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A mutation in \textit{Tpst2} encoding tyrosylprotein sulfotransferase causes dwarfism associated with hypothyroidism

Abbreviated title: \textit{Tpst2} mutation causes hypothyroidism

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The abbreviations used are: cM, centimorgan cM; CCK, cholecystokinin; CF-8, coagulation factor VIII; FSHR, FSH receptor; GP1BA, glycoprotein 1b alpha polypeptide; \textit{grt}, growth-retarded; HE, haematoxylin and eosin; RFLP, restriction fragment length polymorphism; PAPS, 3'-phosphoadenosine 5'-phosphosulfate; PAS, periodic acid schiff; PSGL1, P-selectin glycoprotein ligand-1; TPST, tyrosylprotein sulfotransferases; TSHR, TSH receptor; WT, wild-type.

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ABSTRACT

The growth-retarded (grt) mouse has an autosomal recessive, fetal-onset, severe thyroid hypoplasia related to TSH hyporesponsiveness. Through genetic mapping and complementation experiments, we show that grt is a missense mutation of a highly conserved region of the tyrosylprotein sulfotransferase 2 (Tpst2) gene, encoding one of the two Tpst genes implicated in post-translational tyrosine O-sulfation. We present evidence that the grt mutation leads to a loss of TPST2 activity, and TPST2 isoform has a high degree of substrate preference for TSH receptor (TSHR). The expression of TPST2 can restore TSH-TSHR-mediated cAMP production in fibroblasts derived from grt mice. Therefore, we propose that the tyrosine sulfation of TSHR by TPST2 is crucial for TSH signaling and resultant thyroid gland function.

INTRODUCTION

Normal thyroid function is essential for development, growth, and metabolic homeostasis. Permanent congenital hypothyroidism affects about 1:3000-4000 newborns, and is one of the most common preventable causes of mental retardation. In about 90% of all cases, congenital hypothyroidism is the consequence of thyroid dysgenesis such as a small and sublingual thyroid, or no thyroid tissue. Most of these cases appear sporadically, although a few cases of recurring familial thyroid dysgenesis have been described (1). Molecular genetic analyses have identified four thyroid dysgenesis susceptibility genes in humans, TSHR (2) and the genes for transcription factors TTF1 (3, 4), TTF2 (5) and PAX8 (6) in pathways crucial for the normal development of the thyroid. Studies of the spontaneous mutation and targeted-disruption of Tshr (7), Ttf1 (8), Ttf2 (9), and Pax8 (10) in mice have provided insight into the molecular mechanisms of organogenesis and
thereby form the basis for molecular genetic studies in human patients affected by thyroid
dysgenesis. However, mutations in these genes are found in only 5% of patients with
thyroid dysgenesis. Therefore, the genetic and pathological mechanisms underlying
thyroid dysgenesis are still poorly understood.

The growth-retarded (grt) mouse is a spontaneous mutant exhibiting severe
primary hypothyroidism and dwarfism controlled by a single recessive locus. The gene
responsible for the grt has not yet been cloned, nor has its molecular mode of action been
determined. However, homozygous grt mice exhibit low concentrations of serum T3 and
T4, and a compensatory elevation in the level of circulating TSH demonstrates a normal
pituitary response (11). In fact, measurements of several additional pituitary hormones
revealed no significant differences between normal and grt mice. Growth retardation is
recovered by the administration of T3, indicating that the grt mutation does not affect
anterior pituitary function (12). The thyroid gland of grt mice is defective in TSH
responsiveness, particularly in signaling pathway involving TSH, TSHR, G protein, and
adenylate cyclase (13). These results suggest that the growth-retarded phenotype is due to
an impairment of thyroid glands in the production of thyroid hormone or a response to
TSH. To determine the molecular mechanism of the grt phenotype, we mapped the grt
locus responsible for dwarfism in a 59-centimorgan (cM) region of mouse chromosome 5
(14). No genes responsible for human and rodent dwarfism or thyroid disease have been
mapped to this region. In this report, we narrowed the grt locus to a < 0.1 cM region using
1,084 backcross progenies, sequenced candidate genes located in this region, and finally
identified a causative mutation in the tyrosylprotein sulfotransferase 2 (Tpst2) gene.
Recent papers reported that the posttranslational modification by tyrosine sulfation
regulates many important protein-protein interactions and modulates binding affinity and
specificity. Mutational analysis has demonstrated that TSHR is sulfated on tyrosines in a
motif that is essential for high-affinity binding of TSH and activation of the receptor (15).
In this report, we propose that severe thyroid hypogenesis and consequent dwarfism are due to an impairment of tyrosine sulfation in TSHR by TPST2.

RESULTS

Identification of Tpst2 mutation in grt mutants

To identify the gene underlying the grt mutant phenotype, we created a high-resolution genetic map of the grt locus using 1,084 backcross mice, and analyzed their phenotypes of homozygotes (grt/grt) or heterozygotes (grt/+). To identify the gene underlying the grt mutant phenotype, we created a high-resolution genetic map of the grt locus using 1,084 backcross mice, and analyzed their phenotypes of homozygotes (grt/grt) or heterozygotes (grt/+). By weighing mice at 5 wk of age, homozygotes of both sexes were small with shortened limbs and tails and weighed approximately 40-60% less of the heterozygotes or wild-type (WT) mice. None of the phenotypes of 1,084 backcross progenies had recombination with a microsatellite marker, D5Mit24, indicating a linkage of < 0.1 cM (Fig. 1A). As shown in Fig. 1B, the gene order was from Cryba4 to Asphd2 covering the critical region. In this region, the mouse genome database disclosed seven genes, Cryba4, Crybb1, Tpst2, Tfip11, Srr1, Hps4 and Asphd2. We examined the expression of these possible candidate genes in multiple tissues including thyroid with RT-PCR; there was no qualitative difference observed (data not shown). We next compared the nucleotide sequences of these genes between normal and affected mice, and identified a distinct mutation in Tpst2. The mutation is a single missense mutation with a transversion of C at nucleotide 798 to G, leading to the replacement of a highly conserved histidine with a glutamine at codon 266 in the sulfotransferase domain (Fig. 1B). Sequence alignment of TPST1 and 2 reveals that the histidine residue was highly conserved among TPST family proteins of diverse species (Fig. 2A). In addition, phenotypically affected mice were homozygous for mutant alleles with PCR-restriction fragment length polymorphism (RFLP) genotyping assay,
whereas normal mice were homozygous for wild type alleles or heterozygous in DW/J mouse colony (Fig. 2B). Furthermore, this mutation was not present in ten other laboratory strains (data not shown). These results suggested that the Tpst2 mutation is responsible for the dwarfism.

**Expression of Tpst2 transgene rescues the mutant phenotype**

To confirm this hypothesis, we attempted to complement the grt phenotype using transgenic rescue. We produced transgenic mice carrying Tpst2 cDNA driven by the cytomegalovirus enhancer and chicken actin (CAG) promoter (Fig. 3A). Founder mice expressed the Tpst2 gene in several tissues including thyroid (Fig. 3B). Crosses between grt/+ males with the transgene and grt/+ females generated 17 phenotypically affected mice, all of which were negative for the transgene. Twenty normal mice were homozygous for the grt mutation, and all were positive for the transgene, demonstrating phenotypic rescue of the dwarfism (Fig. 3C and D). Tpst2 overexpression had no influence on growth except in grt/grt mice. Two independent founder lines produced similar results. Next, we investigated the rescue of thyroid function, serum T3, T4, and TSH levels were analyzed in grt/grt mice with or without transgenes. In grt/grt mice, serum levels of TSH were elevated 3 to 5-fold above normal, whereas T3 and T4 levels were less than normal. Rescued mutants demonstrated normal serum levels for T3, T4, and TSH (Fig. 3E). Histologically, the mutant thyroid glands showed hypogenesis characterized by poorly developed follicles that were heterogeneous in size with reduced levels of colloid (Fig. 4A middle, some follicles were unreactive with periodic acid schiff (PAS) staining. Interestingly, the thyroids of rescued mutants revealed no abnormalities in thyroid size, follicular structure, or the morphology of cells (Fig. 4A, bottom). The numbers of cells and follicles per square millimeter were in the normal range, and follicle
size was not significantly different from that of normal mice (data not shown).

Next, to assess thyroid-specific gene expression in grt/grt mice, we performed a semiquantitive RT-PCR analysis. TSH-TSHR signaling is essential for the proliferation and maintenance of the differentiated function of the thyroid follicular cells, but it is not required for early organogenesis and migration of the thyroid anlage (16). TSH-TSHR signaling is essential for the expression of sodium-iodide symporter (Nis), thyroperoxidase (Tpo) and Tshr itself, which are required for thyroid hormone biosynthesis, whereas it is not required for thyroglobulin (Tgn) expression (17). These observations are consistent with our results that the expression of Tshr and Nis is reduced by 3-5-fold, although Tgn was normally expressed in grt/grt thyroids compared with normal controls (Fig. 4B). Previously we have shown that much less iodine is accumulated in the grt/grt thyroid than in the control mice, although the incorporation of iodine into thyroglobulin, glycosylation, and the intracellular transportation of thyroglobulin are all normal (18). The reduction in the amount of iodine accumulated in the grt/grt thyroid might be the major cause of the decrease in levels of thyroid hormones.

Consistent with the results of histopathology and serum hormone levels, the restoration of Tpst2 rescued the levels of expression of Tshr and Nis in the transgenic mutant’s thyroids (Fig. 4B). Thus, on the basis of chromosomal mapping, nucleotide sequence analysis, and genetic complementation study, we concluded that the dwarf-causing grt mutation was contained in the Tpst2 gene.

Loss of enzymatic activity of TPST2 H266Q mutation

TPST catalyzes the transfer of a sulfuryl group from the universal sulfation substrate, 3’-phosphoadenosine 5’-phosphosulfate (PAPS), to a tyrosyl residue within acidic motifs of proteins that transit the Golgi (Fig. 5A). To characterize the enzymatic
activity, we prepared a peptide array presenting the 15-mer peptide substrates, glycoprotein 1b alpha polypeptide (GP1BA) (19), coagulation factor VIII (CF-8) (20), cholecystokinin (CCK) (21), FSH receptor (FSHR), TSHR (15) and P-selectin glycoprotein ligand-1 (PSGL1) (22), that are known acceptors for tyrosine sulfation, and detects the incorporation of $^{35}$S-labeled sulfate into the substrates. TPST1, TPST2 and TPST2 (H266Q) expression vectors were transfected into HEK293T. Immunoblot analysis indicated that each protein was expressed at similar level in the transfected cells (Fig. 5B). Although HEK293T cell expressed both human isozymes, which were detected by RT-PCR, there was no detectable sulfation of all peptide substrates when lysates of mock transfected cell were used (Fig. 5C). Thus, the basal TPST activity was undetectable level for this assay. Fig. 5C shows that all peptides were sulfated by wild-type TPST2. In contrast, the enzyme activity of mutant TPST2 (H266Q) was reduced to undetectable levels, although the same amount of protein was present on each reaction. Thus, histidine 266 must be a critical residue for TPST activity. We investigated the subcellular distribution of normal and mutant TPST2 tagged with FLAG and the effects of mutant TPST2 on the survival of transfected cells, no changes were found (data not shown). This mutation might affect the affinity of TPST protein for acceptor proteins/PAPS or the enzymatic activity itself. Further mutagenesis and crystallographic studies will be needed to address this issue.

**The tyrosine sulfation of TSHR by TPST2 is crucial for TSH signaling**

In mammals, tyrosine sulfation of proteins is catalyzed by two isozymes; TPST1 and 2. A question arose as to whether the isozyme, TPST1, compensates for a deficiency of TPST2, since the mouse TPST1 has 65% homology with TPST2 at the amino acid level. Previously, Northern analyses indicated that both Tpsts were...
ubiquitously expressed in various tissues and cell lines (23, 24). However, it has not been
determined whether the two TPSTs are co-expressed on cells or tissues. To evaluate tissue
expression preferences of Tpst isoforms, we examined the expression of the Tpst1 and
Tpst2 genes in various mouse tissues, the rat thyrocyte cell line FRTL-5 (Fig. 6A), murine
primary fibroblasts, NIH3T3 cells (data not shown). Semiquantitative RT-PCR analysis
was performed concomitantly for both genes in the same tube on each sample, and
revealed that the Tpst isoforms co-expressed in the same tissues and cells including
thyrocytes, although there are some differences in tissue preference. Further, RT-PCR
analyses showed that the grt mutation did not affect steady-state transcription of Tpst1 or
Tpst2 itself in any tissues (data not shown).

In vitro studies using synthetic peptide acceptors indicated that the two TPST
isoforms differed in substrate preference (24, 25). Inactivation of murine Tpst1 did not
result in any abnormality of the thyroid (26). Therefore, we investigated the substrate
specificity of TPSTs. As shown in Fig. 5C and D, TPST1 and 2 similarly sulfated GP1BA,
CF-8, CCK and PSGL1. In contrast, the preference of TPST2 was 5-fold that of TPST1
for TSHR peptides, implying that TPST1 cannot compensate for the deficiency of TPST2
activity.

In the thyroid, the TSH-bound TSHR induces coupling of the receptor to the G
protein and produces intracellular cAMP through the activation of adenylate cyclase (27).
The thyroid gland of the grt mutant exhibited a markedly diminished response of TSH
both in vitro and in vivo, although cAMP production was increased following stimulation
of the thyroid glands with nonspecific adenylate cyclase activators such as forskolin (12,
13). To obtain further evidence, we performed an in vitro TSH signaling assay with
primary fibroblasts obtained from grt/grt mutants. The Lentiviruses encoding Tpst1,
Tsst2 and mutant Tpst2 (H266Q) with TSHR cDNA were introduced into grt/grt
fibroblasts, and TSH-induced production of cAMP was determined. The grt/grt
fibroblasts showed an increase in cAMP production when cultured in the presence of forskolin. Stimulation of cells expressing just TSHR (mock) with TSH resulted in a slight increase in cAMP accumulation (Fig. 6B). The TSH-dependent cAMP accumulation in the cells expressing TPST1 or mutant TPST2 was very low or similar to that of mock-transfected cells. In contrast, in cells expressing TPST2, cAMP accumulation was approximately 2-3 folds higher than that in cells expressing the others. These results strongly suggest that only TPST2 can efficiently mediate the TSH-TSHR signal transduction pathway, whereas TPST1 and TPST2 (H266Q) cannot.

**DISCUSSION**

The present study revealed that *grt* mice carried a recessive loss-of-function mutation in *Tpst2* gene. We have demonstrated that TPST2 is required for normal development and function in thyroid. This conclusion is supported by the following evidence: (a) the position of the *Tpst2* gene closely located to *grt* locus and the perfect co-segregation of the mutations with the phenotype. (b) H266 is highly conserved in TPST family among many species. (c) Q266 substitution causes a loss of enzymatic activity. (d) *Tpst2* transgene rescues the mutant phenotypes *in vitro* and *in vivo*. (e) TPST2 shows high specificity for the substrate toward TSHR. Previously site-directed mutagenesis suggested that sulfation of tyrosine 385 of TSHR is required for high-affinity hormone binding and receptor activation by TSH (15). Further, two kinds of *Tshr* mutant mice, spontaneous and targeted mutants, have been reported (16, 17). Both mutants can produce thyroglobulin, however, they show reduced iodine uptake in the thyroid. Similarly, the uptake of iodine is markedly lower in *grt* mice (18). Combining these observations with our data, we propose that sulfation of tyrosine 385 of TSHR by TPST2 is indispensable for the activation of TSH signaling. Since *grt/grt* mice are unable
to fully respond to TSH, they develop hypothyroidism and dwarfism. The growth of 
grt/grt mice is virtually normal until 2 wk after birth, then is suppressed in the pubertal 
period; however, it gradually catch up with those of normal mice after approximately one 
year. This might be because normal rodent chows provide enough thyroid hormone for 
survival.

Recently, it has been reported that Tpst2 (-/-) mice on the 129/Sv background 
were produced by gene targeting (28). Tpst2 (-/-) mice showed growth retardation at 4-5 
wk of age in both sexes, which resembled that in the grt mutants. However, their report 
could not defined the molecular mechanism of the growth retardation. In our report, we 
determined Tpst2 as the gene responsible for the severe thyroid dysgenesis, which relates 
to TSH hyporesponsiveness through positional cloning and transgenic rescue. 
Furthermore, we show that TPST2 has high degree of substrate specificity for TSHR, and 
is essential for TSH-TSHR signaling. Therefore, our report has clarified the crucial role 
of TPST2 for a particular signaling pathway in vivo for the first time. The maximum 
difference in body weight of Tpst2 (-/-) mice at 5 wk was 20% less compared with normal 
littermates. In contrast, DWJ-grt mice lost up to 50%. It has been reported that thyroid 
dysgenesis were displayed in C57BL/6 mice with Pax8 (+/-)/Ttf1 (+/-) and not in 129Sv 
strain (29). Therefore, Tpst2 (-/-) in the 129 background might be resistant to thyroid 
dysgenesis. These might represent a spectrum of different degrees of severity of the same 
underlying molecular defect. In human, most cases of thyroid dysgenesis are sporadic, 
most patients do not display a clear Mendelian transmission, suggesting the existence of 
several genetic factors which could contribute to the disorder (30). Therefore, this strain 
differences should help us to identify modifier gene(s) involved in morphogenesis, 
growth, and differentiation of the thyroid. The other explanation of phenotypic 
differences between Tpst2-null mutants and grt mice could be through the dimerization of 
TPST proteins, because the many sulfotransferases are homo- and/or heterodimer in
solution (31). The physiological significance of dimerization of cytosolic sulfotransferases is not yet clearly defined. However, it has been reported that both TPST1 and TPST2 form homodimer (23, 32). Therefore, if dimerization is essential for the TPST activity, high expression of TPST2 H266Q mutant may result in a significant decrease in TPST1 activity in a dominant-negative manner. This hypothesis is consistent with the result that the dwarfism of grt mice is more severe than that of Tpst2-null mice.

Hypothyroidism in human is associated with a marked delay in sexual maturation and development (33). The grt males also demonstrate the severe decrease in testicular weight and the numbers of Leydig cells until 5-8 wk age, whereas they gradually acquire normal structure and function of the testis, finally become fertile at 3-4 months (our unpublished data). Thus, we could cross F1 females with grt males to obtain backcrossed progenies. Tpst2 (-/-) mutant males are infertile at 10 wk of age, although spermatogenesis and mating are normal (28). This result is in agreement with previous reports showing a decrease in sperm motility in rats with hypothyroidism (34, 35). However, it has not been described whether Tpst2 (-/-) males become fertile after 10 wk or are infertile throughout their life. If Tpst2 knockout is a phenocopy of the grt mutant, male infertility can be due to dwarfism/hypothyroidism and may also gradually acquire fertility after 10 wk of age.

Tyrosine sulfation was estimated to occur in approximately 1% of all tyrosines of the eukaryotic proteome (36). In mammals, approximately 60 proteins have already been identified (37). The known tyrosine-sulfated proteins include certain adhesion molecules, G protein-coupled receptors, coagulation factors, and extra-cellular matrixes, and hormones. In some of these proteins, tyrosine sulfation has been shown to be required for optimal binding, but in many, a functional role of tyrosine sulfation still has been unclear. Thus, although many proteins are possible to be the substrates of TPST2, our current study mainly focuses on TSHR. However, the possibility that the grt phenotype
reflects a defect in other proteins as the substrate of TPST2 in downstream of the TSHR could not be excluded. Tyrosine sulfation might also affect enzymatic activity, protein transportation, localization and lifespan, possibly being important for hemostasis, chemotaxis, inflammation, and development, as well as for viral and cancer pathogenesis (38).

In the past several decades, the causative genes of inherited hypothyroidism have been identified in humans and rodents. These genes encode transcription factors, hormones, and their receptors. The post-translational processing in the thyroid has been considered to have a significant role in their correct function. However, these processes have not been shown to be mainly involved in the pathogenesis of thyroid until our reports. Although a mutation of TPST has not been found in human disease, our data suggest that TPST2 may be one of the causative genes for thyroid dysgenesis of unknown origin. Further pathological examination of grt, Tpsl (−/−) and Tpst2 (−/−) mutants, and further biochemical examination to find a variety of acceptor proteins will provide new insights into the biological function of tyrosine O-sulfation.

MATERIAL AND METHODS

Pedigree material and haplotype analysis

(C57BL/6J × DW/J-grt)F1 females were mated with DW/J-grt males to obtain backcrossed progenies. Linkage analysis was performed using microsatellite markers, D5Mit314 (MGI:100402), D5Mit240 (MGI:93321), D5Mit24 (MGI:93320) and a single nucleotide polymorphism for Crybb1 (MGI:104992). Mice were maintained under the control of a 12-h light-dark cycle. Research was conducted according to the Guidelines for the Care and Use of Laboratory Animals of both Graduate School of Veterinary Medicine, Hokkaido University and Nagoya City University Medical School. The
experimental protocol was approved by the Institutional Animal Care and Use Committee of both Graduate School of Veterinary Medicine, Hokkaido University and Nagoya City University Medical School.

Vector constructions

The murine *Tpst2*-coding regions (accession number: NM_009419) were amplified by PCR to include the native translation initiation sequence of *Tpst2* with primer pair F/R: 

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  ggctggccatgggcctgtcggtgc/tttcacttatcgtcgtcatccttgtaatccgaactcttaggtgtggggaggtgc.
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The *Tpst1*-coding region (accession number: NM_013837) was amplified with primer pair F/R: 

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  acgtgatatccgttgggaagctgaagcaga/tttcacttatcgtcgtcatccttgtaatccactccacttgtgtcgtcgtgg.
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The sequence coding for the FLAG (FL) peptide YKDDDDK was added to their COOH-terminals and the resultant cDNAs were subcloned in pTriEx 1.1 vector (Novagene) to generate pTriEx/*Tpst1*-FL, pTriEx/*Tpst2*-FL and pTriEx/*Tpst2*(H266Q)-FL.

Genetic complementation test using transgenic mice.

pTriEx/*Tpst2*-FL was prepared for microinjection by digestion with *Sal*I and *Swa*I, followed by electrophoresis and purification of the linearized DNA. Transgenic mice were generated by pronuclear injection of the linear transgene into fertilized zygotes prepared from BDF1 mice. RT-PCR analysis of transgene expression in various tissues was performed using primer pair: A/B, tctgactgaccgcgttact/ggccgcatcctcgtgtggg. (Fig. 3A). The transgenic hemizygotes on *grt/+* were mated to generate the transgenic *grt/grt* homozygotes. The following primer pair was used for genotyping transgenic animals: primers C/D, cacactcaagtcatcgtca/cttgcaacgtgtatacagct (Fig. 3A), which cannot amplify
endogenous Tpst2 locus due to the long distance between both primers. For genotyping Tpst2 alleles, PCR and subsequent RFLP analysis using EcoNI have been used.

Hormonal analysis

To test thyroid function of grt/grt mice with or without transgenes, blood was collected into tubes containing 1 mg/ml EDTA at the final concentration and kept on ice, and plasma was obtained by centrifugation. Serum was removed after centrifugation and stored at -20°C until analysis. Serum levels of free T3 and T4 were measured using ACTIVE Free T3 or T4 EIA kits (Diagnostic System lab). Serum TSH was determined using RAT TSH ELISA kit (Shibayagi) according to the manufacturer’s instructions with minor modification to optimize the signal intensity. Serum samples from grt mice needed to be diluted 10 times with PBS (pH 7.4). NIDDK-rat TSH RP-3 was used as the standard.

mRNA detection methods

For detecting mutations, total RNA from WT and grt tissues were extracted with TRizol and reverse-transcribed with a RT-PCR kit according to the manufacturer's instructions (Invitrogen). For the quantitation of thyroid-specific gene, each mRNA level of murine Tshr, Nis, Tpo, Tgn and Pax8 was determined using specific primers as follows:

Tshr-F/R: acctctttacccgagccactgc/tccaggcgcatggcgaaggtgat,
Nis-F/R: cattcccggatcaacctgatggact/tttagagatgaaaaccagcttccg,
Tgn-F/R: cagaccctagtggtgctgatgtg/gcatagtcgtcttgaggtgct,
For the quantitation of mRNA for *Tpst1* and *Tpst2*, RT-PCR was performed for both genes in the same tube on each sample concomitantly. Primers were designed so that the sense primer was shared for amplification of both *Tpst1* and *Tpst2* genes. The primers used were: tcacggccatgtcttgtaag (*Tpst1* antisense); gtgctgttctggttcacctg (*Tpst2* antisense) and tgcaggccttcattctggaggtgat (common sense). To determine the optimum PCR amplification conditions in the linear range, three amounts of cDNA (10, 50 and 100 ng) were tested for each sample. Each reaction tube contained 2 µl of 10 × Ex Taq buffer (Takara), 2 µl of 2.5 mM dNTP mixture, 0.1 µl of Ex Taq polymerase (Takara), 1 µl (4 µM) of each antisense primer and 2 µl (4 µM) of common sense primer. The total reaction volume was 20 µl. PCR consisted of denaturation at 94°C for 3 min, 25 cycles of denaturation at 94°C for 30 sec, annealing at 55°C for 30 sec and extension at 72°C for 30 sec and a final extension at 72°C for 5 min. The PCR products were electrophoresed on 1.5% agarose gel. All assays were performed in duplicate.

**Cell culture, plasmids and expression of transgenes**

*Tpst1*-FL, *Tpst2* (H266Q)-FL, *Tpst1*-FL and *TSHR* cDNA were expressed by the ViraPower Lentivirus expression system (Invitrogen). The primary skin-derived fibroblasts from *grt* mice were grown in DMEM supplemented with 10% fetal bovine serum and penicillin/streptomycin at 37°C. The expression of these genes were detected with semiquantitative RT-PCR.

**In vitro sulfation of peptide arrays**

pTriEx1.1-*Tpst1*-FL, *Tpst2*-FL, and mutant *Tpst2*-FL plasmids were transfected into 1 ×
$10^7$ HEK293T cells using Lipofectamine 2000 (Invitrogen). After 48 h, each cell was homogenized in 500 µl of ice-cold 10 mM Tris-HCl (pH 7.5) containing 100 mM NaCl, 1 mM EDTA, 1% TritonX-100, 0.1 mM phenylmethylsulfonyl fluoride, 1 mg/ml pepstatin, 2 mg/ml aprotinin and 2 mg/ml leupeptin. Expression of each TPST protein in cell lysates (20 µg) was detected by Western blot analysis using a FLAG antibody (Sigma). TPST activity was determined by measuring the transfer of $^{35}$S-sulfate from $^{35}$S-PAPS to Fmoc-based membrane-immobilized 15-mer peptides (39). Peptide arrays, $^{35}$S-PAPS and 50 ml of TPST lysate were accomplished by incubating 50 µl of the buffer {50 mM piperazine-1,4-bis(2-ethanesulfonic acid), pH 6.9, 0.4 mM EDTA, 1 mM Mg-acetate, 200 mM NaCl, 1 mg/ml BSA, 10 mM dithiothreitol} for 1 h at 37°C. After incubation, peptide arrays were washed with 0.1 M Tris-HCl (pH 8.0) twice. The relative amounts of incorporated radioactivity were visualized and quantified with a BAS2500 Bio-Imaging analyzer (Fuji Film). All assays were performed in duplicate.

Measurement of cAMP

Amount of cAMP was determined according to the instruction manual of the cAMP enzyme immunoassay kit (Cayman chemical). In brief, $2 \times 10^5$ cells of each clone were seeded in a 6-well tissue culture dish. The cells were starved of serum for 18 h, then serum-free medium and bovine TSH (10 mU/ml) was added and incubated for 60 min. To assess the value for nonspecific stimulation of intracellular cAMP levels, cells were incubated with 10 mM forskolin and 0.2 mM 3-isobutyl-1-methylxanthine (Sigma). Each experiment was repeated 2 times.

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FIGURE LEGENDS

Figure 1.
(A) Haplotype map of (C57BL/6J × DW/J-grt)F1 × DW/J-grt backcrossed progenies. Three microsatellite markers shown from chromosome 5 are those that have been typed on 1,084 backcross DNAs. Open squares represent the homozygous for the DW allele; black squares represent the heterozygous for B6 and DW alleles. The asterisks indicate the affected mice. Values at the bottom of the figure are the number of progenies.
(B) Transcript map of the critical region for the grt locus. All genes were screened for mutations. Transcriptional orientations are shown by arrows. Cen, centromeric; Tel, telomeric. In the genomic structure of the mouse Tpst2 gene, exonic sequences that contribute to coding regions are boxed and shaded in stripes, while untranslated regions are open-boxed. The sulfotransferase domain of TPST2 protein is boxed in gray. A transversion of C to G is observed at position 798 in the affected mice, which causes an amino acid substitution (H266Q).

Figure 2.
(A) Multiple alignment of amino acid sequences of TPST1 and 2 in various organisms. C. elegans and D. melanogaster have only one TPST protein. An arrow indicates highly conserved histidine residue among all proteins in TPST family.
(B) The 1115 bp PCR product from genome was digested with restriction endonuclease EcoNI and genetic polymorphism was detected by PCR-RFLP. The wild type fragment was not digested, whereas two fragments of 597 and 518 bp were produced upon EcoNI digestion of the grt sequence. The grt/+ heterozygous shows both band patterns.

Figure 3.
(A) The *Tpst2* amplicon was inserted downstream of the CAG promoter and its intron and upstream to the simian virus 40 polyadenylation signal. Arrows indicate the location and orientation of two primer pairs used. Primers A and B were used for detection of the expression of transgene with RT-PCR, whereas primers C and D for genotyping with genomic PCR.

(B) *Tpst2* transgene is expressed in thyroid, brain, kidney, ovary and testis, but not in liver and spleen of transgenic mice.

(C) Transgenic rescue of *grt* phenotypes. Typical *grt/grt* animals with (+) or without (-) the CAG-*Tpst2* transgene (TG) at 5 wk of age.

(D) Comparison of weights among each genotype at 5 wk of age. 1; *grt/grt* with TG (n = 20), 2; *grtl/grt* (n = 17), 3; *grtl/+* with TG (n = 30), 4; *grtl/+* (n = 30) 5; WT (+/+) with TG (n = 16), 6; WT (+/+) (n = 14). The weight data were presented by combining those of male and female, because no difference was found between them at 5 wk of age. An asterisk indicates a significant difference against other groups (P < 0.0001).

(E) Serum TSH, T3 and T4 values of WT (+/+) (n = 8), *grtl/grt* (n = 8), and transgenic *grtl/grt* (TG-*grt*) mice (n = 8). The values are presented as means ± SEM. They were analyzed for statistical significance with Student's t-test and P < 0.005 was considered to be significant (*).

**Figure 4.**

(A) Histological analysis. Thyroid glands from 8-wk WT, *grtl/grt* and TG-*grt* mice were fixed, sectioned, and stained with haematoxylin and eosin (HE) and PAS. The small boxes at 4 × magnification indicate thyroid glands of each genotype. The *grtl/grt* thyroid is significantly smaller in size. High magnification (40 ×) HE staining showed that the *grtl/grt* thyroid is marked by a reduction in follicular cell number and replacement of some hypoplastic portion by adipose tissue (arrow). In PAS staining of *grtl/grt* thyroid gland at...
40 × magnification, colloid staining was not present in some lumens, suggesting a
decrease in thyroglobulin production and storage. On the other hand, hypogenesis of the
thyroids was completely rescued in transgenic grt/grt mice. (B) Semiquantitative
RT-PCR analysis of thyroid-specific genes, Tgn, Pax8, Nis, Tpo and Tshr from two WT,
grt and TG-grt thyroids each. Beta-actin (Actb) was used as internal control. Tshr and Nis
levels were significantly reduced in grt/grt thyroids.

Figure 5.

(A) TPST catalyzes the transfer of sulfate from the universal sulfate donor PAPS to the
hydroxyl group of a peptidyltyrosine residue to form a tyrosine O-sulfate ester.
(B) Equal amounts of cell extracts were extracted from HEK293T cells, subjected to
Western blot analysis with a FLAG antibody. The bands of approximately 55kDa and
50kDa detected on the blots can be identified as TPST2 (TP2) and TPST1 (TP1),
respectively. Actb was used as a loading control.
(C) Tyrosine-sulfation on the peptide array by TPST1 and 2. Each spot contains the
15-mer peptide substrates, GP1BA, CF-8, CCK, FSHR, TSHR, and PSGL1. The same
input of each TPST was used for in this assay. The tyrosine sulfation of peptides was
revealed by the incorporation of $^{35}$S into the substrates by autoradiography. The putative
sulfation sites are indicated as gray characters. TSHR (Mu) and PSGL1 (Mu) are negative
control peptides, in which a mutation from tyrosine to phenylalanine was introduced. TP1,
TP2 and TP2 (H266Q) indicate TPST1, TPST2 and mutant TPST2 (H266Q),
respectively.
(D) The quantitative evaluation of Fig. 5C. The relative incorporation of $^{35}$S into peptides
was evaluated.

Figure 6.
(A) *Tpst1* and *Tpst2* expression in various mouse tissues. Semiquantitative RT-PCR analysis shows ubiquitous expression of both *Tpst1* and *Tpst2*. *Tpst1* (563 bp) and *Tpst2* (741 bp) were concomitantly amplified in the same tube of each sample. Thyroid and thyrocyte cell lines expressed both genes.

(B) Rescue of TSH-mediated cAMP production in *grt/grt* fibroblasts by TPST2 restoration in response to TSH stimulation. Plus (+) or minus (-) indicates the presence or absence of TSH (10 mU/ml) in the medium. Lentiviruses encoding the TSHR and each TP1, TP2 and TP2 (H266Q) were infected in *grt* cells, alone or in combination. Each bar represents the mean ± SEM.

(C) The expression of each gene in *grt/grt* fibroblasts was confirmed by RT-PCR. The expressions of transgenes increased to approximately two-three fold more than that of endogenous *Tpst* mRNAs.

(D) The expression of TPST proteins in *grt/grt* fibroblasts were evaluated by Western blotting with FLAG antibody. Mock indicates the untransfected-*grt/grt* fibroblasts. Each protein was expressed at similar level in the transfected cells.
A

TPST2

\text{grt}(H266Q) \quad ... \quad \text{AWSDAVLHQEDLIGKP} ...

\text{M. musculus} \quad ... \quad \text{AWSDAVLHEDLIGKP} ...

\text{H. sapiens} \quad ... \quad \text{AWSDAVLHEDLIGKP} ...

\text{R. norvegicus} \quad ... \quad \text{AWSDTVLHEDLIGKP} ...

\text{G. gallus} \quad ... \quad \text{SWSDTVLHEELIGKP} ...

\text{X. laevis} \quad ... \quad \text{PWMDAVLHHEELIGKP} ...

\text{D. rerio} \quad ... \quad \text{PWDTAVLHEQLIGKV} ...

TPST1

\text{M. musculus} \quad ... \quad \text{PWNHSVLLLLHEEMIGKA} ...

\text{H. sapiens} \quad ... \quad \text{PWNHSVLLLLHEEMIGKA} ...

TPST

\text{D. melanogaster} \quad ... \quad \text{PWMDAVLHELLFINKP} ...

\text{C. elegans} \quad ... \quad \text{PWDDKVLHEQLIGKD} ...

B

\begin{align*}
\text{EcoNI} & \quad \text{CCTN\ldots NAGG} \\
\text{DW/J- (+/+)} & \quad \text{CCTGCACCACCG} \\
\text{DW/J-(grt/grt)} & \quad \text{CCTGCACCAGGG} \\
\end{align*}

\begin{tabular}{ccc}
(+/+) & (grt+/) & (grt/grt) \\
1115 & 597 & 518 \text{ bp}
\end{tabular}