Short Communication:
Identification of marine leech and assessment of its prevalence and intensity on cultured hybrid groupers (Epinephelus sp.)

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Abstract. Murwantoko, Negoro SLC, Isnansetyo A, Zafran. 2018. Short Communication: Identification of marine leech and determination of its prevalence and intensity on cultured hybrid groupers (Epinephelus sp.). Biodiversitas 19: 1798-1804. Groper is an important fish species due to its high price both in domestic and international markets. Several hybrid groupers have been produced and can be accepted in market. A major production constraint in grouper culture is mortality due to diseases. Leech is an ectoparasite for groupers which may cause significant loss. The aims of this study were to identify and to assess the prevalence and intensity of leech on hybrid grouper cultured in sea cages at Buleleng waters. Morphological identification was conducted using fresh and stained specimens while molecular identification was conducted using nucleotides sequence of mitochondrial cytochrome oxidase subunit I (COI). The presence of leech was observed by unaided observation of 14 populations of hybrid grouper. Morphological identification showed that the leech belonged to Zeylanicobdella arugamensis. This result was also supported by analysis of COI sequence that showed 100% homology with Z. arugamensis (accession number KY 441721.1) and 90% homology with Aestabdella abditovesiculata (accession number DQ414300.1). Hybrid groupers at sea cages were infected by leeches with prevalence and intensity, respectively, of 100% and 21.2 leeches fish-1. The prevalence and intensity were varied depending on the farm and population. Cantik grouper was more susceptible to leech infection than cantang grouper. The bigger fish tended to have higher leech prevalence and intensity.

Keywords: Cytochrome oxidase, hybrid grouper, identification, leech, Zeylanicobdella arugamensis

INTRODUCTION

Several species of grouper have been cultured in Indonesia and become important fish commodities due to its high price both in domestic and international markets. Several types of hybrid grouper have been developed to increase the quality of fish. Cantang grouper is produced as a result of crossbreed between female tiger grouper (Epinephelus fascioguttatus) and male giant grouper (Epinephelus lanceolatus). The crossbreed between the female tiger grouper and the male brown-marbled grouper (Epinephelus microdon) was named cantik grouper. A crossbreed between mouse grouper (Cromileptes altivelis) and giant grouper was named kustang grouper (Ismi et al. 2013). Cantang grouper culture has developed well from its high price in both domestic and international markets. Several hybrid groupers have been produced and can be accepted in market. A major production constraint in grouper culture is mortality due to diseases. Leech is an ectoparasite for groupers which may cause significant loss. The aims of this study were to identify and to assess the prevalence and intensity of leech on hybrid grouper cultured in sea cages at Buleleng waters. Morphological identification was conducted using fresh and stained specimens while molecular identification was conducted using nucleotides sequence of mitochondrial cytochrome oxidase subunit I (COI). The presence of leech was observed by unaided observation of 14 populations of hybrid grouper. Morphological identification showed that the leech belonged to Zeylanicobdella arugamensis. This result was also supported by analysis of COI sequence that showed 100% homology with Z. arugamensis (accession number KY 441721.1) and 90% homology with Aestabdella abditovesiculata (accession number DQ414300.1). Hybrid groupers at sea cages were infected by leeches with prevalence and intensity, respectively, of 100% and 21.2 leeches fish-1. The prevalence and intensity were varied depending on the farm and population. Cantik grouper was more susceptible to leech infection than cantang grouper. The bigger fish tended to have higher leech prevalence and intensity.

The hirudinea infection on grouper is one problem for parasitic diseases. Hirudinea has four orders, namely Acanthobdellia, Gnathobdellia, Pharyngobdellida, and Rhyynchobdellida. The Rhyynchobdellida Order has three families, i.e., Glossiphonidae, Ozobranchiidae, and Piscicolidae. The Piscicolidae family is characterized by having a symmetrical, flattened cylinder body, an anterior suction and a posterior suction. Their habitats are freshwater and seawater, swimming by extending their body (Sawyer 1986). Species of leeches the family Piscicolidae are often parasitic seawater fish such as Pterobdella amara, Aestabdella leiostomi, Piscicola spp., and Zeylanicobdella arugamensis (Chandra 1991).

Marine leeches are an essential threat to the aquaculture industry (Ravi and Yahaya 2017). Heavily infested fish with leeches often have chronic anemia (Noga 2000). Grouper having infected leeches on its skin will rub the body on objects around it causing injuries and a large ulcer on the skin or in the mouth. Those conditions can cause secondary infection (Noga 2000; Johny and Roza 2006). Fishes mortality usually occurs within a 3-day period following infestation due to secondary infections with pathogenic bacteria such as Vibrio alginolyticus (Ravi and Yahaya 2017). Leeches infection also often transmits microbes and hemoparasites during feeding (Noga 2000). Marine leeches Z. arugamensis have been reported to have the ability to transmit the hemogregarine and trypanosomes
simultaneously between fish (Hayes et al. 2006).

Grouper culture using floating net cages in Pegametan Buleleng waters has been started in 2003. The number of sea cages in these waters is increasing due to the potential and reasonable price of grouper fish and high export demand. An outbreak of leeches was reported in grouper farm on August 2016. In this study, we identified the leech based on morphological and molecular approaches and determined the prevalence and intensity of leeches on hybrid groupers.

MATERIALS AND METHODS

Leeches sampling

Seven farms were selected to represent grouper culture in Pegametan Bay, Buleleng waters in September-October 2016. The position of farms were: Farm A at 8°07'10.7"S 114°36'47.1"E, Farm B at 8°07'03.0"S 114°36'42.2"E, Farm C at 8°07'17.6"S 114°37'04.9"E, Farm D at 8°07'47.3"S 114°36'06.9"E, Farm E at 8°07'27.9"S 114°35'55.5"E, Farm F at 8°07'40.5"S 114°36'09.3"E and Farm G at 8°07'40.9"S 114°35'44.6"E. All fish populations on the selected farm were sampled for the study. We defined population as fishes in a cage which have the same species, the same age and the same source of hatchery when stocked into the cage.

Leech observation

Thirty-six fishes were randomly sampled from each population to meet detection with a minimum prevalence rate of 10% with a 95% confidence level. For one population, fishes were sampled from three cages with twelve fishes in each cage. Fishes were collected from cages using scope net, then kept in a bucket. The species, length, and weight of fishes were recorded. The presence of leeches was observed with unaided eyes from all surface body of fish. The number of parasites was counted, and infected organs were recorded. The prevalence was calculated as the proportion of infected fishes among all the fishes in population. The intensity was calculated as the number of leeches found in the infected fish. For morphological identification, the leeches were collected alive and kept in containers with seawater for further identification. For molecular identification, ten parasites were fixed in 5 ml tubes containing 70% ethanol for further analysis.

Morphological identification

Morphological identification was performed using five fresh samples and five aceticarmine stained samples. Parasites were stained basically from Roberts et al. (2012) with 1% aceticarmine, and then destained using 1% HCl in 70% ethanol. The observations were conducted under a microscope and documented. Identification of species based on morphology and anatomy followed the guidelines of Sawyer et al. (1982) and Chandra (1983).

Molecular identification

The genomic DNA was isolated based on TNES method (Murwantoko et al. 2008). Approximately 50 mg of leech was ground in up 400 μl TNES on the microtube and added with three μl of Proteinase K (Roche) and incubated for 2.5 hours at 37 °C. After incubation, the mixture was centrifuged at 13500 x g for 5 minutes with Sorval Legend Micro 17 Microcentrifuge (Thermo scientific). The supernatant was collected and extracted with 300 μl of PCIAA solution (Phenol Chloroform Isoamyl Alcohol). After centrifugation at 13500 x g for 1 min, the aqueous phase was collected and added with 30 μl 5 M NaCl (1/10 volume of supernatant), and 600 μl cold absolute ethanol (2x volume of supernatant) then incubated for 24 hours in the refrigerator. The mixture was centrifuged at 13500 x g for 5 minutes, the supernatant was discarded, and the pellet was washed with 500 μl of ethanol 70%. After drying, the pellet was resuspended in 100 μl of TE containing 0.5 μl of RNase.

The molecular identification was conducted based on mitochondrial cytochrome oxidase subunit I (COI) gene. LCO 1490 (GGT CAA ATA ATA AAG ATA TTG G) as the forward primer and HCO 2198 (TAA ACT TCA GGG TGA CCA AAA AAT CA) as the reverse primer (Lobo et al. 2013) were used. Amplification was performed in T100TM Thermalcycler (Biorad) with initial denaturation program at 95 °C for 3 minutes, followed by 30 cycles of denaturation at 95 °C for 30 seconds, annealing at 55 °C for 30 seconds, extension at 72 °C for 1 minute and the final extension at 72 °C for 5 minutes. PCR product was electrophoresed in agarose gel in TAE solution with Fluorosave DNA stain (1st Base) using Mupid_2Plus electrophoresis tank (Advance). After electrophoresis, the gel was observed under UVP Transilluminator (Pasificimage Electronic). The PCR products were then sequenced through the sequencing service company. Aligned sequences were also subjected to nucleotide BLAST (Basic Local Alignment Search Tool) search to know the identity. Cluster tree was constructed under unweighted pair group method with arithmetic mean (UPGMA) using MEGA 7 software (Komar et al. 2016).

RESULTS AND DISCUSSION

Grouper culture

The culture of grouper using floating net cages in Pegametan bay has been started since 2003. In 2016 there were 24 farms with approximately 4000 cages as recorded by Association of Coastal Fish Farmer of Buleleng. The number of cages in each farm varied between 40 to 500 cages. The size of each cage ranged from 2 m x 2 m, 3 m x 3 m and 3 m x 6 m. A cage of 3 m x 3 m size was stocked with 500-600 fishes of 11-15 cm length, and cage of 3 m x 6 m was stocked with 700-800 fishes of 11-15 cm length.

The most commonly cultured grouper commodities were cantik hybrid grouper (Epinephelus sp) and cantang hybrid grouper (Epinephelus sp). Tiger grouper (Epinephelus fuscoguttatus), mouse grouper (Cromileptes altivelis), orange spotted grouper (Plectropomus leopardus), malabar grouper (Epinephelus malabaricus), and brown-marbled grouper (Epinephelus microdon) were
cultured in limited number. Based on information from the farms, the leech started to infect groupers with low intensity on few cages in April 2016. In August 2016, when there was a high tide, the leech infection spread to many floating net cages in the waters. Therefore, the sampling conducted around September to October was in condition with relatively high leech infection.

Fish samples
The grouper samples were taken from seven different farmers with total sample of 14 populations. The samples were composed by 9 populations of cantik grouper and 5 populations of cantang grouper. Based on the size, samples can be categorized on small, medium and big with 7, 5 and 2 populations, respectively (Table 1).

Location of leech infection
Leeches were easily observed and founded in mouth, eyes, operculum, skin, dorsal fin, anal fin, pectoral fins and tail (Figure 1). This parasite attaches to its host using its sucker and takes its host blood causing the leeches to become diverse in color as black and brown.

Table 1. Grouper samples from Pegametan cages

| Farm | Pop. | Species | Length (cm) | Weight (g) | Category |
|------|------|---------|-------------|------------|----------|
| A    | A1   | Cantik  | 15.9 ±1.3   | 62.6 ± 14.1| Small    |
|      | A2   | Cantang | 20.2 + 1.7  | 235.7 + 27.4| Medium   |
| B    | B1   | Cantik  | 16.6 + 1.5  | 72.2 + 17.2| Small    |
| C    | C1   | Cantik  | 17.1 + 0.7  | 75.1 + 7.4 | Small    |
|      | C2   | Cantik  | 29.9 + 1.4  | 434.5 + 78.1| Big      |
| D    | D1   | Cantang | 21.2 + 1.9  | 268.0 + 38.4| Medium   |
|      | D2   | Cantik  | 14.1 + 0.9  | 46.2 + 9.7 | Small    |
| E    | E1   | Cantang | 14.4 + 9.6  | 50.9 + 10.3| Small    |
|      | E2   | Cantik  | 21.0 + 14.7 | 304.5 + 19.5| Medium   |
| F    | F1   | Cantik  | 32.2 + 3.0  | 461.6 + 103.8| Big      |
|      | F2   | Cantik  | 13.9 + 1.2  | 43.6 + 9.7 | Small    |
| G    | G1   | Cantang | 14.5 + 1.1  | 51.1 + 10.5| Small    |
|      | G2   | Cantik  | 21.3 + 1.1  | 252.7 + 20.4| Medium   |
|      | G3   | Cantang | 20.4 + 1.5  | 258.7 + 33.5| Medium   |

Figure 1. Infection by leech on fish body part (A: Infected grouper, B: Infection on operculum, C: Infection on eyes, D: Infection on the body surface, E & F: Infection on dorsal fin, G: Infection on caudal fin, H: Infection on pectoral fins and I: Infection on anal fin)
Prevalence and intensity

Leeches were found from all observed farms with different prevalence and intensity (Figure 2A). The average prevalence among farms was 62% with the highest prevalence was 86% (Farm C), and the lowest prevalence was 11% (Farm F). The average intensity among farms was 7.1 leeches fish$^{-1}$ with the highest intensity was 11.2 leeches fish$^{-1}$ (Farm F), and the lowest intensity was 0.9 leeches fish$^{-1}$ (Farm D).

The prevalence and intensity levels of leech infection on each grouper sample population were varied (Figure 2B). Prevalence in the populations was also different even on the same farm. The highest prevalence was in population C2 and F1 grouper (100%), and the lowest was in population D2 (0%). The high prevalence variation among the population in farms occurred in Farm G (population G2 of 72.2% and population G1 of 8.3%) Farm F (Population F1 of 100%, population F2 of 19.4%). The intensity of leech infection in the population was also different even on the same farm. The highest inter-population variation in farms occurred in Farm F with population F1 of 21.2 leeches fish$^{-1}$ and population F2 of 0.3 leeches fish$^{-1}$.

The prevalence and intensity of leech infection on cantik grouper were, respectively, 69% and 9.3 leeches fish$^{-1}$, which were higher than those of cantang grouper with only 42% in prevalence and 2.6 leeches fish$^{-1}$ in intensity (Figure 2C). This result suggests that cantik grouper is more susceptible to leech than cantang grouper. The highest prevalence of 100% was found in large grouper, and then 62% in medium grouper group and the lowest was 46% in small grouper. The highest intensity also showed similar pattern with the highest intensity was found in large grouper and the smallest was found in small grouper (Figure 2D).

Morphological identification

The leeches can be observed on the fish body using unaided observation. The parasite has cylindrical shape, soft, elastic, and smooth body surface with light brown or black (Figure 3.A, 3.B). This parasite attaches to fish using its sucker and sucks its host blood. Adult of this species has a length of about 8-18 mm and a maximum width of the urosome of 0.5-2.0 mm. Anterior sucker has a diameter of 0.3-0.5 mm, and posterior sucker has a larger diameter that of 1.0-1.8 mm.

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**Figure 2.** Prevalence and intensity of leech on the farm (A), population (B) species (C) and grouper size (D)
Figure 3. Leech found in grouper (A; Leech with black color; B: Leech with brown color)

Figure 4. Morphology of leech (1 = Anterior sucker; 2 = Subesophageal ganglion mass; 3 = Proboscis; 4 = Ductus ejaculator; 5 = Ovary; 6 = First testicular ganglion; 7 = First Testis; 8 = Crop Abdomen; 9 = Pulsatile vesicle; 10 = 5th Testis; 11 = Posterior crop caecum; 12 = Posterior sucker)

Table 2. Leech determination according to Chandra (1983)

| No | Description                                      |
|----|--------------------------------------------------|
| 1b | Has eyes and pulsating vesicles                  |
| 4a | Eye pair                                         |
| 5a | Has no lateral branchiae                         |
| 6b | Has pulsating vesicles                          |
| ⊲  | *Zeylanicobdella arugamensis*                    |

Table 3. Leech determination according to Sawyer et al. (1982)

| No  | Description                                      |
|-----|--------------------------------------------------|
| 1b  | Species that live in seawater or brackish         |
| 8b  | Has no gill radius                               |
| 9a  | Has 10-12 pairs of pulsating vesicles along the lateral border of the abdomen |
| 10b | Small size of about 1-2 cm                       |
| 11b | Posterior sucker has a different sleeve with an anterior sucker |
| 12a | The lower body is smooth, about 12 segments in the body |
| ⊲  | *Zeylanicobdella arugamensis*                    |

This species has a pair of eyespots on the anterior sucker, 12 segments in the body, five pairs of testes and a pair of ovaries. The other part of the body of this species consists of subesophageal ganglion mass, proboscis, ovary, crop posterior caecum, pulsatile vesicles, testis (1-5), crop abdomen, testicular ganglion (1-5), and the ductus ejaculator (Figure 2). Posterior sucker (Figure 4(12)) different sleeve and large size than anterior sucker (Figure 4(1)). Based on determination key of Chandra (1983) and Sawyer et al. (1982), this species is *Zeylanicobdella arugamensis* (Table 2 & Table 3)

Molecular identification

The genomic DNA was used as the template to amplify COI gene. The COI gene from leech was successfully amplified as indicated by the presence of a single band of DNA after agarose electrophoresis. This DNA fragment contained 725 nucleotides sequences that have been deposited in Genbank with accession number MH299847. The BLAST analysis showed that the sequence has a 100% identity with *Zeylanicobdella arugamensis* (KY4741721.1), while its homology with *Aestabdella abditovesiculata* (DQ414300.1), *Pterobdella amara* (DQ414334.1), *Myzobdella lugubris* (KY440059.1) was, respectively, 90%, 89%, and 86%. This Indonesian *Z. arugamensis* is closely related with *Z. arugamensis* from Malaysia (Figure 5).

Discussion

The leeches can be seen visually attached to the fins, tail, body, operculum, mouth, and eyes of fishes. Some fishes showed hemorrhagic on their body surfaces, which is in line with the statements of Johnny and Roza (2006) that leech infection was found in the external part of the fish and caused hemorrhages leading to secondary bacterial infection. According to Ravi and Yahaya (2017), the most frequent effect of leech infection in fish are local bleeding and ulceration in fish tissues. This species is attached to the host using anterior and posterior suckers. They suck the blood of their host using their sucker, and in this study, the leech having fish blood was black (Figure 1). Leech that has sucked fish blood will escape from fish to find a place for spawning (Kua et al. 2010). After detached from the host, leeches were able to swim in the sea and able to survive without host for 5-7 days Cruz-Lacierda et al. (2000).

Leech parasites were found from all observed farms in Pegametan bay. The prevalence and intensity of leech that infected grouper in each farm were varied in values. The prevalence and intensity were also varied between population among farms and within farm. The highest prevalence was found in population C2 and F1 (100%), and the lowest prevalence was in population D2 (0%). The results indicated that infestation of leech was affected by location and or cultivation management. The prevalence and intensity of leech varied between species and size. Cantik grouper was more susceptible than cantang grouper. This shows that cantik grouper has a higher risk of disease infection than the cantang grouper. The bigger fish tended to have higher prevalence and intensity.
The average prevalence for these 14 populations was 59%, and average intensity was 6.9 leeches fish\(^{-1}\). This prevalence and intensity were higher than that reported in muddy grouper in the Philippines with a prevalence of 30% and intensity of 2 leeches fish\(^{-1}\) \cite{Cruz-Lacierda2000}, and in red snapper with study of ectoparasite prevalence was 11.5% and intensity of 1.48 leeches fish\(^{-1}\) \cite{Ravi2017}. This prevalence is lower than the that in white snapper in Malaysia (70%) \cite{Kua2006}.

Morphological and molecular identification based on COI sequence consistently showed that the leech belonged to *Zeylanicobdella arugamensis*. The COI sequences of *Z. arugamensis* on Genbank is limited, where up to April 2018, only ten entries were available. *Z. arugamensis* showed the genetic diversity, and at least 3 clusters are shown in Figure 5, Indonesian-Malaysian, Malaysian-China and Iran clusters. This genetic diversity seems to be correlated with the country location. *Z. arugamensis* had been reported to infect brackish-water fish Mozambique tilapia, *Oreochromis mossambicus* in Okinawa Japan \cite{Nagasawa2000}, amphibious goby (*Scartelaos tenuis*) in southern Iran \cite{Polgar2009}. *Z. arugamensis* had been reported to infect cultured marine fish such as orange-spotted grouper (*Epinephelus coioides*) in Philippine \cite{Cruz-Lucierda2000}, crimson snapper (*Lutjanus erythropterus*) in Malaysia \cite{Ravi2017}, orange-spotted grouper, (*Epinephelus coioides*) in Indonesia \cite{Kleinertz2015}. In this study, we report the first time that *Z. arugamensis* can infect cantang and cantik hybrid groupers.

Several authors have documented study on leech infections on fish in Indonesia. Rosa and Johny (2006) have reported infection of the leech on *Epinephelus bleekeri* and *E. polyplakadion*. However, the species of leech was not reported. The intensity of *Piscicola* sp from tiger grouper (*E. fuscoguttatus*) and spotted coral grouper (*Plectropomus maculatus*) has been reported by Diana et al. (2004). The *Z. arugamensis* in Indonesia was reported by Kleinertz and Palm (2015) from orange-spotted grouper, *E. coioides* with the species identification was only based on the morphology using microscope observation. Here we report the first time the identification of *Z. arugamensis* in Indonesia based on the COI nucleotide sequences and the sequences have been deposited in Genbank with accession number MH299847. Improvement of culture management should be addressed to control the leech. The lesions form this infection can cause secondary infection by bacteria, and this *Z. arugamensis* has been reported to be able to transmit the hemogregarine and trypanosomes simultaneously between fish \cite{Hayes2006}.

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