PRDM9 losses in vertebrates are coupled to those of paralogs 2 ZCWPW1 and ZCWPW2

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18 Abstract

In most mammals and likely throughout vertebrates, the gene PRDM9 specifies the 19 20 locations of meiotic double strand breaks; in mice and humans at least, it also aids in their repair. 21 For both roles, many of the molecular partners remain unknown. Here, we take a phylogenetic approach to identify genes that may be interacting with PRDM9, by leveraging the fact that 22 23 PRDM9 arose before the origin of vertebrates, but was lost many times, either partially or entirely-24 -and with it, its role in recombination. As a first step, we characterize PRDM9 domain composition 25 across 446 vertebrate species, inferring at least thirteen independent losses. We then use the 26 interdigitation of *PRDM9* orthologs across vertebrates to test whether it co-evolved with any of 27 241 candidate genes co-expressed with PRDM9 in mice or associated with recombination 28 phenotypes in mammals. Accounting for the phylogenetic relationship among species, we find 29 two genes whose presence and absence is unexpectedly coincident with that of PRDM9: 30 ZCWPW1, which was recently shown to facilitate double strand break repair, and its paralog ZCWPW2, as well as more tentative evidence for TEX15 and FBXO47. ZCWPW2 is expected to 31 32 be recruited to sites of PRDM9 binding; its tight coevolution with PRDM9 across vertebrates suggests that it is a key interactor, with a role either in recruiting the recombination machinery or 33 34 in double strand break repair.

35 Author Summary

36 Our understanding of meiotic recombination in mammals has seen great progress over 37 the past 15 years, spurred in part by the convergence of lines of evidence from molecular biology, 38 statistical genetics and evolutionary biology. We now know that in most mammals and likely in 39 many vertebrates, the gene PRDM9 specifies the location of meiotic double strand breaks and 40 that in mice and humans at least, it also aids in their repair. For both roles, however, many of the 41 molecular partners remain unknown. To search for these, we take a phylogenetic approach, 42 leveraging the fact that the complete PRDM9 has been lost at least thirteen times in vertebrates 43 and thus that its presence is interdigitated across species. By this approach, we identify two genes 44 whose presence or absence across vertebrates is coupled to the presence or absence of *PRDM9*, 45 ZCWPW1 and ZCWPW2, as well as two genes for which the evidence is weaker, TEX15 and 46 FBX047. ZCWPW1 was recently shown to be recruited to sites of PRDM9 binding and to aid in 47 the repair of double strand breaks. ZCWPW2 is likely recruited to sites of PRDM9 binding as well: 48 its tight coevolution with *PRDM9* across vertebrates suggests that it plays an important role either 49 in double strand break formation, potentially as the missing link that recruits the recombination 50 machinery to sites of PRDM9 binding, or in double strand break repair.

51 Introduction

52 Meiotic recombination is initiated by the deliberate infliction of numerous double strand 53 breaks (DSBs) in the genome, the repair of which yields crossover and non-crossover resolutions 54 (reviewed in [1]). In mice and humans, and probably in most mammals, the localization of almost 55 all DSBs is specified through the binding of PRDM9 [2–4]. Yet the presence of a PRDM9 binding 56 site is far from sufficient for a DSB to be made: a number of additional factors modulate whether 57 PRDM9 binds or act downstream of PRDM9 binding [5–7].

58 The mechanism by which PRDM9 directs recombination to the genome is partially 59 understood: it binds DNA through a C2H2 zinc finger (ZF) array and contains a SET domain that 60 tri-methylates histones H3K4 and H3K36 [8,9]. These epigenetic marks together recruit the DSB 61 machinery, notably SPO11 (which makes the DSBs), through intermediates that remain unknown 62 [10]. In addition to the zinc finger binding array and SET domain, most mammalian *PRDM9* genes 63 also have two other domains, KRAB and SSXRD, whose functions are unclear.

The complete PRDM9 protein, with all four domains, originated before the diversification of vertebrates, so has been conserved for hundreds of millions of years [11,12]. Yet the entire gene has also been lost numerous times, including in birds and canids [13–15]. In these species,

67 recombination occurs preferentially around promoter-like features, notably CpG islands [11,15-68 17]. A possible explanation is that in the absence of the histone marks laid down by PRDM9, the 69 recombination machinery defaults to those residual H3K4me3 marks found in the genome, often 70 associated with sites of transcription initiation, or perhaps simply to wherever DNA is accessible 71 [15,18]. The same concentration of DSBs around promoter-like features is seen in $Prdm9^{-1}$ mice 72 [18] and in a woman who carries two loss of function copies of *PRDM9* identical by descent [19]. 73 These findings suggest that mammals that carry an intact PRDM9 retain the mechanism to direct 74 recombination employed by species lacking *PRDM9*, but it is normally outcompeted by PRDM9 75 binding.

76 In addition to complete losses of PRDM9, multiple partial losses have occurred 77 independently (e.g., in platypus and various fish lineages), usually involving the truncation of the 78 N-terminal KRAB and SSXRD domains [11]. Although these partial PRDM9 orthologs evolve 79 under selective constraint and thus must have some conserved function [11], several lines of 80 evidence indicate that they do not direct recombination. For one, only in species with a complete 81 PRDM9 is the ZF unusually rapidly evolving in its binding affinity [11]. Since the rapid evolution of 82 the ZF is thought to arise from the role of PRDM9 in recombination [3,20,21], this evolutionary 83 pattern suggests that all four domains are required for DSB localization. Empirical data support 84 this notion: in swordtail fish carrying one PRDM9 ortholog that lacks KRAB and SSXRD domains 85 as well as in a mouse model in which only the KRAB domain is knocked out, recombination events 86 are concentrated at promoter-like features, as in species lacking PRDM9 altogether [11,22]. 87 Therefore, the KRAB domain at least appears to be necessary for PRDM9 to direct recombination, 88 likely by mediating interactions with other proteins [22,23].

Conversely, the presence of a complete *PRDM9* with a rapidly evolving ZF outside of mammals [11] suggests that PRDM9 also directs recombination to the genome in these species, as has been reported for rattlesnakes [24]. Thus, at least two mechanisms for directing meiotic recombination are interdigitated within mammals as well as seemingly throughout the vertebrate phylogeny.

In addition to specifying the locations of DSBs, PRDM9 has recently been discovered to play a second role, in the downstream repair of DSBs [25–27]. In mice and humans, DSBs at which PRDM9 is bound on both homologs are more likely to be efficiently repaired and to result in a crossover; in contrast, DSBs at which PRDM9 is only bound on one of the two homologs are delayed in their repair [27,28]. If these "asymmetric" DSBs are overwhelming in number—as is the case in certain hybrid crosses in mice—this delay can lead to asynapsis and infertility [29,30].

While this second role of PRDM9 is still poorly understood, recent papers report that it is facilitated by ZCWPW1, which binds H3K4me3 and H3K36me3 [25–27] and is expressed alongside PRDM9 in single cell data from mouse testes [31]. One line of evidence that enabled the discovery of *ZCWPW1* is that although it too has been lost numerous times in vertebrates, it is found in seven clades that carry an intact *PRDM9* [26,27].

105 The important hint provided by the phylogenetic distribution of ZCWPW1 points to the 106 potential power of co-evolutionary tests to identify additional molecular partners of PRDM9. Here, 107 we took this approach more systematically: we considered a set of 241 candidate genes that are 108 either known to be involved in recombination in model organisms [32], associated with 109 recombination phenotypes in a human genome-wide association study [33], or co-expressed with 110 PRDM9 in single cell data from mouse testes [31] and tested for their co-occurrence with PRDM9 111 across 189 vertebrate species. After verifying our initial gene status calls in whole genome data 112 and, for a subset of species, RNA-seq data, we identified the paralog of ZCWPW1, ZCWPW2, as 113 co-evolving with PRDM9 and found more tentative evidence for two additional genes, TEX15 and 114 FBXO47.

115 **Results**

116 A revised phylogeny of PRDM9

117 We previously reported that the complete *PRDM9* gene, including KRAB, SSXRD and 118 SET domains, arose before the origin of vertebrates and was lost independently a number of 119 times, both in its entirety and partially (through the loss of its N-terminal domains; [11]). Here, we 120 leverage the independent losses of *PRDM9* in order to identify genes that are co-evolving with 121 *PRDM9*—specifically, that tend to be present in the same species as *PRDM9* and lost (partially 122 or entirely) when *PRDM9* is no longer complete.

123 As a first step, we characterized the phylogenetic distribution of *PRDM9* in light of new 124 genome sequences published since our initial analysis [11]. To this end, we created a curated 125 dataset of 747 vertebrate *PRDM9* sequences by analyzing publicly available protein sequences 126 from Refseq [34], whole genome sequences, and RNA-seq data from testes samples, as well as 127 four RNA-seq datasets from testes samples that we generated (see Methods, Figure S1, Tables 128 S1-3). For this analysis, we defined *PRDM9* orthologs as complete if they contain both KRAB and 129 SET domains; we did not consider the SSXRD domain, because its short length makes its 130 detection at a given e-value threshold unreliable, or the ZF array, because its repetitive structure 131 makes it difficult to sequence and assemble reliably.

132 Across 446 species, we identified 221 species with at least one complete PRDM9 ortholog 133 and 225 species without a complete *PRDM9* ortholog (Figure 1, Table S4). Notably, we were 134 able to uncover complete *PRDM9* orthologs in a number of species for which we had previously 135 predicted partial or complete losses [11], including in the Tasmanian devil (Sarcophilus harrisii), 136 the atlantic cod (Gadus morhua), and the atlantic herring (Clupea harengus), as well as in a 137 handful of placental mammals that we had previously only investigated using RefSeq (see Table 138 **S4** for details). We also found a complete *PRDM9* ortholog in caecilians and in two species of 139 frogs, suggesting that the previously reported loss of *PRDM9* in amphibians [11] reflects at least 140 one loss in salamanders and more than one independent loss in frogs. We note, finally, that by 141 the approach taken here, the PRDM9 ortholog from the Australian ghostshark (Callorhinchus milii) 142 is considered to be complete (in contrast to in Baker et al. 2017, where we also relied on the 143 SSXRD domain; see Table S4 for details).

144 Given the phylogenetic relationships among species given by the TimeTree tool 145 (http://timetree.org/; [35]), we inferred 23 putative complete or partial losses of PRDM9 across the 146 446 vertebrates considered (Figure 1, Table S4). These putative losses include six previously 147 reported ones [11,13–15], each observed in two or more closely related species: in percomorph 148 fish, cypriniformes fish, characiformes and siluriformes fish, osteoglossomorpha fish, birds and 149 crocodiles, and canids. In addition, independent work supports our finding of a partial loss of 150 PRDM9 in the platypus (Ornithorhynchus anatinus) (J. Hussin and P. Donnelly, personal 151 communication). In turn, the putative losses of PRDM9 in polypteriformes fish, salamanders, and 152 in three clades of frog species (Xenopus, Dicroglossidae and Bufonidae) were each supported by 153 the absence of PRDM9 in the genomes of two or more closely related species. We were further 154 able to verify the absence of PRDM9 in two Xenopus frogs and in two salamanders using RNA-155 seq data from testes: despite sufficient power to detect a set of six highly conserved meiotic genes 156 in each species, we did not detect the expression of any complete PRDM9 orthologs (Table S3).

157 We also failed to find *PRDM9* in RefSeq or the whole genome sequence of the green 158 anole (Anolis carolinensis). We verified this absence of PRDM9 by collecting RNA-seq data from 159 testes in the green anole as well as in the fence lizard (Sceloporus undulatus), for which neither 160 a Refseq nor a genome sequence were available at the time. Despite sufficient power to detect a 161 set of six highly conserved meiotic genes, we were unable to detect PRDM9 expression in either 162 species (Figure S2-3, Table S3). Given the presence of a complete *PRDM9* in bearded dragons 163 (Pogona vitticeps), it appears that this loss of PRDM9 occurred in a lineage basal to the common 164 ancestor of green anoles and fence lizards, over 99 Mya but less than 157 Mya (Figure S5).

165 The remaining 10 putative absences of *PRDM9* are observed in single species; we were 166 unable to verify the calls using testis RNA-seq data, so their *PRDM9* status remains uncertain. 167 Thus, in total, we identified at least 13 independent *PRDM9* losses in vertebrates, and possibly 168 as many as 23 (**Figure 1**, **Table S4**). The 13 losses are all relatively old (**Figure S4**): the most 169 recent case manifest in these data is either the one that happened in the branch leading to 170 platypus or the one in canids, which could be as recent as 14.2 Mya (**Figure S4**).

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172 Identifying genes co-evolving with PRDM9

We selected 193 candidate genes based on their co-expression with PRDM9 in single cell RNA-seq data from mouse testes (specifically, in component 5; see Methods [31]) (**Figure S6A-B**). To this set, we added any gene associated with variation in recombination phenotypes in humans [33] as well as genes known to have a role in mammalian meiotic recombination from functional studies (summarized in [32]). Together, these three sources provided a total of 241 genes to evaluate for possible co-evolution with *PRDM9* (**Table S5**, **Figure S6C**).

179 We evaluated the presence or absence of these 241 genes across the NCBI RefSeg 180 database of 189 species. These 189 species were downsampled from the larger phylogenetic 181 tree to preserve at most three species with high quality genomes below each PRDM9 loss, 182 thereby minimizing phylogenetic signals of genome quality. The phylogeny includes 183 representative species for 11 of the 13 inferred PRDM9 losses (see Methods, Figure S7). Species 184 of Bufonidae frogs and salamanders were not included due to the absence of available gene 185 annotations; moreover, due to the lack of gene annotations for frog species with PRDM9, within 186 these 189 species, the losses in Xenopus and Dicroglossidae frogs cannot be distinguished from 187 a single event.

188 We encoded a gene as present when it contained all the domains found in four 189 representative vertebrates with a complete *PRDM9* and absent if it lacked one or more of those 190 domains (see Methods). Many of the 241 genes are present in every sampled vertebrate and 191 hence provide no information in our co-evolutionary test of presence and absence: specifically, 192 we found apparently complete orthologs for 102 candidate genes in all 189 species used in the 193 phylogenetic test. We therefore focused on the remaining 139 genes, each of which has been 194 lost at least once among vertebrate species evaluated here; the matrix of 189x139 gene status 195 calls is presented in Table S6.

We tested for the co-evolution of *PRDM9* and each candidate gene by comparing a null model with independent rates of gains and losses of *PRDM9* and of the focal gene to an alternative model in which the state transition rates of the two genes are dependent on one

another, using the maximum likelihood approach within *BayestraitsV3* [36,37] (Table S7, Figure
S8). By this approach, we identified nine significant hits at the 5% level (uncorrected for multiple
tests): in order of increasing p-values, *ZCWPW1*, *MEI1*, *ZCWPW2*, *TEX15*, *FBXO47*, *ANKRD31*, *NFKBIL1*, *SYCE1*, and *FMR1NB*. We focused on the top five, for which the false discovery (FDR)
value is below 50% (Table 1, Figure 2A).

204 We sought to verify the phylogenetic distribution of the top genes by developing curated 205 datasets of high confidence orthologs, as we had for PRDM9 (see Methods; Figure 3, Tables S1 206 and **S8**, Figure S8). In doing so, we were able to identify *MEI1* orthologs from the whole genome 207 assemblies of each species missing MEI1 in our initial dataset, resulting in the presence of MEI1 208 in every species considered (**Table S1**); thus, it appears that its inferred co-evolution with *PRDM9* 209 based on Refseq calls is artifactual (see Methods). Rerunning the phylogenetic test on the curated 210 ortholog sets for the remaining four genes, TEX15 is no longer significant at the 5% level 211 (p=0.086), possibly because the curation uncovered an intact TEX15 ortholog in anoles. 212 ZCWPW1 and ZCWPW2 are still highly significant; for FBXO47, the curation did not reveal any 213 discrepancy with the initial calls, so the p-value remains the same.

214 Our approach therefore uncovered two genes with clear-cut evidence of co-evolution with 215 PRDM9, the paralogs ZCWPW1 and ZCWPW2, and more tentative support for two others, TEX15 216 and FBX047. ZCWPW1, ZCWPW2 and TEX15 were among our initial list of 241 candidate genes 217 because they are co-expressed with PRDM9 in single cell testis data from mouse [31] (Figure 218 2B; Figure S6). FBX047 was not included by that criterion but because missense mutations in 219 the gene are associated with recombination rate variation in the total genetic map length in 220 humans, in both males and females [33]. Nonetheless, the expression of FBXO47 in mice is testis-221 specific [38], and the gene is expressed in the component in which PRDM9 had the highest 222 loading (albeit with a smaller loading [31] Figure 2B; see also [39]).

Like *PRDM9*, *ZCWPW1*, *ZCWPW2*, *FBXO47* and *TEX15* are inferred to have been present in the common ancestor of vertebrates. Below we describe the distribution of each of the four genes across the phylogeny of 189 species and the patterns that give rise to the evidence of statistical association with *PRDM9*--in particular, the correspondence between their distributions and that of 11 well supported losses of *PRDM9*, as well as of 9 species for which the status of *PRDM9* is uncertain.

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230 ZCWPW1 and PRDM9 co-evolution

Our finding that *ZCWPW1* is co-evolving with *PRDM9* (*p*=0.0019 in the curated set; **Table**1) is in line with previous reports of an association in vertebrates between the presence and

absence of ZCWPW1 and PRDM9 orthologs [26,27]. Here, we found an even tighter coupling of
PRDM9 and ZCWPW1 than previously documented. Specifically, we inferred 12 losses of
ZCWPW1 among 189 species used in our phylogenetic test, distributed across 17 species that
lack ZCWPW1 entirely and two species carrying partial ZCWPW1 genes (with the PWWP domain
but not the zf-CW domain; Table S8).

238 Seven of the ZCWPW1 losses occur among the 11 well supported losses of PRDM9: in 239 cypriniformes fish, percomorph fish (Euacanthomorphacea), siluriformes fish, polypteriformes 240 fish, osteoglossomorpha fish, birds, and Dicroglossidae frogs. An additional ZCWPW1 loss 241 occurred in the denticle herring (Denticeps clupeoides), a species for which the status of PRDM9 242 is uncertain. The remaining four losses of ZCWPW1 seem to break the pattern, in that they occur 243 in lineages containing a complete *PRDM9* gene. However, three are observed only in a single 244 species and may be spurious. Therefore, across the tree, there is only one well supported case 245 of a taxon with an intact *PRDM9* that has nonetheless lost *ZCWPW1*, supported by two closely 246 related species, the tiger snake (Notechis scutalus) and the eastern brown snake (Pseudonaja 247 textilis) (see Table S8 for details).

In mice as well as human cell lines, ZCWPW1 binds two marks laid down by PRDM9: the zf-CW domain binds H3K4me3 and the PWWP domain H3K36me3 [40–42]. Thus, the coevolution across vertebrates likely reflects a conserved molecular interaction between ZCWPW1 and PRDM9 as reader and writer of these dual histone modifications.

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53 **ZCWPW2** also co-evolves with **PRDM9**

Intriguingly, the strongest association with the presence or absence of *PRDM9* is that of the paralog of *ZCWPW1*, *ZCWPW2* ($p=5x10^{-6}$; **Table 1**). Among the 189 species, there are 12 independent losses, distributed across 21 species that appear to lack *ZCWPW2* altogether and three that contain partial *ZCWPW2* genes (two with the PWWP domain but not the zf-CW domain, and one with the reverse; **Table S8**).

259 Six of the ZCWPW2 losses occur among the 11 well supported losses of an intact PRDM9: 260 in percomorph fish, polypteriformes fish, Xenopus frogs, Dicroglossidae frogs, birds, and the 261 green anole. In order to distinguish whether the absence of ZCWPW2 in Xenopus and 262 Dicroglossidae frogs reflects a single loss or multiple events, we investigated the status of 263 ZCWPW2 in an additional species of frog with PRDM9 (Ranitomeya imitator). We were able to 264 successfully identify a complete ZCWPW2 ortholog in this species, suggesting that ZCWPW2 has 265 indeed been lost at least twice within frogs, possibly coincidentally with *PRDM9* in each case. 266 ZCWPW2 is also absent in a clade encompassing cypriniformes fish and siluriformes fish, as well as the electric eel (*Electrophorus electricus*), which has an intact *PRDM9*. This phylogenetic distribution suggests that the loss of *ZCWPW2* may have occurred before the losses of *PRDM9* in both cypriniformes fish and siluriformes fish. Also suggestive of this order of loss, *ZCWPW2* is absent in osteoglossomorpha fish (the Asian arowana, *Scleropages formosus*); in this case, the gene is also absent from the closest evolutionary relative in the tree, the elephantfish (*Paramormyrops kingsleyae*), which carries *PRDM9*.

Among the nine species for which the status of *PRDM9* is uncertain, *ZCWPW2* is absent in the denticle herring (*Denticeps clupeoides*). The remaining three cases of *ZCWPW2* loss are each observed in a single species carrying an intact PRDM9, without supporting lines of evidence. In summary, in the few cases with *PRDM9* but not *ZCWPW2*, we cannot verify the loss of *ZCWPW2*; conversely, the only species with *ZCWPW2* but that clearly lack *PRDM9* are canids and the platypus, the two lineages that experienced the most recent losses of *PRDM9* (see **Table S8** for details).

Like its paralog, ZCWPW2 contains zf-CW and PWWP domains, predicted to bind H3K4me3 and H3K36me3, respectively (**Figure 4A, Figure S10**). As in ZCWPW1 ([27], [26]), these domains are highly conserved, especially at residues with predicted binding properties (**Figure 4B-C**), suggesting that ZCWPW2 is also recruited to sites of PRDM9 binding.

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The distribution of FBXO47 and TEX15 orthologs

We identified two additional genes, FBX047 and TEX15, that may be co-evolving with 286 287 *PRDM9*: using the curated calls, p=0.016 and p=0.087, respectively (**Table 1**). TEX15 is co-288 expressed with PRDM9 in two components inferred from single cell data from mice, active during 289 pre-leptotene and zygotene (Figure S6). The statistical evidence for co-evolution stems from the 290 fact that TEX15 is missing in two taxa lacking PRDM9: birds and percomorph fish. TEX15 is also 291 absent in the Atlantic cod (Gadus morhua), suggesting that the loss of TEX15 that led to its 292 absence in percomorph fish occurred before that of PRDM9. All of the other 189 species 293 considered have an intact TEX15 (see Table S8 for details).

The statistical evidence is a bit stronger for *FBX047*, which has been lost five times in the absence of *PRDM9*: in cypriniformes fish, osteoglossomorpha fish, siluriformes fish, and in *Xenopus* and *Bufonidae* frogs. Intriguingly, *FBX047* is additionally absent in the electric eel, a species that carries a complete *PRDM9* gene, but lacks both *ZCWPW1* and *ZCWPW2*. Testing for the co-evolution of the candidate genes with each other, a null model in which the state transitions of *FBX047*, *ZCWPW1* and *ZCWPW2* are independent is rejected for all pairs of genes (maximal p<6x10⁻³; Table S9A), and p-values are lower for *FBXO47* and *ZCWPW1*, or *FBXO47*and *ZCWPW2*, than for *FBXO47* and *PRDM9*.

302 In summary, by extending the reconstruction of *PRDM9* to 446 vertebrate species, we 303 identified thirteen losses that are supported by more than one species or by independent 304 evidence, and possibly as many as 23. Focusing on a subset of 189 species that capture eleven 305 state transitions of *PRDM9*, we tested whether *PRDM9* transitions coincide with those of 139 306 candidate genes lost at least once across vertebrates. After carefully vetting the ortholog calls for 307 our top five signals, we identified two genes that are clearly co-evolving in their presence and 308 absence with PRDM9, ZCWPW1 and its paralog ZCWPW2, and two for which the evidence is 309 weaker: FBXO47 and most tentatively, TEX15.

310

311 **Discussion**

312 Dual roles of PRDM9 across vertebrates

We had previously hypothesized that PRDM9 plays a role in directing recombination not only in mammals but across vertebrates, based on the presence of an intact ortholog across vertebrates with a rapidly-evolving zinc finger [11]. Consistent with our prediction, there is tentative evidence for the influence of PRDM9 binding on recombination in rattlesnakes [24]. That a gene with a known role in recombination, *ZCWPW1* co-evolves with *PRDM9* across vertebrates lends further support to this hypothesis.

319 The precise nature of the molecular interactions between PRDM9 and ZCWPW1 remains 320 unknown, but recent evidence suggests that ZCWPW1 interacts with PRDM9 to facilitate the 321 repair of PRDM9-dependent DSBs: notably, Zcwpw1-/- male mice and older female mice are 322 sterile [27,43] and exhibit defects in their ability to repair DSBs [25-27]. In turn, the genomic 323 locations of DSBs are not altered in Zcwpw1-/- mice, indicating that the gene does not play a role 324 in DSB positioning [25–27]. In light of these experimental results, the co-evolution of *PRDM9* with 325 ZCWPW1 across vertebrates indicates that PRDM9 likely plays a role in the efficient repair of 326 DSBs not only in mice and humans [25,26,44,45], but across the vertebrate phylogeny.

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328 Nature of the co-evolution of candidate genes with PRDM9

If a gene interacts with PRDM9 by reading its histone modifications, as is the case for ZCWPW1 [25–27] and likely ZCWPW2 (**Figure 4**), and has no other roles, we would expect that gene to be dispensable in species that no longer have an active PRDM9 SET domain. Previous papers reported that *ZCWPW1* is more likely to be missing from ray-finned fish with substitutions in catalytic tyrosine residues of the SET domain, in addition to clades lacking the entire *PRDM9* gene [26,27]. In our analysis, we find that both *ZCWPW1* and *ZCWPW2* are more likely to be
absent from species carrying only *PRDM9* orthologs with substitutions in at least one catalytic
tyrosine residue, as well as those lacking *PRDM9* altogether (Figure 3).

337 While this pattern suggests a dependence of ZCWPW1 and ZCWPW2 on the intact 338 catalytic activity of PRDM9, the interpretation is complicated by the fact that all species with 339 substitutions at the tyrosine residues in all PRDM9 copies are also carrying only partial PRDM9 340 orthologs lacking KRAB and SSXRD domains, and nearly all species with conserved tyrosine 341 residues also carry a complete copy of *PRDM9*. In that regard, the few exceptions are informative: 342 among species with confident PRDM9 calls, the platypus and siluriformes fish carry PRDM9 343 orthologs putatively missing the KRAB domain but with intact tyrosine residues. ZCWPW2 is 344 absent from all three considered siluriformes fish species while ZCWPW1 is absent from one. 345 Thus, the presence of ZCWPW1 and ZCWPW2 may depend on that of the KRAB domain rather 346 than, or in addition to, the tyrosine residues remaining intact.

347 Similar considerations suggest that in the rare lineages where ZCWPW1, ZCWPW2, 348 FBX047 and TEX15 are present in the absence of PRDM9, we might expect the genes to be 349 under relaxed selective constraint. To examine this prediction, we tested whether $\omega = dn/ds$ was 350 higher in lineages without a complete PRDM9 (where dn is the rate of non-synonymous 351 substitutions and ds the rate of synonymous substitutions; see Methods). For ZCWPW1, there 352 was no evidence for a relaxation of selection (p>0.13; **Table S10**). The intriguing exception is in 353 platypus, one of the few species that has a SET domain with intact tyrosine residues but is lacking 354 the KRAB domain (p=0.038; in all other cases, p>0.13; **Table S10**). This observation lends further 355 support to the notion that the conservation of *ZCWPW1* may depend on the KRAB domain rather 356 than, or in addition to, the tyrosine residues of *PRDM9*.

357 For ZCWPW2, the same test revealed that a model in which all species have the same 358 ω is significantly less likely than one in which ω is elevated in lineages lacking a complete *PRDM9*; 359 in particular, in canids and platypus (p=0.0003; Table S10). In fact, in canid lineages (fox and 360 dogs), for which the loss of PRDM9 is ancestral, ZCWPW2 is no longer under any discernible 361 selective constraint (testing a null model of $\omega = 1$, p = 0.307; **Table S10**). Considered together with 362 the observation that ZCWPW2 is absent from all the other lineages in which a complete PRDM9 363 gene is clearly absent, these evolutionary analyses suggest that ZCWPW2 is dispensable in the 364 absence of a complete PRDM9 ortholog.

The molecular function of ZCWPW2 is to our knowledge unknown. Like its paralog, it could be involved in the processing or repair of DSBs. If so, the observation that *Zcwpw1-/-* mice show defective DSB processing and repair [25–27] suggests that the role of ZCWPW2 cannot be 368 completely redundant with that of its paralog. Alternatively, by reading the dual marks laid down 369 by PRDM9, ZCWPW2 might help to recruit the recombination machinery (in particular SPO11) 370 and thus play an earlier role in the positioning of DSBs. While in yeast, the link between histone 371 modifications (specifically, H3K4me3) and the recruitment of Spo11 is made by Spp1 [46], in 372 mammals, the ortholog of Spp1, CXXC1, is not essential for meiosis [47], and the gene that plays 373 the analogous role has not yet been identified. Our analysis highlights ZCWPW2 as a potential 374 candidate for this role, to be tested experimentally.

375 For TEX15, ω is also higher in lineages where PRDM9 is absent or incomplete (p=0.0036) 376 for fish and p=0.015 for mammals), but remains significantly below 1 (**Table S10**), indicating that 377 in the absence of PRDM9, TEX15 is not dispensable. If TEX15 and PRDM9 are indeed co-378 evolving, the relationship is likely to be indirect; for instance, recent work implicates TEX15 as an 379 effector of piRNA-mediated transposable element (TE) methylation and silencing [48,49]. Male 380 mouse knockouts of Tex15 exhibit a meiotic arrest phenotype associated with the failure to repair 381 DSBs and to undergo chromosomal synapsis [45], as well as the transcriptional activation of TEs 382 [48,49]. This phenotype is similar to those observed in mouse knockouts of other piRNA-pathway 383 genes, such as Miwi or Dnmt3 [50]. In Dnmt3 knockout mice, it has been shown that TEs 384 accumulate both H3K4me3 marks and SPO11-dependent DSBs, suggesting that the methylation 385 of TEs serves not only to silence them, but may also result in preventing their use as sites of 386 recombination [50]. Thus, TEX15 could conceivably play an important but indirect role in 387 preventing the binding of PRDM9 to TEs.

For FBXO47, ω is higher in fish lineages where PRDM9 is absent or incomplete 388 389 (p=0.0023), but remains significantly below 1, while in mammals, there is no evidence for 390 relaxation of constraint (Table S10). Like TEX15, if FBX047 and PRDM9 are co-evolving, the 391 relationship is likely to be indirect. FBX047 is a member of the F-box protein family, which act as 392 recognition subunits of Skp1-Cullin1-F-Box protein (SCF) E3 ubiquitin ligase complexes [38,51]. 393 Recently, FBXO47 has been implicated as a key regulator of the telomere shelterin complex 394 during meiotic prophase I, and in mice is necessary for telomere nuclear envelope attachment 395 and subsequent events, including DSB repair [38]. One possibility for increased conservation of 396 FBX047 in the presence of PRDM9 would be if this role of FBX047 contributes to the formation 397 of a chromatin environment that aids in the repair of PRDM9-dependent DSBs, or possibly in the 398 recruitment of ZCWPW1.

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401 Which loss came first?

402 While PRDM9 has two distinct roles—in specifying the location of DSBs and in facilitating 403 their repair-the four candidate genes that we have identified may only be involved in one of 404 these two roles. If so, the dependencies between the presence of PRDM9 and of these genes 405 may be asymmetric. For instance, if we ignore possible pleiotropic roles of the candidate genes, 406 and assume ZCWPW1 and FBX047 play roles in, or related to, repair but not DSB localization, 407 we would predict that their loss occurs after that of PRDM9 (as appears to have been the case 408 in Tachysurus fulvidraco for both genes, and for FBXO47 in Xenopus laevis; Table S8B). In 409 contrast, if ZCWPW2 is involved in DSB localization but not repair, we would predict it could be 410 lost before *PRDM9* (as was seemingly the case in two lineages of ray-finned fish; **Table S8B**). 411 The phylogenetic data considered here do not allow us to distinguish between these scenarios: 412 there is statistical evidence for a dependence of state transitions of ZCWPW1, ZCWPW2, 413 FBX047 and TEX15 on PRDM9 as well as vice versa (in all tests, maximum p<0.07, testing the 414 null model of no dependence against either dependence as an alternative model; **Table S11**). 415 These scenarios could potentially be distinguished by collecting more fine-grained phylogenetic 416 information to pinpoint the specific lineages in which the first loss occurred, as well as in light of 417 further experimental data.

418

419 **Outlook**

420 Our phylogenetic analysis allowed us to identify novel putative interactors of PRDM9 that 421 are promising candidates for functional studies. For this analysis, the power comes from the 422 repeated losses of *PRDM9*—in our case, from eleven transitions from presence to absence. 423 Confounding these kinds of analyses, however, are issues of data quality and in particular 424 absences of complete PRDM9 orthologs that reflect poor genome quality rather than true losses. 425 To address this issue, we validated any absence in Refseq with whole genome searches and 426 where possible, de novo assemblies from RNA-seq data, leading us to realize that in one case 427 (*MEI1*), the apparent co-evolution with *PRDM9* was in fact spurious.

428 A more subtle but related issue stems from a phylogenetic signal of genome quality, which 429 can lead to apparent clustering of losses. To minimize this issue, we restricted our analysis to 430 genomes that included most "core" eukaryotic genes (Fig. S7) and downsampled our tree to 431 include at most three species below every inferred PRDM9 loss. As genome qualities improve 432 and as their assemblies become more uniform (eg., [52]), these issues should be alleviated. 433 Moreover, as species are added to the phylogeny, additional losses will be identified: as one 434 example, our identification of two species of frogs with a complete PRDM9 revealed that PRDM9 435 had not been lost once in the common ancestor, as had been inferred using fewer species by

Baker et al. (2017), but has instead been lost more than once within amphibians. This discovery
also suggests that frogs may be an interesting clade within which to study the steps by which
PRDM9 and its partners are lost.

439 Beyond the application to *PRDM9* and meiotic recombination, our analysis illustrates how 440 long-standing phylogenetic approaches can now be applied to comparative genomic data to 441 identify novel molecular interactions [53]. Such analyses need not be restricted to measurements 442 of presence or absence of whole genes, as we have done here, but could focus exclusively on 443 specific domains, indicative of specific subfunctions, or consider how rates of evolution in specific 444 domains depend on the presence or absence of other genes. With the explosion of high quality 445 and more representative sets of genomes now coming on line (e.g., [52], [54]), and the 446 development of statistical methods that consider both binary and continuous character evolution 447 jointly, we expect this type of approach to become increasingly widespread.

448

449 Material and Methods

450

1. Identification of PRDM9 orthologs

451 As a first step towards characterizing the distribution of *PRDM9* in vertebrates, we 452 identified putative PRDM9 orthologs in the RefSeq database with a blastp search [30], using the 453 N-terminal portion of the Homo sapiens PRDM9 protein sequence containing KRAB, SSXRD and 454 SET domains as the query sequence (RefSeq accession: NP_001297143; amino acid residues 455 1-364). We downloaded the corresponding GenBank file for 5,000 hits (3,400 unique genes from 456 412 species) and characterized the presence or absence of KRAB, SSXRD and SET domains for 457 each record using the Conserved Domain, Protein Families, NCBI curated and SMART databases 458 (CDD [55]; Pfam (REF); NCBI curated (REF); SMART (REF); accessions cl02581 and cl09744 459 for the KRAB and SSXRD domains respectively, and accessions cl40432 and cl02566 for the 460 SET domain), annotating each domain as present if that domain had an e-value less than 1 in 461 any of the four databases. We then removed alternative transcripts from the dataset by 462 preferentially keeping, for each unique gene, the transcript with the maximal number of annotated 463 domains. When there were multiple transcripts with the same maximal number of domains, we 464 kept the longest one.

Because *PRDM9* shares its SET domain with other PRDM family genes and its N-terminal domains with members of the KRAB-ZF and SSX gene families, many of these hits are potential PRDM9 paralogs. To identify bona fide *PRDM9* orthologs from this initial set of genes, we sought to build phylogenetic trees specific to the KRAB, SSXRD, and SET domains and remove 469 homologs that cluster with genes annotated as distantly related paralogs of *PRDM9*. To this end, 470 we extracted the amino acid sequences for complete KRAB, SSXRD, and SET domains, and for 471 each domain, constructed neighbor-joining trees using Clustal Omega [56]. Utilizing the KRAB 472 and SSXRD domain-based trees, we identified and removed 87 genes that visually cluster with 473 members of the SSX gene family (Fig. S1A-B). Analyzing the SET domain-based tree, we 474 identified and removed 2,637 genes that group with other members of the *PRDM* gene family 475 (Fig. S1C; see figure legend for details). We ultimately retained 625 genes, each of which cluster 476 with *PRDM9* in one or more of these trees.

By this approach, in the 412 species considered, we identified 209 *PRDM9* orthologs containing KRAB, SSXRD and SET domains from 155 species, as well as 13 *PRDM9* orthologs containing KRAB and SET domains for which we were unable to detect an SSXRD domain with an e-value less than 1 from an additional 11 species. For the 246 species for which we were unable to identify a *PRDM9* ortholog spanning KRAB and SET domains in our initial search of the RefSeq database, we sought to verify that *PRDM9* was truly absent using a number of approaches.

484 As a first step, we performed an additional blastp search against the non-redundant 485 protein sequence (nr) database, targeting only those species in order to identify any annotated 486 gene record missed in our initial search of the RefSeq database. We downloaded the 487 corresponding GenBank file for each hit with >55% coverage and >40% identity and, after 488 removing records corresponding to those we had previously identified, annotated domains and 489 removed alternative transcripts as before. We then verified the orthology of the remaining records 490 by blasting each protein sequence against the human RefSeq database, accepting it as a PRDM9 491 ortholog if the top hit was PRDM9 or its paralog PRDM7. This approach enabled the identification 492 of an additional 9 PRDM9 orthologs, including one containing KRAB, SSXRD and SET domains, 493 and one containing KRAB and SET domains.

494 Next, we performed a series of *tblastn* searches of the whole genome of the 244 species 495 remaining using the N-terminal portion of the Homo sapiens PRDM9 protein as a query. When 496 we were unable to retrieve any promising hits with the human protein sequence, we re-performed 497 the *tblastn* search using the N-terminal portion of a *PRDM9* ortholog from a species closely related 498 to the focal species. In order to identify which of the identified contigs corresponded to genuine 499 PRDM9 orthologs (as opposed to paralogs such as PRDM11), we performed blast psearches 500 against the Homo sapiens RefSeg database using the aligned protein sequences as query 501 sequences. Contigs containing the relevant alignments spanning KRAB and/or SET domains 502 were then downloaded and the aligned region including 10,000 of flanking sequence was

503 extracted and input into Genewise [57], using the PRDM9 protein sequence from Homo sapiens 504 or a closely related species as a guide sequence (see **Table S2** for details). In genomes from 10 505 species, we identified separate contigs containing the KRAB domain and the SET domain. In 506 these cases, the contigs were concatenated before use as input in *Genewise*. These approaches 507 enabled us to identify an additional 53 PRDM9 orthologs from 33 species, including 21 PRDM9 508 orthologs containing KRAB, SSXRD and SET domains from 21 species, and 24 PRDM9 orthologs 509 containing KRAB and SET domains but for which we were unable to identify the SSXRD domain 510 from 11 species.

511 These analyses left 210 species for which we were unable to identify a *PRDM9* ortholog 512 with both KRAB and SET domains. For these species, with the exception of 94 birds and 513 crocodiles and 78 percomorpha fish, we additionally searched testis RNA-seq datasets when 514 possible, including those generated for this study (see below; **Table S3**); this approach enabled 515 us to identify two additional *PRDM9* orthologs containing KRAB and SET domains from two 516 species of fish.

517 From this analysis, and given the phylogenetic relationships among species given by the 518 TimeTree tool [35], we inferred 20 putative complete or partial losses of PRDM9 across the 412 519 species represented in the RefSeg database. Of these, 7 losses were supported by the absence 520 of PRDM9 in two or more closely related species: in percomorpha and beryciformes fish, 521 characiformes and siluriformes fish, cypriniformes fish, polypteridae fish, frogs, birds and 522 crocodiles, and canids. The remaining 13 inferred losses each corresponded to an individual 523 species. In order to identify whether or not any of these 13 latter absences could be supported by 524 additional species, and to more accurately infer the dates of each loss, we sought to investigate 525 the status of PRDM9 in species closely related to each putative loss event. To this end, we 526 investigated the whole genomes of an additional 18 species and RNA-seq datasets from an 527 additional 4 species as before, with one species represented by both a whole genome sequence 528 and a corresponding RNA-seq dataset (Ambystoma mexicanum). This approach enabled us to 529 identify an additional 6 PRDM9 orthologs containing KRAB, SSXRD and SET domains from 6 530 species, as well 15 additional species putatively lacking a complete PRDM9 gene. In doing so, 531 we found that 2 additional losses were supported by the absence of PRDM9 in two or more closely 532 related species: in osteoglossomorpha fish, as well as a loss within lizards shared by Anolis 533 carolinensis and Scleropages formosus. For each of the 11 remaining instances in which only a 534 single species was found to be lacking PRDM9, the most closely related species considered 535 possessed a complete PRDM9 ortholog. While independent work supports our finding of a partial 536 loss of PRDM9 in platypus (J. Hussin and P. Donnelly, personal communication), we do not have

537 confirmatory evidence of absence for the remaining 10 species, and therefor treat these species 538 as having an uncertain PRDM9 status. Moreover, we identified two species of frogs carrying 539 complete PRDM9 orthologs. This discovery suggests that PRDM9 has been lost repeatedly within 540 amphibians – at least once in salamanders, and at least three times within frogs (with each of 541 these four putative loss events being supported by the absence of the PRDM9 in two or more 542 closely related species).

543 Lastly, we include in the list of species considered an additional 13 species for which we 544 had previously identified complete PRDM9 orthologs [11] but which were not directly examined 545 here (Table S4). Altogether, this pipeline resulted in the identification of 193 species in which we 546 find a complete *PRDM9* ortholog containing KRAB, SSXRD and SET domains, 26 species for 547 which we identify *PRDM9* orthologs containing KRAB and SET domains but not SSXRD domains, 548 218 species for which we have evidence for the absence of a complete PRDM9 gene, and 9 549 species for which we were unable to make a confident determination (see Tables S1-S4, Figure 550 1).

551 For each of the *PRDM9* orthologs that we identified, we characterized the conservation of 552 three key tyrosine residues that have been shown to underlie the catalytic function of the human 553 SET domain in vitro (i.e., Y276, Y341, and Y357; [58]) and for Y357, in vivo in mouse [10]. To this 554 end, we constructed an alignment of the SET domain using Clustal Omega [56] and extracted the 555 residues aligning to the human tyrosine residues from each of 678 SET domains (**Table S1**).

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2. Verification of genomic calls using RNA-seq data

558 For four species in which we identified no *PRDM9* ortholog or only a partial ortholog, we 559 investigated whether a complete *PRDM9* ortholog may nonetheless be present using RNA-seq 560 data. We therefore sought to verify its absence from *Anolis carolinensis*, a species in which we 561 had been unable to find a *PRDM9* ortholog in the genome assembly or Refseq, as well as a 562 second reptile species, *Sceloporus undulatus*, for which Refseq data and a genome sequence 563 were not available. To this end, we built a *de novo* RNA transcriptome assembly and tested for 564 the expression of PRDM9 in testis and other tissue samples (see below).

565 Similarly, in two species in which we had originally identified only a partial ortholog of 566 *PRDM9* (*Astyanax mexicanus* and *Clupea harengus*), we wanted to verify the incomplete domain 567 structure inferred from the genome sequence by conducting a *de novo* transcriptome assembly 568 (in *Clupea harengus*, this analysis turned out to be unnecessary, as an updated reference 569 genome, GCA_000966335.1, contains a complete *PRDM9*). To this end, we analyzed RNA-seq

570 data from male gonad surface from *Astyanax mexicanus* and liver and testis from *Clupea* 571 *harengus*.

572 Dissected tissue samples preserved in RNAlater were kindly provided to us by Arild 573 Folkvord and Leif Andersson (Clupea harengus), Cliff Tabin (Astyanax mexicanus), Tonia 574 Schwartz and Tracy Langkilde (Sceloporus undulatus), and Athanasia Tzika (Anolis carolinensis). 575 These samples were stored at -20°C until extraction and library preparation. Total RNA was 576 extracted using the Qiagen RNeasy kit (Valencia, CA, USA) following the manufacturer's protocol. 577 RNA was guantified and assessed for guality on a Qubit fluorometer and approximately 1 µg of 578 total RNA was input for library preparation using the Kapa RNA-seq kit. Samples were prepared 579 following the manufacturer's protocol, except that half reactions were used. Briefly, mRNA was 580 purified using manufacturer's beads and chemically fragmented. First and second-strand cDNA 581 was synthesized and end-repaired. Following A-tailing, each sample was individually barcoded 582 with an Illumina index and amplified for 12 cycles. In order to evaluate the library quality and size 583 distribution, libraries were evaluated on an Agilent Tapestation. The libraries were then 584 sequenced over two runs on the NextSeq 550 at Columbia University to collect paired-end 150 585 bp reads.

586 Illumina sequencing reads (248,820,547 2x150 base pair (bp) paired-end reads) were 587 demultiplexed into individual sample fastq files with the software bcl2fastq2 (v2.20.0, Illumina). 588 The FastQC software [59] was used for visual inspection of read quality. Adapters and low-quality 589 reads were trimmed with the Trimmomatic software, which is bundled as a plugin within the Trinity 590 de novo assembler [60] (v2.8.5) and was enabled using the --trimmomatic flag. The default 591 trimming settings (phredscore>=5; slidingwindow:4:5; leading:5, trailing:5; minlen:25) were used 592 following [61] recommendations. The pair-end reads were trimmed and de novo transcriptomes 593 assembled with Trinity (v2.8.5) using the following parameters: --seqType fg --SS lib type FR --594 max memory 100G --min kmer cov 1 --trimmomatic --CPU 32. Details on assembly quality are 595 shown in Fig. S2. Gene expression data for all four species (Anolis carolinensis, Sceloporus 596 undulatus, Clupea harengus, Sceloporus undulatus) are available from the NCBI sequence read 597 archive (Bioproject PRJNA605699, SRA accessions: SRR11050679-SRR11050687).

To evaluate whether PRDM9 was present in the transcriptome data, we conducted a *tblastn* search (e-value \leq 1e-5) against each *de novo* assembly using the human PRDM9 protein sequence (without its rapidly evolving zinc finger array) as a query, and we classified the domain presence of up to five top hits using CDD blast [55]. For a given species, if the KRAB and SET domains were not identified in any transcript, *PRDM9* was considered incomplete. The inability to

603 identify PRDM9 could indicate either that the gene is not expressed or that we lack the appropriate 604 cell types or sequence coverage to detect it. To assess our power to detect PRDM9 from the 605 testis RNA-seq data, we followed methods outlined in [11]. Specifically, for each transcriptome, 606 we evaluated whether we could identify transcripts from six genes with highly conserved roles in 607 meiotic recombination [62] (HORMAD1, MEI4, MRE11A, RAD50, REC114, and SPO11). To 608 identify the transcripts orthologous to each of these genes, we performed a tblastn search (e-609 value \leq 1e-5) of the Homo sapiens reference protein sequence against each de novo transcriptome. We considered PRDM9 to be absent if we detected expression of all six genes but 610 611 not a complete *PRDM9*; by these criteria, we found PRDM9 to be missing from *A. carolinensis*, 612 S. undulatus, and A. mexicanus.

Using the same approach to *de novo* assembly and gene detection, we also analyzed publicly available RNA-seq datasets from testis for 28 additional species (**Table S3**), either to verify the absence of PRDM9 (see above) or of one of the candidate genes (see below).

To estimate the expression levels of the Trinity-reconstructed transcripts, we used RSEM [63] (v1.3.1) implemented through Trinity (v2.8.5). We first aligned the RNA-seq reads from each sample to the newly generated *de novo* assembled transcriptome (see above) using the alignment method bowtie [64] (v1.2.2). We then extracted quantification information for each gene of interest from the RSEM output (in fragments per kilobase of transcript per million mapped reads or FPKM) (**Fig. S3**).

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3. Choice of candidate genes and orthology assignments

624 To identify a set of genes that may co-evolve with *PRDM9*, we relied on three publicly 625 available datasets, namely: (i) 39 genes associated with variation in recombination phenotypes in 626 a genome-wide association study in humans [33]. Of the variants reported to be associated with 627 recombination phenotypes, six were found in intergenic regions; we included the subset of two 628 cases in which the authors assigned these variants to nearby genes (ZNF84 and ZNF140). (ii) 629 193 genes co-expressed with PRDM9 in single cell data from mouse testes. Specifically, we 630 considered the top 1% of genes based on their gene expression loadings in component 5, the 631 component in which PRDM9 has the highest loading [31]. (iii) 36 genes known to have a role in 632 mammalian meiotic recombination based on functional studies [32].

633 Genes co-expressed with PRDM9 in mouse spermatogenesis were converted to human 634 gene symbols using the package biomaRt in R [65]. Fifteen of these genes did not have an 635 orthologous human gene symbol (*Gm7972*, *H2-K1*, *Gm4349*, *Ddx43*, *Atad2*, *XIr4c*, *Gm364*,

Tex16, 4933427D06Rik, Al481877, H2-D1, Trap1a, XIr4a, 2310035C23Rik, and Tmem5) and
eight other genes mapped to more than one human gene symbol (*Msh5, Cbwd1, Nxf2, Cbwd1, Fam90a1b, Srgap2, Cdk11b, Gm15262*). Keeping all mapped gene symbols yielded 185 genes;
combined with the two other sources, 241 genes were tested for their co-evolution with *PRDM9*(**Figure S6**). A supplementary file describing each meiosis candidate gene is available in **Table S5**.

642 For the 241 genes, we characterized whether the ortholog is present in its complete form 643 across vertebrate species. To this end, we first downloaded all the vertebrate RefSeq protein 644 sequences available on the NCBI database (accessed on June 3, 2020), corresponding to 339 645 species. Of these, we filtered out 32 species that were missing 10 or more BUSCO core genes 646 (out of a total of 255 genes) [66], reasoning that their genomes were sufficiently incomplete that 647 they may be missing orthologs by chance (see **Figure S7**). Of the remaining 307 species, we 648 further excluded 29 species in order to remove polytomies observed in the phylogeny; specifically, 649 we removed the minimal number of species necessary to remove each polytomy while preserving 650 any transitions in the state of *PRDM9*. Moreover, to minimize possible phylogenetic signals 651 generated by genome assembly quality, we thinned the tree such that for each PRDM9 loss along 652 the phylogeny, we kept at most three species representing that loss. In cases where a loss was 653 ancestral to more than three species in our dataset, we picked three distantly related species with 654 the best genome assemblies, as measured by the BUSCO score. In the end, we retained 189 655 species: 134 mammals, 3 birds, 6 amphibians, 18 reptiles, 2 percomorph fish, 3 cypriniformes 656 fish, 20 other ray-finned fish, 2 cartilaginous fish, and one jawless fish. This phylogeny includes 657 representative species for 11 of the 13 inferred PRDM9 losses: species of Bufonidae frogs and 658 salamanders were not included due to the absence of available gene annotations: also due to the 659 lack of gene annotations for frog species with PRDM9, within these 189 species, the losses in 660 Xenopus and Dicroglossidae frogs cannot be distinguished from a single event.

661 For each candidate gene in each species, we performed a blastp search of the human 662 ortholog against the RefSeq database of the species and kept up to five top hits obtained at an 663 e-value threshold of 1e-5. We inferred the domain structure of each hit using the Batch CD-Search 664 [67], and considered a domain as present in a species if the e-value was ≤ 0.1 . We considered genes to be complete orthologs if they contained the superfamily domains found in four 665 666 representative species of the vertebrates phylogeny carrying a complete PRDM9 (Esox lucius 667 (fish) Geotrypetes seraphini (caecilian), and Pseudonaja textilis (snake)), at an e-value threshold 668 of 1e-4. For the 15 genes (FANCB, FMR1NB, GPR137C, HAUS8, M1AP, MEI1, SPATA22,

669 *CLSPN*, *FBXO47*, *HMGA2*, *HSF2BP*, *IQCB1*, *LRRC42*, *PRAME*, *SYCE2*) in which no detectable 670 domains were present, we annotated the presence or absence of the gene using the blastp results 671 alone. In the end, we built a matrix of presence or absence across species and candidate genes 672 to be used in the phylogenetic test (see **Table S6**).

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4. Testing for the co-evolution of PRDM9 and candidate genes

To test for the co-evolution of *PRDM9* and each candidate gene, we need to account for the phylogenetic relationships among the species considered. To obtain these relationships and time-calibrated branch lengths, we used the TimeTree resource (<u>http://timetree.org/</u>; [35], accessed on June 10, 2020). Of the 189 species included in the phylogenetic tests, 9 were not present in the TimeTree database; in those cases, we used information from a close evolutionary relative to determine their placement and branch lengths.

For this test, we consider *PRDM9* as present if it contains KRAB and SET domains or incomplete/missing if one of those domains is absent (**Tables S4** and **S8**). We do not rely on the SSXRD domain when making these calls because its short length makes its detection at a given e-value threshold unreliable. Notably, for 19 of the 26 species with *PRDM9* orthologs containing KRAB and SET domains, but not SSXRD domains with an e-value < 1, we are able to detect the SSXRD domain when using an e-value threshold of 1000 (**Table S1**). We additionally do not rely on the ZF array because its repetitive nature makes it difficult to sequence reliably.

688 We tested whether state changes of intact candidate genes were unexpectedly coincident 689 with state changes of the intact *PRDM9* using the software *BayesTraitsV3* [68]. We did so by 690 comparing the statistical support for two models: a null model in which *PRDM9* and a given 691 candidate gene evolve independently of one another along the phylogeny versus an alternative 692 model in which the gain ("1") and loss ("0") of a gene is dependent on the status of *PRDM9* and 693 vice versa. We compared the likelihoods of the two models using a likelihood ratio test with 4 694 degrees of freedom, and reported a p-value uncorrected for multiple tests (Table S7). For each 695 gene and model, 100 maximum likelihood tries were computed and the maximum likelihood value 696 was retained. A quantile-quantile plot was drawn to access the distribution of p-value, and the R 697 package "Haplin" was used to compute pointwise confidence intervals (CI). To control for the false 698 discovery rate (FDR), we computed q-values using the R package "qvalue" and set a 50% FDR 699 threshold.

Given the phylogenetic distribution of *PRDM9*, it is likely that a *PRDM9* ortholog was present in the common ancestor of vertebrates [11,12]. Based on this prior knowledge, we restricted the state of *PRDM9* at the root of the phylogeny to always be present. In turn, for each

candidate gene, we set a prior in which it had 50% probability of being present and 50% probability
of being absent. We also used this prior for the state of *PRDM9* in the 9 species that lack *PRDM9*but where the loss was not supported by a closely related species (i.e., for which we considered
the status uncertain).

For *FBXO47, TEX15, ZCWPW1* and *ZCWPW2*, we also explored restrictions on the rates in the dependent model, such that their state transitions depend on PRDM9 (model X) or the state transitions of *PRDM9* depends on theirs (model Y), rather than both being true. For these tests, we compared the likelihoods of each dependent model against our independent null model using a likelihood ratio test with 2 degrees of freedom. For each gene and model, 100 maximum likelihood tries were computed and the maximum likelihood value was retained.

713 We also explored whether redefining a complete PRDM9 ortholog as containing not only 714 the KRAB and SET domain but also the SSXRD domain would change the statistical significance. 715 By using the improved calls (see below), only *ZCWPW2* remains significant (p=0.004) and 716 *ZCWPW1* marginally so (p=0.056) (**Table S9B-C**).

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5. Improving gene status calls of top candidate genes

719 For the five genes with a FDR \leq 50% (**Figure 2A**), we sought to improve our calls by 720 building phylogenetic trees based on domains in the genes and examining the clustering patterns 721 visually, as well as by searching for orthologs in whole genome assemblies and testis 722 transcriptomes (following the same procedures described for PRDM9). These improved calls 723 were then used to rerun the phylogenetic independent contrast tests, following the same 724 implementation as previously; the p-values for these improved gene models are shown alongside 725 the original ones in **Table 1**. Below we provide an overview of the steps we took for each 726 candidate gene. For each gene we provide descriptions of identified orthologs and how they were 727 identified in **Table S1**, specific details about orthologs identified from whole genome assemblies 728 in **Table S2**, our improved calls per species in **Table S8A**, and a summary of loss events in **Table** 729 S8B.

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731 i. *MEI1*

For *MEI1*, an initial blastp search of the vertebrate RefSeq database using the human sequence as query resulted in the identification of 422 *MEI* orthologs from 372 species. We note that, for *MEI1*, we did not find any domain annotations, and therefore did not perform phylogenetic analysis to support the identification of these orthologs. However, each homolog identified in our initial RefSeq analysis was annotated as either *MEI1* or *MEI1-like*. We thus labeled each species

as having a complete ortholog if an ortholog was present. This approach resulted in the
identification of *MEI1* ortholog for 187 of the 189 species used for our co-evolutionary test. For
the remaining 2 species, we sought to identify *MEI1* orthologs from whole genome sequences
following the same procedures described for *PRDM9*. This approach allowed us to identify a *MEI1*ortholog in every species, revealing that in fact, *MEI1* has not been lost among the vertebrate
species examined (**Tables S1** and **S2**).

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744 ii. ZCWPW1 and ZCWPW2

745 Because ZCWPW1 and ZCWPW2 are paralogs, we performed our analyses of these 746 genes together. To this end, we combined the datasets of genes identified in our initial RefSeg 747 blastp search to create a dataset of 977 putative orthologs from 363 species. We then extracted 748 amino acid sequences and built neighbor-joining trees using Clustal Omega for both the zf-CW 749 and PWWP domains ([56]; accessions cl06504 and cl02554; Figure S9A-B). Utilizing these trees, 750 we removed 573 genes that visually clustered with genes annotated as distantly related paralogs, 751 such as members of the MORC and NSD gene families. We additionally relied on these trees to 752 more confidently label which genes were ZCWPW1 orthologs and which were ZCWPW2 753 orthologs based on where they clustered in the tree. We considered orthologs as complete if they 754 contain both PWWP and zf-CW domains with e-values < 1. This approach resulted in the 755 identification of 193 complete ZCWPW1 orthologs from 188 species, and 187 complete ZCWPW2 756 orthologs from 180 species.

757 Among the 189 species used in our co-evolutionary test, 164 had complete ZCWPW1 758 orthologs and 154 had complete ZCWPW2 orthologs on the basis of this initial search. For the 25 759 species missing a complete ZCWPW1 ortholog, and for the 35 missing a complete ZCWPW2 760 ortholog, we sought to identify the orthologs from whole genome sequences following the same 761 procedures as described for *PRDM9*. This approach enabled us to identify an additional 3 762 complete ZCWPW1 orthologs from 3 species, and an additional 11 complete ZCWPW2 orthologs 763 from 11 species (Tables S1 and S2). For the remaining species, we checked the putative loss of 764 ZCWPW1 or ZCWPW2 using RNA-seg data when available, which led to the identification of an 765 additional 2 complete ZCWPW1 orthologs, but no additional ZCWPW2 orthologs (Tables S1 and 766 S3). We additionally added one ZCWPW1 ortholog from the common shrew (Sorex araneus) from 767 the Ensemble database, which had been identified previously but was absent from NCBI [26,27]. 768 Lastly, we sought to identify ZCWPW2 from the whole genome sequence of a species of frog with 769 PRDM9 (Ranitomeya imitator) not otherwise included in our co-evolutionary test in order to 770 distinguish whether or not the absence of ZCWPW2 in Xenopus and Dicroglossidae frogs

corresponded to a single loss or multiple events. We were able to identify *ZCWPW2* from this
species, suggesting that *ZCWPW2* has been lost multiple times within frogs (**Tables S1**, **S2** and **S8**).

774

775 iii. TEX15

776 For TEX15, our initial blastp search resulted in the identification of 900 putative orthologs 777 from 363 species. We similarly utilized a tree built using the DUF3715 domain to remove 667 778 genes that cluster with distantly related paralogs, in particular, TASOR and TASOR2 (Figure 779 **S9C**). When making our final calls about *TEX15* orthologs, we labeled them as complete if they 780 contained both DUF3715 and TEX15 domains (accessions pfam12509 and pfam15326). This 781 approach resulted in the identification of 179 complete TEX15 orthologs from 175 species. Among 782 the 189 species used for our co-evolutionary test, 150 had complete TEX15 orthologs on the 783 basis of this initial search. For the 39 species missing a complete TEX15 ortholog, we sought to 784 identify the orthologs from whole genome sequences, following the same procedures as 785 described for PRDM9. In this way, we identified an additional 29 complete TEX15 orthologs from 786 28 species (Table S2). For the remaining species, we checked if we could find TEX15 using RNA-787 seq data, when available, and found one additional complete TEX15 ortholog by this approach 788 (Tables S1 and S3).

789

790 iv. FBXO47

791 For FBXO47, an initial blastp search of the vertebrate RefSeg database using the human 792 sequence as query resulted in the identification of 386 putative FBXO47 orthologs from 380 793 species. We did not perform phylogenetic analysis to support the identification of these orthologs: 794 While we detected a domain (F-BOX) in the human FBX047 gene, due to its high e-value in 795 humans (e-value = 0.01), we did not rely on its presence or absence when inferring the whether 796 or not a complete FBXO47 gene was present in each species. However, each homolog identified 797 in our initial RefSeq analysis was annotated as either FBX047 or FBX047-like with the exception 798 of one CWC25 gene, which was removed. We thus labeled each species as having a complete 799 ortholog if an ortholog was present. This approach resulted in the identification of FBX047 800 ortholog for 181 of the 189 species used for our co-evolutionary test. For the remaining 8 species, 801 we sought to identify FBX047 orthologs from whole genome sequences and/or RNA-seq datasets 802 following the same procedures described for *PRDM9*; however, we were unable to identify any 803 additional FBXO47 in this way (Tables S1 and S3).

804

805 6. Conservation of residues in *ZCWPW*2

806 We carried out a residue conservation analysis using an approach proposed by [69], using 807 code score conservation.py available at https://compbio.cs.princeton.edu/conservation/. This 808 approach quantifies the Jensen-Shannon divergence between the amino acid distribution of the 809 focal residue and a "background amino acid distribution." The alignment of ZCWPW2 was 810 produced using Clustal Omega (using default parameters) within MEGA (version 7, [35,70]). As 811 recommended, the overall background amino acid distribution was drawn based on the 812 BLOSUM62 amino acid substitution matrix provided by the software [69]. Any column of the gene 813 sequence alignment with more than 30% gaps was ignored. A window size of 3 was used to 814 incorporate information from sequential amino acids, as recommended by the default settings.

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7. Evidence for relaxed selective constraint in the absence of *PRDM9*

To test for possible relaxed selection in species without a complete *PRDM9*, we used the program *codeml* within PAML [71,72]. *Codeml* uses protein coding sequences to estimate the ratio of non-synonymous to synonymous substitution rates ($\omega = d_N/d_S$). Values of ω significantly less than 1 are indicative of purifying selection, i.e., of the functional importance of the gene.

821 To this end, we considered each major clade (fish, mammals, reptiles, amphibians) 822 separately and extracted and aligned coding nucleotide sequences from NCBI for multiple 823 species. We aligned those sequences in a codon-aware manner using Clustal Omega (using 824 default parameters) within MEGA (version 7, [35,70]) and inspected the codon-aware alignment 825 visually to ensure that the same isoforms were used across species. For each multi-species 826 alignment, we tried two approaches: (i) We estimated ω under a null model assuming the same 827 ω across all branches and an alternative model in which there are two ω allowed: one ω value in 828 species with a complete *PRDM9* and a second ω for the branches in which *PRDM9* is absent or 829 incomplete (including the internal branches on which PRDM9 may have been lost); (ii) We 830 considered the same null model with the same ω across all branches; and an alternative model 831 with one ω value in species with a complete *PRDM9*, a second ω for the branches in which 832 *PRDM9* is absent or incomplete and additional ω values for each branch on which *PRDM9* was 833 inferred to be lost (a different one for each independent loss, as the ω value averaged over the 834 branch will depend on when along the branch PRDM9 was lost). For (i), significance was 835 assessed using a likelihood ratio test with 1 degree of freedom; for (ii), by the number of degrees 836 of freedom corresponded to the number of distinct ω values minus 1. If ω values were found to 837 be significantly higher in species without a complete *PRDM9*, we tested whether or not we could 838 reject $\omega = 1$ for these species. For two cases in which we could not obtain a multi-species

839	alignment that included the whole coding sequence (ZCWPW1 in fish and TEX15 in amphibians),
840	we instead used the pairwise model (runmode: -2 within PAML) on alignments for a pair of
841	species, and tested whether we could reject $\omega = 1$ for species lacking <i>PRDM9</i> by comparing a
842	model allowing ω to vary versus a null model fixing the ω value at 1, with 1 degree of freedom.
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870 Figures

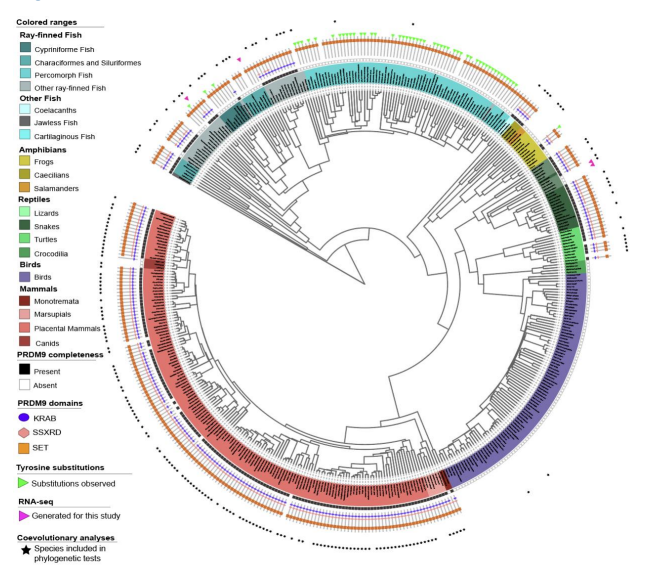
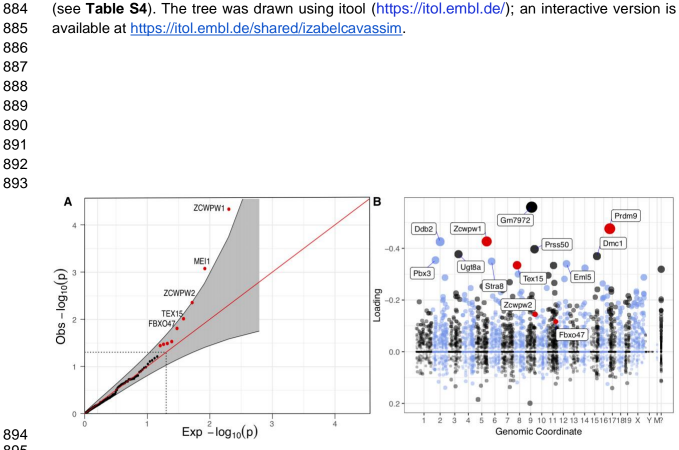




Figure 1. The phylogenetic distribution of PRDM9 and its domain architecture across 872 873 vertebrates. The inferred PRDM9 status of 432 vertebrate species. Branch lengths were 874 computed based on the TimeTree database. For 28 species not present in the database, we used 875 branch length information from a close evolutionary relative; for 14 species in which we made 876 PRDM9 calls, we were unable to find such a substitute, so they are not represented. Different 877 vertebrate clades are indicated by colored segments, with salmon for mammals, cyan for fish, 878 mustard for amphibians, green for reptiles, and purple for birds. In the inner circle, squares 879 indicate whether PRDM9 is complete (filled black) or incomplete/absent (empty black); for species 880 with an uncertain PRDM9 status, no box is shown. The PRDM9 domain architecture of each 881 species is shown with a cartoon, where the presence of a KRAB domain is indicated in blue, of 882 SSXRD in pink, and of the SET domain in orange. Green triangles indicate species that only carry 883 PRDM9 orthologs with substitutions at putatively important catalytic residues in the SET domain

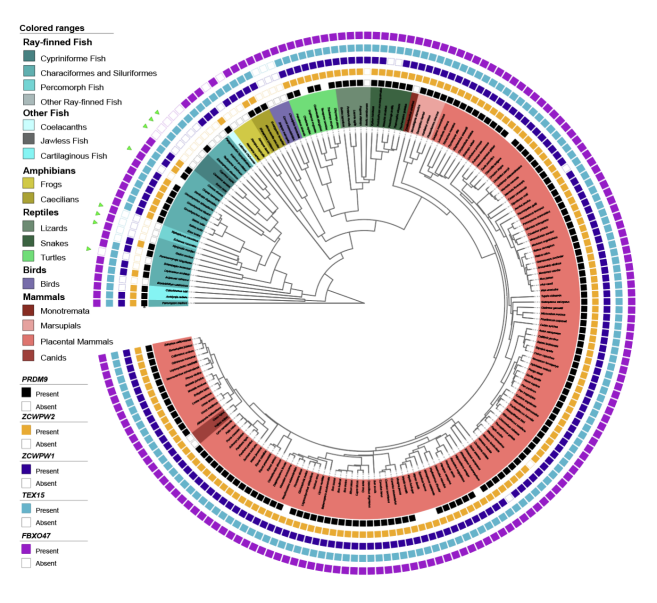


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896 Figure 2. Phylogenetic tests and genes co-expressed with PRDM9 in single cell mouse 897 testes data (A) Quantile-Quantile plot of the p-values obtained from the phylogenetic tests run 898 on 139 genes that appeared to have been lost at least once in the 189 vertebrate species 899 considered. Genes that are significant at the 5% level are shown in red (outside the dashed lines) 900 and a pointwise 95% confidence interval is shown in grey. Genes with a FDR \leq 50% are

901 annotated. (B) Loadings for one of 46 components (component 5) inferred from single cell 902 expression data in mouse testes [31], in which PRDM9 is most highly expressed. The dot sizes 903 are proportional to the square of the absolute value of the loading. PRDM9 and the three genes 904 identified in our phylogenetic tests with p<0.05 are shown in red. Mouse genomic coordinates are 905 displayed. Panel B was made from summary statistics provided by [31], using SDAtools 906 (https://github.com/marchinilab/SDAtools/).

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913 Figure 3. The phylogenetic distribution of *PRDM9* and co-evolving genes across 189 914 species. Filled teal and empty teal squares indicate whether PRDM9 is present or absent, 915 respectively (see Methods). If nothing is indicated, the status of *PRDM9* is uncertain. Likewise, 916 filled orange and empty squares indicate whether ZCWPW2 is present or absent/incomplete; 917 filled and empty navy squares indicate whether ZCWPW1 is present or absent/incomplete; filled 918 and empty light blue squares indicate whether TEX15 is present or absent/incomplete; and filled 919 and empty light purple squares indicate whether FBXO47 is present or absent/incomplete. Green 920 triangles indicate species that only carry *PRDM9* orthologs with substitutions at putatively 921 important catalytic residues in the SET domain (see Table S4). The status of candidate genes 922 (for which FDR \leq 50%; Figure 2A) was re-evaluated based on a search of gene models within 923 whole genome sequences (see Methods); updated p-values for the phylogenetic test are shown

- 924 in Table 1. The tree was drawn using itool (https://itol.embl.de/); an interactive version is available
- 925 at https://itol.embl.de/shared/izabelcavassim.

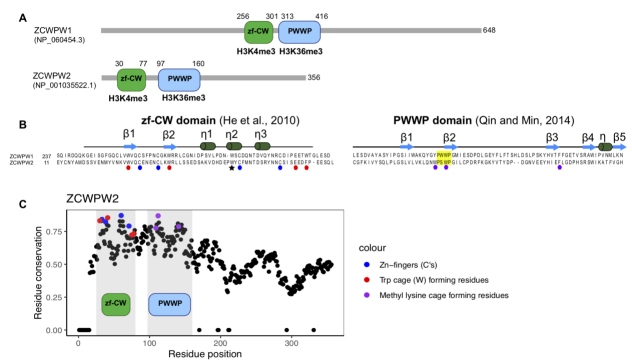


Figure 4. Domain architecture and conservation of ZCWPW1 and ZCWPW2. (A) Amino acid sequence and domain structure composition of genes ZCWPW1 and ZCWPW2 in humans. (B) The ZF-CW domain structure includes the fingers (residues indicated by blue circles) and an aromatic cage (red) expected to bind to H3K4me3 [73], and the star indicates the third Trp residue that is thought to stabilize the fold by hydrophobic interactions [73]. The PWWP domain (yellow) is expected to bind to histone H3K36me3 through a hydrophobic cavity composed of three aromatic residues (purple) [74]. The secondary structures of zf-CW and PWWP domains are represented above sequences. (C) Conservation of residues in ZCWPW2 across vertebrates, with those residues recognizing modifications on the histone tail colored in blue, red and purple. Positions in the ZCWPW2 alignment with > 30% of gaps were ignored and the conservation score was set to 0.

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951 **Tables**

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953 Table 1. Results of phylogenetic tests

We focused on the five genes that had a false discovery rate (FDR) \leq 50%, improved the ortholog status calls, and reran the phylogenetic tests for four of them (all but *MEI1*, which turned out to be present in all species considered; see Methods). Gene source refers to the criterion by which the gene was originally included among our lists of candidates: (1) It is co-expressed with PRDM9 in single cell mouse testes data [31] or (2) variants assigned to the gene are associated with variation in recombination phenotypes in humans [33] or (3) the gene was previously known to have a role in mammalian meiotic recombination from functional studies [32] (see Methods).

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Gene	Position (human coordina te)	Gene source	LogLik H0	LogLik Ha	P-value	FDR	P-value for improved status calls
	chr7: 1						
ZCWPW1	00400826	1	-65.941	-53.35647	4.651e-05	0.0064	1.948e-03
MEI1	chr22: 4 1699503	3,1	-54.200	-44.779	8.442e-04	0.0586	NA
ZCWPW2	chr3: 2834872 1	1	-67.711	-60.146	4.437e-03	0.2055	5.171e-06
TEX15	chr8: 3 0831544	1	-138.430	-131.764	9.760e-03	0.3391	8.682e-02
	chr17: 3893643						
FBXO47	2	2	-53.678	-47.559	1.566e-02	0.4354	1.566e-02

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- 987 <u>https://www.dropbox.com/sh/pihq6a643fz21js/AAANGJWpALT42MCrrsJdlSUHa?dl=0</u>
- 988

989 Author contributions

Conceptualization: M.I.A.C., Z.B., and M.P.; Methodology: M.I.A.C., Z.B., C.H., M.S., and M.P.;
Formal Analysis: M.I.A.C., Z.B.; Investigation: M.I.A.C., Z.B., C.H., M.S.; Resources: M.H.S.,
M.S., M.P.; Data curation: M.I.A.C., Z.B., M.S., and M.P.; Writing - Original Draft: M.I.A.C.
Z.B., M.S., and M.P. Writing - Review and Editing: M.I.A.C., Z.B., C.H., M.S., and M.P;
Visualization: M.I.A.C. and Z.B. Supervision: M.H.S., M.S., and M.P. Project administration:
M.P.; Funding acquisition: M.H.S., M.S., and M.P.

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1001

1002 **Competing interest**

- 1003 The authors declare that they have no competing interests.
- 1004

1005 1006 **References**

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1207 Supplementary tables

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Supplementary Table 1. Genes identified in this study. For *PRDM9* and each candidate gene that was initially identified as significantly coevolving with *PRDM9* (*ZCWPW1*, *MEI1*, *ZCWPW2*, *TEX15*, and *FBX047*), we provide a table detailing, for each ortholog that we identified, which species it is from, how we identified it, its inferred domain architecture, its amino acid sequence, as well as various details about these domains (including their coordinates, e-values, and sequences). For *PRDM9* orthologs, we additionally report which amino acid residues align to three catalytic tyrosine residues in the human SET domain.

1216

Supplementary Table 2. Description of genes found from whole genome sequences. For PRDM9 and each gene initially identified as significantly coevolving with PRDM9 (ZCWPW1, MEI1, ZCWPW2, and TEX15), we provide a table detailing where and how each ortholog identified in our analysis of whole genome assemblies was obtained. FBXO47 is excluded from this table because no FBXO47 orthologs were identified from whole genome sequences.

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Supplementary Table 3. Description of species for which a *de novo* assembly of testis transcriptomes was generated in order to verify the structure and expression of PRDM9 and four significant genes (ZCWPW1, ZCWPW2, TEX15 and FBX047). To this end, we used publicly available RNA-seq data (downloaded from NCBI) [75] and in a subset of cases indicated with a star, generated our own data which are available from the NCBI sequence read archive (Bioproject PRJNA605699, SRA accessions: SRR11050679-SRR11050687, see Methods for further details).

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1231 <u>Supplementary Table 4.</u> The distribution of *PRDM9* orthologs across 446 vertebrate species.
1232 For each species, we describe how many *PRDM9* orthologs we identified of each unique domain
1233 architecture, the domain architecture of the most complete *PRDM9* ortholog from that species,
1234 and whether any *PRDM9* ortholog from that species with the most complete domain architecture

has conserved three catalytic tyrosine residues in the SET domain. We additionally include
columns comparing these results to those previously described in Baker et al. 2017, noting
instances where we have revised our calls of domain architecture.

- 1239 Supplementary Table 5. Description of candidate genes used in the phylogenetic tests. The 241 1240 genes selected for the tests were based on three different sources: (1) genes most highly co-1241 expressed with PRDM9 in mouse testis single cell analyses [31], (2) genes associated with 1242 variation in recombination phenotypes in humans [33], and (3) genes known to have a role in 1243 mammalian meiotic recombination from functional studies (as summarized in the review by [32]). 1244 The genomic coordinates (column named 'Start position') of each gene were based on the 1245 GRCh38/hg38 human reference. The function of each candidate gene is described based on the 1246 definition from its source.
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1248 **Supplementary Table 6.** Presence and absence matrix computed for all candidate genes used 1249 in phylogenetic tests for coevolution with *PRDM9* (139 genes). We defined a gene as complete 1250 ("1") when it contained all the domains observed in four representative vertebrate species with a 1251 complete *PRDM9* sequence, and incomplete ("0") if the gene was not detected in the Refseq 1252 database or if it did not include all the domains shared across four species (see Methods for 1253 details).

1254

1255 Supplementary Table 7. Phylogenetic tests and p-values. P-values were computed by 1256 evaluating the patterns of presence or absence of PRDM9 across 189 vertebrates against the 1257 patterns of presence or absence of candidate genes. Two models were tested using 1258 BayestraitsV3 [36]: a null model in which PRDM9 and a given candidate gene evolve 1259 independently of one another along the phylogeny versus an alternative model in which the gain 1260 ("1") and loss ("0") of the candidate gene is dependent on the status of *PRDM9* and vice versa. 1261 See Pagel, 1994 and the BayesTraitsV3.0.2 manual for further discussion of these models and 1262 rates description.

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1264 Supplementary Table 8. The distribution of PRDM9 and four genes initially found to be significantly coevolving with PRDM9 (ZCWPW1, ZCWPW2, TEX15, and FBXO47) across 189 1265 1266 vertebrate species. MEI1 is not considered because in the curation of the calls, it was found to be 1267 present in all species (see text). (A) Curated calls for the presence or absence of complete genes 1268 based on searches of RefSeq, whole genome assemblies, and RNA-seq data (see Tables S1-1269 **S3**). We additionally include the most complete domain architecture of orthologs from each 1270 species for each gene. (B) Summary of losses inferred for PRDM9, ZCWPW1, ZCWPW2, TEX15 1271 and FBXO47 among the 189 vertebrate species used in our co-evolutionary test.

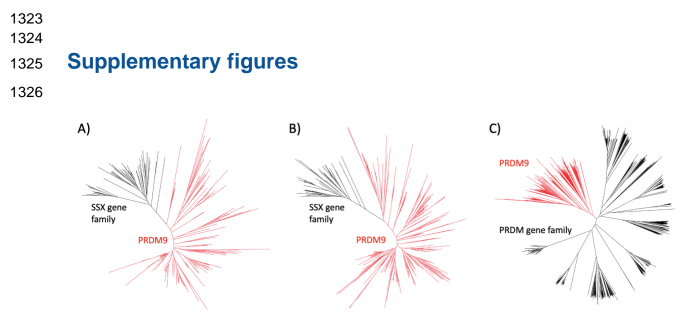
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1273 <u>Supplementary Table 9.</u> (A) Results of phylogenetic tests when considering the pairwise co-1274 evolution of the candidate genes with each other. (B) Results of phylogenetic tests when 1275 considering the SSXRD domain in *PRDM9* classification. P-values were computed by evaluating 1276 the patterns of presence or absence of *PRDM9* across 189 vertebrates against the patterns of 1277 presence or absence of candidate genes. Two models were tested using BayestraitsV3 [36]: a 1278 null model in which *PRDM9* and a given candidate gene evolve independently of one another

along the phylogeny versus an alternative model in which the gain ("1") and loss ("0") of a gene
is dependent on the status of *PRDM9* and vice versa. See [37] and the BayesTraitsV3.0.2 manual
for further discussion of these models and rates description. (**C**) Results of phylogenetic tests
when considering the SSXRD domain in *PRDM9* classification and the curated calls for *ZCWPW1*, *ZCWPW2*, *TEX15*, and *FBXO47*.

Supplementary Table 10. Tests for differences in the rates of amino acid evolution in three significant genes (ZCWPW1, ZCWPW2, TEX15 and FBX047) between representative species with and without a PRDM9 ortholog. To determine whether species lacking a PRDM9 ortholog showed evidence for relaxed selection pressures in co-evolving genes, we estimated ω (dN/dS) using the Branch model within PAML [76] under two models: a null model assuming the same ω across all branches of the phylogeny, and an alternative model in which there are two ω values allowed: one ω value in species lacking a functional PRDM9 and a second ω for the rest of the branches. The clades evaluated in each test are specified. The species used in the alignment for each test are also shown. The log likelihoods for each model, ω estimates and p-values are also provided. See Methods for details.

Supplementary Table 11. Testing the direction of dependency between *PRDM9* and candidate genes ZCWPW1, ZCWPW2, TEX15 and FBX047. Here, we asked whether we could reject a model of independent state transitions of *PRDM9* and a given candidate gene (e.g. ZCWPW1) in favor of a model in which state transitions of the candidate gene depend on those of PRDM9 (model X). Next, we asked whether we could reject the null model in favor of a model in which the state transitions of *PRDM9* depend on those of the candidate gene (model Y). For comparison, we also provide results for the test shown in the main text, in which the alternative considered is that state transitions of PRDM9 depend on those of the candidate gene and vice versa (also shown in Table 1). See Pagel, 1994 and the BayesTraitsV3.0.2 Manual for further description of these models and tests.



1327

Figure S1. Guide trees created from our initial blastp search results for PRDM9 orthologs for (A) KRAB domains, (B) SSXRD domains and (C) SET domains. Genes were removed if they clustered with SSX genes in trees (A) or (B), or if they clustered with PRDM gene family genes other than *PRDM9* or *PRDM7* in the tree (C). Genes clustering with *PRDM9* and retained for subsequent analysis are shown in red.

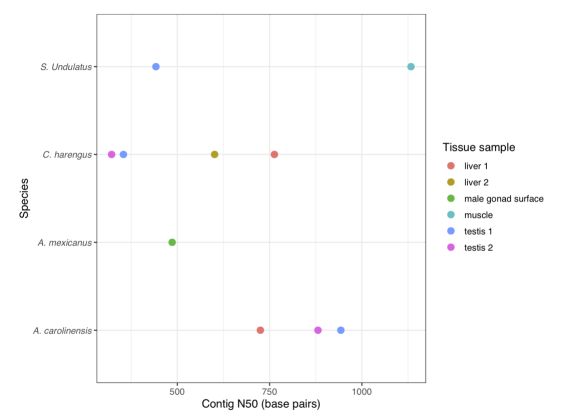
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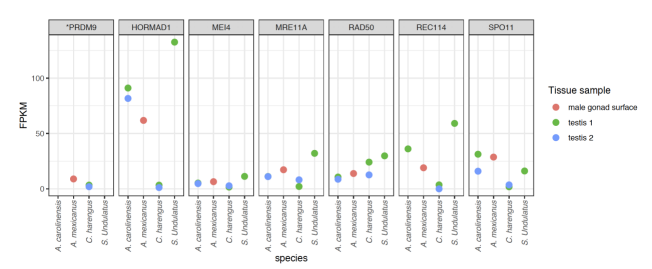


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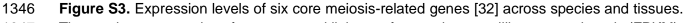
Figure S2. Contig N50 in base pairs as a statistic describing the quality of *de novo* transcriptome
assemblies. Colors represent the different tissues used in the two lizard species (*S. undulatus*and *A. carolinensis*) and two fish species (*C. harengus* and *A. mexicanus*).



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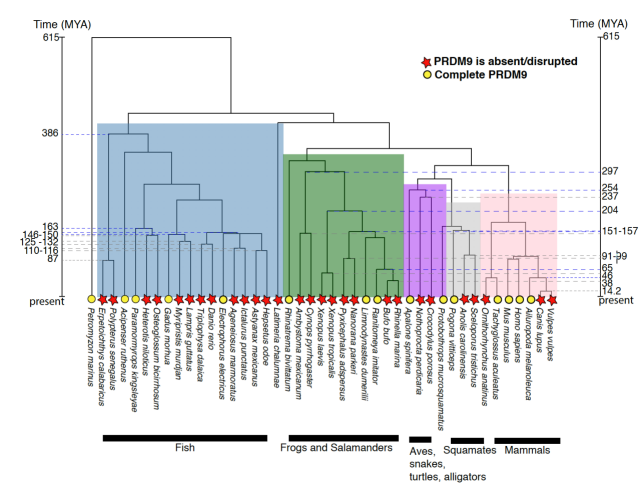


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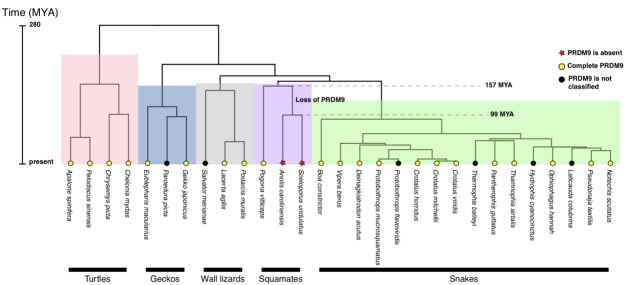
The y-axis corresponds to fragments per kilobase of transcript per million mapped reads (FPKM).
 Despite evidence for expression of the other six core meiotic genes, PRDM9 expression is not

1349 detected in *S. undulatus* and *A. carolinensis*.



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1354 Figure S4. Phylogenetic distribution of *PRDM9* orthologs in vertebrates, using the phylogenetic 1355 tree and divergence dates obtained from Timetree [35]. A complete PRDM9 was found in species 1356 marked with yellow circles. Species marked with a red star are ones for which we were unable to 1357 identify a complete PRDM9. The highlighted dates indicate the inferred timing of the multiple 1358 PRDM9 losses. Dashed lines indicate the earliest (grey) and latest (blue) possible dates for each 1359 loss of PRDM9. The minimum date reflects the time to the most recent common ancestor amongst 1360 species without PRDM9, whereas the 'maximum date' is the time to the first common ancestor 1361 between species without PRDM9 and the most closely related species with PRDM9. The most 1362 recent loss of PRDM9 occurred either in the branch leading to canids, between 14.2 and 46 million 1363 years ago (Mya), or potentially the branch leading to platypus. 1364



1365TurtlesGeckosWall lizardsSquamatesSnakes1366Figure S5. Phylogenetic distribution of *PRDM9* orthologs in reptiles, using the phylogenetic tree1367and divergence dates obtained from Timetree [35]. Species assigned with yellow circles carry a1368complete *PRDM9*. Species indicated with a red star are ones for which we were unable to identify1369PRDM9 expression in testis samples. Species indicated with a black circle are species for which1370Refseq is not available and *PRDM9* classification was therefore not conducted. Based on the1371phylogenetic relationship between *Anolis carolinensis* and *Sceloporus undulatus*, the *PRDM9*1372loss shared by these two species likely occurred between 99 and 157 million years ago (Mya).

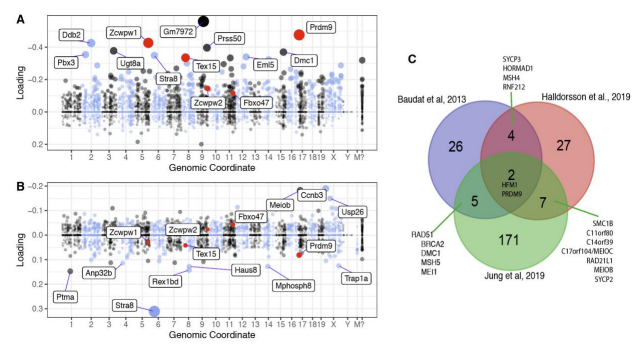




Figure S6. Meiosis-specific candidate genes. [31] inferred 46 principal components from single 1380 1381 cell expression patterns during mouse spermatogenesis, which are thought to loosely correspond 1382 to regulatory programs. Shown in A-B are the two components in which PRDM9 is most highly 1383 expressed. The dot sizes are proportional to the square of the absolute value of the loading, so 1384 are indicative of higher expression within each component. PRDM9 and the five genes with p<0.05 in our phylogenetic analysis are shown in red. Mouse genomic coordinates are displayed. 1385 (A) Component 5 is the one in which PRDM9 has its highest loading; it is associated with double 1386 1387 strand break formation and active during (pre)leptotene [31]. (B) Component 44 is the component in which PRDM9 has its second highest loading; this component is active during zygotene (Jung 1388 1389 et al., 2019). (C) Intersection of candidate genes from three sources: (i) the top 1 percent of 1390 genes with highest loadings in component 5 (ii) genes associated with variation in recombination 1391 phenotypes in humans [33] and (iii) genes known to have a role in mammalian meiotic recombination from functional studies (as summarized in the review by [32]). 1392

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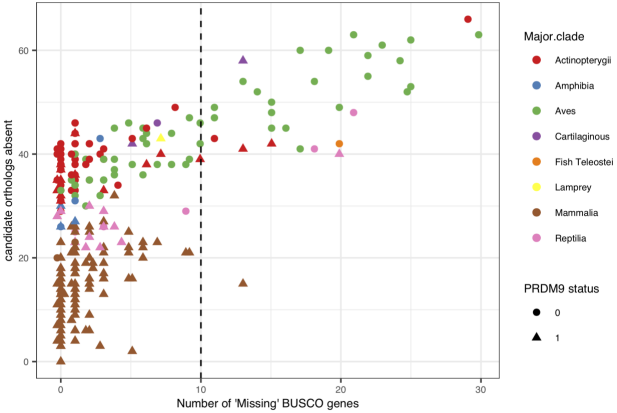




Figure S7. The relationship between the number of candidate genes that were absent in a genome assembly and the number of 'Missing' BUSCO genes [66] for that assembly, across species. BUSCO statistics were computed for the genomes of 339 species. The relationship is significant (Spearman's rank correlation $\rho = 0.5$, p-value < 2.2e-16), suggesting that orthologs of candidate genes of interest might be missed in genomes with low BUSCO scores. In the phylogenetic tests, we therefore considered only species with 10 or fewer missing BUSCO genes (dashed line), leading 32 species to be excluded.

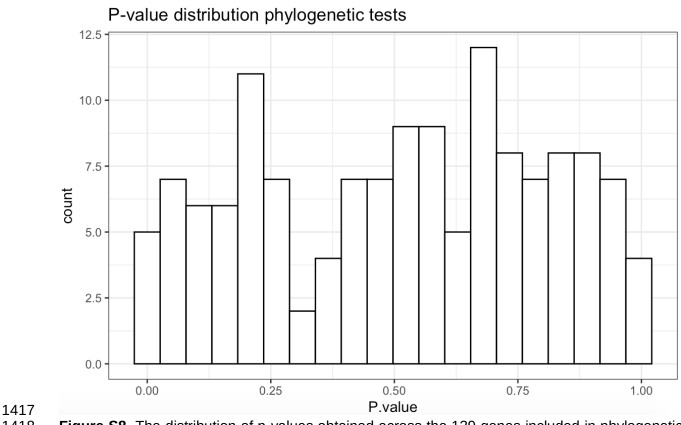


Figure S8. The distribution of p-values obtained across the 139 genes included in phylogenetic
tests (individual p-values are available in Table S7).

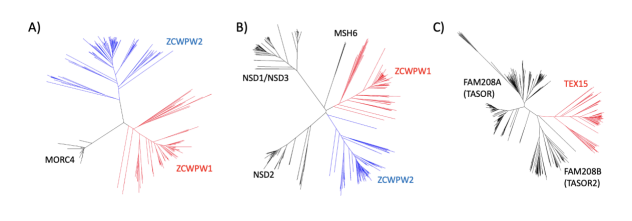


Figure S9. Guide trees created from our initial blastp search results for the zf-CW (A) and PWWP
(B) domains of *ZCWPW1* and *ZCWPW2* orthologs, and the DUF3715 domain of *TEX15*

- orthologs. Genes were removed if they clustered with *MORC4* in tree A, *MSH6*, *NSD1*, *NSD2*, or *NSD3* in tree B, *FAM208A* or *FAM208B* in tree C. Genes clustering with *ZCWPW1*, *ZCWPW2* or *TEX15* and retained for subsequent analysis are shown in red or blue.

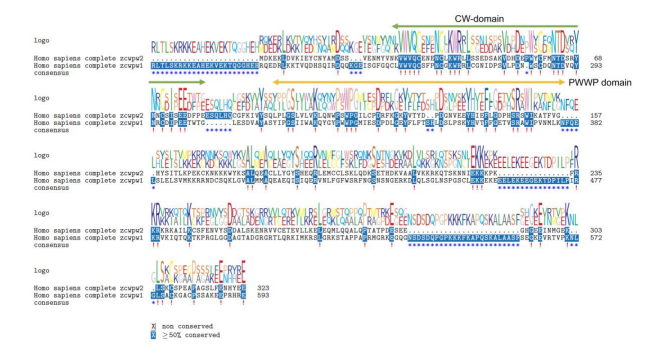


Figure S10: Amino acid sequence alignment between ZCWPW1 and ZCWPW2 proteins from
humans. Superfamily domains are marked. In mice, the CW-domain (green arrow) recognizes
different methylated states of lysine 4 on histone H3 (H3K4me) [77], while the PWWP domain
(yellow arrow) recognizes methylated H3K36 histone tail [74]. The SET domain of PRDM9 trimethylates both histones H3K4 and H3K36 [58].

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