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Genetic variation and temperature affects hybrid barriers during interspecific hybridization

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Abstract

Genomic imprinting regulates parent-specific transcript dosage during seed development and is mainly confined to the endosperm. Elucidation of the function of many imprinted genes has been hampered by the lack of corresponding mutant phenotypes, and the role of imprinting is mainly associated with genome dosage regulation or allocation of resources. Disruption of imprinted genes has also been suggested to mediate endosperm based post-zygotic hybrid barriers depending on genetic variation and gene dosage. Here, we have analyzed the conservation of a clade from the MADS-box type I class transcription factors in the closely related species *Arabidopsis arenosa*, *A. lyrata* and *A. thaliana*, and show that *AGL36-like* genes are imprinted and maternally expressed in seeds of *Arabidopsis* species and in hybrid seeds between outbreeding species. In hybridizations between outbreeding and inbreeding species the paternally silenced allele of the *AGL36-like* gene is

This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as reactivated in the hybrid, demonstrating that also maternally expressed imprinted genes are perturbed during hybridization and that such effects on imprinted genes are specific to the species combination. Furthermore, we also demonstrate a quantitative effect of genetic diversity and temperature on the strength of the post-zygotic hybridization barrier. Remarkably, a small decrease in temperature during seed development increases survival of hybrid F1 seeds, suggesting that abiotic and genetic parameters play important roles in post-zygotic species barriers, pointing at evolutionary scenarios favoring such effects.

Introduction

Seed development is a sophisticated and highly regulated process that requires precise signaling events and interaction between many distinct cell types and tissues. It starts with fusion of the male and female gametes generated in the male and female gametophytes, giving rise to the embryo and endosperm that develop in parallel inside the protective seed coat. The process is initiated when a conspecific pollen grain lands on the stigma of the female reproductive organ and the pollen tube delivers two sperm cells to the female gametophyte. One sperm cell fertilizes the haploid egg cell which develops into the diploid embryo, while the other sperm cell fertilizes the homodiploid central cell generating the triploid endosperm. The endosperm is important for nutrient flow to the embryo but also for coordinating growth of the developing seed (Nowack et al. 2010).

The endosperm has two maternal genome copies and one paternal copy, and a specialized epigenetic phenomenon called genomic imprinting regulates parent-specific gene dosage during seed development, usually occurring in the endosperm (Gehring and Satyaki 2017). Imprinting is manifested by expression of one parental allele, with concurrent silencing of the other allele. The main mechanisms for this process are DNA methylation and histone methylation (Berger et al. 2006). The FERTILIZATION INDEPENDENT SEED-Polycomb Repressive Complex 2 (FIS-PRC2) mediates histone methylation while, in *A. thaliana*, DNA methylation mediated imprinting is maintained by the DNA methyltransferase MET1 (Rodrigues et al. 2015).

A prominent gene family displaying frequent imprinting of its members is the MADS-box transcription factor (TF) family. The MADS-box TFs can be divided into type I and type II by evolutionary relationships. The type I TFs are further divided into M α , M β and M γ phylogenetic subclasses and only share the highly conserved DNA binding MADS (M) domain. The type II TFs have in addition to the M domain, the Intervening (I), the Keratin (K) and the C-terminal (C) domain

and are often referred to as the MIKC-type (Parenicova et al. 2003). The type II class is thought to have evolved from an ancient whole genome duplication as orthologs are found in many other species and the genes are well distributed across all chromosomes in *A. thaliana*. The type I class TFs originate from more recent and smaller scale duplication events and in *A. thaliana* they are mainly concentrated on chromosomes I and V (Parenicova et al. 2003; Airoldi and Davies 2012). As a consequence, MADS-box type I orthologs are uncommon in other species (Masiero et al. 2011). Imprinted genes occur frequently in the type I class, consistent with the hypothesis that recently duplicated genes are more often imprinted to regulate gene dosage (Yoshida and Kawabe 2013). Imprinting is observed mainly in the M α and M γ subgroups and moreover, members of these two subclasses interact extensively in yeast two-hybrid assays, suggesting a common function as heterodimers (de Folter et al. 2005).

Functional studies of the MADS-box type I TFs by genetic dissection, however, are hampered by genetic redundancy. Their roles have also been suggested to have restricted effect and may therefore be involved in specific developmental processes (Nam et al. 2004). Only a few type I genes have been studied phenotypically, including *AGAMOUS LIKE (AGL) 23*, *AGL36*, *PHERES (PHE) 1 (AGL37)* and *PHE2 (AGL38)*, *DIANA (AGL61)*, *AGL62*, and *AGL80* (Bemer et al. 2008; Kang et al. 2008; Steffen et al. 2003; Shirzadi et al. 2011; Colombo et al. 2008; Kohler et al. 2005). To this end, the biological roles of many imprinted genes are still not known, but the role of imprinting is mainly associated with genome dosage regulation or allocation of resources (Dilkes and Comai 2004; Haig and Westoby 1989; Rodrigues et al. 2015).

Imprinting has previously been shown to be disrupted in hybrid crosses of *A. thaliana* and *A. arenosa*. The MADS-box TF *PHE1*, which is imprinted and only paternally expressed in *A. thaliana*, was upregulated in hybrid seeds and it was shown that the expressed *PHE1* was predominantly maternally expressed (Josefsson et al. 2006). Disruption of the expression levels of co-adapted MADS-box TFs in hybrids may thus trigger genome-wide perturbations observed in hybrids (Roth et al. 2019). Furthermore, other MADS-box type I TFs have been shown to be highly upregulated in incompatible hybrid crosses between *A. thaliana* mothers and *A. arenosa* fathers. Using knock-out mutant lines of these genes as *A. thaliana* mother, increased viability in the incompatible hybrid seeds, suggesting that these MADS-box type I TFs partly constitute a genetic basis for the post-zygotic barrier (Walia et al. 2009). Hence, investigation of the imprinting status of these genes and other known imprinted genes in *A. arenosa* and *A. lyrata* will shed light on the role and consequently the evolution of imprinting. It is disputed whether imprinting of specific genes is conserved, and whether the mechanisms behind the establishment and maintenance of imprinting between related and distant

species are preserved (Waters et al. 2013; Klosinska et al. 2016; Hatorangan et al. 2016; Chen et al. 2018).

Diploid *A. arenosa* crossed as father to more than 50 accessions of *A. thaliana* displayed live seeds in the range of 1% to 30% (Burkart-Waco et al. 2012). This suggests that the strength of the post-zygotic barrier can be modulated by genetic variation in accessions. Comparison of *A. arenosa* crossed to different *A. thaliana* accessions, thorough phenotyping (Burkart-Waco et al. 2013) and sequencing of RNA from hybrid seeds (Burkart-Waco et al. 2015), identified perturbation of the imprinting patterns of eight known paternally expressed genes. As these crosses were limited to a specific *A. arenosa* population, we hypothesize that the observed barrier is population dependent. Lafon-Placette et al. (2017) demonstrated that in crosses between *A. lyrata* and *A. arenosa*, the post-zygotic species barrier is is due to endosperm cellularization failure. A similar study in the *Capsella* genus also indicated endosperm failure as the main seed defect in incompatible crosses (Rebernig et al. 2015). A post-zygotic endosperm-based barrier has also been described for rice (Wang et al. 2018; Tonosaki et al. 2018) and tomato (Florez-Rueda et al. 2016).

Here we have investigate the role of genetic variation in the establishment of post-zygotic endospermbased hybrid barriers both in general, using accession, and in a targeted manner, addressing specific MADS-box type I loci. We investigated the function and regulation of a conserved clade of MADSbox type I Mγ class (*AGL34*, *AGL36* and *AGL90*) together with some of their interacting partners. To further elucidate function, we have analyzed the conservation of this clade in the closely related species *A. arenosa*, *A. lyrata* and *A. halleri*, including the imprinting status of *AGL36-like* genes in *A. arenosa*, in *A. thaliana* crossed to *A. arenosa* and in the reciprocal cross of *A. arenosa* and *A. lyrata*. We find that *AGL36-like* genes are imprinted and maternally expressed in seeds of *Arabidopsis* species and in hybrid seeds between outbreeding species. In hybridizations between outbreeding and inbreeding species the paternally silenced allele of the *AGL36-like* gene is reactivated in the hybrid, demonstrating that also maternally expressed imprinted genes are perturbed during hybridization and that such effects on imprinted genes is specific to the species combination.

Moreover, we investigated the role of temperature in hybridization of different genetic backgrounds and specific loci and find a significant positive correlation between lower temperatures and hybrid seed germination rate. We report that just a small change in temperature during seed development is sufficient to increase survival of hybrid F1 seeds, suggesting that abiotic parameters play an important role in post-zygotic, endosperm based species barriers. Crossing mutants of the M γ and M α clades, and their interacting partners to *A. arenosa* to further investigate the effect of these genes on the hybridization barrier identified that lack of *AGL35* significantly aggravate the *A. thaliana A. arenosa* hybrid barrier and that *AGL35* is involved in the temperature dependency of the hybrid barrier.

Results

MADS-box type I Ma and My expression in seed development

In order to investigate the role of imprinted loci in the establishment of endosperm based hybrid barriers, we analysed MADS-box TFs that are closely related to the paternally silenced AGL36 (Shirzadi et al. 2011). We re-analyzed the phylogeny of the M α and M γ classes and assembled them in two groups with several sub-clades (Figure 1A). In the M γ group, AGL36 constitutes a sub-clade together with AGL90 and AGL34, and the latter genes may represent recent, local gene duplication events in A. thaliana since orthologs are not readily identified (Masiero et al. 2011). AGL36 and AGL90 have been shown to be imprinted (Shirzadi et al. 2012, Zhang et al. 2018). The wider sub-clade includes known imprinted, paternally-expressed genes PHE1/AGL37 and AGL92 (Wolff et al. 2011), which cluster together with PHE2/AGL38 and AGL86, respectively. PHE2 expression has previously been demonstrated to be bi-allelic (Villar et al. 2009). The most distant member of the sub-clade is AGL35, which is closely linked to and located between AGL34 and AGL36 on chromosome 5. AGL34, AGL35, AGL36 and AGL90 all map in a 100 Kbp cluster on chromosome 5 (Parenicova et al. 2003), and this clustering makes this sub-clade an especially interesting case to study evolution of imprinted genes. The function of AGL90 and AGL34 is not known, but AGL36 interacts with two Ma MADSbox TFs, AGL28 and AGL62 (Bemer et al. 2010; de Folter et al. 2005) where AGL28 has been shown not to be imprinted (Zhang et al. 2018) or to display accession dependent imprinting (Wolff et al. 2011). AGL62 is biparentally expressed and required for endosperm cellularization (Kang et al. 2008), and for reason of functional study we have included AGL62 and AGL28 in our analysis.

Next, we investigated relative expression of all MADS-box type I TFs at seed developmental stages ranging from one day after pollination (DAP) to 12 DAP (Figure 1B, Figure S1). *AGL36* expression peaked at 4 DAP and coincided with the timing of endosperm cellularization (Shirzadi et al. 2011), hence an RNA-sequencing based differential expression analysis relating all stages to 4 DAP was performed. M α and M γ class TFs are overrepresented in the transcriptome of the developing seed compared to the M β class. All 16 M γ genes and two-thirds of the 25 M α genes are expressed, whereas less than half of the 21 M β genes can be identified (Figure S1B). Ordering the M α and M γ expression profiles according to the branching pattern, displayed a general expression trend with a peak between

4 and 6 DAP (Figure 1B). In the M γ *AGL36* sub-clade, *AGL35*, *AGL36* and *AGL90* display similar profiles with increasing or unchanged expression towards 4 DAP followed by a decline. *AGL34* can only be detected in a small developmental window, but the relative expression pattern is equivalent to *AGL36* and *AGL90* at these stages (Figure 1B, Figure S1B). This supports findings by Zhang and colleagues (2018) and indicates that *AGL34* is not a pseudogene, as previously postulated (Bemer et al. 2010). A similar pattern is found in M α subclades, including *AGL28*. A decrease of *AGL62* levels was observed after the expression maxima observed in M α and M γ classes (Figure 1B). The M α *AGL62* is required for correct timing of endosperm cellularization (Kang et al. 2008), and thus play a putative role in the establishment of endosperm-based hybrid barriers (Lafon-Placette et al. 2017). The M α MADS-box TF subclass are hypothesized to form dimers with the M γ -type (de Folter et al. 2005), and taken together, the co-occurring M α and M γ expression patterns may indicate a possible role for these TFs in the establishment of cellularization-based post-zygotic hybrid barriers.

Imprinting and regulation of My and Ma MADS-box genes

Dosage imbalance caused by imprinted genes has been proposed as a cause for hybrid failure in both plants and animals (Dilkes and Comai 2004; Wolf et al 2014; Brekke et al. 2016). To this end, we successively re-analyzed parent-of-origin expression of the wider AGL36 sub-clade (Figure 1A), including the interacting M α AGL28 (Figure 2). Using Col-0 and Tsu-1 accession specific SNPs, we analysed AGL28, AGL35, AGL36 and AGL90 in 4 DAP Col-0 Tsu-1 hybrid seeds (Figure 2A). Maternal bias from the seed-coat could be excluded, since all transcripts were previously shown to be enriched >8-fold 4DAP in the peripheral endosperm (AGL36, AGL90) compared to all other seed tissues or >8-fold and >5-fold enriched in the chalazal endosperm (AGL35 and AGL28, respectively) (Belmonte at al. 2013; Hornslien at al. 2019). AGL34 was not expressed at a sufficient level in 4 DAP Tsu-1 and was thus omitted. Gene specific RT-PCR products from hybrid crosses were digested with SNP-specific restriction endonucleases (Table S1) and fragments analyzed on a Bioanalyzer 2000 as well as by Sanger sequencing (Figure 2A, Figure S2). The My genes AGL36 and AGL90 and the M α AGL28 were imprinted and maternally expressed. The My AGL35 was biparentally expressed as previously reported (Zhang et al. 2018). AGL36 (Shirzadi et al. 2011; Wolff et al. 2011; Zhang et al. 2018) and AGL90 (Zhang et al. 2018) has previously been shown to be imprinted. Here, we show that AGL90 is maternally biased in its expression but the paternal allele show accession dependent imprinting and is not completely silenced from Col-0 pollen donors (Figure 2A, Figure S2). A similar lack of silencing of the AGL90 paternal Col-0 allele was recently also reported (Hornslien at al. 2019). In contrast, AGL28 was previously reported not to be imprinted (Zhang et al. 2018), or to display accession dependent imprinting in hybrids (Wolff et al. 2011).

To address the regulation of imprinted genes we contrasted 4 DAP parental seed expression from crosses between Tsu-1 and Col-0 accessions versus Tsu-1 and a hemizygous *met1-7* +/- in a Col-0 background to determine if MET1 is involved in maintaining silencing of the the paternal copy of *AGL28*, *AGL36* and *AGL90* (Figure 2B, Table S1, Figure S2). The paternal copy of *AGL28* was shown to be expressed using *met1-7*^{+/-} as pollen donor, suggesting that MET1 is required for silencing of the paternal *AGL28* allele. In our experimental settings, however, lack of *MET1* did not reactivate the paternal copy of *AGL36* and *AGL90*, whereas the *AGL35* biparental control remains unchanged (Figure 2B). In contrast to these findings, the paternal allele of *AGL36* was previously shown to be reactivated in crosses with homozygous and hemizygous *MET1* mutant pollen using the *met1-4* +/- allele (Shirzadi et al. 2011; Saze et al. 2003). Lack of *MET1*, both in homozygous and heterozygous mutants leads to DNA hypomethylation and eventually the accumulation of epimutations, and we attribute the previously observed paternal expression to such effects in the *met1-4* +/- background.. The *met1-7* +/- allele used in our study was kept hemizygous through repeated outcrosses and therefore more likely devoid of such effects (Hornslien et al. 2019).

According to a report investigating the role of small interfering (si) RNA and RdDM in interploidy crosses, several MADS-box type I genes, including AGL36, were deregulated in diploid crosses with NUCLEAR RNA POLYMERASE D1 (NRPD1) mutant mothers, deficient in the largest subunit of RNA polymerase IV, a key components of canonical RdDM (Lu et al. 2012). Using nuclear rna *polymerase d1 (nrpd1)* mutant mothers, AGL36 was upregulated more than 20-fold (Lu et al. 2012). We have previously shown that imprinting of AGL36 do not require paternal DOMAINS REARRANGED METHYLTRANSFERASE 2 (DRM2) or ARGONAUTE4 (AGO4), both part of the RdDM pathway (Shirzadi et al. 2011). The data from Lu and colleagues (2012) suggested that an RdDM dependent mechanism maintain the expression level of AGL36 or is active in maintaining the silencing of the paternal AGL36 allele after fertilization To test the latter hypothesis, we analysed parental expression from 6 DAP seeds, using *nrpd1* both as a maternal and paternal contributor in crosses to wild type (Figure 2C, Figure S2). The AGL36 imprinting pattern was not affected in any cross direction, suggesting that reactivation of the paternal allele is not causing elevated levels of AGL36. In contrast to the previous report (Lu et al. 2012), we could also not detect any significant upregulation of AGL36 by real-time PCR (Figure 2D) in crosses using a homozygous nrpd1 knockout allele (Figure 2E) as maternal cross partner. We conclude that neither the MET1 nor the PolIV RdDM pathway is sufficient to silence the paternal allele of AGL36.

Next, we analysed the effect of PRC2 on MADS-box type I genes. The endosperm cellularization defect observed in *Arabidopsis* interspecies hybrid seeds is highly reminiscent to the failure of endosperm cellularization phenotype observed in mutants of *FIS-PRC2* (Lafon-Placette et al. 2017).

We therefore compared the RNAseq relative expression of all MADS-box type I genes between a *FIS-PRC2* mutant and wild type at seed developmental stages ranging from one DAP to 12 DAP (Figure 3A, Figure S3). Clustering of transcript profiles revealed four main patterns of regulation, ranging from highly regulated to no effect (Figure S3). Overall, the M β class as a group was significantly less regulated by the PRC2 *medea* (*mea*) mutation than the M α and M γ classes, and also displayed the least variation (Figure 3B). This is in accordance with previous observations, that the M β class TF are mainly expressed at low levels or in female gametophytic stages (Bemer et al. 2010).

In the deregulated classes of transcript profiles, consisting of mainly but not exclusively M α and M γ , the wild type profiles are generally characterized by increasing expression that decreases after a peak (Figure S3, left panels). In the *mea* cross, both this pattern and the peak are shifted towards higher expression levels and later developmental stages. In certain cases, decrease is not observed within the analyzed developmental time-frame (Figure S3, right panels). A distinct shift in transcript profiles could also be observed between the profile clusters in *mea*, with one class de-repressed before 6 DAP, while a second class started at 9 DAP and the third class at 12 DAP (Figure 3A, three top clusters). My dominates the two former classes of transcript profiles together with M α whereas the latter constitutes of M α and M β genes. In a recent report, Zhang and colleagues (2018) analyzed MADSbox type I deregulation in a *swinger (swi) mea* double mutant. These authors identified two major expression clusters (C1 and C2) based on difference in temporal expression patterns both in the wild type and in the PRC2 double mutant. The latter cluster was distinguished by the an up-regulation of the expression pattern in the mutant and could be further divided in two clusters (C2.1 and C2.2) based on the timing of downregulation in the wild type (Zhang et al. 2018). The three clusters described in our study (Figure 3A, three top clusters) are well in line with the realtime-PCR based study of Zhang and colleagues (2018). Eight out of 12 genes in the C2.1 cluster are also found in our top cluster, starting deregulation at the earliest stage (Figure 3A, top cluster), whereas three genes are fund in our second cluster (Figure 3A, second top cluster) together with all mea swi up-regulated genes identified in the C2.2 cluster. This also includes AGL91, AGL49 and importantly AGL34 that are upregulated in our study, whereas no upregulation was identified by Zhang et al. In contrast, we could not detect any up-regulation for AGL64, as reported by the other study (Zhang et al. 2018).

We conclude that the *AGL36* sub-clade, including *AGL34* and *AGL90*, as well as the AGL36 interacting M α AGL28 are commonly repressed by MEA from 4-6 DAP. The AGL36 and also the AGL90 interacting M α AGL62 are upregulated in *mea* at 9 DAP, in accordance with the role of AGL62 in in endosperm cellularization (Kang et al. 2008).

Conservation of Arabidopsis AGL36-like imprinting in hybrid seeds is species dependent

Having analyzed the expression and regulation of MADS-box type I genes i A. thaliana, we turned our focus to the expression and role of these genes in the genus Arabidopsis. MADS-box type I genes are often less conserved between model species. For instance, no orthologs of AGL36 were identified in rice or maize (Masiero et al. 2011). AGL36-like genes can be found when analyzing more closely related species such as in the genus Arabidopsis (Figure 4A, Figure S4). Two genomic loci of AGL36*like* genes were identified in *A. arenosa* by assembling online resources (see Materials and methods). Both were verified in various individuals from two A. arenosa populations using PCR amplification (MJ09-1, MJ09-4 (Jorgensen et al. 2011; Lafon-Placette et al. 2017)). The two genes differ in length (1050 bp versus 1008 bp). The shorter does not have a continuous open reading frame and most likely harbor an intron based on two open reading frames spaced by an 88 bp sequence. The 1050 bp locus, but not the 1008 bp AGL36 locus, was confirmed to produce a transcript in 9 DAP seeds, corresponding to the globular-embryo seed stage. Online genome sequencing resources of A. lyrata subsp. lyrata suggest one AGL36-like gene (Figure 4A). In the subspecies A. lyrata subsp. petrea, two loci have been indicated (Yoshida and Kawabe 2013), but by performing Sanger sequencing from the A. lyrata subsp. petrea population MJ09-11 (Jorgensen et al. 2011) combined with online resources, we concluded that A. lyrata contains only one AGL36-like locus (Figure 4A, Figure S4). Notably, although two AGL36-like loci are present in A. arenosa and A. halleri, our analysis indicates that the two duplication events creating the AGL36 subclade (AGL34, AGL36, AGL90) do not exist outside A. thaliana.

To analyze imprinting of *A. arenosa AGL36-like (AaAGL36-like)*, we screened natural populations (MJ09-4 and MJ09-1) for SNPs that could be used to distinguish the parental alleles. We identified one individual that had a SNP in *AaAGL36-like* (I, cf. Figure 4A) that also allowed SNP detection with restriction enzymes (Table S1C). Seed RNA was harvested from reciprocal crosses at 9 DAP followed by RT-PCR of the SNP containing regions from *AaAGL36-like* (Table S2). The PCR products were digested with SNP specific enzymes (Table S1C) and fragments analyzed (Figure 4B). Only maternal expression was found, suggesting that *A. arenosa AGL36-like (AaAGL36 I)* is an imprinted maternally expressed gene.

Next, we analyzed *AGL36-like* imprinting in hybrids of *A. arenosa* and *A. lyrata*. Amplifying *AGL36-like* (Table S1C) from cDNA of reciprocal crosses of *A. arenosa* and *A. lyrata* resulted in one fragment because *AGL36-like* (I and VI, respectively in Figure 4A) from both species are the same length (Figure 4C). When *A. arenosa* is crossed as mother to *A. lyrata*, *AaAGL36-like* is successfully digested by EcoRI (Figure 4C, Table S1C). Using *A. lyrata* as a mother crossed to *A. arenosa*, no EcoRI digestion fragments occur and thus only expression of *AlAGL36-like* was observed (Figure 4C).

As a reciprocal control, we also used *A. lyrata* specific digestion by TauI, giving only digestion fragments in crosses with *A. lyrata* mothers (Figure 4C). To verify the identity of the amplified fragments and the maternal expression pattern, the undigested PCR products were Sanger sequenced and identified as only maternally contributed (Figure S5). In conclusion, only the maternal allele is expressed in reciprocal interspecies crosses between *A. arenosa* and *A. lyrata* indicating that imprinting of *AGL36-like* is preserved in *A. arenosa* x *A. lyrata* hybrid seeds.

Finally, we investigated *AGL36/ AGL36-like* imprinting in *A. thaliana* x *A. arenosa* hybrid seeds, using *A. arenosa* as the paternal cross partner. Upon amplification of cDNA, both maternally expressed *AtAGL36* and paternally expressed *AaAGL36-like* fragments could be identified (Figure 4D, lanes 5-6, see figure legend). Restriction with AlwNI digest the maternally contributed cDNA (Figure 4D, Table S1C). The paternal *AaAGL36-like* fragments remained undigested in hybrid crosses employing two independent *A. arenosa* populations (Figure 4D, lanes7-10). The paternal cDNA was verified by Sanger sequencing and comparison to the *A. arenosa* control (Figure 4D). We thus conclude that in hybrid *A. thaliana* x *A. arenosa* allele is reactivated, demonstrating differential action by the maternal species in the hybrid. Furthermore, this finding demonstrates that the paternal alleles of maternally expressed genes as described in previous studies (Josefsson et al. 2006; Burkart-Waco et al. 2015).

Genetic and environmental factors influence post-zygotic hybrid barriers

Deregulation of MADS-box type I TF genes has been implicated in setting up the post-zygotic hybridization barrier in incompatible hybrid *A. thaliana* x *A. arenosa* (Strecno1) seeds, and *A. thaliana* mutation of some of these genes could partially rescue the severe late seed phenotype in the same hybrid cross from approximately 1 to 10% (Walia et al. 2009). In order to systematically examine if the MADS-box type I TF genes analyzed in this work influence the strength of the hybrid barrier we noted that previous analyses to investigate hybrid barriers in *Arabidopsis* have been performed under slightly different temperature regimes (Josefsson et al. 2006; Walia et al. 2009; Burkart-Waco et al. 2012; Lafon-Placette et al. 2017) and in line with this, the major hypotheses to explain hybrid barriers are centered on genetic factors, not taking environmental variation into account. In Rice, however, it has been demonstrated that temperature affects cellularization of the endosperm (Folsom et al. 2014) and that Type I MADS box TF genes are deregulated during moderate heat stress (Chen et al. 2016). To rule out a temperature effect in our experiments, we therefore repeated crosses first performed by Walia et al. 2009 with Strecno1 and Col-0 using the original temperature regime (22°C) and at slightly lower temperatures (18°C) and also included the *A. arenosa* accession used in this study, MJ09-4. Surprisingly, both the difference in temperature and

genetic variation between *A. arenosa* populations had a major effect on the strength of the postzygotic hybrid barrier (Figure 5A,).

To quantify this observation we first investigated seed survival in the same crosses and temperature conditions. Using *A. arenosa* accessions MJ09-4 and Strecno1 (SN1) in crosses to *A. thaliana* Col-0 at both 18°C and 22°C, a substantial increase in the survival of hybrid seeds at 18°C for both accessions was observed (Figure 5B) with 18°C MJ09-4 replicates showing up to 60% live seeds while at the same time also obtaining the same results as Walia et al. 2009 when crossing Col-0 with Strecno1 at 22°C (live seed count 1%, N=162) (Figure 5B).

In hybrid seed germination experiments, the temperature dependency of the strength of the hybrid barrier became even more evident for the accessions MJ09-4 and SN1 (Figure 5C, p<0,001). Interestingly, when comparing two other *A. arenosa* accessions, MJ09-1 and SN2, in crosses to Col-0, these were found to be insensitive to the temperature change tested here although they display a higher seed survival rate than SN1 crossed to Col-0 (Figure 5C). These accessions may still be affected at larger differences in temperature due do genetic variation and different adaptation. Even though the variation between replicates is high, especially in 18°C crosses, a clear bypass effect of low temperature on the post-zygotic barrier is observed. Furthermore, germination of both 18°C and 22°C crosses demonstrate an increased germination rate of hybrid seeds involving the *A. arenosa* MJ09-4 population as paternal crosses crosses in *A. thaliana* accessions and *A. arenosa* MJ09-4 at 18°C and 22°C displayed no significant difference in germination between the temperatures (Figure S6). Furthermore, tetraploid *A. thaliana* mothers have been shown to alleviate the hybridization barrier, and to exclude this scenario we verified the diploidy of *A. thaliana* Col-0 accessions, the *A. arenosa* MJ09-4 population and the Strecno lines (SN1 and SN2) using flow cytometry (Figure S7).

Our findings indicate that genetic variation between *A. arenosa* populations also influence the success rate of hybridization, as previously demonstrated for different *A. thaliana* genotypes (Burkart-Waco et al. 2012). We therefore further investigated the temperature dependency of the hybrid barrier by varying the maternal *A.thaliana* accession used in the hybrid cross. Burkart-Waco et al. 2012 crossed 56 accessions using the Strecno1 line at 22°C to investigate the effect of the genetic variation on the hybrid barrier and could demonstrate a weaker barrier when using C24, producing 17% normal seeds, while using Ler-1 and WS-2 resulted in 5,2% and 3,5% normal seeds respectively. Using Col-0 they obtained 1,7% normal seeds. Here, we demonstrate that crossing Col-0, Ler-1, C24 and WS-2 accessions to the *A. arenosa* accession MJ09-4 give the effect of elevated seed survival for all accessions except Ws-2 (Burkart-Waco et al. 2012) when comparing to the previous report using SN1 at 22°C (Figure 5D). In addition, the accessions Col-0, C24 and Ws-2 have a significant increase in

seed survival when decreasing the temperature to 18° C (Figure 5D, p<0,05). Ler-1 appeared to be insensitive to the temperature change, similar to the observations using *A. arenosa* accessions MJ09-1 and SN2 in combination with Col-0 (Figure 5C).

Previously published data report that the embryo does not make the transition to the heart stage in crosses between diploid *A. thaliana* Col-0 and *A. arenosa* Strecno1 at 22°C (Burkart-Waco et al. 2013), which we also could confirm for Strecno1 under our laboratory conditions at 22°C (Figure 5). Our analysis at 18°C, however, demonstrates that most seeds develop past this point (Figure 6A-I). There was a clear correlation between the severity of the hybrid barrier in *A. arenosa* accessions and the timing of endosperm cellularization in hybrid seeds (Figure 6A-I) suggesting that endosperm cellularization is the major mechanism for setting up the barrier.

The role of MADS Type I loci in the A. thaliana A. arenosa hybrid barrier.

In order to investigate the specific effect of selected MADS type I loci in establishing or bridging the A. thaliana A. arenosa hybrid barrier, we analyzed insertional mutant alleles of the selected candidate genes (Figure S8A, Table S3). Homozygous mutants could be obtained for all investigated loci except as previously described for AGL62 (Kang et al. 2008), suggesting no vital requirement in seed development. Significantly reduced transcript levels were demonstrated in all lines with the exception of AGL34 where transcript levels were significantly elevated (Figure S8B). Segregation analysis could not detect reduced transmission of the mutant alleles suggestive of a requirement in male or female gametophytes or a recessive effect in embryo or endosperm (Figure S8C, Table S4). We also inspected seed size, seed germination and flowering time. For this analysis agl28-1 was omitted due to mixed Ws-2 Col-0 accession background. Only minor differences were observed in seed size and flowering time (Figure S9A-B) and no difference in germination of mutant seeds was observed (97-100%, N=200). Finally, seed developmental phenotypes in single and higher order mutants were investigated, scoring live, aborted and unfertilized seeds (Figure S9C). Notably, a heterozygous agl62-1 mutation in a double homozygous agl28-1 agl36-1 background did not differ from single agl62 mutants. We concluded that a thorough analysis of seed development in single, double and triple mutants of AGL34, AGL35, AGL36 and AGL90 including their interaction partners AGL28 and AGL62 did not result in any obvious seed developmental phenotypes (Figure S8, Figure S9).

In the case of *agl28-1* a mixed Ws-2 Col-0 accession background did not allow a direct comparison of hybrid seed barrier strength effects, due to the strong effect of the Ws-2 accession (Figure 5D, right panel). Indeed, single *agl28-1* mutants as well as double or triple mutant combinations with *agl36, agl62* and *agl90* crossed with *A. arenosa* all produced significantly lower seed germination rates in 9 out of of 10 cross combinations (Figure S10A). In a direct comparison using Ws-2 wild type as control, compared to Col-0 and *agl28-1* when crossed with *A. arenosa* MJ09-4, revealed no difference

in seed germination rate between Ws-2 and agl28-1(Figure S10B, left). In order to further investigate the role of agl28-1 we generated an AGL28 genomic rescue construct that was transformed into homozygous agl28-1 mutant background. Six homozygous T2 lines were crossed to A. arenosa MJ09-4 and scored for germination (Figure S10B, right). None of the rescue lines were significantly different from agl 28-1 or the Ws-2 control, suggesting that the observed reduced germination is caused by the Ws-2 background alone. Since the Ws-2 background effect co-segregated with the agl28 mutation through repeated introgression to Col-0, we hypothesized a major part of the genetic variation causing the strong Ws-2 A. arenosa hybrid barrier to be linked to the AGL28 locus. In order to genetically map the effect, we backcrossed a Col-0 introgressed hemizygous agl28-1 to Col-0 and genotyped the progeny for the presence of agl28-1, and crossed the two types of plants resulting with A. arenosa MJ09-4 pollen donors (Figure S10C, see legend for detail). Plants wild type for the AGL28 locus had a high probability to be Col-0 in the AGL28 region, and did also regain Col-0 germination rates in crosses with A. arenosa (Figure S10C, NS not significantly different). In addition, plants hemizygous for agl28-1 having a high probability to be Ws-2 Col-0 heterozygous in the AGL28 region, displayed intermediate germination levels and were still significantly different from Col-0 (Figure S10C, p<0,05). This strongly suggest the strong Ws-2 effect on the hybrid barrier to be linked to the AGL28 region on top of chromosome 1.

Finally, various single and double *A. thaliana* MADS-box type I mutants were crossed with the *A. arenosa* MJ09-4 population at the two temperatures established (18°C and 22°C). First, we wanted to investigate the influence of the mutated loci on the strength of the hybrid barrier, as measured by germination rate. Secondly, we wanted to explore if the identified temperature effect on the strength of the hybrid barrier was modulated by the mutated loci, as observed using in both *A. thaliana* and *A. arenosa* accessions (Figure 5C, D).

Hybrid seed phenotypes were inspected for some representative crosses at 18°C (Figure S11A-C). We observed the seed classes normal non-collapsed seeds, collapsed brown seeds and viviparous seeds (Figure S11A). The frequency of non-collapsed seeds in a silique and seed germination frequency were well correlated (Figure S11B). Seed size measurements also displayed variation but no strong effect of the mutant mother background (Figure S11C). The ploidies of *A. thaliana* x *A. arenosa* hybrids with both Col-0 and *agl36-1 agl90-2* as mother were verified using flow cytometry (Figure S7).

In general, none of the single or higher order *A. thaliana* mutants had a significant effect to alleviate the hybrid barrier when crossed to the *A. arenosa* MJ09-4 accession (Figure 7). This is conflicting with previous data reported for *agl62* and *agl90* crossed with an *A. arenosa* Strecno1 population at 22°C (Walia et al. 2009). In the Walia study, mutants of *agl62* and *agl90* used as mothers in the cross

increased the germination frequencies from 1% in crosses with Col-0 to 10% in crosses with *agl62* and *agl90*. In our experiment, the average germination frequencies of of *agl62-1* at 22°C, and the average germination frequencies of *agl34-2* at 22°C were indeed slightly higher than the Col-0 control crossed to *A. arenosa*, but the difference is not significant. We could therefore not reproduce the findings of Walia et al. in crosses using the *A. arenosa* MJ09-4 accession. The reason for this discrepancy may be genetic differences in the *A. arenosa* accessions used (MJ09-4 in this study versus Strecno1 in Walia et al.). As demonstrated here, different *A. arenosa* accessions can modulate the strength of the hybrid barrier (Figure 5C).

We did however see a significant effect of the single mutant *agl35-1* to aggravate the hybrid barrier when crossed to the *A. arenosa* MJ09-4 accession (Figure 7). Significant reduction was observed at both temperatures tested. When the homozygous *agl35-1* mutant was crossed to *A. arenosa* at 18°C, a significant reduction from average values of close to 50% germination to less than 20% was observed (Figure 7, p<0,001), and in the 22°C experiment we found a reduction from more than 25% to close to 15% (Figure 7, p<0,01). These findings suggest that AGL35 may play a role to relieve and bypass the hybrid barrier, or that lack of AGL35 disrupts or lowers the treshold for disruption of endossperm cellularization.

It is noteworthy that the effect of lower temperature to alleviate the hybrid barrier is also is bypassed by mutation of AGL35. We analyzed if any of the loci investigated played a role in establishing the observed temperature effect on the hybrid barrier strength (Figure 7). The Col-0 control and most mutant crosses, including *agl23,agl36, agl62* and *agl90* displayed significant differences in germination frequencies between the two temperatures examined (Figure 7). Crosses with the single mutants *agl34* and *agl35* and the *agl36 agl62* double mutant, however, displayed no significant differences in its germination rate between 18° and 22°C (Figure 7), similar to the previous observation in the L*er*-1 accession cross (Figure 5D). The temperature immunity seen in *agl35* appear to be the most prominent due to low variation in the observations, and may suggest that the *AGL35* links the hybrid block to the temperature effect.

Discussion

The role and regulation of imprinted genes

In this report we have systematically analyzed the function and conservation of imprinting of a subset of MADS-box type I TFs in hybrid crosses in the context of biological and environmental variance. In our expression analysis we observe that M α and M γ class type I TFs are highly represented in the transcriptome of the developing seed, and that both classes contain correlated transcript profiles that peak around the onset of endosperm cellularization. The expression peak occurs at a crucial developmental time point when the endosperm switches from nutrient sink to source for the developing embryo (Lafon-Placette et al. 2016), which is suggestive of a function in this process. Defects in endosperm cellularization are also the mechanistic basis for a post-zygotic reproductive barrier between *Arabidopsis* species and thus support a role for these genes in a hybrid scenario. The M α and M γ class co-regulation is consistent with the notion that M α forms dimers with the M γ -type MADS-box TF class (de Folter et al. 2005). Furthermore, since imprinted genes interact with biparentally expressed genes this would favor the dosage hypothesis for the selection of imprinted genes (Dilkes and Comai 2004). According to this hypothesis imprinting is a means to optimize the expression level of a gene, opposed by the parental conflict theory, where parental conflict over resources leads to selection of genes that promote or restrict resource allocation (Haig and Westoby 1991).

It has been postulated that maternally expressed imprinted genes are regulated by the release of DNA methylation in the central cell by central cell specific action of the DNA glycosylase DEMETER (DME). We could, however, demonstrate that lack of DNA methylation maintenance in the pollen germline does not activate all silenced paternal alleles. There is a discrepancy between the lack of activation of paternal *AGL36* observed in this work and previously published data (Shirzadi et al. 2011). However, the two studies were done using different accessions and also different *met1* mutant alleles. Accession specific effects are less likely since we see paternal reactivation of *AGL28* using the same accessions. However, since the history of zygosity is not known for the *met1* allele used in the previous study, accumulated hypomethylation may explain the inconsistency. Nevertheless, even though the paternal allele of *AGL28* reactivated in our study, paternal silencing of *AGL90* and *AGL36* was not lifted. In this study, *met1-7* was held as heterozygous, thus avoiding parental demethylation. This suggests that maternally expressed imprinted genes may be regulated by other mechanisms or have different regulatory requirements.

Evolution of silencing of the paternal allele of maternally expressed imprinted genes has been associated to global methylation patterns generated towards suppressing transposons (Kim and Zilberman 2014; Anderson and Springer 2018). *De novo* methylation of transposons is mainly performed by the action of RdDM, and in the lack of a mechanistic scenario for the imprinting of maternally expressed genes this pathway is an obvious candidate. Several M α and M γ class type I TFs including *AGL36* have also been suggested to be upregulated in lack of RdDM (Lu et al. 2012). In the case of *AGL36* we could, however, not observe any change in paternal silencing in reciprocal crosses with a PolIV mutant (*nrpd1*), i.e. blocking canonical RdDM. Furthermore, we were also unable to verify the previously reported upregulation of *AGL36* in the same crosses and time points (Lu et al.

2012). In order to determine the global effect, however, a systematic elucidation of the role of RdDM in the regulation of maternally expressed imprinted genes is required.

Although it does not regulate imprinting of maternally expressed genes, we show that repression by PRC2 is specifically targeted towards M α and M γ class type I TFs and acts to repress gene activity concurrent with and at post-cellularization stages. In absence of MEA, only the maternal allele of *AGL36* is upregulated (Shirzadi et al. 2011), indicating that the mode of regulation does not interfere with the actual imprinting mechanism. We hypothesize that DNA methylation of the paternal allele protects from PRC2 repression, as suggested for the paternally expressed imprinted gene *PHE1* (*AGL37*) (Villar et al. 2009; Makarevich et al. 2008; Kohler et al. 2003). In such a scenario the paternal allele of paternally expressed genes should be targeted by PRC2 at cellularization, but the mechanism here remains to be explored. We observe, however, that M α and M γ class type I TFs are regulated by PRC2 in distinct clusters from 6 to 12 DAP and that maternally and paternally expressed genes are co-regulated in the same cluster. While the meaning of the observed gradual repression by FIS-PRC2 in the context of M α and M γ role in seed development remains open to speculation, it seems clear that FIS-PRC2 acts through specific sets of genes at successive stages in seed development, rather than exerting a global effect in the seed concurrent with endosperm cellularization.

Species dependent deregulation of imprinting

We identified orthologs of *AGL36* in *A. arenosa* and *A. lyrata* and studied imprinting in *A. arenosa* and hybrids of *A. arenosa*, *A. lyrata* and *A. thaliana*. *AGL36* is imprinted in *A. thaliana* and maternally expressed, and here we prove that one of the two *AGL36-like* genes in *A. arenosa* is also exclusively maternally expressed, demonstrating conservation of imprinting. The maternal allele of the paternally expressed imprinted genes have previously been shown to be reactivated in the hybrid of *A. thaliana* and *A. arenosa* (Walia et al. 2009; Josefsson et al. 2006; Burkart-Waco et al. 2015). We show that, in hybrid *A. thaliana* x *A. arenosa* seeds, the paternal *AGL36-like* allele is reactivated, suggesting that deregulation in hybrid crosses is not limited to paternally expressed genes as described in previous studies (Walia et al. 2009). This loss of regulation is in strong contrast to the *A. lyrata* x *A. arenosa* hybrid cross where the imprinting of *AGL36-like* is maintained.

The variation or perturbation of the pattern of imprinting might play a role in the endosperm hybridization barrier between species (Florez-Rueda et al. 2016), and in the case described here, different mechanisms may act in the same species depending on the cross partner. Previous imprinting analyses involved crosses between inbreeders and outbreeders, fueling speculation that the mechanisms regulating imprinted genes may differ depending on the mating system (Josefsson et al. 2006; Burkart-Waco et al. (2012); Hatorangan et al. 2016; Klosinska et al. 2016). However,

intraspecific variation of imprinting within *A. thaliana* has previously been demonstrated (Waters et al. 2013; Pignatta et al. 2014). Consistent with the notion that imprinting can vary independently of mating systems, reactivation of the normally imprinted paternal *A. arenosa AGL36* allele depended on the maternal species: it was observed only in crosses to *A. thaliana* but not to *A. lyrata*.

A temperature and population dependent quantitative barrier phenotype

We have identified an important role of temperature in the establishment of endosperm based postzygotic hybrid barriers. This opens for speculation and a multifaceted scenario emerges identifying several parameters; both intrinsic variation in genetic pathways in both parents and external abiotic factors such as temperature may act in concert to mediate the generation of post-zygotic species barriers.

The discovery that lowering the temperature by only 4°C from 22°C to 18°C degrees during the fertilization and development of the *A. thaliana* x *A. arenosa* hybrid seed increases survival was surprising. Nevertheless, incompatibility between diverged individuals can be sensitive to temperature as, for example, in seedling hybrid necrosis (Bomblies and Weigel 2007). In this case, however, appropriate temperatures can ameliorate acute incompatibilities manifested only during early seed development and, therefore, may play a significant role in reducing interspecific hybridization barriers. High environmental sensitivity may also explain inconsistencies with previous studies noted in Results.

Crossing A. thaliana wild type Col-0 and mutants to different A. arenosa accessions demonstrated variation depending on the pollen donor and no significant increase in seed viability could be observed by using the mutants of the MADS-box type I TFs as mothers. The variability caused by natural variation has been demonstrated in A. thaliana by using different accessions. Burkart-Waco et al. (2012) used the diploid A. arenosa accession Strecno-1 and crossed it as the pollen donor to 56 A. thaliana accessions and scored seed viability ranging from 30% normal seeds to close to 2%. Burkart-Waco et al. (2013) compared seed development of A. thaliana accessions Col-0 and C24 crossed to A. *arenosa*, which respectively produced 0 to 1% and \sim 17% live hybrid seeds. Hybrid embryos at all stages beyond 2 DAP were developmentally delayed and similar between Col-0 and C24, with the exception of a few C24 x A. arenosa hybrid embryos displaying developmental progression to heart stage by 6 DAP, whereas no Col-0 embryos made this transition. This is in clear contrast to our findings where most seed survive this stage. Using different A. thaliana accessions in crosses to the A. arenosa accession MJ09-4, we obtained highest seed germination rates in crosses with C24. However, the cross combination with the Ws-2 accession produced similarly low germination rates as the A. arenosa accession Streeno-1 crossed with Col-0. This indicates that rather than specific accession having specific effects, it is the combination of accessions that determines the strength of the hybrid

barrier. Although we could not observe elevated seed viability when single or multiple MADS-box type I mutants were used in crosses to *A. arenosa*, deficiency of *AGL35* resulted in significant reduction of the germination frequency. Notably, mutants of *AGL35* crossed to *A. arenosa* display two effects: first, it increase lethality; second, it decrease or eliminate the temperature effect, suggesting a critical role for this gene in mediating the strength of the hybrid block and the temperature effect.

Weakening of hybridization barriers at lower temperatures might increase fitness of a selfincompatible colonizer by broadening mate choice: few viable seeds are better than no seed. This mechanism could, for instance, have been important for recolonization after the Pleistocene glaciations, a period that was characterized by secondary contact and high amount of hybridization with or without genome duplication (Stebbins 1984; Brochmann et al. 2004). Although A. thaliana is a self-crosser, the mechanism might be ancestral and predate A. thaliana. Its occurrence should be investigated by hybridizing more species under varying temperature. In northern Europe and the Fennoscandian region, A. thaliana with unreduced gametes has most likely hybridized with pollen from tetraploid A. arenosa to create the allotetraploid A. suecica on multiple hybridization events (Novikova et al. 2017). Making a synthetic tetraploid A. *thaliana* and crossing it to A. *arenosa*, can make viable, although unstable, hybrids (Comai et al. 2000). Environmental stress such as heat or cold may increase the rate of unreduced gamete formation (De Storme and Mason 2014) and facilitate hybridization between diploids and tetraploids (Vallejo-Marin and Hiscock 2016). Such instances of genome doubling, however, did not occur in our experiments as flow cytometry of sampled hybrids indicated genome content consistent with reduced gametes of the diploid parents A. thaliana and A. arenosa. Formation of 2N gametes may be under different regulation in Arabidopsis. The temperature sensitive mechanism affecting the endosperm-based barrier and its dependency on AGL function remain an open area of investigation.

In Rice, it has been shown that temperature affects cellularization of the endosperm (Folsom et al. 2014) and that Type I MADS box TF genes are deregulated during moderate heat stress (Chen et al. 2016). The temperature stress tested were much higher than tested here for *Arabidopsis*, but the different species have different temperature adaptations in general. The rice MADS box TF OsMADS87 is a heat-sensitive imprinted gene which is associated with syncytial stage endosperm and regulates rice seed size (Chen et al. 2016). OsMADS87 is a putative ortholog of *Arabidopsis PHE1*. Mutants of OsMADS87 have accelerated endosperm cellularization and lower sensitivity to a moderate heat stress in terms of seed size (Folsom et al. 2014, Chen et al. 2016). Our results indicate that some of the *Arabidopsis thaliana* MADS box mutants hybridized to *A. arenosa*, display a lowered heat sensitivity that may indicate a temperature sensing role for the MADS box genes in the endosperm of *A. thaliana* as well. Considering that many of the Type I MADS box genes are regulated by the PRC2 complex and that the finding that OsFIE1 is imprinted and temperature

sensitive in rice seeds (Folsom et al. 2014), this proposes an epigenetic regulation during hybrid seed development which is altered during environmental perturbations (Folsom et al. 2014). Given the importance of overcoming post-zygotic isolation for the early stages of neo-hybridization (Vallejo-Marin and Hiscock 2016), this temperature effect can be a useful tool for investigating the endosperm based post-zygotic barrier and early speciation.

Materials and methods

Plant material and cultivation

A. thaliana accessions and mutant plant lines were obtained from the Nottingham Arabidopsis Stock Center (NASC) unless specified otherwise. For details on MADS-box type I mutant lines see Table S1. The *met1-7* and *nrpd1* accession numbers are SALK 076522 and SALK 083051, respectively. The *mea/fis1* mutant was kindly supplied and described in Chaudhury et al. (1997). The A. arenosa populations MJ09-1 and MJ09-4 and the A. lyrata MJ09-11 descended from natural populations in central Europe as described, respectively, by Jørgensen et al. (2011) and Lafon-Placette et al. (2017). A. arenosa populations Streeno1 (SN1) and Streeno2 (SN2) were kindly supplied by Kirsten Bomblies (Hollister et al. 2012). Seeds were surface sterilized either by washing steps with 70% ethanol, 20% bleach and wash solution (0.001% Tween20) or by over-night chlorine gas sterilization (Lindsey et al. 2017), sown out on 0.5 MS plates (Murashige and Skoog 1962) supplemented with 2% sucrose and appropriate antibiotics for selection of mutant lines. The seeds were then stratified overnight (A. thaliana) or 1 to 3 weeks (A. arenosa and A. lyrata) at 4°C before transferring to growth chambers with either 18°C or 22°C under long day conditions (16 hr light, 160 µmol/m²/s, relative humidity 60-65%). A. thaliana A. arenosa F1 hybrid seeds were stratified at 4°C for 4-6 days before being placed in growth chambers for scoring of germination as seedling root protruding from the seed regardless of survival at later stages. Germinated seedlings were transferred to soil and grown under long day conditions at 18°C or 22°C. A. arenosa and A. lyrata plants were vernalized at 8°C under short day conditions (10 hr light) for 4-5 weeks to stimulate flowering. To avoid self-pollination, controlled crosses were performed by emasculating unopened flower buds followed by hand pollination after 2 days. Developing or mature seeds were harvested for designated purposes at defined time points. See Table S5 for an overview of interspecies crosses. For flowering time analysis, flowering time was scored as day after stratification and the average number of leaves at stem emergence from the rosette.

Tissue handling, DNA and RNA extraction and cDNA synthesis

Tissue was harvested directly in liquid nitrogen and DNA was isolated using E.Z.N.A. Plant DNA kit (Omega) according to manufacturer's instructions. For total RNA isolation, seeds were hand-dissected from siliques directly into pre-chilled tubes with MagNA Lyser Green Beads (Roche) and grinded in lysis buffer (Sigma Plant total RNA kit) using a MagNA Lyser Instrument (Roche). Isolated RNA was treated with DNase I (Sigma) and cDNA synthesized with Oligo (dT) and Superscript III reverse transcriptase (Invitrogen). Samples were cleaned using a QIAquick PCR purification kit (Qiagen). DNA or RNA concentration was measured using a NanoDrop1000 spectrophotometer or RNA was measured with a Qubit 3 fluorometer (ThermoFisher) using the Qubit RNA BR Assay kit (Invitrogen). All kits were used according to the manufacturers' instructions.

RNA Sequencing and sequence analysis

Total RNA was isolated from dissected seeds 1, 2, 3, 4, 6, 9 and 12 DAP from Ler crossed to Col-0 and from *mea/fis1* crossed to Col-0 in two biological replicates as described previously (Shirzadi et al. 2011). RNA samples were DNase treated before quality checked using an Agilent 2100 Bioanalyzer. Total RNA was prepared to a Strand-specific TruSeqTM RNA-seq library and all 28 samples sequenced over 3 lanes on an Illumina HighSeq 4000, 150 bp paired end reads. Differential expression analysis was performed with RSEM (Li and Dewey 2011) using the edgeR software package (McCarthy et al. 2012). The expression profiles were analyzed and visualized using the Tidyverse 1.2.1, ComplexHeatmap 1.17.1 (Gu et al. 2016), Dendextend 1.7.0 (Galili 2015), Viridis 0.5.0 packages in R version 3.4.3.

Molecular cloning and genotyping

All T-DNA mutant lines were genotyped using specific primers (Table S1, Table S2). Due to high sequence similarities between MADS box genes and AGL36-like genes in different species, primers were optimized to ensure specific amplification and fragments were sequenced for confirmation of identity. The sequencing and characterization of *AGL36*-like genes from MJ09 *A. arenosa* and *A. lyrata* lines were performed by PCR amplification with designated primers (Table S2) using KOD Hot Start DNA polymerase (Sigma) according to manufacturer's instructions with 1,5 mM of MgSO₄ and PCR program with 95°C denaturation, 55°C annealing, and 70°C extension for 35 cycles. The amplified fragment was subsequently cloned into a TOPO Blunt pCR Zero vector (Invitrogen) according to the manufacturers' instructions. The *AGL28* genomic complementation construct was created by nested PCR using primers described in Table S2 containing *att* sites for GateWay cloning (Invitrogen) according to the manufactures' instructions. The genomic *AGL28* fragment was 3531 bp including 2000 bp upstream of the start codon and 500 bp downstream of the stop codon and was cloned into the destination vector pMDC99. The construct was transformed into *Agrobacterium tumefaciens* strain GV3101 pMP90RK which was used to transform *agl28-1* mutant using the floral

dip method (Clough and Bent, 1998) and transformants were selected for by the appropriate resistance encoded in the inserted T-DNA, hygromycin. All sequences from *A. arenosa* and *A. lyrata* generated for this study have been deposited in the National Center for Biotechnology Information Sequence Read Archive (https://www.ncbi.nlm.nih.gov/) with accession numbers MN380433 to MN380437.

Phylogenetic analysis

All alpha and gamma MADS-box genes were extracted from the *A. thaliana* genome (TAIR10 at https://www.arabidopsis.org/). The genomic sequence was translated into amino acid with the AUGUSTUS gene prediction program (Stanke and Morgenstern 2005) using the *A. thaliana* gene model. Nucleotides from the coding regions were aligned based on the protein sequence with PAL2NAL (Suyama et al. 2006). Positions with more than 80% gaps and ambiguously aligned positions were removed from the alignment. A phylogenetic tree was inferred from the resulting alignment using the GTRGAMMA model and the automatic bootstrapping criteria MRE (option -1 autoMRE) in RAxML v8.0.26 (Stamatakis 2014; Pattengale et al. 2011). The genes *AGL34*, *AGL36*, and *AGL90* from *A. thaliana* were used as queries in BLAST against the non-redundant nucleotide database at NCBI (blast.ncbi.nlm.nih.gov/) to find all homologous genes in the *Arabidopsis* genus. In addition, all available *Arabidopsis* Illumina whole genome sequence data from Sequence Read Archive (SRA) was employed in the phylogenetic trees were identified with BLAST. All *AGL36* related genes were aligned and phylogenetic trees were inferred as for the MADS-box genes. In the final analysis only one copy of the gene was kept for each species.

Real-time quantitative PCR

Real-time PCR was performed on a LightCycler 96 instrument using FastStart Essential DNA Probes Master protocol (Roche) and FastStart Essential DNA Green Master protocol (Roche) using designated primers (Table S2). Relative expressions were calculated according to Pfaffl (2001) and are average values of at least two biological replicates. In reactions with low relative expression an E-value of 2.0 was used. All PCR products were sequenced to verify identity of the product amplified.

Single nucleotide polymorphism (SNP) analysis

Seed tissue was sampled at 4 DAP from reciprocal crosses from *A. thaliana*, at 7 DAP from crosses between *A. thaliana* and *A. arenosa*, and at 9 DAP from reciprocal crosses of *A. arenosa* and *A. lyrata*. cDNA was amplified by PCR using designated primers (Table S2) and digested using SNP specific enzymes (Table S4) analyzed on an Agilent 2100 Bioanalyzer using the DNA-1000-LabOnChip system (Agilent Technologies). Images for figures were assembled using Illustrator software (Adobe).

Dry seed phenotyping using ImageJ

For *A. thaliana A. arenosa* hybrid crosses, mature seeds were harvested one silique at a time before imaging using a Nikon D90 and analysis using ImageJ to determine seed size (mm^2) . Viviparous seeds were excluded from the analysis. MADS-box type I mutants were grown and harvested at the same time and conditions. Seed size was measured and analyzed using ImageJ to determine the average seed size (mm^2) per plant. The significance differences between plant lines were tested using Kruskal-Wallis test: p=0,0047 and in pairwise comparison to wildtype (Wilcoxon rank-sum test).

Microscopy and Feulgen staining of seeds

Tissue was harvested from *A. thaliana* and *A. thaliana* x *A. arenosa* at 7 DAP and 10 DAP (seeds from 3 siliques per biological replicate). The seeds were stained with Schiff's reagent (Sigma-Aldrich S5133) following fixation and embedding in LR White (London Resin) as described by Braselton et al. (1996). An Olympus FluoView 1000 Confocal Laser Scanning microscope (BX61WI) with an excitation of 488 nm and emission from 500 to 600 nm was used for imaging.

Ploidy measurements of adult hybrids as well as crossing parents

Ploidy was measured for a selection of the plants with two of the A. arenosa parents (from MJ09-4 population), two Col-0 plant individuals, 14 A. thaliana Col-0 x A. arenosa F1 hybrids and 12 A. thaliana agl36-1 agl90-2 x A. arenosa F1 hybrids. Also, the A. arenosa lines Strecno1 and Strecno2 were included to confirm ploidy. One rosette leaf and one inflorescence were analysed for all samples except for agl36-1 agl90-2 x A. arenosa F1 hybrids where only rosette leaves were analysed. The ploidy was assessed by establishing the genome content by estimating the relative fluorescence intensities by flow cytometry (FCM) and the two-step methodology according to Dolezel et al. (2007). The reference standards for the raw cytometric analysis were Solanum pseudocapsicum for the A. thaliana and A. arenosa hybrid comparison to A. thaliana, and Carex acutiformis for the A. arenosa Strecno1 (N=10) and Strecno2 (N=12). The samples and the internal reference were chopped with a razor blade in 0.5 ml ice-cold Otto I buffer (0.1 M citric acid, 0.5% Tween 20). This was then filtered through a nylon mesh (loop size $0.42 \mu m$), incubated at room temperature for 5 min before being stained with 1 ml of Otto II buffer (0.4 M Na₂HPO₄ \cdot 12 H₂O) supplemented with AT-selective fluorescent dye DAPI (4',6-diamino-2-phenylindol) and 2-mercaptoethanol in final concentrations of 4 µg/ml and 2µl/ml, respectively. After about 5 min of incubation at room temperature, the relative fluorescence intensity for a minimum of 3000 nuclei was recorded using a Partec Space flow cytometer (Partec GmbH, Münster, Germany) equipped with an UV-LED chip (365 nm). The FCM results are the fluorescence intensities relative to unit fluorescence intensity of the internal reference standard.

Accession Numbers

All sequences generated in this study have been deposited in the National Center for Biotechnology Information Sequence Read Archive (https://www.ncbi.nlm.nih.gov/sra/) with project number PRJNA562212.

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The authors declare no conflicts of interest.

List of author contributions:

K.N.B. and P.E.G. conceived the project and wrote the article with contributions of all the authors. K.N.B. designed the experiments. K.N.B., J.B., K.S.H. and I.M.J. performed experiments. K.N.B., J.B. and K.S.H. analyzed the data. K.S.H., I.M.J., A.K.K., Y.S.E., M.K. and R.S. provided technical assistance. K.N.B., J.B. and K.S.H. made figures. J.B., A.K.B. and L.C. contributed to project design and discussion. P.E.G. supervised the project and agrees to serve as the author responsible for contact and ensures communication.

Supplementary legends

Figure S1: Phylogeny and expression of MADS-box type I transcription factors during seed development.

Figure S2: Imprinting analysis of AGL28, AGL35, AGL36 and AGL90.

Figure S3: Clustering of MADS-box type I genes based on expression pattern in wild type and *mea* mutant seeds.

Figure S4: *AGL36* related genes from all available *Arabidopsis* genomes in Sequence Read Archive, GenBank and Phytozome.

Figure S6: Germination rate in self crosses of *A. thaliana* accessions and *A. arenosa* MJ09-4 is not affected by temperature.

Figure S7: Ploidy measurement of Arabidopsis populations and hybrids.

Figure S8: Genetic analysis of selected MADS-box type I transcription factors.

Figure S9: Phenotypic characterization of MADS-box transcription factor mutants.

Figure S10: Characterization of a genetic background effect in mixed *A. thaliana* accessions crossed to *A. arenosa*.

Figure S11: F1 hybrid seed phenotypes from *A. arenosa* crosses to *A. thaliana* Col-0, *agl36-1*, *agl90-1*, *agl90-1* and *agl36-1* agl90-2 at 18°C.

Table S1: Schematic overview of restriction digest set-up.

Table S2 Oligonucleotide name, sequence and description.

Table S3: Characterization of mutants used in this study.

Table S4: Segregation of the mutant alleles *agl28-1, agl34-2, agl35-1, agl36-1, agl90-1* and *agl90-2* in self crosses and in reciprocal crosses to wild type (Col-0).

Table S5: Crossing scheme for various A. thaliana to A. arenosa experiments.

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Figure legends

Figure 1: MADS-box type I transcription factors share similar expression profiles during seed development. A) Maximum likelihood phylogeny of alpha and gamma MADS-box type I genes in *Arabidopsis thaliana*. The tree was inferred using the GTRGAMMA model on 41 genes with 532 unambiguously aligned nucleotides. Scale bar represents the mean number of nucleotide substitutions per site. Only bootstrap values above 65% are shown. **B**) Gene expression profiles of alpha and gamma MADS-box type I genes were ordered in five groups according to the branching pattern. Transcript quantification and differential expression analysis was performed with RSEM and visualized using R. Gene expression profiles for stages ranging from one to 12 days after pollination (DAP) are relative to four DAP using a base-2 logarithmic scale (logFC). Two biological replicas with three technical replicas were analysed. Note that genes within groups show similar gene expression profiles, with a common maximum reached between three and six DAP.

Figure 2: Imprinting and epigenetic regulation of *AGL28*, *AGL35*, *AGL36* and *AGL90*. **A**) Imprinting analysis of *AGL28*, *AGL35*, *AGL36* and *AGL90* using accession specific restriction digest on single nucleotide polymorphisms (SNPs) in reciprocal crosses between accessions Tsu-1 and Col-0. For each panel, the accession specific digestion pattern is indicated. Seeds were harvested for analysis four days after pollination (DAP). Bioanalyzer images of one of three biological replicas is shown. **B**) SNP analysis of *AGL28*, *AGL35*, *AGL36* and *AGL90* in crosses with *met1-7+/-* (Col-0 background) pollen. Only *AGL28* display paternal activation in the *met1-7+/-* mutant. Wild type (WT) crosses are duplicated from **A**) for visualization. Crosses were harvested as **A**). Bioanalyzer images of one of

three biological replicas is shown. **C**) SNP analysis of *AGL36* in reciprocal crosses between WT (Col-0) or *nrpd1* (Col-0) and WT (L*er*). Crosses were harvested six DAP. The *AGL36* imprinting pattern is not changed in the *nrpd1* crosses compared to WT crosses. **D**) Real-time PCR analysis of *AGL36* expression in three biological replicas the *nrpd1* background at six DAP. The relative expression difference is not significant (NS, t-test; p=0,389). **E**) Real-time PCR verifying significant knock-down (t-test; p=0,015, indicated by asterisk) of *NRPD1* in three biological replicas of *nrpd1* homozygous background. Error bar indicate standard deviation (SD).

Figure 3: PRC2 dependent transcriptional repression of MADS-box type I transcription factors during seed development. A) Heat-map clustering of gene expression profiles of MADS-box type I transcription factors (TFs) in the Polycomb Repressive Complex 2 (PRC2) *MEDEA* mutant seeds (*mea*) compared to wild type (WT). Expression profile based clustering, transcript quantification and differential expression analysis was performed for three biological replicas using RSEM and visualized using R. Differential gene expression profiles for stages ranging from one to twelve days after pollination (DAP) are shown using a base-2 logarithmic scale (logFC). Note strong expression change in three out of four expression clusters. B) Box-plot showing class specific expression of MADS-box type I TFs at stages from one to twelve DAP. The MEA dependent repression is shared by the alpha (M α) and gamma (M γ) class TFs, while the beta (M β) class is weakly affected in the *mea* background. Relative expression changes are displayed in a base-2 logarithmic scale.

Figure 4: Conservation and imprinting of AGL36-like genes in Arabidopsis and in selected hybrid crosses. A) Maximum likelihood phylogeny of AGL36-like genes in A. arenosa, A. pedemontana, A. halleri, A. lyrata and A. thaliana. The tree was inferred using the GTRGAMMA model on 15 sequences with 938 unambiguously aligned nucleotides. Because of space limitations branches marked ¹/₂ are shortened to half their original length. Only bootstrap values above 65% are shown. Scale bar represents the mean number of nucleotide substitutions per site. Arabidopsis species are indicated by colors. Roman numerals (right) indicate distinct genes in the respective species and is used as reference in B-D. B) SNP analysis of A. arenosa AGL36 (I) in MJ09-4 background at 9 days after pollination (DAP). Left half, undigested; right half, digested with BceAI. Cross plants where AaAGL36 is digested by BceAI are indicated "A.a.". Cross plants where AaAGL36 is not digested by BceAI are indicated "A.a. SNP". The shorter 475 bp fragment is AaAGL36. AaAGL36 (I) is maternally expressed in A. arenosa as only the undigested AGL36 fragment is present when A.a SNP is the mother, whereas completly digested fragments result when A.a. is the mother. C) Imprinting and maternal expression of AGL36 is conserved in reciprocal crosses between A. arenosa and A. lyrata at 9 DAP. AaAGL36 (I) and AlAGL36 (VI) fragments are both 768 bp of length (lanes 1-2). EcoRI digests AaAGL36 (I) only (lanes 3-4) and TauI digests AlAGL36 (VI) only (lanes 8-9). In

hybrid crosses, only the maternal fragments are digested, and no or very weak undigested fragments are left (lanes 5-7, 10-12). **D**) Imprinting of *AGL36* is lifted in crosses between *A. thaliana* and *A. arenosa*. The undigested fragments of *AtAGL36* (VIII) and *AaAGL36* (I) are 819 and 768 bp, respectively (lanes 1 and 3). An AlwNI restriction site is only present in *AtAGL36* (VIII) (lanes 2 and 4). In hybrid crosses, both fragments are visible (lanes 5-6) and AlwNI restriction digest only the maternal AtAGL36 fragment. Note that the paternal *AaAGL36* (I) allele is expressed (lanes 7-10). EcoRI digests both *AaAGL36* and *AtAGL36* leaving only digested fragments (lanes 11-12). BR, biological replicate; *A.t., A. thaliana; A.a., A. arenosa; A.l., A. lyrata*. All crosses shown are in the order female x male. Two biological replica represent results from three biological replicas tested. DAP stages used are 4 for *A.t.,* 7 for *A.t.xA.a* and 9 DAP for *A.a., A.l.* and the *A.axA.l.* reciprocal cross.

Figure 5: Temperature has a significant effect on the hybrid barrier between A. thaliana and A. arenosa. A) Micrographs of siliques with A. thaliana crossed to A. arenosa hybrid F1 seeds grown at 18°C and 22°C 20 days after pollination (DAP). The crosses were made using two different A. arenosa lines, MJ09-4 and Strecno1 (SN1). Live seeds are green, collapsed seeds are brown or pale green. B) Graph showing percentage live A. thaliana x A. arenosa hybrid F1 seeds from crosses in A). Three biological replicates were tested for each temperature for both A. arenosa lines MJ09-4 and SN1 (N=174, 163, 175, 162, respectively). C) Germination rate of A. thaliana x A. arenosa hybrid F1 seeds. Four A. arenosa accessions were crossed to A. thaliana at 18°C and 22°C, MJ09-1, MJ09-4, SN1 and SN2 (18°C: MJ09-1 N= 18 BR (931 seeds), MJ09-4 N= 18 BR (986 seeds), SN1 N= 9 BR (524 seeds) and SN2 N= 9 BR (475 seeds). 22°C: MJ09-1 N= 12 BR (612 seeds), MJ09-4 N= 36 BR (1482 seeds), SN1 N= 36 BR (1544 seeds), SN2= 12 BR (673 seeds)). D) Germination rate of A. thaliana Col-0, Ler, C24 and Ws-2 crossed to A. arenosa MJ09-4 at 18°C and 22°C (18°C: Col-0 N= 12 BR (572 seeds), C24 N= 8 BR (451 seeds), Ler N= 12 BR (772 seeds), Ws-2 N= 12 BR (622 seeds). 22°C: Col-0 N= 24 BR (1212 seeds), C24 N= 12 BR (462 seeds), Ler N= 12 BR (751 seeds), Ws-2 N= 12 BR (727 seeds)). A.t., A. thaliana; A.a., A.arenosa. Blue colour: 18°C, red colour: 22°C. Outliers are plotted as large points. BR, biological replicas. Significance is indicated for the comparison of lines at 18°C and 22°C (Wilcoxon rank-sum test: NS.: p > 0.05; *: $p \le 0.05$; *: $p \le 0.05$; *: $p \le 0.01$; ***: $p \le 0.001$). Error bar indicate standard deviation (SD).

Figure 6: Variation in endosperm cellularization between *A. thaliana* and *A. arenosa* hybrids. **A-H**) Confocal scanning laser micrographs of endosperm cellularization in hybrid seeds visualized by Feulgen staining. For all crosses, both non-cellularized and cellularized endosperm is observed and micrographs representative for each class are presented in the left and right panels respectively. Openarrow heads point to syncytial endosperm nuclei while closed-arrow heads point to cellularized endosperm nuclei. Scale bar=50µm. **A-B**) *A. thaliana* control 7 days after pollination (DAP) typically at the embryo late heart stage where most seeds display complete endosperm cellularization (**B**). **C-D**) *A. thaliana* x *A. arenosa* MJ09 hybrid seeds at 7 DAP. Embryo development is slower compared to *A. thaliana* controls. Both non-cellularized (**C**) and cellularized endosperm (**D**) was frequently observed. **E-F**) *A. thaliana* x *A. arenosa* MJ09 hybrid seeds at 10 DAP. Only a few seeds fail to cellularize (**E**) and most seeds exhibit completed endosperm cellularization (**F**). **G-H**) *A. thaliana* x *A. arenosa* SN1 hybrid seeds at 10 DAP. A higher fraction of seeds display syncytial stage endosperm (**G**) compared to *A. arenosa* MJ09 hybrid seeds (**E-F**), but some have completed endosperm differentiation (**H**). **I**) Quantification of the described embryo and endosperm stages. All crosses are indicated as female x male. TxT, *A. thaliana* seeds, N=34; TxA, *A. thaliana* x *A. arenosa* MJ09 hybrid seeds, N=81; TxS, *A. thaliana* x *A. arenosa* SN1 hybrid seeds, N=98; em, embryo stages; en, endosperm stages.

Figure 7: Genetic and environmental parameters influence the F1 hybrid barrier. A) Germination rate of seeds from *A. arenosa* MJ09-4 crossed as pollen to the *A. thaliana* (Col-0), single mutants *agl23-1*, *agl34-2*, *agl35-1*, *agl36-1*, *agl62-1*, *agl90-1*, and *agl90-2* and the double mutant *agl36-1 agl62-1* at 18°C and 22°C. Box-plot contains scattered dots representing germination rates observed per silique. Outliers are plotted as large points. Significance is indicated for the comparison of the mutant lines between 18°C and 22°C (Kruskal-Wallis test: p<2,2-16; Wilcoxon rank-sum test: NS.: p > 0.05; *: $p\leq0,05$; **: $p\leq0,01$; ***: $p\leq0,001$). 18°C: Col-0 N= 10 BR (536 seeds), *agl23-1* N= 12 BR (294 seeds), *agl34-2* N= 12 BR (707 seeds), *agl35-1* N= 12 BR (641 seeds), *agl36-1* N= 12 BR (704 seeds), *agl36-1 agl62-1* N= 12 BR (532 seeds), *agl62-1* N= 8 BR (442 seeds), *agl23-1* N= 12 BR (407 seeds), *agl34-2* N= 12 BR (610 seeds), *agl35-1* N= 12 BR (578 seeds), *agl36-1* N= 12 BR (635 seeds), *agl36-1* N= 12 BR (635 seeds), *agl36-1* N= 12 BR (635 seeds), *agl36-1* N= 12 BR (610 seeds), *agl62-1* N= 8 BR (403 seeds), *agl36-1* N= 12 BR (534 seeds), *agl36-1* N= 12 BR (635 seeds), *agl62-1* N= 8 BR (403 seeds), *agl90-1* N= 12 BR (514 seeds), *agl90-2* N= 12 BR (635 seeds). BR, biological replicas.

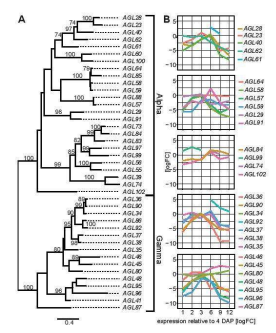


Figure 1: MADS-box type I transcription factors share similar expression profiles during seed development. A) Maximum likelihood phylogeny of alpha and gamma MADS-box type I genes in Arabidopsis thaliana. The tree was inferred using the GTRGAMMA model on 41 genes with 532 unambiguously aligned nucleotides. Scale bar represents the mean number of nucleotide substitutions per site. Only bootstrap values above 65% are shown. B) Gene expression profiles of alpha and gamma MADS-box type I genes were ordered in five groups according to the branching pattern. Transcript quantification and differential expression analysis was performed with RSEM and visualized using R. Gene expression profiles for stages ranging from one to 12 days after pollination (DAP) are relative to four DAP using a base-2 logarithmic scale (logFC). Two biological replicas with three technical replicas were analysed. Note that genes within groups show similar gene expression profiles, with a common maximum reached between three and six DAP.

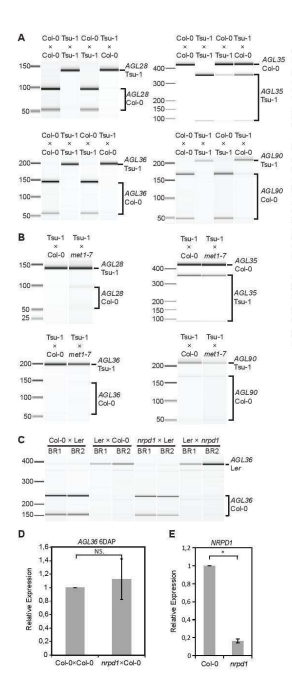


Figure 2: Imprinting and epigenetic regulation of AGL28, AGL35, AGL36 and AGL90. A) Imprinting analysis of AGL28, AGL35, AGL36 and AGL90 using ecotype specific restriction digest on single nucleotide polymorphisms (SNPs) in reciprocal crosses between ecotypes Tsu-1 and Col-0. For each panel, the ecotype specific digestion pattern is indicated. Seeds were harvested for analysis four days after pollination (DAP). Bioanalyzer images of one of three biological replicas is shown. B) SNP analysis of AGL28, AGL35, AGL36 and AGL90 in crosses with met1-7+/ (Col-0 background) pollen. Only AGL28 display paternal activation in the met1-7+1mutant. Wild type (WT) crosses are duplicated from A) for visualization. Crosses were harvested as A). Bioanalyzer images of one of three biological replicas is shown. C) SNP analysis of AGL36 in reciprocal crosses between WT (Col-0) or nrpd1 (Col-0) and WT (Ler). Crosses were harvested six DAP. The AGL36 imprinting pattern is not changed in the nrpd1 crosses compared to WT crosses. D) Real-time PCR analysis of AGL36 expression in three biological replicas the nrpd1 background at six DAP. The relative expression difference is not significant (NS, t-test; p=0,389). E) Real-time PCR verifying significant knock-down (t-test; p=0,015, indicated by asterisk) of NRPD1 in three biological replicas of nrpd1 homozygous background.

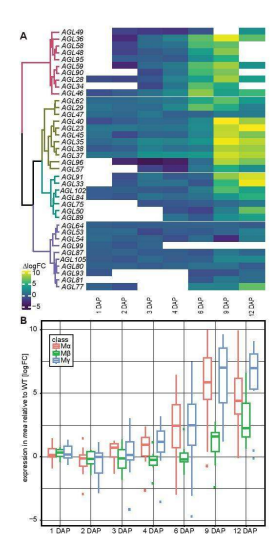
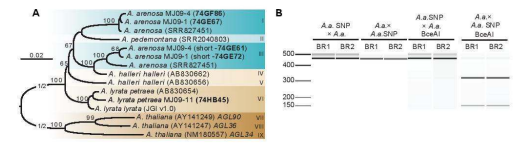


Figure 3: PRC2 dependent transcriptional repression of MADS-box type I transcription factors during seed development. A) Heat-map clustering of gene expression profiles of MADS-box type I transcription factors (TFs) in the Polycomb Repressive Complex 2 (PRC2) MEDEA mutant seeds (mea) compared to wild type (WT). Expression profile based clustering, transcript quantification and differential expression analysis was performed for three biological replicas using RSEM and visualized using R. Differential gene expression profiles for stages ranging from one to twelve days after pollination (DAP) are shown using a base-2 logarithmic scale (logFC). Note strong expression change in three out of four expression clusters. B) Box-plot showing class specific expression of MADS-box type I TFs at stages from one to twelve DAP. The MEA dependent repression is shared by the alpha (M α) and gamma (M γ) class TFs, while the beta (Mβ) class is weakly affected in the mea background. Relative expression changes are displayed in a base-2 logarithmic scale.



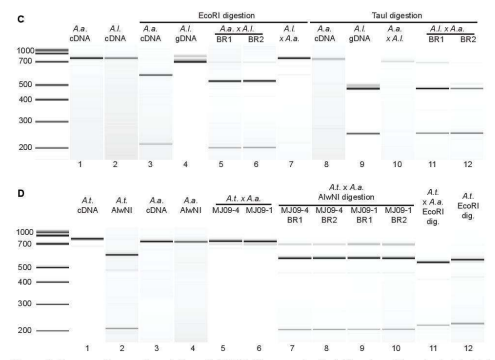


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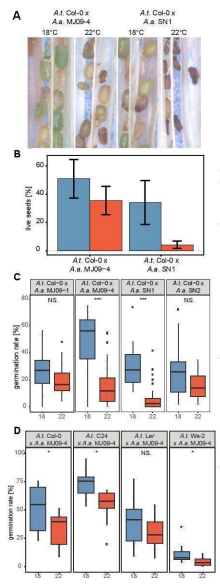


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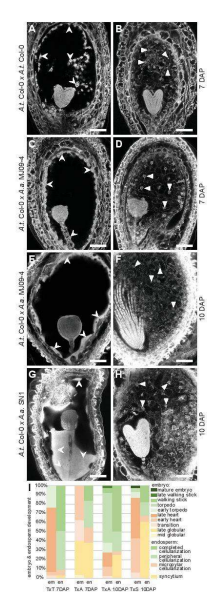


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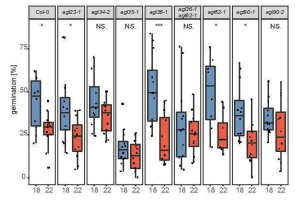


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