Research Article

Differentiation of Human Embryonic Stem Cells to Sympathetic Neurons: A Potential Model for Understanding Neuroblastoma Pathogenesis

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Received 25 June 2018; Revised 17 August 2018; Accepted 16 September 2018; Published 1 November 2018

Guest Editor: Hui Li

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Background and Aims. Previous studies modelling human neural crest differentiation from stem cells have resulted in a low yield of sympathetic neurons. Our aim was to optimise a method for the differentiation of human embryonic stem cells (hESCs) to sympathetic neuron-like cells (SN) to model normal human SNS development. Results. Using stromal-derived inducing activity (SDIA) of PA6 cells plus BMP4 and B27 supplements, the H9 hESC line was differentiated to neural crest stem-like cells and SN-like cells. After 7 days of PA6 cell coculture, mRNA expression of SNAIL and SOX-9 neural crest specifier genes and the neural marker peripherin (PRPH) increased. Expression of the pluripotency marker OCT 4 decreased, whereas TP53 and LIN28B expression remained high at levels similar to SHSY5Y and IMR32 neuroblastoma cell lines. A 5-fold increase in the expression of the catecholaminergic marker tyrosine hydroxylase (TH) and the noradrenergic marker dopamine betahydroxylase (DBH) was observed by day 7 of differentiation. Fluorescence-activated cell sorting for the neural crest marker p75, enriched for cells expressing p75, DBH, TH, and PRPH, was more specific than p75 neural crest stem cell (NCSC) microbeads. On day 28 post p75 sorting, dual immunofluorescence identified sympathetic neurons by PRPH and TH copositivity in 20% of the cell population. Noradrenergic sympathetic neurons, identified by copositivity for both PHOX2B and DBH, were present in 9.4% ± 5.5% of cells. Conclusions. We have optimised a method for noradrenergic SNS development using the H9 hESC line to improve our understanding of normal human SNS development and, in a future work, the pathogenesis of neuroblastoma.

1. Introduction

The neural crest is a transient embryonic cell population which undergoes extensive migration and differentiation to give rise to a diverse range of cell populations in the embryo, ranging from the peripheral nervous system (sensory, enteric, and autonomic (sympathetic and parasympathetic)) to the craniofacial skeleton and pigment cells (reviewed by [1]). Neural crest cells are multipotent stem cells which can self-renew and in humans undergo extensive migration around the third to fourth weeks of gestation [2]. Sympathetic neurons originate from trunk neural crest cells that arrest their migration upon arrival at the dorsal aorta and begin to express the catecholaminergic and
noradrenergic biosynthetic enzymes tyrosine hydroxylase (TH) and dopamine betahydroxylase (DBH), respectively (Figures 1(a) and 1(b)). Bone morphogenetic proteins (BMPs), multifunctional secreted proteins of the transforming growth factor β superfamily, are secreted in the dorsal aorta and the gut [3] and are important for noradrenergic autonomic specification from the neural crest [4, 5].

Neuroblastoma is an embryonal malignancy originating from neural crest cells which give rise to the sympathetic nervous system (SNS) [3]. It is the most common childhood solid tumour outside the central nervous system, and in contrast to many other paediatric malignancies, high-risk neuroblastoma is fatal in around 50% of patients despite intensive multimodal therapy [6]. In vivo and in vitro observations have shown that neuroblastic tumours appear to recapitulate the development of differentiating, predominantly noradrenergic, sympathetic neurons, and chromaffin cells of the adrenal medulla, suggesting that neuroblastoma arises from aberrant or blocked differentiation in normal SNS development (reviewed in [7]). By modelling the normal development of the neural crest and SNS, it may be possible to understand the pathogenesis of neuroblastoma and other abnormalities of the neural crest, e.g., neurocristopathies.

Human embryonic stem cells (hESCs) and induced pluripotent stem cells (iPSCs) have the potential to provide an unlimited source of cells for both disease modelling and cell replacement therapy. The ability to differentiate hESC to neural crest–derived stem-like cells (NCSCs) and autonomic progenitors provides an important tool for modelling human neural crest development.

Kawasaki and colleagues were the first to demonstrate efficient induction of peripheral autonomic neuronal lineages from murine and primate hESC by coculture with PA6 cells, which possess stromal–derived inducing activity (SDIA) [8, 9]. Mizuseki et al. showed that early exposure of cocultured cells to BMP4 inhibited neural differentiation, whereas late exposure to high concentrations of BMP4 (days 5–9) induced differentiation to neural crest cells and autonomic progenitors [9]. Recently, other studies differentiating hESCs have used BMP4 [10] or a feeder layer [11] to help induce SN differentiation.

The aim of this study was to develop an in vitro model using both BMP4 and a stromal feeder layer for efficient differentiation of hESC to noradrenergic sympathetic neurons (Figures 1(a) and 1(b)). We sought to determine the optimum conditions for the differentiation of hESC to SN by comparing different neural differentiation media, sorting methods for neural crest–like cells, and plating conditions for sorted cells.

Understanding normal SNS development in hESC models will enable us to learn more about the SNS as well as neural crest–derived malignancies such as neuroblastoma.

2. Materials and Methods

2.1. Cell Culture. H9 cells were obtained from the WiCell Bank (Wisconsin) following approval from the UK Medical Research Council (MRC) Stem Cell Steering Committee. Undifferentiated H9 hESCs [12] were cultured on either the human foreskin fibroblast cell line (NclFed(R)1A) [13], inactivated with 35 Gy ionising radiation, or irradiated MEF-CF1 standard density cells (AMSBIO, UK). hESCs were cultured in stem cell media (20% KnockOut Serum Replacement (Invitrogen, USA), 0.1% nonessential amino acids (NEAA) (Invitrogen, USA), 0.1 mM β-mercaptoethanol (Invitrogen, USA), 2 mM Glutamax (Invitrogen, USA), and 8 ng/ml FGF2 (Invitrogen, USA) in KO-DMEM (Invitrogen, USA)). Cells were passaged weekly and replated on 6-well plates coated with irradiated feeder cells at a density of 6.5 × 10^3 cells per well. The mouse stromal PA6 cell line was obtained from the Riken Cell Bank (Japan).

All cell lines were checked regularly and found to be free from contamination with Mycoplasma. Karyotypic analysis of H9 cells was also undertaken to confirm their identity using standard Giemsa banding techniques.

SKNAS (S-substrate adherent type, non-MYC amplified) [14], IMR32 (N-neuronal type, MYCN amplified [15]), and SHSY5Y (N type, non-MYC amplified) [16] human neuroblastoma cell lines were used as controls.

2.2. Differentiation to Neural Crest–Like Cells and Sympathetic Progenitors. Neural crest differentiation was induced by coculture of hESC with PA6 cells in neural differentiation media as outlined in Figure 1(c). Cells were detached from Fed1A feeders using 1 mg/ml collagenase IV and incubated for 10 minutes at 37°C to detach hESC colonies. 500–800 cells were transferred to 12- or 24-well plates, each well containing 1 or 0.5 × 10^4 PA6 cells, respectively, and cultured for up to 28 days.

To optimise differentiation to p75+ve neural crest–like cells, two neural differentiation media were compared: (1) neural BHK media (90% BHK-21 medium/Glasgow modified Eagle medium (MEM) with 10% KO-SR, 1% L-Glutamax, 0.5% NEAA, 1% pyruvate, 1% penicillin/streptomycin, and 2 × N2 neuronal supplement) and (2) 90% MACS® neuronal media plus 2% MACS B27 supplement, 1% L-Glutamax, 0.5% NEAA, and 1% penicillin/streptomycin. Conditions required for optimal development of noradrenergic sympathetic neurons were established by comparing the addition or withdrawal of 10 ng/ml BMP4 and 10 ng/ml BMP2 and 4 with 10 ng/ml BMP4 alone, on days 5 to 9 of differentiation. N2 supplement (Life Technologies) and 10 ng/ml nerve growth factor (NGF) (R&D Systems) were added to the media from day 4 of differentiation and 0.1 mM dibutyryl cyclic AMP (dbcAMP) was added on day 8 and withdrawn from the media after 10 days of differentiation. Cells were further differentiated for 3–4 weeks, and media were changed every two days. Differentiation experiments were carried out to n = 3.

2.3. p75 (CD271) Fluorescence–Activated Cell Sorting (FACS) of Differentiating Cells. Two methods of cell sorting for p75-positive cells were used to compare the yield of p75-positive cells obtained.

H9 cells were harvested on day 8 of PA6 coculture, which was found to yield the optimal number of p75-positive cells (data not shown) using 1 mg/ml collagenase, incubated for
10 minutes at 37°C, followed by incubation in Accumax (Stemgent) for 10 minutes with gentle agitation to obtain a single cell suspension. Cells were washed once in FACS wash buffer (1 in 20 dilution of bovine serum albumin (Miltenyi Biotec, UK) in MACs rinsing solution (Miltenyi Biotec) and centrifuged at 150 g for 4 minutes. 10% Fc block in FACS wash buffer was added to the cell suspension and incubated for 10 minutes at room temperature. Anti-p75 (CD 271) primary antibody directly conjugated with phycoerythrin (PE) (Miltenyi Biotec) was added to cells at 1 in 33 dilution. Multiple cell sorts were performed (n = 3) using a FACS Aria II Cell Sorter (BD Bioscience™) and a minimum of 5.5 × 10⁴ p75⁺ and p75⁻ cells plated onto either PA6-coated 24 well plates or BD BioCoat™ poly-l-ornithine/laminin-coated 24 well plates (BD Biosciences). All sorted cells were cultured in MACS neuronal medium containing B27 supplement, 10 ng/ml NGF, 10 ng/ml fibroblast growth factor (FGF2), and 10 ng/ml epidermal growth factor (EGF).
2.4. Neural Crest Stem Cell (NCSC) Microbead Sorting. Differentiating H9 cells were harvested using Trypsin/EDTA and dissociated to a cell suspension. Positive cell enrichment was performed using anti-p75- (CD271) coated NCSC Microbeads, according to the manufacturer’s instructions (Miltenyi Biotec). Sorted cells were cultured in either 24-well PA6-coated plates or BD BioCoat™ poly-l-ornithine/laminin-coated 24-well plates.

2.5. Live Cell Immunofluorescence of Undifferentiated H9 Cells. Undifferentiated H9 hESC colonies were immunostained with 1 : 100 dilutions of TRA-1-60-FITC conjugate (Millipore) and anti-SSEA-4/clone MC-813-70-PE conjugate (Millipore). HESC colonies were incubated with primary antibodies at 37°C for 2 hours followed by a 10-minute incubation with 0.5 μg/ml Hoechst-hESC media solution and twice washed with hESC media to ensure all Hoechst dye was removed. The colonies were then imaged in hESC media under a Nikon eclipse TE2000U inverted microscope after which the media was replaced with fresh hESC media containing 10 μM Rho-associated kinase (ROCK) inhibitor Stem-molecule™ Y27632 [17] (Stemgent, MA, USA).

2.6. Immunofluorescence of Differentiating H9 Cells. H9 cells were immunostained for stem cell and neuronal markers. Cells were washed in PBS and fixed in 4% paraformaldehyde for 10 minutes. After washing in PBS (3 × 5-minute washes), the samples were incubated in blocking solution containing 1% BSA and 10% goat serum. The following antibodies were used at the dilutions indicated: OCT4 1 : 400 (Abcam), NANOG (R&D systems) 1 : 200, neuronal cell adhesion molecule (NCAM) 1 : 200 (Millipore), peripherin (PRPH) 7C5 and C-19 1 : 200 (Santa Cruz Biotechnology), TH 1 : 450 (Millipore), DBH, 1 : 450 (Abcam), and paired like homeobox2B (PHOX2B) 1 : 450 (Santa Cruz Biotechnology). Cells were incubated with primary antibodies overnight at either 4°C or room temperature for 1.5 hours. Secondary antibodies coupled to Alexa Fluor 488 or 568 (Molecular Probes, USA) were used for detection and were used alone as controls for two-colour co-staining as well as comparison with single markers alone. Cells were washed with 3 × 10-minute washes and nuclei stained using 4′,6-diamidino-2-phenylindole, dihydrochloride DAPI (Vectashield) diluted 1 : 10 in PBS in 24-well plates, and coverslips were stained with DAPI. 24-well plates and 4-well chamber slides (Millipore, UK) were viewed and photographed using a Nikon A1r confocal microscope. Percentages of positively immunostained cells were obtained by counting 100 cells each in 3 different areas of the slide and then scoring the number of positive cells. This scoring process was applied to all experimental replicates.

2.7. Time Lapse Photography. Live cell analysis and imaging of p75+ and p75− H9 cells were performed over 4 days using a Nikon Biostation Cell Tracker. Cells were imaged on both PA6-coated 24-well plates and poly-l-ornithine/laminin 24-well plates. The migration rates, including velocity and meandering index, were measured using the Volocity™ software programme (Perkin Elmer, UK).

2.8. RNA Analysis-RT-PCR. RNA was extracted using the RNeasy mini kit (Qiagen) and 0.5 μg reverse transcribed using the iscript cDNA synthesis kit (BioRAD™). RNA was isolated from differentiated H9 cells on day 7, 14, and 21 of differentiation. Undifferentiated stem cells and SKNAS, SHSY5Y, and IMR-32 neuroblastoma cell lines were used as controls. In addition, normal human adrenal cortex, medulla, and dorsal root ganglion tissue from a 14-week gestation fetus were supplied by the Human Developmental Biology Resource (http://www.hdbr.org).

RT-PCR reactions were set up in a total volume of 20 μl containing 10% PCR buffer, 10% magnesium chloride (MgCl₂), 10% dNTPs, 1 μl of 10 μM forward and reverse primers, and 1% Amplitaq Gold™ (Applied Biosystems). RT-PCR was performed for neural crest specifiers (SNAIL, and SOX9), SNS precursors, (PHOX2b and TH), noradrenergic sympathetic neurons (DBH), and other neuronal markers (NCAM, PRPH) on days 7, 14, and 21 of coculture of unsorted cells, as well as on day 8 for p75+ ve FACS sorting (see Supplementary Information 1 for primer sequences). Densitometry was performed using ImageJ software (NIH, USA). Total intensity was calculated from PCR bands and first normalised to GAPDH, mRNA expression for each target was then calculated as fold change relative to undifferentiated H9 stem cells.

2.9. Quantitative Reverse Transcriptase PCR (QRT-PCR). Taqman® gene expression assay primers and probes were used to amplify OCT4, TP53, DBH, and GAPDH. RT-PCR was performed in a total reaction volume of 10 μl containing 5 μl of Taqman universal PCR master mix, 0.5 μl of primers and probes mix (Applied Biosystems), 2.5 μl cDNA, and 2 μl H₂O. Reactions were performed in triplicate then quantified using the ABI Prism 7900HT sequence detection system (Applied Biosystems) relative to GAPDH.

2.10. Statistics. A chi-squared test was used to compare the percentages of p75+ and p75− ve cells obtained using MACS® neuronal media and neural BHK media (n = 3). Continuous velocity data for p75+ cells grown on PA6 cells v poly-l-ornithine/laminin plates was summarised using the mean and standard deviation for normally distributed data and medians with quartiles for skewed data. Log transformations were performed to reduce skewness, and two sample t-tests were used to compare normally distributed data. The statistical package STATA version 14.1 (StataCorp 2015, Stata Statistical Software: Release 14; College Station, TX: StataCorp LP) was used for statistical analyses.

3. Results
3.1. Confirmation of Phenotype and Karyotype of H9-Undifferentiated hESC. Live cell staining using antibodies specific to the pluripotency markers SSEA4 and TRA-1-60 showed intense staining of undifferentiated H9 ES cell colonies (Figures 2(a-i) and 2(b-i)). Hoechst staining of H9 cells was less intense than in the FeedA feeder cell layer, indicative of a “Hoechst dim” phenotype due to efflux of Hoechst dye by stem cells, in contrast to brighter staining
observed on the feeder layer (Figures 2(a-ii) and 2(b-ii)). Kar-
yotyping of H9 cells confirmed a normal female karyotype (Figure S1).

3.2. Sympathetic Neuronal Differentiation of H9 Cells Detected by Immunofluorescence and RT-PCR. Following 7
days of PA6 coculture, morphological changes towards a
neuronal phenotype were observed in H9 cells using phase
contrast microscopy. Immunofluorescence of H9 cells on
day 21 of differentiation for neuronal markers showed
>90% of cocultured cells immunostained positive for NCAM
(Figure 3(a)) with PRPH positivity in 20% (Figure 3(c)). TH-
positive cells were observed in around 10% of the cell popu-
lation (Figure 3(b)). DBH was detected in approximately 5%
of differentiated cells (Figure 3(d)). This cell population
included around 10% PA6 cells, and thus, the percentage of
cells differentiating towards a sympathetic neuronal lineage
is likely to be slightly higher than this.
mRNA expression by RT-PCR of early neural crest specifier genes including SNAIL showed a twofold increase between day 0 and day 14, and SOX-9 expression increased 14-fold by day 14 (Figure 4(a)). The addition of BMP4 (10 ng/ml) on days 5–9 of coculture led to a 12-fold increase in PRPH expression from day 7 to day 21 of differentiation compared to 8-fold without BMP4 (Figure 4(a)). p75 expression was highest between day 7 and day 14 of differentiation and increased upon addition of BMP4 (Figures 4(a) and 4(b)). Low basal expression of p75, DBH, and TH was detected, possibly due to spontaneous differentiation into neuronal cells as has previously been reported for H9 hESC [18]. High expression of p75 was observed in the neural crest-derived control tissues, fetal adrenal gland comprising adrenal cortex and medulla and control dorsal root ganglion (sensory neurons). There was a 5-fold increase in DBH expression and a 6-fold increase in TH expression by day 7 of differentiation compared to control H9 cells. High expression of TH and DBH was observed in the positive controls (IMR32 and SHSY5Y cells). DBH expression on day 7 was comparable with one of the fetal adrenal glands, and as expected, the fetal dorsal root ganglion showed low expression of DBH (Figure 4(b)). The highest expression of the pluripotency gene LIN28B expression was observed in H9 cells on day 7 of differentiation (Figure 4(b)).

3.3. Quantitative RT-PCR of Differentiating H9 Cells. The highest expression of the pluripotency marker OCT4 was observed in undifferentiated H9 cells, decreasing by day 7 through day 21 of differentiation (Figure 5(a)). SHSY5Y and IMR-32 neuroblastoma cell lines expressed low levels of OCT4 compared with undifferentiated H9 cells (Figure 5(a)). TP53 was expressed in pluripotent H9 cells and expression maintained throughout differentiation at levels comparable with the neuroblastoma cell lines (Figure 5(b)). DBH expression was very low in undifferentiated H9 cells and increased to a maximum on day 7 of differentiation before declining during the later stages of differentiation (Figure 5(c)), consistent with the data obtained by RT-PCR where the highest expression of DBH was also on day 7 of differentiation (Figure 4(b)). As expected, control neuroblastoma cell lines expressed very high levels of DBH (Figure 5(c)).
3.4. Enrichment of Sympathetic Progenitor-Like Cells by p75 Sorting. To enrich for NCDSC and sympathetic neuron-like cells, cells were sorted using FACS for the NCSC marker p75 (CD271) on day 8 of coculture (Figure 4(b)). To optimise neural crest stem cell differentiation further, two different neuronal media types were compared: (a) MACS® neuronal media + B27 and (b) neural BHK media + N2 supplement. On day 8 of differentiation, an increased proportion of p75+ cells were observed using MACS® neuronal media 45.0% ± 0.8 (95% confidence interval) compared with 32.7 ± 0.8 (95% confidence interval) using neural BHK media (n = 3, p < 0.0001, chi-squared test) (Figure 6(a)). Preliminary results undertaking FACS for both p75 and DBH on day 8 of differentiation showed that 5.5 and 5.7% of cells were p75 and DBH+, by FACS following culture in BHK and MACS media, respectively (data not shown).

Following p75 sorting, autonomic neuronal populations were enriched when cells were grown in preconditioned media from cultured hESC in the presence of N2 supplement, NGF, and dBCAMP (data not shown). On day 28 post p75 sorting, dual immunofluorescence identified sympathetic neurons by PRPH and TH copositivity cells in H9 cells (Figure 6(b-i)) in up to 20% cells. Noradrenergic sympathetic neurons identified by copositivity for both PHOXB2 and DBH were present in 9.4% ± 5.5% H9 cells (Figure 6(b-ii)). mRNA expression by RT-PCR of p75+ ve H9 cells showed enrichment for p75 and the sympathetic neuronal markers, TH and DBH, together with PRPH where expression was observed exclusively in p75+ ve cells (Figure 6(c)). This was in contrast to the early neural crest specifier SNAIL which was expressed in both p75+ ve and p75− ve cells. Following selection of p75+ ve cells using neural crest stem cell microbeads in differentiating H9 cells, p75 mRNA expression was highly enriched in the p75+ ve cell fraction and DBH was expressed exclusively in p75+ ve cells whereas TH was expressed in both the p75-enriched and the p75-depleted cell populations (Figure 6(d)). This indicates possible contamination of p75+ ve cells in the p75-depleted population.

3.5. Live Cell Imaging (Biostation). p75+ ve-sorted cells plated onto PA6-coated plates cells survived better compared to those on poly-l-ornithine/laminin-coated plates (data not shown).
Figure 6: (a–c) Fluorescence-activated cell sorting (FACS) for p75 (neural crest stem cell marker). (a) FACS plots for H9 cells sorted on day 8 of differentiation. (a-i) 44.7% ± 6.1 p75+ viable cells in MACS® neuronal media. (a-ii) 32.5% ± 1.4 p75+ viable cells in neural BHK media. (b) Immunofluorescence of p75+ FACS-sorted H9 cells at different stages of differentiation. (b-i) PRPH and TH copositive H9 cells at 14 days post cell sort. (b-ii) PHOX2B and DBH copositive H9 cells at 14 days post cell sort. (c) mRNA expression by RT-PCR of FACS sorted H9 p75+ and p75−ve fractions showing high expression of TH, DBH, and PERIPHERIN in p75+ cells and undetectable expression in p75− cells and equivalent SNAIL expression in both populations. (d) RT-PCR of p75-enriched H9 cells isolated using anti-human p75 antibody-coated magnetic MicroBeads showing TH and p75 expression in p75-depleted population. +ve = p75 positive fraction, −ve = p75 negative fraction, and c = negative control.
shown). H9 cells appeared to track and follow PA6 cells adhering to them which in turn led to enhanced survival. Time-lapse microscopy revealed that p75+ ve cells showed increased migration compared with p75− ve cells. A t-test for independent groups with correction for unequal variances showed that the mean logged velocity for p75+ ve cells grown on PA6 cells was higher than that for p75− ve cells (p < 0.001). A similar analysis showed that the mean logged velocity for p75+ ve cells grown on poly-l-ornithine/laminin cells was higher than that for p75− ve cells (p < 0.001) (Figures 7(a) and 7(b); Supplementary online video 1).

4. Discussion

The aim of the current study was to optimise a model of normal human sympathetic neuronal development using hESC, which could be used to understand the normal development of the SNS and in the future the pathogenesis of neuroblastoma and other neural crest-derived malignancies.

In this study, we optimised derivation of neural crest-like cells and noradrenergic sympathetic neuron-like cells using a variety of methods. Initially, we tested conditions for neural crest-derived stem line cell differentiation by comparison of two media types: (1) MACS® neuronal media and (2) neural BHK media, where the former was found to be superior for the generation of p75+ ve cells. We also optimised BMP exposure and showed that BMP4 exposure alone was superior, in agreement with previous studies showing that although early exposure to BMP4 can promote dorsal neural differentiation, when applied at later stages, BMP4 enhances the production of NCSC and autonomic neurons in primate and murine cells [9, 19]. BMP signalling is essential for the initiation of differentiation of neural crest cells into sympathetic neurons in the developing embryo [20, 21]. To our knowledge, there is only one other study differentiating hESC to autonomic neurons which has employed the use of BMP4 during differentiation [10], but this study did not use SDIA which is likely to be at least partly responsible for the higher yields of sympathetic neurons we observed (Table 1).

p75 is the low affinity NGF receptor and a well-characterised marker for neural crest-derived stem-like cells [22]. Using a murine in vivo model, NGF was shown to bind the high-affinity NGF receptor (TRKA) which regulates the expression of both TH and DBH in developing and maturing sympathetic neurons [23]. p75 cell sorting has been used previously to purify neural crest stem cells from hESCs [24, 25], and our work has extended this field by showing that day 8 of differentiation induced by SDIA is the optimal time for p75 cell sorting.

The presence of TH and PRPH costaining or DBH and PHOX2B was used to identify catecholaminergic and noradrenergic sympathetic neurons, respectively. TH is the rate-limiting enzyme in the biosynthesis of dopamine and noradrenaline and is a useful marker for catecholaminergic neurons (Figure 1(b)) [26]. PRPH is expressed in neurons of the developing peripheral nervous system [27].

PHOX2B regulates the expression of PHOX2A and heart and neural crest-derived expressed protein 2 (HAND2). Hand2 is induced by BMPs and is first observed after the onset of Phox2B and Asc-1 expression. Overexpression of Hand2 has been shown to induce the generation of catecholaminergic neurons from neural precursor cells both in vitro and in vivo [4, 5]. Furthermore, germline mutations in PHOX2B have been identified in hereditary neuroblastoma [28, 29]. DBH, a specific marker of noradrenergic sympathetic neurons, is expressed in some neuroblastoma cell lines [30]; it catalyses the conversion of dopamine to noradrenaline in the catecholamine synthesis pathway leading to noradrenergic neurons [31]. In the current study, noradrenergic sympathetic neurons were identified by DBH mRNA expression and immunostaining for DBH alone and DBH/PHOX2B copositivity.

Time-lapse analysis of p75+ ve and p75− ve cells showed that p75+ ve cells had increased migration compared with p75− ve cells, consistent with migratory properties of neural

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**Figure 7:** Migration of p75+ and p75− H9 cells sorted by FACS on (a) PA6 coated wells showing a higher mean logged velocity of p75+ cells compared with p75− cells (p < 0.001), and (b) poly L’ornithine/Laminin coated plates again showing a higher mean logged velocity in p75+ cells (p < 0.001).
crest cells [1]. These results mirror findings observed in vivo using live cell imaging during embryogenesis [32].

QRT-PCR gene expression analysis showed high expression of the pluripotency markers \textit{OCT4} and \textit{LIN28B} in pluripotent H9 cells decreasing with the onset of neural differentiation as expected [35]. \textit{TP53} expression was observed throughout differentiation at levels similar to those in neuroblastoma cell lines consistent with evidence that p53 regulates the proliferation and differentiation of neural progenitor cells independently of its role in the induction of apoptosis. \textit{In vivo} studies using transgenic mouse models have demonstrated a fundamental role for p53 during neural stem cell self-renewal and differentiation [36, 37].

Previous studies have used murine neural crest systems to investigate neuroblastoma development by \textit{MYCN} transformation of primary neural crest cells derived from day 9.5 mouse embryos [38]. Further studies also reported that \textit{MYCN} and common \textit{ALK} mutations exert a role in neuroblastoma tumour initiation using neural crest progenitor cell lines MONC-1 and JoMa1 [39, 40]. Very recently, human NCSC derived from hESC have been transformed by \textit{MYCN} to form neuroblastoma in vivo [11]. The use of human stem cell models of sympathoadrenal development such as ours will develop this field further.

In conclusion, our study describes advancement in the generation of noradrenergic sympathetic neuron-like cells from hESC to improve our understanding of the normal development of the human SNS and abnormalities thereof including neural crest-derived malignancies such as neuroblastoma. This model could later be perturbed by oncogenic transformation of these cells with genes known to be important in the development of neuroblastoma including \textit{MYCN} and/or \textit{ALK} as has been reported for NCDSC, to better understand events leading to the development of neuroblastoma, its cell of origin, and new potential treatment targets.

### Table 1: Comparison of the current study with previously published studies reporting the differentiation of pluripotent stem cells to sympathetic neurons.

| Reference | Methods | Cell type | Markers used to identify SNS progenitors | Yield |
|-----------|---------|-----------|-----------------------------------------|-------|
| Mizuseki et al. [9] | PA6 stromal cells/BMP4 | mESCs, primate ESCs | TH+/PRPH+/Phox2b | Not quantified—very low |
| Pomp et al. [33] | PA6 coculture BHK-21 medium/Glasgow MEM + N2 supplement | HES1, HUES1, HUES7 | TH+/PRPH+ | <1% |
| Lee et al. [25] | MS5 stromal cell line 28 days, FGF2/EGF exposure | H9 | TH+/PRPH+ | 1-2% |
| Jiang et al. [24] | PA6 stromal cell line—7 days | H1, H9 | TH+/PRPH+ | 18% |
| Acevedo et al. [34] | Embryoid bodies (EBs)—onto collagen plates | H9 | Dopa decarboxylase (DDC), TH+, Mash 1 | Not determined |
| Huang et al. [10] | Exposure to retinoic acid, BMP2, BMP4, BMP7 | H1, WTC iPSC | TH+/DBH+ | Not determined |
| This study | PA6 stromal cells, MACS® neuronal media + BMP4 | H9 | TH+/PRPH+, DBH+/PHOX2B+ | 20% and 9.4% ± 5.5%, respectively |

### Abbreviations

- **BMP**: Bone morphogenetic protein
- **hESC**: Human embryonic stem cells
- **DBH**: Dopamine beta hydroxylase
- **NCAM**: Neural cell adhesion molecule
- **SDIA**: Stromal-derived inducing activity
- **PRPH**: Peripherin
- **SNS**: Sympathetic nervous system
- **TH**: Tyrosine hydroxylase

### Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

### Disclosure

This work was presented as a poster presentation at the AACR Special Conference on Developmental Biology and Cancer, Boston, USA.

### Conflicts of Interest

The authors declare that they have no conflicts of interest.

### Acknowledgments

We would like to thank the following for neuroblastoma cell lines: Dr. Jean Bernard (SKNAS) and Prof. Penny Lovat (IMR32 and SHSY5Y). We acknowledge the Newcastle University Flow Cytometry Core Facility (FCCF) for the assistance with the generation of flow cytometry data and in particular are very grateful to Mr. Ian Dimmick, Mr. Ian Harvey, and Dr. Owen Hughes. We thank Dr. Alex Laude for the assistance with the use of the Biostation and confocal microscopy and Mr. Jerome Evans for karyotyping the H9 cell line. We are also grateful to Professor Alison Murdoch...
and staff at the Newcastle Fertility Centre for an additional hESC line (NCL-14) which was used to replicate some of the experimental data obtained with H9 cells (not shown). The human embryonic and fetal material was provided by the Joint MRC/Wellcome Trust Human Developmental Biology Resource (http://www.hdbr.org). This study was supported by Neuroblastoma UK, Children with Cancer UK, Newcastle Healthcare Charity, Wellcome Trust (grant # 099175/Z/12/Z), JGW Patterson Foundation, and the North of England Children’s Cancer Research Fund.

Supplementary Materials

Supplementary 1. Figure S1: G-banded karyotype analysis of H9 hESC showing a normal female karyotype.

Supplementary 2. Video 1: p75+ FACS-sorted H9 cells showing motile p75+ cells tracking the adherent PA6 feeder cells over a 12-hour time period between days 8 and 12 of differentiation.

Supplementary 3. Supplementary Information 1: primer sequences (forward and reverse) including product length and annealing temperatures are detailed below 5’ to 3’.

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