Diagnostic value of the absolute free thyroxine iodine test

Review of 145 cases

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Several investigators¹⁻⁵ have proposed that the thyroid hormone circulates in two parts—a free unbound physiologically active part and a larger protein-bound metabolically inert part. If approximately 5.0 μ g/100 ml of thyroxine iodine is considered 100%, then the free thyroxine is approximately 0.05% of the total amount and is expressed in millimicrograms (nanagrams—n). The normal mean level is approximately 3.0 n/100 ml.

Several methods⁶⁻¹⁰ of measuring the amount of free thyroxine have been proposed, but the few large series^{7, 9, 11, 12} of patients who have been studied to determine the clinical usefulness and diagnostic accuracy of the test have depended upon the protein-bound iodine (PBI)) and the percentage of free thyroxine iodine and the product obtained thereof. Since March 1966, we have measured the absolute free thyroxine iodine based on total serum thyroxine iodine and have tested more than 1,000 patients who had a variety of thyroid and suspected thyroid diseases.† Measurement is similar to the free thyroxine index reported by others, ¹³⁻¹⁵ and our conclusions agree with theirs.

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[†] The serum thyroxine tests were performed in the laboratory of Jack R. Leonards, M.D., Ph.D., Case-Western Reserve University School of Medicine, Cleveland, Ohio.

Materials and methods

Records of 102 patients were used for the main body of this study, and tests on an additional 43 are included in the scatter graphs (Figs. 1–3). Each diagnosis was based on clinical appraisal, standard tests and, when necessary, on results of therapy. Many

EUTHYROID PATIENTS (106)

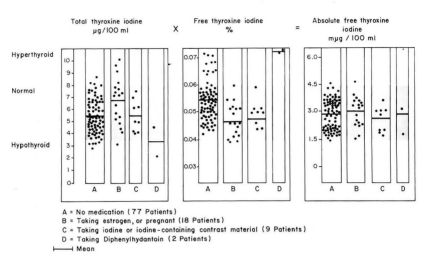


Fig. 1. Effect of exogenous drugs or materials on the absolute free thyroxine iodine levels in euthyroid patients.

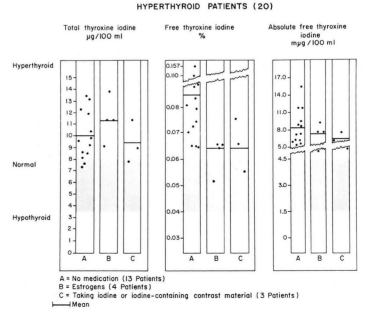


Fig. 2. Effect of exogenous drugs or materials on the absolute free thyroxine iodine levels in hyperthyroid patients.

HYPOTHYROID PATIENTS (19)

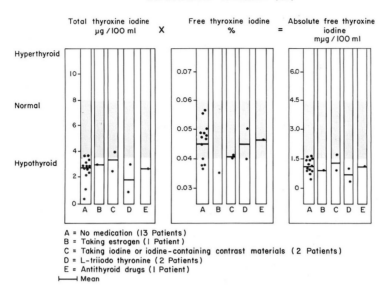


Fig. 3. Effect of exogenous drugs or materials on the absolute free thyroxine iodine levels in hypothyroid patients.

tests which were previously abnormal were repeated.

The effects of iodine contamination, drugs, and estrogens were considered when making a diagnosis. The tests referred to as "standard" included BMR, PBI, and 131 I thyroidal 24-hour uptake, and serum cholesterol determinations. The total serum thyroxine iodine (TT_4^I) and absolute free thyroxine iodine (AFT_4^I) were estimated on blood drawn on the same days as for the other tests.

The 102 patients were classified as follows:

	Patient
1. Clearly active hyperthyroidism	15
of Graves' disease	
2. Possible Graves' disease	12
3. Nodular goiter (toxic 2, sus-	10
pected 1, nontoxic 7)	
4. Myxedema (typical 8, suspected	13
and confirmed 5)	
Possible hypothyroidism	22
6. Euthyroid clinically and by	19
standard tests	
7. Euthyroid continuing various	11
types of treatment	

The total thyroxine iodine was measured by the method of Murphy and Pattee,16-18 with minor modifications; the percentage of free thyroxine was measured by the method of Ingbar et al.7 The quantity of total thyroxine iodine ($\mu g/100$ ml) was multiplied by the percentage of free thyroxine iodine to obtain a product called the absolute free thyroxine iodine. This terminology is similar to that used by Ingbar et al,7 Sterling and Brenner,9 and Anderson,11 except that we used total thyroxine iodine instead of proteinbound iodine, as did Arango et al,12 to avoid the difficulties inherent in the latter test, such as iodine contamination and hormone effect.

Normal values are: total thyroxine iodine, 3.5 to 8.0 $\mu g/100$ ml; free thyroxine iodine, 0.04 to 0.06% absolute free thyroxine iodine, 1.5 to 4.5 n/100 ml. These values represent the mean \pm 2 S D found in 40 medical students. Arango et al¹² reported normal values for AFT₄^I of 0.90 to 5.2 n/100 ml ex-

pressed as thyroxine rather than thyroxine iodine.

Results

The results are outlined in the tables (see appendix).

Table 1 shows the results obtained in patients with confirmed Graves' disease. In all cases the correlation of the clinical diagnosis, PBI, TT₄^I, and AFT₄^I was good. All TT₄^I tests repeated after successful therapy were normal. In case 13 therapeutic control was incomplete; as expected there was a large discrepancy between the PBI and TT₄^I. During continued treatment, however, the TT₄^I was normal but the AFT₄^I remained mildly elevated, suggesting that the latter may be a more delicate index of hyperthyroidism.

Table 2 shows the results obtained in patients with possible Graves' disease. In all cases except case 8 the TT_4^I and the AFT_4^I correlated with the final clinical diagnosis. In case 8 the TT_4^I was elevated and did not support the final impression that the patient was euthyroid.

Table 3 shows the results obtained in patients with nodular goiter. Patients 1 and 2 were hyperthyroid and the TT_4^I and AFT^I levels confirmed this. They were cured by removal of the goiter. Hyperthyroidism was suspected in case 3 but all tests were normal, and the patient was considered to be euthyroid. In the remaining 7 cases the AFT_4^I indicated the euthyroid state. However in one (case 5) the high BMR and the low TT_4^I were at variance with the final impression that the patient was euthyroid.

Table 4 shows the results obtained in patients with confirmed hypothy-

roidism. In all instances there was good correlation between the results of the TT_4^I and AFT_4^I tests and the diagnoses. The response to therapy further confirmed the diagnosis in all instances. In all other cases there was complete correlation of the diagnosis, the TT_4^I , and the AFT_4^I except in case 1, where the AFT_4^I correlated, but the TT_4^I did not.

Table 5 shows the results obtained in patients with possible hypothroidism and patients in whom it was considered necessary to exclude a diagnosis of hypothyroidism.

All 22 patients were considered to be euthyroid and the standard tests were consistent with this diagnosis except in case 19 (PBI 3.0 μ g/100 ml). In patient 22 the TT_4^I and AFT_4 levels were lower than normal. Whether these results are in error or whether the patient may develop hypothyroidism cannot be determined. No adequate explanation can be offered for the AFT_4^I in case 8.

Table 6 shows the results obtained in patients judged to be euthyroid. The PBI, TT₄^I, and AFT₄^I correlate well in 17 of these 19 patients. Estrogen was apparently responsible for raising the PBI and TT₄^I in the first patient, while the AFT₄^I remained normal. In patient 13, judged to be euthyroid clinically and by standard tests, the PBI is near the lower limit of normal; the AFT₄^I and the TT₄^I are abnormally low.

Table 7 shows the results obtained in euthyroid patients on various types of treatment. There is good correlation between the clinical diagnosis and tests in almost all instances. The PBI, TT₄I, and AFT₄I are depressed in case 3 by tri-iodothyronine. The low PBI in case 9 is unexplained and may

be in error. This patient was suspected of having hypothyroidism.

There appears to be a lack of correlation with the results of the TT_4^T in four cases and of the AFT_4^T in one case as compared to the final diagnosis as shown below.

Table	$\mathrm{TT_4^I}$	AFT ₄ I	Diagnosis
Table 3, case 8	0	+	Euthyroid
Table 4, case 3	+	0	Hypothyroid- ism
case 6	0	+	Hypothyroid- ism
Table 5, case 22	+	+	Hypothyroid- ism was sus- pected clini- cally, standard tests normal
Table 6, case	0	+	Alopecia areata
Table 7, case 1	0	+	Euthyroid

Effect of contamination with various nonhormonal organic iodides on the free thyroxine test. The contamination of the blood stream with a variety of iodine-containing materials had no effect on the free thyroxine test, but rendered both the PBI and 131I uptake abnormal. The use of antithyroid preparations and 1-tri-iodothyronine was followed by a low level of total thyroxine iodine, leaving the percentage of free thyroxine iodine normal and thus resulting in a low level of absolute free thyroxine iodine value. Desiccated thyroid and sodium 1-thyroxine had the opposite effect, raising both the total thyroxine iodine and the percentage of free thyroxine iodine, and therefore the absolute free thyroxine iodine. Iodine or iodide had no effect on the tests. The two patients

who had been taking diphenylhydantoin showed low total thyroxine with the percentage of free thyroxine iodide increased slightly over normal, leading to a normal absolute free thyroxine level. All eight patients shown in the tables who had had iodides and PBI levels over 50 had normal levels of TT_4^I varying from 1.6 to 6.6 μ g/ 100 ml and AFT₄^I varying from 0.6 to 4.3 n/100 ml.

Effect of various hormones and pregnancy on the free thyroxine test. Pregnancy or the use of estrogen had no significant effect on the AFT_4 level except in one patient, in whom it was 5.5 n/100 ml. The total thyroxine iodine was moderately increased because of increased binding, but the percentage of free thyroxine iodine was diminished, resulting in an absolute free thyroxine iodine value within normal range. 19

Comment

The use of the absolute free thyroxine iodine test gives an accurate laboratory measurement of thyroid function, except possibly in hyperthyroidism due to tri-iodothyronine.

We have found good correlation between the clinical evaluation of the patient and this test, and also good correlation between it and the standard tests (PBI and 131I uptake), when the latter are not changed by contamination of iodine or influenced by drugs, and usually remain within the normal range during estrogen therapy and pregnancy. The absolute free thyroxine iodine test has advantages over existing tests. It is not affected by contrast materials. It is changed by the effects of antithyroid drugs and thyroid preparations, and corresponds to the clinical state.

Since this review of our earliest cases, the findings of others have confirmed the above conclusions.²⁰ However, there have been several cases in which the values of the free thyroxine did not correlate with the clinical picture or other tests. These have been due in part to laboratory error (we estimate approximately one test in a hundred) or have been completely unexplainable, since the same figures were obtained after repeating the test several times.

It is unfortunate that the determination of the total thyroxine iodine and the percentage of free thyroxine iodine is not easily done in the average clinical laboratory; however, with experience one can obtain reproducible results in most cases without difficulty. Because the tests are technically difficult and do not add clinically useful information in the majority of cases, they are not now performed routinely in our laboratories.

Representative case reports

Case 1. A 37-year-old multigravida was examined in the 7th month of her fourth pregnancy because of a history and physical signs of thyrotoxicosis. She had apparently taken propylthiouracil intermittently before her initial examination. The results of tests were: BMR, +31%; PBI, $15~\mu g/100~ml$; T_3 red blood cell uptake, 15.5%; absolute free thyroxine iodine, 9.1~n/100~ml. The latter figures confirmed the initial clinical impression of hyperthyroidism, and treatment with propylthiouracil was reinstituted with subsequent clinical improvement.

Case 2. A 62-year-old woman appeared to be hypothyroid by history and physical examination. She had recently had a urogram. The PBI was over 50 μ g/100 ml; the ¹³¹I uptake was less than 1%; and absolute free thyroxine iodine was 1.0 n/

100 ml, confirming the diagnosis of hypothyroidism.

Case 3. A 37-year-old woman was thought to be hyperthyroid. She was quite anxious, was taking oral contraceptives, and recently had undergone cholecystography. The BMR was +18%; PBI 50 μ g/100 ml; T₃ red blood cell uptake 15%; ¹³¹I uptake 49%; serum cholesterol 190 mg/100 ml; and absolute free thyroxine iodine, 7.6 n/100 ml. The patient was treated with ¹³¹I, and within 2 months the absolute free thyroxine iodine level had fallen to 1.5 μ g/100 ml and the patient appeared to be euthyroid.

Cases 4, 5 and 6. Three patients with possible thyroid crisis, but with confusing histories, physical findings, and laboratory data, were found to have extraordinarily high values of absolute free thyroxine iodine (from 18 to 24 n/100 ml). These results were of considerable assistance in reaching an accurate diagnosis. The patients were treated for thyroid crisis with improvement in their clinical courses as the levels of free thyroxine decreased toward normal.

Acknowledgments

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APPENDIX

Table 1.—Results in confirmed Graves' disease

Case	Factors possibly affecting	BMR	PBI	¹³¹ I 24-hr uptake	Total serum TT ₄ ¹ µg/100	AFT ₄ 1 n/100	Total serum TT ₄ ^I µg/100 ml	AFT ₄ I n/100 ml
no.	test results	%	100 ml	%	ml	ml	After t	herapy
1	Pregnancy PTU*	+31	15		13.9	9.1		
2	17	+4 5	17.1	2	14.9	19.9		
3			11.2	41	9.3	7.3	4.5	3.0
4		+19	11.4	36	13.3	8.6		
5	PTU therapy thy- roid	+16	16.5		10.4	11.4	6.8	4.5
6	0		12.3	46	9.8	15.4		
7	Estrogen 127I	+18	9.2	4 9	11.4	7.6	3.6	1.5
8	0	+53	14.0	77	13.5	11.9		
9	PTU therapy	+46			8.7	6.1		
10	Estrogen		9.9	33	10.3	6.9	4.8	2.6
11	0	+30	14.5	54	9.6	6.2	6.7	3.5
12	0	+43		59	17.4	9.3	5.0	2.7
13	Pneumonia, KI and PTU ther- apy when tested		17.0	39	7.6	5.5	7.2	5.0
14	- •	+40	7.3	22	8.6	7.1	6.5	4.6
15		+4 9	8.3	7 0	12.5	12.7		

^{*} PTU = propylthiouracil.

Table 2.—Results in possible Graves' disease

Case no.	Factors possibly affecting test results	PBI µg/100 ml	¹³¹ I uptake 24-hr	${{ m TT_4^I} \atop \mu { m g}/100} \atop { m ml}$	AFT ₄ I n/100 ml	Final clinical diagnosis
1	KI	9.9	20%	5.0	3.0	Euthyroid
2		7.8	27%	7.7	3.6	Euthyroid
3	Estrogen	9.2	20%	8.1	3.7	Hyperthyroid
4	Thyroid	6.2	<1%	5.0	2.4	
5	Thyroiditis	7.5	21%	6.6	3.8	Euthyroid
6	•	8.1		7.1	3.9	Euthyroid
7				4.4	2.4	Euthyroid, BMR +3%
8	Estrogen	9.1		10.1	3.9	Euthyroid
9	Estrogen, Na LT4*	6.6		7.2	3.3	Euthyroid
10	<u>-</u>	>25		5.4	3.5	Euthyroid, BMR +13%
11		8.5		5.3	2.7	Euthyroid
12	Pyelogram	>25		7.5	3.7	Euthyroid

^{*} NaLT₄ = sodium levothyroxine.

Table 3.—Results obtained in patients with nodular goiter

Case no.	Thyroid status and factors possibly affecting test results	BMR %	$\begin{array}{c} \mathrm{PBI} \\ \mu\mathrm{g}/100 \\ \mathrm{ml} \end{array}$	¹⁸¹ I 24-hr uptake %	${ m TT_4^I} \ { m \mu g/100} \ { m ml}$	AFT ₄ I n/100 ml
1	Hyperthyroid	+19	11.4	36	13.0	8.6
2	Hyperthyroid; mercury diuretic	+40	7.3*	22	8.6	7.1
3	Euthyroid		25		6.1	2.6
4	Euthyroid, estrogen	-5	14.6†		7.4	4.0
5	Euthyroid	+29	4.8		4.9	2.6
6	Euthyroid	+8		18	6.9	3.6
7	Euthyroid				7.8	4.4
8	Euthyroid		5.4		8.6	3.6
9	Euthyroid		6.5		6.6	3.9
10	Euthyroid; mesantoin		4.2		4.5	3.0

^{*} Possibly affected by the diuretic.

Table 4.—Results obtained in patients with hypothyroidism

Case no.	Factors possibly affecting test results	BMR %	PBI µg/100 ml	Choles- terol mg/100 ml	131 J 24-hr uptake	${{\mathrm{TT}_4}^{\mathrm{I}}}\atop{{\mu\mathrm{g}/100}}$ ml	AFT ₄ I n/100 ml	Follow-up
1	Pituitary neoplasm	-21	3.3		6	3.4	1.25	After treatment TT ₄ ^I 9.2,
2	Arteriogram; 60CO ther- apy to pituitary	-22	>50*			3.2	1.5	NaLT4 therapy; BMR -10%
3	0	-27		455		3.5	1.9	Tests after beginning NaLT4 therapy; BMR +2%, cholesterol 340 mg/100 ml
4	0		2.0		13	0.3	1.0	Improved after therapy
5	0		1.9			2.8	1.1	Improved after therapy
6	Gall bladder dye	-21	9.8	240		3.1	1.5	Improved after therapy
7	IV pyelogram	16	>50*	372	1	1.6	0.6	After therapy, symptom free; TT ₄ ^I 7.2, AFT ₄ ^I 3.9
8	Struma lymphomatosa	-4	3.2	310		3.6	1.2	Thyroid antibod. 1:2500 after therapy, PBI 4.5

^{*} Affected by injected dye.

[†] Possibly affected by the estrogen.

Table 5.—Results obtained in patients with possible hypothyroidism

Case	Factors possibly affecting test results	BMR %	PBI µg/ 100 ml	Choles- terol mg/ 100 ml	131 I 24-hr uptake %	Total serum TT ₄ ^I µg/100 ml	AFT ₄ I n/100 ml	Follow-up
1	None	-1		255		4.6	2.5	
2	None	-14	3.7	160		5.9	3.5	
3	Off TT ₄ I, 1 mon.	-2	7.3	226		7.7	2.2	
4	None	+3	5.0		25	4.6	2.2	
5	Depressed	-10		430		6.2	3.1	
	IV pyelogram							
6	Dilantin	+7				4.5	3.3	All patients con-
7	None	+17	4.4	375	7	4.5	2.9	sidered euthy-
8	90Y implant	+7	25		19	6.7	5.5	roid
9	Estrogen	-5	11.7	295		6.3	2.6	
10	None	+5	4.9			5.2	3.6	
11	None			170		5.4	3.0	
12	None		6.1		35	7.4	3.4	
13	Iodine	-14		27 0		4.2	2.1	
14	None	-10	7.8	130		6.2	3.1	
15	Estrogen	-13		205		7.3	3.4	
16	None		5.4			4.6	2.3	
17	None		9.4			5.2	2.7	
18	None		3.6			4.4	2.2	
19	None		3,0	150		4.2	2.3	
20	None			205		4.0	2.1	
21	None	-9	6.7	250		6.0	3.4	
22	None	-3	8.9	190		3.2	1.0	

Table 6.—Results obtained in patients judged to be euthyroid

Case no.	Clinical impression	Drugs	BMR %	PBI µg/100 ml	Choles- terol mg/100 ml	181] 24-hr uptake %	Total serum TT ₄ I µg/100 ml	AFT ₄ I n/100 ml
1	Hyperthyroidism to be excluded	Estrogen		9.5	-		9.6	3.7
2	Euthyroid	Previous tapazole	+7	7.1		39	6.8	2.8
3	Goiter	0	+2	9.6		20	5.0	2.9
4	Goiter	0		6.4	285		6.6	3.9
5	Thyroiditis	Iodine estrogen	8	13.9	240		4.0	1.9
6	90Y implant	0	+22	5.5		43	4.9	3.5
7	Treated Graves' dis-	PTU T3	-13	6.8	210	6	4.0	2.2
8	Treated Graves' dis-	Previous tapazole		5.5		10	3.8	1.9
_	ease	Lugol's solution	••		055	(4 hr)	= 4	0.5
9	Euthyroid	Estrogen	-19	10.3	255		5.4	2.5
10	Euthyroid	0	-6	6.9	255		6.2	
11	ATS euthyroid	Estrogen				29	5.7	3.4
12	Euthyroid	0				26	5.9	3.2
13	Alopecia	0	-3	4.2	225		3.3	1.7
14	ATS		-2	7.3	220		7.7	4.2
15	Exophthalmos	0	+14	5.5	170	30	5.5	3.3
16	ATS	0	-29	4.3			3.8	2.5
17	Goiter	0	+2	9.6		20	5.0	2.8
18	ATS	Estrogen	-6	13.9			4.0	1.9
19	Post 181I therapy	0	+7	8.5			0.8	0.3

Table 7.—Results obtained in euthyroid patients receiving various types of treatment

$ \begin{array}{cccccccccccccccccccccccccccccccccccc$								1,	1,1	
NaLT ₄ None now post ¹³¹	Clinical impression		Drugs	$^{\rm BMR}_{\%}$	PBI μg/100 ml		131 24-hr uptake %	Total serum TT_4^I $\mu g/100$ mi	$\begin{array}{c} {\rm AFT_4^I} \\ {\rm n/100} \\ {\rm ml} \end{array}$	Follow up
None now post ¹³¹	Myxedema		NaLT4	2	Ē			3.5	1.9	Symptom free
T3 -2, +6 1.8, 0.9 1, 2 3.0 1.2 12 Israngen +8	Euthyroid		None now post ¹³¹ I PTU	-13	6.8		9	4.0	2.2	
137 I setrogen +8 9.6 185 8 4.4 4.3 T3 Imm +11 6.5 4.9 292 4.1 1.9 Tby. Gr. II/day 12.5 3 6.5 3.1 pre Rx Tby. Gr. II/day +14 5.5 170 30 5.5 3.3 during Rx NaLT ₄ 5.4 4.0 1.9 NaLT ₄ 5.4 4.0 1.9 Desic. thy. -4 250 15 6.0 3.4 150 mCi -2 7.1 20 3.7 3.2 Tapazole +18 7.1 20 39 6.8 2.8	Thyroid carcinoma	oma	T3	-2, +6	1.8, 0.9		1, 2	3.0	1.2	No advance 4 yr
T3 lmm +11 6.5 4.9 292 4.1 1.9 Urogram Thy. Gr. II/day +14 5.5 170 30 5.5 3.1 pre Rx NaLT ₄ 5.5 170 30 5.5 3.3 during Rx NaLT ₄ 5.5 170 30 5.5 3.3 during Rx NaLT ₄ 5.5 170 30 5.5 3.3 during Rx NaLT ₄ 5.5 170 30 5.5 3.3 during Rx NaLT ₄ 5.5 170 30 5.5 3.3 during Rx NaLT ₄ 5.5 170 30 5.5 3.3 during Rx NaLT ₄ 5.5 170 30 5.5 3.3 during Rx Tapacole +14 5.5 170 30 5.5 3.4 5.3 7.7 3.2 3.7 3.2 4.0 7.1 220 39 6.8 2.8	Goiter (postoperative)	rative)	127I estrogen	+8	9.6	185	80			
Thy. Gr. II/day -5 4.9 292 4.1 1.9 Urogram 12.5 3 6.5 3.1 pre Rx Thy. Gr. II/day +14 5.5 170 30 5.5 3.3 during Rx NaLT4 +14 5.5 170 30 5.5 3.3 NaLT4 5.4 4.0 1.9 Desic. thy. -4 250 15 6.0 3.4 150 mCi -2 7.1 20 3.7 3.2 Tapazole +18 7.1 220 39 6.8 2.8			T3 1mm	+11	6.5			4.4	4.3	
Urogram 12.5 3 6.5 3.1 pre Rx 1 Thy. Gr. II/day 170 30 5.5 3.3 during Rx 1 NaLT4 5.5 170 30 5.5 3.3 during Rx 1 NaLT4 5.4 4.0 1.9 Desic. thy. -4 250 15 6.0 3.4 130 mCi -2 7.1 20 39 6.8 2.8 Tapazole +18 7.1 220 39 6.8 2.8	90Y implant		Thy. Gr. II/day	-5	4.9	292		4.1	1.9	
1. Thy. Gr. II/day 3.3 during Rx 3.3 during Rx 3.4 mult 4.5 5.5 170 30 5.5 3.3 during Rx 3.4 mult 4.0 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9 1.9	Mental retardation	tion	•		12.5		જ	6.5	3.1 pre Rx	
NaLT ₄ +14 5.5 170 30 5.5 3.3 NaLT ₄ 5.4 4.0 1.9 Desic. thy4 250 15 6.0 3.4 150 mCi -2 7.1 220 39 6.8 2.8	skeletal retardation	dation	•						3.3 during Rx	
Desic. thy4 250 15 6.0 3.4 1.9 Desic. thy2 7.1 220 39 6.8 2.8	Treated Graves' dis-	s' dis-	NaLT4	+14	5.5	170	30	5.5	3.3	BMR -3, Chol. 150, PBI
Desic. thy4 250 15 6.0 131 -2 7.1 Tapazole +18 7.1 220 39 6.8	case									F:0
Desic. thy4 250 15 6.0 131 6.2 150 mCi -2 7.1 Tapazole +18 7.1 220 39 6.8	Treated Graves' dis-	s' dis-	NaLT4		5.4			4.0	1.9	
Desic. thy4 250 15 6.0 131	ease									
150 mCi	Hypothyroid		Desic. thy.	4-		250	15	0.9	3.4	
150 mCi -2 7.1 Tapazole +18 7.1 220 39	Myxedema NaLT4	LT,	1181					6.2	3.7	
Tapazole +18 7.1 220 39	0.3 mg/dy		150 mCi	-2	7.1			7.7	3.2	
	Mild hyperthyroid	roid	Tapazole	+18	7.1	220	39	8.9	2.8	