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The making of gametes in higher plants

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ABSTRACT Higher plants have evolved to be one of the predominant life forms on this planet. A great deal of this evolutionary success relies in a very short gametophytic phase which underlies the sexual reproduction cycle. Sexual plant reproduction takes place in special organs of the flower. In most species the processes of gametogenesis, pollination, syngamy and embryogenesis are sequentially coordinated to give rise to a functional seed in a matter of few weeks. Any of these processes is so intricately complex and precisely regulated that it becomes no wonder that each involves more specific genes and cellular processes than any other function in the plant life cycle. While variability generation - the evolutionary output of the sexual cycle - is the same as in any other Kingdom, plants do it using a completely original set of mechanisms, many of which are not yet comprehended. In this paper, we cover the fundamental features of male and female gametogenesis. While the physiological and cellular bases of these processes have been continuously described since the early nineteenth century, recent usage of *Arabidopsis* and other species as central models has brought about a great deal of specific information regarding their genetic regulation. Transcriptomics has recently enlarged the repertoire and pollen became the first gametophyte to have a fully described transcriptome in plants. We thus place special emphasis on the way this newly accumulated genetic and transcriptional information impacts our current understanding of the mechanisms of gametogenesis.

KEY WORDS: *pollen, embryo sac, gametogenesis, microsporogenesis, macrosporogenesis*

The uniqueness of a life form

One of the most remarkable features of life on earth is diversity, a great deal of which is based on the evolutionary output of sexual reproduction. Unlike animals, in which the primordial germ line develops early during embryogenesis, higher plants alternate the growth of the diploid sporophyte organism with a highly reduced growth form on the plant life cycle, the haploid gametophyte. This is a well-suited strategy for selection because plants spend most of their life on the vegetative phase. The gametophytic stage also presents an opportunity for selection at the haploid level (Ottaviano *et al.*, 1990).

Plant cells don't move and positional information instead of lineage is the primary determinant of cell fate in plants. Meiosis triggers the separation between sporophytic and gametophytic generations involving various genes (Caryl *et al.*, 2003). This spatial pleiotropy of the sexual organs has prompted evolution for the appearance of safety mechanisms to prevent the fusion of incompatible genomes while supporting genetic variability. In higher animals genetic mechanisms for sex determination establish striking developmental differences between males and fe-

males. In contrast, most higher plant species develop both male and female structures within the same flower, allowing self-fertilization. Out-crossing is ensured by self-incompatibility mechanisms, which evolved under precise genetic control controlling self-recognition and cell-to-cell interaction (Thomas and Franklin-Tong, 2004).

The short period that plants spend on their reproductive cycle includes some of the most intriguing and challenging questions in biology, many yet without a mechanistic answer. With the publication of the complete genomic sequence of the flowering plant *Arabidopsis thaliana* (The *Arabidopsis* Genome Initiative, 2000), new foundations for functional and comparative genomics emerged to reveal basic genetic differences between plants and other eukaryotes. Although the logic underlying many of the developmental processes as pattern formation and cell-cell communication are comparable between plants and animals, the master regulators are unrelated or represent novel protein rearrangements which serve identical functions (Meyerowitz, 2002), making many genes unique to plants. It is therefore of no surprise that we find this prediction mirrored on the various stages of sexual plant reproduction.

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Gametogenesis in the major group of vascular plants: the angiosperms

The phylum *Anthophyta*, i.e. the Angiosperms (from the Latin word *angi*, “enclosed” and Greek word *sperma*, “seed”) or flowering plants, is the dominant group of land plants today. They arose about 140 million years ago during the late Jurassic and experienced rapid diversification during the Cretaceous. There are more than 250 000 species, grouped into 12 500 genera and about 300 families and in terms of ecological and nutritional importance they surpass all other groups. Numerous traits contributed to this rise to dominance, including the ability to reach reproductive maturity rapidly and adaptation to animal pollination and seed dispersal (Darwin, 1862). Flowers, the reproductive organs of angiosperms, are more varied than the equivalent structures of any other group of organisms (Barrett, 2002).

The life cycle of higher plants includes seed germination, vegetative growth, flowering, fertilization, development of embryo and seed maturation. Induction of flowering is triggered by environmental cues such as light, temperature and nutrient access, in combination with endogenous signals (e.g. the plant hormone gibberellin and circadian oscillators), for which many flowering time genes are described (Mouradov *et al.*, 2002). The integration of signals directs the activation of meristem identity genes, which specify floral identity and the shoot apical meristem is transformed into an inflorescence. Finally, the floral organ identity genes are activated in different regions of the flower, producing four organ types, sepals, petals, the male stamens, collectively named the androecium and the female carpels, making up the gynoecium, or the pistil, located in the innermost whorl (Smyth *et al.*, 1990). Such are the organs where male and female gametogenesis take place respectively.

The male gametophyte - pollen

The development of the male gametophyte involves a series of events culminating in the production and release of mature pollen grains from anthers (McCormick, 1993, 2004). Mature pollen grains may be considered as a “cell within a cell”, with very specialized functions. Enclosed by a very special cell wall, the vegetative cell harbors an individual cytoplasm with a particular gene expression pattern. Its final fate is to transport the two sperm cells through the female tissues and release them in the vicinity of the egg and central cells (see review by Boavida *et al.*, 2005).

It all begins in stamens, the male reproductive organs, which consist of an anther, where pollen development takes place and a stalk-like filament which provides support, nutrient transport and positions anthers to facilitate pollen dispersal (Scott *et al.*, 1991a; Irish, 1999). Stamens are derived from periclinal divisions of the L2 layer of the shoot apical meristem (Jenik and Irish, 2000).

Pollen development takes place within the anther locus, or pollen sac (Fig. 1D), involving perhaps more genes than any other single process in plant development and depends on interactions between sporophytic and gametophytic cells (Scott *et al.*, 2004). Both are essential for male fertility as revealed by numerous genetic screens (van der Meer *et al.*, 1992; Chen and McCormick, 1996; Aarts *et al.*, 1997; Hulskamp *et al.*, 1997; Taylor *et al.*, 1998; Ariizumi *et al.*, 2003).

Anther differentiation will lead to the production of several specialized cells and tissues involved in reproductive or non-reproductive functions (e.g. support and dehiscence) (Koltunow *et al.*, 1990). The male gametogenesis starts from L2 cells of the shoot apical meristem, that divide to form a primary parietal cell (PPC) and the primary sporogeneous cells (PSC). The PPC later form several concentric layers on the anther wall, which differentiate into the endothecium, middle and the very special inner layers, known as the tapetum (Fig. 1A). PSC give rise to the microsporocytes. Coordination of the process is under tight genetic control (Goldberg *et al.*, 1993).

The development of viable pollen grains is dependent on the presence of the tapetum, which has roles in the nourishment of microspores, formation of exine and deposition of tryphine on the pollen wall (Koltunow *et al.*, 1990; Mariani *et al.*, 1990; Goldberg *et al.*, 1993). A summary of major developmental events on male gametogenesis with the correspondent anther differentiation stages is represented on Figure 1.

When microspores initiate meiosis, coenocytic masses are established by enlarged plasmodesmata, the cytomictic channels (Fig. 1B), which create a cytoplasmic continuum between groups of cells. This way, there is an effective synchronization during microsporogenesis, by allowing exchange of molecules and ions (Heslop-Harrison, 1966; Mascarenhas, 1975). Microsporocytes are encased in an impermeable β -1,3-glucan wall (callose), isolating the meiocytes from the sporophyte cells (Fig. 1E). The callosic wall was proposed to function as a sieve for large molecules whose influence could impair the commitment to an haploid genomic expression (Mackenzie *et al.*, 1967; Mascarenhas, 1975). However, the apparent ability of large molecules to trans-

Fig. 1 (opposite). A generalized overview of microsporogenesis and pollen development. Schematic representations were adapted from Goldberg *et al.* (1993) and McCormick (1993). C, connective; CCC, circular cell cluster; E, epidermis; En, endothecium; PG, pollen grain; PS, pollen sac; St, stamium; T, tapetum; Td, tetrads; V, vascular bundle. See text for description of developmental events. Individual steps are illustrated with transmitted electron microscopy (TEM) micrographs or TEM-prepared semi-thin sections seen with light microscopy of microsporogenesis of the orchid *Ophrys lutea* (Feijó, 1995). **(A)** Individualization of a group of sporogeneous cells by a callose envelope. These cells will divide to final size and enter male meiosis. **(B)** Meiocytes are connected by large channels, named cytomictic channels (CC), through which a general streaming, including organelles like plastids (PI) is believed to contribute to the synchronization of meiosis within the whole population. **(C)** Telophase II, with twin mitotic spindles (f1 and f2) within a callose enclosed tetrad. **(D)** Mid section of an immature flower showing the relative position of the male gametophyte forming organs, the pollen sacs (Pol1 and 2) and the female gametophyte forming organs, the ovules in the ovary (Ov). **(E)** Up until the end of meiosis, each microspore is enclosed on a callose wall and groups of microspores are individualized from surrounding tissues by thick callose walls (Cal), here stained with decolorized aniline blue. **(F)** Formation of individualized microspores within a meiotic tetrad (1,2,3,4). Note the synchronization of one mass of meiocytes in telophase II, while the upper mass is already synchronized in microspore individualization. **(G)** Polarization and vacuolization of the young microspore. The nucleus is strongly pulled out to one end of the cell, while the plastids are on the opposite end. Small vacuoles are formed to account for the enlargement of the cell and later will coalesce in a single big vacuole. **(H)** Synchronized pollen mitosis I. The mass of microspores on the right is still on anaphase and on the left cytokinesis already occurred (arrow), dividing a small generative cell and a big vegetative one. **(I-K)** Process of pinching-off the generative cell.

verse this barrier raised some doubts about its actual function (Scott *et al.*, 2004).

Two meiotic divisions transform then each microsporocyte (or pollen mother cell) into a tetrad of haploid microspores, each with its own callose envelop and all encased in the callose wall of the tetrad (Fig. 1 E,F). At this stage the cytomictic channels disappear, establishing an individual cytoplasm in each microspore cytoplasm. During meiosis the levels of rRNA and mRNA decrease dramatically and plastids and mitochondria populations undergo a cycle of dedifferentiation and replication. These changes at the cytoplasmic level are believed to reset the sporophytic program and launch the gametophytic functional program (Bird *et al.*, 1983). Pollen wall deposition also initiates and a prim(exine) layer develops around each microspore which will become visible at tetrad stage of pollen maturation (Heslop-Harrison, 1963, 1971)(Fig. 2B).

STUD (STD) and *TETRASPORE (TES)* genes were shown to be essential for cytokinesis. The *stud/tes* mutants form large tetranucleate microspores with a common cytoplasm, lacking intersporal walls. However, up to four pairs of sperm cells are observed in *stud* and *tes* pollen grains (Hulskamp *et al.*, 1997; Spielman *et al.*, 1997; Yang *et al.*, 2003), suggesting that meiotic cytokinesis and cytoplasmic isolation are not prerequisites to establish cytoplasmic polarity in each microspore.

In most species, dissolution of the callose wall by callases releases the haploid microspores from the tetrad (McCormick, 1993). The timing of callase secretion appears to be critical for normal pollen development (Frankel *et al.*, 1969; Worrall *et al.*, 1992). The *Arabidopsis* gene, *glucan synthase-like5 (AtGsl5)*, encoding a plasma membrane-localized protein homologous to yeast β -1,3-glucan synthase (callose synthase), has been shown to partially complement the yeast mutant. The gene is developmentally expressed at high levels in flowers and may represent the callose synthase required for deposition of callose in pollen (Ostergaard *et al.*, 2002). In tobacco, the downregulation of *TAG1* gene, which encodes a β -1,3-glucanase expressed in the tapetum, was shown not to affect tetrad dissolution and pollen development (Bucciaglia *et al.*, 2003). A candidate gene required for tetrad dissolution is *QUARTET (QRT)*. In *quartet (qrt)* mutants pollen grains are released as tetrads and callose patches are observed between the fused walls. However, the failure on microspore separation has been traced to the persistence of pectin components on the pollen walls (Preuss *et al.*, 1994; Rhee and Somerville, 1998). *QUARTET 3* was recently cloned, showing homology to an endo-polygalacturonase and expression on the tapetum; moreover, when expressed in yeast, it shows polygalacturonase activity (Rhee *et al.*, 2003), confirming

a role for pectins on tetrad integrity.

After release, haploid microspores undergo cytoplasmic reorganization, small vacuoles coalesce into a single vacuole, which polarizes most of the cytoplasm to one side of the cell periphery (Fig. 1G) and the nucleus on the other side, a process mediated by the cytoskeleton (Brown and Lemmon, 1991; Zonia *et al.*, 1999). This polarization leads to an asymmetric division called pollen mitosis I (PMI) (Fig. 1 G-J), essential for the differentiation of a big vegetative cell (VC) and a small generative cell (GC), which later will generate the sperm cells (Fig. 1 H-J). This division can be viewed as determinative since the two daughter cells will have different cell fates (Horvitz and Herskowitz, 1992). Models have been proposed to explain the different cell fates by polarised distribution of gametophytic regulatory factors, resulting in the repression of the vegetative cell specific genes on the generative cell as a result of asymmetric division (Eady *et al.*, 1995). These models led to the conventional assumption that the generative cell or sperm cell nuclei are transcriptionally repressed (McCormick, 1993).

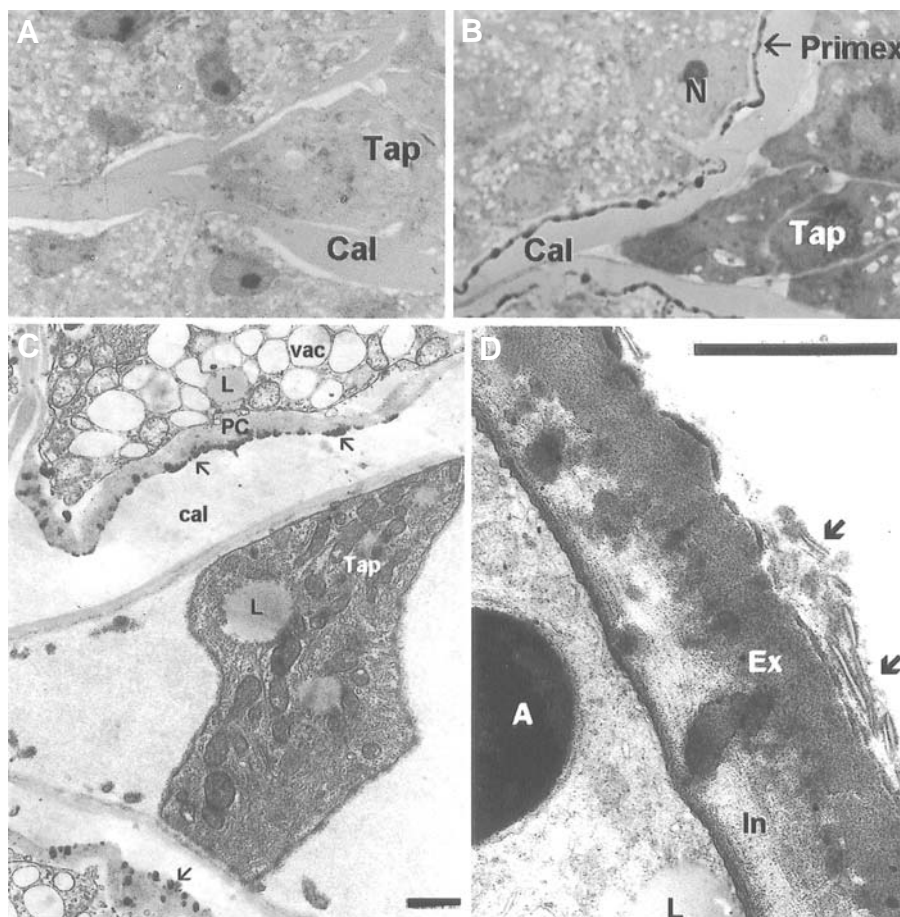


Fig. 2. Formation of the special pollen wall, the exine on *O. lutea*. (A) Young microspore stage, with active amoeboid tapetum cells (Tap) and a thick isolating callose layer (Cal). (B) Late microspore stage, with degenerating tapetum cells and beginning of deposition of the first layer of sporopollenin, named primexin (Primex). (C) Same stage observed with detail using TEM. The first deposition is basically organized in small spots, originating from tapetal secretion. Callose is still very thick. (D) Final structure of the pollen wall, with the pectocellulosic Intine (In), the sporopollenin Exine (Ex) and later deposited membrane like structure which may contain recognition proteins (arrows).

Genetics screens in *Arabidopsis* based on alterations to the typical tricellular pollen morphology yielded mutants affected on cell division. *Sidecar* (*scp*) was the first gametophytic mutant shown to affect microspore division symmetry: the microspore goes through a premature symmetrical division, but one of the daughter cells divides asymmetrically producing a normal VC with two sperm cells, supporting the hypothesis that an asymmetric localization of a polarity factor exists before PMI (Chen and McCormick, 1996). In *two-in-one pollen* (*tio*) mutant the mature pollen grains contain two nuclei, due to a failure of cytokinesis at pollen mitosis I (Twell and Howden, 1998; Twell *et al.*, 1998). However, *tio* microspores undergo maturation and are able to activate a vegetative cell fate marker, indicating that VC fate is the default program in absence of PMI. The *geminipollen 1* (*gem1*) mutant is affected in microspore polarity, asymmetric cell division and cell fate. The gene was shown to encode a microtubule associated protein required for the correct localization of the fragmoplast during cytokinesis (Park *et al.*, 1998; Twell *et al.*, 2002). After PMI the GC cytoplasm becomes isolated from the VC by a thin callose wall that fuses with the inner pollen wall, named intine (Fig. 1 I,J). This wall is then degraded by the mobilization of β -1,3-glucanases, allowing the GC to detach from the intine wall (Fig. 1 J,K). The round GC is then "engulfed" by the VC cytoplasm acquiring a central position on the pollen grain (Fig. 1K). Later the GC undergoes a morphogenetic process, acquiring an elongated or spindle-like shape, which is stabilized by the microtubule arrays aligned along the GC (Fig. 1 L-M). While not possessing a true cell wall, the mature GC does possess an external extracellular matrix of polysaccharide nature (Fig. 1L). On the gametophytic mutant *limpet pollen* (*lip*) GC migration is blocked after PMI and generative or sperm cells remain at the periphery of the pollen wall. Since in *limpet* pollen, the transient wall material that separates the GC from the VC persists, *LIP* is believed to either regulate the delivery of β -1,3 -glucanases or to encode general products required for the GC migration (Howden *et al.*, 1998).

In many species the generative cell divides before pollen release into two identical sperm cells (tricellular pollen), while in others this division only occurs after pollen germination (bicellular pollen). At the end of this division (PMII) the vegetative nucleus and the two sperm cells will assume a specific structural arrangement, the male germ unit (MGU). Recently, the *mud* (*male germ unit displaced*) and *gum* (*germ unit malformed*) male-specific gametophytic mutants were shown to affect the integrity and/or the positioning of the MGU in the mature pollen grain. These mutants show reduced transmission through the male gametophyte (Lalanne and Twell, 2002).

The final steps of pollen maturation are usually coincident with floral anthesis and anther dehiscence, ending with the release of mature dehydrated pollen grains. Anther dehiscence is coordinated with pollen differentiation and involves three major tissue types, the stomium, the endothecium and the circular cell cluster. Several mutants related to jasmonic acid biosynthesis were demonstrated to be affected in anther dehiscence and cause male sterility (Zhao and Ma, 2000; Ishiguro *et al.*, 2001; Park *et al.*, 2002; Hatakeyama *et al.*, 2003; Rieu *et al.*, 2003). A male sterile mutant in *Arabidopsis* affecting the *AtMYB26* transcription factor shows defects in anther dehiscence because the stamens lack cell wall fortifications in the endothelial cells, required for dehiscence (Steiner-Lange *et al.*, 2003). Pollen is, however, viable and

can fertilize when released mechanically from the anthers.

Mature pollen grains of angiosperms are released dehydrated (Fig. 1M), with a water content of 15-30%, which is associated with an almost inactive metabolism (Heslop-Harrison, 1979; Hoekstra and Bruinsma, 1980). Recently, the *ADL1C* gene, a member of the a dynamin-like protein family, was shown to be involved on the plasma membrane and intine morphology of post meiotic male gametophytes, with possible involvement on the formation and maintenance of the pollen surface and viability during desiccation (Kang *et al.*, 2003).

The pollen wall

Spores and pollen of higher plants have enormous value in taxonomic studies, due to the species-specific wall structure and surface pattern. The consistency of the pattern within species suggests a high degree of genetic regulation for which there is no obvious explanation. This issue raised an intensive debate whether the establishment of the pollen wall pattern is a consequence of a self-assembly process due to the physical and chemical behavior of the components and their interactions, or if it depends only on a complex cooperation of cellular components (Sheldon and Dickinson, 1983; Fitzgerald and Knox, 1995; Southworth and Jernstedt, 1995; Paxson-Sowders *et al.*, 1997; Hemsley *et al.*, 2003).

Pollen wall formation begins soon after meiosis is complete (Fig. 2 A,B) and it continues during tetrad and vacuolated stages of microspores to almost completion on the first pollen mitosis with a major contribution from the tapetum (Fig. 2C). There are two major types of tapetum, the more primitive secretory type, considered to be the prevalent type in the majority of plants and the amoeboid type that extends to the microspores in the anther locule for presumable direct delivery of tapetal contents (Fig. 2 A,C) (Furness and Rudall, 2001). The tapetum transfers sporopollenin and tryphine precursors to the pollen wall. Several typical lipid bodies, known as Ubisch bodies or orbicules, found on the fluid of the pollen sacs, were shown to accumulate at the site of exine formation on the surface of the developing microspores after completion of meiosis (Fig. 2 B,C). Its function remains elusive, but they are thought to be either a by-product of tapetal metabolism, or to have a direct functional role on transport of sporopollenin precursors (Heslop-Harrison, 1968a; Staiger and Apel, 1993). Recently, a structural protein named RAFTIN isolated in wheat and rice was shown to be essential for pollen development and it accumulates in Ubisch bodies targeted to microspore exine (Wang *et al.*, 2003).

Angiosperm pollen walls consist of several layers of chemically different materials, an outer sporopollenin layer, the exine, which consists of two sublayers the sexine and the nexine and a pectocellulosic innermost layer, the intine (Fig. 2D).

Intine wall synthesis starts during the free microspore stage (Knox and Heslop-Harrison, 1970) presumably under gametophytic control (Fig. 2 B,C). The intine is composed of cellulose-like components, enzymes and other proteins like AGP-related glycoproteins (Knox and Heslop-Harrison, 1970; Vithanage and Knox, 1976; Vaughn, 1982; Dodds *et al.*, 1993; Hiscock *et al.*, 1994; Li *et al.*, 1995; Aouali *et al.*, 2001).

Exine formation begins with the synthesis of a cellulosic matrix, the primexine, which accumulates as a homogeneous layer

between the plasma membrane and the callose wall of the microspore, except on areas destined to be pollen apertures (Fig. 2 B,C). The position of pollen apertures is believed to be set during microsporocyte cytokinesis (Heslop-Harrison, 1968b). The primexine precursors are synthesized and secreted by the microsporocytes to the pollen surface (Scott *et al.*, 1991a; Fitzgerald and Knox, 1995) and may work as a scaffold for polymerization of fatty acids and phenolics on the surface (Sheldon and Dickinson, 1983; Scott *et al.*, 1991a; Scott, 1994). The chemical composition of the exine is not exactly known, partly due to the high stability and chemical resistant composition of sporopollenin. Various biochemical analyses revealed a mixture of biopolymers of lipids, long chain fatty acids, phenylpropanoids, phenolics and carotenoids (Brooks and Shaw, 1968; Piffanelli *et al.*, 1997; Piffanelli *et al.*, 1998; Wang *et al.*, 2002; Ahlers *et al.*, 2003). Evans *et al.* (1992) have shown that a high rate of lipid biosynthesis in pollen grains start after PMI, later corroborated by the high expression levels observed in the tapetum for several enzymes related to lipid biosynthesis in *Brassica napus* (Piffanelli *et al.*, 1997). The *Arabidopsis male sterile (ms2)* mutant produces non-viable pollen grains with very thin walls, lacking exine. The *MS2* gene is suggested to encode a fatty acyl reductase which converts wax fatty acids to fatty alcohols, with expression observed in the tapetum shortly after the release of microspores from tetrads (Aarts *et al.*, 1997). The mutant *faceless pollen-1 (flp1)*, exhibits conditional male sterility and the *FLP1* protein is likely to be involved in wax fatty acid biosynthesis required for the synthesis of tryphine, sporopollenin of exine and the wax components of stems and siliques (Ariizumi *et al.*, 2003).

Enzymes of the phenylpropanoid pathway also show high levels on tapetal cells (Herdt *et al.*, 1978; Shen and Hsu, 1992). Evidence supports the involvement of phenylpropanoids as components of sporopollenin (Guilford *et al.*, 1988; Scott, 1994). In particular, high activity of chalcone synthase (CHS) and phenylalanine ammonia lyase (PAL) have been correlated with pollen fertility (Kishitani *et al.*, 1993; Taylor and Hepler, 1997; Atanassov *et al.*, 1998). Downregulation of PAL and CHS enzymes using antisense RNA, results in complete male sterility due to abnormal pollen development (van der Meer *et al.*, 1992; Matsuda *et al.*, 1996). An anther-specific transcriptional activator of *PAL (NtMYBAS)* was identified in tobacco and likely is a positive regulator of *PAL* expression and phenylpropanoid synthesis in sporophytic, but not in gametophytic tissues of the anther (Yang *et al.*, 2001). Silencing of the tapetum-specific zinc finger gene *TAZ1 (tapetum development zinc finger protein 1)* causes premature degeneration of the tapetum and pollen abortion in *Petunia*, with reduced flavonol accumulation, defects in pollen wall formation and poor germination (Kapoor *et al.*, 2002). Similarly, the *F3H* gene, encoding a flavone-3-hydroxylase gene in *Zea mays*, was shown to be correlated with flavonol accumulation in anthers (Deboo *et al.*, 1995). The expression of phenylpropanoid genes extends behind the completion of exine formation, indicating that they are also involved in other pollen structures (Shen and Hsu, 1992). However, flavonols may not be essential components of pollen grains in all species, since the flavonol-deficient *Arabidopsis tt4 (transparent testa4)* mutant, which disrupts the chalcone synthase gene known to catalyze the first step of flavonoid biosynthesis, shows normal pollen development (Burbulis *et al.*, 1996; Ylstra *et al.*, 1996). An alternative explanation to account for

the fertility of *Arabidopsis* is the possibility that phenylpropanoids other than quercetin and kaempferol can compensate for a lack of flavonoids (Mitchell *et al.*, 1970; Ylstra *et al.*, 1992; Li *et al.*, 1993; Ylstra *et al.*, 1996).

Several mutants have been isolated in recent years, all affecting the exine. The *Arabidopsis* mutant, *dex1n (defective in exine 1)* is blocked in the normal invagination of the plasma membrane, which disrupts the proper deposition of sporopollenin, providing direct evidence for a critical role of the plasma membrane in the pollen wall pattern. *DEX1* is predicted to encode a membrane protein that contains several potential calcium-binding domains, which could act as nucleation sites for sporopollenin deposition (Paxson-Sowders *et al.*, 1997; Paxson-Sowders *et al.*, 2001). Two *Arabidopsis* male sterile mutants, *ms9* and *ms12*, also show defects on exine deposition. In both, the tapetum degenerates earlier suggesting that the mutation may just affect general functions on the tapetum (Taylor *et al.*, 1998). *Lap1 (less adherent pollen 1)*, which shows a disturbed exine pattern, also presents defects on pollen adhesion to the stigmatic surface (Zinkl and Preuss, 2000).

In addition to the mechanical protection of the exine, a lipid-rich coat, termed pollen coat, pollenkit or tryphine, fills the spaces between the baculae of the exine surface and provides several important functions, such as attachment to pollinators, pollen-stigma interactions, protection of pollen grains from excessive dehydration, UV-radiation or pathogen attack (Pacini, 1997; Zinkl *et al.*, 1999; Dickinson *et al.*, 2000).

Unlike exine, the pollen coat is easily extractable by organic solvents such as cyclohexane (Doughty *et al.*, 1993). *Brassica* and *Arabidopsis* pollen coat has been intensively characterized (Doughty *et al.*, 1993; Preuss *et al.*, 1993; Ross and Murphy, 1996; Rüter *et al.*, 1997c; Murphy and Ross, 1998; Mayfield and Preuss, 2000; Mayfield *et al.*, 2001; Fiebig *et al.*, 2004), showing the presence of non-polar esters of medium and long-chain fatty acids, very long fatty acids (VLFA), small proteins and glycoproteins (Preuss *et al.*, 1993; Piffanelli *et al.*, 1997). Non-polar esters such as triterpene and sterol esters are thought to maintain the fluidity of the pollen coat in order to enclose and hold together proteins and other substances embedded in the pollen coat (Caffrey *et al.*, 1987; Piffanelli *et al.*, 1997). The long chain lipids of tryphine are known to be important during the initial contact with the stigma surface either by acting as signaling molecules themselves or by stabilizing/solubilizing other signal components on the pollen coat or cell walls (Wolters-Arts *et al.*, 1998; Pruitt *et al.*, 2000). Several *eceriferum (cer)* mutants in *Arabidopsis*, which eliminate very long chain lipids from the cuticle surface and, in some cases, from the pollen coat, are conditional male sterile. Pollen grains fail to hydrate when placed on the stigma, but are able to hydrate and grow a pollen tube *in vitro* (Preuss *et al.*, 1993; Aarts *et al.*, 1995; Hulskamp *et al.*, 1995b; Jenks *et al.*, 1995; Fiebig *et al.*, 2000).

Although in species with wind-dispersed pollen the coat is highly reduced or absent, maize was used to characterize the two predominant proteins of the pollen coat. One is an endoxylanase and the other a β -glucanase. They are expressed in the tapetum and are thought to help pollen tube penetration by hydrolyzing the stigma cell wall (Bih *et al.*, 1999; Suen *et al.*, 2003). Some stigmas also present oleosins, which are thought to stabilize the lipids in the pollen coat during hydration and re-hydration and assist on

adhesion of the pollen coat to stigma and subsequent hydration. In *Brassica* several glycine-rich oleosins were shown to be specifically expressed in the tapetum and their expression seems to be regulated by the water content in anthers (Ruiter *et al.*, 1997a). Finally, mutant pollen from *Arabidopsis* deficient in one of the pollen surface glycine-rich proteins (GRP17) could undergo germination and fertilization although hydration on stigma was delayed (Mayfield and Preuss, 2000).

Lipases are also a major constituent of the *Arabidopsis* pollen coat and they were shown to alter lipid composition *in vitro*, making them putative mediators of pollen coat behavior (Mayfield *et al.*, 2001). Similarly, non-specific lipid transfer proteins (LTPs), which are small, soluble, basic proteins from plants, are known to stimulate phospholipid transfer between membranes *in vitro*. They can bind to fatty acids and acyl-CoA esters and are thought to secrete or make the deposition of lipophilic substances in cell walls (Aronel *et al.*, 2000). Several LTP proteins are expressed in the tapetum raising the possibility that these proteins may participate in the transfer of fatty acids and other lipid precursors from the tapetum to the microspores during pollen wall deposition.

The *E2* gene encoding a LTP in *Brassica napus* is exclusively expressed in tapetal cells (Foster *et al.*, 1992). In *Arabidopsis* the LTP1 protein is highly expressed in the cell walls of stigma and pollen grains. The *LTP1* promoter region contains sequences homologous to putative regulatory elements of genes in the phenylpropanoid biosynthetic pathway, suggesting that the expression of the *LTP1* gene may be regulated by the same or similar mechanisms as genes in the phenylpropanoid pathway (Thoma *et al.*, 1994). In *Zea mays* the *MZm3-3* gene was suggested to be involved in pollen coat formation. It encodes a short alkaline protein of 10.6 kDa expressed in anthers with a conserved pattern of eight cysteine residues common to lipid transfer proteins and some male-flower-specific proteins (Lauga *et al.*, 2000). Similarly, the *LHM7* gene from *Lilium henryi* L. is anther specific and also contains the conserved pattern of cysteine residues present in non-specific lipid transfer proteins from *Arabidopsis thaliana*, *Antirrhinum majus* and *Lycopersicon esculentum* (Crossley *et al.*, 1995).

Despite the new insights into the genetics and function of pollen coatings, the mechanisms by which these components are formed in the tapetum and translocated to the pollen grain surface and how they interact with the stigmatic surface remain far from clear.

Gene expression on the male gametophyte

The earliest efforts to dissect the molecular basis of pollen development relied on kinetic analysis of transcriptional and translational activity of anther-specific genes corresponding to different stages of pollen development. The first conclusive evidences that transcription and translation of the haploid genome during pollen development occur independently of the diploid tissues, were obtained from isoenzymatic profiles (Linskens, 1966; Weeden and Gottlieb, 1979; Stinson and Mascarenhas, 1985). In addition, detection of different classes of RNA after incorporation of labelled RNA precursors demonstrated an intense RNA synthesis associated with PMI, which resulted in an accumulation of RNAs upon anther dehiscence. In lily and *Tradescantia* the synthesis of ribosomal and transfer RNA was

shown to occur prior to microspore mitosis, followed by a sharp decrease on synthetic activity leading to inactivation of transcriptional activity during the final stages of pollen development (Steffensen, 1966; Mascarenhas and Bell, 1970; Peddada and Mascarenhas, 1975). Despite these observations, protein synthesis is essential during pollen germination and pollen tube growth, suggesting that it is sustained by the presynthesized RNAs (reviewed by Mascarenhas, 1975). Interestingly, it has been shown in *Tradescantia* that a substantial amount of small molecular weight RNAs, which are not tRNAs, are synthesized upon pollen germination and tube growth (Mascarenhas and Goralnick, 1971).

The knowledge of pollen specific gene expression increased considerably through the use of differential and subtractive hybridization of cDNA libraries from isolated mature pollen grains of several species. Quantitative estimates of the amount of gene expression in the gametophyte generation and of the extent of gametophyte-sporophytic overlap were described (Stinson *et al.*, 1987; Hanson *et al.*, 1989; Brown and Crouch, 1990; Theerakulpisut *et al.*, 1991; Weterings *et al.*, 1992a). In maize, a major switch of gene expression after microspore mitosis was demonstrated by differences on the mRNA populations isolated from different developmental stages (Bedinger and Edgerton, 1990). Mandaron *et al.* (1990) used *in vivo* labelling and two-dimensional gel electrophoresis to show that protein synthesis was extremely active from tetrad stage to the vacuolated stage of pollen development, stopped for a short period during starch accumulation and rapidly increased just before anther dehiscence, indicating that presumably these proteins were required for pollen germination and tube growth.

All these results established the foundation for the generally accepted concept that pollen development is transcriptionally modulated. RNA populations and protein profiles of several species showed that specific mRNAs could be associated with pre- and post-mitotic stages, sustaining the view proposed by Mascarenhas (1990), that two classes of genes were expressed during pollen development. Transcripts from the "early" genes, which are detected soon after meiosis and are reduced on mature pollen grains and transcripts of "late" genes, which are first detected soon after PMI and continue to accumulate as pollen matures, create a stable pool of mRNA thought to be essential for germination and early pollen tube growth. Thus, at the time of anthesis, all the proteins that are required for germination and early tube growth are either already in the pollen grain, or if new, the messenger RNAs for their synthesis already exist in the ungerminated pollen grain. The genetic program of the latter part of pollen maturation prior to anthesis should then be the same as the one present during germination and tube growth.

Because most of the research has been done on pollen-specific genes expressed after pollen mitosis, few transcripts of the so-called "early" genes were isolated. The undifferentiated state of the microspore before mitosis may constitute a reason for the difficulty in isolating microspore-specific genes. A differential screening of cDNA libraries covering the stages of anther development from pre-meiotic microsporocytes to tri-nucleate pollen grains led to the isolation of several microspore-specific clones from *Brassica napus* (Scott *et al.*, 1991b). The *Bp4* gene in *Brassica napus* is expressed from the early unicellular microspore stage until the tricellular stage (Albani *et al.*, 1990). In

situ hybridization of the *NTM19* transcript of *Nicotiana tabacum* was only detected in the unicellular microspore (Oldenhof *et al.*, 1996). However, the expression of the *GUS* (β -glucuronidase) reporter gene under the control of *Bp4* and *NTM19* promoters in a heterologous system (*Nicotiana tabacum*) demonstrated that the *Bp4* promoter becomes active after the first pollen mitosis, but not in the microspores. However, the *NTM19* promoter turned out to be microspore-specific and to direct very high levels of *GUS* expression in unicellular microspores. These results showed the *NTM19* promoter as an excellent tool to direct high levels of transgene expression exclusively to the microspores (Custers *et al.*, 1997).

A good example of the "late" pollen genes, are the *LAT* (*Late Anther Tomato*) genes isolated from *Lycopersicon esculentum*, which have been intensively characterized in the last years. The *LAT52* and *LAT51* genes encode for cysteine-rich proteins and the *LAT56* and *LAT59* show sequence similarity to pectate lyase proteins, all presumably involved on late stages of pollen development (Twell *et al.*, 1989; Ursin *et al.*, 1989). Conclusive evidence of a vegetative cell-specific regulation was obtained for *LAT* genes by linking the gene promoter sequences to a *GUS* reporter gene (Twell *et al.*, 1991; Twell, 1992; Eyal *et al.*, 1995; Bate *et al.*, 1996). The *LAT52* promoter was able to confer a dramatically increased expression of heterologous transcripts in a pollen-specific and strictly developmentally regulated manner during the final stages of pollen maturation and thus turned out to be a suitable promoter to direct transgene expression in mature pollen grains of dicots. Since then it has been widely used in numerous studies (Muschiatti *et al.*, 1994; Liu *et al.*, 1999; Gerola *et al.*, 2000; Cheung, 2001; Faure *et al.*, 2002; Cheung *et al.*, 2003). Several other late genes showing a similar pattern of expression were isolated from several species. In some of them the promoter regions share considerable homology with regulatory elements of *LAT* promoters (Hanson *et al.*, 1989; Guerrero *et al.*, 1990; Albani *et al.*, 1991; Hamilton *et al.*, 1992; Lombardero *et al.*, 1994; Weterings *et al.*, 1995; Hamilton *et al.*, 1998).

Due to their size, sperm RNA is likely to be diluted on the total pollen RNA and thus poorly represented in EST databases. This called for sperm or generative cell cDNA libraries as a way of identifying specific transcripts. The approach was first taken on species where the relative large size of the pollen grain would facilitate generative cell isolation. The *LGC1* gene was isolated from a cDNA library of lily generative cells and was shown to be expressed exclusively at the surface, suggesting a possible role in sperm-egg interactions (Xu *et al.*, 1999). Moreover, analyses of the *LGC1* promoter have shown a regulatory sequence required to direct gametic cell-specific expression (Singh *et al.*, 2003). Recently thousands of cDNA clones isolated from a sperm-cell cDNA library of *Zea mays* revealed the presence of a large diversity of mRNAs (Engel *et al.*, 2003). Most of the sequences are predicted to encode secreted or plasma membrane localized proteins and a large number of transcripts of unknown function, representing potential good candidates to mediate gamete interactions. However, some sperm transcripts were found to be expressed in late microspores and hence could be present on the vegetative cell as well. The authors suggest that certain transcripts may be transcribed early in pollen development and later directed into sperm cells.

A transcriptomic definition of pollen

Despite all the approaches described on the previous paragraph, the genetic basis of pollen development and pollen germination and tube growth was restricted to a total of about 150 pollen-expressed genes in about 28 species (Twell, 2002). The development of technologies allowing true large-scale gene expression profiling has paved the way for the comparison of the transcriptome of the male gametophyte with the transcriptional profiles of sporophytic tissues, which revealed the unique features of the transcriptome of mature pollen grains. Five recent studies used *Arabidopsis thaliana* with two different technologies. While Lee and Lee (2003) used Serial Analysis of Gene Expression (SAGE) technology, two other groups employed GeneChip technology (Becker *et al.*, 2003; Honys and Twell, 2003; Honys and Twell, 2004; Pina *et al.*, 2005). The five studies come to similar general conclusions, but differences arise due to the technology used and the different experimental approaches.

In a first approach Becker *et al.* (2003) and Honys and Twell (2003) used Affymetrix AG GeneChip arrays, which represented approximately 8,200 genes and thus almost one third of the *Arabidopsis* genome (reviewed in da Costa-Nunes, 2003). Both groups compared the transcriptional profile of mature pollen grains to those of sporophytic tissues, but there were significant differences in the number of pollen-expressed genes (1,584 versus 992) and of genes predicted to be selectively expressed in pollen (10% versus 40%). The discrepancy for pollen-selectively expressed genes most probably derived from the differences of sporophytic tissues chosen for comparison. While we compared the transcriptional profile of pollen to those of seedlings, leaves, roots and siliques, Honys and Twell (2003) used developmental stages instead. The latter approach could cause a dilution effect of individual tissues expression patterns and therefore increase the proportion of putatively pollen-selective genes. Discrepancies in pollen-expressed genes and relative expression values between the two studies might also derive from the fact that we developed a protocol for fluorescence activated cell sorting (FACS) of *Arabidopsis* pollen grains to assure that only highly purified, viable pollen grains were used for the extraction of total RNA and in addition a specialized normalization protocol for the GeneChip raw data sets that accounts for the relatively small number of genes expressed in pollen, when compared to the vegetative tissues (Becker *et al.*, 2003). We believe that cell sorting is important, not only to avoid that RNA from non-pollen cells is included in the pollen sample, but also because autolysis, observed in mature *Arabidopsis* pollen grains (Yamamoto *et al.*, 2003), bears the risk of including non-viable pollen with altered RNA levels and ongoing RNA degradation in the sample. The fact that different *Arabidopsis* ecotypes were used in the two studies (*Columbia* and *Landsberg erecta*) might also explain differences between the results obtained. Phenotypic differences between ecotypes and proteome analysis (Chevalier *et al.*, 2004) raise doubts on the value of direct comparisons of transcriptional profiles derived from different *Arabidopsis* ecotypes. This notion is supported by recent studies using ATH1 arrays to compare the transcriptomes of vegetative shoot apices from *Col* and *Ler*, which indicates many genuine expression differences between these two ecotypes (Schmid *et al.*, 2003).

Besides the differences in the two studies by Becker *et al.*

(2003) and Honys and Twell (2003), a general picture of the unique characteristics of the pollen transcriptome emerged. This included a reduced complexity when compared with the sporophyte and a relative over-representation of mRNAs in pollen encoding proteins involved in signalling, cell wall metabolism and cytoskeleton dynamics, contrasted by an under-representation of mRNAs related to energy pathways and translation. These studies comprised an important first step to a better understanding of pollen tube growth and morphogenesis by providing hundreds of new transcripts of potential importance.

The introduction of the ATH1 GeneChip array (Redman *et al.*, 2004) allowed the enlargement of the studies to a nearly full-genome scale, covering more than 80% of the *Arabidopsis* genome. Pina *et al.* (2005) compared the transcriptional profile of cell-sorted, mature pollen grains with those of seedlings, flowers, leaves and siliques (*A. thaliana* ecotype *Columbia*), while Honys and Twell (2004) have analyzed male gametophyte development in *A. thaliana* ecotype *Landsberg erecta* from uninucleate microspores (UNM) over bicellular (BCP) and tricellular (TCP) pollen to mature non-sorted pollen grains (MPG) in comparison with publicly available sporophytic datasets. In another approach, Lee and Lee (2003) used SAGE to profile the transcriptome of mature pollen (*A. thaliana* ecotype *Columbia*) under normal and chilling conditions and in comparison to leaves. The number of identified genes expressed in mature pollen grains ranged from 4,211 in the SAGE study over 6,587 genes in cell-sorted pollen grains to 7,235 genes in non-sorted pollen grains. The significantly lower number obtained in the SAGE study surprises, since this technology does not require prior knowledge of the transcripts and can thus discover previously unknown transcripts, which would not be detected on the ATH1 GeneChip. However, a possible explanation for the lower number is that only the limited amount of 21,237 tags were obtained and sequenced, which in the authors' estimate should have allowed them to detect transcripts present at a minimum of 5 copies per cell (1/20,000 transcripts). In addition, the use of relatively short 10-bp sequence tags in this study complicates the unique assignment to specific genes, particularly when members of a gene family have a high degree of similarity (tag-to-gene ambiguity, reviewed in Meyers *et al.*, 2004). The detection sensitivity of Affymetrix GeneChips is approximately 1/100,000 transcripts and with 11 probe sets representing each transcript most of the even highly homologous sequences can still be distinguished. Thus the number of 6587 different transcripts being expressed in *Arabidopsis* pollen based on our study should be more reliable (Pina *et al.*, 2005). An extrapolation to the 28,000 genes encoded in the *Arabidopsis* genome, would lead to an estimate of about 8,200 transcripts stored in mature pollen grains.

The study by Honys and Twell (2004) reveals the remarkable decline in diversity of mRNA transcripts during the development of the male gametophyte, which is most prominent during the transition from bicellular to tricellular pollen. According to their analysis the transcript diversity slightly increases from 11,565 in microspores to 11,909 different transcripts in tricellular pollen and then drops to 8,788 and 7,235 different transcripts expressed in tricellular and mature pollen, respectively. However, these numbers should be used in perspective for two reasons. Firstly the purity of several stages of the isolated microspores was sub-optimal, ranging from 95% for UNM over 77% for BCP to 88% for

TCP. With the sensitivity of the ATH1 GeneChip estimated at 1 transcript per cell (Redman *et al.*, 2004), transcripts with medium and high expression levels in the impurities will be detected as present. In other words, genes expressed at medium or high levels in bicellular and not in tricellular pollen will have been detected on the TCP GeneChips, because of the 12% of bicellular pollen contained in the tricellular pollen sample. Thus the decline in diversity of transcripts from bicellular to tricellular pollen is most probably underestimated. Secondly, during the data analysis the empirical MAS 4 detection algorithm was used to determine present and absent calls for all GeneChip raw data sets (Honys and Twell, 2004), which yields more false positive calls in comparison to the statistical MAS 5 detection algorithm (Liu *et al.*, 2002). We re-analyzed the data sets for the male gametophyte (J.D. Becker, unpublished) using DNA-Chip Analyzer 1.3, the MAS 5 algorithm for detection calls and the relatively stringent criterion used by Honys and Twell (2004) of calling only genes as Present (expressed), when they were called Present in both replicates; this stringency seems justified with regard to the impurities mentioned and the relatively low correlation of the replicates, especially for BCP and MPG. Based on 22,750 genes being represented on the ATH1 GeneChip, our re-analysis identified 9390 different transcripts to be expressed in UNM, 9,602 in BCP, 6,788 in TCP and 5,075 in MPG. 11,405 genes were expressed in at least one stage of male gametophyte development. Since Honys and Twell (2004) identified 13,977 male-gametophyte expressed genes, the difference of 2568 genes could be seen as potential false positives. The comparison of the gametophytic data set (*Landsberg erecta*) with sporophytic ATH1 data sets derived from the ecotypes *Columbia*, *Wassilewskija* and *C24* adds further complications to this study. Yet, this pioneer study provides a good general overview of the transcriptome changes occurring during the development of the male gametophyte.

The specialization of the male gametophyte transcriptome after pollen mitosis II (PMII) can also be seen in a reduction of the number of diverse transcripts comprising the high abundance class, while at the same time genes related to cytoskeleton, cell-wall and signaling become over-represented in this class. The transition from early to late developmental programs was further analyzed by using cluster analysis, a powerful tool to detect sets of co-regulated genes in time-course data sets. As expected, major clusters containing genes repressed after PMII were identified. Those clusters containing genes up-regulated after BCP/TCP stage should contain genes of importance during pollen germination, tube growth and fertilization. Focusing on genes with potential regulatory roles in the male gametophyte, Honys and Twell (2004) showed that for most of the core translation factors expression is restricted to the early stages. Surprisingly, against this trend, 6 of 7 transcripts encoding poly(A)-binding proteins are expressed in mature pollen grains, some specifically. The analysis of 608 transcription factors (412 after our re-analysis), showed clusters containing transcripts with early, constitutive and late expression throughout development of the male gametophyte. Those transcription factors showing high expression levels in mature pollen grains are of special interest, because studies in several plant species have indicated that the bulk of mRNA needed for pollen germination and early tube growth is stored in mature pollen grains (Mascarenhas, 1989; Guyon *et al.*, 2000;

Wang et al., 2004). Honys and Twell (2004) confirm this strict dependence on translation, but not transcription, for *Arabidopsis thaliana* pollen by using translational and transcriptional inhibitors in pollen cultures. However, their criterion to score pollen-tube growth was that tubes had to be longer than two pollen grain diameters. Thus their study does not allow conclusions on what effect a block of transcription will have on pollen tube growth in later stages. It is likely that at least some of the transcription factors up-regulated in mature pollen grains are involved in *de novo* transcription during pollen tube growth. This idea is supported by differential screens for cDNAs up-regulated or induced after pollen germination in *Petunia* (Guyon et al., 2000), *de novo* synthesis of mRNAs depleted after cold-storage in lily pollen (Wang et al., 2004) and by an initial study comparing the transcriptional profiles of *Arabidopsis* pollen grains and *in vitro* grown pollen tubes using ATH1 GeneChips (Miyazaki et al., 2004).

We compared the transcriptomes of pollen and vegetative tissues of the same ecotype (A.t. Col-0 seedlings, leaves, siliques and flowers) using graphical "Snail-view" representations and Principal Component Analysis to underline the distinctiveness of the pollen transcriptome (Pina et al., 2005). Based on these comparisons we identified 11% of the 6587 genes expressed in mature pollen grains as pollen selectively expressed. Mature pollen grains have not only greater proportions of selectively expressed genes, but also more enriched genes (26%) than any vegetative tissue, confirming previous conclusions from other groups (Lee and Lee, 2003; Honys and Twell, 2004).

Can this relative specialization of the pollen transcriptome be linked to the biological functions of pollen, namely germination, pollen tube growth and fertilization of the female gamete? The three studies come to similar conclusions: using Gene Ontology (GO) terms and adding an analysis of statistical significance we looked at the frequencies of representation of GO categories in the different tissues (Pina et al., 2005). Transcripts encoding proteins in the classes signalling (reviewed in Feijó et al., 2004), vesicle trafficking, cytoskeleton and membrane transport were proportionally over-represented, a tendency also observed in male-gametophyte specific genes during microsporogenesis (Honys and Twell, 2004). These classes become even more prominent when analyzing the functions of pollen-enriched genes (Lee and Lee, 2003; Pina et al., 2005). On the contrary, the functional classes transcription, protein synthesis (e.g. ribosomes) and general and oxidative metabolism are under-represented in mature pollen grains. Although the protein synthesis class is under-represented, translation initiation factors are enriched in pollen (Pina et al., 2005), supporting the notion that an mRNA pool ready for rapid translation upon re-hydration is stored in mature pollen.

A detailed view on gene families and pathways in *Arabidopsis* pollen has become possible due to the extensive representation of transcripts of gene families and pathways on the ATH1 array. As an example expression data for mature pollen grains compared with an average of vegetative tissues are overlaid on a map of general metabolism in *Arabidopsis* (Fig. 3A). In accordance with results of the GO analyses, a high proportion of genes involved in cell wall and lipid metabolism are up-regulated and the opposite was found for transcripts involved in energy metabolism. While the absence or down-regulation for transcripts of light reactions and photorespiration is anticipated, the same tendency

is found in processes like glycolysis and mitochondrial electron transport. Considering the fast growth rate of pollen tubes, how the energy needed is provided becomes an imminent question.

A detailed analysis of transcription factor families hints at an important role for non-classical MADS-box genes, showing an over-representation of pollen-expressed and enriched genes (Fig. 3B). Interestingly, some of these non-classical type I and MIKC* genes have also been detected as specifically expressed during reproductive development in a recent study using ATH1 GeneChips to profile three stages of *Arabidopsis* flower and fruit development (Hennig et al., 2004). As the authors note, their set of identified floral-specific genes is dominated by genes expressed in pollen. The exact functions of the non-classical MADS-box genes are not understood and MADS-box genes are not the only class of transcription factors with enriched transcripts in pollen. We combined and re-analyzed data sets on microgametogenesis with our comparative analysis with the restriction of different ecotypes being used in these studies. As a result, we identified transcription factors up-regulated during the development of the male gametophyte that show at the same time higher expression levels in mature pollen, when compared with vegetative tissues (Fig. 3B and Table 1). These transcription factors might be the ones controlling *de novo* transcription in pollen germination and thus should constitute possible primary targets for reverse-genetic approaches.

Pollen might not only serve as a model to study cell growth and morphogenesis, but also, by omission, of cell cycle control. The vegetative nucleus of pollen is thought to be arrested in G1 of the cell cycle. Again, we have used a combination of pollen expression data with a recent schematic overview of the mechanistic regulation of the G1-S and the G2-M transition in plants (Pina et al., 2005), which indicates that the block is achieved by a combination of absence of transcripts encoding essential proteins for the G1-S transition (CycD, E2F-DP) and an up-regulation of potential repressors (DEL3) and of potential factors of increased cell-cycle duration (CKS2). Surprisingly, pollen seems to feature most of the transcripts needed for the G2-M transition, though probably kept in their inactive state through a relative down regulation of the CDC25 phosphatase. In support of these results, Hennig et al. (2004) have come to the conclusion that S-Phase genes were under-represented and G2/M-Phase genes enriched in the set of genes they identified as specifically expressed during reproduction. In the light of G2 karyogamy in *Arabidopsis* this might indicate a key role for pollen derived transcripts or proteins during the first mitosis after fertilization, although it is unclear how the transcripts or proteins would get to the egg cell unless they are somehow accumulated in the sperm cells.

A surprising finding was the apparent inactivation of small RNA pathways in mature pollen (Pina et al., 2005). The analysis of the expression of genes involved in small RNA pathways in *Arabidopsis* revealed that all of the 15 transcripts analyzed, including those encoding Argonaute 1,2,4 and 7, Dicer-like 1-3 and RNA-dependent RNA polymerase 1,2 and 6, were called Absent in pollen, while the majority of them were expressed in the vegetative tissues. In addition, a specific down regulation of these transcripts occurs during microgametogenesis. The apparent absence of small RNA pathways in *Arabidopsis* mature pollen would affect all small RNA pathways known in plants (Baulcombe, 2004), including the defence against viruses through siRNAs, the regulation of

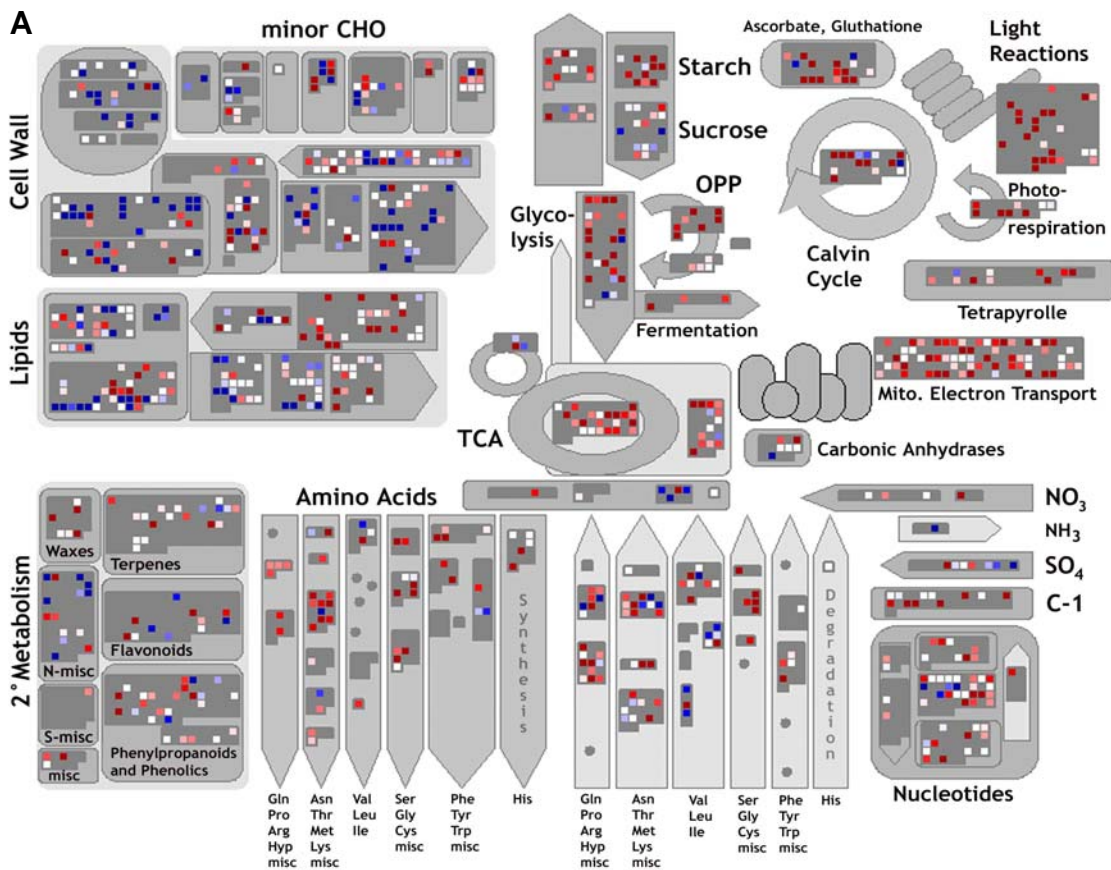
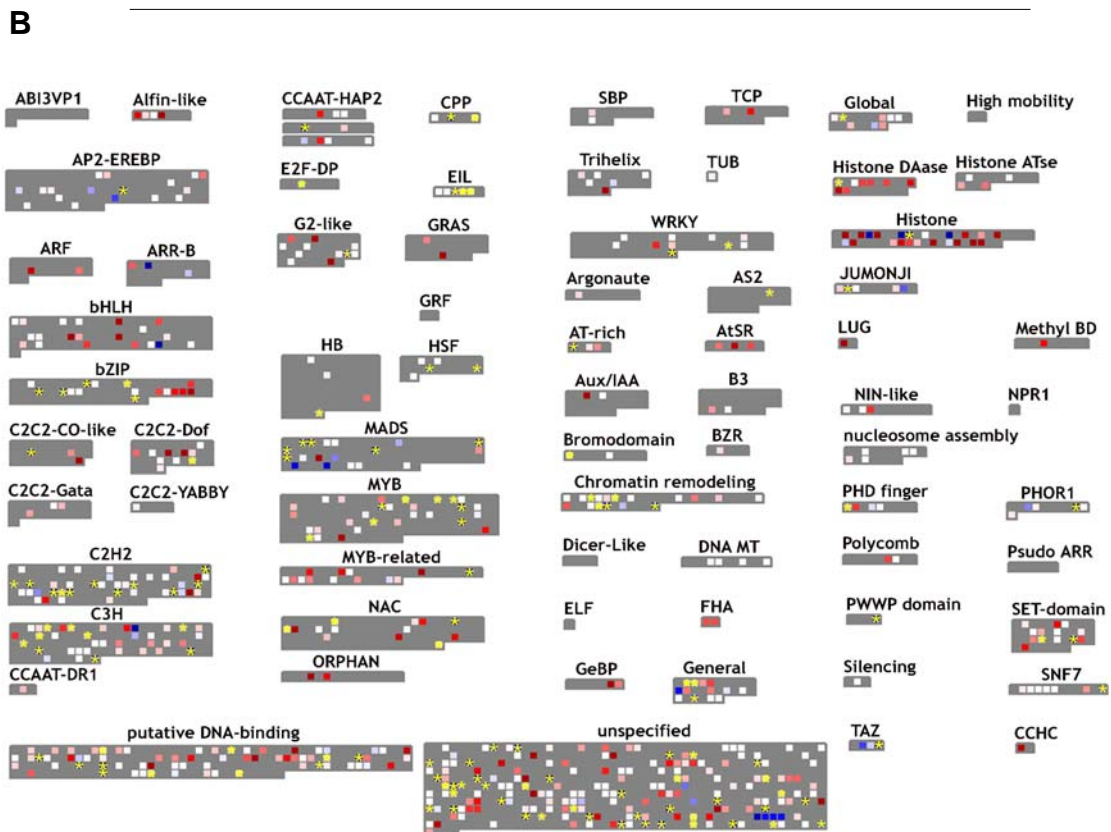


Fig 3. Expression patterns for general metabolism and transcription in the pollen transcriptome. (A) Gene expression data in pollen relative to the vegetative tissues leaves, seedlings and siliques (genomic data set derived from the study of Pina et al., (2005) are depicted in an overlay on a map of general metabolism in Arabidopsis using the MAPMAN tool (Thimm et al., 2004). Genes are symbolized by color-encoded squares (red, down-regulation; blue, up-regulation; white, Present call in pollen, but no change; grey, Absent call in pollen).



(B) Gene expression data in pollen relative to the vegetative tissues on a map of genes involved in transcription in Arabidopsis. Genes are symbolized by color-encoded squares (red, down-regulation; blue, up-regulation; white, Present call in pollen, but no change; grey, Absent call in pollen). In addition those genes showing an up-regulation in this comparison and during Arabidopsis microgametogenesis (mature pollen versus uninucleate microspores; re-analyzed data set of (Honys and Twell, 2004) are marked with yellow asterisks and their detailed expression data are listed in table 1 (except for the groups "putative DNA-binding" and "unspecified").

gene expression through miRNAs and the condensation of chromatin into heterochromatin. This absence of small RNA pathways in mature pollen would provide new possible explanations for genetic and epigenetic phenomena found in the male gametophyte and during/after fertilization.

Though not on a genomic scale, smaller studies in other species, e.g. like those on *Petunia* (Guyon *et al.*, 2000; Cnudde *et al.*, 2003) and lily (Wang *et al.*, 2004), are complementing the *Arabidopsis* data sets and reveal new aspects of pollen genetics and physiology. Transcriptomic studies are providing a quantum leap in information available for the male gametophyte.

The female reproductive organ- pistil and ovule development

Thorough descriptions of the molecular and genetic basis of the female organ development *Arabidopsis* can be found in recent reviews (Gasser *et al.*, 1998; Skinner *et al.*, 2004).

Unlike in other groups of plants, ovules in Angiosperms are enclosed within a carpel and during seed development they produce an embryo-nourishing endosperm (Friedman, 2001b). The evolution of a closed carpel (syncarpy) is believed to be one of the major evolutionary changes, which led to the reproductive success of angiosperms. Carpels are proposed to have evolved from ancestral foliar organs or bract-like structures (Bowman *et al.*, 1999). Carpels may be associated or not in the pistil, or gynoecium, which is the female reproductive unit of the angiosperm flower. Pistils differ widely in form between species (Endress and Igersheim, 1999). Despite the differences in the overall morphology, a set of common structures, characterized by cell types with unique properties, can be distinguished. In *Arabidopsis*, at maturity, most gynoecia are composed of (1) a basal ovary, usually with fused carpels, which encloses the ovules bearing the female gametophyte, (2) a short solid style, which places the stigmatic surface in close contact with the anthers and (3) an apical stigma composed of numerous elongated epidermal cells, where pollen grains adhere and germinate (Sessions and Zambryski, 1995; Bowman *et al.*, 1999). The style varies in length and appearance in association with specific pollination strategies (Barrett, 2002). The transmitting tract is a special layer of cells, which originates in the stigma and spans the entire organ, secreting a mucilaginous extracellular matrix. After fertilization a prominent elongation of the ovary precedes the dispersal of the mature seeds (Ferrandiz *et al.*, 1999).

In *Arabidopsis* pistil development starts with a ring-shaped primordium by stage 6 of flower development. The gynoecium continues to elongate and develops as an open cylinder with a medial ridge, where the placental and septum will form (Fig. 4 A,B). At stage 9, rows of ovules arise from the placental tissue along the margins of the site of carpel fusion, the septum. By stage 10 the septum is formed, resulting in a gynoecium with two congenitally fused carpels. At stage 11, the stigmatic papillae differentiate and ovules develop a funiculus and integument. At stage 12 the gynoecium grows, the transmitting tract differentiates and ovules are completed. Stage 13 defines anthesis, the gynoecium comes to maturity and the flower opens (Smyth *et al.*, 1990; Sessions, 1997; Ferrandiz *et al.*, 1999).

Ovules are simple structures, which consist of three elements: at the top, the nucellus harbors the haploid embryo sac or female gametophyte, a central chalaza, which is characterized by the

integuments initiating at its flanks and at the bottom and the funiculus or stalk that connects the ovule to the carpel tissue (Fig. 4G). The ovules are initiated from divisions occurring in the subepidermal tissue of the placenta and cell proliferation gives rise to finger-like structures that will differentiate in an ovule (Fig. 4 C-E) (Jenik and Irish, 2000). The integuments are two cell layers that will give rise to the seed coat. They cover the nucellus, forming the micropyle, a small aperture through which the pollen tube will penetrate (Fig. 4G). The outer integument grows asymmetrically, shaping the typical curvature to the ovule (Fig. 4G).

Identically to other floral organs, ovule identity is regulated by MADS-box genes. In tobacco the two mutants *Mgr3* and *Mgr9* have carpelloid structures instead of ovules and in *Petunia* the co-suppression of the two genes *FLORAL BINDING PROTEIN7* (*FBP7*) and *FBP11* resulted in a similar phenotype (Evans and Malmberg, 1989; Angenent *et al.*, 1995). Furthermore the ectopic expression of *FBP11* led to ectopic ovuloid expression (Colombo *et al.*, 1995). The identification of these and many other flower organ mutants contributed to elucidate much of the molecular and genetic mechanism of floral organ development, but it should be kept in mind that many of these mutants show pleiotropic effects in early stages of flower development and thus the elucidation of their true effects may be partially disguised.

The female gametophyte - the embryo sac

The female gametophyte supports pollen tube growth down to the ovule, directing the sperm cells to the egg cell and central cell, up to seed, embryo and endosperm development (Chaudhury and Berger, 2001; Chaudhury *et al.*, 2001; Higashiyama, 2002; Johnson and Preuss, 2002; Higashiyama *et al.*, 2003). Sporophytic mutations characterized by female sterility, such as megaspore mother cell development and meiosis are developmental processes originated in the surrounding sporophytic tissues (Gasser *et al.*, 1998; Grossniklaus and Schneitz, 1998). On the other hand, systematic screens based on two essential criteria, abnormal seed set and distorted Mendelian segregation, identified a number of gametophytic mutations disrupting distinct steps of embryo sac ontogenesis in *Arabidopsis* and maize, indicating the requirement of haploid expressed genes (Christensen *et al.*, 1997; Drews *et al.*, 1998; Siddiqi *et al.*, 2000; Christensen *et al.*, 2002).

The ovule primordium initiates as an outgrowth that emerges from the carpel placental tissue (Fig. 4 B-F). Megasporogenesis begins with the differentiation of an archeosporial cell in the hypodermal cell layer at the distal end of the nucellus (Fig. 4D). Usually only a single cell differentiates. In the *mac1* mutant of maize several hypodermal cells develop into archeosporial cells, suggesting that this gene may be required to suppress the differentiation in neighboring cells and putatively controlling the switch of the hypodermal cells from vegetative to meiotic (Sheridan *et al.*, 1996). In general, megasporogenesis in *Arabidopsis* involves the differentiation of one of these cells in the megaspore mother cell (MMC) in the nucellus, which undergoes meiotic reduction giving rise to four haploid nuclei (Fig. 4 A, C-E). *SPOROXYTELESS* (*SPL*) blocks the transition from archeosporial cell into a megasporocyte in ovules (Yang *et al.*, 1999). These defects may be related with the lack of support and nutrient supply to the female gametophyte (Reiser *et al.*, 1995; Elliott *et al.*, 1996; Klucher *et al.*, 1996). In agreement with the role of integuments in the control of

TABLE 1
TRANSCRIPTION FACTORS UP-REGULATED IN MATURE POLLEN AND DURING MICROGAMETOGENESIS

Gene Family	FC	Selective	Probe Set	AGI ID	Annotation	Pollen Call	Leaf	Call	Seedling	Call	Silique	Call	
C2H2 zinc finger	77.6		262291_at	A11g70790	C2 domain-containing protein	16713	P	P	136	P	272	P	
C2H2 zinc finger	49.1	X	253153_at	A14g35700	zinc finger family protein	3950	P	A	61	A	41	A	
C2H2 zinc finger	39.6		255232_at	A14g05330	zinc finger and C2 domain protein, putative	5696	P	P	99	P	115	P	
C2H2 zinc finger	33.6		256576_at	A13g28210	zinc finger protein (PMZ) -related	3363	P	P	120	P	60	A	
C2H2 zinc finger	20.1	X	258967_at	A13g10470	zinc finger family protein	1840	P	A	77	A	60	A	
C2H2 zinc finger	18.8		250128_at	A15g16540	zinc finger protein 3	8592	P	P	398	P	303	P	
C2H2 zinc finger	7.8	X	245705_at	A15g04390	zinc finger transcription factor-related protein	573	P	61	A	62	A	56	A
C2H2 zinc finger	5.9		253526_at	A14g31420	zinc finger protein -related	3623	P	P	528	P	760	P	
C2H2 zinc finger	4		252009_at	A13g52800	zinc finger - like protein	1245	P	P	246	P	237	P	
C2H2 zinc finger	3.8		258689_at	A13g07940	zinc finger and C2 domain protein, putative	987	P	A	205	P	263	P	
C2H2 zinc finger	2.3		253187_at	A14g35280	zinc-finger protein -related	175	P	P	62	A	52	A	
C3H zinc finger	7.2		249483_at	A15g38895	C3HC4-type zinc finger protein family	1754	P	P	149	P	354	P	
C3H zinc finger	5.4		253865_at	A14g27470	C3HC4-type zinc finger protein family	1183	P	P	144	P	186	P	
C3H zinc finger	4.4		247125_at	A15g66070	C3HC4-type zinc finger protein family	1354	P	P	260	P	222	P	
C3H zinc finger	2.5		256326_at	A13g02340	C3HC4-type zinc finger protein family	1298	P	P	397	P	631	P	
C3H zinc finger	2.3		260354_at	A11g69330	C3HC4-type zinc finger protein family	552	P	P	215	P	219	P	
C3H zinc finger	2.2		260671_at	A11g19310	C3HC4-type zinc finger protein family	1257	P	P	339	P	690	P	
C3H zinc finger	2.1		260231_at	A11g74410	C3HC4-type zinc finger protein family	636	P	P	186	P	453	P	
C3H zinc finger	2		262656_at	A11g14200	C3HC4-type zinc finger protein family	502	P	P	190	P	198	P	
C3H zinc finger	2		266016_at	A12g18670	C3HC4-type zinc finger protein family	340	P	P	124	A	129	A	
MYB transcription factor	45.7		267087_at	A12g32460	AtMYB101	5080	P	A	61	A	102	P	
MYB transcription factor	18.1	X	253939_at	A14g26930	AtMYB97	1167	P	A	49	A	38	A	
MYB transcription factor	4.3		265700_at	A12g03470	MYB family transcription factor -related	1891	P	P	307	P	392	P	
MYB transcription factor	2.3	X	263006_at	A11g54240	hypothetical protein	221	P	A	77	A	62	A	
MYB transcription factor	2.1	X	248116_at	A15g55020	AtMYB120	173	P	A	70	A	59	A	
MYB transcription factor	1.9		260999_at	A11g26580	expressed protein	1219	P	P	392	P	904	P	
MYB transcription factor	1.9		262724_s_at	A11g43330	PC-MYB2 -related	200	P	A	70	A	75	P	
MADS-box transcription factor	49.5	X	255952_at	A11g22130	AGL104 (MIKC*)	1569	P	A	23	A	21	A	
MADS-box transcription factor	19.8	X	266994_at	A12g34440	AGL29 (Type I MADS-box)	915	P	A	30	A	32	A	
MADS-box transcription factor	13.7	X	262179_at	A11g77980	AGL66 (MIKC*)	1331	P	A	62	A	54	A	
MADS-box transcription factor	12.1		251623_at	A13g57390	AGL18 (Type II MADS-box)	4993	P	P	318	P	796	P	
MADS-box transcription factor	4.5	X	266793_at	A12g03060	AGL30 (MIKC*)	495	P	A	90	A	89	A	
MADS-box transcription factor	4.5	X	266793_at	A12g03060	AGL30 (MIKC*)	495	P	A	90	A	89	A	
Chromatin remodelling complex subunit R	4.3		247056_at	A15g66750	SNF2 domain/helicase domain-containing protein	345	P	A	94	P	86	P	
Chromatin remodelling complex subunit R	4.3		250601_at	A15g07810	SNF2 domain/helicase domain-containing protein	423	P	A	105	P	96	P	

Expression data for genes up-regulated in pollen relative to vegetative tissues (Pina *et al.*, 2005) and likewise during microgametogenesis (Hony and Twell, 2004) are listed (see genes marked with yellow asterisks in Fig. 3B, except the groups "putative DNA-binding" and "unspecified"). The first column gives the gene family and the second gives the lower confidence bound of the fold change (FC; average of the comparisons of pollen to the three vegetative tissues). The third column depicts, if a gene is selectively expressed in pollen (Present detection call only in pollen), followed by the Affymetrix probe set and the TAIR locus (AGI ID) assigned to this probe set in the fourth and the fifth column. The sixth column depicts the gene annotation. In the following column the expression value of the gene (weighted average of duplicates) and its detection call (Call: Present; P or Absent; A) are given for the respective cell type or tissue.

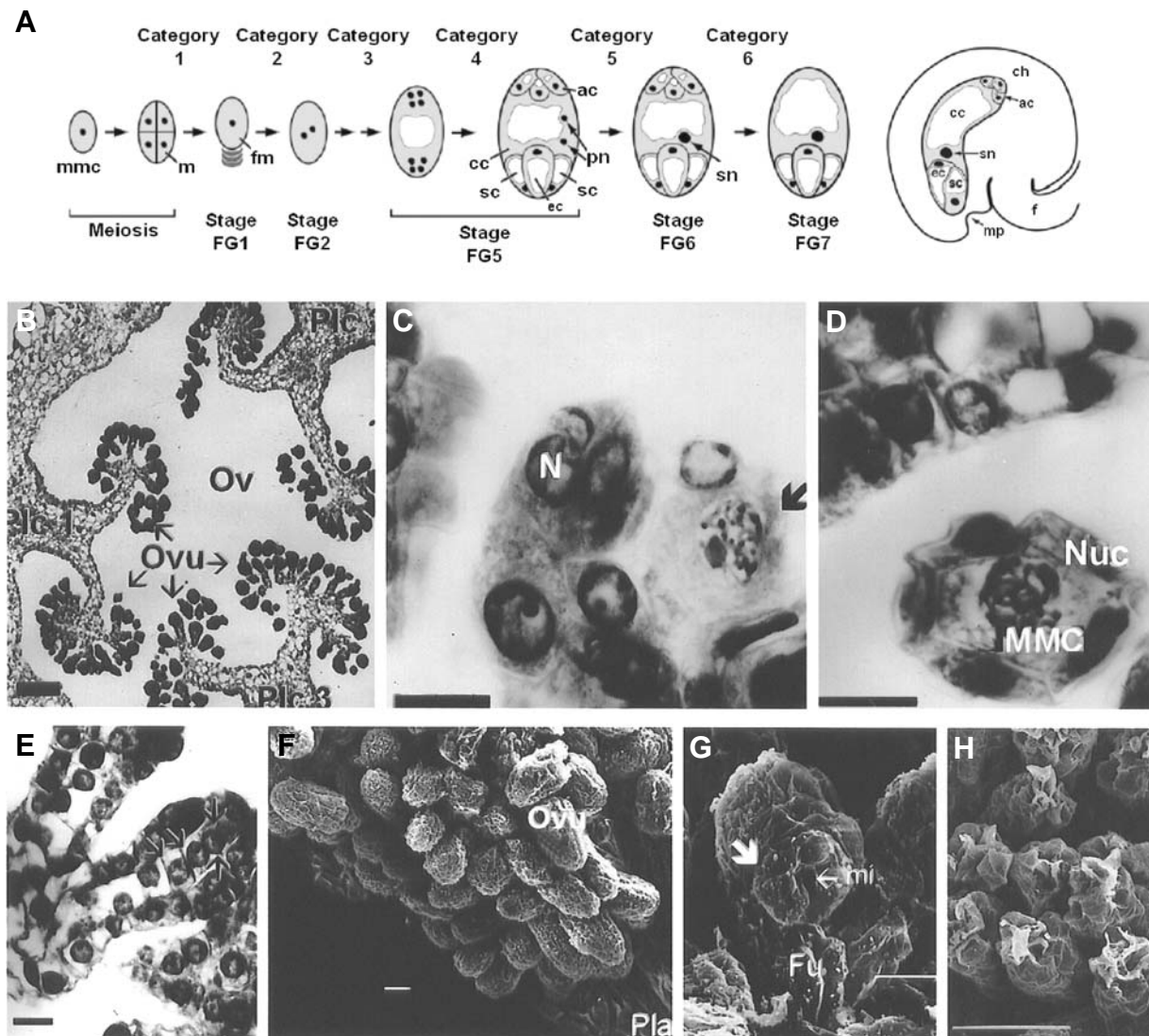


Fig. 4. The development of embryo sac and the ovule. (A) Major events on embryo sac ontogeny. Categories represent checkpoints on embryo sac development based on mutant analysis. mmc megaspore mother cell; fm female megaspore; CC central cell; EC egg cell; SC synergid cell; pn polar nuclei; ac antipodal cell; sn secondary nucleus; ch chalaza; mp mycopyle; f funiculus. Adapted from Drews et al. (1998) and Yadegari and Drews (2004).

(B-E) Embryo sac development in *O. lutea*. The ovary is divided by 3 parietal placentas **(B)**. The apical cell of the nucellus (N or Nuc) divides longitudinally **(C)** and then anticlinally to define a central megaspore mother cell (MMC), that enters meiosis (arrow in **(C,D)**). This cell gives rise to a linear tetrad (arrows in **(E)**), 3 of which will regress and the top one will develop into the mature embryo sac, like depicted in **(A)** (category 3 on). **(F-H)** Ovule development. From a capsule like shape **(F)** ovules (Ov) bend over by differential growth of the integuments to form a pipe-like structure **(G)**. At this stage the integuments also grow to form the pollen tube entry point, the micropyle (mi). At later stages of mature ovules, the micropyle is sometimes covered with secretion **(H)**.

meiotic progression is the *sterile apetala* (*sap*) mutant, where megasporogenesis is arrested during or just after meiosis. Although the *SAP* gene is expressed initially in the nucellar region, it switches to a prominent integument expression later on (Byzova et al., 1999). The *DYAD* gene was reported to be female specific and meiosis is arrested at the dyad stage (Siddiqi et al., 2000). The four haploid cells are surrounded by a very thin cell wall after meiosis, with plasmodesmata connecting the chalazal megaspore to the neighbouring nucellar cells, suggesting that it may provide a positional signal that promotes the degeneration of the other spores to undergo cell death. At the beginning of meiosis in

Arabidopsis the MMC shows cytoplasmic polarity, thought to be important to determine the functional megaspore (Bajon et al., 1999).

Megagametogenesis begins when the surviving megaspore goes through three rounds of mitosis to form a two-nucleated, four-nucleated (Fig. 4E) and subsequently eight-nucleated embryo sac (Fig. 4A; stage FG1-FG4). Different patterns of gametophyte development have been described in several species, differing from each other by variations on cytokinesis during meiosis, on the number of mitotic divisions and on the cellularization pattern. *Arabidopsis* and maize megasporogenesis follows the monosporic

Polygonum type, which is also the most common form in Angiosperms (Maheshwari, 1950; Willemse and van Went, 1984; Huang and Russell, 1992a). At the end of each meiotic stage vacuoles occupy a central location on the cells and microtubules are thought to mediate nuclear migration to position the nuclei at the poles of the coenocytic cell (Webb and Gunning, 1994a). After the third mitotic round non-sister nuclei and the embryo sac begin to cellularize to form the final seven-cell structure. The other two nuclei (the polar nuclei) migrate to a central position. In some species the nuclei fuse to give rise to the secondary endosperm nucleus, in others the polar nuclei just fuse partially prior to fertilization. During cell differentiation the polarity of chalazal-micropyle axis established in the beginning of the female gametophyte development will determine that the nuclei positioned at the micropylar end become specified to develop into an egg and synergids cells and the chalazal nuclei give rise to the antipodal cells. Cell walls are absent in the chalazal end of the egg cell and the synergids, allowing the direct contact of plasma membranes (Willemse and van Went, 1984; Webb and Gunning, 1994b). The polarity of the female gametophyte is believed to be defined by the asymmetry of the surrounding ovule layers and thus to be under sporophytic control, but no factors have yet been identified (Christensen *et al.*, 1997).

Most of the gametophytic mutants, which do not show lesions on the sporophytic tissues of the ovule, fall in different classes corresponding to key developmental events during megagametogenesis, such as mitosis, vacuole formation, nuclear fusion, cellularization and cell death (see categories in Fig. 4) (Feldmann *et al.*, 1997; Schneitz *et al.*, 1997; Christensen *et al.*, 1998; Drews *et al.*, 1998).

Conclusions

Albeit covering most of the recent info, namely the fast accumulating genomic and transcriptomic data, this review mostly highlights the incompleteness of our understanding of many of the fundamental mechanisms underlying the complex puzzle of gametogenesis. Both gametophytes show developmental programs that prepare these tissues for their future interactions. The events that lead to gametophyte development are controlled primarily by the sporophytic tissues, but are ultimately reassigned to singular cells with a narrow set of functions to accomplish. When first released from the sporophyte, the male gametophyte is already committed to a role. The female tissues, on the other hand, form a long and complex pathway, branded by different molecular and cellular contexts. The past years have seen significant advances in our understanding of all these processes, revealing some of the key cellular events, but the identity of many of the molecular components and signaling pathways in which they are involved is still mostly unknown. Moreover, the genetic and functional analysis has mostly progressed on a gene-by-gene basis and relying on the relative serendipity of genetic screens, but transcriptomics offers the promise of major advances through a more deterministic approach. Near future will prove these to be sound basis to a more integrated view of the processes of plant gametogenesis.

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