Isolation and Identification of Biofilm-Forming *Staphylococcus Aureus* in Commercial Cow Milk Products

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Abstract

Food poisoning caused by the contamination from *Staphylococcus aureus* are frequently found in food especially in dairy products. Pasteurization process in milk production was not enough to kill *S. aureus* because it formed biofilm that could survive in high temperature. This research aimed to study the presence of biofilm-forming *S. aureus* in samples from packed commercial milk products in Yogyakarta City, Indonesia. Twenty isolates from dairy products were grown in Brain Heart Infusion (BHI) broth then inoculated into Braid-Parker Agar (BPA) medium to get the candidate of *S. aureus* isolates. These isolate candidates were selected using Mannitol Salt Agar (MSA) and Congo Red Agar (CRA) medium. Another selection was done by carbohydrate fermentation analysis and confirmed using API STAPH. Confirmation analysis showed that eight isolates were identified as *S. aureus*. Another two isolates were identified as *S. xylosus* and *S. haemolyticus*. Therefore, it indicated the presence of *Staphylococcus aureus* as contaminant in dairy products.

Keywords: *Staphylococcus aureus*, packed milk product, biofilm, API STAPH

Introduction

*Staphylococcus aureus* is pathogenic bacteria that easily transferred through food and caused foodborne illness. This phenomenon is globally important. *S. aureus* affected the human and animal health. It infected human through the consumption of contaminated food. In China, 53.7% of food poisoning cases were caused by *S. aureus* in 2015 (Wu et al., 2018). The highest cases of foodborne diseases in United States were reported caused by *S. aureus* with 241,000 cases per year (Kadariya et al., 2014). Dairy milk product was one of the main sources of food poisoning caused by *Staphylococcus*. The bacteria could survive during the pasteurization process and produced enterotoxin that can be involved in the product (Johler et al., 2015; Jin and Yamada, 2016). Isolates of *S. aureus* from pasteurized milk products in China were reported to have the ability to produce biofilm (96.7%) and 66.7% isolates had the virulence factor to cause diseases (Dai, 2019). Castelini (2014) reported that *S. aureus* produced *Staphylococcus* enterotoxin (SEs) in food and if it consumed, it can lead to high fever and vomiting with or without diarrhea and nausea in less than 8 h (between 3 and 4 h).

Research conducted by Qian et al. (2019), showed that from 289 samples collected from food poisoning cases because of goat milk consumption in Shaanxi, China, 68 isolates of *S. aureus* were found and 91.8% had a close relation with its ability to produced biofilm with antibiotic-resistance characteristic. Biofilm formation made *S. aureus* has the ability to colonize, resistance to antibiotic with high level of virulent characteristic (Rohinishree, 2011). In Yogyakarta City, packed-pasteurized dairy milk products are one of the favorite drinks for the society, especially students because of its high nutrition content, relative cheap price, simplicity and easiness to be carried everywhere. The aim of this research was to find the presence of biofilm-forming *Staphylococcus* in packed commercial dairy milk products in Yogyakarta City, Indonesia.

Materials and Methods

Materials

The samples were 20 packs of cow milk products from 5 kinds of milk with different brands that have been randomly collected from shops and supermarkets in Yogyakarta.
City, Indonesia. Brain Heart Infusion (BHI) broth, Baird-Parker Agar (BPA), Mannitol Salt Agar (MSA), Brain Heart Infusion Agar (BHIA), Congo Red Agar (CRA) and peptone water were purchased from Millipore Corp. (Massachusetts, USA). API STAPH was purchased from bioMerieux Company (Marcy-l’Étoile, France). All chemicals and reagents were of analytical grade.

Methods
Isolation step

Each sample (10 ml) was taken and grown in 90 mL BHI broth media then incubated for 16-18 h (Palilu & Budiarso, 2017). Incubated sample (1 ml) was then diluted using 0.1% peptone water (9 ml) into solution with the concentration from $10^{-1}$ to $10^6$ and homogenized using vortex. Diluted sample (0.1 ml) from the concentration of $10^{-4}$, $10^{-5}$ and $10^{-6}$ were inoculated to the surface of BPA medium and spread using drygalsski and incubated for 48 h at 37 °C. Suspected colonies of *Staphylococcus* expressed in dark grey to shining black color in BPA medium (Sutejo et al., 2017). Purification was done for these suspected colonies by taking the separate colonies and inoculating them in BPA medium using streak plate technique to get the single colony. This suspected *Staphyloccous* single colony then streaked into MSA medium and incubated for 24 h. Colony that expressed in yellow color was then separated into another BPA medium to get the single isolate. This isolate was finally grown in BHIA medium and collected as *S. aureus* suspected isolate (Olwal et al., 2018).

Selection of Biofilm-forming *Staphylococcus aureus*

Single suspected isolate of *S. aureus* from BHIA medium was inoculated into the surface of CRA medium by streak plate technique then incubated for 24 h at 37 °C. *Staphylococcus aureus* that produced slime were expressed in black colony. Congo Red Agar medium contains of BHI broth (37 g/L), agar base (10 g/L), saccharose (36 g/L) and Congo Red dye (0.8 g/L) as the biofilm-forming indicator (Bryan, 2017; Casagrande Proietti et al., 2015).

Biochemistry Confirmation Analysis using API STAPH

According to the reference method from Savage et al. (2017) and Vanderhaeghen et al. (2015), there were three steps in confirmation analysis including preparation the API STAPH stripe, inoculum preparation and inoculation process. Incubation process was done for 18-24 h in 37 °C and the observation was conducted for 24 h by adding NIT 1 and NIT 2 reagents into NIT well, Zym A and Zym B reagents into PAL well and VP1 and VP2 reagents into VP well. Color changing was observed and identified using APIWEB software version 1.3.0 (bioMerieux Company, Marcy-l’Étoile, France).

Result
Isolation and Selection of *Staphylococcus aureus*

*Staphylococcus aureus* was isolated from 20 kinds of different cow milk product. Isolation was done by inoculated the culture into BPA medium by adding egg yolk tellurite. The presence indicator of suspected *S. aureus* in the milk product was the black color in the colony with clear zone surrounding it (Figure 1.) The suspected
different kind of carbohydrate sources before confirmation analysis using API STAPH. Carbohydrate fermentation analysis (Table 1.) was done to test the bacteria ability to ferment different kinds of sugar.

Isolates with predicted Staphylococcus aureus Table 1. then analyzed for confirmation analysis using API STAPH. Isolates with different predicted species of Staphylococcus were taken (one isolate per species) for confirmation analysis and the results shows in Table 2. The medium in this analysis consisted of 20 sugar mediums to test the homogeneous suspension of bacteria using McFarland 0.5 standard to see its level of turbidity. Medium containing bacterial suspension was then incubated at 37°C for 18-24 h. The metabolism process produced the color’s changing even after the addition of reagent into the NIT, PAL and VP medium (Langlois et al., 1983).

The suspension from API STAPH from NIT, PAL and VP medium after addition of reagent showed the positive changing color (Fig. 3.) with all medium changed from red into yellow, and ADH and URE from yellow into pink.
Discussion

Suspected colony of *Staphylococcus aureus* in this research was able to appear as black color colony surrounded by clear zone when inoculated in BPA medium. This happened because there was lipolytic activity from *S. aureus* that reduced the tellurite into tellurium. The opaque zone caused by the proteolytic and lipolytic processes with the additional of egg yolk (Capita *et al*., 2001). The growth of *S. aureus* in BPA medium shows that this bacteria still presence in the milk product even though passing through the pasteurization process during the production. These suspected colony then regrowth in MSA medium. The growth of many species of bacteria, except *Staphylococcus*, was inhibited by 7.5 % of sodium chloride. Mannitol is the carbohydrate source that can be fermented by *S. aureus* in MSA medium therefore, it produced yellow color colony in the end of the incubation time. The coagulate protein of negative species of *Staphylococci* and *Micrococci* did not fermented mannitol hence, growth as small red color colony (Pumipuntu *et al*., 2017).

Gram staining technique was done in this research to differentiate the group of positive and negative bacteria. *Staphylococcus aureus* is gram positive bacteria. Violet crystal colored cells will show the shape and color of *S. aureus*. This procedure produced the purple iodine area in the bacteria cytoplasm. Previous cells colored with violet crystal and iodine were then added by the mixture of acetone and alcohol to wash away the stain. The difference between gram-positive and gram-negative bacteria was its cell wall permeability (Elsa *et al*., 2010).

The isolate that has been analyzed for its carbohydrate fermentation was then analyze for its biofilm-forming indicator. Biofilm is a group of microbia cell that is irreversibly and related with the surface and close off matrix that mostly contains polysaccharide (Kim and Han, 2014). Ten isolates from different samples formed biofilm. Black and red colony in the surface of media showed the activity of *S. aureus*. Forming the biofilm is the survival mechanism of *S. aureus*. Black colony produced slime thus, indicated that biofilm is formed by *Staphylococcus*. Slime is produced because of the activity of fermentation enzyme (poly-N-acetylglucosamine; PNAG) that indicated by the appearance of black color in CRA medium. The use of high temperature in production process aimed to kill the opportunist bacteria. Not all of them were killed because of the biofilm formed by bacteria. The strain of bacteria that can
produced slime or biofilm could increase its cell ability to survive from the process involved high temperature (Nermati et al., 2009).

Carbohydrate fermentation analysis was carried out using many kinds of sugar as its carbon source, there were xylose, maltose, sucrose, galactose, mannitol and Voges-Proskauer (VP) analysis. The purpose of this analysis was to convince that the suspected isolates were \textit{S. aureus}. The changing color in the medium showed the forming of acid as the product from fermentation process by bacteria (Ajayi et al., 2017). The result of fermentation analysis showed that there were 6 suspected isolates of \textit{S. aureus}. Those isolates were confirmed using API STAPH analysis. According to Table 1., isolate with code S.B.4.3 was able to ferment xylose but not VP. Thus, it was predicted as \textit{Staphylococcus gallinarum}. Suspected \textit{S. aureus} isolates were able to ferment maltose, sucrose, galactose and mannitol. Other isolates were predicted as \textit{S. saprophyticus}, \textit{S. epidermidis} and \textit{S. lugunensis}. These results need confirmation since it was still prediction. All of the isolates predicted for \textit{S. aureus} were confirmed with API STAPH, while another isolate (\textit{S. saprophyticus}, \textit{S. epidermidis} and \textit{S. lugunensis}) was taken one isolate per species as the representative.

The confirmation analysis using API STAPH was conducted for biochemistry analysis for \textit{Staphylococcus}, \textit{Micrococcus} and \textit{Kocuria}. The result from Table 2. shows that the highest ID percentage was isolate S.K.1.1 from sweetened condensed milk sample. The ID percentage of 98% was confirmed as \textit{S. xylosus}. Some suspected isolates from carbohydrate fermentation showed the positive identification as \textit{S. aureus}. These results showed that \textit{S. aureus} can still survive in dairy milk product even after passed through high temperature during pasteurization process. Isolate S.U.1.1 and S.U.3.4 confirmed as \textit{S. haemolyticus} with ID percentage of 85.0% and \textit{S. xylosus} with ID percentage of 98.7%, respectively. The ability of predicted \textit{Staphylococcus} to ferment sugar medium was different. Not all the predicted \textit{S. aureus} in Table 1. ferment all the sugar medium. It can be seen in Figure 3. that there was different reaction indicated by the changing color after incubated for 18-24 h. All medium was incubated in aerobic condition, except for arginine dihydrolase (ADH) and URE medium were incubated in anaerobic condition. Fermentation results showed that the medium color in well changed from red into yellow and for ADH and URE changed from yellow into pink. The result with 92.5% percentage of ID was identified as \textit{S. aureus} with the rest (3% ID) was \textit{S. haemolyticus}. Thus, it supported the different results after confirmation analysis conducted.

**Conclusion**

There were 8 isolates identified as \textit{S. aureus} with the percentage ID up to 92% and were able to form biofilm. Another isolates identified as \textit{S. xylosus} and \textit{S. haemolyticus}, with the percentage ID of 98.7% and 85%, respectively.

**References**

Adetutu, A.A., Oritsewehinmi, B., Ikhiiwili, O.M., Moradeke, A.O., Odochi, A.S., & Adeola, O.A. (2017). Studies on \textit{Staphylococcus aureus} isolated from pimples. Pakistan. Journal of Biological Science, 20(7), 350-354.

Argudín, M.Á., Mendoza, M.C., & Rodicio, M.R. (2010). Food poisoning and \textit{Staphylococcus aureus} enterotoxins. Toxins, 2(7), 1751-1773. Doi:10.3390/toxins2071751

Bryan, J. (2013). Detection of Adhesion Genes and Slim Production among \textit{Staphylococcus aureus} and \textit{Staphylococcus epidermidis} Isolated from Hemodialysis Patients. College of Medicine, University of Babylon. Hilla, Iraq

Brooks, G.F., Carroll, K.C., Butel, J.S., Morse, S.A., & Mietzner, T.A. (2013). Mikrobiologi Kedokteran Ed.23, Translation of Jawetz, Melnick, and Adelberg’s Medical Microbiology, 23th. McGraw-Hill, US

Casagrande, P.P., Stefanetti, V., Hyatt, D.R., Marenzoni, M.L., Capomaccio, S., Coletti, M., & Passamonti, F. (2015). Phenotypic and genotypic characterization of canine...
pyoderma isolates of *Staphylococcus pseudintermedius* for biofilm formation. *Journal of Veterinary Medical Science*, 77(8), 945–951. doi:10.1292/jvms.15-0043

Capita, R., Alonso-Calleja, C., Moreno, B., & Garcia-Fernandez, M.C. (2001). Occurrence of *Listeria* species in retail poultry meat and comparison of a cultural/immunoassay for their detection. *The Journal of Food Microbiology*, 39, 321-325

Dai, J., Wu, S., Huang, J., Wu, Q., Zhang, F., Zhang, J., Wang, J., Ding, Y., Zhang, S., Yang X., Lei, T., Xue, L., and Wu, H. (2018). Prevalence and characterization of *Staphylococcus aureus* isolated from pasteurized milk in China. *Front. Microbiol*. 10, 641. doi: 10.3389/fmicb.2019.00641

Elsa, H.W., Maria, L.R., Tereza, C.R., Luciane, B.M., Lina, C.A., & Vanerli B. (2010). Assessment of the risk of raw milk consumption related to *Staphylococcal* food poisoning. Accessed 27 March 2015

Jin, W., & Yamada, K. (2016). *Staphylococcal enterotoxins* in processed dairy products. *Food Hygiene and Toxicology in Ready-to-Eat Foods*, 241–258. doi:10.1016/b978-0-12-801916-0.00014-5

Johler, S., Weder, D., Bridy, C., Huguenin, M.-C., Robert, L., Hummerjohann, J., & Stephan, R. (2015). Outbreak of *staphylococcal* food poisoning among children and staff at a Swiss boarding school due to soft cheese made from raw milk. *Journal of Dairy Science*, 98(5), 2944–2948. doi:10.3168/jds.2014-9123

Kadariya, J., Smith, T. C., & Thapaliya, D. (2014). *Staphylococcus aureus* and *Staphylococcal* Food-Borne Disease: An Ongoing Challenge in Public Health. *BioMed Research International*, 2014, 1–9. doi: 10.1155/2014/827965

Kim, M & Han, M. (2014). Characteristics of biofilm development in an operating rainwater storage tank. *Environmental Earth Sciences*, 72(5), 1633–1642. https://doi.org/10.1007/s12665-014-3067-2

Kumar, A., Alam, A., Rani, M., Ehtesham, N. Z., & Hasnain, S. E. (2017). Biofilms: Survival and defense strategy for pathogens. *International Journal of Medical Microbiology*, 307(8), 481–489. https://doi.org/10.1016/j.ijmm.2017.09.016

Langlois, B.E., Harmon, R.J., & Akers, K. (1983). Identification of *Staphylococcus* species of bovine origin with the API Staph-Ident system. *Journal of Clinical Microbiology*, 18(5), 1212–1219.

Nemati, M., Hermans, K., Devriese, L.A., Maes, D., & Haesebrouck, F. (2009). Screening of genes encoding adhesion factors and biofilm formation in *Staphylococcus aureus* isolates from poultry. *Avian Pathology*, 38(6), 513–517. https://doi.org/10.1080/03079450903349212

Olwal, C. O., Ang’lenda, P. O., Onyango, D. M., & Ochieng, D. O. (2018). Susceptibility patterns and the role of extracellular DNA in *Staphylococcus epidermidis* biofilm resistance to physico-chemical stress exposure. *BMC Microbiology*, 18(1), 1–13. https://doi.org/10.1186/s12866-018-1183-y

Palilu, P. T & Budiario, T. Y. (2017). Isolation and identification of *Staphylococcus sp.* in powdered infant milk. AIP Conference Proceedings 1844, 020016 (2017); doi: 10.1063/1.4983427

Pumipuntu, N., Kulpeanpraisit, S., Santajit, S., Tunyong, W., Kong-ngoen, T., Hinthong, W., & Indrawattana, N. (2017) Screening method for *Staphylococcus aureus* identification in subclinical bovine mastitis from dairy farms. *Veterinary World*, 10(7), 721-726.

Qian, W., Shen, L., Li, X., Wang, T., Liu, M., Wang, W., & Zeng, Q. (2019) Epidemiological Characteristics of *Staphylococcus aureus* in Raw Goat Milk in Shaanxi Province, China. *Antibiotics*, 8(3), 141. doi:10.3390/antibiotics8030141

Rohinishree, Y & Negi, P.S. (2011). Detection, Identification and Characterization of *Staphylococci* in Street Vend Foods. *Food and Nutrition*

Scallan, E., Hoekstra, R. M., Angulo, F. J., Robert, V. T., Marc-Alain, W., Sharon, L. R., Jeffrey, L. J., & Patricia, M.G.(2011). Foodborne illness acquired
in the United States—major pathogens. *Emerging Infectious Diseases*. 17, 7–15. doi: 10.3201/eid1701.P11101.

Savage, E., Chothe, S., Lintner, V., Pierre, T., Matthews, T., Kariyawasam, S., & Jayarao, B. (2017) Evaluation of Three Bacterial Identification Systems for Species Identification of *Bacteria* Isolated from Bovine Mastitis and Bulk Tank Milk Samples. *Foodborne Pathogens and Disease*, 14(3), 177–187. doi:10.1089/ fpd.2016.2222

Sutejo, S.V. H., Amarantini, C. & Budiarto, T.Y. (2017) Molecular detection of *Staphylococcus aureus* resistant to temperature in milk and its products. *AIP Conference Proceedings* 1908, 050007 (2017) https://doi.org/10.1063/1.5012731

Vanderhaeghen, W., Piepers, S., Leroy, F., Van Coillie, E., Haesebrouck, F., & De Vliegher, S. (2015). Identification, typing, ecology and epidemiology of coagulase negative *Staphylococci* associated with ruminants. *The Veterinary Journal*, 203(1), 44–51. doi:10.1016/j.tvjl.2014.11.001

Wu, S., Huang, J., Wu, Q., Zhang, F., Zhang, J., Lei, T., & Xue, L. (2018). Prevalence and Characterization of *Staphylococcus aureus* Isolated from Retail Vegetables in China. *Frontiers in Microbiology*, 9. doi:10.3389/fmicb.2018.01263