ANDROGEN-BINDING PROTEIN IN EFFERENT DUCT FLUID OF RAT TESTIS

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Testicular fluid obtained from the rete testis and vasa efferentia has been reported to contain testosterone in concentrations approaching the amount in spermatic venous blood (Voglmayr, Waites & Setchell, 1966; White & Hudson, 1968). Flow rates, which have been established in several mammalian species (Setchell, 1970; Tuck, Setchell, Waites & Young, 1970) would indicate that the exocrine secretion may deliver a substantial quantity of androgen to the caput epididymidis. Testosterone and possibly other androgens in rete testis fluid might also exert a direct influence on spermatozoa as they are transported in this fluid.

In the present report, a specific androgen-binding protein with high affinity for dihydrotestosterone (DHT) and testosterone (T) is demonstrated in rete testis and efferent duct fluid from the rat testis. Fluid was collected after ligation of the vasa efferentia for 18 to 20 hr or for 3 days. The 18- to 20-hr ligations were made as close to the capsule of the testis as possible and rete testis fluid was collected from anaesthetized rats as described by Setchell & Waites (1971). The 3-day ligations were placed mid-way between testis and caput epididymidis. Animals were decapitated and the efferent ducts were dissected free of blood vessels before collection of fluid by opening the distended ducts. Spermatozoa were removed by centrifugation and the fluid was stored at −20°C. Both methods yielded 0·2 to 0·4 ml fluid/testis. Total protein concentrations were measured by the method of Lowry, Rosebrough, Farr & Randall (1951), using a bovine serum albumin standard. Values of the means and individual standard deviations were 2·4±0·6 mg/ml in several samples of rete testis fluid collected 18 to 20 hr after ligation and 8·4±0·4 mg/ml in efferent duct fluid collected 3 days after ligation.

Testicular fluid was diluted in 0·01 M-tris-HCl buffer, pH 7·5, containing 0·0015 M-EDTA, 0·002 M-2-mercaptoethanol, 10% glycerol and 0·5% bovine albumin (TEMGA-buffer). Aliquots of diluted fluid were equilibrated at 0°C for 2 hr with tritium-labelled steroids. Proteins were fractionated by polyacrylamide gel electrophoresis (PAGE) and the radioactivity measured in 2·4-mm slices as described by Ritzén, Nayfeh, French & Dobbins (1971). Text-figure 1 shows the single peak of bound [1,2-3H]dihydrotestosterone...
(\(^{3}\text{H}\)DHT) obtained when 0.15 ml fluid freshly collected from a single testis was diluted to 0.45 ml with TEMGA buffer and equilibrated with 0.1 \(\mu\)Ci (0.66 ng) \(^{3}\text{H}\)DHT for 2 hr at 0\(^{\circ}\)C. A similar aliquot of fluid was diluted in TEMG buffer (albumin omitted) and run in parallel for protein staining with amido black. Samples were layered directly over 6.5% gels and run at 0\(^{\circ}\)C in tris-glycine buffer (pH 8.9), 5 mA/tube for 3 hr.

No qualitative difference was noted in the peaks of bound radioactivity or patterns of protein bands obtained from fluids collected 18 to 20 hr and 3 days after efferent duct ligation. Albumin did not bind DHT and T in this PAGE system but stabilized the specific androgen-binding protein. Extraction of efferent duct fluid with charcoal (1 mg/mg protein) at 0\(^{\circ}\)C for 12 hr was shown to increase the binding of trace amounts of [1,2-\(^{3}\text{H}\)]T, suggesting the extraction of endogenously bound androgen.

The electrophoretic mobility of the binding component in testicular fluid was identical to that of the androgen-binding component previously described in a 105,000 \(g\) supernatant of rat epididymis homogenate (Ritzén et al., 1971). In both efferent duct fluid and epididymis supernatant, binding activity was destroyed by prior treatment with the proteolytic enzyme Protease Type VII (Sigma). Detailed comparisons of the physico-chemical properties of the high-affinity androgen-binding protein (ABP) in rat testis and epididymis indicate they are identical to the ABP in rete testis and efferent duct fluids (Ritzén, Dobbins, French & Nayfeh, 1972).

A quantitative estimation of binding sites was made (Text-fig. 2), utilizing the Sephadex gel equilibration assay of Pearlman & Crèpy (1967) modified as previously described by Ritzén et al. (1971). Rete testis fluid (2.0 ml), collected from four rats 18 to 20 hr after efferent duct ligation, was diluted
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1:3 in TEMGA buffer (containing 0.5% human albumin) and extracted with charcoal (1 mg/mg albumin) at 0°C for 12 hr. Charcoal was removed by centrifugation repeated several times. Sephadex G-25 (200 mg) was placed in glass-stoppered tubes with 1 ml TEMG buffer (albumin omitted) and allowed to swell overnight at 25°C. To each assay tube was added 0.25 ml diluted rete testis fluid, 0.25 ml TEMG-buffer containing [3H]DHT (0.04 μCi, 0.25 ng), and 0.50 ml TEMG buffer containing 0, 1, 3, 6 or 12 ng DHT. Tubes were shaken horizontally for 1 hr at 0°C and placed upright for 1 hr.

Radioactivity was counted in aliquots of the clear supernatants. Total bound steroid was plotted on the abscissa and the ratio of bound/free on the ordinate. Charcoal-extracted TEMGA-buffer was assayed separately and the corrected Scatchard plot of DHT-binding to the high-affinity sites was obtained graphically as described by Sandberg, Rosenthal, Schneider & Slaunwhite (1966). The concentration of ABP-binding sites in this rete testis fluid, 4.2 x 10^-8 M, was calculated from the x-axis intercept of the corrected Scatchard plot.

Binding affinities and concentrations of ABP-binding sites for DHT and T in charcoal-extracted efferent duct fluid were compared by PAGE and by Sephadex gel equilibration (Text-fig. 3). For PAGE, the fluid was diluted.
1:4-5 in TEMGA buffer (containing 0.5% bovine serum albumin) and extracted with charcoal as in Text-fig. 1. Aliquots of 0.5 ml were equilibrated with 0.8 µCi (5.28 ng) [3H]DHT or [3H]T and 10.72 ng DHT or T, amounts of the androgens which would completely saturate the binding sites, before fractionation (Text-fig. 3a). For Sephadex gel equilibration, the fluid was diluted 1:7 in TEMGA buffer (containing 0.5% bovine serum albumin). To each assay tube was added 0.25 ml efferent duct fluid, [3H]DHT or [3H]T (0.04 µCi, 0.25 ng) in 0.25 ml TEMG buffer and 0.50 ml TEMG buffer containing 0, 1, 3 or 6 ng of DHT or T. Charcoal-extracted TEMGA buffer was assayed separately and the individual values subtracted to obtain corrected values for the specific androgen-binding protein (Text-fig. 3b).

The relative amounts of DHT (2.44 ng) and T (1.06 ng) remaining bound after PAGE fractionation (Text-fig. 3a) closely resembled relative values of the equilibrium constants of association (Kₐ) for DHT and T as determined by Sephadex gel equilibration (Text-fig. 3b). The Kₐ for DHT was 2.5 times greater than the Kₐ for T. The similar concentrations of binding sites (Text-fig. 3b) and identical electrophoretic mobilities strongly suggested that DHT and T bind to the same site. Further evidence for a common site was obtained previously when it was demonstrated by competitive binding assay that [3H]DHT bound to ABP sites in epididymis supernatant could be displaced by T (Ritzén et al., 1971). A high degree of specificity for T and DHT was exhibited.
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by the ABP in this PAGE system. There was a relatively small affinity for [1,2-3H]5α-androstane-3α,17β-diol, but no binding of [1,2-3H]androstenedione, [1,2-3H]progesterone, or [2,4,6,7-3H]estradiol was noted.

The concentration of androgen-binding sites in fluid collected 3 days after efferent duct ligation was 8.5 x 10^-8 M which reflects the larger quantity of total protein in this fluid. The position of the ligature in these preparations probably permitted fluid to be absorbed in the efferent ducts, and the concentration of protein may indicate a preferential absorption of water in this region of the ducts. Resorption of fluid was also demonstrated by formation of sperm plugs in the ligated efferent ducts.

In studies reported elsewhere (French & Ritzén, 1972), it was demonstrated that the ABP is formed in the testis and may serve as a carrier of androgen from testis to epididymis. During its passage through the caput epididymidis, a major portion of the ABP either loses its binding activity or disappears from the soluble supernatant because of absorption from the lumen and uptake by an insoluble subcellular organelle of epididymal cells.

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REFERENCES