Stimulation of Ovarian Development in a Tropical Damselfish by Prolonged Photoperiod using Pellets Containing Long-afterglow Phosphorescent Pigment

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Abstract
The present study examined whether light emitted by long-afterglow phosphorescent pigments (LumiNova) would stimulate gonadal development in fish during the nonbreeding season. Pellets containing LumiNova powder (treatment group) were prepared and placed on the calvaria of specimens of the sapphire devil Chrysiptera cyanea, a reef-associated damselfish that requires long days for gonadal recrudescence. A pellet without LumiNova powder was placed on the calvaria of the control fish (control group). Fish were reared at 26°C under a light–dark cycle (12 h photophase, 12 h scotophase; LD 12:12) for 4 weeks. No difference in the gonadosomatic index (GSI) or ovarian histology was observed among the control, sham-operation, and treatment groups 1 week after the start of the experiment. After 4 weeks, the GSI of the control and sham-operation groups remained at low levels, and ovaries contained immature oocytes at the perinucleolus stage. In contrast, the treatment group exhibited significantly higher values of GSI as well as developed ovaries with fully vitellogenic oocytes. These results demonstrated that long-day conditions were produced by light emitted from the LumiNova pellets, thus stimulating ovarian development in the damselfish. Therefore, long-afterglow phosphorescent pigments can be used as an alternative to standard light sources for purposes of artificial stimulation of gonadal development in fish.

Key words: Chrysiptera cyanea, Gonadal development, LumiNova, Photoperiod, Ovarian development

Introduction
Seasonal changes in day length are critical factors affecting the initial cueing, timing, and subsequent synchronization of gonadal development in various fish species (Bromage et al., 2001; Migaud et al., 2010). Reproductive strategies involving day length are likely to be diverse among species. For example, a steady increase in day length stimulates gonadal development in the black seabass Centropristis striata (Howell et al., 2003), the Japanese amberjack Seriola quinquera-diata (Hamada and Mushiake, 2006), and the Eurasian perch Perca fluviatilis (Migaud et al., 2003), whereas a decrease in day length serves as the trigger for gonadal development in the Atlantic cod Gadus morhua (Skjaeraasen et al., 2004) and most salmonid species (Bromage et al., 2001). In both of these cases of photoperiodicity in fish, changes in day length (i.e.,
long-day or short-day conditions) are believed to be perceived by the photosensory organ(s) and transduced as an internal signal, and gonadal development is subsequently stimulated by the cascade activation of the hypothalamus–pituitary–gonadal (HPG) axis at an appropriate time of a year. Melatonin secreted by the pineal organ appears to function as a potent transducer of photic information to central and peripheral organs, as demonstrated by seasonal changes in diurnal melatonin profiles in some fish species (Randall et al., 1995). However, how this indoleamine hormone regulates the HPG axis remains unclear (Bromage et al., 2001).

Artificial stimulation or suppression of gonadal development by manipulating photoperiodic conditions offers potential industrial advantages for efficiently breeding fish with high commercial value at a desired time of year and for reducing the excessive use of drugs and hormones for maturation. Previous studies have used incandescent, fluorescent, metal-halide, and tungsten-halogen lamps as standard light sources and have successfully controlled the gonadal development of various fish species (Bromage et al., 2001). In terms of long-operating life and low-energy consumption, light-emitting diodes (LEDs) have recently been used as a new light source to stimulate gonadal development (Bapary et al., 2011; Leclercq et al., 2011) and growth (Yamanome et al., 2009). However, waterproofing and barotolerance issues limit the usage of this new light technology in aquaculture.

Recently, long-afterglow phosphorescent pigments (LumiNova) have been used as a light source, and light emitted by LumiNova sheets was able to prolong the reproductive season of the sapphire devil Chrysiptera cyanea (Bapary et al., 2012), a reef-associated damselfish that requires long days for gonadal recrudescence (Bapary et al., 2009). Additional advantages of the glow-in-the-dark pigments of LumiNova include the following: 1) elimination of the need for an electrical energy supply, 2) their activation by a broad band of wavelengths, 3) their high emission level relative to the amount of radiation, 4) their chemical stability and lack of hazardous and radioactive substances, and 5) their light- and waterproof qualities (NEMOTO, 2000). The objective of the present study was to improve the methodology for LumiNova application and to stimulate gonadal development in the sapphire devil during the nonbreeding season.

Materials and Methods

Fish and experimental design

Adult sapphire devils of 3.5 ± 0.4 cm mean body length and 1.3 ± 0.6 g mean body mass were collected from coral reef lagoons (26°25′74.7″N, 127°68′91.3″E) in Urasoe, Okinawa, Japan, using a small round haul net during daytime low tide. Fish were transferred to and reared in aquaria equipped with filtering and aeration systems under short-day conditions (10 h photophase and 14 h scotophase; LD 10:14, light on at 08:00 and off at 18:00) and a water temperature of 26°C. A fluorescent bulb (14 W) was used as a light source, and the light intensity at the water surface was 3.6 W/m². Fish were fed daily at 10:00 with commercial pellets (EP1; Nisshin-Marubeni, Tokyo, Japan).

LumiNova (G-300M; Nemoto Lumi-Materials, Tokyo, Japan), epoxy resin (craft resin) and hardener (Nisshin Resin KK, Yokohama, Japan) were mixed at a ratio (w/v) of 2:1:1. The mixture was poured into a silicone tube (internal diameter = 2 mm, external diameter = 3 mm) and left for 6 h at room temperature. After drying and hardening, the silicone tube containing LumiNova was cut into approximately 2-mm sections (Fig. 1A and 1B), which were kept at room temperature until use. Control pellets were prepared by mixing only the epoxy-resin and its hardener at a ratio of 2:1.

The experiment was conducted for 1 month beginning in December 2011, during the nonbreeding season of the damselfish and when their ovaries are exclusively occupied by immature oocytes (Bapary et al., 2012). After fish (n = 60, male:female = 1:5) were anesthetized with ethyl-3-amino-benzoatemethanesulfonic acid (MS-222; Sigma-Aldrich, St. Louis, MO, USA), a pellet was sewn on the head of each individual (Fig. 1C). Prepared fish were separated into three groups: the treatment group (3 males and 15 females), control pellet group (3 males and 15 females), and sham-operation group (4 males and 20 females). Each group was housed in an...
aquarium equipped with a filtering and aeration system under LD 12:12 conditions (lights on at 06:00 and off at 18:00) and a water temperature of 26°C. All fish were fed daily with EP1. Subsequently, 80- to 100-mm-diameter pipes were placed on the bottom of each aquarium as a spawning nest. Fish (five females per aquarium) were sampled at 1 and 4 weeks after the start of the experiment. Ovaries from five females were also collected from the sham-operation group just before the onset of the experiment. The original ratio of males to females was maintained by removing males at each sampling time. Fish were anesthetized with 2-phenoxyethanol (Kanto Chemical Co., Tokyo, Japan) and decapitated. The body mass and body length of each individual were recorded. The ovaries were removed from the body cavity, weighed, and subsequently fixed in Bouin’s solution. The gonadosomatic index (GSI) was calculated using the following formula: GSI = (ovarian mass/body mass) × 100.

All experiments were conducted in compliance with the guidelines of the Animal Care and Use Committee of the University of the Ryukyus and the regulations for the care and use of laboratory animals in Japan.

Histological procedures

A portion of the fixed ovary was dehydrated in an ethanol series, permutated with xylene, and then embedded in paraffin (Nacalai Tesque Inc., Kyoto, Japan). Serial sections (7 µm) were prepared and stained using Mayer’s hematoxylin–eosin for microscopic observations. Oocyte development was classified into six stages: the perinucleolus stage, oil droplet stage, yolk vesicle stage, primary yolk stage, secondary yolk stage, and tertiary yolk stage (Bapary et al., 2009).

Statistical analysis

The GSI results were expressed as the mean ± standard error of the mean (SEM). A two-way analysis of variance (ANOVA) followed by a Tukey–Kramer test was used for comparing the mean GSI among fish groups (P < 0.05 for a statistically significant difference).

Results

No fish died during the experimental period. The GSI of the initial control was 0.64 ± 0.12. The GSI of the treatment group increased to 0.70 ± 0.10 after 1 week and to 1.88 ± 0.37 after 4 weeks. The GSIs of the control pellet group and sham-operation group did not change over time and were 0.98 ± 0.11 and 0.92 ± 0.07 after 4 weeks, respectively. After 4 weeks, the GSI of the treatment group was significantly higher than that of the other two groups (Fig. 2).

Ovaries of the initial control contained immature oocytes at the perinucleolus stage. Similar ovarian features were confirmed in the control pellet and sham-operation groups at 1 week after the start of the experiment (Fig. 3A). In contrast, ovaries of the treatment group had oocytes at the perinucleolus and oil droplet stages after 1 week (Fig. 3B). After 4 weeks, ovaries of the control pellet and sham-operation groups were occupied only by oocytes at the perinucleolus stage (Fig. 3C), whereas those of the treatment group contained fully vitellogenic oocytes at the tertiary yolk stage (Fig. 3D).

Discussion

The present study clearly demonstrated that light emitted by long-afterglow phosphorescent pigments (LumiNova) stimulated ovarian development in the sapphire devil during the nonbreeding season. No ovarian development was induced during the experimental period in either control or sham-operation fish. Similar results have been obtained using the same species: fish in aquaria covered with LumiNova sheets continued active oocyte growth and repeated spawnings even after the reproductive season of naturally reared fish had terminated (Bapary et al., 2012). Thus, LumiNova emits wavelengths of light that can be perceived and utilized by the fish.

Histological observations have indicated that ovarian development of the sapphire devil in Okinawan water initiates in March and peaks in May, when both the photoperiod and water temperature are increasing (Bapary et al., 2009). When females were reared at 25°C under experimental conditions of LD 10:14, LD 12:12, and LD 14:10 using fluorescent bulbs as a light source, ovaries with fully vitellogenic oocytes were only observed under long-day conditions (Bapary et al., 2009). In addition, exposing female sapphire devils to long-
altivelis altivelis. In addition, Northern blot and reverse transcription–polymerase chain reaction (RT-PCR) analyses have revealed that rhodopsin is expressed in the brain of the ayu sweetfish (Masuda et al., 2003). When ophthalmectomized sapphire devils were reared under long-day conditions (LD 14:10), vitellogenic oocytes appeared in the ovary (Bapary, 2011). These previous findings indicate that extraretinal photoreceptor(s) play a role in inducing gonadal development. In situ hybridization using the sapphire devil brain revealed that long-wavelength sensitive cone opsin (LWS) mRNA is expressed in the third ventricle periventricular area in the anterior hypothalamus and that the strong signals of this LWS mRNA are predominantly observed in the ventromedial thalamic nucleus (VM), anterior periventricular nucleus (NAPv), and suprachiasmatic nucleus (NSC) (Takeuchi et al., 2011). In addition, the expressions of both middle-wavelength sensitive cone (MWS) and rhodopsin mRNA were detected in the brain of this species using RT-PCR (Takeuchi, 2012). Taken together, these opsins are likely to be involved in the perception of green light emitted by LumiNova. However, the possibility that the eyes play a role in transducing light signals from LumiNova cannot yet be excluded, as light from the pellet on day conditions (LD 14:10) with red (627 nm) and green (530 nm) LED lights induced ovarian development, whereas exposure to blue (455 nm) and white LED lights did not (Bapary and Takemura, 2010). These previous reports clearly suggest that the sapphire devil perceives long-day conditions from LED lights and begins ovarian development during the natural nonbreeding season. In addition, mid- to long-wavelength lights are preferable for initiating ovarian development in this species. Because LumiNova emits green light (530 nm) in the dark after energy absorption and emits light for several hours thereafter (NEMOTO, 2000), fish are likely exposed to the desired wavelengths of light after lights are turned off. Because the photoperiodic conditions of LD 12:12 were used in the present study, the additional emission of light by LumiNova produced long-day conditions, which stimulated gonadal development.

Because the pellet was placed on the calvaria of each individual, the majority of light emitted by LumiNova likely stimulated the extraretinal photoreceptors. For example, Masuda et al. (2005) demonstrated that following pinealectomy and ophthalmectomy, gonadal development was induced under short-day conditions in the ayu sweetfish Plecoglossus altivelis altivelis. In addition, Northern blot and reverse transcription–polymerase chain reaction (RT-PCR) analyses have revealed that rhodopsin is expressed in the brain of the ayu sweetfish (Masuda et al., 2003). When ophthalmectomized sapphire devils were reared under long-day conditions (LD 14:10), vitellogenic oocytes appeared in the ovary (Bapary, 2011). These previous findings indicate that extraretinal photoreceptor(s) play a role in inducing gonadal development. In situ hybridization using the sapphire devil brain revealed that long-wavelength sensitive cone opsin (LWS) mRNA is expressed in the third ventricle periventricular area in the anterior hypothalamus and that the strong signals of this LWS mRNA are predominantly observed in the ventromedial thalamic nucleus (VM), anterior periventricular nucleus (NAPv), and suprachiasmatic nucleus (NSC) (Takeuchi et al., 2011). In addition, the expressions of both middle-wavelength sensitive cone (MWS) and rhodopsin mRNA were detected in the brain of this species using RT-PCR (Takeuchi, 2012). Taken together, these opsins are likely to be involved in the perception of green light emitted by LumiNova. However, the possibility that the eyes play a role in transducing light signals from LumiNova cannot yet be excluded, as light from the pellet on

Fig. 3. Changes in ovarian histology of the sapphire devil (Chrysiptera cyanea) with a pellet on the calvaria. Ovaries were collected at 1 and 4 weeks after the onset of the experiment. A cross section of an ovary (CSO) of (A) the control pellet group after 1 week, (B) CSO of the treatment group after 1 week, (C) CSO of the control pellet group after 4 weeks, and (D) CSO of the treatment group after 4 weeks. ODS, oil droplet stage; PNS, peri-nucleolus stage; TYS, tertiary yolk stage. Scale bar = 200 μm.
the head of one individual may be perceived through the eyes of other individuals.

Our series of trials using tropical damselfish is the first report to demonstrate the effectiveness of long-afterglow phosphorescent pigments in the artificial control of gonadal activity. Previous approaches have involved irradiation of the entire aquarium by light emitted from LumiNova (Bapary et al., 2012), whereas the present study used irradiation focused onto the calvaria of fish. Despite this technical difference in the application of LumiNova, this approach offers a valuable tool for controlling gonadal activity in fish and can serve as an energy-free and environmentally safe method in aquaculture. Further studies are necessary to examine whether LumiNova could be widely applicable to other important fisheries species.

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References


