

1 *DNA METHYLTRANSFERASE 3 (MET3) is regulated by Polycomb Group*
2 *complex during Arabidopsis endosperm development*

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6

7 **Key message**

8 ***The DNA METHYLTRANSFERASE MET3 is controlled by Polycomb group complex during***
9 ***endosperm development.***

10 **Abstract**

11 Complex epigenetic changes occur during plant reproduction. These regulations ensure the proper
12 transmission of epigenetic information as well as allowing for zygotic totipotency. In *Arabidopsis*, the
13 main DNA methyltransferase is called MET1 and is responsible for methylating cytosine in the CG
14 context. The *Arabidopsis* genome encodes for three additional reproduction-specific homologs of
15 MET1, namely MET2a, MET2b and MET3. In this paper, we show that the DNA methyltransferase
16 MET3 is expressed in the seed endosperm and its expression is further restricted to the chalazal
17 endosperm. MET3 is biallelically expressed in the endosperm but displays a paternal expression bias.
18 We found that MET3 expression is regulated by the Polycomb complex proteins FIE and MSI1. Seed
19 development is not impaired in *met3* mutant, and we could not observe significant transcriptional
20 changes in *met3* mutant. Interestingly, we found that MET3 regulates gene expression in a Polycomb
21 mutant background suggesting a further complexification of the interplay between H3K27me3 and
22 DNA methylation in the seed endosperm.

23 **Keywords**

24 DNA methylation, *Arabidopsis*, endosperm, Polycomb, MET3

25

26 *Introduction*

27 Sexual reproduction in Angiosperm is initiated by a double fertilization event. Fertilization of the
28 haploid egg cell by one of the sperm cells gives rise to the diploid embryo whereas fertilization of the
29 homodiploid central cell gives rise to the triploid endosperm (Berger 2003; Costa et al. 2004). The
30 endosperm represents a nourishing tissue supporting embryo growth and is therefore key for proper
31 seed development. Cell divisions in the endosperm are initiated very rapidly following fertilization.
32 These divisions are initially occurring without cellularization and form a syncytium that will later
33 cellularize (Brown et al. 1999, 2003; Boissard-Lorig et al. 2001). An additional peculiarity of the
34 endosperm, beyond its triploid syncytial nature, is being the seat of interesting epigenetic phenomena
35 and complex epigenetic regulation.

36 Endosperm development and cellularization are indeed regulated by the FIS Polycomb group
37 complex known to mediate Histone H3 Lysine 27 tri-methylation, a key silencing epigenetic mark.
38 Some of the key members of the FIS Polycomb group complex (FIS-PcG) are MEA, FIS2, FIE, and MSI1
39 (Chaudhury et al. 1997; Luo et al. 1999; Kiyosue et al. 1999; Yadegari et al. 2000; Köhler et al. 2003;
40 Guittou et al. 2004). In mutants affecting those genes, the endosperm fails to cellularize, resulting in
41 an arrest of embryo development and eventually seed abortion. Additionally, several genes were
42 found to be imprinted, *i.e.* only one parental allele is expressed whereas the other is epigenetically
43 silent, in the endosperm. This is the case, for example, of genes such as *FIS2*, *FWA*, *MEA* or *PHE1*
44 (Kinoshita et al. 1999, 2004; Luo et al. 2000; Köhler et al. 2005; Jullien et al. 2006). The silencing of
45 those endosperm imprinted genes relies principally on two epigenetic mechanisms: either regulation
46 by the FIS-PcG itself like for *MEA* or *PHE1* or silencing by DNA methylation like for *FWA* and *FIS2* (Jullien
47 and Berger 2009; Gehring 2013; Batista and Köhler 2020). Another epigenetic singularity of the
48 endosperm is of being relatively hypomethylated compared to other plant tissues (Gehring et al. 2009;
49 Hsieh et al. 2009). This hypomethylation is due in part to the activity of a DNA glycosylase called
50 DEMETER (Choi et al. 2002; Gehring et al. 2006; Hsieh et al. 2009) and likely also to the low expression
51 of canonical actors of the DNA methylation pathway (Jullien et al. 2012).

52 DNA methylation is a key epigenetic mark regulating gene expression and protecting genome
53 integrity by repressing transposons. In plant genomes, DNA methylation is found in three cytosine
54 contexts: CG, CHG and CHH (where H is any base except C). Methylation on these different contexts
55 relies on specific DNA methyltransferases. DNA methylation on CG sites relies on maintenance DNA
56 METHYLTRANSFERASE (MET) where the main ubiquitous enzyme is MET1. DNA methylation on CHG
57 sites relies on CHROMOMETHYLASE3 (CMT3) and an interplay with histone methylation (Lindroth et
58 al. 2001). DNA methylation on CHH site, due to its non-symmetrical nature, relies on the constant *de*
59 *novo* methylation pathway involving small RNA molecules as well as DOMAIN REARRANGED
60 METHYLTRANSFERASE2 (DRM2) (Cao and Jacobsen 2002). In centromeric sequences, CHH
61 methylation also relies on CHROMOMETHYLASE2 (CMT2) (Stroud et al. 2013).

62 Although we know a lot about the main actors of this pathway, the *Arabidopsis*'s genome encodes
63 multiple copies of DNA methyltransferase genes (4 *METs*, 3 *CMTs* and 3 *DRMs*) some of which might
64 have a more complex or similar function in discreet cell types. For example, *CMT1* (the third
65 CHROMOMETHYLASE encoded by the *Arabidopsis* genome) is principally detected in reproductive
66 tissue (Henikoff and Comai 1998; Klepikova et al. 2016) and the reconstitution of a full-length
67 transcript relies on the splicing out of a transposable element situated in its 13th exon (Yadav et al.
68 2018). The *DOMAIN REARRANGED METHYLTRANSFERASE1 (DRM1)* seems to also solely play a role in
69 reproductive tissue, where a redundancy between *DRM1* and *DRM2* was observed in the early embryo
70 (Jullien et al. 2012). Similarly, data concerning the potential function of non-canonical *METs* are scarce.
71 *MET2a* and *MET2b* are detected in the central cell, but their function is unknown (Jullien et al. 2012).
72 Nonetheless, correlative evidence suggest *MET2a* might be important to regulate transposon
73 reactivation in wild *Arabidopsis* accessions (Quadrana et al. 2016) and involved in fungal response
74 (Salvador-Guirao et al. 2018).

75 As mentioned, little is known about the DNA methyltransferase *MET3*. *MET3* is also named
76 *MATERNAL EFFECT EMBRYO ARREST 57 (MEE57)* as a transposon insertion associated with the *MET3*

77 locus led to an arrest in endosperm development (Pagnussat et al. 2005). *MET3* is also reported to be
78 the sole *MET* expressed in the endosperm (Jullien et al. 2012). Here, we show that *MET3* is specifically
79 expressed in the endosperm in a biallelic fashion with a paternal bias. *MET3* expression is controlled
80 by the FIS-PcG complex. Despite the initial report of a seed arrest phenotype in the *mee57* line, we
81 did not see any seed developmental phenotype in two independent *met3* mutant alleles. Additionally,
82 we could not see major changes in the seed transcriptome of *met3* mutant. Nevertheless, we could
83 see an effect on the seed transcriptome in a *fie* mutant background suggesting that *MET3* might
84 interplay with PcG gene regulation in the developing endosperm.

85 *Material and Methods*

86 *Plant Materials, Growth Conditions and Genotyping.*

87 The wild-types Col-0 and Gr-1, the *MET3* mutant lines *met3-3* (GABI404F04), *met3-4* (GABI659H03)
88 and PRC2 mutant lines *fie-362* (GABI_362D08) (Bouyer et al. 2011) and *msi1* (SAIL_429_B08) were
89 provided by the Nottingham Arabidopsis Stock Center (NASC). *pMET3:H2B-RFP* line was previously
90 described (Jullien et al. 2012). After sowing, plants were stratified in the dark at 4°C between 2 and 4
91 days. Plants were germinated and grown in growing chambers under long-day conditions (16h light
92 22°C / 8h dark 18°C). For the transmission analysis (Fig. S5a), plants were grown on Murashige and
93 Skoog (MS1/2) media agarose plate in long-day conditions for 12 days before genotyping. Primers for
94 genotyping are listed in Table S1.

95 *Microscopy and phenotype observation*

96 DIC seed phenotype and GUS observations were done using a Leica DM2000 as previously described
97 (Jullien et al. 2006). For seed development observation and counting (Fig. 4a) , plants were
98 synchronized by removing all open flowers from the inflorescences. After 6 days, the two first siliques
99 situated above the previously removed flowers were picked for each inflorescence. We refer to this
100 stage as 6 days after synchronization (6 DAS). Siliques were dissected, cleared using chloride hydrate

101 solution and mounted on a slide for observation. *pMET3:H2B-tdTomato* and *pMET3:H2B-RFP* reporter
102 lines were imaged using a laser scanning confocal microscope (Leica SP5). When necessary, brightness
103 and contrast were uniformly modified by using ImageJ.

104 *Cloning and transformation*

105 *pMET3:H2B-tdTomato* and *pMET3:H2B-GUS* were generated using the Gateway Cloning System
106 (Invitrogen). All PCR fragments were amplified by PCR using the Phusion High-Fidelity DNA Polymerase
107 (Thermo). Primer sequences used for cloning can be found in Table S1. All plasmids were transformed
108 into wild-type Columbia-0 plants the by floral dipping method (Clough and Bent 1998). At least ten
109 transgenic lines were analyzed per construct, which showed a consistent fluorescence expression
110 pattern. An Illustration of the constructs can be found in Fig. S2a.

111 *RNA extraction, qPCR & RT-PCR*

112 Total RNAs were extracted using RNeasy Plant Minikit (Qiagen). All samples were treated with DNase
113 I (ThermoScientific) at 37°C for 30 minutes. DNase I was subsequently inactivated by the addition of
114 EDTA and heat treatment (65°C for 10 minutes). First-strand cDNAs were synthesized using between
115 500 and 1000 ng of DNase treated total RNAs as a template. The RT reaction was performed using
116 either Maxima First Strand cDNA Synthesis Kit (Fig. 1a, Fig. 2a) (ThermoScientific) , containing both
117 oligo-dT and random hexamer primers, or RevertAid RT Reverse Transcription Kit (Fig. 3e, Fig. S4b)
118 (ThermoScientific) , containing only oligo-dT primers. The qPCR reactions were performed with a
119 QuantStudio 5 thermocycler (ThermoScientific) using SYBR green (KAPA SYBR FAST qPCR Master Mix
120 or ORA qPCR Green ROX H Mix). The qPCR mix was prepared according to the manufacturer's protocol.
121 An RNA equivalent of 25 ng of cDNA templates was used for each reaction. The qPCR program was as
122 follow: 95 °C for 3 minutes followed by 45 cycles of 95 °C for 5 seconds and 60 °C for 30 seconds.
123 *ACTIN2 (AT3G18780)* expression was used to normalize the transcript level in each sample. For each
124 condition, RNA abundance of target genes was calculated from the average of three independent
125 biological replicates with three qPCR technical replicates. Real-time PCR primers used in this study are

126 listed in Table S1. For the allele-specific RT-PCR (Fig.2a) , cDNAs were amplified for 22 cycles for *ACT2*
127 primers and 35 cycles for *MET3* specific primers. Half of the *MET3* PCR product was digested for 1h30
128 at 37°C with XbaI restriction enzyme (Sigma-Aldrich). *ACT2* amplification was used as a control.

129 *RNA sequencing and Bioinformatics*

130 Total RNAs were extracted and DNaseI treated as previously mentioned. mRNA libraries were
131 prepared and sequenced by Novogene (<https://en.novogene.com/>). Bioinformatic analyses were
132 performed on the Galaxy web platform (<https://usegalaxy.org>) (Afgan et al. 2018). Our Galaxy
133 workflow including the exact parameters and tool versions used can be downloaded on
134 <https://usegalaxy.org/u/pej/w/pejrnaseq> and be freely reused. Briefly, Paired-end raw mRNA
135 sequencing reads were controlled using FastQC (Galaxy version 0.72) and trimmed using Trimmomatic
136 (Galaxy version 0.36.6) (Andrews 2010; Bolger et al. 2014). Clean reads were aligned to the *Arabidopsis*
137 *thaliana* TAIR 10 genome assembly using HISAT2 (Galaxy version 2.1.0+galaxy4) (Kim et al. 2015).
138 Aligned sequencing reads were assigned to genomic features using featureCounts (Galaxy version
139 1.6.2) (Liao et al. 2014). Differential expression was analyzed using DESeq2 default parameters (Galaxy
140 Version 2.11.40.6+galaxy1) (Love et al. 2014). Differentially expressed genes (DEGs) were defined by
141 an absolute logFC > 2 and an FDR < 0.05. Gene ontology (GO) enrichment analysis has been performed
142 on PANTHER (Mi et al. 2019) and visualized using REVIGO (Supek et al. 2011). All plots have been
143 generated using R-studio (www.rstudio.com). Raw data are deposited on the European Nucleotide
144 Archive under reference PRJEB46544 (<http://www.ebi.ac.uk/ena/data/view/PRJEB46544>).

145 *Results*

146 *MET3 is expressed biallelically with a paternal bias in the endosperm*

147 Our previous analysis has shown that *MET3* is expressed in the endosperm of developing
148 *Arabidopsis* seeds (Jullien et al. 2012). However, the detail and exclusivity of its expression pattern
149 remain to be investigated. To get a better characterization of *MET3* expression pattern, we performed

150 a qPCR of *MET3* transcript in different tissue types of wild-type Col-0 (Fig. 1a). Our result shows that
151 *MET3* is principally expressed in siliques and its expression peaks at 5 Days After Pollination (DAP).
152 These results could be confirmed using publicly available transcriptome datasets (Fig. S1a) (Klepikova
153 et al. 2016). To gain a better spatial and temporal characterization of *MET3* expression, we generated
154 two new transcriptional *MET3* reporter constructs, encompassing 2kb of the *MET3* promoter driving
155 either H2B-tdTomato or H2B-GUS (Fig. S2a). The analysis of the *pMET3:H2B-GUS* in different plant
156 tissues confirmed the specificity of *MET3* expression to the seed endosperm (Fig. S1b-d). To
157 characterize in detail the temporal expression of *MET3* in the endosperm, we performed confocal
158 microscopy on the *pMET3:H2B-tdTomato* lines at different stages of seed development. We could
159 detect *pMET3:H2B-tdTomato* expression from as early as the four nuclei stage endosperm (Fig. 1b).
160 *pMET3:H2B-tdTomato* remains express throughout the endosperm (Fig. 1c) until the globular stage of
161 embryo development where its expression starts to be higher in the chalazal endosperm and chalazal
162 cyst (Fig. 1d). At later stages, *pMET3:H2B-tdTomato* expression is limited to the chalazal endosperm
163 and chalazal cyst (Fig. 1e). From 7 DAP, *pMET3:H2B-tdTomato* expression can no longer be detected.
164 A similar expression pattern was observed with *pMET3:H2B-GUS* (Fig. S2b) and *pMET3:H2B-RFP* (Fig.
165 S2c) as well as online transcriptomic data (Fig. S2d) (Belmonte et al. 2013). *MET3* protein expression
166 and localization could not be determined as we, so far, failed in expressing a fluorescently tagged
167 *MET3* protein in *Arabidopsis* (LT personal communication).

168 Such endosperm expression pattern is common in imprinted genes, like *FWA*, *FIS2*, *MEA* or *PHE1*
169 (Kinoshita et al. 1999, 2004; Köhler et al. 2005; Jullien et al. 2006). In order to examine if *MET3* is
170 biallelically or mono-allelically expressed, we performed allele-specific RT-PCR. We are making use of
171 a Short Nucleotide Polymorphism (SNP) consisting of a substitution from a C to a T within *MET3* 9th
172 exon in the Gr-1 ecotype which is abolishing a XbaI restriction site present in Col-0. We did reciprocal
173 crosses using Col-0 and Gr-1 ecotypes and analyzed *MET3* parental expression at 5 DAP following XbaI
174 digestion (Fig. 2a). We could observe, for both reciprocal crosses, bands corresponding to *MET3*
175 transcript from Col-0 (505 and 329 bp) and from Gr-1 (834 bp) with a bias toward the paternal allele.

176 This result shows that *MET3* is expressed from both maternal and paternal allele but displays a
177 paternal bias of expression. *MET3* paternally biased expression was also observed using the
178 *pMET3:H2B-RFP* transgene at 1 DAP (Fig.2b). Taking together our results shows that *MET3* expression
179 is biallelic with a paternal bias and confined to the endosperm. *MET3* expression, initially throughout
180 the endosperm, becomes restricted to the chalazal pole at later stages.

181 *MET3 expression is regulated by Polycomb group proteins*

182 Beyond imprinted genes, *MET3* expression pattern is reminiscent of the expression of genes
183 controlled by the endosperm FIS Polycomb group complex (PcG) composed of FIE, MSI1, FIS2 and MEA
184 (Guitton et al. 2004). To investigate if *MET3* could be regulated by PcG, we introgressed a *MET3*
185 transcriptional reporter into *fie* and *msi1* mutant background (Fig. 3a-d). We could observe increased
186 *pMET3:H2B-RFP* reporter expression in 49% of the seeds (n=249) in *fie/+* mutant and 44% (n=240) in
187 *msi1/+* mutant background characteristic of the maternal gametophytic effect of those mutations (Fig.
188 3d). In *msi1* and *fie* mutants, *pMET3:H2B-RFP* expression is higher and observed throughout the
189 endosperm (Fig. 3b-c) whereas at the same developmental stage the expression of *pMET3:H2B-RFP* is
190 already restricted to the chalazal pole in wild-type seeds (Fig. 3a). In order to confirm that the
191 regulation of *MET3* by *MSI1* and *FIE* was not only restricted to our transgene, we performed a RT-qPCR
192 of *MET3* expression in wild-type and mutant selected seeds at 10 DAP (Fig. 3e). We could observe a
193 clear upregulation of *MET3* expression in both *fie* and *msi1* seeds compare to wild-type seeds (t-test
194 p-value of 0.0543 and 0.0114 respectively). Using publicly available data, we could see that the
195 upregulation of *MET3* is also observed in other PcG mutants (Fig. S3a-b). *MET3* upregulation was
196 observed in silique samples of *clf* mutant (Fig. S3a) (Liu et al. 2016) and in *fis2* seeds (Fig. S3b)
197 (Weinhofer et al. 2010).

198 Subsequently, we wanted to know if the effect of the PcG complex was direct or indirect. PcG
199 complexes are known to repress gene expression by tri-methylating the Lysine 27 of the Histone H3
200 tail (H3K27me3) inducing a closed chromatin state at the targeted loci. We, therefore, analyzed

201 available H3K27me3 genome-wide Chromatin Immunoprecipitation (ChIP) data. We could see that
202 the *MET3* locus is covered by H3K27me3 in *Arabidopsis* seedling samples (Fig. 3f) (Zhang et al. 2007).
203 Additionally, using the RepMap2020 tool (Chèneby et al. 2020) , we could observe H3K27me3 on the
204 *MET3* locus in several independent ChIP experiments including some performed on endosperm tissue
205 (Fig. S3c). We conclude that *MET3* expression is directly repressed by PcG complex induced H3K27me3
206 in the endosperm.

207 *MET3 does not affect seed development*

208 Considering that *MET3* is specifically expressed in the developing endosperm during seed
209 development, we then ask if *MET3* function influences seed and/or endosperm development. We
210 characterized two mutant alleles from the GABI collection: *met3-3* (GABI_404F04) and *met3-4*
211 (GABI_659H03) (Fig. S4a). The mutations are located on the 10th and the 2nd exon respectively and are
212 expected to abolish *MET3* function. To confirm the downregulation of *MET3* in the mutants, we
213 performed RT-qPCR. We could observe that *MET3* is downregulated in both mutant alleles (Fig. S4b).
214 To investigate if the *met3* mutation could result in seed lethality we first investigated the presence of
215 aborted seeds at the green seed stage (~12DAP). We could not see any significant seed abortion in
216 both *met3-3* and *met3-4* alleles (Fig. S5b). To further confirm the absence of defects, we analyzed the
217 transmission rate of the *met3* mutations in *met3-3/+* and *met3-4/+* selfed progeny. We could not
218 observe any segregation distortion from the Mendelian ratio (Fig. S5a). Mutations affecting the main
219 *Arabidopsis* DNA methyltransferase *MET1* display variation in seed size. We, therefore, investigated if
220 *met3* mutants would display a seed size phenotype. We could not see any significant variation in seed
221 size using both *met3* alleles (Fig. S5c-d). We conclude that *MET3* mutation does not severely impair
222 seed development. Additionally, *met1* mutant are known to display increased developmental defect
223 through generation (Mathieu et al. 2007). To assess potential transgenerational effect of the *met3*
224 mutation we have maintained *met3* homozygotes mutants for five generations. However, we could
225 not observe such increased developmental phenotype with *met3* mutants after five generation of
226 inbreeding (*met3*^{G5}) (Fig. S5e-g). The *met3*^{G5} did not show difference when compared to wild-type

227 either in term of rosette size (Fig. S5f) nor flowering time (Fig. S5g). We conclude that *MET3* mutations
228 do not severely impair seed development and do not accumulate transgenerational developmental
229 defects.

230 As shown above, *MET3* expression is regulated by PcG complex in the endosperm, we then
231 analyzed if *MET3* mutation could influence the PcG *fie* mutant phenotype. To answer this question,
232 we generated *fie/+;met3-3* and *fie/+;met3-4* double mutants and analyzed their seed development
233 phenotype using DIC (Fig. 4a-f). In order to minimize the stress to the plant due to handling during
234 emasculatation and crossing, we used “synchronized seeds”. In practice, we remove the open flowers
235 of the day, and we wait an x number of days before collecting two siliques above our cutting. This is
236 allowing us to have age synchronized siliques without the physical disturbance of
237 emasculatation/pollination and is, therefore, closer to normal growth and fertilization. We are using the
238 term Day After Synchronization (DAS). To our experience, 6 DAS is corresponding to around 4-5DAP.
239 At 6 DAS, we could not see any delay in embryo development and endosperm cellularization
240 comparing Col-0 to *met3-3* and *met3-4* mutants (Fig. 4a-c, 4g-h). In *fie/+*, we could clearly see the
241 delayed endosperm cellularization characteristic of FIS-PcG mutants (Fig. 4d-f and h) (56% n=407)
242 (Ohad et al. 1999; Sørensen et al. 2001). In the double mutants, we could not see variations in the
243 quantification of the *fie/+* phenotype for both embryo development (Fig. 4g) and endosperm
244 cellularization (Fig. 4h). We conclude that *MET3* mutations do not influence *fie* mutant seed
245 phenotype.

246 *Effect of MET3 mutation on the seed transcriptome*

247 In order to investigate the potential effect of *met3* mutation on the seed transcriptome, we
248 performed a mRNA deep sequencing experiment of 3 DAP (Fig. S6) and 10 DAS (Fig. 5a) seeds (*i.e.*
249 seeds attached to the septum). We then compared the seed transcriptome of *met3* seeds to wild-type
250 seeds. At 3 DAP, we could only detect one differentially expressed gene: *ESM1* (*AT3G14210*). *ESM1*
251 was up-regulated in both *met3-3* and *met3-4* mutant seeds (Fig. S6a-b). At 10 DAS, we could also only

252 see very minor changes to the seed transcriptome (Fig. 5a). We could identify only 16 differentially
253 expressed genes with an absolute $\log_{2}FC > 2$ and $FDR < 0.05$. These results show that reminiscent of the
254 absence of seed phenotype in *met3* mutants, the seed transcriptome is also mostly unaffected by the
255 *met3* mutation. As *MET3* expression is regulated by the FIS-PcG complex in the endosperm, we then
256 ask if *MET3* mutation could influence the *fie* transcriptome. We, therefore, sequenced the
257 transcriptome of 10 DAS seeds where we selected for *fie* mutant seeds under the dissecting
258 microscope (white seeds). We compared the transcriptome of *fie* seeds to the transcriptome of
259 *fie;met3-3* double mutant seeds. We could observe 87 differentially expressed genes (DEG) with an
260 absolute $\log_{2}FC > 2$ and $FDR < 0.05$ (Fig 5b). A Goterms enrichment analysis revealed that these genes are
261 enriched for genes involved in pectin metabolism (GO:0045490, $FDR = 4.46E-10$; GO:0045488,
262 $FDR = 6.59E-09$) and cell-wall related processes (GO:0042545, $FDR = 8.97E-09$; GO:0071555, $FDR = 4.87E-$
263 10 , GO:0071554, $FDR = 5.67E-08$) (Fig. 5c and Table S2). We then analyzed if among these 87 DEGs
264 some are also modified in *fie* mutant compared to wild-type. We could find that a large proportion of
265 either *met3* DEGs (Col vs *met3*, 11/16) and *met3;fie* DEGs (*fie* versus *met3fie*, 44/87) are miss-
266 regulated in *fie* mutant seeds (Fig. 5d). We conclude that the *met3* mutation does not drastically
267 change the seed transcriptome, but a set of genes are miss-regulated by both *met3* and *fie* mutants.

268 *Discussion*

269 Our study highlights an additional connection between DNA Methylation pathways and Polycomb
270 group H3K27 tri-methylation in the seed endosperm via the regulation of *MET3* by the FIS-PcG
271 complex. *MET3* is specifically expressed in the endosperm, and its expression becomes restricted to
272 the chalazal pole at later stages of endosperm development. In our study, using two independent
273 insertion lines, *met3-3* and *met3-4*, we could not observe any major seed developmental phenotype.
274 Both mutants show a significant decrease in *MET3* expression by qPCR suggesting they both represent
275 knockout mutants. It was previously documented that a mutation affecting *MET3* named *mee57*
276 displayed a strong seed developmental phenotype (Pagnussat et al. 2005). The *mee57* mutation shows

277 an early maternal embryo and endosperm arrest. The discrepancy between our lines and the
278 previously published *mee57* could have several causes: the mutagenesis method used (T-DNA versus
279 transposition) , the presence of additional genetic modifications, or a difference between the two
280 ecotypes used, Columbia-0 in our case and Landsberg for *mee57*. If the latter is true, *MET3* function
281 could vary between different *Arabidopsis* accessions.

282 In this work, we show that repression of *MET3* expression at later stages of endosperm
283 development is linked to the direct action of the FIS-PcG complex on the *MET3* locus. *MET3* is over-
284 expressed in PcG mutant seeds, such as *fie* and *msi1* mutant seeds. Interestingly, it was previously
285 shown that *fie* mutant endosperm display higher CG methylation compared to wild-type endosperm
286 and lower CHG and CHH (Ibarra et al. 2012). This increased CG methylation in *fie* mutant seems to be
287 restricted to the endosperm as it is not observed in *fie* seedling or *fie* embryo methylome (Bouyer et
288 al. 2017). We propose that higher expression of *MET3* in the *fie* endosperm could be the cause for the
289 increase in CG methylation specifically in the endosperm. Indeed, in our study, we could not observe
290 any change in *MET3* tissue expression pattern in PcG mutants but the increased expression is still
291 restricted to the endosperm. In further studies, the investigation of the endosperm methylome in
292 *met3* and *met3;fie* mutants will allow to test this hypothesis. Additionally, further experiments are
293 required to determine if *MET3* is a functional DNA methyltransferase.

294 As we previously mentioned, *MET3* function could vary in between wild *Arabidopsis* accessions.
295 Similarly, to *CMT1*, several SNPs can be found at the *MET3* locus in-between different ecotypes
296 suggesting that *MET3* might not be fully functional in all of them. Additionally, looking closer at the
297 *MET3* locus, we could detect the presence of a Class 2 DNA transposon (*AT4TE34810*) situated in the
298 third intron of the *MET3* gene. In addition to PcG regulation, *MET3* expression and possibly function
299 could be influenced by the presence or regulation of *AT4TE34810* like it is the case for *CMT1* transcript
300 (Yadav et al. 2018). The study of *MET3* function and imprint in different accession could reveal more
301 about its function in natural growth conditions. One hint of *MET3* potential function in Columbia-0

302 came from the study of methylome stability across several generations (Becker et al. 2011). In this
303 study, methylome stability was investigated in several Col-0 selfed lineages (30 generations). The line
304 accumulating the most methylation polymorphisms had concomitantly acquired a SNP in the *MET3*
305 gene. It was therefore suggested that *MET3* could be the cause of such methylome instability (Becker
306 et al. 2011; Schmitz and Ecker 2012). Analyzing the methylome of selfed *met3* mutants after 30
307 generations would allow to test this hypothesis. If true and taking into account the endosperm
308 specificity of *MET3* expression, it would indicate that the endosperm methylome influences the
309 embryonic methylome, a hypothesis that has stimulated a lot of interest over the last 10 years but
310 remains to be demonstrated.

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317 *Author contributions*

318 PEJ conceived the study. LT performed the experiments. LT and PEJ analyzed the data. PEJ wrote
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320 *Conflicts of interest*

321 Authors state that there is no conflict of interest.

322

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- 465
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467 **Figure legends**

468 **Figure 1. *MET3* expression pattern**

469 (a) *MET3* expression measured by RT-qPCR. DAP = Day After Pollination. The histogram displays the
470 mean, and each dot represents a biological replicate. *ACT2* is used as a normalizer. RQ = Relative
471 Quantification (b-e) Single-plan Confocal images representing the expression of *pMET3:H2B-*
472 *tdTomato* construct in *Arabidopsis* 1 Day After Synchronization (DAS) seed (b), 2 DAS seed (c), 3 DAS
473 (d) and 5 DAS (e). cze = chalazal endosperm, endo = endosperm nuclei. Scale bars represent 25 μ m.

474 **Figure 2. *MET3* is biallelically expressed with a paternal bias**

475 (a) Allele specific RT-PCR of *MET3* parental expression in 5DAP silique samples. The *Xba*I restriction
476 enzyme digests the Col-0 *MET3* transcript but not the Gr-1 transcript. *ACT2* is used as loading control.
477 (b) Single-plan confocal images of *pMET3:H2B-RFP* parental expression in the endosperm of 1 DAP
478 seeds. endo = endosperm nuclei. Scale bars represent 25 μ m. ♂ symbol indicates the genotype of the
479 male parent while ♀ indicates the genotype of the female parent.

480 **Figure 3. *MET3* expression is controlled by *FIE* and *MSI1* Polycomb proteins**

481 (a-c) Confocal images representing the expression of *pMET3:H2B-RFP* construct in *Arabidopsis*
482 wildtype (a), *fie* (b) and *msi1* (c) selfed seeds at 5 days after pollination (DAP). Scale bars represent 50
483 μ m. (d) Proportion of seeds with high RFP signal in the endosperm at 5DAP in wild-type, *fie* and *msi1*
484 mutants. Grey bars represent the seeds with a restricted RFP expression to the chalazal endosperm
485 (as illustrated in (a)). Black bars represent the seeds with ectopic expression of *pMET3:H2B-RFP*
486 throughout the endosperm (as illustrated in (b-c)). (e) *MET3* expression measured by RT-qPCR in 10
487 DAP selected seeds of wild-type, *fie* and *msi1*. The histogram displays the mean, and each dot
488 represents a biological replicate. *ACT2* is used as a normalizer. (f) Snapshot showing that the *MET3*
489 locus contains H3K27me3 and LHP1 but no H3K4me1, H3K4me2, H3K4me3 and H3K9me2 in seedlings.
490 Data from <http://epigenomics.mcdb.ucla.edu/H3K27m3/> (Zhang et al. 2007).

491 **Figure 4. MET3 does not influence fie phenotype**

492 (a-f) Seed developmental phenotype observed after clearing by Difference Interference Contrast (DIC)
493 of wild-type (a), *met3-3* (b), *met3-4* (c), *fie/+* (d), *met3-3;fie/+* (e) and *met3-4;fie/+* (f) at 6 days after
494 synchronization (DAS). Scale bars represent 50 μm . (g-h) Quantification of the embryo developmental
495 stages (g) and endosperm cellularization (h) in wild-type, *met3-3*, *met3-4*, *fie/+*, *met3-3;fie/+* and
496 *met3-4;fie/+* at 6 DAS. mce = micropylar endosperm.

497 **Figure 5. Transcriptome of *met3* and *met3;fie* mutant seeds**

498 (a-b) Volcano plot depicting differentially expressed genes (DEGs) comparing Col-0 to *met3-3* (a) and
499 *fie* to *met3-3;fie* in 10 DAS seeds (b). Red dots represent up-regulated DEGs and blue dots represent
500 down-regulated DEGs. The top 10 DEGs are annotated. We use a threshold of [absolute logFC > 2,
501 FDR < 0.05]. (c) GO term analysis for the 87 DEGs obtained with the *fie* vs *met3-3;fie* contrast. (d)
502 Venn diagram showing the overlapping DEGs between the different contrasts.

503 **Figure S1. MET3 is only expressed in seeds**

504 (a) Snapshot of the *MET3* expression pattern obtained from ebar using the data from Klepikova *et al.*
505 (Winter *et al.* 2007; Klepikova *et al.* 2016)(b-d) The absence of *pMET3:H2B-GUS* expression observed
506 in *Arabidopsis* seedling (b), stamen (c) and ovule (d).

507 **Figure S2. MET3 reporters and expression**

508 (a) Representation of *pMET3:H2B-GUS*, *pMET3:H2B-tdTomato* and *pMET3:H2B-RFP*. (b) *pMET3:H2B-*
509 *GUS* expression observed in 1 DAP and 3 DAP seeds. (c) *pMET3:H2B-RFP* expression observed in 1 DAP
510 and 3 DAP seeds. Scale bars represent 50 μm . (d) Snapshot of expression data from LCM dissected
511 seeds from http://seedgenenetwork.net/plotprobe?name=254720_at (Belmonte *et al.* 2013).

512 **Figure S3. MET3 regulation by Polycomb group**

513 (a) *MET3* expression from an RNA-seq experiment in inflorescence, root, shoot and siliques from wild-
514 type plants and *clf28* mutant plants, data from Liu *et al* 2016 (Liu *et al.* 2016). (b) *MET3* expression

515 from transcriptomic data obtained from 3 DAP seeds using MicroArray comparing wild-type to *fis2-1*
516 endosperm, data from Weinhofer *et al* 2010 (Weinhofer et al. 2010). (c) Snapshot of H3K27me3 ChIP
517 data from several tissues including endosperm (colored in blue), flowers (colored in yellow) and
518 various sporophytic (colored in green). Green line represent the presence of ChIP peaks of H3K27me3.
519 Data from ReMap (Chèneby et al. 2020).

520 **Figure S4. Characterization of MET3 mutants**

521 (a) Representation of the *MET3* locus. Blue triangles are representing the T-DNA insertion site of both
522 *met3-3* (GABI_404F04) and *met3-4* (GABI_659H03) (b) *MET3* expression measured by RT-qPCR in Col-
523 0 and *met3* mutants seeds at 3DAP. The histogram displays the mean, and each dot represents a
524 biological replicate. *ACT2* is used as a normalizer. 2 stars indicate a p value <0.01.

525 **Figure S5. MET3 mutants do not show developmental defect**

526 (a) *met3* mutants transmission in self progeny. +/+ represent wild-type, +/- heterozygotes and -/
527 homozygotes plants for the mutations. (b) Percentage of seed abortion in Col-0 and *met3* mutants
528 (*met3-3* and *met3-4*) siliques. (c) Seed size measurement for Col-0 and *met3* mutants dry seeds. No
529 significant differences observed. (d) Pictures of Col-0 and *met3* mutants dry seeds. The scale bars
530 represent 1 mm. (e) Pictures of Col-0 and *met3* mutants rosette at first (top panel) and fifth generation
531 (bottom panel). The scale bars represent 1 cm. (f) Rosette area measurement in mm² of Col and *met3*
532 mutants at fifth generation. n=12. (g) Flowering time (number of days between transplanting and
533 bolting) of Col and *met3* mutants at fifth generation. n=12.

534

535 **Figure S6. Transcriptome of met3 mutant seeds at 3DAP**

536 (a-b) Volcano plot depicting one differentially expressed gene comparing Col-0 to *met3-3* (a) and
537 *met3-4* (b) at 3DAP.

538

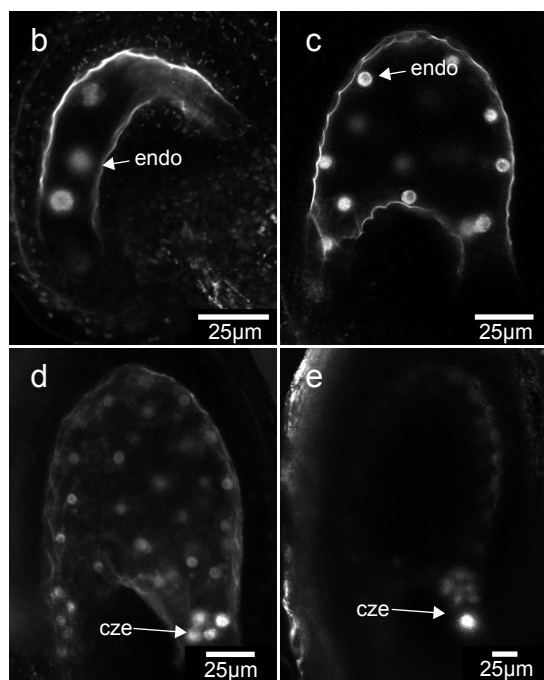
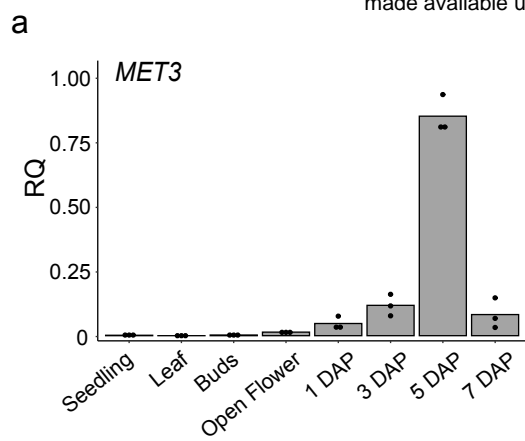


Figure 1. *MET3* expression pattern

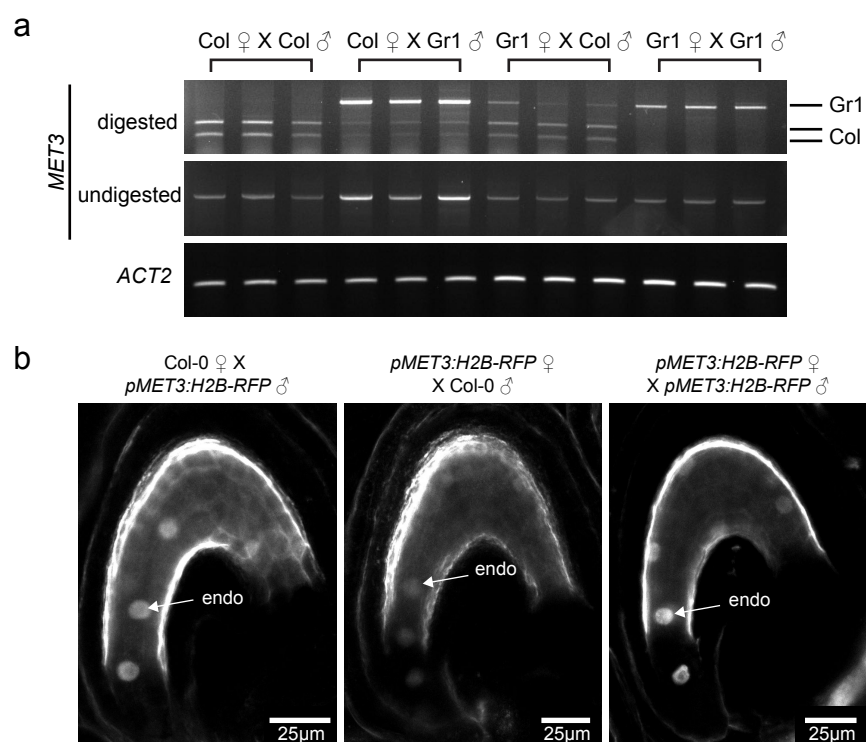


Figure 2. *MET3* is biallelically expressed with a paternal bias

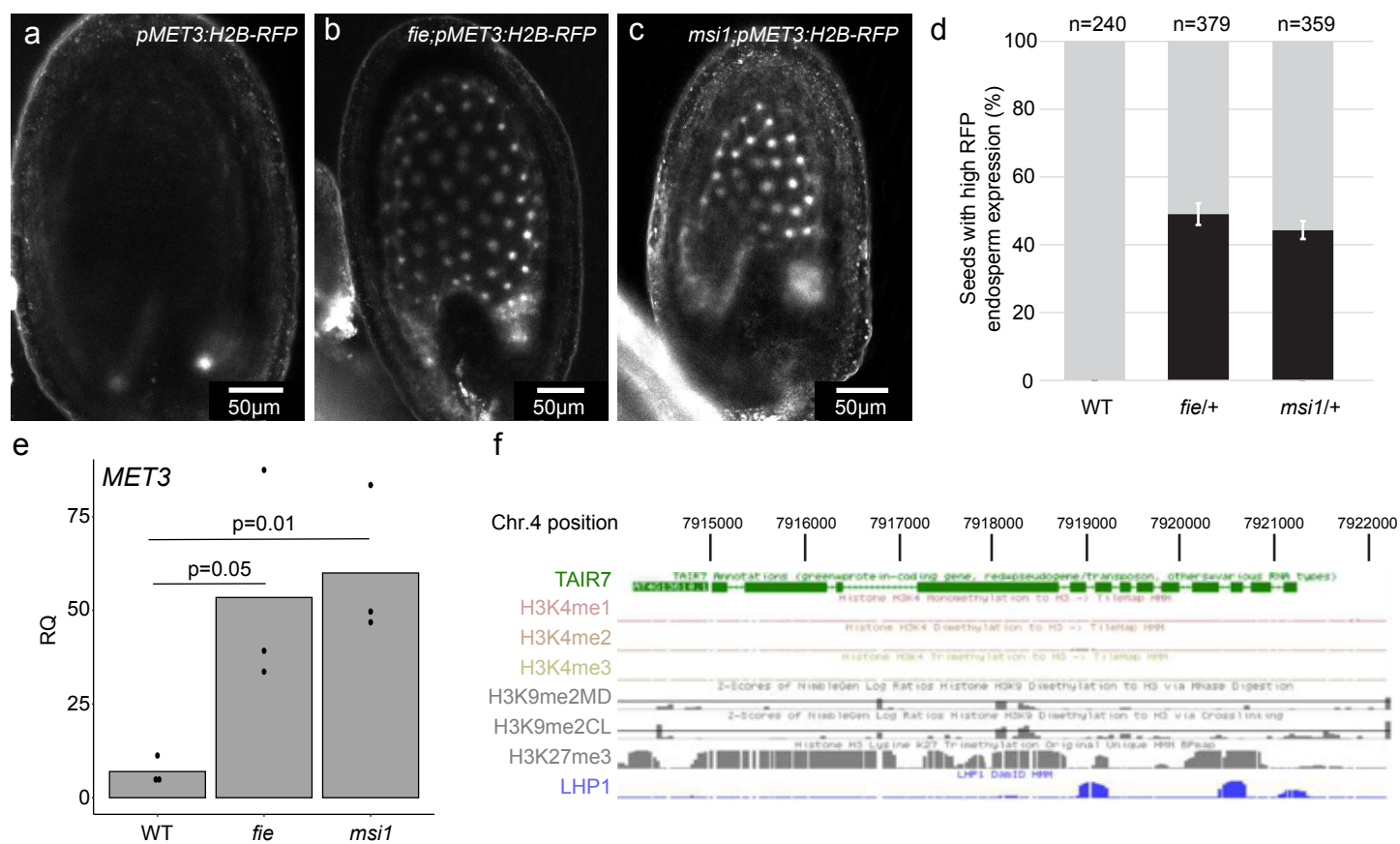


Figure 3. *MET3* expression is controlled by *FIE* and *MSI1* Polycomb proteins

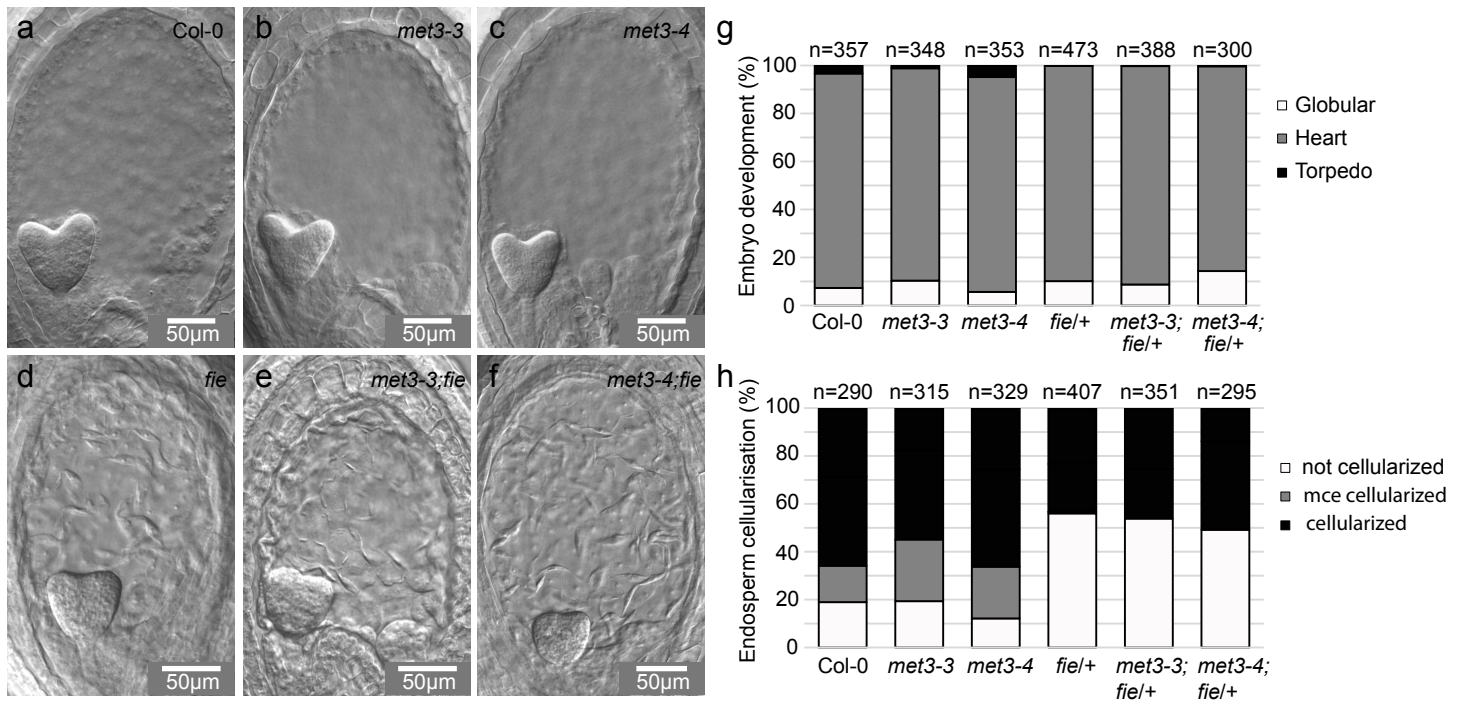


Figure 4. *MET3* does not influence *fie* phenotype

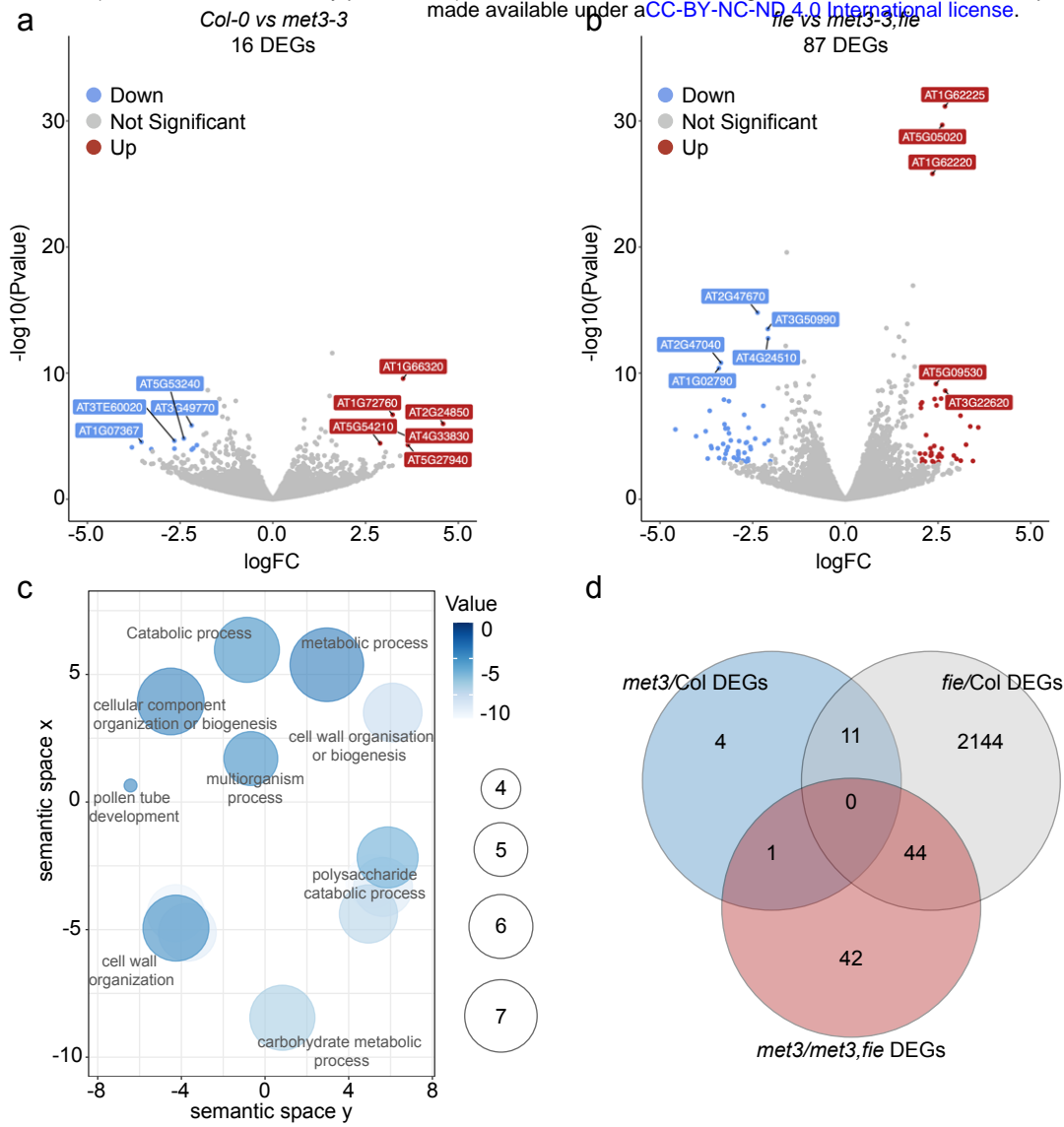


Figure 5. Transcriptome of *met3* and *met3fie* mutant seeds