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Mutations in *CDC45*, Encoding an Essential Component of the Pre-Initiation Complex, Cause Meier-Gorlin Syndrome and Craniosynostosis

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Abstract

DNA replication precisely duplicates the genome to ensure stable inheritance of genetic information. Impaired licensing of origins of replication during the G₁ phase of the cell cycle has been implicated in Meier-Gorlin syndrome (MGS), a disorder defined by the triad of short stature, microtia and a/hypoplastic patellae. Biallelic partial loss-of-function mutations in multiple components of the pre-replication complex (preRC; ORC1, ORC4, ORC6, CDT1 or CDC6) as well as *de novo* stabilizing mutations in the licencing inhibitor, GMNN, cause MGS. Here we report the identification of mutations in CDC45 in 15 affected individuals from 12 families with MGS and/or craniosynostosis. CDC45 encodes a component of both the pre-initiation (preIC) and CMG helicase complexes, required respectively for initiation of DNA replication origin firing and ongoing DNA synthesis during S-phase itself, hence is functionally distinct from previously identified MGS-associated genes. The phenotypes of affected individuals range from syndromic coronal craniosynostosis to severe growth restriction, fulfilling diagnostic criteria for Meier-Gorlin syndrome. All mutations identified were biallelic and included synonymous mutations altering splicing of physiological CDC45 transcripts, as well as amino acid substitutions expected to result in partial loss of function. Functionally, mutations reduce levels of full-length transcripts and protein in subject cells, consistent with partial loss of CDC45 function and a predicted limited rate of DNA replication and cell proliferation. Our findings therefore implicate the preIC as an additional protein complex involved in the etiology of MGS, and connect the core cellular machinery of genome replication with growth, chondrogenesis and cranial suture homeostasis.

Introduction

Replication of DNA during eukaryotic cell division is an essential process, which requires a complex apparatus of conserved proteins operating under tight temporal and regulatory control. Although the duplication process itself occurs during the S (synthesis) phase of the cell cycle, the initial components assemble on DNA much earlier, during the late mitotic stages and in G1 phase.

In the first stage, the pre-replication complex (preRC) is formed by the 6-subunit origin recognition complex (ORC) binding to replication origins distributed throughout the genome (Figure 1).¹ ORC recruits CDT1 and CDC6, which leads to the binding of the inactive MCM2-7 helicase as a double hexamer at replication origins.² At the G₁/S transition the preinitiation complex (preIC) proteins assemble in a two-step DDK- and CDK-dependent manner³ and through interaction with the MCM helicase enable binding by the CDC45 and GINS1-4 proteins. This creates the activated CMG helicase, an eleven-subunit complex that possesses essential DNA unwinding activity, allowing polymerases access to DNA and enabling replication to commence.^{4, 5, 6, 7, 8} CDC45 has single-stranded DNA binding activity, facilitating DNA strand displacement at the replication fork.⁹ Hence CDC45 plays a central role both in initiation of DNA replication origin firing (preIC) and ongoing DNA synthesis (CMG helicase), and genetic studies demonstrate that it is essential in both yeast and mice.^{10, 11, 12, 13, 14} Both *in vitro* and *in vivo* data indicate that CDC45 is loaded onto chromatin specifically in the S phase of the cell cycle, after the assembly of the preRC complexes.^{11, 14, 15, 16, 17}

Several Mendelian syndromes have been associated with mutations in components of the DNA replication machinery. Meier-Gorlin syndrome (MGS) (MIM 224690) is characterised by short stature, microtia (small ears) and aplasia or hypoplasia of the patellae.¹⁸ Biallelic mutations in multiple components of the preRC (*ORC1* [MIM 601902],

ORC4 [MIM 603056], *ORC6* [MIM 607213], *CDT1* [MIM 605525] and *CDC6* [MIM 602627]) were identified in individuals with MGS, ^{19, 20, 21} and among them, mutations in these genes account for approximately 70% of cases.²² Recently, *de novo* mutations in three individuals were reported in the CDT1 inhibitor, *GMNN* (MIM 602842), resulting in the omission of a degron domain that stabilises GMNN levels and is consequently predicted to impair licensing in subject cells.^{23, 24}

Here, we provide genetic and functional evidence that mutations in *CDC45* (MIM 603465) cause human disease. We describe 15 individuals with biallelic partial loss of function mutations in *CDC45* and demonstrate a phenotype that extends from syndromic craniosynostosis to classical MGS.

Subjects and Methods

Clinical studies

The clinical studies were approved by Oxfordshire Research Ethics Committee B (reference C02.143), London Riverside Research Ethics Committee (reference 09/H0706/20), Scottish Multicentre Research Ethics Committee (04:MRE00/19) and the Medisch Ethische Toetsingscommissie of the Leiden University Medical Center (LUMC) (P14-029). Subjects were enrolled into the craniosynostosis cohort based on referral from a craniofacial unit or clinical genetics department, with craniosynostosis proven on either plain radiographs or computed tomography (CT) of the skull. Individuals in the MGS cohort were clinically diagnosed by the referring clinician; in those from whom clinical data were available, all demonstrated hypoplastic or absent kneecaps, small or simple ears, and facies typical of MGS (small mouth and full lips). All participants gave informed consent, and separate permission was obtained for publication of clinical photographs.

Whole genome and exome sequencing

Subjects P1 (Family 1, II-3 in Figure 2A) and P2 (Family 2, II-1 in Figure 2A), from a craniosynostosis cohort, were subjected to whole genome sequencing and exome sequencing, respectively. Exome sequencing of subject P4 (Family 4, II-1 in Figure 2A) was undertaken by Oxford Gene Technology, as part of a study of subjects with primordial dwarfism (characterised by *in utero* growth retardation, and postnatal head circumference and height more than 4 standard deviations below the age- and gender-matched population mean). Exome sequencing of Family 12 was undertaken as part of a diagnostic approach to mutation identification. All details of sequencing library preparations and platforms, as well as software tools for mapping, alignment and variant calling and prioritization are presented in Table S1.

Genomic analysis of CDC45

To investigate further the significance of the *CDC45* variants, primers were designed for amplification of genomic DNA (GenBank: NT_011520.13) and cDNA (GenBank: NM_003504.4; see Supplemental Data, Note on accession numbers for *CDC45*). We used dideoxy-sequencing to confirm the identity of *CDC45* variants identified during whole genome and exome sequencing, and analyzed their segregation in parents and unaffected siblings. Primer sequences and experimental conditions used for PCR amplification and sequencing are provided in Table S2. To analyze a larger number of individuals for *CDC45* mutations, we used Fluidigm/Ion Torrent resequencing or dideoxy-sequencing, respectively, to screen *CDC45* in DNA panels from subjects with craniosynostosis (467 samples) or MGS (34 samples), in whom mutations in known causative genes had not previously been

identified (Table S2). Multiplex ligation-dependent probe amplification (MLPA) was used to confirm the intragenic deletion in Family 12 (Table S2).

Analysis of CDC45 splicing

Total RNA was extracted from lymphoblastoid cell lines using Trizol reagent (Invitrogen) and the RNeasy kit (Qiagen) according to the manufacturer's instructions. DNA was removed by treatment with DNase I (Qiagen). cDNA was generated using random oligomer primers and superscript II (Thermo Fisher) or the RevertAid First Strand Synthesis kit (Fermentas). Primers used for RT-PCR and qRT-PCR are listed in Table S3. cDNA levels were quantified by qRT-PCR using Brilliant II SYBR green qPCR master mix with passive ROX reference dye (Agilent Technologies) on the ABI Prism HT7900 Sequence Detection System. The relative expression of target genes to a control was calculated using the comparative CT method ($2^{-\Delta\Delta CT}$) method.

Cell lines, chemicals, plasmids and transfections

Lymphoblastoid cell lines were cultured in RPMI-1640 supplemented with 15% fetal bovine serum (FBS) and 1% penicillin and streptomycin. Amniocytes were cultured in Amnio-MAX C-100 Complete Medium (Gibco). All reagents were purchased from Life Technologies unless otherwise stated.

Antibodies and Western blotting

Whole cell lysate was prepared using urea buffer (9 M urea, 50 mM Tris HCl, pH 7.3, 150 mM β -mercaptoethanol). Primary antibodies used were mouse anti-human CDC45 (sc55569, Santa Cruz Biotech) at 1:1000 and mouse anti-human TUBULIN (B512, Sigma) at 1:10000.

Protein alignments and structure prediction

Clustal Omega²⁵ was used to generate multi-sequence alignments of orthologous CDC45 proteins. Accession numbers are: *Homo sapiens* NP_003495.1, *Pan troglodytes* XP_009436063.1, *Canis lupus* XP_543547.3, *Mus musculus* NP_033992.2, *Gallus gallus* XP_415070.2, *Danio rerio* NP_998551.1, *Drosophila melanogaster* NP_569880.1, *Anopheles gambiae* XP_320573.1, *Caenorhabditis elegans* NP_497756.2, *Saccharomyces cerevisiae* NP_013204.1, *Schizosaccharomyces pombe* NP_594693.1. The structural model of human CDC45 was modelled on the 3.7Å cryo-electron microscopy structure of *S. cerevisiae* Cdc45.²⁶ It was created using Modeller²⁷ and is presented using Pymol (Schrodinger).

Results

Exome and Genome Sequencing and Targeted Resequencing Identify Biallelic Variants in *CDC45*

As part of a large multi-disease study,²⁸ clinical whole genome sequencing of a trio comprising a male individual (P1, Family 1:II-3 in Figure 2A) with syndromic unilateral coronal craniosynostosis and his unaffected parents, initially identified several candidate variants for further scrutiny. These were a single *de novo* missense mutation (*FLRT1* [MIM: 604806] c.137C>T, p.T46M [NM_013280.4]), two very rare (absent in dbSNP135, ESP or 1000 Genomes) hemizygous nonsynonymous variants (*OPHN1* [MIM 300127] c.1559C>T, p.Thr520IIe [NM_002547.2] and *GEMIN8* [MIM 300962] c.500G>A, p.Arg167His [NM_017856.2]) and two very rare compound heterozygous genotypes: *CDC45* c.[318C>T; c.677A>G], p.[Val106=; Asp226Gly], and *MYO10* [MIM 601481] c.[1491G>A; 5459A>C], p.[Glu497=; His1820Pro] (NM_012334.2). *FLRT1* was initially prioritized for further analysis, but resequencing this gene in a panel of 230 individuals with craniosynostosis revealed no further mutations. During concurrent exome sequencing of female subject P2 (Family 2:II-1 in Figure 2A) with syndromic bilateral coronal craniosynostosis, variant analysis under a recessive model of inheritance also identified two very rare predicted missense changes in *CDC45* (c.226A>C, p.Asn76His and c.469C>T, p.Arg157Cys); dideoxy-sequencing of parental samples showed that the *CDC45* variants were present in a compound heterozygous (biallelic) state. Given the identification of two individuals with unilateral or bilateral coronal or craniosynostosis, each of whom had two rare biallelic variants in *CDC45* (Table 1), resequencing of *CDC45* was undertaken in a further 467 subjects with craniosynostosis. This revealed one further individual (P3, Family 3:II-1 in Figure 2A) with two rare *CDC45* variants (Table 1), which analysis of parental samples showed were also biallelic. Consistent with a causal role in the disease state, three unaffected siblings of P1 and P2 inherited different combinations of *CDC45* alleles (Figure 2A).

In addition to craniosynostosis, clinical features variably present in individuals P1-P3 included mild short stature, microcephaly and ear anomalies (Figure 2B, Table 2). In view of the previous association of replication-associated genes with growth restriction,^{19, 20, 21, 22} we interrogated exome sequencing data from a cohort of individuals with primordial dwarfism (n = 52) for variants in *CDC45*. One individual (P4, Family 4:II-1 in Figure 2A), diagnosed with MGS, was found to harbor two rare *CDC45* variants (Table 1).

Given the presentation of subject P4 with MGS, we then undertook dideoxy-sequencing of 34 further subjects with MGS, in whom causative mutations had not previously been found in preRC components. This identified seven further families (Family 5- Family 11) (Table 1), in which nine individuals have biallelic mutations in *CDC45*. All variants had either a very low allele frequency in control populations with no homozygotes, appropriate for a rare recessively inherited Mendelian disorder, or were absent from the ExAC dataset. The

autosomal recessive inheritance was confirmed by appropriate segregation in all parents where available (Figure 2A, Table 1).

Concurrently, exome sequencing was undertaken in a nonconsanguineous Dutch family (Family 12) in which two consecutive fetuses displayed craniosynostosis and additional anomalies *in utero*. The recurrence within the family suggested an autosomal recessive inheritance, however no candidate disease genes were identified following bioinformatic analysis and filtering of the exome data. Given the previous findings, *CDC45* and other autosomal recessive craniosynostosis disorder associated genes were manually re-examined; this identified a single paternally transmitted missense variant in *CDC45*, c.893C>T encoding p.Ala298Val. Visual analysis of the sequencing reads using Integrated Genomics Viewer²⁹ suggested lower coverage of exon 5 in both the two affected samples and the maternal sample compared with the paternal sequencing (Figure S1A). Analysis using MLPA probes confirmed a deletion encompassing only exon 5 of *CDC45* in these three samples (Figure S1B). This would be predicted to delete a highly conserved portion of the protein, but translation would remain in-frame and therefore transcripts from this allele would not be predicted to undergo nonsense-mediated decay. This was confirmed by analysis of cDNA derived from maternal blood (Figure S1C).

The human CDC45 mutational spectrum indicates partial loss of protein function

CDC45 is an essential component in the preIC that establishes active replication at licensed origins (Figure 1). The majority of variants identified in *CDC45* lead to amino acid substitutions (Figure 3A), which bioinformatic analysis predict to be damaging; we examined conservation of these residues as a measure of deleteriousness, and found all are conserved in vertebrates, with residues at several sites conserved through to yeasts (Figure 3B).

One conserved substitution, c.203A>G, p.Gln68Arg, occurred at the penultimate nucleotide in exon 3 suggesting that it might also impact on splicing in addition to any protein-level consequence (see next section). The three subjects (P4, P5, P8) with a truncating/null mutation (nonsense, frameshift deletion, or initiation codon mutation) on one allele all segregate an identical missense variant, c.1388C>T, p.Pro463Leu present on the *trans* allele; given the South Asian ancestry in all three subjects (ExAC South Asian allele frequency 0.000062), the presence of this variant is compatible with a founder effect.

All individuals had at least one allele compatible with some residual function; none were biallelic for null mutations, consistent with the early embryonic lethality reported in a *Cdc45* knockout mouse (*Cdc45* homozygotes die before E7.5, due to impaired proliferation of the inner cell mass¹³). Individuals who were compound heterozygotes for a loss-of-function and a substitution allele (p.Pro463Leu or p.Ala298Val) were particularly severely affected.

Structural consequences of CDC45 mutations

Missense mutations did not cluster within a specific domain of the protein (Figure 3A). However, a homology model derived from the recent high resolution cryo-electron microscopy study of the yeast CMG helicase complex³⁰ demonstrated that a number of substituted residues are located within the core of the CDC45 protein (Figure 4A, B). This core region corresponds to the known catalytic site of the bacterial RecJ orthologs of CDC45 and its DHH phosphoesterase homologs.^{31, 32} Such substitutions would therefore be likely to perturb protein stability. In particular, the recurrent p.Pro463Leu substitution replaces a conserved proline residue with a likely essential role in N-capping of an alpha-helix (Figure 4C). Likewise, within the protein core the p.Asn76His substitution is expected to introduce a positive charge within, and thus disrupt, the predicted divalent cation coordination sphere

present in many members of the DHH phosphoesterase superfamily (Figure 4D).^{33, 34, 35} Predictions of functional consequences for point mutations at other sites in the homology model were not possible, aside from the p.Pro321Thr substitution that is expected to disrupt the interaction of CDC45 with MCM2 (Figure 4E).

Hypomorphic *CDC45* Mutations act by Disrupting Transcript Splicing and Reducing Protein Levels

Five of the alleles (in Family 1, 3, 9 and 10) were hypothesized to affect splicing (Table S4-, S5) and this was confirmed in the three samples available for extraction of RNA, in which we analyzed the effects of putative splicing mutations on cDNA.

In two families (Family 1, 9), the only putative second disease variants present after filtering to remove common variants were synonymous, which generally are less likely to be pathogenic. These two variants (c.318C>T and c.333C>T) lie close together in exon 4 which is skipped for transcripts in which alternative splicing from exon 3 directly to exon 5 generates a naturally occurring shorter isoform.³⁶ This evolutionarily conserved instance of exon skipping is predicted to generate an in-frame protein product that lacks part of the highly conserved DHH phosphoesterase domain.³² RNA derived from these subjects (P1: c.318C>T, P9-1: c.333C>T) exhibited an increased ratio of exon 3/5 transcript to exon 3/4/5 cDNA product compared to controls (Figure 5). *In silico* tools (Human Splice Finder and Sroogle)^{37, 38} predict the c.318C>T variant to disrupt exon splice enhancer sites, and the c.333C>T variant to enhance potential cryptic exonic splice silencer sites, both mechanisms by which these variants may promote the skipping of exon 4 (Table S5). The apparently nonsynonymous variant c.203A>G (mutation present in P9-1 and P9-2), which lies at the penultimate nucleotide of exon 3, was also shown to cause complete skipping of exon 3 in the cell line from P9-1 (Figure 5Bi).

We next sought to investigate the consequence of these mutations on the CDC45 protein by assessing protein levels in cells from five subjects by immunoblotting for endogenous CDC45. In the three cell lines available from affected individuals (lymphoblastoid cell lines derived from P1, P2 and P9-1) and in primary amniocytes derived from P12-1 and P12-2, CDC45 levels were significantly lower than in multiple control cells or cell lines (P < 0.0001). Moreover, CDC45 levels were reduced below 50% in all cases, indicating a contribution from both alleles to reduced protein levels. This suggests that the missense mutations present in these cells or cell lines (P1, p.Asp226Gly; P2, p.Asn76His and p.Arg157Cys; P12, p.Ala298Val) each result in destabilization of CDC45 protein.

Taken together with the nature of the mutations identified, the key demonstration of decreased cellular levels in several affected individuals confirm that variants identified were pathogenic and the mutations would most likely result in destabilization of CDC45 protein.

Spectrum of Phenotypes associated with *CDC45* Mutations and Correlation with Genotype

In total we identified 15 subjects from 12 families with variants in *CDC45* (Table 1, Figure 2A). Individuals with biallelic *CDC45* mutations exhibit substantial variation in severity, in part explained by ascertainment from two distinct cohorts of affected subjects (craniosynostosis for P1-P3, P12 and MGS for P4-P11). Nevertheless they manifest a recognizable phenotypic spectrum with overlapping clinical features (Table 2, Figure 2B-D). For example, although subjects P4-P11 all presented with the classical triad of MGS features (short stature from -1.3 to -7.7 SD, microtia and absent or hypoplastic patellae), craniosynostosis was also frequent, in contrast to MGS associated with mutations in preRC components. The severity of craniosynostosis varied widely from unilateral or bilateral coronal synostosis to multiple suture involvement (Figure 2C). The discordance for

craniosynostosis in two siblings with identical mutations (P9-1 and P9-2), indicates that this is not a fully penetrant phenotype. Conversely, individuals with a primary presentation of craniosynostosis also exhibited mild MGS features, such as hypoplastic ears (P1, P3, P12-1), mild short stature, and a similar facial gestalt including a small mouth (Figure 2B). The MGS feature of patellar hypoplasia was however not observed in any of the three individuals with craniosynostosis who presented postnatally (P1-P3). Thin eyebrows were present in subjects presenting with either craniosynostosis or MGS, a feature that has not been previously highlighted in these or related developmental disorders. Additionally anal abnormalities (imperforate anus or anterior placement) were present in seven subjects drawn from both cohorts. Growth failure and microcephaly and height reduction were progressive throughout childhood. Carrier parents and heterozygous siblings were phenotypically normal, a point of particular note since *CDC45* is located on 22q11.21, within the interval deleted in the 22q11.2 deletion syndromes.³⁹

Discussion

Here we identify multiple mutations in *CDC45*, implicating it is involved in the pathogenesis of both MGS and craniosynostosis. Previous work had established MGS as a disorder of replication licensing, with biallelic mutations in *ORC1*, *ORC4*, *ORC6* and accessory proteins encoded by *CDC6* and *CDT1*. Furthermore, a recent report has identified stabilizing mutations in *GMNN*, which likely act to inhibit CDT1 and consequently reduce licensing in G_{1} .²³ No mutations in any of the MCM subunits have been discovered in individuals with MGS, previously suggesting that MGS was limited to the upstream preRC components; indeed, a founder *MCM4* mutation has been described causing a distinct entity - a genome instability syndrome of short stature, adrenal insufficiency and primary immunodeficiency.^{40,}

^{41, 42} The identification of *CDC45* is therefore surprising, given the well-established function of the CDC45 protein in replication activation and fork progression (Figure 1). Associating *CDC45* mutations with MGS therefore expands understanding of the pathogenic basis of MGS, and given the number of individuals reported here, establishes *CDC45* as a common cause of MGS syndrome, alongside *ORC1* and *CDT1*. These mutations in individuals with MGS also raise the possibility that mutations in genes coding for other proteins involved in DNA replication may be implicated in MGS and/or craniosynostosis. ^{17, 43, 44, 45}

Following the initial identification of MGS mutations, several studies have proposed disease mechanisms distinct from the canonical role of the preRC in replication, such as dysregulated cilia formation or centrosome duplication.^{46, 47} Our identification of MGS mutations in distinct replication machinery, with similar growth and clinical characteristics seen in the preRC MGS individuals (including those with *GMNN* mutations), strongly favors a replication-based etiology for MGS. Notably, biallelic mutations in the helicase *RECQL4* (MIM 603780, also required for preIC formation and replication initiation^{44, 45}), can cause a phenotype that overlaps with MGS; RAPADILINO syndrome (MIM 266280) is characterised by growth retardation and patellar agenesis, although diarrhoea, palatal anomalies and radial ray defects are also evident.⁴⁵ Since CDC45 has been shown to bind several additional proteins in the preIC, for example TOPBP1⁴⁸ and MCM10,⁴⁹ mutations of these other components may in future be implicated in MGS or disorders exhibiting overlapping features.

A distinctive feature of MGS caused by *CDC45* mutations is the frequent association with craniosynostosis. Only two previous MGS subjects have been reported with craniosynostosis, both with *ORC1* mutations (P1 and P3 in Bicknell et al., 2011)¹⁹, therefore it is likely that the presence of craniosynostosis in MGS subjects is a strong predictor for *CDC45* mutations. Craniosynostosis is also a feature of Baller-Gerold syndrome (MIM

218600), another *RECQL4* disorder,⁴⁵ further implicating replication initiation in the etiology of premature cranial suture closure.

Disruption of the canonical function of CDC45 in replication initiation is consistent with the previously proposed models to explain the growth phenotype, which suggested that impaired replication in affected individuals disrupts cell division during periods of rapid proliferation in development, ultimately impairing organism growth and leading to a 'hypocellular' dwarfism.^{20, 50} Partial loss of function mutations in *CDC45*, like those in preRC components, further indicate that strict regulation of replication is particularly required for the development of specific cartilaginous structures (ear, patella, trachea or bronchial tree),⁵¹ but the exact mechanism remains elusive and future studies with model organisms will be required. Similarly, the frequent occurrence of craniosynostosis in this cohort, along with the established premature suture fusion seen in subjects with *RECQL4* mutations, highlights the importance of normal replication initiation and progression to maintain correct proliferation-differentiation balance in the cranial sutures.⁵²

In summary, we present genetic and functional evidence that mutations in *CDC45* cause Meier-Gorlin syndrome with craniosynostosis. The identification of these mutations in a DNA replication protein functioning downstream to the previously identified MGS genes reinforces the premise that the growth and cartilaginous phenotypes of MGS, as well as the premature cranial suture fusion seen here, are due to replication dysfunction. This identification also suggests new candidate genes in which mutations might underlie this disorder or component phenotypes.

Supplemental Data

Supplemental Data contains Supplementary Notes, Tables S1-S5 and Figure S1.

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Web Resources

The URLs for data presented herein are as follows: 1000 Genomes, http://browser.1000genomes.org dbSNP, http://www.ncbi.nlm.nih.gov/projects/SNP/ ExAC Browser, http://exac.broadinstitute.org/ NHLBI Exome Sequencing Project (ESP) Exome Variant Server, http://evs.gs.washington.edu/EVS/

OMIM, http://www.omim.org/

Pymol, http://www.pymol.org

RefSeq, http://www.ncbi.nlm.nih.gov/RefSeq

GoNL, http://www.nlgenome.nl/

References

- 1. Bell, S.P., and Stillman, B. (1992). ATP-dependent recognition of eukaryotic origins of DNA replication by a multiprotein complex. Nature *357*, 128-134.
- 2. Tanaka, S., and Diffley, J.F. (2002). Interdependent nuclear accumulation of budding yeast Cdt1 and Mcm2-7 during G1 phase. Nat. Cell. Biol. *4*, 198-207.
- 3. Tanaka, S., and Araki, H. (2013). Helicase activation and establishment of replication forks at chromosomal origins of replication. Cold Spring Harb. Perspect. Biol. *5*, a010371.
- 4. Costa, A., Renault, L., Swuec, P., Petojevic, T., Pesavento, J.J., Ilves, I., MacLellan-Gibson, K., Fleck, R.A., Botchan, M.R., and Berger, J.M. (2014). DNA binding polarity, dimerization, and ATPase ring remodeling in the CMG helicase of the eukaryotic replisome. Elife *3*, e03273.
- 5. Ilves, I., Petojevic, T., Pesavento, J.J., and Botchan, M.R. (2010). Activation of the MCM2-7 helicase by association with Cdc45 and GINS proteins. Mol. Cell *37*, 247-258.
- 6. Li, N., Zhai, Y., Zhang, Y., Li, W., Yang, M., Lei, J., Tye, B.K., and Gao, N. (2015). Structure of the eukaryotic MCM complex at 3.8 A. Nature *524*, 186-191.
- Moyer, S.E., Lewis, P.W., and Botchan, M.R. (2006). Isolation of the Cdc45/Mcm2-7/GINS (CMG) complex, a candidate for the eukaryotic DNA replication fork helicase. Proc. Natl. Acad. Sci. U S A 103, 10236-10241.
- Pacek, M., Tutter, A.V., Kubota, Y., Takisawa, H., and Walter, J.C. (2006). Localization of MCM2-7, Cdc45, and GINS to the site of DNA unwinding during eukaryotic DNA replication. Mol. Cell 21, 581-587.
- Szambowska, A., Tessmer, I., Kursula, P., Usskilat, C., Prus, P., Pospiech, H., and Grosse, F. (2014). DNA binding properties of human Cdc45 suggest a function as molecular wedge for DNA unwinding. Nucleic Acids Res. 42, 2308-2319.
- 10. Hopwood, B., and Dalton, S. (1996). Cdc45p assembles into a complex with Cdc46p/Mcm5p, is required for minichromosome maintenance, and is essential for chromosomal DNA replication. Proc. Natl. Acad. Sci. U S A *93*, 12309-12314.
- 11. Mimura, S., and Takisawa, H. (1998). Xenopus Cdc45-dependent loading of DNA polymerase alpha onto chromatin under the control of S-phase Cdk. EMBO J. *17*, 5699-5707.
- 12. Miyake, S., and Yamashita, S. (1998). Identification of sna41 gene, which is the suppressor of nda4 mutation and is involved in DNA replication in Schizosaccharomyces pombe. Genes Cells *3*, 157-166.
- 13. Yoshida, K., Kuo, F., George, E.L., Sharpe, A.H., and Dutta, A. (2001). Requirement of CDC45 for postimplantation mouse development. Mol. Cell. Biol. *21*, 4598-4603.
- Zou, L., Mitchell, J., and Stillman, B. (1997). CDC45, a novel yeast gene that functions with the origin recognition complex and Mcm proteins in initiation of DNA replication. Mol. Cell Biol. 17, 553-563.

- 15. Heller, R.C., Kang, S., Lam, W.M., Chen, S., Chan, C.S., and Bell, S.P. (2011). Eukaryotic origindependent DNA replication in vitro reveals sequential action of DDK and S-CDK kinases. Cell *146*, 80-91.
- 16. On, K.F., Beuron, F., Frith, D., Snijders, A.P., Morris, E.P., and Diffley, J.F. (2014). Prereplicative complexes assembled in vitro support origin-dependent and independent DNA replication. EMBO J. *33*, 605-620.
- 17. Yeeles, J.T., Deegan, T.D., Janska, A., Early, A., and Diffley, J.F. (2015). Regulated eukaryotic DNA replication origin firing with purified proteins. Nature *519*, 431-435.
- Bongers, E.M., Opitz, J.M., Fryer, A., Sarda, P., Hennekam, R.C., Hall, B.D., Superneau, D.W., Harbison, M., Poss, A., van Bokhoven, H., et al. (2001). Meier-Gorlin syndrome: report of eight additional cases and review. Am. J. Med. Genet. *102*, 115-124.
- 19. Bicknell, L.S., Bongers, E.M., Leitch, A., Brown, S., Schoots, J., Harley, M.E., Aftimos, S., Al-Aama, J.Y., Bober, M., Brown, P.A., et al. (2011). Mutations in the pre-replication complex cause Meier-Gorlin syndrome. Nat. Genet. *43*, 356-359.
- Bicknell, L.S., Walker, S., Klingseisen, A., Stiff, T., Leitch, A., Kerzendorfer, C., Martin, C.A., Yeyati, P., Al Sanna, N., Bober, M., et al. (2011). Mutations in ORC1, encoding the largest subunit of the origin recognition complex, cause microcephalic primordial dwarfism resembling Meier-Gorlin syndrome. Nat. Genet. *43*, 350-355.
- Guernsey, D.L., Matsuoka, M., Jiang, H., Evans, S., Macgillivray, C., Nightingale, M., Perry, S., Ferguson, M., LeBlanc, M., Paquette, J., et al. (2011). Mutations in origin recognition complex gene ORC4 cause Meier-Gorlin syndrome. Nat. Genet. 43, 360-364.
- 22. de Munnik, S.A., Hoefsloot, E.H., Roukema, J., Schoots, J., Knoers, N.V., Brunner, H.G., Jackson, A.P., and Bongers, E.M. (2015). Meier-Gorlin syndrome. Orphanet J. Rare Dis. *10*, 114.
- 23. Burrage, L.C., Charng, W.L., Eldomery, M.K., Willer, J.R., Davis, E.E., Lugtenberg, D., Zhu, W., Leduc, M.S., Akdemir, Z.C., Azamian, M., et al. (2015). De novo GMNN mutations cause autosomal-dominant primordial dwarfism associated with Meier-Gorlin syndrome. Am. J. Hum. Genet. *97*, 904-913.
- 24. Yoshida, K., Oyaizu, N., Dutta, A., and Inoue, I. (2004). The destruction box of human Geminin is critical for proliferation and tumor growth in human colon cancer cells. Oncogene *23*, 58-70.
- Sievers, F., Wilm, A., Dineen, D., Gibson, T.J., Karplus, K., Li, W., Lopez, R., McWilliam, H., Remmert, M., Soding, J., et al. (2011). Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. Mol. Syst. Biol. 7, 539.
- 26. Yuan, Z., Bai, L., Sun, J., Georgescu, R., Liu, J., O'Donnell, M.E., and Li, H. (2016). Structure of the eukaryotic replicative CMG helicase suggests a pumpjack motion for translocation. Nat. Struct. Mol. Biol. *23*, 217-224.
- 27. Sali, A., and Blundell, T.L. (1993). Comparative protein modelling by satisfaction of spatial restraints. J. Mol. Biol. *234*, 779-815.
- Taylor, J.C., Martin, H.C., Lise, S., Broxholme, J., Cazier, J.B., Rimmer, A., Kanapin, A., Lunter, G., Fiddy, S., Allan, C., et al. (2015). Factors influencing success of clinical genome sequencing across a broad spectrum of disorders. Nat. Genet. 47, 717-726.
- 29. Robinson, J.T., Thorvaldsdottir, H., Winckler, W., Guttman, M., Lander, E.S., Getz, G., and Mesirov, J.P. (2011). Integrative genomics viewer. Nat. Biotechnol. *29*, 24-26.
- Abid Ali, F., Renault, L., Gannon, J., Gahlon, H.L., Kotecha, A., Zhou, J.C., Rueda, D., and Costa, A. (2016). Cryo-EM structures of the eukaryotic replicative helicase bound to a translocation substrate. Nat. Commun. 7, 10708.
- Krastanova, I., Sannino, V., Amenitsch, H., Gileadi, O., Pisani, F.M., and Onesti, S. (2012). Structural and functional insights into the DNA replication factor Cdc45 reveal an evolutionary relationship to the DHH family of phosphoesterases. J. Biol. Chem. 287, 4121-4128.
- 32. Sanchez-Pulido, L., and Ponting, C.P. (2011). Cdc45: the missing RecJ ortholog in eukaryotes? Bioinformatics *27*, 1885-1888.

- 33. He, Q., Wang, F., Liu, S., Zhu, D., Cong, H., Gao, F., Li, B., Wang, H., Lin, Z., Liao, J., et al. (2016). Structural and Biochemical Insight into the Mechanism of Rv2837c from Mycobacterium tuberculosis as a c-di-NMP Phosphodiesterase. J. Biol. Chem. 291, 3668-3681.
- Wakamatsu, T., Kim, K., Uemura, Y., Nakagawa, N., Kuramitsu, S., and Masui, R. (2011). Role of RecJ-like protein with 5'-3' exonuclease activity in oligo(deoxy)nucleotide degradation. J. Biol. Chem. 286, 2807-2816.
- 35. Yamagata, A., Kakuta, Y., Masui, R., and Fukuyama, K. (2002). The crystal structure of exonuclease RecJ bound to Mn2+ ion suggests how its characteristic motifs are involved in exonuclease activity. Proc. Natl. Acad. Sci. U S A *99*, 5908-5912.
- 36. Kukimoto, I., Igaki, H., and Kanda, T. (1999). Human CDC45 protein binds to minichromosome maintenance 7 protein and the p70 subunit of DNA polymerase alpha. Eur. J. Biochem. *265*, 936-943.
- Desmet, F.O., Hamroun, D., Lalande, M., Collod-Beroud, G., Claustres, M., and Beroud, C. (2009). Human Splicing Finder: an online bioinformatics tool to predict splicing signals. Nucleic Acids Res. 37, e67.
- 38. Schwartz, S., Hall, E., and Ast, G. (2009). SROOGLE: webserver for integrative, user-friendly visualization of splicing signals. Nucleic. Acids. Res. *37*, W189-192.
- Delio, M., Guo, T., McDonald-McGinn, D.M., Zackai, E., Herman, S., Kaminetzky, M., Higgins, A.M., Coleman, K., Chow, C., Jalbrzikowski, M., et al. (2013). Enhanced maternal origin of the 22q11.2 deletion in velocardiofacial and DiGeorge syndromes. Am. J. Hum. Genet. *92*, 439-447.
- 40. Casey, J.P., Nobbs, M., McGettigan, P., Lynch, S., and Ennis, S. (2012). Recessive mutations in MCM4/PRKDC cause a novel syndrome involving a primary immunodeficiency and a disorder of DNA repair. J. Med. Genet. *49*, 242-245.
- Gineau, L., Cognet, C., Kara, N., Lach, F.P., Dunne, J., Veturi, U., Picard, C., Trouillet, C., Eidenschenk, C., Aoufouchi, S., et al. (2012). Partial MCM4 deficiency in patients with growth retardation, adrenal insufficiency, and natural killer cell deficiency. J. Clin. Invest. *122*, 821-832.
- 42. Hughes, C.R., Guasti, L., Meimaridou, E., Chuang, C.H., Schimenti, J.C., King, P.J., Costigan, C., Clark, A.J., and Metherell, L.A. (2012). MCM4 mutation causes adrenal failure, short stature, and natural killer cell deficiency in humans. J. Clin. Invest. *122*, 814-820.
- 43. Im, J.S., Ki, S.H., Farina, A., Jung, D.S., Hurwitz, J., and Lee, J.K. (2009). Assembly of the Cdc45-Mcm2-7-GINS complex in human cells requires the Ctf4/And-1, RecQL4, and Mcm10 proteins. Proc. Natl. Acad. Sci. U S A *106*, 15628-15632.
- 44. Sangrithi, M.N., Bernal, J.A., Madine, M., Philpott, A., Lee, J., Dunphy, W.G., and Venkitaraman,
 A.R. (2005). Initiation of DNA replication requires the RECQL4 protein mutated in Rothmund-Thomson syndrome. Cell *121*, 887-898.
- 45. Siitonen, H.A., Sotkasiira, J., Biervliet, M., Benmansour, A., Capri, Y., Cormier-Daire, V., Crandall, B., Hannula-Jouppi, K., Hennekam, R., Herzog, D., et al. (2009). The mutation spectrum in RECQL4 diseases. Eur. J. Hum. Genet. *17*, 151-158.
- 46. Hossain, M., and Stillman, B. (2012). Meier-Gorlin syndrome mutations disrupt an Orc1 CDK inhibitory domain and cause centrosome reduplication. Genes Dev. *26*, 1797-1810.
- 47. Stiff, T., Alagoz, M., Alcantara, D., Outwin, E., Brunner, H.G., Bongers, E.M., O'Driscoll, M., and Jeggo, P.A. (2013). Deficiency in origin licensing proteins impairs cilia formation: implications for the aetiology of Meier-Gorlin syndrome. PLoS Genet. 9, e1003360.
- 48. Schmidt, U., Wollmann, Y., Franke, C., Grosse, F., Saluz, H.P., and Hanel, F. (2008).
 Characterization of the interaction between the human DNA topoisomerase Ilbeta-binding protein 1 (TopBP1) and the cell division cycle 45 (Cdc45) protein. Biochem J. 409, 169-177.
- Di Perna, R., Aria, V., De Falco, M., Sannino, V., Okorokov, A.L., Pisani, F.M., and De Felice, M. (2013). The physical interaction of Mcm10 with Cdc45 modulates their DNA-binding properties. Biochem J. 454, 333-343.

- 50. Klingseisen, A., and Jackson, A.P. (2011). Mechanisms and pathways of growth failure in primordial dwarfism. Genes Dev. 25, 2011-2024.
- 51. de Munnik, S.A., Bicknell, L.S., Aftimos, S., Al-Aama, J.Y., van Bever, Y., Bober, M.B., Clayton-Smith, J., Edrees, A.Y., Feingold, M., Fryer, A., et al. (2012). Meier-Gorlin syndrome genotype-phenotype studies: 35 individuals with pre-replication complex gene mutations and 10 without molecular diagnosis. Eur. J. Hum. Genet. 20, 598-606.
- 52. Twigg, S.R.F., and Wilkie, A.O.M. (2015). A genetic-pathophysiological framework for craniosynostosis. Am. J. Hum. Genet. *97*, 359-377.
- 53. Newell, N.E. (2015). Mapping side chain interactions at protein helix termini. BMC Bioinformatics *16*, 231.
- 54. Berti, M., and Vindigni, A. (2016). Replication stress: getting back on track. Nat. Struct. Mol. Biol. 23, 103-109.
- 55. Aravind, L., and Koonin, E.V. (1998). The HD domain defines a new superfamily of metaldependent phosphohydrolases. Trends Biochem. Sci. 23, 469-472.
- 56. Cole, T.J., Freeman, J.V., and Preece, M.A. (1998). British 1990 growth reference centiles for weight, height, body mass index and head circumference fitted by maximum penalized likelihood. Stat. Med. *17*, 407-429.
- 57. Fenton, T.R., and Kim, J.H. (2013). A systematic review and meta-analysis to revise the Fenton growth chart for preterm infants. BMC Pediatr. *13*, 59.

Figure Legends

Figure 1. Pre-Replication and Pre-Initiation Complexes in DNA Replication, Showing Components Mutated in MGS

Previously identified MGS-associated genes (labelled with orange lettering) encode members of the pre-replication complex (preRC, upper cartoon); these components are involved in the licensing of replication origins during the G_1 phase of the cell cycle. GMNN acts during other cell cycle phases to inhibit CDT1 but is degraded in late M (mitosis) phase (indicated by GMNN with dashed outline), permitting free CDT1 to participate in origin licensing in G_1 . In contrast, CDC45 contributes at the next major step in DNA replication, in which the coordinated action of many replication initiation factors including RECQL4 forms the preinitiation complex (PreIC, lower cartoon) to support the interaction of CDC45 and GINS1-4 with the MCM helicase, converting the latent form to an active helicase and initiating the unwinding of DNA.

Figure 2. Clinical Features of Individuals with Mutations in CDC45

(A) Pedigrees of families segregating mutations in *CDC45*. Note consanguinity in Family 5,7 and 11 and that all parents and unaffected siblings available for testing were heterozygousfor only one variant, consistent with autosomal recessive inheritance.

(B) Facial appearance of individuals with *CDC45* mutations. Many individuals demonstrate the facial characteristics of MGS including microtia, small mouth, full lips and micrognathia, but there is a marked spectrum of severity across the cohort. Note the consistent appearance of thin eyebrows.

(C) CT head scans and plain skull radiographs of individuals with *CDC45* mutations demonstrating premature fusion of cranial sutures. Note the discordance in suture fusion

between sibs P9-1 and P9-2. Imaging of P8 indicates progressive suture closure (P8 left panel, age 2 yr 4 m, P8 right panel, age 3 yr 8 m). Skull radiography of P5 demonstrates a copper-beaten appearance of the skull.

(D) Individuals with *CDC45* mutations have below average stature and many exhibited intrauterine growth retardation and microcephaly (both *in utero* and postnatal). SD, standard deviation; Lgt, length; OFC, occipitofrontal circumference; Wgt, weight; Hgt, height. Blue, individuals derived from craniosynostosis cohort; black, individuals derived from MGS cohort.

Figure 3. Mutations in CDC45, and Conservation of Substituted Residues in Orthologous Proteins

(A) Mutations identified in *CDC45* include variants predicted or shown to affect splicing (blue, position in gene shown in upper panel), predicted missense substitutions (black), or truncating mutations (purple). $\Delta ex5$ refers to the intragenic deletion encompassing exon 5 of *CDC45* in family P12 (green). Mutations do not cluster within any specific domain of CDC45. Domain architecture of human CDC45 indicated as described in Yuan *et al.*²⁶ The two alpha-beta domains are shown in purple and orange, respectively. The protruding helical motif (PHM) inserted in the first alpha-beta domain is shown in red. The middle all-helix interdomain (ID) is shown in blue, and the two linkers (L1 and L2) are shown in gray.

(B) Clustal Omega alignments of eukaryotic CDC45 proteins indicate mutated residues are well conserved, with several residues (such as p.Asn76, p.Arg157, p.Pro321 and p.Pro463) conserved as distantly as fungi. Blue shading indicates proportion of conserved amino acids.

Figure 4. Structural Consequences of CDC45 Substitutions

(A) Structural model of human CDC45 demonstrating the spatial distribution of substitutions in the protein. Red boxes indicate position of missense mutations, with side-chains of these substituted residues displayed in green.

(B) CDC45 in context of the *S. cerevisiae* CMG complex from the 3.8Å cryo-electron microscopy structure.²⁶

(C-E) Modeling of substitutions that cluster in the central core of CDC45. (C) Proline 463 is an evolutionarily conserved position, with an essential role in N-capping of the alpha-helix localized between Pro542 and Cys556 in yeast Cdc45 (Pro463 and Phe477 in human). Given the location of Pro463 in the CDC45 structural core, the substitution to leucine is predicted to have severe effects on the stability of this helix⁵³ and therefore overall stability of the protein. (D) Asn76 is predicted to be part of the divalent cation coordination sphere. While the DHH domain of CDC45 is currently considered to be catalytically inert,^{30, 54} residues D26, D28, D99 and H101, corresponding to the catalytic core of prokaryotic RecJ exonucleases and the DHH superfamily of phosphoesterases, are conserved (highlighted by black boxes).^{26, 32, 55} This suggests the continued presence of a divalent cation at this position (predicted divalent cation indicated by yellow sphere). Asn76 is therefore thought to be part of a divalent cation coordination sphere which, in other homologs, is the phosphoesterase catalytic centre. The substitution to histidine would introduce a positive charge into the putative divalent cation coordination sphere, thereby disrupting the network of interactions among water molecules, charged residues and divalent cation/s observed in different members of the superfamily of DHH phosphoesterases.^{33, 34, 35} (E) Proline 321 (corresponding to Pro369 in yeast) is a key part of the CDC45 interacting surface with Mcm2,²⁶ given that in yeast it lies within 3.5Å of Ile289 of Mcm2 (corresponding to Ile211 in human MCM2). Additionally, this proline has a key role in the N-capping of the alpha-helix localized between Leu370 and Gln374 in yeast

Cdc45.⁵³ Therefore, substitution of proline 321 to threonine is likely not only to have effects on CDC45-MCM2 interaction but also on CDC45 protein stability. The interacting pair Pro321 (in CDC45) and Ile211 (corresponding to Ile289 in Mcm2) are shown in sticks (green) inside their van der Waals surfaces.

Figure 5. Synonymous *CDC45* Variants Alter Levels of Alternative Splicing of Exon 4 in Lymphoblastoid Cell Lines

(A) The synonymous variant c.318C>T present in P1 increases the ratio of exon3/5 transcript compared to controls. (i) RT-PCR demonstrating an enrichment in the exon3/5 lower band. (ii) Results of qRT-PCR to assay the relative amounts of exon 4 in P1, P2 and four unaffected controls. Products were amplified with primers in exons 3 and 4, and compared with products generated from primers in exons 1 and 3. cDNA from P1 showed a 0.43-fold reduction in exons 3-4 relative to exons 1-3. Levels of exons 5-6 compared with exons 1-3 are also shown, no differences were observed. Error bars, standard deviation. n=3 experimental replicates of qPCR. (B) The synonymous variant c.333C>T together with the trans c.203A>G, p.Glu68Arg splice site variant identified in P9-1 and P9-2, increases aberrant splicing across this region of the gene. (i) RT-PCR demonstrating an increase in exon3/5 transcript and the creation of a novel transcript where exon 3 has been skipped due to the c.203A>G mutation. (ii) Total RNA extracted from LCLs derived from P9-1 was analyzed by qRT-PCR. Full length only transcripts were amplified using a forward primer (F 2/3) spanning the exon 2exon 3 boundary and a reverse primer (R 4/5) spanning the exon 4-exon 5 boundary. The transcript missing exon 4 was specifically amplified using the F 2/3 primer paired with a reverse primer (R 3/5) which spanned the exon 3-exon 5 boundary. Total amount of minus exon 4 (-Ex4) and full length (FL) transcript was normalized to GAPDH for P9-1 and control.

Error bars indicate standard deviation (n = 3 experiments). A two tailed, unpaired student's ttest was performed and p-values are indicated.

Figure 6. CDC45 Protein Levels are Significantly Reduced in Lymphoblastoid Cell Lines and Primary Amniocytes

Cell lines from affected individuals (P1, P2, P9-1) and amniocytes derived from affected fetuses (P12-1 and P12-2) demonstrate a significant reduction in endogenous CDC45 levels compared to three independent controls (LCL samples - one way ANOVA comparing all LCL samples against C1, NS, p > 0.01, ****p < 0.0001, n = 5 biological replicates; amniocyte samples - one way ANOVA comparing all amniocyte samples - one way ANOVA comparing all amniocyte samples against C4. ***p < 0.001, ****p < 0.0001, n = 3 biological replicates).

Subject	Country of Origin	Allele 1	Allele 2	Segregation
P1ª	UK	c.318C>T, p.Val106= (SE)	c.677A>G, p.Asp226Gly	Het, M, P, (2) ^b
P2	UK	c.226A>C, p.Asn76His	c.469C>T, p.Arg157Cys	Het, M, P, (1)
P3	USA	c.653+5G>A (SE)	c.1487C>T, p.Pro496Leu	Het, M, P
P4	Bangladesh	c.1A>C, p.Met1?	c.1388C>T, p.Pro463Leu	nps
P5	Bangladesh	c.1270C>T, p.Arg424*	c.1388C>T, p.Pro463Leu	Het, M, P
P6	Belgium	c.791C>A, p.Ser264Tyr	c.791C>A, p.Ser264Tyr	Hom, M, P
P7-1	Egypt	c.1660C>T, p.Arg554Trp	c.1660C>T, p.Arg554Trp	Hom, M (P deceased)
P7-2	Egypt	c.1660C>T, p.Arg554Trp	c.1660C>T, p.Arg554Trp	Hom, M, P
P8	Sri Lanka	c.1388C>T, p.Pro463Leu	c.1532delC, p.Pro511GInfs*36	Het, M, P
P9-1	Turkey	c.203A>G, p.GIn68Arg (SE)	c.333C>T, p.Asn111= (SE)	Het, M, P
P9-2	Turkey	c.203A>G, p.GIn68Arg (SE)	c.333C>T, p.Asn111= (SE)	Het, M, P
P10 [°]	New Zealand	c.[464A>G; 961C>A], p.[Glu155Gly; Pro321Thr]	c.1440+14C>T (SE)	Het, M, P
P11	Egypt	c.1660C>T, p.Arg554Trp	c.1660C>T, p.Arg554Trp	Hom, M, P
P12-1	Netherlands	c.(485+1_486-1)_(630+1_631-1)del, p.lle115_Glu162del	c.893C>T, p.Ala298Val	Het, M, P
P12-2	Netherlands	c.(485+1_486-1)_(630+1_631-1)del, p.lle115_Glu162del	c.893C>T, p.Ala298Val	Het, M, P

Table 1. Mutations Identified in CDC45.

Refseq Accession: NM_003504.4 SE: splicing effect. Het, compound heterozygous in affected individual, nps, no parental samples available, M, mutation identified in mother, P, mutation identified in father, Hom, homozygous in affected individual. ^aMutation previously reported²⁸ (note mutation numbering differs in previous publication due to different reference transcript). ^bNumbers in brackets indicate number of clinically unaffected siblings shown not to have inherited the compound heterozygous genotype. ^cSubject P10 segregates two rare missense variants on one allele, in combination with a splice region mutation on the second allele. Based on conservation of CDC45 residues across different species the p.Pro321Thr variant is more likely to be deleterious as this residue is conserved through to *S. cerevisiae* (Figure 3).

				Birth	, SD		Cu	urrent E	xam, SD)	MGS	S Features				
Subject	Consan	Gender	Gest.	Lgt (cm)	OFC (cm)	Wgt (kg)	Age	Hgt (cm)	OFC (cm)	Wgt (kg)	Microtia	a/hypoplastic patella	Thin Eyebrows	Developmental Delay	Craniosynostosis	Other Clinical Features
P1	N	Μ	40	NA	NA	-0.7 (3.2)	28y	-0.7 (173)	-0.4 ^b (55.5)	+1.3 (85.7)	+	-	+	none	right unicoronal	urethral stricture, mild 2/3/4 syndactyly of hands
P2	N	F	38	NA	NA	-1.2 (2.5)	4y 4m	-0.9 (100)	-3.1 (47.6)	-0.8 (15.3)	-	-	+	none	bicoronal	anterior anus, bilateral strabismus
P3	N	Μ	38	NA	NA	-1.7 (2.4)	16y 3m	-3.6 (147)	-3.7 (50.5)	-3.9 (35.3)	+	-	+	mild	bicoronal	R choanal atresia, high palate, imperforate anus, vesicoureteral reflux, hypospadias, 2-3 syndactyly of toes, scoliosis, C1-C3 and C4-C7 fusion, bilateral radial head dislocation
P4	N	F	36	NA	-2.9 (28.5)	-3.2 (1.45)	5y 5m	-7.7 ° (48)	-8.3° (33)	-6.1 (9.9)	+	+	+	severe	large anterior fontanelle (x-ray)	cleft palate, exorbitism, bilateral radial head dislocation, thoracic vertebral segmentation defects, digital clubbing, pulmonary hypoplasia, anterior anus, cliteromegaly
P5	3rd	F	40	NA	NA	-4.7 (1.5)	6у	-4.6 (93)	-8.1 (42.5)	-7.5 (9.2)	+	+	+	none	copper-beating on skull radiograph	cleft palate, bowed legs, anorectal malformation
P6	N	Μ	34	0.3 (46.5)	-1.1 (30)	-1.2 (1.88)	22y	-4.8 (145)	-5.4 (48)	-8.4 (32.5)	+	+	+	severe	bicoronal and bilambdoid	AV block grade II, SN hearing loss, joint laxity, chiari I malformation, myopia
P7-1	1st	Μ	term	-4 (43)	-4 (30)	-2.5 (2.4)	3y 6m	-4.1 (83)	-5.5 (44)	-6 (8.5)	+	+	+	none	coronal, sagittal	anal stenosis, budlike mouth, ASD, VSD, undescended testes, hypospadius
P7-2	1st	F	term	-4.9 (41)	-2.9 (31)	-2.2 (2.45)	5y 2m	-3.5 (94)	-4 (47)	-4.2 (11.5)	+	+	+	none	coronal, sagittal	AV canal
P8	N	F	37	-0.4 (47)	-4.6 (27)	-1.7 (2.14)	4y	-4.9 (82)	-10.4 (38.6)	-7.6 (7.5)	+ lowset	NA	+	NA	bicoronal and bilambdoid, broad patent sagittal with bifid metopic suture	vestibular, anterior anus, hearing mild conduction delay, proptosis, VSD, duodenal stenosis
P9-1	N	М	42	-1.4 (50)	NA	-1.5 (3.2)	7у	-1.3 (115)	-2.3 (50)	-0.4 (22)	+	+	+	none	none	high arched palate, micropenis

Table 2. Clinical Characteristics of Individuals with Mutations in CDC45.

P9-2	Ν	Μ	term	0.5 (52)	-1.7 (33)	-1.2 (3.0)	16m	-2.5 (73)	-4.7 (43)	-3.7 (7.5)	+	N/A	+	none	bicoronal with widely patent metopic and sagittal sutures	none
P10	Ν	F	36	NA	-1.8 (30)	-1.5 (2.04)	25y	-2.5 (149)	-4.7 (49)	NA	+	+	+	none	bicoronal	high palate, hearing loss, breast agenesis, heart block
P11	1st	F	40	NA	NA	-4.0 (1.75)	4y 7m	-2.5 (95)	-4.6 (-4.6)	-3.2 (12)	+	+	+	none	coronal and lambdoid	small ASD
P12-1	Ν	F	22+3ª	-1.9 (25)	-3.9 (15.5)	-3.0 (0.33)		то	P		+	Not reported	+	N/A	Yes	anterior anus
P12-2	Ν	М	30+5ª	-0.4 (39.1)	-2.9 (24)	-1.1 (1.15)		то	P		lowset	Not reported	+	N/A	bicoronal, partial lambdoid, squamous and lateral parts of the occipital bone	pre-axial polydactyly of both hands, small VSD, meconium peritonitis, multiple nodules of ectopic thymic tissue, high palate

SD, standard deviation. Calculated using LMSgrowth⁵⁶ and adjusted for sex and gestation, with the exception of ^a, where Z-scores were calculated using the Fenton 2013 Growth Calculator.^{57 b}measurement at 15 years of age, after surgery. ^cmeasurement at 5 months of age. NA, not available; TOP, termination of pregnancy.







В

A. C. S. S.

H. sapiens	66 KEQFHYFILINCGANVDLLD
P. troglodytes	66 REQERVETLINCGANVOLLD
C. familiaris	66 KEQFHYFYL INCGANVDLLD
M. musculus	66 KEQESYFILINCGANVOLLD
G. gallus	66 KDQFKYFVFINCGANVDLLE
D. rerio	66 KEQYKYFVLINCGANVDLLE
D. melanogaster	66 QGDVKYVVLV <mark>NCG</mark> GCVDIVE
A. gambiae	66 RGD I RYVLLVNCGGC I DIVD
C. elegans	69 QEQKCHYILINCGATRSLTR
S. cerevisiae	75 DDNINSLLLVGFGGVIDLEA
S. pombe	76 NEDIKFIILLNCGTMVDLNN
	p.Ser264Tyr
H saniens	257 EENTLSVDCTRISFE
P. troalodytes	257 E ENTLSVDCTRISFE
C. familiaris	260 EEHAUSVDCTRISFE
M musculus	257 EENMLSVDCTRISEE
G gallus	257 EENSUS LDCMR LAFE
D rerio	263 E ENSUS LDCMR INEE
D melanogaster	273 QNTMSASKITEE
A gambiae	273 QALQTSVKLVFF
C elecans	260 N Q- GKYDDLLHLTEG
S cerevisiae	
S nombe	289 NOLVGRAHGKTPHDOSLELE
0. рошье	
	p.Pro463Leu
H. sapiens	453 GTPDVMLFSRPASLSLLSKH
P. troalodytes	453 GTPDVMLFSRPASLSLLSKH
C. familiaris	456 GTPDIVLESKPASLSLLSRH
M. musculus	453 GTPDVTLESKPASLSLLSRH
G. gallus	453 GTPDVKLESKPVSLCLLSKY
D. rerio	459 GTPDVKLESKPLSLTLLSKY
D. melanogaster	465 EHAFESYPYALGULARE
	LOG CONTRACTOR AND

p.Met1? p.Gln68Arg

p.Asn76His

p.Val106= p.Asn111=

Δex5

KEQFHYFILI	N	CGANVDLLDI 86
KEQFHYFILI	N	CGANVDLLDI 86
KEQFHYFVL I	N	CGANVDLLDI 86
KEQFSYFILI	Ν	CGANVDLLDI 86
KDQFKYFVF I	N	CGANVDLLEI 86
KEQYKYFVL I	Ν	CGANVDLLET 86
QGDVKYVVLV	N	CGGCVDIVEL 86
RGDIRYVLLV	N	CGGCIDIVDV 86
QEQKCHYILI	N	CGATRSL TRL 88
DDNINSLL	G	FGGVIDLEAF 95
NEDIKFILL	N	CGTMVDLNNY 95

Diens 257 EENTLSVDCTRISFEY 2	74
Ilodytes 257 E ENTLSVDCTRISFEY 2	74
niliaris 260 E EHALSVDCTRISFEY 2	77
ISCUIUS 257 E ENMLSVDCTRISFEY 2	74
lus 257 EENSLSIDCMRIAFEY 2	74
0 263 EENSLSIDCMRINFEY 2	80
lanogaster 273 Q NTM SASKITER 2	87
mbiae 273 QAIQTSVKIVFES 2	87
gans 260 NQ-GKVDDLLHITFGR 2	76
evisiae 308NSVKTPDTLTLNIQP 3	22
nbe 289 NQIVGRAHGKTPHDQSIRLED 3	11

								1.						
sapiens	453	GТ	PD	VМ	LF	SR	Ρ	AS	L	SL	Ľ٤	s KH	۱L	473
troglodytes	453	GТ	PD	VM	LF	SR	Ρ	AS	L	SL	LS	3KH	IL	473
familiaris	456	GТ	PD	١V	LF	SK	Ρ	AS	L	SL	LS	SR F	۱L	475
musculus	453	GТ	PD	VT	LF	sĸ	Ρ	AS	L	SL	LS	SR H	IL	473
gallus	453	GТ	PD	٧K	LF	sk	Ρ	VS	L	CL	LS	3 KY	Ĺ	473
rerio	459	GТ	PD	٧K	LF	sk	Ρ	LS	L	ΤL	Ľ۶	3KY	Ĺ	479
melanogaster	465		- E	HA	FF	SY	Ρ	ΥA	L	GL	L,A	ARF	L	483
gambiae	465		- E	٧S	FF	sc	Ρ	ΥG	L	LL	L/	١RF	L	483
elegans	455	SΙ	DE	DM	V -	DS	R	ΗF	L	ΥN	TA	٩GF	M	474
cerevisiae	533	G-	PD	LD	LΥ	RN	Ρ	LΤ	L	LR	LC	SNW	٧L	552
pombe	516	G-	PD	LΚ	١F	MH	Ρ	LA	L	ΤK	MS	SLW	٧I	535

p.Arg157Cys

p.Arg424* p.Pro511GInfs*36

													_										
150	-	-	-	G	s	E	Ρ	s	Е	k	R	Т	R	L	Е	Е	Е	Г	V	Е	Q	16	7
150	-	-	-	G	s	E	Ρ	s	Е	k	R	Т	R	L	Е	E	Е	L	V	Е	Q	16	7
153	-	-	-	G	s	E	Ρ	s	E	k	R	Т	R	L	E	E	Е	V	V	A	Q	17	0
150	-	-	-	G	s	E	Ρ	s	Е	k	R	Т	R	L	Е	E	E	Г	V	Е	R	16	7
150	-	-	-	G	s	E	Ρ	s	D	k	R	R	R	F	Е	E	Е	V	I	E	R	16	7
155	-	-	-	S,	Ą	Е	Ρ	s	G	k	R	R	R	F	D	E	D	A	L	E	R	17	2
167	-	-	-	Q,	A	Ρ	R	S	R	k	L	S	-	-	R	L	E	R	Н	Е	Q	18	2
167	-	-	-	G	G	A	Ρ	K	Ρ	Α	Ν	N	-	-	R	I,	Q	R	١	Е	Q	18	2
158	-	-	-		-	-	-	-	-	-	S	Y	Е	Q	R	М	Е	K	I	R	R	16	8
202	-	÷	-	ΕI	D	E	Т	I	S	Ν	K	R	G	Ν	S	S	I	G	Ρ	-	Ν	21	8
178	Y	S	S	SI	D	Y	Q	A	R	S	R	R	R	F	S	Е	E	Т	Т	-	Q	19	7

p.Ala298Val

288	D	S	L	С	N	Т	S	Y	Т	A	A	R	F	K	Ľ	W	S	V	н	G	Q	30	8
288	D	S	L	С	N	T	S	Y	Т	A	A	R	F	K	Ľ	W	S	V	Н	G	Q	30	8
291	D	S	L	С	N	Т	С	Y	Т	A	A	R	F	K	Ľ	W	S	V	Н	G	Q	31	1
288	Е	S	Ĺ	Y	N	Т	S	Y	Т	A	A	R	F	K	Ľ	W	S	V	Η	G	Q	30	8
288	Е	S	L	С	N	Т	S	Y	Т	S	A	Т	L	K	Ľ	W	S	V	Q	G	Q	30	8
294	Е	S	I	С	N	S	С	Y	Т	S	С	S	F	K	Ľ	W	S	I	N	G	Q	31	4
301	Е	S	Μ	R	Y	S	R	Y	S	s	С	Q	L	K	Ľ	W	Т	L	R	G	Е	32	1
301	D	S	L	R	Y	S	Y	Y	Ρ	A	С	R	L	K	Ľ	W	Т	Н	K	G	D	32	1
290	S	A	M	Μ	V	S	E	Y	F	s	I.	K	Т	K	N	W	Т	Q	K	G	D	31	0
336	D	S	F	Y	Y	S	Ν	Y	V	N	A	K	L	S	Ľ	W	N	E	N	G	K	35	6
325	D	S	М	L	н	S	A	Y	V	G	s	R	L	н	1	W	S	Е	Е	G	R	34	5

p.Pro496Leu

						_		_		_	_		
486	CKL	LPL	VM.	AA	Ρ	L	SM	E	Η-	G	T١	/T	505
486	CKL	LPL	VM.	AA	Ρ	L	sv	E	Η-	G	T۱	/Τ	505
489	CKL	LPL	VM.	AA	Ρ	L	sv	E	Q -	G	T١	/T	508
486	CKL	LPL	VM.	AA	Ρ	L	sv	E	Q -	G	T١	/T	505
486	CKL	LPL	ΙM.	AA	Ρ	M	DV	E	Q-	G	T١	Ζ.	505
492	CKI	LPL	I M	AA	Ρ	L	DI	E	K-	G	T١	Ζ.Γ	511
496	ASD	LPL	ΙA	SC	Ρ	L	NA	S	E -	Gl	v10	CL	515
496	AQE	LPL	I A'	VA	Ρ	L	DL	A	R-	G	13	SL	515
487	T	KPL	I AI	MF	Ρ	L	ΤG	E	SA	G	ΝL	$_{\rm V}$	505
562	KQL	LPM	VL.	AS	T	-	D-	E	ΝT	D	T١	1L	580
549	LRH	LPL	VL.	AA	F	÷	٧-	E	ΕK	NF	۲۶	7L	567

p.Asp226Gly

19

18

Splicing

216	М	L١	Ŵ	N.	A	1	V	G	L	Т	D	Q	w	V	Q	D	K	I	Т	Q	М	236	
216	M	LV	Ņ	N.	A	l	V	G	L	Т	D	Q	w	V	Q	D	ĸ	I	Т	Q	М	236	
219	M	L١	W	N.	A	Ĩ	٧	G	L	Т	G	Q	w	V	Q	D	K	I	Т	Q	М	239	
216	M	LV	W	N,	A	ł	٧	G	L	Т	D	a	w	V	Н	D	K	L	Т	Q	М	236	
216	M	LV	M	N.	A	1	٧	G	L	T	D	Q	w	V	Q	D	K	L	Т	Н	L	236	
222	M	L٧	Ŵ	N	s	1	I	G	L	Т	D	a	w	V	Н	D	ĸ	I	Ρ	Н	М	242	
232	L	L٧	W	N.	A	I	V	G	I	T	Е	Q	LI	L	L	G	K	T	E	S	G	252	
232	L	LV	W	N.	A	I	V	G	V	Т	E	Q	KI	L	L	G	K	V	E	S	А	252	
218	Т	M	V	F	Τ.	A	٧	G	L	Ν	S	A	M,	A)	D	K	L	L	S	I	Е	238	
269	Ν	LV	V	LĮ	Ν	1	L	G	Т	T	S	L	D	Ι.	A	Y	-	-	-	A	Q	286	
248	M	L١	V	L	A	L	۷	G	L	Т	С	L	Е	I	Н	С	Q	S	S	K	Κ	268	

p.Pro321Thr

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311	L	Q	Е	F	L	A	D	М	GI	_ F	L	K	Q	V	K	Q	KI	F	Q.	A	331
311	L	Q	E	F	L	A	D	M	GI	F	L	K	Q	V	K	Q	KI	F	Q,	A	331
314	L	Q	Е	F	L	A	D	V	GI	F	L	K	Q	V	K	Q	KI	F	Q	S	334
311	L	Q	Е	F	L	A	D	М	GI	_ F	L	K	Q	V	K	Q	KI	F	Q	S	331
311	L	Q	E	F	L	A	D	M	Gl	F	L	K	Q	V	K	Q	KI	F	N	s	331
317	L	Q	Е	F	L	A	D	М	GI	F	L	K	Q	V	R	Q	KI	F	N	S	337
324	L	Н	E	L	L	L	E	М	GI	_ F	L	V	Н	A	R	Q	T	Y	G.	A	344
324	M	Ν	Е	L	L	V	D	M	Gl	F	L	٧	Q	A	K	Q	Т	Ŧ	N.	A	344
313	I	R	Н	L	L	A	Q	L	G	I T	L	Н	Е	Т	K	Q	KI	-	Ε.	A	333
359	L	Н	K	M	F	A	R	M	G	F	L	s	Т	A	Q	Е	T١	N	Ľ	Y	379
348	L	н	K	L	L	A	K	M	GI	S	L	V	Е	С	K	Q	Τì	Y	1	Н	368

p.Arg554Trp

		_	_	_	_	_	_		_	_	_		_	_	_	_	_	_			_	
544	L	s	V	I	Е	L	K.	A	E	D	R	S	K	F	L	D	A	L	L	s	L	564
544	L	s	V	1	Е	L	K.	А	E	D	R	s	K	F	L	D	A	L	L	S	L	564
547	L	s	٧	I	Е	L	K	A	E	D	R	s	K	F	L	D	A	L	V	S	L	567
544	L	S	V	Ĭ	Е	L	K.	A	E	D	R	s	K	F	L	D	A	L	V	S	L	564
544	Μ	s	I	1	Е	L	K	Т	E	D	R	s	K	F	L	D	A	L	I	s	L	564
550	Т	s	V	Ì	Е	L	K	M	E	D	R	G	K	F	L	D	A	L	L	Т	L	570
553	Ρ	A	V	V	Q	L	R	Q	s	D	L	Т	R	F	L	D	s	L	Т	V	L	573
552	Т	A	V	I	Q	I	K	Q	s	D	С	Т	K	F	L	D	A	L	Т	V	М	572
548	S	D	I	1	R	L	K	s	E	D	R	Т	R	F	I	D	F	L	Q	Т	V	568
627	S	s	1	I	Е	1	RI	R	E	D	L	s	Ρ	F	L	Е	K	L	Т	L	S	647
615	A	s	V	Ĩ	E	С	Q	ĸ	s	D	L	G	V	F	L	Е	s	L	s	F	ĸ	635









Supplementary Note.

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Note on accession numbers for CDC45

Human *CDC45* encodes a 20-exon gene that has been annotated as including three major spliceforms. As well as the full-length version which encodes a protein of 598 amino acids, alternative splicing to skip either exon 7 alone or both exons 4 and 7 leads to shorter isoforms with in-frame deletions of 32 and 78 amino acids, respectively. Whereas exon 4 shows high conservation in mammalian species both in sequence and occurrence of exon skipping, neither is the case for exon 7, so the physiological significance of the isoform including exon 7 is currently uncertain. Virtually all previous publications investigating the structure and function of CDC45 employ the numbering of the middle-sized, 566 amino acid isoform, so for consistency with prior work all numbering used here refers to NM_003504.4.



Figure S1. Family 12 segregates a genomic deletion encompassing exon 5 of CDC45.

(A) Manual interrogation of exome sequencing reads across the CDC45 gene suggested a decrease in coverage across exon 5 in both affected samples and the carrier mother, compared to the father, who carries the c.893C>T, p.Ala298Val variant. While the reduction in coverage was not statistically significant, the presence of a rare variant on the *trans* allele in both affected samples prompted further investigation. BAM files visualized using IGV. (B) MLPA, comparing the quantity of DNA in a test sample vs a synthetic control with multiple probes across CDC45, was performed in both affected samples and the mother, and all samples showed a deletion of exon 5. (Bi) Peak traces of control (red) and affected (blue). (Bii) Calculated ratio of DNA. Both probes for exon 5 demonstrate an approximately 50% reduction, consistent with a heterozygous deletion of exon 5. Probes for exon 4 and exon 6 do not show a reduction, indicating the breakpoints for the deletion lie within intron 4 and intron 5, respectively. Reports generated using GeneMarker Software (SoftGenetics Inc.). (C) RT-PCR and sequencing of cDNA from carrier parents in Family 12. (Ci) RT-PCR of exons 3-6 in CDC45 from cDNA derived from parents in Family 12. There is an increase in a lower migrating band in the mother, who carries the exon 5 deletion allele (a faint co-migrating lower band visible in all samples is caused by physiological skipping of exon 4, see figure 5). (Cii) Sequencing of each band from the maternal sample indicates the lower migrating band (labelled Band 2) comprises exons 3, 4 and 6, with complete skipping of exon 5, resulting in an in-frame transcript.

					Analysis Software					
Family	Clinical Diagnosis/ Study Focus	Design	Library Kit	Platform	Mapping	Variant Calling	Annotation	Coverage	Filtering parameters	Control populations
1	Craniosynostosis (SC_2930), clinical genome sequencing	Genome	NEBNext DNA Sample Prep Master Mix Set 1 Kit	Illumina HiSeq2000	BWA ¹ , Stampy (v1.0.12-1.0.22) ²	Platypus $(v.0.2.4)^3$	ANNOVAR ⁴	28x	Minor allele absent	1000 Genomes, ESP
2	Craniosynostosis	Exome	TruSeq V2	Illumina HiSeq2000	Novoalign (Novocraft Technologies)	SAMtools (v0.1.19)	ANNOVAR	55x	Minor allele absent	1000 Genomes, ESP
4	Primordial dwarfism	Exome	SureSelect V4	Illumina HiSeq2500	BWA, Stampy	SAMtools, Dindel	Ensembl VEP	50x	MAF < 0.005, ≥ 1 deleterious consequence prediction	1000 Genomes, ESP
12	Diagnostic	Exome	SureSelect ^{XT} V5	Illumina HiSeq2500	Custom in-house pipeline MAGPIE (based on BAM-MEM and GATK)			70x	SeattleSeq, LOVDplus	GoNL, 1000 Genomes

Table S1. Experimental details for next-generation sequencing undertaken in this study.

MAF, minor allele frequency; GoNL, genomes of The Netherlands.

Table S2. Primers and amplification conditions for resequencing.

Dideoxy-sequencing primers to confirm variants in the craniosynostosis cohort ^a					
	_		Primer sequence $5' \rightarrow 3'$	-	
Variant	F	Primers	Forward	Reverse	Product size (bp)
c.226A>C c.318C>T	4	4F/4R	TAGTGATGAAGGAAAAGGGGGCCTCCTCG	CTGTTCCCAGTCCCACAGGGTAGTCAG	314
c.469C>T 5F/5R		5F/5R	ACCACGTATGGTGTAACTCTGGTGCCTCAC	CTTGGCCTGGCAGGCTTCAGGATGAC	436
c.653+5G>A 8F/8R		8F/8R	CATGAGCCTTAGACTTCTCTGCTTCCTTAC	ATCACACATACCCAGAAAGGGGCTGCA	240
c.677A>G	9	9F/9R	GAGAGAGGCCACCTGACTGAGGCAAG	CAGTCCTCACTTCTCAATAGGACCCTA	335
c.1487C>T	1	16F/16R	CATAGACAAAGAACCGGCGCTGCAAACTG	GACACTAGAGGCAAATACCACTCTACTCAG	223
Dideoxy-se	equencing p	primers for	<i>CDC45</i> in the MGS cohort ^b		
Exon(s)	Forward			Reverse	Product size (bp)
1-2	GATTTGG	CGGGAGTC	TTG	ACGCAGCACCCTCACCTC	382
3	TCTTTTCA	AAGGTAAA	ITGCTAATGTC	TGGAAATAGAATGCTGTGTCAAG	230
4	AGTGATG	GAAGGAAAA	AGGGGC	CCAAGAACACCAGGTGAGAC	274
5	TCAGCTT	ATTAGGAA	ATGATAAGATTC	CCACTGCCCTCAGTACACC	343
6	CTGGGCC	CTACTGACT	ICTGC	TCACTTCAAGCTAGAATAACCTTTC	519
7	TTTTCCAA	AAATATTTC	GGCTTCC	AGGACACTGACCCCTGGAG	174
8	TCAAGTC	CAGTCTGG	CTGC	TTATGTGGCACTGACCAAGG	201
9	CCTCATTO	GAGCCCAG	GTG	GTGGCTATGGTCAGGCTGTG	179
10	CTGCCTG	GTAAGAGC	TGGAG	ACCCACAGAGGGTGACAAAG	257
11-12	CCACCTG	GCTGGAGTT	ACGAG	CCCTTCACTTTAATGAACCTGG	610
13	GCAGTGA	AGAGCTTGC	CCAC	CCCGTAACTGACCTCCCTG	293
14-15	CAAGCAA	ACTATTTG	TGACTTTGG	GGAGCTCTCCAAGCACCTG	437
16-17	CTCTGAG	TGTTGAGC	IGGGG	AACTTGCTTGGTTCCAGGG	506
18	TTCTGTG	CCCTGTCTT	GTG	AGCAGGGCATCAGGGTC	199

	Primers for Fluidigm resequencing in the craniosynostosis cohort ^c		
ID	Forward	Reverse	Length (bp, excluding universal adaptors)
CDC45_1	ACACTGACGACATGGTTCTACAAGTCTTGACCGCCGCC	TACGGTAGCAGAGACTTGGTCTCCCCTCATCCCTTCCCCA	173
CDC45_2	ACACTGACGACATGGTTCTACACCTCGGACGTGGATGCT	TACGGTAGCAGAGACTTGGTCTCTCACTTGCTGTTCACCTCCT	186
CDC45_3	ACACTGACGACATGGTTCTACAGTCCAGAGCCAGGTGACG	TACGGTAGCAGAGACTTGGTCTCTGAAGGATCTTGCACGCAC	210
CDC45_4	ACACTGACGACATGGTTCTACAAGATAGGCTTTGAAAAACACTCTCTCT	TACGGTAGCAGAGACTTGGTCTTGCTGTGTCAAGAAATTACAATTGAGTA	186
CDC45_5	ACACTGACGACATGGTTCTACAATGAAGACACTATATTCTTTGTGTGTG	TACGGTAGCAGAGACTTGGTCTTAGTCAGACACACACCTTACCAAG	167
CDC45_6	ACACTGACGACATGGTTCTACATCAGAGGCTTAGTGATGAAGGAAAA	TACGGTAGCAGAGACTTGGTCTATTGACGACATTGACTGGCCTAT	192
CDC45_7	ACACTGACGACATGGTTCTACACCGTAGAAAGAGGAGAGGGCTTAATT	TACGGTAGCAGAGACTTGGTCTGATAGCTGTGAGAGACCCCAAACT	215
CDC45_8	ACACTGACGACATGGTTCTACAGGCCTACTGACTTCTGCCAAA	TACGGTAGCAGAGACTTGGTCTGGCTTCTGGTGCTGGGAG	193
CDC45_9	ACACTGACGACATGGTTCTACAGTGCCACCCATACCTCTGAC	TACGGTAGCAGAGACTTGGTCTCCATGGTCTCTAATGGATGG	188
CDC45_10	ACACTGACGACATGGTTCTACACTTCCTGTACTGCTACTTCCTGTT	TACGGTAGCAGAGACTTGGTCTACATTTGAGGACACTGACCCC	164
CDC45_11	ACACTGACGACATGGTTCTACAGCCTTAGACTTCTCTGCTTCCTT	TACGGTAGCAGAGACTTGGTCTGTGGCACTGACCAAGGCA	145
CDC45_12	ACACTGACGACATGGTTCTACACATCATCTCACTCCATCCCCC	TACGGTAGCAGAGACTTGGTCTGCTATGGTCAGGCTGTGCA	200
CDC45_13	ACACTGACGACATGGTTCTACATTCCCGCCACAACCACC	TACGGTAGCAGAGACTTGGTCTCACCTGCTTTCCCTCCTGG	191
CDC45_14	ACACTGACGACATGGTTCTACACTGCCTGGTAAGAGCTGGAG	TACGGTAGCAGAGACTTGGTCTGAGTGTGTTCTCCTCATCCTCG	141
CDC45_15	ACACTGACGACATGGTTCTACATTCAAGCTGTGGTCTGTGCA	TACGGTAGCAGAGACTTGGTCTTTCTGGGATCTAGGGCAAGGA	194
CDC45_16	ACACTGACGACATGGTTCTACAGCTTGCATGCTCCTTGACTG	TACGGTAGCAGAGACTTGGTCTCTCCTGGAGCCGCTTCTG	144
CDC45_17	ACACTGACGACATGGTTCTACAGTCGGTGTGTGGTGACCTC	TACGGTAGCAGAGACTTGGTCTCTTTAATGAACCTGGATGTAATGAAGATAAATC	200
CDC45_18	ACACTGACGACATGGTTCTACAGCTGTGTTTGCTCCCATGAC	TACGGTAGCAGAGACTTGGTCTCAAACCCACAGCCCTCCT	207
CDC45_19	ACACTGACGACATGGTTCTACAAACTCGCCAAGAAGCAGCT	TACGGTAGCAGAGACTTGGTCTAGACGAAGCCACTGGTGC	161
CDC45_20	ACACTGACGACATGGTTCTACATTACCAAGAGTTCCTGACAAGCA	TACGGTAGCAGAGACTTGGTCTAATGGTCTGCTGGGTGGC	127
CDC45_21	ACACTGACGACATGGTTCTACACAGTGGCTTCGTCTGACCAT	TACGGTAGCAGAGACTTGGTCTGGCCAGCATCTAACTCCCC	201
CDC45_22	ACACTGACGACATGGTTCTACACTCTGAGTGTTGAGCTGGGG	TACGGTAGCAGAGACTTGGTCTGGACGAGTGGAAGCTGCTC	200
CDC45_23	ACACTGACGACATGGTTCTACACATGTCTTGTGTGTCAGCAGC	TACGGTAGCAGAGACTTGGTCTCCAGCCATGCCAACTTGC	187
CDC45_24	ACACTGACGACATGGTTCTACAGTGAGCAGCTTCCACTCGT	TACGGTAGCAGAGACTTGGTCTCTTCTCAAACGCCCTCCCAA	196
CDC45_25	ACACTGACGACATGGTTCTACATCCCTTCTCACGGCTGTTTTT	TACGGTAGCAGAGACTTGGTCTCCAAGTAGAAGCCTCCGTTGA	201
	·	·	

MLPA Primer Sequences ^d						
ID	Sequence					
CDC45-EX4-5	GGGTTCCCTAAGGGTTGGAGGCCAGTCAATGTCGTCAATGTATA					
CDC45-EX4-3	CAACGATACCCAGGTACTTTTTGTGTCTAGATTGGATCTTGCTGGCAC					
CDC45-EX3-5	GGGTTCCCTAAGGGTTGGAGATAGGCTTTGAAAACACTCTCTCCCAGGC					
CDC45-EX3-3	CTTGTTCCAGTGTGACCACGTGCAATATACGTCTAGATTGGATCTTGCTGGCAC					
CDC45-EX5B-5	GGGTTCCCTAAGGGTTGGAGCGCACACGGTTAGAAGAGGTGAGTTTGGGTCTCT					
CDC45-EX5B-3	CACAGCTATCCCAGAGGAACTTGCACTCCCAGAGGTCTAGATTGGATCTTGCTGGCAC					
CDC45-EX7-5	GGGTTCCCTAAGGGTTGGACTACGAGCAGTATGAATATCATGGGACATCGGTAAGTAT					
CDC45-EX7-3	GAATAGGTGGAACTCACTATAAAGTTCTGACTCCAGGGGTCTAGATTGGATCTTGCTGGCAC					
CDC45-EX5A-5	GGGTTCCCTAAGGGTTGGAGATGATGACCTTGAAGTTCCCGCCTATGAAGACATCTTCAG					
CDC45-EX5A-3	GGATGAAGAGGAGGATGAAGAGCATTCAGGAAATGACAGTGTCTAGATTGGATCTTGCTGGCAC					
CDC45-EX6-5	GGGTTCCCTAAGGGTTGGAGCTGTGGGGGATAAATTCCTGGTGCATTTGCTCCACCTTTTGTT					
CDC45-EX6-3	CTCTTTGTCCCTGTATCAGGAGATAGTGGAGCAAACCATGCGGTCTAGATTGGATCTTGCTGGCAC					

^aDNA was extracted from either venous blood collected into EDTA, lymphoblastoid cell lines or cultured fibroblasts established from skin biopsies obtained from the scalp incision at the time of surgical intervention. All DNA was extracted using the Nucleon Blood and Cell Culture (BACC) DNA extraction kit (Gen-Probe Inc.) according to the manufacturer's instructions. PCR amplification of DNA was performed in a volume of 20 μ l, containing 15 mM TrisHCl (pH 8.0), 50 mM KCl, 2.5 mM MgCl₂, 100 μ M each dNTP, 0.4 μ M primers, and 0.75 units of FastStart Taq DNA Polymerase (Roche). Cycling conditions comprised a 5 min initial denaturation step at 94°C, followed by 35 cycles of 94°C for 30 s, annealing at 63°C for 30 s and extension at 72°C for 30 s, with a final extension at 72°C for 7 min. For dideoxy-sequencing, amplified PCR products were treated with *ExoI* (NEB) and shrimp alkaline phosphatase (FastAP; Thermo Scientific) to remove PCR primers and dNTPs respectively, by incubating at 37°C for 30 min, followed by 85°C for 15 mins to denature the enzymes. Sequencing was then carried out using the BigDye Terminator v3.1 cycle sequencer system (Applied Biosystems).

^bPCR amplification of DNA was performed in a volume of 10 μ l, using ReddyMix PCR master mix with 1.5 mM MgCl₂ (Thermo Scientific), 0.5 μ M primers and 10 ng DNA. Cycling conditions employed a touchdown approach and were as follows: 5 min initial denaturation step at 95°C, then 3 cycles of 94°C for 30 s, 65°C for 30 s, 72°C for 45 s, then 3 cycles of 94°C for 30 s, 62°C for 30 s, 72°C for 45 s, then 3 cycles of 94°C for 30 s, 62°C for 30 s, 72°C for 45 s, then 3 cycles of 94°C for 30 s, 72°C for 45 s, then 3 cycles of 94°C for 30 s, 72°C for 45 s, then a final 35 cycles of 94°C for 30 s, 56°C for 30 s, 72°C for 45 s, then a final extension of 72°C for 10 min. Dideoxy-sequencing was performed as in ^a.

^cPrimer pairs were designed to amplify 25 target regions of approximately 200 bp, covering all 19 exons and intron/exon boundaries for use with the Access ArrayTM IFC system (Fluidigm). Universal CS1 and CS2 adaptor sequences were included on the 5' ends of all target-specific forward and reverse primers, respectively. Primers were multiplexed into pools of 4-5 optimised pairs, with a final concentration of 1 μ M per primer. Amplicon tagging, thermal cycling using the Biomark HDTM system (Fluidigm) and product harvesting were performed following the manufacturer's instructions. Ion PGMTM sequence-specific adaptors and sample indexes were attached to each amplicon pool for DNA extracted from 48 individual craniosynostosis patients using a DNA Engine Dyad® Peltier thermal cycler (Bio-Rad). Indexed PCR products were pooled, purified with AMPure XP beads (Beckman Coulter Ltd.) and quantified using an Agilent High Sensitivity DNA Kit with the 2100 Bioanalyzer (Agilent Technologies, Inc) or

the 2200 TapeStation (Agilent Technologies) with a High Sensitivity D1000 ScreenTape. In total, 467 patient samples were processed for downstream sequencing applications. Pooled and indexed PCR products were sequenced using the Ion PGMTM System (Life Technologies Ltd.). Pooled libraries were diluted to a final concentration of 10 pM and emulsion PCR and enrichment were performed with the Ion PGMTM Template OT2 200 Kit (Life Technologies Ltd.) according to the manufacturer's instructions. Enriched target regions were sequenced with the Ion PGMTM Sequencing 200 Kit v2 for 125 cycles, using an Ion 318 chip.

Underperforming amplicons (CDC45_2, CDC45_3 and CDC45_24) were re-sequenced independently. PCR products were amplified from 40 ng DNA with Q5[®] Hot Start High Fidelity polymerase (NEB) in the recommended standard reaction mixture, with the inclusion of Q5 High GC Enhancer. Cycling conditions were denaturation at 98°C for 30 s, followed by 35 cycles of 98°C for 10 s, 60°C for 30 s and 72°C for 30 s, followed by a final extension at 72°C for 2 min. Amplification products were diluted 100-fold and used in a second PCR reaction to incorporate Ion PGMTM sequence-specific adaptors and sample indexes as above, for 9 cycles. Pooled samples were purified and sequenced for 100 cycles as described above, using an Ion 314 chip. Sequence read alignment and variant analysis were performed using the Ion Torrent platform-specific software v4.2.1. In total 361 samples fulfilled minimum coverage parameters (x10 reads) for all amplicons.

^dThe MLPA reactions were performed according to a protocol based on the methods described previously.^{5, 6} Products were separated by capillary electrophoresis on the ABI 3130 (Applied Biosystems) and data analyzed using GeneMarker (SoftGenetics Inc). Threshold ratios for deletion and duplication were set at <0.75 and >1.3, respectively. The colored primer sequences are universal for the MLPA procedure.

Target Primers	Primer se	Product size (bp)	
	Forward	Reverse	
Exon 1-3	TCCGATTTCCGCAAAGAGTTCTACG	CAGCGTATATTGCACGTGGTCACA	138
Exon 3-4	GGTGGCAAGAACTTGAAACTG	ACATTGACGACATTGACTGGCCTAT	166
Exon 3-5	GGTGGCAAGAACTTGAAACTG	ATGTCTTCATAGGCGGGAAC	238/100 if exon 4 skipped
Exon 5-6	GTTCCCGCCTATGAAGACAT	CTCCTCCGCATGGTTTGCTC	137
Exon 2F-5R	GACGTGGATGCTCTGTGTGC	GCTTCTCAGAAGGCTCTGACC	394/301 if exon 3 skipped /256 if exon 4 skipped
Exon 2_3F-5_4R	AGATCCTTCAGGCCTTGTTCC	TAATTTGATCTGGGTATCGTTGTATAC	251
Exon 2_3F-5_3R	AGATCCTTCAGGCCTTGTTCC	TAATTTGATCTGTTCTTTATGCTCAAG	113
GAPDH	GGACTCCACGACGTACTCAGCGCCAGC	GTGGATATTGTTGCCATCAATGACC	213

Table S3. Primer sequences used for cDNA analysis of CDC45.

Patient(s)	Mutation	Allele	Sequence	MAXENT Score	Sum Difference	Sum Variation	Significance Threshold	Significant ^a	
Р3	a 652 + 5C > A	G (WT)	tcagtaa <mark>g</mark> g	8.61	5.05	-69%	-30%	Yes – 5´ donor site	
	c.035+5G>A	A (MUT)	tcagtaaag	2.66	5.95			lost	
P9-1 & P9-2	c.203A>G, p.Gln68Arg	A (WT)	c <mark>a</mark> ggtattg	8.35	2 49	-42%	200/	Yes – 5´ donor site lost	
		G (MUT)	cgggtattg	4.87	5.46		-30%		
P10	a 1440±14C≥ T	C (WT)	cagg <mark>c</mark> gggt	0.81	7 75	11570/	200/	Yes – cryptic 5' donor site created	
	c.1440+14C>1	T (MUT)	caggtgggt	8.56	-1.15	+1137%	+30%		

Table S4. In silico prediction of mutation effects on canonical splice sites.

^aSignificance based on thresholds applied by prediction software, Alamut Batch (Interactive Biosciences Inc) and Human Splicing Finder.⁷

Table S5. In silico splicing predictions for synonymous mutations in CDC45.

Patient(s)	Mutation	Motif affected	Predicted Consequence
P1	c.318C>T, p. V106V	Multiple potential silencing motifs created Exonic splice enhancer motif lost	Skipping of exon 4
P9-1 & P9-2	c.333C>T, p.N111N	SRp40 exonic splice enhancer motif lost	Skipping of exon 4

Predictions calculated using Human Splicing Finder.⁷

References

- 1. Li, H., and Durbin, R. (2009). Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics *25*, 1754-1760.
- 2. Lunter, G., and Goodson, M. (2011). Stampy: a statistical algorithm for sensitive and fast mapping of Illumina sequence reads. Genome Res. *21*, 936-939.
- 3. Rimmer, A., Phan, H., Mathieson, I., Iqbal, Z., Twigg, S.R., Consortium, W.G.S., Wilkie, A.O., McVean, G., and Lunter, G. (2014). Integrating mapping-, assembly- and haplotype-based approaches for calling variants in clinical sequencing applications. Nat. Genet. *46*, 912-918.
- 4. Wang, K., Li, M., and Hakonarson, H. (2010). ANNOVAR: functional annotation of genetic variants from high-throughput sequencing data. Nucleic Acids Res. *38*, e164.
- Schouten, J.P., McElgunn, C.J., Waaijer, R., Zwijnenburg, D., Diepvens, F., and Pals, G. (2002). Relative quantification of 40 nucleic acid sequences by multiplex ligation-dependent probe amplification. Nucleic. Acids. Res. *30*, e57.
- White, S.J., Vink, G.R., Kriek, M., Wuyts, W., Schouten, J., Bakker, B., Breuning, M.H., and den Dunnen, J.T. (2004). Two-color multiplex ligation-dependent probe amplification: detecting genomic rearrangements in hereditary multiple exostoses. Hum. Mutat. 24, 86-92.
- Desmet, F.O., Hamroun, D., Lalande, M., Collod-Beroud, G., Claustres, M., and Beroud, C. (2009). Human Splicing Finder: an online bioinformatics tool to predict splicing signals. Nucleic Acids Res. 37, e67.