Cross-regulation of Connexin43 and β-catenin influences differentiation of human neural progenitor cells

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Connexin43 (Cx43) is the most widely and abundantly expressed gap junction (GJ) protein and it is strongly associated with the regulation of cell cycle progression. Emerging roles for Cx43 in cell adhesion and migration during neural differentiation have also been recently recognized, and this has emphasized the involvement of Cx43 in different physiological processes beyond its role as a GJ protein. In this study, we explore the function of Cx43 in the differentiation of human neural progenitor cells (hNPCs) using viral vectors that mediate the overexpression or knockdown of the protein. Results showed that in the absence of this protein fetal cortex-derived hNPCs differentiated toward a neuronal phenotype at expenses of a glial phenotype. Furthermore, the silencing of Cx43 did not affect hNPC proliferation rate or numbers of apoptotic cells. The increase in the number of neurons was not recapitulated when GJ intercellular communications were pharmacologically blocked, and this suggested that Cx43 was influencing hNPCs differentiation with a GJ-independent effect. In addition, Cx43 knockdown significantly increased β-catenin signaling, which has been shown to regulate the transcription of pro-neuronal genes during embryonic neural development. Our results add further support to the hypothesis that Cx43 protein itself regulates key signaling pathways during development and neurogenesis beyond its role as GJ protein.

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Connexin43 (Cx43) protein has been found to be expressed early during mouse embryogenesis, at the preimplantation stages. Indeed when organogenesis occurs, the protein is reported to be preferentially upregulated in the ectoderm rather than mesodermal and endodermal layer. Interestingly during neurogenesis, Cx43 is both temporally and spatially regulated with a prominent expression in neural progenitor cells followed by a decrease when neurons start to differentiate. The presence of Cx43 at the early stage of development together with its dual regulation during neurogenesis strongly suggest this protein is greatly involved in development events and the precise regulation of the protein expression may be crucial to ensure a proper maturation. Cx43 is one of the proteins that form Gap junction (GJ) channels between adjacent cells, and this type of cellular communication is critical to several processes ranging from tissue homeostasis and synchronization of cellular activity, to the establishment and maintenance of microenvironments or niches. Several Cx proteins have been identified that form GJs, intriguingly Cx43 is the most widely and abundantly expressed isoform with a unique long C-terminal (CT) tail. This has been reported to be the binding site of multiple proteins, which dynamically interact, and participate in the regulation of physiological events (for review, see Giepmans5). The importance of Cx43 in regulating cell adhesiveness and directionality of migration has recently been described (for review, see Matsuuchi and Naus9). Cx43 is present on radial glia fibers at the points where they contact with migrating neurons and Cx43 conditional knockout mice show delayed neuronal migration together with impairment in the lamination of the neocortex. A brain transcriptome analysis identified gene transcripts altered in Cx43 null and Cx43 heterozygous mice, highlighting a change in proteins regulating cell growth, neuronal survival and differentiation. Therefore, it is clear that together with physically mediating neurogenesis events, Cx43 might also act as a central role in the control of growth.

Neural stem cells have been identified in both the developing and adult nervous system. In this study, we use neural progenitor cells (NPCs) derived from human fetal brain to unravel the role of Cx43 in neuronal differentiation. We found that Cx43 protein knockdown increased the number of migrating neurons 7 and Cx43 conditional knockout mice show delayed neuronal migration together with impairment in the lamination of the neocortex. A brain transcriptome analysis identified gene transcripts altered in Cx43 null and Cx43 heterozygous mice, highlighting a change in proteins regulating cell growth, neuronal survival and differentiation. Therefore, it is clear that together with physically mediating neurogenesis events, Cx43 might also act as a central role in the control of growth.

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of neurons differentiated from human NPCs (hNPCs). We further demonstrated that the knockdown of the protein enhanced the transcriptional activity of β-catenin, which is a key mediator of the canonical WNT pathway and it has been shown to regulate transcription of pro-neuronal genes.18–20

Results

Human Cx43 expression in an in vitro model of neurogenesis. We analyzed the spatiotemporal expression of the human Cx43 (hCx43) protein in hNPCs. Those cells provide a valuable source of neural tissue and an in vitro model for studying neurogenesis.21 RT-PCR was performed in multipotent hNPCs, as well as following 7 and 14 days differentiation. Results indicated a specific pattern of Cx43 mRNA expression during hNPCs differentiation with higher levels of Cx43 mRNA in undifferentiated and 7 days differentiated cells (0.93 ± 0.2; 0.69 ± 0.1), followed by a decrease at 14 days of differentiation (0.3 ± 0.2; Figures 1a and b). There was no change in the mRNA levels of the neural markers Mash1 and Pax6 between undifferentiated and 7 days differentiated hNPCs (undifferentiated Pax6 1.7 ± 0.8, MASH1 1.7 ± 0.9; 7 days differentiated Pax6 1.5 ± 0.2, MASH1 1.4 ± 0.2). At 14 days of differentiation, we observed a small reduction in Pax6 expression (1.0 ± 0.2), whereas Mash1 levels remained consistent with previous

Figure 1  Cx43 expression in an in vitro model of neurogenesis. (a) Representative RT-PCR bands of undifferentiated, 7 and 14 days differentiated hNPCs for Cx43, Pax6 and MASH1 mRNAs. GAPDH is used as housekeeping gene. (b) Quantification of RT-PCR bands for hCx43, Pax6 and MASH1 genes. Values are the means of three independent experiments and error bars represent S.E.M. Values were expressed as arbitrary units normalized to the GAPDH values. (c) Fluorescence microscopy of undifferentiated hNPCs stained with Cx43 (green), nestin or Tuj1 (red) antibodies. Cell nuclei are indicated by DAPI (blue) staining. Scale bar 50 μm. (d) Higher magnification of the area around 7 days differentiated sphere. Cx43 (red) on GFAP (green)-positive cells expressed at the contact points with migrating Tuj1 (blue) neurons (arrowheads in boxed areas). Scale bar 50 μm
expression (1.4 ± 0.3; Figures 1a and b). Those results suggested that as differentiation progressed cell were maintaining a neural phenotype with a decrease in Cx43 expression.

Multipotent hNPCs revealed an intense Cx43 immuno-labeling throughout the neurosphere, these cells were all undifferentiated neuroepithelial cells as indicated from the nestin staining (Figure 1c). Following 7 days of differentiation, immunostaining revealed presence of TuJ1 and GFAP-positive cells with the Cx43 protein specifically localized at the contact points of migrating neurons and GFAP-positive process (Figure 1d).

**Lentiviral vector mediated specific knockdown of hCx43.** To induce a specific hCx43 protein knockdown, we designed four short hairpin RNA (shRNA) constructs (Supplementary Table 1) and cloned them into pCR 2.1 TOPO. The efficiency of knockdown mediated by each construct was tested in HEK293 cells by co-transfection of shRNA plasmid together with an hCx43-overexpressing vector. This co-transfection was required as HEK293 cells do not express detectable levels of endogenous Cx43 protein. Immunoblot analysis was performed to detect the shRNA sequence allowing the maximum hCx43 protein knockdown (Supplementary Figures 2A and B). Subsequently, the shRNA sequence mediating the most effective knockdown was cloned into a lentiviral vector backbone to generate viral particles (Figure 2a). Viral efficacy was tested using multiplicity of infections (MOIs) of 1 and 5 in HEK293 cells. Again an hCx43shRNA virus was used together with an hCx43-overexpressing virus (Figures 2b and c). High transduction efficiency was obtained at both MOIs tested (Figure 2b), however, western blot results revealed a higher hCx43 protein knockdown at MOI 5 (Figure 2c).

**Knockdown of hCx43 regulates hNPC neuronal differentiation.** We sought to investigate the contribution of hCx43 to neurogenesis by knocking down this protein in hNPCs and then inducing their differentiation. Figure 3a illustrates the experimental model used in which viral transduced hNPCs were expanded for 10 days from the day of lentiviral vector transduction followed by differentiation, which was carried out for 2 h and for 7 days, respectively. Strong enhanced green fluorescent protein (EGFP) fluorescence was detectable at all-time points tested (Figure 3b) indicating continuous shRNA expression. Immunocytochemistry performed in 7 days differentiated cells revealed a robust knockdown of hCx43 protein compared with EGFP control (Figure 3c). Quantification of nestin-positive cells after 2 h of differentiation revealed that the majority of cells still retained an immature neural phenotype (Supplementary Figure 3A). The shRNA-mediated knockdown of hCx43 resulted in a significant increase in the number of TuJ1-positive cells compared with cells transduced with either EGFP or a scrambled control (SC; Figures 4a and b and Figures 5a and b) for both 2 h (shCx43 6 ± 0.9%; EGFP 2 ± 0.3%; SC 1 ± 0.2%; P < 0.01) and 7 days (shCx43 31.4 ± 5.0%; EGFP 9.1 ± 0.4%; SC 7.0 ± 0.7%; P < 0.01) differentiation. The number of GFAP-positive cells was significantly reduced in cells differentiated for 2 h compared with EGFP control (shCx43 66 ± 1.0%; EGFP 85.6 ± 2.8%; SC 84.4 ± 3.0%; P < 0.01) and in 7 days differentiated cells compared with EGFP and to SC transduced cells (shCx43 66 ± 1.0%; EGFP 85.6 ± 2.8%; SC 84.4 ± 3.0%; P < 0.01), (Figures 5c and d). Characterization of neurons derived from 7 days differentiated hNPCs revealed a great majority of GABA-positive neurons (Supplementary Figure 4). However, hCx43 knockdown did not affect the number of actively dividing cells as indicated by Ki67 expression (Figures 5e and f). As we detected an alteration in cell fate commitment already at 2 h of hNPCs differentiation, we decided to investigate whether Cx43 knockdown could affect the viability and proliferation at this earlier stage.
Figure 3 Experimental model and validation of ShRNAhCx43 virus in neural stem cell culture. (a) Schematic representation of experimental method used to transduce hNPCs. hNPCs were transduced at day 0 (D0) and expanded until D10 (green line). Subsequently, they were differentiated for 2 h or 7 days, respectively, (red lines). (b) EGFP expression in undifferentiated, 2 h and 7 days differentiated hNPCs. Scale bar 50 μm. (c) Fluoresce microscopy for Cx43 (red) and EGFP (green) of 7 days differentiated hNPCs in EGFP and shRNAhCx43 transduced cells. Cell nuclei are indicated by DAPI (blue) staining. Scale bar 50 μm

To assess cell proliferation, 5'-bromo-2-deoxyuridine (BrdU) was added to hNPC medium at the moment of viral transduction and left in culture for the expansion time (10 days). Cells were differentiated for 2 h and analysis indicated that hCx43 absence did not affect the proliferation (Supplementary Figure 3B). For cell viability analysis, TUNEL assay was performed on cells differentiated for 2 h, and this also revealed no difference between conditions (Supplementary Figure 3C). Finally, morphometric analysis of neurons derived from hCx43 and control transduced hNPCs did not reveal any change in neuronal complexity between groups (Supplementary Figure 5), indicating that the absence of Cx43 increased the number of differentiated neurons, however, it did not advance neuronal maturation.

Increase in neuronal differentiation is caused by the absence of hCx43 protein. In order to evaluate whether the increase in neuronal number was due to the specific absence of hCx43 protein or to a general reduction of hCx43-mediated GJ intercellular communication, we used a pharmacological blockade of GJ using a well-established GJ blocker carbenoxolone (CBX). CBX used at 50 μM largely impaired the spread of GJ permeable dye Cascade blue (Supplementary Figure 6), thus we incubate hNPCs for 10 days with CBX and we used its inactive analog glycyrrhizic acid (GZA) as control. Cells were differentiated for 2 h and results showed in Figure 6 indicated that CBX did not affect the number of Tuj1 or GFAP-positive cells (Figure 6a and b), nor cell proliferation (Figure 6c). Thus, these results suggested that the effect on hNPCs neuronal fate was independent of GJ intercellular communications, but dependent on the expression of the hCx43 protein itself.

hCx43 contributes to the regulation of β-catenin transcription. We confirmed a role for hCx43 in hNPCs differentiation and we also demonstrated that the protein itself was responsible for the increase in the number of neurons and a concomitant decrease in glial cells. We next wanted to investigate a possible contribution of hCx43 to the canonical Wnt/β-catenin pathway, which has been shown to be involved in many aspects of embryonic development and neurogenesis. In addition, β-catenin has also been reported to regulate hCx43 transcription. We first performed immunocytochemistry to determine the levels of hCx43 and β-catenin proteins in hNPCs derived neurospheres, which had been differentiated for 2 h. Results revealed intense immunolabeling of both proteins, which was mainly localized in the cell membrane (Figure 7a). We then assessed if the knockdown of endogenous hCx43 or its forced expression was affecting β-catenin protein expression. We transduced hNPCs with our shCx43 or our viral vector containing the hCx43 full-length sequence or our EGFP control and we collected proteins for immunoblot analysis. Results indicated a significant increase in β-catenin protein expression when hCx43 was silenced (Figures 7b–d; shCx43 2.2 ± 0.14; EGFP 1 ± 0.0; P ≤ 0.01). Then, we tested the hypothesis that in the absence of hCx43 the increase in β-catenin protein expression observed was due to an increase in the uncomplexed form of the protein. Free cytoplasmic β-catenin accumulates in the cytoplasm and then translocates to the nucleus where it mediates gene transcription binding to the TCF/LEF transcription factor motifs. Therefore, to measure β-catenin transcriptional activity we used a well-established gene reporter assay.

In the assay, a plasmid containing the TCF/LEF-binding site upstream of a Firefly luciferase ORF (Addgene, Cambridge, MA, USA) is used (TOPFLASH), together with a mock control plasmid (FOPFLASH). In the absence of hCx43, we found a significant increase in TCF/LEF luciferase activity when stimulated with WNT1 agonist CHIR (6-[2-[[4-(2,4-dichlorophenyl)-5-(5-methyl-1H-imidazol-2-yl)-2-pyrimidinyl]amino], ethyl[amino]-3-pyrindinecarbonitrile) at 1.5 μM (Figure 7e), compared with control EGFP transduced cells (shCx43 2.2 ± 0.7; EGFP 1.1 ± 0.3) and compared with hCx43-overexpressing cells (shCx43 2.2 ± 0.7; hCx43 0.6 ± 0.4; P ≤ 0.05). Interestingly, we also found a significant decrease
in TCF/LEF luciferase activity when hCx43 was overexpressed compared with EGFP control (hCx43 0.6 ± 0.4; EGFP 1.1 ± 0.3; P < 0.05). Thus, we can conclude that the knockdown of Cx43 protein increased β-catenin protein transcriptional activity, which enhanced neuronal differentiation.

**Discussion**

Cx proteins are differently regulated during neurogenesis. Cx43 is one of the major isoforms in the developing brain, being highly expressed in (E10.5) multipotent neural stem cells and in proliferating cells during...
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Figure 6 Quantification of Tuj1, GFAP and Ki67 expression in hNPCs expanded in the presence of CBX. Number of Tuj1 (a), GFAP (b) and Ki67 (c) expressing cells in 2 h differentiated cells. GJs were blocked by addition of CBX 50/μM. GZA was used as inactive analog of CBX. Values are the means of three independent experiments and error bars represent S.E.M. Values were expressed as % on total cell number.

Figure 6a: Screening of Tuj1 expression
Figure 6b: Screening of GFAP expression
Figure 6c: Screening of Ki67 expression

a

b

c

Development,27 furthermore, its levels decrease as NPCs differentiate into neurons,33 thus suggesting Cx43 has an important role in neurogenesis. This is a very dynamic process orchestrated by proliferation and differentiation of NPCs, which contribute to the development and maturation of a highly complex neural network. We used human NPCs isolated from the developing cortex as an in vitro model of human neurogenesis.34 hNPCs are EGF and FGF-2 responsive and can be expanded for >200 days in culture while retaining the ability to differentiate into neurons and glial cells.35 In this study, we confirmed that the expression of Cx43 was found in undifferentiated hNPCs but decreased as differentiation progressed out to 14 days. This is in keeping with what has been reported for the rodent brain.33 Following 7 days of differentiation, Cx43 was only present in GFAP-positive cells, indicating that when cells start to differentiate Cx43 expression is restricted to the glial phenotype. Interestingly, we found Cx43 punctate immunostaining at contact points between GFAP fibers and Tuj1-positive cells. This was in line with other rodent studies in which Cx43 was found to mediate tangential and radial migration of newborn neurons from the ventricular zone to the cortical plate.7,30,36 We designed and validated an shRNA construct to specifically knockdown hCx43 protein expression and then we generated lentiviral particles to deliver this construct into hNPCs. Cx43 knockdown resulted in a significant increase in the number of neurons derived from hNPCs both after 2 h and 7 days of differentiation. Interestingly, we also found a significant decrease in the number of GFAP-positive cells at the two time points examined. In vivo neurogenesis involves differentiation of neurons first, followed by glial cells,37 which suggests that in our system the increase in the number of neurons might reduce the pool of progenitor cells available to acquire an astrocytic phenotype at later time points. Alternatively as Cx43 in the mature brain is only expressed in astrocytes, it could be possible that the absence of the protein impaired or delayed astrocytic differentiation. Morphometric analysis revealed that the increase in neuronal phenotype when Cx43 was silenced did not correspond to an increase of neuronal complexity including cell body size, number and length of neurites. This suggests that this protein is not involved in the process of neuronal maturation, findings similar to what we recently reported for Cx36 in neurons derived from rat NPCs.38 In addition, we show that knockdown of Cx43 does not affect cell proliferation or cell death. Taken together, these results suggest that Cx43 knockdown increases neuronal commitment rather than influencing neuronal complexity, cell death or NPC proliferation.

Similarly, in vivo models of neurogenesis have shown that hCx43 is highly expressed at early stages of neurogenesis30,39 and Cx43 has been reported to mediate GJ coupling between neuroblasts in the hippocampus in mice.26 However, as progenitors start to differentiate, the degree of cellular communication together with Cx43 protein expression is decreased.29 The reduction in Cx43 expression and consequent disruption in neural progenitor coupling as differentiation occurred, suggests that Cx43 and GJ intracellular communication are important in the early stages of neurogenesis to maintain progenitors in an undifferentiated.29 Therefore, we investigated if the increased neuronal fate commitment we observed following hCx43 knockdown was due to a decrease in cellular communications or to the absence of the protein itself. Pharmacological blockade of GJICs with a generic GJ blocker (CBX) did not recapitulate the effects of Cx43 knockdown, suggesting a channel-independent role of Cx43 in the regulation of neuronal differentiation. Previous studies have established that the Cx43 adhesive, scaffolding and cell signaling-mediating properties are active factors during neurogenesis.7–9,36,40,41 Interestingly, Cx43-CT region contains multiple phosphorylation sites and binding domains, which not only modulate Cx43 protein functions but also allow Cx43 interaction with cytoskeletal elements and other cell membrane proteins including ZO1-2, adherens junction (AJ) proteins, cadherins and β-catenin (reviewed in Matsuuchi and Naus).28 β-Catenin signaling has a key role in neuronal differentiation,18 indeed, β-catenin through the TCF/LEF complex regulates the transcription of genes involved in cell adhesion, proliferation and differentiation such as E-cadherin, neuronal cell adhesion molecule, c-MYC and cyclin D1.42,43 Furthermore, Cx43 promoter contains three TCF/LEF binding sites.23 We stained Cx43 and β-catenin in hNPCs differentiated for 2 h and found that both proteins were abundantly expressed. We also detected areas in which the two proteins were colocalized, indicating a possible functional cross-regulation. Further experiments showed an increase in β-catenin protein expression following hCx43 knockdown and
more importantly no changes in β-catenin were observed using different shRNA constructs, confirming direct link between hCx43 knockdown and β-catenin protein expression. We also confirmed that the transcriptional active form of the β-catenin protein was enhanced in the absence of hCx43. In the cell, β-catenin is found at the cell membrane in complex with E-cadherin to form AJs. In the absence of WNT ligands, cytoplasmic β-catenin levels are kept low as the protein is phosphorylated by glycogen synthase kinase 3b (GSK-3b) and then targeted for ubiquitination and degradation. The activation of WNT pathway induces β-catenin dephosphorylation, nuclear translocation and the formation of a transcriptionally active complex with LEF and TCF sites. In cardiac cells, a complex of Cx43/β-catenin proteins have been described at the cell membrane following stimulation of WNT pathway.44 This evidence together with the presence of β-catenin binding sites in the Cx43 promoter region raised the possibility of a dual regulation between the two proteins. Cx43 may serve to negatively regulate its own transcription through β-catenin, as well as participate in the regulation of other β-catenin targets such as pro-neuronal genes18–20 by sequestering β-catenin to the cell membrane. Thus, in hNPCs, hCx43 knockdown disrupted the protein junctional complex with β-catenin leading to an increase in β-catenin nuclear translocation, transcription of pro-neuronal genes and an increase in the number of neurons. A dual regulation between gene transcription and cell adhesion molecules has already been described in a model proposed by Orsulic et al.,45 in which E-cadherin antagonized β-catenin signaling activity. In conclusion, in our investigation we have provided evidence for an interaction occurring between Cx43 and β-catenin proteins during neurogenesis. Furthermore, our results suggest a novel approach to increase the number of neurons from hNPCs, opening novel frontiers for research applications.

Materials and Methods
Design and construction of hCx43 shRNA lentiviral vector. The hCx43 sequence (accession no. NM.000165.3) was used to generate four hCx43 shRNAs (Supplementary Table 1). Sh expression cassette comprised a sense
and an antisense (antisense specific to the target mRNA) strand spaced by a loop sequence, a RNA polymerase III transcription termination signal and XhoI restriction site. A sequence coding for the mouse U6 promoter was inserted upstream of the antisense strand (Supplementary Figure S1A). A PCR-based method was used to generate double-stranded shRNA expression cassettes using the pSilencer 1.0-U6 expression vector (Life Technologies, Carlsbad, CA, USA) as a transcription template and reverse primer encoding the shRNA oligonucleotides. shRNA constructs were cloned into pCR2.1-TOPO vector as previously described26 and validated for specific hCx43 protein knockdown (Supplementary Figures S2A and B). Successful shRNA cassette was then excised by digestion with EcoRI and inserted into the lentiviral backbone plasmid via MfeI site generating the pRLSinscpt.U6.hCx43.shRNA.CMV.EGFP.wpre construct (Figure 2a).

**Construction of lentiviral vectors expressing hCx43, EGFP or SC.** The hCx43 coding sequence was excised from pDesH0.hCx43 plasmid (GlaxoSmithKline, Harlow, UK) using Ndel and AgeI restriction endonucleases and cloned into pRLSinscpt.CMV.polyA.B27EGFP.wpre construct via Ndel and AgeI sites generating pRLSinscpt.CMV.hCx43.IRES2.EGFP.wpre (Figure 2a). Lentiviral vector coding for EGFP under CMV promoter pRLSinscpt.CMV. EGFP.wpre and a sh SC sequence under a U6 promoter pRLSinscpt.U6.SC.CMV. EGFP.wpre were constructed as previously described.27

**Human neural stem cell cultures and lentivector vector transduction.** hNPCs were derived from 9 to 10 weeks old (post-menstrual) human fetus in line with the UK Department of Health guidelines and according to the Local Research Ethics Committee approval. hNPCs were expanded as neurospheres in presence of growth factors EGF and FGF-2 (20 ng/ml) and kept in the medium until the cells were harvested. The following day cells were co-transfected using Fugene 6 (Roche) with 0.99 μg of hCx43 shRNA constructs together with 0.2 μg of the CBX inactive analog DNA. Samples were cleaned up using the RNeasy minelute kit (Qiagen) and RNA quantity and quality was determined by spectrophotometry. RNA (1 μg) was retro-transcribed using the Superscript III system (Life Technologies) according to manufacturer’s instructions. PCR was carried out using the GoTaq system (Promega, Madison, WI, USA) according to manufacturer’s instructions. Primer sequences are indicated in Supplementary Table 3. PCR products were separated on 1.5% agarose gel containing 200 μg/ml ethidium bromide (Sigma), and visualized using a GelDoc-IT UV imaging system (UVF, Upland, CA, USA). PCR band intensity was analyzed by densitometry using ImageJ software and mRNA levels were normalized to the GAPDH mRNA. The mean of three different experiments was plotted and transcript level expressed as arbitrary units.

**Pharmaceutical blockade of GJ intercellular channels.** hNPCs were treated with 50 μM of CBX (Sigma) and 50 μM of the CBX inactive analog GZA (Sigma) for 10 days. CBX or GZA containing media was replaced every 3 days to avoid drug depletion. hNPCs were differentiated for 24 h.

**Luciferase reporter assay.** β-Catenin transcription was assessed using the TOP FLASH/FOP FLASH reporter luciferase assay.24 For TOP FLASH/FOP FLASH reporter luciferase assay, hNPCs were dissociated with Accutase (37 °C; Sigma) and plated onto poly-c-lysine-coated 24-well plates at 10^5 cells per well in differentiation medium. WNT pathway was stimulated adding 1.5 μM CHIR (WNT1 agonist, TOCRIS Biosciences, Minneapolis, MN, USA) at the time of plate down and kept in the medium until the cells were harvested. The following day cells were co-transfected using Fugene 6 (Roche) with 0.99 μg of β-catenin-LEF/TCF-sensitive (TOP) or β-catenin-LEF/TCF insensitive (FOP) reporter vector (Addgene) together with 0.1 μg of a constitutively active Renilla luciferase construct (Promega) for normalization of transfection efficiency. Cells were processed 48 h after co-transfection for Luciferase reporter activity using the Dual Luciferase Reporter System (Promega) according to the manufacturer’s instructions.

**Validation of hCx43 shRNA constructs.** Knockdown activity of hCx43 shRNA constructs was tested in HEK293 cells by transient transfection. Briefly, cells were plated at 7 x 10^4 cells per well and co-transfected with 0.8 μg of hCx43 shRNA constructs together with 0.2 μg of pDesH0.hCx43 transduced using Fugene 6 (Roche). Cells were harvested 48 h after transfection following and processed for western blot analysis.

**Image acquisition and data analysis.** Fluorescence microscopy was performed on 4% PFA fixed cells using Leica DM IRB inverted or LEICA DM RB upright microscopes (Leica, Wetzlar, Germany). Images were taken using ×20, ×40 and ×60 lenses. Neuronal cell body area and neurite branching were analyzed using the ImageJ plugin Simple neutile tracer (http://rsb.info.nih.gov/ij/). All experiments were performed at least three times on samples derived from three different fetuses. Values were expressed as mean ± S.E.M. P-values were calculated using Graphpad Prism 4 software (GraphPad, San Diego, CA, USA) by one-way ANOVA followed by Newman–Keuls post-hoc test. *P < 0.05, **P < 0.01.

**Conflict of Interest**

The authors declare no conflict of interest.

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Author contributions

Federica Rinaldi: conception and design of the study, performed the experiments and wrote the paper. Elizabeth M Hartfield: collection and assembly of data, final approval of the manuscript. Lucy A Crompton: data analysis and interpretation, final approval of the manuscript. Jennifer L Badger: provision of study material, final approval of the manuscript. Colin P Glover: provision of study material, final approval of the manuscript. Claire M Kelly: provision of study material, final approval of the manuscript. Anne E Rosser: provision of study material, final approval of the manuscript. James B Uney: conception and design, financial support, data analysis and interpretation, final approval of the manuscript. Maae A Caldwell: conception and design, financial support, data analysis and interpretation, manuscript writing, final approval of the manuscript.

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