression to provide parameter estimates
- **Replication** in an independent data set is crucial

- Multifactor Dimensionality Reduction (Ritchie et al. (2001) AJHG 69:138-147)
  - Divide data into 10 equal parts
  - Fit model using  $\frac{9}{10}$  of data, use remaining  $\frac{1}{10}$  to assess performance, repeat for each  $\frac{9}{10}/\frac{1}{10}$  partition
  - Pick best-fitting model from all considered partitions
- Within each partition, perform exhaustive search over all single-locus models, 2-locus models, 3-locus models...
  - Computationally prohibitive for large numbers of loci
  - Use in conjunction with initial filtering method e.g. TuRF
- Has been used to detect potential interacting loci in breast cancer, type 2 diabetes, rheumatoid arthritis and coronary artery disease
  - Require replication in an independent data set

- Model construction based on classifying genotype combinations at the  $n$  loci as 'high' or 'low' risk
  - Based on the number of cases and controls in each cell
- Equivalent to fitting saturated genotype model



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	2	1	0
2	$\beta_0 + \beta_{G_2} + \beta_{H_2} + \beta_{22}$	$\beta_0 + \beta_{G_2} + \beta_{H_1} + \beta_{21}$	$\beta_0 + \beta_{G_2}$
1	$\beta_0 + \beta_{G_1} + \beta_{H_2} + \beta_{12}$	$\beta_0 + \beta_{G_1} + \beta_{H_1} + \beta_{11}$	$\beta_0 + \beta_{G_1}$
0	$\beta_0 + \beta_{H_2}$	$\beta_0 + \beta_{H_1}$	$\beta_0$

- But then reduce resulting  $3^n$  dimensional model to 2-dimensional model (high/low risk)

# Other model-search based approaches

- Random forests
  - Based on classification and regression trees (CART)
- Penalized regression methods
  - E.g. Zhu et al. (2014) Genet Epid 38:353-368
- Entropy-based methods
  - e.g. MECPM (maximum entropy conditional probability modelling)
    - Miller et al. (2009) Bioinformatics 25:2478-2485
    - Performed extremely well in comparison to other approaches in comprehensive simulation study by Chen et al. (2011) BMC Genomics 12:344
- Bayesian model selection
  - Involves specifying prior distributions for the number of loci and their effect sizes (=regression coefficients) including interactions
  - Use MCMC techniques to search through space of possible models, find model that maximizes likelihood
    - See also a recent Bayesian network approach (LEAP) which uses a heuristic search algorithm; outperformed MECPM in the study by Jiang and Neapolitan (2015) Genet Epid 39:173-184.



- Zhang and Liu (2007) Nat Genet 39:1167-1173
- Loci divided into 3 groups:
  - Not associated with disease
  - Contribute via main effects only
  - Contribute via saturated interaction model
- Use MCMC to jump through space of possible models (divisions of loci)
- Generates posterior probabilities for each SNP of being in each group
  - Or test via B-statistic
- Method has been recently extended/improved (BEAM2, BEAM3)
  - Zhang et al. (2011) Ann Appl Stat 5:2052-2077
  - Zhang (2012) Genet Epidemiol 36:36-47

# Empirical evidence for epistasis

- Hypothesis-based studies
  - Several papers by Combarros et al., most notably those part of “The Epistasis Project”
    - An attempt to replicate previous findings of epistasis in Alzheimer's disease, or discover new findings through restricting to candidate genes
    - Some success, but replication evidence quite weak: recommend cautious interpretation

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- Epistasis among *HLA-DRB1*, *HLA-DQA1*, and *HLA-DQB1* in multiple sclerosis (Lincoln et al. 2009 PNAS 106:7542-7547)
- Epistasis between *BANK1* and *BLK* in SLE (Castillejo-Lopez et al. 2012)
- Epistasis between *HLA-C* and *ERAP1* in psoriasis (Strange et al. 2010)
- Epistasis between *HLA-B27* and *ERAP1* in ankylosing spondylitis (Evans et al. 2011)

# Empirical evidence for epistasis (cont.)


- Hypothesis-free studies



- Exhaustive searches in WTCCC (Wan et al. 2010; Lippert et al. 2013) generally find lots of interactions within MHC for type 1 diabetes and rheumatoid arthritis
  - Could represent haplotype effects?
  - Require replication?
- Prabhu and Pe'er (2012) used SIXPAC to identify a pair of interacting SNPs in Bipolar disorder
  - Regions replicated, though actual discovery SNPs did not
- Gusareva et al. (2014) “Genome-wide association interaction analysis for Alzheimer’s disease” found a reasonably convincing (partially replicating) interaction between SNPs on chromosome 6 (*KHDRBS2*) and 13 (*CRYL1*)

# Empirical evidence for epistasis (cont.)

- Hypothesis-free studies

- Hemani et al. 2014 (Nature 508:249-253) found 501 instances of epistatic effects on gene expression, of which 30 could be replicated in two independent samples 
  - Many SNPs are close together, could represent haplotype effects?
  - Or the effect of a single untyped variant?
  - See Wood et al. (2014) Nature 514(7520):E3-5. PMID:25279928
- Brown et al. 2014 (eLIFE 3:e01381) found 508 'candidate' SNPs showing potential interactions ( $G \times G$  or  $G \times E$ ) on gene expression, based on their effect on trait variance
  - Twin studies suggested  $G \times E$  played a role in 70% of these findings (but we don't know what the relevant environmental factors are)
  - 57  $G \times G$  interactions (between specific SNP and gene) replicated in a smaller data set

# Conclusions

- Gene-gene and gene-environment interactions can be modelled in genetic (including genome-wide) association studies
- Computationally intensive if considering large numbers of loci: may need to filter down
- May be worth doing in some situations to increase power to detect effects (but further work needed on optimal search strategies)
  - Utility depends heavily on true underlying genetic model
  - Potentially useful for *detection* of interacting loci
  - Biological interpretation complex...and perhaps better addressed via alternative experiments

