ASYMMETRIC EXPRESSION OF ARGONAUTES IN ARABIDOPSIS REPRODUCTIVE TISSUES

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ABSTRACT

During sexual reproduction, development of a totipotent zygote from the fusion of highly differentiated gametes is accompanied by a dynamic regulation of gene expression. This notably involves RNA silencing operated by Argonautes (AGO) effector proteins. While AGOs’ roles during Arabidopsis somatic life have been extensively investigated, less is known about their expression during reproduction, which proceeds via double-fertilization of an egg and a central cell, leading respectively to the embryo and a supportive tissue known as endosperm. Using full-locus translational reporters for all ten Arabidopsis AGOs, we uncover cell-specific expression patterns and AGO-intrinsic subcellular localizations in reproductive tissues. However, while some Arabidopsis AGOs are enriched in both male and female gametes, i.e. sperm and egg cells, they are comparably low-expressed in accessory, i.e. vegetative and central cells. Likewise, following fertilization, several AGOs are expressed in the early embryo, yet below detection in the early endosperm. Thus, there is pre- and post-fertilization asymmetry between the embryo and endosperm lineages. Later during embryo development, AGO9, AGO5 and AGO7 are restricted to the apical embryonic meristem in contrast to AGO1, AGO4, AGO6 and AGO10. Beside shedding some light onto Arabidopsis reproduction, the plant material generated here should constitute a valuable asset to the RNA silencing community by enabling functional AGOs studies.
INTRODUCTION

In most eukaryotes, sexual reproduction involves specialized cellular structures and entails complex orchestration of the timing and spatial localization of gene expression. In the model plant Arabidopsis, reproduction involves the development of a haploid structure called gametophyte in both male and female reproductive organs. The mature male gametophyte, or pollen grain, contains three cells: one vegetative cell and two gametes known as sperm cells. The mature female gametophyte, on the other hand, contains seven cells: three antipodal cells of unknown function, two synergides involved in pollen tube reception and guidance, one central cell and one egg cell. Fertilization of the egg cell by one of the sperm cells forms the embryo proper that will later develop into the next-generation plant. Fertilization of the homodiploid central cell by the second sperm cell forms the endosperm, a nourishing tissue required for proper seed development. In addition, the female gametophyte as well as the developing endosperm and embryo are surrounded by 4 layers of maternal integuments connected to the mother plant’s vasculature at the chalazal pole.

Small RNAs (sRNA) are key regulators of gene expression. Their importance during sexual reproduction has been highlighted by the discovery of a class of reproduction-specific sRNA known as Piwi-interacting RNAs (piRNAs) in animals (Castel and Martienssen, 2013; Weick and Miska, 2014). Likewise, the importance of reproduction-specific small RNAs has also been recognized in plants (Mosher and Melnyk, 2010; Van Ex et al., 2011). Beyond the roles of microRNAs (miRNAs) in embryonic development, small interfering (si)RNA tame transposon in the pollen (Slotkin et al., 2009) and during ovule development (Olmedo-Monfil et al., 2010) but also could be linked to hybrid lethality as they were shown to be involved the regulation of parental genome dosage (Borges et al., 2018; Martinez et al., 2018). RNA silencing in plants can be divided into Post-Transcriptional Gene Silencing (PTGS) and Transcriptional Gene Silencing (TGS). Both pathways rely on the generation of small (s)RNAs of 21, 22 or 24 nucleotides in length by DICER-LIKE enzymes, of which there are four paralogs in Arabidopsis (DCL1-4). These sRNA execute PTGS or TGS upon their loading into ARGONAUTE (AGO) effector proteins. Despite the established impact of sRNA-pathway mutations in plant biology, little is known of the expression profiles of silencing-pathway proteins in reproductive and post-reproductive tissues. In particular, the cell-specific expression and subcellular localization patterns of AGOs remain largely unknown.
The *Arabidopsis* genome encodes ten AGO genes divided into three phylogenetic clades (Mallory and Vaucheret, 2010) conserved among flowering plants (Fang and Qi, 2015; Zhang et al., 2015). Clade I consist of AGO1/5/10, with AGO1 being the ubiquitously-expressed member. *ago1* mutants display aberrant sporophytic phenotypes likely due to AGO1’s key role in executing miRNA functions. Embryonic *ago1* mutant phenotypes were only observed, however, from the torpedo stage (Lynn et al., 1999). The second clade I member, AGO10, has been implicated in the control of shoot apical meristem (SAM) identity from late stages of embryo development (Lynn et al., 1999; Moussian et al., 1998), despite being expressed from very early stages (Tucker et al., 2008). In fact, early embryo phenotypes were only observed in the double *ago1ago10* mutant, thereby suggesting functional redundancy between the two proteins at this stage, possibly in executing miRNA-directed silencing. Supporting this notion, early embryo phenotypes are observed in dcl1 mutants exhibiting compromised miRNA processing (Nodine and Bartel, 2010; Seefried et al., 2014; Willmann et al., 2011). Contrasting with that of AGO1 and AGO10, AGO5 expression is substantially enriched in reproductive tissues (Tucker et al., 2012). A dominant AGO5 allele (*ago5-4*) arrests female gametophyte development altogether (Tucker et al., 2012) while *ago5* recessive mutants display early flowering (Roussin-Léveillée et al., 2020).

The AGO Clade II is constituted of AGO7, AGO2 and AGO3. AGO7 has a well-established role in leaf development yet no overt function in reproductive tissues has been describe thus far. Despite AGO2 and AGO3 displaying higher expression levels in reproductive tissue, their role(s) during reproduction remains, likewise, elusive because no developmental phenotypes were observed (Jullien et al., 2018). Clade III includes AGO4/6/9/8, which, with the exception of the truncated AGO8 (Takeda et al., 2008), have all been implicated in DNA methylation commonly referred as the RNA-dependent-DNA-methylation (RdDM) pathway (Matzke and Mosher, 2014). The RdDM pathway is required for the establishment of new DNA methylation pattern as well as the maintenance of cytosine methylation in the CHH context. AGO4 and AGO6 are mostly ubiquitously expressed, whereas AGO8 and AGO9 seem specific for reproductive tissue (Havecker et al., 2010; Olmedo-Monfil et al., 2010; Wuest et al., 2010). In particular, AGO9 is involved in meiosis, megaspore mother cell differentiation and transposon silencing in the egg cell (Oliver et al., 2014; Olmedo-Monfil et al., 2010). Megaspore mother cell differentiation is also impeded in *ago4*, *ago6* and *ago8* mutants (Hernández-Lagana et al., 2016) suggesting an important role for RdDM during this process.
To help deciphering the roles of sRNA pathways in reproductive processes, we have generated stable transgenic lines expressing full length fluorescently tagged AGOs under their cognate endogenous promoter and analyzed their expression as well as intracellular localization in reproductive tissues.

RESULTS AND DISCUSSION

ARGONAUTES TRANSLATIONAL REPORTERS

In order to investigate the expression pattern and intracellular localization of the ten Arabidopsis AGO proteins, we generated full-locus reporter constructs in which the open reading frames (ORFs) of fluorescent proteins were fused to each AGO coding sequence in their genomic contexts, using multiple Gateway™ cloning. We engineered N-terminal translational fusions, since N-terminal, but not C-terminal, tagging preserves Arabidopsis AGOs’ functionality (Carbonell et al., 2012). Each AGO construct was cloned under the corresponding presumptive promoter (1.3kb to 2.5kb upstream start codons) and terminators (467bp to 1kb downstream of stop codons). This generated pAGO:FP-AGO constructs, where “FP” corresponds to either the Green Fluorescent Protein (GFP) or mCherry (mCh). For the sake of simplification, the constructs will be referred to FP-AGOX (where X is the number of each AGO1-10) in the main text and FPX in the figures. For example, pAGO1:mCherry-AGO1 will be shortened to mCherry-AGO1 in the main text and to mCh1 in figures. Construct were transformed in their respective mutant background, at least 6 T1 were screened for each construct and all show a consistent expression pattern in the tissue examined. A detail map of the constructs can be found in Fig. S1 and a summary of their expression pattern in Fig.S11. Functional complementation was validated for AGO1, AGO4, AGO5, AGO6 and AGO7 and can be found in Fig. S2, Fig. S3, Fig. S4, Fig. S5 and Fig. S6 respectively.

AGO EXPRESSION PATTERNS IN THE MATURE FEMALE GAMETOPHYTE

To gain insights into AGO protein expression patterns in the mature female gametophyte, we analyzed fluorescent signals of each protein fusion, using confocal microscopy. Almost all AGOs accumulate in the mature female gametophyte (Fig. 1A-K) except GFP-AGO10 (Fig. 1D) and mCherry-AGO3 (Fig. 1J), which are only detected in the maternal integument. The clade I mCherry-AGO1 and mCherry-AGO5 reporters display similar accumulation patterns in the female gametophyte (Fig. 1B-C). They are mainly detected in the egg cell and, to a lower extent, in the central cell. mCherry-AGO1 and mCherry-AGO5 also accumulate in the maternal
integument with mCherry-AGO1 being detected in both the inner and outer integument, and mCherry-AGO5 solely in the former, as previously reported (Tucker et al., 2012). Both mCherry-AGO1 and mCherry-AGO5 signals are particularly pronounced in the nucellus at the chalazal pole of the ovule. Clade I, GFP-AGO10 mainly accumulates in the inner-integument of the ovule with a stronger signal at the chalazal seed coat and in vascular tissues of the funiculus (Fig. 1D, Fig. S7A-B).

The main clade III *i.e.* RdDM AGO's, mCherry-AGO4 and mCherry-AGO6, accumulate ubiquitously in both integuments and female gametophyte, with stronger signals in the egg cell (Fig. 1E-F). The reproduction-stage-specific RdDM AGO's, mCherry-AGO9 and mCherry-AGO8, show more spatially restricted expression patterns (Fig. 1G-H). mCherry-AGO9 accumulates mainly in the egg cell within the female gametophyte but can be detected, albeit at lower levels, in the central cell. Strong mCherry-AGO9 accumulation is also detected in the nucellus and funiculus (Figure1H and Fig. S8A), consistent, for the latter, with previous *in situ* hybridization results (Olmedo-Monfil et al., 2010). mCherry-AGO8 is specifically detected in the mature egg cell (Fig. 1G). Egg cell expression of both AGO8 and AGO9 is supported by conclusions drawn from transcriptional fusions for both AGO9 and AGO8 (Sprunck et al., 2019) as well as transcriptomic data (Fig. S9B, (Wuest et al., 2010)). However, mCherry-AGO9 protein fusion expression in the egg cell do not agree with previously published results obtained by immuno-fluorescence (Olmedo-Monfil et al., 2010). Like mCherry-AGO8, the clade II-member mCherry-AGO2 is also specifically detected in the egg cell (Fig.11) while clade II-member mCherry-AGO3 is solely detected in the chalazal seed coat of the ovule (Fig. 1J, (Jullien et al., 2018)). The last clade II-member, GFP-AGO7, is detected in all cell types of the female gametophyte and surrounding integument, with significant enrichment in the egg cell, similarly to AGO1 (clade I), AGO4 and AGO6 (clade III) (Fig. 1K).

Overall, our analysis shows that all ten *Arabidopsis* AGOs are detected in mature ovules before fertilization. Within the female gametophyte their accumulation seems to be particularly enriched in the egg cell compared to the central cell. Preferential AGO expression in the egg cell was confirmed using previously published female gametophyte transcriptome data obtained after laser-capture microdissection (Fig. S9A, (Wuest et al., 2010)). Our data together with the potential ago8 phenotype during MMC development (Hernández-Lagana et al., 2016) suggest that AGO8 might not be a pseudogene. AGO8 was thought to be a pseudogene as it possesses a premature stop codon predicted to result in a truncated protein containing a PAZ domain but not the catalytically active PIWI domain (Takeda et al., 2008). Our observation of...
mCherry-AGO8 protein specifically in the egg cell (Fig. 1G) could result from the accumulation of a truncated mCherry-AGO8 protein in this particular cell type. However, we could not test this hypothesis using Western blot due to the very low expression of mCherry-AGO8. It is nevertheless tempting to hypothesis that a truncated AGO8 protein could act as a dominant negative sRNA regulator specifically in the egg cell as it was shown for the dominant ago5-4 allele that similarly express a truncated AGO5 including the PAZ but not the PIWI domain (Tucker et al., 2012). Together our results suggest a complex regulation of sRNA loading and function in the egg cell.

**AGO EXPRESSION PATTERNS IN THE MALE GAMETOPHYTE**

Analysis of the translational reporters in mature pollen grain (Fig. 2A-H) revealed a preferential enrichment of some AGOs in sperm cells. Indeed, mCherry-AGO1, mCherry-AGO2, GFP-AGO7, mCherry-AGO4 and mCherry-AGO9 were solely detected in those cells (Fig. 2 B, D,F-H). mCherry-AGO5 and mCherry-AGO6 where mainly detected in sperm cells, but also in the vegetative cell, albeit only at substantially lower levels (Fig. 2 C, E). AGO5 sperm cell expression was previously reported (Borges et al., 2011; Tucker et al., 2012). The mCherry-AGO3, mCherry-AGO8 and GFP-AGO10 reporters signals are not detected in mature pollen.

In order to address if paternally-expressed AGOs could be potentially transmitted during fertilization, we investigated the presence of the protein fluorescent signals in germinated pollen (Fig. 2I-P). All AGOs showing expression in sperm cells were indeed also detected in germinated pollen, suggesting that AGO-loaded sRNAs of paternal origin could be transported to the egg cell and potentially regulate gene expression in the zygote at, or shortly after, fertilization. Such phenomena are now proven to be important for embryonic development in mammals (Conine et al., 2018; Sharma et al., 2018; Yuan et al., 2016). Data highlighting such phenomena are rare in *Arabidopsis*. Indeed, most mutants affecting embryo development are rather sporophytic recessive rather than showing a paternal gametophytic effect (Meinke, 2020) which is the case for mutants affecting DCL1 the main micro RNA processing enzyme in *Arabidopsis* (Nodine and Bartel, 2010). However, recently, a microRNA mutant for miR159 was found to have a paternal effect on endosperm development via the downregulation of the MYB33 transcription factor (Zhao et al., 2018b). The high expression of miR159 in sperm cell as well as the very early nature of the observed phenotype suggest that miR159 could be delivered paternally at fertilization.

**AGO EXPRESSION PATTERNS IN THE EARLY SEED**
To gain insights into AGO accumulation during early seed development, we observed fluorescent signals in seeds at one Day-After-Pollination (1 DAP), using confocal microscopy (Fig. 3A-I). At this stage, we could detect expression of eight out of ten Arabidopsis AGOs, while signals from mCherry-AGO2 and mCherry-AGO8 were below detection limit. Clade I mCherry-AGO1 is detected in both inner and outer integument layers in the sporophytic tissue as well as in the embryo, but is below detection in the endosperm (Fig. 3A). Similarly, clade I mCherry-AGO5 is detected in the embryo and inner (but not outer) integument, while it is excluded from the endosperm (Fig. 3B). Despite not being detected in the egg cell, clade I GFP-AGO10 could be detected in the early embryo but not in the endosperm (Fig. 3C). Strong GFP-AGO10 accumulation is also observed in the funiculus’ vasculature and the seed/ovule vascular termination (Fig. S7). As previously described, accumulation of the clade II mCherry-AGO3 is limited to the chalazal seed coat (Fig. 3H; (Jullien et al., 2018)). Clade II GFP-AGO7 is detected in all cell types except the endosperm, and its accumulation is particularly strong in the chalazal seed coat (Fig. 3I).

Among the clade II AGOs involved in RdDM, mCherry-AGO4 shows the strongest accumulation, especially in the sporophytic tissues (Fig. 3D). mCherry-AGO4 can be detected in the early embryo (Fig 3E) but is not detected in the endosperm. The expression pattern of mCherry-AGO6 is similar to that of mCherry-AGO4 (Fig.3F), although the signal is of lower intensity. This result agrees with the known redundancy of AGO4 and AGO6 in mediating DNA methylation and TGS at some genetic loci (Stroud et al., 2012; Zheng et al., 2007). mCherry-AGO9 could be detected in the embryo and also in the endosperm, although not in the integuments (Fig. 3G). In fact, mCherry-AGO9, is visible from the first nuclear division of the endosperm, upon which its signal decreases in intensity but it is still detected at the 4-cells embryo stage (Fig. 3G and Fig. S8B-C). Based on our reporter constructs, AGO9 appears as the only AGO detected during endosperm development.

To conclude, we observed a strong asymmetry of AGOs’ patterns between the endosperm and the embryonic lineages. This difference is supported by LCM transcriptomic data from developing seeds (Fig. S8) and suggests a much less active involvement of RNA silencing pathways in the endosperm compared to the embryo, during early seed development. In Arabidopsis, the RdDM pathway is responsible for Cytosine DNA methylation in the CHH context. It relies on the constant *denovo* DNA methylation by the *de novo* DNA methyltransferases (DRM1 and DRM2) and loading of sRNA by Class II AGOs. In the case of Class II AGOs similarly to what was observed with the *de novo* DNA methyltransferases...
the asymmetry of expression observed between embryo and endosperm lineage could contribute to the high CHH methylation observed during embryo development compared to the endosperm (Bouyer et al., 2017; Gehring et al., 2009; Hsieh et al., 2009; Kawakatsu et al., 2017).

**AGO EXPRESSION PATTERNS IN HEART-STAGE EMBRYO**

In order to investigate AGO accumulation patterns in the differentiated zygote, we dissected heart-stage embryos where most early developmental decision already took place and performed confocal imaging using the fluorescently tagged AGO transgenic lines (Fig. 4A-H). As previously reported (Du et al., 2019; Lynn et al., 1999), clade I mCherry-AGO1 is expressed in all cells of the heart-stage embryo (Fig. 4B). Similarly, we observed a ubiquitous expression of the clade II mCherry-AGO4 and mCherry-AGO6 (Fig. 4E-F). By contrast, clade III mCherry-AGO2, mCherry-AGO3 and clade II mCherry-AGO8 are could not be detected in the heart-stage embryo. Clade I GFP-AGO10 displays its previously reported pattern (Du et al., 2019; Tucker et al., 2008), with fluorescent signals observed in the adaxial part of cotyledons and in the pre-vasculature (Fig 4D).

The three remaining AGOs, AGO5 (clade I), AGO9 (clade III) and AGO7 (clade II) accumulate specifically in the shoot apical meristem region (SAM) of the heart-stage embryo (Fig. 4C, G, H), a pattern already documented for AGO5 using a YFP fluorescent reporter (Tucker et al., 2012). In agreement with our results, AGO5, AGO7 and AGO9 transcript were found to be enriched in meristematic stem cell both at the embryonic and adult stage (Gutzat et al., 2018). Although AGO7 and AGO9 function in meristem remain to be investigated, recent work from Roussin-Leveillee et al suggest that AGO5 function in the meristem is involved in regulating flowering (Roussin-Léveillée et al., 2020). Indeed, ago5 mutant display an early flowering phenotype but do not affect leaf morphology (Fig.S4, (Roussin-Léveillée et al., 2020)). It is thought to do so by interacting with miR156 leading to the repression of SPL transcription factors. However miR156 was shown to affect both flowering and leaf morphology (Wu et al., 2009) which suggest a more complex regulation of this pathway possibility via differential AGO loading and function of miR156 or miR157 isoform (Ebhardt et al., 2010; He et al., 2018) in meristematic region.

**AGO INTRACELLULAR LOCALIZATION PATTERNS**

*Arabidopsis* AGO1 and AGO4 have been shown to shuttle between the cytoplasm and the nucleus although their steady-state subcellular localizations seem to reflect their involvement in either PTGS (clade I and II) and TGS/RdDM (Clade III), respectively: AGO1 is mostly
cytoplasmic and AGO4 nuclear (Bologna et al., 2018; Ye et al., 2012). In agreement with their involvement in Clade I and II, mCherry-AGO1, mCherry-AGO5, mCherry-AGO2, mCherry-AGO3, GFP-AGO10 are mostly localized in the cytoplasm in reproductive cells (Fig. 1B-D, Fig.1I-J, Fig.3A-C, Fig3H, Fig. 4B-D). However, we observed that mCherry-AGO1, mCherry-AGO5, mCherry-AGO2, mCherry-AGO3, GFP-AGO10 are mostly localized in the cytoplasm in reproductive cells (Fig. 1B-D, Fig.1I-J, Fig.1K-L). However, caution should be exerted in interpreting these observations given that artefactual aggregation has been reported with mCherry-tagged proteins (Cranfill et al., 2016; Landgraf et al., 2012). It is likely that these AGO foci occur in tissues where AGO1, AGO5 and AGO2 are particularly highly expressed. Further work is required to assess if the mCherry-AGO1, mCherry-AGO5, and mCherry-AGO2 aggregates represent relevant biological entities.

The main Clade II AGOs, mCherry-AGO4 and mCherry-AGO6, are localized in the nucleus in all reproductive cell types analyzed (Fig. 1E-F, Fig. 2D-E, Fig. 3D-F, Fig. 4E-F). Occasionally, cytoplasmic localization could be observed in the ovule integument or embryo likely due to disruption of the nuclear membrane during cell division. Nuclear localization for AGO4 and AGO6 was previously observed in other tissues (Ye et al., 2012; Zheng et al., 2007). Perhaps more strikingly, mCherry-AGO9 displays nuclear localization in somatic tissues (Fig. 1H, Fig. 3G, Fig. 4G, Fig. S8) but appears to be also partially localized to the cytoplasm in the central and egg cells (Fig. 1H). Combined cytoplasmic and nuclear localizations for AGO9 was previously observed by immunolocalization in ovule primordia (Rodríguez-Leal et al., 2015; Zhao et al., 2018a). However, in those studies, AGO9 was found to accumulate in cytoplasmic foci which we could not observe with our reporter construct in the tissues examined. The mCh-AGO8 translational reporter unlike other RddM AGOs, mainly localize to the cytoplasm of the egg cell (Fig. 1G). As discussed above, this unusual localization for a Clade II AGO could be the result of AGO8 being a truncated protein.

One of the most intriguing intracellular localization patterns was that of GFP-AGO7. GFP-AGO7 is mainly localized to the nucleus. However, in some cells of the integument, in the egg cell and in the central cell, clear cytoplasmic localization is observed, with the presence of cytoplasmic foci (Fig. 1K-L). Localization of GFP-AGO7 to cytoplasmic foci named “sRNA bodies” was previously observed in Nicotiana benthamiana leaves during transient overexpression of GFP-AGO7 (Jouannet et al., 2012). Nuclear GFP-AGO7 accumulation was not reported in this study. The discrepancy could be explained by several factors such as the difference in the promoter used (p35S versus cognate pAGO7 promoters) and also, importantly,
the tissues analyzed (mature leaves or roots versus reproductive tissue). Interestingly, retention of AGO7 in the nucleus using an NLS signal-peptide fused to AGO7 in stable Arabidopsis transformant did not complement the so called leaf zippy phenotype of ago7 mutant (Hunter et al., 2003; Jouannet et al., 2012) showing that cytoplasmic localization is necessary for AGO7 function during leaf development. Similarly to AGO7, it was shown that a nuclear retention of AGO1 do not complement the ago1 mutant phenotype as AGO1 shuttling between the nucleus and the cytoplasm is required for its function in the micro RNA pathway (Bologna et al., 2018). Given that AGO7 zippy mutant phenotype rely on the loading by AGO7 of the micro-RNA miR390 and that our construct also rescues the of ago7 mutant Arabidopsis (Fig. S6) and displays both nuclear and cytoplasmic localization in the inspected tissues, it is most likely that AGO7, similarly to AGO1, shuttles between the nucleus and the cytoplasm and that this process could be regulated intracellularly during development. Additionally, nuclear AGO7 in reproductive tissues could have as yet uncharacterized function as shown recently for nuclear AGO1 (Liu et al., 2018).

To conclude, we have generated full locus fluorescently N-term tagged constructs of all Arabidopsis AGOs under their native endogenous promoter. We have analyzed their expression pattern as well as intracellular localization in reproductive tissues. These constructs or transgenic lines can be used by the community to study AGOs regulation in a developmental context as well as during stresses. Fluorescent AGOs reporter were previously reported in Arabidopsis however inconsistencies are present such as the use of overexpression promoters or C-terminal fusion known to affect the functionality of AGOs protein. We hope these lines will be beneficial to the community and provide a uniform set of tools for further analysis.

**MATERIALS AND METHODS**

**PLANT MATERIAL AND GROWTH CONDITIONS**

After three days at 4°C in the dark, seeds were germinated and grown on soil. Plants were grown under long days at 20-21°C (16h light/8h night). All plants were in Columbia (Col-0) accession. The mutants and lines described in this work correspond to the following: ago1-36 (SALK_087076,(Baumberger and Baulcombe, 2005)), ago2-1 (salk_003380,(Lobbes et al., 2006)), ago3-3 (GABI-743B03,(Jullien et al., 2018)), ago4-5 (WiscDsLox338A0, (Stroud et al., 2012)) , ago5-1 (salk_063806, (Katiyar-Agarwal et al., 2007)), ago6-2 (salk_031553,
(Zheng et al., 2007), ago7-1 (salk_037458, (Vazquez et al., 2004)), ago8-1 (salk_139894, (Takeda et al., 2008), ago9-1 (salk_127358, (Katiyar-Agarwal et al., 2007)) and ago10-1 (SALK_000457). The insertion lines were provided by The Nottingham Arabidopsis Stock Centre (NASC) (http://arabidopsis.info/). Pollen were germinated in Pollen growth medium at 21°C in the dark 5 hours to over-night (Hamamura et al., 2011).

**Microscopy**

Fluorescence images were acquired using laser scanning confocal microscopy (Zeiss LSM780 or Leica SP5). Brightness was adjusted using ImageJ (http://rsbweb.nih.gov/ij/) and assembled using ImageJ or Adobe Illustrator.

**Plasmid Construction and Transformation**

All DNA fragments were amplified by PCR using the Phusion High-Fidelity DNA Polymerase (Thermo). Primer sequences can be found in Supplementary Table S1. All plasmids were transformed into their respective mutants and for some also in LIG1-GFP marker line (Andreuzza et al., 2010). All constructs were generated using Multisite Gateway technology (Invitrogen). *A. thaliana* transformation was carried out by the floral dip method (Clough and Bent, 1998). At least six to ten transgenic lines (T1) were analysed and showed a consistent fluorescence expression pattern using a Leica epifluorescence microscope or a Leica SP5. One to three independent lines with single insertions, determined by segregation upon BASTA selection, were used for further detailed confocal analysis.

**Author Contributions**

PEJ conceived the study. PEJ, NP and JAS generated the transgenic lines. PEJ and DMVB performed the imaging. DMVB and JAS tested the complementation. PEJ wrote the manuscript, which was further edited and amended by OV.

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COMPETING INTERESTS
The authors have no conflicts of interest to declare.

SUPPLEMENTARY INFORMATION
Additional Supporting Information may be found in the online version of this article.


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Figure Legends

Fig. 1. AGOs accumulation in the mature female gametophyte. (A) schematic representation of a mature female gametophyte of *Arabidopsis thaliana* illustrating the three major cell types: the central cell (cc) in green, the egg cell (ec) in orange, the synergides (syn) in blue. The mature female gametophyte is surrounded by maternal sporophytic tissue represented in grey, including the inner integument (ii), outer integument (oi) and the nucellus (nuc). (B-K) Confocal images representing the expression of the 10 *Arabidopsis* AGOs in mature female gametophytes: mCherry-AGO1 (B), mCherry-AGO5 (C), GFP-AGO10 (D), mCherry-AGO4 (E), mCherry-AGO6 (F), mCherry-AGO8 (G), mCherry-AGO9 (H), mCherry-AGO2 (I), mCherry-AGO3 (J) and GFP-AGO7 (K). (L) Confocal image of the intra-cellular localization of GFP-AGO7 in the egg cell and central cell. Scale bars represent 10 µm.

Fig. 2. AGOs accumulation in the pollen. (A-H) AGOs in the mature male gametophyte. (A) Schematic representation of a mature pollen grain of *Arabidopsis thaliana* illustrating the two major cell types: the vegetative cell (v) and the two sperm cells (s). (B-H) Confocal images representing the expression of seven *Arabidopsis* AGOs accumulating in the pollen grain: mCherry-AGO1 (B), mCherry-AGO5 (C), mCherry-AGO4 (D), mCherry-AGO6 (E), mCherry-AGO9 (F), mCherry-AGO2 (G) and GFP-AGO7 (H). (I-P) AGO accumulation in germinating pollen tube. (I) Schematic representation of a growing pollen tube of *Arabidopsis thaliana* illustrating the two sperm cells (s). (J-P) Confocal images of the seven *Arabidopsis* AGOs in germinated pollen grain: mCherry-AGO1 (B), mCherry-AGO5 (C), mCherry-AGO4 (D), mCherry-AGO6 (E), mCherry-AGO9 (F), mCherry-AGO2 (G) and GFP-AGO7 (H). Scale bars represent 5 µm.

Fig. 3. AGOs accumulation in the developing seed. (A-I) Confocal images of the height *Arabidopsis* AGOs expressed in the developing seeds 2 days-after-pollination (DAP): mCherry-AGO1 (A), mCherry-AGO5 (B), GFP-AGO10 (C), mCherry-AGO4 (D-E), mCherry-AGO6 (F), mCherry-AGO9 (G), mCherry-AGO3 (H) and GFP-AGO7 (I). Scale bars represent 10 µm. Abbreviation: emb (embryo), endo (endosperm), inner integument (ii), outer integuments (oi), Chalazal seed coat (czsc).
**Fig. 4.** AGOs accumulation in the heart-stage embryo. (A) A simplified representation of a heart-stage embryo of *Arabidopsis thaliana* illustrating the different cell types. (B-H) Confocal images of the seven Arabidopsis AGOs accumulating in the heart-stage embryo: mCherry-AGO1 (B), mCherry-AGO5 (C), GFP-AGO10 (D), mCherry-AGO4 (E), mCherry-AGO6 (F), mCherry-AGO9 (G), and GFP-AGO7 (H). Scale bars represent 10 µm.

**Fig. S1.** Schematic representation of the constructs used in this study. The different features are represented by arrows: promoter (purple), UTRs (yellow), fluorescent protein (red), exon (green) and additional annotations (blue).

**Fig. S2.** Complementation of the *ago1-36* mutant *Arabidopsis* by *pAGO1:mCherry-AGO1*. Representative pictures showing the rescue of the string developmental phenotype in Col-0 (A) and mCh1 #9 *ago1-36/-* (C) compared to *ago1-36/-*. scale bar represents 1mm.

**Fig. S3.** Complementation of *ago4-5* mutant by *pAGO4:mCherry-AGO4*. qPCR showing the absence of *AtSN1* ectopic expression in rosette leaves of seven independent lines expressing the construct *pAGO4:mCherry-AGO4* (*mCh4*) in *ago4-5* background compared to Col-0 and *ago4-5*. Actin2 was used as endogenous control. p indicates the p value obtained by a Student’s T-Test.

**Fig. S4.** Complementation of *ago5-1* mutant *Arabidopsis* by *pAGO5:mCherry-AGO5*. (A) Representative pictures of the early flowering phenotype of *ago5-1/-* compared to Col-0 and mCh5 #29 *ago5-1/-*. (B) Quantification of *ago5-1/-* flowering phenotype in *ago5-1/-* (n=27), Col-0 (n=26) and mCh5#29 *ago5-1/-* (n=27).

**Fig. S5.** Complementation of *ago6-2* mutant by *pAGO6:mCherry-AGO6*. qPCR showing the absence of *AtSN1* ectopic expression in rosette leaves of seven independent lines expressing the construct *pAGO6:mCherry-AGO6* (*mCh4*) in *ago6-2* background compared to Col-0 and
ago6-2. Actin2 was used as endogenous control. p indicates the p value obtained by a Student’s T-Test.

Fig. S6. Complementation of ago7-1 mutant Arabidopsis by pAGO7:GFP-AGO7. Illustrative pictures of the leaf “zippy” phenotype of ago7-1/− (B) compared to Col-0 (A) and GFP-AGO7 ago7-1/− (C). n= 7

Fig. S7. Additional pictures of pAGO10:GFP-AGO10. (A-B) Picture of the accumulation in ovules of GFP-AGO10 in the vascular tissue of the funiculus (funi) and at the vascular termination of the chalazal seed coat (czsc). Scale bars represent 10 μm.

Fig. S8. Additional pictures of pAGO9:mCh-AGO9. (A) Pictures of the expression in ovules of mCherry-AGO9 in the funiculus (funi) and in the chalazal seed coat (czsc). (B-C) Accumulation of mCherry-AGO9 in developing seeds, in early embryo and endosperm, 17 hours after pollination (HAP) (B) or 24 HAP (C). Scale bars represent 10 μm.

Fig. S9. Arabidopsis AGO transcription patterns extracted from microarray data of LCM-dissected female gametophytes (Wuest et al, 2010) confirming the general enrichment of AGO transcripts in the egg cell (EC) compared to central cell (CC) or synergids (Syn). (A) Violin plot representing the general enrichment of AGO transcripts in the EC. (B) AGOs individual expression boxplot in the different cell types. p values of a Wilcoxon test are indicated.

Fig. S10. Arabidopsis AGO transcription patterns extracted from microarray data of LCM-dissected seeds at the pre-globular stage (Belmonte et al., 2013) confirming the general enrichment of AGO transcripts in the embryo compared to the peripheral endosperm. (A) Violin plot representing the general enrichment of AGO transcripts in the embryo. (B) AGOs individual expression boxplot in the different cell types. p values of a Wilcoxon test are indicated. AGO6 and AGO8 probes are not present in these data.
Fig S11. Summary table of AGO's expression pattern in reproductive tissues.
Fig. 1. Argonaute expression in the mature female gametophyte. (A) schematic representation of a mature female gametophyte of Arabidopsis thaliana illustrating the three major cell type: the central cell (cc) in green, the egg cell (ec) in orange, the synergides (syn) in blue. The mature female gametophyte is surrounded by maternal sporophytic tissue represented in grey including the inner integument (ii), outer integument (oi) and the nucellus (nuc). (B-K) Confocal images representing the expression of the 10 Arabidopsis AGOs in mature female gametophyte: mCherry-AGO1 (B), mCherry-AGO5 (C), GFP-AGO10 (D), mCherry-AGO4 (E), mCherry-AGO6 (F), mCherry-AGO8 (G), mCherry-AGO9 (H), mCherry-AGO2 (I), mCherry-AGO3 (J) and GFP-AGO7 (K). (L) Confocal image illustrating the intra-cellular localization of GFP-AGO7 in the egg cell and central cell. Scale bars represent 10μm.
Fig. 2. Argonaute expression in pollen. (A-H) AGOs in the mature male gametophyte. (A) Schematic representation of a mature pollen grain of Arabidopsis thaliana illustrating the two major cell type: the vegetative cell (v) and the two sperm cells (s). (B-H) Confocal images representing the expression of 7 Arabidopsis AGOs expressed in the pollen grain: mCherry-AGO1 (B), mCherry-AGO5 (C), mCherry-AGO4 (D), mCherry-AGO6 (E), mCherry-AGO9 (F), mCherry-AGO2 (G) and GFP-AGO7 (H). (I-P) Argonaute expression in germinating pollen tube. (I) Schematic representation of a growing pollen tube of Arabidopsis thaliana illustrating the two sperm cells (s). (J-P) Confocal images representing the 7 Arabidopsis AGOs expressed in in germinated pollen grain: mCherry-AGO1 (B), mCherry-AGO5 (C), mCherry-AGO4 (D), mCherry-AGO6 (E), mCherry-AGO9 (F), mCherry-AGO2 (G) and GFP-AGO7 (H). Scale bars represent 5 μm.
Fig. 3. Argonaute expression in developing seed. (A-I) Confocal images representing Arabidopsis AGOs expressed in the developing seeds at 2 day after pollination: mCherry-AGO1 (A), mCherry-AGO5 (B), GFP-AGO10 (C), mCherry-AGO4 (D-E), mCherry-AGO6 (F), mCherry-AGO9 (G), mCherry-AGO3 (H) and GFP-AGO7 (I). Scale bars represent 10 μm. Abbreviation: emb (embryo), endo (endosperm), inner integument (ii), outer integuments (oi), Chalazal seed coat (czsc).
Fig. 4. Argonaute expression in heart stage embryo. (A) A simplified representation of a heart stage embryo illustrating the different cell types. (B-H) Confocal images representing the 7 Arabidopsis AGOs expressed in heart stage embryo: mCherry-AGO1 (B), mCherry-AGO5 (C), GFP-AGO10 (D), mCherry-AGO4 (E), mCherry-AGO6 (F), mCherry-AGO9 (G), and GFP-AGO7 (H). Scale bars represent 10 μm.
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Fig. S3. Complementation of ago4-5 mutant by pAGO4:mCherry-AGO4. qPCR showing the absence of AtSN1 ectopic expression in rosette leaves of seven independent lines expressing the construct pAGO4:mCherry-AGO4 (mCh4) in ago4-5 background compared to Col-0 and ago4-5. Actin2 was used as endogenous control. p indicates the p value obtained by a Student’s T-Test.
Fig. S4. Complementation of ago5-1 mutant by pAGO5:mCherry-AGO5. (A) Illustrative pictures of the early flowering phenotype of ago5-1/- compared to Col-0 and mCh5 # 29 ago5-1/-. (B) Quantification of ago5-1/- flowering phenotype in ago5-1/- (n=27), Col-0 (n=26) and mCh5#29 ago5-1/- (n=27).
Fig. S5. Complementation of ago6-2 mutant by pAGO6:mCherry-AGO6. qPCR showing the absence of AtSN1 ectopic expression in rosette leaves of seven independent lines expressing the construct pAGO6:mCherry-AGO6 (mCh4) in ago6-2 background compared to Col-0 and ago6-2. Actin2 was used as endogenous control. p indicates the p value obtained by a Student’s T-Test.
Fig. S6. Complementation of ago7-1 mutant by pAGO7:GFP-AGO7. Illustrative pictures of the leaf phenotype of ago7-1/- (B) compared to Col-0 (A) and GFP-AGO7 ago7-1/- (C).
Fig. S7. Additional pictures of pAGO10:GFP-AGO10. (A-B) Picture representing the expression in ovules of GFP-AGO10 in the vascular tissue of the funiculus (funi) and at the vascular termination in the chalazal seed coat (czsc). Scale bars represent 10μm.
Fig. S8. Additional pictures of pAGO9:mCh-AGO9. (A) Pictures representing the expression in ovules of mCherry-AGO9 in the funiculus (funi) and in the chalazal seed coat (czsc). (B-C) Pictures representing the expression in developing seeds of mCh9 in early embryo and endosperm at 17 hours after pollination (HAP) (B) and at 24 HAP (C). Scale bars represent 10 μm.
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