The Porcine Mitochondrial Transcription Factor a Gene: Molecular Characterization, Radiation Hybrid Mapping and Genetic Diversity among 12 Pig Breeds

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Abstract: Problem statement: Mitochondrial transcription factor A (TFAM) is a nucleus-encoded protein that is a key activator of mitochondrial transcription as well as a major participant in mitochondrial genome replication. Genomic characterization of the porcine TFAM gene is, therefore, necessary to determine its involvement in regulation of fat depots and meat quality traits in pigs.

Approach: Genomic DNA sequence was determined using a comparative in silico annotation approach. RT-PCR was used for analysis of alternative splicing. Genome location was determined using Radiation Hybrid (RH) mapping. Genetic marker was identified by sequencing and genotyped by the PCR-RFLP method with SacI. GENEPOP version 3.3 software was used for statistic analysis.

Results: We determined both full-length cDNA and genomic DNA sequences of the porcine TFAM gene. Gene expression analysis revealed an alternative 5' splice site, which excludes exon 4 of the pig gene. We assigned this gene to porcine chromosome 14 (SSC14). A G/A substitution was detected in intron 1 of porcine TFAM gene and genotyped on a total of 252 animals, including 165 from seven Chinese and 87 from five Western pig breeds. The Bayesian analysis via MCMC (Markov chain Monte Carlo) revealed that these two groups of pigs were well separated at this locus during the breed history; 95% of the posterior difference of TFAM allelic frequency between these two pig groups was greater than zero.

Conclusion/Recommendations: All these data provided basic genomic information needed for further functional characterization of the porcine TFAM gene. Because marked differences in fat and lean tissue deposition exist between Western and Chinese pig breeds, the G557A mutation in the TFAM gene deserves further evaluation to determine its phenotypic effect on fattening and carcass traits in commercial pig populations.

Key words: alternative splicing, annotation, diversity, mapping, pig, TFAM

INTRODUCTION

Mitochondrial transcription Factor A (TFAM; previously known as mtTFA) is an integral part of the basal mitochondrial transcription machinery in mammals. It is required for transcription of mammalian mitochondrial DNA (mtDNA) from the light-strand and heavy-strand promoters in the presence of mitochondrial transcription factors B1 (TFB1M), B2 (TFB2M) and mitochondrial RNA polymerase (POLRMT). Transcription from the light-strand produces a short RNA primer, which is required for mtDNA replication. TFAM is also involved in the regulation of the mtDNA copy number. For example, mice heterozygous for a targeted mutation of the Tfm gene exhibited reduced mtDNA copy number, while homozygous knockout embryos exhibited severe mtDNA loss and embryonic lethality, probably due to abolished transcription dependent priming of mtDNA replication. In mammals, the TFAM gene has already been isolated and characterized in human, mouse, rat, cattle and silvered leaf-monkey. The TFAM gene sequences for orangutan, chimpanzee and dog were also deposited in the GenBank database. Here we report annotation, alternative splicing identification, potential miRNA target site detection, mapping and genetic diversity of porcine TFAM gene.
MATERIALS AND METHODS

Molecular characterization of the porcine TFAM gene:

Comparative annotation: The comparative approach for annotation of the porcine TFAM gene was described previously [9]. Basically, this annotation process consists of three steps. In step 1, we used the well-annotated human mRNA sequence as reference for BLAST searches to obtain the pig orthologous Expressed Sequence Tag (EST) sequences from the public database. Some of these ESTs were then selected for assembly and used for BLAST searches against the pig EST database for further extension of the cDNA sequence. In step 2, this extended cDNA sequence was further used as a query to retrieve genomic DNA sequence of the porcine TFAM gene and their alignment revealed its genomic organization. In step 3, the genomic DNA was used as a template against the EST databases for an electronic rapid amplification of cDNA ends (e-RACE) to generate a full-length cDNA sequence for the porcine gene. Such a process led to a comprehensive annotation of the porcine TFAM gene at both cDNA and genomic DNA levels.

Detection of alternative splicing forms: The total RNA was extracted by homogenization of 0.5 g of adipose and muscle tissues using TRIzol reagent according to the manufacturer’s instructions (Gibco BRL, Gaithersburg, MD). The mRNA was then reverse-transcribed to generate cDNA (RT-Superscript, Gibco BRL, Gaithersburg, MD). One µl of each cDNA mixture was amplified using the primers: 5'-CCG TTC AGT TTT GCG TAT GTA-3' (upstream primer; covering the border between the exons 1 and 2) and 5'-GCT CAA CCT TCT ATT TCA ACA CT-3' (downstream primer; exon 7). PCR was performed for 30 cycles under the conditions of denaturation at 94°C for 45 sec, annealing at 57°C for 45 sec and extension at 72°C for 1 min using the MJ Research PTC-200 DNA Engine Thermal Cycler (MJ Research, Waltham, MA). PCR product was electrophoresed through a 1% agarose gel and sequenced on ABI310 genetic analyser. Primer specificity for the RH panel was verified using pure pig and hamster DNA in PCR amplifications. Primer pairs that amplified hamster DNA were omitted and redesigned.

RH panel typing: The RH panel source, marker typing procedure and RH map construction strategy were described previously [10]. In brief, DNA from a porcine/hamster RH panel was purchased from ResGen Invitrogen Corporation (Huntsville, AL, USA), which consisted of 94 cell lines, a porcine and a hamster DNA control. PCR reactions with a final volume of 10 µL were performed under the conditions as follows: 95°C for 10 min, 32 cycles of 94°C for 30 sec, 56, 57 or 61°C (Table 1) for 30 sec and 72°C for 30 sec, followed by a further 5 min extension at 72°C. PCR products were then examined by electrophoresis on a 1.5% agarose gel in 1X TBE buffer. Gels were stained with ethidium bromide and photographed. Each gel was scored twice in order to eliminate possible scoring errors. The bands were designated into three classes: Positive (1), negative (0) and faint (?) for each gel image.

RH map construction: The RH maps were constructed using RHMAP (3.0), a statistical package for multipoint RH mapping developed by Boehnke et al [11]. The RH2PT program was used to estimate the locus-specific retention probabilities and two-point LOD scores for linkage of the various marker pairs and to generate linkage groups with a LOD score of 4.0, 6.0 and 8.0. Framework maps were first established by RHMAPLIK at odds greater than 1000:1 (SAVMAX = 15 and PRTMAX = 5) with a stepwise locus ordering strategy. The best comprehensive map was constructed with information from both RHMINBRK and RHMAXLIK analysis using the same ordering strategy at an equal retention model (SAVMAX = PRTMAX = 5).

Genetic diversity of TFAM gene among 12 pig breeds:

Animals and DNA preparation: A total of 252 animals, including 165 from seven Chinese pig breeds and 87 from five Western pig breeds were used in this study.
Table 1: Primer sequences and PCR conditions for radiation hybrid mapping

<table>
<thead>
<tr>
<th>Gene symbol</th>
<th>Gene description</th>
<th>HSA10 (Mb)</th>
<th>Forward primer (5'→3')</th>
<th>Reverse primer (5'→3')</th>
<th>Size (bp)</th>
<th>Tm (°C)</th>
<th>Accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCDH15</td>
<td>Protocadherin 15</td>
<td>55.28</td>
<td>TTTTGGATGGCAAACTACTCG</td>
<td>GAGAGCGCAACAAAACACAAA</td>
<td>212</td>
<td>56</td>
<td>BP151124</td>
</tr>
<tr>
<td>ZWINT</td>
<td>ZW10 interactor</td>
<td>57.78</td>
<td>GTAGCTGACCAAAACCACCAA</td>
<td>AGCTTCCAGGTGGTTTATGATTGA</td>
<td>314</td>
<td>61</td>
<td>CF36573</td>
</tr>
<tr>
<td>TFAM</td>
<td>Mitochondrial transcription factor A</td>
<td>59.82</td>
<td>CTGCTGAAAGAATGAAAGTTCGTT</td>
<td>GCTCAACCTCTTTTCAACACT</td>
<td>150</td>
<td>61</td>
<td>AY923074</td>
</tr>
<tr>
<td>PHYHIPL</td>
<td>Phytanoyl-CoA hydroxylase interacting protein-like</td>
<td>60.61</td>
<td>AGAATTATCGACACTTCCAAATG</td>
<td>AAGCAGACTAGATCAGAGGCAATTA</td>
<td>362</td>
<td>57</td>
<td>BQ599786</td>
</tr>
<tr>
<td>CDC2</td>
<td>Cell division cycle 2 protein</td>
<td>62.21</td>
<td>ATGGCAGCTTATCATCCATATTTT</td>
<td>GTTGTCTACGATGGGAATCTCAAAACA</td>
<td>261</td>
<td>56</td>
<td>AJ600106</td>
</tr>
</tbody>
</table>

Tm: Annealing temperature

Genomic DNA of the Chinese pig breeds was extracted at Nanjing Agricultural University from blood samples of Jinhua (n = 19), Neijiang (n = 28), Xian pig (n = 24), Ming pig (n = 25), Wan (n = 20), Jiangquhai (n = 24), Ming pig (n = 24), and Taoyuan (n = 18). As for Western pigs, skin tissue was sampled from five different pig breeds: Swedish Landrace (n = 21), German Landrace (n = 19), Piétrain (n = 21), Yorkshire (n = 18) and Krskopolje pig (n = 8). These samples were collected from farms in Slovenia under the National Pig Improvement Program. Genomic DNA was prepared from skin tissue samples by standard phenol-chloroform-isooamyl alcohol (25:24:1) extraction.

**Marker development:** Primers from exon 1 and 2 were designed to amplify the sequence of entire intron 1: forward primer, 5'-GTC ATT GTT GGG GTT GTT CTC T-3' and reverse primer: 5'-GTA TGA AGT CAT AGG CTT CTT TGG-3'. A G557A substitution in intron 1 of the porcine TFAM gene (GenBank acc. no. AY923075) was detected by direct sequencing of PCR products on Chinese and Western DNA pools and then assayed on a total of 252 animals. The PCR reactions were performed under the same conditions as described above for RH panel genotyping. The PCR products were then digested with 2U of SacI (MBI Fermentas, Vilnius, Lithuania) at 37°C for 4 h and examined by electrophoresis on a 1.5% agarose gel with 1× TBE buffer. The gels were stained with ethidium bromide and photographed.

**Statistical analysis:** GENEPOP version 3.3 [12] software was used to calculate allele frequency, observed (H_o) and expected (H_e) heterozygosity, Wright’s F statistics at the TFAM locus in single pig populations and in pooled Chinese and Western pigs, respectively. Further, Bayesian Poisson modeling was used to reveal differentiation of TFAM gene frequency between Chinese and Western pigs by assuming that they are random samples from two distinct overall populations, respectively, using the following model:

\[ x_i \sim \text{Poisson}(\theta_i n_i), \quad i = 1, 2; \quad j = 1, 2, \ldots, n_i \]  

Where:

- \( x_{ij} \) and \( n_{ij} \) = The number of allele A and the total number of alleles (A and G) in population j from region i (i.e., \( i = 1 \) for Chinese pigs and \( i = 2 \) for western pigs)
- \( \theta_i \) = The Poisson mean for either region of pigs.
- Conjugate gamma prior distribution is used for the allelic frequency parameter: \( \theta_i \sim \text{Gamma}(\alpha, \beta) \)

During the MCMC (Markov chain Monte Carlo) sampling, we ran a chain of 100,000 updates after a burn-in of 1,000 updates, with chain values saved at every 10th update. Thus, a total of 10,000 posterior samples were obtained to infer the distribution of allelic frequency parameter.

**RESULTS**

**cDNA sequences compilation:** The BLAST searches using the human TFAM cDNA (NM_003201) as a reference sequence retrieved fourteen porcine ESTs from the “ESTs_others” database at NCBI that are orthologous to the human gene. Two ESTs (BW961355 + BP167625) were chosen for initial assembly, which was further extended by adding CV877499 to its 5’ end and DB806415 to its 3’ end. As such, these four ESTs formed a complete coding sequence of 1,798 bp for the porcine TFAM gene, including 353 bp of the 5’UTR region, 741 bp of the coding sequence and 704 bp of the 3’UTR region.

The coding sequence of 741 bp was translated into a protein sequence of 246 amino acids for pig TFAM, the length being the same as in human, chimpanzee, cattle and dog, but two amino acids longer than in rat, three amino acids longer than in mouse and 38 amino acids shorter than in orangutan. The amino acid sequence showed 91% identity with cattle [7], 88% with...
dog (NW_139850), 71% with human (M62810), chimpanzee (XM_521481) and orangutan (CR857400), 64% with rat (AJ312746), 62% with mouse (U57939), 42% with frog (U35728), 40% with chicken (AB059657), 22% with fruit fly (AB045318) and 17% with yeast (M73753), respectively.

Genomic organization: The cDNA sequence of 1,798 bp further retrieved a genomic DNA sequence of 171,235 bp (CU041308) from the GenBank database that contains the porcine TFAM gene. Alignment of both cDNA and genomic DNA sequence revealed that, like in other mammals, the pig gene has seven exons with six introns all completely sequenced. The size of each intron is: intron 1, 543 bp; intron 2, 2,052 bp; intron 3, 645 bp; intron 4, 4,587 bp; intron 5, 7,993 bp and intron 6, 485 bp, respectively. The e-RACE further extended the 3'UTR from 704 bp to 2,651 bp with support of more than 50 ESTs. Therefore, the TFAM gene spans a region of at least 20,050 bp from transcriptional start site to transcriptional stop site in the porcine genome (CU041308).

Alternative splicing and miRNA target site: RT-PCR reactions amplified two amplicons, the predominant one with 660 bp and another weaker amplicon with 558 bp. Sequencing analysis indicated that the latter amplicon does not contain a coding sequence of 102 bp in exon 4 of the porcine gene (Fig. 1). The alternative splicing follows a “GT” site as a splicing donor site, which results in the shortening of exon 4 from 150 bp to 48 bp. Therefore, the seven-exon structure remains preserved in the alternative splicing form of pig TFAM gene, but at the protein level it is 34 amino acids shorter compared to the long version.

The TargetScan search for conserved miRNA target sites revealed a highly conserved miR-27ab binding site in the 3'UTR region of the TFAM gene, which is conserved in nine species (Fig. 2). The sequence alignment revealed the conservation of the 3'UTR region in pig as well.

Chromosome assignment: Mapping of the porcine TFAM was performed by using a porcine/hamster RH panel, which consisted of 94 radiation hybrid cell lines plus pig and hamster DNA as controls. All five selected genes were successfully typed on the panel with a retention rate of 26.6%, 21.5%, 22.3%, 21.3% and 21.3% for CDC2, PHYHPL, TFAM, ZWINT and PCDH15, respectively. The RH2PT program analysis assigned these five genes to one linkage group with a LOD score greater than 5. As the porcine CDC2 gene has been previously placed on SSC14 by Nonneman and Rohrer[13], any member of this linkage group is accordingly assigned to the same pig chromosome, including the porcine TFAM gene (Fig. 3). These five genes spanned a total of 129.5 cR in distance.

Genetic diversity of TFAM gene among 12 pig breeds: A 792 bp fragment containing the first intron and a part of the first and the second exon of the porcine TFAM gene was amplified from DNA of 252 animals representing seven Chinese pig breeds and five Western pig breeds. The G557A substitution in the intron 1 (GenBank acc. no. AY923075) was identified using restriction enzyme SacI. The frequency of allele A was high (from 0.917-1.000) in the five Western pig breeds, whereas it was moderate (from 0.500-0.750) in Chinese pig breeds except Jinhua and Taoyuan (Table 2).
Table 2: Allele frequency, heterozygosity and Wright’s F statistic (F<sub>IS</sub>) of TFAM gene in Chinese and Western pig populations

<table>
<thead>
<tr>
<th>Genotypes</th>
<th>Allele frequency</th>
<th>Heterozygosity</th>
<th>F&lt;sub&gt;IS&lt;/sub&gt;</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>G</td>
<td>A</td>
<td>H&lt;sub&gt;O&lt;/sub&gt;</td>
</tr>
<tr>
<td>Pig breed</td>
<td>N</td>
<td>GG</td>
<td>GA</td>
</tr>
<tr>
<td>Chinese pig breeds</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neijiang</td>
<td>28</td>
<td>2</td>
<td>17</td>
</tr>
<tr>
<td>Xian pig</td>
<td>24</td>
<td>4</td>
<td>16</td>
</tr>
<tr>
<td>Ming pig</td>
<td>25</td>
<td>7</td>
<td>9</td>
</tr>
<tr>
<td>Wan pig</td>
<td>20</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>Jiangduai</td>
<td>31</td>
<td>8</td>
<td>11</td>
</tr>
<tr>
<td>Taoyuan</td>
<td>18</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Jinhua</td>
<td>19</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Chinese</td>
<td>165</td>
<td>22</td>
<td>63</td>
</tr>
<tr>
<td>European pig breeds</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Swedish</td>
<td>21</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Landrace</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Piétrain</td>
<td>21</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Yorkshire</td>
<td>18</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Krskopolje pig</td>
<td>8</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>German</td>
<td>19</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Landrace</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Western</td>
<td>87</td>
<td>0</td>
<td>4</td>
</tr>
</tbody>
</table>

1: W and C: F<sub>IS</sub> calculated as in Weir and Cokerham[18]; R and H: F<sub>IS</sub> calculated as in Robertson and Hill[19]; 2: H<sub>O</sub>: Observed heterozygosity, H<sub>E</sub>: Expected heterozygosity

**DISCUSSION**

In the present study, we developed a simple, but reliable procedure to characterize the genomic features of porcine TFAM gene. First, we used a comparative approach to retrieve cDNA sequence of the porcine gene from the EST database: four ESTs (BW961355, BP167625, CV877499 and DB806415) contributed to the formation of a full-length cDNA sequence. Second, this full-length cDNA sequence was then used to search for its genomic DNA sequence against the pig genome sequencing database, which revealed that the pig gene contains seven exons. Third, mRNA extracted from two tissues was used to examine whether this gene has alternative splicing forms. Two splicing forms were confirmed by sequencing. Last, bioinformatics search revealed a miR-27 target site in the 3'UTR that was conserved in ten species including pig. Interestingly, the role of miR-27 in adipogenesis has been reported just recently[14]. However, the experimental validation of the predicted miR-27ab target in the porcine TFAM 3'UTR region as well as in other species needs to be performed in the future.
RH mapping assigned the TFAM and four neighbor genes on SSC14. The gene order for PCDH15, ZWINT, TFAM, PHYHIP1 and CDC2 was conserved in both human and pig genomes, but the orientation of the gene cluster in both species needs to be investigated by further mapping efforts. Interestingly, this linkage group of five genes falls into a region where the QTL for backfat thickness has been reported by Rohrer and Keele[15,16]. Recently, Jiang and colleagues[7] have reported a significant association of TFAM promoter polymorphisms with marbling scores and subcutaneous fat depth in Wagyu × Limousin cattle F2 crosses. This may imply that the porcine TFAM gene may be a strong candidate gene for this backfat QTL on SSC14, indicating that the TFAM gene plays an important role in lipid metabolism, fat deposition in muscle and obesity in mammals in general. Observed (H0) and expected heterozygosity (H0) was not dramatically different in these pig populations except for Taoyuan pigs, indicating no significant departure from Hardy-Weinberg equilibrium. This observation is in agreement with the calculated FIS statistic at this locus, which was low in most breeds except in Taoyuan pigs. The FIS statistic measures the reduction in heterozygosity in an individual due to nonrandom mating within a population. Consistently, Hardy-Weinberg test indicates significant heterozygote deficit in Taoyuan pigs. This might be due to so-called Wallhund effect, which is the result of the presence of subpopulations in the samples representing this breed. However, given that only 18 Taoyuan pigs were sampled, the significant heterozygote deficit could be due to sampling error as well. In addition, we noted that both H0 and H0 were higher in Chinese breeds than in Western breeds, which is similar to some previous reports[8,17]. As heterozygosity is a measure of genetic diversity, this result indicates that Chinese breeds exhibited higher within-population variation than Western pigs. Bayesian estimation indicated a clear allele differentiation between Chinese and Western pigs at this locus. Differentiation of allele frequency of the TFAM gene could reflect the fact that genes involved in energy metabolism were affected by different selection strategies in both groups of pig breeds.

CONCLUSION

The present study represents the first comprehensive investigation of the pig TFAM gene for its genomics features, such as cDNA sequence, genomic organization, alternative splicing and miRNA target site. We placed this gene to Sus scrofa chromosome 14 and developed a single nucleotide polymorphism in its intron 1. Because marked differences in fat and lean tissue deposition exist between Western and Chinese pig breeds, the mutation G557A in the TFAM gene deserves further evaluation to determine its phenotypic effect on fattening and carcass traits in commercial pig populations.

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REFERENCES