A novel SoxB2 gene is required for maturation of sperm nucleus during spermiogenesis in the Chinese mitten crab, *Eriocheir sinensis*

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**SRY**-related HMG box (Sox) genes are characterized by the presence of a DNA-binding HMG domain and involved in a diverse range of developmental processes. In this study, we identified a novel Sox gene, designated as *EsSoxB2-1*, from the Chinese mitten crab *Eriocheir sinensis*. The *EsSoxB2-1* encodes a protein of 259 amino acids, sharing the highest identity with the beetle *Tribolium castaneum* SOX21b. Unlike insect *Sox21b*, however, *EsSoxB2-1* is intronless and exhibits a gonad-specific expression pattern at both mRNA and protein level. Two core promoters in 5' flanking region were demonstrated to be essential for inducing transcriptional regulatory activity. The transcription of *EsSoxB2-1* mRNA begins in spermatogonia stage, while the translation of *EsSOXB2-1* protein initiates at spermiogenesis stage. Interestingly, *EsSOXB2-1* protein was exclusively localized in the nucleus of spermatid and spermatozoa even at the end of acrosome reaction, and was bound to the uncondensed chromatin in nucleoplasm of mature spermatozoa. Knockdown of *EsSoxB2-1* by RNAi leads to abnormal transformation of the nucleus during spermiogenesis. Together, these findings demonstrated the requirement of *EsSoxB2-1* for the spermatozoa nucleus maturation and also suggested that *EsSoxB2-1* would be delivered into fertilized eggs along with chromatins as a paternal transcription factor for regulating early embryonic development.

**SRY** was first discovered in human Y chromosome as a sex-determining factor, and possesses a conserved high-mobility group (HMG) box**1,2.** The HMG box contains 79 amino acids and is responsible for binding the minor groove of DNA at the site of specific target sequences, (A/T)(A/T)CAA(A/T)G**3,4, to regulate transcription of downstream genes in the sex-determining cascade by altering chromatin structure**5,6.** Although **SRY** homologues are found only in placental mammals and marsupials and are not evolutionally conserved in the animal kingdom**7,8, a large **SRY**-related HMG box (Sox) gene family of transcription factors was subsequently identified in both vertebrates and invertebrates**2,8,9. The members of Sox family play pivotal roles in cell differentiation and embryonic organogenesis including gonadogenesis, neurogenesis, chondrogenesis, and oligodendrocyte development**10-12. To date, over 30 members have been isolated and classified into ten groups (A-J) mainly based on their sequences similarity of HMG box. These groups are: A, SRY/Sry; B, Sox1, Sox2, Sox3, Sox14 and Sox21; C, Sox4, Sox11, Sox12, Sox22 and Sox24; D, Sox5, Sox6, Sox13 and Sox23; E, Sox8, Sox9, and Sox10; F, Sox7, Sox17, and Sox18; G, Sox15 and Sox20; H, Sox30; I, Sox31; J, Sox J. Members of the same group usually share over 70% amino acids identity both within and outside the HMG domains**7. Although all five members of the group B genes are intronless in vertebrates, sequence analysis and functional studies suggested that the group B Sox genes can be subdivided into two further groups; B1; Sox1, Sox2 and Sox3; and B2; Sox14 and Sox21**13. The three subgroup B1 members act as transcriptional activators, while the B2 members (SOX14 and -21) are transcriptional repressors**13. In insects, four group B genes, SoxNeuro (SoxN), *Dichaete*, Sox21a and Sox21b, have been identified so far**6. SoxN is associated with group B1 and the latter three are physically linked in the genome and assigned to group B2. However, *SoxN* and *Dichaete* are intronless while *Sox21a* and *Sox21b* bear introns**14.

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Gonad or sex differentiation is the complicated differentiation models of organogenesis entailing biochemical and morphological reconstruction of the germ cell. A number of Sox genes have been implicated in these complicated procedures. Besides Sry acting as a master sex determination gene in mammals, some other Sox genes are involved in sex differentiation and germ cell development based on the data of their expression profiles. In mice, Sox3 is located on X chromosome and is regarded as the ancestor of Sry. Sox3 is not required for sex determination, but is needed for normal gametogenesis such as male testis differentiation. Targeted deletion of Sox3 caused abnormal development of spermatogonia. Sox9 is expressed shortly after Sry in pre-Sertoli cells and was confirmed to be the only target of SRY in mediating a switch from the ovarian pathway to the testicular pathway. And the specific expression of Sox3 in the normal testes, but not in the germ cell-deficient testes, suggesting the involvement of Sox3 in the differentiation of mouse male germ cells. Despite Sox genes have been cloned and characterized in a wide variety of taxonomic groups, rare molecular data of Sox genes has been documented in crustaceans so far. In the present study, we identified a novel Sox gene, termed EsSoxB2-1, which displayed gonad-specific expression in the Chinese mitten crab, Eriocheir sinensis. The EsSOXB2-1 protein was exclusively localized in the nucleus of spermatogenic germ cells and was revealed to be involvement in sperm nucleus maturation.

**Results**

Full-length sequence of the crab *EsSoxB2-1* cDNA. A cDNA fragment of 220 bp was amplified from the ovary by degenerate RT-PCR. The amplified fragment contained a conserved HMG box and was shown to be a Sox homologue by Blast analysis. Then the full-length cDNA of this Sox homologue was generated by 5′ and 3′ RACE. This cDNA is 951 bp in length and contains a 5′ untranslated region (UTR) of 54 bp, a open reading frame (ORF) of 777 bp, and a 3′ UTR of 111 bp with a poly(A) tail. The ORF encodes a polypeptide of 259 amino acids (aa) with a predicted molecular weight of 28.55 KD. Amino acid sequence alignment showed the HMG box of the crab Sox homolog the highest (95%) identity with those of *Tribolium castaneum* SOX21b, but there is little (about 36%) sequence similarity outside the HMG box between them. The HMG box contains a consensus sequence RPMNAFMVW and four histidine residues (22, 49, 83, and 87aa) that are thought to be essential for the DNA-binding properties (Supplementary Figure S1). The crab Sox homolog also contains two nuclear localization signals (NLS) K24RPMNAFMVWSRMQRRK40 and R93PRRKPKT100, a nuclear export-signal (NES) and a small hydrophobic leucine-rich motif (ISKRLGEWKL), but lack a subgroup B motif (Fig. 1A), which appears next to the HMG box in vertebrate Sox group. Unlike *Tribolium castaneum* SOX21b, the crab Sox homolog contained one poly-alanine stretches at the carboxyl terminus (Fig. 1A). Accordingly, we designated this novel Sox homolog as *EsSoxB2-1*.

Phylogenetic analysis. In an effort to determine the phylogenetic affinities between the crab *EsSoxB2-1* and other members of Sox family, an unrooted phylogenetic tree was constructed by the NJ method using the multialignment of complete protein sequences of various metazoans Sox including *Mus musculus* SRY. As shown in Fig. 1B, the previously established groupings of Sox are supported by the tree. Various Sox groups were assigned into different clades. The crab *EsSoxB2* falls into SoxB group and was first clustered with the beetle *Tribolium castaneum* and the fruit fly *Drosophila melanogaster* SOX21b, and then they were together clustered with SOX21a and Dichaete in Sox B2 group, although with less confidence.

Gene structure and promoter activity of the crab *EsSoxB2-1*. The genomic sequence of *EsSoxB2-1* gene was amplified by PCR using a pair of gene-specific primers set at the 3′ and 5′ ends of the cDNA sequence. The retrieved genomic sequence was in excellent accord with its corresponding cDNA sequence (Supplementary Figure S1), indicating that the novel *EsSoxB2-1* gene contains no intron.

The 5′-flanking sequence of *EsSoxB2-1* gene was obtained by genome walking method. Sequence analysis showed that the 1509 bp 5′-flanking region contained two basal core promoters (−628/−577 bp and −493/−444 bp) and a 377 bp CpG island (−299/+78 bp). Many potential transcription factor binding sites were identified in the region from the transcription start site (TSS) to the core promoters. Some of binding sites were given more attention such as SRY/Sox, CATA, GATA-1, E2F, CREB and TATA (Supplementary Figure S2). The promoter activity in 5′-flanking region of the *EsSoxB2-1* gene was assayed using Dual-Luciferase Reporter Assay System (Promega). To determine which fragment within the 5′-flanking region responsible for transcriptional regulatory activity, various lengths of the 5′-flanking sequences F1 (−189/+88 bp), F2 (−430/+88 bp), F3 (−545/+88 bp), F4 (−714/+88 bp) and F5 (−1483/+88 bp) were cloned into the promoterless pGL3-Basic plasmid containing luciferase genes, respectively (Supplementary Figure S2). Significantly high activities were detected in F3, F4, and F5 (Fig. 2). The promoter activity of F4 (−714/+88 bp) was much higher than that of the full length of the 5′-flanking region F3 (−1483/+88 bp), indicating the existence of silencing sequence elements within the fragment from −1483 to −714 bp. When excluding one core promoter (−628/−577 bp), the promoter activity of F3 displayed significant reduction compared to F4 (−714 to +88) (Fig. 2). When excluding both of the two core promoters (−628/−577 bp and −493/−444 bp), the promoter activity of F2 (−430/+88 bp) and F1 (−189/+88 bp) became extremely low similar with the pGL3 Basic empty plasmid (Fig. 2). These data demonstrated that the two core promoters (−628/−577 bp) are essential for inducing transcriptional regulatory activity.

**Tissue distribution of *EsSoxB2-1* mRNA and *EsSOXB2-1* protein.** Interestingly, RT-PCR analysis of seven types of adult tissues showed that *EsSoxB2-1* was exclusively expressed in testis and ovary, but not found in any other somatic tissues examined. The amount of *EsSoxB2-1* transcripts is higher in ovary than in testis, with especially higher in immature ovary and testis. As an internal reference, β-actin was simultaneously amplified using the same cDNA samples, and similar amounts of amplified product were obtained from all of the tissues (Fig. 3A).
Western blot analysis was performed by using the rabbit anti-\( \text{EsSOXB2-1} \) serum. A specific target band of about 30KD protein was detected in the testis and ovary (Fig. 3B), while no signal was seen in the negative control using the normal (preimmune) rabbit serum instead of the primary specific antibody (data not shown).
As a loading control, β-Actin protein was detected in all the tissues examined. These data demonstrated that EsSOXB2-1 protein also displayed specific expression in the testis and ovary. However, unlike mRNAs, the EsSOXB2-1 protein is expressed much higher in testis than in ovary (Fig. 3B), suggesting that EsSoxB2-1 could mainly function in testes. Thereafter we focused on the characterization of its potential role in testis.

**Localization of the EsSoxB2-1 mRNA and EsSOXB2-1 protein in the testes.**

Like most decapods, the crab testis is formed by numerous seminiferous tubules that contain germ cells at various developmental stages. As shown in Fig. 4, the spermatogenesis process consisted mainly of four developmental stages: (1) spermatogonia, (2) spermatocytes, (3) spermatids and (4) spermatozoa. Spermatogonia were larger in size and each spermatogonium contains a thin rim of cytoplasm around a vesicular nucleus (Fig. 4A). The spermatocytes have an irregularly shaped nucleus that is larger than that of the spermatogonia and is stained by hematoxylin (Fig. 4B), while the cytoplasm is indistinct and acidophilous. Spermatids are smaller and round in shape. Their nuclei show deep stain while the cytoplasm stained grey with eosin (Fig. 4C). In spermiogenesis, the spermatids transform into spermatozoa. An acrosomal vesicle appears in the vicinity of the nucleus and is gradually surrounded by the nucleus. Finally, the cup-shaped nucleus is positioned in the periphery and the acrosomal complex located in the central region of the mature spermatozoa (Fig. 4D,E).

To examine spatio–temporal expression of the RNA and protein of EsSOXB2-1 in the crab spermatogenesis, tissue sections of testes in a breeding period were subjected to in situ hybridization and immunohistochemical analysis, respectively. When using a DIG-labeled antisense RNA probe, strong signals were detected in early development germ cells including spermatogonium and spermatocyte (Fig. 4E,G), but not in spermatid and spermatozoa (Fig. 4H). No signal was detected in a negative control using a sense RNA probe (Fig. 4I). Contrary to its corresponding mRNA localization, the crab EsSOXB2-1 protein was detected only in the nucleus of spermatid and spermatozoa (Fig. 4L,M), whereas no immuno–signal was found in spermatogonia (Fig. 4J) and spermatocyte (Fig. 4K) as in the negative controls (Fig. 4N).

**Subcellular localization of the EsSOXB2-1 protein with immunogold labeling.**

To further examine the subcellular distribution of EsSOXB2-1 protein in the nucleus of spermatozoon, immuno-electron microscope technic (IEM) was performed. As shown in Fig. 5, the mitten crab spermatozoa are allagellated. Under
electron microscopy, the crab spermatozoon contains a complicated acrosome surrounded by an uncondensed nucleus with radial arms. The acrosomal complex is composed of a lot of subcellula structures like acrosomal tubule, apical cap, acrosomal vesicle, etc (Fig. 5A). In the nucleus, the uncondensed chromatins appear as loose fibrous evenly suspended in the nucleoplasm. The immuno-gold granules of EsSOXB2-1 protein were localized on the uncondensed chromatin fibers (Fig. 5B,C). No gold signal was detected in the IEM control section incubated with normal (preimmune) rabbit serum (Fig. 5D).

Knockdown expression of EsSoxB2-1 by RNAi. To determine the role of EsSoxB2-1 in spermiogenesis, RNAi was performed by in vivo injection of dsRNA targeting the coding region of EsSoxB2-1. The knockdown expression levels of EsSoxB2-1 in testis were assayed by qPCR and Western blot analysis. At 24 hours
post injection, a notably reduction of \textit{EsSoxB2-1} mRNA (about 46%) and protein (about 43%) was detected in \textit{EsSoxB2-1}-dsRNA injected group (Fig. 6A,B), while the expression of \textit{EsSoxB2-1} has no change in control groups after GFP-dsRNA or PBS injection. Subsequently, a lower expression of \textit{EsSoxB2-1} protein was found at 48 hours post injection of \textit{EsSoxB2-1}-dsRNA (Fig. 6B). The phenotype effects of RNAi knockdown were further observed after \textit{in vivo} repetitive injection for a month. In comparison with normal testicular development in control groups, the size and histological morphology of testes seems to be similar in \textit{EsSoxB2-1}-dsRNA injected group (data not shown). Under transmission electron microscope, however, the mature spermatozoa display abnormal structure. In normal spermatozoa, the cup-shaped nucleus generally extends into radial arms (Fig. 7A), whereas the nucleus arms degraded into many high dense electronic granules after RNAi (Fig.7B–D). Therefore, we concluded that the \textit{EsSoxB2-1} plays an essential role in forming/maintaining nucleus arms of spermatozoa.

Given that histones H3 and H4 were previously revealed to be involved in maintaining uncondensed nucleus\textsuperscript{25}, the histones H3 and H4 mRNA levels were additionally assayed at 24 hours and 48 hours post injection. No change was found after knockdown of \textit{EsSoxB2-1} expression (Supplementary Figure S3), indicating that the mRNA expression of histones H3 and H4 is not regulated by \textit{EsSoxB2-1}.

**The detection of \textit{EsSOXB2-1} protein in spermatozoa during acrosome reaction.** To investigate whether \textit{EsSOXB2-1} protein has a potential role in fertilization, \textit{EsSOXB2-1} protein was traced during the acrosome reaction induced \textit{in vitro} with CaCl\textsubscript{2} (Fig. 8A–J). The procedure of acrosome reaction is divided into four typical steps as described by Du \textit{et al.}\textsuperscript{26}, namely (1) protrusion of the apical cap (Fig. 8B), (2) eversion of the acrosomal vesicle and contraction of the nuclear cup (Fig. 8D), (3) disappearance of the acrosomal vesicle and the completion of the reaction (Fig. 8E). Immunocytochemistry analysis showed that the immuno-signals of the \textit{EsSOXB2-1} protein were precisely localized in the nuclear throughout all the four stages of acrosome reaction (Fig. 8G–J), indicating that the \textit{EsSoxB2-1} is not involved in the process of acrosome reaction.
Discussion

The group B Sox genes attract particular interest since they are most closely related to SRY and appear to be functionally conserved during evolution among mammals\textsuperscript{2,14}. In the present study, we identified a novel Sox homolog \textit{EsSoxB2-1} from the mitten crab, \textit{Eriocheir sinensis}. The encoding protein contains a conserved HMG box sharing the highest (about 95\%) identity with the beetle \textit{T. castaneum} SOX21b, but there is little similarity in sequence outside the HMG box between them. Further, similar to vertebrate \textit{Sox21}, \textit{EsSoxB2-1} is intronless in the coding region, that is contrary to insect \textit{Sox21} gene with multi-intron structure\textsuperscript{14,27}. Also, the putative \textit{EsSOXB2-1} protein contains a polyalanine stretch at C-termini, whereas the polyalanine stretch is absent in the fruit fly \textit{D. melanogaster} SOX21b (Fig. 1A). Unlike vertebrate group B Sox genes, however, both the crab \textit{EsSOXB2-1} and insect \textit{SOX21} lack a unique motif for subgroup B members (Fig. 1A). In terms of the gene expression pattern, insects and vertebrates \textit{Sox21} mainly expressed in nervous system\textsuperscript{10,22,23,26,28}, whereas the crab \textit{EsSoxB2-1} exhibits gonad-specific expression pattern as revealed by RT-PCR (Fig. 3). Given that insect \textit{Dichaete} also contains no intron\textsuperscript{14}, we constructed a phylogenetic tree using full length sequence of SOX proteins, in order to investigate whether \textit{EsSOXB2-1} is an insect \textit{Dichaete} homolog. The tree showed that \textit{EsSOXB2-1} first clustered with the beetle \textit{T. castaneum} and the fruit fly \textit{D. melanogaster} SOX21b rather than \textit{Dichaete} (Fig. 1B), implicating that \textit{EsSOXB2-1} is closely related to insect SOX21 in sequence. Taken together, we concluded that \textit{EsSoxB2-1} is a novel \textit{SoxB2} homolog and most likely represents a specific \textit{SoxB2} form of crustacean in the evolution of Sox genes.

Spermiogenesis is a highly complicated differentiation process from spermatid to mature spermatozoa. The differentiation of spermatogenic cells appears to be regulated by many nucleus-resident proteins\textsuperscript{25}. In vertebrates, haploid spermatids undergo dramatic changes in morphology including reduction of the nuclear size, elongation of sperm tail and condensation of chromatin. Histones are progressively replaced by protamine to pack genomic DNA, thereby producing more compact chromatin\textsuperscript{29}, and along with the removal of cytoplasm\textsuperscript{30}. The transcription and translation of many key regulatory genes are switched off to silence all cellular process that are not relevant to fertilization\textsuperscript{31}. These sequential changes result in spermatozoa maturation that generates sperm-specific mobility and fertility. Contrary to most species sperm, crab sperm is typically aflagellated and

Figure 6. Knockdown effect of \textit{EsSoxB2-1} mediated by RNAi. Expression level of \textit{EsSoxB2-1} mRNA and \textit{EsSOXB2-1} protein was assayed by qPCR (A) and Western blot analysis (B), respectively. Three individuals were randomly collected in each group at 24 and 48 hours (hr) post injection of PBS, GFP-dsRNA and \textit{EsSoxB2-1}-dsRNA. The relative expression level was determined using the beta actin as an internal control.
immotile, containing a spherical acrosome surrounded by the uncondensed nucleus. The nucleus of a mature sperm keeps the similar size with spermatids. Spermatid differentiation is characterized by chromatin decondensation. During chromatin decondensation, most histones are reduction and only small amounts of histones H2B and H3 remain in the nucleus of mature spermatozoa in the blue swimming crab *Portunus pelagicus*, which could lead to the disruption of nucleosomal organization and consequently the decondensation of sperm chromatin26. However, the molecular mechanism for maturation of the uncondensed sperm nucleus remains unknown in the crab. Our immunocytochemical analysis data showed that *Es*SOXB2-1 protein was not detected in spermatogonia and spermatocyte, but exclusively localized in the nucleus of the crab spermatid and spermatozoa during spermiogenesis (Fig. 4), suggesting involvement of *Es*SOXB2-1 protein in the crab spermiogenesis. To test this hypothesis, we further performed in vivo RNAi of *Es*SoxB2-1. Abnormal transformation of the nucleus was observed in the spermiogenesis (Fig. 7), indicating that *Es*SoxB2-1 is required for maturation of sperm nucleus. Additionally, our RNAi data showed that the expression of histones H3 and H4 had no significant change after knockdown of *Es*SoxB2-1 expression (Supplementary Figure S3), implicating that the mRNA expression of histones H3 and H4 is not regulated by *Es*SoxB2-1. *Es*SoxB2-1 could function as a transcription factor through controlling other protein expression to mediate the movement of histones between nucleus and cytoplasm, although there is no direct interaction between the *Es*SoxB2-1 and the histones H3/H4. Further study is needed to identify more RNAi-responsive genes and examine their relationships in sperm nucleus maturation. Intriguingly, several potential SRY/Sox binding sites were identified in the promoter region of *Es*SoxB2-1 (Supplementary Figure S2), suggesting *Es*SoxB2-1 could be regulated by other Sox homologs. Testing this hypothesis would require identifying more Sox protein(s) that can interact specifically with *Es*SoxB2-1. It will be of interesting to further elucidate the regulatory mechanism of *Es*SoxB2-1 in spermiogenesis.

Previous studies have shown that the SRY nuclear localization signals (NLSs) are highly conserved during evolution among mammals, and the mutation of NLS can lead to inefficient transportation into the cell nucleus12. This means NLSs are essential for translocation of SRY from cytoplasm to nucleus. Like SRY, *Es*SoXB2-1 also has two NLSs at the C- and N-terminal in its HMG domain (Supplementary Figure 1). This may explain why *Es*SOXB2-1 protein display nucleus localization in spermiogenic germ cells during spermiogenesis. Given that some nucleus-resident proteins such as extracellular signal-regulated kinases (ERKs) translocated from the nucleus to the acrosomal tubule during acrosome reaction of the crab spermatozoa22,32, we extended our studies to trace the *Es*SOXB2-1 protein in acrosome reaction. Interestingly, immunocytochemical analysis showed that

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**Figure 7.** Ultrastructural observation of spermatozoa in PBS group (A) and *Es*SoxB2-1-dsRNA group (B–D) after one month of repetitive injection. Blue arrow heads point high dense electronic granules generated from the degradation of nucleus arms. The scale bar indicates 0.5 μm. ls, lamellar structure; ml, middle layer; n, nucleus; po, percutor organ; ra, radial arm.
EsSOXB2-1 protein retained in nuclear without translocation to any other site (Fig. 8). Therefore, we conclude that EsSOXB2-1 protein is not involved in acrosome reaction and will remain in the sperm nucleus till fertilization. In fertilization, conventionally, the sperm cell delivered only the paternal haploid genome to the oocyte, which contributed the maternal haploid genome and all the other components required for early zygotic development, such as yolk protein, ooplasm and organelles. However, recent accumulating evidence showed that spermatozoon RNA was present in the zygotic of rat, mouse, and human. Furthermore, the paternal Wnt4 and Foxg1 can be translated into protein in zygotes. All these data demonstrated that spermatozoon delivered not only paternal haploid genome to the oocyte but functional mRNA and protein as well when fertilization. These paternal components include the spermatozoon centriole, transcription factors, and signaling molecules, which are required for early embryonic development. Accordingly, the maintenance EsSOXB2-1 protein in the sperm nucleus at the end of acrosome reaction also implicated that EsSOXB2-1 could be delivered into fertilized eggs along with chromatins functioned as a paternal transcription factor in regulating early embryonic development.

Conclusion
This study represents the first report on identification and functional characterization of a Sox gene in decapod species. Different with known Sox B genes in other species, EsSoxB2-1 has unique gene structure and were found to be specifically expressed in the gonads. The EsSOXB2-1 protein is predominantly expressed in the testes and exclusively localized in the nucleus of spermatid and spermatozoa even at the end of acrosome reaction, suggesting that EsSOXB2-1 could be delivered into fertilized eggs as the paternal transcript factor in regulating early embryonic development. RNAi knockdown of EsSoxB2-1 leads to abnormal transformation of the nucleus during spermiogenesis, demonstrating a role for the EsSoxB2-1 in sperm nucleus maturation.

Materials and Methods
Animals and tissues.
The mitten crabs were collected from a local fisheries farm. Various tissues, including testis, ovary, heart, muscle, liver, gill and thoracic ganglion, were sampled and immediately frozen in liquid nitrogen and stored at −80 °C until used. Testes were also fixed in Bouin’s fixative (15% saturated picric acid, 5% formalin, and 1% glacial acetic acid) for histological observation. For in situ hybridization and immunohistochemical analysis, testes were fixed with 4% paraformaldehyde in phosphate buffered saline (PBS) solution overnight at 4 °C and stored in methanol at −20 °C after washing three times with PBS. The testis stages were classified as described by Zhang and Qiu.

Total RNA isolation.
Total RNA was isolated from the tissues using TRIzol reagent according to manufacturer’s instruction (Invitrogen, USA). The potential contamination of genomic DNA was excluded by treating with RNase-free DNase I (Promega, USA). The integrity of RNA was assayed by agarose gel electrophoresis and the quantity of RNA was measured by a Nanodrop 2000 spectrophotometer (Nanodrop Technologies, USA).

Degenerate RT-PCR.
A pair of degenerated primers of crab Sox gene, sense, (5’-AAGCGACCCCATGAA(C/T) GC(A/G)(C/T)TT(C/T)AT-3’) and anti-sense (5’-TC(T/C)ACGAGGTCGATA(C/T)TT(A/G)TA(A/G)T-3’), were designed according to the conserved HMG box of different Sox genes. A total volume of 25 μL PCR mixture
Es were incubated with rabbit anti-

St. Louis, MO, USA) bearing gold particles at 25 °C for 2 hours. Specimens were examined by a transmission

electron microscope (JEM-1400, JEOL, Japan).

Preparation of antibody of

SoxB2-1 protein was produced from rabbits as described previously. The antibody of

SoxB2-1 protein, obtained from BL21 strain after induced by

isopropyl-β-D-thiogalactoside (IPTG), was purified through affinity chromatography. The antibody of

SoxB2-1 protein overnight at 4 °C. For negative control, a normal rabbit serum

was used instead of the primary specific antibody. Sections were then incubated with goat anti-rabbit IgG (Sigma,

St. Louis, MO, USA) bearing gold particles at 25 °C for 2 hours. Specimens were examined by a transmission

electron microscope (JEM-1400, JEOL, Japan).

Full-length cDNA amplification and sequencing. The full-length cDNA of the crab Sox gene

was retrieved with 3′ and 5′ RACE method using the Marathon cDNA Amplification Kit (Clontech, USA).

The specific primers (5′-GGGCTGGAATGGGCGCTTGTA-3′ for 5′ end amplification and 5′-ATGCTG

GAGATGATGCTGATGGA-3′ for 3′ end amplification) were designed based on the cDNA sequence of the
degenerated RT-PCR product. The amplification parameters were 94 °C for 30 s; five cycles of 94 °C for 5 s, 58 °C

for 4 min; five cycles of 94 °C for 5 s, 70 °C for 4 min; five cycles of 94 °C for 5 s, 72 °C for 4 min; twenty-five cycles

of 94 °C for 5 s, 68 °C for 4 min. RACE products cloned and sequenced were conducted as described above.

Phylogenetic analysis. The deduced amino acid sequence of the crab Sox gene was aligned with those of

other metazoan Sox genes from GenBank database using the ClustalW (http://www.ebi.ac.uk/Tools/clustalw2/

index.html). A phylogenetic tree was constructed based on the multiple alignment with the neighbor-joining (NJ)

method using MEGA 6.06 package.

Cloning of the genomic sequence of EsSoxB2-1 and analysis of promoter activity. Genomic DNA

was isolated from the crab muscle using phenol extraction method, and the 5′-flanking region of EsSoxB2-1

was obtained with the genome walking method using the Universal Genome Walker 2.0 Kit (Clontech, USA).

The sequence of 5′-flanking region was analyzed using the promoter prediction software and the transcription factor

binding sites prediction software. Then varying lengths of the 5′ flanking sequences were amplified by PCR using

five forward primers F1-F5 and a universal reverse primer at 3′ end (Supplementary Figure S2). The amplicorns of the

PCR were cloned into the promoterless pGL3-Basic Vector (Promega, USA). Sequence integrity and ori-

entation of cloned inserts were confirmed by sequencing. Transfections to HEK 293T cells were performed using

Lipofectamine 2000 Reagent (Invitrogen, USA) when the cells reached about 90% confluence. Cells transfected

with the empty pGL3-Basic plasmid were used as a negative control. Cell lysates were harvested 48 hours after

transfection, and then promoter activity assay was assayed by measuring luminescence signal intensity of both the

firefly luciferase and sea pansy luciferase using Dual Luciferase Reporter Assay System (Promega, USA). The ratio

of firefly luciferase activities and sea pansy luciferase activities were analyzed by one-way ANOVA followed by

Tukey’s test using the statistics software SPSS 10.0.

Tissue distribution of EsSoxB2-1 mRNA. Tissue distribution of EsSoxB2-1 mRNA was examined with

RT-PCR method. Equal amounts (500 ng) of each total RNA from ovary, testis, thoracic ganglia, heart, liver, muscle and
gill were reverse transcribed into first-strand cDNA using M-MLV Reverse Transcriptase (Takara, Japan). Target gene

and the reference gene β-actin were amplified with gene-specific primers: Sox (5′-CTCCAGAAGAGGGGCTACA-3′),

Sox (5′-CGTACATCTTGGTCATGCTT-3′); β-actinF (5′-CGACGGGTAGGGTCACTAC-3′) and

β-actinR (5′-CGTACATCTTGGTCATGCTT-3′). The amplification parameters were 94 °C for 4 min; thirty cycles of 94 °C for

30 s, 54 °C for 30 s, and 72 °C for 1 min; 72 °C for 10 min; stored at 4°C.

In situ hybridization (ISH). DIG-labeled cRNA probes were generated from the clone of a 281 bp fragment

(position 633–899 bp) of EsSoxB2-1 cDNA using a DIG RNA Labeling Kit (SP6/T7) (Roche Diagnostics,

Germany). The fixed testes tissues were routinely embedded in paraffin and sectioned. In situ hybridization on the

tissue sections was performed as previously described.

Preparation of antibody of EsSOXB2-1 protein. The open reading frame (ORF) of crab EsSoxB2-1

gene was cloned into an expression vector pGEX-5X, and then the recombinant plasmid was trans-
formed into BL21 strain. Recombinant EsSOXB2-1 protein, obtained from BL21 strain after induced by

isopropyl-β-D-thiogalactoside (IPTG), was purified through affinity chromatography. The antibody of

EsSOXB2-1 protein was produced from rabbits as described previously.

Western blotting and immunocytochemical analysis. The specificity of the antibody and the tissue
distribution of EsSOXB2-1 protein were determined by Western blotting. Total proteins of testis, heart, muscle,
liver, thoracic ganglion, gill were extract, 30 μg of total protein of different tissues were used for SDS-PAGE, pro-
teins in polyacrylamide gel were electrotransferred onto a nitrocellulose membrane, the membrane was treated in

blocking solution and incubated with primary antibody overnig at 4°C followed by a secondary antibody for

2 hours at room temperature. A negative control was set by using the normal (preimmune) rabbit serum. After

rinsed in Tris-buffered saline, the membrane was colored in the reagent of DAB Horseradish Peroxidase Color

Development Kit (Boster, Wuhan, China). Immunocytochemistry analysis on testis sections was performed as

previously described by Qiu et al.

Immunogold electron microscopy. Mature testes were fixed using 4% paraformaldehyde and 0.5% glutar-
aldehyde in 0.1 M sodium phosphate and sodium phosphate dibasic buffer (pH 7.05) at 4°C over night, post-fixed

for one hour at 4°C in 1% osmium tetroxide, and then embedded in resin and ultrathin-sectioned. The sections

were incubated with rabbit anti-EsSOXB2-1 protein overnight at 4°C. For negative control, a normal rabbit serum

was used instead of the primary specific antibody. Sections were then incubated with goat anti-rabbit IgG (Sigma,

St. Louis, MO, USA) bearing gold particles at 25 °C for 2 hours. Specimens were examined by a transmission

electron microscope (JEM-1400, JEOL, Japan).
Knockdown of EsSoxB2-1 expression by RNAi. To synthesize dsRNA targeting EsSoxB2-1, a 617bp fragment was amplified using specific primers (EsSoxF4, 5′-tcccccagtgCCACCAAAAGAGGAAAG-3′ and EsSoxF4, 5′-ggactagTCGAGGAACGTAC-3′) and cloned into L4440 plasmid vector with two T7 promoters. Recombinant plasmids were transformed into HT115 (DE3). E. coli were used to produce sense and antisense RNA of EsSoxB2-1. Total RNAs were extracted using TRIZol reagent (Invitrogen, USA), and then heated at 70 °C for 10 min, cooled down slowly to room temperature for annealing. After purified using phenol/chloroform, the resultant EsSoxB2-1-dsRNA was assayed using agarose gel electrophoresis, and quantified using a Nanodrop2000c spectrophotometer (Thermo, USA). Meanwhile a green fluorescent protein (GFP) gene was also cloned for production of GFP-dsRNA as a control.

RNAi experiment was performed by in vivo injection of dsRNA. Individuals that finished reproductive molting were selected and randomly divided into three groups: EsSoxB2-1-dsRNA, GFP-dsRNA and PBS injection groups. Each individual was injected at the base of the fifth pleopods. The injection of EsSoxB2-1-dsRNA, GFP-dsRNA and PBS injection was assayed by real-time qPCR using a SYBR Premix Ex Real Time PCR kit (Takara, Japan). mRNA levels was assayed using real-time PCR and subjected to one-way analysis of variance (one-way ANOVA) using SPSS statistics software. EsSoXB2-1 protein level was assayed by Western blotting. The intensity of target band was quantified using Quantity One software (Bio-Rad, USA) and the significant difference was analysed using SPSS statistics software.

Induction of acrosome reaction. The spermatzoa were collected from spermatophores in the seminal vesicles. To trace the EsSoxB2-1 protein in spermatzoa during acrosome reaction, acrosome reaction of the crab spermatzoa was induced by 0.1% CaCl2. Samples were collected and fixed using 4% paraformaldehyde in a PBS (pH 7.4) overnight at 4 °C and stored in methanol at −20 °C after washing three times with PBS. The histological observation and immunocytochemical analysis were conducted as above.

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Author Contributions
G.-F. Q. conceived and designed the experiments, analyzed the data and finished the final version of the manuscript. Z.-Q. L., X.-H. J., H. Y. Q. and L.-W. X. conducted the experiments, gathered the data and drafted the manuscript.

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