SHORT COMMUNICATION

Diversity of Amylase-Producing *Bacillus* spp. from "Tape" (Fermented Cassava)

TATI BARUS*, AMANDA KRISTANI, ADI YULANDI

Department of Biology, Faculty of Biotechnology, Atma Jaya Catholic University, Jalan Jenderal Sudirman No. 51, Jakarta 12930, Indonesia

Received March 13, 2013/Accepted May 27, 2013

Fermented cassava or "Tape" is one of traditional Indonesian fermented food. The quality of "Tape" is determined by microorganisms involved during fermentation process. It was reported that *Bacillus subtilis* determined the quality of cassava "Tape". The most common way to identify species is by using 16S rRNA gene. This gene contains conserved regions as unique sequence which is relative among species. It has been widely used as a reliable molecular marker for phylogeny identification. Therefore, the aim of this research was to study diversity of amylase-producing *Bacillus* spp. from "Tape" based on 16S rRNA gene sequences. *Bacillus* spp. were isolated from "Tape" from several area in Indonesia i.e. Jakarta, Bandung, Cianjur, Subang, Rangkas Bitung, and Kediri. Amplification of 16S rRNA gene used 63f and 1387r primers. This research showed that based on 16S rRNA gene sequences, twenty-six of amylase-producing *Bacillus* spp. isolates were divided into four groups. All isolates were identified as species either *B. megaterium, B. subtilis, B. amyloliquefaciens*, or *B. thuringiensis*.

Keywords: Bacillus sp., cassava fermentation, diversity, amylase

INTRODUCTION

Most people in the world consume fermented foods. Many fermented foods contain of ingredients that are good for health such as stimulating intestinal immunity and improving the balance of microbial population in gastrointestinal tract. Therefore, fermented foods will become even more important in our diet for maintaining the health (Farnworth 2003).

"Tape" is one of the famous fermented food from Indonesia. It is made from steamed cassava (Manihot utilissima), then mixed with starter commonly referred as "Ragi Tape" (Barus & Wijaya 2011). "Tape" is produced using traditional methods which have some drawbacks, such as not standardized manufacturing processes and the products (Pawiroharsono 2007). This could be resulted from the inconsistency of microbial composition in starter, also the influence of environmental factors. The quality of "Tape" depends on quality of cassava, preparation method, and microbes. Starter of "Tape" comprise a consortium of microbes consist of molds, yeasts and bacteria. These microbes will determine the quality of "Tape" due to their role during fermentation process.

The role of *Bacillus* spp. in improving quality of fermented food was reported, such as in Indian kinema (Sarkar *et al.* 2002), Korean cheonggukjang (Kwon *et al.* 2009), African dawadawa (Terlabie *et al.* 2006) and soumbala (Sarkar *et al.* 2002). It was also reported that *Bacillus subtilis* determined the quality of cassava "Tape" (Barus & Wijaya 2011). However, there is very limited of information on diversity of *Bacillus* strains present in conventional prepared "Tape".

A range of molecular biological approaches has been applied to study genetic diversity of microbes, predominantly based on the analysis of 16S rDNA genes. Therefore, this study aimed to study the diversity of amylase-producing *Bacillus* spp. from cassava "Tape" based on 16S rRNA gene sequences. The results will be used as basis for further analysis of the role of *Bacillus* strain in determining the quality of the "Tape".

MATERIALS AND METHODS

Isolation of *Bacillus* spp. Genome. Twentysix of amylase-producing *Bacillus* spp. were isolated

^{*}Corresponding author. Phone: +62-21-5703306 Fax: +62-21-5719060, E-mail: tati.barus@atmajaya.ac.id

from "Tape". The samples of "Tape" were obtained from several area in Indonesia i.e. Pasar Bendungan Hilir-Jakarta, Pasar Kopro-Jakarta, Pasar Kelapa primer, and 4 μl DNA te

from several area in Indonesia i.e. Pasar Bendungan Hilir-Jakarta, Pasar Kopro-Jakarta, Pasar Kelapa Gading-Jakarta, Pasar Petukangan-Jakarta, Pasar Kebon Jeruk-Jakarta, Bandung, Cianjur, Subang, Rangkas Bitung, and Kediri. Amylase activity test was done according to the method of Oguntoyinbo *et al.* (2006). Bacterial cultures were grown overnight at 30 °C in 50 ml of Luria-Bertani broth. Cells were recovered by centrifugation at 13,000 × g for 3 min. Cell pellet was resuspended in1 mL of 10 mM Tris– HCl, pH 8.0, 10 mM EDTA, 100 mM NaCl, 2% (w/ v) SDS. Genomic DNA was isolated using *Fermentas*® *Genomic DNA Purification Kit* (Fermentas, Lithuania) based on the manufacturer's protocol.

Amplification and DNA Sequencing of 16S rRNA Gene. Amplification of 16S rRNA genes sequence of *Bacillus* spp. was performed in GeneAmp® PCR System 2700 (Applied Biosystems, Carlsbad, CA, USA) using the universal primers, comprises the forward primer 63F (5'-CAGGCCTAA CACATGCAAGTC-3') and the reverse primer 1387R (5'-CAGGCCTAACACATGCAAGTC- 3') (Marchesi *et al.* 1998). The primers were targeted to conserved regions and permitted the amplification of an aproximately 1,300 bp rDNA fragment. PCR master mix (50 μl) contained 25 μl *GoTaq Green* (Promega, Madison, USA), 17 µl Nuclease Free Water (Promega, Madison, USA), 2 µl of each primer, and 4 µl DNA template [± 100 ng]. PCR conditions were as follows: pre-denaturation at 95 °C for 5 minutes was followed by 30 cycle of denaturation at 95 °C for 1 minute, annealing at 58 °C for 5 minutes, extension at 72 °C for 1 minute, and post extension at 72 °C for 10 minutes. PCR products were observed using 1% electrophoresis agarose gel (Promega, Madison, USA) then stained with ethidium bromide (Sigma-Aldrich, USA). UV transilluminator was routinely used to visualize DNA in gel electrophoresis. PCR products were then partially sequenced in Macrogen Inc., Republic of Korea. The DNA sequencing results were aligned with 16S rRNA genes sequence database provided by GenBank (www.ncbi.nlm.nih.gov) using Basic Local Alignment Search Tool (BLAST) (http:// blast.ncbi.nlm.nih.gov/Blast.cgi). Phylogenetic tree was constructed using MEGA 5 software (Tamura et al. 2011). Neighbour joining method was used to develop phylogenetic tree.

RESULTS

A total twenty-six of amylase-producing *Bacillus* spp. were obtained from "Tape". The *Bacillus* spp. have diverse amylase activities with range from 3.40

Maximum Accession Amylase activity Sampel origin Isolate code Isolate code/homology identity(%) number (units/ml) B. amyloliquefaciens strain SWI4b Bandung Bdg 1-163f 99 JX203228.1 24.18 Bdg 1-2163f B. amyloliquefaciens strain BVC13 96 JQ660596.1 22.36 Cianjur Cjr 1163f B.amyloliquefaciens strain NBRC 15535 99 NR_041455 28.55 Cjr 4163f B. amyloliquefaciens isolate 003114 98 NR_027552 11.36 Cjr 5163f 96 HQ204325.1 B. subtilis strain M34 31.04 97 HQ336298.1 Jakarta Timur Pkg 2163f B. thuringiensis strain Bi51 23.57 Pkg 3163f B.thuringiensis strain IAM 12077 98 NR_043403 16.28 Pkg 4163f B. subtilis subsp. subtilis strain DSM 10 100 NR_027552 24.97 Pkj 2163f B.subtilis strain CCGE2066 EU867368.1 96 21.63 Pkj 3163f B. subtilis subsp. subtilis strain DSM 10 98 NR_027552 24.48 Jakarta Pusat Psb 1-1163f B. megaterium strain IAM 13418 99 NR_043401 29.16 Psb 1-2163f 99 KC441855.1 14.70 B. amyloliquefaciens strain W49 Psb 2-1163f 99 KC441855.1 4.80 B.amyloliquefaciens strain W49 Jakarta Barat Psk 1163f B.megaterium strain IAM 13418 99 NR_043401 15.19 Psk 3163f B. megaterium strain IAM 13418 97 NR_043401 27.34 Psk 4163f B. megaterium strain IAM 13418 99 NR_043401 32.02 Ptk 3163f B. subtilis subsp. subtilis strain DSM 10 98 NR_027552 24.00 Rangkas Bitung Rbt 2163f B. megaterium strain IAM 13418 96 NR_043401 35.97 99 NR_027552 Rbt 3163f B. subtilis subsp. subtilis strain DSM 10 32.93 NR_043401 Subang Sbg 1163f B. megaterium strain IAM 13418 96 11.18 Sbg 3163f B. amyloliquefaciens strain W49 98 KC441855.1 6.14 Kediri 99 NR_041455 Smk 1163f B. amyloliquefaciens strain NBRC 15535 4.62 Smk 3163f B. amyloliquefaciens strain NBRC 15535 100 NR_041455 29.10 Smk 4163f B. amyloliquefaciens strain NBRC 15535 99 NR_041455 3.16 99 7.17 Smk 5163f B.subtilis subsp. subtilis strain DSM 10 NR_027552 B.amyloliquefaciens strain NBRC 15535 Smk 8163f 98 NR_041455 3.40

Table 1. Characteristics of amylase-producing Bacillus spp. from cassava "tape"

to 35.97 unit/ml at 37 °C (Table 1). Genome of all *Bacillus* spp. isolates have been successfully isolated from cell cultures using protocol kit of Fermentas® Genomic DNA Purification Kit. PCR amplification of 16S rRNA gene sequences yielded DNA fragments with single band at 1,300 bp for each *Bacillus* sp. strains (data not shown).

BLASTN results of the partial sequence of 16S rRNA gene (about 800 to 950 nucleotides) showed high similarity with *Bacillus* spp. with maximum identities for each isolate in range of 94-100% with E-value 0 (Table 1). The distribution of all isolates was only on four species of *Bacillus* sp. such as *B. megaterium*, *B. subtilis*, *B. amyloliquefaciens*, and *B. thuringiensis*.

Neighbour joining tree based on 16S rRNA gene sequences was successfully constructed. Phylogenetic tree showed the relation among twenty-six of *Bacillus* sp. isolates (Figure 1). It showed that the isolates were divided into four groups (Figure 1). Group 1 (11 isolates), group 2 (seven isolates), group 3 (two isolates), and group 4 (six isolates) were quite related to *B. amyloliquefaciens*, *B. subtilis*, *B. thuringiensis*, and *B. megaterium* references strains respectively (Tabel 1).

DISCUSSION

This is a preliminary study of amylase-producing *Bacillus* spp. from "Tape". A total nine samples of "Tape" examined contained of amylase-producing *Bacillus* spp. *Bacillus* spp. are the organisms which responsible for any food fermentations and spoilage of foods in general, due to their versatile metabolism and heat resistant spores. Several fermented products rely on the participation of various *Bacillus* species,

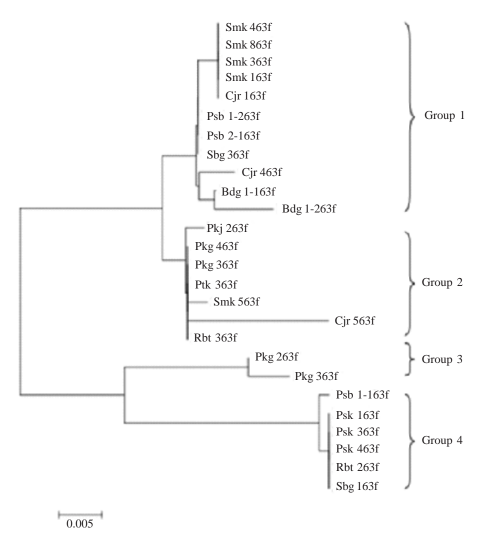


Figure 1. Dendogram showing genetic diversity among the partial 16S rDNA sequence of 26 amylase - producing *Bacillus* spp. strains from cassava fermented "Tape". The neighbour joining (NJ) tree was constructed using Mega 5 with 100 bootstrap analysis of replication.

including cassava fermented. It is widely distributed in many country such as: *B. cereus* and *B. subtilis* in Lafun – African (Padonou *et al.* 2009), *Bacillus* spp. in "attie'ke'" – Abidjan (Assanvo *et al.* 2006), *Bacillus* sp. in "fufu" – Nigeria (Achi & Akomas 2006), and *B. subtilis* in akyeke – Ghana (Obilie *et al.* 2003).

Bacillus spp. have an important role in cassava fermentation. During fermentation, many *Bacillus* produce enzymes, which are hydrolyzed oligosaccharides into easily digestible sugars (Chantawannakul *et al.* 2002; Joo *et al.* 2007). *Bacillus* spp. influenced the taste and aroma of "fufu" (Okafor *et al.* 2008). *Bacillus* sp. was also reported involve in the textural modification of cassava tissue during fermentation (Amoa-Awua & Jakobsen 1995; Obilie *et al.* 2003). It was reported that *B. subtilis* produced flavors of "Tape" most preferred by panelists (Barus & Wijaya 2011).

The results of this study showed that there were *B. subtilis*, *B. amyloliquefaciens*, *B. megaterium*, and *B. Thuringiensis* found in "Tape". Amylase assay (Table 1) showed that all *Bacillus* spp. isolates produced diverse amylase activity. *Bacillus* spp. isolates in each group also have diverse amylase activities. Example, group 1 (11 isolates) have amylase activities with range from 4.80 to 29.10 unit/ ml at 37 °C (Table 1). The Rbt 2, Rbt 3, and Smk 3 isolates that had highest amylase activity were closely related to *B. megaterium*, *B. subtilis*, and *B. amyloliquefaciens* respectively. The isolates will be studied further to determine their amylase activity and role in improving qualty and taste of tape.

Phylogenetic dendogram based on 16S rRNA gene sequences (Figure 1) generated four groups. The groups were dominated by *B. amyloliquefaciens*, *B. subtilis*, *B. thuringinensis*, and *B. megaterium* (Table 1). The isolates in the same group were closely related to one reference species of *Bacillus* with maximum identities in range of 94-100%.

In this study, sequences of the 16S rDNA can be used to reveal diversity of *Bacillus* spp. accordance with those reported by Wahyudi *et al.* (2010)

ACKNOWLEGEMENT

This study was supported by Competitive Grant Program from Atma Jaya Catholic University and Competitive Grant Program from Directorate General of Higher Education-Indonesia to Tati Barus.

REFERENCES

- Achi OK, Akomas NS. 2006. Comparative assessment of fermentation techniques in the processing of fufu, a traditional fermented cassava product. *Pak J Nutr* 5:224-229. http://dx.doi.org/10.3923/pjn.2006.224.229
- Amoa-Awua WKA, Jakobsen M. 1995. The role of *Bacillus* species in the fermentation of cassava. *J Appl Bacteriol* 79:250-256. http://dx.doi.org/10.1111/j.1365-2672.1995.tb03134.x
- Assanvo JB, Agbo GN, Behi YEN, Coulin P, Farah Z. 2006. Microflora of traditional starter made from cassava for "attie'ke"" production in Dabou (Co^ te d_Ivoire). *Food Control* 17:37-41. http://dx.doi.org/10.1016/ j.foodcont.2004.08.006
- Barus T, Wijaya LN. 2011. *Mikrobiota dominan dan perannya dalam cita rasa "Tape" singkong. Biota* 16:354-361.
- Chantawannakul P, Oncharoen A, Klanbut K, Chukeatirote E, Lumyong S. 2002. Characterization of proteases of *Bacillus* subtilis strain 38 isolated from traditionally fermented soybean in Northern Thailand. Sci Asia 28:241-245. http:// /dx.doi.org/10.2306/scienceasia1513-1874.2002.28.241
- Farnworth ER. 2003. Handbook of Fermented Functional Foods. Ed ke-2. Florida: CRC Pr.
- Joo MH, Hur SH, Han YS, Kim JY. 2007. Isolation, identification, and characterization of *Bacillus* strains from the traditional Korean soybean-fermented food, chungkookjang. *J Appl Biol Chem* 50:202-210.
- Marchesi JR, Sato T, Weightman AJ, Martin TA, Fry JC, Hiom SJ, Wade WG. 1998. Design and evaluation of useful bacterium-specific PCR primers that amplify genes coding for bacterial 16S rRNA. *Appl Environ Microbiol* 64:795-799.
- Obilie EM, Tano-Debrah K, Wisdom Kofi Amoa-Awua. 2003. Microbial modification of the texture of grated cassava during fermentation into akyeke. *Int J Food Microbiol* 89:275-280. http://dx.doi.org/10.1016/S0168-1605(03) 00294-0
- Oguntoyinbo FA, Sanni AI, Franz CMAP, Holzapfel WH. 2006. In vitro fermentation studies for selection and evaluation of *Bacillus* strains as starter cultures for the production of okpehe, a traditional African fermented condiment. *Int J Food Microbiol* 113:208-218. http://dx.doi.org/10.1016/ j.ijfoodmicro.2006.07.006
- Okafor N, Ijioma B, Oyolu C. 2008. Studies on the microbiology of cassava retting for foo-foo production. *J Appl Microbiol* 56:1-13. http://dx.doi.org/10.1111/j.1365-2672.1984. tb04691.x
- Ouoba LI, Diawara B, Amoa-Awua Wk, Traoré AS, Møller PL. 2004. Genotyping of starter cultures of *Bacillus subtilis* and *Bacillus pumilus* for fermentation of African locust bean (Parkia biglobosa) to produce Soumbala. *Int J Food Microbiol* 90:197-205. http://dx.doi.org/10.1016/S0168-1605(03)00302-7
- Padonou SG, Nielsen DS, Hounhouigan JD, Thorsen L, Nago MC, Jakobsen M. 2009. The microbiota of Lafun, an African traditional cassava food product. *Int J Food Microbiol* 133:22-30. http://dx.doi.org/10.1016/j.ijfoodmicro.2009.04. 019

- Pawiroharsono S. 2007. Potensi pengembangan industri dan bioekonomi berbasis makanan fermentasi tradisional. *J Ilmu Kefarmasian Indonesia* 5:85-91.
- Sarkar PK, Hasenack B, Nout MJR. 2002. Diversity and functionality of *Bacillus* and related genera isolatd from spontaneously fermented soya bean (Indian kinema) and locust beans (African soumbala). *Int J Food Microbiol* 77:175-186. http://dx.doi.org/10.1016/S0168-1605(02) 00124-1
- Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S. 2011. MEGA5: Molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. *Mol Biol Evol* 28:2731-2739. http://dx.doi.org/10.1093/molbev/msr121
- Terlabie NN, Sakyi-Dawson E, Amoa-Awua WK. 2006. The comparative ability of four isolates of *Bacillus subtilis* to ferment soybeanss into dawadawa. *Int J Food Microbiol* 106:145-152. http://dx.doi.org/10.1016/j.ijfoodmicro.2005. 05.021
- Wahyudi AT, Prasojo BJ, Mubarik NS. 2010. Diversity of antifungal compound-producing Bacillus spp. isolated from Rhizosphere of soybean plant based on ARDRA and 16S rRNA. *HAYATI J Biosci* 17:145-150. http://dx.doi.org/ 10.4308/hjb.17.3.145