Identification of More Than Two Paternal Haplotypes of the Ovine Fatty Acid-Binding Protein 4 (FABP4) Gene in Half-Sib Families: Evidence of Intragenic Meiotic Recombination

Wei Yan¹, Huitong Zhou², Yuzhu Luo¹⁎, Jiang Hu¹, Jon G. H. Hickford²⁎

¹ Gansu Key Laboratory of Herbivorous Animal Biotechnology, Faculty of Animal Science and Technology, Gansu Agricultural University, Lanzhou, China, ²Gene-Marker Laboratory, Faculty of Agriculture and Life Sciences, Lincoln University, Lincoln, New Zealand

Abstract

The fatty acid binding protein 4 (FABP4) plays an important role in the regulation of lipid metabolism in mammals. In this study, two regions of ovine FABP4 spanning exon 2-intron 2 and exon 3-intron 3 were investigated in four hundred and twenty lambs derived from seven sires that were previously typed as having heterozygous genotypes in both these regions of the gene. These regions have been shown to be variable, with three SNPs plus one indel and four SNPs respectively constituting five and four allele variants in the two regions. Across these regions, fourteen haplotypes have been identified. The lambs were typed using a Polymerase Chain Reaction Single-Stranded Conformational Polymorphism (PCR-SSCP) method to identify the haplotypes inherited from the sires. Between three and four paternally-derived haplotypes were identified in the progeny of six of the seven sires, suggesting that meiotic recombination occurs within ovine FABP4. A number of sequence motifs associated with recombination “hotspots” were detected in the two regions of the gene that were analyzed and these may facilitate the recombination.

Citation: Yan W, Zhou H, Luo Y, Hu J, Hickford JGH (2014) Identification of More Than Two Paternal Haplotypes of the Ovine Fatty Acid-Binding Protein 4 (FABP4) Gene in Half-Sib Families: Evidence of Intragenic Meiotic Recombination. PLoS ONE 9(2): e88691. doi:10.1371/journal.pone.0088691

Editor: Qinghua Shi, University of Science and Technology of China, China

Received August 11, 2013; Accepted January 9, 2014; Published February 11, 2014

Copyright: © 2014 Yan et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work is funded by the Chinese International Cooperation and Exchanges Program, the Gansu Province Creative Research Groups Program and the Lincoln University Gene-Marker Laboratory. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

⁎ E-mail: luoyz@gsau.edu.cn (YL); jon.hickford@lincoln.ac.nz (JGH)

Introduction

The fatty acid binding protein 4 (FABP4; also known as adipocyte fatty-acid binding protein) belongs to a fatty-acid binding protein family that is comprised of at least nine members in mammals [1]. The protein is predominantly expressed in white and brown adipose cells; but it is also found in many other cells and tissues, including blood, liver, heart, brain, skin and muscle [2–4]. As an intracellular protein, it is thought to be involved in lipid metabolism and acts by binding and transporting long-chain fatty acids [5].

In humans, the FABP4 gene has been shown to be associated with the risk of acquiring hyper-triglyceridemia, type 2 diabetes and cardiovascular disease [6]. Studies of farm animals have revealed that it may have an impact on growth in pigs [7], fat deposition, marbling and carcass weight in cattle [8,9], and fleece-rot resistance in sheep [10].

FABP4 consists of four small coding exons and three introns. The gene has been described in many species, but genetic variation has only been reported in some [6,8,11]. An unexpectedly high level of polymorphism has been reported in pigs, with approximately one nucleotide substitution in every 50 bp [7]. Little is known of the mechanism(s) that create(s) this genetic diversity.

We have recently reported five and four sequence variants, containing three SNPs plus one indel and four SNPs respectively, in two separate regions of ovine FABP4 (Figure 1) [11]. Fourteen haplotype sequences were constructed across these two regions [11] and some of them appear to have been created by intragenic recombination events. These regions contain the majority of the FABP4 coding sequence and some of the encoded amino acids appear to be important for the functional domains of the protein in the tertiary structure [12]. Sequence similarity is observed between species in these regions (Figure 1). To confirm whether meiotic recombination occurs within the gene, we investigated the inheritance of paternal haplotypes in 359 lambs derived from seven sires.

Materials and Methods

Sheep blood and DNA samples

Blood samples from commercially farmed sheep on private farms were collected onto FTA cards (Whatman, Middlesex, UK) by the owners. The collection of blood drops by nicking sheep ears, is covered by Section 7.5 Animal Identification and Minimal Standard No. 13 - Identification, of the Animal Welfare (Sheep and Beef Cattle) Code of Welfare 2010; a code of welfare issued under the Animal Welfare Act 1999: Public Act 1999 No. 142,
New Zealand (NZ) Government. Blood samples are routinely collected for commercial DNA testing for various traits. Blood samples were collected from all the sheep onto FTA cards (Whatman, Middlesex, UK) and genomic DNA was purified using a two-step procedure [13]. Initially, by typing NZ Romney sires, seven were identified that were heterozygous in both regions of FABP4. These sires produced 420 lambs that were subsequently typed for two regions of FABP4. The lambs and sires were also typed for variation in PRNP [14], ADRB3 [15] and the MHC-DQA2 - DQA2-like region [16] to confirm their sire group. The sires and lambs were all derived from the ongoing NZ Romney progeny test 2006-2013. This genetic trial involves the single-sire mating of selected NZ Romney rams to approximate 60 ewes each, with lambs identified to ram and ewe at birth and phenotypic data collected for growth and carcass traits. DNA is collected from all the sheep and lambs annually.

Genotyping of ovine FABP4

Two regions of ovine FABP4 were separately typed using a Polymerase Chain Reaction Single-Stranded Conformational Polymorphism (PCR-SSCP) method as described previously [11]. Briefly, region 1 (exon 2 - intron 2) and region II (exon 3-intron 3) by (a) PCR-SSCP and (b) DNA sequencing [11]. The FABP4 sequences from other species used to construct the homology trees (c) are: NC_007312 (cattle), EF661481 (pig), HQ384160 (human), ENSFCAG00000032028 (cat) and ENSMUSG00000062515 (mouse). The nucleotide numbering follows the nomenclature described on www.hgvs.org/mutnomen/. doi:10.1371/journal.pone.0088691.g001

Figure 1. Polymorphism of ovine FABP4 and sequence homology with other species. Five and four allele variants are respectively identified in region I (exon 2-intron 2) and region II (exon 3-intron 3) by (a) PCR-SSCP and (b) DNA sequencing [11]. The FABP4 sequences from other species used to construct the homology trees (c) are: NC_007312 (cattle), EF661481 (pig), HQ384160 (human), ENSFCAG00000032028 (cat) and ENSMUSG00000062515 (mouse). The nucleotide numbering follows the nomenclature described on www.hgvs.org/mutnomen/. doi:10.1371/journal.pone.0088691.g001

After denaturation and rapid cooling on wet ice, PCR amplicons were electrophoresed in 14% (for region 1), or 10% (for region 2) polyacrylamide gels for 19 hours at 7.5°C and 390 V (for region 1) or 300 V (for region 2). Amplifications were performed in a 15-μL reaction containing the DNA on one 1.2 mm punch of FTA card, 0.25 mM of each paired primer set, 150 μM dNTPs (Bioline, London, UK), 2.5 mM Mg2+, 0.5 U Taq DNA polymerase (Qiagen, Hilden, Germany) and 1X the reaction buffer supplied with the enzyme. The thermal profiles consisted of 2 min at 94°C, followed by 35 cycles of 30 s at 94°C, 30 s at 60°C and 30 s at 72°C, with a final extension of 5 min at 72°C.

Determination of paternal haplotypes

At each region, by comparing the progeny and sire genotypes, the sequence variants inherited from the sires could be determined.
for the progeny. The exception was with lambs that had the same genotypes as their sires. These lambs were excluded from analysis as the variants inherited from the sires could not be resolved, and maternal DNA was not available to assist in this resolution.

Having determined the sequence variants inherited from the sire for each of the two regions, the paternal haplotypes across the regions could be constructed.

Results

Of the 420 progeny investigated, 188 (44.8%) possessed the same genotypes as their sires at either one or both regions of FABP4, and hence the haplotypes inherited from the sires across these regions could not be determined. For the remaining 232 (55.2%) of the progeny, the parental haplotypes could be determined. More than two paternal haplotypes were identified in all but one of the sire-lines (Table 1). In each of the sire-lines where more than two paternal haplotypes were observed, only two haplotypes were common, while the other/others was/were detected at a lower frequency (Table 1). These less frequent haplotype(s) could be generated from the main paternal haplotypes by meiotic DNA recombination between the two gene regions that were amplified.

The paternity of all the lambs was confirmed by typing for variation in PRNP, ADRB and the MHC-DQA2 - DQA2-like region (results not shown). Within the ovine FABP4 regions analysed, we found a number of simple sequence motifs that have been reported to be associated with recombination hotspots in both prokaryotes and eukaryotes.

Discussion

In this study, we report the detection of three or four haplotypes of ovine FABP4 inherited from a single sire. Of these, two haplotypes were commonly found, and the less common haplotypes appear to have come-about from meiotic recombination between the two regions of the gene that were amplified. In the absence of meiotic recombination, only two sire haplotypes would typically be observed, whereas theoretically four different paternal haplotypes including two un-rearranged and two recombined haplotypes would be expected in the progeny if a single recombination event occurred within a gene. If we assume that the rate of recombination is the same across all the sires, then based on the recombination that could be observed in the 232 typeable progeny, we would guess the average recombination rate is 11/232 genotypes or 4.7%. This is considerably higher than one might expect given the proximity of the two regions typed in this gene, but caution is needed in making this interpretation given that only 232 out of 420 progeny could be typed.

### Table 1. Paternal haplotypes of ovine FABP4 detected in individual sire-lines.

| Sire ID | Sire genotype | Total number of progeny | Progeny with paternal haplotype identified |
|---------|---------------|-------------------------|------------------------------------------|
| Totaranui 376/02 | A1/D1 A2/B2 | 67 | A1-B2 23 (57.5%) |
|           |               |             | D1-A2 17 (42.5%) |
| Waidale 618/04 | C1/D1 A2/C2 | 68 | C1-C2 22 (46.8%) |
|           |               |             | D1-A2 24 (51.1%) |
|           |               |             | C1-A2 1 (2.1%) |
| Snowlea 192/02 | B1/C1 B2/C2 | 65 | B1-B2 16 (45.7%) |
|           |               |             | C1-C2 18 (51.4%) |
|           |               |             | C1-B2 1 (2.9%) |
| Mana 83/04 | B1/C1 B2/C2 | 62 | B1-B2 17 (54.8%) |
|           |               |             | C1-C2 13 (41.9%) |
|           |               |             | C1-B2 1 (3.2%) |
| Doughboy 45/04 | B1/C1 A2/B2 | 69 | B1-B2 22 (52.4%) |
|           |               |             | C1-A2 16 (38.1%) |
|           |               |             | C1-B2 4 (9.5%) |
| Glenleith 25/02 | A1/B1 A2/B2 | 42 | A1-A2 11 (57.9%) |
|           |               |             | B1-B2 7 (36.8%) |
|           |               |             | B1-A2 1 (5.3%) |
| Mana 90/01 | A1/C1 A2/B2 | 47 | A1-B2 8 (44.4%) |
|           |               |             | C1-A2 7 (38.9%) |
|           |               |             | A1-A2 2 (11.1%) |
|           |               |             | C1-B2 1 (5.6%) |

* Recombined minor sire haplotypes.
doi:10.1371/journal.pone.0088691.t001

These included: 1) a 14-mer sequence (GCTGGTGCTGGTGTA) consisting of two partially overlapping chi-like sequences (GCTGGTGCA and GCTGGTGTA) in region 2; 2) a CRE-like sequence (ATGAAGTCA) in region 1; 3) a CCTCCCT motif approximately 2 kb upstream of region 1 and variants of this sequence in region 1 and region 2; and 4) a number of copies of a CCAAT motif in region 1, region 2, and other regions of the gene (Figure 2).
The detection of only one recombined haplotype in some sire-lines may be because the paternal haplotypes in approximately half of the progeny could not be determined using the PCR-SSCP technique (the progeny typed the same as the sire) and/or because the recombined haplotypes occurred at a low frequency. It seems likely that other recombined paternal haplotypes could be detected if all the progeny could be effectively typed or additional sire lines were typed.

Meiotic recombination occurs frequently in some regions of various genomes and these have been called “recombination hotspots” [18]. The presence of a number of motifs that have been associated with “recombination hotspots” in ovine FABP4 (Figure 2) supports our contention that meiotic recombination occurs with this gene.

Firstly two partially overlapping chi-like sequences (GCTGGTGC and GCTGGTGA) were found in region 2 of the gene. The chi sequence, GCTGGTGG, is the most notable hotspot known to enhance homologous recombination in E. coli [19]. It serves as a stimulator of DNA double-strand break repair in bacteria, and the resulting single-stranded DNA (ssDNA) is bound by multiple molecules of RecA protein that facilitate “strand invasion” in which one strand of a homologous double-stranded DNA is displaced by the RecA-associated ssDNA [20]. A homolog of bacterial RecA exists in eukaryotes including mice and humans [21,22]. The presence of two chi-like sequences in ovine FABP4 may be important for DNA recombination, as chi-like sequences may be implicated in the activity of recombination hotspots [23]. It is interesting that these two chi-like sequences are partially overlapped in a 14-mer sequence GCTGGTGC-TGTTGA, and that the presence of a C at the eighth position of the 14-mer motif eliminates a chi site, but creates two partially over-lapping chi-like sequences. This 14-mer motif has been previously observed for recombined fimA sequences from Dichlobacter nodosus [24].

A CRE-like sequence (ATGAAGTCA) was found in region 1 of the gene. The CRE sequence (ATGACGTCA), also known as M26, is the binding site for a hetero-dimeric transcription factor Aff1-Pcr1, and it acts as a recombination hotspot in the yeast Schizosaccharomyces pombe (S. pombe) [25,26].

Thirdly, the presence of a CCTCCCT motif and variants of this motif in both regions of the gene may promote recombination. In humans, the CCTCCCT motif is present more frequently in recombination hotspots than elsewhere, and chromosomes active for recombination contain this motif, with the “suppressor” mutation being a change from T to C in its third position [27].

Lastly repeats of a CCAAT motif were present in the gene. The CCAAT motif appears to be associated with recombination hotspot activity in S. pombe [28].

It is also notable that the human FABP4 gene sequence GenBank accession number HQ384160, appears to contain an ALU-like sequence in intron 2 (nucleotides 5801-6089). Ovine intron 2 (accession number NC_019466) and human intron 2 (accession number HQ384160) share approximately 38% homology (results not shown), a major difference being that the ovine intron is much shorter and does not contain a region comparable to the ALU-like sequence. ALU sequences are proposed to facilitate local recombination [29], but the absence of this sequence in sheep, would suggest it is not responsible for the observed recombination.
The presence of these potential “recombination hotspots” in ovine \textit{FABP4} supports the contention that intragenic recombination has occurred in this gene. This is consistent with our previous finding that five sequence variants in region 1 and four variants in region 2 generate 14 different haplotypes across these two regions of the ovine FABP4 gene \cite{11}.

\section*{References}

1. Storch J, Concirro B (2000) The emerging functions and mechanisms of mammalian fatty acid-binding proteins. Annu Rev Nutr 20:73–95
2. Pelton PD, Zhou L, Demarest KT, Burris TP (1999) PPAR-\gamma activation induces the expression of the adipocyte fatty acid-binding protein gene in human macrophages. Biochem Biophys Res Commun 261:436–458
3. Fischer H, Gustafsson T, Sundberg CJ, Norrobin J, Elkan M, et al. (2006) Fatty acid binding protein 4 in human skeletal muscle. Biochem Biophys Res Commun 346:125–130
4. Genecards website. Available: http://www.genecards.org/cgi-bin/carddisp.pl?gene=FABP4. Accessed 2013 Dec 08.
5. Furushashi M, Hotamisligil GS (2006) Fatty acid-binding proteins: role in metabolic diseases and potential as drug targets. Nat Rev Drug Discov 7:409–403
6. Tuncman G, Erbay E, Hom X, De Vivo I, Campos H, et al. (2006) A genetic polymorphism of the bovine Fatty acid binding protein 4 gene. Mol Biol Rep 39:10621–10625
7. Ojeda A, Rozas J, Folch JM, Perez-Enciso M (2006) Unexpected high polymorphism at the FABP4 gene unveils a complex history for pig populations. Genetics 174:219–2227
8. Barendse W, Bunch RJ, Thomas MB, Harrison RE (2009) A splice site single nucleotide polymorphism of the fatty acid binding protein 4 gene appears to be associated with intramuscular fat deposition in longissimus muscle in Australian cattle. Anim Genet 40:770–773
9. Lee SH, van der Werf JJH, Park EW, Oh SJ, Gibson JP, et al. (2010) Genetic polymorphisms of the bovine Fatty acid binding protein 4 gene are significantly associated with marbling and carcass weight in Hanwoo (Korean Cattle). Anim Genet 41:442–444
10. Smith WJ, Li Y, Ingham A, Collis E, McWilliam SM, et al. (2010) A genomics-informed, SNP association study reveals FBLN1 and FABP4 as contributing to resistance to fleece rot in Australian Merino sheep. BMC Vet Res 6:27
11. Byun SO, Fang Q, Zhou H, Hickford JG (2008) Rapid genotyping of the ovine ADRB3 gene by polymerase chain reaction-single-strand conformation polymorphism (PCR-SSCP). Mol Cell Probes 22:69–70
12. Hickford JG, Zhou H, Slow S, Fang Q (2004) Diversity of the ovine DQA2 gene. J Anim Sci 82:1553–1563
13. Byun SO, Fang Q, Zhou H, Hickford JG (2009) An effective method for silver-staining DNA in large numbers of polyacrylamide gels. Anal Biochem 383:174–175
14. Zhou H, Hickford JGH, Fang Q (2005) Technical Note: Determination of alleles of the ovine \textit{PKRN} gene using PCR-single-strand conformational polymorphism. J Anim Sci 83:745–749
15. Byun SO, Fang Q, Zhou H, Hickford JG (2008) Rapid genotyping of the ovine ADRB3 gene by polymerase chain reaction-single-strand conformation polymorphism (PCR-SSCP). Mol Cell Probes 22:69–70
16. Hickford JGH, Zhou H, Slow S, Fang Q (2004) Diversity of the ovine DQA2 gene. J Anim Sci 82:1553–1563
17. Byun SO, Fang Q, Zhou H, Hickford JG (2009) An effective method for silver-staining DNA in large numbers of polyacrylamide gels. Anal Biochem 383:174–175
18. Lichten M, Goldman AS (1995) Meiotic recombination hotspots. Annu Rev Genet 29:423–444
19. Smith GR (1994) Hotspots of homologous recombination. Experientia 50:234–241
20. Kowalczykowski SC, Dixon DA, Eggleston AK, Lauder SD, Rehrauer WM (1994) Biochemistry of homologous recombination in \textit{Escherichia coli}. Microbiol Rev 58:401–465
21. Morita T, Yoshimura Y, Yamamoto A, Murata K, Mori M, et al. (1993) A mouse homolog of the \textit{Escherichia coli} recA and \textit{Sacharomyces cerevisiae RAD51} genes. Proc Natl Acad Sci U S A 90:6577–6580
22. Shiohara A, Ogawa H, Matsuda Y, Ushio N, Ike K, et al. (1993) Cloning of human, mouse and fission yeast recombination genes homologous to \textit{RAD51} and \textit{radC}. Nat Genet 4:239–243
23. Giordano M, Marchetti C, Chiorboli E, Bona G, Momigliano Richiardi P (1997) Evidence for gene conversion in the generation of extensive polymorphism in the promoter of the growth hormone gene. Hum Genet 100:249–255
24. Zhou H, Hickford JG (2000) Novel flanking subunit genes of \textit{Dichelobacter nodosus} recombination in vivo. J Anim Sci 82:1553–1563
25. Kon N, Kraschuk MD, Warren BG, Smith GR, Wahl WP (1997) Transcription factor Ms1/Mts2 (Aft1/Per1, Gad7/Per1) activates the M26 metiotic recombination hotspot in \textit{Saccharomyces cerevisiae}. Proc Natl Acad Sci U S A 94:13756–13767
26. Schuchert P, Langford M, Kaslin E, Kohli J (1991) Meiotic recombination hotspots of recombination rates and hotspots across the human genome. Science 252:321–324
27. Myers S, Bottole L, Freeman C, McVean G, Donnelly P (2005) A fine-scale map of meiotic recombination rates and hotspots across the human genome. Science 310:321–324
28. Steiner WW, Steiner EM, Girvin AR, Plewik LE (2009) Novel nucleotide sequence motifs that produce hotspots of meiotic recombination in \textit{Dichelobacter nodosus}. Vet Microbiol 134:125–130
29. Batzer MA, Dweinling PL (2002) Alu repeats and human genomic diversity. Nat Rev Genet 3:370–379

\section*{Acknowledgments}

The authors thank Qian Fang, Seung OK Byun and Hua Gong for technical assistance.

\section*{Author Contributions}

Conceived and designed the experiments: WY HZ JH YL JGH. Performed the experiments: WY HZ. Analyzed the data: WY HZ JGH. Wrote the paper: WY HZ JH YL JGH.