Systems Medicine

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OUTLINE

- Why looking at "interactions"?
- Case study: pancreatic cancer
- MB-MDR: a decade's work; Lessons learned
- Implications
 - Risk prediction
 - Molecular reclassification of disease
 - Omics integration
- Take-home message



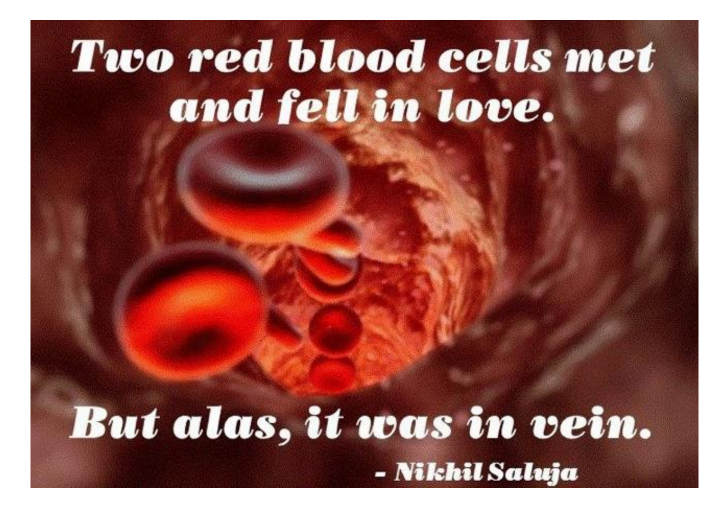




Why looking at "interactions"?

including SNP based

A tale of ... multiple ... stories



Biological interactions

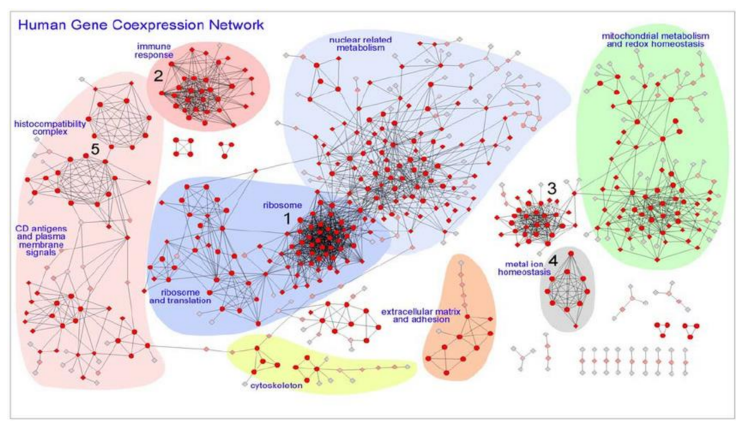
- Biological interactions are the effects that the organisms in a community have on one another. In the natural world no organism exists in absolute isolation, and thus every organism must interact with the environment and other organisms.
- An organism's interactions with its environment are fundamental to the survival of

that organism and the functioning of the ecosystem as a whole.



Gene-gene interactions

• Inference about gene-gene interactions using microarray data

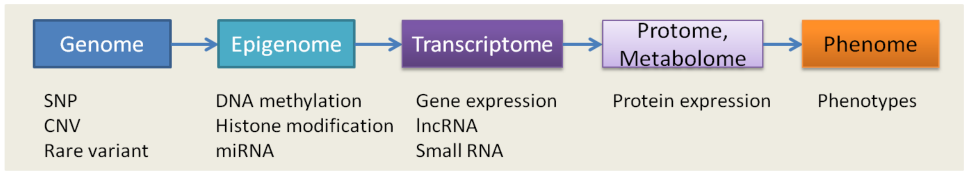


(Prieto et al. 2008)

Omics data as a starting point

- Roughly, omics data is a generic term that describes genome-scale data sets that emerge from high-throughput technologies
- These data describe virtually all biomolecules in a cell (e.g., proteins, metabolites)

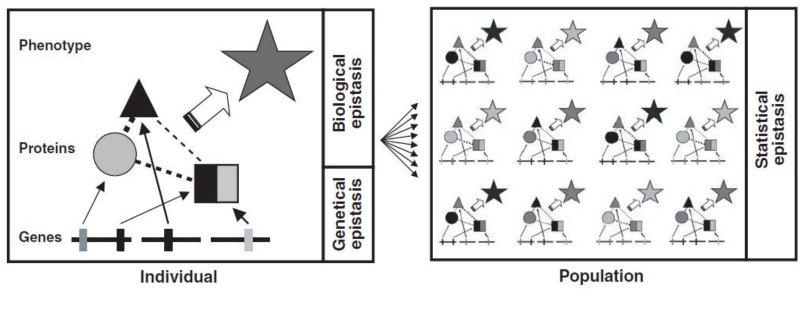
(Joyce and Palsson 2006)



(courtesy figure Maggie Wang)

DNA-DNA interactions

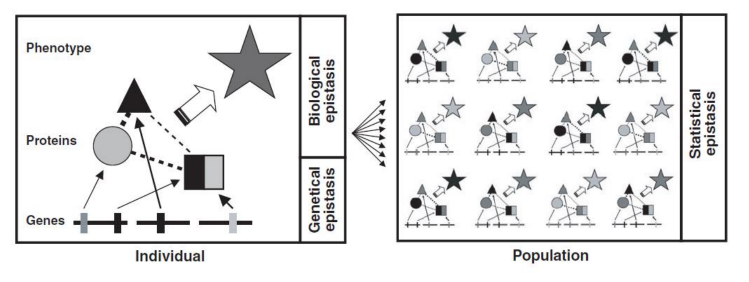
 Two or more DNA variations may "interact" either directly to change transcription or translation levels, or indirectly by way of their protein product (to alter disease risk separate from their independent effects)



(Moore 2005)

Formal definition of epistasis (Moore 2005; Moore and Williams 2005)

- The original definition (**driven by biology**) refers to a variant or allele at one locus preventing the variant at another locus from manifesting its effect (William Bateson 1861-1926).
- A later definition of epistasis (**driven by statistics**) is expressed in terms of deviations from a model of additive multiple effects (Ronald Fisher 1890-1962).



Occurrences of "epistasis" – model organisms

- During HSPH post-doc 2003-2005
 - Epistatic QTLs without individual effects have been found in various organisms, such as birds, mammals, insects (Drosophila Melanogaster – fruit fly) and plants.
 - Other similar studies have reported only low levels of epistasis or no epistasis at all, despite being thorough and involving large sample sizes^{35–37}.
- No single mode of inheritance can be expected to be the rule in all populations and traits; "complex" complex trait regulation; epistatic heterogeneity, which is likely to be contextual

(Carlborg et al. 2004)

Occurrences of epistasis – humans

 Canalization is a form of stabilizing selection to explain the buffering of phenotypes to genetic and environmental perturbations

(Waddington 1942)

Evolution tends to keep our blood pressure and glucose levels within healthy ranges (i.e., evolution of the "system" to a robust level), resistant to most genetic and environmental stimuli

 The consequence is an underlying genetic architecture that is comprised of networks of genes that are redundant and robust (*trans – epistasis*)

(Moore et al. 2009)

Deviations from these healthy ranges are often categorized as "disease", such as hypertension and diabetes

Unexplained heritability (from GWAs)

- The **statistical definition** for heritability defines it as the proportion of phenotypic variance attributable to genetic variance.
- The "sensical" definition defines it as the extent to which genetic individual differences contribute to individual differences in observed behavior (or phenotypic individual differences).
- The proportion of heritability explained by a set of variants is the ratio of (i) the heritability due to these variants (numerator), estimated directly from their observed effects, to (ii) the total heritability (denominator), inferred indirectly from population data.

(Maher 2008, Zuk et al. 2012)

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Unexplained heritability

Explanation	Rationale	Comments
Overestimated heritability	These estimates are typically	Limiting pathway modeling
estimates	performed in the absence of	suggests that epistasis could
	gene-gene or gene-	account for missing
	environment interactions	heritability in complex
	(Young et al. 2014)	diseases (Zuk et al. 2012)
Rare genetic variants	Resequencing studies (e.g.,	Limited evidence for rare
	WES) could identify rare	variants of major effect in
	genetic determinants of large	complex diseases accounting
	effect size (Zuk et al. 2014)	for large amount of genetic
		variation – most rare variants
		analysis methods currently
		suffer from increased type I
		errors (Derkach et al. 2014)
Phenotypic and genetic	Most complex diseases are	Improvements in phenotyping
heterogeneity	like syndromes with multiple	of complex diseases will be
	potentially overlapping	required to understand
	disease subtypes	genetic architecture.

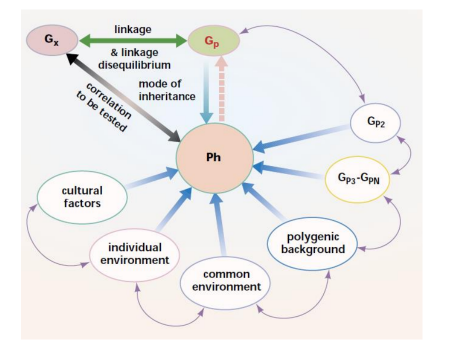
Explanation	Rationale	Comments
Interactions	Gene-gene and gene- environment interactions are likely to be important for complex diseases (Moore et al 2005) Roughly 80% of the currently	Limited <i>replicated</i> evidence for statistical interactions in complex diseases; network-based approaches may be helpful (Hu et al. 2011)
	missing heritability for Crohn's disease could be due to genetic interactions, if the disease involves interaction among three pathways (Zuk et al. 2012)	

(adapted from Silverman et al. 2012)



(Hayden 2010) « Life is Complicated »)

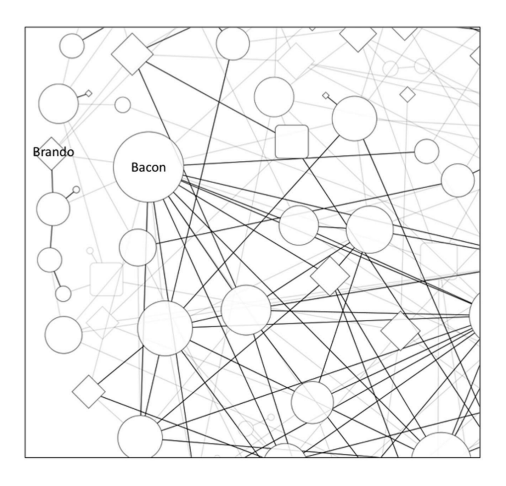
The complexity of complex diseases



(Weiss and Terwilliger 2000)

There are likely to be *many* susceptibility genes each with combinations of rare and common alleles and genotypes that impact disease susceptibility primarily through *non-linear* interactions with genetic and environmental factors (Moore 2008)

Disappointing results for human complex traits [... to date with SNPs]



Edges represent small gene–gene interactions between SNPs. Gray nodes and edges have weaker interactions. Circle nodes represent SNPs that do not have a significant main effect. The diamond nodes represent significant main effect association. The size of the node is proportional to the number of connections.

(McKinney et al. 2012)

Disappointing results for human complex traits [... to date with SNPs]

- Expectations of the first hour seem to be poorly met:
 - Different schools: heritability composition additivity (Polderman et al. 2015: Meta-analysis of the heritability of human traits based on fifty years of twin studies vs relevance of SNP-based interactions) vs epistasis oriented starting point (incl. "Moore et al")

• "LDAK":
$$E[H_{SNP}^2] = (MAF_{SNP}(1 - MAF_{SNP}))^{1+\alpha} w_{SNP}r_{SNP}$$

- "GCTA": $w_{SNP} = 1$; $r_{SNP} = 1$; "LDSC": $\alpha = -1$ + extra parameters

"LDSC" will typically have standard errors 25–100% higher than those from
 "GCTA" (Bulik-Sullivan 2015)
 (Speed et al. 2017)

•
$$H_I^2 = H^2 - H_{M,SNP1}^2 - H_{M,SNP2}^2$$
 (Winham et al. 2012)

$$H_{M,SNP1}^{2} = \frac{1}{P(1-P)} \sum_{i=0}^{2} \left[\sum_{j=0}^{2} Prob(G_{ij}) \right] \left[\sum_{j=0}^{2} Prob(D|G_{ij}) Prob(G_{ij}) - P \right]^{2}$$

Disappointing results for human complex traits [... to date with SNPs]

- Expectations of the first hour seem to be poorly met
 - There is an abundance of methodological approaches
 - Interestingly, widely accepted protocol to perform a Genome-Wide Association Interaction Study (GWAIS) is still lacking
 - Possible explanations:
 - many difficulties (technical, statistical, computational) involved in performing large-scale epistasis screening
 - and in inferring biological evidence from statistical findings

Human Genetics (2019) 138:293–305 https://doi.org/10.1007/s00439-019-01987-w

REVIEW



How to increase our belief in discovered statistical interactions via large-scale association studies?

K. Van Steen^{1,2} · J. H. Moore³

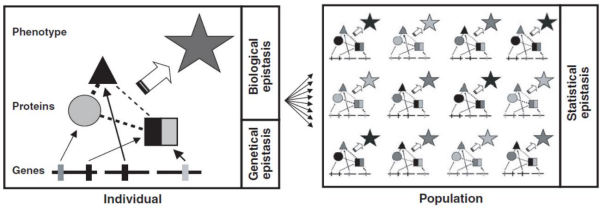
Received: 26 July 2018 / Accepted: 20 February 2019 / Published online: 6 March 2019 © The Author(s) 2019

Abstract

The understanding that differences in biological epistasis may impact disease risk, diagnosis, or disease management stands in wide contrast to the unavailability of widely accepted large-scale epistasis analysis protocols. Several choices in the analysis workflow will impact false-positive and false-negative rates. One of these choices relates to the exploitation of particular modelling or testing strategies. The strengths and limitations of these need to be well understood, as well as the contexts in which these hold. This will contribute to determining the potentially complementary value of epistasis detection workflows and is expected to increase replication success with biological relevance. In this contribution, we take a recently introduced regression-based epistasis detection tool as a leading example to review the key elements that need to be considered to fully appreciate the value of analytical epistasis detection performance assessments. We point out unresolved hurdles and give our perspectives towards overcoming these.

What's in a name ?

• The original definition (**driven by biology**) refers to a variant or allele at one locus preventing the variant at another locus from manifesting its effect (William Bateson 1861-1926).



(Moore 2005)

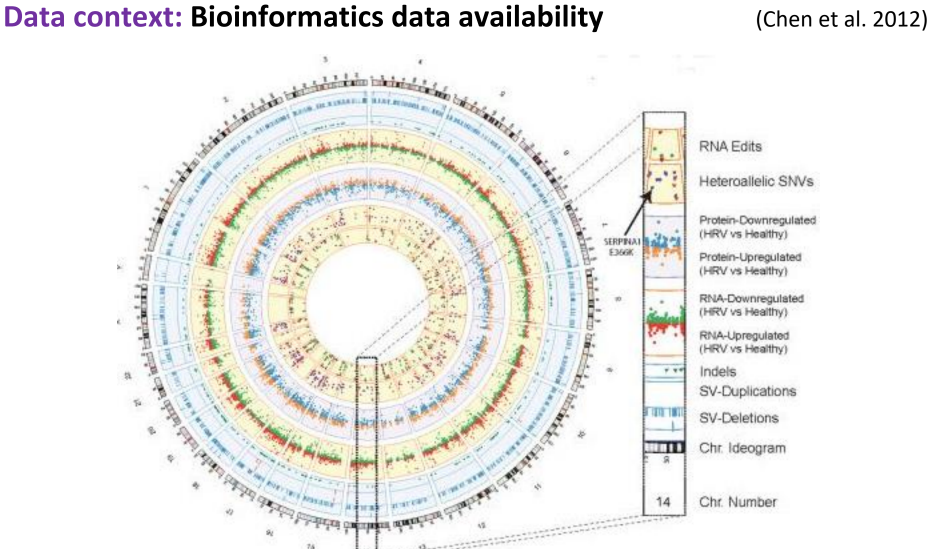
• Grown into a more general theory and applications framework for

the analysis of interactions across and between -omics strata.

Link to "the" interactome

The **interactome** refers to the entire complement of interactions between DNA, RNA, proteins and metabolites within a cell. These interactions are influenced by genetic alterations and environmental stimuli. As a consequence, the interactome should be examined or considered in *particular contexts*.

Case study: Pancreatic Cancer (opportunity)



Disease context: complex "complex diseases"

THE ALARMING RISE OF PANCREATIC GANGER DEATHS IN THE INITED STATES: WHY WE NEED TO STEM THE TIDE TODAY



Addressing complexity in "complex diseases" - pancreatic cancer

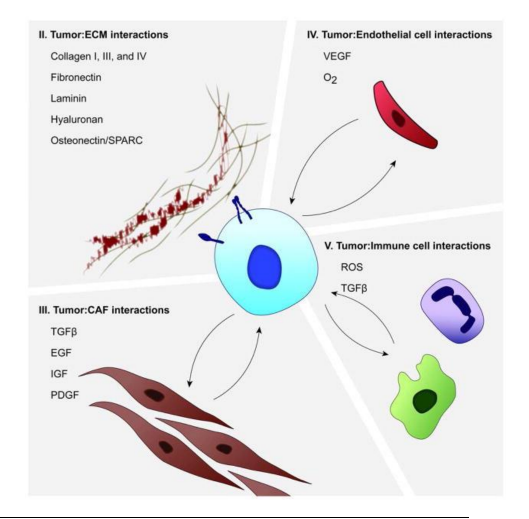
"Because effective systemic therapy capable of controlling the aggressive pancreatic cancer biology is currently lacking, the need for a better understanding of detailed mechanisms underlying pancreatic cancer development and progression is **URGENT**"

(Xie and Xie 2015)

Examples of interactions in pancreatic cancer

Tumor-stromal interactions

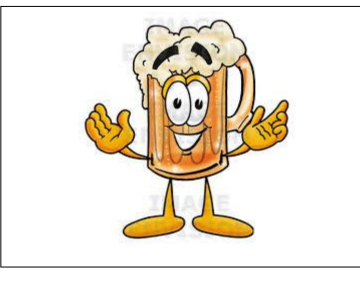
- Treatments focusing on pancreatic cancer cells alone have failed to significantly improve patient outcome over many decades
- Research efforts have now moved to understanding the pathophysiology of the stromal reaction and its role in cancer progression



(Whatcott et al. 2014)

Gene-environment interactions

(Jansen et al. 2015)









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Formal definition of gene-environment interactions

- Also gene-environment interactions can be defined in a statistical or a biological way.
- A **biological gene-environment** interaction occurs when one or more genetic and one or more environmental factors participate in the same causal mechanism in the same individual (Yang and Khoury 1997; Rothman et al. 2008)
- As with gene-gene interactions, a **statistical gene-environment** interaction does not imply any inference about a specific biological mode of action. It is based on modeling a sample of individuals.

Formal definition of epistasis

- In practice, when modeling or testing, it may only be possible to detect **effect modification** from real-life data and not **interaction**, or interaction but not effect modification.
- Whereas an interaction effect for "exposures" X_1 and X_2 relies on a symmetric role for both X_1 and X_2 , an effect modification relies on a conditioning argument (for instance on X_2) (VanderWeele 2009a)
- The distinction between both effect types is often concealed in regression analysis ... (Robins et al. 2000; North et al. 2005)

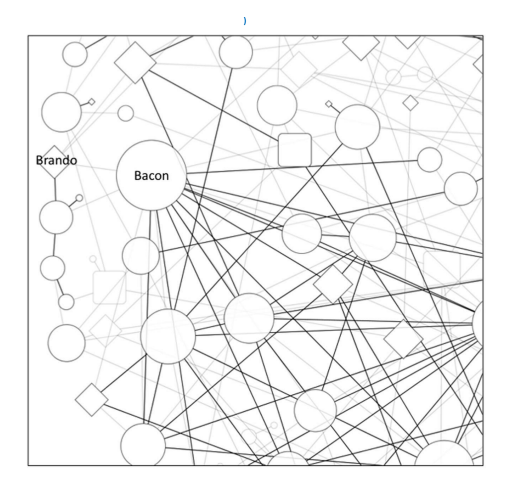
Comparison between gene-gene and gene-environment issues

- Conceptually many similar issues in terms of definition and mathematical modelling.
- In practice, some clear differences emerge.
- For G x E:
 - We generally have to decide which environments to measure / test; these are typically only a few (often < 100)
 - Measurement error (lifestyle) and unknown confounding
 - Risk estimation, important for screening strategies and public health interventions

Comparison between gene-gene and gene-environment issues

- For G x G
 - Assuming we have GWAS data, we have already measured the genetic factors of interest
 - Adequate error rates (except for newer sequencing technologies)
 - (Hundred) thousands of variants
 - Higher-order interactions may reflect the complex biological wiring of complex diseases (whereas G x E often restricts attention to pairwise interactions)

Looking for higher-order interactions



Edges represent small gene-gene interactions between SNPs.

Gray nodes and edges have weaker interactions.

Circle nodes represent SNPs that do not have a significant main effect. The diamond nodes represent significant main effect association.

The size of the node is proportional to the number of connections.

(McKinney et al 2012)

Some references

Published in final edited form as: *Hum Genet.* 2012 October ; 131(10): 1591–1613. doi:10.1007/s00439-012-1192-0.

Challenges and Opportunities in Genome-Wide Environmental Interaction (GWEI) studies

Hugues Aschard¹, Sharon Lutz^{2,*}, Bärbel Maus^{3,4,*}, Eric J. Duell⁵, Tasha Fingerlin², Nilanjan Chatterjee⁶, Peter Kraft^{1,7}, and Kristel Van Steen^{3,4}

Hum Genet (2014) 133:1343–1358 DOI 10.1007/s00439-014-1480-y

REVIEW PAPER

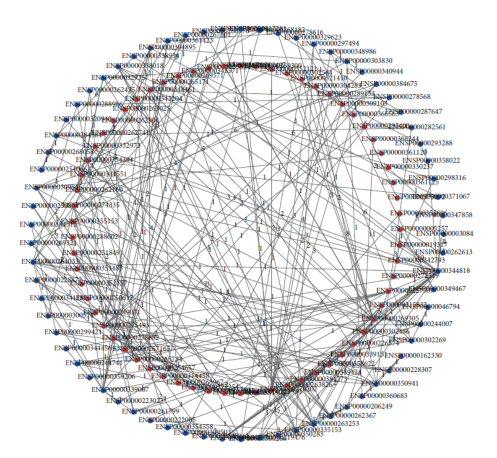
Practical aspects of genome-wide association interaction analysis

Elena S. Gusareva · Kristel Van Steen

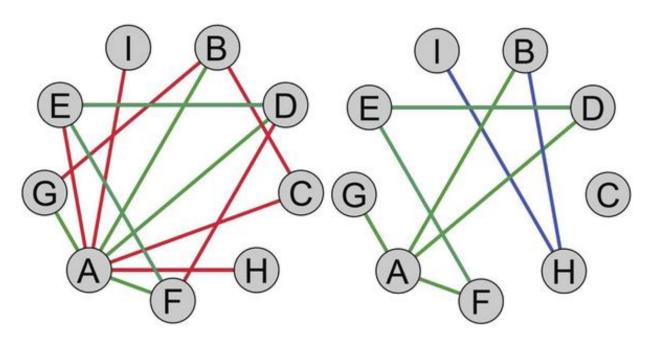
Protein-protein interactions

A graph consisting of 2,080 shortest paths:

- The nodes on the inner circle (red nodes) represent 65 PC-related genes.
- The nodes on the outer circle (blue nodes) represent 69 shortest path genes.
- The numbers on the edges represent the weights of the edges.

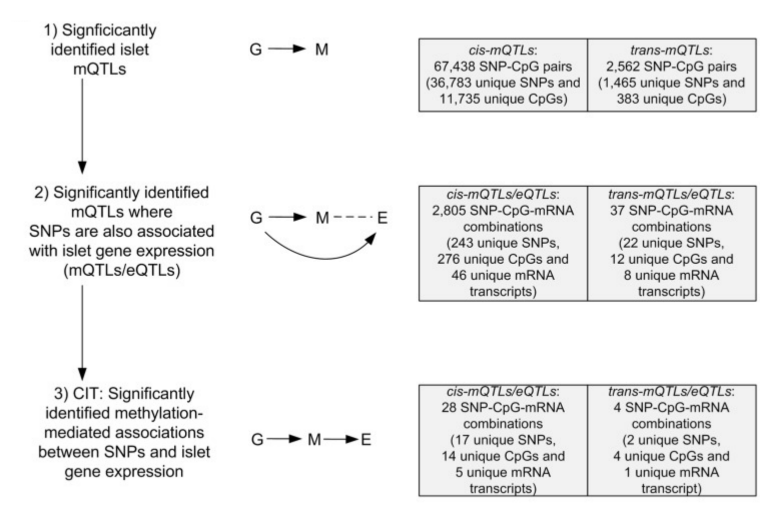


Gene-coexpression networks



- Healthy condition on the left and disease-affected tissue on the right. Green links remain unchanged in the two phenotypes
- Red connections are loss from healthy to cancer network
- Blue edges are novel connections in the cancer tissue

Genetic-epigenetic mechanistic interactions (pancreatic islets)



Gene-gene interactions using SNPs?

(Olsson et al. 2014)

Wolpin BM (PMID: 25086665) 2014-08-03 Nat Genet Genome-wide association study identifies multiple susceptibility loci for pancreatic cancer. 10 • Initial sample description 1,582 European ancestry cases, 5,203 European ancestry controls European (U.S., Australia, France, Germany, Netherlands, Denmark, Finland, Norway, Sweden, U.K., Greece, Italy, Spain) 6,101 European ancestry cases, 9,194 European ancestry controls Replication sample description 6,101 European ancestry cases, 9,194 European ancestry controls Replication sample description 6,101 European (Canada, U.S., France, Germany, Netherlands, Denmark, Finland, Norway, Sweden, U.K., Greece, Italy, Spain) Platform [SNPs Illumina [608202]			
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passing QC]		Platform [SNPs passing QC]	Illumina [608202]

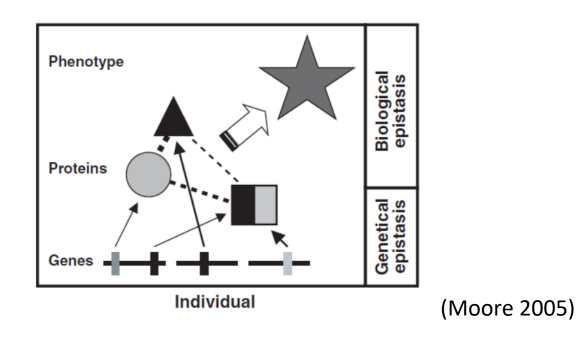
GWAS Catalogue – "Pancreas Cancer"

(http://www.ebi.ac.uk/gwas/search?query=pancreas%20cancer#study)

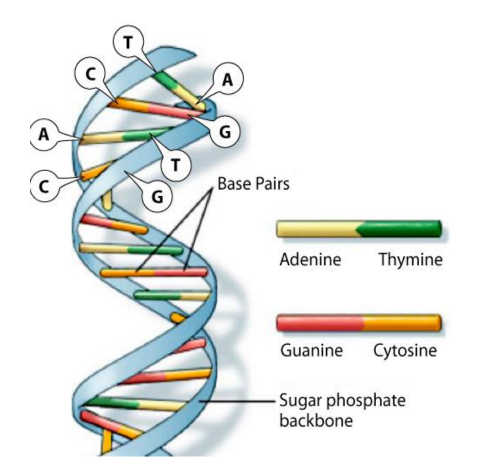
Model-Based Multifactor Dimensionality Reduction a decade's work

Recall DNA-DNA interactions: biological viewpoint

 Two or more DNA variations may interact either directly to change transcription or translation levels, or indirectly by way of their protein product (to alter disease risk separate from their independent effects)



Recall DNA-DNA interactions: non-biological viewpoint



- Epistasis (driven by statistics) is expressed in terms of deviations from a model of additive multiple effects.
- This might be on either a linear or logarithmic scale, which implies different definitions (Ronald Fisher 1890-1962).

(Logistic) Regression

• Alternatively, we can assume additive effects of each allele at each locus, leading to a single interaction term (instead of 4 next!)

		Locus H	
Locus G	2	1	0
2	$\beta_0 + 2\beta_G + 2\beta_H + 4\beta$	$\beta_0 + 2\beta_G + \beta_H + 2\beta$	β_0 + 2 β_G
1	$\beta_0 + \beta_G + 2\beta_H + 2\beta_H$	$\beta_0 + \beta_G + \beta_H + \beta$	$\beta_0 + \beta_G$
0	β_0 + 2 β_H	$\beta_0 + \beta_H$	$eta_{ ext{o}}$

• This corresponds in statistical analysis packages to the model

$$\log\left(\frac{p}{1-p}\right) = \beta_0 + \beta_G X_1 + \beta_H X_2 + \beta X_1 X_2$$

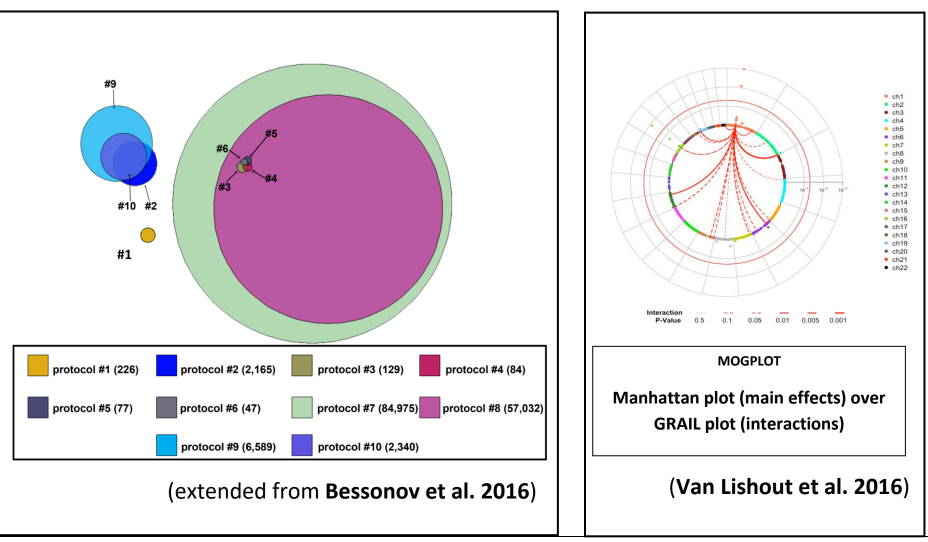
and dosage encoding for X1 and X2.

(Logistic) Regression

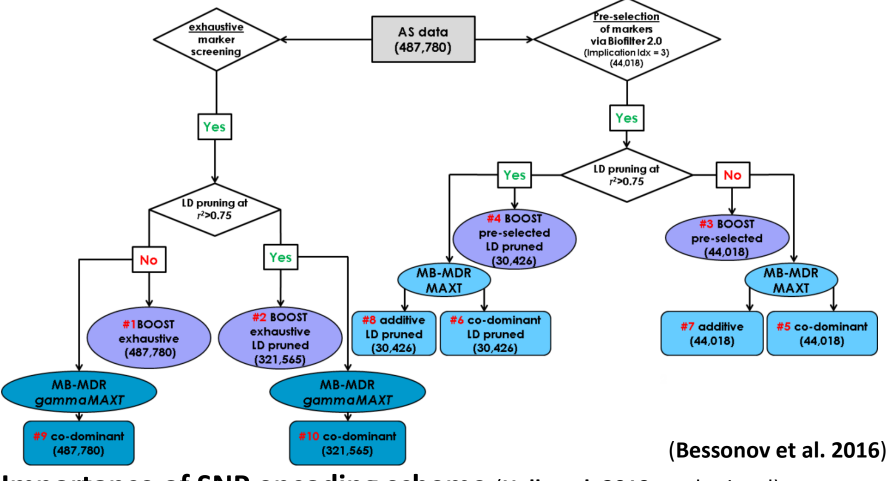
- Most general saturated (9 parameter) genotype model allows all 9 penetrances to take different values
- Log odds is modelled in terms of a baseline effect (β_0), main effects of locus *G* (β_{G1} , β_{G2}), main effects of locus *H* (β_{H1} , β_{H2}), 4 int. terms
- This corresponds in statistical analysis packages to encoding X1, X2 (0,1,2) as a "factor"

		Locus H	
Locus G	2	1	0
2	$\beta_0 + \beta_{G2} + \beta_{H2} + \beta_{22}$	$\beta_0 + \beta_{G2} + \beta_{H1} + \beta_{21}$	eta_0 + eta_{G2}
1	$\beta_0+\beta_{G1}+\beta_{H2}+\beta_{12}$	$\beta_0+\beta_{G1}+\beta_{H1}+\beta_{11}$	eta_0 + eta_{G1}
0	$\beta_0+\beta_{H2}$	eta_0 + $eta_{_{H1}}$	eta_0

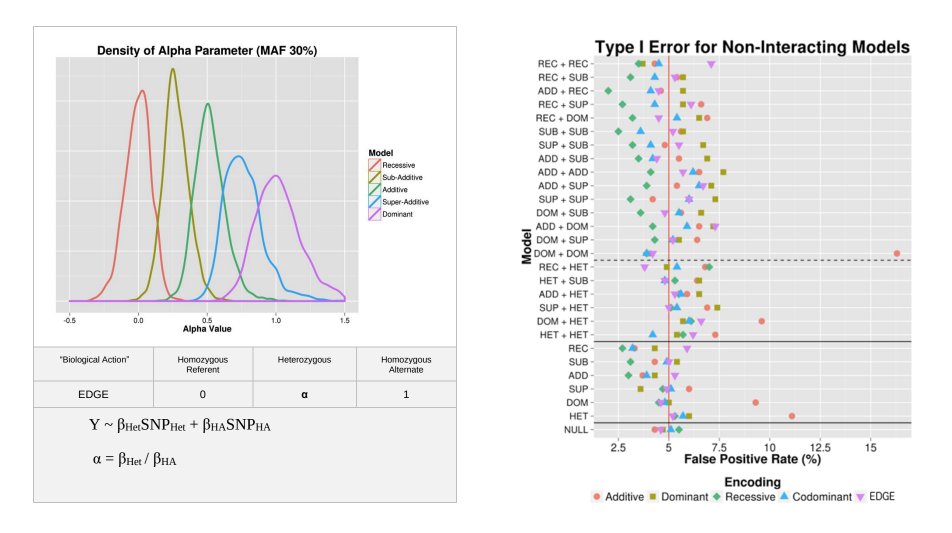
Importance of SNP encoding scheme (Ankylosing Spondylitis; WTCCC2 - ~2000 cases + 5000 controls)



Stability of results: analytic REPLICATION



Importance of SNP encoding scheme (Hall et al. 2019 – submitted)



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Encoding (Hall et al. 2020 – under review)

Table 1. Examples of possible proportional genotype risk

Biological Action	Homozygous Referent (AA)	Heterozygous (Aa)	Homozygous Alternate (aa)
Recessive (REC)	0%	0%	100%
Sub-Additive (SUB	0%	25%	100%
Additive (ADD)	0%	50%	100%
Super- Additive (SUP)	0%	75%	100%
Dominant (DOM)	0%	100%	100%

underlying genetic loci

Binary genomic markers???

Bioinformatics, 33(12), 2017, 1820–1828 doi: 10.1093/bioinformatics/btx071 Advance Access Publication Date: 14 February 2017 Original Paper

Genetics and population analysis

Genome-wide genetic heterogeneity discovery with categorical covariates

Felipe Llinares-López^{1,2,*,†}, Laetitia Papaxanthos^{1,2,*,†}, Dean Bodenham^{1,2}, Damian Roqueiro^{1,2}, COPDGene Investigators³ and Karsten Borgwardt^{1,2,*}

¹Machine Learning and Computational Biology Lab, Department of Biosystems Science and Engineering, ETH Zurich, Basel, Switzerland, ²SIB Swiss Institute of Bioinformatics, Lausanne, Switzerland and ³COPDGene® Study

Note about analytic comparisons: conceptual differences

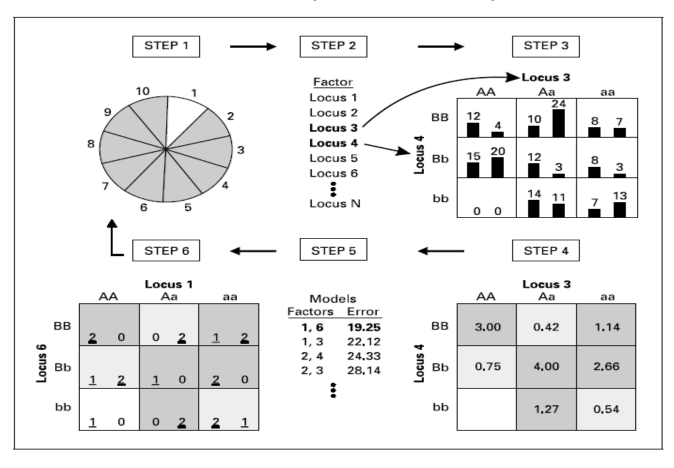
- **Regression modelling:** predicting the average response from covariates [maximizing predictive power] *versus* characterizing relationships [control of confounders, account for effect modifiers]
- **Deep neural networks** have been recognized as some of the best performing machine learning methods:

Methods	Accuracy
Deep learning	68.78
RF	55.85
LR	67.07
Naïve Bayes	62.68
GBM	65.85

(Uppu et al. 2016)

Historical notes about MB-MDR

• Start: Multifactor Dimensionality Reduction by MD Ritchie et al (2001)



Which dimensions are reduced?

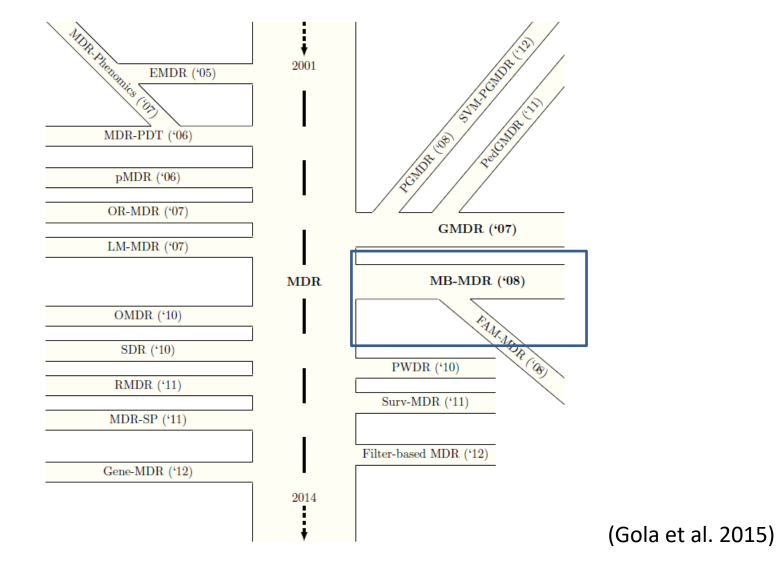
• The estimated degrees of freedom for MDR and LR using K=1, 2 and 3 factors (standard errors in parentheses). LR exact refers to the asymptotic exact degrees of freedom

	Number of Factors K *						
Method	1	2	3				
MDR	1.9 (0.13)	5.6 (0.20)	17.4 (0.37)				
LR	2.1 (0.4)	8.0 (0.26)	26.8 (0.53)				
LR exact	2	8	26				

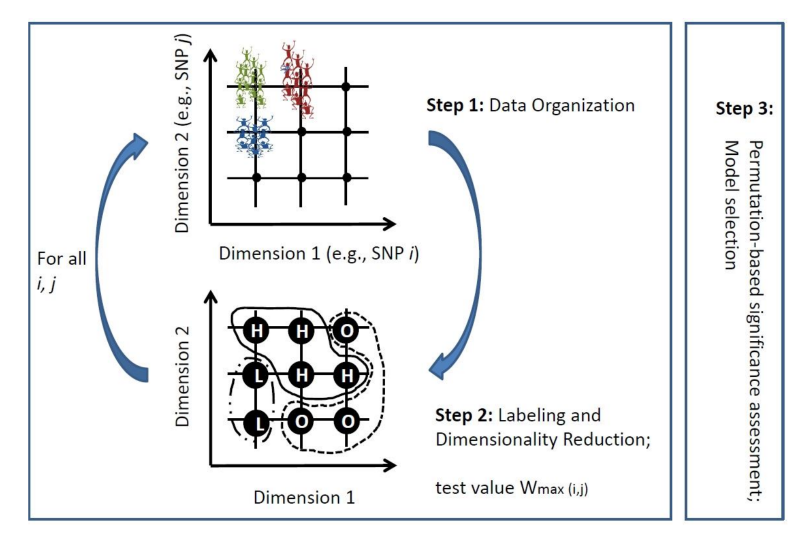
(Park and Hastie 2007)

* "K-way" interaction

Several MDR roads lead to Rome







Current versions of MB-MDR

- Computation time is invested in
 - optimal association tests to label multi-locus genotype combinations and
 - in statistically valid permutation-based methods to assess joint statistical significance of multiple SNP pairs
- Labels are related to substantially improve/worsen trait values (H/L).
 In case there is **no** such **evidence**, the multi-locus label is not forced to be H or L (but will be O).
- In the **presence of main effects**, MB in MB-MDR ensures false positive control at 5%

Global versus specific modeling

• Model-Based MDR by Calle et al (2008a,b)

SNP 40 x SNP 252 genotypes	Cases	Controls	OR	p-value	Category
c1 = (0,0)	88	77	1.01	0.9303	0
c2 = (0,1)	102	114	0.73	0.0562	L
c3 = (0,2)	38	34	0.98	1.0000	0
c4 = (1,0)	50	59	0.76	0.1229	0
c5 = (1,1)	96	37	2.68	0.0000	Н
c6 = (1,2)	18	28	0.55	0.0675	L
c7 = (2,0)	12	6	1.99	0.3399	0
c8 = (2,1)	14	18	0.67	0.3668	0
c9 = (2,2)	6	6	0.84	1.0000	0

 Table 3. MB-MDR first step analysis for interaction between SNP 40

 and SNP 252 in the bladder cancer study

H: High risk; L: Low risk; 0: No evidence

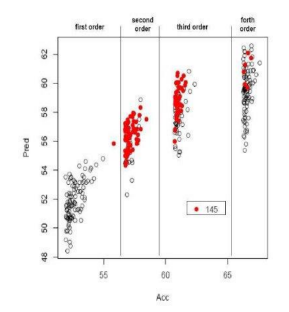
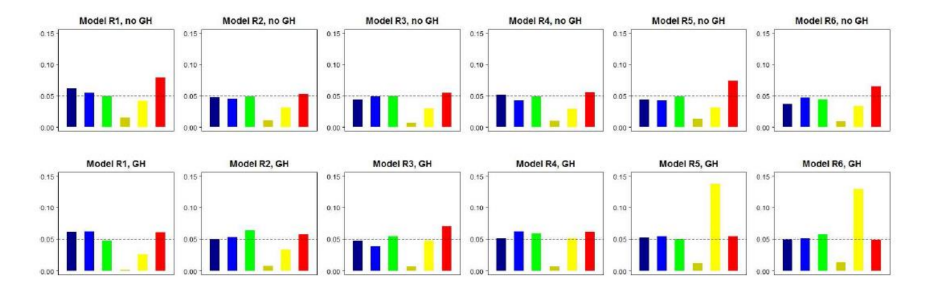


Fig. 1. Average Balanced Training accuracy (Acc) versus Average Balanced Predictive accuracy (Pred) for the 100 models with higher balanced training accuracy for the whole sample. First, second, third and forth order interactions are considered.

Comparative performance of 2-locus MB-MDR

• False positives

(example: pure epistasis scenario's; unpublished - 2010)



BOOST (dark blue) EpiCruncher optimal options (light blue) MB-MDR (green) PLINK epistasis (dark yellow) PLINK fast epistasis (light yellow) EPIBLASTER (red)

Comparative performance of 2-locus MB-MDR

• Power performance

Model R1, no GH Model R2, no GH Model R3, no GH Model R4, no GH Model R5, no GH Model R6, no GH 1.0 1.0 1.0 0.8 0.8 0.8 0.8 0.8 0.8 0.6 0.6 0.6 0.6 0.6 0.4 0.4 0.4 0.4 0.2 0.2 0.2 0.2 Model R1, GH both Model R2, GH both Model R3, GH both Model R4, GH both Model R5, GH both Model R6, GH both 1.0 1.0 1.0 -1.0 1.0 1.0 0.8 0.8 0.8 0.8 0.8 0.8 0.6 0.6 0.6 0.6 0.6 0.6 0.4 0.4 0.4 0.4 0.4 0.4 0.2 0.2 0.2 0.2 0.2 0.2

(example: pure epistasis scenario's; unpublished - 2010)

BOOST (dark blue)

EpiCruncher optimal options (light blue) MB-MDR (green) PLINK epistasis (dark yellow) PLINK fast epistasis (light yellow) EPIBLASTER (red)

Performance summary

Mode	1	Mode	2	Mode	13	Mode	e l 4	Mode	el 5	Mode	el 6
	False Positives (%)										
MB	MDR	MB	MDR	MB	MDR	MB	MDR	MB	MDR	MB	MDR
6	9	4	5	6	17	5	13	5	21	5	23
					Pow	er (%)		·			
MB	MDR	MB	MDR	MB	MDR	MB	MDR	MB	MDR	MB	MDR
100	99	100	100	100	95	100	93	93	62	97	73
MB-MDR (MB): $p_c = 0.1$, T = H vs L test; MDR: default options, screening over 1-5 order models											

(Cattaert et al. 2011)

1	Model 1	p = 0.	5
	BB	Bb	bb
AA	0	0.1	0
Aa	0.1	0	0.1
aa	0	0.1	0

Model 3, p = 0.25 BB Bb bb 0.08 0.07 0.05 AA 0.1 0 Aa 0.1 0.03 0.1 0.04 **aa**

Model 4, n = 0.25

	BB	Bb	bb
AA	0.07	0.05	0.02
Aa	0.05	0.09	0.01
aa	0.02	0.01	0.03

Model 2, p = 0.5

0.1

AA Aa

aai

Í	BB	Bb	bb
	0	0	0.1
	0	0.05	0

0

0

model +, p = orau						
	BB	Bb	bb			
AA .	0	0.01	0.09			
Aa	0.04	0.01	0.08			
aa	0.07	0.09	0.03			

Model 6, $p = 0.1$

	BB	Bb	Bb
AA	0.09	0.001	0.02
Aa	0.08	0.07	0.005
aa	0.003	0.007	0.02



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Model-Based Multifactor Dimensionality Reduction for Rare Variant Association Analysis

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Systems and Modeling Unit, Montefiore Institute, and Bioinformatics and Modeling, GIGA-R, University of Liège, Liège, Belgium



Learning from data

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- Calle M, Urrea V, Malats N, Van Steen K. (2008) Improving strategies for detecting genetic patterns of disease susceptibility in association studies Statistics in Medicine 27 (30): 6532-6546 [MB-MDR with Wald tests and MAF dependent empirical test distributions]
- Calle ML, Urrea V, Van Steen K (2010) mbmdr: an R package for exploring gene-gene interactions associated with binary or quantitative traits. Bioinformatics Applications Note 26 (17): 2198-2199 [first MB-MDR software tool, in R]
- Cattaert T, Urrea V, Naj AC, De Lobel L, De Wit V, Fu M, Mahachie John JM, Shen H, Calle ML, Ritchie MD, Edwards T, Van Steen K. (2010) FAM-MDR: a flexible family-based multifactor dimensionality reduction technique to detect epistasis using related individuals, PLoS One 5 (4). [first implementation of MB-MDR in C++, with improved features on multiple testing correction and improved association tests + recommendations on handling family-based designs]

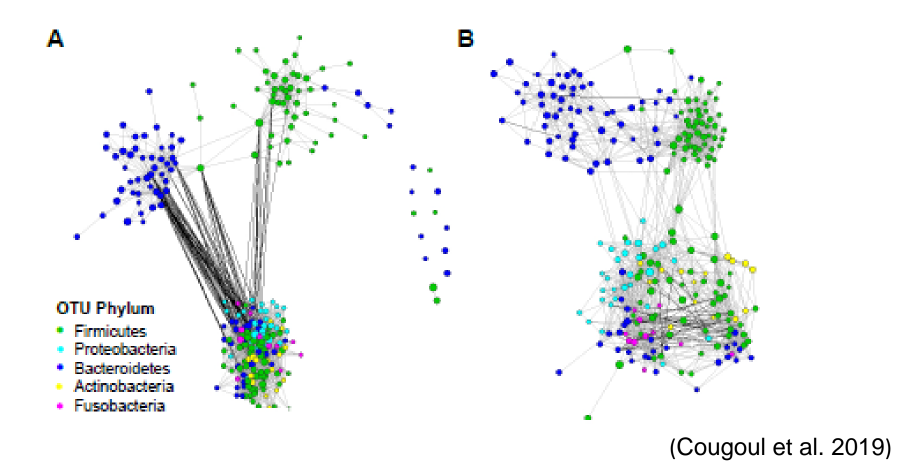
- Cattaert T, Calle ML, Dudek SM, Mahachie John JM, Van Lishout F, Urrea V, Ritchie MD, Van Steen K (2010) Model-Based Multifactor Dimensionality Reduction for detecting epistasis in case-control data in the presence of noise *(invited paper)*. Ann Hum Genet. 2011 Jan;75(1):78-89 [detailed study of C++ MB-MDR performance with binary traits]
- Mahachie John JM, Cattaert T, De Lobel L, Van Lishout F, Empain A, Van Steen K (2011) Comparison of genetic association strategies in the presence of rare alleles. BMC Proceedings, 5(Suppl 9):S32 [first explorations on C++ MB-MDR applied to rare variants]
- Mahachie John JM, Cattaert T, Van Lishout F, Van Steen K (2011) Model-Based Multifactor Dimensionality Reduction to detect epistasis for quantitative traits in the presence of errorfree and noisy data. European Journal of Human Genetics 19, 696-703. [detailed study of C++ MB-MDR performance with quantitative traits]
- Van Steen K (2011) Travelling the world of gene-gene interactions *(invited paper)*. Brief Bioinform 2012, Jan; 13(1):1-19. [positioning of MB-MDR in general epistasis context]
- Aschard et al. ... Van Steen K (2012) Challenges and opportunities in genome-wide environmental interaction (GWEI) studies. Hum Genet. 2012 Oct;131(10):1591-613
 [connection between GxG and GxE interaction detection]

- Mahachie John JM , Cattaert T , Van Lishout F , Gusareva ES , Van Steen K (2012) Lower-Order Effects Adjustment in Quantitative Traits Model-Based Multifactor Dimensionality Reduction. PLoS ONE 7(1): e29594. doi:10.1371/journal.pone.0029594 [recommendations on lower-order effects adjustments]
- Mahachie John JM, Van Lishout F, Gusareva ES, Van Steen K (2013) A Robustness Study of Parametric and Non-parametric Tests in Model-Based Multifactor Dimensionality Reduction for Epistasis Detection. BioData Min. 2013 Apr 25;6(1):9 [recommendations on QT analysis]
- Van Lishout F, Mahachie John JM, Gusareva ES, Urrea V, Cleynen I, Theâtre E, Charloteaux B, Calle ML, Wehenkel L, Van Steen K (2012) An efficient algorithm to perform multiple testing in epistasis screening. BMC Bioinformatics. 2013 Apr 24;14:138 [C++ MB-MDR made faster!]
- Gusareva ES, Van Steen K (2014) Practical aspects of genome-wide association interaction analysis. Hum Genet 133(11):1343-58 [GWAI analysis protocol]
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 Dimensionality Reduction for Rare Variant Association Analysis. Hum Hered 79(3-4):157-67
 [aggregating based on similarity measures to deal with DNA-seq data or multi-omics]
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- Bessonov K, Gusareva ES, Van Steen K (2015) A cautionary note on the impact of protocol changes for genome-wide association SNP x SNP interaction studies: an example on ankylosing spondylitis. Human Genetics 134(7): 761-773.
- Ritchie MD and Van Steen K (2018) The search for gene-gene interactions in genome-wide association studies: challenges in abundance of methods, practical considerations, and biological interpretation. Ann Transl Med 6:157.
- Gusareva et al. ... Van Steen K (2018) Male-specific epistasis between WWC1 and TLN2 genes is associated with Alzheimer's disease. Neurobiol Aging 72:188.e3-188.e1 (2018).

What have we learned?

Tower of Babel - Microbiome interactions (Gusareva)

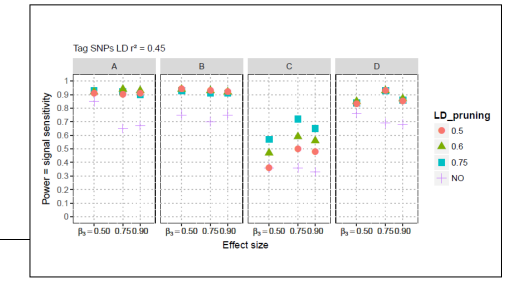


Linkage disequilibrium (LD) as a merit AND a nuisance

Merit: increased signal sensitivity

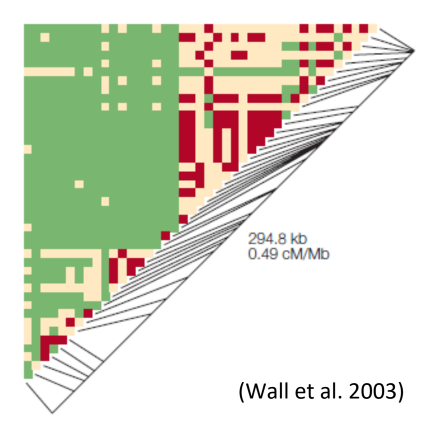
- LD pruning to avoid/reduce confounding between LD and epistasis (*redundant epistasis*)
- LD blocks to capture signals (Joiret et al 2019 under review):
 - Exact signal sensitivity may be low when actual actors were pruned out
 - No pruning gives the lowest signal sensitivity
 - Sufficient pruning gives acceptable signal sensitity

Lowest power when DSLs
 reside at the boundaries of
 LD regions (scenario C)



Different ways to determine LD blocks

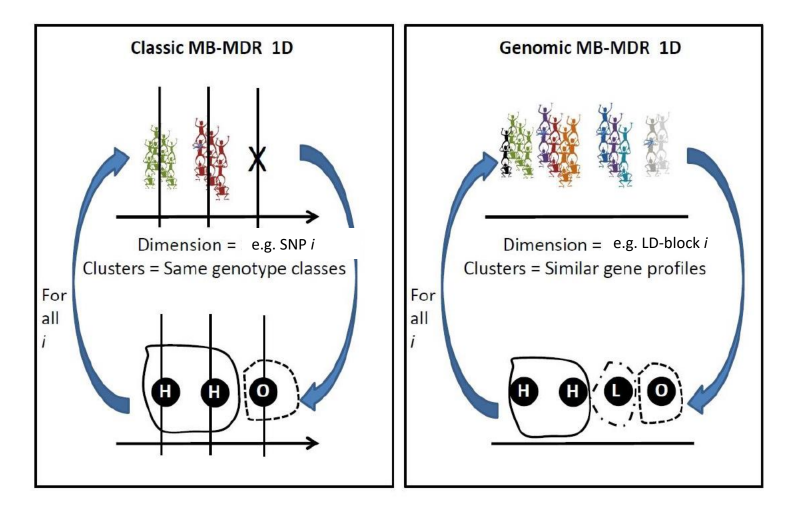
- LD-block computation methods
 - often do not allow
 intermediate regions of low
 LD between strongly
 associated SNP pairs:
 - small blocks,
 - high between-block
 correlations (Kim et al 2018)



Different ways to determine LD blocks (Junior et al.)

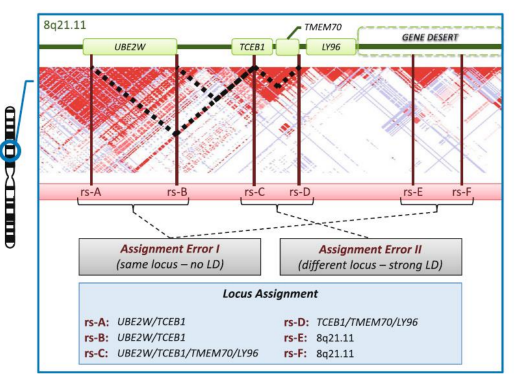
- **Big-LD** produces larger LD blocks compared to existing methods (e.g. MATILDE, Haploview, MIG ++, or S-MIG ++); LD blocks better agree with recombination hotspot locations determined by sperm-typing experiments; per population (Kim et al. 2018)
 - Adaption in progress to facilitate downstream analysis and processing multi-ethnic groups jointly (e.g., multi-population GWAIS):
 - Population-corrected r², using genetic origin / admixture proportions of individual genomes (Mangin et al. 2012)

Big-LD and MB-MDR ← → "phantom epistasis (de los Campo et al. 2019)"

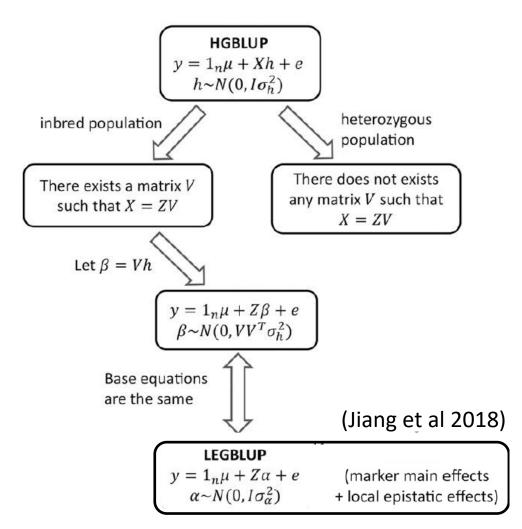


Big-LD and MB-MDR $\leftarrow \rightarrow$ classical enrichment (Junior et al.)

- LD based locus assignment and its error sources
 - whole genomic region
 captured by SNPs in strong
 LD (say r² ≥0.8) with the
 marker originally reported
 in a GWAS
 - loci: genes located within this region
 - (Arnold et al 2012)



Big-LD and MB-MDR $\leftarrow \rightarrow$ within block epistasis

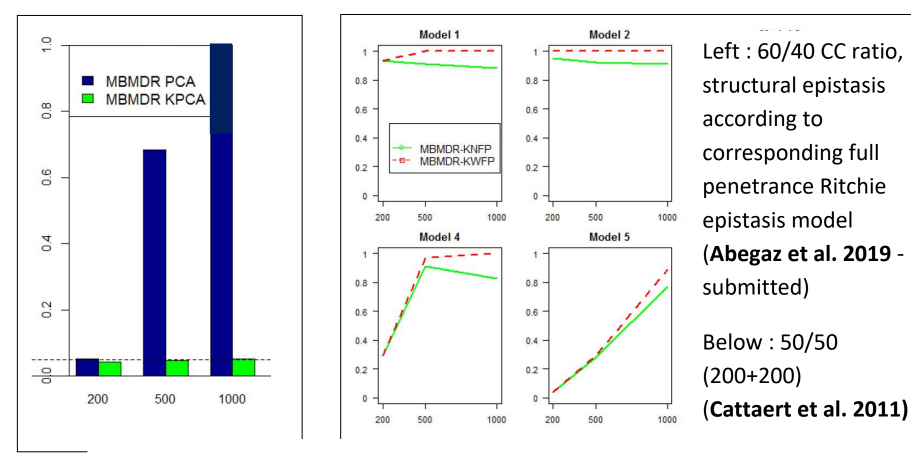


Published in final edited form as: *Nature*. 2014 April 10; 508(7495): 249–253. doi:10.1038/nature13005.

in humans Detection and replication of epistasis influencing transcription

Metspalu⁴, Powell^{1,2,} Henders⁷ Gibran Hemani^{1,2,*} , Allan F. Lude Franke³, Grant W. Montgomery^{7,+}, , Konstantin Shakhbazov^{1,2}, Harm-Jan Westra³, T McRae^{1,2}, Jian Yang¹, Greg Gibson⁸, Nicholas G. Peter M Visscher^{1,2,+}, and Joseph E Tonu Esko^{4,5,6}, Anjali K Martin⁷, Andres

Nuisance: Inadequate capturing of population structure (no cat cov!)



	Model 1		Model 2		Model 3		Model 4		Model 5		Model 6	
Noise	MB	MDR										
None	100	99	100	100	100	95	100	93	93	62	97	73

Van Steen K

Non-continuous axes of genetic variation ???

Bioinformatics, 33(12), 2017, 1820–1828 doi: 10.1093/bioinformatics/btx071 Advance Access Publication Date: 14 February 2017 Original Paper

Genetics and population analysis

Genome-wide genetic heterogeneity discovery with categorical covariates

Felipe Llinares-López^{1,2,*,†}, Laetitia Papaxanthos^{1,2,*,†}, Dean Bodenham^{1,2}, Damian Roqueiro^{1,2}, COPDGene Investigators³ and Karsten Borgwardt^{1,2,*}

¹Machine Learning and Computational Biology Lab, Department of Biosystems Science and Engineering, ETH Zurich, Basel, Switzerland, ²SIB Swiss Institute of Bioinformatics, Lausanne, Switzerland and ³COPDGene® Study

Lack of obvious correspondence between biology and statistics

- From the literature (~ interaction-specific vs two-locus hypotheses):
 - Siemiatycki and Thomas (1981) Int J Epidemiol 10:383-387
 - ...
 - Moore and Williams (2005) BioEssays 27:637–646
 - Phillips (2008) Nat Rev Genet 9:855-867
 - Clayton DG (2009) PLoS Genet 5(7): e1000540
 - Wang, Elston and Zhu (2010) Hum Hered 70:269-277

- ...

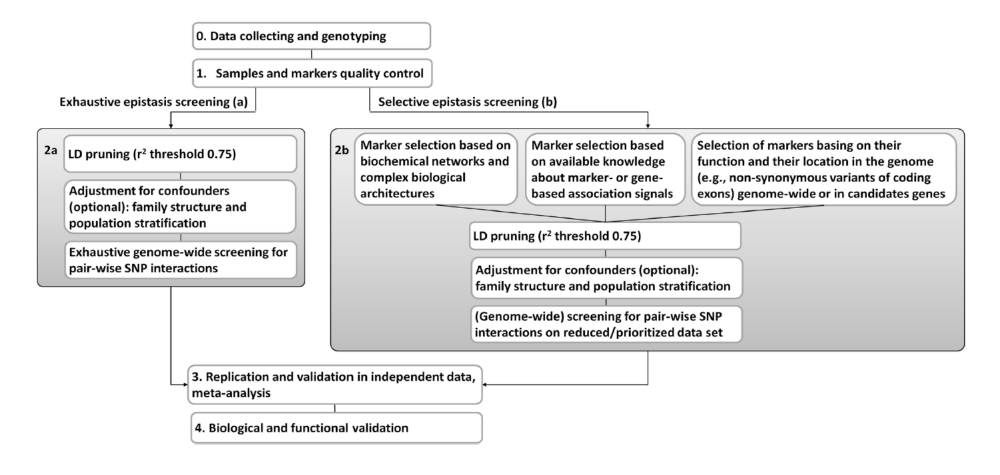
- Van Steen et al. (2012) Brief Bioinform. 13(1):1-19
- Aschard et al. (2012) Hum Genet 131(10):1591-1613
- Gusareva and Van Steen (2014) Hum Genet 133(11):1343-58
- In either case: statistical interactions DO imply JOINT involvement

Understanding molecular mechanisms of epistasis [→ Vidal lab]

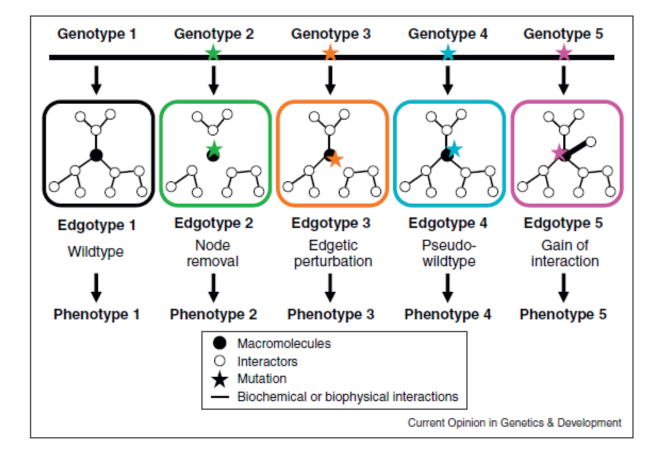
- Best evidence for pervasiveness of epistasis (wrt strong LOF mutations) are derived from large-scale reverse genetics screens
 Pairs of mutations (or RNA interference treatments) are systematically combined and effects on viability or growth are determined (yeast: Costanzo et al. 2010)
- Conclusion:
 - For majority of LOFs the effect can be influenced by perturbing the activity of many additional genes
 - Mutations with weaker effects on proteins are understudied. It needs to be determined what their potential epistatic role is.
 - Multiple molecular mechanisms underly similar epistatic interactions

(Lehner 2011)

A priori use of biological (functional) knowledge



(Gusareva et al. 2014)



Knowledge boosting: PPI perturbations

(Sahni et al. 2013)

The space of PPI perturbations

Multi-scale perturbations of protein interactomes reveal their mechanisms of regulation, robustness and insights into genotype-phenotype maps

Marie Filteau,* Hélène Vignaud,* Samuel Rochette,* Guillaume Diss,* Andrée-Ève Chrétien,* Caroline M. Berger and Christian R. Landry

Corresponding author: Christian Landry, Département de Biologie, Institut de Biologie Intégrative et des Systèmes, Université Laval, Room 3106, Pavillon Charles-Eugène-Marchand 1030, Avenue de la Médecine, Québec (Québec) G1V 0A6, Canada. Tel.: 418-656-3954; Fax: 418-656-7176; E-mail: Christian.landry@bio.ulaval.ca

*These authors contributed equally to this work.

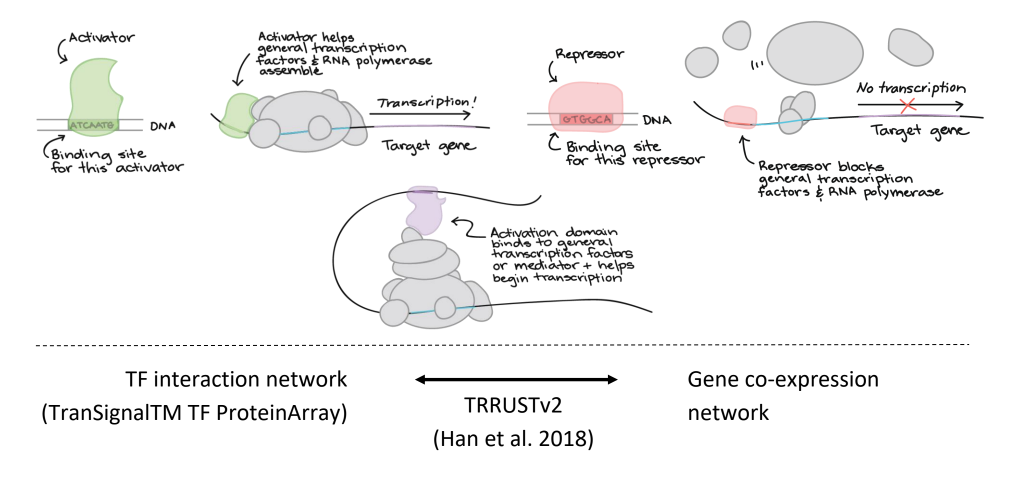
(Filteau et al. 2015)

The space of PPI perturbations

- Zhong et al. 2009: Interactomes of 29 mutant alleles of genes implicated in five human Mendelian disorders. Each allele was cloned and its interactions tested with ~ 8100 other proteins (edgetic perturbations)
 - Different mutant alleles of the same protein can cause different perturbations on their PPI profiles
- Sahni et al. 2015: ~ 2500 mutant alleles of proteins implicated in human diseases and their ~1000 corresponding wild-type proteins;
 Y2H to test the interactions of these proteins with a set of ~ 7200 human ORFs
 - ~ One third of the mutations, the cause of the disease may result from perturbations of PPIs

"Message passing" between layers of information

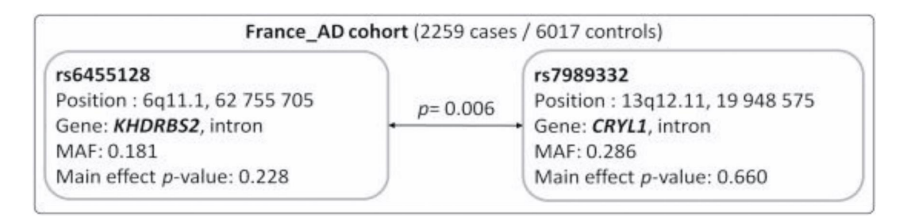




Replication and validation

Genome-wide association interaction analysis for Alzheimer's disease

Elena S. Gusareva^{1,2}, Minerva M. Carrasquillo³, Céline Bellenguez^{4,5,6}, Elise Cuyvers^{7,8}, Samuel Colon³, Neill R. Graff-Radford⁹, Ronald C. Petersen¹⁰, Dennis W. Dickson³, Jestinah M. Mahachie Johna^{1,2}, Kyrylo Bessonov^{1,2}, Christine Van Broeckhoven^{7,8}, The GERAD1 Consortium, Denise Harold¹¹, Julie Williams¹¹, Philippe Amouyel^{4,5,6}, Kristel Sleegers^{7,8}, Nilüfer Ertekin-Taner⁹, Jean-Charles Lambert^{4,5,6}, and Kristel Van Steen^{1,2}



Which extent of replication is required?

4. Replication analysis with alternative methods for epistasis detection: follow up the selected set of markers (MB-MDR_{2D} analysis, SD plot,

logistic regression-based methods)

5. Replication of epistasis in the independent data and biological validation "This study in particular demonstrates an alternative approach to elucidate the functional repercussions of epistasis."

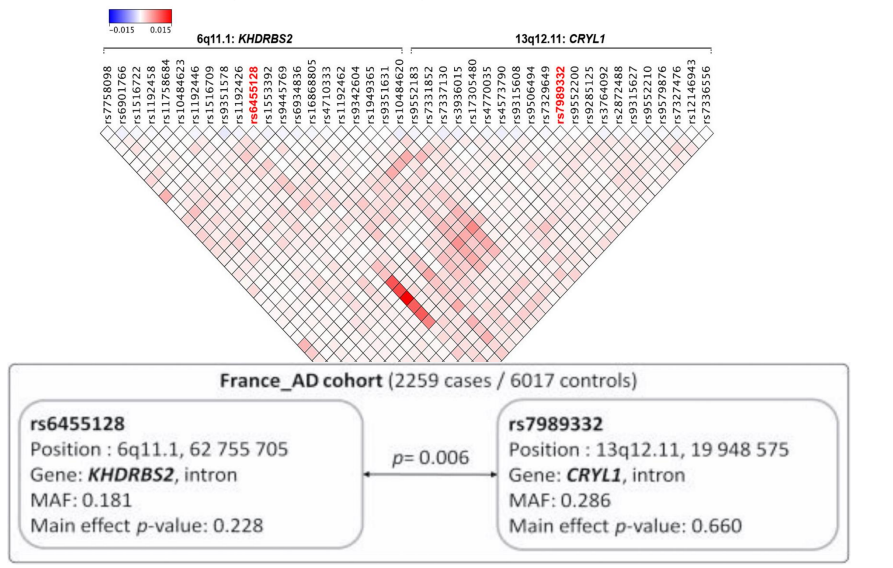
Review Article

Bridging the Gap between Statistical and Biological Epistasis in Alzheimer's Disease

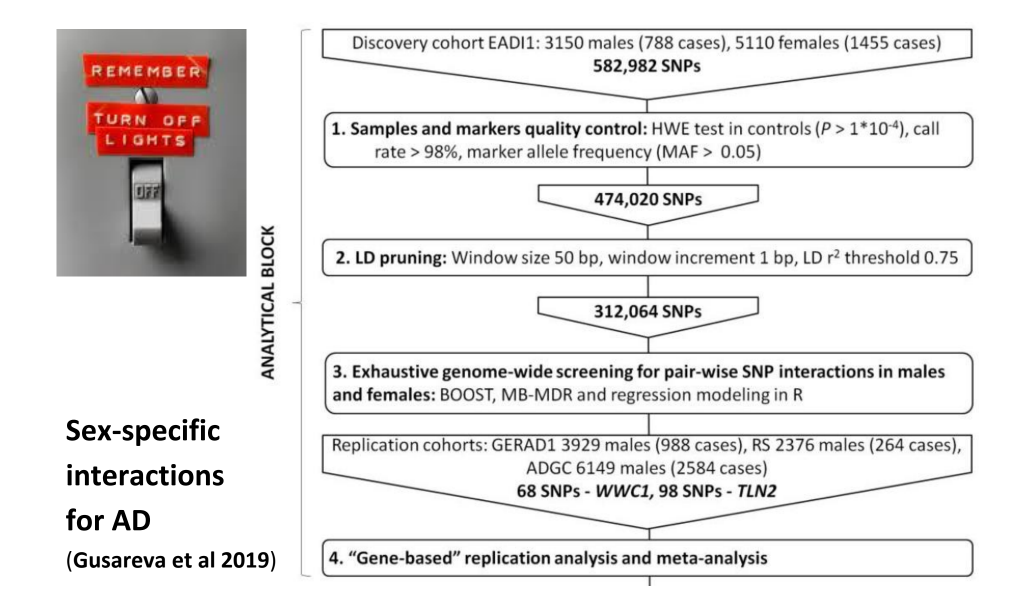


Mark T. W. Ebbert, Perry G. Ridge, and John S. K. Kauwe

Which extent of replication is required?



Van Steen K



Which extent of (experimental) validation is required?

4. "Gene-based" replication analysis and meta-analysis

5. Biological validation of statistical epistasis (series of functional analyses): Transcriptome analysis to assess co-expression of WWC1 and TLN2 in brain tissues of AD and non-AD subjects

Experiments in model organisms (i.e., Tau toxicity in the Drosophila eye) to test whether WWC1 and TLN2 can modulate AD physiopathology

Immunofluorescence and confocal microscopy to confirm presence of WWC1 and TLN2 in human brain cells and to assess their co-localization in common cellular compartments

Immunoprecipitation analysis to confirm physical interaction between WWC1 and TLN2 in a real biological system

Protein docking and molecular dynamics analysis to get more inside into mechanisms of the physical interaction between WWC1 and TLN2

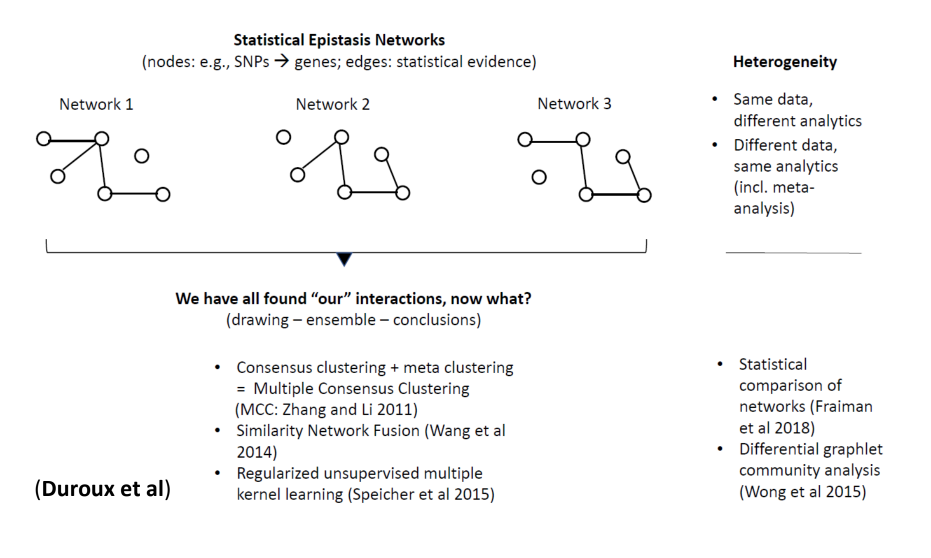
(Gusareva et al 2019)

Functional in-silico analyses - pairs or unique SNP set from pairs

- Hemani et al. 2014:
 - Are the SNPs in transcriptionally active regions?
 - Cell-type specific enrichment of active chromatin
 - Colocalization of corresponding chromosomal regions in a cell?
 - Which genomic features are covered by the index SNPs (LD r²>0.8;
 0.5Mb)
 - Predicted promoter region including TSS, Predicted promoter flanking region, Predicted enhancer, Predicted weak enhancer or open chromatin cis regulatory element, CTCF enriched element, Predicted transcribed region, or Predicted Repressed or Low Activity region positions.
 - Enrichment of transcription factor binding sites around index SNPs?
- Final protocol paper: pre (Moore et al.) / post in silico (Hemani et al.);

BIOGRID with stringent validation criteria; and minimal lab work (Gusareva et al.)

Epistasis meta-analysis is hampered by analytic heterogeneity



Computational efficiency – pragmatic consideration

- Graphics processing units (GPUs), as alternative powerful and cost-effective parallel processing units (Putz et al. 2013)
- Cloud computing infrastructures, although these do not offer unlimited possibilities (Wang et al. 2011)
- Hardware oriented solutions, such as those based on field-programmable gate array (FPGA) architecture (Gundlach et al. 2016)

Computational feasibility

Multiple testing correction via "MAXT" in MBMDR-3.0.3:

	Sequential version	Sequential version	Parallel workflow	Parallel workflow
SNPs	Binary trait	Continuous trait	Binary trait	Continuous trait
10^{2}	$45 \mathrm{sec}$	$1 \min 35 \sec$	< 1 sec	< 1 sec
10^{3}	1 hour 16 min	2 hours 39 min	$38 \sec$	$1 \min 17 \sec$
10^{4}	5 days 13 hours	11 days 19 hours	1 hour 3 min	2 hours 14 min
10^{5}	≈ 1.5 year	≈ 3 years	$4~{\rm days}~9~{\rm hours}$	$\approx 9 \text{ days}$

The parallel workflow was tested on a cluster composed of 10 blades, containing each four Quad-Core AMD Opteron(tm) Processor 2352 2.1 GHz. The sequential executions were performed on a single core of this cluster. The results prefixed by the symbol " \approx " are extrapolated.

(Van Lishout et al. 2013)

Computational feasibility: approximating vs exact

Multiple testing correction via "gammaMAXT" in MBMDR-4.2.2:

	Sequential version	Parallel workflow	Sequential version	Parallel workflow
SNPs	Binary trait	Binary trait	Continuous trait	Continuous trait
10^{3}	$13 \min 33 \sec$	$20 \sec$	$13 \min 18 \sec$	18 sec
10^{4}	$52 \min 15 \sec$	$1 \min 05 \sec$	$56 \min 14 \sec$	$53 \mathrm{sec}$
10^{5}	64 hours 35 min	$22 \min 15 \sec$	70 hours 03 min	$20 \min 28 \sec$
10^{6}	$\approx 270 \text{ days}$	25 hours 12 min	$\approx 290 \text{ days}$	24 hours 06 min

The parallel workflow was tested on a 256-core computer cluster (Intel L5420 2.5 GHz 1333 MHz FSB). The sequential executions were performed on a single core of this cluster. The results prefixed by the symbol " \approx " are extrapolated.

(Van Lishout et al. 2015)

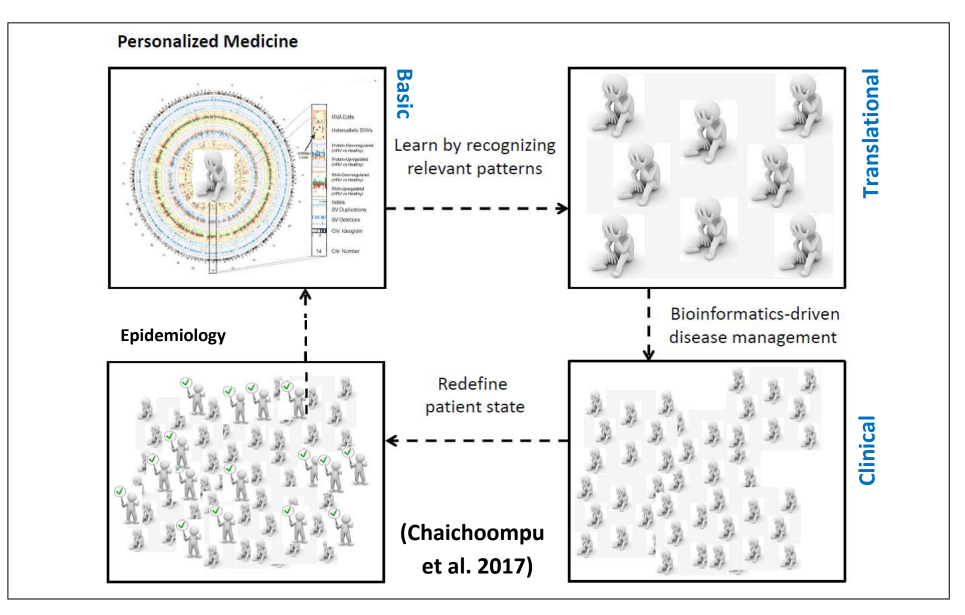
What are the implications?

Individual risk (trait) prediction – deep MBMDR (~McGill,CA)

- MB-MDR was never tested for its predictive ability
 - Note: MDR uses cross-validation and prediction accuracy as measure to select the most optimal interaction model
- Collaborators extended **MB-MDR** to generate **prediction rules**
- The new algorithm (available in R) can use information hidden in interactions more efficiently than two other state-of-the-art algorithms; it clearly **outperforms Random Forest and Elastic Net** if interactions are present.
- The performance of these algorithms is comparable if no interactions are present

(Gola et al. 2019)

Molecular reclassification of disease [→ Sharma lab, Michael Cho]



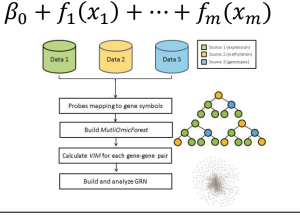
Integration - GRNs by integrating multi-omics data

Integrative networks approaches

- Adapt Lasso? Nine "Lasso's" compared incl new LABNet for gene expression networks (paper: Gadaleta)
 - Multicollinearity + highdimensionality are concern for all
 - GO LABNet (MCC: correl. coefficient between observed and predicted binary classifications)
- New computational approach to analyse gene expression and methylation profiles via regression analysis and network-based techniques - Regression2Net (paper: Gadaleta; Bessonov)
 - SNF between EE-net and EM-net

Conditional Inference Forests (CIFs)

- RF VIMs: bias towards correlated predictor variables. Created **new VIMs for CIFs**
 - Optimal performance of CIF_{cond}
 followed by CIF_{mean} (paper: Bessonov)
 - Bring EXTRA trees idea in CIF
- MultiOmicForest (thesis : Bessonov)
 - Extends CIF_{mean} to 3-omics (diff scales)
 - Non-linearity via GAMs: g(E(Y)) =

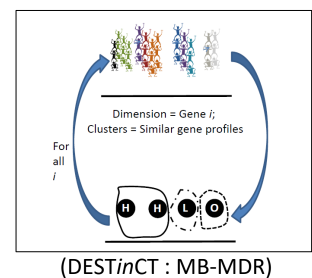


BIO3's approach

• Data integration (heterogeneous data types) – WELL PROGRESSING

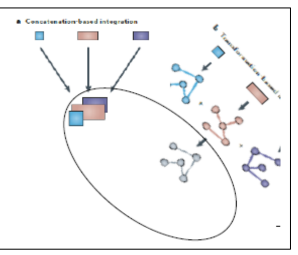
Ex: MB-MDR + defining a smaller "system"

to create omics integrative units of analysis



- Component-based
- Kernel-based (big)
- Network-based

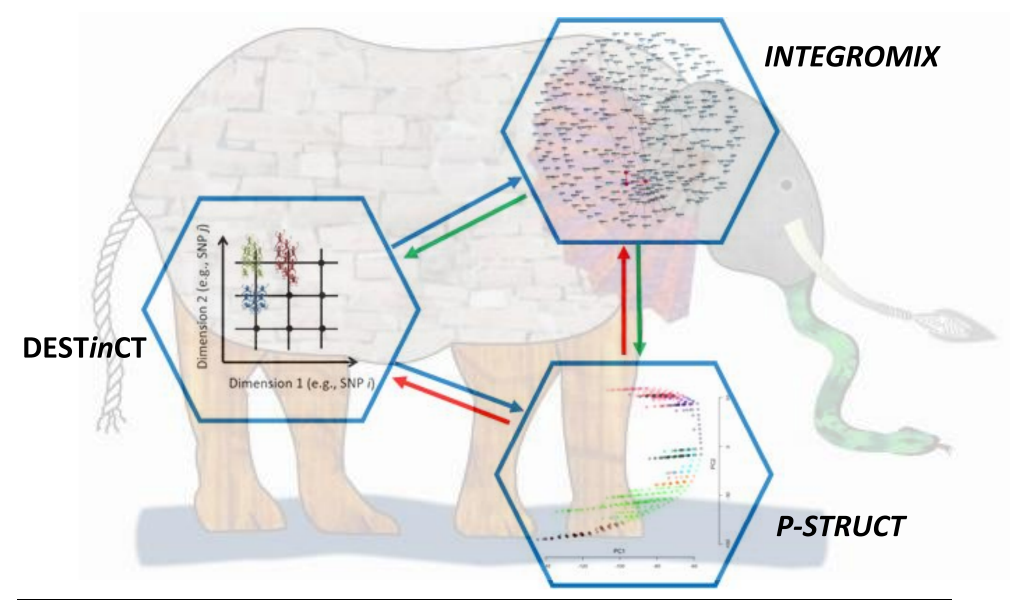
(PhD thesis Fouladi 2018)



(Ritchie et al. 2015)

• Analytic integration (modelling paradigms) – INFANCY

Take-home messages



Through the looking-glass

Alice doesn't play by the conventional rules of a little girl during the 1800s; she's up for whatever comes her way and is willing to take a chance on the unexpected with brilliant results. (Lewis Carroll)



In Michael Guriade

Questions?

Main supporting doc to this class (complementing course slides)

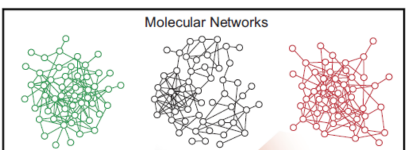


Cell Systems

Article

Systematic Evaluation of Molecular Networks for Discovery of Disease Genes

Graphical Abstract



Authors

Justin K. Huang, Daniel E. Carlin, Michael Ku Yu, Wei Zhang, Jason F. Kreisberg, Pablo Tamayo, Trey Ideker

Correspondence jkh013@ucsd.edu

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Human Genetics (2019) 138:293-305 https://doi.org/10.1007/s00439-019-01987-w

REVIEW



How to increase our belief in discovered statistical interactions via large-scale association studies?

K. Van Steen^{1,2} · J. H. Moore³