

CG020 Genomika
BI7201 Základy genomiky

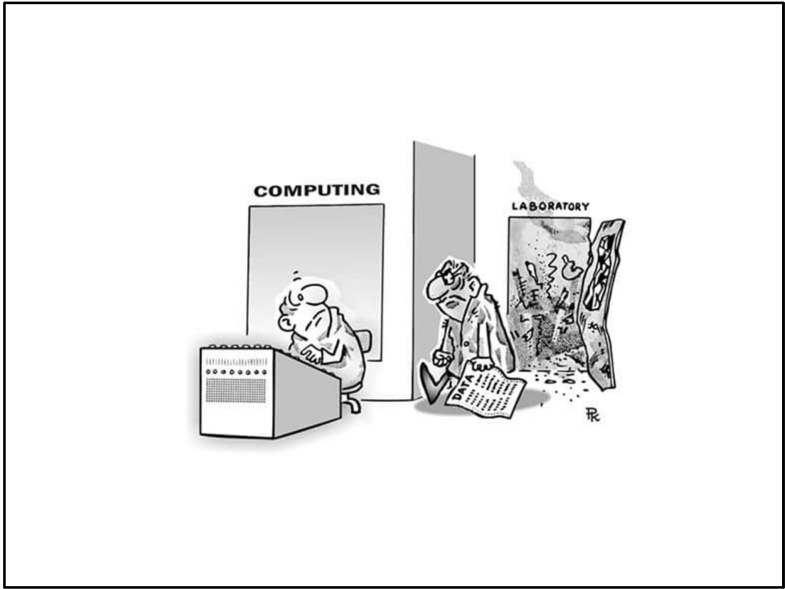
High throughput approaches Systems biology

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Funkční genomika a proteomika rostlin,
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Přehled

- High throughput biology
 - Automation
 - Omics
 - Transcriptomics and high throughput transcriptomics
 - High throughput interactomics and how to read it
 - High throughput of anything
 - 1000(+1) genomes, GWAS
 - ENCODE
 - Epigenome and epitranscriptome
- Little about Systems biology
 - Omics
 - Holism and modules
 - Gene regulation in *E. coli*



Examples of automation in human history



blacksmith



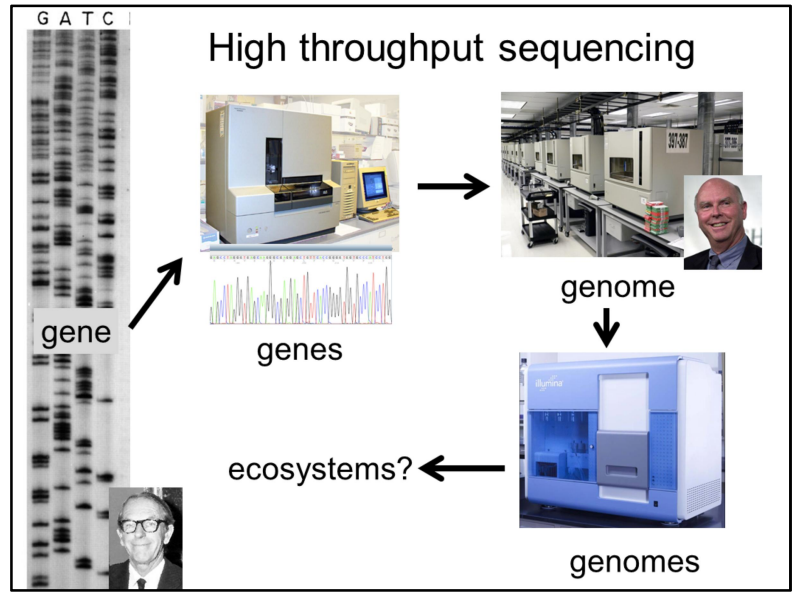
manufacture



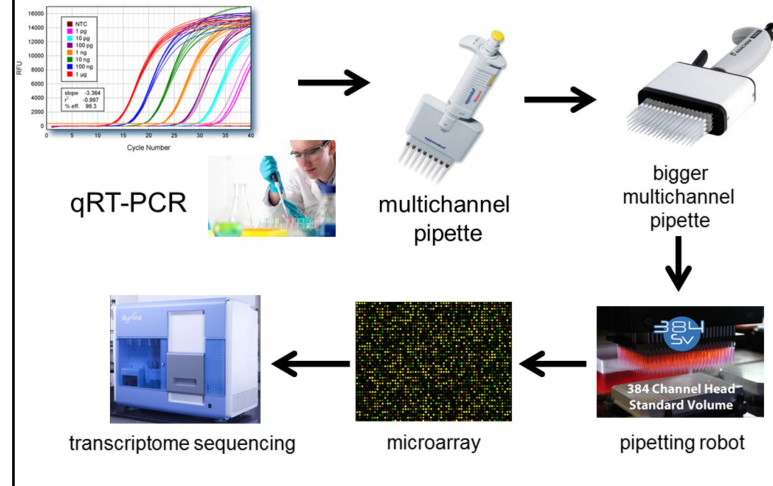
robotic automation



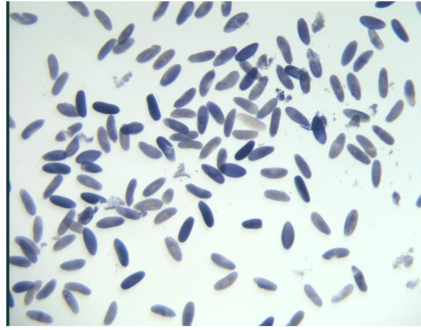
assembly
line



Automation in transcriptomics

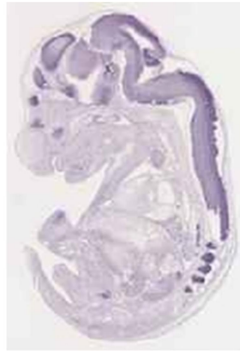


Fl(2)D gene in Drosophila embryos



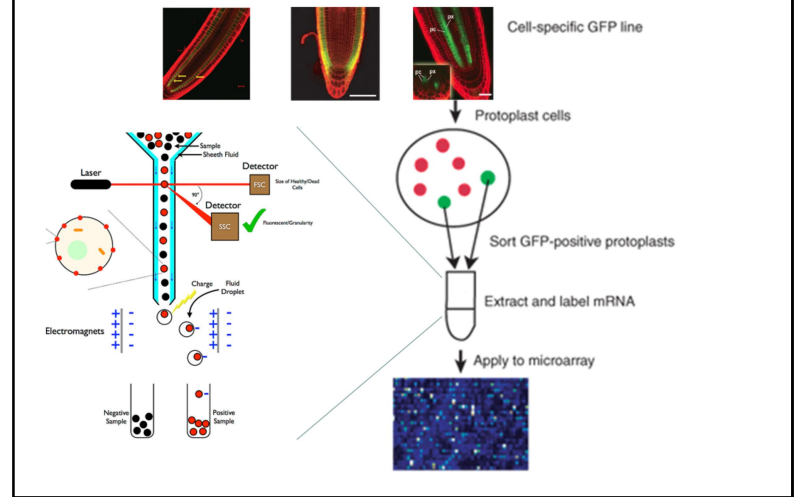
insitu.fluitfly.org

KIAA1841 in mouse expressed in neurons

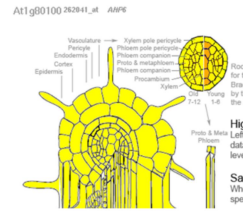
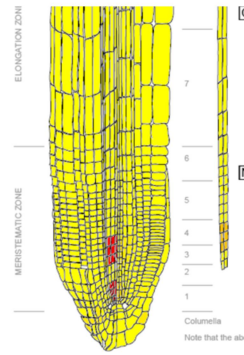
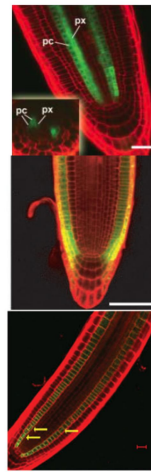


emouseatlas.org

Protoplasting/cell sorting

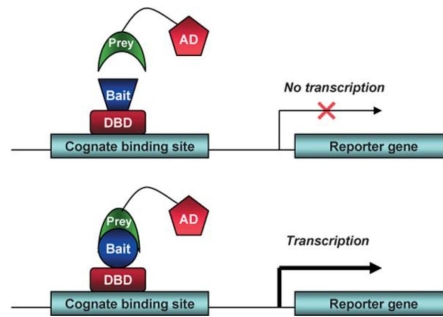


eFP browser



<http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi>

Yeast two-hybrid (Y2H) summary protein-protein interaction hunt



High throughput yeast two hybrid for various organisms

articles

A comprehensive analysis of protein–protein interactions in *Saccharomyces cerevisiae*

(2000)

Peter Uetz¹, Lolic Giot¹, Gerard Cagney¹, Traci A. Mansfield¹, Richard S. Judson¹, James R. Knight¹, Daniel Lockshon¹,
Valbhav Narayan¹, Malithreyan Srinivasan¹, Pascale Pochart¹, Alia Qureshi-Emili¹, Ying Li¹, Brian Godwin¹, Diana Conover¹,
Theodore Kalbitzsch¹, Gowindan Vijayamodar¹, Meijia Yang¹, Mark Johnston¹, Stanley Fields¹ & Jonathan M. Rothberg¹

A Protein Interaction Map of *Drosophila melanogaster*

L. Giot,^{1*} J. S. Bader,^{1**†} C. Brouwer,^{1*} A. Chaudhuri,^{1*}
B. Kuang,¹ Y. Li,¹ Y. L. Hao,¹ C. E. Ooi,¹ B. Godwin,¹ E. Vitols,¹
G. Vijayamodar,¹ P. Pochart,¹ H. Machineni,¹ M. Welsh,¹
Y. Kong,¹ B. Zerhusen,¹ R. Malcolm,¹ Z. Varrone,¹ A. Collis,¹
M. Minto,¹ S. Burgess,¹ L. McDaniel,¹ E. Stimpson,¹ F. Spriggs,¹
J. Williams,¹ K. Neurath,¹ N. Joime,¹ M. Agee,¹ E. Voss,¹
W. Furtak,¹ R. Renzulli,¹ N. Aanensen,¹ S. Carroll,¹
Nickelhaupt,¹ Y. Lazovatsky,¹ A. DaSilva,¹ J. Zhong,²
Itanyon,² R. L. Finley Jr.,² K. P. White,² M. Braverman,¹
D. R. Vrie,¹ S. Gold,¹ M. Leach,¹ J. Knight,¹ R. A. Shimkets,¹
M. P. McKenna,¹ J. Chant,^{1,†} J. M. Rothberg¹

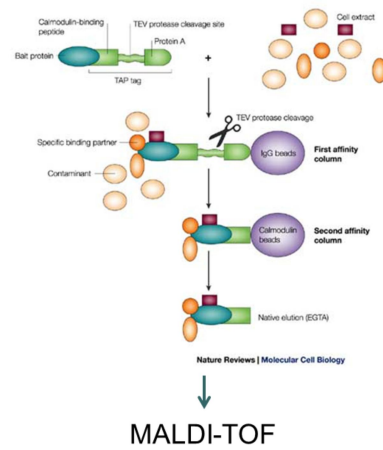
Evidence for Network Evolution in an *Arabidopsis* Interactome Map

(2009)

Arabidopsis Interactome Mapping Consortium[†]

(2005)

TAP purification affinity purification interaction hunt



So, far high throughput affinity purification
approach slightly less popular

**Functional organization of the yeast
proteome by systematic analysis of
protein complexes** (2002)

Anno-Claudio Gavin · Markus Bösch · Roland Krause · Paola Grandi · Martina Marzocchi · Andreas Bauer · Jörg Schmitt ·
Alex M. Beck · Anne-Marie Michon · Cristina-María Ducual · Marita Rumber · Christian Hellert · Margherita Schiader · Miro Brajncević ·
Heide Ruffner · Alejandro Morán · Karin Kuster · Mamada Nishikawa · David Dickson · Tatjana Rodi · Volker Ogris · Angela Basch ·
Goran Borstnik · Bettina Huber · Christina Luchinat · Marie-Kline Heuvelink · Richard B. Goppey · Joseph Edman · Erik Sauerblich ·
Vladimir Rybin · Gerard Drewes · Manfred Ralda · Tavis Bowmester · Peer Bork · Bertrand Seraphin · Bernhard Kuster ·
Gitta Heuvelink · & Giulio Superti-Furga¹

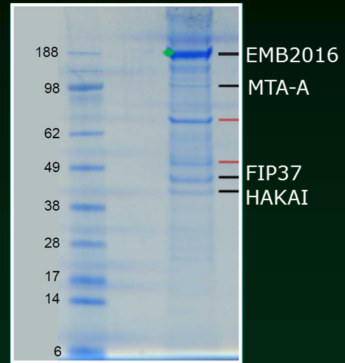
**A Protein Complex Network
of *Drosophila melanogaster*** (2011)

K.G. Gurusarsha,^{1,4} Jean-François Rual,^{1,4} Bo Zhai,^{1,4} Julian Mintseris,^{1,4} Pujita Vaidya,¹ Namita Vaidya,¹
Chapman Beekman,¹ Christina Wong,¹ David Y. Rhee,¹ Odise Cenaj,¹ Emily McKillip,¹ Saumini Shah,¹ Mark Stapleton,²
Kenneth H. Wan,² Charles Yu,² Bayan Parsa,² Joseph W. Carlson,² Xiao Chen,² Bhavoon Kapadia,² K. VijayRaghavan,²
Steven P. Gygi,³ Susan E. Celniker,² Robert A. Obar,^{1,5} and Spyros Artavanis-Tsakonas^{1,6}

thebiogrid.org - highly relevant for searching for interactors,
but look also elsewhere!

Interactors of EMB2016

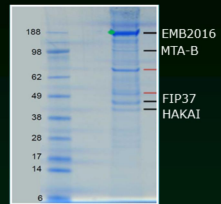
use databases if you have a conserved complex



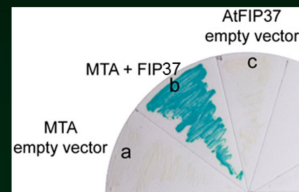
tandem affinity purification

Geert de Jaeger lab

EMB2016 interactors – RNA methylase



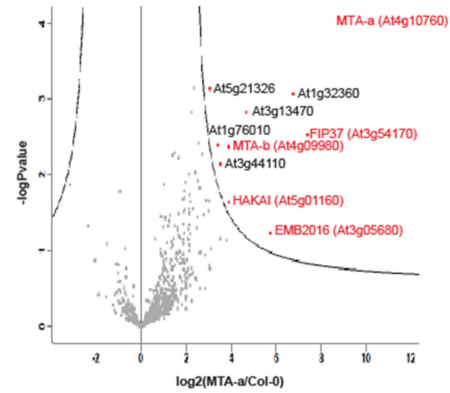
RING finger/HAKAI was also shown to associate with splicing factors (human)



MTA-A – homolog of MTA

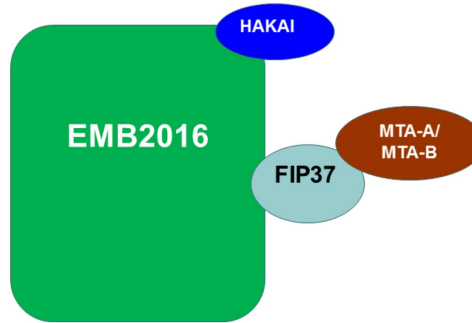
Zhong et al. 2009

All guys back here when using MTA-A as bait

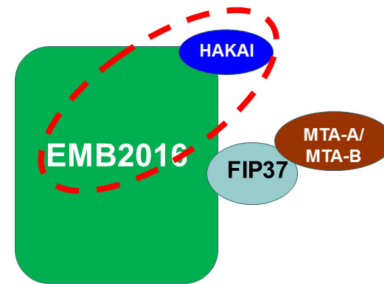


(Immunoprecipitation)

Inferred protein complex



Inferred protein complex



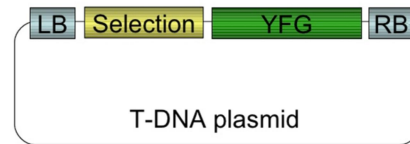
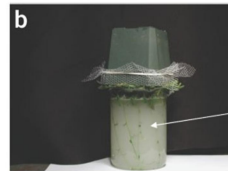
Flybase: EMB2016 interacts with HAKAI (no data on Biogrid)

Summary of Physical Interactions

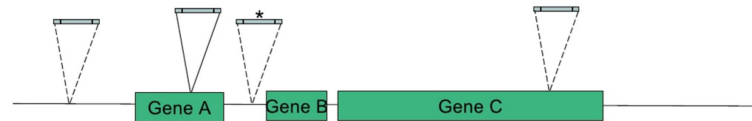
RNA-protein		
Interacting group	Assay	References
vir - stau	anti bait coimmunoprecipitation, partial dna sequence identification by hybridization	(Laver et al., 2013)
protein-protein		
Interacting group	Assay	References
vir - CG7358	experimental knowledge based	(Guruharsha et al., 2011)
vir - Hakai	experimental knowledge based	(Guruharsha et al., 2011)
vir - fl2jd	experimental knowledge based	(Guruharsha et al., 2011)



T-DNA insertion at random locations in the genome

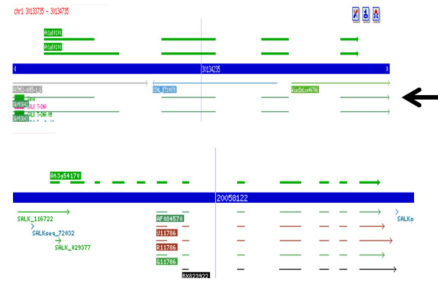


Examples of possible insertions:



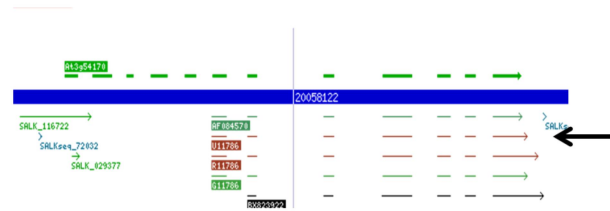
- RNAi/amiRNA (can be also ordered)
- CRISPR

You can order your cDNA clone from the stock center



the same for yeast,
Drosophila, mouse etc.
signal.salk.edu

You can order your cDNA clone from the stock center



even basic fusions (GFP, myc, TAP etc.) often ready for you

signal.salk.edu

You can order antibodies against your protein

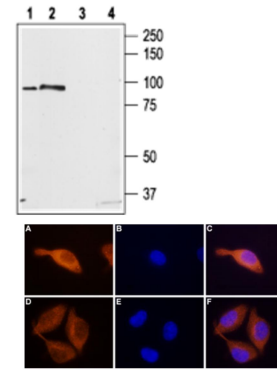
several human proteins
providers:

<http://www.scbt.com/>

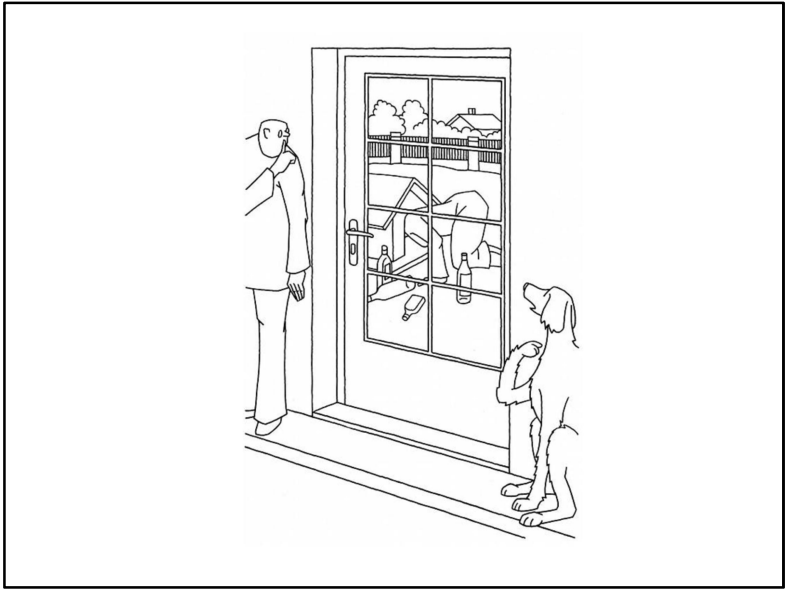
www.acris-antibodies.com/

etc.

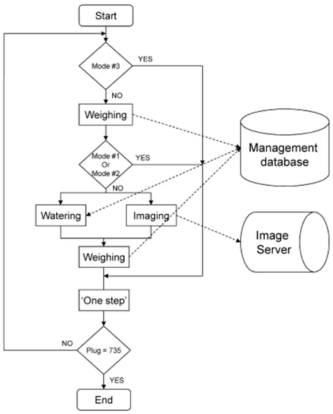
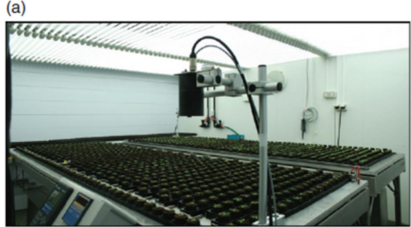
- even get western and
immunocytochemistry in
advance



Arabidopsis so far lagging – agrisera.com perhaps little bit.
Rather commercial service.



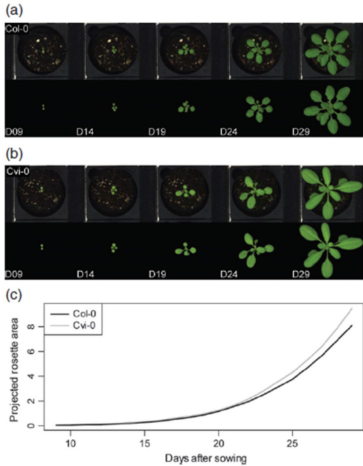
Phenoscope



[PHENOSCOPE: an automated large-scale phenotyping platform](#)

Thisne et al. 2013

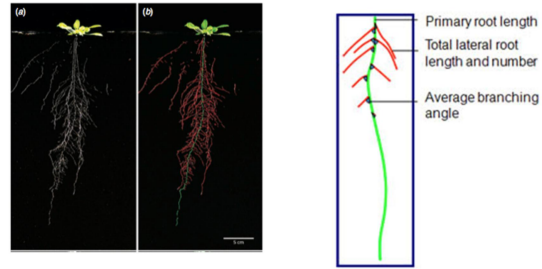
Phenoscope



Phenoscope

- leaf area (camera)
- photosynthesis (spectra)
- weight
- temperature (thermo camera)
- in a dynamic manner
- ...
- various ecotypes only, so far
- commercially promising

Phenoscope – perhaps in future
adaptation on other tissues certainly possible



GrowScreen-Root software

Check your phenotype online

seedgenes.org

– database of plant embryonic mutants (in-dept)

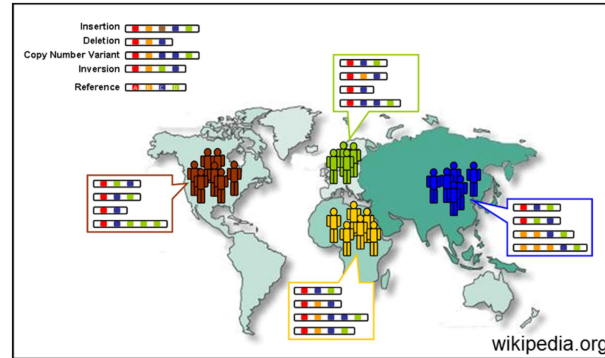


<http://rarge.psc.riken.jp/phenome/>

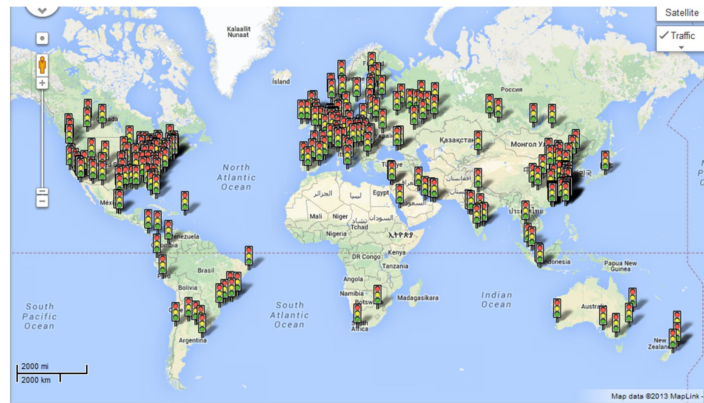
- RIKEN Arabidopsis Phenome Information
Database (kind of attempt on adult plant)

1000 genomes

1000 human genomes over the world



1001 genomes - Arabidopsis



<http://1001genomes.org/>

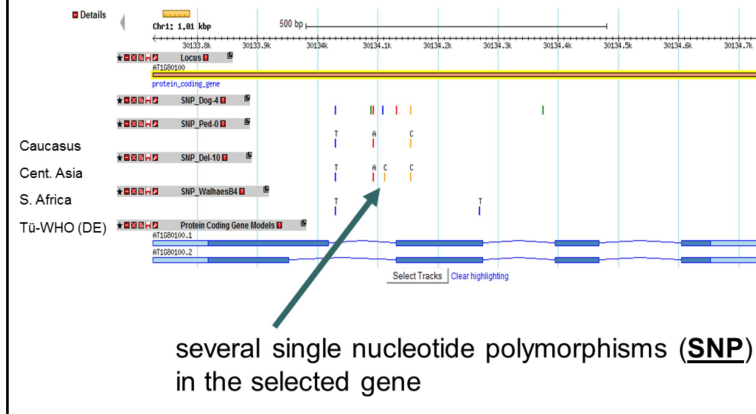
in both cases, much more lines already sequenced

How the ecotypes are collected



Olivier Loudet web page

1001 genomes user interface



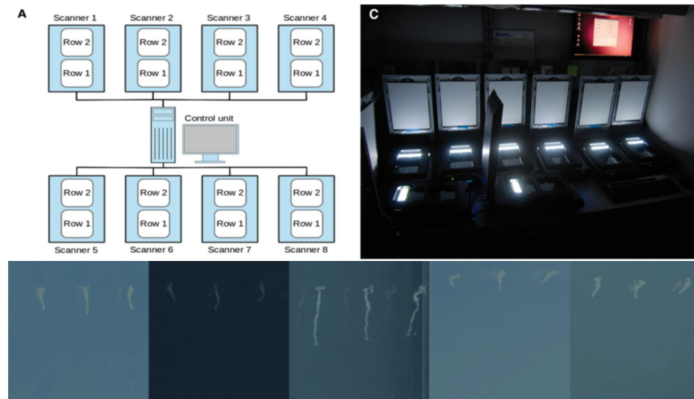
What could be natural variation good for?

What could be natural variation good for?

Quantitative trait loci (QTL)

- nature makes genetic screen for you
- QTL is analogous to gene in genetic screen

Genome wide association studies (GWAS)



Slovak et al. 2014, Busch lab, Vienna

Genome wide association studies (GWAS)

Trait No.	Trait
1	Total length
2	Euclidian length
3	Root tortuosity
4	Root growth rate
5	Relative root growth rate
6	Root angle
7	Root direction index
8	Root horizontal index
9	Root vertical index
10	Root linearity
11	Average root width
12	Root width 20
13	Root width 40
14	Root width 60
15	Root width 80
16	Root width 100

163 accessions (ecotypes),
several replicates (8 x 3)



searching for those different
(say how different they might be!)

Slovak et al. 2014

Genome wide association studies (GWAS)

Trait No.	Trait
1	Total length
2	Euclidian length
3	Root tortuosity
4	Root growth rate
5	Relative root growth rate
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16	Root width 100

163 accessions (ecotypes),
several replicates (8 x 3)



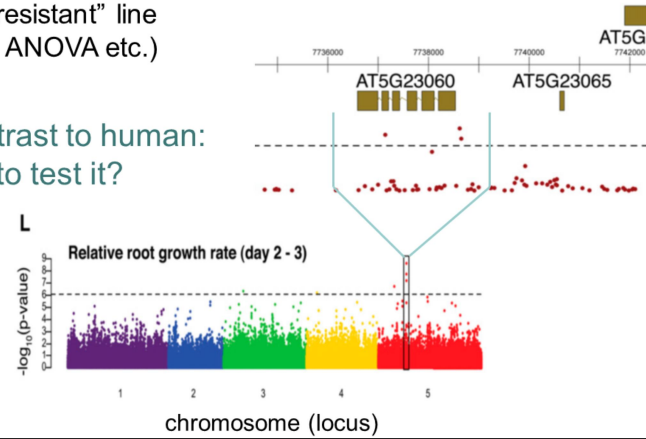
searching for those different
(e. g. root growth, slim root,
resistant to exogenous treatment)

Slovak et al. 2014

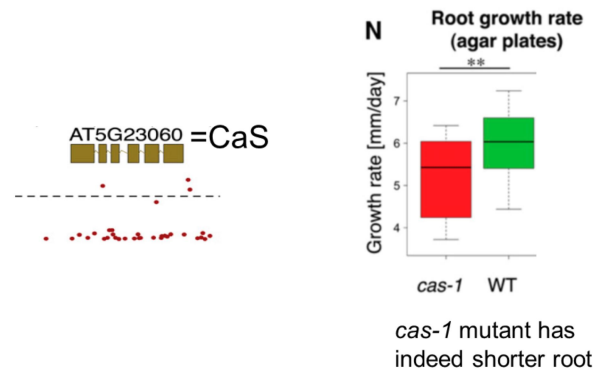
Genome wide association studies (GWAS)

high p-value => SNP specifically
in the "resistant" line
(*N*-way ANOVA etc.)

In contrast to human:
- how to test it?

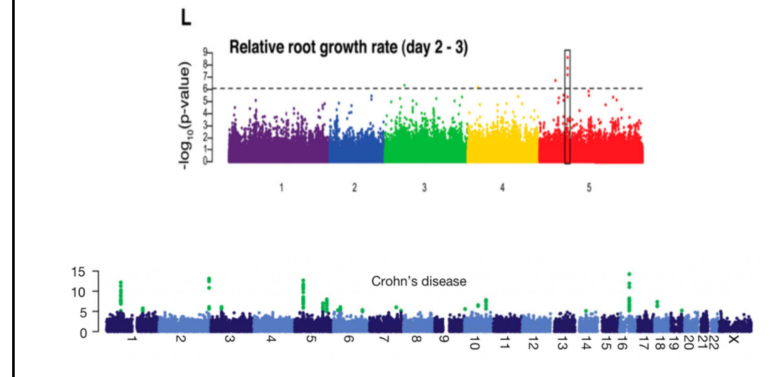


Genome wide association studies (GWAS)



Slovak et al. 2014

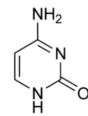
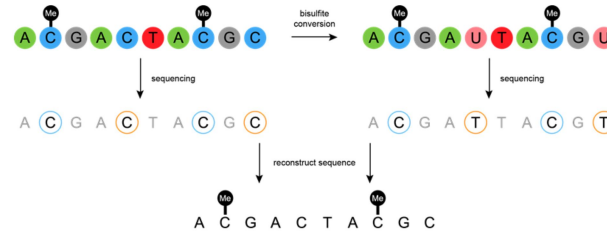
Genome wide association studies (GWAS) Manhattan plot by human



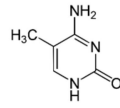
How to find methylated bases in genome?

Which bases are methylated?

How to sequence methylation of genome?

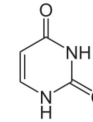


cytosine



5-methylcytosine

bisulfite sequencing



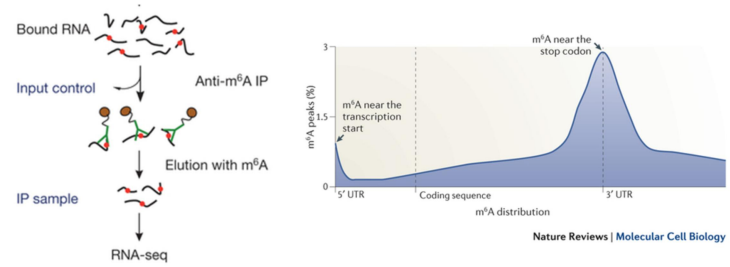
uracil

What is methylation of cytosine good for?

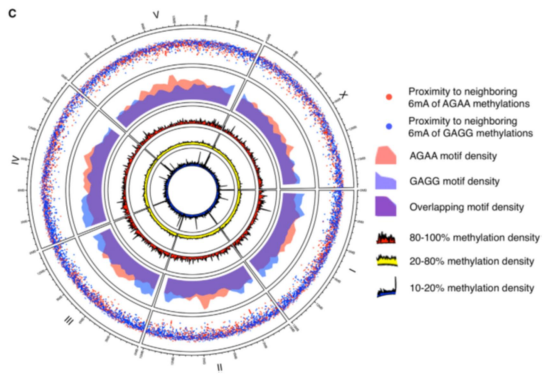
Are there other covalent modifications?

>100 base modification detected in nucleic acids, incl. RNA
N6-methyl adenosine most common in mRNA
(0.5 – 5 % adenosines methylated)

MeRIP – detecting adenine methylation on RNA



Similar technique also adapted on DNA in *C. elegans*



Greer et al. DNA Methylation on N6-Adenine in *C. elegans*, Cell 2015



The ENCODE project
The Encyclopedia of DNA Elements

Is really only ~1 % human genome functional?

1 % = gene coding regions

September 2012

ENCODE – think big

- 80 million dollars (1/2 yearly GAČR budget)
- 1,640 data sets
- 147 cell types
- Nature (6), Genome Biology (18), Genome Research (6 papers)

The ENCODE project

Mainly cancer cells, lymphocytes etc.

RNA transcribed regions:

RNA-seq, CAGE, RNA-PET and manual annotation

Protein-coding regions:

mass spectrometry

Transcription-factor-binding sites:

ChIP-seq, DNase-seq

Chromatin structure:

DNase-seq, FAIRE-seq, histone ChIP-seq and MNase-seq

DNA methylation sites:

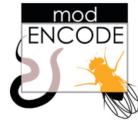
RRBS assay (cheaper version of bisulfite seq)

ENCODE - summary

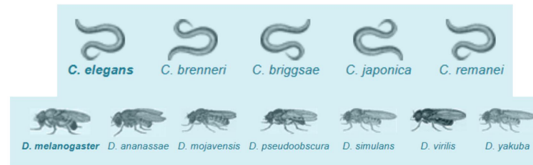
~80 % genome associated with biochemical function:

- enhancers, promoters
- transcribed to non-coding RNA
- 75 % genome transcribed, at least little bit
- number of recognition sequences of DNA binding proteins doubled

E. g. 75 % meaningful number?



ModENCODE on the way



Drosophila tissue sources:

- Adult eclosion + several days
- Adult female
- Adult male
- Embryos 0-1, 0-2, 0-12, 10-12 hr etc
- Larvae in various instars
- Pupae in various stages
- Mated males or females
- etc.

<http://www.modencode.org/>

Question: where do you see the limits of high throughput biology?

Cons

- sometimes low quality data or artifacts
- occasionally data missing
- biological material is quite complex
- what to do with so many data?
- where is the idea?

What is systems biology

- next name for something between biology and chemistry?
biochemistry -> proteomics
molecular biology -> (functional) genomics
- a real new concept?



“Multidimensional biology”

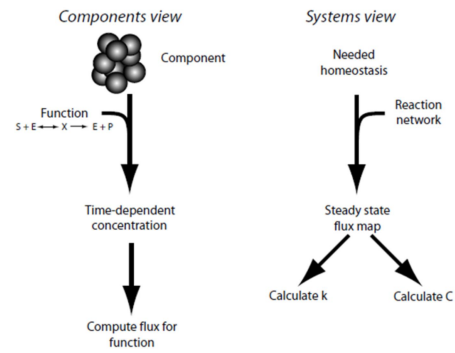
- Genomics
- Epigenomics
- Transcriptomics
- Epitranscriptomics
- Translatomics / Proteomics
- Metabolomics
- Interactomics
- Fluxomics
- NeuroElectroDynamics
- Phenomics
- Biomics

Systems theory

Forget about reductionism, think holistically.

ὅλος [hol'-os] – greek. all, the whole, entire, complete

Reductionism vs. holism



Ludwig von Bertalanffy

(1901-1972)

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GENERAL SYSTEM THEORY

Gathered here are Ludwig von Bertalanffy's writings on general system theory, selected and edited to show the evolution of systems theory and to present its applications to problem solving. An attempt to formulate common laws that apply to virtually every scientific field, this conceptual approach has had a profound impact on such widely diverse disciplines as biology, economics, psychology, and demography.

A German-Canadian biologist and philosopher, Ludwig von Bertalanffy (1901-1972) was the creator and chief exponent of general system theory. He is the author of ten books including *Robots, Men, and Minds* and *Modern Theories of Development*, both which have been published in several languages.

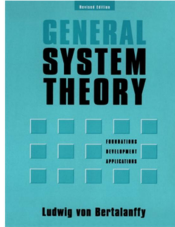
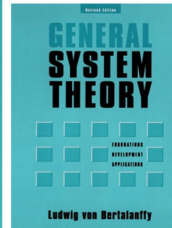
Also available from George Brazzler, Inc.

The Systems View of the World
ISBN 0-8076-0636-7, pb, \$7.95

The Relevance of General Systems Theory
ISBN 0-8076-0609-6, hb, \$8.95

Hierarchy Theory
ISBN 0-8076-0674-X, hb, \$7.95

GEORGE BRAZZLER, INC.
171 Madison Avenue
New York, NY 10016
ISBN 0-8076-0403-4

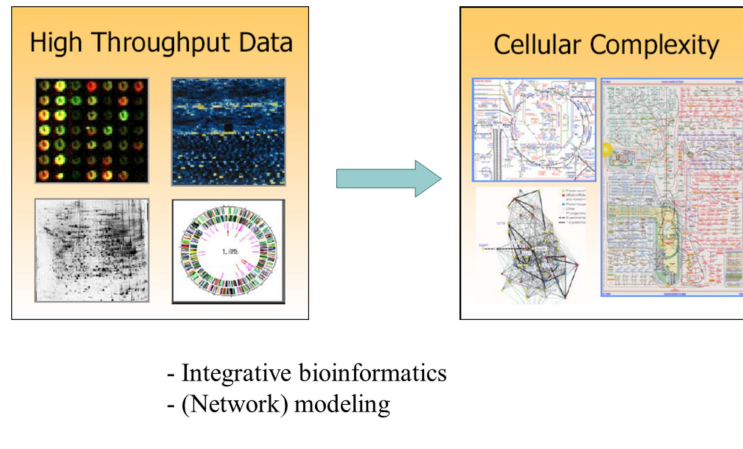



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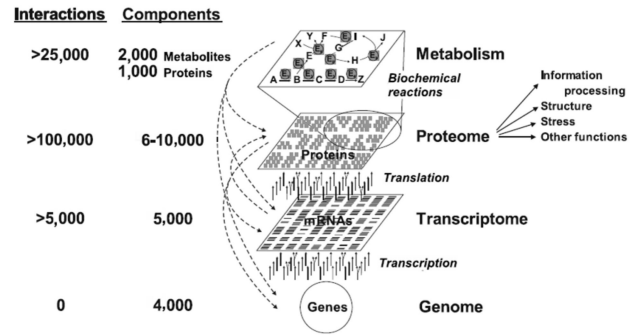
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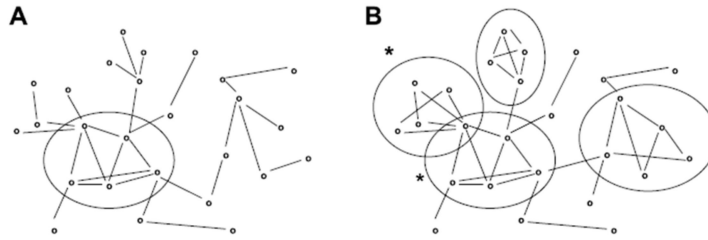
Omic-revolution shifts paradigm to large systems



E. coli genome and proteome is small



Reductionism within holism



Lets e.g. assume that transcription and translation is one module.

Conclusions

- computing capacities allow handling large data sets
- fashionable
- modelling whole cell processes in silico?
- story frequently missing, there will be always question marks

Great web sites

http://www.yeastgenome.org/	<i>S. cerevisiae</i>
http://www.pombase.org/	<i>S. pombe</i>
http://flybase.org/	<i>Drosophila</i>
http://www.wormbase.org/	<i>C. elegans</i>
http://www.arabidopsis.org/	<i>A. thaliana</i>

Also nice web sites

<http://encodeproject.org/>

<http://www.thebiogrid.org/>

<http://www.genemania.org/>

<http://string-db.org/>

...and many others

...pay attention, if they are kept alive and curated

Additional literature

- Venter, J.C. (2008). A life decoded: my genome, my life (London: Penguin).
- Albert-László Barabási (2005) V pavučině síti. (Paseka) (znamenitá kniha o matematické síti, dynamicky se rozvíjejícím oboru od předního světového vědce)
- PA052 Úvod do systémové biologie, Přednášky, Fakulta Informatiky MU
- http://www.youtube.com/watch?v=Z_BHVFPOLk and further – excellent talks about systems biology from Uri Alon (Weizman Institute) – absolutely best
- <http://www.pnas.org/content/110/29/11952> (paper which challenges something conclusions in ENCODE)