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# Bi-allelic Variants in *RALGAPA1* Cause Profound Neurodevelopmental Disability, Muscular Hypotonia, Infantile Spasms, and Feeding Abnormalities

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Ral (Ras-like) GTPases play an important role in the control of cell migration and have been implicated in Ras-mediated tumorigenicity. Recently, variants in *RALA* were also described as a cause of intellectual disability and developmental delay, indicating the relevance of this pathway to neuropediatric diseases. Here, we report the identification of bi-allelic variants in *RALGAPA1* (encoding Ral GTPase activating protein catalytic alpha subunit 1) in four unrelated individuals with profound neurodevelopmental disability, muscular hypotonia, feeding abnormalities, recurrent fever episodes, and infantile spasms . Dysplasia of corpus callosum with focal thinning of the posterior part and characteristic facial features appeared to be unifying findings. RalGAPA1 was absent in the fibroblasts derived from two affected individuals suggesting a loss-of-function effect of the *RALGAPA1* variants. Consequently, RalA activity was increased in these cell lines, which is in keeping with the idea that RalGAPA1 deficiency causes a constitutive activation of RalA. Additionally, levels of RalGAPA1 deficiency clearly increased cell-surface levels of lipid raft components in detached fibroblasts, which might indicate that anchorage-dependence of cell growth signaling is disturbed. Our findings indicate that the dysregulation of the RalA pathway has an important impact on neuronal function and brain development. In light of the partially overlapping phenotype between *RALA* and *RALGAPA1*-associated diseases, it appears likely that dysregulation of the RalA signaling pathway leads to a distinct group of genetic syndromes that we suggest could be named RALopathies.

The Ral guanine nucleotide exchange factor (RalGEF)-Ral GTPase cascade is one of three major downstream pathways that become activated upon Ras signaling. The others include the Raf/mitogen-activated protein kinase (MAPK) cascade and the phosphoinositide 3-kinase (PI3K)-dependent phosphoinositide pathway.<sup>1</sup> *RALGAPA1* (previously referred to as *TULIP1* or *GARNL1* [MIM: 608884]) encodes the catalytic subunit of Ral GTPase-activating protein (GAP) complex for two highly related small G proteins, RalA and RalB.<sup>1,2</sup> Ral proteins draw significant research interest as promising cancer drug targets downstream of Ras.<sup>3,4</sup> The role of RalA in normal cell physiology centers around the regulation of intracellular-vesicle trafficking. RalA mediates post-Golgi vesicle trafficking toward the plasma membrane by rearranging the cytoskeleton and via its interaction with the tethering complex exocyst, which positions vesicles for docking and fusion at the plasma membrane.<sup>5,6</sup> Thus, RalA is implicated in cellular processes, which depend on directional membrane trafficking, such as cell migration, establishment of cell polarity, neuronal polarity, and ciliogenesis. Consequently, RalA is essential for normal brain development. *Rala*-null mouse embryos display exencephaly, which recently has been classified as ciliopathy, between embryonic day 10.5 (E10.5) and E19.5.<sup>7</sup>

RalA activity is regulated in response to extracellular cues. For example, RalA mediates the delivery of vesicles that contain the glucose transporter GLUT4 toward the

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plasma membrane in adipocytes treated with insulin and also mediates the formation of lamellipodia in fibroblasts treated with TNFa.<sup>8,9</sup> As is true for a typical G protein, RalA activity depends on its nucleotide-binding status. Binding to GTP activates RalA, whereas binding to GDP renders RalA inactive.<sup>10</sup> Thus, RalA is regulated by factors that facilitate nucleotide exchange (guanine exchange factors [RalGEFs]) or GTP hydrolysis (RalGAP). In its basal state, RalA is maintained in an inactive GDP-bound form by the RalGAP complex and can be activated either by Ral-GEF (e.g., downstream of Ras) or by the inactivation of Ral-GAP (as was described for insulin signaling).<sup>10</sup> The RalGAP complex is a heterodimer of a catalytic subunit, either Ral-GAPA1 or RalGAPA2 (the expression of which is cell-typespecific), and a scaffolding subunit, RalGAPB.<sup>2</sup> It has been demonstrated in several mouse models that the loss of either RalGAP subunit results in constitutive activation of Ral proteins, and physiological outcomes are highly cell-type- and development-stage-specific. It is important to note that the ability to cycle between GTP- and GDPbound states might be important for RalA function.<sup>10</sup> Thus, although constitutive RalA activation might facilitate certain signaling cascades, it might disrupt the others.

Although the Ral GTPase pathway is well known in cancer biology, its implication in Mendelian disorders has yet to be elucidated. Pathogenic variants in RALA (MIM: 179550) have been identified as a cause of an autosomaldominant inherited syndrome typified by speech and motor delays, intellectual disability, early-onset epilepsy, short stature, facial dysmorphism, and finger and toe abnormalities (e.g., long fingers with extensible joints, finger overlap, and toe clinodactyly).<sup>11</sup> RALGAPA1 is ubiquitously expressed; the highest expression levels are in the brain, ovaries, and uterus (GTEx: phs000424.v7.p2 on 02/07/ 2019). RALGAPA1 has been suggested as a candidate gene for neurodevelopmental disorders because microdeletions, including deletion of RALGAPA1, were identified in individuals with developmental delay, microcephaly, seizures, muscular hypotonia, and mild brain atrophy. In these cases, RALGAPA1 was the only deleted gene with high expression in the brain.<sup>12,13</sup> A morpholino oligonucleotide knockdown of the homolog tulip1 in a zebrafish model resulted in severe developmental impairment of the brain and comparably mild abnormalities of the whole body.<sup>12</sup>

Here, by identifying bi-allelic variants in *RALGAPA1* in four unrelated families, we establish that the loss of *RALGAPA1* disrupts the RalGAP complex, leads to constitutive RalA activation, and causes a severe neurodevelopmental disease with profound muscular hypotonia, infantile spasms, and feeding difficulties.

Legal guardians of all individuals investigated gave written informed consent for genetic testing and functional studies from tissue material (families A and B). Permission to include clinical photographs was obtained. The study was performed according to the declaration of Helsinki. The local ethical committees at the Heinrich-Heine-University Düsseldorf (#4957), the Technical University in Munich (#5360/12S), King Faisal Specialist Hospital and Research Center (RAC #2080006), and the Sheba Medical Center (#7786-10-SMC) approved the genetic studies and the studies on tissue samples derived from individuals 1 and 2, where applicable.

The clinical phenotype of the affected individuals is summarized in Table 1. Detailed case reports can be found in the Supplemental Information.Pedigrees and clinical images are depicted in Figure S1 and Figure 1, respectively.

All individuals were born after uneventful pregnancies. Only one child was born preterm. Birth parameters were normal in two individuals, and two children were born small for gestational age. All four children presented with respiratory distress requiring transient non-invasive ventilation directly after birth. One child required a tracheostomy as a result of persistent respiratory problems. Congenital profound muscular hypotonia was a unifying clinical feature. During the first year of life, three individuals had frequent episodes of high fever. A dysregulation of central temperature control was suspected. In two individuals, recurrent viral infections were noted, but there were no abnormalities on immunologic work-up. In one child, hypohidrosis was diagnosed. All individuals had feeding abnormalities requiring tube feeding. Infantile spasms starting between the ages of 8 and 12 months were observed in three out of four children. In this context, it should be noted that the only child without clinical evidence of epilepsy is still very young (8 months at last followup appointment). Seizures responded to adrenocorticotropic hormone (ACTH) treatment in one child, and two children suffered from drug-resistant epilepsy. Profound developmental delay was observed in all children, none of whom reached any major developmental milestones. Brain MRIs in the affected individuals revealed dysplasia of the corpus callosum; focal thinning of the posterior part (splenium) was a unifying CNS finding (Figure 1 and Figure S2). In addition, a wide spectrum of unspecific abnormalities, including gray-matter heterotopias, ectopic posterior pituitary, signal abnormalities in basal ganglia, and stratum subependymale, was observed (see Figure S3). Only one child showed signs of global brain atrophy (Figure 1). Interestingly, moderate dysmorphic facial features were observed in all of the children. They had a rather consistent phenotype of upslanting palpebral fissures, mild epicanthus, a short nose with a flattened nasal bridge, and slightly anteverted nares (see Figure 1). The oldest child is now 5 years old and is severely disabled. He is unable to sit, crawl, or pull up. Head control is still inadequate, and truncal muscle tone is profoundly reduced. He did not acquire language, and he never interacted with his environment.

Exome sequencing (ES) was performed at four clinical sites in Munich, Tel-Aviv, Chapel Hill, and Riyad. For family A, trio ES was performed with the Sure Select Human All Exon kit V5 (index individual) and V6 (parents; Agilent) for exome enrichment and a HiSeq2500 (index individual)

| Family                     | Family A  | Family B  | Family C   | Family D  |
|----------------------------|---|---|--|---|
| ID                         | Individual 1  | Individual 2  | Individual 3   | – –<br>Individual 4   |
| Age at last examination    | 4 1/2 years   | 8 months  | 18 months  | 17 months   |
| Gender                     | male  | female  | male   | male  |
| Genome molecular           | exome sequencing  | exome sequencing  | exome sequencing                                       | exome sequencing  |
| analysis                   |   |   |  |   |
| Variant (NM_194301.4)      | c.3227A>G;1126C>T,<br>(p.Asn1076Ser;Arg376*)                      | c.4992del;4992del,<br>(p.Phe1664Leufs*11;<br>Phe1664Leufs*11)                       | c.610G>T;610G>T,<br>(p.Glu204*;Glu204*)                | c.5732C>G);(5732C>G),<br>(p.Ser1911*;Ser1911*)                                |
| Pathogenicity              | pathogenic  | pathogenic  | pathogenic   | pathogenic  |
| Familial History and Per   | inatal Period   |   |  |   |
| Ancestry                   | German  | Palestinian   | German and Irish                                       | Saudi   |
| Consanguinity              | no  | yes   | no   | yes   |
| Pregnancy                  | unremarkable  | unremarkable  | poor maternal<br>weight gain                           | preterm delivery  |
| Prematurity                | no  | no  | no   | born at 34 weeks  |
| Perinatal period           | neonatal respiratory<br>distress                                  | neonatal respiratory<br>distress  | neonatal respiratory<br>distress                       | neonatal respiratory distress   |
| Growth Parameters          |   |   |  |   |
| Birth length               | 56 cm (97 <sup>th</sup> centile)                                  | N/A   | 45.5 cm (5 <sup>th</sup> centile)                      | 43 cm (<5 <sup>th</sup> centile)  |
| Birth weight               | 3,875 g (79 <sup>th</sup> centile)                                | 3,000 g (20 <sup>th</sup> centile)  | 2,240 g (5 <sup>th</sup> centile)                      | 2,115 g (<5 <sup>th</sup> centile)  |
| Head circumference         | 36.5 cm (93 <sup>th</sup> centile)                                | N/A   | 33.5 cm (13 <sup>th</sup> centile)                     | 33.5 cm (10 <sup>th</sup> centile)  |
| Age at examination (y)     | 4 1/2 years   | 3 months  | 18 months  | 17 months   |
| Current height             | 111 cm (75 <sup>th</sup> centile)                                 | 51 cm (<1 <sup>st</sup> centile)  | 78.1 cm (6 <sup>th</sup> centile)                      | 75 cm (5 <sup>th</sup> centile)   |
| Current weight             | 16.4 kg (25 <sup>th</sup> centile)                                | 2.9 kg (<1 <sup>st</sup> centile)   | 9.52 kg (7 <sup>th</sup> centile)                      | 12.5 kg (75 <sup>th</sup> centile)  |
| Current head circumference | 51.5 cm (52 <sup>th</sup> centile)                                | 35 cm (<1 <sup>st</sup> centile)  | 46.8 cm (7 <sup>th</sup> centile)                      | 48 cm (37 <sup>th</sup> centile)  |
| Motor development          | severely delayed  | severely delayed  | severely delayed                                       | severely delayed  |
| Speech development         | none  | none  | none   | none  |
| Neurological Findings      |   |   |  |   |
| Muscle tone                | profound muscular<br>hypotonia                                    | profound muscular<br>hypotonia  | profound muscular<br>hypotonia                         | profound muscular<br>hypotonia  |
| Head control               | poor  | poor  | poor   | poor  |
| Spasticity                 | no  | yes   | no   | yes   |
| UL/LL tendon reflexes      | reduced/reduced   | reduced/reduced   | reduced/reduced  | increased   |
| Seizures                   | infantile spasms  | no  | infantile spasms                                       | infantile spasms  |
| Brain MRI                  | focal thinning of corpus<br>callosum, gray matter<br>heterotopias | focal thinning of corpus<br>callosum, thin pituitary<br>stalk, global brain atrophy | focal thinning of corpus callosum, ectopic pituitary   | focal thinning of corpus callosum   |
| Skin findings              | dry skin and hypohidrosis   | none  | none   | none  |
| Ophthalmologic<br>findings | bilateral cataract  | none  | cortical vision impairment<br>(no cataracts)           | suspected cortical vision<br>impairment (not confirmed)                       |
| General Clinical Finding   | s   |   |  |   |
| Heart                      | normal  | normal  | normal   | mild tricuspid valve<br>regurgitation and mild<br>mitral valve regurgitation. |
| Lungs                      | normal  | tracheostomy  | laryngomalacia, oxygen by<br>nasal canula during sleep | recurrent apnea,<br>tracheomalacia with<br>intermittent stridor               |

(Continued on next page)

| Family<br>ID                                   | Family A<br>Individual 1  | Family B<br>Individual 2                      | Family C<br>Individual 3  | Family D<br>Individual 4  |
|--|---|---|---|---|
|  |   |   |   |   |
| Urinary tract                                  | bilateral vesicoureteric<br>reflux grade II   | normal  | normal  | vesicoureteric reflux, right<br>grade III / IV and left grade III   |
| Recurrent fever /<br>frequent viral infections | yes   | yes   | no  | yes   |
| Limbs  | drop foot   | normal  | soft tissue syndactyly of fingers<br>3 and 4 bilaterally, partial<br>syndactyly of toes 2, 3, and 4<br>on the right foot  | normal  |
| Craniofacial features                          | thick eyebrows, deep-set<br>eyes, short nasal bridge<br>with anteverted nares,<br>large mouth with<br>prominent lower lip,<br>protruding tongue | bushy eyebrows and a<br>low anterior hairline | flat occiput and<br>brachycephaly,<br>high anterior hair line with<br>frontal balding, tall forehead,<br>full cheeks, medial eyebrow<br>flare, left epicanthal fold,<br>pointed superior helices of<br>ears, upturned nasal tip | horizontal eyebrows,<br>low-set ears, depressed<br>nasal bridge, and large<br>mouth with prominent<br>lips, and slight retrognathia |
| Additional clinical findings                   | gingival hyperplasia  | mild hepatomegaly                             | No  | gingival hyperplasia  |

and HiSeq4000 (parents) system (Illumina) for sequencing. The average read depth was  $131 \times$ . Reads were aligned to the UCSC human reference assembly (hg19) with the Burrows-Wheeler algorithm (BWA v.0.5.9). More than 98% of the target sequences were covered to a read depth of at least  $20 \times$ . Single-nucleotide variants (SNVs) as well as small insertions and deletions were detected with SAMtools v.0.1.19. In-house custom Perl scripts were used for variant annotation.

In family B, trio ES was performed with the Sure Select Human All Exon Kit V6 on a HiSeq2500 sequencing machine (Illumina). Reads were aligned to the UCSC human reference assembly (hg19) with the BWA-MEM algorithm (BWA v.0.7.15). Variants were called with Genomic Analysis Toolkit (GATK) variant caller version 3.8; best practices as recommended by the Broad Institute were used.<sup>14</sup> Variant annotation was performed with KGGSeq.<sup>15</sup>

For family C, trio ES was done on a clinical basis through GeneDx with XomeDxPlus. A *RALGAPA1* homozygous variant was identified but classified as a variant of uncertain significance because no disease was known to be associated with it at that time. 100% of the coding region was covered for this gene at a minimum of 10×, with no indication of a multi-exon deletion or duplication involving this gene.

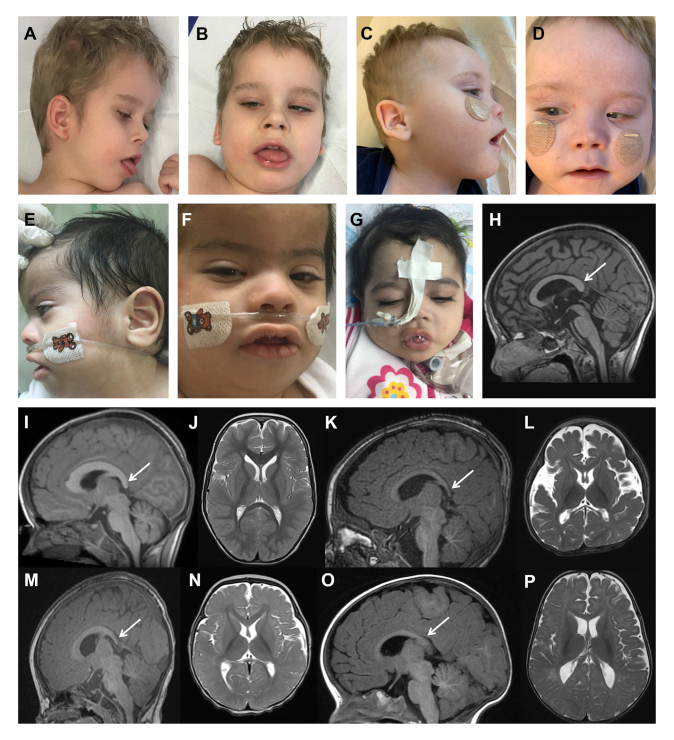
For family D, index-only ES was performed in parallel with array-based autozygosity mapping as described before.<sup>16</sup>

In family A, we filtered for homozygous and compoundheterozygous variants with a minor-allele frequency <0.1% in our in-house database of 15,000 individuals as well as for *de novo* variants that had a minor-allele frequency <0.01%.

A search for bi-allelic variants in family A identified changes in six genes (Table S1), whereas a filter for de novo variants did not identify protein altering variants. Among the biallelic variants, variants in NEB and BRCA1 were associated with autosomal recessive inherited conditions and were classified as variants of uncertain significance. Because there was no phenotypic overlap between the condition of our index individual and previously published individuals these variants were considered to not be associated with the child's condition. Of the remaining variants, only the compound-heterozygous variants in RALGAPA1 (GenBank: NM\_194301.4) were predicted to result in a loss of function c.3227A>G(p.Asn1076Ser) and c.1126C>T(p.Arg376\*). The parents were heterozygous carriers for one of the variants each. Variants were confirmed by Sanger sequencing in the index individual and his parents (Figure S1).

We next sought to identify further families with biallelic variants in *RALGAPA1* by using the web-based collaboration platform GeneMatcher.<sup>17</sup> We identified additional individuals who came from three unrelated families and had an overlapping phenotype and bi-allelic loss-of-function variants in *RALGAPA1* in Israel (family B), the USA (family C), and Saudi Arabia (family D, Figure 1). In family B, a homozygous frameshift variant, c.4992delT(p.Phe1664Leufs\*11), was detected; in family C, the homozygous nonsense variant c.610G>T(p.Glu204\*) was detected.

In family D, the combination of ES and array-based homozygosity mapping revealed a homozygous nonsense variant, c.5732C>G(p.Ser1911\*), within an autozygosity block (see Figure S4) that is unique to the affected



#### Figure 1. Clinical Details of Affected Individuals

(A-G) Clinical photographs of individuals with pathogenic variants in *RALGAPA1*. (A and B) Individual 1 at the age of 4  $\frac{1}{2}$  years. (C and D) Individual 3 at the age of 18 months. (E and F) Individual 4 at the age of 17 months. (G) Individual 2 at the age of 8 months. All children suffered from severe developmental disability with profound muscular hypotonia and feeding abnormalities. Facial features included upslanting palpebral fissures, mild epicanthus, a short nose with a flattened nasal bridge, and slightly anteverted nares. (H) Brain MRI image (T1-weighted sagittal view) of a healthy individual at the age of 4 1/2 years. The white arrow indicates the posterior part (splenium) of the corpus callosum.

(I–P) Brain MRI images of individuals with bi-allelic variants in *RALGAPA1* (T1-weighted sagittal as well as T2-weighted axial views). (I and J) Individual 1 (age 4 1/2 years). (K and L) Individual 2 (age 8 months). (M and N) Individual 3 (age 10 months). (O and P) Individual 4 (age 8 months). Of note, in all individuals with pathogenic variants in *RALGAPA1*, a moderate dysplasia of the corpus callosum with focal thinning of the posterior part was detectable (white arrows; compare with normal findings in panel [H]). Further details (including age-related normal values) regarding this finding are depicted in Figure S2. In addition, individual 2 showed global brain atrophy (L). Other variable or unspecific CNS abnormalities in individuals with RalGAPA1 deficiency are depicted in Figure S3. individuals. Additional information on the *RALGAPA1* variants can be found in Table S2.

To further support the relevance of *RALGAPA1* variants, we used living skin fibroblasts derived from the affected individuals from families A and B. Cell lines were cultured in Dulbecco's modified Eagle's medium (DMEM; life technologies) supplemented with 10% fetal bovine serum (FBS)(Life Technologies) and 1% penicillin-streptomycin (Life Technologies) at 37°C in a humidified atmosphere of 5% CO<sub>2</sub>.

For immunoblotting, whole-cell lysates were prepared with cell-lysis reagent CelLyticM (Sigma-Aldrich). A monoclonal antibody against RalGAPA1 (rabbit; 1:3000; EPR13592-38; Abcam) was used. GAPDH served as a loading marker (mouse; 1:4000; Thermo Fisher Scientific, catalog number AM4300). BM chemiluminescence blotting substrate (Roche) was used for visualization. Methodological details can be also found in the Supplemental Information. As depicted in Figure 2, a RalGAPA1 protein signal was nearly absent in fibroblasts of affected individuals, suggesting a loss-of-function effect in the investigated cell lines. To determine whether the variants result in nonsense-mediated decay (NMD), we additionally performed qRT-PCR for RALGAPA1 expression for individual 1, and this analysis revealed a reduction to 54% of the transcriptional level (Figure S5). This indicates that transcripts with the nonsense variant undergo NMD, whereas the missense variant is expressed, and reduced protein amounts are rather a consequence of protein instability.

As a next step, we aimed to gain further insight into the functional consequences of RalGAPA1 deficiency. Of note, the RalGAP complex facilitates GTP hydrolysis on RalA and thus converts RalA into an inactive GDP-bound form. The loss of RalGAP subunits in mice results in constitutively active GTP-bound RalA.<sup>18</sup> This can be exemplified by the deletion of the catalytic subunit RalGAPA2 in the bladder or RalGAPA1 in muscles, as well as the deletion of scaffolding subunit RalGAPB in adipose tissue.<sup>8,18,19</sup> Thus, we tested RalA activation status in human fibroblasts with RalGAPA1 deficiency. For RalA activity measurements, the fibroblasts were starved in FBS-free media and then in PBS. Fresh cell lysates were used for GTP-bound RalA pulldown with RalBP1 beads (Millipore) according to the manual. Samples were diluted in SDS sample buffer with 50 mM DTT, boiled for 5 min, resolved on SDS-PAGE, and transferred to nitrocellulose membranes (Bio-Rad). Individual proteins were detected with the specific antibodies and visualized on film via horseradish-peroxidaseconjugated secondary antibodies and Western Lightning Enhanced Chemiluminescence (Perkin Elmer Life Sciences). Further details are explained in the Supplemental Material and Methods.

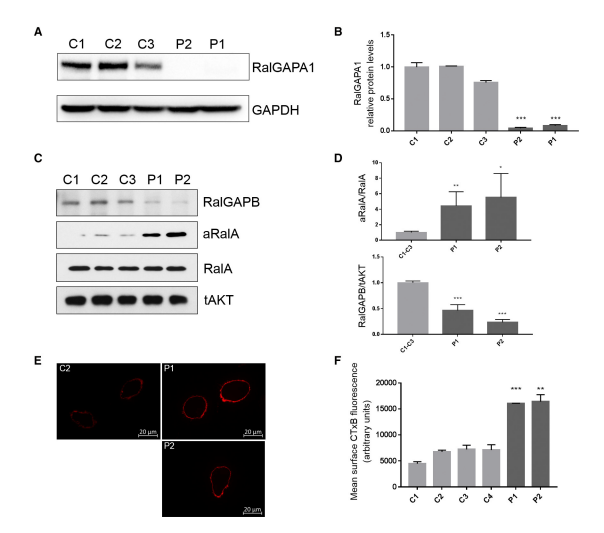
In fibroblasts with RalGAPA1 deficiency, RalA activity was increased in comparison to that in control fibroblasts (Figure 2). This finding is in line with the idea that RalGAPA1 deficiency causes constitutive activation of RalA. Additionally, amounts of RalGAPB, a scaffolding sub-

unit of RalGAP complex, were dramatically reduced in samples without the RalGAPA1 subunit, indicating a dysfunctional RalGAP complex and the necessity of RalGAPA1 for RalGAPB stability. *RALGAPB* expression analysis via RT-qPCR revealed no downregulation at the transcriptional level, supporting this hypothesis (see Figure S6).

Constitutive activation of the RalA pathway might have important consequences for cell physiology. Importantly, it has been shown that constitutively active RalA retains lipid-raft microdomains in detached cells at the plasma membrane and promotes anchorage-independent growth signaling.<sup>20</sup> It was suggested that this mechanism might play a role in tumor and metastasis formation. To further investigate this aspect in individuals with RalGAPA1 deficiency, we analyzed the localization of lipid-raft components in detached fibroblasts by using the fluorescence marker CTxB-Alexa 594 (for details, see Supplemental Information). Of note, CTxB (cholera toxin B subunit) constitutes the membrane-binding subunit of cholera toxins and serves as a well-established marker of lipid rafts.<sup>21</sup> CTxB Alexa 594 cell-surface fluorescence was visualized by ZEISS ApoTome imaging, and quantitative data were obtained by fluorescence-activated cell sorting (FACS) analysis. As depicted in Figure 2, fibroblasts with variants in RALGAPA1 retained significantly higher levels of surface lipid-raft components than did several control cell lines. Raw data that were used for quantification can be found in Figure S7. FACS analysis of CTxB-stained fibroblasts confirmed these results with a shift toward higher fluorescence intensity, indicating plasma membrane retention of lipid rafts in RalGAPA1-deficient fibroblasts (see also Figure S8). This finding is in agreement with previous research and indicates that cell-growth signaling might be altered in individuals with RalGAPA1 defiency.<sup>20</sup>

Taken together, the experiments described above confirmed a loss-of-function character of the RALGAPA1 variants and are in line with an autosomal-recessive inheritance pattern. Given the loss-of-function nature of the variants we describe here and the strikingly consistent associated phenotype, we conclude that the loss of RalGAPA1 in the individuals reported here is most likely responsible for the observed clinical abnormalities. This hypothesis is strengthened by the gnomAD geneconstraint metrics showing a depletion of loss-of-function and missense variants (observed-expected ratio of loss-offunction variants in RALGAPA1 = 0.12; 0.69 for missense variants).<sup>22</sup> This finding indicates that RALGAPA1 is a disease-associated gene. In light of the constraint metrics, the identification of one missense variant in our study group suggests that both loss-of-function and missense variants can cause disease.

Regarding the functional consequences of RalGAPA1 deficiency, it is suggestive that alterations in intracellular trafficking downstream of constitutively active RalA and possibly RalB are involved in disease pathophysiology. However, cell physiological abnormalities caused by



#### Figure 2. Functional Consequences of RalGAPA1 Deficiency

(A) Representative immunoblot image of fibroblasts derived from two individuals with pathogenic variants in *RALGAPA1* (P1 [derived from individual 1] and P2 [derived from individual 2]) and three control fibroblast lines (C1–C3). GAPDH served as a loading marker. Molecular weight: RalGAPA1 = 230 kDa; GAPDH = 36 kDa.

(B) Quantitated density of immunoblot bands from four independent experiments. Average values are presented as the mean  $\pm$  SEM; \*\*\*p < 0.001 significantly different from controls. In P1 and P2 fibroblasts, RalGAPA1 amounts are drastically reduced.

(C) Representative immunoblot image of the amounts of active GTP-bound RalA (aRalA) (these amounts were determined by a pulldown of aRalA with RalBP1 beads) in comparison to RalGAPB, RalA, and tAKT. Molecular weight: RalGAPB = 165.2 kDa; RalA = 23.5 kDa; and AKT = 55.7 kDa.

(D) Quantification of immunoblot results from three independent experiments. Average values are presented as means  $\pm$  SEM; \*p < 0.05, \*\*p < 0.01, and \*\*\*p < 0.001 significantly different from controls (C1–C3, pooled control). Amounts of GTP-bound RalA are increased in P1 and P2 cell lines compared to controls (C1–C3). Moreover, amounts of RalGAPB are reduced in P1 and P2 cell lines. The amounts of AKT (AKT serine-threonine kinase 1) are not altered.

(E) Microscope images (optical sections) of CTxB Alexa 594 cell-surface fluorescence in detached human skin fibroblasts. Images were obtained with an Axio Observer Z1 microscope with ApoTome 2 ( $63 \times$  objective lens, Zeiss). Images depict typical examples of control fibroblasts (C2) and two fibroblast lines derived from affected individuals (P1 and P2).

(F) Quantification of CTxB Alexa 594 cell-surface fluorescence by FACS analysis. Results depicted represent data from three independent experiments. Average values are presented as means  $\pm$  SEM; \*\*p < 0.01 and \*\*\*p < 0.001 significantly different from controls. P1 and P2 fibroblasts show higher levels of CTxB Alexa 594 fluorescence, indicating a disturbance of lipid-raft clearance from the plasma membrane in detached cells.

constitutively active RalA will be highly dependent on cell type. Which cell types are affected is dependent on whether RalGAPA1 is a major catalytic subunit of the RalGAP complex in the respective tissue because RalGAPA2 might be predominant. In light of the above results, RalGAPA1 deficiency appears to have a crucial impact on neuronal function and development. Directly after birth, all individuals identified in this study displayed neuromuscular abnormalities (e.g., respiratory problems, feeding difficulties, muscular hypotonia) that later developed into severe psychomotor disability. Three individuals developed early-onset epilepsy. All children had mild-tomoderate dysmorphic facial features, and one had soft tissue syndactyly of fingers (see Figure 1 and Figure S9). Brain MRIs revealed dysplasia of the corpus callosum as well as additional variable CNS manifestations (e.g., gray-matter heterotopias, ectopic posterior pituitary) that suggested an impact on brain maturation (Figure 1 and Figures S2 and S3). Characterization of the *RALGAPA1* homolog in mice (*Ralgapa1*, also known as *GRIPE*) indicated that its expression is upregulated in the developing brain during embryogenesis and remains high in the adult brain, especially in neurons of the *cortex* and hippocampus.<sup>23</sup> In addition, knockdown of the *RALGAPA1* homolog in zebrafish resulted in severe brain developmental delay.<sup>12</sup> Finally, RalGAPB knockout (which also leads to RalA activation) in mice is embryonic lethal (unpublished data). These data highlight the importance of *RALGAPA1* for neuronal development and differentiation.

On the basis of the complexity of the RalA pathway and its downstream effectors, the mechanisms behind the impact of disturbed RalA signaling on the nervous system will be difficult to dissect. Nevertheless, one important experimental result might be the disturbance that we observed in lipid-raft microdomains in RalGAPA1-deficient fibroblasts. It is well known that lipid remodeling is crucial for the development of the central nervous system via compartmentalization, signal transduction, cell adhesion, and axonal spreading.<sup>24,25</sup> Accordingly, altered lipid-raft trafficking might be a critical aspect of neuronal development in individuals with *RALGAPA1* variants.

Interestingly, the pathophysiology of RalGAPA1 deficiency might be at least in part similar to the mechanism of RALA variants published by Hiatt et al., 2018.<sup>11</sup> The majority of RALA variants cluster in the GDP-GTP binding pocket and are thus predicted to impair GDP-GTP binding. Three out of four RALA mutants that were tested in biochemical assays had a dramatic reduction in effector binding, and all mutants had a significantly reduced GTPase activity. The question of whether reduced effector binding together with decreased GTPase activity increases RalA signaling was not examined by the authors, but one RALA mutant was definitely constitutively activated, a finding similar to the RalA activation that we observed in RalGAPA1-deficient fibroblasts in this study. In general, the phenotype of the individuals with RALA variants was milder than in the children reported here. However, the core phenotype observed in individuals with variants in RALGAPA1 resembles the phenotype of individuals with RALA variants in previous studies (e.g., this phenotype includes intellectual disability, developmental delay, muscular hypotonia, and early-onset epilepsy). Furthermore, facial abnormalities (e.g., a short nose, a flattened nasal bridge, and gingival hypertrophy) appear to be a common feature. In addition, more than half of the children reported as having RALA variants had a dysplastic corpus callosum (shortening and/or thinning), which appears to be also a unifying MRI feature in individuals with RalGAPA1 deficiency. Interestingly, in some of the individuals with RALA variants, finger abnormalities, as also

seen in individual 3, were observed.<sup>11</sup> These striking similarities also pose an additional argument that *de novo* mutations in *RALA* might have the same downstream effect as the loss of *RALGAPA1*.

In light of the overlapping phenotype between *RALA*and *RALGAPA1*-associated diseases, it becomes evident that dysregulation of the RalA signaling pathway leads to a distinct group of genetic syndromes that we suggest could be named RALopathies. This group of disorders could be expanded by the identification of further *RALA*related disease genes.

In addition to the neurological disease manifestation, profound muscular hypotonia was a unifying clinical feature of affected individuals. This is of interest because RalA and the RalGAP complex have been implicated in the regulation of glucose uptake in response to insulin.<sup>8</sup> In mice, muscle-specific Ralgapa1 deletion caused RalA activation, leading to increased glucose and the uptake of free fatty acids by muscles, thus improving whole body glucose handling and lipid homeostasis. However, the study did not address the impact of RalGAPA1 deletion on muscle physiology or contractile ability. Additionally, the genetic approach used in that study did not allow the authors to address the role of RalGAPA1 in muscle development because Cre recombinase under myosin light chain 1f promoter is expressed in mature muscle only. No abnormalities in glucose metabolism were identified in the children we report on here.

Another disease feature of RalGAPA1-deficient individuals was recurrent episodes of fever with or without viral infections, as was reported in three individuals. This might be related to the severe developmental disability (e.g., increased risk for infections as a result of immobility; risk of aspirations). However, fever episodes and viral infections started so early in life and occurred at such a high frequency that they triggered extensive immunological work-up in two of the individuals. In this context, it is important to note that the *RALGAPA1* homolog in mice binds E12, a transcriptional regulator of immunoglobulin genes.<sup>23</sup> Therefore, although no specific immune defect could be identified in the affected individuals, a possible role of *RALGAPA1* in immune function might be expected.

Finally, on the basis of the overactivation of RalA in autosomal-recessive RalGAPA1 deficiency, it is tempting to speculate that inhibition of the Ral pathway via the inhibition of Ral prenylation<sup>26</sup> or the phosphorylation of RalA by Aurora A or protein kinase  $C^{27}$  might have a therapeutic effect. However, it will be very difficult to titrate the effect of such inhibitor approaches (especially against the background wherein RalA downregulation might also cause a neurological phenotype). Moreover, in light of the neonatal disease onset, it is suggestive that the variant already affects prenatal development in a way that might not be compensated for by a postnatal start of treatment. Further studies in knockout animal models of *RALGAPA1* will be helpful in addressing this issue.

#### Accession Numbers

The accession numbers for the data reported in this paper are as follows: for individual 1, ClinVar: VCV000691798 and VCV000691794; for individual 2, ClinVar: VCV000691795; for individual 3, ClinVar: VCV000691796; and for individual 4, ClinVar: VCV000691797.

### Supplemental Data

Supplemental Data can be found online at https://doi.org/10. 1016/j.ajhg.2020.01.002.

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## **Declaration of Interests**

The authors declare no competing interests.

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### Web Resources

ClinVar, https://www.ncbi.nlm.nih.gov/clinvar/ GeneMatcher, https://www.genematcher.org gnomAD Browser, https://gnomad.broadinstitute.org/ OMIM, https://www.omim.org/

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