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2	Functional impact of the H2A.Z histone variant during meiosis in
3	Saccharomyces cerevisiae
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43 Abstract

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45 Among the collection of chromatin modifications that influence its function and structure, the 46 substitution of canonical histones by the so-called histone variants is one of the most 47 prominent actions. Since crucial meiotic transactions are modulated by chromatin, here we investigate the functional contribution of the H2A.Z histone variant during both unperturbed 48 49 meiosis and upon challenging conditions where the meiotic recombination checkpoint is 50 triggered in budding yeast by the absence of the synaptonemal complex component Zip1. We 51 have found that H2A.Z localizes to meiotic chromosomes in an SWR1-dependent manner. 52 Although meiotic recombination is not substantially altered, the *htz1* mutant (lacking H2A.Z) 53 shows inefficient meiotic progression, impaired sporulation and reduced spore viability. 54 These phenotypes are likely accounted for by the misregulation of meiotic gene expression 55 landscape observed in *htz1*. In the *zip1* mutant, the absence of H2A.Z results in a tighter 56 meiotic arrest imposed by the meiotic recombination checkpoint. We have found that Mec1-57 dependent Hop1-T318 phosphorylation and the ensuing Mek1 activation are not significantly 58 altered in zip1 htz1; however, downstream checkpoint targets, such as the meiosis I-59 promoting factors Ndt80, Cdc5 and Clb1, are drastically down-regulated. The study of the checkpoint response in *zip1 htz1* has also allowed us to reveal the existence of an additional 60 61 function of the Swe1 kinase, independent of CDK inhibitory phosphorylation, which is relevant to restrain meiotic cell cycle progression. In summary, our study shows that the 62 63 H2A.Z histone variant impacts various aspects of meiotic development adding further insight 64 into the relevance of chromatin dynamics for accurate gametogenesis.

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66 Introduction

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Sexual reproduction relies on a specialized cell division, meiosis, which reduces 68 chromosome ploidy by half and is usually accompanied by cell differentiation processes that 69 70 culminate in the formation of gametes. The reduction in chromosome complement is achieved 71 by two consecutive rounds of nuclear division preceded by a single round of DNA replication. 72 Premeiotic S phase is followed by a long prophase I in which, prior to the first meiotic 73 division, homologous chromosomes (homologs) pair, synapse and recombine. Meiotic 74 recombination is initiated by programmed DNA double-strand breaks (DSBs) generated by 75 the Spoll protein and a cohort of regulatory factors (Keeney et al. 2014). During the repair of 76 a subset of these meiotic DSBs, crossovers between homologs are formed, which are essential 77 for correct distribution of chromosomes to the meiotic progeny. Alignment of homologous 78 chromosomes -pairing- and the stabilization of these interactions by the synaptonemal 79 complex (SC) -synapsis- influence meiotic recombination outcomes (Hunter 2015). These 80 crucial meiotic events are monitored by the so-called meiotic recombination checkpoint 81 (MRC), an evolutionarily-conserved surveillance mechanism that senses defective synapsis 82 and/or recombination and imposes a block or delay in meiotic cell progression providing time to fix the faulty process in order to prevent aberrant chromosome segregation. The meiotic 83 84 checkpoint network also operates in unperturbed meiosis to ensure the proper sequential 85 execution of events (MacQueen and Hochwagen 2011; Subramanian and Hochwagen 2014).

86 In this work, we have used the *zip1* mutant of the budding yeast Saccharomyces 87 cerevisiae as a genetic tool to activate the MRC. Zip1 is a major structural component of the 88 SC central region and ZIP1 deletion impairs synapsis and crossover recombination (Dong and 89 Roeder 2000; Borner et al. 2004; Voelkel-Meiman et al. 2015); as a consequence, the zip1 90 mutant experiences a significant MRC-dependent delay in the prophase to meiosis I transition (Herruzo et al. 2016). The zip1-induced defects are detected by the Mec1-Ddc2^(ATR-ATRIP) 91 92 complex resulting in phosphorylation of the Hop1 checkpoint adaptor at several residues, 93 including T318 (Carballo et al. 2008; Refolio et al. 2011; Penedos et al. 2015). The Hop1 94 protein is a component of the lateral elements of the SC; its abundance, dynamics and 95 phosphorylation state at chromosome axes in response to checkpoint activation are finely 96 tuned by the AAA+ ATPase Pch2 (Herruzo et al. 2016). Phosphorylated Hop1 recruits the 97 meiosis-specific Mek1 protein to chromosomes facilitating the activation of this Rad53/Chk2-98 related kinase containing an FHA domain in two steps: first by Mec1-dependent

99 phosphorylation and subsequently by in trans autophosphorylation of Mek1 dimers on its 100 activation loop (Niu et al. 2005; Ontoso et al. 2013). In turn, active Mek1 stabilizes Hop1-101 T318 phosphorylation at chromosomes (Chuang et al. 2012). Mek1 promotes interhomolog 102 recombination bias by the direct phosphorylation of the recombination mediator Rad54 at 103 T154 to attenuate its interaction with the strand-exchange Rad51 protein (Niu et al. 2009). 104 Also, the phosphorylation of Hed1 at Thr40 stabilizes this protein stimulating its inhibitory 105 action on Rad51 (Callender et al. 2016). Mek1 also exerts a spatial control on recombination 106 bias by a synapsis-dependent mechanism involving Pch2 (Subramanian et al. 2016). In 107 addition, Mek1 is essential for the meiotic checkpoint response to the accumulation of 108 unrepaired DSBs and to the *zip1*-induced synapsis and/or recombination defects (Xu et al. 109 1997; Ontoso et al. 2013; Prugar et al. 2017). The arrest or delay at meiotic prophase I 110 imposed by the MRC is established by two interconnected mechanisms: down-regulation of the Ndt80 transcription factor and inhibitory phosphorylation of Cdc28^{CDK1} (Subramanian and 111 112 Hochwagen 2014). Ndt80 is a master regulator of yeast meiotic development that activates the 113 transcription of a number of genes involved in meiotic divisions and spore formation (Winter 114 2012). Among the gene products regulated by Ndt80, the polo-like kinase Cdc5 and the type-115 B Clb1 cyclin are crucial factors to promote exit from prophase (Tung et al. 2000; Sourirajan 116 and Lichten 2008; Acosta et al. 2011; Argunhan et al. 2017). Inhibition and nuclear exclusion 117 of Ndt80 by the checkpoint prevents the wave of meiotic induction of Clb1 required for entry into meiosis I (Wang et al. 2011). In addition, stabilization of Swe1 by MRC action also 118 maintains Cdc28^{CDK} inhibited by Tyr19 phosphorylation (Leu and Roeder 1999). In sum, the 119 lack of Clb1 induction together with the inhibitory phosphorylation of Cdc28 restrains 120 121 prophase I exit by keeping in check CDK activity levels.

122 Most of DNA meiotic transactions occur in the context of highly specialized 123 chromosome and chromatin structures. Chromatin dynamics can be modulated by several 124 processes, including posttranslational modification (PTM) of histones and incorporation of 125 histone variants. Among the myriad of histone PTMs described to date, a meiotic function has 126 been ascribed to a number of them (Brachet et al. 2012; Wang et al. 2017). In particular, 127 H3K79 methylation and H4K16 acetylation are involved in the budding yeast MRC (Ontoso 128 et al. 2013; Cavero et al. 2016). Much less is known about the meiotic functional contribution 129 of histone variants; in particular, one of the most prominent such as H2A.Z, a variant of the 130 canonical histone H2A conserved in evolution from yeast to human.

131 In vegetative yeast cells, H2A.Z is involved in multiples processes, including 132 transcription regulation (both positively and negatively), maintenance of genome stability and

chromatin silencing (Billon and Cote 2013; Weber and Henikoff 2014). H2A.Z is 133 134 preferentially found in the vicinity of promoters at nucleosomes flanking a nucleosome-135 depleted region containing the transcription start site (Raisner et al. 2005). Nevertheless, not 136 all the functions of H2A.Z are necessarily related to transcription; for example, H2A.Z is also 137 deposited at persistent DSBs promoting their anchorage to the nuclear periphery and 138 stimulating resection (Kalocsay et al. 2009; Adkins et al. 2013; Horigome et al. 2014). The 139 incorporation of H2A.Z to chromatin is carried out by the SWR1 complex, which utilizes the 140 energy of ATP hydrolysis to exchange canonical H2A-H2B by H2A.Z-H2B dimers in 141 particular nucleosomes (Krogan et al. 2003; Kobor et al. 2004; Mizuguchi et al. 2004).

142 The number of studies addressing the role(s) of H2A.Z during meiosis is much scarce, 143 although H2A.Z also appears to perform meiotic functions in several model organisms. In 144 Arabidopsis thaliana, H2A.Z is associated to meiotic recombination hotspots and colocalizes 145 with chromosomal foci of the Dmc1 and Rad51 recombinases; moreover, meiocytes from the 146 *arp6* mutant (lacking a component of the SWR1 complex) show reduced number of Dmc1. 147 Rad51 and Mlh1 foci suggesting a role for H2A.Z in the formation and/or processing of 148 meiotic DSBs (Choi et al. 2013). Meiotic gene expression is also altered in the arp6 mutant of 149 A. thaliana (Qin et al. 2014). During mouse spermatogenesis, H2A.Z is first detected at 150 pachytene, but excluded from the sex-body, where it accumulates at later stages. Based on the 151 dynamics of chromosomal distribution during mammalian spermatogenesis a role for H2A.Z 152 in meiotic sex chromosome inactivation has been proposed (Greaves et al. 2006; Ontoso et al. 153 2014). Recently, a transcription-independent function of H2A.Z in meiotic DSB generation 154 by modulating chromosomal architecture in the fission yeast Schizosaccharomyces pombe has 155 been reported (Yamada et al. 2018).

156 In contrast to most organisms where the absence of H2A.Z is not compatible with life, 157 the *htz1* deletion mutant in S. cerevisiae (lacking H2A.Z) is viable, allowing us to directly 158 assess its meiotic functional impact. In most cases the role of H2A.Z in other organisms has 159 been inferred indirectly by analyzing mutants of the SWR1 complex or by cytological 160 observations. In this work, we demonstrate that H2A.Z is important for meiosis in the 161 budding yeast S. cerevisiae. We show that the htzl mutant displays impaired meiotic 162 progression and sporulation and that spore viability is compromised, although meiotic 163 interhomolog recombination does not appear to be strongly affected. The landscape of gene 164 expression during meiotic prophase is substantially altered in the absence of H2A.Z, likely 165 contributing to at least some of the *htz1* meiotic phenotypes. Finally, we report that H2A.Z 166 also functions during the meiotic checkpoint response induced by the *zip1* mutant impacting

167 on the regulators of the prophase to meiosis I transition, such as the Ndt80 transcription factor

and the CDK inhibitory kinase Swe1. Our study reveals the existence of novel functionalconnections between these cell cycle regulators.

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172 Materials and Methods

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174 Yeast strains

175 Yeast strains genotypes are listed in Table S1. All the strains, except the ones used in 176 Figure 1F and Figure S1, are isogenic to the BR1919 background (Rockmill and Roeder 177 1990). The *htz1::hphMX4*, *swr1::natMX4*, *swr1::hphMX4*, *spo11::natMX4*, *sum1::natMX4*, 178 mer3:hphMX4, swe1::natMX4, [hta1-htb1]::kanMX6 and [hta2-htb2]::natMX4 gene 179 deletions were made using a PCR-based approach (Longtine et al. 1998; Goldstein and 180 McCusker 1999). The *htz1::URA3* deletion was made using the pTK17 plasmid digested with 181 HindIII-SalI (Santisteban et al. 2000). The zip1::LYS2, mek1::kanMX6, ddc2::TRP1, 182 *sml1::kanMX6*, *spo11::ADE2*, *swe1::LEU2* and *rad51::natMX4* gene deletions were 183 previously described (Leu and Roeder 1999; San-Segundo and Roeder 1999; Refolio et al. 184 2011; Ontoso et al. 2013; Herruzo et al. 2016). HTZ1-GFP and MIH1-GFP were made by 185 PCR using pKT127 (Sheff and Thorn 2004) and pFA6a-kanMX6-GFP (Longtine et al. 1998), 186 respectively. The P_{GALI} -ZIP1-GFP and P_{GDP1} -GAL4(848).ER constructs were obtained from 187 Amy Macqueen (Wesleyan University, CT, USA) (Voelkel-Meiman et al. 2012). Strains 188 carrying Swe1 tagged with 3 copies of the MYC epitope at the N terminus and strains 189 carrying ZIP1-GFP have been previously described (Leu and Roeder 1999; White et al. 190 2004). The kinase-dead *swe1-N584A* allele was generated using the *delitto perfetto* approach 191 (Stuckey et al. 2011). Strains carrying the cdc28-AF mutation, in which Thr18 and Tyr19 of 192 Cdc28 have been changed to alanine and phenylalanine, respectively, were generated by 193 transformation with the plasmid pR2042 digested with BlpI (Leu and Roeder 1999). The 194 htb1-Y40F mutant strains in which the Y40 of histone H2B has been mutated to 195 phenylalanine carry the deletion of the HTA1-HTB1 and HTA2-HTB2 genomic loci and a 196 centromeric plasmid (pSS348) expressing HTA1-htb1-Y40F. These strains were generated as 197 follows. A diploid heterozygous for [hta1-htb1]::kanMX6 and [hta2-htb2]::natMX4 198 containing the URA3-based pSS345 plasmid expressing wild-type HTA1-HTB1 was 199 transformed with the TRP1-based pSS347 or pSS348 plasmids expressing wild-type HTA1200 HTB1 (as control) or HTA1-htb1-Y40F, respectively (see plasmid construction below). These 201 diploids were sporulated and Ura- Trp+ haploid segregants harboring [htal-htbl]::kanMX6 202 and [hta2-htb2]::natMX4 genomic deletions and the pSS347 or pSS348 plasmid as the only 203 source for H2A-H2B or H2A-H2BY40F, respectively, were selected. In all cases, gene 204 deletions, mutations and tagging in haploid strains were made by direct transformation with 205 PCR-amplified cassettes and/or digested plasmids, or by genetic crosses and sporulation 206 (always in an isogenic background) followed by selection of the desired segregants. Diploids 207 were made by mating the corresponding haploid parents and isolation of zygotes by 208 micromanipulation.

209

210 Plasmids

211 The plasmids used in this work are listed in Table S2. The 2µ-based high-copy 212 pSS248 plasmid contains the meiosis-specific HOP1 promoter driving the expression of GFP. 213 In-frame cloning of a gene ORF after the GFP in pSS248 leads to overproduction of the GFP-214 fusion specifically during meiotic prophase. pSS248 was constructed as follows. First, the 215 HOP1 promoter (650 bp) was amplified from genomic DNA and cloned into the BglII-PacI 216 sites of pFA6a-kanMX6-GAL1-GFP (Longtine et al. 1998) replacing the GAL1 promoter by 217 the HOP1 promoter to generate pSS232. Then, the P_{HOP1} -GFP fragment from pSS232 was 218 amplified by PCR with oligonucleotides containing the appropriate restriction sites and cloned into SpeI-NotI of pYES2 (Invitrogene) to replace P_{GAL1} by P_{HOP1} -GFP generating 219 220 pSS248. The pSS265 plasmid to overexpress *MIH1* during meiosis was constructed by PCR 221 amplification of the MIH1 ORF flanked by NotI-SpeI sites and cloning into NotI-XbaI of 222 pSS248. For meiotic overexpression of BDF1 the ORF flanked by NotI-SphI sites was 223 amplified by PCR and cloned into the same sites of pSS248 to generate pSS354. Plasmid 224 pSS263 was generated by cloning a 2.7-kb NotI-SalI fragment from pSS200 (=p1-1) (Pak and 225 Segall 2002) containing NDT80 plus the promoter and 3'UTR regions into the same sites of 226 the high-copy vector pRS426. The HTA1-HTB1 genomic region containing the genes 227 encoding histones H2A and H2B expressed from a common divergent promoter including 228 also 285 and 540 bp of the flanking 3'UTR sequences was amplified by PCR from genomic 229 DNA and cloned into the BamHI-SalI sites of the centromeric vectors pRS316 and pRS314 to 230 generate plasmids pSS345 and pSS347, respectively. The htb1-Y40F mutation was introduced 231 by site-directed mutagenesis of pSS347 to generate pSS348. All constructs were verified by 232 sequencing. Oligonucleotide sequences are available upon request.

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234 Meiotic time courses, sporulation efficiency and spore viability

235 For meiotic time courses, BR strains were grown in 3.5 ml of 2xSC medium for 20 -236 24 hours (2% Glucose, 0.7% Yeast Nitrogen Base without amino acids, 0.05% Adenine and 237 Complete Supplement Mixture from Formedium at twice the particular concentration 238 indicated by the manufacturer), then transferred to YPDA (2.5 ml) and incubated to saturation 239 for additional 8 hours. Cells were harvested, washed with 2% potassium acetate (KAc), 240 resuspended into 2% KAc (10 ml) and incubated at 30°C with vigorous shaking to induce 241 meiosis and sporulation. Both YPDA and 2% KAc were supplemented with 20 mM adenine 242 and 10 mM uracil. The culture volumes were scaled-up when needed. To score meiotic 243 nuclear divisions, samples were taken at different time points, fixed in 70% Ethanol, washed 244 in PBS and stained with 1 mg/ml DAPI for 15 min. At least 300 cells were counted at each 245 time point. Meiotic time courses were repeated several times. To induce ZIP1-GFP from the 246 P_{GAL1} promoter in strains expressing GAL4.ER, 1 mM β -estradiol (Sigma E2257; dissolved in ethanol) was added to the cultures. Sporulation efficiency was quantitated by microscopic 247 248 examination of asci formation after 3 days on sporulation plates. Both mature and immature 249 asci were scored. At least 300 cells were counted for every strain. Spore viability was 250 assessed by tetrad dissection. At least 144 spores were scored for every strain.

251

252 Western blotting

Total cell extracts were prepared by TCA precipitation from 5-ml aliquots of sporulation cultures as previously described (Acosta *et al.* 2011). Analysis of Mek1 phosphorylation using Phos-tag gels was performed as reported (Ontoso *et al.* 2013). The antibodies used are listed in Table S3. The ECL or ECL2 reagents (ThermoFisher Scientific) were used for detection. The signal was captured on films and/or with a ChemiDoc XRS system (Bio-Rad).

259

260 Fluorescence microscopy

Immunofluorescence of chromosome spreads was performed essentially as described (Rockmill 2009). For analysis of spindle formation by whole-cell immunofluorecence, the following protocol was used. Cells from meiotic cultures (1.5 ml) were fixed with 3.7% formaldehyde for 45 minutes, washed twice with solution A (1.2 M Sorbitol, 0.05 M KH₂PO₄) and resuspended into the same solution containing 0.1 mg/ml 20T Zymolyase, 0.1% Glusulase and 0.001% β -mercaptoethanol. Samples were incubated at 37°C for 20-30 minutes monitoring spheroplast formation. After two washes with ice-cold Solution A, cells were 268 resuspended into 50 µl of this solution. 25 µl were deposited onto a polylysine-coated 8-well 269 glass slide and let stand for 30 minutes. Liquid was carefully aspirated and the slide was 270 submerged into -20°C methanol for 6 minutes and -20°C acetone for 30 seconds using a 271 Coplin jar. The wells were successively rinsed with 1% BSA in PBS, 1% BSA 0.1% NP-40 in 272 PBS (twice) and 1% BSA in PBS, and incubated overnight with the anti-tubulin antibody in 273 1% BSA-PBS at 4°C. Wells were then rinsed as described above, incubated with the 274 secondary antibody for 2 hours at room temperature and rinsed again. A drop of Vectashield 275 containing DAPI (Vector Laboratories, H-1200) was deposited and extended with a coverslip 276 sealed with nail polish. The antibodies used are listed in Table S3. Images of spreads and 277 fixed whole cells were captured with a Nikon Eclipse 90i fluorescence microscope controlled 278 with MetaMorph software and equipped with a Hammamatsu Orca-AG CCD camera and a 279 PlanApo VC 100x1.4 NA objective. Images of fluorescent spores as well as ZIP1-GFP and 280 HTZ1-GFP live cells were captured with an Olympus IX71 fluorescence microscope 281 equipped with a personal DeltaVision system, a CoolSnap HQ2 (Photometrics) camera and 282 100x UPLSAPO 1.4 NA objective. For Zip1-GFP and Htz1-GFP, stacks of 10 planes at 0.4 283 um intervals were captured. Maximum intensity projections of deconvolved images were 284 generated using the SoftWorRx 5.0 software (Applied Precisions). DAPI images were 285 collected using a Leica DMRXA fluorescence microscope equipped with a Hammamatsu 286 Orca-AG CCD camera and a 63x 1.4 NA objective.

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288 **Recombination frequency**

289 To measure genetic distances in a chromosome VIII interval we used a spore-290 autonomous fluorescence assay in SK1 strains as previously described (Thacker et al. 2011). 291 Basically, diploid SK1 cells were patched on YEP-glycerol plates, streaked on YPD plates 292 and single colonies were inoculated into 2 ml of liquid YPD incubated at 30°C for 20 h. Cells 293 were transferred to 10 ml of YPA (1% yeast extract, 2% peptone, 2% KAc), incubated at 294 30°C for 14 h and sporulated in 10 ml of 2% potassium acetate containing 0.001% 295 polypropylene glycol to prevent aggregation. Asci with fluorescent spores were imaged after 296 48 h in sporulation. Samples were sonicated for 15 sec before imaging. The "cell counter" 297 plugin of ImageJ (http://imagej.nih.gov/ij/plugins/cell-counter.html) was used to manually 298 score the tetrads of each type. Genetic distances (cM) were calculated using the Perkins 299 equation: cM = (100 (6NPD + T))/(2(PD + NPD + T)), where PD is the number of parental ditypes, NPD is the number of nonparental ditypes and T is the number of tetratypes. 300

301

302 Meiotic transcriptome analysis.

303 Global analysis of gene expression during meiotic prophase was carried essentially as 304 described (Morillo-Huesca et al. 2010). Briefly, gene expression profiles were determined 305 using the "GeneChip[™] Yeast Genome 2.0 Array" of Affymetrix at CABIMER Genomics 306 Unit (Seville, Spain). Total RNA from meiotic prophase cells (15 hours after meiotic 307 induction) was isolated using the RNeasy® Midi kit (Oiagen) and its quality confirmed by 308 Bioanalyzer® 2100 (Agilent technology). Synthesis, labeling and hybridization of cDNA was performed with RNA from 3 independent cultures of each strain following Affymetrix 309 310 protocols (http://www.affymetrix.com/ analysis/index.affx). Probe signal intensities were 311 extracted from the scanned images and analyzed with the GeneChip Operating Software 312 1.4.0.036 (Affymetrix). The raw data (CEL files) were preprocessed and normalized using the 313 Robust Multichip Average (RMA) method. Fold-change values (log2) and their FDR-adjusted 314 p-values were calculated with Limma (Linear Models for Microarray Analysis using the 315 affvlmGUI interface. All the statistical analysis was performed using R language and the packages freely available from "Bioconductor Project" (http://www.bioconductor.org). Fold-316 317 change cutoffs were analyzed at 95% confidence levels (FDR-adjusted p-values<0.05). All 318 data is MIAME compliant and the raw data have been deposited at the Miame compliant 319 Gene Expression Omnibus (GEO) database at the National Center for Biotechnology 320 Information (<u>http://www.ncbi.nlm.nih.gov/geo/</u>) and are accessible through accession number 321 GSE110022. Gene ontology and functional clustering analyses were performed using DAVID 322 tools (Database for Annotation, Visualization and Integrated Discovery) (Huang et al. 2007).

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324 Quantitative RNA analysis.

The amount of mRNA was determined by real-time PCR (RT-PCR) amplification of cDNA generated by reverse transcription and RNaseH treatment (SuperScript II Reverse Transcriptase; Invitrogen) of the RNA samples obtained for the microarray hybridization analysis. Amplification of *ACT1* was used to normalize for differences in the amount of input RNA. Similar results were obtained after normalization with *NUP84* (data not shown). Primers were designed using the Primer Express software (Applied Biosystems) and their sequence is available upon request.

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333 Statistics

Unless specified, to determine the statistical significance of differences a two-tailed Student *t*-test was used. *P*-Values were calculated with the GraphPad Prism 5.0 software. The nature of errors bars in graphical representations and the number of biological replicates (n) is
indicated in the corresponding figure legend. For analysis of statistical significance in Venn
diagrams a hypergeometric test was applied.

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340 Data availability

All relevant data necessary for confirming the conclusions presented are within the article and Supplemental Information (GSA Figshare). Strains and plasmids are available upon request. Microarray raw data are deposited at GEO repository under GSE110022 accession number (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE110022).

- 345
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347 **Results**

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349 H2A.Z localizes to meiotic prophase chromosomes in a SWR1-dependent manner

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351 To investigate the localization of H2A.Z during meiotic prophase we generated a 352 functional version of this histone variant tagged with the green fluorescent protein (GFP). Live wild-type cells observed by fluorescence microscopy 15 h after meiotic induction (peak 353 354 of prophase in the BR strain background) displayed the H2A.Z-GFP signal along elongated 355 structures likely corresponding with zygotene-pachytene chromosomes. In contrast, the *swr1* 356 mutant showed diffused H2A.Z-GFP throughout the nucleus (Figure 1A). To explore H2A.Z 357 localization in more detail we performed immunofluorescence of meiotic chromosome 358 spreads (Figure 1B). In wild-type pachytene chromosomes, H2A.Z decorated all chromatin, 359 except a particular region of the genome corresponding to the rDNA region, as demonstrated 360 by the presence of the nucleolar-enriched Pch2 protein (Herruzo et al. 2016) (Figure 1B, 361 arrows). In contrast, pachytene chromosomes of the swrl mutant were largely devoid of 362 chromatin-associated H2A.Z (Figure 1B). These observations indicate that, like in vegetative 363 cells (Krogan et al. 2003; Kobor et al. 2004; Mizuguchi et al. 2004), the SWR1 complex is 364 also required for the deposition of H2A.Z into meiotic chromatin. Occasionally, discrete dots 365 of H2A.Z-GFP accumulation could be observed in *swr1* nuclei. The nature and possible 366 functional implication of this SWR1-independent localization of H2A.Z will be described 367 elsewhere. Western blot analysis revealed that global levels of H2A.Z remained fairly 368 constant throughout the whole meiotic program in the wild type; however, they were 369 gradually diminishing in the *swr1* mutant (Figure 1C), suggesting that chromatin
370 incorporation stabilizes H2A.Z during meiosis.

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372 Meiotic progression and sporulation are impaired in the *htz1* mutant

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374 To determine whether H2A.Z plays a role in meiotic progression, we followed the 375 kinetics of meiotic divisions by DAPI-staining of nuclei in wild type and htz1 strains. 376 Completion of meiotic divisions was less efficient in the *htz1* mutant compared to the wild 377 type (Figure 2A). Likewise, sporulation efficiency and spore viability were also reduced in 378 *htz1*, and asci morphology was altered (Figure 2B-2D). These observations imply that H2A.Z 379 function is required for normal meiotic development. The htzl mutant showed a random 380 pattern of spore death, with no predominance of four-, two- and zero-spore-viable tetrads 381 (Figure 2E), suggesting that the reduced spore viability in *htz1* is not resulting, at least 382 exclusively, from meiosis I non-disjunction events. We also examined crossover 383 recombination in a chromosome VIII interval between CEN8 and THR1 using a microscopic 384 fluorescence assay that is independent of spore viability (Thacker et al. 2011). Recombination 385 frequency in this interval, measured as map distance (cM), was not altered in the htzl mutant 386 compared to the wild type. As a control, a crossover-defective *mer3* mutant was also included 387 in the assay (Figure 2F and Figure S1). To assess whether the inefficient meiotic progression 388 of *htz1* was a consequence of the activation of the meiotic recombination checkpoint we 389 combined the absence of H2A.Z with that of Spo11 (lacking recombination-initiating meiotic 390 DSBs) and with that of Mek1 (lacking the main checkpoint effector kinase). The *htz1* delay in 391 meiotic progression was maintained in the *htz1 spo11* and *htz1 mek1* double mutants (Figures 392 3A and 3B, respectively). Moreover, the dynamics of various indicators of checkpoint 393 activity, such as Hop1-T318 phosphorylation (Herruzo et al. 2016) and Mek1 activation, as 394 assessed both by Mek1 autophosphorylation (Ontoso et al. 2013) and phosphorylation of its 395 H3-T11 target (Cavero et al. 2016; Kniewel et al. 2017), was similar in wild type and htz1 396 (Figure 3C). These results indicate that the lower overall efficiency of meiotic divisions in 397 htzl does not stem from activation of the meiotic recombination checkpoint, and it is 398 consistent with the observation that meiotic recombination (CO) does not appear to be 399 significantly affected in the absence of H2A.Z. To explore the possibility that the absence of 400 H2A.Z affects meiotic entry rather than (or in addition to) meiotic progression we used ZIP1-401 GFP as a reporter for early meiotic gene expression and analyzed the percentage of cells 402 showing nuclear fluorescence in the wild-type and *htz1* strains shortly after meiotic induction.

We found that the kinetics of appearance of Zip1-GFP fluorescence was slightly, but reproducibly, delayed in the *htz1* mutant, although eventually it reached nearly wild-type levels (Figure 3D). This observation likely reflects a delay in the onset of the meiotic program in the absence of H2A.Z and may account, at least in part, for the checkpoint-independent impaired completion of meiotic divisions in the *htz1* mutant.

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409 The SWR1 complex partially impairs meiosis in the absence of H2A.Z

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411 The SWR1 complex is required to replace H2A-H2B by H2A.Z-H2B dimers at 412 particular nucleosomes. It has been proposed that the SWR1 complex exerts a deleterious 413 effect on chromatin integrity in the *htz1* mutant due to the attempt to replace the canonical 414 histone H2A by H2A.Z in the absence of this histone variant creating nucleosome instability 415 (Morillo-Huesca et al. 2010). As a consequence, deletion of SWR1 (encoding the catalytic 416 component of the SWR1 complex) totally or partially suppresses some of the multiple 417 phenotypes of *htz1* in vegetative cells (Morillo-Huesca *et al.* 2010). Thus, we analyzed the 418 kinetics of meiotic divisions, sporulation efficiency and spore viability in *swr1* and *htz1 swr1* 419 mutants. We found that meiotic progression was faster (Figure S2A) and that asci formation 420 and spore viability were somehow improved in *htz1 swr1* compared to *htz1* (Figure S2B and 421 S2C), although they did not reach wild-type levels. The *swr1* single mutant showed an 422 intermediate phenotype between the wild type and *htz1* mutant in meiotic progression and 423 sporulation efficiency (Figure S2A and S2B). These observations imply that some, but not all, 424 meiotic phenotypes of *htz1* result from the pathogenic action of SWR1 in the absence of 425 H2A.Z. Moreover, the fact that SWR1 deletion only partially suppresses the meiotic defects of 426 htzl also supports a direct impact of H2A.Z chromatin deposition on proper meiotic 427 development.

428

429 Meiotic gene expression is altered in the *htz1* mutant

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431 Several studies have demonstrated that mutation of *HTZ1* causes transcriptional 432 misregulation during vegetative growth (Billon and Cote 2013). To assess the influence of 433 H2A.Z on general meiotic gene expression we used whole-genome microarray analysis to 434 compare the transcription profile of wild-type and *htz1* meiotic prophase cells (15 hours after 435 meiotic induction). We found 611 genes showing differential expression in *htz1* compared to 436 wild type (1.5-fold cutoff, p<0.05) (Table S4); of those, 339 genes were up-regulated and 272

down-regulated. Among the genes whose expression was increased in the absence of H2A.Z, 437 438 genes encoding ribosomal proteins were on the top of the list ordered by linear-fold change 439 (LFC) (Table S4). On the top positions of the genes whose expression was significantly 440 down-regulated in the *htz1* mutant we found genes involved in the MEN (Mitotic Exit 441 Network) pathway (BFA1, LTE1) and PP1 phosphatase regulators (GIP14, GAC1) (Table S4). 442 Although there were no meiosis-specific genes among those whose mRNA levels showed a 443 strong change, it was possible to find some genes with meiotic functions, chromatin, DNA 444 damage response and cell-cycle related events with a LFC>1.5 (Table 1). The reduced 445 expression of some of these genes in the *htz1* mutant was verified by RT-PCR analysis of the 446 same mRNA samples used in the microarrays (Figure S3A). Moreover, gene ontology and 447 clustering analyses of the genes with decreased expression showed a significant enrichment of 448 functional categories related to both mitotic and meiotic cell cycle regulation (Table S4). On 449 the contrary, genes with increased expression in *htz1*, cluster mainly in ribosome biogenesis, 450 translation and metabolic processes (Table S4). Since genes encoding ribosomal proteins are 451 rapidly repressed upon meiotic induction (Chu et al. 1998), this observation is consistent with 452 the slight delay in meiosis entry of the *htz1* mutant (Figure 3D). Interestingly, 133 of 611 genes ($p=5x10^{-5}$) with differential level of expression between wild type and *htz1* during 453 454 meiotic prophase identified in this study overlap with those affected by htz1 (948 genes) in 455 mitotically growing cells (Morillo-Huesca et al. 2010) (Figure S3B).

Thus, these analyses revealed that the meiotic transcriptional landscape is significantly disturbed in *htz1*, suggesting that the pleiotropic phenotypes of the *htz1* mutant (aberrant morphology, inefficient meiotic development, low spore viability...) could stem from the more or less subtle alteration of multiple mechanisms.

460

461 The *zip1 htz1* mutant displays a tight checkpoint-dependent meiotic arrest

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463 Next, we sought to explore the possible role of H2A.Z during challenged meiosis; that 464 is, under conditions in which meiotic defects trigger the meiotic recombination checkpoint. 465 We used the *zip1* mutant, which is defective in CO recombination and SC formation, to 466 induce the checkpoint. The *zip1* mutant arrests in prophase I for a long period, but eventually, 467 at late time points, a fraction of the culture completes the meiotic divisions to generate largely 468 inviable spores (Figure 4A) (Ontoso et al. 2013). Strikingly, we found that meiotic 469 progression was completely blocked in the *zip1 htz1* double mutant as most cells remained 470 uninucleated during the whole time course (Figure 4A). This observation suggests that H2A.Z 471 may have a role during prophase I exit because its absence, combined with that of Zip1,472 provokes a strong meiotic arrest.

473 Like in the wild-type (Figure 1), chromatin incorporation and stability of H2A.Z also 474 depended on SWR1 in the *zip1* mutant (Figure S4A-S4B). To determine whether the impact 475 of *htz1* on the inability to resume meiotic progression in *zip1* was a consequence of the 476 deleterious effect of SWR1 as explained above, we analyzed the kinetics of meiotic divisions 477 in *zip1 swr1* and *zip1 htz1 swr1* mutants. Interestingly, like *zip1 htz1*, the *zip1 swr1* and *zip1* 478 htz1 swr1 mutants also showed a tight meiotic block (Figure S4C). Since the swr1 single 479 mutant is able to complete meiosis, albeit with a small delay compared to the wild type, these 480 results indicate that the strong meiotic arrest of *zip1 htz1*, *zip1 swr1* and *zip1 htz1 swr1* stems 481 from the lack of H2A.Z chromatin deposition and does not result from the indirect toxic effect 482 of SWR1 in the absence of H2A.Z.

483 To ascertain whether the *zip1 htz1* block was caused by the meiotic recombination 484 checkpoint we generated the *zip1 htz1 spo11* mutant, in which meiotic DSBs are not formed 485 (Keeney et al. 1997), and the zip1 htz1 ddc2 mutant, in which meiotic recombination intermediates are not sensed (Refolio et al. 2011). We found that meiotic divisions and 486 487 sporulation were largely restored in the *zip1 htz1 spo11* and *zip1 htz1 ddc2* mutants (Figure 488 4B) generating mostly dead spores (5.6% and 1.5% spore viability for *zip1 htz1 spo11* and 489 *zip1 htz1 ddc2*, respectively; n=72), thus confirming that the meiotic prophase block in *zip1* 490 *htz1* is imposed by the meiotic recombination checkpoint.

491

492 The *zip1 htz1* mutant does not accumulate additional unrepaired DSBs

493

494 One possible explanation for the more robust meiotic arrest of *zip1 htz1* compared to 495 that of *zip1* is that the absence of H2A.Z may provoke additional defects that, combined to 496 those resulting from the lack of Zip1, could lead to further hyperactivation of the meiotic 497 recombination checkpoint and, therefore, a tighter prophase I block. To test this possibility, 498 we used immunofluorescence of spread nuclei to analyze the presence of Rad51 foci as an 499 indirect marker for unrepaired DSBs (Joshi et al. 2015) in zip1 and zip1 htz1 mutants. The 500 zip1 htz1 spo11 mutant was also included as a control for the absence of meiotic DSBs. Due 501 to the different kinetics of meiotic progression of the strains analyzed (Figure 4A-4B), only 502 prophase I nuclei, as assessed by the bushy morphology of tubulin staining, were scored 503 (Figure 4C). We found that the *zip1 htz1* double mutant did not display more Rad51 foci than 504 *zip1* (Figure 4C), suggesting that the absence of H2A.Z together with that of Zip1 does not

505 generate more unrepaired meiotic DSBs. We also performed immunofluorescence of spread 506 nuclei using an antibody that recognizes phosphorylated S/T-Q motifs as an additional assay 507 for Mec1/Tel1-dependent DNA damage signaling during meiotic prophase. We found that 508 phospho-S/T-Q foci were significantly increased in a *dmc1* mutant, used as control, that 509 accumulates hyper-resected DSBs (Bishop et al. 1992), but similarly decorated prophase 510 chromosomes of *zip1* and *zip1 htz1* (Figure S5). These observations do not favor the 511 possibility that the accumulation of additional DNA damage is responsible for the 512 exacerbated meiotic arrest of *zip1 htz1*.

513

514 Dynamics of upstream checkpoint activation-deactivation is normal in *zip1 htz1*

515

516 To pinpoint what event in the *zip1*-induced MRC pathway is impacted by H2A.Z we 517 used a battery of molecular markers to analyze checkpoint status during meiotic time courses 518 of wild type, *zip1* and *zip1 htz1* strains (Figure 4D). Activation of the Mec1-Ddc2 sensor 519 complex by unrepaired DSBs (and perhaps other types of meiotic defects) is one of the first 520 events in the meiotic checkpoint pathway (Refolio et al. 2011; Subramanian and Hochwagen 521 2014). Active Mec1 phosphorylates Hop1 at various sites, including T318 (Carballo et al. 522 2008; Penedos et al. 2015). In the zip1-induced checkpoint, Hop1-T318 phosphorylation is 523 critical to sustain activation of the Mek1 effector kinase (Herruzo et al. 2016), and serves as 524 an excellent readout for Mec1 activity. Since unrepaired DSBs promote Mec1 activation, 525 Hop1 phosphorylation has been also used as an indirect assay for DSB formation (Chen et al. 526 2015). In the wild type, there was a weak and transient phosphorylation of Hop1-T318 527 coincident with the peak of prophase I and ongoing recombination. In contrast, Hop1-T318 528 phosphorylation was very robust and sustained in the *zip1* mutant (Figure 4D), although at 529 late time points phospho-Hop1-T318 declined coincident with completion of meiotic 530 divisions in a fraction of the culture (Figure 4A). Remarkably, despite the tight meiotic arrest 531 (Figure 4A), the kinetics of Hop1-T318 phosphorylation in the *zip1 htz1* double mutant was 532 similar to that of *zip1* (Figure 4D), further supporting that the turnover of meiotic DSBs is not 533 significantly affected by htz1.

We also monitored the activity of the downstream Mek1 effector kinase using three different readouts: Mek1 autophosphorylation (Ontoso *et al.* 2013), Hed1 phosphorylation at T40 (Callender *et al.* 2016) and histone H3 phosphorylation at T11 (Cavero *et al.* 2016). As shown in Figure 4D, the dynamics of Mek1 activation paralleled that of Hop1-T318 phosphorylation (that is, Mec1 activity) and, again, was similar in both *zip1* and *zip1 htz1*,
except for a slight persistence of phospho-H3-T11 in *zip1 htz1* at the latest time point.

540 These results, together with the analysis of Rad51 foci, indicate that the robust meiotic 541 block in *zip1 htz1* does not arise from the persistence of unrepaired recombination 542 intermediates sustaining permanent upstream checkpoint activation.

543

544 H2A.Z is required for reactivation of the cell cycle checkpoint targets

545

546 We next analyzed the downstream targets that are inhibited by the checkpoint to 547 prevent cell cycle progression while recombination and/or synapsis defects persist. In 548 particular, we examined the production of various meiosis I-promoting factors: the Ndt80 549 transcriptional inductor, the Clb1 cyclin and the Cdc5 polo-like kinase (Acosta et al. 2011). In 550 addition, we also monitored the levels of the Swe1 kinase and its activity: the inhibitory 551 phosphorylation of Cdc28 (CDK) at tyrosine 19 (Leu and Roeder 1999). In the wild type, 552 after the recombination process is completed and the transient activation of Mek1 disappears. 553 the program for meiosis I entry is turned on with the production of Ndt80, Clb1 and Cdc5, as 554 well as the reduction of the inhibitory phosphorylation at Y19 of Cdc28 (Figure 4D). In the 555 *zip1* mutant, the induction of Ndt80, Clb1 and Cdc5 were significantly delayed and high 556 levels of the Swe1 kinase promoting Cdc28-Y19 phosphorylation persisted for as long as 557 Mek1 was active. However, as Mek1 activation eventually declined, Ndt80 and Cdc5 were 558 induced, and Swe1 and phospho-Cdc28-Y19 diminished, thus sustaining entry into meiosis I 559 of at least a fraction of the cells (Figure 4A, 4D). In contrast, we found that although Mek1 560 was down-regulated in *zip1 htz1* with similar kinetics to that in *zip1*, Ndt80, Clb1 and Cdc5 561 production remained largely inhibited, and Swe1 and phospho-Cdc28-Y19 levels stayed high 562 at late time points (Figure 4D), consistent with the inability of *zip1 htz1* cells to exit prophase 563 I (Figure 4A). These results indicate that the main cell cycle targets of the checkpoint are 564 misregulated in the absence of H2A.Z and suggest that this impairment is responsible for the 565 strong block in meiotic progression of *zip1 htz1*.

566

567 HA2.Z contribution to checkpoint recovery

568

569 To determine whether H2A.Z is required to re-start meiotic cell cycle progression 570 when the *zip1* defects that initially triggered the checkpoint are corrected we used a 571 conditional system in which *ZIP1-GFP* expression is controlled by β -estradiol. *ZIP1-GFP*

572 was placed under control of the GAL1 promoter in strains producing a version of the Gal4 573 transcriptional regulator fused the β -estradiol receptor (Gal4[848].ER) (Benjamin *et al.* 2003; 574 Voelkel-Meiman et al. 2012). As depicted in Figure 5A, meiotic cultures of both wild-type 575 and *htz1* strains were initiated without β -estradiol; that is, in the absence of Zip1, to induce 576 the checkpoint response. After 24 h, when the cells are blocked in prophase by the 577 checkpoint, β -estradiol was added to half of the culture and the other half was maintained in 578 the absence of the hormone as control. Recovery from the arrest after ZIP1 induction was 579 monitored at the cytological level (Zip1-GFP chromosome incorporation and DAPI staining 580 of nuclei) and at the molecular level (western blot analysis of various checkpoint markers) 581 (Figure 5B-5D).

582 In the absence of β -estradiol ("ZIP1 OFF"), the checkpoint was activated in the wild 583 type as shown by the prominent H3-T11 and Hed1-T40 phosphorylation, but eventually the 584 phosphorylation of these markers decreased concomitant with Ndt80 activation, Cdc5 production and Cdc28-Y19 dephosphorylation (Figure 5D); thus sustaining meiotic 585 586 progression (Figure 5C). Note that, for unknown reasons, the meiotic delay induced by the 587 checkpoint in this "ZIP1 OFF" situation is less pronounced than in a zip1 mutant (Figure 4A), 588 perhaps due to a leaky, but undetectable, expression of *GAL1-ZIP1* even in the absence of β -589 estradiol. In the *htz1* mutant without β -estradiol, the checkpoint was also heavily activated but, albeit with a slightly slower kinetics, the levels of H3-T11 and Hed1-T40 590 phosphorylation were also finally reduced. However, like in *zip1 htz1* mutants. Ndt80 591 592 production was not induced and Cdc28-Y19 remained phosphorylated at late points (Figure 593 5D); as a consequence, meiotic progression was robustly blocked (Figure 5C). Thus, this 594 "ZIP1 OFF" situation phenocopies ZIP1 deletion in *htz1* (Figure 4A-4D).

595 When β -estradiol was added, ZIP1-GFP expression was induced, and 3 h after 596 hormone addition Zip1-containing chromosomes were detected in nuclei of both wild type 597 and htz1 (Figure 5B). ZIP1-GFP induction was slightly less efficient in the htz1 mutant 598 (Figure 5D), perhaps due to the effect of H2A.Z on GAL1 promoter regulation (Santisteban et 599 al. 2000). In the wild type, the checkpoint was rapidly turned off upon Zip1 production: Mek1 600 signaling drastically disappeared, Ndt80 and Cdc5 were sharply induced and Cdc28-Y19 601 phoshosphorylation was erased (Figure 5D; "ZIP1 ON"). Consistently, prophase-arrested 602 wild-type cells immediately underwent meiotic divisions after ZIP1 expression (Figure 5C; 603 "ZIP1 ON"). In the htz1 mutant the checkpoint was also down-regulated upon ZIP1 induction, 604 but with a slower kinetics than that of the wild type. Consistently, a fraction of *htz1* cells 605 resumed meiotic divisions (Figure 5C, "*ZIP1* ON"); thus, H2A.Z is not essential to re-start 606 meiotic cell cycle progression when the defects that triggered the checkpoint are resolved, but 607 contributes to an efficient recovery from the cell-cycle arrest.

608

609 NDT80 overexpression partially alleviates *zip1 htz1* meiotic arrest

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611 Since *zip1 htz1* shows a dramatic reduction in Ndt80 levels, and Cdc5 production is also impaired (Figure 4D), we examined whether an artificial increase in CDC5 and NDT80 612 613 expression could restore meiotic progression in *zip1 htz1*. As reported (Acosta *et al.* 2011), 614 *CDC5* overexpression from a high-copy plasmid partially suppressed the meiotic delay of the 615 zip1 single mutant (Figure 6A); however, it had little effect on zip1 htz1 (Figure 6B). In 616 contrast, NDT80 overexpression did promote more efficient meiotic progression in both zip1 617 and *zip1 htz1* (Figure 6A-6B). These observations indicate that, in part, the strong meiotic 618 block of the zip1 htz1 mutant results from the drastic reduction in Ndt80 production and 619 suggest that the relevant Ndt80-dependent event responsible for the arrest is not the inability 620 to efficiently activate CDC5 expression.

621

622 Deletion of SWE1, but not mutation of Cdc28-Y19, suppresses the *zip1 htz1* meiotic 623 block

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625 We have found that the levels of both the Swe1 kinase and the phosphorylation of its 626 target, Cdc28-Y19, remain high at late time points in the *zip1 htz1* meiotic cultures. To assess 627 the relevance of Cdc28-Y19 inhibitory phosphorylation to impose the tight *zip1 htz1* meiotic 628 arrest (Figure 7A), we generated three situations in which this phosphorylation event is either 629 abolished or drastically reduced (Figure 7B): 1) SWE1 deletion, 2) cdc28-AF mutation 630 (carrying the threonine 18 and tyrosine 19 of Cdc28 changed to alanine and phenylalanine, 631 respectively), and 3) overexpression of the *MIH1* gene from the prophase I-specific HOP1 632 promoter in a high-copy plasmid (Figure S6A).

Remarkably, deletion of *SWE1* conferred a notable suppression of the *zip1 htz1* meiotic arrest (Figure 7C) although it did not reach wild-type kinetics; however, the elimination of Cdc28-Y19 phosphorylation by other means, such as Mih1 overproduction or *cdc28-AF* mutation had none or only a subtle effect on meiotic progression as most cells remained uninucleated (Figure 7C); only about 10% of *zip1 htz1 cdc28-AF* cells segregated their nuclei. In contrast, *MIH1* overexpression or *cdc28-AF* mutation did accelerate meiotic 639 progression in a *zip1* single mutant (Figure S6B). A kinase-dead *swe1-N584A* allele (Harvey 640 *et al.* 2005) conferred the same suppression of the checkpoint meiotic arrest as the *SWE1* 641 deletion both in *zip1* and *zip1 htz1* strains (Figure S6C), ruling out the possibility of a direct 642 inhibitory effect exerted by the physical interaction of Swe1 with CDK independent of Tyr19 643 phosphorylation. Thus, these results strongly suggest that the Swe1 kinase must impact an 644 additional mechanism, independent of CDK phosphorylation, which is particularly relevant in 645 the absence of H2A.Z to maintain the *zip1*-induced checkpoint arrest.

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CLB1 overexpression restores meiotic progression in zip1 htz1 cdc28-AF

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To further explore the checkpoint response in *zip1 htz1* and the effect of CDK phosphorylation we analyzed by western blot various molecular markers in the *swe1* and *cdc28-AF* mutants. According with its meiotic progression (Figure 7C), the checkpoint was deactivated in *zip1 htz1 swe1*, as manifested by the disappearance of phospho-Hop1-T318 and phospho-Hed1-T40. Concurrently, the meiosis I-promoting factors Ndt80, Clb1 and Cdc5 were produced, albeit with slower kinetics than in the wild type (Figure 7D).

Like in *zip1 htz1*, upstream checkpoint signals were also down-regulated in *zip1 htz1 cdc28-AF*; in contrast, Ndt80, Clb1 and Cdc5 accumulated at higher levels at later time points in this mutant (Figure 7D and Figure S6D). The presence of meiosis I-promoting factors suggests that the *zip1 htz1 cdc28-AF* triple mutant is proficient to undergo the prophase to meiosis I transition, but does not efficiently complete chromosome segregation. Indeed, about 40% of *zip1 htz1 cdc28-AF* cells assembled meiotic spindles at late time points (Figure 7E) despite their marked impairment to undergo meiotic divisions (Figure 7C).

Notably, *CLB1* overexpression from a high-copy plasmid restored substantial meiotic progression in *zip1 htz1 cdc28-AF* phenocopying *zip1 htz1 swe1* (Figure 7C-7D). In sum, these observations suggest that, in addition to phosphorylate Cdc28 at tyrosine 19 to prevent exit from prophase I, Swe1 regulates timing and/or abundance of Clb1 production to restrain meiotic progression in *zip1 htz1* at a later stage in meiotic development.

667

668 **Discussion**

669

670 The H2A.Z histone variant is a ubiquitous determinant of chromatin structure playing 671 crucial roles in genome stability and gene expression in mitotically dividing eukaryotic cells. bioRxiv preprint doi: https://doi.org/10.1101/316133; this version posted May 8, 2018. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

However, only a limited number of studies in a few model organisms have addressed the relevance of H2A.Z in meiosis, often using indirect approaches. In this article, we have focused on the direct functional contribution of H2A.Z during meiosis in the budding yeast *S*. *cerevisiae*, a widely used model system for meiotic studies.

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677 H2A.Z is required for proper meiotic development

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679 We report here that the htzl mutant of S. cerevisiae lacking the H2A.Z histone 680 completes the meiotic program albeit less efficiently than the wild type. The htz1 mutant 681 shows delayed entry into meiosis, impaired sporulation and reduced spore viability indicating 682 that H2A.Z is required to sustain accurate meiosis. The persistence of recombination 683 intermediates or incomplete synapsis triggers the so-called pachytene checkpoint or meiotic 684 recombination checkpoint (MRC) that delays meiotic progression. We found that checkpoint 685 elimination by deleting *MEK1* or abolishing DSB formation by deleting *SPO11* do not restore 686 normal levels of meiotic nuclear divisions in htz1 indicating that the faulty events resulting in 687 impaired completion of meiotic development are not sensed by the MRC and likely do not 688 involve recombination.

689 In fission yeast, H2A.Z participates in the initiation of meiotic recombination by 690 promoting the association of Spo11 and accessory proteins to chromatin (Yamada et al. 691 2018). We have found a modest reduction in the number of Rad51 foci in *zip1 htz1* compared 692 to *zip1* (Figure 4C) that could be compatible with reduced number of initiating DSBs, 693 although a slightly defective loading of Rad51 to DSBs in the absence of H2A.Z or a delayed 694 onset of DSB formation cannot be ruled out. A possible role for H2A.Z in DSB generation 695 could be also inferred from the presence of H2A.Z at promoters (at least in vegetative cells) 696 (Raisner et al. 2005) where most DSBs occur in S. cerevisiae (Pan et al. 2011). However, our 697 results suggest that, in budding yeast, the functional contribution of H2A.Z to DSB formation, 698 if any, is only minor: 1) dynamics of Hop1 phosphorylation at T318, which serves as an 699 indirect reporter for meiotic DSBs, is similar in wild type and htzl. 2) A reduction in DSB 700 formation provoked by the absence of H2A.Z would result in a less stringent checkpoint 701 response; however the *zip1 htz1* double mutant displays a more robust checkpoint arrest 702 compared to *zip1*. 3) Crossover recombination in a particular interval of chromosome VIII is not significantly affected by htz1. It is formally possible that recombination could be altered 703 704 in other chromosomal regions and/or that CO homeostasis could compensate for a reduced 705 number of initiating events (Martini et al. 2006), but this would imply at best a subsidiary

706 function for H2A.Z in DSB formation. In sum, we do not favor the scenario in which the 707 meiotic phenotypes of the *htz1* mutant could be solely explained by impaired DSB formation. 708 Our genome-wide study of meiotic gene expression in the *htz1* mutant reveals that many 709 down-regulated genes cluster in several functional categories related to mitotic and meiotic 710 cell cycle and chromosome segregation events (Table 1 and Table S4). We propose that, in 711 unperturbed conditions, H2A.Z is not essential to perform any critical meiotic event, but the 712 massive transcription misregulation that occurs in the absence of this histone variant may 713 impact on various processes resulting in a less accurate and efficient completion of the 714 meiotic program.

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716

H2A.Z is essential to resume meiotic progression in the absence of Zip1

717

718 Certain chromatin modifications are crucial for checkpoint activity. Dot1-mediated 719 trimethylation of H3K79 controls Pch2 chromosomal distribution and sustains Hop1 720 phosphorylation and the ensuing Mek1 activation in *zip1* mutants. As a consequence, deletion 721 of DOT1 or mutation of H3K79 suppresses the meiotic arrest/delay of zip1 (San-Segundo and 722 Roeder 2000; Ontoso et al. 2013). The Sir2 histone deacetylase is also essential for the zip1-723 induced MRC. One of the main targets of Sir2 is acetylated H4K16. In zip1 sir2 mutants, as 724 well as in *zip1 H4-K16Q* mutants (mimicking constitutive H4K16 acetylation), the *zip1* block 725 is bypassed (San-Segundo and Roeder 1999; Cavero et al. 2016). At least in vegetative cells, 726 Dot1 and the SIR complex collaborate with H2A.Z in delimiting the boundaries between 727 euchromatin and telomeric heterochromatin (Dhillon and Kamakaka 2000; Meneghini et al. 728 2003). However, these chromatin modifications perform opposite functions in the MRC; 729 while in *zip1 dot1* and *zip1 sir2* the meiotic delay is suppressed, *zip1 htz1* shows a stronger 730 meiotic arrest. Our results imply that, in contrast to Dot1 and Sir2, H2A.Z is not required for 731 checkpoint activation, but it is involved in regulation meiotic progression at least in a zip1 732 mutant.

733 We show that the *zip1* mutant exhibits a pronounced meiotic delay, but eventually, 734 checkpoint signaling declines, as manifested by the drop in Hop1 phosphorylation and in 735 Mek1 activation at late time points, and at least a fraction of the culture resumes meiotic 736 progression and completes sporulation. In principle, checkpoint deactivation and resumption 737 of cell cycle progression can occur by two related but conceptually different phenomena: 'checkpoint adaptation' and 'checkpoint recovery'. Adaptation takes place when, despite the 738 739 persistence of the defects that initially triggered the checkpoint, its activity declines after a prolonged period and the cell cycle resumes without previous elimination of the damage. This process of adaptation has been extensively documented in vegetative budding yeast responding to the presence of an irreparable DSB (Pellicioli *et al.* 2001). In contrast, checkpoint recovery involves the disappearance or repair of the initial problems that stimulated the checkpoint, resulting in decreased signaling and cell cycle progression.

745 Previous studies suggest that the eventual checkpoint deactivation and recovery of 746 meiotic progression in *zip1* is consequence of the disappearance of the initial defects (likely 747 unrepaired DSBs) presumably by using the sister chromatid instead of the homolog as 748 template for DNA repair. This is based on the observation that deletion of RAD51, which 749 fundamentally compromises sister chromatid recombination (Liu et al. 2014; Callender et al. 750 2016), leads to a permanent arrest in *zip1* (Herruzo *et al.* 2016) (Figure S7A). In this work we 751 report that, like *zip1 rad51*, the *zip1 htz1* double mutant also shows a tight meiotic block; 752 however, the analysis of various checkpoint markers reveals that the cause of the arrest is 753 different in *zip1 rad51* and *zip1 htz1*. In the *zip1 rad51* mutant, high levels of Hop1-T318 754 phosphorylation and Mek1 activity persist until late time points, consistent with the 755 accumulation of unrepaired recombination intermediates that signal to the checkpoint. 756 Consequently, Cdc28-Ty19 phosphorylation remains high and Cdc5 production is inhibited, 757 thus explaining the meiotic arrest (Herruzo et al. 2016) (Figure S7B). In contrast, we show 758 here that in *zip1 htz1*, Hop1 and Mek1 activation eventually decline with similar kinetics to 759 that observed in the *zip1* single mutant, although meiosis I promoting factors (i.e., Ndt80, 760 Cdc28, Cdc5, Clb1) remain largely inhibited. These observations imply that the 761 disappearance of the initial signal stimulating the checkpoint is not impacted by *htz1*, placing 762 H2A.Z function downstream in the pathway.

763

764 Influence of H2A.Z on Ndt80 and CDK activity

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766 In our molecular analysis of the *zip1*-induced MRC pathway at various levels, the 767 main alterations detected resulting from the absence of H2A.Z were the dramatic reduction in 768 Ndt80 levels and the persistence of both the Swe1 kinase and phosphorylation of its substrate 769 Cdc18-Y19. The observation that *NDT80* overexpression partially suppresses the *zip1 htz1* 770 arrest raises the possibility that H2A.Z could be directly or indirectly controlling NDT80 gene 771 expression. It has been recently described that Bdfl, a subunit of the SWR1 complex involved 772 in the interaction with certain histone marks at particular nucleosomes (Altaf *et al.* 2010), is 773 required for meiotic progression and sporulation. Bdf1 binds to the NDT80 promoter through 774 the BD1 and BD2 bromodomains promoting its transcription (Garcia-Oliver et al. 2017). 775 Nevertheless, several observations suggest that H2A.Z does not control Ndt80 levels via 776 Bdf1. The interaction of Bdf1 with the *NDT80* promoter is independent of the SWR1 complex 777 (Garcia-Oliver et al. 2017), consistent with our observation that meiotic progression is not 778 significantly affected in the swr1 single mutant (Figure S2A-S2B). However, the meiotic 779 checkpoint function of H2A.Z does depend on SWR1 since both *zip1 htz1* and *zip1 swr1* 780 show meiotic arrest (Figure S4D). In addition, strong BDF1 overexpression does not promote 781 sporulation in *zip1 htz1* (Figure S8A). Moreover, we did not find a significant change in 782 *NDT80* transcript levels in our genome-wide expression analysis of the *htz1* mutant during 783 meiosis. Regulation of NDT80 expression is quite complex and also involves the elimination 784 of the Sum1 repressor binding to the middle-sporulation elements (MSE) in its promoter. The 785 displacement of Sum1 from the MSE requires the competition with Ndt80 and also the 786 phosphorylation of Sum1 by Ime2 and CDK (Winter 2012). We found that, like *zip1 htz1*, the 787 *zip1 htz1 sum1* triple mutant remains blocked in meiosis (Figure S8B) indicating that H2A.Z 788 does not exert its effect on Ndt80 levels via Sum1. In addition, activation of Ndt80 requires 789 its phosphorylation in the nucleus; stimulation of the MRC results in cytoplasmic 790 sequestration of Ndt80 (Wang et al. 2011). It is tempting to speculate that H2A.Z could be 791 involved, directly or indirectly, in the nuclear import of Ndt80 when the signal stimulating the 792 checkpoint by the absence of Zip1 declines. The contribution of H2A.Z to the nuclear 793 transport of other proteins has been reported in yeast (Gardner et al. 2011), but the almost 794 undetectable levels of Ndt80 in *zip1 htz1* complicate this analysis with the tools currently 795 available.

796 Our results also show that, in *zip1 htz1*, Swe1-dependent inhibitory phosphorylation of 797 Cdc28-Y19 persists longer than in *zip1*, suggesting that H2A.Z action may be impinging on 798 CDK activity. In fact, deletion of SWE1, which abolishes Cdc29-Y19 phosphorylation, 799 significantly suppresses *zip1 htz1* arrest. Since *MIH1*, the gene encoding the phosphatase that 800 reverts Cdc28-Y19 phosphorylation, was found among the genes whose meiotic expression 801 decreases in the *htz1* mutant (Table 1), it is plausible to postulate that lower levels of the 802 Mih1 phosphatase in *zip1 htz1* could explain the accumulation of phosphorylated Cdc28-Y19 803 and the impaired meiotic progression. However, we demonstrate that strong overproduction 804 of Mih1, which results in negligible Cdc28-Y19 levels, does not restore meiotic nuclear 805 divisions in *zip1 htz1*. This observation, together with the fact that a non-phosphorylatable 806 cdc28-AF mutant also has a minimal impact on the kinetics of meiotic progression of zip1

htz1, strongly suggest that Swe1 must possess another target in addition to CDK to restrain
meiosis in *zip1 htz1*.

809 Besides CDK, only a limited number of substrates for Swe1/Wee1 have been 810 described. One attractive candidate is Y40 of histone H2B, which is phosphorylated by Swel 811 in yeast (or H2B-Y37 phosphorylated by Wee1 in mammals) to control transcription of 812 histone genes (Mahajan et al. 2012). H2A.Z interacts with H2B in the nucleosomes; therefore, 813 it is formally possible that the conformational change induced by SWR1-dependent substitution of histone H2A by H2A.Z could modulate the phosphorylation of H2B-Y40 by 814 815 Swe1. To explore if this chromatin modification has an impact on the MRC, we have 816 generated and analyzed a non-phosphorylatable *htb1-Y40F* mutant and found that the *zip1* 817 htz1 htb2A htb1-Y40F mutant displays the same meiotic arrest as zip1 htz1 (Figure S8C), indicating that this additional Swe1 target is not relevant for the checkpoint response. 818

819 It is surprising that in the *zip1 htz1 cdc28-AF* mutant we observe the induction and 820 accumulation of the proteins involved in meiosis I entry, such as Ndt80, Clb1 and Cdc5, but 821 most cells remain uninucleated (Figure 7). This situation (i.e., accumulation of Ndt80, Cdc5 822 and Clb1) is reminiscent of the metaphase I arrest induced by a meiotic-depletion P_{CLB2} -cdc20 823 mutant (Okaz et al. 2012) and suggest that, at least some, zip1 htz1 cdc28-AF cells are 824 capable of exiting prophase and may arrest at a later stage, such as the metaphase to anaphase 825 I transition. Remarkably, CLB1 overexpression in zip1 htz1 cdc28-AF allows completion of 826 meiotic divisions to a similar degree as does the *zip1 htz1 swe1* mutant. This observation is 827 consistent with the notion that, in the absence of CDK inhibitory phosphorylation (i.e., cdc28-828 AF), Swe1 negatively controls CLB1 levels in zip1 htz1 likely by inhibiting a CLB1-829 promoting factor. We note that overexpression of *CLB1* from a high-copy plasmid not only 830 increases the global amount of Clb1, but also accelerates its production being detected at 831 earlier time points in the meiotic kinetics. Execution of proper prophase to meiosis I transition 832 is under tight temporal control by a number of events including the sequential degradation 833 and accumulation of mitotic and meiotic factors, respectively (Okaz et al. 2012). We show 834 that CLB1 overexpression in zip1 htz1 cdc28-AF partially restores the proper scenario for 835 timely execution of meiotic transitions. Clb1 is phosphorylated in a CDK- and Cdc5-836 dependent manner and it is imported to the nucleus by a mechanism that depends on CDK, 837 but not Cdc5 activity. Although Clb1 nuclear localization is not essential for meiotic nuclear 838 divisions it contributes to efficient meiosis I exit (Tibbles et al. 2013). On the other hand, the 839 biological relevance of Clb1 phosphorylation remains to be established, but it correlates with 840 the induction of Cdc5. What is the identity of the CLB1-promoting factor negatively

841 controlled by Swe1? We speculate that Swe1 could be acting, directly or indirectly, on Ndt80 842 to inhibit its activity especially in the absence of H2A.Z. We propose a model in which Swel 843 action could impact both CDK and Ndt80 activity to restrain meiotic progression (Figure 8A). 844 A cross-talk between CDK and Ndt80 activation in checkpoint-inducing conditions has been 845 also documented (Acosta et al. 2011). This model would explain the following situations. 1) 846 In the *zip1 htz1* mutant overexpressing NDT80, exogenous levels of this transcription factor 847 could partially overcome Swe1 inhibitory effect on Ndt80, resulting only in a partial release 848 of the meiotic arrest (Figure 6) because Swe1-dependent Cdc28-Y19 phosphorylation would 849 persist (Figure 8B). 2) In the *zip1 htz1 cdc28-AF*, the inhibition of CDK by Swe1 is released 850 because the phosphorylation target is mutated, but the timing of Clb1 induction is incorrect 851 due the opposite effect of CDK and Swe1 on Ndt80 preventing proper meiotic progression 852 (Figure 8C). 3) In the *zip1 htz1 swe1*, both inhibitions on CDK and Ndt80 disappear 853 sustaining meiotic progression (Figure 8D).

854 In summary, the detailed analysis of the MRC in the *zip1 htz1* allowed us to discover 855 novel functional interactions between the downstream components of the pathway driving 856 meiotic cell cycle progression. Why these aspects are particularly manifested in the absence 857 of H2A.Z? We show here that a number of genes involved in different cell cycle events are 858 misregulated in the *htz1* mutants. A feasible explanation is that the unbalanced levels in cell 859 cycle regulators creates more stringent conditions in *zip1 htz1* for meiosis I entry in 860 comparison with *zip1*, thus revealing more subtle aspects of the molecular mechanisms 861 regulating the prophase to meiosis I transition when the MRC has been deactivated. 862 Additional work will be required to pinpoint the relevant factors targeted by H2A.Z.

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1092 Figure Legends

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1095 Figure 1. Localization of H2A.Z during meiotic prophase depends on SWR1. (A) 1096 Representative images of wild type and *swr1* live cells, at 15 hours after meiotic induction 1097 (peak of prophase I), expressing HTZ1-GFP. (B) Immunofluorescence of spread meiotic 1098 chromosomes from wild type and swrl stained with DAPI (red) to visualize chromatin, anti-1099 GFP (green) to detect H2A.Z, and anti-Pch2 (blue) to mark the nucleolar region (arrow). (C) 1100 Western blot analysis of H2A.Z production during meiosis detected with anti-GFP antibodies. 1101 Tubulin was used as a loading control. Strains are DP840 (HTZ1-GFP) and DP841 (swr1 1102 HTZ1-GFP).

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1104 Figure 2. H2A.Z is required for proper meiotic development. (A) Time course analysis of 1105 meiotic nuclear divisions; the percentage of cells containing two or more nuclei is 1106 represented. Error bars: SD; n=6. (B) Sporulation efficiency, determined by microscopic 1107 counting, as the percentage of cells forming mature or immature asci after 3 days on 1108 sporulation plates. Error bars: SD; n=3. (C) Representative DIC images of asci. (D) Spore viability determined by tetrad dissection. At least 288 spores were scored for each strain. 1109 1110 Error bars: SD; n=5. (E) Distribution of tetrad types. The percentage of tetrads with 4, 3, 2, 1 1111 and 0 viable spores (4-sv, 3-sv, 2-sv, 1-sv and 0-sv, respectively) is represented. Error bars: 1112 SD; n=3. (F) Recombination frequency, expressed as map distance (cM), in a chromosome VIII interval (see Figure S1). Error bars: range; n=2. Strains used in (A)-(E) are DP421 (wild 1113 1114 type) and DP630 (htz1). Strains used in (F) are DP969 (wild type), DP973 (htz1) and DP974 1115 (*mer3*).

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1117 Figure 3. The inefficient meiotic progression of the *htz1* single mutant does not result 1118 from activation of the meiotic recombination checkpoint. (A, B) Time course analysis of 1119 meiotic nuclear divisions; the percentage of cells containing two or more nuclei is 1120 represented. Error bars: SD; n=3. (C) Western blot analysis of Hop1-T318 phosphorylation 1121 and Mek1 activity at the indicated time points in meiosis. PGK was used as a loading control. 1122 Strains in (A-C) are DP421 (wild type), DP630 (htz1), DP713 (mek1), DP1523 (spo11), 1123 DP1144 (htz1 spo11) and DP1259 (htz1 mek1). (D) Time course analysis of ZIP1-GFP 1124 induction. The percentage of cells showing Zip1-GFP nuclear fluorescence during early time

points after transfer to sporulation conditions is represented. Strains are: DP437 (wild type)
and DP838 (*htz1*). Error bars: SD; n=3.

1127

Figure 4. Robust checkpoint-dependent meiotic arrest in zip1 htz1. (A), (B) Time course 1128 1129 analysis of meiotic nuclear divisions; the percentage of cells containing two or more nuclei is 1130 represented. Error bars: SD; n=3. Strains are DP421 (wild type), DP422 (zip1), DP776 (zip1 1131 htz1), DP1524 (zip1 spo11), DP815 (zip1 htz1 spo11) and DP816 (zip1 htz1 ddc2). (C) Localization and quantification of Rad51 foci as markers for unrepaired DSBs on spread 1132 1133 meiotic nuclei of zip1 (DP449), zip1 htz1 (DP776) and zip1 htz1 spo11 (DP815) after 16 h of meiotic induction. Only prophase I nuclei, as assessed by tubulin staining, were scored. 1134 1135 Representative images are shown. (D) Western blot analysis of the indicated molecular 1136 markers of checkpoint activity at different levels in the pathway. PGK was used as a loading 1137 control. Strains are DP421 (wild type), DP422 (zip1) and DP631 (zip1 htz1). For detection of 1138 Myc-tagged Swe1, the strains used are DP1353 (wild type), DP1354 (zip1) and DP1414 (zip1 1139 htz1).

1140

1141 Figure 5. Analysis of meiotic checkpoint recovery. (A) Schematic representation of the 1142 experimental setup for conditional ZIP1 induction in wild type (DP1185) and htz1 (DP1186) cells containing the GAL4-ER transcriptional activator regulated by β -estradiol and P_{GAL1}-1143 1144 ZIP1-GFP. (B) Representative fluorescence microscopy images showing SC incorporation of 1145 Zip1-GFP. Cells were imaged 3 hours after β -estradiol addition. (C) Time course analysis of 1146 meiotic nuclear divisions; the percentage of cells containing two or more nuclei is 1147 represented. The arrow indicates β -estradiol addition (blue lines and symbols). Error bars: 1148 range; n=2. (D) Western blot analysis of the indicated molecular markers of checkpoint 1149 activity. PGK was used as a loading control.

1150

Figure 6. *NDT80* overexpression partially suppresses *zip1 htz1* meiotic arrest. (A), (B) Time course analysis of meiotic nuclear divisions; the percentage of cells containing two or more nuclei is represented. Strains are DP422 (*zip1*) in (A) and DP1017 (*zip1 htz1*) in (B), transformed with vector alone (pRS426) or with high-copy plasmids overexpressing *CDC5* (pJC29) or *NDT80* (pSS263), denoted as *OE-CDC5* and *OE-NDT80*, respectively. Error bars: SD; n=3.

- 1157
- 1158

1159 Figure 7. Impact of Cdc28-Y19 phosphorylation and Clb1 levels on *zip1 htz1* meiotic 1160 arrest. (A) Schematic representation of the regulation of CDK activity by Cdc28-Y19 phosphorylation controlled by the opposite action of the Swe1 kinase and the Mih1 1161 phosphatase. (B) Western blot analysis of Cdc28-Y19 phosphorylation in the indicated 1162 1163 strains. Total Cdc28 is also shown as control. (C) Time course analysis of meiotic nuclear 1164 divisions; the percentage of cells containing two or more nuclei is represented. Error bars: 1165 SD; n=3. (D) Western blot analysis of the indicated molecular markers of checkpoint activity. Swe1 was detected with anti-myc antibodies. PGK was used as a loading control. Strains in 1166 1167 (B-D) are: DP1353 (wild type), DP1414 (zip1 htz1), DP1113 (zip1 htz1 swe1) and DP1416 1168 (*zip1 htz1 cdc28-AF*). To overexpress *MIH1* and *CLB1*, the DP1414 and DP1416 strains were 1169 transformed with high-copy plasmids pSS265 (OE-MIH1) and pR2045 (OE-CLB1), 1170 respectively. (E) Whole-cell immunofluorescence using anti-tubulin antibodies in *zip1 htz1* 1171 (DP1017) and zip1 htz1 cdc28-AF (DP1154) cells at 48 hours in meiosis. Representative 1172 nuclei of prophase, meiosis I and meiosis II stages are shown. The quantification is presented 1173 in the graph. 169 and 119 nuclei were scored for zip1 htz1 and zip1 htz1 cdc28-AF, 1174 respectively. Error bars: range; n=2.

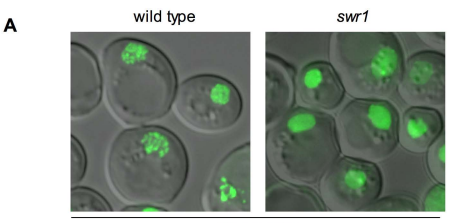
1175

1176 Figure 8. Exit from prophase I in S. cerevisiae. (A) A model for the regulation of the prophase to meiosis I transition by the meiotic recombination checkpoint. See text for details. 1177 1178 The discontinuous line connecting Mek1 and Swe1 indicates that there is no evidence for direct phosphorylation of Swe1 by Mek1. A functional connection or dependency between 1179 1180 DSB repair by sister chromatid recombination and entry into meiosis I is represented by 1181 dotted lines. (B), (C) and (D) Impact on meiotic progression resulting from the mutant 1182 conditions indicated. Green and red colors represent the predominant positive and negative 1183 effects, respectively.

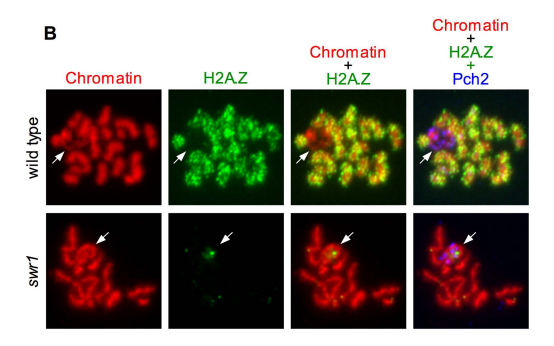
5	
GENE	LFC (>1.5)
Cell cycle	
BFA1	2.393400
LTE1	2.277104
GIP4	2.268020
BUB2	1.704342
MCK1	1.586536
MIH1	1.583281
CDC7	1.536321
Meiotic genes	
RME1	1.852768
RPD3	1.799528
HFM1	1.683362
REC8	1.666158
MEK1*	1.599045
MRE11	1.597116
SKI8	1.570898
IME4	1.531359
ZIP2	1.511252
SPO22	1.504426
HOP1	1.499766
DNA damage response	
SRS2	1.898810
RAD17	1.705357
TOF1	1.612276
MEC1	1.509767
IRC6	1.505446
Chromatin	
SWC3	2.864716
SWI3	2.070301
RSC8	1.850621
SPT20	1.631780
HFI1	1.629896
CHD1	1.514238
LEC: linear fold change (>1.5)	

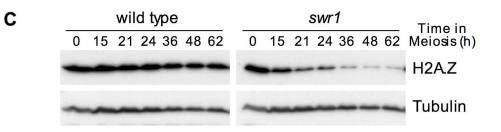
Table 1. Subset of genes with decreased meiotic prophase expression in htz1 (p<0.05)

LFC: linear fold change (>1.5) *p=0.07



H2A.Z-GFP





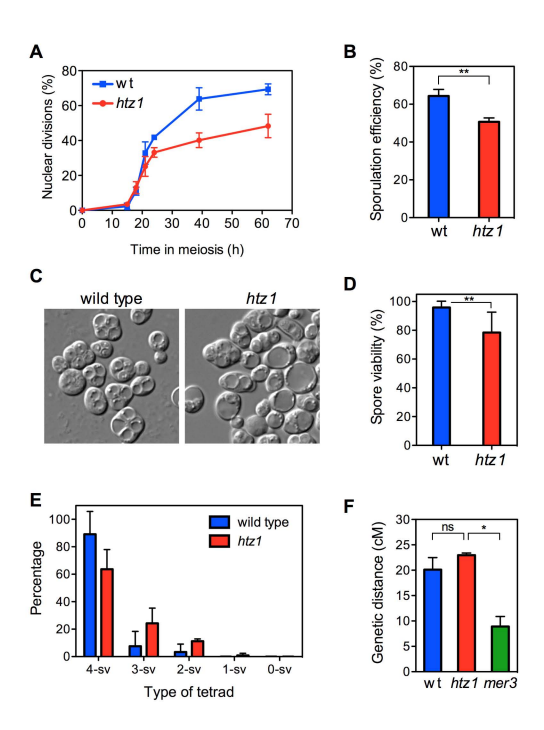
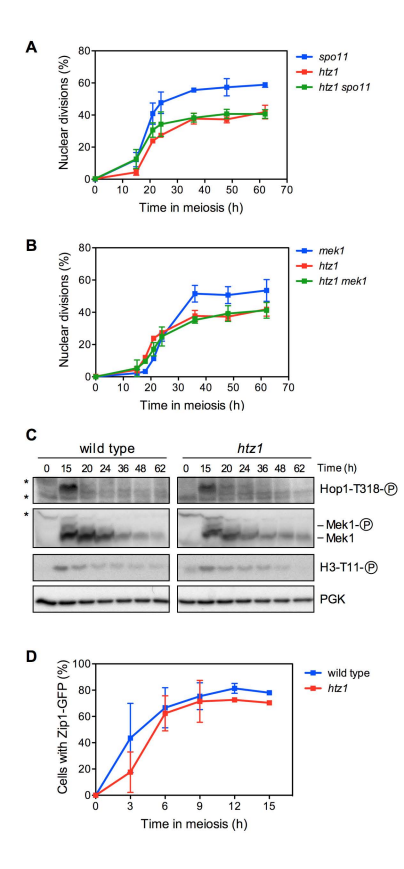


Figure 2



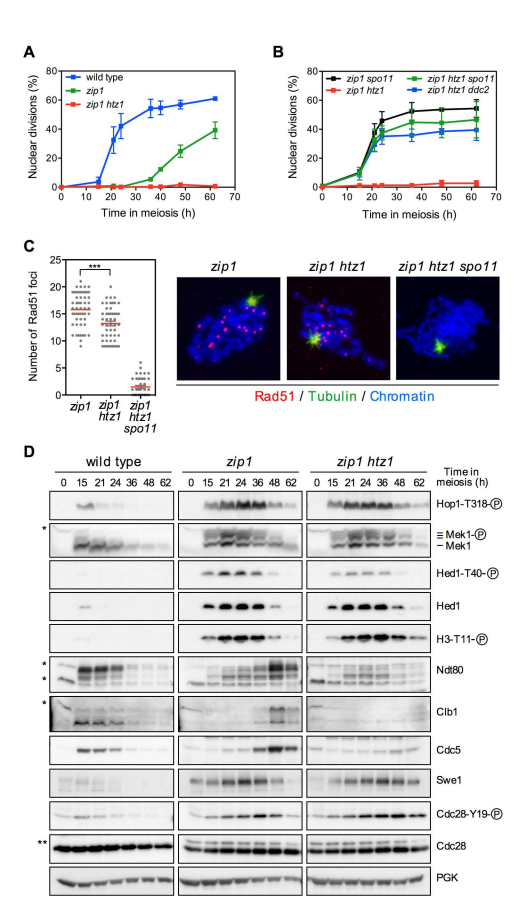
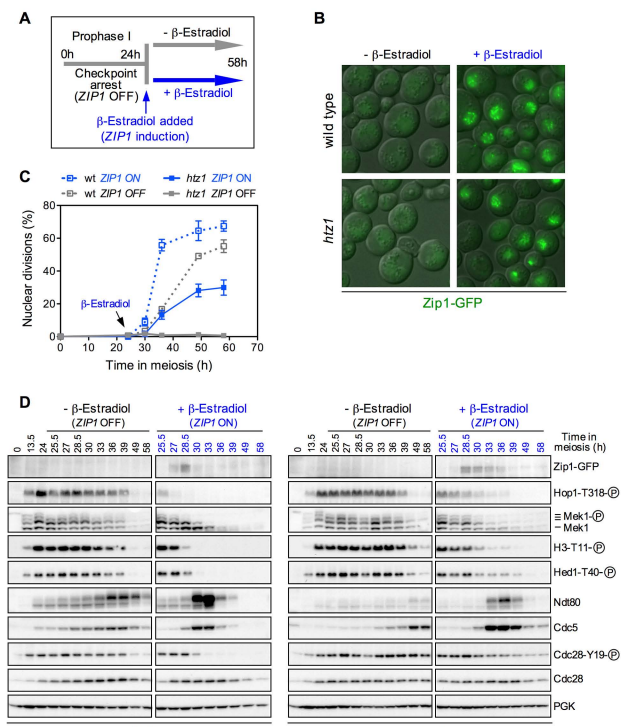
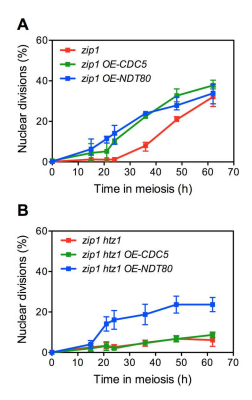


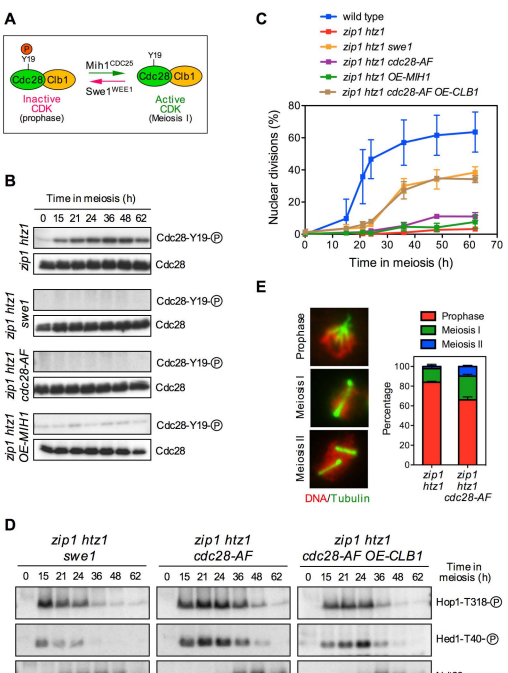
Figure 4

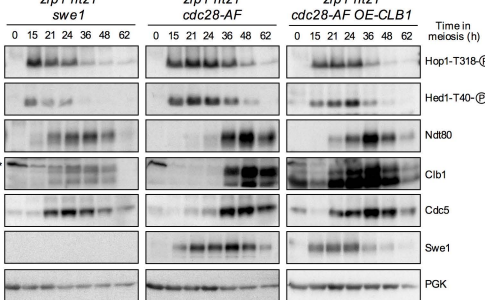


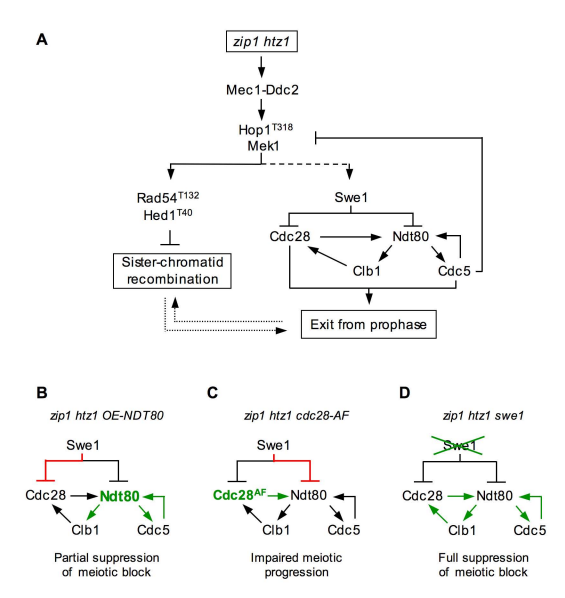
wild type











SUPPLEMENTAL DATA

Functional impact of the H2A.Z histone variant during meiosis in Saccharomyces cerevisiae

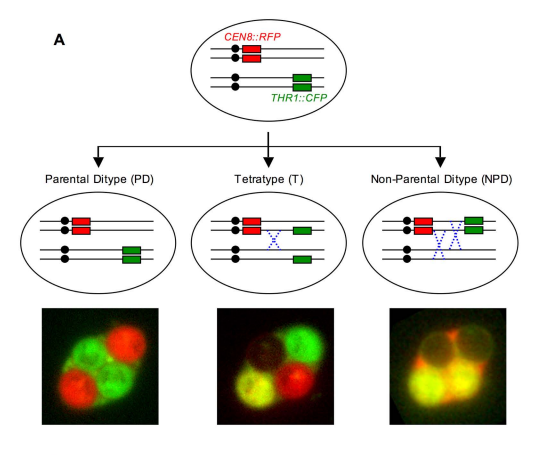
Sara González-Arranz*, Santiago Cavero*, Macarena Morillo-Huesca[†], Eloisa Andújar[‡], Mónica Pérez-Alegre[‡], Félix Prado[†] and Pedro San-Segundo*, ¹

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Supplemental Figures (Figures S1-S8) Table S1 (Strains list) Table S2 (Plasmids list) Table S3 (Antibodies list)



В

	PD(%)	T(%)	NPD(%)	Genetic distance (cM)	n
wild type	61.62	38.00	0.37	20.11	2245
htz 1	55.92	43.72	0.38	22.98	1027
mer3	73.71	16.13	0	8.91	302

Figure S1. Analysis of crossover frequency by spore-autonomous fluorescence assay. (A) Cartoon depicting the potential configuration of the fluorescent reporter markers on Chromosome VIII and representative images of the different types of asci, according to (Thacker *et al.* 2011). Dashed crosses represent possible recombination events leading to each configuration. (B) The table shows the frequency of PD, T and NPD tetrads, map distance expressed in centiMorgan (cM) and the number of tetrads scored (n) pooled from two independent experiments. Strains are DP969 (wild type), DP973 (*htz1*) and DP974 (*mer3*).

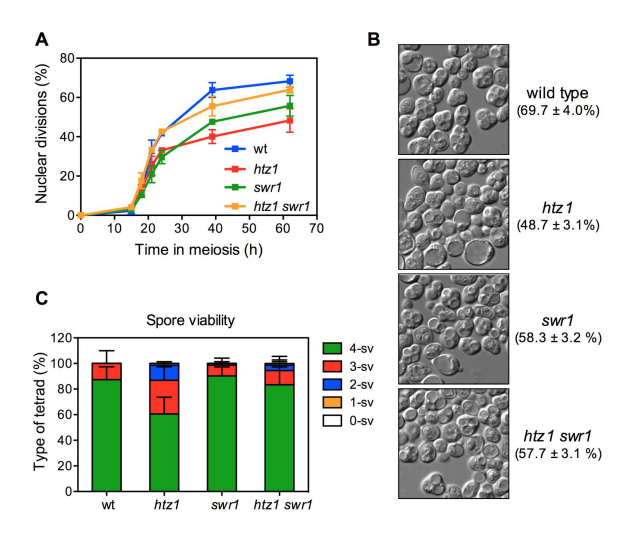
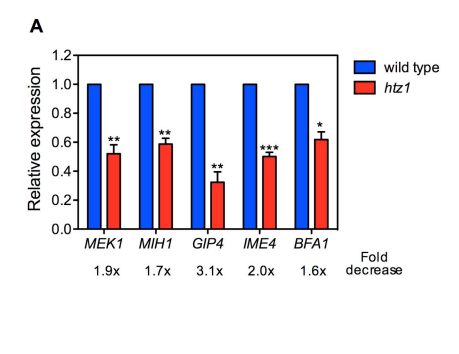
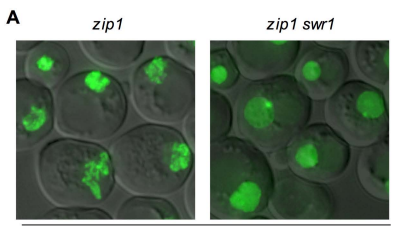


Figure S2. Meiotic impact of the SWR1 complex in the presence or absence of H2A.Z. (A) Time course analysis of meiotic nuclear divisions; the percentage of cells containing two or more nuclei is represented. Error bars: range; n=2. (B) Representative DIC images of asci. The sporulation frequency after 62 hours in liquid sporulation medium is shown in parentheses. (C) Spore viability determined by tetrad dissection. The distribution of tetrad types as the percentage of tetrads with 4, 3, 2, 1 and 0 viable spores (4-sv, 3-sv, 2-sv, 1-sv and 0-sv, respectively) is represented. At least 288 spores were scored for each strain Error bars: range; n=2. Strains are DP421 (wild type), DP630 (*htz1*), DP1174 (*swr1*) and DP1056 (*htz1 swr1*).



B Vegetative prophase 815 133 478 p=5x10⁻⁵

Figure S3. Altered meiotic gene expression in the *htz1* mutant. (A) RT-PCR analysis of mRNA levels of the indicated genes at 15 hours in meiosis. The graph shows relative levels in *htz1* normalized to those in the wild type. Error bars: SD; n=3 (except for *BFA1*; n=2). (B) Venn diagram showing the number of overlapping genes misregulated by *htz1* (1.5-fold cutoff) in vegetative and meiotic cells (15 h). The data for vegetative cells was obtained from (Morillo-Huesca *et al.* 2010). The *p* value was calculated by a hypergeometric test. Strains are DP421 (wild type) and DP1016 (*htz1*).





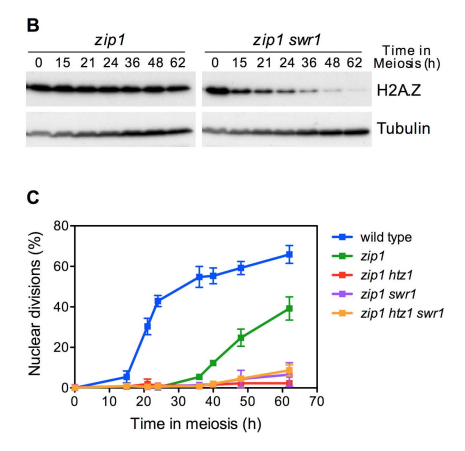


Figure S4. SWR1-dependent chromatin incorporation of H2A.Z is required for normal *zip1-induced meiotic checkpoint response.* (A) Representative images of *zip1* and *zip1 swr1* cells expressing *HTZ1-GFP* at 15 hours after meiotic induction. (B) Western blot analysis of H2A.Z production during meiosis detected with anti-GFP antibodies. Tubulin was used as a loading control. Strains in (A-B) are DP839 (*zip1 HTZ1-GFP*) and DP842 (*zip1 swr1 HTZ1-GFP*). (C) Time course analysis of meiotic nuclear divisions; the percentage of cells containing two or more nuclei is represented. Error bars: SD; n=3. Strains are DP421 (wild type), DP422 (*zip1*), DP776 (*zip1 htz1*), DP804 (*zip1 swr1*) and DP777 (*zip1 swr1 htz1*).

Α Phospho-(S/T)Q Tubulin Chromatin

zip1

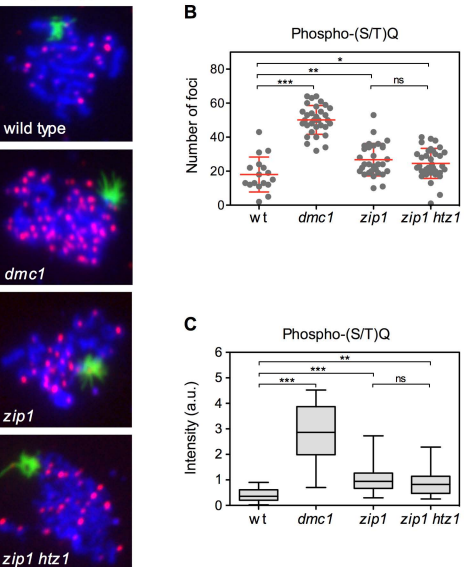


Figure S5. The *zip1 htz1* mutant does not sustain additional Mec1/Tel1-dependent DNA damage signaling compared to zip1. (A) Immunofluorescence of prophase meiotic chromosomes stained with DAPI (blue), and anti-phospho-(S/T)-Q (red) and anti-tubulin (green) antibodies. The *dmc1* mutant was included as a positive control for the accumulation of extensive meiotic DNA damage (unrepaired resected DSBs). Representative nuclei are shown. (B) Quantification of the number of phospho-(S/T)-Q foci per nucleus. Error bars: SD. (C) Quantification of the fluorescence intensity of the phospho-(S/T)-Q signal per nucleus. Whiskers in the box plot represent the maximum to minimum values. The strains used and the number of nuclei scored in (B-C) are: DP421 (wild type; n=17), DP590 (*dmc1*; n=32), DP1525 (*zip1*; n=31) and DP1526 (*zip1 htz1*; n=34).

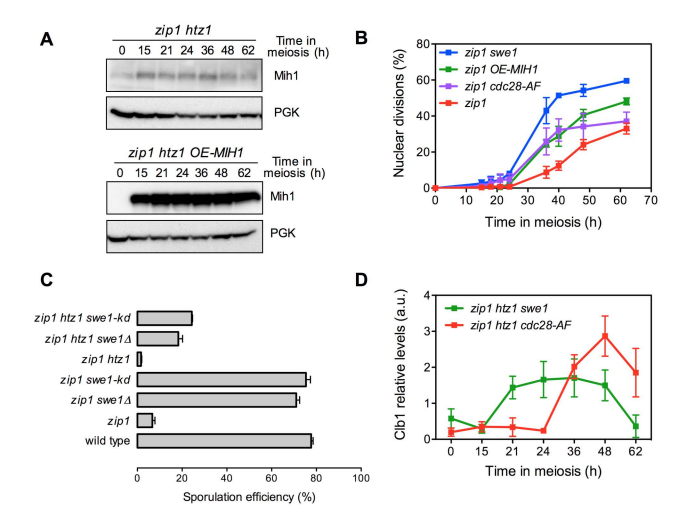


Figure S6. Additional analysis of the role of CDK inhibitory phosphorylation in the meiotic checkpoint. (A) Western blot showing *MIH1-GFP* overexpression from the *HOP1* promoter in a high-copy plasmid. The DP1134 strain (zip1 htz1 MIH1-GFP) was transformed with vector alone (left panels) or with the pSS265 plasmid (right panels). PGK was used as a loading control. Mih1 was detected with anti-GFP antibodies. (B) Faster meiotic progression in zip1 cdc28-AF and zip1 OE-MIH1 compared to zip1. Time course analysis of meiotic nuclear divisions; the percentage of cells containing two or more nuclei is represented. Strains are DP1157 (zip1 swe1), DP1153 (zip1 cdc28-AF) and DP422 transformed with empty vector pSS248 (zip1) or with pSS265 (zip1 OE-MIH1). Error bars: SD; n=3. (C) The kinase-dead swe1-N584A mutant (swe1-kd) phenocopies SWE1 deletion. The sequence changes introduced to generate the *swe1-N584A* mutation in the genomic locus by *delitto perfetto* are shown. The graph represents the sporulation efficiency determined by microscopic counting as the percentage of cells forming mature or immature asci after 3 days on sporulation plates. Error bars: SD; n=3. Strains are: DP1353 (wild type), DP1354 (*zip1*), DP1157 (*zip1 swe1*) and DP1467 (zip1 swe1-kd), DP1414 (zip1 htz1), DP1113 (zip1 htz1 swe1) and DP1468 (zip1 htz1 swel-kd). (D) Quantification of Clb1 levels, normalized to PGK, throughout meiosis in the indicated strains. Error bars: SD; n=3. Strains are DP1113 (zip1 htz1 swe1) and DP1416 (zip1 htz1 cdc28-AF).

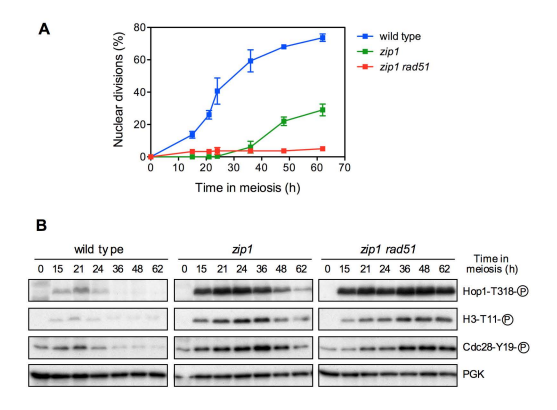


Figure S7. Deletion of *RAD51* leads to sustained checkpoint activation and meiotic arrest in *zip1*. (A) Time course analysis of meiotic nuclear divisions; the percentage of cells containing two or more nuclei is represented. Error bars: SD; n=3. (B) Western blot analysis of the indicated molecular markers of checkpoint activity. PGK was used as a loading control. Strains are DP1359 (wild type), DP1360 (*zip1*) and DP1364 (*zip1 rad51*).

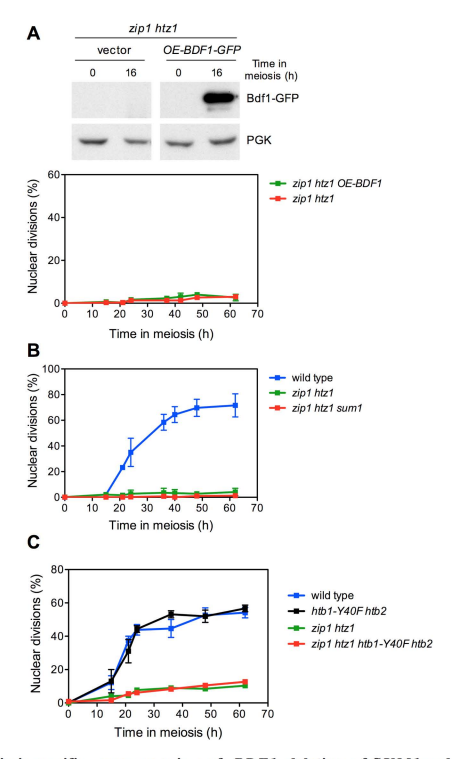


Figure S8. Meiosis-specific overexpression of *BDF1*, deletion of *SUM1* and mutation of *H2B-Y40* do not suppress *zip1 htz1* arrest. (A) Upper panel; western blot analysis of *zip1 htz1* cells (DP1017) transformed with vector (pSS248) or with a 2-micron plasmid expressing *BDF1-GFP* from the *HOP1* promoter (*OE-BDF1-GFP*; pSS354). Extracts were prepared at time zero or 16 hours after meiotic induction (around the peak of prophase) and analyzed with anti-GFP antibodies. PGK was used as a loading control. Lower panel; time course analysis of meiotic nuclear divisions in the same strains. The percentage of cells containing two or more nuclei is represented. Error bars: SD; n=3. (B) Time course analysis of meiotic nuclear divisions in the indicated strains. Wild type (DP421), *zip1 htz1* (DP630) and *zip1 htz1 sum1* (DP1441). Err or bars: range; n=2. (C). Time course analysis of meiotic nuclear divisions in the indicated strains. Wild type (DP1445), *htb1-Y40F htb2* (DP1446), *zip1 htz1* (DP1449) and *zip1 htz1 htb1-Y40F htb2* (DP1450). Error bars: range; n=2.

Tabla S1	Saccharomyces	corovisiao	strains
Table S1.	Succharomyces	cerevisiue	SUI AIIIS

Strain	Genotype	Source
DP421	MATa/MATa leu2,3-112 his4-260 thr1-4 ura3-1 trp1-289 ade2-1 lys2 Δ NheI	PSS Lab
DP422	DP421 zip1::LYS2	PSS Lab
DP437	DP421 ZIP1-GFP	PSS Lab
DP449	DP421 zip1::LYS2 DDC2-GFP::TRP1	PSS Lab
DP590	DP421 dmc1::hphMX4 p306(BrdU-Inc)::URA3/ura3-1	PSS Lab
DP630	DP421 htz1::URA3	This work
DP631	DP421 zip1::LYS2 htz1::URA3	This work
DP713	DP421 mek1::kanMX6	PSS Lab
DP776	DP421 zip1::LYS2 htz1::URA3 DDC2-GFP::TRP1	This work
DP777	DP421 zip1::LSY2 htz1::URA3 swr1::natMX4 DDC2-GFP::TRP1	This work
DP804	DP421 swr1::natMX4 zip1::LSY2 DDC2-GFP::TRP1	This work
DP815	DP421 zip1::LYS2 htz1::URA3 spo11::natMX4 DDC2-GFP::TRP1	This work
DP816	DP421 zip1::LYS2 htz1::URA3 ddc2::TRP1 sml1::KanMX6	This work
DP838	DP421 ZIP1-GFP htz1::URA3	This work
DP839	DP421 zip1::LYS2 HTZ1-GFP::kanMX6	This work
DP840	DP421 HTZ1-GFP::kanMX6	This work
DP841	DP421 swr1::natMX4 HTZ1-GFP::kanMX6	This work
DP842	DP421 zip1::LYS2 swr1::natMX4 HTZ1-GFP::kanMX6	This work
DP969	SK1 CEN8::tdTomato-LEU2/CEN8 THR1:mCerulean-TRP1/THR1	S. Keeney
DP973	SK1 CEN8::tdTomato-LEU2/CEN8 THR1:mCerulean-TRP1/THR1 htz1::hphMX4	This work
DP974	SK1 CEN8::tdTomato-LEU2/CEN8 THR1:mCerulean-TRP1/THR1 mer3::hphMX4	This work
DP1016	DP421 htz1::hphMX4	This work
DP1017	DP421 zip1::LYS2 htz1::hphMX4	This work
DP1056	DP421 htz1::URA3 swr1:: natMX4	This work
DP1113	DP421 zip1::LYS2 htz1::hphMX4 swe1::natMX4	This work
DP1134	DP421 zip1::LYS2 htz1::hphMX4 MIH1-GFP::kanMX6	This work
DP1144	DP421 htz1::URA3 spo11::ADE2	This work
DP1153	DP421 zip1::LYS2 cdc28-AF::LEU2	This work
DP1154	DP421 zip1::LYS2 htz1::hphMX4 cdc28-AF::LEU2	This work

DP1157	DP421 zip1::LYS2 swe1::LEU2	This work
DP1174	DP421 swr1::hphMX4	This work
DP1185	DP421 $TPR1-P_{GALI}$ -ZIP1-GFP ura3:: P_{GPD1} -GAL4(848)ER::URA3/ura3-1	This work
DP1186	DP421 TPR1-P _{GAL1} -ZIP1-GFP P _{GPD1} -GAL4(848)ER::URA3/ura3-1 htz1::hphMX4	This work
DP1259	DP421 htz1::URA3 mek1::kanMX6	This work
DP1353	DP421 3MYC-SWE1	This work
DP1354	DP421 zip1::LYS2 3MYC-SWE1	This work
DP1359	DP421 TUB1/TUB1-GFP::TRP1	PSS Lab
DP1360	DP421 zip1::LYS2 TUB1/TUB1-GFP::TRP1	PSS Lab
DP1364	DP421 zip1::LYS2 rad51::natMX4 TUB1/TUB1-GFP::TRP1	PSS Lab
DP1414	DP421 zip1::LYS2 htz1::hphMX4 3MYC-SWE1	This work
DP1416	DP421 zip1::LYS2 htz1::hphMX4 cdc28-AF::LEU2 3MYC-SWE1	This work
DP1441	DP421 zip1::LYS2 htz1::URA3 sum1::natMX4	This work
DP1445	DP421 [hta1-htb1]::kanMX6 [hta2-htb2]::natMX4 pSS347 (HTA1-HTB1)-TRP1	This work
DP1446	DP421 [hta1-htb1]::kanMX6 [hta2-htb2]::natMX4 pSS348 (HTA1-htb1-Y40F)-TRP1	This work
DP1449	DP421 zip1::LYS2 htz1::hphMX4 [hta1-htb1]::kanMX6 [hta2-htb2]::natMX4 pSS347 (HTA1-HTB1)-TRP1	This work
DP1450	DP421 zip1::LYS2 htz1::hphMX4 [hta1-htb1]::kanMX6 [hta2-htb2]::natMX4 pSS348 (HTA1-htb1-Y40F)-TRP1	This work
DP1467	DP421 zip1::LYS2 3MYC-swe1-N584A	This work
DP1468	DP421 zip1::LYS2 htz1::hphMX4 3MYC-swe1-N584A	This work
DP1523	DP421 spo11::ADE2	This work
DP1524	DP421 zip1::LYS2 spo11::ADE2	This work
DP1525	DP421 zip1::LYS2 p306(BrdU-Inc)::URA3/ura3-1	This work
DP1526	DP421 zip1::LYS2 htz1::hphMX4 p306(BrdU-Inc)::URA3/ura3-1	This work

* All strains are diploids isogenic to BR1919 (Rockmill and Roeder 1990), except strains DP969, DP973 and DP974, which are diploids isogenic to SK1. The haploid parents of DP969 (SK1) were obtained from S. Keeney (Thacker *et al.* 2011). Unless specified, all strains are homozygous for the indicated markers. DP421 is a *lys2* version of the original BR1919-2N.

Rockmill, B., and G. S. Roeder, 1990 Meiosis in asynaptic yeast. Genetics 126: 563-574.

Thacker, D., I. Lam, M. Knop and S. Keeney, 2011 Exploiting spore-autonomous fluorescent protein expression to quantify meiotic chromosome behaviors in *Saccharomyces cerevisiae*. Genetics 189: 423-439.

Plasmid	Vector	Relevant parts	Source/Reference
pTK17	pUC19	htz1::URA3	(SANTISTEBAN et al. 2000)
pJC29	pRS426	2µ URA3 CDC5	(JASPERSEN et al. 1998)
pR2042	pRS305	LEU2 cdc28-AF	(LEU and ROEDER 1999)
pR2045	pRS426	2µ URA3 CLB1	(LEU and ROEDER 1999)
pSS263	pRS426	2µ URA3 NDT80	This work
pSS248	pYES2 derivative	2µ URA3 P _{HOP1} -GFP	This work
pSS265	pSS248	2μ URA3 P _{HOP1} -GFP-MIH1	This work
pSS345	pRS316	CEN6 URA3 HTA1-HTB1	This work
pSS347	pRS314	CEN6 TRP1 HTA1-HTB1	This work
pSS348	pRS314	CEN6 TRP1 HTA1-htb1-Y40F	This work
pSS354	pSS248	2µ URA3 P _{HOP1} -GFP-BDF1	This work

Table S2. Plasmids

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- LEU, J. Y., and G. S. ROEDER, 1999 The pachytene checkpoint in *S. cerevisiae* depends on Swelmediated phosphorylation of the cyclin-dependent kinase Cdc28. Mol Cell **4:** 805-814.
- SANTISTEBAN, M. S., T. KALASHNIKOVA and M. M. SMITH, 2000 Histone H2A.Z regulates transcription and is partially redundant with nucleosome remodeling complexes. Cell **103**: 411-422.

Antibody	Host and type	Application* (Dilution)	Source / Reference
Cdc2-Y15-P (Cdc28-Y19-P)	Rabbit polyclonal	WB (1:1000)	Cell Signaling Technology #9111
Phospho-(S/T)Q	Rabbit polyclonal	IF (1:200)	Cell Signaling Technology #2851
Cdc5	Goat polyclonal	WB (1:1000)	Santa Cruz Biotechnology sc-6733
Cdc28 (PSTAIRE)	Rabbit polyclonal	WB (1:1000)	Santa Cruz Biotechnology; sc53
Clb1	Goat polyclonal	WB (1:100)	Santa Cruz Biotechnology; sc-7647
Hed1	Rabbit polyclonal	WB (1:20000)	N. Hollingsworth (CALLENDER <i>et al.</i> 2016)
Hed1-T40-P	Rabbit polyclonal	WB (1:50000)	N. Hollingsworth (CALLENDER <i>et al.</i> 2016)
Hop1-T318-P	Rabbit polyclonal	WB (1:1000)	J. Carballo (PENEDOS <i>et al.</i> 2015)
H3-T11-P	Rabbit polyclonal	WB (1:2000)	Abcam ab5168
Mek1	Rabbit polyclonal	WB (1:1000)	PSS Lab (ONTOSO <i>et al.</i> 2013)
Ndt80	Rabbit polyclonal	WB (1:5000)	M. Lichten (BENJAMIN <i>et al.</i> 2003)
Rad51	Rabbit polyclonal	IF (1:300)	Santa Cruz Biotechnology sc-33626
Мус	Rabbit polyclonal	WB (1:1000)	Sigma c3956
Pch2	Rabbit polyclonal	IF (1:400)	PSS Lab / R. Freire
GFP (JL-8)	Mouse monoclonal	WB (1:2000-10000)	Clontech 632381
PGK (22C5D8)	Mouse monoclonal	WB (1:2000)	Invitrogene 459240
Tubulin (TAT1)	Mouse monoclonal	WB (1:10000) IF (1:500)	K. Gull (ACOSTA <i>et al.</i> 2011)

 Table S3. Primary antibodies

*WB, western blot; IF, immunofluorescence

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