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Role of transcription factors in early male gametophyte development of *Arabidopsis*

Ph.D. Thesis

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Annotation: In the presented work the relationship between transcription factors and male gametophyte development was studied. The Ph.D. Thesis covers selection of candidate genes, wide-scale screening of T-DNA mutant lines and detailed analysis of a selected transcription factor on pollen development.

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Declaration of originality

The coauthors listed below fully acknowledge that David Reňák performed the major works in respective papers with the mention of his contribution.

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1. Cell biology of the male gametophyte

1.1. Stamen development

The developmental events leading to a stamen formation follows the pattern of floral meristem identification. Stamen primordia are initiated on the meristem surface within the third whorl as a result of specific floral homeotic MADS-box genes co-expression, including *APETALA3 (AP3)*, *PISLILLATA (PI)*, *AGAMOUS (AG)*, and *SEPALLATA1/2/3/4 (SEP1/2/3/4)* in *Arabidopsis*. Misexpression of participating transcriptional regulators is well known to produce messy floral architecture, for instance additional stamens replacing carpels in the neighbouring fourth whorl, if male part is considered in *superman* mutation (Bowman et al. 1992). Stamen primordia appear after the outer sepal and petal primordia have initiated but prior to the initiation of inner carpel primordia. Later, the stamen primordium prolongs and differentiates into two compartments, filament and anther. The filament stands as a holder carrying conducting tissue while the anther contains both nonreproductive and reproductive tissues that are responsible for the production and release of mature pollen grains (Scott et al. 2004).

The anther forms two bilaterally symmetrical pollen sacks with two loculi in each. Developmentally, the anther surrounding pollen sacks consists of several cellular layers originating from three layers called L1, L2, and L3 of the floral meristem that can be recognized histologically and functionally and that play a role in the anther differentiation process. The outermost L1 layer establishes the epidermis and stomium which is responsible for splitting of anther wall during dehiscence. Thus the programmed cell death (PCD) of the stomial cells is necessary for pollen release and general plant fertility. The middle L2 layer produces archesporal cells that divide periclinally into outer parietal cells and inner sporogenous cells. Parietal cells by further divisions form the outer part of the tapetum, middle wall layer(s) and endothecium, which enforces the power for mechanical opening of the anther by the unique mechanism of the cell wall thickening (Scott et al. 2004). The mutation in *MYB26* gene causes the male sterility by defective cell wall fortification in this endothelial layer which unables the opening (Steiner-Lange et al. 2003). The sporogenous cells give rise by further divisions to microsporocytes or meiocytes and such differentiation establishes the male germ lineage (Yang and Sundaresan 2000). The last L3 layer contributes

to the formation of a connective, vascular bundle, and the inner part of tapetum (Goldberg et al. 1993). The tapetum is a nutritive tissue forming the inner wall of the pollen sack and supporting the development of microspore mother cells due to numerous lateral interconnections by plasmodesmata (Heslop-Harrison 1971a, 1971b). However, they lack plasmodesmata on their face towards middle layer. Sugars destined to pollen in the loculus have then to cross the middle layer and the tapetal layers by the apoplasmic pathway; it is suggested that these two envelopes could be involved in the control of sugar transport from the outer anther wall layers to the locular fluid (Clément and Audran 1995).

Several genes have been identified genetically that are critical for early anther development and microsporogenesis (reviewed by Wilson and Yang 2004). One of the earliest acting gene is the *SPOROCTELESS/NOZZLE (SPL/NZZ)* playing a role at early anther differentiation with strong expression in microsporocytes and tapetum (Fig. 1). *spl/nzz* mutant plants are male-sterile producing no sporogenous and tapetal cells (Schiefhale et al. 1999, Yang et al. 1999, Ito et al. 2004). This gene encodes a transcription factor that directly regulates a gene of *EXCESS MICROSPOROCTES1/EXTRA SPOROGENOUS CELLS (EMS/EXS)*. Its mutation causes the formation of additional male sporocytes along with lack of tapetal cells but resulting in non-viable pollen, which implicates the irreplaceable role of tapetum (Canales et al. 2002, Zhao et al. 2002). *EMS/EXS* gene encodes a putative leucine-rich repeat receptor protein kinase. Another mutation in the *TAPETUM DETERMINANT1 (TPD1)* gene showed the same phenotype as that of *ems/exc* (Yang et al. 2003) and it is known to be a putative signal peptide for secretion. Taken together, *EMS/EXC* and *TPD1* may interact as a receptor and a ligand determining the microsporocytes and tapetal cell formation (Ma 2005, Wilson and Zhang 2009). Next, *DYSFUNCTIONAL TAPETUM (DYT1)*, a putative bHLH transcription factor, is likely to be involved in regulation of *AMS* and *MS1* (Zhang et al. 2006). *ABORTED MICROSPORES (AMS)* is a member of MYC class bHLH transcription factor family. Although *ams* plants have both tapetum and male sporocytes that undergo meiosis, both tapetum and microspores degenerate soon afterwards (Sorensen et al. 2003). *MALE STERILITY (MS1)*, a nuclear protein with a PHD-finger domain accompanying RING finger and putative leucine zipper motifs, is also required for early microspore development and proper exine formation (Wilson et al. 2001, Ito and Shinozaki 2002, Ariizumi et al. 2004, Vizcay-Barrena and Wilson 2006, Ito et al. 2007). *MYB33/65* are GYMYB-like TFs regulated by *SPL/NZZ* either directly or via *DYT1* (Fig. 1). Their mutations caused premeiotic abortion of

microsporocytes probably due to aberrant tapetum development (Millar and Gubler 2005). MYB103 is regulated by EMS/EXS and it is necessary for proper tapetum and microspore development (Zhang et al. 2007). In addition, MYB32 is expressed in tapetum and its mutation leads to aberrant pollen development and partial male sterility (Preston et al. 2004) (Fig. 1).

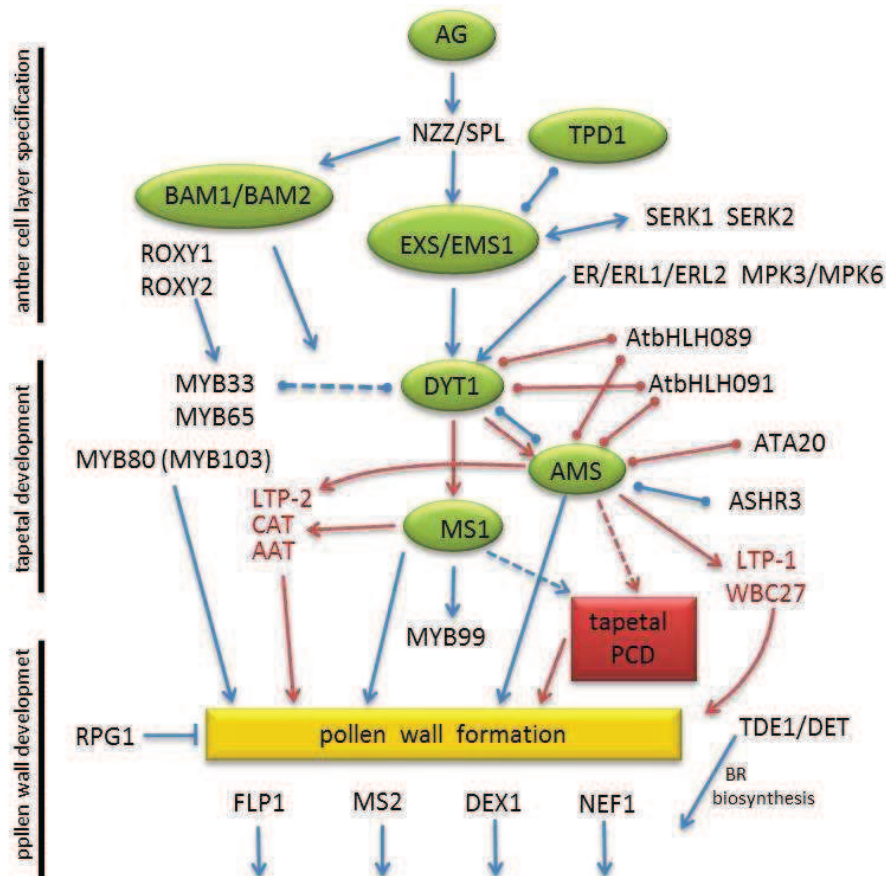


Fig 1. Regulatory network determining the anther cell wall specification, tapetum and pollen wall development indicating downstream regulation (→), and a protein-protein interaction (-) compiled from two sources: Wilson and Zhang, 2009 (blue lines) and Xu et al. 2010 (red lines).

After microsporogenesis, tapetal cells play an important role in the production of callase that helps to release individual microspores from the tetrads. In addition, tapetal cells secrete and deposit various compounds, including flavonoids and structural polymers such as sporopollenin, the most resistant biopolymer in the nature making the exine of the pollen wall. The tapetal cells eventually disintegrate and the material is deposited as a part of the pollen coat. Some of their cytoplasmic contents, including lipids and proteins, may cause allergic reaction to a human.

1.2. Pollen development

Microsporogenesis is the initial process of the male gametophyte development producing haploid microspores from the diploid microsporocytes. It is initiated by meiotic divisions I and II giving rise to tetrads of microspores. The tetrad is enclosed in a callose (β -1,3-glucan) cell wall that physiologically isolates the nascent microspores from the surrounding sporophytic tissue for the establishment, as believed, of the new developmental programme. Thick callose capsule is subsequently digested by callase secreted by tapetal cells. Such events demonstrate the tight co-operation between sporophytic and gametophytic cells for a successful process of pollen formation. At this point, *quartet3* mutant plants produce mature pollen grains remaining attached in tetrad position for the mutation in polygalacturonase which is normally secreted by tapetum.

As microspore grows, the initial three-lobed cell surface gradually evens and smoothes. The nucleus of an early microspore is located in the centre but later it is pushed towards the cell wall due to rapid vacuole biogenesis. The large central vacuole seems to be important for further haploid pollen mitosis I, an extremely asymmetric cell division producing larger vegetative and smaller generative cell. The vegetative cell makes the pollen grain itself and includes most of the former cytoplasm enriched by organelles and contains a large disperse and transcriptionally active nucleus while the generative cell encloses only very limited cytoplasm and rather condense and less active nucleus. Disruption of cell polarization dramatically changes the developmental programme and may even lead to a haploid embryogenesis as often used in plant biotechnologies (Hause et al. 1993, Eady et al. 1995). After pollen mitosis I, the generative cell migrates inward the pollen grain that then represents the unique cell-within-a-cell structure. In number of plants including *Arabidopsis*, the generative cell undergoes pollen mitosis II to form two sperm cells within a pollen grain (three-celled pollen). However, in most species (producing two-celled pollen) the second mitosis occurs during pollen tube growth when penetrating style tissues. Considering the male germline, the generative cell and later derived sperm cells involve processes that trigger different development and cell specification including chromatin rearrangement and histone modification plus germline specific histone H3.3 incorporation and transcriptional and post-transcriptional regulation by small RNAs pathways (Jones-Rhoades et al. 2006, Berger and Twell 2011, Borges et al. 2011, Twell 2011). On the cellular level, generative cell and sperm cells are characterized by condensed chromatin and very limited cytoplasm.

During the maturation of *Arabidopsis* pollen, the vegetative cell accumulates carbohydrate and lipid reserves for later demand during pollen tube growth (Pacini 1996) and similarly synthesizes mRNAs and proteins which are functionally required for rapid pollen tube growth after pollination as described earlier in other species (Tupý 1982, Mascarenhas et al. 1984, Mašek et al. 2000) (Fig. 2).

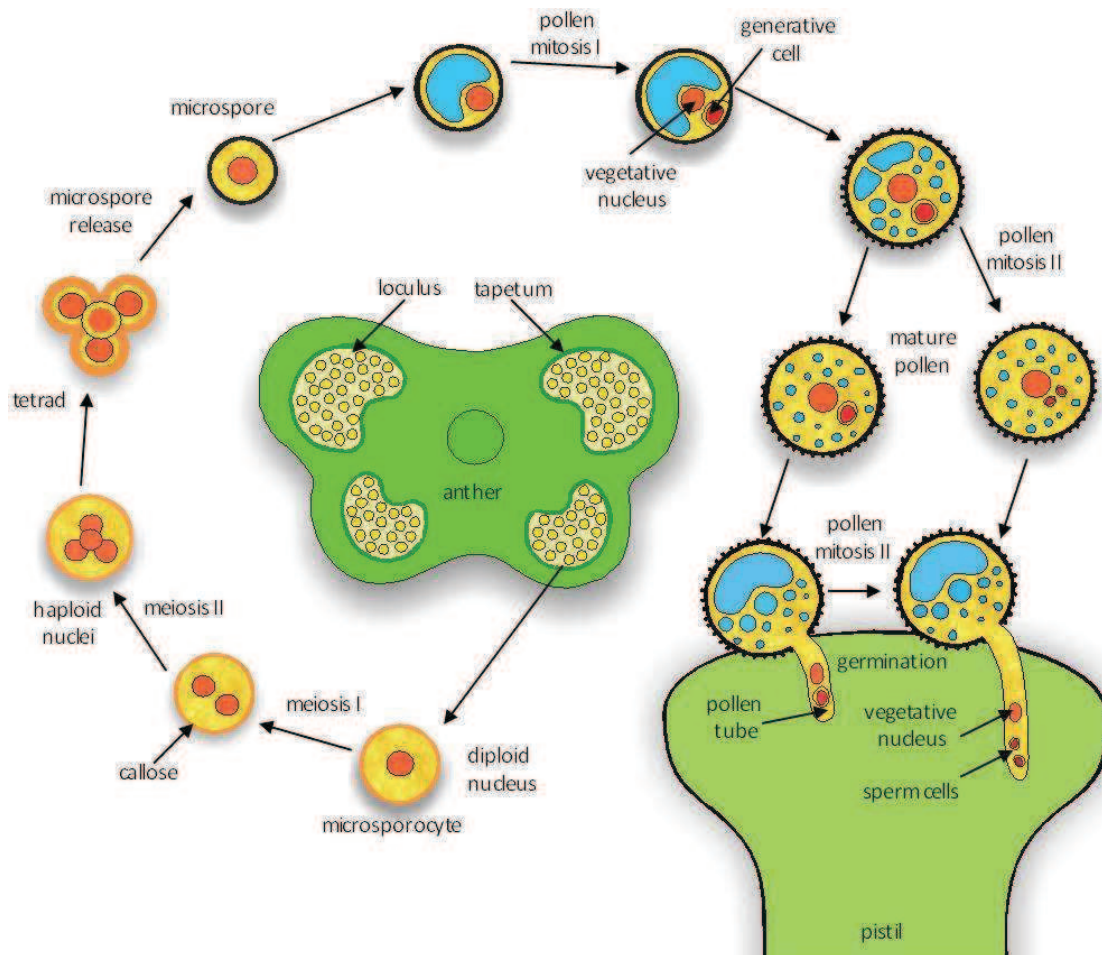


Fig. 2. Male gametophyte development. Diploid microsporocyte undergoes meiosis to produce a tetrad of haploid microspores embedded in callose wall. After enzymatic digestion by tapetal callase, individual microspores are released. After two haploid mitosis, the mature pollen grain is ready to land on stigma (*Arabidopsis*, tricellular pollen) and growth into the pollen tube delivering two sperm cells into female gametophyte. In case of bicellular pollen (on the left), the second mitosis of generative cell occurs later within the pollen tube.

Pollen cell wall displays a complex structure of different materials arranged in several distinguished layers. The internally laid intine is enriched with cellulose and pectines while outer exine, having additional sublayers of nexine and sexine, is made predominantly of sporopollenin, the complex of phenylpropanoids, phenolics, fatty acids, and carotenoids. The

exine often exhibits a highly decorated pattern characteristic for each plant species. The origin and genetic control of all pollen wall layers is different. Synthesis of exine is under the sporophytic control; however, the pattern is initiated by primexine at the tetrad stage. On the contrary, intine is completely formed and maintained from gametophytic side of the pollen wall. Exine does not form a completely enclosed layer; at specific spots (apertures) it is weakened to enable the outgrowth of a pollen tube. The final stage of pollen development is significant for pollen dehydration and production of osmoprotectants such as sugars, proline, and glycine-betain (Schwacke et al. 1999).

1.3. Pollen tube growth

Rehydration of a pollen grain after the landing on a stigma is a crucial initiation step for following vegetative cell activation. Rapid water uptake is accompanied by massive efflux of ions, sugars, amino acids, enzymes and other compounds. It boosts metabolic activity, respiration and various subcellular events including formation of vesicles, lipid droplets, vacuoles, and endomembrane biogenesis. Dynamic cytoskeletal rearrangements, mainly actin microfilaments, support the aperture, which was already determined during pollen maturation, for the outgrowth of intine when pollen germinates. Other significant features include higher dictyosome activity, massive RNA and protein synthesis (Raghavan 1997), and polysome formation to provide a translational site for stored mRNA/mRNP molecules, which are essential for pollen germination and early tube growth (Čapková et al. 1994, 1997, Honys et al. 2000 and 2009). There is the great evidence that the application of translational inhibitors (cycloheximide) arrests pollen tube growth (Čapková et al. 1980) unlike the transcriptional inhibitors (actinomycin D) that have no or very mild effect on pollen tube growth (Lafleur and Mascarenhas 1978, Honys and Twell 2003).

After germination, the vegetative cell becomes highly polarised and self-arranged into four subcellular zones for the whole period of tube growth: apical, subapical, nuclear and vacuolar zones from the tip backwards (Raghavan 1997). The apical zone is typical of the presence of vesicles derived from dictyosomes carrying carbohydrates and fusing with plasma membranes in the process of exocytosis (Pierson et al. 1990) and endocytic vesicles bringing various materials into the cell. The very tip is free of cytoskeleton and keeps sharp but oscillating gradient of Ca^{2+} ions (Malhó et al. 1994, Pierson et al. 1994) which correlates with oscillating vesicle-trafficking system and depositing methyl-esterified, and deesterified

and Ca²⁺-bound pectins into the cell wall and oscillating tip growth (Pierson et al. 1995, Geitmann et al. 1996). The subapical zone is enriched by most organelles especially dictyosomes and mitochondria, while vegetative nucleus and sperm cells stay in nuclear zone, and the last vacuolar zone is occupied by large vacuoles regularly separated by callose plugs (Cresti et al. 1977) to keep the cytoplasm towards in constant volume when pollen tubes grows. Pollen tube zonation and intracellular movements are directly generated by molecular motors (myosin, kinesin) with association to cytoskeletal elements (Pierson et al. 1985, Lancelle et al. 1987, Raghavan 1997). The composition of the cell wall depends on the location along with pollen tube; at the very tip it is composed of pectins providing very elastic property for the active tip growth, subapically the inner cellulose wall is added for mechanical support and further backwards callose is deposited from the inner site to add even more resistance during the growth through transmitting tissue making the cell wall finally three-layered.

2. Molecular biology and genetics of the male gametophyte

2.1. Transcriptional regulation

Initiation:

Gene expression is the fundamental process to support the life of the cell and it involves multiple steps of regulation occurring in various subcellular compartments. The process has been traditionally divided into following events: signalling, chromatin remodelling, transcription, mRNA processing, post-transcriptional regulation and distribution of mRNA, translation, post-translational regulation, and degradation. It is obvious that such cascade of information flow represents the complex machinery and needs to be very well orchestrated. Each step also allows the cell to regulate the temporary need according to the internal or external signals at multi-level regulatory manner and therefore each step plays its own important role during the gene expression. However, the transcriptional control is considered as the primary level of regulation of the fate of a single cell/multicellular organism. The key point of the transcriptional regulation is the initiation of transcription that is tightly associated with the chromatin remodelling. The whole process is controlled by a diversity of proteins involved in specific DNA-protein and protein-protein interactions that finally help to place the RNA polymerase on promoters of regulated genes. In eukaryotes, three different enzymes (*RNA polymerase I, II, and III*) catalyse the transcription of different RNA molecules. *RNA polymerases I*, located in the nucleolus, generate precursors of ribosomal RNA (45S pre-rRNA) later processed into 25/28S (plant/animal), 5.8S and 18S rRNAs. *RNA polymerase II* transcribes all protein-coding genes producing pre-mRNA. The regulation of transcription by *RNA polymerase II* is the considered subject of this chapter. Finally, *RNA polymerase III* catalyses the origin of tRNAs, 5S rRNA, and a variety of small RNAs including spliceosomal U6 and SRP-specific RNA.

The expression of protein-coding genes, as mentioned, is regulated by multiple protein-binding DNA sequences. These include promoter elements, promoter-proximal elements and distant enhancers. The promoter is a regulatory part overlapping tens of nucleotides upstream of translational initiation site and includes TATA box, CpG islands, and other elements that control the position of RNA polymerase II for the transcription. The sequence which is recognized as a TATA box usually lies 25-35 bp upstream of the start site and is very conserved. Even a single mutation significantly decreases the level of

transcription. Instead of TATA-box, some eukaryotic genes contain an alternative but less conserved promoter element called the initiator. Despite those well-defined elements, many protein-coding genes, typically low-rate expressing house keeping genes require GC-rich stretch, the islands of 20-50 nucleotides situated within about 100 base pairs upstream of the start region. Besides those elementary promoter elements, there are other sequences known as promoter-proximal elements lying within 100-200 nucleotides upstream of the start site important for cell-type specific gene expression in terms of tissue and developmental stage delimitations. Additionally, the transcription of many genes can be remarkably stimulated by enhancers located thousands/tens-of-thousands base pairs from the start site but also they are frequently found downstream in 5'UTRs, intron or exon sequences or even behind the stop codon of the controlled gene. Like promoter-proximal elements, many enhancers are cell-type-specific. Taking together, it is notable that (not only) plant promoters harbour surprisingly high amount of variation, mostly due to specific *cis*-elements, that significantly affect expression levels and support the stability yet plasticity of plant development under various biotic and abiotic conditions.

Proteins controlling the pre-mRNA biogenesis can be classified into four different functional groups: (1) chromatin-associated proteins, (2) the basic transcription apparatus and intrinsic associated factors, (3) large multi-subunit coactivators and other cofactors, and (4) gene-specific DNA-binding transcriptional factors.

(1) ***Chromatin-associated proteins***. Chromatin structure can operate in two distinct states: euchromatin which is open and accessible for transcriptional machinery and heterochromatin which is closed and inactive for gene expression. The ability of the flexible change between these two states is the presumption for active transcriptional regulation. Proteins responsible for such dynamic rearrangement include factors that covalently modify histones and remodelling complexes necessary for chromatin reorganization. Covalent modification – histone acetylation – promotes chromatin unfolding and is characteristic for actively transcribed regions. It is catalysed by acetyltransferase and associated with coactivators. On the other hand, histone deacetylation triggers chromatin condensation and deacetylase activity is often associated with corepressors (Li et al. 2002). Chromatin remodelling complexes (such as SWI/SNF and ISWI complexes) actively hydrolyze ATP for local reorganization of nucleosomes in two pathways; by sliding nucleosomes in *cis* or displacing nucleosome octamers in *trans* manner. SWI/SNF are placed with the help of

enhancer-binding factor operating upstream of the initiation site. Increasing evidence suggests that the putative plant SWI/SNF complexes control transcription of genes involved in many developmental processes such as floral organ identification and embryogenesis (Hurtado et al. 2006, Archacki et al. 2009).

(2) The **basic transcription apparatus and intrinsic associated factors** represent the general transcriptional factors that play an important role in correct placing the RNA polymerase II on promoter sequences. These general transcription factors comprise TFIIA, TFIIB, TFIID, TFIIE, TFIIF, and TFIIH. Most of them are multimeric proteins. The largest general TF is TFIID, which consists of TATA box-binding protein (TBP) and thirteen TBP-associated factors (TAFs). TBP is extremely conserved in all eukaryotes and binds the TATA box in a saddle-shape position. Once TBP has bound, it recruits TFIIB and subsequently a preformed complex of TFIIE, TFIIF, and TFIIH. The last has a kinase activity necessary for phosphorylation of the C-terminal domain (CTD) of RNA polymerase II, its separation from the associated factors and transcription start. Moreover, TFIIH harbours two key activities. First, it has a helicase activity important for DNA accessibility and second, it exhibits a histone acetylase activity and so it can function as a transcription coactivator by histone acetylation in the TATA box vicinity. The TAF subunits of TFIID appear to play a role in initiating of the transcription from promoters lacking the TATA box.

(3) **Large multi-subunit coactivators and other cofactors** represent heterogenous class of regulatory proteins including Mediator complex of about 20 subunits that essentially act as a molecular bridge between activation domains and RNA polymerase II by binding to those components directly and thus contributing to the integration of signals from several activators at a single promoter.

(4) **Gene-specific DNA-binding transcriptional factors** can act as activators or repressors and can interact with chromatin-remodelling complexes. In principle, these specific transcription factors bound to gene specific *cis*-elements and themselves are expressed in a tissue/cell-type-, temporal- or stimulus-dependent-specific manner and are therefore responsible for the specificity of gene expression. Transcription factors can interact directly with different components of the general machinery and with coactivators, affecting complex formation and provide, by combinational manner, another level of transcription regulation. Specific transcription factors are modular proteins with distinct and functionally separable domains, such as DNA-binding and activation domains. According

their DNA-binding domain and other properties they can be grouped into distinct families that are described later in an extra chapter.

Elongation and termination:

Once the polymerase (RNPII) has transcribed about 25 bases and pre-mRNA emerged from the protein complex, the 5' end is modified by capping enzyme complex, the C-terminus of RNPII is phosphorylated and further elongation is highly processive until it passes the polyadenylation signal that directs pre-mRNA 3' end cleavage and enables its further polyadenylation (Phatnami and Greenleaf 2006). RNA polymerase then can terminate at multiple sites located over a distance of 0.5-2 kb of this poly(A) additional tail. The protein complex responsible for such action is associated with phosphorylated carboxyl-terminal domain of the RNA polymerase II which serve as a flexible functional platform not only for polyadenylation but also other post-transcriptional modification including capping of nascent pre-mRNA, splicing of introns and cleavage behind polyadenylation signal.

2.2. Regulatory sequences

Up to date, a bulk of pollen-specific genes has been identified, making it possible to analyse their promoters including the potential regulatory sequences driving the specificity of their expression. Several pollen-specific *cis*-elements such as AGAAA, TCCACCATA and GTGA, have been already identified in many plant species (Twell et al. 1989, Twell et al. 1991, Eyal et al. 1995, Bate and Twell 1998, Hamilton et al. 1998, Rogers et al. 2001). Indeed, precise analysis of function of such motifs is a fruitful approach to reveal and understand the mechanisms of cell-specific gene regulation during pollen development in plants.

AGAAA and TCCACCATA motifs were well described within tomato *LAT52* promoter by Bate and Twell (1998). The promoter is active from pollen mitosis I but the accumulation of respective transcripts substantially increases during pollen maturation. Sequence analysis of the *LAT52* promoter revealed the presence of all major *cis*-regulatory elements required for pollen-specific transcription within the upstream region of -492 to -52. This region was shown to comprise three independent activator domains A, B, and C, each sufficient for transcriptional regulation. The PBII motif (TGTGGTT) within the domain B functions as pollen-specific enhancer element. Domain C contains an activator unit -72 to -52 (sub-domain C2) embodying two novel co-dependent regulatory elements AGAAA and

TCCACCATA. It was demonstrated that the transcriptional activity of *LAT52* promoter is controlled by a complex of pollen-specific *cis*-regulatory elements in a highly cooperative manner.

Similarly, AGAAA and GTGA motifs were described to regulate anther- and pollen stage-specific expression of two distinct genes, *OsIPA* and *OsIPK*. *OsIPA* encodes a protein similar to expansins, pollen allergens, and is activated during the late stage of pollen development, whereas, *OsIPK* promoter is active in developing anther till the pollen maturation and encodes a CDPK, a calcium-dependent protein kinases (Gupta et al. 2007).

In addition to known regulatory sequences, recently described *cis*-elements in *SBgLR* gene of *Solanum tuberosum* were demonstrated to drive the late pollen specific expression. A detailed promoter dissection analysis revealed that the region from -345 to -9 (counted upstream from ATG as +1 site) was sufficient in restricting gene expression specifically in pollen (Zhou et al. 2010). Moreover, the region between -345 and -269 was found as an enhancer while the newly described motif lies in the region of -253 to -227 and suppresses the function of nearby palindrome TTTCTATTATAATAGAAA in the region of -227 to -209. Interestingly, two pollen-specific palindrome motifs here TTTCT and AGAAA surround a putative TATA box and from this site downstream to ATG they are presented in additional nine copies plus one of GTGA. Besides the two known motifs (AGAAA/TTTCT and GTGA), the authors showed their spatial and combinational distribution within the promoter to affect the specific regulation of *SBgLR* gene expression. Its protein shares an 81% homology to SB401, encoded by a pollen-specific gene, that bounds to microtubules and actin filaments. Hence the *SBgLR* may be involved in cytoskeletal organization during pollen development (Zhou et al. 2010).

Detailed analysis of another well-described tobacco pollen-specific gene *NTP303* revealed an AAATGA regulatory element of which TGA triplet was shown to comprise an active part of the motif (Weterings et al. 1995, Verelst et al. 2007). This motif is completely conserved within the similarly regulated promoter of the *Bp10* gene from *Brassica napus* encoding the *NTP303* homologue. In addition, distal region of *NTP303* promoter shows an extensive overlap (CTTGTGTGGTTAAT) with a BPIII region of tomato *LAT52* promoter.

In *Arabidopsis*, three nuclear-encoded genes for mitochondrial Complex I (nCI) subunits (PSST, TYKY, and NADH binding protein) were found to share promoter regulatory sequences responsible for enhancement of their expression in pollen and anther wall. In

those genes, the common motif TGTGGTT involved in such specific regulation has a similarity to a “pollen box” (52/56 box) in inverted orientation. It is present in two tomato pollen-specific genes *LAT52* and *LAT56* (Twell et al. 1990, 1991) and in promoters of two pollen-specific pectin esterase genes from *Brassica campestris* (Kim et al. 1997). Obviously, regulatory sequences in those three mitochondrial *Complex I* genes play an essential role for correct spatial and temporal activity since anther/pollen development is highly energy-demanding process in plants.

2.3. Specific transcription factors

The genome-wide comparison of transcription factors among eukaryotic organisms revealed the evolutionary consequence of diversity in the regulation of transcription reflecting the biology, life style, strategy, and adaptation of various species. Each of individual eukaryotic kingdoms (*plants, animals, fungi*) has its own set of particular transcription factor families and genes.

Among 19 families that are shared between animals and plants, more than 14 are larger in plants than in animals. Such dramatic expansion in plants is not only as a result of gene/genome duplication but also as a frequent adaptive response to selection pressure in plants. Interestingly, bioinformatic analysis indicated that TF families have higher duplicability than genes involved in most functions in *Arabidopsis* (Shiu et al. 2005, Qu and Zhu 2006). In most cases, the TF family counterparts in *Arabidopsis* and rice genomes are of similar sizes. There are, however, a few exceptions in which the numbers of members within a family differ substantially between *Arabidopsis* and rice including WRKY, NAC, bZIP, MADS, ALFIN-like, GRAS and C2C2-dof families (Qu and Zhu 2006). There are about 20 families represented specifically in plants and several of them are at the same time the most abundant families in *Arabidopsis* at all, namely AP2/EREBP, NAC, and WRKY. That clearly reflects the uniqueness of the plant kingdom and the evolutionary adaptation for plant-specific functions (considering settled lifestyle, balancing environmental changes etc.). Members of AP2/EREBP families are involved in development, cell proliferation, secondary metabolism, abiotic and biotic stress response and ABA/ethylen response. Genes of NAC specifically regulate developmental processes including pattern formation and organ separation, whereas WRKY transcription factors are mostly involved in defence response.

The other highly represented and plant-specific gene families represent Dof, CO-like, YABBY (all from C2H2 superfamily), GRAS, Trihelix, TCP, ARF, C3H-type, SBP, Win-like, AB13/VP1, Alfin-like, EIL, LFY, and ARF-Aux/IAA, HB, MADS TFs. On the other hand, there are several gene families comprising transcription factors playing roles specifically in animals and therefore not represented in the Arabidopsis genome like NHR (C8-Zn₂), Adf-1, T-Box, ETS, and Fork head. In addition to other abundant plant families, the MYB superfamily (consisted of MYB (R1)R2R3 and MYB-related) which is otherwise present in all model species, is dominant just in plants where it represents the largest superfamily of all counting about 180 gene members.

The classification of transcription factor families follows the functional organization of the factors with the most significance attributed to the DNA-binding domain playing the cognitive role in targeting specific *cis*-elements. The other prominent feature is the protein-protein interaction domain enabling the interaction with the RNA polymerase II or with other co-factors, either positively or negatively. Regarding the protein secondary structure, several distinct motifs were identified to mediate DNA binding properties. Such motifs have been used to classify transcription factors into families.

(1) **Helix-turn-helix** was the first described motif in a large number of various transcription factor genes of different classes suggesting its importance in regulatory activities including homeobox proteins, MYB, HSF factors (Riechmann et al. 2000, Stracke et al. 2001, von Koskull-Döring et al 2007). It contains two α -helices separated by three amino acid spacer in rigid 120° angle. The complete factor is set together as homo- or heterodimer. This structure is embedded in a homeodomain of 60 amino acids that consists of three α -helices and a flexible N-terminal arm. Helix III, the recognition helix, packs against the DNA major groove and is responsible for specific contacts with DNA (Gehring et al. 1994, Wolberger 1996). Homeodomain-encoding genes share not only sequence similarity, but in general they play related roles *in planta* as well, predominantly during development.

(2) **Zinc-finger motif** is a proposed structure in which a loop of two β -sheets and one α -helix contains twelve amino acids including, in the typical case, two invariant cysteine and two invariant histidine residues that coordinate one zinc atom, so called class I C2H2 finger (Ciftci-Yilmaz and Mittler 2008). However, this structure can be modified into class IV C3H by replacing one histidine residue with another cysteine; this group is typical for plants (Wang et

al. 2008). Alternatively, there is a pair of fingers binding two zinc atoms in class II C8 or class III C6 found in yeast. Zinc-finger factors operate as homodimers.

(3) **Leucine zipper/bZIP TFs** contain characteristic leucine-rich region in which successive leucine residues occur at every seventh position supporting an α -helix structure. Moreover, this long side chain tends to dimerize into parallel coiled-coil structure, the leucine zipper (Oas et al.1990, Zhou et al. 1992). Hence, the leucine zipper is not responsible only for DNA binding but also for such protein-protein interaction. However, the zipper is not itself the DNA binding domain, unlike zinc finger or helix-turn-helix, but it helps DNA binding by facilitating of adjacent region of basic amino acids which interact with DNA directly. In general, bZIP TFs bind DNA as homo- or heterodimers (Ellenberger et al. 1992). Heterodimers were shown to convey synergistic activation properties to target genes, suggesting that heterodimerization serves as an efficient mechanism of signal integration (Sib ril et al. 2001, Jakoby et al. 2002, Weltmeier et al. 2006, Alonso et al. 2009).

(4) **Helix-loop-helix/bHLH TFs** contains two α -helices spaced by a loop of several amino acids. This motif is distinct from the helix-turn-helix in that it can form two amphipathic helices, containing all charged amino acids on one side of the helix. Helix-loop-helix motif plays a role similar to that of leucine zipper allowing dimerization of the transcription factors and thereby facilitating binding to DNA by the basic motif (Murre et al. 1989). Again, these factors pair as a homodimers or restrict their heterodimerization activity to closely related members of the family (Toledo-Ortiz et al. 2003). However, the others can dimerize with bZIP factors or members of R2R3 MYB proteins providing extensive combinational and various functional activities (Kuras et al. 1997, Heim et al. 2003).

2.4. Male gametophytic gene functions

After releasing microspores from tetrads they need to undergo certain structural and functional events during the development till the pollen maturation. Those includes cytoskeletal rearrangement, migration of nucleus, two haploid mitoses, synthesis of reserve metabolites, cell wall development, etc. Up to date, there are tens of genes playing role during the male gametophyte development (Table 2), whose mutations, functions and coding genes were mostly described. Probably the most striking event in pollen development is the cell cycle regulation which will be described more. It controls the essential asymmetric division of microspore nucleus during pollen mitosis I producing a

vegetative and generative nucleus. The later is further divided into two sperm cells during pollen mitosis II. Microspore enlarges and produces a large vacuole that pushes the nucleus towards the cell wall with the cooperation of microtubule rearrangement that essentially establishes the asymmetrical division. This event seems to be critical for the formation of germline since induced equal division results in two daughter cells that both exhibit vegetative cell fate (Eady et al. 1995). Several mutants have been isolated that demonstrate importance of asymmetric division including *sidecar pollen (scp)*, *gemini pollen1 (gem1)*, *gemini pollen2 (gem2)*, and *two-in-one (tio)*.

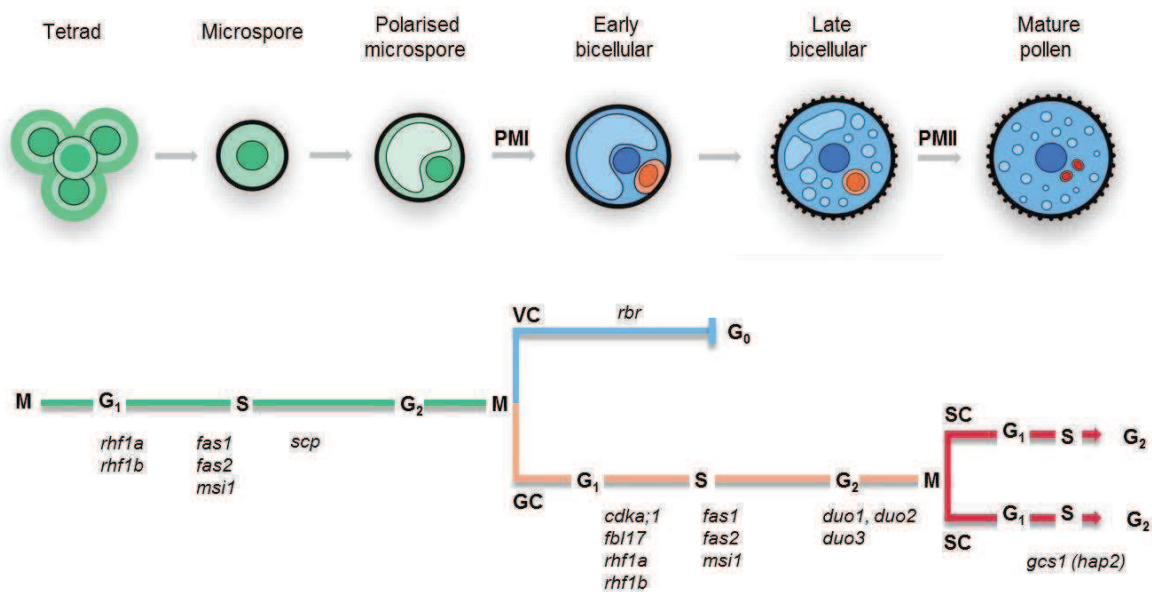


Fig. 3. Male gametophyte development and mutants affecting cell division, patterning and cellular fate in *Arabidopsis*. During microgametogenesis, the haploid microspores undergo a highly asymmetrical division, called pollen mitosis I (PMI), to produce a bicellular pollen within one vegetative nucleus (blue) and one generative cell (orange). Later, the generative cell enters further mitotic division, pollen mitosis II (PMII) to make two sperm cells (red). Compiled from two sources: Borg et al. 2009 and Twell 2011.

sidecar pollen is a male-specific mutant characterized by symmetrical division, followed by asymmetric division of only one of the daughter cells to produce mature pollen with an additional vegetative cell. *gemini pollen1* displays a range of microspore division phenotypes including equal, unequal, and partial division (Park et al 1998). GEM1 is identical to MOR1 (Whittington et al. 2001) which belongs to MAP215 family of microtubule-associated proteins and plays a vital role in microspore polarity and cytokinesis by stimulating growth of

the interface spindle and phragmoplast microtubule arrays (Twell et al. 2002). *gem2* displays the similar defects in establishing germ cell fate (Park et al. 2004). In the *tio* mutant, microspores complete nuclear division but fail to complete cytokinesis resulting in binucleate pollen grains. TIO is a plant homologue of a Ser/Thr protein kinase FUSED (Oh et al. 2005), which is localised to the phragmoplast where it plays an essential role in centrifugal cell plate expansion. Similar role to TIO could be linked with Kinesin-12A/12-B, functional redundant microtubule motor kinesins, localised also to the phragmoplast midline (Oh et al. 2010). TUBG1 and TUBG2, functionally redundant γ -tubulins, are required for spindle and phragmoplast organization in Arabidopsis microspores (Pastuglia et al. 2006). Microtubules form a basket-like nuclear cap connected to nuclear envelope and cell cortex and could be therefore involved in generating asymmetrical position of nucleus prior to division.

After pollen mitosis I, the fate of vegetative nucleus is co-determined by RBP (RETINOBLASTOMA-RELATED PROTEIN), the conserved repressor of cell proliferation. *rbp* mutation results in hyper-proliferation of the vegetative cell and to a lesser extent the germline, it perturbs cell differentiation to various degrees leading to pollen with four sperm cells (Chen et al. 2009). Hyperproliferation in the absence of RBP is dependent on CDKA;1 activity since introduction of a *cdka;1* mutant allele prevents cell proliferation in *rbp* pollen. This places RBP repression of the E2F-DP pathway downstream of KRP-dependent CDKA;1 inhibition, so that both mechanisms may cooperate to enforce cell cycle exit that is associated with commitment to vegetative cell fate (Fig 3, Fig. 4).

Considering the generative cell, the cell cycle is co-regulated by A-type cyclin-dependent kinase (CDKA;1) mentioned above, which mutation results in retarded S-phase, when germ cell enters mitosis during pollen germination (Iwakawa et al. 2006, Nowack et al. 2006) and pollen tube growth (Aw et al. 2010). Disruption of the *F-box-Like17* (*FBL17*) gene, which is only transiently expressed in the male germ cell was shown to phenocopy the *cdk;1* germ cell defect (Kim et al. 2008, Gusti et al. 2009). FBL17 normally targets the CDK inhibitors KRP6 and KRP7 for proteasome-dependent degradation (Fig. 3, Fig. 4), enabling the germ cell to progress through S-phase (Kim et al. 2008). Conversely, the persistence of KRP6/7 in the vegetative cell is proposed to maintain the inhibition of CDKA activity and vegetative cell cycle progression. Germline specific expression of FBL17 thus enables differential control of the cell cycle in the germ and vegetative cells, effectively licensing

germ cell for progression through S-phase (Twell 2011). Similarly, the proteasome-mediated degradation of KRP6 by two RING-finger E3 ligases, RHF1a and RHF2a is required for regular progression through both microspore and the subsequent germ cell mitotic cycle. RHF1a and RHF2a are proposed to reduce the level of KRP6 accumulated during meiosis, thereby preventing the inhibition of CDK activity in microspores and germ cells (Liu et al. 2008).

Recent analysis of Chromatin assembly Factor-1 (CAF-1) pathway mutants (*fas1*, *fas2*, *msi1*) indicates that chromatin integrity is also important for germ cell division. The mutants display a range of phenotypes with some failing to divide at pollen mitosis I, some failing to divide at pollen mitosis II, and some successfully dividing to produce tricellular pollen. Surprisingly, pollen grains with single germ cell are able to fertilize either the egg or central cell (Chen et al. 2008).

A single germ cell phenotype is also present in *duo pollen (duo)* mutants. In these mutants, asymmetric microspore division at pollen mitosis I is completed, however, the resulting germ cell fails to undergo cell division at pollen mitosis II (Durberry et al. 2005, Borg et al. 2009), *duo1* and *duo2* pollen then contains only single germ cell and vegetative nucleus. DUO1 encodes a novel R2R3 MYB transcription factor specifically expressed in germline cell (Rotman et al. 2005, Singh et al. 2008). Unlike *fbl17*, *cdk;1* and CAF-1 pathway-deficient mutant pollen, *duo1* pollen cannot fertilize. DUO3 is required for G2/M transition and for the expression of a subset of DUO1 target genes. However, unlike DUO1, the requirement for DUO3 in the germline during G2/M transition acts independently of CYCB1;1 (Brownfield et al. 2009). The mechanism by which DUO3 influences the expression of a subset of DUO1 targets remains unknown, but it has been proposed that DUO1 and DUO3 may cooperate in a transcriptional complex (Fig. 4).

Considering the sperm cells, their surface proteins are likely to play important roles in the guidance, recognition, and/or fusion of gametes during double fertilization. This is the case of Hap2 that has been identified in *Arabidopsis* (von Besser et al. 2006). It is a homolog of GCS1 (Generative Cell-Specific1), encoding a gamete surface protein required for pollen tube guidance and fertilization (Mori et al. 2006, von Besser et al 2006). In addition to the gene expression, germline and sperm cell specific histon H3 variant H3.3 (MGH3) is expressed under the control of DUO1 transcription factor. Histon H3.3 is incorporated into chromatin and corregulates its rearrangement to support the global reprogramming events (Twell 2011).

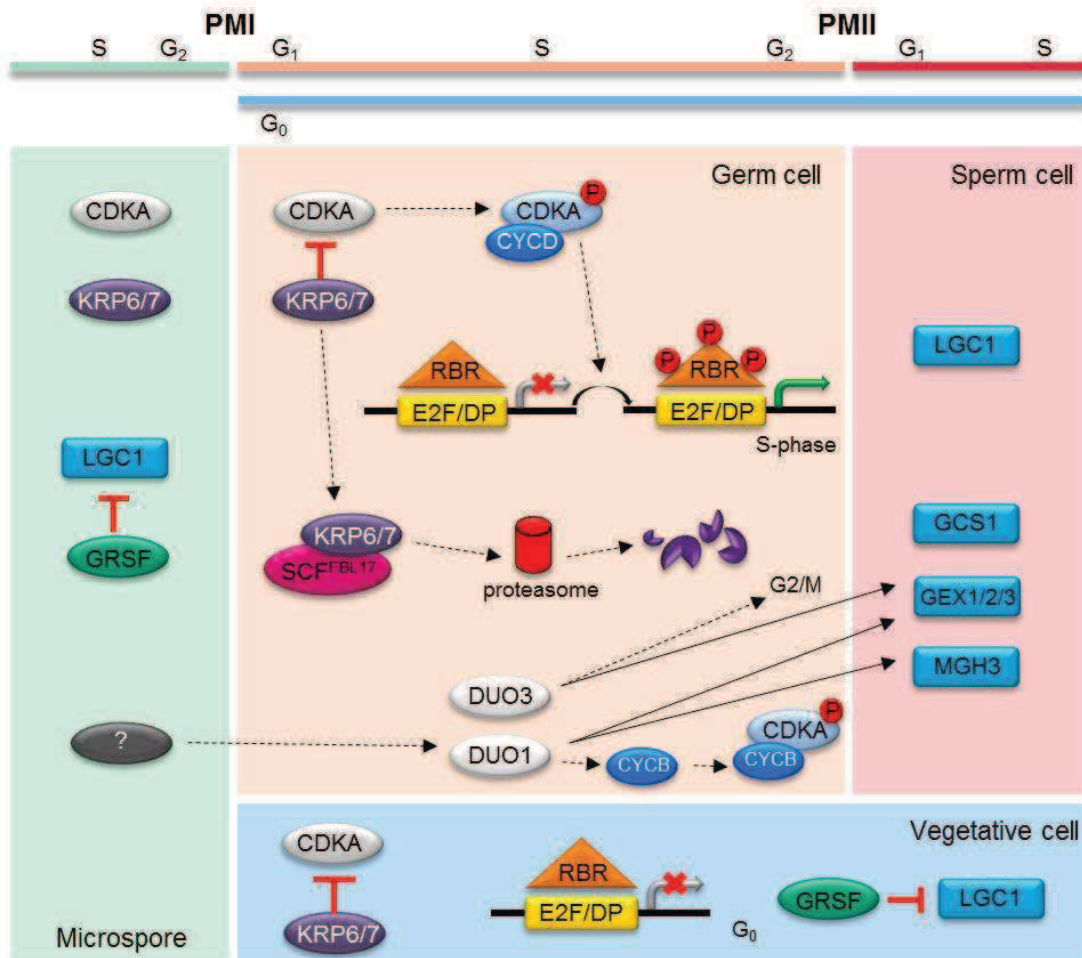


Fig. 4. A model of male germline specification and maintenance. After PMI, the cell cycle inhibitors KRP6 and KRP7 are present in both the germ and vegetative cells. Transient expression of FBL17 in the germ cell, unlike in vegetative cell, leads to the degradation of these inhibitors, enabling CDKA activation and entry into S-phase. Once S-phase has been completed, the DUO1-dependent activation of G₂/M phase regulators coupled to CDKA activation, promotes the germ cell to progress through the G₂/M checkpoint and enter mitosis. Ultimately, the co-ordinated association of these parallel pathways results in a pair of fully differentiated sperm cells. Compiled from Borg et al. 2009, Berger and Twell 2011, Twell 2011.

During the male gametophyte development, the structural and functional changes are obviously connected to regulation of transcriptome dynamics. Pollen-specific MIKC class of MADS-domain transcription factors play a significant role in such regulation. The Arabidopsis MIKC genes can be subdivided into two groups whose members make functional heterodimers; S-class involving AGL66, AGL67 and AGL104 etc., and P-class involving AGL30, AGL 65 and AGL94 etc. AGL30, AGL65, AGL66, AGL94 and AGL104 are expressed specifically in pollen while AGL18, AGL29, AGL49 and AGL84 appear to be enriched in pollen. Five

heterodimers have been postulated to form in pollen: AGL65- AGL66, AGL65- AGL104, AGL30- AGL66, AGL30- AGL104 and they may be considered as the major regulators of pollen maturation programs (Verelst et al. 2007, Adamczyk and Fernandez 2009). It was intriguing that even though more than 1,300 transcripts were misregulated in a triple mutant pollen (*agl65/66/104*), the basic cellular organization and characteristic properties of mature pollen and the ability to germinate and fertilize ovules appeared to be largely unaffected (Verelst et al. 2007). It proves the great plasticity and potential dynamics in pollen transcriptome.

3. Male gametophyte transcriptomics and microarray databases

3.1. Male gametophyte developmental transcriptomics

The male gametophyte represents a unique plant structure and plays an unsubstitutable role during plant life cycle. This fact is reflected in various cellular events including specific gene expression pattern. Despite the long-term knowledge of overlapping gene expression during pollen development based on isozyme (Tanksley et al. 1981, Sari-Gorla et al. 1986, Pedersen et al. 1987) and hybridization studies (Stinson et al. 1987, Mascarenhas 1990), it was advanced microarray technology that helped to reveal the transcriptional profile of male gametophyte in great detail (Becker et al. 2003, Honys and Twell 2003, Pina 2005, Borges et al. 2008). For pollen developmental transcriptomics, it was necessary to carry out gene expression profiling over the whole developmental process, including microspores (UNM), bicellular pollen (BCP), tricellular pollen (TCP), and mature pollen grain (MPG). Additionally, the potential to effectively isolate RNA from pollen developmental stages, pollen tubes and sperm cells in required quantity and purity enabled further extensive transcriptomic studies (Honys and Twell 2004, Borges et al. 2008, Wang et al. 2008, Qin et al. 2009).

With the use of ATH1 Genome Array harbouring 22,591 genes that represent 80.7% of estimated 28,000 protein coding genes in *Arabidopsis*, it was revealed that 13,977 genes were actively transcribed providing significant and consistently positive expression signal in at least one stage of male gametophyte development. It represented 61.9% genes on the chip and suggested about 17,000 genes playing a role during pollen development. This huge number was in accord with previous estimations. However, it evoked a surprising finding that such specialized cell lineage with unique gametophytic function shared almost two thirds of its genes involved in sporophytic regulation as well (Honys and Twell 2004). The uniqueness of the pollen transcriptome was dissected by analysing the frequencies of representation of gene ontology categories. The most significant categories included general and cell wall metabolism, cytoskeleton, signalling, transport and vesicle trafficking. It clearly demonstrated the functional specialization of pollen and its predetermination to the pollen tube growth (Honys and Twell 2003, Pina et al. 2005).

Among all 13,977 genes expressed in pollen, 1,355 genes were considered pollen-specific based on their expression at least one stage of pollen development and no expression signal in any sporophytic tissue. The second key finding was the quantification of the gradual decrease of number of expressed genes from early to late phase. In numbers, UNM – BCP – TCP – MPG – PT4 – SPC stages expressed 11,565 – 11,909 – 8,788 – 7,235 – 6,148 – 5,801 genes, respectively (Fig. 5). The bulk of genes transcribed during early pollen development might reflect the intense metabolism and cell growth of microspores and young pollen. On the contrary, during pollen maturation, the steady decrease could be connected to the cessation of metabolic activity towards mature pollen grain until pollen germination and tube growth. However, the proportion of pollen-specific genes expressed over male gametophyte development steadily increased as calculated 6.9% - 7.2% - 8.0% - 8.6% - 10.9% - 11.5% for each respective stage (Fig. 5). The rising proportion of pollen-specific genes expressed in later pollen phase may be associated with desiccation and other specific pollen functions during maturation (Honys and Twell 2004).

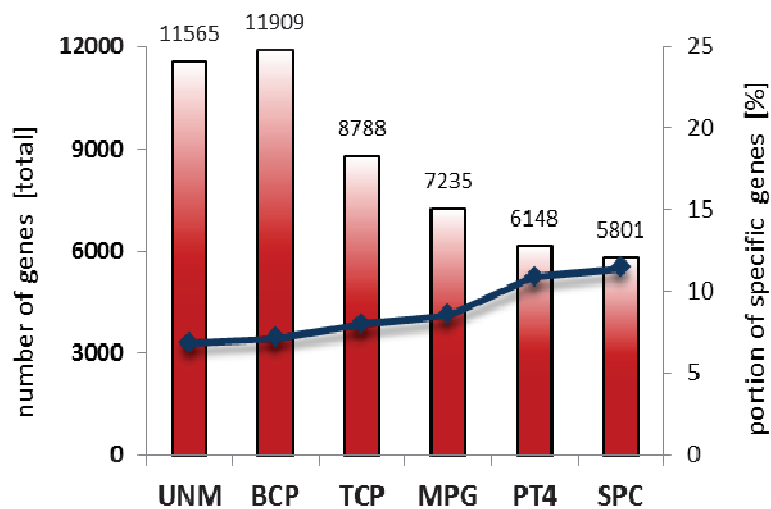


Fig. 5. Transcriptome profile of total expressed genes during male gametophyte development with the portion of specific genes active in each developmental stage. Columns from UNM to PT4 (4hr pollen tubes) represent the transcriptome of vegetative nucleus while SPC represents the transcriptome of sperm cells. Data from: Honys and Twell 2004, Borges et al. 2008, Wang et al 2008, Qin et al 2009.

Another striking phenomenon of pollen transcriptomic was a dramatic change of two subsequent global developmental programmes between early and late stages. There is a significant group of genes expressed specifically/predominantly in UNM and BCP with sharp decrease in expression at later stages and vice versa. Basically, such reprogramming reflects

the transition between BCP and TCP stages at the cellular level. When comparing UNM and BCP by scatter plot analysis, relatively high correlation coefficient of $R=0.96$ indicated close relation between these stages (Fig. 6). Similarly, the coefficient of $R=0.86$ between TCP and MPG also suggested great similarity in terms of expression profile of respective genes. On the contrary, BCP and TCP showed very poor correlation and even less similarity between UNM and MPG (Fig 6). It simply reflected the change in “cell status” from relatively undifferentiated and proliferating population of microspores into highly differentiated mature pollen (Honys and Twell 2004).

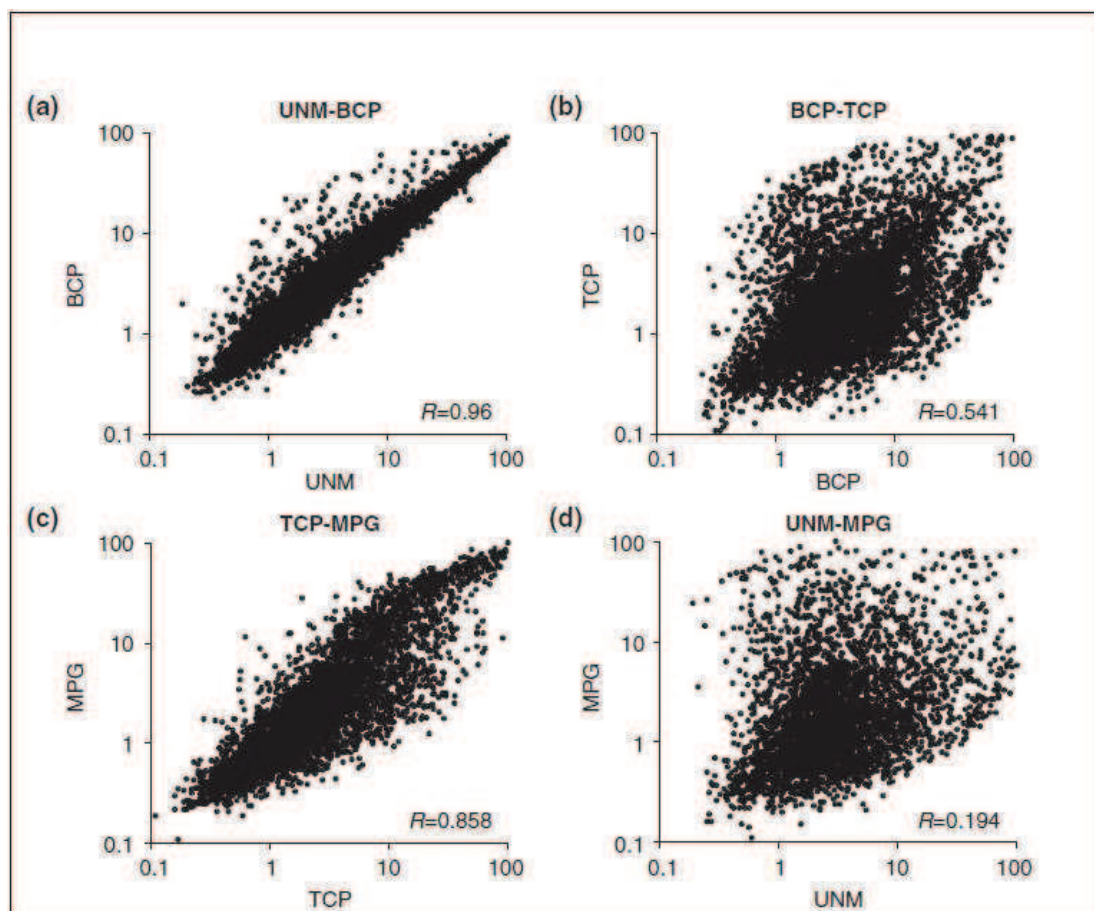


Fig. 6. Scatter-plots comparing gene expression between two respective developmental stages. (a) UNM versus BCP, (b) BCP versus TCP, (c) TCP versus MPG, (d) UNM versus MPG. Honys and Twell, 2004.

Considering transcription factors (TF), ATH1 harbours 1,350 genes that represent about 85% of total 1,594 annotated TF genes (Honys and Twell 2003). Of them, 608 (45%) TF genes were expressed at least one stage of male gametophyte development and 54 genes (15.7%) were pollen-specific. The family size also mattered rather than proportionally equal representation of genes in each TF family. There was a sharp difference between over-

represented families including p-coumarate 3-hydroxylase (C3H) family (67%), CAAT family (64%), C2H2 zinc finger proteins (57%), WRKY family (53%), bZIP family (51%), TCP family (50%), GRAS family (50%) and, in contrast, families under-represented such as AUX/IAA (20%), HSF (33%), bHLH (34%), NAC (34%), AP2-EREBP (35%), HB (36%), R2R3-MYB (37%), MADS (37%), and C2C2 zinc finger gene family (37%).

Besides pollen (vegetative cell) transcriptome, the data from isolated Arabidopsis sperm cells are also available (Borges et al. 2008). With the use of ATH1 Genome arrays (22,392), the amount of 5,829 (27%) gene transcripts was detected to be expressed in sperm cells. The functional classification of genes revealed that most represented genes were associated with DNA repair, cell cycle progression and ubiquitin-mediated proteolysis. Moreover, significant presence of components involved in small RNA metabolism and DNA methylation suggested their convincing role in epigenetic regulation. It may not be surprising that the vast majority of genes (3,813) expressed in sperm cells were also detected in pollen vegetative cell. However, the overlap between sperm cells and seedlings comprised 4,757 genes, representing almost the whole sperm-cell transcriptome with the half showing predominant expression in sperm cells. In relation to transcriptional regulation, the TF families were rather weakly represented with the exception for MYB-type TF family with relatively high expression of their members in sperm cells. Families of homeobox and bHLH genes, which were otherwise very abundantly expressed in sporophyte, were represented only by one or few genes in sperm cells, respectively. Focusing on transcription factors managing specific sperm cell functions, one candidate (At4g35700) could be traced into zinc finger C2H2 family, which was specifically expressed just in sperm cells and at the highest expression level of all TF genes on top of that, and possibly one member (at2g42300) of bHLH family being expressed specifically in male gametophyte and was not detected in pollen vegetative cell.

3.2. Microarray databases

The gene expression in its specific combinatorial pattern and dynamics during whole development represents the essential background of the life with direct connection to structural and physiological behaviour of the organism. Therefore the study of transcriptome as a spatial and temporal result of inner and outer stimuli is of the great interest. In past decades, number of techniques have been used to discover the secret of transcriptional

activity using Northern blot analysis, quantitative reverse transcription-PCR, cDNA library screening, serial analysis of gene expression, manual/mechanical spotting of probes on nylon/glass macro/microarrays, etc. (Schena et al. 1995, Velculescu et al. 1995, Lockhart et al. 1996, Lipshutz et al. 1999). Those approaches were able to detect obviously very limited number of genes. It was the introduction of sophisticated high-throughput microarray technologies that significantly supported the parallel large-scale transcriptome analyses. Nowadays, microarray ATH1 GeneChip developed by Affymetrix covering 76% of Arabidopsis thaliana genes is considered as a standard tool in genome wide transcriptomic studies in plants.

Not surprisingly, the technological reliability, reproducibility, price availability and last but not least the experimental requirement have contributed to the massive production of transcriptomic data. Along with continuous data accumulation and the customers' need for the access to the gene expression information, numerous databases have been established. Currently, there are dozens of different databases covering different species, experimental conditions, and developmental stages which may result in navigation difficulties. Over last years, many articles have been published to deal with microarray databases, providing relevant introduction and help researchers direct to resources and tools they require. Basically, there are two types of microarray databases; one serves as a publicly available data repository and the others provide some additional data analysis tools. In both cases, experimental data comes directly from Affymetrix Service via Nottingham Arabidopsis Stock Centre (NASC) or from extensive bioinformatic projects run by large institutions like European Bioinformatics Institute (EMBL-EBI) or National Centre for Biotechnology Information (NCBI) as the largest data repositories. Alternatively data originates from numerous laboratories' projects dealing with transcriptomic analyses. Bioinformatic tools associated with most databases represent the main instrument to effectively select and analyse data of interest. Essentially, such facilities make the difference among each database and may attract different scientific questions. To mention several key data mining tools, it would include normalization and data filtering tools, global statistical data analysis, analysis of co-regulated genes, and hierarchical or K-means clustering tools. Some specialized databases offer visualizing tools, promoter element detection or potential upstream elements. Other helpful possibilities represent the selection from various experimental setups including physiological and stress conditions, different developmental stages and

tissue/cell lines. Moreover, most databases permit data download, either for free or charged, for further data processing if needed. Because of the diverse data sources and experimental conditions, the large number of databases is strictly MIAME-compliant, asking for the Minimum Information About Microarray Experiment (MIAME) to enable comparability across numerous datasets. Further criteria also comprise the assessment of expression data quality (QC, Quality Control) to evaluate the relevance of providing data. Ongoing microarray experiments and continuous generation of transcriptomic data have led to establish various microarray databases. Many applications are being supplemented with more specialized products offering various visualization and data mining tools to fulfil growing demands of researchers. It was one aim of this Ph.D. thesis to overview the most frequent databases (Honys et al. 2008) and setup a new portal using own expressional data including pollen transcriptome as described in Results of the thesis.

4. Aims of the study

The study of male gametophyte represents a very interesting field in the plant science for the uniqueness of the object of interest including structural features (cell-in-cell organization, cell wall composition) and functional properties (unique transcriptional pattern, posttranscriptional regulation, tip growth) not speaking about amazing cellular and life-span reduction during the gametophyte evolution in plants. Due to massive use of microarray technologies, 608 TF genes were identified to be expressed during pollen development and some of them have been functionally characterized. However, the complete understanding of the whole regulatory network is still rather limited.

Hence, the main goal of this Ph.D. thesis was to employ large-scale screening of T-DNA mutant lines in specific transcription factors to bona fide determine their importance in pollen development. Therefore this work intended to provide knowledge on TF mutant phenotype in terms of their structural and functional defects as a basement for further detailed studies of their regulatory roles. In addition, this thesis also reviewed the male gametophyte development on the whole and reflected gene expression data and transcriptomic databases. The particular aims were:

(1) To summarise the male gametophyte development with the respect to the structure and function of pollen grain and growing pollen tube, mutants affecting male gametophyte development and pollen transcriptomics.

(2) To select TF genes with either specific or enriched expression pattern during pollen development based on microarray data with the special focus on early stages of male gametophyte development and to perform large-scale screening of T-DNA mutant lines in selected TF genes and their evaluation regarding their effect on male gametophyte development.

(3) To participate on the characterization of a candidate gene *AtbZIP34* selected from previous large-scale screening as a case study including structural and functional analysis of the mutant.

(4) To identify promoter sequences driving gene expression in early stages of pollen development and their use as a molecular tool for the manipulation of gene expression.

(5) To build a gene expression database based on microarray data and to make it public to a broad scientific community in user-friendly and graphic environment.

Moreover, to compile and compare available transcriptomic databases world-wide to help potential users to target their questions on particular gene expression.

5. The list of publications

In accord to the aims of this Ph.D. thesis, I participated in preparation of several articles and book chapters.

(1) Honys D, **Reňák D**, Twell D. Male gametophyte development and function. Global Science Books, pp.76-87, 2007.

(2) **Reňák D**, Dupl'áková N, Honys D (2011) Wide-scale screening of T-DNA lines for transcription factor genes affecting male gametophyte development in *Arabidopsis*. Plant Sex Rep, *resubmitted*.

(3) Gíbalová A, **Reňák D**, Matczuk K, Dupl'áková N, Cháb D, Twell D, Honys D (2009) *AtbZIP34* is required for Arabidopsis pollen wall patterning and the control of several metabolic pathways in developing pollen. Plant Mol Biol (2009) 70: 581-601.

(4) Honys D, Oh SA, **Reňák D**, Donders M, Šolcová B, Johnson JA, Boudová R, Twell D (2006) Identification of microspore-active promoters that allow targeted manipulation of gene expression at early stages of microgametogenesis in *Arabidopsis*. BMC Plant Biol 6: 31-39.

(5) Dupl'áková N, **Reňák D**, Patrik Hovanec, Honysová B, Twell D, Honys D (2007) Arabidopsis Gene Family Profiler (aGFP) – user-oriented transcriptomic database with easy-to-use graphic interface. BMC Plant Biol 7: 39-46.

(6) Honys D, Dupl'áková N, **Reňák D**. Online tools for presentation and analysis of plant microarray data. Nova Science Publishers, Inc. In: Oligonucleotide array sequence analysis, eds: M.K. Moretti and L.J. Rizzo, pp. 265-295, 2008.

6. Results

This chapter considers four author's research papers on wide-scale screening of T-DNA lines, *AtbZIP34* TF functional analysis, identification of microspore-active promoters, and building the aGFP database, only (excluding two book chapters). Since all results are fully presented in each of respective papers, this part is only to summarise them in a united manner.

Aiming to identify TFs affecting male gametophyte development, we screened 74 T-DNA insertion lines for pollen phenotypic defects. We focused on structural and functional defects occurring during pollen development using bright field and epifluorescence microscopy. In comparison to the wild-type phenotype, five complex disorder classes were observed. Those included (A) pollen abortion, (B) presence of cytoplasmic inclusions, (C) pollen grain size and cell wall structure, (D) cell cycle defects, and (E) MGU (male germ unit) organization. However, the precise phenotype analyses have led to refined classification of above dysfunctions into tens categories: A: (1) abortion; B: (2) inclusions; C: (3) small grain, (4) oval grain, (5) deformed cell wall; D: (6) one-celled pollen, (7) two-celled pollen; E: (8) eccentric MGU, (9) separated MGU, and (10) linear MGU. We mostly observed abnormalities connected to cytoskeletal organization and function leading to the defective arrangement of MGUs with their eccentric, separated or linear position in the pollen grain unlike their characteristic central arrangement in wild-type pollen. Because of the sole positional variation we described those phenotypes simply as misarranged MGU. Other classes of mutations were likely to affect cell cycle and cell division. Namely, PMI defects resulted in one-celled pollen and/or PMII defects resulted in two-celled pollen with one vegetative nucleus and one generative cell. Cytoplasmic inclusions represented more specific class of phenotype defects. The most serious and complex phenotypic defect(s) led to a complete cell abortion with squashed pollen wall. Considering the pollen shape, we observed unusual pollen grain size and shape such as small and oval pollen grains. However, these phenotypic defects were rare. Only one insertion line produced pollen with cell wall deformation with rough cell surface. Among 61 confirmed lines, about a half (29 lines) showed strong phenotypic changes (i.e. $\geq 25\%$ aberrant pollen). Yet only few lines producing heavily disturbed pollen grains were observed reaching very high penetrance. The most striking phenotypes were observed in lines of SALK_140819 with 100% abnormal pollen (At2g20400;

G2-like), 80-85% in SALK_018864 (At2g42380; bZIP), 70-75% in SALK_084050 (At3g15740; C3H), 70-75% in SALK_010704 (At5g54470; C2C2-CO-like), and 50-55% in SALK_090677 (At4g05330; C2C2) (Table 1).

AGI	T-DNA line	TF family	mutant phenotype
At2g20400	SALK_140819	G2-like	100%
At2g42380	SALK_018864	bZIP	80-85%
At3g15740	SALK_084050	C3H	70-75%
At5g54470	SALK_010704	C2C2-CO-like	70-75%
At4g05330	SALK_090677	C2C2	50-55%

Table 1. The list of the most affected lines by T-DNA insert in pollen development

But generally, in the population of homozygous plants we mostly identified remarkable proportion of pollen with wild-type or mild phenotype defects counting the average penetrance in between 10-30%. Several T-DNA lines with strong phenotype defects were further genetically tested using backcross analysis. Out of eight selected lines, six ones segregated close to the Mendelian 1:1 ratio. Remaining two lines performed heavy segregation distortion of 1:0.04 (SALK_002235) and 1:0.39 (SALK_043690) indicating the significant role of the TFs in male gametophyte development.

Our wide-screening led to an evaluation of selected TF mutant lines in terms of impact on pollen development and hence served as a good information basement for further studies. Based on this fact, we have selected a few lines with considerable impact on male gametophyte development which have been analysed in our laboratory. One of them was the transcription factor *AtbZIP34*, published by our laboratory (Gibalová et al. 2009). *AtbZIP34* belongs to a large family (75 annotated genes) of bZIP transcription factors whose members have been shown to be involved in male gametophyte development (Iven et al. 2010). *AtbZIP34* was confirmed to be specifically and increasingly expressed during pollen maturation both by RT-PCR and promoter:GUS analyses, that were in agreement with previously published microarray data (Honys and Twell 2004). Pollen grains of mutant *atbzip34* plants showed several phenotypic features including 15% aborted pollen and 27% malformed/misarranged MGU. In addition, the typical sign was the presence of lipid/oil

bodies. Electron microscopy confirmed the abundance of such inclusions in mutant pollen accompanied by under-developed ER and vacuoles increased in number and size. Farther observation revealed abnormal exine patterning with collapsed baculae and tectae and with areas of extra material deposition plus characteristic wrinkled intine detached from the outer nexine layer. Furthermore, its dysfunction was significant in progamic phase as well, as demonstrated by *in vitro/in vivo* pollen tube growth and segregation ratio distortion. Pollen bearing mutant allele showed weaker germination (85% reduction compared to wild-type) and slower pollen tube growth rate. According to microarray experiment, AtbZIP34 affected expression of genes involved in various metabolic pathways besides other functions. Over 1,300 genes were upregulated and, in contrast, over 800 genes were downregulated; the later subset included most significant changes in functional categories representing transcriptome, transport, metabolism and protein fate. Among the strongest downregulated genes belonged those encoding proteins involved in lipid catabolism and two lipid transfer proteins, and ABC transporter. All those findings indicated to the missing role of AtbZIP34 TF during male gametophyte development and therefore its unsubstitutable role in the regulatory network. Moreover, it is the first up to date described factor being involved in both sporophytic and gametophytic regulation.

Manipulation of gene expression during pollen development using differently active promoters may serve as a convenient tool for functional analysis. For such need, Lat52 as well characterised promoter active in late pollen development and in progamic phase has been a favourite tool for decades. In addition to Lat52 isolated from tomato, LGC1 promoter was characterized in lily generative cell (Singh et al. 2003, Singh and Bhalla 2008). However, promoters active in early pollen development were mostly missing. Therefore we focused on the identification of promoters driving expression in early stages and then isolated three microspore-active promoters enabling the spatially and temporally controlled expression pattern. Based on previous transcriptomic data from four pollen developmental stages (UNM, BCP, TCP, MPG) and several sporophytic tissues, a group of seven genes was selected exhibiting strict expression patterns only at early stages of male gametophyte development. Those candidate genes were verified by RT-PCR analysis but only three of them (At5g59040, At5g46795, and At4g26440) passed strict selective criteria on specific early expression and were named MSP1, MSP2, and MSP3, respectively. The activity of those promoters was tested using GUS:GFP markers. MSP1 showed earlier signal in microspores followed by a

sharp decrease in mature flowers while MSP2 and MSP3 initiated expression in microspores but accumulated in mature pollen as well. To evaluate MSP promoters as a suitable tool for early pollen expression *in planta*, two constructs for functional complementation using MSP1 and MSP2 promoters and *TIO* coding sequence were prepared. *tio* heterozygous plants produced 50% aberrant pollen and MSP1 construct were able to significantly (more than MSP2 construct) restore the phenotype confirming the usability of MSP1 promoter for targeted manipulation of gene expression at early pollen development.

With the increasing demand for transcriptomic studies, loads of expression data have been generated by number of research groups round the world including our and collaborative laboratory as published previously (Hony and Twell, 2003, 2004). These datasets included unique expression profiles of four pollen successive stages (UNM, BCP, TCP, MPG) of male gametophyte development plus numerous sporophytic tissues. To access those data to wide scientific community and provide it with an alternative web-based toolbox, we established a database named Arabidopsis Gene Family Profiler (aGFP). The database works with transcriptomic datasets from NASC Array and AtGenExpress and presents them in user friendly graphic environment introducing a unique virtual plant concept with coloured organs and tissues in a scale reflecting the level of gene expression. Furthermore, the data can be searched according to AGI codes, BAC loci, gene names, gene families or as custom genesets with the further possibility of keyword search. The user can also choose from MAS 4.0 or MAS 5.0 normalised datasets and from NASC Array/AtGenExpressed source to name some options. aGFP database covers only data from Affymetrix ATH1 arrays using wild-type *Arabidopsis* of different ecotypes grown under normal physiological (AtGenExpressed) or various (NASCArrays) conditions.

7. Discussion

The wide-screen analysis of T-DNA insertional lines brought in a sense surprising result. The first was, despite of numerous 21 TF families represented by 61 confirmed insertion lines, relatively small variance among individual pollen mutant phenotypes. Such limited portfolio of pollen defects may have seemed unexpected at this wide-scale screening but one should regard the role of transcription factors in complex networks when different regulators can trigger various biochemical processes that finally share and often overlap similar cellular events and functions like cytoskeleton organization, cell cycle, transport, cell wall metabolism etc. Such pleiotropic activity, but overlapping when comparing different mutants, has repeatedly been reported (Twell et al. 2006, Verelst et al. 2007, Gibalová et al. 2009). Second, we could not detect any significant bias towards any particular phenotypic category when comparing early and late TFs. Moreover, there were no preferences among individual TF families in disrupting cellular events. The common cellular disruption over all TF families affected MGU position resulted into three phenotypic defects: eccentric, separated or linear MGU. Third, among 61 verified lines we did not find many lines with considerably high penetrance with the exception of four lines reaching over 70% of aberrant pollen. On the contrary, the vast majority of lines showed wild-type-like appearance or mild phenotype defects in a range of 10-30%. Many authors suggested gametophytic mutations as pleiotropic, incompletely penetrant and displaying variable expressivity due to functional redundancy (Feldmann et al. 1997, Bonhomme et al. 1998, Grini et al. 1999, Drews and Yadegari 2002, Verelst et al. 2007) or due to a conditional-based phenotype. This phenomenon was especially apparent among transcription factors (Blanc et al. 2000, Verelst et al. 2007). Indeed, the vast majority of several hundred knockouts in *Arabidopsis* did not give rise to visible, directly informative phenotypes (Bouché and Bouchez 2001). However, out of six selected T-DNA lines for further genetic analysis using backcross, two lines performed heavy segregation distortion counting 1:0.04 (SALK_002235) and 1:0.39 (SALK_043690) in comparison to the Mendelian 1:1 segregation. It is likely that the corresponding genes (At3g20910, CCAAT-HAP and At5g54680, bHLH, respectively) play significant regulatory role in the male gametophyte development.

Further study on one selected transcription factor, AtbZIP34, revealed characteristic phenotypic and genetic transmission defects providing several lines of evidence that support

both sporophytic and gametophytic roles of this factor in male gametophyte development and function (Gibalová et al. 2009). Various phenotypic features including cell wall organization, changes in endomembrane system, vacuole and vesicle content in addition to handicapped pollen tube growth, were enlightened by microarray analyses. As a result, we identified over 1,300 genes up regulated and over 800 genes down regulated which disclosed the key position of the AtbZIP34 factor in male gametophyte regulatory network. For example, the abnormal presence of lipid/oil bodies in mutant pollen nicely correlated with significant down regulation of genes encoding proteins involved in lipid catabolism, lipid transfer and ABC transporter. Similar links could be associated to the cell wall biogenesis as the mutant pollen sustained in the structure of both intone (gametophytic management) and exile wall patterning (sporophytic management). The further analysis of downstream regulated genes including the regulator MYB97 may be promising to reveal new connections in the cellular network that control male gametophyte development.

For the functional analysis of transcription factors, manipulated gene expression using a set of promoters with various spatial and temporary activities is a very useful approach. Yet only two *Arabidopsis* promoters (*BCP1* and *AtSTP2*) were known to be active in microspores (Xu et al. 1995, Truernit et al. 1999). However, *BCP1* also showed the expression in tapetum and *AtSTP2* was active from the tetrad stage. Similarly for other species (rapeseed, potato), microspore-active promoters have been identified but functionally overlapping to other stages (Fourgoux-Nicol et al. 1999, Maddison et al. 1999) with the exception for tobacco *MTM19*, whose promoter is highly microspore-specific (Clusters et al. 1997). Because of the absence of such tool for an early expression in *Arabidopsis* pollen development, three early active promoters were selected, isolated and verified by RT-PCR and GUS expression. Moreover, two of them, MSP1 and MSP2, were functionally confirmed by complementation analysis using MSP promoter and TIO coding sequence. Interestingly, the expression in early stages of male gametophyte development was steadily accompanied with the expression in tapetum, which tightly surrounds developing microspores in anther microsporangia. In this regard, the co-regulation of gene expression in both microspores and tapetum at early stages of anther development is not surprising since they have the same origin in archesporic tissue, tapetal initials develop side by side with pollen mother cells the microsporogenesis itself is vitally dependent on secretory activity of tapetal cells.

Since the massive use of microarray experiments, the wealth of transcriptomic datasets have been widely expanding and it has been naturally followed by the introduction of transcriptomic databases collecting such data and presenting them in various manners either as a repository or with a web-based tools (Honys et al. 2008). aGFP database launched by our lab provides data from AtGenExpress and NASCArrays in easy intuitive graphic environment, which makes it unique among the others. The selection from those data source provides researchers transcriptomic data from Columbia-0 of numerous developmental stages grown under comparable conditions (AtGenExpress) and data coming from different ecotypes grown under various conditions (NASCArrays). Furthermore, none of the databases had till then offered instruments such as profiling according to gene families. In addition, interactive “virtual plant” represents another innovation helping users simply check transcriptomic data. The option between MAS 4.0 or MAS 5.0 normalized data are also rare among other databases round the world, which enables direct comparison of the influence of the detection algorithm. Empirical MAS 4.0, unlike the statistical MAS 5.0, is thought to yield more false-positive calls, our analysis of four pollen developmental stages showed that MAS 5.0 detection algorithm tended to eliminate a number of genes identified by MAS 4.0 as expressed genes and which were experimentally verified to be so, highlighting the extra value of MAS 4.0 detection algorithm. The uses can also switch between virtual plant or simple bar chart (standard or log-scaled) or tabulated display. The database is freely available on the internet and hopefully providing quick and easy steps to mine transcriptomic data as needed.

8. References

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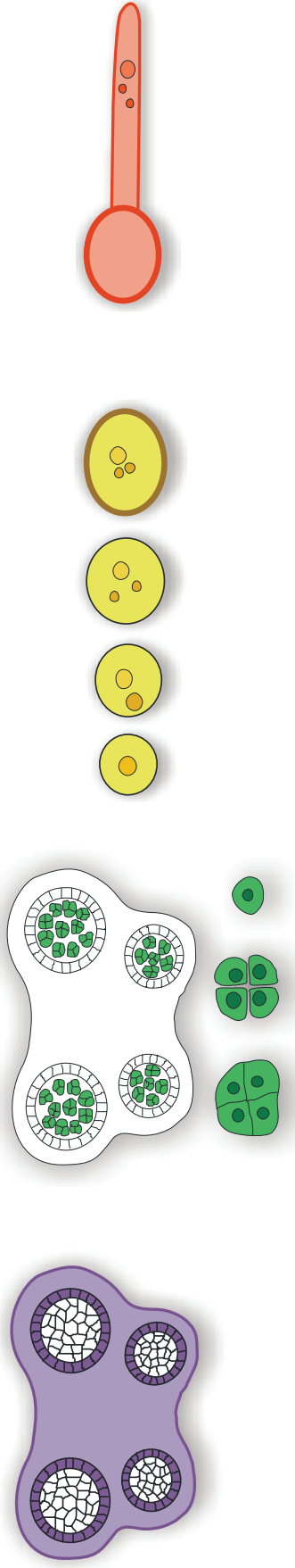


Fig. 7. Development of male sporophytic tissue (purple) and gametophyte structures (green, yellow, pink), colours correspond with Table 2.

Table 2. The list of genes affecting development of stamen, microsporogenesis, pollen development and pollen tube growth. TF genes are in red.

AGI	Gene	Mutant	Mutant phenotype	Protein Identity	Protein Function	Reference
A2g06050	<i>DDE1/OPR3</i>	<i>DELAYED DEHISCENCE 1</i>	anther do not dehiscence	12-oxophytodienoate reductase that is required for jasmonate biosynthesis	jasmonic acid biosynthetic process	Sanders et al. 2000
A2g39940	<i>CO1</i>	<i>coronatine insensitive1</i>	short filament, anther no dehiscence, conspicuous pollen	protein containing Leu-rich repeats and a degenerate F-box motif	SCF-dependent proteasomal ubiquitin-dependent protein catabolic process	Feys et al. 1994
A2g44810	<i>DAD1</i>	<i>defective anther dehiscence1</i>	defects in anther dehiscence, pollen maturation	chloroplastic phospholipase A1	catalyzes the initial step of jasmonic acid biosynthesis	Ishiguro et al. 2001
A3g13890	<i>AtMYB26</i>	<i>AtMYB26, male sterile 35</i>	male sterile because of anther no dehiscence, fertile pollen	MYB transcription factor	transcriptional activation	Steiner-Lange et al. 2003
A3g23130	<i>SUP</i>	<i>superman</i>	extra stamen at the expense of carpel	similar to zinc finger transcription factors	negative regulator, suppression of AP3 and Pl in fourth flower whorl	Gaiser et al. 1995
A3g27810	<i>AtMYB21</i>	<i>AtMYB21</i>	male-sterile, short filaments, delayed dehiscence	MYB TF	regulation of jasmonate pathway, anther maturation	Mandaokar et al. 2006
A4g24972	<i>TPD1</i>	<i>tapetum determinant1</i>	excess of sporocytes, lack of tapetal cells	small putative signal peptide	hypothesized ligand for EMS1/EXC, meiotic regulation	Yang et al. 2003
A4g25640	<i>FFT</i>	<i>flower flavonoid transporter</i>	anther dehiscence perturbed, male sterility	multidrug and toxin efflux family transporter	involved in flavonoid transport network	Thompson et al. 2010
A4g27330	<i>SPZ/NZZ</i>	<i>sporocytless/nozzle</i>	defect in anther wall senescence and sporocytes development	nuclear protein, similar to MADS box TFs	regulation of sporocytes development	Yang et al. 1999
A5g07280	<i>EMS1/EXC</i>	<i>excess male sporocytes/extra sporogenous cells</i>	additional sporocytes, lack of tapetal cells	leucine-rich repeat receptor protein-kinase	required for meiotic cytokinesis, normal tapetum development	Canales et al. 2002, Zhao et al. 2002
A5g40350	<i>AtMYB24</i>	<i>AtMYB24</i>	double mutant with MYB21: complete male-sterile	MYB TF	regulation of jasmonate pathway	Mandaokar et al. 2006
A5g56110	<i>AtMYB103</i>	<i>AtMYB103</i>	tapetum degeneration, male sterility	putative MYB TF	required for normal tapetum and pollen development	Higginson et al. 2003, Zhang et al. 2007
unknown	<i>FAT TAPETUM</i>	<i>fat tapetum</i>	enlarged tapetum and middle layer at the time of meiosis	unknown	may be critical for proper differentiation of middle layer	Sanders et al. 1999, Sorensen et al. 2002
unknown	<i>GNE1</i>	<i>gus-negative1</i>	similar to <i>fat tapetum</i>	unknown	unknown	Sorensen et al. 2002
unknown	<i>GNE4</i>	<i>gus-negative4</i>	similar to <i>fat tapetum</i>	unknown	unknown	Sorensen et al. 2002

Gene ID	Gene Name	Gene Description	Phenotype	Protein Function	Biological Process	References
At1g06520	AtGPA11	glycerol-3-phosphate acyl transferase1	abnormal tapetum, defects in pollen wall formation	membrane bound glycerol-3-phosphate acyl transferase	unknown	Zheng et al. 2003
At1g07340	AtSTP2	atstp2	not observed ?	monosaccharide carrier, sugar transporter	uptake of glucose from degrading callose	Truernit et al. 1999
At1g13330	AHP2	arabidopsis homologue pairing 2	reduced, fertility, defects in chrom. pairing and bivalent formation	similar to yeast protein involved in homolog pairing, HP2 homolog	involved in chromosome synapsis and bivalent formation	Schommer et al. 2003
At1g14750	SDS	solo dancer	reduced fertility, defective in synapsis and bivalent formation	meiotic cyclin-like protein	involved in chromosome synapsis and recombination	Azumi et al. 2002
At1g63990	SPO11-2	sporulation11-2	male and female meiosis defects	homolog of topoisomerase subunit	homologue synapsis, recombination and bivalent formation	Keeney et al. 1997
At1g66170	DUET*	duet	defects in meiosis telophase II, pollen abortion	PHD-domain containing protein	regulation of meiotic cell cycle progression	Reddy et al. 2003, Yang et al. 2003
At1g66170	MMD1*	male meiocyte death1	defects in meiosis telophase II, pollen abortion	PHD-domain containing protein	regulation of meiotic cell cycle progression	Reddy et al. 2003, Yang et al. 2003
At1g67370	ASY1	asynaptic1	reduced, fertility, meiotic defects, reduced fertility	homolog to HOP1 important in chrom. synapsis in yeast	involved in chromosome synapsis and recombination	Ross et al. 1997, Armstrong et al. 2002
At1g75950	ASK1	arabidopsis skp1-like1	polyads production of variable size and content	homologous to yeast SKP1 subunit of SCF complex	chromosome separation and segregation	Yang et al. 1999
At1g77320	MEI1	meiosis defective1	abnormal tetrad, variable size and content	BRCT-domain-containing protein, similar to acrosin-trypsin inhibitor	involved meiotic cell cycle	He and Mascarenhas 1998
At1g77390	TAM1	tardy asynchronous meiosis1	defects in timing of meiotic division	core cell cycle protein involved in meiosis II during microsporogenesis	may control entry into metaphase II	Magnard et al. 2001, Wang et al. 2004
At2g31970	ARAD50	atrads50	chromosome fragmentation, hypersensitivity to DNA damage	homolog of RAD50, component of meiotic recombination complex	involved in chromosome synapsis and recombination	Gallego et al. 2001
At3g09090	DEX1	defective in exine formation1	defective exine, microspore degeneration	hypoth. membrane and calcium bonding protein	involved in sporopollenin deposition	Paxson-Souders 1997
At3g11440	MYB65	myb65	myb64, myb33	nucleus, pollen sperm cell differentiation	regulation of gene expression	Millar and Gubler 2005
At3g11980	MS2	male sterile2	thin pollen wall, reduce exine layer	similar to yoyoba protein, fatty acid reductase	may promote synthesis of long chain aliphatic molecules	Aarts et al. 1997
At3g13170	AtSPO11-1	atspo11-1	production of polyads, defective chromosome pairing	homolog of the archaeal DNA topoisomerase VIA subunit (topo VIA)	involved in chromosome synapsis and recombination	Grelon et al. 2001
At3g22880	AtDMC1	atdmc1	low male fertility, defects in bivalent formation	similar to meiosis-specific yeast DMC gene	bivalent formation	Couteau et al. 1999
At3g25100	AtCDC45	atcdc45	polyads production, reduced male fertility	meiotic regulator	involved in DNA synthesis	Stevens et al. 2004
At3g43210	STD*	stud	blocked meiotic cytokinesis, failure in callose wall formation after PMII	kinesin with n-terminal motor domain	regulation of cytokinesis after meiosis	Hulskamp et al. 1997, Spielman et al. 1997
At3g43210	TES*	tetraspore	giant microspore with 4 nuclei, blocked meiotic cytokinesis	kinesin with n-terminal motor domain	regulation of cytokinesis after meiosis	Hulskamp et al. 1997, Spielman et al. 1997
At4g14220	RHF1	ring-h2 group f1	male/female gamet formation	RING-type E3 ubiquitin ligase implicated in gametogenesis	cell cycle regulation	Liu et al. 2008
At4g20900	MSS/TDM1	male sterile5/three-division mutant	microspores in polyads, degeneration, third division after meiosis II without DNA replication	similarity with SC protein and regulatory subunit of CDK	meiosis/cytokinesis control	Glover et al. 1998, Sanders et al. 1999
At4g21270	KATA/ATK1	arab. thaliana kinesin1	abnormal meiosis	kinesin-like protein	possibly involved in meiotic spindle formation	Chen et al. 2002
At4g34990	MYB32	Myb32	defective in meiosis, abnormal microspores, partial male sterility	R2R3 MYB transcription factor	highly expressed in tapetum,	Preston et al. 2004
At5g05490	SYN1/DIF1	determinate infertile1	multiple meiotic defects, abnormal chromosome pairing	homolog to yeast meiotic-specific adhesion subunit	role in sister chromosome cohesion	Bai et al. 1999, Bhatt et al. 1999
At5g06100	MYB33	myb33	double mutant-premeiotic abortion of meiocytes	membrane/transporter protein	during anther development	Millar and Gubler 2005
At5g13390	NEF1	no exine formation1	lack of sporopollenin deposition, reduce lipid content in tapetal plastids		accumulation/transport of lipids in tapetum	Arizumi et al. 2004
At5g20850	ARAD51	atrads51	low male fertility, defects in chromosome pairing	encodes a homolog of yeast RAD51	chromosome repairing, synapsis	Li et al. 2004
At5g22000	RHF2	ring-h2 group f2	male/female gamet formation	RING-type E3 ubiquitin ligase implicated in gametogenesis	cell cycle regulation, targeting of ICK4/KRD6, a CDK inhibitor	Liu et al. 2008

microsporogenesis

Gene ID	Gene Name	Gene Description	Phenotype	Protein Function	Cellular Location	Reference
At1g14830	<i>ADL1C</i>	<i>Arabidopsis dynamin-like1C</i>	pollen abortion during maturation	dynammin-like protein	pollen plasma membrane maintenance, cell wall	Kang et al. 2003
At1g19890	<i>MGH3</i>	<i>male-gamete-specific histone H3</i>	no phenotype due to redundancy	male-germline-specific histone H3.3	germ cell and sperm cell specific gene expression	Okada et al. 2005, Ingouff et al. 2007
At1g24520	<i>AGP1</i>	<i>arabinogalactan protein1 (AtBACP1)</i>	pollen aborts and collapsed at bicellular stage	arabinogalactan protein	tapetum, microspore and bicellular pollen viability	Xu et al. 1995
At1g30450	<i>HAP5</i>	<i>hapless5</i>	aborted mature pollen	cation-chloride cotransporter	ion homeostasis during development	Johnson et al. 2004
At1g50230	<i>TIO/SOLO POLLEN</i>	<i>two-in-one/solo pollen</i>	failure of cytokinesis at PMI	homology to Ser/Thr protein kinase FUSED	essential in centrifugal cell plate expansion	Oh et al. 2005
At1g64570	<i>DUO3</i>	<i>duo pollen3</i>	germ cell fails to divide or delay	nuclear regulatory protein	required for G2/M transition	Brownfield et al. 2009
At1g65470	<i>FAS1</i>	<i>fasciata1</i>	microspore and germ cycle arrested at G2/M	chromatin assembly factor 1 (CAF-1)	nucleosome/chromatin assembly during S phase	Chen et al. 2008
At1g78900	<i>VHA-A</i>	<i>vacuolar ATPase V1 subunit A</i>	pollen aborted from bicellular stage, swollen ER	vacuolar-ATPase V1 subunit A	Golgi organization, Ph homeostasis, secondary active transport	Dettmer et al. 2005
At2g21870	<i>MGP1</i>	<i>gametophyte defective1</i>	destruction of mitochondria, pollen death	subunit of F1F0 ATP-synthase	co-regulation of ATP-synthase activity	Li et al. 2010
At2g29980	<i>FAD3</i>	<i>fatty acid desaturase3</i>	<i>fad3 fad7</i> triple mutant:	ER enzyme responsible for the synthesis of 18:3 fatty acids	omega-3 fatty acid desaturase activity	McConn and Browse 1996
At2g34550	<i>UNG4</i>	<i>ungud4</i>	pollen aborts at bicellular stage	GA-2-oxidase	regulation of gibberelins	Lalanne et al. 2004b
At2g35630	<i>GEM1</i>	<i>gemini pollen1</i>	binucleate pollen, defective cytokinesis at PMI	MOR1/GEM1 microtubule associated protein	microspore polarity/cytokinesis	Park 1998, 2001, Twell et al 2002, Lee et al. 2007
At2g42380	<i>AtZFP34</i>	<i>atzfp34</i>	defects in cell wall patterning and lipid metabol.	BZIP transcription factor	Regulation of gene expression	Giblová et al. 2009
At3g11170	<i>FAD7</i>	<i>fatty acid desaturase7</i>	pollen abortion	chloroplastic enzyme responsible for the synthesis of 16:3 and 18:3 fatty acids	omega-3 fatty acid desaturase activity	McConn and Browse 1996
At3g12280	<i>RBR</i>	<i>retinoblastoma-related</i>	microspore proliferation, cell fate defects	cell-cycle repressor protein	G1/S transition of mitotic cell cycle, cell fate specification	Chen et al. 2009
At3g23670	<i>Kinesin-12Bk1</i>	<i>kinesin-12b</i>	microspores fail to complete PMI with spindle defects	kinesin-12 family	phragmoplast microtubule formation	Lee et al. 2007
At3g45150	<i>TCP16</i>	<i>tcp16</i>	microspore nuclear DNA loss and abortion	bHLH protein, TCP PCF-subfamily	regulator of microspore gene expression	Takeda et al. 2006
At3g48750	<i>CDK4;1</i>	<i>cyclin-dependent kinase A-1</i>	bicellular pollen, germ cell division fails, S-phase delayed	cyclin-dependent kinase;1	cell cycle regulation, germ cell S-phase progression	Iwakawa et al. 2006, Nowak et al. 2006
At3g54650	<i>FBL17</i>	<i>F-box-like17</i>	bicellular pollen, S-phase inhibited	E3 ubiquitin protein ligase, F-box protein	targeted proteolysis of CDKA inhibitor KRP6	Kim et al. 2008
At3g57390	<i>AGL18</i>	<i>agl18</i>	not given	MADS-box containing protein	potential role in microspore gene expression	Alvarez-Buylla et al. 2000, Varelt et al. 2007
At3g60460	<i>DUO1</i>	<i>duo pollen1</i>	mature pollen at bicellular stage, germ cell fails to enter PMI	novel R2R3 MYB protein in GC (MYB125)	regulation of gene expression for PMII entry, GC division	Twell and Howden 1998ab, Durbarry et al. 2005, Rotman et al. 2005
At4g14450	<i>Kinesin-12A</i>	<i>kinesin-12a</i>	microspores fail to complete PMI with phragmoplast defects,	kinesin-12 family, microtubular motor kinesin	phragmoplast microtubule formation	Lee et al. 2007
At4g26440	<i>WRKY34</i>	<i>wrky34</i>	less sensitive to cold stress	WRKY transcription factor	mediates cold sensitivity	Honyts et al. 2006, Zou et al. 2010
At4g36900	<i>HAP12</i>	<i>hapless12</i>	aborted mature pollen	contains AP2 domain	refulator of pollen gene expression	Johnson et al. 2004
At5g05580	<i>FAD8</i>	<i>fatty acid desaturase8</i>	reduced germination and PT growth	temperature sensitive plastidic fatty acid desaturase	omega-3 fatty acid desaturase activity	McConn and Browse 1996
At5g22260	<i>MS1</i>	<i>male sterile1</i>	microspore degeneration after releasing from tetrad	PHD finger domain	required for early microspore development	Yang et al. 2007
At5g39400	<i>PTEN</i>	<i>arabidopsis phosphatase, tensin homologue</i>	pollen death after pollen mitosis II	phosphatase and tensin homologue (tyrosine/PIP3 phosphatase)	pollen maturation	Gupta et al. 2002
At5g44860	<i>AGM</i>	<i>abnormal gametophytes</i>	pollen degenerates at late microspore stage	putative transmembrane protein	microspore development and/or division	Sorensen et al. 2004
At5g49150	<i>GEX2</i>	<i>gamete expressed2</i>	affected pollen tube growth and fertility	plasma membrane associated, 6 transmembrane domains predicted	potentially involved in pollen tube guidance and fertilization	Engel et al. 2005

pollen development

Gene ID	Gene Name	Gene Description	Phenotype	Function	Protein Description	Role	Reference
At5g1330	SWITCH1/DYAD	<i>switch1/dyad</i>	uneven segregation of chromatids	member of sister chromatid complex	member of sister chromatid complex	role in sister chromosome cohesion	Siddiqi et al. 2000, Mercier et al. 2001
At5g54260	AtMRE11	<i>atmre11</i>	chromosome fragmentation, hypersensitivity to DNA damage	DNA repair and meiotic recombination protein	DNA repair and meiotic recombination protein	double strand break repair	Puizina et al. 2004
At5g54800	GPT1	<i>glucose-6-phosphate translocator1</i>	aborted pollen, reduced lipid bodies, vesicles and vacuoles	glucose-6-phosphate translocator	glucose-6-phosphate translocator	Glc6P import for plastic starch and fatty acids biosynthesis	Niewiadomski et al. 2005
At5g55490	GEX1	<i>gamete expressed1</i>	affected pollen tube growth and fertility	transmembrane domain containing protein expressed in sperm cells	transmembrane domain containing protein expressed in sperm cells	potentially involved in gamete guidance/interaction	Engel et al. 2005
At5g57800	FLP1	<i>faceless pollen1</i>	defective sporopollenin and abnormal tryphine	transmembrane protein with similarity to the sterol desaturase family	transmembrane protein with similarity to the sterol desaturase family	role in synthesis of tryphine and sporopollenin	Arizumi et al. 2003
At5g58230	MSI1	<i>multicopy suppressor of IRA1</i>	microspore and germ cycle arrested at G2/M	chromatin assembly factor 1 (CAF-1) p48 subunit/pRbAp48	chromatin assembly factor 1 (CAF-1) p48 subunit/pRbAp48	role in DNA replication	Chen et al. 2008
At5g5r350	AHA3	<i>arabidopsis H+-ATPase3</i>	collapsed and aborted mature pollen	plasma membrane K+-ATPase	plasma membrane K+-ATPase	microspore/pollen maturation	Robertson et al. 2004
At5g64630	FAS2	<i>fasciata2</i>	microspore and germ cycle arrested at G2/M	chromatin assembly factor 1 (CAF-1)	chromatin assembly factor 1 (CAF-1)	nucleosome/chrom. assembly during S phase	Chen et al. 2008
unknown	APG	<i>apg</i>	defective pollen wall	unknown	unknown	involved in intine wall formation	Roberts et al. 1991
unknown	DSY1	<i>dsynaptic1</i>	reduced fertility	unknown	unknown	involved in chromosome synapsis and recombination	Peirson et al. 1996, Ross et al. 1997
unknown	BCP1	<i>bcp1</i>	aborted pollen development, cell death	unknown	unknown	structural component of pollen wall	Xu et al. 1995
unknown	DSY10	<i>dsynaptic10</i>	male sterile, meiosis defects	unknown	unknown	involved in chromosome synapsis and recomb.	Ross et al. 1997
unknown	DUO2	<i>duo pollen2</i>	bicellular pollen grain, germ cell arrested at prometaphase	unknown	unknown	mitotic progression at PMII, prometaphase	Durbarry et al. 2005
unknown	GEM2	<i>geminipollen2</i>	defective cytokinesis at PMI, range of pollen phenotype	unknown	unknown	asymmetric division and cytokinesis at PMI	Park et al. 2004
unknown	GUM	<i>germ unit malformed</i>	dislocation of the MGU	unknown	unknown	MGU organization and integrity	Lalanne and Twell 2002
unknown	LIP	<i>limpet pollen</i>	defective migration of GC into the centre after PMI	unknown	unknown	may regulate delivery of β -1,3-glucans/factor involved in cell migration	Howden et al. 1998
unknown	MAD1	<i>male gametophytic defective1</i>	variable phenotype of generative cell	unknown	unknown	essential for PMII, GC division	Grini et al. 1999
unknown	MAD2	<i>male gametophytic defective2</i>	generative cell defects	unknown	unknown	essential for PMII, GC division	Grini et al. 1999
unknown	MAD3	<i>male gametophytic defective3</i>	formation of intine wall, PMI defect, double DNA content	unknown	unknown	essential for PMI, wall patterning	Grini et al. 1999
unknown	MS33	<i>male sterile33</i>	defect in intine formation and tryphine deposition, abnormal pollen desiccation	unknown	unknown	unknown	Fei and Sawhney 2001
unknown	MUD	<i>mgu displaced</i>	dislocation of the MGU	unknown	unknown	MGU organization and integrity	Chen and McCormick 1996
unknown	SCP	<i>sidecar pollen</i>	extra celled-pollen, abnormal divisions at PMI	unknown	unknown	involved in pollen asymmetric division	Chen and McCormick 1996
At1g02140	HAP1	<i>hapless1</i>	disrupted guidance of pollentube growth, failure to leave the septum	high homology to Mago nashi	high homology to Mago nashi	mRNA metabolism/localization	Johnson et al. 2004
At1g04950	AtTAF6	<i>ataf6</i>	short pollen tube growth	TBP-associated factor 6	TBP-associated factor 6	basal transcription factor	Lago et al. 2005
At1g20200	HAP15	<i>hapless15</i>	short pollen tube growth, failure to exit style	26S proteasome regul. subunit protein S3	26S proteasome regul. subunit protein S3	protein degradation	Johnson et al. 2004
At1g60780	HAP13	<i>hapless13</i>	short pollen tube growth, failure to exit style	clathrin adaptor complexes medium subunit family protein	clathrin adaptor complexes medium subunit family protein	membrane trafficking	Johnson et al. 2004
At1g66570/	HAP3	<i>hapless3</i>	short pollen tube growth,	sucrose transporter SUC1/	sucrose transporter SUC1/	saccharide metabolism/	Johnson et al. 2004
At1g71270	POK	<i>poly pollen tube</i>	short pollen tubes	Vps52/Sac2 family protein	Vps52/Sac2 family protein	polar tube growth, component of vesicle transport machinery	Lobstein et al. 2004
At1g71880	AtSUC1	<i>sucrose-proton symporter</i>	pollen tube growth arrest	plasma membrane-H ⁺ -sucrose symporter	plasma membrane-H ⁺ -sucrose symporter	sucrose import/cell specific modulation of water potential	Stadler et al. 1999, Feuerstein et al. 2010
At2g03070	SETH10	<i>seth10</i>	reduced competitive ability, slower pt growth	putative protein	putative protein	unknown	Lalanne et al. 2004b

pollen tube growth													
At2g17800	Rop3At	rop3at	inhibition of pollen tube growth	member of ROP GTPase family	regulation of pollen tube growth	Li et al. 1998, 1999							
At2g25600	ATK6	spik	impaired pollen tube growth	shaker family inward K ⁺ channel	ion transport	Mouline et al. 2002							
At2g34680	UNG9	ungu9	reduced male transission	auxin-induced protein (AIR9), similar to leucine-rich repeat family protein	cell wall metaolism/signalling	Lalanne et al. 2004b							
At2g34980	SETH1	seth1	strongly reduced germination, abnormal callose deposition	phosphatidylinositol-glycan synthase subunit C	Component of GPI-anchor biosynthetic pathway	Lalanne et al. 2004b							
At2g35650	AtCSLA7	cellulose synthase-like7	slower pollen tube growth, defects in embryo development	glycosyltransferase	polysaccharides	Goubet et al. 2003							
At2g41930	SETH7	seth7	strongly reduced pollen germination and pollen tube growth	Ser/Thr protein kinase	initiation of pollen germination	Lalanne et al. 2004b							
At2g43040	MPG1	no pollen germination	nongerminating pollen, no pollen tubes	calmodulin-binding protein	binding to calmodulin in calcium-dependent manner	Golovkin and Reddy 2003							
At2g47040	VGDI	vanguard1	retarded pollen tube growth	pectine methyltransferase-homologous protein	cell wall metaolism	Jiang et al. 2005							
At2g47860	SETH6	seth6	strongly reduced pol. germination and ptgrowth	RPT2/NPH3-like protein	scaffold protein within signalling pathway	Lalanne et al. 2004b							
At3g04080H	AtAPY1	apyrase1+2	nongerminating pollen,	apyrase	mediation of g'ignalling +	Stiebrunner et al. 2003							
At3g08970	TMS1	thermosensitive male sterile1	lowered thermotolerance of pollen tube	DnaJ domain protein localized in ER lumen, homology to Hsp40	required for thermotolerance in progamic phase	Yang et al. 2009							
At3g10380	SEC8	exocyst complex	nongerminating pollen, no pollen tubes	subunits of exocyst complex	polarised exocytosis ofsecretory vesicles	Cole et al. 2005							
At3g45100	SETH2	seth2	strongly reduced germination, abnormal callose deposition	GPI-GnT catalytic subunit PIG-A	Component of GPI-anchor biosynthetic pathway	Lalanne et al. 2004ab							
At3g51300	Rop1At	rop1at	inhibition of pollen tube growth	pollen-specific Rop GTPase	intracellular Ca ²⁺ -gradient, regulation of pollen tube growth	Li et al. 1998, 1999							
At3g52590	HAP4	hapless4	chaotic pollen tube growth in the ovary	ubiquitin extension protein1 (UBQ1)/60S ribosomal protein L40	preteosynthesis/ubiquitination	Johnson et al. 2004							
At3g54090	UNG10	ungud10	reducedmale transmittsion	fructokinase-like protein	sugar metabolism	Lalanne et al. 2004b							
At3g54690	SETH3	seth3	reduced pollen germination and pt growth	arabinose-5-phosphate isomerase	synthesis of rare rhamnogalactouronans	Lalanne et al. 2004b							
At4g00800	SETH5	seth5	strongly reduced pol. germination and growth	unknown transmembrane protein	unknown	Lalanne et al. 2004b							
At4g04710	UNG6	ungud6	reduced male transission	calcium-dependent protein kinase	calcium-dependent signalling	Lalanne et al. 2004b							
At4g11720	HAP2/GCS1	hapless2	disrupter pollen tube guidance, blocked gamete fusion and fertilization	GCS1-like hypothetical protein, likely transmembrane protein	unknown	Johnson et al. 2004, von Besser et al. 2006							
At4g21150	HAP6	hapless6	short pollen tube growth, failure to exit style	ribophorin II (RPN2) family protein	membrane trafficking	Johnson et al. 2004							
At4g34940	SETH4	seth4	strongly reduced pollen germination	ARM repeat containing protein	unknown	Lalanne et al. 2004b							
At4g35950	Rop5At	rop5at	inhibition of pollen tube growth	member of ROP GTPases gene family-like	regulation of pollen tube growth	Li et al. 1998, 1999							
At5g12250	SETH9	seth9	reduced competitive ability, slower pollen tube growth	alcohol dehydrogenase-like protein	alcohol metabolism	Lalanne et al. 2004b							
At5g13650	SETH8	seth8	reduced competitive ability, slower pt growth	GTP-binding tyra-related protein	unknown	Lalanne et al. 2004b							
At5g18280	AtAPY2	double mutant	no pollen tube	apyrase	nucleotide metabolism	Stiebrunner et al. 2003							
At5g47020/	HAP11	hapless11	normal pollen tube growth,	unknown protein/	unknown/	Johnson et al. 2004							
At5g47030	KIP	kinky pollen	failure to enter the micropyle	mitochondrial ATP synthase δ chain	energy metabolism	Johnson et al. 2004							
At5g49680	HAP8	hapless8	kinky-shaped pollen tubes	SABRE-like protein	component of the secretory pathway	Procissi et al. 2003							
At5g56250	unknown	mad4	short pollen tube growth, failure to exit style	expressed protein	unknown	Johnson et al. 2004							
unknown			defective pollen tube elongation	unknown	unknown								
unknown	RTG	raring-to-go	pollen germinated precociously within the anther	unknown	unknown	Johnson and McCormic 2001							

Honys D, Reňák D, Twell D.

Male gametophyte development and function.

Global Science Books, pp.76-87, 2007.

Abstract:

Male gametophyte development in higher plants is a complex process that requires the coordinated participation of various cell and tissue types and their associated specific gene expression patterns. The male gametophytic life cycle can be divided into a developmental phase leading to the formation of mature pollen grains, and a functional or progamic phase, beginning with the impact of the grains on the stigma surface and ending at double fertilisation. Pollen ontogeny is also an excellent model in which to dissect the cellular networks that control cell growth, polarity, cellular differentiation and cell signaling. Here we present an overview of important cellular processes in male gametophyte development and recent advances in genetics and genomic approaches that are advancing the field significantly. Genetic approaches have identified a growing number of gametophytic mutants affecting discrete steps during the developmental or progamic phases that are now beginning to uncover some of the key molecular processes involved. With recent technological advances pollen transcriptomic studies now provide the first genome-wide view of male reproductive cell development in *Arabidopsis*. These studies reveal at least two successive global gene expression programs and the identity of a large number of male gametophyte-specific genes and putative transcriptional regulators. Transcriptome analysis has revealed a striking overrepresentation of cell wall metabolism, cytoskeleton and signaling genes in preparation for the progamic phase. This quantum leap in gene-centered knowledge highlights the functional specialization of this pathway and offers many new opportunities for the dissection of cellular processes that control male reproductive success.

Abstrakt:

Vývoj samčího gametofytu u vyšších rostlin je složitý vývojový proces, který vyžaduje koordinovanou účast rozličných buněk a pletiv a jejich specifickou genovou expresí. Životní cyklus vývoje samčího gametofytu může být rozdělen do několika vývojových fází vedoucích k produkci zralých pylových zrn a k progamické fázi, která začíná po dopadu pylového zrna na bliznu a jeho klíčení a růst v pylovou láčku zajišťující dvojité oplození samičího vajíčka. Vývoj pylu je také výjimečným modelem umožňujícím studium kontroly buněčného růstu, polarity, diferenciaci a buněčné signalizace. Toto review se zabývá důležitými životními procesy během vývoje samčího gametofytu a pokroky v genetice a genomice na tomto poli. Bylo identifikováno množství gametofytických mutací postihujících jednotlivé fáze vývoje pylu a pylové láčky, které jsou postupně identifikovány na molekulární úrovni. S využitím nejnovějších technologických přístupů, transkriptomika pylu nyní poskytuje první ucelený pohled v širí celého genomu na vývoji samčího gametofytu u *Arabidopsis*. Tato studia odhalila přinejmenším dva následné expresní vývojové programy a velké množství pylově specifických genů a předpokládaných transkripčních regulátorů. Transkriptomická studia ukázala překvapující vysokou expresi genů účastnících se metabolismu buněčné stěny, cytoskeletu a signalizace, jakožto příprava na pozdější progamickou fázi. Tyto poznatky ukazují na vysokou funkční specializaci tohoto vývoje a nabízí mnoho nových příležitostí ke studiu buněčných procesů, které řídí samčí reprodukční úspěšnost.

Reňák D, Dupláková N, Honys D.

Wide-scale screening of T-DNA lines for transcription factor genes affecting male gametophyte development in *Arabidopsis*.

Plant Sex Rep, (2011), resubmitted.

Abstract:

Male gametophyte development leading to the formation of a mature pollen grain is precisely controlled at various levels during its whole progression. Transcriptomic studies exploiting genome-wide microarray technologies revealed the uniqueness of pollen transcriptome and the dynamics of early and late successive global gene expression programs. However, the knowledge of transcription regulation is still very limited. In this study, we focused on the identification of pollen-expressed transcription factor (TF) genes involved in the regulation of male gametophyte development. To achieve this, reverse genetic approach was used. 74 T-DNA insertion lines were screened representing 49 genes of 21 TF families active in either early or late pollen development. In the screen, ten phenotype categories were distinguished affecting various structural or functional aspects including pollen abortion, presence of inclusions, variable pollen grain size, disrupted cell wall structure, cell cycle defects and male germ unit organization. Thirteen lines were not confirmed to contain the T-DNA insertion. Among 61 confirmed lines, about a half (29 lines) showed strong phenotypic changes (i.e. $\geq 25\%$ aberrant pollen) including four lines producing remarkable high proportion (70-100%) of disturbed pollen. However, remaining 32 lines exhibited mild defects or resembled wild type appearance. There was no significant bias towards any phenotype category amongst early and late TF genes and interestingly within individual TF families. Presented results have a potential to serve as a basal information resource for future research on the importance of respective TFs in male gametophyte development.

Abstrakt:

Vývoj samčího gametofytu vedoucí k produkci zralých pylových zrn je precizně regulován na různých úrovních během celého vývoje. Transkriptomická studia odhalila jedinečnost pylové genové exprese a její dynamiku během raného i pozdního vývojového programu. Přese všechno, znalost regulace transkripce je stále velmi omezené. V této studii se zaměřujeme na identifikaci pylově exprimovaných transkripčních faktorů, které hrají úlohu v regulaci vývoje samčího gametofytu. K tomuto účelu byl využit přístup reverzní genetiky. Bylo analyzováno 74 T-DNA linií odpovídající 49 sledovaným genům z 21 rodin transkripčních faktorů aktivních jak v raných tak pozdních stádiích vývoje pylu. Bylo detekováno 10 fenotypových kategorií vyznačujícími se strukturními i funkčními poruchami vývoje, jako jsou: aborce, přítomnost inkluzí, různá velikost pylových zrn, poškozená struktura buněčné stěny, defekty v buněčném cyklu a organizaci vegetativního jádra a spermatických buněk. U 13 linií nebyl T-DNA inzert detekován. Mezi 61 potvrzenými liniemi asi polovina (29 linií) ukázala významné fenotypové změny (tj. $\geq 25\%$ poškozeného pylu) včetně čtyř linií vyznačujícími se vysokou produkcí (70-100%) poškozeného pylu. Nicméně zbývající 32 linie ukazovaly jen mírně poškozený pyl nebo vzhledově připomínající pyl divokých rostlin. Nebyla shledána žádná souvislost mezi fenotypovou kategorií a mutanty genů raných a pozdních TF ani genovou rodinou TF. Předkládané výsledky práce mohou být dobře využity jako informační základna k dalším detailním analýzám významnosti vybraných TF ve vývoji samčího gametofytu.

Gibalová A, Reňák D, Matczuk K, Dupláková N, Cháb D, Twell D, Honys D.

***AtbZIP34* is required for Arabidopsis pollen wall patterning and the control of several metabolic pathways in developing pollen.**

Plant Mol Biol (2009) 70: 581-601.

Abstract:

Sexual plant reproduction depends on the production and differentiation of functional gametes by the haploid gametophyte generation. Currently, we have a limited understanding of the regulatory mechanisms that have evolved to specify the gametophytic developmental programs. To unravel such mechanisms, it is necessary to identify transcription factors (TF) that are part of such haploid regulatory networks. Here we focus on bZIP TFs that have critical roles in plants, animals and other kingdoms. We report the functional characterization of *Arabidopsis thaliana* AtbZIP34 that is expressed in both gametophytic and surrounding sporophytic tissues during flower development. T-DNA insertion mutants in AtbZIP34 show pollen morphological defects that result in reduced pollen germination efficiency and slower pollen tube growth both in vitro and in vivo. Light and fluorescence microscopy revealed misshapen and misplaced nuclei with large lipid inclusions in the cytoplasm of *atbzip34* pollen. Scanning and transmission electron microscopy revealed defects in exine shape and micropatterning and a reduced endomembrane system. Several lines of evidence, including the AtbZIP34 expression pattern and the phenotypic defects observed, suggest a complex role in male reproductive development that involves a sporophytic role in exine patterning, and a sporophytic and/or gametophytic mode of action of AtbZIP34 in several metabolic pathways, namely regulation of lipid metabolism and/or cellular transport.

Abstrakt:

Sexuální reprodukce rostlin závisí na produkci a diferenciaci funkčních gamet produkovaných haploidní gametofytickou generací. Poznání regulačních mechanismů specifického gametofytického programu je v současné době stále omezené. Pro odhalení těchto mechanismů je nezbytná identifikace transkripčních faktorů (TF), které jsou součástí regulační sítě. V této práci jsme se zaměřili na bZIP TF, který hraje významnou úlohu ve vývoji rostlin, živočichů i dalších říších. V této práci předkládáme výsledky funkční charakterizace genu AtbZIP34 u *Arabidopsis*, který je exprimován jak v gametofytu tak i okolních sporofytických pletivech během vývoje květu. T-DNA inzerce v genu AtbZIP34 způsobuje morfologické defekty ve vývoji pylu, které zapříčiňují sníženou klíčivost pylu a omezenou rychlost růstu pylových láček jak v in vitro tak in vivo podmínkách. Světelná i fluorescenční mikroskopie odhalila změny v organizaci vegetativního jádra a spermatických buněk a velké lipidové inkluze v cytoplazmě mutantního pylu. Elektronová mikroskopie navíc odhalila změny ve stavbě exiny a redukci endomembránového systému. Několik takových důkazů včetně průběhu exprese genu AtbZIP34 a fenotypových defektů ukazuje na složitou roli daného faktoru v samčím reprodukčním vývoji, jak v úloze sporofytu při tvorbě exiny tak i kordinovanou úlohu sporofytu a gametofytu v několika metabolických drahách jako je regulace lipidového metabolismu a/nebo buněčného transportu.

Honyš D, Oh SA, Reňák D, Donders M, Šolcová B, Johnson JA, Boudová R, Twell D.
Identification of microspore-active promoters that allow targeted manipulation of gene expression at early stages of microgametogenesis in *Arabidopsis*.
BMC Plant Biol (2006) 6: 31-39.

Abstract:

Background: The effective functional analysis of male gametophyte development requires new tools enabling the spatially and temporally controlled expression of both marker genes and modified genes of interest. In particular, promoters driving expression at earlier developmental stages including microspores are required.

Results: Transcriptomic datasets covering four progressive stages of male gametophyte development in *Arabidopsis* were used to select candidate genes showing early expression profiles that were male gametophyte-specific. Promoter-GUS reporter analysis of candidate genes identified three promoters (MSP1, MSP2, and MSP3) that are active in microspores and are otherwise specific to the male gametophyte and tapetum. The *MSP1* and *MSP2* promoters were used to successfully complement and restore the male transmission of the gametophytic *two-in-one* (*tio*) mutant that is cytokinesis-defective at first microspore division.

Conclusion: We demonstrate the effective application of MSP promoters as tools that can be used to elucidate gametophytic gene functions in microspores in a male-specific manner.

Abstrakt:

Pozadí: Efektivní funkční analýza vývoje samčího gametofytu požaduje nové nástroje umožňující místně i časově specifickou genovou expresi jak markerových genů, tak vlastních studovaných genů. Pro tyto účely je nezbytná zejména exprese řízená promotory aktivními v raných vývojových stádiích.

Výsledky: Byla využita transkriptomická databáze zahrnující čtyři vývojová stadia vývoje samčího gametofytu *Arabidopsis* k výběru kandidátních genů s ranou a pylově specifickou expresí. S pomocí promotor-GUS reportérového systému byly identifikovány tři promotory (MSP1, MSP2, and MSP3) aktivní ve stádiu mikrospory ale i tapeta. Promotory MSP1 a MSP2 byly úspěšně využity v komplementaci a obnově transmise gamet prostřednictvím *two-in-one* (*tio*) mutantu, který je defektní v prvním dělení mikrospory.

Závěr: Prokázali jsme efektivní aplikaci MSP promotorů jakožto nástrojů, které mohou být použity k analýzám gametofytických genových funkcí ve stádiu mikrospory pylově specifickým způsobem.

Dupl'áková N, Reňák D, Patrik Hovanec, Honysová B, Twell D, Honys D.

Arabidopsis Gene Family Profiler (aGFP) – user-oriented transcriptomic database with easy-to-se graphic interface.

BMC Plant Biol (2007) 7: 39-46.

Abstract:

Background: Microarray technologies now belong to the standard functional genomics toolbox and have undergone massive development leading to increased genome coverage, accuracy and reliability. The number of experiments exploiting microarray technology has markedly increased in recent years. In parallel with the rapid accumulation of transcriptomic data, on-line analysis tools are being introduced to simplify their use. Global statistical data analysis methods contribute to the development of overall concepts about gene expression patterns and to query and compose working hypotheses. More recently, these applications are being supplemented with more specialized products offering visualization and specific data mining tools. We present a curated gene family-oriented gene expression database, Arabidopsis Gene Family Profiler (aGFP; <http://agfp.ueb.cas.cz>), which gives the user access to a large collection of normalised Affymetrix ATH1 microarray datasets. The database currently contains NASC Array and AtGenExpress transcriptomic datasets for various tissues at different developmental stages of wild type plants gathered from nearly 350 gene chips.

Results: The Arabidopsis GFP database has been designed as an easy-to-use tool for users needing an easily accessible resource for expression data of single genes, pre-defined gene families or custom gene sets, with the further possibility of keyword search. Arabidopsis Gene Family Profiler presents a user-friendly web interface using both graphic and text output. Data are stored at the MySQL server and individual queries are created in PHP script. The most distinguishable features of Arabidopsis Gene Family Profiler database are: 1) the presentation of normalized datasets (Affymetrix MAS algorithm and calculation of model-based gene-expression values based on the Perfect Match-only model); 2) the choice between two different normalization algorithms (Affymetrix MAS4 or MAS5 algorithms); 3) an intuitive interface; 4) an interactive "virtual plant" visualizing the spatial and developmental expression profiles of both gene families and individual genes.

Conclusion: Arabidopsis GFP gives users the possibility to analyze current Arabidopsis developmental transcriptomic data starting with simple global queries that can be expanded and further refined to visualize comparative and highly selective gene expression profiles.

Abstrakt:

Pozadí: DNA-čipové technologie nyní patří ke standardním genomickým nástrojům a jejich masivní rozvoj vedl ke zvýšení genomového pokrytí, přesnosti a spolehlivosti. Počet experimentů využívajících těchto technologií v posledních letech významně vzrůstá. Současně s tímto rozvojem stoupá množství on-line nástrojů zjednodušujících jejich následné využití. Globální statistické analýzy dat přispívají k rozvoji celkových pohledů na strukturu genové exprese a tvorbu pracovních hypotéz. V poslední době jsou tyto aplikace poskytovány se specializovanými produkty nabízejícími vizualizační a jiné nástroje ke zpracování dat. Nabízíme databázi orientovanou na transkripci genových rodin Arabidopsis Gene Family Profiler (aGFP; <http://agfp.ueb.cas.cz>), která nabízí uživateli přístup k široké sbírce

normalizovaných dat z Affymetrix ATH1. Databáze nabízí data z NASC Array a AtGenExpress zdrojů pokrývajících různá vývojová stadia divokých rostlin pocházejících z téměř 350 experimentů.

Výsledky: database aGFP byla designována jako jednoduše ovladatelná databáze se snadným přístupem k datům jednotlivých genů, genových rodin nebo zvolené skupiny genů s možností hledání podle klíčových slov. Arabidopsis Gene Family Profiler nabízí uživatelsky přístupné rozhraní s grafickým i textovým výstupem. Data jsou uložena na MySQL server a jednotlivé dotazy jsou tvořeny v PHP skriptu. Nejcharakterističtějšími rysy Arabidopsis Gene Family Profiler database jsou: 1) prezentace normalizovaných dat (Affymetrix MAS algoritmus a výpočet genových expresních dat podle Perfect Match-only model u); 2) výběr mezi dvěma odlišnými normalizovanými algoritmy (Affymetrix MAS4 or MAS5); 3) intuitivní rozhraní; 4) interaktivní virtuální rostlina znázorňující místně a vývojově expresní profil jak genových rodin, tak jednotlivých genů.

Závěr: Arabidopsis GFP nabízí uživateli možnost analyzovat současné transkriptomická data Arabidopsis začínajících u jednoduchých celkových dotazů, které mohou být dále rozšířeny a upřesněny k vizualizaci srovnávacích a vysoce selektivních genových profilů.

Honys D, Duplákova N, Reňák D.

Online tools for presentation and analysis of plant microarray data.

Nova Science Publishers, Inc. In: Oligonucleotide array sequence analysis, eds: M.K. Moretti and L.J. Rizzo, pp. 265-295, 2008.

Abstract:

The complete sequencing and annotation of the *Arabidopsis thaliana* genome represented a major step in biological research. This knowledge enabled gene prediction, assignment of functional categories and gave an opportunity to characterize global gene expression patterns at the transcriptome level at different developmental stages and under various physiological and stress conditions. For the discovery of partial or whole regulatory or functional networks, the development of high-throughput technologies was inevitable with genome-wide transcriptomic studies providing an essential input. DNA microarray technologies are among the most frequently used methods for parallel global analysis of gene expression. Microarray technologies now belong to the standard functional genomics toolbox and have undergone massive development leading to increased genome coverage, accuracy and reliability. Whole genome microarrays developed by Affymetrix in collaboration with Syngenta represented the first standard in genome-wide transcriptomic studies in plants. Whole genome Affymetrix ATH1 GeneChips covers about 76% of the *Arabidopsis thaliana* genome. Moreover, the introduction of the Minimum Information About Microarray experiments (MIAME) has increased the value and reproducibility of microarray experiments and has become a standard in documentation of array experiments and in the creation of databases of comparable transcriptomic experiments. The number of experiments exploiting microarray technology has markedly increased in recent years. Not surprisingly, there are potential difficulties in the orientation in available data sets. Microarray expression data are deposited on servers, many of which are publicly accessible. Currently, these databases store several thousands of individual datasets and some of these offer online tools for data normalization, filtering, statistical testing and pattern discovery. In parallel with the rapid accumulation of transcriptomic data, on-line analysis tools are being introduced to simplify their use. Global statistical data analysis methods contribute to the development of overall concepts about gene expression patterns and to query and compose working hypotheses. More recently, these applications are being supplemented with more specialized products offering visualization and specific data mining tools. In this chapter, an overview of available on-line databases and web-based applications using microarray data is presented together with the information about their structure and elementary principles.

Abstrakt:

Kompletní sekvence a anotace genomu *Arabidopsis thaliana* představuje velký pokrok v biologickém výzkumu. Tato znalost umožňuje předpověď genů, zařazení do funkční kategorie a nabízí možnost funkčně charakterizovat celkový profil genové exprese různých vývojových stádií během fyziologických i stresových podmínek. K objevení částečné nebo celkové regulační funkční sítě, vývoj přesných DNA čipových technologií byl nezbytný spolu s celogenomovými transkriptomickými studiemi. DNA čipové technologie patří k nejčastěji používaným metodám ke globální paralelní analýze genové exprese. Tyto technologie nyní patří ke standardním funkčním genomickým nástrojům, které prošly masivním vývojem vedoucím k vyšší genomové pokryvnosti, přesnosti a spolehlivosti. Vývoj

celogenomových DNA čipů pokrývá 75% genomu *Arabidopsis thaliana*. Navíc, zavedení Minimum Information About Microarray experiments (MIAME) umožnilo vzrůst jejich hodnoty a reproducibility a stal se standardem v dokumentaci těchto experimentů a tvorbě databází srovnatelných transkriptomických experimentů. Počet experimentů využívajících DNA čipy v posledních letech významně vzrostl. Není překvapením, že to znesnadňuje orientaci v dostupných datových zdrojích. Expresní data jsou ukládána na serverech a mnoho z nich jsou veřejně přístupná. V současné době tyto databáze obsahují data z několika tisíc experimentů, některé z nich nabízejí on-line nástroje k normalizaci dat, filtrování, statistickému testování a odhalování expresních profilů. Současně s hromaděním transkriptomických dat přibývá také on-line nástrojů k zjednodušení nakládání s nimi. Metody globální statistické analýzy dat přispěly k rozvoji celkových pohledů na strukturu genové exprese a tvorbu pracovních hypotéz. Tyto aplikace jsou v poslední době poskytovány s více specializovanými produkty nabízejícími vizualizační a speciálními nástroji k získávání dat. V této kapitole nabízíme přehled dostupných on-line databází a webových aplikací využívajících expresní data spolu s informací o jejich struktuře a základních principech.