The spermiation period in the rainbow trout (*Salmo gairdneri*). Plasma gonadotropin and androgen levels, sperm production and biochemical changes in the seminal fluid

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Summary. Plasma immunoreactive t-GTH and androgens, sperm production and biochemical changes in seminal fluid were studied during the spermiation period in rainbow trout. The volume of sperm collected by hand-stripping was very low (< 0.1 ml) at the onset of spermiation, increased slowly during the next 4 weeks and sharply thereafter. Plasma t-GTH was high (6 ng/ml) at the onset and then decreased; the sharp rise in sperm production started when circulating androgens had reached maximum values. In a second phase (6-12 weeks), the elevation in sperm production was strongly correlated with plasma t-GTH, while plasma androgens fluctuated. Spermatocrit did not vary significantly during the period studied so that increased sperm production corresponded to an elevation in spermatozoal production. During spermiation Na⁺ ionic concentration in the seminal fluid increased sharply and that of K⁺ only slightly. Total protein in the seminal fluid decreased significantly after 8 weeks.

Introduction.

There is little information in the literature on spermiation in fish, and especially on hormonal control as related to the biochemical composition and quantitative production of sperm. In the present work, we have studied variations of plasma gonadotropin (GTH) and androgens as well as differences in some seminal fluid parameters, such as spermatocrit and protein and mineral concentration, which could serve as sperm quality criteria during spermiation in rainbow trout.

Material and methods.

Experimental procedure. — Three-year old males weighing 875 ± 75 g were purchased from a commercial hatchery and kept in a recycled, bacteria-filtered water system from September until February under seasonal photoperiod and temperature ranging between 8 and 12 °C. After a 1-month acclimation to laboratory conditions, the animals were weighed and their blood sampled on 15 October. From this date,
the fishes were hand-stripped every week to detect the onset of spermiation. Once it was detected, blood and sperm samples were taken on anesthetized fishes (propoxate R7464, Janssen Pharmaceutica, 2 mg/I) every 2 weeks for a period of 12 weeks.

Measurements. — Plasma gonadotropin (GTH) levels were measured by a double specific radioimmunoassay (RIA) (Breton et al., 1975). Plasma androgen levels were determined by a double antibody RIA without chromatography and with a non-specific antibody (gift of Mr. Terqui), mainly binding testosterone (100 p. 100) and 11 Keto-testosterone (110 p. 100) but less androstene-dione (57 p. 100) and 5x-DHT (37 p. 100); adrenosterone, 17α-hydroxy-20α-dihydroprogesterone and 17β-estradiol were not bound (< 0.5 p. 100).

The volume of sperm released was determined by milt-stripping every time the animals were sampled. The spermatocrit was measured only when the volume of stripped sperm was at least 0.5 ml. Total protein concentration in the seminal fluid, obtained after 20 min. of centrifugation at 1 500 g, was determined using Lowry's

**FIG. 1.** — Changes in plasma immunoreactive t-GTH, total androgen levels and sperm production during the 12 weeks after the onset of spermiation in rainbow trout. Graphs show the mean value and standard deviation for each parameter. Difference between values are expressed as: N.S. : non-significant ; * : P < 0.1 ; ** : P < 0.05 ; ***: P < 0.02 ; **** : P < 0.01 ; (n) : number of fishes.
method modified by Hartree (1972); sodium and potassium were measured by flame spectrophotometry (Eppendorf model).

The statistical methods used were variance analysis and t-tests; the data were compared with Couple's method.

Results.

Plasma t-GTH and androgen levels (fig. 1).

Plasma GTH was high at the onset of spermiation (6.3 ng/ml), decreased during the following 6 weeks, then augmented between weeks 6 and 12, reaching a maximum value at 12 weeks (10 μg/ml). Androgen evolution was the reverse, increasing during 4 weeks from 34 to 104 ng/ml while t-GTH decreased. Total androgen levels then fluctuated, showing a slight decrease.

Sperm analysis.

The volume of milt produced between 2 samplings increased significantly during the period studied (fig. 1). Sperm production was very low at the onset of spermiation...
(0.1 ml) and rose steadily to reach a maximal value of 17.16 ml at 12 weeks. Spermato-
crit did not vary significantly during this period, fluctuating between 28.3 p. 100 at the
beginning and 25.4 p. 100 at the end (fig. 2). Spermatozoal production (fig. 2) there-
fore followed that of sperm (fig. 1).

Total protein. — Seminal fluid protein concentration was at its highest level
(1.74 to 1.89 mg/ml) at the onset of spermiation. It decreased significantly afterwards
and reached a concentration of 0.8 mg/ml at week 12 (fig. 2).

Sodium and potassium (fig. 3.) — Na+ concentration increased steadily from
1.400 mg/ml at 6 weeks, when first measured, to 2.034 mg/ml at 12 weeks. Variations
in K+ concentration during the same period were different; K+ increased from 784.6 mg
ml at 6 weeks to 1 156 at 8 weeks (P < 0.02), but decreased to 853.9 mg/ml at 10 weeks
(P < 0.01) to increase again at 12 weeks to 1 113.6 mg/ml (P < 0.02). The relation
Na+/K+ varied also ; its minimal value was 1.79 and its maximal value 2.36.

Discussion.

These data elucidate hormonal changes, sperm production and variation in semi-
nal fluid composition during the 12 weeks following the onset of spermiation. When
sperm release is first detected, low amounts of sperm are produced and high levels
of t-GTH are recorded in the plasma. Significant sperm production only occurred
several weeks later when t-GTH decreased and androgens had reached a maximum
level (fig. 1). At the onset of spermiation, the effect of GTH and androgen on sperm
production were not clear; GTH might be involved in the initiation of spermiation and the rise of circulating androgens which, in turn, could reduce the t-GTH secretion, thus suggesting a negative androgen feedback on t-GTH. Later, between weeks 6 and 12, sperm production was more closely correlated with the plasma GTH level. This agrees with data in the literature stating that gonadotropin is involved in sperm release (Clemens and Grant, 1965; Yamazaki and Donaldson, 1968). A high level of GTH has also been observed at the end of gametogenesis and during the spawning season in salmonids (Crim, Watts and Evans, 1975; Breton et al., unpublished data). Plasma androgens reach a maximum value at the same period (Schreck, Lackey and Hopwood, 1972; Idler, Horne and Sangalang, 1971). Androgens could also be involved in spermiation, and high levels may be required to stimulate sperm release (fig. 1). Whether only one specific active androgen is involved, or whether there are several, cannot be determined since the antibody used was not specific for any androgen. From the present data two stages are determined during spermiation, (1) initiation and preparatory stage lasting several weeks followed by (2) an active stage of sperm production when androgens are high and GTH is rising. Exogenous androgens alone stimulate spermatogenesis in vivo only when used in very large doses of 100 or 200 µg/g (Billard, 1974). This observation may also apply to spermiation. The spermatocrit stability already reported in this species (Chemayel, 1975) shows that stimulation of sperm production between weeks 6 and 12 is consequently a true stimulation of spermatozoal production.

The drop in seminal fluid protein concentration corresponds to a sharp increase of sperm production, and may suggest that protein synthesis capacity is limited and cannot deal with the copious seminal fluid secretion.

Ionic concentration of the seminal fluid is very high and increases mainly in Na⁺ during the period studied and to a lesser extent in K⁺. The rise in Na⁺ concentration is strongly correlated with elevated sperm production and plasma t-GTH, suggesting hormonal regulation.

Résumé. Une étude réalisée durant la période de spermiation chez la truite Arc-en-ciel a porté sur les taux plasmatiques de t-GTH et d’androgènes immunoréactifs sur la production de sperme et les changements dans la composition du liquide séminal. Le volume de sperme recueilli après massage abdominal est très faible au début (0,1 µl), augmente très légèrement au cours des quatre premières semaines et très fortement ensuite. Les teneurs plasmatiques en t-GTH sont élevées lorsque débute la spermiation et diminuent ensuite. La forte augmentation de production de sperme débute lorsque les teneurs en androgènes circulants ont atteint leur valeur maximum. Dans une seconde phase (6-12 semaines) l’augmentation de production de sperme est fortement corrélée avec les teneurs en t-GTH tandis que les niveaux d’androgènes fluctuent (fig. 1). La concentration du sperme en spermatozoïdes (spermatocrit) ne varie pas significativement pendant la période d’observation, de sorte que l’augmentation de production de sperme correspond à une augmentation de la production de spermatozoïdes (fig. 2). Pendant la période de
spermiation la teneur du plasma séminal augmente fortement dans le cas de Na⁺ et légèrement dans le cas de K⁺. Par contre la teneur en protéines totales diminue significativement après la 8e semaine.

References

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