Analysis of Genetic Variation in RYR 1 and H-FABP in PSS Pigs using SNP Genotyping

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Abstract: The present study investigated the genetic variations in the Ryanodine Receptor 1 (RYR1) and Heart Fatty Acid-Binding Protein (H-FABP) genes, responsible for Porcine Stress Syndrome (PSS) observed during auction in pigs. A total of 655 pigs were sampled during either transportation or auction period. A SNPShot assay, combining multiplex PCR with multiplex primer extension assay was used for targeted detection of all mutations in one reaction. The most frequent RYR1 genotype was the recessive nn, followed by the heterozygous Nn and the dominant NN. Investigation of the H locus of the 5' upstream region in H-FABP showed that the genotype frequencies of HH, Hh and hh were 0.82, 0.16 and 0.02, respectively. Furthermore, analysis of the D and A loci in the intron 2 region of H-FABP showed that the genotype frequencies of DD, Dd and dd were 0.03, 0.39 and 0.59 and those of AA, Aa and aa were 0.47, 0.51 and 0.02, respectively. Thus, a high incidence of PSS was associated with HH, DD, Dd, Aa and AA and a low rate with Hh, hh, DD and aa. In conclusion, a positive correlation was found between the PSS development and SNP genotypes of RYR1 but not H-FABP.

Key words: Auction market, heart fatty acid-binding protein, porcine stress syndrome, ryanodine receptor, SNPs, H-FABP

INTRODUCTION

During transport to the slaughter plant, nonambulatory pigs with no injuries cause severe economic loss for the swine industry. This has raised serious concerns about animal welfare and pre-slaughter management such as transportation, lairage condition, and marketing process in last decades (Gregory, 2008; Johnson et al., 2013). In Taiwan, market-weight pigs are sold alive in auction markets. It has been reported that the percentage of pigs rendered nonambulatory during transportation and auction process was about 1.15% and their mortality rate was 0.21-0.45% (Huang et al., 2011). Besides, environmental and management factors, the well-known disease related to pre-slaughter loss is Porcine Stress Syndrome (PSS) which is associated with a point mutation of Cytosine (C) to Thymine (T) at nucleotide 1843 in the gene RYR1 (Fuji, et al., 1991). RYR1 has a negative impact on pre-slaughter death and pork quality, however, homozygous and heterozygous pigs show higher lean yield and better feed efficiency (De Smet et al., 1996; Band et al., 2005; Leach et al., 1996; Murray and Johnson 1998). Besides RYR1, Heart Fatty Acid-Binding Protein gene (H-FABP) is considered to be important for carcass and meat quality, especially for intramuscular fat and eye muscle area (Gerbens et al., 1999, Kocwin et al., 2006; Lin et al., 2002; Zhang et al., 2001).

Genotyping of RYR1 and H-FABP for mutations has been successfully employed as a means of genetic selection in pig breeding programs (Robinson and Buhl, 2005). Previously, genotyping was performed either by sequencing or Polymerase Chain Reaction-Restriction Fragment Length Polymorphism (PCR-RFLP) (O’Brien et al., 1993; Zeng et al., 2005). However, these methods are either expensive or time consuming. But a kind of Single Nucleotide Polymorphism (SNP) genotyping technique has been discussed in animal genetics (Garvin et al., 2010; Gupta et al., 2008;
Ragoussis, 2009). This approach is included within a single or multiple allele-discrimination principles, following analyzed with signal detection mechanisms. SNP markers have been further used to breed high values of economic traits by breeders. Advantages of SNP genotyping techniques which was rapid, high throughput and accurate was developed for determining genetic variation (Huang et al., 2015, Syvanen, 2001). The purpose of this study was to use this assay for determining nucleotide variations in RYR1 and H-FABP and to investigate the frequency of these genotypes in pigs exhibiting PSS during transportation and auction period.

**MATERIALS AND METHODS**

The protocol used in this study was approved by the institutional animal care and use committee of national chiayi university.

**Animals and DNA preparation:** Six hundred and fifty five pigs that were exposed to stress due to transportation and auction and hence, prone to PSS were sampled from the main auction market in central Taiwan during summer season (June-August). These pigs were either Landrace × Yorkshire × Duroc or Landrace × Duroc hybrids. The lot contained three hundred and twenty seven females and three hundred and twenty eight male pigs. The transport time from farms to the auction market was approximately 1 h and pigs were auctioned after at least 3 h of laringage. No pigs were given sedatives or treated with electric prods during loading, unloading or the auction process. PSS and normal pigs were assessed among the pigs arriving in the laringage in the auction market (transportation group) or those being auctioned (auction group) and blood samples were collected by the veterinarian of the auction market. PSS pigs were defined as pigs with muscle and tail tremor, labored and irregular breathing, blanching and reddening of skin, a rapid rise in body temperature, the ones that collapsed and showed muscle rigidity. Blood samples were collected from the jugular vein of each animal and stored in tubes containing sterile-EDTA as the anticoagulant. Genomic DNA was isolated from the pig blood as described by Shian and Huang (1997) and quantified by spectrophotometry at S2000 UV/VIS (Tomy Seiko Co., LTD, Tokyo, Japan). The genomic DNA was then diluted with double distilled water to a concentration of 50 ng/μL and used for PCR.

**SNPShot method:** DNA fragments from RYR1 and portions of the 5'-upstream region and intron 2 Section of H-FABP were amplified in a single multiplex PCR. Each reaction was carried out in a total volume 10 μL which contained 25 ng genomic DNA as the template, 5 mM MgCl₂, 2.5 mM dNTPs, 50 pmol/μL each of the six primers (Table 1 and 0.2 U of DyNAzyme II polymerase (Finnzymes Inc., Finland). Reactions were carried out in a PCR thermocycler (GeneAmp® 2700, Applied Biosystems), using the following conditions: an initial denaturation step at 94°C for 5 min, followed by 35 cycles consisting of denaturation at 94°C for 30 sec, annealing at 60°C for 1 min, and extension at 72°C for 1 min with a final extension at 72°C for 7 min. Unincorporated dNTPs and excess primers were inactivated and degraded by the addition of 0.2 U exonuclease 1 (Epicycle®) and 0.5 U shrimp alkaline phosphatase (SAP; Promega) in a final reaction volume of 2 μL which contained 1.5 μL of the PCR product. This degradation reaction mixture was incubated at 37°C for 2 h and the enzymes were subsequently, inactivated at 75°C for 15 min.

These treated PCR products were then processed in a multiplex mini-sequencing reaction where four SNP loci were simultaneously analyzed. The SNP-specific primers designed to detect nucleotide variations by mini-sequencing at the selected SNP sites are listed in Table 1. Conditions were created so that, when the template DNA annealed next to each SNP site, extension by DNA polymerase added just a single complementary deoxyribonucleoside Triphosphate (dNTP) to the nucleotide at the polymorphic site (Table 1) Each of the four dNTPs was labeled with a spectrally distinct

<table>
<thead>
<tr>
<th>Gene name</th>
<th>PCR primer sequences 5’-3’</th>
<th>Product length (bp)</th>
<th>Extension primer sequences 5’-3’</th>
</tr>
</thead>
<tbody>
<tr>
<td>RYR 1</td>
<td>GACGCGCTCTCTTCAGTTAGAGAT</td>
<td>67</td>
<td>CRC-1804: GCAATAGTGCGCCTGTTG (T/C)</td>
</tr>
<tr>
<td>H-FABP 5’ upstream</td>
<td>H-FABP upst F2: GGACCGGAGGTCGACGCAG CG</td>
<td>709</td>
<td>H-FABP upst-1324: CAGCGCTCTCCTCAGAT (C/T)</td>
</tr>
<tr>
<td>H-FABP intron 2</td>
<td>H-FABP intr F2: AAAGCGCGAATCAGCTATGGGATG (C/G)</td>
<td>816</td>
<td>H-FABP intr-1489: GACTGACCTACACTTCCTCAGGA (C/T)</td>
</tr>
</tbody>
</table>

Table 1: Sequences of primers used for the amplification and extension of the RYR1 and H-FABP genes in the mini-sequencing reactions
fluorophore. In addition to the labeled ddNTPS, the primer length differed at the 5' end of each SNP site and non-homologous d (GATC). Two fluorophores were used to analyze each SNP (Table 2). After the multiplex PCR, the nucleotide variation was verified by fluorescence and the length of PCR product using capillary electrophoresis.

Each multiplex mini-sequencing reaction contained 2 µL treated multiplex PCR product, various concentrations of the minisequencing primers, 0.7 µL SNaPshot™ multiplex ready reaction mix AmpliTaq® DNA polymerase, fluorescently labeled ddNTPs and reaction buffer (Applied Biosystems) in a total volume 4 µL. The reaction mixture was subjected to 25 single-base extension cycles comprising denaturation at 96°C for 10 sec, primer annealing at 50°C for 5 sec and primer extension at 60°C for 10 sec. The unincorporated fluorescent ddNTPs were removed by adding 0.35 U of shrimp alkaline phosphatase to the mix and incubating at 37°C for 1 h, followed by heat inactivation at 75°C for 15 min. Automated capillary electrophoresis was used to determine the nucleotide sequence at each SNP which was carried out using a mixture of 0.5 µL of treated multiplex minisequencing products, 9.25 µL of HiDi™ formamide and 0.25 µL of GeneScan-120 LIZ size standard (Applied Biosystems). The products were resolved on an ABI PRISM® 3100 Genetic Analyzer (Applied Biosystems) for 45 min and any nucleotide variation was detected by Gene Scan™ Version 3.7 Application Software (Applied Biosystems). Point mutations were verified by their fluorescent color and length (Table 2).

### RESULTS AND DISCUSSION

The nucleotide variations of four SNPs in RYR and H-FABP were detected unambiguously and simultaneously by the primer extension method. The migration distance of the labeled multiplex PCR product after electrophoresis depended on the length of the designed primer and the molecular weight of the labeled fluorescent product. Moreover, homozygous or heterozygous genotypes were distinguished by the expression of peaks of one or two different colors, respectively, at each SNP site (Table 2 and Fig. 1). For example, the RYR1 product containing 1843C-T polymorphism migrated more slowly than the one containing C; thus, the SNP for the RYR1 1843 C/T heterozygous sample displayed two peaks having a minimum overlap (Fig. 1b). The C/G heterozygous genotype in the SNP 1811 of H-FABP showed two overlapping peaks that could be distinguished by the variations in the fluorescence of ddGTP (blue) and ddCTP (black) (Table 2, Fig. 1a-c). We found that the genotype nn (a homozygote with T at 1843) of RYR1 was the most common allele in pigs that developed PSS during transport and auction period. The frequency of the nn genotype was higher in the transportation (0.6) and auction group (0.7) than in the normal group (0.01) (Table 3). Three SNP sites in H-FABP were also examined. The genotype frequency of each locus showed no difference in the normal, transport and auction groups. However, it is interesting to note that the major genotype was different among these loci. For the H allele (5' upstream region at position 1324), the HH genotype (T/T) was the major genotype with a frequency of 0.82. For the D allele (intron 2 at position 1811), the frequency was 0.04 for the DD genotype (C/C) 0.42 for the Dd genotype (C/G) and 0.56 for the dd genotype (G/G). In contrast to the former two alleles mentioned above, the heterozygous genotype (Aa, C/T) of the A allele (intron 2 at position 1489) was most commonly encountered with a frequency of 0.52. The occurrence of the homozygous recessive genotype (aa, T/T) was rare (0.02).

The transportation and auction processes present a combination of physical and unfamiliar experiences such as high ambient temperature, noise and pig-human interaction that can be stressful for pig. These sequential stressors can induce PSS which results in acute death and poor pork quality (Johnson et al., 2013; Warris, 2004). Besides, its association with environmental factors, PSS is closely related to C to T polymorphism in RYR1 at nucleotide position 1843 (De Smet et al., 1996). In Taiwan, market-weight pigs are auctioned alive. After transportation and lairage, pigs are forced to walk on a narrow route (about 10 m length and 0.5 m width) one by one. As the pigs walk through the route, competitive bidding is held within 1 min. Body conformation and type are the most important criteria assessed by buyers who prefer pigs with a lean-meat
phenotype. Therefore, pig farmers or breeders tend to select herds with lean-meat phenotype and high meat production. This breeding program focused on lean-meat phenotype is coupled with better growth performance and high susceptibility to exertional myopathy (Altrock and Holleben, 1999); however, it might potentially result in an increase in then genotype and in the incidence of PSS and Pale, Soft, Exudative meat (PSE). Studies have shown that Nn pigs have better feed efficiency and carcass characteristics than NN pigs (Leach et al., 1996). Fisher et al. (2000) concluded that the inclusion of halothane gene had positive impacts on carcass characteristics such as carcass length and lean content of certain cuts but was associated with a higher incidence of PSE (Fisher et al., 2000). In our study, pigs which developed PSS had a greater (0.9) incidence of the n RYR1 genotype. The result was similar to that reported by Murray and Johnson (1998) who showed that more than 50% of pre-slaughter deaths were mainly caused by the presence of halothane (n) gene in Western Canadian commercial packing plants. However, in our study, pigs that were identified to have PSS showed serious physical symptoms and were slaughtered before their death. It is interesting that PSS was present not only in the nn carrier pigs but also partially in the NN or Nn pigs. This result was similar to that by Ritter et al. (2008) who indicated that the pre-slaughter loss including norrambulatory and dead pigs was not only attributable to the nn genotype of RYR1 but also to the non-genetic
factors. The H-FABP genotype which is another important gene for pig industry is a candidate gene for determining the muscle quality and might be associated with the degree of Intramuscular Fat (IMF) (Gerbens et al., 1997, 1998). The genotypes of H-FABP were observed to have significant effects on IMF at sites such as the eye muscle area (Gerbens et al., 1999; Lin et al., 2002; Zhang et al., 2001) but not on the backfat thickness (De Koning et al., 1999). A correlation between the genotype and variation in fat content was also found in the duroc breed (Gerbens et al., 1999). In this study, the frequencies of H-FABP genotype showed that there was a higher frequency of PSS development in the HH, dd, Dd, Aa and AA genotypes whereas lower frequencies were associated with the Hh, hh, DD and aa genotypes. Our data also showed that the frequencies of the dominant genotypes were 0.98, 0.42 and 0.98 for H (HH and Hh) D (DD and Dd) and A (AA and Aa) alleles, respectively. These genotype frequencies were inconsistent with those reported in previous studies (Gerbens et al., 1999; Lin et al., 2002) using pure-breed pigs. However, the genotype frequency of H-FABP and its effect on IMF differed among the pig breeds and populations (Nechtelberger et al., 2001; Fang et al., 2006). In our study, samples were collected from hybrid pigs in which the variation in the genotype frequencies was more significant than in the pure-breed pigs. Therefore, a large population was needed to investigate the correlation between PSS and the genotype frequency of H-FABP.

The advantages of the primer extension method such as SnaPshot assay are accuracy and automation for high throughput genotyping and therefore, this method is efficient for a large survey project and is ideal for a central laboratory (Gwee et al., 2003; Pati et al., 2004; Syvanen, 2001). Moreover, microarray analysis of SNPs has been used in commercial pig breeding (Kaminski et al., 2008). Therefore, a primer extension method for simultaneous screening of five important mutations on two genes was developed in this study.

CONCLUSION

In conclusion, PSS was correlated with the SNP genotypes of RYR1 in pigs during transportation and auction period but not with H-FABP. A controlled breeding program to eliminate the nm pigs would be effective in reducing the pre-slaughter loss; however, improvements in humane handling and minimization of the stressors would be also necessary in the auction market.

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REFERENCES


