Defective high-affinity thiamine transporter leads to cell death in thiamine-responsive megaloblastic anemia syndrome fibroblasts

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We have investigated the cellular pathology of the syndrome called thiamine-responsive megaloblastic anemia (TRMA) with diabetes and deafness. Cultured diploid fibroblasts were grown in thiamine-free medium and dialyzed serum. Normal fibroblasts survived indefinitely without supplemental thiamine, whereas patient cells died in 5–14 days (mean 9.5 days), and heterozygous cells survived for more than 30 days. TRMA fibroblasts were rescued from death with 10–30 nM thiamine (in the range of normal plasma thiamine concentrations). Positive terminal deoxynucleotide transferase–mediated dUTP nick end-labeling (TUNEL) staining suggested that cell death was due to apoptosis. We assessed cellular uptake of [3H]thiamine at submicromolar concentrations. Normal fibroblasts exhibited saturable, high-affinity thiamine uptake (Km 400–550 nM; Vmax 11 pmol/min/106 cells) in addition to a low-affinity unsaturable component. Mutant cells lacked detectable high-affinity uptake. At 30 nM thiamine, the rate of uptake of thiamine by TRMA fibroblasts was 10-fold less than that of wild-type, and cells from obligate heterozygotes had an intermediate phenotype. Transfection of TRMA fibroblasts with the yeast thiamine transporter gene THI10 prevented cell death when cells were grown in the absence of supplemental thiamine. We therefore propose that the primary abnormality in TRMA is absence of a high-affinity thiamine transporter and that low intracellular thiamine concentrations in the mutant cells cause biochemical abnormalities that lead to apoptotic cell death.


Introduction

The syndrome called thiamine-responsive megaloblastic anemia (TRMA) with diabetes and deafness (OMIM 249270) was first described by Rogers et al. (1). This autosomal recessive disorder with early childhood onset has recently been localized to chromosome 1, band 1q23.23.3, in four large, geographically and ethnically distinct kindreds (2). Data from several additional families have sharpened the TRMA interval to a 4-cM region and provided evidence for locus homogeneity (3), suggesting that a mutation in a single gene is responsible for the triad of features that define this disease. More than a dozen case reports have described the clinical manifestations of the disorder in 31 patients (2). In all reported cases, the anemia is corrected by pharmacologic doses of thiamine, although macrocytosis (2) and defects in erythroid differentiation in vitro (4) persist after therapy. The disorder represents a novel form of childhood non–type I diabetes mellitus. No anti-insulin or anti-islet antibodies have been found in patients, and in some cases, pharmacologic doses of thiamine have reduced or eliminated insulin requirements. Hearing loss is irreversible in all patients, although some investigators have reported that thiamine therapy prevented further progression of deafness (5). In addition to the definitive clinical characteristics of TRMA, several patients have had strokelike episodes and/or arrhythmias, and abnormalities of the retina and optic nerve (2).

Normal thiamine homeostasis involves the following steps: (a) uptake of thiamine from the gut, (b) transport to tissues and into cells, (c) conversion of thiamine into the cofactor form thiamine pyrophosphate (TPP) by thiamine pyrophosphokinase (TPK; EC 2.7.6.2), and (d) binding of TPP to apoenzymes. Studies of human intestinal thiamine absorption (6–8) suggest that two uptake pathways exist: transport by a saturable, high-affinity/low-capacity carrier, as well as passive intake by a low-affinity/high-capacity system. Because thiamine concentrations in the intestinal lumen are low, Laforenza et al. (8) propose that most thiamine is absorbed into the body by way of the high-affinity transporter. A similar pattern of thiamine uptake has also been seen in human erythrocytes (5, 9, 10). Whether the erythroid and intestinal thiamine uptake activities are identical is unknown. After intracellular thiamine is converted to the active cofactor TPP, TPP is incorporated into four known mam-
malian enzymes: the pentose phosphate shunt enzyme transketolase and three multisubunit, multicofactor enzyme complexes involved in oxidative decarboxylation reactions — pyruvate dehydrogenase, α-ketoglutarate dehydrogenase, and branched-chain keto-acid dehydrogenase. The genes encoding the TPP-binding E1 subunits of each of these enzymes have been localized by other investigators and none map to chromosome 1q (2).

Several investigators have proposed a variety of putative defects of thiamine metabolism in patients with TRMA. On the basis of low levels of α-ketoglutarate dehydrogenase found in lymphocytes from a patient with TRMA, Abboud et al. (11) proposed that a defective TPP binding site on the enzyme leads to TRMA. Others (12, 13) have cited mildly reduced levels of TPK activity as a primary defect. Poggi et al. (5) first noted low TPP content of TRMA erythrocytes (~50% of controls) and postulated that lack of a high-affinity thiamine transporter may be associated with the syndrome (5, 9, 10, 14). The magnitude of the reported defect in thiamine uptake of erythrocytes was subtle, ~60% of normal. Additionally, these authors suggested that decreased TPK activity contributes to TRMA, as they observed a 20%–30% decrease in TPK activity in TRMA erythrocytes compared with controls.

No mammalian thiamine transporter has been isolated. A thiamine transporter gene from Saccharomyces cerevisiae has been identified (15, 16). The THI10 gene encodes a protein with multiple membrane spanning domains, which confers high-affinity thiamine transport in yeast.

In the present study, we demonstrate that fibroblasts from patients with TRMA die in the absence of exogenous thiamine. Using normal or mutant cells, we have explored the thiamine uptake system and the ability of the yeast transporter to correct the lethality in thiamine-depleted cultures. This has enabled us to clarify previous data and to test earlier hypotheses regarding possible defects in TRMA. On the basis of these experiments, we propose a model for the pathogenesis of TRMA in which absence of a very-high-affinity thiamine transporter causes a decrease in cell survival.

**Methods**

*Patients.* Samples were obtained from members of a large, inbred Alaskan kindred reported previously (2). Fibroblasts were obtained from skin biopsies of three patients, several obligate heterozygotes (parents), and non–blood relatives (wild-type). Informed consent was obtained from all subjects and their parents.

*Cell culture and cell lines.* Diploid fibroblasts were grown from skin samples in thiamine-replete α-MEM (1 mg/l thiamine; Gibco BRL, Grand Island, New York, USA), 15% FCS, 50 U/ml penicillin, 300 μg/ml streptomycin sulfate (GIBCO BRL) in 5% CO2 at 37°C. α-MEM minus thiamine (thi– medium) with 15% dialyzed FCS (GIBCO BRL) was used for thiamine starvation. Thiamine depletion was begun after cells were washed three times in PBS and passaged into thi– medium. Experiments shown were performed on two TRMA cell lines (first cousins), two lines from obligate heterozygotes, and two normal controls.

*Thiamine uptake assays.* [3H]thiamine hydrochloride (specific activity 15 Ci/mmol; American Radiolabeled Chemicals Inc., St. Louis, Missouri, USA) was used for uptake assays. Thiamine uptake was assayed in extracellular concentrations ranging from 33 nM to 3 mM. Cells were washed with PBS, trypsinized, and plated at a density of 1–2 × 105 cells/35-mm well in 1 ml of thi– medium. Uptake was measured in cells incubated at 37°C in thi– medium without FCS. In unlabeled thiamine competition assays, thiamine- HCl (Sigma Chemical Co., St. Louis, Missouri, USA) was added at the appropriate concentration. After cell labeling, cells were washed three times with PBS and harvested with 0.05% trypsin-EDTA (GIBCO BRL). The amount of cell-associated [3H]thiamine was determined by scintillation counting in 5 ml Ecolome scintillation fluid (ICN Radiochemicals Inc., Costa Mesa, California, USA) using a Beckman LS 3801 instrument (Beckman Instrument Inc., Palo Alto, California, USA). Incubation at 0°C was used as a control for passive, cell-associated radioactivity.

*Assay for α-ketoglutarate dehydrogenase.* The method was adapted from a pyruvate dehydrogenase assay described by Hansen (17). α-ketoglutarate dehydrogenase catalyzes the formation of succinyl CoA from [1-14C]α-ketoglutarate, resulting in release of 14CO2, which is trapped and measured. The mitochondrial fraction (10,000-g pellet of postnuclear supernatant) was incubated with 1 μCi [14C]α-ketoglutarate (Du Pont NEN Research Products, Boston, Massachusetts, USA) diluted with nonlabeled α-ketoglutarate to 1 mM, 1 mM coenzyme A (li- thium salt; Sigma Chemical Co.), 1.6 mM nicotinamide adenine dinucleotide, 40 mM sodium phosphate (pH 7.4), 0.5 mM dihydrothreitol, and 5 mg/ml BSA, in the presence or absence of 1 mM TPP. The reaction was carried out in a 5-cc vial sealed

<table>
<thead>
<tr>
<th>Concentration of thiamine required to prevent toxicity in TRMA fibroblasts</th>
<th>0.3 nM</th>
<th>0.9 nM</th>
<th>3 nM</th>
<th>9 nM</th>
<th>30 nM</th>
<th>300 nM</th>
<th>900 nM</th>
</tr>
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<tbody>
<tr>
<td>TRMA</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>±</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td>Control</td>
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Cells were washed, trypsinized, and re-plated in varying thiamine concentrations and then maintained for 30–40 days. Plus sign indicates cell survival, minus sign indicates cell death, and plus over minus indicates morphologic changes without cell death. Normal human plasma contains 18–32 nM free thiamine. Replete MEM contains >3,000 nM. Results are from duplicate wells in three individual experiments. TRMA, thiamine-responsive megaloblastic anemia.
fibroblasts were plated at a density of 10^6 cells/100-mm dish and incubated in thi– medium. The following day, the medium was changed on day 0, and then medium was collected and the protein was removed with 50 mM Tris (pH 8.0). Free thiamine ran in the void volume, and ketoisovalerate, 2-keto-3-methylvalerate, 2-ketoisocaproate, and 2-ketoglutarate. As an internal standard for each reaction, 10 μg of pentadecanonic acid was added to each medium sample before extraction and derivatization, to control for recovery. Standards for authentic α-ketoacids were obtained from Sigma Chemical Co. Results (peak heights) were compared by normalizing to the size of the internal standard peak and corrected for cell number in each dish.

Expression of yeast thiamine transporter gene in TRMA fibroblasts. The yeast thiamine transporter gene THI10 was amplified with Pfu DNA polymerase (Stratagene Inc., La Jolla, California, USA) from total yeast genomic DNA (Promega Corp., Madison, Wisconsin, USA) using primers flanking the open reading frame YRL237w. The primers contained EcoRI restriction sites to aid in subsequent cloning steps. Amplified products were subjected to restriction sites to aid in subsequent cloning steps. Amplified products were subjected to restriction enzyme digestion, electrophoresis, and transformation of E. coli. The restriction products were resolved by FPLC anion exchange chromatography as described in Methods. Determinations are means ± SD of triplicate determinations. FPLC, fast protein liquid chromatography; thi– medium, α-MEM minus thiamine; TPP, thiamine pyrophosphate.

Table 2
Uptake and conversion of [14C]thiamine to TPP at steady state by normal and TRMA–/– fibroblasts

<table>
<thead>
<tr>
<th>Total thiamine</th>
<th>TPP</th>
<th>Conversion (%)</th>
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<tbody>
<tr>
<td>pmol/10^6 cells</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+/+</td>
<td>12</td>
<td>4.1</td>
</tr>
<tr>
<td>–/–</td>
<td>9.2</td>
<td>5.3</td>
</tr>
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Cells were washed and plated in thi– medium overnight before incubation with 2.4 mM [14C]thiamine for 18 h. Cell-associated radioactivity was extracted in 5% TCA and resolved by FPLC anion exchange chromatography as described in Methods. Determinations are means ± SD of triplicate determinations. FPLC, fast protein liquid chromatography; thi– medium, α-MEM minus thiamine; TPP, thiamine pyrophosphate.
THI10 was cloned into the mammalian expression vector pEF1α-neo and confirmed by restriction and sequence analysis. Fibroblasts from patients with TRMA as well as normal controls were cultured as already described here. On the day before transfection, confluent 100-mm tissue culture dishes were split 1:2 and plated in replete medium lacking antibiotics. Lipofectamine Plus Reagent (4 mg; GIBCO BRL) was used for the transfection of either pEF1 α-neo/THI10 or pEF1 α-neo alone into the TRMA fibroblasts in serum-free medium. Three hours after transfection, medium with serum was added so that the final concentration of FCS was 15% and thiamine was 3 μM. Transfected cells were maintained at 37°C and 5% CO2. The medium was replaced 24 h after transfection with medium containing 300 μg/ml G418 (GIBCO BRL) to select transformants. Medium was changed every 2–3 days. Three weeks after transfection, G418-resistant cells were replated to be 50%–75% confluent and were maintained in medium with or without thiamine at no more than 90% confluence.

**Results**

**Thiamine deprivation of TRMA fibroblasts.** Fibroblasts from patients with TRMA and from normal controls were assessed for a thiamine-dependent phenotype by growing cells in thiamine-replete medium. After 5–14 days of thiamine starvation, the morphology of TRMA fibroblasts began to change (Fig. 1a). The mean time to abnormal morphology was 9.5 days (± 5 days, SD) in eight experiments. Cells maintained in thiamine-replete medium remained indistinguishable from wild-type (Fig. 1b). Within 1–3 days of morphological evidence of toxicity, cells in thiamine-deficient medium detached or died on the plates (Fig. 1c). However, addition of thiamine (3 μM) to TRMA fibroblasts when adverse changes first became evident rescued the remaining cells (Fig. 1d). Rescued cells reverted to normal morphology within 24 hours and could then be grown as before thiamine starvation. Normal fibroblasts in thiamine-replete medium showed some decrease in growth rate but did not die under these conditions (experiments terminated at 40 days). Heterozygous cells died after 30 and 32 days on two occasions. These experiments suggest that TRMA fibroblasts are more sensitive to thiamine starvation and can be distinguished from normal fibroblasts by their requirement for exogenous thiamine. We postulate that normal cells survive because of the trace amounts of vitamin (below fluorometric assay) remaining in the dialyzed serum.

The concentration of thiamine necessary to sustain TRMA fibroblasts was investigated by growing cells in medium with thiamine concentrations ranging from 0 to 900 nM. Results are shown in Table 1. Nine nanomolars of thiamine rescued mutant cells to a variable degree, whereas 30 nM vitamin always rescued completely. This concentration is approximately that in plasma and is 100-fold less than the amount in replete tissue culture media (3 μM). We concluded that the defect in TRMA cells might be observable only at extremely low thiamine concentrations.

A representative time course experiment. Fibroblasts were incubated in 66 nM [3H]thiamine over a period of two hours. Values were corrected for cell-associated counts in cells incubated in an ice bath. These 0°C controls were always <0.1 pmol/10^6 cells. However, these results are not corrected for nonspecific, or unsaturable, uptake. Uptake was approximately linear for 30 minutes and approached a plateau by approximately two hours in serum-free medium. Subsequent assays were done at 30 minutes. As demonstrated by this experiment, mutant fibroblasts show a significant defect in thiamine uptake, and heterozygous cells are intermediate in degree of uptake.

We next assessed uptake at varying concentrations of thiamine in 30-minute assays. A substantial difference between mutant and normal thiamine uptake was observed at the lowest concentrations tested (Fig. 2b). These results are corrected for nonspecific uptake by subtracting counts obtained in the presence of 3 mM unlabeled thiamine (50,000 to 100,000-fold excess). The mutants exhibit ~5% of the wild-type uptake at 33 and 66 nM thiamine under these conditions. This is consistent with either a severe defect or absence of a high-affinity transporter. Laforenza et al. (18) have observed a pH-sensitive thiamine transport system in rat intestine that is augmented by omeprazole. The fibroblast high-affinity uptake system is apparently unaffected by omeprazole or pH changes in the 6.5–8.0 range (data not shown).

**Kinetic properties of high-affinity thiamine transport by fibroblasts.** Pilot studies suggested that thiamine uptake became unsaturable at concentrations <1 μM. We therefore performed detailed analysis of uptake by normal fibroblasts.
in the nanomolar range as shown in Fig. 3. The properties of uptake by growing fibroblasts proved highly reproducible from experiment to experiment and among cell lines from the different normal and mutant cell lines tested. We analyzed only specific, or saturable, uptake by subtracting counts associated with cells in 500-fold excess of unlabeled thiamine. Under these conditions, Lineweaver-Burke analysis revealed a saturable process with apparent $K_m$ values from 400 to 550 nM, and a maximum transport rate of 11 pmol/10^6 cells/30 min. We were unable to detect a saturable component in the TRMA fibroblasts in several attempts, although a small amount of residual transport activity could still be present. However, the unsaturable component of thiamine uptake at thiamine concentrations >1 μM was essentially the same in TRMA as in wild-type cells (not shown).

**Assays of other candidate enzymes for TRMA.** We also examined the function of reported candidate genes, α-ketoglutarate dehydrogenase and TPK. We assayed α-ketoglutarate dehydrogenase in mitochondrial pellets of fibroblasts from two patients with TRMA or normal controls, as described in Methods, by measuring the coenzyme A–dependent release of 14CO₂ from [1-14C]α-ketoglutarate. Results expressed as nmol CO₂ release/mg protein/h (mean of duplicate determinations ± range) were as follows: control, 250 (± 42); TRMA patient no. 1, 144 (± 39); TRMA patient no. 2, 273 (± 13). The differences were not statistically significant and indicated that this enzyme is not defective in the TRMA fibroblasts.

We next measured conversion of thiamine to TPP, as an indirect measurement of TPK activity in vivo, in overnight incubation of mutant or wild-type cells as described in Methods. We found no defect in conversion of thiamine to TPP. Indeed, the degree of conversion of radiolabeled thiamine to TPP (Table 2) was consistently greater in the mutants (58% ± 7% vs. 45% ± 6%; mean ± SD, $P = 0.043$ by two-tailed $t$ test). We speculate that the difference is due to smaller intracellular pools of unlabeled thiamine in the mutant compared with the wild-type cells during these experiments. Thiamine monophosphate was a minor product in both wild-type and mutant cells.

**Apoptosis assay.** Once the phenotype of TRMA fibroblasts in thi– medium became apparent, the mechanism of cell death was investigated. Using the TUNEL assay for fragmented genomic DNA, TRMA and control fibroblasts depleted of thiamine were assayed (Fig. 4). Cells were assayed on day 10 of incubation in thi– medium. Both normal and TRMA fibroblasts are TUNEL negative when grown in thiamine-replete media (Fig. 4, a and b, respectively). Normal fibroblasts remain negative for TUNEL staining in thi– medium (Fig. 4c). However, as shown in Fig. 4d, TRMA cells become TUNEL positive as the morphology becomes abnormal. These results suggest that TRMA fibroblasts die by programmed cell death. We were unable to detect a DNA “laddering” pattern of apoptosis in dishes of TRMA cells approaching cell death in thi–medium. We postulate that only a small number of cells undergo death at any given time. It is also possible that the positive TUNEL staining represents a mechanism for DNA fragmentation other than apoptosis.

**Organic acid accumulation in medium of TRMA cells subjected to thiamine starvation.** We postulated that the thiamine uptake defect in TRMA cells should lead to more rapid appearance of a thiamine-depleted biochemical phenotype than wild-type cells. Table 3 shows the results of organic acid analysis of medium from TRMA and control cells after one and six days of thiamine starvation. Compared with thi– medium alone, both mutant and wild-type medium show very little increase of the critical organic acids on day 1. By day 6, however, all of these compounds are significantly higher in the mutant cells. This implies lower internal stores of TPP in the mutant cells within a few days, despite the very high concentrations of thiamine in standard medium. Although pyruvate is present in the medium as a supplement, it accumulates severalfold in the mutant cells by day 6. Lactate rose in both cell lines.

**Rescue of thiamine-dependent lethality in TRMA fibroblasts by S. cerevisiae THI10 gene.** On the basis of the apparent deficiency of a thiamine transporter in TRMA cells, we tested the ability of S. cerevisiae THI10 gene (thiamine transporter) to allow survival of mutant fibroblasts in thiamine-depleted conditions. The THI10 gene was cloned into expression vector pEF1α-neo as described in Methods and was stably transfected into TRMA cells. As shown in Fig. 5, THI10-transfected cells, but not vector-only controls, were able to survive for more than six weeks in thi– medium. Vector-only controls died after 13 days in thi– medium, and the experiment was terminated after 44 days. These results strongly support the hypothesis that the primary TRMA defect is a thiamine transporter. These results do not prove that the transport defect is across the plasma membrane (as opposed to an internal membrane), but we suspect that this is the case based on the uptake assays described.

**Discussion**

We have demonstrated that TRMA fibroblasts are uniquely sensitive to thiamine depletion. Cells die after several days in thi– medium, with DNA fragmen-
tation as detected by TUNEL assay consistent with apoptosis or programmed cell death. To our knowledge, this is the first report of a genetic disorder of intermediary metabolism causing apoptosis. Labeling studies using [3H]thiamine of high specific activity demonstrate that TRMA fibroblasts take up ~5%–10% of wild-type amounts of thiamine at nanomolar concentrations. Nonspecific (unsaturable) thiamine uptake is not impaired in the mutants. Thus, in standard tissue culture medium (3 μM thiamine plus the rich supply in serum), the metabolic defect is obscured, whereas in thi– medium, characteristic organic acids accumulate faster in TRMA cells than in normal controls. Our results support the hypothesis that the primary defect in TRMA is a recessive mutation in the gene encoding a high-affinity thiamine transporter. Rescue of the lethal phenotype by a known yeast transporter provides additional support for this hypothesis.

Rindi and colleagues (9, 10) have previously defined a subtle defect of thiamine uptake in erythrocytes of patients with TRMA, ~60% of control. There are at least two possible explanations for the differences observed between their studies and the present work, in which we have found greater than 10-fold reduction in [3H]thiamine uptake at low concentrations. First, the erythrocyte and fibroblast high-affinity systems might be similar or identical, but experimental variables allow the defect to be more readily discerned in the present fibroblast studies. For example, we have used thiamine of specific activity 20-fold higher than that used in previous studies (15 vs. 0.75 Ci/mmol; refs. 9, 10). This facilitates assays at low substrate concentrations, at which differences between mutant and normal cells are greatest (Fig. 2). The $K_m$ of the high-affinity system described here, ~0.5 μM, is at the lower end of concentrations studied by Rindi et al. (9). Second, there may be more than one specific thiamine uptake system, operating together or individually in different cell types. If erythrocytes normally have two systems, and fibroblasts only one, the degree of abnormality would appear more severe in the fibroblasts. The current data do not distinguish these possibilities.

Cell death by apparent apoptosis in severely thiamine-depleted TRMA fibroblasts raises several questions about the pathogenesis of the disease. First, why are patients with TRMA not severely ill without pharmacologic thiamine therapy? With a single exception (19), subjects with TRMA have not had symptoms suggestive of severe tissue thiamine deficiency, and they have not had organic aciduria (5, 11). Red blood cell transketolase is also normal. We find that 10–30 nM thiamine is enough to rescue fibroblasts (Table 1) and that thiamine depletion does lead to organic acidosis in the cells from patients after a few days of thiamine depletion (Table 3). Normal plasma levels of free thiamine are on the order of 18–33 nM (5, 13). This implies that patients with TRMA on an adequate diet without pharmacologic supplementation are metabolically stable, with thiamine uptake solely by the low-affinity system.

Second, why is the deafness in TRMA progressive, whereas the anemia, and to some degree the diabetes, are reversible? We postulate that the thiamine requirement of cochlea or acoustic nerve cells is substantially higher than that of fibroblasts, which have very low energy requirements compared with neurons, myocytes, and other cell types. (We have not examined this phenomenon in other TRMA cell types.) If so, even pharmacologic doses of enteral thiamine may not prevent a thiamine-deficient phenotype in cell types with high energy usage at all times. On the basis of our observations in fibroblasts, we propose that this would lead to occasional cell death in the most sensitive tissues. This, in turn, may result in optic atrophy and progressive deafness.

### Figure 4

Apoptosis of fibroblasts subjected to thi– medium. TUNEL stain was performed as described in Methods. Normal (a and b) or TRMA mutant cells (c and d) were incubated for 10 days in the presence (a and c) or absence (b and d) of thiamine. (a) Wild-type cells, thiamine-replete. (b) Wild-type cells, thiamine-depleted. (c) TRMA mutant cells with thiamine. (d) TRMA fibroblasts, thiamine-depleted. ×40. TUNEL, deoxynucleotide transferase–mediated dUTP nick end-labeling.

### Figure 5

Complementation of TRMA fibroblasts defects with yeast thiamine transporter gene $THI10$. Cells transfected with the $THI10$ gene and vector alone were grown in the presence or absence of thiamine as described in Methods. TRMA fibroblasts transfected with vector alone and grown in thi– medium died after 13 days. The experiment was terminated after 44 days. (a) TRMA fibroblasts transfected with $THI10$ gene, thiamine-replete. (b) TRMA fibroblasts transfected with vector alone, thiamine-replete. (c) TRMA fibroblasts transfected with $THI10$ gene, thi– medium. (d) TRMA fibroblasts transfected with vector alone, thi– medium.
marrow, in contrast, may be less sensitive to this effect for two reasons. First, as long as the pluripotent hematopoietic stem cells are not sensitive to cell death, the erythroid lineage can be repopulated when patients are thiamine replete. Persistent macrocytosis (2) and poor erythroid colony growth in vitro (4) suggest that even thiamine-replete erythroid progenitors are somewhat sensitive to the thiamine transport defect. Diabetes in TRMA patients has been variably reversible with thiamine. Persistent macrocytosis (2) and hematopoietic stem cells are not sensitive to cell death, for two reasons. First, as long as the pluripotent marrow, in contrast, may be less sensitive to this effect present throughout the gut), with apparent recent reports of a thiamine uptake system from human liver and muscles, which would be responsible for organic acidemia in severe thiamine deficiency, must be resistant to cell toxicity in the ambient thiamine concentration found in untreated patients with TRMA.

It is conceivable that deafness in patients with TRMA could be prevented by parenteral, early (even prenatal) thiamine therapy. Laforenza and colleagues (8) have recently reported a thiamine uptake system from human intestinal biopsies (maximum rate in duodenum but present throughout the gut), with apparent \( \frac{K_m}{V_{max}} \approx 4 \mu M \). Whether this system is due to the same gene product defective in TRMA remains to be determined. The transport across the intestinal epithelium into the abluminal space was not measured. If either of these processes depends on the TRMA gene, it is possible that even pharmacologic doses of oral thiamine cannot provide supra-normal levels of the vitamin in the blood and tissues. For cells with high energy requirements, this may not be sufficient in TRMA. We propose that if transintestinal transport is also defective in TRMA, patients may do substantially better if treated from birth (or even prenatally) with parenteral thiamine.

It is not yet clear how the vitamin transport defect in TRMA is linked to programmed cell death. Matsushima et al. (20) recently reported apoptotic cell death in the thalassemia of thiamine-deficient rats. This suggests that the phenomenon is not limited to TRMA. However, it does not clarify the relationship between cellular thiamine deficiency and activation of apoptosis pathways. Fibroblasts in thi- medium release organic acids expected from defects in the TPP requiring enzymes, i.e., branched chain \( \alpha \)-ketoacids and pyruvate (Table 3). This suggests that thiamine deprivation can lead to dysfunction in the Krebs cycle, shutting down mitochondrial energy production. We speculate that TRMA cells in thi- medium will be relatively more impaired at these steps than normal fibroblasts. Mitochondrial changes, in turn, may play a pivotal role in activation of apoptosis. A number of possible mechanisms have been proposed (reviewed in ref. 21). Mitochondrial dysfunction might lead to leakage of caspase activators, cytochrome C, or apoptosis-inducing factor, for example. It will be of interest in the future to examine whether TRMA cells from other kindreds exhibit the same uptake kinetics and thiamine-dependent survival as cells from the Alaskan patients tested here. Independent mutations have apparently arisen on different haplotypes in each TRMA family (2, 3), and it is possible that differences in phenotype will correlate with different genotypes.

In light of our present results, we believe that a candidate gene for TRMA would be a thiamine transporter, responsible for vitamin uptake at low extracellular thiamine concentration. Database homology searches have not yielded such a gene. Isolation of the TRMA gene will facilitate further studies of this unique biochemical disorder.

Acknowledgments

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