Genotyping of Kappa-Casein Gene of Buffalo in Indonesian

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ABSTRACT

Casein (KCN) is a milk protein with four variants of alpha S1, alpha S2, beta and kappa which has different allele of each. Kappa Casein (CSN3) gene lies at exon 4 associating with fat and protein milk contents. Genetic variation of CSN3 gene influences quality and composition of milk. The aim of this research was to genotype the CSN3 gene of Buffalos from several places in Indonesia. Amount of 29 buffalo samples was used in this research, those consisted of 15 collected from Medan, 3 samples of Banyuwangi, 10 samples of Baluran and one sample of South-East Nusa Tenggara (NTT). A Polymerase Chain Reaction-Restriction Fragment Length Polymorphism (PCR-RFLP) method was used for CSN3 gene analysis. The CSN3 gene was amplified at 56°C of annealing with 30 cycles and resulted a right fragment of 379 bp. The fragments were digested with HindIII at 37°C during for four hours for genotyping analysis. The result showed that BB genotypes were found at the locus of CSN3 gene in all samples. Those BB genotypes merged the pattern of digested fragments with sizes of 255 and 154 bp. This finding suggests that genotyping using HindIII reveals monomorphics of BB types associated with milk content of casein.

Key Words: KCN Gene, Genotyping, PCR-RFLP, Indonesian Buffalo

INTRODUCTION

Indonesia is known as a country with its mega biodiversity among species, breed and population in term of animal genetic resources (FAO 2013). Domesticated ruminant animals have been contributing to this country in supplying of protein both from meat and milk sources. Up to present, protein consumption of people is dominated by cattle and not popular yet fulfilled by milk either from goat or buffalo. In 2014, buffalo population in Indonesia is very low (1,320,616) compared to cattle (14,703,000) population (DGLSAH 2014). Both buffalo and cattle have a chance to contribute both in meat and milk consumption, however buffalo has not been exposed yet.

Up to present Indonesia still imports 30% meat and 60% milk (FAO 2013). Milk production in cattle is higher than in Buffalo. Lactation period in cattle is up to 305 days (Syrdstad 1993) with dry period of 40-50 days (Univ. Illions 2010) while in buffalo is lower (262-295 days) with a longer dry period of 2-3 months (Tamil Nandu Univ. 2009). Even though a lower milk production in buffalo than in cattle, however chemical contents of buffalo milk is higher than in cattle (Damayanthi et al. 2014). Therefore, milk buffalo has a prospect in future as a source of better milk quality. Milk protein components in river buffalo is higher (Damayanthi et al. 2014) compared to cattle (Susilorini & Sawitri 2007; Pandey & Voskuil 2011). As reported by Damayanthi et al. (2014) that protein milk content of swamp buffalo is higher (5.14±0.37%) than in river buffalo (4.68±0.41%).

Milk is an important source of essential nutrients besides for lactating calves it also as a key raw material for human food preparations (Reinhardt et al. 2012). All over the world people fulfill approximately 13% of their protein requirements from milk and dairy products. Indonesian buffalo has not been explored yet to improve their genetic merit for milk traits. In the context of achieving self-sufficient of meat and milk in Indonesia,
buffalo needs to be explored for their potential in both meat and milk production traits. Milk production relates to protein contents. Milk protein is important trait since influences the milk quality.

Milk protein is divided into two fractions, those are soluble protein named whey protein with constitutes of α-Lactoalbunin and β-lactoglobulin. Another is insoluble fraction named casein which has four variants, i.e., α-S1, α- S2, β-casein and kappa casein (El-Rafey & Darwish 2007; Dogru et al. 2007; Rachagani & Gupta 2008; Deb et al. 2014). Kappa casein (CSN3) is one of the most important milk proteins that controlled by a gene with 5 exon and 4 intron, the CSN3 protein has 19.8 kDa molecular weight and 169 amino acids (Abassi et al. 2009). The kappa casein gene lies at chromosome 6 of bovine and at chromosome 4 of sheep and goat. Casein fraction of milk proteins significantly influences the composition and physico-chemical properties of milk (Grosclaud 1988) and milk production (Ghafoor et al. 2014).

Breaktrhugh in molecular genetics, it contributes animal breeders that have greatly and effectively manipulated the genomes of livestock and enhanced production traits in their herds by selecting individuals with superior traits. Those selected individuals are as sources for the next generations. Therefore, there is a need to use selection methods with based on genomic studies (André 2012). Genes associating with performance parameters can improve the estimation of breeding value and hence can contribute as a suitable inputs for conventional breeding procedures. Genotyping or genetic polymorphism relates to the differences in animal performance, it can be taken into account in the selection process.

This CSN3 gene is important to be explored for looking the best casein content in milk buffalo. Therefore this study was designed to focus on genotyping of kappa casein gene in several Indonesian buffalos to find the best buffalo bearing the gene with marker technology approach. This information is necessary for selection process of the gene relating to the casein content and buffalo milk production based on genomic study.

MATERIAL AND METHODS

Blood sample collection

Fresh blood samples were collected from base point of tails. Amount of 29 buffalo (river and swamp) originated from several places in Indonesia. The blood was collected in 5ml per head by a 21-G needle into a vacuumed-tube containing EDTA. Those 29 buffaloes samples were originated from Medan (15), Banyuwangi (3), Baluran (10), and one sample from East Nusa Tenggara (NTT), see Table 1.

Table 1. List of Indonesian buffalo samples used in the research

<table>
<thead>
<tr>
<th>Buffalo (Bubalus bubalis)</th>
<th>Number (head)</th>
<th>Origin of buffalo</th>
<th>Origin of buffalo</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Medan (Sumatra)</td>
<td>Banyuwangi (East Java)</td>
</tr>
<tr>
<td>River (2n = 50)</td>
<td>15</td>
<td>15 (3M; 12F)</td>
<td>-</td>
</tr>
<tr>
<td>Swamp (2n = 48)</td>
<td>14</td>
<td>-</td>
<td>3 (2M; 1F)</td>
</tr>
<tr>
<td>Total</td>
<td>29</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

M: Male; F: Female
DNA extraction

Genome DNA was manually extracted from the whole blood cells according to a method of Montgomery & Sise (1990). Quality and concentration of the extracted DNA were measured through a spectofotometer of GenQuant Pro (Bioscience, USA). DNA concentration was calculated at λ 260 nm wave length of absorbancy while DNA quality was calculated at λ 260/280 nm. DNA concentration was prepared at 50 ng/µl for PCR work.

Polymorphism chain reaction

Amplification of CSN3 gene (379 bp) was conducted by applying a pair of KCN primer for cattle (Mitra et al. 1998). Those sequence primers of KCN-F (5’-CACGTCACCCA CACCCACATTTATC-3’) dan KCN-R (5’-TAATTAGCCCAT TTCGCCTTCTCTGT-3’), see Figure 1. Total volume of 20 µl containing 10 µl PCR Master Mix of DreamTaq Green PCR Master Mix (2x), 1 µl of 10 pmol/µl each primer, 6 µl DDW and 2 µl of 50 ng/µl DNA template was used as a PCR reaction. Amplification was carried out in a Thermalcycler (Eppendorf, USA) with following condition: pre-denaturation 95°C for 1 min, denaturation 95°C for 1 min, annealing 56°C for 1 min, elongation 72°C for 1 min, was repeated up to 30 cycles followed a final extension 72°C for 5 min, then stepped down temperature up to 10°C. PCR products were analyzed in 1% agarose gel at 100 volt for 1 hour, then immersed in Ethidium Bromide solution. Visualisation of bands used UV transilluminator (MUV21, MajorScience, USA) with a DNA ladder of VC 100 bp Plus (Vivantis, USA).

Restriction fragment length polymorphism

PCR products were digested with restriction enzyme of HindIII (Fermentas, Germany), at SNP 353 A>C (Figure 1). Final of 10 µl reaction volume consisted of 5 µl PCR products, 1 µl of buffer R with 10x BSA (Fermentas), 0.2 µl of 10 unit HindIII and 3.8 µl of DDW. The reaction mixture was incubated at 37°C for four hours in a water bath (Trade Raypa). Following incubation, the digested fragment was analyzed by electrophoresis in a 2% agarose gel and run with 100 volt for 1 hour then stained with Ethidium Bromide (AppliChem, Germany). The digested fragment (polymorphic locus) was compared with a 100 bp DNA ladder in 2% agarose gel. Bands were visualized with a UV transilluminator exposure (MUV21, MajorScience, USA) to observe polymorphic locus, then documented by a digital camera (Nikon, Japan).

Figure 1. KCN primer sequence and restriction site of HindIII (SNP 353 A>C; red sign), based on GenBank AJ841946.1
RESULTS AND DISCUSSION

Kappa-casein gene

Gene of kappa casein (CSN3) was detected in all buffalo samples with fragment size of 379 bp (Figure 2). The length of CSN3 gene in this study is similar to CSN3 gene size reported in cattle (Mitra et al. 1998). This homology might be that buffalo is classified as the same family of Bovidae with subfamily Bovinae and tribe of Bovini (MacEachern et al. 2009).

![Image of DNA ladder and buffalo samples](image)

M: 100 bp DNA ladder; Line 2 to 9: Indonesian buffalo samples

**Figure 2.** Kappa-casein (KCN) gene found in all buffalo samples at 379 bp

The KCN gene of bovine located at chromosome 6 p31 with full length of the gene is 13 kb (Othman et al. 2011). This variant of kappa casein gene locating at exon 4 has alleles A and B. The allele B was reported associating with the high content of protein and fat milk (Rachagani & Gupta 2008; Gangaraj et al. 2008). The KCN gene is associated with the yield and component of milk. Caseins amount to nearly 80% of the protein output in cow milk (Patel et al. 2007a). Caseins are biologically important proteins and they are also a raw material for the cheese making industry (El-Gawad & Ahmed 2011).

**Genotyping of KCN gene**

Genotyping of KCN gene digested by HindIII yielded 2 fragments of 154 bp and 225 bp in all samples (Figure 3A). Those fragment patterns were categorized as genotypes BB in all samples. None of genotype A found in this study which has fragment size of 379 bp and different from genotype BB with 2 fragments (Figure 3B). Those fragments are termed as alleles. This finding is different from cattle which has polymorphic for the kappa casein gene (Doosti et al. 2011; Khaizaran & Al-Razem 2014; Cinar et al. 2016).
This finding of genotype BB in all buffalo samples (swamp and river) in Indonesia is similar to previous reports of allele frequencies which found in most of buffalo (Table 2).

Table 2. Allel frequencies of KCN gene in buffalos

<table>
<thead>
<tr>
<th>Breed of buffalo</th>
<th>Origin of buffalo</th>
<th>Allele frequencies</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water buffalo</td>
<td>Guangxi, China</td>
<td>A: 0, B: 1</td>
<td>Ren et al. (2011)</td>
</tr>
<tr>
<td>Water buffalo</td>
<td>Khouzestan, Iran</td>
<td>A: 0, B: 1</td>
<td>Abbasi et al. (2009)</td>
</tr>
<tr>
<td>River buffalo</td>
<td>Egypt</td>
<td>A: 0.125, B: 0.875</td>
<td>Gouda et al. (2013)</td>
</tr>
<tr>
<td>Nili-Ravi buffalo</td>
<td>Pakistan</td>
<td>A: 0, B: 1</td>
<td>Ghafoor et al. (2014)</td>
</tr>
<tr>
<td>River buffalo</td>
<td>India</td>
<td>A: 0, B: 1</td>
<td>Gangaraj et al. (2008)</td>
</tr>
<tr>
<td>Pandharpuri buffalo</td>
<td>India</td>
<td>A: 0, B: 1</td>
<td>Shende et al. (2009)</td>
</tr>
<tr>
<td>River buffalo</td>
<td>Egypt</td>
<td>A: 0, B: 1</td>
<td>Dayem et al. (2009)</td>
</tr>
<tr>
<td>Murrah, Jaffarabadi, Surti, Pandharpuri</td>
<td>India</td>
<td>A: 0.016, B: 0.984</td>
<td>Patel et al. (2007b)</td>
</tr>
<tr>
<td>River breed</td>
<td>Egypt</td>
<td>A: 0, B: 1</td>
<td>Dayem et al. (2009)</td>
</tr>
<tr>
<td>Nili-Ravi buffalo</td>
<td>Pakistan</td>
<td>A: 0, B: 1</td>
<td>Riaz et al. (2008)</td>
</tr>
<tr>
<td>Anatolian water buffalo</td>
<td>Turkey</td>
<td>A: 0, B: 1</td>
<td>Cinar et al. (2016)</td>
</tr>
<tr>
<td>Egyptian buffalo</td>
<td>Egypt</td>
<td>A: 0, B: 1</td>
<td>Othman et al. (2011)</td>
</tr>
</tbody>
</table>

This finding of genotype BB in the kappa casein gene is associated with protein content in milk. It was known that BB types have a significant influencing on cheese making properties (Shende et al. 2009). It was reported that genotype BB yields more protein than genotype AB and even more milk yields than genotype AA (Rachagani & Gupta 2008; Azevado et al. 2008). As stated by El Nahas et al. (2013), types of BB are differing by mutation points in nucleotide sequence encoding amino acid 136 and 148 bp. Position of 136 which is threonine (ACC) changed into isoleucin (ATC) and position of 148 is arginine (GAT) changed into alanine (GCT) for A and B types, respectively.
Observation of BB genotypes in all samples seems that there was no crossed with other buffalo breed in the most buffalo of Indonesia. It figures that most buffalo in Indonesia have a potential in a higher casein protein content of milk production.

CONCLUSION

A Kappa casein gene was amplified at the right size of 379 bp in all buffalo samples. All swamp and river buffalos used in this research showed monomorphic patterns of the kappa-casein gene with only had allel B and presented in homozygote of BB genotypes. It seems that most buffalo in Indonesia have a potential in a higher casein protein content of milk production. This finding contributes to the breeding program especially in milk protein content of buffalo.

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