11p13 deletions can be more frequent than the PAX6 gene point mutations in Polish patients with aniridia

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Received: 7 December 2012 / Revised: 29 April 2013 / Accepted: 20 May 2013 / Published online: 13 June 2013
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Abstract Aniridia is a rare, bilateral, congenital ocular disorder causing incomplete formation of the iris. The visual impairment caused by iris aplasia/hypoplasia might be enhanced by several ocular anomalies, such as cataracts, glaucoma, conical pannus, optic nerve hypoplasia, macular hypoplasia, or ectopia lentis. In the majority of cases, it is caused by mutation in the PAX6 gene, but it can also be caused by microdeletions that involve the 11p13 region. Twelve unrelated patients of Polish origin with a clinical diagnosis of aniridia were screened for the presence of microdeletions in the 11p13 region by means of multiplex ligation probe amplification (MLPA). Additionally, the coding regions of the PAX6 gene were sequenced in all probands. MLPA examination revealed different size deletions of the 11p13 region in five patients. In three cases, deletions encompassed the entire PAX6 gene and a few adjacent genes. In one case, a fragment of the PAX6 gene was deleted only. In the final case, the deletion did not include any PAX6 sequence. Our molecular findings provide further evidence of the existence of the distant 3′ regulatory elements in the downstream region of the PAX6 gene, which is known from other studies to influence the level of protein expression. Sequence analysis of the PAX6 gene revealed the three different point mutations in the remaining four patients with aniridia. All the detected mutations were reported earlier. Based on accomplished results, the great diversity of the molecular basis of aniridia was found. It varies from point mutations to different size deletions in the 11p13 region which encompass partly or completely the PAX6 gene or cause a position effect.

Keywords Aniridia · PAX6 gene · 11p13 microdeletions · MLPA analysis

Introduction

Aniridia (MIM:106210) is a rare, bilateral, congenital ocular disorder causing incomplete formation of the iris. The visual impairment caused by iris aplasia/hypoplasia might be enhanced by several ocular anomalies, such as cataracts, glaucoma, conical pannus, macular hypoplasia, nystagmus, and photophobia. The prevalence of aniridia in the general population is approximately 1 in 64,000 to 1 in 96,000 live births. Two-thirds of cases are familial, with autosomal dominant mode of inheritance, complete penetrance, and variable expressivity. One-third of cases are sporadic and result from de novo mutations (Kokotas and Petersen 2010).

Mutations in the PAX6 gene

PAX6 (MIM:607108; NM_001604.4) was identified as a candidate gene for aniridia by positional cloning in 1991 (Ton et al. 1991). Mutations in the PAX6 gene are the major cause of congenital aniridia. PAX6 is a member of the paired box gene family, located on chromosome 11p13, and consists of 14 exons that span over 22 kb in length. It encodes a 422-amino acid transcriptional factor, involved in oculeogenesis and other developmental processes. The protein contains three functional domains: paired domain (PD) and homeodomain (HD), separated by the proline/serine/threonine-rich transactivation domain (PST). The paired domain comprises two structurally distinct DNA-binding subdomains: the N-terminal subdomain (NTS) and the C-terminal subdomain (CTS) (Kokotas and Petersen 2010). The protein is abundantly expressed in the forebrain, early eye structures, ventral spinal cord, and endocrine pancreas (Lauderdale et al. 2000). Pathogenic alterations in the PAX6 gene reported to date include: nonsense, splicing, frame-shifting insertions or...
deletions, in-frame insertions or deletions, missense, and run-on mutations (D’Elia et al. 2007). Human PAX6 mutations are archived in the PAX6 Allelic Variant Database (LOVD: Leiden Open Variation Database, http://lsdb.hgu.mrc.ac.uk/home.php). The majority of mutations in the PAX6 gene are nonsense mutations, which are considered to produce truncated transcripts that activate the nonsense-mediated decay (NMD) process, resulting in haploinsufficiency of PAX6 (Fantes et al. 1995).

11p13 deletions

Aniridia can be also caused by chromosomal rearrangements that involve the 11p13 region. A significant proportion of chromosomal rearrangements are submicroscopic microdeletions (Crolla and van Heyningen 2002). There have been described cases of aniridia with deletions not involving the structure of the gene but with the removal of 3’ regulatory elements essential for adequate PAX6 expression and known as a “position effect” (Fantes et al. 1995).

Chromosomal alterations in the chromosome 11p, encompassing the PAX6 and WT1 (*607102) genes, may cause WAGR syndrome (MIM:194072). WAGR syndrome is a rare contiguous gene deletion syndrome, clinically associated with Wilms’ tumor, aniridia, genitourinary anomalies, and mental retardation (Fischbach et al. 2005). Wilms’ tumor is one of the most common solid tumors of childhood, which affects approximately 1 in 10,000 children. In patients with WAGR syndrome, the risk of Wilms’ tumor will develop reaches 45 % (Crolla and van Heyningen 2002).

The aim of this study was to analyze the genetic background of disease in Polish patients with aniridia.

Materials and methods

Patients

This study comprises 12 unrelated patients of Polish origin, with a clinical diagnosis of aniridia, based on ophthalmologic examination and family history. Among them, there are five patients with positive family history, six sporadic cases, and one patient (p6) who was adopted, for whom we have no information about the patient’s biological parents. The ophthalmologic findings of the patients are summarized and compared in Table 1. The local ethics committee approved the study and a written informed consent was obtained from all subjects or their legal guardians.

Methods

Genomic DNA was extracted from venous blood samples using a standard salting-out procedure. Patients were screened for structural rearrangements in the 11p13 region, using multiplex ligation probe amplification (MLPA) analysis. MLPA allowed in the first instance to exclude WT1 gene deletion, associated with a high risk of Wilms’ tumor. MLPA reaction was performed using the P219-B1 commercial kit (MRC-Holland, Amsterdam, Netherlands), following the manufacturer’s instructions and using an exactly defined amount of genomic DNA (200 ng). The kit contains probes for the PAX6 gene (with the exception of exon 6) and for the following genes located upstream, WT1, HIPK3, LMO2, EHF, CD44, and RCN1, and downstream, BDNF, FSHB, DCC1, and ELP4, of PAX6. Data were intra-normalized by dividing the area of the peak for each amplification product by the total area of the reference probes’ products in the probemix. Inter-sample normalization was achieved by dividing the intra-normalized probe ratio of all reference samples. Data normalization was performed within one experiment (MRC-Holland). Peak areas in the range 0.65–1.31 were considered normal, below 0.65 as a deletion, and above 1.31 as a duplication. Each MLPA analysis included several control samples (DNA from healthy individuals).

The coding regions of the PAX6 gene were sequenced (exon 5a included) to exclude point mutations. PCR products were separated on an ABI 3130xl Capillary Sequencer (Applied Biosystems). Additionally, in patients p2, p3, and p6, the PAX6 gene promoter was sequenced. The sequenced fragment included 5’UTR and 1,500 nucleotides upstream of the 5’UTR fragment of the PAX6 gene. The sequences of the primers used in the amplification and sequencing of the PAX6 gene and PAX6 promoter are given in Table 2. Subsequently, to uncover probable, smaller, rare copy number variations (CNVs), array comparative genomic hybridization analysis (using NimbleGen Human CGH 6x630K, Roche) of these three patients was performed. The DNA labeling reaction and hybridization were carried out according to the manufacturer’s instructions.

Results

MLPA examination revealed different size heterozygous deletions of the 11p13 region in five patients. In three cases, deletions encompassed the PAX6 gene and a few adjacent genes. The heterozygous deletion in patient p8 contains five annotated genes: PAX6, doublecortin domain-containing protein 1 (DCDC1, MIM:608062), elongation factor protein 4 (ELP4, MIM:606985), brain-derived neurotrophic factor (BDNF, MIM:113505), and beta polypeptide of follicle-stimulating hormone (FSHB, MIM:136530) (Fig. 1a). The heterozygous deletion in patient p9 harbors the following six genes: PAX6, DCDC1, ELP4, FSHB, reticulocalbin 1 (RCN1, MIM:602735), and DKFZp686K1684. In patient p11, a large heterozygous deletion of the 11p13 region
(WT1 gene included) was found (Fig. 1b), which confirmed the diagnosis of WAGR syndrome. In one patient (p12), a heterozygous deletion of several contiguous exons (exons 1–8) of the PAX6 gene was found (Fig.1c). In patient p10, MLPA revealed a heterozygous deletion of approximately 0.6 Mb on chromosome 11. The deletion did not include the PAX6 sequence. The following four genes located downstream of PAX6 are present in the deleted region: DCDC1, zinc finger CLS domain containing protein 3 (DPH4, MIM:611072), IMP1 inner mitochondrial membrane peptidase (IMMP1L, MIM:612323), and ELP4 (published data, Wawrocka et al. 2012).

The analysis of the coding region of the PAX6 gene by direct sequencing revealed three different point mutations in

| Table 1  | Clinical features of the patients |
|----------|----------------------------------|
| Patient  | Aniridia | Congenital cataract | Congenital glaucoma | Nystagmus | Photophobia | Other features |
| p1       | +        | +                  | +                  | +        | +          | –             |
| p2       | +        | +                  | –                  | –        | –          | Complex microphthalmia, lens subluxation |
| p3       | +        | –                  | –                  | –        | –          | Atrophy of the eyeballs, blindness |
| p4       | +        | +                  | –                  | +        | +          | Atrophy of the optic nerves, persistent hyaloid artery |
| p5       | + (partial) | –           | –                  | +        | +          | Atrophy of the optic nerves, psychomotor developmental delay |
| p6       | +        | –                  | –                  | +        | +          | Facial dysmorph, left kidney agenesis |
| p7       | +        | +                  | –                  | –        | –          | –             |
| p8       | +        | –                  | –                  | +        | –          | Astigmatism |
| p9       | +        | +                  | –                  | –        | –          | –             |
| p10      | + (partial) | +          | –                  | –        | –          | Lens subluxation, macular hypoplasia |
| p11      | +        | –                  | +                  | +        | –          | Psychomotor developmental delay, muscular hypotony, cryptorchidism (WAGR syndrome) |
| p12      | +        | +                  | +                  | +        | +          | Atrophy of the eyeballs, uveal coloboma |

| Table 2  | Primers used for the amplification and sequencing of the PAX6 gene and PAX6 gene promoter |
|-----------|-----------------------------------------------|
| PAX6 gene | Sense primer | Antisense primer | Product size (bp) |
| Exon 4   | AACAGAGCCCCATATTCGAG | GAGTCCCTGTGCTCTCCCA | 143 |
| Exon 5   | CTCTTCTTCCCTACCTTG | GAAATGAGAGAGGCCTTG | 268 |
| Exon 5a  | CTCTACAGTAAGTCTCTACCC | GGAAGTGAAGAGAGACAGAAACCC | 197 |
| Exon 6–7 | TTTGATGCATCTTCCAGCCAG | GACTCCACACACTTGACTCC | 748 |
| Exon 8   | TTGGAATGACTTGCTCTCCAG | ATGTTGAGAGAGTAGGAG | 300 |
| Exon 9   | TTAAGACTCAACCAGGCC | TGAAGATGGAGCAATTCTT | 294 |
| Exon 10–11 | TTGATGCACACAGTTGTGCAAC | GAGAGACAGGCGCC | 602 |
| Exon 12  | TAAATCCTGTGGCTCTCC | AAGCTCTCAAGGGTGCA | 282 |
| Exon 13  | CTTGTTGGCGAGAGTCTCCTCG | AGAGTTAGGCAACACGCCCTCC | 347 |
| Exon 14  | CCATGTCTGTTGCTCCAAGG | AAGCTCAACTGTTGCTCC | 222 |

PAX6 gene promoter:

- Product 1: AAGAGGTCTAGCTACAGCTCCAAGTCCT | CTGACACCACACACAGAAGTCT | 288
- Product 2: TGGAGTCTGTGCTGTGCT | GGTATGTCAAGCGTGTCCTG | 185
- Product 3: CAGCGTCACTCAGGAGCT | AAGCCAGGGGAGATAAGG | 156
- Product 4: AGAGAATCCCTCAGCAGATT | TCCTACCAACACACTCACA | 163
- Product 5: CAGCTATGATAGTCAACA | CGCCTCTTCCTTCTCCTT | 195
- Product 6: GCTTCCGAAAAGGAGAAAAGGA | CCCCTGGAAGAAAGAAGAAA | 216
- Product 7: CCCCTTCTCCTCTCCTCAA | CAAATGTCCTGTTTGTGGA | 386
- Product 8: ACCAGAAACCTCTACTTGAC | AGCAAACACTTCCTCCTGC | 416
Fig. 1 The normalized multiplex ligation probe amplification (MLPA) results of the three probands. A patient p8, B patient p11, C patient p12. The light gray columns represent the deleted probes.
the remaining four patients. All the detected mutations were previously reported. In patient p1, we identified a c.718C>T heterozygous nonsense mutation, which occurred in the protein homeodomain (HD), resulting in a change of arginine to stop codon (p.Arg240*). In patients p4 and p7, another heterozygous nonsense mutation at position c.949C>T was identified. It is present in the PST domain coding region, and results in a premature termination of the PAX6 protein (p.Arg317*). According to the database, the identified mutations are common deleterious mutations in the PAX6 gene and they probably encode a non-functional protein. Bidirectional sequencing of exon/intron boundaries for patient p5 resulted in the identification of a nucleotide substitution at the splice donor sequence of intron 6, c.357+1G > A (IVS6+1G > A) (Fig. 2).

Sequencing of the PAX6 gene promoter in patients p2, p3, and p6 did not identify any mutation in this region. So, the molecular methods used in this study did not allow to detect any mutations in patients p2, p3, and p6. Therefore, array comparative genomic hybridization analysis of these three patients was performed. However, array comparative genomic hybridization (aCGH) analysis also did not reveal any causative mutation. The patients’ molecular findings are summarized in Table 3.

Discussion

This is the first molecular study carried out on Polish patients with aniridia. In this study, we identified the alterations that cause the disease in 9 of 12 examined (75 %) patients with aniridia. This is also worth mentioning that, in our group of patients, aniridia is more frequently caused by large deletions than by the point mutations. This statement is in contrast to larger studies done by Ton et al. (1991), Kokotas and Petersen (2010), and Park et al. (2012). Those researchers stated that heterozygous mutations in the PAX6 gene are found in about 40–80 % of all non-syndromic aniridia patients. However, we must remember that our results are based on a small group, thus, our conclusion requires further verification with a larger cohort of aniridia patients.

The large deletion containing not only PAX6 but also several other genes and the RCNI gene was detected by MLPA analysis in patient p9 (Table 3). The RCNI gene is a Ca2+ binding protein, participating in the secretory pathway. Favor et al. (2009) reported that mouse, containing a heterozygous deletion of the Alox and RCNI genes, presents an extreme microphthalmia. They suggest that RCNI may have an influence on the eye phenotype in PAX6 contiguous gene deletions. A familial case of aniridia resulting from a deletion containing PAX6, RCNI, and DKFZp686k1684 was reported by Zhang et al. (2011). Severe visual impairment observed in the described family is compatible with the mouse phenotype reported by Favor et al. (2009). Patient p9 contains a very similar deletion of the downstream flanking region of PAX6 to the family described by Zhang et al. (2011). However, our patient exhibits no significant phenotype differences in comparison to other aniridia patients presented in this study. Microphthalmia was not observed.

When the deletion does not include the PAX6 sequence, then the so-called “position effect” is the underlying pathogenic mechanism of aniridia in these patients. Such a case is observed in patient p10, who has been diagnosed to have aniridia, despite the fact that the structure of the PAX6 gene was normal. Submicroscopic deletions not involving the PAX6 gene in the patients with aniridia were previously described by several authors (Lauderdale et al. 2000; DeLia et al. 2007; Davis et al. 2008; Wawrocka et al. 2012). MLPA analysis revealed in this patient a heterozygous deletion encompassing several genes, including the ELP4 gene. Kleinjan et al. (2006) confirmed that cis-regulatory elements located within the introns of the ELP4 gene are PAX6-specific long-range control elements, essential for normal expression of the gene. Functional studies in human cells and animal models confirmed the presence of 3’ regulatory elements in the downstream region of PAX6, which are essential for normal expression of the gene. Our molecular findings provide further evidence of the existence of the distant 3’ regulatory elements in the downstream region of the PAX6 gene, which are known from other studies to influence the level of protein expression.

DNA sequencing of the PAX6 gene revealed three different mutations: two nonsense mutations, c.718C>T and c.949C>T (in two patients), and one splice-site mutation, c.357+1G > A. It was previously described that the mutation c.357+1G > A lead to the use of a cryptic splice donor site within exon 6, leading to an in-frame deletion of 36 amino acids from the PD (Hingorani et al. 2009). The heterozygous mutation c.718C>T (p.Arg240*) in patient p1 occurred in the protein HD. In two unrelated patients (p4 and p7), identical heterozygous substitution c.949C>T (p.Arg317*) in the PST domain coding region was discovered. In accordance to the PAX6 mutation.

![Fig. 2 Forward sequence analysis of exon 6 and adjacent intronic sequence of the PAX6 gene in patient p5. The arrow indicates the heterozygous G/A transition in intron 6 at position +1 c.357+1G > A](image)

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**Fig. 2** Forward sequence analysis of exon 6 and adjacent intronic sequence of the PAX6 gene in patient p5. The arrow indicates the heterozygous G/A transition in intron 6 at position +1 c.357+1G > A.
database: c.718C>T and c.949C>T are the most frequent 
PAX6 mutations causing aniridia. Despite the presence of the 
same mutation, both patients represent quite divergent pheno-
types. Singh et al. (1998) showed that the variable phenotype 
could be explained by differences in PAX6 protein levels, the 
ratio of mutant and normal protein, and interaction with dif-
ferent molecular factors. It was reported that mutations behind 
the HD could lead to more severe phenotypes in comparison 
with truncating mutations in the PD or HD domains (Kokotas 
and Petersen 2010; Park et al. 2012). This partly agrees with 
our observations that patient p4 (mutation in the PST domain) 
showed a more severe phenotype of aniridia in comparison to 
patient p1 (mutation in the HD) (Table 1).

We have not determined any genetic causes of aniridia in 
three patients (p2, p3, and p6). The disease can be caused by 
abnormalities in the PAX6 gene promoter or in the regula-
tory elements that affect the normal expression of the gene 
but are not routinely subjected to sequencing. Therefore, we 
declined to sequence the PAX6 gene promoter in these 
patients. However, the sequencing did not reveal any muta-
tion in the PAX6 promoter. Another possible explanation for 
this fact is that chromosomal rearrangements that involved a 
gene or genes in the 11p13 region do not include the probes 
used in the MLPA analyses. Patient p2 suffers from aniridia and 
microphthalmia. The presence of microphthalmia with 
aniridia suggests the possibility of genocopy. Therefore, 
another gene involved in oculogenesis can be responsible 
for the disease. Possibly, the damage of the iris only resembles 
aniridia. aCGH analysis of patient p2 was performed, as 
well as in two other patients, in whom the genetic background 
of aniridia was not determined (p3, p6). This analysis allows 
to uncover smaller, rare CNVs that cannot be detected by 
lower resolution molecular techniques. However, no causative 
mutation was found.

Ophthalmic findings revealed nearly no phenotypic vari-
abilities of the aniridia. Patients having deletion 3′ to the 
PAX6 gene exhibit no significant phenotypic differences 
compared to the patients with point mutations in the PAX6 
gene. Due to quite the homogenous clinical symptoms and a 
small number of patients, we were not able to define any 
genotype–phenotype correlations. The findings of the current 
study are consistent with those of D’Elia et al. (2007), Lim et al. 
(2012), and Park et al. (2012), who did not find any phenotypic 
differences according to the identified genotype. Defining the 
genotype–phenotype correlation requires the analysis of a larg-
er cohort of patients.

In connection to the great variability of mutations in anir-
idia patients, it is recommended to perform several analyses to 
detect the genetic background of aniridia. First of all, we care 
about the exclusion of life-threatening Wilms’ tumor (associ-
ated with the WT1 gene deletion) and then the detection of 
genetic causes of aniridia. Therefore, we decided to perform 
MLPA analysis first (to detect structural rearrangements in the 
PAX6 gene and 11p13 region) and then PAX6 gene sequencing 
to identify point mutations). Additionally, gene sequencing is 
not usually used to detect copy number changes; therefore, in 
our opinion, it is better to begin the molecular analysis with 
MLPA or aCGH.
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