Apoptosis of Sertoli cells after conditional ablation of murine double minute 2 (Mdm2) gene is p53-dependent and results in male sterility

S Fouchécourt1,2,3,4, G Livera5, S Messiaen5, B Fume1,2,3, A-S Parent6, J-C Marine7,8 and P Monget1,2,3,4

Beside its well-documented role in carcinogenesis, the function of p53 family has been more recently revealed in development and female reproduction, but it is still poorly documented in male reproduction. We specifically tested this possibility by ablating Mdm2, an E3 ligase that regulates p53 protein stability and transactivation function, specifically in Sertoli cells (SCs) using the AMH-Cre line and created the new SC-Mdm2−/− line. Heterozygous SC-Mdm2+/− adult males were fertile, but SC- Mdm2−/− males were infertile and exhibited: a shorter ano-genital distance, an extra duct along the vas deferens that presents a uterus-like morphology, degenerated testes with no organized seminiferous tubules and a complete loss of differentiated germ cells. In adults, testosterone levels as well as StAR, P450c17 (Cyp17a1) and P450sc (Cyp11a1) mRNA levels decreased significantly, and both plasma LH and FSH levels increased. A detailed investigation of testicular development indicated that the phenotype arose during fetal life, with SC-Mdm2−/− testes being much smaller at birth. Interestingly, Leydig cells remained present until adulthood and fetal germ cells normally initiated meiosis. Inactivation of Mdm2 in SCs triggered p53 activation and apoptosis as early as 15.5 days post conception with significant increase in apoptotic SCs. Importantly, testsis development occurred normally in SC-Mdm2−/− lacking p53 mice (SC-Mdm2+/−p53−/−) and accordingly, these mice were fertile indicating that the aforementioned phenotypes are entirely p53-dependent. These data not only highlight the importance of keeping p53 in check for proper testicular development and male fertility but also certify the critical role of SCs in the maintenance of meiotic repression.

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Sertoli cells (SCs) are the supporting cell lineage of the male germ line and their function is critical for male fertility. SCs differentiate early during fetal life; they are the first cells that differentiate in the testes and they drive the sexual differentiation of the gonad (for review, see Svingen and Koopman1). In the mouse, these cells start forming cords by enclosing germ cells as early as 12 days post conception. SCs are responsible for providing germ cells with a specialized environment to promote their survival and orchestrate their differentiation throughout life. They also allow the differentiation of Leydig cells through paracrine signaling, including Desert hedgehog signaling, for example,2 and promote their activity.3 SCs are involved in masculinizing the embryo as they produce high levels of anti-Müllerian hormone (AMH) that triggers the regression of the future female genitalia. Strikingly, SCs only proliferate during fetal and post-natal life, and definitively cease cycling around 10 days postpartum. Interestingly, these cells are relatively resistant to apoptosis in response to DNA damage. It has been proven in developing rat testes that these cells easily survive a high dose of radiation exposure.4 The mechanisms underlying the poor responsiveness of SCs to DNA damage remains elusive. In the human embryonic testis, the mild apoptotic response of SCs following radiation is decreased by pharmacological inhibition of p53 with pifithrin alpha.5

The role of p53 has been largely documented as the ‘guardian of the genome’; owing to its deregulation in numerous cancers, it is considered as a major tumor suppressor. From a model organism, it appears that p53 homologs are also key regulators of development and reproduction. Interestingly in mammals, p53-related proteins (i.e., p63 or p73) are also involved in the development of the central nervous system. Inactivation of any of the three p53 family members impairs mouse female fertility.6,7 On the other hand, the data relating p53 to male fertility are still scarce. So far, in testis, the function of p53 is little documented, except in testicular germ cell tumors8 and germ cell apoptosis (apoptosis of prenatal sperm cells9 and of damaged sperm in the adult10). The murine double minute 2 gene Mdm2 belongs to a large family of RING finger-containing proteins,
and functions mainly as an E3 ligase regulating the activity of various substrates by ubiquitylation (mono- or poly- ubiquitylation). Among the substrates is the p53 tumor suppressor, which is responsible for transcriptional activation of genes involved in the cell cycle, apoptosis and cell aging.\textsuperscript{11–13} Overexpression of Mdm2 is one of the mechanisms that leads to p53 inactivation by promoting its proteasome-dependent degradation after ubiquitylation in the tumors that retain wild-type p53 (for review, see Marine and Lozano\textsuperscript{14}). In addition, Mdm2 can bind the p53 transactivation domain and directly interfere with p53 transcriptional regulatory mechanisms. Mdm2-null mice are not viable because of early embryonic lethality (E3.5), but are viable in a p53-null background.\textsuperscript{15,16} To test whether the suppression of p53 function is required for testicular development and male fertility, we inactivated specifically Mdm2 in SCs using the AMH-Cre line.\textsuperscript{17}

**Results**

Mdm2 mRNA was detected in testes (adult and P10) and was present in SC-enriched fractions from prepubertal P10 mice (Supplementary data 1). To test the hypothesis of the critical role of Mdm2 in SCs, we created a transgenic line with targeted invalidation of Mdm2 specifically in SCs by crossing AMH-Cre and Mdm2\textsuperscript{LL} lines, and named this new line SC-Mdm2\textsuperscript{−/−}. The contemporary Cre-Mdm2\textsuperscript{LL} littermate controls were called Cre-. We verified that the allele lacking exons 5 and 6 (Mdm2\textsuperscript{δexons5/6}) after the Cre recombination event (226 bp) was detected in testes but not in any other tissues (Supplementary data 2).

At adulthood (3 months), SC-Mdm2\textsuperscript{−/−} males were infertile and had very small testes (Figures 1a–d; testis weight: SC-Mdm2\textsuperscript{−/−} 5.35 mg\textsuperscript{***} ± 1.40 versus Cre- 112.8 mg ± 4.5, \textit{P}<0.001), an abnormal external genital tract and a shorter ano-genital distance compared with Cre- (Figures 1g and h). Conversely, heterozygous SC-Mdm2\textsuperscript{−/+} males were normally fertile and had a normal genital tract (not shown). Testicular histology in adult SC-Mdm2\textsuperscript{−/−} (Figure 2A) was characterized by the absence of seminiferous tubule organization and differentiated germ cells.

**Figure 1** Internal genital tract of SC-Mdm2\textsuperscript{−/−} males (a and b) compared with Cre- (c and d) and normal female tracts (e and f). Black circles are positioned on testes, stars on the caput of the epididymis, and arrowheads on the extra ducts present in SC-Mdm2\textsuperscript{−/−} (a, b) and the uterus in females (e). SV: seminal vesicles; rSV: reduced seminal vesicles; VD: vas deferens. Ov: ovary (E: white circles); uT: uterus. External genital tract of SC-Mdm2\textsuperscript{−/−} males (g/h). Arrows show the AGD in SC-Mdm2\textsuperscript{−/−} (g) and Cre- (h) males. a = anus, p = penis. Bar represents 1 cm.
versus Cre- 48.6 mg ± 2.2, \( P < 0.001 \) (Figure 1a–d). Interestingly, we observed an extra duct along the vas deferens with the apparent aspect of a uterus (Figures 1a, b, e and f) and histology similar to the uterus (Figures 2E and F); the histology of the vas deferens was normal (Figures 2C and D); no sperm was present in the lumen (*) of the vas deferens contrary to Cre- (C, c and D, d). The histology of the extra duct was similar to the uterus (E, e and F, f). Bar in large windows represents 100 μm; magnification is four times greater in small windows (a, c, d, e, f).

In correlation with a decrease in seminal vesicle weight, plasma testosterone levels significantly decreased in SC-Mdm2\(^{-/-}\) (Figures 3a, \( n=8 \) for each genotype), along with mRNA levels per testis of Cyp17a1 (P450c17), Cyp11a1 (P450sc) and STAR (Figure 3b, \( n=5 \) for each genotype). Consequently, LH levels significantly increased by a factor of around 7 in SC-Mdm2\(^{-/-}\) compared with Cre- (Figure 3c; \( P<0.001 \)). Interestingly, FSH levels also significantly increased in SC-Mdm2\(^{-/-}\) (Figure 3d; \( P<0.001 \)).

As AMH-Cre is expected to induce the deletion of the Mdm2 allele during fetal life (around E15), we sought to determine the precise kinetics of the appearance of this phenotype. Testes were harvested at various stages during fetal (E15, E16, E18) and post-natal life (P1) (Figures 4–7). The analysis of

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**Figure 2**  Histology sections (hematoxylin staining) of testes and ducts in adult SC-Mdm2\(^{-/-}\) males (A, C, E) compared with Cre- males for testes and vas deferens (B, C) and females for uterus (E, F). Testicular cells were not organized in seminiferous tubules in adult SC-Mdm2\(^{-/-}\) testes (A, a) and no differentiated germ cells were present. No sperm was present in the lumen (*) of the vas deferens contrary to Cre- (C, c and D, d). The histology of the extra duct was similar to the uterus (E, e and F, f). Bar in large windows represents 100 μm; magnification is four times greater in small windows (a, c, d, e, f).
sections at P1 already indicated a pronounced reduction in gonad size (0.10 ± 0.01 vs 0.38 ± 0.02 μl, n = 4, P<0.01, respectively, for SC-Mdm2−/− and the control) with an almost total loss of cord structures. Testicular cord disappearance was similarly pronounced at E18, while a few cords could still be retrieved at E16 and E15. To understand this phenomenon, apoptosis was investigated by observing TUNEL (Figures 4 and 5) and cleaved-caspase 3 (Figure 6). Both methods similarly revealed numerous apoptotic cells in the SC-Mdm2−/− testes at all the stages examined (e.g., 29 ± 11 versus 150 ± 22 cells/mm², n = 4, respectively, for the control and SC-Mdm2−/− at P1). Interestingly, double AMH-TUNEL staining clearly indicated that dying cells were mostly SCs at E15 and E16 (Figure 4). Quantification of AMH-TUNEL cells evidenced a significantly increased apoptosis (1.2 ± 0.7 versus 37.2 ± 2.7%, n = 3–5, respectively, for the control and SC-Mdm2−/− at E15). Of interest, we observed no evidence for senescence based on the absence of p21 or heterochromatin foci in these cells (data not shown). As Mdm2 invalidation is expected to activate p53, double AMH-p53 staining was performed at E15. In control testes (Cre−), hardly any AMH-positive cells expressed detectable levels of p53, whereas in SC-Mdm2−/− testes, we observed that many AMH-positive cells were clearly stained for p53 (Figure 5).

To clarify the identity and fate of the cells remaining in the absence of SCs, germ cell and Leydig cell populations were analyzed in developing SC-Mdm2−/− testes. Germ cells were identified using DDX4 staining (Figure 6) at E15 and P1. As expected, they aggregated in the cords in the control testes. A similar gathering of germ cells (despite the absence of cords or lack of SCs) was retrieved in E15 and E16 testes in SC-Mdm2−/− testes; consequently, germ cells tended to be more dispersed throughout the testes (E18 and P1). Surprisingly, we could still observe many DDX4-positive cells in P1 testes, indicating that the loss of the supporting SCs did not immediately trigger germ cell death. As SCs are believed to produce a meiosis-repressing environment for germ cells during fetal life, we analyzed the possible presence of the meiotic marker, SYCP3 (Figure 6). In control testes, SYCP3-positive cells were virtually absent, whereas in SC-Mdm2−/− testes, we detected numerous SYCP3-positive cells, all equally positively stained for DDX4. A close examination of the localization of SYCP3 using immunofluorescence (Figures 6i and j) indicated that SYCP3 formed fine threads in the germ cell nuclei, probably along the chromosomal axis, as expected in meiotic cells. Leydig cells were detected using 3βHSD staining (Figure 7). As expected, they were retrieved between the cords in the control testes and formed well-organized clusters around birth. In SC-Mdm2−/−

\[ \text{Figure 3} \] SC-Mdm2−/− hormonal status. (a) Plasma testosterone levels in adult SC-Mdm2−/− males, in basal conditions and after human chorionic gonadotropin (hCG) stimulation. In both SC-Mdm2−/− and Cre− (8 males for each genotype), hCG induced a significant increase (***/P<0.001) in testosterone secretion, as expected. Plasma testosterone levels decreased significantly in SC-Mdm2−/− compared with Cre− (**P<0.01). Data shown as mean +/- S.E.M.; statistical analyses: two-way ANOVA followed by Bonferroni’s post hoc test; hatching: basal levels; white: hCG stimulated levels. (b) SC-Mdm2−/− tests exhibit a decrease in mRNA levels of Cyp17a1 (P450c17), Cyp11a1 (P450scc) and StAR. The amounts of mRNA of the three steroidogenic genes were compared between SC-Mdm2−/− and Cre− (n = 5 mice for each genotype, tested in triplicate) using TaqMan quantitative real-time PCR as described in Materials and Methods. Normalization was performed with two housekeeping genes, β-actin and GAPDH, that both exhibited similar expression between samples. Results are presented as % of mRNA level in Cre− (100 is arbitrary level for whole Cre− testes) and a non-parametric test (***/P<0.001). White: Cre−; hatching: SC-Mdm2−/−. (c) LH levels in SC-Mdm2−/− males: plasma LH concentrations expressed in ng/100 μl in Cre− (n = 13) and SC-Mdm2−/− (n = 19) determined by RIA. (d) FSH levels in SC-Mdm2−/− males: plasma FSH concentrations expressed in ng/100 μl in Cre− (n = 16) and SC-Mdm2−/− (n = 23) determined by RIA. Data shown as mean +/- S.E.M. Statistical analysis: Mann-Whitney test (***/P<0.001). White: Cre−; hatching: SC-Mdm2−/−.
testes, the density of Leydig cells was normal. However, they did not form tight clusters. Interestingly, these cells were still retrieved in adult mutant testes that appeared literally packed with $\beta$HSD-positive cells. This clearly indicated that the presence of SCs was dispensable for Leydig cell maintenance in this model.

To verify the implication of the $p53$ tumor suppressor gene in the phenotype, we created a new line by introgressing the SC $Mdm2$ invalidation in the $p53^{-/-}$ line, as described in Materials and Methods ($p53^{-/-}$ being normally fertile). Mating was conducted until at least four individuals of each genotype of interest and their respective controls were obtained (Table 1). As expected, SC-$Mdm2^{-/-}p53^{+/+}$ were infertile with the same phenotype as SC-$Mdm2^{-/-}p53^{-/-}$, whereas their controls Cre-$p53^{+/+}$ were fertile. Conversely, SC-$Mdm2^{-/-}$ lacking $p53$ (SC-$Mdm2^{-/-}p53^{-/-}$) were normally fertile proving that the phenotype due to $Mdm2$ invalidation involves $p53$. One $p53$ allele (SC-$Mdm2^{-/-}p53^{+/+}$) is sufficient to maintain the infertility phenotype.

**Discussion**

To date, few studies have examined the critical factor for SC survival and the development of a gonad devoid of SCs. We report here that Mdm2 is critical to prevent SC apoptosis and that its specific invalidation triggers a rapid loss of this lineage in testes. Without SCs, Leydig cells persisted, albeit with reduced steroidogenic activity and germ cells did not differentiate properly and only survived a few days.

$Mdm2$ is a key regulator of $p53$-mediated apoptosis and is proven to be a major downregulator of $p53$ in many cell types such as kidney cells, cardiomyocytes, neuronal cells, smooth muscle cells, and hepatocytes. In testes, the role of $Mdm2$ has already been documented in germ cells. $Mdm2$ is one mechanism leading to $p53$ inactivation in male GCs. It is also suggested that $p53$ mediates spermatogonal apoptosis after DNA damage or overheating. Contrary to GCs, SC apoptosis is poorly documented as it is suspected that this phenomenon is rare. $p53$ was suggested to be involved in an intrinsic apoptotic program in SCs, but poorly active.
Figure 5  SC apoptosis is correlated with p53 activation. The activation of P53 was investigated in SCs at E15 in control (Cre-) (a, c) and SC Mdm2<sup>-/-</sup> (b, d) testes through double p53/AMH immunostaining. Upper panel represents a global view of a testis section and lower panel a higher magnification of testicular cords. No p53 was observed in control SCs (a, c), while many p53-positive cells (brown) were detected in cord-like structures in SC Mdm2<sup>-/-</sup> (b, d). Interestingly, most were clearly positive for AMH staining (purple) as well. Arrowheads point to p53-negative SCs (c and d) and arrows point to p53-positive SCs (d: double positive). Bars represent 10 μm.

Figure 6  Maintenance of germ cells and premature meiotic entry in absence of SC. DDX4 was investigated by immunostaining (purple) or immunofluorescence (red) in E15 (a–d, i, j) and P1 (e–h) testes from control (Cre-) and SC Mdm2<sup>-/-</sup> animals to follow the germ cell population. DDX4/SYCP3 double immunostaining was performed in E15 testes: immunohistochemistry (a–d: purple for DDX4 / brown for SYCP3; c, d insets provide a higher magnification of germ cells), and immunofluorescence (i, j: DDX4 in red, SYCP3 in green forming obvious filaments in germ cell nuclei) indicated that no SYCP3-positive cells were present in the control testes, while many were retrieved in the SC Mdm2<sup>-/-</sup> testes. Double DDX4 (purple)/cleaved caspase3 (brown) immunostaining was performed in P1 testes. Upper panels (e, f) represent a global view of a testis section and lower panels a higher magnification (g, h). In both control and SC Mdm2<sup>-/-</sup> testes, numerous germ cells and few apoptotic cells were retrieved (i.e., positive for caspase 3). In H7, the arrow points to an apoptotic germ cell. Bars represent 10 μm.
Nevertheless, the role of Mdm2 in SCs had not been examined prior to this work. The inactivation of Mdm2 using AMH-Cre led to a rapid increase in the p53 protein and apoptotic markers triggering an almost total loss of SCs during fetal life. As expected, this phenotype was alleviated when p53 was inactivated in the p53<sup>−/−</sup> genetic background. This demonstrates that p53 activity is strongly repressed in fetal SCs, and may explain the relative poor radiosensitivity of this cell type. 26 As p53 is expressed in various testicular cells (and not only SCs), we cannot exclude the p53 effect in other testicular cell types leading to SC apoptosis. Targeted deletion in SCs (using SC-Mdm2<sup>−/−</sup>/p53<sup>loxP/loxP</sup>) would allow to reach this point.

Fetal SCs are critical for the masculinization of the embryo, both indirectly through androgen production by LC and directly as they produce AMH, the hormone inducing the regression of the female genitalia. We report here that SC-Mdm2<sup>−/−</sup> males displayed a pronounced decrease in AMH staining as early as E15. The extra duct persisting at adulthood in those mice is probably due to this drop. This lack of female duct regression has been observed previously in AMHR2KO and AMHKO males.28,29

SCs are essential for the differentiation of fetal Leydig cells.30 Of interest, we report here that even in the absence of SCs and testicular cords, LC were observed in adult testes. This suggests that SCs are dispensable for the increase and maintenance of the LC population during late fetal and postnatal life. Future work will definitively need to address the nature of the numerous steroidogenic cells observed in

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**Table 1** Individual male number (#) and fertility status tested with 2 or 4 normal females (f), of the three genotypes corresponding to the introgression of SC-Mdm2<sup>−/−</sup> in p53: SC-Mdm2<sup>−/−</sup> p53<sup>−/−</sup> or +/−, and their corresponding controls: Cre- p53<sup>−/−</sup> or +/−.

| SC-Mdm2<sup>−/−</sup> p53<sup>−/−</sup> | Cre- p53<sup>−/−</sup> |
|---------------------------------|---------------------|
| #3638  fertile (2f)            | #3629  fertile (2f) |
| #3630  fertile (2f)            | #3631  fertile (2f) |
| #3634  sterile (4f)            | #3632  fertile (2f) |
| #3881  fertility (2f)          | #3879  fertile (2f) |

| SC-Mdm2<sup>−/−</sup> p53<sup>+/−</sup> | Cre- p53<sup>−/−</sup> |
|---------------------------------------|---------------------|
| #3880  sterile (4f)                   | #3990  sterile (4f) |
| #4248  sterile (4f)                   | #3991  sterile (4f) |
| #3989  sterile (4f)                   | #3988  sterile (2f) |
| #4933  sterile (4f)                   | #3633  sterile (2f) |

| SC-Mdm2<sup>−/−</sup> p53<sup>+/+</sup> | Cre- p53<sup>+/−</sup> |
|---------------------------------------|---------------------|
| #3992  sterile (4f)                   | #4934  sterile (2f) |
| #4245  sterile (4f)                   | #4246  sterile (2f) |
| #4783  sterile (4f)                   | #4247  sterile (2f) |
| #4935  sterile (4f)                   | #0922  fertile (2f) |

Males are contemporary.

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SC-Mdm2−/−. It is also noticeable that SC-Mdm2−/− adult testes produced low levels of androgens, according to the testosterone assay and the measurement of the expression of steroidogenic enzymes. This decrease in testosterone production also probably occurred perinatally, as a decrease in ano-genital distance was observed in SC-Mdm2−/− mice. Although ano-genital distance was recently reported to conserve some degree of plasticity at adulthood,31 most of it is determined by testosterone production during late fetal life. Altogether, this proves that the presence of SCs is critical for sustaining LC activity and the production of androgens during both fetal and post-natal life. Some androgens were probably produced during the development of SC-Mdm2−/− males, as the male genitalia were formed properly (e.g., presence of epididymis), and testicular descent, which is partially dependent upon androgens, occurred correctly. Testes are indeed formed in the abdominal cavity during fetal life and descend to reach a scrotal position in post-natal life. This depends on the production of two main LC hormones: first, INSL3, a peptide involved in the initial steps of the descent, and second, testosterone.32 We observed no cryptorchidy in SC-Mdm2−/− males, which proves that enough testosterone and INSL3 were produced during fetal life. Thus, this secretion of LC is probably not dependent upon SC regulation.

Quite surprisingly, we observed that the loss of SCs did not trigger an immediate extinction of the germ lineage. In SC-Mdm2−/−, numerous germ cells were retrieved in neonatal testes several days after SC loss. However, germ cells were completely lost at adulthood confirming the absolute requirement of a supporting lineage for the long-term maintenance of germ cells. Fetal SCs are known to govern the orientation toward the male fate of germ cells through the production of Fgf9 and Cyp26b1, and thus to prevent meiotic entry during fetal life.33 In this context, our report of a premature meiotic entry is not surprising. However, the sex of germ cells is determined between E12 and E14, so one would not expect meiotic entry owing to a late loss of SCs (i.e., at E15). We may thus consider that SCs are required both to impose and to maintain a male fate in germ cells. In addition, the seemingly proper differentiation of LC and the altered one of germ cells fits well with fetal LC differentiation occurring prior to germ cell differentiation (i.e., determination of fate during fetal life).

Very recently, another study successfully produced SC-deficient testes.34,35 This work used the diphtheria toxin expressed by SCs and the same AMH-Cre system as in our work. Our SC-Mdm2−/− model proposed here is complementary as it precludes the risk related to using a toxin with possible side effects. Moreover, Rebourcet and colleagues indicate a variable phenotype with, in most cases (95%), a more 60% reduction in the size of adult testes, possibly owing to a partial ablation of SCs in response to expression of the toxin. The invalidation of Mdm2 thus offers a more efficient and reproducible ablation of SCs, as we systematically obtained (around 30 males examined for the entire study) a 95% reduction in testis weight. The overall conclusions of both studies converge toward SCs being required to sustain germ and Leydig cell activity with a total loss of germ cells and the presence of a population of LC in adult testes. A slight difference related to the masculinization (e.g., differentiation of epididymis and vas deferens owing to testosterone production) was observed that did not seem to occur following expression of the toxin, whereas it is obvious following Mdm2 invalidation. We believe this may be due to the partial loss of SCs in the former system. Finally, these authors also reported the persistence of SOX9-positive cells in the rete area, as observed here, due to epithelial cells in the rete only partially expressing Cre.

As a whole, our work provides a clear demonstration of the crucial role of Mdm2/p53 in the regulation of SC apoptosis and opens new possibilities for studying the role of this cell type, thanks to efficient ablation during fetal life. Future works will need to characterize the precise kinetics of dysfunctions in LC in the absence of SCs.

Materials and Methods

Animals. Mice were fed a standard laboratory diet and tap water ad libitum, and maintained under 12 L:12D photoperiods in a temperature-controlled room (21–23 °C). All animal studies were conducted in accordance with the guidelines for the care and use of laboratory animals issued by the French Ministry of Agriculture and with the approval of a local ethical committee (number 2011-12-3, Comité d’Ethique en Expérimentation Animale Val de Loire – n°19). All efforts were made to minimize animal stress and suffering.

Generation of SC-Mdm2−/− and SC-Mdm2−/−p53−/− and their respective controls. Mdm2−/− mice were previously described by Grier et al.36 and contain a floxed Mdm2 allele with a loxP site in intron 4, followed by a neomycin cassette, and a second loxP site in intron 6. Recombination yields an Mdm2 allele lacking exons 5 and 6 (Mdm2Δeoxons5/6) after the Cre recombination event, resulting in a loss of most of the p53-binding domain.36 Mdm2−/− were crossed with heterozygous AMH-Cre mice on a C57BL/6 background, and the resulting Cre+Mdm2−/− offspring were backcrossed to Mdm2−/− mice to obtain Cre+Mdm2−/−, so called SC-Mdm2−/− controls were contemporary Cre-Mdm2−/− littermates, called Cre− here for simplicity.

Mice with the genotype Cre−Mdm2−/−p53−/− (null) were obtained from the Jackson laboratory (Sacramento, CA, USA) and were normally fertile. Cre+Mdm2−/− males were bred with p53−/− males, and the resulting Cre+Mdm2−/−p53−/− offspring were intercrossed to obtain homozygous mice for invalid Mdm2 alleles (Mdm2−/−), followed by intercrossing to obtain males with key genotypes: SC-Mdm2−/−p53−/−, SC-Mdm2−/−p53−/−, SC-Mdm2−/−p53−/−, and their respective contemporary littermate controls: Cre-p53−/−, Cre-p53−/−, Cre+p53−/−, Cre+p53−/−.

A high predisposition to malignancy was described for homozygous p53−/−, with a greatly accelerated rate of tumorigenesis.37 To avoid the risk of cancer, we used these mice for breeding and experiments before the age of 3.5 months.

Male fertility test. For each genotype of interest (SC-Mdm2−/−, SC-Mdm2−/−p53−/−, SC-Mdm2−/−p53−/−, SC-Mdm2−/−p53−/− and heterozygous SC-Mdm2−/−), each adult male (at least four per genotype) was mated with two or four primiparous Swiss female mice.

Mouse genotyping. PCR genotyping of Cre, Mdm2 and p53 alleles have been described previously (Lécureuil et al.,17 Montes et al.35 and Jones et al.16 respectively). DNA was extracted using the Tissue PCR Kit (Sigma-Aldrich, Saint Quentin Fallavier, France). GABR detection (200 bp) was used to control DNA quality. To verify that excision was restricted to the testes, DNA was extracted from various tissues, including testes from prepubertal mice, and PCR was performed using primers previously described36 to detect the deleted allele lacking exons 5 and 6 (Mdm2Δeoxons5/6): forward: 5′-TGGAGAAACAGTTACTTC-3′; reverse 5′-TGAGTGGACAGGCTCTG-3′. The annealing temperature was 50 °C.

Gene expression analysis. RNA extraction. Total RNA was isolated from whole testes (adult or P10) and from SC-enriched fractions obtained at P10 and prepared as described previously38 using RNeable reagent (Eurobio, Courtaboeuf, France). RNA (1 μg) was reverse-transcribed using the RNasey kit (QIAGEN, Courtaboeuf, France) according to the manufacturer’s instructions (the kit includes genomic DNA elimination with DNase). The sample with RNA but without RT (RT−) was the negative control.
RT-PCR for Mdm2 mRNA detection: Classic PCR was performed with Taq polymerase from Eurobio to detect Mdm2 mRNA in SC-enriched fractions. Mdm2 mRNA-specific primers (band at 180 bp), described in a previous study were forward 5'-CGGAAAGGCGCGGGAAGTA-3'; reverse 5'-TTCCGAGCTGGAARCTGTGAGGTGC-3'. The annealing temperature was 60 °C. For actin detection (band at 111 bp), used to control DNA quality, the primers were: forward 5' TAC GAC CAG AGG CAT ACA GG 3' reverse 5' TGA CCC AGA TCA TTG AGA 3'. The annealing temperature was 55 °C.

Quantitative real-time PCR analysis for the investigation of steroidogenesins: Three steroidalogenes genes were analyzed using the TaqMan assay with primers and probes invented by Applied Biosystems (Fisher Scientific, Illkirch, France). Normalization was performed using two internal standards, β-actin (Mm00472902_m1) and GAPDH (Mm9999915_g1) from the same sample, and the normalized cDNA was compared between the two genotypes.

Blood collection and hormonal assay. Around 500 μl of blood were obtained by retro-orbital sampling in anesthetized mice (with diazepam and ketamine), and collected in a tube containing EDTA.

Plasma testosterone levels: Mice were treated with an intraperitoneal injection of 15 IU per animal of human chorionic gonadotropin (Chorulon, MSD Santé Animale, Beaucouze, France). Blood was collected before (basal level) and 2 h after injection (stimulated level). The plasma was stored at −20 °C until tritium-based testosterone competitive radio-immunoassays, which are carried out regularly in our lab, were performed in our lab as described previously for other transgenic lines. The sensitivity of the assay was 0.125 ng/ml and the intraassay coefficient of variation was 7.5%. Briefly, samples (two dilutions per sample) or testosterone dilutions (to determine the range) were incubated for 1 h at 40 °C (0.1 M phosphate buffer with 0.1% gelatin) with tritiated testosterone plus the anti-testosterone antibody. A secondary antibody was then added and the mixtures incubated overnight at 4 °C. Immuno-precipitation was then performed with PEG (Polyethylene Glycol) 4000 and the radioactivity counted (Packard C2900, TriCarb, PerkinElmer, Villebon S/Yvette, France).

Plasma FSH levels: Plasma FSH levels were determined in 100 μl aliquots using a double Ab method and a RIA kit (rFSH RIA), kindly supplied by the National Institutes of Health (Dr A. F. Parlow, National Hormone and Peptide Program, Torrance, CA, USA). Rat FSH antigen (NIDDK-rFSH-I) was labeled with 125I using the chloramine-T method and the hormone concentration was expressed using the rat FSH reference preparation (NIDDK-rFSH-SP-2) as the standard. Intraassay and interassay coefficients were less than 7% and 10%, respectively. The sensitivity of the assay was 0.125 ng/100 μl.

Plasma LH levels: Plasma LH levels were determined in 100 μl aliquots using a double Ab method and a RIA kit (mLH Ria), kindly supplied by the National Institutes of Health (Dr A. F. Parlow, National Institute of Diabetes and Digestive and Kidney Diseases, National Hormone and Peptide Program). Rat LH-110-1 (AFP-11536B) was labeled with 125I using the chloramine-T method and the hormone concentration was expressed using the mouse LH reference preparation (AFP-S306-A) as the standard. Intraassay and interassay coefficients were less than 7% and 10%, respectively. The sensitivity of the assay was 4 pg/100 μl.

Histology and immuno-detection of AMH, DDX4, 3βHSD, C-CAS-PASE3, SYCP3 and p53: Testis histology (semiferous tubule organization) was analyzed after fixing in Bouin’s fluid and embedding in paraffin. For microscopic observation, 4-μm sections were stained with hematoxylin.

Immunostaining was performed as described previously with minor modifications. Fetal, postnatal and adult testes were fixed overnight in 10% neutral formalin (Carlo Erba Reagents, Val de Reuil, France) before being dehydrated, embedded in paraffin wax and cut into 5-μm sections. Adult testes were stained with hematoxylin and eosin; fetal and postnatal testes were used for immunostaining. The primary antibodies used in this study were either rabbit polyclonal anti-DEAD/H box polypeptide 4 (DDX4/VASA; 1:200, Abcam, Paris, France), rabbit polyclonal anti-cleaved caspase-3 (1:200, Cell Signaling/Ozyme, Montigny-Le Bretonneux, France), goat polyclonal anti-AMH (1:200, Santa Cruz Biotechnology, Heidelberg, Germany), mouse monoclonal anti-BetaHSD (1:200, Transgenic Inc., Chuo-Ku, Kobe, Japan), mouse monoclonal anti-SYCP3 (Abcam 1/2000), mouse monoclonal anti–P21 (1:200, BD Biosciences, Le Pont de Claix, France) or rabbit polyclonal anti-TP53 (1/500, Novoceastra, Leica Microsystems, Nanterre, France). The same protocol was used for antigen retrieval. After testis section dewaxing and rehydration, antigen retrieval was performed in HIER citrate buffer pH 6 (Zytomed, Diagnostics, Blagnac, France) in an autoclave (Reverter 2100, Proteogénie, Mundolsheim, France). Sections were then washed in distilled water and incubated for 15 min in 3% H2O2 at room temperature. After 30 min in 2.5% normal Horse serum (Vector laboratories, EuroBio, Les Ulis, France), primary antibodies diluted in PBS were incubated for 1 h at 37 °C. Primary antibodies were revealed using the secondary antibody IMPRESS kit (Vector Laboratories, EuroBio). Peroxidase activity was visualized using either 3,3’-diaminobenzidine (brown) or vip (purple) as a substrate. Sections were counterstained with hematoxylin. TUNEL staining was performed using the Apoptag peroxidase kit (MP Biomedicals, Illkirch, France) in accordance with the manufacturer’s recommendations. For cell counting, at least 300 cells were counted in three different sections for each testis. For immunofluorescence detection, primary antibodies were detected with specific secondary antibody species Alexa Fluor 488- and Alexa Fluor 594-conjugated antibodies (1:500, Life Technologies, Life Technologies, Euromedex, Saint-Aubin, France).

Statistical analysis. All data are presented as means ± S.E.M. To compare means between two groups, the Student’s t-test or the Mann-Whitney U test, in case of differences in variance (Fisher test), was used. Other comparisons were performed using a two-way ANOVA followed by the Bonferroni post-test. P < 0.05 was considered significant.

Conflict of Interest

The authors declare no conflict of interest.

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