Transcriptomic and Proteomic Insights into Amborella trichopoda Male Gametophyte Functions

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Flowering plants (angiosperms) are characterized by pollen tubes (PTs; male gametophytes) carrying two immobile sperm cells that grow over long distances through the carpel toward the ovules, where double fertilization is executed. It is not understood how these reproductive structures evolved, which genes occur de novo in male gametophytes of angiosperms, and to which extent PT functions are conserved among angiosperms. To contribute to a deeper understanding of the evolution of gametophyte functions, we generated RNA sequencing data from seven reproductive and two vegetative control tissues of the basal angiosperm Amborella trichopoda and complemented these with proteomic data of pollen grains (PGs) and PTs. The eudicot model plant Arabidopsis thaliana served as a reference organism for data analysis, as more than 200 genes have been associated with male gametophyte functions in this species. We describe methods to collect bicolellar A. trichopoda PGs, to induce their germination in vitro, and to monitor PT growth and germ cell division. Transcriptomic and proteomic analyses indicate that A. trichopoda PGs are prepared for germination requiring lipids, energy, but likely also reactive oxygen species, while PTs are especially characterized by catabolic/biosynthetic and transport processes including cell wall biosynthesis and gene regulation. Notably, a number of pollen-specific genes were lacking in Arabidopsis, and the number of genes involved in pollen signaling is significantly reduced in A. trichopoda. In conclusion, we provide insight into male gametophyte functions of the most basal angiosperm and establish a valuable resource for future studies on the evolution of flowering plants.

During the evolution of seed plants and the colonization of dry environments, male gametes (sperm cells) have lost their mobility and thus need a vehicle to be transported toward the major female gamete (egg cell). This is achieved by the development of a male gametophyte (pollen) capable of forming a tip-growing tube. In angiosperms (flowering plants), the pollen tube (PT) must grow over long distances through the maternal tissues of the carpel to reach the female gametophyte (embryo sac) inside an ovule that is sheltered within the ovary. Together with the double fertilization process involving two distinct female gametes (egg and central cell), these innovations represent main characteristics of angiosperms distinguishing them from all other plants, including gymnosperms. PTs carrying two immobile sperm cells over long distances thus not only play a key role during double fertilization, they are also highly interesting to understand the successful evolution of angiosperms, now representing about 90% of all living land plants (Sauquet et al., 2017). Notably, comparative studies of male gametophytes in gymnosperms and angiosperms also revealed remarkable differences in pollen grain (PG) morphology, PT walls, and growth rates, indicating that pollen–pollen played a central role in the rapid diversification of angiosperms (Williams, 2008, 2012).

In this context, analyses of the most basal living angiosperm, Amborella trichopoda, a sister of all other extant angiosperms, is especially interesting to understand the evolution of pollen–pollen function(s) and morphology.
that contributed to the success of flowering plants. *A. trichopoda* diverged very early on (about 150 million years ago) from all the other extant ~300,000 species of angiosperms and is now considered as the most basal living angiosperm (Drew et al., 2014). It therefore has a unique position to shed light on the evolution of sexual organs and gametophyte development and the associated functions in angiosperms. *A. trichopoda* is a dioecious woody shrub endemic to New Caledonia. Once per year, male individuals produce thousands of small flowers (Fig. 1) that generate large numbers of PGs. Detailed morphological and histological studies have been performed to examine its PGs and to follow PT growth in vivo (Hesse, 2001; Williams, 2009). However, nothing is known about the underlying molecular and physiological processes. Although the *A. trichopoda* genome was published a few years ago (Amborella Genome Project, 2013), transcriptomic data are currently very limited, and comparisons with well-studied angiosperm model plants like Arabidopsis (*Arabidopsis thaliana*) are mostly restricted to predicted genes and gene families, respectively. In contrast, a considerable amount of information has been accumulated during the past 30 years to understand male gametophyte development and function in Arabidopsis and to a lesser extent in other angiosperm species including crops like maize (*Zea mays*) and rice (*Oryza sativa*; Zhou and Dresselhaus, 2019). Among other discoveries, the research community has learned that Arabidopsis has lost genetic mechanisms to avoid self-fertilization (Nasrallah, 2019) and that rapid PT germination and proper navigation through the transmitting tract is critical for reproductive success (Johnson et al., 2019; Lopes et al., 2019). Furthermore, fundamental features and underlying molecular mechanisms of PT tip growth were elucidated, and mechanisms and players regulating cell wall rigidity of this fastest growing plant cell have been identified. Most progress has been made in the past years to understand the mechanisms how PTs exit the transmitting tract in Arabidopsis, how they are guided toward and inside the ovule, and how their burst is regulated to release their sperm cell cargo. Interactions of polymorphic peptide ligands with receptor-like kinases have been shown to play key roles during male-female interactions (Qu et al., 2015; Zhong and Qu, 2019). While mechanisms of polar PT tip growth appear conserved in angiosperms, different proteins are used, for example, for PT guidance and burst (Zhou and Dresselhaus, 2019). Studies involving the basal angiosperm *A. trichopoda* therefore have a high potential to reveal the evolution, conservation, and specificity of above-mentioned reproductive processes and their roles during reproductive isolation and thus speciation during angiosperm diversification.

As a first step to obtain a deeper understanding of the evolution of male gametophytes and their functions in angiosperms, we describe here methods to collect and purify mature bicellular PGs of *A. trichopoda* and how they can be germinated and monitored in vitro. PT growth was measured and timing of germ cell division determined. In addition, we generated transcriptomic data from nine different *A. trichopoda* tissues, including PGs as well as bicellular and tricellular PTs and studied transcriptome dynamics. Proteomic profiles were generated from PGs and tricellular PTs and compared with corresponding transcriptomic data. By using Arabidopsis as a reference angiosperm, we searched for putative orthologs of key proteins involved in male gametophyte function. In conclusion, we provide a valuable resource of transcriptomic and proteomic data of *A. trichopoda* for usage of evolutionary comparative studies of plant sexual reproduction. Furthermore, we identified putative conserved signaling pathways, many genes with yet-unknown functions as well as an the expansion of signaling components in Arabidopsis PTs, suggesting evolutionary diversification over the course of angiosperm evolution.

**RESULTS**

**Bicellular *A. trichopoda* Pollen Generates Two Sperm Cells (Tricellular Stage) within 14 h during PT Growth In Vitro**

It is difficult to cultivate *A. trichopoda* successfully outside their natural environment in New Caledonia.
For many years, the Botanical Garden in Bonn (Germany) has optimized growth conditions for dioecious *A. trichopoda* plants (Grosse-Veldmann et al., 2011). We collected PG and ovules during the flowering period (September to early December) from male and female flowers born on male and female plants as well as vegetative plant material from a male plant through four successive years (2015–2018). To study *A. trichopoda* PGs and PTs (representing the male gametophyte), we first established protocols to isolate mature PGs in large quantities and to induce their germination in vitro. Despite the small size of *A. trichopoda* flowers (Fig. 1, A and B), bulk harvesting of mature PGs from anthers (Fig. 1, C and D) was successful using the procedure described in Figure 1E. In brief, this procedure consists of collecting 20 male flowers in a 50-mL tube, adding 25 mL of *Camellia japonica* pollen germination medium (CJ-PGM) and vortexing the tube for 45 s to release PGs into the solution. Afterward, the suspension is first filtered through a 1-mm pore size mesh to remove large debris and then subsequently filtered through strainers of 70-, 30-, and 15-micron pore size to remove small debris and to separate contaminating fungal spores that we found in all PG preparations. Finally, clean PGs with a diameter of about 25 μm (Fig. 1D) are retained on the 15-micron strainer, recovered from the surface, and transferred to a 1.5-mL collection tube for further analysis.

To induce in vitro pollen germination, five different PGMs were tested (Supplemental Table S1) and the germination ratio evaluated 10 h after incubation. The maximum germination rate (83.17%) was observed using liquid CJ-PGM (Fig. 2A; Nakamura et al., 1982). We next determined that *A. trichopoda* pollen tube growth rate (PTGR) on solid CJ-PGM was 100.46 ± 0.5 μm h⁻¹ (Fig. 2B; Supplemental Table S2). Time-lapse imaging revealed that PT growth in *A. trichopoda* is linear in vitro and without recognizable oscillations as indicated in a x/t-projection (kymograph; Fig. 2C).

We then analyzed nuclei content and cell wall material of PGs and PTs. The 4′,6-diamino-phenylindole staining confirmed that *A. trichopoda* generates bicellular PGs including a single generative cell (GC) and a tube cell whose vegetative nucleus contains less compact chromatin compared with the GC (Fig. 3A). The presence and location of pectin in the cell wall of PGs and PTs was explored by staining with ruthenium red. The highest amounts of pectin were detectable at the aperture zone of the PG as well as in the tip of growing PTs but were also present at significant levels at the shank of the whole PT (Fig. 3B). Aniline blue staining showed the presence of callose deposition in the PT wall and the formation of extended callose plugs during progression of PT growth (Fig. 3F; Supplemental Table S2). SYBR GREEN I staining in combination with the membrane counter stain FM 4-64 revealed that on average the length of sperm cells in the PT is 10 μm. Compared with the chromatin of the GC and the vegetative tube cell, the chromatin of sperm cells is highly condensed (Fig. 3G). About 90% germinated PTs are tricellular 14 h after incubation on CJ-PGM. This time point was used to sample PTs at the tricellular stage for further experimentation.

**Transcriptomes of *A. trichopoda* Male Gametophytes at Different Stages Are Highly Similar and Show Low Dynamics**

In order to visualize differences in gene expression patterns before and during PT germination, we investigated the *A. trichopoda* transcriptomes of mature pollen grains (PGs) as well as PTs at the bicellular (PT-Bi)
and tricellular (PT-Tri) stages. After RNA extraction, library preparation, and Illumina sequencing, a total of 46.6 million reads were generated from PGs, 63.88 million reads from PT-Bi, and 58.01 million reads from PT-Tri stages and uniquely mapped to the *A. trichopoda* genome v1 (Amborella Genome Project, 2013), which was predicted to contain 27,313 genes. Additionally, RNA sequencing (RNA-seq) data were generated from *A. trichopoda* ovaries, leaves, male and female flowers, tepals, and roots, resulting in 8.91, 22.55, 80.25, 81.96, 50.17, and 18.60 million mapped reads, respectively. An overview of the RNA-seq data obtained for all these tissues is provided in Table 1. RNA-seq data were validated by RT-qPCR by selecting nine genes with high, medium, and low log$_2$ fold change (log$_2$FC) values among the tissue comparisons (Supplemental Fig. S1). A master table showing RNA-seq gene expression levels of all *A. trichopoda* genes in transcripts per million (TPM) values for each of the three biological replicates in the seven tissues is provided in Supplemental Table S3.

To determine the number of genes expressed in stages PG, PT-Bi, and PT-Tri, we considered a gene to be expressed at TPM values $\geq 1$. A Venn diagram analysis revealed overlaps of 90.0\%, 93.1\%, and 89.4\% within biological replicates from PG, PT-Bi, and PT-Tri samples, respectively (Fig. 4A). With the exception of ovule samples showing only an overlap of 45.3\% between biological replicates and thus only 6,144 expressed genes in all samples, other control samples (leaves, roots, flowers, and tepals) showed an overlap between 84.4\% to 91.9\% and on average $>15,000$ expressed genes (Fig. 4B). We assume that the lower overlap between the three ovary samples that consequently resulted in a lower number of genes present in all samples results from tissue collection at slightly different developmental stages. There were 10,134 genes expressed in all PG samples, while PT-Bi showed 10,384 and PT-Tri 11,002 expressed genes. Additionally, principal component analyses (PCAs) were performed to analyze the similarity between the biological replicates and among the three pollen developmental stages and the control tissues. As shown in Figure 4C, with the exception of the PT-Tri stage, which still contained about 10\% of PTs at stage PT-Bi, all biological replicates cluster closely together and the three male gametophyte stages differ considerably from each other. These findings support data validity and accurate sampling. A PCA of all nine samples, including leaves, roots, flowers, tepals, and ovaries, shows that the three male gametophyte stages group very closely together (Fig. 4D). In conclusion, the high overlap among biological replicates and the presence of different clearly separable subgroups in the PCAs underlines the high reproducibility among the different samples and indicates that transcriptomes of *A. trichopoda* male gametophytes at different developmental stages are highly similar.
Next, we aimed to identify differentially expressed genes (DEGs) among PG, PT-Bi, and PT-Tri stages. For that purpose, we compared PG versus PT-Bi, PG versus PT-Tri, and PT-Bi versus PT-Tri by using the DESeq2 program ($q < 0.05$; Supplemental Table S4A). We consider those genes as DEGs whose TPM levels are $\geq 10$ in all replicates of at least one of the compared tissues and a $\log_{2}FC \leq -2$ and $\log_{2}FC \geq 2$, respectively (Supplemental Table S4B). A total of only 301 DEGs were detected between PG and PT-Bi, 220 between PG and PT-Tri, and 26 between PT-Bi and PT-Tri. In order to obtain further insights into functional information during male gametophyte transition from PGs to tri-cellular PTs, we analyzed biochemical processes defined by MapMan bins (Usadel et al., 2005) in a pairwise manner with identified DEGs (Supplemental Table S4). Supplemental Figure S2 shows a summary bar chart representing the number of DEGs that are down- or up-regulated in each comparison for a given bin. Despite the fact that many DEGs in each comparison are not assigned to any biological process, especially genes related to cell wall and cytoskeleton metabolism, protein degradation and synthesis as well as transcription regulation, signaling, and transport are up-regulated in PTs in PT-Bi and PT-Tri in comparison to PG. Only a few unregulated DEGs were identified in a PT-Bi-PT-Tri comparison, which are primarily associated to stress responses.

Additionally, we sought to explore the transcriptome dynamics between PG, PT-Bi, and PT-Tri stages and identify those genes with a similar expression pattern by a k-mean clustering analysis (Fig. 5; Supplemental Table S5A). The three male gametophyte stages show an overlap of 9,759 expressed genes and distribute into seven k-mean clusters. The largest cluster 2 containing 2,594 genes (26.58% of all overlapping genes) shows comparable expression levels in all samples. Stage-specific patterns are present in six out of the seven clusters, although, in general, transcript levels are only moderately changed (e.g. down-regulated to 50% in PTs compared to PGs in cluster 3). Cluster 1 and cluster 4 (together about 25% overlapping genes) show genes up-regulated during PT growth. These encode genes homologous to Arabidopsis genes coding for RALF, LRX, and BUPS, as well as other receptor kinases like PRKs (see next section and Supplemental Table S5B for details about gene identifiers). Cluster 6 and cluster 7 (together about 22% overlapping genes) display genes transiently upregulated at the PT-Bi stage. Among these genes there are a polygalacturonase QRT3 with a reported role in degrading the pollen mother cell wall during microspore development, a pollen-specific aquaporin (NIP4;1) required for pollen development, and a SET DOMAIN GROUP (SDG) gene reported to be involved in H3K4 methylation and important for gametophyte development. Additionally, these genes include a protein similar to the $\alpha$-fucosyltransferase AIOFT1, whose mutation produces lower seed set and reduced fertility. Finally, cluster 3 and cluster 5 (together about 26% overlapping genes) show genes downregulated during PT growth. These include homologs of transcriptional regulators like AtPV42a and AGL104, members of MIKC MADS box genes, and other transcription factors like BZIP34. Genes associated with the cell cycle such as a regulator of the anaphase-promoting complex/cyclosome (SAMBA) or regulators of actin organization such as a cyclase associated protein1 (CAP1) are also downregulated, while homologs of signaling proteins like pollen-specific receptor kinases (PRK3 and PRK6) and pollen-specific RHO-RELATED PROTEIN FROM PLANTS1 GTPase (ROP1) appear only slightly downregulated.

We also explored which genes are enriched in male gametophytic tissues in comparison to sporophytic samples. To do so, A. trichopoda genes with an average expression of TPM $\geq 1$ in a given tissue and TPM value $\geq 3$ in all the sporophytic tissues (ovule, leaf, tepal, and root) were considered as enriched or preferentially

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Table 1. Overview of RNA-seq data obtained from nine A. trichopoda tissues

<table>
<thead>
<tr>
<th>Tissue Type</th>
<th>Sample</th>
<th>Mapped Reads</th>
<th>No. of Expressed Genes</th>
<th>No. of Preferentially Expressed Genes$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male Gametophytic Tissues</td>
<td>PG</td>
<td>46.60</td>
<td>10,134 (37.10)</td>
<td>470 (1.72)</td>
</tr>
<tr>
<td></td>
<td>PT-Bi</td>
<td>63.88</td>
<td>11,384 (41.67)</td>
<td>652 (2.38)</td>
</tr>
<tr>
<td></td>
<td>PT-Tri</td>
<td>58.01</td>
<td>11,002 (40.28)</td>
<td>611 (2.23)</td>
</tr>
<tr>
<td>Flower Tissues</td>
<td>MALE</td>
<td>80.25</td>
<td>15,611 (57.15)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>FEMALE</td>
<td>81.96</td>
<td>16,802 (61.51)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>OVARY</td>
<td>8.91</td>
<td>6,144 (22.49)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>TEPAL</td>
<td>50.17</td>
<td>14,396 (52.70)</td>
<td>–</td>
</tr>
<tr>
<td>Vegetative Tissues</td>
<td>LEAF</td>
<td>22.55</td>
<td>15,003 (54.93)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>ROOT</td>
<td>18.60</td>
<td>16,174 (59.22)</td>
<td>–</td>
</tr>
</tbody>
</table>

$^a$Genes with a TPM $\geq 1$ in a given tissue and TPM $\leq 3$ in sporophytic tissues (ovary, leaf, tepal and root; see Supplemental Table S4).
expressed in that tissue. Despite the fact that this criterion could potentially identify genes with the same TPM value (e.g. TPM = 2) as enriched in a given tissue, in our data there are no genes that present this overlapping TPM. According to this criterion, the number of genes preferentially expressed in male gametophytic tissues are 470 in PG, 652 in PT-Bi, and 611 in PT-Tri (Table 1; see Supplemental Table S6A for details). These genes include pollen-specific genes described above and below, but again overall expression levels are similar in the three stages. In summary, the transcriptomes of the three different male gametophyte-specific stages are very similar and the vast majority of genes expressed at comparable levels.

Putative Orthologs of Essential Genes for Arabidopsis Male Gametophyte Functions Can Be Identified in Amborella trichopoda

To understand Amborella trichopoda-specific male gametophyte functions, we next explored the 30 most strongly expressed genes in the three developmental stages (Supplemental Table S6B). These genes are very highly expressed with TPM values up to 88,000. Expression values in the three stages were comparable. The six most highly expressed genes display TPM values between 25,000 to around 80,000. The corresponding predicted proteins do not have significant similarities with other described proteins and thus are prime candidates for functional studies. Notably, among the 30 most strongly expressed genes in PG, PT-Bi, and PT-Tri samples, we found 15, 13, and 14 genes lacking homology to known proteins, respectively (Supplemental Table S6B). Other highly expressed genes are homologous to genes coding for RAPID ALKALINIZATION FACTORS (RALFs), lipases/acylhydrolases with a Gly-Asp-Ser-Leu-like motif, allergen-like proteins, invertases, and pectin lyases. Because many genes essential for male gametophyte functions, including PT polarity, PT cell wall, and transport, physiology, growth, and guidance, as well as sperm release, have been characterized in the model flowering plant Arabidopsis, we next investigated the expression profiles of the respective homologous genes in Amborella trichopoda. Based on extensive literature data, we searched homologs for more than 200 genes with known functions in PGs and PTs in Arabidopsis. Following a BLASTp approach based on “best-hit,” many homologous genes were identified (Supplemental Table S7). Among the top 30 mostly strongly expressed Amborella trichopoda homologs, we identified genes encoding cell wall proteins like pectin methyl-esterases including AtPPME1 and Vanguard1 (VGD1), a RALF4/19 homolog, and one gene encoding a
LEU-RICH REPEAT/EXTENSIN (LRX) protein known to interact with RALFs (Mecchia et al., 2017). Most genes from the top 30 homology list encode proteins involved in signaling, including a number of receptor kinases like BUDDHAS PAPER SEAL1 (BUPS1) and BUPS/2 and small GTPases such as RHO-RELATED PROTEIN FROM PLANTS1 (ROP1) and ARABIDOPSIS RAB GTPASE HOMOLOG D2B (RABD2B) and RABD2C, but we also found MADS box keratin binding domain (MIK) C-type MADS box transcription factors. We could not identify homologs for 253 genes expressed in PG, 346 genes in PT-Bi, and 331 genes in PT-Tri by BLASTp searches, and about one-quarter of the 210 putative homologous genes listed in Supplemental Table S7 are lowly expressed in pollen with TPMs.

As shown in Figure 3, the cell wall of A. trichopoda consists of pectin and callose, similar to Pts of Arabidopsis and other eudicots and monocots (Williams, 2008). It is thus not surprising that cell wall synthesis and modifying enzymes appear conserved, but so do general signaling components like small GTPases. To study conservation of key signaling pathways during pollen functions in A. trichopoda, we next focused on the presence and detailed expression pattern of different classes of receptor-like kinases (RLKs). To identify homologous and putative orthologous RLKs expressed in Arabidopsis and A. trichopoda pollen, we first compared annotated protein sequences phylogenetically and studied the expression pattern of the corresponding genes (Fig. 6). In Arabidopsis, especially members of the largest RLK subfamily, LRR-RLKs, have been shown to possess key functions in PT growth and guidance. These include pollen-specific receptor kinases (PRKs), MALE DISCOVERER1/2 (MDIS1/2), and MDIS-interacting RLKs (MIKs), as well as BAK1-ASSOCIATING RECEPTOR-LIKE KINASE1 (BARK1), STRUBBELIG-RECEPTOR FAMILY4/5 (SRF4/5), CYS-RICH RLK 1 (CRK1), and RECEPTOR-LIKE KINASE IN FLOWERS1 (RKF1). These were aligned with all 95 LRR-RLKs that were identified in the A. trichopoda genome (Fig. 6A; Liu et al., 2016). Based on the phylogenetic tree and expression pattern, we identified AMTR_s00130p00117400 as a putative ortholog of Arabidopsis MDIS1 and MDSI2. AMTR_00025p00206890 appears as a single homolog of PRK1, PRK2, PRK4, PRK5, and PRK7, while AMTR_00024p00252010 is the only homolog of PRK3, PRK6, and PRK8. This finding indicates an amplification of pollen-specific LRR-RLK genes in Arabidopsis. Notably, the genes coding for A. trichopoda proteins phylogenetically related to Arabidopsis MIK1 and MIK2 as well as to SRF4 and SRF5 do not show a remarkable expression pattern in Pts, similar to proteins in the same species.
clade with Arabidopsis CRK1 and RFK1. The *A. trichopoda* LRR-RLK phylogenetically related to Arabidopsis BARK1 (AMTR_00117p00096300) shows only a weak expression level in PTs. Some *A. trichopoda* RLK genes such as AMTR_00010p00238570 and AMTR_00077p00153450, which are moderately expressed in PTs do not possess predominantly pollen-expressed LRR-RLK homologs in Arabidopsis. Most of the other *A. trichopoda* LRR-RLK genes are either not expressed in the tissues investigated or are expressed in vegetative tissues.

Another subclass of RLKs called *Catharanthus roseus*-RLK1-like kinases (CrRLK1Ls) is shown in Figure 6B. This subclass has 17 members in Arabidopsis and nine in *A. trichopoda*. Expression pattern and sequence similarity suggests that AMTR_00024p00049240 and AMTR_00045p00071050 are putative orthologs of BUPS1 and BUPS2 from Arabidopsis, while AMTR_s00016p00259140 appears to be the only putative ortholog of ANX1 and ANX2. The higher number of Arabidopsis RLK family members appears to be a general feature and suggests that gene duplication and diversification especially contributed to the expansion of RLK genes in evolutionary younger angiosperm species, but not *A. trichopoda*.

Genes encoding RALFs, which interact as mobile ligands with CrRLK1Ls, but also with cell-wall-located LRX proteins, confirm this finding. While the Arabidopsis genome encodes 37 RALF genes, *A. trichopoda* contains only nine genes, of which four are significantly expressed in male gametophytes (Fig. 6C). AMTR_s0022p00078480 and AMTR_s00044p00139580 are like the putative RALF4/19 orthologs regulating cell wall stability during PT growth, while AMTR_s00045p00207290 and AMTR_s00067p00137560 are members of a unique clade and together with egg apparatus expressed AMTR_s00067p00138490. Function(s) for these RALFs have not been elucidated. In conclusion, the combination of phylogenic and expression pattern analyses was successful in identifying candidate orthologs of Arabidopsis RLKs and their ligands with key functions in PT germination, growth, guidance, and burst in *A. trichopoda*.

Proteomic Studies Reveal That *A. trichopoda* PGs Are Well Prepared for Germination, While PTs Are Characterized by Catabolic, Biosynthetic, and Transport Processes

To obtain an additional level of insight into *A. trichopoda* male gametophyte functions, a proteomic analysis was conducted to compare mature *A. trichopoda* PGs and tricellular PTs. Three biological replicates each of PGs and PTs were processed and analyzed by liquid chromatography-tandem mass spectrometry (LC-MS/MS). Total protein extracts from three replicates for both samples were separated by SDS-PAGE and silver stained (Fig. 7A). PG and PT samples clearly displayed a different banding pattern pointing toward significant differences in protein levels. In total, 2,632 protein groups (Supplemental Table S8) were detected by LC-MS/MS. There were 1,667 protein groups were identified in PG and 2,288 protein groups in PT samples. A Venn diagram shows an overlap of 64.1% and 68.4% among the three PG and PT replicates, respectively (Fig. 7B). An additional PCA analysis groups the three biological replicates of PG and PT closely together and significantly apart from each other (Fig. 7C). These results indicate that the proteome of PG and PT are significantly different, while the biological replicates are very similar and thus reproducible.

For further analysis, we considered only those proteins that were simultaneously detected in all three biological replicates from either PG (1,086 protein groups) or PT (1,581 protein groups). Among these, 270 protein groups were significantly enriched in PG and 926 in PT samples (q < 0.05, fold change > 2). However, only 11% and 15% expressed genes, respectively, have been detected as proteins, and thus it is difficult to compare transcriptome and proteome data. In order to elucidate the corresponding biological processes that these enriched proteins represent, a functional category analysis was performed by PANTHER v14.1 (Fig. 7D; Supplemental Table S9, A and B). The enriched functional categories were grouped into main biological processes. Within each biological process, the total number of protein accessions from PG (in red) and PT (in blue) is provided. Redox processes appear especially important for PG, which is enriched in proteins related to the electron transport chain, proteins involved in reactive oxygen species (ROS) biosynthesis and response to oxidative stress. Additionally, proteins related to oxylipin biosynthesis and lipid transport as well as carbohydrate metabolism are enriched. Altogether, this indicates that the PG is well prepared to start germination requiring a lot of energy, lipids, and sugars. Proteins involved in cell wall organization/biosynthesis further support this hypothesis. PT is especially characterized by processes related to biosynthesis, catabolism, and transport. Many proteins are required for carbohydrate, nucleotide, fatty acid, carboxylic acid, and amino sugar catabolism, while simultaneously biosynthetic processes play a major role to generate lipids, proteins, and macromolecules in general, including cell wall material. Intracellular transport and Golgi vesicle transport are further major functional categories. Finally, a large number of identified proteins are involved in transcription, indicating that gene expression is regulated during PT growth.

Notably, with exceptions like PG cell wall biosynthesis, many proteins involved in PG/PT-specific functions such as signaling processes were not detected in the proteome data. This is not surprising, taking into consideration that only 11% and 15% of the expressed genes were represented in the proteomic data and that soluble proteins and highly abundant proteins, including those for housekeeping enzymes, are enriched. However, from the list of identified putative homologs of more than 200 genes with known
Figure 6. Phylogenetic tree and heatmap of *A. trichopoda* RLK and RALF genes to identify their putative Arabidopsis orthologs. A, LRR-RLKs predominantly expressed in PTs of Arabidopsis (24 genes) and all LRR-RLKs identified in *A. trichopoda* (95 genes). Arabidopsis genes involved in PT growth and guidance are indicated by filled dots, while their putative orthologs are shown by empty dots. B, *CrRLK1L* receptor genes identified in *A. trichopoda* (nine genes) and Arabidopsis (17 genes). Filled triangles...
functions in PGs and PTs in Arabidopsis (Supplemental Table S7), we have also detected peptides from 28 potential orthologs in the proteome data (Table 2). These include cell wall proteins like pectin methyl-esterase VGD1-like and AtPPME1-like, as well as a single homolog of LRR-expansin-like LRX-like9 to LRX-like11 proteins. The single putative A. trichopoda ortholog of PRK-like1, PRK-like2, PRK-like4, PRK-like5, and PRK-like6 RLKs was also detected. Peptides from other highly expressed signaling proteins like RALFs or BUPS/ANX RLKs as well as other PRKs and membrane transporters were not detected in the proteome data. These might be detectable in pollen membrane and cell wall fractions or might not have been detected due to low protein amounts despite high transcript levels. Finally, a few proteins were identified, like dynamin-like and actin filament bundling protein or small GTPases, which are lowly expressed, but which were detected in all samples of either PG or PT, indicating their high protein stability and low turnover rate. In summary, the proteomic analyses show that PGs seem to be well prepared for germination requiring lipids, energy, but likely also ROS molecules for signaling, while PTs are especially characterized by catabolic/biosynthesis and transport processes, including gene expression. Further experimentation involving fractionation of PG and PT samples is required to increase protein numbers and to detect proteins in specific subcompartments.

DISCUSSION

The A. trichopoda Male Gametophyte Shows Typical Features of Other Extant Angiosperm Groups

In order to understand the evolution of male gametophyte functions in flowering plants, studies using A. trichopoda are unique, as this species diverged already about 130 million years ago from all other living angiosperms and thus may still contain, for example, ancient molecular mechanisms that may have been lost or have been amplified in sister clade angiosperms, for example, during reproductive isolation. Histochemical studies of A. trichopoda male gametophytes confirmed that PGs are bicellular at maturity as observed in A. trichopoda and other basal angiosperms from the Amborellales, Nymphaeales, Austrobaileyales clade like Trithuria bibracteata, Nuphar advena, or Austrobaileya scandens (Williams et al., 2014). Notably, bicellular PGs are also characteristic for a number of eudicots like Solanum lycopersicum (Chaturvedi et al., 2013) or Nicotiana tabacum (Bokvaj et al., 2015). We reported that division of the GC during in vitro PT growth is completed within 14 h after germination, while in S. lycopersicum and N. tabacum, for example, GCs produce sperm cells about 10 h after germination in vitro (Yang et al., 2005; Lu et al., 2015). Notably, it has been reported in N. tabacum that the GC does not produce two sperm cells unless reduced forms of nitrogen like specific amino acids or ammonium chloride are present in the medium (Read et al., 1993). CJ-PGM used for A. trichopoda PT growth does not contain reduced forms of nitrogen, and therefore it will be interesting to test in future studies whether its addition will accelerate mitosis in A. trichopoda GCs. Moreover, the vegetative nucleus always precedes the GC during the PT translocation phase and the two sperm cells during further PT growth, respectively, which is similar to many sister clade angiosperms (Heslop-Harrison and Heslop-Harrison, 1989; McCue et al., 2011; Palevitz, 1993).

According to a comparative study of PTGRs in different seed plants, PTs of A. trichopoda and other basal angiosperms grow much faster than those from gymnosperm species but slower than later-diverging angiosperms (Williams, 2008). Furthermore, when comparing studies of PTGRs in different seed plants both in vivo and in vitro systems, PTGRs tend to be significantly higher in vivo than in vitro (see Supplemental Table S2). However, we identified conditions allowing A. trichopoda PTs to grow in vitro even about 25% faster than in vivo, indicating that either support by the maternal tissues of the ovary is less important in A. trichopoda or that the germination medium selected mirrors quite well the natural environment of the ovary tissues. We further observed that PT growth appeared linear without noticeable oscillations that were reported in fast growing PTs like Lilium longiflorum and N. tabacum (Lassign et al., 2014). This does not exclude the existence of oscillation of secondary messengers like Ca2+ and ROS and the development of pH gradients, which have been reported to play a key role for polar tip growth in PTs of sister clade angiosperm species (Damineli et al., 2017; Mangano et al., 2018). Based on the above reported in vitro growth system, a transient transformation system for PTs could now be established to also address such questions in a basal angiosperm species.

Figure 6. (Continued.)

indicate Arabidopsis CRLK1Ls involved in maintenance of the PT integrity, and open triangles indicate putative A. trichopoda orthologs. The heatmap shows expression values (in TPM) of A. trichopoda CRLK1Ls. C, RALFs identified in A. trichopoda (nine genes) and Arabidopsis (37 genes). Filled squares indicate Arabidopsis RALFs involved in maintenance of the PT integrity, and empty squares indicate putative A. trichopoda orthologs. Black arrowheads point toward highest expression in male gametophyte tissues. The phylogenetic tree was generated by using full-length protein sequences and the maximum likelihood method with 100 bootstraps. Female flower, EA(S), EA(M), and EA(L) represent small, medium, and large egg apparatus cells from A. trichopoda described in Flores-Tornero et al. (2019).

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**A. trichopoda** Male Gametophyte Transcriptomes Are Less Complex Compared with Those of Eudicots and Monocots

The richest resource to elucidate and compare the evolution of male gametophyte functions are the transcriptomic data reported above. Similar to other species like Arabidopsis (Becker et al., 2003; Honys and Twell, 2003), rice (Wei et al., 2010), or maize (Davidson et al., 2011), the transcriptome of PGs and PTs of **A. trichopoda** is less complex (about 40% of all protein encoding genes) compared with those of vegetative tissues such as roots and leaves (expression of almost 60% of all protein encoding genes). Around 2% expressed genes are enriched in male gametophytes, which is lower compared with reports for other species like Arabidopsis, rice, *N. tabacum*, or Glycine max showing 4% to 11% enriched genes (Rutley and Twell, 2015). There exists a very high overlap between the PG and PT transcriptomes of **A. trichopoda** and even gene expression levels appear quite similar: less than 40% variation in expression levels were observed for more than two-thirds of expressed genes. This is different in Arabidopsis, rice, or *N. tabacum*, showing a significant and progressive decrease in the diversity of transcripts during different stages of pollen development and germination (Rutley and Twell, 2015). In Arabidopsis, it was also shown that the PT transcriptome changes significantly depending on its growth in vitro or in vivo, as it was reported that more than 300 genes related to signaling, cell extension, and transcription were only found in semi-in vivo PTs in comparison to those germinated in vitro (Qin et al., 2009). Thus, it
Table 2. Overview about A. trichopoda homologs of Arabidopsis genes reported to play a role in PG and PT functions, which were identified by using proteomic analyses.

TPM values represent the average expression of three biological replicates and are listed according to their expression level in PT-Tri. Significant enrichment of a given protein in PT or PG tissues is represented with an "x" (see Supplemental Tables S7 and S8 for details). Gene symbols according to putative Arabidopsis homologs.

<table>
<thead>
<tr>
<th>A. trichopoda Gene ID</th>
<th>Gene symbol</th>
<th>E-value</th>
<th>Description</th>
<th>Reference</th>
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<tr>
<td>AMTR_s00040p00188020</td>
<td>VGD1</td>
<td>1.00E–132</td>
<td>Rectin methyltransferase</td>
<td>Jiang et al., 2005</td>
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<td>AMTR_s00007p00264370</td>
<td>APPM1</td>
<td>5.00E–119</td>
<td>Rectin methyltransferase</td>
<td>Tian et al., 2006</td>
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<tr>
<td>AMTR_s00029p00245270</td>
<td>LXR10, LXR11, LXR8, LXR9</td>
<td>4.00E–47, 7.00E–45, 3.00E–48, 2.00E–46</td>
<td>Leu-rich repeat extensin-like protein</td>
<td>Mecchia et al., 2017; Fabrice et al., 2018; Wang et al., 2018</td>
</tr>
<tr>
<td>AMTR_s00025p00206890</td>
<td>PRK1</td>
<td>0.00E+00</td>
<td>Leu-rich repeat RLK</td>
<td>Takeuchi and Higashiyama, 2016</td>
</tr>
<tr>
<td>AMTR_s00066p00182490</td>
<td>COBL10</td>
<td>0.00E+00</td>
<td>GPI-anchored protein</td>
<td>Li et al., 2013</td>
</tr>
<tr>
<td>AMTR_s00007p00221730</td>
<td>ATAPY1</td>
<td>0.00E+00</td>
<td>Apyrase</td>
<td>Wolf et al., 2007</td>
</tr>
<tr>
<td>AMTR_s00099p00037820</td>
<td>ATP, GAPCp1, GAPCp2</td>
<td>3.00E–98, 7.00E–116</td>
<td>Mitochondrial ATP synthase subunit</td>
<td>Geisler et al., 2012</td>
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<td>AMTR_s00047p00209120</td>
<td>RGP1, RGP2</td>
<td>0.00E+00, 0.00E+00</td>
<td>UDP- L-arabinose mutase</td>
<td>Drakakaki et al., 2006</td>
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<td>AMTR_s00102p00131940</td>
<td>QRT3</td>
<td>2.00E–155</td>
<td>Polygalacturonase</td>
<td>Rhee et al., 2003</td>
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<tr>
<td>AMTR_s00153p00086670</td>
<td>EXO70C2</td>
<td>5.00E–158</td>
<td>Putative exocyst subunit</td>
<td>Synek et al., 2017</td>
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<tr>
<td>AMTR_s00024p00207140</td>
<td>PDAT1</td>
<td>0.00E+00</td>
<td>Phosphodiacylglycerol acyltransferase</td>
<td>Zhang et al., 2009</td>
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<tr>
<td>AMTR_s00055p00193060</td>
<td>FIM5</td>
<td>0.00E+00</td>
<td>Actin bundling factor</td>
<td>Wu et al., 2010</td>
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<td>AMTR_s00044p00070480</td>
<td>UGP1, UGP2</td>
<td>0.00E+00, 0.00E+00</td>
<td>UDP-Glc pyrophosphorylase</td>
<td>Park et al., 2010</td>
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<tr>
<td>AMTR_s00010p00252880</td>
<td>AuUtR1, AuUtR3</td>
<td>0.00E+00, 0.00E+00</td>
<td>UDP-Gal/ Glc transporter</td>
<td>Reyes et al., 2010</td>
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<td>AMTR_s00080p00116860</td>
<td>DFRP2A</td>
<td>0.00E+00</td>
<td>GTase</td>
<td>Backues et al., 2010</td>
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<tr>
<td>AMTR_s00010p00159850</td>
<td>PGM2</td>
<td>0.00E+00</td>
<td>Phosphoglucomutase</td>
<td>Egli et al., 2010</td>
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<td>RGP5</td>
<td>1.00E–137</td>
<td>UDP- L-arabinose mutase</td>
<td>Drakakaki et al., 2006</td>
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<td>AMTR_s00176p00051440</td>
<td>DRIP1C</td>
<td>0.00E+00</td>
<td>Dynamin-like protein</td>
<td>Kang et al., 2003</td>
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<td>AMTR_s00039p00217390</td>
<td>GACPc1, GACPc2</td>
<td>0.00E+00, 0.00E+00</td>
<td>Phosphate dehydrogenase</td>
<td>Muñoz-Bertomeu et al., 2010</td>
</tr>
<tr>
<td>AMTR_s00067p00130250</td>
<td>MGP1</td>
<td>7.00E–116</td>
<td>Mitochondrial F1F0-ATP synthase</td>
<td>Li et al., 2010</td>
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<td>AMTR_s00019p002040770</td>
<td>VLN5</td>
<td>0.00E+00</td>
<td>Actin filament bundling protein</td>
<td>Zhang et al., 2010</td>
</tr>
<tr>
<td>AMTR_s00002p00227840</td>
<td>ARGH1</td>
<td>0.00E+00</td>
<td>Arginase</td>
<td>Brownfield et al., 2008</td>
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<tr>
<td>AMTR_s00013p00172160</td>
<td>ATKDAS2</td>
<td>0.00E+00</td>
<td>Phosphate synthetase</td>
<td>Delmas et al., 2008</td>
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<td>AMTR_s00003p00020850</td>
<td>MIRO1</td>
<td>0.00E+00</td>
<td>GTPases</td>
<td>Sormo et al., 2011</td>
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<td>AMTR_s00019p00246290</td>
<td>PUR4</td>
<td>0.00E+00</td>
<td>Ribonucleotide synthase</td>
<td>Berthomé et al., 2008</td>
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<td>AMTR_s00155p00043060</td>
<td>IPMDH2, IPMDH3</td>
<td>0.00E+00, 0.00E+00</td>
<td>3-Isopropylmalate dehydrogenase</td>
<td>He et al., 2011</td>
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<td>AMTR_s00022p00227570</td>
<td>RABA4D</td>
<td>8.00E–138</td>
<td>Rab GTPase</td>
<td>Szmulanski and Nielsen, 2009</td>
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<tr>
<td>AMTR_s00103p00075300</td>
<td>STP6</td>
<td>0.00E+00</td>
<td>Hexose transporter</td>
<td>Scholz-Starke et al., 2003</td>
</tr>
</tbody>
</table>
cannot be ruled out that such differences in expression will also occur in *A. trichopoda* semi-in vivo-grown PTs. All together, these findings suggest that the regulation of male gametophyte functions in sister clade angiosperms is likely more complex and imply that the *A. trichopoda* male gametophyte is more similar to that of the most recent common ancestor of all angiosperms. Likewise, the egg apparatus cells (egg and synergids) of *A. trichopoda* are morphologically highly similar and contain a rather large overlap of jointly expressed genes (Flores-Tornero et al., 2019).

A detailed analysis by comparing the *A. trichopoda* male gametophyte transcriptomes with that of other angiosperms identified homologs of a large number of well-known and characterized proteins with male gametophyte functions. Many conserved genes encode proteins involved in cell wall metabolism and composition (Wallace and Williams, 2017; Fabrice et al., 2018), including pectin methyltransferases, polygalacturonases, and LRR-extensin-like proteins that were also detected in the proteomic data. This finding confirms the histological data showing that the composition and arrangement of pectin and callose in the pollen cell wall in *A. trichopoda*, as well as the formation of extensive callose plugs, is typical for angiosperms, especially as described for *C. japonica* and *L. longiflorum*, respectively (Hasegawa et al., 2000). Notably, this is different from gymnosperm PT cell walls that show a different arrangement and composition, such as arbovirilacatan protein abundance and absence of pectins and callose depositions (Yatomi et al., 2002). As another remarkable finding, we detected high transcript levels of some genes that were not detected at the proteomic level and vice versa obtained enriched proteins in PGs or PTs, whose transcript levels were very low. Low correlation between protein abundance and mRNA expression levels has been reported before (e.g. Greenbaum et al., 2003; Walley et al., 2016) and can be explained as follows: on the one hand, the male gametophyte of plants shows a high level of translational regulation and thus transcripts are stored in the mature PG and are only translated during pollen germination in vivo (e.g. Fila et al., 2017). Some proteins are very stable and thus are only represented by low transcript levels. Moreover, technical issues associated with proteomic studies still have a strong impact on detected proteins as a whole battery of protocols would be necessary to detect all proteins that are located at different membranes, organelles, cell wall, etc. Therefore, transmembrane proteins like BUPS1/2 and ANX1/2 or small secreted proteins (e.g. RALFs) can easily be missed or are underrepresented in proteomic studies. Usually, cytoplasmic proteins are particularly enriched in proteomic studies (Walley et al., 2016).

### Identification of Genes Likely Essential for PT Functions

Both transcriptomic and proteomic data indicated that processes related to lipid and protein biosynthesis, as well as to transport mechanisms, are significantly upregulated during PT growth, which has also been shown in other angiosperms like rice (Dai et al., 2006), Arabidopsis (Zou et al., 2009), *S. lycopersicum* (Chaturvedi et al., 2013), or *N. tabacum* (Ischebeck et al., 2014). Moreover, the RALF-ANX/BUPS-LRR signaling pathway regulating PT cell wall integrity (Boisson-Dernier et al., 2009; Miyazaki et al., 2009; Ge et al., 2017, 2019) appears conserved, as homologs of all components could be clearly identified in the combined phylogenetic expression data. However, *A. trichopoda* additionally encodes a male/female gametophyte-specific RALF clade (Flores-Tornero et al., 2019) that does not contain homologs in other angiosperms (Galindo-Trigo et al., 2016) and may possess an ancient function or a function specific to basal angiosperms from the Amborellales, Nymphaeales, Austrobaileyales clade. Relatively few of the 94 LRR-RLKs encoded by the *A. trichopoda* genome (Liu et al., 2016) are expressed in male gametophytes. These include homologs of the Arabidopsis plasma membrane receptors PRK6 and MDISI1/2 that interact with female gametophyte-secreted PT attractant LURE1 peptides (Takeuchi and Higashiyama, 2016; Wang et al., 2016). Notably, putative homologs of further LURE1 interactors MIK1 and MIK2 (Wang et al., 2016) are not significantly expressed in the male gametophytes of *A. trichopoda*. It should also be emphasized that the number of PT-expressed PRKs including PRK1 and PRK3 that contribute to LURE1 binding is considerably expanded in Arabidopsis. The higher number of Arabidopsis RLK family members appears to be a general feature and suggests that gene duplication and diversification especially contributed to the expansion of RLK genes in evolutionary younger angiosperm species, but not *A. trichopoda*. Thus, taking into consideration that *A. trichopoda* lacks homologs of PT attractants like Arabidopsis LURE1 or maize EA1 peptides (Márton et al., 2005, 2012) and that Arabidopsis, for example, contains significantly more RLKs and candidate signaling peptides like CRPs expressed in male gametophytes, indicates that communication of PTs with the maternal tissues and the female gametophyte is less complex and more ancient in *A. trichopoda* and points to the possibility that amplification of these signaling mechanisms including polymorphic peptide ligands contributed significantly to reproductive isolation and thus angiosperm diversity.

### Outlook

It will now be exciting to elucidate components and mechanisms regarding how and whether *A. trichopoda* can discriminate self from alien PGs, whether PT guidance mechanisms exist, and which are the roles of the highly and specifically expressed genes in *A. trichopoda* male gametophytes lacking homologs in sister clade angiosperms. The data generated in this study can be used for comparative gene expression studies, but also to select candidate genes and protein domains for
investigations about protein evolution during angiosperm reproduction and diversification and can be used to complement, for example, corresponding mutants in Arabidopsis and other species. Functional studies of *A. trichopoda* male gametophyte-specific genes is still very difficult, as stable transformation systems are not available and cultivation of the species is quite challenging. However, overexpression studies of *A. trichopoda* male gametophyte-specific genes in other species, the development of a transient PT transformation system, and complementation approaches of reproductive mutants in sister clade angiosperms like Arabidopsis and maize with *A. trichopoda* genes could significantly help to better understand the evolution of reproductive gene families and mechanisms and thus could significantly help to better understand the evolution of the reproductive mechanism in this fascinating plant species and in flowering plants in general.

**MATERIALS AND METHODS**

### Plant Material and Growth Conditions

Plant material was harvested from male and female flowers of *Amborella trichopoda* at the Botanical Garden in Bonn (Germany). Plants were grown in a shaded place inside a greenhouse under controlled conditions of 16°C to 18°C, constant humidity of 66%, and 12-h photoperiods. Tissues including male and female flowers, tepals, leaves, and roots were immediately frozen in liquid nitrogen after harvest and stored at −80°C. Fully opened male flowers were gathered in 50-mL Falcon conical tubes (Thermo Fisher), placed without lid in a hermetically sealed plastic box containing a bed of silica gel, and stored at 4°C until pollen isolation. Ovaries were isolated from fully opened female flowers, immediately frozen in liquid nitrogen, and stored at −80°C.

### Pollen Isolation and Germination

As shown in Figure 1, male flowers were collected in 50-mL Falcon conical tubes (Thermo Fisher) up to a volume of 25 mL. Next, tubes containing flowers were transferred without lid to a moisture chamber at room temperature for up to 30 min to allow rehydration. Afterward, 25 mL of fresh sterile-filtered liquid of CJ-PCM (5% [w/v] Suc, 300 μg mL−1 Ca(NO3)2, 200 μg mL−1 MgSO4, 100 μg mL−1 KNO3, 100 μg mL−1 H3BO3, pH 5.0) was added. The mixture was vortexed for 30 s at maximum speed to release PGs. Subsequently, the liquid germination medium containing PGs was filtered through a 1-mm pore size steel mesh to remove flowers tissues and debris. The pollen suspension was further filtered in a 50-mL Falcon conical tube containing a扑室ininer 70-μm polyester mesh (PluriSelect) and a connector ring (PluriSelect) in which the sample is force-filtered using manual pressure from a syringe. After this filtration step, small debris were retained on the surface of the mesh and the pollen suspension was collected and refiltered through a stack of 70-30-15-micron pore size polyester strainers (PluriSelect) to remove cellular debris and separate bacterial and fungal spores from PGs. Finally, PGs were obtained as a beige-white clean film on the 15-micron strainer and removed from the membrane using a spatula placed in a 1.5-mL collection tube and resuspended in 0.5-mL CJ-PCM. For germination in liquid medium, 1 mL of PG suspension of OD490 = 1 was resuspended in a final volume of 5 mL CJ-PCM, poured into 5.5-cm diameter glass petri dishes, covered with a lid, and incubated in a moisture chamber at room temperature for up to 20 h on the laboratory bench. To estimate PTGR in solidified medium, a small square plastic chamber (2 × 2 cm) was built on a glass slide, filled with 3 mL sterile CJ-PCM with 0.5% (w/v) low melting agarose (Carl Roth) and 0.2 μmol NaH2PO4, 10 mM EDTA, 0.05% (w/v) Triton X-100, pH 7.2. Liquid CJ-PCM was used to germinate mature PGs, and PTs were visualized with a Nikon Eclipse TE2000-S inverted microscope. To detect the presence of pectin in PGs and PTs, liquid CJ-PCM with 0.01% (w/v) ruthenium red (Sigma-Aldrich) was used. For callus plug staining, decolorized aniline blue solution (0.1% [w/v] aniline blue in 108 mM K3PO4, pH 11.0) was added to liquid medium. A Zeiss Axioskop was used to visualize stained PGs and PTs. The presence of sperm cells and vegetative nuclei in PTs was determined by staining with SYBR Green I nucleic acid stain solution (Thermo Fisher) in a 1:1000 solution in combination with the membrane dye FM 4-64 (N-(3-triethylammoniumpropyl)-4-(6-dimethylaminophenyl)hexatrienyldibromide; Thermo Fisher) and visualized using a Leica TCS S8 confocal laser scanning microscope.

### RNA Isolation, cDNA, and Library Preparation

Total RNA was isolated from three biological replicates of control tissues (tepals, leaves, and roots) as well as from PGs, PTs 10 h after germination (PT-Bi), and PTs 20 h after germination (PT-Tri) by using a Spectrum plant total RNA kit (Sigma-Aldrich) according to manufacturer’s instructions. Two hundred fifty nanograms of total RNA was used for each library construction. RNA-seq was carried out as described in the Illumina TruSeq stranded mRNA sample preparation guide for the Illumina HiSeq 1000 System (Illumina) and the KAPA library quantification kit (KAPA Biosystems). For ovary samples, three biological replicates containing 11 ovaries each were used to extract total RNA according to the “Purification of total RNA from animal and human cells” protocol of the RNeasy Plus micro kit (Qiagen). The SMARTer ultra low input RNA kit for sequencing v4 (Clontech Laboratories) was used to generate first strand cDNA from 0.5 ng total RNA. Double-stranded cDNA was amplified by LD PCR (15 cycles) and purified via magnetic bead cleanup. Library preparation was carried out as described in the Illumina Nextera XT Sample Preparation Guide (Illumina). One hundred fifty picograms of input cDNA were tagmented (tagged and fragmented) by the Nextera XT transposome. Products were purified and amplified via a limited-cycle PCR program to generate multiplexed sequencing libraries. For the PCR step, 1.5 dilutions of index 1 (G7) and index 2 (G5) primers were used. Libraries were quantified using the KAPA SYBR FAST ABI prism library quantification kit (Sigma-Aldrich). Equimolar amounts of each library were used for cluster generation on the cBot (TruSeq SR cluster kit v3; Illumina). Sequencing runs were performed on a HiSeq 1000 instrument using the indexed, 2 × 100 cycles paired-end protocol and the TruSeq SBS v3 kit. Image analysis and base calling were performed in “bcl” format and converted into “fastq” files using the CASAVA1.8.2 software (Illumina). Illumina deep sequencing was carried out at the genomics core facility of the University of Regensburg (Center for Fluorescent Bioanalytics KFB).

### RNA-seq Data Analysis

RNA-seq data were processed using the LSTRAP-Lite pipeline (Tan and Mutwil, 2020), where Trimomatic with default settings was used to trim sequence reads. All samples passed LSTRAP’s quality control. Reads were mapped to the *Amborella* genome v1.0 (Amborella Genome Project, 2013) obtained from Ensembl Plants (Kersey et al., 2016). Log2FC and Benjamini and Hochberg-corrected *P* values (Benjamini and Hochberg, 1995) for DEGs were calculated using DESeq2 (package version 1.10.1; Love et al., 2014). Significant DEGs (*P* < 0.05 after correction) were used to identify significantly enriched MapMan bins (MapMan v 3.6.0RC1; Thimm et al., 2004). Finally, gene expression levels were estimated using TPM values (Conesa et al., 2016) and clustered using Python Seaborn package with ‘Euclidean’ distance and ‘complete’ method. To estimate similarities between transcriptomes of the various samples, we used Jaccard index defined as: A(X,Y) = |X∩Y| / |X∪Y|, where X and Y are differentially regulated (e.g. up-regulated) genes in the two compared samples. K-means clustering and finding the optimal k with sum of squared distances was calculated using Python package scikit-learn (Pedregosa et al., 2011). Genes with TPM values ≥ 1 were considered as expressed. A comprehensive overview on normalized expression data are shown in Supplemental Table S3. RNA-seq data are deposited at the European Bioinformatics Institute under accession number E-MTAB-9190 (https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-9190/).
RT-qPCR Validation of RNA-seq Data

For RNA-seq data validation, RT-qPCR was performed with nine randomly selected genes, including two genes with similar expression levels in all three tissues as reference genes and seven DEGs representing high, medium, and lowly expressed genes, respectively. RNA from PG, PT-Bi, and PT-Tri samples was isolated as previously described, and 100 ng of each sample was used to prepare cDNA using SuperScript III reverse transcriptase (Thermo Fisher) according to the manufacturer’s instructions. Primers used for RT-qPCR are listed in Supplemental Figure S1. RT-qPCR was performed in 10-μL volumes with 1 μL cDNA by using KAPA SYBR FAST universal kit (Merck). Reactions were performed in three technical replicates per sample. RT-qPCR was performed using the Mastercycler Realplex2 (Eppendorf) in a 96-well reaction plate according to the manufacturer’s recommendations. Data were normalized to the expression levels of the two reference genes, and relative expression was calculated using the comparative Ct method (2^ΔΔCt). ΔCt values were compared to log2FC (TPM in tissue A versus tissue B) using Pearson’s correlation. The correlation of RNA-seq and RT-qPCR data are shown in Supplemental Figure S1.

Protein Extraction and LC-MS/MS Analysis

PG pellets were obtained from three independent experiments. Fifty milligrams of pellets from each isolation were immediately frozen in liquid nitrogen and stored at −80°C. The rest of the pellets from each isolation were resuspended in 1 mL liquid CJ-PGM until OD600 = 1, diluted to a final volume of 5 mL germination medium, and incubated at room temperature for 20 h in a moisture chamber. PTs were collected by filtration through a 40-μm strainer (PluriSelect) and washed twice with liquid germination medium to remove nongerminated PGs. Fifty milligrams of each isolation of PT pellet was collected from the surface of the strainer with a spatula, placed in a 1.5-mL collection tube, and immediately frozen in liquid nitrogen. Total proteins were extracted by grinding samples with liquid nitrogen in a mortar and subsequent addition of 250 μL of ice-cold extraction buffer (50 mM Tris/HCl, 150 mM NaCl, 0.1% [w/v] sodium deoxycholate, 0.1% [v/v] Triton-X100, 1 mM phenylmethylsulfonyl fluoride, pH 8.0). Pollen protein extracts were cleared twice in a bench centrifuge (15,000 rpm, 10 min, 4°C). Dilutions of equal protein concentrations were prepared in triplicate and analyzed in the Microcon device with molecular weight cut-off 30 kD (Merck Millipore) and used only those genes with TPM > 1. PCA analysis was done by using MetaboAnalystR 3.0 online tool (Pang et al., 2020) with already normalized data. Colored regions from each tissue represent areas with 95% confidence to consider the replicates reproducible. GO term enrichment was performed online by using PANTHER v14.1 with a significance level of P < 0.05 and correction of false discovery rate < 0.05. See the “RNA-seq Data Analysis” section for statistical analysis related to DEGs. The LRR-RLK genes from Amborella and Arabidopsis (Arabidopsis thaliana) were used for phylogenetic analysis was obtained from Liu et al. (2016) and Takeuchi and Higashiyama (2016) respectively, whereas the CrRLK1L genes were all obtained from Galindo-Trigo et al. (2016). Phylogenetic analyses were performed with MEGA7 software (Kumar et al., 2016) by using full-length protein sequences, aligning them by selecting the ClustalW option and using the maximum likelihood method with 500 bootstraps. Heatmaps were generated in Excel by using TPM values obtained from CoNeKt online database (https://europro.db.sbs.nus.edu.sg).

Accession Numbers

The raw sequencing data sets generated and analyzed during this study are available at the European Bioinformatics Institute under accession number E-MTAB-9190. Please see Supplemental Table S3 for detailed expression values of all annotated A. trichopoda genes.

Supplemental Data

The following supplemental materials are available.

- Supplemental Figure S1. Validation of RNA-seq data by RT-qPCR.
- Supplemental Figure S2. DEG dendrogram of A. trichopoda PG and PT transcriptomes using MapMan ontology analysis.
- Supplemental Table S1. PGM tested by using A. trichopoda PGs.
- Supplemental Table S2. A. trichopoda PTGR observed in this study compared with another A. trichopoda study and those for other seed plants.
- Supplemental Table S3. Gene expression values of all A. trichopoda tissues described in this report.
- Supplemental Table S4. DEGs in male gametophyte tissues.
- Supplemental Table S5. K-means clustering of all 9759 genes expressed in A. trichopoda PG, PT-Bi, and PT-Tri.
- Supplemental Table S6. Most strongly expressed genes detected in male gametophytic tissues of A. trichopoda.
- Supplemental Table S7. List of 210 Arabidopsis genes reported to play a role in PG and PT functions, their putative A. trichopoda orthologs, and average gene expression pattern.
- Supplemental Table S8. Identification of proteins expressed in A. trichopoda PG and PT by mass spectrometry, and differentially expressed proteins (DEP) in pairwise comparisons between PG and PT.
- Supplemental Table S9. List of gene ontology terms of significantly enriched proteins in A. trichopoda PG and PT.

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