

CG020 Genomika

Přednáška 12

Nástroje systémové biologie
Modelové organismy, PCR a zásady navrhování primerů

Jan Hejátko

Funkční genomika a proteomika rostlin,
Mendelovo centrum genomiky a proteomiky rostlin,
Středoevropský technologický institut (CEITEC), Masarykova univerzita, Brno
hejatk@sci.muni.cz, www.ceitec.muni.cz



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Genomika 12

▪ Zdrojová literatura

- Wilt, F.H., and Hake, S. (2004). *Principles of Developmental Biology*. (New York ; London: W. W. Norton)
- Roscoe B. Jackson Memorial Laboratory., and Green, E.L. (1966). *Biology of the laboratory mouse*. (New York: Blakiston Division) <http://www.informatics.jax.org/greenbook/index.shtml>
- Eden, E., Navon, R., Steinfeld, I., Lipson, D., and Yakhini, Z. (2009). GOrilla: a tool for discovery and visualization of enriched GO terms in ranked gene lists. *BMC Bioinformatics* 10, 48.
- The Arabidopsis Genome Initiative. (2000). Analysis of the genome sequence of the flowering plant *Arabidopsis thaliana*. *Nature* 408, 796-815.
- Gregory, S.G., Sekhon, M., Schein, J., Zhao, S., Osoegawa, K., Scott, C.E., Evans, R.S., Burridge, P.W., Cox, T.V., Fox, C.A., Hutton, R.D., Mullenger, I.R., Phillips, K.J., Smith, J., Stalker, J., Threadgold, G.J., Birney, E., Wylie, K., Chinwalla, A., Wallis, J., Hillier, L., Carter, J., Gaige, T., Jaeger, S., Kremitzki, C., Layman, D., Maas, J., McGrane, R., Mead, K., Walker, R., Jones, S., Smith, M., Asano, J., Bosdet, I., Chan, S., Chittaranjan, S., Chiu, R., Fjell, C., Fuhrmann, D., Girn, N., Gray, C., Guin, R., Hsiao, L., Krzywinski, M., Kutsche, R., Lee, S.S., Mathewson, C., McLeavy, C., Messervier, S., Ness, S., Pandoh, P., Prabhu, A.L., Saeedi, P., Smailus, D., Spence, L., Stott, J., Taylor, S., Terpstra, W., Tsai, M., Vardy, J., Wye, N., Yang, G., Shatsman, S., Ayodeji, B., Geer, K., Tsegaye, G., Shvartsbeyn, A., Gebregeorgis, E., Krol, M., Russell, D., Overton, L., Malek, J.A., Holmes, M., Heaney, M., Shetty, J., Feldblyum, T., Nierman, W.C., Catanese, J.J., Hubbard, T., Waterston, R.H., Rogers, J., de Jong, P.J., Fraser, C.M., Marra, M., McPherson, J.D., and Bentley, D.R. (2002). A physical map of the mouse genome. *Nature* 418, 743-750.
- Benitez, M. and Hejatkó, J. Dynamics of cell-fate determination and patterning in the vascular bundles of *Arabidopsis thaliana* (submitted)



MINISTERSTVO ŠKOLSTVÍ, MLÁDEŽE A TĚLOVÝCHOVY

OP Vzdělávání
pro konkurenceschopnost



EVROPSKÝ SOCIÁLNÍ FOND
EVROPSKÝ SOCIÁLNÍ FOND
a státním rozpočtem České republiky

Osnova

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**
- Modelové organismy
 - *Mus musculus*
 - *Arabidopsis thaliana*
- Vybrané **metody molekulární biologie**
 - Příprava transgenních organismů
 - PCR
 - Design a příprava primerů (Dr. Hana Konečná)



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Osnova

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**



EVROPSKÁ UNIE

esf



MINISTERSTVO ŠKOLSTVÍ,
MLÁDEŽE A TĚLOVÝCHOVY



OP Vzdělávání
pro konkurenceschopnost



UNIVERSITA
MASARYKŮV
PRAHA

INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Results of -omics Studies vs Biologically Relevant Conclusions

- Results of **-omics studies** are represented by **huge amount of data**, e.g. differential gene expression. But how to get any **biologically relevant conclusions**?

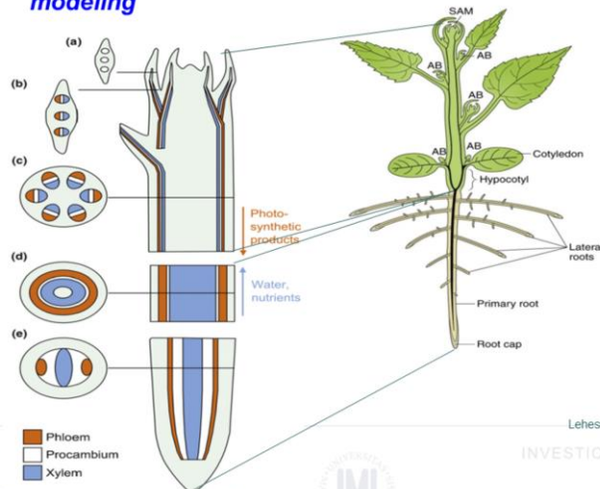
Ddii et al., unpublished

gene	locus	sample_1	sample_2	status	value_1	value_2	log2(fold_change)	test_stat	p_value	q_value	significant
AT1G07795	1.2414285-2414967	WT	MT	OK	0	1.1804	1.79769e+308	308	6.88885e-05	0.00039180	1yes
HRS1	1.4556891-4558708	WT	MT	OK	0	0.895583	1.79769e+308	308	6.61994e-0605	4.57708e-	yes
ATML014	1.9227472-9232296	WT	MT	OK	0	0.514609	1.79769e+308	308	9.74219e-05	0.00053505	5yes
NRT1.6	1.9400663-9403789	WT	MT	OK	0	0.877895	1.79769e+308	308	3.2592e-08 07	3.50131e-	yes
AT1G27570	1.9575425-9582376	WT	MT	OK	0	2.0829	1.79769e+308	308	9.76039e-066.647e-05	9.84992e-	yes
AT1G60095	1.22159735-22162419	WT	MT	OK	0	0.688588	1.79769e+308	308	9.95901e-0807	1.79769e+	yes
AT1G03020	1.698206-698515	WT	MT	OK	0	1.78859	1.79769e+308	308	0.00913915	0.0277958	yes
AT1G13609	1.4662720-4663471	WT	MT	OK	0	3.55814	1.79769e+308	308	0.000216830	0.0108079	yes
AT1G21550	1.7553100-7553876	WT	MT	OK	0	0.562868	1.79769e+308	308	0.001155820	0.0471487	yes
AT1G22120	1.7806308-7809632	WT	MT	OK	0	0.617354	1.79769e+308	308	2.48392e-0605	1.91089e-	yes
AT1G31370	1.11238297-11239363	WT	MT	OK	0	1.46254	1.79769e+308	308	4.83523e-05	0.00028514	3yes
APUM10	1.13253397-13255570	WT	MT	OK	0	0.581031	1.79769e+308	308	7.87855e-0605	5.46603e-	yes
AT1G48700	1.18010728-18012871	WT	MT	OK	0	0.556525	1.79769e+308	308	6.53917e-05	0.00037473	6yes
AT1G59077	1.21746209-21833195	WT	MT	OK	0	138.886	1.79769e+308	308	0.001227890	0.0496816	yes
AT1G60050	1.22121549-22123702	WT	MT	OK	0	0.370087	1.79769e+308	308	0.00117953	0.0048001	yes
AT4G15242	4.8705706-8706997	WT	MT	OK	0.00930712	17.9056	10.9098	-4.40523	1.05673e-057	1.3983e-05	yes
AT5G33251	5.12499971-12500433	WT	MT	OK	0.0498375	52.2837	10.0349	-9.8119	0	0	yes
AT4G12520	4.7421055-7421738	WT	MT	OK	0.0195111	15.8516	9.66612	-3.90043	9.60217e-050	0.000528904	yes
AT1G60020	1.22100655-22105076	WT	MT	OK	0.0118377	7.18823	9.24611	-7.50382	6.19504e-141	4.9898e-12	yes
AT5G15360	5.4987235-4989182	WT	MT	OK	0.0988273	56.4834	9.1587	-10.4392	0	0	yes

Example of an output of transcriptional profiling study using Illumina sequencing performed in our lab. Shown is just a tiny fragment of the complete list, comprising about 7K genes revealing differential expression in the studied mutant.

Molecular Regulatory Networks Modeling

- **Vascular tissue** as a developmental model for **GO analysis** and **MRN modeling**



Lehesranta et al., *Trends in Plant Sci* (2010)



MINISTERSTVO ŽIVOTNÍHO
PROSTŘEDÍ A STAVBY
OP Inovace
pro konkurenceschopnost

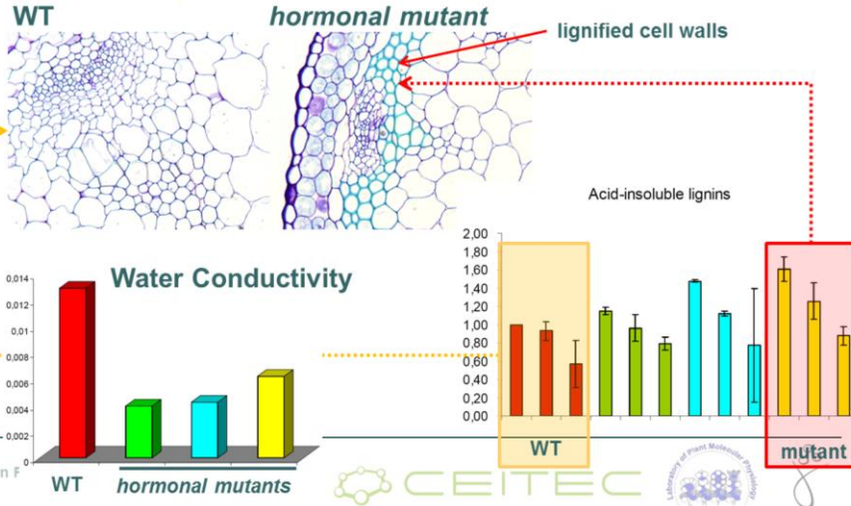


INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Hormonal Control Over Vascular Tissue Development

- Plant **Hormones Regulate Lignin Deposition** in Plant Cell Walls and **Xylem Water Conductivity**

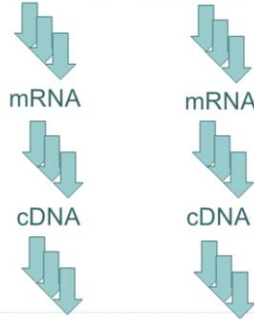
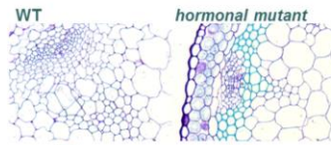


Hormonal Regulation in F



Hormonal Control Over Vascular Tissue Development

□ *Transcriptional profiling* via *RNA sequencing*



Sequencing by Illumina and **number of transcripts** determination

Hormonal Regu...

CETEC



Results of -omics Studies vs Biologically Relevant Conclusions

- Transcriptional profiling yielded more than **7K differentially regulated genes**...

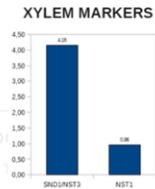
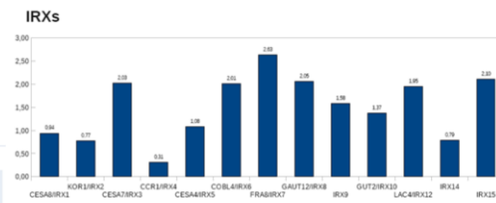
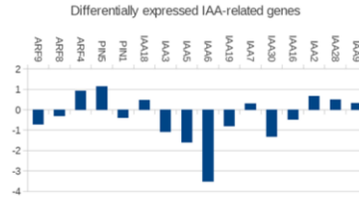
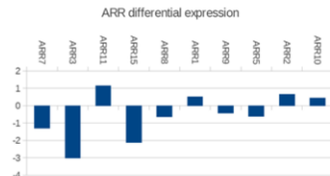
Ddii et al., unpublished

gene	locus	sample_1	sample_2	status	value_1	value_2	log2(fold_change)	test_stat	p_value	q_value	significant
AT1G07795	1.2414285-2414967	WT	MT	OK	0	1.1804	1.79769e+308	308	6.88885e-05	0.00039180	1yes
HRS1	1.4556891-4558708	WT	MT	OK	0	0.896583	1.79769e+308	308	6.61994e-0605	4.67708e-	yes
ATML014	1.9227472-9232296	WT	MT	OK	0	0.514609	1.79769e+308	308	9.74219e-05	0.00053505	5yes
NRT1.6	1.9400663-9403789	WT	MT	OK	0	0.877895	1.79769e+308	308	3.2692e-08	3.50131e-	yes
AT1G27570	1.9575425-9582376	WT	MT	OK	0	2.0829	1.79769e+308	308	9.76039e-066.647e-05	9.84992e-	yes
AT1G60095	1.22159735-22162419	WT	MT	OK	0	0.688588	1.79769e+308	308	9.95901e-0807		yes
AT1G03020	1.698206-698515	WT	MT	OK	0	1.78859	1.79769e+308	308	0.00913915	0.0277958	yes
AT1G13609	1.4662720-4663471	WT	MT	OK	0	3.55814	1.79769e+308	308	0.000216830	0.0108079	yes
AT1G21550	1.7553100-7553876	WT	MT	OK	0	0.562868	1.79769e+308	308	0.001155820	0.0471487	yes
AT1G22120	1.7806308-7806932	WT	MT	OK	0	0.617354	1.79769e+308	308	2.48392e-0605	1.91089e-	yes
AT1G31370	1.11238297-11239363	WT	MT	OK	0	1.46254	1.79769e+308	308	4.83523e-05	0.00028514	3yes
APUM10	1.13253397-13255570	WT	MT	OK	0	0.581031	1.79769e+308	308	7.87855e-0605	5.46603e-	yes
AT1G48700	1.18010728-18012871	WT	MT	OK	0	0.556525	1.79769e+308	308	6.53917e-05	0.00037473	6yes
AT1G59077	1.21746209-21833195	WT	MT	OK	0	138.886	1.79769e+308	308	0.001227890	0.0496816	yes
AT1G60050	1.22121549-22123702	WT	MT	OK	0	0.370087	1.79769e+308	308	0.00117953	0.0048001	yes
AT4G15242	4.8705796-8706997	WT	MT	OK	0.00930712	17.9056	10.9098	-4.40523	1.05673e-057	1.3983e-05	yes
AT5G33251	5.12499971-12500433	WT	MT	OK	0.0498375	52.2837	10.0349	-9.8119	0	0	yes
AT4G12520	4.7421055-7421738	WT	MT	OK	0.0195111	15.8516	9.66612	-3.90043	9.60217e-050	0.000528904	yes
AT1G60020	1.22100651-22105276	WT	MT	OK	0.0118377	7.18823	9.24611	-7.50382	6.19504e-141	4.9898e-12	yes
AT5G15360	5.4987235-4989182	WT	MT	OK	0.0988273	56.4834	9.1587	-10.4392	0	0	yes

Example of an output of transcriptional profiling study using Illumina sequencing performed in our lab. Shown is just a tiny fragment of the complete list, comprising about 7K genes revealing differential expression in the studied mutant.

Gene Ontology Analysis

- One of the possible approaches is to study **gene ontology**, i.e. previously demonstrated **association** of genes to **biological processes**



Ddii et al., unpublished

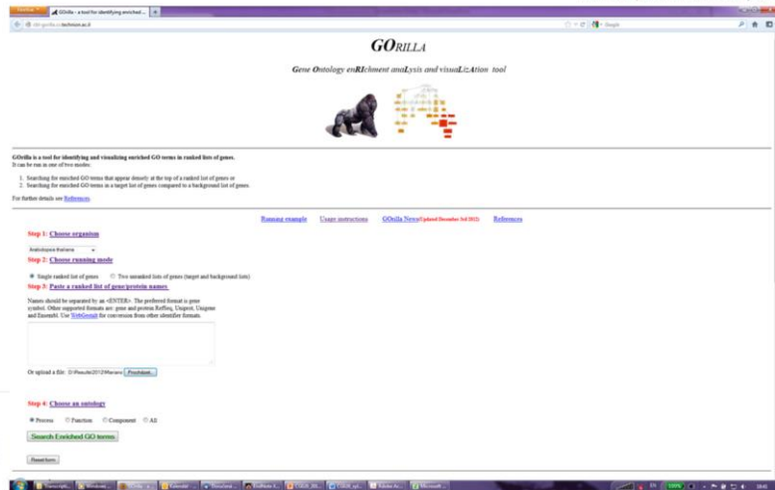


INVESTICE DO
 AVÁNI
 financována
 n fondem
 a státním rozpočtem České republiky

Gene Ontology Analysis

- Several tools allow **statistical evaluation** of **enrichment** for genes **associated with specific processes**

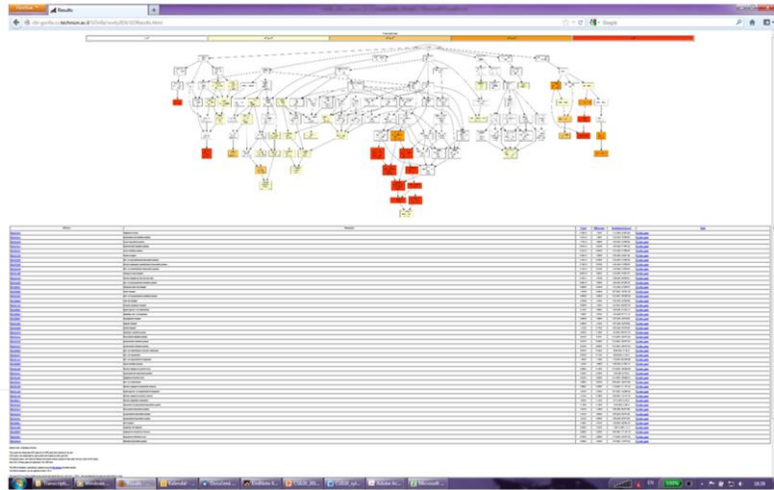
Eden et al., *BMC Bioinformatics* (2009)



One of such recent and very useful tools is Gorilla software, freely available at <http://cbl-gorilla.cs.technion.ac.il/>.

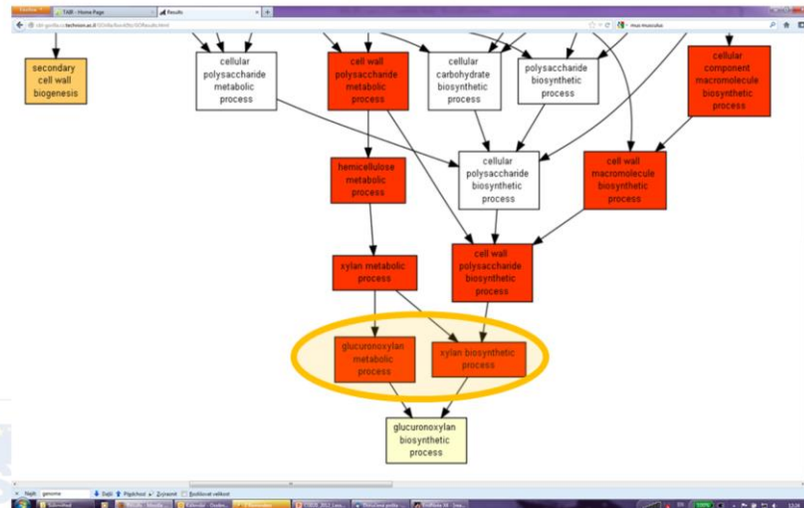
Gene Ontology Analysis

- Several tools allow **statistical evaluation** of **enrichment** for genes **associated with specific processes**



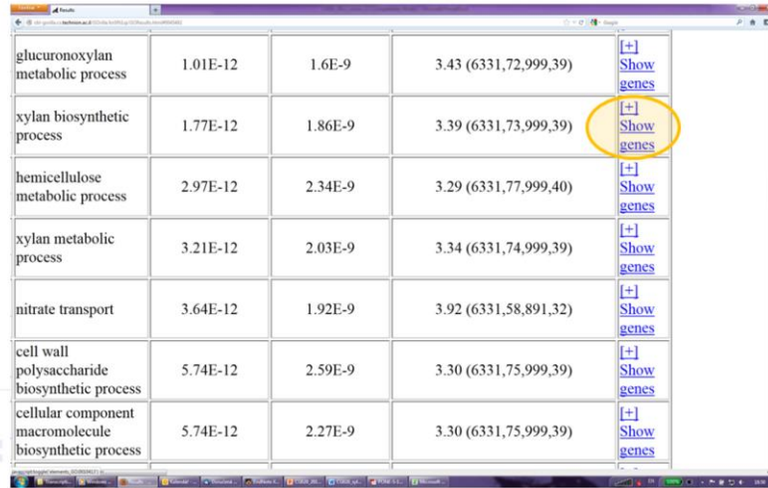
Gene Ontology Analysis

- Several tools allow **statistical evaluation** of **enrichment** for genes **associated with specific processes**



Gene Ontology Analysis

- Several tools allow **statistical evaluation** of **enrichment** for genes **associated with specific processes**



glucuronoxylan metabolic process	1.01E-12	1.6E-9	3.43 (6331,72,999,39)	[+] Show genes
xylan biosynthetic process	1.77E-12	1.86E-9	3.39 (6331,73,999,39)	[+] Show genes
hemicellulose metabolic process	2.97E-12	2.34E-9	3.29 (6331,77,999,40)	[+] Show genes
xylan metabolic process	3.21E-12	2.03E-9	3.34 (6331,74,999,39)	[+] Show genes
nitrate transport	3.64E-12	1.92E-9	3.92 (6331,58,891,32)	[+] Show genes
cell wall polysaccharide biosynthetic process	5.74E-12	2.59E-9	3.30 (6331,75,999,39)	[+] Show genes
cellular component macromolecule biosynthetic process	5.74E-12	2.27E-9	3.30 (6331,75,999,39)	[+] Show genes



Gene Ontology Analysis

- Several tools allow **statistical evaluation** of **enrichment** for genes **associated with specific processes**

Description	P-value	FDR q-value	Enrichment (N, B, n, b)	Genes
response to nitrate	4.76E-13	1.5E-9	4.13 (6331,55,891,32)	[+] Show genes
glucuronoxylan metabolic process	1.01E-12	1.6E-9	3.43 (6331,72,999,39)	[+] Show genes
				[+] Hide genes
				<ul style="list-style-type: none"> GGT2 - putative glycosyltransferase PGAP3 - plant glycoprotein-like starch synthase protein 3 FRAB - mucinase-like protein GAUT12 - alpha 1,4-galactosyltransferase AT1G02140 - bifunctional inhibitor/ligand transfer protein sand storage 2a subunit-like protein AT1G04180 - proteinase 64 AT1G04916 - ring-B2 finger protein at72 LAC11 - lactase 11 KNAT1 - homeobox protein knotted 1-like 7 NAAG12 - nar domain-containing protein 12 IR39 - nucleotide-diphosphate sugar transferase-like protein AT1G01090 - protein tyrosine-like protein C58A4 - cellulose synthase a catalytic subunit 4 [judg-forming] AT1G08140 - rho gppase activating protein with pak-box p21-psi-binding domain CTL2 - chitinase-like protein 2 IR36 - xylanase-like protein 4 MYB84 - myb domain protein 43 PGAP2 - plant glycoprotein-like starch synthase protein 1 AT1G04140 - putative alpha-acetyltransferase AT1G01710 - hypothetical protein AT1G05200 - aspartyl proteinase-like protein AT1G09440 - protein kinase family protein AT1G04600 - polyphenol oxidase domain-like protein AT1G01090 - targeting protein for ubiq-like protein AT1G04710 - hypothetical protein AT1G04230 - hlx-poc domain-containing protein AT1G01910 - hypothetical protein IR409 - putative polyphosphatase sem catalytic subunit pd30 MAP70-3 - microtubule-associated protein 70-3 AT1G02020 - hypothetical protein AGL44 - protein aquaporin-like 44 IR312 - lectin-4 NAAG71 - nar domain-containing protein 71 IR31 - cellulose synthase a catalytic subunit 1 [judg-forming] AT1G02143 - hypothetical protein MYB46 - myosin domain protein 46 AT1G02220 - ring-B2 finger protein at54 IR33 - zinc ribbon family protein AT1G01880 - hypothetical protein
xylan biosynthetic process	1.77E-12	1.86E-9	3.39 (6331,73,999,39)	



Osnova

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**



EVROPSKÁ UNIE

esf



MINISTERSTVO ŠKOLSTVÍ,
MLÁDEŽE A TĚLOVÝCHOVY



ÚIV
NÁRODNÍ ÚSTAV
PRO VĚDECKÝ VÝZKUM V
OBlastI VĚDEK O VĚDOVÉ
KVALITĚ



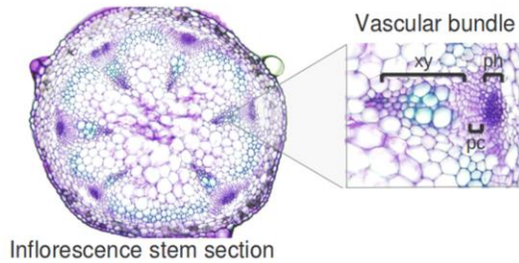
UNIVERSITA
MASARYKŮV
PRAHA

INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Molecular Regulatory Networks Modeling

- **Vascular tissue** as a developmental model for **MRN modeling**



Benitez and Hejato, *submitted*



OP VĚLÁVÁNÍ
úspěšně financována
základním fondem
Evropské unie a
státem České republiky



Molecular Regulatory Networks Modeling

- **Literature search** for published data and creating own database

Interaction	Evidence	References
A-ARRs – CK signaling	Double and higher order type-A ARR mutants show increased sensitivity to CK.	[27]
	Spatial patterns of A-type ARR gene expression and CK response are consistent with partially redundant function of these genes in CK signaling.	[27]
	A-type ARR decreases B-type ARR6-LUC.	[13]
	Note: In certain contexts, however, some A-ARRs appear to have effects antagonistic to other A-ARRs.	[27]
AHP6 – AHP	ahp6 partially recovers the mutant phenotype of the CK receptor WOL.	[9]
	Using an in vitro phosphotransfer system, it was shown that, unlike the AHPs, native AHP6 was unable to accept a phosphoryl group. Nevertheless, AHP6 is able to inhibit phosphotransfer from other AHPs to ARR.	[9]

Signaling and Hormones

Benitez and Hejatko, submitted

www.nature.com/scientificdata/



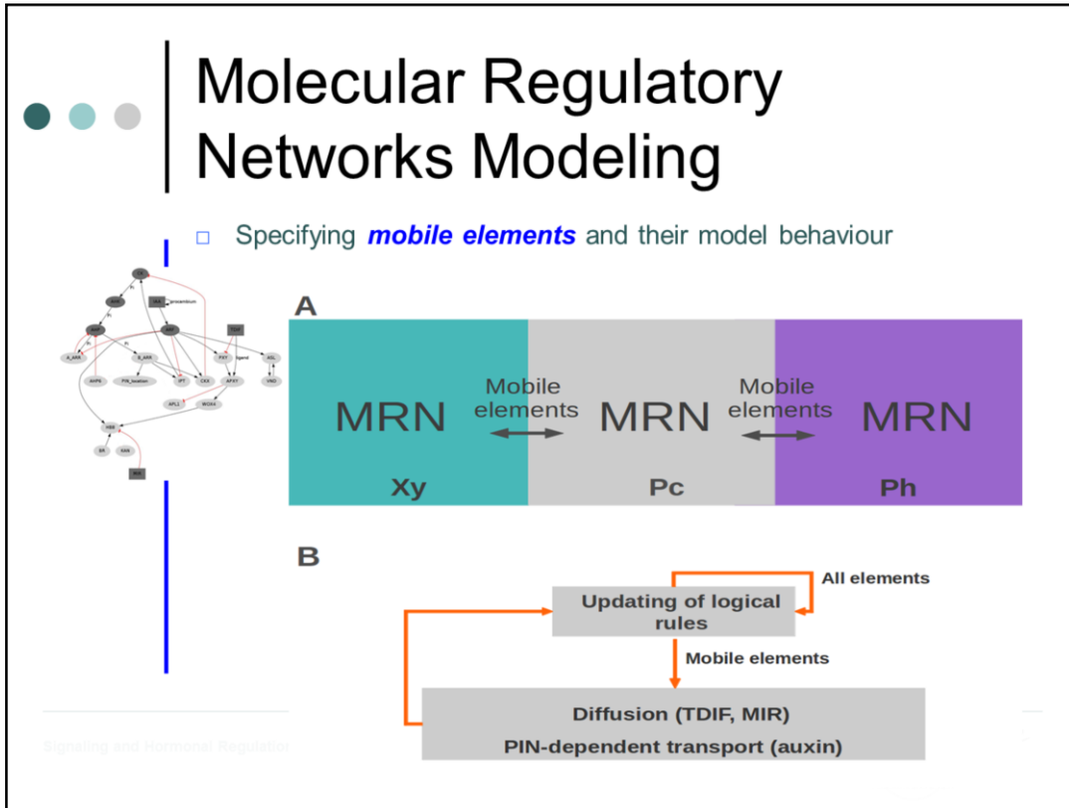


Molecular Regulatory Networks Modeling

- Formulating **logical rules** defining the **model dynamics**

Network node	Dynamical rule
CK	2 If ipt=1 and cxx=0 1 If ipt=1 and cxx=1 0 else
CKX	1 If barr>0 or arf=2 0 else
AHKs	ahk=ck
AHPs	2 If ahk=2 and ahp6=0 and aarr=0 1 If ahk=2 and (ahp6+aarr<2) 1 If ahk=1 and ahp6<1 0 else
B-Type ARRs	1 If ahp>0 0 else
A-Type ARRs	1 If arf<2 and ahp>0 0 else

Benitez and Hejatko, submitted



According to experimental evidence for the system under study, the hormone IAA, the peptide TDIF, and the microRNA MIR165/6 are able to move among the cells. In the case of TDIF and MIR165/6, the mobility is defined as diffusion and is given by the following equation:

$$g(t+1)T[i] = H(g(t)[i] + D(g(t)[i+1] + g(t)[i-1] - N(g(t)[i])) - b) \quad (2),$$

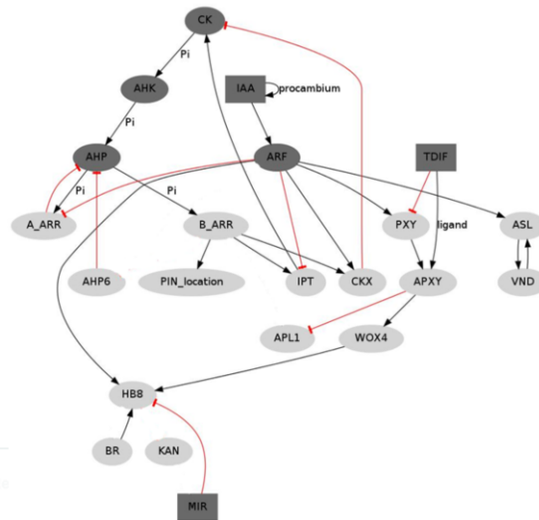
where $g(t)T[i]$ is the total amount of TDIF or MIR165 in cell (i). D is a parameter that determines the proportion of g that can move from any cell to neighboring ones and is correlated to the diffusion rate of g . b is a constant corresponding to a degradation term. H is a step function that converts the continuous values of g into a discrete variable that may attain values of 0, 1 or 2. N stands for the number of neighbors in each cell. Boundary conditions are zero-flux. In the case of IAA, the mobility is defined as active transport dependent on the radial localization of the PIN efflux transporters and is defined by the equation:

$$iaa(t+1)T[i] = H(iaa(t)[i] + Diaa(pin(t)[i+1])(iaa(t)[i+1]) + Diaa(pin(t)[i-1])(iaa(t)[i-1]) - N(Diaa)(pin(t)[i])(iaa(t)[i]) - b) \quad (3),$$

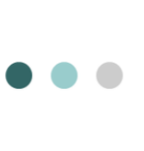
where $Diaa$ is a parameter that determines the proportion of IAA that can be transported among cells. The transport depends on the presence of IAA and PIN in the cells and b corresponds to a degradation term. As in equation 2, H is a step function that converts the continuous values to discrete ones and N stands for the number of neighbors in each cell. Boundary conditions for IAA motion are also zero-flux.

Molecular Regulatory Networks Modeling

- Preparing the *first version* of the model and its *testing*



The proposed model considers data that we identified and evaluated through an extensive search (up to January 2012). It takes into account molecular interactions, hormonal and expression patterns, and cell-to-cell communication processes that have been reported to affect vascular patterning in the bundles of Arabidopsis. The model components and interactions are graphically presented in the figure above. In the network model, nodes stand for molecular elements regulating one another's activities. Most of the nodes can take only 1 or 0 values (light gray nodes in the figure), corresponding to "present" or "not present," respectively. Since the formation of gradients of hormones and diffusible elements may have important consequences in pattern formation, mobile elements TDIF and MIR, as well as members of the CK and IAA signaling systems, can take 0, 1 or 2 values (dark gray nodes in the figure above) Benitez and Hejatko, submitted.



Molecular Regulatory Networks Modeling

- Specifying of missing interactions via *informed predictions*

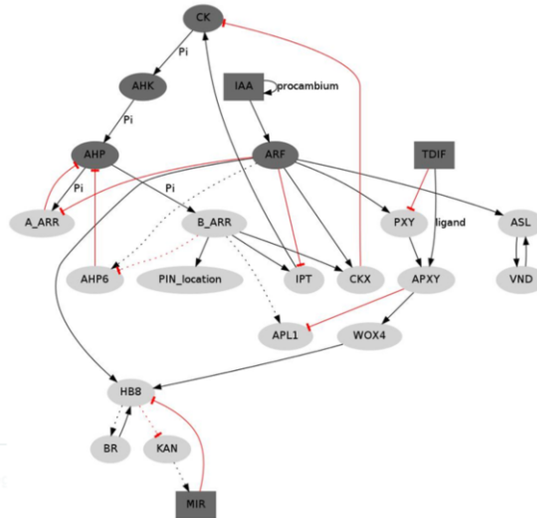
Interaction	Evidence	References
CK → PIN7 radial localization	Predicted interaction (could be direct or indirect)	
	Informed by the following data:	
	During the specification of root vascular cells in <i>Arabidopsis thaliana</i> , CK regulates the radial localization of PIN7.	[18]
CK → APL	Expression of PIN7::GFP and PIN7::GUS is upregulated by CK with no significant influence of ethylene.	[18,20]
	In the root, CK signaling is required for the CK regulation of PIN1, PIN3, and PIN7. Their expression is altered in <i>wol</i> , <i>cre1</i> , <i>ahk3</i> and <i>ahp6</i> mutants.	[19]
CK → APL	Predicted interaction (could be direct or indirect)	
	Consistent with the fact that APL overexpression prevents or delays xylem cell differentiation, as does CKs.	[21]
	Partially supported by microarray data and phloem-specific expression patterns of CK response factors.	(TAIR, ExpressionSet:1005823559, [22])

Signaling and Hormon

PLASMA MEMBRANE

Molecular Regulatory Networks Modeling

- Preparing the *next version* of the model and its *testing*



Benitez and Hejatko, *PlosONE*, 2013

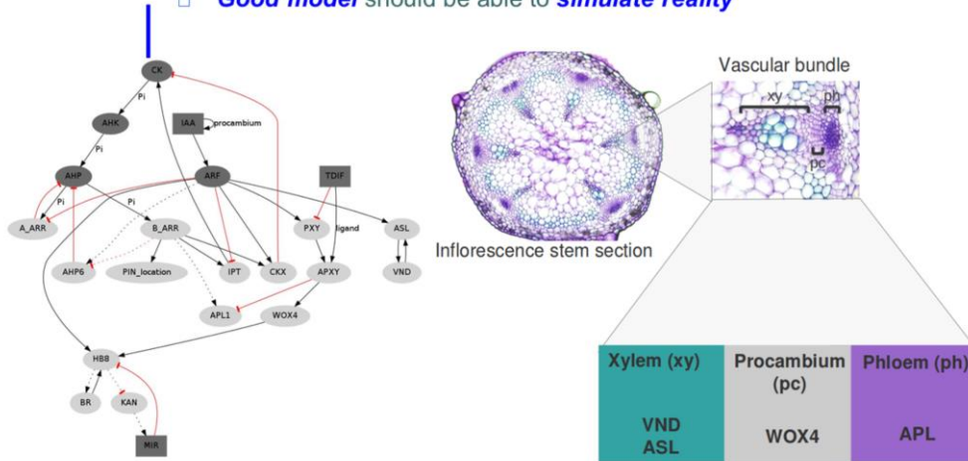
Signaling and Hormonal Reg.



In comparison to the model shown on slide 21, the final version of the model contains the predicted interactions (dashed lines).

Molecular Regulatory Networks Modeling

- *Good model* should be able to *simulate reality*



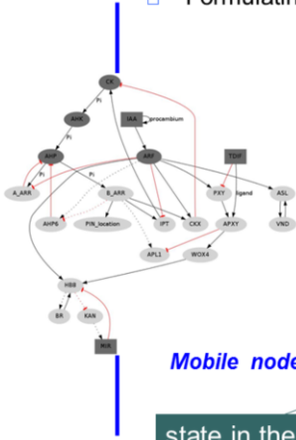
Signaling and Hormonal Regulation of Plant Development

Benitez and Hejatk, PlosONE, 2013



Molecular Regulatory Networks Modeling

- Formulating **equations** describing the **relationships** in the model



logical rule function

state in the time t

Static nodes: $g_n(t+1) = F_n(g_{n1}(t), g_{n2}(t), \dots, g_{nk}(t))$

state in the time $t+1$

Amount of TDIF or MIR165 in cell i

Mobile nodes: $g_{(t+1)T[i]} = H(g_{(t)T[i]} + D(g_{(t)T[i+1]} + g_{(t)T[i-1]} - N(g_{(t)T[i]})) - b)$

state in the time $t+1$

constant corresponding to a degradation term

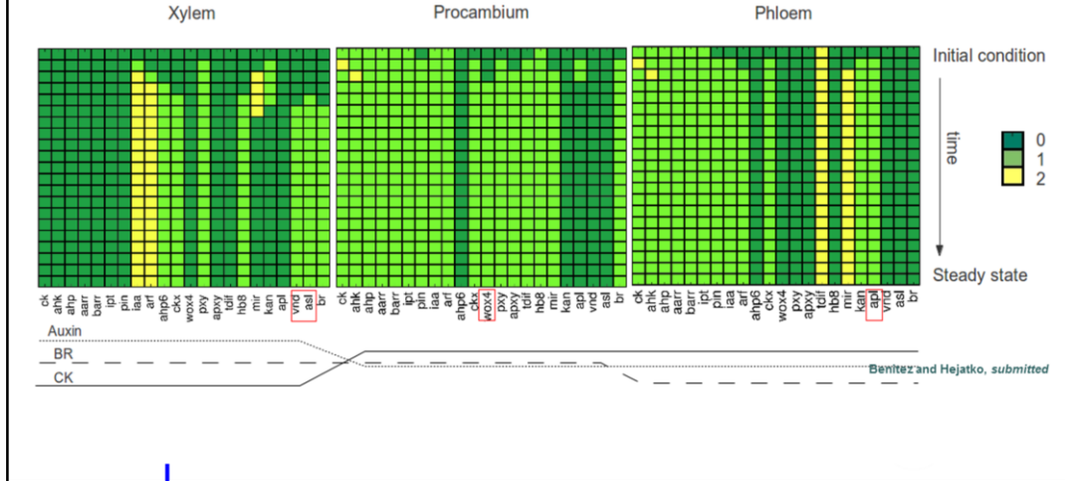
proportion of movable element

Molecular Regulatory Networks Modeling

- **Good model** should be able to **simulate reality**

Static nodes: $g_n(t+1) = F_n(g_{n1}(t), g_{n2}(t), \dots, g_{nk}(t))$

Mobile nodes: $g_{(t+1)T[i]} = H(g_{(t)[i]} + D(g_{(t)[i+1]} + g_{(t)[i-1]} - N(g_{(t)[i]})) - b)$



The initial conditions specify the initial state of some of the network elements (figure above) and are the following :

I) In the procambial position (central compartment), CK is initially available and there is an initial and sustained IAA input or self-upregulation. This condition is supported by several lines of evidence. Also *HB8*, a marker of early vascular development that has been found in procambial cells, is assumed to be initially present at this position. These conditions are not fixed, however. After the initial configuration, all the members of the CK and IAA signaling pathways, as well as *HB8*, can change their states according to the logical rules.

II) In the xylem and phloem positions, it is assumed that no element is initially active except for the CK signaling pathway and TDIF, both in the phloem position. The level of expression for a given node is represented by a discrete variable g and its value at a time $t+1$ depends on the state of other components of the network (g_1, g_2, \dots, g_N) at a previous time unit. The state of every gene g therefore changes according to:

$$g_n(t+1) = F_n(g_{n1}(t), g_{n2}(t), \dots, g_{nk}(t)) \quad (1).$$

In this equation, $g_{n1}, g_{n2}, \dots, g_{nk}$ are the regulators of gene g_n and F_n is a discrete function known as a logical rule (logical rules are grounded in available experimental data, for example see slide 20). Given the logical rules, it is possible to follow the dynamics of the network for any given initial configuration of the nodes expression state. One of the most important traits of dynamic models is the existence of steady states in which the entire network enters into a self-sustained configuration of the nodes state. It is thought that in developmental systems such self-sustained states correspond to particular cell types.

According to experimental evidence for the system under study, the hormone IAA, the peptide TDIF, and the microRNA MIR165/6 are able to move among the cells. In the case of TDIF and MIR165/6, the mobility is defined as diffusion and is given by the following equation:

$$g(t+1)T[i] = H(g(t)[i] + D(g(t)[i+1] + g(t)[i-1] - N(g(t)[i])) - b) \quad (2),$$

where $g(t)T[i]$ is the total amount of TDIF or MIR165 in cell (i). D is a parameter that determines the proportion of g that can move from any cell to neighboring ones and is correlated to the diffusion rate of g . b is a constant corresponding to a degradation term. H is a step function that converts the continuous values of g into a discrete variable that may attain values of 0, 1 or 2. N stands for the number of neighbors in each cell. Boundary conditions are zero-flux. In the case of IAA, the mobility is defined as active transport dependent on the radial localization of the PIN efflux transporters and is defined by the equation:

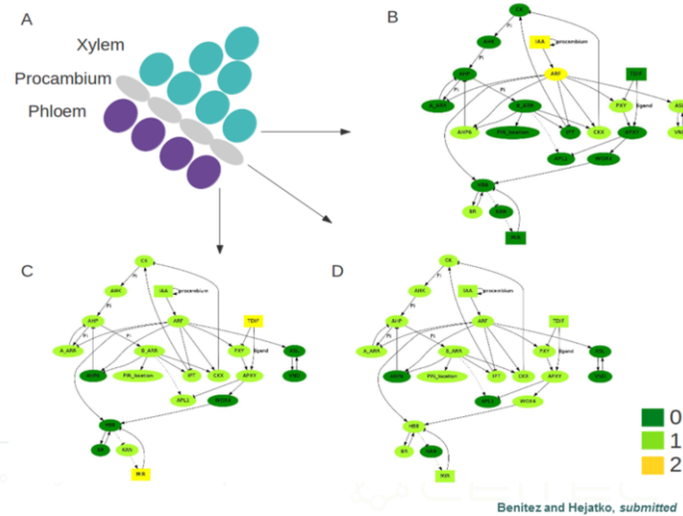
$$iaa(t+1)T[i] = H(iaa(t)[i] + Diaa(pin(t)[i+1])(iaa(t)[i+1]) + Diaa(pin(t)[i-1])(iaa(t)[i-1]) - N(Diaa)(pin(t)[i])(iaa(t)[i]) - b) \quad (3),$$

where $Diaa$ is a parameter that determines the proportion of IAA that can be transported among cells. The transport depends on the presence of IAA and PIN in the cells and b corresponds to a degradation term. As in equation 2, H is a step function that converts the continuous values to discrete ones and N stands for the number of neighbors in each cell. Boundary conditions for IAA motion are also zero-flux.

Using the logical rules, equations 1–3, and a broad range of parameter values (not shown here), it is possible fully to reproduce the results and analyses reported in the following sections (see the figure above for the simulation time course).

Molecular Regulatory Networks Modeling

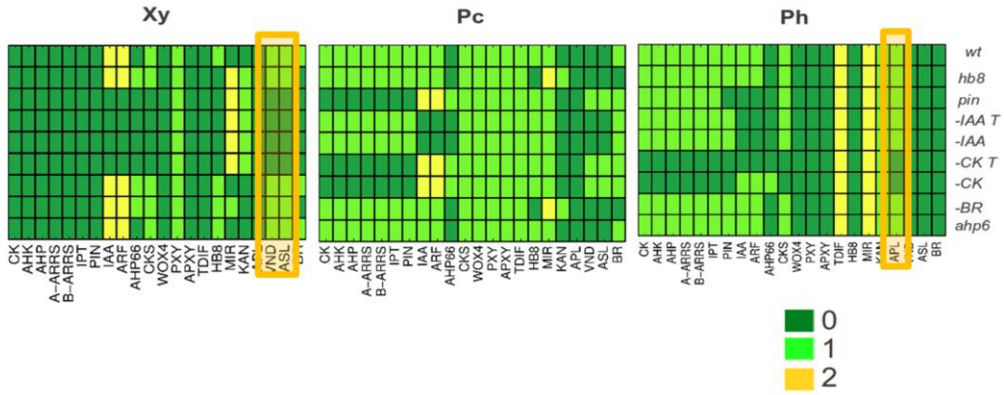
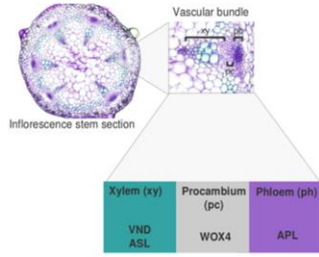
- The **good model** should be able to **simulate reality**



Another representation of the distinct expression profiles in the individual vascular bundle compartments (phloem, procambium and xylem).

Molecular Regulatory Networks Modeling

□ Simulation of *mutants*





Osnova

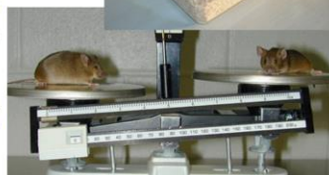
- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**
- Modelové organismy
 - *Mus musculus*



Mus musculus

myš domácí, house mouse

- malé nároky na chovnou plochu
- relativně velké množství mláďat (3-14, v průměru 6-8)
- velikost genomu se blíží velikosti genomu člověka (cca 3000 Mbp), podobně jako počet genů (cca 24K)
- 20 chromozomů (19+1)
- vhodná pro široké spektrum fyziologických experimentů (anatomicky i fyziologicky podobná člověku)
- možno poměrně snadno získávat K.O. mutanty i transgenní linie



INVE

NI

na

am

by

More info about mouse at

<http://www.informatics.jax.org/greenbook/index.shtml>.

Mus musculus

myš domácí, house mouse

- Genom známý od roku 2002 (<http://www.ncbi.nlm.nih.gov/projects/genome/assembly/grc/mouse/>)

The screenshot displays the Mouse Genome Reference Consortium (GRC) website. The main heading is "Mouse Genome Overview" with the subtitle "Information concerning the continuing improvement of the mouse genome." Below this, there is an ideogram of the mouse genome with chromosomes 1 through 19, X, and Y. Red arrows point to specific regions on chromosomes 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, X, and Y. A legend indicates that red arrows represent "Regions containing alternate-loci" and red squares represent "Regions containing fix patches." An orange box highlights the "Next assembly update" information: "The next assembly update (patch release 2) will be a minor update (only patches) and will happen in March 2013." The page also features a "Getting Data" section with links to GRCm38 p1, GRCm38, and MGSv37. On the right side, there is a "GRC Blog" section with articles such as "The GRC and the 10th International Zebrafish Genetics and Development Meeting" and "Hidden assembly problems." The bottom of the page includes logos for the European Union (EVROPSKÁ UNIE) and the Ministry of Education, Youth and Sports (MŠMT) of the Czech Republic, along with the text "Vzdělávání" and "Evropským sociálním fondem a státním rozpočtem České republiky".

Osnova

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**
- Modelové organismy
 - *Mus musculus*
 - *Arabidopsis thaliana*



EVROPSKÁ UNIE

esf



MINISTERSTVO ŠKOLSTVÍ,
MLÁDEŽE A TĚLOVÝCHOVY



OP Vzdělávání
pro konkurenceschopnost



UNIVERSITA
MASARYKŮV
PRAHA

INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Arabidopsis thaliana

huseníček polní, mouse-ear cress

- malé nároky na kultivační plochu
- velké množství semen (20.000/rostlinu a více)
- malý a kompaktní genom, (125 MBp, cca 25.000 genů, prům. velikost 3 kb)
- 5 chromozomů
- vhodná pro široké spektrum fyziologických experimentů
- velká přirozená variabilita (cca 750 ekotypů (Nottingham Arabidopsis Seed Stock Centre))



Columbia 0

Landsberg 0

Wassilewskija 0

<http://seeds.nottingham.ac.uk/>



INVESTICE DO F

EVROPSKÁ UNIE
a státním rozpočtem České republiky

Arabidopsis thaliana huseníček polní, mouse-ear cress

- Genom známý od roku 2000 (<http://www.arabidopsis.org/>)

The screenshot shows the homepage of the Arabidopsis Information Resource (TAIR) website. The page features a navigation menu with options like Home, Help, Contact, About Us, and Login/Register. Below the navigation, there is a search bar and a main content area with several sections:

- The Arabidopsis Information Resource:** A paragraph describing the database's scope, including genome sequence, gene structures, and gene product information.
- Breaking News:** A section with links to subscribe to a news feed, follow on Twitter, and join a Facebook group.
- New Set of Confirmed T-DNA Lines Available:** A notice dated November 28, 2012, regarding the availability of new T-DNA lines.
- New from ABRC Education and Outreach:** A notice dated October 31, 2012, about a re-designed education and outreach website.
- 2012 MASRC Report Now Available:** A notice dated July 11, 2012, about a report from the Multinational Arabidopsis Steering Committee.
- New Protein Chip and Cell Cultures at ABRC:** A notice dated May 9, 2012, about a new protein chip and cell cultures.

At the bottom of the page, there is a call to action: "Click here to try our new online submission form and submit the molecular function (e.g. protein kinase), biological process (e.g. seed development), localization (e.g. plasma membrane) or interacting partner of your favorite gene".



Osnova

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**
- Modelové organismy
 - *Mus musculus*
 - *Arabidopsis thaliana*
- Vybrané **metody molekulární biologie**
 - Příprava transgenních organismů



EVROPSKÁ UNIE



MINISTERSTVO ŠKOLSTVÍ,
MLÁDEŽE A TĚLOVÝCHOVY



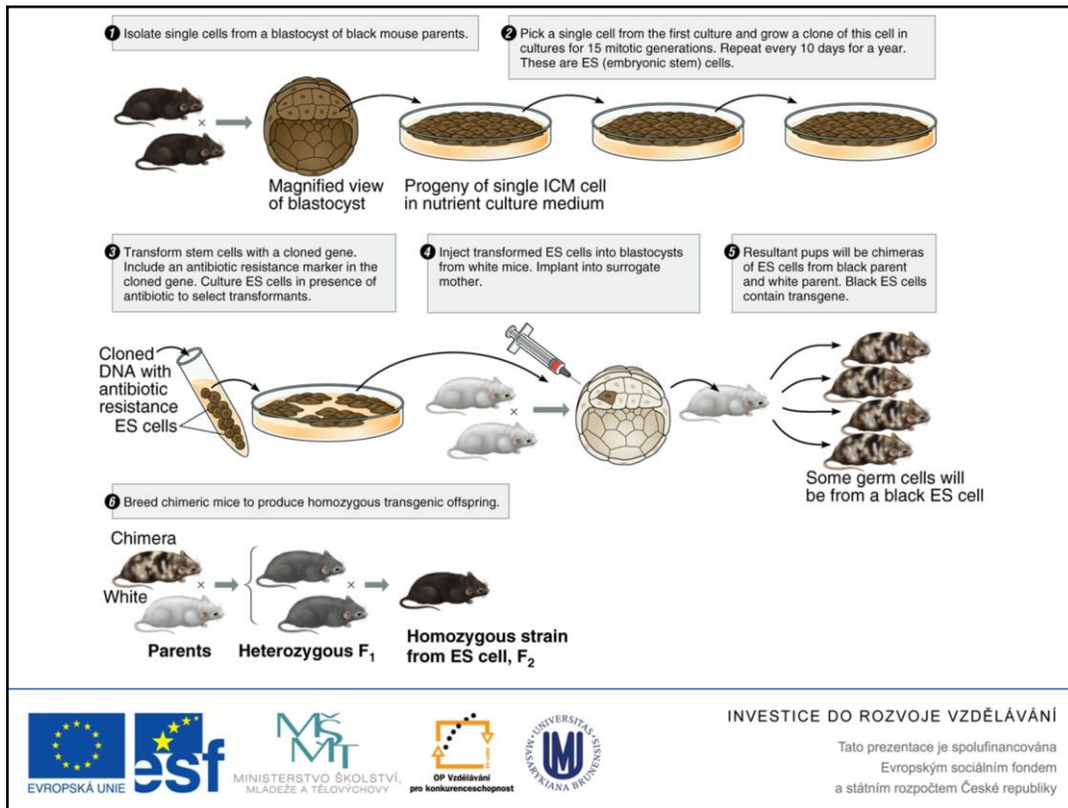
OP Vzdělávání
pro konkurenceschopnost



UNIVERSITA
MASARYKŮV
PRAHA

INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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



Individual ICM cells of the embryo could be isolated and later re-introduced into the new embryo. These ICM cells are called **embryonic stem (ES) cells**. It is a very important technique that allows production of transgenic mice.

The isolated ES cells are transformed via foreign DNA construct and it is injected within the embryo. The transformed cell becomes a part of the embryo and might result in the formation of different tissue types, among them the spermatogonia or oogonia, i.e. the tissue that provides progenitor for sperm or egg cells in the resulting chimera. Thus, the progeny of those chimeras will inherit the modified cell with certain probability and these individuals will carry the transgene in every cell of their body. Thus, the transgenic mice will be produced.

This is very important mainly with regard to the knockout mutant (K.O.) production. In the modified ES, the genes might be specifically eliminated via DNA recombination. In that way, the function of many of the mouse genes was identified.

E.g. the gene *NODAL* is expressed in the anterior portion of the primitive streak that is equivalent to the Hensen's node. *nodal/nodal* embryos are lethal; they do not undergo gastrulation and form almost no mesoderm.

**Transformace *Arabidopsis* prostřednictvím
*Agrobacteria tumefaciens***



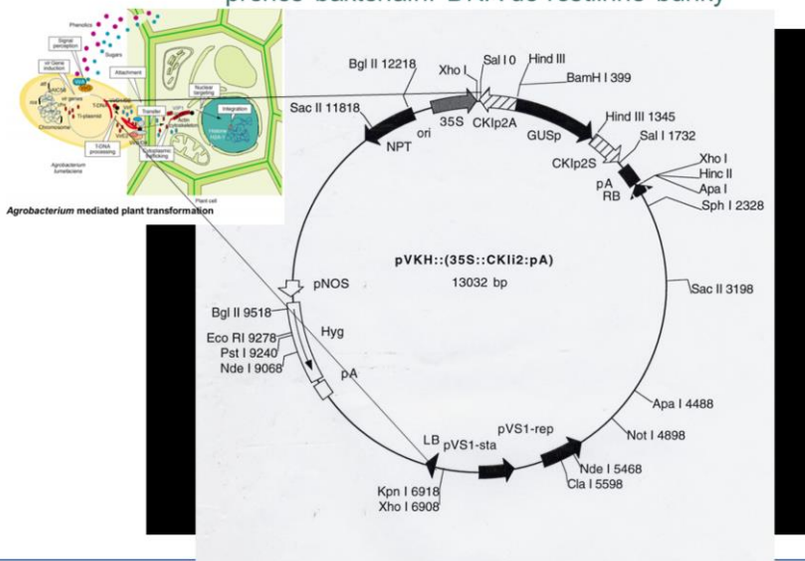
Crown gall of raspberry caused by *Agrobacterium tumefaciens*.



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Transformace *Arabidopsis* prostřednictvím *Agrobacterium tumefaciens* přenos bakteriální DNA do rostlinné buňky



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Transformace kokultivací listových disků



a Non-transgenic plant in a Murashige-Skoog (MS) medium. Excise leaf explants for co-cultivation.

b Excising a leaf from an Arabidopsis thaliana plant. Done in a laminar flow hood.

c Punching leaf discs. Done with sterile instrument in a laminar flow hood.

d Overnight (customer-specific) culture of Agrobacterium tumefaciens.

e Leaf discs floated on Agrobacterium suspension.

f Leaf discs discolored on culture induction media (2 days after co-cultivation). Photo taken 2 days after plating. Some bacterial growth (yellow) on surface of discs.

g Callus production on leaf disc explants. Some primary explants still show bacterial growth on the induction media.

h Discs discolored on leaf disc explants. Done in the presence of a selective agent.

i Excised discs inserted into root-induction media (RIM).

j Roots beginning to form on excised discs. Done in the presence of a selective agent.

k Extensive root production on excised discs.

l Fully regenerated transgenic plants growing in a Murashige-Skoog medium.



EVROPSKÁ UNIE



MINISTERSTVO ŠKOLSTVÍ, MLÁDEŽE A TĚLOVÝCHOVY

pro konkurenceschopnost

ANNA BR

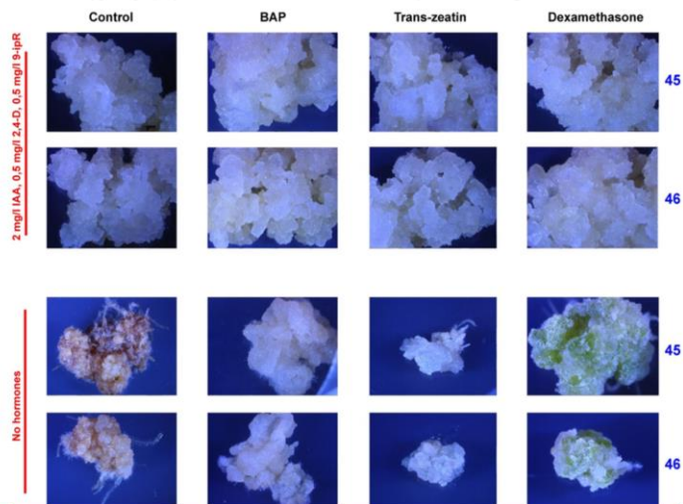
ÁVÁNÍ

ncována

i fondem

a státním rozpočtem České republiky

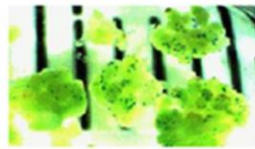
Transformace kokultivací kalusů



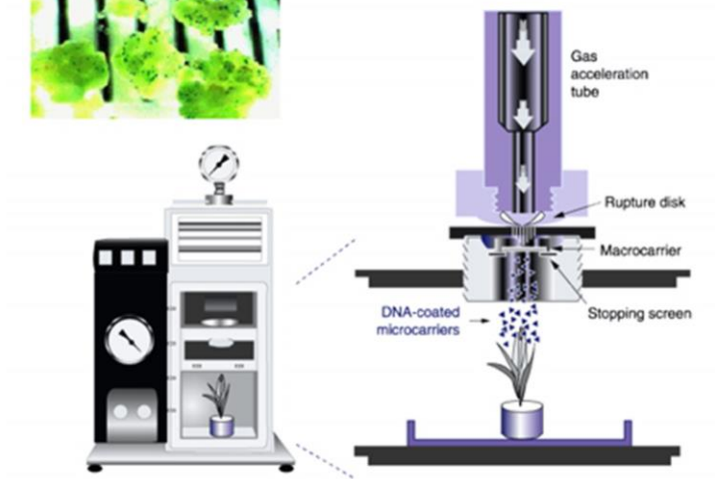
INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Transformace „nastřelováním“ DNA



Biolistic delivery of DNA



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Transformace květenství



When plants have primary bolts 5-15 cm they are ready to infiltrate. Clipping of primary bolts is not necessary.



After infiltration, pots are placed on their sides to allow for drainage and are covered with plastic wrap. Plants are returned, in this state, to the growth chamber for 24 hours. After 24 hours, they are turned upright into a fresh flat.



Plants are allowed to grow to maturity. They are staked to avoid seed loss and facilitate plant harvesting.
NOTE: Leaves degenerate within 2 weeks of infiltration. This is normal and does not affect seed set.



After infiltration, pots are placed on their sides to allow for drainage and are covered with plastic wrap. Plants are returned, in this state, to the growth chamber for 24 hours. After 24 hours, they are turned upright into a fresh flat.

EVROPSKÝ FOND
<http://www.bch.msu.edu/pamgreen/green.htm>



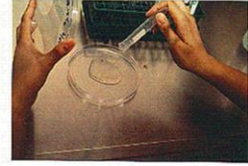
INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Transformace květenství



Sterilize seed in bleach solution.



Transfer seed to 100 µm mesh screen and wash.



<http://www.bch.msu.edu/pamgreen/green.htm>
medium (a 40mg/l kanamycin plate is shown)



Plant transformed seedlings in soil.

VANI
environmental
quality

Osnova

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**
- Modelové organismy
 - *Mus musculus*
 - *Arabidopsis thaliana*
- Vybrané **metody molekulární biologie**
 - Příprava transgenních organismů
 - PCR



EVROPSKÁ UNIE



MINISTERSTVO ŠKOLSTVÍ,
MLÁDEŽE A TĚLOVÝCHOVY



NIBB
NÁRODNÍ ÚSTAV
PRO BIOTECHNOLOGICKÉ VÝZKUMY



UNIVERSITA
MASARYKŮV
PRAHA

INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

PCR



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Osnova

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**
- Modelové organismy
 - *Mus musculus*
 - *Arabidopsis thaliana*
- Vybrané **metody molekulární biologie**
 - Příprava transgenních organismů
 - PCR
 - Design a příprava primerů (Dr. Hana Konečná)



EVROPSKÁ UNIE

esf



MINISTERSTVO ŠKOLSTVÍ,
MLÁDEŽE A TĚLOVÝCHOVY



ÚIV
NÁRODNÍ ÚSTAV
PRO VĚDECKÝ VÝZKUM V OBLASTI
VĚDEK O VĚDOVÉ ČINNOSTI



UNIVERSITA
MASARYK
UNIVERSITY

INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Shrnutí

- Nástroje **systemové biologie**
 - Analýza **genové ontologie**
 - **Modelování molekulárních regulačních sítí**
- Modelové organismy
 - *Mus musculus*
 - *Arabidopsis thaliana*
- Vybrané **metody molekulární biologie**
 - Příprava transgenních organismů
 - PCR
 - Design a příprava primerů (Dr. Hana Konečná)



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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

Diskuse



INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

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