Deep exploration of a CDKN1C mutation causing a mixture of Beckwith-Wiedemann and IMAGE syndromes revealed a novel transcript associated with developmental delay

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ABSTRACT
Background Loss-of-function mutations in CDKN1C cause overgrowth, which is, Beckwith-Wiedemann syndrome (BWS), while gain-of-function variants in the gene’s PCNA binding motif cause a growth-restricted condition called IMAGE syndrome. We report on a boy with a remarkable mixture of both syndromes, with developmental delay and microcephaly as additional features.

Methods Whole-exome sequencing and ultra-deep RNA sequencing of leucocyte-derived fibroblast-derived mRNA were performed in the family.

Results We found a maternally inherited variant in the IMAGE hotspot region: NM_000076.2(CDKN1C) c.822_826delinsGAGCTG. The asymptomatic mother had inherited this variant from her mosaic father with mild BWS features. This delins caused tissue-specific frameshifting resulting in at least three novel mRNA transcripts in the boy. First, a splice product causing CDKN1C truncation was the likely cause of BWS. Second, an alternative splice product in fibroblasts encoded IMAge-associated amino acid substitutions. Third, we speculate that developmental delay is caused by a change in the alternative CDKN1C-201 (ENST00000380725.1) transcript, encoding a novel isoform we call D (UniProtKB: A6NK88). Isoform D is distinguished from isoforms A and B by alternative splicing within exon 1 that changes the reading frame of the last coding exon. Remarkably, this delins changed the reading frame back to the isoform A/B type, resulting in a hybrid D–A/B isoform.

Conclusion Three different cell-type-dependent RNA products can explain the co-occurrence of both BWS and IMAGE features in the boy. Possibly, brain expression of hybrid isoform D–A/B is the cause of developmental delay and microcephaly, a phenotypic feature not previously reported in CDKN1C patients.

INTRODUCTION
The imprinted and predominantly maternally expressed cell cycle inhibitor CDKN1C located to 11p15.5 encodes a negative growth regulator also called p57kip1. This gene is associated with two contrasting phenotypes: the overgrowth disorder Beckwith-Wiedemann syndrome (BWS [MIM: 130650]) and the growth-restricted IMAGE syndrome (MIM: 614732), an acronym for Intrauterine growth restriction, Metaphysial dysplasia, Adrenal insufficiency and Genital abnormalities.1 IMAGE and rare cases of Silver-Russel syndrome (SRS) are caused by gain-of-function (GoF) mutations in the PCNA-binding motif of CDKN1C, while loss-of-function (LoF) mutations are found in 8% of BWS cases.2–4

CDKN1C is a cyclin-dependent kinase inhibitor (CDI) belonging to the CIP/Kip family and encoded by a small gene with three to four exons of which two to three are protein coding. Other members are p21Cip1 (CDKN1A) and p27Kip1 (CDKN1B), which inhibit cell growth in G1 and can cause cell-cycle arrest. The ENSEMBL database (ensembl.org) contains several CDKN1C transcripts encoding different isoforms; CDKN1C-201 encodes the 316aa isoform A, CDKN1C-203 encodes an alternative isoform A with different UTRs, CDKN1C-202 encodes the 305aa isoform B with an alternative start site and CDKN1C-201 encodes the 131aa A6NK88 isoform that we call isoform D (see figure 1). CDKN1C has an N-terminal CDI (cyclin-dependent kinase inhibitor) domain, a central PAAP (proline-alanine) repeat and C-terminally the PCNA (proliferating cell nuclear antigen)–binding motif and an overlapping putative KRRK-containing nuclear localisation signal.

BWS is most often caused by sporadic loss-of-methylation (LOM) of imprinting centre 2 (IC2) on the maternal chromosome 11 that results in diminished CDKN1C expression, found in about 50%–60% of BWS, followed by paternal segmental uniparental disomy (UPD) 11p also involving the paternally methylated imprinting centre 1 (IC1) of the IGF2-H19 domain in about 20%.5 IC1 gain-of-methylation (GoM) has been reported in 5%–10% of the patients, whereas microdeletions of IC1 and CDKN1C variants are important causes of familial BWS. Other chromosomal CNVs, balanced rearrangements, multilocus imprinting disturbances and mosaic genome-wide paternal UPD are rare causes of BWS. Typical features are omphalocele/umbilical hernia, macroglossia, neonatal hyperinsulinism, Wilms tumour, lateralised overgrowth, macrosomia, ear creases and pits, large internal organs, adrenal fetal cortex cytomegaly and renal abnormalities.5 Genotype-phenotype correlations are reported as CDKN1C mutations have higher rates of preterm birth, cleft palate, abdominal wall defect, capillary malformations and ear signs,
while intrauterine overgrowth and tumours are less common.\(^7^,\,^8\) CDKN1C-restricted BWS features are also reported, including maternal pre-eclampsia, genital abnormalities, polydactyly, polydactyly and posterior fossa abnormalities.\(^7^,\,^9^–\,^12\)

Maternally inherited GoF variants can cause IMAGe syndrome, an SRS-like phenotype,\(^13\,^\text{and}^\) shorter stature with diabetes,\(^14\) but such variants are rare causes of growth restriction.\(^15^–\,\text{and}^\text{17}\) IMAGe was first described in 1999\(^\text{and}^\text{linked to}^\text{CDKN1C in 2012.}^\) Besides, SRS phenotype with salt-wasting due to adrenal insufficiency mimicking IMAGe was described in two patients with 11p15 maternal duplications.\(^19\) So far, only 12 molecularly confirmed IMAGe and BWS patients have both GoF and LoF effects and to find a potential explanation for the affected brain function. Our work revealed that a novel CDKN1C transcript with a non-canonical C-terminal reading frame encoding isoform A6NK88 (named isoform D by us) was also affected by the delins, and as isoform D is also expressed in the brain, this could be the cause of his developmental delay.

**METHODS**

The boy and his family were clinically examined, given genetic counselling and consented to RNA analysis to investigate the molecular consequences of the CDKN1C variant.

**Sanger sequencing**

We used CDKN1C reference sequence NM_000076.2 encoding a 316 aa protein (isoform A). Sequence pilot (JSI Medical Systems, Ettenheim, Germany) software was used for interpretation. DNA was isolated from peripheral blood obtained from the boy, his parents, his mother’s siblings, maternal grandparents and maternal grandfather’s parents, and sequenced. To explore the mosaic pattern in the maternal grandfather, we analysed additional DNA obtained from buccal swab, urine and fibroblasts from four distinct skin biopsies.

**RNA isolation, sequencing of cDNA and qPCR**

Total RNA was isolated from fibroblasts from biopsies of healthy skin (two biopsies from the boy, one from his mother and four from his grandfather), one epidermal nevus in the boy and also from blood samples (from the boy, both parents and the grandfather) by using the RNeasy Mini Kit (QIAGEN, Hilden, Germany) for fibroblasts and PAXgene Blood RNA Kit (QIAGEN) for blood samples. cDNA was synthesised following the manufacturer’s protocol. The variant was amplified by quantitative qRT-PCR using different primer sets taking alternative acceptor site usage into account and also spanning the variant.
affecting transcript ENST00000380725.1 CDKN1C-201 (for a full list of transcripts, see figure 1). The primer sequences are given in online supplemental file. We examined expression levels of wt (wild type) and mutant (delins) CDKN1C, and wt CDKN1C-201 by qRT-PCR on RNA isolated from both blood and cultured fibroblasts from the boy (all biopsies), his mother and grandfather (where we picked the biopsy with highest mutation load), and blood from his father. Primers and probes were from Applied Biosystems, Life Technologies, and assessment of B2M (beta-2-microglobulin) RNA (Applied Biosystems) served as endogenous controls. At least two runs per experiment were performed, and two different probes for both CDKN1C-201 and mutant CDKN1C were analysed.

**Total RNA deep sequencing**

We used blood-derived RNA from the boy, his mother and grandfather, and an unrelated healthy adult female, and fibroblast RNA from the boy’s skin biopsy #2. Whole transcriptome sequencing libraries were generated using the Illumina TrueSeq Stranded Total RNA kit with Ribo-Zero Gold depletion for fibroblasts and Ribo-Zero Globin depletion for blood, according to the manufacturer’s protocols. Libraries were quality checked on the Agilent Bioanalyzer system and accurately quantified using the KAPA qPCR quantification kit. Libraries were paired-end sequenced on the Illumina HiSeq4000 system with a read length of 2 x 75 nt. Sequencing was performed ultra-deep for all libraries (~ 250 million reads per sample). RNA-sequencing reads were aligned to the human genome reference (GRCh38) using HISAT2 (V2.0.5). Reads aligned within the coding part of the genome (adequate Gencode gtf annotation file) were counted using featureCounts. To resolve alternative CDKN1C transcripts, we created supplementary reference files in FASTA format and realigned reads. Data were visualised in IGV (Integrative Genomics Viewer V2.3.74) (online supplemental file). We only considered splice junction tracks (SJTs) when counting reads and manually checked all reads in a region of interest. All samples were run in the same flow cell.

**Whole-exome sequencing**

Whole-exome sequencing (WES) was performed on genomic DNA isolated from blood from the patient and parents. DNA samples were prepared using SeqCap EZ MedExome Target Enrichment Kit (Roche NimbleGen, Madison, WI) and followed by paired-end 150 nt sequencing on an Illumina NextSeq500. The paired-end reads were aligned using the Burrows-Wheeler Alignment tool and variant calling was performed using the Genome Analysis Toolkit (GATK; Broad Institute, Cambridge, MA) according to GATK’s Best Practices guidelines. Mean #reads per base pair in the exome was 83X with 97.7% of the base pairs covered at least 10 times. Data annotation and interpretation were performed using the NGS module of Cartagenia Bench Lab software (Agilent, Santa Clara, CA).

**CNV and methylation analysis**

The boy’s DNA isolated from peripheral blood and fibroblasts (two from normal skin and one benign nevus) was analysed for 11p15 CNVs and methylation aberrations by SALSA MS-MLPA Probes (Roche NimbleGen, Madison, WI). Also, DNA from blood was tested for other imprinted diseases using the SALSA MS-MLPA Probes ME032-A1 UPD7-UPD14, and for genomic CNVs by CytoScanHD Array (Thermo Fisher Scientific, Waltham, MA).

**Microsatellite analysis**

To determine CDKN1C haplotypes and allele segregation, and to explore on which parental allele the grandfather’s de novo variant arose, an informative haplotype was set up by simple tandem repeat markers amplified by PCR and size determined. There were two informative markers upstream and four downstream of CDKN1C (online supplemental file). The haplotype analysis was performed on DNA from peripheral blood from the patient, his parents, his mother’s siblings, maternal grandparents and the grandfather’s parents.

**RESULTS**

**Clinical history and findings in the boy**

Clinical findings in the patient are summarised in table 1 and illustrated in figure 2. Omphalocele was discovered by routine ultrasound in week 19 of gestation, anomicentesis was performed, and the pregnancy was complicated by maternal hyperemesis, pre-eclampsia, diabetes from week 29 and later polyhydramnios. He was born after caesarean section in gestation week 34 +2 with birth weight 2160 g (~1.1 SDS), birth length 42 cm (~2.5 SDS) and head circumference 32 cm (0 SDS) (all measurements adjusted for prematurity). Placental weight was on the 90th
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percentile. Apgar score was 3/5/7 after 1, 5 and 10 min, respectively. He had mild hypoglycaemia (1.5 mmol/L) and compromised respiration. The giant omphalocele also contained liver tissue, and he underwent several surgeries before the abdominal wall finally was closed at 3 months of age. He suffered from apnoeic spells and needed CPAP until he was discharged from the hospital 5 months of age. He was exclusively fed by nasogastric tube for the first 5 months, and the tube was removed after 12 months. At the age of 3 years, he had been through more than 20 surgeries, including for patent ductus arteriosus, inguinal hernia and cryptorchidism. He had dysmorphic features as illustrated in figure 2, small intra-abdominal testes, multiple benign epidermal nevi and mild hypotonia. A follow-up at 25 months of age, his length was 0.8 SDS, weight −2.0 SDS and he was now microcephalic (−2.7 SDS). Motor milestones were delayed; he could sit at 15 months and walk without support at 2 years and 10 months of age. At age 3 years, he could speak a few simple words, and at age 5 years he had 3–4 words sentences with articulation problems. He had normal neurological examination, hearing, vision and social development, but marked global delayed yet without formal assessment. He received speech, education and physical therapy. Cerebral MRI was normal, repeated abdominal ultrasounds from the age of 4 months showed small kidneys bilaterally and no organomegaly. At 2.5 years, an ACTH-stimulation test did not reveal adrenal insufficiency, and skeletal X-ray showed long and slender diaphyses, broad metaphyses, a fusion of left fourth and fifth ribs, and a 1.5-year delay in bone maturation. At age 5 years, his height was −1.5 SDS, head circumference centiles unchanged, confirming a postnatal/secondary microcephaly (−2.5 SDS), and he still had a low BMI of 12.9 (−2.8 SDS). He was scheduled to cancer surveillance every sixth month until 7–8 years of age.

Family history
The boy’s phenotypically normal parents are non-consanguineous, the mother was 28 and the father 42 years at his birth. Clinical examination and birth history in the mother were negative for BWS features. Head circumferences were at 99th percentile (mother) and 60th percentile (father). The maternal grandfather was >5 kg at birth, and a large tongue was commented on neonatally, suggesting macroglossia. At examination at age 58 years, he was 181 cm tall with a wide mouth, slightly short halluces, no asymmetry, a normal skeletal X-ray survey and normal cognition.

Diagnostic workup
During pregnancy, a normal male karyotype 46,XY was found in amniotic fluid cells. After birth, clinical suspicion of BWS in the boy warranted MS-MLPA of the BWS region, which was normal in DNA from blood, skin and epidermal nevus. However, sequencing of CDKN1C verified BWS (see next section). Due to borderline short stature, developmental delay and postnatal microcephaly, further analyses were performed with normal results: high-resolution CNV analysis, UPD screening and trio-based WES.

Interpretation of the CDKN1C variant
Sanger sequencing of CDKN1C (NM_000076.2/ENST00000414822.7) revealed a heterozygous delins variant in the sequence encoding the PCNA binding motif affecting the 5’-end of exon 2: c.822_826delinsGAGCTG. The delins caused frameshifting and a premature termination codon (PTC) about 100 nucleotides upstream of the exon 2–3 junction. This change should either lead to nonsense-mediated mRNA decay

Figure 2  Boy at age 3 months (A, C, D), 6 months (B) and 27 months (G, F), with clinical features of both Beckwith-Wiedemann syndrome (ie, omphalocele, midface retrusion, large cheeks, hypertelorism with down-slanted palpebral fissures, infraorbital and ear creases, a fading glabellar capillary malformation, and a long and marked philtrum with a thin upper lip, wide mouth with a high palate and macroglossia) and IMAGe (slender habitus, feeding difficulties, frontal bossing, broad nasal bridge and a wide tip, low set and posteriorly rotated ears, long and slender finger and toes). Radiographs of leg (G) showing long and slender diaphysis and broad metaphysis, with a delayed bone age of fingers (F).
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(NMD) or synthesis of a truncated protein Asp274GlufsTer12, that is, an LoF change that fits with BWS phenotype. However, the delins also introduced two new putative splice junction AG acceptor sites, in addition to the consensus AG site (figure 3, upper panel). Use of alternative acceptor site I will also lead to a frameshift and a new stop codon, but in the very last protein-coding exon, that is, Asp274GlufsTer47, which should escape NMD (figure 3, lower panel). The corresponding protein product contains a modified C terminus including substitutions of four IMAGe-associated amino acids and introduce the known GoF variant Phe276Ser.4 Use of alternative acceptor site II would be in line with the canonical reading frame of the protein's C-terminal part, but also include deletion of three and insertion of two new amino acids in the PCNA-binding motif: 274_276delinsAlaVal. This would modify one IMAGe residue (Asp274Ala), but also introduce a previously reported IMAGe-associated substitution: Phe276Val (figure 3).4 This alternative transcript can be compared with known IMAGe-mutated transcripts, resulting in increased protein stability or affected PCNA binding.

Figure 3 Predicted impact of the delins variant c.822_826delinsGAGCTG on DNA and isoform A. Upper panel shows the CDKN1C intron 2/exon 3 junction with reference amino acids on top and the reference nucleotide sequence (ENST00000414822.7, CDKN1C-202) just below. Intron sequences are written in lower case letters and exon sequences in capital letters. The five deleted nucleotides are in green. The bottom line shows the mutated sequence with the six inserted nucleotides in red, the three adjacent AGs (potential acceptor sites/AS) underlined and the out-of-frame sequence marked in blue. The lower panel shows details of the predicted amino acid sequence and changes introduced by the delins variant in amino acid numbers according to protein isoform A (316aa—NP_000067.1). On top reported IMAGe (black) and Silver-Russel syndrome (SRS)—like variants (magenta) from literature with the corresponding amino acid change, then the reference protein product, followed by the mutated products using the consensus AG (p.Asp274GlufsTer12), the protein product if alternative acceptor site I is used (p.Asp274GlufsTer47) and the protein product if alternative acceptor site II is used (p.Asp274_276delinsAlaVal). Parentheses (…) represent 21 amino acids omitted from the figure.

Figure 4 Overview of the predicted protein products. From top, wild-type isoform A (encoded by transcripts ENST00000414822.7=CDKN1C-202 and ENST00000430149.2=CDKN1C-203), wild-type isoform B (ENST00000440480.6=CDKN1C-204), followed by three predicted CDKN1C protein variants of mutated isoform A depending on the splice acceptor site used; consensus acceptor site and alternative acceptor sites I or II. At the bottom, isoform D (A6NK88, encoded by ENST00000380725.1=CDKN1C-201, with a reading frame different from isoforms A and B) and the mutated isoform D (hybrid isoform D—A/B), consensus acceptor site used. The colours correspond to regions in figures 1 and 3, where intron sequences are written in lowercase letters and exon sequence in capital letters. The five deleted nucleotides are in green. The six inserted nucleotides in the mutated (delins) sequences are shown in red, and the out-of-frame sequence marked in blue. † denotes a predicted transcript that was not found on RNA sequencing.
The predicted out-of-frame C-terminal product from alternative acceptor site 1, p.(Asp274GlufsTer47) (figures 3 and 4), was analysed by BLAST and found to be almost identical to CDKN1C-201, encoding a 131 amino acid isoform we call isoform D (UniProtKB accession A6NK88). Isoform D consists of the common CDI domain with a unique C-terminal D-tail, marked in red in figure 4, and its alternative splicing bypasses the PAPA-repeat region (figure 1). In addition to the PCNA-related amino acid substitution, the delins could also affect isoform D by introducing a frameshift that changed the reading frame back to the canonical reading frame (marked in yellow in figure 4). The predicted hybrid protein had elements from two distinct isoforms, D and A/B, where the D-type splicing causes direct connection of the CDI domain to an altered PCNA-binding motif, and where delins-induced frameshifting causes the C-terminus to be of A/B type and not D type.

Diagnostic analysis of the family

The boy’s asymptomatic mother was also heterozygous for the delins, compatible with a paternally imprinted allele in her, and further family studies including haplotype segregation analysis confirmed that the variant arose de novo on the grandfather’s maternal allele. The mild BWS features in the grandfather indicated mosaicism, which was confirmed by DNA analyses of multiple tissues; only wt was present in fibroblasts from three out of four skin biopsies and a buccal swab, while the delins was found in blood, urine sediment and skin biopsy from the left leg. cDNA Sanger sequencing and qPCR cDNA sequencing confirmed the presence of both wt and (at a lower level) the delins allele in the boy, his mother and grandfather. Gene expression levels appeared generally low. Only in fibroblasts from the boy, the delins seemed to be more abundant than the wild-type transcript. Primers were also designed to cover the use of the two alternative splice acceptor sites illustrated in figure 3, but corresponding isoform-A/B-derivated transcripts were not detected. However, sequencing demonstrated that wild-type isoform D (CDKN1C-201) transcript was present in blood from all family members and controls, and the delins-CDKN1C-201 transcript was present in blood from the boy.

Semi-quantitative real time-PCR (qPCR) verified the presence of both CDKN1C (isoform A/B) and CDKN1C-201 (isoform D) transcripts in blood samples from the patient, parents and grandfather. Also, isoform A/B was found in fibroblasts from all family members. Transcript levels were low compared with the B2M control and considerably lower in fibroblasts than in blood. Expression of delins-CDKN1C was seen in blood from the boy and grandfather, but not the mother, while expression in fibroblast was very low in the boy and undetectable in others. Results from cDNA sequencing and qPCR are not shown.

NGS-based cDNA sequencing

Ultra-deep RNA-seq was done for a more quantitative non-biased analysis of all transcripts, including the delins variant. The results are illustrated in figure 5. In summary, the analysis confirmed that the level of delins was 40%–50% in the boy (higher in fibroblast than in blood), 25% in the mosaic grandfather and 6% in the mother; that activation of alternative acceptor site 1 occurred in the boy’s fibroblasts; and that CDKN1C-201 represented 16%–42% of the total CDKN1C mRNA reads but accounted for the majority of the delins reads. Neither standard nor ultra-deep RNA sequencing could verify the presence of a predicted fifth transcript CDKN1C-206 (ENST000000647251.1) that encodes a putative 175 aa protein product labelled isoform C (CCDS86169.1).

All RNA-seq samples had in total >340 million reads after filtering, and the numbers of CDKN1C tracks spanning splice junctions (SJT) were between 35 and 204 (figure 5A). Visualization of the alignment to NCBI Build 38 is presented with the counting of SJTs (online supplemental file). No delins reads were mapped against Build 38, and we suspected mapping problems. We then created three new FASTA reference files with the delins for accurate mapping (online supplemental file), one based on the consensus splice site and two with alternative acceptor sites I and II. Alignment to one of these putative delins splice sites unmasked the delins-specific tracks, while the other two alignments did not add more data and were therefore not further analysed. Figure 5A shows the result from the mapping of transcripts CDKN1C-202, CDKN1C-203, CDKN1C-204 (combined) and CDKN1C-201 against the reference sequence (green columns), and the distribution was compared with the sum of other CDKN1C transcripts. Likewise, the pink columns show the distribution of delins-mapped reads. The total distribution of delins and CDKN1C-201 is also presented. SJTs presented as arcs in IGV (online supplemental file) showed that the vast majority of expressed transcripts have splicing from 5’-UTRs of different lengths, including CDKN1C-201 and CDKN1C-204, and both use the alternative Met11 amino acid downstream and have four exons. In contrast, CDKN1C-202 and CDKN1C-203 always start at Met1 and have three exons. CDKN1C-203 uses an alternative acceptor splice site five nucleotides downstream in the last exon and can be distinguished from other transcripts. The coding part of CDKN1C-201 continues into exon 4, while this exon is making a 3’-UTR in other transcripts.

DISCUSSION

This family, where the index case has a mixture of diametrically opposed growth-related syndromes due to the same delins variant in CDKN1C, illustrates that the biological complexity of this small gene goes far beyond imprinting. Due to intricate gene splicing with frameshifting, this delins has an LoF effect on some isoforms and GoF effect on another. The latter is unexpected for a five against six nucleotides exchange and is due to the creation of a new splice acceptor site two nucleotides downstream of the canonical AG. Furthermore, we demonstrate that an alternative isoform with a different C-terminal reading frame (that we call isoform D, corresponding to transcript CDKN1C-201) is also affected, which turns the C-terminus of isoform D back to the isoform A/B reading frame. If the hybrid D–A/B isoform transcript is expressed in the brain, this could be the cause of his developmental delay.

The asymptomatic carrier mother expressed the delins at a low level, as expected for a leaky paternally imprinted allele. The maternal grandfather was mosaic for the delins on his maternal allele, in agreement with an intermediate expression level. As a cautious note, the relative levels of the transcripts are not indicative of the degree of imprinting due to variable and poor expression, also affected by presumably inconsistent and non-uniform NMD. Variation in NMD efficiency within mRNA with the same PTC and subpopulations of mRNA escaping decay are reported, and this can contribute to explaining variation between individuals and between different tissues.11–12

Ultra-deep RNA sequencing proved to be superior to conventional RNA analysis, particularly on fibroblast RNA. Such unbiased sequencing verified the use of alternative acceptor site I and enabled us to quantify the expression of different transcripts,
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which again suggested which isoforms were most abundant. RNA sequencing as a first choice in a diagnostic pipeline would have been unsuccessful in this family as the delins could not be mapped with a single track against the reference sequence, but we solved this by mapping against dedicated reference files.

In summary, the boy fulfilled the clinical criteria of BWS—an overgrowth condition—despite being rather small and microcephalic, with a total score of 8, with two cardinal features (omphalocele and macroglossia) and four suggestive features (see table 1).

Figure 5  Result from NGS-based cDNA sequencing. (A) The number of reads spanning exon junctions (splice junction tracks, SJTs) after mapping to the reference sequence (from Genome Build 38, green background) and the delins-containing sequence (manually designed reference sequence, pink background). Reads with delins are shown in red. All reads were manually checked, and only SJTs spanning exons 2–3 are counted. Numbers are derived from figures as in the last part of the online supplemental file 1. Of note, reference sequence mapping did not detect any delins reads, probably because they were filtered out due to the complexity of the delins variant. (B) Sector diagrams illustrating the distributions of documented transcripts from RNA sequencing of RNA derived from blood samples and fibroblast. The percentages are calculated from the numbers in figure 5A. It was not possible to discriminate all CDKN1C transcripts by RNA sequencing, so CDKN1C-202, CDKN1C-203 and CDKN1C-204 are collectively referred to as CDKN1C in this figure, but CDKN1C-201 transcripts (isoform D, UniProt: A6NK88) are identified. Green and blue colours represent wt reads of CDKN1C and CDKN1C-201, respectively, and red and purple colours represent delins reads using the consensus acceptor site in CDKN1C and CDKN1C-201, respectively. A fifth transcript (yellow) with delins CDKN1C using alternative acceptor site I, p.(Asp274GlufsTer47), was only documented in RNA from fibroblasts in the boy. The predicted or hypothetical clinical consequences of each of the four different mutated transcripts are noted in parentheses.

insufficiency could have been masked by continuous intravenous treatment of a leaking omphalocele. Genital abnormalities are described in both IMAGe and BWS, but small testes in IMAGe only. Even though the renal size is not discussed in IMAGe (only fetal renal cortex size), his renal hypoplasia could very well be an IMAGe mirror of renal hyperplasia in BWS. Skeletal abnormalities are important findings in IMAGe, but besides asymmetry less important in SRS and are not features of BWS. Clinically, he did not pass the Netchine-Harbinson SRS score, having 3/6 criteria which is below the 4/6 limit.33 His being small for gestational age, feeding difficulties, mild skeletal abnormalities and small testes are compatible with IMAGe, but this diagnosis is not definite as adrenal hypoplasia and metaphyseal dysplasia were not documented. The clinical findings point to


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dual molecular effects with both LoF (BWS) and GoF (IMAGE) in CDKN1C. The latter can be explained by p.Asp274_276delinsAlaVal encoding transcript from alternative acceptor site II, but we could not confirm expression of this transcript in the two tissues tested. Another hypothetical GoF candidate is the p.Asp274GlufsTer47 encoding transcript from alternative acceptor site I with a verified GoF substitution Phe276Ser.43 This transcript encodes an alternative C-terminal abolishing the PCNA-binding site, but the product might also be non-functional or degraded. Probably NMD is inefficient as a protective mechanism in CDKN1C as the truncated transcript from the consensus splice site, that is, encoding p.Asp274GlufsTer12 which is well within theoretical criteria for NMD, was present in both blood and fibroblasts.34 This mutant product also substitutes IMAGE-related amino acids (figure 3) and might also have GoF effects.

A combination of opposing phenotypes due to a single mutation in the same gene is not reported earlier for CDKN1C, but for GNAS, another imprinted gene, this is not new.35 Other genes where a variant is reported to cause both GoF and LoF effects are RET, FLNA, GDF5 and SMCHD1.36–39 In general, multinoucleotide variants are potentially more damaging than single-nucleotide variants, illustrated by our family.40

The most intriguing finding was that the delins also affected a hitherto not confirmed transcript using a non-canonical reading frame—CDKN1C-201—which represented ~40% of all CDKN1C transcripts in the family (figure 5A). In contrast to wt CDKN1C-201, delins-mutated CDKN1C-201 used the canonical reading frame, resulting in a hybrid D–A/B product that may be the cause of the unusual features in the boy. Alternative non-canonical reading frames are common in prokaryote and mitochondrial genes, but less so in eukaryotic and human genes, and another cell-cycle inhibitor gene, CDKN2A, is perhaps the most known example. This gene encodes at least two different proteins from different reading frames, p16(INK4) and p14(ARF). Both of these have splice variants, p16-gamma and p12, respectively, with tissue-specific expression.41,42 Maybe this kind of regulatory complexity is a peculiarity of cyclin-dependent kinase inhibitors.

Our study identifies CDKN1C-204 (isoform B) as the most abundant transcript, followed by CDKN1C-201 (isoform D). The GTEx portal (gtexportal.org) suggests CDKN1C-201 to be the most abundant isoform, particularly in the brain, followed by the isoforms encoded from CDKN1C-204 and CDKN1C-202, but this could be explained by poor coverage of the PAPA-repeat region not included in the CDKN1C-201, that is, be artefactual. CDKN1C-201 is only reported in humans and some higher primates.

Despite high CDKN1C expression in the embryonic brain, developmental delay and microcephaly are not considered features of either BWS or IMAGE, although brain abnormalities are reported in some patients.43–46 However, behavioural and emotional difficulties and autistic spectrum disorders were found to be more frequent in BWS than controls, especially among patients with IC2 defects.47 Moreover, in a cancer predisposition syndrome clinic, half of the patients with BWS or isolated hemihypertrophy needed special therapy.48 Protein studies in embryonic brains of mice and rats suggested that Cdkn1c regulate newly formed migrating neurons,49 and played a CD1-independent role in the maturation of midbrain dopaminergic neurons.50 Differential spatial and temporal Cdkn1c expression in embryonic brain could be important for neurogenesis and gliogenesis, differentiation and generation of adult neural stem cells.48–50 During cortical development, the differentiation of projection neurons was regulated via Cdkn1c.51 According to a recent study, increased neuronal Cdkn1 expression resulted in abnormal social behaviour, implying that tightly regulated monoallelic expression is beneficial for neurological function.52 Unexpectedly, brain-specific conditional deletion of the imprinted paternal cdk1c allele in mice resulted in microcephaly and thinning of the neocortex, despite low paternal expression (1%–2% of maternal levels) in control brains.53

The delins-mutated hybrid D–A/B transcript is our best explanation for the child’s developmental delay. Brain damage due to neonatal hypoglycaemia or an untreated adrenal crisis was highly unlikely, and his brain MRI was normal, but prematurity and long hospitalisation might be contributing factors. Comparison with other IMAGE patients is difficult as many are not molecularly confirmed and could include phenocopies like MIRAGE syndrome.44 Neither a high-resolution CNV analysis nor a trio-WES revealed alternative explanations for his developmental delay, but this cannot be excluded before further similar cases confirm a neurological association to CDKN1C-201. We searched the literature for other variants that may affect CDKN1C-201 and came across a BWS family with a variant c.821–9C>A with an unknown function.1 Unfortunately, no RNA was available to check the molecular consequence (Brioude, personal communication).

In summary, our findings demonstrate that a single delins variant can give rise to three distinct phenotypes through different molecular pathomechanisms. A hitherto not reported transcript CDKN1C-201, encoding an isoform D (A6NK88) with abundant brain expression, is also affected by the variant. Further functional studies or additional patients are needed to address the role of isoform D in brain development and to verify if changes to this isoform may cause a third face of the CDKN1C spectrum: developmental delay.

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REFERENCES


Genotype-phenotype correlations


