

The Ubiquitin-Conjugating DNA Repair Enzyme HR6A Is a Maternal Factor Essential for Early Embryonic Development in Mice

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The *Saccharomyces cerevisiae* RAD6 protein is required for a surprising diversity of cellular processes, including sporulation and replicational damage bypass of DNA lesions. In mammals, two RAD6-related genes, HR6A and HR6B, encode highly homologous proteins. Here, we describe the phenotype of cells and mice deficient for the *mHR6A* gene. Just like *mHR6B* knockout mouse embryonic fibroblasts, *mHR6A*-deficient cells appear to have normal DNA damage resistance properties, but *mHR6A* knockout male and female mice display a small decrease in body weight. The necessity for at least one functional *mHR6A* (X-chromosomal) or *mHR6B* (autosomal) allele in all somatic cell types is supported by the fact that neither animals lacking both proteins nor females with only one intact *mHR6A* allele are viable. In striking contrast to *mHR6B* knockout males, which show a severe spermatogenic defect, *mHR6A* knockout males are normally fertile. However, *mHR6A* knockout females fail to produce offspring despite a normal ovarian histology and ovulation. The absence of *mHR6A* in oocytes prevents development beyond the embryonic two-cell stage but does not result in an aberrant methylation pattern of histone H3 at this early stage of mouse embryonic development. These observations support redundant but dose-dependent roles for HR6A and HR6B in somatic cell types and germ line cells in mammals.

Repair of damaged DNA requires a number of complementary, partially overlapping mechanisms, including mismatch repair, nucleotide and base excision repair, and repair pathways that deal with double-strand breaks. S-phase arrest caused by unrepaired DNA damage is prevented by a mechanism known as postreplication repair, which we prefer to rename replicational damage bypass (RDB). This bypass mechanism allows replication across lesions instead of repairing lesions (20, 26).

Based on the mutant phenotype, RAD6 is one of the key players in RDB in yeast. *rad6* mutants are extremely sensitive to a wide range of DNA-damaging agents and show an increased spontaneous mutation frequency and a loss of DNA damage-induced mutagenesis (41, 46). In addition, RAD6 is also involved in a number of other cellular processes in yeast, including the degradation of proteins via the N-end rule, retrotransposition, gene silencing, meiosis, and sporulation (14, 15, 28, 52). RAD6 was identified as a ubiquitin-conjugating enzyme (29), and all of the functions of RAD6 in yeast depend on its ability to transfer ubiquitin molecules to a protein substrate (66). The ubiquitin-conjugating pathway requires the consecutive activity of ubiquitin-activating (E1), -conjugating (E2), and -ligating (E3) enzymes and results in mono- or polyubiquitination of substrate proteins. The ubiquitination serves as a signal for modification, activation, or proteolytic degradation via the 26S proteasome. The identification of 11 ubiquitin-conjugating enzymes in yeast and at least 20 genes encoding

such proteins in the human genome, as well as the existence of a vast number of E3s, suggests that high substrate specificity can be obtained by combining different E2s and E3s (24). RAD18, UBR1, and BRE1 are E3 proteins that interact with RAD6 and are involved in RDB, degradation of proteins via the N-end rule, and chromatin modification, respectively (7, 15, 77). Ubiquitination of histone H2B at lysine residue 123 by RAD6 is essential for sporulation in yeast (54). Recently, it was shown that ubiquitination of H2B is a prerequisite for the methylation of histone H3 at specific lysine residues, with BRE1 being the putative E3 involved in this process (11, 16, 48, 65). PCNA (proliferating cell nuclear antigen) has only recently been identified as an important target for RAD6 and as having relevance to the process of RDB and resistance to DNA damage. The presence of a functional RAD18 is essential for this (mono)ubiquitination (25). Mammals contain two RAD6-homologous genes, designated *HR6A* and *HR6B*, encoding 152-amino-acid proteins that are structurally and functionally highly conserved; the proteins share approximately 70% sequence identity with yeast RAD6 and around 95% identity with each other (33). The mouse homologs *mHR6A* and *mHR6B* are 100% identical to the human proteins *hHR6A* and *hHR6B*, respectively. Both mammalian homologs are ubiquitously expressed and are able to complement a *rad6* mutant strain for repair and mutagenesis functions but not for the sporulation defect which appears to depend on the acidic tail that is present only in yeast RAD6 (31, 53).

Previously, the generation and characterization of *mHR6B* knockout mice, the first animal model defective in the ubiquitin-conjugating pathway, were described (55). The *mHR6B* knockout mice show a male-limited sterility phenotype, with spermatogenic derailment which is most pronounced during

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postmeiotic condensation of chromatin in elongating spermatids (55). This step in spermatogenesis coincides with histone-to-protamine replacement, essential for normal sperm development. As was shown, the level of HR6B exceeds that of HR6A in spermatids in rats and mice, which is in agreement with an important role in postmeiotic cells (6, 32). In addition, a defect in the meiotic prophase in spermatocytes of *mHR6B* knockout male mice has been identified (6).

In the present report, we describe the effect of using gene targeting in mice to inactivate the other *RAD6* homolog, *mHR6A*. In addition, we have evaluated the viability of mice with different numbers of intact *mHR6A* and *mHR6B* alleles. The results obtained are discussed with respect to the specificity and redundancy of the two encoded ubiquitin-conjugating enzymes.

MATERIALS AND METHODS

Isolation of cDNAs and genomic clones. A 1.7-kb human HR6A cDNA fragment (33) was used to screen a λ ZAP mouse brain library (Stratagene, La Jolla, Calif.). Out of a number of positive clones, we selected one (designated B4) that contained an insert of approximately 1.7 kb and resembled the major transcript detectable in all tissues (32). This cDNA was sequenced by using T7 polymerase (Amersham Biosciences, Piscataway, N.J.) according to the manufacturer's guidelines. An EcoRI subclone encompassing the first 301 nucleotides was subsequently used to screen an EMBL-3 λ phage-based 129Sv mouse genomic CCE library. Two genomic clones were isolated. These clones were designated X2 and X4, and both contained the complete gene.

Construction of the targeting vector and disruption of *mHR6A* in ES cells. A 5.26-kb ScaI fragment containing exons 1 to 3 was selected as the starting point for a targeting construct and cloned in pIC20R (43). The resulting plasmid was digested with SalI and BamHI, and a cassette containing the neomycin resistance gene expressed under the control of the thymidine kinase promoter (69) was inserted as a XhoI/BamHI fragment, thereby replacing the coding part of exon 1 with the dominant selectable marker (Fig. 1A). The resulting plasmid was digested with Asp718I and EcoRV, and the insert was recovered from agarose by electrophoresis. Approximately 10 μ g of DNA was electroporated into E14 embryonic stem (ES) cells (a kind gift of A. Berns, NKI, The Netherlands) as described previously (81). Individual neomycin-resistant clones were picked and expanded. Genomic DNA was isolated as described previously (39), digested with EcoRI, and screened for homologous integration by Southern blot analysis.

Generation of *mHR6A*-deficient mice and fibroblasts. Individual ES clones with the correct karyotype and a correctly integrated *mHR6A*-targeting construct were injected into C57BL/6J blastocysts by using standard procedures (9). Male chimeras were bred with wild-type female FVB/n animals, and germ line transmission was recognized on the basis of a gray coat color of the offspring. Tail DNA was isolated as described previously (39), digested, and analyzed by Southern blotting. Starting with the F₂ generation, animals were genotyped by multiplex PCR. With primers *mHR6A.7B* (5'-CTCGGCTCCGGAGCGGCAGT-3') and *mHR6A.3B* (5'-CCCTCATGAGGCGGCCGG-3'), a 153-bp fragment representing the wild-type allele was amplified. Primers *mHR6A.7B* and *NEOAS* (5'-TAGCCGAATAGCCTCTCCACCC-3') resulted in the amplification of a 462-nucleotide fragment representing the targeted allele. Primary mouse embryonic fibroblasts (MEFs) were isolated as described elsewhere (71). The UV sensitivity of MEFs was determined by assaying the incorporation of [³H]thymidine at various doses of UV (61).

RNA isolation and RT-PCR. Total RNA was isolated from testes by using LiCl-urea (4), and cDNA was made by using 20 U of AMV-reverse transcriptase (RT) (Promega Benelux, Leiden, The Netherlands) with random hexamer primers (Roche Diagnostics, Almere, The Netherlands) according to the manufacturer's protocol in the presence of 40 U of RNAsin (Promega). *mHR6A* cDNA was amplified by using forward primer *mHR6A.7B* in exon 1 and reverse primer *mHR6A.10* (5'-TCATAGGTTGGACTCCAACGG-3') in exon 5.

Gel electrophoresis and immunoblotting. Two-dimensional gel electrophoresis was done essentially as described by O'Farrell et al. (49). In short, oocytes or ovaria were sonicated in buffer A (0.5% [wt/vol] sodium dodecyl sulfate, 4 M urea, 1 mM dithiothreitol [DTT], 1% [vol/vol] Nonidet P-40, 0.1% [vol/vol] Bio-Lyte [pH 3 to 5; Bio-Rad, Veenendaal, The Netherlands], 0.1% [vol/vol] Bio-Lyte [pH 4 to 6; Bio-Rad]) and spun at 4°C at 13,000 rpm for 10 min, and the supernatant was collected. The protein concentration was determined by using

the bichinchoninic acid assay (Pierce, Rockford, Ill.). Protein lysates to be separated in the first dimension were diluted in solution B (9 M urea, 1% [vol/vol] Bio-Lyte [pH 3 to 5], 1% [vol/vol] Bio-Lyte [pH 4 to 6], 1% Nonidet P-40, 100 mM DTT). Samples were boiled for 3 min and cooled on ice. Samples (40 μ l) containing approximately 10 μ g of protein were loaded on gels consisting of 4% acrylamide-bisacrylamide in solution B without DTT in 12-cm glass tubes. Samples were overlaid with 10 μ l of overlay buffer (8 M urea, 1% [vol/vol] Bio-Lyte [pH 3 to 5], 1% [vol/vol] Bio-Lyte [pH 4 to 6]) and run for 16 h at 175 V in a Bio-Rad 175 tube cell (Bio-Rad) with 0.02 M NaOH in the lower chamber (cathode) and 0.01 M H₃PO₄ in the upper chamber (anode). After the first dimension was run, gels were removed from the glass tubes and either directly used for second-dimension polyacrylamide gel electrophoresis or stored at -20°C. Approximately 2 cm of the gel was cut off at the acidic side, and the rest was equilibrated in sample buffer (10% [vol/vol] glycerol, 2% sodium dodecyl sulfate, 10 mM DTT, 0.01% bromophenol blue, 62.5 mM Tris-HCl [pH 6.8]). Equilibrated gels were placed on top of a 15% phosphonoacetic acid gel with a 4% stacking layer and electrophoresed by using the Mini Protean II system (Bio-Rad, Hercules, Calif.). Subsequently, proteins were blotted on nitrocellulose (0.45 μ m; Schleicher & Schull, Dassel, Germany), and *mHR6A* and *mHR6B* were detected by using α -HR6A/B antibody as described previously (5).

Superovulation of female mice and isolation of oocytes and embryos. To isolate unfertilized oocytes for biochemical analysis, follicle growth in female mice was induced by intraperitoneal injection of 5 IU of Folligonan (Intervet, Boxmeer, The Netherlands). Forty-eight hours later, the mice were killed by cervical dislocation and the ovaries were isolated. Isolated ovaries were put in 100- μ l drops of IVF medium [98.37 mM NaCl, 4.56 mM KCl, 0.69 mM MgSO₄ · 7H₂O, 22.54 mM NaHCO₃, 0.86 mM NaH₂PO₄ · 2H₂O, 0.085% (wt/vol) glucose, 0.0581% (wt/vol) Ca-L(+) -lactate, and 0.1485% (wt/vol) powdered F-10 supplemented with pasteurized plasma solution (final concentration, 3.5 g/liter)] and extensively punctured with two 30-gauge needles. Oocytes were collected and washed once in fresh IVF medium, pelleted by a quick spin in a table centrifuge, snap frozen, and stored at -80°C pending further use.

To obtain fertilized oocytes and preimplantation embryos, female mice were injected intraperitoneally with 5 IU of Chorulon (Intervet) 48 h after Folligonan injection and housed overnight with male mice. One-cell embryos (fertilized oocytes up to and including the zygote stage), two-cell embryos, and blastocysts were isolated from the oviduct or the uterus at 0.5, 1.5, and 3.5 days postcoitum (dpc) (E0.5, E1.5, and E3.5), respectively, as described previously (27). E1.5-stage embryos were placed in 50- μ l droplets of IVF medium under mineral oil in a 5% CO₂ incubator at 37°C and cultured for 72 h.

Immunofluorescence. One-cell-stage embryos were isolated as described above, collected in phosphate-buffered saline (PBS), fixed in 4% paraformaldehyde in PBS for 8 min at room temperature, and subsequently permeabilized in 0.5% Triton X-100 in PBS for 8 min at room temperature. To reduce background fluorescence, embryos were treated with 1% [wt/vol] bovine serum albumin in PBS-0.5% Tween 20 (blocking reagent) for 30 min at room temperature. Primary antibody incubations in blocking reagent were carried out overnight at 4°C, followed by three washes in blocking reagent at room temperature for 10 min. Secondary antibodies were incubated for 2 h at room temperature in blocking reagent, followed by three 10-min washes in blocking reagent at room temperature in the dark. Embryos were mounted on slides in Vectashield (Vector Laboratories, Burlingame, Calif.) containing DAPI (4',6'-diamidino-2-phenylindole). Images were taken with a fluorescence microscope (Axioplan 2; Carl Zeiss, Jena, Germany) equipped with a digital camera (Coolsnap-Pro; Photometrics, Waterloo, Canada).

Confocal imaging. Images were made with a confocal scanning laser microscope (Zeiss LSM510NLO; Carl Zeiss). During the observation period, fertilized oocytes were kept in a drop of IVF medium on a round cover glass mounted in a special chamber. Observations were made by using a \times 63 oil immersion lens, and fertilized oocytes were imaged in optical planes with a 2- μ m distance. DNA was made visible by using the DNA vital stain Syto16 (Molecular Probes Europe BV, Leiden, The Netherlands) added to the IVF medium. A 488-nm argon laser was used for excitation, and the emission was measured with a band-pass filter between 500 and 550 nm. Images were obtained by using the LSM510 image software (Carl Zeiss).

Nucleotide sequence accession number. The B4 HR6A cDNA sequence was submitted to GenBank/EMBL under accession no. AF089812.

RESULTS

Isolation and characterization of mouse *mHR6A* cDNA. To isolate the *mHR6A* cDNA, we screened a mouse brain library

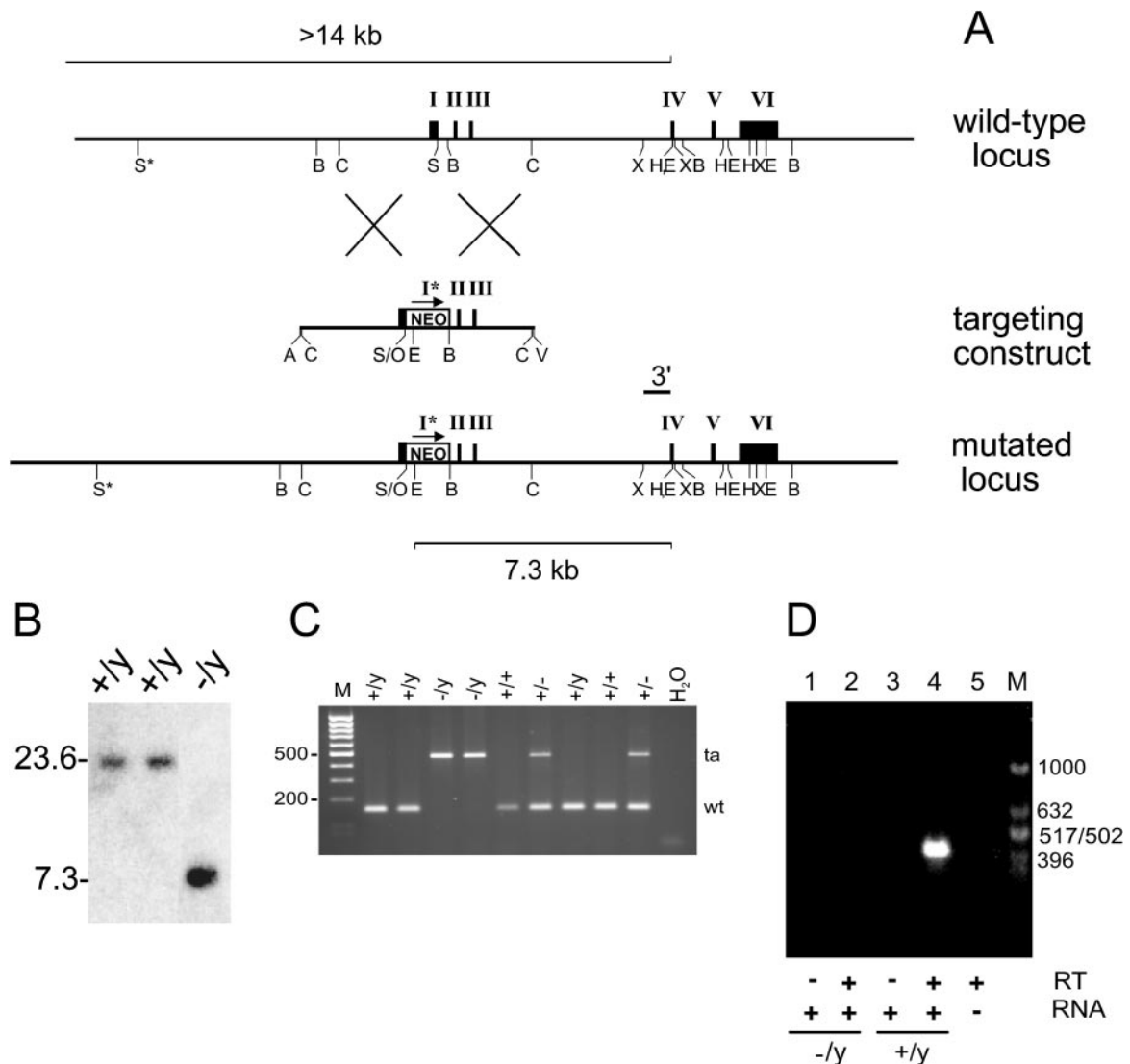


FIG. 1. Targeted disruption of the *mHR6A* gene by homologous recombination. (A) Genomic organization and disruption strategy for *mHR6A*; shown are the gene, the targeting construct, and the targeted *mHR6A* allele. The *neo* cassette is inserted between the SalI site of exon 1 and the BamHI site in the first intron, introducing a diagnostic EcoRI site. Note that the insertion of the dominant marker disrupts the gene immediately behind the ATG translation initiation codon and deletes the coding part of exon 1. Shown are the relevant restriction sites (E, EcoRI; S, SalI; B, BamHI; C, ScaI; X, XmnI; H, HinDIII; O, XhoI; A, Asp718I; V, EcoRV). The position of the 3' probe for Southern blot analysis is indicated above the mutated locus. Lines at the top and bottom indicate the estimated lengths of the fragments detected in Southern blot analysis of EcoRI-digested DNA. Roman numerals mark the exons. The 3' deletion of exon 1 in the targeted allele is indicated with an asterisk. (B) Southern analysis of EcoRI-digested DNA from three neomycin-resistant ES clones after hybridization with the 3' probe. The positions of the wild-type allele (23.6 kb) and the targeted allele (7.3 kb) are indicated. Note that the male ES line contains only one X-chromosomal *mHR6A* gene copy. “-/*y*” indicates a homologous recombinant ES clone, and “+/*y*” indicates a wild-type ES clone. The difference in signals between the targeted band and the wild-type band is presumably due to difficulty in the transfer of large fragments. (C) PCR-based genotyping of nine littermates. Primers *mHR6A.7B* and *mHR6A.3B* amplify a 152-bp fragment representing the wild-type allele (wt), and primers *mHR6A.7B* and *NEOAS* amplify a 473-nucleotide fragment representing the targeted allele (ta). Note that male animals (+/*y* and -/*y*) possess only one allele due to the X-chromosomal location of the allele. (D) RT-PCR with total mouse testis RNA (lanes 1 to 4) from an *mHR6A*^{-/*y*} animal (lanes 1 and 2) and an *mHR6A*^{+/*y*} male littermate (lanes 3 and 4). Reactions were performed in the absence (lanes 1 and 3) or in the presence (lanes 2, 4, and 5) of RT. Lane 5 contains no RNA. M indicates the marker lane, with fragment lengths indicated on the right.

with human cDNA as a probe. A mouse cDNA fragment of ~1.7 kb, probably representing the major transcript (32), was isolated. Sequencing revealed 73% sequence identity with the human transcript overall and 93% sequence identity in the putative open reading frame (ORF). The encoded gene product, 152 amino acid residues long as translated from the sub-

mitted mouse sequence (GenBank accession no. AF089812), was 100% identical in sequence to the protein specified by the human cDNA from the public expressed sequence tag and human genome project databases, underlining the importance of this protein. In contrast to the 3' untranslated regions of *mHR6B* and *hHR6B*, no stretch of conserved nucleotides

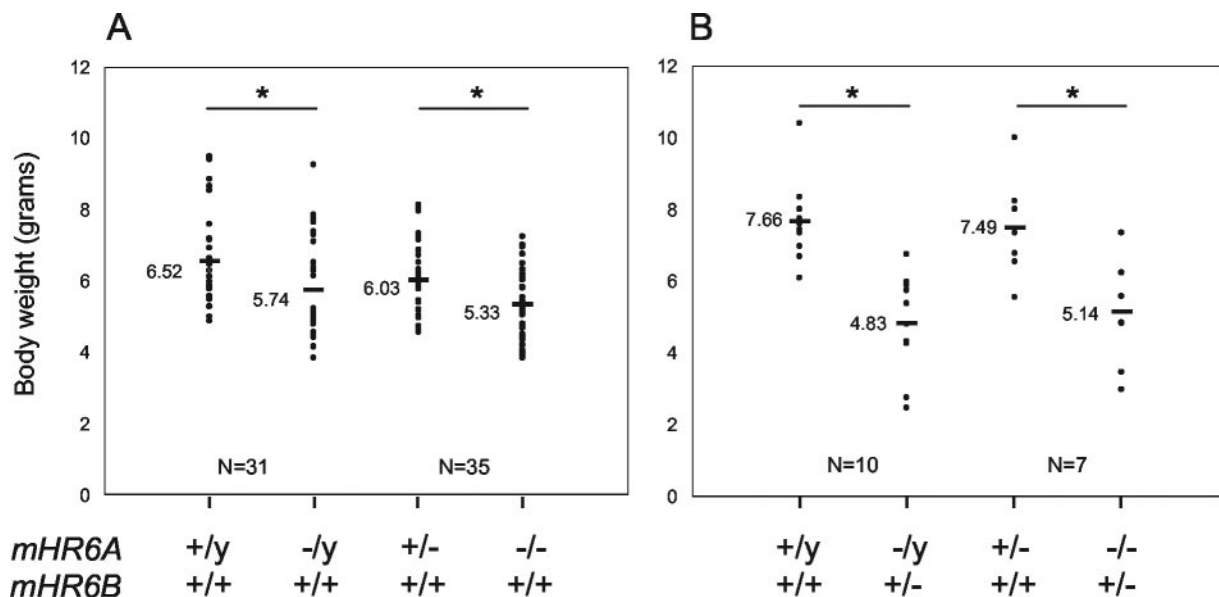


FIG. 2. Paired *t* test of the difference between total body weights of mice with a specific genotype and control littermates 10 ± 1 days after birth. Each dot represents the average body weight of all animals with a specific genotype present in one litter. The mean body weights are indicated by horizontal bars. The asterisks indicate statistically significant differences between the indicated groups ($P \leq 0.05$). N is the number of litters used in the analysis. (A) Total body weights of *mHR6A*-deficient animals and control littermates. Included are litters that contain both *mHR6A*^{+/y} and *mHR6A*^{-/y} male mice and/or both *mHR6A*^{+/-} and *mHR6A*^{-/-} female mice. (B) Total body weights of mice with one single functional *mHR6B* allele and control littermates. Included are only litters that contain both *mHR6A*^{+/y}/*mHR6B*^{+/+} and *mHR6A*^{-/y}/*mHR6B*^{+/+} male animals and/or both *mHR6A*^{+/-}/*mHR6B*^{+/+} and *mHR6A*^{-/-}/*mHR6B*^{+/+} female animals. The genotypes of both genes are indicated under the x axis.

was present in the 3' untranslated regions of *mHR6A* and *hHR6A*, suggesting that this is a specific feature of the *HR6B* gene (55).

Isolation of genomic sequences and targeting of ES cells.

Two overlapping clones of mouse strain 129Sv DNA, one of which contained the complete gene as outlined in Fig. 1A, were isolated. The gene spans a region of ~ 10 kb and is composed of six exons. A 5.36-kb ScaI genomic fragment that was present in both isolated *mHR6A* clones was selected. This fragment contains the first three exons, including the start codon ATG of the ORF. A dominant-selectable neomycin resistance marker was inserted by using the Sall and the BamHI restriction sites. In this way, the ORF was disrupted immediately after the start codon (Fig. 1A). In addition, amino acids 3 to 14 were deleted, thereby strongly reducing the possibility that an aberrant protein might be formed. Transfection of the targeting construct by electroporation and selection for stable integration of the dominant marker resulted in a frequency of targeted, single integrations of 2.8% (4 out of 146 analyzed), as confirmed by Southern blot analysis. Hybridization with the 3' external probe visualized a large EcoRI fragment of 23.6 kb in the case of the wild-type allele and a 7.3-kb EcoRI fragment for a targeted allele (Fig. 1B).

Generation of *mHR6A*-deficient mice. Two of the targeted ES clones (clones 49 and 102) had a correct karyotype and were used for injection into C57Bl/6 blastocysts to generate chimeras. Four chimeras were obtained, but only one (from clone 49) transmitted the mutated allele as determined by coat color and Southern blot analysis with the 3' probe (data not shown). Because the *mHR6A* gene, like the human ortholog, is located on the X chromosome (34, 56), the male chimera transmits the targeted allele only to female pups of the F₁

offspring. Indeed, all of the F₁ females that had the coat color associated with the 129Sv/FVB hybrid genomic background were heterozygous.

Next, the F₁ males and females were bred, resulting in a large number of knockout (*mHR6A*^{-/y}) male mice in the F₂ offspring. The absence of a functional *mHR6A* allele in these animals was confirmed by RT PCR with total RNA isolated from testis material of an *mHR6A*^{-/y} animal and a wild-type animal. No *mHR6A* transcripts spanning the (partially) deleted exon 1 could be detected in testis cDNA from the *mHR6A*^{-/y} mouse, while a product of the right size (420 nucleotides) was readily amplified from testis cDNA of the wild-type male (Fig. 1D). This finding indicates that targeted integration of the knockout construct resulted in the absence of the original transcripts containing the complete ORF as well as any alternatively spliced mRNAs.

***mHR6A*-deficient mice and embryonic fibroblasts.** A program of breeding between *mHR6A*^{+/-} females and *mHR6A*^{-/y} males was set up. The offspring showed a Mendelian representation of knockout and heterozygous female animals (91 and 91 animals, respectively) but a significant underrepresentation of knockout male animals compared to wild-type males (71 and 116 animals, respectively) (χ^2 test, $P = 0.001$).

To compare the body weights of the different genotypes, litters that contained animals with both male and/or both female genotypes were included in a paired *t* test analysis. Mice were weighed at the time of tailing (10 ± 1 days after birth). The average body weight of all animals with the same genotype and sex within one litter was determined and included in the analysis. On average, the body weight of knockout male and female mice at 10 days of age was significantly lower (paired *t* test, $P < 0.01$), showing a reduction of approximately 12%

TABLE 1. Average numbers of pups per litter with various parental *mHR6A* combinations

Parental <i>mHR6A</i> genotypes	No. of successful matings	Litter size (mean no. of pups \pm SD)
+/+ \times +/Y	ND ^a	
+/+ \times -/Y	7	5.9 \pm 2.6
+/- \times +/Y	35	8.7 \pm 2.6
+/- \times -/Y	79	8.6 \pm 2.7
-/- \times +/Y	11	0
-/- \times -/Y	13	0

^a ND, not determined.

compared to that of the respective control littermates (Fig. 2A).

To investigate the possible occurrence of pathological abnormalities, most organs of three adult knockout animals and three control littermates were fixed and hematoxylin and eosin stained, but no aberrations were found (data not shown).

In view of the role of the RAD6 protein in RDB in yeast, *mHR6A*-deficient MEF cell lines were tested for their growth rates and cellular survival after irradiation with different doses of UV, as measured by [³H]thymidine incorporation. No differences between *mHR6A*-deficient and *mHR6A*-proficient cell lines were observed (data not shown). This finding demonstrates that UV resistance in *mHR6B*-deficient cells (55) does not depend on a functional *mHR6A* allele but is presumably caused by redundant activities of *mHR6A* and *mHR6B* in somatic cell types.

***mHR6A*^{-Y} males are normally fertile, but *mHR6A*^{-/-} females are not able to produce offspring.** In contrast to the phenotype observed in *mHR6B* knockout mice, whereby females are normally fertile and males are sterile, we observed the reverse for *mHR6A*-deficient animals; *mHR6A*^{-Y} male mice were normally fertile, as shown by their ability to sire litters (Table 1) as well as by a precise analysis of male reproductive parameters (data not shown), but *mHR6A*^{-/-} female mice appeared infertile. We investigated whether these females were receptive to mating with both *mHR6A*^{-Y} and wild-type males. Successful matings, indicated by the presence of copulatory plugs, were observed in *mHR6A*^{-/-} females regardless of which males were used. The numbers of pups per litter that resulted from the breeding of heterozygous females with wild-type and knockout males were not significantly different ($P = 0.81$, two-tailed Student's *t* test). Ovaries were then isolated from knockout and heterozygous littermates, fixed, sectioned, stained, and microscopically analyzed for the presence of the various stages of follicular development. Ovaries from *mHR6A*^{-/-} females appeared indistinguishable from those of *mHR6A*^{+/-} littermates, with the presence of a normal number of growing follicles and corpora lutea (data not shown). This finding indicates that the ovaries of *mHR6A*-deficient females are capable of follicular development and ovulation. To follow up on the initial observation, we superovulated *mHR6A*^{-/-} and *mHR6A*^{+/-} females and housed them overnight with FVB/n wild-type males. Fertilized oocytes were recovered from the oviducts and cultured in vitro. The one-cell embryos recovered from *mHR6A*^{-/-} females appeared morphologically normal and comparable with the embryos isolated from the heterozygous control females. All of the embryos contained two pronuclei (Fig. 3A to D), indicating

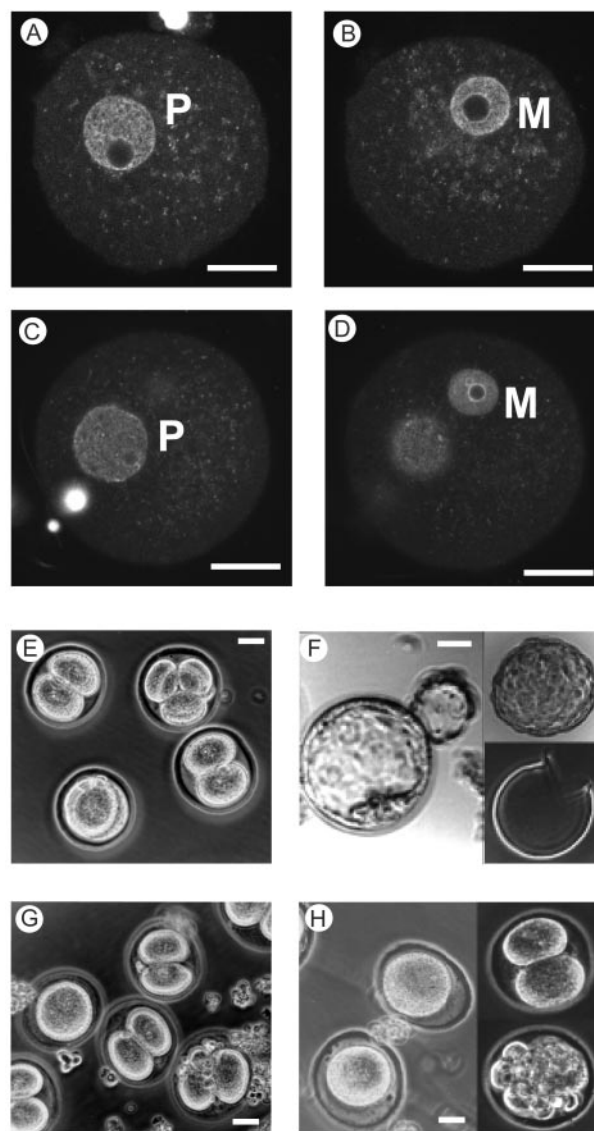


FIG. 3. Developmental stages of preimplantation embryos obtained from *mHR6A*^{+/-} (A, B, E, and F) and *mHR6A*^{-/-} (C, D, G, and H) female mice fertilized by wild-type males. Images obtained with confocal laser scanning microscopy (A to D) show embryos from an *mHR6A*^{+/-} heterozygous mother (A and B) and an *mHR6A*^{-/-} knockout mother (C and D) approximately 8 h after fertilization. Two planes are displayed for the fertilized egg, one showing the paternal pronucleus (P in panels A and C) and one showing the maternal pronucleus (M in panels B and D). Phase-contrast microscopy shows examples of embryos obtained from *mHR6A*^{+/-} (E and G) and *mHR6A*^{-/-} (G and H) female mice at 1.5 dpc (E and G) and cultured for 3 days (F and H). (E) Embryos at various stages of development, obtained from an *mHR6A*^{+/-} female. (F) Hatching (left) and hatched (upper right) embryos and an empty zona pellucida (lower right). (G) Embryos obtained from an *mHR6A*^{-/-} female. (H) One-cell (left), two-cell (upper right), and fragmented (lower right) embryos. Scale bar, 20 μ m.

that they were normally fertilized and did not develop parthenogenetically after fertilization with wild-type sperm. The one-cell embryos recovered from *mHR6A*^{+/-} and *mHR6A*^{-/-} females developed up to the two-cell stage in vitro but were not able to overcome a two-cell-stage block. This situation is not unusual for noninbred strains (22) but prevented us from de-

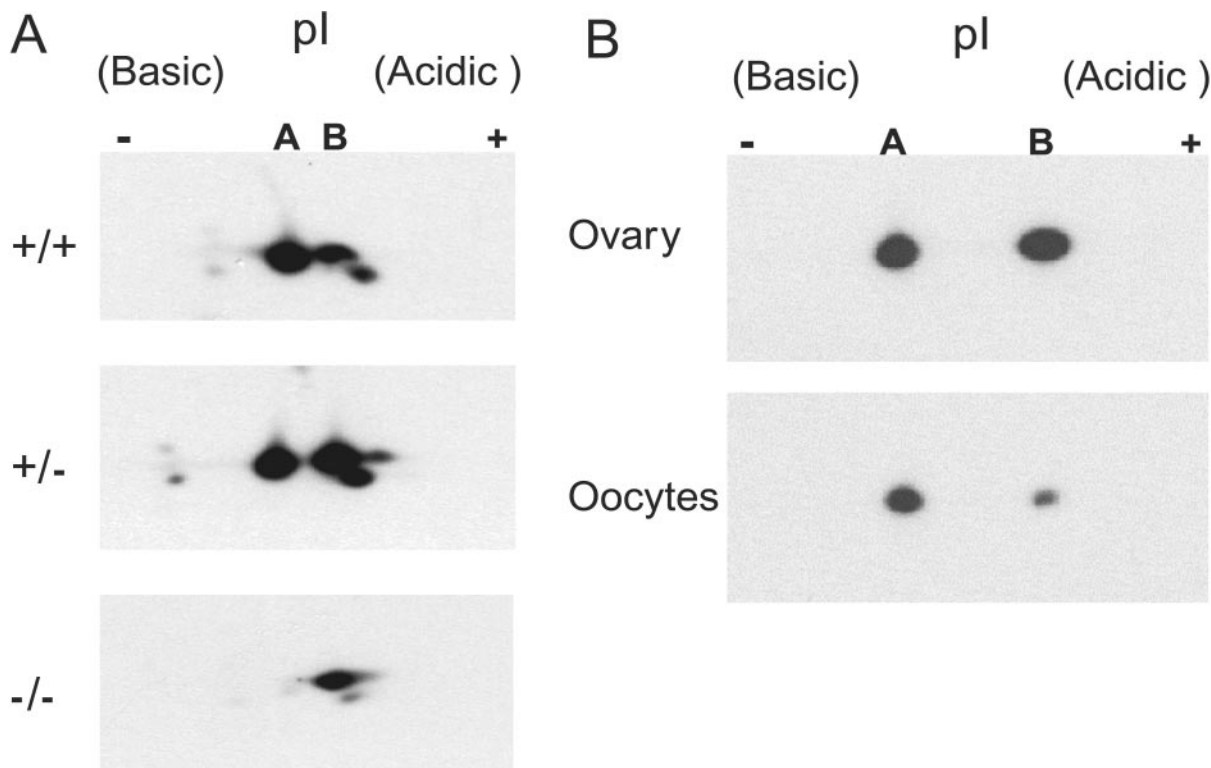


FIG. 4. mHR6A and mHR6B protein expression. Total protein extracts were separated by using two-dimensional gel electrophoresis. (A) Comparison of the amounts of mHR6A and mHR6B proteins (indicated by "A" and "B," respectively) in ovaries from $mHR6A^{+/+}$, $mHR6A^{+/-}$, and $mHR6A^{-/-}$ mice. (B) Expression of mHR6A and mHR6B in oocytes isolated from wild-type mice compared to the amounts of the two proteins in whole ovaries.

terminating the end point of the development of embryos derived from an $mHR6A^{-/-}$ mother. Therefore, embryos were recovered by flushing the oviducts 1.5 dpc, when developing embryos are at the two-cell stage, and subsequently these embryos were cultured for three additional days before reexamination. Total numbers of 37 two-cell embryos and 1 four-cell-stage embryo were recovered from four $mHR6A^{+/-}$ females, and all of these embryos appeared normal (Fig. 3E). All two-cell and four-cell embryos developed into blastocysts, most of which started hatching, and some of these embryos were already completely hatched, leaving behind an empty zona pellucida (Fig. 3F). From four $mHR6A^{-/-}$ females, 13 phenotypically normal two-cell-stage embryos, 2 two-cell-stage embryos with cytoplasmic fragments, 11 fertilized oocytes (of which 5 showed signs of fragmentation), and 1 three-cell embryo were retrieved (Fig. 3G). None of these embryos isolated from the oviducts of the $mHR6A^{-/-}$ females grew out to the blastocyst stage. Eight of them reached or remained in the two-cell stage, and others remained unicellular or showed severe cytoplasmic fragmentation and degeneration (Fig. 3H). This result was not a culture artifact, since embryos isolated at 3.5 dpc from $mHR6A^{-/-}$ female mice were also found to have developed no further than the two-cell stage (data not shown). This finding points to an important role of maternal mHR6A in the very early stages of embryonic development.

The mHR6A protein level is high compared to that of mHR6B in oocytes. In a number of tissues and cell types that were analyzed, the levels of mHR6A and mHR6B were com-

parable (32). In the testis, the postmeiotic cells, the spermatids, showed a relative increase of mHR6B compared to mHR6A (6, 32). This increase might be caused by the inactivation of the X-chromosomal $mHR6A$ gene in spermatogenesis, when the X and Y chromosomes form the heterochromatic sex body during the meiotic prophase (62). In oogenesis, the two X chromosomes are transcriptionally active during the meiotic prophase, which might result in a relatively high level of mHR6A compared to mHR6B (35). To test this possibility, we first prepared protein extracts from ovaries of wild-type and $mHR6A$ heterozygous and knockout mice. The protein contents of these samples were separated by two-dimensional gel electrophoresis and blotted, and mHR6A and mHR6B proteins were visualized by using the α -HR6A/B antibody as described (5). This analysis confirmed the absence of mHR6A in ovaries from $mHR6A^{-/-}$ females (Fig. 4A) but also showed that in the wild-type whole ovary, the levels of mHR6A and mHR6B were about equal. Differential expression of the two proteins in oocytes may go undetected when whole ovary tissue is analyzed. Hence, to determine the relative levels of mHR6A and mHR6B proteins in wild-type oocytes, oocyte maturation was induced in six female mice by injecting the mice with 5 IU of Folligonan. Approximately 240 oocytes were isolated, lysed, and analyzed by two-dimensional gel electrophoresis. It was found that the level of mHR6A in isolated oocytes was relatively high compared to that of mHR6B (Fig. 4B).

Methylation of histone H3 in pronuclei in wild-type and

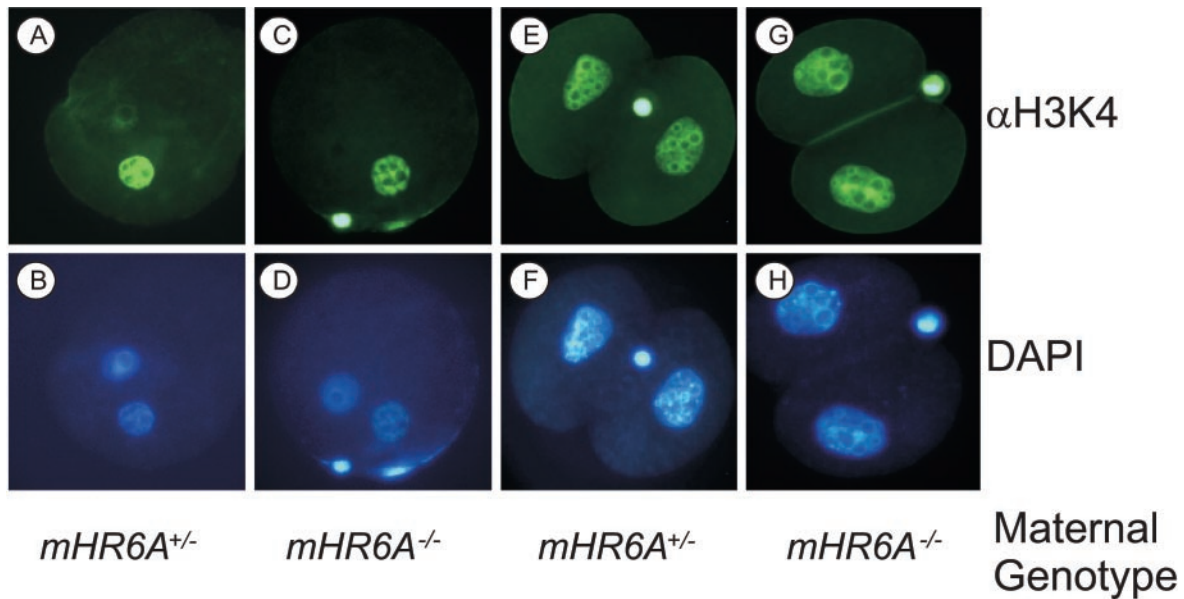


FIG. 5. Immunofluorescent images of one-cell (A to D) and two-cell (E to H) embryos stained with an antibody against methylated H3K4 (A, C, E, and G) and DAPI (B, D, F, and H). The genotype of the mother is indicated on the right. The intense bright spots in panels C to H are caused by the remnants of a polar body. Staining of only one of the two pronuclei with α -methylated H3K4 is observed (A and C), while both nuclei of the two-cell embryo stain equally intensely (E and G).

mHR6A-deficient oocytes. In the very early stages of mammalian development, dramatic epigenetic changes that are involved in spatial and temporal control of gene expression occur (3, 16, 45, 58). These changes include demethylation and de novo methylation of CpG dinucleotide sequences. In addition, much recent evidence shows that epigenetic histone modifications are tightly regulated to assure control of gene expression but also DNA repair and recombination (for a review, see reference 18). Recently, it was shown that *Saccharomyces cerevisiae* RAD6 plays an important role in the posttranslational modification of histones. Monoubiquitination of histone H2B at lysine 123 is a prerequisite for the methylation of lysine 4 of histone H3 (H3K4) (16, 65), a modification that is associated with active gene expression, cell growth, telomeric and ribosomal DNA silencing, and sporulation (10, 57, 64). In higher eukaryotes, H3K4 methylation is associated with active chromatin domains, whereas methylated H3K9 associates with heterochromatic subdomains and gene silencing (reviewed in reference 38). Using specific antibodies against methylated H3K4 and methylated H3K9, we analyzed the role of mHR6A in controlling the methylation status of histone H3 during fertilization and early embryonic development. Embryos were isolated approximately 12 and 36 h postcoitum from *mHR6A*^{-/-} and *mHR6A*^{+/-} females. At 12 h postcoitum, immunofluorescent staining for H3K9 was predominantly associated with the maternal pronucleus (data not shown), a finding that is in full agreement with earlier data (3, 13). Two-cell embryos, isolated 36 h postcoitum, show clear and overall staining of the nuclei of both blastomeres irrespective of the genotype of the mother (data not shown). This finding suggests that H3K9 methylation is not severely affected by the absence of mHR6A during the formation of the first two blastomeres. In a similar experiment, it was observed that methylated H3K4 was also predominantly associated with the maternal pronucleus in one-cell embryos

obtained from *mHR6A*^{+/-} females (Fig. 5A), and no overt difference was observed for one-cell embryos from the *mHR6A* knockout females (Fig. 5C). After the first cell division, both nuclei fluoresce with similar intensities irrespective of the genotype of the mother (Fig. 5E and G). Immunofluorescent staining of one- and two-cell embryos for histone H3 clearly demonstrates the presence of histones in both pronuclei at this stage (data not shown). These data suggest that there are no marked changes in the methylation status of histone H3 in early embryos that can be attributed to the absence of maternal mHR6A.

Mice with losses of *mHR6A* and *mHR6B* alleles. The severe, pleiotropic phenotype of a *rad6*-null yeast strain (for a review, see reference 40), compared to the relatively mild phenotype that results from either mHR6A or mHR6B deficiency, suggests a strong functional redundancy between these two proteins. To investigate the effect of the complete loss of both RAD6 homologs in mice, a breeding program was set up to obtain double-knockout (DKO) animals. In this breeding program, double heterozygosity has no gross deleterious effect on embryonic development, and *mHR6A*^{+/-}/*mHR6B*^{+/-} female mice were mated with *mHR6A*^{+/-}/*mHR6B*^{+/-} males. As shown in Table 2, no DKO animals were found, whereas approximately 11 DKO mice were to be expected (χ^2 test, $P < 0.001$). In addition, female *mHR6A*^{+/-}/*mHR6B*^{-/-} animals were also absent. This result is most likely due to random inactivation of the X chromosome early in development, thereby leaving only 50% of the cells in these mHR6B-deficient mice with one functional wild-type X-chromosomal *mHR6A* allele.

With the exception of *mHR6A*^{-/-}/*mHR6B*^{-/-} animals (no functional allele) and the *mHR6A*^{+/-}/*mHR6B*^{-/-} mice (only one intact allele, which is subject to X inactivation), all of the other genotypes were represented in the offspring. Mice

TABLE 2. Possible genotypes and observed numbers of pups with genotypes for various parental combinations

Genotype	Observed (expected) no. of pups with genotype for parental combination ^a			
	A ^{+/Y} /B ^{+/-} × A ^{+/-} /B ^{+/-}	A ^{-Y} /B ^{+/+} × A ^{+/-} /B ^{+/-}	A ^{-Y} /B ^{+/-} × A ^{+/-} /B ^{+/+}	A ^{-Y} /B ^{+/-} × A ^{+/-} /B ^{+/-}
A ^{+/+} /B ^{+/+}	11 (11)			
A ^{+/+} /B ^{+/-}	29 (22)			
A ^{+/+} /B ^{-/-}	13 (11)			
A ^{+/-} /B ^{+/+}	11 (11)	75 (59)	7 (9)	30 (17)
A ^{+/-} /B ^{+/-}	20 (22)	70 (59)	11 (9)	56 (34)
A ^{+/-} /B ^{-/-}	0 (11)			0 (17)
A ^{-/-} /B ^{+/+}		61 (59)	8 (9)	16 (17)
A ^{-/-} /B ^{+/-}		16 (59)	3 (9)	20 (34)
A ^{-/-} /B ^{-/-}				0 (17)
A ^{+Y} /B ^{+/+}	12 (11)	87 (59)	11 (9)	27 (17)
A ^{+Y} /B ^{+/-}	37 (22)	83 (59)	15 (9)	54 (34)
A ^{+Y} /B ^{-/-}	17 (11)			29 (17)
A ^{-Y} /B ^{+/+}	16 (11)	60 (59)	10 (9)	24 (17)
A ^{-Y} /B ^{+/-}	11 (22)	23 (59)	6 (9)	11 (34)
A ^{-Y} /B ^{-/-}	0 (11)			0 (17)
Total	177	475	73	267

^a A, *mHR6A*; B, *mHR6B*. Boldface numbers are significantly lower than the expected numbers.

lacking an intact *mHR6A* allele and being heterozygous for *mHR6B* (*mHR6A*^{-Y}/*mHR6B*^{+/-}), however, were clearly underrepresented (χ^2 test, $P < 0.025$) and showed a pronounced decrease in body weight that was already visible at birth. To investigate this result in more detail, *mHR6A*^{+/-}/*mHR6B*^{+/-} females were mated with *mHR6A*^{-Y}/*mHR6B*^{+/+} males and the offspring were genotyped and weighed. Mice that contained only one wild-type *mHR6B* allele were significantly underrepresented in the offspring (χ^2 test, $P < 0.001$) (Table 2). To quantify the loss of body weight, we compared the body weights of wild-type versus *mHR6A*^{-Y}/*mHR6B*^{+/-} males and those of *mHR6A*^{+/-}/*mHR6B*^{+/+} versus *mHR6A*^{-/-}/*mHR6B*^{+/-} females. Only litters that contained pups with both male and/or both female genotypes were included. The average body weights of *mHR6A*^{-Y}/*mHR6B*^{+/-} male and *mHR6A*^{-/-}/*mHR6B*^{+/-} female animals at 10 days of age were significantly lower (paired *t* test, $P < 0.01$) than those of the control littermates, with a reductions of 37 and 31%, respectively (Fig. 2B). Interestingly, *mHR6A*^{-Y}/*mHR6B*^{+/-} male animals were found to be normally fertile. The average number of pups per breeding sired by these males was not significantly lower than that of pups sired by *mHR6A*^{-Y}/*mHR6B*^{+/+} males ($P = 0.387$, two-tailed Student's *t* test) mated with *mHR6A*^{+/-}/*mHR6B*^{+/-} females (7.1 ± 2.7 and 7.6 ± 2.6 pups, respectively). This result indicates that, in the testis, the level of *mHR6B* protein from one intact and functional allele can compensate for the complete loss of *mHR6A* and/or that *mHR6B* has a specific function in spermatogenesis.

DISCUSSION

Over the last few years, a number of mammalian homologs of yeast RDB genes that exhibit a high degree of sequence and functional conservation have been identified, suggesting that RDB mechanisms might be conserved throughout eukaryotic evolution (21, 30, 33, 42, 44, 47, 68, 72, 78–80). However, some

remarkable functional differences between yeast and mammalian proteins (8, 17, 73, 76) and the identification of additional Y family DNA polymerases in mammalian species (for a review, see reference 19) forbid simple extrapolation from yeast to mammals. Also, the identification of two mammalian *RAD6* homologs provides a complicating factor. Previously, we demonstrated that the inactivation of the *mHR6B* gene, one of the two mammalian homologs of yeast *RAD6*, induces only a relatively mild phenotype in mice compared to that induced by a *rad6*-null allele in yeast. Whereas yeast *rad6* mutants display extreme sensitivity to all kinds of genotoxins, low growth rates, and impaired sporulation, the main phenotype of *mHR6B*^{-/-} mice is a severe derailment of spermatogenesis (55). This observation raised the question whether the second homolog, *mHR6A*, is responsible for the greater part of the somatic RAD6-like functions in mice or whether there is extensive functional redundancy between the two mammalian proteins.

RAD18 is the E3 that interacts with RAD6 in RDB in yeast. Both human RAD6 homologs can interact with human RAD18, and the overexpression of a dominant-negative human *RAD18* gene results in increased UV sensitivity (68, 79). Analysis of *mHR6A*-deficient MEFs, however, failed to reveal any difference between their response to UV irradiation and that of wild-type cells. This outcome is comparable with results of UV survival assays of *mHR6B*-deficient MEFs, which also showed no increased UV sensitivity (55). In addition, *mHR6A*-deficient mice appeared normal, apart from a slight reduction in body weight, and preliminary studies did not reveal an increased cancer predisposition up to 1 year of age.

Besides RAD18, UBR1 is another E3 capable of interacting with RAD6 (15). The human and mouse homologs of *UBR1* have been cloned, and their roles in the N-end rule pathway and interaction with *mHR6A* and *mHR6B* have been confirmed (36, 37). One of the observed phenotypes in *Ubr1*^{-/-} mice is a small but significant reduction in body weight (37). Similar reductions in body weight were observed for *mHR6A*-deficient animals (present data) and for *mHR6B*-deficient animals (unpublished data). *Ubr1*^{-/-} mice are, however, normally fertile. This finding indicates that the infertility phenotype of both *mHR6B*^{-/-} male and *mHR6A*^{-/-} female mice is not determined by the interaction of *mHR6A* and/or *mHR6B* with *Ubr1* and does not involve dysregulation of protein degradation via the N-end rule.

In the present breeding program, in addition to the absence of DKO mice, we found that *mHR6A*^{-/-}/*mHR6B*^{+/-} females and *mHR6A*^{-Y}/*mHR6B*^{+/-} males were underrepresented and runted and often died around birth. The *mHR6A*^{-Y}/*mHR6B*^{+/-} males that survived were normally fertile. This finding indicates that the level of *mHR6B* protein obtained from one functional *mHR6B* allele is sufficient to support spermatogenesis, even in the absence of *mHR6A* protein. On the one hand, the results obtained indicate that *mHR6A* and *mHR6B* have completely redundant activities both in somatic cell types and in germ line cells. On the other hand, the observed female and male infertility phenotypes in *mHR6A*- and *mHR6B*-deficient mice, respectively, indicate that germ line cells are subject to a dose-dependent effect which involves spermatogenesis- and oogenesis-specific differences in expression levels of *mHR6A* and *mHR6B*. However, the possibility that *mHR6A* and *mHR6B* have different functional properties

at the molecular level cannot be completely excluded. Several observations that point in this direction have been described. Ubiquitin-protein ligases (E3s) can be found among proteins that contain either HECT or RING motifs, such as RAD18 and UBR1 (reviewed in reference 51), but also among proteins with U-box domains. Hatakeyama et al. (23) showed that the U-box motif-containing cyclophilin CYC4/hCyP-60 needs the E2 enzyme HR6B, but not HR6A, for polyubiquitination. Cyclophilins are thought to have chaperone functions, being involved in the proper folding of protein substrates *in vivo*. Although ubiquitously expressed, transcript levels of cyclophilins are elevated and differ in size in the testis (74). This finding suggests the possibility that the spermatogenic derailment observed in *mHR6B*^{-/-} but not in *mHR6A*^{-/-} males is not caused by the relatively low level of mHR6A but by the inability of mHR6A to interact with cyclophilin. Additional data on possible specificity come from the identification of mHR6A, and not mHR6B, as an interacting partner of Rfp14 in a yeast two-hybrid screen (67). Rfp14 is present in maturing oocytes and remains expressed up to the eight-cell-stage embryo in the mouse. The protein shows the characteristics of a RING domain containing E3 ubiquitin ligase and can specifically interact with mHR6A. This protein could be useful in trying to identify the specificity-determining amino acid residues of mHR6A and mHR6B; mHR6A and mHR6B differ in only seven positions.

The most prominent effect of disrupting *mHR6A* is the observed complete failure of mHR6A-deficient females to produce offspring, resulting from developmental arrest after the first embryonic cell division. It is unlikely that the phenotype of the female *mHR6A*^{-/-} mice is due to aberrant levels of gonadotropins, since vaginal smears for mHR6A-deficient females revealed regular 4- to 5-day cycles and histological examination of fixed and stained ovary sections showed the presence of all stages of follicular development. In addition, the animals mated readily, the number of fertilized oocytes was similar to that for *mHR6A*^{+/-} littermates, and pseudopregnancy was induced. These data indicate a normal hypothalamic-hypophyseal-gonadal axis. Measurable deviations from normal serum gonadotropin levels would have resulted in dysregulated cycles.

The present results identify *mHR6A* as one of the few mammalian genes that encode a factor essential for the very first steps in embryonic development. Thus far, only *Mater* and *Zar1* have been shown to encode proteins that have their major, if not exclusive, functions in early postfertilization development (12, 70).

The mechanism that is disrupted by the absence of mHR6A is still unclear but could be related to the regulation of transcription and/or chromatin rearrangements. Just recently, it has started to become clear how, and what kind of, core histone tail modifications play a role in chromatin remodeling, higher-order chromatin structure, and transcription (see reference 63 and references therein). These modifications include phosphorylation, acetylation, methylation, and ubiquitination. Robzyk et al. (54) showed that the ubiquitination of histone H2B in yeast is dependent on a functional *RAD6* gene. Mutation of lysine residue 123 of H2B, which is essential for ubiquitination, prevents the methylation of H3K4 and induces

defects in sporulation and gene silencing. Other *RAD6*-dependent processes, such as RDB, appear unaffected (54).

During fertilization, the chromatin in the paternal pronucleus is reorganized, with histones of maternal origin replacing protamines. From that point on, acetylated forms of histone H4 are detectable in the paternal pronucleus, associated with the onset of transcription (1, 2). The methylation of histone H3 at lysine residue 9 (H3K9), known to be involved in the formation of stable repressive heterochromatin (50), was found in the maternal pronucleus but could hardly be detected in the paternal pronucleus (3, 13). In addition, H3K4 methylation, a hallmark of active chromatin (64), was also observed predominantly in the maternal pronucleus rather than the paternal pronucleus. The antibody used in the present study recognizes the dimethylated form of H3K4. The presence of trimethylated H3K4 has been shown to be a prerequisite for active genes, and the role of dimethylated H3K4 in mammalian cells is not clear, but the presence of dimethylated H3K4 may correlate with a permissive state of chromatin in which genes are potentially active (59, 60). The observed H3K4 staining in the maternal pronucleus may mark regions that will become active shortly after the pronuclear stage. However, we did not observe a change in the histone H3 methylation pattern in the pronuclei in the mHR6A-deficient oocytes during fertilization.

Both *mHR6A* and *mHR6B* cDNAs encode proteins that are able to induce H2B ubiquitination and subsequent methylation of H3K4 in *rad6* mutants (65), but there is a clear difference between ubiquitination of H2B in yeast and that in humans. In yeast, ubiquitinated H2B is most abundant and ubiquitinated H2A is undetectable, whereas in human somatic cells about 10% of H2A and only 1 to 1.5% of H2B is ubiquitinated (75). In spermatogenesis, no change in histone ubiquitination patterns has been detected in *mHR6B* knockout testis (5), but this result does not exclude the possibility of a low, but functionally significant, rate of H2A and/or H2B ubiquitination by mHR6B. Similarly, the possibility that the ubiquitin-conjugating enzyme mHR6A takes part in chromatin remodeling during early embryonic development through low rates of ubiquitination of H2A and/or H2B, which then might affect the methylation of histone H3 in a manner which has not been detected in the present experiments, cannot be excluded. Finally, it is important to note that the ubiquitination of H2B by *RAD6* in yeast appears to be unrelated to RDB mechanisms (54) and that the role of mHR6A as a maternal factor for early embryonic development might involve a molecular mechanism of action outside the context of DNA repair.

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