Spontaneous Graft-Induced Dyskinesias Are Independent of 5-HT Neurons and Levodopa Priming in a Model of Parkinson’s Disease

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ABSTRACT: Background: The risk of graft-induced dyskinesias (GIDs) presents a major challenge in progressing cell transplantation as a therapy for Parkinson’s disease. Current theories implicate the presence of grafted serotonin neurons, hotspots of dopamine release, neuroinflammation and established levodopa-induced dyskinesia.

Objective: To elucidate the mechanisms of GIDs.

Methods: Neonatally desensitized, dopamine denervated rats received intrastriatal grafts of human embryonic stem cells (hESCs) differentiated into either ventral midbrain dopaminergic progenitor (vmDA) (n = 15) or ventral forebrain cells (n = 14).

Results: Of the eight rats with surviving grafts, two vmDA rats developed chronic spontaneous GIDs, which were observed at 30 weeks post-transplantation. GIDs were inhibited by D2-like receptor antagonists and not affected by 5-HT1A/1B/5-HT6 agonists/antagonists.

Conclusions: These findings argue against current thinking that rats do not develop spontaneous GID and that serotonin neurons are causative, rather indicating that GID can be induced in rats by hESC-derived dopamine grafts and, critically, can occur independently of both previous levodopa exposure and grafted serotonin neurons. © 2021 The Authors. Movement Disorders published by Wiley Periodicals LLC on behalf of International Parkinson and Movement Disorder Society

Key Words: Parkinson’s disease; dopamine; cell therapy; graft-induced dyskinesias; neuroinflammation; microglia; 5-HTL-dopa

Clinical trials using fetal dopamine (DA) cells as a neuroreparative strategy for Parkinson’s disease (PD) have produced remarkable long-term recovery of function in some patients1 and graft survival for over 20 years.2 However, the results have been variable and a proportion of patients in three major trials developed persistent abnormal involuntary movements (AIMs), now termed “graft-induced dyskinesias” (GIDs), occurring in the absence of levodopa (l-dopa).3-5 Pre-existing l-dopa induced dyskinesias (LID) have been identified as a potential risk factor for the development of GIDs, but other potential risk factors include aberrant immune responses,6-7 incomplete graft innervation,8 5-HT neurons in the graft,9 abnormal ratios of 5-HT/DA receptors,10-12 or activity of specific 5-HT receptors.13 Investigations of the mechanisms underlying GIDs have been hampered by rodent models that do not replicate the spontaneous nature of the behaviors. Rather they rely on administration of l-dopa or amphetamine to generate acute, transient GID expression14,15 in transplanted animals.

With the evolution of human stem cell (hESC)-derived sources of cells for transplantation, it is now possible to generate ventral midbrain DA (vmDA) grafts free of 5-HTergic neurons and to assess if GIDs can occur also in the absence of 5-HT. Intrastralatal grafts of hESC-derived vmDA cells have been widely reported to survive, integrate, release DA, and alleviate functional impairments in rodent models of PD,16-19 but GIDs have not been explored. To ensure the safety of new cell products for transplantation, it is imperative that we evaluate and understand the potential risk of side effects.

In the process of studying the long-term functional efficacy of hESC-derived vmDA grafts in 6-hydroxydopamine (6-OHDA) lesioned rats, we unexpectedly observed spontaneous, continuous abnormal AIMs in a subset of rats. The
Materials and Methods

Experiments were conducted in compliance with the United Kingdom (UK) Animals (Scientific Procedures) Act 1986 under Home Office License No. 30/2498, with the approval of the local Cardiff University Ethics Review Committee.

Sprague–Dawley rat pups (female, n = 29) were neonatally desensitized at 2 days post-birth with mixed hESC-derived neural progenitors and mature neurons (Fig. 1A), as described elsewhere. Rats were housed in groups of 3–4, with a 14-hour:10-hour light:dark cycle. At 20 weeks old, rats received MFB 6-OHDA lesions, as previously described. Rats were sorted into matched groups based on amphetamine-induced rotations (2.5 mg/kg) conducted at 4 weeks post-lesion (mean net rotations/minute: vmDA = 15.3 ± 0.8; vFB = 14.6 ± 1.0). Rats received intrastriatal transplants of hESC-derived cells (vmDA or vFB). H9 cells were differentiated for 16 days and made into a suspension of 60,000 cells/μL, as previously described. For both transplanted groups, 4 μL (240,000 total cells/graft) was injected into the neostriatum at the following coordinates: (1) AP: +0.5, ML: −3.0; (2) AP: +1.2, ML: −2.7; (3) DVs: −4/−5.

Rotational behaviors were measured for 90 minutes (Rotorat, Med Associates) after 2.5 mg/kg methamphetamine (Sigma, cat. no. M8750), for 60 minutes after 0.05 mg/kg apomorphine (Sigma, cat. no. Y0001465) and after administration of other pharmacological agents (Fig. 1B). Observations from 30 weeks post-graft revealed spontaneous AIMs (as described in Breger et al) were scored every 10 minutes for 60 minutes post-administration of saline or drug. The assessor was blind to both the substance administered and the rat group allocation. During AIMs scoring, the experimenter noted whether normal locomotion and exploratory behaviors were evident in the rotameters, to give confidence that any reduction in GID expression was not the consequence of reduced activity overall. Rotational data were collected in an unbiased manner using automated rotameters.

At 52 weeks post-graft, rats were terminally anaesthetized and transcardially perfused (4% paraformaldehyde [PFA]). Brains were processed for peroxidase-based immunohistochemistry as previously described. CD4+, CD8+, and 5-HT+ cells were counted manually within and at the border of the graft. TH+, HuNu+, OX42+/CD11b+, and GFAP+ cells were estimated using unbiased stereology. Two-dimensional stereology was performed (Olympus BX50 microscope, Olympus C.A.S.T. image-analysis software). OX42 was also measured by optical density (Image, NIH).

Pharmacological challenges and rotation data were analyzed using Kruskal–Wallis non-parametric test with Group (non-GID vs. GID) as the factor. Histological data were analyzed by one-way ANOVA with Group (vFB, vmDA non-GID, and vmDA GID) as the factor. Histological and behavioral data were correlated using Spearman’s ρ. All statistical analyses were conducted using IBM SPSS Statistics 25 software.

Results

Dopaminergic Grafts Can Induce Spontaneous GIDs

To obtain long-term survival of human grafts in a rodent model without the need for daily immunosuppression, we applied a model of neonatal desensitization in rats, which required subcutaneous injection of human cells at P0-5. In adulthood, the desensitized rats received unilateral lesions and subsequently hESC-derived vmDA or vFB intrastriatal transplants. Only a subset of the transplanted rats (4/15) had surviving vmDA grafts and, of these, two displayed spontaneous contralateral rotations, a trend for l-dopa-induced rotations and reduced amphetamine-induced rotations (Fig. 1D). These two rats also developed spontaneous AIMs at 30 weeks post-graft (Fig. 1C and Supporting Data). These were compared to rats with smaller surviving vmDA grafts (n = 2) and rats with non-DA, vFB grafts (n = 4/14) (neither the smaller vmDA nor the vFB graft groups displayed spontaneous AIMs or reduced amphetamine-induced rotations).

To elucidate the neurobiological basis of the GIDs, pharmacological challenges were undertaken using receptor agonists and antagonists. AIMs were observed in both the home cage and rotameters (Supplementary Video S1), and scored blind twice post-saline administration (at 30 and 52 weeks post-graft) and twice post-l-dopa administration (Fig. 1C). These sequential observations suggest that GIDs were chronic, stable, and minimally affected by stress or repeated exposure to rotameters. For GID rats, 0.3 mg/kg buspirone (5-HT1A agonist/D2-like antagonist), fenfluramine (5-HT reuptake inhibitor), 8-OH-DPAT+CP94253 (5-HT1A/1B agonists) and SB399885 (5-HT6 antagonist) did not alter expression of the AIMs behaviors. High dose 1 mg/kg buspirone and eticlopride (D2/D3 antagonist) largely eliminated GID expression without reducing normal motor activity.
Larger vmDA Grafts Induce Neuroinflammation and GIDs

Larger vmDA grafts were identified in GID-expressing rats and small vmDA grafts in non-GID rats. The vFB grafts were a similar volume to large vmDA grafts, containing similar numbers of human cells, but with an absence of mature DA neurons (Fig. 2A). Similar numbers of CD4+ and CD8+ t-lymphocytes were present in large vFB and vmDA grafts, but more reactive astrocytes were observed in GID vmDA rats (Fig. 2A,B). No 5-HT+ cells were observed in any of the grafts (Fig. 2A), despite positive control staining evident within the raphe nuclei. Large vmDA grafts were associated with high levels of microglial activation in both hemispheres, whereas small vmDA grafts induced modest microglial activation in both hemispheres. In contrast, large vFB grafts induced almost no microglial activation (Fig. 2A,B). Neither graft volume, total HuNu+...
cells, GFAP, CD4⁺, nor CD8⁺ t-lymphocyte infiltration correlated with the development of spontaneous AIMS (Fig. 2C). High DA neuron content (tyrosine hydroxylase [TH]) and markedly increased microglial activation (Ox42) correlated significantly with GIDs (Fig. 2C).
Discussion

This is the first report of measurable spontaneous GIDs occurring following transplantation of hESC-derived vmDA neurons into an animal model. These behaviors occurred without prior exposure to l-dopa or LID “priming” and with grafts devoid of 5-HT neurons. Given that they were abolished by the D₂ receptor antagonists eticlopride and by buspirone, which acts as a D₂ receptor antagonist at high doses, this suggests that DA itself, and particularly D₂-like family receptors, play a role in mediating GIDs.

These long-term (1 year post-graft) pilot data are important insofar as they suggest that GID emergence is possible in clinical trials of hESC-derived vmDA grafts. This is despite the evidence that hESC-derived cell preparations for clinical use will contain no/limited 5-HT neurons and patient selection will prioritize patients without severe LIDs. Important parallels can be drawn with recently published details on five patients who developed persistent GIDs in a double-blind United States (US) fetal cell transplant study. Both of these datasets challenge current assumptions in the field, which suggest that GIDs expression may be dependent on 5-HT expression. Dopamine is directly implicated in both studies: (1) the presentation of GIDs resembles some elements of classic LIDs; (2) anti-dopaminergic drugs reduce GIDs; (3) GIDs correlate with measures of dopamine (fluorodopa in clinical trials/TH immunohistochemistry here), and (4) GIDs occur in association with functional improvements (Unified Parkinson Disease Rating Scale [UPDRS] in clinical studies, amphetamine rotations here). Although in the present study, grafts contained ~14,000 TH⁺ cells (whereas rats have ~10–12,000 TH⁺ cells/hemisphere), in patients GIDs have been induced from grafts of ~37,000 TH⁺ cells (human brain harbors ~170,000 substantia nigra pars compacta [SNpc] DA neurons) and without evidence of excess dopamine in PET scans.

There has remained an unanswered question about the temporal pattern of GID emergence clinically. One supposition is that it relates to the withdrawal of immunosuppression, potentially triggering inflammatory drivers, which would be consistent with data demonstrating that DA can mediate immune responses. In the current study, the large microglial inflammatory response was specific to brains with vmDA-containing grafts although a direct causal relationship between neuroinflammation and GIDs could not be determined. Alternatively, GID emergence could relate to the time that it takes for DAergic neurons to begin significant maturation and outgrowth as a similar timescale occurred in a trial, which did not immunosuppress graft recipients.

The use of neonatal desensitization makes this the first behavioral study of extended (12 months) human vmDA graft survival times in immunologically intact animals (in the absence of immunosuppression). Graft survival was low in this study (n = 4 grafts/group), but variability within this model has been reported previously. Moreover, all published literature using this model has reported graft survival for a maximum of 6 months post-graft. The data presented here are, to our knowledge, the first report of human-to-rat graft survival up to 12 months post-graft. Even if protection conferred by neonatal desensitization may reduce with time, this model has nevertheless been highly effective in revealing a previously unobserved phenomenon not readily investigated in standard immune-compromised/suppressed animals.

The severity of spontaneous GIDs in these rats were mild-to-moderate in magnitude when compared to LID established in standard protocols in rodents and did not appear to worsen over the observed time. Importantly, if LIDs are not the sole driver for GIDs, then the risks to late-stage patients may not be significantly elevated, whereas their benefits, such as reductions in LID, could be substantially more than early-stage patients. Therefore, this therapy may be open to a broader population of people with PD.

Importantly for future clinical trials, although stem-cell based therapies can be designed to be completely devoid of 5-HT neurons and, therefore, may be considered safer from a GID perspective, our data suggest that this may not completely eliminate the risk for GIDs. The mechanism of action of spontaneous GIDs needs further investigation in larger studies, but if they can be mediated through excess dopamine and/or inflammatory responses, it may be possible to mitigate these through design of the therapeutic delivery.

Conclusion

This is the first study to demonstrate that stem cell-based therapies can induce spontaneous GIDs in a rodent model of PD, simultaneous with functional recovery. The data demonstrate that GID onset can occur independently of l-dopa exposure and 5-HT neurons in the graft. Instead, the data presented here suggest involvement of D₂-like family receptors and suggest that GIDs may be associated with graft inflammation.

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Data Availability Statement
Data available on request from the authors

References


Supporting Data
Additional Supporting Information may be found in the online version of this article at the publisher’s web-site.