Abstract: The first described patients with pyridox(am)ine 5'-phosphate oxidase deficiency all had neonatal onset seizures that did not respond to treatment with pyridoxine but responded to treatment with pyridoxal 5'-phosphate. Our data suggest, however, that the clinical spectrum of pyridox(am)ine 5'-phosphate oxidase deficiency is much broader than has been reported in the literature. Sequencing of the PNPO gene was undertaken for a cohort of 82 individuals who had shown a reduction in frequency and severity of seizures in response to pyridoxine or pyridoxal 5'-phosphate. Novel sequence changes were studied using a new cell-free expression system and a mass spectrometry-based assay for pyridoxamine phosphate oxidase. Three groups of patients with PNPO mutations that had reduced enzyme activity were identified: (i) patients with neonatal onset seizures responding to pyridoxal 5'-phosphate (n = 6); (ii) a patient with infantile spasms (onset 5 months) responsive to pyridoxal 5'-phosphate (n = 1); and (iii) patients with seizures starting under 3 months of age responding to pyridoxine (n = 8). Data suggest that certain genotypes (R225H/C and D33V) are more likely to result in seizures that respond to treatment with pyridoxine. Other mutations seem to be associated with infertility, miscarriage and prematurity. However, the situation is clearly complex with the same combination of mutations being seen in patients who responded and did not respond to pyridoxine. It is possible that pyridoxine responsiveness in PNPO deficiency is affected by prematurity and age at the time of the therapeutic trial. Other additional factors that are likely to influence treatment response and outcome include riboflavin status and how well the foetus has been supplied with vitamin B6 by the mother. For some patients there was a worsening of symptoms on changing from pyridoxine to pyridoxal 5'-phosphate. Many of the mutations in PNPO affected residues involved in binding flavin mononucleotide or pyridoxal 5'-phosphate and many of them showed residual enzyme activity. One sequence change (R116Q), predicted to affect flavin mononucleotide binding and binding of the two PNPO dimers, and with high residual activity was found in Groups (ii) and (iii). This sequence change has been reported in the 1000 Genomes project suggesting it could be a polymorphism but alternatively it could be a common mutation, perhaps responsible for the susceptibility locus for genetic generalized epilepsy on 17q21.32 (close to rs72823592). We believe the reduction in PNPO activity and B6-responsive epilepsy in the patients reported here indicates that it contributes to the pathogenesis of epilepsy.

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Epilepsy due to PNPO mutations: genotype, environment and treatment affect presentation and outcome

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The first described patients with pyridox(am)ine 5’-phosphate oxidase deficiency all had neonatal onset seizures that did not respond to treatment with pyridoxine but responded to treatment with pyridoxal 5’-phosphate. Our data suggest, however, that the clinical spectrum of pyridox(am)ine 5’-phosphate oxidase deficiency is much broader than has been reported in the literature. Sequencing of the PNPO gene was undertaken for a cohort of 82 individuals who had shown a reduction in frequency and severity of seizures in response to pyridoxine or pyridoxal 5’-phosphate. Novel sequence changes were studied using a new cell-free expression system and a mass spectrometry-based assay for pyridoxamine phosphate oxidase. Three groups of patients with
PNPO mutations that had reduced enzyme activity were identified: (i) patients with neonatal onset seizures responding to pyridoxal 5′-phosphate (n = 6); (ii) a patient with infantile spasms (onset 5 months) responsive to pyridoxal 5′-phosphate (n = 1); and (iii) patients with seizures starting under 3 months of age responding to pyridoxine (n = 8). Data suggest that certain genotypes (R225H/C and D33V) are more likely to result in seizures that to respond to treatment with pyridoxine. Other mutations seem to be associated with infertility, miscarriage and prematurity. However, the situation is clearly complex with the same combination of mutations being seen in patients who responded and did not respond to pyridoxine. It is possible that pyridoxine responsiveness in PNPO deficiency is affected by prematurity and age at the time of the therapeutic trial. Other additional factors that are likely to influence treatment response and outcome include riboflavin status and how well the foetus has been supplied with vitamin B₆ by the mother. For some patients there was a worsening of symptoms on changing from pyridoxine to pyridoxal 5′-phosphate. Many of the mutations in PNPO affected residues involved in binding flavin mononucleotide or pyridoxal 5′-phosphate and many of them showed residual enzyme activity. One sequence change (R116Q), predicted to affect flavin mononucleotide binding and binding of the two PNPO dimers, and with high residual activity was found in Groups (ii) and (iii). This sequence change has been reported in the 1000 Genomes project suggesting it could be a polymorphism but alternatively it could be a common mutation, perhaps responsible for the susceptibility locus for genetic generalized epilepsy on 17q21.32 (close to rs72823592). We believe the reduction in PNPO activity and B₆-responsive epilepsy in the patients reported here indicates that it contributes to the pathogenesis of epilepsy.

**Keywords:** pyridoxal 5′-phosphate (PLP); pyridoxine; pyridox(am)ine 5′-phosphate oxidase (PNPO); seizures; epilepsy

**Abbreviations:** FMN = flavin mononucleotide; PLP = pyridoxal 5′-phosphate; PMP = pyridoxamine phosphate

### Introduction

Pyridox(am)ine phosphate oxidase (PNPO, EC 1.4.3.5) is a flavin mononucleotide (FMN)-dependent oxidase required for synthesis of pyridoxal 5′-phosphate (PLP) from pyridoxine (and its phosphate, PNP, and glucoside) and from pyridoxamine (and its phosphate, PMP) in the diet. It is also required for recycling PMP to regenerate PLP, the active cofactor. PLP is the cofactor for >140 enzyme-catalyzed reactions in man, including many involved in synthesis or degradation of amino acids or amines that serve as neurotransmitters or neuromodulators in the brain (Garcia-Cazorla et al., 2012; Mills et al., 2012).

Neonatal epileptic encephalopathy is characterized by the onset, shortly after birth, of drug-resistant seizures associated with severe neurological dysfunction that can be fatal. PNPO deficiency (OMIM 6032870) is an autosomal recessive inborn error of metabolism that leads to a seizure disorder, presenting in the newborn period (neonatal epileptic encephalopathy) or early infancy, that can be treated with pyridoxal 5′-phosphate but (classically) not pyridoxine. Mills et al. (2005) showed that mutations in PNPO led to reduced enzyme activity when expressed in Chinese hamster ovary cells.

A review by Garcia-Cazorla et al. (2012) of 16 cases with PNPO deficiency (eight families) revealed that clinical features included in utero seizures, 3/16; foetal distress before delivery, 5/16; premature birth, mean 32 weeks gestation; low Apgar scores/requiring intubation at delivery, 5/16; onset of seizures—in first 24 h, 11/14; between 24 h and 72 h, 2/14; between 72 h and 2 weeks 1/14; burst suppression pattern on EEG, 10/11; seizures completely resistant to antiepileptic drugs, 13/16; completely resistant to pyridoxine, 7/10; metabolic acidosis 6/16; raised blood lactate 8/16; distressing spasms (dystonia), 3/16; anaemia, 3/16; hepatomegaly, 3/16; abdominal distension, 2/16; and hypoglycaemia 2/16.

In the small number of patients tested, CSF PLP concentration was low (Ormazabal et al., 2008). However, other disorders can also lead to low CSF PLP (Footitt et al., 2011).

In addition to the small number of infants in the literature whose PLP-responsive neonatal epileptic encephalopathy has been shown to be a result of PNPO deficiency, a larger number of infants has been described for whom empirical clinical trials demonstrated that their severe epilepsy was better controlled with PLP than with pyridoxine (Wang et al., 2005). This suggests that PNPO deficiency might be a cause, not only of neonatal epileptic encephalopathy, but also of other later-onset seizure disorders.

A genome-wide association study of patients of European ancestry identified an important susceptibility locus for genetic generalized epilepsies as a whole at 17q21.32 (rs72823592), the closest gene to which is PNPO (EPICURE Consortium et al., 2012). This data suggests that mild PNPO deficiency could be a susceptibility factor for genetic generalized epilepsies presenting at various ages.

Mutations in PNPO known to be associated with neonatal epileptic encephalopathy for which expression studies have shown reduced enzyme activity are: IVS3-1g> a, X262Q, R229W (Mills et al., 2005) and R95H (Khayat et al., 2008). Other probable neonatal epileptic encephalopathy-causing PNPO mutations include R95C, D33V, c.246delT (Hoffmann et al., 2007), A174X (Ruiz et al., 2008), R225C (Veerapandiyan et al., 2011) and G118R (Pearl et al., 2012).

The effects of R229W on PNPO catalytic function and crystal structure have been studied by Musayev et al. (2009). This variant is 850-fold less efficient than the wild-type enzyme because of a 192-fold decrease in pyridoxine 5′-phosphate affinity and a 4.5-fold decrease in catalytic activity. There is also a 50-fold reduction in affinity for the FMN cofactor. The decrease in affinity for
pyridoxine 5'-phosphate suggests that, for this mutation, significantly increased synthesis of PLP might be achieved by high dose pyridoxine. In fact, in the premature infants affected by the R229W mutation, administration of pyridoxine did lead to partial improvement in clonic contractions and lip-smacking automatisms; in contrast, patients harbouring mutations with no residual activity showed no response to pyridoxine. Musayev et al. (2009) suggested that, for the R229W mutation at least, another treatment likely to be beneficial would be riboflavin, which would supply extra FMN and partially overcome the effect of reduced FMN binding.

Thus, a picture is emerging of PNPO sequence variations that produce epilepsy that only responds to treatment with PLP, through variations that may respond to pyridoxine and/or riboflavin to PNPO variations that might be insufficient to produce epilepsy alone but might do so in concert with other factors determining brain PLP levels such as genes influencing blood PLP levels [e.g. the TNSALP gene (now known as ALPL) encoding tissue non-specific alkaline phosphatase (Hazra et al., 2009; Tanaka et al., 2009)] and dietary intakes of B6 vitamers and riboflavin.

In this study we looked for PNPO sequence variations in children with epilepsy that responded to treatment with pyridoxine or PLP. We characterized the phenotypic spectrum of PNPO deficiency further and demonstrated that some children with PNPO deficiency respond to treatment with pyridoxine. To determine the effect of sequence variations on PNPO enzyme activity we set up a new, cell-free system for expression of the mutant proteins and a new mass spectrometry-based enzyme assay for measurement of PNPO activity.

Materials and methods

Patients

This study was approved by the Ethics Committee of UCL Institute of Child Health and Great Ormond Street Hospital (04/Q0508/81). Patients included satisfied the following criteria: (i) improved seizure control following administration of pyridoxine or PLP; and (ii) exclusion of antitin deficiency by measurement of urinary α-aminoisobutyric semialdehyde and/or sequencing of ALDH7A1 (Mills et al., 2010). Detailed histories and the questionnaire sent to clinicians are available in the Supplementary material.

Chemical reagents

All chemicals unless mentioned specifically were from Sigma Aldrich. D2 PLP was kindly supplied by Professor Coburn, Department of Chemistry, Indiana University-Purdue University, Fort Wayne, Indiana, USA.

PNPO sequencing

Mutation analysis of genomic DNA and complementary DNA was as described previously (Mills et al., 2005).

Site-directed mutagenesis of PNPO and expression studies

Wild-type PNPO complementary DNA had been cloned into pSP72 previously (Mills et al., 2005). Site-directed mutagenesis was carried out using the QuikChange XL Site-Directed Mutagenesis Kit (Stratagene), according to manufacturer’s instructions. The complementary DNA was then amplified using ProofStart™ DNA polymerase (Qiagen) using primers detailed in Supplementary Table 1. EcorI digestion products were subcloned into the pT7CFE1-CHis expression vector (Thermo Scientific). PNPO was expressed using the Thermo Scientific 1-Step Human Coupled IVT Kit according to manufacturer’s instructions; this expression system uses a HeLa cell lysate. The reaction was incubated for 6 h at 30°C. PNPO enzyme activity was measured with PMP as substrate. Expressed protein (8 μl) was incubated at 37°C in the dark in 20 mM potassium phosphate (pH 7.6) containing 2.5 μM PNP and 1.5 μM FMN in a final volume of 120 μl. The reaction was stopped by addition of 120 μl 0.3 N trichloroacetic acid containing d2-PLP and incubated on ice for 30 min before centrifugation (10000 rpm, 10 min at 4°C). The supernatant containing the B6 vitamers was analysed by HPLC-MS/MS (Footitt et al., 2013). Enzyme activity was calculated as pmol PLP synthesized/mg protein/min and expressed as a percentage of the activity obtained with the wild-type enzyme.

Results

Mutations

Sequencing of the PNPO gene was undertaken for 82 individuals that had shown some reduction in frequency and severity of seizures in response to pyridoxine or PLP. Mutations were identified in PNPO for 15 patients (Table 1) from 14 families. Five of these were novel; three missense mutations and two deletions/insertions. The novel missense mutations E120K, P213S and R225H have not been reported as polymorphisms in Ensembl (http://www.ensembl.org). All novel missense mutations were tested for pathogenicity using both SIFT (Sim et al., 2012) and the PolyPhen web server (Ramensky et al. 2002) and were predicted to be damaging or probably damaging, respectively (Supplementary Table 2).

Previously reported mutations included D33V, R95H, R95C and R225C. E50K, which has been reported as a single nucleotide polymorphism (Mills et al., 2005), was detected in Patient 1. The sequence variant R116Q (rs 17679445) was detected in 5 of 15 patients. This is reported on Ensembl as a single nucleotide polymorphism in the majority of populations with the exception of Asian ancestry (Supplementary Table 3) with a prevalence of 4–10% depending on the subpopulation investigated. R116Q was found in some patients in combination with other mutations that we believe are pathogenic and which have not been reported previously in the general population. However, R116Q was the only sequence change detected in Patients 7 and 8. Comparison of the frequency of R116Q in our PNPO-deficient patients relative to 1000 Genomes allele frequencies (http://www.ensembl.org) suggests that R116Q is over-represented in our
Clinical phenotypes

Our series of PNPO patients has identified some new clinical presentations, including seizures with onset under 3 months responsive to pyridoxine and seizures with onset after 3 months responsive to PLP, suggesting that the phenotype of this disorder is much broader than reported previously. Patient clinical features are summarized in Table 2 and individual histories are given in the Supplementary material.

Five patients were identified with neonatal onset seizures responsive to PLP and one affected sibling was treated prophylactically, in utero and from birth (Patient 4). Immediate seizure control was seen upon administration of PLP in three of five of these patients and seizure control was achieved in Patients 6 and 3 after treatment with PLP for 12 h and 3 days, respectively. Patient 7 did not present with seizures until 5 months of age. Initial doses of PLP ranged from 10–85 mg/kg/day and maintenance doses range from 10–72 mg/kg/day. Individual treatment regimes are detailed in the Supplementary material.

To our surprise, 62% of patients in this series that had an onset of seizures under 3 months showed a dramatic response to treatment with pyridoxine (Patients 8–15); initial and maintenance doses ranging from 18–55 and 6–26 mg/kg/day, respectively.

Reported seizure types (Table 2) included clonic (45%), myoclonic jerks (55%), tonic (30%), generalized tonic-clonic (79%) and focal (45%). EEG abnormalities included burst suppression and hypsarhythmia (Table 2, Supplementary material). A cranial ultrasound at 5 h showed bilateral grade 3 intraventricular haemorrhages with associated moderate ventricular dilatation in one patient. The MRI findings (Supplementary material) varied from normal to extensive white matter oedema and a possible haemorrhage and changes reported as consistent with hypoxic ischaemic encephalopathy. Two patients on PLP treatment have persistently deranged liver function tests.

Expression studies

In vitro expression studies using the HeLa cell lysate system showed that wild-type PNPO activity was readily measurable. Several PNPO sequence variants that had been expressed previously using the Chinese hamster ovary cell system (R229W, X262Q and E50K) (Mills et al., 2005) were investigated using this more rapid approach so that we could compare the two expression systems. Using this new system we investigated the effects of R116Q on PNPO activity to determine if the single
nucleotide polymorphism is ‘possibly damaging’ as predicted by PolyPhen, or ‘tolerated’ as predicted by SIFT (Supplementary Table 2). The effects of R225H, R225C and D33V were also studied in more detail as these sequence variants had been found in various combinations with R116Q.

Characterization of wild-type PNPO activity and the various mutant constructs revealed a pre-steady state period where the rate of PLP synthesis was slower than that seen between 10–20 min. This was true for all constructs investigated and was also apparent when PNPO was overexpressed in Chinese hamster ovary cells previously (unpublished data).

Figure 1 shows that the presence of R229W, X262Q, R225H and R225C led to a reduction in PLP synthesis to 8–15% of the rate catalyzed by wild-type PNPO. The effect of D33V on PNPO activity was less dramatic with ~C245% PNPO activity being retained. E50K, which had no effect on PNPO activity when over-expressed in Chinese hamster ovary cells, resulted in decreased PNPO activity by ~C25% when compared to wild-type activity whereas the presence of R116Q decreased PNPO activity by 17%.

Discussion

Mutations
This article takes the number of published disease-causing mutations in PNPO to 14. However, if we include sequence variants reported previously to be single nucleotide polymorphisms and shown here to result in reduced PNPO activity i.e. E50K and
### Expression studies

Characterization of the PNPO wild-type and mutant proteins revealed a slower pre-steady state period (Fig. 1A). Ordinarily transient periods in enzyme catalyzed reactions only occur for a few seconds whilst the enzyme and substrate form a complex, however, the lag period here persisted for the first 5 min. PNPO has been shown to be a sluggish enzyme with a turnover number of only 0.19/s and 0.20/s for PNP and PMP, respectively and it is self-regulated by its product PLP (Musayev et al., 2003). PNPO is a dimeric enzyme that contains a non-catalytic site on each of the monomers that binds PLP tightly (Safo et al., 2005). Crystallographic studies suggested that a tunnel may exist between the active site and this secondary non-catalytic site acting to protect PLP from nucleophiles and channeling it to vitamin B6 apoenzymes (Safo et al., 2005). If the endogenous PLP in the cell free expression system (Time 0; Fig. 1A) is bound at the non-catalytic site, the active site would be open and the enzyme reaction would proceed with steady-state kinetics. However, if the endogenous PLP is tightly bound at the active site in the closed position, the substrate PMP would have to bind to an allosteric site to enable a conformational change to occur. This would allow the tightly bound PLP at the active site to be released and catalysis to proceed (Safo et al., 2005). This may explain the slower rate of reaction observed initially.

Comparison of PNPO activity for two mutations that have been over-expressed previously in Chinese hamster ovary cells, i.e. R229W and X262Q, to that obtained using the new more rapid cell-free expression system confirmed that these sequence variants are pathogenic. Previously these mutations produced ~30% and undetectable PNPO activity, respectively (Mills et al., 2005) whereas in the new system these mutations resulted in 15% and 12% residual activity, respectively. The discrepancy between these two systems is perhaps not surprising; the LC-MS/MS based system is more sensitive than the spectrophotometric assay and also allows for PLP product inhibition. E50K which had previously been shown to have no effect on PNPO activity when over-expressed in Chinese hamster ovary cells (Mills et al., 2005), did have 30% lower activity than wild-type. E50K has been identified (previously and in Patient 1) in cis with the splice site mutation c.364-1G>A (IVS3-1g>a). Previous expression studies concluded that c.364-1G>A is a pathogenic mutation (Mills et al., 2005) and that E50K was a polymorphism. E50 is conserved across mammalian species (Supplementary Fig. 2) and is predicted by SIFT and PolyPhen to have a ‘damaging effect’ and to be ‘probably damaging’ to protein activity, respectively (Supplementary Table 2). Whether this sequence variant—on its own or in combination with environmental factors—would prove sufficient to result in B6-responsive epilepsy remains to be seen.

The effects of R225H, R225C and D33V on PNPO activity were also investigated as these sequence variants were found in various combinations with R116Q. Expression of R225H and R225C dramatically reduced PNPO activity to 9% and 8% of wild-type activity, respectively (Fig. 1). The effect of D33V on PNPO activity was investigated as Kang et al., (2004) reported that deletion of the first 56N-terminal amino acid residues of human PNPO affected neither binding of coenzyme nor catalytic activity. In our expression system D33V dramatically reduced enzyme activity by ~60%. This was not surprising as D33V accounts for 15% of sequence changes found in PNPO patients to date. We conclude therefore that D33V is pathogenic.

![Figure 1](http://brain.oxfordjournals.org/)

**Figure 1** (A) PNPO activity (conversion of PMP to PLP) was measured by LC MS/MS and activity expressed as pmol PLP/mg protein. (B) Amount of PLP synthesized over a 40-min period by PNPO containing various mutations/sequence variants was compared with that of wild-type PNPO.
The presence of R116Q decreased PNPO activity by ~20% when compared with wild-type (Fig. 1). R116 is not only conserved in mammals but also in Escherichia coli (Supplementary Fig. 2). This residue contributes to one of two pairs of intersubunit salt-bridges binding the two monomeric PNPO subunits to constitute the functional dimer (Musayev et al., 2003). R116Q is also predicted to affect FMN binding. The prevalence of this sequence change within populations suggests that this polymorphism might be a common contributor or susceptibility allele to epilepsy that responds to treatment with PLP. R225H, R225C and D33V were present in tandem with R116Q in Patients 12, 14 and 15. We have yet to study these combinations. When two mutations occur in cis in the same allele in cystic fibrosis the combination may either improve or exacerbate the phenotype (Polizzi et al., 2011).

**Clinical phenotype**

Our data suggest that the clinical spectrum of PNPO deficiency is much broader than has been reported. The first described patients did not respond to treatment with pyridoxine; cessation of seizures only occurred with PLP. Recently, however, Pearl et al. (2012) reported a patient with PNPO mutations who showed a transient response to pyridoxine. Three groups of patients could be identified in this study with regard to their response to treatment: (i) patients that had neonatal onset seizures responding to treatment with PLP (n = 6); (ii) one child with infantile spasms (onset 5 months) responsive to PLP; and (iii) patients with seizures that responded to pyridoxine (n = 8). Patient 11, from Group (iii), had improved seizure control with a switch from pyridoxine to PLP treatment at the age of 19 years. However, for three patients in this group symptoms deteriorated with this change in treatment. The worsening of seizure control with attempting to switch to PLP therapy is noteworthy. High doses of PLP can cause seizures in experimental animals and in infants (Ishioka et al., 1995; Hammen et al., 1998). PNPO plays a role in controlling intracellular PLP levels through inhibition of the enzyme activity by high concentrations of PLP. It is possible that a mutant enzyme with residual activity could show impaired inhibition by PLP thus increasing the risk of development of toxic levels. Alternatively, it may be the build-up of another metabolite e.g. PMP that has an adverse effect on some patients on PLP treatment.

Patient 12 in Group (iii), at the age of 19 years, found that a regular morning aura could be prevented by taking a multivitamin preparation in addition to pyridoxine. The tablet contains 100 mg riboflavin (precursor of FMN, the cofactor for PNPO) and it is possible that this boosted residual PNPO activity. Patients 12 and 14 in Group (iii) had autistic features.

Patient 10, being treated with pyridoxine, developed a severe neuropathy with persistent loss of ankle jerks. This is the first patient with PNPO deficiency that we have encountered to develop a severe neuropathy. Neuropathy is well recognized in normal adults taking doses >200 mg/day of pyridoxine and we see it in some patients with antiquitin deficiency on similar doses. Analysis of plasma B6 vitamer levels in two antiquitin-deficient patients treated with pyridoxine and showing mild peripheral neuropathy, revealed that plasma PLP remained low whereas pyridoxal, pyridoxamine, PMP and pyridoxine were increased (Footitt et al., 2013). Animal experiments have suggested that pyridoxine may be more toxic to the PNS than other B6 vitamers (Levine et al., 2004).

Two patients in this series that are treated with PLP have persistently deranged liver function tests. Similar findings have been documented in a child with PNPO deficiency receiving a high dose (100 mg/kg/day) PLP treatment (Mills et al., 2012). A liver biopsy showed early cirrhosis, which was attributed to the PLP therapy. The PLP dose has now been reduced to 60 mg/kg/day and although the liver function tests have improved, they have not normalized and there is persistent evidence of hepatic fibrosis and portal hypertension. Liver toxicity secondary to high dose PLP (1000 mg/day) has also been reported for a child with homocystinuria (Yoshida et al., 1985). It is uncertain whether the cause of this toxicity is directly due to the high dose PLP or to degradation products of PLP that may form by photochemical reactions. Liver function should be monitored in all children treated with high doses of PLP.

Premature delivery (<37/40) has been documented in the majority of genetically confirmed PNPO-deficient cases reported in the literature with many patients also having low Apgar scores at delivery (often requiring intubation). In this series 54% of the patients were born at term and, although two patients required assistance with breathing (continuous positive airway pressure/mechanical ventilation) and one patient required resuscitation, most of the patients were in good condition at birth.

In the majority of reported cases, seizures commenced within the first hours of life. While seizures commenced in 10/14 of the individuals reported here at <24 h, three patients did not present with seizures until later (2, 3 and 8 weeks old, respectively) and Patient 7 did not present until 5 months of age (Table 3). Patient 7 remains seizure-free on a dose of PLP of 10 mg/kg/day; most of the other patients treated with PLP have required 2–10 times this dose.

A history of infertility should alert the clinician to a possible diagnosis of PNPO deficiency as heterozygous couples seem to have reduced rates of conception. Four of eight families reported previously have undergone several attempts at in vitro fertilization treatment and/or suffered early pregnancy losses. In this study four families reported miscarriages, treatment for infertility and/or a molar pregnancy.

As with many rare diseases, it is difficult to draw conclusions about the long-term outcome. If untreated, PNPO deficiency is usually fatal with a single child surviving with severe epilepsy and psychomotor delay to 3 years of age (Hoffmann et al., 2007). In the seven cases where treatment with PLP was initiated, only one child died and this was as a result of fungal sepsis (Ruiz et al., 2008). The other six survived with varying degrees of disability. Here 3 of 12 patients have marked motor delay; for Patients 11 and 13 this may relate to the duration of poor seizure control. Six of twelve patients show no evidence of any disability. Three patients reported here are now older than 21 years and the eldest patient is 41 and his neurological abnormalities are limited to dyslexia and Asperger’s syndrome.
Table 3  Possible effects of genotype and environmental factors on patient response to pyridoxine, prematurity and developmental outcome

<table>
<thead>
<tr>
<th>Group</th>
<th>Patient</th>
<th>Genotype</th>
<th>Born prematurely</th>
<th>Ethnicity</th>
<th>Mother given B6</th>
<th>Neonatal feeding</th>
<th>Seizure onset</th>
<th>Time taken to control seizures</th>
<th>Response to PN</th>
<th>Current age</th>
<th>Outcome</th>
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<tbody>
<tr>
<td>(i)</td>
<td>1</td>
<td>R95H + E50K; Splice errors(^a)</td>
<td>+</td>
<td>Asian</td>
<td>?</td>
<td>30 min</td>
<td>5 d</td>
<td>-</td>
<td>-</td>
<td>2 y 8 m</td>
<td>Global delay</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>D33V/D33V</td>
<td>-</td>
<td>Caucasian</td>
<td>B</td>
<td>6 h</td>
<td>8 w</td>
<td>-</td>
<td>-</td>
<td>4 y</td>
<td>Normal</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>P213S/P213S</td>
<td>-</td>
<td>Caucasian</td>
<td>B/F(^*)</td>
<td>90 min</td>
<td>3–6 d</td>
<td>-</td>
<td>-</td>
<td>2 y 7 m</td>
<td>Normal</td>
</tr>
<tr>
<td></td>
<td>4(^**)</td>
<td>P213S/P213S</td>
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<td>Caucasian</td>
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<td>None</td>
<td>No seizures</td>
<td>Not tried</td>
<td>Not tried</td>
<td>4 m</td>
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</tr>
<tr>
<td></td>
<td>5</td>
<td>R95C/R95C</td>
<td>+</td>
<td>Turkish</td>
<td>F</td>
<td>2 h</td>
<td>7 d</td>
<td>-</td>
<td>-</td>
<td>1 y</td>
<td>IQ 66 at 9 m. Mild truncal</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>hypotonia and speech delay</td>
</tr>
<tr>
<td></td>
<td>6(^d)</td>
<td>Q214fs/()</td>
<td>+</td>
<td>Caucasian</td>
<td>PN/F</td>
<td>5 h</td>
<td>15 d</td>
<td>-</td>
<td>-</td>
<td>4 y</td>
<td>Normal</td>
</tr>
<tr>
<td>(ii)</td>
<td>7</td>
<td>R116Q/R116Q</td>
<td>?</td>
<td>Caucasian</td>
<td>B</td>
<td>5 m</td>
<td>24 h</td>
<td>-</td>
<td>-</td>
<td>4 y</td>
<td>Normal/advanced</td>
</tr>
<tr>
<td>(iii)</td>
<td>8</td>
<td>R116Q/R116Q</td>
<td>-</td>
<td>Pakastani</td>
<td>?</td>
<td>3 h</td>
<td>2.5 m</td>
<td>+</td>
<td>7 y</td>
<td>Minimal delay</td>
<td></td>
</tr>
<tr>
<td></td>
<td>9</td>
<td>D33V/D33V</td>
<td>-</td>
<td>Caucasian</td>
<td>F</td>
<td>3 w</td>
<td>1 w</td>
<td>+</td>
<td>2 y</td>
<td>1 y</td>
<td>Minimal delay</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>D33V/E120K</td>
<td>?</td>
<td>Caucasian</td>
<td>?</td>
<td>2 m</td>
<td>4 m</td>
<td>+</td>
<td>21 y</td>
<td>Mild intellectual disability</td>
<td></td>
</tr>
<tr>
<td></td>
<td>11</td>
<td>D33V/Splice errors</td>
<td>+</td>
<td>Caucasian</td>
<td>F</td>
<td>3 h</td>
<td>6 m</td>
<td>+</td>
<td>21 y</td>
<td>Severe delay/no language</td>
<td></td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>D33V + R225C + R116Q(^b)</td>
<td>-</td>
<td>Caucasian</td>
<td>F</td>
<td>14 d</td>
<td>3.5 m</td>
<td>+</td>
<td>41</td>
<td>IQ 93. Dyslexia and Aspergers</td>
<td></td>
</tr>
<tr>
<td></td>
<td>13</td>
<td>R225H/R225H</td>
<td>?</td>
<td>?</td>
<td>?</td>
<td>24 h</td>
<td>6 m</td>
<td>+</td>
<td>7 y</td>
<td>Spastic quadriplegia with good social contact</td>
<td></td>
</tr>
<tr>
<td></td>
<td>14</td>
<td>R225H,R116Q + R225H,R116Q(^c)</td>
<td>-</td>
<td>Caucasian</td>
<td>?</td>
<td>30 min</td>
<td>2 w</td>
<td>+</td>
<td>2 y 7 m</td>
<td>Minimal delay</td>
<td></td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>R225H,R116Q + R225H,R116Q(^c)</td>
<td>+</td>
<td>Kosovan</td>
<td>?</td>
<td>10 h</td>
<td>5 d</td>
<td>+</td>
<td>8 y</td>
<td>DQ 65</td>
<td></td>
</tr>
</tbody>
</table>

\(\(^{a}\)\) Neonatal onset seizures responding to pyridoxal 5’-phosphate.
\(\(^{b}\)\) Infantile spasms (onset 5 months) responsive to pyridoxal 5’-phosphate.
\(\(^{c}\)\) Seizures starting under 3 months of age responding to pyridoxine.

\(B =\) breast-fed; \(F =\) formula fed; \(PN =\) parenteral nutrition; \(- = \) no; \(+ = \) yes

\(\(^{*}\)\) Formula from 2 weeks

\(\(^{\ast\ast}\) Sibling of Patient 3.

\(\(^{\ast\ast\ast}\) Formula from 2 weeks of Patient 3.

\(\(^{\ast\ast\ast\ast}\) Sibling of Patient 3.

\(\(^{c}\)\) [148G > A] and [364-1G > A] were inherited in \(c/s,\) this was confirmed by analysing parental DNA.

\(\(^{d}\)\) No parental DNA was available to ascertain which mutation R116Q was in \(c\) with.

\(\(^{e}\)\) Assume that R116Q has been inherited in \(c\) with R225H, no parental DNA was available to confirm this.

\(\(^{f}\)\) Second mutation not found.
Influence of environmental factors on phenotypes of individuals with identical or similar genotypes

D33V: a predominantly pyridoxine responsive genotype?

The response of Patients 8–15 to pyridoxine suggests that they have sufficient enzyme activity to allow synthesis of PLP from pyridoxine. Within this group, the missense mutation D33V was common. Expression studies showed that D33V did not totally abolish PNPO activity (44% residual activity). The response to treatment with pyridoxine is therefore perhaps not surprising. However, D33V was also found in Patient 2 whose seizures showed no response to pyridoxine. The situation is obviously complex. It is possible that pyridoxine responsiveness in PNPO deficiency is affected by prematurity and age and riboflavin status at the time of the therapeutic trial. Patient 9 (homozygous for D33V) responded to treatment with pyridoxine and was born at term, formula fed and tested for pyridoxine responsiveness at 3 weeks of age. Patient 10 (compound heterozygote for D33V and E120K) who showed a good response to pyridoxine (started at 6 months) in terms of both seizure control and psychomotor development and did not develop seizures until after the neonatal period (2 months). Patient 11 (compound heterozygote for D33V and c.264-21_264-1delinsC) resembled Patient 10 with regard to treatment response but differed in terms of neuropsychological outcome. Perhaps the poorer outcome in Patient 11 relates to the poorly controlled seizures in the first 2 months of life (Jonas et al., 2005). Seizures in Patient 12 (D33V in combination with R116Q and R225C) also responded well to treatment with pyridoxine. Conversely seizures in Patient 2 (homozygous D33V) failed to respond to a 10-day trial of 100 mg pyridoxine/day and previously reported patients [Patient 4; homozygous D33V (Goyal et al., 2013) and Patient 3; heterozygous D33V/heterozygous c.246deT (Hoffmann et al., 2007)] failed to respond to shorter trials of pyridoxine. Perhaps Patient 2 did not respond to pyridoxine because she was breast fed; breast feeding can be associated with a relative deficiency of riboflavin (Hovi et al., 1979) or perhaps because she has a second genetic factor compromising PLP homeostasis or neurotransmitter metabolism.

R225H/C genotype: a pyridoxine responsive phenotype?

Patients 14 and 15 had the same variant sequences (homozygous for R225H + R116Q) strongly suggesting that this is a pyridoxine-responsive genotype. Patient 13 (homozygous for R225H) also responded to treatment with pyridoxine, as did Patient 12 who was heterozygous for R225C. Expression of R225H and R225C revealed 8% residual activity—presumably sufficient to allow synthesis of PLP from pyridoxine. Further work is required to determine whether, in the R225H/C variants, pyridoxine phosphate oxidase activity is better preserved than pyridoxamine phosphate oxidase activity. These enzyme activities can be differentially affected by inhibitors (Takeuchi et al., 1985).

R95H, R95C and E50K with c.[364-1G > A]: alleles associated with infertility, miscarriage, premature birth and PLP-responsive seizures

Patient 1 was compound heterozygous for E50K plus c.[364-1G > A] on the paternal allele and R95H on the maternal allele. Homozygosity for E50K plus c.[364-1G > A] produced a similar clinical picture (parental infertility, premature birth and PLP-responsiveness) in two siblings (Mills et al., 2005). Patient 5 (homozygous for R95C) was premature and PLP-responsive and his parents had a history of miscarriage. An infant born at 36 weeks gestation with R95H/R95H in another series had severe neonatal epileptic encephalopathy with a brief, incomplete response to pyridoxine but still a fatal outcome (Khayat et al., 2008). In the Chinese hamster ovary cell expression system, PNPO enzyme activity for E50K plus c.[364-1G > A] and R95H was undetectable and 18% of wild-type, respectively. Why with one or both mutations showing 18% residual activity, should there be no or minimal response to pyridoxine? One possibility is that all of these patients were premature and tested for pyridoxine responsiveness in the first week of life. PNPO transcript is low in the foetus compared to adults (30% adult levels in liver; 2% adult levels in brain) (Ngo et al., 1998; Kang et al., 2004). An analysis of B6 vitamins in very preterm infants has shown that CSF may contain significant amounts of pyridoxine alongside a low PLP concentration (Albersen et al., 2012). Thus PNPO activity is already critically low in infants born before 30 weeks. PNPO mutations can therefore be expected to lead to more severe effects in a preterm than in a term infant. Further, this may suggest a therapeutic role for PLP in seizures in preterm infants without PNPO mutations. In addition newborns can develop riboflavin deficiency in the first days of life (Hovi et al., 1979) and this could further reduce PNPO activity.

Does the maternal supply of B6 to the developing foetus affect phenotype?

Patient 8, who was in poor condition at birth and started having seizures 3 h after birth, is homozygous for the sequence variant R116Q—the same genotype as Patient 7 who was normal until the age of 5 months. Therefore, whilst a predisposition to B6-responsive seizures may be determined by the R116Q/R116Q phenotype, other factors determine whether a child is in poor condition at birth and develops seizures in the first 24 h. The comparison of Patients 3 and 4 (siblings) indicates that the most likely additional factor is how well the foetus has been supplied with vitamin B6 by the mother (Schenker et al., 1992). In the case of Patient 4, the mother took a multivitamin preparation during pregnancy (2.6 mg/day of pyridoxine and 1.8 mg/day of riboflavin) and an additional PLP supplement just before delivery, which appeared to prevent intratuerine seizures and early post-natal seizures. A recent study suggests that PLP levels in Pakistani adults are lower than in other ethnic groups (Iqbal et al., 2009). If we look at the backgrounds of patients born prematurely, and/or those that had low Apgar scores and/or those that needed resuscitation, 17/21 are from a non-Caucasian background whereas for patients not meeting these criteria, the figure is only 1 in 8 (Table 4). Differences in maternal plasma PLP and pyridoxal levels may
explain the phenotypic differences seen in Patients 7 and 8, both of whom are homozygous for R116Q.

Prevalence of R116Q variant

The presence of R116Q-decreased PNPO activity by 20% when compared with wild-type; however, R116Q is listed as a single nucleotide polymorphism in Ensembl (Supplementary Table 3). It has been suggested that low PNPO activity in erythrocytes may confer resistance to malaria (Anderson et al., 1993). A second enzyme important in maintaining high levels of PLP in the red cell is pyridoxal kinase. Mean red blood cell pyridoxal kinase activity in black Americans is 40% that of white Americans (Chern and Beutler, 1976) and it has been suggested that this lower activity was favoured by natural selection because it conferred resistance to malaria (Martin et al., 1978; Flanagan and Beutler, 2006). Plasmodium produces enzymes for de novo synthesis of PLP and for phosphorylation of pyridoxal, but it is possible that erythrocyte schizogony requires some PLP to be provided by the host. A gene encoding PNPO, and therefore capable of regenerating PLP from PMP, has not been convincingly demonstrated in plasmodium (Müller and Kappes, 2007). Furthermore, at least one parasite enzyme that is important in erythrocyte schizogony (ornithine decarboxylase) gradually loses activity that can be restored by the addition of PLP (Bitonti et al., 1987).

Conclusion

Three groups of patients with PNPO mutations that reduced enzyme activity have been identified: (i) patients with neonatal onset seizures responding to PLP; (ii) a patient with infantile spasms (onset 5 months) responsive to PLP; and (iii) patients with seizures starting before 3 months of age responding to pyridoxine. One sequence variant, R116Q, a single nucleotide polymorphism that has been reported in the general population, was found to have an effect on PNPO activity. We believe the reduction in PNPO activity and B6-responsive epilepsy in the patients reported here indicates that it is a variant that contributes to the pathogenesis of epilepsy. It is possible that R116Q is responsible for the susceptibility locus for genetic generalized epilepsy on 17q21.32 (close to rs72823592). Although additional studies will be necessary to delineate this further, our findings support the use of DNA tests for PNPO deficiency in a wide range of infants with epilepsy.

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Supplementary material

Supplementary material is available at Brain online.

References

Epilepsy due to PNPO mutations

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