

Genetic characterization of a missense mutation in the X-linked *TAF7L* gene identified in an oligozoospermic man[†]

Li Ling^{1,‡}, Fangfang Li^{1,‡}, Pinglan Yang^{1,‡}, Robert D. Oates², Sherman Silber³, Cornelia Kurischko⁴, Francis C. Luca⁴, N. Adrian Leu⁴, Jinwen Zhang¹, Qiuling Yue¹, Helen Skaletsky⁵, Laura G. Brown⁵, Steve G. Rozen⁶, David C. Page⁵, P. Jeremy Wang^{4,*} and Ke Zheng^{1,*}

¹State Key Laboratory of Reproductive Medicine, Nanjing Medical University, Nanjing, China

²Department of Urology, Boston University Medical Center, Boston, MA, USA

³Infertility Center of St. Louis, St. Luke's Hospital, St. Louis, MO, USA

⁴Department of Biomedical Sciences, University of Pennsylvania School of Veterinary Medicine, Philadelphia, PA, USA

⁵Howard Hughes Medical Institute, Whitehead Institute, and Department of Biology, Massachusetts Institute of Technology, Cambridge, MA, USA

⁶Centre for Computational Biology, Duke-NUS Graduate Medical School, Singapore

*Correspondence: State Key Laboratory of Reproductive Medicine, Nanjing Medical University, Nanjing 211166, China. E-mail: kezheng@njmu.edu.cn or Department of Biomedical Sciences, University of Pennsylvania School of Veterinary Medicine, Philadelphia, PA 19104, USA. E-mail: pwang@vet.upenn.edu

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[‡]Li Ling, Fangfang Li and Pinglan Yang contributed equally to this work.

Abstract

Although hundreds of knockout mice show infertility as a major phenotype, the causative genetic mutations of male infertility in humans remain rather limited. Here, we report the identification of a missense mutation (D136G) in the X-linked *TAF7L* gene as a potential cause of oligozoospermia in men. The human aspartate (D136) is evolutionally conserved across species, and its change to glycine (G) is predicted to be detrimental. Genetic complementation experiments in budding yeast demonstrate that the conserved aspartate or its analogous asparagine (N) residue in yeast TAF7 is essential for cell viability and thus its mutation to G is lethal. Although the corresponding D144G substitution in the mouse *Taf7l* gene does not affect male fertility, RNA-seq analyses reveal alterations in transcriptomic profiles in the *Taf7l* (D144G) mutant testes. These results support *TAF7L* mutation as a risk factor for oligozoospermia in humans.

Summary Sentence

A sequencing screen of infertile men identifies a missense mutation in the human *TAF7L* gene.

Keywords: infertility, oligozoospermia, spermatogenesis, X chromosome, *TAF7L*, transcription factor, Characterization of missense mutation in *TAF7L* gene

Introduction

An estimated 15% of couples suffer from infertility worldwide, half of which is attributed to male factors [1–4]. Spermatogenic failure represents a common cause of male infertility [5]. About 50% of male infertility cases are idiopathic, and idiopathic male infertility is believed to be largely genetic in etiology [6, 7]. Studies of mouse models have defined many pathways that regulate reproduction in mammals [7, 8]. At least 2000 genes are preferentially expressed in mouse spermatogenesis [2], and more than 400 of them are known to regulate mouse fertility [3, 4]. Exome-sequencing of infertile patients has identified mutations in a number of genes in humans as the cause of male sterility [9, 10]. Despite these recent advances, our knowledge about the genetic etiology underlying male infertility in humans remains rather limited.

The X chromosome is enriched for spermatogonally expressed testis-specific genes in mouse [11]. *Taf7l* (TBP associated factor 7 like) is an X-linked testis-specific homologue of TBP associated factor 7 (*Taf7*) [11]. *Taf7* is an autosomal

retrotransposon and encodes a ubiquitously expressed component of the transcription factor transcription factor IID (TFIID) that is required for the transcription of protein-coding genes by RNA polymerase II [12]. Like TAF7 in somatic tissues, *TAF7L* interacts with TBP associated factor 1 (TAF1) and is associated with TATA box-binding protein (TBP) in the testes, indicating that *TAF7L* is a bona fide TBP-associated factor (TAF) [13]. Moreover, biochemical studies have shown that *TAF7L* replaces TAF7 in the TFIID complex to modulate the transcription in spermatogenesis [13]. Inactivation of *Taf7l* in mice causes a reduction in the sperm count, sperm motility, testis weight, and male fertility [14]. A backcross of *Taf7l* knockout mice to the C57BL/6J background produced knockout males that are essentially sterile [15]. The deletion of testis-expressed gene 11 (*Tex11*), an X-linked germ cell-specific gene, leads to meiotic arrest in males [16]. Interestingly, the *Taf7l Tex11* double mutant males exhibit much severe defects in meiosis than either single mouse mutant, suggesting the synergistic regulation of

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spermatogenesis by these two X-linked genes [17]. In addition to its role in spermatogenesis, TAF7L acts as a transcription regulator in adipocyte differentiation [15, 18].

Based on the phenotype of *Taf7l* knockout mice, we reasoned that mutations in human TAF7L gene would most likely cause reduced sperm count—oligozoospermia. Sequence variants in TAF7L were previously reported in oligozoospermic men [19]. However, these TAF7L variants were either found in fertile controls or not verified [20], leaving their functional consequence unknown [7]. To determine if TAF7L plays a role in human fertility, we screened 113 infertile men and identified a single amino acid substitution (D136G) in TAF7L in an infertile man with low sperm count. This aspartic residue is conserved in TAF7 from yeast to human and in TAF7L in mammals. We genetically characterized the functional impact of this amino acid substitution in TAF7 and TAF7L in yeast and mouse, respectively.

Materials and methods

Ethics statement

The human study was approved by the Institutional Review Board of Massachusetts Institute of Technology. Informed consent was obtained from all participants. Mice were maintained under veterinarian care from the University Laboratory Animal Resources at the University of Pennsylvania or Nanjing Medical University. All animal experimental procedures were approved by the Institutional Animal Care and Use Committees at University of Pennsylvania and Nanjing Medical University.

Patients and controls

In this study, 113 patients with severe oligozoospermia (sperm count $<5 \times 10^6/\text{ml}$) were included. We excluded patients with any of the following conditions or treatments that cause or predispose to spermatogenic failure: Y-chromosomal deletions [21–25]; a 47, XXY karyotype [26]; cryptorchidism; orchitis; radiotherapy; or chemotherapy. We also studied 175 controls, consisting of men known to be fertile ($n=93$) and men of unknown fertility selected to represent worldwide genetic diversity ($n=82$, samples from the National Institutes of Health polymorphism discovery panel, Coriell Cell Repositories, and from our own collection) [27]. This collection of 175 control men was previously used in the *TEX11* genetic study [28]. Genomic DNA was prepared from peripheral blood leukocytes or EBV-transformed lymphoblastoid cell lines.

Mutation screening

The TAF7L exon/intron structure was determined by the alignment of human TAF7L cDNA sequence (NM_024885) with the genomic sequence [11]. We polymerase chain reaction (PCR)-amplified the TAF7L coding exons (exons 2 through 13) with 12 primer pairs (Supplementary Table S1). PCR was performed in a 25- μl reactor with 12.5 ng of genomic DNA (94 °C, 30 s; 56 °C, 30 s; 72 °C, 90 s; 35 cycles). PCR products were purified by Sephadex S-300 gel filtration. Then, 12.5 μl of the purified PCR product was sequenced in a 25- μl reactor using one of the PCR primers and ABI BigDye on an ABI 3700 sequencer according to the manufacturer's instructions. Sequence data were analyzed using the Sequencher software (Gene Codes Corporation, Ann Arbor, USA), and sequence variants were identified by an inspection of the aligned sequences in Sequencher.

Pathogenicity prediction of the human TAF7L D136G substitution

Four algorithms, Mutation Taster [29], PolyPhen2 [30], PROVEAN [31], and SIFT [32], were used to predict the possible effects of the missense mutation D136G found in the human TAF7L gene. The web links are as follows: Mutation Taster (<http://www.mutation-taster.org>); PolyPhen2 (<http://genetics.bwh.harvard.edu/pph2>); PROVEAN (<http://provean.jcvi.org/index.php>); and SIFT (<https://sift.jcvi.org/>).

Plasmid constructs and yeast strains

The wild type (WT) *Saccharomyces cerevisiae* TAF7 gene (scTAF7-N147) was PCR-amplified from the yeast genomic DNA and cloned into the CEN plasmid pRS416 (URA3+). Two TAF7 mutants harboring N147D and N147G point mutations respectively were generated by PCR-based mutagenesis [33] and subcloned into pRS415 (LEU2+) to yield pRS415-scTAF7-N147D and pRS415-scTAF7-N147G plasmids. pRS416-scTAF7-N147 (WT) was transformed into TAF7/*taf7* Δ ::*kanMX6* heterozygous diploid yeast cells [34, 35], and tetrad analysis was performed to test whether the scTAF7 (WT) plasmid rescues the lethality of the yeast *taf7* deletion. Sc *taf7* Δ segregants containing pRS416-scTAF7-N147 were viable and plasmid dependency for viability was confirmed by assaying for growth on 5-FOA plates, which counter-selects the URA3-containing pRS416-scTAF7 plasmid [36]. To determine whether scTAF7-N147D or cTAF7-N147G rescued the *taf7* Δ mutation, haploid *taf7* Δ cells containing pRS416-scTAF7-N147 (WT) were transformed with pRS415-scTAF7-N147D or pRS415-scTAF7-N147G and the viability of the transformants was assessed after counter-selecting the pRS416-scTAF7-N147 plasmid on 5-FOA plates.

Generation of *Taf7l* knockin (D144G) mutant mice

We mutated codon 144 from GAT [aspartate (D)] to GGT [glycine (G)] in the mouse *Taf7l* gene to generate a mutation analogous to the D136G mutation in the human TAF7L gene. Codon 144 is located in exon 4 (Figure 2A). We generated this point mutation in the targeting vector by an overlapping PCR-mediated mutagenesis strategy as described previously [33]. Using a *Taf7l*-containing BAC clone (RP22-415C9) as the template, two homologous arms (2.4 and 2.1 kb) were amplified by high-fidelity PCR (Figure 2A). The floxed neomycin selection marker was cloned into intron 4. V6.5 ES cells were electroporated with the linearized targeting vector and were selected for integration in the presence of G418 (350 $\mu\text{g}/\text{ml}$) and gancyclovir (2 μM). In G418-positive clones screened for homologous recombination, 7 out of 184 were homologously targeted. Exon 4 was PCR-amplified from positive ES cell clones and sequenced to confirm the presence of the point mutation. Two ES cells harboring the *Taf7l* D144G allele were injected into BALB/c blastocysts that were subsequently transferred to the uteri of pseudopregnant females. The resulting male chimeras were bred with C57BL/6J females to obtain germline transmission of the mutant allele. Offspring were genotyped by PCR assays. WT (599 bp) and *Taf7l* mutant (471 bp) alleles were assayed by PCR with the *Taf7l*-KI forward and reverse primers (Supplementary Table S2). *Taf7l* knockin mice were crossed with *Actb*-Cre to remove the floxed PGK-Neo selection cassette [37]. The *Actb*-Cre transgene was removed during subsequent breeding. *Taf7l* knockin mice were backcrossed to the C57BL/6J strain (Jack-

son Laboratory) for 10 generations. The backcrossed mice were used in this study.

Quantitative real-time PCR (qRT-PCR) expression analysis

Total RNA was isolated from WT and *Taf7l* mutant adult (3 months old) mouse testes using TRIzol reagent (15 596 026, Invitrogen, USA). One microgram of total RNA was reverse-transcribed using the PrimeScript RT Master Mix (RR036A, Takara, Japan). The obtained cDNA was diluted five times and then RT-qPCR was performed using the TB Green Premix Ex Taq II (RR820A, Takara, Japan) on a StepOnePlus RT-PCR machine (4 376 592, Applied Biosystems, USA). The relative expression of target mRNA was determined after normalization to the *36B4* gene. Student *t*-test was used for statistical analysis.

Western blot analysis

Protein extracts were prepared from fresh testes of adult mice. Testicular protein (5 μ g) from each sample was loaded and separated on a 10% SDS-PAGE gel. After transferring onto a polyvinylidene fluoride membrane (1 620 177, Bio-Rad, USA), the blots were incubated with the following primary antibodies: anti-TAF7L (DF2403, Affinity, China; 1:1000) or anti-GAPDH (AC033, Abclonal, Wuhan, China; 1:20 000), and secondary antibodies: HRP goat antimouse IgG (H+L) (AS003, Abclonal, Wuhan, China; 1:10 000) or HRP goat antirabbit IgG (H+L) (AS014, Abclonal, Wuhan, China; 1:10 000).

Histology

The testes and epididymides from 3-month-old mice were fixed in Bouin's solution (HT10132-1 L, Sigma, USA) overnight. The tissues were dehydrated through gradient ethanol, embedded in paraffin, sectioned at 5 μ m, and stained with hematoxylin and eosin.

Mating test

Each WT or *Taf7l*^{D144G/Y} male (five males per genotype) was housed with two *Taf7l*^{+/-D144G} females for 10 months. Mice were 2 months old at the start of the mating test. The date and size of litters were recorded.

Sperm count

Cauda epididymides were dissected. Each epididymis was minced in 4 ml 1 \times phosphate-buffered saline (PBS) solution to disperse the sperm and fixed with 2 ml of 4% paraformaldehyde (PFA) for 10 min. Sperms were counted using a hemocytometer.

Computer-assisted sperm analysis (CASA)

Three small openings were cut on the freshly harvested cauda epididymis from adult mice. The epididymis was placed in 37 °C preheated sperm culture medium (10% fetal bovine serum (FBS) in DMEM) and incubated for 5 min to allow the sperms to swim out. Then, 10 μ l of sperm suspension was subjected to CASA analysis (IVOS, Hamilton Thorne, USA).

Transfection and coimmunoprecipitation

HEK293T cells were transfected with plasmids for the expression of HA-TBP and FLAG-tagged TAF7L (WT or KI). Then, 48 h after transfection, the cells were harvested with lysis buffer (20 mM Tris pH 7.4, 150 mM NaCl, 0.5% Triton X-100, 0.5% sodium deoxycholate, 1 mM dithiothreitol (DTT),

protease inhibitor). Approximately 10 μ g of the indicated antibody was added into cell lysates with constant agitation at 4 °C overnight; 50 μ l of protein G dynabeads slurry (Invitrogen; 10004D) was added to pull down the immunocomplexes. Beads were collected with a magnetic grate and washed seven times with cold buffer (20 mM Tris pH 7.4, 150 mM NaCl, 0.5% Triton X-100, 1 mM EDTA). 50 μ l of 2 \times SDS loading buffer was added and the sample was boiled at 95 °C for 10 min. Protein samples were subjected to immunoblotting with appropriate antibodies.

Isolation of germ cell populations

Pachytene spermatocytes (pacSC) and round spermatids (rST) were isolated twice from adult testes using a STA-PUT apparatus within a 4 °C refrigerator [38]. For each isolation, three pairs of *Taf7l* WT or KI testes were subjected to first enzymatic digestion with 1 mg/ml collagenase IV (Gibco) in a 37 °C water bath for 15 min, followed by a second enzymatic digestion with 1 mg/ml trypsin (Sigma) and 1 mg/ml DNase I (Sigma) in a 37 °C water bath for 8 min. Then cells were resuspended in 25 ml of 0.5% BSA solution and filtered through a 100 mm mesh to remove cell aggregates and a 40 mm mesh to remove sperm tails. Next, 25 ml of cell suspensions was loaded and stirred gently. The stirrer worked under the 2% BSA solution and artery clips were removed so that both 2 and 4% BSA solutions were introduced to the chamber (~30 min). Artery clips were replaced and the stirrers were switched off, and a gradient of BSA was formed after 2 h of sedimentation in the standard cell sedimentation chamber. In order to collect different cell types with 15-ml centrifuge tubes, the first fraction collected was designated as 1 and the rest were numbered up to 100. The cells in each fraction were collected by centrifugation at 500 *g* for 5 min. The cells were resuspended in 0.2 ml of cold PBS. An aliquot of each fraction was examined under a DIC microscope to assess cellular integrity and identify the cell types. Fractions containing cells of similar size and morphology were pooled together and prepared for RNA extraction and RNA-seq analysis.

RNA-seq analysis

Total RNA was extracted from 3-month-old WT and *Taf7l*^{D144G/Y} mouse testes or isolated spermatogenic cells using TRIzol reagent (15 596 026, Invitrogen, USA). The RNA quality was assessed using an Agilent 2100 RNA 6000 Nano kit. Strand-specific RNA-seq libraries (three biological replicates per genotype) were constructed using the TruSeq stranded total RNA sample preparation kit (Illumina, USA) according to the manufacturer's instructions and sequenced on the Illumina HiSeqTM 4000 platform at Genedenovo Biotechnology Co., Ltd (Guangzhou, China). Low-quality read filtering and adaptor trimming were performed using fastP [39]. High-quality reads were mapped to the mouse genome (mm10) using HISAT2 [40] with a GTF file downloaded from the Ensembl genome browser 104. First, we used StringTie [41] to reconstruct the transcript and calculate the mRNA expression level of all genes in every sample. Second, CPC2 [42] and CNCI [43] were used to predict the coding ability of new transcripts. Transcripts without coding potential were classified as long noncoding RNAs (lncRNAs). Finally, unmapped reads with high quality were processed using the find_circ [44] pipeline to identify circular RNAs (circRNAs). For differential gene expression analysis of mRNAs and lncRNAs, we used DESeq2 [45] to normalize the read count and calculate the *P*-value and

Table 1. Summary of sequence variants in *TAF7L* found in oligozoospermic and control men

Position	Nucleotide change ^a	Resultant change ^a	113	175
Exon 4	c.407A→G	Missense mutation, D136G	1 ^b	0
Exon 5	c.633G→A	Silent	1	0
Intron 2	-47G→C	Intronic alteration	2	0
Intron 4	+42A→C	Intronic alteration	1	0
Polymorphisms				
Exon 9	c.922A→G	Missense mutation, S222G	19	25
Exon 10	c.1026del(TGAGGA)	Del(ED) in a stretch of EDED	22	40
Exon 13	c.1373G→A	Missense mutation, C370Y	2	4
Intron 7	-57C→G	Intronic alteration	1	1
Intron 8	+52A→G	Intronic SNP: rs3788764	44	61
Intron 10	+91A→G	Intronic SNP: rs3761514	40	60

^aReference sequences: human *TAF7L* cDNA, NM_024885; human TAF7L protein, NP_079161. ^bNumber of males with the given sequence variant.

false discovery rate value but used edgeR [46] for circRNA. Significant differentially expressed genes were identified as those with *P*-value <0.05 and fold-change >1.5.

Statistical analysis

Statistical analysis was performed using the two-tailed unpaired Student's *t*-test (ns: not statistically significant; **P* < 0.05; ***P* < 0.01; ****P* < 0.001).

Results

Mutation screening in the *TAF7L* gene in infertile men

TAF7L is a component of the transcription factor TFIID complex. *Taf7l* is an X-linked single-copy testis-specific gene in both mice and humans [11]. Targeted disruption of *Taf7l* in mice causes significant reduction in sperm count and sperm motility [14]. The most common form of male infertility is low sperm count. As the X chromosome is in the hemizygous state in men, we hypothesized that mutations in *TAF7L* could cause oligozoospermia (reduced sperm count) in humans. To screen for *TAF7L* mutations in infertile men, we sequenced 12 exons of *TAF7L* in 113 severely oligozoospermic men (sperm count <5 × 10⁶/ml) and 175 control men. We found a total of 10 sequence variants (Table 1). Six of these occurred in two or more samples with similar frequencies in both affected and control men and thus were likely to be polymorphisms. Among the six polymorphisms, two resulted in amino acid changes in exons 9 and 13, respectively, and another variant led to a deletion of two amino acids in exon 10 (Table 1). These three missense variants were also identified in infertile and fertile men in previous studies of nonobstructive azoospermic men and thus were not associated with infertility [19, 20].

However, four variants were observed only in one or two infertile men: one missense mutation (D136G), one silent mutation, and two intronic mutations. Therefore, only the D136G mutation is most likely to be deleterious (Figure 1A). The patient with this point mutation (D136G) (WHT3493, Figure 1A) and his female partner had experienced infertility for several years. The evaluation showed sperm counts ranging 4–6 million/ml, far lower than the normal sperm count of >15 million/ml. His karyotype and Y chromosomal microdeletion assay were both noninformative. Bilateral microsurgical varicocelectomy was performed with no demonstrable change in semen parameters. However, over the next several years, the couple had three naturally conceived pregnancies: two miscarriages on either side of the birth of

their healthy daughter. Unfortunately, the patient developed unilateral testis cancer but only required orchiectomy. Subsequent sperm counts settled out at an average of 3 million/ml. The couple did not return for follow-up visits or assessment.

D136 is highly conserved in the TAF7/TAF7L gene family

Although *Taf7l* is present only in mammalian species, *Taf7*, the ubiquitously expressed homologue of *Taf7l*, is present in all eukaryotes examined [14]. Aspartic acid (D136) is conserved not only in TAF7L but also in TAF7 proteins from diverse species including fission yeast (*Schizosaccharomyces pombe*), worms, fly, fish, and mammals (Figure 1B). In contrast, the residues adjacent to D136 are not as well conserved. TAF7 and TAF7L are components of the basal transcription factor TFIID complex that interacts with transcription activators to activate transcription. The residue D136 is located in the TAF7/TAF7L domain that is known to interact with transcriptional activators [12]. Therefore, the D136G mutation could disrupt the interaction between TAF7L/TAF7 and transcriptional activators, thereby inhibiting downstream transcription programs necessary for sperm production. The evolutionary conservation of D136 in TAF7L and TAF7 suggests that it is a functionally important residue.

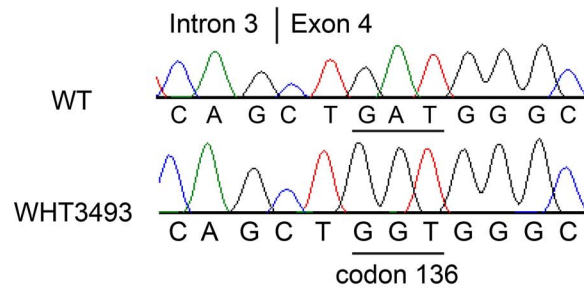
The D136G substitution in TAF7L is predicted to be deleterious

To gain more insights before direct functional studies, we used bioinformatic tools to evaluate the possible impact of the D136G substitution on the function of TAF7L. Four bioinformatics tools including SIFT, MutationTaster, PolyPhen-2, and PROVEAN were employed. Among them, the SIFT software is based on sequence homology and physical properties of amino acids. PolyPhen-2, MutationTaster, and PROVEAN predict the impact of an amino acid change on the structure and function of the protein based on a large number of sequences, evolutionary relationships, and structural features. All four analyses predicted that the D136G substitution was potentially detrimental to the function of TAF7L (Table 2).

The conserved D/N residue is required for the TAF function in budding yeast

The D136 (human TAF7L) is conserved in TAF7L and TAF7 in all species examined except *S. cerevisiae* (budding yeast) (Figure 1B). Budding yeast is the only species

A



B

Species	Accession No.	Sequence alignments
TAF7L		
		136
Homo sapiens	NP_079161	EHACTVRNLRASQSVKMKDKLKDILLPDRGHAVVEVEDVPLAAKLVDLPCVI
Macaca mulatta	XP_001092487	EHACTVRNLVHSQSVKMKDKLKDILLPDRRHAVVEVEDVPLAAKLVDLPCVI
Equus caballus	XP_001493193	EHASTVRKLLRSRSVSMKDKLKDIDSPDGRHAVVEVEAVSLTAKLVDLPCVI
Canis familiaris	XP_538106	EHASTVRKIVHSRSVAMKDKLKDIDSSDGRHAIIVEVEEISLAAKLVDLPCVI
Bos taurus	XP_872680	EHASTVRKMVHSGSSSMKNKLDLFS DTRHAVVEVDNVSLAAKLVDLPCVI
Rattus norvegicus	XP_001078826	EHASTVRKIIRSGNAAMRDKLDLSPDSRHAIVQVDDVLSAKVVDLPCVI
Mus musculus	NP_083234	EQAYAVRKIIHSRNAAWKDKLKDIDFSPDGHHAVVQVDNVSLPAKLVDLPCVI
TAF7		
Homo sapiens	NP_005633	EYASTVRRVQSGHVNLKDRLTIELHPDGRHGIVRVDRVPLASKLVDLPCVM
Mus musculus	NP_786964	EYAATVRRVQSGHVNLKDKLSIELHPDGRHGIVRVDRVPLAAKLVDLPCVT
Rattus norvegicus	NP_001100864	EYAATVRRVQSGHVNLKDRLSIELHPDGRHGIVRVDRVPLAAKLVDLPCVT
Bos taurus	NP_001039493	EYASTVRRVQSGHVNLKDRLSIELHPDGRHGIVRVDRVPLAAKLVDLPCVM
M. domestica	XP_001369129	EYASTVRRVQSGSVNLKDKLTIELHADGRHGIVRVDRAPLAAKVVLDLPCVT
O. anatinus	XP_001513417	ESAATVRRVQSSASVSPKDRVTIELHPDGRHGIVRVDRVPLAAKLVDLPCSI
Gallus gallus	XP_420187	EYASTVRRVQSGSVNLKDRLTIELHADGRHGIVRVDRVPLAAKLVDLPCII
Xenopus laevis	NP_001086631	EYASTVRRVQSGSINTKDRLSIELHPDGRHGIVRVDRVPLAAKLVDVPCVT
Salmo salar	NP_001133720	EYASTVRCIAQSSMNLKDRLTIELHADGRHGIVRVDRVPLACKLVDLPCIL
Equus caballus	XP_001918058	EYASTVRRVQSGHVNLKDRLTIELHPDGRHGIVRAHCAPLIK---NLPWVM
T. guttata	XP_002187302	EYASTVRRVQSGNVNLKDRLTIELHADGRHGIVRVDRVPLAAKLVDLPCIT
T. nigroviridis	CAG09835	EYASTVRQIAQSSMNMKDRLTIELHPDGRHGIVRVDRVPLACKLVDLPCMI
Danio rerio	NP_775367	EYASTVRRVQSGSINTKDRLSIELHPDGRHGIVRVDRVPLACKLVDLPCIL
A. thaliana	NP_175926	VSERIDRLLEDASTSDEIPLDLFFSEDRNGTFMIGNDEFASLLDLPAVV
D. melanogaster	AAC39086	ELADTVHEAINAGTI--KDRLTIQLDPLDRLYGEVRIIDDQIILYTKLVDLPTVV
C. elegans	NP_499647	DCVGRIEKIIQSDGK--HEEFSLNLSN DARNSTVRIGNQLLNGKILDLPTIT
S. pombe	NP_593650	EDCEYVRKAIENREVGGRADIWVKFK-DQRRAVVHVNGHLYAAKLVDLPCII
S. cerevisiae	NP_013954	IQLFVKNLSLES---GDYSGISIKWK-NERHAVVTINDVMYGAILVDLPTVI
Conservation		. . . : : : . . . : : *

Figure 1. Identification of a missense mutation (D136G) in human *TAF7L* gene in an oligozoospermic man. (A) Sequencing chromatograms of human wild type and mutant *TAF7L* alleles. Patient WHT3759 harbors a missense mutation in exon 4 of *TAF7L*, resulting in the D136G amino acid substitution. (B) Conservation of the aspartic acid at 136 (D136) in *TAF7L* and *TAF7* proteins among eukaryotes. The position of the conserved aspartic acid/asparagine residue: D136 (human *TAF7L*), N147 (budding yeast *TAF7*), and D144 (mouse *TAF7L*). Sequence alignment was performed with Clustal X.

Table 2. Bioinformatic prediction of possible functional impacts of the missense mutation (c. 407A>G; p.D136G) in the human *TAF7L* gene

Bioinformatic tools	Prediction
Mutation Taster	Disease causing
PolyPhen-2	Probably damaging
PROVEAN	Deleterious
SIFT	Deleterious

that we found to contain asparagine (N) instead of aspartic acid (D) at this position. Notably, asparagine and aspartic acid are structurally similar amino acids. We found that *TAF7*-null yeast mutants were not viable, showing that *TAF7* is an essential gene in budding yeast (Table 3). We next performed tetrad analysis to determine whether the N147 in yeast *TAF7* (corresponds to D136 in human *TAF7L*) is required for its function. We cloned WT yeast *TAF7* (sc*TAF7*-N147) into a

Table 3. Yeast (*S. cerevisiae*) complementation assay

Plasmid	<i>taf7Δ</i> (null) yeast
None	nonviable
scTAF7-N147(WT)	viable
scTAF7-N147D	viable
scTAF7-N147G	nonviable

low-copy CEN plasmid pRS416. As expected, the pRS416-scTAF7-N147 rescues the lethality of TAF7-null yeast cells (Table 3). We made two TAF7 mutants harboring N147D and N147G point mutations respectively, resulting in two constructs pRS415-scTAF7-N147D and pRS415-scTAF7-N147G. We found that scTAF7-N147D, like the WT TAF7-N147, rescued the lethality of TAF7-null yeast cells. This was expected because the aspartate residue is conserved in all other species, including the fission yeast (*S. pombe*). Notably, however, the scTAF7-N147G (analogous to the infertility-associated human mutation) did not rescue the lethality caused by TAF7 deletion in yeast (Table 3), showing that this conserved D/N residue is required for the function of TAF7.

Taf7l^{D144G/Y} male mice exhibit apparently normal fertility

To assess if this point mutation in TAF7L contributes to human male infertility, we generated *Taf7l*^{D144G} knockin mice, which harbor the D144G point mutation (analogous to the human D136G mutation) (Figure 2A). The point mutation was located in the fourth exon of *Taf7l* (Figure 2A). The WT and knockin *Taf7l* alleles were assayed by PCR (Figure 2B). Sanger sequencing of PCR products confirmed the point mutation in mice (Figure 2C). The *Taf7l*^{D144G} allele was backcrossed to the C57BL/6J genetic background for 10 generations. Quantitative RT-PCR and western blot analyses of adult testes (3-month-old) showed no significant difference in mRNA and protein levels of *Taf7l* between WT (*Taf7l*^{+/+}) and *Taf7l*^{D144G/Y} testes (Figure 2D and E).

Taf7l^{D144G/Y} male mice displayed no overt developmental defects and sired offspring. The testis size of *Taf7l*^{D144G/Y} males was comparable to that of WT males (Figure 3A). Body weight, testis weight, and epididymis weight were similar between WT and *Taf7l*^{D144G/Y} males (Figure 3B–D). CASA showed no difference in epididymal sperm count or the percentage of sperm with progressive motility between WT and *Taf7l*^{D144G/Y} males (Figure 3E and F). All other sperm motility parameters for *Taf7l*^{D144G/Y} males were normal as well (Supplementary Table S3). Mating tests were performed by mating WT or *Taf7l*^{D144G/Y} males with *Taf7l*^{D144G/+} females for 10 months. The total number of litters sired was similar between WT and *Taf7l*^{D144G/Y} males (Figure 3G). The average litter size was lower in *Taf7l*^{D144G/Y} males than in WT males, but the difference was not statistically significant (Figure 3G). Histological analysis of testes revealed apparently normal spermatogenesis in seminiferous tubules from *Taf7l*^{D144G/Y} testes, which contained the full spectrum of spermatogenic cells: spermatogonia, spermatocytes, and post-meiotic spermatids (Figure 3H). Mature sperm were abundant in the cauda epididymides of WT and *Taf7l*^{D144G/Y}

males (Figure 3H). These results indicated that the fertility of *Taf7l*^{D144G/Y} males was normal.

To test whether the D144G substitution in TAF7L affects its interaction with TBP, we constructed expression plasmids for HA-TBP, FLAG-TAF7L (WT), and FLAG-TAF7L (D144G). Following transfection in HEK293T cells, co-immunoprecipitations coupled with the western blot assay showed that the TAF7L (D144G) protein was still able to interact with TBP (Figure S1).

Alterations in the transcriptome of *Taf7l*^{D144G/Y} testes

As TAF7L is a transcription factor, we performed RNA-seq analysis of WT and *Taf7l*^{D144G/Y} testes to address whether the D144G substitution causes transcriptomic changes. Principal component analysis showed that the three biological replicates for each genotype were apparently clustered, except for a relative divergence of one sample per genotype (mRNAs, Figure 4A; lncRNAs, Supplementary S2A; circRNAs, Figure S2B). The intergenotype divergence and intragenotype convergence were clearly shown by the sample correlation analyses by Pearson's method (mRNAs, Figure 4B; lncRNAs, Supplementary Figure S2C; circRNAs, Supplementary Figure S2D). We identified differentially expressed protein-coding genes (DE genes) in *Taf7l*^{D144G/Y} testis with the cutoff of >1.5 fold-change and $P < 0.05$: 165 upregulated genes and 77 downregulated genes (Figure 4C). In addition, our analysis revealed the altered expression of 507 lncRNAs (Supplementary Figure S2E) and 135 circRNAs (Supplementary Figure S2F) in *Taf7l*^{D144G/Y} testis. Furthermore, we prepared populations of pacSC and rST from WT and *Taf7l*^{D144G/Y} testes by the STA-PUT method (Supplementary Figure S3), and performed RNA-seq analysis (Supplementary Figures S4 and S5). For both cell types, the intragenotype replicates were much more uniform than the intergenotype groups (Supplementary Figure S4A–D). We identified 38 upregulated and 227 downregulated genes in *Taf7l*^{D144G/Y} pacSC (Supplementary Figure S5A and C). In *Taf7l*^{D144G/Y} rST, 42 genes were upregulated and 80 downregulated (Supplementary Figure S5B and D). A selection of differentially expressed DE genes in *Taf7l*^{D144G/Y} rST were validated by qRT-PCR (Supplementary Figure S5E and F; Supplementary Table S4). We then analyzed the overlap of DE genes in the *Taf7l*^{D144G/Y} testes with those reported in the *Taf7l* knockout testes [14, 15] and found minimal overlaps (Figure 4D and E). These results indicate that the D144G substitution alters the testicular transcriptome, and these alterations are not severe enough to impact fertility in the *Taf7l*^{D144G/Y} males.

Discussion

Here we identified a missense mutation (D136G) in TAF7L in an oligozoospermic man. This aspartate residue is critical for TAF7 function in budding yeast. Interestingly, a single amino acid substitution (G690D) in TAF1, an X chromosome-encoded TAF that is the largest subunit of TFIID, confers a temperature-sensitive phenotype in the hamster ts13 cell line [47]. TAF1L is an autosomal retrotransposed homologue of TAF1 and is testis-specific in humans [33]. The WT TAF1L rescued the temperature-sensitive lethality of hamster ts13 cells but a mutant

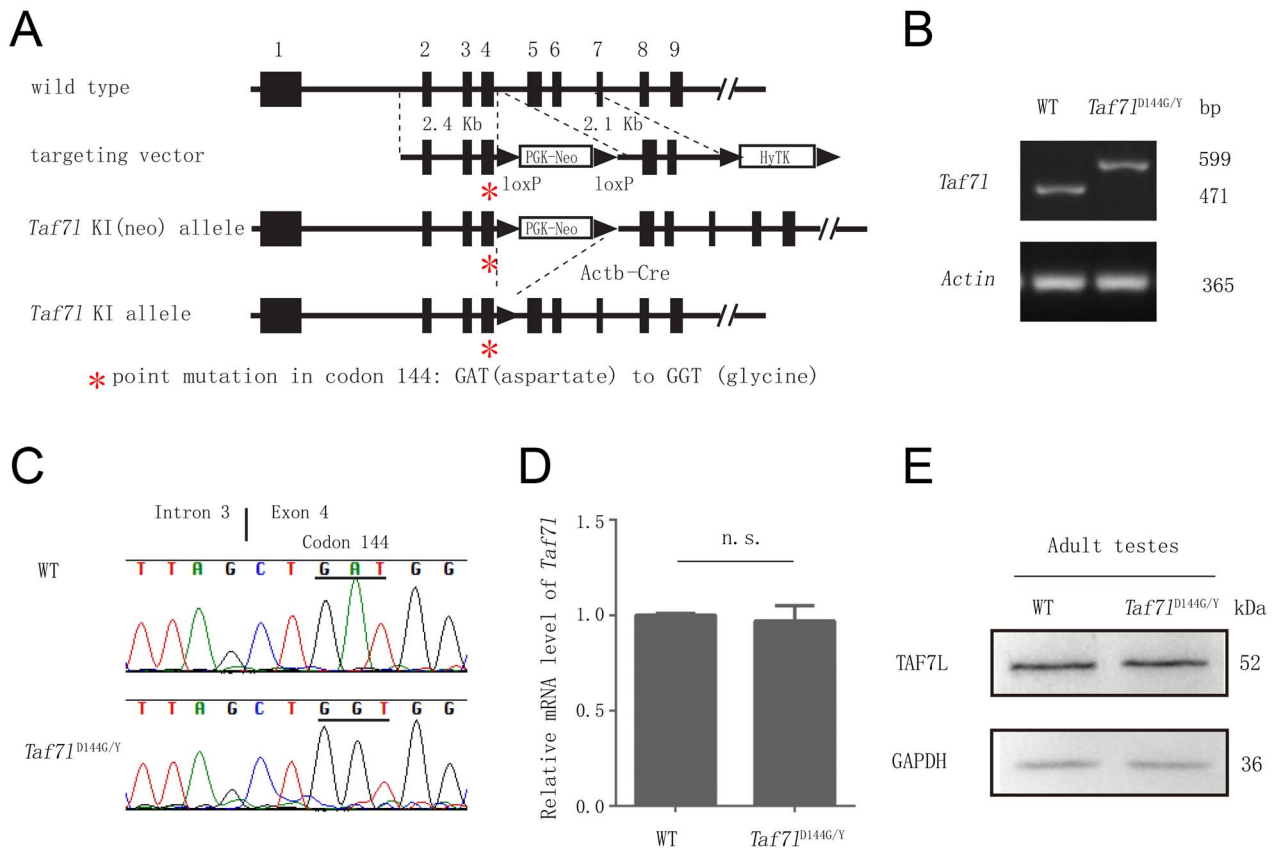


Figure 2. Generation of *Taf7*^{D144G} knockin mice. (A) Schematic of mouse *Taf7* gene structure, targeting vector, and various alleles. *Taf7* consists of 13 exons. Only the first nine exons are shown in rectangles. Exons 10 to 13 are not shown. The asterisk in Exon 4 denotes the point mutation. PGK-Neo encodes a neomycin selection marker. HyTK enables positive selection with hygromycin (not used) and negative selection with ganciclovir. pOG231 expresses Cre recombinase. (B) Genotyping of *Taf7* alleles. (C) Sequencing of the wild type and *Taf7*^{D144G} alleles. (D) qRT-PCR analysis of the *Taf7* transcript in wild type and *Taf7*^{D144G/Y} testes from 3-month-old mice. n.s.: nonsignificant, by Student's *t*-test. (E) Western blot analysis of TAF7L in 3-month-old wild type and *Taf7*^{D144G/Y} testes. GAPDH serves as a loading control.

TAF1L (G714D; analogous mutation) did not, indicating that TAF1L and TAF1 are functionally equivalent [33]. These studies point to the selection pressure (from meiotic silencing of sex chromosomes in male mammals) on X-linked TAFs and the continuous evolution of testis-specific TAF homologues.

Although the *Taf7* knockout male mice are subfertile with reduced sperm count [14], the *Taf7*^{D144G/Y} males have normal sperm count. The difference in the phenotypes of these two mouse mutants (knockout versus point mutation) could reflect functional compensation by TAF7. TAF7 compensation might be sufficient to restore fertility in *Taf7*^{D144G/Y} mice with compromised TAF7L function but not in knockout mice with a complete loss of TAF7L. TAF7 is essential for embryonic development [48]. Therefore, it might be informative to generate *Taf7*^{+/-} *Taf7*^{D144G/Y} mice or *Taf7*^{+/-} *Taf7*^{L^{-/-}} mice in future studies to determine whether *Taf7* heterozygosity leads to more severe spermatogenic defects in *Taf7* mutant mice.

As exome and whole-genome sequencing has become increasingly cost-effective, a large number of human patients can be screened and as a result, an overwhelming number of sequence variants have been identified. However, the challenge lies in the prediction and interpretation of the functional

consequences of these variants. Although the deleterious effect of nonsense mutations is predictable, the functional relevance of missense mutations or intronic variants is difficult to infer. Bioinformatic tools provide predictions as to the possible pathological impact of variants but cannot substitute for experimental validation. Although variants for somatic functions can be tested in cell lines, this becomes more challenging for testing variants in human germ cells because a human in vitro gametogenesis system is yet to be developed [49]. Therefore, animal modeling of sequence variants identified in infertile patients remains the only option for functional studies. We have investigated the functional consequences of the human D136G substitution in TAF7L identified in an oligozoospermic man by modeling this substitution in budding yeast and mouse. While aspartic acid and asparagine are functionally interchangeable in the yeast TAF7 gene, G at this position leads to lethality in yeast (Table 3). We also modeled this mutation in *Taf7*^{D144G/Y} mice. Although *Taf7*^{D144G/Y} testes exhibit transcriptomic changes, *Taf7*^{D144G/Y} mice have normal sperm count, normal sperm motility, and normal fertility. There are several possible explanations for the difference in phenotypes for this amino acid substitution between mouse and human. First, there might be differences in species-specific requirements for TAF7L. Although the human

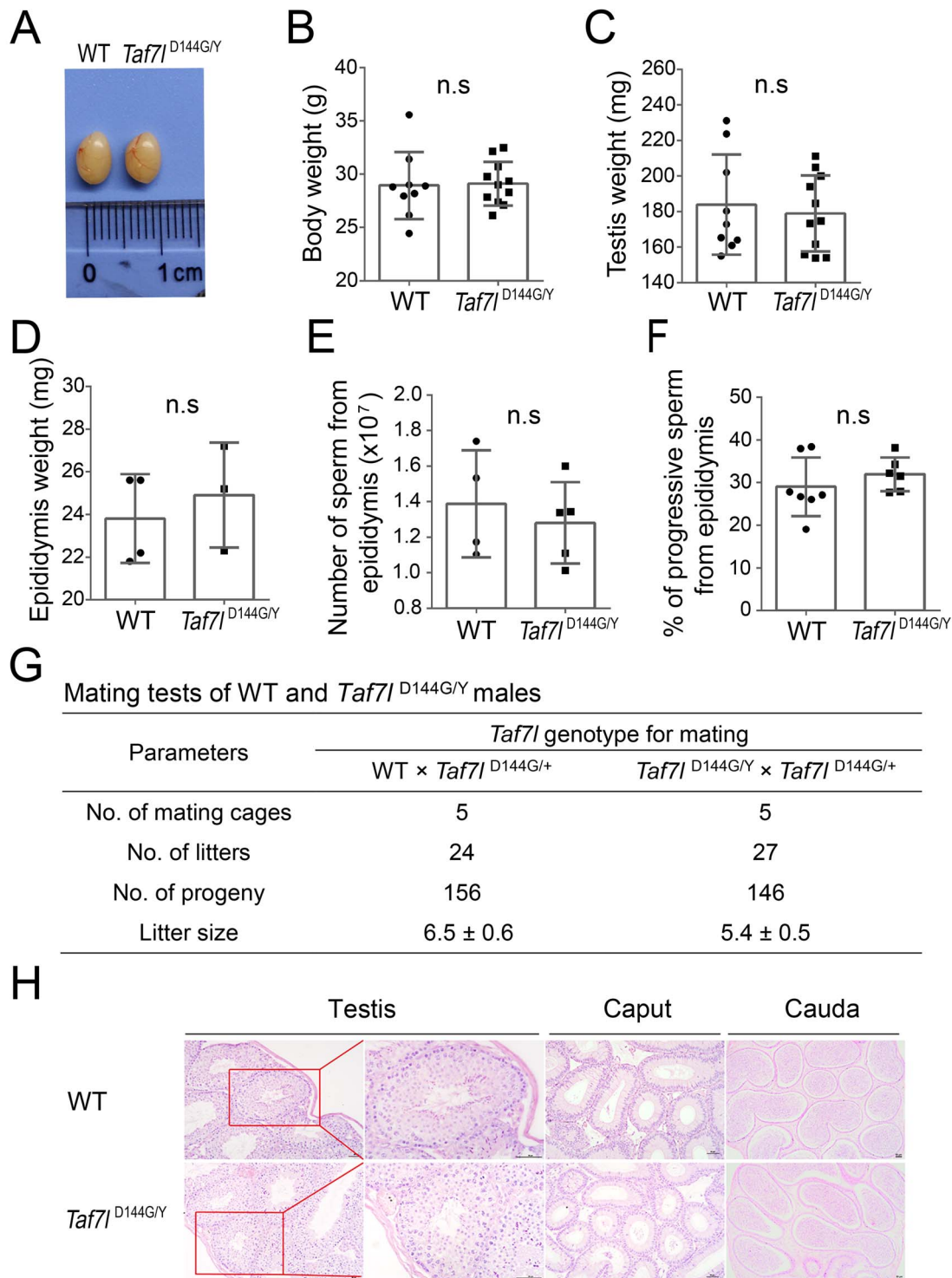


Figure 3. Characterization of the *Taf7l*^{D144G/Y} male mice. (A) Images of testes from WT and *Taf7l*^{D144G/Y} males (3 months old). (B–F) Body weight (B), testis weight (C), epididymis weight (D), epididymal sperm count (E), and percentage of epididymal sperm with progressive motility (F) of 3-month-old WT and *Taf7l*^{D144G/Y} mice. The mean \pm SEM is plotted. *n*, number of mice. n.s.: nonsignificant. (G) Mating test results. Each male was housed with two females for 10 months. (H) Histology of testis, caput epididymis, and cauda epididymis from 3-month-old WT and *Taf7l*^{D144G/Y} males. Scale bar: 50 μ m.

and mouse TAF7 proteins share 90% sequence identity, the human and mouse TAF7L proteins exhibit only 57% sequence identity, showing that the TAF7L proteins are more diverged. Thus, it is possible that human spermatogenesis is more dependent on TAF7L than mouse spermatogenesis. Second,

the human patient may have variants in other genes that have synthetic deleterious effects with the TAF7L D136G substitution. Third, given dysregulation of some noncoding RNAs (lncRNAs and circRNAs) in the *Taf7l*^{D144G/Y} testis, a fertility defect might only be induced under certain stress

Data availability

The RNA-seq data are available from the National Center for Biotechnology Information's GEO database (www.ncbi.nlm.nih.gov/geo/) under accession number GSE186296.

Authors' contributions

K.Z., L.L., F.F.L., P.L.Y., and Q.L.Y. carried out the experiments. R.D.O. and S.S. provided patient samples. P.J.W., H.S., L.G.B., and S.R. carried out the mutation screening. P.J.W., D.C.P., and K.Z. analyzed data. C.K. and F.C.L. performed the yeast experiments. N.A.L. performed blastocyst injection. J.W.Z. performed RNA-seq analyses. P.J.W., K.Z., L.L., and F.F.L. wrote the manuscript.

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Conflict of interest

The authors have declared that no conflict of interest exists.

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