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The genetic diversity of *Drosophila* flies based on cytochrome-c oxidase subunit 1 gene from North Sulawesi

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Abstract

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Background: *Drosophila* sp. is a fruit fly species that can spread quickly worldwide. A short life cycle, small body size, and fast adaptation to new habitats allow fruit flies to live in various parts of the world. This study characterized the genetic variation of the cytochrome oxidase c subunit (CO1) gene in *Drosophila* species from different North Sulawesi regions.

Methods: Fruit fly samples were collected from six districts: Central Minahasa, Southeast Minahasa, South Minahasa, North Minahasa, Bolaang Mongondow, and Sitaro. DNA extracted from thoracic tissue. The COI gene was amplified by polymerase chain reaction (PCR) and sequenced by the Sanger method. Sequence characterization using BioEdit and MEGA XI programs.

Result: The results showed that the consensus CO1 gene sequence length was 688 bp to 700 bp. Divergent evolution based on disparity analysis showed CO1 Bolaang had the farthest sequence characteristic differences from the other five CO1 genes in North Sulawesi. Genetic distance analysis showed that Bolaang's gene sequence has the farthest genetic distance. CO1 gene consensus alignment analysis with ClustalW showed high genetic variation. The phylogenetic construction showed that CO1 Bolaang had the most significant differences in sequence characteristics from the other five sequences in a monophyletic group with different nodes. Phylogenetic reconstruction with the 21 most similar sequences from BLAST showed similarities in the four *Drosophila* species, namely *D. atriplex*, *D. melanogaster*, *D. lacteicornis*, and *D. pandora*.

Conclusion: Based on the CO1 gene, there are significant variations in fruit flies in North Sulawesi.

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diversity genetics; phylogeny; CO1; *Drosophila*; Sulawesi Utara



Introduction

Drosophilae is widely used in the study of biological phenomena, including developmental biology and genomics [1,2], microbiome [3], human disease mechanism [4, 5], and genetic studies [6, 7]. Understanding intraspecies and interspecies phylogenetic relationships within this family greatly influences studying these biological phenomena. However, studies on the genetic diversity of fruit flies are still underreported, primarily based on their habitat origin and biogeographic regions. There are still no research reports on the genetic diversity of fruit flies in Indonesia, more specifically in the biogeographical region of Wallacea. Fruit flies are found in tropical to subtropical areas. This causes high intraspecies genetic variation in fruit flies around the world. Furthermore, the short life cycle, small body size, and high reproduction speed reinforce the process of divergent evolution in fruit flies [8-10].

North Sulawesi Province is the northernmost part of Sulawesi Island, Indonesia. However, Sulawesi is included in the Wallacea zone, which has specific flora and fauna biodiversity, in contrast to the western part of Indonesia, which has oriental biogeographical characteristics, and the eastern part of Indonesia, which has Australian biogeographical characteristics [11, 12]. The species endemism in Sulawesi is relatively high. Many endemic fruit plant species are also food sources for fruit flies. Some endemic plant species include Pakoba (*Syzygium luzonense* (Merr.)), Nutmeg, Beringin Minahasa (*Ficus minahassae*), Langsat forest (*Lansium domesticum* L.), Forest Mangosteen (*Garcinia mangostana* L.), tome (*Flacourtia inermis* L.) etc. Insects can coevolve with plant food sources, thereby affecting their genetic diversity. It is suspected that this process took place in *Drosophila* sp. Research on the genetic diversity of fruit flies is important in North Sulawesi, a region that produces various types of local fruits.

Morphological characteristics have historically been used to identify taxonomy and genetic diversity. In the last two decades, molecular markers for animal identification have been widely used, namely the Cytochrome oxidase subunit 1 (CO1) gene. DNA barcoding data can be used for more than just taxonomic studies. It will have significant effects on a wide range of biological disciplines, such as ecology (quick examination of the food chain and biodiversity), conservation biology (surveillance of protected species), biosafety (identifying invasive pest species early), pharmaceuticals (identifying medically important infections and their carriers), and pharmacology (the detection of active substances). The CO1 gene has been widely used for the molecular marker in the *Drosophila* family, so its accuracy is believed [13-15].

Based on previous morphological studies, there were differences in morphology, especially eye color, abdomen color, wing length, and average body length of *Drosophila* sp. from island locations and the Sulawesi mainland. The difference in the average body length of fruit flies affects the shape and size of the reproductive organs. This can lead to reproductive isolation [16]. Geographical isolation has caused morphological variations of *Drosophila* sp. Demographic history, geographic isolation, and environmental factors together form the genetic structure of the population of *D. melanogaster* [17, 18]. *Drosophila* sp. can coevolve with food source plants in chemoreceptors [19].

However, little is known about research on the genetic diversity of fruit flies in the Sulawesi (Wallacea) biogeographical zone. Therefore, a study was conducted to characterize the CO1 gene of the North Sulawesi fruit fly from various locations. The findings of this study provide excellent material for evolutionary studies and phylogenetic relationships, particularly at the species level of *Drosophila* sp. in specific areas.

Methods

Collection samples *Drosophila* sp.

Adult *Drosophila* sp. samples were obtained directly from several locations in North Sulawesi (Figure 1). At each location, ten adult individuals of *Drosophila* sp. were isolated and preserved in 95% alcohol for 24 hours. After 24 hours, the fruit flies were transferred to a new bottle containing 95% alcohol. Sample collection was carried out from March to October 2022.

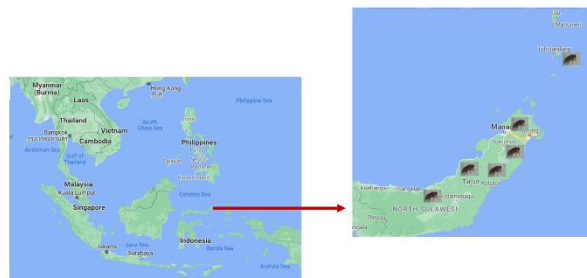


Figure 1: Map of Indonesia, the location of the *Drosophila* sp. collection in North Sulawesi Province is labeled with a picture of a *Drosophila* sp.

Extraction, Purification of DNA, PCR, and Sequencing.

Fruit fly genomic DNA was extracted from the thoracic tissue (Fig. 2). The tissues were homogenized (SPEX Sample Prep 1600 Mini G) before being digested overnight at 56°C. A Quick-DNATM Miniprep Kit from Zymo Research USA was used to extract DNA according to the manufacturer's instructions. Extracted total DNA was stored at -20°C for use in the CO1 gene amplification stage. The 700 bp CO1 gene fragment was amplified using the primer pairs listed below: HCO, 5'-TAAACTTCAGGGTGACCAAAAAATCA-3' and LCO,

5'GGTCAACAAATCATAAAAAGATATTGG-3' (Folmer et al., 1994) [20]. The PCR process used 2x MyTaq HS Red Mix Bioline (USA), DNA template, ddH₂O and CO1 primer. The PCR component was 2x MyTaq HS Red Mix Bioline (USA) 25 µl; primer forward 1 µl; primer reverse 1 µl; DNA template 2 µl, ddH₂O 21 µl. PCR conditions was denaturation 94°C (60 seconds), Annealing 50°C (30 seconds), Extension 72°C (30 seconds) and Final Extension 72°C (60 seconds) [21].



Figure 2: *Drosophila* sp. samples used for DNA extraction, 200x with Hirox KH8700 stereomicroscope.

Amplicon Visualization and Sequencing

Amplicons of the CO1 gene of *Drosophila* produced at the PCR were visualized using conventional electrophoresis (1% TBE agarose—M, 100bp ladder (loaded 2.5 L). Sequencing uses ABI PRISM 3730xl Genetic Analyzer developed by Applied Biosystems, USA, through Singapore FIRST BASE sequencing services. The sequencing output in a seq file was analyzed using MEGA XI and BioEdit.

Data analysis

Sequence Analysis

Partial sequences of the CO1 gene 600-700 bp long were collected from 6 North Sulawesi, Indonesia districts. Forward and reverse sequences were analyzed using BioEdit to obtain consensus sequences. Each sequence is first translated into an amino acid sequence to detect and remove any stop codons in the middle of the sequence. In addition to careful manual checking, sequence checking, and editing were done using the BioEdit software (Hall, 1999). Sequence characteristics were analyzed for the CO1 gene, and the amino acids were performed using BioEdit. Each sample sequence was validated using the GenBank NCBI's online Basic Local Alignment Search Tool (BLAST) (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>).

For phylogenetic tree reconstruction, sums of the 21 accessions from GenBank NCBI were selected as in-group and outgroup (Table 1). In MEGA XI software

version 10.2.6, multiple sequences were reassigned using the ClustalW Algorithm [22].

No	Accession Number	Species	Origin
1	Select seq MK659835.1	<i>Drosophila pandora</i>	China
2	Select seq OK037194.1	<i>Drosophila atriplex</i>	Malaysia
3	Select seq MK659836.1	<i>Drosophila parapallidosa</i>	China
4	Select seq MK659805.1	<i>Drosophila ananassae</i>	USA
5	Select seq MK659811.1	<i>Drosophila bipectinata</i>	China
6	Select seq MK659826.1	<i>Drosophila malerkotliana</i>	France
7	Select seq AY757282.1	<i>Drosophila parabiptectinata</i>	Canada
8	Select seq OK175852.1	<i>Drosophila ananassae</i>	Khandesh
9	Select seq AY757280.1	<i>Drosophila pseudoananassae</i>	Canada
10	Select seq MK455900.1	<i>Drosophila</i> sp.	India
11	Select seq AB907180.1	<i>Drosophila melanogaster</i>	India
12	Select seq AB830535.1	<i>Drosophila lacteicornis</i>	Japan
13	Select seq MK659817.1	<i>Drosophila ficusphila</i>	China
14	Select seq AB669729.1	<i>Drosophila bocki</i>	Japan
15	Select seq MK659806.1	<i>Drosophila anomalata</i>	China
16	Select seq MK659806.1	<i>Drosophila anomalata</i>	China
17	Select seq AB669702.1	<i>Drosophila neosahinai</i>	Japan
18	Select seq AB669709.1	<i>Drosophila tani</i>	Japan
19	Select seq AB669704.1	<i>Drosophila rufa</i>	Japan
20	Select seq AF200849.1	<i>Drosophila simulans</i>	USA
21	Select seq AB669699.1	<i>Drosophila lacteicornis</i>	Japan

Table 1: BLAST results for phylogeny construction with *Drosophila* sp. CO1 sequences from North Sulawesi.

Phylogenetic tree reconstruction

The Maximum Composite Likelihood model was used to compute the CO1 gene disparity matrix of the Sulawesi fruit fly. The North Sulawesi fruit fly CO1 gene sequence was aligned using the ClustalW model in the MEGA X program version 11.2.6 and online at the European Bioinformatics Institute (<https://www.ebi.ac.uk>). On 27 sequences, a phylogenetic tree reconstruction based on partial CO1 gene sequences was performed (in-group and outgroup accessions obtained from GenBank). MEGA X version 11.2.6 was used to reconstruct the phylogenetic tree. Meanwhile, the phylogenetic tree reconstruction methods used are Neighbor-Joining (N.J.) and Maximum Likelihood (ML) (ML). The NJ reconstruction was computed using the Kimura 2-Parameter substitution model (K2P) [22]. Bootstrap 1000 replicates were used to evaluate phylogenetic tree reconstruction.

Results

Characterization of Sequences Obtained

The amplicons of the sequenced results were classified as good, as evidenced by the chromatogram of the sequenced results, which showed that the bands of the nitrogenous base types did not coincide much (Fig. 3). Sequencing of the CO1 gene sequence of fruit flies revealed an average length of 600 bp – 700 bp (Table 2). Characteristics of *Drosophila* sp. DNA CO1 gene has a G+C ratio of 30.20% to 32.82% (Appendix 1). At the same time, the A+T ratio is between 67.01% and 65.78%. The consensus CO1 gene molecular length is between 688 bp to 716 bp. All the characteristics of the *Drosophila* sp. CO1 gene from various regions in North Sulawesi

analyzed with the BioEdit Program are shown in Appendix 1.

No.	Samples	Sequence length (bp)		Consensus sequence length (Bio Edit v.7.2.5 & MEGA 11)
		LCO 1490	HCO 2198	
1	L1 (Minahasa)	683	700	716
2	L2 (Minteng)	683	696	688
3	L3 (Minut)	682	684	692
4	L4 (Minsel)	685	684	713
5	L5 (Sitaro)	685	684	697
6	L5 (Bolaang)	684	687	685

Table 2: The sequence length of the *Drosophila* sp CO1 gene from North Sulawesi.

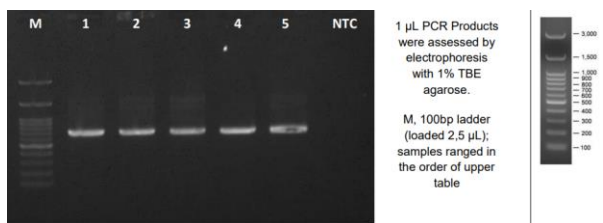


Figure 3: *Drosophila* sp. from North Sulawesi CO1 DNA profile amplified by primers LCO 1490 and HCO 2198. Electrophoresis of 1 L PCR products with 1% TBE agarose M, 100bp ladder (loaded 2.5 L) was performed; 1): 1: L1 (Minahasa), 2: L2 (Southeast Minahasa), 3: L3 (North Minahasa), 4: L4 (South Minahasa), 5: L5 (Siau Tangelandang Biaro), 6: L6 (Bolaang Mongondow).

Use of sequence analysis model Genetic Disparity and Distance Matrix

The disparity matrix analysis was used to estimate divergent evolution between sequences. The number of base substitutions per site between sequences is displayed. The Maximum Composite Likelihood model was used in the analysis. This investigation involves five nucleotide sequences. The included codon positions are 1st+2nd+3rd+Noncoding. According to the disparity matrix analysis results, the longest distance was found between the Bolaang CO1 gene sequence and the other five sequences. In contrast, the closest distances are CO1 Minsel and CO1 Sitaro (Table 3).

	1	2	3	4	5	6
1. Minteng		1.000	1.000	0.066	0.084	0.000
2. Minduk	0.000		1.000	0.204	0.268	0.000
3. Minut	0.000	0.000		0.022	0.016	0.000
4. Sitaro	0.138	0.056	0.197		1.000	0.000
5. Minsel	0.141	0.047	0.201	0.000		0.000
6. Bolmong	4.267	4.523	4.562	5.634	5.723	

Table 3: Disparity Matrix of CO1 Gene DNA, *Drosophila* sp. from North Sulawesi.

The ratio of genetic differences between species or populations is depicted by genetic distance. The Bolaang Mongondow CO1 gene sequence had the most significant genetic distance between the CO1 genes of the Sulawesi fruit fly. The Minsel CO1, Sitaro CO1, and

Main Minahasa CO1 genes followed this. CO1 Sitaro and CO1 Minsel have the closest genetic distance (Table 4).

	Minteng	Minduk	Minut	Sitaro	Minsel	Bolaang
Minteng	0.0000	0.0029	0.0227	0.1140	0.1105	1.2270
Minduk	0.0029	0.0000	0.0335	0.1297	0.1357	1.2377
Minut	0.0227	0.0335	0.0000	0.0924	0.0889	1.2208
Sitaro	0.1140	0.1297	0.0924	0.0000	0.0000	1.2680
Minsel	0.1105	0.1357	0.0889	0.0000	0.0000	1.2617
Bolaang	1.2270	1.2377	1.2208	0.1268	0.1261	0.0000

Table 4: The genetic distance of *Drosophila* sp CO1 gene from North Sulawesi.

Furthermore, Maximum Likelihood (ML) analysis was used to estimate nucleotide substitution. Each entry represents the likelihood of substitution (r) from one base (row) to another base (column). The Tamura-Nei model was used to estimate turnover patterns and rates. Transitional substitution levels are bold, and transversion substitutions are in italics. When evaluating r, the relative instantaneous value must be taken into account. For clarity, the sum of the r values was set to 100. The nucleotide frequencies were A = 30.43%, T/U = 37.03%, C = 15.57%, and G = 16.97%. (Table 5). To estimate the ML value, the tree topology is calculated automatically. The maximum possible Log for this calculation is -2222.697. Six nucleotide sequences are used in this analysis. The included codon positions are 1st+2nd+3rd+Noncoding. The final dataset contains 736 positions in total. MEGA X1 was used for evolution analysis.

Base	A	T/U	C	G
A	-	8.97	<i>3.77</i>	6.88
T/U	<i>7.37</i>	-	9.58	4.11
C	<i>7.37</i>	22.78	-	4.11
G	12.34	<i>8.97</i>	<i>3.77</i>	-

Table 5: Substitution Matrix Maximum Likelihood Estimate

Alignment Analysis with ClustalW

The CO1 gene sequences of the North Sulawesi fruit fly were aligned using the ClustalW, model ClustalW is a common method for aligning multiple nucleotide sequences or homologous proteins. ClustalW uses the progressive alignment method for multi-sequence alignment. In this case, the sequence with the highest alignment score is aligned first. Then, progressively more distant sequence groups are aligned until a global alignment is collected. The alignment results indicated the sites of nucleotide differences between the 6 CO1 gene sequences of fruit flies from North Sulawesi. The Bolaang CO1 gene sequence showed the highest nucleotide variation. The Minteng CO1, Minduk CO1, and Minut CO1 gene sequences followed this. The CO1 Minsel and CO1 Sitaro gene sequences show many nucleotide similarities (Figure 4).

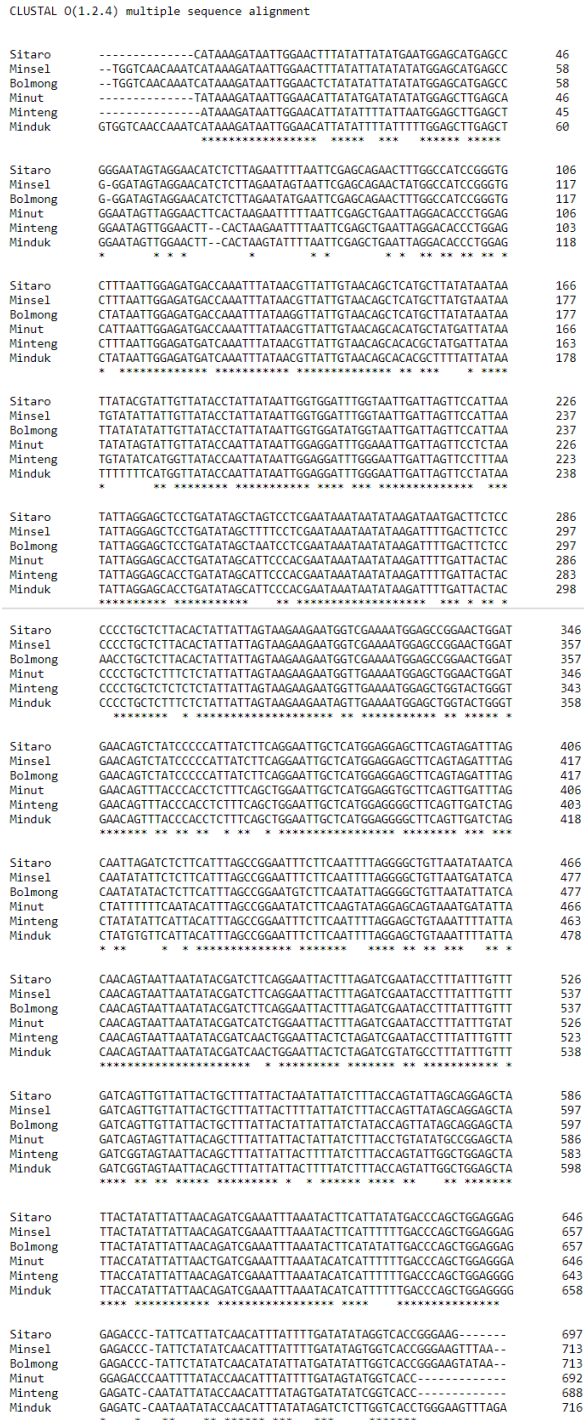


Figure 4: Alignment of CO1 gene sequences of North Sulawesi Fruit fly Figure 4. Alignment of *Drosophila* sp. CO1 gene sequences from North Sulawesi.

Characteristics of the amino acid sequences that make up CO1 protein

The characteristics of the CO1 gene protein have a molecular weight between 58939.43 Daltons to 62173.10 Daltons. The highest molecular weight consensus sequence is the Mansell CO1 gene, while the lowest is

the Mitra CO1 gene. Complete characteristics of CO1 amino acids from North Sulawesi fruit flies based on analysis with the BioEdit Program are shown in Appendix 2.

CO1 Protein Genetic Distance Matrix

The genetic distance of the amino acid protein CO1 *Drosophila* sp. from Sulawesi shows that Bolaang CO1 has the most significant genetic distance with the other five CO1s. Then CO1 Minahasa Tenggara, CO1 Minahasa and CO1 South Minahasa. The closest genetic distance is directed by South Minahasa CO1 and Siau Tagulandang Biaro CO1 (Table 6).

	Minteng	Minduk	Minut	Sitaro	Minsel	Bolaang
Minteng	0.0000	0.0221	0.0936	0.2318	0.2267	1.2678
Minduk	0.0221	0.0000	0.1032	0.2451	0.2534	1.2586
Minut	0.0936	0.1032	0.0000	0.2206	0.2155	1.3104
Sitaro	0.2318	0.2451	0.2206	0.0000	0.0049	1.3327
Minsel	0.2267	0.2534	0.2155	0.0049	0.0000	1.3297
Bolaang	1.2678	1.2386	1.3104	1.3327	1.3297	0.0000

Table 6: Genetic Distance of CO1 amino acids *Drosophila* sp from North Sulawesi.

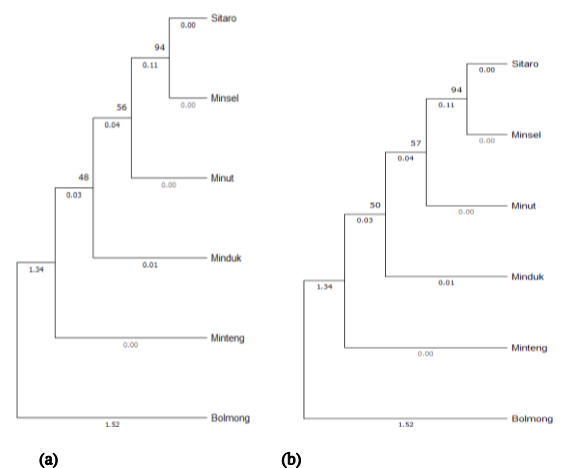


Figure 5: North Sulawesi *Drosophila* sp. phylogeny construction (a) Neighbor-Joining method, (b). Minimum Evolution; boosted 1000x, 2-Parameter Kimura model

Construction of Phylogeny

The North Sulawesi fruit fly CO1 gene phylogeny was constructed using the neighbor-joining method, the minimum evolution method, and the maximum composite likelihood substitution model. The phylogenetic tree topology resulting from the construction of the two methods or three methods was not significantly different. The phylogenetic construction showed *Drosophila* sp. from Bolaang as an outgroup. *Drosophila* sp. Sitaro, Minsel, Minut, Minduk, and Minteng were in one monophyletic group. *Drosophila* sp. sitaro forms a node with minsel. *Drosophila* sp. minut is in a monophyletic group with sitaro and minsel but at different nodes, so do *Drosophila* sp. minduk and minteng. Based on the phylogenetic tree formed, *Drosophila* sp. minsel has the

closest phylogenetic relationship with *Drosophila* sp. Sitaro (Fig. 5).

Phylogenetic construction based on amino acids, CO1 minut was in the outgroup while CO1. CO1 Minteng, CO1 Minduk, CO1 Minsel, and CO1 Sitaro formed a monophyletic group with different nodes. Minteng, Minduk, Minsel, and Sitaro were in the same node. CO1 gene DNA phylogeny construction differed from the construction based on amino acids (Fig. 6).



Figure 6: Phylogeny based on CO1 Neighbor-Joining/UPGMA amino acid sequences using BioEdit.

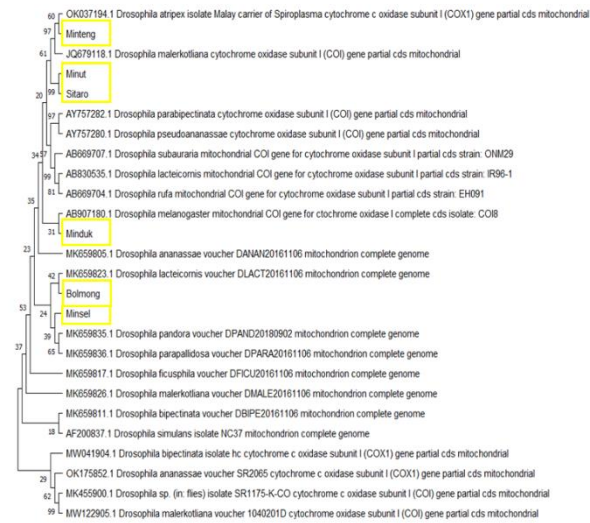


Figure 7: Position of the Sulawesi *Drosophila* sp. based on the CO1 gene and NCBI BLAST results.

Discussion

The visualization of the amplicon with a stable band on electrogram supports amplification of the *Drosophila* sp. CO1 gene from various regions in Sulawesi was successfully carried out using thoracic tissue by modifying proteinase K soaking time which effectively produces high purity and concentration in insects [23, 24]. Nucleotides sorted by Sequencing produce Forward and Reverse sequences which are well proven by the chromatograms of each type of nitrogenous bases that are not overlapping visualized using the BioEdit and MEGA X programs. The consensus sequences are at a length of 685 to 716 bp, indicating the length of the CO1 gene sequence [25, 26, 27]. The amplification and sequencing results were confirmed as the CO1 gene. Consensus sequences are the most frequently occurring subsequences in a DNA sequence [28]. This sequence is useful for determining the location of the protein.

Consensus sequences were determined using the BioEdit Program. Consensus sequences were verified for accuracy by BLAST on the NCBI website. The BLAST results showed that the most similar sequence was *Drosophila* sp. Alignment of the six sequences of fruit flies from North Sulawesi using the Clustal W method showed polymorphic sites. The most polymorphic sites were found in the Bolaang fruit fly compared to the CO1 sequences of other North Sulawesi fruit flies. The genetic range of CO1 amino acids in fruit flies from North Sulawesi is consistent with the genetic range of CO1 DNA sequences. Where *Drosophila* sp. from Bolaang showed the farthest genetic distance with five other *Drosophila* sp. However, *Drosophila* sp. Sitaro and *Drosophila* sp. Minsel had closest phylogenetic relationship based on amino acids encoded by the CO1 gene.

Phylogenetic reconstruction based on the sixth CO1 gene of *Drosophila* sp. from North Sulawesi was built using two methods, namely Neighbor-Joining and Minimum Evolution, to compare and ascertain the topography of the phylogenetic tree formed [29, 30]. The research results proved that the phylogenetic tree's topography was not significantly different between the two methods. This proves that using the CO1 gene accurately determines the position of the species *Drosophila* sp. Thus, the position and phylogenetic relationship of each *Drosophila* sp. from North Sulawesi based on the CO1 gene was ascertained. *Drosophila* sp. from Bolaang showed the highest variation in the CO1 gene with other sequences, so it occupies the outgroup on both phylogeny trees.

Interestingly, this study found that *Drosophila* sp. Sitaro and *Drosophila* sp. Minsel are in the same node. Geographically Sitaro and Minsel are in different areas. Thus it or both have the closest phylogenetic relationship compared to other *Drosophila* sp. in North Sulawesi. *Drosophila* sp. Sitaro is in the islands, while *Drosophila* sp. Minsel is on the mainland of North Sulawesi. This indicates that it is necessary to use other marker genes to analyze the phylogeny of *Drosophila* sp.

The phylogenetic construction based on the amino acids of the six CO1 genes of fruit flies from North Sulawesi is not much different from the phylogenetic construction based on DNA sequences. This shows the consistency of the phylogeny formed based on DNA and the amino acids encoded by the CO1 gene. Both phylogenetic constructs based on CO1 gene DNA and amino acids are consistent with the results of genetic distance analysis.

Reconstruction of the fruit fly phylogeny from North Sulawesi with 21 sequences from the BLAST results showed variations in species similarity. *Drosophila* sp. Minteng forms a node with *D. atriplex* (OK037194.1);

thus, *Drosophila* sp. Minteng based on the CO1 gene has the closest resemblance to *D. atriplex*. *Drosophila* sp. minut and Sitaro from the same node. *Drosophila* sp. Minahasa forms a node with *D. melanogaster* [AB907180.1]. Based on the CO1 gene, *Drosophila* sp. Minahasa has the closest resemblance to *D. melanogaster*. *Drosophila* sp. Bolaang forms a node with *D. lavteicomis* [MK659823.]. Thus, based on the CO1 gene, Bolaang fruit fly has the closest resemblance to *D. lavteicomis* [MK659823.1]. At the same time, *Drosophila* sp. from Minsel forms a node with *D. pandora* [MK659835.1] and *D. parapallidosa* [mk659836.1] (Figure 7), which means that the CO1 gene of the Minsel fruit fly was similar, closest to the two species.

This study found that the intraspecies variation of the CO1 gene in fruit flies in North Sulawesi was high. Future research is needed using multi-gene barcodes to confirm the status of fruit fly species from North Sulawesi. This study also proved that the CO1 gene has not explicitly confirmed intraspecies differences; what is proven states that clearly) it also means that the hypothesis of intraspecific has failed due to geographical variations. This is the first report on genetic variation and evolutionary relationship reconstruction based on the CO1 gene of *Drosophila* sp. from North Sulawesi. Although they have high morphological similarities, the genetic variation based on the CO1 gene of fruit flies from North Sulawesi is high demonstrated this point in this paper. The findings of this research reveal that a molecular approach based on the partial sequence of the CO1 gene supports morphological identification results in *Drosophila* sp. found in North Sulawesi. Future studies can use a morphological approach to strengthen the identification of *Drosophila* sp.

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Author Contributions

Herry Maurits Sumampouw: constructed the experiments, interpreted the results, and review the article; Mokusuli Yermia Samuel: constructed the experiments, produced the report, statistically analyzed the data, generated illustrations and wrote the article

Conflicts of interest

The authors declare no conflict of interest.

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