Disc1 Point Mutations in Mice Affect Development of the Cerebral Cortex

Frankie H. F. Lee,1,2 Marc P. Fadel,1,5 Kate Preston-Maher,6 Sabine P. Cordes,3,7 Steven J. Clapcote,8 David J. Price,6 John C. Roder,3,7 and Albert H. C. Wong1,2,4,5

1Centre for Addiction and Mental Health, Toronto, Ontario, Canada M5T 1R8, Departments of 2Pharmacology and 3Molecular Genetics and 4Institute of Medical Science, University of Toronto, Toronto, Ontario, Canada M5S 1A8, 5Department of Psychiatry, University of Toronto, Toronto, Ontario, Canada M5T 1R8, 6Centre for Integrative Physiology, University of Edinburgh, Edinburgh EH9 9XD, United Kingdom, 7Samuel Lunenfeld Research Institute, Mount Sinai Hospital, Toronto, Ontario, Canada M5G 1X5, and 8Institute of Membrane and Systems Biology, University of Leeds, Leeds LS2 9JT, United Kingdom

Disrupted-in-Schizophrenia 1 (DISC1) is a strong candidate gene for schizophrenia and other mental disorders. DISC1 regulates neurodevelopmental processes including neurogenesis, neuronal migration, neurite outgrowth, and neurotransmitter signaling. Abnormal neuronal morphology and cortical architecture are seen in human postmortem brain from patients with schizophrenia. However, the etiology and development of these histological abnormalities remain unclear. We analyzed the histology of two Disc1 mutant mice with point mutations (Q31L and L100P) and found a relative reduction in neuron number, decreased neurogenesis, and altered neuron distribution compared to wild-type littermates. Frontal cortical neurons have shorter dendrites and decreased surface area and spine density. Overall, the histology of Disc1 mutant mouse cortex is reminiscent of the findings in schizophrenia. These results provide further evidence that Disc1 participates in cortical development, including neurogenesis and neuron migration.

Introduction
Schizophrenia (SZ) is a common psychiatric disorder characterized by reduced hippocampal and cortical volume (Ross et al., 2006), abnormal cytoarchitecture (Kovalenko et al., 2003), reduced neuronal density in superficial cortical layers (Akbarian et al., 1993), decreased neuron size (Sweet et al., 2003), and reduced dendritic arborization (Young et al., 1998) and dendritic spine density (Garey et al., 1998). Many potential schizophrenia susceptibility genes have been identified (Wong and Van Tol, 2003; Ross et al., 2006), including Disrupted-in-Schizophrenia 1 (DISC1), first identified in a large Scottish family carrying a balanced (1q42.1:11q14.3) translocation cosegregating with major mental illnesses including SZ, bipolar disorder, and major depressive disorder (Millar et al., 2000). The DISC1 locus shows genetic linkage with SZ, and DISC1 variants show genetic association with SZ (Nakata et al., 2009; Rastogi et al., 2009; Schumacher et al., 2009). DISC1 acts as a regulatory scaffold protein, interacting with multiple binding partners including cytoskeletal and signaling proteins that link DISC1 to important brain developmental functions such as neurogenesis, neuron migration, neurite outgrowth, spine development, and neurotransmitter signaling (Cameron et al., 2007; Brandon et al., 2009).

The cortical histology of transgenic mice expressing various truncated mouse or human DISC1 fragments is similar to that seen in postmortem brain samples from patients with SZ (Hikida et al., 2007; Li et al., 2007; Pletnikov et al., 2008). Mice engineered to express truncated DISC1 have morphological alterations in medial prefrontal cortex and hippocampus (Kvajo et al., 2008). Because the truncated DISC1 gene is unique to the original translocation family and because common disease-associated DISC1 variants are single-nucleotide polymorphisms (SNPs) (Chubb et al., 2008), we sought to characterize brain histology in mice with Disc1 SNPs. Our group described previously two mutant Disc1 mice, each with a different SNP: Q31L (127A/T) and L100P (334T/C) (Clapcote et al., 2007). Both mutants have reduced brain volume, deficits in spatial working memory, and decreased prepulse inhibition. In addition, the Q31L mutants have abnormalities in social behavior and the forced swim test, while the L100P mice have increased open field activity (Clapcote et al., 2007). While the point mutations in our mice are not identical to human disease-associated variants, they may still provide important mechanistic insights into Disc1 SNP effects in contrast to the more drastic translocation mutations.

We undertook a comprehensive histological analysis of the cerebral cortex of Disc1 Q31L and L100P mutant mice. Our mutants have fewer neurons, decreased neuronal proliferation, and altered cortical layer positioning compared to wild-type (WT) littermates. Golgi staining showed shorter pyramidal neuron dendrite length in frontal cortex and reduced spine density in both frontal cortex and hippocampus. Our findings are similar to...
the abnormalities seen in postmortem human studies of SZ, and with transgenic Disc1 mutant mouse models. Our results provide evidence for the effects of DISC1 SNPs on neurodevelopment and cortical structure that may be useful for interpreting DISC1 genetics in the general SZ patient population, and represent a starting point for further investigations of molecular disease mechanisms in SZ.

Materials and Methods

Mice. N-ethyl-N-nitrosurea-mutagenized Disc1 mutant mouse lines on a C57BL/6 background (Q31L and L100P homozygous / − / ) were generated as described previously (Clapcote et al., 2007), and additional mice were bred for histological analysis at the Toronto Centre for Phenogenomics (TCP) (Toronto, Canada). WT littermates from both Q31L and L100P groups were combined and used as controls. All mouse protocols were approved by the TCP Animal Care Committee.

Bromodeoxyuridine labeling. Timed pregnant female mice were injected with bromodeoxyuridine (Brdu) (i.p., 50 mg/kg) at embryonic day 14 (E14) for proliferation experiments and at E12, E15, or E18 for investigation of neuronal positioning. Embryonic brains were harvested 24 h after BrdU injection at E15 and embedded in paraffin.Brains at postnatal day 21 (P21) were fixed in 4% paraformaldehyde overnight, cryoprotected in 30% sucrose, and frozen at −80°C before further processing.

Immunohistochemistry. Paraffin-embedded and frozen coronal sections of 5 and 10 μm thickness, respectively, were cut using a microtome cryostat system (Bright Instruments 5030). All sections were initially incubated in blocking solution (0.1 υ PBS, 1% Triton X-100, 0.5% Tween 20, 2% skim milk) or serum-free protein block (DakoCytomation) for 1 h at room temperature to reduce nonspecific background and then incubated with primary and secondary antibodies overnight at 4°C. The following primary antibodies were used: anti-NeuN (1:200; Millipore), anti-BrdU (1:200; BD Biosciences or Abcam), anti-Ki67 (1:200; Neomarkers), anti-Cux1 (1:200; Santa Cruz Biotechnology), and anti-Brn2 (1:200; Santa Cruz Biotechnology). Fluorescent secondary antibodies conjugated to Alexa 488 or Rhodamine Red-X (1:200; Invitrogen) or Cy3 (1:100; Jackson ImmunoResearch Laboratories) were used for detection of primary antibodies.

Golgi–Cox staining. Golgi–Cox staining was performed as described previously (Gibb and Kolb, 1998). In brief, adult mice (age 6–8 weeks) were anesthetized with xylazine/ketamine (10 ml/kg) and intracardially perfused with 0.9% saline. Brains were removed and immersed in Golgi–Cox solution in the dark for 14 d before transfer to 30% sucrose solution and frozen at −80°C be- fore further processing.

Analysis of immunohistochemistry: neuron number, distribution within the cortex, and neurogenesis. Immunohistochemistry (IHC) images of the whole cortex were captured using a confocal microscope (Zeiss LSM510 Meta) at 5 × or 25 × magnification. All fluorescently labeled images were converted to gray scale and normalized to background staining. Sections chosen for analysis were anatomically matched along the rostral-caudal axis for all samples. Regions of interest (ROIs) were positioned over the cortex as sampling windows. A two-dimensional cell counting approach of random sampling was used to provide accurate estimates of cell densities (Benes and Lange, 2001). For all IHC images, neurons were counted using the ITCN plugin for ImageJ (http://rsb.info.nih.gov/ij/) (ICTN parameters: width, 10 pixels; minimum distance, 2 pixels; threshold, 1 pixel). Specific procedures for defining areas of analysis differed slightly for each antibody, since each was chosen to address different questions. These procedures are described in detail below.

NeuN antibody-labeled images were used to examine overall neuron numbers in the cortex. Eight rectangular ROIs of fixed size (500 μm high × 250 μm wide), with the long axis perpendicular to the pial surface, were outlined throughout the neocortex from medial to lateral (see Fig. 1A). Each ROI was straightened with ImageJ software, and the number of neurons in the ROI was counted. BrdU and Ki67 were used to assess neurogenesis in both the embryonic and the postnatal cerebral cortex. For neuronal progenitor proliferation, fluorescently labeled cells in the ventricular zone (VZ) and subventricular zone (SVZ) were counted in a fixed area ROI of 100 × 120 μm. Similarly with P21 brains, BrdU-labeled cells in the frontal cortex were counted in an ROI of fixed width (500 μm) but of variable length, corresponding to the thickness of the cortex. Frontal cortical regions were defined according to the Golgi Atlas of the Postnatal Mouse Brain (Valverde, 1998). These BrdU ROIs were subdivided into 10 deciles along the axis perpendicular to the pia to assess the distribution of each wave of newly born neurons within the layers of cortex.

Cux1 and Brn2 antibodies were used to investigate whether neurons normally destined for the superficial cortical layers II and III would instead be seen in deeper layers (IV–VI) in Disc1 mutant mice. In both Cux1- and Brn2-labeled sections, two ROIs with a fixed width of 500 μm and of variable length spanning the thickness of the cortex were delineated. Each ROI was subdivided into eight equal regions from the pia to the inner border of the cortex to assess neuron distribution across the layers of the cortex.

Neuron morphology and dendritic spines. For morphometric analyses of individual neurons, Golgi images at 40 × magnification were captured under brightfield illumination with a Nikon Eclipse E600 microscope.
Neurons were chosen based on the following criteria: (1) fully visible and characterized by clear, distinct morphology; (2) all dendrites seen within the 40× magnification field; and (3) only pyramidal neurons in layers III and V of the frontal cortex and CA1 area of the hippocampus, as shown in the Golgi Atlas of the Postnatal Mouse Brain (Valverde, 1998). A z-stack of different focal lengths for each individual neuron was generated to capture the three-dimensional dendritic branching tree in different planes. Acquisition parameters were kept the same for all images. The neurites of each neuron were traced, and the length and surface area were estimated using Neuroromantic software (http://www.rdg.ac.uk/neuroromantic). All parameters were further normalized to soma surface area for comparison.

Sholl analysis provides a quantitative measure of the radial distribution of neuronal dendritic arborization (Sholl, 1953). Using ImageJ, we created 15 concentric and equidistant circles (8 μm separation of each radius) centered at the perikaryon and then counted the number of dendritic intersections at each circle of increasing radius. The log of the number of intersections per circle area versus the circle radius was plotted (the semilog Sholl method). The slope of the regression line (κ = Sholl regression coefficient) is a measure of the decay rate of the number of branches with distance from the soma (Sholl, 1953). The Schoenen ramification index (maximum number of intersections/number of primary dendrites), a measure of the ramification richness for each neuron (Schoenen, 1982), and the number of dendritic bifurcations provide important information on the degree of dendritic branching complexity.

Spine density was measured with Golgi-stained images captured at 100× magnification (Nikon Eclipse E600). Spines were counted only on the apical dendrites of pyramidal neurons in layers III and V of frontal cortex and CA1 of hippocampus. Spine density was expressed as the number of spines per dendritic length (micrometer). All images for quantification were blinded before analysis. Statistical analysis. Statistical differences among different mutant lines and across genotypic groups against various measured parameters were determined using one-way or two-way ANOVA (SPSS 13.0), followed by Bonferroni’s correction for multiple testing. To further confirm significance, Student’s two-tailed t test was performed in comparing two sets of data. Data are expressed as mean ± SEM. A significance level of p < 0.05 was used for all analyses.

Results
Fewer NeuN-positive neurons in Q31L and L100P (−/−) mutant mice
Experimental evidence is accumulating for the role of DISC1 in neuronal proliferation and neuronal migration (Kamiya et al., 2005; Mao et al., 2009; Singh et al., 2010). We therefore examined the number of neurons throughout the neocortex along the medial-lateral axis. Neurons were labeled with antibodies to NeuN, a neuronal marker, and then counted in WT, Q31L, and L100P mice (Fig. 1A). Although NeuN does not label some types of neuron such as Purkinje cells, the high specificity and dense labeling of cortical neurons and interneurons provide a suitable measure of neuronal density (Wolf et al., 1996). We observed significantly fewer NeuN-labeled neurons in both Q31L (7260 ± 863; p < 0.001)
and L100P mutant mice (7638 ± 346; p < 0.001) across the neocortex when compared with WT mice (8636 ± 522) (Fig. 1B). This observation of fewer NeuN-labeled neurons in Q31L and L100P compared to WT was also seen within each individual ROI, spanning from medial to lateral (Fig. 1C).

Reduced neuronal proliferation in Q31L and L100P compared to WT

We next determined whether the differences in the number of NeuN-positive neurons could be related to differences in neurogenesis between WT, Q31L, and L100P mice. First, BrdU was injected into pregnant dams at E14 and embryonic brains were collected at E15 to analyze neural progenitor proliferation in SVZ/VZ of the embryonic cortex (Fig. 2A). Disc1 mutants had fewer BrdU + cells when compared to WT controls (Q31L: 70 ± 18 and L100P: 66 ± 24 vs WT: 102 ± 33; p < 0.001) (Fig. 2B, C). A similar pattern was observed with Ki67 immunostaining, further confirming a decrease in proliferation (Fig. 2D). As previous studies have shown that DISC1 knockdown causes premature neuronal differentiation (Mao et al., 2009), we investigated whether our mutations have similar effects. We identified cells that had left the cell cycle as BrdU positive and Ki67 negative. The percentage of BrdU+/Ki67− cells was only slightly increased in L100P mutants (97.41 ± 2.91%; p = 0.018) but not Q31L (96.75 ± 3.04%) compared to WT (95.26 ± 3.00%) (Fig. 2E). These subtle effects suggest that our Disc1 mutations may not have strong effects on the timing of neuronal differentiation.

To observe the numbers and eventual location of neurons born at different times during embryonic corticogenesis, BrdU was injected at three different time points, E12, E15, and E18, and analyzed at P21. Coinmunostaining with NeuN confirmed the neuronal identity of BrdU-labeled cells (Fig. 3A). The average total number of BrdU-positive cells was significantly lower in Q31L and L100P cortices compared to WT at E12 (Q31L: 61 ± 14 and L100P: 64 ± 14 vs WT: 149 ± 25; p < 0.001) and at E15 (Q31L: 185 ± 21 and L100P: 199 ± 40 vs WT: 254 ± 31; p < 0.001) (Fig. 3B). Interestingly, E18-injected BrdU + neurons were only significantly decreased in Q31L (Q31L: 37 ± 17 vs WT: 54 ± 7; p < 0.05) but not in L100P mutants (42 ± 21) (Fig. 3B). Thus our results suggest that these single point mutations within Disc1 are associated with decreased neuronal proliferation.

Because neurons destined for more superficial layers are born later and because there is evidence that DISC1 may affect neuronal migration (Marin and Rubenstein, 2003; Kamiya et al., 2005; Young-Pearse et al., 2010), we next examined the cortical distribution and position of BrdU-labeled cells for each time point of BrdU injection. Confocal imaging and quantification revealed that BrdU + cells were located in deeper cortical layers for E12 and E15 time points in both mutants when compared to WT (p = 0.01). In contrast, the distribution of E18-injected BrdU-labeled cells was similar in all cortical layers between WT and mutants, with only a small number of BrdU-positive cells observed in both mutants in deep cortical layers but not in control mice. All data are presented as mean ± SEM; *p < 0.05, **p < 0.01 versus WT, I—VI, Cortical layers I–VI.

Figure 3. Reduced BrdU incorporation and mispositioning of cortical neurons in Disc1 mutant mice. A, BrdU was injected into E12, E15, and E18 pregnant females of WT, Q31L, and L100P mutants. Mice were killed at P21 for BrdU and NeuN immunostaining. Scale bar, 250 μm. B, The total number of BrdU + cells was significantly lower in both E12 and E15 BrdU-injected Q31L and L100P mutants, but only Q31L showed fewer labeled cells with BrdU injection at E18 when compared to WT (n = 5–20; t test, p < 0.01). C, The distribution of BrdU-labeled cells was analyzed across cortical layers in all BrdU-injected time points. A rectangular ROI of fixed width (500 μm), spanning the thickness of the frontal cortex, was defined in each slice. This rectangle was then divided into 10 equal deciles along the axis perpendicular to the pial surface. The number of BrdU-positive cells in each decile was counted, and is shown as a percentage of the number in entire ROI. There were more BrdU-positive cells in deeper layers for E12 and E15 time points in both mutants when compared to WT (p = 0.01). Two-way ANOVA, p < 0.01. D, At E18, the distribution of BrdU-labeled cells was similar in all cortical layers between WT and mutants, with only a small number of BrdU-positive cells observed in both mutants in deep cortical layers but not in control mice. All data are presented as mean ± SEM; *p < 0.05, **p < 0.01 versus WT, I—VI, Cortical layers I–VI.
To further examine the relationship between DISC1 and neuronal positioning, we performed IHC with two layer II/III-specific protein markers, Cux1 and Brn2 (Molyneaux et al., 2007). When compared to WT, both Cux1- and Brn2-labeled neurons in Q31L and L100P animals were further away from the pia (Fig. 4A). To quantify the positions of Cux1- and Brn2-labeled cells, two rectangles were positioned over the neocortical region of fluorescent staining, and each was divided into eight equal octants (spanning superficial to deep layers) (Fig. 4B). The percentage of Cux1-labeled cells was significantly higher in both octant 1 and octant 3 of WT littermates (octant 1: 12.83 ± 3.55%; octant 3: 37.37 ± 4.52%) compared to Q31L (octant 1: 5.99 ± 1.76%; octant 3: 31.42 ± 4.07%, p < 0.01) and L100P (octant 1: 7.43 ± 2.61%, p < 0.01; octant 3: 35.11 ± 4.81%, p < 0.05) mice. In contrast, WT controls (12.85 ± 4.84%) displayed a significantly lower proportion of Cux1+ cells in octant 4 than in Q31L (22.53 ± 5.98%, p < 0.01) and L100P (22.04 ± 9.11%, p < 0.01) (Fig. 4B). For Brn2, WT mice showed a significantly higher proportion of Brn2+ cells in superficial cortical layers (octants 1 and 2), and significantly fewer such cells in the deep layers of octants 4, 5, and 6 when compared to both mutants (Fig. 4C). Moreover, we measured the ratio of distance between the inner edge of Cux1fluorescently labeled cells and pia mater versus total cortical thickness and saw a higher ratio in Q31L (0.433 ± 0.039, p < 0.01) and L100P (0.432 ± 0.041, p < 0.01) mutants versus WT (0.417 ± 0.031; p < 0.01) (Fig. 4D). These data suggest that the localization of cortical neurons in Q31L and L100P mice is altered compared to WT mice.

Differences in frontal cortical dendritic morphology in Q31L and L100P mice
There is strong evidence that DISC1 regulates neurite outgrowth and dendritic arborization; therefore, we performed a detailed morphological analysis of dendritic trees for individual neurons in Disc1 Q31L and L100P mutants and WT mice. Golgi staining provides a clear and complete image for a subgroup of neurons without interference by neighboring neurons. Representative neurons from WT, Q31L, and L100P mice are shown in Figure 5A. We observed a significantly shorter apical dendritic length (ADL) in Q31L (208.9 ± 68.9 μm, p < 0.01) and L100P mutants (242.4 ± 94.9 μm, p = 0.03) when compared to WT (328.3 ± 55.5 μm). Consistently, basal dendritic length (BDL) showed a similar trend with significant differences in Q31L (690.6 ± 100.5 μm, p < 0.01) and L100P (786.8 ± 90.1 μm, p = 0.021) versus WT (884 ± 109.6 μm) (Fig. 5B). We also found a significantly lower total dendritic surface area (DSA) of
Q31L (4250.9 ± 514.1 μm², p < 0.001) and L100P mutants (4966.3 ± 550.7 μm², p = 0.005) when compared to WT (5708.5 ± 439.5 μm²) (Fig. 5B).

Cell soma size has been reported to correlate with dendritic structure (Somogyi and Klausberger, 2005). Thus, we normalized the measured parameters described above to soma surface area. Interestingly, neurite outgrowth was significantly lower by 40.5% (p = 0.014), 23% (p = 0.042), and 27.7% (p = 0.007) in ADL, BDL, and DSA, respectively, within Q31L mutants only, while L100P mice showed a similar but nonsignificant trend (Fig. 5C).

We evaluated dendritic arbor complexity in our Disc1 mutant mice versus WT via Sholl analysis. There were no significant differences in the ramification index (Q31L: 2.3 ± 0.2, p = 0.37; L100P: 2.8 ± 0.3, p = 0.73; vs WT: 2.6 ± 0.3), the dendritic bifurcations (Q31L: 7.2 ± 2.1, p = 0.39; L100P: 7.5 ± 0.5, p = 0.48; vs WT: 8.1 ± 2.0), Sholl’s regression coefficient (Q31L: 0.089 ± 0.005, p = 0.97; L100P: 0.091 ± 0.007, p = 0.92; vs WT: 0.090 ± 0.004), or the number of dendritic intersections per radial segments between Q31L and L100P and WT controls (Fig. 5D–F).

In addition to the frontal cortex, the hippocampus is another important brain area of interest in SZ (Harrison, 2004). Hence, we extended our morphological analyses to pyramidal neurons in hippocampal CA1 regions. In our study, we did not observe any alterations in dendritic length, surface area, ramification index, number of bifurcations of dendrites, and Sholl’s regression coefficient in either mutant line within the hippocampus (Fig. 6).

**Lower spine density in frontal cortex and hippocampus in Q31L and L100P Disc1 mutant mice**

Cognitive deficits associated with SZ have been attributed to altered synaptic transmission and plasticity in which dendritic spines play a critical role (Calabrese et al., 2006). DISC1 is involved in dendritic spine development via Kalirin-7 (Penzes and Jones, 2008; Hayashi-Takagi et al., 2010). To determine whether our Disc1 point mutations affect dendritic spine development, we measured dendritic spine density in the frontal cortex and hippocampus of Q31L, L100P, and WT mice. Golgi images of 100× magnification were captured, and all spine types were counted (Fig. 7A). Both Disc1 mutants had significantly reduced spine density in the frontal cortex with an average spine density of 0.39 ± 0.063 spines/μm² in Q31L (p = 0.001) and 0.391 ± 0.057 spines/μm² in L100P (p < 0.001) compared to 0.449 ± 0.071 spines/μm² in WT (Fig. 7B). Similarly, spine density was also reduced in the hippocampus (Q31L: 0.363 ± 0.087 spines/μm² and L100P: 0.427 ± 0.085 spines/μm² vs WT: 0.543 ± 0.11 spines/μm²; p < 0.001). Thus, DISC1 mutations appear to affect spine density in both the frontal cortex and hippocampus.

**Discussion**

There is increasing evidence for a strong association between DISC1 and several major mental illnesses. The mechanism by which DISC1 gene variants produce both cellular and behavioral...
abnormalities is still unclear. We used two previously described mouse lines with point mutations in Disc1 that have behavioral changes relevant to SZ and depression (Clapcote et al., 2007). In this study, we report a relative decrease in neuron number and decreased neuronal proliferation in the Disc1 mutants compared to WT mice. The mutant mice have differences in neuron positioning and morphology similar to some findings in human SZ histopathological studies (Akbarian et al., 1993; Ross et al., 2006). However, other histological abnormalities observed in postmortem schizophrenia brain, such as interneuron deficits, have not yet been investigated in our Disc1 mutant mice.

We found relatively fewer neurons and decreased neuronal proliferation in Q31L and L100P mutant mice compared to WT. Decreased neuronal density is a common finding of postmortem studies on patients with SZ. Reductions in neuron density in the primary visual cortex (Dorph-Petersen et al., 2007) and reduced glutamatergic neurons in the orbitofrontal cortex have been reported (Garey, 2010). Presently, the DISC1 status of the patients reported in those studies is unknown. Human studies combining genetic markers and histopathological analysis are required.

DISC1 has been well established as a regulator of neurogenesis (Mao et al., 2009). DISC1 participates in a glycogen synthase kinase 3β (GSK3β) signaling pathway involving β-catenin via a direct interaction with GSK3β at two different domains of DISC1, spanning amino acids 1–220 and 356–595. Moreover, DISC1 knockdown results in a reduction of proliferation progenitors likely caused by early cell cycle exit (Mao et al., 2009). However, our Disc1 mutations showed a decrease in neuronal proliferation but not premature neuronal differentiation. It is possible that the Q31L and L100P mutations in DISC1 may only affect part of its interaction with GSK3β and that premature cell cycle exit may not be the sole determinant of neuronal proliferation. Recently, the L100P Disc1 mutant mouse was shown to have reduced interaction with both GSK3α and β (Lipina et al., 2011). Furthermore, genetic and pharmacological inhibition of GSK3 activity rescued DISC1-mediated behavioral effects in these mice.

Examination of neuronal distribution using layer-specific protein markers revealed altered neuron location in Q31L and L100P mice compared to WT. Cortical neuronal positioning can be affected by changes in neurogenesis and neuronal migration. Later-born neurons migrate to more superficial layers of the cortex through radial migration (Marín and Rubenstein, 2003). As DISC1 regulates neurogenesis and neuronal migration, we hypothesized that aberrant neuronal distribution in the cortex may be due to DISC1-mediated effects on both processes (Kamiya et al., 2005; Mao et al., 2009). It was recently shown that DISC1 may participate in neurogenesis and neuronal migration via separate and distinct signaling pathways (Singh et al., 2010). Abnormal cortical cytoarchitecture may also result from malfunctioning of the cytoskeletal machinery mediating neuronal migration. Recent studies have reported several important DISC1-interacting candidates with critical roles in the regulation of neuronal migration, including pericentriolar material 1 (PC1M1) (Kamiya et al., 2008), amyloid precursor protein (APP) (Young-Pearse et al., 2010), neuregulin-1/ErbB4 (Jaaro-Peled et al., 2009), and LIS1/NDEL1 (Morris et al., 2003; Wynshaw-Boris, 2007). However, complete details of
the mechanisms by which neurogenesis and neuronal migration interact to modulate cortical cytoarchitecture remain to be determined.

The observed differences in frontal cortical neuron morphology in the *Disc1* mutants compared to WT may also be mediated by DISC1 interactions with the actin and microtubule cytoskeleton (Ishizuka et al., 2006). Loss of normal DISC1 function or expression of mutant DISC1 disrupts its interaction with NDEL1 and causes abnormal neurite outgrowth in PC12 cells (Ozeki et al., 2003; Kamiya et al., 2005). Consistent with this, transgenic mice expressing truncated *Disc1* have an inhibition of neurite outgrowth and a reduction of apical dendritic length (Kvajo et al., 2008; Shen et al., 2008). We reported similar findings of significant reductions in dendritic length and surface area in our *Disc1* mutants, further supporting the role of DISC1 in neurodevelopment. Recent studies with transgenic *Disc1* mice have shown a disturbance in neuronal arborization both in the developing cerebral cortex and hippocampus (Kamiya et al., 2005; Li et al., 2007; Niwa et al., 2010).

In contrast, Kvajo reported no significant changes in dendritic complexity with their mice expressing truncated *Disc1* (Kvajo et al., 2008). Our *Disc1* point mutations also showed no alterations in dendritic branching pattern, consistent with Kvajo et al. Dendritic arbor development is a complicated and strictly regulated multistep process involving the following: (1) neurite initiation, outgrowth and guidance; (2) branching and synapse formation; and (3) cytoskeleton stabilization (Urbanska et al., 2008). Proper formation and stabilization of dendritic arbors requires various intrinsic and extrinsic signals (Urbanska et al., 2008).

Hippocampal neurons in SZ have been reported to have subtle morphological changes in size, organization, and perhaps shape (Harrison, 2004). Other *Disc1* mutant mouse models show similar features (Li et al., 2007; Shen et al., 2008), and human SNPs in *DISC1* have been associated with altered hippocampal structure and cognitive function (Callicott et al., 2005). However, not all SZ postmortem studies are consistent in finding abnormal hippocampal neuronal morphology (Benes et al., 1998). We did not detect any significant changes in hippocampal neuronal morphology with our *Disc1* mutations, similar to the findings from Kvajo (Kvajo et al., 2008). Duan et al. (2007) recently demonstrated an acceleration of neuronal integration when downregulating DISC1 in adult hippocampal neurons. As the developmental origin of hippocampus is distinct from the cerebral cortex, DISC1 may modulate different developmental programs in the hippocampus.

Synaptic pathology has been proposed as a cause for cognitive deficits in SZ. Dendritic spines are the postsynaptic targets for synaptic transmission, and decreased spine density in prefrontal cortical and subventricular pyramidal neurons has been reported in human postmortem SZ studies (Garey et al., 1998; Glantz and Lewis, 2000; Rosoklija et al., 2000). Spine density can be used as a measure of neural connectivity (Benes, 2000). DISC1 interacts with Kalirin-7 to modulate Rac1, an important regulator of dendritic spine development and functional plasticity (Penzes and Jones, 2008; Hayashi-Takagi et al., 2010). Long-term suppression of DISC1 can lead to spine shrinkage in primary cortical neurons (Hayashi-Takagi et al., 2010). Our observations are consistent with these previous data, since we see a significant reduction in spine density in both mutants for frontal cortex and hippocampus. Our results further confirm that DISC1 regulates spine development and may represent a possible link between DISC1 genetic variants and cognitive deficits observed in SZ.

Intriguingly, although Q31L and L100P mutant mice have distinct behavioral abnormalities, they have similar histological deficits. Subtle disruption of neuronal architecture and connections can have diverse effects on complex behaviors and on the activity of other brain regions. Q31L and L100P mutants also have different phosphodiesterase 4B (PDE4B) activity and Disc1-PDE4B binding (Clapcote et al., 2007). The discrepancy between histology and behavior may require further histological and biochemical characterization of the effects of these point mutations. Our study provides a general overview of cortical histology, development, and neuronal morphology in two independent *Disc1* single-point mutant mouse lines. The function of *Disc1* in neuron development has mostly been investigated by observing the effect of drastic reductions in DISC1 expression or by expression of a truncated protein. Our study is novel in characterizing histopathological findings in two mouse lines with *Disc1* SNPs. Although the human disease-associated *DISC1* SNPs are not the same as the Q31L and L100P mutations in our mice, we argue that our mouse SNPs are more similar to the *DISC1* SNPs in the general human population than the truncated Disc1 mutants. Previous studies with truncated DISC1 or severe suppression of DISC1 expression are more relevant to understanding the pathophysiology of the original Scottish translocation pedigree. Ongoing experiments to further understand the molecular mechanisms by which DISC1 regulates brain development are...
required to understand the causal links between DISC1 variants and susceptibility to schizophrenia.

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