SUMMARY

Histone variants play crucial roles in gene expression, genome integrity, and chromosome segregation. We report that the four H2A variants in Arabidopsis define different genomic features, contributing to overall genomic organization. The histone variant H2A.W marks heterochromatin specifically and acts in synergy with heterochromatic marks H3K9me2 and DNA methylation to maintain transposon silencing. In vitro, H2A.W enhances chromatin condensation by promoting fiber-to-fiber interactions via its conserved C-terminal motif. In vivo, H2A.W is required for heterochromatin condensation, demonstrating that H2A.W plays critical roles in heterochromatin organization. Similarities in conserved motifs between H2A.W and another H2A variant in metazoans suggest that plants and animals share common mechanisms for heterochromatin condensation.

INTRODUCTION

All eukaryotic genomes are packaged into chromatin by nucleosomes, which consist of 146 base pairs of DNA wrapped around an octamer of histones H2A, H2B, H3, and H4 (Luger et al., 1997). Chromatin is organized into transcriptionally active euchromatin and more compact inactive heterochromatin. In somatic cells of Arabidopsis, heterochromatin can be visualized as dense structures called chromocenters by DAPI staining (Lysak et al., 2006). In Schizosaccharomyces pombe and mammals, heterochromatin is demarcated by histone H3 lysine 9 methylation (H3K9me3) (Jenuwein and Allis, 2001). Similarly, in Arabidopsis, heterochromatin is marked by H3K9me2, which is deposited by the histone methyltransferases KRYPTONITE (KYP) and SUVH5 and 6 (Bernatavichute et al., 2008; Jackson et al., 2002; Malagnac et al., 2002). DNA methylation plays an important function in regulating heterochromatin (Law and Jacobsen, 2010). The plant-specific DNA methyltransferase CHROMOMETHYLASE 3 (CMT3) binds to H3K9me2 and methylates cytosine residues at CHG sites (where H is A or C or T) (Du et al., 2012; Lindroth et al., 2001). In addition, DNA methylation in CG contexts is carried out by METHYLTRANSFERASE 1 (MET1) (Finnegan and Dennis, 1993; Law and Jacobsen, 2010). DOMAINS REARRANGED METHYLTRANSFERASE 2 (DRM2) and the recently discovered CHROMOMETHYLASE 2 (CMT2) perform methylation in CHH contexts (Haag and Pikaard, 2011; Zemach et al., 2013; Stroud et al., 2014). DNA methylation and H3K9me2 act together to silence genes and transposable elements (TEs) in Arabidopsis (Law and Jacobsen, 2010).

Histone variants carry out diverse chromatin functions, including transcription, DNA repair, and chromosome segregation (Bönnisch and Hake, 2012; Millar, 2013; Szenker et al., 2011; Talbert and Henikoff, 2010). The vast majority of known histone variants are homologs of H2A and H3 histones. A phylogenetic analysis of histone families supports the notion that genes encoding ancestral histone H3.3 and H2A.Z duplicated and evolved to encode histone variants with distinct properties (Talbert et al., 2012). In the case of H3, the cell-cycle-controlled H3.1 class evolved independently in most phyla, acquiring similar properties in plants and animals (Filipescu et al., 2013; Stroud et al., 2012; Wollmann et al., 2012). Similarly, the canonical H2A, thereafter referred to simply as “H2A,” as well as H2A.X, evolved independently from the ancestral H2A.Z in...
almost all eukaryotes (Talbert et al., 2012). In all eukaryotes, the centromeric H3 variant (cenH3) is functionally conserved, specifically recruited at the centromere, and instrumental for kinetochore function (De Rop et al., 2012; Talbert et al., 2008).

Genome-wide profiling of histone variants using chromatin immunoprecipitation followed by deep sequencing (ChIP-seq) revealed that some histone variants decorate specific functional regions of the genome (Talbert and Henikoff, 2010). In yeast and animals, H2A.Z is enriched at transcriptional start sites (TSS) or 5’ ends of genes (Guillemette et al., 2005; Li et al., 2005; Raisner et al., 2005). On the other hand, H3.3 is enriched over gene bodies with inclination toward the 3’ end of genes in Drosophila and humans (Goldberg et al., 2010; Mito et al., 2005). Histone variants also colocalize with posttranslational modifications on their own N-terminal tails. For example, H3.3 tends to be associated with H3K4me2/3, whereas H3.1 tends to be enriched at heterochromatin marked by H3K9me2 (Loyola and Almouzni, 2007). Although H2A.Z and H3.3 are predominantly linked to transcriptional activation, they are also associated with heterochromatin maintenance. In yeast, H2A.Z is enriched in subtelomeric regions and prevents spreading of telomeric heterochromatin into euchromatic regions (Meneghini et al., 2003). Similarly, in humans, H3.3 is localized to telomeric heterochromatin (Goldberg et al., 2010; Wong et al., 2009). In contrast, in Arabidopsis, H2A.Z and H3.3 are excluded from heterochromatin. H2A.Z is enriched at the TSS of expressed genes and also enriched in gene bodies of response genes, which are differentially activated during development or stress (Aceituno et al., 2008; Coleman-Derr and Zilberman, 2012; Zilberman et al., 2009). Similarly, H3.3 enrichment is confined toward the 3’ end of genes and is correlated positively with gene expression (Stroud et al., 2012; Wollmann et al., 2012). Hence, histone variants H2A.Z and H3.3 appear to exclusively define euchromatin in Arabidopsis. This observation led us to question whether a specific combination of histone variants defines heterochromatin in plants, and if so, what their impact might be on chromatin organization.

To understand the roles of histone H2A variants in chromatin regulation in Arabidopsis, we generated a comprehensive localization map of all four types of H2A variants and found that they demarcate different features of the genome. Interestingly, heterochromatin was specifically marked by the previously uncharacterized H2A variant H2A.W. We find that heterochromatin-specific localization of H2A.W does not depend on H3K9me2 or DNA methylation and that H2A.W and DNA methylation cooperatively silence TEs through independent pathways. In vitro nucleosome array experiments suggest that H2A.W causes higher-order chromatin condensation by promoting chromatin fiber-to-fiber interactions. Finally, our findings further demonstrate that H2A.W is both necessary and sufficient for heterochromatin condensation in vivo.

**RESULTS**

**Genome-wide Profiling of H2A Variants Identifies the Heterochromatin-Specific H2A.W**

The Arabidopsis genome comprises 13 genes encoding histone H2A variants, which are grouped into four classes according to C-terminal conserved motifs (Figure S1A available online) (Talbert et al., 2012). We obtained antibodies specific and representative of each H2A variant class (Figure S1B). We performed genome-wide profiling by chromatin immunoprecipitation for H2A variant-bound DNA followed by deep sequencing (ChIP-seq) (Figures 1 and S1C). H2A.X covered the entire genome (Figures 1A and S1C). H2A.X showed a relative depletion at pericentromeric chromatin, as well as over TEs and islands of H3K9me2 present in chromosome arms outside pericentromeric heterochromatin (Figures 1A, 1C, 1D, 1E, and S1C). Both H2A.X and H2A.W were enriched over gene bodies (Figure 1B). In contrast with the rather uniform profiles of H2A and H2A.X over gene bodies, H2A.Z was enriched specifically at the 5’ end of genes (Figure 1B) and was highly depleted over all heterochromatic features (Figures 1C and 1D), as reported previously (Coleman-Derr and Zilberman, 2012). In addition, we analyzed whether enrichment of each H2A variant over gene bodies depends on the level of gene expression. Comparing quintiles of genes defined by their expression level, we observed gradual depletion of H2A and H2A.X and, in turn, enrichment of H2A.Z when expression levels decreased (Figure 1F). Hence, highly expressed genes are defined by a marked enrichment in H2A.Z and H2A at the 5’ end and 3’ end, respectively, whereas inactive genes are covered by H2A.Z.

Unlike all other H2A variants, H2A.W was strongly depleted from gene bodies (Figure 1B). H2A.W was specifically enriched in pericentromeric heterochromatin, TEs, and islands of H3K9me2 (Figures 1A, 1C–1E, and S1C). The Pearson correlation between H3K9me2 and H2A.W within TEs across the genome was 0.7, indicating a very high degree of overlap between H2A.W and H3K9me2. Overall, we conclude that antagonistic localizations of H2A.Z and H2A.W distinguish active and inactive domains of chromatin.

**H2A.W Indexes Heterochromatin**

H2A.W carries an extended C-terminal tail with a conserved SPKK (in general T/SPXK) motif (Figure S1A) (Bönisch and Hake, 2012). The H2A.W variant class evolved and is highly conserved in seed-bearing plants (i.e., gymnosperms and angiosperms) (Figure S2A). In Arabidopsis, the H2A.W family comprises three proteins—H2A.W.6, H2A.W.7, and H2A.W.12—which are encoded by the genes HTA6, HTA7, and HTA12, respectively (Figure S1A). HTA6 is expressed at higher levels than the other two H2A.W homologs (Figure S2D) and was used for the ChIP-seq analyses described above. In order to investigate whether the properties of H2A.W.6 revealed by ChIP-seq analysis were representative of the entire H2A.W class, we analyzed the localization of each member of the H2A.W family in vivo. We fused the RED FLUORESCENT PROTEIN (RFP) to each gene encoding H2A.W and expressed the fusion protein under the control of its respective promoter in transgenic plants that were also expressing the CENTROMERIC HISTONE 3 (CenH3) fused to the GFP. CenH3 is located at the centromere of each chromosome and is surrounded by pericentromeric chromatin that forms heterochromatin domains called chromocenters (Fang and Spector, 2005; Talbert et al., 2002; Fransz et al., 2002). Each H2A.W-RFP localized primarily to domains corresponding to the ten chromocenters, each containing a CENH3-GFP patch in its center (Figures
Figure 1. The Histone Variant H2A.W Is Specifically Localized to Heterochromatin

(A) Distribution of ChIP-seq reads along chromosome 1 for each class of H2A variants and total H3, relative to input DNA. The gray bar at the bottom indicates pericentromeric heterochromatin.

(B) Average enrichment of each class of H2A variants and total H3 over genes delimited by a TSS and a transcription termination site (TTS).

(C) Average enrichment of each class of H2A variants and total H3 over TEs.

(legend continued on next page)
2A, S2B, and S2C). H2A.W.7-RFP and H2A.W.12-RFP also colocalized with H3K9me2 (Figures S2E and S2F), as demonstrated for H2A.W.6 (Figure 2B, wild-type [WT] row, and 1D). In conclusion, all members of the H2A.W family are tightly associated with heterochromatin in somatic cells.

Recruitment of H2A.W Is Independent of H3K9me2 or DNA Methylation
The colocalization between H2A.W and H3K9me2 suggested a possible mechanistic link between the deposition of these two heterochromatic features. Preventing H3K9me2 deposition by mutating the three methyltransferases, KYP, SUVH5, and SUVH6, did not significantly alter the pattern of H2A.W.6 (Figure 2B). Similarly, H2A.W.6 localization was not perturbed significantly by the absence of CMT3, the factor that methylates cytosine residues in CHG contexts at heterochromatic loci marked by H3K9me2 (Figure 2B). Because H2A.W localization does not depend on H3K9me2 or the associated CHG methyl-

Condensation in met1 and ddm1 genetic backgrounds, H2A.W.6 was still colocalized with dispersed heterochromatin marked with H3K9me2 (Figures S2G–S2I). These results suggest that the recruitment or maintenance of H2A.W to heterochromatin depend neither on DNA methylation nor on H3K9me2 and that a mechanism independent from these markers of heterochromatin is responsible for H2A.W deposition. However, other unknown components of heterochromatin might participate in H2A.W localization.

Loss of H2A.W Causes Heterochromatin Decondensation
To study the function of H2A.W in vivo, we obtained knockout lines for all three genes encoding H2A.W (Figure S3A). In single mutants, we did not observe any obvious visible phenotype. Significant growth defects were observed in h2a.w.6 h2a.w.7 and h2a.w.6 h2a.w.12 double mutants, and plant growth was severely affected in h2a.w.6 h2a.w.7 h2a.w.12 (h2a.w) triple mutants (Figure 3A and data not shown). The strong effect of...
h2a.w.6 is likely due to its high level of expression among the three H2A.W encoding genes (Figure S2D). Triple mutant plants failed to flower and did not give rise to the next generation, suggesting that H2A.W is essential. Because H2A.W is specifically localized to heterochromatin, we analyzed heterochromatin structure in nuclei of h2a.w knockout plants. In the double and triple mutants, a significant proportion of nuclei showed variable degrees of physical expansion of heterochromatin marked by H3K9me2 (Figure 3B), reaching in some nuclei a dispersed state in which chromocenters were no longer defined by condensed DAPI staining (Figure 3C). We conclude that H2A.W is required for heterochromatin condensation.

Because heterochromatin was disorganized in h2a.w mutant, we analyzed the localization of the conserved heterochromatic mark H3K9me2 in h2a.w triple mutants. In this genetic background, H3K9me2 was still localized to DAPI-rich regions even in nuclei with dispersed heterochromatic chromocenters, indicating that H3K9me2 deposition does not require H2A.W (Figure S3B). The independent colocalization of H2A.W and H3K9me2 prompted us to examine the genetic interaction between these two marks of heterochromatin. In the H3K9 methyltransferase mutant kyp (Figure S3C), H3K9me2 is reduced but does not show a significant effect on plant growth (Jackson et al., 2002), as SUVH5 and 6 also contribute to H3K9me2

Figure 3. H2A.W Is Required for Heterochromatin Condensation
(A) Morphological phenotypes observed in h2a.w double and triple mutants growing side by side with WT plants. The triple mutant shown in the main panel is magnified in the inset.
(B) Decondensed chromocenters observed in leaf nuclei stained with DAPI and immunostained against the heterochromatin marker H3K9me2. Scale bars represent 2 μm.
(C) The bar chart represents percentages of nuclei belonging to each class of nuclei shown in (B) based on the degree of decondensation of chromocenters observed in DAPI staining. Error bars represent SD.
(D) Morphological phenotype observed in the h2a.w.6 h2a.w.7 kyp mutant growing side by side with a WT plant. The triple mutant shown in the main panel is magnified in the inset.
See also Figure S3.
We found that simultaneous mutation of KYP and H2A.W in h2a.w.6, h2a.w.7, kyp plants caused severe growth defects compared to growth defects observed in h2a.w.6, h2a.w.7 mutant plants (Figure 3D). Similarly, we also observed an increase in the proportion of nuclei with decondensed heterochromatin (Figure 3C). These results indicate that H2A.W and KYP participate in parallel pathways controlling heterochromatin condensation.

H2A.W Represses Transposable Elements in Synergy with CMT3-Mediated CHG Methylation

Decondensation of heterochromatin in mutants such as met1 has been previously shown to be associated with increased transcription of TEs and genes in the proximity of heterochromatic loci (Vaillant et al., 2008; Reinders et al., 2009; Law and Jacobsen, 2010). However, RNA sequencing (RNA-seq) analyses of the h2a.w triple mutant did not show a substantial impact on expression of TEs (Figure S4A). This suggests that, in the absence of H2A.W, other silencing pathways might play compensatory roles in maintaining the transcriptional repressive status of TEs. Indeed, DNA methylation analysis by whole-genome bisulfite sequencing (BS-seq) in the h2a.w triple mutant showed a significant increase in CHG methylation over TEs and regions enriched in H3K9me2, whereas DNA methylation over gene bodies was not affected (Figure 4A). A notable upregulation of the expression of CMT3 encoding the primary CHG DNA methyltransferase likely accounts for the increase in CHG methylation (Figure 4B). DNA methylation at CG and CHH sites was not significantly affected, suggesting a specific association of H2A.W with CHG methylation (Figure 4A). These results suggested the possibility that increased CHG methylation may compensate for the lack of H2A.W, resulting in maintenance of the transcriptional repression of TEs. To test this hypothesis, we combined the h2a.w.6, h2a.w.7, and cmt3-11 null mutant alleles. This combination led to male and female gamete abortion (Figures S4B and S4C), and we were unable to obtain any triple homozygous plants out of 704 progeny segregating from the self-fertilized triple heterozygous mutant, indicating that CMT3 and H2A.W pathways are synergistically required for viability. CMT3 methylates heterochromatic DNA by binding to H3K9me2 (Du et al., 2012), and consistently, CHG methylation is reduced in kyp mutants (Jackson et al., 2002; Stroud et al., 2013). Accordingly, the synergistic action of H2A.W and KYP was also observed on expression of solo long terminal repeat

Figure 4. H2A.W Represses the Transcription of Heterochromatic Elements in Synergy with DNA Methylation Mediated by CMT3

(A) Metaplot analysis from whole-genome bisulfite sequencing showing different types of methylation levels in WT and h2a.w.6 h2a.w.7 h2a.w.12 triple mutant over major genomic features. Graphs were plotted as in Figure 1A.

(B) Expression levels of CMT3 in h2a.w.6 h2a.w.7 compared to WT. Expression values were obtained from RNA-seq data.

(C) Expression levels of SDC and solo LTR in h2a.w.6 h2a.w.7 kyp mutant compared with WT. The data from quantitative real-time RT-PCR were normalized to ACTIN7. Error bars represent SD based on three independent biological replicates. See also Figure S4.
TE and the gene SDC, which were both dramatically increased in the h2a.w.6 h2a.w.7 kyp triple mutant (Figure 4C). These data suggest that genetic pathways of H2A.W and CHG methylation interact to ensure heterochromatin integrity.

H2A.W Promotes Chromatin Condensation through Its Conserved C-Terminal Motif In Vitro

To further understand the potential mechanism of H2A.W in chromatin condensation, we tested whether H2A.W is able to cause condensation of nucleosome arrays in vitro. Chromatin condensation in vitro is composed of two independent structural transitions. Nucleosomes undergo short-range interactions with neighboring nucleosomes to form locally folded chromatin fibers. In addition, chromatin fibers self-associate to form supramolecular oligomers that share many of the properties of large-scale structure in condensed chromosomes. These two reversible structural transitions rely on different types of nucleosome-nucleosome interactions (Hansen, 2002) and can be studied in a model system by analyzing the magnesium-dependent solution-state properties of defined 12-mer nucleosomal arrays in vitro. To understand the mechanism of H2A.W in chromatin condensation in vitro, we tested whether H2A.W affects self-association or folding. We assembled nucleosome arrays with Arabidopsis H2A.1, H2A.W.6, or a truncated version of H2A.W.6 (H2A.W-CT) that lacks the C-terminal tail containing the SPKK motif (Figure 5). Because folding and self-association depend on the number of nucleosomes per array, the three types of arrays were closely matched for nucleosome saturation. We ensured that all arrays contain on average 11 to 12 nucleosomes per DNA molecule, i.e., every nucleosome positioning sequence repeat is occupied (Figures 5A and 5B). We also verified equimolar distribution of all histone components by SDS-PAGE (Figure 5C). We then determined the extent of self-association as a function of salt concentration and found that H2A.W.6 arrays engaged in oligomerization at significantly lower MgCl2 concentrations than H2A.1 or H2A.W.6-CT (Figure 5E). This suggests that H2A.W.6 promotes long-range nucleosome-nucleosome interactions in chromosomes and that the C-terminal tail of H2A.W.6 contributes to this behavior. Intriguingly, both arrays with H2A.W.6 and H2A.W.6-CT exhibited no sign of folding at 1 mM MgCl2, whereas arrays with canonical H2A.1...
behave like other nucleosomal arrays and were moderately folded under these conditions (Figure 5D). Altogether, this suggests that H2A.W is able to enhance large-scale chromatin condensation by promoting long-range chromatin fiber-to-fiber interactions through its conserved C-terminal tail but is deficient in the short-range interactions that form the 30 nm fiber, suggesting a role in promoting highly condensed heterochromatin in vivo.

H2A.W Promotes Chromatin Condensation In Vivo

Because there was a gain of DNA methylation upon loss of H2A.W in h2a.w triple mutants (Figure 4A), it was difficult to address directly whether H2A.W is sufficient to promote heterochromatin condensation in somatic cells. However, we observed that none of the genes encoding H2A.W were expressed in the central cell, one of the two female gametes of flowering plants (Figures S5A–S5C). The absence of H2A.W in this cell type correlated with the complete absence of condensed heterochromatic domains as indicated by the uniform distribution of H2A.X.5-RFP (Figure 6A), which marks equally heterochromatin and euchromatin in somatic cells (Figures 1A–1D). In order to test whether the lack of H2A.W was responsible for the absence of condensed heterochromatin in the central cell nucleus, we ectopically expressed H2A.W.6 using the promoter of FWA, which is specifically active in the central cell and the endosperm (Kinoshita et al., 2004). The chromatin of the central cell expressing pFWA-H2A.W.6-RFP showed domains of varying intensity with distinct zones of higher condensation (Figure 6B). We could distinguish two types of organization. A minor group of nuclei showed one or two strong patches of dense chromatin. The most representative class of central cell nuclei showed several patches of dense chromatin assembled in the periphery of the nucleus (Figure 6B), contrasting with the uniform WT pattern of H2A.X-RFP that marks both euchromatin and heterochromatin (Figure 6A). To test whether the assembled condensed domains are heterochromatic, we immunostained whole ovules against H2A.X to report the general chromatin organization and against H3K9me2 to mark heterochromatin. We also used the POLYDACTYL ZINC FINGER (PZF)-GFP that binds to centromeric 180 bp repeats in somatic cells (Lindhout et al., 2007). In the WT, chromocenters marked by chromatin condensation and H3K9me2 were clearly observed in the egg cell nucleus (Figures 6C), whereas the central cell chromatin organization was more diffuse with elongated patches of H3K9me2 lining the periphery of the nucleus (Figure 6C). In contrast, central cells expressing H2A.W-6-RFP clearly showed dense chromatin domains marked by H3K9me2, indicating that these domains are heterochromatic. In addition, within these domains, we detected a single dot of centromeric marker PZF-GFP (Figure 6D, bottom), indicating that condensed chromatin domains caused by ectopic H2A.W are indeed pericentromeric heterochromatin foci. The absence of visible dots of PZF-GFP in WT central cell nuclei (Figure 6D, WT) was likely explained by the decondensed status of the pericentromeric and centromeric domains in the central cell and the absence of the cenH3 assembly reported previously (Ingouff et al., 2010). These observations indicate that H2A.W is sufficient to cause heterochromatin condensation in vivo.

DISCUSSION

Comprehensive genomic profiling showed that each H2A variant is characterized by a specific profile. In contrast to H2A, H2A.Z, and H2A.W, the distribution of H2A.X is relatively ubiquitous, which is in keeping with the idea that H2A.X is involved in DNA repair (Downs et al., 2007; Soria et al., 2012). H2A uniformly marks gene bodies similar to that described for H3.1 (Stroud et al., 2012; Wollmann et al., 2012). Transcription of both H3.1 and H2A is highly coupled with S phase (Talbert et al., 2012), suggesting that H2A and H3.1 are likely subject to common dynamics related to the loading of new nucleosomes over replicated DNA. However, unlike H3.1, H2A enrichment over gene bodies positively correlates with transcription with a gradual replacement by H2A.Z in genes expressed at low levels, suggesting that transcription is coupled to an exchange between H2A and H2A.Z over gene bodies. We confirmed that, in highly expressed genes, H2A.Z enrichment is confined to the 5’ end of the gene bodies, similar to that described in yeast, Drosophila, and human (Zilberman et al., 2008). This enrichment appears to be a mirror image of H3.3 distribution, confined to the 3’ end of gene bodies (Stroud et al., 2012; Wollmann et al., 2012). Hence, the boundaries of gene bodies appear to be defined by differential enrichment of H2A, H2A.Z, and H3.3 variants. In sharp contrast to euchromatin, heterochromatin depletion is lowered down by H2A.W and H2A.Z and is occupied by H2A.W, suggesting that H2A variants restrict the deposition of each other over larger genomic domains defining euchromatin and heterochromatin. Therefore, combinatorial localizations of H2A and H3 variants define the main genomic features in Arabidopsis.

We show that H2A.W recruitment to chromatin is independent of the pathways that deposit heterochromatric marks H3K9me2 and DNA methylation. However, H2A.W acts together with these pathways to maintain heterochromatin condensation into higher-order domains and to silence TEs. The lethal genetic interaction observed in our study suggests that H2A.W and H3K9me2/CHG methylation are absolutely essential for plant life. The lethality observed in mutants deficient for both pathways likely results from a lack of heterochromatin integrity, which could impact cell division. In response to H2A.W depletion, we also observe a crosstalk between these pathways as both CMT3 expression and CHG methylation increase in heterochromatin.

In vitro micrococcal nuclease digestion data showed that nucleosomes containing H2A.W protect a longer stretch of DNA (162 bp) than nucleosomes containing H2A (146 bp) (Lindsey et al., 1991), suggesting that H2A.W prevents accessibility to DNA. Consistently, we find that H2A.W represses TE expression in synergistic action with H3K9me2. The acidic patch present in the histone core region of H2A affects the folding ability of nucleosome arrays (Kalashnikova et al., 2013). For example, euchromatic H2A.Z shows higher chromatin folding than H2A as a consequence of its extended acidic patch (Fan et al., 2004). The acidic patch in H2A.W and canonical H2A are identical, and H2A.W shows a lower folding capacity than H2A, suggesting that other domains of H2A participate in chromatin folding. Our results show that H2A.W promotes chromatin condensation through long-range fiber-to-fiber interactions in vitro. These
Figure 6. H2A.W Is Sufficient to Cause Chromatin Condensation In Vivo

(A) Uniform pattern of pH2A.X.5-H2A.X.5-RFP expressed in the central cell nucleus indicating the absence of defined chromocenters. N points the nucleolus.

(B) Impact of ectopic expression of H2A.W.6-RFP in the central cell nucleus (CCN). We observe mild or severe chromatin condensation (arrowheads), and the proportion of each type of nuclei is indicated in the top right of each image. N points the nucleolus.

(C) Impact of ectopic expression of H2A.W.6-RFP in the CCN on H2A.X and H3K9me2 immunolocalization. Top: the egg cell nucleus (ECN) shows well-defined chromocenters reported by H2A.X condensation and marked with H3K9me2 and serves as a reference point for comparison with the CCN. Middle: ectopic expression of H2A.W.6 in the CCN causes condensation of chromatin domains reported by H2A.X. Bottom: these domains are marked with H3K9me2 (lower panel).

(D) A CCN-expressing PZF-GFP that recognizes the centromeric 180 bp repeats. Top: in the WT, PZF-GFP marks condensed centromeres in somatic cells (S, insert, a nucleus from root cells) but does not mark any condensed structure in the CCN. Bottom: in a central cell expressing pFWA::H2A.W.6-RFP, arrowheads point the position of condensed chromatin domains, which contain a dot of PZF-GFP marking the centromeric location. Scale bars in confocal sections represent 5 μm.

See also Figure S5.
data are supported by the contrasting impacts of loss or gain of function of H2A.W in vivo on heterochromatin organization. We propose that H2A.W is necessary and sufficient to organize heterochromatin into higher-order functional domains such as chromocenters, which are essential for kinetochore assembly and mitosis (Bian and Belmont, 2012; Luger et al., 2012).

H2A.W-specific chromatin condensation properties rely on the extended C-terminal domain that contains the conserved SPKK motif. This is consistent with previous in vitro studies of the function of the SPKK motif in other histone variants (Khadake and Rao, 1997). The SPKK motif binds preferentially to the minor groove of DNA at A/T rich sites (Churchill and Suzuki, 1989), which are generally enriched in satellite repeats of heterochromatin (Brutlag, 1980). During evolution, the SPKK motif not only became associated with H2A.W in seed plants but was also recruited to other histones, which are associated with condensed chromatin, the linker H1 in most eukaryotes, and the sea urchin testis-specific H2B (Suzuki, 1989; Buttinelli et al., 1999; Poccia and Green, 1992). Motifs similar to SPKK are also present in the linker region of the vertebrate-specific macroH2A that separates the core histone domain from the C-terminal macrodomain (Figure S6). MacroH2A is highly enriched on the X chromosome and is linked to transcriptional repression (Costanzi and Pehrson, 1998; Gamble et al., 2010; Gaspar-Maia et al., 2013). In vitro, the linker region of macroH2A causes chromatin condensation (Muthurajan et al., 2011), and macroH2A devoid of the macrodomain stabilizes nucleosomes (Chakravarthy et al., 2012). However, how macroH2A affects chromatin architecture in vivo remains unclear. The similarity between H2A.W and macroH2A devoid of its macrodomain suggests that the properties of H2A.W described in this study may reveal conserved aspects of histone variant functions that may not only aid in our understanding of plant heterochromatin but also shed light on mechanisms controlling higher-order heterochromatin condensation in other eukaryotes.

**EXPERIMENTAL PROCEDURES**

**Plant Material and Growth Conditions**

All mutants are in Columbia-0 (Col) ecotype. h2a.w.6 (SALK_024544.32), h2a.w.7 (GK_149508), and h2a.w.12 (SAIL_667) T-DNA lines were obtained from ABRC at Ohio State University and NASC at The University of Nottingham, respectively. drm2-2 (SALK_150863.37.35), cmrt3-11 (SALK_148381), kyp-6 (SALK_041474), suvh4.5,6,7 triple mutant (SALK_041474, Gabi_263C05, SAIL_1244_F04), cmrt3-11 (SALK_148381), dmr2-2 (SALK_150863), met1-3, and the EMS-induced SNP in ddm1-2, met1-3, ddm1-2, and kyp suvh5 suvh6 were previously described (Johnson et al., 2008; Saze et al., 2003; Vongs et al., 1993). The 180 bp repeat reporter PDZ-GFP was a kind gift from B. van der Zaal. h2a.w.6 and h2a.w.7 T-DNA insertions were confirmed by PCR-based genotyping. Primer sequences are described in Table S1. Arabidopsis plants were grown under long day conditions—i.e., 16 hr light and 8 hr dark.

**H2A Variants CHIP-Seq Analyses**

CHIP was performed as previously described (Wollmann et al., 2012) with few modifications. Two grams of seedlings were ground in liquid nitrogen and fixed in 1% formaldehyde. Sonicated chromatin was incubated overnight with anti-H2A, anti-H2A.X, anti-H2A.W.6, anti-H2A.Z.9, and anti-H3 (ab1791, Abcam) antibodies or IgG (ab46540-1, Abcam). Immunoprecipitated DNA was treated with RNase A (Fermentas) and purified with the QiAquick purification kit (QiAGEN). CHIP-seq libraries were generated as per manufacturer’s instructions (Illumina). Libraries were sequenced on a HiSeq2000 (Illumina). Reads were mapped to the TAIR8 genome, and analysis was performed as previously described (Stroud et al., 2012).

**DNA Methylation Analyses**

For genome-wide bisulfite sequencing (BS-seq), plants were grown under continuous light, and samples were collected from 3-week-old plants. Genomic DNA was extracted using Plant DNeasy Mini kit (QiAGEN). BS-seq libraries were generated, sequenced, and analyzed exactly as previously described (Stroud et al., 2013), except that reads were mapped to the TAIR8 genome.

**Nucleosome Array Assembly**

Nucleosomal arrays were reconstituted onto a DNA fragment that comprises 12 repeats of 207 base pairs “601” positioning sequence (Lowary and Widom, 1998). A two-step reconstitution method was applied to assemble nucleosome arrays (Muthurajan et al., 2011). First, different amounts of (H3-H4)2 tetramers were titrated to DNA. The tetrasomes were reconstituted by step dialysis (4–6 hr each step) starting from 1 M NaCl to 0.75 M NaCl and finally 2.5 M NaCl (in a buffer containing 10 mM Tris [pH 7.5] and 0.25 mM EDTA). The best ratio between H3-H4 and DNA for tetrasome formation was determined by sedimentation velocity AUC. Once the S20.w midpoint of reconstitution reached around 19S, the ratio of tetramer to DNA was fixed for the next step. Varying amounts of variant H2A.H2B dimers were added to the optimized ratio of tetramer to DNA. The saturation levels of arrays were determined by sedimentation velocity AUC (SV-AUC). To reach a S20.w of 29S (characteristic of a fully saturated array), the ratio of dimer to tetrasome was around 2.0. EcoRI digestion also was used to monitor the saturation status of nucleosome arrays as described previously (Muthurajan et al., 2011).

**Self-Association Assay**

Nucleosomal arrays (the equivalent of ~1 μg DNA) were combined with various concentrations of MgCl2 (from 0 to 5 mM). The mixtures were incubated at room temperature for 5 min and then spun at 14,000 g for 10 min. The percentage of arrays remaining in the supernatant was determined at 259 nm and was plotted against the concentration of MgCl2.

**Folding Assay by Sedimentation Velocity**

Nucleosome arrays containing ~10 μg DNA were treated with 1 mM MgCl2-SV-AUC was applied to monitor the impact of MgCl2 on array folding. Sedimentation velocity experiments were done in a Beckman XL-A or XL-I ultracentrifuge at 20,000 rpm and 20°C. The absorbance at 259 nm was scanned continuously with a radial increment of 0.003 cm. All results were processed and analyzed by UltraScanIII. By the improved van Holde-Weischat method in UltraScanIII, the diffusion-corrected integral distribution (Q(s)) of sedimentation coefficients over the entire boundaries was analyzed and plotted.

**ACCESSION NUMBERS**

The sequencing data are deposited at Gene Expression Omnibus ( GEO) with the accession number GSE50942.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, six figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2014.06.006.

**AUTHOR CONTRIBUTIONS**

K.Z., U.M.M., and K.L. performed experiments reported in Figures 5 and S5; S.F. and X.Z. performed the first series of CHIP-seq for H2A.W.6; S.F. performed genome-wide bisulfite sequencing; H.S. performed the RNA-seq and all bioinformatic analyses; S.H. participated in H2A variants CHIP; S.H., T.K., and F.B. produced the data shown in Figures 6 and S6; X.N. performed preliminary transcriptome analyses; M.G. provided Figure S2H; and R.Y. performed...
all other experiments and analyses. F.B. conceived the initial project, which was developed with the help of a fruitful collaboration with the S.E.J. lab. R.Y., F.B., and S.E.J. wrote the manuscript with inputs from all other authors.

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Cloning of pH2A::H2A-RFP for Making Transgenic Plants and Generation of Constructs of H2As for Bacterial Protein Expression

For each of the H2A.W genes, a genomic fragment containing the promoter region until the codon before the termination codon was amplified with primers attB1-F-H2A and attB2-R-H2A (Table S1) using the KOD-plus-PCR kit (Toyobo). PCR fragments were cloned directionally after a BP reaction (Invitrogen) into the plasmid pDONR/Zeo (Invitrogen) to generate a pDONR/Zeo-H2A entry vector. Recombination LR reactions were performed between a pAlli-2 destination vector and the pDONR/Zeo-H2A vector to generate a recombined destination plasmid with an H2A gene with its N-terminal fused to a gene encoding red fluorescent protein (pH2A::H2A-RFP). These constructs were transformed into WT (Col-0 accession) plants by Agrobacterium tumefaciens–mediated floral dip method. Primary transformants were selected based on GFP positive signal in seeds. For bacterial protein expression constructs, protein-coding sequences were amplified with primers described in Table S1 using cDNA as template. The fragments were digested with Ndel and BamHI and cloned in to pET3A vector.

H2A Class-Specific Antibodies
All anti-H2As class specific antibodies are affinity-purified rabbit polyclonal antibodies made by GenScript USA Inc (Piscataway, NJ) against peptides CPKKAGASKPSADE, GGRKPPGAPKTKSVC, CKVGNKGDIGSASQ, and KPSGSDKDKKKPC for H2A, H2A.W, H2A.X, and H2A.Z, respectively. Antibodies are characterized by Western blot on total protein extracts for their specificity. For all samples, proteins are extracted from 10-day old seedlings except for H2A.Z.8-RFP and H2A.W.12-RFP. For H2A.Z.8-RFP protein extract is made from shoot meristem and for H2A.W.12-RFP protein extract is made from ovules. For detecting RFP and H3, anti-DsRed (Clontech, Cat.No.632496) and anti-H3 (Abcam, Cat.No. ab1791) antibodies were used.

Immunofluorescence
Immunofluorescence experiments examining the protein localization and chromatin condensation in the nuclei were performed as previously described (Jacob et al., 2009), with the following modifications. Leaves from three-week-old plants (For mutant nuclei phenotyping, leaves were collected from first generation plants. For H2A.W.7-RFP, H2A.W.12-RFP, and H3K9me2 colocalization, nuclei were isolated from roots) were fixed in 4% paraformaldehyde in TRIS buffer (10 mM TRIS pH 7.5, 10 mM EDTA, 100 mM NaCl) for 20 min and washed twice in TRIS buffer. Leaves were chopped in 400 μl lysis buffer (15 mM TRIS pH 7.5, 2 mM EDTA, 0.5 mM spermine, 80 mM KCI, 20 mM NaCl, 0.1% Triton X-100) and filtered through a 40-micron cell strainer. Nuclei were concentrated by centrifugation at 500 g for 3 min. Nuclei pellet was resuspended in 100 μl of lysis buffer and from this 10μl was air-dried on coverslips. Nuclei were post-fixed in 4% paraformaldehyde in PBS for 30 min. Coverslips were washed three times in PBS and incubated in blocking buffer (1% BSA in PBS) for 30 min at 37°C. Nuclei were incubated at 4°C overnight in respective antibodies (Mouse Anti-H3K9me2 Abcam ab1220, 1:100; Rabbit Anti-H2A.W.6, 1:100). Coverslips were washed three times with PBS and incubated with goat anti-mouse 488 and anti-rabbit 555 (Invitrogen, 1:200) for 30 min at 37°C. Following three PBS washes, nuclei were counterstained and mounted in Vectashield mounting media with DAPI (Vector H-1200).

Whole-mount immunostaining of ovules was performed as described (Pilot et al., 2010).

RNA Analyses
Total RNA was prepared with RNeasy Mini kit (QIAGEN, Chatsworth, CA) from ten-day-old seedlings. DNase treatment was done on 2 μg of total RNA with DNase Free Kit (Ambion, Austin, TX). RNA-seq was performed using TruSeq RNA Sample Prep Kit (Illumina) and sequenced on a HiSeq 2000 (Ilumina). RNA-seq reads were processed and analyzed as previously described, where only uniquely mapping reads were retained and differentially expressed TEs and genes were defined with a four-fold cutoff and p < 0.05 as described (Stroud et al., 2012). Reads were mapped to the TAIR8 genome. Expression values of genes and TEs were calculated as RNA-seq reads per kilobase per million mapping reads.

For characterizing mutants, total RNA was isolated with Trizol (Life Technologies, Carlsbad, CA) from 100 mg fresh weight of 10-day old seedlings. 2 μg of total RNA were treated with DNaseI (Life Technologies, Carlsbad, CA) and half of the DNaseI-treated samples, proteins are extracted from 10-day old seedlings except for H2A.Z.8-RFP and H2A.W.12-RFP. For H2A.Z.8-RFP protein extract is made from shoot meristem and for H2A.W.12-RFP protein extract is made from ovules. For detecting RFP and H3, anti-DsRed (Clontech, Cat.No.632496) and anti-H3 (Abcam, Cat.No. ab1791) antibodies were used.

Immunofluorescence experiments examining the protein localization and chromatin condensation in the nuclei were performed as previously described (Jacob et al., 2009), with the following modifications. Leaves from three-week-old plants (For mutant nuclei phenotyping, leaves were collected from first generation plants. For H2A.W.7-RFP, H2A.W.12-RFP, and H3K9me2 colocalization, nuclei were isolated from roots) were fixed in 4% paraformaldehyde in TRIS buffer (10 mM TRIS pH 7.5, 10 mM EDTA, 100 mM NaCl) for 20 min and washed twice in TRIS buffer. Leaves were chopped in 400 μl lysis buffer (15 mM TRIS pH 7.5, 2 mM EDTA, 0.5 mM spermine, 80 mM KCI, 20 mM NaCl, 0.1% Triton X-100) and filtered through a 40-micron cell strainer. Nuclei were concentrated by centrifugation at 500 g for 3 min. Nuclei pellet was resuspended in 100 μl of lysis buffer and from this 10μl was air-dried on coverslips. Nuclei were post-fixed in 4% paraformaldehyde in PBS for 30 min. Coverslips were washed three times in PBS and incubated in blocking buffer (1% BSA in PBS) for 30 min at 37°C. Nuclei were incubated at 4°C overnight in respective antibodies (Mouse Anti-H3K9me2 Abcam ab1220, 1:100; Rabbit Anti-H2A.W.6, 1:100). Coverslips were washed three times with PBS and incubated with goat anti-mouse 488 and anti-rabbit 555 (Invitrogen, 1:200) for 30 min at 37°C. Following three PBS washes, nuclei were counterstained and mounted in Vectashield mounting media with DAPI (Vector H-1200).

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For characterizing mutants, total RNA was isolated with Trizol (Life Technologies, Carlsbad, CA) from 100 mg fresh weight of 10-day old seedlings. 2 μg of total RNA were treated with DNasel (Life Technologies, Carlsbad, CA) and half of the DNasel-treated RNA was used for cDNA synthesis using SuperScript III First-Strand Synthesis System (Life Technologies, Carlsbad, CA). 1 μl of cDNA (+RT) or 1 μl of DNasel-treated total RNA (-RT) were used for RT-PCR analysis of WT and mutant alleles with primers flanking the EMS-induced SNP in ddm1-2 or sites of T-DNA insertion in the other mutants. Ubiquitin (Ub10) or GAPDH was used as control for the RT-PCR.

Real-time PCR assays were performed using a PCR Master Mix (Applied Biosystems, Foster City, CA) according to manufacturer’s instructions using 1 μl of RT product. The specificity of the amplification product was determined by performing a dissociation curve analysis. The PCR reaction and quantitative measurements were done on 7900HT Fast Real-Time PCR System (Applied Biosystems, Foster City, CA). We performed three biological replicates, with three technical replicates for each sample. For each PCR reaction, the ΔCt was calculated using ACT7 gene as endogenous control. Relative Quantitation values (RQ) were calculated using the 2^(-ΔΔCt) method (RQ = 2^(-ΔΔCt)).
ChIP-Seq Analyses of H2As

ChIP was performed as previously described with few modifications (Wollmann et al., 2012). 2 g of leaves were ground in liquid nitrogen and fixed in 1% formaldehyde for 10 min. The reaction was stopped by adding 0.125 M glycine. Nuclei were extracted by filtration through Miracloth and iteratively washed and centrifuged at 2,000 g. Nuclei were lysed in SDS buffer. DNA was sonicated for 10 cycles of 0.5 min on and 1 min off with an UCD-200TM-EX Bioruptor (Diagenode) on medium power, at 0 to 4°C. Sonicated chromatin was incubated overnight with either anti-H2As, anti-H3 (ab1791, Abcam) antibodies, or IgG (ab46540-1, Abcam). After pre-clearing, magnetic protein A-beads (Dynabeads protein A, Invitrogen) were incubated with the antibodies-chromatin mix for 3 hr. After precipitation of the beads on a magnetic rack (MagnaRack, Invitrogen) and washes with increasing stringency, DNA was eluted at 65°C and reverse cross-linked with proteinase K (Fermentas). Immunoprecipitated DNA was treated with RNase A (Fermentas) and purified with the QIAquick purification kit (QIAGEN). ChIP-seq libraries were generated per manufacturer’s instructions (Illumina). Libraries were sequenced on a HiSeq2000 (illumina). Reads were mapped to the TAIR8 genome by allowing up to 2 mismatches, and only retaining uniquely mapping reads. Reads mapping to identical locations were collapsed in to one read and analysis performed was done as previously described (Du et al., 2012).

DNA Methylation Analyses

For genome-wide bisulfite sequencing (BS-Seq), plants were grown under long day condition and samples were collected from 4-week-old plants. Genomic DNA was extracted using Plant DNeasy Mini kit (QIAGEN). BS-seq libraries were generated, sequenced, and analyzed exactly as previously described (Stroud et al., 2013), except that reads were mapped to the TAIR8 genome.

Confocal Microscopy

The localization pattern of H2A.W-RFP in root nuclei and immunostained nuclei were observed with laser scanning confocal microscope (Leica SPE) using 63X magnification objective. The expression pattern of H2A.W-RFP in ovules was observed using a laser scanning confocal microscopes Zeiss LSM5 Exciter and 510. Images were processed using ImageJ (http://rsbweb.nih.gov/ij/). Colocalization analysis was done using auto settings in Imaris (Bitplane) software.

Alignment of Protein Sequences and Phylogeny Analysis

Arabidopsis thaliana histone protein sequences were obtained from TAIR website. Histone H2A variants of Arabidopsis thaliana were aligned using ClustalW2 (www.ebi.ac.uk/Tools/msa/clustalw2/). Phylogenetic tree analysis of histone H2As was done using http://www.phylogeny.fr/version2_cgi/index.cgi (Dereeper et al., 2008). Protein sequences were obtained from http://www.uniprot.org/. Following Uniprot ID sequences were used for phylogenetic tree analysis. Hs_mH2A (O23628), Hs_H2A.X (P16104), Hs_H2A.Z (POC0S5), Mm_H2A.X (P27661), Mm_H2A.Z (POC0S6), Mm_H2A (POC0S8), Mm_mH2A (Q9QZQ8), Dm_H2A.V (P08985), Dm_H2A (P84051), Sc_H2A (P04911), Sc_H2A.Z (Q12692), Pp_H2A.X (A9U2Y0), Pp_H2A.Z (A9S4I9), Pp_H2A.Z (A9TYS8), Zm_H2A.W (P40280), Zm_H2A.X (B6T543), Zm_H2A.Z (B4FX82), Zm_H2A (B4FI6A), At_H2A.X (Q9S9K7), At_H2A (Q9LHQ5), At_H2A.W (Q9FJE8), At_H2A.Z (O23628). Pica abies H2A sequences were obtained from http://congenie.org/.

Protein Expression and Purification

Arabidopsis H2A variants H2A.1, H2A.W.6, and H2A.W.6ΔCT (C-terminal region deleted, also called CT-deletion in this paper) were expressed and purified as previously described (Muthurajian et al., 2011). All histones were prepared by the Protein Expression and Purification Facility. Arabidopsis H2A.1, H2A.W.6, and H2A.W.6ΔCT were refolded together with Xenopus H2B to form various H2A-H2B dimers. Xenopus histone H3 and H4 are reconstituted into tetramer.

SUPPLEMENTAL REFERENCES


Figure S1. Genome-wide Profiling of H2A Variants, Related to Figure 1

(A) Alignment of histone H2A variants of Arabidopsis thaliana generated with ClustalW2. H2A variants are classified into four classes based on their conserved C-terminal tails. Open boxes represent conserved motifs and asterisks represent acidic patch.

(B) Western blot confirming the specific binding of each antibody. The antibodies are tested on whole cell protein extracts. Protein samples were run on two gels (Gel 1: WT, H2A and H2A.X, Gel 2: H2A.W and H2A.Z) in a consistent order for all antibodies. For each antibody Western blot analysis, both gels were treated in an exactly same reagents and conditions. In H2A-RFP transgenic protein samples both respective WT H2A and H2A-RFP fusion proteins are present. H2A.1 and H2A.13 are recognized by the anti-H2A antibody. The anti-H2A.X antibody recognized both members of the family. The anti-H2A.Z antibody is specific for H2A.Z.9. The anti-H2A.W antibody is specific for H2A.W.6. No cross reactivity is observed to RFP fusion proteins of other class members, confirming specificity of the antibodies. For H2A.Z and H2A.W no band was observed in respective T-DNA knocks out lines (*), further confirming the specificity of anti-H2A.Z and anti-H2A.W antibodies.

(C) Distribution of ChIP-seq reads along chromosomes for each H2A variant and H3 relative to input DNA. Grey bars at the bottom indicate pericentromeric heterochromatin.
Figure S2. H2A.W Is Conserved in All Seed Plants, Related to Figure 2

In Arabidopsis, all H2A.W class members are localized to heterochromatin consistently and H2A.W.6 recruitment is independent of DNA methylation and DDM1


(B) Confocal section of a root nucleus of transgenic plants expressing pCENH3::CENH3-GFP and pH2A.W.7::H2A.W.7-RFP.

(C) Confocal section of an integument cell nucleus of transgenic plants expressing pCENH3::CENH3-GFP and pH2A.W.12::H2A.W.12-RFP.

(D) Expression levels of H2A.W.6, H2A.W.7, and H2A.W.12. Reads per kilobase per million reads (RPKM) values were obtained from RNA-seq data of WT.

(E) Confocal section of a root nucleus immunostained for RFP and H3K9me2 with DAPI counterstain.

(F) Confocal section of an ovule integument nucleus immunostained for RFP and H3K9me2 with DAPI counterstain.

(G) WT, met1-3, drm2-2, and ddm1 leaf nuclei immunostained for H3K9me2 and H2A.W.6 with DAPI counterstain. H2A.W.6 is co-localized with H3K9me2 in all the genetic backgrounds.

(H) WT and mutant alleles were analyzed with primers flanking the sites of T-DNA insertion in kyp-6 suvh5 suvh6 (kyp/suuh6) triple mutant (SALK_041474, GABI_263C05, SAIL_1244_F04), cmt3-11 (SALK_148381), drm2-2 (SALK_150883), met1-3, or the EMS-induced SNP in ddm1-2. All mutants lacked transcripts corresponding to the WT. The mutation of the splice donor site in ddm1-2 leads to miss-spliced transcripts (Jeddeloh et al., 1999). cDNA (+RT) or DNaseI-treated RNA (-RT) were used as PCR template. Ubiquitin (Ub10) was used as control for the RT-PCR. PCR products were separated in agarose gels and visualized by ethidium bromide staining.

(I) Quantification of H2A.W.6 and H3K9me2 co-localization in WT, cmt3-11, met1-3, drm2-2, and ddm1 using Imaris software (Bitplane). The graph is plotted using Pearson’s coefficient in ROI volume values on y axis. Error bars represent standard deviation based on a minimum of three independent data sets. In confocal sections, scale bars represent 2 μm.
Figure S3. H3K9me2 Localization Is Not Dependent on H2A.W, Related to Figure 3
(A) Semiquantitative RT-PCR of H2A.W.6, H2A.W.7, and H2A.W.12 in h2a.w.6 h2a.w.7 h2a.w.12 triple mutant and WT. GAPDH was used as an endogenous control.
(B) WT and h2a.w.6 h2a.w.7 h2a.w.12 leaf nucleus immunostained for H3K9me2 and H2A.W.6 with DAPI counterstain. We did not detect H2A.W.6 protein in h2a.w triple mutant chromatin. In confocal sections, the scale bar represents 2 μm.
(C) RT-PCR analysis of WT and kyp-6 (SALK_041474) T-DNA insertion line using primers flanking the sites of T-DNA insertion. Ubiquitin (Ub10) was used as control.
Figure S4. H2A.W and CMT3 Genetically Interact to Repress Heterochromatic Elements, Related to Figure 4

(A) Box plots showing the expression levels of TEs. TEs with 4 fold higher H2A.W.6 ChIP-seq read density compared to H3 and with a non-zero RPKM value for either WT or h2a.w.6 h2a.w.7 h2a.w.12 triple mutant were included in the analysis.

(B) Abortion of pollen in anther of h2a.w.6/+ h2a.w.7/+ cmt3-11/+ triple mutant compared with WT.

(C) Abortion of ovule development in h2a.w.6/+ h2a.w.7/+ cmt3-11/+ triple mutant compared with WT.
Figure S5. H2A.W Is Not Expressed in the Female Gamete Central Cell, Related to Figure 6

(A) Absence of pH2A.W.6::H2A.W.6-RFP expression in the nucleus of the WT central cell (CCN) marked by pFWA::GFP.

(B) The expression of pH2A.W.7::H2A.W.7-RFP is not detected in the central cell nucleus (CCN).

(C) The expression of pH2A.W.12::H2A.W.12-RFP is not detected in the central cell nucleus (CCN). The arrowhead and the dotted circle indicate the position of the central cell nucleus. Scale bars represent 5 μm.
The amino acid sequence of human macroH2A is compared with the three Arabidopsis H2A.W proteins. Note the SPKK like motif in the linker region of the macroH2A and similarities in amino acids in between the extended tail of H2A.W and macroH2A linker region.

**Figure S6. Alignment of Arabidopsis H2A.W Protein with Human macroH2A.1, Related to Discussion**

The amino acid sequence of human macroH2A is compared with the three Arabidopsis H2A.W proteins. Note the SPKK like motif in the linker region of the macroH2A and similarities in amino acids in between the extended tail of H2A.W and macroH2A linker region.