Activating RAC1 variants in the switch II region cause a developmental syndrome and alter neuronal morphology

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RAC1 is a highly conserved Rho GTPase critical for many cellular and developmental processes. De novo missense RAC1 variants cause a highly variable neurodevelopmental disorder. Some of these variants have previously been shown to have a dominant negative effect. Most previously reported patients with this disorder have either severe microcephaly or severe macrocephaly.

Here, we describe eight patients with pathogenic missense RAC1 variants affecting residues between Q61 and R68 within the switch II region of RAC1. These patients display variable combinations of developmental delay, intellectual disability, brain anomalies such as polymicrogyria and cardiovascular defects with normocephaly or relatively milder micro- or macrocephaly. Pulldown assays, NIH3T3 fibroblast spreading assays and staining for activated PAK1/2/3 and WAVE2 suggest that these variants increase RAC1 activity and over-activate downstream signalling targets. Axons of neurons isolated from Drosophila embryos expressing the most common of the activating variants are significantly shorter, with an increased density of filopodial protrusions. In vivo, these embryos exhibit frequent defects in axonal organization. Class IV dendritic arborization neurons expressing this variant exhibit a significant reduction in the total area of the dendritic arbour, increased branching and failure of self-avoidance. RNAi knock down of the WAVE regulatory complex component Cyfip significantly rescues these morphological defects. These results establish that activating substitutions affecting residues Q61–R68 within the switch II region of RAC1 cause a developmental syndrome. Our findings reveal that these variants cause altered downstream signalling, resulting in abnormal neuronal morphology and reveal the WAVE regulatory complex/Arp2/3 pathway as a possible therapeutic target for activating RAC1 variants. These insights also have the potential to inform the mechanism and therapy for other disorders caused by variants in genes encoding other Rho GTPases, their regulators and downstream effectors.

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Introduction

RAC1 is a highly conserved Rho GTPase that switches between an inactive GDP-bound state and an active GTP-bound state, in which it can bind to and regulate the function of numerous cellular proteins.1 RAC1 plays several roles in the development and function of the nervous system, including regulation of neuronal morphology and migration.2 Functional importance of RAC1 is emphasized by its high mutational constraint in the general population.3 We recently described de novo missense RAC1 variants, spread across the gene, resulting in a highly variable neurodevelopmental disorder (RAC1-NDD) (OMIM 617751).4 Notably, the head circumferences of the patients with RAC1-NDD ranged from −5 to +4.5 SD. We provisionally divided RAC1-NDD patients into three broad phenotype-based groups of microcephalic, normocephalic and macrocephalic. We showed that some ‘microcephaly RAC1 variants’ are likely to have dominant-negative effects. Macrocephaly was noted in two patients, both with RAC1 substitutions affecting the residue V51. Normocephaly was noted in only one patient with a RAC1 Y64D variant. This variant was located in switch II, a region of RAC1 that undergoes significant structural changes when RAC1 transitions between the GTP and GDP bound states. This Y64D substitution was the only variant we found to be likely activating in our previous study.

Here, we describe eight patients with five distinct RAC1 variants leading to substitution of three different residues all within a region of switch II and show that this group of variants results in a developmental syndrome. Using Rac1-GTP pulldown assays, immunofluorescence and NIH3T3 spreading assays, we demonstrate that substitutions at these sites result in a higher proportion of RAC1 in the active GTP-bound state, increased RAC1 activity and signalling via multiple downstream effectors, including the RAC1/WAVE Regulatory Complex (WRC)/Arp2/3 pathway, and altered cell morphology. Using a Drosophila model, we show that the most common of these variants, Y64D, induces morphological changes in embryonic neurons, causes axon fasciculation defects in the embryonic CNS and increases branching of sensory neurons. Finally, we show that Rac1-Y64D-induced neuronal branching defects can be rescued in Drosophila by knockdown of the WRC component Cyfip, a homolog of human CYFIP1/2.

Materials and methods

Patient identification

We identified patients with RAC1 missense variants affecting the residues in the switch II region of RAC1 through the Deciphering Developmental Disorders study5 and via personal collaborations. The Central Manchester and Cambridge South NHS Research Ethics Committees approved this study (02/CM/238 and 10/H0305/83, respectively). InterVar (http://wintervar.wglab.org/) was used to apply the 2015 American College of Medical Genetics and Genomics (ACMG) guidelines for variant interpretation.6 A customized proforma was completed to collate relevant clinical information. Informed consent from patients or their legal representatives was obtained in all cases.
Rac1-GTP pulldown assays

Mutations were introduced into the coding region of human RAC1 using Quikchange Lightning (Agilent) according to the manufacturer’s instructions and subcloned in pRK5-myc using BamH1 and EcoR1. RAC1-GTP pulldown assays were performed as previously described. Briefly, HEK293 cells were transfected using Jet-OPTIMUS transfection reagent (Polyplus). Transfected cells were then incubated for 24 h before being lysed and active RAC1 isolated using the recombinant Cdc42- and RAC-interactive binding (CRIB) domain of p21 activated kinase (PAK) fused to GST. The ratio of isolated myc-RAC1-GTP to total myc-RAC1 was then assessed using western blotting, probing for the myc-tag (Cell Signalling Technology #2278), followed by a goat anti-rabbit HRP secondary antibody (BioRad) and visualized using enhanced chemiluminescence.

NIH3T3 spreading assays and immunofluorescence

Subconfluent NIH3T3 cells in a 24-well plate were transfected with RAC1 variants in pRK5-myc using Fugene 6 transfection reagent (Promega) according to the manufacturer’s instructions. Twenty-four hours later, cells were trypsinized and replated on 13 mm coverslips coated with fibronectin. Thirty minutes after replating, cells were fixed with 4% formaldehyde in PBS, then permeabilized with 0.1% Triton X-100 in PBS. Cells were then blocked with 1% BSA in PBS followed by incubation with primary antibodies in blocking buffer and secondary antibodies in PBS. Finally, coverslips were mounted using Prolong Gold antifade reagent with DAPI (Invitrogen) and imaged using a Leica DL600 widefield compound microscope with a 63×/1.40 NA oil objective. Antibodies used: Anti-myc mouse monoclonal (9E10) (Sigma-Aldrich) 1:500; Anti-phospho-Ser141 PAK-1/2/3 rabbit polyclonal (Thermo Fisher, 44-940G) 1:100; Anti-WAVE-2 (D2C8) rabbit monoclonal antibody (Cell Signalling Technology, 3659) 1:100; Alexa-568 donkey anti-rabbit secondary antibody (Invitrogen) 1:500; Alexa-488 donkey anti-mouse (Invitrogen) 1:500; Alexa-568-phalloidin (Invitrogen) at 1:500 and Alexa-568-phalloidin (Invitrogen) at 1:200 in PBT then mounted in Prolong Gold containing DAPI (Thermo Fisher). Cultures were imaged using a Nikon A1R confocal microscope, with a 60x Plan-Apo VC oil objective. Images were analysed using ImageJ.

Drosophila embryo immunohistochemistry

Embryos were dechorionated by immersion in 50% sodium hypochlorite (Sigma Aldrich) for 2 min, washed with water, then fixed in 1:1 4% formaldehyde in PBS:heptane for 20 min at room temperature. The aqueous phase was removed and one volume of methanol added to the heptane phase. The tube was shaken for 1 min and the heptane/methanol removed. Embryos were washed with methanol then PBT, then blocked with 2% BSA in PBT. Embryos were stained with anti-FasII (DSHB) at 1:100 followed by anti-mouse Alexa-488 (Invitrogen) at 1:500, Alexa-568-phalloidin (Invitrogen) at 1:200 in PBT then mounted in Prolong Gold containing DAPI (Thermo Fisher). Embryos were imaged using a Nikon A1R confocal microscope, with a 20x Plan-Apo VC objective. Images were analysed and processed using ImageJ and Adobe Photoshop. Segments were scored positive for a fasciculation defect if the defect extended ≥ half segment length.

Data availability

Experimental data will be shared on reasonable request from qualified investigators.
Results

Substitutions in the Q61-R68 region of RAC1 cause a developmental syndrome

Switch II comprises approximately residues 57–75 of RAC1 and is a conserved structural feature of small GTPases that undergoes a conformational change on exchange of GDP for GTP.\(^6\) Based on our previous analysis of a single normocephalic patient with a Y64D RAC1 variant, we hypothesized that some variants in the switch II region may cause a disorder distinct from the microcephalic and macrocephalic forms of RAC1-NDD.\(^4\) Through collaborations and the Deciphering Developmental Disorders study, we identified eight patients (including the original patient in our previous study\(^5\)) with missense variants affecting residues between Q61 and R68 of RAC1 (Table 1). These included four patients with the identical Y64D variant and one patient each with Q61E, Y64C, R68S and R68G variants. In all individuals, where investigation of inheritance was possible, the variants were found to have occurred de novo. All the variants affect amino acids that are highly conserved across orthologs, paralogs and the wider RAS superfamily (Fig. 1A). No variants affecting these residues are found in control databases. All variants affecting residues 61 or 64 were classified as pathogenic according to the ACMG variant interpretation guidelines (Supplementary Table 1). The two variants affecting residue R68 were initially classified as variants of uncertain significance because both parents could not be tested to confirm the variants to be de novo in these cases (Table 1). However, if the results of functional studies described later in the manuscript are taken into account, these variants could be reclassified as likely pathogenic (Supplementary Table 1). Switch II is the primary binding site for upstream regulators and also contains the G3 box which is involved in GTP hydrolysis.\(^10\) Q61, mutated to glutamate in one patient, has a well-established role in GTP hydrolysis, and substitution of this residue has frequently been observed to activate small GTPases, including RAC1.\(^11-13\) Y64 is not directly involved in nucleotide binding or hydrolysis, but its side chain is exposed on the surface of switch II (Fig. 1B), so substitution of this residue to aspartate or cysteine may affect the ability of RAC1 to interact with upstream regulators. The side chain of R68 is predicted to form hydrogen bonds with several other residues within and close to switch II (Fig. 1B), so substitution of R68 to glycine or serine is likely to alter the structure of switch II, potentially affecting GTP hydrolysis or interaction with upstream regulators. These data suggest that RAC1 variants affecting the residues Q61, Y64 and R68 identified in this study are pathogenic or likely pathogenic.

Next, we studied the clinical phenotypes of the eight patients, including the one patient with Y64D reported in a previous publication.\(^4\) This cohort comprises of five females and three males aged between 20 months and 15 years. Birth head circumference was known in only one case and was within the normal range. Most recent head circumferences were reported to be within normal range in six out of eight patients. One girl with R68S, born pre-maturely in the 26th week of gestation, had mild microcephaly of –2.2 SD at 4 years 5 months of age. One boy with a Y64D variant had mild macrocephaly of +3.06 at 14 years of age. Age of independent walking was delayed in five out of seven children for whom information on motor milestones was available. Although formal assessments were not available for these patients, the recruiting clinicians reported cognitive delay or intellectual disabilities ranging from mild to severe in all patients. None of the affected individuals were reported to have epilepsy. Brain MRI scans showed polymicrogyria in two patients, one with Y64C and one with the Y64D variant (Fig. 1C). Brain MRI scans were not performed in two individuals. Cardiovascular anomalies, such as ventricular septal defects, atrial septal defects and patent ductus arteriosus, were noted in four of eight patients. The patient with Q61E variant was also noted to be significantly obese, with a weight of >6 SD at the age of 2.5 years.

Comparison with previously published V51 RAC1-NDD patients showed that, none of the patients in the current cohort had severe macrocephaly (Table 2). Comparison with all other previously reported (likely dominant negative) RAC1-NDD patients showed that none of the patients in the current cohort had severe macrocephaly or epilepsy (Table 2). Although the patient numbers in the three groups are still small, collectively, these data suggest that RAC1 variants affecting residues Q61, Y64 and R68 in the switch II region result in a developmental syndrome with a phenotypic spectrum that is different from RAC1-NDD caused by other known RAC1 variants.

RAC1 variants affecting Q61–R68 of switch II increase RAC1 activity

To investigate whether the variants described above alter the proportion of cellular RAC1 in an active state, we performed RAC1-GTP pulldown assays using lysates of unstimulated HEK293 cells expressing patient variants or control RAC1 constructs. Three of the five patient variants were selected for this analysis: Y64D, Y64C and R68G. The level of RAC1-GTP pulled down by the RAC binding domain of PAK was analysed by western blot and normalized to the total level of RAC1 in the lysate. A previously characterized constitutively active variant of RAC1 (Q61L) exhibited increased levels of GTP-binding relative to RAC1 wild-type (WT), while a known dominant negative variant (T17N) exhibited reduced GTP binding. All three of the patient variants analysed exhibited significantly increased GTP binding relative to RAC1-WT (Fig. 2A and B). The increase in RAC1-GTP levels varied between the different variants, with the R68G variant exhibiting the largest increase. One of the two patient variants not analysed here, Q61E, has previously been shown to increase RAC1-GTP levels in a similar pull-down assay.\(^13\) We, therefore, conclude that patient variants affecting residues Q61, Y64 and R68 all increase levels of the activated GTP-bound form of RAC1.

RAC1 promotes the formation of lamellipodia and membrane ruffles around the cell periphery as cells spread, so the morphology of spreading cells is a useful cellular readout of RAC1 function. Constitutively active RAC1 (Q61L) promotes the formation of lamellipodia around the whole cell periphery, causing fibroblasts to exhibit a circular morphology, while dominant negative RAC1 (T17N) suppresses lamellipodia, leading to a stellate morphology dominated by filopodia (Fig. 2C and D). We expressed RAC1-Y64D, -Y64C and -R68G in NIH3T3 fibroblasts and examined their effect on the morphology of cells spreading on a fibronectin-coated surface. We found that cells expressing these variants typically exhibited a circular morphology reminiscent of cells expressing constitutively active RAC1, with the cell periphery dominated by lamellipodia and membrane ruffles (Fig. 2C and D). Cell circularity index (4π x area/perimeter\(^2\)) provides a simple means of quantifying cell morphology, and we found that RAC1-Y64D, -Y64C and -R68G all induced significant increased circularity relative to RAC1-WT, with circularity values for Y64C and R68G close to those induced by Q61L (Fig. 2D). The variants all exhibited increased localization to the cell periphery relative to RAC1-WT (Fig. 2C, arrows), and...
Table 1 Clinical features of patients with activating RAC1 variants in switch II region

| Patient No. | 1 (from Reijnders et al.) | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
|-------------|---------------------------|---|---|---|---|---|---|---|
| Sex, age    | Female, 2 years 6 months | Male, 12 years | Female, 5 years | Male, 14 years | Female, 3 years | Female, 3 years | Female, 5 years | Male, 15 years |
| RAC1 variant | De novo c.181C>G [p.(Gln61Glu)] | De novo c.190T>G [p.(Tyr64Asp)] | One of non-identical twins. Mild neonatal hypotonia. | De novo c.190T>G [p.(Tyr64Asp)] | None | De novo c.191A>G [p.(Tyr64Cys)] | RSV bronchiolitis at age 1 month | c.302C>A [p.(Arg68Set)] |
| Significant prenatal, birth or neonatal history | Neonatal feeding difficulties | Neutropenia | De novo | None | De novo | De novo | Born at 25 weeks 4 days. Neonatal feeding difficulties. Grade 1 IVH | c.302C>G [p.(Arg68Gly)] |
| Height, age (SD) | 95.9 cm, 2 years 6 months (+1.6) | 159.4 cm, 12 years (+0.9) | 88.6 cm, 3 years 4 months (−2.0) | 165 cm, 14 years (+0.3) | 93.5 cm, 3 years 9 months (−1.7) | 87.5 cm, 3 years (−1.7) | 95.5 cm, 5 years 1 month (+2.3) | 148.5 cm, 13 years (−0.8) |
| Weight, age (SD) | 24.4 kg, 2 years 6 months (+5.0) | Unknown | 11.7 kg (3 years 4 months) (−2.2) | 65 kg (14 years) (+1.4) | 11.7 kg (3 years 9 months) (−2.5) | 10.4 kg (3 years 2 months) (−1.3) | 14.2 kg, 5 years 2 months (−2.2) | 33.4 kg, 13 years (−1.5) |
| OFC cm (age) (SD) | 47.4 cm (2 years 6 months) (−2.0) | 56.5 cm (12 years) (+0.7) | 50.8 cm (5 years) (−0.8) | 61.5 cm (14 years) (+3.1) | 49 cm (3 years 9 months) (−0.2) | 49.0 cm (22 months) (−1.6) | 54.7 cm (4 years 5 months) (−2.2) | 54 cm (13 years) (−0.9) |
| Independent sitting | 10 months | 19 months | Unknown | Unknown | Delayed | Unknown | 12 months | Unknown |
| Independent walking | 20 months | 4 years | Unknown | 2 years 6 months | 1 year 6 months | Unknown | 2 years | 3 years |
| DD or ID | Mild-moderate | Severe | Hand stereotypes | Moderate Hypotonia and poor balance | Mild Autistic features, dyspraxia, ADHD | Mild Sleep difficulties | Mild Hypotonia, poor motor coordination, frequent falls, stereotypies | Moderate Mild sensory issues and hypotonia |
| Behavioural and neurological features | | | | | | | | |
| Brain MRI features | Normal | PMG | Not performed | Chiari I malformation | Not performed | Bilateral PMG, thin CC, thick WM | Short CC, prominent superior vermis fissures and 4th ventricular outlet | Bilateral CP cysts, prominent perivascular spaces, increased WM T2 FLAIR signal intensity |
| Vision or hearing features | Normal | Mild visual impairment, bilateral SNHL | Normal | Unilateral mild-moderate SNHL | Unilateral congenital cataract | Anisocoria, ROP | Frequent middle ear effusions | None |
| CVS features | ASD, VSD | Prominent metopic suture and nasal bridge, wave shaped PSs, low columella, dysplastic ear, high palate | None | Prominent glabella, macrostomia, micrognathia | Chronic OM, mild-moderate CHL | Unilateral microphthalmia | Arched eyebrows, broad nasal bridge, broad uvula, retrognathia | None |
| Cerebrofacial features | Normal | Mild hypotonia, prominent metopic suture and nasal bridge, small ears and orbits, dysplastic ears, high palate | Normal | Unilateral mild-moderate SNHL | Prominent glabella, mild hypertelorism, epicanthus | Anisocoria, ROP | None | None |
| Other observations | Keratosis pilaris, milia | Keratosis pilaris, milia | Keratosis pilaris, milia | Keratosis pilaris, milia | Keratosis pilaris, milia | Keratosis pilaris, milia | Keratosis pilaris, milia | Keratosis pilaris, milia |
| ADHD = attention deficit hyperactivity disorder; CC = corpus callosum; CHL = conductive hearing loss; CLD = chronic lung disease; CF = choroid plexus; CVS = cardiovascular; DD = developmental delay; FLAIR = fluid attenuated inversion recovery; GOR = gastro-esophageal reflux; ID = intellectual disability; IVH = intraventricular haemorrhage; KFS = Klippel-Feil syndrome; OFC = occipital frontal circumference; OM = otitis media; FDA = Patent ductus arteriosus; PF = palpebral fissure; PFO = patent foramen ovale; PMG = polymicrogyria; ROP = retinopathy of prematurity; RSV = respiratory syncytial virus; SNHL = sensorineural hearing loss; TI = tricuspid valve insufficiency; WM = white matter. | | | | | | | |
| aNM_006908.4. | | | | | | | |
| bBoth parents not available for testing. | | | | | | | |
| cVariant not present in mother and father not available for testing. | | | | | | | |
this peripheral localization was particularly pronounced for the Y64D and R68G variants.

Collectively, these data suggest that all the patient variants studied here increase the proportion of cellular RAC1 in an active GTP-bound state, increase RAC1 activity in cellular conditions and alter cell morphology.

Activating RAC1 switch II variants promote activation of downstream WRC and PAK signalling in fibroblasts

The WAVE regulatory complex is a downstream effector of RAC1 involved in regulating cell morphology in a variety of cell types including fibroblasts and neurons. Binding of active RAC1 enables the WRC to activate the Arp2/3 complex, a nucleator of actin filaments, and this RAC1/WRC/Arp2/3 pathway is implicated in regulating lamellipodia assembly in fibroblasts and axonal growth as well as initiation of collateral branches on axons and dendrites in neurons. To test the effect of expressing the activating switch II variants on this signalling pathway, we stained spreading NIH3T3 fibroblasts with antibodies against WAVE2, a component of the WRC. In cells expressing WT or dominant negative RAC1, WAVE2 was detected in the perinuclear region of the cell, but rarely at or close to the cell periphery. In contrast, cells expressing constitutively active RAC1, RAC1-Y64D, Y64C and R68G exhibited prominent WAVE2 localization at the cell periphery (Fig. 3A, arrowheads), consistent with increased recruitment and activation of the WRC at the lamellipodial leading edge by the activating switch II variants.

The PAK family of protein kinases are activated by binding of active RAC1 and are implicated in regulating a variety of cellular processes including cell survival, proliferation and metabolism. To test if this pathway is stimulated by the activating switch II variants we stained NIH3T3 fibroblasts expressing Y64D, Y64C and R68G with an antibody specific for the activated form of PAK1/2/3 phosphorylated at S141. While some staining was detected for activated PAK in cells expressing WT or dominant negative RAC1, this staining was restricted to the perinuclear region (Fig. 3B). In contrast, activated PAK was detected throughout cells expressing constitutive active RAC1 and all the switch II variants tested, including punctate accumulations at and close to the cell periphery (Fig. 3B, arrowheads), where little or no active PAK was observed in cells expressing WT or dominant negative RAC1 (Fig. 3B). Quantification of activated PAK staining close to the cell periphery suggested that levels of activated PAK were significantly elevated in this region for all switch II variants tested relative to WT RAC1, with levels highest for Y64C and R68G (Fig. 3C).

Collectively, these data provide further evidence that all the activating switch II variants studied here increase signalling through multiple downstream pathways including the WRC and PAK family kinases.

Rac1-Y64D induces morphological changes in Drosophila embryonic neurons

Given that all the activating switch II variants tested induce significant morphological changes in fibroblasts (Fig. 2C–E), we hypothesized that these variants may also alter the morphology of neurons and that this could contribute to disease phenotypes. Model organism studies have previously demonstrated a variety of defects in neuronal morphology, including axonal misrouting and altered dendritic branching when RAC1 function is impaired. The Drosophila homolog of RAC1, named Rac1, is 92% identical to its human equivalent, and Drosophila has proved an excellent model for dissecting the functions of RAC1 in the nervous system, so we chose this system to model the effects of activating switch II variants on neuronal morphology. Since Y64D is the most frequently observed variant in our cohort, it was chosen as an exemplar for activating switch II variants in this analysis. We thus generated a Drosophila strain harbouring a transgenic copy of Drosophila Rac1-Y64D under the control of the conditional UAS-Gal4 system, allowing targeted expression of this variant.

To examine the effect of Rac1-Y64D on axonal growth and morphology, we isolated neurons from stage 11 embryos expressing Rac1-Y64D under the control of the pan-neuronal elav-Gal4 driver and allowed the neurons to grow in culture for 6 h. At this developmental stage, axons are just beginning to extend, allowing us to capture the earliest effects of Rac1-Y64D on axonal growth and morphology. We found that the axons of neurons expressing Rac1-Y64D were around two-fold shorter than controls or those expressing Rac1-WT (Fig. 4A and B). Interestingly, Rac1-Y64D expression also resulted in an increase in the density of filopodial protrusions along the axon shaft, structures previously identified as precursors of collateral axonal branches (Fig. 4A and C, arrows). These results suggest that Rac1-Y64D reduces the rate of axonal extension and promotes increased formation of collateral branch precursors.

Rac1-Y64D induces axon fasciculation defects in the Drosophila embryonic CNS

Our finding that axonal growth and morphology is perturbed in cultured neurons expressing Rac1-Y64D prompted us to investigate whether we could detect axonal defects in the intact embryonic nervous system. To examine axonal organization within the CNS, we stained embryos expressing this Rac1 variant under the control of the pan-neuronal driver elav-Gal4 with an antibody against Fasciculin II (FasII). The axons of neurons expressing FasII bundle into six fascicles that run longitudinally within the nerve cord and any deviation from this arrangement is indicative of defects in axonal morphology or routing. We found that while expression of UAS-Rac1-WT did not cause any obvious changes in the fasciculation of FasII-positive axons, Rac1-Y64D expression resulted in frequent defects in axonal organization, including absence of fascicles, crossing of axons between fascicule tracks and defasciculation of axons (Fig. 4D and E).

Rac1-Y64D increases branching of sensory neurons

Since our data suggested that Rac1-Y64D may increase the formation of neuronal branch precursors, we investigated the effect of this variant on branching in more detail using an established model system for this process; class IV dendritic arborization (cIVda) neurons. This neuron class forms highly branched dendritic arbors, but since they exhibit self-avoidance and tilting, the morphology of individual cells can be accurately imaged and analysed in vivo. Rac1-Y64D was expressed specifically in cIVda neurons using the ppk-Gal4 driver alongside a fluorescent membrane marker to allow the dendritic arbor of these cells to be easily visualized. Expression of Rac1-Y64D resulted in significant changes in the morphology of cIVda neurons (Fig. 5A and B). Firstly, the total area of the dendritic arbor was reduced in Rac1-Y64D expressing cells. Secondly, the density of branching close to the cell body was elevated compared to controls (Fig. 5D and E). Thirdly, crossing...
Figure 1 RAC1 switch II variants cause a neurodevelopmental disorder. (A) Alignment of switch II region of RAC1 and related small GTPases. Residues in RAC1 affected by described variants are indicated in bold. (B) Structure of RAC1 bound to GTP analogue (Protein Database ID 1MH1). Switch II region indicated. Hydrogen bonds predicted to be formed between R68 and other residues in RAC1 are indicated by dashed lines. Image preparation and hydrogen bond prediction performed using UCSF ChimeraX. (C) T1-weighted brain MRI images of Patient 3 with the Y64C variant illustrating bilateral perisylvian polymicrogyria (white quadrilaterals) and thin corpus callosum (arrow).
over of dendrites was observed, demonstrating failure of self-avoidance, suggesting defective dendrite pathfinding.

Rac1-Y64D-induced neuronal branching defects can be rescued by knockdown of the WRC component Cyfip

A recent report demonstrated that dendritic branching by cIVda neurons is initiated by activation of the actin-nucleating Arp2/3 complex downstream of Rac1 and the WRC. Since our cell culture studies had suggested that Rac1-Y64D increases WRC activation, we hypothesized that the increased branching density observed on expression of Rac1-Y64D might result from excessive activation of the branch-promoting WRC/Arp2/3 complex pathway. To test this hypothesis, we used RNAi to knock down expression of the WRC component Cyfip (homolog of human CYFIP1/2) in Rac1-Y64D expressing neurons. We found that Cyfip knock-down rescued the morphological defects induced by Rac1-Y64D, in particular the increase in branches close to the cell body (Fig. 5A–F). We quantified dendritic architecture using Sholl analysis, and this indicated that the branching of neurons expressing Rac1-Y64D is significantly different to that of both control and Rac1-Y64D + Cyfip RNAi expressing neurons. Control neurons are not significantly different to Rac1-Y64D + Cyfip RNAi neurons, demonstrating that knockdown of Cyfip rescues the Rac1-Y64D phenotype (Fig. 5G). These results suggest that Rac1-Y64D induces defects in dendrite morphology by over-activating the WRC and that reducing WRC activity can suppress morphological defects induced by Rac1-Y64D.

Discussion

We had previously reported a number of germline RAC1 variants as causes for RAC1-NDD. The RAC1-NDD patients could be divided into broad categories according to their head circumferences. Although we presented multiple patients with either micro- or macrocephaly, we had identified only one patient, with a Rac1 Y64D variant, located in the switch II region, with a head circumference within the normal range. We now describe a total of eight patients with variants in other regions of RAC1. Four out of eight patients described here are less than 6 years of age and Patient No. 7 (Table 1) was born prematurely in the 26th week of gestation. This makes it difficult to compare the phenotypes of patients in this cohort. However, even within this group of patients with activating variants, there appears to be a substantial phenotype variability. For example, the developmental delay ranged from mild to severe and the head circumferences were between −2.2 and −3.0 SD. There were differences in severity of the clinical features of patients with identical Y64D variants, which suggests possible roles for genetic background or environmental factors in determining the phenotype of this condition.

In our original study, our only functional data regarding Y64D was that it induced morphological changes in cultured fibroblasts reminiscent of those induced by constitutively active Rac1, leading us to propose that this could be an activating variant. The current study considerably extends the functional understanding of this variant and others affecting nearby residues. We confirm that all of the switch II variants analysed here are activating by showing that they all increase levels of GTP-bound Rac1, all promote fibroblast spreading and all stimulate downstream signalling pathways. In our previous report, we described three variants affecting residues closely flanking the region of Rac1 explored in this study (V51M, V51L and P73L), and none of these variants showed any evidence of increased Rac1 activation. Overall, these observations suggest that variants affecting residues Q61–R68 form a distinct class of activating Rac1 variants with shared mechanistic features. Activating Rac1 mutations affecting residues outside of switch II have been identified in various cancers, so it is possible that the syndrome described here could potentially also be caused by variants outside of the Q61–R68 region.

Recently, a de novo germline E62K variant of RAC2, a Rac1 paralog expressed primarily in the hematopoietic cell lineages, was shown to cause severe T- and B-cell lymphopenia, myeloid dysfunction and recurrent respiratory infections through gain-of-function effects (OMIM 618986). Interestingly, de novo germline Q61L and E62K variants in RAC3 have also been shown to cause a neurodevelopmental disorder with structural brain anomalies and dysmorphic faces (OMIM 618577). However, no functional studies have yet been performed for the Rac3 variants. Germline Y64C, R66G and R68Q variants in CDC42, another Rho GTPase closely related to Rac1, have been described to cause Takenouchi-Kosaki syndrome (OMIM 616737), another neurodevelopmental disorder. Interestingly, Q61–R68 has also been identified as a somatic mutational hotspot in KRAS in colon and rectal cancers. Overall, our and previous results suggest that the Q61–R68 region of switch II may be prone to activating mutations throughout the RAS superfamily.

Table 2 Group-wise comparison of selected clinical features of RAC1-NDD patients

| Group | Q61–R68 RAC1-NDD | V51 RAC1-NDD | All other RAC1-NDD |
|-------|------------------|-------------|-------------------|
| Number of individuals | 8 | 2 | 4 |
| Head circumference SD | −2.2 to +3.06 | +4.1 to +4.5 | −2.5 to −5.0 |
| Epilepsy | None | None | Two |
| Main brain MRI findings | Polymicrogyria in two; Chiari 1 malformation in one; thin white matter anomalies in two | Cerebellar anomalies, thin corpus callosum and mega cisterna magna in three; enlarged lateral or 4th ventricles in two; thin brain stem in two | Two |

Table 1

| Variants | Number of individuals | Head circumference SD | Epilepsy | Main brain MRI findings |
|----------|-----------------------|-----------------------|----------|-------------------------|
| Q61E, Y64C, Y64D, R68S, R68G | Eight | −2.2 to +3.06 | None | None |
| V51L, V52M | Two | +4.1 to +4.5 | None | None |
| C18Y, N39S, P73L, C157Y | Four | −2.5 to −5.0 | Two | Two |

Table 2

| Variants | Number of individuals | Head circumference SD | Epilepsy | Main brain MRI findings |
|----------|-----------------------|-----------------------|----------|-------------------------|
| Q61E, Y64C, Y64D, R68S, R68G | Eight | −2.2 to +3.06 | None | None |
| V51L, V52M | Two | +4.1 to +4.5 | None | None |
| C18Y, N39S, P73L, C157Y | Four | −2.5 to −5.0 | Two | Two |
Figure 2 RAC1 switch II variants increase levels of GTP-bound RAC1 and alter fibroblast morphology. (A) Western blot of myc-RAC1-GTP (top panel) pulled down from lysates of HEK293 cells expressing indicated RAC1 variant using PAK-CRIB probed with anti-myc. Lower two panels show blots of raw lysates used in pulldowns probed with anti-myc to show total myc-RAC1 levels or anti-actin as loading control. Uncropped images of these blots are shown in Supplementary Fig. 1. (B) Quantitation of relative GTP-RAC1 levels for indicated RAC1 variants. Calculated by dividing GTP-RAC1 band intensity by total RAC1 intensity for each sample then normalizing to value obtained for constitutively active (Q61L) RAC1 in the same dataset. Bars indicate mean ± SEM. n = 7 independent experiments for all variants, except Y64C and R68G where n = 4. Data analysed in Graphpad Prism using mixed effects model. *P < 0.05. (C) Spreading NIH3T3 fibroblasts expressing indicated RAC1 variant stained with Alexa568-Phalloidin to label F-actin and anti-myc to label expressed RAC1 variant. Second row shows magnification of a section of the cell periphery in the above image. Arrows indicate localization of RAC1 variants to cell periphery. Scale bars in first column indicate 10 µm and apply to all images in the row. (D) Circularity of cells expressing indicated RAC1 variant. n > 50 cells pooled from three independent experiments. Line indicates mean value. Data statistically analysed using Kruskal-Wallis test with Dunn’s correction for multiple comparisons. ****P < 0.0001, ***P < 0.001, **P < 0.01, *P < 0.05, ns P > 0.05 relative to RAC1 WT. (E) Categorization of cell morphology based on predominant protrusion type. Cell scored as >50% lamellipodia or filopodia if this protrusion type occupies greater than 50% of cell periphery. n > 50 cells pooled from three independent experiments. CA = constitutively active; DN = dominant negative.
There are two likely explanations for pathogenic switch II variants causing activation: firstly, since switch II contains residues with a direct role in GTP hydrolysis, variants that change these residues or alter their position by inducing local conformational changes are likely to reduce the efficiency of GTP hydrolysis, thus increasing the proportion of cellular RAC1 in an activated.
GTP-bound state. Of the variants described here, Q61E is most likely to directly affect GTP hydrolysis since Q61 has a well-established role in the hydrolysis process.\textsuperscript{11} Although we did not perform functional studies for the Q61E variant, this substitution has previously been shown to increase RAC1 activity.\textsuperscript{13} The second possible mechanism is via altering interactions with regulators; for example, reduced affinity for GTPase activating proteins (GAPs) or guanine nucleotide dissociation inhibitors would be expected to increase basal activity of RAC1. For example, a E62K variant in RAC2 abolishes the ability of GAPs to accelerate GTP hydrolysis.\textsuperscript{25} The crystal structure of RAC1 bound to the GAP sptP identifies Y64 as a direct point of contact between the two proteins; therefore, it is possible that Y64D and Y64C variants increase RAC1 activity by impairing interaction with GAPs.\textsuperscript{31} R68 is predicted to form several hydrogen bonds that stabilize the structure of switch II, so substitution of this residue could impact GTP hydrolysis and/or regulator

Figure 4 Expression of Rac1-Y64D alters axon morphology and organization in Drosophila embryos. \textit{(A)} Representative images of cultured Drosophila embryonic neurons extracted from stage 11 embryos of indicated genotype. Stained for tubulin, F-actin and DAPI. Arrows indicate filopodia. Scale bar = 10 µm and applies to all three images. \textit{(B} and \textit{C}) Quantitation of axon length of cultured Drosophila embryonic neurons and density of filopodia along axon shaft. Eighty-five neurons analysed for each genotype extracted from 24 stage 11 embryos. Lines indicate mean values. Data statistically analysed using Kruskal–Wallis test with Dunn’s correction for multiple comparisons. * $P < 0.05$, **** $P < 0.0001$, ns $P > 0.05$ relative to control. \textit{(D)} Representative images of ventral nerve cord of stage 16 embryos with indicated genotype stained with anti-FasII to reveal axon fasciculation in developing CNS. Lower panels show magnified region from upper panels. Arrowheads show defasciculation and asterisks show fascicle breaks. Scale bars in first column = 10 µm and apply to all images in the row. \textit{(E)} Quantitation of number of segments in which fasciculation defects are observed in ventral nerve cord of stage 16 FasII-stained embryos. At least 19 embryos analysed for each genotype. Lines indicate mean values. Data statistically analysed using Kruskal–Wallis with Dunn’s correction for multiple comparisons. **** $P < 0.0001$, ns $P > 0.05$ relative to control.
binding by inducing local structural changes. Both the R68G and R68S variants would be expected to disrupt hydrogen bonding within switch II, so while we did not functionally analyze R68S, it is likely to have a similar effect on RAC1 activity to R68G. While there are clear mechanistic similarities between variants investigated here, the extent and nature of changes in GTP hydrolysis and protein interactions induced by each variant will differ, so their disease mechanisms are unlikely to be identical.

The increased activity observed for all of these variants is likely to be significant as increased or decreased RAC1 activation has also been detected in neurodevelopmental disorders caused by variants in TRIO and HACE1, supporting the notion that tight regulation of the level of RAC1 activity is important for proper neurodevelopment and function. The increase in RAC1-GTP levels we observe for Y64D, Y64C and R68G is smaller than for the constitutively active Q61L mutant, which is unable to hydrolyze GTP and is therefore

Figure 5 Expression of Rac1-Y64D alters dendritic branching in Drosophila sensory neurons. (A–C) Class IVda sensory neurons from dorsal surface of segments A1–A4 of L3 larvae of indicated genotype. Neurons visualized using CD8-mCherry expressing under the control of ppk-Gal4. Scale bar indicates 40 µm and applies to all three images. (D–F) Magnification of the area around the cell body for the cells shown in A–C. Scale bar = 40 µm and applies to all three images. (G) Sholl analysis of dendritic organization in which the number of times dendrites intercept a semicircle originating at the cell body is plotted against the radius of the semicircle. The semicircle comprises the region of the neuron that is dorsal to the cell body, corresponding to approximately the top half of the images shown in A–C. Graph shows mean ± SEM of ∼15 neurons for each data set. Statistical analysis by two-way ANOVA. ****P < 0.0001, ns P > 0.05 relative to control.
always in an active state. This mirrors the relative effects of these mutants on fibroblast morphology and PAK activation and suggests that the variants found in patients are still able to undergo a GTPase cycle, albeit with a higher percentage of molecules in an active state at any one time relative to WT RAC1. This likely also applies to Q61E, which has previously been shown to activate RAC1, but less strongly than Q61L. Although we did not functionally characterize every variant in this study, there is no obvious correlation between the extent of RAC1 activation in our functional studies and the severity of disease phenotypes for the variants studied.

In our Drosophila studies, we found that the Rac1-Y64D alters the morphology and growth of neuronal axons and dendrites. Specifically, we find that overall extension of axons and dendrites is reduced, while branching is increased. Rac1 activity is known to trigger branching of axons and dendrites, leading us to propose that the elevated levels of active Rac1-Y64D results in increased frequency of branch initiation events at the expense of axon/dendrite extension. Disorganization of axon fascicles in the CNS of Drosophila embryos expressing Rac1-Y64D is also consistent with altered axon morphology and/ or pathfinding being a contributing factor in phenotypes caused by this variant. Notably, the dendritic branching phenotype induced by Rac1-Y64D in Drosophila and humans and component of the WRC. This result suggests that the Rac1-Y64D branching phenotype is caused by overactivation of the WRC/Arp2/3 complex pathway. Germline variants in the genes encoding WRC components (e.g. CYFIP2 and WASF1) or for ubiquitous actins (e.g. ACTB and ACTG1) have been implicated in a variety of neurodevelopmental disorders. Of note, overactivation of the WRC/Arp2/3 complex pathway has been shown to underpin NDDs caused by variants in CYFIP2. As well as increasing our mechanistic understanding of the cellular and developmental effects of switch II variants, our ability to genetically rescue phenotypes induced by Rac1-Y64D identifies the WRC/Arp2/3 pathway as a possible pharmacological target for treatment of this and possibly other disorders in which RAC1 activity is elevated. However, further research is needed to explore whether these results can be reproduced in higher organisms and establish a possible therapeutic window for such interventions.

In summary, our results establish activating substitutions within the switch II region of RAC1 as a distinct cause of neurodevelopmental delay and delineate their clinical consequences, provide an insight into the underlying disease mechanism and reveal a possible therapeutic target. Furthermore, our findings suggest a potential mechanistic and therapeutic convergence between variants in RAC1 and several other related neurodevelopmental disease-genes involved in regulation of Rho-GTPases and the WRC/Arp2/3 pathway.

Acknowledgements
We are thankful to all patients and families for participating in the study. We are thankful to the Deciphering Developmental Disorders (DDD) study for the invaluable collaboration. We thank the University of Manchester Bioimaging and Fly Facilities for technical assistance.

Funding
The Deciphering Developmental Disorders Study (Cambridge South REC approval 10/H0305/83 and the Republic of Ireland REC GEN/284/12) presents independent research commissioned by the Health Innovation Challenge Fund (grant number HICF-1009-003), a parallel funding partnership between the Wellcome Trust and the Department of Health and the Wellcome Trust Sanger Institute (grant number WT098051). The views expressed in this publication are those of the author(s) and not necessarily those of the Wellcome Trust, BBSRC or the Department of Health. The research team acknowledges the support of the National Institute for Health Research, through the Comprehensive Clinical Research Network, UK. T.M. acknowledges financial support from the Wellcome Trust Grant no. 201958/Z/16/Z.

Competing interests
The authors report no competing interests.

Supplementary material
Supplementary material is available at Brain online.

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