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HIF 1 inhibits STAR transcription and testosterone synthesis in murine Leydig cells

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Abstract

Hypoxia-inducible factor-1 (HIF1) is a critical transcription factor involved in cell response to hypoxia. Under physiological conditions, its ‘a’ subunit is rapidly degraded in most tissues except testes. HIF1 is stably expressed in Leydig cells, which are the main source of testosterone for male, and might bind to the promoter region of steroidogenic acute regulatory protein (STAR), which is necessary for the testosterone synthesis, according to software analysis. This study aims to identify the binding sites of HIF1 on Star promoter and its transcriptional regulation of STAR to affect testosterone synthesis. Testosterone level and steroid synthesis-related proteins were determined in male Balb/C mice exposed to hypoxia (8% O2). While HIF1 was upregulated, the testosterone level was significantly decreased. This was further confirmed by in vitro experiments with rat primary Leydig cells or TM3 cells exposed to hypoxia (1% O2), CoCl2 or DFX to raise HIF1. The decline of testosterone was reversed by pregnenolone but not cAMP, indicating the cholesterol transport disorder as the main cause. In agreement, STAR expression level was decreased in response to HIF1, while 3b-hydroxysteroid dehydrogenase, 17b-hydroxysteroid dehydrogenase and p450scc did not exhibit significant changes. By ChIP, EMSA supershift and dual-luciferase reporter assays, HIF1 was found to bind to the Star promoter region and repress the expression of STAR. Mutation assays identified three HIF1-binding sites on mouse Star promoter. These findings indicate that HIF1 represses STAR transcription through directly binding to the Star promoter at −2082/−2078, −2064/−2060 and −1910/−1906, leading to the negative regulation of testosterone synthesis.

Introduction

Hypoxia-inducible factor-1 (HIF1), a critical transcription factor, regulates cellular hypoxic response in oxygen-lacking cells and is rapidly degraded through the ubiquitin-proteasome pathway under normoxic conditions (Semenza 2011, 2012, 2017, Ivan & Kaelin 2017). HIF1 consists of two subunits, the oxygen-dependent HIF1α and the stably expressed HIF1β (Wang et al. 1995). HIF1α is hydroxylated by prolyl hydroxylase domain-containing protein 2 (PHD2) and then bound by the VHL protein, which recruits ubiquitin ligase that targets HIF1α for proteasome
degradation under well-oxygenated conditions (Ivan et al. 2001). PHD2 uses O$_2$ and a-ketoglutarate as substrates for hydroxylation. Thus, oxygen deprivation stabilizes HIF1 protein. HIF1α is also induced by deferoxamine (DFX) and cobalt chloride (CoCl$_2$), which inhibits hydroxylases via removing Fe(II) from the catalytic center (Kotake-Nara & Saida 2006). It has been reported that HIF1 is associated with the initiation and development of certain diseases, including hereditary erythrocytosis, cancer, traumatic shock, pulmonary arterial hypertension and obstructive sleep apnea (Semenza 2012).

Under normal physiological conditions, HIF1α is unstable in most tissues and degrades rapidly with sufficient oxygen. Thus, current knowledge about how HIF1 functions was mostly achieved in tumor cells or other pathologically hypoxic cells. One interesting phenomenon is that HIF1α can be stable under physiological conditions in the tests. Marti et al. first reported that HIF1α is expressed in spermatocytes, spermatids and spermatozoa in mouse testes (Marti et al. 2002). A following study reported that low partial pressure of oxygen in the testes was a stimulant of HIF1α and that HIF1α might play a role in spermatogenesis (Maxwell & Salnikow 2004). It was also found that HIF1α is expressed in rat testes and increased with the period of testicular ischemia (Powell et al. 2002). Furthermore, Palladino et al. identified Leydig cells, adjacent to the seminiferous tubules in the testis, as the major source of HIF1α in both normoxic and hypoxic testes, suggesting that HIF1 might play a critical role in Leydig cell functions (Palladino et al. 2011). However, the mechanism of HIF1 affecting/regulating Leydig cell functions is not clearly elucidated.

The main function of Leydig cells is synthesizing and secreting testosterone, producing 95% of the total amount in males (Svechnikov et al. 2010). Testosterone plays crucial roles in the development of secondary sex characters, maintenance of bone and sex drive (Mooradian et al. 1987). Testosterone deficiency could cause cardiovascular disease, erection dysfunction and male hypogonadism (Basaria 2014, Kloner et al. 2016, Skakkebaek et al. 2016, Elkhoury et al. 2017, Holgersson et al. 2017). Several factors such as age (Nguyen et al. 2015), exercise (Vingren et al. 2010), nutrients (Livera et al. 2002) and bodyweight (Koehler et al. 2009) affect the serum testosterone level and sexual function. Physiological testosterone level is strictly regulated by the hypothalamic–pituitary–testicular axis (Swerdloff et al. 1992). Luteotropic hormone (LH) secreted from the pituitary binds to LH/human chorionic gonadotropin (hCG) receptor and stimulates protein kinase A (PKA) activation through cAMP in Leydig cells. PKA phosphorylates steroidogenic acute regulatory protein (STAR), which transports cholesterol into mitochondria, where cholesterol is transformed into pregnenolone under the catalysis of P450 cholesterol side chain cleavage (P450scc). Pregnenolone is converted to testosterone under the catalysis of a series of enzymes, including 3β-hydroxysteroid dehydrogenase (3b-HSD) and 17b-HSD in the smooth endoplasmic reticulum (Häggström & Richfield 2014). Here, STAR-mediated transportation of cholesterol into mitochondria is the rate-limiting step, and thus, STAR is the rate-limiting protein for testosterone synthesis (Miller & Bose 2011). Several proteins have been described to functionally interact with STAR for cholesterol movement across mitochondrial membranes (Bose et al. 2008, Elustondo et al. 2017). The transcriptional level of STAR directly affects the synthesis of testosterone.

Internal homeostasis of testosterone levels is of great importance to the body. Previous studies focused on the regulation of testosterone synthesis by the hypothalamic–pituitary system, but little is known about the endogenous regulatory mechanism within Leydig cells. It should be noticed that a lower oxygen concentration in testical tissue was reported to cause significant reduction of serum testosterone level (Gonzales 2013). Since it is known that hypoxia can stabilize HIF1 protein, we speculated that the increased HIF1 level caused a decline of testosterone synthesis, and thus, reduced the testosterone level. Star was identified as a potential target gene of HIF1 through the analysis with software JASPAR. Kowalewski et al. also reported that HIF1 binds to the Star promoter region by performing the chromatin immunoprecipitation (ChiP) assay in granulosa cells (Kowalewski et al. 2015), although the exact binding sites and function of HIF1 on Star have not been determined. Thus, HIF1 may play a key role in regulating testosterone synthesis via the transcriptional regulation of STAR in Leydig cells. The purpose of this study is to determine the binding sites and regulation mode of HIF1 for testosterone synthesis.

Materials and methods

Animals and treatment

Adult male Balb/c mice weighing 25–30g, provided by the Experimental Animal Center of Nantong University, were accommodated for 1 week before experiments with a 12-h light–dark cycle. Mice were treated in an 8.0% O$_2$ hypoxic chamber for 4, 8 or 12 h (n=10 for each time point) and were anesthetized by intraperitoneal injection of chloral.
hydrate. Blood samples were collected from abdominal aorta. Testis tissues were harvested for real-time PCR, Western blot, and immunofluorescence staining. All the studies reported here were submitted to the Ethics Committee on Animal Experimentation of Nantong University and all procedures were approved according to the Animal Care and Use Committee of Nantong University and the Jiangsu Province Animal Care Ethics Committee (Approval ID: SYXK(SU)2007-0021).

Cell culture

Male Sprague–Dawley rats weighing 180–220g, provided by the Experimental Animal Center of Nantong University, were used for isolation of primary Leydig cells as described previously (Klinefelter et al. 1987). Decapsulated rat testes were softly digested by 0.1% collagenase at 37°C. Then, the seminiferous tubules were removed by filtration through the 200-mesh to get cell suspension. The collected cells were resuspended in the medium of 27:3:20 (volume ratio) of Percoll: 9% NaCl saline solution: DMEM-F12 medium and density gradient centrifuged at 20,000 g for 60 min at 4°C. Isolated Leydig cells were cultured at 5 × 10^4 cells/mL in DMEM-F12 medium containing 5% newborn calf serum (Gibco) at 37°C in humidified atmosphere composed of 95% air and 5% CO₂ for subsequent analysis. In addition, 75 IU/L hCG were added for the stimulation of testosterone synthesis and cell growth. The cultured cells were characterized with immunofluorescence staining. Qualified cell samples should consist of more than 98% cells positive of STAR expression.

TM3 cell line (CRL-1714, ATCC, RRID:CVCL_4326) was established from Mus musculus (mouse) and was characterized by the androgen receptor. Cells were cultured in 1:1 mixture of Ham-12 and Dulbecco’s MEM, containing 5% horse serum, 2.5% fetal bovine serum at 37°C in humidified atmosphere composed of 95% air and 5% CO₂. In addition, 200IU/L hCG were added for the stimulation of testosterone synthesis.

Cell treatments and transfection

In the physical treatment group, cells were cultured in the medium which was hypoxia equilibrated for 24 h, and were incubated in 1% O₂ and 5% CO₂ at 37°C. In the chemical treatment groups, 200μM CoCl₂ or DFX were added into medium to induce HIF1 expression. For inhibition of HIF1 expression, TM3 cells were exposed to 200μM digoxin or shRNA at 95% air and 5% CO₂ for 2 h and then at 1% O₂ and 5% CO₂ for 4 h. The supernatant was collected and used for the determination of testosterone levels, and the remaining cells were used for protein extraction, RNA extraction with TRIzol or for immunofluorescence with addition of 4% paraformaldehyde.

HEK293T cells were cultured in DMEM containing 10% fetal bovine serum. HEK293T cells or TM3 cells were seeded in six-well plates in antibiotic-free medium the day before transfection. The constructed plasmid DNAs were transfected into HEK293T cells using Lipofectamine 2000 (11668019, Invitrogen) according to the provided instructions or transfected into TM3 cells by electroporation.

Total RNA isolation and real-time PCR

Testicular tissue total RNA was isolated by using the column animal RNAout kit (TianDZ, Beijing, China) according to the manufacturer’s instructions. Cellular total RNA was isolated by TRIzol reagent. 500ng purified total RNA was then reverse-transcribed using HiScript 1st Strand cDNA Synthesis Kit (Vazyme, China) according to the instructions. Real-time PCR was performed using SYBR premix (Roche) with the 7500 real-time PCR system (Applied Biosystems). The reaction procedure was 98°C for 4 min followed by 40 three-step cycles of 98°C for 25 s, 60°C for 25 s and 72°C for 25 s. All the primers used for real-time PCR are shown in Supplementary Table 1 (see section on supplementary data given at the end of this article). The relative amount of gene expression normalized to the reference gene was calculated using ΔΔCt method, where Ct is the threshold cycle of PCR reaction. The evaluation of contamination of both RNA sample and water sample were provided through agarose gel electrophoresis (Supplementary Fig. 1). PCR products were sequenced and verified (Supplementary Fig. 1). The amplification efficiency was within the scope from 0.9 to 1.1. Although we did not choose to fully present our housekeeping gene (at internal controls), we examined multiple control genes including 18s rRNA, Actb and Gapdh, which gave consistent results. In our experiments, we made sure that Ct for internal controls are between 14 and 17, and Ct for target genes are between 20 and 30 according to reagent manufacturer’s suggestions for reliable quantification. None of the samples gave results outside those ranges.

Western blot

Testicular tissues and cells were collected in the protein extraction reagents, placed on ice for 30 min, followed by 15,000 g centrifuging at 4°C for 15 min. Protein
concentration was determined by bicinchoninic acid assay. Proteins were isolated by sodium dodecyl sulfate polyacrylamide gel electrophoresis and transferred to polyvinylidene fluoride membranes using a semi-dry or wet trans-blot unit. The membranes were incubated for 1 h with 5% nonfat dry milk and then probed overnight at 4°C with anti-HIF1α (GTX127309, GeneTex, AB_2616089), anti-VEGFA (#AB882, Millipore, AB_10806337), anti-STAR (8449, CST, AB_10889737), anti-P450scc (14217, CST, AB_2631970), anti-3beta-HSD (sc-30820, Santa Cruz, AB_2279878) and anti-B-actin (ab8227, Abcam, AB_2305186). Binding of primary antibodies was visualized with donkey anti-rabbit HRP-conjugated secondary antibody (305-005-003, Jackson, AB_2339376) or rabbit anti-goat HRP-conjugated secondary antibody (111-005-003, Jackson, AB_2337913) and the ECL-Plus system. Grayscale analysis was carried out using ImageJ (https://imagej.nih.gov/ij/).

**Immunofluorescence**

Cultured cells were fixed with 4% paraformaldehyde for 15 min at room temperature, permeabilized with Triton (0.1% in TBS) for 30 min and blocked with 5% BSA in PBS for 1 h at room temperature. Cells were then incubated overnight at 4°C with anti-HIF1α (AF1935, R&D, AB_355064) and anti-STAR (8449, CST, AB_10889737). Binding of primary antibodies was visualized with Alexa Fluor 488-conjugated anti-rabbit IgG (R37118, Thermo Fisher Scientific, AB_2556546) or Alexa Fluor 555-conjugated anti-goat IgG (A-21432, Thermo Fisher Scientific, AB_2535853) for 90 min. Finally, the samples were incubated in DAPI for 10 min (Life Technologies) for nuclear counterstain. Paraffin-embedded testicular tissue was sectioned at 5 μm following dewaxing hydration treatment for immunofluorescence staining. Images were acquired using a Leica SP8 confocal microscope with 63x oil immersion objective.

**Chromatin immunoprecipitation (ChIP) assay**

ChIP was performed using the SimpleChIP Enzymatic Chromatin IP Kit (Magnetic Beads) (CST, #9003) according to the manufacturer’s instructions. To crosslink proteins to DNA, TM3 cells were fixed in culture medium containing 1% formaldehyde at room temperature for 10 min. Cells were lysed and collected, followed by adding 0.5 μL of micrococcal nuclease per IP preparation and incubating for 20 min at 37°C to digest DNA to lengths of approximately 200–500bp. The sonicated lysate was then immunoprecipitated with 2.5 μg of HIF1α antibody (AF1935, R&D, AB_355064) or a negative control IgG at 4°C for overnight. The pulled-down chromatin was washed, reverse-cross linked and purified. The purified DNA was amplified by PCR with the corresponding primers listed in Supplementary Table 2. The reaction procedure was 98°C for 5 min followed by 45 three-step cycles of 98°C for 25 s, 60°C for 30 s and 72°C for 25 s. Products were analyzed by 1.5% agarose gel electrophoresis and quantified by real-time PCR. Fold enrichment = 2^{Ct (IgG)}−Ct (HIF1α).

**Electrophoretic mobility shift assay (EMSA) and supershift assays**

EMSA was carried out to determine HIF1 interactions with putative binding elements on mouse Star promoter as previously described (Luo et al. 2003). Briefly, oligonucleotide probes with putative HIF1-binding sites on mouse Star promoter and mutations were designed, synthesized, annealed and labeled by biotin. Each probe was incubated with 5 μg TM3 cell nuclear extracts (Promega) and processed for EMSA. Supershift assays were performed as follows. In each reaction, 2.5 μg of HIF1α antibodies (AF1935, R&D, AB_355064) were added to the probe/nuclear extract mixture and incubated for 20 min at room temperature. For competition, 100-fold excess of unlabeled cold oligo probes were incubated with nuclear extract before adding labeled oligonucleotide. Shift reactions were loaded onto 6% polyacrylamide gel and run at 170V (at 4°C) for 5 h in 0.5× TBE buffer. Results were obtained in the chemical system.

**Dual-luciferase reporter assays**

Fragments of mouse Star promoter region (−2227/−1893, −1000/+78) were cloned into the KpnI and HindIII restriction sites of the pGL3-basic vector (Promega). The HIF1-binding sites on mouse Star promoter were mutated respectively. All constructs were verified by sequencing. HEK293T cells (4 × 10^6) were cultured in 24-well plate in antibiotic-free culture medium the day before transfection. Cells were cotransfected with 500 ng STAR reporter plasmids and 5 ng of Renilla reporter plasmid (pRL-TK, Promega) as an internal control by Lipofectamine 2000 reagent (1.5 μL) and then treated with 4 h hypoxia, CoCl$_2$ or DFX. Cells were lysed and the luciferase activities were detected by Dual-Luciferase Reporter Assay System.
(Promega). Firefly luminescence signal was normalized by Renilla luminescence signal.

**Testosterone determination**

Mouse plasma and cell supernatant were collected and diluted to 20% according to the requirements of the instructions of the kit. The testosterone level was determined following the manufacturer’s instructions of the testosterone parameter assay kit (R&D, KGE010). First, primary antibody (50 µL per well) was incubated for 1 h at room temperature on a microplate shaker at 800 g. The standard, control or sample (100 µL each) was added after three times washing. Then, testosterone conjugate (50 µL) was added into each well, incubated for 3 h at room temperature. Finally, stop solution (200 µL) was added after 30-min incubation of substrate solution.

The optical density was determined at 450 nm and 540 nm wavelengths. Standard curve was created by performing a four-parameter logistics curve-fit with Origin 8.5 (http://www.originlab.com/). The sensitivity of testosterone assay is 0.03 ng/mL. The intra-assay precision of these results is between 2.2 and 3.7%, and the inter-assay precision of is between 4.2 and 5.5%.

**Statistical analysis**

All the data were presented as mean±s.d. Means of two samples were compared using unpaired Student’s t-test, and means of multiple groups were compared using ANOVA followed by Bonferroni’s test. Differences were considered statistically significant if \( P < 0.05 \). The statistical software used for analyzing the data was SPSS 17.0 (https://www.ibm.com/analytics/us/en/technology/spss/).

**Figure 1**

Testosterone level decline is accompanied by the accumulation of HIF1α protein. (A) After 4-, 8- or 12-h hypoxia (8% \( O_2 \)) treatment, mouse testis HIF1α protein levels were determined by Western blot, and serum testosterone concentrations were examined by ELISA (\( n = 10 \), ***\( P < 0.001 \) vs 0 h). (B) Histopathological changes were determined by H&E staining of testis from mice after hypoxia (8% \( O_2 \)) treatment for 0, 4, 8 or 12 h (\( n = 10 \)). (C) Rat primary Leydig cells were exposed to 1% \( O_2 \) for 0, 1, 2, 4, 8 and 16 h. HIF1α protein levels were determined by Western blot, and supernatant testosterone concentrations were examined by ELISA (\( n = 6 \), *\( P < 0.05 \), **\( P < 0.01 \), and ***\( P < 0.001 \) vs 0 h). DFX, deferoxamine. A full colour version of this figure is available at https://doi.org/10.1530/JME-18-0148.
Results

Upregulation of HIF1 by hypoxia, DFX and CoCl₂ decreases testosterone synthesis in murine Leydig cells

To evaluate the hypoxia effect on testosterone synthesis, male mice were treated with 8.0% O₂ for 0, 4, 8 and 12 h. Western blot results showed that the protein level of testis HIF1 significantly increased, while ELISA results indicated a decline trend of serum testosterone concentration (Fig. 1A). The histological structure of the testis was not significantly altered as illustrated by H&E staining (Fig. 1B). Rat primary Leydig cells and TM3 cells were treated with 1% O₂ and both showed significant increases of HIF1 protein level and significant decline of supernatant testosterone concentration (Fig. 1C and D). To confirm that the reduction of testosterone was caused by HIF1, we added CoCl₂ or DFX to stabilize HIF1 in TM3 cells and detected testosterone level in supernatant. Similar to the response to hypoxia, testosterone level significantly decreased when HIF1 increased (Fig. 1E and F), indicating that HIF1 could reduce testosterone synthesis.

HIF1-induced testosterone reduction is recovered by pregnenolone but not cAMP

In testosterone synthesis pathway, cAMP is induced by LH receptor and activates PKA which then stimulates STAR phosphorylation and cholesterol transport. We added cAMP to the cell culture medium in synchronism with hypoxia (1% O₂), 200nM DFX, or 200nM CoCl₂ for 4 h. Results showed that cAMP could not recover testosterone synthesis in these cells as in normoxic cells, indicating that the interference of HIF1 on testosterone synthesis was not prior to the transport of cholesterol (Fig. 2A). We also added pregnenolone, an intermediate product produced by p450scc in mitochondria, into cell culture medium with hypoxia, DFX or CoCl₂ treatments for 4h, where testosterone synthesis was recovered in a dose-dependent trend, indicating that HIF1-mediated interference on testosterone synthesis was upstream of pregnenolone synthesis (Fig. 2B).

HIF1 accumulation by hypoxia, DFX or CoCl₂ downregulates STAR

Next, we determined the protein and mRNA levels of factors related to testosterone synthesis after hypoxia, DFX or CoCl₂ treatments in vivo and in vitro. Mice were exposed to 8% O₂ for 0, 4, 8 and 12h. As shown in Fig. 3A and B, VEGF-A and HIF1α in mouse testes significantly increased after hypoxia treatment (8% O₂). Protein and mRNA levels of STAR significantly decreased, while other factors including P450scc and 3b-HSD remained unchanged. Similarly, in both rat primary Leydig cells
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and mouse Leydig cell line TM3, 1% O2, 200nM CoCl2, or 200nM DFX for 4h upregulated HIF1α and VEGF-A whereas downregulated STAR, but did not affect P450scc or 3b-HSD (Fig. 3C, D, E and F).

Furthermore, immunofluorescence indicated that hypoxia, CoCl2 and DFX induced accumulation and nuclear transposition of HIF1α, with STAR levels depressed simultaneously in vivo (Fig. 4A) and in vitro (Fig. 4B).

Repression of HIF1α recovers STAR and testosterone synthesis in both normoxia and hypoxia condition

To determine whether only HIF1 or other factors were involved in the decrease of testosterone, we repressed HIF1α by PX-478 and digoxin under normoxia for 0, 1, 2, 4, 8 and 16h. Western blot and real-time PCR results showed a significant decrease of HIF-1 and increase of STAR, which were highest at 2 and 4h (Fig. 5A and C). Testosterone level continued to rise over time (Fig. 5B). Similar results were shown when TM3 cells were exposed to HIF-1α shRNA or 200nM digoxin under hypoxia.

While HIF1α protein level significantly decreased in these cells, VEGFA mRNA level (Fig. 5E) significantly decreased. Both mRNA and protein levels of STAR (Fig. 5D) significantly increased synchronously with testosterone concentration (Fig. 5F). Immunofluorescence results confirmed that the repression of HIF1α by digoxin was accompanied by an increase in STAR level (Fig. 5G). These results indicated that silencing HIF1α upregulated STAR expression under normoxia and hypoxia, demonstrating HIF1α as the main factor to suppress STAR.
HIF1 transcriptional inhibition of STAR

The abovementioned results indicated that HIF1α decreased testosterone synthesis by inhibiting STAR expression in TM3 cells under hypoxia. The exact mechanism of HIF1 participation in STAR regulation was unknown. Bioinformatics analysis of JASPAR 8.0 (http://jaspar.genereg.net/) showed that there were four predicted hypoxia response elements (HREs) in the Star promoter region, i.e. −2082/−2078, −2064/−2060, −1910/−1906, −215/−218. Probes of these target sequences (probe 1: −2090/−2055, probe 2: −1920/−1899 and probe 3: −222/−201) were synthesized as shown in Fig. 6A. EMSA experiments were performed to evaluate the combination of HIF1α and probes. As shown in Fig. 6B, probes 1, 2 and 3 combined to the nuclear exact in line 2, 7 and 12 and were competed by cold probes in line 3, 8 and 13, indicating that all these sequences combined with HIF1α protein. However, only probes 1 and 2 combined to HIF1α in the supershift assays in line 4, 9 and 14 with addition of HIF1α antibody (Fig. 6B). ChIP assay was used to confirm the binding of HIF1α to the Star promoter in vivo. Agarose gel electrophoresis results indicated that HIF1α co-precipitated with the DNA fragment −2337/−1893 under both normoxia and hypoxia, and with the fragment −550/−170 under hypoxia (Fig. 6C). Real-time PCR results showed that the fold enrichment of HIF1α to IgG in −2337/−1893 was above 40 times under hypoxia and 10 times under normoxia, while that in −550/−170 was above six times under hypoxia and about one-fold under normoxia (Fig. 6D). To determine whether such binding of HIF1α to Star promoter regions was functional, Star promoter fragments (−2337/−1893, −1000/+78) were cloned into luciferase reporter vector, which was transfected into HEK293T cells, followed by hypoxia (1% O₂), 200nM CoCl₂ or 200nM DFX treatments. Luciferase assay results showed that upregulation of HIF1α significantly decreased relative luciferase activity in HEK293T cells with luciferase under the control of Star promoter −2337/−1893, while the other group of Star promoter −1000/+78 did not show any significant difference (Fig. 6E).

**Mutations of the binding sites relieve HIF1α inhibition of STAR**

To verify the specific binding between HIF1α and the Star promoter region, probes with mutations in the binding sites were designed and synthesized as shown in Fig. 7A. WT probes and their mutations were incubated with nuclear exacts from hypoxia-treated TM3 cells (1% O₂, 4h). EMSA results showed higher combinations of HIF1α and WT probes in hypoxia-treated TM3 cells. The binding activity of the mutated probes (mutated site 1: −2082/−2078, mutated site 2: −2064/−2060, mutated site 3: −1910/−1906) was significantly reduced compared to the WT (Fig. 7B). To determine the transcriptional activity of Star with mutations on HIF1-binding sites, luciferase reporter vectors were transfected into HEK293T cells, followed by 1% O₂ or 200nM CoCl₂ treatments. As shown in Fig. 7D and E, transcriptional activities of STAR with mutations on HIF1-binding sites were significantly higher than the WT. In particular, transcriptional activity of Star with mutated site 3 (−1910/−1906) was stronger than with two other sites (Fig. 7D and E). These results indicated that HIF1α bound to the Star promoter region −2337/−1893 and inhibited STAR transcription.

**Discussion**

In this work, we investigated the molecular mechanism for the transcriptional regulation of STAR through HIF1α and the consequent effects on the synthesis and secretion of testosterone in Leydig cells. We used physiological or chemical hypoxia (CoCl₂ or DFX) to upregulate the protein level of HIF-1. To verify the upregulation of HIF-1 regulatory pathway, we detected the expression of VEGFA, which is one of the major target genes of HIF-1 and is the most commonly used positive reference for HIF-1 (Semenza 2001). We found that upregulation of HIF1α caused a significant decrease in STAR and testosterone both in vivo.
and in vitro. The STAR level and testosterone concentration were negatively correlated with HIF1, which is consistent with the results from Fadhillah et al. who have reported that hypoxia (1 and 5% O₂) could inhibit steroid synthesis though STAR in ovarian granulosa cells (Fadhillah et al. 2017). It was also reported that intermittent hypoxia (10% O₂ for 17 days) could cause abnormal synthesis of STAR protein and inhibit testosterone secretion in rats (Zhang et al. 2013). In vivo studies with zebrafish also showed that upregulation of HIF1 expression could inhibit the synthesis of STAR protein (Tan et al. 2017). Although these studies have proved that HIF1 represses STAR expression, the underlying molecular mechanism was not clarified. Our work reveals the exact binding sites for HIF1 on the Star promoter region (−2082/−2078, −2064/−2060 and 1910/−1906) and confirms the negatively regulatory effect of HIF1 on STAR and testosterone synthesis. One interesting finding was that a HRE in the Star promoter region −222/−201 bound to HIF1, while HIF1 was not able to bind to this site and any change of HIF1 did not affect the transcription activity of this site (Fig. 6). We speculate that there might be an actual HRE in the region −222/−201 where HIF1 binding depends on the help of other DNA elements and associated protein complexes.

Figure 5
Repression of HIF1 upregulates STAR and testosterone. (A) Protein levels of HIF1 α, STAR and B-actin in TM3 cells were determined by Western blot. (B) mRNA levels of Star were detected by real-time PCR and normalized to Actb (n = 6, *P < 0.05, ***P < 0.001). (C) In situ STAR and HIF1 α in TM3 cells were examined by immunofluorescence (n = 4). STAR was stained with Alexa 488 (green), HIF1 α with Alexa 555 (red) and nucleus with DAPI (blue). (D) Protein levels of HIF1 α, STAR, and B-actin in TM3 cells were determined by Western blot (n = 4, *P < 0.05, ***P < 0.001). (E) mRNA levels of Star and Vegfa were detected by real-time PCR and normalized to Actb (n = 6, *P < 0.05, ***P < 0.001). (F) Testosterone levels in the supernatant of TM3 cells were determined by ELISA (n = 6, ***P < 0.001). (G) In situ STAR and HIF1 α in TM3 cells were examined by immunofluorescence (n = 4). STAR was stained with Alexa 488 (green), HIF1 α with Alexa 555 (red) and nucleus with DAPI (blue). A full colour version of this figure is available at https://doi.org/10.1530/JME-18-0148.
which lie away from the region covered by the EMSA probes and the cloned promoter fragments.

It is known that HIF1 is stably expressed in Leydig cells under normal physiological conditions. We think that the transcriptional inhibitory effect of HIF-1 on STAR does not completely prevent the testosterone synthesis. The results of WB and ELISA also prove this. Stable HIF-1 expression may play a negative regulatory role in normoxic condition. Maximum decrease in testosterone concentration occurs after 4 h of hypoxia, CoCl$_2$, DFX in TM3 cells and after 8 h of hypoxia in rat primary Leydig cells. However, the levels increase thereafter. It is interesting that the expression level of HIF-1 is also highest at 4 h and then gradually decreases. We speculate that the increase in testosterone after 4 h is due to a decrease in the level of HIF-1. Hwang et al. (2009) found intermittent hypoxia (10% O$_2$, 8h per day) to male mice for 1 day, the testosterone decreased. However, testosterone was recovered and increased further when mice were treated for 3–14 days. But after 28-day treatments, the serum testosterone levels were significantly decreased and continued to decline, consistent with the decrease of serum testosterone in patients with clinical respiratory disorder during sleep. We inferred that the increase in testosterone in the medium term may be a stress response.

From the synthesis of cholesterol to testosterone, there are three rate-limiting proteins, of which STAR is the first rate-limiting protein, while P450scc and 3b-HSD are the other two (Aspden et al. 1998). Previous studies reported that 3b-HSD and P450scc are target genes for HIF1α in murine Leydig cells (Lysiak et al. 2009, Kumar et al. 2014). But in our work, neither protein nor mRNA levels of P450sc and 3b-HSD were significantly changed.

Figure 6
HIF1α directly binds to the Star promoter region. (A) Probe sequences used. Numbers indicate the positions upstream of transcription start site. Capital letters show the putative HIF1-binding sites. (B) EMSA supershift experiments. Probes were labeled with biotin. Each probe was incubated with nuclear extract of TM3 cells. Shift arrows indicate the complex positions (lane 2, 7 and 12). Unlabeled probes were used as competitions and co-incubated with labeled probes and nuclear extract (lane 3, 8 and 13). HIF1 antibody was added to distinguish the combinations of HIF1 and probes. Supershift arrows indicate the complex positions (lane 4, 9 and 14). IgG was added into the mixture as negative control for HIF1 antibody (lane 5, 10 and 15). (C and D) ChIP experiments. IgG groups were operated as negative controls for HIF1 groups. PCR and real-time PCR were performed to amplify the ChIP products. Numbers in C indicate positions upstream of transcription start site. Vegfa was examined as positive control for ChIP. Enrichment effects of ChIP were performed in D by comparison to IgG (n = 3). (E) Luciferase reporter gene assay following transient transfection with Star (−1000/+38Luc or −2337/−1893Luc). The pRL-TK vector was co-transfected to normalize transfection efficiencies. Results were presented as a luciferase/Renilla ratio (n = 6, ***p < 0.001 vs Nor). DFX, deferoxamine; Hyp, hypoxia; Nor, normoxia.
under hypoxia, while STAR mRNA and protein levels were significantly reduced. Addition of pregnenolone recovered the testosterone synthesis decline induced by HIF1, suggesting that the testosterone synthesis pathway in cytoplasm was not affected. One possibility is that the shorter duration of hypoxia (4h) in our study was not long enough to show the regulation on 3b-HSD and P450scC levels. cAMP is an upstream signaling factor in testosterone synthesis pathway (Kolena et al. 1978). Addition of cAMP into TM3 cells did not recover testosterone synthesis, indicating the factor affected by HIF1 should be downstream of cAMP. Thus, the changes in testosterone synthesis pathway might involve cholesterol transport and pregnenolone synthesis, where STAR and p450scC are the two key factors, respectively.

External environment changes may lead to a low level of testosterone. Previous clinical work has demonstrated that respiratory hypoxia, e.g. chronic obstructive pulmonary disease (COPD), and sleep apnea are related to serum testosterone depression (Semple et al. 1980). Decreased serum testosterone level is also associated with high altitude living (Gonzales 2013), which implies that hypoxia may induce serum testosterone depression. In this study, we used hypoxia as a powerful means to increase the HIF1 level, which significantly decreased the testosterone level, consistent with clinical results (Fig. 1). It suggests the transcriptional repression of STAR by HIF1 as one mechanism for the reduction of testosterone under hypoxia condition. Hypoxia condition in tissues and organs mediated by low oxygen supply is a critical factor that affects many physiological and pathological processes, particularly in the endocrine system (Trayhurn et al. 2008, Anju et al. 2011, Olufsen & Arukwe 2015).

For instance, STAR is necessary for syntheses of steroid hormones (Sewer & Waterman 2003). Considering that STAR not only existed in Leydig cells but also is expressed in all steroid cells including granulosa cells and adrenal cells as an essential factor in steroidogenesis from cholesterol, we infer that abnormal HIF1 levels in hypoxic tissues, tumors and degradation pathway disorders may affect steroidogenesis and thus damage endocrine system and related organs. However, it was reported that long-term exposure to the high altitude of low oxygen environment, or low blood oxygen partial pressure induced by respiratory diseases, could cause lower levels of the blood testosterone and estrogen and significantly higher corticoid and aldosterone levels (Newby et al. 2015, Verratti et al. 2017). It was inferred that HIF1-mediated STAR repression was not the only mechanism for the hypoxia-induced steroidogenesis disorder. Our other work on STAR transcriptional regulation indicated that nuclear respiratory factor 1 (NRF1) stimulated STAR transcription, which was downregulated under chronic hypoxia (Wang et al. 2017). Since NRF1 increase in adrenal gland was observed (unpublished), we infer that stimulation of STAR by NRF1 might be another mechanism underlying hypoxia-induced steroidogenesis disorder.

Steroid hormones are extremely important in circulatory system, cardiovascular system, respiratory system, reproductive system, etc. As the key male hormone, testosterone is crucial for male secondary sexual characteristics of the development, maintenance of sexual desire and skeletal muscle growth. The incidence of male COPD is much higher than that of women, and the
severity of disease is positively correlated with the decrease of testosterone level (Semple et al. 1980). Our study of the reduction of testosterone secretion by HIF1 provides a new insight into the entire testosterone synthesis disorder field. Although the testosterone replacement therapy has made great development, the clinical effects still need to be improved. We will further study the effects of testosterone on the pathogenesis of COPD, in order to provide new ideas for the clinical treatment.

Supplementary data
This is linked to the online version of the paper at https://doi.org/10.1530/ JME-18-0148.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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