Diversity of antibiotic-producing Actinomycetes in mangrove forest of Torosiaje, Gorontalo, Indonesia

YULIANA RETNOWATI1,2,*, LANGKAH SEMBIRING3, SUKARTI MOELJOPAWIRO3, TJUT SUGANDAWATY DJOHAN3, ENDANG SUTARININGSIH SOETARTO3**

1 Graduate Study Program of Biology, Faculty of Biology, Universitas Gadjah Mada. Jl. Teknika Selatan, Sleman 55281, Yogyakarta, Indonesia.
2Department of Biology, Faculty of Mathematics and Natural Sciences, Universitas Negeri Gorontalo. Jl. Jenderal Sudirman No. 6, Kota Gorontalo 96128, Gorontalo, Indonesia. Tel.: +62-435-821125; Fax.: +62-435-821752. *email: yuliana_r@yahoo.com
3Faculty of Biology, Universitas Gadjah Mada. Jl. Teknika Selatan, Sleman 55281, Yogyakarta, Indonesia. Tel./Fax.: +62-274-580839, **email: annisah-endang@ugm.ac.id

Abstract. Retnowati Y, Sembiring L, Moeljopawiro S, Djohan TS, Soetarto ES. 2017. Diversity of antibiotic-producing actinomycetes in mangrove forest of Torosiaje, Gorontalo, Indonesia. Biodiversitas 18: 1453-1461. Actinomycetes for antibiotic production have been studied at various extreme environments. Mangrove forest of Torosiaje in Gorontalo Province, Indonesia has unique geomorphological conditions where the forest is surrounded by karst ecosystem consisting of fringe and overwash types. Therefore, the objective of this study was to analyze the distribution and diversity of antibiotic-producing Actinomycetes in various rhizosphere including different locations and mangrove species. Samples of rhizosphere soil were collected in the depth of 0-10 cm, which was then subjected to detailed physicochemical analysis. Actinomycetes were collected through heating pre-treatment (60°C for 15 min) followed by culturing in the Starch Casein Agar medium supplemented with cycloheximide and nystatin. The screening process of antibiotic-producing Actinomycetes was based on Agar block method against pathogenic microorganisms. Grouping of Actinomycetes was determined by ARDRA fingerprinting analysis. The diversity of Actinomycetes was analyzed based on sequencing of 16S rDNA. The results showed that the distribution of Actinomycetes was found in overwash type, middle zone and upper zone of fringe type including rhizosphere of 7 species of mangrove. The highest population of Actinomycetes was found in rhizosphere of R. mucronata at the overwash type, and the lowest one found in rhizosphere of R. apiculata at the middle zone of fringe type. A total of 77 isolates amongst 167 isolate collection showed antibacterial activities. Forty seven representatives from 77 antibacterial-activities isolates were selected using ARDRA for partial characterization according to their phylogenetic diversity. Sequencing and analysis of 16S rDNA from selected-representative isolates displayed the presence of members associated with Actinomycetes genera such as Streptomyces, Amycolatopsis, Saccharomonospora, and Nocardiopsis. The member of genus Streptomyces such as Streptomyces qinglanensis and Streptomyces champavatii were distributed across locations. Genus Saccharomonospor and Nocardiopsis were mostly found at the overwash type, while Amycolatopsis was found at the upper zone of fringe type.

Keywords: Actinomycetes, diversity, fringe, mangroves, overwash

INTRODUCTION

Actinomycetes are Gram-positive bacteria that produce bioactive compound (Amrita et al. 2012). 70-80% of bioactive compounds are produced by Actinomycetes (Kanna et al. 2011), especially Streptomyces (Berdy 2005; Chaudary et al. 2013), and a part of bioactive compounds is produced by rare-Actinomycetes (Berdy 2005). The exploration of Actinomycetes that produce bioactive compounds has been focused on the extreme environment due to the rediscovery of bioactive compounds from terrestrial environment (Nolan and Cross 1988; Mangamuri et al. 2014). Mangrove forest is one of interesting ecosystems to explore Actinomycetes (Santhi et al. 2010; Baskaran et al. 2011; Khanna et al. 2011; Naikpatil and Rathod 2011; Ravikumar et al. 2011; Mangamuri et al. 2012; Mangamuri et al. 2014).

Mangrove ecosystem is a habitat of various of Actinomycetes. The Actinomycetes of mangrove ecosystem produce 122 bioactive compounds, where 73 of them have been identified as new kinds of bioactive compounds, and 49 compounds have not been identified, yet (Xu et al. 2014). The bioactive compounds from mangrove Actinomycetes demonstrate antibacterial activity (Sharma et al. 2011; Hunadanamra et al. 2013; Das et al. 2014; Phongsopitanum et al. 2014), anti fungal activity (Suthindiran and Kannabiran 2010; Sharma et al. 2011; Phongsopitanum et al. 2014; Das et al. 2014), and anti viral activity (Waksman 1984) and anti cancer activity (Hong et al. 2009; Suthindiran and Kannabiran 2010; Becerril-Espinosa et al. 2012; Ravikumar et al. 2012; Anupa et al. 2013; Xu et al. 2014).

Actinomycetes producing bioactive compounds in mangrove ecosystem are distributed on rhizosphere and non rhizosphere (Xiao 2009; Ravikumar et al. 2011), which rarely live as endophytics on stems and leaves of mangrove (Ravikumar et al. 2011; Gayathri and Muralikrishnan 2013; Govindasamy and Franco 2013). The number of Actinomycetes population in rhizosphere are more than those in non rhizosphere. The various number of Actinomycetes is also depended on types of mangrove (Bafagurunathan et al. 2010; Ravikumar et al. 2011;
Ramirez-Elias et al. 2014).

Mangrove ecosystem of Torosiaje is karst mangrove ecosystem including two types of forest, i.e., fringe and overwash types. The fringe type is divided into two zones, i.e. middle and upper zone. The middle zone was dominated by Rhizophora apiculata and Bruguiera gymnorrhiza while the upper zone consists of Sonneratia alba, Avicennia marina, Xylocarpus sp., Ceriops tagal and Sonneratia alba (Katili et al. 2014). Based on the uniqueness of Torosiaje mangrove ecosystem and some other important aspects; however, the diversity and distribution of Actinomycetes in this location had not been comprehensively explored, yet. Therefore, the objective of this study was to analyze the diversity and distribution of antibiotic-producing Actinomycetes in rhizosphere of various species mangrove located at mangrove ecosystem of Torosiaje, Gorontalo, Indonesia.

MATERIALS AND METHODS

Study area and sampling of soil rhizospheres. This study was conducted from April 2016 to April 2017. Samples of mangrove rhizospheres soil were collected from mangrove forest at Torosiaje, Pohuwato District, Gorontalo Province, Indonesia (Figure 1). The total of eight soil samples was collected from 3 different location i.e overwash type, middle zone and upper zone of fringe type, including 7 species of mangrove rhizospheres (Rhizophora mucronata, Rhizophora apiculata, Bruguiera gymnorrhiza, Avicennia marina, Xylocarpus sp., Ceriops tagal and Sonneratia alba). Sampling sites were located at 3 different coordinates i.e N. 00°.28'.21.9"; E.121°.26'.55.7"; N. 00°.28'.50.5"; E. 121°.27'.56.8" and N. 00°.28'.47.2; E. 121°.27'.56.2". Soil samples were collected on the depth of 0-10 cm by modified soil core with 4 replication.

Soil physicochemical character analysis. The physicochemical analysis of soil samples (salinity, soil texture, soil pH, C-organic, total-N, N-NO₃, Fe, Mn, Zn, Cu, phosphor, K₂O and total S) was conducted at Soil Chemical laboratory, Faculty of Agriculture, Gadjah Mada University, Yogyakarta, Indonesia.

The isolation of actinomycete isolates. The initial treatment of rhizosphere soil samples was based on dry wet method on 60°C for 15 minutes in combination of sea water and aquadest (1: 1 v/v) (Mangamuri et al. 2012). Soil suspension was serial-diluted up to 10⁻⁶. Soil suspension as much as 100 µL was inoculated in Starch Casein Agar (SCA) medium supplemented by 25 µg/mL of cycloheximide and 25 µg/mL of nystatin (Baskaran et al. 2011). Then incubated on 28 ± 2 °C for 10-28 days. Colony Forming Unit (CFU) was calculated and isolated in slant ISP 2 medium.

Screening of actinomycete isolates for potential antibiotics production. The screening of Actinomycetes for potential antibiotics production was based on Agar Block Method. The medium Muller Hinton agar got 5 holes and each hole was 5 mm diameters. After 28 days of age, actinomycete isolates cultured on ISP2 medium were then clipped around 5 mm diameters and moved to the holes. The Actinomycetes were incubated on 28°C ±2 for 3-7 days. The cell mass of microorganisms test (Escherichia coli, Staphylococcus aureus and Bacillus subtilis) on Nutrient Broth medium was equated by spectrophotometry on OD 0.6. Afterward, 300 µL of microorganisms test was inoculated in the around of actinomycete growing by surface plate method, then incubated on 37°C for 2 x 24 hours.

Extraction of genomic DNA. The spores of actinomycete isolates were suspended in aquadest. The suspension of spore was inoculated in 20 mL of ISP2 broth, then incubated in shaker incubator at 150 rpm for 4 to 7 days. Cell pellet in 20 mL of ISP2 broth was separated by centrifugation at 13,000 rpm for 5 min. The cell pellet was added with 20 mg of glass bead and 1 mL of buffer lysis (100mM Tris HCl pH 8, 50 mM EDTA pH 8, 2% SDS and 100 mM NaCl) then shaken out vigorously by vortex for 2 min. Proteinase K (20 mg/mL) as much as 10 µL was added then shaken out vigorously for 30 min. Then 60 µL of lysozyme (100 mg/mL) was added and incubated on 55°C for 30 min in water bath. The suspension was centrifugated at 13,000 rpm for 5 min, then cell pellet was discarded and the supernatant was resuspended by phenol (1: 1 v/v). The suspension of supernant and phenol were shaken vigorously for 30 minutes, then centrifugated on 13,000 rpm for 10 min. The top layer was moved to new microtube and added with cold chloroform (1: 1 v/v), then shaken out slowly for 20 minutes. Then they were centrifugated on 13000 rpm for 10 minutes. The top layer was moved to new microtube and added with cold absolute ethanol (1: 1 v/v), and saved on -80°C for 2 hours or -20°C for overnight. The DNA genome pellet was separated by centrifugation at 13000 rpm for 10 minutes, cleaned up with 70% cold ethanol then dried for 2 hours. The DNA genome was suspended in 50 µL of TE (pH 8). The suspension of DNA genome was added by 3 µL of RNAase (10µg/mL) and incubated on 37°C for 1 hour in water bath. The genomic DNA was applied or saved over -20°C. The genomic DNA were examined by 2% agarose gel electrophoresis. The gels were stained by syber and visualized under UV transilluminator (Magarvey et al. 2004).

Amplification of 16S rDNA. The 16S rDNA was amplified from genomic DNA samples using eubacterial universal primers 27 F (5'-AGAGTTTGATCCTGGCTCAG-3') and 1492R (5'-TACGGCTACCTTGTTACGACTT-3'). The amplification was carried out in 25 µL volume by using 1 µL of genomic DNA as template with 12.5 µL volume by using 1 µL of genomic DNA as template with 12.5 µL of 2X GoTaq (R) Green Master Mix [buffer reaction, deoxynucleotide triphosphate (dNTPs), Taq DNA polymerase, and MgCl₂], 9.5 µL dH2O, and 1 µL universal primer 27F and 1 µL 1492R. The thermal cycling conditions were as follows: initial denaturation at 96°C for 5 min; 29 cycles at 95°C for 1 min, 54°C for 1 min, 72°C for 2 min and post-extension at 72°C for 10 min. The amplification reaction was performed by Bio-Rad thermal cycler (MyCycler, Bio-Rad, USA) and the amplified products were examined by 2% agarose gel electrophoresis (Mangamuri et al. 2012).
Figure 1. Location of mangrove forest of Torosiaje, Gorontalo, Indonesia. The alphabet A, B and C indicating the sampling site of mangroves soil rhizosphere at different mangrove forest type. A. overwash type (N. 00°28’.21,9”;E. 121°26’.55,7”), B. middle zone of fringe type (N. 00°28’.50,5”; E. 121°27’.56,8”), and C. upper zone of fringe type (N. 00°28’.47,2; E. 121°27’.56,2”)

Amplified ribosomal DNA restriction analysis (ARDRA). To identify the number of polymorphic groups and to select representative strains among the actinomycete isolates, aliquots of purified 16S rDNA amplicons were subjected to amplified ribosomal DNA restriction analysis. The purified PCR products were digested separately with HaeIII and MspI, in 10 μL reaction volume by using manufacture’s recommended buffer and temperature. The digested restriction fragments were electrophoresed in 2% agarose gels using TE buffer. The gels were stained with syber and visualized under UV transilluminator. Strong and clear bands were scored in a binary data used for similarity and clustering analysis in numerical taxonomy analysis program package, MVSV Plus (Multivariate Statistical Package) Version 3.1A. Similarities among the isolates were calculated by Jaccard's coefficient, and the dendrogram was constructed using UPGMA (Unweighted Pair Group Method with Arithmetic Averages) method (Jose and Jebakumar 2012). If the strains formed a cluster defined at the 72% similarity level, they would be assigned to a single species (Meng Yuan et al. 2014).

16S rDNA sequencing and phylogenetic analysis. The PCR amplicons of 47 actinomycete isolates with representative ARDRA profiles were purified by using PCR product purification spin kit (BigDye® Terminator v3.1, USA) and sequenced by ABI PRISM 3730xl Genetic Analyzer develop by Applied Biosystems using same primary set as used in PCR amplification. All the sequences obtained from sequencing phase were analyzed and edited by using BioEdit soft-ware (Jose and Jebakumar 2012). Initially, all the 16S rDNA sequences were compared to sequences in GenBank by using the online service of Basic Local Alignment Search Tool to determine the approximate phylogenetic position. Sequences were aligned by using ClustalW with representative actinomycete 16S rDNA sequences, and a phylogenetic tree was constructed by using the Molecular Evolutionary Genetics Analysis (MEGA) software, version 6.06 (Yuan et al. 2014).
RESULTS AND DISCUSSION

Physico-chemical characteristics of soil, population and distribution of Actinomycetes

Mangrove forest ecosystem is a habitat for various types of tolerant microorganisms to an extreme environmental conditions. Mangrove forest ecosystems are rich in organic, nitrogen and sulfur materials that required for the grow of microorganisms (Malek et al. 2014). Mangrove ecosystem of Torosiaje, Gorontalo consist of two type mangrove forest i.e., overwash type and fringe type. Fringe type divided into 2 zone i.e., middle zone and upper zone. Each of mangrove forest type and zone were show the specific physicochemical characters. The sediment salinity level was in range of 23.1 - 31.3 ppt. The physico-chemical characteristics of soil samples collected from overwash mangrove, middle and upper zone of fringe mangrove at mangrove forest of Torosiaje, Gorontalo were studied using standard protocols and summarized in Table 1. Several parameters, such as pH, salinity of soil, soil texture, macroelements and microelements have been measured. The results show that there were differences across all measurable characters in all three locations. Overwash type mangroves forest is washed periodically by tides of sea water so that it affects to the physicochemical character of the soil to be quite extreme, including salinity and high level of Fe, while the organic-C content was relatively low. Middle zone and upper zone in fringe type mangrove forest were shown the differences physico-chemical characters amongst both of them. The upper zone tends to be at least exposed by sea tides was shown sandy soil conditions with low salinity and neutral pH.

The physicochemical character of the 3 sampling sites were responded by actinomycete as one of the members of the mangrove ecosystem. The responses of actinomycete were indicated by the various number of population in the three locations. The highest population of Actinomycetes was found on overwash type, and the lowest one was found on mudle zone of fringe type (Table 2). The periodically tides of overwash type is causes a continuously flow of macro-element and micro-element that supporting the growth of soil microorganisms. Otherwise, the geographical condition of the sloping fringe mangrove forest causes the middle zone is more exposed by the tides than the upper zone. This condition might cause the nutrient in the middle zone was more transported out of the zone.

Actinomycetes exploration at mangrove forest of Torosiaje is more emphasized on rhizosphere soil samples of 7 types mangroves at the depth of 0-10 cm. Nevin et al. (2000) and Mangamuri et al. (2011) claimed that the most Actinomycetes population were found in mangrove rhizosphere at the depth of 0-20 cm and apparently varied in each species of mangroves. The high population of Actinomycetes in the rhizosphere was due to the root exudation activities. That activity provided the space for actinomycete colonization and depended on the availability of amino acids and high concentrations of sugars and low concentrations of tannin content in rhizosphere (Atlas and Barta 1998; Badri and Vivanco 2009; Mangamuri et al. 2011). Mangamuri et al. (2011) also reported that the

<table>
<thead>
<tr>
<th>Physicochemical characters of soil</th>
<th>Fringe type</th>
<th>Overwash type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Salinity (ppt)</td>
<td>26.56</td>
<td>31.3</td>
</tr>
<tr>
<td>pH</td>
<td>3.16</td>
<td>6.04</td>
</tr>
<tr>
<td>Soil texture:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sand (%)</td>
<td>62</td>
<td>62</td>
</tr>
<tr>
<td>Dust (%)</td>
<td>31</td>
<td>10</td>
</tr>
<tr>
<td>Clay (%)</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td>C-organic (%)</td>
<td>12.63</td>
<td>2.18</td>
</tr>
<tr>
<td>N (ppm)</td>
<td>102.94</td>
<td>610.40</td>
</tr>
<tr>
<td>Total-S (%)</td>
<td>0.38</td>
<td>0.43</td>
</tr>
<tr>
<td>N-NO3 (ppm)</td>
<td>24.09</td>
<td>62.31</td>
</tr>
<tr>
<td>P2O5 (ppm)</td>
<td>63.33</td>
<td>3.09</td>
</tr>
<tr>
<td>K (me/100g)</td>
<td>1.74</td>
<td>1.23</td>
</tr>
<tr>
<td>Fe (ppm)</td>
<td>427.69</td>
<td>441.24</td>
</tr>
<tr>
<td>Mn (ppm)</td>
<td>4.76</td>
<td>8.08</td>
</tr>
<tr>
<td>Zn (ppm)</td>
<td>4.31</td>
<td>11.03</td>
</tr>
<tr>
<td>Cu (ppm)</td>
<td>0.37</td>
<td>1.07</td>
</tr>
</tbody>
</table>

Table 1. Physico-chemical characteristic of soil samples from overwash type, middle zone and upper zone of fringe type at mangrove forest of Torosiaje, Gorontalo, Indonesia

<table>
<thead>
<tr>
<th>Mangrove forest type</th>
<th>Zone</th>
<th>Mangrove type</th>
<th>Actinomycete population (CFU/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overwash</td>
<td>R. mucronata</td>
<td>23.5 x 10^3</td>
<td>78,4x10^3</td>
</tr>
<tr>
<td>Total on overwash</td>
<td>B. gymnorrhiza</td>
<td>3.5 x 10^1</td>
<td></td>
</tr>
<tr>
<td>Fringe</td>
<td>R. apiculata</td>
<td>10,1 x 10^3</td>
<td>50,7 x 10^2</td>
</tr>
<tr>
<td>Middle</td>
<td>B. gymnorrhiza</td>
<td>34 x 10^2</td>
<td></td>
</tr>
<tr>
<td>Sub total</td>
<td>S. alba</td>
<td>33,6 x 10^3</td>
<td>26,3 x 10^2</td>
</tr>
<tr>
<td>Upper</td>
<td>A. marina</td>
<td>26,3 x 10^2</td>
<td>18,6 x 10^2</td>
</tr>
<tr>
<td>Sub total</td>
<td>Xylocarpus</td>
<td>96,1 x 10^2</td>
<td>73,4 x 10^2</td>
</tr>
<tr>
<td>Total on fringe</td>
<td>C. tagal</td>
<td>96,1 x 10^2</td>
<td>73,4 x 10^2</td>
</tr>
</tbody>
</table>

Table 2. Population of Actinomycetes at 7 types mangrove rhizosphere
tannins, alkaloids and various secondary metabolites (Atlas and Barta 1998; Walker et al. 2003). Exudation activities of various compounds in plant roots can regulate the soil microbial community by mediating soil-microbial root interactions, altering soil chemical physical properties and inhibiting the growth of plant competitors (Badri and Vivanco 2009). Each of mangrove species was produce a specific kind of root exudates. The adult of *Avicennia marina* is produce root exudates with different levels of glucose and fructose composition (Lie et al. 2014) and root exudates of *Bruguiera gymnorrhiza* contain of aminopyrine, palmitic acid, stearic acid, di-n-propyl-ether, and 2.5-angeloglobulon (Kumar et al. 2009).

**Screening of antibiotic-producing Actinomycetes based on agar block method**

Mangrove forests are interesting ecosystems for the discovery of antibiotic-producing Actinomycetes (Santhi et al. 2010; Baskaran et al. 2011; Khanna et al. 2011; Naikpatil and Rathod 2011; Ravikumar et al. 2011; Mangamuri et al. 2012; Mangamuri et al. 2014). The extreme conditions of mangrove environment are expected to stimulate the actinomycete to produce antibiotics with specific character (Basilio et al. 2003). The bioactive compound were shown the activity as antibacterial, antifungal, antiviral and anticancer (Suthindhiran et al. 2010; Ravikumar et al. 2011; Amrita et al. 2012; Doroghazi and Metcalf 2013; Zotchev 2014). This study proves that amongst 167 actinomycete isolates, there were 77 actinomycete isolates that show the antibacterial activity against pathogenic bacteria. The highest number of antibiotic-producing actinomycete isolates was found at rhizosphere of *R. mucronata* in overwash type, and the lowest one at rhizosphere of *C. tagal* in upper zone of fringe type (Figure 3). While the high ability of actinomycete isolates to against pathogenic microbes was found in rhizosphere of *A. marina* and *Xylocarpus sp.* at upper zone of fringe type. That was shown that mangrove forest of Torosiasje, Gorontalo was potential to exploration antibiotic-producing Actinomycetes.

**Amplification of 16S rDNA and ARDRA.** PCR amplification of the 16S rDNA using a set of universal eubacterial specific primers: 27 F and 1492R yielded a single amplicon of ~1500 bp for 77 the antibiotic-producing actinomycete isolates. The amplicons of 16S rDNA were used to Amplified Ribosomal DNA Restriction Analysis (ARDRA) with restriction enzymes endonuclease *Hae* III and *Msp*I. Digestion of these amplicons with *Hae* IIII and *Msp*I were generated almost similar profile for several of isolates. Whereas, when amplicons were digested by restriction enzyme *Msp*I, different number of fragments were produced about 5 to 9 fragments with varied sizes about 30 to 900 bp for different isolates.

The ARDRA pattern was used for dendrogram construction using UPGMA and Jaccard's coefficient algorithms (Figure 4). The 77 of antibiotic-producing actinomycete isolates were divided into 47 groups in a dendrogram inferred from ARDRA pattern obtained from *Msp*I restriction enzymes. The groups consisted of 1 to 5 actinomycete isolates. The groups formed on the dendrogram provide information about the presence of the same species found in different location and different kinds of mangrove. If the isolates formed a group/cluster defined at the 72% similarity level, they would be assigned to a single species (Meng Yuan et al. 2014).

**Figure 2.** The number of Actinomycete isolates in 7 types of mangrove rhizosphere at mangrove forest of Torosiasje Gorontalo, Indonesia. O. Overwash; FM. Fringe/middle zone; FU. Fringe/upper zone

**Figure 3.** The number of antibiotic-producing Actinomycete isolates at 7 kind of mangrove rhizosphere. O. Overwash; FM. Fringe/middle zone; FU. Fringe/upper zone
Actinomycetes community composition and phylogenetic analysis. The 16S rDNA of 47 representative isolates that belong to different clusters established by ARDRA was sequenced and used to determine the diversity of isolates. Based on similar criteria of 98% at the 16S rRNA gene, the 47 isolates were sorted into 6 phylotypes, belong to 4 different genera. The four genera were Streptomyces, Saccharomonospora, Nocardiopsis and Amycolatopsis (Figure 5). The most abundant group of isolates was affiliated with the genus Streptomyces, represented by four ribotypes accounting for 95% of the total actinomycete population. The ribotypes ORm2-a was closely related to type strain Streptomyces qinglanensis (GenBank: NR_109303.1) with 99.9% sequence similarity, representing 83% of total isolates. The other ribotypes of Streptomyces, OBg1-3 (representing 5% of isolates) and FMRa1-1 (representing 7% of all isolates) were closely related to type strain Streptomyces sanyensis (GenBank: NR_116599.1) with 99% sequence similarity and Streptomyces champavatii (GenBank: NR_115669.1) with 97% sequence similarity, respectively. The second most dominant group in the isolates was the genera Nocardiopsis, representing 3% of total actinomycete isolates. Ribotype ORm2-f was closely related to type strain Nocardiopsis lucentensis (GenBank: NR_026342.1) with 99% of sequence similarity. Ribotype FUXy2-d and OBg4-e representing 1% of total actinomycete isolates, respectively. Ribotype FUXy2-d was closely related to type strain Amycolatopsis sacchari (GenBank: NR_112696.1) with 94% of sequence similarity. While, ribotype OBg4-e was closely related to type strain Saccharomonospora amisosensis (GenBank: NR_109529.1) with 97% of sequence similarity.

The four genus of antibiotic-producing Actinomycetes were distributed at overwash type, middle zone and upper zone of fringe type of mangrove forest of Torosiaje, Gorontalo (Figure 6). Several studies have been conducted in various mangrove ecosystems around the world and managed to discover various new types of Actinomycetes from rhizosphere of mangrove, non rhizosphere and endophytes in stems, roots and leaf of mangrove (Mangamuri et al. 2011). Several types of members of the Actinomycetes were isolated from habitats with moderate to high salt concentration such as saline soil or marine sediment (Sabry et al. 2004) and slattern (Chun et al. 2000).

In conclusion, distribution and diversity of Actinomycetes were explored from mangrove ecosystem at Torosiaje, Pohuwato District, Gorontalo Province, Indonesia. 167 actinomycete isolates were isolated from 7 types of mangrove rhizosphere. They were distributed at overwash mangrove, middle and upper zone of fringe mangrove. Among 167 actinomycete isolates, there were 77 actinomycete isolates shown antibacterial activity against pathogenic bacteria. The isolates which show high antibacterial activity were found at rhizosphere of A. marina and Xylocarpus sp. The Actinomycetes that produce antibiotics were assigned to four genera i.e Streptomyces, Saccharomonospora, Amycolatopsis and Nocardiopsis on the basis of their16S rDNA sequences. This is the first step towards better understanding of actinomycete from mangrove ecosystem at Torosiaje, Gorontalo.
Figure 4. UPGMA dendrogram inferred from ARDRA patterns of Actinomycetes isolated from mangrove rhizosphere. The dendrogram shows the clustering of 77 antibiotic-producing actinomycete isolates generated from amplified ribosomal DNA restriction analysis with restriction endonuclease MSP1, using the UPGMA algorithm and the Jaccard's coefficient. Number 1 to 47 represent the forty seven clusters obtained in the analysis.
Figure 5. Neighbour-joining phylogenetic tree inferred from 16S rRNA gene sequences. The phylogenetic tree shows the phylogenetic relationship of isolates with related genera. Bootstrap values are expressed as percentages of 1000 replications. Bootstrap values, >50% are shown at branch points. Score bar represents 1 nucleotide substitution per 100 nucleotides

ACKNOWLEDGEMENTS

The authors are grateful for the funding from “Research Funds Assistance” of Department of Education and Sports, Government of Gorontalo Province, Indonesia and “Hibah Penelitian Disertasi Doktor” of The Indonesian Ministry of Research and Technology in 2017. We are also thankful to Umar Pasandre as environmentalist of mangrove forest at Torosiaje, Gorontalo, Indonesia.

REFERENCES


Li TY, Xu F, Yan CL. 2014. Quantitatively analyzing monosaccharides in the root exudates of gray mangrove as its response to cadmium and copper stress. WIT Trans Built Env 156: 265-271


