



TILLING (Targeting Induced Local Lesions IN Genomes)

Richard Thompson, Christine Le Signor, Myriam Sanchez, Brigitte Darchy,
Gregoire G. Aubert, Karine Gallardo, Christine Saffray, Marion Dalmais,
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TILLING (Targeting Induced Local Lesions IN Genomes)

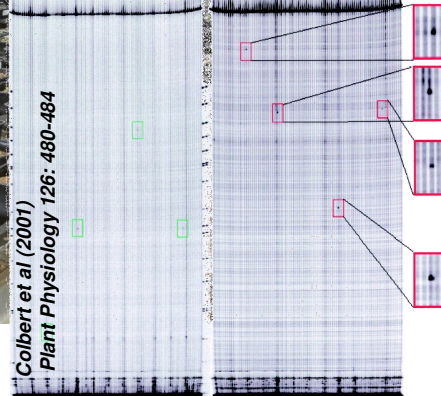
Christine Le Signor, Myriam Sanchez, Brigitte Darchy, Grégoire Aubert, Karine Gallardo, Richard Thompson (INRA-UMR Agroécologie, Dijon)

Christine Saffray, Marion Dalmais, Abdelhafid Bendahmane (IPS Paris-Saclay)

TILLING: A method of producing and identifying mutations in any gene of interest

- EMS-generated population of point mutants
- Make DNAs from each plant (~4500), arrange in pools
- Screen for lines of interest by gene-specific detection
- Purify away from other mutations by back-crossing

Photo serre URLEG-INRA Dijon C. Le Signor



TILLING

Targeting Induced Local Lesions in Genomes

Photo serre URLEG-INRA Dijon C. Le Signor

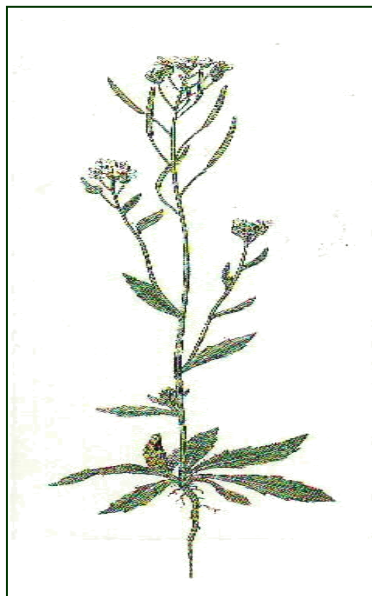


Method of producing
and identifying
mutations in genes of
interest

Origin of TILLING

Henikoff's lab

Basic Sciences Division, Fred Hutchinson Cancer Research Center, Seattle, USA.



Arabidopsis
(génome 125 Mb)

Targeted screening for induced mutations.

McCallum et al.

Nat Biotechnol. 2000; 18(4): 455-7

High-throughput TILLING for functional genomics.

Till BJ et al.

Methods Mol Biol. 2003; 236:205-20.

Application of TILLING

Cereals:



Rice

Génome de
500 Mb



Maize

2500 Mb

 *Crop Pathology and Genetics Research*
UC- Davis, USA

 *Henikoff's lab*
Seattle, USA

Legumes:



Lotus

470 Mb



Medicago
Jemalong A17

520 Mb



Pois

4000 Mb

 *John Innes Centre*
Norwich, UK

 *D. Cook's lab*
UC Davis, USA

 *INRA Dijon*
France

 *INRA Dijon et URGV- INRA*
Evry France

Advantages of TILLING:



Applicable to all organisms (small or large genomes)

if utilises a chemical mutagen which generates an allelic series of point mutations (mutation frequency 1 mutation/300 Kb).



Easily set up:

if Rapid screening of a collection of mutagenized plants for the identification of point mutations in genes of interest

Steps in TILLING procedure

 **Mutagenesis of genome using Ethyl Methane Sulfonate (EMS)**

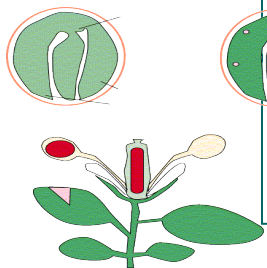
 **Production of an M2 population**

 **Screening of the M2 population**

 **Isolation of lines bearing mutations in gene of interest and back-crossing to eliminate unwanted mutations**

Assessment of mutation efficiency

M1



Chimaera

All the mutations are in the heterozygous state

Mutation efficiency:

Percentage of embryos aborted
in pods of M1 plants ~15 à 20

%

1

M2



The plant is has a
mutation of interest

M2 plant carrying one or several mutations in its genome

50% mutations heterozygote

25% mutations homozygote

25% homozygote wild-type

Recessive mutations (99%) -
no phenotype associated

For recessive mutations: 25% of plants show
phenotype

For dominant mutations: 75% of plants show
phenotype



(2003) Genetics 164: 731-740

Spectrum of Chemically Induced Mutations From a Large-Scale Reverse-Genetic Screen in Arabidopsis

Elizabeth A. Greene,* Christine A. Codomo,* Nicholas E. Taylor,* Jorja G. Henikoff,*
Bradley J. Till,* Steven H. Reynolds,* Linda C. Eman,* Chris Bartner,* Jessica E. Johnson,†
Anthony R. Odden,* Luca Comai,* and Steven Henikoff*,‡,§

*Fred Hutchinson Cancer Research Center and †Howard Hughes Medical Institute, Seattle, Washington 98109 and ‡Department of Biology,
University of Washington, Seattle, Washington 98195
Manuscript received January 18, 2003
Accepted for publication March 26, 2003

ABSTRACT
Chemical mutagenesis has been a valuable tool of traditional genetics, but it has not been possible to determine the underlying rate and distribution of mutations from phenotypic screens. Here we report a large-scale reverse-genetic screen for recessive mutations in the *Arabidopsis thaliana* target genes from a large-scale TILLING reverse-genetic project, about two orders of magnitude larger than previous screens. We find that the distribution of mutations is highly skewed, with the highest frequency of mutations occurring at the first position of codons. This reflects the occurrence and randomness of chemically induced mutations. We provide evidence that we have detected all mutations in the target genes. We also show that the distribution of mutations is highly throughput TILLING method; therefore, any deviations from randomness can be attributed to selection or mutational biases. Our data detect twice as many heterozygotes as homozygotes, as expected; however, for most genes, the predicted ratio of heterozygotes to homozygotes is 1.6, indicating a 6% reduction of heterozygotes. An overview of the distribution of mutations by EMS.

Mutation frequency: 1 mutation/300 kb (M2)

For a genome size similar to *M. truncatula* (500 Mb), the frequency would be ~1500 mutations/plant M2)

Therefore backcrosses to WT line needed

Mutation distribution:

TABLE I
Distribution of missense and truncation mutations in heterozygotes and homozygotes

	All	Silent	Missense	Truncation
% Distribution	1890	851	946	93
% expected	100	44.4	48.3	5.3
% observed	1276	566	637	49
Heterozygous	1276	566	637	73
Homozygous	614	285	309	20
Ratio	2.08	1.99	2.06	3.6 ($P \leq 0.05$)

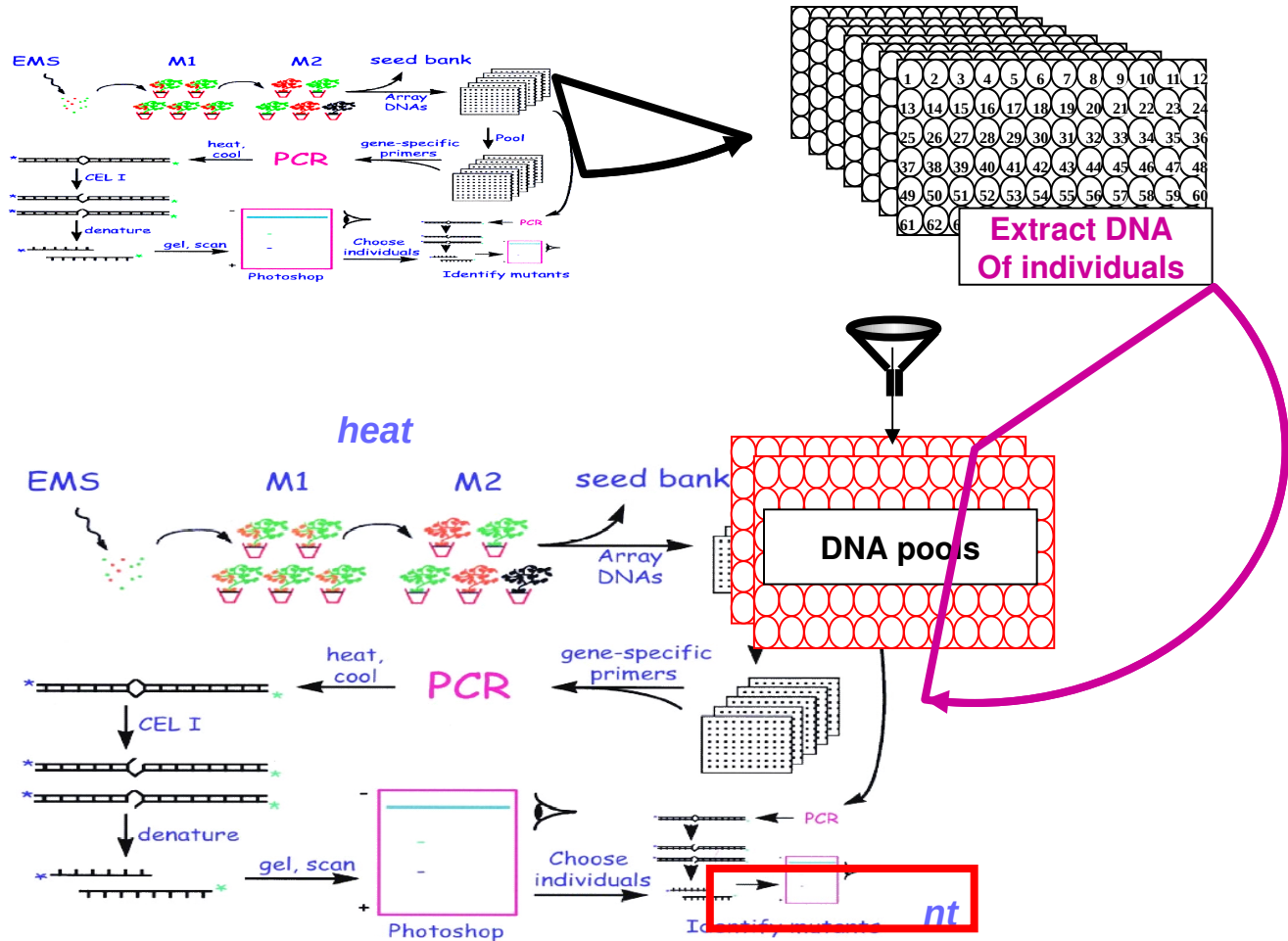
caused by seed contamination of the TILLED population

Silent mutations : 45%
Missense mutations : 50%
Stop codon Mutations: 4,9%

Therefore need to isolate several mutants for each gene

Summary of TILLING method

From Colbert et al. (2001). Plant Physiology, 126: 480-484.



Screening carried out on DNA pools from 8-12 individuals on microtitre plates and subsequently on individual DNA samples.

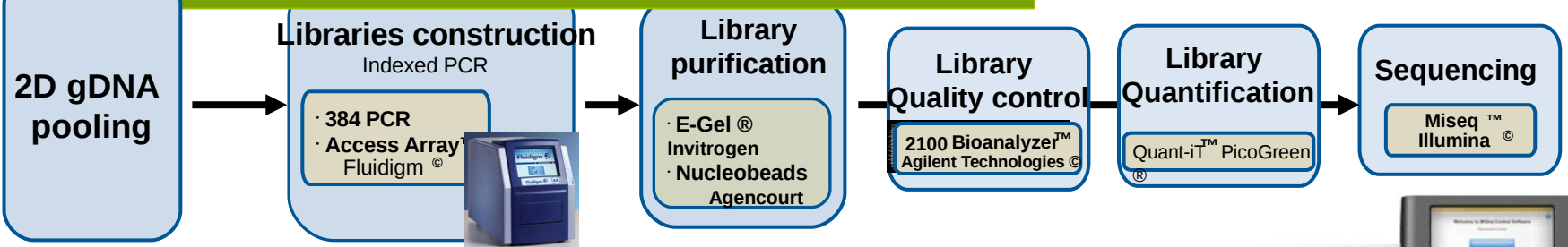


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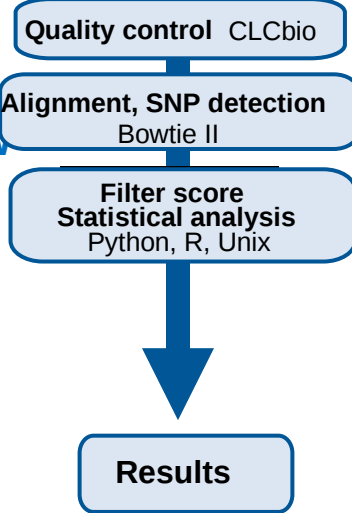


Towards a high throughput screening based on NGS (URGV Paris-Saclay)

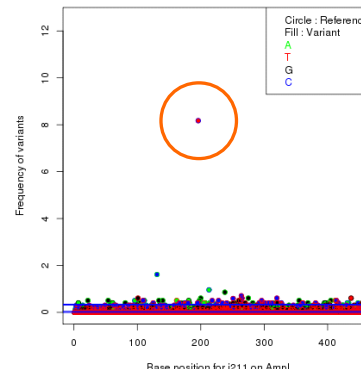
2013 : MiSeq detection system – general workflow



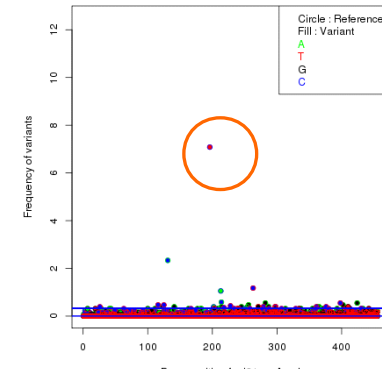
NGS data analysis : Bioinformatic workflow



Output in
1st pooling dimension

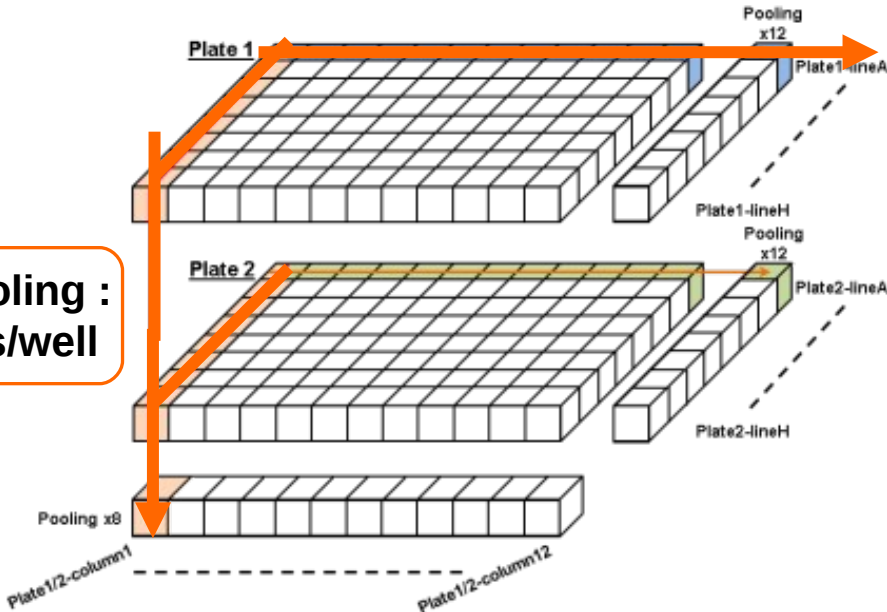


Output in
2nd pooling dimension





Column pooling :
16 families/well



Row pooling :
12 families/well

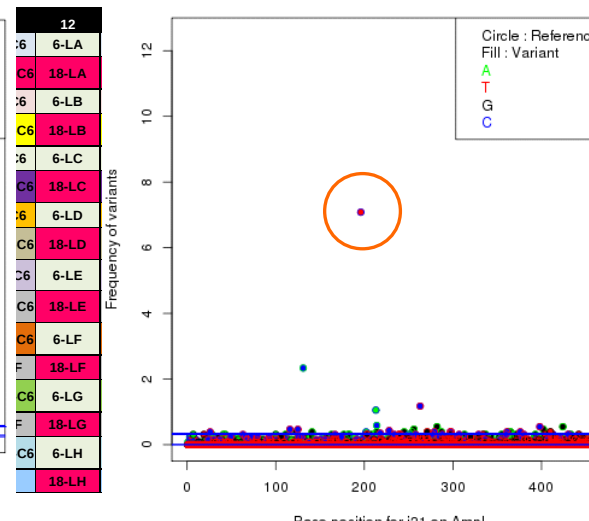
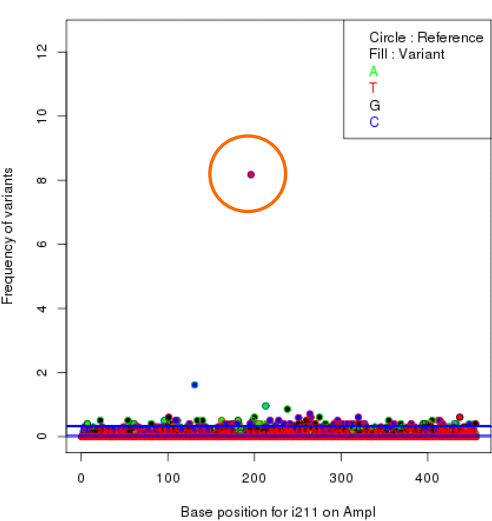
1 family = 2 coordinates

384 well-plate gDNA :

First pooling dimension

Second pooling dimension

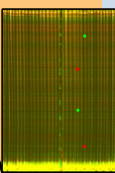
	1	2	3
A	1_2-C1	1-LA	1-LB
B	17_18-C1	13-LA	17-LB
C	3_4-C1	1-LB	3-LA
D	19_20-C1	13-LB	19-LA
E	5_6-C1	1-LC	5-LB
F	21_22-C1	13-LC	21-LB
G	7_8-C1	1-LD	7-LA
H	23_24-C1	13-LD	23-LB
I	9_10-C1	1-LE	9-LA
J	25_26-C1	13-LE	25-LB
K	11_12-C1	1-LF	11-LA
L	25-LA	13-LF	25-LB
M	13_14-C1	1-LG	13-LA
N	26-LA	13-LG	26-LB
O	15_16-C1	1-LH	15-LA
P	WT	13-LH	16-LB



	22	23	24
11-LA	1_2-C12	12-LA	
23-LA	17_18-C12	24-LA	
11-LB	3_4-C12	12-LB	
23-LB	19_20-C12	24-LB	
11-LC	5_6-C12	12-LC	
23-LC	21_22-C12	24-LC	
11-LD	7_8-C12	12-LE	
23-LD	23_24-C12	24-LD	
11-LE	9_10-C12	12-LE	
23-LE	25_26-C12	24-LE	
11-LF	11_12-C12	12-LF	
23-LF		24-LF	
11-LG	13_14-C12	12-LG	
23-LG		24-LG	
11-LH	15_16-C12	12-LH	
23-LH		24-LH	



Cel1/Endo1 and NGS TILLING methods compared (URGV Paris-Saclay)

	Cel1/Endo 1	NGS- MiSeq - Illumina	
		PCR 384	Fluidigm
DNA pooling	1D	2D (rows + columns pooling)	
Amplicon	1000 bp	500 bp	500 bp
Detection system	<ul style="list-style-type: none">· Endonuclease digestion· Detection of labelled DNA fragments on Li-Cor· Sanger sequencing		<p>Pair-end sequencing 2 x 250bp □ 2 x 300bp</p> <p>Sequence analysis</p> 
Amplicon/run	1	25	48
Time/ experiment	1 month	3 months	2-3 months
Mutation frequency	1 mutant/200 kb	1 mutant/130 kb	1 mutant/130 kb
Families / screen	5000	384-well plate = 2500 families	384-well plate = 2500 families



Pea TILLING populations

CAMEOR
spring pea collection

Terese *rms4*
Spring pea collection

4704 M2 families
60 genes screened (80 targets)
1150 mutants
NGS: 1 mutation / 120kb

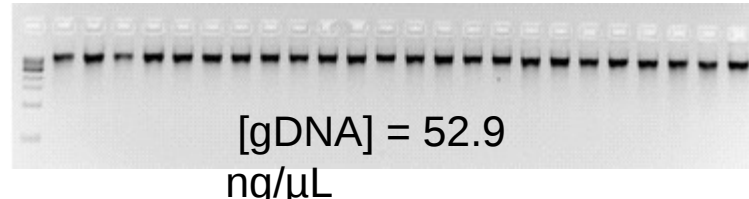
4200 M2 families
10 genes screened
84 mutants
NGS: 1 mutation / 359kb

Dalmais et al., Genome Biology, 2008
<http://urgv.evry.inra.fr/UTILLdb>)

Hr Ps336/11 genotype
PEAMUST Winter pea collection

- ➡ 2014 : DNA samples for 946 low-seeds M2 families
- ➡ 2015 : DNA samples for 960 high-seeds M2 families
- ➡ 2016-2017 : end of population production

4500 M2 families





Variation in mutation frequency within treated seed lot

→ Comparison of mutation frequency in low- and high-yield families

Hr Ps336/11 genotype

**PEAMUST Winter pea
collection**

- ↳ 2014 : DNA samples for 946 low-seeds M2 families (1-7 seeds/plant)
- ↳ 2015 : DNA samples for 960 high-seeds M2 families (20-100 seeds/plant)

→ Mutation frequency
1/72 kb

→ Mutation frequency
1/334 kb

- Low-yield families had a mutation frequency 4-fold higher, but probability of losing the mutant allele also high → aim for intermediate mutation frequency



Recovery of mutant line for phenotyping

- TILLING lines contain 100s-1000 or more induced point mutations, so the mutation of interest should be backcrossed to a non-mutagenized parental line (>2 generations)
- Mutant allele is followed using a genotyping method such as dCAPS
- If two different mutant alleles available, an allelism test can help associate mutation and an eventual phenotype





Exploitation of TILLING via new genomics resources in Pea

The Pea Gene Atlas <http://bios.dijon.inra.fr/FATAL/cgi/pscam.cgi>

Fichier Édition Affichage Historique Marque-pages Outils 2

Débuter avec Firefox Dictionnaire de Cuisine et Gastronomie - ... jburstin @ Pisum sativum Cameor +

debs.dijon.inra.fr/FATAL/cgi/pscam.cgi?_wb_url=4edoc06edoc06edoc00128edoc558edoc5C28edoc3edoc528edoc328edocC28edocX048edoc328edocC28edocPT68328ec gousse pois

Les plus visités Hotmail Personnaliser les liens Windows Media Windows

jburstin @ Pisum sativum Cameor

INRA Lipm UMP Ecog

Peptides Clusters Fasta Blast PatScan RNA-seq@BIOS

PsCam008801_1_AA

Permalink : http://bios.dijon.inra.fr/FATAL/cgi/pscam.cgi?peptideid=PsCam008801_1_AA

		Get Fasta Seq	MultAliN Multalin		
--	--	---------------	-----------------------------	--	--

ID PsCam008801_1_AA
AC PsCam008801_1_AA
IP Auxin efflux carrier
EC
GO Biological Process: transmembrane transport (GO:0055085)
Cellular Component: integral to membrane (GO:0016021)
CC
LN 352 aa

....10.....20.....30.....40.....50
MISLANVFHRVTTKTVPLVTFMLAYVSIXWFKLFTQE~~QCSGINKFVAKF'S~~
IPLLSPFIQISSNNIYKMSLKLHMADF~~IQKLLAFLILLTAIIKRRGGGLKW~~
ITIGLSLSTPLNTLII~~GIPVPLKAMYND~~EAVVLLAQFWFLQS~~MNWYNLLLF~~
~~TKEPPDAAKTILSAPPSETAESIQSKEEEEDDEESVGTGRKHKLYPILVTVG~~
~~KKLIRNPNTFAASLLGIWSSIHFRWGTHMPVEVNQSIELLSNHGLGMF~~
~~SIGLFMHSQSSIAICGARNTHMVAIGLKVLVGPALMASLSIVIGLRNLTFK~~
~~VAIVQALPQGIVPFVFAKEYNHWPSTLSTAIIGLMILALPVELAFYELL~~
AL

Domain decomposition

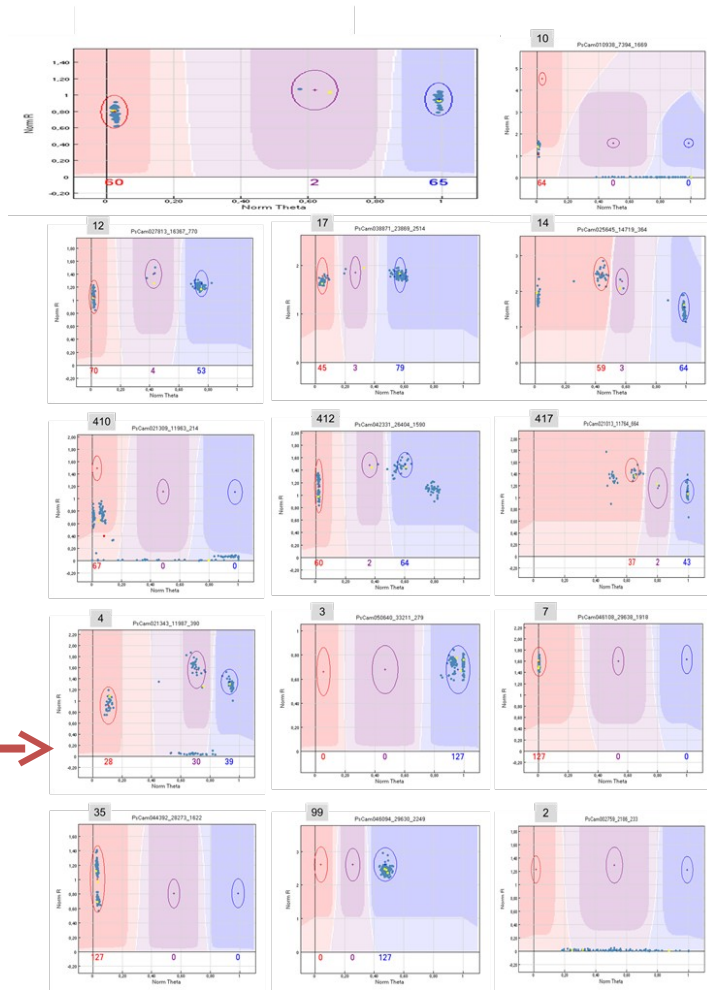
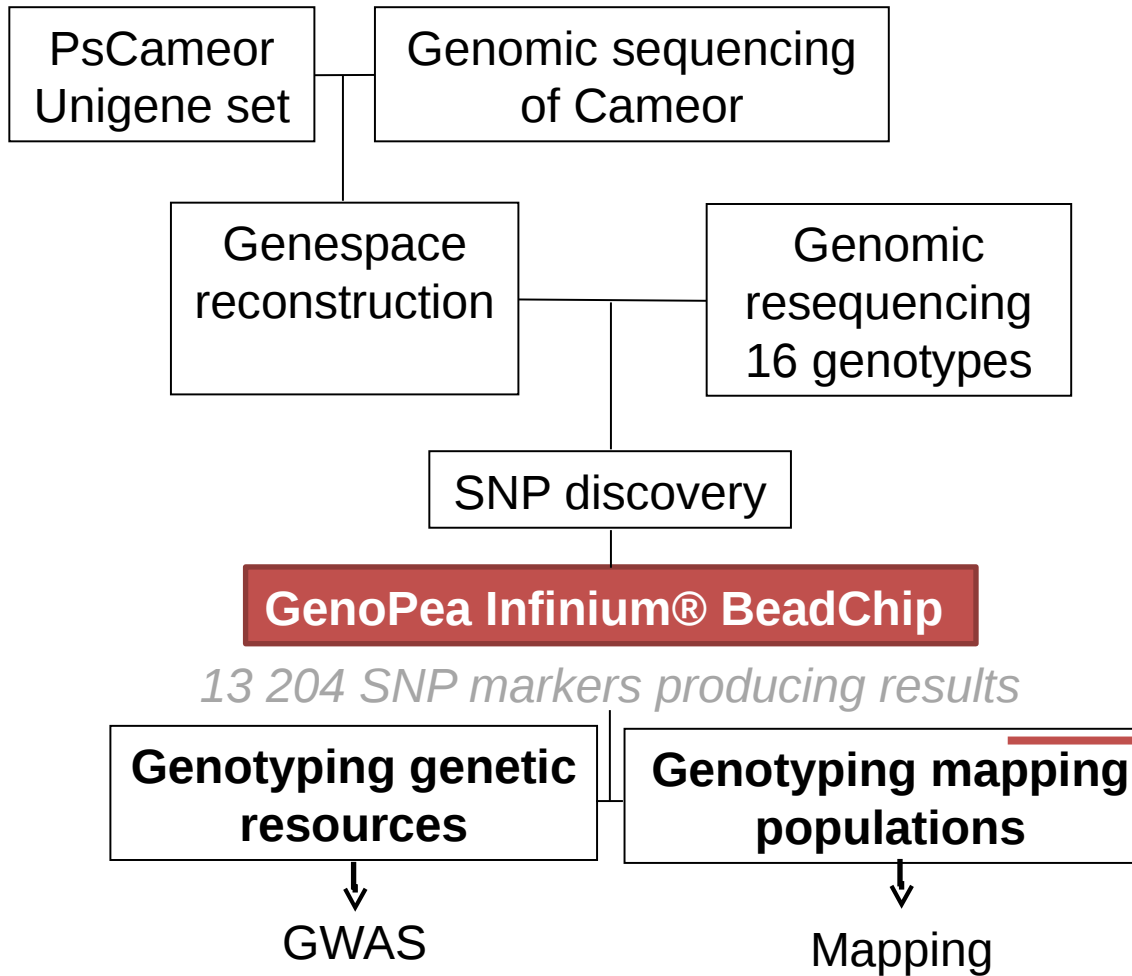
PsCam016724_1_AA PsCam016724_1_AA 1 352 S=1290 I=1290 E=6.52466e-149 IPR004776:Auxin efflux carrier; pscam

PsCam07118_1_AA 4 250 S=650 I=2914 E=1.22222e-149 IPR004776:Auxin efflux carrier; pscam

démarrer Courier entrant... Brno-2012 Présentation T1 Cadrage_Journées... jburstin @ Pisum s... FR 17:16

Search any gene of interest

Exploitation of TILLING via new genomics resources in Pea



Aluome et al. submitted Tayeh et al. 2015, *The Plant J.*



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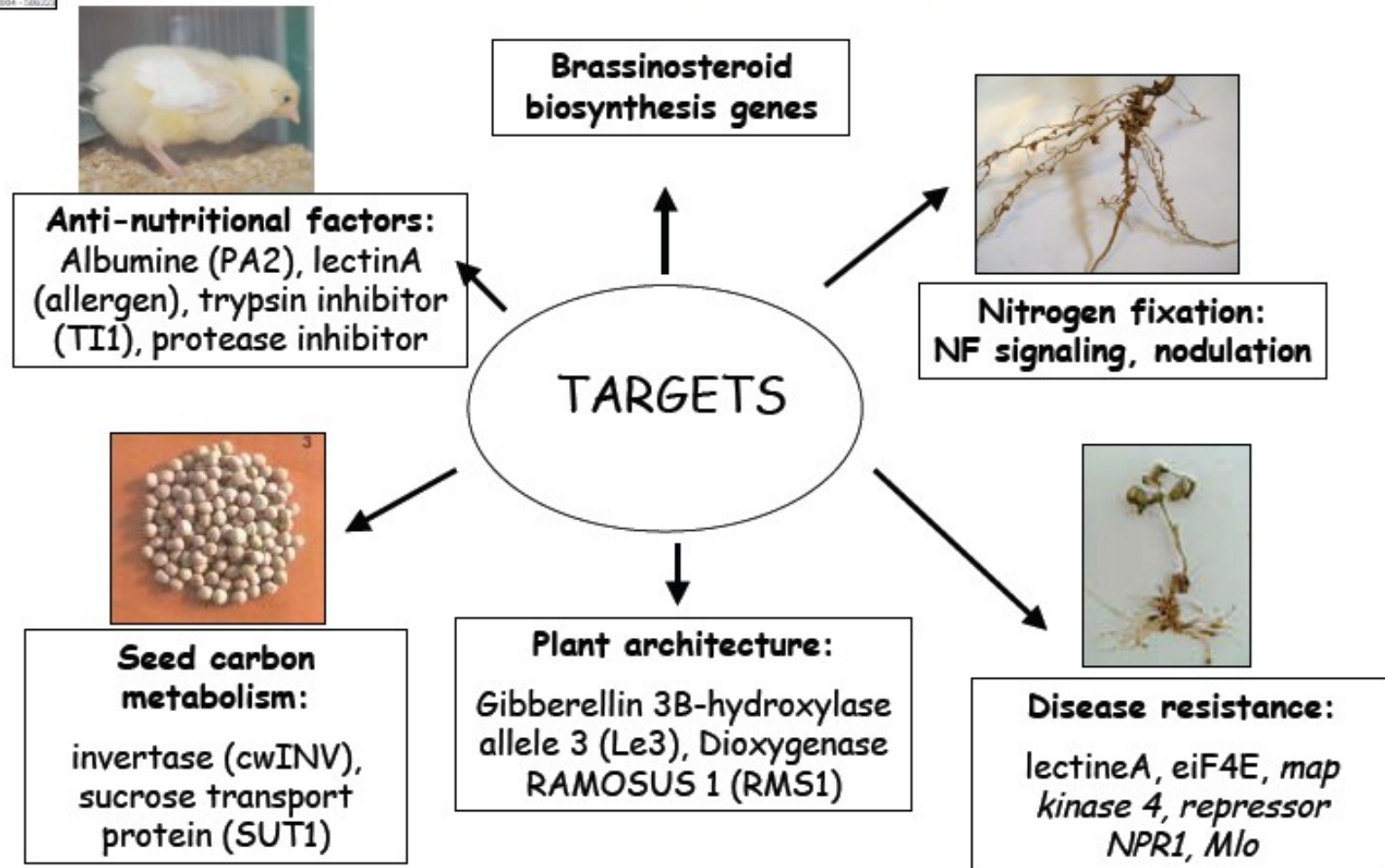
INRA

UNIB
UNIVERSITÉ
BOURGOGNE FRANCHE-COMTÉ

(Judith
Burstin et al.)

TILLING: Widely used in Pea since 2008

FOOD-CIT-2004-50022



Examples of TILLED genes in our project on seed development and composition

Model
M. truncatula



GWAS & Comparative QTL mapping

Candidate genes

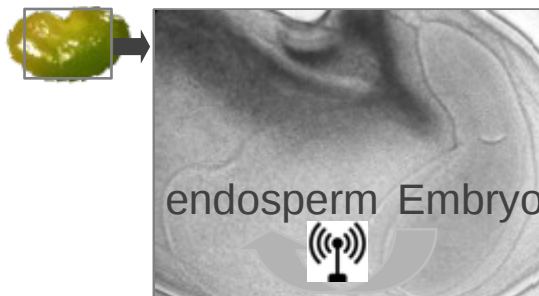
Crop
P. sativum



1-seed weight

Putative pea orthologs

Seed
development



Communication &
interaction ?

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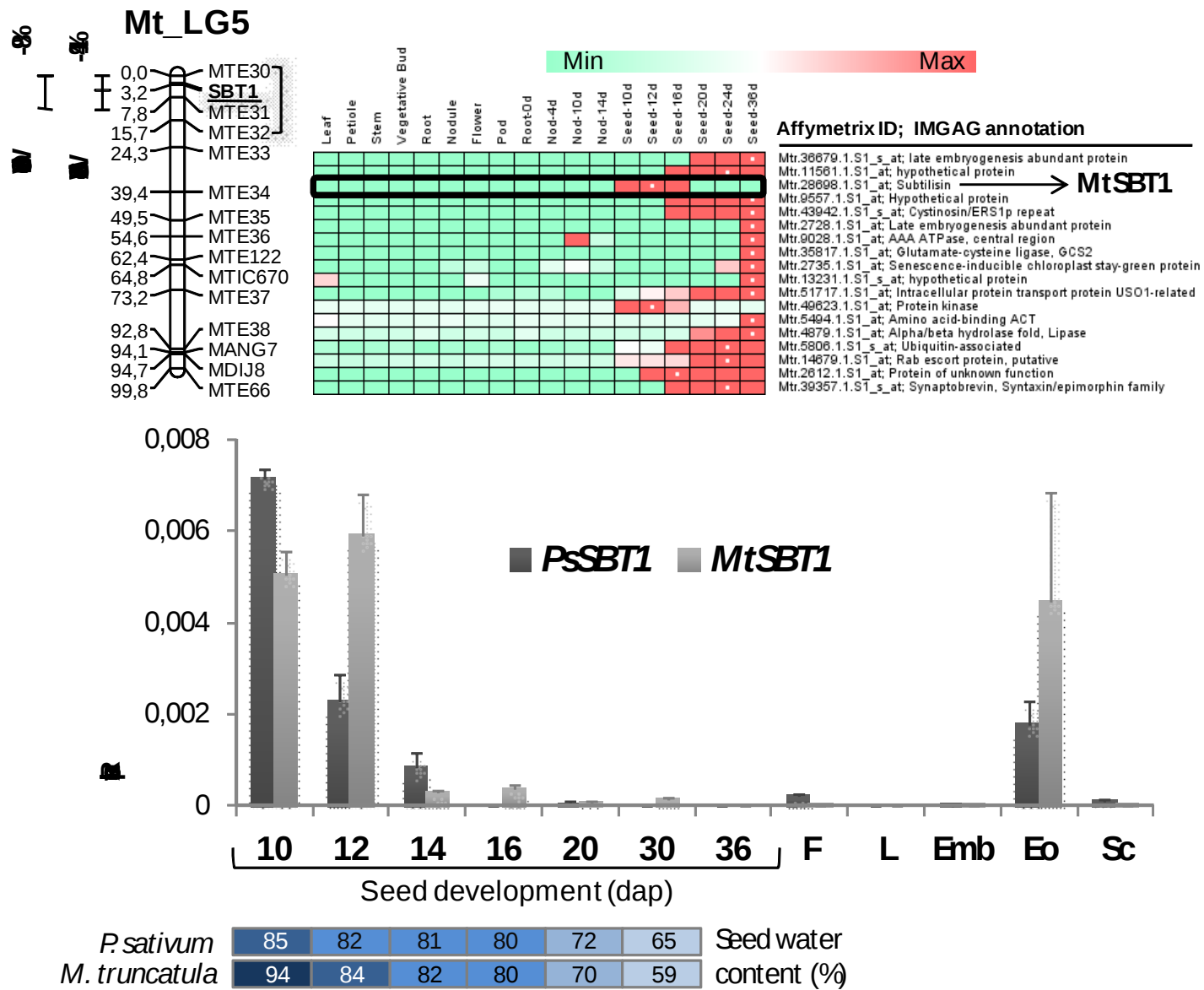
Noguero et al., 2013 Plant Sci.

Noguero et al. 2015 Plant J.

Two genes with an endosperm-specific expression during embryogenesis determine final seed weight through the regulation of embryo cell division :

- ↳ **a Subtilisin-like serine protease**  **Phytosulfokines ?**
D'Erfurth et al., 2012 New Phytol.
- ↳ **a DNA-binding with One zinc-Finger (DOF)-type transcription factor**

A gene encoding a seed-specific subtilisin colocalizes with a *Medicago* seed weight QTL and is expressed at the interval embryogenesis-seed filling

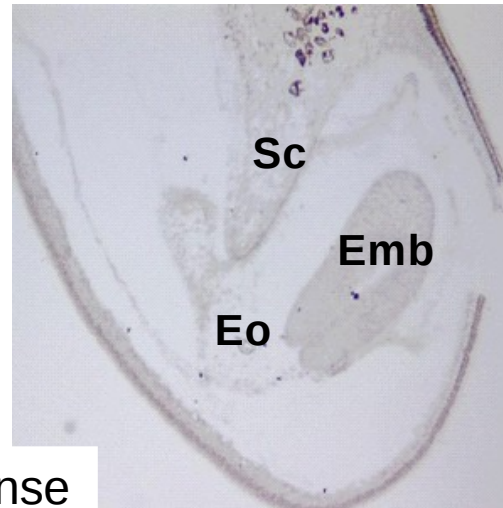


In situ hybridization of MtSBT1.1 to 10 and 12 DAP seeds

10 dap



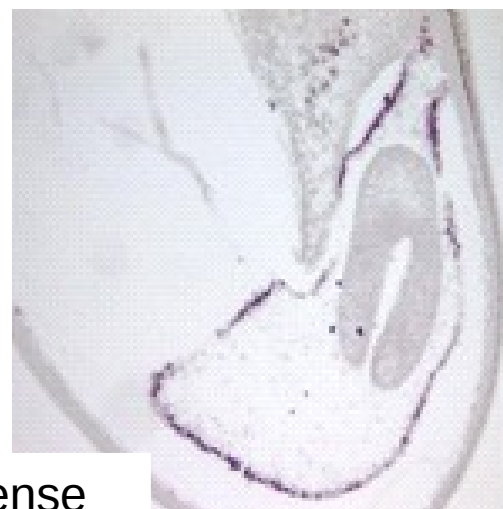
12 dap



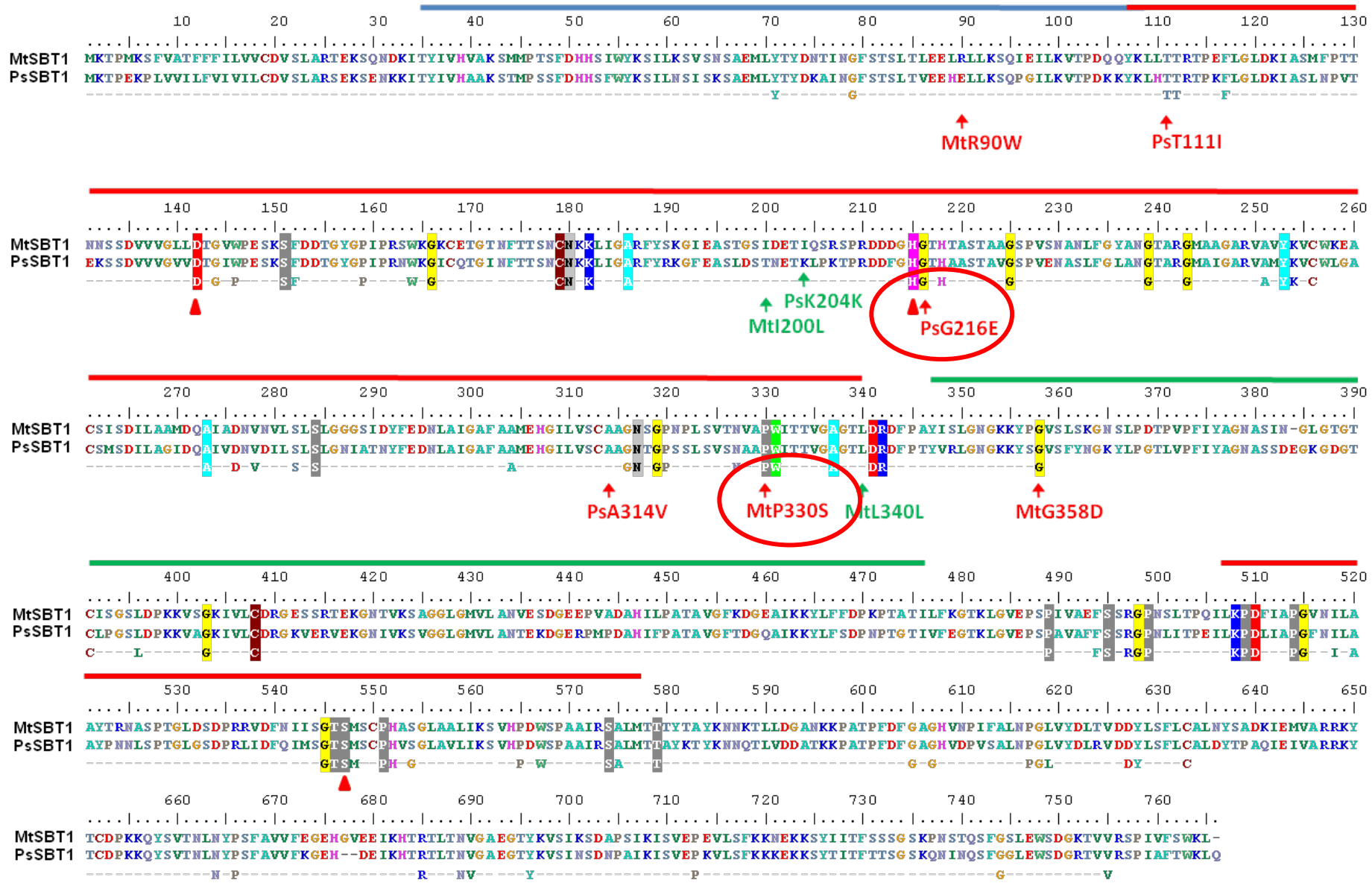
sense



antisense



Alignment of MtSBT1.1 AND PsSBT1.1 showing mutants available



Production de légumes secs en France

Surfaces de tontilles et de pois échappés en France



Production de légumes secs en France

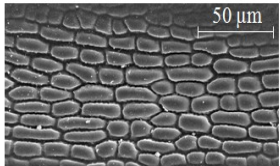
Surfaces de tontilles et de pois échappés en France



Surface area and number of cotyledon cells in mature seeds of the homozygous MtP330S mutant versus wild-type.



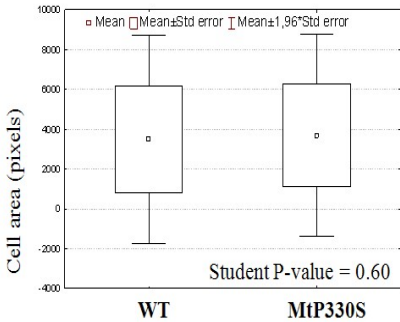
A



Genotype	WT	MtP330S
Mean epidermal cell surface (pixels)	1906,5	1918,8
Standard error	177,1	169,7

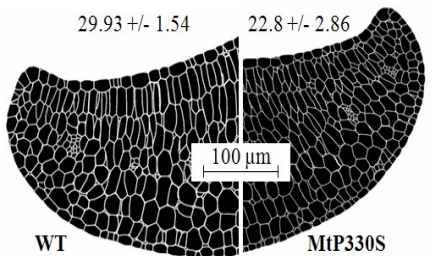
B

Mean area of cotyledon cells



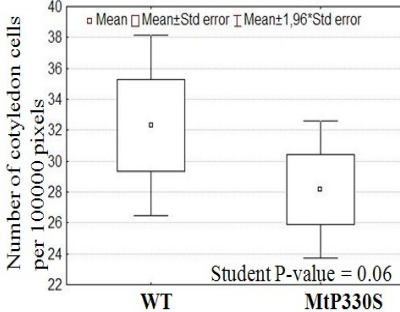
C

Number of cells per 100000 pixels in 2 blocs :



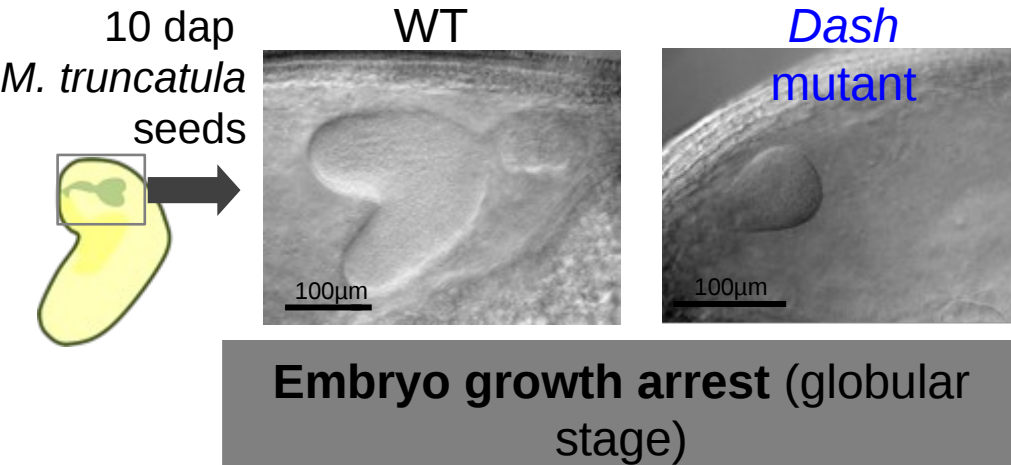
D

Mean cotyledon cell number



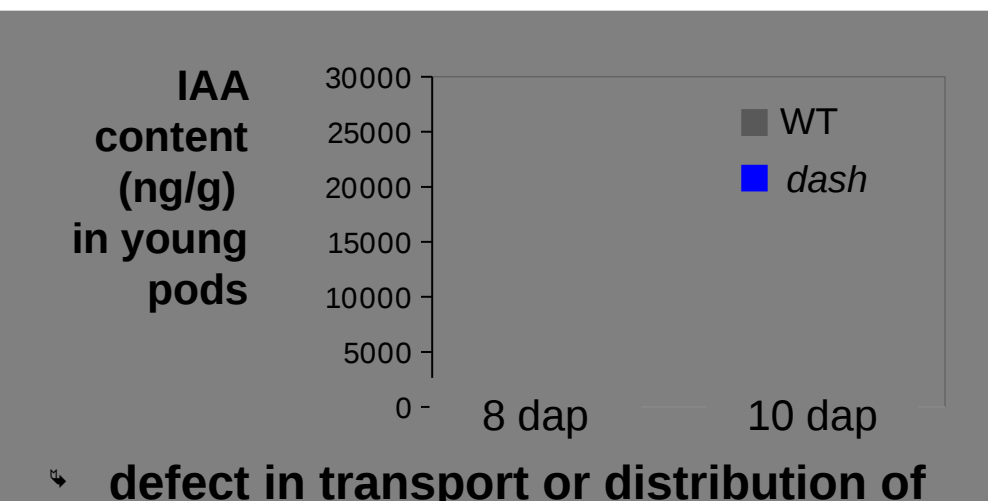
Endosperm-specific genes controlling seed quality

DASH, a DOF TF Acting in Seed embryogenesis and Hormone regulation



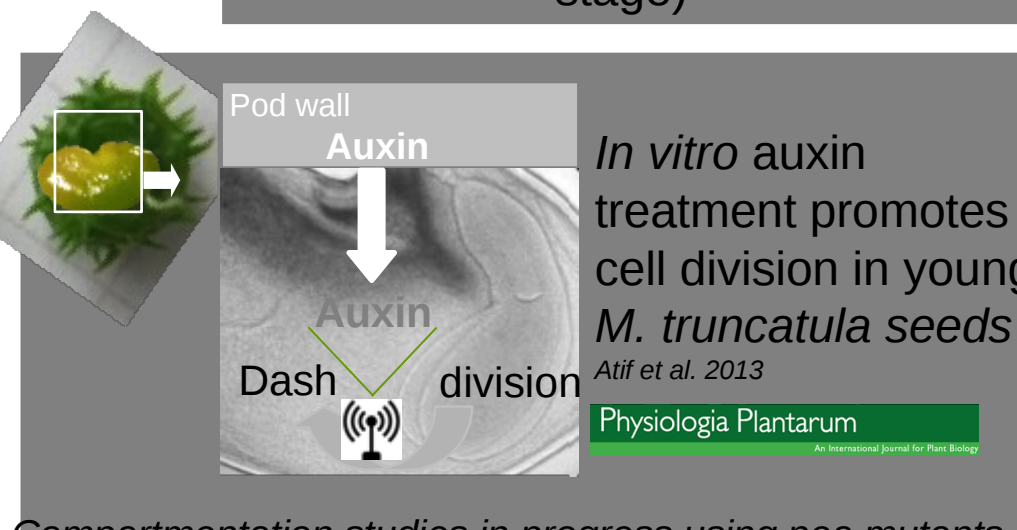
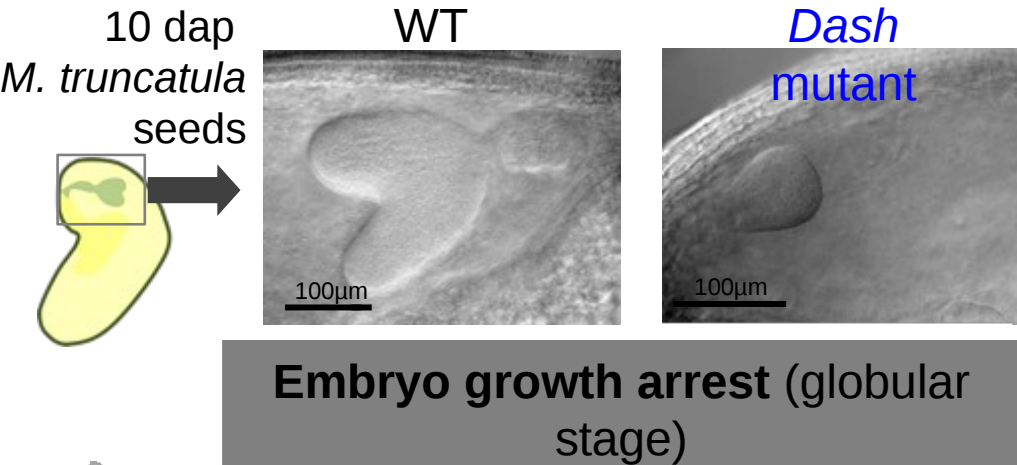
PageMan display of the functional classes down-regulated in the dof mutant

protein synthesis	-6,8	
hormone metabolism.auxin.induced-regulated-activated	-6,0	
hormone metabolism.auxin	-5,8	
cell	-5,4	
cell.cycle	-5,2	
DNA.synthesis/chromatin structure	-5,1	
cell.organisation	-3,4	



Auxin-transport genes
Auxin-responsive genes

Endosperm-specific genes controlling seed quality



Compartmentation studies in progress using pea mutants
the plant journal

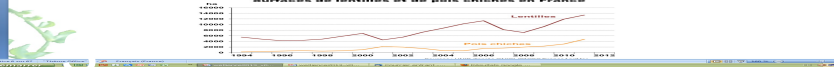


protein synthesis	-6,8	
hormone metabolism.auxin.induced-regulated-activated	-6,0	
hormone metabolism.auxin	-5,8	
cell	-5,4	
cell.cycle	-5,2	
DNA.synthesis/chromatin structure	-5,1	
cell.organisation	-3,4	

Auxin-transport genes
Auxin-responsive genes

Acknowledgements

Production de légumes secs en France



Melanie Noguero, Christine Lesignor, Brigitte Darchy, Greg Aubert,
Myriam Sanchez, Judith Burstin, Karine Gallardo, (AE, Dijon)
Jérôme Verdier (Shanghai Center for Plant Stress Biology)



C. Saffray, M. Dalmais,
A. Bendahmane (IPS Saclay)



Pea/Mtr TILLING: Publications arising

Resource development

- Triques, K., et al. (2008). "Mutation detection using ENDO1: Application to disease diagnostics in humans and TILLING and Eco-TILLING in plants." *BMC Molecular Biology* 9.
- LeSignor C., (2009) Optimizing TILLING populations for reverse genetics in *Medicago truncatula*. *Plant Biotechnology J.* 7 : 430-441
- Dalmais, M., UTILLdb, a *Pisum sativum* in silico forward and reverse genetics tool. *Genome Biol.* (2008) 9(2): R43.

Examples of Applications

- Hofer, J., L. Turner, et al. (2009). "Tendril-less Regulates Tendril Formation in Pea Leaves." *Plant Cell* 21(2): 420-428.
- Laura de Lorenzo, (2009) A Novel Plant Leucine-Rich Repeat Receptor Kinase Regulates the Response of *Medicago truncatula* Roots to Salt Stress *The Plant Cell* 21:668-680
- Plet et al, (2011) MtCRE1-dependent cytokinin signaling integrates bacterial and plant cues to coordinate symbiotic nodule organogenesis in *Medicago truncatula*. *Plant J.* 65: 622-633.
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