Expression of Cre Recombinase in Alveolar Epithelial Cells of the AQP2-Cre Transgenic Mini-Pigs

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Key Words
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Abstract

\textbf{Background/Aims:} Optimal use of Cre mediated recombination in conditional animal models depends on well characterized Cre driver lines. Unfortunately, some Cre driver lines exhibit unexpected expression patterns hindering their utility in Cre/loxP systems. Thus, systematic assessment of new Cre lines is essential for generating useful Cre driver lines for future studies. 

\textbf{Methods:} Here, we describe a Cre Transgenic (Tg) mini-pig line in which the expression of Cre is directed by a 3-kb 5' fragment of the kidney-specific aquaporin 2 (AQP2); however, the AQP2-Cre Tg mini-pig line exhibits expression of Cre in alveolar epithelial cells (AECs) instead of collecting duct cells. The specificity of the AQP2-Cre plasmid was validated in vitro, and indicating that the AQP2-Cre was specifically expressed in the transfected LLC-PK1 cells. Absolute quantitative real-time PCR (qRT-PCR) and inverse PCR were performed to determine the copy numbers and integration sites of the AQP2-Cre transgene. Relative qRT-PCR was performed to evaluate variation in Cre expression levels over time. \textbf{Results:} Our data indicated that this AQP2-Cre Tg mini-pig line exhibits stable expression of Cre recombinase over time and in subsequent generations, even though the AQP2-Cre transgene was segregated and reduced in subsequent generations. \textbf{Conclusion:} Combined with our previous studies of the activity of this Cre, we conclude that this Cre Tg mini-pig line will provide a reliable tool for generating lung-specific gene targeting mini-pig models, thereby allowing the investigation of gene functions in lung development and studying the molecular mechanisms of human lung disease.
Introduction

The Cre/LoxP system is the most widely used conditional gene targeting strategy for spatial and temporal deletion of genes. This strategy exhibits a number of advantages compared with conventional gene targeting methods [1, 2]. It can be used to mutate target sites at precise positions or in specific development stages. The conditional animals also avoid potential consequences due to complete gene inaction, such as developmental defects and systemic phenotypes. In addition, this system is simple and efficient, without the need of additional cofactors or sequence elements. Thus, the Cre/loxP recombination system is widely used in yeast [3], mammalian cells [4], and plants [5].

The utility of recombination in Cre/LoxP animal models is based on the availability of well-characterized Cre driver lines. By controlling the expression of Cre recombinase within a particular type of cell or at a specific development stage, it is possible to control gene deletion in corresponding cells or stages in animal models [6, 7]. Advances in mouse embryology and molecular genetics, have allowed for the generation of various kinds of Cre recombinase expressing mouse lines [8, 9]. These mice show great promise for studying tissue-specific gene deletion and for designing more accurate models of human diseases.

It is important to note that a simple sounding goal such as generating a transgenic Cre expressing mouse line with cell type specific expression is not particularly simple to achieve. Indeed, even after a line is generated some Cre lines display activity at inappropriate times or in an unintended cell type [10]. For example, the promoter of the insulin 2 (Ins2) gene is known to direct expression only within the beta-cell population [11]; however, the Ins2-Cre Tg mouse lines showed widespread Cre activity in the brain [12]. In addition, the efficiency of Cre mediated recombination might vary at different genomic locations (LoxP sites) and segregation of the Cre Tg in different generations may lead to expression variation [13, 14]. Moreover, many Cre Tg animals display unanticipated characteristics that are not always obvious in the founders [15]. Also, reduced Cre activity has been correlated with the number of generations [16, 17]. These unexpected or varied expression patterns are major limitations in generating conditional gene-targeted mouse models. Thus, an important aspect of generating Cre driver lines is testing the spatial and temporal expression of Cre as well as its efficiency.

To obtain kidney-specifically expressed Cre driver mini-pigs, we generated two Cre Tg mini-pig lines expressing Cre recombinase under the control of 3.0-kb and 8.0-kb fragment of the mini-pig AQP2 5’-flanking region respectively. AQP2 is a water channel that is associated with renal water excretion and is thought to be preferentially expressed in the kidney collecting duct of human, mouse, and rat [18, 19]. Previous reports have shown that Cre recombinase, when driven by mouse or human AQP2 5’-flanking region, can mediate gene targeting specifically in mouse collecting duct cells [20]. As expected, the Tg mini-pigs in which the Cre expression was directed by 8-kb 5’-flanking region of AQP2 exhibit expression of Cre only in kidney collecting duct cells [21]. To our surprise, the 3-kb AQP2 5’-flanking region directed Cre expression was specifically observed in alveolar epithelial cells (AECs) instead of collecting duct cells.

The AECs participate in many essential functions including synthesis, secretion, and recycling of surfactant; and the transport of ions and water [22, 23]. Although these AQP2-Cre Tg mini-pigs express Cre recombinase outside of the intended cell type, it offers an option for investigating gene functions in AECs. In present study, we characterized the expression of Cre to confirm the utility of this Cre Tg mini-pig line and further clarified the correlation between Cre expression and genomic transgene integration. Our results demonstrate that despite the unexpected expression domain of Cre in the lungs, the expression is stable and specific over time. Combined with our previous studies of Cre recombinase activity, we believe that this Cre Tg mini-pig line will provide a reliable tool for developing AECs specific gene targeting mini-pig models.
Materials and Methods

Ethics statement

All animal studies were conducted according to the experimental practices and standards approved by the Animal Welfare and Research Ethics Committee at Jilin University (Approval ID: 2011010).

Construction of the AQP2-Cre expression plasmid

A 3 kb mini-pig AQP2 5’ fragment was amplified from the mini-pig genome, introducing NheI and Scal sites at the ends using primers AQP2-F/R (All primers used throughout this study are listed in Table 1). The PCR reaction was performed according to standard protocols, with the following parameters: 94 °C for 3 min, followed by 30 cycles of 30 s at 95 °C, 30 s at 66 °C, and 3 min at 72 °C, with a final extension step at 72 °C for 5 min. The amplified products were subjected to electrophoresis on 1 % agarose gel and purified using Gel Band Purification Kit (Tiangen, Beijing, China). Then, the AQP2 5’ fragment was used as an AQP2-Cre vector and corresponding endogenous genomic DNA sequence around the integration site

| Name  | Primers         | Sequence (5’→3’)                  | Product Size (bp) |
|-------|-----------------|-----------------------------------|-------------------|
| AQP2-F| AQP2-F          | TCTAGAAGGCTACGCTGACGTCGCTTCCT    | 3,990             |
| AQP2-R| AQP2-R          | GAAGATCTGGCGGCCGCCTGCTGACTTCT    | 2,684             |
| AC-F1 | AC-F1           | GGGGCGTCCGACCACTAGTGAATAGAGT     | 1,200             |
| Cre-1 | Cre-1           | GATGTTCAACTGCGCTCGTACGTTAGCG     | 1,044             |
| Cre-2 | Cre-2           | CTAGTCCTTGCGGCGTCGCG            | 127               |
| GAPDH | GAPDH-F         | CACATGGGGTCTGGCTATTCCACCTT      | 117               |
| GAPDH-R| GAPDH-R        | TCTCCAATGCTGAGTACGCACTAGCA      | 132               |
| TFRF-C| TFRF-C          | TCGGACTGGCTGATATGTTAGCAAT        | 132               |
| Cre-F1| Cre-F1          | CAGATTGAGAAGTGAAGGCGAGAAGAATCT  | 1,044             |
| Cre-F2| Cre-F2          | TCATTTCAACAAAGAAGGTTCTGCTCT     | 2,684             |
| Cre-R1| Cre-R1          | ACGCTTTGAAAGAAGGCGAGAAGAATCT    | 1,200             |
| Cre-R2| Cre-R2          | GGACATTATCTGAGCCTGAGAAATCT     | 127               |
| HindIII (3’-end) | F1 | TGATTTCAACAAAGAAGGTTCTGCTCT | 2,684 |
| F2   | ACTGGTAAAGAAGGCGAGAAGAATCT | 1,200 |
| R1   | GCCGATAATATCTGAGCCTGAGAAATCT | 127 |
| F1   | GCCGATAATATCTGAGCCTGAGAAATCT | 127 |
| Ecoro (5’-end) | R1 | GCCGATAATATCTGAGCCTGAGAAATCT | 127 |
| F2   | GCCGATAATATCTGAGCCTGAGAAATCT | 127 |
| R2   | GCCGATAATATCTGAGCCTGAGAAATCT | 127 |
| Specific primers (SP) for the 4 integration site | No.1 | CGCGATTAAATATCTGAGCCTGAGAAATCT | 638 |
| No.2 | CGCGATTAAATATCTGAGCCTGAGAAATCT | 638 |
| No.3 | CGCGATTAAATATCTGAGCCTGAGAAATCT | 638 |
| No.4 | CGCGATTAAATATCTGAGCCTGAGAAATCT | 638 |

Porcine nuclear transfer was performed as described by Lai et al. [25]. To identify Tg mini-pigs, genomic DNA samples from tail tissues of the newborn mini-pigs were extracted for Southern blotting and PCR analysis. All genomic DNA samples were extracted using TIANamp Genomic DNA Kit (Tiangen) in this study. For Southern blotting analysis, 20 μg of genome DNA was digested with BamHI/EcoRI restriction enzymes. The BamHI cuts once in the 3kb AQP2 5’ element and the EcoRI cuts once in the expression vector pET28a(+)-Cre backbone, resulting in digesting the pig genome efficiently and forming a 3, 527 bp fragment (Fig. 1A). The hybridization probe used to detect the Cre gene (1, 044 bp) was synthesized by PCR using primers Cre-F1/R1 and labeled by PCR DIG Probe Synthesis Kit (Roche). The digested DNA was separated in a 0.8 %
agarose gel and transferred to a nylon membrane (Roche). Hybridization and chemiluminescence detection were performed following the procedures of DIG-High Prime DNA Labeling and Detection Kit II (Roche). The PCR analysis was performed using primers AC-F/R as described above. The PCR products were analyzed by electrophoresis on a 1% agarose gel and sequenced to confirm the integrity of the Cre coding sequence.

**RT-PCR analysis**

Total RNA was extracted from cells or tissues using the Trizol A kit (Tiangen). RNA from each sample (1 µg) was treated with DNase I (Fermentas) and then reverse-transcribed using the BioRT cDNA First Strand Synthesis Kit (Bioer, Hangzhou, China). The cDNA samples were examined using primers Cre-F2/R2 to detect Cre gene respectively. Pig GAPDH was used as an internal control and amplified with the primers GAPDH-F/R. The PCR was conducted using 30 cycles of denaturation at 94 °C for 30 s; annealing at 59 °C for Cre and 61 °C for GAPDH for 30 s; and extension at 72 °C for 15 s. The sizes of the PCR products were 127 bp for Cre-F2/R2 and 117 bp for GAPDH-F/R. The PCR products were subjected to electrophoresis on a 1.5% agarose gel.

**Western blotting analysis**

About 3 × 10⁶ cells were lysed in 80μl IP lysis buffer (1% Triton X-100, 150 mM NaCl, 0.25% sodium deoxycholate, 20 mM Tris-HCl, pH 7.5) to extract protein samples. For tissue samples, 5 mg homogenized tissue was disintegrated in 100 μl IP lysis buffer. The protein concentrations were measured using the BCA Protein Assay Kit (Beyotime, Haimen, China). After protein samples (25 µg) were resolved on 10% SDS-PAGE gels, western blotting was performed according to a standard protocol. The membranes were incubated with goat-anti Cre antibody (1:2000, Santa Cruz, Dallas, TX, USA) or mouse-anti GAPDH antibody (1:2000, Bioss, Beijing, China) at 4°C overnight. Then, the blots were incubated with corresponding HRP-conjugated antibodies HRP-conjugated goat anti-rabbit secondary antibodies (1:8,000, Beyotime), donkey anti-goat (1:8000, Beyotime) or HRP-conjugated goat anti-mouse IgG (1:8000, Beyotime). Bands were detected using an ECL kit (Beyotime) based on the manufacturer’s protocol.

**Immunohistochemistry analysis**

Paraffin sections were pretreated with citrate buffer (0.01 M, pH 6.0) in 95 °C for 10 min and blocked with normal goat serum. The sections were incubated with goat-anti Cre polyclonal antibody (1:800, Santa Cruz) at 4°C overnight and then incubated with biotin labeled rabbit anti-goat IgG antibody (1:1000, Bioss) for 20 min at room temperature. Sections were washed with PBS and incubated with avidin/HRP complex. Immunohistochemical staining was carried out using UltraSensitive™ SP IHC Kit (Maxim, Fuzhou, China). Diaminobenzidine (Maxim, Fuzhou, China) was used for the color reaction, and the sections were analyzed by microscopy (Olympus CellSens Dimension).

**Testing the AQP2-Cre vector in vitro**

Initial in vitro analysis of the AQP2-Cre plasmid was performed two rounds. The first round was performed following the plasmid construction. The AQP2-Cre plasmid was transiently transfected into LLC-PK1 cells to estimate the expression of Cre recombinase. The second round in vitro expression analysis was performed after confirming Cre expression patterns in Tg mini-pigs. In the second round, the plasmid was transiently transfected into PK15, Hela, A549, HepG-2, and LLC-PK1 cells to assess the specificity of the AQP2-Cre construct. PK15 (pig kidney epithelial cell lines), Hela (human cervical fibroblasts lines), and HepG-2 (human hepatocellular carcinoma HepG 2 cell lines) cell lines, obtained in our laboratory, were purchased from American Type Culture Collection (ATCC); LLC-PK1 (pig kidney proximal tubule cell lines) cell lines were obtained from ATCC; A549 (human alveolar basal epithelial cell lines) cell lines were generously provided by Professor Ziyi Li at Jilin University. Briefly, cells were maintained in DMEM (Fermentas, Ottawa, Canada) supplemented with 5% fetal bovine serum (FBS) (Life Technologies, Grand Island, NY, USA), 1% penicillin/streptomycin (Gibco, Grand Island, NY, USA), at 37 °C, 5% CO₂ in humidified air. For transient transfections, cells were seeded at 1.2 × 10⁵ cells/cm² in six-well plates (Nunc, Roskilde, Denmark). Six hours after seeding, the cells were transfected using the FugeneHD (Roche, Basel, Switzerland) according to the manufacturer’s instructions. A total of 3.5 µg of circular AQP2-Cre plasmid and 10 µl of FugeneHD were added to 86.5µl of DMEM and incubated at room temperature for 15 min before the mixture was added to the culture medium. The cells were respectively harvested at 48 and 72 hours in the first round and at 48
in the second round after the transfection. The expression of Cre recombinase was determined by RT-PCR and western blot analysis.

**Relative qRT-PCR analysis**

To examine variations in Cre expression over time and in subsequent generations, qRT-PCR was performed. Extraction and treatment of total RNA were described as above. The Cre expression was examined using primers Cre-F2/R2 and GAPDH was used as an internal control. The qRT-PCR was performed using SYBR Premix Ex Taq Kit (Bioer) and the Bio-Rad Real-Time PCR System (Bio-Rad, Hercules, CA, USA). Amplification was performed under the following conditions: one cycle at 94 °C for 2 min, then 40 cycles at 94 °C for 10 s, and 60 °C for 30 s. For each test sample, a standard qPCR was performed in six replicates. A melting curve analysis was also conducted following amplification to test for primer dimers and other non-specific products. The delta CT for both the Cre and GAPDH are means of six replicates.

**Analysis of copy number by absolute qRT-PCR**

Copy number analysis was performed as described by Ingham et al. [26]. Briefly, Cre was detected using Cre-F2/R2 primers, and the transferrin receptor (TFRC) gene was amplified as a reference with the primers TFRC-F/R. Two rounds of absolute qRT-PCR, based on the standard curves with different resolution, were performed. In the first round, the standards were prepared by mixing different amounts of AQP2-Cre plasmid with 50 ng wild-type mini-pig genomic DNA to achieve 1, 2, 4, 8, 16, 32, 64 and 128 copies of the Cre gene per haploid genome. To prepare a standard sample containing one Cre gene copy per haploid genome, the mass of AQP2-Cre plasmid required for mixing with the 50 ng wild-type genomic DNA was determined by:

\[ \text{ng} = \frac{a \times b \times 0.5}{2} \times 10^{b} \]

where 'a' represents the plasmid size in kb, and 'b' represents the mass of wild-type genomic DNA in ng. Then, the different masses (in ng) of AQP2-Cre plasmid were calculated and mixed with 50 ng of wild-type genomic DNA. The qRT-PCR assays were performed as described above. Amplification was performed under the following conditions: one cycle at 94 °C for 2 min, then 40 cycles at 94 °C for 10 s, and 60 °C for Cre and 61.5 °C for TFRC for 30 s. For each standard and test sample qPCR was performed in triplicate. A standard curve was constructed by plotting the \( \Delta C_{\Delta} \) (\( \Delta C_{\Delta} = C_{\Delta \text{Cre}} - C_{\Delta \text{TFRC}} \)) values against the log of the Cre gene copies from the corresponding standard templates.

In the second round, a standard curve with proportionate resolution was used according to the Cre copy number values detected in the first round of qRT-PCR. The standards preparation and the qRT-PCR assays were performed as described above.

**Integration site analysis by inverse PCR**

One microgram of genomic DNA was digested with restriction endonucleases at 37 °C overnight. The restriction digestions were performed using HindIII (for 3'-end) and EcoRV (5'-end). After restriction enzyme digestion, the genomic fragments were treated with T4 DNA polymerase followed by ligation using T4 DNA ligase. Then, 100 ng of circularized DNA was amplified and 2 μl of this primary PCR product was used as the template in the second round of PCR using nested primers (NH-F/R and NE-F/R). The PCR was conducted using 30 cycles of denaturation at 94 °C for 30 s; annealing at 57.5 °C for primary PCR and 63.5 °C for the second round PCR for 30 s; and extension at 72 °C for 4 min. PCR products were gel-extracted and sequenced. The sites of integration were determined by BLAST-searching the pig genome database on NCBI.

To verify the integration sites, the genomic DNA samples from all Tg mini-pigs were further analyzed by PCR using specific primers (SP). These specific primers were designed according to the 3'- or 5'-flanking sequence of AQP2-Cre (forward primers) and corresponding endogenous genomic DNA sequence around the integration site (reverse primers). PCR was performed with the following parameters: 94 °C for 3 min, followed by 30 cycles of 30 s at 95 °C, 30 s at 59 °C for SP1, 62 °C for SP2, 57 °C for SP3 and 59 °C for SP4, and 15 s at 72 °C, with a final extension step at 72 °C for 5 min. PCR products were gel-extracted and sequenced.

**Statistical analyses**

Statistical analysis was performed using SPSS 17.0. All results are shown as mean ± SD values. Relative gene expression was determined using the 2^(-ΔΔCT) method built into the thermocycler software. One-way ANOVA followed by a Student–Newman–Keuls test was used to assess the differences between groups. Values of P = <0.05 were considered statistically significant.
Results

AQP2-Cre plasmid construction and in vitro expression analysis

The AQP2-Cre plasmid was constructed as shown in Fig. 1A. After verified by sequencing, the AQP2-Cre plasmid was transiently transfected into LLC-PK1 cells to estimate the expression of Cre recombinase. As shown in Fig. 1B, the Cre recombinase can be detected at transfected LLC-PK1 cells, showing that a 3-kb fragment of the pig AQP2 5'-flanking region can efficiently direct the expression of Cre recombinase in LLC-PK1 cells.

Generation and identification of AQP2-Cre Tg mini-pigs

Cells from a single surviving colony of mini-pig transfected fibroblasts were selected as nuclear donors for pig somatic cell nuclear transfer (SCNT) and a total of 1427 reconstructed embryos were transferred to five recipient female pigs. Two recipients miscarried all embryos during pregnancy and the other three produced twelve male mini-pigs. Of these mini-pigs, two died during delivery (No.0665 and No.0681) and four died soon after birth (No.0671, No.0679, No.0685, and No.0687) leaving six founder piglets (Table 2). The piglets appeared to be normal in size, morphology, and behavior (Fig. 2A). The average birth weight of those 12 cloned mini-pigs was 0.95 kg.

Tail tissue from all 12 newborn cloned mini-pigs was collected for transgene identification and fibroblast isolation. The integration of Cre recombinase gene was confirmed by Southern blotting and PCR analysis, and the results revealed that all of the mini-pigs were positive for the transgene (Fig. 2B, 2C). Sequencing results showed that all of them had integrated the complete Cre CDS.

To generate F1 transgenic mini-pigs, five month old Tg mini-pig, No.0677, was mated with a female wild-type mini-pig. Five F1 mini-pigs were born from one litter (Fig. 2A). One female (No.2448) and two male (No.2513 and No.2517) mini-pigs were confirmed to be transgenic by PCR analysis (Fig. 2C). Here, the genotype of AQP2-Cre Tg mini-pig lines segregated in subsequent generations, which is a common observation in Tg animals.
Expression pattern of Cre recombinase in various organs of the F0 and F1 Tg mini-pigs

The Cre expression pattern in the Tg mini-pigs was analyzed by RT-PCR and western blotting. Tissues from the four founder Tg mini-pigs, No.0679, No.0681, No.0685 and No.0687, that died during delivery or shortly after birth, and from one of the male offspring (No.2513) were used for analysis. Total RNA and protein samples were extracted from the heart, liver, spleen, lung, kidney, skeletal muscle, blood vessel, and testes. Our data showed that of the tissues analyzed, Cre was only expressed in lung tissue of founder Tg mini-pigs (Fig. 3-left and right). The F1 generation Tg mini-pig, No.2513, also showed exclusively lung expression of Cre (Fig. 3-middle).

To determine which cells were specifically expressing Cre in the lung, we performed IHC analysis on various tissues of mini-pigs No.0679 and lung tissue of No.2513 (Fig. 4). Cre staining was observed in AECs of F0 and F1 Tg mini-pigs (Fig. 4D and 4I). Of note, Cre was located in the nucleus and cytoplasm of the epithelial cells (indicated by red arrows) (Fig. 4J).
Specificity analysis of AQP2-Cre plasmid in vitro

As the AQP2-Cre Tg mini-pigs exhibit unanticipated Cre expression in lung tissues, we subsequently performed the in vitro expression analysis of the AQP2-Cre plasmid. Five cell lines named PK15, Hela, A549, HepG-2 and LLC-PK1 were transiently transfected with AQP2-
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Fig. 5. Expression of AQP2-Cre in different cell lines. Five cell lines, PK15, Hela, A549, HepG-2, and LLC-PK1 were transiently transfected with the AQP2-Cre plasmid. Cre expression was evident in AQP2-Cre transfected LLC-PK1 cells both in RT-PCR and western blotting (WB) analysis. The expected sizes of PCR products were 127 bp for Cre-F2/R2 and 117 bp for GAPDH-F/R primer pairs. GAPDH was used as a positive control in each test; water served as negative control (Control) in RT-PCR analysis.

Analysis of Cre copy number in Tg mini-pigs
We next analyzed the copy number of the AQP2-Cre transgene in the mini-pigs. To estimate the copy number of the AQP2-Cre transgene more accurately, we performed two rounds of absolute qRT-PCR using two standard curves with different resolution respectively. In the first round, we confirmed that the Cre copy number values of founders ranged from 8.20 to 11.41. Therefore, a standard curve with 15 points where standards ranged from 1 to 15 copies was used in the second round. Genomic DNA samples, extracted from tail tissues from each Tg mini-pig and from the other tissues (as above) harvested from piglets No.0679, No.0685, and No.2513 were analyzed. As showing Table 3 and 4, the Cre copies of founders and various tissues of a single founder mini-pig were ranged from 8 to 11. These data revealed that the Cre copies varied in the different founder Tg mini-pigs and various tissues, but that these differences were not significant (P>0.05) (Fig. 6A, 6B), which is consistent with previous reports [27]. The analysis of the F1 Tg mini-pigs showed only 5 to 6 Cre copies present in the different Tg mini-pigs and in the various tissues of No.2513 (Table 3 and 4). The observed Cre copies were varied, but the difference between the lowest and highest values (5 and 6 copies, respectively) was not significant (P>0.05) (Fig. 6A and 6B).

Table 3. Cre copy number values in different newborn Tg mini-pigs. The standard curve was defined as: \[ \log_2 N = \Delta C_t + 2.561 (r^2 = 0.9882) \]

| Pigs     | Cre copies | Pigs | Cre copies |
|----------|------------|------|------------|
| No.0665  | 8.52±1.35  | No.2448 | 5.08±0.96  |
| No.0667  | 9.85±1.37  | No.2513 | 6.97±0.98  |
| No.0669  | 10.96±1.31 | No.2517 | 5.32±1.39  |
| No.0671  | 9.16±0.73  |       |            |
| No.0673  | 8.58±0.88  |       |            |
| No.0675  | 9.24±0.97  |       |            |
| No.0677  | 10.39±0.87 |       |            |
| No.0679  | 10.45±0.69 |       |            |
| No.0681  | 11.16±1.09 |       |            |
| No.0683  | 10.53±0.97 |       |            |
| No.0685  | 9.88±0.93  |       |            |
| No.0687  | 9.57±1.16  |       |            |

Table 4. Cre copy number values in various tissues. The standard curve was defined as: \[ \log_2 N = -0.9045\Delta C_t + 1.9571 (r^2 = 0.9809) \]

| Tissue         | No.0679 | No.0685 | No.2513 |
|----------------|--------|--------|--------|
| Heart          | 9.38±1.23 | 9.74±1.59 | 5.80±1.45 |
| Liver          | 10.43±1.13 | 8.89±1.46 | 6.08±0.76 |
| Spleen         | 8.94±1.27 | 10.17±2.26 | 6.21±0.72 |
| Lung           | 10.28±1.49 | 9.91±1.37 | 5.93±1.61 |
| Kidney         | 9.78±1.18 | 10.39±1.19 | 6.50±1.27 |
| Muscle         | 10.18±0.98 | 10.98±1.21 | 5.97±1.00 |
| Blood vessel   | 10.58±1.22 | 9.18±1.53 | 5.80±0.86 |
| Testes         | 8.75±1.20 | 10.53±1.49 | 7.07±0.80 |

Analysis of Cre expression changes with aging and in subsequent generations
To confirm whether this Tg mini-pig line will stably express Cre recombinase with aging, the founder Tg mini-pigs, No.0687 (two days old), No.0683 (one month old), No.0677 (five months old), No.0675 (ten months old), and No.0673 (fifteen months old) were used to obtain lung tissues. Firstly, we analyzed the genomic integration copy number of different mini-pigs by absolute qRT-PCR. Genomic DNA samples extracted from these lung tissues...
Table 5. Cre copy number values in lung tissue from Tg mini-pigs at different ages.

| Age (months) | Newborn | 1    | 5    | 10   | 15   |
|--------------|---------|------|------|------|------|
| Cre copies   |         |      |      |      |      |
| No.0685      | 10.88±0.43 | 10.46±0.99 | 10.06±1.37 | 9.46±0.97 | 9.09±1.62 |
| No.0685      | NS      |      |      |      |      |
| No.0685      | NS      |      |      |      |      |
| No.2513      | NS      |      |      |      |      |

The standard curve was defined as: log\(^2\)N = -0.9312ΔC\(t\) + 2.062 (\(r^2 = 0.9901\)).

were examined. The results indicated that the Cre copy number in different Tg mini-pigs varied, but the difference was not significant (Table 5) (P>0.05) (Fig. 7A). Based on the Cre copy number analysis results, we analyzed the relative Cre expression levels in these founder Tg mini-pigs by relative qRT-PCR analysis. Total RNA samples extracted from the lung tissue of the mini-pigs described above (2 days old – 15 months old; No.0687, No.0683, No.0677, No.0675, and No.0673) were used in Cre expression analysis by relative qRT-PCR. The data revealed that the Cre expression levels in the founder Tg mini-pigs varied over time, but not significantly (P=0.075>0.05) (Fig. 7C).

We also analyzed the relative Cre expression changes between a founder mini-pig (No.0677, at 5 months old) and its offspring (No.2513, newborn). Although the genomic Cre copies between tails from mini-pig No.0677 and his offspring showed a significant decrease after passage (P<0.05) (Fig. 7B), the Cre expression levels didn’t significantly changed after passage (P=0.381>0.05) (Fig. 7D). These findings suggested that this Tg mini-pig line can stably express Cre recombinase in founders and the first generation.

Analysis of AQP2-Cre integration sites in Tg mini-pigs

In this study, the Cre expression vector was introduced randomly into the host genome. The unexpected expression domain of the AQP2-Cre could be caused by positional effects of random insertion. Therefore, we performed inverse PCR to analyze the integration sites of the AQP2-Cre vector in Tg mini-pigs. After two round inverse PCR, a total of eight PCR products were sequenced, and four insertion sites were identified (Table 6). We observed that the AQP2-Cre plasmids integrated in both intergenic regions and within genes. Plasmid
vectors tended to integrate within introns if the integration sites were in genes. Then, genomic DNA samples from the 12 founders and 3 offspring Tg mini-pigs were assayed by PCR using specific primers. We determined that all four integration sites were present in the 12 founders and 2 male offspring (No.2513 and No.2517), while only three integration sites (ID 1, 2, 3) were detected in piglet No.2448 (Fig. 8).

**Discussion**

Pigs are similar to humans in organ size, structure, and physiology; thus developing conditional pig models will be an exciting biomedical feat [28]. Recent advances in pig SCNT have made it possible to utilize Cre/loxP strategy in developing pig models. Leuchs S et al. described a targeted pig line carrying an inducible TP53 R167H allele orthologous
to the human TP53 R175H mutation [29]. This line can be used to investigate Li Fraumeni syndrome and to decipher the oncogenic characteristics of p53 mutations found in human cancers. In addition, the generation and investigation of other Cre Tg pigs has recently been reported [24, 30, 31]. The development of diagnoses and therapies for human diseases requires convincing animal model data. Therefore, well characterized Cre Tg pig lines with stable and specific expression are urgently needed.

The aim of this study was the generation of a kidney specific Cre mini-pig line. Previous reports have demonstrated that AQP2 is expressed in the kidney collecting duct cells of mice [32], rats [33], and humans [18]. Skowronski and colleagues reported that AQP2 expression occurred in the inner medullary collecting ducts of female pig kidney [34]. Our previous study revealed that endogenous AQP2 was expressed in kidney tissues of male pigs [21]. We therefore selected mini-pig AQP2 5′-flanking region in hopes of creating Tg mini-pigs in which Cre recombinase was specifically expressed in kidney collecting duct cells.

How long mini-pig AQP2 5′-flanking fragment is sufficient to direct the kidney-specific expression of Cre in Tg mini-pigs? Previous studies suggested that transcriptional regulators (TRs), binding to cAMP-responsive element-binding proteins (CREB), retinoid X receptor (RXR), Homeobox (HOX) and GATA elements, are involved in cell-specific regulation of AQP2 gene expression. These TRs binding elements are present in two highly conserved regions (513 and 224 bp upstream from transcription start site) of human and mouse AQP2 5′-flanking [35]. Sequence conservation analysis suggested that about 1500 bp upstream from the transcription start site of AQP2 gene is highly conserved promoter region. In this study, a 3-kb mini-pig AQP2 5′-flanking region was used to include all species-specific transcriptional regulators binding elements (TRBEs) in pig AQP2 promoter region. In addition, Cre recombinase has been shown to mediate cell-specific gene inactivation in mouse collecting duct cells under the control of the 11-14 kb mouse or human AQP2 5′-flanking region [20]. For instance, Guan and colleagues developed an AQP2-Cre mouse strain, using 14 kb of the mouse AQP2 5′-flanking region, to specifically delete the peroxisome proliferator-activated receptor subtype gamma (PPARγ) in collecting duct cells [36]. Similarly, Zhang et al. used a Tg mouse strain for Cre recombinase driven by 11 kb of the mouse AQP2 5′-flanking region to remove the PPARγ exclusively from collecting duct cells [37]. Accordingly, we constructed two Cre Tg mini-pig lines by using 3.0-kb and 8.0-kb pig AQP2 5′-flanking region respectively.

To our surprise, these two Cre Tg mini-pig lines characterize with absolutely different Cre expression patterns. The Cre Tg mini-pigs with 8-kb AQP2 5′-flanking region display an expected expression pattern of Cre within kidney collecting duct [21], while the 3-kb AQP2 promoter region directed Cre expression was unanticipated present in AECs of Cre Tg mini-
pigs. Moreover, characterization of Cre expression revealed that this Cre Tg mini-pig line exhibits stable expression within the AECs in founders and the first generation.

The AQP2 5′-flanking region directed Cre expression in AECs has not previously been reported. We speculated that potential regulation pathway(s) might be present in the AECs involved in the unexpected expression of AQP2-Cre in Tg mini-pigs. We therefore performed second round in vitro expression analysis. In this regard, a human alveolar basal epithelial cell line, A549, and other four cell lines were used. As a result, only AQP2-Cre transfected LLC-PK1 cells expressing Cre recombinase, whereas the others did not. Analogously, previous studies revealed that a 3-kb human AQP2 promoter region can efficiently direct the expression of reporter gene in LLC-PK1 cells, but not in rat IMCD cells, UT-87, Hep G2, HT-29, or NIH/3T3 cells [38]. These in vitro studies including expression pattern analysis of endogenous AQP2 in mini-pigs indicated that intracellular pathway(s) of AECs cannot regulate the AQP2 or AQP2-Cre expression.

Unexpected or varied expression of Cre recombinase has often been observed in Cre Tg mouse lines. For example, in an effort to generate a retinal pigmented epithelium (RPE)-specific Cre line, Ueki et al. used a 3.0 kb human vitelliform macular dystrophy-2 (VMD2) promoter to direct Cre expression, but they detected unanticipated Cre activity in the Müller cells during retinal development [39]. The Thy1-Cre Tg mouse line was initially created to express Cre recombinase in the postnatal cortex and hippocampal neurons [40]. Heffner et al. went on to demonstrate that it was also expressed in many additional tissues such as the myocardium, vascular endothelium, lung alveoli and bronchiole cells, renal tubules, skeletal muscle, skin, hair follicles, testis, and bladder [10]. These ectopic or varied cellular expression of Cre recombinase in the mouse and pig, suggest that unexpected expression of Cre recombinase is common in generating Tg Cre lines.

However, it is important to note that the most published studies focus on Cre expression in the tissues of interest, while ectopic or varied expression of Cre in other tissues are rarely reported. This means that a Cre line with varied expression is generally unavailable to potential users since the data were unreported. For example, the promoter fragment of the rat insulin gene was previously used to direct expression of a reporter gene in endothelial cells [41]. However, Gustafsson et al. reported that ectopic expression was observed in the adult rat brain [42]. Critically, the ectopic or varied expression of Cre may lead to the misinterpretation of phenotypes and could have a major impact on the final conclusions drawn from experiments [43, 44]. Though the generated Cre Tg mini-pigs characterize with unexpected expression of Cre, the systematical investigation is considered to be essential for further studies.

In this study, all 12 founder Tg mini-pigs were generated from a single cell clone using pig SCNT. The Cre copy number and integration site analysis indicated that these Tg mini-pigs all had similar integration patterns. Previously, Kong et al. proposed that Tg animals generated using SCNT, when they are derived from a single donor colony, should have a single genotype [27]. Our observations support this proposition. Furthermore, as the Cre copies did not significantly decline over time in the different aged mini-pigs, these Tg mini-pigs offer an option for investigating Cre expression variation with aging. We found that the expression levels of Cre were similar at all ages evaluated.

Uneven transgene passage between generations is a common observation in Tg animals, and previous reports concluded that this may influence expression [45, 46]. We did note a significant decrease in the number of Cre copies transmitted between founder (parent) and offspring, while Cre expression was stable. Notably, one integration site, ID.4, was absent from the genome of piglet No.2488; however, the Cre expression was readily detectable. This observation indicates that exogenous may be easily lost during fertilization, but expression can be maintained.

Although the AQP2-Cre mini-pig line exhibits exclusive and stable expression of Cre recombinase within the AECs, several fundamental questions remain unanswered. For instance, advanced testing of the AQP2-Cre Tg mini-pig lines for recombination efficiency and specificity will be necessary. Previous reports showed that the Cre-mediated recombination
might vary between generations or genomic target locations [47-49]. Therefore, further studies demonstrating the efficiency of the Cre to promote floxed-specific recombination are required, which could be accomplished by mating the Cre Tg mini-pig with a "reporter" Tg mini-pig, notably the Tg mini-pigs containing pICE-STOP [30]. In addition, Tg animals may segregate the integration patterns when mated with non-Tg animals [50, 51]. Although heterozygous Cre expression mouse lines have been used to generate conditional gene targeting mouse models [52-54], the sustained analysis of Cre integration patterns and expression in subsequent generations may be required for further applications of the AQP2-Cre Tg mini-pigs. Furthermore, the identified integration sites were shown to be on distinct genome loci in this study. Notably, integration site ID3 was located within an intron of the transmembrane protein 163 (TMEM163) gene. In rodents, TMEM163, is expressed in a wide extension of the brain [55]. Previous reports showed that TMEM163 is predicted to form an integral membrane protein with six transmembrane helices and involved in cell growth and migration [56]. Whether the integrated AQP2-Cre plasmids influence TMEM163 expression needs to be further analyzed. All these factors must be taken into account when choosing a cre driver strain for conditional knockout experiments.

In this study, the AQP2-Cre Tg mini-pigs were generated by a random integration. The unexpected expression domain of Cre may be due to positional effects that can occur in Tg animals [57, 58]. Therefore, we initially analyzed the genes located 50 kb up- and downstream of the insertion sites (Table 6). However, we found no potential promoters or TRBEs present in this region that may influence the expression of AQP2-Cre.

In addition to the position effect, several other hypotheses about the unique expression of transgene are possible. Firstly, we cannot exclude some complex mechanisms, such as long-distance regulation or interaction between exogenous and locus regulatory regions, that could lead to varied expression of transgene [59, 60]. For example, mutations in a conserved non-coding region in intron 5 of the Lmbr1 locus, which is 1 Mb away from the sonic hedgehog (Shh) coding sequence, caused a complete loss of Shh expression in the limb buds [61]. Previous studies also showed that the transgene can interact with locus control regions and thereby disrupt normal interactions between local and long-distance regulatory regions within a tissue-specific domain [62]. Additionally, comparing the Cre expression patterns of 8-kb and 3-kb AQP2 5'-flanking region directed Cre driver lines, we suppose that the 3-kb AQP2 5'-flanking region may miss regulatory elements crucial in controlling the tissue-specific expression of Cre. It is also important to note that some regulation mechanisms, such as cAMP, NFATc, and NF-kappaB signal pathways, were found to be critical for AQP2 expression in vivo or in vitro [38, 63, 64]. Thus, a precise understanding of the regulation elements and molecular mechanisms of cell-specific expression of AQP2 awaits further studies. Moreover, DNA methylation plays an important role in altered expression of transgene [65, 66]. Future studies will be needed to determine the methylation status of the AQP2 fragment used in this Tg mini-pig line.

Here, the mechanisms causing unexpected expression of transgene are poorly understood [67, 68]. The exact reason for the unexpected expression domain of the AQP2-Cre is beyond the focus of our study. Indeed, we successfully generated a lung-specific Cre expression mini-pig tool and the current study highlights several potentially issues that must be considered when designing specific Cre driver strains. This AQP2-Cre Tg mini-pig line showed no apparent defect, and the expression pattern of Cre recombinase was exclusive and stable between generations. Importantly, the coding sequence of Cre used in this study has been shown to be effective in previous studies. Thus, we predict that this Cre mini-pig line will be a valuable tool for investigating gene function in AECs.

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