The genetics of human infertility by functional interrogation of SNPs in mice

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Infertility is a prevalent health issue, affecting ~15% of couples of childbearing age. Nearly one-half of idiopathic infertility cases are thought to have a genetic basis, but the underlying causes are largely unknown. Traditional methods for studying inheritance, such as genome-wide association studies and linkage analyses, have been confounded by the genetic and phenotypic complexity of reproductive processes. Here we describe an association- and linkagefree approach to identify segregating infertility alleles, in which CRISPR/Cas9 genome editing is used to model putatively deleterious nonsynonymous SNPs (nsSNPs) in the mouse orthologs of fertility genes. Mice bearing "humanized" alleles of four essential meiosis genes, each predicted to be deleterious by most of the commonly used algorithms for analyzing functional SNP consequences, were examined for fertility and reproductive defects. Only a Cdk2 allele mimicking SNP rs3087335, which alters an inhibitory WEE1 protein kinase phosphorylation site, caused infertility and revealed a novel function in regulating spermatogonial stem cell maintenance. Our data indicate that segregating infertility alleles exist in human populations. Furthermore, whereas computational prediction of SNP effects is useful for identifying candidate causal mutations for diverse diseases, this study underscores the need for in vivo functional evaluation of physiological consequences. This approach can revolutionize personalized reproductive genetics by establishing a permanent reference of benign vs. infertile alleles.

CRISPR/Cas9 | meiosis | spermatogenesis | cyclin | genome editing

espite the high incidence of infertility, autosomal genetic Causes of infertility attributable to gametogenesis defects are poorly characterized. In males, the most common known genetic causes of infertility are Y chromosome microdeletions, thought to be responsible for 6-18% of nonobstructive azoospermia (NOA) or severe oligozoospermia cases (1). In females, most of the known genetic causes are linked to syndromes that also affect the soma (e.g., Kallmann, Turner) or the neuroendocrine axis (2). In rare cases, families segregating infertility alleles have been mapped by linkage (3-6). Several candidate gene resequencing studies have implicated mutations or SNPs as being causative for azoospermia (7–13), but in the absence of genetic data, only a few reports (e.g., ref. 14) have made a compelling case. Recently, hemizygous deletions of the TEX11 gene, presumably catalyzed by unequal recombination between repetitive elements in the locus, have been linked to maturation arrest and infertility in azoospermic men (15).

Gene knockout and molecular genetic studies in mice have shown that germ cell development is genetically complex. A screen of the Mouse Genome Informatics database using MouseMine (www.mousemine.org) identifies 728 genes currently associated with infertility. Clearly, many more genes required for fertility remain to be identified. Furthermore, infertility is genetically heterogeneous; scores of distinct genes cause grossly identical phenotypes when mutated in mice (2, 16). This likely explains why genome-wide association studies (GWAS) have not been effective even in stratified cohorts, with only two reporting significant associations with NOA in Chinese populations (11, 17). Even if associations could be readily obtained, identification and validation of causative variants would remain problematic. Finally, the proportion of Mendelian infertilities that are caused by de novo mutations vs. segregating polymorphisms is unknown. Clearly, different approaches are needed to address the genetics of human infertility.

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Here we describe a reverse genetics approach for identifying infertility alleles segregating in human populations that does not require linkage or association data; rather, it combines in silico prediction of deleterious allelic variants with functional validation in CRISPR/Cas9-edited "humanized" mouse models. We modeled four nonsynonymous human SNPs (nsSNPs) in genes that are essential for meiosis in mice. Each of these nsSNPs has been predicted to be deleterious to protein function by several widely used algorithms. Only one of the nsSNPs was found to cause infertility, highlighting the importance of experimentally evaluating computationally predicted disease SNPs.

Results

To address the question of whether human infertility can be caused by segregating Mendelian alleles, we applied a precision genome editing approach (Fig. 1*A*) to model variants of meiosis genes that when knocked out in mice cause the drastic infertility phenotype of NOA from meiotic arrest (also called "maturation arrest" in humans). To identify candidate infertility alleles in such genes, we screened the dbSNP database for potentially deleterious nsSNPs that met the following criteria: (*i*) they exist as minor (low frequency, preferably <1–2%) alleles in humans; (*ii*) they alter evolutionarily conserved amino acids; and (*iii*) the changes are predicted to be functionally deleterious by multiple algorithms.

Meiosis gene nsSNPs were first ranked using two of the most commonly used computational tools for predicting the likelihood of a variant being deleterious: SIFT (18) and Polyphen-2 (19). Four of the highest-scoring SNPs identified by both algorithms, one each in the genes *Cyclin-dependent kinase* 2 (*Cdk2*), *MutL*

Significance

Although infertility is a highly prevalent disease with a major genetic component, the underlying genetic causes are unknown in the vast majority of patients. The complexity of sperm and egg production has confounded most efforts to address the genetics of human infertility. We have developed an experimental strategy for identifying infertility-causing mutations that exist in human populations. This strategy exploits the expanding databases of human genetic variation to computationally identify potential infertility mutations, then mimic them in mice by CRISPR/Cas genome editing technology. The mice are then evaluated for whether the human mutation renders them infertile. This technology can be used to establish a permanent resource of validated disease-causing genetic variants in infertility and other diseases.

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A	nsSNPs in known meiosis genes Predicted deleterious SNPs dbSNP MAF <1-2%				→ rs3087335 CRISPR/Cas9 rs63750447 rs61735519 rs147871035 Fertility Analysis		
В	Gene dbSNP ID		AA alt	Allele	KO Phenotype		
				Frequency	Male	Female	
	CDK2	CDK2 rs3087335 Y15S		0.006	pairing, non-homologous synapsis unrepaired DSBs	Infertile	
	MLH1	rs63750447	V384D 0.009 Infertile; crossover failure, Metaphase arrest		Infertile; crossover failure, Metaphase arrest	Infertile; crossover failure	
	SMC1B	rs61735519	Infertile; pachytene arrest, short SCs, abnormal sister chromatid cohesion, crossover failure		Infertile; identical defects as in males		
	TEX15	rs147871035	T2181I	0.001	Infertile, zygotene/pachytene arrest, unrepaired DSB, synapsis failure	Fertile	

homolog 1 (Mlh1), Structural maintenance of chromosomes protein 1B (Smc1b), and Testis-expressed gene 15 (Tex15) (Fig. 1B), were then evaluated with six additional tools that predict functional consequences of SNPs: Mutation Assessor, PANTHER, PhD-SNP, SNPs&GO, SNAP, and CADD (20–25). Overall, each of these four nsSNPs was predicted to be deleterious by at least five of the eight computational algorithms used (Table 1). These four genes have crucial roles in meiosis, affecting processes including recombination, homolog synapsis, DNA repair, and chromatid cohesion (Fig. 1B).

For each of the four SNPs, the orthologous amino acid changes were introduced into the mouse genes using CRISPR/ Cas9-mediated genome editing in mouse embryos. Specifically, single-celled embryos were comicroinjected with Cas9 mRNA, sgRNA, and a synthetic single-stranded oligodeoxynucleotide (ssODN) containing nucleotide changes (Fig. 2B, Fig. S1 A-C, and Table S1) that, via homologous recombination stimulated by targeted Cas9 double-strand breaks (DSBs), introduced the desired amino acid change. We identified edited founder mice (Materials and Methods) carrying at least one desired humanized allele of Cdk2, Mlh1, Smc1b, and Tex15 at frequencies of 44.4%, 29.1%, 21.4%, and 56.2%, respectively (Table S2). In the experiments that generated the Cdk2 and Tex15 alleles used here, a Next-Gen sequencing strategy was used to confirm that there was no evidence of off-target editing at degenerate sgRNA recognition sites (26).

To determine whether the humanized alleles compromised reproduction, we bred each to homozygosity, tested fertility by mating, and performed a series of histological and cytological analyses. In males, we examined epididymal sperm number and morphology at progressive ages. In addition, we assessed testis

Algorithm	D Min	CDK2 rs3087335 Cdk2 ^{Y155}	MLH1 rs63750447 Mlh1 ^{v384D}	SMC1B rs61735519 Smc1b ^{F1055L}	TEX15 rs147871035 Tex15 ^{T21811}
SIFT	≤ 0.05	D (0)	D (0)	D (0)	D (0)
PolyPhen 2	>0.5	D (1)	D (0.99)	D (0.99)	D (0.92)
Mutation	>0.65	D (2.3)	D (3.2)	D (3.2)	D (1.8)
Assessor					
PANTHER	≥0.5	D (0.8)	D (0.5)	D (0.7)	T (0.4)
PhD-SNP	≥0.5	D (0.9)	D (0.9)	T (0.4)	D (0.7)
SNPs&GO	≥0.5	D (0.9)	D (0.9)	T (0.4)	D (0.7)
SNAP	≥5	D (5)	D (5)	T (2)	T (3)
CADD*	>15	D (23)	D (19.5)	D (25.5)	D (24.3)

The allele for each SNP (rs no.) modeled is indicated. D Min refers to the minimum score for deleterious classification; D, deleterious; T, tolerated. *CADD scores based on v1.1.

Fig. 1. Generating humanized mouse models to test potential human infertility SNPs. (A) Workflow for identifying and functionally testing potential infertility alleles, including the four (in red) studied here. (*B*) Candidate infertility SNPs modeled and tested in this study. MAF, minor allele frequency; AA alt, amino acid alteration; KO, knockout.

size, weight, and histology. In the cases of adult males homozygous for the humanized alleles of rs63750447 ($Mlh1^{V376D}$), rs61735519 ($Smc1b^{F1055L}$), and rs147871035 ($Tex15^{2181I}$), histological analysis of testes of these mutants did not reveal any phenotypic defects; the seminiferous tubules exhibited normal spermatogenesis that was indistinguishable from that in wild type (WT) (Fig. S1). As predicted from the testes histology, the mutants exhibited no significant difference in epididymal sperm counts compared with WT, an assay that also reflects sperm motility (Fig. S2), and a homozygote of each was proven to be fertile in matings.

Null alleles of these genes, which encode proteins involved in meiotic chromosome cohesion, crossing-over, and repair of meiotic DSBs, respectively, cause meiotic arrest with characteristic defects in meiotic prophase I chromosomes (27-29). To determine whether there was evidence of any such defects in these mice, we immunolabeled surface-spread spermatocyte nuclei for proteins diagnostic of key meiotic events. For each humanized allele, patterns of yH2AX (present at meiotic DSBs and on asynapsed chromosomes), SYCP1 (chromosome synapsis marker), SYCP3 (a synaptonemal complex axial element protein), and MLH1 foci (which mark sites of chiasmata) revealed no differences between WT and mutants (Fig. S3 A-L), including in the number of MLH1 foci (Fig. S3M). In addition, whereas mouse knockouts of Smc1b and Mlh1 cause female infertility from meiotic failure, the female homozygotes for each humanized allele were fertile and normally fecund (Fig. S44). These data indicate that rs61735519 (SMC1B), rs63750447 (MLH1), and rs147871035 (TEX15) do not impact fertility, despite the high algorithm prediction scores.

In contrast, male mice homozygous for the CDK2-Y15S alteration, mimicking human rs3087335, severely disrupted spermatogenesis. $Cdk2^{Y15S/Y15S}$ males (n = 13) had markedly smaller testes than WT siblings (n = 12), weighing ~80% less (Fig. 3 A and B). These mutants were devoid of sperm in their epididymides, and failed to impregnate WT female partners despite plugging them. Histological examination of $Cdk2^{YI5S/YI5S}$ testes revealed an arrest of meiotic progression at the pachytene spermatocyte stage; no metaphase spermatocytes or postmeiotic spermatids were observed (Fig. 3 C-E), consistent with the phenotype of Cdk^2 null mice (30, 31). Furthermore, spermatocyte meiotic prophase I chromosomes exhibited abnormalities common in null mutants (32), including telomere fusions, XY body defects, and unrepaired DSBs, all of which prevent mutant spermatocytes from progressing past the midpachytene stage (Fig. 4). There were three differences from the knockout, however. First, unlike null females, *Cdk2*^{YI5S/YI5S} females were fertile, delivering an average of approximately seven pups per litter (Fig. S4A). The mutant ovaries had grossly normal histology and numbers of follicles (both primordial pool and developing follicles) that were not significantly different from that in WT at



Fig. 2. CRISPR/Cas9 modeling of *CDK2* SNP rs3087335. (*A*) Domains in the hCDK2 protein and location of rs3087335 (red arrow). (*B*) CRISPR/Cas9 editing strategy for making the Y15S alteration. The Tyr15 site is conserved in humans and mice. PAM nucleotides are in green. Underscored nucleotides match the CRISPR gRNA. To avoid Cas9 recutting of an already-edited locus, two silent mutations (blue) were included into the ssODN recombination template, only part of which is shown (Table S1). (*C*) Sanger sequencing chromatograms of control and mutant mice showing correct CRISPR/Cas9 targeting.

postnatal day (P) 21 (Fig. S4 *B* and *C*). Second, whereas $Cdk2^{-/-}$ seminiferous tubules retained spermatogonia that continued to initiate waves of spermatogenesis until at least P120 (31), $Cdk2^{Y15S/Y15S}$ seminiferous tubules became devoid of germ cells, marked by mouse vasa homolog (MvH), by P90; they exhibited a Sertoli cell-only phenotype (Fig. 3*E* and Fig. S5*B*). As discussed later, this is indicative of spermatogonial stem cell maintenance or a proliferation defect. We also noticed pyknotic cells (presumably spermatogenic wave progress as far as the pachytene spermatocyte stage (Fig. 3*C*).

The third distinction between the $Cdk2^{Y15S}$ and null alleles appeared in heterozygous males. At P90, $Cdk2^{Y15S/+}$ males exhibited smaller testes of approximately one-half the weight of WT testes (Fig. 3 *A* and *B*), abnormal seminiferous tubule histology characterized by large multinucleate cells and decreased luminal elongated spermatids (Fig. 3*E*), and a ~3.5-fold reduction in epididymal sperm count compared with WT littermates (Fig. 3*F*). The seminiferous tubules became more disorganized by P180 (Fig. S5 *C* and *D*). Mice heterozygous for a *Cdk2* null allele do not exhibit meiotic chromosome abnormalities or infertility (32), and the $Cdk2^{Y15S}$ allele did not affect CDK2 levels (Fig. 3*G*). Thus, $Cdk2^{Y15S}$ appears to be a semidominant, gain-of-function allele.

Discussion

In this paper, we introduce an association- and linkage-free approach to studying the genetics of human infertility that can be applicable to other diseases, implemented either in a purely reverse genetic paradigm as described here or as a validation tool for candidate SNPs implicated by genetic mapping. The speed and efficacy with which CRISPR/Cas9 can be used to generate precisely engineered mouse models enables powerful and physiologically relevant functional assessment of human genetic variants.

This is especially important in cases of infertility, for which accurate in vitro systems are lacking (33).

A surprising outcome of the present study was the poor accuracy of computational tools designed to predict functional consequences of SNPs. Whereas all eight algorithms correctly predicted that the SNP in CDK2 would disrupt gene function, all classified the MLH1 SNP as being functionally damaging, and most also incorrectly predicted the same for the SNPs in SMC1B and TEX15. SNAP, a neural network-based tool (24), was the most accurate (75% correctly predicted) for the small sample size tested here. Nonetheless, it remains possible that the SNPs affect protein function in ways that do not grossly impact germ cell function. It also should be cautioned that the SNPs may be deleterious in humans but not in mice. Also noteworthy is the fact that the CDK2 SNP was the only SNP tested in which the affected amino acid resided in an annotated functional domain of the protein, and in this case, the amino acid itself was known to be a target of WEE1 kinase (Fig. 24 and Fig. S1 A-C). We surmise that manual scrutiny of nsSNPs will complement algorithm predictions. Our finding that $CDK2^{Y15S}$ disrupts spermatogenesis is re-

Our finding that $CDK2^{PDS}$ disrupts spermatogenesis is remarkable in two respects. First, it provides evidence that semidominant infertility alleles can segregate in human populations. This particular allele exists at a low frequency, having been identified in only 2 of 688 alleles sampled (dbSNP build 142). Because it affects only male mice (with full penetrance in homozygotes and with a partial, age-related germ cell depletion phenotype in heterozygotes), transmission may occur only through women. If men heterozygous for $CDK2^{YISS}$ are fertile (perhaps only at earlier ages), then homozygous males would occur at a rate of ~1 in 236,672. The allele frequencies in different demographic populations remain unclear, however. In mice, a fully penetrant, dominant male-specific sterile allele of the meiotic recombination gene Dmc1 has provided evidence



Fig. 3. The CDK2^{Y155} alteration encoded by rs3087335 causes male infertility. (*A*) Testes (90 d old) from littermates of indicated genotypes. (*B*) Testes weights of P90 heterozygous and homozygous mice were 54.6% and 81.5% lower, respectively, than those of WT. (*C–E*) H&E-stained seminiferous tubule cross-sections of 15-, 25-, and 90-d-old testes. Arrowheads indicate degenerating cells. (*F*) Sperm counts of 3- and 6-mo-old WT and heterozygous males. (*G*) Western blot analysis of CDK2 in spleen. Genotypes are indicated, as are approximate molecular weights of the detected species with anti-CDK2 antibody. Spleens were used instead of testes, because differences in cellularity of mutant testes would compromise interpretation of relative protein amounts. *****P* < 0.0002; ****P* < 0.0001; ***P* < 0.001. Data are presented as mean \pm SD of at least three animals. Ser, Sertoli cells. (Scale bars: 100 µm).

that dominant infertility mutations can arise in genes that have a recessive character as null alleles, and that such mutations can be transmitted through the unaffected sex (34). Finally, these cases indicate the difficulty in predicting the mode of inheritance and associated phenotypes purely by computational means. Second, the $CDK2^{YISS}$ allele reveals a novel role for the function

of CDK2 in vivo. Histological and cytological analyses indicated that, along with the null-like meiotic arrest of $Cdk2^{Y15S/Y15S}$ spermatocytes during the first wave of spermatogenesis, homozygotes became germ cell-depleted, unlike the null alleles. Given that phosphorylation of Tyr15 by WEE1 kinase inhibits CDK2 activity and thus cell cycle progression in mitotic cells (35-37), we speculate that mutation of the inhibitory phosphorylation site in this allele disturbs the proper cell cycle control of spermatogonial stem cells, affecting the regulation of self-renewal and differentiation of stem cells. It will be of interest to determine whether Cdk2Y15S allele triggers inappropriate proliferation or differentiation of spermatogonial stem cells at the expense of proliferation or renewal of the stem cell population. Consistent with this, a previous study found that when both of the CDK2 inhibitory phosphorylation sites at threonine 14 and tyrosine 15 were changed to alanine and phenylalanine, respectively, heterozygous male mice were rendered subfertile or sterile, and that somatic cells bearing these mutations entered S phase improperly (38). We suspect that in Cdk2 null mice, control of the spermatogonial cell cycle or meiotic entry is accomplished by another cyclin, such as Cdk1 (39). The absence of a self-renewing oogonial stem cell population may explain the fertility of Cdk^{YI5S} homozygous females. We caution that our alleles were generated in a different genetic background (FvB × C57BL/6) than that of mice bearing null alleles (mixes of 129 and C57BL/6, as far as can be judged from the literature) (30–32).

The complexity of male germ cell differentiation, coupled with limited information on the biochemistry of germ cell-specific genes, complicates the task of identifying true segregating infertility alleles in humans. Our results demonstrate that whereas computational predictions of SNP effects are useful for identifying candidate causal mutations for diverse diseases, including those mapped by GWAS, functional evaluation in vivo is essential to define true physiological consequences. It is likely that computational tools can be improved with increased knowledge of protein structure and function, coupled with a larger body of training data generated by methods described in this report; however, a present shortcoming for identifying human disease SNPs in mice is that this is limited to nsSNPs that alter conserved amino acids, and that addressing noncoding SNPs will be more difficult. Nevertheless, the efficiency of this approach can revolutionize reproductive genetics by establishing a permanent reference of deleterious vs. benign alleles, which will help clarify the molecular basis of idiopathic infertilities in individual patients. Furthermore, because this approach can be



Fig. 4. Meiotic defects in $Cdk2^{\gamma_{155}/\gamma_{155}}$ spermatocytes. (*A*–*J*) Immunolabeling of surface-spread spermatocyte nuclei. Genotypes, antibodies used, and pachytene substages are indicated. Histone H1t appears in mid-pachynema. Note abnormal nonsynaptic fusions (arrowheads in *B* and *D*) and abnormal XY body (arrows in *B* and *D*). Meiotic DSBs (indicated by γ H2AX foci) in WT spermatocytes (*G*) are repaired by late pachynema (*I*); however, mutant chromosomes retain γ H2AX (which marks both unrepaired DSBs and areas of asynapsed chromosomes) and these spermatocytes do not progress beyond a mid/late pachynema-like stage (*J* and *J*). Boxed areas are enlarged to highlight specific abnormalities (*A*′–*J*′). XY, XY body (also known as the "sex body"); arrow, abnormal XY body; arrowhead, telomeric fusion; pachy, pachynema. (Scale bars: 10 µm.)

applied to any genetic disease, it should be a powerful tool in the era of personalized genomic medicine.

Materials and Methods

Computational Identification of Potential Infertility SNPs. The Ensembl Genome Browser (40) was used to extract lists of nsSNPs in several meiosis genes that were predicted to be deleterious based on precomputed scores from the SIFT (v5.0.2) (18) and PolyPhen-2 (v2.2.2) (19) algorithms. The SNPs selected were limited to those with frequency information from multiple sources (e.g., 1000 Genomes Project, HapMap Project) or multiple individuals. The highest-scoring SNPs were screened with the other six algorithms listed in Table 1.

Production of CRISPR/Cas9-Edited Mice. sgRNAs (Table S2) were designed and produced as described previously (26). The ssODN source was "Ultramer" oligos (Integrated DNA Technologies). The plasmid pST1374-NLS-flag-linker-Cas9 (44758; Addgene) was used for generating Cas9 mRNA after linearization with Agel. In vitro transcription and capping were performed using the mMESSAGE mMACHINE T7 Transcription Kit (AM1345; Life Technologies). Embryo microinjections into zygotes [F1 hybrids between strains FVB/ NJ and B6(Cg)-Tyr^{c-2J}/J] were performed as described previously (26) using the reagent concentrations listed in Table S1. Edited founders were identified either by Next-Gen sequencing or by subcloning followed by Sanger sequencing as described previously (26). Genotyping was performed by

sequencing or by restriction enzyme analysis of PCR products produced from DNA isolated from ear punch biopsy specimens. Primer sequences and their respective amplicon sizes for genotyping for each mutant mouse line are listed in Table S3. Annealing temperature was 64 °C for *Cdk2*, 60 °C for *Mlh1*, 62 °C for *Smc1b*, and 60 °C for *Tex15*.

Mice and Reproductive Phenotypes. The alleles generated in this study and their abbreviations (in parentheses) were as follows: $Cdk2^{em1/cs}$ ($Cdk2^{Y155}$), $Mlh1^{em1/cs}$ ($Mlh1^{V384D}$), $Smc1b^{em1/cs}$ ($Smc1b^{F1055L}$), and $Tex15^{em1/cs}$ ($Tex15^{T2181}$). For female fertility tests, 8-wk-old animals were housed with age-matched FVB/NJ males. For the $Cdk2^{Y155}$ allele, there was perfect genotype–phenotype correlation; all 13 homozygous males (P25–P180) and 8 heterozygous males (P90 and P180) examined, but none of the 12 WT littermates, had small gonads.

Histology and Immunohistochemistry. Testes and ovaries were fixed for 24 h at 23 °C in Bouin's solution, paraffin-embedded, sectioned at 5 μ m, and then stained with H&E. Ovarian follicle quantification was done by counting the every fifth section, as described previously (41). For immunofluorescence, testes were fixed in 4% paraformaldehyde for ~24 h, paraffin-embedded, sectioned at 7 μ m, deparaffinized, and then boiled for 30 min in 10 mM citric acid (pH 6.0) for antigen retrieval. Sections were blocked in PBS containing 5% goat serum for 30 min at 23 °C, followed by incubation with primary antibody (rabbit anti-MVH; 1:500, ab13840; Abcam) for 12 h at 4 °C and detection

by the secondary antibody (goat anti-rabbit IgG 488; 1:1,000, A11008; Molecular Probes).

Immunocytology of Meiotic Chromosomes. Preparation of surface-spread spermatocyte chromosomes and immunolabeling was done as described previously (42, 43). Primary antibodies were rabbit anti-CDK2 (1:50, ab7954-1; Abcam), rabbit anti-SYCP3 (1:500, ab15093; Abcam), mouse anti-SYCP3 (1:500, ab97672; Abcam), rabbit anti-SYCP1 (1:500, ab15087; Abcam), mouse anti-γH2AX (1:500, 05-636; Millipore), guinea pig anti-H1t (1:1,000, a gift from M. A. Handel, The Jackson Laboratory, Bar Harbor, ME), and mouse anti-mLH1 (1:50, 550838; BD Pharmingen). Secondary antibodies were goat anti rabbit-IgG 488 (1:1,000, A11008; Molecular Probes), goat ant-rabbit IgG 594 (1:1,000, A11012; Molecular Probes), goat anti-guinea pig IgG 647 (1:1,000, A21450; Molecular Probes). Nuclei were counterstained with DAPI, and slides were mounted with ProLong Gold antifade reagent (P36930; Molecular Probes).

Western Blot Analysis. Protein was extracted from spleens using RIPA buffer (9806S; Cell Signaling Technology) supplemented with protease inhibitors (cOmplete, Mini, 11836153001; Roche). Extracts were separated by SDS/PAGE (12% acrylamide), followed by electrotransfer to a PVDF membrane (Immobilon-P membrane, IPVH00010; EMD Millipore), and then blocked in 5% nonfat milk.

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Membranes were probed with rabbit anti-CDK2 (1:200, ab7954-1; Abcam) for 2 h at room temperature and then washed with Tris-buffered saline with 0.1% Tween-20, followed by a 1-h incubation in anti-rabbit IgG HRP-linked secondary antibody (70745; Cell Signaling Technology). Signal was detected using Luminata Classico Western HRP substrate (WBLUC0100; EMD Millipore).

Imaging. Surface-spread preparations were imaged with an Olympus BX51 microscope with a $60\times/0.90$ NA infinity/0.11–0.23 objective, and Magnafire 2.0 software (Olympus). Histological sections were imaged with the same microscope using a $20\times(20\times/0.05;$ NA infinity/0.17) objective. Cropping, color, and contrast adjustments were made with Adobe Photoshop CC 2014, using identical background adjustments for all images.

Sperm Counts. Cauda epididymides (3–6 mo old) were minced in 2 mL of PBS and then incubated for 15 min at 37 °C to allow spermatozoa to swim out. The sperm were diluted 1:5 in PBS, incubated for 1 min at 60 °C, and then counted with a hematocytometer.

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