Molecular Diversity of *Bacillus thuringiensis* and Bioinformatics Analysis of Local Isolate of Auky Island, Padaido District in Biak Numfor Papua as a Control of Anopheles Mosquito Larvae

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Abstract

This research aims to analyze the level of similarity and diversity among local isolates of *B. thuringiensis* Auky Island Padaido District in Biak Numfor Regency with NCBI gene bank base, the basis of which is to obtain *B. thuringiensis* isolates from Jayapura local isolates that can act as controllers of Anopheles mosquito larvae. Several steps in the research are 16s gene amplification, PCR product purification, cloning using pTA2 vectors and transformation into competent *E. coli* Zymo 5α cells, confirmation with PCR colonies, recombinant plasmid isolation, sequencing analysis and phylogenetic tree construction. The isolates of ABNP8, ABNP9, ABNP11, ABNP12 and ABNP18 have been detected as local isolates from in Auky Island Padaido District in Biak Numfor Papua Regency that have great potential as bioinsecticides, and capable of controlling and killing *Anopheles* mosquito larvae. Of the five isolates, ABNP8 isolates had unique diversity and characteristics and were different from the four other isolates. Based on the similarity analysis in the MEGA7 program, the similarity rate reached 84%. Its diversity can be seen from the uniqueness of the sequence and its position in different branching dendrograms.

Keywords: Anopheles, *Bacillus thuringiensis*, Genetic diversity, Auky Island, local isolates, Larva controller

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INTRODUCTION

*Bacillus thuringiensis,* which is one of the many facultative anaerobic bacteria, can produce spores, has gram-positive properties, and is an indigenous bacterium in soil, water, plant surfaces, dead insects, and seeds. This bacterium is known to be able to encode crystalline proteins (cry proteins) during the sporulation period. In order to obtain new *B. thuringiensis* isolates capable of producing cry proteins, the isolation process of several *B. thuringiensis* isolates has been transformed into routines in many industries. *B. thuringiensis* has been studied and is known to have a wide specificity in various orders of insects. *B. thuringiensis* has been studied and is known to have a wide specificity in various orders of insects. *B. thuringiensis* has been studied and is known to have a wide specificity in various orders of insects. *B. thuringiensis* has been studied and is known to have a wide specificity in various orders of insects. *B. thuringiensis* has been studied and is known to have a wide specificity in various orders of insects.

**Amplification of 16S genes by PCR**

The 16S gene was amplified using the PCR method. The total volume of the master mix is 90L divided into five tubes. The primers used are 16S rRNA primers (forward: 5' AGA GTT TGA TCC TGT CTC AG 3', and reverse: 3' ACG GCT ACC TTG TTA CGA CTT 5'). The DNA of PCR products was purified by Zymoclean Gel DNA Recovery Kit from Zymo Research, and then ligated and transformed in the 16S gene cloning process.

**Toyobo pTA2 plasmid ligation and transformation to E. coli Zymo 5α competent cells**

The ligation process of Gen16 is conducted through the making of ligation reaction on ice consisting of 1μL (0.05 pmol) pTA2 vector cloning (50 ng/μL), 1 μL T4 DNA ligase (1-3 U/μL), 16S of purification results 3 μL, and 2x rapid ligation buffer as much as 5 μL. Next, the solution went through divortex and centrifuged process for 3-5 minutes. The result of ligation process then incubated at temperature (4°C) for 18 to 24 hours. Eventually, it was transformed by through Sambrook method.

**PCR colony and recombinant plasmid isolation**

Positive colonies containing 16S Genes were amplified by PCR using T3 and T7 KOD FX Neo (Toyobo) primers at annealing temperature of 56°C. Isolation of recombinant plasmid using ZR Plasmid MiniPrep (Zymo Research)

**Sequencing analysis and phylogenetic tree construction**

The raw data from the sequencing was edited with the Seqtrace 0.9.0 program and Bioedit 7.2.5. Next, homologous species were searched based on its 16S gene sequences on NCBI. From the BLAST result, we selected species similar to *Bacillus thuringiensis* isolate Jayapura to be collected/inserted in one group. Sequence of consensus area 16S *B. Thuringiensis* isolates Jayapura and sequences from NCBI were dialigned by the Muscle method. The phylogenetic tree construction uses the Neighbor Joining algorithm, with a bootstrap value of 1000 in the MEGA version 6 program. These result of...
this process then presented through narrative descriptive to provide a clear understanding of the research.

RESULT AND DISCUSSION

Results of PCR amplification with 16S primers

The amplification process with 16S primary primer uses the colony PCR method with annealing temperature of 56°C. The amplified 16S gene was then electrophoresed using agarose gel with a concentration of 1%. The PCR results can be seen in Fig. 1.

At this stage, the 16S gene was successfully amplified, as evidenced by the band pattern profile after electrophoresis with a size of ± 1500 bp. The 16S target gene was successfully detected in five samples, namely codes 1–5. The visibility of these five samples shows a specific clear, and bright bands. These five samples which detected by the 16S gene were proven by the presence of DNA band fragments. Qualitative test uses a standard method namely DNA migration through agarose gel electrophoresis. Several factors that supported the successfulness of the electrophoresis process are: size and conformation of DNA molecules, agarose concentration, electric current, and temperature. Additionally, the DNA fragments also separated in the gel. Thus, the presence of EtBr as a dye will be bound between DNA double strands. The DNA band in agarose will fluoresce because EtBr contains fluorocene. The DNA-EtBr bond will be exposed to UV light so the qualitative test results show that the ribbon profile looks bright. Purified PCR products are prepared with the aim of eliminating all contaminants and potential inhibitors that can prevent the continuation of the cloning process16.

The results of cloning to the toyobo pTA2 vector plasmid and transformation to E. coli Zymo 5α

PCR amplifiers that have been transformed into host E. coli ZYMO5a are such that replication can occur when they have become recombinant plasmids. T4 DNA ligase is an enzyme used in the ligation process because it can catalyze the formation of phosphodiester bonds between 3'-OH groups of target DNA fragments and 5'-P groups in vector DNA. The success of cloning is influenced by: (1) Plasmids have an ori role as an initiator in replication, (2) the ampR gene that carries ampicillin resistance, (3) the MCS side helps form new recombinant DNA combinations and (5) the lacZ gene that plays a role in blue white screening. CCMB that is done when competent cell preparation will affect the cell wall and bind DNA to the cell surface17. On the other hand18 stated that E. coli cells treated with CCMB would be able to take DNA from bacteriophages and plasmid DNA. With the content of ampicillin in the medium, then E. coli cells that can grow are cells that only contain pTA2 plasmids because these plasmids have properties that are resistant to ampicillin.

PCR colonies were successfully performed on isolates ABNP8, ABNP9, ABNP11, ABNP12, and ABNP18, with 8 colonies each. ABNP8 isolates showed that most of them produced thick positive bands. Seen the size of ± 1.5 kb of an amplicon that is the same size as the target gene. The profile of the fragment band in the same position as the ladder (marker) is a strong indication of the detection of the 16S gene. Positive colonies that do not produce specific bands are caused by false positives, that is, they appear to be positive colonies, but in fact the plasmids do not contain insert genes. In general, confirmation of the presence of 16S genes in pTA2 resulting from transformation by PCR colonies was successfully carried out. The tape on the electrophoresis results measuring ± 1.5 kb shows that the 16S gene was

Fig. 1. PCR result of five isolates.
Note: M: Marker DNA ladder 1Kb, 1-5: sample ABNP8, ABNP9, ABNP 11, ABNP 12, and ABNP18 respectively
successfully inserted into pTA2. However, to be more convincing and validate the existence of the target gene is correct, then the sample is then sequenced to obtain DNA sequences which will then be aligned with the NCBI database.

Observation of transformant bacterial colonies containing recombinant plasmids

Based on observations, the transformation process was successfully carried out. The key factor to the success of the transformation is the proper

![Fig. 2. PCR results from recombinant plasmid PCR colonies](image)

Note: M: 1Kb DNA ladder markers, 1-8: eight colonies in each ABNP 8, ABNP 9, ABNP 11, ABNP 12, ABNP 18 isolates.

![Fig. 3. Results of alignment of five B. thuringiensis local isolates from Auky Island Padaido District in Biak Numfor Papua.](image)
process of heat shock transformation so that the plasmid that has been inserted with the target DNA enters the competent cell. The target colony is a white colony containing transformants, where MCS is inserted by 16S fragments so that it is not expressed and causes the colony to remain white. Thus, colonies containing recombinant plasmids are white colonies. Negative control is intended to be able to find out contamination during transformation. To find out the presence of insert genes in an isolated plasmid, it must be confirmed by PCR. The purpose of confirming the transformation results is to know the existence of the 16S gene in pTA2. This is also a way to see the successful integration of insert genes in plasmids by using primers that are specific to insert genes. To be more convincing and validate the existence of the insert gene is correct then the sample needs further analysis process, namely by sequencing to obtain the DNA sequence to be aligned with the data bank.

**Sequencing analysis**

Sequencing results in the form of chromatograms show a certain spectrum and intensity. The higher the plasmid concentration, the better the spectrum of sequencing results. The size of the target DNA is ± 1500 bp, so that the sequencing is done in two directions, so that all nucleotide DNA targets can be known. After going through the contig, reverse complement process, DNA sequences without gaps are obtained which are called sustainable areas or consensus. The results of the Multiple Sequence Alignment of some 16S Auky Island Padaido District in Biak Numfor Papua Regency isolates detected showed that Auky Island Padaido District in Biak Numfor Papua Regency local isolates categorized as ABNP8, ABNP9, ABNP11, ABNP12, and ABNP18 had sustainable areas (conserve) or consensus in the same position but adapted areas that varied from size 16S ± 1500bp. The colored shaded area in Fig. 4 above shows the same nitrogenous bases in all four isolates. Thus the results of preliminary analysis showed that DNA sequences from the 16S *B. thuringiensis* marker gene in Auky Island Padaido District in Biak Numfor Papua Regency local isolates were detected as different isolates, in addition to those previously discovered by many researchers.

**Phylogenetic tree construction results**

Phylogenetic tree construction is intended to know phylogeny and show evolutionary relationships between organisms (kinship relationships), where the sample has properties similar to the closest relatives because of a kind. To
carry out phylogenetic tree construction analysis in this study, outgroups were determined at different levels with all sequences compared\textsuperscript{19}.

Based on the input data in the MEGA 6 program, a phylogenetic tree construction was obtained which illustrates the similarity between fellow Bacillus thuringiensis Auky Island Padaido District in Biak Numfor Papua Regency isolates and the kinship of several isolates in the gene bank database. In this Fig., 4 large clusters were formed, namely from various isolates randomly located in one cluster. It can be seen in Fig. 3 that based on the 16S gene markers, successfully placed isolates ABNP8, ABNP9, ABNP11, ABMP12, and ABNP18 are in one cluster with 84% safety percentage showing credibility, consistency and degree of robustness from its position in the phylogenetic tree. Specifically ABNP8 has specificity because it is separate and has its own branching between other Jayapura local isolates namely ABNP 9, ABNP11, ABNP12, and ABNP18. It is very interesting that there is a possibility that ABNP8 isolates are really different, have diversity or combination alleles that cause variation, while isolates in the NCBI gene database such as Kerstersia, Bordetella and proteus in other clusters. Likewise, Bacillus cereus and Bacillus subtilis are located in the same cluster with a similarity percentage level reaching mencapai 80%. Phylogenetic trees are also intended to find out phylogeny, where samples that have similar properties will be in an adjacent position because they have identical DNA sequences or have many similarities. In this phylogenetic tree, outgroups are determined to be at different levels with all the sequences compared, namely *Trametes hirsuta*, in different clusters from the 3 previous clusters, because *Trametes hirsuta* has been positioned as outgroup samples that are indeed different at the species level. The phylogenetic tree was constructed using the Neighbor Joining algorithm method, and a bootstrap value of 1000

CONCLUSION

Based on the results of research that has been done, it can be concluded as follows: The isolates of ABNP 8, ABNP9, ABNP11, ABMP12 and ABNP18 have been detected as local isolates from Auky Island Padaido District in Biak Numfor Papua Regency that have great potential as bioinsecticides to control and kill the Anopheles mosquito larvae. The similarity of the five isolates reach 84% based on MEGA7 program. Of the five isolates, ABNP 8 isolate had an unique characteristics and diversity compared to the other four isolates.

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CONFLICT OF INTEREST
The authors declare that there is no conflict of interest.

AUTHORS’ CONTRIBUTION
All listed author(s) have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

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DATA AVAILABILITY
All datasets generated or analyzed during this study are included in the manuscript.

ETHICS STATEMENT
This article does not contain any studies with human participants or animals performed by any of the authors.

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