Mutations in *PIK3C2A* cause syndromic short stature, skeletal abnormalities, and cataracts associated with ciliary dysfunction


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**Abstract**

*PIK3C2A* is a class II member of the phosphoinositide 3-kinase (PI3K) family that catalyzes the phosphorylation of phosphatidylinositol (PI) into PI(3)P and the phosphorylation of PI(4)P into PI(3,4)P2. At the cellular level, *PIK3C2A* is critical for the formation of cilia and for receptor mediated endocytosis, among other biological functions. We identified homozygous loss-of-function mutations in *PIK3C2A* in children from three independent consanguineous families with short stature, coarse facial features, cataracts with secondary glaucoma, multiple skeletal abnormalities, neurological manifestations, among other findings. Cellular studies of patient-derived fibroblasts found that they lacked PIK3C2A protein, had impaired cilia formation and function, and demonstrated reduced proliferative capacity. Collectively, the genetic and molecular data implicate mutations in *PIK3C2A* in a new Mendelian disorder of PI metabolism, thereby shedding light on the critical role of a class II PI3K in growth, vision, skeletal formation and neurological development. In particular, the considerable phenotypic overlap, yet distinct features,
between this syndrome and Lowe’s syndrome, which is caused by mutations in the PI-5-phosphatase **OCRL**, highlight the key role of PI metabolizing enzymes in specific developmental processes and demonstrate the unique non-redundant functions of each enzyme. This discovery expands what is known about disorders of PI metabolism and helps unravel the role of **PIK3C2A** and class II PI3Ks in health and disease.

**Author summary**

Identifying the genetic basis of rare disorders can provide insight into gene function, susceptibility to disease, guide the development of new therapeutics, improve opportunities for genetic counseling, and help clinicians evaluate and potentially treat complicated clinical presentations. However, it is estimated that the genetic basis of approximately one-half of all rare genetic disorders remains unknown. We describe one such rare disorder based on genetic and clinical evaluations of individuals from 3 unrelated consanguineous families with a similar constellation of features including short stature, coarse facial features, cataracts with secondary glaucoma, multiple skeletal abnormalities, neurological manifestations including stroke, among other findings. We discovered that these features were due to deficiency of the **PIK3C2A** enzyme. **PIK3C2A** is a class II member of the phosphoinositide 3-kinase (PI3K) family that catalyzes the phosphorylation of the lipids phosphatidylinositol (PI) into PI(3)P and the phosphorylation of PI(4)P into PI(3,4)P2 that are essential for a variety of cellular processes including cilia formation and vesicle trafficking. This syndrome is the first monogenic disorder caused by mutations in a class II PI3K family member and thus sheds new light on their role in human development.

**Introduction**

Identifying the genetic basis of diseases with Mendelian inheritance provides insight into gene function, susceptibility to disease, and can guide the development of new therapeutics. To date, ~50% of the genes underlying Mendelian phenotypes have yet to be discovered [1]. The disease genes that have been identified thus far have led to a better understanding of the pathophysiological pathways and to the development of medicinal products approved for the clinical treatment of such rare disorders [2]. Furthermore, technological advances in DNA sequencing have facilitated the identification of novel genetic mutations that result in rare Mendelian disorders [3,4]. We have applied these next-generation sequencing technologies to discover mutations in **PIK3C2A** that cause a newly identified genetic syndrome consisting of dysmorphic features, short stature, cataracts, and secondary glaucoma, multiple skeletal abnormalities, neurological manifestations including stroke, among other findings. We discovered that these features were due to deficiency of the **PIK3C2A** enzyme. **PIK3C2A** is a class II member of the phosphoinositide 3-kinase (PI3K) family that catalyzes the phosphorylation of the lipids phosphatidylinositol (PI) into PI(3)P and the phosphorylation of PI(4)P into PI(3,4)P2 that are essential for a variety of cellular processes including cilia formation and vesicle trafficking. This syndrome is the first monogenic disorder caused by mutations in a class II PI3K family member and thus sheds new light on their role in human development.
central to the pathophysiology underlying cancer, metabolic disease, and host-pathogen interactions [6].

The functions of class II PI3Ks are poorly understood relative to many other kinases and phosphatases that regulate PI metabolism, in part because there was no causal link between any class II PI3K and a monogenic human disease. In contrast, a number of disorders of PI metabolism have previously been described that have provided invaluable insight into the physiological functions of specific PI metabolizing enzymes [9]. These include Charcot-Marie-Tooth type 4J (FIG4) [10,11], Centronuclear X-linked myopathy (MTM1) [12], and primary immunodeficiency (PIK3CD) [13,14], among others. As just one example, detailed studies of FIG4 subsequent to its identification as a cause of Charcot-Marie-Tooth type 4J have revealed both genetic and physiological interactions with VAC14 and PIKFYVE, which together generate PI(3,5)P2 and are required for melanosome homeostasis, oligodendrocyte differentiation, and remyelination [15–18]. Collectively, the array of PI metabolism disorders is striking for its phenotypic diversity, affecting a wide range of organ systems including those described above as well as others that lead to neuromuscular, skeletal, renal, eye, growth, and immune disorders. The diversity of phenotypic manifestations resulting from PI metabolism defects highlights the lack of functional redundancy between genes that regulate nominally the same enzymatic transformation of PIs.

PIK3C2A has previously been attributed a wide-range of biological functions including glucose transport, angiogenesis, Akt activation, endosomal trafficking, phagosome maturation, mitotic spindle organization, exocytosis, and autophagy [19–28]. In addition, PIK3C2A is critical for the formation and function of primary cilia [23,26]. However, as mentioned above, there is as yet no link between PIK3C2A or any class II PI3K and a Mendelian disorder. Here, we describe the evidence that homozygous loss-of-function mutations in PIK3C2A cause a novel syndromic disorder involving neurological, visual, skeletal, growth, and occasionally hearing impairments.

Results

Five individuals between the ages of 8 and 21 from three unrelated consanguineous families were found by diagnostic analyses to have a similar constellation of clinical features including dysmorphic facial features, short stature, skeletal and neurological abnormalities, and cataracts (Fig 1, Table 1, S1 Table). The dysmorphic facial features included coarse faces, low hairline, epicanthal folds, flat and broad nasal bridges, and retrognathia (S1 Table). Skeletal findings included scoliosis, delayed bone age, diminished ossification of femoral heads, cervical lordosis, shortened fifth digits with mild metaphyseal dysplasia and clinodactyly, as well as dental findings such as broad maxilla incisors, narrow mandible teeth, and enamel defects (Fig 1B and 1C, S1 Table, S1 Fig). Most of the affected individuals exhibited neurological involvement including developmental delay and stroke. This was first seen in individual I-II-2 when she recently started having seizures, with an EEG demonstrating sharp waves in the central areas of the right hemisphere and short sporadic generalized epileptic seizures. Her brain MRI showed a previous stroke in the right corpus striatum (Fig 1E). Hematological studies were normal for hypercoagulability and platelet function (S2 Table). In addition, brain MRI of patient II-II-3 showed multiple small frontal and periventricular lacunar infarcts (S1E Fig). Unclear episodes of syncope also led to neurological investigations including EEG in individual III-II-2, without any signs of epilepsy. Her brain MRI showed symmetrical structures and normal cerebrospinal fluid spaces but pronounced lesions of the white matter (S1E Fig). Other recurrent features included hearing loss, secondary glaucoma, and nephrocalcinosis.
Fig 1. Pedigrees and pictures of the individuals studied. (A) Pedigree of three consanguineous families studied. Black boxes indicated affected individuals. Roman numerals indicating the generation are on the left and Arabic numerals indicating the individual are below each pedigree.
In addition to the shared syndromic features described above in all three families, both affected daughters in Family I were diagnosed with congenital adrenal hyperplasia (CAH), due to 17-alpha-hydroxylase deficiency, and were found to have a homozygous familial mutation: NM_000102.3:c.286C>T; p.(Arg96Trp) in the \textit{CYP17A1} gene (OMIM #202110) [29,30]. The affected individuals in Families II and III do not carry mutations in \textit{CYP17A1} or have CAH, suggesting the presence of two independent and unrelated conditions in Family I. The co-occurrence of multiple monogenic disorders is not uncommon among this highly consanguineous population [31].

To identify the genetic basis of this disorder, enzymatic assays related to the mucopolysaccharidosis subtypes MPS I, MPS IVA, MPS IVB, and MPSVI were tested in Families I and II and found to be normal. Enzymatic assays for mucolipidosis II/III were also normal and no pathogenic mutations were found in galactosamine-6-sulfate sulfatase (\textit{GALNS}) in Family I. Additionally, since some of the features of patient II-II-3 were reminiscent of Noonan syndrome, Hennekam syndrome, and Aarskog-Scott syndrome, individual genes involved in these disorders were analyzed in Family II, but no pathogenic mutation was identified. In patient III-II-2, Williams-Beuren syndrome was excluded in childhood. Additionally, direct molecular testing at presentation in adulthood excluded Leri-Weill syndrome, Alstrom disease, and mutations in \textit{FGFR3}.

### Table 1. Phenotypic characteristics of \textit{PIK3C2A} deficient patients.

<table>
<thead>
<tr>
<th>Family</th>
<th>I</th>
<th>I</th>
<th>II</th>
<th>II</th>
<th>III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient</td>
<td>II-1</td>
<td>II-2</td>
<td>II-2</td>
<td>II-3</td>
<td>II-2</td>
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<tr>
<td>Age (years)</td>
<td>13</td>
<td>8</td>
<td>12</td>
<td>10</td>
<td>20</td>
</tr>
<tr>
<td>Gender</td>
<td>female</td>
<td>female</td>
<td>male</td>
<td>male</td>
<td>female</td>
</tr>
<tr>
<td>Origin</td>
<td>Israel (Muslim-Arabic)</td>
<td>Israel (Muslim-Arabic)</td>
<td>Syria</td>
<td>Syria</td>
<td>Tunisia</td>
</tr>
<tr>
<td>Consanguineous</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Height</td>
<td>-3.3 SD</td>
<td>-2.3 SD</td>
<td>-2.5 SD</td>
<td>-4.8 SD</td>
<td>-2.5 SD</td>
</tr>
<tr>
<td>Weight</td>
<td>-0.2 SD</td>
<td>-1.7 SD</td>
<td>-0.2 SD</td>
<td>-3.9 SD</td>
<td>-1.9 SD</td>
</tr>
<tr>
<td>Head circumference</td>
<td>-0.25 SD</td>
<td>N.D.</td>
<td>+0.9 SD</td>
<td>-1.1 SD</td>
<td>+0.5 SD</td>
</tr>
<tr>
<td>Congenital cataract</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Secondary glaucoma</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Hearing loss</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Scoliosis/skeletal abnormalities</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Teeth abnormalities</td>
<td>+</td>
<td>+</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
</tr>
<tr>
<td>Developmental delay</td>
<td>+</td>
<td>+</td>
<td>N.D.</td>
<td>N.D.</td>
<td>N.D.</td>
</tr>
<tr>
<td>Elevated urine MPS levels</td>
<td>+</td>
<td>+</td>
<td>N.D.</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

"+" indicates presence of trait, "-" indicates absence of trait, N.D., not done.
Given the negative results of targeted genetic testing, WES and CNV analysis was performed for the affected individuals from all three families. Five homozygous candidate variants were identified in Family I, including the CYP17A1 (p.Arg96Trp) mutation that is the cause of the CAH [29,30], but is not known to cause the other phenotypes. The remaining four variants affected the genes ATF4, DNAH14, PLEKHA7, and PIK3C2A (S3 Table). In Family II, homozygous missense variants were identified in KIAA1549L, METAP1, and PEX2, in addition to a homozygous deletion in PIK3C2A that encompassed exons 1–24 out of 32 total exons (S3 Table). The deletion was limited to PIK3C2A and did not affect the neighboring genes. Sequence analysis of Family III showed a homozygous missense variant in PTH2R, nonsense variant in DPRX, and splice site variant in PIK3C2A (S3 Table).

Sequencing analyses revealed that all affected family members in the Families I, II, and III were homozygous for predicted loss-of-function variants in PIK3C2A, and none of the unaffected family members were homozygous for the PIK3C2A variants (Fig 1A and 1G). The initial link between these three families with rare mutations in PIK3C2A was made possible through the sharing of information via the GeneMatcher website [3]. The PIK3C2A deletion in Family II was confirmed by multiplex amplicon quantification (S2A Fig). The single nucleotide PIK3C2A variants in Families I and III were confirmed by Sanger sequencing (S2B and S2C Fig).

In Family I, the nonsense mutation in PIK3C2A (p.Tyr195*) truncates 1,492 amino acids from a protein that is 1,686 amino acids. This is predicted to eliminate nearly all functional domains including the catalytic kinase domain, and is expected to trigger nonsense-mediated mRNA decay [25]. Accordingly, levels of PIK3C2A mRNA are significantly decreased in both heterozygous and homozygous individuals carrying the p.Tyr195* variant (Fig 2A). The deletion in Family II eliminates the first 24 exons of the 32-exon PIK3C2A gene and is thus predicted to cause a loss of protein expression. This is consistent with a lack of PIK3C2A mRNA expression (Fig 2B). The variant in PIK3C2A in Family III affects an essential splice site (c.1640+1G>T) that leads to decreased mRNA levels (Fig 2C). Deep sequencing of the RT-PCR products revealed 4 alternative transcripts in patient-derived lymphocytes (p.[Asn483_Arg547delinsLys, Ala521Thrfs*4, Ala521_Glu568del, and Arg547SerinsTyrIleIle*]) of which the transcript encoding p. Asn483_Arg547delinsLys that skips both exons 5 and 6 was also observed in patient’s fibroblasts (S3 Fig). Although this transcript remains in-frame, no PIK3C2A protein was detected by Western blotting (Fig 2D and 2F). This is consistent with Families I and II, for which Western blotting also failed to detect any full-length PIK3C2A in fibroblasts from the affected homozygous children (Fig 2E and 2F). Thus, all three PIK3C2A variants likely encode loss-of-function alleles. Importantly, among the 141,456 WES and whole genome sequences from control individuals in the Genome Aggregation Database (gnomAD v2.1) [32], none are homozygous for loss-of-function mutations in PIK3C2A, which is consistent with total PIK3C2A deficiency causing severe early onset disease.

To test whether the observed loss-of-function mutations in PIK3C2A cause cellular phenotypes consistent with loss of PIK3C2A function, we examined PI metabolism, cilia formation and function, and cellular proliferation rates. PIK3C2A deficiency in the patient-derived fibroblasts decreased the levels of PI(3,4)P2 throughout the cell (Fig 3A) as well as decreased the levels of PI(3)P at the ciliary base (Figs 3B and S4A). The reduction in PI(3)P at the ciliary base was associated with a reduction in ciliary length (Fig 4A), although the percentage of ciliated cells was not altered (Fig 4B). Additional cilia defects include a reduction in ciliary length (Fig 4A), although the percentage of ciliated cells was not altered (Fig 4B). Additional cilia defects include a reduction in the levels of RAB11 at the ciliary base (Figs 4C and S4B), which functions within a GTPase cascade culminating in the activation of RAB8, which together with ARL13B selectively traffics ciliary proteins to the cilium [33]. Additionally, there was increased accumulation of IFT88 along the length of the cilium (Figs 4D and S4C), which is a component of the intraflagellar transport
sub-complex IFT-B, and is essential for the trafficking of ciliary protein cargoes along the axonemal microtubules [34,35]. Together, these findings are suggestive of defective trafficking of ciliary components. Finally, the proliferative capacity of PIK3C2A deficient cells was reduced relative to control cells (Fig 5).

Fig 2. Protein and mRNA levels of PIK3C2A in patient-derived cells. PIK3C2A mRNA levels were detected by qRT-PCR in patient derived fibroblasts from (A) Family I, (B) Family II, and (C) Family III. The WT sample in Fig 2A is based on n = 1 individual, the WT sample in Fig 2B is based on n = 3 individuals, and the WT sample in Fig 2C is based on n = 1 individual, totaling n = 5 different WT samples. qRT-PCR data is represented as mean ± SEM (n = 3–4 technical replicates per sample). (D, E) Whole cell lysates from fibroblasts of healthy controls (WT), heterozygous parents, and affected individuals from (D) Family III and (E) Families I and II were analyzed by Western blotting for PIK3C2A and the loading controls Actin or GAPDH. Immunogen of anti-PIK3C2A antibodies (AB1-AB4) are detailed in S5 Table. (F) Densitometry of Western blot results was performed using ImageJ. Individual samples are shown with the data combined from the four different PIK3C2A antibodies used, with the exception of the WT samples which includes fibroblasts from two individuals. * indicates p < 0.05, ** indicates p < 0.01, *** indicates p < 0.0001.

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Mutations in PIK3C2A cause syndromic short stature
As PIK3C2A is a member of the class II PI3K family, we tested whether the expression of the other family members PIK3C2B and PIK3C2G were altered by PIK3C2A deficiency. The expression of PIK3C2G was not detected by qRT-PCR in either patient-derived or control primary fibroblasts. This is consistent with the relatively restricted expression pattern of this gene in the GTEx portal [36], with expression largely limited to stomach, skin, liver, esophagus, mammary tissue, and kidney, but absent in fibroblast cells and most other tissues. In contrast, PIK3C2B expression was detected, with both mRNA and protein levels significantly increased in PIK3C2A deficient cells (Fig 6A–6D). Downregulation of PIK3C2A using an inducible shRNA in HeLa cells also resulted in elevated levels of PIK3C2B (Fig 6E). Together, these data are consistent with increased levels of PIK3C2B serving to partially compensate for PIK3C2A deficiency.

**Discussion**

Here we describe the identification of three independent families with homozygous loss-of-function mutations in PIK3C2A resulting in a novel syndrome consisting of short stature, cataracts, secondary glaucoma, and skeletal abnormalities among other features. Patient-derived fibroblasts had decreased levels of PI(3,4)P2 and PI(3)P, shortening of the cilia and impaired
Fig 4. Ciliary defects due to PIK3C2A deficiency in patient-derived fibroblasts. (A) Cilia length and (B) cilia number were determined in primary fibroblasts from two affected individuals and three unrelated controls. Data is represented as mean ± SEM (n>300/sample). Raw data for cilia length is provided in S6 Table. (C) Immunofluorescence analysis of RAB11 localization at the base of the primary cilium. Results showed that RAB11 is significantly reduced at the base of the primary cilium in -/- cells compared with +/+ and or +/- cells. n = 3 independent experiments and 15 cells.
ciliary protein localization, and reduced proliferation capacity. Thus, based on the loss-of-function mutations in *PIK3C2A*, the phenotypic overlap between the three independent families, and the patient-derived cellular data consistent with previous studies of PIK3C2A function, we conclude that loss-of-function mutations in *PIK3C2A* cause this novel syndrome.

The identification of *PIK3C2A* loss-of-function mutations in humans represents the first mutations identified in any class II PI-3-kinase in a disorder with a Mendelian inheritance, and thus sheds light into the biological role of this poorly understood class of PI3Ks [7,37]. This is significant not only for understanding the role of *PIK3C2A* in rare monogenic disorders, but also the potential contribution of common variants in *PIK3C2A* in more genetically complex disorders. There are now numerous examples where severe mutations in a gene cause a rare Mendelian disorder, whereas more common variants in the same gene, with a less deleterious effect on protein function, are associated with polygenic human traits and disorders [38–40]. For example, severe mutations in *PPARG* cause monogenic lipodystrophy, whereas less severe variants are associated with complex polygenic forms of lipodystrophy [41,42]. In the case of PIK3C2A deficiency, the identification of various neurological features including developmental delay, selective mutism, and the brain abnormalities detected by MRI (S1

![Fig 5. PIK3C2A deficiency causes delayed proliferation rates in patient-derived fibroblasts.](https://doi.org/10.1371/journal.pgen.1008088.g005)
Table) may provide biological insight into the mechanisms underlying the association between common variants in PIK3C2A and schizophrenia [43–45].

Other monogenic disorders of phosphoinositide metabolism include Lowe’s syndrome and Joubert syndrome, which can be caused by mutations in the inositol polyphosphate 5-phosphatases OCRL and INPP5E, respectively [46]. All three of these disorders of PI metabolism affect some of the same organ systems, namely the brain, eye, and kidney. However, the phenotype associated with mutations in INPP5E is quite distinct, and includes cerebellar vermis hypo-dysplasia, coloboma, hypotonia, ataxia, and neonatal breathing dysregulation [47]. In contrast, the phenotypes associated with Lowe’s syndrome share many of the same features with PIK3C2A deficiency including congenital cataracts, secondary glaucoma, kidney defects, skeletal abnormalities, developmental delay, and short stature [9,48]. The enzyme defective in Lowe’s syndrome, OCRL, is functionally similar to PIK3C2A as well, as it is also required for membrane trafficking and ciliogenesis [49]. The similarities between Lowe’s syndrome and PIK3C2A deficiency suggest that similar defects in phosphatidylinositol metabolism may underlie both disorders. In addition to Lowe’s syndrome, there is partial overlap between

Fig 6. PIK3C2B levels are increased by PIK3C2A deficiency. PIK3C2B mRNA levels were detected by qRT-PCR in (A) Family I, (B) Family II, and (C) Family III. qRT-PCR data is represented as mean ± SEM (n = 3 technical replicates per sample). The WT sample in Fig 6A is based on n = 1 and the WT sample in Fig 6B is based on n = 3, totaling n = 4 different WT samples. (D) PIK3C2B protein levels were detected by Western blotting in Family I from 2 independent experiments. (E) PIK3C2A and PIK3C2B protein levels were analyzed in HeLa cells by Western blotting following doxycycline inducible shRNA mediated knockdown of PIK3C2A. Data shown is representative from 3 independent experiments. * indicates p < 0.05, ** indicates p < 0.01. *** indicates p < 0.0001.

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PIK3C2A deficiency and yet other Mendelian disorders of PI metabolism such as the early-onset cataracts in patients with INPP5K deficiency [50,51], demonstrating the importance of PI metabolism in lens development.

The viability of humans with PIK3C2A deficiency is in stark contrast to mouse Pik3c2a knockout models that result in growth retardation by e8.5 and embryonic lethality between e10.5–11.5 due to vascular defects [20]. One potential explanation for this discrepancy is functional differences between human PIK3C2A and the mouse ortholog. However, the involvement of both human and mouse PIK3C2A in cilia formation, PI metabolism, and cellular proliferation suggests a high degree of functional conservation at the cellular level [26,28]. An alternate possibility is that the species viability differences associated with PIK3C2A deficiency result from altered compensation from other PI metabolizing enzymes. For instance, there are species-specific differences between humans and mice in the transcription and splicing of the OCRL homolog INPP5B that may uniquely contribute to PI metabolism in each species [52].

Alternately, PIK3C2B levels were significantly increased in human PIK3C2A deficient cells, including both patient-derived cells and HeLa cells surviving PIK3C2A deletion, suggesting that this may partially compensate for the lack of PIK3C2A in humans, although it remains to be determined whether a similar compensatory pathway exists in mice.

It is intriguing that both PIK3C2A and OCRL have important roles in primary cilia formation [26,53,54]. Primary cilia are evolutionary conserved microtubule-derived cellular organelles that protrude from the surface of most mammalian cell types. Primary cilia formation is initiated by a cascade of processes involving the targeted trafficking and docking of Golgi-derived vesicles near the mother centriole. They play a pivotal role in a number of processes, such as left-right patterning during embryonic development, cell growth, and differentiation. Abnormal phosphatidylinositol metabolism results in ciliary dysfunction [55], including loss of PIK3C2A that impairs ciliogenesis in mouse embryonic fibroblasts, likely due to defective trafficking of ciliary components [26]. The importance of primary cilia in embryonic development and tissue homeostasis has become evident over the two past decades, as a number of proteins which localize to the cilium harbor defects causing syndromic diseases, collectively known as ciliopathies [56,57].

Hallmark features of ciliopathies share many features with PIK3C2A deficiency and include skeletal abnormalities, progressive vision and hearing loss, mild to severe intellectual disabilities, polydactyly, and kidney phenotypes. Many of these disorders, including Bardet-Biedl Syndrome, Meckel Syndrome, and Joubert Syndrome are also associated with decreased cilium length [58], as seen in PIK3C2A deficient cells. Ciliary length is a function of both axoneme elongation and cilium disassembly, and is molecularly regulated by intraflagellar protein transport, including the velocity of transport and cargo loading, as well as soluble tubulin levels and microtubule modifications [59,60]. As defects in intraflagellar protein transport were likely indicated by abnormal IFT88 localization along the length of the cilium in PIK3C2A deficient cells, this may represent a potential mechanism underlying the shortened cilium. Further work and the identification of additional patients with mutations in PIK3C2A will continue to improve our understanding of the genotype-phenotype correlation associated with PIK3C2A deficiency. However, the identification of the first patients with PIK3C2A deficiency establishes a role for PIK3C2A in neurological and skeletal development, as well as vision, and implicates loss-of-function PIK3C2A mutations as a potentially new cause of a cilia-associated disease.

**Materials and methods**

**Ethics statement**

The study was approved by the Helsinki Ethics Committees of Rambam Health Care Campus (#0038-14-RMB), the University Hospital Institutional Review Board for Case Western
Reserve University (#NHR-15-39), the Ethics Committee of the Friedrich-Alexander University Erlangen-Nürnberg (#164_15 B), and was in accordance with the regulations of the University Medical Center Groningen’s ethical committee. Written informed consent was obtained from all participants.

**Whole exome sequencing**

Whole exome sequencing (WES) of two patients from Family I was performed using DNA (1μg) extracted from whole blood and fragmented and enriched using the Trueq DNA PCR Free kit (Illumina). Samples were sequenced on a HiSeq2500 (Illumina) with 2x100bp read length and analyzed as described [61]. Raw fastq files were mapped to the reference human genome GRCh37 using BWA [62] (v.0.7.12). Duplicate reads were removed by Picard (v. 1.119) and local realignment and base quality score recalibration was performed following the GATK pipeline [63] (v. 3.3). The average read depth was 98x (I-II-1) and 117x (I-II-2). HaploTypeCaller was used to call SNPs and indels and variants were further annotated with Annovar [64]. Databases used in Annovar were RefSeq [65], Exome Aggregation Consortium (ExAC) [32] (v. exac03), ClinVar [66] (v. clinvar_20150330) and LJB database [67] (v. ljb26_all). Exome variants in Family I were filtered out if they were not homozygous in both affected individuals, had a population allele frequency greater than 0.1% in either the ExAC database [32] or the Greater Middle East Variome Project [68], and were not predicted to be deleterious by either SIFT [69] or Polyphen2 [70].

Whole exome sequencing was performed on the two affected individuals of Family II and both their parents essentially as previously described [71]. Target regions were enriched using the Agilent SureSelectXT Human All Exon 50Mb Kit. Whole-exome sequencing was performed on the Illumina HiSeq platform (BGI Europe) followed by data processing with BWA [62] (read alignment) and GATK [63] (variant calling) software packages. Variants were annotated using an in-house developed pipeline. Prioritization of variants was done by an in-house designed ‘variant interface’ and manual curation.

The DNAs of Family III were enriched using the SureSelect Human All Exon Kit v6 (Agilent) and sequenced on an Illumina HiSeq 2500 (Illumina). Alignment, variant calling, and annotation were performed as described [72]. The average read depth was 95x (III-II-2), 119x (III-I-1) and 113x (III-I-2). Variants were selected that were covered by at least 10% of the average coverage of each exome and for which at least 5 novel alleles were detected from 2 or more callers. All modes of inheritance were analyzed [72]. Variants were prioritized based on a population frequency of 10^{-3} or below (based on the ExAC database [32] and an in-house variant database), on the evolutionary conservation, and on the mutation severity prediction.

All candidate variants in Families I, II, and III were confirmed by Sanger sequencing (primers listed in S4 Table).

**Copy number variant (CNV) analysis**

Microarray analysis for CNV detection in Family I was performed using a HumanOmniv5--Quad chip (Illumina). SNP array raw data was mapped to the reference human genome GRCh37 and analyzed with GenomeStudio (v. 2011/1). Signal intensity files with Log R ratio and B-allele frequency were further analyzed with PennCNV [73] (v. 2014/5/7). In Family III the diagnostic chromosomal microarray analysis was performed with an Affymetrix CytoScan HD-Array and analyzed using Affymetrix Chromosome analysis Suite-Software, compared with the Database of Genomic Variants and 820 in house controls. All findings refer to UCSC Genome Browser on Human, February 2009 Assembly (hg19), Human Genome built 37.
CNV analysis on the WES data of Families II and III were performed using CoNIFER [74]. Variants were annotated using an in-house developed pipeline. Prioritization of variants was done by an in-house designed ‘variant interface’ and manual curation as described before [75]. Subsequent segregation analysis of the pathogenic CNV in Family II was performed with MAQ by using a targeted primer set with primers in exons 3, 10, 20 and 24 which are located within the deletion and exons 28, 32, 34 which are located outside of the deletion (Multiplex Amplicon Quantification (MAQ); Multiplicom).

**Cell culture**

Human dermal fibroblasts were obtained from sterile skin punches cultured in DMEM (Dulbecco’s Modified Eagle’s Medium) supplemented with 10–20% Fetal Calf Serum, 1% Sodium Pyruvate and 1% Penicillin and streptomycin (P/S) in 5% CO2 at 37˚C. Control fibroblasts were obtained from healthy age-matched volunteers. Fibroblasts from passages 4–8 were used for the experiments. To measure cell proliferation, cells were detached using trypsin and counted with an Automated Cell Counter (ThermoFisher). Cells (n = 2500) were plated in triplicate in 96-well plates. Viability was measured at day 2, 4, 6 and 8. Each measurement was normalized to day 0 (measured the day after plating) and expressed as a fold increase. Viability was assessed by using CellTiter-Glo Luminescent Cell Viability Assay (Promega). Three independent experiments were performed.

**Inducible knockdown of PIK3C2A**

HeLa cells were infected with lentiviral particles containing pLKO-TET-PI3KC2A-shRNA or pLKO-TET-scramble-shRNA in six-well plates (n = 50,000 cells). After two days, the medium containing lentiviral particles was replaced with DMEM 10% FBS, 1.5 μg/ml puromycin. After 7 days of selection, cells were detached and 100,000 cells were plated in six-well plates in triplicate in the presence of doxycycline (0.5, 1 and 2 μg/ml). Medium containing doxycycline was replaced every 48 hours. After 10 days of doxycycline treatment, cells were lysed and analysed by Western blot.

**cDNA and quantitative real-time-PCR**

Total RNA was purified from primary fibroblasts using the PureLink RNA purification kit (ThermoFisher) or RNPure peqGOLD (Peqlab). RNA was reverse transcribed into complementary DNA with random hexamer using a high-Capacity cDNA Reverse Transcription Kit (ThermoFisher). RT-PCR from lymphocytes to detect exon-skipping in family III was performed using primers flanking exon 6. The resulting product was sequenced on an Illumina HiSeq2500 (Illumina) to detect splicing variants with high sensitivity. Gene expression was quantified by SYBR Green real-time PCR using the CFX Connect Real-Time System (BioRad). Primers used are detailed in S4 Table. Expression levels were calculated using the ΔΔCT method relative to GADPH.

**Western blotting**

Protein was extracted from cultured primary fibroblast cells as described [76,77]. Extracts were quantified using the DC protein assay (BioRad) or the BCA method. Equal amounts of protein were separated by SDS-PAGE and electrotransferred onto polyvinylidene difluoride membranes (Millipore). Membranes were blocked with TBST/5% fat-free dried milk and stained with antibodies as detailed in S5 Table. Secondary antibodies were goat anti-rabbit
Immunostaining

Primary fibroblasts were grown on glass coverslips to approximately 80% - 90% confluency in DMEM + 10% FCS + 1% P/S, at which time the medium was replaced with DMEM without FCS for 48 hours to induce ciliogenesis. Cells were fixed in either methanol for 10 minutes at -20°C or 4% paraformaldehyde for 10 minutes at room temperature (RT). Fixed cells were washed in PBS, and incubated with 10% normal goat serum, 1% bovine serum albumin in PBS for 1 hour at RT. If cells were fixed with paraformaldehyde, blocking solutions contained 0.5% Triton X-100. Cells were incubated with primary antibody overnight at 4°C, washed in PBS, and incubated with secondary antibody including 4',6-diamidino-2-phenylindole (DAPI) to stain nuclei for 1 hours at RT. Coverslips were mounted on glass slides with fluoromount (Science Services) and imaged on a confocal laser scanning system with a 63x objectives (LSM 710, Carl Zeiss MicroImaging). Primary antibodies are detailed in S5 Table.

Cilia analysis

To induce ciliogenesis, cells were grown in DMEM with 0–0.2% FCS for 48 hours. Cells were washed in PBS, then fixed and permeabilized in ice-cold methanol for 5 minutes, followed by extensive washing with PBS. After blocking in 5% Bovine Serum Albumin, cells were incubated with primary antibodies for 1.5 hours at RT and extensively washed in PBS-T. Primary antibodies used for Centrin and ARL13B are detailed in S5 Table. To wash off the primary antibody, cells were extensively washed in PBS-T. Subsequently, cells were incubated with secondary antibodies, Alexa Flour 488 (1:800, Invitrogen) and Alexa Fluor 568 (1:800, Invitrogen), for 45 min followed by washing with PBS-T. Finally, cells were shortly rinsed in ddH2O and samples were mounted using Vectashield with DAPI. Images were taken using an Axio Imager Z2 microscope with an Apotome (Zeiss) at 63x magnification. Cilia were measured manually using Fiji software taking the whole length of the cilium based on ARL13B staining. At least 300 cilia were measured per sample. Cilia lengths were pooled for 3 control cell lines and compare to 2 patient-derived samples (II-II-2 and II-II-3). Statistical significance was calculated using a Student t-test. PI(3)P at the ciliary base was detected in randomly chosen cells using the same exposure for each acquisition. A specific anti-PI(3)P antibody (Echelo Z-P003) was used to quantify the PI(3)P by measuring the green fluorescent intensity around the ciliary base in a region with a diameter of 8 μm and a depth of 10 μm as previously illustrated and described [26].

Supporting information

S1 Fig. Images of individuals with PIK3C2A deficiency. Photographic images of (A) teeth, (B) hands, and (C) feet are shown from the five individuals with PIK3C2A deficiency. (D) X-Ray images of the pelvis and (E) MRI images of the brain are shown when available. White arrows in the MRI images indicate regions of altered signal intensity.

S2 Fig. Homozygous loss-of-function mutations in PIK3C2A. (A) CNV analysis confirmed a homozygous deletion encompassing exons 1–24 out of 32 total exons of PIK3C2A, indicated with the red line (B) Sanger sequencing confirmed homozygosity for the PIK3C2A c.585T variant in Family I. (C) Sanger sequencing confirmed homozygosity for the PIK3C2A c.1640+1
G>T variant in Family III.

S3 Fig. The c.1650+1G>T mutation in PIK3C2A disrupts the splice donor site in intron 6. (A) Deep sequencing of RT-PCR products revealed 4 alternative transcripts in lymphocytes (red lines) compared to control samples (blue lines): "*: r.1561_1704del; p.Ala521_Glu568del, "**: r.1640+1_1640+27; p. Arg547SerinsTyrIleIle; "***: r.1561_1640del; p. Ala521Thrfs*4; ****: r.1449_1640del; p. Asn483_Arg547delinsLys. Exon/intron structure of PIK3C2A (exons: yellow boxes) with transcripts of the patient of family III compared to controls. (B) Example of sequenced RT-PCR products from cDNA of fibroblasts from wild-type control and the patient of family III using primers located in exons 3 and 10. Skipping of exons 5 and 6 is the result of the mutation. Positions of primers are indicated by orange arrows and position of the splice site mutation is indicated by a black arrow.

S4 Fig. Quantification of fluorescence in patient-derived fibroblasts. Quantification of fluorescence intensity for (A) PI(3)P at the ciliary base, (B) RAB11 at the ciliary base, and (C) ciliary IFT88. Data is shown separately for each individual family as indicated.

S1 Table. Detailed phenotypic characteristics of patients in Families I, II, and III.
S2 Table. Hematological evaluation of patients in Family I.
S3 Table. Homozygous candidate variants identified by whole exome sequencing.
S4 Table. List of primers used in this study.
S5 Table. List of antibodies used in this study.
S6 Table. Raw data underlying Fig 3 and Fig 4A.

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References


