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1	Stage-specific transcriptomes and DNA methylomes indicate an early and
2	transient loss of transposon control in Arabidopsis shoot stem cells
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16 In contrast to animals, postembryonic development in plants is modular, and aerial 17 organs originate from stem cells in the center of the shoot apical meristem (SAM) throughout life. Descendants of SAM stem cells in the subepidermal layer (L2) will give 18 19 also rise to male and female gametes ¹and therefore can be considered primordial germ 20 cells. In these cells, transmission of somatic mutations including virus and TE insertions 21 must be avoided. Despite their essential role for plant development and intergenerational 22 continuity, a comprehensive molecular analysis of SAM plant stem cells has been missing, 23 due to their low number, deep embedding among non-stem cells and difficult isolation. 24 Here we present a comprehensive analysis of stage-specific gene expression and DNA 25 methylation dynamics in Arabidopsis SAM stem cells. This revealed that stem cell expression signatures are mostly defined by development, but we also identified a core set 26 27 of differentially expressed stemness genes. Surprisingly, vegetative SAM stem cells 28 showed increased expression of transposable elements (TEs) relative to surrounding cells, 29 despite high expression of genes connected to epigenetic silencing. We also find increasing 30 methylation at CHG and a drop in CHH methylation at TEs before stem cells enter the 31 reproductive lineage, indicating an onset of epigenetic reprogramming at an early stage. 32 Transiently elevated TE expression is reminiscent of that in animal primordial germ cells 33 (PGCs)² and demonstrates commonality of transposon biology. Our results connect SAM 34 stem cells with germline development and transposon evolution and will allow future 35 experiments to determine the degree of epigenetic heritability between generations.

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In contrast to short and straight germlines in animals, plants have multiple germlines in the form of meristems, and, if necessary, these can even be generated *de novo* from differentiated cells. Meristems are tissue in regions of plant growth and contain a few stem cells that give rise to various derivatives, including gametes. In Arabidopsis, the SAM stem cell niche is marked 41 by expression of the CLAVATA3 (CLV3) gene. Other transcription factors, signaling molecules (including CLV3), and receptors (for review see e.g. ^{3,4}) are necessary for stem cell 42 maintenance, however, our knowledge of the characteristics of "stemness" and the molecular 43 44 signatures of plant stem cells remains very limited. In order to overcome technical difficulties 45 associated with stem cell isolation, the number of *CLV3*-expressing cells was increased using an *ap1-1;cal1-1* double-mutant and enabled the first stem cell transcriptome ^{5,6}. However these 46 47 plants show stem cell-related phenotypic aberrations, and the analysis was limited to floral 48 meristems. A recent study used the INTACT method to gain insights into gene expression and 49 histone H3 modification dynamics of the entire SAM, but nonetheless included different cell 50 types and gene expression domains ⁷. With the same method, nuclei from SAM stem cells and differentiated leaves were isolated and compared for chromatin accessibility⁸. We wanted to 51 52 obtain information about gene expression and DNA methylation of pure SAM stem cell 53 fractions and generated Arabidopsis plants expressing a transcriptional fusion of the CLV3 promoter ⁹ and mCherry-labelled histone H2B. Microscopic analysis demonstrated correct and 54 55 specific expression of the pCLV3:mCherry-H2B marker in nuclei of approximately 20-40 stem 56 cells in 14-day-old seedlings (Fig. 1a). We applied fluorescence-activated nuclear sorting (FANS)¹⁰ to nuclei isolated from tissue manually enriched for shoot apical meristems and 57 58 collected mCherry-positive and -negative nuclei, with non-transgenic plants as controls (Fig. 59 1b, Supplementary Fig. 1a, Supplementary Table 1). Microscopic analysis confirmed that all sorted nuclei from the positive channel appeared intact and displayed red fluorescence, 60 61 validating the purity of the fraction (Supplementary Fig. 1b). The transcript level of endogenous 62 CLV3 was more than 1000-fold higher in mCherry-positive versus mCherry-negative nuclei or 63 whole seedlings (Fig. 1c). General RNA expression of sorted nuclei was highly correlated with 64 total RNA expression (Pearson correlation coefficient = 0.94; Supplementary Fig. 1c), indicating that nuclear RNA represents well the transcriptome of whole cells. Therefore, we 65

66 will refer to samples from mCherry/CLV3-positive nuclei as stem cells and to mCherry/CLV3-

67 negative samples from the closest neighboring tissue as non-stem cells.

68 We generated and sequenced RNA expression libraries from stem and non-stem cells isolated 69 from heart through torpedo stage embryos (E), 7 day- (D7), 14 day- (D14), and 35 day- (D35) 70 old plants (Supplementary Fig. 2a,b, Supplementary Table 2), aiming (i) to identify expression signatures in stem cells preceding major developmental switches, (ii) to find genes that are 71 72 involved in epigenetic resetting and germline formation, and (iii) to detect "stemness core 73 genes" whose expression would characterize stem cells independent of development. 74 Normalized read counts demonstrated high enrichment of CLV3 and mCherry transcripts in 75 stem cells at all developmental stages (Fig. 2a). High expression of the meristem marker genes 76 STM and KNAT1 relative to nuclei of 14-day-old whole seedlings (S14) confirmed the 77 meristematic features of the non-stem cells (Fig. 2a).

78 Transcriptome-wide clustering analysis showed that the expression signature of stem cells is 79 dominated by the respective developmental stage rather than by cell type (Fig. 2b). Pairwise 80 comparison between stem cells with the respective non-stem cells revealed differentially 81 expressed genes (DEGs, q < 0.05) at all four timepoints, respectively (Fig. 2c), the majority 82 upregulated in stem cells (with the exception of the embryo samples). Interestingly, GO term 83 analysis revealed that GOs describing reproductive processes, floral organ development, and 84 inflorescence development were already enriched in E, D7, and D14 (Supplementary Table 3, 85 Supplementary Fig. 3), while their absence in D35 stem cells was likely due to the low number of DEGs. 86

Overlap analysis between samples (Fig. 2d, Supplementary Fig. 4) revealed many stagespecific DEGs but also identified a set of 10 core genes (including *CLV3*) that were more highly expressed in stem cells of all four stages (Fig. 2a, Supplementary Fig. 4 and 5, Supplementary Table 4), and 23 genes with elevated expression in three out of the four stages (Supplementary Table 4). Twelve out of these 33 genes encode transcription factors (p-value for enrichment:
1.24e-08). Seven have already been connected with a meristem- or stem cell-related function
(Supplementary Table 4) leaving the remaining 26 as candidates for a potential role in stem cell
maintenance.

95 We could not detect significant overlap with transcript analysis in the SAM during flower induction⁷, probably due to differences in experimental set up and tissue type. The meristem 96 transcriptome of the *ap1-1;cal1-1* double-mutant⁶ had limited but significant overlap for 97 98 upregulated genes (Supplementary Table 5). Comparison with transcriptome data for different types of root meristem cells¹¹ resulted in an overlap especially with upregulated genes with 99 WOX5-expressing cells of the quiescent center (Supplementary Table 5). Also noticeable was 100 101 an overlap between upregulated stem cell DEGs with genes related to DNA methylation or siRNAs highly expressed in meristematic tissue ¹² (Fig. 3a). Among these are transcripts of two 102 103 Argonaute proteins (AGO5 and AGO9), two histone methyltransferases (SUVH4 and SUVR2), 104 the nucleosome remodeler DDM1, and three putative RNA-dependent RNA polymerases 105 (RDR3, 4, and 5). This indicated that specific family members of prominent epigenetic 106 components were upregulated in stem cells. Since AGO9, SUVH4, and DDM1 (among others) 107 are necessary for TE repression ¹³⁻¹⁵, we asked whether TEs were downregulated in stem cells relative to the surrounding cells. Indeed, several Arabidopsis TE families ¹⁶ were 2-fold less 108 109 expressed in stem versus non-stem cells through the four stages (Fig. 4 and Supplementary 110 Table 6). Surprisingly, with the same significance threshold, we found other TE families that 111 were more highly expressed in stem than in non-stem cells (Fig. 4, Supplementary Table 6). 112 Strikingly, at D7, the number of highly expressed TE groups coincided with the lowest number 113 of downregulated other groups, indicating a transient loss of control over TE expression in stem 114 cells at this early stage of vegetative growth, followed by resilencing towards generational 115 transition.

TEs overexpressed in D7 were mostly COPIA LTR-retroelements and Mutator-like DNA transposons but also included Helitrons, gypsy-like LTR elements, and SINEs (Supplementary Table 6). As LTR retroelements are more prevalent within pericentromeric regions, whereas SINEs and Helitrons are distributed on chromosome arms ¹⁷, TE expression in stem cells occurred independently of chromosomal localization. We could not find a bias for TEs that were recently mobile in natural populations ¹⁸, nor for transposons with new insertions in DNA methylation-deficient mutants ^{19,20}.

123 To determine whether TE expression was influenced by changing DNA methylation, we 124 performed whole-genome bisulfite sequencing of genomic DNA from D7, D14, and D35 stem and non-stem nuclei, with material from 7 d- and 14 d-old seedlings as reference. Modification 125 of cytosines in plants (reviewed in ²¹) at CG sites (mCG) is mainly achieved by MET1 and 126 127 occurs in repetitive sequences as well as along the gene body of protein-coding genes. Cytosine 128 methylation at CHG sites (mCHG) (H = A, C, or T) is installed by CMT2 and CMT3. Cytosine 129 methylation at CHH sites (mCHH) is established by DRM1 and DRM2 as well as CMT2. 130 mCHG and mCHH are mostly restricted to repetitive sequences and important for TE silencing. 131 mCHG is recognized by the histone methyltransferase SUVH4 which methylates histone H3 132 on lysine 9, a binding site for CMT3, and thereby reinforces DNA methylation in heterochromatic domains ²². 133

Analysis of DNA methylation distribution revealed pronounced differences around the centromeres for mCHG and mCHH, with the highest mCHG and lowest mCHH portion in stem cells of D35 (Fig. 5a and Supplementary Fig. 6). Congruent with the distribution along the chromosomes, metaplot analyses revealed that these methylation differences were found at TEs, while protein-coding genes were not affected (Fig. 5b). mCHG levels increased with developmental age, and TEs in stem cells had consistently higher mCHG levels than the respective non-stem cells, reaching a maximum at D35. Conversely, mCHH decreased withdevelopmental age, most pronounced in stem cells (Fig. 5b).

While TE groups varying in genomic location, cytosine content, structure and localization of repeats, and siRNA targeting sites ¹⁷ showed similarly increasing mCHG and decreasing mCHH in stem cells over developmental time (Supplementary Fig. 7), there was a correlation with their length: plotting methylation levels of TEs against their size range (Fig. 5c) revealed that mCHG in older meristems increased more in long TEs (>2.5 kb), parallel to decreasing mCHH. This suggests a contribution of DDM1, as this chromatin remodeler mediates methylation preferentially at long TEs ²³.

149 In order to understand which DNA methylation components are involved in methylation 150 dynamics in stem cells, we identified differentially methylated regions (DMRs) for each 151 timepoint and compared them with DMRs of mutants lacking different epigenetic 152 components²⁴. Increased mCHG in stem cells was especially pronounced at D14 in hypo-153 DMRs of *ddm1*, *suvh4*, *cmt3*, and *suvh456*; DMRs with reduced mCHH overlapped with those 154 of cmt2, suvh456, ddm1, and met1 (Fig. 6). This suggested a concerted action between the 155 chromatin remodeler DDM1 and the reinforcing heterochromatin formation of CMT3 and 156 histone methyltransferases to establish strong CHG methylation in stem cells entering the 157 reproductive phase. Furthermore, the reduction of CHH methylation in *cmt2* and *suvh456* 158 DMRs indicated a functional interference of CMT3 activity in stem cells with the related 159 CMT2.

The elevated TE expression at D7 correlated with a minimum of *AGO5* and *AGO9* transcript expression (Fig. 3b). While they belong to different clusters of the AGO clade ²⁵, both have been previously identified to be expressed in meristematic tissue of embryos ^{26,27} or in gametes or gametophytes ^{28,29}. AGO5 has not been connected with RNA-directed DNA methylation (RdDM) of TEs ³⁰, and neither *ago5* nor *ago9* showed many DMRs in DNA of whole

seedlings²⁴. However, AGO9 can restore methylation in an *ago4* mutant if accordingly 165 expressed ²⁶, suggesting that it can substitute RdDM-related functions. Although their exact 166 167 molecular functions and the subpopulation of bound small RNAs in the stem cells remain to be 168 determined, their expression anticorrelated with active TEs could hint to a specific protection 169 of germline precursor cells from virus and/or TE invasions. The transient loss of TE control in 170 early vegetative stages might even provide the sequence-specific information via small RNAs 171 that are then available for stem cell-enriched or -specific silencing components at later stages. 172 It has often been noted, but not explained, that mutants lacking major components of the RNA-173 directed DNA methylation (RdDM) pathway have no or only mild developmental defects. 174 Reinforced silencing in stem cells during development, involving additional specific factors 175 like RDR3, 4, or 5, may be responsible for this resilience. Such a barrier might break down only 176 upon special conditions, as indicated by stress-induced transposition prior to flower formation that occurs only in RdDM-compromised mutants ¹⁹. Interestingly, a recent study showed that 177 male premeiotic meiocytes also exhibit high mCG and mCHG methylation and low mCHH³¹. 178 179 This raises the intriguing possibility that cells of the central zone of the SAM enter a germline 180 DNA methylation state long before they can be cytologically distinguished. Alternatively, our 181 data could also suggest the presence of several cell types within the central domain of the SAM. 182 The possibility to extend the isolation of stem cells at different stages, from mutants and under 183 different external conditions, will enable future experiments to shed more light on epigenetic 184 maintenance and dynamics in germline precursor cells.

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186 Data access

187 DNA bisulfite and RNA-seq data have been deposited in the ArrayExpress database at EMBL188 EBI (www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-5478 and E-MTAB189 5479.

190

- 191 Methods
- 192 Plant material

193 All experiments were performed with Arabidopsis thaliana ecotype Col-0, wild type or 194 transgenic for pCLV3:H2B-mCherry. The pCLV3:H2B-mCherry construct was generated as 195 follows: the coding sequence of the H2B gene was PCR-amplified with primer H2B-forward 196 and H2B-reverse (Supplementary Table 7) from cDNA prepared from 14 d-old seedlings. The vector pCLV3:erCFP⁹ was cut with BamHI and SacI, and the H2B amplicon was inserted (In-197 198 Fusion, Clontech) into the open vector. The resulting plasmid was opened with SacI and In-199 Fusion-filled with a PCR-amplified mCherry-coding fragment using the primers mCherry-200 fusion-F1 and mCherry-fusion-R1 (Supplementary Table 7). Correct sequence of the resulting 201 vector pCLV3:H2B-mCherry was confirmed by Sanger-sequencing. The construct was used to generate transgenic plants by the floral dip method ³². Primary transformants were selected with 202 203 glufosinate (Merck) and their progeny screened for lines with a segregation ratio of 3 resistant 204 to 1 sensitive plant. Homozygous offspring were propagated for seed amplification.

205 Growth conditions

All plants were grown either *in vitro* on GM medium with or without selection or on soil under a 16 h light/8 h dark regime at 21°C. Material was always harvested at the same time of the light period.

209 Microscopic analysis and immunostaining

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For wide-field microscopy, plant material was immersed in PBS buffer and imaged with a Zeiss Axio Imager epifluorescence microscope. Isolated nuclei were imaged with an LSM780 Axio Observer, and Images were deconvolved using Hyugens Core (Scientific Volume Imaging) with a theoretical PSF. Immunostaining was performed according to ³³, with an additional clearing step using ScaleA ³⁴ and DAPI as counterstain. Anti-mCherry nanobodies were purchased from Chromotek (#rba594-100). Immunostains of meristems were imaged using the Airyscan mode on an LSM880 Axio Observer.

217 Fluorescence-activated nuclei sorting (FANS)

218 For 7D/14C/35D samples, 200-800 apexes (depending on size) of soil-grown plants with the 219 corresponding age were collected. For embryo samples, ovules from siliques of a few 220 representative plants were analyzed to contain early heart till early torpedo stage embryos, and 221 developmentally identical siliques were used to dissect 3000-4000 ovules. Collected material 222 was immediately transferred into nuclei isolation buffer on ice (NIB: 500 mM sucrose, 100 mM KCl, 10 mM Tris-HCl pH 9.5, 10 mM EDTA, 4 mM spermidine, 1 mM spermine and 0.1% v/v 223 2-mercaptoethanol, prepared just before use ³⁵). The material was then transferred into a tube 224 225 containing 1.8 ml of nuclear extraction buffer (NEB of the Sysmex CyStain® PI Absolute P kit 226 (#05-5022) plus 1% v/v 2-mercaptoethanol) and disrupted with the TissueRuptor (Qiagen) at 227 the lowest speed for 1 min. The suspension was filtered (30 µm filter nylon mesh, Sysmex # 228 04-0042-2316) and centrifuged for 10 min at 4000 rcf at 4°C. The nuclear pellet was 229 resuspended in Precise P staining buffer (Sysmex #05-5022; plus 1% v/v 2-mercaptoethanol 230 and DAPI to a final concentration of 5 μ g/ul), incubated for 15 min and again filtered (30 μ m) 231 into tubes (Sarstedt #55.484.001). Sorting was performed on a BD FACSAriaTM III cell sorter 232 (70 µm nozzle). Forward/Side scatter and DAPI and mCherry gating were adjusted with wild 233 type nuclei (DAPI-positive, mCherry-negative) as reference. The mCherry gate was adjusted 234 so that a maximum of 1/10 of mCherry events occurred in wild type compared to the 235 pCLV3:mCherry-H2B line. For DNA extraction, nuclei were directly sorted into Genomic Lysis Buffer (Quick-DNA Microprep Kit, Zymo Research, #D3020,), and DNA was purified 236 237 according to the suppliers' protocol for whole blood and serum samples. DNA was quantified 238 using pico-green on a NanoDrop fluorospectrometer (Thermo Scientific). For RNA isolation, 239 NIB, NEB, and staining buffer were complemented with RiboLock RNase inhibitor (Thermo 240 Scientific #EO0381, final concentration 1 U/µl) and nuclei were directly sorted into TRIzol LS 241 (Ambion, #10296028). RNA was prepared according to the manufacturers' recommendation, 242 except that nuclease-free glycogen (Thermo Scientific) was added during an overnight 243 precipitation at -20°C. Amount and quality of RNA was determined on an RNA 6000 pico-chip (Bioanalyzer/Agilent Technologies). For DNA and RNA extraction, DNA-LoBind tubes 244 245 (Eppendorf, #022431021) were used.

qPCR analysis

For qPCR and enrichment analysis, RNA was extracted with TRIzol LS (Ambion) either from
sorted nuclei or from shock-frozen and ground tissue material. RNA was treated with DNAse
(Thermo Scientific, #79254) and reverse-transcribed with iScript (Biorad, #172-5038). qPCR
assays were performed with Universal ProbeLibrary (UPL) assays (Roche, # 06402682001)
with primers and probes described in Supplementary Table 7.

252 Library preparation and sequencing

For RNA library preparation, total RNA of biological duplicates was extracted either from nuclei directly sorted into TRIzol LS or from shock-frozen ground material and used to generate cDNA libraries with the SMART-Seq v4 Ultra Low Input RNA Kit (Clontech). For the comparison with the nuclear RNA transcriptome, RNA was extracted from DAPI-stained FANSed nuclei isolated from 14 d-old p*CLV3:mCherry-H2B* seedlings with the same protocol as for cDNA production. cDNA populations were paired-end sequenced on a HiSeq 2500 Illumina sequencing platform. For bisulfite library preparation, at least 200 pg of DNA was used. Libraries were prepared with the Pico Methyl-Seq Library Prep Kit (Zymo Research
#D5456) according to the manufacturer's protocol.

262 Analysis of the RNA-sequencing data

For the analysis of nuclear to total RNA expression correlation, Tophat ³⁶ was used for mapping
to the TAIR10 reference genome after removal of low-quality bases with Trimmomatic ³⁷
(parameters: LEADING:8 TRAILING:10 SLIDINGWINDOW:4:15 MINLEN:50;).
Cuffdiff ³⁸ was used for normalization.

267 For all other analysis, RNA-seq reads were first adapter- and quality-trimmed with Trim 268 Galore! (Krueger F. Trim galore, v0.4.1, with default parameters). The reads were then aligned to the TAIR10 reference genome (including the mCherry sequence) with STAR 39 (v2.5.2a) 269 (Supplementary Table 8). Alignment parameters for STAR were set by the quantifier RSEM ⁴⁰ 270 271 (v1.2.30), which are based on previous ENCODE standards. The annotation used for 272 quantification was Araport11. RSEM was run with default settings. To correct for possible 273 positional biases in the data, we activated RSEM's positional bias correction option (--estimate-274 rspd). The resulting gene expression tables were imported into R (v3.4) via the tximport package ⁴¹ (v1.4.0). Consecutive differential gene expression analysis was performed with 275 DESeq2⁴² (v1.16). Samples of the same stages were analyzed pairwise via DESeq2s Wald test 276 (FDR < 0.05). To detect genes that are differentially regulated in stem cells across all 277 278 timepoints, we made use of DESeq2's model-based likelihood ratio test (LRT, FDR < 0.05). 279 The LRT allowed us to investigate how well the expression of a gene is recapitulated by 280 different models. DESeq2 compares two models, one full model and a reduced model. Our full 281 model factored in the cell type, the stage, and the interaction of both, while our reduced model 282 did not factor in the interaction, leaving us with a set of differentially expressed genes whose 283 variation can be explained by a combination of cell type and time. The RNA-seq pipeline is available under https://gitlab.com/nodine-lab/rsem-rna-seq-pipeline/. GO enrichments were 284

calculated using the AmiGO2 tool and the PANTHER classification system
 (<u>http://amigo.geneontology.org/rte</u>)⁴³. Visualization and clustering of the data was achieved
 using the R packages "gplots" and "gclus".

288 **DEG TE-Families**

All RNAseq samples were quality-trimmed using cutadap (v1.14) (Marcel Martin; Cutadapt removes adapter sequences from high-throughput sequencing reads; EMBnet.journal; Vol17, No1) and trimmomatic ³⁷ (v0.36). STAR ³⁹ (v2.5.2a) (Col-0 Arabidopsis reference genome, the Araport11 gene and TE annotations) was used as reference to map the reads, allowing multiple hits (--outFilterMultimapNmax 100 and --winAnchorMultimapNmax 100). TEtranscripts from the TEToolkit ⁴⁴ (v1.5.1) was used in multi-mode to find DEG TE-families.

295 Analysis of the bisulfite-sequencing data

296 Illumina HiSeq 2500 sequencing data was obtained from three stages (D7, D14, and D35) each 297 in three different settings (+: FANS-sorted stem cell tissue, -: non-stem cell but meristematic 298 tissue, s; whole seedling). Samples D14 and D35 were sequenced with 125 bp paired end reads, 299 D7 with 50 bp paired end reads (Supplementary Table 8). The data were quality-checked 300 (fastqc) and trimmed with TrimGalore (Krueger F. Trim galore, v0.4.1, default settings with stringency = 1) and trimmomatic 37 (v0.36, sliding window: 4:20, leading: 20). Bismark 45 301 302 (v0.18.1 with Bowtie2 v2.2.9) was used to map the reads to the Arabidopsis thaliana Col-0 303 reference genome (including mitochondria and chloroplast genomes) in the non-directional 304 mode with a mapping stringency of L,0,-0.6. A mapping-position-based removal of duplicates 305 (Bismark) was applied, and the C-to-T conversion rate was calculated using the reads mapped 306 to the chloroplast genome (ranging from 98.9 to 99.5%). Methylation was called (Bismark), 307 ignoring the first bases according to the M-Bias plots. Samples with same stages and settings 308 were pooled to a single sample, resulting in genome coverages for the nuclear genome from 309 16,4x to 53,9x.

310 DMR analysis

Differentially methylation positions (DMP) were identified by Fisher's exact test. Their positions were clustered together based on a minimum distance of 50 bp between DMPs to call a differential methylated region (DMR). DMR calling was done using methylpy (https://github.com/yupenghe/methylpy.git) version 1.1.9. We used custom R and python scripts for further analysis of these DMRs.

316

317 Author contributions

- 318 R.G. and O.M.S. designed the study and wrote the manuscript, R.G., N.D., A.G., G.B., N.L.,
- and M.D. performed the experiments, K.R., T.N., R.P., and F.H. analyzed the data, Ma.N. and
 Mi.N. discussed the results and commented on the manuscript.

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328 **Competing interests**

329 The authors declare no competing financial interest.

330

331

Figure legends

333 Figure 1: Establishment of FANS for stem cells of the shoot apical meristem (SAM).

- (a) Expression of H2B-mCherry under control of the *CLV3* promoter in 14 d-old seedlings.
- 335 Whole-mount immunostaining using α -mCherry antibodies and laser scanning microscopy
- 336 (scale bar 10 μm). (b) Example of a FANS experiment: mCherry-positive (+) and mCherry-
- 337 negative (-) gates of DAPI-gated nuclei. Numbers indicate total number and percent of DAPI
- events. (c) Enrichment of *CLV3* transcript in mCherry-positive nuclei determined by qRT-PCR.

Figure 2: Differential RNA expression in SAM stem cells during development.

- 340 (a) Expression of CLV3, mCherry, and the meristem marker genes STM and KNAT1. (b)
- Hierarchical clustering of expression data. (c) Number of DEGs between stem and non-stem cells at each timepoint. The banded portion of the bars indicates the number of transcription factor genes (also in parenthesis). (d) Overlap of genes with higher expression in stem cells (excluding *mCherry*). s = stem cells; n = non-stem cells.

Figure 3: Expression analysis of genes related to epigenetic regulation.

(a) Expression heatmap (in alphabetical order of gene acronyms). (b) Expression of
significantly upregulated DNA methylation-related genes in stem cells, marked with # in (a).
Asterisks indicate timepoints of significantly different expression between stem and non-stem
cells. s = stem cells; n = non-stem cells.

350 Figure 4: Expression analysis of transposable elements.

- 351 (a) Heatmap of expression differences for all 318 Arabidopsis TE groups in stem cells relative
- to non-stem cells at different timepoints. (b) Number of TE groups with at least 2x expression
- 353 difference at the different timepoints.

Figure 5: DNA methylation analysis of stem cells at different developmental stages.

- 355 (a) CG, CHG, and CHH methylation at chromosome 3 in stem and non-stem cells. (b)
- 356 Metaplots of DNA methylation at CG, CHG, and CHH for genes and transposons. (c) Locally
- 357 weighted scatterplot smoothing fit of CG, CHG, and CHH methylation levels in stem cells and

358	non-stem cells	plotted on TE	size. D7. D14 a	and $D35 = sorted$	nuclei 7. 14.	and 35 d.a.g.,	S7 and
220							

359 S14 = above-ground seedlings 7 and 14 d. a. g., s = stem cells; n = non-stem cells.

Figure 6: DMR analysis of stem cells.

361 DNA methylation differences between stem and non-stem cell nuclei within DMRs of different 362 epigenetic mutants. For each category, the scale denotes the number of standard deviations of 363 differential methylation in relation to the rest of the genome.

364

365 Supplementary Figure 1: Isolation of stem cell nuclei and RNA comparison.

(a) Gating strategy used for FANS of stem cells. Representative FANS plots are shown. Events
are gated for DAPI (top row) and next either for mCherry+ or mCherry- (bottom row). For
numbers see also Supplementary Table 1. (b) Examples of mCherry-positive nuclei after FANS
(scale bar 5 µm). (c) Correlation of log10-normalized FPKM values of nuclear and total RNA
extracted from 14 d-old seedlings.

371 Supplementary Figure 2: Growth stages used for genome-wide expression and DNA 372 methylation analysis in stem and non-stem cells.

373 (a) Developmental stages of representative plants (scale bars 1 cm). (b) Wide-field microscopic 374 images with RFP filters. (c) LSM pictures of representative plants. For better visualization 375 DAPI was used as counterstain in E, D7 and D14. IM = Inflorescence meristem. FM = Floral 376 meristem. Scale bars in (b): 60 μ m for the embryo; 1 mm for the other three stages. Scale bars 377 in (c): 20 μ m.

378 Supplementary Figure 3: Clustered heatmap displaying GO-term enrichment.

- 379 Color codes represent the negative ln of the Bonferroni corrected p-value for enrichment of
- ach GO-term. A p-value of 0.05 corresponds approximately to 3. See also Supplementary
- 381 Table 3 for exact values. s = stem cells; n = non-stem cells.
- **382** Supplementary Figure 4: Overlap of DEGs at different timepoints.

- 383 (a) Venn diagrams for genes up- and (b) downregulated in stem cells, respectively. (c) p-values
- 384 (hypergeometric tests) for likelihood of overlap of upregulated genes in different pairs of
- 385 timepoints.
- 386 Supplementary Figure 5: Expression of core stemness genes.
- 387 Bar plots of expression of genes that are significantly upregulated in SAM stem cells throughout
- 388 development. s = stem cells; n = non-stem cells.
- 389 Supplementary Figure 6: DNA methylation analysis of stem cells on all five Arabidopsis
- 390 chromosomes in stem and non-stem cells at different developmental stages.
- 391 s = stem cells; n = non-stem cells.
- 392 Supplementary Figure 7: DNA methylation analysis of different TE classes in stem and
- 393 non-stem cells at different developmental stages.
- 394 s = stem cells; n = non-stem cells.
- 395
- 396 Supplementary Table 1: Examples of FANS data
- 397 Supplementary Table 2: RNA expression data
- 398 Supplementary Table 3: GO-term annotations
- 399 Supplementary Table 4: Overlapping DEGs
- 400 **Supplementary Table 5: Comparison with other data sets**
- 401 Supplementary Table 6: TE expression data
- 402 Supplementary Table 7: Primer sequences
- 403 Supplementary Table 8: Mapping statistics
- 404

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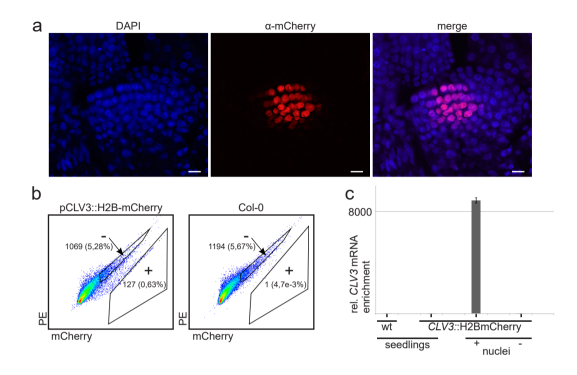


Figure 1 | Establishment of FANS for stem cells of the shoot apical meristem (SAM). (a) Expression of H2B-mCherry under control of the *CLV3* promoter in 14 d-old seedlings. Whole-mount immunostaining using α -mCherry antibodies and laser scanning microscopy (scale bar 10 μ m). (b) Example of a FANS experiment: mCherry-positive (+) and mCherry-negative (-) gates of DAPI-gated nuclei. Numbers indicate total number and percent of DAPI events. (c) Enrichment of *CLV3* transcript in mCherry-positive nuclei determined by qRT-PCR.

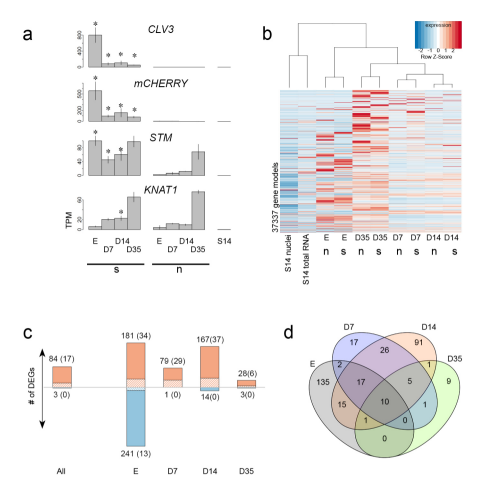


Figure 2 | **Differential RNA expression in SAM stem cells during development.** (a) Expression of *CLV3, mCherry* and the meristem marker genes *STM* and *KNAT1.* (b) Hierarchical clustering of expression data. (c) Number of DEGs between stem and non-stem cells at each timepoint. The banded portion of the bars indicates the number of transcription factor genes (also in parenthesis). (d) Overlap of genes with higher expression in stem cells (excluding *mCherry*). s = stem cells; n = non-stem cells.

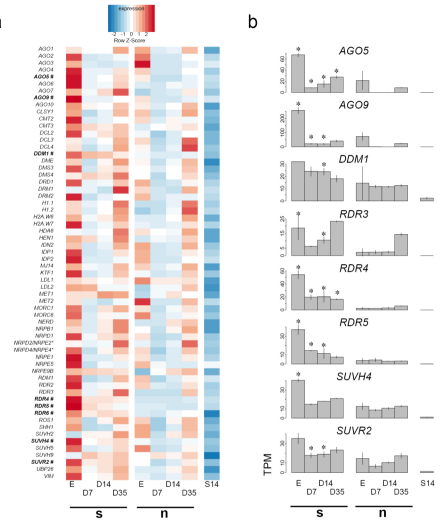


Figure 3 | Expression analysis of genes related to epigenetic regulation. (a) Expression heatmap (in alphabetical order of gene acronym). (b) Expression of significantly upregulated DNA methylation-related genes in stem cells, marked with # in (a). Asterisks indicate timepoints of significantly different expression between stem and non-stem cells. s = stem cells; n = non stem cells.

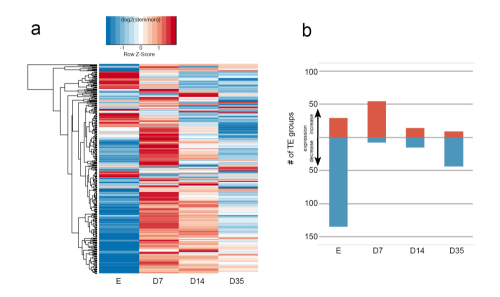


Figure 4 | Expression analysis of transposable elements. (a) Heatmap of expression differences for all 318 Arabidopsis TE groups in stem cells relative to non-stem cells at different timepoints. (b) Number of TE groups with at least 2x expression difference at the different timepoints.

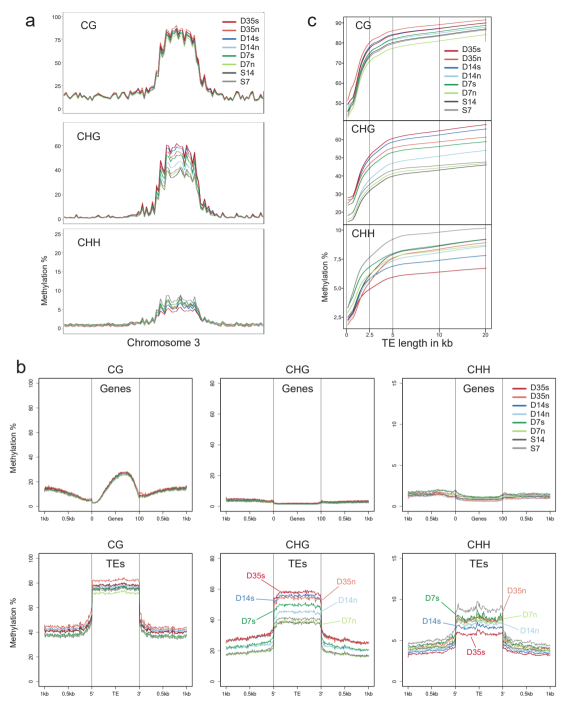


Figure 5 | DNA methylation analysis of stem cells at different developmental stages. (a) CG, CHG and CHH methylation of chromosome 3 in stem and non-stem cells. (b) Metaplots of DNA methylation at CG, CHG, and CHH for genes and transposons. (c) Locally weighted scatterplot smoothing fit of CG, CHG and CHH methylation levels in stem cells and non-stem cells plotted on TE size. D7, D14 and D35 = sorted nuclei 7, 14, and 35 d.a.g., S7 and S14 = above-ground seedlings 7 and 14 d. a. g., s = stem cells; n = nonstem cells.

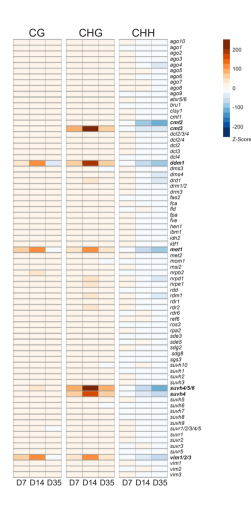
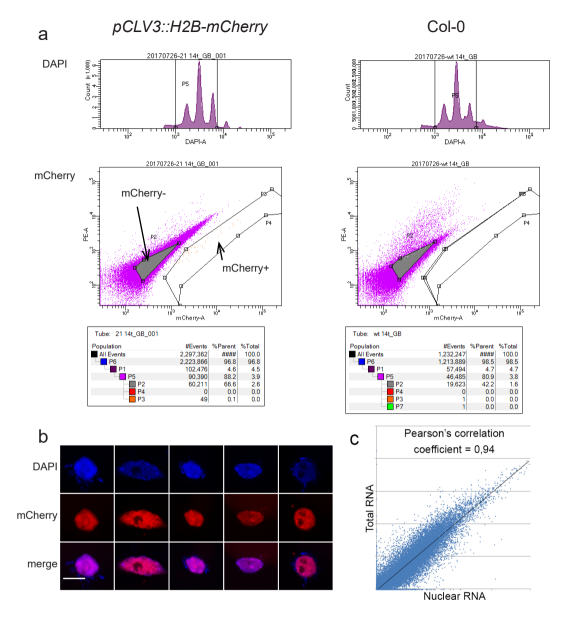
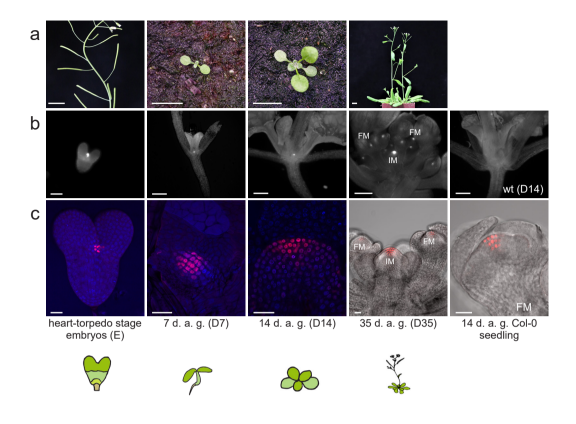


Figure 6| DMR analysis of stem cells.

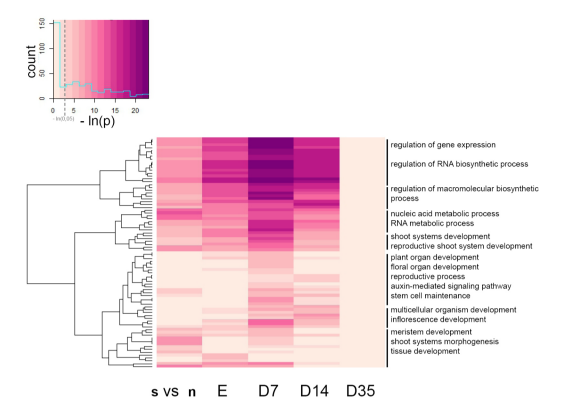
DNA methylation differences between stem and non-stem cell nuclei within DMRs of different epigenetic mutants. For each category, the scale denotes the number of standard deviations of differential methylation in relation to the rest of the genome.



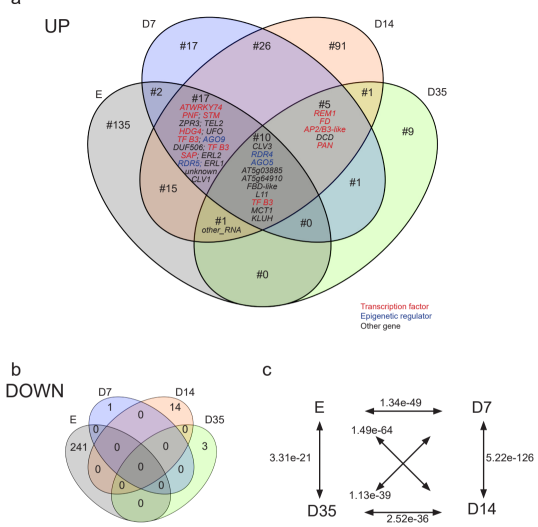
Supplementary Figure 1 | Isolation of stem cell nuclei and RNA comparison. (a) Gating strategy used for FANS of stem cells. Representative FANS plots are shown. Events are gated for DAPI (top row) and next either for mCherry+ or mCherry- (bottom row). For numbers see also Supplementary Table S1. (b) Examples of mCherry-positive nuclei after FANS (scale bar 5 μ m). (c) Correlation of log10-normalized FPKM values of nuclear and total RNA extracted from 14 d-old seedlings.



Supplementary Figure 2 | Growth stages used for genome-wide expression and DNA methylation analysis in stem and non-stem cells. (a) Developmental stages of representative plants (scale bars 1 cm). (b) Wide-field microscopic images with RFP filters. (c) LSM pictures of representative plants. For better visualization DAPI was used as counterstain in E, D7 and D14. IM = Inflorescence meristem. FM = Floral meristem. Scale bars in (b): 60 μ m for the embryo; 1 mm for the other three stages. Scale bars in (c): 20 μ m.

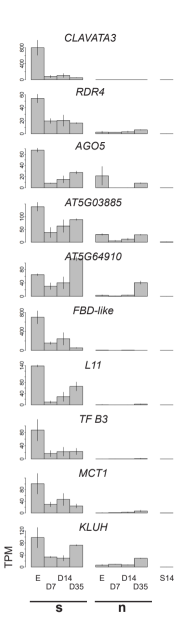


Supplementary Figure 3 | Clustered heatmap displaying GO-term enrichment. Color codes represent the negative In of the Bonferroni corrected p-value for enrichment of each GO-term. A p-value of 0.05 corresponds approximately to 3. See also Supplementary Table S3 for exact values. s = stem cells; n = non stem cells.

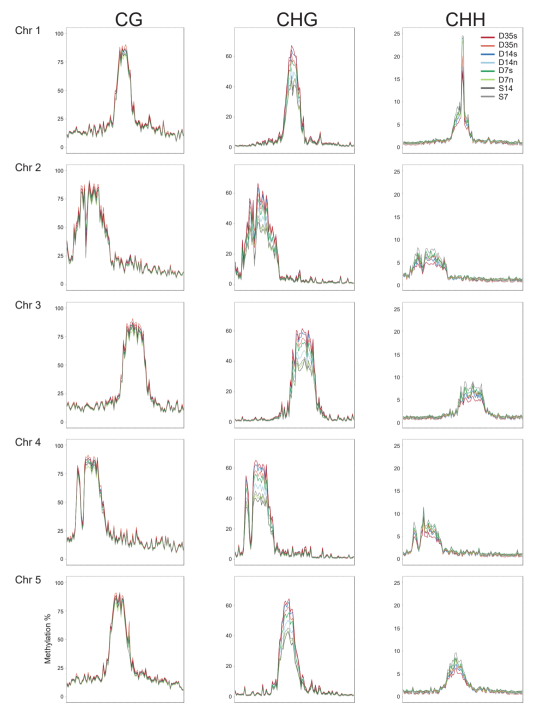


Supplementary Figure 4 | Overlap of DEGs at different timepoints. (a) Venn diagrams for genes up- and (b) downregulated in stem cells, respectively. (c) p-values (hypergeometric tests) for likelyhood of overlap of upregulated genes in different pairs of timepoints.

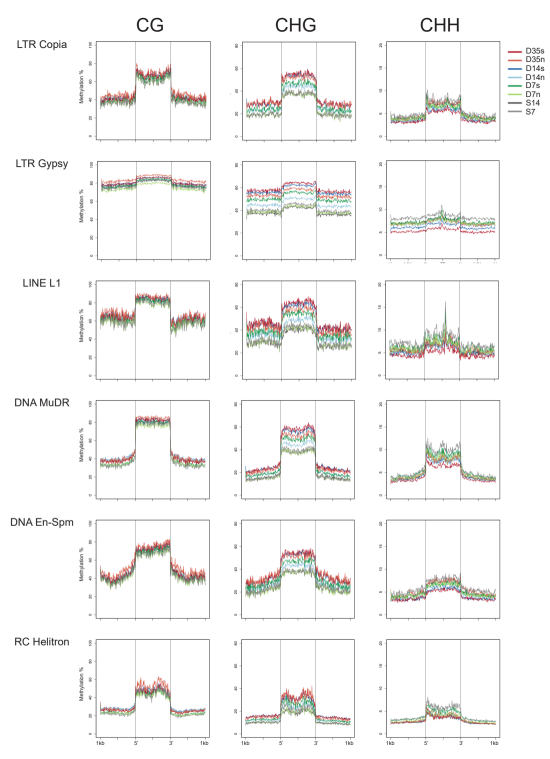
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Supplementary Figure 5 | Expression of core stemness genes. Bar plots of expression of genes that are significantly upregulated in SAM stem cells throughout development. s = stem cells; n = non stem cells.



Supplementary Figure 6| DNA methylation analysis of stem cells on all five Arabidopsis chromosomes in stem and non-stem cells at different developmental stages. s = stem cells; n = non-stem cells.



Supplementary figure 7| DNA methylation analysis of different TE classes in stem and non-stem cells at different developmental stages. s = stem cells; n = non-stem cells.