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Identical mutations in unrelated families with generalized resistance to thyroid hormone occur in cytosine-guanine-rich areas of the thyroid hormone receptor beta gene. Analysis of 15 families.

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## Research Article

Generalized resistance to thyroid hormone (GRTH) is a syndrome of variable reduction of tissue responsiveness to thyroid hormone. 28 different point mutations in the human thyroid hormone receptor beta (TR beta) gene have been associated with GRTH. These mutations are clustered in two regions of the T3 binding domain of the TR beta (codons 310-347 and 417-453). We now report point mutations in the TR beta gene of six additional families with GRTH and show that three mutations occurred each in three families with GRTH, and that three other mutations were each present in two families. In 11 of these 15 families, lack of a common ancestor could be confirmed by genetic analysis. 28 of the 38 point mutations so far identified, including all those occurring in more than one family, are located in cytosine-guanine-rich areas of the TR beta gene. Differences in clinical and laboratory findings in unrelated families harboring the same TR beta mutation suggest that genetic variability of other factors modulate the expression of thyroid hormone action.

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# Identical Mutations in Unrelated Families with Generalized Resistance to Thyroid Hormone Occur in Cytosine-Guanine-rich Areas of the Thyroid Hormone Receptor Beta Gene

**Analysis of 15 Families** 

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#### **Abstract**

Generalized resistance to thyroid hormone (GRTH) is a syndrome of variable reduction of tissue responsiveness to thyroid hormone. 28 different point mutations in the human thyroid hormone receptor beta  $(TR\beta)$  gene have been associated with GRTH. These mutations are clustered in two regions of the T<sub>3</sub> binding domain of the TR $\beta$  (codons 310-347 and 417-453). We now report point mutations in the TR\$\beta\$ gene of six additional families with GRTH and show that three mutations occurred each in three families with GRTH, and that three other mutations were each present in two families. In 11 of these 15 families, lack of a common ancestor could be confirmed by genetic analysis. 28 of the 38 point mutations so far identified, including all those occurring in more than one family, are located in cytosine-guanine-rich areas of the TR\$\beta\$ gene. Differences in clinical and laboratory findings in unrelated families harboring the same TR\$\beta\$ mutation suggest that genetic variability of other factors modulate the expression of thyroid hormone action. (J. Clin. Invest. 1993. 91:2408-2415.) Key words: dinucleotide repeat polymorphism • allele specific amplification • thyrotropin • free thyroxine

#### Introduction

Generalized resistance to thyroid hormone  $(GRTH)^1$  is an inherited condition of reduced target tissue responsiveness to thyroid hormone characterized by a nonsuppressed thyrotropin (TSH) in spite of high concentrations of thyroid hormone (1). It has long been suspected that the condition is due to abnormalities at the level of the thyroid hormone receptor (TR) (2, 3). The detection of two genes  $(TR\alpha)$  and  $TR\beta$  located on different chromosomes) that encode nuclear proteins with identi-

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1. Abbreviations used in this paper: FT<sub>4</sub>, free thyroxine; GRTH, generalized resistance to thyroid hormone; TR, thyroid hormone receptor; TSH, thyrotropin; TT<sub>4</sub>, total thyroxine; TT<sub>3</sub>, total triiodothyronine.

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cal thyroid hormone binding properties (4) led to the demonstration of linkage of the  $TR\beta$  gene with GRTH (5). Subsequently, a single basepair substitution in the hormone binding domain of the  $TR\beta$  was described in affected members of a family with GRTH (6). 28 point mutations have been since described in families with GRTH (6–22). With the exception of one mutation (16), all cluster in two regions in the triiodothyronine ( $T_3$ ) binding domain of the  $TR\beta$  (codons 310–347 and 417–453). Codons are numbered according to the corrected coding sequence of the  $TR\beta$  gene, which adds 5 amino acids to the amino terminus (23).

Each family with GRTH up until the present report was believed to have a unique mutation. Based on this latter finding, it was anticipated that differences in functional impairment of the mutant TRs would explain the variable manifestations of GRTH among families. Such relation has been demonstrated in vitro but not in vivo. Mammalian cells transfected with mutant TRs having lower binding affinity for T<sub>3</sub> required higher concentrations of the hormone to produce transactivation of a cotransfected thyroid hormone responsive reporter gene (24, 25). Surprisingly, this correlation was not found in vivo in terms of the severity of clinical manifestations. More specifically, subjects expressing the mutant  $TR\beta$  with fivefoldreduced T<sub>3</sub>-binding affinity had more severe impairment of mental function and growth than individuals from another family expressing a mutant TR with more than 100-fold-reduced T<sub>3</sub>-binding affinity (6, 7, 26). This discrepancy was present despite similar levels of endogenous thyroid hormone and the presence of unaltered expression of both mutant and normal TR $\beta$  genes in the heterozygous affected subjects (27).

Sequence analysis of the  $TR\beta$  gene in more families with GRTH has led to the recognition of identical mutations in apparently unrelated families. We now report identical nucleotide substitution in the  $TR\beta$  gene occurring in affected members of different families with GRTH. Three of these mutations were each found in three families and three others, each in two families. In 11 of the 15 families, lack of a common ancestor was confirmed by genetic analysis. The mutations in six of the families have not been previously reported. All identical mutations occurred in cytosine(C)-guanine(G)-rich regions of the  $TR\beta$  gene. Differences in clinical and laboratory findings in unrelated families harboring the same  $TR\beta$  mutation suggest that additional factors modulate the expression of the thyroid hormone resistance.

# Methods

Reagents. Providers of biological reagents, including modifying and restriction enzymes, were Sigma Immunochemicals (St. Louis, MO), GIBCO BRL (Gaithersburg, MD), and New England BioLabs, Inc.

(Beverly, MA). Sequenase version 2.0 DNA sequencing kits were purchased from U.S. Biochem. Corp. (Cleveland, OH). [ $\alpha$ -35S]dCTP was obtained from Amersham Corp. (Arlington Heights, IL). Synthetic oligonucleotides were synthesized by  $\beta$ -cyanoethyl chemistry using a DNA synthesizer (308B; Applied Biosystems, Inc., Foster City, CA).

Subjects. 61 affected subjects belonging to 15 families are included in the study. Data from thyroid function tests and, when applicable, references to the first description of the families appear in Table I. All subjects had inappropriately normal TSH with elevated thyroid hormones, indicating reduced pituitary sensitivity to thyroid hormone. They had no symptoms or signs of thyrotoxicosis. The majority of subjects had goiters, and those tested failed to respond normally to the administration of supraphysiologic doses of  $T_3$  or  $T_4$ .

DNA sequencing. DNA was isolated from peripheral blood leukocytes and/or from cultured skin fibroblasts (33). It was used for subsequent analyses and for  $TR\beta$  gene sequencing of at least one affected member of each family. Exons 6, 7, and 8, comprising 75% of the  $T_3$ -binding region of the  $TR\beta$ , were separately amplified using the PCR (35). For details on the structure of the  $TR\beta$  gene, see Takada et al. (34). The conditions of the reactions and the sequences of the oligonucleotides were those published by Usala et al. (36). The amplified DNA fragments were cloned into the M13 bacteriophage vector and sequenced by the dideoxynucleotide chain termination method (37) using Sequenase.

Confirmation of mutations. Several techniques were used to determine the presence or absence of each mutation in family members. Allele-specific amplification (6) was used to confirm the mutations in families F52 and F89. The identification of families (F followed by a number) is according to the tabulation of all published families with GRTH (38). DNA from each individual was subjected to two PCRs, one with a pair of oligonucleotide primers, 5'-GGAGAT-CATGTCCCTTCGCG-3' ("normal primer") and 5'-AAAGCT-CTTTGGATCCCCACTAACGAG-3' (primer α) and the other with a pair of primers, one of which contained the mutant nucleotide (5'-GGAGATCATGTCCCTTCGCA-3', mutant nucleotide underlined).

Allele-specific amplification was not successful in confirming all mutations due to nonspecific amplification. Furthermore, not every mutation resulted in the creation or loss of a restriction endonuclease site. An alternative and novel approach to the confirmation of mutations was the synthesis of degenerate oligonucleotide primers complementary to sequences near the mutated nucleotide that produced a novel specific restriction site only if the DNA template contained the expected nucleotide substitution. Following amplification of the subjects' DNA by PCR, the DNA fragments were submitted to electrophoresis in acrylamide gel or in 3% NuSieve/1% agarose gel, before and after digestion with the specific enzyme. Partial cleavage of a DNA fragment indicated that the mutant nucleotide was present in one of the two alleles. A 20mer oligonucleotide primer was used for confirming the mutation in family F88 resulting in a new DraIII site (degenerated nucleotides are underlined): 5'ATGTCCCTTCGCGCACTGT-3' and in families F29 and F106 resulting in a new Ncol site: 5'ATGGGGAAATGGCAGTGCCA-3'. The antisense oligonucleotide primer used in these reactions was the same (5'-AAAGCTCG-GATCCCCACTAACGAG-3'). In family F95, however, the mutation itself resulted in a new restriction site (ApaLI), and the primers used were the same as for amplifying the exon for sequencing.

Confirmation of genetic unrelatedness by dinucleotide repeat polymorphism. The presence of dinucleotide (CA) repeat polymorphism in the TR $\beta$  intron flanked by exons 2 and 3 (39) was used to determine whether members from different families harboring identical TR $\beta$  mutations had a common ancestor. A 197–209 bp fragment containing the region of variable lengths of CA repeats was amplified by PCR using oligonucleotide primers labeled at their 5' end with <sup>35</sup>P by the kinase reaction. The resulting products were resolved on a polyacrylamide sequencing gel as described by Sakurai et al. (39).

Tests of thyroid function. Sera were available from individuals belonging to families F29, F45, F52, F88, F89, F95, and F106. Total thyroxine (TT<sub>4</sub>) and triiodothyronine (TT<sub>3</sub>) were measured by radio-immunoassays (Diagnostic Products Corp., Los Angeles, CA) and TSH by ELISA-2 (Nichols, San Juan Capastrano, CA). Free thyroxine

Table I. Hormonal Data and Family Identification\*

	Family ID <sup>‡</sup>	T <sub>3</sub> -Affinity of TR $oldsymbol{eta}^{oldsymbol{s}}$	TT <sub>4</sub>	TT,	TSH	FT <sub>4</sub>	$N_{\parallel}$	Reference
			(nmol/liter)	(nmol/liter)	(mU/liter)	(%)		
M1 <sup>1</sup>	F100 (E-D)**	0.2	15	9.9	4.7	_	1 (1)	8
	F89 (XVIII)**	_	360	3.1	4.2	_	1 (1)	9##
	F52 (XIII)	_	297±4	5.3±0.0	$3.9 \pm 0.1$	292	2 (2)	28,29##
M2	F54 (WR)	0.49±0.10	196±29	3.7±0.5	$3.2 \pm 1.4$	120±23	13 (14)	18,30
	F88 (XIX)	_	247±19	$3.3 \pm 0.7$	3.3±0.6	182±31	3 (3)	9,29##
M3	F67 (K-Cl)	0.51±0.29	192±24	4.3±0.4	$2.2 \pm 1.3$	137±13	6 (7)	12
	F95 (XVII)	_	186±22	$3.2 \pm 0.4$	$2.5 \pm 1.1$	118±29	8 (8)	29##
M4	F29 (V)	_	283±21	$5.2 \pm 1.2$	$2.8 \pm 1.9$	213±19	3 (3)	31##
	F106	_	241±35	4.3±0.8	$3.6 \pm 2.6$	200±13	3 (4)	##
	F110 (K-T)	0.21	268	4.6	4.6	296	1(1)	17
M5	F45 (XII)	0.14±0.21	216±41	5.1±1.1	3.8±2.9	216±80	2 (2)	9,32
	F68	_	_	_	1.3±0.7	181±16	3 (4)	10
	F111	0.2					0 (2)	21
M6	F85	$0.46 \pm 0.02$	237±3	$3.9 \pm 0.2$	1.7±0.9	183±39	3 (3)	15
	F105 (Q-W)	0.41	196	3.5	1.9	194	1 (6)	8

<sup>\*</sup> Data for TT<sub>4</sub>, TT<sub>3</sub>, and TSH are expressed as mean±SD, and for FT<sub>4</sub> as percent of the upper limit of normal. Except for F100 and F89, all values are from untreated patients.

<sup>&</sup>lt;sup>‡</sup> Family identification according to chronology of publication or presentation in our clinic. In parentheses is the ID used in the respective reference.

<sup>§</sup> Expressed as the ratio of the mutant-to-normal association constants of the TRs expressed in vitro.

Number of affected subjects for whom hormonal data was available. Total number of affected individuals in each family are in parentheses.

<sup>&</sup>lt;sup>1</sup> M1-M6 refer to the 6 different TR  $\beta$  gene mutations in this report.

<sup>\*\*</sup> Treated with T3 (F100) and T4 (F89) because of previous thyroid gland ablation. # Subjects or mutations not previously published.

(FT4) was estimated from the free  $T_4$  index utilizing a resin  $T_4$  uptake test (40). Results of affected subjects from each family were expressed as mean $\pm$ SD. Published data from families F54, F67, F68, F85, F105, and F110 were used and normalized using the corresponding upper limits of normal. Statistical significance was determined by analysis of variance and by unpaired t tests. P values  $\geq 0.05$  were considered not to be statistically significant.

### Results

Six distinct mutations were found in 15 families, each present in at least two families (Table II). Seven of these families were studied in our laboratory, six of which have not been previously reported (see Table I). In the latter families, the presence of a nucleotide substitution in the  $TR\beta$  gene of all subjects affected by GRTH but not in normal relatives was confirmed. Allele-specific amplification was used to demonstrate the substitution of the normal G-1234 by an A (M1) in affected subjects of families F52 and F89. As shown in Fig. 1, this mutation was absent in the nonaffected natural parents of the 9-yr-old twin boys of family F52, indicating a de novo mutation. An identical mutation was found by sequencing and confirmed by the allele-specific amplification in a woman with GRTH (F89) who was 31 yr old at the time of diagnosis. This patient was adopted and no other family members were available for testing.

The creation of a new ApaLI restriction site by the substitution of G-1244 by A (M3) was used to confirm the mutation in all affected members of family F95. An alternative approach, that of the loss of the recognition site for HhaI, was used by Cugini et al. (12) to identify the identical mutation in another family (F67). Confirmation that two other mutations were present only in subjects affected by GRTH necessitated the use of a degenerate oligonucleotide primer (see Methods), which together with the mutated nucleotide created a specific restriction site. This is illustrated in Fig. 2 for the detection of the substituted C-1243 for T (M2) in family F88, creating a new recognition site for DraIII. The same strategy was used to confirm the presence of specific nucleotide substitutions in the

Table II. Sites of Identical Mutations in HTR $\beta$  Gene of Families with GRTH

		Nucleotide number	Codon			
Mutation	Family ID		Number	Normal	Mutant	
M1	F52	1234	317	GCT (Ala)	ACT (Thr)	
	F89					
	F100					
M2	F54	1243	320	CGC (Arg)	TGC (Cys)	
	F88					
M3	F67	1244	320	CGC (Arg)	CAC (His)	
	F95					
M4	F29	1297	338	CGG (Arg)	TGG (Trp)	
	F106					
	F110					
M5	F45	1598	438	CGC (Arg)	CAC (His)	
	F68					
	F111					
<b>M</b> 6	F85	1642	453	CCT (Pro)	ACT (Thr)	
	F105			. ,	, ,	

A

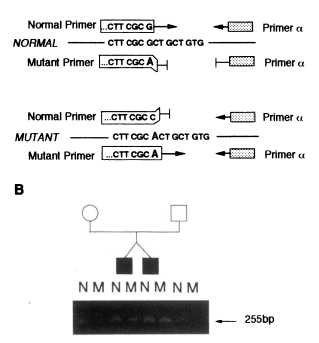


Figure 1. Allele-specific amplification confirming the replacement of G-1234 by an A in codon 317 of the TR $\beta$  gene in family F52. (A) Oligonucleotide primers were complementary to the normal and mutant allele with the last nucleotide at the 3' end aligned with the mutated nucleotide. DNA from each individual was subjected to two PCR reactions, one with the normal primer and one with the mutant primer. Primer  $\alpha$  was common for both reactions. A normal allele would only amplify with the normal primer and a mutant allele would only amplify with the mutant primer. (B) Pedigree of F52 showing that the affected twin boys (black symbols) had both a normal (N) and a mutant (M) allele confirming their heterozygous state. Both alleles of the unaffected parents (open symbols) were normal.

 $TR\beta$  gene of families F29 and F106. In these two families with an identical mutation, C-1297 replaced by T (M4), DNA amplified with the degenerate oligonucleotide primer (see Methods) created a recognition site for NcoI only in the presence of the mutant nucleotide (data not shown).

Each of the six mutations under consideration (M1 through M6), their location on the  $TR\beta$  gene, and the resulting amino acid replacement are shown in Table II. Mutations M1, M4, and M5 were each detected in three families (a total of nine families), while mutations M2, M3, and M6 were each found in two families (a total of six families). All mutations were clustered in two distinct areas of the  $T_3$ -binding domain of the  $TR\beta$ , nucleotides 1214–1325 and 1598–1661. It is of interest to note that all mutations occurring in more than one family involved guanines or cytosines substituted for adenines or thymidine.

Since it is possible for two presumably unrelated families to have a common ancestor, we undertook to demonstrate by genetic means that identical mutations in the  $TR\beta$  gene occurred independently. Analysis of families harboring mutations M3 (F67 and F95) and M4 (F29 and F106) relied on CA repeat polymorphisms known to occur in the intron flanked by exons 2 and 3 of the  $TR\beta$  gene (39). Results from such analysis of two families harboring an identical nucleotide substitution

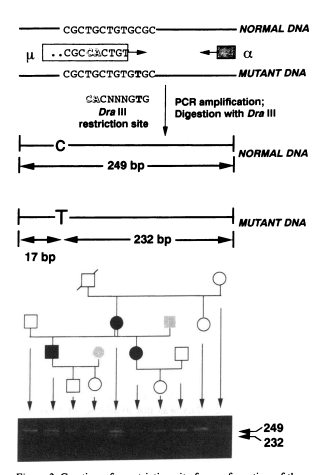


Figure 2. Creation of a restriction site for confirmation of the replacement of C-1243 by a T in codon 320 of the  $TR\beta$  gene in family F88. The mutation M2 in F88 did not create or delete a site for a restriction endonuclease. Therefore, a degenerate oligonucleotide primer  $(\mu)$  was synthesized that would change the sequence of the PCR product such that a restriction site was formed only if the template DNA had the specific base substitution M2. Amplification using primer  $\mu$  changed the TG in the patients' DNA to CA (outlined). The degenerate primer amplifies both normal and mutant alleles and does not overlap the mutation site. However, the presence of the mutant nucleotide, T, results in creation of a DraIII site. When the PCR product from each individual was subjected to DraIII digestion, a 249-bp (uncut) fragment was visualized by 3% NuSieve/1% agarose electrophoresis in unaffected subjects (open symbols). The presence of a 249 bp (uncut) and 232 bp (cut) fragments in affected subjects (black symbols) confirmed the presence of a mutation in one of their two  $TR\beta$  alleles. The 17-bp fragment was too small to be visualized.

in the coding region of the  $TR\beta$  gene are illustrated in Fig. 3. The mutant M4 allele present in affected subjects of family F29 contained the polymorphic band 4, while the M4 allele of family F106 contained band 6. These results confirm that the mutation M4 occurred independently in the two families. Furthermore, M4 was recently identified in an individual with GRTH (F110) in whom this mutation appeared de novo (17). Therefore, F110 is also unrelated to F29 and F106. DNA samples from family F67 (12), kindly provided by Dr. Stephen J. Usala, showed that the polymorphic band 7 was present in the allele containing mutation M3, while the identical mutation in family F95 had band 4. We previously found a silent mutation (T-1535 replaced by a C), preserving amino acid Phe-417, on the same allele and in the same exon as M5 in family F45 (9).

The allele harboring the M5 mutation in family F68 (10) maintained the wild type T-1535 (C. V. Boothroyd, personal communication) confirming that mutation M5 in the two families also occurred independently. M5, recently detected in another family (F111), occurred de novo and was transmitted to a single affected child (21), indicating that this family with GRTH is not related to F45 and F68.

Evidence for the de novo occurrence of mutation M1 in families F52 and F100 is based on the observation that the parents of families F52 (Fig. 1) and F100 (8) were not affected. Since affected members of these families were 11 yr old or younger and thus not in the reproductive age, the identical mutation found in the adopted subject of family F89 has occurred independently of the mutations in families F52 and F100. The criteria used to determine the genetic relatedness in all families with identical mutations in the  $T_3$ -binding domain of the  $TR\beta$  gene are shown in Table III.

Because clinical observations suggested that differences in the manifestations of GRTH existed not only among families with different mutations but also between families with the same mutation, we attempted to establish the presence of heterogeneity by more objective means. Since mean serum TSH levels were normal and not significantly different among the 10 families for whom sufficient data were available, we examined the corresponding mean serum  $T_4$  and  $T_3$  concentrations re-

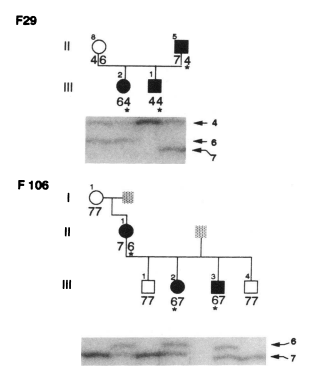


Figure 3. Determination of CA repeats polymorphism in the exclusion of founder's effect for the identical  $TR\beta$  gene mutation (M4) in families F29 and F106. The black symbols represent affected individuals harboring the M4 mutation, the open symbols, unaffected individuals devoid of the M4 mutation, and the gray symbols, spouses that were not tested. Three different alleles were identified in F29, represented by bands 4, 6, and 7, and the 4 allele harbored the M4 mutation (asterisk). In contrast, the M4 mutation in family 106 was contained in the allele having the 6 CA-repeat pattern (asterisk). The roman numerals identify the generation and the number above each symbol the subject.

Table III. Criteria Used to Determine Genetic Unrelatedness among Families with Identical TR\$ Mutations

Mutation	Family	Criteria
MI	F52 and F100	De novo mutations (both parents unaffected)
	F89	Adopted
M2	F54*	Scottish descent, living in USA
	F88	English/Irish/German/Amerindian descent, living in USA
M3	F67	Mutant allele had CA-repeats band "7"
	F95	Mutant allele had CA-repeats band "4"
M4	F29	Mutant allele had CA-repeats band "4"
	F106	Mutant allele had CA-repeats band "6"
	F110*	De novo mutation
M5	F45	Silent substitution C-1535 (Phe-417) <sup>‡</sup>
	F68	Wild type T-1535 (Phe-417)
	F111*	De novo mutation
M6	F85	Japanese descent
	F105	Caucasian

<sup>\*</sup> DNA not available.

quired to maintain comparable TSH levels. One pair of families (F54 and F88) with mutation M2 had significantly different mean  $TT_4$  values (P < 0.02), and another family pair (F67 and F95) with mutation M3 showed significant difference (P < 0.02) in their mean TT<sub>3</sub> concentration (Table I). Free T<sub>4</sub> level rather than TT<sub>4</sub>, TT<sub>3</sub>, or free T<sub>3</sub> correlates best with the feedback regulation of TSH and is a measure of pituitary sensitivity to thyroid hormone (41, 42). We therefore compared the mean FT4 values among the families. Values were expressed as percent above the upper limit of normal in order to reduce the error in comparing data obtained by different laboratories. As shown in Fig. 4, significant differences were observed in free T<sub>4</sub> levels between families with mutation M3 and those with M4 and M5. More surprisingly, a highly significant difference (P < 0.02) in mean free T<sub>4</sub> values was also found between families F54 and F88, both harboring the same M2 mutation.

#### **Discussion**

A previous search for mutations in the  $TR\beta$  gene identical to those found in two patients with GRTH indicated that mutations were probably unique to affected members of each unrelated family (29). Subsequent detection of  $TR\beta$  mutations in additional families with GRTH and in particular two studies

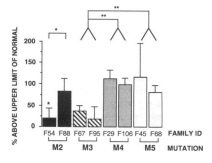


Figure 4. FT<sub>4</sub> levels in families with GRTH. Results are expressed as mean percent above the upper limit of normal $\pm$ SD. \*P = 0.002; \*\*P < 0.01. No data was available for comparison of families with M1 and M6.

each reporting seven and nine families, respectively (8, 9), also failed to show identical mutations in different families. We currently report six different mutations each present in 2 or 3 of 15 unrelated families. Obviously the need to confirm that the families with identical mutations have no common ancestor is not trivial. A physician may have sent patient material to more than one laboratory or the common ancestor may be distant and unknown to the families in question. Indeed, the mutation in the  $TR\beta$  gene in one individual was reported by Parilla et al. (8) as case 1 of family F-W, and by us as subject VII-1 (9), independently. This subject with GRTH was first reported by Kaplowitz et al. (43) and then studied by us (29, 44, 45). After subsequent adoption, he moved to another state and came to the attention of Parilla et al. (8).

The criteria used to determine the genetic relation between families harboring identical mutations are listed in Table III. In 11 of the 15 families harboring four of the six mutations (M1, M3, M4, and M5), it was possible to confirm by direct gene analysis that the mutations occurred independently. In two families with M1 and one family each with M4 and M5 the mutations occurred de novo. In seven families (one with mutation M1 and two each with mutations M3, M4, and M5), although the lineage could not be traced back far enough to date the occurrence of their mutations, their independent development could be confirmed. The independent occurrence of mutation M2 and M6, each in two families, could not be assessed by genetic means since DNA was not available for analysis. Their different ethnic origins is suggestive but does not exclude a common ancestor.

37 of the 38 mutations so far reported, including the 6 described herein, cluster in the two regions of the T<sub>3</sub>-binding domain of the  $TR\beta$  previously identified (8, 9). Their nature and exact location in the  $TR\beta$  gene are depicted in Fig. 5. 71% of these mutations occur in 9 of the 16 CG-rich segments, with 4 or more Cs or Gs, located in a 448 bp segment of the  $TR\beta$ coding sequence. Furthermore, of the six mutations found to occur independently in 15 unrelated families, five (M1, M2, M3, M4, and M5) were in regions of CG repeats and one (M6) in an area of C repeats. This is significant because such sequences are thought to be mutational hot spots in humans since they are frequent sites of polymorphisms (46). Regions with the dinucleotide CG in the Factor IX gene had a 77-fold increase in the mutation rate compared with the rates of other regions (47). In a review of single base-pair substitutions that cause human genetic diseases (excluding hemophilia), 32% of point mutations were CG to TG or CG to CA transitions (48). This represents a 12-fold higher frequency than that predicted to occur at random. The mechanism of this phenomenon appears to be methylation of cytosine 5' to guanine and the subsequent spontaneous deamination of 5-methylcytosine to thymine, which is not excised by the enzyme DNA-uracil glycosidase (49). GRTH, therefore, seems similar to other genetic diseases in humans in the frequency of this type of mutation, which was present in 14 of the 38 reported mutations (37%) with 27 of 38 (71%) occurring in CG-rich regions. Of the 34 point mutations recently described in the human androgen receptor (50-54), 15 were T and A transitions occurring in CGdinucleotides (53%) but only 6 (21%) occurred in CG-rich areas as defined above.

It is of interest that mutations have not been described between nucleotides 1326 and 1588 though this DNA segment

<sup>&</sup>lt;sup>‡</sup> Same allele and exon as mutation M5.

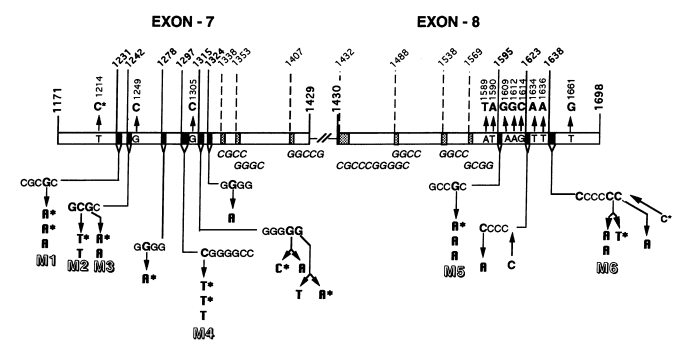


Figure 5. Summary of reported mutations in  $TR\beta$  gene of subjects with GRTH. Mutations analyzed in this report are M1 through M6. \*Mutations studied in our laboratory. CG-rich regions, defined as four or more adjacent Cs and Gs, are indicated in black and gray boxes. The black boxes are those areas where mutations have been identified, and the gray boxes are areas without known mutations. The first nucleotide of each CG repeat sequence is identified. The position of the mutated nucleotides outside the CG-rich areas are also indicated above.

contains CG rich sequences. This region of the  $TR\beta$  gene encodes the heptad repeats forming a "leucine zipper" structure that is involved in receptor dimerization (55–57). Failure to detect individuals with mutations in this region can be explained by the inability of the mutant  $TR\beta$  to exert a "dominant negative effect" through formation of stable mutant  $TR\beta$  homodimers (58), thus escaping clinical recognition. This hypothesis is consistent with the absence of clinical or laboratory abnormalities in heterozygous patients with complete deletion of the  $TR\beta$  gene (34).

The recent observation that the degree of thyroid hormone resistance, as manifested by the clinical severity of GRTH, does not necessarily correlate with the degree of T<sub>3</sub>-binding impairment of mutant receptors (26, 27) is supported by the current data. In contrast to former studies, we had the opportunity to compare the free T<sub>4</sub> levels required to maintain the same TSH value in four pairs of families, each with identical mutations of the TR $\beta$  gene. In one pair of families, F54 and F88, the mean free  $T_4$  concentration was significantly different at P < 0.02(Fig. 4). Although differences in other clinical characteristics, such as hyperactivity and bone maturation, were also observed between families with identical point mutations, they were difficult to quantify because of differences in subjects' ages. This is not to say that the degree of impairment of T<sub>3</sub>-binding does not contribute to the severity of the hormone resistance. It should be noted that the two families with mutation M3 had significantly lower free T<sub>4</sub> values than families with mutation M5, as well as three-fold higher affinity for T<sub>3</sub> of their M3-mutant  $TR\beta$  (Table I, Fig. 5).

The possibility that the observed biochemical and clinical differences among families harboring identical mutations in the  $T_3$ -binding domain of the  $TR\beta$  may be due to a second

mutation in the TR $\beta$  deserves serious consideration. Such mutations have not been detected in four families with GRTH in whom the entire coding region of the  $TR\beta$  gene has been sequenced (6, 9, 15, 16) and in two families in whom sequencing comprised the T<sub>3</sub>-binding, DNA-binding, and hinge regions (12, 13). In addition, screening for mutations upstream of the T<sub>3</sub>-binding domain, including 75% of the DNA binding region, in six of the families described herein (F25, F45, F52, F88, F89, and F95) also failed to show an additional mutation (9). Furthermore, it can be speculated that functionally important mutations in the DNA-binding domain are less likely to have a significant biological effect in the heterozygous state since, as shown by site-directed mutagenesis (59), such mutations have no dominant negative effect. Thus, the most likely explanation for the apparent  $TR\beta$  mutation independent differences in the degree of thyroid hormone resistance among families is the genetic variability of other factors that are involved in the expression of thyroid hormone action. Current evidence indicates that another thyroid hormone receptor (TR $\alpha$ , located on chromosome 17), a number of transactivation protein cofactors (RXR $\alpha$  and RXR $\beta$ ), and less well-defined auxiliary TR proteins interact in the expression of thyroid hormone action (56, 60, 61). The interaction results in the formation of stable complexes that bind to specific DNA sequences (thyroid hormone response elements) present in genes regulated by thyroid hormone. Undoubtedly, each of the involved factors and elements is subject to genetic polymorphism, which modulates its structure, interaction, and function. The genetically determined form and expression of a particular coregulator may diminish or augment the hormonal resistance imparted by a particular mutation in the  $TR\beta$  gene and thus contribute to the modulation of the phenotype of GRTH.

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