

1 **Title:** MORC proteins regulate transcription factor binding by mediating chromatin compaction in
2 active chromatin regions

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18

19 **Abstract**

20 **Background:** The Microrchidia (MORC) proteins are a family of evolutionarily conserved GHKL-
21 type ATPases involved in chromatin compaction and gene silencing. *Arabidopsis* MORC proteins
22 act in the RNA-directed DNA methylation (RdDM) pathway, where they act as molecular tethers
23 to ensure the efficient establishment of RdDM and *de novo* gene silencing. However, MORC
24 proteins also have RdDM-independent functions; although, their underlying mechanisms are
25 unknown.

26 **Results:** In this study, we examined regions of MORC binding where RdDM does not occur in
27 order to shed light on the RdDM-independent functions of MORC proteins. We found that MORC
28 proteins compact chromatin and reduce DNA accessibility to transcription factors (TFs), thereby
29 repressing gene expression. We also found that MORC-mediated repression of gene expression
30 was particularly important under conditions of stress. We showed that MORC proteins regulate

31 TFs through either direct or indirect interactions, and these TFs can in some cases regulate their
32 own transcription, resulting in feedforward loops.

33 **Conclusions:** Our findings provide insights into the molecular mechanisms of MORC-mediated
34 chromatin compaction and transcription regulation.

35 **Keywords:** Microrchidia, MORC, chromatin compaction, TF binding, transcription regulation

36

37 **Background**

38 The MORC proteins are a family of highly conserved GHKL-type ATPases involved in gene
39 silencing and chromatin compaction [1]. In *C. elegans*, MORC-1 can compact DNA through a
40 through topological entrapment [2], while in humans, MORC2 is recruited by the human silencing
41 hub (HUSH) complex for H3K9me3 deposition, chromatin compaction, and gene silencing [3]. In
42 mice, MORC1 is involved in germline transposon silencing [4], and MORC3 is essential for
43 transposon silencing in embryonic stem cells [5].

44 The *Arabidopsis* genome encodes six MORC proteins: MORC1, 2, 4, 5, 6, and 7 (MORC3 being
45 a pseudogene) [6]. These six proteins are functionally redundant, but colocalize with sites of RNA-
46 directed DNA methylation (RdDM) genome-wide [7], where they are critical for establishing
47 efficient RdDM and *de novo* gene silencing [7]. MORC7, when tethered to DNA using an artificial
48 zinc finger, can target RdDM to ectopic sites. MORC7 is also required for the silencing of a newly
49 integrated *FWA* transgene [7]. MORC proteins also act downstream of DNA methylation to
50 suppress gene expression, and are also involved in plant immunity — protecting against potential
51 pathogens by interacting with plant resistance (R) proteins [8,9]. However, the molecular
52 mechanisms underlying these RdDM-independent functions remain unknown. We previously
53 observed MORC binding sites where RdDM does not occur (MORC-unique sites) [7]; and by
54 studying these sites, we aim to shed light on the mechanisms underlying the RdDM-independent
55 functions of MORC proteins.

56 TOPLESS (TPL) and LEUNIG (LUG) are both Grocho (Gro)/TLE-type transcriptional co-
57 repressors in plants. They are characterized by a conserved glutamine-rich C-terminal domain
58 and an N-terminal WD-repeat domain [10]. The glutamine-rich domain participates in protein
59 oligomerization, and the WD-repeat domain interacts with downstream transcriptional regulators
60 [10]. The functional counterpart of the Gro/TLE family of proteins in yeast, Tup1, was originally
61 identified as a co-repressor that occupied the binding sites of transcriptional activators [11,12].

62 However, evidence now shows that Tup1 can switch from a co-repressor to a co-activator in
63 response to stress, and is required for the activation of certain genes related to the stress
64 response [12,13].

65 Here, we use MORC-unique sites to study the RdDM-independent functions of MORC proteins.
66 We show that MORC proteins compact chromatin and reduce DNA accessibility to TFs, thereby
67 repressing the transcription of stress-responsive genes.

68

69 **Results**

70 **MORC proteins bind to active chromatin regions devoid of RdDM**

71 We previously reported that approximately 80% of MORC7 binding regions overlap with sites of
72 RdDM [7]. MORC7 is recruited to these sites by the RdDM machinery, where it then facilitates
73 the efficiency of the RdDM pathway. However, the remaining 20% of MORC7 binding sites are
74 devoid of RdDM, as evidenced by a lack of Pol V occupancy [7]. The mechanisms underlying the
75 function of MORC7 within these RdDM-depleted regions remain unknown.

76 Mouse MORC3 recognizes and localizes to regions of H3K4me3-marked chromatin through its
77 CW domain [14]; however, Arabidopsis MORCs do not contain CW domains. To determine
78 whether Arabidopsis MORCs co-localize with specific chromatin features, we used the
79 ChromHMM method to investigate correlations between MORC7 and several well-characterized
80 chromatin features (H3K9ac, H3K27ac, H4K16ac, H3K4me1, H3K4me3, H3K36me2, H3K36me3,
81 H3K9me2, H3K27me3, Pol II, and Pol V). We analyzed chromatin states using a similar method
82 as previously reported [15] but also included Pol V ChIP-seq data. We found 13 different
83 chromatin states ([Supplementary Fig. 1](#)). MORC7 showed a strong correlation with Pol V (a
84 known indicator of RdDM sites), which was consistent with our previous findings (State 11,
85 [Supplementary Fig. 1](#)). Chromatin state 12 included sites enriched with MORC7 but depleted of
86 Pol V — indicative of MORC7-unique regions. We did not observe enrichment of histone marks
87 in these MORC7-unique regions ([Supplementary Fig. 1](#)).

88 Within these MORC7-unique regions, we identified two subgroups: MORC7A and MORC7B. The
89 ChIP-seq data for MORC7, Pol V, ATAC-seq and transposable element (TE) density indicated
90 that MORC7A was within a region of high chromatin accessibility and low TE density ([Fig. 1a, b](#)).
91 Consistent with the ChromHMM analysis, MORC7A displayed low levels of histone occupancy
92 and histone modification, although its flanking regions were enriched for active histone

93 modifications. This suggests that MORC7A is located within an active chromatin compartment
94 between genes (Fig. 1b).

95 MORC7B contained a high density of TE with no apparent active histone marks, reflective of its
96 heterochromatic location (Fig. 1b). We found that MORC7A regions had low levels of DNA
97 methylation, while MORC7B regions had high levels of methylation (Fig. 1c, d). These results
98 suggest that MORC7 binds to active and deep heterochromatic regions of DNA, where RdDM
99 does not occur, suggesting that it regulates gene expression at these sites through RdDM-
100 independent mechanisms.

101

102 **MORC7 preferentially binds to the promoters of TFs**

103 The genomic distribution enrichment data showed enrichment of MORC7A peaks over promoters
104 (Fig. 2a), but no enrichment of MORC7B peaks — consistent with their deep heterochromatic
105 localization. The functional annotation of the genes proximal to MORC7A suggested that they
106 were enriched with TFs (Table 1). The *Arabidopsis* genome encodes approximately 1491 TF
107 genes (5.5% of the genome) [16]. Of the genes proximal to MORC7A, 23% were TFs (p-value =
108 3.12E-36); these included PHYTOCHROME INTERACTING FACTOR (PIF), ethylene and auxin-
109 responsive transcriptional factors and Myb transcriptional factors (Supplementary Table 1). This
110 enrichment was more significant than enrichment for MORC7-Pol V common (p-value = 0.001)
111 and Pol V-unique (p-value = 1.6E-5).

112 Gene Ontology (GO) term analysis of genes proximal to MORC7A showed an enrichment of
113 negative regulation of auxin metabolic process (~80 fold), shade avoidance (~30 fold), and the
114 primary shoot apical meristem specification pathway (~30 fold) (Fig. 2b). The primary shoot apical
115 meristem specification pathway (GO0010072) is responsible for the growth of all post-embryonic,
116 above-ground plant structures [17]. In *Arabidopsis*, this pathway includes several *topless*-related
117 genes [17]. Interestingly, we found that MORC7 specifically bound to 12 of the 19 genes in this
118 pathway (Fig. 2c, Supplementary Fig. 2), and co-localized with Pol V at an additional three. We
119 show examples of MORC7 enrichment over the promoter regions for the four *TOPLESS* genes in
120 Supplementary Fig. 2.

121

122 **MORC7 closely co-localizes with some TFs**

123 To investigate the protein interaction network of MORC7 with chromatin, we re-analyzed
124 previously published crosslinked IP-MS data of MORC7 [7]. We identified 494 proteins (FDR <
125 0.05, FC > 2) that interacted with MORC7 (Fig. 3a), and found that many of these were involved
126 in either chromatin-related pathways or development (Fig. 3b). We also identified 68 TFs from the
127 MORC7 interacting proteins (68/494, p = 7.89E-12) (Supplementary Table 2). To further test
128 whether MORC7 co-localizes with TFs, we obtained binding site information for 200 TFs from the
129 DNA Affinity Purification and sequencing (DAP-seq) database [18], and performed pairwise peak
130 overlap analysis with MORC7 peaks. We found that MORC7A showed stronger co-localization
131 with TFs compared to MORC7B, MORC7-Pol V common and Pol V-unique regions (Fig. 3c). This
132 indicates that MORC7A peaks are associated with TF binding sites. We also re-analyzed three
133 factors in particular because published ChIP-seq data was available, PIF4 [19], ARF6 [20], and
134 TPR1 [21]. Metaplot analysis showed that with ChIP-seq data for MORC7A-unique, MORC7B-
135 unique, MORC7-Pol V Common, and Pol V-unique regions showed the strongest co-localization
136 with MORC7A (Fig. 3d, Supplementary Fig. 3).

137

138 **MORC7 influences TF binding through chromatin compaction**

139 To understand how MORC7 affects chromatin conformation, we performed an Assay for
140 Transposase-Accessible Chromatin with high-throughput sequencing (ATAC-seq) in *morc4*
141 *morc7*, *morc6*, and *morc hextuple* (*morchex*, in which all functional MORCs are knocked out)
142 mutants [6]. We plotted ATAC-seq data across the four groups, and found that MORC7A,
143 MORC7B and MORC7-Pol V common showed greater chromatin accessibility changes in the
144 mutants, particularly over the MORC7A regions (Fig. 4a, b). This phenotype is consistently
145 observed in *morc4* *morc7*, *morc6* and *morchex* — with *morchex* showing the most pronounced
146 phenotype (Fig. 4a). Interestingly, for Pol V-unique sites, DNA compaction was not reduced, but
147 actually became slightly increased (Fig. 4a). Consistently, we also observed an increase in DNA
148 methylation for Pol V-unique sites in the mutants (Fig. 1d). This suggests that Pol V may be
149 redistributed from the MORC7-Pol V common sites to Pol V-unique sites in the absence of MORC
150 proteins. This is consistent with our previous findings that suggested MORC proteins function as
151 molecular tethers to facilitate the recruitment of RdDM components [7].

152 To examine whether MORC-mediated DNA compaction affects TFs, we analyzed the ATAC-seq
153 data for TF footprints. When a TF binds to DNA, it inhibits the integration of DNA by Tn5
154 transposes, causing the binding motif to exhibit lower DNA accessibility, and the flanking regions

155 to exhibit higher DNA accessibility [22]. The footprints of 572 TFs downloaded from JASPAR were
156 analyzed in the *morc4* *morc7*, *morc6* and *morchex* mutants [23]. Many TFs showed substantially
157 stronger apparent binding within the MORC7A regions in the mutants. There were some
158 increases in binding within the MORC7B regions (although to a lesser degree than in MORC7A
159 regions) (Fig. 4c, d), while TF binding over RdDM sites was largely unaffected (Fig. 4e, f). The
160 metaplot of ATAC-seq signals over the TF binding sites for the MORC7A regions confirmed that
161 these TFs have stronger apparent binding in *morc4* *morc7*, *morc6* and *morchex* mutants – with
162 *morchex* showing the strongest binding changes, and the random control regions showing no
163 differences (Fig. 4g-l).

164 We previously showed that targeting either MORC7 or MORC6 ectopically in the *fwa-4* epiallele
165 background using ZF108 can trigger the silencing of *FWA* [7,24]. In addition to the *FWA* locus,
166 ZF108 can also bind thousands of off-target sites [24]. These off-target sites are preferentially
167 localized to promoter regions, and therefore provide an excellent opportunity to test whether the
168 presence of MORC proteins can affect TF binding. We compared TF footprints between ZF-
169 MORC6 and *fwa-4* and found a substantial decrease for many of the TF footprints in ZF-MORC6
170 plants. This supports the hypothesis that MORC proteins affect TF binding (Fig. 4j). Together,
171 these results suggest that MORCs inhibit TF binding by altering chromatin accessibility.

172

173 **MORC influences gene expression downstream of the TFs**

174 To understand whether MORC proteins regulate gene expression, we performed RNA-seq with
175 the *morchex* mutant. As MORC7A co-localizes strongly with PIF4 (Fig. 3d) – a central regulator
176 in temperature signaling [25] – we applied heat treatment to the *morchex* mutant. We first
177 compared the expression of genes proximal to MORC7A peaks in wild type (WT) and *morchex*
178 mutant without treatment. This showed that the genes proximal to MORC7A were slightly up-
179 regulated in the *morchex* mutant without treatment (Fig. 5a), including the TFs SEP3, PIF4,
180 ARF6/8, TPR1, LUG and SEU (Fig. 5b). After heat treatment, *morchex* displayed a stronger
181 response compared to the WT, with significantly more upregulated genes (Fig. 5c, d). Genes
182 proximal to MORC7A were enriched in shoot apical meristem specification pathways, and
183 consistently, we observed stronger upregulation of these genes in *morchex* after heat treatment
184 (Supplementary Fig. 4).

185

186 To confirm that MORC proteins affect TF binding, and to understand how they affect downstream
187 gene expression, we selected TFs TOPLESS (TPL) and LEUNIG (LUG) for ChIP-seq analysis,
188 because they were present in the MORC7 IP-MS data [7]. We expressed TPL and LUG fused
189 with a 3XFLAG-tag in both WT and *morchex*. Consistent with the TF footprint analysis, both TPL
190 and LUG displayed stronger binding at MORC7A regions, while only a slight increase in binding
191 was noted for the MORC7-Pol V co-binding sites in *morchex* (Fig. 5e, f) — consistent with an
192 increase in chromatin accessibility in *morchex* (Fig. 5g; Supplementary Fig. 5). We ranked TPL
193 and LUG binding sites based on the MORC ChIP-seq signals and divided them into three groups;
194 high, middle and low. Overall, in the *morchex* mutant, we observed increased TPL and LUG
195 binding, as well as increased chromatin accessibility across the regions with stronger MORC7
196 signals (Fig. 5g). We found that MORC7A-bound genes were downregulated in the *lug* mutant,
197 suggesting that LUG may facilitate expression of these genes (Fig. 5a). Using ChIP-seq data
198 together with RNA-seq data in the *lug* mutant, we identified 95 genes that appeared to be directly
199 regulated by LUG (Supplementary Table 3). We found that these LUG-regulated genes were
200 upregulated in *morchex*, particularly after heat treatment (Fig. 5h).

201

202

203 **Discussion**

204 We previously reported that MORC proteins are localized to sites of RdDM throughout the
205 genome, and function as molecular tethers to facilitate the efficient establishment of RdDM [7].
206 We showed that this RdDM-related function of MORC proteins is critical for *de novo* transgene
207 silencing [7,24]; however, this model does not explain other functions of MORC proteins. For
208 example, MORC1 and MORC6 were shown to work downstream of DNA methylation to repress
209 the expression of both the endogenous *SDC* gene and an *SDC* transgene, as well as other DNA
210 methylated targets in the genome [9]. In addition, *morc* mutants display various disease
211 phenotypes; for example, Kang et al. [8] reported that *morc1* is susceptible to Turnip Crinkle Virus
212 (TRV), while Harris et al. [6] reported that *morchex* is susceptible to the *Hyaloperonospora*
213 *arabidopsis* (Hpa) strain, Emwa1. However, the molecular mechanisms underlying the
214 additional functions of the MORC proteins remain unknown.

215 Here, we investigated the function of MORC7 in regions where RdDM does not occur, particularly
216 those near genes where no DNA methylation is present. We found that MORC proteins reduce
217 chromatin accessibility within these regions. Previous *in vitro* studies showed that *C. elegans*
218 MORC1 homodimers can topologically entrap and condense DNA through further oligomerization
219 of MORC1 proteins [2]. In addition, *Arabidopsis morc* mutants display pericentromeric
220 heterochromatin decondensation [9], which takes place with minimal losses of DNA methylation
221 throughout the genome. This indicates that MORC proteins contribute to chromatin compaction
222 independently of DNA methylation [9]. We show here that MORC proteins reduce chromatin
223 accessibility in methylation-free promoter regions of DNA, which may explain their mechanism for
224 methylation-independent gene regulation. We suggest that *Arabidopsis* MORCs may use a similar
225 mechanism of chromatin compaction to that of *C. elegans* MORC1 – compacting chromatin by
226 topological entrapment, thereby reducing its accessibility to TFs.

227 Plant MORCs have been implicated in plant pathogen responses. MORCs promote resistance in
228 some plant species and inhibit defense responses in others [8]. Upregulation of protein-coding
229 genes was previously shown in *morc4 morc7*; although, the underlying mechanisms of this was
230 unknown [6]. Here, we report that MORC proteins regulate gene expression by compacting
231 chromatin in promoter regions, thereby preventing access by TFs. In addition, MORCs
232 preferentially bound to the promoter regions of TFs, contributing to their regulation, and our
233 crosslinked IP-MS data suggested that MORCs are located in close proximity to some TFs.
234 Interestingly, we observed that many of these TFs bind to their own promoters – suggesting

235 regulation by a feedforward loop, which may amplify the effects of MORCs on transcriptional
236 networks.

237 Finally, we showed that MORC proteins are important for the regulation of gene expression,
238 particularly under stress conditions. We also found altered expression of heat-responsive genes
239 in *morchex*. Like with its role in plant pathogen defense response, it seems likely that the role of
240 MORCs in stress responses relates to its chromatin compaction of promoter regions and affects
241 in TF networks.

242 **Conclusions**

243 MORC proteins have a broad binding spectrum in the genome, and appear to participate in at
244 least three separate processes. They co-localize to sites of RdDM, facilitating efficient DNA
245 methylation establishment [7], they are needed to repress DNA methylated areas of
246 pericentromeric heterochromatin in a DNA methylation independent manner [9], and they co-
247 localize with TFs in unmethylated promoter regions, regulating TF binding and gene expression
248 by altering chromatin accessibility (Fig. 5i). Although it seems likely that MORC act in each of
249 these processes by topologically entrapping DNA, there are likely mechanistic differences that
250 can explain the localization and function of MORCs in these three different epigenetic
251 environments in the genome.

252

253 **Methods**

254 **Plant materials and growth conditions.**

255 All plants in this study were grown in standard greenhouse conditions (22 — 25 °C, 16 hrs light/8
256 hrs dark). The following plant materials were used in this study: *morchex* consisting of *morc1-2*
257 (SAIL_893_B06), *morc2-1* (SALK_072774C), *morc4-1* (SALK_051729), *morc5-1*
258 (SALK_049050C), *morc6-3* (GABI_599B06), and *morc7-1* (SALK_051729). For heat treatments,
259 plants were grown under 37 °C for 0.5 hours and put back to normal temperature for 48 hours for
260 recovery.

261 **Epitope-tagged transgenic lines**

262 Full-length genomic DNA fragments, including native promoter sequences, were cloned into
263 pENTR/D vectors (Invitrogen), followed by modified destination vectors carrying 3xFLAG with LR
264 Clonase (Invitrogen). All primers used in this study are available in **Supplementary Table 4**.

265

266 **Nuclei extraction and ATAC-seq library preparation**

267 The nuclei collection process from inflorescence and meristem tissues was performed in
268 accordance with previously described methods [26,27]. Freshly isolated nuclei were used for
269 ATAC-seq, as described elsewhere [28]. Inflorescence tissues were collected for extraction of
270 nuclei as follows: 5g (approximately) of inflorescence tissue was collected and immediately
271 transferred into the ice-cold grinding buffer (300mM sucrose, 20mM Tris pH 8, 5mM MgCl₂, 5mM
272 KCl, 0.2% Triton X-100, 5mM β-mercaptoethanol, and 35% glycerol); the samples were then
273 ground with Omni International General Laboratory Homogenizer at 4°C, and filtered through a
274 two-layer Miracloth using a 40-μm nylon mesh Cell Strainer (Fisher). Samples were spin filtered
275 for 10 min at 3,000 g, the supernatant was discarded, and the pellet was resuspended with 25ml
276 of grinding buffer using a Dounce homogenizer. The wash step was performed twice in total.
277 Nuclei were then resuspended in 0.5ml of freezing buffer (50mM Tris pH 8, 5mM MgCl₂, 20%
278 glycerol, and 5mM β-mercaptoethanol). Nuclei were then subjected to a transposition reaction
279 with Tn5 (Illumina). For the transposition reaction, 25μl of 2 x DMF (66mM Tris-acetate pH 7.8,
280 132mM K-Acetate, 20mM Mg-Acetate, and 32% DMF) was mixed with 2.5μl Tn5 and 22.5μl nuclei
281 suspension at 37°C for 30 min. The transposed DNA fragments were then purified with ChIP DNA
282 Clean & Concentrator Kit (Zymo). Libraries were prepared with Phusion High-Fidelity DNA
283 Polymerase (NEB), in a system containing: 12.5μl 2 x Phusion, 1.25μl 10mM Ad1 primer, 1.25μl
284 10mM Ad2 primer, 4μl ddH₂O, and 6μl purified transposed DNA fragments. The ATAC-seq
285 libraries were sequenced on the NovaSeq 6000 platform (Illumina).

286

287 **RNA-seq library preparation**

288 Total RNAs were extracted from 100mg (approximately) of flower buds using TRIzol and the
289 Direct-zol RNA Miniprep kit (Zymo, R2050). Sequencing libraries were prepared using the TruSeq
290 Stranded mRNA Library Prep kit (Illumina), according to the manufacturer's instructions, and
291 sequenced on a NovaSeq 6000 sequencer (Illumina).

292

293 **ChIP-seq library preparation**

294 10g of inflorescence and meristem tissues were used for ChIP-seq. ChIP assays were performed
295 as has been described previously [29]. Briefly, 2-4g of flower tissue was collected from 4 — 5-

296 week-old plants, and ground with liquid nitrogen. 1% formaldehyde containing a nuclei isolation
297 buffer was used to fix the chromatin for ten minutes. Freshly prepared glycine was then used to
298 terminate the crossing reaction. Shearing was performed via Bioruptor Plus (Diagenode), and
299 immunoprecipitations with antibodies were performed overnight at 4°C. Magnetic Protein A and
300 Protein G Dynabeads (Invitrogen) were added and incubated at 4°C for two hours. The reverse
301 crosslink was performed overnight at 65°C. The protein-DNA mix was then treated with Protease
302 K (Invitrogen) at 45°C for four hours. The DNA was purified and precipitated with 3M Sodium
303 Acetate (Invitrogen), glycoBlue (Invitrogen) and Ethanol overnight at -20°C. The precipitated DNA
304 was then used for library preparation using the Ovation Ultra Low System V2 kit (NuGEN), which
305 was then sequenced using an Illumina NovaSeq sequencer. The anti-FLAG M2 (Sigma) antibody
306 was used in this study. Libraries were prepared using the NuGen Ovation Ultra Low System V2
307 kit, in accordance with the manufacturer's instructions.

308

309 **ATAC-seq analysis**

310 ATAC-seq read adaptors were removed using trim_galore. The reads were then mapped to the
311 Arabidopsis thaliana reference genome, TAIR10, using Bowtie2 (-X 2000 -m 1) [30]. Reads of
312 chloroplast and mitochondrial DNA were filtered out and duplicate reads were removed using
313 Samtools [31]. ATAC-Seq open chromatin peaks of each replicate were called using MACS2 with
314 parameters of -p 0.01 --nomodel --shift -100 --extsize 200. Consensus sets of chromatin peaks
315 for all samples were merged by bedtools (v2.26.0) intersect allowing a distance of 10 base pairs
316 [32]. Following this, edgeR was used to define significant changes between peaks [Fold Change,
317 (FC) > 2 and False Discovery Rate, (FDR) < 0.05] [33]. ATAC-seq peak distributions were
318 annotated using ChIPseeker [34]. TF footprints were analyzed by TOBIAS [22] with 572 plant TF
319 motifs downloaded from JASPAR (<http://jaspar.genereg.net/>) [23].

320

321 **RNA-seq analysis**

322 Cleaned short reads were aligned to the reference genome, TAIR10, by Bowtie2 (v2.1.0) [30].
323 Expression abundance was then calculated by RSEM using the default parameters [35].
324 Heatmaps were visualized using the R package pheatmap. Differential expression analysis was
325 conducted using edgeR [33]. A threshold of p-value < 0.05 and Fold Change > 2 were used to
326 decide whether there were any significant differences in expression between samples.

327

328 **ChIP-seq analysis**

329 ChIP-seq data was aligned to the TAIR10 reference genome with Bowtie2 (v2.1.0) [30], only
330 including uniquely mapped reads without any mismatches. Duplicated reads were removed by
331 Samtools. ChIP-seq peaks were called by MACS2 (v2.1.1) and annotated using ChIPseeker [34].
332 Differential peaks were called by the bdgdiff function in MACS2 [36]. ChIP-seq data metaplots
333 were plotted by deeptools (v2.5.1) [37]. Correlation of MORC7 with ChIP-seq data was conducted
334 with ChromHMM [38]. H3K9ac, H3K27ac, H4K16ac, H3K4me1, H3K4me3, H3K36me2,
335 H3K36me3, H3K9me2, H3K27me3, Pol II, and Pol V, as published previously, were included in
336 this analysis [26,39–44] (Supplementary Table 5).

337

338 **Whole-genome bisulfite sequencing (BS-seq) analysis**

339 Previously published whole-genome bisulfite sequencing data for *morc*-mutants and wild type
340 was reanalyzed [6]. Briefly, Trim_galore
341 (http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) was used to trim adapters. BS-
342 seq reads were aligned to the TAIR10 reference genome by BSMAP (v2.90), allowing two
343 mismatches and one best hit (-v 2 -w 1) [45]. Reads with three or more consecutive CHH sites
344 were considered to be unconverted reads and were filtered out. DNA methylation levels were
345 defined as #C/ (#C + #T).

346

347 **Supplementary Information**

348 The online version contains supplementary material available at xxxx.

349

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364

365 **Availability of data and materials**

366 Data supporting the findings of this work are available within the paper and its Supplementary
367 Information files. All high-throughput sequencing data generated in this study are accessible at
368 NCBI's Gene Expression Omnibus (GEO) via GEO Series accession number GSE212801
369 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE212801>). The customized codes used
370 in this study are available upon reasonable request.

371 **Authors' contributions**

372 ZZ, YX, and SEJ conceived the study. ZZ, YX, and CJH performed experiments assisted by MW,
373 ZL, and YK. SF performed high throughput sequencing. YJA and JAW performed IP-MS. ZZ, YX,
374 and SEJ wrote the manuscript with help from all authors. All authors approved the final version of
375 the manuscript and agree on the content and conclusions.

376 **Declarations**

377 **Ethics approval and consent to participate**

378 Not applicable.

379 **Consent for publication**

380 Not applicable.

381 **Competing interests**

382 The authors declare no competing interests.

383

384 **References**

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527

528 **Table**

529 **Table 1** Number of TFs among the proximal genes of MORC7A-unique, MORC7B-unique,
530 MORC7-Pol V Common, and Pol V-unique regions.

Peak	TF	non TF	Total	Percentage	p-value
MORC7A	99	326	425	23.29%	3.12E-36
MORC7B	6	27	33	18.18%	0.002
MORC7-Common	114	1475	1589	7.17%	0.001
Pol V	98	1080	1178	8.32%	1.60E-05
Whole genome	1491	25681	27172	5.49%	NA

531

532

533 **Figure Legends**

534 **Fig 1. MORC7 binds to regions devoid of RdDM. a.** Screenshots of ChIP-seq data for
535 MORC7A-unique, MORC7B-unique, MORC7-Pol V Common, and Pol V-unique regions. **b.**
536 Metaplots of ChIP-seq data for MORC7, Pol V, H3K27ac, H3K4me3, ATAC-seq, and
537 transposable element (TE) density over regions of MORC7A-unique, MORC7B-unique, MORC7-
538 Pol V Common, and Pol V-unique. **c.** Metaplot showing methylation levels of CG, CHG, and CHH,
539 over regions of MORC7A-unique, MORC7B-B unique, MORC7-Pol V Common, and Pol V-unique.
540 **D.** Metaplot showing methylation levels of CG, CHG, and CHH methylation changes (*morchex*
541 minus WT) over regions of MORC7A-unique, MORC7B-unique, MORC7-Pol V Common, and Pol
542 V-unique.

543

544 **Fig 2. MORC7-unique regions preferentially localize to the promoter regions of TFs. a.**
545 Genomic distribution enrichment data for MORC7A-unique, MORC7B-unique, MORC7-Pol V
546 Common, and Pol V-unique regions. **b.** Gene ontology enrichment data for the proximal genes of
547 MORC7A, MORC7-Pol V Common, and Pol V-unique regions. **c.** MORC7 and Pol V binding on
548 promoters of genes in the primary shoot apical meristem specification pathway.

549

550 **Fig. 3 MORC7 associates with some TFs. a.** Volcano plot showing proteins that have significant
551 interactions with MORC7, as detected by crosslinked IP-MS. **b.** Protein-protein interaction
552 networks of MORC7. **c.** A graph showing the degree of overlap between the DAP-seq peaks of
553 approximately 200 TFs with MORC7A-unique, MORC1B-unique, MORC7-Pol V Common, and

554 Pol V unique regions. **d.** Metaplot of PIF4 ChIP-seq data [19] over MORC7A-unique, MORC7B-
555 unique, MORC7-Pol V Common, and Pol V unique regions.

556

557 **Fig. 4 MORC proteins influence TF binding through chromatin compaction. a.** Metaplot
558 showing chromatin accessibility changes in MORC7A-unique, MORC7B-unique, MORC7-Pol V
559 Common, and Pol V-unique regions profiled by ATAC-seq. **b.** A representative screenshot
560 showing higher chromatin accessibility at the promoter of SAUR6 in the *morchex* mutant. **c.**
561 Volcano plot showing changes in TF footprints in MORC7A regions, comparing *morchex* and wild
562 type. **d.** Volcano plot showing changes in TF footprints in MORC7B regions, comparing *morchex*
563 and wild type. **e.** Volcano plot showing changes in TF footprints in MORC7-Pol V Common regions,
564 comparing *morchex* and wild type. **f.** Volcano plot showing changes in TF footprints at Pol V-
565 unique regions, comparing *morchex* and wild type. **g.** Metaplot showing TF footprint changes for
566 MORC7A-unique regions in the *morc4morc7* mutant. **h.** Metaplot showing TF footprint changes
567 for MORC7A-unique regions in the *morc6* mutant. **i.** Metaplot showing TF footprint changes for
568 MORC7A-unique regions in the *morchex* mutant. **j.** Volcano plot showing TF changes for ZF off-
569 target sites, comparing ZF-MORC6 and *fwa-4* plants.

570

571 **Fig. 5 MORC influence TF binding through chromatin compaction.**

572 **a.** Violin plot showing expression levels of genes proximal to MORC7A with Col-0, *morchex*
573 mutant, Ler (wild type background for *lug* mutant), and *lug* mutant. **b.** Expression levels of
574 transcriptional factors: SEP3, PIF4, ARF6/8, TPR1, LUG and SEU (TFs with MORC7A peaks in
575 their promoter regions), with Col-0 and *morchex* mutants following heat treatment. **c.**
576 Transcriptomic changes of *morchex* mutants under normal conditions. **d.** Transcriptomic changes
577 of *morchex* mutants after 30 minutes of heat treatment. **e.** TPL and LUG binding over MORC7A-
578 unique, MORC7B-unique, MORC7-Pol V Common, and Pol V unique regions. **f.** A representative
579 screenshot showing increased binding of TPL and LUG on MORC7A-unique regions in the
580 *morchex* mutant. **g.** Correlation of TPL/LUG binding and ATAC-seq alterations with MORC7
581 binding intensity in *morchex* mutant. **h.** Boxplot showing the expression levels of genes directly
582 regulated by LUG in Col-0 and *morchex* mutants following heat treatment for 30 minutes (T30). **i.**
583 A proposed model of the RdDM-independent functions of MORC proteins.

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587

588 **Supplementary figures:**

589 **Supplementary Fig. 1 Chromatin states of MORC7.** ChIP-seq analysis was performed for
590 H4K16ac, H3K4me3, H3K27ac, H3K9ac, H3Ac, H3K36me3, H3K4me1, H3K36me2, H3K9me2,
591 H3K27me3, Pol II, Pol V, and MORC7.

592

593 **Supplementary Fig. 2** Examples showing MORC7 enrichment over the promoter regions of the
594 TOPLESS genes.

595

596 **Supplementary Fig. 3. MORC7 associates with some TFs. a.** Metaplot of ARF6 ChIP-seq data
597 [20] over MORC7A-unique, MORC7B-unique, MORC7-Pol V Common, and Pol V-unique regions.
598 **b.** Metaplot of TPR1 ChIP-seq data [21] over MORC7A-unique, MORC7B-unique, MORC7-Pol V
599 Common, and Pol V-unique regions. **c.** A screenshot showing MORC7A co-localization with TPR1.

600

601 **Supplementary Fig. 4** Expression levels of genes in the primary shoot apical meristem
602 specification pathway, with and without heat treatment, in Col-0 and *morchex* mutants.

603

604 **Supplementary Tables**

605 **Supplementary Table 1** Genes proximal to MORC7A-unique, MORC7B-unique, MORC7-Pol V
606 Common, and Pol V-unique peaks.

607 **Supplementary Table 2** List of MORC7 interacting proteins.

608 **Supplementary Table 3** Expression level of LUG directly regulated genes in Ler and lug mutant.

609 **Supplementary Table 4** Primers used in this study.

610 **Supplementary Table 5** Published ChIP-seq data used for ChromHMM states analysis.

611

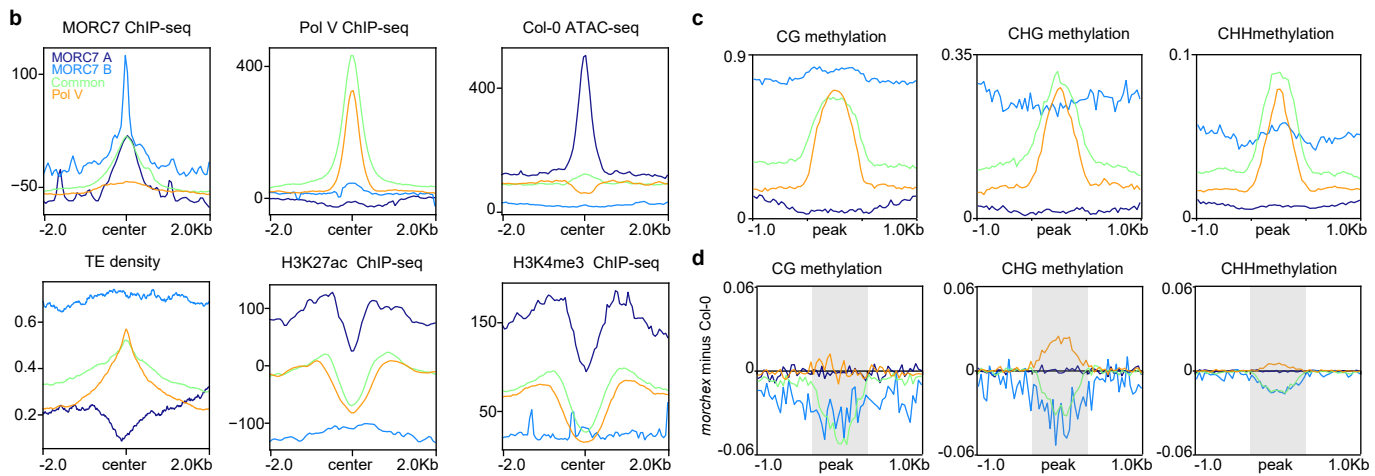
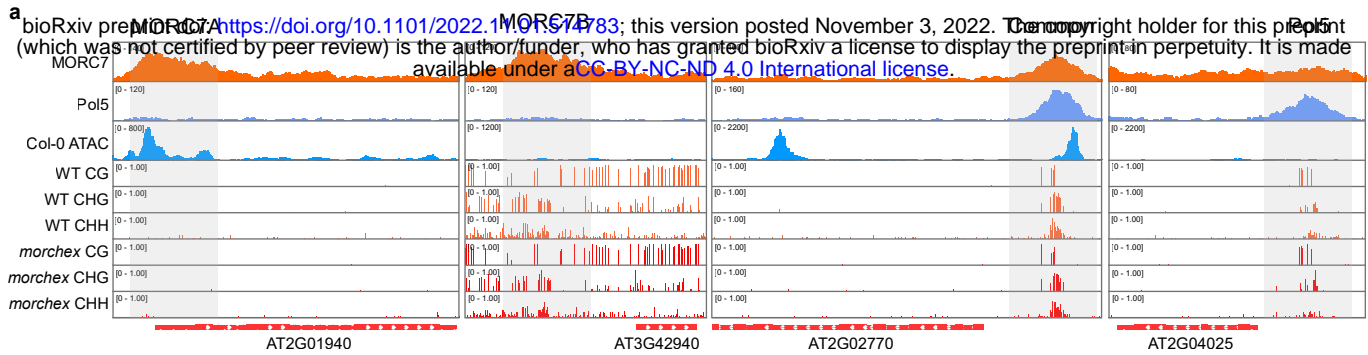


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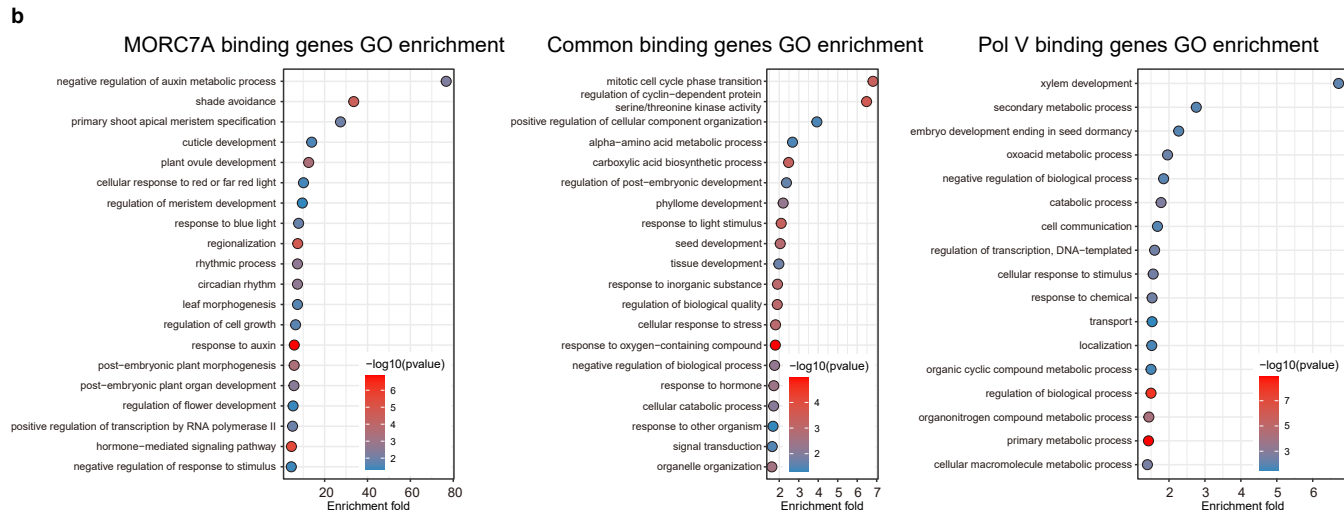
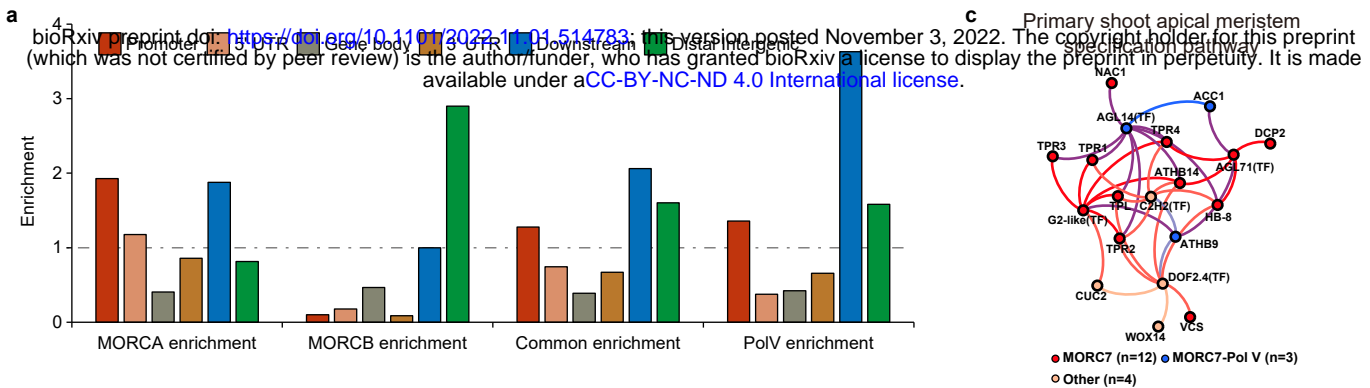


Fig 2. MORC7-unique regions preferentially localize to the promoter regions of TFs. a.

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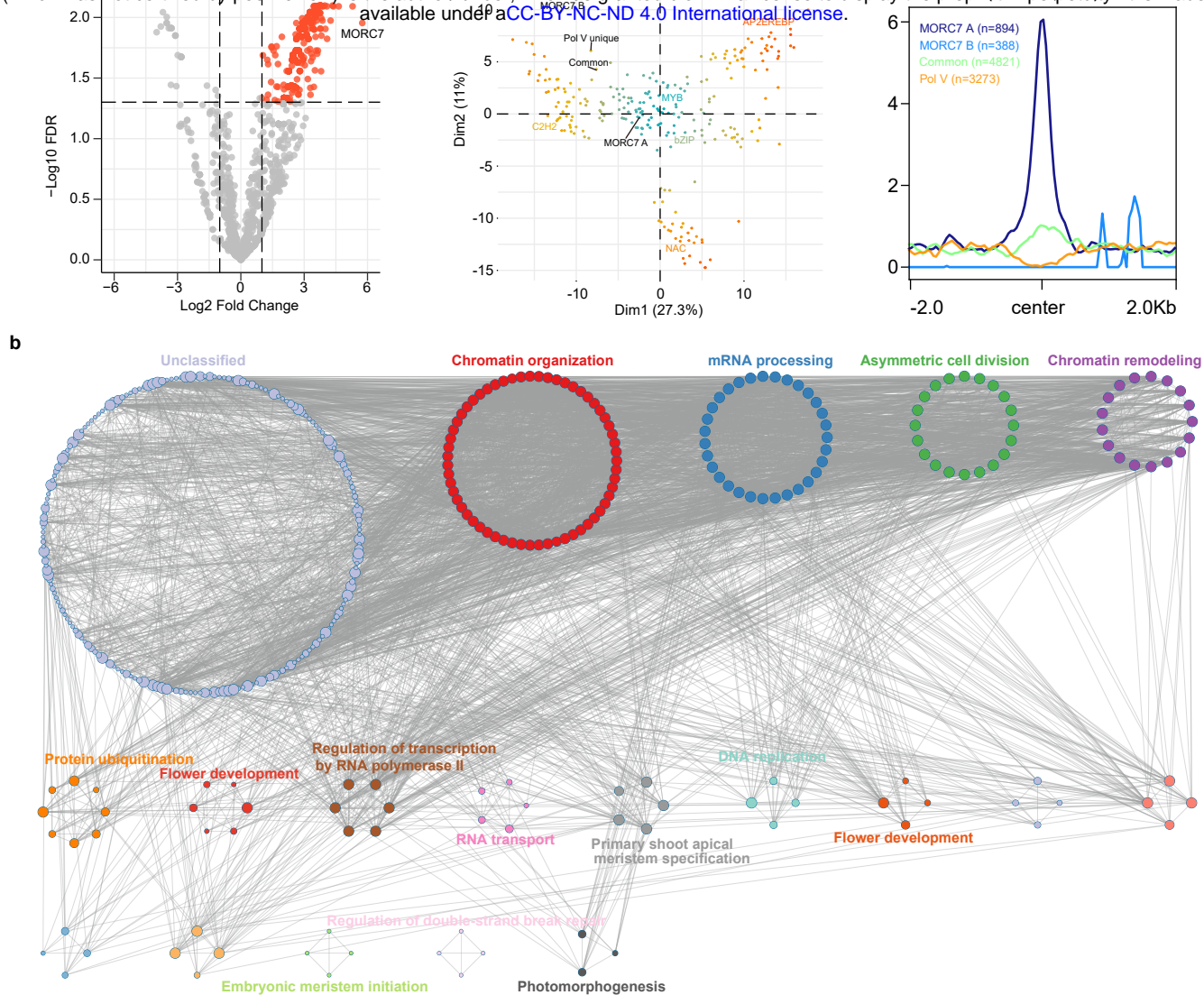


Fig. 3 MORC7 associates with some TFs. **a.** Volcano plot showing proteins that have significant interactions with MORC7, as detected by crosslinked IP-MS. **b.** Protein-protein interaction networks of MORC7. **c.** A graph showing the degree of overlap between the DAP-seq peaks of approximately 200 TFs with MORC7A-unique, MORC7B-unique, MORC7-Pol V Common, and Pol V unique regions. **d.** Metaplot of PIF4 ChIP-seq data [19] over MORC7A-unique, MORC7B-unique, MORC7-Pol V Common, and Pol V unique regions.

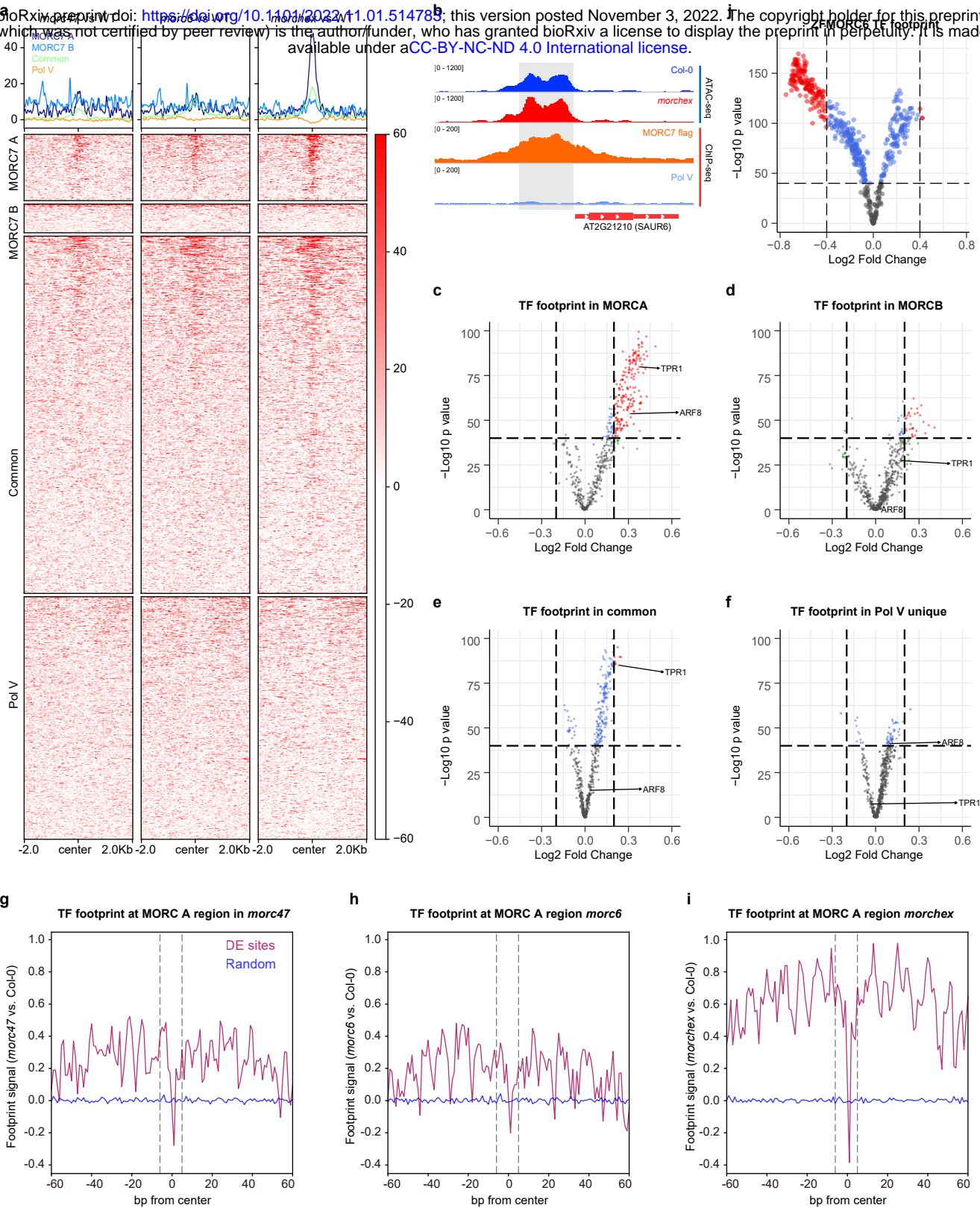


Fig. 4 MORC proteins influence TF binding through chromatin compaction. **a.** Metaplot showing chromatin accessibility changes in MORC7A-unique, MORC7B-unique, MORC7-Pol V Common, and Pol V-unique regions profiled by ATAC-seq. **b.** A representative screenshot showing higher chromatin accessibility at the promoter of SAUR6 in the morchex mutant. **c.** Volcano plot showing changes in TF footprints in MORC7A regions, comparing morchex and wild type. **d.** Volcano plot showing changes in TF footprints in MORC7B regions, comparing morchex and wild type. **e.** Volcano plot showing changes in TF footprints in MORC7-Pol V Common regions, comparing morchex and wild type. **f.** Volcano plot showing changes in TF footprints at Pol V-unique regions, comparing morchex and wild type. **g.** Metaplot showing TF footprint changes for MORC7A-unique regions in the morc4morc7 mutant. **h.** Metaplot showing TF footprint changes for MORC7A-unique regions in the morc6 mutant. **i.** Metaplot showing TF footprint changes for MORC7A-unique regions in the morchex mutant. **j.** Volcano plot showing TF changes for ZF off-target sites, comparing ZF-MORC6 and fwa-4 plants.

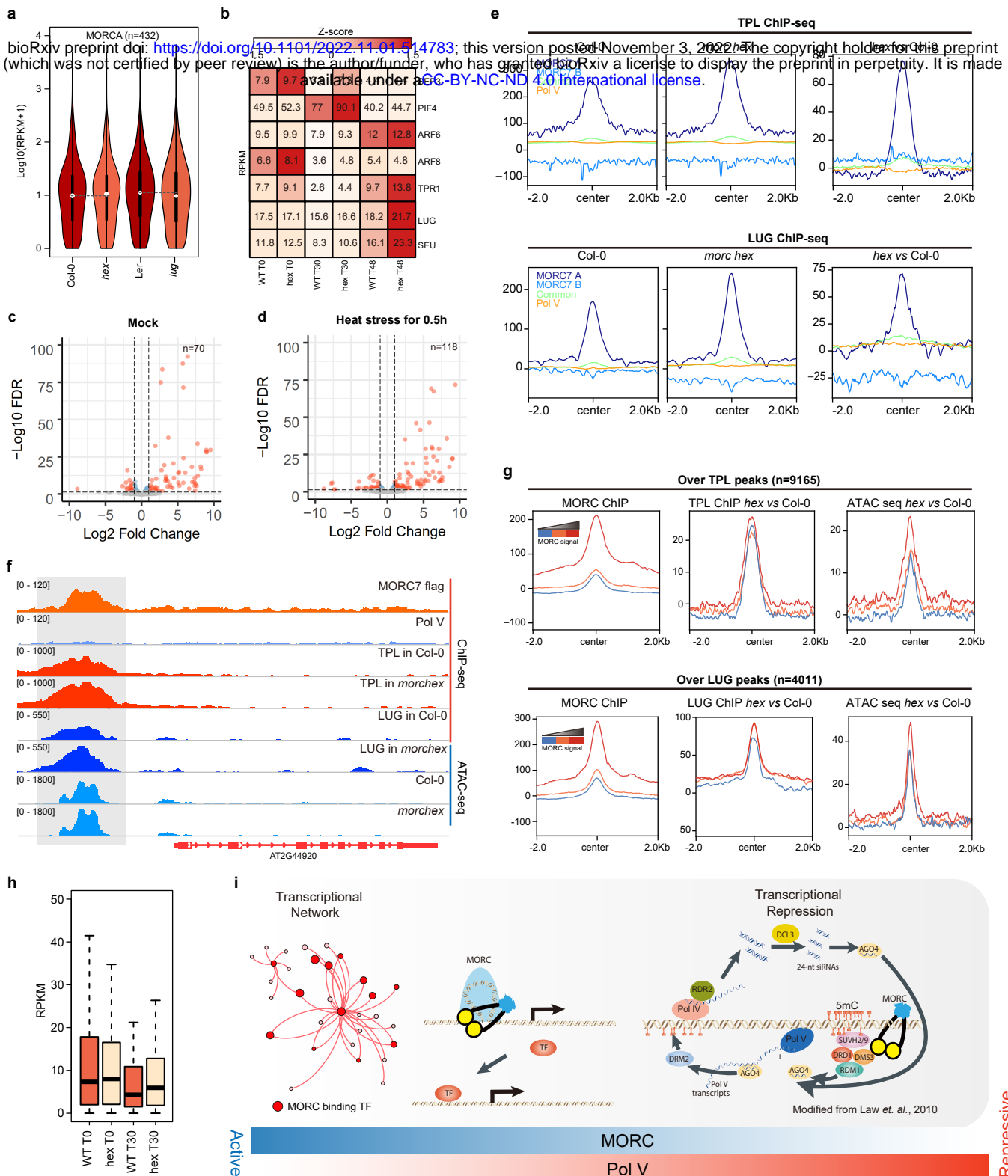


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a. Violin plot showing expression levels of genes proximal to MORC7A with Col-0, morchex mutant, Ler (wild type background for lug mutant), and lug mutant. **b.** Expression levels of transcriptional factors: SEP3, PIF4, ARF6/8, TPR1, LUG and SEU (TFs with MORC7A peaks in their promoter regions), with Col-0 and morchex mutants following heat treatment. **c.** Transcriptomic changes of morchex mutants under normal conditions. **d.** Transcriptomic changes of morchex mutants after 30 minutes of heat treatment. **e.** TPL and LUG binding over MORC7A-unique, MORC7B-unique, MORC7-Pol V Common, and Pol V unique regions. **f.** A representative screenshot showing increased binding of TPL and LUG on MORC7A-unique regions in the morchex mutant. **g.** Correlation of TPL/LUG binding and ATAC-seq alterations with MORC7 binding intensity in morchex mutant. **h.** Boxplot showing the expression levels of genes directly regulated by LUG in Col-0 and morchex mutants following heat treatment for 30 minutes (T30). **i.** A proposed model of the RdDM-independent functions of MORC proteins.

Repressive