

**Figure 1. Comparison of the protamines of the liverwort *M. polymorpha* to other nonchordate and chordate protamines.** *a*, electrophoresis (acetic acid urea-PAGE) of the SNBPs from the following: lane 1, *M. polymorpha* (26); lane 2, *Oncorhynchus keta* (salmon, salmine (SL)) (84); lanes 3 and 6, *M. californianus* (California mussel) used as a marker (85, 86); lane 4, *Chaetopterus variopedatus* (parchment worm) (23); and lane 5, *Styela montereyensis* (Monterey stalked tunicate) (22). *b*, amino acid sequences of the *O. keta* salmine (SL1–4) protamines (84) and the P2 protamine of *Styela plicata* (sea squirt) (22).

presence of histone H1 ontogenically-related protamines. The results described below conclusively show that in *Marchantia* the post-translational processing of three highly-specialized H1 histones of the PL type, which are expressed in antheridophores, gives rise to the protamines that are found in the sperm of this organism.

## Results

### Initial evidence for the presence of protamines in *Marchantia polymorpha* sperm

The first evidence of protamines in *M. polymorpha* comes from a study published in 1978 by Reynolds and Wolfe (24). Fig. 1D of that paper shows an acetic acid urea-PAGE depicting a pattern of four small molecular mass proteins that is almost identical to that shown in Fig. 1a, lane 1. In both instances, the electrophoretic pattern is characterized by the complete absence of histones, and as it was initially concluded: "... histones are completely lost and replaced by proteins ... resembling animal protamines." (see Ref. 24). Coarse electrophoretic fractionation of these proteins revealed a composition enriched in basic amino acids that was also important in reaching this conclusion. In a subsequent paper, the same authors extended their study to a broader spectrum of plants, including algae, bryophytes, and ferns, for which representative species containing protamines were also identified.

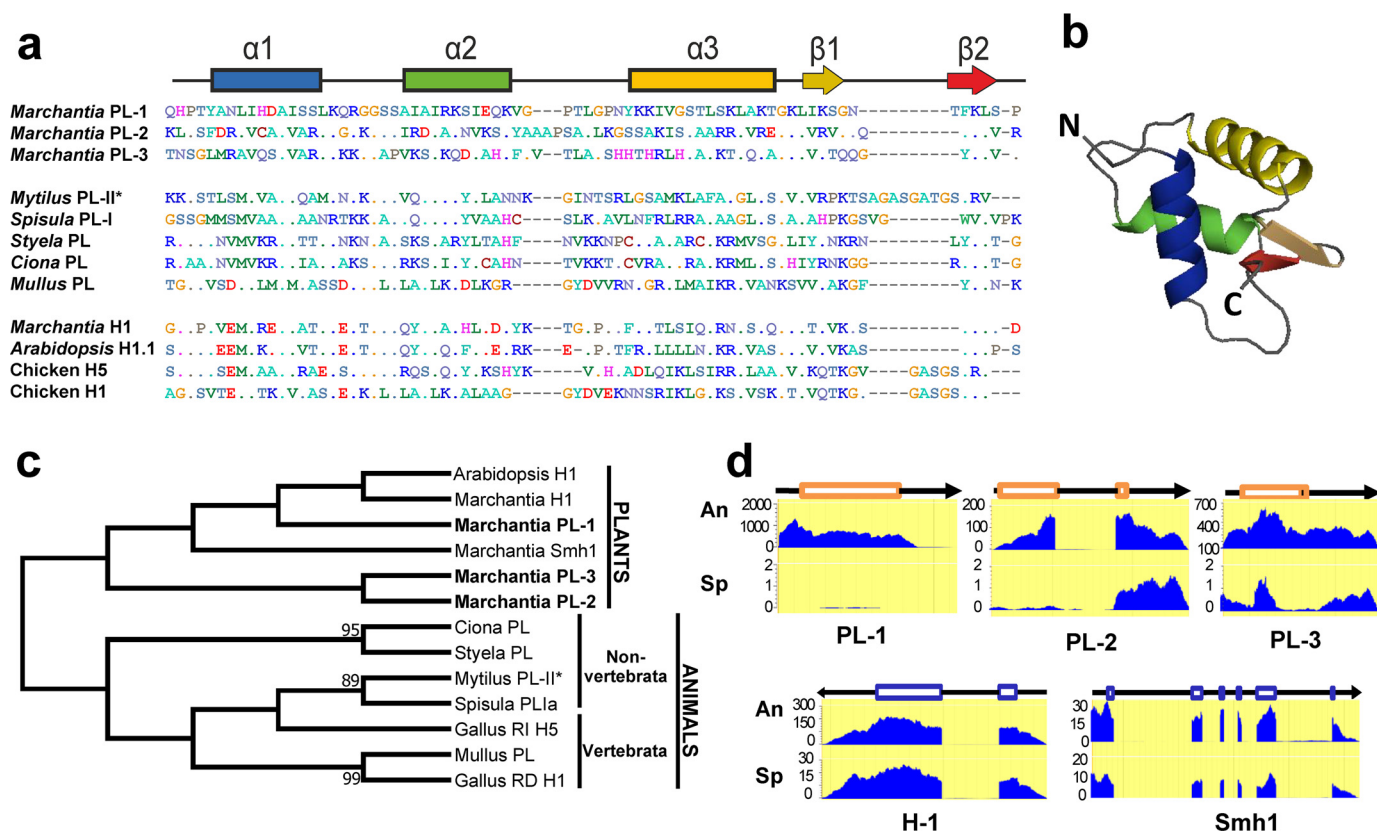
Fig. 1 shows the electrophoretic pattern observed in the sperm of *M. polymorpha* SNBPs compared with those of some

representative nonchordate and chordate animals. The amino acid sequence of the protamines from these organisms is arginine-rich (Fig. 1b), and in the nonchordate organisms a marine worm, *Chaetopterus variopedatus* (lane 4) (23), and a tunicate, *Styela montereyensis* (lane 5) (22), a WHD-containing PL structurally related to chordates exists (P1 in Fig. 1a, lanes 4 and 5). The high electrophoretic mobility of *M. polymorpha* SNBPs resembles that of the small protamines present in salmon sperm (Fig. 1a, lane 2). A very similar arginine-rich composition is observed in protamines of other nonchordate and chordate organisms which, like in salmon, exclusively contain P-type SNBPs in their sperm (4, 7, 25).

### *Marchantia* genome and male transcriptome indicate a relation of its sperm protamines to histone H1

In 2014, we used SNBPs isolated from *M. polymorpha* sperm and Edman degradation N-terminal amino acid sequencing to publish a partial amino acid sequence corresponding to protamine P5 (26) (Fig. 1a, lane 1). This sequence information was more recently used to identify a histone H1 gene of the SNBP PL-type (termed MpPRM) from the analysis of the male gametogenesis transcriptome of this organism which is expressed during spermiogenesis (27). The protein contains the characteristic histone H1 WHD (Fig. 2, a, PL-I, and b) in addition to several arginine-rich clusters at its C-terminal end.

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**Figure 2. Genome of *M. polymorpha* reveals the presence of three histone H1-related PL proteins.** *a*, amino acid sequence alignment of the protein regions corresponding to the WHD of *M. polymorpha* PL1–PL3 compared with the amino acid sequences corresponding to the WHD of nonchordate and chordate organisms as well as the WHD of some representative somatic H1 histones, including the H5 histone, which is in the terminally differentiated erythrocytes of nonmammalian vertebrates: *M. californianus* PL-II\* (86); *S. solidissima* (surf clam) PL-I (87); *S. plicata* PL (22); *C. intestinalis* (transparent sea squirt) PL (22); and *M. surmuletus* (striped ret mullet) PL (88). *b*, predicted tertiary structure organization of the *M. polymorpha* PL-1 WHD. *c*, phylogenetic analysis of the proteins shown in *a*. *d*, levels of gene expression in antheridiophores (An) and in sporophytes (Sp) of *M. polymorpha* PL1–PL3. In the schematic representation, the boxes indicate exon domains. *Smh1*, single *myb* histone 1 (first identified in maize) that binds to telomere DNA repeats in plants (41).

The more recent availability of the *M. polymorpha* genome (28) has allowed us to identify two additional arginine-rich PL proteins also containing a WHD (Fig. 2*a*, PL-2/PL-3). The phylogenetic relationship of these three PL proteins and their evolutionary connection to other *M. polymorpha* histone H1s, as well as to animal PLs, is shown in Fig. 2*c*. The PLs identified in this way are mainly expressed in antheridiophores (the *M. polymorpha* male sex organ), in contrast to the histone H1 counterparts, which are more similarly expressed in both antheridiophores and sporophytes (Fig. 2*d*).

### Determination of the amino acid sequence of *M. polymorpha* SNBPs provides evidence for their PL origin

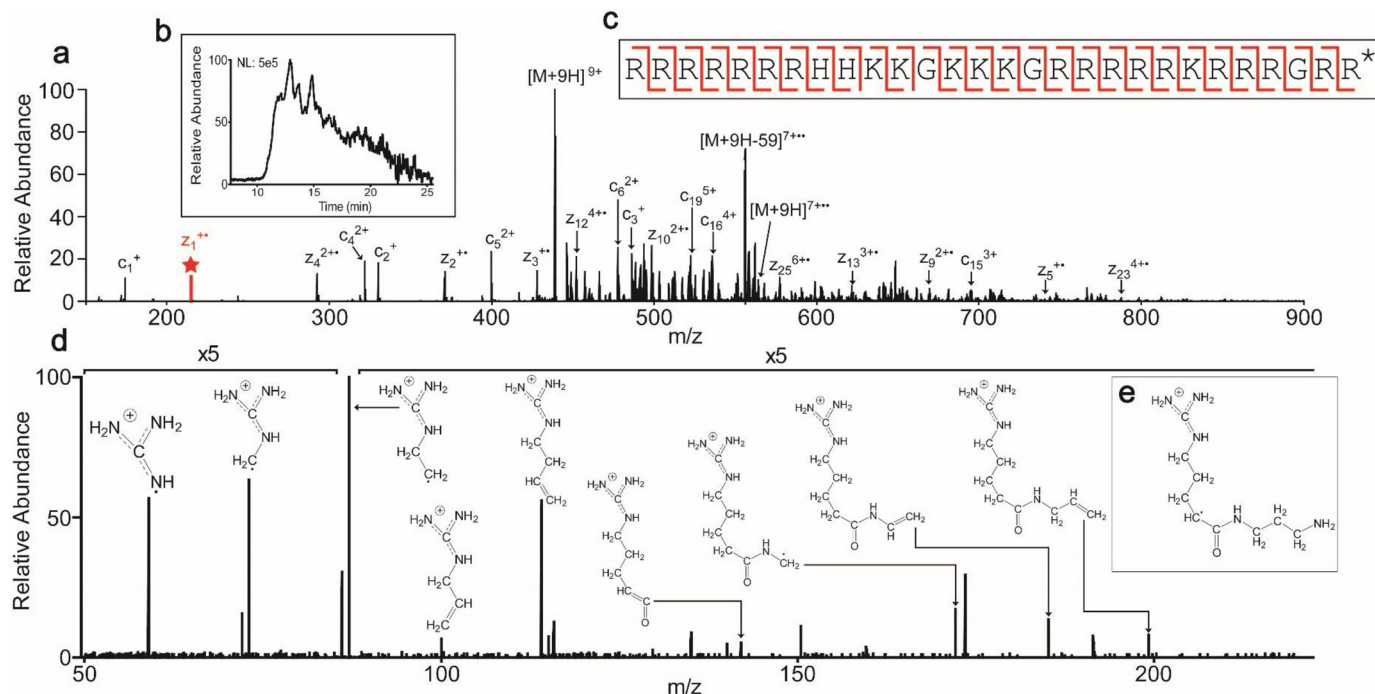
The sequences for the rest of the *M. polymorpha* SNBPs (P1–P4 in Fig. 1*a*) were determined using HPLC-coupled MS. The HPLC-fractionated proteins were fragmented using electron transfer dissociation (ETD) and collisionally-activated dissociation (CAD) (Fig. 3, *a–e*, see also Table 1 and Table S1) (29–31). The sequences obtained (see Table 1), primarily using ETD, confirmed the arginine-rich protamine nature of these proteins, and they revealed their structural relation to the three PL proteins that were bioinformatically identified above. These results conclusively show that *M. polymorpha* SNBPs are the post-translational cleavage

products of three independent, histone H1-related, and protamine-like precursors.

### *M. polymorpha* protamines are di-aminopropanelated at their C termini

Interestingly, in multiple ETD MS<sup>2</sup> scans, a c-ion series could be deduced that covered the majority of the protein. However, there was a strong ion at 215.1740 *m/z* (*z* = 1) present in multiple species denoted by the *star* in Fig. 3*a*. From this ion, a series of amino acids was determined that complemented the c-ion series. Therefore, the ion at 215.1740 *m/z* must be a z' ion. Based on the sequence deduced from the c-ion series, the database BLAST search provided the protein from which the observed species originated as well as the final residue in the protein. This ion corresponded to an arginine with an unknown modification of 56.0740 Da greater than an unmodified arginine. This unknown modification was also observed on C-terminal lysines.

A theoretical elemental composition was calculated to assist in the determination of the modification present on the z<sub>1</sub><sup>+</sup> ion. The molecular formula C<sub>9</sub>H<sub>21</sub>N<sub>5</sub>O was determined to be the closest combination based on parts/million error. Interestingly, there was only one oxygen atom present in the theoretical composition. This would place the modification on the C terminus because an unmodified C terminus would



**Figure 3. Detection and fragmentation of the protamine RRRRRRHHKKGKKGRRRRRKR-RGR\* (where \* = di-aminopropanelated C terminus) (this peptide corresponds to amino acids 102–129 of *Marchantia* PL-1 (MpPRM) NCBI accession number BAU71552).** *a*, precursor ions at 438.07 *m/z* (*z* = 9) were fragmented by ETD to produce the MS<sup>2</sup>. Selected abundant ions are labeled (*M* = precursor mass; 59 = mass of guanidinium group). The peak in red with a star above represents the *z*<sub>1</sub><sup>+</sup> ion. *b*, extracted chromatogram of the protamine. Because the peak elutes for over 10 min, multiple MS<sup>2</sup> scans were averaged to produce the final scan shown in *a*. *c*, sequence coverage by ETD. The cleavages denoted in red depict the *c* and *z'* ions observed giving unambiguous sequence coverage across the length of the protamine. *d*, HCD MS<sup>3</sup> fragmentation spectrum of the *z*<sub>1</sub><sup>+</sup> ion at 215.1740 *m/z* (*z* = 1). Proposed structures for the major ions are depicted. *e*, proposed structure of the *z*<sub>1</sub><sup>+</sup> fragment ion.

**Table 1**  
**Identified protamines**

Asterisk indicates the di-aminopropanylated C terminus. The lower part of the table corresponds to protamines of lower abundance. Amino acids in parentheses do not contain fragment ions confirming their location. These residues can be interchanged within the parentheses.

M + H	Sequence	Protein name	First amino acid residue no.	Last amino acid residue no.	NCBI accession no.
3955.50	RRRRRKSRRRRRRRRRSKGSRSPPRR*	PL-1 (MpPRM)	182	209	BAU71552
3799.41	RRRRRKSRRRRRRRRRSKGSRSPPRR*	PL-1 (MpPRM)	182	208	BAU71552
3942.56	RRRRRRHHKKGKKGRRRRRKR-RGR*	PL-1 (MpPRM)	102	129	BAU71552
3313.16	RRRRRRRRRRRRRRRRRRGRK*	PL-1 (MpPRM)	146	167	BAU71552
7466.75	RRRRRRHGRKGGKPKRHHRRRRRRRRGR KAHRGRKKRG-GRRKGRHHRRRRR*	PL-3	162	216	PTQ32223
4082.51	RSTSRSRGRRRRRRRRRRRRRRRRGRK*	PL-2	207	236	PTQ35141
3780.36	RRRRRKSRRRRRRRRR (SKGSH) SPRR*	PL-1 (MpPRM)	182	208	BAU71552
4139.33	RRRRRKSRRRR (RRTVDRRRRRRRRRR)	PL-1 (MpPRM)	182	209	BAU71552
4366.65	RRRRRKSRRRRRRRRRSKGSRSPPRRHSSR	PL-1 (MpPRM)	182	213	BAU71552
7819.01	RRRRRRHGRKGGKPKRHHRRRRRRRRGR KAHRGRKKRGRRKGRHHRRRRR (GKSH)	PL-3	162	220	PTQ32223

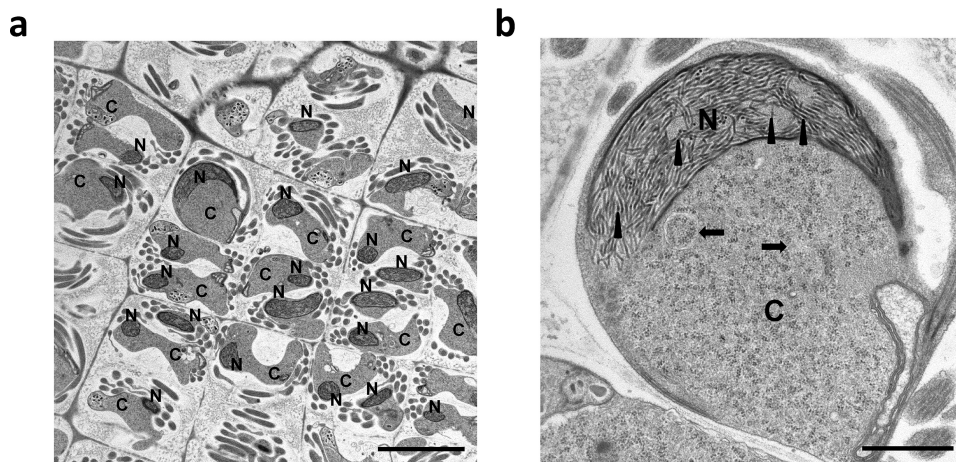
have two oxygens. After accounting for the arginine side chain (C<sub>4</sub>H<sub>11</sub>N<sub>3</sub>) and the protein backbone (C<sub>2</sub>HO), the elements C<sub>3</sub>H<sub>9</sub>N<sub>2</sub> remain.

To determine the structure of the modification, the ion at 215.1740 *m/z* was first formed by ETD and then re-isolated for fragmentation by high-energy collisional dissociation (HCD) to give the MS<sup>3</sup> scan shown in Fig. 3*d*. Because of the guanidino group present on the arginine side chain, the charge should be stabilized within this group. Therefore, the structure can be determined from additions to the arginine side chain. As shown in Fig. 3*d*, the ions present confirm the structure shown in Fig. 3*e*. All identified protamines contained this modification. The modification was absent in other observed proteins from *Marchantia* (results not shown), indicating that this modification is specific to protamines.

### Chromatin organization during *M. polymorpha* spermiogenesis

Organisms where SNBPs of the protamine type undergo a post-translational processing involving extensive protein cleavage often exhibit a characteristic nuclear organization during this process. A transition from lamello-fibrillar chromatin organization to a complete electron-dense highly-compacted organization in their mature sperm is observed (7). Such a transition has been observed both in internally fertilizing invertebrate (26, 32) and vertebrate organisms (33–35).

Fig. 4 shows an electron microscopic image of spermatids in developing *M. polymorpha* antheridia (36) (the haploid structure producing male gametes). These spermatids have been shown to contain PL-1-type protamines until the late stages of spermiogenesis (27). As can be seen in Fig. 4, chromatin adopts



**Figure 4. Electron micrographs of *M. polymorpha* spermatids.** *a*, chromatin condensing nucleus (N) exhibiting a shrinking cytoplasm (C). The scale bar is 5  $\mu\text{m}$ . *b*, electron micrograph (greater magnification) of a condensing spermatid showing the characteristic  $24 \pm 3\text{-nm}$  fibers (arrowheads). The arrows point to autophagosomes. The scale bar is 1  $\mu\text{m}$ .

a distinct fibrillar organization with dispersed uniform fibers of  $24 \pm 3\text{-nm}$  thickness. The fibers appear clearly dispersed and further coalesce into a uniform, highly-condensed, and electron-dense opaque nucleus in the mature sperm (37). This type of chromatin organization has also been observed in the liverwort *Blasia pusilla*, a bryophyte (26). The peculiar transitional chromatin organization before complete condensation observed in these organisms has been proposed to be mediated by the liquid-liquid phase separation mechanism of spinodal decomposition (26). The process is framed on a physicochemical model that involves “kinetic, equilibrium, and structural aspects of a system in route to equilibrium.” In it, the less electron-dense nucleoplasm and the chromatin appear to be continuous rather than one disperse and one continuous phase (7).

## Discussion

### Protamines in plants

The occurrence of protamines in the sperm of *Marchantia* may have broader implications for the overall evolution of SNBPs. As pointed out in earlier work, the shift to the convergent presence of SNBPs in the sperm of both plants and animals might have involved selection pressure to reduce sperm nuclear weight and volume for a swimming male gamete (24) in internally fertilizing organisms. Indeed, protamines appear to be present in more primitive lower plants with motile sperm and are much less ubiquitous in the male nuclei of the pollen grain (3). In this regard, it is interesting that while in animals protamines may be present in both internal and external fertilizers, the last type of fertilization represents a SNBP evolutionary bottleneck. Once organisms from a clade have acquired this fertilization mode, there is no return to the histone or PL SNBP types (38, 39). Moreover, evolutionarily-driven interspecific changes in the regulation of protamine genes and amino acid sequence in mice and mammals have been shown to confer inter-specific sperm competition (fertility) advantage (40, 42, 43). Thus, in addition to providing genome compaction/protection against DNA damage and somatic histone epigenetic clearance (6, 11), protamines might have evolved to fine-tune the structural chromatin features of the nucleus (44). These

changes optimize the swimming potential of internal fertilizers, not only in the animal kingdom but also in plant internal fertilizers, such as in the case of bryophytes, including liverworts, mosses, and hornworts (45).

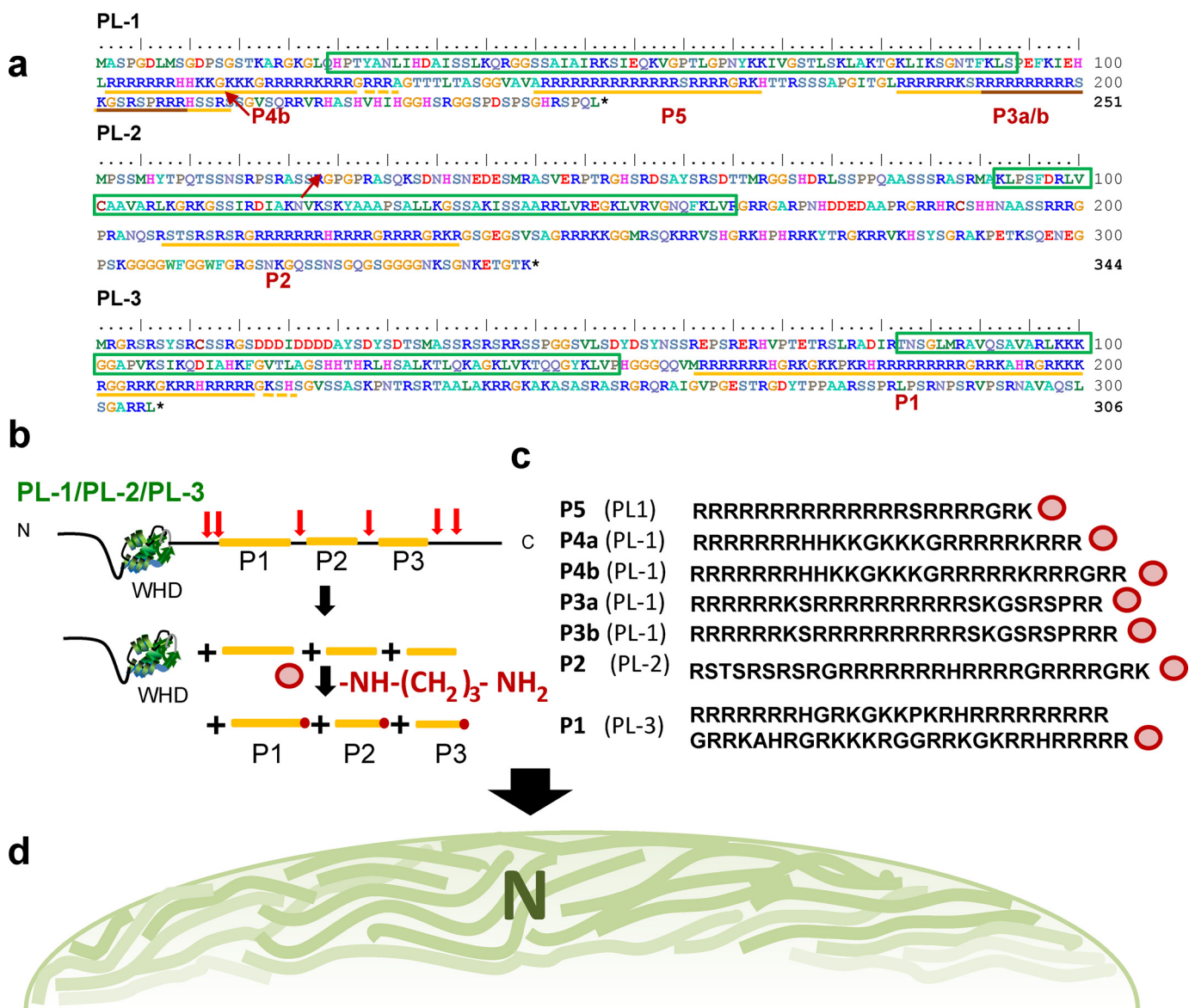
### Protamines and histone H1

Fig. 5 summarizes the relationship between the SNBPs of *M. polymorpha* (Fig. 1a, lane 1) and their histone H1-related PL precursors. Once more, these results emphasize the evolutionary relatedness of protamines to histone H1, not only in animals (4, 16) but also in plants (Fig. 2c). Although protamines are the only SNBPs found in *M. polymorpha* mature sperm, their PL histone H1-related precursors are present in the spermatids in the early stages of spermiogenesis (27). Moreover, *M. polymorpha* protamines present in the final sperm are the product of post-translational cleavage of their precursors (Fig. 5b).

Post-translational cleavage of protamine precursors is quite a general occurrence both in protostome and deuterostome species, which appears to be independent of their histone H1 origin, and it also appears to have been the product of evolutionary convergence (46). For instance, the protamines of cephalopods (*Sepia officinalis* (cuttlefish) (47)/*Loligo opalescens* (squid) (46)) and the protamine P2 in mammals (mouse (48) and human (49)) are the products of a gene encoding a protein with an N-terminal leading domain consisting of a mixture of neutral/polar amino acids and a highly-arginine-rich C-terminal end. The N-terminal leading sequence is gradually removed during the late stages of spermiogenesis following DNA binding (6, 48), and hence it is hypothesized to play an important role in proper chromatin deposition and condensation.

### Protamine post-translational processing and sperm chromatin condensation

The post-translational modifications (PTMs) undergone by *M. polymorpha* protamine precursors include cleavage as well as C-terminal di-aminopropanelation (Fig. 5, b and c). Di-aminopropane is a product of the oxidation of spermine and spermidine that is catalyzed by the FAD-dependent polyamine



**Figure 5. Protamines of *M. polymorpha* sperm are the post-translational cleavage products from three PL histone H1 precursors PL-1, PL-2, and PL-3.** *a*, amino acid sequences of the three PL precursors. The boxes with green borders indicate the WHD, which is characteristic of H1 histones. The regions underlined in yellow correspond to the peptides identified in mature sperm. The region of PL-1 underlined in brown represents the first *M. polymorpha* protamine sequence determined using conventional Edman degradation sequencing (26). *b*, schematic representation of the post-translational cleavage undergone by *M. polymorpha* PLs. Precursor SNBP histone H1-related PLs consisting of an N-terminal (N) winged helix domain (WHD) and an arginine-rich C-terminal domain (C) are post-translationally cleaved (red arrows) into smaller fragments (yellow) and further C-terminally di-aminopropanelated (dark red dots) of the protamines (Fig. 1a, lane 1) identified by MS. *d*, schematic representation of the intermediate liquid phase chromatin condensation stage observed in spermatids (Fig. 4).

oxidases commonly found in monocotyledonous plants (50, 51) and that in *A. thaliana* have been shown to participate in polyamine metabolism with important involvement in plant development (50). Upon activation of the arginine  $\alpha$ -carboxyl group in the *M. polymorpha* protamine, it can readily acylate 1,3-diaminopropane to produce the PTM observed here. Of note, oxidation of spermine by diamine oxidase occurs in human seminal plasma and is related to sperm fertility (52, 53).

The occurrence of PTMs in mammalian protamines has been well-documented (6), where phosphorylation, acetylation, and methylation of different residues have been described (54). Although the functional role of these PTMs is not well-understood, serine phosphorylation was initially proposed to regulate the proper interaction of the highly positively-charged prota-

mines with DNA, therefore ensuring a proper chromatin assembly (55–57). In those instances where cleavage of a protamine precursor takes place, like in mammalian protamine P2, phosphorylation does not occur until the initiation of the cleavage process (56).

There is not much information about C-terminal PTMs due mainly to the experimental difficulty of their analysis (58), and hence there is not much information about their functional involvement. However, half of the biologically active peptides and peptide hormones are  $\alpha$ -amidated at their C termini where the neutralization of the negative charge enhances their hydrophobicity and improves their receptor-binding activity (59). In this regard, the role of the carboxyl end di-aminopropanelation, observed for the first time here in chromosomal proteins,

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remains elusive. The possibility exists that as in the case of phosphorylation, it has an important role in the chromatin transitions undergone during spermiogenesis. In particular, it might play a role during the PL to protamine transition as it might neutralize the negative charge of the carboxyl end which, like in the hormone peptides (59), might otherwise interfere with proper protamine deposition onto DNA from a complex heterogeneous mixture of very small arginine-rich proteins (Figs. 1a, lane 1, and 5c).

Protamine precursor trimming and phosphorylation have been involved in chromatin patterning of the developing spermatid nucleus of those organisms undergoing liquid-phase condensation (60) through a process of spinodal decomposition (7, 32, 61). These events precede the final highly-compacted chromatin structure that is characteristic of mature sperm. The protein processing observed in *M. polymorpha* appears to play a similar role as its spermatids exhibit this pattern type (Figs. 4 and 5d), in which the intrinsically disordered nature of the proteins (8) involved (histone H1 and protamines) play a critical role in this process (10, 62–64). This is very similar to the phase separation process recently described for the heterochromatin domain formation by *Drosophila* HP1a (65) and other chromatin subcompartments (66). Indeed, sperm chromatin is a salient example of fully heterochromatinized nuclei.

Paraphrasing the title of the paper reporting the genome of *M. polymorpha* (28), the analysis of its SNBP composition and transitions, made possible through the analysis of its genome, provides a powerful evolutionary insight that transcends that of plants and has implications for the evolution of these protamines and their conserved chromatin organization encompassing the plant and animal kingdoms.

## Experimental procedures

### Materials

Male accession of *M. polymorpha*, Takaragaike-1 (Tak-1) (67), was grown and induced to sexual reproduction as described in Minamino *et al.* (36).

LCMS-grade water and LCMS-grade acetonitrile were purchased from VWR Scientific. LCMS-grade 2-propanol and formic acid were purchased from Thermo Fisher Scientific. Acetic acid, vasoactive intestinal peptide, and angiotensin I were purchased from Sigma.

### Protein extraction

*M. polymorpha* sperm from antheridiophores was resuspended in 100 mM Tris-HCl (pH 7.5), 0.5% Triton X-100 containing 10 µg/ml tosyl lysine chloromethyl ketone buffer (at a ratio of 20 µl/liter antheridiophore) and centrifuged at maximum speed in an Eppendorf microcentrifuge. The pellet obtained in this way was homogenized in 0.4 N HCl (at a ratio of 10 µl/1 antheridiophore) and homogenized in a small Dounce with 10 strokes. The HCl extract was precipitated overnight with 6 volumes of acetone at –20 °C, and the next day it was centrifuged at maximum speed in an Eppendorf microcentrifuge. The protein pellet was washed with acetone at room temperature, centrifuged again under the same conditions, and

dried in a speedvac. The protein pellet thus obtained was resuspended in water for further use.

### Gel electrophoresis

Gel electrophoresis was carried out on acetic acid (5%), urea (2.5 M), and (15%) PAGE-polymerized using thiourea and hydrogen peroxide (68), as described previously (69).

### Genome screening

*M. polymorpha* proteins containing a histone H1 globular domain were retrieved from the *M. polymorpha* Genome Database (<http://marchantia.info/><sup>4</sup> (28)). Gene structure and levels of expression were analyzed using the Genome Browser implemented in this database. PL proteins from the mussel *M. californianus* (PL-II\*; accession no. AAB24707), the surf clam *Spisula solidissima* (PLIa; accession no. AAT45384), the tunicates *S. montereyensis* (P1; accession no. AAQ01227) and *Ciona intestinalis* (PL; accession no. XP\_002130983), the fish *Mullus surmuletus* (PL; accession no. Q08GK9), histone H1.1 from *Arabidopsis thaliana* (accession no. AAF63139), and histones H1 and H5 from the chicken *Gallus gallus* (accession nos. P09987 and AAA48798, respectively) were retrieved from the GenBank™ database.

### Multiple sequence alignments

These were performed using MAFFT version 7.310 (70) in the Jalview2 package (71). The evolutionary history of the winged-helix globular domains (WHD) was inferred by using the maximum likelihood method based on the Le Gascuel model (72) in the software MEGA version 7 (73). The bootstrap consensus tree inferred from 1000 replicates is taken to represent the evolutionary history of the taxa analyzed. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) is shown next to the branches (only bootstrap values higher than 80% are shown). A discrete Gamma distribution was used to model evolutionary rate differences among sites (five categories (+G, parameter = 2.3546)). The analysis involved 13 amino acid sequences. All positions with less than 95% site coverage were eliminated. That means fewer than 5% alignment gaps, missing data, and ambiguous bases were allowed at any position. There were a total of 65 positions in the final dataset.

### Tertiary structure prediction

The tertiary structures of the globular domains of histone H1 and PL proteins were modeled in the SWISS-MODEL server (74) using the most closely-related structure, automatically searched by the software, as a template in each case. The obtained structures were rendered using the PyMOL Molecular Graphics System version 1.2r1.

### Sample preparation

One µl of the acetone-precipitated protein-HCl extract suspension was diluted 10-fold with 0.1% acetic acid in water. 300 nl of the dilution, corresponding to ~0.2% of the total suspen-

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sion, was pressure-loaded onto a reversed-phase column containing 10 cm of 3- $\mu$ m diameter, 300 Å PLRP-s packing material (Polymer Laboratories) within a 360  $\times$  75- $\mu$ m fused silica capillary integrated with an electrospray tip (75, 76). Additionally, 100 fmol of both vasoactive intestinal peptide and angiotensin I were loaded onto the column as internal standards.

### Chromatography and mass spectrometry

An Agilent Technologies (Palo Alto, CA) 1100 Series binary HPLC system coupled to a Thermo Fisher Scientific Orbitrap Fusion Tribrid mass spectrometer (San Jose, CA) operating in standard pressure mode was used to characterize the proteins in the sample (77). Proteins that were unable to be retained on the column due to high hydrophilicity were identified using an isocratic elution at 100% solvent A (0.3% formic acid in water) for 25 min at a flow rate of  $\sim$ 50 nl/min (Fig. S1). Following the identification of proteins not retained on the column, the column was washed with solvent A for an additional 25 min at  $\sim$ 100 nl/min to remove any additional salts. The remaining proteins were then eluted using a gradient of 0–10<sup>−30</sup>–70–100% solvent B (72% acetonitrile, 18% isopropenyl acetate, 10% water, and 0.3% formic acid) in 0–10<sup>−20</sup>–30–35 min at a flow rate of  $\sim$ 100 nl/min (Fig. S2).

Proteins were selected for fragmentation based on a 60,000 resolution Orbitrap MS<sup>1</sup> scan. Using top speed mode with a 3-s cycle time, proteins with a charge state of 4 and higher were isolated by the quadrupole with a width of 1.5 *m/z* and fragmented by ETD and CAD (29–31). All MS<sup>2</sup> scans were analyzed in the Orbitrap at 60,000 resolution with an AGC target of 2.0e5 and a maximum injection time of 100 ms. ETD was performed on ions with *m/z* 300–700 using calibrated reaction times (78, 79) (Table S2). CAD was performed on ions with *m/z* 400–1500 at 30% normalized collision energy. Ions were placed on an exclusion list for 30 s after three scans in 30 s. For only the isocratic identification, all scans included an in-source dissociation energy of 40 to break up any adducts that may be present due to the acetone precipitation.

### Determination of C-terminal di-aminopropanelation by MS3

A targeted MS3 method was used to determine the structure of the z<sub>1</sub><sup>+</sup> ion present in many species because it did not correspond to a known amino acid (80, 81). First, the precursor mass of 439.18 *m/z* (*z* = 9) was isolated with the quadrupole at a width of 1.5 *m/z*, an AGC target of 5.0e5, and a maximum injection time of 300 ms. These ions were then fragmented by ETD using 2.0e5 reagent ions and a reaction time of 12 ms. The fragment ion at 215.174 *m/z* (*z* = 1) was then isolated with a width of 1.5 *m/z* and fragmented by HCD using 30% HCD collisional energy. The resulting fragment ions were analyzed in the Orbitrap at 60,000 resolution and a scan range of 50–220 *m/z*.

### MS data analysis

The data files were manually inspected for the species present. All fragmentation scans from the major species were manually sequenced *de novo* by averaging all MS<sup>2</sup> scans under the peak using Qual Browser version 4.0.27.10 (Thermo Fisher Scientific). Sequenced proteins were identified by using the NCBI

Protein Blast Nonredundant protein sequences database (as of 03/22/2019) against *M. polymorpha* (82, 83).

### EM

EM was carried out as described previously (36). Briefly, *M. polymorpha* antheridia were fixed overnight with paraformaldehyde (2%) and glutaraldehyde (2%) in 0.05 M cacodylate buffer (pH 7.4) at 4 °C. The samples were next washed three times with cacodylate buffer and post-fixed with osmium tetroxide (2%) in the same buffer for 3 h at 4 °C. The samples were dehydrated by rinsing them for 30 min with increasing concentrations (50, 70, and 90%) of ethanol at 4 °C, and this was repeated four times with 100% ethanol at room temperature. The samples were finally left overnight at room temperature in 100% ethanol. The samples were subsequently embedded twice in propylene oxide (PO) for 30 min and then transferred to a 70:30 mixture of PO and Quetol-651 resin (Nisshin EM Co.) for 1 h. After evaporation of PO, the samples were transferred to 100% resin, and polymerization was allowed to proceed for 48 h at 60 °C. Ultra-thin sections were stained with 2% uranyl acetate and lead stain solution (Sigma) visualized on a JEM1400 Plus (Jeol Ltd.) transmission electron microscope.

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