

De Novo Mutations Affecting the Catalytic C α Subunit of PP2A, PPP2CA, Cause Syndromic Intellectual Disability Resembling Other PP2A-Related Neurodevelopmental Disorders

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Type 2A protein phosphatases (PP2As) are highly expressed in the brain and regulate neuronal signaling by catalyzing phospho-Ser/Thr dephosphorylations in diverse substrates. PP2A holoenzymes comprise catalytic C-, scaffolding A-, and regulatory B-type subunits, which determine substrate specificity and physiological function. Interestingly, *de novo* mutations in genes encoding A- and B-type subunits have recently been implicated in intellectual disability (ID) and developmental delay (DD). We now report 16 individuals with mild to profound ID and DD and a *de novo* mutation in *PPP2CA*, encoding the catalytic C α subunit. Other frequently observed features were severe language delay (71%), hypotonia (69%), epilepsy (63%), and brain abnormalities such as ventriculomegaly and a small corpus callosum (67%). Behavioral problems, including autism spectrum disorders, were reported in 47% of individuals, and three individuals had a congenital heart defect. *PPP2CA de novo* mutations included a partial gene deletion, a frameshift, three nonsense mutations, a single amino acid duplication, a recurrent mutation, and eight non-recurrent missense mutations. Functional studies showed complete PP2A dysfunction in four individuals with seemingly milder ID, hinting at haploinsufficiency. Ten other individuals showed mutation-specific biochemical distortions, including poor expression, altered binding to the A subunit and specific B-type subunits, and impaired phosphatase activity and C-terminal methylation. Four were suspected to have a dominant-negative mechanism, which correlated with severe ID. Two missense variants affecting the same residue largely behaved as wild-type in our functional assays. Overall, we found that pathogenic *PPP2CA* variants impair PP2A-B56(δ) functionality, suggesting that PP2A-related neurodevelopmental disorders constitute functionally converging ID syndromes.

Introduction

Neurodevelopmental disorders (NDDs) represent a collection of clinically heterogeneous disorders—including intellectual disability (ID), epilepsy, and various behavioral disor-

ders—and can manifest as an isolated cognitive defect or in combination with other comorbidities, such as congenital abnormalities.^{1,2} Until recently, identifying the genetic etiology of NDDs remained challenging because of their extensive clinical and genetic heterogeneity. Recent advances

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in whole-exome sequencing (WES) and whole-genome sequencing have highlighted *de novo* mutations as a major cause of ID,^{2–9} resulting in an increase in new clinical NDD entities associated with ID and developmental delay (DD).¹

Recently, recurrent *de novo* mutations in genes encoding specific subunits of type 2A protein phosphatases (PP2As) have been reported,^{3,4,10–12} establishing this phosphatase family as a novel and important player in the etiology of ID and DD. PP2A enzymes catalyze dephosphorylation of phospho-Ser and phospho-Thr residues in a large variety of substrates, thereby counterbalancing Ser/Thr-specific protein kinases and thus playing essential regulatory roles in cellular signaling.¹³ Structurally, PP2As are holoenzymes comprising three subunits: a catalytic C-type, a scaffolding A-type, and a regulatory B-type subunit, the latter of which is of major importance in determining physiological functions, substrate specificity, and regulation of the complex.^{14–16} PP2A complexes are encoded by 19 different genes, and mutations in three of these genes—*PPP2R1A* (MIM: 605983; subunit A α), *PPP2R5C* (MIM: 601645; subunit B56 γ), and *PPP2R5D* (MIM: 601646; subunit B56 δ)^{3,4,10–12}—have been conclusively identified as causes of ID and DD syndromes; for two others—*PPP2R5B* (MIM: 601644; subunit B56 β)¹¹ and *PPP2R2C* (MIM: 605997; subunit B55 γ)—variants in individuals with ID and DD have been observed.¹⁷

So far, detailed studies on *de novo* mutations in both genes encoding the C subunit (*PPP2CA* and *PPP2CB*) have not been reported. Yet, ID- and DD-related mutations in genes involved in C subunit regulation, such as *IGBP1* (MIM: 300472),¹⁸ *MID1* (MIM: 300552),¹⁹ and *BOD1* (MIM: 616745),^{20,21} have been identified. Together with mutations in other PP2A subunits, these observations make the genes encoding the C subunit themselves excellent candidates for ID and DD disorders. Here, we report that *de novo* mutations in *PPP2CA*, encoding the PP2A catalytic C α subunit, cause an ID and DD syndrome phenotypically characterized by mild to severe ID and DD, behavioral problems, variable types of epilepsy, hypotonia, and structural brain abnormalities. In addition, we provide functional evidence for a mutation-specific pathophysiological mechanism resulting in at least a common impairment of PP2A-B56 δ holoenzyme functionality. Our data further highlight the importance of PP2A function in neurodevelopment.

Material and Methods

Generation of *PPP2CA* Mutants

To study the functional consequences of *de novo* missense mutations, we cloned the coding region of wild-type (WT) C α cDNA into an N-terminally HA-tagged eukaryotic expression vector (pMB001) by using XbaI and BamHI sites. Mutated C α constructs were directly generated from this pMB001 vector through PCR-based site-directed mutagenesis (Stratagene) with proofreading Pwo polymerase (Roche Applied Science) and complementary DNA oligonucleotide primers (Sigma Genosys) containing the desired point mutations (primer sequences in Table S1). To generate the C α p.Phe308dup variant, we directly incorporated

the mutation into the reverse C α stop primer. We cloned the coding region, PCR-amplified with a C α start primer and this mutated stop primer (primer sequences in Table S1), into pMB001 by using XbaI and BamHI restriction sites. All mutations were confirmed by sequencing (LGC Genomics).

Cellular PP2A Binding Assays

HEK293T cells were transfected with PEI transfection reagent according to a standard protocol. 48 hr after transfection, cells were rinsed with PBS, lysed in 200 μ L NET buffer (50 mM Tris HCl [pH 7.4], 150 mM NaCl, 15 mM EDTA, and 1% Nonidet P-40) containing protease and a phosphatase inhibitor cocktail (Roche Applied Science), and centrifuged for 15 min at 13,000 \times *g*. If the experiment required measuring of phosphatase activity, TBS was used instead of PBS, and phosphatase inhibitors were omitted from the lysis buffer.

For pull-down experiments, cell lysates were incubated at 4°C for 1 hr with 500 μ L NENT100 buffer (20 mM Tris HCl [pH 7.4], 1 mM EDTA, 0.1% Nonidet P-40, 25% glycerol, and 100 mM NaCl) containing 1 mg/mL bovine serum albumin (BSA) and 30 μ L monoclonal anti-HA-Agarose antibody beads (Sigma-Aldrich; for HA pull-down), 30 μ L glutathione-Sepharose beads (GE Healthcare; for GST pull-down), or 30 μ L GFP-trap-A beads (ChromoTek; for GFP trapping) on a rotating wheel. The beads were washed twice with 0.5 mL NENT100 containing 1 mg/mL BSA and twice with 0.5 mL NENT300 containing 300 mM NaCl.

In all cases, bound proteins were eluted by the addition of 2 \times NuPage sample buffer (Invitrogen) and boiling. The eluted proteins were subsequently analyzed by SDS-PAGE on 4%–12% (w/v) Bis-Tris gels (Bio-Rad) and immunoblotting. The membranes were blocked in 5% milk solution in TBS and 0.1% Tween-20 for 1 hr at room temperature (RT) and subsequently incubated with the primary antibody overnight at 4°C. The following primary mouse antibodies were used: anti-HA (Sigma-Aldrich), anti-GST (Sigma-Aldrich), anti-GFP (generously supplied by Dr. P. Parker at the Francis Crick Institute, London, UK), anti-PP2A-A subunit (generously supplied by Dr. S. Dilworth at Middlesex University, London, UK), anti-demethyl-PP2A clone 1D7 (BioLegend), anti-FLAG M2 (Sigma-Aldrich), anti-vinculin (Sigma-Aldrich), and anti-alpha4 clone 5F6 (Millipore).

After being washed in TBS and 0.1% Tween-20, the membranes were incubated at RT for 1 hr with horseradish-peroxidase-conjugated secondary antibodies (Dako) and developed on an ImageQuant LAS 4000 system (GE Healthcare) with a WesternBright ECL detection kit (Advansta). All densitometric quantifications were done with ImageJ software.

HA Peptide Elution

For binding experiments with the HA-tagged p.Arg214* C α mutant, anti-HA-agarose-bound proteins were eluted for 30 min at 4°C after the final washing step by the addition of HA peptide (Sigma-Aldrich) at 200 ng/ μ L. The remaining beads were spun down, and the eluate was transferred for further use.

PP2A Activity Assays

After HA pull-down, beads were washed once more with 20 mM Tris HCl (pH 7.4) and 1 mM DTT (Tris-DTT) and finally resuspended in 80 μ L Tris-DTT solution. All assays were performed with 20 μ L of this “phosphatase suspension” and 9 μ L of 2 mM stock of K-R-pT-I-R-R phosphopeptide for 20–40 min at 30°C (still in the linear range of the assay). The released free phosphate was

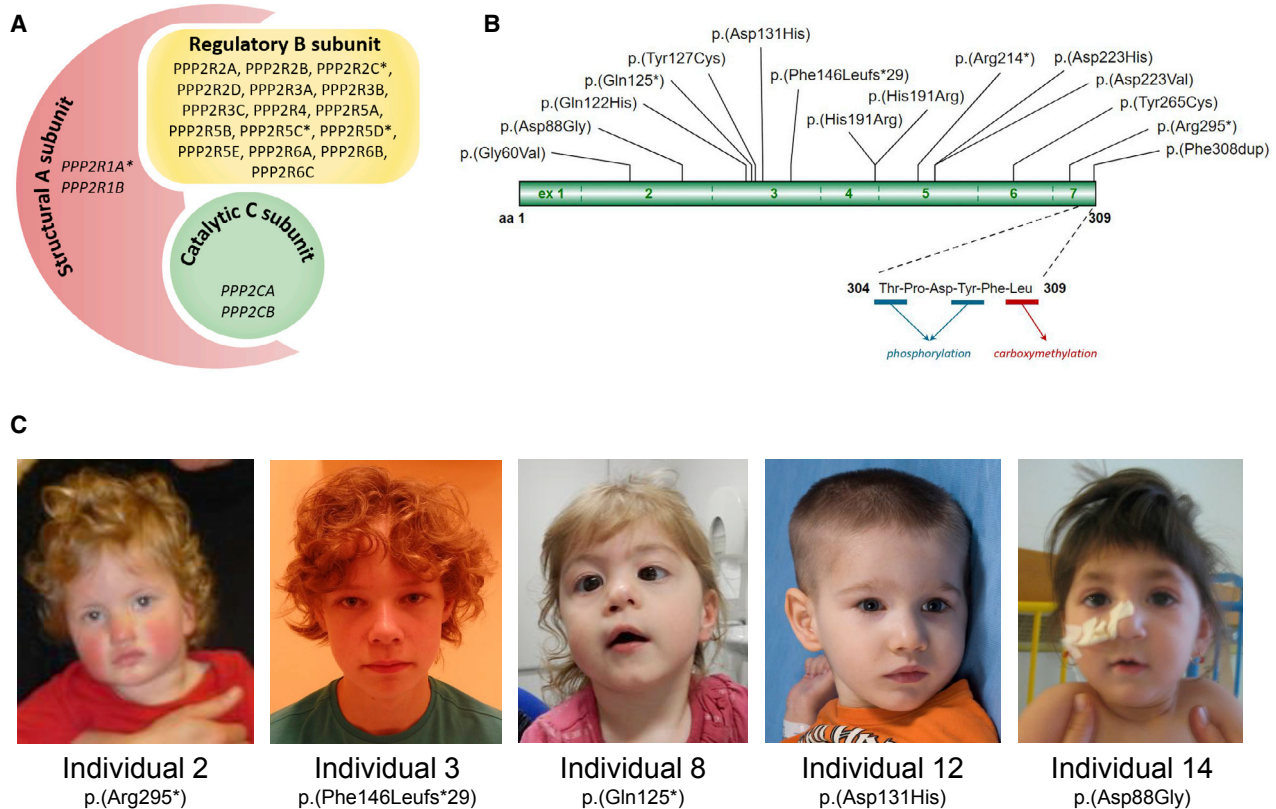


Figure 1. Schematic Overview of the PP2A Complex, Variants in the PP2A Catalytic C α Subunit, and Photographs of Individuals
 (A) Overview of the PP2A complex based on the canonical subunits. Gene names encoding the subunit components are listed in each subunit. *PPP2CA* encodes the catalytic subunit alpha, highlighted in green. Subunits known for their involvement in ID and DD are annotated with an asterisk.
 (B) *De novo* mutations identified in individuals with ID and DD are shown with respect to their exonic and C α protein localization; exon notation is indicated in the ribbon protein structure. The C-tail sequence involved in carboxymethylation and phosphorylation is enlarged for visualization purposes. Note that the partial deletion is not depicted.
 (C) Facial features of individuals with a *de novo* mutation in *PPP2CA*. Some individuals had mild dysmorphisms, but a consistent facial gestalt was not observed. Informed consent was provided by the parents.

determined by the addition of BIOMOL Green (catalog no. BML-AK111-0250, Enzo). After 20 min of incubation at RT, absorbance at 620 nm was measured in a multi-channel spectrophotometer. We subsequently obtained specific phosphatase activity by correcting the measured absorbance for the input of HA-tagged C α , as determined by immunoblotting with anti-HA antibodies and signal quantification by ImageJ.

Statistics

Statistical analysis of biochemical data was assessed with one-sample Student's *t* tests in which the data were compared with WT values that were set at 100% in each experimental replicate. *p* values below 0.05 were considered significant.

Results

Clinical Spectrum Associated with *PPP2CA* Mutations

Through international collaboration,^{2,22} we recruited 16 individuals with a *de novo* mutation in *PPP2CA* (Figures 1A and 1B). This study was approved by the local institutes under the realm of diagnostic testing. Detailed clinical information is provided in the [Supplemental Note](#) and sum-

marized in [Table 1](#). Individuals were 1.5–23 years old and had mild to profound ID and DD, and ten had severe language impairment that was worse than expected in several individuals given their level of development. Behavioral problems, of which autism spectrum disorders (ASDs) were most frequently diagnosed, were noted in 7 of 15 individuals (47%). One individual (individual 7) also had psychotic episodes after which she had progressive cognitive dysfunction and worsening behavioral problems. Regression of development was also noted in individual 3. Ten individuals (63%) had seizures, albeit in different forms: two showed fever-related seizures, one was diagnosed with epileptic encephalopathy, two had focal seizures (one of whom was diagnosed with focal electrical status epilepticus during sleep), two had generalized tonic-clonic seizures, one was diagnosed with Jeavons syndrome, one had hypsarrhythmia, and one had an unknown type of seizures. Brain imaging was performed in 15 individuals and showed dilated ventricles (4/15), delayed myelination (2/15), and dysplasia or absence of the corpus callosum (4/15). Two individuals had macrocephaly, one of whom was also tall, and five individuals had

Table 1. Main Clinical Features of Individuals

General Information	Individual 1	Individual 2	Individual 3	Individual 4	Individual 5	Individual 6	Individual 7	Individual 8
Age	6 y	7 y	13 y	3 y, 10 m	19 y	2 y, 4 m	23 y	2 y, 1 m
Gender	male	female	male	male	female	female	female	Female
Chr change (hg19)	g.133536680T>C	g.133533511dup	g.133537587del	g.133533469_133533471dup	g.133536124G>A	g.133536680T>C	g.133536096T>A	g.133537652G>A
GenBank: NM_002715.2	c.572A>G	c.882dup	c.438del	c.922_924dup	c.640 C>T	c.572A>G	c.668A>T	c.373C>T
Protein change	p.His191Arg	p.Arg295*	p.Phe146Leufs*29	p.Phe308dup	p.Arg214*	p.His191Arg	p.Asp223Val	p.Gln125*
Mutation type	missense	nonsense	frameshift	1 amino acid insertion	nonsense	missense	missense	nonsense
Inheritance	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>
Other variants? ^a	–	+	–	–	+	–	+	+
Proposed mechanism	haploinsufficiency of PP2A-B56δ complexes	dominant-negative effect on PP2A-B56δ complexes	haploinsufficiency: null allele (no expression)	dominant-negative effect on all PP2A-B56 complexes	haploinsufficiency: null allele (latent complex with alpha 4)	haploinsufficiency of PP2A-B56δ complexes	unclear	haploinsufficiency: null allele (no expression)
Growth								
Height	+2 SD	+2.5 SD	+1.3 SD	+1.7 SD	–0.2 SD	0 SD	+0.7 SD	–1.3 SD
Head circumference	+2.5 SD	+0.5 SD	–1.5 SD	–3.9 SD	NK	–0.2 SD	+2.8 SD	–0.8 SD
Weight	–0.2 SD	0 SD	–1.1 SD	+0.7 SD	>+2.5 SD	NK	NK	–0.6 SD
Neurological								
DD and ID (degree)	+ (mild)	+ (severe)	+ (mild)	+ (moderate)	+ (mild to moderate)	+ (mild)	+ (mild)	+ (mild)
Severe language delay	+	+	+	+	+	NK	–	–
Regression	–	–	+	–	–	–	+ (after period with psychoses)	–
Behavioral problems	–	+ (automutilation, stereotypic behavior)	+ (PDD-NOS)	–	+ (ASD, ADHD)	NK	+ (ASD, psychoses)	–
Hypotonia	–	+	+	–	+	–	+ (mild)	+
Epilepsy (type)	+ (related to fever)	+ (tonic-clonic seizures, absences)	+ (focal left temporal, mostly at night)	–	+ (generalized tonic-clonic seizures, absences)	+ (related to fever, tonic-clonic seizures)	–	–
Epilepsy syndrome	NA	epileptic encephalopathy	focal ESES	NA	Jeavons syndrome	NA	NA	NA
Age of onset	11 m	6 m	10 y	NA	6 y	NK	NA	NA
Age of offset	18 m	ongoing	ongoing	NA	ongoing	NK	NA	NA
Brain abnormalities	+ (ventriculomegaly)	+ (ventriculomegaly, gracile corpus callosum, delayed myelination)	–	+ (posterior hypoplasia of corpus callosum, unmyelinated left temporal lobe, nonspecific periventricular white-matter hyperintensities)	–	+ (mildly underdeveloped pons, mesencephalon, mild dilated lateral and third ventricles)	–	+ (reduced white matter [supratentorial and infratentorial], some global atrophy of the brain [pons and corpus callosum], bilateral plexus choroideus cysts)
Other	–	sleeping difficulties	–	–	burning pain from neck down to left leg	–	–	broad-based gait

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Individual 9	Individual 10	Individual 11	Individual 12	Individual 13	Individual 14	Individual 15	Individual 16	Total (Percentage)
1 y, 9 m	8 y	2 y, 4 m	11 y	12 m	4 y	3 y, 4 m	7 y	NA
male	female	male	male	male	female	male	male	9 males, 7 females
g.133537645T>C	g.133534840T>C	deletion at chr5: 133,546,961–133,667,321	g.133537634C>G	g.133537659C>G	g.133541662T>C	g.133541746C>A	g.133536097C>G	NA
c.380A>G	c.794A>G	NA	c.391G>C	c.366G>C	c.263A>G	c.179G>T	c.667G>C	NA
p.Tyr127Cys	p.Tyr265Cys	NA	p.Asp131His	p.Gln122His	p.Asp88Gly	p.Gly60Val	p.Asp223His	NA
missense	missense	partial deletion	missense	missense	missense	missense	missense	NA
<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	<i>de novo</i>	NA
–	–	+	–	+	–	–	–	NA
haploinsufficiency of PP2A-B56 and PR72 complexes; dominant-negative effect on PP2A-B55 and PP2A-STRN complexes	haploinsufficiency: significant impairment of any trimer formation	haploinsufficiency (deletion)	haploinsufficiency of PP2A-B56 complexes	haploinsufficiency?	dominant-negative effect on PP2A-B56 γ and B56 δ complexes	haploinsufficiency: poor expression	unclear	NA
+0.4 SD	–2 SD	–3 SD	–1.6 SD	–1.9 SD	–0.8 SD	0 SD	–2 SD	NA
–3 SD	–2.3 SD	–3 SD	–1.1 SD	–4.6 SD	<–2 SD	+1.2 SD	0 SD	NA
–0.2 SD	–1.5 SD	–2 SD	–2.1 SD	–1.9 SD	–0.6	–2 SD	–1.5 SD	NA
+ (severe)	+ (mild)	+ (mild)	+ (profound)	+ (moderate)	+ (severe)	+ (mild)	+ (severe)	16/16 (100%)
+	+	–	+	NA	+	–	+	10/14 (71%)
–	–	–	–	–	–	–	–	2/16 (13%)
–	+ (ASD, ADHD)	–	–	–	–	+	+ (ASD)	7/15 (47%)
+	+	–	+	+	+	–	+ (mild)	11/16 (69%)
–	+ (NK)	–	+ (staring spells, head drops, and tonic-clonic seizures)	+ (focal epilepsy with secondary generalization)	+ (generalized tonic-clonic seizures)	–	+ (generalized tonic-clonic seizures)	10/16 (63%)
NA	NK	NA	hypsarrhythmia (West syndrome)	not specified	not specified	NA	not specified	NA
NA	3 y	NA	8 m	5 m	1 w	NA	1 y	NA
NA	ongoing	NA	ongoing	ongoing	ongoing	NA	ongoing	NA
+ (pontocerebellar hypoplasia, mild ventriculomegaly)	+ (nonspecific thinning of the corpus callosum, mild volume loss of the cerebellum)	NK	–	+ (microcephaly, enlarged subarachnoid spaces, nonspecific findings of cerebral underdevelopment)	+ (slightly dilated external and internal subarachnoid spaces, diffuse but discrete atrophy, adequate myelination)	–	+ (bilateral dilatation of perivascular spaces along the corona radiata)	10/15 (67%)
optic-nerve anomaly	NK	–	choreiform movements	–	stereotypic movements	–	–	6/15 (40%)

(Continued on next page)

Table 1. Continued

General Information	Individual 1	Individual 2	Individual 3	Individual 4	Individual 5	Individual 6	Individual 7	Individual 8
Facial								
Eyes	hypertelorism, prominent eyes	–	periorbital fullness restricted to upper eyelids, long eyelashes	ptosis, short palpebral fissures	–	prominent eyes, epicanthic folds	small palpebral fissures, deep-set eyes	megalocornea
Nose	full nasal tip, low-hanging columella	broad nasal bridge	full nasal tip	broad nasal tip	–	–	broad nasal tip	asymmetrical nostrils and bifid nasal tip
Philtrum	short philtrum	short philtrum	–	short philtrum	–	–	–	–
Other dysmorphic features	broad forehead, frontal bossing, square shape of both ears	open fontanel, plagiocephaly, some frontal bossing, prominent upper lip, sacral dimple	–	trigonocephaly, double hair whorl, high palate, low-set and anteverted ears, retrognathia	crowded teeth	plagiocephaly	broad forehead, freckling around the mouth	plagiocephaly
Other								
Extremities	single left palmar crease, long fingers and toes, bilateral sandal gap	bilateral fetal finger pads, single left palmar crease	–	poorly developed flexure fold on the third right finger, incomplete single transverse palmar crease, adducted thumbs (reducible); X-ray: ovoid appearance of vertebrae, coxa valga, slender distal phalanges	hypermobile knees	mild shortened fifth digits	–	hyperlaxity
Feeding difficulties	+	+	+	–	–	–	–	–
Vision problems	–	–	–	–	periods of sudden vision loss and pain in left eye	megalocornea, moderated excavated papillae, mild hypermetropia (both eyes +2)	NK	megalocornea, strabismus
Other	4 café-au-lait spots (max 5 mm)	–	vitamin B12 deficiency	elevated right diaphragm	recurrent ear and urinary-tract infections, heart murmur, fractures after minimal trauma	diastasis recti	frequent airway infections as a child	muscular ventricular septal defect

Abbreviations are as follows: +, present; –, not present; w, weeks; y, years; m, months; NK, not known; NA, not applicable; SD, standard deviation; DD, developmental delay; ID, intellectual disability; PDD-NOS, pervasive developmental disorder not otherwise specified; ASD, autism spectrum disorder; ADHD, attention deficit hyperactivity disorder; ESES, electrical status epilepticus during sleep; CVI, cerebral visual impairment.

^aAdditional information on variant(s) in other gene(s) is in the [Supplemental Note](#).

microcephaly. A recognizable facial gestalt could not be observed, but some individuals showed mild dysmorphic features, such as a full nasal tip and megalocornea, and one individual had an asymmetrical face (Figure 1C). Other clinical observations included hypotonia (11/16 [69%]), feeding difficulties (9/15 [60%]), vision problems (6/14 [43%]), and congenital heart defects, the last of which included a muscular ventricular septal defect, atrial septal defect, and trileaflet aortic valve in three separate individuals.

Mutational Spectrum Is Diverse and Suggests Loss of PPP2CA Subunit Functioning

In total, 15 different *de novo* mutations in 16 individuals were identified through routine clinical diagnostic proced-

ures, including WES (15 individuals) and microarray-based copy-number analysis (one individual; Figure 1B). The latter detected a 120 kb deletion (chr5: 133,546,961–133,667,321; GRCh37) including *PPP2CA* exon 1 and the 3' coding region of *CDKL3* in individual 11. The variants identified by WES included a recurrent missense mutation, c.572A>G (p.His191Arg) (which is predicted to be deleterious by MutationTaster²³ and affects a highly conserved nucleotide and amino acid position in the “loop switch” of PP2A-C²⁴) in individuals 1 and 6. The conformation of the loop switch determines the conformation of the active site and the binding of catalytic metal ions.²⁴ In addition, His191 is one of four residues forming a hydrophobic cage accommodating binding of pharmacologic PP2A inhibitors, such as okadaic acid and microcystin.²⁵ Gln122

Individual 9	Individual 10	Individual 11	Individual 12	Individual 13	Individual 14	Individual 15	Individual 16	Total (Percentage)
bilateral epicanthal folds	–	periorbital fullness, upslanting palpebral fissures	–	–	almond-shaped eyes	–	–	NA
–	–	–	–	–	–	broad nasal tip, small alae nasi	–	NA
–	–	–	–	–	–	–	–	NA
prominent metopic suture, mild micrognathia	–	full cheeks, thin upper lip, everted lower lip, square configuration of the lobule of the ear (right bigger than left)	high arched palate	–	small mouth, high palate	–	–	NA
bridged palmar crease, bilateral ulnar deviation of wrists and fingers, partially adducted thumbs, right hand greater than left	NK	bilateral fifth-finger clinodactyly	hyperlaxity	bilateral single palmar crease, bilateral adducted thumb	atypical single left palmar crease, pes planus, overlapping toes	single left palmar crease	–	NA
+	NK	+	+	+	+	+(infancy)	–	9/15 (60%)
CVI	NK	immature retina at the age of 1 m	does not track or blink to threat	–	–	–	–	6/14 (43%)
mild bilateral sensorineural hearing loss	NK	inguinal hernia, umbilical hernia	constipation, atrial septal defect, low bone mineral density	trileaflet aortic valve, multiple café-au-lait macules	–	sacral haemangioma	–	NA

is another of those four residues²⁵ and was identified to be substituted into a histidine in individual 13 (p.Gln122His). Two other missense mutations, c.380A>G (p.Tyr127Cys) and c.794A>G (p.Tyr265Cys), affect two tyrosine residues that are within the active site and are involved in direct contacts with okadaic acid and microcystin (Tyr265)²⁵ and with major PP2A cellular activators, such as PTPA (Tyr127 and Tyr265)²⁶ and the PP2A methyltransferase LCMT-1 (Tyr127).²⁷ The c.391G>C (p.Asp131His) variant harbors a charge reversal in an amino acid residing nearby the “helix switch” that is directly involved in binding to LCMT-1²⁷ and the PP2A inhibitor TIPRL1.²⁸ The c.179G>T (p.Gly60Val) and c.263A>G (p.Asp88Gly) variants are in close proximity to amino acids Asp57, His59, and Asp85, which are directly involved in the binding of catalytic metal ions in the active site of PP2A-C.²⁵

Moreover, His59 is involved in direct binding to okadaic acid and microcystin,²⁵ LCMT-1,²⁷ and PTPA.²⁶ The two remaining missense variants, 667G>C (p.Asp223His) and c.668A>T (p.Asp223Val), affect an amino acid of unknown functional significance.

If not degraded by nonsense-mediated RNA decay, the frameshift c.438del (p.Phe146Leufs*29) and two of the nonsense mutations, c.373C>T (p.Gln125*) and c.640C>T (p.Arg214*), are predicted to result in severely C-terminally truncated proteins. The remaining nonsense mutation c.882dup (p.Arg295*) is located in the ultimate exon and is predicted to result in a small, 14 amino acid truncation of the C α C-terminal tail. Importantly, this highly conserved C tail represents a focal point of PP2A regulation²⁹ and is reversibly modified by phosphorylation³⁰ and carboxymethylation^{31–34} within the six most

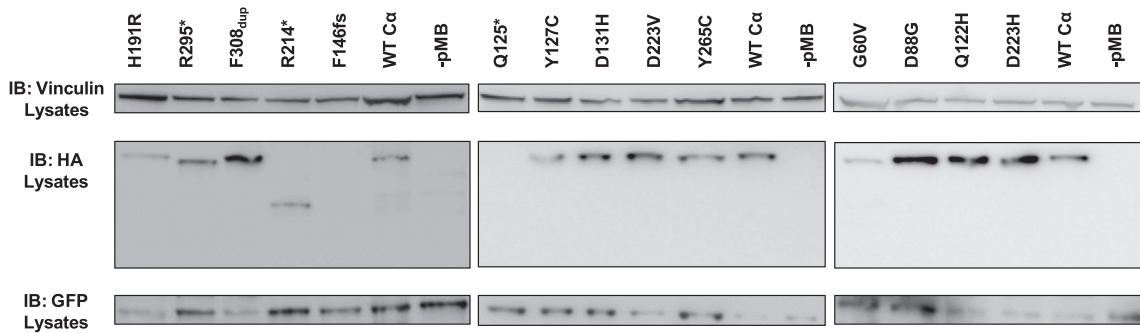


Figure 2. Levels of Mutant PP2A C α Subunits in HEK293T Cells

HA-tagged WT and mutant C α proteins were co-expressed with GFP (transfection control) in HEK293T cells. Protein levels were monitored in total protein lysates by immunoblotting with anti-HA and anti-GFP antibodies. Vinculin was used as a loading control. Abbreviated amino acid changes are as follows: H191R, p.His191Arg; R295*, p.Arg295*; F308dup, p.Phe308dup; R214*, p.Arg214*; F146fs, p.Phe146fs; Q125*, p.Gln125*; Y127C, p.Tyr127Cys; D131H, p.Asp131His; D223V, p.Asp223Val; Y265C, p.Tyr265Cys; G60V, p.Gly60Val; D88G, p.Asp88Gly; Q122H, p.Gln122His; and D223H, p.Asp223His.

C-terminal residues (304-T-P-D-Y-F-L-309), affecting PP2A holoenzyme assembly and activity (Figure 1B).^{35–37} Interestingly, individual 4 harbors a duplication of the phenylalanine in this sequence (T-P-D-Y-F-F-L), rendering the C-tail 1 amino acid longer and possibly hindering PP2A assembly and/or activation (Figure 1B).

Population metrics based on large databases (such as the Exome Aggregation Consortium [ExAC] Browser) indicate that, after sequence context and mutability are considered, *PPP2CA* is depleted of loss-of-function (LoF) variants according to multiple LoF metrics (pLI = 0.99; Z score = 4.47). Whereas these population signatures cannot be considered evidence of causality on their own, they support a hypothesis that *PPP2CA* variants leading to a functional loss are under strong purifying selection in the human population and that their occurrence could contribute to clinical phenotypes.

Both Haploinsufficiency and Non-haploinsufficiency Mechanisms Might Underlie the ID and DD Syndrome Caused by *De Novo* Variants in *PPP2CA*

We next set out to analyze both the catalytic capacity and the subunit binding of the *de novo* mutations affecting *PPP2CA*. We first attempted to determine protein levels of 14 different *de novo* mutants (one frameshift, three nonsense, one in-frame duplication, and nine missense mutants) in HEK293T cells, a frequently used model for PP2A biochemistry studies. We co-expressed the N-terminally HA-tagged C α mutants with GFP to control for transfection efficiency. Anti-HA immunoblotting using total protein lysates of the transfected cells consistently showed the presence of protein for 12 of 14 constructs (Figure 2); that is, despite the clear presence of GFP in all conditions tested, the p.Gln125* and p.Phe146Leufs*29 mutants failed detection (Figure 2), suggesting that these mutants might not be stable. Thus, we concluded that for the p.Gln125* and p.Phe146Leufs*29 variants, haploinsufficiency is the likely underlying pathophysiological mechanism. In addition, a persistent lower protein level was seen

for p.Gly60Val, leading to an inefficacy in generating reliable biochemical data. This could imply that this mutant is not stable, prompting us to advocate for a haploinsufficient mechanism in this case as well. Given the clear protein levels of all other variants examined, a dominant mechanism remained possible, and we further investigated this in additional functional assays.

PPP2CA Mutations Severely Compromise Phosphatase Activity in Most Cases

To determine intrinsic catalytic activity of the 11 C α mutants that resulted in stable protein levels, we isolated the proteins from total lysates of transfected HEK293T cells by pull-down on HA-agarose beads and washed and incubated them with a short phospho-peptide (K-R-pT-I-R-R) for varying times (20–40 min) at 30°C, after which we used BIOMOL Green to measure the release of free phosphate. An inactive C α mutant (p.Asp85Asn) was included as a negative control.³⁴ Whereas essentially no phosphatase activity was found for p.Asp85Asn, the activity of p.Gln122His, p.His191Arg, p.Asp223His, and p.Arg295* was severely decreased to 30%–50% of WT activity (Figure 3 and Table 2). In addition, PP2A activity was virtually absent for the p.Asp88Gly, p.Tyr127Cys, p.Arg214*, p.Tyr265Cys, and p.Phe308dup mutants (Figure 3). Together with the absence of the p.Gln125* and p.Phe146Leufs*29 variants and the low levels of p.Gly60Val, these results further suggest that the underlying pathophysiological mechanism is a functional loss.

On the other hand, p.Asp131His and p.Asp223Val variants displayed similar or even higher activity, indicating a divergent mechanism in these cases.

Carboxymethylation Assay Further Suggests PP2A Functional Defects

To further underscore the above activity data, we addressed potential changes in carboxymethylation of the C α mutants by using commercially available demethyl-specific antibodies. It has previously been demonstrated that

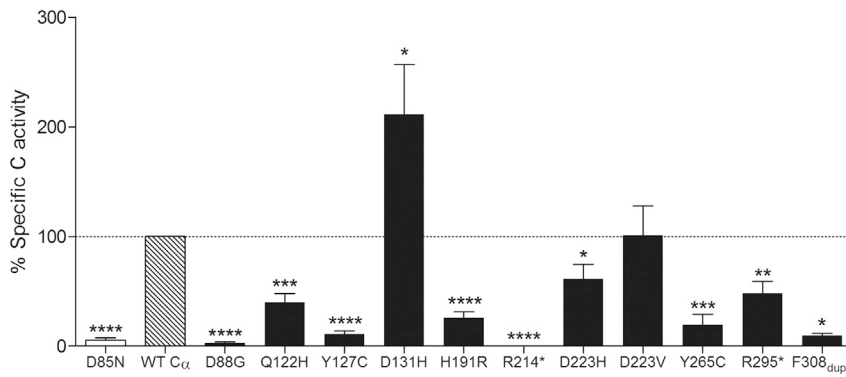


Figure 3. Phosphatase Activity of Mutant PP2A C α Subunits

HA-tagged WT and mutant C α proteins were purified from transfected HEK293T cells by HA pull-down, and absolute PP2A activity was determined on the K-R-pT-I-R-R phosphopeptide substrate with BIOMOL Green. Specific PP2A activities were calculated via correction of the measured activities for actual C α inputs, determined by anti-HA immunoblotting. Results represent the average specific activity \pm SEM for a given C α mutant in relation to the specific activity of WT C α (set at 100% in each experiment), as determined in at least three independent experiments ($n \geq 3$). A one-sample t test (compare to 100%) was used for assessing statistical significance (* $p \leq 0.05$, ** $p \leq 0.01$, *** $p \leq 0.001$, **** $p \leq 0.0001$).

loss of the single C-terminal Leu309 residue is already sufficient to prevent methylation.^{27,35} Hence, it is reasonable to predict that methylation of the truncated p.Arg214* and p.Arg295* mutants is no longer possible. In contrast, predicting a potential methylation defect of the p.Asp88Gly, p.Gln122His, p.Tyr127Cys, p.Asp131His, p.His191Arg, p.Asp223His, p.Asp223Val, p.Tyr265Cys, and p.Phe308dup mutants is more difficult. We therefore introduced the latter mutants, as well as WT C α and the p.Arg295* mutant, as HA-tagged fusion proteins and subsequently assessed their methylation status in HA pull-downs.

Anti-demethyl immunoblots indicated that the p.Asp88Gly, p.Gln122His, p.Tyr127Cys, p.Asp223Val, and p.Tyr265Cys variants had lower carboxymethylation ($p = 0.001, 0.03, 0.02, 0.03, \text{ and } 0.02$, respectively) than WT C α (Figures 4A and 4B and Table 2), whereas the p.His191Arg variant showed a trend toward increased demethylation ($p = 0.095$). For the p.Asp131His and p.Asp223His variants, however, there were no significant differences ($p > 0.15$). The decrease in methylation was most pronounced for the p.Asp88Gly, p.Tyr127Cys, and p.Tyr265Cys mutants, in which the mutated residues are part of the catalytic center and are in close or direct contact with the PP2A methyltransferase LCMT-1.²⁷ The observed PP2A activity defects in these variants (Figure 3) are in accordance with their inability to be properly methylated, given that effective C subunit carboxymethylation requires an active conformation of PP2A-C.²⁷

For the p.Phe308dup variant, the anti-demethyl antibodies revealed no signal at all, as was the case for the C tail lacking the p.Arg295* variant, suggesting that the antibodies are probably unable to recognize this mutant (Figure 4A). Hence, we could draw no conclusions regarding potential methylation defects of the variant with a 1 amino acid extension of the C tail. However, given the severe effect on activity (Figure 3), we speculate that this mutant might have impaired methylation as well.

PPP2CA Mutations Alter PP2A Subunit Binding in a Mutation-Specific Manner

To investigate whether the PPP2CA missense mutations affect PP2A subunit interactions, we first assessed binding of the HA-tagged C α mutants to the endogenous A subunit. After HA pull-down of HA-tagged C α mutants introduced in HEK293T cells, the presence of the A subunit in the bead fractions was monitored by anti-A immunoblotting. Whereas A subunit binding for the p.Gln122His, p.Tyr127Cys, p.His191Arg, p.Asp223His, and p.Asp223Val mutants was not obviously impaired in comparison with that of WT C α , structural A subunit binding was significantly decreased for p.Asp88Gly, p.Asp131His, Tyr265Cys, p.Arg295*, and p.Phe308dup mutants (30%–60% of WT C α binding) and even completely abolished for the p.Arg214* mutant (Figures 5A–5C and Table 2). For the latter, however, we noticed strong binding to endogenous alpha 4 (Figure 5D and Table 2), a non-canonical PP2A subunit whose binding is mutually exclusive with the A subunit. Alpha 4 binding stabilizes the C subunit in a latent form,^{24,36} which is consistent with our observation of the complete lack of phosphatase activity for p.Arg214* as shown in our assays (Figure 3).

We next assessed binding of the C α mutants to at least one representative of the regulatory B55, B56, B'', and B'''/striatin subunit classes. To this end, N-terminally GST-tagged B55 α (encoded by PPP2R2A), B56 α –B56 ϵ (encoded by PPP2R5A–PPP2R5E), and B''/PR72 (encoded by PPP2R3A) (Figure 6A, Table 2, and Figure S1) or N-terminally GFP-tagged B'''/SG2NA (encoded by STRN3) (Figure 6B and Table 2) was co-expressed with HA-tagged WT or mutant C α . The retrieved C α proteins in GST pull-downs or GFP traps were subsequently visualized by anti-HA immunoblotting.

Whereas essentially no B-type subunit binding defects could be identified for the p.Gln122His, p.Asp223His, and p.Asp223Val variants in GST pull-downs (Figure 6C and Figures S1E–S1G), a highly diverse aberrant B-type subunit binding pattern was detected for the other mutants. The p.His191Arg mutant significantly lost binding

Table 2. Overview of Mutation-Specific Biochemical Defects

PPP2CA Variant	Protein Detected	Activity (%)	A Binding (%)	Alpha 4 Binding	Demethylation	B55 α Binding (%)	B56 α Binding (%)	B56 β Binding (%)	B56 γ 1 Binding (%)	B56 δ Binding (%)	B56 ϵ Binding (%)	PR72 Binding (%)	STRN3 Binding (%)
WT C α	+	100	100	NA	100	100	100	100	100	100	100	100	100
p.Asp88Gly	+	3****	37****	NA	1379**	0****	17****	20****	88	195	41**	18**	52**
p.Gln122His	+	40***	78	NA	183*	176	228**	177	135	285	254*	118	384*
p.Tyr127Cys	+	11****	82	NA	1109*	108	15**	27*	33*	18**	14****	31*	273
p.Asp131His	+	211*	48****	NA	200	185	0****	0****	0****	5****	0****	149	296
p.His191Arg	+	26****	102	NA	218	199	99	73	57	14****	34	65	652*
p.Asp223Val	+	101	83	NA	243*	239	165	160	254	180	163*	479	72
p.Asp233His	+	61*	81	NA	138	129	97	105	111	105	147**	108	142
p.Tyr265Cys	+	19**	30***	NA	1046*	1****	6***	3****	24*	42*	7****	2****	8****
p.Arg295*	+	48**	36**	NA	ND	0****	0****	0****	0****	18***	0****	0****	235
p.Phe308dup	+	10****	62*	NA	ND	15****	64	166	69	206	198	5****	372*
p.Gly60Val	±	ND	ND	NA	ND	ND	ND	ND	ND	ND	ND	ND	ND
p.Gln125*	–	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
p.Phe146fs	–	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
p.Arg214*	+	0****	0****	yes	ND	0****	0****	0****	0****	0****	0****	0****	NA

Abbreviations are as follows: ND, not determinable; NA, not analyzed; +, present; ±, detected at very low level; –, not detected; *p ≤ 0.05; **p ≤ 0.01; ***p ≤ 0.001; ****p ≤ 0.0001.

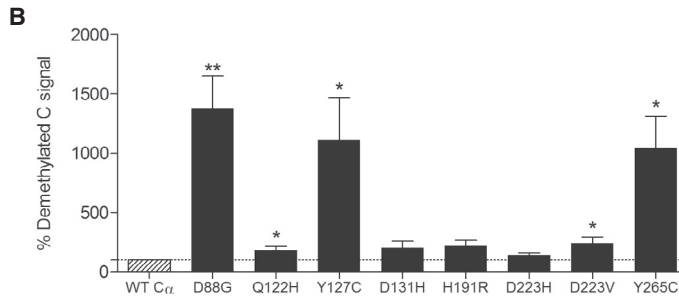
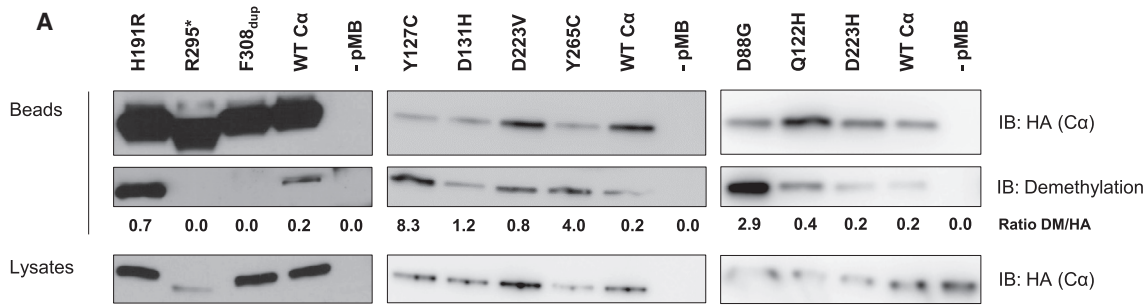


Figure 4. Carboxyterminal Methylation of Mutant PP2A C α Subunits

(A) HA-tagged WT and mutant C α proteins were isolated from transfected HEK293T cells by HA pull-down. PP2A C subunit methylation was subsequently assessed by immunoblotting with a demethyl-specific anti-C monoclonal antibody. Immunoblot data from a single representative experiment are shown ($n \geq 4$).

(B) The average ratios of the quantified demethyl-specific signals over the quantified anti-HA signals \pm SEM are displayed ($n \geq 4$) in relation to those of WT C α (set at 100%). A one-sample t test (compare to 100%) was used for assessing statistical significance (* $p \leq 0.05$, ** $p \leq 0.01$).

to only B56 δ (Figure 6C and Figure S1A). The p.Arg295* mutant lost binding to all B55, B56, and B'' subunits tested, except for B56 δ , to which binding was, although significantly decreased, still slightly retained (Figure 6C and Figure S1A). The p.Asp88Gly and p.Tyr265Cys variants showed a similar pattern of severe binding defects to all B55, B56, and B'' subunits but a higher overall binding level to the B56-type subunits than p.Arg295*, particularly to the B56 γ and B56 δ subunits (Figure 6C and Figures S1C and S1F). The p.Asp131His variant specifically lost binding to all B56 subunits, whereas binding to B55 and B'' subunits was entirely preserved (Figure 6C and Figure S1D). On the contrary, mutant p.Phe308dup completely lost binding to B''/PR72, showed decreased binding to B55 α , and retained binding to all B56 isoforms (Figure 6C and Figure S1B). For the p.Tyr127Cys variant, binding was diminished for all B56 and B'' subunits tested and intact for B55 α (Figures 6A and 6C). As expected from its complete lack of A subunit binding, p.Arg214* failed to bind any B-type subunit tested (data not shown).

Interestingly, the majority of C α mutants retained binding to the B''/striatin subunit STRN3, and for the p.Gln122His, p.His191Arg, and p.Phe308dup variants, this interaction was even significantly enhanced (Figures 6B and 6C). This observation was also previously made for ID-associated mutations in *PPP2RIA*, encoding the A α

subunit.³⁸ However, for the p.Asp88Gly and p.Tyr265Cys mutants, STRN3 binding was diminished or completely abolished (Figures 6B and 6C).

Discussion

PP2A is an important holoenzyme for which broad substrate specificity is orchestrated by different subunit compositions: in particular, the B subunits play a pivotal role in determining substrate specificity, regulation, and hence physiologic functions of the different PP2A complexes. With this broad specificity, PP2A is involved in diverse cellular processes, and variants in either of these subunits could therefore cause disease. Here, we report the identification of a syndrome caused by *de novo* mutations in *PPP2CA* and characterized by mild to profound ID and DD, behavioral problems, variable types of epilepsy, hypotonia, abnormal head circumference, and brain abnormalities such as corpus callosum abnormalities and ventriculomegaly. Some individuals had mild dysmorphic features, but a recognizable facial gestalt could not be observed.

Mutations in several other PP2A-subunit-encoding genes have been reported before and result in an ID and DD phenotype. Interestingly, albeit not unsurprisingly,

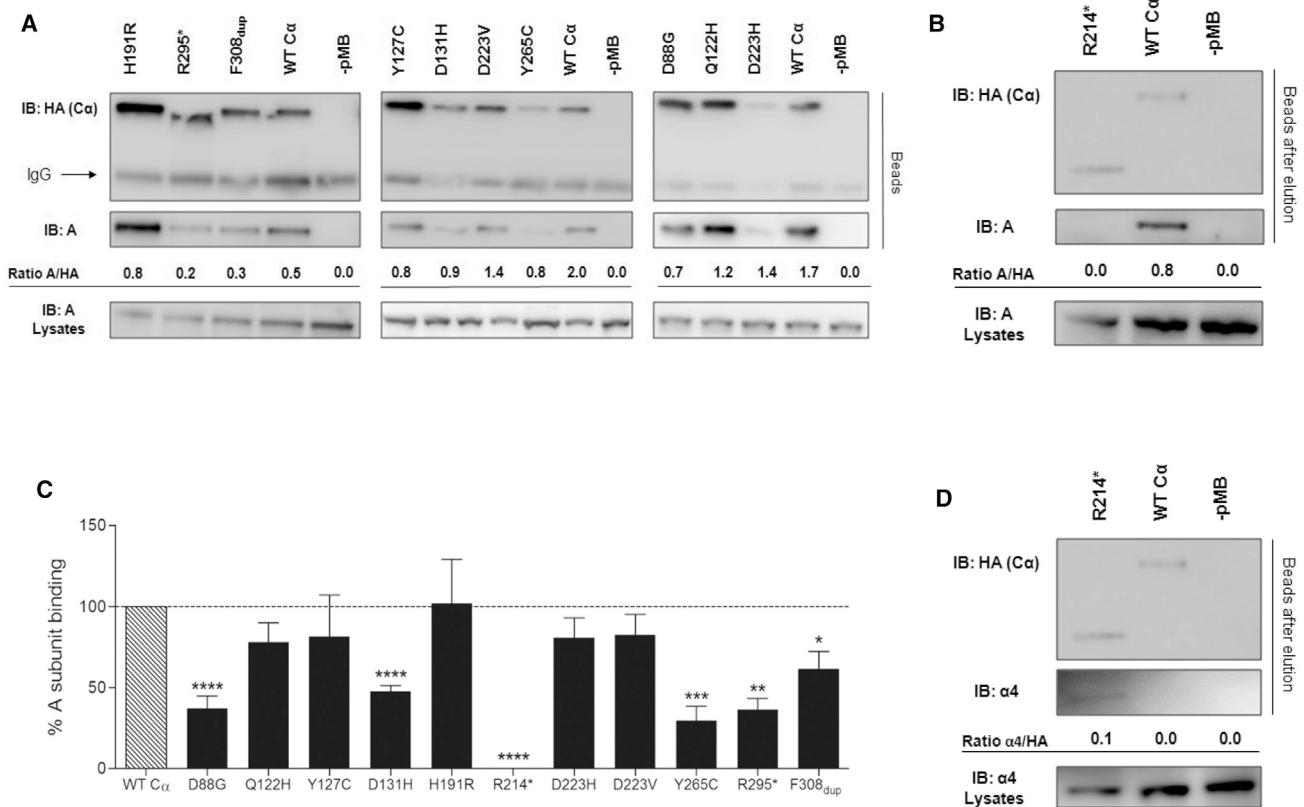


Figure 5. Binding of Mutant PP2A C α Subunits to the A Subunit

(A) A subunit binding. HA-tagged WT and mutant C α proteins were purified from transfected HEK293T cells by HA pull-down, and the presence of endogenous A subunit in the complexes was determined by anti-A immunoblotting. Abbreviated amino acid changes are as follows: H191R, p.His191Arg; R295*, p.Arg295*; F308dup, p.Phe308dup; Y127C, p.Tyr127Cys; D131H, p.Asp131His; D223V, p.Asp223Val; Y265C, p.Tyr265Cys; D88G, p.Asp88Gly; Q122H, p.Gln122His; and D223H, p.Asp223His.

(B) Same experimental setup as in (A), but in this case, HA-agarose-bound proteins were eluted from the beads with an excess of free HA-tagged peptide before anti-A immunoblotting of the eluates. This setup was necessary for enabling unambiguous visualization of HA-tagged p.Arg214*, which co-migrated with immunoglobulin light chains at approximately 25 kDa.

(C) Quantified values of A subunit binding. Results represent the average value \pm SEM of the ratios of the quantified anti-A signal to the quantified anti-HA signal for a given C α mutant in relation to those of WT C α (set at 100% in each experiment), as determined in at least four independent experiments ($n \geq 4$). A one-sample t test (compare to 100%) was used for assessing statistical significance (* $p \leq 0.05$, ** $p \leq 0.01$, *** $p \leq 0.001$, **** $p \leq 0.0001$).

(D) Alpha 4 subunit binding. HA-tagged WT and p.Arg214* mutant C α proteins were purified from transfected HEK293T cells by HA pull-down and eluted from the beads by excess HA peptide, and eluates were subjected to immunoblotting with anti-alpha 4 monoclonal antibodies for the detection of binding of endogenous alpha 4.

individuals with a *de novo* mutation in *PPP2CA* showed a phenotypic resemblance to those individuals harboring an ID- and DD-related pathogenic variant in other PP2A subunits^{10–12} (Table S2). This overlap consisted of ID, DD, and hypotonia (in all individuals with *PPP2R1A* and *PPP2R5D* mutations), epilepsy and ventriculomegaly (in some individuals with *PPP2R1A* and *PPP2R5D* mutations), ASD (in some individuals with a *PPP2R5D* mutation), corpus callosum agenesis (in all individuals with a *PPP2R1A* mutation and some individuals with a *PPP2R5D* mutation), and congenital heart disease (ventricular and atrial septal defects in two individuals with a mutation in *PPP2R5D*). Severe speech delay was also noted in multiple individuals and for multiple subunits, which might be partially related to the severity of ID but was more severe than expected in several individuals. A noticeable difference from mutations in other subunits is the absence of

overgrowth, which was reported for individuals with pathogenic mutations in *PPP2R5B*, *PPP2R5C*, and *PPP2R5D*¹¹ but was observed in only one individual with a *de novo* mutation in *PPP2CA*. In addition, although two individuals were macrocephalic, five other individuals had microcephaly, a phenotype also reported for cases of *PPP2R1A* mutations.¹⁰ Recently, brain-specific *Ppp2ca* knockout was reported to result in mice with obvious morphological and behavioral abnormalities, including microcephaly, cortical neuronal atrophy, plastic synapse deficits, and learning and memory impairments at an early postnatal stage.³⁹ These features were also partly observed in individuals with *de novo* mutations in *PPP2CA*, and although the phenotypes of the heterozygous mice (if any) were not further documented, these mice could be an excellent *in vivo* model for further studies. Likewise, congenital heart defects, specifically affecting septum

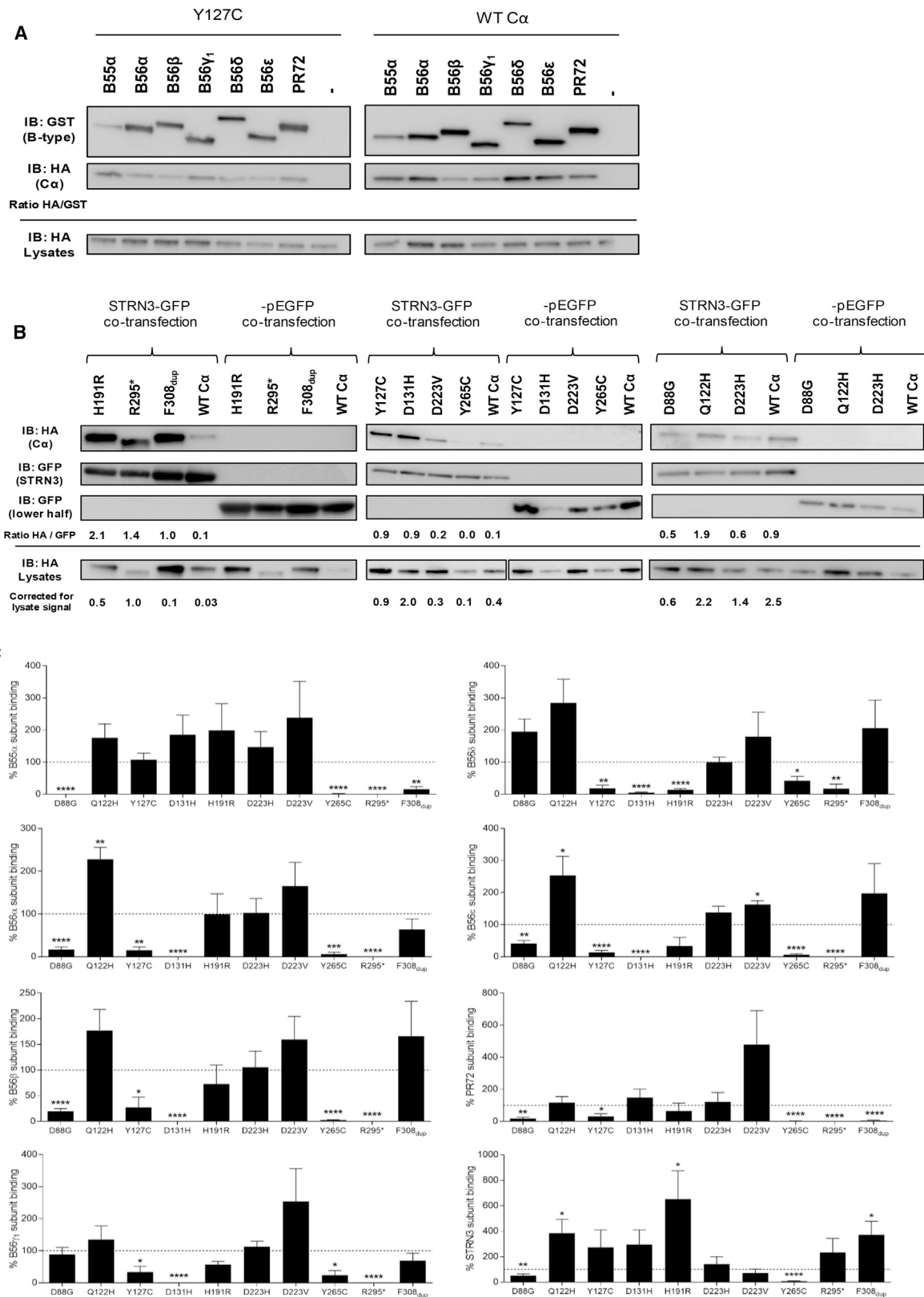


Figure 6. Binding of Mutant PP2A C α Subunits to B-type Subunits

(A) GST-tagged B-type subunits were co-expressed with HA-tagged WT or mutant p.Tyr127Cys C α protein in HEK293T cells. The presence of the PP2A C variant was subsequently assessed in GST pull-downs by anti-HA immunoblotting. Representative blots of similar assays with other C α variants can be found in [Figure S1](#).

(B) GFP-tagged B α /STRN3 or GFP alone was co-expressed with HA-tagged WT or mutant C α proteins in HEK293T cells. The presence of C α variants was subsequently assessed in GFP traps by anti-HA immunoblotting.

(legend continued on next page)

formation, have been reported in B56 γ -knockout mice and in transgenic mice specifically overexpressing a dominant-negative mutant of $A\alpha$ in (cardiac) muscle.^{40,41} All functionally characterized and clinically described variants in subunits of PP2A so far and our data on *PPP2CA* variants collectively seem to constitute overlapping syndromic forms of ID involving more than one gene but functionally converging to PP2A deficiency, and they could therefore be redefined as “PP2A-related neurodevelopmental syndrome.”

Notably, for most individuals, WES was used to identify the *PPP2CA* mutation, so other rare (*de novo*) mutations were also identified in 6 of 16 individuals. Whereas these most likely reflect the background per-generation *de novo* mutation rate, a contribution of these *de novo* variants to the individuals' phenotypes cannot be excluded. Similarly, involvement of the partial *CDKL3* deletion to the phenotype of individual 11 cannot be ruled out given that a balanced *CDKL3*-disrupting translocation leading to decreased expression of *CDKL3* has previously been reported in an individual with mild ID.⁴² To collect further insight into the clinical spectrum, we established a website on the Human Disease Genes platform for the collection of clinical data (see [Web Resources](#)).

By far most reported individuals with PP2A variants so far harbor *de novo* mutations in *PPP2R5D*, encoding the regulatory B56 δ subunit.^{10–12} Biochemical characterization has revealed that the vast majority of these variants show decreased binding to the A and C subunits, hinting at a dominant-negative mode of action.¹⁰ Similarly, characterization of *de novo* mutations in *PPP2R1A*, encoding the scaffolding $A\alpha$ subunit,¹⁰ has identified defective binding of nearly all B-type subunits, except for B56 δ . Yet for the latter variants, the $A\alpha$ -B56 δ complexes have proved catalytically impaired,¹⁰ potentially because of increased recruitment of a cellular PP2A inhibitor, TIPRL1.³⁸ Our functional assays, focusing on the functional defects following *PPP2CA* mutations, showed similar defects—that is, our binding assays suggested that some $C\alpha$ variants, although catalytically impaired, were indeed still able to form trimeric complexes with specific B-type subunits, providing a mechanistic basis for their potential dominant-negative mode of action. Clearly, such a dominant-negative mechanism would affect only a set of specific holoenzymes rather than the whole PP2A spectrum. This scenario is most likely the case for the p.Asp88Gly, p.Tyr265Cys, p.Arg295*, and p.Phe308dup variants, which all showed significantly decreased catalytic activity but still retained binding to at least B56 δ . Moreover, the p.Asp88Gly and p.Arg295* variants occurred in individuals

with severe ID, and the p.Phe308dup variant caused moderate ID—all in line with a potential dominant-negative effect on PP2A-B56 δ complexes (for p.Arg295*), on PP2A-B56 γ and B56 δ complexes (for p.Asp88Gly), or on all PP2A-B56 complexes (for p.Phe308dup). In contrast, the individual with the p.Tyr265Cys variant had mild ID (with comorbid epilepsy, hypotonia, brain MRI abnormalities, ASD, and ADHD), suggesting that in this case, the 50% retained binding to B56 δ might not be causally involved, but rather the significantly decreased binding to all B-type subunits is the most important determinant, bringing us into a scenario of haploinsufficiency in this particular individual. Likewise, haploinsufficiency would be the proposed mechanism for the partial *PPP2CA* deletion and the p.Gln125*, p.Phe146Leufs*29, and p.Arg214* variants, all of which lead to complete null alleles (due to the lack of expression in the first three cases or expression of a completely dysfunctional variant in the last case), as well as for the very poorly expressed p.Gly60Val variant. All of these individuals had mild ID often accompanied by severe epilepsy. Notably, the remaining 50% of WT $C\alpha$ in those five individuals might no longer be sufficient to assemble all PP2A holoenzymes, as previously suggested for specific cancer-associated $A\alpha$ (*PPP2R1A*) mutants^{43,44} and haploinsufficiency of *PPP2R4*, encoding the PP2A activator PTPA.⁴⁵ Upon loss of one functional allele for those two genes, selective PP2A dysfunctions were observed for B56-subunit-containing complexes, specifically B56 γ , B56 δ , and B56 ϵ (whereas B55 complexes remained fully functional), as a result of a competition mechanism between B subunits for binding the remaining WT $A\alpha$ subunit or fully activated C subunit. In other words, both *PPP2R1A* and *PPP2R4* haploinsufficiency caused a selective PP2A-B56 defect. In this respect, it is interesting to point out that three B-type-subunit-encoding genes hitherto reported to be involved in the etiology of ID and DD all encode subunits of the B56 family (δ , γ , and β isoforms).^{10,11} Thus, in those *PPP2CA* cases where we suspect haploinsufficiency as the pathological mechanism, the eventual PP2A dysfunctions could be highly similar and mainly affect B56-type holoenzymes. On the basis of our biochemical assays, such a selective B56 defect also seems to be the case for the catalytically competent p.Asp131His variant, which retained binding to all B-type subunits tested, except for all B56 isoforms. The p.Tyr127Cys variant also lost binding to all B56 subunits, as well as to B''/PR72. However, in this case, the catalytic activity and methylation of this variant were also significantly impaired, suggesting that a dominant-negative mechanism toward PP2A-B55 and PP2A-STRN

(C) Quantified values of C subunit binding. Results represent the average value \pm SEM of the ratios of the quantified anti-HA signal to the quantified anti-GST or anti-GFP (in the case of STRN3) signal for a given $C\alpha$ mutant in relation to those of WT $C\alpha$ (set at 100% in each experiment, dotted line), as determined in at least three independent binding experiments ($n \geq 3$). A one-sample t test (compare to 100%) was used for assessing statistical significance (* $p \leq 0.05$, ** $p \leq 0.01$, *** $p \leq 0.001$, **** $p \leq 0.0001$).

Abbreviated amino acid changes are as follows: H191R, p.His191Arg; R295*, p.Arg295*; F308dup, p.Phe308dup; Y265C, p.Tyr265Cys; Y127C, p.Tyr127Cys; D131H, p.Asp131His; D223V, p.Asp223Val; D88G, p.Asp88Gly; Q122H, p.Gln122His; and D223H, p.Asp223His.

complexes might additionally be in place—further explaining the severe ID in this individual. Despite a relatively large number of available *PPP2CA*-knockout mouse models, heterozygous *Cα*-null mice have unfortunately never been phenotypically or biochemically examined.⁴⁶ In any case, our data provide evidence that *PPP2CA* is a haploinsufficient gene in humans.

The only recurrent *PPP2CA* variant in our cohort, c.572A>G (p.His191Arg), showed a (non-significant) increase in demethylation signal (218%; $p = 0.095$) and a decrease in activity (48%; $p = 0.001$), whereas no changes in A or B subunit interaction could be detected, except for B56δ (for which binding was decreased) and for B'''/STRN3 (for which binding was increased). Both individuals with this variant had a mild ID phenotype and fever-related seizures. Structural studies suggested that the His191 residue is important in PP2A-C's interaction with alpha 4²⁴ and thereby induces displacement of the catalytic center and subsequent reduction in activity.²⁸ Additionally, in a yeast model, the p.His191Arg variant was incapable of exhibiting a dominant-negative effect on WT human PP2A *Cα* in a growth assay.⁴⁷ This could indicate a potential haploinsufficient mechanism of the p.His191Arg variant, which mainly affects B56δ-containing holoenzymes. The biochemical profile of the p.Gln122His variant appeared very similar to that of p.His191Arg, except for the retained binding to B56δ in p.Gln122His. Both affected residues are indeed two of four residues that are directly involved in binding the hydrophobic part of pharmacologic PP2A inhibitors, okadaic acid, and microcystin.²⁵

Finally, the p.Asp223His and p.Asp223Val variants did not seem to have a significant impact on PP2A assembly and function in any of our assays tested, except for a slight increase in B56ε binding (150%), a slight decrease in activity (for p.Asp223His), and a light increase in demethylation signal (for p.Asp223Val). The latter might suggest that the clinical phenotype might be milder, but this was not the case for individual 16 (p.Asp223His), who had severe ID. Also, although the other individual (individual 7, p.Asp223Val) initially had a normal IQ, she showed speech delay and ASD and additionally developed progressive cognitive dysfunction after a psychotic episode at the age of 17 years. In the absence of a clear pathogenic effect for these two variants in the majority of our biochemical assays tested, it could be speculated that another (yet unidentified) genetic variant contributes to the disease manifestation of individuals 7 and 16.

Many of the observed biochemical defects in specific *Cα* variants can be rationalized by structural data from crystallographic studies or are sustained by previously reported biochemical and functional studies of PP2A C mutants in yeast. The variants affecting Asp88 and two tyrosine residues in the catalytic center—p.Tyr127Cys and p.Tyr265Cys—both induced a drastic decrease in methylation and activity. Most likely the variants induce

an inactive *Cα* conformation, which subsequently prevents methylation. In yeast, a p.Tyr127Asn variant was also identified as LoF because it could not rescue the growth defect of a PP2A-C deletion strain.⁴⁷ For the p.Asp131His, p.His191Arg, and p.Asp223His variants, the effects were less clear, although there might have been a trend toward slightly decreased methylation for the p.His191Arg variant. On the other hand, a slight but significant decrease in methylation of the p.Asp223Val mutant was detected, whereas the p.Asp223His variant, affected in the same residue, behaved like the WT. Besides a charge difference between the mutated amino acids, a straightforward structural explanation for this observation could not be found. A similar increase in demethylated signal could be observed for the p.Gln122His variant. Although it's impossible to measure with the available antibodies, we expect methylation of p.Phe308dup to be diminished as well. On the basis of the “deep and narrow cleft” conformation of the active center of the PP2A methyltransferase LCMT-1 and the fact that several residues in the C tail need to form stabilizing contacts with the LCMT-1 active-site residues for efficient methylation to occur, a tail that is 1 amino acid too long might no longer fit in and thus no longer be modified.

The c.640 C>T (p.Arg214*) nonsense mutation is located in *PPP2CA* exon 5. Migueleti et al. reported on the existence of a shorter *PPP2CA* mRNA, in which this specific exon can be skipped, in peripheral-blood mononuclear cells under specific growth conditions (serum exhaustion), and this mRNA resulted in a slightly shorter *Cα* isoform.⁴⁸ Although we do not favor potential exon 5 skipping in the c.640 C>T (p.Arg214*) variant given our data and observed molecular weight of the protein detected, it is remarkable that this shorter isoform was reported as catalytically inactive without any binding to the scaffolding A subunit and increased binding to alpha 4, very much like our data for the truncated p.Arg214* protein. This would suggest an important role for exon-5-encoded amino acids in A subunit binding. Tang et al. further demonstrated that expression of the shorter *Cα* isoform had no effect on the expression or the modification of the longer *Cα* variant, thus disfavoring a dominant-negative effect of the shorter variant.⁴⁹

In summary, we have identified 16 individuals with a syndrome characterized by ID, DD, hypotonia, epilepsy, behavioral problems, and structural abnormalities of the brain and with a *de novo* mutation in or partial deletion of *PPP2CA*, encoding one of the catalytic subunits of PP2A. Functional characterization of all the *Cα* variants in HEK293T cells showed a diverse spectrum of biochemical distortions that were unique to each variant but always justified the conclusion of a functional loss and affected at least the functionality of the PP2A-B56δ complexes. Only for both p.Asp223 missense variants, our results failed to reveal a major biochemical defect. Our data further expand the repertoire of PP2A subunits involved in the etiology of ID and DD and underscore

the importance of PP2A in neurodevelopmental processes and brain function. Hence, we propose that *de novo* mutations in *PPP2CA*, *PPP2R1A*, and *PPP2R5D* constitute a spectrum of overlapping ID syndromes characterized by mild to severe phenotypes and functional convergence by severe PP2A dysfunction, and we propose that these collectively be called “PP2A-related neurodevelopmental disorders.”

Supplemental Data

Supplemental Data include a Supplemental Note, one figure, and two tables and can be found with this article online at <https://doi.org/10.1016/j.ajhg.2018.12.002>.

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Declaration of Interests

H.M.L. is an employee of GeneDx Inc., a wholly owned subsidiary of OPKO Health Inc.

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Web Resources

DECIPHER, <https://decipher.sanger.ac.uk/>
denovo-db, <http://denovo-db.gs.washington.edu/denovo-db/>
ExAC Browser, <http://exac.broadinstitute.org/>
GeneMatcher, <https://genematcher.org/>
Human Disease Genes: *PPP2CA*, <http://www.humandiseasesgenes.com/PPP2CA/>
OMIM, <http://www.omim.org/>

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