CG920 Genomics

Lesson 5

Gene Expression and Chemical Genetics

Jan Hejátko

Functional Genomics and Proteomics of Plants,

Mendel Centre for Plant Genomics and Proteomics, Central European Institute of Technology (CEITEC), Masaryk University, Brno <u>hejatko@sci.muni.cz</u>, <u>www.ceitec.muni.cz</u>





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Literature

• Literature sources for Chapter 05:

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- Lecuyer, E., Yoshida, H., Parthasarathy, N., Alm, C., Babak, T., Cerovina, T., Hughes, T.R., Tomancak, P., and Krause, H.M. (2007). Global analysis of mRNA localization reveals a prominent role in organizing cellular architecture and function. Cell 131, 174-187.
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- Methods of gene expression analysis
 - Qualitative analysis of gene expression
 - Preparation of transcriptional fusion of promoter of analysed gene with a reporter gene
 - Preparation of translational fusion of the coding region of the analysed gene with reporter gene
 - Use of the data available in public databases
 - Tissue- and cell-specific gene expression analysis
 - Quantitative analysis of gene expression
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- Regulation of gene expression in the identification of gene function by gain-of-function approaches
 - T-DNA activation mutagenesis
 - Ectopic expression and regulatable gene expression systems
- Chemical genetics

- Methods of gene expression analysis
 - Qualitative analysis of gene expression
 - Preparation of transcriptional fusion of promoter of analysed gene with a reporter gene



- **Transcriptional fusion with a promoter region**
 - Identification and cloning of the promoter region of the gene
 - Preparation of recombinant DNA carrying the promoter and the reporter gene (uidA, GFP)
 - Preparation of transgenic organisms carrying this recombinant DNA and their histological analysis





GUS reporter in mouse embryos





LacZ activity marks the cells of the developing myotome.

- Methods of gene expression analysis
 - Qualitative analysis of gene expression
 - Preparation of transcriptional fusion of promoter of analysed gene with a reporter gene
 - Preparation of translational fusion of the coding region of the analysed gene with reporter gene



intron exon 9

- Translational fusion of coding region of analysed gene with a reporter gene
 - Preparation of transgenic organisms carrying the recombinant DNA and their histological analysis
 - Compared to transcriptional fusion, translation fusion allows analysis of e.g. Intercellular localization of gene product (protein) or its dynamics



PIN1-GFP in Arabidopsis



Histone 2A-GFP in Drosophila embryo by PAM

- Methods of gene expression analysis
 - Qualitative analysis of gene expression
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 - Use of the data available in public databases

Analysis of expression using Genevestigator (AHP1 and AHP2, Arabidopsis, Affymetrix ATH 22K Array)

🛑 258184_at



Analysis of expression using Genevestigator (AHP1 and AHP2, Arabidopsis, Affymetrix ATH 22K Array)

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Analysis of expression using Genevestigator (AHP1 and AHP2, Arabidopsis, Affymetrix ATH 22K Array)



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 - Use of the data available in public databases
 - Tissue- and cell-specific gene expression analysis



Brady et al., Science, 2007

Expression Maps - RNA

High-Resolution Expression Map in Arabidopsis Root





(A) The majority of enriched GO terms (hierarchically clustered) are associated with individual cell types (blue bar). A smaller number are present across multiple cell types (green bar). (fig. S2) (B) GO category enrichment for hair cells confirms a previous report (15). Enriched ciselements and an enriched TF family were also identified. (C) From the top 50% of varying probe sets, 51 dominant radial patterns were identified. Pattern expression values were meannormalized (rows) and log₂ transformed to yield relative expression indices for each marker line (columns). Marker line order is the same for all figures; see table S1 for marker line abbreviations. (D) Pattern expression peaks were found across one to five cell types. (E to G) Patterns where expression is enriched in single and multiple cell types support transcriptional regulation of auxin flux and synthesis. In all heat maps with probe sets, expression values were mean-normalized and log₂ transformed. Expression is false-colored in representations of a root transverse section, a cut-away of a root tip, and in a lateral root primordium. (E) Auxin biosynthetic genes (CYP79B2, CYP79B3, SUPERROOT1, and SUPERROOT2) are transcriptionally enriched in the QC, lateral root primordia, pericycle, and phloem-pole pericycle $(P = 1.99E^{-11})$, pattern 5). All AGI identifiers and TAIR descriptions are found in table S14. (F) Auxin amido-synthases GH3.6 and GH3.17 that play a role in auxin homeostasis show enriched expression in the columella, just below the predicted auxin biosynthetic center of the QC (P = 8.82E⁻⁴, pattern 13). (G) The expression of the auxin transporter, *PIN-FORMED2*, and auxin transport regulators (PINOID, WAG1) are enriched in the columella, hair cells, and cortex (P = 1.03E⁻⁴, pattern 31).



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Expression Maps - Proteins

Human Protein Atlas







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Schematic flowchart of the Human Protein Atlas. For each gene, a signature sequence (PrEST) is defined from the human genome sequence, and following RT-PCR, cloning and production of recombinant protein fragments, subsequent immunization and affinity purification of antisera results inmunospecific antibodies. The produced antibodies are tested and validated in various immunoassays. Approved antibodies are used for protein profiling in cells (immunofluorescence) and tissues (immunohistochemistry) to generate the images and protein expression data that are presented in the Human Protein Atlas (Ponten et al., *J Int Med*, 2011).





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Expression Maps - Proteins

 Human Protein Atlas (http://www.proteinatlas.org/)

THE HUMAN PROTEIN ATLAS

ABOUT & HELP

| SEARCH ? » | | | | |
|--|----|------|-------|----------|
| e.g. CD44, ELF3, KLK3, or use Fields to search specific fields such as protein_class:Transcription factors or chromosome:X | Se | arch | Clear | Fields » |





The Human Protein Atlas project is funded by the Knut & Alice Wallenberg foundation.



Expression Maps - Proteins

 Human Protein Atlas (http://www.proteinatlas.org/)



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 - DNA and protein chips

DNA Chips

DNA chips

- Method, which provides quick comparison of a large number of genes/proteins between the test sample and control
- Oligo DNA chips are used the most



- There are commercialy available kits for the whole genome
 - company Operon (Qiagen), 29.110 of 70-mer oligonucleotides representing 26.173 genes coding proteins, 28.964 transcripts and 87 microRNA genes of *Arabidopsis thaliana*
 - Possibility of use for the preparation of photolithography chips facilitation of oligonucletide synthesis e.g. for the whole human genome (about 3,1 x 10⁹ bp) jit is possible to prepare 25-mers in only 100 steps, by this technique



Chips not only for the analysis of gene expression, but also for e.g. Genotyping (SNPs, sequencing with chips, ...)

Affymetrix ATH1 Arabidopsis genome array

| Number of arrays | One |
|--------------------------------|--|
| Number of sequence represented | >24,000 gene sequences |
| Feature size | 18µm |
| Oligonucleotide probe length | 25-mer |
| Probe pairs/sequence | 11 |
| Control sequences | E. coli genes bioB, bioC, bioD. B. subtilis gene lysA. Phage P1 cre gene. Arabidopsis maintenance genes GAPDH, Ubiquitin, and Actin |
| Detection sensitivity | 1:100,000* |

DNA Chips

- DNA chips, analysis of results
 - For the correct interpretation of the results, good knowledge of advanced statistical methods is required
 - It is necessary to include a sufficient number of controls and repeats
- Control of accuracy of the measurement (repeated measurements on several chips with the same sample, comparing the same samples analysed on different chips with each other)
- Control of reproducibility of measurements (repeated measurements with different samples isolated under the same conditions on the same chip – comparing with each other)
- Identification of reliable measurement treshold
- Finally comparing the experiment with the control or comparing different conditions with each other -> the result



 Currently there's been a great number of results of various experiments in publicly accessible databases Che et al., 2002

Protein Chips

- Protein chips
 - Chips with high density containing 10⁴ proteins
 - Analysis of protein-protein interactions, kinase substrates and interactions with small molecules
 - Possibility of using antibodies more stable than proteins





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Protein Chips

- Identification of proteins interacting with integrin $\alpha_{IIb}\beta_3$ cytoplasmic domain of platelets
 - Expression of cytoplasmic part as a fusion peptide biotin-KVGFFKR
 - Analysis of binding to the protein chip containing 37.000 clones of *E. coli* expressing human recombinant proteins
 - Confirmation of interaction by pulldown analysis of peptides and by coprecipitation of whole proteins as well (e.g. chloride channel lcln)
 - Other use: e.g. in the identification of kinase substrates, when substrates are bound to the chip and exposed to kinases in the presense of radiolabeled ATP (786 purified proteins of barely, of which 21 were identified as CK2α kinase substrates; Kramer et al., 2004)



Lueking et al., 2005







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Next Gen Transcriptional Profiling

Transcriptional profiling via RNA sequencing



Results of –omics Studies vs Biologically Relevant Conclusions

Transcriptional profiling yielded more then 7K differentially regulated genes...

Ddii et al., unpublished

| gene | locus | sample 1 | sample_2 | status v | /alue 1 | value 2 | log2(fold_change) | test_stat | p value | a value | significant |
|-----------|-------------------------|---------------|------------|----------|------------|-----------|-------------------|------------------|-------------|-----------------|-------------|
| 3 | | | <u>-</u> - | | _ | | | 1.79769e+ | P | 0.0003918 | - |
| AT1G07795 | 1:2414285-2414967 | WT | MT | ОК | (| 1,1804 | 1.79769e+308 | 308 | 6.88885e-05 | | 1 yes |
| | | | | | | | | 1.79769e+ | | 4.67708e- | |
| HRS1 | 1:4556891-4558708 | WT | MT | ОК | (| 0,696583 | 1.79769e+308 | 308 | 6.61994e-06 | | yes |
| | 1 0007 170 000000 | | | | | | | 1.79769e+ | | 0,0005350 | |
| ATMLO14 | 1:9227472-9232296 | VV I | MT | ок | (| 0,514609 | 1.79769e+308 | 308 1.79769e+ | 9.74219e-05 | 3.50131e- | 5 yes |
| NRT1.6 | 1:9400663-9403789 | WT | МТ | ок | ſ | 0 877865 | 1.79769e+308 | 308 | 3.2692e-08 | | yes |
| | 1.9400003-9403789 | VVI | | UK | L. L. | 0,077003 | 1.797096+300 | 1.79769e+ | 5.20928-00 | 07 | yes |
| AT1G27570 | 1:9575425-9582376 | WТ | MT | ок | (| 2.0829 | 1.79769e+308 | 308 | 9.76039e-06 | 6.647e-05 | ves |
| | 1:22159735- | | | | | , | | 1.79769e+ | | 9.84992e- | , |
| AT1G60095 | 22162419 | WT | MT | OK | (| 0,688588 | 1.79769e+308 | 308 | 9.95901e-08 | 07 | yes |
| | | | | | | | | 1.79769e+ | | | |
| AT1G03020 | 1:698206-698515 | WT | MT | ОК | (| 1,78859 | 1.79769e+308 | 308 | 0,00913915 | 5 0,027795 | 8 yes |
| 1710/0000 | | | | | | 0 0 / / | | 1.79769e+ | | | _ |
| AT1G13609 | 1:4662720-4663471 | VV I | MT | ОК | (| 3,55814 | 1.79769e+308 | 308 1.79769e+ | 0,00021683 | 3 0,0010807 | 9 yes |
| AT1G21550 | 1:7553100-7553876 | \ // Т | МТ | ок | ſ | 0 562868 | 1.79769e+308 | 1.79769e+ 308 | 0,00115582 | 0 0047140 | 7.000 |
| A11021000 | 1.7333100-7333870 | VVI | | UK | · · · | 0,302000 | 1.797096+300 | 1.79769e+ | 0,00113302 | 1.91089e- | r yes |
| AT1G22120 | 1:7806308-7809632 | WТ | MT | ок | C | 0.617354 | 1.79769e+308 | 308 | 2.48392e-06 | | yes |
| | 1:11238297- | | | | | -, | | 1.79769e+ | | 0,0002851 | |
| AT1G31370 | 11239363 | WT | MT | OK | (| 1,46254 | 1.79769e+308 | 308 | 4.83523e-05 | | 3 yes |
| | 1:13253397- | | | | | | | 1.79769e+ | | 5.46603e- | |
| APUM10 | 13255570 | WT | MT | ОК | (| 0,581031 | 1.79769e+308 | 308 | 7.87855e-06 | | yes |
| 1710/0700 | 1:18010728- | | | | | | | 1.79769e+ | | 0,0003747 | |
| AT1G48700 | 18012871 1:21746209- | WТ | MT | ок | (| 0,556525 | 1.79769e+308 | 308 1.79769e+ | 6.53917e-05 | | 6 yes |
| AT1G59077 | 21833195 | wт | МТ | ок | C | 138 886 | 1.79769e+308 | 1.79769e+ 308 | 0,00122789 | 0 00/0681 | Sves |
| A11033077 | 1:22121549- | VV I | | OK | (| 100,000 | 1.1910961000 | 1.79769e+ | 0,00122708 | 0,0043001 | 0 ye3 |
| AT1G60050 | 22123702 | wт | MT | ок | C | 0.370087 | 1.79769e+308 | 308 | 0,00117953 | 3 0.004800 | 1 ves |
| | | | | | | -, | | | -, | , | ., |
| AT4G15242 | | WT | MT | OK | 0,00930712 | 17,9056 | 10,9098 | -4,40523 | 1.05673e-05 | 7.13983e-0 | 5 yes |
| | 5:12499071- | | | | | | | | | | |
| AT5G33251 | 12500433 | WT | | OK | 0,0498375 | - , | , | -9,8119 | | | 0 yes |
| AT4G12520 | 4:7421055-7421738 | WT | MT | OK | 0,0195111 | 15,8516 | 9,66612 | -3,90043 | 9.60217e-05 | 0,0005289 | 04 yes |
| AT1G60020 | 1:22100651- 22105276 | WT | MT | ОК | 0,0118377 | 7,18823 | 9,24611 | 7 50200 | 6.19504e-14 | 1 10000 10 | 100 |
| AT5G15360 | 5:4987235-4989182 | | | OK | 0.0988273 | , | , | -10.4392 | | 1.4988e-12) | · · |
| A13G13300 | 5.4907235-4909162 | VVI | IVII | UN | 0,0900273 | 5 50,4654 | 9,1007 | -10,4392 | . (|) | 0 yes |

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Gain-of-Function Approaches

- Methods for identification of gene function using gain-of-function approaches
 - T-DNA activation mutagenesis
 - Method enabling isolation of dominant mutants by random insertion of constitutive promoter, resulting in overexpression of the gene and therefore in corresponding phenotypic changes
 - First step: preparation of mutant library prepared by tansformation of a strong constitutive promoter or enhancer
 - Next step: search of interesting phenotypes
 - Identification of the affected gene, e.g. by plasmid-rescue





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Activation Mutagenesis









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Isolation of CKI1 gene

- Tatsuo Kakimoto, Science 274 (1996), 982-985 *
- Isolation of the gene using activation mutagenesis

- Mutant phenotype is a phenocopy of exogenous application of cytokinins (*CKI1*, <u>*CYTOKININ INDEPENDENT 1*</u>) K1

plasmid K2 35S::*CK1* rescue cDNA

t-zeatin

no hormones



*

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 - Ectopic expression and regulatable gene expression systems

Regulated Expression Systems

- Regulatable gene expression systems
 - Time- or site-specific regulation of gene expression, leading to a change in phenotype and thereby identification of the natural function of the gene
 - pOP system





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reporter

Regulated Expression Systems



reporter

Regulated Expression Systems



Regulated Expression Systems

- Regulatable gene expression systems
 - Time- or site-specific regulation of gene expression, leading to a change in phenotype and thereby identification of the natural function of the gene
 - pOP system
 - UAS system





Outline

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- Chemical genetics

New trends

- Chemical genetics
- "chemical genetics" more than 50.000/89.631 records in PubMed database (16.10. 2008/29.10. 2015, an increase of 65 %)

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- New trends
 - Chemical genetics
 - "chemical genetics" more than 50.000/89.631 records in PubMed database (16.10. 2008/29.10. 2015, an increase of 65 %)
 - Like in the case of genetics, there are also "forward" and "reverse" genetics approaches
 - Unlike in "classical" genetics approaches, the subject of study is not a gene, but a protein
 - Chemical genetics tries to identify either the target protein after a chemical treatment and after following phenotypic changes ("forward" chemical genetics) or chemicals able to interact with protein of interest ("reverse" chemical genetics)
 - For that purpose there are carried out searches in the libraries of various chemicals (thousands of entries, comercially available)
 - example: **analysis of endomembrane transport** in plants

- Analysis of mechanisms of endomembrane transport by chemical genetics approaches
 - In plants cells there occurr very dynamic processes mediated mainly by endomembrane transport (see film, GFP targeting to the ER)



- Analysis of mechanisms of endomembrane transport by chemical genetics approaches
 - In plants cells there occurr very dynamic processes mediated mainly by endomembrane transport (see film, GFP targeting to the ER)
 - Endomembrane transport is an important regulatory mechanism in signal transduction and regulation of cellular processes









Richter et al., E J Cell Biol (2010)









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Tato prezentace je spolufinancována Evropským sociálním fondem a státním rozpočtem České republiky In the figure, there is simplified scheme of vesicle trafficking pathways, regulated by GNOM and its closest relative, GNOM-LIKE1 (GNL1).

Secretory and membrane proteins are synthesised at the ER (blue) and passed onto the Golgi apparatus (green) by anterograde trafficking in COPII-coated vesicles.

The retrograde route from the Golgi apparatus to the ER is regulated by the ARF-GEFs GNOM (GN) and GNL1, which regulate the recruitment of COPI coats to the Golgi membrane. On the secretory route, proteins are transported to the sorting station, the trans-Golgi network (TGN; lilac).

From there, proteins are either transported to the vacuole (grey) via multivesicular bodies (MVB, also called prevacuolar compartment, PVC, which corresponds to the late endosome; deep blue) or trafficked to the plasma membrane (PM).

Plasma membrane proteins like the auxin efflux carrier PIN1 (red), which accumulates at the basal PM at steady state, are continually internalised and trafficked to the TGN, which resembles the early endosome (EE) in plants.

From the TGN, PIN1 is recycled to the plasma membrane via the recycling endosome (RE; light blue). This pathway is regulated by the ARF-GEF GNOM.



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Tato prezentace je spolufinancována Evropským sociálním fondem a státním rozpočtem České republiky

- Analysis of mechanisms of endomembrane transport by chemical genetics approaches
 - By searching in the "library" of chemicals there were identified those, that lead to the secretion of enzyme (carboxypeptidase Y) in yeast (*S. cerevisiae*) – this enzyme is normally transported to the vacuole via the endomembrane transport
 - Analysis of changes in secretion using dotblot and immunodetection of carboxypeptidase Y in the culture medium with monoclonal antibodies
 Chemical structure of sortins



Detection of vacuole phenotype (tonoplast shape) of yeast by staining with a specific color (MDY-64)





Zouhar et al., 2004

- Analysis of mechanisms of endomembrane transport chemical genetics approaches
 - By searching in the "library" of chemicals there were identified those, that lead to the secretion of enzyme (carboxypeptidase Y) in yeast (*S. cerevisiae*) – this enzyme is normally transported to the vacuole via the endomembrane transport
 - Analysis of changes in secretion using dotblot and immunodetection of carboxypeptidase Y in the culture medium with monoclonal antibodies
 - Identified compounds ("sortins") were able to induce similar changes in *Arabidopsis* as well – transport mechanisms are conserved in yeast and in plants
 - For detailed identification of the molecular proces affected by one of the identified "sortins", the analysis of its influence on a secretion of a marker protein (AtCPY) was performed – sortin 1 specifically inhibits only this secretory pathway
 - Identifcation of mutants with altered sensitivity to sortin 1 (hyper- or hypo-sensitive mutants) by EMS mutagenesis



bv

Shape of plant vacuoles using EGFP:-TIP

Image: state of plant vacuoles using EGFP: state of plant vacuoles using EGFP:

Е

Phenotype of seedlings in the presence of sortins

Zouhar et al., 2004

- Analysis of mechanisms of endomembrane transport by chemical genetics approaches – summary
 - GFP::d-TIP vacuole membrane (tonoplast) labelling and identification of mutations leading to altered tonoplast morphology
 - Chemical genetics in combination with classical genetics – identification of proteins participating in regulation of endomembrane transport
 - Proteomics approaches identification and analysis of vacuole proteome

