

International Journal of **Dairy Science**

ISSN 1811-9743



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International Journal of Dairy Science

ISSN 1811-9743 DOI: 10.3923/ijds.2019.29.35



Research Article Microbiological Qualities of Goat Milk Obtained under Different Milking Systems at a Smallholder Dairy Farm in Yogyakarta, Indonesia

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Abstract

Background and Objective: In Indonesia, smallholder dairy farms contribute 20.37% of domestic market. A small portion of these farms is from goat dairy farms that are traditionally managed with poor sanitation during milking and improper storage management. This system causes contamination that can affect consumer health and cause financial loss. This study aimed to evaluate the microbiological quality of goat milk obtained under different milking systems at a smallholder dairy farm in Yogyakarta, Indonesia. **Materials and Methods:** Samples were collected from 20 crossbred dairy goats divided into two groups: Group A and B. In Group A, milking was conducted manually. In Group B, milking was conducted using a bucket milking machine. Total Plate Count (TPC) and the presence of Enterobacteriaceae (EB), Shiga toxin-producing *Escherichia coli* (STEC) and *Salmonella* were assessed. Data obtained were analyzed using a t-test. **Results:** The results showed that TPC and EB were higher in Group A than in Group B. In Groups A and B, 80% (8/10) of samples contained STEC and 30% (3/10) contained *Salmonella*. Phylogenetic analysis showed that partial sequencing of amplified genomic DNA using *stx1* primers had more than 90% similarity with several sequences of *Escherichia coli* O157:H7 strain Shiga toxin subunit 18 (*stx*1B). Moreover, partial sequencing of amplified genomic DNA using 16S rRNA primers had more than 90% similarity with several sequences of *S. enterica*. **Conclusion:** The results conclude that hygienic and sanitary practices in smallholder dairy goat farming are still poor as shown by the presence of pathogenic bacteria.

Key words: Goat milk, microbiological quality, shiga toxin-producing E. coli, Salmonella

Citation: Ismiarti, Yustina Yuni Suranindyah and Widodo, 2019. Microbiological Qualities of goat milk obtained under different milking systems at a smallholder dairy farm in Yogyakarta, Indonesia. Int. J. Dairy Sci., 14: 29-35.

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Competing Interest: The authors have declared that no competing interest exists.

Data Availability: All relevant data are within the paper and its supporting information files.

INTRODUCTION

Among milk from other species, goat milk has special characteristics; it can be easily digested because its fat globules are smaller with more short chain fatty acids and it is less allergenic than other milk and safe to be consumed by people with lactose intolerance^{1,2}. Microbiological gualities of goat milk are determined by its composition and hygienic practices during milking and condition during storage and distribution³. Microbial contamination of milk can occur from direct contact between fresh milk and contaminated equipment surfaces during milking⁴. Improper handlings or storage of fresh milk can represent a transmission hazard for bacterial pathogens that responsible for foodborne illness⁴. A number of bacterial pathogens was previously reported to be responsible for milk-borne illness and it includes Shiga toxin-producing *Escherichia coli* (STEC)⁵ and *Salmonella* sp.⁶. Global milk-borne illness caused by STEC and Salmonella sp. has previously been reported⁶⁻⁸. In 2010, the estimated global diseases of STEC was 2,841,511 cases with 269 people were deaths9. Healthy dairy ruminants commonly carrying STEC in their feces¹⁰. Transmission of this bacterial pathogen human mainly occurs to through less-cooked meat, unpasteurized milk, feces-contaminated water and improper hygienic and sanitary practices during milking.

Smallholder dairy farms in Indonesia contribute 20.37% of milk needed domestically¹¹ of which, a small portion is from goat dairy farms. Generally, smallholder dairy farms are traditionally managed such as milking by hand with improper storage management¹². In many developing countries, poor sanitation during milking causes contamination that can affect consumer health and cause financial loss¹³. It is therefore, important to investigate microbiological quality of goat milk harvested from two different milking systems from smallholder dairy farm in Indonesia. The data obtained during this study will be used as a recommendation for better practices in managing smallholder dairy farm and improving goat milk qualities.

This study was conducted to evaluate microbiological qualities of goat milk based on total plate count (TPC), Enterobacteriaceae (EB) count and the presence of pathogens such as STEC and *Salmonella* in milk from a smallholder dairy goat farm managed under different milking systems. The TPC shows all living mesophilic microbes in goat milk that can grow as forming colonies on plate count agar (PCA) medium. The EB are indicators of fecal contamination and an

effective measurement for the environmental sanitary program, such as in powdered milk and ready-to-serve food¹⁴.

MATERIALS AND METHODS

Sampling of goat milk farms: Goat milk was sampled in January to April, 2018 from two goat farms in Yogyakarta Indonesia that applied different milking systems. In Group A, milking was carried out after cleaning around the stall. Milking in Group A was conducted by hand without cleaning the udders and dipping and farmers washed hands before milking. The first milk flow in Group A was discarded, the milk was collected in receptacle drinking-water bottles. Milk samples in Group A were collected from the bottles and moved to a sterile plastic container. In Group B, milking was carried out after cleaning around the stall and goats from colony stalls were moved to milking stalls. Farmers washed their hands before milking and the goat's udder was cleaned using warm water. The first milk flows were discarded. Milking was conducted using a bucket milking machine connected to tubes on a vacuum pump that fastens on teats, so milk flowed from tubes and was collected in a milk bucket. Samples were carried out from the milk bucket to a sterile plastic container. Samples from Groups A and B were put in a box with ice, which was quickly transported to the laboratory to be analyzed.

Microbiological analysis of goat milk: Microbiological qualities of goat milk were determined using three parameters, TPC to measure total amount of mesophilic aerobic microbes was conducted using plate count agar (PCA) media¹⁵ isolation of EB to measure enteric bacteria using violet red bile glucose (VRBG) media¹⁶, isolation of STEC¹⁷⁻¹⁹ and *Salmonella*¹⁹ to measure the presence of pathogen. Isolates found were identified using specific primers *stx*1 and *stx2* for STEC and 16S rRNA sequencing for *Salmonella* determination.

TPC: Goat milk samples were serially diluted from 10^{-1} - 10^{-3} and then plated onto PCA medium and incubated at 37° C for 24 h. Total colonies on PCA medium were counted¹⁵ as CFU mL⁻¹.

EB: Goat milk samples were serially diluted from 10^{-1} - 10^{-3} and then plated onto VRBG medium and incubated at 37°C for 24 h. Predictive EB colonies that were reddish or purplish in color were counted.

Pathogen isolation

STEC: Isolation of Escherichia coli was carried out using an enrichment broth medium of Brilliant Green Lactose Bile (BGLB) and incubated at 37°C for 24 h. A positive sample was detected with turbidity and then inoculated onto a selective medium, Eosin Methylene Blue (EMB) agar and incubated at 37°C for 24 h. A positive sample was detected when a colony appeared with a metallic sheen or dark chocolate color. Positive samples on EMB were then streaked onto a selective medium, Sorbitol MacConkey (SMAC) agar and incubated at 37°C for 24 h. A positive sample of STEC was detected as a colorless colony on the surface of SMAC medium. Predictive STEC colonies were inoculated onto Lactose Broth medium.

Salmonella: Isolation of *Salmonella* was carried out with pre-enrichment using buffered peptone water (BPW) and incubated at 37°C for 24 h. Turbid samples were enriched into rappaport-vassiliadis soy (RVS) broth and incubated at 37°C for 24 h. Turbid samples from the enrichment were then streaked onto xylose lysine deoxycholate (XLD) agar, hektoen enteric (HE) agar and bismuth sulfite agar (BSA) and incubated at 37°C for 24 h. Positive samples were pink with or without black spots on XLD, green or dark green with or without dark spots on HE and gray or dark gray on BSA. Positive isolates were tested for their biochemical characteristics using tryptic soy iron (TSI) agar and lysine iron agar (LIA). Positive isolates were then transferred into nutrient broth.

Molecular detection: One milliliter of culture of putative STEC and Salmonella isolates were transferred into DNA isolation kit (Favorgen[™]). Genomic DNA of STEC isolates was amplified using specific primers for stx1 (F: 5 -ATAAATCGCCATTCGTT GACTAC-3 and R: 5 - AGAACGCCCACTGAGATCATC-3) and stx2 (F: 5-GGCACTGTCTGAAACTGCTCC-3 and R: 5-TCGCCAGTTAT CTGACATTCTG-3). Genomic DNA of putative Salmonella isolates were amplified using universal primers of 16S rRNA [F: 5-AGA GTTTGAT(C/T)(A/C)TGGCTCAG-3 and R: 5-CA(G/T)AAAGGAGGTGATCC-3]. Amplification of STEC genomic DNA using *stx*1 primers has a 180 bp target product and stx2 has a 255 bp target product¹⁶. Amplification of Salmonella genomic DNA using universal 16S rRNA primers has a 1500 bp target product. Amplification was conducted using a thermal cycler 2720 (Applied Biosystems[™]). Amplicons produced were visualized using agarose gel electrophoresis. The PCR products were sequenced using ABI PRISM[™] 3730-XL 1406-022.

Table 1: The T	FPC of goat milk	from Groups A and B

Sample	Group A	Group B
codes	TPC (CFU g^{-1})	TPC (CFU g ⁻¹)
1	6.02×10 ⁷	5.75×10 ⁴
2	3.68×10 ⁷	1.34×10⁵
3	1.77×10 ⁷	2.56×10 ⁵
4	1.80×10 ⁷	1.19×10⁵
5	6.36×10 ⁷	7.95×10^{4}
6	5.99×10 ⁷	2.86×10 ⁵
7	5.40×10 ⁷	2.06×10 ⁵
8	2.50×10 ⁷	2.95×10⁵
9	3.99×10 ⁷	2.43×10 ⁵
10	2.93×10 ⁷	2.92×10 ⁵
Average	4.05×10 ^{7a}	1.97×10 ^{5b}

TPC: Total plate count. The number followed by different superscript means different number (p<0.05)

Table 2: The EB count in goat milk from Groups A and B

Group A	Group B
EB (CFU g ⁻¹)	EB (CFU g ⁻¹)
1.60×10 ⁴	4.90×10 ³
4.35×10 ⁴	1.79×10^{4}
2.30×10 ⁴	2.50×10^{3}
1.20×10 ⁴	7.90×10^{3}
5.10×10 ⁴	1.01×10^{4}
1.11×10 ⁵	1.60×10^{4}
2.80×10 ⁴	1.31×10^{4}
1.28×10 ⁵	5.50×10^{3}
1.36×10 ⁵	1.06×10^{4}
8.95×10 ⁴	2.95×10^{3}
6.38×10 ^{4a}	9.15×10 ^{3b}
	$\frac{\text{EB} (\text{CFU g}^{-1})}{1.60 \times 10^4}$ 4.35×10^4 2.30×10^4 1.20×10^4 5.10×10^4 1.11×10^5 2.80×10^4 1.28×10^5 1.36×10^5 8.95×10^4

EB: Enterobacteriaceae. The number followed by different superscript means different number (p<0.05)

RESULTS

Total plate count (TPC): The TPC of goat milk from Groups A and B is presented in Table 1. Table 1 shows that TPC was higher in Group A than in Group B (p<0.05). The average TPC in Groups A and B was 4.05×10^7 and 1.97×10^5 CFU g⁻¹, respectively. The highest amount of TPC in Group A and B were 6.36×10^7 and 2.95×10^5 CFU g⁻¹, respectively. The lowest were 1.77×10^7 and 5.75×10^4 CFU g⁻¹, respectively.

Enterobacteriaceae (EB): There was different amount of EB between Group A and B. Table 2 shows that the EB count was higher in Group A than in Group B (p<0.05). The average EB count in Groups A and B was 6.38×10^4 and 9.15×10^3 CFU g⁻¹, respectively. According to Standar Nasional Indonesia, the maximum total EB count in raw milk is 1×10^3 CFU mL⁻¹. A high EB count in the two groups indicated poor sanitation.

Shiga toxin-producing *Eschericia coli* (STEC) and *Salmonella*: Regarding the detection of STEC, all samples from both groups showed positive growth in BGLB and EMB, while samples grown in SMAC showed 8 positive growth.

Int. J. Dairy Sci., 14 (1): 29-35, 2019

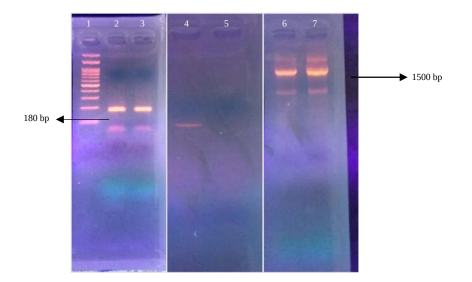


Fig. 1: Electrophenogram of amplified genomic DNA; 1: DNA ladder 100 bp, 2: Amplified DNA from Group A with primer *stx*1,3: Amplified DNA from Group B with primer *stx*1, 4: Unamplified DNA from Group A with primer *stx*2, 5: Unamplified DNA from Group B with *stx*2 primer, 6: Amplified DNA from Group A with 16S rRNA primer, 7: Amplified DNA from Group B with primer 16S rRNA

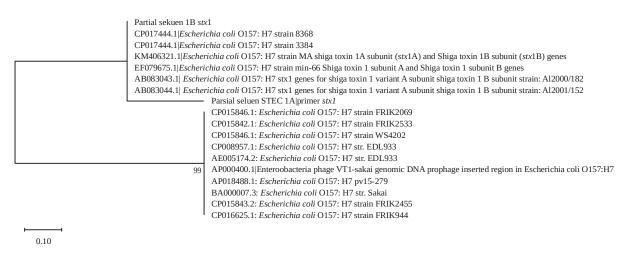


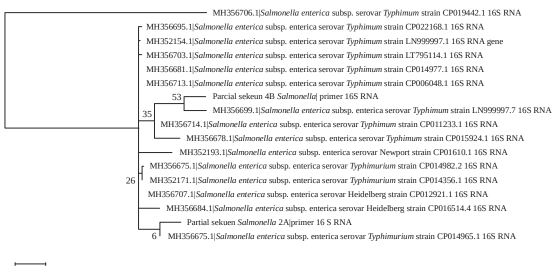
Fig. 2: Phylogenetic tree of partial sequence of STEC Groups A and B using primer *stx*1 Note: Sequence accession number provided before culture identity

Meanwhile, the detection of *Salmonella* in BPW and RVS showed all samples with positive growth. In Group A, 1 sample was positive in XLD, another 1 sample was positive in HE and 2 samples were positive in BSA. In Group B, no sample was positive in XLD, 3 samples were positive in HE and no sample was positive in BSA. Based on biochemical tests, all putative *Salmonella* isolates were positive in TSI and all isolates were negative in LIA.

Molecular detection of STEC and *Salmonella* was carried out by PCR and the products were visualized using 2% agarose gel electrophoresis (Fig. 1). Figure 1 shows that

amplification with *stx*1 primers resulted in a 180 bp product; however, *stx*2 primers had no amplification products, indicating that these primers did not recognize the corresponding sequence of the STEC genomic DNA. BLASTN analysis showed that amplified DNA using *stx*1 primers in Groups A and B had more than 90% similarity with several identified sequences as *E. coli* O157:H7. Further analysis using a phylogenetic tree showed that amplified sequences in Groups A and B were aligned with the sequence of *E. coli* O157:H7 strain Shiga toxin subunit 1A (*stx*1A) and 1B (*stx*1B) (Fig. 2).

Int. J. Dairy Sci., 14 (1): 29-35, 2019



0.020

Fig. 3: Phylogenetic three of partial sequence of *Salmonella* Groups A and B using primer 16s rRNA Note: Sequence accession number is provided before culture identity

Genomic DNA of putative *Salmonella* species was amplified using 16S RNA primers and resulted in a 1500 bp product (Fig. 1). The BLASTN analysis showed that amplified DNA using 16S RNA primer of putative *Salmonella* in Groups A and B had more than 90% similarity with several sequences of *S. enterica* (Fig. 3).

DISCUSSION

Different milking systems between Group A and B had different amount of TPC and EB. Based on Standar Nasional Indonesia, good-quality milk has TPC <1 \times 10⁶ CFU mL⁻¹, hence, goat milk from Group A did not meet this standard²⁰. The higher TPC in Group A was due to low hygienic and sanitation practices, such as no cleaning of the udder and teats before milking. Improper hygienic practices trigger microbial contamination during milking. Mohammadi et al.8 reported that milk quality is determined by its composition and hygienic practices that are applied during milking processes, such as cleanliness of milking equipment, conditions of storage and transportation and cleanliness of the udder of the individual goat. Suranindyah et al.¹² also reported that improving environmental sanitation during milking and dipping of teats can reduce total microbes in raw milk. This is in line with the present study results. Smallholder dairy farms with better sanitation produce raw milk with lower total microbes. The TPC in raw milk indicated growth of total microbes¹⁴.

The lower EB in Group B was due to better hygienic practices such as conducting milking in a milking stall and cleaning the udder before milking. In Group A, milking was conducted in an individual stall without cleaning the udder before milking. This finding was in line with a study conducted by Kyozaire *et al.*²¹, which reported that milking with a bucket-system milking machine produces milk with the lowest TPC and Coliform, compared with manual milking and pipeline system milking. Good handling practices, especially for fresh milk are essential in controlling microbial contamination before (internal) and after (external) milking²². Hence, improving sanitation is important to reduce EB contamination in milk. According to Farrokh et al.¹⁰, poor hygiene and sanitation of the milking system caused STEC from feces to adhere in teats, leading to contamination during milking.

The presence of pathogenic *Salmonella* sp., indicated that the stall environment had low hygiene and sanitation. This brings the possibility of *Salmonella* sp., to transfer from feces to the fresh milk harvested through air, equipment and during milking. The presence of *Salmonella* in fresh milk leads to food-borne disease known as Salmonellosis²³. Several *S. enterica* isolates from warm-blooded animals were known to cause fever, gastroenteritis, bacteremia and other symptoms^{23,24}. Based on these findings, better hygienic and sanitary practices have to be applied in dairy farms to prevent pathogenic transmission from ruminants to humans. The presence of STEC and *Salmonella* sp. in goat milk obtained from smallholder farms in Indonesia confirms the necessity of milk processing, such as pasteurization, before consumption.

CONCLUSION

Goat milk obtained in a milking system with better hygiene and sanitation results in a better microbiological qualities. However, the presence of STEC, *Salmonella* sp. and high EB count shows the hygiene and sanitation during milking is not good enough to prevent pathogenic contamination and needs to be improved.

SIGNIFICANCE STATEMENT

This study discovers that pathogenic bacteria were present in goat milk obtained under poor hygienic and sanitation system in smallholder dairy farms. This study recommend a better milking systems with better hygienic and sanitation condition have to be applied in smallholder dairy farms in Indonesia to avoid milk-borne illness.

ACKNOWLEDGMENT

Ismiarti was a recipient of the Indonesian Endowment Fund for Education from the Ministry of Finance, Republic of Indonesia.

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