Wnt5a Regulates Midbrain Dopaminergic Axon Growth and Guidance

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Abstract

During development, precise temporal and spatial gradients are responsible for guiding axons to their appropriate targets. Within the developing ventral midbrain (VM) the cues that guide dopaminergic (DA) axons to their forebrain targets remain to be fully elucidated. Wnts are morphogens that have been identified as axon guidance molecules. Several Wnts are expressed in the VM where they regulate the birth of DA neurons. Here, we describe that a precise transitory-spatial expression of Wnt5a accompanies the development of nigrostriatal projections by VM DA neurons. In mice at E11.5, Wnt5a is expressed in the VM where it was found to promote DA neurite and axonal growth in VM primary cultures. By E14.5, when DA axons are approaching their striatal target, Wnt5a causes DA neurite retraction in primary cultures. Co-culture of VM explants with Wnt5a-overexpressing cell aggregates revealed that Wnt5a is capable of repelling DA neurites. Antagonism experiments revealed that the effects of Wnt5a are mediated by the Frizzled receptors and by the small GTPase, Rac1 (a component of the non-canonical Wnt planar cell polarity pathway). Moreover, the effects were specific as they could be blocked by Wnt5a antibody, sFRPs and RYK-Fc. The importance of Wnt5a in DA axon morphogenesis was further verified in Wnt5a/−/− mice, where fasciculation of the medial forebrain bundle (MFB) as well as the density of DA neurites in the MFB and striatal terminals were disrupted. Thus, our results identify a novel role of Wnt5a in DA axon growth and guidance.

Introduction

Dopamine (DA) neurons within the ventral midbrain (VM) project to the striatum and prefrontal cortex forming the nigrostriatal, mesocortical and mesolimbic pathways, which are important for motor and cognitive functions. DA neuron dysfunction is associated with a number of neurological and psychiatric disorders. Abnormal development of the nervous system may contribute to these disorders; hence, the importance of understanding the processes involved in DA neuron maturation and connectivity. Whilst the cues that orchestrate the birth of midbrain DA neurons are well established, the signals regulating DA neurite morphogenesis (including neurite growth, axon guidance and synaptogenesis) are less well defined.

Several studies have identified cellular and molecular signals that participate in establishing these pathways (see review by [1]), including Ephrins [2–4], Semaphorins [5–9], Netrins and Slits [10,11], Engrailed-1 [12,13], and Sonic hedgehog [14]. In this study we asked whether Wnts also regulate DA axon morphogenesis.

Wnt1 and Wnt5a are important morphogens for VM development, regulating proliferation, differentiation and survival of DA neurons [15–24]. Wnts also participate in axon guidance elsewhere in the central nervous system [25–30]. Specifically, Wnt5a repels corticospinal axons [31–33], commissural axons [34] and cortical axons in the corpus callosum [35,36], and promotes neurite elongation of cortical neurons [33].

Wnt5a is a highly conserved diffusible protein whose signal is transduced by Frizzled (Fz) receptors and/or co-receptors including the atypical tyrosine kinases Ryk and Ror2. Dependent on the receptor and cell type, Wnt5a has been shown to activate three signaling pathways: the Wnt/β-catenin/canonical pathway, the Wnt/calcium/non-canonical pathway, and the Wnt/planar cell polarity (PCP)/non-canonical pathway [37–39]. However, little is known about which of these pathways and downstream signaling components mediate Wnt5a’s influence on axon growth and guidance. Moreover, it is not known whether Wnt5a promotes neuritogenesis and axonal growth of DA axons in the nigrostriatal system. We therefore set out to determine whether Wnt5a plays a role in DA axon growth and guidance and examined the
involvement of some of the candidate Wnt5a receptors and Wnt signaling components.

**Materials and Methods**

**Animals**

**Ethics statement.** This study conformed to the Australian National Health and Medical Research Council’s published Code of Practice for the Use of Animals in Research, and experiments were approved by the Florey Neuroscience Institutes animal ethics committee (#07-040).

Embryos were isolated from timed-mated C57BL/6 mice or Sprague Dawley rats. Animals were time mated overnight and visualization of a vaginal plug on the following morning was taken as embryonic day (E) 0.5. B6;129S7-Wnt5a<sup>tm1.8m</sup>/J (subsequently referred to as Wnt5a<sup>+/−</sup> mice) were obtained from Jackson Laboratories (JAX, Strain 004758) and maintained on a mixed B6:129 background [40]. Wnt5a embryos were collected at E12 and E18.

**In situ hybridization and immunohistochemistry**

Embryos were isolated in ice-cold PBS, fixed overnight in 4% paraformaldehyde, followed by overnight immersion in 30% sucrose in PBS. Embryonic day 11.5 (E11.5), E12 and E14.5 embryos were cryosectioned on either a sagittal or coronal plane at a thickness of 14 μm. E18 embryos were cryosectioned at 16 μm. In situ hybridization (ISH) was performed as previously described [41], using a DIG-labelled single-stranded RNA probe for Wnt5a [40]. Following ISH, the tissue was again fixed using 4% paraformaldehyde, followed by overnight immersion in 30% sucrose in PBS. Embryonic day 11.5 (E11.5), E12 and E14.5 embryos were cryosectioned on either a sagittal or coronal plane at a thickness of 14 μm. E18 embryos were cryosectioned at 16 μm. Wnt5a embryos were collected at E12 and E18.

**Quantitative real-time PCR**

Given the lack of reliable antibodies to detect many of the Wnt ligands and receptors histochemically, we relied on quantitative real-time PCR (Q-PCR) to assess the expression of Wnt5a, Ryk and Fz3 within VM, and more specifically within DA neurons. Ventral midbrains were isolated and dissociated from E11.5 Tyrosine Hydroxylase-GFP (TH-GFP) reporter mice, in which all DA neurons express GFP [42]. Dissections are described in further detail below. At least five TH-GFP<sup>+</sup> embryos were used for each dissection with four independent dissections performed. Using previously described methods [43], fluorescence-activated cell sorting (FACS) was used to separate GFP<sup>+</sup> cells (dopamine neurons) from GFP<sup>−</sup> cells (non-TH<sup>+</sup> neurons within the VM) in order to identify the source of Wnt5a, Ryk and Fz3 in the midbrain. Following sorting, total RNA was isolated using the PicoPure kit (Arcturus). Alternatively, the ventral midbrain (VM), dorsal midbrain (DM) and the rest of the embryo (E) were microdissected from four independent E11.5 mouse litters. Following tissue isolation, total RNA was isolated using the RNeasy Micro kit (Qiagen).

RNA was reverse transcribed using Superscript III First-Strand Synthesis supermix for qRT-PCR (Invitrogen) and Q-PCR was carried out using the SYBR GreenE<sup>TM</sup> qPCR SuperMix Universal (Invitrogen) on an ABI7700 sequence detection system (Applied Biosystems, Foster City, CA) using the comparative ΔCT method [44]. Oligonucleotide sequences were as follows: HPRT forward, 5′- TCTTGTGACCTGCTGGATT-3′; HPRT reverse, 5′- TAGTCGCCCCGTGGACTGAT-3′; Wnt5a forward, 5′- AATAACCTGTTACAGATGTCA-3′; Wnt5a reverse, 5′- TACTCTAGTGGTCTCTGATA-3′; Ryk forward, 5′- CGGTCGTGCTTAAACCTGG-3′; Ryk reverse, 5′- CGGATCAATCTTTTTAATC-3′; Fz3 forward, 5′- CAGTCCTGCAATGGAGTGG-3′; Fz3 reverse, 5′- CGGACTAATATTGTCACCT-3′.

**Ventral Midbrain Primary Cultures**

The ventral midbrain of E11.5 and E14.5 mouse (or E13.5 rat) embryos was microdissected in chilled L15 media (Invitrogen). Note, stages in development of the dopamine systems occur approximately 2 days later in rats than mice, hence E13.5 rat is considered equivalent to E11.5 mouse. Whilst initial studies were performed in mice, they were verified later in rats. Rat embryos were used in all antagonism studies as greater volumes of VM primary neurons can be obtained, necessary for the outlined antagonism studies that required multiple conditions. The isolated ventral midbrains were enzymatically dissociated in HBSS containing 0.05% trypsin and 0.1% DNase for 12 minutes at 37 °C. Cells were subsequently centrifuged and resuspended in serum-free N2 medium consisting of a 1:1 mixture of F12 and MEM supplemented with 15 mM HEPES buffer, 1 mM gluta mine, 6 mg/ml glucose (Sigma-Aldrich), 1 mg/ml bovine serum albumin and N2 supplement (all purchased from Invitrogen). Cells were seeded at a density of 125,000 cells per well in a 48-well plate at 37 °C, 5% CO<sub>2</sub> for 72 hours.

Wnt5a recombinant protein (R&D Systems) was added to the cultures at the time of cell seeding. For antagonism experiments using secreted frizzled-related protein 1 (sFRP1; 5 μg/ml, R&D Systems), Wnt5a antibody (αWnt5a; 2 μg/ml, R&D Systems), human RYK-Fc (3 μg/ml, see details below), goat anti-Frizzled3-CRD (αFz3-CRD; 3 μg/ml, R&D Systems), Dickkopf-1 (Dkk1; 500 ng/ml, R&D Systems), casein kinase 1 inhibitor (D4476, 500 ng/ml, Roche) or Rac-1 inhibitor (NC23976, 500 nM, Calbiochem), Wnt5a and the antagonist were added to the wells 15 minutes prior to seeding the VM cells.

To generate the RYK-Fc, the human RYK WIF domain (residues 60–195 of Genbank accession number NP_002949.2) was cloned by PCR into pApex-3.Fc.FLAG, between an IL-3 signal peptide and the human IgG<sub>1</sub> Fc domain, to create a fusion protein with a carboxyl-terminal FLAG epitope tag. CHO-K1 cells were transfected with pApex-3.hrYKWD.Fc.FLAG using FuGENE 6 (Invitrogen) and selection applied after 24 h (200 μg/ml hygromycin B; Invitrogen). Stable colonies were picked after 7–9 days. The stable cell line hRYKWD.Fc.FLAG/CHO was seeded into a medium FiberCell cartridge, 20 kDa (FiberCell Systems), using DMEM (Invitrogen), 10% fetal bovine serum (FBS) and 100 μg/ml hygromycin B. Extracapillary space media from the FiberCell cartridge was collected every 2–3 d, filtered using 0.22 μm filters (Millipore), and secreted protein was purified using anti-FLAG M2 affinity gel (Sigma) as previously described [45].

TH-immunoreactive (TH<sup>+</sup>) neurons from each primary VM culture were analyzed from 3–5 independent cultures. Under all culture conditions, sampling was commenced in the second field of view from the left-hand side of the culture well. The first 30 TH<sup>+</sup> cells found to be measurable (neurites intact and distinguishable from other stained neurites, i.e. not intertwined with other TH<sup>+</sup> neurons) from GFP<sup>+</sup> cells (non-TH<sup>+</sup> neurons within the VM) in order to identify the source of Wnt5a, Ryk and Fz3 in the midbrain. Following sorting, total RNA was isolated using the PicoPure kit (Arcturus). Alternatively, the ventral midbrain (VM), dorsal midbrain (DM) and the rest of the embryo (E) were microdissected from four independent E11.5 mouse litters. Following tissue isolation, total RNA was isolated using the RNeasy Micro kit (Qiagen).

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neurites) were quantified in order to avoid any potential sampling bias. In each experiment, data was compared to the mean normalized control value (set at 100%) to account for inter-experimental variation. Photomicrographs of each DA neuron (identified by TH+) were taken using a 20x objective (Olympus IX71) and the following measurements obtained using NeuronJ software [ImageJ, NeuronJ plugin, NIH]: the total numbers of neurites per DA neuron, the number of neurite branches, total length of all neurites per neuron and the length of the dominant neurite (the longest, most dominant neurite arising from the soma, and thereby presumably the axon [46]).

VM primary cultures were also performed from Wnt5a+/− and littermate Wnt5a−/+ and Wnt5a+/− mice. Given that single VM were required for each culture, and the low yield of neurons generated, each dissected VM was dissociated as described above and plated into a 96-well plate (50,000 cells/well). Cells were cultured for three divisions (3DIV) prior to staining and measurements of TH+ neurons. Wnt5a−/+ cultures were compared to Wnt5a−/+ and Wnt5a+/− littermates.

Immunoblotting
SN4741 cells were cultured in DMEM, 10% FBS, L-glutamine (2 mM), penicillin/streptomycin (50 U/ml) and glucose (0.6%). For analysis of intracellular Wnt signaling, 100,000 cells were seeded in 12-well plates, grown overnight in the absence of serum and stimulated for 2 hours in the same media with Wnt5a (0, 30, 100, 300, 1000 ng/ml; R&D Systems), Wnt5a (300 ng/ml) + RYK-Fc (3 μg/ml) or Wnt5a (300 ng/ml) + Fz3-CRD (3 μg/ml; R&D Systems). Preparation of lysates and immunoblotting were carried out as previously described [47]. The following primary antibodies were used: rabbit anti-Dvl2 (1:500, Santa Cruz) and mouse anti-β-actin (1:3000, Sigma).

Co-culture explant assays
Neuronal c17.2 cells over-expressing Wnt5a (or the parental cell line containing the empty vector i.e. mock) were cultured as previously described [48]. Aggregates of the c17.2 cells were generated by plating 50,000 cells per 20 μl droplet onto the inverted lid of a 60 mm culture plate containing 1 ml PBS (to maintain humidity). Cell aggregates formed within 48 hours and were floated in N2 media prior to co-culture with VM explants. The ventral midbrain of E11.5 or E14.5 mice was isolated in L15 media. Cell aggregates formed within 48 hours and were floated in N2 media prior to co-culture with VM explants. The number of TH+ fibers in the striatum. The MFB was delineated in approximately 7 sections (16 μm thickness, 1:10 series, i.e. approximately 1120 μm in length) from each brain, with the area and total volume estimated using StereoInvestigator software. The number of dopaminergic fibers (TH+) in the MFB and the density of TH+ varicosities in the lateral striatum were estimated using previously described fractionator methods [49–51]. The density of TH+ fibers was assessed at two independent levels along the MFB, (i) 320 μm and (ii) 800 μm rostral to the midbrain DA neurons. TH+ fiber counts were made at regular pre-determined intervals (x = 50 μm, y = 50 μm). These counts were derived by means of a grid program, through which a systematic sample of the area occupied by the fibers was made from a random starting point. An unbiased counting frame of known area (7 μm × 7 μm = 49 μm²) was superimposed on the image of the tissue sections viewed under a 100×, N.A. 1.30 oil immersion objective. The number of TH+ fibers at each level was counted in 16 μm thick, 1:10 serial sections.

TH+ varicosities in the lateral 400 μm of the striatum were counted from 16 μm serial sections, 1:10 series, with four sections sampled from each striatum. Counts of TH+ varicosities were made at regular predetermined intervals (x = 150 μm, y = 150 μm) using an unbiased counting frame of known area (6 μm × 6 μm = 36 μm²). TH+ varicosities were identified as predominantly round swellings in association with axonal processes. TH+ varicosity counts were expressed as terminal density, with comparisons made between Wnt5a+/−, Wnt5a−/+ and Wnt5a+/− mice. For TH+ fiber counts and TH+ varicosity numbers, the coefficients of error (CE) and coefficients of variance (CV) were calculated as estimates of precision, and values of less than 0.1 were accepted [50,52,53].

Statistical analysis
One-way ANOVAs with Tukey post-hoc tests or Student’s t-tests were used to identify statistically significant changes. Statistical significance was set at a level of p<0.05. Data represents mean ± s.e.m.

Results
Wnt5a is expressed along the developing nigrostriatal pathway
The first ventral midbrain DA neurons are born in mice at E10.5 and shortly thereafter, at E11.5, the first DA neurites appear. Initially, these neurites project dorsally towards the dorsal
midbrain (Fig. 1D) and are subsequently deflected rostrally towards their forebrain targets [54]. The first DA neurites approach the border of the ventrolateral ganglionic eminence at E14.5 (Fig. 1K) and increase in number without further elongation (axonal stalling). Subsequently these fibers enter the ventral areas of the lateral ganglionic eminence (LGE; the future striatum), followed by lateral, and finally medial and dorsal regions of the LGE [1]. A smaller subset of DA axons arising from the VM...
Wnt5a regulates dopaminergic axon morphogenesis

Wnt5a mRNA expression was more closely examined in VM cells isolated by FACS from the TH-GFP reporter mouse [12]. Quantitative real-time PCR (Q-PCR) performed on the GFP+ fraction (DA neurons) and GFP- fraction (other VM cells) revealed that Wnt5a mRNA expression was significantly higher in the GFP+ fraction (four-fold increase, p = 0.042) compared to the GFP- fraction (Fig. 1O). These findings were in accordance with Wnt5a in situ hybridization, with expression greatest in non-TH+ cells (Fig. 1N’, filled arrow-head). These results are also in agreement with previous studies showing greater expression of Wnt5a in glial cells (radial glia first and later astrocytes) compared to neurons in the developing VM [22,56].

Wnt5a increases dominant DA neurite length and reduces DA neurite branching in VM cultures at E11.5, but not at E14.5

As the temporal-spatial pattern of expression of Wnt5a was appropriate for a role in DA neurite development, we tested the effect of Wnt5a on DA neurite growth by applying recombinant Wnt5a protein to VM primary neuron cultures isolated from E11.5 and E14.5 mouse embryos and examining the neurites of tyrosine hydroxylase (TH; rate-limiting enzyme in DA synthesis and marker of DA neurons) immunoreactive neurons. A dose-response curve revealed that Wnt5a promotes DA neurite elongation in a dose-dependent manner in E11.5 VM cultures. Maximal elongation, as measured by total neurite length, was achieved with a dose of 300 ng/mL of Wnt5a (Fig. 2A). Immunoblots were performed in a dopaminergic cell line (SN44741) in order to confirm that Wnt5a induced intracellular activation of Wnt signaling. We found that 100, 300 and 1000 ng/mL of Wnt5a induced dishevelled-2 (Dvl2) phosphorylation (visible by Western blot as a mobility shift of the protein) in a dose-dependent manner (Fig. 2A). The effects were optimal at 300 ng/mL and this dose was used for further assessment of the neurite arbors of TH+ cells. Wnt5a treatment of E11.5 VM cultures increased total neurite length compared to control treated cultures (295%±12%, p<0.001; Fig. 2B-D). Other morphological changes were also observed in Wnt5a treated cultures. The dominant neurite was significantly longer compared to controls (330%±17%, p<0.001; Fig. 2E).

Furthermore, Wnt5a treatment resulted in fewer neurites (80%±2%, p<0.001; Fig. 2B,C,F) and branches (53%±29%, p = 0.012; Fig. 2B,C,G) compared to controls, suggesting that Wnt5a promotes the extension of DA axons, rather than the elaboration of shorter neurites or dendritic trees [46]. These findings were also replicated in E13.5 rat cultures (comparable in age to mouse E11.5), demonstrating conservation of the Wnt5a effect across species (data not shown).

We next examined the specificity of the effects of Wnt5a by examining the neurite length of BIII-tubulin immunoreactive (TUJ1+) neurons within the culture, knowing that TH+ cells represent approximately only 5% of the neurons in the VM culture. The total length of TUJ1-labeled neurites in cultures treated with Wnt5a were not significantly longer than neurites in control cultures (117%±14%, and 100%±10%, respectively, p = 0.309; Fig. 2H-J), confirming that Wnt5a selectively affected DA neurites.

Surprisingly, when the activity of Wnt5a (300 ng/mL) was examined on older (E14.5) VM primary cultures, the effects on DA neurite length were reversed. Total neurite length was significantly reduced (69%±4%, Fig. 3A, E-F) and the length of the dominant process (axon) was also decreased compared to control-treated cultures (65%±5%; Fig. 3B). Furthermore, Wnt5a treatment affected neither the number of DA neurites nor their branching (Fig. 3C,D). Collectively, these results indicate that Wnt5a differentially regulates DA neurite growth and morphology during development.

The effects of Wnt5a protein on DA neuritogenesis are specific and mediated by Frizzled

To confirm the specificity of the effects of Wnt5a on DA neurite development, we treated primary VM cultures with different Wnt blocking tools and subsequently evaluated TH+ neurites. Given the maintained effect of Wnt5a on DA neurites in both mice and rats, we performed these antagonism experiments in rats due to the increased yield of VM tissue, and the numerous antagonists to be employed. Figure S1 provides a schematic representation of the site of action of these various antagonists. Secreted Frizzled-related proteins (sFRPs) modulate Wnt signaling by preventing Wnt from interacting with membrane-bound receptors. In the absence of exogenous Wnt5a, sFRP-1 reduced neurite length to 62%±4% compared to untreated cultures (Fig. 4A-C), presumably through antagonism of endogenous Wnt signaling within the VM. This was confirmed by using a Wnt5a blocking antibody (αWnt5a), which also reduced neurite length to 67%±8% (Fig. 4A-D). In the presence of Wnt5a, increased neurite length of TH+ cells (267%±31%) was completely blocked by co-administration of sFRP-1 (101%±10%) or the αWnt5a (70%±11%; Fig. 4A,E-G). Interestingly, sFRP-2, but not sFRP-3 (data not shown), also antagonized the effects of Wnt5a on neuritogenesis. These results indicated that the effects of exogenously supplied Wnt5a on TH+ cells are specific and suggest a role for Wnt5a in DA neuritogenesis.

Previous studies have shown that Wnt5a modulates axon growth and guidance in other systems through interactions with Frizzled receptors and the atypical tyrosine kinase receptor, Ryk [31,33,35]. Moreover, Fz3 has been found to be relevant to the development of the DA nigrostriatal pathway as Fz3 expression increases at the time of DA axon extension in the VM [43] and the nigrostriatal pathway was absent in Fz3−/− mice [57,58]. We thus first examined the expression of Fz3 and Ryk in the VM by Q-PCR and found elevated expression of both receptors in the VM compared to the dorsal midbrain (DM) and the rest of the embryonic E1 (Fig. 4H,I,G). Further, Q-PCR performed on the GFP+ and GFP− fraction of VM tissue isolated from TH-GFP mice,
revealed that these receptors were expressed on the DA neurons (GFP\(^+\)) and not surrounding cells (GFP\(^-\)) within the VM (Fig. 4H'' and 4I'').

Subsequently, we examined whether E13.5 rat VM cultures treated with antibodies against Fz3 (\(\alpha Fz3\)-CRD) or in the presence of a Ryk construct containing the human RYK WIF domain (RYK-Fc), blocked the effects of Wnt5a (Fig. 4K–Q). Interestingly, the increase in neurite length produced by the addition of Wnt5a (166\(\%\) ± 10\(\%\)) was significantly attenuated in the presence of a RYK-Fc, to levels not significantly different to control (97\(\%\) ± 4\(\%\)), yet they had no effect on neurite number (data not shown). Similarly, but more modestly, \(\alpha Fz3\)-CRD also reduced the effect of Wnt5a on total neurite length (from 166\(\%\) ± 10\(\%\) to 129\(\%\) ± 6\(\%\)), but not neurite number (data not shown). Since RYK-Fc can bind Wnt, we interpret that the blocking by RYK-Fc likely mediated by Wnt5a binding. However, as the RYK-Fc construct is not capable of interacting directly at the receptor membrane level, this data should be interpreted with caution. In

Figure 2. Wnt5a promotes DA axon elongation and alters neuron complexity during the period of initiation of neurite outgrowth. (A) Wnt5a recombinant protein promoted TH\(^+\) neurite elongation in a dose-responsive manner in mouse E11.5 VM primary cultures. (A') Wnt5a activated Dvl2 in a dose-responsive manner in the SN4741 dopaminergic cell line. Note the mobility shift of the Dvl2 protein with increasing doses of Wnt5a. (B) Photomicrographs illustrating the complexity of DA neurons under control conditions, and (C) following Wnt5a treatment. (D) Wnt5a induced a three-fold increase in total neurite length compared to control. (E) The effect of Wnt5a was specific to the dominant neurite (presumably the DA axon). Wnt5a protein reduced the number of (F) DA neurites and (G) DA neuritic branches per neuron. (H) Immunocytochemistry for TUJ1 revealed that the effects of Wnt5a within the VM were specific to DA neurons, with no change in neurite length observed for other neurons in culture. (I) Compared to control cultures, (J) Wnt5a had no effect on neurite length of TUJ-labeled cells. Cells were analyzed after 3DIV. Scale bar = 25 \(\mu\)m.

Data represents mean ± s.e.m., \(n=4\)--5 cultures; * \(p<0.05\), ** \(p<0.01\), *** \(p<0.001\).

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fact, a recent study has shown no changes in the morphology or trajectory of DA axons in Ryk(-/-) embryos [55]. We believe that additional experiments, employing selective functional blocking antibodies against the Ryk receptor, as well as Ryk-Wnt5a double knock out mice will be required to ascertain whether Ryk plays a role in Wnt5a mediated DA axon growth and guidance.

In contrast, since the αFz3-CRD binds directly to the receptor (with its specificity and function verified by the manufacturer and others Endo 2008), our results indicate that the action of Wnt5a on DA neurite morphology is mediated, at least in part, by the Frizzled-3 receptor. Interestingly, we also found that these proteins antagonized the Wnt5a-mediated Dvl-2 phosphorylation in a dopaminergic cell line (SN4741 cells, Fig. 4J), reinforcing the idea that the effects of Wnt5a require Wnt5a binding and are mediated by Frizzled. It is also important to note that RYK-Fc and αFz3-CRD had no effect on non-DA neurites within the culture, illustrating the specificity of the effects of Wnt5a for DA neurites and the lack of toxicity of the proteins used here (Figure S2).

**Wnt5a regulates neurite morphogenesis in midbrain DA neurons via Rac1**

Depending on the cell type and context, Wnt5a activates either Wnt/β-catenin or Wnt/PCP signaling. However, we previously reported that in dopaminergic cell lines and expanded VM cultures [19,20] Wnt5a does not activate Wnt/β-catenin signaling. Moreover, Wnt5a mediates axon guidance through non-canonical Wnt pathways in other neuronal systems [33,59]. To characterize the pathway that mediates the effects of Wnt5a on DA neuritogenesis, we employed an antagonist of the canonical pathway, Dickkopf-1 (Dkk1), to prevent Wnt interaction with the Fz/LRP receptor complex [60], and a casein kinase 1 antagonist, D4476, which blocks both the canonical and non-canonical pathways [61]. While Dkk1 had no effect on Wnt5a-mediated neurite length of TH+ cells in E13.5 rat VM cultures (Fig. 5A,C,D), D4476 significantly inhibited the effects of Wnt5a on neurite length (Fig. 5A,C,E). Importantly, D4476 and Dkk1 had no effect on neurite length of TUJ+ neurons (data not shown), verifying the lack of toxicity of the antagonists. These results suggested that the action of Wnt5a on DA neurite length was mediated by non-canonical Wnt signaling.

We next examined whether blocking of the small GTPase, Rac1, a downstream component of the Wnt/PCP pathway in DA neurons [23] affected the total length of DA neurites in primary VM cultures. Treatment of control cultures with the Rac1 inhibitor, NSC23376 reduced TH+ neurite length (39%±4% reduction), presumably due to antagonism of endogenous Wnt signaling as well as other potential axon guidance pathways for which Rac1 is a downstream component. However, co-treatment

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**Figure 3. Wnt5a causes DA neurite retraction in older ventral midbrain cultures.** (A) At a time when DA axons would normally be approaching their striatal targets (mouse E14.5), treatment with Wnt5a protein caused retraction of TH+ neurites, and more specifically (B) DA axons (dominant neurite length). Wnt5a had no effect on the complexity of DA neurons, as assessed by (C) neurite number and (D) neurite branching. Photomicrographs illustrating examples of neurite retraction following Wnt5a application (F), compared to control (E). Cells were analyzed after 3DIV. Scale bar = 25 μm. Data represents mean ± s.e.m., n = 4–5 cultures, **p<0.001. doi:10.1371/journal.pone.0018373.g003
Figure 4. The effects of Wnt5a protein on DA neuritogenesis are specific and mediated by Frizzled. (A) Whilst sFRP-1 and Wnt5a blocking antibody (αWnt5a) were capable of reducing neurite length in E13.5 VM rat primary cultures under control conditions (presumably due to antagonism of endogenous Wnt signaling), in the presence of Wnt5a they significantly reduced neurite length compared to Wnt5a alone. (B–G)
Examples of Wnt antagonism in VM cultures ± Wnt5a treatment. Q-PCR analysis revealed that the Wnt-related receptors Fz3 (H) and Ryk (I) were highly expressed in the VM compared to the dorsal midbrain (DM) and whole embryo (E). Furthermore, Fz3 (H*) and Ryk (I*) showed significantly higher expression within DA neurons (GFP) compared to other cells (GFP) isolated from the VM of TH-GFP mice. (J) Wnt5a (300 ng/ml) activated Dvl2 in SN4741 cells, an effect that could be blocked by Fz3-CRD (3 μg/ml) or Ryk-Fc (3 μg/ml). (K) E13.5 rat primary VM cultures showed that the effects of Wnt5a on DA neurite length were specific, illustrated by antagonism with Ryk-Fc, and mediated through the Fz3 receptors, illustrated by blocking of Fz3 with Fz3-CRD. (L-Q) Photomicrographs illustrating the effects of Wnt5a ± Fz3-CRD or Ryk-Fc on DA neurite length. Cells were cultured for 3DIV. Scale bar = 100 μm. Data represents mean ± s.e.m., n = 4–5 cultures. ** p < 0.05, *** p < 0.001. Significantly different from Wnt5a: * p < 0.05. doi:10.1371/journal.pone.0018373.g004

With Wnt5a and NSC23766, significantly reduced total TH+ neurite length even further (60% ± 4%; Fig. 5A,C,F) compared to Wnt5a treatment alone. These findings suggest that at least in part Rac1, downstream of Wnt5a and thereby the PCP pathway, regulates DA neurite length. Finally, treatment with Wnt5a + D4476, but not Wnt5a + NSC23766, increased the number of TH+ neurites per explant (Fig. 5B), suggesting that while PCP signaling increases neurite length, other non-canonical Wnt signaling such as the Ca2+ pathway may cooperate to reduce the number of neurites.

Wnt5a repels DA neurites in E11.5 and E14.5 explant cultures

After characterizing the effects of Wnt5a on neuritogenesis, we examined the chemorepulsive or chemorepellent effect of Wnt5a on developing DA neurites in VM explants. In other CNS regions, Wnt5a repels neurites during development [31,33,35], but its effect on DA neurons has not yet been examined. Mouse VM explants (E11.5) were co-cultured with either mock-transfected or Wnt5a over-expressing cell aggregates for 72 hours and the number of TH+ fibers in the distal (D) and proximal (P) quadrants of the explant, with respect to the cell aggregate, were counted, as depicted in Fig. 6A. TH+ neurites from explants co-cultured with mock-transfected cell aggregates radiated out in all directions from the explant (Fig. 6D), showing a proximal to distal ratio of neurites close to 1 (1.19 ± 0.06, n = 30; Fig. 6B). However, most TH+ neurites in VM explants cultured with Wnt5a-overexpressing cell aggregates emanated from the distal aspect of the explant, with a proximal to distal ratio of 0.70 ± 0.09 (n = 29, p < 0.001; Fig. 6B,E), indicating a repulsive effect. We confirmed that the effects of Wnt5a were specific to TH+ neurites by staining all neurites in culture (TUJ1+). Whilst TH+ fibers were repelled by Wnt5a (Fig. 6F), TUJ1+ neurites were observed emanating from all aspects of the same explants (Fig. 6F). This effect of Wnt5a on DA neurites was ablated by bath application of the casein kinase 1 inhibitor D4476, the Rac1 inhibitor NSC23766, and anti-Fz3-CRD, indicating that the repulsive effects of Wnt5a on TH+ neurites are mediated by PCP/Wnt signaling via the Fz3 receptor (Fig. 6B,G–I). These findings have recently been supported by Fenstermaker et al (2010), who illustrated that DA neurons in Fz3(−/−) and Celsr3 (−/−) mice (Celsr3 being an additional component of the PCP pathway) were non-responsive to Wnt5a [55]. Given the contrasting effect of Wnt5a on neurite extension at differing developmental ages (E11.5 and E14.5, Fig. 2 and Fig. 3, respectively), we asked whether the chemorepulsion effect of Wnt5a on DA neurites was maintained in older VM explants (E14.5) and found that this was the case (Fig. 6C). We next investigated whether Wnt5a also regulates the development of midbrain DA axons in vivo and therefore examined the Wnt5a−/− mouse.

Deletion of Wnt5a increases the number of TH+ fibers in the medial forebrain bundle and the innervation of the striatum in vivo

The contribution of Wnt5a to DA axon growth and guidance was further examined by inspecting the trajectory of TH+ axons in the Wnt5a−/− mouse at E18. Gross examination of the pathway highlighted a broadening of the MFB and elaborated innervation in the dorsal striatum of Wnt5a−/− mice, not seen in Wnt5a+/+ mice (Fig. 7A,B). Stereological quantification of the fiber bundle...
volume revealed that the Wnt5a^-/- mice possessed a significantly enlarged MFB (9.708±0.6107 vs. 7.081±0.4207) compared to heterozygotes (7.081±0.1403) and wildtype (5.770±0.1007) littermates (Fig. 7C). Furthermore, there were more fibers within the caudal MFB (320 mm rostral to the TH+ cells in the VM) of Wnt5a^-/- mice than in Wnt5a^+/+ mice (11,487±632 and 8296±549 TH+ fibers, respectively; Fig. 7D-G'). The increase in TH+ fiber number was maintained within the MFB, with more rostral aspects of the bundle (800 mm rostral to the TH+ cells in the VM) also showing significant increases in fibers (data not shown).

We next used the density of TH+ synaptic varicosities in the striatum as a measure of the capacity of DA fibers to innervate their target structures. The lateral striatum of mutant and wildtype E18 embryos, which is innervated by both substantia nigra pars compacta (SNpc) and ventral tegmental area (VTA) DA neurons [62,63] was examined. The terminal density in the lateral striatum of Wnt5a^-/- mice (8.10±0.6102 terminals/um^3) was significantly greater than in Wnt5a^+/+ mice (6.11±0.1303 terminals/um^3, Fig. 7E-H,J) and innervation of the dorsal striatum in Wnt5a^-/- mice (Fig. 7B,B') was notably denser than in wildtype littersmates (Fig. 7A,A'). These results suggest that Wnt5a is required for the correct distribution of TH+ fibers in the dorso-lateral striatum, the target area of substantia nigra DA neurons.

Finally, in light of observed DA axon defects in E18 Wnt5a^-/- embryos, and our expression gradients identified in Figure 1, we examined an earlier time point (E12) to determine the effect of Wnt5a ablation on the establishment of the DA pathways. We observed that axons were notably shorter in Wnt5a^-/- embryos (Figure S3A' and 3B', arrow heads), confirming the importance of Wnt5a in initial neurite elongation. Furthermore, broadening of

Figure 6. Wnt5a acts as a chemorepellant for DA neurites in VM explants. (A) Schematic representation of the explant cultures. VM explants were plated in collagen adjacent to mock- or Wnt5a-transfected cell aggregates. After three days in culture, the number of DA neurites (TH+) radiating from the proximal and distal sides of the VM explant (relative to the aggregate) were counted and expressed as a ratio. (B) In co-cultures of E11.5 mouse VM explants with Wnt5a-transfected cells, the majority of TH+ neurites emanated from the distal side of the explant, an effect that could be ablated by bath application of casein kinase 1 inhibitor D4476, Rac1 inhibitor NSC23766 and anti-Fz3-CRD. (C) The ability of Wnt5a to induce repulsion of DA neurites was maintained in older cultures (E14.5 mouse explants). Photomicrographs of E11.5 VM explants co-cultured with (D) mock-transfected cell aggregates and (E) Wnt5a-transfected cell aggregates. (F) VM explant co-cultured with Wnt5a cell aggregate, illustrating the specificity of Wnt5a to repel TH+ fibers (F) but not total neurite fibers (F', Tuj1-labeled fibers). (G-I) Images illustrating the effects of Wnt5a on neurite chemotaxis could be ablated by (G) D4476, (H) NSC23766 and (I) anti-Fz3-CRD. Dashed line demarcates the border of the cell aggregate. Arrow indicates direction of ligand signal (i.e. cell aggregate) relative to explant. Scale bar = 200 mm.

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Figure 7. *Wnt5a*−/− mice display abnormal fasciculation of the DA axons in the median forebrain bundle (MFB) and changes in DA fiber and terminal density. (A) Sagittal image of the Wnt5a+/+ and (B) Wnt5a−/− brain demonstrating morphological changes within the dopaminergic pathway. Note the broadening of the MFB in Wnt5a−/− compared to Wnt5a+/+ mice, indicated by arrows, as well as increased terminal innervation in the dorsal striatum (A' and B'). (C) Wnt5a−/− mice showed defasciculation of the MFB as revealed by the increased volume occupied by TH+ fibers. (D) Within the MFB, Wnt5a−/− mice had significantly more TH+ fibers at proximal (320 μm) levels of the MFB. (E) Wnt5a−/− mice also had significantly more TH+ terminals within the lateral striatum than Wnt5a+/+ mice. (F) Image illustrating the level of the MFB at which neurite counts were performed (HP, hippocampus; Ctx, cortex). (I) Image illustrating the area of the lateral striatum delineated for stereological estimates of TH+
the axon bundle was observed with fibers present in the intermediate zone and encroaching on the ventricular zone of the VM (Figure S3A′ and 3B′), similar to E18 and validating Wnt5a’s role in axon fasciculation.

**Discussion**

Understanding the intricate and precise pattern of connectivity achieved during brain development has been, and remains, an outstanding challenge. This is particularly true for the dopaminergic pathways that arise from the ventral midbrain and innervate distant targets such as the striatum and cortex. Changes in the connectivity of these pathways and in the availability of their neurotransmitter, dopamine, underpin a number of neurological disorders including Parkinson’s disease, schizophrenia and addiction. In this context, our study demonstrates a novel function for Wnt5a in the establishment of the dopaminergic pathways originating in the VM. First, we found that the spatial expression pattern of Wnt5a correlates with the development of the nigrostriatal/mesolimbic DA pathways. Second, we report that Wnt5a selectively regulates the axon length of TH+ cells in primary VM cultures in a time-dependent manner, promoting axon extension at E11.5 (and rat E13.5) and axon retraction at E14.5. Third, we identify Wnt5a as a chemoattractant of DA neurites in mouse E11.5 and E14.5 explant cultures. We confirm that these growth and chemotactic effects of Wnt5a are mediated by Fz3 and involve the activation of Rac1, a component of the Wnt/PCP pathway. Fourth, analysis of the Wnt5a−/− mice revealed that Wnt5a is required for fasciculation of DA axons in the MFB, and for the innervation of the dorsolateral striatum, the target area of substantia nigra DA neurons.

Wnt5a expression within the VM and caudal region of the MFB was maintained during development of the DA pathways. More precise temporal and spatial gradients reflect the functions of this protein in DA axon morphogenesis. We speculate that at E11.5, the high expression of Wnt5a within the ventricular zone and in the rostral part of the ventral midbrain, (Fig. 1A,E) may prevent axons from entering the ventricular zone and from taking a premature anterior direction. In support, we in vitro results at E11.5 indicate that Wnt5a increases neurite length during initial DA axon development (Fig. 2D,E), and repels them away from the source of Wnt5a (Fig. 6B,D,F). Additionally, E12 Wnt5a−/− embryos show reduced neurite length and disorganization of DA neurites, with fibers seen within intermediate layers of the VM (Figure S3). We therefore suggest that the initial effect of Wnt5a is to contribute to DA axonal elongation and to maintain axons within the VM, but out of the ventricular zone (Fig. 8A, D). Later in development, by E14.5, the high rostral expression of Wnt5a is down-regulated and expression is higher in the caudal VM (Fig. 1L). This gradient shift may facilitate DA axons taking a forebrain trajectory by being repelled away from the caudal VM, and preventing their entry into the hindbrain (Fig. 3B). Interestingly, whilst not observed by us, Fenstermaker et al (2010) reported the appearance of DA axons in the hindbrain of Wnt5a−/− mice at E12.5, a phenotype that was lost later in development [55], yet supports our theory of Wnt5a ensuring DA axons maintain their rostral trajectory.

Previous studies have shown that Fz3 is involved in PCP signaling in the context of axon guidance [64] and that Fz3-deficient mice, whilst possessing normal numbers of DA neurons, have no nigrostriatal pathway [57,58]. Moreover, Stereberner et al. has recently reported that Fz3 and Fz6 cooperate to regulate midbrain morphogenesis [65]. Similarly, we have previously reported that Wnt5a mutant mice have a near normal number of DA neurons, but show a clear defect in midbrain morphogenesis [23]. We hereby report that, similar to Fz3, Wnt5a promotes DA neuritogenesis in vitro and that Wnt5a−/− mice exhibit defects in DA axonogenesis. Moreover, we found that Wnt5a effects are mediated by Fz3 or Fz3-like receptors. Fenstermaker et al. (2010) has since illustrated notable defects within the trajectory of DA fibers in Fz3−/− mice, observing several fibers projecting caudally and failing to make striatal contacts [55]. Surprisingly, we observed increased fiber innervation in Wnt5a−/− mice suggesting that Fz3 is only partially responsible for our observations. In the future, analysis of double mutant mice for Fz3 and Wnt5a may more clearly identify these roles and highlight the need to investigate other Wnt related receptors.

Signaling mediated through Frizzled receptors is commonly associated with axon elongation and attraction while, in contrast, signaling through Ryk induces axon repulsion [25–30,66]. However, Ryk receptors activated by Wnt5a gradients promoted elongation of cortical axons in the callosal and corticospinal tract [31,33,35], and chemorepelled these same axons via Wnt5a signaling through both Ryk and Frizzled receptors. These differing actions were mediated by Ca2+ changes involving IP3 receptors and/or TRP channels [33]. Whilst we can not directly attribute the effects of Wnt5a to Ryk binding, the high expression of Ryk within DA neurons of the VM, combined with RYK-Fc antagonism in a DA cell line and existing literature in other pathways, suggests that Ryk may also play a role in regulating DA morphogenesis. Analysis of Ryk-deficient mice as well as double mutants (Wnt5a and Ryk) or siRNA experiments may shed more light on this receptor’s involvement in these processes.

Whilst a number of guidance molecules have been recognized for their role in DA neurite development, few studies have verified these functions in vivo. In our study, we examined the nigrostriatal DA pathway in Wnt5a−/− mice, validating a number of our in vitro findings and the importance of Wnt5a in DA connectivity. In wildtype mice, TH+ axons remain tightly fasciculated within the MFB as they project towards their forebrain targets. In contrast, DA axons in the Wnt5a−/− mice were clearly defasciculated throughout the MFB, with broadening of the bundle observed from caudal to rostral levels of the pathway. Interestingly, defasciculation and broadening of the MFB has also been observed in Sema3F−/− mice [9] and Ryk−/− mice showed defasciculation of the callosal axon bundle crossing the midline [35]. These findings indicate that Wnt5a is required to fasciate DA axons in vivo.

Surprisingly, despite the role of Wnt5a in promoting axonal growth, repelling DA axons and regulating fasciculation of the DA fiber bundle, DA axons in Wnt5a−/− mice maintained their appropriate caudo-rostral trajectory, indicating that Wnt5a alone is not necessary to initiate neurite growth or provide directional...
absence of Wnt5a, at E11.5-12, TH of axonal stalling, resulting in increased striatal innervation in Wnt5a \\
Wnt5a
Since we previously reported that the number of neurons in rostral trajectory of these DA axons. (B) At E14.5, a higher caudal to lower rostral Wnt5a gradient (red), combined with the maintained repulsive action of Wnt5a at this age, may ensure TH axons maintain their forebrain trajectory. The ability of Wnt5a to induce axon retraction in vitro (as shown in Fig 3) suggests a plausible role in axon stalling, a key developmental event that prevents axons prematurely entering their forebrain striatal targets. The dotted red line indicates the border of the ganglionic eminence where axon stalling occurs. (C) At E18, low expression of Wnt5a is maintained within the VM. At this stage in development TH axons are present within the GE and are making increasing synaptic contacts. (D) In the absence of Wnt5a, at E11.5-12, TH axons maintain a rostral projection but are shorter and show a lack of organization. (E) By E18, the disorganization of TH fibers is evident by the defasciculation of the MFB in Wnt5a−/− mice. Additionally, axons prematurely enter the target, presumably due to loss of axonal stalling, resulting in increased striatal innervation in Wnt5a−/− mice. Lv, lateral ventricle; Aq, aquaduct; GE, ganglionic eminence; MFB, medial forebrain bundle.

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in vivo. Collectively these findings suggest that other guidance molecules or Wnts might be capable of compensating for the lack of Wnt5a during the establishment of this pathway. However, more axons were detected within the MFB of Wnt5a−/− mice. Since we previously reported that the number of neurons in Wnt5a−/− and Wnt5a+/+ mice is different [23], the increased fiber density in the MFB of Wnt5a−/− mice suggests a possible increase in DA neurites per DA neuron, or alternatively, increased branching of the DA axons. Interestingly, our in vitro data supports both possibilities, but the greater effect on branching suggests a predominant role of Wnt5a on DA neurite branching, rather than on the number of neurites (Fig. 2F,G).

Finally, it is important to note that despite DA axons arriving at the border of the ganglionic eminence at approximately E15 in rats, they do not enter the striatum, but rather increase in numbers until approximately E17 (comparable to mouse E15) [1]. In support of a possible role for Wnt5a in axon stalling, we found that Wnt5a had a negative effect on DA neurite length in vitro, at E14.5 (approximately equivalent to rat E16.5) (Fig. 3), and that axonal innervation of the dorsolateral striatum was accelerated in the Wnt5a−/− mice. These results indicate that Wnt5a may be the repulsive signal that prevents premature entry of DA axons into the striatum. Indeed, both the increased fiber number in the MFB and the dense striatal innervation of the Wnt5a−/− mice may reflect a premature maturation of the pathway (i.e. loss of axonal stalling). Thus our results suggest that one of the functions of Wnt5a in the nigrostriatal system would be to prevent the premature maturation of this pathway.

In summary, our findings identify a number of key roles for Wnt5a in the development of DA axons in vitro and in the maturation of the MFB in vivo. Wnt5a promotes DA axon elongation, retraction and repulsion in a time-dependent manner, as well as maturation and fasciculation of the MFB. These effects, at least in part, are mediated through the Frizzled receptors and downstream activation of the Wnt/PCP pathway. Whilst broadening our knowledge of dopamine development, an understanding of the regulation and promotion of DA axonal growth and guidance may have significant implications for a number of neurological disorders in which the development of nigrostriatal DA axons are affected, as well as enhancing integration of grafted dopamine neurons into the Parkinsonian brain.

Supporting Information

Figure S1 Schematic representation of the site of action of the antagonists employed to identify the pathways mediating the effects of Wnt5a. sFRP, zWnt5a and RYK-Fc act to sequester Wnt5a out of circulation, thereby preventing its interaction with Wnt-related receptors. zFz3-CRD binds directly to the frizzled-3 receptor, thus preventing Wnt-receptor interaction. Dkk1 does not bind Wnt but affects the interaction of Wnt with the LRP co-receptor, thereby affecting canonical Wnt signaling. D4476, a casein kinase 1 antagonist, blocks the Wnt activity-dependent phosphorylation of Dishevelled and thereby prevents downstream canonical and non-canonical Wnt signaling. NSC23766 is a Rac1 antagonist and thereby an inhibitor of the
non-canonical Wnt/PCP pathway, CK1, casein kinase 1; Dikk1, Dickkopf-1; Fz, Frizzled; LRP5/6, low density lipoprotein receptor-related protein 5/6; Fz3-CRD, Fz3 antibody; Wnt5a, Wnt5a antibody.

**Figure S2** Effects of Wnt receptor antagonism on neurites of all VM neurons. Immunocytochemistry for all neurons (TUJ1) revealed that (A) Fz3-CRD and (B) Ryk-Fc had no effect on neurite length of non-TH+ neurons within the VM, indicating that the effects seen in Fig. 5 were specific to DA neurites. Furthermore, the absence of an effect of Fz3-CRD and Ryk-Fc on the general neuronal population verifies the lack of toxicity of these proteins at the doses selected. (TIF)

**Author Contributions**

Conceived and designed the experiments: CLP. Performed the experiments: BDB CRB CVF SAS MLM EA CLP. Analyzed the data: BDB MKH CRB. Wrote the paper: BDB MKH CRB CVF SAS MLM EA CLP.

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