Endocrine Control of Embryonic Diapause in the Australian Sharpnose Shark Rhizoprionodon taylori

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Abstract

The reproductive cycle of the Australian sharpnose shark, Rhizoprionodon taylori, includes a temporary suspension of development at the commencement of embryogenesis termed embryonic diapause. This study investigated levels of 17β-estradiol (E2), testosterone (T) and progesterone (P4) in plasma samples of mature wild female R. taylori captured throughout the reproductive cycle and correlated them with internal morphological changes. Levels of T were elevated through most of the embryonic diapause period, suggesting a role of this hormone in the maintenance of this condition. Increasing plasma T concentrations from late diapause to early active development were associated with a possible role of androgens in the termination of embryonic diapause. As in other elasmobranchs, a concomitant increase of E2 with ovarian follicle size indicated a direct role of this hormone in regulating vitellogenesis, while a peak in P4 suggested this hormone is associated with preovulation and ovulation. Additionally, significant correlations between photoperiod or water temperature and maximum follicular diameter and hepatosomatic index suggest that these abiotic factors may also play a role triggering and regulating the synchrony and timing of reproductive events.

Introduction

The ability to evolve and adapt reproductive strategies in response to changes in environmental conditions helps ensure the evolutionary success of any species [1]. As such, it has been postulated that environmental seasonality allows for the adjustment of reproductive events to occur over the most favourable period of the year maximizing survival and reproductive success [2–4]. By anticipating less favourable conditions, seasonal breeders are able to modify their morphology, physiology and/or behaviour [4]. In order to cope with unfavorable conditions and reduce energy waste, species have evolved several reproductive mechanisms (e.g. delayed fertilization, reduced gestation period), including embryonic diapause [2].

Embryonic diapause is the temporary ceasing or retardation of development during any stage of embryogenesis [3,6]. The phenomenon has been widely reported for vertebrates and invertebrates. Hypothetically, for live-bearing species delaying gestation allows parturition to occur when the environment is most likely to favour the survival of young [7,8]. Alternatively, it allows females to restore their energetic reserves after events of ovulation, mating, gestation and birth, without compromising the success of the following reproductive period [9].

In elasmobranchs, embryonic diapause is believed to have evolved independently in at least 16 species displaying three different types of viviparity [9]. This includes species where embryonic development relies solely on yolk (lecithotrophy) or additional nutrient input (matrotrophy) through uterine secretions (aplacental) or directly from the mother (placental) [10]. Aside from the occasional description of embryonic diapause in a number of elasmobranchs, there has been no systematic study of this phenomenon. Investigations have generally been limited to the observation of uterine eggs for extended periods (e.g. [7,11]). However, the physiological mechanisms controlling this process have never been investigated within this group of vertebrates. Furthermore, the selective pressures that have resulted in the evolution of embryonic diapause, as well as its benefits to the species, remain largely unknown (see review in [9]).

As in other vertebrates, reproductive events in elasmobranchs are associated with periodic cycles of steroid hormones. Therefore, understanding elasmobranch endocrinology is important for accurately delineating reproductive processes [12–15]. Testosterone (T), 17β-estradiol (E2) and progesterone (P4) are the main steroid hormones produced by the ovary [16] and are involved in controlling the reproductive cycle of female elasmobranchs. 17β-estradiol is known to play a role in vitellogenesis [17,18], while P4 can cause an antagonistic effect on E2 [19]. Other roles attributed to P4 are a regulation of ovulatory events [15,20,21], parturition [15,21,22], implantation and the inhibition of myometrial contractions [20,23]. The roles of T are less clear as it shows no distinct patterns among species and therefore it has been associated with several reproductive events (e.g. sperm storage, courtship, ovulation) [20,24,25]. Although previous investigations of these three key hormones have led to a better understanding of
the physiology of female elasmobranchs, their role in controlling embryonic diapause in this group remains unknown.

The control of embryonic diapause has been better studied in mammals [26] and reptiles [27]. Within viviparous reptiles, low temperature is necessary to arrest the late embryonic development, thus delaying parturition [28]. Although the endocrine control of this trait is not well understood in viviparous reptiles, injections of P4 have been shown to delay oviposition [29]. Similarly, concentrations of P4 are directly associated with the onset, maintenance and termination of this reproductive trait in mammals [6], in which P4 is required to maintain pregnancy and the suppression of its production induces diapause.

Though the control of embryonic diapause seems to be strongly linked to the mother’s endocrine functions, previous studies have shown it depends on a complex series of events. An environmental cue, usually photoperiod or temperature, is necessary to trigger the endocrine pathways that regulate the development of the corpora lutea [30], a temporary endocrine gland capable of synthesizing significant amounts of P4 [23,31]. In many cases the administration of P4 alone does not induce the termination of diapause. Investigations in mustelids and rodents have shown that a combination of P4 with some other unknown lutefugal protein(s) [32,33] or estrogen [34], respectively, is necessary to induce the termination of diapause. There is no evidence in the literature for the involvement of T in the control of embryonic diapause in other vertebrate groups.

In this context, the aim of this study was to investigate the role of reproductive hormones, and their correlation with abiotic factors, in the control of diapause and reproductive events in the Australian sharpnose shark Rhizoprionodon taylori. This species is a placental viviparous carcharhinid shark endemic to waters across northern Australia and southern Papua New Guinea [35]. Rhizoprionodon taylori is an annual reproducer with a gestation period of 11.5 months during which embryonic development is halted at blastodisc stage for a seven-month period (between February and September) of embryonic diapause [8]. This study describes the circulating steroid hormone levels of E2, P4 and T throughout the female reproductive cycle of R. taylori in relation to changes in gonadal morphology and environmental parameters. This is the first time that steroid hormone levels have been described in a diapausing elasmobranch species.

Materials and Methods

Sampling

All research activities were conducted under the permit requirements of the Great Barrier Reef Marine Park Authority - GBRMPA (G09/31573.1 and G10/33227.1) and the Queensland Department of Primary Industries and Fisheries – DPI (90911 and 144482). This study was conducted in compliance with the National Health and Medical Research Council (NHMRC) and Queensland Animal Care and Protection Ethics Committee. The protocol was approved by the James Cook University Animal Ethics Committee (permit No. A1508).

Females of Rhizoprionodon taylori were collected at night using 10 cm stretched mesh monofilament gillnets in Cleveland Bay, north Queensland (19°14’S, 146°48’E), monthly from February 2010 to February 2012. The nets were checked at 15-minute intervals. Shortly after capture, blood samples (3 ml) were collected from each specimen through caudal venipuncture using pre-heparinised syringes fitted with 22 gauge needles. Blood samples were preserved on ice for no longer than five hours and then centrifuged for five minutes at 1000 x g. The separated plasma was stored at -15°C until analysed for hormone levels through radioimmunoassay (RIA).

A total of 132 females were captured. Of these, 29 did not survive capture and were retained for examination of the reproductive organs. Blood samples were collected from the remaining 123 animals, of which 23 were released. All retained animals were euthanised by immersion in benzocaine bath for dissection. Specimens with vitellogenic follicles in the ovary, or embryos or ova in the uterus, were considered mature. Their total stretched length (STL; mm) was measured and total weight (g) and liver weight (g) obtained to calculate the hepatosomatic index (HSI = liver mass/body mass x 100). For the analysis of reproductive organs, ovaries were extracted, weighed (g) and the diameter of the largest follicle (MFD) measured to the nearest millimetre; the total number of ova or embryos in each uterus was counted, and total uterus weight (g) recorded, the ova or embryos weighed (g) and measured to the nearest millimetre. Females were classified into five reproductive stages (Table 1).

The presence of atretic vitellogenic follicles (AF) and postovulatory follicles (POF) were recorded. Macroscopic and microscopic examinations of ovarian follicles throughout the reproductive cycle were undertaken and correlated to existing literature of the chondrichthyan, especially elasmobranch, ovarian structures. Atresia can also occur in follicles prior to vitellogenesis, however, due to their morphological similarities with POF, only AF were used in the statistical analysis.

Environmental data for the study site were obtained from open online databases. Daily mean water temperature and air pressure data were obtained from the Australian Institute of Marine Science [36], and data on day length throughout the study period were obtained from U.S. Naval Observatory Astronomical Applications Department [37].

Radioimmunoassay

Plasma levels of P4, T and E2 were determined by RIA in 123 plasma samples. Plasma aliquots of 100 μl were extracted with ethyl acetate (1 ml) and 100 μl aliquots were then analysed. The antiserum and [1,2,6,7-3H] for E2 and P4 were purchased from Sigma-Aldrich (Australia). The E2 and P4 antiserum were reconstituted by adding 5 ml of Tris buffer (pH 8, 0.1 m HCl) and 50 μl of [1,2,6,7-3H] T were diluted in 5 ml of 100% ethanol and kept as stock for the assay. Duplicate standards (0–800 pg/tube authentic E2 and P4 in ethanol) and sample extracts were dried down and 200 μl of the reconstituted antiserum, diluted 1:10 in assay buffer (containing 0.1% of gelatin and 0.01% of Thimerosal), and 100 ul of the E2 and P4 stock, diluted 1:9 in assay buffer, added to each tube. Samples were placed in a bath at 37°C for an hour. Bound and free fractions were then separated using dextran-coated charcoal and aliquots of the supernatants counted in a Beckman LS 5801 liquid scintillation counter. All assays were validated by the evaluation of the slope of serially diluted extractions of plasma against the assay standards. Extraction efficiency was determined from recovery of 3H-labelled steroid added to pooled aliquots of plasma and assay values corrected accordingly. Extraction efficiency was 86% (E2) and 90% (P4). The detection limit for all assays was 0.02 ng.ml⁻¹. Intra and interassay variability was determined by including in each assay replicates of three levels of commercially available human control serum (CON 4, CON5, and CON 6 DPC). Interassay variability was 7% (E2) and 9% (P4), and intrassay variability less than 5%.

The T antiserum was Siroscera C-6050 (Bioquest) and [1,2,6,7-3H] T was purchased from Sigma-Aldrich (Australia). Duplicate standards (0–800 pg/tube authentic testosterone in
ethanol) and sample extracts were dried down and the assay protocol used was as described by Nicol et al. [38]. The assay was validated by the evaluation of the slope of serially diluted extractions of plasma against the assay standards. The detection limit for the assay was 0.02 ng.mL\(^{-1}\). Extraction efficiency was 90\%, interassay coefficient of variation was 11% and intrassay variability was less than 4%.

### Statistics

All statistical analyses were conducted using R (version 2.15.0) [39] with a critical probability level of 0.05. Hormone concentrations failed the Kolmogorov-Smirnov normality test and therefore nonparametric statistic tests were used. Differences in sample medians (steroid hormone, STL and HSI) over time and different stages were evaluated by Kruskal-Wallis one-way ANOVA on ranks followed by a pairwise Wilcoxon rank sum test with Bonferroni adjustment. The Spearman correlation [40] and ANOVA on ranks were used to determine the existence of associations between hormone levels, environmental parameters and morphometric parameters.

### Results

#### Reproductive cycle

Females ranging from 517 to 975 mm STL (683±4 mm; mean ± SE) were captured in all months of the year, except in June. Mean STL did not differ significantly between months (Kruskal Wallis, \(\chi^2[5] = 9.329, p = 0.10\)).

The average litter size per female was 5.3±0.2 embryos (range: 2–10); Soon after parturition (January), diapausing ova consisting of bright-yellow dense yolk masses concealed in brown egg cases measuring 23.9±0.3 mm (range: 21–29 mm) were observed in the uterus from mid February to mid September. Upon termination of diapause (stage 1), uterine ova became flaccid and slightly increased in size (29±0.2 mm, range: 25–32 mm). At this stage in September embryos from 15 to 17 mm were first visible. Embryos grew continuously to ~212 mm in January, when parturition occurs (Figure 1A).

The development of ovarian follicles commenced in late diapause (August). This initially slow process sped up in the two months prior to parturition (December to January; Figure 1B), when the largest embryonic size was recorded. Independent of the litter size, a single AF was present in the ovary throughout the diapausper period. The maximum size of these structures (26 mm) was observed in postovulatory females, after which it continuously reduced in size. The identification of AF at the end of the diapausper period (August – September) was difficult and sometimes not possible due to its small size, similar to POF (1–2 mm).

The HSI was significantly different between stages throughout the reproductive cycle, ranging from 1.2% in post partum females, to 9.6% in late diapause females (\(\chi^2[5] = 36.341, p<0.001\)) (Figure 2).

#### Serum steroid analysis

All reproductive hormones showed distinct cycles throughout the year (Figure 3) and no significant correlation was observed among them (P4 vs. E2: \(r_s[123] = -0.110, p = 0.23\); P4 vs. T: \(r_s[123] = -0.011, p = 0.90\); E2 vs. T: \(r_s[123] = 0.150, p = 0.10\)). Restricted sample sizes of different stages in January and February did not allow statistical determination of median differences amongst steroids at different stages over this period. However, differences in hormone levels between stages at each given month were examined when necessary.

17β-Estradiol

Plasma of E2 (1.42±0.13; range: 0.03–6.2 ng.mL\(^{-1}\)) varied significantly between stages through the reproductive cycle. Concentrations remained low through most of the diapausper period, starting to increase at the beginning of follicular development. Levels gradually increased to a peak at late stage 2 and post partum females before declining prior to ovulation (Figure 4). A Kruskal Wallis test revealed significant differences between E2 levels at different reproductive stages (\(\chi^2[5] = 46.32, p<0.01\)), with significant differences between levels of this hormone in early diapause and late diapauses (stage 2 and stage 3, respectively).

17β-estradiol levels were significantly correlated to ovarian structures. This association was positive with MFD (\(r_s[100] = 0.656, p<0.001\); Figure 5A) and negative when correlated with the size of AF observed during embryonic diapause, from February to October (\(r_s[34] = -0.479, p<0.001\); Figure 5B). Hence, it appears that E2 may be associated with the control of ovarian functions.

Progestosterone

Serum levels of P4 (0.900±0.082; range: 0.02–5.49 ng.mL\(^{-1}\)) in female R. taylori remained relatively low and were not significantly different between months (Figure 3). However, animals at different reproductive stages in January and February showed distinct differences in P4 concentration. The only stage 2 female captured in January showed a P4 level that was almost undetectable (0.02 ng.mL\(^{-1}\)), being much lower than the three post partum females (1.81±0.53, range: 0.97–2.80 ng.mL\(^{-1}\)). Progestosterone levels then increased in February to a peak (3.55 ng.mL\(^{-1}\)) in one ovulating female before a sharp decrease observed in six females in early diapause (0.43 ng.mL\(^{-1}\)±0.17, range: 0.02–0.85 ng.mL\(^{-1}\)). Although there was no significant correlation of this hormone with the female’s reproductive stage (\(\chi^2[5] = 9.329, p = 0.10\)), MFD

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**Table 1.** Classification of reproductive stages for female Rhizoprionodon taylori, where the numbers within the ovary column represent the range of maximum follicular diameter during each reproductive stage.

| Stage       | Ovary                                      | Uteri                              |
|-------------|--------------------------------------------|------------------------------------|
| Post partum | Vitellogenic follicles (10–14 mm)          | Empty                              |
| Ovulating   | Vitellogenic follicles (~12.0 mm)          | Ova half-way down the oviduct and/or uterus |
| Diapause    | Only previtellogenic follicles (1–3 mm)    | Presence of uterine ova            |
| 1           | Previtellogenic follicles (3–5 mm)         | Flaccid uterine ova (not attached to uterus) and visible embryo |
| 2           | Previtellogenic and vitellogenic follicles (2.5–13 mm) | Visible embryo attached to the uterus by a placenta |

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Testosterone

Plasma T levels (1.86 ± 0.17, range: 0.02–7.20 ng.ml⁻¹) varied widely throughout the year (Figure 3). In January and February, where sample sizes of different stages were too low to allow statistical comparisons some distinct variations were observed. In January, a rapid increase in T level was observed from one stage 2 female (0.33 ng.ml⁻¹) compared to three post partum females (2.01 ± 0.62; range: 0.81–2.84 ng.ml⁻¹), before returning to basal levels in February at ovulation and early diapause (0.60 ± 0.13; range: 0.06–1.13 ng.ml⁻¹). Therefore, it appears that T levels may undergo a small ephemeral post partum peak before declining at ovulation.

Testosterone varied significantly throughout different reproductive stages ($\chi^2[5] = 29.97, p<0.01$; Figure 4). Levels of this hormone were significantly different between late diapause and stages 2 and 3, nevertheless, it is important to note that high temporal variability of T concentrations during each given reproductive stage influenced this result. In fact, T concentrations remained elevated through most of the embryonic diapause period, after which two distinct peaks were observed (Figure 3). The first one, in September, revealed a significant difference in plasma T levels at different stages occurring in September ($\chi^2[2] = 13.9, p=0.01$). Significant differences in T in late diapausing females (1.09 ± 0.20; range: 0.39–2.93 ng.ml⁻¹, n = 11) compared to those at active development stages 1 and 2 (3.46 ± 0.33; range: 0.53–7.09 ng.ml⁻¹, n = 23) were observed (Figure 4). This result suggests a possible role for T in the termination of diapause. The second peak occurred at Stage 2, in November, and was not correlated to any observed morphological change. Testosterone levels showed a weak positive correlation to MFD ($r_{[100]} = 0.22, p=0.027$; Figure 5C) and negative to AF size ($r_{[95]} = -0.24, p=0.018$; Figure 5D), thus indicating that to some extent this hormone may be required in the regulation of ovarian functions.

Abiotic factors and reproduction

The water temperature in Cleveland Bay varied in close association with day length. A maximum seawater temperature of 30.1°C was recorded in February 2010 and January 2011 and after this peak, the temperature dropped gradually to a minimum of 20.8°C and 19.6°C during winter in July 2010 and June 2011 respectively. Day length at the study site latitude varied from 13.3 h in December to 10.9 h in June. Statistical analysis showed significant correlation between the environmental variables and E2 (day length: $r_{[123]} = 0.487, p<0.001$, Figure 6A; water temperature: $r_{[127]} = 0.360, p<0.001$, Figure 6B), but not with T or P4. Among the measured biological parameters, environmental parameters have also shown a significant correlation with HSI (day length: $r_{[96]} = -0.557, p<0.001$; and water temperature: $r_{[96]} = -0.664, p<0.001$) and MFD (day length: $r_{[100]} = 0.656, p<0.001$; and water temperature: $r_{[98]} = 0.539, p<0.001$).

Figure 1. Average monthly size (± SE) of (A) embryos and (B) maximum ovarian follicles in *Rhizoprionodon taylori*. Numbers in brackets denote sample size.
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Discussion
This is the first assessment of hormone levels of a diapausing elasmobranch. It provides a quantitative description of the main circulating reproductive hormones throughout the annual reproductive cycle. The amplitude of hormonal change in female *R. taylori* in association with reproductive events suggests some involvement of E2, T and P4 in the regulation of these events. Unlike mammals and reptiles, T and P4 (to a lesser degree) are probably involved in the control of embryonic diapause, but only T seems to be required at the release from embryonic diapause and reactivation of active development. Although E2 does not appear to have a direct role in controlling diapause, evidence suggests that photoperiod influences levels of this hormone, which could possibly be involved in the regulation and synchronization of reproductive events.

Late pregnancy and ovulation events
Although only small sample sizes were obtained from animals at the two stages occurring in both January (late stage 2 and post partum) and February (ovulating and diapause), the patterns of hormonal change were similar to previously studied elasmobranchs, allowing some assumptions to be made. Preovulatory peaks of E2 and T levels followed by increments in P4 concentrations at ovulation (when E2 and T levels sharply drop), observed in *R. taylori*, have been previously reported in a number of oviparous (*Raja erinacea*: [41,42]) and viviparous elasmobranchs (*Neyprion brevirostris*: [22]; *S. tiburo*: [20]; *S. acanthias*: [41]). A surge of T during preovulatory events is commonly linked with reproductive behaviour, ovulation and courtship events [13,20–22], while a peak in P4 is associated with the termination of the vitellogenic process. Therefore, it seems that in *R. taylori*, increased P4 and T levels are necessary to induce ovulation and inhibit further enlargement of vitellogenic follicles stimulated by E2.

The liver is known to be an important energetic reservoir, utilized in the development and maturation of large follicles and the nourishment of the developing embryos [43,44]. Commonly, mating and fertilization quickly follow parturition in annual breeders with concurrent ovarian and gestation cycles, such as *R. taylori* [8,45]. Nonetheless, all seasonally breeding elasmobranchs with consecutive ovarian cycles and gestation periods (*Carcharhinus isodon*: [46]; *Carcharhinus plumbeus*: [47]) require a ‘resting stage’, usually one year, to recover their hepatic reserves and produce large vitellogenic follicles in preparation for the following gestation [45,46].

This study demonstrates that levels of HSI in *R. taylori* steadily increased from minimum, at the early stages of pregnancy and diapause, to maximum at late diapause. Therefore, it seems that a diapausing period, as the resting stage, could allow females to restore energetic reserves in preparation for the subsequent ovarian cycle and embryonic nourishment, once development is restarted. Although more studies are required to confirm this theory, it seems that embryonic diapause allows females to restore their fitness between gestations without undergoing a biennial cycle.

![Figure 2. Hepatosomatic index variation throughout reproductive cycle in female *Rhizoprionodon taylori*. Values are medians ±SE. Letters depict statistical similarity using pairwise Wilcox post-hoc test. Missing letters represent groups that are too small for statistical analysis. Numbers in brackets denote sample size. doi:10.1371/journal.pone.0101234.g002](image-url)
Maintenance of diapause

Moderate levels of P4 in *R. taylori* throughout most of the diapausing period (except in May) indicate a possible role of this steroid in maintaining this phase of arrested development. Progesterone is closely related with vertebrate gestation and is generally required to allow embryonic development of mammals. In fact, where diapause occurs within this group, P4 production is inhibited during the period of arrested development, only allowing enough secretion to maintain the viability of the embryos [5,48]. However, this steroid hormone seems to have evolved different roles among elasmobranchs, as only a few species have been shown to elevate P4 levels through the gestation period (e.g. *Sphyrna tiburo*; [20]; *Squalus acanthias*; [49]). The role of P4 in *R. taylori* seems to be diametrically opposed to its role in mammals playing a role more related with the maintenance than the inhibition of diapause.

Nevertheless, it is also possible that P4 does not play a direct role in the inhibition of active embryonic development. Another role attributed to P4 in elasmobranchs is the inhibition of spontaneous uterine contractions, maintaining a quiescent uterus for recently formed embryos in *S. acanthias* [50]. Sorbera and Callard [50] suggest that uterine contractions allow the flushing of seawater into the uterine cavity, providing oxygenated water and the removal of waste materials. Assuming that, as in other vertebrates, diapausing embryos of *R. taylori* have very low metabolic rates [6,51], the exchange of water and removal of waste material is unnecessary and could potentially damage the delicate embryos. Therefore it seems possible that, as in *S. acanthias*, sustained moderate P4 levels in diapausing *R. taylori* could play a role in inhibiting uterine contractions, providing embryos with a quiescent uterus through this period.

Figure 3. Seasonal variation of 17β-estradiol (E2), progesterone (P4) and testosterone (T) in female *Rhizoprionodon taylori*. Letters depict statistical similarity using pairwise Wilcoxon post-hoc test. Progesterone levels were not significantly different throughout the year. The graph at the base of the figure indicates the timing of each reproductive event in females and numbers in brackets denote sample size.

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The major site of P4 synthesis in elasmobranchs has been postulated to be the CL, which can be formed after ovulation from the remnants of POF or from AF [52]. Histochemical studies have identified luteinizing tissues in POF [53] and AF [54], or both [55], in several elasmobranch species. The presence of both POF and AF persisting in the ovary through most of the diapausing period suggests these structures could be responsible for the observed sustained P4 levels during embryonic diapause. The lack of a significant correlation between AF size and levels of P4 in *R. taylori* could indicate that the synthesis capacity of this temporary gland is not directly associated with its size. Alternatively, this possibly indicates that the POF or other structures could potentially be more important for P4 production in this species. Studies in reptiles have demonstrated the capacity of P4 synthesis within the placental and ovarian tissues [56,57], suggesting that in *R. taylori* other sites of P4 production may exist.

Elevated T concentrations through most of the diapausing period (April to July) suggest this hormone may also function in the maintenance of embryonic diapause. Androgens have never been reported to have an involvement in the regulation of diapause of mammals or reptiles. Within non-diapausing elasmobranchs,
concentrations of T are generally low during pregnancy and do not seem to have a specific role during this period. The exception is *Dasyatis sabina* [21], where T was associated with changes in uterine events (embryonic nourishment). Histotroph secretion (uterine nourishing secretions) in *R. taylori* only starts at the release from embryonic diapause [58], therefore increased T levels during this period could suggest this androgen is required to maintain embryo viability while in the arrested development period.

Contrary to P4 and T, which seem to be associated with the regulation of some reproductive events, including embryonic diapause, the major role of E2 appears to be follicular growth and maturation. The present data demonstrates a strong linkage between increasing levels of E2 and vitellogenesis, which increase concomitantly from late diapause to a maximum in *post partum* animals. Similar patterns have been reported for several elasmobranchs studied to date [15,21,23,24,41], corroborating earlier *in vitro* studies where the E2 synthesis capacity by the elasmobranch ovarian follicles has been demonstrated (Fileti and Gallard, 1990 apud [59,60]). Once synthesised and secreted, E2 has been shown to induce the liver to produce vitellogenin, the main yolk precursor [16,41]. Thus, the strong correlation between E2 levels and MFD observed in this study reflects this vitellogenin production by the liver [41,43]. Therefore, it seems that this sex steroid is not directly involved in the regulation of embryonic diapause but, as in other elasmobranchs, it is essential for the control of vitellogenesis of *R. taylori*.

**Termination of diapause and active development**

Among the measured hormones, the termination of embryonic diapause and reactivation of embryonic development in *R. taylori* possibly only requires the presence of T. Unlike previous hypotheses that, as in other diapausing vertebrates, P4 would play an important role at the reactivation of normal development in elasmobranchs [9], the data in the present study do not show a peak in P4 prior to or at the release from embryonic diapause in September. Instead, a 3-fold increase in levels of T is observed in females from late diapause to stages 1 and 2, suggesting this hormone is possibly required for the reactivation of embryonic development and therefore terminating diapause. In addition, the new ovarian cycle starts as diapause is restarted, so ovarian follicles are very small at early embryonic stages until large (ready to ovulate) at parturition, supporting the idea that T may play a role in the reactivation of embryo development than triggering the initial vitellogenesis stages. Although the role that T plays in viviparous elasmobranch remains unclear, T has been associated with the later stages of follicle growth in oviparous elasmobranchs [42,61,62]. However, further studies are required to confirm whether these levels are specifically associated with embryonic development or other reproductive process, such as follicular development.

A further decline in P4 levels following the reactivation of active development supports the suggestion that this sex steroid could be playing an inhibitory role in embryonic development and/or regulation of uterine contractions. Testosterone, on the other hand, remains moderately elevated through the remaining part of gestation. A peak in November is not associated with any change in morphometric parameters, however previous observations by Simpfendorfer [58] suggest that the timing could be associated with the transition of embryonic nourishment (from histotroph to placenta). Testosterone has been previously associated with changes in embryonic nutrition in at least one elasmobranch. Embryos of *D. sabina* are nourished by yolk followed by uterine secretions and this transition is associated with a peak in T [21]. Thus, T could possibly play a role in the uterine events related to embryonic nutrition.

**Figure 4. Mean plasma steroid hormone concentrations at different reproductive stages of females *Rhizoprionodon taylori*.** Letters depict statistical dissimilarity using pairwise Wilcox post-hoc test. Missing letters in E2 and T graphs represent groups with sample sizes too small for statistical analysis. Progesterone levels were not significantly different among reproductive stages. Numbers in brackets denote sample size.

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Environmental factors

The present data indicates that abiotic factors may play a role as reproductive cues to synchronize reproductive events in *R. taylori*. Strong relationships were shown between the abiotic factors and MFD and HSI, suggesting that a certain level of dependency could exist between these variables and the changes in environmental factors. Additionally, the correlation of abiotic factors and E$_2$ supports the hypothesis that activating the synthesis of this hormone in this species possibly depends on environmental conditions.

Although the use of external factors as cues to regulate the timing of reproductive cycle is well known in other vertebrates

Figure 5. Relationship between plasma levels of hormones and ovarian follicles in *Rhizoprionodon taylori*. Changes in 17β-estradiol were significantly correlated with (A) maximum follicular diameter, and (B) atretic vitellogenic follicles observed from February to October. A weak correlation was observed between testosterone and (C) maximum follicular diameter, and (D) atretic vitellogenic follicles observed from February to October. Graphs show the correlation, including regression line and confidence interval of 95% represented by the shaded area.

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Figure 6. Seasonal changes in 17β-estradiol levels in *Rhizoprionodon taylori* were positively correlated with changes in environmental parameters: (A) day length and (B) water temperature. Graphs show the correlation, including regression line and confidence interval of 95% represented by the shaded area.

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[3,26], this strategy is not well documented in male and female elasmobranchs. While an association between abiotic factors and hormone levels has been demonstrated in at least three elasmobranchs (*Hemisquilla ocellata* [63]; *Urobatis hallieri*; [64,65]; *Scyliorhinus canicula*; [61]), controlled experiments demonstrating such dependence has only been shown in male *U. hallieri* [64], in which temperature has a role in elevating T levels.

Whether photoperiod or water temperature acts as reproductive cues controlling reproductive events (including embryonic diapause) in *R. taylori* and other elasmobranchs requires further investigation in controlled environments. Thus, it seems likely that highly synchronous populations, such as *R. taylori* in Cleveland Bay, require a fairly stable element to allow changes in reproductive stages to occur a short period apart among individuals.

**Conclusions**

This study provides the profile of hormones throughout the reproductive cycle of a diapausing elasmobranch and suggests that *R. taylori* may have evolved distinct endocrine pathways from other studied diapausing vertebrates. The data in the present study suggest the involvement of P4 and T in the control of embryonic development, while only T appears to be required for the reactivation of embryonic development after diapause. However, further research will be necessary to confirm the presence of androgen targets on the female and embryo, as well as the effects of this androgen in embryonic development. As in other elasmobranchs, levels of E2 in this species are closely associated with follicle development. This investigation provides support for the regulation of reproductive events by environmental variables. It provides indication that environmental factors such as day length and temperature may be playing a role in triggering events such as vitellogenesis and embryonic diapause in *R. taylori*.

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**Author Contributions**

Conceived and designed the experiments: DW CAA CAS. Performed the experiments: DW. Analyzed the data: DW. Contributed reagents/materials/analysis tools: DW SMJ CAA. Wrote the paper: DW.

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