Gonadal Degeneration and Hepatopancreas Alteration in Orange Mud Crab *Scylla olivacea* Infected With *Sacculina beauforti* (Crustacea; Rhizocephala; Sacculinidae)

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Infestation of *Sacculina beauforti* on an aquaculture species, mud crab *Scylla olivacea* is alarming due to its high prevalence and the extreme morphological changes in hosts. To further understand its pathological effect on growth and reproduction of *S. olivacea*, gonadal and hepatopancreatic histological changes of infected individuals were compared with healthy individuals. Also, the histological characteristics of *S. beauforti*’s mature externa was described. Hepatopancreases of infected individuals were loosely packed and rootlets were observed in the intertubular spaces. Although hepatopancreatic tubule count was significantly lower, tubule diameters were unaffected. Gonads, however, were severely affected. No germ cells were found in the infected testes (except for remnants of spermatozoa), indicating the arrest of spermatogenesis. Rootlets were also present in testes of infected individuals. Ovarian tissues of infected females were severely damaged with no rigid cell structures. Gonadosomatic index (GSI) of infected males and females were lower, but the hepatosomatic index (HSI) were higher than their healthy counterparts. No significant differences in GSI and HSI values were observed among infected males and females. Multiple regression analysis revealed that carapace width, GSI and HSI are statistically important for the prediction of infection status. Given the high prevalence of *S. beauforti* infection and its disruptive effects on the morphology and internal physiology of *S. olivacea*, this study, albeit fundamental and incomprehensive, highlights to farmers and researchers the emergence of a castrating parasite and the urgency for the development of preventive measures and treatments for this disease in an economically important aquaculture species.

**Keywords:** sacculinid, rhizocephalan, mud crab, *Scylla olivacea*, histology, externa
INTRODUCTION

Orange mud crab *Scylla olivacea* is widely distributed around Southeast Asia (SEA) countries, Australia, Japan and Pakistan (Keenan et al., 1998; Alberts-Hubatsch et al., 2016). Together with other *Scylla* species, i.e., *S. paramamosain*, *S. serrata* and *S. tranquebarica*, mud crabs are economically important, not only as a valuable commodity to support the livelihood of coastal communities but also as a major aquaculture crustacean species that contribute greatly to a country’s economic growth. According to FAO (2019), the global aquaculture production of mud crabs (*Scylla* spp.) was doubled within 3 years, from 41,460 tons in 2013 to 89,390 tons in 2016. The aquaculture production of mud crabs, however, still relies almost entirely on wild-caught seeds and broodstocks (Waiho et al., 2018). In most countries including Malaysia where wild population of mud crabs are still abundant, most aquaculture activities involve fattening and soft-shell crab farming using juveniles. They are often held in earthen ponds that rely on spring tide for water exchange or directly cultured in pens and cages in their natural environment (mangroves, lagoons or estuarine areas) (Ihwan et al., 2013). Due to the increasing domestic and international demand for mud crabs, hatchery and nursery productions are also gaining attention. Private farmers, industrial players and research institutions are working toward optimization of mud crab larval rearing, although broodstocks are still sourced from the wild (Waiho et al., 2018).

Thus far, parasites infecting mud crabs, especially *S. olivacea*, in Malaysian waters include cuculliaceans, nematodes, copepods and barnacles (*Octolasmis* spp.). Being the most abundant among other parasites (71.1% prevalence) (Ihwan et al., 2015), *Octolasmis* barnacles are often found in the respiratory chambers of mud crabs (Jeffries et al., 1989) and pose minimal threat to their hosts as they do not feed on host tissues (Gannon and Wheatly, 1992). Recently, however, the occurrence of rhizocephalan barnacles *Sacculina beauforti* (*Sacculinidae*) on mud crab *Scylla olivacea* was reported, with a high prevalence rate of 42.27% (Waiho et al., 2017b). Unlike barnacles of other superorders, i.e., Thoracica (filter-feeding barnacles) and Acrothoracica (burrowing barnacles), rhizocephalan barnacles are parasitic barnacles that exhibit extremely reduced adult forms (absence of any organ structures, centralized nervous system, feeding appendages and segmentation) (Glenner and Hebsgaard, 2006) and cause severe behavioral changes, and morphological and physiological damages to their hosts (Hoeg, 1995; Yang et al., 2014). To date, this rhizocephalan species has only been reported in two *Scylla* species, i.e., *S. olivacea* and *S. serrata* (not found in Malaysian waters) (Fazhan et al., 2017). Although the other two species (*S. tranquebarica* and *S. paramamosain*) live sympatrically with *S. olivacea*, *S. beauforti* seems to be selective of its host species as no infestation has been found (Waiho et al., 2017b).

All rhizocephalans infect other crustaceans (Hoeg, 1995; Hoeg and Lützen, 1995). Their unique life cycle begins with the hatching of sexually dimorphic nauplii, which upon several molts, metamorphose into male and female cyprids (Trédez et al., 2017). Female cyprids would develop into kenetrogon larvae and infect hosts by penetrating hosts’ integument, thereby transferring parasitic material (vermigon) into the hemolymph (Glenner and Hoeg, 1994; Glenner, 2001). Eventually, an interna within the host is formed. It possesses numerous rootlets that infiltrate host’s organs for nutrient absorption (Nagler et al., 2017). During this period of infection, the external morphology of the host remains normal. After several molts, the host starts to exhibit female-like characters and molting usually stops when an externa emerges on the soft ventral surface of the host’s abdomen (Kristensen et al., 2012). Unlike their female counterparts, male cyprids are free-living. Mature externae will start to release nauplii not long after dwarf males have been implemented into a pair of receptacle-like structures formed in the visceral mass of each externa (Hoeg, 1995; Glenner et al., 2000).

The adverse effects of *S. beauforti* infection on the external morphology of *S. olivacea* are obvious. Among them includes feminization of male hosts (broadened and darkened female-like abdomens), reduction in copulatory appendages (gonopods in males and pleopods in females) and significantly smaller body size (Waiho et al., 2017b). These changes profoundly decrease their market values as they are often regarded as immature females and traded off at a much lower market price (personal communication). However, due to the limited reports on their occurrence (Boschma, 1949; Waiho et al., 2017b; Fazhan et al., 2018), the precise biology and life cycle of *S. beauforti*, and the effects they have on their hosts remain largely unknown. As the growth and reproduction of crustaceans are primarily regulated by hepatopancreas and gonads (Wang et al., 2014), we investigated the pathological changes of the *S. beauforti* infection had on these two organs of *S. olivacea*. With the current reliance of mud crab aquaculture sector on wild stocks, the results of this study, though fundamental, serve as essential baseline data for future prevention, management and treatment of this disease.

MATERIALS AND METHODS

Orange mud crabs, *Scylla olivacea* were collected from local fishermen in Marudu Bay, Sabah, Malaysia. Healthy mature adults (carapace width, CW range = 90–105 mm, n = 30 for each sex) (Waiho et al., 2016; Fazhan et al., 2018) and sacculinid-infected crabs with mature externa (CW range = 90–105 mm, n = 21 for females, n = 20 for males) were obtained, with the former healthy individuals serving as controls. Species identity was validated based on the taxonomic key provided by Keenan et al. (1998). The sex of infected crabs was determined based on the presence of gonopods (males) or pleopods (females) (Waiho et al., 2017b; Fazhan et al., 2018). Crabs were classified as “healthy” based on the absence of (1) externa, (2) visible morphological alterations on sexual organs or abdomen, and (3) interna upon dissection (Waiho et al., 2017b). All animal experimental procedures were reviewed approved by the Institute of Tropical Aquaculture and Fisheries, Universiti Malaysia Terengganu.

Measured parameters were CW (to the nearest 0.01 mm), body weight (BW, to the nearest 0.1 g), and the total weight of gonad (whole ovary for females; testis and vas deferens for males)
and hepatopancreas (to the nearest 0.001 g). Gonadosomatic index (GSI) and hepatosomatic index (HSI) were calculated based on the following equations:

\[ GSI = \frac{\text{total weight of gonad}}{\text{BW}} \times 100 \]

\[ HSI = \frac{\text{total weight of hepatopancreas}}{\text{BW}} \times 100 \]

Gonadal tissues of 20 individuals (healthy males and females, and infected males and females, with five specimens each) and hepatopancreas tissues of 10 individuals (five healthy individuals and five infected individuals) were extracted and fixed in Davidson’s solution for 24 h (Waiho et al., 2017c). In addition, a mature externa (slightly cut open using scalpel) and its stalk base were similarly fixed for 48 h to allow full penetration of fixative. Fixed tissues were then transferred to 70% ethanol for 24 h before being dehydrated in ascending (70–100%) ethanol concentrations. The dehydrated tissues were subsequently cleared in xylol, embedded in paraffin wax and sectioned to 5 μm thickness using a rotary microtome (Leica RM2255). Standard Mayer’s Hematoxylin-Eosin (HE) staining was applied to all sectioned slides. After staining, sections were mounted on slides using DPX and viewed using Nikon Eclipse 80i Advance Research Microscope. Hepatopancreatic tubules on one transverse cross-section were counted \((n = 10 \text{ sections})\) and randomly measured \((n = 50)\). Further, the diameter of \(S. \text{beauforti}\)’s rootlets were also measured \((n = 30)\). All counts and measurements were conducted under 4× magnification.

All data analyses were performed using Microsoft Excel 365 and IBM SPSS Statistic ver. 25. Data were checked for homogeneity of variance using Levene’s test and Welch’s correction was applied during one-way analysis of variance (ANOVA) if homogeneity of variance is violated (Welch, 1947). Subsequent differences among treatments were detected using Games Howell posthoc test (Kirk, 1995). Multiple regression was conducted, with infection status as dependent variable and sex, CW, BW, GSI and HSI as independent variables to determine if these variables are significant in predicting infection status in \(S. \text{olivacea}\). All data were expressed in the form of mean ± standard deviation (s.d.) and a significance level at \(\alpha = 0.05\) was used in all statistical analyses in this study.

**RESULTS**

The body of the externa is laterally compressed. A thin-walled, muscular mantle surrounds a visceral mass, which contains the connective tissue and the reproductive apparatus. A cuticle clad mantle cavity is located between the mantle cavity and the mantle, and it communicates with the surrounding seawater by a mantle opening (aperture). The aperture is situated at the anterior margin of the externa, opposite the stalk, which through the ventral, abdominal integument of the host connects the external part of the parasite, the externa, with the internal root-like system of the parasite, the interna (Figure 1A). A pair of strait receptacles are located in the muscular basal region of the stalk outside the visceral mass. In lateral cross-section of the basal stalk reveals that it is filled with tubes (Figure 1D). The ovaries in the visceral mass are filled with developing ovules and is traversed by muscle fibers (Figure 1B). The mantle of the externa consists of two layers of collagen fibers enveloping two layers of radial muscles with a longitudinal muscle layer situated in the middle (Figure 1C).

To determine the effect of \(S. \text{beauforti}\)’s infection on the host’s gonadal and hepatopancreatic physiology and development, the gonadal and hepatopancreatic histological sections of healthy and infected individuals were compared. Densely packed hepatopancreatic tubules with clear star-shaped luminae were observed in the hepatopancreas of healthy individuals (Figure 2A). A thin layer of connective tissues was found in the intertubular spaces. In the hepatopancreas of infected crabs, hepatopancreatic tubules were loosely packed and \(S. \text{beauforti}\)’s rootlets were found among tubules. The \(S. \text{beauforti}\)’s rootlets had a range of 40.27–125.68 μm in diameter \((\text{mean} = 80.00 \pm 22.74 \text{μm})\) (Figure 2B). The hepatopancreatic tubule count in healthy individuals \([\text{count range } (n) = 89–135; \text{mean} = 109.00 \pm 14.78]\) was statistically significantly higher than those infected with \(S. \text{beauforti}\) \([\text{count range } (n) = 20–44; \text{mean} = 33.60 \pm 7.46]\) (Mann-Whitney U; \(U = 0, P < 0.001\)). However, the hepatopancreatic tubule diameters were not significantly different between healthy and infected individuals \((\text{Student’s } t \text{ test}; F_{1,98} = 3.099, P = 0.081)\). Resorptive cells (R-cells) and blister-like cells (B-cells) were still present in the hepatopancreatic tubules of infected individuals.

All four types of germ cells, i.e., spermatogonia, spermatocytes, spermatids and spermozoa were present in a healthy testis (Figure 2C). However, severe degradation of testis was observed in infected individuals (Figure 2D), with no germ cells of any types but only remnants of spermozoa could be found. This indicates that spermatogenesis process in the testis of infected individuals was completely arrested. Further, similar to the case of hepatopancreas of infected individuals, infected testis was heavily infiltrated with \(S. \text{beauforti}\)’s rootlets.

Several types of cells could be found in the ovarian tissues of healthy females, i.e., oogonia, primary oocytes, early-maturing oocytes, and late-maturing oocytes, all of which are densely packed and surrounded by follicle cells (Figure 2E). The ovarian tissues of infected individuals, however, were acutely damaged (severe necrosis) and no rigid cell structures could be observed (Figure 2F).

In line with the degradation of gonad and disruption of gametogenesis, significant reduction in gonad overall sizes were observed among healthy and infected males and females (ANOVA; \(F_{3,47.291} = 83.335, P < 0.001\)). The GSI of infected males and females were significantly smaller than their healthy counterparts (Games Howell; \(P_{\text{male}} < 0.001\)). However, although hepatopancreases of crabs with mature externa were also disrupted, opposite trend was observed in their overall sizes (Figure 3). Differences in HSI values among healthy and infected males and females were obvious (ANOVA; \(F_{3,48.877} = 125.061, P < 0.001\)), with that of infected males and females significantly larger than their healthy counterparts (Games Howell; \(P_{\text{male}} = 0.001\)) and no significant difference was observed between the HSI values of infected males.
and females (Games Howell; $P = 0.4245$). A multiple regression was conducted to predict infection status from sex, CW, BW, GSI, and HSI. These variables significantly predicted infection status, $F_{5, 95} = 119.867, P < 0.001, R^2 = 0.856$. However, only three variables (CW, $P = 0.001$; GSI, $P = 0.003$; HSI, $P < 0.001$) added statistically significance to the prediction.

**DISCUSSION**

The high prevalence of *S. beauforti* in a natural mud crab population has been alarming, as this population is known to supply crabs, both domestically and internationally, to fisheries and aquaculture sectors (Waiho et al., 2017b). *Sacculina beauforti* belongs to the family Sacculinidae, a family with the most abundant rhizocephalan species (Høeg, 1995).

The internal anatomy of externa provides valuable information on the taxonomy and reproductive biology of the invading parasite (Glenner and Hebsgaard, 2006). Understanding it could provide clues on the future development of prevention or treatment methods, especially in such case where the host (*S. olivacea*) is an economically important and highly valuable aquaculture species. The externa anatomy of *S. beauforti* conforms with the classical externa description of most sacculinids (Lützen and Takahashi, 1997; Alvarez et al., 2009; Kobayashi et al., 2018). The double-layered mantle near the mantle opening of *S. beauforti* provides additional muscle tissues to perform strong contractions during oviposition, fertilization and expulsion of nauplii (Hoeg and Lützen, 1995; Alvarez et al., 2009).

Hepatopancreas plays an essential role in nutrient absorption and storage in crustaceans (Wu et al., 2020). In addition, hepatopancreas stores lipid, an essential nutrient that could provide huge amount of energy during critical moments, such as during starvation, molting, maturation or reproduction (Wang et al., 2014). It is also involved in the ovarian development by regulating the synthesis of vitellogenin and vital sex steroid hormones, including that of mud crab genus *Scylla* (Warrier et al., 2001; Jia et al., 2013). Due to these critical roles, the whole organ is rich in nutrient and the rhizocephalan *S. beauforti* takes advantage of this – penetrating intertubular spaces within the hepatopancreas of its host with numerous rootlets, potentially absorbing nutrient directly from it. Similar finding of sacculinid rootlets in the hepatopancreas of its host was also reported in portunid blue crab *Callinectes sapidus* infected by *Loxothylacus texanus* (Bortolini and Alvarez, 2008) and shore crab *Carcinus maenas* infected by *Sacculina carcini* (Powell and Rowley, 2008). The adverse metabolic effect of *S. beauforti* on *S. olivacea* is postulated to be at the maximum when externa matures (Robles et al., 2002). Both healthy and infected individuals exhibited similar tubule diameters and with the presence of tubular R- and B-cells. R-cells are involved in nutrient absorption and...
FIGURE 2 | The hepatopancreas and gonad histological cross-sections of healthy (A,C,E) and infected Scylla olivacea (B,D,F) (Mayer’s Hematoxylin-Eosin staining). (A) Healthy hepatopancreas with normal tubule shape and arrangement, magnification 10×; (B) Infected hepatopancreas with the presence of rhizocephalan rootlets (RL) and collapsed tubules (CT), magnification 10×; (C) Healthy testis with the presence of spermatogonia (SPG), spermatocytes (SPC), spermatids (SPT) and spermatozoa (SPZ), magnification 10×; (D) Infected testis with the presence of RL, almost no spermatogenesis activity, cell lysis and only remnants of SPZ could be found, magnification 10×; (E) Healthy ovary with the presence of oogonia (OO), primary oocytes (PO), early-maturing oocytes (EO) and late-maturing oocytes (LO), with follicle cells (FC) surrounding oocytes, magnification 20×; (F) Infected ovary appeared degenerated and no rigid cell structures were observed, magnification 4× (large panel) and 20× (upper right panel). Connective tissues (C), R-cells (RC), B-cells (BC).
FIGURE 3 | Boxplots showing the gonadosomatic index (GSI, %) and hepatosomatic index (HSI, %) of healthy and infected (A) males and (B) females. All comparisons showed significant differences ($P < 0.05$, marked with asterisk *). Infection by Sacculina beauforti resulted in significant reduction in GSI but increment in HIS of Scylla olivacea.

metabolization, lipid and glycogen storage, and is crucial in the detoxification process and uric acid excretion (Vogt and Quinitio, 1994; Johnston et al., 1998; Vogt, 2019); B-cells play important role in intracellular digestion and assimilation (Al-Mohanna and Nott, 1986) as they might be involved in the production and recycling of fat emulsifiers (Vogt, 2019). This implies that although S. beauforti's infection and the infiltration of numerous rootlets might occupy most intertubular spaces and resulted in an increase in mass (as evident by the significant increase of infected individuals’ HSI and decrease in tubular counts), the tubular structure integrity of some hepatopancreatic tubules remains intact and might still be functional. The preservation of the function of host's hepatopancreas is vital for the survival of rhizocephalans as it is their main nutrient absorption site and its destruction will result in the death of the host.

The gonads of hosts, however, were even more affected. Unlike healthy gonads with various types of germ cells (Quintiio et al., 2007; Azmie et al., 2017; Waiho et al., 2017c), the testes of infected individuals were significantly reduced in size, degenerated and non-functional (no active germ cells were found), indicating that spermatogenesis was completely inhibited. In addition, ovaries of infected S. olivacea were so severely damaged that no rigid cell structures could be observed. Similar gonadal atrophy and disruption of vitellogenesis and spermatogenesis are commonly reported in sacculimid-infected hosts (Rubiliani, 1983; Hoeg, 1995; Isaeva et al., 2001). The extent of damage, however, varies among species. For example, Polyascus polygenea disrupted spermogenesis but not spermatogenesis in the testis of its host Hemigrapsus sanguineus (Isaeva et al., 2001; Glenner et al., 2003); and in C. maena parasitized by S. carcini, virtually no weight difference could be found in the testes of infected and healthy crabs, - and nothing indicated that spermatophores of infected crabs would not be functional (Zetlmeisl et al., 2011). The severe damage observed in the ovaries of infected S. olivacea in this study is similar to that reported in spider crab Macropodia longirostris infected by sacculinid Drepanorchis neglecta (Hartnoll, 1962), where disintegration of ovarian cells and total ovary degeneration (in some cases) were observed. The size of testes and ovaries of infected hosts (S. olivacea) with mature externa were extremely reduced, to the extent that the mean of their GSI values (mean_{male} = 0.06 ± 0.02%; mean_{female} = 0.26 ± 0.14%) were even smaller than the mean GSI values of immature males (mean = 0.10 ± 0.04%) (Waiho et al., 2017c) and females (mean = 2.46 ± 1.31%) (Azmie et al., 2017) of the same species reported in previous studies. Based on the size (CW) of the hosts collected in this study, they should be sexually mature with fully developed gonads (Waiho et al., 2016; Fazhan et al., 2017). However, S. beauforti infestation resulted in gonadal atrophy of both sexes, to such degree that the hosts are physiologically infertile.

Gonadal development and maturation are essential processes in the life history of crustaceans (Waiho et al., 2017a). Aside from being directly involve in reproduction, gonads are responsible for the morphological differentiation of most crustacean species.
which leads to sexual dimorphism (Parvizi et al., 2017; Waiho et al., 2019). Therefore, it is possible that rhizocephalans induce morphological changes (feminization) by negatively regulating the development of gonads in hosts as gonads play vital role in sex differentiation of an organisms, including crustaceans (Waiho et al., 2019). For example, doublesex gene (Dsx)—a male sex-determining gene—is found in the testes of crustaceans and involves directly in the regulation of sexual dimorphism and the silencing of Dsx gene in male Daphnia magna embryos is known to induce the development of female secondary characteristics (Kato et al., 2011). Another important organ in crustaceans that is involved in sexual dimorphism is the androgenic gland (AG). As demonstrated in intersex individuals of red-claw crayfish Cherax quadricarinatus (Rosen et al., 2010) and juveniles of freshwater prawn Macrobrachium rosenbergii (Ventura et al., 2009), the silencing of an AG specific insulin-like gene (Cq-IAG in C. quadricarinatus; Mr-IAG in M. rosenbergii) resulted in feminization (impede the regeneration of male secondary sexual characteristics) and severe testicular degeneration – two characteristics similar to the effects induced by rhizocephalans on their respective hosts. By injecting healthy male crabs Rhithropanopeus harrisii with crude extracts of roots from sacculinid Loxothylacus panopei, Rubiliani (1983) showed that proteinaceous substances from rootlets are responsible, not only for the inhibition of spermatogenesis, but cytolysis of androgenic gland and sinus gland as well. Thus, future research on the potential effect of S. beauforti infestation on S. olivacea's AG and in-depth study on the molecular changes in gonads and AG after infected are recommended to uncover the regulatory mechanism of rhizocephalans that would result in feminization and gonad degeneration in infected hosts.

The mud crab S. olivacea serves as important fisheries commodity and aquaculture candidate in Southeast Asia where this species is abundantly found. Infestation of rhizocephalan barnacles (S. beauforti) on S. olivacea did not only affect external morphologies, but also severely damaged their internal organs, rendering them infertile. This could adversely affect their market price as mature females with ripe ovaries are known to fetch significantly higher price compared to males of the same size (Waiho et al., 2020). Further, since aquaculture activities such as crab fattening, soft-shell crab production and hatching rearing of mud crab still rely heavily on sourcing juveniles and broodstocks from the wild, they could be render fruitless if infected individuals are accidentally sourced as infected juveniles are more susceptible to other infections and have a higher chance of mortality (Keogh et al., 2017) while broodstocks are not able to produce any larvae due to the degenerated gonads. Thus, farmers would endure great losses—financial, effort and time—if infected crabs were unknowingly cultured. Also, as most mud crab aquaculture activities are conducted in large earthen ponds with direct water intake from nearby brackish mangrove estuary ecosystems, and infected hosts might not be aware of the presence of infected individuals being co-cultured with healthy individuals. Eventually, this could lead to an outbreak of S. beauforti infestation within the culture ponds if mature externae were fertilized by free-living male cyprids (Glenner et al., 2003; Kobayashi et al., 2018) potentially introduced via the water intake system. Therefore, if infected individuals were detected during culture (i.e., with the presence of externa), especially in facilities where water intake is unfiltered and untreated, it is important to remove and discard (kill) them to prevent unwanted outbreaks (Waiho et al., 2021).

CONCLUSION

Although it is still early to demonstrate, unequivocally, the total adverse effect of rhizocephalan barnacle S. beauforti have on its host S. olivacea, the high prevalence of this parasite on an economically important aquaculture crustacean species, and the severe damage it causes on two essential organs—hepatopancreas and gonads—warrant an urgent report to highlight to researchers, farmers and other industry players on its emergence and manifestation. It is hoped that through this report, occurrence of S. beauforti on mud crab genus Scylla in other geographically distinct populations would be reported to aid in understanding S. beauforti's distribution, seasonal abundance, life cycle and pathology, all of which are important for future research into its prevention and cure.

DATA AVAILABILITY STATEMENT

The datasets generated for this study are available on request to the corresponding author.

AUTHOR CONTRIBUTIONS

HF and KW conceived and designed the research, performed the field work, and wrote the manuscript. MH and JM processed the data. HG contributed to the further analysis of data. HG, MH, and MI revised the manuscript and contributed to the writing. All authors contributed to the article and approved the submitted version.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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