Heimler Syndrome Is Caused by Hypomorphic Mutations in the Peroxisome-Biogenesis Genes PEX1 and PEX6

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Heimler syndrome (HS) is a rare recessive disorder characterized by sensorineural hearing loss (SNHL), amelogenesis imperfecta, nail abnormalities, and occasional or late-onset retinal pigmentation. We ascertained eight families affected by HS and, by using a whole-exome sequencing approach, identified biallelic mutations in PEX1 or PEX6 in six of them. Loss-of-function mutations in both genes are known causes of a spectrum of autosomal-recessive peroxisome-biogenesis disorders (PBDs), including Zellweger syndrome. PBDs are characterized by leukodystrophy, hypotonia, SNHL, retinopathy, and skeletal, craniofacial, and liver abnormalities. We demonstrate that each HS-affected family has at least one hypomorphic allele that results in extremely mild peroxisomal dysfunction. Although individuals with HS share some subtle clinical features found in PBDs, the diagnosis was not suggested by routine blood and skin fibroblast analyses used to detect PBDs. In conclusion, our findings define HS as a mild PBD, expanding the pleiotropy of mutations in PEX1 and PEX6.

Introduction

Heimler syndrome (HS [MIM: 234580]) is a rare autosomal-recessive disorder that was first described in 1991 in two siblings, born to healthy and non-consanguineous parents, with sensorineural hearing loss (SNHL), enamel hypoplasia of the secondary dentition, and nail abnormalities.1 Subsequently, five additional cases have been reported.2–4 In 2011, Lima et al.5 reported retinal pigmentation in one of the original individuals diagnosed with HS.1 The genetic cause of HS had not been identified, but it had been suggested that the syndrome could be due to mutations in a gene affecting derivatives of the ectodermal tissue, given that the described abnormalities have a common embryological origin.1 In contrast, because of the spectrum of clinical features, Lima et al.5 classified HS as a ciliopathy. In this study, we ascertained eight HS-affected families and analyzed them in a hypothesis-free way by whole-exome sequencing (WES). We identified biallelic PEX1 (MIM: 602136) or PEX6 (MIM: 601498) mutations in six of them. Further functional studies showed that HS is not a ciliopathy but rather a PBD with an atypical mild phenotype that shows limited clinical overlap with other PBDs.

Material and Methods

WES

We used standard techniques4 to collect blood samples of different family members (Figure 1A) and isolate DNA from blood or saliva. All subjects or their legal representatives provided written informed consent for this study, which was performed in accordance with the Declaration of Helsinki protocols and approved by the local institutional review boards.

WES was undertaken for one or more individuals from families 1–5, 7, and 8. DNA enrichment for WES was achieved with the

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Figure 1. Pedigrees of the Families Affected by HS, Illustration of Clinical Characteristics of Individual F5-II:2, and Air-Conduction Audiograms of HS Individuals

(A) Affected individuals are shaded. Arrows indicate individuals in whom WES was performed and blood and/or fibroblasts were analyzed. For family 1, additional exomes were sequenced for members I-1, II-1, and II-3. DNA for cosegregation analysis was available from family members with an asterisk.

(B) An orthopantogram of individual F5-II:2 shows severe amelogenesis imperfecta.

(C) A retinal photograph taken when individual F5-II:2 was 20 years old shows marked mottling of the retinal pigment epithelium.

(D) A fingernail of individual F5-II:2 shows evidence of onychoschizia and Beau’s lines.

(E) An air-conduction audiogram of individual F5-II:2.

(F) An air-conduction audiogram of individual F5-II:3.
were selected. The effect of the variant on protein function were filtered out. Only non-synonymous, frameshift, nonsense, present in our in-house control database, including 770 exomes, Sequencing Project Exome Variant Server (ESP6500). Variants (v.137), 1000 Genomes (April 2012), and the NHLBI Exome frequency (MAF) above 1% were filtered out on the basis of dbSNP bias smaller than 20. Next, common variants with a minor allele quality above 50, (2) quality by depth above 4.8, (3) mapping-performed according to the following parameters: (1) mapping or ABI 3730 DNA sequencer (Applied Biosystems).

NeutraX Rapid Capture Exome 38 Mb Kit (Illumina) for family 1 and with the SureSelect Human All Exon Kit v.5 (Agilent) for families 2–5, 7, and 8. Paired-end sequencing (100 bp) was run on an Illumina HiSeq 1500 or 2500. A minimum of 4.5 Gb of sequence was generated for each individual, yielding a mean depth of coverage ranging from 75× to 163× and 88.3%–98% of target bases sequenced at 20× coverage. The sequence data were mapped to the human reference genome (UCSC Genome Browser hg19) with the Burrows Wheeler Aligner (BWA). Variant calling was performed with the Genome Analysis Toolkit (GATK) v.2.4.7. VariantDB was used for variant annotation and filtering. Variant annotation was based on information from GATK, SNPeff, ANNOVAR, and Gene Ontology. Quality-based filtering was performed according to the following parameters: (1) mapping quality above 50, (2) quality by depth above 4.8, (3) mapping-quality rank sum between –3 and 3, and (4) Fisher-scaled strand bias smaller than 20. Next, common variants with a minor allele frequency (MAF) above 1% were filtered out on the basis of dbSNP (v.137), 1000 Genomes (April 2012), and the NHLBI Exome Sequencing Project Exome Variant Server (ESP6500). Variants present in our in-house control database, including 770 exomes, were filtered out. Only non-synonymous, frameshift, nonsense, and splice-site variants and genes containing biallelic variants were selected. The effect of the variant on protein function was predicted by PolyPhen-2, SIFT, and MutationTaster with dbNSFP for non-synonymous variants and with multiple tools for potential splice-site mutations. Confirmation of the putative disease-causing variants and their cosegregation with the disease phenotype and analysis of PEX6 for variants in family 6 were performed by standard Sanger dyeoxy sequencing on an ABI 3130XL or ABI 3730 DNA sequencer (Applied Biosystems).

Analysis of STRs and Healthy Control Population

For family 1, we amplified, pooled, and analyzed short tandem repeat (STR) markers via capillary electrophoresis on an ABI 3130XL DNA sequencer (Applied Biosystems). We analyzed data with GeneMapper v.3.7 (Applied Biosystems). We used NCBI Map Viewer (annotation release 105) to search for STRs in a 10 Mb region surrounding the PEX1 mutation.

We collected blood samples from umbilical cords of 250 unrelated newborns originating from different regions of Morocco. The Moroccan origin of their parents and grandparents was confirmed. We obtained informed consent for DNA analysis from the parents. We used a standard salting-out method to extract genomic DNA from 3 ml blood. We developed a real-time PCR (Applied Biosystems 7500 Fast Real-Time PCR Systems) assay by using TaqMan probes for the PEX1 c.3750G>A (p.Tp1250*) nonsense mutation (Table 2) and validated the assay by using homozygous and heterozygous members of family 1.

Electron Microscopy

We cultured fibroblasts to confluence and fixed them with 4% formaldehyde and 2.5% glutaraldehyde in 0.1 M HEPES buffer (pH 7.2). We postfixed the cells with 1% osmium tetroxide and 1.5% potassium ferrocyanide in 0.1 M cacodylate buffer (pH 7.2) for 1 hr, then in 1% tannic acid in 0.1 M cacodylate buffer (pH 7.2) for 1 hr, and finally in 1% uranyl acetate in water for 1 hr. The samples were dehydrated in ethanol series, infiltrated with TAAB 812 resin, and polymerized for 24 hr at 60°C. Ultrathin sections were cut with a Reichert Ultracut ultramicrotome and visualized with a FEI Tecnai 12 Biotwin microscope at 100 kV accelerating voltage. Images were taken with a Gatan Oris SC1000 CCD camera.

Biochemical and Enzyme-Activity Assays

We measured peroxisomal parameters in plasma (very-long-chain fatty acids [VLCFAs], bile acid intermediates, piperocic acid, phytic acid, and pristanic acid), in erythrocytes (plasmalogens), and in skin fibroblasts (VLCFAs, C26:0 and pristanic acid β-oxidation, phytic acid α-oxidation, and dihydroxyacetonephosphate acyltransferase [DHAPAT] activity). Immuno-blots analyses assessed the processing of thiolase and acyl-CoA oxidase 1 (ACOX1) in fibroblasts.

Molecular Cloning

We introduced the different PEX1 or PEX6 variants identified in the individuals with HS in the mammalian expression vector pcDNA3 containing full-length PEX1 or PEX6 cDNA, respectively, by site-directed mutagenesis (QuickChange Site-Directed Mutagenesis Kit, QuIAGEN) according to the manufacturer’s instructions. We obtained the pcDNA3 vector containing PEX6 c.1930C>T (p.Arg644Trp) by amplifying PEX6 cDNA spanning nucleotides c.1856 to c.1961 (with flanking restriction sites for Nhel and Kpnl) from total RNA isolated from the fibroblasts of family 5 individual II:2 (F5-II:2). We subsequently subcloned the ampiclons into the pcDNA3 vector containing full-length wild-type PEX6 cDNA. We confirmed successful introduction of the variants by sequence analysis of the cDNAs. To exclude unintentional mutations in the vector backbone during site-directed mutagenesis, we either used several clones per construct for further analyses or recloned the mutated cDNA into pcDNA3 plasmids.

Cell Culture and Transfection

We used primary skin fibroblasts from individuals with HS and primary skin fibroblast cell lines completely deficient of PEX1 (compound heterozygous for p.[Thr263Ilefs*6];[Ile700Tyrfs*42], c.[788_789del];[2097dup]) or PEX6 (homozygous for p.Gly135 Aspfs*23 [c.-402del]). Cells were cultured in DMEM with L-glutamine (Bio-Whittaker) supplemented with 10% fetal bovine serum (Bio-Whittaker), 25 mM HEPES buffer (BioWhittaker), 100 U/ml penicillin, 100 μg/ml streptomycin (Life Technologies), and 250 ng/ml Fungizone (Life Technologies) in a humidified atmosphere of 5% CO2 at 37°C or 40°C. Transfections were performed with the AMAXA NHDF Nucleofector Kit (Lonza) according to the manufacturer’s instructions (program U23). The medium was changed 24 hr after transfection, and the cells were imaged 72 hr after transfection.

Immunofluorescence Assays

We analyzed peroxisomal appearance in skin fibroblasts from HS individuals by immunofluorescence microscopy. The cells were cultured on glass slides to a confluency of 50%–70%. For fixation, we treated the cells with 2% paraformaldehyde (Merck) in PBS for 20 min at room temperature and permeabilized them with 0.5% Triton X-100 (BioRad) for 5 min. The peroxisomal matrix protein catalase was labeled with the monoclonal antibody α-catalase (Map 17E10, own production), biotinylated α-mouse antibodies (E 433, Dako), and streptavidine-FITC (F 422, Dako). Peroxisomal membranes were labeled with antibodies against PMP70 (ABCD3) (PMP70, no. 718300, Zymogen) and Alexa Fluor 555 goat anti-rabbit (Invitrogen). The slides were fixed on mounting medium.
Assays of Genetic and Functional Complementation

We performed genetic complementation of fibroblasts by transfecting the cells from HS individuals with PEX cDNA as described in Ebberink et al. To test the functionality of the PEX variants, we co-transfected pcDNA3-PEX1 or -PEX6 plasmids with the peroxisomal matrix marker pEGFP-SKL into skin fibroblasts deficient in PEX1 or PEX6. Cells transfected with only pEGFP-SKL served as negative controls, whereas co-transfections of the marker with pcDNA3 vectors containing the respective wild-type PEX cDNA served as positive controls. We subsequently analyzed the localization of the fluorescent signal 3 days after transfection by using the fluorescence microscope Zeiss Axio Observer A1.

To evaluate the effect of the variants found in the affected individuals, we determined per transfection the percentage of cells showing a punctate GFP signal (indicating “peroxisome-positive” or “complemented” cells) of the total number of 100–200 transfected cells. These ratios were normalized to the complementation efficiency of the positive control (set as 100%) and averaged per construct (n = 5–7). We used the one-sample Wilcoxon signed-rank test to test the statistical significance of deviations from the positive control.

## Results

### Identification of PEX1 and PEX6 Variants

In order to unravel the genetic cause and the pathological mechanism of HS, we ascertained eight HS-affected families, including three previously described families (family 2, family 6, and family 8) (Figure 1A and Table 1). The HS individuals were all characterized by a homogeneous phenotype of severe to profound pre-lingual bilateral SNHL most pronounced at high frequencies and amelogenesis imperfecta (Table 1 and Figures 1B–1F). The retinal-pigmentation phenotype was highly variable such that some individuals showed no evidence of retinal pigmentation at 16 and 21 years (in families 1 and 7, respectively), whereas the affected individual in family 4 (F4-II:1) had progressive visual impairment with no peripheral vision and no night vision at age 6 years. Visual assessment was not available from individuals in families 3 and 8. All affected individuals had normal intellect. After the initial WES findings, a single individual (F1-II:3) had a brain MRI scan, which was normal.

We performed WES on at least one affected individual from each family. WES of affected individuals from families 1–5, 7, and 8, as well as selection of biallelic rare or previously reported pathogenic non-synonymous, frameshift, nonsense, or splice-site variants consistent with recessive inheritance co-segregating with the disease phenotype, did not identify variants in a single gene common to all unrelated affected individuals. However, we noted that affected individuals from families 1–4 each had biallelic putative disease-causing variants in PEX1 (GenBank: NM_000466.2), whereas in family 5 a mutation in PEX6 (GenBank: NM_000287.3) was found. PEX1 and PEX6 are two related genes involved in peroxisome...
biogenesis (Figure 2). In family 1, we identified a previously unreported homozygous exon 23 nonsense variant, c.3750G>A (p.Trp1250*), only 19 bases from the last exon-exon boundary of PEX1. The resultant transcript is assumed to escape nonsense-mediated decay and thus lead to expression of a truncated protein. The homozygous variant was surrounded by a region identical by descent from a common ancestor (Figure S1) and absent in 250 ethnically matched healthy control individuals. In families 2 and 4, we identified a heterozygous, previously reported pathogenic PEX1 c.2097dup variant resulting in a p.Ile700Tyrfs*42 frameshift in trans with a very rare missense variant (in families 2 and 4) on the other allele (Table 2 and Figure S2). In family 3, we identified a pathogenic heterozygous splice-site PEX1 variant, c.1239+1G>T,23 also previously reported in individuals with a severe peroxisome-biogenesis disorder (PBD) and in trans with an ultra-rare missense variant. In family 5, the two affected individuals were compound heterozygous for a previously reported pathogenic c.821C>T (p.Pro274Leu)24 variant in PEX6 and an ultra-rare missense variant on the other allele. We did not perform WES in family 6, but Sanger sequence analysis of PEX1 and PEX6 in the affected twins identified a missense variant, c.1802G>A (p.Arg601Gln), in PEX6 and a single-nucleotide deletion predicted to result in a frameshift on the other allele. The c.1802G>A (p.Arg601Gln) variant has been reported previously20,23 and is associated with milder PDB phenotypes (data not shown). The variants identified in the individuals with HS and not previously reported were absent in all public databases and in-house databases including 770 exomes. Two missense variants, PEX1 c.1742G>C and PEX6 c.1930C>T, were present in the ExAC Browser at a MAF of <0.000033. All variants were confirmed by Sanger sequencing (Figure S2) and predicted to be damaging (Table 2). The variants segregated with the

Table 2. Variants Found in PEX1 and PEX6 in Individuals with HS

| Family | Gene   | Variant          | Type of Variant (DNA Level) | Amino Acid Change | SIFT   | PolyPhen-2       | MutationTaster | Reference                           |
|--------|--------|------------------|----------------------------|-------------------|--------|------------------|----------------|-------------------------------------|
| 1      | PEX1   | c.3750G>A        | nonsense SNV               | p.Trp1250*        | deleterious | probably damaging | disease causing | not published                      |
| 2      | PEX1   | c.2114T>G        | non-synonymous SNV         | p.Leu705Trp       | tolerated | probably damaging | disease causing | not published                      |
| 2 and 4| PEX1   | c.2097dup        | single-nucleotide insertion| p.Ile700Tyrfs*42  | NA      | NA               | disease causing | Collins and Gould22                 |
| 3 and 4| PEX1   | c.1742G>C        | non-synonymous SNV         | p.Arg581Pro       | deleterious | probably damaging | disease causing | ExAC: 1 in 121,398 alleles          |
| 3      | PEX6   | c.1239+1G>T      | splice-altering SNV        | –                 | NA      | NA               | disease causing | Yik et al.23                        |
| 5      | PEX6   | c.821C>T         | non-synonymous SNV         | p.Pro274Leu       | deleterious | benign            | disease causing | Steinberg et al.24                 |
| 5      | PEX6   | c.1930C>T        | non-synonymous SNV         | p.Arg644Trp       | deleterious | probably damaging | disease causing | ExAC: 4 in 121,396 alleles          |
| 6      | PEX6   | c.1802G>A        | non-synonymous SNV         | p.Arg601Gln       | deleterious | probably damaging | disease causing | Yik et al.23                        |
| 6      | PEX6   | c.1841del        | single-nucleotide deletion | p.Leu614Argfs*5   | NA      | NA               | disease causing | not published                      |

For each variant, its predicted pathogenetic effect is stated, and in the last column, it is indicated whether the variant has been described before. "Not published" indicates that it has not been published and is also absent from the ExAC database. The following abbreviation is used: NA, not applicable.

*Ensembl: ENST00000248633 or GenBank: NM_000466.2.
**HUGO nomenclature is based on Ensembl: ENST00000248633 or GenBank: NM_000466.2, which is dissimilar to the protein nomenclature (p.Ile700Tyrfs*41) used in Collins and Gould.22
†Ensembl: ENST00000304611 or GenBank: NM_000287.3.
disease phenotype, such that all affected individuals had biallelic variants in PEX1 or PEX6, whereas unaffected siblings and parents were heterozygous or wild-type for the variants. In families 7 and 8, we did not identify a putative candidate gene, and no variants were identified in PEX1, PEX6, or any of the other currently known PEX genes.25

Peroxisomal Parameters of Individuals with HS
The finding of biallelic variants in PEX1 and PEX6 in the six HS-affected families suggested a peroxisomal defect in the affected individuals. We therefore analyzed plasma and erythrocytes from affected individuals with PEX1 and PEX6 variants (F1-II:3, F5-II:2, and F5-II:3) for peroxisomal parameters. These were all within the normal range and did not indicate peroxisomal dysfunction. Moreover, we also did not identify any significant peroxisomal biochemical aberrations in cultured skin fibroblasts from affected individuals F1-II:3 and F5-II:2 (Tables 3 and 4). However, previous studies have shown that individuals with very mild PBDs do not necessarily demonstrate significant biochemical abnormalities in plasma and/or fibroblasts.26,27 Thus, on the basis of these biochemical findings, we could not exclude a peroxisomal defect. Other evidence of clinical effect due to peroxisome dysfunction was not investigated because there was no clinical indication.

Peroxisomal Studies in the Cultured Fibroblasts of Individuals with HS
Because it is often a more sensitive indicator for a mild PBD,25,28 we also analyzed the peroxisomal phenotype in the cultured fibroblasts at 37°C from individuals F1-II:3 and F5-II:2 by immunofluorescence (IF) microscopy by using antibodies against the membrane protein PMP70 (ABCD3) and the peroxisomal matrix protein catalase (Figure 3). For both cell lines, we observed a so-called “mosaic” peroxisomal pattern. We saw different types of cells, including cells with normal peroxisomal staining, cells with a reduced number of peroxisomes, and cells with only peroxisomal membrane remnants, referred to as “ghosts,” but no import of matrix protein. Peroxisomal mosaicism has been described previously for hypomorphic variants in PEX1 and PEX6 and is typically associated with mild PBDs.29,30 Consistent with cells displaying peroxisomal mosaicism, the cells from individuals F1-II:3 and F5-II:2 showed a more severe peroxisomal phenotype when cultured at an elevated temperature (40°C), and the vast majority of cells lacked catalase-positive peroxisomes (Figure 3). Electron microscopy of fibroblasts cultured at normal temperature revealed no striking ultra-structural abnormalities of peroxisomes (Figure S3).

PEX-cDNA-Transfection Complementation Assay
To confirm that the variants found in PEX1 and PEX6 are the cause of the aberrant peroxisomal phenotype, we performed a genetic complementation assay. We cultured fibroblasts of individuals F1-II:3 and F5-II:2 at 40°C and then transfected these with control PEX1 and PEX6 cDNAs. The introduction of PEX1 cDNA rescued the impaired peroxisome biogenesis in cells

| Table 3. Peroxisomal Parameters in Blood of Individuals with HS |
|------------------|------------------|------------------|
|                  | Family 1          | Family 5          |
|                  | II:3             | II:2             | II:3             |
| Plasma           |                  |                  |                  |
| VLCFA concentration (µmol/l) | C22:0 | 47.57 (40–119)  | NA               | NA               |
|                  | C24:0            | 38.97 (33–84)  | NA               | NA               |
|                  | C26:0            | 0.67 (0.45–1.32) | 3.91 (0.3–4)    | 3.47 (0.3–4)    |
| VLCFA ratio      | C24:0/C22:0      | 0.82 (0.57–0.92) | 0.77 (0.35–1.1) | 0.76 (0.35–1.1) |
|                  | C26:0/C22:0      | 0.01 (0–0.02)   | 0.026 (0.003–0.033) | 0.026 (0.003–0.033) |
| Bile acids (µmol/l) | DHCA            | 0 (0)           | NA               | NA               |
|                  | THCA             | 0 (0–0.1)       | NA               | NA               |
| Phytanic acid (µmol/l) |            | 2.8 (0.5–9.9)  | 5.25 (0–16)     | 8.1 (0–16)      |
| Pristanic acid (µmol/l) |            | 0.4 (0.1–3)   | 0.18 (0–5)      | 1 (0–5)         |
| Pipecolic acid (µmol/l) |            | 1.5 (0.1–7)   | NA               | NA               |
| Erythrocytes     |                  |                  |                  |
| Plasmalogens (%) | C16:0 DMA        | 7.7 (6.8–11.9) | 4.61 (4.8–12)  | 5.27 (4.8–12)  |
|                  | C18:0 DMA        | 17.5 (10.6–24.9)| 9.3 (8.9–27)   | 10.7 (8.9–27)  |

Numbers in parentheses indicate the normal range according to the labs in which the assays were performed. Parameters were determined in different labs and reflect different reference values. Abbreviations are as follows: DHCA, 3β,7α-dihydroxycholestanolic acid; DMA, dimethylacetal; NA, not assessed; THCA, 3α,7α,12α-trihydroxycholestanolic acid; VLCFA, very-long-chain fatty acid.
from individual F1-II:3, and PEX6 cDNA rescued peroxisome biogenesis in cells from individual F5-II:2 (complementation in 14% or 20% of cells, respectively; Figure S4). These results confirm that the variants in PEX1 and PEX6 are the cause of the peroxisomal defects in the individuals with HS.

To determine whether, and to which degree, each of the identified PEX1 and PEX6 variants affect peroxisome biogenesis, we co-transfected cDNAs harboring the different variants with fluorescent peroxisomal marker EGFP-SKL into cells completely deficient in PEX1 or PEX6. We compared the capability of these cDNAs to functionally complement the peroxisome-deficient cells to the complementation capability of the control cDNAs (Figure 4). Because PEX1 c.1239+1G>T causes a splicing defect and consequently does not produce a functional protein, we did not test it in the complementation assay. Transfection with the constructs containing the previously reported pathogenic variants PEX1 c.2097dup and PEX6 c.821C>T, as well as the variant PEX6 c.1841del, resulted in no (0%) to minimal (1.3%) functional complementation. These results are consistent with the severe clinical presentation and complete absence of import of peroxisomal matrix protein in individuals homozygous for these variants (data not shown). In contrast, transfection with the other variants resulted in rescue of peroxisomal biogenesis in between 23% and 58% of cells, indicating that these variants are associated with significant residual activity. Thus, all affected individuals possess at least one PEX variant with residual activity in peroxisomal biogenesis.

**Discussion**

PEX1 and PEX6 encode two interacting proteins that belong to the peroxisomal import machinery and that are involved in the shuttling of PEX5, the cytosolic receptor for peroxisomal matrix proteins.32,33 Biallelic pathogenic variants in PEX1, PEX5, PEX6, or any of the other 11 PEX genes result in a PBD, which is characterized by defective peroxisome assembly due to impaired import of proteins into the peroxisomal matrix or membrane.25 Peroxisomes are found in virtually all human cells and play a crucial role in a number of metabolic pathways.34 PBDs usually have a severe progressive multi-systemic clinical presentation, including developmental delay, seizures, SNHL, retinopathy, peripheral neuropathy, leukodystrophy, and skeletal, craniofacial, and liver abnormalities.35–37 Dependent on the underlying genetic defect, however, the clinical presentation and survival of individuals with a PBD can show a wide variability ranging from the severe, early-childhood lethal Zellweger syndrome to milder phenotypes, including isolated progressive ataxia.26,27 To reflect this clinical variability, the PBDs are often referred to as Zellweger spectrum disorders.19 Accounting for 60% and 16%, respectively, of diagnosed cases,19 mutations in PEX1 and PEX6 represent the most common causes of PBDs. Our combined findings show that HS is caused by compound heterozygosity for a loss-of-function allele and a hypomorphic allele in PEX1 or PEX6. Alternatively, in family 1 a homozygous hypomorphic allele also results in the HS phenotype. Consequently, although PBD-affected individuals with mild or normal

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**Table 4. Peroxisomal Parameters in Primary Skin Fibroblasts of Individuals with HS**

| Parameter                       | Control Range           | Zellweger Spectrum Range | Family 1 II:3 | Family 5 II:2 |
|---------------------------------|-------------------------|--------------------------|---------------|---------------|
| VLCFA concentration (µmol/l)    | C22:0                   | 3.84–10.2                | 2.36–5.59     | 4.43          | 3.67          |
|                                 | C24:0                   | 7.76–17.66               | 5.41–13.39    | 8.36          | 8.58          |
|                                 | C26:0                   | 0.18–0.38                | 0.59–3.38     | 0.24          | 0.22          |
| VLCFA ratio                     | C24:0/C22:0             | 1.55–2.3                 | 2.08–3.4      | 1.89          | 2.34          |
|                                 | C26:0/C22:0             | 0.03–0.07                | 0.11–1.17     | 0.05          | 0.06          |
| DHAPAT activity [µmol/(2h × mg protein)] | 5.4–10.6               | 0.1–0.9                  | 7.1           | 14            |
| ACOX1 immunoblot                | 72 kDa                  | +                        | +             | +             | +             |
|                                 | 52 kDa                  | +                        | –             | +             | +             |
|                                 | 20 kDa                  | +                        | –             | +             | +             |
| Thiolase immunoblot             | 44 kDa                  | –                        | +             | –             | –             |
|                                 | 41 kDa                  | +                        | –             | +             | +             |
| α oxidation activity [µmol/(h × mg protein)] | phytanic acid substrate | 28–95                    | 0–10          | 34            | 39            |
| β oxidation activity [µmol/(h × mg protein)] | C16:0 substrate        | 3,330–7,790              | 3,330–7,790   | 7,162         | 5,959         |
|                                 | C26:0 substrate        | 800–2,040                | 50–350        | 1,088         | 1,013         |
|                                 | pristanic acid substrate| 790–1,690                | 0–30          | 1,092         | 756           |

Abbreviations are as follows: ACOX1, acyl-CoA oxidase I; DHAPAT, dihydroxyacetonephosphate acyltransferase; VLCFA, very-long-chain fatty acid.
peroxisome functions in blood and fibroblasts and normal intellect have been described before, HS represents a discrete phenotypic entity at the mildest end of the PBD clinical spectrum. The characteristic presentation overlaps some of the clinical features observed in affected individuals with PBDs. Indeed, SNHL is a common feature of PBDs, and tooth and nail abnormalities have been described in PBD-affected individuals with prolonged survival, but always in association with additional and more severe features. Importantly, in contrast to individuals with PBDs at the severe end of the clinical spectrum, the individuals with HS showed no identifiable dysmorphic or additional neurological features. Other evidence of clinical effect due to peroxisome dysfunction was not investigated, given that there was no clinical indication. However, future assessments of individuals with HS should consider other features well described in individuals with PBDs, including brain imaging, testing for adrenal insufficiency, liver-function tests, and clinical assessment for evidence of (progressive) peripheral neuropathy. Compound heterozygosity of the hypomorphic PEX6 c.1802G>A allele has been reported previously in seven individuals with a Zellweger spectrum disorder. In all reported individuals, the p.Arg601Gln allele was in trans with a severe PEX6 allele that, when homozygous or in trans with another severe PEX6 allele, causes a severe peroxisomal phenotype (six individuals) or that is predicted to be deleterious (one individual). For three of the seven individuals, who were diagnosed at an advanced adult age, studies in fibroblasts revealed peroxisomal mosaicism at 37°C and slightly elevated C26:C22 levels (unpublished results). Other peroxisomal parameters were normal. No clinical data are available for determining whether these individuals showed a HS-like phenotype. However, these findings suggest that the PEX6 c.1802G>A allele is a risk allele for mild PBD when in trans with a severe PEX6 allele. Because

![Figure 3](image-url)

**Figure 3. Peroxisomal Phenotype of Cells from Individuals with HS**

Skin fibroblasts of individuals F1-III:3 and F5-II:2 and control fibroblasts were cultured at 37°C or 40°C and immunolabeled for PMP70 (ABCD3) (red, peroxisomal membrane protein) and catalase (green, peroxisomal matrix protein). In cells of individuals with HS, the staining at 37°C revealed a mosaic pattern showing cells with a normal peroxisomal phenotype (positive for both PMP70 and catalase) and cells with import-incompetent peroxisomes ("ghosts," positive for PMP70 and negative for catalase). At 40°C, the vast majority of these cells showed import-incompetent peroxisomes (positive for PMP70 and negative for catalase). Control cells showed a normal peroxisomal phenotype in all conditions.
the PEX6 c.1802G>A allele has a frequency of 0.41% in the European population (see ExAC Browser in the Web Resources), we expect that future WES studies will identify additional individuals who have a mild PBD due to compound heterozygosity of the PEX6 c.1802G>A allele and a severe PEX6 allele and who have not been suspected of or analyzed for a peroxisomal disorder on the basis of clinical diagnosis.

Because standard biochemical screening of blood for evidence of a peroxisomal disorder would not have provided a diagnosis in the individuals with HS, our study used a genomic approach to diagnose a rare inborn error of metabolism. Our findings could also be relevant for the development of future therapy for PBDs, because they indicate that partial restoration of the function of altered PEX proteins would lead to a phenotype consistent with HS. Notably, the normal intellectual development and lack of severe hepatic and neurological impairment should be instructive in the expected outcomes of future therapeutic trials. Because of the SNHL and retinal pigmentation, HS is also an important differential diagnosis for Usher syndrome (MIM: 276900). Our study allows precise molecular differentiation of the two diagnoses and indicates that all individuals with SNHL and retinal pigmentation are candidates for mutation analysis of PEX genes. Our data indicate that HS is a clinically and genetically heterogeneous condition due to biallelic variants in PEX1 or PEX6. We did not identify PEX variants in families 7 or 8, nor did we find another genetic cause. However, there are phenotypic differences between these families and those in whom PEX1 or PEX6 variants were identified. Although amelogenesis imperfecta was present in all affected individuals in families 7 and 8, the SNHL in the individuals in family 7 was less severe than in the other affected individuals. In the previously reported affected individual in family 8, the SNHL was unilateral, whereas it was bilateral in all the other affected individuals. In addition, only individual F8-II:1 presented with subtle Beau’s lines, individuals in family 7 did not show any nail abnormalities, and ocular features were not present in any of the individuals in families 7 or 8. So, these phenotypic differences might account for the lack of PEX1 or PEX6 mutations, indicating both clinical and genetic heterogeneity.

The development of genomic medicine has stimulated an active debate about the interpretation of sequence variants and the challenges of pleiotropy. Our data highlight the complexity of the clinical interpretation of genomic data by showing that different mutations in PEX1 and PEX6 result in strikingly different clinical outcomes. In addition, the results of this study further emphasize the power of functional laboratory tests in the evaluation of rare variants in known disease-associated genes.

Supplemental Data

Supplemental Data include four figures and can be found with this article online at http://dx.doi.org/10.1016/j.ajhg.2015.08.011.

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Web Resources

The URLs for data presented herein are as follows:

1000 Genomes, http://www.1000genomes.org/
ANNOVAR, http://www.openbioinformatics.org/annovar/
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