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The Candidate Schizophrenia Risk Gene *DGCR2* Regulates Early Steps of Corticogenesis

Aude Molinard-Chenu and Alexandre Dayer

ABSTRACT

BACKGROUND: Alterations in early steps of cortical circuit assembly are thought to play a critical role in vulnerability to schizophrenia (SZ), but the pathogenic impact of SZ-risk mutations on corticogenesis remains to be determined. DiGeorge syndrome critical region 2 (*DGCR2*) is located in the 22q11.2 locus, whose deletion is a major risk factor for SZ. Moreover, exome sequencing of individuals with idiopathic SZ identified a rare missense mutation in *DGCR2*, further suggesting that *DGCR2* is involved in SZ.

METHODS: Here we investigated the function of *Dgcr2* and the pathogenic impact of the SZ-risk *DGCR2* mutation in mouse corticogenesis using in utero electroporation targeted to projection neurons.

RESULTS: *Dgcr2* knockdown impaired radial locomotion and final translocation of projection neurons, leading to persistent laminar positioning alterations. The *DGCR2* missense SZ-risk mutation had a pathogenic impact on projection neuron laminar allocation by reducing protein expression. Mechanistically, we identified *Dgcr2* as a novel member of the Reelin complex, regulating the phosphorylation of Reelin-dependent substrates and the expression of Reelin-dependent transcriptional targets.

CONCLUSIONS: Overall, this study provides biological evidence that the SZ-risk gene *DGCR2* regulates critical steps of early corticogenesis possibly through a Reelin-dependent mechanism. Additionally, we found that the SZ-risk mutation in *DGCR2* has a pathogenic impact on cortical formation by reducing protein expression level, suggesting a functional role for *DGCR2* haploinsufficiency in the 22q11.2 deletion syndrome.

Keywords: 22q11, Corticogenesis, *DGCR2*, Neuronal migration, Reelin, Schizophrenia

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Alterations in the formation of cortical circuits are involved in the pathophysiology of developmental disorders, including schizophrenia (SZ) (1,2). Numerous genes are associated with SZ (3,4) and many of them regulate cellular events involved in cortical circuit assembly (5,6). However, the precise biological mechanisms through which SZ-risk genes alter early cortical circuit formation remain unclear (7). Here, we investigate the impact of a novel SZ-risk gene, DiGeorge syndrome critical region 2 (*Dgcr2*), on cortical projection neuron (PN) development. *Dgcr2* is expressed throughout brain development (8), encodes for an activity-dependent adhesion protein (9,10), and is associated with SZ (3,11). *DGCR2* is located in the 22q11.2 locus, in the minimal critical region for 22q11.2 microdeletion syndrome (22q11.2DS) (12). This microdeletion is one of the highest known risk factors for SZ (13), and it has been suggested that *DGCR2* haploinsufficiency may contribute to increased risk for SZ in 22q11.2DS (11). Additionally, exome sequencing of family trios identified a rare de novo *DGCR2* mutation (P429R) in a patient with idiopathic SZ (3). Here we used in utero electroporation in mice (14,15) to determine whether *Dgcr2* impacts early steps of PN development and assessed the pathogenic impact of the *DGCR2* SZ-risk mutation in this model system.

Early cellular events involved in the development of cortical PNs are increasingly well understood (16,17). After being generated asymmetrically from radial glia in the ventricular zone, upper-layer PNs undergo sequential migratory stages until they reach their final laminar position within the cerebral cortex. A key stage in neuronal migration is radial glia-guided locomotion during which PNs migrate from the intermediate zone toward upper cortical layers (18). The fine positioning of PNs relies on an ultimate migratory step defined as final translocation (19). The precisely orchestrated migration of PNs appears to be a vulnerable cellular process that is affected by different types of psychiatric-related genetic insults (20–22).

Multiple cell-extrinsic regulators of PN migration have been identified (23,24). Among these factors, Reelin is an important signaling protein secreted by Cajal-Retzius cells (25). It controls the inside-out lamination of cortical PNs (26), and alterations in downstream signaling components of the Reelin pathway, such as *DAB1*, disturb radial glia-guided locomotion (27) and final translocation (20,28,29). Interestingly, the Reelin signaling cascade is associated with several SZ-related processes including *N*-methyl-D-aspartate receptor hypofunction (30–32) and SZ-like cognitive impairments (33–35).

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In this study, we find that short hairpin RNA (shRNA)-mediated knockdown (KD) of *Dgcr2* as well as an SZ-risk mutation in *DGCR2* affects the migration of PNs. Moreover, we identify *DGCR2* as a novel member of the Reelin complex and show that the phosphorylation of Reelin-dependent signaling targets is reduced following *Dgcr2* KD. Finally, using RNA sequencing of *Dgcr2*-KD PNs, we identify *Dgcr2*-dependent target genes, which are significantly enriched in Reelin-dependent genes. Overall, this study provides evidence that the SZ-risk gene *Dgcr2* regulates PN migration and positioning possibly through Reelin-dependent mechanisms.

METHODS AND MATERIALS

Plasmids, antibodies, and primers are discussed in [Supplemental Methods](#).

In Utero Electroporation

Animal experiments were conducted according to the Swiss and international guidelines and were approved by the local animal care committee. Embryos from time pregnant E14.5 CD1 mice or C57BL/6 (RNA sequencing experiment and [Figure 1A](#)) were electroporated in the lateral ventricular zone of the dorsal pallium as described previously ([36](#)). Sequential electroporations were performed at 24-hour intervals at E14.5 and E15.5.

Tissue Processing, Immunohistochemistry, and Proximity Ligation Assay

Dissected brains were fixed, stored in a cryoprotective solution if required, and processed for immunohistochemistry as described previously ([22,36](#)) and for proximity ligation assay (PLA) as described in [Trifileff et al. \(37\)](#) with Duolink reagents (Sigma-Aldrich, St. Louis, MO).

Cell Culture and Transfection

Human embryonic kidney (HEK) 293T cells were cultured in Dulbecco's modified Eagle medium (Life Technologies, Carlsbad, CA) supplemented with 10% fetal bovine serum (Biochrom, Berlin, Germany), 10 U/mL penicillin/streptomycin mixture (Life Technologies) at 37°C under 5% CO₂. Cells were transfected using lipofectamin-2000 (Thermo Fisher Scientific, Waltham, MA). Primary neuronal cultures were prepared as previously described ([36](#)).

Western Blot and Immunoprecipitation

Cells were scraped and lysed in cell lysis buffer B. The lysate was clarified by centrifugation and 50 µg of proteins were processed as in [Riccio et al. \(36\)](#). For the coimmunoprecipitation experiment, Reelin-enriched medium was produced and collected as described in [D'Arcangelo et al. \(38\)](#) and after 1-hour incubation with Reelin-enriched or mock medium, transfected HEK 293T cells were resuspended in 0.4% paraformaldehyde for 10 minutes at room temperature, centrifuged, and lysed with cell lysis buffer B. A total of 1 mg of proteins from the lysate was incubated overnight in phosphate-buffered saline (PBS) 0.01% Tween-T, with 20 µL of hemagglutinin (HA) tag (C29F4) monoclonal antibody-conjugated magnetic beads (Cell Signaling, Danvers, CT). Beads were washed 4 times with PBS 0.01% Tween-T before elution in Laemmli buffer. Eluates

were boiled at 100°C for 5 minutes, and Western blot was performed as previously described ([22](#)). The disabled homolog 1 (DAB1) phosphorylation assay was performed as in [Lee et al. \(39\)](#), except Reelin-enriched medium was used to stimulate primary cortical neurons instead of purified Reelin.

Fluorescence-Activated Cell Sorting, Reverse Transcriptase Quantitative Polymerase Chain Reaction, and RNA Sequencing

E17.5 cortices from embryos previously electroporated at E14.5 together with the fluorescent reporter tdTomato were dissected and processed until dissociation as previously described for neuronal cultures. Cells were resuspended in PBS, filtered through a cell strainer (70 µm, BD, Franklin Lakes, NJ), and sorted on a BD FACSAria II Cell Sorter (488–633-nm laser, 100-µm nozzle at 20 psi; BD); 1 ng of RNA per sample was used for complementary DNA preparation, according to the manufacturer instructions (PrimeScript RT, Takara Bio, Kusatsu, Japan), and a preamplification (TaqMan PreAmp Master Mix, Applied Biosystems, Foster City, CA) was performed. Real-time quantitative polymerase chain reaction (SDS 7900 HT instrument, Applied Biosystems) using the SYBR Green master mix (Applied Biosystems) was used to assess the expression level of the relevant genes. Nine RNA libraries were prepared using Nextera XT DNA (Illumina, San Diego, CA) kit corresponding to the triplicates of the three experimental conditions (*Scram* shRNA, *Dgcr2* shRNA, Rescue). Libraries were multiplexed and sequenced on an Illumina HiSeq2000 with a setup that generates pair-end reads of 2x50 bp at an average depth of 35 million read-pairs per library.

Phosphorylation Assay: Flow Cytometry

E17.5 electroporated cortices previously electroporated at E14.5 with *Scram* shRNA + green fluorescent protein (GFP) or *Dgcr2* shRNA + tdTomato were dissected and dissociated as described previously. tdTomato and GFP samples were then mixed and split into two tubes where they were washed in 10 mL of prewarmed L-15 (2% D-Glucose, Sigma-Aldrich) and centrifuged. Cells were resuspended in 1 mL of Reelin-enriched or control Dulbecco's modified Eagle medium and incubated for 30 minutes at 37°C; 500 µL of 16% prewarmed paraformaldehyde was added to the cells for a 15-minute incubation at 37°C. Cells were centrifuged and resuspended in 70% ice-cold methanol for a 30-minute incubation on ice, before being centrifuged and resuspended in 100 µL of PBS. A total of 20 µL of RAC-alpha serine/threonine-protein kinase (AKT) and anti-extracellular signal-related kinase 1/2 (ERK1/2) antibodies were added to the cells, and the mixture was incubated for 1 hour at room temperature on a rotating wheel. Cells were washed in PBS and filtrated through a cell strainer (70 µm, BD). Cells were analyzed by flow cytometry on a CyAn ADP analyzer (Beckman Coulter, Brea, CA).

Time-Lapse Imaging

E18.5 acute brain slices were prepared as described previously ([22](#)) from embryos electroporated at E14.5. Migrating

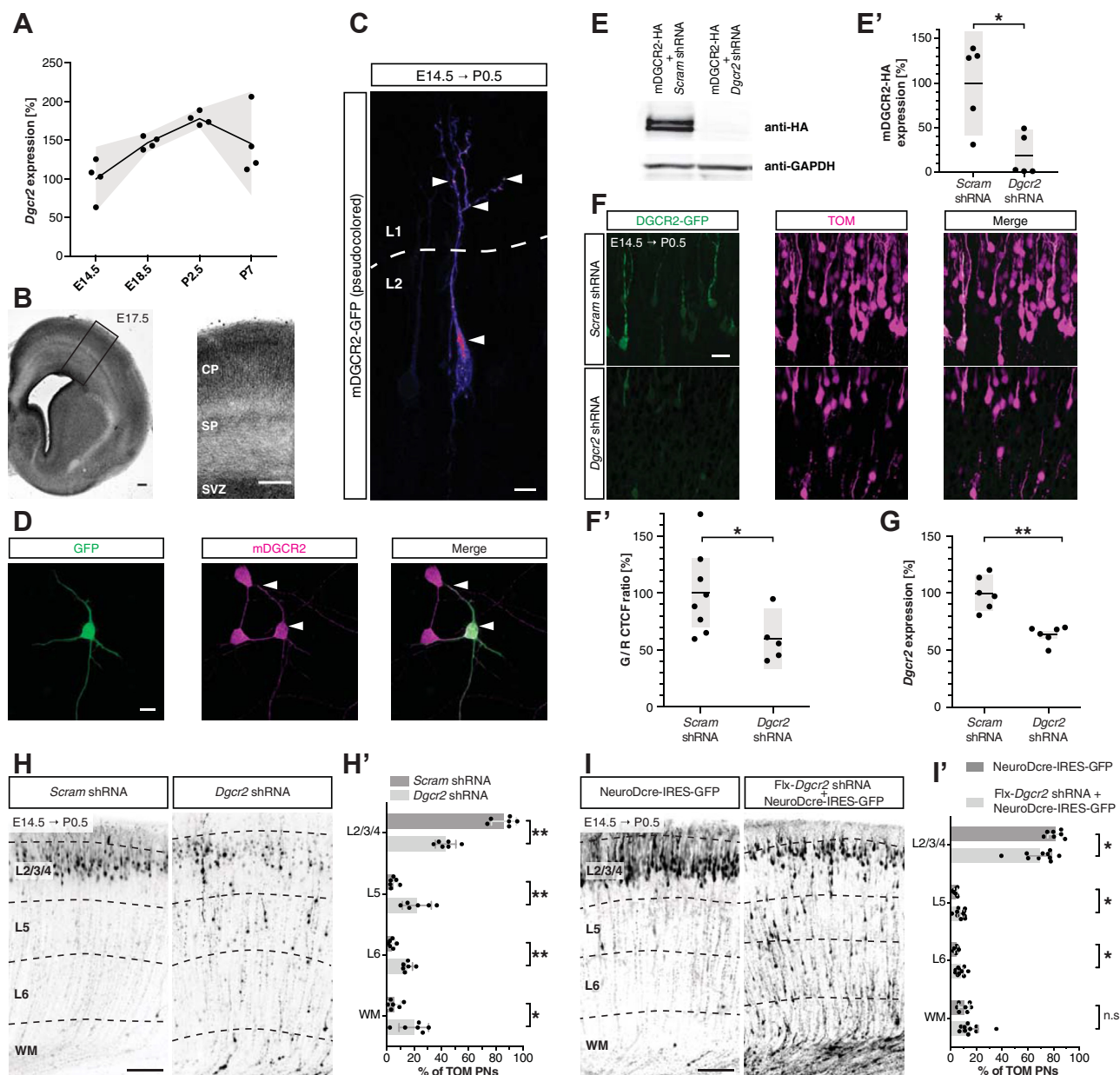


Figure 1. *Dgcr2* knockdown (KD) impairs the laminar positioning of cortical projection neurons (PNs). **(A)** Reverse transcriptase quantitative polymerase chain reaction indicates that *Dgcr2* is expressed during corticogenesis. $n = 4$ brains per time point from ≥ 3 separate litters. Error bar indicates 95% confidence interval (CI). **(B)** In situ hybridization reveals strong labeling of *Dgcr2* in the cortical plate (CP), subplate (SP), and subventricular zone (SVZ) of E17.5 brains. Scale bar = 100 μ m. **(C)** In utero electroporation at E14.5 of a green fluorescent protein (GFP)-tagged *Dgcr2* construct in PNs. Arrowheads indicate a puncta-shaped expression of mouse DiGeorge syndrome critical region 2 (mDGCR2) (pseudocolored) localized in the cytoplasm and apical processes. Scale bar = 10 μ m. **(D)** Immunohistochemistry on cultured cortical neurons demonstrates endogenous protein expression of mDGCR2 in PNs (arrowheads indicate mDGCR2 puncta). Scale bar = 10 μ m. **(E/E')** Western blot (anti-hemagglutinin [HA]) of overexpressed mDGCR2-HA in human embryonic kidney (HEK) 293T cells reveals a significant reduction of the mDGCR2-HA protein concentration, when cotransfected with a short hairpin RNA (shRNA) targeted against *Dgcr2*. $n = 5$; $p < .05$, Mann-Whitney. **(F/F')** The corrected total cell fluorescence (CTCF) ratio between mDGCR2-GFP and tdTomato (TOM) is reduced in vivo by coelectroporation of an shRNA targeted against *Dgcr2*. $n = 5$ –8 brains per condition from ≥ 3 separate litters. Error bars indicate 95% CI. * $p < .05$, Mann-Whitney. Scale bar = 20 μ m. **(G)** Reverse transcriptase quantitative polymerase chain reaction on RNA extracts of TOM PNs isolated by fluorescence-activated cell sorting at E17.5 after in utero electroporation at E14.5 shows that the total amount of *Dgcr2* is reduced by *Dgcr2* shRNA. $n = 6$ –7 litters per condition. Error bars indicate 95% CI. ** $p < .01$, Mann-Whitney. **(H/H')** shRNA-mediated *Dgcr2* KD by in utero electroporation at E14.5 dramatically impairs the laminar positioning of PNs in the somatosensory cortex at P0.5. $n = 6$ brains per condition from ≥ 3 separate litters. Error bars indicate 95% CI. * $p < .05$, ** $p < .01$, Mann-Whitney. Scale bar = 100 μ m. **(I/I')** Conditional expression of *Dgcr2* shRNA in postmitotic PNs by coelectroporation of neurogenic differentiation-cre-internal ribosome entry site-GFP (NeuroDcre-IRES-GFP) and a lox-flanked (lox) *Dgcr2* shRNA mimics the migratory deficit observed with constitutive *Dgcr2* KD at P0.5. PNs were visualized by immunohistochemistry against GFP. $n = 7$ –10 brains per condition from ≥ 3 separate litters. Error bars indicate 95% CI. * $p < .05$, Mann-Whitney. GAPDH, glyceraldehyde 3-phosphate dehydrogenase. n.s., nonsignificant; WM, white matter.

PNs located in the lower part of the cortical plate (CP) at the beginning of the recording were tracked during 10 hours.

Analysis and Statistics

All described image analyses were done using Fiji (40), using the following plugins: cell counter, MtrackJ (41). The corrected total cell fluorescence was quantified on Fiji as described in Burgess *et al.* (42). Except for the RNA sequencing data, all statistical analyses were performed on GraphPad Prism software, version 7 (GraphPad Software, La Jolla, CA). The sample sizes and relevant statistical tests are specified for each result in the Results. For the detailed methods of the RNA sequencing analysis, see the [Supplemental Methods](#).

RESULTS

Dgcr2 was found to be expressed during the late embryonic phase of corticogenesis (Figure 1A). In situ hybridization revealed that *Dgcr2* is expressed in the CP, subplate, and subventricular zone at E17.5, a period during which superficial PNs migrate toward the CP (Figure 1B). We next used in utero electroporation targeted to the E14.5 lateral ventricular zone to specifically manipulate the expression of *Dgcr2* in PN progenitors migrating toward superficial layers of the somatosensory cortex. Overexpression of a GFP-tagged *Dgcr2* construct (mouse *DGCR2*-GFP [m*DGCR2*-GFP]) in PNs showed a puncta-shaped expression in the apical processes of PNs (Figure 1C). The endogenous protein expression of *Dgcr2* was confirmed by immunohistochemistry on neuronal cultures of E14.5 electroporated PNs (Figure 1D). An shRNA targeted against *Dgcr2* significantly reduced the protein expression level of a HA-tagged *Dgcr2* (m*DGCR2*-HA) overexpressed in vitro in HEK 293T cells (Figure 1E). Furthermore, shRNA efficiency targeting *Dgcr2* was confirmed in vivo by measuring the fluorescence intensity of m*DGCR2*-GFP overexpressed in PNs (Figure 1F) and by performing reverse transcriptase quantitative polymerase chain reaction on in utero electroporated PNs isolated by fluorescence-activated cell sorting at E17.5 (Figure 1G). Overall, the shRNA-mediated KD of *Dgcr2* reduced by around 40% its messenger RNA expression.

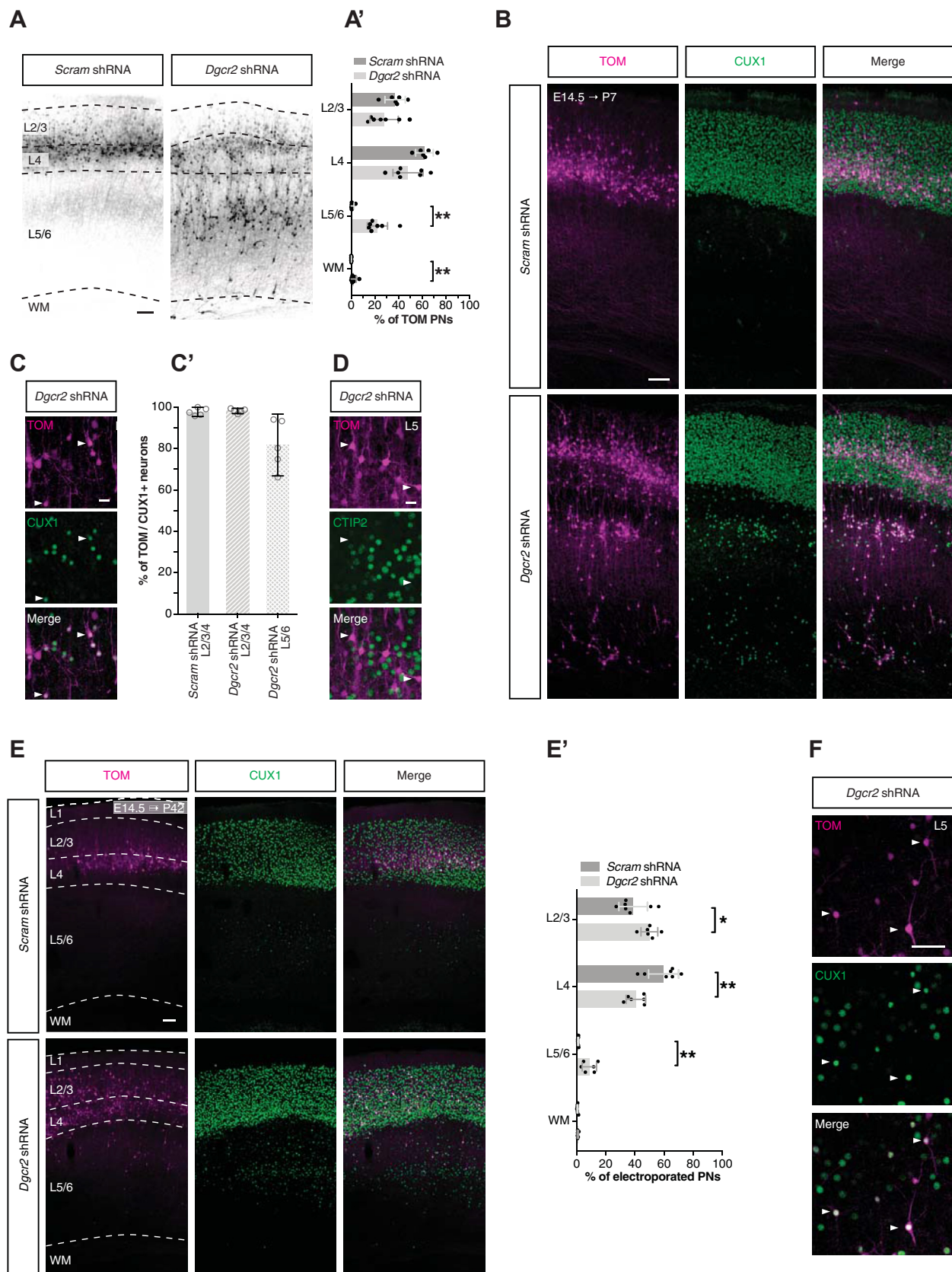
Dgcr2 KD induced a significant alteration in the laminar positioning of PNs at P0.5, compared with a control shRNA (*Scram* shRNA). *Dgcr2*-KD PNs were significantly misplaced in the layers 5 and 6, and in the white matter (Figure 1H). This mispositioning phenotype was replicated using a second shRNA against *Dgcr2* (Supplemental Figure S1C). Furthermore, conditional expression of *Dgcr2* shRNA in postmitotic PNs induced a similar type of mispositioning phenotype (Figure 1I). *Dgcr2* KD did not alter the laminar positioning of PNs at the earlier embryonic ages of E17.5 and E18.5 (Supplemental Figure S1A, B). In addition, *Dgcr2* KD did not appear to alter the proliferation rate of PN progenitors (Supplemental Figure S2A) or their early differentiation process (Supplemental Figure S2B–E). At E15.5, the percentage of PNs that expressed the intermediate progenitor marker T-box brain protein 2 (43) in the subventricular zone was similar in the *Scram* and *Dgcr2* shRNA conditions (Supplemental Figure S2B). At E17.5, *Dgcr2*-KD PNs expressed the upper-layer markers special AT-rich sequence binding protein 2 and

BRN2 (44,45), in the same proportion as control PNs (Supplemental Figure S2C, D). In addition, the percentage of electroporated PNs that expressed the deep-layer marker T-box brain protein 1 (46) remained similarly low between *Dgcr2* shRNA- and *Scram* shRNA-electroporated PNs (Supplemental Figure S2E). Overall, *Dgcr2* KD had a major impact on late PN migration but did not affect the proliferation or the early differentiation process of PN progenitors.

Dgcr2 KD induced a persistent laminar positioning deficit (Figure 2). At P7, misplaced *Dgcr2*-KD PNs were observed in deep cortical layers (Figure 2A, B) and preserved their upper-layer cut like homeobox protein cut-like 1 (CUX1) molecular identity, whereas they did not express the deep-layer marker COUP-TF-interacting protein 2 (47) (Figure 2B–D). In addition, no difference was observed in the percentage of electroporated PNs that express CUX1 in the upper layers between the *Dgcr2*-KD and control conditions (Figure 2C). Finally, at P42, misplaced *Dgcr2*-KD PNs were still observed in deep cortical layers, as opposed to control PNs (Figure 2E), and maintained expression of the upper-layer marker CUX1 (Figure 2F).

Given that the mispositioning phenotype appeared between E18.5 and P0.5, we next assessed dynamically the impact of *Dgcr2* KD on neuronal migration using confocal time-lapse imaging of PNs in E18.5 cortical slices. At this developmental time point, E14.5-born PNs are undergoing radial-guided locomotion from the upper part of the intermediate zone toward the CP (22). Quantification revealed that *Dgcr2* KD dramatically decreased the migratory speed of PNs (Figure 3A–D, Supplemental Movies S1, 2). At P0.5, *Dgcr2* KD significantly shifted the distribution of PNs to the lower CP, suggesting that *Dgcr2* KD affects the process of terminal translocation (16,20), the final step of PN migration (Figure 4A, B). To further assess the impact of *Dgcr2* KD on terminal translocation, we performed sequential in utero electroporations at E14.5 to label a reference population of control PNs and at E15.5 to assess the effects of *Dgcr2* KD on later-born PNs, which translocate in more superficial layers compared with E14.5-born PNs (20). Strikingly, E15.5-born *Dgcr2*-KD PNs were intermingled with the E14.5-born control PNs, whereas E15.5-born *Scram*-shRNA PNs were distinctly laminated in superficial layers of the CP (Figure 4C). Overall, these results indicate that radial-guided locomotion and terminal translocation of PNs are regulated by *Dgcr2*.

We next aimed to determine the molecular mechanisms through which *Dgcr2* regulates PN migration. To do this we used an in vivo electroporation rescue strategy with mouse shRNA-resistant human *DGCR2* constructs. First, we found that the *Dgcr2*-KD mispositioning phenotype was rescued by the wild-type (WT) human (h) *DGCR2* construct (Rescue WT) (Figure 5A), thus providing molecular specificity for the *Dgcr2*-KD PN migratory phenotype. Additionally, overexpression of *DGCR2* per se did not affect the laminar positioning of electroporated PNs (Supplemental Figure S1D). We then took advantage of this rescue strategy to investigate the migratory function of different subdomains of *DGCR2* (Figure 5A). Mutated constructs with deletions of the cytoplasmic C-terminal domain of *DGCR2* (C'-Del) or the extracellular low-density lipoprotein receptor class A (LDL-A) domain (LDL-Del) (Supplemental Figure S3A) did not rescue the *Dgcr2*-KD PN



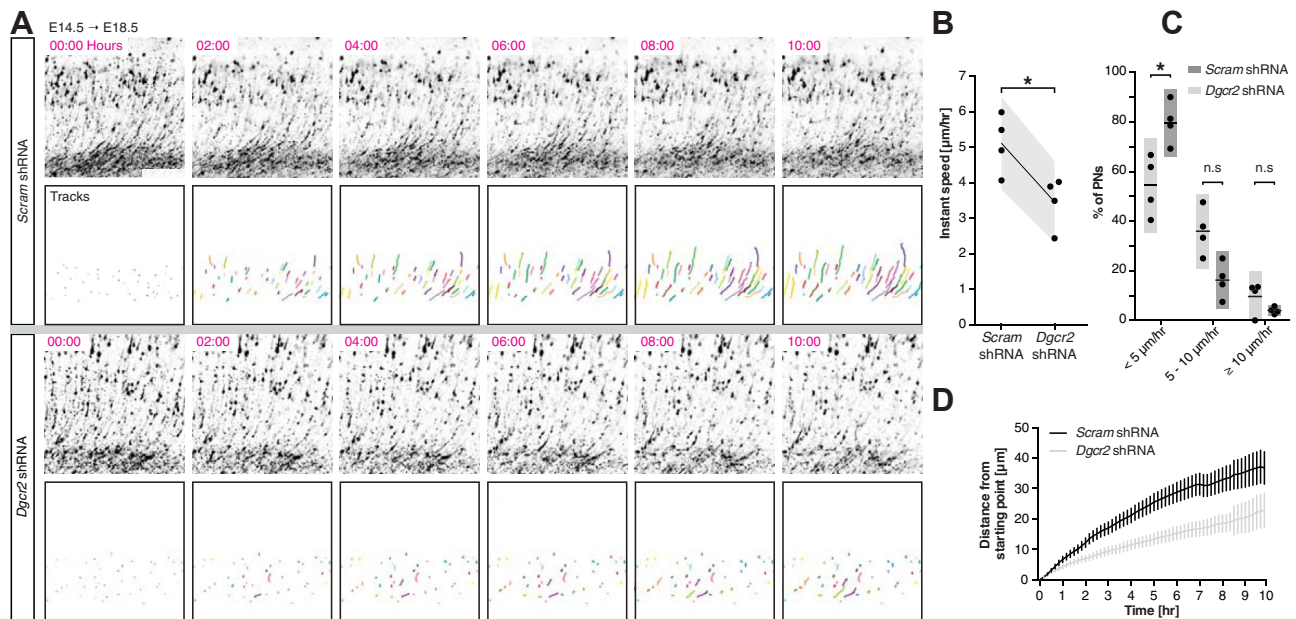


Figure 3. *Dgcr2* knockdown reduces the migratory speed of projection neurons. (A–D) Confocal time-lapse imaging at E18.5 of migrating projection neurons electroporated at E14.5 with tdTomato (TOM) and *Dgcr2* short hairpin RNA (shRNA) or *Scram* shRNA shows that *Dgcr2* knockdown reduces (B) the average instant speed, (C) the percentage of cells moving at high speed, and (D) the distance from the starting point as compared with control condition. $n = 4$ brains per condition from ≥ 3 separate litters, 183–196 tracked cells per condition. Color coding indicates illustrative tracked cells. Error bars indicate 95% confidence interval. $*p < .05$, Mann-Whitney. Scale bar = 100 μm . n.s., nonsignificant.

migratory phenotype (Figure 5A), thus indicating that these molecular subdomains are required for human DGCR2 (hDGCR2) function in PN migration. In a more translational perspective, we also assessed the functional consequence of the SZ-risk mutation P429R (3). Strikingly hDGCR2-P429R failed to rescue the *Dgcr2*-KD migratory phenotype, providing evidence that this SZ-risk mutation is pathogenic for PN migration. Interestingly, western blots of transfected HEK 293T cells with hDGCR2-WT-HA or hDGCR2-P429R-HA revealed a significantly lower protein expression of hDGCR2-P429R-HA in contrast to WT (Figure 5B). Furthermore, overexpression of hDGCR2-P429R-HA in mouse primary cortical neurons indicated that the protein expression of hDGCR2-P429R-HA was reduced (Supplemental Figure S3B), whereas messenger RNA expression of *Dgcr2* was not modified compared with the control condition (Supplemental Figure S3C). Proteasome inhibition using MG 132, an inhibitor of the proteasome activity (48), led to a significant increase (about 60%) of hDGCR2-

P429R-HA (Supplemental Figure S3D), suggesting that the SZ-risk mutation affects protein expression of DGCR2 through increased proteasome degradation.

We next investigated the molecular mechanisms mediating the effects of DGCR2 on PN migration. Interestingly the extracellular LDL-A domain of *DGCR2* carries structural similarities with LDL-A domains found in the two canonical Reelin receptors very low-density lipoprotein receptor and apolipoprotein E receptor 2 (49). As reported (50), the canonical very low-density lipoprotein receptor coimmunoprecipitated with both full-length Reelin and the Reelin central fragment and was used here as a positive control condition (Figure 6A). Coimmunoprecipitation of Reelin with DGCR2 revealed that full length Reelin and the monomeric form of the central fragment of Reelin coimmunoprecipitated with mDGCR2 and hDGCR2, whereas no Reelin was detected in the negative control condition (GFP) (Figure 6A). Molecular proximity between DGCR2 and Reelin was further assessed in vivo using a PLA on E18.5

Figure 2. *Dgcr2* knockdown persistently affects the positioning of projection neurons (PNs) until P42 but not their laminar molecular identity. (A/A') *Dgcr2* knockdown impairs the laminar positioning of E14.5-born PNs at P7. $n = 6$ –7 brains per condition from ≥ 3 separate litters. Error bars indicate 95% confidence interval (CI). $**p < .01$, Mann-Whitney. Scale bars = 100 μm . (B) Immunohistochemistry of the upper-layer neuronal marker cut like homeobox 1 (CUX1) at P7, after in utero electroporation of *Dgcr2* short hairpin RNA (shRNA) at E14.5, compared with *Scram* shRNA. Scale bars = 100 μm . (C/C') High magnification captures of mispositioned PNs in layer 5 in the *Dgcr2* shRNA condition shows that at P7 the vast majority of misplaced PNs express the upper-layer marker CUX1. Arrowheads show colocalized PNs. $n = 5$ brains per condition from ≥ 3 separate litters. Error bars indicate 95% CI. Scale bars = 20 μm . (D) Mispositioned PNs do not express the deep-layer marker COUP-TF-interacting protein 2 (CTIP2). Arrowheads indicate electroporated PNs. Scale bars = 20 μm . (E/E') *Dgcr2* knockdown impairs the laminar positioning of E14.5-born PNs at P42. $n = 6$ –7 brains per condition from ≥ 2 separate litters. Error bars indicate 95% CI. $**p < .01$, Mann-Whitney. Scale bars = 100 μm . (F) High-magnification image of mispositioned PNs in layer 5 in the *Dgcr2* shRNA condition shows that at P42, misplaced PNs maintain expression of the upper-layer marker CUX1. Arrowheads show colocalized PNs. Scale bars = 50 μm . TOM, tdTomato; WM, white matter.

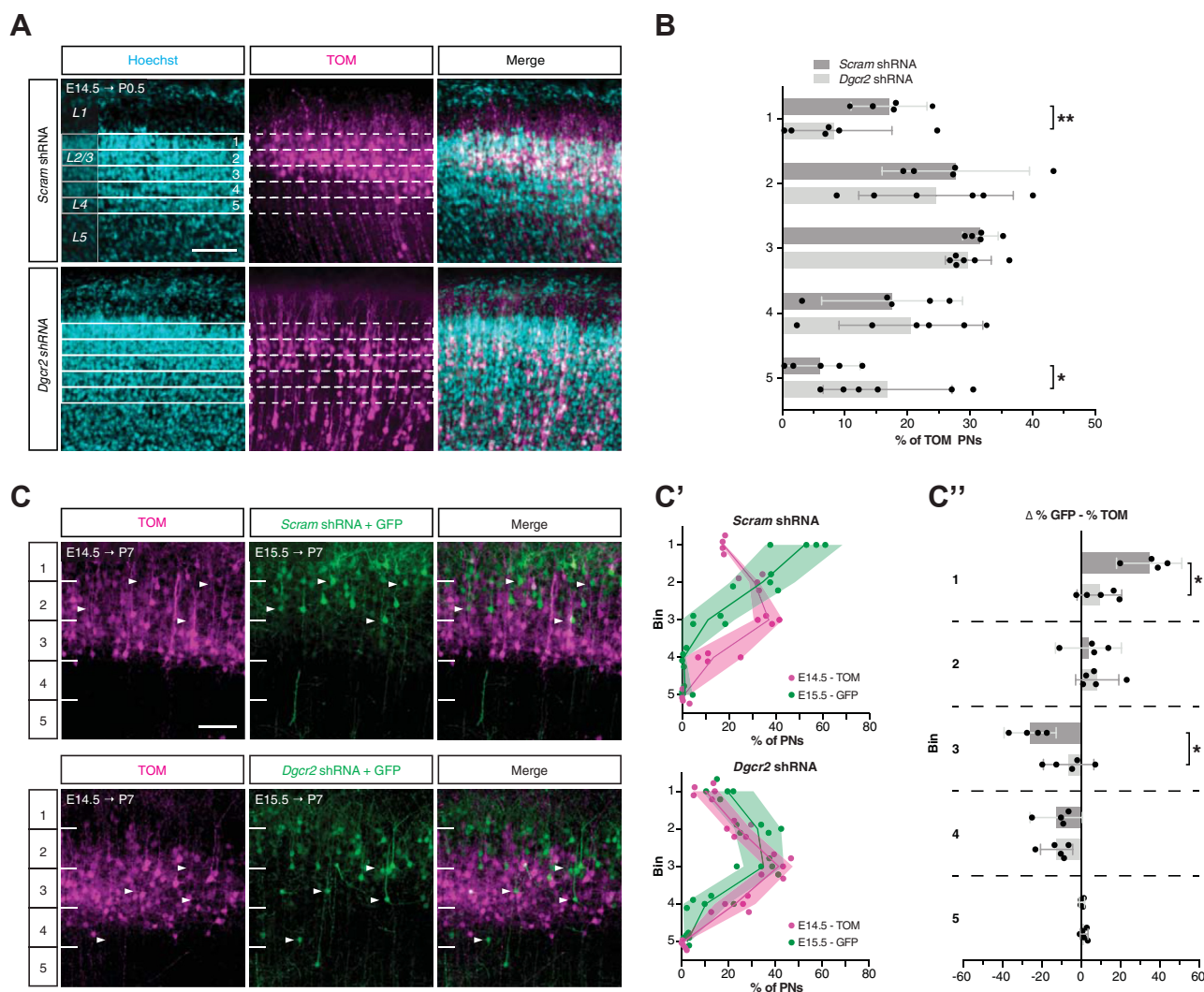


Figure 4. *Dgcr2* knockdown (KD) impairs the terminal translocation of upper-layer projection neurons (PNs). **(A, B)** *Dgcr2* KD reduces the fraction of E14.5 electroporated PN reaching the outermost region of the cortical plate (CP) at P0.5 (Bin 1) as compared with *Scram* short hairpin RNA (shRNA). Same dataset as in Figure 1. $n = 5-6$ brains per condition from ≥ 3 separate litters. $*p < .05$, $**p < .01$, Mann-Whitney. Error bars indicate 95% confidence interval. **(C)** Sequential in utero electroporations of PN at E14.5 with tdTomato (TOM) and at E15.5 with green fluorescent protein (GFP) + *Scram* or *Dgcr2* shRNA demonstrates that at P7 the inside-out laminar assembly of PN is disrupted by *Dgcr2* KD. Arrowheads indicate E15.5 born PN. $n = 4-5$ brains per condition from ≥ 3 separate litters. $*p < .05$, Mann-Whitney. Error bars indicate 95% confidence interval. Scale bar = 100 μm .

cortical slices. We observed that the overall density of PLA puncta was highest in the marginal zone (MZ) as compared with other cortical compartments (Figure 6C), in line with observations that Reelin is mainly secreted by Cajal-Retzius cells in the MZ (51). Strikingly, we found that *Dgcr2* KD decreased PLA puncta density as compared with *Scram* shRNA condition in the MZ (Figure 6C), whereas the total protein expression of Reelin was unaffected by *Dgcr2* KD (Supplemental Figure S3E). These results provide novel evidence that DGCR2 and Reelin form a protein complex in vitro and in vivo.

To determine whether *Dgcr2* regulates Reelin signaling, we investigated the impact of *Dgcr2* KD on DAB1, a Reelin-dependent phosphorylation target controlling PN migration (52). DAB1 phosphorylation was assessed in primary neuronal

cultures after stimulation with Reelin-enriched medium or mock medium using DAB1 immunoprecipitation followed by p-Y Western blot. For this experiment, *Dgcr2* KD was obtained by lentiviral infection of *Dgcr2* shRNA (Supplemental Figure S3F) in cultured neurons. In each of the five replicates of this experiment, Reelin-dependent phosphorylated DAB1 levels were decreased by *Dgcr2* KD (Figure 7A). We next performed a flow cytometry-based phosphorylation assay to assess AKT and ERK1/2, two additional Reelin-dependent phosphorylation targets (39,53). Following in utero electroporation at E14.5, control or *Dgcr2*-KD PN were exposed at E17.5 to a Reelin-enriched medium or a control medium and the phosphorylation status of AKT (pS473) and ERK1/2 (pT202/pY204) was assessed using flow cytometry (Figure 7B).

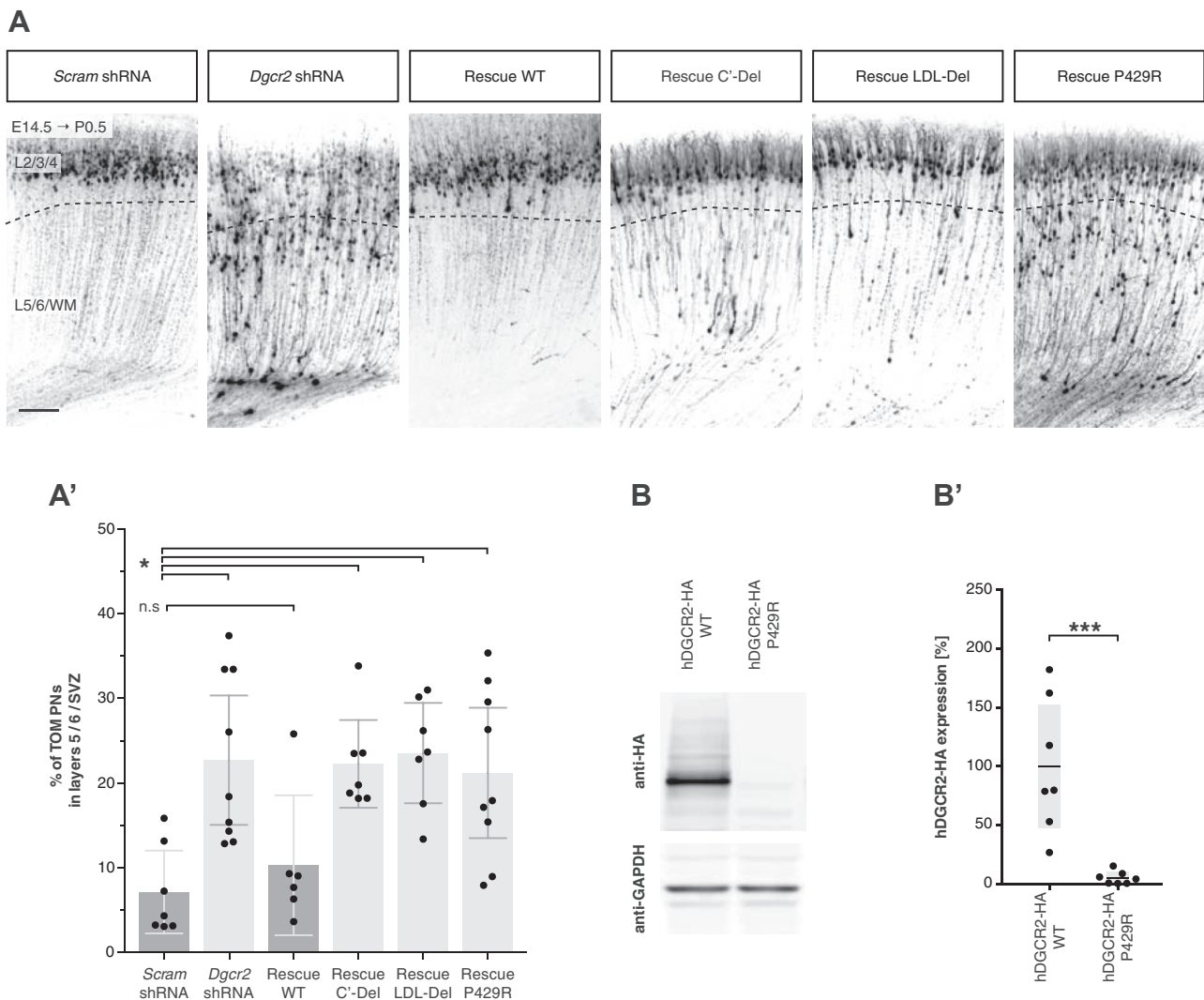


Figure 5. The human DiGeorge syndrome critical region 2 (*hDGCR2*) schizophrenia (SZ)-risk mutation does not rescue the *Dgcr2* knockdown (KD) mispositioning phenotype. **(A/A')** Overexpression of *DGCR2* is able to rescue the positioning of projection neurons (PNs) in upper layers at P0.5, when coelectroporated with *Dgcr2* short hairpin RNA (shRNA) at E14.5 (Rescue wild-type [WT]). Deletions of the C' terminal domain (Rescue C'-Del) or the low-density lipoprotein receptor class A (LDL-A) domain (Rescue LDL-A-Del) of *DGCR2*, as well as the missense SZ-risk mutation P429R (Rescue P429R), do not rescue the mispositioning phenotype. $n = 6-9$ brains per condition from ≥ 3 separate litters. $*p < .05$, Kruskal-Wallis + Dunn's. Error bar indicates 95% confidence interval. Scale bar = 100 μm . **(B/B')** Western blot (anti-hemagglutinin [HA]) of protein lysates from human embryonic kidney (HEK) 293T cells overexpressing hDGCR2-WT-HA or hDGCR2-P429R-HA shows that P429R is associated with a reduced total amount of hDGCR2. $n = 7$. Error bars indicate 95% confidence interval. $***p < .001$, Mann-Whitney. n.s., nonsignificant; TOM, tdTomato; WM, white matter.

Reelin-induced phosphorylation of phosphorylated AKT (Figure 7C) and phosphorylated ERK1/2 (Figure 7D) were decreased in all replicates in the *Dgcr2*-KD condition as compared with the control condition. Overall, these experiments indicate that *DGCR2* is a novel member of the Reelin complex, regulating Reelin-dependent signaling targets in PNs.

To identify genetic pathways dysregulated by *Dgcr2* KD in migrating PNs, RNA sequencing was performed on PNs isolated at E17.5 using fluorescence-activated cell sorting, after E14.5 in utero electroporation. *Dgcr2* was among the top downregulated genes in the *Dgcr2*-KD condition, validating the

experimental approach. Expression levels of 98 genes were found to be specifically altered in the *Dgcr2*-KD condition in contrast to the control and rescue conditions (Figure 8A, Supplemental Table S1). Interestingly, Reelin target genes (32) were significantly enriched in our dataset ($p = 1.60 \times 10^{-4}$, hypergeometric testing; Figure 8A, D). Some of the *Dgcr2*-KD dysregulated genes were associated with mental disorders (54) (Figure 8B), and among them some were positively associated with psychiatric disorders such as SZ and bipolar spectrum disorders (55) (Figure 8C). Finally, Gene Set Enrichment Analysis (56) revealed that dysregulated *Dgcr2*-KD target genes were enriched in biological pathways linked to developmental

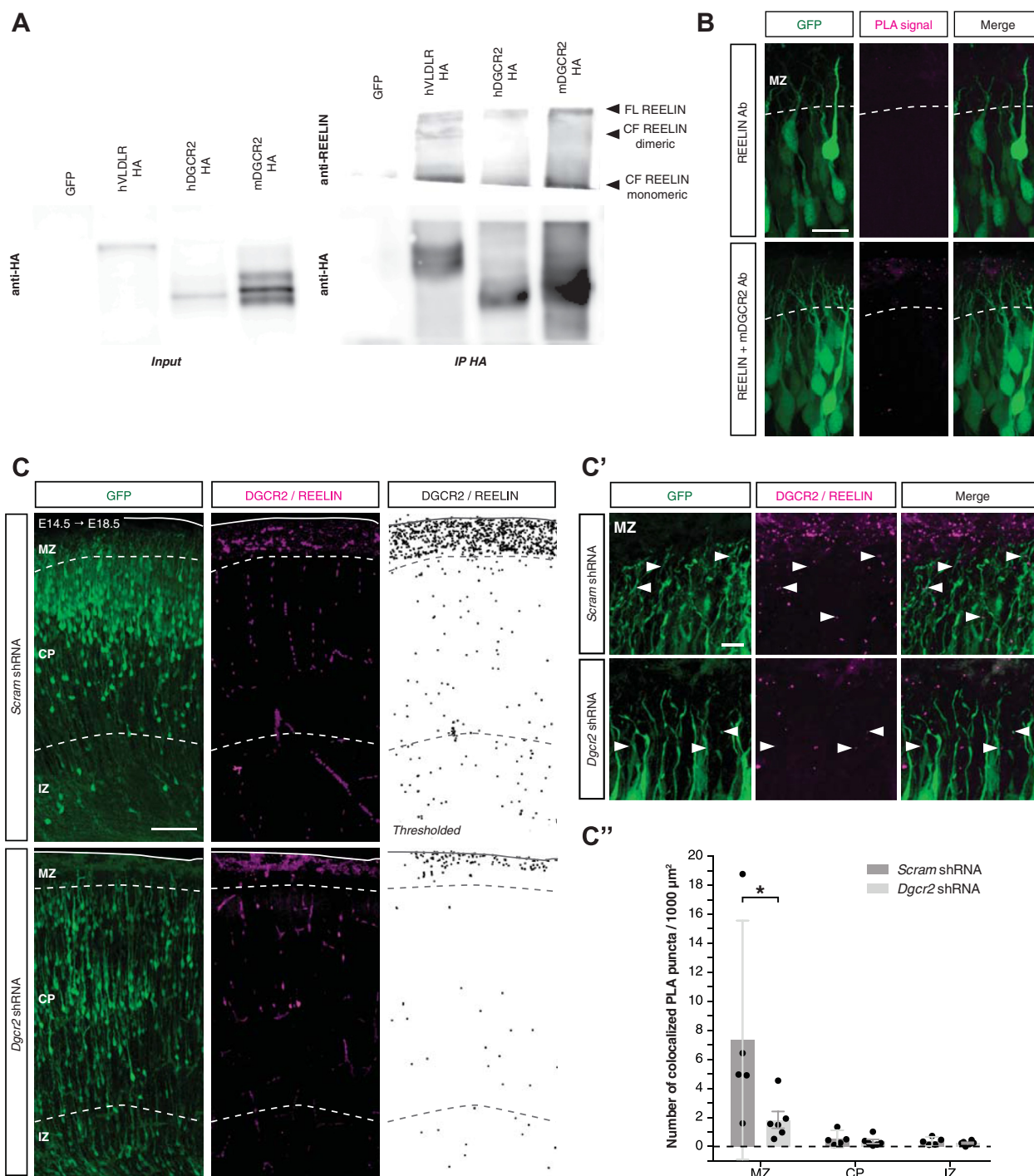


Figure 6. DiGeorge syndrome critical region 2 (*DGCR2*) is part of the Reelin complex. **(A)** Protein lysates from human embryonic kidney (HEK) 293T cells overexpressing green fluorescent protein (GFP), human very low-density lipoprotein receptor hemagglutinin (hVLDLR-HA), mouse *DGCR2* HA (m*DGCR2*-HA), or human *DGCR2* HA (h*DGCR2*-HA) were exposed to a Reelin-enriched medium and immunoprecipitated (IP) with HA. Reelin and HA were revealed by Western blot. Full-length (molecular weight > 440 kDa) Reelin and its central fragment (molecular weight = 180 kDa) coprecipitated with hVLDLR, m*DGCR2*, and h*DGCR2* whereas the 320-kDa dimeric form of the Reelin central fragment only coprecipitated with hVLDLR. **(B)** Illustrative images of proximity ligation assay (PLA) signal on E18.5 brain slices where PLA was performed without m*DGCR2* antibody (Ab) (Reelin Ab) as a negative control, compared with the test condition in which both Reelin and m*DGCR2* Ab are used. Scale bar = 20 μm . **(C)** PLA on E18.5 brains electroporated at E14.5 with GFP and *Scram* or *Dgcr2* short hairpin RNA (shRNA) reveals a greater density of PLA puncta in the marginal zone (MZ) as compared with other layers of the cortex. Thresholded image

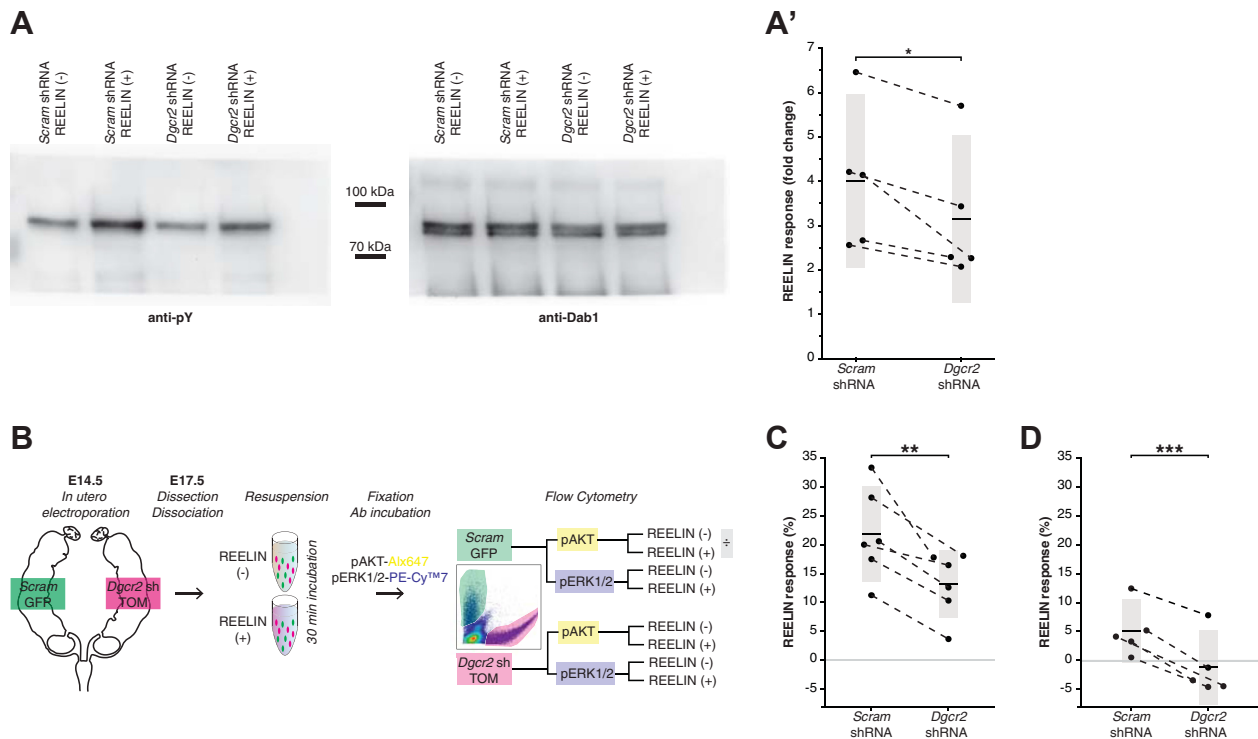


Figure 7. *Dgcr2* knockdown (KD) affects Reelin-dependent phosphorylation of disabled homolog 1 (DAB1), RAC-alpha serine/threonine-protein kinase (AKT), and extracellular signal-related kinase 1/2 (ERK1/2). **(A)** Western blot for pY and DAB1 of DAB1 immunoprecipitates shows an increased DAB1 phosphorylation in cortical neurons after exposure to a Reelin-enriched medium in the control condition, but the difference is fainter in the *Dgcr2*-KD condition. **(A')** Quantification of the Reelin-induced phosphorylation of DAB1 in *Dgcr2*-KD cortical neurons compared with control cortical neurons, calculated as pDAB1^{REELIN(+)} / pDAB1^{REELIN(-)} normalized on corresponding total DAB1, shows a decreased Reelin response in *Dgcr2*-KD cortical neurons. **p* < .05, paired *t* test. **(B)** Schematics of the flow cytometry phosphorylation assay for phosphorylated AKT (pAKT) and phosphorylated ERK1/2 (pERK1/2). Cortical projection neurons were obtained at E17.5 from somatosensory cortices of brains from the same litter, electroporated at E14.5 with green fluorescent protein (GFP) + *Scram* short hairpin RNA (shRNA) or tdTomato (TOM) + *Dgcr2* shRNA. Projection neurons were dissociated, exposed to Reelin-enriched or control medium for 30 minutes, fixed, and incubated with pAKT-Alexa 647 and pERK1/2-PE-Cy7 for flow cytometry analysis. The Reelin response was assessed by the ratio of pAKT or pERK1/2 mean fluorescence between Reelin (+) and Reelin (-) conditions, expressed in percent of increase in Reelin (+). **(C)** *Dgcr2* KD reduces the Reelin-induced phosphorylation of pAKT. *n* = 5 separate litters. ***p* < .01, paired *t* test. **(D)** *Dgcr2* KD reduces the Reelin-induced phosphorylation of pERK, although the average Reelin response in the control condition is only around 5%. *n* = 5 separate litters. ****p* < .001, paired *t* test. pDAB1, phosphorylated disabled homolog 1.

processes and neuronal specific functions (Figure 8D, Supplemental Figure S3G, Supplemental Table S1).

DISCUSSION

Overall, this study reveals that the SZ-risk gene *Dgcr2* regulates the migration and laminar positioning of upper-layer cortical PNs. Using mutated *DGCR2* constructs, we find that the extracellular LDL-A and the cytoplasmic C'-terminal domains of *DGCR2* are important molecular domains required for the migratory function of the protein. In addition, our results indicate that the missense SZ-risk mutation P429R (3) has a pathogenic effect on PN migration by leading to a state of *DGCR2* protein insufficiency. Mechanistically, we show that

DGCR2 is part of the Reelin complex and that *Dgcr2* KD regulates Reelin-dependent signaling pathways. Finally, using RNA sequencing we identify in migrating PNs a set of *Dgcr2*-dependent genes, which are significantly enriched in previously reported Reelin-dependent genetic targets (32).

The migration of upper-layer PNs is controlled by a variety of factors (23,24), including Reelin (57). Using time-lapse imaging and sequential electroporation in vivo we find that *Dgcr2* regulates both radial-guided locomotion as well as terminal translocation. Strikingly, the *Dgcr2*-KD PN mispositioning phenotype observed in the CP phenocopies the migratory phenotype described following in utero KD of DAB1, a key Reelin effector (20), thus suggesting that *DGCR2* regulates PN migration through the Reelin signaling pathway.

represents *DGCR2*/Reelin PLA puncta as isolated particle <5 μm^2 and offset by 5 pixels. PNs were visualized by immunohistochemistry against GFP. **(C'/C'')** In the MZ the number of PLA puncta that colocalize with the electroporated PNs is reduced by *Dgcr2* KD as compared with the control condition. Arrowheads depict *DGCR2*/Reelin PLA puncta colocalizing with GFP. *n* = 5–6 brains per condition from ≥ 3 separate litters. **p* < .05, Mann-Whitney. Error bars indicate 95% confidence interval. **(C)** Scale bar = 100 μm . **(C')** Scale bar = 10 μm . CP, cortical plate; IZ, intermediate zone.

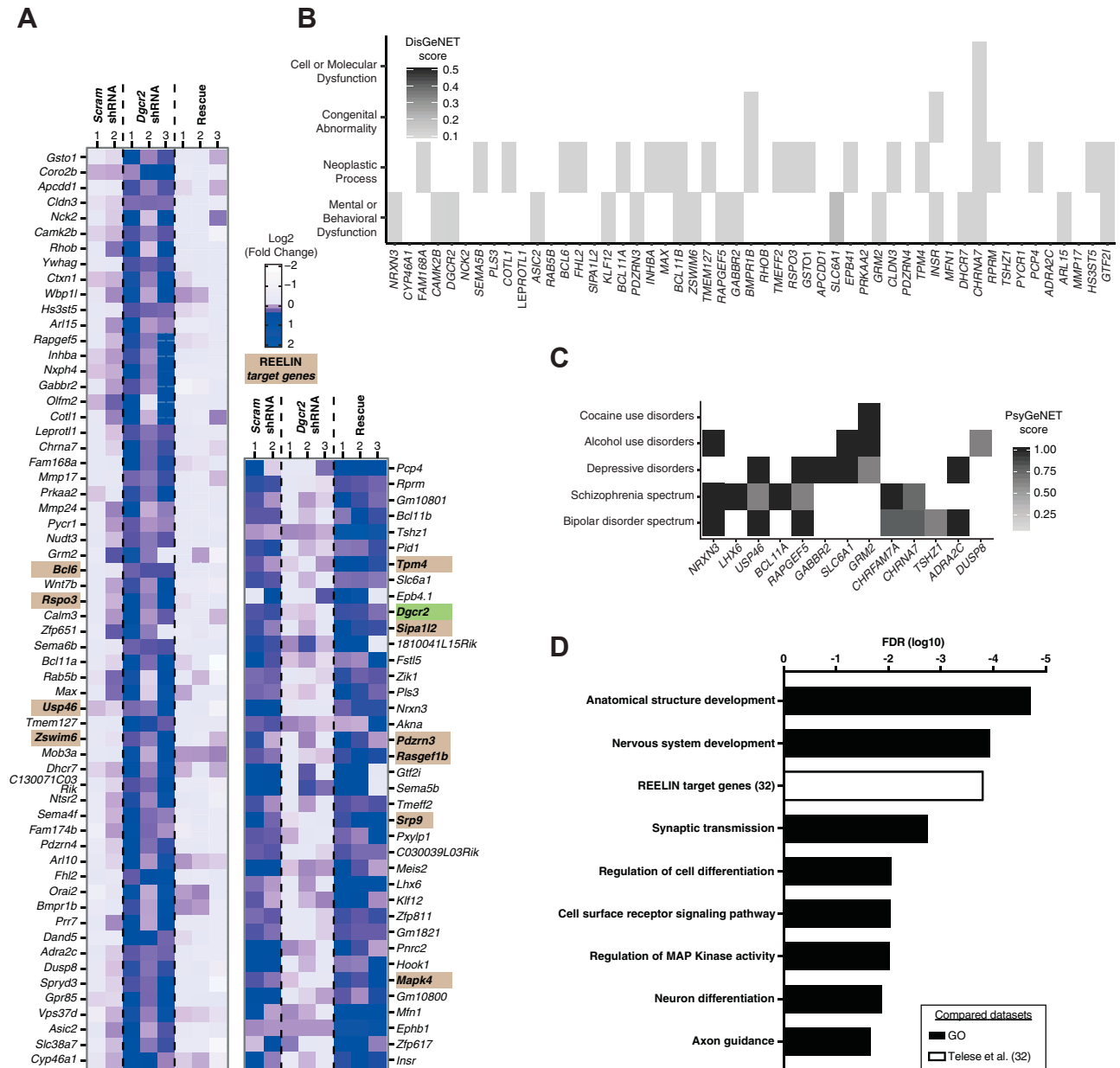


Figure 8. Transcriptional changes induced by *Dgcr2* knockdown (KD) in migrating projection neurons. **(A)** E14.5 electroporated projection neurons were isolated by fluorescence-activated cell sorting at E17.5 for RNA sequencing. Ninety-eight genes were identified as differentially expressed in the *Dgcr2*-KD condition compared with *Scram* short hairpin RNA (shRNA) or *Dgcr2* shRNA + human DiGeorge syndrome critical region 2 (hDGCR2) (Rescue). Sixty genes displayed an increased expression associated with *Dgcr2* KD and 38 displayed a decreased expression following *Dgcr2* KD. Highlighted in brown are 10 transcripts previously identified as Reelin target genes. *Dgcr2* is highlighted in green. **(B)** Selected disorders associated with *Dgcr2*-KD dysregulated genes, as identified by the DisGeNET database, encompass mental and behavioral disorders as well as congenital diseases. **(C)** PsyGeNET score highlights genes in our gene set that were positively associated with schizophrenia spectrum disorder and bipolar disorder spectrum. **(D)** Gene Set Enrichment Analysis revealed selected biological processes enriched in *Dgcr2*-KD differentially expressed genes. FDR, false discovery rate; GO, gene ontology; MAP, mitogen-activated protein.

Using coimmunoprecipitation and PLA, we reveal that DGCR2 is part of the Reelin complex. Mechanistically, we find that DGCR2 regulates Reelin-dependent phosphorylation of key intracellular effectors such as DAB1, AKT, and ERK1/2. Finally, recent work has revealed key transcriptional changes induced

by Reelin activation (32). Here, we identify a core set of genes whose expression is specifically dysregulated following *Dgcr2* KD. The mitogen-activated protein kinase pathway (58,59) was enriched in the pathway analysis of *Dgcr2*-dependent genes and in this context, *Mapk4* represents an interesting

link between *Dgcr2* and Reelin (32). In addition, a significant proportion of these genes were enriched for Reelin target genes previously identified (32). Overall, these results provide evidence that *DGCR2* is functionally linked to the Reelin pathway and suggests that the *Dgcr2*-KD migratory phenotype could be due to defective Reelin signaling in PNs.

Our experimental approach is based on shRNA-induced downregulation in PNs using targeted in utero electroporation. Given that this method has been reported to induce off-target effects (60), we performed rescue experiments using an *hDGCR2* construct that did not contain the mouse shRNA target sequence. The *hDGCR2* construct induced a full rescue of the shRNA-induced mispositioning phenotype, indicating the specificity of our *Dgcr2*-KD approach. In addition, this rescue approach was used to control for potential off-target gene expression changes and allowed the identification specific *Dgcr2*-KD target genes in our RNA sequencing dataset. Finally, our quantitative analysis did not detect major changes in proliferation and early neuronal differentiation following *Dgcr2* KD. However, a role for *Dgcr2* in these early developmental processes cannot be fully excluded because the efficiency of shRNA-induced KD may not be sufficiently high at these early developmental time points.

In a disease perspective, allelic variations of Reelin have been associated with SZ (61–64) and endophenotypes of SZ (65). However, single nucleotide polymorphisms and de novo mutations alone have a relatively modest effect size on the risk of developing SZ (66), compared with rare copy number variations such as 22q11.2DS (67). Among the genes encoded in this locus, *DGCR2* has been associated with SZ (3,11), although a lack of replication has been reported (68,69). More recently, an exome-sequencing study identified a potentially disruptive de novo mutation in *DGCR2* associated with idiopathic SZ (3). Here we found that this SZ-risk mutation (P429R) had a pathogenic effect on PN migration by decreasing *DGCR2* protein but not transcripts levels. Decreased levels of *DGCR2*-P429R could be linked to increased protein degradation by the proteasome, but other mechanisms such as defective subcellular targeting or a reduced translation rate due to codon usage bias might also be involved. Overall, our results indicate that the P429R mutation could phenocopy a state of haploinsufficiency as observed in 22q11.2DS, thus suggesting that the rare de novo *DGCR2* mutation detected in a single case of idiopathic SZ (3) shares a common pathogenic mechanism with the 22q11.2DS. In a more general perspective, although numerous genetic studies provide support for an association between rare de novo mutations and SZ (3,5,70–72), this study provides, to our knowledge, the first insight into the biological pathogenicity of one of these rare mutations in vivo. SZ is conceived as neurodevelopmental disorder due to multiple pathogenic hits operating at early embryonic stages and at the later adolescent period (73). Here we display evidence that a human rare SZ-risk *DGCR2* mutation affects early steps of cortical assembly, further indicating the relevance of early embryonic hits in SZ.

The long-term effects of *Dgcr2* KD on the connectivity and circuit function of PNs will have to be assessed in future studies. Altered embryonic migration of PNs due to genetic insults has been proposed to impair their integration into cortical circuits and lead to network dysfunction as well as

SZ-related behavioral deficits (74). For example, transient and selective KD of the SZ-risk gene *Disc1* in cortical PNs has been reported to affect neuronal migration (75), impair the electrophysiological properties of PNs as well as their response to dopamine pharmacological challenge, and induce SZ-relevant behavioral abnormalities, which could be reversed by clozapine (76). In addition, several studies using mouse models of 22q11.2DS have revealed developmental alterations including decreased progenitor proliferation of superficial layer PNs (77), delayed migration of hippocampal dentate progenitors (78), altered migration and distribution of parvalbumin-expressing cortical interneurons (78,79), and decreased density of superficial cortical PNs (77,80). Furthermore, at the functional level, mouse models of 22q11.2DS have revealed an imbalance between excitation and inhibition in local cortical circuits upon dopamine challenge (81), have disrupted axonal branching and synaptic transmission (82), have altered spine turnover and short-term plasticity (80), and have impaired activity patterns in neuronal ensembles (83) as well as theta- and gamma-band oscillation deficits (83,84). Whether *DGCR2* plays a role in these SZ-relevant functional phenotypes remains to be determined. Finally, alterations in the distribution or density of CUX1+ superficial PNs were observed in the *Dgcr2*-KD model and in a 22q11.2DS mouse model (77,80). Given that the recent use of laminar markers revealed patches of abnormal cortical microstructure in post-mortem tissue from children with autism spectrum disorders (85), similar types of investigations should be performed in human brains of 22q11.2DS patients.

In conclusion, we report that the SZ-risk gene *Dgcr2* regulates PN migration and acts through the Reelin pathway by modulating the phosphorylation of downstream effectors, and by regulating the expression of Reelin-dependent genes. Moreover, we found that a rare SZ-associated missense mutation in *DGCR2* has a pathogenic effect on PN migration, possibly by leading to a state of *DGCR2* haploinsufficiency as seen in 22q11.2Del carriers. This study adds support to the hypothesis that SZ risk variants confer vulnerability to SZ by affecting early embryonic events involved in cortical circuit assembly.

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AM-C and AD designed the experiments and wrote the paper. AM-C performed the experiments and analyzed the data with the technical help of Dr. Julien Prados, Dr. Sabine Bavarian, Christiane Aubry-Deuel, Joan Badia Cabré, and Aurélie Flaive. The bioimaging, flow cytometry, and genomics core facilities of the University of Geneva also provided valuable technical help for this project.

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