Introduction

Interleukin-7 (IL7) was first characterized in human as a growth factor of B lineage cells [1], while now it is a well-known multifunctional cytokine. IL7 plays an active role in the development, survival, maintaining and restoring homeostasis of mature T lymphocytes [2,3]. It is also a key regulator of the commitment, survival, proliferation and maturation of B cells during development [4]. Furthermore, IL7 can improve the antiviral function and expansion of natural killer (NK) cells [5,6] and regulate the development and differentiation of dendritic cells [7]. IL7 is produced by stromal cells in bone marrow and thymus [8,9] as well as other types of cell, such as keratinocytes [10], hepatocytes [11], and epithelial cells [12]. Besides its pleiotropic role in the immune system, IL7 has been reported as a regulator of the development of central nervous system [13] and myogenesis and skeletal muscle cell migration [14].

Tree shrews (Tupaia belangeri) are squirrel-like animals inhabiting in the tropical shrubs or forests of South and Southeast Asia [15], as well as South China [16]. It has the highest brain-to-body mass ratio of known mammals. Because tree shrews share some characteristics of primates and insectivores, the exact taxonomic position of tree shrew has been on debate [17–20]. The viewpoint that tree shrew has a close affinity with primates has been recently supported by genome sequencing of a Chinese tree shrew and comparison with 14 other species [20]. Due to these unique characteristics of experimental animals, such as small body size, short reproductive cycle and life span, and low-cost of maintenance, tree shrew has been proposed to be an alternative experimental animal to primates in biomedical research [16]. Indeed, there are some spontaneous diseases, e.g. diabetes and tumor, in captured tree shrews [21,22]. So far, tree shrew has been reported to be susceptible to infection with a wide range of human pathogenic viruses [23], including HBV [24–26], HCV [27], and HSV [28]. However, there are still many obstacles, especially low efficiency of infection and unknown mechanism, which disabled our attempts to establish a repeatable and stable tree shrew model for these human viruses. To collect more basic knowledge about the immune system and important genes that are related to pathogen infection and surveillance in tree shrew will undoubtedly pave the way to fulfill our ambitious task.
In this study, *IL7* and its mRNA transcripts were characterized in Chinese tree shrew. We analyzed their expression pattern in eight tissues of adult Chinese tree shrews and evaluated expression levels in tree shrew primary renal cells in response to poly(I:C) of different lengths. In addition, subcellular localization of overexpressed *IL7* isoforms was also investigated. Our results provide valuable information on understanding the key regulator *IL7* in Chinese tree shrew.

**Materials and Methods**

**Experimental Animals and Ethics Statement**

Chinese tree shrews were introduced from the experimental animal core facility of the Kunming Institute of Zoology, Chinese Academy of Sciences. After lethally anesthetized by diethyl ether, we collected eight different tissues, including heart, liver, spleen, lung, kidney, intestine, skeletal muscle and brain. Tissue samples were quickly dissected, immediately frozen in liquid nitrogen and were stored at −80°C. All efforts were made to minimize the suffering of animals.

The study protocol was reviewed and approved by the Institutional Animal Care and Use Committee of Kunming Institute of Zoology, Chinese Academy of Sciences.

**Total RNA Extraction and Reverse-Transcription (RT)**

Total RNA was extracted from eight tissues and primary renal cells of Chinese tree shrews using RNAsimple Total RNA Kit (TIANGEN, Beijing) according to the manufacturer’s instruction. The A260/A280 ratio of total RNA was measured on a photometer (Eppendorf, Germany) and only these samples with a value of 1.8–2.0 were used for subsequent reverse transcription. We also evaluated the quality and integrity of RNA samples based on the 28S and 18S rRNA bands on a 1% agarose gel. Around 2 μg total RNA was used to synthesize cDNA by using oligo-dT18 primer and M-MLV reverse transcriptase (Promega, USA).

**tl7 Transcripts Cloning**

Based on the predicted *IL7* sequences of tree shrew retrieved from the Ensembl (http://www.ensembl.org/index.html) and the genome information of Chinese tree shrew [20] which is available at the tree shrew database (http://www.treeshrewdb.org/), a pair of primers tIL7-F and tIL7-R (Fig. 4A and Table 1) was designed to amplify the entire *IL7* gene sequence. About 1 μl cDNA synthesized from total RNA (from spleen) or pooled RNA (from all eight tissues) was used as the template. The reaction was performed in a volume of 20 μl containing 0.4 μM of each primer, 200 μM dNTPs, 1U of LA Taq DNA polymerase (TaKaRa, Dalian, China) and 2 μl 10×Buffer. We used the following PCR conditions: one denaturation cycle at 95°C for 2 min, 35 cycles of 94°C for 30 s, 55°C for 30 s and 72°C for 30 s, followed by one cycle of 72°C for 5 min. Purified PCR products were cloned into the PMD 19-T simple vector (TaKaRa, Dalian). Five positive clones of each insert were directly sequenced.

**Reverse Transcription Quantitative Real-Time PCR (RT-qPCR)**

In order to investigate mRNA expression profile of *IL7* and its alternative splicing transcripts in tissues and cells, transcript-specific primer pairs were designed (Table 1) and RT-qPCR was performed using SYBR green Premix Ex Taq II (TaKaRa, Dalian) on an MyIQ2 Two-Color Real-Time PCR Detection system (Bio-Rad, USA). In brief, a volume of 20 μl containing 0.4 μM of each forward and reverse primer, 1 μl of cDNA product, and 10 μl of 2×SYBR green Premix Ex Taq II were used for the RT-qPCR reaction. The tree shrew housekeeping gene β-actin was used as the reference gene for normalization. The cycling condition consisted of an initial denaturation cycle for 5 min at 95°C, 35 cycles of 30 s at 94°C, 40 s at 55°C, and a final extension step at 72°C for 15 s. In order to verify no non-specific amplification, following the completion of RT-qPCR, melting curve analysis was performed. The melting protocol consisted of heating from 55 to 95°C at a rate of 0.5°C per step, and each step was held for 1 s for data acquisition. Standard curves were generated using 10⁻⁷–10⁻¹⁰ dilution series of PCR product for each of the *IL7* transcripts and β-actin gene.

**Plasmids Construction**

The CDS regions of four *tl7* transcripts (*tl7c, tl7-sv1, tl7-sv2 and tl7-sv3*) were amplified by two primer pairs to introduce restriction endonuclease sites (Xho I and BamH I) and to cover signal peptide region. PCR products were cloned into pEGFP-N2 (Clontech, USA; Primer pair: tl7F1 and tl7R2c), respectively. Another primer pair, tl7F3c and tl7R2c (Table 1), was designed to amplify the *tl7* transcripts without signal peptide, and PCR fragments were inserted into pEGFP-N2. The CDS regions of human *IL7* gene (*hIL7*) with (Primer pair: hILF and hILR2) and without (Primer pair: hILF2 and hILR2) signal peptide were also cloned into pEGFP-N2 (Table 1). All constructs were verified by sequencing.

**Cell Transfection and Immunofluorescence**

HeLa cell was bought from the Kunming Cell Bank, Kunming Institute of Zoology, which was initially introduced from ATCC. Cells were cultured in (Invitrogen, USA) at 37°C in 5% CO2. In brief, cells (1x10⁴ per well) were seeded in 12-well plate with coverslips and grown to 50% confluence. For each well, a total volume of 50 μl mixture containing 1 μg EGFP-tagged plasmid DNA and 2.5 μl FuGENE HD Transfection Reagent (Roche, USA) was incubated at room temperature for 20 min. Meanwhile, culture medium was removed and washed once with the OPTI-MEM medium (Invitrogen, USA). DNA/FuGENE HD complex was added to each well, together with an additional 430 μl Opti-MEM. After an incubation for 6 h, 1 ml of growth medium was added to each well. 48 h after transfection, cells were fixed with 4% paraformaldehyde for 10 min. Nuclei were stained with DAPI (Roche, USA). Subcellular localization of *tl7-EGFP* were visualized by using an Olympus Fluoview 1000 confocal microscope (Olympus, Meadow, NY, USA).

**Isolation and Culture of Tree Shrew Primary Renal Cells and Poly(I:C) Transfection**

Primary renal cells were established from 3 Chinese tree shrews with age range from 1 to 4 months. Briefly, tree shrew was sacrificed and a pair of renal was dissected. The intact renal was
minced into small pieces (about 1 mm3) in cold PBS, and the pieces were transferred into a 50 mL sterile plastic tube containing a 1 mg/mL DNAse (Sigma, USA) and 5 mg/mL collagenase Type IV (Invitrogen, USA) solution for 30 min in 37 °C water bath. After digestion, the solution was filtered through a 200-mesh sieve to remove tissue pieces. The primary renal cells were transferred into a 50 mL sterile plastic tube containing 105 cells/well) were transfected with Lipofectamine 2000 (Invitrogen, USA) with unpaired Student's t-test. Statistical analysis was performed using GraphPad software (GraphPad Software, La Jolla, CA, USA) with unpaired Student's t-test.

**Phylogenetic Analysis**

To infer the phylogenetic position of Chinese tree shrew based on the IL7 gene sequences, we retrieved IL7 mRNA sequences of 16 species from GenBank and/or Ensembl (Table S1). Both the coding DNA sequences (CDS) and amino acid sequences were used for phylogenetic analyses. The puffer fish fugu (Takifugu rubripes) was used as the outgroup to root the phylogenetic tree. Trees were reconstructed using the neighbor-joining [NJ] method, maximum likelihood (ML), and minimum evolution (ME) by MEGA5.0 [29]. Since protein sequences used for phylogenetic analysis are shorter than 200 amino acid residues, we chose the Kimura 2-parameter and Poisson as the models for nucleotide sequences and amino acid sequences, respectively. Accuracies and statistical tests of phylogenetic trees were measured by bootstrap method with 1000 replications. MrBayes 3.1.2 [30,31], which implements a Poisson model with Markov chain Monte Carlo method, was also used to obtain a phylogenetic tree. The run started with one cold chain and three heated chains for 2 million generations and every 100 sample was retained to get the final consensus tree.

**Statistical Analysis**

For measurement of expression pattern of tIL7 mRNA and its transcripts in primary renal cells with and without poly(I:C) stimulation, each assay was independently performed three times to validate the consistency of results. Data were presented as mean ± SD of three independent tests. Statistical analysis was performed using GraphPad software (GraphPad Software, La Jolla, CA, USA) with unpaired Student’s t-test.

**Results**

**Tree Shrew IL7 Cdna Sequence and Its Amino-Acid Sequence**

According to the predicted sequence information of tree shrew’s IL7 gene in Ensembl and the Chinese tree shrew genome sequence generated by our own [20], we inferred that the IL7 gene is consisted of 6 exons. Our sequencing data showed that the full-length of tIL7 transcript (tIL7c) is 1817 bp, with a 634 bp 5'-UTR, and a 647 bp 3'-UTR (including a poly-A tail) (Fig. 1). A potential polyadenylation signal AATAAA was located at 18 bp upstream of the poly-A sequence. The open reading frame (ORF) consists of 534 nucleotides and encoded a putative polypeptide of 177 amino acid residues (Fig. 1). In the deduced gene product of tIL7, there is a signal peptide with 25 amino acid residues in the N-terminal (Fig. 1). Three potential N-glycosylation sites Asn56-Ala-Ser

**Table 1. Primers for cloning and quantification of IL7 transcripts in Chinese tree shrew.**

| Primer    | Sequence (5'-3') | Application          |
|-----------|-----------------|----------------------|
| tIL-7F    | GCCGTGGACATATTAGCAAC | PCR for cloning tIL7 |
| tIL-7R    | ATCAAAATAGCTGACGGTCAG | PCR for cloning tIL7 |
| tIL7 F674 | CTCCTCTGAGTCTTGCTCTTG | 3’ RACE          |
| tIL7 GSP-R1 | CCGTTCCTACGAGGGTGTCGGAG | 5’ RACE          |
| tIL7 F1239 | CAGTGGTGGAGGAGGAGG | 3’ RACE nested PCR |
| tIL7 GSP-R2 | CAGGGGGGGGGCGCAACACAC | 5’ RACE nested PCR |
| tIL7 SSP-F3 | TGCATTGGAAGTTAAGTTCCTA | qRT-PCR for tIL7-sv1, tIL7-sv3 |
| tIL7 SSP-F4 | AAAATTCATGTGATGATAATAA | qRT-PCR for tIL7c |
| tIL7 SSP-F7 | CATGTGTGAAATAAGGAGTTT | qRT-PCR for tIL7-sv2 |
| tIL7 SSP-R1 | TTGTCTCCTCCTGCTGGTG | qRT-PCR for tIL7-sv2, tIL7-sv3 |
| tIL7 SSP-R5 | TCTTGGTGGAGGACTTTATG | qRT-PCR for tIL7-sv1, tIL7c |
| tβ-actin F | ATTTTGAGATGACGGCACCC | qRT-PCR for tβ-actin |
| tβ-actin R | AGGTAAACCGTCGCTCCTC | qRT-PCR for tβ-actin |
| tIL7F1e   | CCCTGCAGGACACGTCGGTCTTTC | PCR for plasmids construction |
| tIL7F2e   | CCCTGGAGATGATGATGATGGTACAGGTT | PCR for plasmids construction |
| tIL7R2e   | CGCGTACCGGTGTTTACGACCTCTC | PCR for plasmids construction |
| hIL7F     | CCCTGGAGATGATGATGATGGTACAGGTT | PCR for plasmids construction |
| hIL7F2    | CGCGTACCGGTGTTTACGACCTCTC | PCR for plasmids construction |
| hIL7R2    | CGCGTACCGGTGTTTACGACCTCTC | PCR for plasmids construction |

*Restriction endonuclease sites introduced by PCR are underlined. RT-qPCR, quantitative real-time PCR. doi:10.1371/journal.pone.0099859.t001
Figure 1. Nucleotide and deduced amino acid sequence of the IL7 gene in Chinese tree shrew. The six exons were marked by arrows and alternative splicing fragment of transcript tIL7-sv6 in the 5'-UTR was shaded. Potential polyadenylation signal AATAAA was marked with a box. Three predicted N-glycosylation sites were marked with dots below the respective amino acid. Three single nucleotide polymorphisms were underlined in this gene and were marked by "=".

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Evolutionary Analysis of the tIL7 Gene

The Chinese tree shrew IL7c transcript had a considerably high nucleotide identity with human IL7 gene (80.2%) (Table S2). In order to evaluate the evolutionary conservation of IL7 protein, sequences of multiple species were aligned together. The most conserved region was the signal peptide, in which 24 out of 25 amino acids were identical among the analyzed species except for chicken (*Gallus gallus*) (Fig. 2). In general, IL7 appeared to be a protein only highly conserved in lineage-specific species, such as primates. However, some amino acid residues were highly conserved (e.g. six cysteine residues) in all analyzed mammals (Fig. 2). These six cysteine residues in human IL7 protein can form 3 disulfides and have been reported to be important for the stability of protein three dimensional structure [32].

Identification of tIL7 Transcripts

To identify potential mRNA transcripts in Chinese tree shrew tissues, we performed sequencing for 230 cDNA clones from total RNA isolated from spleen tissue and pooled total RNA from eight tissues. Besides the above canonical transcript, a total of eleven different transcripts of the tIL7 gene were recognized in Chinese tree shrew tissues (Fig. S1), suggesting that the result was robust.
Expression Pattern of tIL7 mRNA and its Transcripts

To characterize different tIL7 transcripts, we first quantified mRNA expression profiles of tIL7c, tIL7-sv1, tIL7-sv2, and tIL7-sv3 in eight different tissues from 20 adult Chinese tree shrews. These transcripts were chosen because of their relatively high abundance in above-mentioned cloning analysis. The overall expression profiles of transcripts tIL7c, tIL7-sv1, and tIL7-sv2 were roughly same, whereas tIL7-sv3 had a remarkable difference (Fig. 5D). Transcripts tIL7c, tIL7-sv1 and tIL7-sv2 were mainly detected in tissues related to the immune system such as intestine and spleen, as well as lung (Fig. 5A, 5B and 5C), where alveolar macrophages were widely distributed to protect the host from invading pathogens. Moderate expression levels of these transcripts were observed in heart, skeletal muscle, liver and kidney, whereas the brain tissue had the lowest expression level (Fig. 5). The mRNA expression of transcript tIL7-sv3 was distinguished from the other three transcripts: (1) it had a high expression level in heart and skeletal muscle rather than in tissues related to immune system; (2) brain tissue had a considerably high mRNA expression of tIL7-sv3 (Fig. 5D). Of all analyzed tissues, tIL7c was the main transcript that accounted for more than 50% of all generated mRNA (excluding the brain tissue), whereas tIL7-sv3 had an obviously prevalent expression in brain (Fig. 5E). This result is consistent with the abundance of transcripts revealed by the above cloning sequencing (Table S8).

We next assessed mRNA expression levels of tIL7c, tIL7-sv1, tIL7-sv2 and tIL7-sv3 in primary renal cells transfected with a viral dsRNA mimic, poly(I:C), of short (0.2–1 kbp) and long (1.5–8 kbp) lengths for 6, 12 and 24 h. We found that both short and long poly(I:C) obviously induced mRNA expression of tIL7-sv1 at 6, 12 and 24 h. However, stimulation with poly(I:C) caused a trough of the tIL7-sv1 mRNA expression at 12 h compared to 6 h and 24 h. Similar tendency was also observed for mRNA expression profile of tIL7c, tIL7-sv2, tIL7-sv3, and tIL7 receptor (tIL7R) (Fig. 5F and Fig. S2). Short poly(I:C) had a better induction effect on these transcripts than long poly(I:C) at 6 h and 12 h, but this induction effect was similar (excluding tIL7c) for short poly(I:C) and long poly(I:C) at 24 h post-transfection (Fig. 5F). This pattern might reflect different reactions and signaling pathways of short and long poly(I:C) stimulation.

It should be mentioned that different expression profiles of tIL7c and tIL7-sv1 transcripts were found between kidney tissue and primary renal cells. The mRNA expression level of tIL7c was higher than tIL7-sv1 in kidney tissue (Fig. 5A and 5B), but tIL7c was lower than tIL7-sv1 in primary renal cells (Fig. 5E and 5F). The exact reason for this discrepancy might be due to different types of cells in kidney tissue.

Cellular Localization of tIL7 and its Transcripts

IL7 functions as a cytokine when released in extracellular medium. Protein function is strongly influenced by subcellular localization, and immunofluorescence microscopy was employed...
to determine cellular localization of the immature and mature tIL7c. EGFP-tagged tIL7c and tIL7 vectors were transfected into HeLa cells and tree shrew primary renal cells. As shown in Figure 6 and Figure S4, tIL7c without a signal peptide (tIL7c-SP-) was mainly localized to cytoplasm and presented a dot distribution, which is consistent with the pattern of hIL7-SP-. The fluorescence distribution of tIL7c-SP- was higher than the one with the signal peptide (tIL7c-SP+), Fig. 6, possibly because of the release of mature tIL7 to culture medium under the guidance of signal peptide. The other three isoforms (tIL7-sv1, tIL7-sv2, and tIL7-sv3) without a signal peptide had a similar distribution pattern with tIL7c-SP-.

**Discussion**

IL7 is an important molecule in the immune system and regulates the development, differentiation, survival, and matura-

Figure 4. Schematic structure of *IL7* mRNA and its transcripts in Chinese tree shrew. (A) Eleven mRNA transcripts of *tIL7* gene. All transcripts were amplified by using primer pair tIL7F and tIL7R. Exons were indicated as boxes. Broken lines indicated alternative splicing of exons in tIL7 transcripts. (B) A 21-bp insertion between exons 3 and 4 in transcripts tIL7-sv9 and tIL7-sv10 would result in a truncated peptide in the C-terminal of predicted protein. Transcripts tIL7-sv6, tIL7-sv9 and tIL7-sv10 complied with the splicing rule and were GT-AG introns. doi:10.1371/journal.pone.0099859.g004
mouse, pig, rabbit, horse, sheep, and fish were well characterized [40–46]. However, the existence of an \( IL7 \) homologue in tree shrew has not been well determined so far. Because tree shrew harbored some characteristics sharing with both the ancestral and modern primates, as well as unique features as an experimental animal, it has long been proposed as an alternative animal model to primates in biomedical research [20,23,47]. However, lack of basic knowledge regarding the immune system genes of Chinese tree shrew has disabled our efforts to create a stable and successful animal model for infectious disease.

Figure 5. Expression patterns of \( tIL7 \) and its transcripts in eight different tissues from 20 adult Chinese tree shrews. Relative mRNA levels of \( tIL7c \) (A), \( tIL7-sv1 \) (B), \( tIL7-sv2 \) (C), \( tIL7-sv3 \) (D) were normalized to the amount of \( \beta\text{-actin} \) mRNA. (E) Overall expression profile of the four transcripts of \( tIL7 \). (F) mRNA expression of the \( tIL7c \) and its transcripts in primary renal cells transfected with 1 \( \mu g/mL \) short and long poly(I: C) at 6, 12 and 24h. NC – non-transfected cells, poly(I:C) L – long poly(I:C), poly (I:C) S – short poly(I:C). The graph shows the mean ± SD of three independent tests. doi:10.1371/journal.pone.0099859.g005

Figure 6. Subcellular localization of EGFP-tagged \( tIL7c \) and \( tIL7-sv \) isoforms in HeLa cells. HeLa cells were transfected with pEGFP-N2 empty vector and pEGFP-N2 vector with insert of \( tIL7c \) or each of the three \( tIL7c \) transcripts (\( tIL7-sv1 \), \( tIL7-sv2 \) and \( tIL7-sv3 \)) with (SP+) and without (SP-) the signal peptide. Immunofluorescence images were taken at 48 h after transfection. The scale marked in each section of the figure referred to 20 \( \mu m \). doi:10.1371/journal.pone.0099859.g006
In this study, we cloned the *IL7* gene and identified a total of eleven alternative splicing transcripts. The splicing region of transcripts *tIL7-sv2*, *tIL7-sv4*, and *tIL7-sv5* were similar to IL785 (hIL7 transcript lacking exon 5), IL783/4 (hIL7 transcript lacking exons 3 and 4) and IL783/4/5 (hIL7 transcript lacking exons 3, 4 and 5) in human, respectively. However, there are no homologues of transcripts *tIL7-sv1*, *tIL7-sv6*, *tIL7-sv7*, *tIL-sv8*, *tIL7-sv9* and *tIL7-sv10* in human, suggesting that the alternative splicing of the *IL7* gene might be different between Chinese tree shrew and human. The uniqueness of the *IL7* gene in Chinese tree shrew could be further demonstrated by the phylogenetic trees of available *IL7* gene sequences (Fig. 3), in which tree shrew showed a divergent relationship to primates.

Transcripts resulted from alternative splicing usually had tissue- and/or time-specific expression patterns and played important roles in certain tissues and/or developmental stage [40]. *tIL7*, *tIL7-sv1*, *tIL7-sv2*, and *tIL7-sv3* were highly expressed in the immune system and presented somewhat different tissue expression patterns, suggesting their active roles in immune-regulation rather than being nonfunctional. Human IL7 and its isoforms were reported to be regulators of central nervous system and impacted on neuronal tissue development and plasticity [13]. Moreover, IL7 and its isoforms could act as a myokine to affect myogenesis and migration [14]. The relatively high mRNA expression of *tIL7* in the heart and skeletal muscle may indicate their roles in these related systems. The distinctly high expression level of *tIL7-sv3* in brain relative to other tissues may imply a key role of this transcript in tree shrew’s central nervous system (Fig. 5).

Cellular localization of these *tIL7* isoforms showed no specificity of certain isoform (Fig. 6). It should be noted that mRNA levels of these *IL7* transcripts might not be fully correlated with protein expression levels in tissues or cells, but we do not have the necessary antibodies to recognize each *tIL7* isoform.

To characterize potential function of different *tIL7* transcripts, we made several attempts, including (1) determination of mRNA expression levels of *tIL7* transcripts in primary renal cells in response to stimulation by different drug (lipopolysaccharide [LPS], poly(I:C), phytohaemagglutinin [PHA], rotenone, vitamin K3, carbonyl cyanide m-chlorophenylhydrazone [CCCP]); (2) testing for the proliferation rate of tree shrew spleen cells and primary renal cells in the presence of culture supernatant of HEK293 cells transfected with tIL7, tIL7-sv1, tIL7-sv2, or tIL7-sv3. Unfortunately, we did not obtain useful information to answer the critical question regarding the potential function of different *tIL7* transcripts. With the exception of poly(I:C) stimulation (Fig. 5F), other drugs had no apparent stimulation effect on mRNA expression levels of *tIL7* transcripts. We obtained inconsistent results regarding mRNA expression levels of *tIL7* transcripts in response to LPS treatment in renal cells from different tree shrew individuals: there was a seemingly delay of induction effect on mRNA expression of *tIL7* transcripts compared with poly(I:C) stimulation in some cells, but other cells had no response to LPS treatment (data not shown). There was no obvious difference of the proliferation rate of tree shrew spleen cells and renal cells cultured in the supernatant of HEK293 cells transfected with each of the four transcripts (*tIL7*, *tIL7-sv1*, *tIL7-sv2*, and *tIL7-sv3*) in comparison to the supernatant of HEK293 cells transfected with empty vector (Fig. S3). One potential reason for these negative observations would indicate that our system might not be optimal for distinguishing the effect of *tIL7* isoforms. Several laboratories have documented the expression of *IL7* in primary and secondary lymphoid organs using *IL7* reporter mice [8,9,48–50]. It may be more proper to work on thymic mesenchymal or epithelial cells. More efforts should be carried out to further define the function of these *tIL7* isoforms.

Some lines of evidence showed an effect of poly(I:C) on the induction of *IL7*. Maternal exposure to poly(I:C) in C57BL/6J pregnant mice (gestational day 16) induced expression of *IL7* in fetal mouse brain [51]. Treatment with poly(I:C) in salivary gland epithelial cells caused a significant increase of the *IL7* gene expression and protein production [32]. In the Japanese pufferfish, expression of the *IL7* gene in head kidney cells increased significantly upon treatment with poly(I:C) after 4 h [44]. Concordantly, we demonstrated that poly(I:C) had an upregulation effect on mRNA expression levels of *tIL7c* and its transcripts, in particular for *tIL7-sv1* (Fig. 5F). Future studies will be performed to characterize the in vivo effect of poly(I:C) on the induction of different IL7 isoforms and the signaling pathway underlying this upregulation effect.

In summary, we characterized expression pattern of alternative splicing variants of the *IL7* gene in Chinese tree shrew. The identification of diverse *tIL7* transcripts in Chinese tree shrew offered more food for thought: why Chinese tree shrew owns such a variety of *IL7* transcripts? What is potential function of different *tIL7* transcripts? How *tIL7* splicing is regulated during infection? How the tIL7 protein and its isoforms are modified in vivo? Functional study should be performed to answer these questions and to further define the regulation of alternative splicing of *tIL7* and the exact biological role of these transcripts in Chinese tree shrews.

**Supporting Information**

Figure S1 ML tree (A), ME tree (B) of IL7 amino acid sequences, with 1000 bootstrap replications and complete deletion in Gaps/Missing data. The Bayesian tree (C) using a Poisson model with mcmc method.

**Figure S2** Quantitative real-time PCR analysis of the *tIL7R* gene in primary renal cells stimulated with poly(I:C) of different lengths. Real-time PCR was performed using primer pair *tIL7R* F (5'-GAAGATTATCCAGACAAAACT-3')/tIL7R R (5'-TGACAGGAGGCATGAG-3') and cDNA synthesized from primary renal cells transfected with 1 μg/mL short or long poly(I:C) at 6, 12 and 24 h. The tree shrew housekeeping gene β-actin was used as the reference gene for normalization. NC—non-transfected cells, poly(I:C) L—long poly(I:C), poly(I:C) S—short poly(I:C). The graph shows the mean ± SD of three independent tests.

**Figure S3** Proliferation of tree shrew spleen cells and renal cells in response to *tIL7* isoforms and hIL7. 293T cells were transfected with 10 μg each of the four transcripts (*tIL7*, *tIL7-sv1*, *tIL7-sv2*, and *tIL7-sv3*) and hIL7 or an empty vector (pcDNA3.1) in a 10 cm dish (2 × 10⁶ cells/dish). Cell culture medium without FBS was replaced at 24 h post-transfection. Cells were incubated at 37°C for another 24 h, and then cell culture medium was harvested and added to tree shrew spleen cells seeded at 2 × 10^5^ cells/well or renal cells seeded at 2 × 10^5^ cells/well in 96-well plates. Proliferation of tree shrew spleen and renal cells was determined by MITT assay at 48 h. Data are presented as the mean ± SD deviation of triplicate samples.

**Figure S4** Subcellular localization of EGFP-tagged *tIL7c* and *tIL7-sv* isoforms in tree shrew primary renal cells. Cells were transfected with pEGFP-N2 empty vector and pEGFP-N2 vector with insert of *tIL7c* or each of the three *tIL7c* transcripts (*tIL7-sv1*, *tIL7-sv2* and *tIL7-sv3*) with (SP+) and without (SP-) the signal
peptide. Immunofluorescence images were taken at 48 h after transfection. The scale marked in each section of the figure referred to 20 μm.

**Table S1** 17 species used in the present analyses.

**Table S2** Homology analysis of the IL7 gene in 17 mammalian species.

**Table S3** Percentage of clones with tIL7 and its transcripts in mRNA isolated from tree shrew tissues.

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**Author Contributions**

Conceived and designed the experiments: YGY. Performed the experiments: DL XH JY. Analyzed the data: DL XH JY. Contributed reagents/materials/analysis tools: LBL. Contributed to the writing of the manuscript: DL XH JY.

**Table S1** 17 species used in the present analyses.

**Table S2** Homology analysis of the IL7 gene in 17 mammalian species.

**Table S3** Percentage of clones with tIL7 and its transcripts in mRNA isolated from tree shrew tissues.
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