TOTAL BACTERIA AND IDENTIFICATION OF PROTEOLYTIC RUMINAL AND RETICULUM BACTERIA FROM LOCAL KACANG GOAT

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INTRODUCTION

Kacang goat is one of the original Indonesian goat whose population quite high and widespread in Indonesia. These goats have a small body with shoulder height of about 50 cm to 60 cm. Some of the benefits of Kacang goat are: efficient in the conversion of grass into meat, resistant to disease, rapid breeding, relatively easy maintenance, not requiring extensive land, the meat is in great interest after beef (Fitra et al., 2009; Sudewo et al., 2012) and able to adapt to tropical environment such as in Aceh. However, this potential is not yet optimal due to the slow growth of the goats and posing a challenge in the effort to improve their productivity, particularly their growth.

Some factors influence the growth of goats, such as the quality and quantity of feed, as well as the process of feed degradation by microbes in the gastrointestinal tract. In farmer community of Aceh, goat feed was not generally given additional concentrate; the forage contain only cellulose, protein, and other nutrients. The crude protein content (CP) of forage varies considerably, ranging from 30 g/kg of dry weight in old grass, to more than 300 g/kg in well-grown young grass (Haryanto, 2012; Yang et al., 2014). The feed was degraded in the rumen, reticulum, and goat omasum, where the digestion in the reticulorumen is very intensive. Protease enzymes, produced by proteolytic microbes, hydrolyze feed proteins in the rumen (Kim et al., 2011; McSweeney and Mackle, 2012; Muslim et al., 2014).

Protease is a group of enzymes which hydrolyzes polypeptide bonds to oligopeptides and amino acids, and then hydrolyzed further to produce ammonia (NH3). Ammonia is utilized by microbes as the primary source of nitrogen for protein synthesis (Kumar et al., 2012). According to Genzenbu and Tesfay (2015), bacteria capable of utilizing nitrogen in the rumen are classified as proteolytic bacteria. Bacteria are found in largest population inside the rumen, and 38% of the total bacteria in the rumen are proteolytic bacteria. The availability and ability of proteolytic bacteria to degrade proteins will affect the amount of amino acids that enter the bloodstream.

Eschenlauer et al. (2002) showed that the total of proteolytic bacteria in the goat rumen is 10^9 cells/g, whereas the number and type of proteolytic bacteria in the reticulum has not been reported. Proteolytic bacteria in goat rumen that have been identified are:
Proteus vulgaris, Streptobacillus, Staphylococcus, Streptococcus, Clostridium butylicum, Clostridium sporogens, Lactobacillus pentosus, Propionibacterium pentosaceum, Corynebacteria, Bacteroides amylophilus, Bacteroides ruminicola, Butyrivibrio fibrisolvens, Butyrivibrio alactadigenus, Selenomonas ruminantium var. ruminantium, Selenomonas ruminantium var. lactilytica, Fibrobacter succinogenes, Butyrivibrio fibrisolvens, Ruminobacter amylophilus, and Streptococcus bovis (Kamra, 2005; McSweeney and Mackle, 2012; Hungate, 2013; Krause et al., 2014). Until present, no information on the total and types of proteolytic bacteria in Kacang goat in Aceh has been reported, requiring studies to be conducted on the number and types of proteolytic bacteria in rumen and reticulum of Kacang goat.

MATERIALS AND METHODS

Isolation of Proteolytic Bacteria and Total of Proteolytic Bacteria

Five goats were slaughtered at slaughter house and had its rumen and reticulum fluid sampled. Subsequently the liquid sample was diluted with a 10⁻¹ to 10⁻³ dilution. One ml solution of each dilution was inserted into sterile petri dish, and then 20 mL of skim milk agar medium was added by pour plate method in duplicate before incubated for 48 hours at 39°C. Total proteolytic bacteria were calculated from the bacterial colonies that grow from the 10⁻² to 10⁻³ dilution by multiplying the number of colonies with the dilution factor. The colonies that grow was Gram stained and its clear zone was measured. The pure colonies with the largest proteolytic index were cultured into liquid medium and incubated for 48 hours. The incubation result was centrifuged at 7000 g and its supernatant was disposed. The DNA of the cell pellet was then isolated.

Total DNA Isolation

Total DNA isolation used Presto™ Mini gDNA Bacteria kit (Geneaid). Cell pellet was added with 180 µL extraction buffer and 20 µL Proteinase K and was then incubated at 60°C. About 200 µL GB buffer is added to the sample. The sample was mixed and incubated at 70°C. Afterward, 200 µL of absolute ethanol was added and the sample was inserted into the tube column and centrifuged at 14000-16000 x g for 1 minute. A total of 400 µL of buffer W1, was added into the tube column and centrifuged 14000-16000 x g for 30 seconds. The supernatant was discarded and the column was added with 600 µL buffer wash, centrifuged at 14000-16000 x g for 30 seconds and its supernatant discarded. The column was then added with 30-50 µL elution buffer and incubated at room temperature for 3-5 minutes, and then centrifuged at 14000-16000 x g for 1 minute.

Amplification of 16S rRNA Gene

The gene was amplified with PCR. One of the primers (BacF) used was a complement to conserved areas in the bacterial domain, and the other primary domains (UniB) is based on universal conserved area of the 16S rRNA E. coli gene (Baker et al., 2003). A total of 25 µL PCR reaction mixture was used for DNA amplification. The PCR cycle was 1 minute for denaturation at 94°C, 1 minute for annealing at 50°C and 2 minute for primary elongation at 72°C. The final stages of primary elongation were carried out for 8 minute at 72°C (Safika et al., 2013). Electrophoresis was then performed to the PCR result on 1% agarose gel.

Determination of DNA Sequence

DNA sequencing is performed through commercial services at Macrogen Inc., Korea. The process of determination is using the Dye Terminator method (3'-dye labelled dideoxynucleotide triphosphate) which includes several stages, namely preparation of templates, sequencing reactions, PCR product purification, and electrophoresis with scanning fluorescence.

Phylogenetic Analysis

The sequencing results are compared using program Basic Local Alignment Search Tool (BLAST) program at NCBI http://www.ncbi.nlm.nih.gov. Homology analysis of the 16S rRNA gene sequence used the data in GenBank. The phylogenetic tree was constructed using a neighbor-joining model matrix from the MEGA 7.0 program (Kumar et al., 2016), with the maximum composite likelihood substitution method (Tamura et al., 2004) with a bootstrap analysis 1000 times.

RESULTS AND DISCUSSION

The Morphology of Proteolytic Bacteria

The growth of proteolytic bacteria colonies from rumen and reticulum fluid of Kacang goat were seen spreading inside the skim milk agar (SMA) medium. Colonies that grow in the media indicated that the bacteria are facultative anaerobic bacteria. The growing colonies have a variety of morphologies. The result of isolation of proteolytic bacteria can be seen in Figure 1. Proteolytic colonies were observed its morphologies, such as, color, shape, position, fringe, and colony diameter. Based on the observations made, bacterial colonies that grow were generally white with rounded shape, and smooth edges, with varying proteolytic index (IP) as can be seen in Table 1.

The IP in goat rumen fluid ranges from 1.0 to 2.5 mm and in the reticulum fluid ranges from 1.0-2.0 mm. The IP is the diameter of the clear zone divided by the diameter of bacterial colonies (Baehaki et al., 2011). The clear zone formed is an indicator that the growing bacteria have proteolytic activity because these bacteria are able to produce protease enzymes that hydrolyze casein (Rudwitasari and Wikandari, 2014). Skim milk contains casein in which casein is the main protein of the milk, a macromolecule composed of amino acid subunits associated with peptide bonds. Casein serves as a substrate for protease enzymes. Casein hydrolysis...
was used to show the hydrolytic activity of proteases (Susanti, 2003; Yuanita and Wikandari, 2014). Increased enzyme activity is evident with the increase in the diameter of the clear zone present around bacterial colonies on skim milk media. Based on the morphological observations of Gram staining (Figure 2), the bacteria found are generally gram-positive coccus or basil bacteria and Gram-negative basil bacteria in both the rumen and the reticulum fluid (Table 1).

**Table 1. Morphology of bacterial colonies on skim milk agar medium**

| Colony | Shape | Color | Position | Edge | Elevation angle | Proteolytic index (mm) | Gram staining |
|--------|-------|-------|----------|------|-----------------|-----------------------|--------------|
| **RUMEN** |       |       |          |      |                 |                       |              |
| Ru 1   | Round | White | Deep     | Smooth | Convex          | 2.0                   | Coccus (+)   |
| Ru 2   | Round | White | Deep     | Smooth | Convex          | 1.5                   | Coccus (+)   |
| Ru 3   | Round | White | Deep     | Smooth | Convex          | 2.5                   | Bacillus (+) |
| Ru 4   | Round | White | Deep     | Smooth | Convex          | 1.3                   | Coccus (+)   |
| Ru 5   | Round | White | Deep     | Smooth | Convex          | 1.0                   | Bacillus (+) |
| **RETIKULUM** |       |       |          |      |                 |                       |              |
| Re 1   | Round | White | Deep     | Smooth | Convex          | 2.0                   | Bacillus (-) |
| Re 2   | Round | White | Deep     | Smooth | Convex          | 1.5                   | Coccus (+)   |
| Re 3   | Round | White | Deep     | Smooth | Convex          | 1.2                   | Coccus (+)   |
| Re 4   | Round | White | Deep     | Smooth | Convex          | 1.3                   | Bacillus (+) |
| Re 5   | Round | White | Deep     | Smooth | Convex          | 1.0                   | Bacillus (+) |

**Figure 1.** Colonies of bacteria growing on the Skim Milk Agar medium. A= Colonies of rumen fluid at dilution $10^5$; B= Colonies of the reticulum fluid at dilution $10^7$.

Total of Proteolytic Bacteria

Colonies of proteolytic bacteria on all petri dish from each dilution was counted. Dilution reduced the quantity of bacteria that grows on the media. The greater the dilution result in the lesser the number of colonies growing on the media. Dilution can also help facilitate the calculation of the colony. The total of proteolytic bacteria from the rumen fluid and kacang goat reticulum grown on skim milk agar after incubation for 48 hours can be seen in Table 2.

In this study, the number of proteolytic bacteria in the local kacang goat reticulum was greater ($2.382 \times 10^7$ CFU/mL) than in the rumen ($6.416 \times 10^6$ CFU/mL). The total of bacterial results of this study is smaller than Eschenlauer et al. (2002), which stated that the total of proteolytic bacteria in the rumen is $10^9$ cells/g, but the total of proteolytic bacteria in goat or cow reticulum has not been reported. The existence of a number of varying bacteria is influenced by the type and content of feed consumed by the livestock. In general, nutrient content in forages consists of cellulose (35-50%), hemicellulose (20-35%), lignin (50-30%) and protein (18-30%) (Kamra, 2005; Haryanto 2012; Yang et al., 2014). High-fiber feeding, such as forage in large quantities without supplementation with protein-rich feed, causes fewer proteolytic bacteria or microbes in the rumen of Kacang goat. When more animals are fed rough fiber feeds, the populations of fiber-degrading microbes will be dominant (Sari et al., 2017). In addition, according to Jones et al. (1994) and McSweeney et al. (1999), forage feeds such as legume contain many protein-tannin complexes (polyphenols). This tannin reduces the availability of nitrogen in the rumen and inhibits the growth of dominant bacteria in the rumen, especially proteolytic bacteria, due to lack of nitrogen. This is presumed to be one of the causes of the low number of proteolytic bacteria in Kacang goat in Aceh, where the feed was given in the form of forage without the addition of concentrate.

Based on this study, bacteria proteolytic index in rumen is higher, but the number of bacteria is lower than in the reticulum. This is because the rumen is where protein degradation, such as protein hydrolysis by protease enzyme into peptide and amino acid, takes
place. The peptide and amino acid can be utilized directly by microflora or further degraded by peptidase and deaminase enzyme into short chain fatty acid and ammonia. Rumen and reticulum are connected by a channel called reticuloruminal fold, so the food ingesta can flow from the rumen to the reticulum or vice versa. Feed that has not been digested in the rumen will be digested in the reticulum. Proteins consumed by ruminants are not entirely degraded in the rumen. It takes longer for the protein to be digested in the rumen, so undigested proteins are transported to the reticulum for further processing (Nuswantara, 2003; Chiba, 2014; Safika et al., 2017). The presence of proteins in the reticulum causes an increase in number of proteolytic bacteria to hydrolyze proteins by producing protease enzymes.

**DNA Bacteria and Amplicon Gen 16S rRNA**

Isolation of bacterial DNA is an important step in molecular analysis in the field of microbiology. The result of DNA isolation electrophoresis can be seen in Figure 3a. The isolated DNA is then amplified using PCR and the result of can be seen in Figure 3b. In Figure 3b, the intact and non-fragmented DNA bands indicated the existence of specific bacteria DNA and have a good purity. The amplified band is parallel to the 1500 bp marker band.

**Homology of Bacteria**

Homology analysis from BLAST revealed that the DNA sequence of proteolytic bacteria in the rumen of Kacang goat had 93% sequence similarity with *Bacillus aereus*. While the Re1 isolate of Kacang goat reticulum had 92% similarity with *Tatumella punctata*. Phylogenetic analysis using MEGA 7.0 Ru3 isolates was close to the Bacillus group and the Re1 isolate was close to Tatumella (Figure 4).

Identification of bacteria using the 16S rRNA gene sequence is a widely practiced technique, although with limitations for closely related taxa members (Kim et al., 2011). If the sequence has a similarity of greater than 98% with a known bacterial 16S rRNA gene, this species is considered a member of the species, but if less than 97% then it is considered different species (Vandamme et al., 1996; Janda and Abbott, 2007). The homology of 92% sequence for Ru3 and 93% for Re1

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**Table 2. The average number of proteolytic bacteria in the rumen and peanut goat’s reticulum**

| No | Kacang goat | Rumen | Reticulum |
|----|-------------|-------|-----------|
| 1  | Goat 1      | 3.55 x 10^6 | 3.07 x 10^7 |
| 2  | Goat 2      | 6.35 x 10^6 | 1.24 x 10^7 |
| 3  | Goat 3      | 9.40 x 10^6 | 2.82 x 10^7 |
| 4  | Goat 4      | 6.53 x 10^6 | 1.95 x 10^7 |
| 5  | Goat 5      | 6.25 x 10^6 | 2.83 x 10^7 |
| **Means** | **6.416 x 10^6** | **2.382 x 10^7** |
in this study indicated that both isolates are either new or unconfirmed. Further research is required to determine whether the isolates are new species such as DNA-DNA hybridization analysis, G + C content and free fatty acid analysis. Microbial protease enzymes account for about 60% of the total production of enzymes in the industry. Protease is one of the most important groups of industrial enzymes with extensive applications including animal feed, pharmaceuticals, detergent manufacture, cheese making, bread making, waste management and silver recovery. The industrial sector often uses Bacillus subtilis for the production of various enzymes. All proteases, especially the alkaline proteases produced by the Bacillus species, are very important in the industry because they have a stable temperature and a high pH. Bacillus is a major producer of extracellular proteases. The isolation and characterization of new strains of Bacillus in the enzyme production process are promising. This
bacteria are rod-shaped, resistant to extreme environmental conditions, aerobic or facultative anaerobes and include normal flora although some species are pathogenic (Suganthi et al., 2013: Pant et al., 2015).

*Tatumella* spp. is also produce proteases which can grow at pH 5.0-7.0 (Rodarte et al., 2011). Tatumella is a Gram-negative, short-shaped, facultative anaerobes, negative oxidase, and catalase-positive bacteria. This bacteria included in family Erwiniaeaeae, order Enterobacteria, class Gammaproteobacteria and phyla Proteobacteria. Types of bacteria *Tatumella* spp. can grow at pH 5.0-7.0 and is also known to produce protease (Rodarte et al., 2011).

**CONCLUSION**

The amount of proteolytic bacteria with protease activity in reticulum fluid of Kacang goat is higher (2.382x10^7 CFU/mL) compared to the amount of proteolytic bacteria in rumen of Kacang goat (6.416x10^6 CFU/mL). The bacteria with the widest proteolytic index in the fluid of rumen goat had similarity to the Bacillus bacteria (93%), whereas the bacteria in the reticulum fluids have close relation to the Tatumella bacteria (92%). There is a possibility that both isolate is a new or unconfirmed species.

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