GC-1 mRHBDD1 knockdown spermatogonia cells lose their spermatogenic capacity in mouse seminiferous tubules

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Abstract

Background: Apoptosis is important for regulating spermatogenesis. The protein mRHBDD1 (mouse homolog of human RHBDD1)/rRHBDD1 (rat homolog of human RHBDD1) is highly expressed in the testis and is involved in apoptosis of spermatogonia. GC-1, a spermatogonia cell line, has the capacity to differentiate into spermatids within the seminiferous tubules. We constructed mRHBDD1 knockdown GC-1 cells and evaluated their capacity to differentiate into spermatids in mouse seminiferous tubules.

Results: Stable mRHBDD1 knockdown GC-1 cells were sensitive to apoptotic stimuli, PS341 and UV irradiation. In vitro, they survived and proliferated normally. However, they lost the ability to survive and differentiate in mouse seminiferous tubules.

Conclusion: Our findings suggest that mRHBDD1 may be associated with mammalian spermatogenesis.

Background

Spermatogenesis generates functional sperm cells from initially undifferentiated germ cells. This involves the proliferation of spermatogonia, meiosis of spermatocytes and the differentiation of spermatids into spermatozoa. It is a complex developmental program in which myriad events take place to ensure that the germ cells reach their proper stages of development at the appropriate times. Normal spermatogenesis requires a well-regulated balance of several processes, including cell proliferation, differentiation and apoptosis.

Apoptosis is a key phenomenon during spermatogenesis. For instance, an early, massive wave of germ cell apoptosis occurs at puberty. This event takes place during postnatal weeks 2 to 4 in mice, with a peak after 3 weeks [1-3]. It is estimated that 75% of spermatogenic cells undergo apoptosis during development [4,5], ensuring the maintenance of a critical ratio between maturing germ cells and Sertoli cells [2,6]. Sporadic apoptosis also occurs, primarily in spermatogonia and spermatocytes [2], eliminating defective germ cells with mutated DNA [7].
The Rhomboid family comprises polytopic membrane proteins, which may be the most widely-conserved membrane proteins identified to date [8]. They share conserved biochemical properties in all biological kingdoms. Rhomboid proteases, which have been well studied in Drosophila, appear to regulate EGF receptor signalling pathways, thereby controlling growth and development [9,10]. In addition, some yeast Rhomboid proteases have been found to play important roles in mitochondrial membrane remodelling [11], while some parasite proteases containing a Rhomboid domain are important for invasiveness [12].

Vertebrate rhomboid genes have been grouped into three classes: (1) active cellular rhomboids, including RHBDL1, RHBDL2, RHBDL4 and RHBD1 [13]; (2) inactive cellular rhomboids, including RHBDL5 and RHBDL6; (3) a mitochondrial rhomboid, PARL [14]. However, the physiological functions of these proteases remain to be clarified. Recently, RHBDL2 was found to cleave thrombomodulin and ephrin B3 [15,16]. An exciting finding in this field was the discovery of PARL (Presenilin-associated rhomboid-like), the function of which appears to be associated with apoptosis [17,18]. Drosophila Rhomboid 7 is believed to be associated with spermatogenesis, as male mutants tend to be afflicted with familial sterility [19].

mRHBD1 is the first Rhomboid serine protease known to be involved in mammalian spermatogenesis. We found that RHBD1, which contains a Rhomboid domain, is highly expressed in testes and is involved in regulating apoptosis [13]. In the present study, the roles of mRHBD1 in germ cell apoptosis and mouse spermatogenesis were investigated. We found that it is involved in regulating apoptosis in a mouse spermatogonia cell line, GC-1, and is essential for their capacity to differentiate into spermatids in mouse seminiferous tubules.

Results

mrbdd1 and rrhbdd1 are polytopic genes that are highly expressed in testis

mrbdd1 and rrhbdd1 were expressed in a variety of mouse and rat tissues, and the highest levels of gene products were found in the testis (Figure 1A and 1B). Previously, a human homolog, rrhbdd1, was also shown to be highly expressed in testes [13]. These findings suggest that this Rhomboid-containing protein may be involved in regulating mammalian spermatogenesis.

Apoptotic regulation of mRHBD1 in a mouse spermatogonia line

In order to investigate this putative role of mRHBD1 in spermatogenesis, we studied its expression in a mouse-derived spermatogonia cell line, GC-1 spg (ATCC number: CRL-2053; Figure 2A). Stable mRHBD1 knockdown shRNA and negative control shRNA GC-1 cells were constructed (Figure 2B–E). Because RHBD1 regulates apoptosis in some somatic cells [13], the occurrence of apoptosis in mRHBD1 knockdown GC-1 cells was determined after challenges with apoptotic stimuli, PS341 and UV irradiation. Procaspase 3 cleavage was indicated by an increase in p17 or p12, its cleavage products during apoptosis. As shown in Figure 2F and 2G, these apoptosis markers were more prominent in the mRHBD1 knockdown GC-1 cells than in the negative control GC-1 cells after exposure to apoptosis inducers, PS341 and UV irradiation. We also confirmed by FACS analysis that apoptosis increased in stable mRHBD1 knockdown GC-1 cells after UV stimulation (Figure 2H). These results indicate that mRHBD1 normally inhibits apoptotic activity in spermatogonia.

Endogenous mRHBD1 knockdown in GC-1 cells does not affect their proliferation

To investigate the proliferation of mRHBD1 knockdown GC-1 cells, MTS and CCK-8 assays were performed. Proliferation did not differ between these cells and the negative control cells (Figure 3A and 3B). Thus, mRHBD1 knockdown GC-1 cells possess normal survival and proliferation capacities under in vitro conditions.

Stable mRHBD1 knock-down GC-1 cells lose survival and differentiation capacities in mouse seminiferous tubules

The above experiments demonstrating the anti-apoptotic activity of mRHBD1 in GC-1 cells suggested that this factor is associated with the survival of spermatogonia. To investigate an in vivo role for mRHBD1, mRHBD1 knockdown GC-1 cells and negative control GC-1 cells were transplanted...
Apoptosis in mRHBDD1 knock-down GC-1 cells. A. mRHBDD1 protein levels in GC-1 cells and mouse testis were examined by Western blotting; the same blot was immunoblotted with anti-β-actin antibody as internal control. Protein (30 μg) was separated on a 12% polyacrylamide gel as described in the Methods section. B and C. Screening for effective RNAi plasmids against mRHBDD1. Western blotting indicated that pRNAT-H1.1/Hygro-4# and 6# were both candidates; pRNAT-H1.1/Hygro-negative was selected. GFP was a loading control. D and E. Identification of stable mRHBDD1 knockdown GC-1 cell clones by Real-Time PCR and Western blotting. Clone 2# was selected for the following experiments. F and G. Western blot analysis of caspase 3 cleavage in 2# and negative control GC-1 cells after PS341 or UV treatment. Control cells were subcultured from a selected stable GC-1 cell clone expressing a control negative shRNA. H. FACS assay to analyse apoptosis in 2# and negative control GC-1 cells after UV irradiation. The percentage of apoptotic cells (% of total cells) was determined using the program EXPO32-ADC and shown as a bar chart. * (p < 0.05)

Proliferation of mRHBDD1 knockdown GC-1 cells and negative control cells. GC-1/2# cells and GC-1/negative cells were seeded at a density of 1000/well in 96-well plates in DMEM with 10% FBS. Measurements of absorbance at 490 nm for the MTS assay (A) and at 450 nm for the CCK-8 assay (B) were made once per day for 5 days. Results are the means ± SD of three independent experiments.
**Figure 4**
Confocal images of mRHBDD1 knockdown and control mouse testis sections. A. Autologous control tubules from the right epididymis of one mouse. pRNAT-H1.1/Hygro-negative control (encoding GFP) stably-expressing GC-1 cells were transplanted into the seminiferous tubules of the right testis. These cells differentiated into spermatids with GFP in the epididymal tubules. They had acrosomes that were stained with PNA-linked Rhodamine. I. PNA-stained acrosomes. II. Hoechst 33258-stained nuclei. III. GFP-labelled spermatids. IV. Merge. B. Heterologous control (seminiferous tubule). Stable pRNAT-H1.1/Hygro-negative control (encoding GFP) GC-1 cells were transplanted into the seminiferous tubules of testes from another animal, capable of producing spermatids in the seminiferous tubules, and were tagged with GFP. They contained acrosome structures stained with PNA-linked Rhodamine. I. PNA-stained acrosomes. II. Hoechst 33258-stained nuclei. III. GFP-labelled spermatids. IV. Merge. C. Test group tubules from the left epididymis of the same animal as in A. pRNAT-H1.1/Hygro-6# containing an effective RNAi plasmid against stable mRHBDD1-expressing GC-1 cells (cell line 2#) was transplanted into the seminiferous tubules of the left testis. The transplanted GC-1 cells and spermatids were not found in the seminiferous or epididymal tubules, indicating that these stable germ cells had died or lacked the ability to differentiate. I. PNA-stained acrosomes. II. GFP-labelled spermatids. III. Hoechst 33258-stained nuclei. D. Residual endogenous germ cells that had resisted busulfan treatment in mouse testes differentiated into sperm cells in the epididymal tubules of the left testis, indicating that these busulfan-treated seminiferous tubules could still support the generation of spermatids. Note: PNA specifically stains the acrosomes of spermatids. I. PNA-stained acrosomes. II. Hoechst 33258-stained nuclei. III. Merge. White triangles are PNA-stained acrosomes (AI, BI and DI). Open triangles are GFP-labelled spermatids with GFP tag derived from stable GC-1 cells (AIII, BIII).
into mouse seminiferous tubules and their survival and differentiation into spermatids were studied. In this in vivo model, mRHBDD1 knockdown GC-1 cells with pRNAT-H1.1/Hygro-6# and negative control cells with pRNAT-H1.1/Hygro-negative (both GFP positive) were transplanted into the seminiferous tubules of, respectively, right and left mouse testes, from which most of the endogenous germ cells had been eliminated by busulfan treatment. As shown in Figure 4A and 4B, autologous and heterologous transplanted control cells, tagged by GFP, differentiated into spermatids, as shown by the presence of acrosomes (stained red by PNA-labelled Rhodamine). However, mRHBDD1-suppressed GC-1 cells did not survive or differentiate into spermatids, as shown by the absence of GFP and acrosomes from the epididymal tubules (Figure 4C). In contrast, the mouse residual germ cells (no GFP tag) that resisted busulfan treatment did differentiate into spermatids, as shown by the presence of PNA-Rhodamine-labelled acrosomes (Figure 4D). Thus, mRHBDD1 knockdown apparently influences the survival of germ cells and their differentiation into sperm cells in vivo.

Discussion

It is well established that the regulation of apoptosis is a key control mechanism for spermatogenesis [2,6]. An early wave of germinal cell apoptosis during the first round of spermatogenesis in the immature testis is probably a critical requirement during testis development because it maintains a balanced ratio of germ cells to Sertoli cells [20]. Moreover, the elimination of supra-optimal production of spermatogonia by apoptosis is common in the adult testis, where it is primarily restricted to spermatogonia [21]. However, apoptosis of defective germ cells is also essential for maintaining genetic fidelity in the development and survival of offspring. RHBDD1/mRHBDD1 may play a role in regulating these events.

The testis cell transplantation method has been developed as a powerful approach for studying spermatogenesis. It is also used to examine spermatogenic defects, to correct male infertility and to generate transgenic animals [22]. Primary germ cells are the best model to perform these experiments, but there are some puzzles: 1) It is difficult to obtain enough spermatogonia cells from seminiferous tubules since these cells are a minority among germ cells. 2) The transfection efficiency of pRNAT-H1.1/Hygro-6# into spermatogonia is very low. 3) Most of spermatogonia transfected by plasmids will undergo death. To resolve the above problems, we have established a method for studying mammalian spermatogenesis using a GC-1 spermatogonia line transplantation model. Its distinct advantage is to make it much easier to obtain highly-purified GC-1 spermatogonia cells for experiments than by purifying primary spermatogonia cells from mouse seminiferous tubules.

The results of the current study show that transplanted control GC-1 cells can differentiate from spermatogonia into spermatids, indicating that they possess the properties of germ stem cells, capable of differentiating in an in vivo microenvironment. Furthermore, the appearance of spermatids derived from control GC-1 cells and from the residual endogenous germ cells that had resisted busulfan treatment in the seminiferous tubules showed that these damaged tubules maintained the ability to generate spermatids from spermatogonia. Thus, the disappearance of the mRHBDD1 knockdown GC-1 cells from the seminiferous tubules 8 weeks after transplantation is the direct result of endogenous mRHBDD1 suppression.

mRHBDD1 has anti-apoptotic potential, and its suppression in GC-1 cells vitiates spermatogonia survival and differentiation in vivo. The mRHBDD1 knockdown GC-1 cells did not disappear from the mouse seminiferous tubules because they entered a morbid state and died; they showed normal survival and differentiation in vitro like the negative control GC-1 cells. Although the mechanism by which mRHBDD1 regulates mammalian spermatogenesis needs to be clarified, it seems quite logical that it occurs by regulating germ cell apoptosis, because mRHBDD1 knockdown GC-1 cells were more sensitive to apoptotic stimuli than negative control cells. An apparently indispensable role for mRHBDD1 during this process has been demonstrated by the present in vivo results. This warrants additional studies to determine whether malfunction of this enzyme may be a basis of male infertility. This information may be useful for the development of a male contraceptive.

Conclusion

The mRHBDD1 gene may be associated with the regulation of mammalian spermatogenesis by regulating apoptosis in spermatogonia.

Methods

Cell culture and transfection

GC-1 cells, a mouse-derived spermatogonia line, were purchased from ATCC (ATCC number: CRL-2053) and cultured in fresh Dulbecco’s modified Eagle’s medium supplemented with 10% foetal bovine serum in 5% CO₂ incubators at 37°C. Adherent cells were passaged every 2–3 days with 0.5 mg/ml trypsin (1:250) and 0.53 mM EDTA. All transfections were performed using Lipofectamine 2000 (Invitrogen).

Plasmid construction

An RNAi targeting sequence against mrhbdd1 (mouse homolog of rhbd1) designated 6# (sense primer: CATCCGAATCAGCGCATGACATCAAATCAGAGATTTGAAGT CACGGCTGATCCTTTTTTA; antisense primer: AGCTTAAAA AAGAATCAGCGCATGACATCAAATCAGAGATTTGAAGTCA CGGCTGATCCTTTTTTA) was cloned into pRNAT-H1.1/Hygro.

RT-PCR

Male BALB/c mice and Sprague-Dawley rats were obtained from the Laboratory Animal Centre of PUMC (Beijing, China).
China). The Animal Ethics Committee of the National Research Institute for Family Planning Beijing approved the animal experimentation protocols and all animal experiments were performed according to the Guidelines for the Care and Use of Laboratory Animals established by the Chinese Council on Animal Care. The mice and rats were anesthetized under ether. The following organs were excised: intestine, muscle, spleen, liver, lung, heart, kidney, brain, epididymis, stomach and testis. The tissues were immediately immersed in liquid nitrogen and stored at -80°C pending use. Trizol reagent (Invitrogen) was used to isolate total RNA following the manufacturer's instructions. A SuperScript™ First-Strand Synthesis System (Invitrogen) was used for RT-PCR following the manufacturer's instructions.

**Western blotting**

Thirty μg of protein was loaded on each lane of a 12% SDS-polyacrylamide gel, separated and then electrochemically transferred on to a PVDF membrane (GE Healthcare). The membrane was blocked in 5% skim milk for 1 h at room temperature and then incubated with polyclonal rabbit anti-RHBDD1 (Sigma), anti-HA (Sigma), anti-GFP (Santa Cruz), anti-caspase 3 (Santa Cruz) or anti-β-actin (Santa Cruz) overnight at 4°C. The membrane was incubated with anti-rabbit or anti-mouse HRP-IgG (Santa Cruz) for 1 h at room temperature. Chemiluminescence was detected using an ECL blot detection system (Santa Cruz).

**Induction of apoptosis in GC-1 cells**

For UV-induced apoptosis, mRHBDD1 knockdown or negative control stable GC-1 cells were exposed to UV irradiation (65 μl). The medium was changed and cells were harvested 4, 5, 6, 7, 8 or 12 h post UV treatment. For PS341-induced apoptosis, cells were treated with 1 μM PS341 and were harvested 12 h post treatment.

**Flow cytometry**

Cells were removed from the plates using 0.25% trypsin-EDTA at 37°C for 5 min, collected, and fixed with 75% ethanol for 30 min at 4°C. They were stained with 50 μg/ml propidium iodide and 100 μg/ml RNase A in PBS at 37°C for 20 min. The DNA content of 10,000 cells was analyzed using a COULTER flow cytometer (EPICS-XL) with EXP322-ADC software. The percentage of apoptotic cells (% of total cells) was analyzed by the program EXP322-ADC and shown as a bar chart (Microsoft Excel).

**Cell proliferation assay**

Cells were seeded at a density of 1000/well in 96-well plates in DMEM with 10% FBS. The medium was changed regularly, and measurements of proliferating cells were performed once per day for 5 days. For the MTS assay (Promega), 20 μl of a combined MTS/PMS solution was added to each well containing 100 μl culture medium and incubated for 3 h at 37°C, then the absorbance was measured at 490 nm using an ELISA plate reader. For the CCK-8 assay (Dojindo Molecular Technologies), 10 μl of CCK-8 solution was added to each well containing 100 μl culture medium and incubated for 3 h at 37°C, then the absorbance was measured at 450 nm using an ELISA plate reader.

**Regeneration of spermatogenesis by GC-1 stable transfectant cells in mouse seminiferous tubules**

BALB/c male mice were obtained from the Laboratory Animal Centre of PUMC (Beijing, China). The Animal Ethics Committee of National Research Institute for Family Planning Beijing approved the animal experimentation protocols and all animal experiments were performed according to guidelines (Guidelines for the Care and Use of Laboratory Animals) established by the Chinese Council on Animal Care. The mice were anesthetized under ether. Animals were maintained at 24 ± 1 °C, 55 ± 1% humidity with a 14 h Light: 10 h Dark cycle. Food and water were provided ad libitum. Male mice aged 6–7 weeks were treated with Busulfan (Sigma), 40 mg/kg i.p., to destroy the endogenous germ cells at 4–5 weeks prior to the transplantation. A recipient mouse was anesthetized and the testis was exteriorized through a midline abdominal incision. A sample of approximately 10 μl of cell suspension was injected into the recipient mouse testis by the rete testis, whereby approximately 60%–90% of the surface tubules were filled with cells. For each recipient mouse, one testis was used as the test site for the transplanted stably expressed pRNAT/H1.1/Hygro-negative and the contralateral testis was used for transplantation of the stably expressed pRNAT/H1.1/Hygro-6#.

At 8 weeks after transplantation, the treated and control testes were excised for histological examination. The testis tissues were embedded in O.C.T. compound, snap-frozen in liquid nitrogen and cryosectioned. Forty sections were prepared from each testis and examined by fluorescent microscopy (Microphot Nikon) to identify the donor-derived germ cells. Sections were selected, fixed with 4% paraformaldehyde in PBS, stained with TRITC/Rhodamine conjugated peanut agglutinin (PNA) (Sigma) to visualize acrosomes, followed by staining with Hoechst 33258 (Sigma) to visualize nuclei and re-examined by confocal laser microscopy (Leica, Germany).

In our experiment, approximately 80% of the surface tubules were filled with cells in the control and test testes.

**Statistical analysis**

The results are expressed as means ± SD. Statistical comparisons used Student’s t-test. P < 0.05 was considered statistically significant.
Authors’ contributions
YW, WS, SL, and XG carried out all experiments. SZ observed and analyzed the slides. YW and SSK wrote and revised the manuscript. YW, SM and LW conceived of the study and participated in its design and coordination. All authors read and approved the final manuscript.

Acknowledgements
This work was supported by grants from the National Program for the Important Research Plan (2006CB944002) and the National Program for the key Basic Research Project (2006CB504001) from the Ministry of Science and Technology of China, and a grant to the Creative Research Group (No. 30721063) from the National Natural Science Foundation of China, and the Research Fund for the Doctoral Program of High Education (20080230029). The experimental study was performed in compliance with the laws of the People’s Republic of China.

References
1. Wang RA, Nakane PK, Koji T: Autonomous cell death of mouse male germ cells during fetal and postnatal period. Biol Reprod 1998, 58(5):1250-1256.
2. Rodriguez I, Ody C, Araki K, Garcia I, Vassalli P: An early and massive wave of germinal cell apoptosis is required for the development of functional spermatogenesis. Embo J 1997, 16(9):2262-2270.
3. Mori C, Nakamura N, Dix DJ, Fujikoa M, Nakagawa S, Shiota K, Eddy EM: Morphological analysis of germ cell apoptosis during postnatal testis development in normal and Hsp 70–2 knock-out mice. Dev Dyn 1997, 208(1):125-136.
4. Billig H, Furuta I, Rivier C, Tapanainen J, Parvinen M, Husej AJ: Apoptosis in testis germ cells: developmental changes in gonadotropin dependence and localization to selective tubule stages. Endocrinology 1995, 136(1):5-12.
5. Barke A: Apoptosis of male germ cells, a generalized or a cell type-specific phenomenon? Endocrinology 1995, 136(1):3-4.
6. Print CG, Loveland KL: Germ cell suicide: new insights into apoptosis during spermatogenesis. Bioessays 2000, 22(5):423-430.
7. Schaller CE, Wang CL, Beck-Engeser G, Goss L, Scott HS, Anderson MS, Wabl M: Expression of Aire and the early wave of apoptosis in spermatogenesis. J Immunol 2008, 180(3):1338-1343.
8. Koonin EV, Makarova KS, Rogozin IB, Davydov I, Letelier MC, Pellegriini L: The rhomboids: a nearly ubiquitous family of intramembrane serine proteases that probably evolved by multiple ancient horizontal gene transfers. Genome Biol 2003, 4(3):R19.
9. Lee JR, Urban S, Garvey CF, Freeman M: Regulated intracellular ligand transport and proteolysis control EGF signal activation in Drosophila. Cell 2001, 107(2):161-171.
10. Urban S, Lee JR, Freeman M: Drosophila rhomboid-1 defines a family of putative intramembrane serine proteases. Cell 2001, 107(2):173-182.
11. McQuibban GA, Saurya S, Freeman M: Mitochondrial membrane remodelling regulated by a conserved rhomboid protease. Nature 2003, 423(6939):537-541.
12. Howell SA, Hackett F, Jongco AM, Withers-Martinez C, Kim K, Carruthers VB, Blackman MJ: Distinct mechanisms govern proteolytic shedding of a key invasion protein in apicomplexan pathogens. Mol Microbiol 2005, 57(9):1342-1356.
13. Wang Y, Guan X, Fok KL, Li S, Zhang X, Miao S, Zong S, Koide SS, Chan HC, Wang L: A novel member of the Rhomboid family, RHBD1, regulates BIK-mediated apoptosis. Cell Mol Life Sci 2008, 65(23):3822-3829.
14. Urban S: Rhomboid proteins: conserved membrane proteases with divergent biological functions. Genes Dev 2006, 20(22):3054-3068.
15. Pascall JC, Brown KD: Intramembrane cleavage of ephrinB3 by the human rhomboid family protease, RHBDL2. Biochem Biophys Res Commun 2004, 317(1):244-252.
16. Lohi O, Urban S, Freeman M: Diverse substrate recognition mechanisms for rhomboids; thrombomodulin is cleaved by Mammalian rhomboids. Curr Biol 2004, 14(3):236-241.
17. Gottlieb E: OPA1 and PARL keep a lid on apoptosis. Cell 2006, 126(1):27-29.
18. McQuibban GA, Lee JR, Zheng L, Juusola M, Freeman M: Normal mitochondrial dynamics requires rhomboid-7 and affects Drosophila lifespan and neuronal function. Curr Biol 2006, 16(10):982-989.
19. Allan DJHB, Kerr JFR: Cell death in spermatogenesis. London: Oxford university Press; 1987.
20. Allan DJ, Harmon BV, Roberts SA: Spermatogonial apoptosis has three morphologically recognizable phases and shows no circadian rhythm during normal spermatogenesis in the rat. Cell Prolif 1992, 25(3):241-250.
21. Brinster RL: Germine stem cell transplantation and transgenesis. Science 2002, 296(5576):2174-2176.

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