Characterization of two novel intronic OPA1 mutations resulting in aberrant pre-mRNA splicing

Bolognini, Ramona; Gerth-Kahlert, Christina; Abegg, Mathias; Bartholdi, Deborah; Mathis, Nicolas; Sturm, Veit; Gallati, Sabina; Schaller, André

Abstract: BACKGROUND We report two novel splice region mutations in OPA1 in two unrelated families presenting with autosomal-dominant optic atrophy type 1 (ADOA1) (ADOA or Kjer type optic atrophy). Mutations in OPA1 encoding a mitochondrial inner membrane protein are a major cause of ADOA. METHODS We analyzed two unrelated families including four affected individuals clinically suspicious of ADOA. Standard ocular examinations were performed in affected individuals of both families. All coding exons, as well as exon-intron boundaries of the OPA1 gene were sequenced. In addition, multiplex ligation-dependent probe amplification (MLPA) was performed to uncover copy number variations in OPA1. mRNA processing was monitored using RT-PCR and subsequent cDNA analysis. RESULTS We report two novel splice region mutations in OPA1 in two unrelated individuals and their affected relatives, which were previously not described in the literature. In one family the heterozygous insertion and deletion c.[611-37
6
11−38insACTGGAGAATGTAAAGGGCTTTT;611−6
6
11−16delCATATTATCT] was found in all investigated family members leading to the activation of an intronic cryptic splice site. In the second family sequencing of OPA1 disclosed a novel heterozygous deletion c.2012+47delAGT resulting in exon 18 and 19 skipping, which was not detected in healthy family members. CONCLUSION We show the importance of transcript analysis to determine the consequences of unclear intronic mutations in OPA1 in proximity to the intron-exon boundaries.

DOI: https://doi.org/10.1186/s12881-017-0383-x

Originally published at: Bolognini, Ramona; Gerth-Kahlert, Christina; Abegg, Mathias; Bartholdi, Deborah; Mathis, Nicolas; Sturm, Veit; Gallati, Sabina; Schaller, André (2017). Characterization of two novel intronic OPA1 mutations resulting in aberrant pre-mRNA splicing. BMC Medical Genetics, 18(1):22. DOI: https://doi.org/10.1186/s12881-017-0383-x
Characterization of two novel intronic OPA1 mutations resulting in aberrant pre-mRNA splicing

Ramona Bolognini 1,5, Christina Gerth-Kahlert 2, Mathias Abegg 4, Deborah Bartholdi 1, Nicolas Mathis 1, Veit Sturm 3, Sabina Gallati 1 and André Schaller 1*

Abstract

Background: We report two novel splice region mutations in OPA1 in two unrelated families presenting with autosomal-dominant optic atrophy type 1 (ADOA1) (ADOA or Kjer type optic atrophy). Mutations in OPA1 encoding a mitochondrial inner membrane protein are a major cause of ADOA.

Methods: We analyzed two unrelated families including four affected individuals clinically suspicious of ADOA. Standard ocular examinations were performed in affected individuals of both families. All coding exons, as well as exon-intron boundaries of the OPA1 gene were sequenced. In addition, multiplex ligation-dependent probe amplification (MLPA) was performed to uncover copy number variations in OPA1. mRNA processing was monitored using RT-PCR and subsequent cDNA analysis.

Results: We report two novel splice region mutations in OPA1 in two unrelated individuals and their affected relatives, which were previously not described in the literature. In one family the heterozygous insertion and deletion c.[611-37_611-38insACTGGAGAATGTAAAGGGCTTT;611-6_611-16delCATATTATCT] was found in all investigated family members leading to the activation of an intronic cryptic splice site. In the second family sequencing of OPA1 disclosed a de novo heterozygous deletion c.2012+4_2012+7delAGTA resulting in exon 18 and 19 skipping, which was not detected in healthy family members.

Conclusion: We identified two novel intronic mutations in OPA1 affecting the correct OPA1 pre-mRNA splicing, which was confirmed by OPA1 cDNA analysis. This study shows the importance of transcript analysis to determine the consequences of unclear intronic mutations in OPA1 in proximity to the intron-exon boundaries.

Keywords: OPA1, Splice site mutation, Autosomal dominant optic atrophy, Kjer type optic atrophy, Optic neuropathies

Background

Autosomal dominant optic atrophy type 1 (ADOA1, MIM#165500), also known as Kjer-type optic atrophy [1, 2], is the most frequent form of inherited optic neuropathies (ION), with a prevalence ranging from 1:10,000 [3, 4] in Denmark to 1:50,000–30,000 [3–6] worldwide. ADOA is typically characterized by childhood-onset insidious moderate to severe progressive bilateral visual loss, color vision deficits, centrocecal scotoma, temporal optic nerve pallor and macular ganglion cell loss [7], which often leads to legal blindness [8–10]. Although ADOA is genetically heterogeneous [11–13], the vast majority of ADOA patients (approx. 75%) harbor a mutation in the OPA1 gene, less frequent mutations in OPA3 and WFS1 are observed in ADOA patients. OPA1 codes for a mitochondrial dynamin-like GTPase [14–16], which is an inner membrane protein targeted to mitochondria by a highly basic N-terminal import sequence and is anchored to mitochondrial cristae where it faces the intermembrane space [17]. OPA1 mRNA is 6492 bp (largest isoform) in length and has been mapped to chromosome 3q28-q29 [15, 18, 19]. OPA1 consists of 31 exons of which the last one is non-coding [15, 19]. Through alternative splicing of
OPA1 is involved in many different processes. It is the key player during mitochondrial inner membrane fusion, it plays a role in mitochondrial DNA, respiratory chain and membrane potential maintenance, cristae organization by controlling their shape and structure and it is further involved in cytochrome c mediated apoptosis [22–25]. More than 340 unique DNA variants have been allocated to OPA1 (http://mitodyn.org/home.php?select_db=OPA1) [26]. Half of these mutations result in haploinsufficiency of OPA1 by premature termination of translation leading to a truncated protein. Most of these mutations are located in the GTPase domain, whereas no mutation was found in the alternatively spliced exons (4, 4b and 5b). The phenotypic expression of OPA1 mutations in ADOA shows a high variability between families as well as members of the same family [27], often with an incomplete penetrance ranging from asymptomatic carriers and mild visual impairment to legally blind individuals [28]. ADOA can be classified either as a non-syndromic or isolated form, presenting with pure ophthalmologic features or as syndromic form associated with extra-ocular neurological manifestations, known as ADOA plus syndrome [5, 29]. Around 20% of ADOA patients develop additional neuro-muscular features, often in the third decade of life, which are sometimes present at a subclinical level. These include sensorineural deafness, ataxia, axonal sensory-motor polyneuropathy, mitochondrial myopathy and external ophthalmoplegia, whereas in one case parkinsonism and dementia were reported [30]. Many different mutations have been identified in OPA1 responsible for ADOA, including nonsense mutations, missense mutations, frameshift mutations, in-frame deletions and splice site or splice region mutations [14, 28, 31]. Since the vast majority of OPA1 mutations are predicted to result in a premature stop codon, mainly caused by splice site or splice region, nonsense or frameshift mutations, the generation of a truncated protein leading to haploinsufficiency is proposed to be the major cause for the pathogenesis of ADOA [28]. The majority of transcripts containing a premature termination codon (PTC) are degraded by a mechanism called nonsense-mediated decay (NMD), which is used to protect the cell from translation of potentially harmful proteins [32]. In this study we analyzed all coding exons including the intron-exon boundaries of OPA1 by DNA sequencing of two unrelated families presenting with clinical features of ADOA. We report two novel splice region mutations in OPA1, which we characterized at RNA level, thereby expanding the mutational spectrum of OPA1.

Methods

Patients

Family 1

The index patient (III:1, Fig. 1a) was referred to our institutions at age of 9 years because of reduced vision and optic disc pallor. The child is the first child of unrelated parents of Swiss origin. She is normally developed and besides her vision disorder a healthy child. On the first examination, best corrected visual acuity (BCVA) for distance was reduced to 0.3 and 0.5, which dropped to 0.2 and 0.32 at 10 years in the right and left eye, respectively. Temporal optic atrophy was present in both eyes. The apparently unaffected mother (II:2) showed minor optic atrophy, which was confirmed by the reduced retinal nerve fiber layer thickness in the papillo-macular bundle (Fig. 2a). BCVA was normal in the left eye but reduced to 0.8 in the right eye due to anisometropic amblyopia (Fig. 2a). The maternal grandmother (I:2) was already diagnosed with optic atrophy with mild (age 55: BCVA 0.7 in both eyes) but progressive vision loss (age 62: BCVA 0.4 and 0.2).

The diagnosis of ADOA was established and molecular genetic testing performed.

Family 2

The index patient (II:1, Fig. 1b) is the first child of unrelated parents of Swiss and Italian origin. The parents (I:1 and I:2) and her younger sister (II:2) did not complain of any visual dysfunction. There is a possible history of a mitochondrial disorder with myopathy and ptosis in the paternal family (paternal grandmother), but no genetic testing performed.

Personal history: The patient was born after an uneventful pregnancy and psychomotor development was normal. At age 14 years she noticed a decrease of visual acuity at distance. This led to the diagnosis of a myopia of -1.0 diopters in both eyes and glasses were prescribed. However, despite best correction, visual acuity remained at 0.63 in both eyes. Visual field showed few unspecific defects in the central 30°, mean visual field defect was 1.1 dB and 1.5 dB in the right and the left eye respectively. The patient did not notice difficulties with color vision and she correctly identified 14 and 13 out of 15 Ishihara color plates with the right eye and the left eye, respectively. Ophthalmoscopic examination demonstrated bilateral temporal optic disc pallor (Fig. 2b) and peripapillary optical coherence tomography showed a thinning of the retinal nerve fibre layer most pronounced in the papillomacular bundle (Fig. 2b). Visual...

exon 4, 4b and 5b, eight different mRNA isoforms with different expression patterns in different tissues can be produced [20]. The OPA1 protein of the main isoform counts 960 amino acids and contains 5 different domains, the N-terminal hydrophobic mitochondrial targeting domain, a N-terminal coiled-coil domain, a GTPase domain, a dynamin central region and a proposed GTPase effector domain and the C-terminal coiled-coil domain [14, 21]. OPA1 is ubiquitously expressed in human tissues with the highest levels found in retinal ganglion cells and the brain [20, 21]. OPA1 is involved in many different processes. It is the key player during mitochondrial inner membrane fusion, it plays a role in mitochondrial DNA, respiratory chain and membrane potential maintenance, cristae organization by controlling their shape and structure and it is further involved in cytochrome c mediated apoptosis [22–25]. More than 340 unique DNA variants have been allocated to OPA1 (http://mitodyn.org/home.php?select_db=OPA1) [26]. Half of these mutations result in haploinsufficiency of OPA1 by premature termination of translation leading to a truncated protein. Most of these mutations are located in the GTPase domain, whereas no mutation was found in the alternatively spliced exons (4, 4b and 5b). The phenotypic expression of OPA1 mutations in ADOA shows a high variability between families as well as members of the same family [27], often with an incomplete penetrance ranging from asymptomatic carriers and mild visual impairment to legally blind individuals [28]. ADOA can be classified either as a non-syndromic or isolated form, presenting with pure ophthalmologic features or as syndromic form associated with extra-ocular neurological manifestations, known as ADOA plus syndrome [5, 29]. Around 20% of ADOA patients develop additional neuro-muscular features, often in the third decade of life, which are sometimes present at a subclinical level. These include sensorineural deafness, ataxia, axonal sensory-motor polyneuropathy, mitochondrial myopathy and external ophthalmoplegia, whereas in one case parkinsonism and dementia were reported [30]. Many different mutations have been identified in OPA1 responsible for ADOA, including nonsense mutations, missense mutations, frameshift mutations, in-frame deletions and splice site or splice region mutations [14, 28, 31]. Since the vast majority of OPA1 mutations are predicted to result in a premature stop codon, mainly caused by splice site or splice region, nonsense or frameshift mutations, the generation of a truncated protein leading to haploinsufficiency is proposed to be the major cause for the pathogenesis of ADOA [28]. The majority of transcripts containing a premature termination codon (PTC) are degraded by a mechanism called nonsense-mediated decay (NMD), which is used to protect the cell from translation of potentially harmful proteins [32]. In this study we analyzed all coding exons including the intron-exon boundaries of OPA1 by DNA sequencing of two unrelated families presenting with clinical features of ADOA. We report two novel splice region mutations in OPA1, which we characterized at RNA level, thereby expanding the mutational spectrum of OPA1.

Methods

Patients

Family 1

The index patient (III:1, Fig. 1a) was referred to our institutions at age of 9 years because of reduced vision and optic disc pallor. The child is the first child of unrelated parents of Swiss origin. She is normally developed and besides her vision disorder a healthy child. On the first examination, best corrected visual acuity (BCVA) for distance was reduced to 0.3 and 0.5, which dropped to 0.2 and 0.32 at 10 years in the right and left eye, respectively. Temporal optic atrophy was present in both eyes. The apparently unaffected mother (II:2) showed minor optic atrophy, which was confirmed by the reduced retinal nerve fiber layer thickness in the papillo-macular bundle (Fig. 2a). BCVA was normal in the left eye but reduced to 0.8 in the right eye due to anisometropic amblyopia (Fig. 2a). The maternal grandmother (I:2) was already diagnosed with optic atrophy with mild (age 55: BCVA 0.7 in both eyes) but progressive vision loss (age 62: BCVA 0.4 and 0.2).

The diagnosis of ADOA was established and molecular genetic testing performed.

Family 2

The index patient (II:1, Fig. 1b) is the first child of unrelated parents of Swiss and Italian origin. The parents (I:1 and I:2) and her younger sister (II:2) did not complain of any visual dysfunction. There is a possible history of a mitochondrial disorder with myopathy and ptosis in the paternal family (paternal grandmother), but no genetic testing performed.

Personal history: The patient was born after an uneventful pregnancy and psychomotor development was normal. At age 14 years she noticed a decrease of visual acuity at distance. This led to the diagnosis of a myopia of -1.0 diopters in both eyes and glasses were prescribed. However, despite best correction, visual acuity remained at 0.63 in both eyes. Visual field showed few unspecific defects in the central 30°, mean visual field defect was 1.1 dB and 1.5 dB in the right and the left eye respectively. The patient did not notice difficulties with color vision and she correctly identified 14 and 13 out of 15 Ishihara color plates with the right eye and the left eye, respectively. Ophthalmoscopic examination demonstrated bilateral temporal optic disc pallor (Fig. 2b) and peripapillary optical coherence tomography showed a thinning of the retinal nerve fibre layer most pronounced in the papillomacular bundle (Fig. 2b). Visual...
Fig. 1  

**a** and **b** Pedigrees of the described and analyzed ADOA families. Index patients are indicated with an arrow. Black symbols represent affected individuals. Symbols containing 'n.a.' (not analyzed) indicate individuals with unknown genotype and/or phenotype (not described in this study).  

**c** Sequence analysis of the OPA1 gene at the genomic level. All affected family members of family 1 carrying a heterozygous intronic 22 bp insertion c.611-37_611-38insACTGGAGAATGTAAAGGGCTTT and a 11 bp deletion c.611-6_611-16delCATATTTATCT (indicated with the arrow).  

**d** Sequence analysis of OPA1 in the index patient of family 2 discloses a heterozygous intronic 4 bp deletion c.2012+4_2012+7AGTA which was absent in healthy parents and the control.
evoked potentials were not recorded and brain MRI scan ruled out a compression of the optic nerves. There was no hearing impairment but diffuse myalgia, which became manifest about the same time as the decrease in visual acuity. A thorough neurological exam did not reveal any abnormal findings apart from the visual impairment. Therefore, the tentative diagnosis of ADOA was made at age 14 years and genetic analysis of the OPA1 gene was initiated.

**Molecular genetic studies**

**Mutation screening and DNA Sequencing**

Genomic DNA (gDNA) was isolated from peripheral white blood cells of EDTA-stabilized venous blood, using Prepito DNA Blood600 kit according to the manufacturer’s instructions (PerkinElmer, Waltham, MA, USA). All coding exons (including exon 4b and 5b) of OPA1 as well as exon-intron boundaries were amplified from gDNA by PCR (primers and cycling conditions are listed in Additional file 1: Table S1). All PCR reactions were performed in a Veriti thermal cycler (Thermo Fisher, Waltham, MA, USA) and fragments were analyzed using Qiagen DNA High Resolution Cartridge (Qiagen GmbH, Hilden, Germany). PCR products were ExoSAP (Thermo Fisher, Waltham, MA, USA) purified and sequenced using BigDye Terminator chemistry v3.1 (Applied Biosystems, Foster City, CA, USA). Sequencing of both strands was performed on an ABI Prism 3500XL Genetic Analyzer (Applied Biosystems, Foster City, CA, USA). Sequence data were analyzed with SeqAnalysis 6 and SeqScape 3 (Applied Biosystems, Foster City, CA, USA). MLPA reactions were performed using the P229 kit from MRC Holland (Amsterdam, Netherlands) according to the manufacturer’s instructions. Products of the MLPA reaction were separated on an ABI Prism 3500XL Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) and were analyzed using GeneMarker Software v2.6.3. (SoftGenetics, State College, PA, USA)

GenBank transcript NM_130837 was used for variant numbering using A in ATG as number 1.

**OPA1 cDNA analysis**

Total RNA was obtained from PAX-blood (family 1, family 2 and a healthy unrelated control subject) using the PAXgene Blood RNA kit according to the manufacturer’s recommendations (PreAnalytiX GmbH, Hombrechtikon, Switzerland). Random oligohexamer primed RNA (up to 1 μg) was reverse-transcribed using the SuperScript II First-Strand Synthesis System (Invitrogen, Carlsbad, CA, USA) in a total volume of 25 μl, according to the manufacturer’s protocol. 1 μl of the single-stranded cDNA was used as template for amplification of fragments extending from OPA1 exon 4b to 5b (243 bp) and exon 17 to 23 (605 bp), using the following primer pairs: OPA1-Ex4bF (5’-CTCAGGTCACAAATTTGGTTAGT-3’), OPA1-Ex5bR (5’-CAACAGAAGCGCAAGGTGTC-3’), and OPA1-Ex17F
(5'-TCAAGGCTCTTAAAGACAAGC-3') and OPA1-Ex23R (5'-AGCTTGAGGGTTATTCACA-3'). PCR conditions using HotStar Taq DNA Polymerase (Qiagen GmbH, Hilden, Germany) included an initial denaturation step of 15 min at 95 °C, 40 cycles of denaturation at 94 °C for 30 s, annealing at 55 °C for 30 s, polymerization at 72 °C for 1 min and a final extension step at 72 °C for 10 min RT-PCR products were separated on a 12% polyacrylamide gel and extracted overnight in dH₂O at 4 °C. Isolated RT-PCR products were rated on a 12% polyacrylamide gel and extracted over-night in dH₂O at 4 °C. Isolated RT-PCR products were sepa-rated for subsequent sequencing as described above.

Results

Mutation analysis in OPA1

All coding exons of OPA1, including exon-intron boundaries, were sequenced and analyzed in two ADOA patients from two unrelated families (family 1: III:1 and family 2: II:1) and their family members (Fig. 1a and b). Sequence analysis of patient 1 (III:1, family 1) revealed a heterozygous 22 bp insertion c.611-37_611-38insACTGGAGAATG TAAAGGGCTTT proximal of a 11 bp deletion c.611-6_611-16delCATATTTATCT in intron 5 (Fig. 1c), whereas the insertion c.611-37_611-38insACTGGAGAATGTTAAAGGGCTTT could be a potential result of a tandem duplication (Fig. 4). This sequence alteration is likely to abolish the 3’ acceptor splice site located in intron 5, which was supported by different in silico prediction tools (Table 1). The patient’s mother presents with mild visual impairment and optic atrophy (II:2, family 1) and a grandmother with reduced vision and optic atrophy (II:2, family 1). Both of them were found to harbor the same heterozygous c.[611-37_611-38insACTGGAGAATGTTAAAGGGCTTT;611-6_611-16delCATATTTATCT] mutation (Fig. 1c).

In index patient 2 (II:1, family 2) we identified a heterozygous 4 bp deletion c.2012 + 4_2012 + 7delAGTA deletion in patient 1 occurred therefore de novo and is likely to be pathogenic, since it is absent in both healthy parents.

Table 1 In silico prediction of splice score changes of the natural splice site

| Prediction Tool | Threshold | Acceptor Site | Donor Site |
|-----------------|-----------|---------------|------------|
| SSF             | ≥70       | 87.40 – – – – | 87.36 – – – |
| MaxEnt          | ≥0        | 7.78 – - – - | 10.47 – 2.39 (-77.2%) |
| NNSPLICE        | ≥0.4      | 0.93 – - – - | 1.00 – – – – |
| GeneSplicer     | ≥0        | n.a.          | 2.26 – – – – |

n.a. not analysed, SSF SpliceSiteFinder-like [39], MaxEnt MaxEntScan [40], NNSPLICE Splice Site Prediction by Neural Network [41], GeneSplicer [42]; Threshold: minimal value to recognise a putative splice site; - splice site abolished

suspicous for ADOA (http://www.mitodyn.org/variants.php?select_db=OPA1&action=view&view=0000543%2C0000353%2C0) and classified as variant of unknown clinical significance (VOS). This mutation was predicted to abolish the 5’ splice donor site (Table 1). We analyzed further both unaffected parents (I:1 and I:2, family 2), who were negative for the heterozygous c.2012 + 4_2012 + 7delAGTA deletion (Fig. 1d), suggesting the mutation occurred de novo in the index patient. MLPA analysis did not give any evidence for the presence of the detected intronic mutations. Furthermore, in both index patients, the three LHON mutations m.11778G > A, p.(Arg340His) in MTND4, m.3460G > A, p.(Ala52Thr) in MTND1 and m.14484 T > C, p.(Met64Val) in MTND6 were not detected. Also no mutations were found in the nuclear genes OPA3, TMEM126A and WFS1, thus no other reason for an opticusatrophy could be identified.

OPA1 cDNA analysis

Both identified deletions have been predicted to alter the splice sites using different in silico prediction tools (Table 1). Since OPA1 is expressed in many tissues, including the white blood cells [21, 28], we investigate the impact of the deletions at RNA level using PAX-blood samples of various family members (family 1 (I:2, II:2 and III:1) and family 2 (I:1, I:2 and II:1)). cDNA analysis in family 1, showed two transcript species in all investigated and affected family members (family 1: index patient III:1, mother II:2 and grandmother I:2), including the wildtype form and an additional larger transcript (Fig. 3a). Sequence analysis of the transcripts showed that the c.[611-37_611-38insACTGGAGAATGTTAAAGGGCTTT;611-6_611-16delCATATTTATCT] results in the activation of a cryptic acceptor splice site, which is located in the wildtype sequence 166 bp upstream of the canonical 3’ splice acceptor site. Activation of this cryptic splice site leads to an addition of 177 intronic bases in the aberrantly spliced mRNA inserting a premature stop codon in the context of the appropriate reading frame of this transcript, including the 22 bp insertion c.611-37_611-38insACTGGAGAATGTTAAAGGGCTTT (Fig. 4), which is also encoded by the gDNA (Fig. 1c).

OPA1 cDNA analysis in family 2 revealed two different cDNA species, including the wildtype transcript, which was present in all analyzed individuals including the control. An additional smaller transcript was reported in the index patient (Fig. 3b). The unaffected parents, who do not harbor the deletion showed only the wildtype form (Fig. 3b). Sequence analysis of the additional smaller band found in the index patient disclosed a skipping of exons 18 and 19 (Fig. 5). The heterozygous c.2012 + 4_2012 + 7delAGTA deletion in patient 1 occurred therefore de novo and is likely to be pathogenic, since it is absent in both healthy parents.
Discussion

*OPA1* encodes a protein located in the inner mitochondrial membrane and is involved in a number of different cellular processes. It plays a key role during mitochondrial inner membrane fusion, further it is involved in mitochondrial DNA, respiratory chain and membrane potential maintenance, cristae organization by controlling their shape and structure and it has been shown to play a role in apoptosis [22–25]. Over 300 pathogenic mutations have been reported in *OPA1* [33] of which most mutations lead to the generation of a PTC and result in a loss of function of one allele (haploinsufficiency), which is
proposed to be the major mechanism underlying the pathogenesis of ADOA [20, 28]. The OPA1 protein consists of a hydrophobic transmembrane domain, a GTPase domain, a middle domain and two C-terminal coiled-coil domain. Most of the reported mutations are located within the GTPase domain and the C-terminal coiled-coil domain (exon 27 and 28) known as mutational hotspots of OPA1.

Our study indicates that ADOA in two unrelated patients and their affected family members is caused by two novel splice region mutations in OPA1. We screened the entire coding and flanking region of OPA1 in the two index patients and we identified two novel splice region mutations. The heterozygous c.[611-37_611-38insACTGGAGAATGTAAAGGGCTTT;611-6_611-16delCATATTTATCT] leads to the activation of a cryptic splice site resulting in intron retention and a de novo heterozygous 4 bp deletion c.2012 + 4_2012 + 7delAGTA leads to an in-frame exon 18 and 19 skipping. Both heterozygous splice region mutations, c.[611-37_611-38insACTGGAGAATGTAAAGGGCTTT;611-6_611-16delCATATTTATCT] in intron 5 and c.2012 + 4_2012 + 7delAGTA in intron 19 of the OPA1 gene, have not been described before.

The novel splice region mutation c.[611-37_611-38insACTGGAGAATGTAAAGGGCTTT;611-6_611-16delCATATTTATCT] identified in family 1 is likely to cause premature termination of translation of OPA1 by the formation of a PTC, leading to a truncated protein that lacks around 80% of the OPA1 protein. Furthermore, formation of a premature stop codon located within and not at the end of the transcript, more than 50-55 nucleotides upstream of the last exon-exon junction, raises the possibility that non-sense mediated decay (NMD) might be involved in degrading aberrant transcripts [32, 34]. Since we identified a mutant transcript species without inhibiting NMD, we conclude that NMD was not sufficient enough to entirely prevent the expression of truncated OPA1 and that these small amounts could interfere with both, the structural as well as the regulatory function of OPA1 in a negative way.

The de novo splice region mutation c.2012 + 4_2012 + 7delAGTA found in the affected child of family 2, leads to an in-frame skipping of exon 18 and 19 which affects the middle domain of the OPA1 protein. The middle domain of dynamins has been shown to be possibly involved in self- and higher-order assembly [35, 36]. Therefore, disruption of the middle domain of OPA1 may result in impaired mitochondrial localization which further lead to a decreased mitochondrial inner membrane fusion and a subsequent mitochondrial network fragmentation. Fragmentation of the mitochondrial network triggers decreased energy supply due to the loss of mitochondrial bioenergetics integrity on one hand and an increased production of ROS on the other hand. This leads finally to mitochondrial dysfunction and to cell death, initiated by the release of cytochrome c from mitochondria [24].

We suggest that the activation of a cryptic splice site and the in-frame exon skipping might affect the OPA1 protein stability. Therefore, we reason that the mutant OPA1 protein is unstable and gets rapidly degraded, either through non-sense mediated decay or another degradation mechanism leading to haploinsufficiency. A further speculative possibility, especially for the de novo splice region mutation c.2012 + 4_2012 + 7delAGTA, leading to an in-frame exon skipping, could
lead to an altered protein structure and function instead of resulting in haploinsufficiency, however, this would need to be elucidated by additional functional studies.

The exact pathomechanism remains unknown and therefore, further studies, preferably in vivo, would shed light on the pathogenic mechanisms playing a role during development of ADOA caused by OPA1 mutations.

We suggest that in both analyzed families the altered protein length and structure is likely the mechanism for ADOA in the affected individuals, therefore, analysis at the genomic as well as at the RNA level should be performed to investigate OPA1 defects.

Because of the mitochondrial dysfunction and the resulting impaired energy supply, the optic nerve could be primarily affected due to its higher sensitivity to energy deficiency leading to optic atrophy [37] in ADOA. To understand the pathophysiology of ADOA is important for the development of novel therapeutic strategies. To this day, there is no cure for ADOA, but a study performed by Barboni et al. in 2013, showed for the first time an improved visual function, after application of a synthetic analogue of coenzyme Q10 (Idebenone), an antioxidant used to reduce reactive oxygen species (ROS) levels in Leber’s hereditary optic neuropathy (LHON) patients, in ADOA patients with OPA1 mutations [38]. Further, identification of new mutations and their associated clinical presentation would enable an appropriate genetic counseling of the patients and their family members. Our current knowledge about the exact function and the pathomechanism of OPA1 is far from complete, but identification of new mutations associated with ADOA and their functional characterization would give us new insights into the pathogenesis of ADOA.

Conclusion

Many of the reported OPA1 mutations have only been characterized at the genomic level, only a few splice site or splice region variants have been investigated at their transcript level. In our study, we were able to characterize two novel splice region mutations at their genomic, as well as at their RNA level. Our results showed the importance of cDNA analysis to evaluate the effect of intronic DNA variants of unknown clinical significance encompassing the splice regions.

The study further expands the mutational spectrum of OPA1 in ADOA by the discovery of two novel splice region mutations and their characterization at the RNA level in individuals presenting with the clinical picture of an isolated ADOA, since there was no evidence of extra-ocular features. We could further confirm the variability of intrafamilial phenotypic expression associated with OPA1 mutations, which was shown in family 1 with its incomplete penetrance, ranging from nearly subclinical, mildly affected individuals to symptomatic family members presenting with visual impairment and optic atrophy. The clinical phenotype could be dependent on the quantity of aberrant OPA1 protein disturbing normal mitochondrial function. However, the final clinical manifestation is likely to be influenced by other endogenous and/or environmental modifying factors.

Additional file

Additional file 1: Additional 5’ M13 Sequence: F: 5’-GTAAAACGACGGCCAGT-3’ R: 5’-CAGGAAACAGCTATGAC-3’ PCR was performed according to standard PCR conditions. (DOCX 77 kb)

Abbreviations

ADOA: Autosomal dominant optic atrophy; BCVA: Best corrected visual acuity; cDNA: Complementary DNA; gDNA: Genomic DNA; LHON: Leber’s hereditary optic neuropathy; MLPA: Multiplex ligation-dependent probe amplification; NMD: Nonsense-mediated decay; PTC: Premature termination codon; ROS: Reactive oxygen species; RT-PCR: Reverse transcription polymerase chain reaction; VOUS: Variants of unknown significance

Acknowledgments

We are indebted to the families participating in this study.

Funding

This work is supported by a grant from Novartis Foundation for medical-biological research (AS).

Availability of data and material

All data generated or analysed during this study are included in this published article (and its supplementary information files) or are available from the corresponding author on reasonable request.

Authors’ contributions

AS designed and supervised the study and contributed in drafting the manuscript. RB performed molecular genetic studies and drafted the manuscript. Clinical part was written by CGK and DB. CGK, MA, DB and VS performed clinical investigations and contributed in drafting the manuscript. SG contributed in supervising and drafting the manuscript. NM participated in performing the molecular genetic studies. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Consent for publication

All participants or parents gave written informed consent for publication.

Ethics approval and consent to participate

The study was approved by the local ethical committee of the Canton of Berne and written informed consent was obtained by all participants or parents.

Author details

1Department of Ophthalmology, Cantonal Hospital St. Gallen, St. Gallen, Switzerland. 2Department of Ophthalmology, University Hospital Zurich, Zurich, Switzerland. 3Department of Ophthalmology, Cantonal Hospital St. Gallen, St. Gallen, Switzerland. 4Department of Ophthalmology, Inselspital, Bern University Hospital, University of Bern, Bern, Switzerland. 5Graduate School for Cellular and Biomedical Sciences (GCB), University of Bern, Bern, Switzerland.

Additional file

Additional file 1: Additional 5’ M13 Sequence: F: 5’-GTAAAACGACGGCCAGT-3’ R: 5’-CAGGAAACAGCTATGAC-3’ PCR was performed according to standard PCR conditions. (DOCX 77 kb)
References

1. Kjer P. Infantile optic atrophy with dominant mode of inheritance: a clinical and genetic study of 19 Danish families. Acta Ophthalmol Suppl. 1959; 164(Supp 54):1–147.

2. Delettre C, Lenaers G, Pelloquin L, Belenguer P, Hamel CP. OPA1 (Kjer type) dominant optic atrophy: a novel mitochondrial disease. Mol Genet Metab. 2002;75(2):97–107.

3. Kivlin JD, Lovrien EW, Bishop DT, Maumenee IH. Linkage analysis in dominant optic atrophy. Am J Hum Genet. 1983;35(3):1190–5.

4. Kjer B, Eiberg H, Kjer P, Rosenberg T. Dominant optic atrophy mapped to chromosome 3q region. II. Clinical and epidemiological aspects. Acta Ophthalmol Scand. 1996;74(1):3–7.

5. Amati-Bonneau P, Milea D, Bonneau D, Chevrollier A, Ferre M, Guillet V, Gueguen N, Loiseau D, de Crescenzo MA, Verny C, et al. OPA1-associated disorders: phenotypes and pathophysiology. Int J Biochem Cell Biol. 2009;41(10):1865–65.

6. Cohn AC, Toomes C, Potter C, Towns KV, Hewitt AW, Inglehearn CF, Craig JE, Mackey DA. Autosomal dominant optic atrophy: penetrance and expressivity in patients with OPA1 mutations. Am J Ophthalmol. 2007;143(4):656–62.

7. Abegg M, Zinkernagel M, Wolf S. Re: Ronnback et al.: Imaging of the macula and fine mapping of dominant optic atrophy locus on 3q28-qter by linkage disequilibrium method: a study of 38 British Isles pedigrees. Hum Genet. 1998;102(1):79–86.

8. Olichon A, Emorine LJ, Descoins E, Pelloquin L, Bichese L, Gas N, Guillou E, Valette A, Belenguer P, Lenaers G. Loss of OPA1 perturbs the mitochondrial inner membrane structure and integrity, leading to cytochrome c release and apoptosis. J Biol Chem. 2003;278(10):7743–6.

9. MacLear T, Langer T. OPA1 processing in cell death and disease - the long and short of it. J Cell Sci. 2016;129(12):2297–306.

10. Merlo P, Armani Bonne P, Tourmen Y, Malthiery Y, Reynier P. eOPA1: an online database for OPA1 mutations. Hum Mutat. 2005;25(5):423–8.

11. Votruba M, Fitzke FW, Holder GE, Carter A, Bhattacharya SS, Moore AT. Clinical features in affected individuals from 21 pedigrees with dominant optic atrophy. Arch Ophthalmol. 1998;116(6):29–30.

12. Delettre C, Moore AT, Bhattacharya SS. Clinical features, molecular genetics, and pathophysiology of dominant optic atrophy. J Med Genet. 1998;35(10):793–800.

13. Yu-Wai-Man P, Griffiths PG, Hudson G, Chinnery PF. Inherited mitochondrial optic neuropathies. J Med Genet. 2009;46(3):145–58.

14. Yu-Wai-Man P, Griffiths PG, Chinnery PF. Mitochondrial optic neuropathies - disease mechanisms and therapeutic strategies. Prog Retin Eye Res. 2011;30(2):111–14.

15. Kerrison JB, Armour VL, Ferraz Sallum JM, Vagefi MR, Barmada MM, Li Y, Zhu D, Maumenee IH. Genetic heterogeneity of dominant optic atrophy, Kjer type: Identification of a second locus on chromosome 18q12-2.3. Arch Ophthalmol. 1999;117(6):805–10.

16. Reijer P, Armani Bonne P, Votruba M, Olichon A, Simard G, Guichet A, Bonnemains C, Malecaze F, Malinge MC, Pelletier JB, et al. OPA1 gene mutations responsible for autosomal dominant optic atrophy and cataract. J Med Genet. 2004;41(10):e110.

17. Barber F, Hakiki S, Orsaas L, Gerber S, Perrault I, Hanein S, Dufier JL, Munnich A, Kaplan J, et al. A third locus for dominant optic atrophy on chromosome 22q. J Med Genet. 2005;42(1):e1.

18. Delettre C, Lenaers G, Pelloquin L, Belenguer P, Hamel CP. OPA1 (Kjer type) dominant optic atrophy mapped to chromosome 3q28-3q29, within a 3-Mb YAC contig. Hum Genet. 1997;99(1):115–20.

19. Delettre C, Griffon JM, Kaplan J, Dolfu H, Lorentz B, Faivre L, Lenaers G, Belenguer P, Hamel CP. Mutation spectrum and splicing variants in the OPA1 gene. Hum Genet. 2001;109(6):584–91.

20. Alexander C, Votruba M, Pesch UE, Thistleth DL, Mayer S, Moore A, Rodriguez M, Kellner U, Leo-Kottler B, Aeburger G, et al. OPA1, encoding a dynamin-related GTPase, is mutated in autosomal dominant optic atrophy linked to chromosome 3q28. Nat Genet. 2000;26(2):211–5.

21. Belenguer P, Pellegrini L. The dynamin GTPase OPA1: more than mitochondria? Biochim Biophys Acta. 2013;1833(1):176–83.

22. Alavi MW, Fuhrmann N. Dominant optic atrophy, OPA1, and mitochondrial quality control: understanding mitochondrial network dynamics. Mol Neurodegener. 2013;8:32.

23. Olichon A, Baricault L, Gas N, Guillou E, Valette A, Belenguer P, Lenaers G. Loss of OPA1 perturbs the mitochondrial inner membrane structure and integrity, leading to cytochrome c release and apoptosis. J Biol Chem. 2003;278(10):7743–6.

24. Mackey DA. Autosomal dominant optic atrophy: penetrance and expressivity in patients with OPA1 mutations. Am J Ophthalmol. 2007;143(4):656–62.

25. Mackey DA. OPA1 processing in cell death and disease - the long and short of it. J Cell Sci. 2016;129(12):2297–306.

26. Merlo P, Armani Bonne P, Tourmen Y, Malthiery Y, Reynier P. eOPA1: an online database for OPA1 mutations. Hum Mutat. 2005;25(5):423–8.

27. Votruba M, Fitzke FW, Holder GE, Carter A, Bhattacharya SS, Moore AT. Clinical features in affected individuals from 21 pedigrees with dominant optic atrophy. Arch Ophthalmol. 1998;116(6):29–30.

28. Petta M, Lin X, Sabzberg SL. GeneSplicer: a new computational method for splice site prediction. Nucleic Acids Res. 2001;29(15):E185–65.

Received: 6 December 2016 Accepted: 17 February 2017
Published online: 28 February 2017