

Loss-of-Function Ferrochelatase and Gain-of-Function Erythroid-Specific 5-Aminolevulinate Synthase Mutations Causing Erythropoietic Protoporphyrria and X-Linked Protoporphyrria in North American Patients Reveal Novel Mutations and a High Prevalence of X-Linked Protoporphyrria

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Erythropoietic protoporphyria (EPP) and X-linked protoporphyria (XLP) are inborn errors of heme biosynthesis with the same phenotype but resulting from autosomal recessive loss-of-function mutations in the ferrochelatase (*FECH*) gene and gain-of-function mutations in the X-linked erythroid-specific 5-aminolevulinate synthase (*ALAS2*) gene, respectively. The EPP phenotype is characterized by acute, painful, cutaneous photosensitivity and elevated erythrocyte protoporphyrin levels. We report the *FECH* and *ALAS2* mutations in 155 unrelated North American patients with the EPP phenotype. *FECH* sequencing and dosage analyses identified 140 patients with EPP: 134 with one loss-of-function allele and the common IVS3-48T>C low expression allele, three with two loss-of-function mutations and three with one loss-of-function mutation and two low expression alleles. There were 48 previously reported and 23 novel *FECH* mutations. The remaining 15 probands had *ALAS2* gain-of-function mutations causing XLP: 13 with the previously reported deletion, c.1706_1709delAGTG, and two with novel mutations, c.1734delG and c.1642C>T(p.Q548X). Notably, XLP represented ~10% of EPP phenotype patients in North America, two to five times more than in Western Europe. XLP males had twofold higher erythrocyte protoporphyrin levels than EPP patients, predisposing to more severe photosensitivity and liver disease. Identification of XLP patients permits accurate diagnosis and counseling of at-risk relatives and asymptomatic heterozygotes.

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INTRODUCTION

Erythropoietic protoporphyria (EPP) (OMIM 177000), the most common ery-

thropoietic porphyria, is an autosomal recessive inborn error of heme biosynthesis because of the deficient activity (<35% of

normal) of ferrochelatase (*FECH*) (E.C.4.99.1.1), the enzyme in the heme biosynthetic pathway that inserts ferrous iron into protoporphyrin to form heme. The enzyme deficiency results in the accumulation of protoporphyrin, primarily in marrow reticulocytes (1–3). Accumulated protoporphyrin is released from the marrow and circulating erythrocytes into the plasma, leading to its deposition in vascular cells as well as the liver, which is primarily responsible for its excretion.

Clinically, EPP is characterized by the onset of acute cutaneous photosensitivity

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early in childhood (1,4). Exposure to sunlight or ultraviolet radiation for even a few minutes causes extreme pain that can be excruciating and is not relieved by narcotic analgesics. After prolonged sunlight exposure, the pain may persist for several days and full recovery may take a week. The exposed skin often becomes erythematous and edematous, but does not blister, although scarring may occur with multiple exposures.

Pathogenetically, sunlight photoactivates protoporphyrin in the superficial vessels, presumably triggering a singlet oxygen-mediated free radical reaction that results in cutaneous damage and painful symptoms. Protoporphyrin is insoluble in water and is not secreted into the urine, but is normally secreted into the bile. Excess biliary protoporphyrin is presumably responsible for the increased prevalence of gallstones in EPP patients. Accumulated hepatic protoporphyrin can precipitate in hepatocytes and bile canaliculi, causing hepatotoxicity, decreased bile formation and flow, and cholestatic liver failure in some patients (5).

The diagnosis of the EPP phenotype is based on the characteristically painful acute cutaneous photosensitivity and biochemical confirmation of markedly elevated levels of erythrocyte protoporphyrin. There is no specific treatment, so patients rigorously avoid sunlight exposure, which markedly limits their quality of life (6). Liver transplantation has been undertaken in patients with liver failure, and bone marrow transplantation has prevented recurrence (7,8). Recently, clinical trials of an α -melanocyte stimulating hormone analog (afamelanotide) to darken the skin have resulted in decreased photosensitivity, increased pain-free exposure to sunlight and improved quality-of-life for these patients (9,10).

FECH mutation analyses in Western Europeans with the EPP phenotype revealed that the phenotype is genetically heterogeneous (11,12). In Europe, about 92% of biochemically diagnosed patients with the EPP phenotype had autosomal recessive EPP with either two *FECH*

loss-of-function mutations or, most often, one *FECH* loss-of-function mutation and the common *FECH* low expression mutation, *IVS3-48T>C*. The *IVS3-48T>C* allele creates a cryptic upstream acceptor site in intron 3 that modulates the alternative splicing of the normal *FECH* mRNA, generating about 25% of wild-type *FECH* mRNA and an mRNA that contains 63 intronic bases resulting in premature truncation and nonsense-mediated decay (13). The *IVS3-48T>C* allele frequency varies widely with ethnicity and is ~10% among Western European Caucasians (14). Over 160 *FECH* loss-of-function mutations have been reported in the Human Gene Mutation Database (www.hgmd.cf.ac.uk) (15).

Whatley *et al.* (11) reported that 2–4% of unrelated UK patients with the EPP phenotype did not have *FECH* mutations but had a two- or four-base deletion in the last exon of the X-linked erythroid-specific 5-aminolevulinic synthase (*ALAS2*) gene, either c.1699-1700delAT (p.M567EfsX2; designated Δ AT) or c.1706-1709delAGTG (p.E569GfsX24; designated Δ AGTG) (16). Both deletions altered the carboxyl-terminal region of the enzyme such that its activity and/or stability were increased. These gain-of-function mutations markedly increased the erythroid synthesis of 5-aminolevulinic acid (ALA) resulting in the erythroid accumulation of protoporphyrin and the EPP phenotype (16). This X-linked disorder is termed X-linked protoporphyria (XLP). Of note, 6% of the UK patients with the EPP phenotype did not have *FECH* or *ALAS2* mutations (11), indicating the occurrence of additional genetic heterogeneity.

In EPP, >90% of the elevated erythrocyte protoporphyrin is not zinc chelated. In contrast, affected XLP males have ~15–50% zinc-chelated protoporphyrin, despite normal *FECH* activity. The heterozygous females in the European XLP families had increased erythrocyte protoporphyrin and photosensitivity (16), although mutation-confirmed XLP heterozygotes can be asymptomatic, as expected for X-linked traits (16,17). In

addition, about 17% of XLP patients had overt liver disease (16), higher than that reported for EPP (16,18). To date, in limited studies of unrelated North American patients with EPP (19), only two with XLP were reported (20). Here, we report the *FECH* and *ALAS2* mutations in 155 unrelated North American patients with the EPP phenotype. Of these, ~90% (140 of 155) had *FECH* mutations causing EPP, including 23 novel loss-of-function lesions. All 15 remaining patients had *ALAS2* exon 11 mutations causing XLP, including 13 with the Δ AGTG deletion and two with novel exon 11 lesions, a deletion and a nonsense mutation. Thus, about 10% of unrelated North American patients with the EPP phenotype had XLP, five times that reported for UK EPP phenotype patients (11), and at least twice that found in Western European patients with the EPP phenotype (12). Clearly, genotyping patients with the EPP phenotype is important to identify at-risk XLP family members, particularly the asymptomatic heterozygous females who can transmit the disease-causing genes to their offspring.

MATERIALS AND METHODS

Patients with the EPP phenotype were referred for mutation analyses by the Porphyrins Consortium of the National Institutes of Health (NIH)-sponsored Rare Disease Clinical Research Network or by their physicians. Peripheral bloods or cheek swabs were obtained with informed consent. Erythrocyte protoporphyrin levels were determined by commercial laboratories or at the Porphyrin Center (University of Texas Medical Branch, Galveston, TX, USA). Clinical information, including medical histories and family pedigrees, symptoms and results of biochemical testing were obtained. *FECH* and *ALAS2* exon 11 mutation analyses were performed by the Porphyrin Molecular Diagnostic Testing Laboratory at the Mount Sinai School of Medicine in New York City.

Genomic DNAs were extracted using the QIAGEN[®] DNA Isolation Kit (QIAGEN, Valencia, CA, USA). For *FECH*

gene analyses, ~500 base pairs (bp) of the promoter, all exons and ~30–40 bp adjacent intronic or flanking sequences were amplified by using the GeneAmp PCR System 9700 (PE Applied Biosystems, Foster City, CA, USA) with previously obtained oligonucleotide primers (14) with the following modification. Primer 9R had a known intronic polymorphism (rs58628398 in the NCBI dbSNP [single nucleotide polymorphism database]), which deleted seven bases (AGGACAC) in 10% of European, 5% of African and 0% of East Asian chromosomes (21). Therefore, a new 9R primer was synthesized to avoid the polymorphism (Supplementary Table S1). The IVS3-48T>C region was also amplified with exon 4, and the T>C genotype was determined (22). For *ALAS2*, the entire exon 11 coding region and ~30–40 bp intronic and 3'-untranslated sequence were amplified by using the indicated polymerase chain reaction (PCR) primers (Supplementary Table S1). Sequences were analyzed by using Sequencer version 4.8 software (Gene Codes) (reference sequences: *FECH* transcript NM_000140.3; *FECH* gene: NG_008175; *ALAS2* transcript GenBank NM_000032.4). Gene dosage analysis to detect large deletions that included entire exons was performed as described (23). Splicing mutations were analyzed by using SplicePort to predict effects on *FECH* function (<http://spliceport.cbcb.umd.edu/>) (24).

Additional genotyping included *FECH* intragenic single nucleotide polymorphisms (SNPs). These SNPs were 5'-252A>G, tightly linked to IVS1-23C>T and IVS3-48T>C, as well as c.798C>G (P266P) linked to c.921A>G. The unlinked SNPs included c.287G>A (p.R96Q), IVS6-198C>T, IVS8+94T>A, IVS10-64G>A and 3'+248C>T.

Structural Characterization of *FECH* Missense Mutations

The potential impact of each novel missense mutation was evaluated by visual inspection of the environment of the altered residue in the *FECH* crystal structures of the catalytic structural states,

that is, "open" (PDB #2QD3), "closed" with bound substrate (2QD1) and "release" with bound product (2QD2). Attention was directed to identify residues for which available structural data indicate a role in catalysis and/or macrocycle binding. The propensity for the observed mutations to be tolerated or not were assessed by the SIFT (<http://sift-dna.org>) and PolyPhen programs (25,26). Low scores indicate deleterious mutations predicted by the SIFT program, whereas high scores (1.0 = max) indicate deleterious mutations predicted by the PolyPhen program. The PyMol program was used to produce a figure highlighting the locations of the novel *FECH* mutations (The PyMOL Molecular Graphics System, Version 1.2; Schrödinger LLC, New York, NY, USA).

All supplementary materials are available online at www.molmed.org.

RESULTS

Mutation Analysis of the *FECH* Gene in Patients with the EPP Phenotype

Genomic DNAs from 155 North American patients with the EPP phenotype (that is, photosensitivity and markedly elevated erythrocyte protoporphyrin levels) were analyzed for *FECH* loss-of-function mutations and the IVS3-48T>C allele by PCR amplification and sequencing. Of the 155 patients, 140 (~90%) had *FECH* loss-of-function mutations causing autosomal recessive EPP. All 15 remaining patients were found to have *ALAS2* exon 11 gain-of-function mutations causing XLP (see below). Among the 140 patients with EPP, three had two loss-of-function *FECH* alleles: genotypes c.348A>C (p.S151F)/c.1001C>T (p.P334L), c.643C>T (p.R219W)/c.901_902delTG (p.W301Afs*22) and c.820G>A (p.D274N)/c.913G>T (a splicing mutation). Three patients were homozygous for the low-activity allele but also had a second mutation on one allele [genotypes: c.485G>A(G162E) + IVS3-48T>C/IVS3-48T>C, c.577C>T(Q193X)+IVS3-48T>C/IVS3-48T>C and c.794T>G(L265R)+IVS3-48T>C/IVS3-48T>C]. The remaining 132

patients had a loss-of-function mutation on one allele and the low-expression IVS3-48T>C lesion on the other. Altogether, 71 *FECH* mutations were detected, including 48 previously reported and 23 novel mutations (Figures 1 and 2, Table 1).

Previously Reported *FECH* Mutations

The previously reported *FECH* mutations, identified in 111 patients, are indicated in Figures 1 and 2 and included 9 missense, 5 nonsense, 12 small insertions or deletions, 18 splicing mutations and 4 large deletions, with the latter 4 detected by SNP homozygosity and identified by breakpoint sequencing (Figure 2). Among the most common of the previously reported mutations were the following: IVS3+2T>G found in 13 patients, c.1232G>A (p.C411G) in 12 patients, the large deletion del10,379insTTCA (23) in 6 patients and splicing defects IVS1+1G>A, c.913G>T and c.1135A>T each in 6 patients. These mutations were all present *in trans* with the IVS3-48T>C allele. Of interest, one patient had two splicing mutations (IVS5+1G>T and IVS5-3C>T) on the same allele and the IVS3-48T>C mutation on the other, as defined by family studies. These two mutations were reported previously, but singly, in different probands (15).

Reverse transcriptase (RT)-PCR studies of the previously suspected splicing mutation c.1077G>A (p.E359E) (27) demonstrated that the G>A transition in the last nucleotide of exon 9 altered the 5' acceptor site consensus sequence from GAGgtaaat to GAAgtaaat caused 100% exon 9 skipping (data not shown), as predicted by the SplicePort program (24).

Novel *FECH* Mutations

The 23 novel *FECH* loss-of-function mutations (Table 1, Figures 1 and 2) included eight missense and nine nonsense mutations, three splice site lesions, one small deletion and two complex large deletion/insertions. Of the 17 novel missense and nonsense mutations, one (p.R215W) occurred at a cytosine guanine dinucleotide (CpG), a

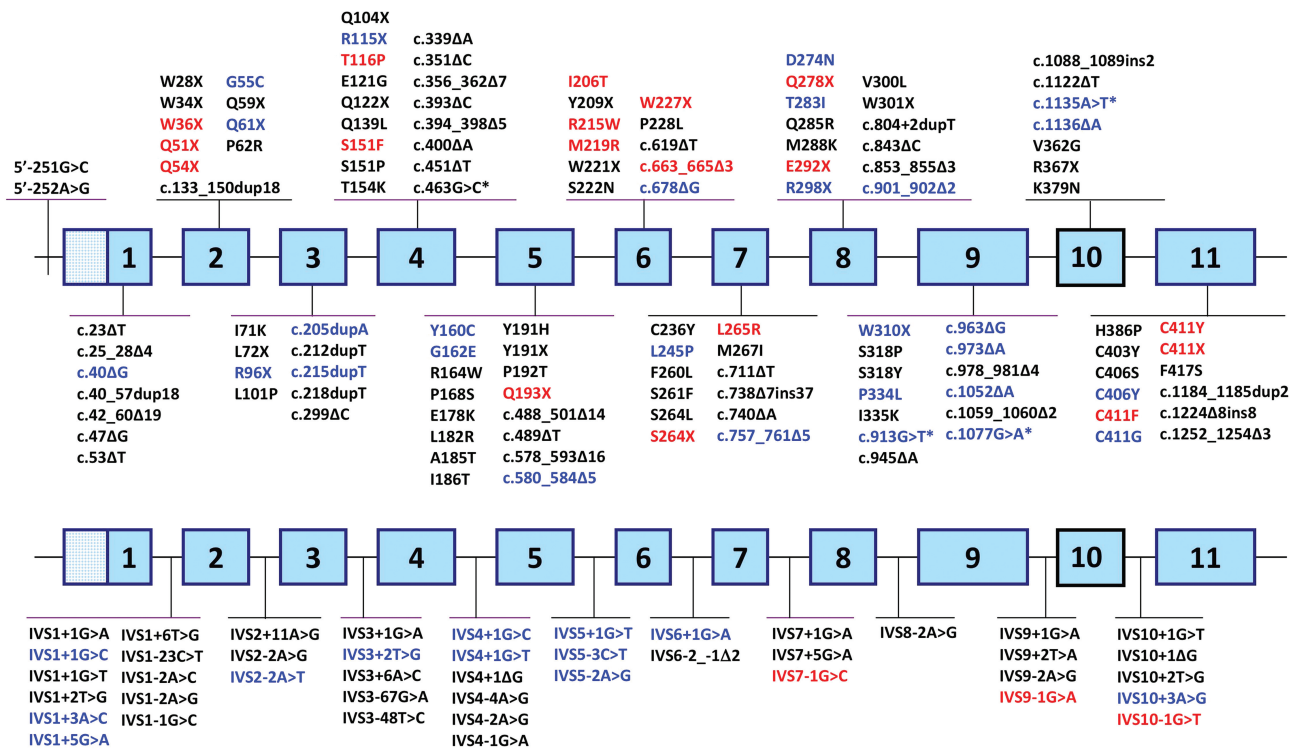


Figure 1. *FECH* loss-of-function mutations causing autosomal-recessive EPP. Mutations in black and blue type were previously reported and listed in the Human Gene Mutation Database, release date 29 June 2012. Previously published mutations in blue were those found in the patients of this study. Novel mutations are indicated in red. Asterisks indicate splicing mutations caused by exonic mutations. The gene depicted in the top line shows exonic mutations, and mutations in intronic sequences are shown on the bottom line.

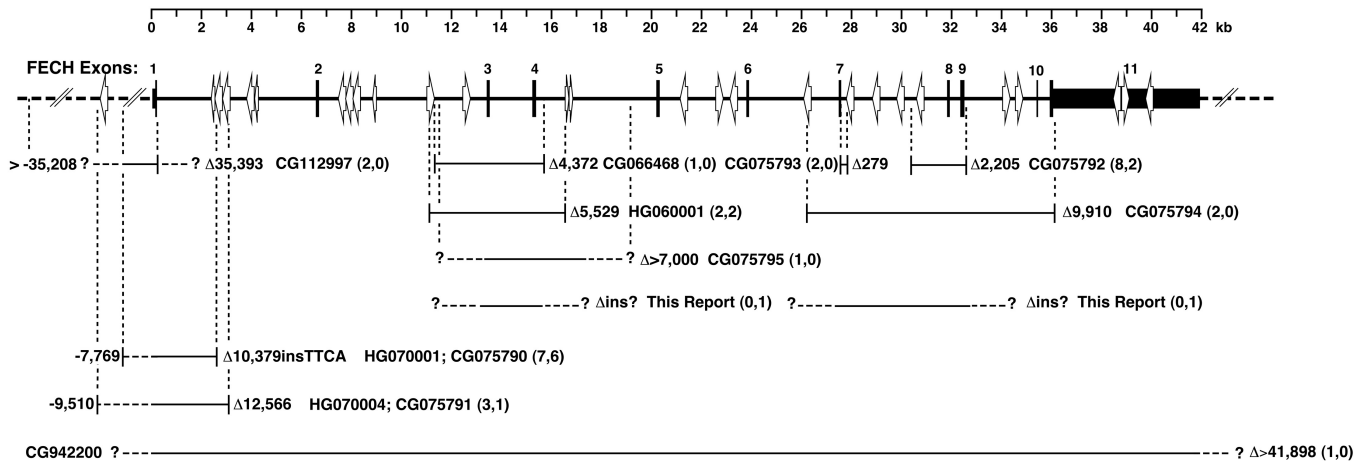


Figure 2. Large *FECH* deletions and complex deletion/insertion rearrangements causing EPP. The Human Gene Mutation Database (HGMD) accession numbers are provided, and the extent of the deletion is listed to the right of each deleted region. In parentheses, the number of patients previously reported with the deletion is followed by the number of patients with the deletion in the present study. Novel mutations are those with only one incidence in this study. Question marks indicate the unknown positions of the breakpoints. *Alu* sequences and their orientations are indicated with open arrows. The published breakpoints have been mapped to the *FECH* sequence provided in GenBank accession number NG_008175 from the February 2009 HG-19 human genome assembly. HGMD accession numbers for mutations with identical breakpoints have been listed together with the respective deletion region.

Table 1. Novel *FECH* loss-of-function mutations in EPP patients.

Subject	Sex	Novel <i>FECH</i> mutations ^a (novel allele)	Known <i>FECH</i> mutations ^a (other allele)
Missense mutations			
1	M	p.T116P (c.346A>C)	IVS3-48T>C
2	F	p.S151F (c.452C>T)	p.P334L (c.1001C>T)
3	M	p.I206T (c.617T>C)	IVS3-48T>C
4	M	p.R215W (c.643C>T)	p.W301Afs*22 (c.901_902delTTG)
5	F	p.M219R (c.656T>G)	IVS3-48T>C
6	M	p.L265R (c.794T>G) and IVS3-48T>C (c.315-48T>C)	IVS3-48T>C
7	M	p.C411Y (c.1232G>A)	IVS3-48T>C
8	M	p.C411F (c.1232G>T)	IVS3-48T>C
Nonsense mutations			
9	M	p.W36X (c.107G>A)	IVS3-48T>C
10	M/F	p.Q51X (c.151C>T)	IVS3-48T>C
11	F	p.Q54X (c.160C>T)	IVS3-48T>C
12	M	p.Q193X (c.577C>T) and IVS3-48T>C (c.315-48T>C)	IVS3-48T>C
13	F	p.W227X (c.681G>A)	IVS3-48T>C
14	M/F	p.S264X (c.791C>G)	IVS3-48T>C
15	M/F	p.Q278X (c.832C>T)	IVS3-48T>C
16	M	p.E292X (c.874G>T)	IVS3-48T>C
17	M	p.C411X (c.1233C>A)	IVS3-48T>C
Splice site mutations			
18	M	IVS7-1G>C (c.805-1G>C)	IVS3-48T>C
19	M/F	IVS9-1G>A (c.1078-1G>A)	IVS3-48T>C
20	M	IVS10-1G>T (c.1138-1G>T)	IVS3-48T>C
Deletion mutations			
21	F	p.W221_S222delinsC (c.663_665delGAG)	IVS3-48T>C
22	F	del exon 3-4 + ins (breakpoints unknown)	IVS3-48T>C
23	F	del exon 7-9 + ins (breakpoints unknown)	IVS3-48T>C

^aProtein change (cDNA change).

known hot spot for mutation (28), of which there are 38 in the *FECH* coding sequence.

Of the eight novel missense mutations, six occurred at residues having 96–100% amino acid conservation in 26 species from insects to humans (Supplementary Table S2). The novel *FECH* missense mutations were assessed on the *FECH* 3D crystal structure (29) (Figure 3) to determine their predicted affects on the structure and/or function of the enzyme (Table 2). Although none was predicted to be tolerated by both the SIFT and PolyPhen programs (Table 2), on the basis of the scores for both programs, the

R215W mutation was relatively more tolerated, consistent with its low conservation (27%).

The novel small deletion c.663_665delGAG (p.W221_S222delinsC) (Figure 1) lost three bases across two codons, thereby substituting a cysteine for tryptophan in codon 221 and deleting the serine codon 222, after which it retained the normal amino acid sequence. Two “sequencing cryptic” deletions were identified by dosage analyses. These studies revealed two additional novel large rearrangements in three patients, involving exons 3 and 4 or exons 7, 8 and 9 (Figure 2).

FECH Single Nucleotide Polymorphisms

Of the 906 single nucleotide polymorphisms (SNPs) in the *FECH* gene recorded in the dbSNP database, about 125 were common (minor allele frequency [MAF] >5%). Of these, 11 were characterized in the EPP phenotype patients, since they were present in the promoter and exonic amplicons that were sequenced. These included rs17963905 (5'-252A>G) and rs7243988 (IVS1-23C>T), which are usually linked to the *IVS3-48T>C* mutation. Five other polymorphisms were relatively frequent (minor allele frequencies of 14–31% in Caucasians), including three exonic polymorphisms: rs1041951 (c.287G>A, p.R96Q), rs536765 (c.798C>G; P266P) and rs536506 (c.921A>G, p.P307P). Monitoring these SNPs for apparent homozygosity was used to detect possible “sequencing cryptic” *FECH* gene deletions that were further delineated by *FECH* gene dosage analyses (23).

ALAS2 Mutations Causing XLP

Of the 15 unrelated probands who did not have two *FECH* mutations causing EPP, 13 had the previously reported four-base *ALAS2* deletion, ΔAGTG (16); one had a novel c.1734delG (p.Q581SfsX13) single-base deletion (designated ΔG); and one had a novel nonsense mutation, c1642C>T (p.Q548X). With the exception of an African-American and a Pacific Islander, the ΔAGTG XLP patients had ancestors of Western European origin, particularly from the United Kingdom. The novel ΔG mutation altered the seven carboxyl-terminal wild-type amino acids (QYVTTYA) to SMSPPMP and extended the enzyme polypeptide sequence by five residues (EKPA A) (Figure 4). Of note, the last 14 altered carboxyl-terminal residues of the elongated ΔAGTG and ΔG mutations were identical. The c.1642C>T (p.Q548X) nonsense mutation altered *ALAS2* codon 548 from CAG encoding a glutamine to TAG, a chain termination signal that prematurely deleted the 40 wild-type carboxyl-terminal residues (548–587) of the enzyme polypeptide.

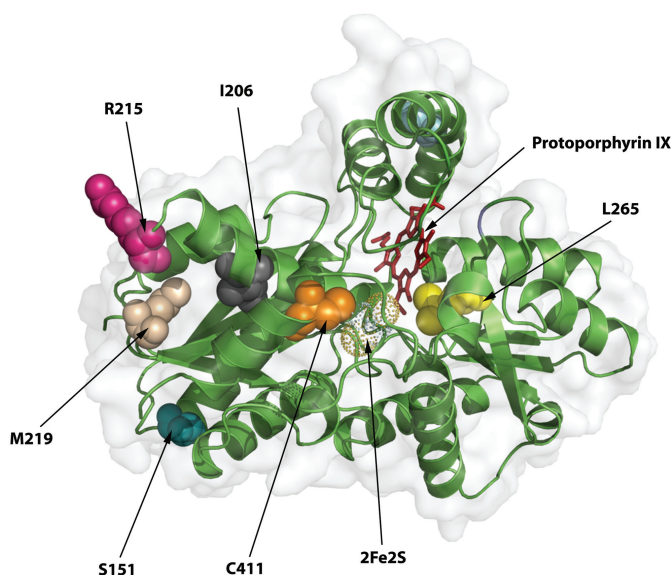


Figure 3. FECH three-dimensional structure and the location of novel deleterious *FECH* mutations. The crystal structure of one of the homodimeric FECH monomers is shown by using secondary structure cartoons in green and the surface in light gray. The view is from the side of the protein that is membrane-bound, looking into the active site region containing protoporphyrin IX and the iron-sulfur cluster. The residues changed by the novel mutations are shown as space-filling spheres and are the indicated native residues, not the mutations.

EPP and XLP Genotype and Protoporphyrin Levels

Assuming that the erythrocyte and plasma protoporphyrin concentrations can be correlated with (or predict) disease severity, the available erythrocyte and plasma levels in EPP and XLP patients were stratified by mutation type (Table 3). For 120 EPP patients, the mean and median erythrocyte protoporphyrin levels were 2,220 and 1,740 $\mu\text{g}/\text{dL}$, respectively, and ranged from 407 to 8,240 $\mu\text{g}/\text{dL}$, with the highest levels mostly in patients with liver disease or liver transplants (Table 3, Supplementary Figure S1). The mean erythrocyte protoporphyrin levels in males and females were 2,113 and 2,344 $\mu\text{g}/\text{dL}$, respectively, which were similar. There were no effects of age on erythrocyte protoporphyrin levels (data not shown), but there was a significant increase in risk of liver disease above a threshold of 2,500–3,000 $\mu\text{g}/\text{dL}$. The mean percentage of erythrocyte metal-

Table 2. Predicted effect of novel missense mutations on FECH structure and function.

Mutation	SIFT ^a score/ tolerated	PolyPhen ^a score/ tolerated	Predicted effect
p.T116P	0.19/Yes	1.000/No	T116 is in a critical position at the end of a small helix that moves to close the active site upon porphyrin binding. Replacement with proline will make this rigid and/or create a kink so that the protein should have subnormal function. Arginine 115, which is important for substrate binding, is located within this helix and its position is likely to be affected by the T116P-induced conformation change.
p.S151F	0.17/Yes	1.000/No	This serine residue is on the backside of the enzyme, away from the active site, and is predicted by SIFT to be tolerated. However, it is located at the end of a helical structure, where it forms a hydrogen bond with threonine 154, stabilizing a tight β turn. The substitution of the bulky hydrophobic benzyl ring of the mutant phenylalanine for the smaller hydrophilic serine might cause protein instability because of misfolding.
p.I206T	0.00/No	1.000/No	This is an internal residue that is in the middle of a helix. The mutation would disrupt local structure and alter overall tertiary structure.
p.R215W	0.19/Yes	0.985/No	This is a surface residue located on the side of the protein. It is not clear how it would affect the structure of the protein, but if this region is involved in protein-protein interactions, substitution of a large bulky residue for a charged residue should have an impact on such an association.
p.M219R	0.01/No	0.998/No	This residue is at the end of a β sheet and would cause structural perturbations because of the addition of the bulky guanido moiety with its positive charge.
p.L265R	0.00/No	1.000/No	L265 is located adjacent to the 2-vinyl group in the substrate-bound structures, and substitution of an arginine would be expected to have a significant negative impact on macrocycle binding in the active site.
p.C411F/Y	0.01/No	1.000/No	The 411F and Y mutation would not have the (2Fe-2S) cluster required for enzyme activity, since C411 is one of the four essential ligands.

^aSIFT and PolyPhen are *in silico* phenotyping tools, as described in Materials and Methods.

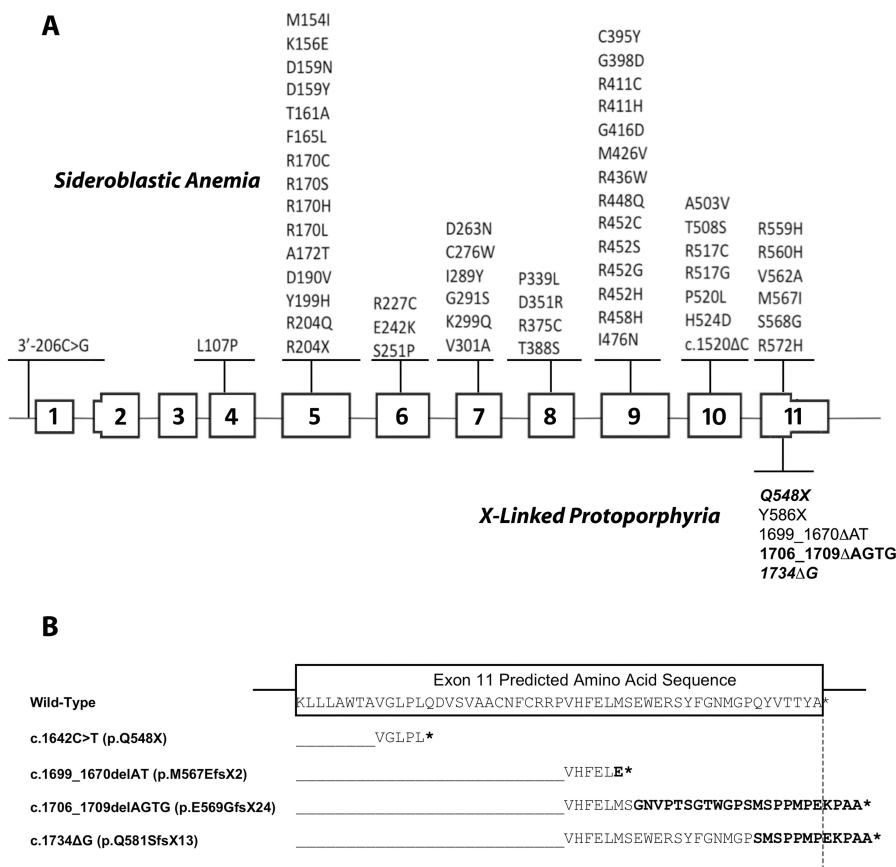


Figure 4. XLP mutations and gene structure. (A) *ALAS2* loss-of-function and gain-of-function mutations causing X-linked sideroblastic anemia and EPP. Mutations in black and blue are those reported in the Human Gene Mutation Database, release date 29 June 2012. The mutations in bold are published mutations also found in patients of this study. The mutations in bold italics are the novel mutations. (B) Variations in the C-terminal sequences of the XLP gain-of-function mutations. The partial wild-type exon 11 *ALAS2* sequence is boxed. The hybrid sequences of wild-type *ALAS2* and the sequence following the mutation sites are aligned below the wild-type sequence. The termination codons are denoted by asterisks. Note that for the pE569GfsX24 and pQ581SfsX13 mutations, the last 12 mutated residues are identical to each other and different from the wild-type.

free protoporphyrin for the different *FECH* genotypes in Table 3 ranged from 92 to 97% of the total erythrocyte protoporphyrin with a range of 84 to 98%. The 25 patients with missense mutations (including unrelated patients with the same mutation), which may have residual activity, had the lowest mean erythrocyte protoporphyrin concentration (1,430 ± 660 μg/dL), whereas patients with a splicing defect, nonsense mutation or small or large deletion had higher mean concentrations ranging from 2,170 to

3,580 μg/dL. Similarly, the mean plasma protoporphyrin levels were lowest in patients with missense mutations (5.7 μg/dL) and higher in the other *FECH* genotype groups (8.7–14.0 μg/dL). Of note, the patient with two missense alleles (S151F and P334L) and the patient with two low-activity alleles (one having a missense mutation [G162E]) had concentrations of 3,210 and 7,750 μg/dL, respectively.

For eight XLP male patients with the ΔAGTG mutation, the mean erythrocyte protoporphyrin level was 5,153 μg/dL and

the range was from 2,293 to 10,650 μg/dL, about twofold higher than that in the 120 EPP patients (Supplementary Figure S1). Notably, the mean level of free erythrocyte protoporphyrin for patients with the ΔAGTG mutation was 67%, indicating an average ratio of 67/34 for free versus zinc-chelated protoporphyrin. Of note, the Q548X proband had a ratio of 92/8, which was similar to those of EPP patients.

DISCUSSION

Fifty years ago, Magnus *et al.* (30) reported the first patients with EPP, described the photo-induced cutaneous manifestation as “solar urticaria” and identified the presence of accumulated protoporphyrin in the patient’s erythrocytes and stools. Their description of the photo-induced cutaneous manifestations and the protoporphyrin accumulation not only characterized the clinical and biochemical phenotype of the disease, but remains today the basis for its diagnosis (1). Subsequently, the partially deficient *FECH* activity (<35% of normal in cultured cells and < 25% of normal in bone marrow) was shown in EPP patients (31,32).

Early studies of the inheritance of EPP suggested that the disease was either an autosomal-dominant trait with reduced penetrance (33) or an autosomal-recessive disorder (34). Initially, EPP was thought to be an autosomal-dominant disorder when the *FECH* cDNA and genomic sequences were isolated and characterized (35,36) and the causative mutations were identified (15,23,37,38)

(www.hgmd.cf.ac.uk). Over 90% of symptomatic patients had only one loss-of-function *FECH* mutation supporting autosomal-dominant inheritance. The small percentage of EPP patients with two *FECH* mutations was classified as having “recessive” EPP. The fact that symptomatic patients had 15–35% of normal activity was a dilemma that led to various hypotheses concerning the molecular genetics of the disease (1).

The inheritance was clarified by Gouya *et al.* (13,14,22,39), who discov-

Table 3. EPP and XLP genotypes and their erythrocyte and plasma protoporphyrin (PROTO) concentrations.

Genotype	Total erythrocyte PROTO ($\mu\text{g}/\text{dL}$)		Free erythrocyte PROTO (%)		Plasma porphyrin ^a ($\mu\text{g}/\text{dL}$)	
	Mean \pm SD (range)	n	Mean \pm SD (range)	n	Mean \pm SD (range)	n
EPP						
Missense/IVS3-48T>C	1,430 \pm 660 (407–2,940)	25	92 \pm 4 (84–97)	16	5.7 \pm 6.4 (0.7–28)	17
Splicing/IVS3-48T>C	2,170 \pm 1,550 ^a (562–7,170)	44	94 \pm 3 (86–98)	27	8.7 \pm 9.9 (0.5–44)	29
Nonsense/IVS3-48T>C	2,350 \pm 1,320 ^b (845–5,190)	16	95 \pm 3 (87–97)	11	11.0 \pm 11.8 (0.5–47)	14
Small deletion/IVS3-48T>C	3,580 \pm 2,410 ^c (1,060–8,240)	10	97 \pm 1 (96, 97 ^d)	2	14.0 \pm 9.8 (6.2–28)	4
Large deletion/IVS3-48T>C	2,610 \pm 1,200 (1,230–5,160)	13	95 \pm 4 (84–98)	12	10.3 \pm 7.7 (0.2–26)	12
p.L265Re+IVS3-48T>C/IVS3-48T>C	7,750 ^e	1		1		1
p.Q193X+IVS3-48T>C/IVS3-48T>C	926	1	92	1	11.7	1
p.S151F/p.P344L	3,210	1		1	25.5	1
p.R215W/p.W301Afs*22	664	1		1		1
p.D274N/c.913G>T (splicing defect)	2,520	1		1		1
XLP						
c.1706delAGTG (<i>FECH</i> Wt/Wt)	5,153 \pm 3,162 (2,293–10,650)	8	66 \pm 8.3 (50–78)	8	11.7 \pm 7.2 (5.8–23)	5
c.1706delAGTG (<i>FECH</i> IVS3-48T>C/Wt)	3,891	1	75	1	4.7	1

^aFive patients with liver transplants had protoporphyrin levels from 2,680 to 6,690. If excluded, mean \pm standard deviation (SD) was 1,980 \pm 1,460.

^bOne patient with a liver transplant and two with liver disease had protoporphyrin levels from 2,840 to 5,190. If excluded, mean \pm SD was 1,900 \pm 864.

^cThree patients with liver transplants had protoporphyrin levels from 5,410 to 8,240. If excluded, mean \pm SD was 2,770 \pm 1,770.

^dTwo specific values.

^eThe patient had liver disease with cirrhosis.

ered the common intronic *FECH* low expression allele. With the identification of this allele, it was quickly realized that over 90% of EPP patients had a loss-of-function *FECH* mutation on one allele and the IVS3-48T>C mutation on the other (14), consistent with autosomal-recessive inheritance.

To further complicate the inheritance, it was noted that loss-of-function *FECH* mutations were absent in about 5–10% of symptomatic patients with elevated erythrocyte protoporphyrin. The inheritance pattern in some of these patients suggested X-linkage. While loss-of-function mutations in the *ALAS2* gene cause X-linked sideroblastic anemia (Figure 4) (40), Whatley *et al.* in 2008 (16) reported two deletions in exon 11 of the *ALAS2* gene that resulted in frameshift mutations and increased *ALAS2* activity. Notably, both affected and asymptomatic heterozygotes for the same *ALAS2* gain-of-function mutation were reported, demonstrating that it is not a dominant disorder (17).

Here, we report mutation analyses of 155 unrelated North American patients

with the EPP phenotype that identified 140 with two *FECH* mutations causing EPP and 15 with *ALAS2* mutations causing XLP. These studies identified 23 novel *FECH* loss-of-function mutations and two novel *ALAS2* exon 11 mutations: a small deletion (ΔG) and a nonsense mutation (p.Q548X). These novel mutations add to the previously reported 164 *FECH* (Figures 1 and 2, Table 1) and two *ALAS2* exon 11 mutations causing XLP (and over 60 loss-of-function *ALAS2* mutations that cause X-linked sideroblastic anemia) (Figure 4). Of the 140 patients with EPP, most (96%) had a loss-of-function *FECH* allele and the IVS3-48T>C allele, similar to the findings in Western Europe, where ~95% of EPP patients had a loss-of-function mutation and the IVS3-48T>C allele, and only ~5% had two loss-of-function alleles (11,16).

Of the North American EPP patients with the 23 novel *FECH* mutations, eight had missense mutations that mostly replaced highly conserved amino acids. Structural modeling predicted that these mutations would likely alter *FECH* func-

tion and/or stability (Table 2, Figure 3). The *FECH* coding sequence contains 38 CpGs, which are mutational hot spots for C→T or G→A transitions (29). However, of the 60 previously published missense and nonsense mutations, only eight occurred at CpGs, whereas only one of the 17 novel base substitutions reported here was at a CpG. Thus, EPP mutations do not appear to favor these hot spots. Of the 44 previously reported and three novel splicing mutations, 1–12 occurred in each of the 10 introns (Figure 1).

Gene dosage analyses were required to identify four additional large *FECH* deletions involving entire exons and not detectable by sequence analysis, thereby confirming the EPP diagnoses in these patients. In total, we found about 8% of the EPP phenotype patients had large deletions, similar to the frequency of large deletions (10%) reported in European patients (11,23).

Of the 15 patients with the EPP phenotype who did not have *FECH* mutations, all had *ALAS2* exon 11 gain-of-function mutations. Thirteen had the ΔAGTG

deletion and two had novel mutations: a single-base deletion (c.1734delG), which altered the nine carboxyl-terminal amino acids and elongated the enzyme polypeptide by five residues, and a nonsense mutation (p.Q548X), which deleted the terminal 40 amino acids (Figure 4). For these mutations, the mechanism(s) by which the structural changes are responsible for their increased enzymatic activity and/or stability remain to be determined. It has been hypothesized that the carboxy-terminal amino acids may normally restrict release of the product, since it was demonstrated that numerous hyperactive variants created by mutations in this region resulted in increased rates of conformational change leading to increased ALA synthesis (41). It is notable that the human *ALAS2* exon 11 encodes 54 terminal amino acids, of which the terminal 30 amino acids are only present in vertebrates and that this region was recently shown to be required for binding of the precursor enzyme succinyl-CoA synthetase (42). In addition, other mutations in this region can cause X-linked sideroblastic anemia (Figure 4) by alterations in enzyme stability *in vivo* (43). Thus, these terminal residues play a critical role in the regulation of erythroid heme biosynthesis in humans.

Notably, the three exon 11 deletions (Δ AGTG, Δ AT and Δ G) all occur within a span of 36 nucleotides. The Δ AGTG deletion was at a direct repeat (AGTG AGTG), which only occurred at this location in the coding region of the human *ALAS2* gene. The Δ AT mutation occurs three bases upstream from Δ AGTG, and the Δ G mutation occurs downstream in a region containing four guanine residues followed by four cytosines. The mutational mechanism responsible for the Δ AGTG and Δ G mutations may be homologous, but unequal, with crossing-over of their repeats during recombination. Comparison of the distribution of the erythrocyte protoporphyrin levels in the available EPP and XLP patients (Supplementary Figure S1) indicated a higher concentration in the XLP compared with EPP patients. The finding of higher pro-

toporphyrin levels may correlate with disease severity (particularly increased cutaneous photosensitivity and risk for liver damage, with the latter due to delivery of excess hepatotoxic protoporphyrin to the liver for biliary excretion). Thus, physicians should closely monitor liver function in EPP and especially in XLP patients. Notably, the erythrocyte zinc protoporphyrin in XLP males and most XLP females averaged >30% of the total protoporphyrin, compared with ~5% for EPP patients, a previously noted finding of diagnostic value (16).

Although all 155 North American patients with the EPP phenotype were found to have a causative *FECH* or *ALAS2* mutation, that was not the case in the Western European patients (16), where *FECH* or *ALAS2* mutations were not detected in ~5% of patients with the EPP phenotype. This finding suggested that other gene defects may cause the EPP phenotype (11,12). Notably, the percentage of patients with the EPP phenotype who had XLP in our North American cohort was ~10%, which was two- to fivefold greater than that in the Western European studies (11,16).

CONCLUSION

ALAS2 mutation analysis should be undertaken in patients with the EPP phenotype who have no loss-of-function *FECH* mutations. Identification of the XLP patients will permit accurate diagnosis of at-risk family members, particularly asymptomatic heterozygotes who have a 50% risk for affected male offspring. Moreover, the finding of higher mean erythrocyte protoporphyrin levels in XLP probands compared with EPP probands, suggests that XLP probands may have a more severe disease course (particularly, an increased risk of liver damage).

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