MANCHESTER



#### SOC-B CENTRE FOR DOCTORAL TRAINING IN **BIOSOCIAL RESEARCH**

MODULE 2: 'OMICS, SYSTEMS BIOLOGY & BIO **INFORMATICS** 

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## INTRODUCTIONS

## Overview

- What is biological information?
- Types of 'omic data
- Basic analysis and tools
- Network analysis of 'omic data

# WHAT IS BIOLOGICAL INFORMATION?

## Genotype vs. Phenotype

#### Genotype

- The genetic makeup of an organism
  - An organisms complete set of genes
  - Instructions for building and maintaining
    - Formation of proteins, regulation of metabolism
  - Genetic traits
  - Internally coded- not observed
  - Copied during cell division & reproduction

#### Phenotype

- Observable physical properties of an organism
  - Appearance, development & behavior



Phenotype is determined by an organisms genotype and ALSO environmental factors

**Genes + Environment = Phenotype** 

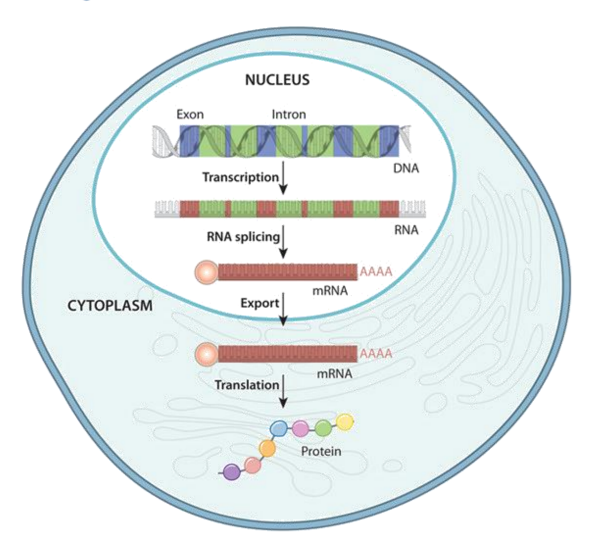


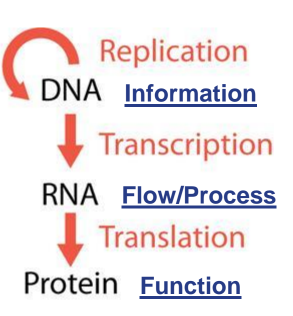
### **Central Paradigm of Molecular Biology**

Genes to messenger to proteins

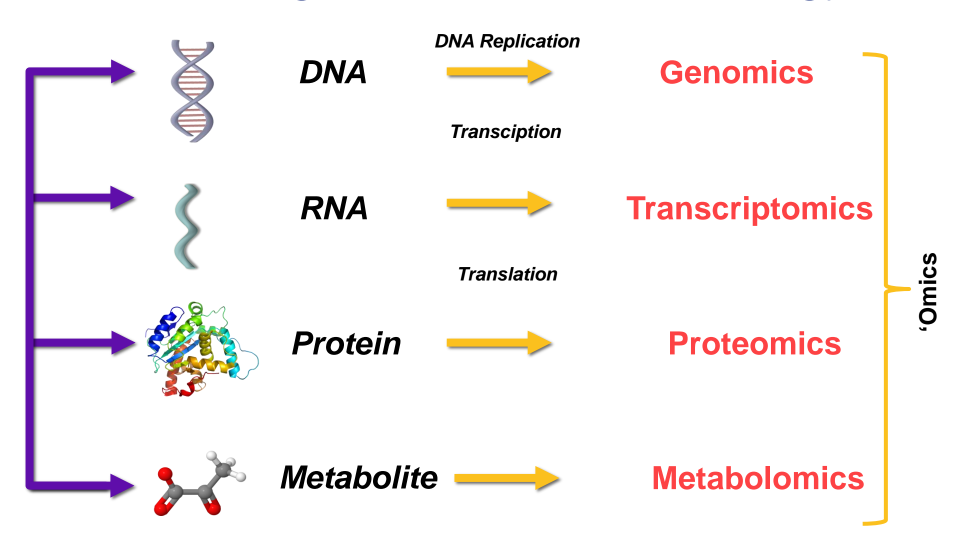
DNA → Protein

**Biological information flow** 





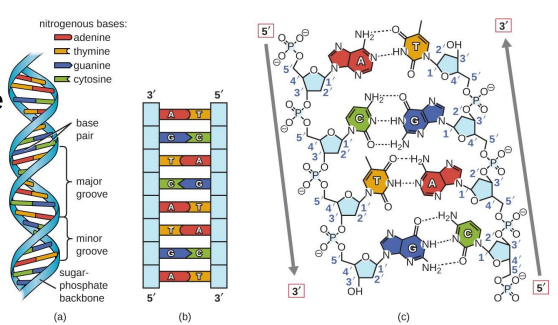
## Central dogma of molecular biology





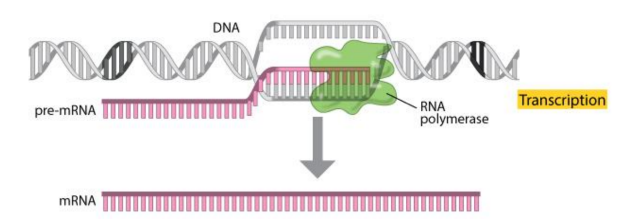
## **DNA & Genomics**

- Deoxyribonucleic acid (DNA)
  - Polymer of nucleotides
  - Sequence of nucleotides is responsible for carrying and retaining hereditary information in a cell – Base Sequence
- Double helix of complementary base pairs
- Nitrogenous base
  - Adenine and Thymine
  - Cytosine and Guanine
- Phosphate group
- Deoxyribose



## **RNA & Transciptomics**

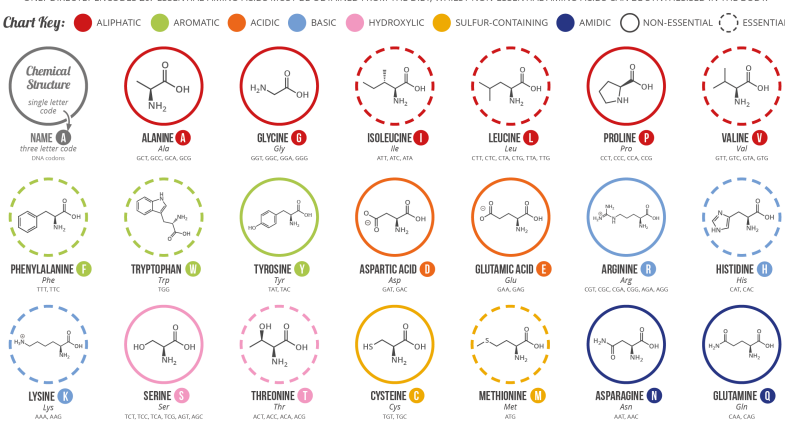
- Ribonucleic acid (mRNA)
  - Result of TRANSCRIPTION
  - Information encoded with the DNA sequence of one or more genes is TRANSCRIBED into a strand of RNA –RNA transcript
    - Single stranded
    - A,G,C,U (T)





# Proteins & Proteomics A GUIDE TO THE TWENTY COMMON AMINO ACIDS

AMINO ACIDS ARE THE BUILDING BLOCKS OF PROTEINS IN LIVING ORGANISMS. THERE ARE OVER 500 AMINO ACIDS FOUND IN NATURE - HOWEVER, THE HUMAN GENETIC CODE ONLY DIRECTLY ENCODES 20. 'ESSENTIAL' AMINO ACIDS MUST BE OBTAINED FROM THE DIET, WHILST NON-ESSENTIAL AMINO ACIDS CAN BE SYNTHESISED IN THE BODY.



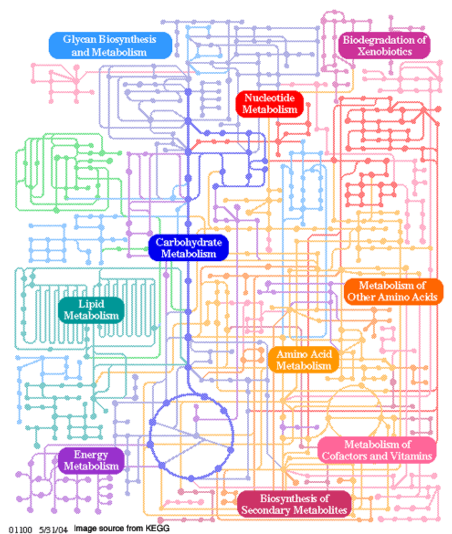
**Note:** This chart only shows those amino acids for which the human genetic code directly codes for. Selenocysteine is often referred to as the 21st amino acid, but is encoded in a special manner. In some cases, distinguishing between asparagine/aspartic acid and glutamine/glutamic acid is difficult. In these cases, the codes asx (B) and glx (Z) are respectively used.





## Metabolites & Metabolomics

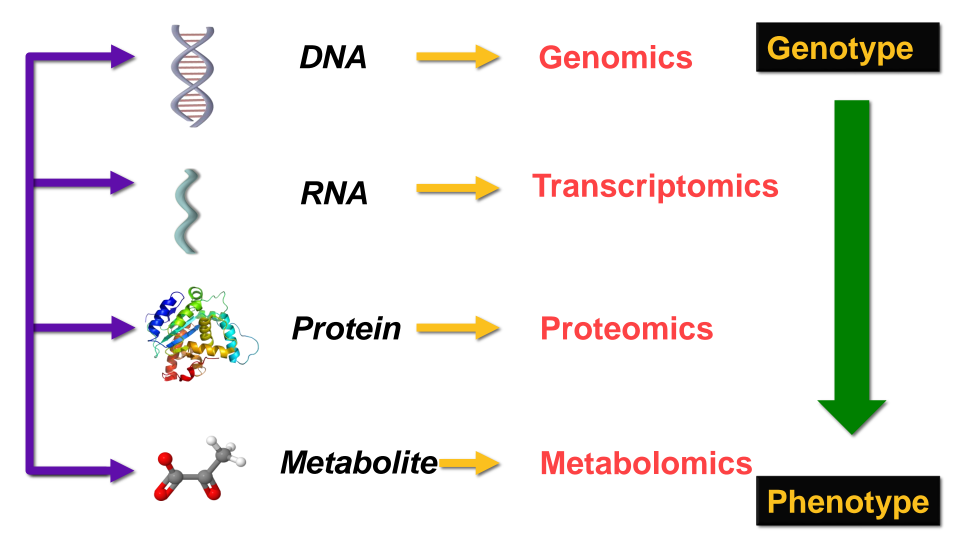
- Low molecular weight produ
- Metabolism is a complex intaction that occur within cells
- Diverse roles
  - Energy metabolism
  - Amino acid metabolism
  - Lipid metabolism
- Primary metabolism –direct development
- Secondary metabolism er
  - Antibiotics, steroids etc



BIOSOCIAL CDT 13/02/2020 01100 5/31/04 Image source from KEGG



## Central dogma of molecular biology



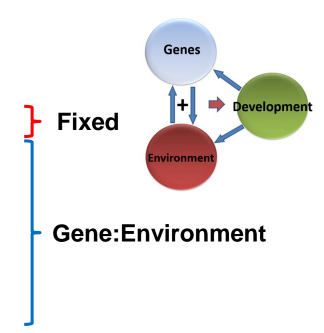


### What is 'Omic Data

The University of Manchester

#### Omic data sets include:

- Genetics (SNPs, CNVs)
- Transcriptomics (Affymetrix, RNAseq)
- Epigenomics (DNA methylation, histone mods)
- ChIPseq
- Metabolomics
- Proteomics
- Phosphoproteomics



All have specific quality control (QC) issues and difficulties in analysis All rely on the use of a false discovery rate correction (FDR) for analysis





When one and one gives more than two: challenges and opportunities of integrative omics

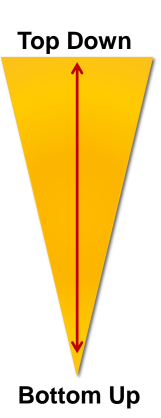
Hyungwon Choi<sup>1</sup>\* and Norman Pavelka<sup>2</sup>\*

Multi-omic analysis is the integration of different omic data sets

# TYPES OF 'OMICS DATA?

## Why 'omics?

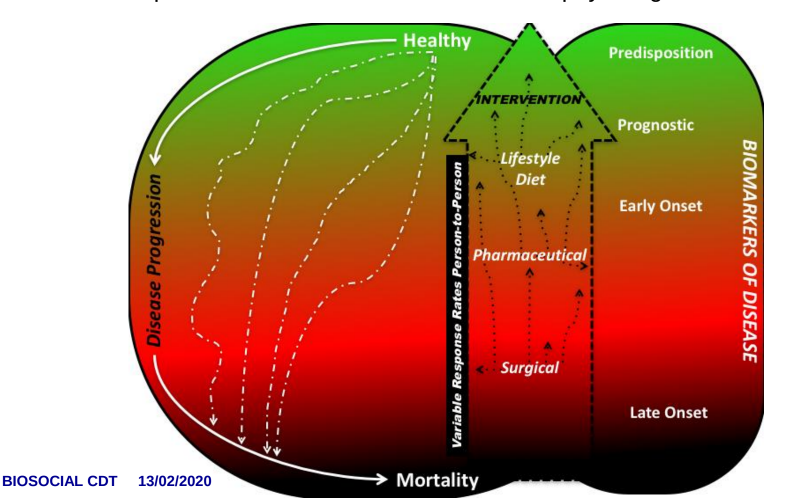
- Adopts an holistic view of all 'molecules' that make up a cell, tissue or organism
- Universal approach/Hypothesis-generating
- No analysis bias
- Many applications
  - 'BIOMARKER' discovery
  - Early detection/population screening
  - Increasing understanding of disease aetiology
  - Drug discovery & toxicity and efficacy screens





## Biomarkers of disease

In medicine, a **biomarker** is a measurable indicator of the severity or presence of some disease state. More generally a **biomarker** is anything that can be used as an indicator of a particular disease state or some other physiological state of an organism.

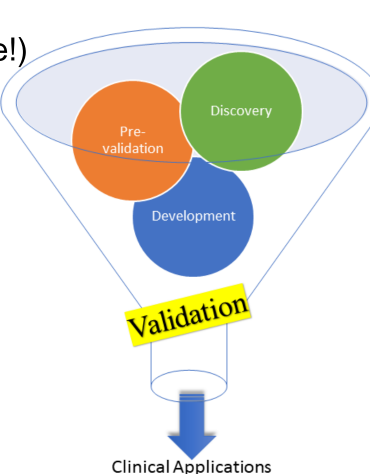


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## Experimental Design

Case vs. Control (an appropriate one!)

- Samples numbers?
  - How many is sufficient?
  - Test cohort + validation cohort?
- Quality of samples
  - Collection/storage/processing
- Confounding factors?
  - Age, gender, environment



## Is this good experimental design?

 Metabolomics investigation of liver failure from plasma samples

**Table 1.** Demographic Information of the Healthy Group and Liver Failure Patient Group Investigated<sup>a</sup>

	healthy group $(n = 23)$	patient group $(n = 24)$
Gender (male/female)	15/8	21/3
HBsAg	Negative	Positive
Age (year)	$27.39 \pm 9.24$	$46.77 \pm 13.35$
ALT (U/L)	< 40	$172.63 \pm 147.49$
TB (µmol/L)	<12	$457.33 \pm 135.48$
PT (s)	< 14	$26.06 \pm 15.14$
MELD score	/	$24.68 \pm 8.38$

<sup>&</sup>lt;sup>a</sup> Abbreviations: ALT, alanine aminotransferase; TB, total bilirubin; PT, prothrombin time; MELD, model for end-stage liver disease. The value is represented as the form of mean  $\pm$  SD.



## Or what about this one?

Molecular Classification of Cancer: Class Discovery and Class Prediction by Gene Expression Monitoring

T. R. Golub<sup>1,2,\*,†</sup>, D. K. Slonim<sup>1,†</sup>, P. Tamayo<sup>1</sup>, C. Huard<sup>1</sup>, M. Gaasenbeek<sup>1</sup>, J. P. Mesirov<sup>1</sup>, H. Coller<sup>1</sup>, M. L. Loh<sup>2</sup>, J. R. Downing...

+ See all authors and affiliations

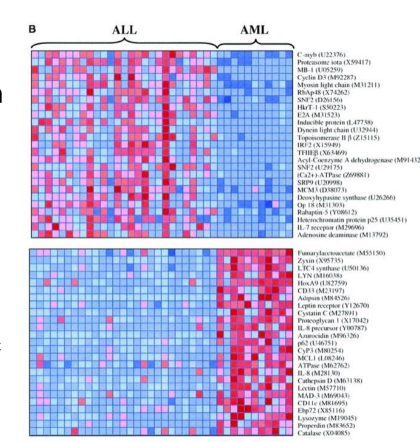
Science 15 Oct 1999: Vol. 286, Issue 5439, pp. 531-537 DOI: 10.1126/science.286.5439.531

Generic approach to cancer classification based on gene expression monitoring by DNA microarrays applied to human acute leukemias

38 Affymetrix microarrays with 6,817 probes

27 from childhood acute lymphoblastic leukemia

11 from adult acute myeloid leukemia



-2.5 -2 -1.5 -1 -0.5 0 0.5 1 1.5 2

Normalized Expression



## Or this?

The University of Manchester

## Novel biomarkers for pre-eclampsia detected using metabolomics and machine learning

Louise C. Kenny<sup>a,\*</sup>, Warwick B. Dunn<sup>b</sup>, David I. Ellis<sup>b</sup>, Jenny Myers<sup>a</sup>, Philip N. Baker<sup>a</sup> and the GOPEC Consortium, and Douglas B. Kell<sup>b,\*</sup>

 Pre-eclampsia - Pregnancy-induced hypertension which may affect mother and foetus

Table 1
Demographic data for patients from whom plasma samples were taken

	Normal outcome $n=87$	Preeclampsia $n = 87$
Age	30 (19–43)	31 (19–41)
Parity	0 (0-2)	0 (0-2)
BMI (weight/height <sup>2</sup> )	25 (19-46)	26 (18-46)
Max (S) BP (mm Hg)	122 (96-147)	162 (138-220)*
Max (D) BP (mm Hg)	80 (60–93)	110 (90-140)*
Delivery gestation	40 + 4	37+0*
(weeks + days)	(34+3  to  42+0)	(26+3  to  41+1)
Birth weight (g)	3420 (2380-4420)	2410 (590-4300)*
IBR (centile)	34 (10-99)	8 (0-99)*

Median (range).

Pre-eclampsia vs normal outcome.

p < 0.0001.

## Genomics

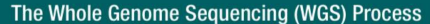
- Genome = total DNA of a cell or an organism
- Human genome = 3.2 billion bases and estimated > 30,000 protein coding genes
- Seeking mutations or alterations that may contribute towards a certain disease!
  - *i.e.* the genes BRCA1 and BRCA2 cause 60% of all cases of hereditary breast and ovarian cancers
  - BUT not a single mutation- there are >800 different mutations in BRCA1 alone

#### **TOOLS: Whole Genome Sequencing**

https://theanalyticalscientist.com/fields-applications/the-tools-behind-genomics



## Sequencing



WGS is a laboratory procedure that determines the order of bases in the genome of an organism in one process. WGS provides a very precise DNA fingerprint that can help link cases to one another allowing an outbreak to be detected and solved sooner.

#### **Bacterial Culture**



Scientists take bacterial cells from an agar plate and treat them with chemicals that break them open, releasing the DNA. The DNA is then purified.



Scientists make many copies of each DNA fragment using a process called polymerase chain reaction (PCR). The pool of fragments generated in a PCR machine is called a "DNA library."



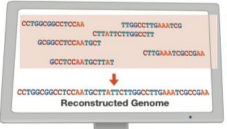


DNA is cut into short fragments of known length, either by using enzymes "molecular scissors" or mechanical disruption.



4. DNA Library Sequencing

sequencer. The combination of nucleotides (A, T, C, and G) making up each individual fragment of DNA is determined, and each result is called a "DNA read."



ACTGRACTGACTG CTGACTGACTGACT

CTOGNARCTCCARG



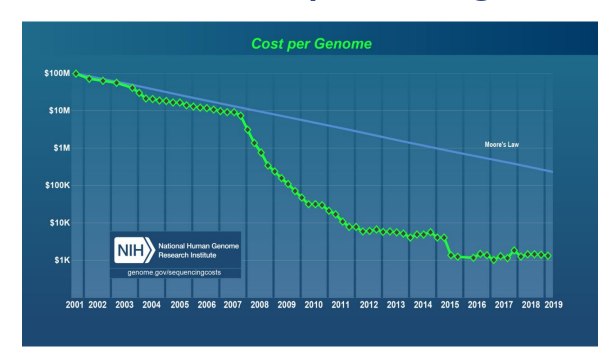
The sequencer produces millions of DNA reads and specialized computer programs are used to put them together in the correct order like pieces of a jigsaw puzzle. When completed, the genome sequence containing millions of nucleotides (in one or a few large pieces) is ready for further analysis.

DNA

Reads



## Cost of Sequencing





#### PacBio Sequencer

£300K

Single-Molecule, Real-Time (SMRT) technology

BIOSOCIAL CDT 13/02/2020 WWW.pacb.com 2

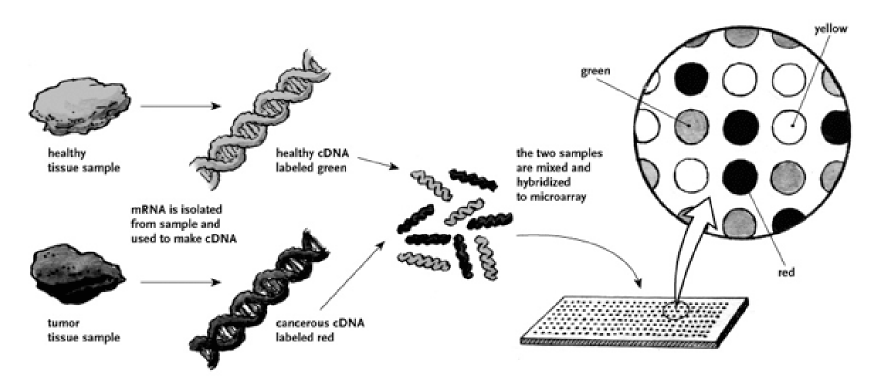
## **Transcriptomics**

- Measuring the transcriptome
  - All the mRNA that is transcribed at a given point in a given cell or organism
- Provides direct knowledge of gene regulation and protein content information
  - Which genes are actively expressed
- Began in 1990s and is now a widespread discipline
  - Many technological advances
  - Now a routine, 'simple' process

TOOLS: Microarrays & RNA-Seq

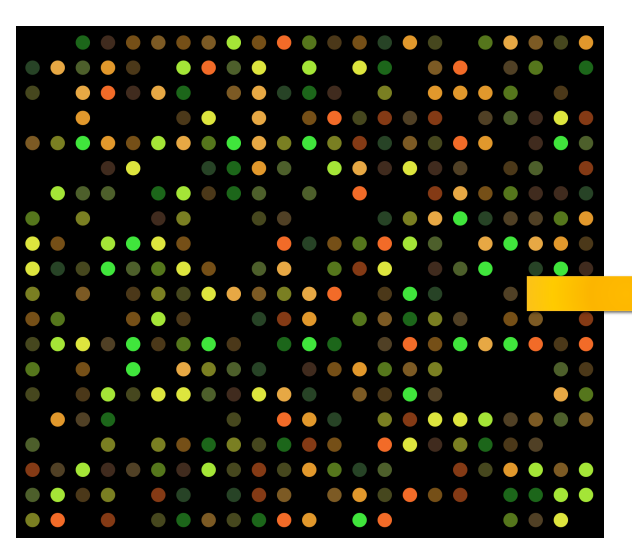


## **DNA Microarrays**



A microarray is an orderly arrangement of rows and columns on a surface like a glass slide. Each of the spots on an array contains single-stranded DNA molecules that correspond to a single gene. An array can contain a few, or thousands, of genes.

## **DNA Microarrays**



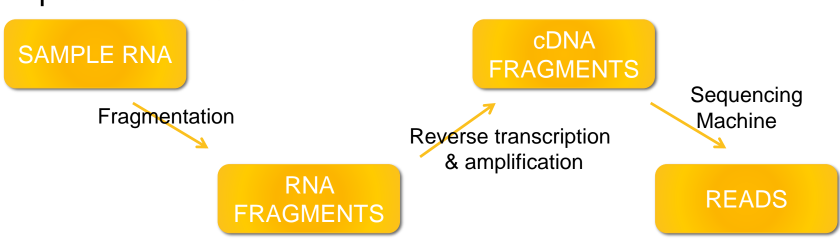
Converted to numerical values for interpretation

Limitation: Results limited to what probes you have on your chip



## RNA Seq

- High throughput sequencing with computat
- No reference sequence
- Large dynamic range
- Sequences every RNA molecule and profiles the expression of a particular gene by counting the number of times its transcript has been sequenced
- Expression levels!





## **Proteomics**

- Investigating the entire complement of proteins within a cell, tissue or organism.
- >100,000 proteins
- Large dynamic range
- Some questions
  - When & where proteins are expressed
  - The involvement of proteins with particular phenotypes
  - How proteins are modified or how they interact with each other

Blood, urine, tissues



## **Proteomic Tools**

Mass spectrometry







Proteins for top-down analyses come from a variety of sources.



#### EXTRACTION OF PROTEINS

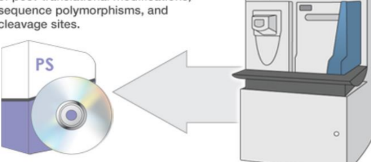
Proteins are extracted and denatured.



Proteins are separated, most often by molecular weight, to reduce sample complexity and ensure maximal identification of intact proteins.

#### AUTOMATED DATA ANALYSIS

Intact proteins are identified in an automated fashion using ProSightPC software, including characterization of post-translational modifications, sequence polymorphisms, and cleavage sites.



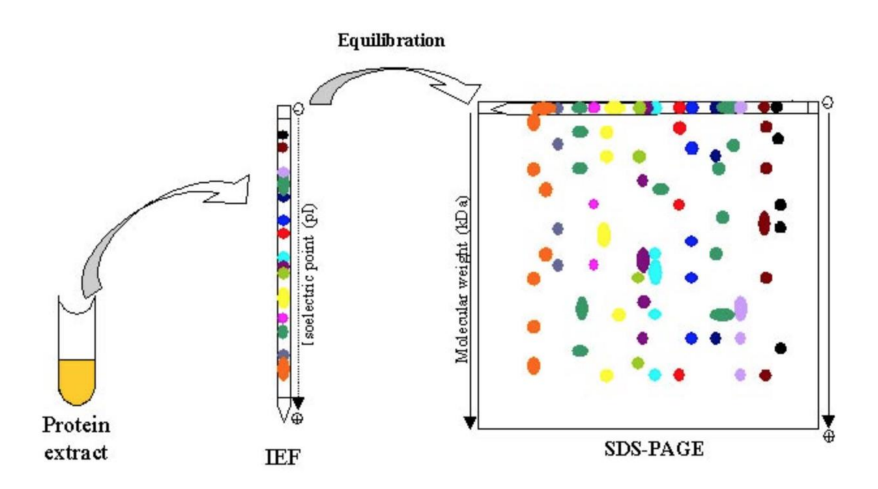
#### ANALYSIS BY LC-MS/MS

Intact proteins are analyzed by LC-MS/MS on Orbitrap-based mass spectrometers.





## **Proteomic Tools**

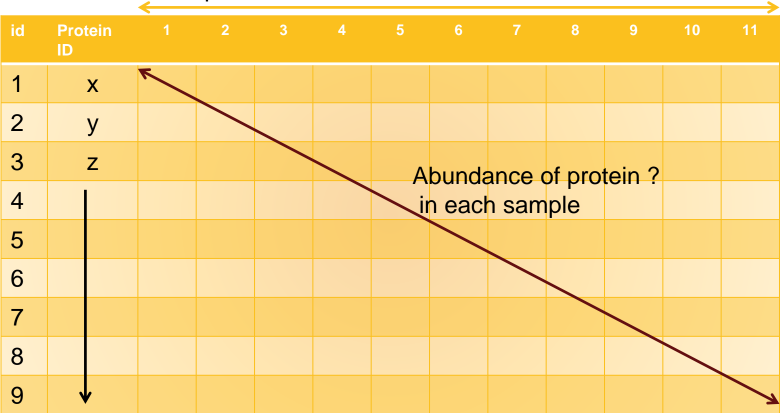


100's

**Detected Proteins -**

## **Proteomics Data**



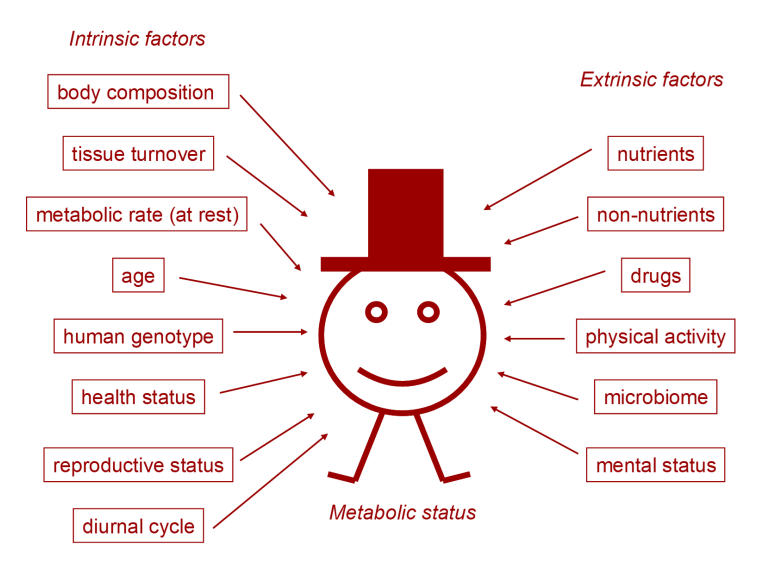


Do statistical analysis to decide which features change between classes

## **Metabolomics**

- Investigating the entire complement of low molecular weight molecules in an sample of interest.
- ~ 5000 metabolites in a biological sample from a human
  - GREATLY impacted by environment!
  - GREATLY impacted by time!
- Experimental design & sample collection is VERY important!





Goodacre, R. (2007) Metabolomics of a superorganism. *Journal of Nutrition* 137, 259S-266S.



## **Biological Matrices**

## Primary Sources

## Secondary Sources

## **Additional Sources**

Serum

Plasma

**CSF** 

BAL

Saliva

Saliva

Semen Urine

Faeces

Tissue Biopsies

**Brain** 

Nerve

Lung

**Pancreas** 

Liver

Heart

Gut

Skin

**Animal Models** 

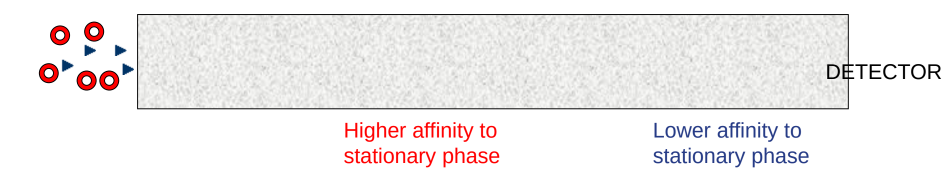
Mammalian Cell Culture

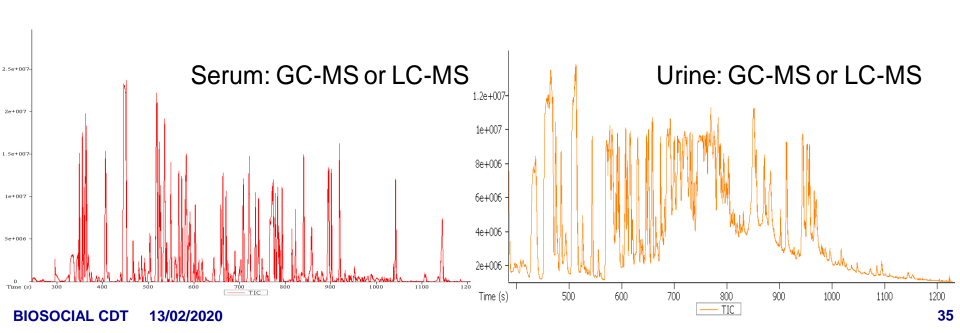
IVF culture medium

Sweat Breath



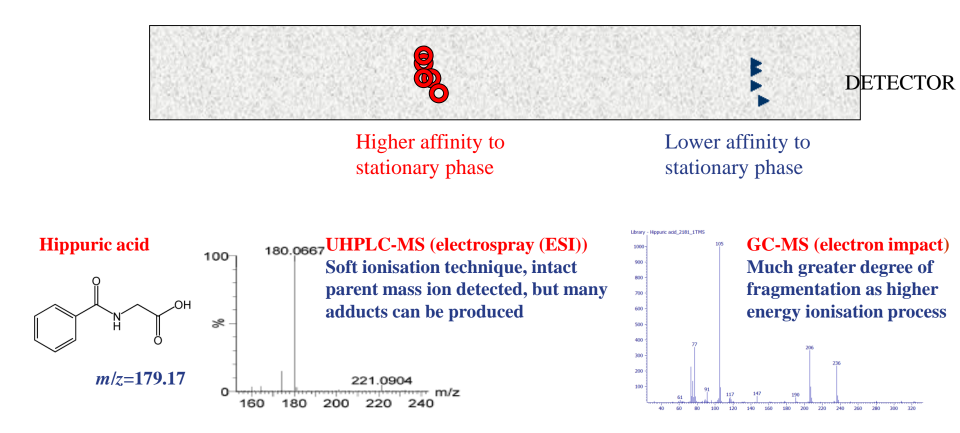
## Chromatography linked to Mass Spectrometry







## Chromatography linked to Mass Spectrometry



Matching of the chromatographic retention time and fragmentation mass spectra between a sample analyte and a reference standard is required for definitive id. We have *ca.* 1600 analytes in our GC-MS library

Sumner L.W. et al. (2007) Metabolomics 3, 211-221

1000's

Detected Features-

## **Metabolomics Data**

Sample Names – often 100's with different

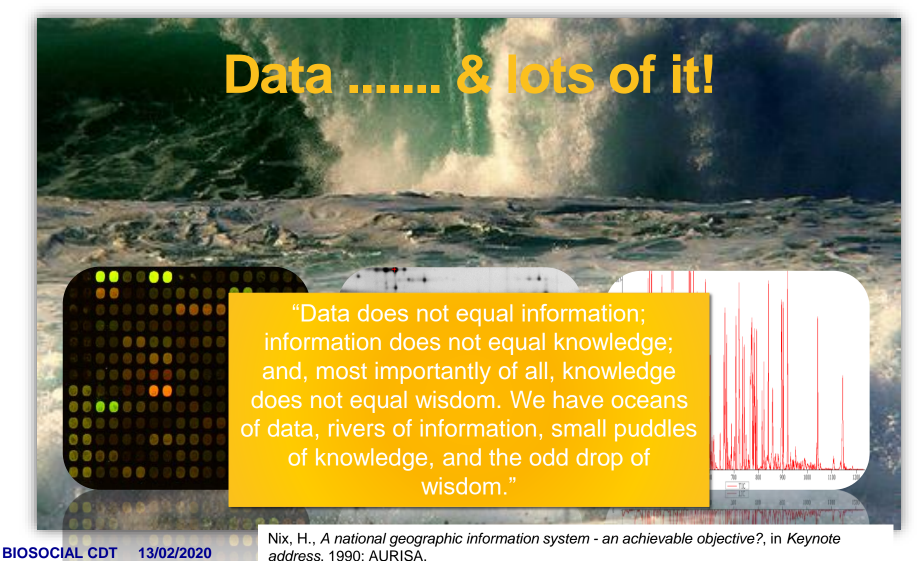


Use RT & m/z to identify the metabolite – MAJOR CHALLENGE!

Do statistical analysis to decide which features change between classes



## Common Theme



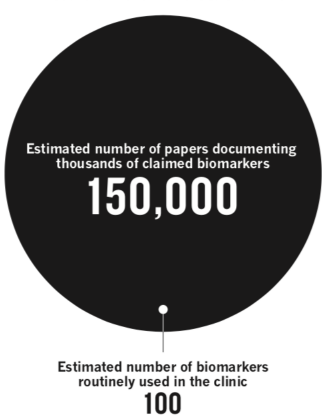
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# The challenge of biomarkers?

Poste, G., Bring on the biomarkers. Nature, 2011.
 469(7329): p. 156-157.

Few of the numerous biomarkers so far discovered have made it to the clinic.

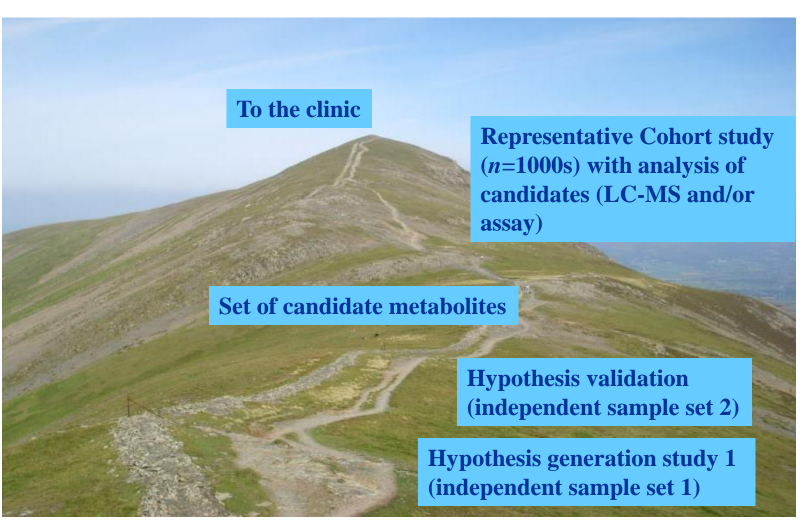
Multi-Centre, large cohort studies





## Biomarker Discover: From lab to bedside







# SOME EXAMPLES.....

## Personalised/Precision Medicine

- Is this the future of healthcare?
  - 2015 State of the Union address President Obama announced that he was launching the Precision Medicine Initiative
- Assess the genotype (SNPs) and phenotype (metabolome) of a patient before they undergo any treatment
  - Population monitoring & data collation
  - Seeking cures & preventative screening
- Offering a well-designed screening program at a reasonable cost may not always be possible due to the numerous associated challenges
  - monetary limitations (labour and consumable costs) as well as ethical, legal and social considerations for an opt-in test

# (Metabol)'omics for the masses?

- Wearable technologies smartphones, smart-watches, health bands, necklaces, glucose monitoring contact lenses
  - Innovations for collecting personal information
- mPower mobile Parkinson's Disease study that attempts to research the occurrence, presentation and management of PD symptoms via survey telemetry data using a smartphone app
  - Bot, B.M., et al., The mPower study, Parkinson disease mobile data collected using ResearchKit. Scientific Data, 2016. 3: p. 160011.
- Smart-phone app to monitor the association between pain and the weather for people suffering from rheumatoid arthritis
  - Reade, S., et al., Cloudy with a Chance of Pain: Engagement and Subsequent Attrition of Daily Data Entry in a Smartphone Pilot Study Tracking Weather, Disease Severity, and Physical Activity in Patients With Rheumatoid Arthritis. JMIR Mhealth Uhealth, 2017. 5(3): p. e37.
  - Dixon, W.G., et al. How the weather affects the pain of citizen scientists using a smartphone app. Npj Digital Medicine (2019) 2:105

CLOUDY



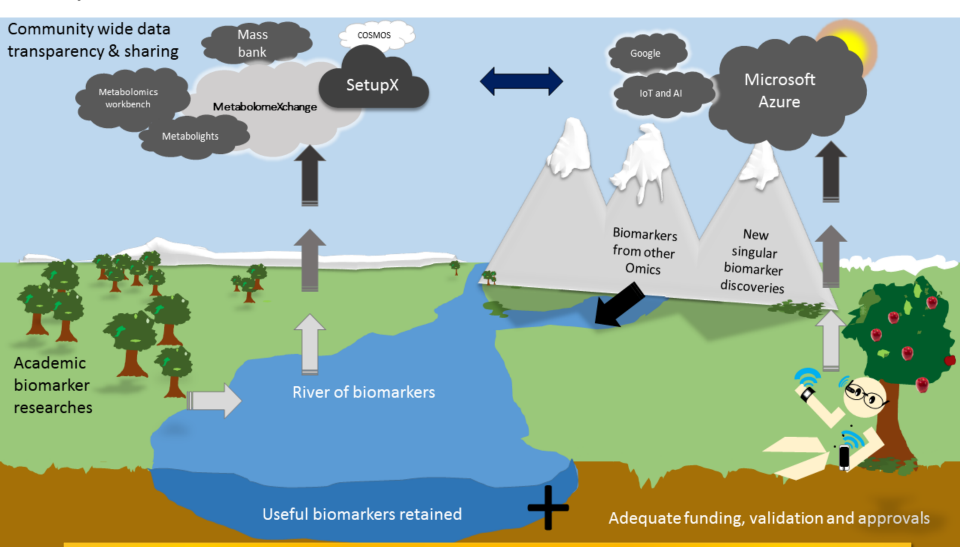




Patients with chronic pain commonly believe their pain is related to the weather. Scientific evidence to support their beliefs is inconclusive, in part due to difficulties in getting a large dataset of patients frequently recording their pain symptoms during a variety of weather conditions. Smartphones allow the opportunity to collect data to overcome these difficulties. Our study *Cloudy with a Chance of Pain* analysed daily data from 2658 patients collected over a 15-month period. The analysis demonstrated significant yet modest relationships between pain and relative humidity, pressure and wind speed, with correlations remaining even when accounting for mood and physical activity. This research highlights how citizen-science experiments can collect large datasets on real-world populations to address long-standing health questions. These results will act as a starting point for a future system for patients to better manage their health through pain forecasts.

## MANCHESTER 1824

The University of Manchester



The future cycle of metabolomics precision medicine-based research and healthcare where academia, industrial partners, corporate data analytics work with patients' wearable data collection devices to provide health monitoring solutions.



## TWINS UK



FOR TWINS ☑



# Twin research for a healthy future

Researching the link between our genes, the environment, and common diseases



<u> 14,274</u> Twins



**76** Studies



800+ Publications



**59** Researchers

### Looking to collaborate?

We aim to facilitate and encourage the sharing of TwinsUK data and samples with the world's scientific community to promote and contribute to scientific research and generate new knowledge. Find out more by visiting our data access pages below.

Collaborate

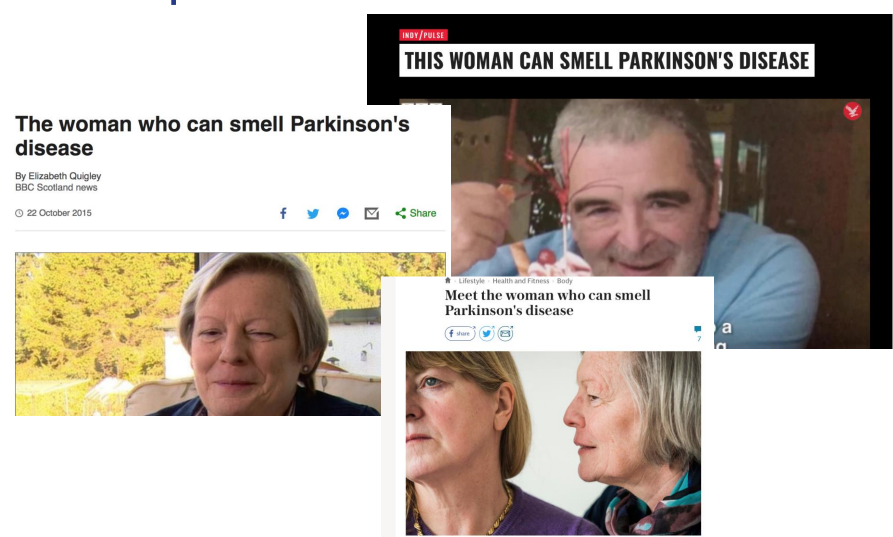
L Vision

### **Research Areas** TWINS UK **Omics** □ Epigenomics Comprehensive study □ Expression-omics □ Genetics □ Glycomics Unique design with internal controls Well documented & controlled **Disease & Ageing** ∟ Bone Wealth of scientific publications □ Brain □ Cardiovascular □ Diet ⊢ Hearing Pain ∟ Skin

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# The super smeller



BIOSOCIAL CDT 13/02/2020 48

Joy Milne, right, gives the author a sniff CREDIT: CHRIS WATT



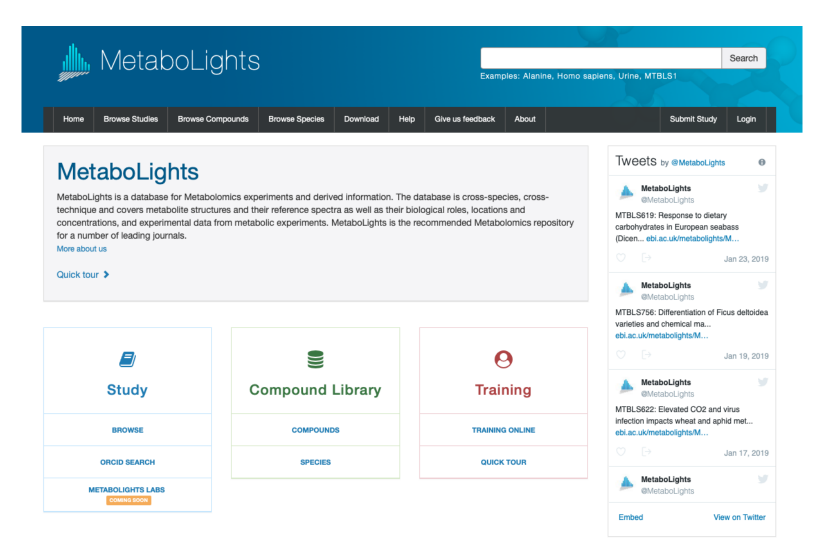
## Biomarker detection



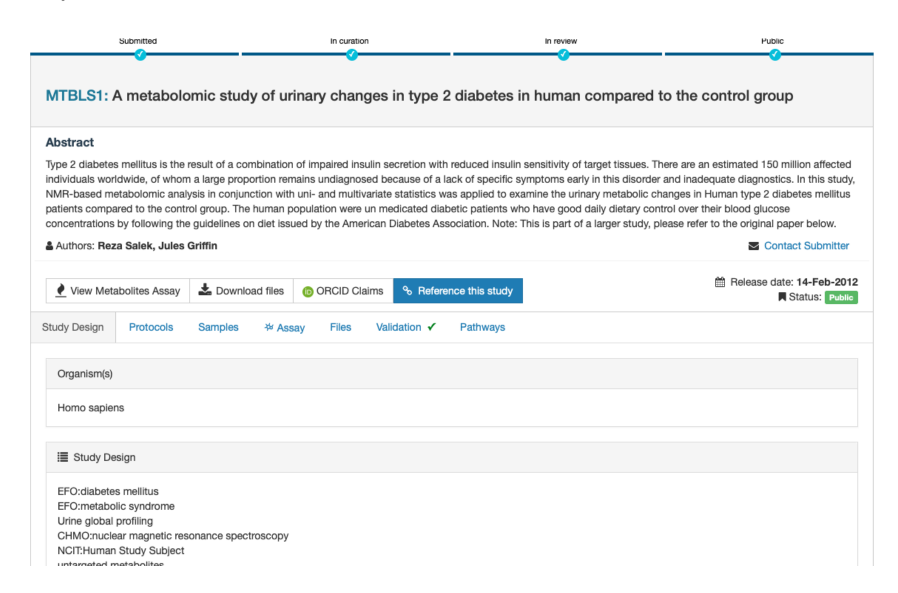
Gaining distance on biomarker discovery.....



# Data Repositories







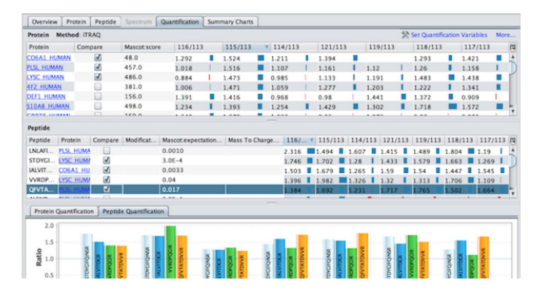




PRIDE > Archive

## PRIDE Archive - proteomics data repository

The PRIDE PRoteomics IDEntifications (PRIDE) database is a centralized, standards compliant, public data repository for proteomics data, including protein and peptide identifications, post-translational modifications and supporting spectral evidence. PRIDE is a core member in the ProteomeXchange (PX) consortium, which provides a single point for submitting mass spectrometry based proteomics data to public-domain repositories. Datasets are submitted to PRIDE via ProteomeXchange and are handled by expert biocurators.



#### Datasets

- 8159 projects
- 76426 assays



JAN @CellCellPress

"Retrovirus-like Gag
Protein Arc1 Binds RNA
and Traffics across Synaptic
Boutons" link Dataset

#PXD008136

JAN (2/2) ... corresponding dataset #PXD004736 available proteomexchange @pride\_ebi

proteomexchange @pride\_ebi link

JAN

JAN (1/2) @naturemethods

"Biotinylation by antibody recognition—a method for proximity labeling" link





#### PEPTIDEATLAS HOME

Seattle Proteome Center

#### PEPTIDEATLAS: Overview Contacts Data Contributors

**Publications** Software

Database Schema Feedback FAQ

#### ATLAS DATA:

Data Repository Human Plasma (Farrah, et al.) **HPPP Data Central** PeptideAtlas Builds Search Database

Contribute Data Genome Browser Setup

RELATED: **SRMAtlas** 

**PASSEL** Phosphopep Uninen



Search PeptideAtlas:

GO

**Expanded Search** 

PeptideAtlas is a multi-organism, publicly accessible compendium of peptides identified in a large set of tandem mass spectrometry proteomics experiments. More...



**PeptideAtlas** Chromosome **Explorer** (Human only)

### **SRMAtlas**

**SRMAtlas** interface for selection of best available **SRM transitions** 



**PeptideAtlas** Raw Data Repository



(PASSEL)



Submit

your raw data to PeptideAtlas



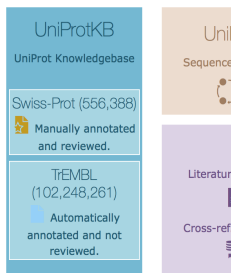
PeptideAtlas and the Chromosome-Centric **Human Proteome Project** 

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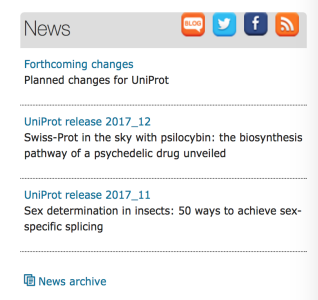




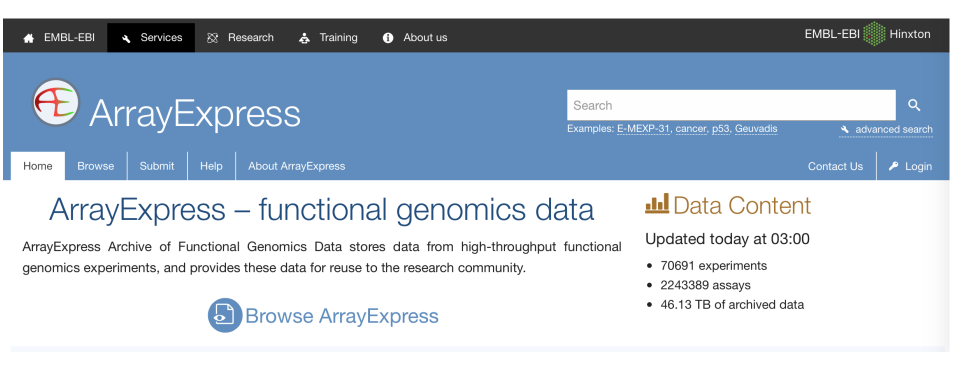
The mission of UniProt is to provide the scientific community with a comprehensive, high-quality and freely accessible resource of protein sequence and functional information.





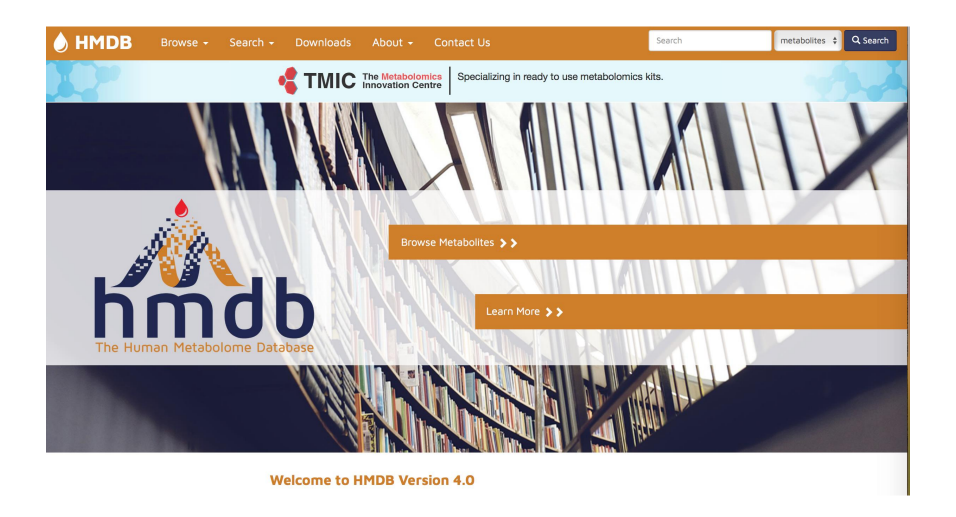






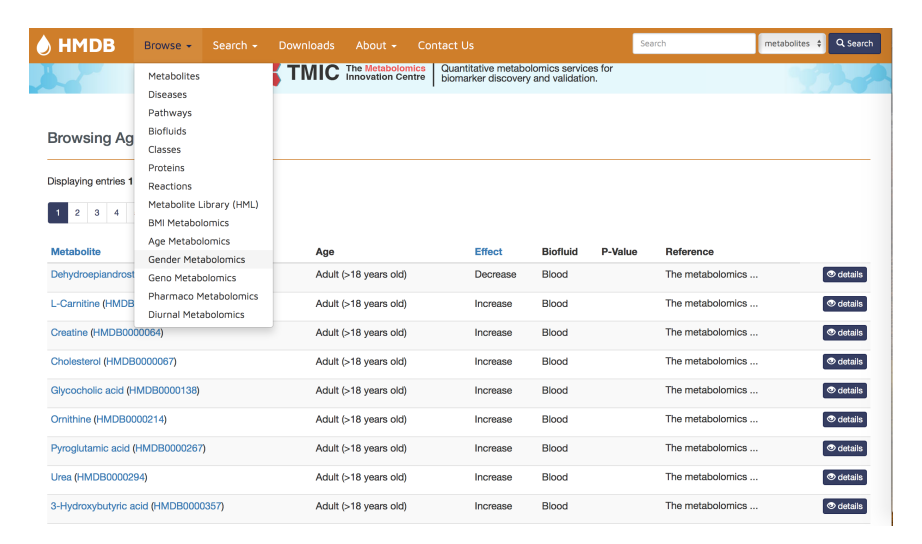


## Human Metabolome Database





# **Search Parameters**





## Standardisation

- MSI formed in 2005 to unify and to engage with the growing metabolomics community so that experiments can be reproduced by others and are based on solid sample collection, analysis and data processing. — Complete Transparency!
- Working group now working on how to perform experimental design better.
- Pre-requisite for publication in Metabolomics

Metabolomics (2007) 3:175–178
DOI 10.1007/s11306-007-0070-6

BRIEF REPORT

The metabolomics standards initiative (MSI)

Oliver Fiehn · Don Robertson · Jules Griffin ·
Mariet van der Werf · Basil Nikolau · Norman Morrison ·
Lloyd W. Sumner · Roy Goodacre · Nigel W. Hardy ·
Chris Taylor · Jennifer Fostel · Bruce Kristal ·
Rima Kaddurah-Daouk · Pedro Mendes ·

Ben van Ommen · John C. Lindon · Susanna-Assunta Sansone

Metabolite Identification MSI Level 1 – Definitive MSI Level 2 - Putative

# **NETWORK ANALYSIS**



## **Individual Omic Data Set Analysis**

The University of Manchester

	varID	Gene Symbol	q-Value	Fold change
1	203820_s_at	IGF2BP3	1.19574e-16	3.03628
2	203819_s_at	IGF2BP3	9.37733e-11	2.62628
3	240143_at		2.07997e-08	1.76953
4	206659_at		3.21346e-07	1.32887
5	228988_at	ZNF711	1.25012e-05	1.56114
6	201417_at	SOX4	1.33199e-05	1.71911
7	207996_s_at	C18orf1	1.33199e-05	1.38872
8	203038_at	PTPRK	1.33317e-05	1.65887
9	201310_s_at	C5orf13	1.38711e-05	1.42943
10	222344_at		1.67269e-05	1.40778
11	213808_at	ADAM23	1.88826e-05	1.61604
12	214744_s_at	RPL23	2.49258e-05	1.29967

Age group 2 Vs Rest, ANOVA, q<0.1 = 1524 probe-sets

## Lists and Cutoffs!!

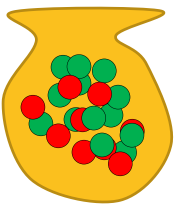
What does a p-value threshold mean?

What does a Fold change cut off mean?



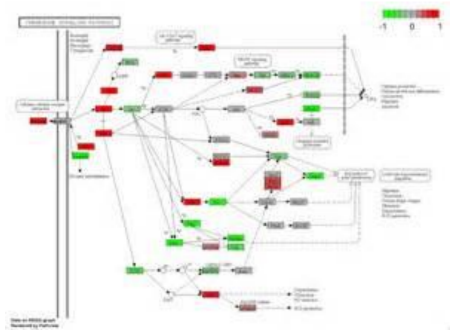
# Fisher's Exact Test / Hypergeometric Test





Pick 5 marbles

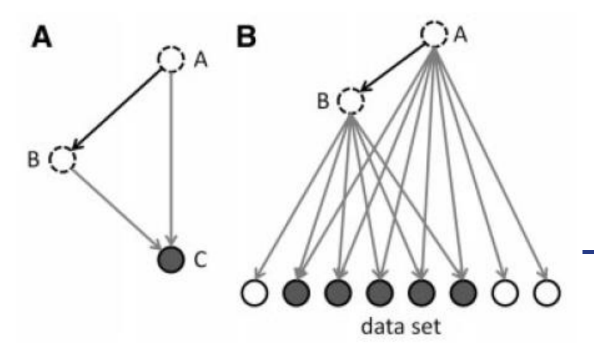
P(2 are red)





## **Upstream Regulator Analysis**

The University of Manchester



Enrichment of 'causal transitive triangles'

### **Literature Driven analysis**



Interactive pathway analysis of complex 'omics data

BIOSOCIAL CDT

13/02/2020

#### **BIOINFORMATICS**

ORIGINAL PAPER

2014, pages 1–8 doi:10.1093/bioinformatics/btt703

Systems biology

Advance Access publication December 13, 2013

#### Causal analysis approaches in Ingenuity Pathway Analysis

Andreas Krämer<sup>1,\*</sup>, Jeff Green<sup>1</sup>, Jack Pollard, Jr<sup>2</sup> and Stuart Tugendreich<sup>1</sup>

<sup>1</sup>Ingenuity Systems, 1700 Seaport Boulevard, Redwood City, CA and <sup>2</sup>Translational and Experimental Medicine—Bioinformatics, Sanofi-Aventis, 270 Albany Street, Cambridge, MA, USA

Associate Editor: Jonathan D, Wren



## What is **Network Biology**

The University of Manchester

### Biological networks are:

- "Scale free"
- Resistant to random error
- Exhibit "small world" properties

*Nature* **393**, 440-442 (4 June 1998) | doi:10.1038/30918; 1998

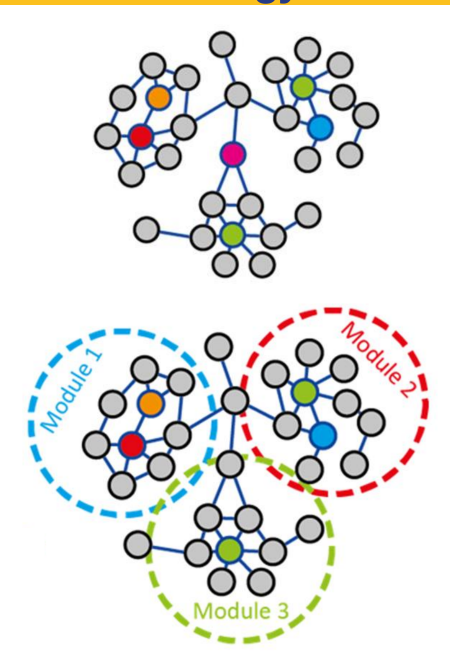
Collective dynamics of 'small-world' networks

Duncan J. Watts<sup>2</sup> & Steven H. Strogatz<sup>1</sup>

## $L \propto \log N$

typical distance L between two randomly chosen nodes (the number of steps required) grows proportionally to the logarithm of the number of nodes N in the network

small-world network - most nodes are not neighbours of one another, but most nodes can be reached from every other by a small number of steps.



< Previous Article

Volume 159, Issue 5, p1212-1226, 20 November 2014

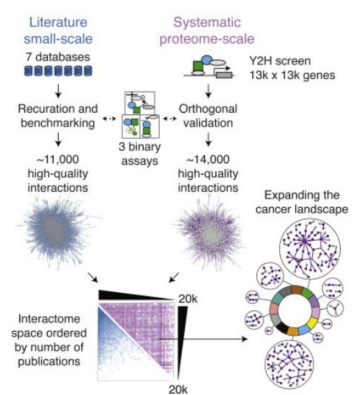
Resource Switch to Standard View

### A Proteome-Scale Map of the Human Interactome Network

Thomas Rolland<sup>19</sup>, Murat Taşan<sup>19</sup>, Benoit Charloteaux<sup>19</sup>, Samuel J. Pevzner<sup>19</sup>, Quan Zhong<sup>19</sup>, Nidhi Sahni<sup>19</sup>, Song Yi<sup>19</sup>, Irma Lemmens, Celia Fontanillo, Roberto Mosca, Atanas Kamburov, Susan D. Ghiassian, Xinping Yang, Lila Ghamsari, Dawit Balcha, Bridget E. Begg, Pascal Braun, Marc Brehme, Martin P. Broly, Anne-Ruxandra Carvunis, Dan Convery-Zupan, Roser Corominas, Jasmin Coulombe-Huntington, Elizabeth Dann, Matija Dreze, Amélie Dricot, Changyu Fan, Eric Franzosa, Fana Gebreab, Bryan J. Gutierrez, Madeleine F. Hardy, Mike Jin, Shuli Kang, Ruth Kiros, Guan Ning Lin, Katja Luck, Andrew MacWilliams, Jörg Menche, Ryan R. Murray, Alexandre Palagi, Matthew M. Poulin, Xavier Rambout, John Rasla, Patrick Reichert, Viviana Romero, Elien Ruyssinck, Julie M. Sahalie, Annemarie Scholz, Akash A. Shah, Amitabh Sharma, Yun Shen, Kerstin Spirohn, Stanley Tam, Alexander O. Tejeda, Shelly A. Trigg, Jean-Claude Twizere, Kerwin Vega, Jennifer Walsh, Michael E. Cusick, Yu Xia, Albert-László Barabási,

Lilia M. Iakoucheva, Patrick Aloy, Javier De Las Rivas, Jan Tavernier, Michael A. Calderwood<sup>20</sup>, David E. Hill<sup>20</sup>, Tong Hao<sup>20</sup>, Frederick P. Roth — Marc Vidal — Ware Vidal — Marc Vidal — Ware Vidal — Marc Vidal — Ware Vidal

DOI: http://dx.doi.org/10.1016/j.cell.2014.10.050



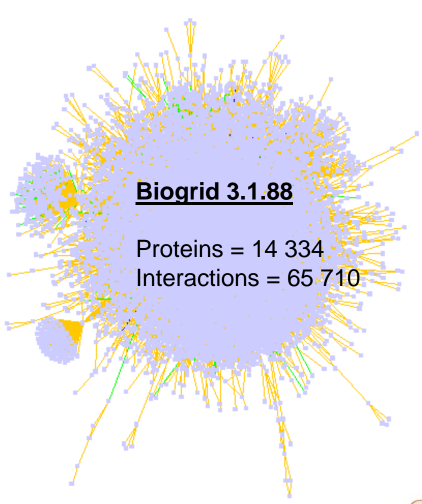
<sup>19</sup> Co-first author

<sup>20</sup> Co-senior author



## **Human Interactome.**

The University of Manchester



### **Biogrid 3.2.105**

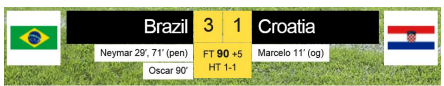
Proteins = 18 107 Interactions = 217 927

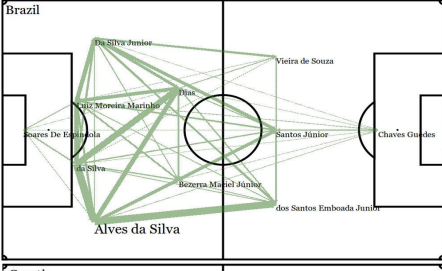


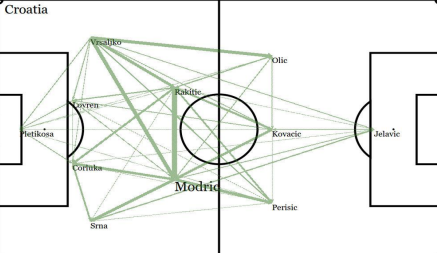


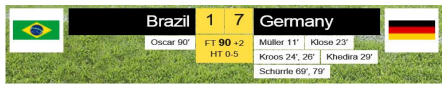
## MANCHESTER What is a network model FIFA World Cup 2014

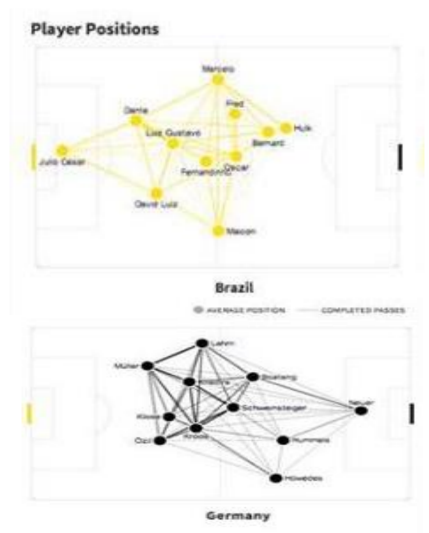
The University of Manchester









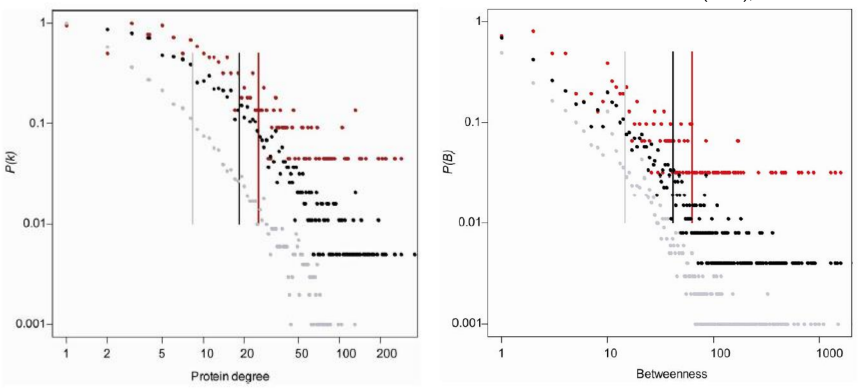




### Network Topology Is Associated with Essential Function

The University of Manchester





Red = Cancer Genes, Black = Essential Genes, Grey = Control Genes

Sun and Zhao BMC Genomics 2010, 11(Suppl 3):S5 http://www.biomedcentral.com/1471-2164/11/S3/S5

RESEARCH



Open Access

A comparative study of cancer proteins in the human protein-protein interaction network

Jingchun Sun<sup>1,2</sup>, Zhongming Zhao<sup>1,2,3\*</sup>

Chavali et al. BMC Systems Biology 2010, 4:78 http://www.biomedcentral.com/1752-0509/4/78



#### RESEARCH ARTICLE

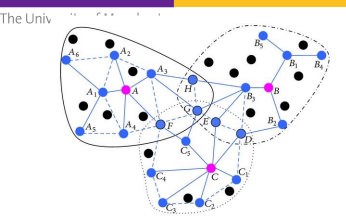
**Open Access** 

Network properties of human disease genes with pleiotropic effects

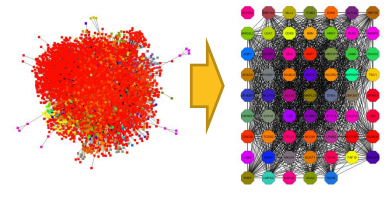
Sreenivas Chavali\*+, Fredrik Barrenas+, Kartiek Kanduri and Mikael Benson



### **Network Clusters are associated with hierarchy of biological function**



"overlapping modules"



"Overlapping Community Clustering

#### Hierarchy of functions and pathways ranked by centrality of cluster

Cluster	Hierarchy of Functions and Pathways	Core nodes of cluster
SUMO <sub>2</sub>	Cellular responses to stress, Apoptosis, Circadian clock, DNArepair, Metabolism of Proteins (q<1x10 <sup>-3</sup> ). G1/S DNA damage checkpoint (q<6.2x10 <sup>-8</sup> )	SUMO2, UBC, MDM2, TP53, EEF1A1, PRKDC, RPS4X, SRRM2, RPS13, RPS20
SKP1	Cell Cycle, Circadian Clock (q<1x10 <sup>-5</sup> ). Wnt & Prolactin signalling (q<1.0x10 <sup>-5</sup> ).	<b>SKP1</b> , BTRC, CUL1,GSK3B, CTNNB1, SKP2, NFKBIA, CLSPN, FBXW11, FBXO6
GSK3B	Apoptosis, Signal transduction (q<0.01). Wnt, PI3/AKT signalling (q<0.01).	GSK3B, AXIN1, APP, AKT1, MAPT, CTNNB1, ELAVL1, KIF5B, AXIN2, HIPK2
NCOA1	Transcriptional regulation of white adipocyte differentiation, Regulation of lipid metabolism by PPARalpha, Circadian Clock, Mitochondrial biogenesis (q<0.01).	NCOA1, NCOA6, ESR1, PPARG, MLL3, RXRA, ESR2, MLL4, NCOR2, VDR
EZH2	Cellular Senescence, Epigenetic regulation of gene expression (q<1.0x10 <sup>-4</sup> )	<b>EZH2,</b> EED, SUZ12, EZH1, JARID2, SON, SRSF7, NRF1, FBL, HDAC1
TCEB2	Cellular response to hypoxia (q<1x10 <sup>-5</sup> ). TGF-beta Signalling (q<0.05)	TCEB2, VHL, CUL5, TCEB1, CUL2, TCEB3, ASB9, STK16, NEDD8, COPS6
NCOR2	Transcriptional regulation of white adipocyte differentiation, Regulation of lipid metabolism by PPARalpha, Circadian Clock, Mitochondrial biogenesis (q<0.01).	NCOR2, NCOA6, HDAC3, BCL6, RARA, AR, ANKRD11, THRB, HDAC1, KDM5B
DIABLO	Immune system, Apoptosis (q<0.05) Toll-Like Receptors Cascades (q<1.0x10 <sup>-4</sup> )	<b>DIABLO</b> , XIAP, BIRC2, BIRC6, UBE2D4, BIRC3, BIRC7, TRAF2, BIRC5, ELAVL1
NBN	DNA Repair, Cell Cycle (q<1.0x10 <sup>-4</sup> ) G2/M DNA damage checkpoint (q<6.7x10 <sup>-4</sup> )	NBN, MRE11A, MDC1, RAD50, BRCA1, H2AFX, FANCD2, ATM, TP53BP1, ATR
rsci	Cellular response to stress (q<0.01). Insulin, IGF1, mTOR, PI3K signalling (q<1.0x10 <sup>-5</sup> ).	<b>TSC1</b> , RHEB, TSC2, YWHAE, MTOR, YWHAB, RAF1, RPTOR, BECN1, MAST3

Bioinformatics (2012) 28: 2202-2204

Data and text mining

ModuLand plug-in for Cytoscape: determination of hierarchical layers of overlapping network modules and community centrality

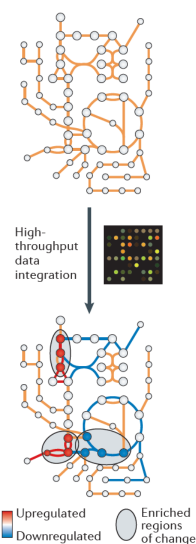
Máté Szalay-Bekő<sup>1,‡</sup> and Robin Palotai<sup>1,‡</sup>, Balázs Szappanos<sup>2</sup>, István A. Kovács<sup>1,3</sup>, Balázs Papp<sup>2</sup> and Péter Csermely<sup>1,\*</sup>



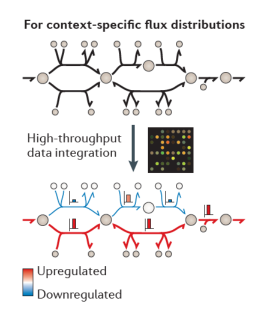
### **Constraint Based Modelling**

The University of Manchester

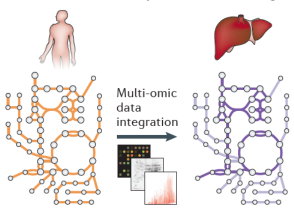
a Topological enrichment



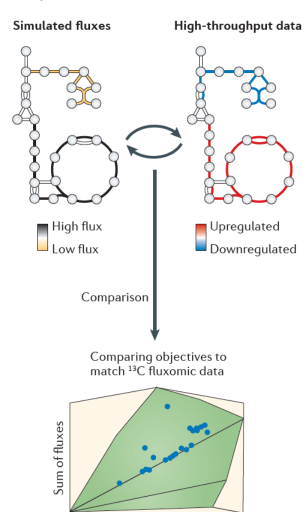
**b** Constraining the solution space



For cell- and tissue-specific model building



c Comparison



Biomass

yield

ATP

yield

NATURE REVIEWS | GENETICS VOLUME 15 | FEBRUARY 2014 | 107

### Why are methods of prioritising Omic data important?

The University of Manchester

**Answer:** Lists don't necessarily deliver the solution!

Haem oxygenase is synthetically lethal with the tumour suppressor fumarate hydratase

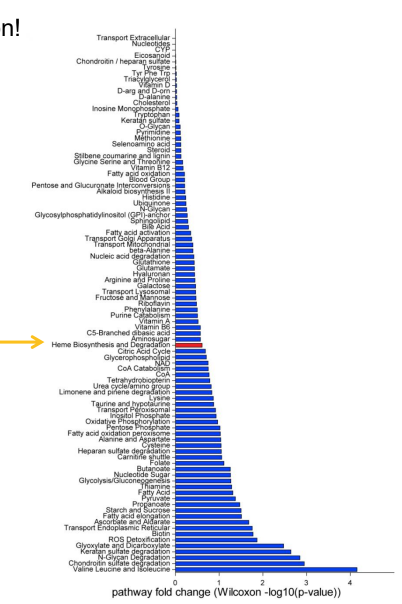
Christian Frezza, Liang Zheng, Ori Folger, Kartik N. Rajagopalan, Elaine D. MacKenzie, Livnat Jerby, Massimo Micaroni, Barbara Chaneton, Julie Adam, Ann Hedley, Gabriela Kalna, Ian P. M. Tomlinson, Patrick J. Pollard, Dave G. Watson, Ralph J. Deberardinis, Tomer Shlomi, Eytan Ruppin & Eyal Gottlieb

Affiliations | Contributions | Corresponding author

Nature 477, 225–228 (08 September 2011) | doi:10.1038/nature10363 Received 13 July 2010 | Accepted 11 July 2011 | Published online 17 August 2011

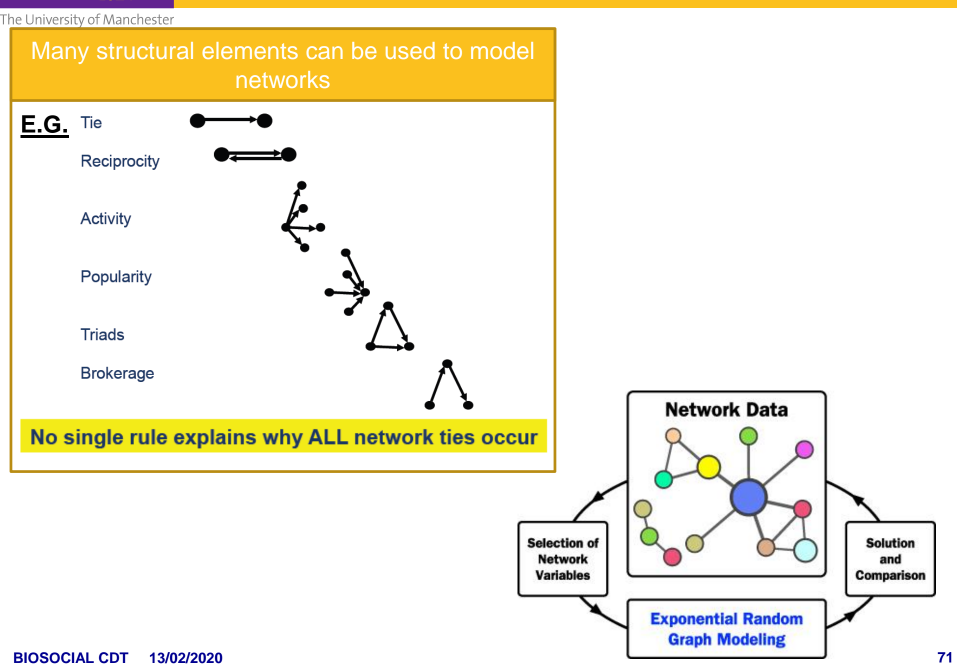
Gene set analysis of Transcriptomic data

Systems Analysis by constraint based modelling predicts correctly





## **Exponential Random Graphs – Network statistics**





### **Integration of Multi-Omic Data**



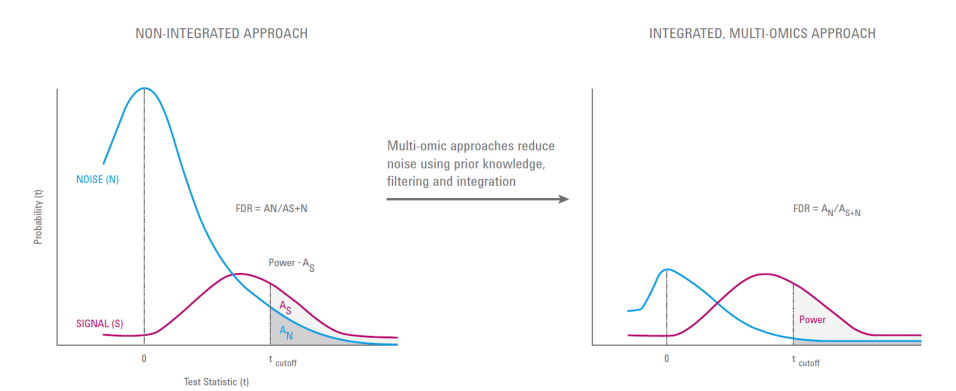
Volume 144, Issue 6, p860-863, 18 March 2011

Leading Edge Minireview

# Boosting Signal-to-Noise in Complex Biology: Prior Knowledge Is Power

Trey Ideker,1,2,\* Janusz Dutkowski,1 and Leroy Hood3

### "Network Biology is a primary tool"



The statistical power of omics experiments can be enhanced through bioinformatics methods that decrease noise through the use of (1) complementary datasets, and (2) incorporation of prior knowledge about the system (e.g., aggregating measurements from entities that belong to the same pathway). This results in an effective decrease in the False Discovery Rate (FDR), at a given t statistic cutoff, or in the ability to relax such cutoff while maintaining the same FDR.



### **Integration of Networks to Enhance Clinical Prediction**

The University of Manchester

Phenotype-gene bilayer network

Phenotype-phenotype similarity

Phenotype-gene association

Gene-gene similarity

NATURE METHODS | VOL.11 NO.3 | MARCH 2014

## Similarity network fusion for aggregating data types on a genomic scale

Bo Wang<sup>1,5</sup>, Aziz M Mezlini<sup>1,2</sup>, Feyyaz Demir<sup>1,2</sup>, Marc Fiume<sup>2</sup>, Zhuowen Tu<sup>3</sup>, Michael Brudno<sup>1,2</sup>, Benjamin Haibe-Kains<sup>4,5</sup> & Anna Goldenberg<sup>1,2</sup>

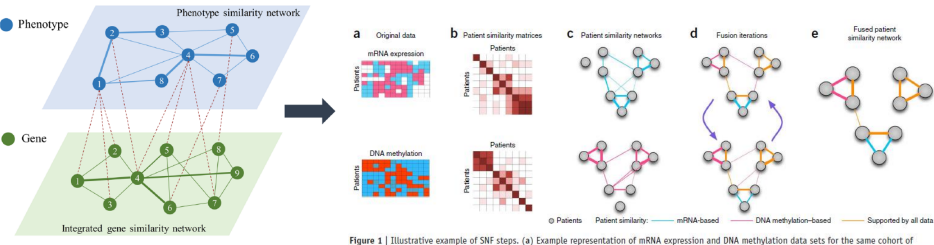
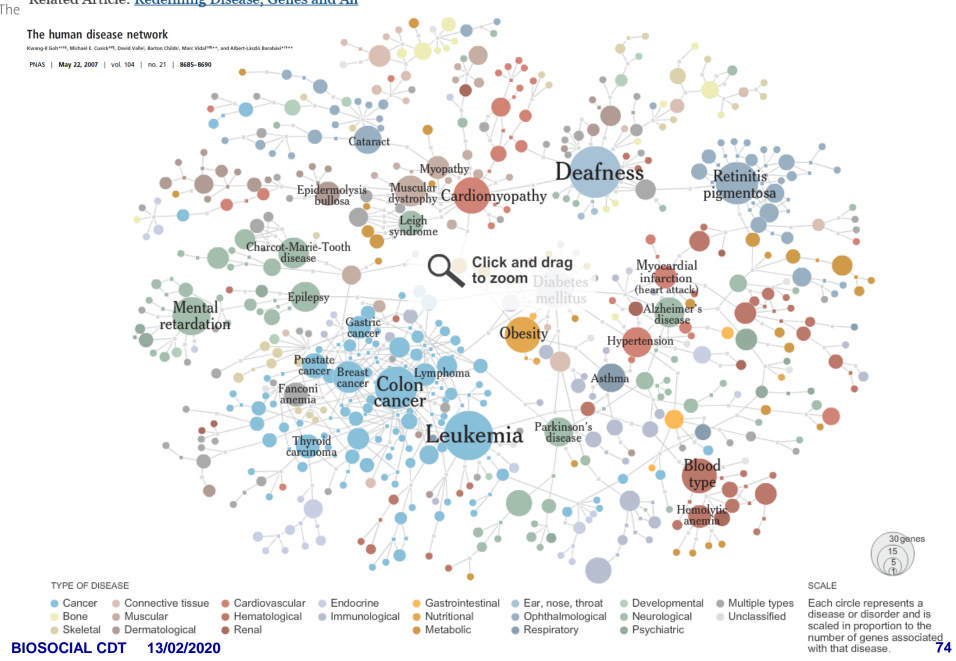


Figure 1 | Illustrative example of SNF steps. (a) Example representation of mRNA expression and DNA methylation data sets for the same cohort of patients. (b) Patient-by-patient similarity matrices for each data type. (c) Patient-by-patient similarity networks, equivalent to the patient-by-patient data. Patients are represented by nodes and patients' pairwise similarities are represented by edges. (d) Network fusion by SNF iteratively updates each of the networks with information from the other networks, making them more similar with each step. (e) The iterative network fusion results in convergence to the final fused network. Edge color indicates which data type has contributed to the given similarity.

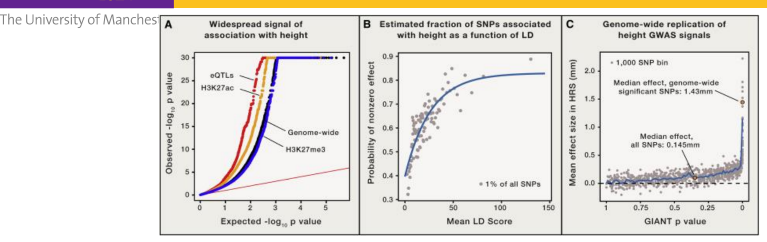
## Mapping the Human 'Diseasome'

Researchers created a map linking different diseases, represented by circles, to the genes they have in common, represented by squares. Related Article: Redefining Disease, Genes and All



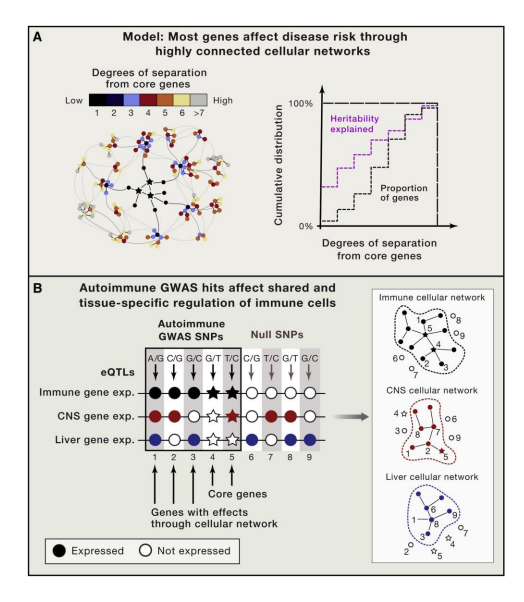


### An Expanded View of Complex Traits: From Polygenic to Omnigenic



- A central goal of genetics is to understand the links between genetic variation and disease.
- Intuitively, one might expect disease-causing variants to cluster into key pathways that drive disease etiology.
- But for complex traits, association signals tend to be spread across most of the genome—including near many genes without an obvious connection to disease.
- We propose that gene regulatory networks are sufficiently interconnected such that all genes expressed in disease-relevant cells are liable to affect the functions of core disease-related genes and that most heritability can be explained by effects on genes outside core pathways.
- We refer to this hypothesis as an "omnigenic" model.

**BIOSOCIAL CDT** 





# Take home messages?