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UNUSUAL BEHAVIOUR OF FROG AND HUMAN SERUM PROTEINS LOADED WITH ¹³¹I-LABELLED THYROID HORMONES DURING PAPER ELECTROPHORESIS

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It has been reported earlier that the serum proteinbound iodine in the South African toad (Xenopus laevis) is higher than that of most animals." Since it is thought that the serum proteins to which thyroid hormones are bound, determine to some extent the speed with which the hormones enter the cells," a greater affinity of the specific binding proteins for thyroid hormones or a slower turnover rate of the binding proteins in cold-blooded animals could explain in part the relatively high serum proteinbound iodine and, therefore, the decreased metabolic rate in the frog.

During studies on the binding properties of serum proteins with ¹³¹I-labelled thyroid hormones, the association of 101I-labelled tri-iodothyronine (101Ta) and thyroxine (111Ta) with the proteins of frog serum was compared with that of human serum. The specific binding proteins, in most cases, were overloaded with 131T3 and 131T4. The hormones were labelled with 131 chemically by exchange³ and also endogenously by injecting 200µc of 131I into a mouse. After 48 hours the thyroid was digested enzymically. The hydrolysate was chromatographed, radio-autographed, and the radioactive bands corresponding to tri-iodothyronine and thyroxine markers were eluted (Fig. 1*). The "I-labelled thyroid hormones were added in vitro to serum and electrophoresed on paper in barbiturate-acetate buffer at pH 8.6.

The electrophoresed frog serum, unlike human serum, gave only 3 distinct bands on the anode side of the origin (Fig. 2). The albumin band of frog serum shows a greater electrophoretic mobility than that of human serum. Midway between the albumin and the origin, a broad band appeared in frog serum which corresponded to the area between the α , and the α_1 globulin of human serum. This protein band is referred to as the β -globulin, although it may not necessarily be the same as β -globulin of human serum. In some cases the y-globulin is fairly well defined, but in most cases it is just visible when 20µl. of frog serum is used.

The radioactivity from endogenously prepared ^{III}T₄ added to serum was associated with both the inter-alpha globulins and albumin when human serum was used. On the other hand, in frog serum it was mainly associated with β -globulin and to a lesser extent with albumin (Fig. 3).

In order to gain some information about the binding capacity of frog and human serum proteins, chemically prepared ¹⁸¹T, was added in known concentrations to serum, and electrophoresed. At a concentration of 0.05µg. "T₄/ml. of serum the bulk of radioactivity was again associated with the inter a-globulins in human serum and with the β -globulins in frog serum (Fig. 4). However, when the chemically prepared "T+ was increased to 0.5µg. "T+/ml. of serum, most of the radioactivity shifted onto the human serum albumin, whereas the albumin of frog

* Figs. 1 - 8 are on p. 1047.

serum had not taken up much 131T4. In frog serum such high concentrations of ^{III}T₄ were mainly associated with β -globulin and another radioactive peak appeared ahead of the albumin (Fig. 5).

When endogenously prepared 131T3 was added to frog and human sera, an unusual electrophoretic migration of the radioactivity was noted. With human serum the major portion of the radioactivity moved during electrophoresis for a distance of about 5 cm. from the origin towards the cathode, while smaller portions were associated mainly with albumin and pre-albumin. With frog serum the major portion of the radioactivity was similarly located as an intense band about 5 cm, on the cathode side of the origin whereas the rest of the activity coincided with the positions of albumin and β -globulin (Fig. 6).

Staining of the electrophoretograms showed no protein bands corresponding to the ¹³¹I-labelled substance which migrated towards the cathode. Electrophoresis of the same sera loaded with endogenously prepared ^{III}T₃ was repeated and the same results were obtained.

In an attempt to identify the radioactive substance which migrated towards the cathode, the human and frog sera containing the endogenously prepared ${}^{131}\mathrm{T}_3$ and ${}^{131}\mathrm{T}_4$ were chromatographed one-dimensionally in butanol: dioxan: 2-N NH₄OH (4:1:5) with carriers T₃ and T₄ (Fig. 7). The darkened areas on the radio-autograph of the chromatogram indicated that the 131Ts corresponded exactly with carrier T₁ in frog and human sera, but that the ¹³¹T₃ was slightly ahead of the carrier T₃ spots. The concentration of carrier T3 was greater than that of carrier T4, and this may have caused the 131T3 to move slightly ahead of carrier T3.

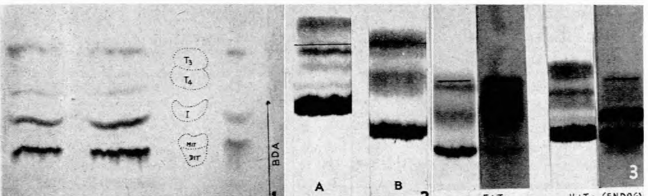
The section of the electrophoretogram corresponding to the ¹³¹I-labelled substance, which migrated towards the cathode, was cut out, eluted, and the extract chromatographed two-dimensionally in butanol:dioxan:2-N NH4OH and in butanol:acetic acid:water (120:30:50). Most of the activity corresponded to non-radioactive T₃ and iodide (Fig. 8). Other carriers tested out, like 3-mono-iodothyronine, 3:3'-di-iodothyronine, and 3:5:3'-tri-iodothyroacetic acid, did not coincide in two-dimensional chromatograms with the radioactive spots.

It is concluded that (1) frog serum, unlike human serum, carries thyroxine on the β -globulins, (2) the binding capacity of the β -globulins of frog serum for thyroid hormones is greater than that of the inter- α -globulins of human serum, and (3) the bulk of the 131I-labelled substance, which migrated towards the cathode during electrophoresis of frog and human sera under the conditions of the experiment, was in fact 132T3.

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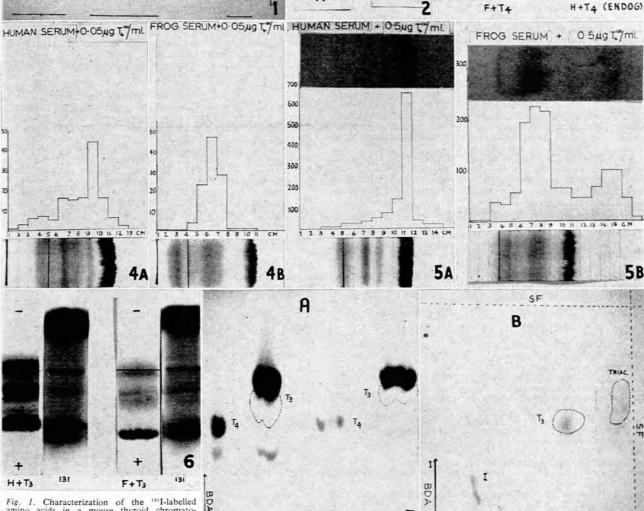


Fig. 1. Characterization of the ¹³¹I-labelled amino acids in a mouse thyroid chromato-graphed in butanol: dioxan:2-N NH₄OH. The dotted areas indicate the positions of the inactive carriers di-iodothyrosine (DIT), mono-iodothyrosine (MIT), iodide (I), thyroxine (T_4) and tri-iodothyronine (T_5).

Fig. 2. Stained electrophoretic patterns of (A) normal human serum, and (B) frog serum. Fig. 3. The association of endogenously prepared ¹³¹T₄ with frog (F+T₄) and human (H+T₄) sera. The radio-autograms are shown to the right of the stained electrophoretograms in each case. Fig. 4. The association of 0.05 μ g. ¹³¹T₄/ml. with (A) human serum, and (B) frog serum. The histograms represent the radioactive counts per cm.

C

D

BAW

B

A

length of the electrophoretograms.

Fig. 5. The association of 0.5 μ g. T₄/ml. with (A) human serum, and (B) frog serum. The radio-autogram (above) and the graph indicate qualitatively

and quantitatively the activity associated with the various protein fractions on the electrophoretogram below the graph. Fig. 6. The association of endogenously prepared ¹¹³T, with human (H) and frog (F) serum proteins. The stained electrophoretogram is indicated on the left of each pair while the radio-autogram is on the right. -Fig. 7. Radio-autogram of chromatographic analyses of frog (A and B) and human (C and D) sera to which endogenously prepared ¹¹³T, and ¹¹³T, had been added. Fig. 8. Characterization of the endogenously prepared radioactive substance which migrated to the cathode on electrophoresis, indicating the positions of non-radioactive carriers:iodide (I'), tri-iodothyronine (T_a) and tri-iodothyroacetic acid (TRIAC). Solvents: 1. Butanol:dioxan:2-N NH₄OH (BDA) 2. Butanol:acetic acid; water (BAW).

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