Plasticity of Locomotor Activity Permits Energy Homeostasis During Reproduction in a Female Sea Cucumber

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Behavioral plasticity in animals allows for moment-by-moment behavioral adjustments to biotic and abiotic uncertainties in the environment. For most aquatic animals, reproduction involves gonadal development and other physiological changes, causing increasing demands in nutrients and energy for females. The aim of this report was to determine how the female sea cucumber, Apostichopus japonicus, accommodates increasing energy demands during reproduction by adopting a behavioral energy conservation strategy. Dynamic changes in feeding activity, total body mass, locomotor activity, energetic condition, and metabolic performance of the females were measured from the non-breeding stage up to the mature stage. Routine metabolic rate analysis showed that reproduction caused a significant increase in energy demands in the adult. However, reproduction also suppressed the appetite of adults resulting in decreased energy intake. Interestingly, combining time-lapse camera and behavioral analysis software, the results showed that sea cucumbers down-regulated their locomotor activities in order to conserve energy effectively. Energy budget analysis and unchanged cortisol level revealed that the reduction in motility offset the increased energy demand for reproduction while helping to maintain energy homeostasis in the maternal body. Altogether, our study suggests that plasticity in the locomotor behavior allows sea cucumbers to cope with the high energy demands of reproduction.

Keywords: sea cucumber, behavioral plasticity, energetic physiology, energy homeostasis, reproduction

INTRODUCTION

Reproduction is one of the most important events of all organisms. However, reproduction is energetically costly (Speakman, 2008). For example, gonadal development, metabolic maintenance, behavioral thermoregulation, and brood care require the expenditure of considerable amounts of energy in many aquatic animals (Baeza and Fernández, 2002; Fossette et al., 2012; McBride et al., 2015; Duisan et al., 2021). As a result, activities associated with reproduction pose energetic challenges, particularly to females.
Maintaining energy homeostasis is critical for development, growth, reproduction, and survival (Unniappan et al., 2020). Interestingly, as a response to variations in energy or nutrient demands at different life stages (i.e., reproduction), animals have evolved the capacity to adjust their behavioral and physiological functions in order to accommodate changing energy demands (Nie et al., 2015). For example, plasticity in physiological adjustment involves remodeling gut structure, and adjusting digestive strategy to increase digestive and absorptive efficiency (Krockenberger and Hume, 2007; Reiff et al., 2015). In addition, behavioral adjustments may involve expanding the home range, increasing foraging time, and shifting prey choice to increase energy intake (Barclay, 1989; Rodriguez et al., 2010; Sokolov et al., 2014).

Locomotor ability is a crucial necessity for survival. However, the development of eggs or fetuses places an additional physical burden on females (McLean and Speakman, 2000; Miles et al., 2000; Pruitt and Troupe, 2010). As a result, loss of or decrease in locomotor performance has been widely reported in terrestrial animals over the past several decades. Due to buoyancy, growing eggs may have no effect on locomotor ability in aquatic animals, especially in marine invertebrates that have a large body surface area. However, decreased locomotor activity under certain circumstances has been observed in many aquatic organisms during reproduction, including fish, echinoderms, aquatic snakes, and sea turtles (James and Johnston, 1998; Winne and Hopkins, 2006; Fossette et al., 2012; Ru et al., 2017). Recently, the behavioral energy conservation hypothesis has been proposed in order to explain unexpected behavior in aquatic animals (Fossette et al., 2012). This hypothesis states that plasticity of locomotor behavior allows animals to down-regulate their motility in order to conserve energy for other functions (such as reproduction and growth), thereby maximizing their fitness (Fossette et al., 2012; Okuyama et al., 2013). However, there is little physiological evidence to support this hypothesis.

In the present study, the sea cucumber, Apostichopus japonicus, was used as model system to test the behavioral energy conservation hypothesis during reproduction. Recent studies have reported that A. japonicus shows a series of behavioral and physiological changes after the onset of the annual reproductive cycle (Ru et al., 2017, 2021). Since both locomotion and egg production are high energy demand functions, we hypothesized that decreasing locomotor activity can compensate for the increased energy demand of reproduction. The behavioral energy conservation strategy might also have additional benefits for breeding females, such as maintaining maternal energy homeostasis. However, the relationship between reproduction, locomotor behavior, and energetic physiology remains unclear and requires further testing.

To elucidate this point, we analyzed the dynamic interaction of feeding activity, locomotor activity, metabolic performance, growth function, gonadal development, and energetic condition in female sea cucumber during reproduction. We also used a mechanistic model to show how plasticity of locomotor function maintains energy homeostasis during reproduction. Our study provides new insights into the interaction of locomotor behavior and energetic physiology during reproduction in sea cucumber.

MATERIALS AND METHODS

Animals and Rearing Conditions

In this study, sea cucumbers were collected in May 2017 from Rushan Bay, China (36° 46′ N, 121° 34′ E). Animals were induced to spawn using the air-drying method (Liu et al., 2015). Only females were used in this study, since feeding activity, immune function, and locomotor activity of females may be affected by the pheromones of males (Jiang et al., 2017, 2020; Marquet et al., 2018). Identification of females was conducted by observation of oocyte release during spawning. After spawning, the females were identified, selected, and reared in three 5 m × 3 m × 2 m cages in an open-air pond located in the National Algae and Sea Cucumber Engineering Technique Research Center of China until November. Prior to the start of the annual reproductive cycle, 34 females (126.93 ± 5.02 g) were collected randomly as the experimental animals.

The experiment lasted for 100 days. Prior to the study, 10 sea cucumbers were sampled as the non-breeding group. During the study, procedures were conducted every 25 days as follows: (1) All individuals (n = 24) were weighted to assess growth dynamics. (2) Six adults were randomly selected for behavioral activity recordings. (3) Six adults were randomly selected to measure metabolic performance.

During the study, sea cucumbers were reared in four 350 L plastic aquaria, with the stocking density of six individuals per tank. The animals were fed to satiation at 18:00. Feces were siphoned from each tank at 8:00 am daily, and then frozen at −20°C for future analysis. Additional conditions: water temperature was 14°C controlled by a running water system, pH was ∼7.9, dissolved oxygen concentration was 8.0 mg L⁻¹, and the light intensity was 25 lux.

Locomotor Activity Analysis

At each stage, six females were randomly selected and individually placed in a white plastic cylinder (50 cm in diameter) for 24 h. Movement of each animal was recorded by a TLC 200 Brinno time-lapse camera. Environmental factors have a significant effect on the behavioral performance of sea cucumbers (Pan et al., 2015; Sun et al., 2018). Therefore, water temperature and light intensity in the cylinders were kept the same as rearing tanks throughout the recording.

Videos were saved as AVI files of 680 × 480 pixels and analyzed by XT Ethovision 10.1 software (Noldus Information Technology, Wageningen, Netherlands). The color difference between the animal and the background was used to catch and trace the movement of the sea cucumber during the behavioral analysis. The total locomotor distance in 24 h and total locomotor time in 24 h were used as functional parameters to assess the locomotor activity of sea cucumbers during the experiment.

Measurement of Metabolic Performance

The routine metabolic rate (RMR) of sea cucumbers was measured using an established method (Dong and Dong, 2006). Six sea cucumbers were randomly selected and each individual was placed in a 2.8 L sealed respiratory chamber (Subo Co.,
was calculated following the equation (Dong and Dong, 2006):

\[
RMR(\mu gO_2g^{-1}h^{-1}) = 1000(D_f - D_o)V/WT
\]

where \(D_f\) is the change in oxygen level (mg O_2 L^{-1}) in the test respiratory chamber with animal, \(D_o\) is the change in oxygen level (mg O_2 L^{-1}) in the test respiratory chamber without animal, \(V\) is the volume of the respiratory chamber (L), \(W\) is the total body weight of the test sea cucumber (g), and \(T\) is the test time (h).

**Feeding Activity**

Daily food consumption (FC) and ingestion rate (IR) were used to assess the feeding activity of females during reproduction. FC and IR were determined using acidic insoluble ash markers (AIA, a natural indicator of food intake) in feces and diet according to an established protocol (Ru et al., 2018). A 1 g sample was carbonized using a muffle furnace for 2 h at 300°C to remove organic matter. The ash was acidized, filtered, and then burned again at 600°C for 15 h to obtain the AIA value.

**Sample Collection and Determination of Cortisol Level**

All individuals were euthanized by decollation on day 100. The gonad, body wall, respiratory tree, and gut of each animal were separated immediately under sterile conditions. In addition, a total of 10 individuals were randomly selected for sample collection (i.e., two, three, three, and two individuals were sampled from tank1, tank2, tank3, and tank4, respectively). 2 mL of coelomic fluid from each individual was removed with a sterile syringe for cortisol determination, and the whole gonad and somatic tissues (i.e., body wall, gut and respiratory tree) of each individual were weighted and frozen at −20°C for measurement of energy density.

Cortisol levels were determined by the enzyme linked immunosorbent (ELISA) assay method. We purchased the commercial ELISA Assay Kit from Nanjing Jiancheng Bioengineering Institute (Nanjing, China), and the determination was performed under the manufacturer's guidelines. The ELISA Assay Kit has been validated in many previous studies (Xu et al., 2020).

In details, prior to assay, coelomic fluid was clarified by centrifugation at 2,000 g for 10 min at 4°C. And then 40 µL supernatant of each sample was mixed with 10 µL cortisol antibody and 50 µL streptavidin-HRP in the test well. After the test well was sealed, the processed sample was incubated for 60 min in a water bath at 37°C. After the procedures of washing and chromogen for 10 min, the color development was terminated by stop solution. Finally, the cortisol level was determined at 450 nm using a DG5033A microplate reader (Huadong Electronics Group, Nanjing, China).

**Energy Budget**

Energy density of all samples, including somatic tissues, gonadal tissues, diet and feces, were measured using an IKA C600 calorimeter (IKA®-Werke GmbH & Co., KG, Staufen, Germany).

The daily activities of sea cucumbers are very simple, they move randomly to search food at night, and rest under a shelter during the day. A. japonicus is an income breeder (Ru et al., 2021). With the exception of egg development, energy from ingested food during reproduction also is invested into physiological metabolism, locomotor function, and growth function. Most unused nutrients were discharged from the body as feces (Yuan et al., 2007).

Therefore, an energy budget equation was constructed to investigate how sea cucumbers adjust their locomotor function to maintain energy homeostasis during reproduction as follows (Yuan et al., 2007, 2016):

\[
C = F + R + B + P + G 
\]

where \(C\) (J ind^{-1} day^{-1}) is the energy intake from ingested resources during reproduction, \(F\) (J ind^{-1} day^{-1}) is discharged energy as feces, \(R\) (J ind^{-1} day^{-1}) is energy allocated into reproduction (i.e., gonadal growth), \(P\) (J ind^{-1} day^{-1}) is energy lost in maintaining metabolic function, \(B\) (J ind^{-1} day^{-1}) is energy used in locomotor function, and \(G\) (J ind^{-1} day^{-1}) is the deposited energy as somatic reserves. The six parameters were calculated as follows:

\[
C = FC \times E_{\text{diet}}; F = FP \times E_{\text{feces}} 
\]

where \(E_{\text{diet}}\) and \(E_{\text{feces}}\) are the energy density of diet and feces, \(FC\) is the daily food consumption, and \(FP\) is the daily feces production.

\[
R = \Delta W_g \times E_{\text{gonad}}; G = \Delta W_s \times E_{\text{soma}} 
\]

where \(E_{\text{gonad}}\) and \(E_{\text{soma}}\) are the energy density of gonadal tissues and somatic tissues, and \(\Delta W_g\) and \(\Delta W_s\) are the increased weight of gonadal and somatic tissues, respectively. The growth rate of somatic tissues and gonadal tissues of sea cucumbers at different breeding stages (every 25 days over 100 days) were 48.81, 28.53, 13.4, 9.27% and 7.53, 23.66, 33.79, and 35.02%, respectively (Ru et al., 2018).

Energy used to maintain routine metabolic function was evaluated by nitrogen excretion. Sea cucumbers use 24,830 J of energy to excrete 1 g nitrogen (Yuan et al., 2016). The ratio of oxygen consumption to nitrogen excretion is 1:15 (Yuan et al., 2007).

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\[
P = RMR \times 15 \times 24830 \times 24 
\]

Lastly, the value of \(B\) was calculated by:

\[
B = C - F - R - P - G 
\]
Statistical Analysis

Sea cucumbers show great individual variations in growth and physiological condition (Liang et al., 2010). Therefore, total body mass, body wall mass, gonad mass, metabolic rate, and locomotor activity were analyzed at the individual level. It is not possible to distinguish each cucumber for food ingestion and feces production in a same tank. Therefore, feeding activity and parameters in energy budget equation were analyzed between different tanks at different breeding stages. In order to avoid the pseudoreplication for the final analysis, data of cortisol level and energy density from the same tank were averaged and treated as one replicate.

All data were examined for normality distribution and homogeneity of variance by the Kolmogorov–Smirnov test and the Levene's test. Differences in body wall mass, gonad mass and cortisol levels at the non-breeding stage and the mature stage were analyzed by Student’s t-test.

For normal data, changes in feeding activity, locomotor time, and parameters in the energy allocation model over time were analyzed by one-way ANOVA followed by Tukey's post hoc comparisons. For data with unequal variances, total body mass, R, and locomotor distance were analyzed by the Kruskal–Wallis test.

Body size has a significant effect on physiological traits in marine organisms. Thus, the correlation between body weight and routine metabolic rate was analyzed using the Pearson correlation analysis among all individuals. The routine metabolic rate at different breeding stages were also analyzed by one-way analysis of covariance with the least significant difference test with changing body mass as a covariate.

RESULTS

Body Growth and Gonad Development

The total body mass of A. japonicus showed a continuous increase during reproduction (Figure 1A; Kruskal–Wallis test: $H_4 = 71.557, P < 0.001$). The gonad mass reached 37.90 g over 100 days (Figure 1C; independent-samples t-test; $t_{32} = −19.301, P < 0.001$), suggesting that gonads were in the mature stage. It was found that the body wall mass at the mature stage was also significantly higher than that at the non-breeding stage (Figure 1B; $t_{32} = −13.007, P < 0.001$).

Feeding Activity

Feeding activity, including daily food consumption rate and ingestion rate, is shown in Table 1. Both daily food consumption (one-way ANOVA: $F_{3,12} = 40.111, P < 0.001$) and ingestion rate ($F_{3,12} = 63.033, P < 0.001$) decreased significantly during reproduction.

Routine Metabolic Rate and Cortisol Levels

Pearson correlation analysis indicated that the routine metabolic rate was positively correlated with the total body mass (Figure 2A; $r = 0.568, P < 0.001$). As expected, the routine metabolic rate gradually increased during reproduction (ANCOVA: $F_{4,25} = 8.261, P < 0.001$). However, the cortisol levels did not differ significantly between the non-breeding stage and the mature stage (Figure 2B; $t = −0.319, df = 12, P = 0.755$).

Locomotor Activity

Dynamic change in locomotor activity of sea cucumbers during reproduction was assessed by the total locomotor distance and the total locomotor time in 24 h. However, both locomotor distance (Figure 3A; Kruskal–Wallis test: $H_4 = 22.400, P < 0.001$) and locomotor time (Figure 3B; one-way ANOVA: $F_{4,25} = 31.726, P < 0.001$) showed continuous decline during reproduction.

Energy Intake and Allocation Patterns

Energy acquisition in females ranged from 11,253.74 to 6,656.89 J ind$^{−1}$ day$^{−1}$ during reproduction, showing a continuous downward trend (Figure 4A; one-way ANOVA: $F_{3,12} = 40.111, P < 0.001$). Increasing energy was discharged from the body as feces over gonadal growth (Figure 4B; $F_{3,12} = 18.536, P < 0.001$). Energy allocated to metabolic functions increased significantly during reproduction (Figure 4B; $F_{3,12} = 56.767, P < 0.001$). Energy allocated to reproduction increased significantly at different breeding stages (Figure 4B; Kruskal–Wallis test: $H_3 = 13.257, P = 0.004$). Energy allocated to locomotor function and somatic growth decreased significantly over time (Figure 4B; $F_{3,12} = 8.307, P = 0.003; F_{3,12} = 32.663, P < 0.001$).

DISCUSSION

Increasing Energy Demand and Decreased Appetite During Reproduction

Metabolic rate is a key indicator reflecting the state of energy expenditure in aquatic organisms (Huang et al., 2020; Pang et al., 2020). Compared to the non-breeding stage, the increase in sea cucumber metabolic rates during the breeding stage indicates that reproduction induces an increase in energy demand. Sea cucumbers, as ectotherms, are sensitive to both abiotic and biotic challenges. For example, variations in ambient temperature lead to an increase in metabolic rate (Dong et al., 2006). In addition, internal states, such as starvation and estivation, lead to a decrease in metabolic rate (Bao et al., 2010). In our study, adults were reared in suitable and stable conditions. Therefore, the growing gonad is the key factor inducing the increased metabolic rate in sea cucumbers. Similar results were also reported in fish, shellfish, and crustaceans (Boehlert et al., 1991; Baeza and Fernández, 2002; Kraffe et al., 2008). There are two reasons for increasing energy expenditure during reproduction. First, active egg growth requires high amounts of energy directly (from 228.83 to 1,063.72 J day$^{−1}$ at different breeding stages), and second the excretion of metabolic wastes (i.e., ammonia) from egg development through the maternal circulation also consumes energy indirectly (from 121.11 to 242.14 J day$^{−1}$ at different breeding stages).
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FIGURE 1 | Dynamic changes in total body mass (A), body wall mass (B) and gonad mass (C) of sea cucumbers during reproduction. Different letters indicate significant differences on different days (P < 0.05). Dotted lines represent 95% confidence intervals. Asterisk indicates a significant difference between the non-breeding stage (NBS) and the mature stage (MS). ***P < 0.001.

Sea cucumbers feed on sediments (Zhang et al., 2016). When food is easy to obtain, breeding animals may up-regulate their feeding activity to accommodate the increasing energy demand (Xu et al., 2012). Conversely, although females were provided a sufficient diet, their feeding activity showed a downward trend over gonad growth without interference from the courtship behavior of males (Marquet et al., 2018; Ru et al., 2018). This unexpected result suggests that reproduction suppresses the sea cucumber appetite. The contradictory phenomenon between increasing energy demand and suppressed feeding behavior has been widely reported for marine animals. Interestingly, many species such as octopus (*Amphioctopus fangia*), leatherback turtle (*Dermochelys coriacea*), and Atlantic salmon (*Salmo salar*) cease feeding completely during reproduction (Kadri et al., 1995; Plot et al., 2013; Tziouveli and Yokoyama, 2017). In our study, potential reasons for the loss of appetite in breeding females were not explored.

However, we found that there were no changes in cortisol levels between the non-breeding and breeding stages in sea cucumbers. Cortisol level is associated with the endocrine response to energy homeostasis in sea cucumbers (Pei et al., 2012). The unchanged cortisol levels during reproduction suggests that the increased energy demand does not cause a negative effect on maternal energy homeostasis, even though sea cucumbers lose their appetite.

Plasticity of Locomotor Function and Energy Homeostasis During Reproduction

Loss of locomotor ability during reproduction has been widely considered a cost of reproduction (Seigel et al., 1987). The physical burden hypothesis has been proposed to explain these observations (Miles et al., 2000; Shine, 2003). This hypothesis suggests that the growing egg or embryo presents an additional burden for females. Therefore, reproductive status causes inevitable inhibition of locomotor function. However, sea cucumbers have a fluid-filled body cavity, due to buoyancy, the growing eggs do not add to the physical burden of breeding sea cucumbers (Hamel et al., 2019). Furthermore, in order to avoid potential interference on locomotor activity from courtship behavior and thermoregulation (Fossette et al., 2012; Ru et al., 2017; Marquet et al., 2018), all individuals used in this study were females and the ambient temperature was maintained at 14°C. However, the locomotor activity of females still decreased gradually with egg growth.

Sea cucumbers do not possess a well-developed digestive system for physical chewing or chemical digestion of ingested...
food (Massin, 1982). As a result, most of the ingested energy from food was excreted out of the body as feces (Yuan et al., 2007). However, we showed that sea cucumbers allocate the highest proportion of assimilated energy, an average of 26.48–42.55%, to locomotor function at any reproductive stage. Feeding activity was continuously decreasing over gonad growth. However, both locomotion and reproduction (egg growth and metabolic function associated with reproduction) are energy-demanding functions. As a consequence of energy trade-off between reproduction and locomotion, the females gradually reduce unnecessary locomotory activity in order to conserve energy for reproductive success (Fossette et al., 2012). Thus, our results suggest that a decrease in locomotor function in breeding females might serve as a behavioral compensation for the decreased energy acquisition but increased energy demand during reproduction. There is also physiological evidence to support the hypothesis of behavioral energy conservation strategy. The energetic state of longitudinal muscle determines the locomotor activity in sea cucumbers, our recent study has reported that the supply of energy in longitudinal muscles decreases gradually during reproduction (Ru et al., 2017).
In fact, behavioral plasticity allows animals to adjust their behavioral function to both abiotic and biotic challenges. In particular, plasticity of locomotor behavior is a crucial capacity in sea cucumbers to cope with various challenges on different life history stages. For example, juvenile sea cucumbers can decrease their locomotor activity in order to maintain their survival when faced with strong water flow or intraspecific competition (Pei et al., 2014; Pan et al., 2015). In our study, the results showed that adults could decrease their locomotor activity in order to conserve energy for reproductive success and maintaining energy homeostasis. Similar results also reported in sea turtles. For example, juvenile sea turtles actively decrease their locomotor activity when they are satiated in order to conserve energy for somatic growth (Okuyama et al., 2013). In adults, in order to conserve energy for egg development, turtles also adjust their locomotor activity at low levels to conserve energy for maximum reproductive output (Fossette et al., 2012).

Lastly, this study showed high behavioral (i.e., locomotor activity) and physiological (i.e., routine metabolic rate) variations among individuals in the same environment during reproduction. As a result, sea cucumber gonad mass also showed individual variations within the population. Thus, these results indicate that behavioral and physiological polymorphisms might be the basis for differences of individual reproductive outputs in sea cucumbers. Functionally, the behavioral and physiological polymorphisms in marine invertebrates play an important role in dealing with the uncertainty of the complex marine environment (Dong and Zhang, 2016; Dong et al., 2017). Therefore, the significance of these polymorphisms in maternal traits of A. japonicus requires future studies.

**CONCLUSION**

In conclusion, we found that reproduction caused a significant increase in the energy demand in female sea cucumbers. However, the appetite of sea cucumber showed a downward trend over the course of gonad growth. Interestingly, the plasticity of locomotor behavior allows sea cucumbers to decrease locomotor activity in order to conserve energy, thereby accommodating for the increasing energy demand and maintaining energy homeostasis during reproduction. Our study provides new insights into how sea cucumbers accommodate reproductive demands by adopting a behavioral energy conservation strategy.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

**ETHICS STATEMENT**

All of the procedures in this study, including animal collection, animal rearing, and euthanasia, were performed under the Guidelines of Ethical Regulations of Animal Welfare of the IOCAS (IOCAS 2013.3).

**AUTHOR CONTRIBUTIONS**

XR performed the research. LZ and HY supervised this study. All authors contributed to the article and approved the submitted version.

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