# Mechanisms of Adaptation to Iodine Deficiency in Rats: Thyroid Status Is Tissue Specific. Its Relevance for Man

Pablo Enrique Pedraza, Maria-Jesus Obregon, Hector Francisco Escobar-Morreale, Francisco Escobar del Rey, and Gabriella Morreale de Escobar

Endocrinología Molecular, Instituto de Investigaciones Biomédicas, Centro mixto Alberto Sols, Consejo Superior de Investigaciones Científicas and Facultad de Medicina, Universidad Autónoma de Madrid, 28029 Madrid, Spain

Many animals, man included, live in areas providing insufficient iodine (1) for optimal health. Degrees of I deficiency (ID) vary from mild-moderate to very severe, with quali- and quantitatively different negative consequences. To understand the mechanisms involved in adaptation to different grades of ID, we fed rats a low-iodine diet, plus additions resulting in a 250-fold range of I daily available to the thyroid, ranging from 5  $\mu$ g (adequate) down to 0.02  $\mu$ g I. We measured thyroid weight, total I, T<sub>4</sub>, T<sub>3</sub>, and type I 5' iodothyronine deiodinase (D1) activity, TSH, T<sub>4</sub>, free T<sub>4</sub>, and T<sub>3</sub> in plasma, T<sub>4</sub> and T<sub>3</sub> in 11 tissues, and two 5' deiodinase isoenzymes in four. TSH-independent thyroid autoregulation plays an important role in addition to TSH-dependent mechanisms in the adaptation to ID, avoiding a decrease of T<sub>3</sub> in plasma and most tissues,

**I** ODINE (I) IS a prerequisite for the synthesis of thyroid hormones,  $T_4$  and  $T_3$ . Although relatively abundant in sea water, it is often very scarce in terrestrial areas of the world, in which animals have developed a highly specialized structure, the thyroid follicle, to adapt to situations in which the supply of this element is inadequate to meet thyroid hormone requirements. The thyroid is optimized for the efficient use and storage of I in the form of I-containing compounds and for its intrathyroidal recycling, as a result of which severe systemic deficiency of  $T_3$  is prevented, even during relatively long periods of I deficiency (ID). A chronic and severe ID might, however, lead to a situation in which compensatory autoregulatory mechanisms can no longer avoid a decrease in  $T_4$  and  $T_3$  secretion, and the organism might suffer the ensuing negative consequences.

Today ID is still the single most important cause of preventable mental defects and cerebral palsy. It has been established that more than 1 billion people are living in conditions of chronic ID of varying degrees and are at risk of suffering some, or all, of the various ensuing ID disorders (IDDs) (1, 2). Their severity and irreversibility in man depends on both the degree of the ID and the period during development in which it is suffered. An adequate supply of

First Published Online February 2, 2006

despite a marked decrease of plasma  $T_4$ , whereas extrathyroidal responses of D2 mitigate  $T_3$  deficiency in tissues in which  $T_3$  is mostly generated from  $T_4$ . We focused on mild and moderate ID, the least investigated experimentally, despite its current frequency in industrialized countries. The novel and important finding of our study is that thyroid status cannot be defined for the animal as a whole: at all grades of ID,  $T_3$  is simultaneously elevated, normal, and low in different tissues. Present findings in mild-moderate ID draw attention to the importance, for man, of the resulting hypothyroxinemia that may affect mental functions and neurodevelopment of the inhabitants, even when they do not have the increased TSH or clinical hypothyroidism, often wrongly attributed to them. (*Endocrinology* 147: 2098–2108, 2006)

I to children and pregnant women are now considered basic human rights.

The epidemiological, clinical, and biochemical findings reported from different ID areas are quali- and quantitatively heterogeneous. Definition of different grades of ID (grade I, mild; grade II, moderate; grade III, severe) has been most helpful in clarifying apparently conflicting reports from different regions (3). Although the most severe irreversible IDD, such as the birth of neurological cretins, are usually found in areas with severe (grade III) ID (4), impaired mental functions are frequently found in the general population of even mild to moderate ID areas (5).

Most of our present understanding of the adaptation of human adults to chronic ID has been derived from experimental models that limited the I intake of rats. Such studies have usually involved comparison of results from animals on a low-iodine diet (LID) with those from rats receiving the same diet supplemented with enough I to meet physiological thyroid hormone requirements or eating the stock diet (*i.e.* Refs. 6–13). (For other related references, see supplemental data published on The Endocrine Society's Journals Online Web site at http://endo.endojournals.org.) It is often difficult to compare results because the nutritional composition and I content of the various LID diets differ or are often not even reported and so do the sex and strain of the animals. Few of these previous studies have involved graded degrees of ID, measured the concentrations of  $T_4$  and  $T_3$  in more than a few selected tissues, or evaluated some parameter of thyroid hormone action other than serum TSH. Despite this, such studies have permitted identification of many, but not all, of the intra- and extrathyroidal mechanisms involved in the

Abbreviations: BAT, Brown adipose tissue; BW, body weight; D1, type 1, 5' deiodinase; D2, type 2, 5' deiodinase; DTT, dithiothreitol; FT<sub>3</sub>, circulating free T<sub>3</sub>; FT<sub>4</sub>, circulating free T<sub>4</sub>; I, iodine; ID, I deficiency; IDD, ID disorders; LID, low I diet; LID', low I diet + KClO<sub>4</sub> (0.005%); PTU, 6-*N*-propyl-2-thiouracil; Tx, thyroidectomized.

*Endocrinology* is published monthly by The Endocrine Society (http:// www.endo-society.org), the foremost professional society serving the endocrine community.

Endocrinology, May 2006, 147(5):2098-2108 2099

response to ID, most of which cannot be investigated directly in man for obvious constraints.

We want to point out that the experimental design used here aimed at a difference in the amount of I available to the thyroid as the single controlled variable between groups. It should be an adequate model for inhabitants of areas in which ID is the sole cause of goiter and thyroid tissue is functionally unaffected, even in the neurological cretin. It is not, however, an appropriate model for those ID areas in which other nutritional or environmental factors may lead to thyroid necrosis, primary hypothyroidism, and myxedematous cretinism, such as seen in central Africa (14).

The aim of the present study was to assess the relative roles of intra- an extrathyroidal mechanisms in the response of individual tissues to different grades of chronic ID, from mild to moderate, to severe and very severe, with special focus on the concentrations of  $T_3$ . These were measured in a larger number of tissues than previously studied over such a wide range of ID. Although the  $T_3$  concentration was taken as an index of possible thyroid hormone effectiveness at the individual tissue level (15), some biological end points of thyroid hormone action, other than circulating TSH, were also measured in selected tissues. The present approach shows that the thyroid status of ID rats cannot be defined for the animal as a whole because it is eminently tissue specific: at all grades of ID, elevated, normal, and low concentrations of  $T_3$  are simultaneously found in different tissues of the same animal.

# **Materials and Methods**

#### Experimental design

Young adult female Wistar rats were used. They were housed under humane conditions, with alternating 12-h light, 12-h dark cycles and at 22–24 C, under veterinary control according to European Community guidelines and after approval by the ethics committee of our institution. They were fed a stock pelleted diet for rats (Sandermus; Sanders, S.A., Barcelona, Spain) with an I content varying between 0.15 and 0.35  $\mu$ g I/g, which would provide 3–7  $\mu$ g I per day per rat (assuming a daily food intake of 20 g). When they were approximately 50 d of age and weighed 120–150 g, they were switched from the stock diet to a diet with a very low iodine content (LID) and given 1% KClO<sub>4</sub> as drinking water for 1 wk to block further uptake of circulating iodide and accelerate depletion of thyroid stores and I-containing compounds (16). This treatment minimized previous differences between animals. From then on, animals were separated into different groups of five to six animals each and fed the different diets described for experiments A and B.

Although we prepare the LID with components obtained from the same commercial sources, its final I content is variable over time. To ensure that at least one experimental group is severely ID, we often include animals fed LID supplemented with very low amounts of KCIO<sub>4</sub>, at a concentration of 0.005% (LID' group) (17, 18). This concentration is 200 times lower that that used during the initial week before separating the animals into different experimental groups. The purpose is to decrease the availability to the thyroid of the small amounts of I present in the LID and generated during intra-thyroid I recycling.

The LID was prepared by mixing thoroughly 6 kg corn flour (Productos Hercosa, Barcelona, Spain), 2.5 kg wheat gluten (Herman Kröner GMBH, Ibbenbürn, Germany), 1 kg brewer's yeast (Vitalevor, Strasbourg-Neuhof, France), 0.15 kg NaCl, and 0.15 kg CaCO<sub>3</sub> (Carlo Erba, Milano, Italy). Sufficient amounts were prepared to cover a complete experiment with the same batch. Daily rations were prepared by mixing the dry LID with distilled water containing the different additions (KClO<sub>4</sub> or KI solutions) in a proportion of 0.7:1.0 (water-dry LID) and dividing it into individual aliquots, which were kept frozen until use.

*Experiment A.* A preliminary experiment was performed to investigate whether these small amounts of  $KClO_4$  (0.005%) would exert *per se* 

effects on thyroid hormone economy other than impairing the uptake of the minute amounts of I contained in the LID and intrathyroid I recycling. For this, two groups of six animals each were fed LID (LID groups) and two other groups LID supplemented with KI (LID + I groups). One each of these two groups received LID + 0.005% KClO<sub>4</sub>: LID' and LID' + I groups, respectively. The amount of KI was such that a daily intake of 20 g of the LID + I mixture would provide 10  $\mu$ g I per animal.

*Experiment B.* This comprised five groups of five animals each that drank distilled water and were fed the basic LID diet with the additions specified in Table 1.

The I content of the LID diet itself was below the limit of detection of the analytical procedure used to determine I in food (9). We measured the 24-h urinary I excretion of LID and LID' rats during 5 consecutive days. Mean values ( $\pm$  SEM) were 0.052  $\pm$  0.012 and 0.083  $\pm$  0.009  $\mu$ g, respectively. Although these results indicated that the amount of I ingested daily by animals on LID or LID' was 0.052  $\pm$  0.012  $\mu$ g, the amount actually available to the thyroid of the LID' animals was 0.031  $\mu$ g less, namely 0.021  $\mu$ g I. Table 1 shows the estimated I intakes of the different groups; they are merely tentative because neither the amount of food ingested individually nor the fecal excretion of I-containing compounds was measured. The amount of I available to the thyroids of the different groups of rats appeared to differ 100-fold or more between the two extreme groups (LID' *vs.* LID + 5.0).

After 3 months on these different diets, the rats were slightly anesthetized with ether, bled extensively, and perfused (15). Samples of plasma, interscapular pads of brown adipose tissue (BAT), pituitary, brain, cerebral cortex, cerebellum, liver, kidney, heart, lung, adrenals, ovaries, and muscle (musculus quadriceps femoris) were dissected out and frozen for the determination of  $T_4$  and  $T_3$ . Aliquots of cerebral cortex, BAT, liver, and lung were kept frozen at -80 C for the determination of iodothyronine deiodinase activities.

The thyroid was dissected out, weighed, divided into three aliquots, and kept frozen for the determination of type I 5' deiodinase (D1) activity and total I,  $T_4$ , and  $T_3$  contents.

#### Determinations

*I content*. This was determined in aliquots of thyroid glands, urine, or LID by modifications of a chloric acid digestion procedure (16).

 $T_4$  and  $T_3$  in plasma and extrathyroidal tissues. Total  $T_4$  and  $T_3$  were measured by specific and highly sensitive RIAs, after extraction with methanol and extensive purification of the iodothyronines, as detailed elsewhere (15, 19).  $T_4$  and  $T_3$  concentrations in a given type of sample from the five experimental groups were determined in the same extraction run and in a single RIA for each hormone. To increase recovery of very small tissues (*i.e.* pituitary, adrenals, ovaries), the initial methanol extract was purified directly through the resin columns, omitting the methanol chloroform extraction and back-extraction procedure.

The plasma free  $T_4$  (FT<sub>4</sub>) was calculated from the total  $T_4$  concentration and the percentage of added tracer amounts of [<sup>125</sup>I]T<sub>4</sub> that was not bound to serum transport proteins. The latter was determined by ultracentrifugation of undiluted samples through Centricon-10 microconcentrators (Amicon GmbH, Witten, Germany) as detailed (20).

 $T_4$  and  $T_3$  in the thyroid. The contents of  $T_4$  and  $T_3$  in the thyroid were measured separately in two different fractions to which we refer here, respectively, as the "Free"  $T_4$  and "Free"  $T_3$  pools and as the "Total"  $T_4$  and  $T_3$  pools. When applied to the thyroid, the adjectives free and total have a different meaning from plasma. In the thyroid, "Free"  $T_4$  and

**TABLE 1.** Estimation of the amounts of iodine available daily to the thyroid of the rats from the different experiment B groups

Group	Diet + supplements	I, $\mu$ g/d per rat <sup>a</sup>		
LID'	LID'	$0.021^{b}$		
LID	LID	$0.052^b$		
LID + 0.5	$LID + 0.5 \ \mu g I/20 g$	0.50		
LID + 1.0	LID + 1.0 $\mu$ g I/20 g	1.0		
LID + 5.0  (controls)	LID + 5.0 $\mu$ g I/20 g	5.0		

<sup>*a*</sup> Theoretically available for thyroidal uptake.

<sup>b</sup> Based on 24-h urinary I excretion data.

"Free"  $T_3$  correspond to the iodothyronines present in the gland as amino acids, no longer incorporated into proteins by peptidic bonds, and presumably available for secretion as hormones into the bloodstream. "Total"  $T_4$  and  $T_3$  correspond to the iodothyronines residues still incorporated by peptidic bonds into thyroglobulin and other proteins. The concentrations of "Free"  $T_4$  and "Free"  $T_3$  were obtained using the methanol extracts of the thyroid homogenates, then processed as other tissue extracts. The thyroid pools of "Total"  $T_4$  and  $T_3$  were measured in methanol extracts of proteolytic hydrolysates of the pellets remaining after the initial methanol extraction, as described (21)

*lodothyronine deiodinases.* D1 activity was assayed in liver and lung homogenates using 400 nm rT<sub>3</sub> and 2 nm dithiothreitol (DTT) for liver and 2 nm rT<sub>3</sub> and 20 mm DTT for lung in 100 mm potassium phosphate buffer (pH 7.0) and 1 mm EDTA. The reaction time was 10 min for liver and 60 min for lung. D1 activity was also assayed in an aliquot of the thyroid gland, using 0.8–1.0  $\mu$ m rT<sub>3</sub> and 2 mm DTT for 10 min.

Type II 5'-iodothyronine deiodinase (D2) activity was assayed in the cerebral cortex and BAT using 2 nm  $T_4$  + 1  $\mu$ m  $T_3$  and 20 mm DTT in the presence of 1 mm PTU, and the reaction time was 60 min.

All samples were homogenized in buffer containing 0.32 M sucrose, 10 mM HEPES (pH 7), and 1 (for D1) or 10 mM (for D2) DTT. Before each assay [ $^{125}I$ ]rT<sub>3</sub> or [ $^{125}I$ ]T<sub>4</sub> was purified, and the assays were performed as described (22). The protein content was usually 150–200  $\mu$ g/tube for most tissues but was 10-fold lower when liver or thyroid were assayed.

*Circulating TSH.* TSH was measured in plasma using immunoreactants kindly provided by the Rat Pituitary Agency of the National Institute of Diabetes, Digestive and Kidney Diseases (National Institutes of Health, Bethesda, MD) as described elsewhere (16). Results are expressed in weight equivalents of the National Institute of Diabetes and Digestive and Kidney Diseases rTSH RP-3 preparation.

#### Drugs and reagents

 $T_4$ ,  $T_3$ , 3,5-diiodothyronine, 6-*N*-propyl-2-tiouracil, and DTT were obtained from Sigma Chemical Co. (St. Louis, MO).  $rT_3$  and 3',3-diiodothyronine were obtained from Henning Berlin GMBH (Berlin, Germany).

High specific activity  $[^{131}I]T_4$ ,  $[^{125}I]T_3$ ,  $[^{125}I]T_4$ , and  $[^{125}I]rT_3$  (3000  $\mu$ Ci/ $\mu$ g) were synthesized in our laboratory, as previously described (19) and used for highly sensitive  $T_4$  and  $T_3$  RIAs, as recovery tracers for plasma and tissues extractions, for the determination of plasma FT<sub>4</sub>, and as substrates for D1 and D2.

#### Statistical analysis

One-way ANOVA and the protected least significant difference *post hoc* test were used for multiple comparisons after validation of the homogeneity of variances by the Bartlett-Box F test. Square root or logarithmic transformations usually ensured homogeneity of variances when this was not found with the raw data. Results are expressed as means  $\pm$  sEM. *P*  $\leq$  0.05 was considered significant in all comparisons. Whenever it is stated that a difference was found between groups, it implies that it is statistically significant. To be able to compare the degree of changes in different parameters, all present results are expressed as percentages of the mean value obtained for the LID + 5.0 group (also referred to as control, C, group), used as 100%. Multiple regressions and partial correlation analyses between different parameters were also performed. All calculations were done using the SPSS for Macintosh (version 6.1.1; SPSS Inc., Chicago, IL).

#### Results

# Experiment A

 $T_4$ ,  $T_3$ , and TSH concentrations in plasma as well as the concentrations of  $T_4$  and  $T_3$  in liver, brain, BAT, kidney, lung, heart and skeletal muscle, and D2 activities in the cerebral cortex were measured for the four groups. No differences were found between the LID + I *vs.* LID'+I groups. (For detailed data, see table in supplemental data published on The Endocrine Society's Journals Online Web site at http://

endo.endojournals.org). The values found in the LID and LID' groups were significantly different from those of their respective I-supplemented groups, LID + I and LID' + I. There were also significant differences between values found in LID', compared with LID animals: plasma  $T_3$  and TSH, liver, brain, kidney, and muscle  $T_4$  and brain, lung, heart, and muscle  $T_3$ . Present results confirm that these very small amounts of KClO<sub>4</sub> do not have any detectable effect on thyroid hormone status when the I intake is adequate. On the contrary, when the intake is very low, as in rats on LID, thyroid hormone status is affected by the addition of 0.005% of KClO<sub>4</sub>. The changes observed in LID' rats *vs*. those on LID are those that would be expected from a greater degree of ID.

# Experiment B

*Body weight.* During the 3 months of treatments, the animals on KI-supplemented LID, namely the LID + 0.5, LID + 1.0, and LID + 5.0 groups, increased similarly in body weight (BW) by  $64.4 \pm 2.0$ ,  $60.80 \pm 2.3$ , and  $63.4 \pm 8.9$  g, respectively. The BW of the LID and LID' groups increased significantly less, by  $51.5 \pm 2.0$  and  $36.2 \pm 2.7$  g, respectively; the difference between the latter two groups was also significant.

*Thyroid gland.* Figure 1 shows the results corresponding to the thyroids of the five groups of experiment B. In this and successive figures, the results have been normalized by expressing them as percentages of the mean values for the C group, which are detailed in the figure legend.

All values shown in Fig. 1 for the LID + 1.0 group were already different from those of C animals including the weight of the gland. The exceptions were the "Total" T<sub>3</sub> to "Total" T<sub>4</sub> ratio and the activity of the D1 isoenzyme, which, however, were different from C animals in LID + 0.5 rats. The weight of the thyroid increased up to 2.5-fold in the LID' rats, with a 2-fold increase being observed already in the LID + 1.0 group. The total I content of the gland decreased with I intakes to an LID' value that was 5% of that of the C animals. The "Total" T<sub>4</sub> and "Free" T<sub>4</sub> also decreased markedly and to a similar degree, reaching values of 1.5 and 1.8% of the C value, respectively, in LID' rats. In contrast, the "Total" T<sub>3</sub> and "Free" T<sub>3</sub> followed different patterns of change from those observed for  $T_4$ . After a marked decrease in LID + 1.0, compared with C animals, the "Total" T<sub>3</sub> and "Free" T<sub>3</sub> remained relatively constant despite a marked decrease in I content. In LID' rats, "Total" T<sub>3</sub> and "Free" T<sub>3</sub> were, respectively, 8.5 and 23% of C value, much higher than those for "Total" T<sub>4</sub> and "Free" T<sub>4</sub>.

The ratios of both "Total"  $T_3$  and "Free"  $T_3$  to  $T_4$  increased with decreasing I availability. The increase was, however, much more marked (17-fold) for the ratio of "Free"  $T_3$  to  $T_4$ than that found for the ratio between "Total"  $T_3$  to  $T_4$  (7.5fold). This discrepancy may, at least partly, be accounted for by the increase in D1 activity in the gland that could generate "Free"  $T_3$  from the "Free"  $T_4$  but would not affect the iodothyronine residues still incorporated by peptidic bonds into thyroglobulin and other proteins.

# Circulating $T_{4}$ , $FT_{4}$ , $T_{3}$ , and TSH

The changes occurring in the plasma of the different groups of rats as I availability decreases are shown in Fig. 2.

FIG. 1. Changes in thyroid weight, total iodine, T<sub>4</sub>, and T<sub>3</sub> contents as well as T<sub>3</sub> to T<sub>4</sub> ratios and total D1 activities in groups of rats with a decreasing iodine intake. "Total" T<sub>4</sub> and T<sub>3</sub> correspond to residues incorporated into thyroglobulin and other proteins by peptidic bonds, whereas "Free" T<sub>4</sub> and "Free" T<sub>3</sub> are those present in the gland as iodoamino acids, available for secretion as hormones into the bloodstream. All values are normalized by taking as 100% the mean value of the corresponding variable in the controls (LID + 5.0 group of animals); weight:  $23 \pm 1.2$  mg; total I: 3.19  $\pm$  0.85  $\mu g/gland;$  "Total"  $T_4$ and T<sub>3</sub>: 4053  $\pm$  1297 and 212  $\pm$  47 ng/ gland; "Free" T<sub>4</sub> and "Free" T<sub>3</sub>: 139  $\pm$  46 and  $15.7 \pm 4.3$  ng/gland; "Total" T<sub>3</sub> to "Total"  $T_4$  ratio: 0.045  $\pm$  0.007; "Free"  $T_3$  to "Free"  $T_4$  ratio: 0.111  $\pm$  0.026; and D1:  $1288 \pm 175$  pmol/min per gland. Arrows identify statistically significant increases or decreases vs. the C group; & and & identify statistically significant differences between the mean values for LID' and LID animals. The black arrows and bold & correspond to the variables shown as *black bars*, the *white arrows* and & to those represented by white or striped bars.



Plasma  $T_4$  is lower in LID + 0.5 rats, compared with controls (LID + 5.0 group), but the difference was not statistically significant for the LID + 1.0 animals. It further decreases with increasing ID, reaching values in LID' animals that were only 5% of C levels. FT<sub>4</sub> is already decreased in the LID + 1.0 rats, compared with controls. The pattern of changes in circulating  $T_3$  is quite different from that described for  $T_4$ . When the I intake decreases from LID + 5.0 to LID + 1.0, circulating  $T_3$  actually increases and then de-



FIG. 2. Changes in plasma  $T_4,\ FT_4,\ and\ T_3,\ in$  the plasma  $T_3$  to plasma  $T_4$  ratio and circulating TSH in groups of rats with decreasing I availability. All values are normalized by taking as 100% the mean value of the corresponding variable in the controls (LID + 5.0 group of animals;  $T_4$ : 22.9  $\pm$  3.9 ng/ml; FT\_4: 29.6  $\pm$  0.4 pg/ml;  $T_3$ : 0.31  $\pm$  0.02 ng/ml;  $T_3$  to  $T_4$ : 0.0135  $\pm$  0.0015; TSH: 1.25  $\pm$  0.16 ng rTSH RP-3/ml. The meaning of symbols is the same as in Fig. 1.

creases again to C values in the LID + 0.5 and LID groups. Only in the LID' animals did plasma  $T_3$  decrease, to 45% of C values, a change that contrasts with the much more pronounced decrease observed for circulating  $T_4$ . As a consequence, the  $T_3$  to  $T_4$  ratio is already increased above C values in the LID + 1.0 animals and further increased, more than 10-fold, in LID' rats.

The mean circulating TSH value was higher in the LID + 1.0 animals, compared with C values (1.69  $\pm$  0.27 vs. 1.25  $\pm$ 0.16 ng rTSH RP-3/ml, respectively), but the difference did not reach statistical significance (P = 0.198) with present sample sizes. It then increased progressively, 10-fold in LID', compared with C animals, with a 7-fold increase in the LID + 0.5 group. Plasma TSH was negatively correlated to T<sub>4</sub> (P =0.0012) but not to  $T_3$ . Partial correlation analysis disclosed that the relation between circulating TSH and T<sub>4</sub> was independent from T<sub>3</sub> levels. Plasma TSH, however, was negatively correlated to both pituitary  $T_4$  and  $T_3$  (P < 0.000 for both). Moreover, the degree of increase in TSH was unexpectedly small, even in the LID' animals, compared with that observed in rats with a similar degree of hypothyroxinemia caused by primary thyroid failure, when T<sub>4</sub> and T<sub>3</sub> decrease concomitantly, and circulating TSH is negatively and independently correlated to both plasma T<sub>4</sub> and T<sub>3</sub>. This is illustrated in Fig. 3, in which the changes in plasma TSH of the present animals are compared with those of sex- and weight-



FIG. 3. A, Relationships between plasma TSH and  $T_4$  in rats under two different experimental conditions, namely experiment B and those from a model of primary thyroid failure, namely Tx age- and weight-paired rats infused with different doses of  $T_4$  (15). B, Results on a log scale for TSH. *Symbols in black* identify groups with circulating  $T_3$  that was decreased with respect to the corresponding C value: the LID + 5.0 group for the LID experiment B and intact rats for the Tx +  $T_4$  experiment.

paired animals that had been thyroidectomized (Tx) and infused with different doses of  $T_4$  (15). The latter animals showed plasma  $T_3$  changes parallel to those of  $T_4$ , whereas in the LID animals,  $T_3$  was still normal with plasma  $T_4$  and  $FT_4$  levels that decreased to 25 and 15%, respectively, of C values. Figure 3B shows that the plasma TSH increase is blunted, more than 10-fold, in the LID animals, compared with that observed in the Tx rats on  $T_4$ .

Extrathyroidal tissues. Figure 4 illustrates the changes of the concentrations of T<sub>4</sub> and T<sub>3</sub> in the liver, lung, brain, and BAT occurring with decreasing I availability as well as the D1 activities in liver and lung and the D2 activities in brain and BAT. In these tissues the concentrations of T<sub>4</sub> decreased following a pattern similar to that observed for plasma T<sub>4</sub> or FT<sub>4</sub>. The concentrations of T<sub>3</sub> in liver and BAT followed patterns similar to those of circulating T<sub>3</sub>, with an initial increase from the LID + 5.0 to LID + 1.0 animals, normal concentrations in the LID + 0.5 and LID groups, and lowerthan-normal levels in the LID' rats. In the lung the increase in T<sub>3</sub> concentrations persisted in all groups, including the severely iodine-deficient LID' animals that had a decreased plasma  $T_3$ . The pattern of changes for  $T_3$  in the brain were different from those in liver, lung, and BAT: T<sub>3</sub> decreased with decreasing I availability and already did so in groups with normal circulating  $T_3$  but low  $T_4$ .

D1 activity in liver and lung showed only small changes in the LID group, which did not persist in LID' rats. In contrast, the activity of D2 in the cerebral cortex and BAT increased, with the differences between the LID + 1.0 and the C groups being statistically significant. The activities of the D2 isoenzyme increased up to 9.5- and 7-fold in the cerebral cortex and BAT, respectively, of LID', compared with C rats.

Figure 5 summarizes the changes observed in  $T_4$  and  $T_3$  in cerebellum, pituitary, kidney, ovary, adrenal, heart, and muscle with decreasing I availability.  $T_4$  decreased in all

FIG. 4. Changes in the concentrations of  $T_4$  and  $T_3$  and D1 and D2 activities in the liver, lung, brain, and BAT with decreasing I availability. The values corresponding to controls (LID + 5.0 group) were: liver, 40.00  $\pm$  4.23 ng  $T_4/g$ , 3.59  $\pm$  0.12 ng  $T_3/g$ , 47.5  $\pm$  1.4 pmol/min per mg protein (D1); lung, 8.08  $\pm$  1.35 ng  $T_4/g$ , 1.71  $\pm$  0.15 ng  $T_3/g$ , 528  $\pm$  84 fmol/h per mg protein (D1); brain, 1.81  $\pm$  0.09 ng  $T_4/g$ , 2.44  $\pm$  0.20 ng  $T_3/g$ , 10.9  $\pm$  1.4 fmol/h-mg protein (D2); BAT, 2.04  $\pm$  0.21 ng  $T_4/g$ , 3.02  $\pm$  0.18 ng  $T_3/g$ , 42  $\pm$  6 fmol/h-mg protein (D2). The meaning of symbols is the same as in Fig 1.



FIG. 5. Changes in the concentrations of  $T_4$  and  $T_3$  in the cerebellum, pituitary, kidney, ovary, adrenal, heart, and muscle with decreasing I availability. The upper left panel shows the corresponding plasma concentrations for comparison. The values corresponding to controls (LID + 5.0 group) were: cerebellum, 3.98  $\pm$  0.31 ng T<sub>4</sub>/g and 2.18  $\pm$ 0.14 ng T<sub>3</sub>/g; pituitary, 54  $\pm$  10 pg T<sub>4</sub>, 93  $\pm$  6 pg T\_3; kidney, 20.70  $\pm$  0.80 ng T<sub>4</sub>/g, 7.08  $\pm$  0.79 ng T<sub>3</sub>/g; ovary, 1.18  $\pm$ 0.07 ng T<sub>4</sub>,  $0.15 \pm 0.01$  ng T<sub>3</sub>; adrenal,  $0.30 \pm 0.04$  ng T<sub>4</sub>,  $0.20 \pm 0.07$  ng T<sub>3</sub>; heart, 3.82  $\pm$  0.49 ng T<sub>4</sub>/g, 1.54  $\pm$  0.18 ng T<sub>3</sub>/g; muscle, 1.91  $\pm$  0.19 ng T<sub>4</sub>/g, 1.18  $\pm$  0.06 ng T\_3/g. The values shown for the pituitary, ovary, and adrenal are the total  $T_4$  and  $T_3$  contents because the recorded weights were considered unreliable. The meaning of symbols is the same as in Fig. 1.



these tissues following patterns similar to those of plasma T<sub>4</sub> or FT<sub>4</sub>. As already described for the liver, lung, brain, and BAT (Fig. 4), the patterns of change in  $T_3$  varied greatly among these other tissues. The greatest difference was found between the patterns for the ovary and adrenal. In the ovary there is a very remarkable increase of  $T_3$  in the animals on LID + 1.0, LID + 0.5, and LID, compared with LID + 5.0, with an almost 2-fold increase in the LID + 0.5group. Ovarian T<sub>3</sub> is still higher than that of the controls, even in the LID' group, in which serum T<sub>3</sub> was decreased by about 50%.<sup>1</sup> On the contrary, T<sub>3</sub> in the adrenal decreases steadily with decreasing I availability, almost in parallel to adrenal  $T_4$ , and more markedly than circulating  $T_3$ : values in the adrenals of LID' animals are only 17% of those of the C group, whereas plasma  $T_3$  is still 46%. As already described above for the lung, the muscle and heart maintained normal T<sub>3</sub> concentrations, even in LID' animals. In these animals, T<sub>3</sub> decreased only in the cerebellum, pituitary, and kidney and did so only to 67-73% of C values, less than the decrease in circulating T<sub>3</sub>.

## Discussion

# The experimental design

The I intake of the female rats on LID + 5.0  $\mu$ g/d was sufficient for adequate growth. The I content of the stock pelleted diet used in our breeding facilities was determined

and found to vary between 3 and 7  $\mu$ g I per 20 g. The daily thyroidal secretion rate of strain-, sex-, and age-paired rats on this diet is estimated as being equivalent to 0.9  $\mu$ g T<sub>4</sub> + 0.15  $\mu$ g T<sub>3</sub> per 100 g BW (23), which together contain 0.59  $\mu$ g I from T<sub>4</sub> + 0.09  $\mu$ g I from T<sub>3</sub> = 0.68  $\mu$ g of I per 100 g BW. Present rats weighed less than 300 g at the end of the experiment, and turnover of hormonal I should be 2.04  $\mu$ g I or less per day. Thus, a daily supplement of 5.0  $\mu$ g I ought to cover physiological requirements, a conclusion in agreement with previous findings by others (8), namely that an I intake of 3  $\mu$ g/d or higher already ensures maximum levels of T<sub>4</sub> and T<sub>3</sub> in rat tissues.

The composition of the LID was the same for the five groups of experiment B, with the I content of the daily rations being the single controlled variable from the LID + 5.0 to the LID group. Results of experiment A also support that the amount of I available to the thyroid is the single variable between the LID' group and the other animals because the very low amount of added KClO<sub>4</sub> (1 mg/d) did not have, *per se*, any effect on thyroid hormone status in the I-sufficient (LID + I) groups.

Our experimental design therefore avoided differences between the experimental groups that could be related to sex, strain, and nutritional composition of the diet, all of which are known to affect thyroid hormone status.

As far as could be assessed from the increment in BW, the decrease in I availability did not affect the general condition of the animals until it became severe (LID and LID' groups). Despite their smaller increase in BW, even the animals of the LID' group did not show clinical signs of hypothyroidism comparable with those of animals that are thyroidectomized and stop growing within a few weeks. This is in agreement with previous results from this laboratory using the same strain of animals and type of LID (16, 18).

<sup>&</sup>lt;sup>1</sup>As indicated in *Materials and Methods*, the ovarian extracts were purified by a modified protocol to increase recovery of the iodothyronines. The possibility that interfering substances had been carried over into the  $T_3$  RIA samples was tested. Extracts of the ovaries from the LID' animals were tested at six different successive 2-fold dilutions. The plot of the specifically antibody-bound tracer *vs.* the log of the aliquot volume was parallel to that of obtained with the  $T_3$  standard. This result confirmed that  $T_{3\nu}$  and not artifacts, was being measured in the ovaries.

## Response to different grades of ID

Present findings fully confirm that the mechanisms involved are clearly dependent on the degree of I availability to the thyroid. For this reason, we will discuss them separately for animals with mild (LID + 1.0 group), moderate (LID + 0.5 group), severe (LID group), and very severe (LID' group) ID (Table 2).

# Mild ID: LID + 1.0 group

Intrathyroidal response mechanisms predominate when the daily I intake is reduced from 5 to 1  $\mu$ g, no longer sufficient to compensate for daily requirements. This is confirmed by a notable increase in thyroid weight that already accounts for a major part of the total increase observed with decreasing I availability. There is also an increase in the thyroid "Free" T<sub>3</sub> to "Free" T<sub>4</sub> ratios. We wish to point out that these changes occur without an increase in circulating TSH. This finding initially surprised us until we reviewed earlier studies in rats on an I intake similar to that of our LID + 1.0 group (6, 9, 12, 24–27): An increase in vascularity, blood flow, I trapping, acinar cell height, and hyperplasia of the gland have all been described to occur without any significant increase in circulating TSH. The same occurs with the changes in intrathyroid I metabolism, which lead to a preferential synthesis and secretion of T<sub>3</sub> over T<sub>4</sub>, and an increased  $T_3$  to  $T_4$  ratio in the circulation. These changes are correlated to the degree of I depletion and are the opposite of the TSH-independent response of the thyroid exposed to a sudden I excess (28). Their independence from serum TSH has been confirmed in hypophysectomized rats fed LID (24, 26). Our present data, therefore, confirm an important role of thyroid autoregulatory responses in the efficient adaptation to a mild degree of ID.

The mechanism(s) involved in these autoregulatory processes have not yet been identified. It is possible that a reduced availability of I decreases the thyroid content of iodolactones, which are involved in TSH-independent hyperplasia of the gland (29). A possible role of the sodium/ iodide symporter in ID has hardly been addressed experimentally in rats, with the exception of a report (30) that its expression increased in the thyroid of ID fetuses with normal plasma TSH.

Extrathyroidal response mechanisms involved in these mildly ID rats are less evident than the intrathyroidal autoregulatory ones that result in the higher plasma  $T_3$  to  $T_4$  ratio, resulting mainly from the preferential secretion of  $T_3$  over  $T_4$ . The patterns of the changes in the concentrations of  $T_4$  and

 $T_3$  in the tissues studied here, however, appear to be tissue specific and not easily predictable from the change in circulating  $T_4$  and  $T_3$ . A similar conclusion was reached years ago by Heninger and Albright (8), who measured the concentrations of  $T_4$  and  $T_3$  in several tissues of rats on a diet with an I content similar to that of the present LID + 1.0 group: they also found that  $T_3$  increased in plasma and many tissues but not all (*i.e.* the brain) and that the degree of change was tissue specific. Extrathyroidal responses appear to be involved, as shown by the increased D2 activities in brain and BAT.

The increase in T<sub>3</sub> concentration in many of the tissues studied here, such as the liver, lung, kidney, and muscle, were predictable from their known major dependency on plasma-derived T<sub>3</sub>, Similarly, the lack of an increase in the brain, cerebellum, and pituitary would be expected from their dependency on locally generated T<sub>3</sub> by D2 deiodination of T<sub>4</sub> (31). BAT, however, presented some unexpected features, being usually included among the tissues that derive intracellular  $T_3$  locally from  $T_4$  by D2. In these tissues D2 activity increases when T<sub>4</sub> decreases. Indeed, this is what occurred in the BAT of mildly I-deficient rats (see Fig. 4), as expected. Not anticipated, however, was the marked increase in BAT T<sub>3</sub>, comparable with that in plasma, and not observed in other experimental models (32). Whichever the mechanisms involved, the increased generation of  $T_3$  in BAT may actually contribute, together with the preferential secretion of  $T_3$  by the gland, to the marked increase in systemic  $T_3$  of the LID + 1.0 rats (33).

An increased activity of D2 in the brain had been expected from the decrease in brain  $T_4$ , a response amply shown in rats on LID with markedly decreased plasma  $T_4$  (34), but present results show that D2 responds to a much milder degree of ID than previously described.

The lack of increase of  $T_3$  content of the adrenals had also not been anticipated because this tissue has been considered dependent on serum-derived  $T_3$  (31, 32). The marked increase in the concentration of  $T_3$  in the ovary, a tissue not previously included in other studies (31), suggests its dependence on plasma-derived  $T_3$ .

In summary, both intra- and extrathyroidal mechanisms are involved in the response of the rat to mild ID: the former are autoregulatory and very effective in avoiding  $T_3$  deficiency in most tissues, and the latter occur in tissues in which D2 is important for local generation of  $T_3$ . In mild ID, hypothyroidism, as inferred from the concentrations of  $T_3$ , is avoided in all tissues studied.

**TABLE 2.** Mean values of thyroid weight and I content and circulating  $T_4$ , FT4,  $T_3$ , and TSH, expressed as percent of the control values (LID + 5.0 group), used in the present study to define different grades of ID

Grade of ID	Group	I, $\mu g/d^a$	Thyroid weight	Thyroid I	Plasma ${\rm T}_4$	Plasma $\mathrm{FT}_4$	Plasma ${\rm T}_3$	Plasma TSH
No ID	LID + 5.0 (controls)	5.0	100	100	100	100	100	100
Mild	LID + 1.0	1.0	187 ↑	$61\downarrow$	82	$57 \downarrow$	<b>139</b> ↑	135
Moderate	LID + 0.5	0.50	196 ↑	$23 \downarrow$	$51\downarrow$	$32 \downarrow$	105	<b>593</b> ↑
Severe	LID	0.052	$204$ $\uparrow$	$5\downarrow$	$25 \downarrow$	$15 \downarrow$	96	$634$ $\uparrow$
Very severe	LID'	0.021	$248$ $\uparrow$	6 ↓	$5\downarrow$	$2\downarrow$	$46 \downarrow$	976 ↑

<sup>*a*</sup> Estimated values; see Table 1.

 $\uparrow$  and  $\downarrow$ , Significant increase or decrease *vs.* LID + 5.0 group.

## Moderate ID: LID + 0.5 group

There is a further decrease in the I, "Total"  $T_4$ , and "Total"  $T_3$  contents of the thyroid to about 25% of control values. A new intrathyroidal-adaptive mechanism becomes evident, namely an increase in D1 activity, which could deiodinate the "Free"  $T_4$  fraction and the plasma  $T_4$  entering the gland. This could prevent a further decrease of the "Free"  $T_3$  content and contribute to maintenance of a normal plasma  $T_3$  as effectively as, or more than, the preferential intrathyroidal synthesis of  $T_3$ . We do not know whether the increase in D1 activity is an autoregulatory mechanism or the consequence of the increase in circulating TSH observed in this group. In contrast, thyroid weight is hardly affected by the increase in TSH, a finding consistent with the concept that thyroid growth is mainly determined by autoregulatory processes in both mild and moderate ID.

Plasma  $T_3$  was no longer elevated in the moderately ID animals but remained normal despite a 50% decrease of  $T_4$ .  $T_3$  deficiency was prevented in tissues that derive  $T_3$ mostly from plasma and also in BAT and cerebellum. Unexpectedly, despite normal  $T_3$  and very low  $T_4$  in plasma, some tissues maintained high  $T_3$  levels, most notably the lung, ovary, and muscle. The underlying mechanisms have not been identified: in the lung, for instance, an increase in D1 activity was not involved. In the brain  $T_3$  decreased, despite the increased D2 activity, and so did pituitary  $T_3$ . As already noted in the mild ID group,  $T_3$ decreased in the adrenal in parallel with  $T_4$ .

In summary, the case of moderate ID, intra- and extrathyroidal responses are still adequate to prevent low  $T_3$  levels in plasma and most tissues, despite a reduction of the I intake to 25% of that of controls. Some tissues even maintain elevated  $T_3$  concentrations, whereas others are markedly (adrenal) or moderately (brain and pituitary)  $T_3$  deficient.

# Severe I deficiency: LID group

The intra-thyroidal response mechanisms operative in previous groups continue to minimize the effects of a further marked decrease of the I intake and circulating  $T_3$  remains normal. A role is also likely to be played by the marked increase in thyroid D1 activity, which would avoid a further decrease of the "Free"  $T_3$  concentration. This increase occurs without a further concordant increase in TSH; it might be an autoregulatory process, but the influence of the higher than normal serum TSH cannot be excluded.

Despite a major decrease in plasma  $T_4$  to 25% of control values,  $T_3$  concentrations not only remained normal in plasma, but also in most tissues. The role of a further increase of D2 activity is evident in those tissues where it was measured. Again, the most unexpected and striking results are those obtained for the concentration of  $T_3$  in the ovary and lung, where it is much higher, and in the adrenal, where it is much lower, than expected from the normal circulating  $T_3$ .

In summary, despite a 100-fold decrease in I availability, a combination of intra- and extrathyroidal adaptive mechanism still mitigates  $T_3$  deficiency and presumably hypothyroidism in most but not all tissues.

# Very severe ID: LID' group

Intrathyroidal adaptive mechanisms are no longer sufficient in LID' rats to ensure a normal  $T_3$ , which decreases in plasma to 45% of C values, and also in many tissues that depend on plasma-derived  $T_3$ , including the liver. Despite the marked increase in D2 activity, the brain, cerebellum, pituitary, and BAT are  $T_3$  deficient, probably because of the very low availability of plasma  $T_4$  that has decreased to 5% of normal values. We have previously shown that the brain, pituitary, and liver of such animals are indeed hypothyroid, as assessed from several biological end points of thyroid hormone action (11). Tissue  $T_3$ , however, decreases less than would be expected from the circulating  $T_3$  level (adrenals again excepted), and some tissues continue having normal (muscle, heart) or even elevated (lung, ovary)  $T_3$  concentrations.

In summary, the threshold I availability below which most tissues are  $T_3$  deficient appears to be reached when the intraand extrathyroidal adaptive mechanisms are no longer capable of ensuring a normal circulating  $T_3$ . But even then, adaptive mechanisms that protect most tissues, and especially the heart, muscle, and ovary, become operative from the degree of  $T_3$  that would be expected from the decrease in plasma  $T_3$ .

In the present study, we did not measure the activities of the iodothyronine deiodinase isoenzymes in most tissues or activities of other enzymes, such as sulfotransferases and sulfatases, or the concentrations of  $T_4$  and  $T_3$  sulfates (35) that might regulate the local availability of  $T_3$  in different tissues. Also not investigated were the possible adaptive roles of changes in tissue uptake and/or exit rates of the iodothyronines (36) that might further minimize  $T_3$  deficiency in tissues. In this context, it is interesting that tissue to plasma  $T_3$  and  $T_4$  ratios were increased in some tissues of the LID rats and in most of those of the LID' group (data not shown).

# $T_4$ , $T_3$ , and TSH feedback in ID

We wish to draw attention to the findings illustrated in Figs. 2 and 3 that clearly show that in conditions of decreased I availability, plasma-derived T<sub>3</sub> plays a very decisive role in the regulation of circulating TSH. The TSH response is one tenth, or less, that observed in rats with primary thyroid failure and comparable degrees of hypothyroxinemia but with parallel decreases in plasma  $T_3$  (15). We believe it is very important and probably quite relevant for inhabitants of areas with mild and moderate ID to realize that a normal plasma TSH does not exclude, *per se*, a selective T<sub>3</sub> deficiency in tissues, such as the brain, that are affected by the decreased availability of  $T_4$  as substrate for the local generation of  $T_3$ . There are also other experimental situations, in which serum TSH is more closely correlated to circulating  $T_3$  than  $T_4$  (37). Such findings also point to a more important role of plasmaderived  $T_3$  in the inhibition of TSH release, compared with that of  $T_3$  generated locally from  $T_4$  by D2.

Present results constitute a clear contradiction to the previous idea that findings in I-deficient animals and humans, in which a close negative correlation is found between circulating TSH and  $T_4$ , despite normal  $T_3$ , are a prime example of the beneficial adaptive role of the local generation of  $T_3$  from  $T_4$ : it was, moreover, difficult to explain the adaptive advantage of the observed increase in D2 activity in the pituitary because it would increase locally generated  $T_3$ , shutting down the compensatory mechanism, namely an increase in plasma TSH.

# Conclusion

ID rats are endowed with numerous and very efficient adaptive mechanisms, most of which require a fully functioning normal thyroid gland, and are thus lost in animals with primary thyroid failure. ID rats are often considered to be either hypothyroid, because of their low circulating T<sub>4</sub> and increased TSH, or euthyroid, because of their normal (or increased) plasma  $T_{3}$ , but present results stress that neither assumption is correct: thyroid hormone status is not only related to the degree of depletion of I availability to the thyroid but is also eminently tissue specific. As discussed elsewhere (15), few tissue-specific direct effects of thyroid hormone action are available, and we measured the concentration of  $T_3$  in the tissues as a first step in assessing their thyroid hormone status. With a moderate, or even severe, I deficiency, most tissues depending on plasma T<sub>3</sub> would have normal, or even slightly elevated, T<sub>3</sub> concentrations, most notably the ovary, lung, heart, and muscle. However, those tissues that depend to an appreciable extent on T<sub>4</sub> for the local generation of  $T_3$  are protected from  $T_3$  deficiency to a lesser degree. As a consequence, in the latter type of tissue, thyroid hormone-sensitive functions are more likely to be affected than those characteristic of tissues depending on plasma-derived T<sub>3</sub>. Such is the case of the brain, for instance, and cerebral functions may already be impaired in moderate ID because brain  $T_3$  is already decreased. In the mildly ID group, in which plasma TSH was normal and plasma FT<sub>4</sub> was slightly decreased, total brain T<sub>4</sub> was decreased and D2 activity was increased. This prevented a decrease of  $T_3$  in the total brain but not necessarily in all brain areas (13, 38). Present results also indicate that the degree of increase in D2 activity in different cerebral structures of ID rats does not permit, per se, conclusions to be drawn regarding their protection from T<sub>3</sub> deficiency because the latter also involves the amount of  $T_4$  available in each area (13).

Circulating  $T_3$  has to decrease before many  $T_3$ -dependent tissues become  $T_3$  deficient. This appears to occur when circulating  $T_4$  decreases from 25 to 5% of normal values. But even under such conditions, the many known and as-yetundefined intra- and extrathyroidal adaptive mechanisms are efficient enough to maintain euthyroidism in muscle and heart and even slightly elevated  $T_3$  in lung and ovary. The findings in the ovary may underlie the observation that even very severely I-deficient animals are easily mated, do not show decreased fertility, and bear litters of normal size (16, 18), in contrast to Tx or goitrogen-treated hypothyroid females (19).

## Possible implications for man

As already pointed out in the introductory text, the present study is relevant only for inhabitants of areas in which ID is the sole cause of goiter but not for areas in which other environmental factors may lead to destruction of the gland and therefore to clinical hypothyroidism. We believe present results in mild and moderate grades of decreased I availability are especially relevant for man because such conditions are still widespread in Western industrialized countries (39).

Many of the findings in rats, described here and by others, have also been described in people from areas with an adequate I intake who are changed to an I-deficient diet or for inhabitants of the ID areas defined above. Thus, for instance, the gland responds within a few days with a striking increase in blood flow, occurring before any change is detected in plasma  $T_4$  or TSH (40). Increased I trapping and circulating  $T_3$  to  $T_4$  ratios have also been shown (3). In simple sporadic goiter, the increase in thyroid volume occurs without a necessary increase in circulating TSH (41-44). Even in very severe ID, the increase in circulating TSH is markedly blunted, compared with that usually observed in hypothyroid patients (43-46). Missler et al. (45) reported that in 304 children from an ID area, 60% had an enlarged gland, but only 9% had TSH greater than 4.5 mU/liter. In the seminal studies by Glinoer (47) on thyroid function in pregnant women from a population with mild to moderate ID, TSH levels above the normal reference range were hardly ever found at any stage of pregnancy, even among the women with the lowest first-trimester FT<sub>4</sub> levels. The same was observed in pregnant women from an area with very mild ID (48, 49).

Western-trained physicians dealing mostly with patients with primary thyroid failure rely on an increased circulating TSH for their classification of overt or subclinical hypothyroidism and are often unfamiliar with the concept of hypothyroxinemia: a decreased T<sub>4</sub> without an increase of TSH above normal. The TSH-independent autoregulatory mechanisms controlling thyroid function in ID are often overlooked, as these mechanisms are better known in the case of iodine excess. The present experimental data obtained with mild to moderate grades of ID stress the primary importance of autoregulatory mechanisms that protect many tissues from overt  $T_3$  deficiency. Even in very severe ID (2), people are able to sustain heavy physical work and have normal cardiac function, observations that might be related to the present findings in muscle and heart, which maintain normal  $T_3$  concentrations, even in the LID' group.

As discussed elsewhere in more detail (49), it is inaccurate to assume that inhabitants of ID areas are clinically hypothyroid individuals. The present experimental model supports the epidemiological findings that inhabitants of ID areas are not clinically hypothyroid individuals because their normal circulating  $T_3$  ensures normal  $T_3$  concentration in most tissues. But, as shown here, this might not avoid selective  $T_3$  deficiency of those tissues, such as the brain, that depend mostly on  $T_4$  for their intracellular  $T_3$  supply. In man, this might already negatively affect mental functions in mild ID (5, 39, 46, 49–51), in which inhabitants are often described as dull (52).

It is often assumed that eradication of severe ID is enough to avoid the most important IDD, including those affecting mental processes. Present experimental results obtained in moderately ID rats stress that this is not so, and countries with areas of moderate ID should actively correct it: an important proportion of their inhabitants, and their future progeny, may still suffer from easily preventable impairments of mental functions and the consequent socioeconomic implications (49) as well as the increased incidence of thyroid disorders ensuing from thyroid hyperplasia.

#### Acknowledgments

We thank Mrs. Socorro Duran and Mrs. Maria Jesus Presas for their very competent technical involvement in this study, including all determinations of I,  $T_4$ ,  $FT_4$ ,  $T_3$ , and TSH in plasma and  $T_4$  and  $T_3$  in thyroid and tissues.

Received October 18, 2005. Accepted January 23, 2006.

Address all correspondence and requests for reprints to: Dr. G. Morreale de Escobar, Ph.D., Instituto de Investigaciones Biomédicas Alberto Sols, Arturo Duperier, 4, 28029 Madrid, Spain. E-mail: gmorreale@ iib.uam.es.

This work was supported by Fondo de Investigaciones Sanitarias RCMN (C03/08) from the Instituto de Salud Carlos III and Plan Nacional SAF 2001/2243.

Disclosure summary: None of the authors has potential conflicts of interest.

#### References

- 1. Hetzel BS 1983 Iodine deficiency disorders (IDD) and their eradication. Lancet ii:1126-1129
- Hetzel BS 2002 Eliminating iodine deficiency disorders—the role of the International Council in the global partnership. Bull World Health Organ 80: 410–413
- Delange F 2000 Iodine deficiency. In: Braverman LE, Utiger RD, eds. Werner and Ingbar's the thyroid. Philadelphia: Lippincott Williams and Wilkins; 295– 315
- 4. Dunn JT, Delange F 2001 Damaged reproduction: the most important consequence of iodine deficiency. J Clin Endocrinol Metab 86:2360–2363
- Delange F 2001 Iodine deficiency as a cause of brain damage. Postgrad Med I 77:217–220
- 6. **Studer H, Greer M** 1965 A study of the mechanisms involved in the production of iodine-deficient goiter. Acta Endocrinol (Copenh) 49:610–628
- Silva E 1972 Disposal rates of thyroxine and triiodothyronine in iodine-deficient rats. Endocrinology 91:1430–1435
- 8. Heninger RW, Albright EC 1975 Alteration in tissue and serum concentrations of TSH, iodide, T4 and T3 induced by various dietary iodide levels. Proc Soc Exp Biol Med 150:137–141
- 9. Riesco G, Taurog A, Larsen PR 1976 Variations in the response of the thyroid gland of the rat to different low-iodine diets, correlation with iodine content of diet. Endocrinology 99:270–280
- Okamura K, Taurog A, Krulich L 1981 Elevation of serum 3,5,3'-triiodothyronine and thyroxine levels in rats fed Remington diets; opposing effects of nutritional deficiency and iodine deficiency. Endocrinology 108:1247–1256
- Santisteban P, Obregón MJ, Rodríguez-Peña A, Lamas L, Escobar del Rey F, Morreale de Escobar G 1982 Are iodine-deficient rats euthyroid? Endocrinology 110:1780–1789
- 12. Michalkiewicz M, Huffman L, Connors JM, Hedge GA 1989 Alterations in thyroid blood flow induced by varying levels of iodine intake in the rat. Endocrinology 125:54–60
- Peeters R, Fekete C, Goncalvez C, Legradi G, Tu HM, Harney JW, Bianco AC, Lechan RM, Larsen PR 2001 Regional physiological adaptation of the central nervous system deiodinases to iodine deficiency. Am J Physiol Endocrinol Metab 281:E54–E61
- 14. Contempré B, Morreale de Escobar G, Denef JF, Dumont JE, Many MC 2004 Thiocyanate induces cell necrosis and fibrosis in selenium- and iodine-deficient rat thyroids: a potential experimental model for myxedematous cretinism in central Africa. Endocrinology 145:994–1002
- Escobar-Morreale HF, Escobar del Rey F, Obregon MJ, Morreale de Escobar G 1995 Replacement therapy for hypothyroidism with thyroxine alone does not ensure euthyroidism in all tissues, as studied in thyroidectomized rats. J Clin Invest 96:2828–2838
- Escobar del Rey F, Mallol J, Pastor R, Morreale de Escobar G 1987 Effects of maternal iodine deficiency on thyroid hormone economy of lactating dams and pups: maintenance of normal cerebral 3,5,3'-triiodo-L-thyronine concentrations in pups during major phases of brain development. Endocrinology 121: 803–811
- 17. Morreale de Escobar G, Pedraza PE, Escobar del Rey F, Obregón MJ 1996 Thyroidal and extrathyroidal adaptation to graded degrees of iodine deficiency: an experimental rat model for the study of neurological iodine defi-

ciency disorders (IDD). In: Braverman LE, Köhrle J, Eber O, Langsteger W, eds. Thyroid and trace elements. Vienna: Blackwell; 113–126

- Lavado-Autric R, Ausó E, Garcia-Velasco JV, Arufe MC, Escobar del Rey F, Berbel P, Morreale de Escobar G 2003 Early maternal hypothyroxinemia alters histogenesis and cerebral cortex cytoarchitecture of the progeny. J Clin Invest 111:1073–1082
- Morreale de Escobar G, Pastor R, Obregón MJ, Escobar del Rey F 1985 Effects of maternal hypothyroidism on the weight and thyroid hormone content of rat embryonic tissues. Endocrinology 117:1890–1901
- Calvo RM, Jauniaux E, Gulbis B, Asunción M, Gervy C, Contempré B, Morreale de Escobar G 2002 Fetal tissues are exposed to biologically relevant free thyroxine concentrations during early phases of development. Possible consequences of maternal hypothyroxinemia. J Clin Endocrinol Metab 87: 1768–1777
- Calvo R, Morreale de Escobar G, Escobar del Rey F, Obregón MJ 1997 Maternal non-thyroidal illness and fetal thyroid hormone status, as studied in the streptozotocin-induced diabetes mellitus rat model. Endocrinology 138: 1159–1169
- 22. Obregón MJ, Ruiz de Oña C, Calvo R, Escobar del Rey F, Morreale de Escobar G 1991 Outer ring iodothyronine deiodinases and thyroid hormone economy: responses to iodine deficiency in the rat fetus and neonate. Endocrinology 129:2663–2673
- 23. Escobar-Morreale HF, Escobar del Rey FE, Obregon MJ, Morreale de Escobar G 1996 Only the combined treatment with thyroxine and triiodothyronine ensures euthyroidism in all tissues of the thyroidectomized rat. Endocrinology 137:2490–2502
- Chapman A 1941 The relation of the thyroid and pituitary glands to iodine metabolism. Endocrinology 29:680–685
- 25. Vanderlaan WP, Caplan R 1954 Observations on a relationship between total thyroid iodine content and the iodide-concentrating mechanism of the thyroid gland of the rat. Endocrinology 54:437–447
- Halmi NS, Sirtos BN 1955 Analysis of the modifying effects of dietary iodine levels on the thyroidal response of hypophysectomized rats to thyrotropin. Endocrinology 56:157–162
- Greer MA, Rockie C 1969 Effect of thyrotropin and the iodine content of the thyroid on the triiodothyronine:thyroxine ratio of newly synthesized iodothyronines. Endocrinology 85:244–250
- Nagataki S, Yokoyama N 1996 Autoregulation: effects of iodide. In: Braverman LE, Utiger RD, eds. Werner and Ingbar's the thyroid. Philadelphia: Lippincott-Raven Publishers; 241–247
- Gärtner R, Dugrillon A, Bechtner G 1996 Iodolipids and thyroid function and growth. In: Nauman J, Glinoer D, Braverman LE, Hostalek U, eds. The thyroid and iodine. Stuttgart, Germany: Schattener; 19–30
- Schröder-van der Elst J, van der Heide D, Kastellin J, Rousset B, Obregón MJ 2001 The expression of the sodium/iodide symporter is up-regulated in the thyroid of fetuses of iodide-deficient rats. Endocrinology 142:3736–3741
- van Doorn J, Roelfsema F, van der Heide D 1985 Concentrations of thyroxine and 3,5,3'-triiodothyronine at 34 different sites in euthyroid rats as determined by an isotopic equilibrium technique. Endocrinology 117:1202–1208
- 32. Escobar-Morreale HF, Obregón MJ, Escobar del Rey F, Morreale de Escobar G 1999 Tissue-specific pattern of changes in 3,5,3'-triiodo-L-thyronine concentrations in thyroidectomized rats infused with increasing doses of the hormone. Which are the regulatory systems? Biochimie (Paris) 81:453–462
- Silva JE, Larsen PR 1985 Potential of brown adipose tissue type II thyroxine 5'-deiodinase as a local systemic source of triiodothyronine in rats. J Clin Invest 76:2296–2305
- Obregón MJ, Escobar del Rey F, Morreale de Escobar G 2005 The effects of iodine deficiency on thyroid hormone deiodination. Thyroid 15:917–929
- Visser TJ 1994 Sulfation and glucuronidation pathways of thyroid hormone metabolism. In: Wu S, Visser TJ, eds. Thyroid hormone metabolism. London: CRC Press; 117
- Friesema EC, Jansen J, Visser TJ 2005 Thyroid hormone transporters. Biochem Soc Trans 33:228–232
- Emerson CH, Lew R, Braverman LE, DeVito WJ 1989 Serum thyrotropin concentrations are more highly correlated with serum triiodothyronine concentrations than with serum thyroxine concentrations in thyroid hormoneinfused thyroidectomized rats. Endocrinology 124:2415–2418
- 38. Kester MH, Martinez de Mena R, Obregon MJ, Marinkovic D, Howatson A, Visser TH, Hume R, Morreale de Escobar G 2004 Iodothyronine levels in the human developing brain: major regulatory roles of iodothyronine deiodinases in different areas. J Clin Endocrinol Metab 89:3117–3128
- Vitti P, Delange F, Pinchera A, Zimmermann M, Dunn JT 2003 Europe is iodine deficient. Lancet 361:1226
- Arntzenius AB, Smit LJ, Schipper J, van der Heide D, Meinders AE 1991 Inverse relation between iodine intake and thyroid blood flow: color Doppler flow imaging in euthyroid humans. J Clin Endocrinol Metab 73:1051–1055
- Barakat RM, Ingbar SH1965 The effect of acute iodine depletion on thyroid function in man. J Clin Invest 44:1117–1124
- Vagenakis A, Koutras D, Burger A, Malamos B, Ingbar S 1973 Studies of serum triiodothyronine, thyroxine and thyrotropin concentrations in endemic goiter in Greece. J Clin Endocrinol Metab 37:485–488

#### 2108 Endocrinology, May 2006, 147(5):2098-2108

- Patel YC, Pharoah OD, Hornabrook RW, Hetzel BS 1973 Serum triiodothyronine, thyroxine and thyroid-stimulating hormone in endemic goiter: a comparison of goitrous and nongoitrous subjects in New Guinea. J Endocrinol Metab 37:783–789
- 44. Pharaoh POD, Lawton NF, Ellis SM, Williams ES, Ekins RP 1973 The role of triiodothyronine (T3) in the maintenance of euthyroidism in endemic goitre. Clin Endocrinol (Oxf) 2:193–199
- 45. Missler U, Gutekunst R, Wood WG 1994 Thyroglobulin is a more sensitive indicator of iodine deficiency than thyrotropin: development and evaluation of dry blood spot assays for thyrotropin and thyroglobulin in iodine-deficient geographical areas. Eur J Clin Chem Clin Biochem 32:137–143
- Huda SN, Grantham-McGregor SM, Rahman KM, Tomkins A 1999 Biochemical hypothyroidism secondary to iodine deficiency is associated with poor school achievement and cognition in Bangladesh. J Nutr 129:980–987
- Glinoer D 1997 The regulation of thyroid function in pregnancy: pathways of endocrine adaptation from physiology to pathology. Endocr Rev 18:404–433

- 48. **Morreale de Escobar G, Escobar del Rey F** 2003 Consequences of iodine deficiency for brain development. In: deVijlder J, Morreale de Escobar G, eds. The thyroid and the brain. Stuttgart, Germany: Schattauer Verlag; 33–56
- Morreale de Escobar G, Obregón MJ, Escobar del Rey F 2004 Role of thyroid hormone during early brain development. Eur J Endocrinol 151:U25–U37
- Vitti P, Aghini-Lombardi F, Chiovato L, Ferretti G, Pinchera A 2003 Neuropsychological assessment in humans living in mild to moderate iodine deficiency. In: DeVijlder J, Morreale de Escobar G, eds. The thyroid and the brain. Stuttgart, Germany: Schattauer Verlag; 57–63
- 51. Vermiglio F, Lo Presti VP, Moleti M, Sidoti M, Tortorella G, Scaffidi G, Castagna MG, Violi MA, Crisà A, Artemisia A, Trimarchi F 2004 Attention deficit and hyperactivity disorders (ADHD) in the offspring of mothers exposed to iodine deficiency: a possible novel iodine deficiency disorder in developed countries? J Clin Endocrinol Metab 89:6054–6060
- Dunn JT 1992 Iodine deficiency. The next target for elimination? N Engl J Med 326:267–268

*Endocrinology* is published monthly by The Endocrine Society (http://www.endo-society.org), the foremost professional society serving the endocrine community.