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CONTEXT

Dadih is a traditional Indonesian fermented buffalo milk drink with numerous health benefits associated with several probiotic strains. The current process of dadih production is still mostly artisanal and does not rely on good hygiene practices. The Dadih Initiative was started in Indonesia in 2017 to standardize and optimize production and quality control of dadih. However, before expanding the production of dadih, additional insights into the nature of microorganisms present in the final product are needed. With the advent of next generation sequencing technologies (NGS), it is now possible to explore the microbiota composition of dadih samples with an unprecedented level of complexity. In this work, the microbiota composition of artisanal dadih samples was determined using 16S-rRNA-gene amplicon-sequencing of the V3-V4 region. Several sites of collection and production methods were evaluated, including samples made from pasteurized or raw milk, and using back-slopping (a practice which involves using previous fermentation as a starter culture) or not.

MATERIALS

- Precellys 24 tissue homogenizer (Bertin Technologies, Montigny-le-Bretonneux, France)
- Quick-DNA™ Fecal/Soil Microbe Miniprep Kit (Zymo Research, Irvine, CA)





MICROBIOTA ANALYSIS OF FERMENTED MILK SAMPLES WITH NEXT GENERATION SEQUENCING

PROTOCOL & RESULTS

- Dadih samples were collected from two different regions in Bukittinggi city (Gadut 2 separate producers and Palupuh), and in Padang Panjang town. There were differences in back-slopping practice, where previous fermentations were used as a starter culture (five samples) or not (three samples). **Table 1** gives an overview of the different samples obtained. Duplicate samples were stored frozen.
- DNA extraction: the samples were centrifuged at 14 500 g and subsequent DNA extraction from the pellet was performed using the Quick-DNA™ Fecal/Soil Microbe Miniprep Kit (Zymo Research, Irvine, CA) according to manufacturer's instructions, and the Precellys 24 tissue homogenizer (Bertin Technologies, Montigny-le-Bretonneux, France), with 3 cycles of 30s at 6500 rpm, with 5 min cooling on ice in between.
- PCR-amplifying the V3–V4 region of the 16S rRNA gene and next generation sequencing: Illumina 16S rRNA gene amplicon libraries were generated and sequenced at BaseClear (Leiden, the Netherlands). In short, barcoded amplicons from the V3–V4 region of 16S rRNA genes were generated using a 2-step PCR.
- Sequence processing and analyses: The sequencing run was analysed with the Illumina CASAVA pipeline (v1.8.3) with demultiplexing based on sample-specific barcodes. Results can be found in Figure 1.

	Site of collection	Pasteurized	Back slopping	Note
A	Gadut	Yes	Yes	Producer 1
В	Gadut	No	Yes	Producer 1
С	Palupuh	No	Yes	
D	Palupuh	Yes	No	
E	Padang Panjang	No	No	
F	Padang Panjang	Yes	No	
G	Gadut	Yes	Yes	Producer 2
н	Gadut	No	Yes	Producer 2

Table 1: Origin and characteristics of the collected dadih samples. From (1)

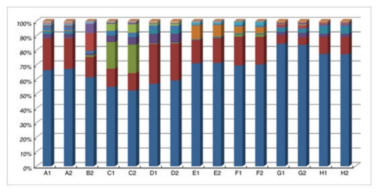


Figure 1: Relative abundance of major OTUs (at least 0·1% in any of the samples) in the duplicate dadih samples (coded according to Table 1). Legend for microbial OTUs (at genus level, or higher if not available at genus level, indicated by f_: family, o_: order. When the same names are given, these are still different OTUs): Lactococcus; Klebsiella; f_Bifidobacteriaceae; Leuconostoc; f_Enterobacteriaceae; Streptococcus; f_Enterobacteriaceae; f_Lactobacillaceae; Lactobacillus; f_Lactobacillaceae Acinetobacter; f_ Oxalobacteraceae; Vagococcus; Acetobacter; Proteus; f_Enterococcaceae; o_Lactobacillales, and Corynebacterium. From (1)





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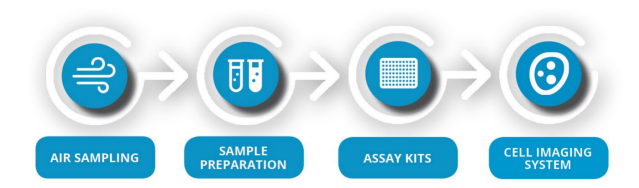
CONCLUSION

NGS results show that Lactococcus (52–83%) predominated in all samples, followed by Klebsiella (5–26%), and Lactobacillaceae, Bifidobacterium, Streptococcus, and Leuconostoc . Back-slopping practice was significantly correlated with higher levels of Lactobacillaceae, Pediococcus, species of the order Burkholderiales, and Serratia, but with lower levels of other Enterobacteriaceae (including Klebsiella), Streptococcaceae, Staphylococcus, and Brachybacterium. Taken together, fermentation results vary significantly from producer to producer and back-slopping practice should be recommended. These findings are instrumental in implementing future good manufacturing practices to produce a safe and healthy yoghurt-like dadih. The Precellys homogenizer is the ideal instrument for food quality control workflows, giving access to high-yield, high-quality nucleic acids suitable for Next Generation Sequencing (NGS) analysis.



(1) VENEMA, Koen et SURONO, Ingrid. S. Microbiota composition of dadih–a traditional fermented buffalo milk of West Sumatra. Letters in applied microbiology, 2019, vol. 68, no 3, p. 234-240.

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