Biometric and genetic differences in kelabau (*Osteochilus* spp.) as revealed using cytochrome c oxidase subunit 1 [version 2; peer review: 2 approved, 1 approved with reservations]

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Abstract

**Background:** Kelabau (*Osteochilus* spp.) is a freshwater fish commonly found in the rivers of Riau, Indonesia. Researchers believe that these are *Osteochilus kelabau*; however, accurate taxonomic determination of these fish in Riau waters has not been made. The purpose of this study was to facilitate the identification of the kelabau based on its morphology and genetics using biometric and cytochrome c oxidase subunit 1 (*CO1*) analyses, respectively.

**Methods:** Fish samples were collected from the Siak, Kampar and Rokan rivers in Riau Province, Indonesia. The DNA of 90 fish was extracted from the caudal fins using a DNA extraction kit, after which it was amplified using primers Fish-F1 and Fish-R1. Sequencing was conducted by Applied Biosystems Macrogen Korea, and the DNA sequences were then edited and aligned using MEGA v. 7. All samples were BLAST-searched for identification using the National Center for Biotechnology Information and BOLD System. Phylogenetic trees were constructed, and the similarity index was calculated using accession numbers AP011385.1 and KC631202.1 in GenBank.

**Results:** Analysis of the consensus barcode sequence for 86 species revealed a high percentage of barcode matches (96%–97% in GenBank and 96.6%–96.76% in the BOLD System). The nucleotide distance between groups of kelabau from the different rivers based on the Kimura 2-parameter model gave the following results: 0.05% between groups from the Siak and Kampar rivers, 0.09% between those from the Siak and Rokan rivers and 0.05% between those from the Kampar and Rokan rivers. The nucleotide distance between the groups in the Siak (0.09%), Kampar (0.00%) and Rokan (0.10%) Rivers indicated that the kelabau in those rivers were related to each other.

**Conclusions:** Based on the results of the research data using *CO1* and biometric analyses, the kelabau were confirmed to be *O. melanopleurus*.
Keywords
DNA barcoding, Kelabau Fish, Common Rivers of Riau, Population Structure

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Author roles: Asiah N: Data Curation, Formal Analysis, Investigation, Project Administration, Software, Writing – Original Draft Preparation, Writing – Review & Editing; Junianto J: Conceptualization, Supervision, Validation, Writing – Review & Editing; Yustiati A: Conceptualization, Methodology, Supervision, Validation, Writing – Review & Editing; Sukendi S: Conceptualization, Methodology, Supervision, Validation, Writing – Review & Editing; Fahmi MR: Conceptualization, Formal Analysis, Investigation, Methodology, Validation, Writing – Review & Editing; Muchlisin ZA: Validation, Writing – Review & Editing; Kadapi M: Methodology, Supervision, Validation

Competing interests: No competing interests were disclosed.

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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First published: 12 Feb 2019, 8:177 (https://doi.org/10.12688/f1000research.17319.1)
Introduction
Kelabau are ancient fish belonging to genus *Osteochilus* of family Cyprinidae. Kelabau fish are distributed throughout Thailand, Vietnam, Peninsular Malaysia, Borneo and Sumatra. In Sumatra Island in Indonesia, the fish is commonly found in the Siak, Kampar and Rokan rivers in Riau Province.

According to local fishers in Riau, kelabau are divided into two types on the basis of morphology; although, there is no detailed information about these fish types. The local fishers said that the first type of kelabau fish is the fish found in this study. The morphological characteristic of this fish is brownish body, with brighter bottom, dark hazy blotches present above the pectoral fins, and the size ranges from 15 to 2,778.93 gr. The second type of fish is larger sized and yellowish color. However, during the study time, from 2017 to 2018, this type was not found. Thus, a study was needed to identify the species using morphological and molecular methods to determine these types in the Siak, Rokan and Kampar rivers. Identification of any species using morphological traits can be difficult and can lead to errors; therefore, owing to morphological similarities among *Osteochilus* spp., molecular markers, such as DNA barcodes, are important to identify the fish species uses a specific sequence region (i.e. *cytochrome c oxidase subunit 1* (*CO1*)) to identify a species and is a technique that can identify taxonomic units as well as biodiversity for determining species of several organisms. Unlike molecular phylogeny used to determine relationships among species, the purpose of DNA barcoding is to identify unknown or undetermined species into phylogeny. The common mitochondrial (mt) DNA region used as a barcode in protists and animals comprises 600 bp. In addition, *CO1* is one of the genetic markers used to identify insects, birds, primates and fish to species. *MiDNA CO1* is selected as a target in DNA barcoding because it is a highly conserved site. This method has advantages over the morphological identification approach in that it is fast, reliable and it can be used for all types of samples because it uses a single gene along with mutations in the nucleotides to acknowledge the taxonomic features of each species.

The study on DNA barcoding for freshwater fish has been widely practiced in various countries, including Nigeria, Malaysia, Philippines, Canada, and Indonesia. The method has been successfully validated for the taxonomic status within *Rasbora* in Lake Laut Tawar; *Anguillidae* in Aceh waters; Ornamental fish from Peat lands; *Channidae* in Peninsular Malaysia, Sarawak, Sumatera, Borneo, Myanmar, Vietnam, India, Germany, Singapore and the United Kingdom; and *Cichlidae* in northeastern Nigeria; therefore, it can be used to equally successfully validate the taxonomic unit of the kelabau using its morphology supported by molecular data. This information is crucial for designing a remedial course of action about the conservation strategy for this species in the Siak, Kampar and Rokan rivers in Riau Province, Indonesia.

Methods
Ethics
The study population was collected and sampled according to the guidelines on the use of living organisms for research from the Laboratory of the Faculty of Fisheries and Marine, Riau University, Indonesia.

Sampling sites and collection
A total of 90 kelabau (30 fish from each river) were collected from the Siak, Kampar and Rokan rivers (Figure 1). Fish were caught using a Gill net 3 m deep and 20 m long with a 12.7-cm mesh. The Gill nets were installed in the river water close to the riverbank and remained for 24 h from 08:00 to 08:00 the following day. The fish collected were counted using hand-counter and cleaned using freshwater. A number of 50-mm caudal fin tissue samples were taken using a sterile scissors and preserved in ethanol, after which a photo of each fish sample was taken for documentation using a digital camera.

All samples were preserved in 3-kg sample bags which were labeled according to site location, date and serial number. Before preservation, the fish samples were injected with 10% formalin. The fish samples were then transported to the laboratory for further evaluation. The morphologies of the collected fish were identified up to species level using the identification book produced by the Indonesian Institute of Sciences ichthyology museum. The fish morphologies observed were
length, color, shape of scales, mouth shape, barbels, number of fins and special marks on the body.

**Biometrics**

Biometric analyses were used to measure morphological characteristics in this study. This tool is considered conventional for identifying organisms. Molecular identification using CO1 gene sequences has been supported for providing additional organism classification.

**DNA isolation and amplification**

DNA was extracted using the spin-column method from the gSYNC DNA Extrusion Kit (Geneaid Catalogue No. GS 300, Taiwan). The extracted DNA was then transferred to a 1X Tris–borate ethylenediaminetetraacetic acid (TBE) solution with a 1.5% agarose gel and Pegreen gel dye (PEQLAB Biotechnologies GmbH, Erlangen, Germany). The quantity of DNA was visualized with the help of a GeneQuant Spectrophotometer by adding 78 μL nuclease-free water in a cuvette along with 2 μL DNA. The DNA was then amplified using the universal primer Fish-F1 (5’-TCA-ACC-AAC-CAC-AAA-GAC-ATT-GGC-AC-3’) and Fish-R1 (5’-TAG-ACT-TCT-GGG-TGG-CCA-AAG-AAT-CA-3’) with a target of 707 bp and 655 bp, respectively. The amplification thermocycling conditions as follow: the PCR condition using pra PCR (94°C for 5 min), 35 cycles of denaturation (94°C for 30 s), annealing (56.6°C for 30 s) and extension (72°C for 30 s), followed by post-PCR extension (72°C for 5 min) and hold (4°C for 5 min). PCR results were analyzed using 1.5% agarose gel at 100 V to assess the bands, and only the clear products were sent to Applied Biosystems Macrogen Korea for sequencing.

**Controlling molecular samples and sequence quality**

The PCR amplicon was 707 bp, which implied that no sequence of DNA was derived from mtDNA nuclear mitochondrial DNA segments (NUMTs), because a NUMT barely
reaches 600 bp. The selected CO1 sequences were entered into GenBank and the BOLD System databases to compare the alignment of nucleotide sequences and 99%–100% values with that with no insertions/deletions. All sequences were aligned using ClustalW with MEGA v.7.

**Data analysis**

**Blasting of CO1 by NCBI (GenBank) and BOLD System (online)**

The entire nucleotide sequence obtained from the sequence chromatogram was assembled using DNA Baser Assembler, aligned and then analyzed using MEGA 7. It was further aligned (multiple alignments) using the reference NCBI GenBank accession numbers AP011385.1 and KC631202.1. Similarly, the percentages of CO1 sequences were blasted using NCBI Blast and BOLD Systems databases.

**Nucleotide variations**

Nucleotide variations among samples were analyzed using dnaSP v.5. The parameters of these calculations were haplotype number, variable site, parsimony site, haplotype diversity and nucleotide diversity.

**Phylogenetic tree**

Phylogenetic trees were estimated using all samples from the three populations and calculated according to the Tamura-Nei model using MEGA 7.

**Results**

**Morphological identification**

The morphological traits of all kelabau used in this study matched those of *O. melanopleurus*. We used the important morphological traits to identify these fish according to Kottelat *et al.* The morphological characteristics measurement of *O. melanopleurus* showed that the fish have 16–19 branched dorsal rays, the number of scales was ranged from 10.5 to 12.5 in between dorsal origin and lateral line, the number of circum peduncu-lar rows of scale was ranged from 20 to 24 and lips covered with folds and plicae and there was no hard tubule at the tip of the mouth (Figure 2a). This species has one pair of barbels at above and one pair at bottom, dark hazy blotches near above of the pectoral fins. The body is brownish, with the bottom brighter than the top and the type of steroid scales (Figure 2b). Raw biometric data are available on OSF.

**Genetic analysis**

A sequence amplified by Fish-F1 primer was successfully identified in 86 of 90 samples of mtDNA fish. The base length of the CO1 nucleotide obtained from the formulation process and electrophoresis (Figure 3).

Based on genetic analysis using the Tamura-Nei model, there was an unequal distribution of all nucleotides with the following frequencies: adenine (A), 26.73%; thymine (T), 30.44%; cytosine (C), 25.93% and guanine (G), 16.90% (Table 1). The rates ratio between transition and transversion was 10.257 purines and 1.915 pyrimidines, and the overall transition and transversion bias were R= 2.499 based on Tamura-Nei model. The pattern of nucleotide distribution of A, T, C and G was T > A > C > G, but the carps which cultivated in India have the following pattern of nucleotide distribution: T > C > A > G.

The nucleotide distances between nucleotide bases within the groups indicated that the values of the nucleotide base sequences within the fish population were 0.0009, 0.0000 and 0.0010 in the Siak, Kampar and Rokan rivers, respectively. The evolutionary distance between the nucleotides of the groups had a comparative difference in the nucleotide sequences of 0.0005 for fish from the Siak and Kampar rivers, of 0.0009 in those from the Siak and Rokan rivers and of 0.0005 for those from the Rokan and Kampar rivers. Based on the nucleotide distance, the fish were identified as being from the same species (0.06%) (Table 2).

CO1 had 612 conserved sites (98%), 9 variable sites (1.45%), 4 informative parsimony (0.64%) and 5 singleton sites (0.81%). The highest nucleotide variation was in the Rokan river population (0.00100 ± 0.00032); whereas, the Kampar river population had no nucleotide variation based on DnaSP5 calculations (Table 3). Using the NCBI database with accession numbers
Figure 3. DNA amplified using Fish-F1 and Fish-R1 primers; M= Marker 100 bp (Vivantis, Malaysia); 199, 200, 201, 202, 203, 204, 205, 206, 207, 208, 209 and 210= Rokan River; 211= Kampar River.

Table 1. Maximum composite likelihood estimates of the pattern of nucleotida substitution.

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>T</th>
<th>C</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>-</td>
<td>4.03</td>
<td>3.34</td>
<td>22.95</td>
</tr>
<tr>
<td>T</td>
<td>3.54</td>
<td>-</td>
<td>6.57</td>
<td>2.24</td>
</tr>
<tr>
<td>C</td>
<td>3.54</td>
<td>7.72</td>
<td>-</td>
<td>2.24</td>
</tr>
<tr>
<td>G</td>
<td>36.29</td>
<td>4.03</td>
<td>3.34</td>
<td>-</td>
</tr>
</tbody>
</table>

i Bold: different transitional substitutions; italic: transversional substitutions.

Table 2. Nucleotide distances among the populations.

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Siak</td>
<td>-</td>
<td>0.0005</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Kampar</td>
<td>0.0005</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Rokan</td>
<td>0.0009</td>
<td>0.0005</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GenBank (AP011385.1)</td>
<td>0.0367</td>
<td>0.0366</td>
<td>0.0368</td>
<td>-</td>
</tr>
<tr>
<td>GenBank (KC631202.1)</td>
<td>0.0351</td>
<td>0.0350</td>
<td>0.0351</td>
<td>0.0048</td>
</tr>
</tbody>
</table>

i Source: Kimura estimation of 1980: “Evolutionary Divergence over Sequence Pair Between Groups”.

Table 3. Nucleotide variation in mtDNA CO1 of Osteochilus melanopleurus by DnaSP5.

<table>
<thead>
<tr>
<th>Sampling location</th>
<th>Number of sites</th>
<th>Number of sequences</th>
<th>Haplotype number</th>
<th>Variable site</th>
<th>Parsimony site</th>
<th>Haplotype diversity</th>
<th>Nucleotide diversity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Siak</td>
<td>621</td>
<td>28</td>
<td>6</td>
<td>6</td>
<td>2</td>
<td>0.439 ± 0.114</td>
<td>0.00090 ± 0.00029</td>
</tr>
<tr>
<td>Kampar</td>
<td>621</td>
<td>30</td>
<td>1</td>
<td>6</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Rokan</td>
<td>621</td>
<td>28</td>
<td>6</td>
<td>6</td>
<td>2</td>
<td>0.437 ± 0.113</td>
<td>0.00100 ± 0.00032</td>
</tr>
</tbody>
</table>

i Source: Nei, 1987 for haplotype and nucleotide diversity.

AP011385.1 and KC631202.1, the DNA sequence of Kelabau was identified as belonging to O. melanopleurus with 96%–97% accuracy, query coverage of 99%–100% and an E-value of 0.0. While based on the BOLD System, the identity of all samples was 96.60%–96.93% accurate.

In the phylogenetic tree consisted of two major groups (Figure 4). The first group was Cirrhinus moltonela (Gen-Bank KC631192.1) and it was divided from O. melanopleurus. The second group was differentiated into two sub groups, O. melanopleurus from GenBank (AP011385.1 and C631202.1) and 86 fish samples from Kampar, Siak and Rokan rivers. The 86 samples have BLASTN similarity values with O. melanopleurus of 96%–97%.

Discussion

Overall, the morphological traits and DNA barcoding showed that the majority of, if not all, kelabau fish in the three rivers at Riau Province were O. melanopleurus. there were 2 types of kelabau fish present in Riau. The first type is relatively small (15 gr to 2,778.93 gr) fish with dark colored dorsal and whitish ventral. This type of fish is used in this study. The second type of kelabau fish is larger, with yellowish color. This fish was not found during the study.

Environmental changes can cause fish death orbmigration to suitable habitats. Overfishing using both legal and illegal methods has also triggered the decline of certain species. In addition, our results suggested that there was little nucleotide diversity among O. melanopleurus in the Siak, Rokan and Kampar rivers in Riau Province, particularly the fish in the Kampar river.

The lack nucleotide diversity of O. melanopleurus from the three rivers was likely to be caused by limited opportunities for kelabau migration, so that the genetic exchanges with other...
populations are very small\(^5\); moreover, the lack nucleotide diversity is believed to be caused by inbreeding, and overfishing\(^3,7\).

In addition, Kelabau from the Siak and Rokan rivers were designated as one sub-sub group in group two (Figure 4) because both rivers are geographically connected, allowing for hybridisation, whereas, there is inbreeding of these fish in the Kampar River, which causes a nucleotide diversity value of 0.0. Geographically, these three rivers (Siak, Kampar and Rokan rivers) have different environmental condition, that cause variations in morphology in fish (O. melanopleurus) based on

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**Figure 4.** Phylogenetic tree of kelabau (Osteochilus melanopleurus) based on the neighbour-joining model.
Truss morphometric\textsuperscript{9}. Factors causing morphological variations are environmental differences, habitat pollution levels, long-term isolation and crossbreeding in populations\textsuperscript{39,40}. The same deductions were drawn from previous studies on Desmopuntius pentazona and D. rombochelatus, although the taxonomies of the two fish are different. Nevertheless, based on a genetic difference of only 0.4%, the two species were grouped into one cluster\textsuperscript{1}. These are distributed throughout Asia, the Mekong and Chao Praya river basins, Peninsular Malaysia, Sumatra and Borneo\textsuperscript{12}.

The identity of a species was derived using the morphological identification method to distinguish between species or organisms with a different ancestry\textsuperscript{41,42}. Basically, the genetic identification of a species can be done using mtDNA COI\textsuperscript{13}, a more effective approach than using rRNA\textsuperscript{35}. The nucleotide location and mutations were used as references to conduct DNA barcoding in all fish samples\textsuperscript{14}. Previous studies have identified several species using DNA bar-coding, such as ornamental fish of wetlands\textsuperscript{15}, wetland fish larvae\textsuperscript{16}, rainbow fish\textsuperscript{17}, Cyprinidae fish\textsuperscript{18}, salmon and trout\textsuperscript{19} and freshwater fish\textsuperscript{20,21}. Furthermore, the phylogenetics of COI sequences can effectively show congeneric and confamilial species\textsuperscript{22}.

The phylogenetic trees could describe the line of biological evolution from species or organisms with a different ancestry\textsuperscript{23}. Nonetheless, the results of all these species did not show a 100% undistinguishable identity. The branch length between species leading to a gap in the pairwise distance distribution is referred to as the barcoding gap in COI\textsuperscript{24}. Intra-species relationships were quite high in all samples, which confirmed that kelabau melanopleurus were native in the three rivers and could adapt to changes in environmental conditions\textsuperscript{25}.

Moreover, the existence of inter-nucleotide patterns and distances between A, T, C and G in the chromosomes showed the characteristics and genetic signs that distinguished each of the individuals, even though they belong to the same species\textsuperscript{26}. This is reinforced by referencing the phylogenetic tree made using the neighbor-joining model.

The identification of fish species is normally conducted using morphological characteristics such as dorsal fins, pelvic fins, pectoral fins, anal fins, lineal lateral fins, upper lineal lateral fins, lower lineal lateral fins, around body fins, and caudal peduncle fins; however, in this study, we used 12 morphological traits as described by the classification system of Kottelat et al.\textsuperscript{1}. These results supported the classification using biometric data that all fish in the three rivers were O. melanopleurus. The morphological characteristics were consistent with the species having a relatively large body with a standard length of 119–560 mm, lips covered with folds and plicae, no tubercles on the snout, a pair of maxillary barbels, and a pair of lower jaw barbels. The body is brownish, with the bottom brighter than the top. Dark hazy blotches near above of the pectoral fins, which is a special trait of O. melanopleurus.

However, this method can be difficult, and molecular identification is necessary. In particular, using mtDNA COI was an effective approach\textsuperscript{43}. The results from nucleotide distance data based on the Kimura 2-parameter model indicated that the nucleotide distance among the fish was short in intraspecific species using mtDNA COI\textsuperscript{27}, which was supported by data showing that the percentage identity in O. melanopleurus species ranged between 96% and 97%. The Kelabau fish from the three sample sites were identified as O. melanopleurus by percentage identity, supported by an E-value of 0.0 and a 99%–100% query cover. The p-value indicated that the BLASTN results contained no errors. Besides, the low nucleotide distance values (<3%–5%) among the samples of O. melanopleurus from the Siak, Kampar and Rokan rivers, indicated that all samples were monophyletic.

**Conclusion**

Based on our findings, we concluded that 86 of the 90 samples of kelabau from the Siak, Kampar and Rokan rivers in Riau were O. melanopleurus, as revealed by their morphological traits and the molecular analyses.

**Data availability**

**Underlying data**

COI gene sequences and raw biometric data of Osteochilus melanopleurus from Riau rivers can be found on OSF.

COI gene sequence DOI: https://doi.org/10.17605/OSF.IO/XGEZD\textsuperscript{37}

Raw biometric data DOI: https://doi.org/10.17605/OSF.IO/CFGM8\textsuperscript{38}.

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

The raw COI sequence data were also deposited in GenBank and can be found under sequential accession numbers MH430827-MH430854 and MH459085-MH459142.

**Acknowledgements**

The authors thank Neli Safrina as a staff Laboratory of Fisheries and Marine Faculty, Riau University and Gema Wahyu Dewantoro, as a staff researcher at the Laboratory of Ichthyology in the Field of Zoology, Research Center for Biology of the Indonesian Institute of Sciences (LIPI) on the identification of the Kelabau fish using morphological traits. Also, thanks to Ornamental Fish Aquaculture Research and Development Center, Ministry of Marine and Fisheries Republic of Indonesia, Depok for providing laboratory facilities.

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\textsuperscript{1}Department of Biology, Faculty of Science, Universitas Indonesia, Depok for providing laboratory facilities.

\textsuperscript{37}Center, Ministry of Marine and Fisheries Republic of Indonesia, Ornamental Fish Aquaculture Research and Development Center.

\textsuperscript{38}Dewantoro, as a staff researcher at the Laboratory of Ichthyology in the Field of Zoology, Research Center for Biology of the Indonesian Institute of Sciences (LIPI).
Open Peer Review

Current Peer Review Status: ✔️  ❓  ✔️

Version 2

Reviewer Report 10 February 2020

https://doi.org/10.5256/f1000research.24129.r59013

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Indra Junaidi Zakaria
Departement of Biology, Faculty of Math and Science, Andalas University, Padang, Indonesia

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Fish Biology and Fisheries biology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 08 Feb 2020

Nur Asiah, Padjadjaran University, Indonesia, Sumedang, Indonesia

Dear Mr. Indra Junaidi Zakaria,

Thank you for your suggestions on our manuscript. Through this letter I would like to explain the revision of my article titled “Biometric and genetic differences in kalabau (Osteochillus spp) as revealed using Cytochrome C Oxidase subunit 1”. I have revised these following points:

- Grammatical and spelling mistakes throughout the paper have been revised.
- The introduction has been improved by adding information on the difference of the characteristics of two species of kelabau fish (Osteochillus spp). As there are 2 types of fish that are named as “kelabau fish” by local people, information on the difference of those fishes was obtained by interviewing local fishermen. The presence of this information may be useful to give more explanation about the difference and the similarity of kelabau fish present in Riau in general.
- A phrase “The demand for it as …” until “… the population of these as well as many other fish” has been deleted as it is not correlate with the theme of this manuscript.
- Finally, the author has also added a quote in the following paragraph: “The study population was collected and sampled …” until “… of the Faculty of Fisheries and Marine Science, Riau University, Indonesia”.

Page 10 of 16
The discussion section has been improved by comparing the result of this study with more articles available.

Thank you for your suggestions and we hope that our article will be approved.

Best Regards,

Nur Asiah and the authors

**Competing Interests:** No competing interests were disclosed.

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**Reviewer Report 07 February 2020**

https://doi.org/10.5256/f1000research.24129.r59015

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Pallipuram Jayasankar
Fish Genetics and Biotechnology Division, ICAR-Central Institute of Freshwater Aquaculture, Bhubaneswar, Odisha, India

I have gone through the revision/corrections effected by the authors in the manuscript “Biometric and genetic differences in kelabau (*Osteochilus* spp.) as revealed using cytochrome c oxidase subunit 1”. The manuscript has been improved as per my suggestions. I recommend to consider its indexing.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Author Response 08 Feb 2020**

**Nur Asiah**, Padjadjaran University, Indonesia, Sumedang, Indonesia

Dear Mr. Pallipuram Jayasankar,

Thank you very much for your approved on our article, and thank you for your suggestions for the good of our article.

Best regards,

Nur Asiah and the authors

**Competing Interests:** No competing interests were disclosed.
Ja’afar Nuhu Ja’afar
1 Department of Biotechnology, Modibbo Adama University of Technology (MAUTECH), Yola, Nigeria
2 Chevron Biotechnology Center, Modibbo Adama University of Technology (MAUTECH), Yola, Nigeria

In version 1 of the manuscript, I have highlighted some sections that may require the authors' attention. However, none of those have been effected. Perhaps, the authors did not get the script I have corrected. It is available if needed.

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Molecular diagnostics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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Pallipuram Jayasankar
Fish Genetics and Biotechnology Division, ICAR-Central Institute of Freshwater Aquaculture, Bhubaneswar, Odisha, India

DNA barcoding is an important and useful tool to ratify species status in circumstances where morphology-based taxonomy fails or nearly-fails to perform unambiguous identification of species. The authors in the present study have taken up an important group of freshwater fish. There are many grammatical errors and some technical flaws in the manuscript. I felt one major drawback could be their total silence on the nominal species of *Osteochilus kelabu*. Of course using barcoding they have proved that the species is *O. melanopleurus*. However, they should have shown a phylogenetic comparison of these two nominal species to rule out confusion in identification. Morphological method followed does not appear to be robust. They are advised to use TRUSS MORPHOMETRICS which can be performed using
user-friendly software nowadays. In the phylogenetic tree the majority of bootstrap values do not appear to be robust; hence it could reflect a spurious relationship.

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
No

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: I have been working on molecular taxonomy of fish, shellfish and marine mammals since 1996, thus 23 years of experience in this field; also gained expertise in freshwater aquaculture technology during the past 11 years.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Reviewer Report 04 October 2019
https://doi.org/10.5256/f1000research.18938.r53842

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Generally, the article needs to be proofread by an expert. There are a lot of areas I have highlighted to guide the authors.

In paragraph three of the discussion section, the authors should be more scientific in providing reasons...
for limited variation such as inbreeding within the species due to limited migration. More so, if COI is known to be conserved within a species, it is not expected to change within a small time frame of genetic event. In general, the discussion should be rewritten to explain the results and interpretations of the study.

In addition, I suggest the authors contact an expert that can explain/interpret the results obtained using DNASP. The explanation of the results from the software should be more than presenting the values.

The gel result will be more appreciable if the marker has been properly labelled.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Partly

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Molecular diagnostics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Reviewer Report 04 March 2019

https://doi.org/10.5256/f1000research.18938.r44418

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**Indra Junaidi Zakaria**

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**Generally:**
- There are still a lot of grammatical and spelling mistakes throughout the paper. Therefore, the authors should remove all of the grammatical and typographical errors as well as spelling mistakes. Use past tense and rephrase the sentences.

Introduction:
- Inconsistency in writing in the first paragraph, the first line: it is written as “genus” but in the second row it is written as “species”. Kelabau are ancient fish belonging to the genus Osteochilus of family Cyprinidae. The species are distributed throughout Thailand, Borneo and Sumatra (etc.).
- According to local fishers in Riau, kelabau are divided into two types on the basis of morphology. Although there is no detailed information about these fish types, please, the authors should explain what the differences in the characteristics are of the two species which the fishermen know.
- Sentences from: “The demand for it as…” until “…the population of these as well as many other fish.” I suggest deleting it, because it has no correlation with the theme of this manuscript which explains the verification of the species of fish kelabau using cytochrome c oxidase - it is not about increasing demand and the problem of decreasing populations of kelabau fish.

Methods:
- Cite references in the following paragraph: “The study population was collected and sampled…” until “…of the Faculty of Fisheries and Marine, Riau University, Indonesia.”

Results:
- Ok.

Discussion:
- The Discussion is not up to the mark, especially in paragraphs one and two; please rewrite back with an explanation of how the characteristics of the two species of fish differ according to fishermen and the reason that these differences were caused by environmental changes and overfishing.

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes
**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Fish Biology and Fisheries biology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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