Quantitative Proteomics Reveals the Essential Roles of Stromal Interaction Molecule 1 (STIM1) in the Testicular Cord Formation in Mouse Testis*

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Testicular cord formation in male gonadogenesis involves assembly of several cell types, the precise molecular mechanism is still not well known. With the high-throughput quantitative proteomics technology, a comparative proteomic profile of mouse embryonic male gonads were analyzed at three time points (11.5, 12.5, and 13.5 days post coitum), corresponding to critical stages of testicular cord formation in gonadal development. 4070 proteins were identified, and 338 were differentially expressed, of which the Sertoli cell specific genes were significant enrichment, with mainly increased expression across testis cord development. Additionally, we found overrepresentation of proteins related to oxidative stress in these Sertoli cell specific genes. Of these differentially expressed oxidative stress-associated Sertoli cell specific protein, stromal interaction molecule 1, was found to have discrepant mRNA and protein regulations, with increased protein expression but decreased mRNA levels during testis cord development. Knockdown of Stim1 in Sertoli cells caused extensive defects in gonadal development, including testicular cord disruption, loss of interstitium, and failed angiogenesis, together with increased levels of reactive oxygen species. And suppressing the aberrant elevation of reactive oxygen species could partly rescue the defects of testicular cord development. Taken together, our results suggest that reactive oxygen species regulation in Sertoli cells is important for gonadogenesis, and the quantitative proteomic data could be a rich resource to the elucidation of regulation of testicular cord development. Molecular & Cellular Proteomics 14: 10.1074/mcp.M115.049569, 2682–2691, 2015.

Male gonadogenesis is a complex process that requires the formation and assembly of several cell types that come together to form a functional organ. These cell lineages coordinate to maintain testicular cord development but do not differentiate independently (1, 2). Shortly after the activation of Sox9, when the genital ridges are still long and very thin, pre-Sertoli cells start to aggregate around germ cell clusters and form cords; they are then referred to as Sertoli cells. Partitioning of this mass of cells into cord-forming units coincides with endothelial cell invasion and expansion of interstitial space (3, 4). In mice, organization of the testicular cords begins with aggregate of germ cell and Sertoli progenitors in the gonad. Previous studies using confocal analysis and three-dimensional modeling have reported that testicular cord formation involves three basic steps (5, 6): pre-Sertoli cells and germ cells coalesce between 10.5 and 12.5 days post coitum (dpc); cords partition at 12.5 dpc with a clear basal lamina surrounding the cords, and all cords are characterized as “external” cords, defined as a single transverse loop located just under the celomic epithelium that surrounds the gonad at this stage; and refinement of cords continues at 13.5 dpc. Although Sertoli cells acting as an organizing center in testicular cord formation has been well known (3) and studies in knockout mouse models have revealed several genes associated with testicular cord formation (7–10), how these cell types assemble into a functional organ remains to be explored (2, 11).

Proteomics technology has been widely used in postnatal testis development and function research in mice (12–16). Two proteomics studies have been carried out in the fetal gonads in mice, and identified more than 1000 proteins expressed in gonads (17, 18), however, the temporal proteome changes have not been elucidated during gonadogenesis. Additionally, mRNA abundance may not always predict the quantity of the corresponding functional protein, and pro-

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* The abbreviations used are: dpc, days post coitum; IF, immunofluorescence; ROS, reactive oxygen species; TMT, tandem mass tag; SCX, strong cation exchange; DE, differentially expressed; STIM1 (Stim1), stromal interaction molecule 1; NAC, N-acetylcysteine.

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Proteomic approach can provide a systemic view of protein level regulation in a large scale (18). Therefore, this study aimed to obtain a better understanding of male gonadogenesis by establishing a first temporal proteomic profile during the initiation of gonad development in male mice. After confirming the specific time point by immunofluorescence (IF) staining, we performed a comparative proteomic analysis of samples of male mouse gonads obtained at 11.5, 12.5, and 13.5 dpc. Bioinformatics analysis and functional studies demonstrate that reactive oxygen species (ROS) regulation in Sertoli cells may be important for testicular cord formation, and functional characterization of the stromal interaction molecule 1 (stim1), a Sertoli cell specific protein, supported this hypothesis. Our categorized protein lists can serve as a useful resource for further exploring the molecular mechanisms involved in gonadal development.

**EXPERIMENTAL PROCEDURES**

**Animals and Sample Collection**—All animal experiments were approved by the Ethics Committee of Nanjing Medical University (China). Timed mating of the ICR strain with noon of the day on which the mating plug was observed designated 0.5 dpc. The sex of the embryo was determined by observation at 12.5 and 13.5 dpc. At 11.5 dpc, the embryos were genotyped to determine the sex using primers to detect Sry, and β-actin was used as an internal control. The following primers were used: Sry, GTTCAGGCTCATACGGAC (forward) and CCAGGGGCACACCTTAA (reverse); β-actin, GGGGTTATCCTCCATCAG (forward) and TCCTATGGAAGGGCACA (reverse). Approximately 800 gonad pairs at 11.5 dpc, 500 gonad pairs at 12.5 dpc, and 500 gonads at 13.5 dpc (with meseocephri removed) were resected from male mouse embryos for use in the proteomics screen.

**Gonad Culture**—Gonads of 12.0 dpc embryos were isolated by dissection in physiological saline at room temperature, and collected into warmed Dulbecco’s Minimal Eagle’s Medium (DMEM) containing 10% fetal calf serum (FCS) and 50 μg/ml ampicillin at 37 °C with 5% CO2. Stim1-Translation-Blocking-Vivo-Morpholino (Oligo Sequence CAAACGGGCGCAGACATCTGAC) and Standard Control Oligo (Oligo Sequence CTTCTACCTCAGGATGATTATA) were purchased from Gene Tools, LLC. After culture for 30 h, gonads were collected for further study. ROS and AT detection were performed by ROS advanced fluoro assay kit (Gennmed Sciences, DE) and ATP Assay Kit (Beyotime, Beijing, China).

**Protein Extraction and Digestion**—Gonads from 11.5, 12.5, and 13.5 dpc were kept separate and collected in ice-cold PBS, and proteins were extracted using protein extraction buffer consisting of 7 M urea, 2 M Thio urea, 65 mM dithiothreitol (DTT) and 1% (v/v) protease inhibitor mixture. After incubation for 1 h on ice (vortexing every 10 min), the supernatants were collected by centrifugation at 40,000 × g for 60 min at 4 °C. The protein content of the supernatants was measured using Bradford assay. Prior to tandem mass tag (TMT) labeling, 70 μg protein was used for trypsic digestion for each TMT Label reagent. After reaction at 37 °C with trypsin in a 1:50 enzyme:protein ratio. The experiments were repeated for four times, with 12 samples in total for gonads of three stages.

**TMT Labeling**—TMT 6-plex labeling was performed according to the manufacturer’s protocol with minor modifications. In brief, TMT Label Reagents were equilibrated to room temperature. Each aliquot was suspended in 41 μl of anhydrous acetonitrile, and 42 μl of the reagent was added to the digestion dissolved in 200 mM triethylammonium bicarbonate (TEAB). After reaction at room temperature for 1 h, 8 μl of 5% hydroxylamine was added to each tube, and the mixture was incubated for another 15 min. The aliquots were then combined, and the pooled sample was evaporated in a vacuum. A total of two TMT six-plex labeling experiments were performed, and each labeled six samples involving two replicates.

**Strong Cation Exchange (SCX) Fractionation**—The labeled peptide mixture was resuspended in SCX chromatography Buffer A (10 mM NH4COOH and 5% ACN; pH 2.7) and loaded onto an SCX column (1 mm internal diameter × 10 cm, packed with Poros 10S; Dionex, Sunnyvale, CA) in an UltiMate® 3000 HPLC systems at a flow rate of 50 μl/min. The following linear gradient was used: 0% to 56% B (800 mM NH4COOH and 5%ACN, pH 2.7) in 30 min; 56% to 100% B for 1 min; 100% B for 3 min; 100% to 0% B for 1 min; and 0% B for 20 min before the next run. The effluents were monitored at 214 nm based on the UV light trace, and fractions were collected every 2 min (19–21).

**Mass Spectrometry Analysis**—Seventeen fractions were sequentially loaded onto a µ-proclemellt® cartridge (0.3 × 5 mm2, 5 μm, 100 Å; Dionex) at a flow rate of 20 μl/min. The trap column effluent was then transferred to a reverse-phase microcapillary column (0.075 × 150 mm2; Acclaim® PepMap100 C18 column, 3 μm, 100 Å; Dionex). The reverse-phase separation of peptides was performed using buffer A (2% ACN and 0.5% acetic acid) and buffer B (80% ACN and 0.5% acetic acid); a 193-min gradient (4% to 9% buffer B for 3 min, 9% to 33% buffer B for 170 min, 33% to 50% buffer B for 10 min, 50% to 100% buffer B for 1 min, 100% buffer B for 8 min, and 100% to 4% buffer B for 1 min) was used.

Peptides were detected on LTQ Orbitrap Velos (Thermo Fisher Scientific, Waltham, MA) using a data-dependent acquisition mode. The MS3 method was programmed as previously described (16). For each cycle, one full MS scan of mass/charge ratio (m/z) = 350–1800 was acquired in the Orbitrap at a resolution of 60,000. After each full scan, eight of the most intense ions were selected for collision-induced dissociation (CID) fragmentation (CE 35%), and the most intense product ion from the MS2 step was selected for higher-energy collisional dissociation (HCD) fragmentation (NCE 68%). Wideband activation was enabled. Lock mass, with siloxane (m/z = 445.120025), was used in all runs to calibrate orbitrap MS precursor masses. The mass spectrometry proteomics data are available via ProteomeXchanger with identifier PXD002583.

**Protein Identification and Quantification**—The raw files were processed with MaxQuant (22) software (version 1.2.2.9) using the International Protein Index (IPI) (23) mouse proteome database (version 445.120025), was used in all runs to calibrate orbitrap MS precursor masses. The mass spectrometry proteomics data are available via ProteomeXchanger with identifier PXD002583.

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groups, the cutoffs of fold change and Q value were set to 2 and 0.01, respectively, using the significance analysis of microarrays (SAM) algorithm performed by J-expression (25).

**Western Blot Analysis**—Western blotting was performed as described previously (14, 26) with a modification. Briefly, the proteins were separated by SDS-PAGE and transferred onto a polyvinylidene difluoride membrane. The membrane was blocked with 5% nonfat milk in TBST solution for 2 h at room temperature, and incubated overnight at 4 °C with primary antibodies. The membranes were washed with TBST buffer three times and incubated at room temperature for 2 h with secondary antibodies. The signals of the detected proteins were visualized on SuperSignal Western blotting detection system (Thermo Scientific).

**IF Staining**—Paraffin-embedded gonad sections were dewaxed and dehydrated. Endogenous peroxidase activity was arrested by incubating the sections with 3% hydrogen peroxide for 15 min. Antigen retrieval was performed by boiling samples in 0.01 M sodium citrate buffer (pH 6.0) for 10 min. The sections were incubated with 5% bovine serum albumin in PBS for 2 h at room temperature. The sections were incubated with primary antibody at 4 °C overnight. After washing with PBS, secondary antibody was applied for 2 h, and the nucleus was stained with DAPI. For whole mount staining, gonads were fixed with 4% paraformaldehyde for 30 min, blocked with 5% bovine serum albumin for 2 h and incubated with primary antibodies overnight at 4 °C, as described previously (16). On the second day, the gonads were washed thrice, and incubated with secondary antibody for 2 h, mounted on slides, and examined. Cell counting was performed by previous reported method (27, 28). Briefly, images of the three most interior sections of each whole mount-stained gonad were collected using the confocal microscope (LSM 710, Carl Zeiss, Oberkochen, Germany). To avoid counting the same cells in consecutive sections, sections were spaced at least 5 mm intervals. Cells with positive signals were counted in three sections and averaged for each gonad. Sample number for each group was at least 3.

**RESULTS**

**Identification of Specific Time Points During the Formation of Testicular Cords**—IF analysis was used to identify the dynamic changes involved in early testis development (Fig. 1A). Germ cells were labeled with DDX4 and OCT4; and Sertoli cells, with SOX9. At 11.5 dpc, pre-Sertoli cells labeled by SOX9 and germ cells labeled by OCT4 and DDX4 were evenly distributed. From 12.5 dpc, germ cells started forming clusters (asterisks, yellow) and were surrounded by Sertoli cells. At 13.5 dpc, Sertoli cells enclosed germ cell populations in distinct testicular cords (asterisks, white). Scale bars: 50 μm. B, Totally 45 DE proteins were in the cluster of Pattern 1, 94 in Pattern 2, and 22 in Pattern 3, as well as 177 in Pattern 4. For each stage, there were 4 replicates.

![Fig. 1. Expression patterns of differentially expressed proteins during testicular cords formation. A, At 11.5 dpc, pre-Sertoli cells labeled by SOX9 and germ cells labeled by OCT4 and DDX4 were evenly distributed. From 12.5 dpc, germ cells started forming clusters (asterisks, yellow) and were surrounded by Sertoli cells. At 13.5 dpc, Sertoli cells enclosed germ cell populations in distinct testicular cords (asterisks, white). Scale bars: 50 μm. B, Totally 45 DE proteins were in the cluster of Pattern 1, 94 in Pattern 2, and 22 in Pattern 3, as well as 177 in Pattern 4. For each stage, there were 4 replicates.](image-url)
supplemental Fig. S1, the largest proportion of DE proteins identified was annotated as cytoplasm (191), followed by membrane (138), nucleus (85), and mitochondrion (52).

Based on expression patterns, using k-means clustering algorithm (31, 32), these 338 DE proteins were classified into four clusters (supplemental Table S3) by J-Express software (25). The four clusters indicated four distinct expression patterns (Fig. 1B): increase at 13.5 dpc (45 proteins, Pattern 1), decrease at 12.5 dpc and remain unchanged at 13.5 dpc (94 proteins, Pattern 2), increase at 12.5 dpc but decrease at 13.5 dpc (22 proteins, Pattern 3), and steady increase across all time points (177 proteins, Pattern 4).

In this study specific cell lineages were not separated, because of the large amount of proteins required for proteomic studies, it is difficult to get enough proteins of specific cell lineage in embryonic gonad for proteomics analysis. However, Jameson et al. analyzed transcriptome of each cell lineage in embryonic testis, and according to mRNA expression values, they assembled the lineage specific genes that were specifically enriched in each cell lineage, including Sertoli, interstitial cells, germ cells, and endothelial cells, relative to the other lineages in embryonic testis (33). Although mRNA expression does not necessarily translate to protein levels, studies have shown that as high as 65–79% (correlation of $r^2$, 0.65–0.79) of the protein level changes can be explained by mRNA levels (34, 35). The lineage-specific gene expression at mRNA levels implies potential important functions of certain gene in the corresponding lineage. Analysis of the distribution of these lineage specific genes showed that the lineage specific genes tended to be differentially-expressed across developmental stages at protein level, there was significant enrichment of lineage specific genes as a total in DE proteins compared with non-DE proteins (31.66% versus 24.14%, $p < 0.01$, Fig. 2A). Furthermore, we analyzed the distribution of lineage specific genes from each lineage separately (supplemental Table S4) and found that for the Sertoli and interstitial cell specific genes, there was significant enrichment in the DE proteins compared with those in non-DE proteins (9.47% versus 5.33%, $p < 0.01$ and 6.21% versus 1.18%, $p < 0.001$, respectively, Fig. 2B, 2C). And no significant enrichment was observed for endothelial cell and germ cell specific genes ($p > 0.05$, supplemental Fig. S2). In addition, among the DE proteins, Sertoli cell specific genes (69%) and interstitial cells specific genes (48%) mainly have steady increased expression during testicular cord formation at protein level (Pattern 4; Fig. 2D, 2E), which indicated the importance of these proteins in testicular cord formation. To verify results of the TMT quantification, several proteins with commercially available antibodies from the cluster of Pattern 4 were chosen for Western blot analysis. As shown in supplemental Fig. S3, Western blot and TMT quantification showed consistent results. Thus, this quantitative proteomic results are of high confidence.

As Sertoli cell and interstitial cell specific genes are enriched in DE proteins during testis development. We annotated the functions of these proteins according to different expression patterns by the Pathway Studio software, and found that in Pattern 4, there was significant enrichment of oxidative stress-associated proteins in proteins with Sertoli cell specific gene expression compared with those in the rest
of Pattern 4 proteins (31.82% versus 9.68%, \( p < 0.01 \)) (Fig. 2F, supplemental Table S5). In the interstitial cell lineage, however, there was no such enrichment. These data suggest that regulation of oxidative stress in Sertoli cells might be important in the regulation of testicular cord formation during male gonadal development.

Expression and Distribution of Stim1 in Male Gonads—As bioinformatics analysis showed possible roles of oxidative stress regulation in testicular cord formation, we selected STIM1, an oxidative stress-associated protein from cluster of Pattern 4 of steady increased expression with enrichment of Sertoli cell specific proteins, for further functional study. Western blot data indicated that, at protein level, STIM1 was weakly expressed at 11.5 dpc, but increased at 12.5 and 13.5 dpc (Fig. 3A), which was consistent with our TMT quantification. However, in contrast to its protein level, we found that Stim1 decreased at 13.5 dpc at mRNA level according to real-time reverse transcriptase polymerase chain reaction (real-time PCR) analysis (Fig. 3B). In addition, our real-time PCR results were highly consistent with Jameson’s microarray data (Fig. 3C). Together, these results showed post-transcriptional regulations during testicular cord development, and that mRNA and protein of the same gene may exhibit discrepant or even reversed temporal expression patterns, with Stim1 as a good example of this phenomenon. Thus, it is crucial to study the functions of genes at protein level, and mRNA level studies only are not enough to elucidate mechanisms of male gonadogenesis.

Next, we performed IF staining to analyze the STIM1 distribution in embryonic testis. As shown in Fig. 3D, STIM1 is a Sertoli cell specific protein, and is mainly expressed in the cytoplasm of Sertoli cells. Quantification of the fluorescence intensity of STIM1 (Fig. 3E) showed similar trend of expression changes during embryonic testis development as those shown in the Western blot (Fig. 3A) and the TMT quantification.

Suppression of Stim1 causes severe phenotypes in gonadogenesis—To explore the function of Stim1 in gonadogenesis, we used morpholino oligo against Stim1, and successfully suppressed Stim1 translation in vitro. Western blotting analysis revealed about 55% suppression efficiency (Fig. 4A, 4B).

After Stim1 knockdown, severe disruption in testicular cord development was observed. As shown in Fig. 5, the testicular cords in the knockdown group were disorganized and showed significant reduction in the number of Sertoli cells (labeled by SOX9 and WT1; Fig. 5A, 5B, 5G, 5H), and germ cells (labeled by OCT4 and DDX4; Fig. 5A–5C, 5E, 5f, 5J).
Moreover, to test whether knockdown of Stim1 in Sertoli cells could affect the development of endothelial cells and interstitial cells, we also examined the histological architecture of PECAM1-labeled endothelial cells (Fig. 5D) and 3β-HSD-labeled interstitial cells (Fig. 5E). Both cell types were malformed and decreased in numbers after Stim1 knockdown in Sertoli cells. To determine the reason for this reduction in cell number, we detected cell proliferation and apoptosis by detecting the expression levels of cleaved KI67 and CASPASE-3, respectively. There was a significantly higher number of cleaved CASPASE-3-positive cells in the knockdown group (Fig. 5F, 5K) than in the normal and control groups.

Fig. 4. A, Verification of 5 μm translation-blocking vivo-morpholino interference efficiency on the STIM1 protein level. B, Western blot results showed about 55% suppression efficiency. Statistical analysis was performed by one way ANOVA. *** p < 0.001.

Fig. 5. Suppression of Stim1 led to defects in testicular cord formation. In the Stim1 knockdown group, there was an obvious reduction in the Sertoli cells labeled by SOX9 (A) and WT1 (B) and germ cells labeled by OCT4 (A, B) and DDX4 (C, E). Statistical data were displayed in (G–J). Stim1-deficient gonads showed failed angiogenesis (D) and development of interstitial cells labeled by 3β-HSD (E). PECAM1 was distributed in both germ cells (D, asterisks) and endothelial cells (D, arrows). Cell apoptosis test results showed that the cleaved CASPASE3-positive cells were significantly higher in the Stim1 knockdown group (F, arrows), compared with the normal and control groups. Statistical data were presented in (K). Statistical analysis was performed by one way ANOVA. n= 3. Scale bars: 50 μm.*** p < 0.001.
groups, whereas there was no obvious change in the number of KI67-positive cells between the knockdown, normal, and control groups (supplemental Fig. S4A, S4B), suggesting that the reduction in the cell number was a result of cell apoptosis.

**Suppression of Stim1 Showed no Obvious Change in Phenotype in Early Female Gonads**—Due to the phenotypes caused by Stim1 knockdown in male gonads, we wondered whether Stim1 played a role in female gonads. Interestingly, we found no obvious morphological changes in the female knockdown group compared with the normal and control female gonads (supplemental Fig. S5A, S5B). Western blot analysis showed different expression patterns of STIM1 between XY and XX gonads, with an up-regulated expression pattern in XY (Fig. 3A) and a down-regulated expression at 13.5 dpc in XX (supplemental Fig. S6). These results indicate that Stim1 might not be required for female gonads differentiation at this stage.

**Reduction of Increased Oxidative Stress Can Partially Rescue Phenotypes Caused by Stim1 Knockdown**—As Stim1 is related to oxidative stress, we sought to assess whether oxidative stress could contribute to the phenotypic defects. The ROS content was markedly higher in the knockdown group than that in the normal and control groups (p < 0.001) (Fig. 6A). Furthermore, the antioxidant response genes Gclc, Gclm, Homx1, Prdx3, and Ucp2 were significantly up-regulated in the knockdown group (Fig. 6B), suggesting that Stim1 deficiency causes constitutive oxidative stress. However, when the knockdown group was treated with the antioxidant scavenger N-acetylcysteine (NAC), the ROS levels markedly decreased in the knockdown group, although they did not...
Western blot data indicated that β-catenin, ERK, and P38 were activated, and NAC treatment reduced their expression levels (A). Akt and NF-κB pathways showed no apparent alterations in the knockdown and NAC treated knockdown groups (B).

reach the normal levels (Fig. 6A). And addition of NAC led to a reduction in the expression levels of the antioxidant response genes, although the reduction in the Gclc and Ucp2 levels was not significant (Fig. 6B). The defects in the testicular morphology were partially rescued after treatment with NAC (Fig. 6C–6I). The numbers of Sertoli and germ cells in the NAC-treated knockdown group were significantly higher than those in the knockdown group (Fig. 6C, 6G, 6H). The numbers of endothelial (Fig. 6D) and interstitial (Fig. 6E) cells were higher in the NAC-treated knockdown group than in the knockdown group. Additionally, we observed that the number of cleaved CASPASE-3-positive cells was lower in the NAC-treated knockdown group than in the knockdown group (Fig. 6F, 6I). Together, these results demonstrate that the aberrant elevation of ROS can cause defects in testis development, whereas the reduction of ROS can partially rescue them.

ROS is well known to activate many important signaling pathways including mitogen-activated protein kinase (MAPK) cascades, nuclear factor-kB (NF-kB), P-I-3-kinase-Akt, and wnt/β-catenin (36–38). We found that β-catenin, ERK, and P38 were activated (Fig. 7A), whereas NAC addition reduced their expression levels. The Akt and NF-kB pathways showed no obvious changes in both knockdown and NAC treatment knockdown groups (Fig. 7B).

**DISCUSSION**

Emerging evidence has illustrated that Sertoli, interstitial, and endothelial cells all participate in the development of testicular cords. Deletions of genes that are specifically expressed in the Sertoli and interstitial cells have been shown to cause severe disruption of testicular cords, and blocking the migration of endothelial cells can impede organization of the testis (10, 39, 40). Testicular cords appear normal in mutant mice lacking germ cells, indicating that germ cells are not vital for cord formation (41). In order to get a systemic view of regulation of testicular cord formation at protein levels, we generated a comparative proteomic profile of male mouse gonads at three time points spanning the critical period for testicular cord formation. Several of the DE proteins obtained in our profile have been previously reported to function in gonadal development (supplemental Table S6). For example, Amh (anti-Müllerian hormone) is crucial for regression of the Müllerian duct and for the promotion of testis differentiation (42, 43).

Comparison with lineage specific genes compiled by Jameson et al. (33) based on mRNA expression data showed that Sertoli cell specific genes were enriched in Pattern 4 with steady increased expression. Oxidative stress related proteins were over-represented in Sertoli cell specific DE proteins; therefore, these results suggest that oxidative stress regulation in Sertoli cells may be important to the formation of testicular cords. In order to verify this hypothesis, STIM1, a Sertoli cell specific protein in Pattern 4 related to oxidative stress regulation, was selected for further study. STIM1 is a transmembrane ER protein containing multiple functional domains, and it serves as an ER Ca²⁺ sensor and activator of store-operated Ca²⁺ entry (SOCE) (44–46). Studies have shown that Stim1 is associated with oxidative stress and mitochondrial bioenergetics (47). We found that suppression of Stim1 in Sertoli cells in the cultured testis in vitro increased ROS production and caused a global disruption of gonads, evidenced by disorganized testicular cord formation, obstruction of angiogenesis, and loss of interstitium. The severe phenotypes observed by Stim1 knockdown demonstrated that Stim1 in Sertoli cells played an essential role in testicular cord formation. Further study indicated that molecules downstream of ROS generation, such as β-catenin, ERK, and P38, were significantly activated in the knockdown group, and treatment of the knockdown group with the antioxidant scavenger NAC could inhibit ROS production, consequently down-regulating the expression of β-catenin, ERK, and P38, and partially rescuing the defects in testicular cord formation.

ROS is an essential regulator of cell processes and is generated in all cells. Although ROS is well known to be toxic, they also function as signaling molecules (48). Oxidative stress during spermatogenesis is generally associated with male infertility. Mice deficient in the transcription factor nuclear factor-erythroid 2-related factor 2 (Nrf2), which regulates the basal and inducible transcription of genes encoding enzymes important for protection against ROS, have been
shown to have disruptive spermatogenesis in an age-dependent manner (49), whereas moderate ROS levels are beneficial for the self-replacement of mouse spermatogonial stem cell (SSC) (50). In humans, physiological ROS is beneficial for the cyclic waves of spermatogenesis, whereas excess ROS are harmful and can cause human infertility (51–53). Our present study indicated that aberrant elevation of ROS caused by suppression of STIM1 in Sertoli cells in early mouse gonads could impair testicular cord formation.

Wnt/β-catenin signaling is one of the fundamental mechanisms that direct cell proliferation, cell polarity, and cell fate specification during embryonic development and tissue homeostasis (54, 55). Activation of the Wnt pathway leads to transcription of downstream target genes such as Myc and Cyclin D1 (56, 57). In the last few years, the role of β-catenin in fetal testis development has been gradually established (58). It has been shown that the constitutively active form of β-catenin in embryonic Sertoli cells in mouse causes testis malfomation, including testicular cord disruption and germ cell depletion (59, 60). It should be noted that our ex vivo gonad culture experiments are not substitutes of in vivo experiments, the use Stim1 knockout mouse models are preferred in future studies in order to elucidate its functions in testis development. With the ex vivo studies, our results demonstrated that Stim1 could regulate β-catenin in gonadal development by ROS signaling.

In conclusion, our comparative proteomic profile of mouse testicular cord formation and functional studies revealed the important roles of oxidative stress regulation in Sertoli cells in the testicular cord formation during early testis development.

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§ This article contains supplemental Tables S1 to S6 and Figs. S1 to S6.

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