Male patients affected by mosaic PCDH19 mutations: five new cases

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Abstract
Pathogenic variants in the PCDH19 gene are associated with epilepsy, intellectual disability (ID) and behavioural disturbances. Only heterozygous females and mosaic males are affected, likely due to a disease mechanism named cellular interference. Until now, only four affected mosaic male patients have been described in literature. Here, we report five additional male patients, of which four are older than the oldest patient reported so far. All reported patients were selected for genetic testing because of developmental delay and/or epilepsy. Custom-targeted next generation sequencing gene panels for epilepsy genes were used. Clinical data were collected from medical records. All patients were mosaic in blood for likely pathogenic variants in the PCDH19 gene. In most, clinical features were very similar to the female phenotype, with normal development before seizure onset, which occurred between 5 and 10 months of age, clustering of seizures and sensitivity to fever. Four out of five patients had mild to severe ID and behavioural problems. We reaffirm the similarity between male and female PCDH19-related phenotypes, now also in a later phase of the disorder (ages 10–14 years).

Keywords PCDH19 · Mosaicism · Epilepsy · Intellectual disability

Introduction
Pathogenic variants in PCDH19 are associated with early onset, clustered epileptic seizures often provoked by fever, intellectual disability (ID) that can be present in variable degrees and behavioural disturbances such as autistic features,
attention deficit, hyperactivity and aggression. The clinical features may resemble those of Dravet syndrome (phenotype MIM # 300088) [1–5].

PCDH19 is located at Xq22.1 and codes for protocadherin-19, a transmembrane protein involved in neuronal organization and migration and cell-cell and cell-matrix adhesion [6–10]. It is highly expressed in the central nervous system [1]. PCDH19-related epilepsy shows a remarkable inheritance pattern: generally, females carrying heterozygous pathogenic variants are affected, whereas hemizygous male carriers are asymptomatic or only show psychiatric or behavioural symptoms [1, 2, 11, 12]. However, in 2009, the first affected male mosaic for a PCDH19 pathogenic variant was described [2]. This finding gave rise to a theory of cellular interference as disease mechanism: disease occurs when two different cell populations exist (cells expressing the normal PCDH19 protein and cells expressing a mutant form of the protein), as is true for heterozygous and mosaic pathogenic variants, but not for hemizygous pathogenic variants in males. These non-homogeneous cell populations are likely to disrupt cell-cell interactions, leading to disease [2].

Only four affected mosaic male patients have been described in literature until now, the oldest being 7 years old [2, 13, 14]. Here, we report five additional male patients with mosaic PCDH19 mutations, of ages 2, 10, 13, 13 and 14 years old. We compared the male and female phenotypes.

Methods

Patients and molecular analysis

All five patients were selected for diagnostic genetic testing because of developmental delay and/or epilepsy. Genetic testing on DNA from lymphocytes was performed using a custom targeted next generation sequencing (NGS) gene panel for epilepsy genes (see online resource 1 for details). Mosaicism of PCDH19 variants was determined based on the simultaneous presence of a variant allele and the reference allele, as PCDH19 is located on the X-chromosome and all patients were male. The percentage of mosaicism was based on the percentage of reads that showed the alternate allele. PCDH19 mosaic male patients from different centres were collected through personal communication between authors, meaning no structured cohort was tested. Detailed clinical data were collected from medical records. The parents of all patients gave informed consent for the publication of clinical data.

Literature search

A literature study was carried out in the PubMed database to identify previously described patients with PCDH19 pathogenic variants.

Results

Molecular analysis

All five patients carried a PCDH19 variant in various degrees of mosaicism (patient A: c.1864G > C, p.Gly622Arg, 60%; patient B: c.840C > G, p.Tyr280*, 22%; patient C: c.462C > G, p.Tyr154*, 65%; patient D: c.1682C > G, p.Pro561Arg, 78%; patient E: c.799G > T p.Glu267*, 20%, RefSeq NM_001184880). According to American College of Medical Genetics and Genomics (ACMG) criteria, the variant of patient A was classified as likely pathogenic, and the variants of patient B-D were classified as pathogenic. The variants of patient B, C and E lead to a premature stop codon; the variants of patient A and D were both predicted probably damaging and damaging by PolyPhen and SIFT, respectively. None of the variants was present in control databases (Database of Single Nucleotide Polymorphisms (dbSNP), NHLBI Exome Sequencing Project (ESP) and the the Exome Aggregation Consortium (ExAC) database). The Pro561Arg variant has been previously described in two affected female siblings with ID, microcephaly and seizures and was paternally transmitted [15]; the other variants are novel. All variants were confirmed de novo, except for the variant in patient D, whose parents were not tested. No other variants that could explain the phenotypes were found.

Clinical characteristics of the newly reported and previously described male patients with de novo PCDH19 pathogenic variants are summarized and compared to those of previously published females in Table 1. See online resource for extensive clinical descriptions (online resource 2). Overall, all patients had normal development before seizure onset, occurring between 5 and 10 months of age, except for patient E. This patient had a delay in speech development, like his father, before onset of seizures and seizure onset occurred later, at 31 months. First seizure types were generalized clonic or clonic–absence seizures. Later seizures were mainly complex partial seizures and primary or secondary generalized tonic–clonic seizures. In all five patients, seizures tended to cluster and could be provoked by fever. Four out of five patients had mild to severe ID, autism and additional behavioural problems.

Discussion

We here report five male patients with mosaic PCDH19 likely pathogenic variants, which raises the total number of described male patients to nine [2, 13, 14]. Four of the currently described male patients are the oldest reported so far (ages 10, 13 twice and 14 years old), which gives the opportunity to investigate whether PCDH19-related phenotypes evolve the same way in male and female patients. Our current findings
Table 1  Clinical description of male and female patients carrying *PCDH19* likely pathogenic variants

| Patient | A     | B     | C     | D     | E     | F [2] | G [13] | H [14] | I [14] | Female patients |
|---------|-------|-------|-------|-------|-------|-------|--------|--------|--------|----------------|
| Age at inclusion (years) | 10    | 13    | 14    | 24 months | 13    | 7     | 6      | 4      | 3.5    | 1–54          |
| Variant* | c.1864G > C, p.Gly622Arg | c.840C > G, p.Tyr280* | c.1682C > G, p.Pro561Arg | c.799G > T, p.Glu267* | del *PCDH19* | c.605C > A, p.Ser202* | c.918C > G, p.Tyr306* | c.1352C > T, p.Pro451Leu | PCDH19 deletions, duplications, miscense and nonsense variants |
| % of mosaicism | 60% (blood) | 22% (blood) | 65% (blood) | 78% (blood) | 20% (blood) | 100% (lymphocytes), 47% (fibroblasts) | 50% (blood), 70% (buccal cells), 100% (urine sediment) | Detection by exome sequencing, estimation of % mosaicism by Sanger | 10% (lymphocytes, saliva, hair) | 90% (lymphocytes, urine) |
| Technique used (number of alternate alleles/total read depth at base position) | NGS gene panel (92/153 reads) | NGS gene panel (77/380 reads) | NGS gene panel (38/59 reads) | NGS gene panel (351/450 reads) | NGS gene panel (19/93 reads) | Detection by microarray, estimation of % mosaicism by FISH (100 cells) | Detection by exome sequencing, estimation of % mosaicism by Sanger | NGS gene panel (157 reads) | NGS gene panel (135 reads) | Various |
| Sex | Male | Male | Male | Male | Male | Male | Male | Male | Male | Female |
| Exam at birth | Meconium stained amniotic fluid, bradycardia | Normal | Normal | Normal | Normal | Speech delay | Normal | Normal | Normal | Normal |
| Development prior to sz onset (months) | 5     | 10    | 10    | 7     | 31    | 12    | 9      | 9      | 10    | 3–70 (most <12) |
| Sz onset age (months) | 7     | 3     | 31    | ?     | ?     | ?     | ?      | ?      | ?     | ?              |
| Sz type at onset | Generalized, clonic (fs) | Generalized clonic | Cluster of CP seizures | Clusters of focal sz, generalizing to GTC and tonic seizures | Clusters of focal sz, generalizing to one hemisphere | GTC (fs, prolonged, repetitive) | Focal myoclonic, tonic-clonic, FSsG | Afebrile hypotonic seizure with hypopnea (40 min) | 24 h cluster of febrile sz with fixed gaze, loss of contact, upper limb hypertonia, and jerks | 24 h cluster of febrile sz with fixed gaze, loss of contact, upper limb hypertonia, and jerks (30–40 s.) |
| Later sz types | CP, secondary generalized. Focal with affective symptoms, secondary generalized. Primary generalized | Generalized clonic | CP, secondary tonic clonic, tonic, status | Focal, febrile, GTC, tonic, CP | CP, febrile | Hemiclonic, GTC, myoclonic jerks | Focal myoclonic, tonic-clonic, rapid secondary generalization | FS, focal tonic-vibratory | FS and afebrile seizure clusters, tonic. Often fearful screaming at start | FS or afebrile; generalized clonic, tonic-clonic or tonic; hemiconvulsion; focal, FSsG; CP |
| Clusters of sz | + (fever related) | + | + | + | + | + | ? | ? | + | + |
| Fever sensitivity | VPA: -, LTG: -, CBZ: + | VPA: + | VPA: + | VPA: + | VPA: + | VPA: + | VPA: + | VPA: + | VPA: + | VPA: + |
| AEDs used and response | VPA: ?, LEV: +/−, LTG: - | VPA: + | VPA: + | VPA: ?, LEV: +/−, LTG: - | VPA: + | VPA: + | VPA: + | VPA: + | VPA: + | VPA: + |
| Development | | | | | | | | | | |
Table 1 (continued)

| Patient | A | B | C | D | E | F [2] | G [13] | H [14] | I [14] | Female patients[d] |
|---------|---|---|---|---|---|------|--------|--------|--------|-------------------|
| Current AEDs[^b] | LEV, OXC, CLB, TPM | VPA | LEV | OXC, VPA, PHB, LEV | OXC | ? | ZNS | – | PHB, VPA | ZNS, NTZ, VGB, KBR, LZP: different responses |
| Sz outcome | 1 cluster of sz per year (5–10 sz/cluster) | Last sz at age 10 years | Ongoing sz | Ongoing sz | Ongoing sz during febrile illness | Persistence of febrile sz in spite of treatment | Seizure free for 20 months | Seizure free age 14–42 months (no AEDs), since then 1 cluster of sz | 4–5 clusters per year | Often seizures less frequent or seizure free at certain age (4–36 years) |
| EEG at onset | 7–8 months: normal. 10–11 months: asymmetrical background activity, non-specific high voltage delta-activity occipital right > left. | Frequent generalized epileptic discharges | 2 years: normal | Multiple focal discharges, right centroparietal, secondary generalization. Mild diffuse background slowing | Focal epileptic discharges left parieto-temporal with generalization to one hemisphere (ictal) | ? | Slower rhythm for age, mild diffuse disturbance. Infrequent right frontal, and rare left temporal sharp wave discharges, suggestive of epileptiform activity | Normal | Rare right frontotemporal sharp waves | Mostly normal |
| EEG at follow-up | 3,5 years: normal | – | 3 years: No epileptic activity, generalized and focal slow activity | – | – | ? | ? | ? | Bilateral centroparietal onset of seizures | Normal, interictal spikes, generalized poly-spikes waves, slow waves, slow background |
| Last EEG | 6 years: normal background activity, diffuse fast activity mainly frontal. No epileptic activity. | – | 4 years: No epileptic activity, generalized and focal slow activity | – | 32 months: interictal EEG normal | ? | ? | ? | Normal interictal – | – |
| ID[^c] | ++ (estimated: slowed PMD, special education) | + (IQ 66 at 5 years) | ++ to +++ | – (only delayed speech development, but no true ID yet, based on clinical evaluation) | + (IQ 55 at 9 years) | ++ to +++ | +/− (IQ = 76) | +/− (GDQ 78 at 46 months, 72 at 52 months) | – (GDQ 101 at 30 months and 103 at 40 months) | – to +++ |
| Developmental stagnation/−regression | – | – | – | – | – | ? | ? | ? | – | Regression in some cases |
| Language (words/sentences) | Delayed; words, sentences | Not delayed. First words at 12 months. Two-word sentences, stereotyped phrases | Mildly delayed speech development | Mildly delayed speech development | delay: words-sentences | ? | ? | ? | Sometimes normal, often delayed, words–sentences, in rare cases absent |
Table 1 (continued)

| Patient | A | B | C | D | E | F [2] | G [13] | H [14] | I [14] | Female patients^d |
|---------|---|---|---|---|---|-------|--------|--------|--------|-------------------|
| Behavioural/psychiatric disturbances | Autism, agitation, behavioural problems, ADHD | Autism, mood disorder | Autism spectrum disorder, anxiety | Behaviour problems resembling autism spectrum disorder, short attention span | Behavioural problems, autistic features | Irritability, aggression, rigidity, poor sleep, ADHD, anxiety, OCD, ODD | Compulsive and stereotyped behaviours | Prominent behavioural problems in most cases (autism, attention deficit, hyperactivity, aggression, emotional lability, impulsiveness, anxiety, jealousy, obsession, depression psychogenic nonepileptic sz) | Mostly normal; hypotonia, dyspraxia, ataxia or motor delay in some | Usually normal (mild atrophy/cortical dysplasia is rarely reported) |
| Neurological examination | Crouched gait | Hypotonia | Motor delay and balance problems | Balance problems (medication induced, improving) | Reduced coordination | Motor delay, ataxia | ? | ? | ? |
| MRI images (age) | Expanded perivascular spaces (8 years) | Normal (7 years) | Widen peripheral subarachnoid spaces | Normal (7 months) | Normal (2.5 years) | Normal | ? (CT: normal) | Normal (10 months) | Usually normal (mild atrophy/cortical dysplasia is rarely reported) |
| Additional comments | – | Pes plano valgus, obesity (BMI 22.3; +2.3 SD) | – | Hand foot mouth disease | – | dysmorphic features: plagiocephaly occipital-parietal area, cupped ears, intradigital webbing of phalanges. Severe myopia |

SZ seizure(s), AED anti-epileptic drug, PMD psychomotor development, GTC generalized tonic-clonic, CP partial complex, FS fever sensitive, SE status epilepticus, FSG focal seizure with secondary generalization, CBZ carbamazepine, CLB clobazam, CLN clonazepam, KBR potassium bromide, LEV levetiracetam, LSM lacosamide, LTG lamotrigine, LZR Lorazepam, NTZ nitrazepam, OXC oxcarbazepine, PHB phenobarbital, PHT phenytoin, STP stiripentol, TPM topiramate, VGB vigabatrin, VPA valproic acid, ZNS zonisamide, GDQ Griffiths Developmental Quotient

^a RefSeq NM_001184880

^b “+” not effective, “+/−” slight effect, “+” good effect

^c “−” absent, “+/−” borderline, “+” mild, “++” moderate, “+++” severe

^d [2–5, 11, 12, 15–29]
confirm previously reported observations of similar clinical features in male and female patients, also for older children [2, 13, 14]. Focal seizures with affective symptoms (fearful screaming) are very common in female patients and become more prevalent with an increasing age [26]. This distinctive seizure type is also reported in three male patients [14, current study]. These similarities suggest that there are no differences in clinical consequences between phenotypes caused by postzygotic PCDH19 pathogenic variants in mosaic males, and phenotypes caused by heterozygous PCDH19 pathogenic variants in females. This lends further support to the hypothesis that cellular interference is the main disease mechanism, as proposed by Depienne et al. [2] The increasing number of affected mosaic male patients undermines the theory of a compensating effect by the nonparalogous PCDH11Y gene in male patients, as proposed by Dibbens et al. [1]. It is highly unlikely that this gene would only compensate for a complete absence of PCDH19 in hemizygous affected males, but not for a partial loss of PCDH19 in mosaic males.

In female patients, seizure frequency often diminishes around puberty [3, 5, 11, 28–30], possibly due to hormonal changes [30]. Our 13-year old patient has been seizure free since the age of 10 years old; our 10-year-old patient only has one cluster of seizures per year. However, our other 13- and 14-year-old patients still have ongoing seizures. Although four of our male patients are the oldest reported until now, none of them has reached adolescence yet, and the numbers are still small, making it hard to draw definite conclusions. Nevertheless, since some of the few reported male patients already show a reduction in seizure frequency with increasing age, it seems unlikely that a declining seizure frequency is exclusively occurring in females and is related to female specific hormones.

We hypothesised that males with a mutated allele percentage around 50% in the brain, who would have an inherent high level of cellular interference, may show a more severe clinical picture than males with a lower or higher percentage of mosaicism. High or low percentages of mosaicism would resemble skewed X-inactivation in female patients, which has also been suggested to lead to a milder phenotype [5, 15], although no clear correlation has been shown [25, 29]. Indeed, in our cohort, the three patients with the lowest and highest percentages of mosaicism in blood are the least severely affected (patients B, D and E), and two previously described patients, one with borderline ID and one with normal intelligence showed percentages of mosaicism of 10 and 90%, respectively (patients H and I). However, patient G shows an equal number of mutated and wild-type alleles in blood and is also mildly affected, and patient F shows no mosaicism in blood cells at all, only in fibroblasts, but has moderate to severe ID and ongoing seizures. It is thus not possible to predict the phenotype based on the percentage of mosaicism in blood, most likely because it does not necessarily equal the percentage of mosaicism in the brain.

The number of identified male patients mosaic for PCDH19 pathogenic variants increases [2, 13, 14], probably due to our improving abilities to detect mosaicism in general by using NGS techniques. It is now clear that PCDH19 pathogenic variants can cause epilepsy both in males and females and that mosaicism for PCDH19 pathogenic variants in males might be more common than previously thought. Because our five described patients were gathered through personal communication between authors from different diagnostic centres, no structured cohort with clearly defined inclusion criteria was tested, which makes it difficult to estimate the frequency of mosaic pathogenic PCDH19 variants in male patients. Since the PCDH19 and Dravet syndrome phenotypes show many similarities, testing a cohort of SCN1A-negative, male Dravet syndrome patients for mosaic PCDH19 pathogenic variants could give more insight in the true incidence. In this cohort and in males with clinical features characteristic of a PCDH19-related disorder, NGS techniques with high coverage should be used to look for PCDH19 pathogenic variants, as traditional Sanger Sequencing is not sensitive enough to reliably detect mosaicism. This overview helps create more knowledge about the disease course in male patients, which is extremely relevant for counselling those affected and their families. Reporting on more (older) patients in the future is essential for establishing a good understanding of prognosis in male patients.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflicts of interest. HAN and KLH are employed by Ambry Genetics; PCDH19 sequencing is among its commercially available tests. This study was supported by the “Stichting Vrienden WKZ” (project 1614054) on behalf of Stichting Panta Rhei. Funders had no involvement in the study design, the collection, analysis, and interpretation of data, the writing of the report and in the decision to submit the paper for publication.

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