Therapeutic Microinjection of Autologous Adult Human Neural Stem Cells and Differentiated Neurons for Parkinson's Disease: Five-Year Post-Operative Outcome

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Abstract: Object. Neural stem cell-derived neurons offer new cellular therapeutic alternatives for diseases of the central nervous system. Selective neural repair can be particularly valuable in progressive degenerative diseases with discrete cell loss, like Parkinson’s disease. Some benefits were previously demonstrated following transplantation of fetal embryonic tissue. This approach, however, carries inherent risks of immunological reactions, infectious transmission, and intractable dyskinesias, in addition to serious ethical concerns.

Methods. Cortical and subcortical tissue samples were obtained during neurosurgical procedures. Neural stem cells were isolated and expanded in vitro for several months. Safety, differentiation and functional studies were performed during the expansion phases. Nine months after harvesting, autologous cell suspensions containing differentiated dopaminergic and GABAergic neurons were microinjected unilaterally in a patient with advanced Parkinson’s disease. 18F-dopa PET studies and neurological evaluations were performed serially (pre/post-operatively).

Results. Over the next 36 months, the overall Unified Parkinson’s Disease Rating Scale (UPDRS) improved by 81% while “on” medication and 83% while “off” medication. At five-years post-operatively, clinical motor scores returned to baseline. At three and twelve months post-operatively, 18F-dopa PET studies showed a 55.6% and 33.2% increase in dopamine uptake in the implanted left putamen.

Conclusions. Adult neural stem cells derived from a patient’s cerebral tissue can become a source of differentiated neurons, useful for grafting in the treatment of Parkinson’s disease. The combined GABAergic and dopaminergic cells produced a long lasting motor improvement. This approach has the potential to make neural stem cell therapy acceptable and available to a large number of patients.

Key Words: Neural stem cell, Parkinson’s disease, Autologous stem cell transplantation, Adult stem cells, Cellular therapy, Clinical translation.

INTRODUCTION

The progressive loss of midbrain dopaminergic neurons innervating the striatum is the major cause of idiopathic Parkinson’s Disease [1-3]. Cellular striatal transplants using human fetal mesencephalic tissue provide some clinical benefits [4-6], but carry inherent risks of immunological reaction [7-9], uncontrolled dyskinesias [10], risks of infectious transmission, in addition to ethical concerns [11]. The identification of neural stem cells (progenitor cells) in the adult brain [12-15] was followed by several studies demonstrating the feasibility of propagating adult human neural stem cells in vitro [16-19].

Neural transplantation in Parkinson’s disease (PD) is based on the assumption that grafted cells will supply dopamine to substitute for the lost nigrostriatal neurons [20, 21]. We have harvested neural progenitor stem cells from several patients’ prefrontal cortical-subcortical region during craniotomy and succeeded in inducing their differentiation into dopamine-secreting neurons [22]. In this report, we present the long-term clinical results following the autologous transplantation of differentiated neural stem cell-derived neurons unilaterally into a patient with Parkinson’s disease.

MATERIALS AND METHODOLOGY

Clinical Material

This 57-year old right-handed patient was diagnosed with idiopathic Parkinson’s disease at age 46. Family history was negative for neurodegenerative disorders. He received medical therapy with dopaminergic agents with mild initial improvement of his symptoms of rigidity, bradykinesia and tremor. His Hoehn and Yahr stage was 4 while “off” (at least 12 hours drug-free) and 2.5 while “on” (1 hour after the usual morning medication) at the time of enrolment in this study.
The equivalent daily intake of levodopa was 600 mg/day. The tremor involving the right extremities became severely disabling and refractory to any increases in medication or changes in medical therapy. The tremor was initially controlled with the insertion of a left thalamic stimulator into the ventro-intermediate nucleus of the thalamus. During this procedure, a small cylindrical biopsy was obtained, (approximately 90 mm³), sampling the prefrontal cortical and subcortical region along the trajectory of the electrode implant. Cedars-Sinai Medical Center Institutional Review Board approved this study and the patient gave informed consent.

Neural Stem Cell Harvesting and Tissue Culture

Specimens were obtained in accordance with the Cedars-Sinai Medical Center Institutional Review Board. Biopsies were placed in sterile ice-cold DMEM/F-12 (GIBCO) containing penicillin-streptomycin, for further dissection. The tissue was cut into small pieces and trypsinized (0.02 mg/ml trypsin in Verseen (GIBCO) at 37 °C for 10 minutes). After adding trypsin inhibitor mixture (Clonetics), tissues were mechanically tritutrated. Cell suspensions were centrifuged at 400 rpm for 5 minutes; pellets were washed once with DMEM/F-12 and plated at density of 5000-10,000 viable cells/ml in the media composed of DMEM/F-12, B27 supplement (GIBCO) and growth factors bFGF (20 ng/ml; Peprotech), EGF (20 ng/ml; Peprotech).

Cell Expansion

Following their isolation, human neural stem cells were grown in F12/DMEM serum free media (GIBCO), supplemented with B27 growth supplement (GIBCO), 20ng/ml of human recombinant bFGF and EGF. Neural stem cells were grown as neurospheres in 25 cm² or 75 cm² Falcon tissue culture dishes in F12/DMEM serum free medium (GIBCO), supplemented with B27 growth supplement (GIBCO), 10-6 M all-trans retinoic acid, 1 mM dibutyryl cyclic AMP, FGF8 (20 ng/ml), and GDNF (20 ng/ml) and cultured 14 days to promote post-mitotic neurons formation and dopaminergic phenotype differentiation. Prior to implantation, cells were removed from the expansion media and matured in the cell differentiation media for 5 minutes at room temperature. Immunological staining using antibodies against tyrosine hydroxylase (TH) and dopa decarboxylase (DDC) were used to evaluate dopaminergic differentiation. Additionally, dopamine secretion was evaluated using HPLC analyses. These studies provide an approximate profile of the final cell suspension to be introduced by microinjection after the initiation of differentiation at the time of implantation.

Immunostaining

Cell cultures were fixed for 20 min at room temperature with 4% paraformaldehyde in PBS, washed 3 times in PBS, pH 7.4, permeabilized using a10 min incubation with 0.1% TritonX-100, and washed again with PBS. Cultures were then incubated in 3% normal goat serum in PBS with 0.1% Tween 20 for at least 1 hour at room temperature. Blocking was followed by incubation with primary antibodies in 1% goat serum + 0.1% Tween 20 for at least 3 hours at room temperature. Antibodies against type III β-tubulin (1:100, Chemicon) and against tyrosine hydroxylase (1:100, Sigma) were used to detect neurons. Antibodies against gamma amino acid decarboxylase (GAD) were used to identify GABAergic neurons (1:1000, Chemicon), anti L-glutamate to detect glutamatergic neurons (1:50, Signature Immunologies), anti glycine to detect glycineric neurons (1:100, Signature Immunologies) and anti choline acetyl transferase (CHAT) to detect cholinergic neurons (1:100, Chemicon). The cultures were washed in PBS at least 3 times, and incubated with secondary antibodies diluted in 1% goat serum with 0.2% Tween 20 for 1 hour at room temperature in the dark. The secondary antibodies were goat anti-mouse FITC (1:200, Sigma) and goat anti-mouse rhodamine (1:200, Boehringer). For double staining, cells were incubated in second primary antibody for required time at room temperature, then washed in PBS/Tween-20 three times for 5 minutes each, then incubated in secondary antibody for 30 minutes at room temperature. For double staining, cells were mixed with both secondary antibodies together. DAPI 300nm was added for 5 minutes at room temperature, then washed in PBS/Tween-20 for 5 minutes at room temperature then removed, washed and allowed slides to dry completely, then mounted with Prolong Antifade mounting media.

Dopamine Synthesis

1 ml of growth media from control or KCl stimulated (50 mM KCl for 30 minutes) cultures (to induce dopamine secretion by adding 50mM KCl for 30 minutes) was collected. Dopamine was immediately stabilized by adding to the culture media 88 μl of 85% orthophosphoric acid and 4.4 mg of metabisulfite. Samples were sent to an HPLC facility where analysis was performed. Dopamine was extracted from samples using aluminum extraction method and analyzed with a reverse-phase C18 column in a MD-TM mobile phase (Esa Inc). Results were validated by co-elution with dopamine standards.

Pre-Implantation Procedures

Prior to implantation, cells were removed from the expansion media and matured in the cell differentiation media to begin post-mitotic differentiation. Cells were induced to differentiate for 3 days, then collected by mild trypsinization (0.01% trypsin in Verseen, 5 minutes at room temperature), washed twice with F12/DMEM medium, resuspended in Dulbecco-modified phosphate-balanced salt solution (GIBCO), washed twice again with F12/DMEM medium and sent to the operating room prior to transplantation in a sterile cell incubator at 37º. Differentiation and dopamine synthesis prior to transplantation were analyzed in vitro from a separate batch of mature cells derived from the same process.

Safety Testing

All specimens were serially tested for sterility over the course of cell expansion and prior to transplantation. Cultures were performed to rule out any viral or bacterial contamination, mycoplasma or endotoxins. Potential tumor-
genicity was ruled out by transplantation of the stem cell isolates into eight nude mice brains (cerebral cortex and striatum). 200,000 stem cells were transplanted per injection site, and brains were analyzed at 5 months post-transplantation. Serial sections of mice brains were analyzed after hematoxylin-eosin staining. Gross morphological, histological and karyotype studies revealed no tumor formation at the site of implant.

Clinical Protocol

The patient was entered in the clinical study for autologous neural stem cell transplantation for idiopathic Parkinson’s disease. Our clinical protocol followed the guidelines from the Core Assessment Program for Intracerebral Transplantation (CAPIT) protocol. Timed testings (pronation-supination, finger dexterity, stand-walk-sit) and UPDRS scores were administered while “off” and “on” medication. The patient demonstrated a positive response to L-dopa following a practically defined “off” examination. One examination was videotaped pre-operatively and post-operatively. Three separate, baseline independent neurological evaluations and medication regimen remained unchanged over six months prior to transplantation. Baseline Mini-Mental status and full neuropsychometric battery were administered. Pre-operative MRI and 18F-DOPA PET scan were obtained within two weeks of surgery. Post-operative clinical assessments were performed at 3, 6, 9, 12, 18, 24, 36, 48 and 60 months by neurological evaluations. Post–transplantation MRI with contrast enhancement and PET scan were obtained at 3 months and 12 months. At 60 months, a brain computerized tomography was obtained.

Functional Imaging

Fluoro-DOPA PET studies were performed after administration of 10.7 mCi of fluorine-18 L-DOPA intravenously one hour after ingestion of 200mg of carbidopa. The patient stopped medication for a least 12 hours before the studies. Tomographic images were acquired on a Siemens/CTI PET scanner and were reconstructed with attenuation correction. Parametric images were generated from regional analysis of the caudate and putamen using a cerebellum region as the free unbound F-DOPA activity in plasma. Studies were obtained two weeks prior to implantation, at 3 and 12 months post-operatively.

Stereotactic Micro-Injection

The second stage of the procedure was performed nine months after harvesting. Under local anesthesia, stereotactic magnetic resonance imaging studies (MRI) were acquired in the Leksell-G stereotactic frame. Surgical planning was made using computerized software, and targets were selected within the left putamen using stereotactic MRI superimposed to the pre-operative Dopa-PET study. Following a single stereotactic craniotomy (burr hole) while awake, 6 microinjections were delivered at a rate of 5 μl/minute with a micro syringe mounted on a hydraulic micro-drive. Sterile cellular suspensions containing approximately six million differentiated neural stem cells (neurons and glial cells) were microinjected into six targets, 4mm apart, in the left post-commissural putamen. Each trajectory delivered 50μl of cell suspension, or approximately one million cells, over 10 minutes per site. No steroids or immunosuppressants were given during or after surgery. The patient was continued on his medication and sent home one day after the surgery.

RESULTS

Neural Stem Cell Culture and Expansion

Neural stem cells were isolated at the time of cortical-subcortical biopsies from several patients and propagated as neurospheres in the presence of epidermal growth factor (EGF) and beta fibroblast growth factor (bFGF) for six months (Figs. 1a and 2). Cells in the neurospheres reproducibly express nestin and musashi mRNA, characteristic markers for neural stem cells, and proliferate with the doubling time of approximately 4 days.

Neural Stem Cell Differentiation

We differentiated several isolates of human adult neurospheres (AC1, AC2, AC3) and analyzed expression of cell-type specific markers βIII-tubulin, glial fibrillar acidic protein (GFAP), and galactocerebroside (GalC). 11% to 35% of differentiated cells consisted of neurons (β-III-tubulin+), 20% to 49% of astrocytes (GFAP +), and 1% to 3% of oligodendrocytes (GalC+) (Table 1). Differentiated neurons (Figs. 1b, c and 3) and tyrosine hydroxylase (TH)-positive cells (Figs. 1d and 4) developed in cultures where growth factors were removed and substituted with a mixture of 10⁻⁶ M all-trans retinoic acid, 1 mM dibutyryl cyclic AMP, FGF8 (20 ng/ml), and GDNF (20 ng/ml). In addition to dopaminergic neurons, differentiated cultures contain astrocytes, oligodendrocytes and several types of neurons, including gamma-aminobutyric acid (GABA)ergic, cholinergic, glycnergic and glutamatergic neurons (Table 2).

Dopamine Synthesis and Secretion

Both immunostaining results and dopamine secretion demonstrate the presence of functional dopaminergic neurons in the treated cultures. Reverse-phase HPLC analysis demonstrated synthesis and secretion of dopamine (Table 2). Dopamine concentration in culture media of differentiated cells 5 days after initiation of differentiation was 100 ± 45 pg/ml. Stimulation of dopamine secretion by exposing these cultures to 50 mM KCl for 30 minutes lead to an approximately three-fold increase in dopamine levels in the culture media (345 ± 74 pg/ml, n=3).

Stereotactic Micro-Injection

The characteristics of the differentiated neuronal transplant from this specific patient cell line “AC2” is shown in Table 2. Our pre-clinical studies have shown a cell survival rate of 10 to 30% at 60 days post-transplantation in rat models. Based on these findings, we could extrapolate that approximately 630,000 neurons, including up to 93,000 dopaminergic and 378,000 GABAergic neurons, from the 6 million cells transplanted in the left putamen, survived the procedure. This number of cells remains well below the estimated number of dopaminergic afferent neurons from the substantia nigra (estimated at 250,000 per hemisphere) needed to fully restore this circuitry.
Table 1. Differentiation of Astrocytes, Oligodendrocytes and Neurons from Adult Human Neural Stem Cell Lines from Patient AC2 After Treatment with $10^{-6}$ M All-Trans Retinoic Acid (RA), 1 mM Dibutyryl Cyclic AMP, FGF8 (20 ng/ml), and GDNF (20 ng/ml)

<table>
<thead>
<tr>
<th>Adult Neural Stem Cell Lines (Cortico-Subcortical Biopsy)</th>
<th>AC2</th>
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<tbody>
<tr>
<td>Astrocytes (GFAP+)</td>
<td>49±3%</td>
</tr>
<tr>
<td>Oligodendrocytes (GalC+)</td>
<td>5±1%</td>
</tr>
<tr>
<td>Neurons (βIII Tubulin+)</td>
<td>35±6%</td>
</tr>
</tbody>
</table>

Clinical Evaluation

The most significant finding of this study was the continued clinical improvement during the first year, which persisted for the following 36 months, but subsequently declined during the 4th and 5th year. The pre-operative UPDRS score of 37/60 (“on/off”) improved to 7/10 at one year post-operatively (81%/83%). The motor UPDRS score before transplantation of 26/41 (“on/off”) improved to 2/5 (92%/88%) one year after transplant. These improvements persisted at 24 and 36 months after transplantation (Fig. 5). At 3 and 6 months post-transplantation when the thalamic stimulator was turned “off,” the baseline right-sided tremor returned to a pre-operative score of 4 (motor UPDRS tremor rating scale). At 9 months post- and up to 48 months post-
transplantation, the tremor score remained at 1 while “off” stimulation and “off” medication. Since the tremor was refractory while “on” medication, the UPDRS improvement while “on” is due to the complete cessation of tremor. The equivalent daily intake of levodopa was reduced to 300mg/day. At 48 months he began to develop rigidity and slowness of movement on the non-operated side. At 60 months, the implanted side regressed to baseline and the non-operated side continued to progress as reflected by the worsening of his motor scores.

**Imaging Studies**

Two weeks before transplantation, the DOPA-PET study showed an asymmetrical deficit in the uptake of the labeled marker, significantly more severe in the left striatum than the right, which correlated with the clinical presentation (Fig. 7a). At three months post-operatively, the dopamine-uptake within the left putamen increased 55.6% above baseline, while other regions of interest (right putamen, left and right caudate) remained within their initial value (Fig. 7b). At twelve months after surgery, the left putamen uptake value remained at 33.2% above baseline and the contralateral putamen increased at 28% above baseline; other striatal regions remained unchanged (Fig. 6). The post-operative MRI at 3 months and one year showed no structural changes within the implanted putamen, no enhancing changes, hypodensity

**Table 2. Characterization of Differentiated Neural Stem Cells Derived from AC2 Cell Line Prior to Implantation.**

<table>
<thead>
<tr>
<th>Differentiated Neurons from AC2 Cell Line</th>
<th>35±6(%)</th>
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<tr>
<td>Tyrosine Hydroxylase(+) neurons</td>
<td>15±3(%)</td>
</tr>
<tr>
<td>DopaDecarboxylase (+)neurons</td>
<td>12±3(%)</td>
</tr>
<tr>
<td>Dopamine baseline secretion(pg/ml)</td>
<td>100±40</td>
</tr>
<tr>
<td>Dopamine secretion after stimulation (pg/ml)</td>
<td>350±82</td>
</tr>
<tr>
<td>GABA neurons</td>
<td>60±8(%)</td>
</tr>
<tr>
<td>Glutamate neurons</td>
<td>20±3(%)</td>
</tr>
<tr>
<td>Cholinergic neurons</td>
<td>2±1(%)</td>
</tr>
<tr>
<td>Glycine neurons</td>
<td>3±1(%)</td>
</tr>
</tbody>
</table>
or mass lesion suggestive of tumor formation. At 60 months, a brain computerized tomography again showed no structural changes in the implanted putamen.

**DISCUSSION**

The use of autologous differentiated neural stem cells for Parkinson’s disease in humans has not previously been documented. In the present study, we demonstrate prolonged clinical regression after autologous differentiated neural stem cell therapy for Parkinson’s disease at five-years post-operatively.

Neural transplantation for Parkinson’s disease aims at replacing the loss of nigrostriatal neurons. Following the equivocal results of two double-blind studies using fetal mesencephalic tissue [4, 23], different cellular therapeutics have since been investigated to better refine and characterize the types of neural replacement. The potential use of neural stem cells as therapeutic agents for neurodegenerative disorders has been extensively reviewed elsewhere [24-26].

Neural stem cells are present in adult avian and mammalian brains [12, 13, 15, 27]. Active neurogenic regions showing continuous dynamic neural cell production have been shown in the granular zone of the hippocampal dentate gyrus and the periventricular subependymal zone [28, 29]. Non-neurogenic regions of the brain can harbor “dormant” neural progenitor cells in many regions, including the white matter,

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**Fig. (4).** Differentiation of human neural stem cells *in vitro*: (a) and (b) Double staining with GFAP, TH and GFAP, TH and DAPI (Bar=100μm) (c) Phase contrast photomicrograph at 10 day following differentiation. (d,e,f) TH-immunostaining of differentiated dopaminergic neurons, GFAP and TH double staining and DAPI, GFAP and TH immunostaining (40X) (Bar=50μm).
demonstrating self-renewal and multipotency in vitro [14, 30]. Active neocortical neurogenesis remains controversial [31-33] and may depend on local permissive factors. Local microenvironment has been modified in vivo to induce intrinsic neurogenesis [34] but has not been shown yet to have significant neural repair potential.

Multiple neural stem cell subpopulations have regional specificity and potential for therapeutic application [25]. We have identified neural stem cells within the prefrontal cortical and subcortical region during neurosurgical procedures. These human adult neural stem cells have certain similarities to embryonic neural stem cells in their multipotentiality, proliferation rate, and expression of regulatory genes [17]. We believe these neural stem cells harvested from the prefrontal cortical-subcortical region represent radial glial progenitor stem cells originating from the primitive neuroepithelium [35, 36].

Our current study is based on the selective isolation, expansion and differentiation of these neural stem cells with epigenetic factors, replicating certain steps of human neuro-
Transplantation of Differentiated Human Adult Neural Stem Cells

The Open Stem Cell Journal, 2009, Volume 1

27

genesis by inducing a shift in the expression pattern of homeobox genes in vitro. Unlike embryonic stem cells, adult neural stem cells have a committed path to the formation of tissue from the central nervous system. In addition, embryonic stem cells have previously been shown to differentiate into neural cell types, but can maintain delayed expression of other phenotypes and can form teratomas [37].

The controlled manipulation of gene expression of adult neural stem cells in vitro allows for the induction of neuronal fate and phenotypic differentiation. Thus, several millions of committed post-mitotic autologous neurons are produced and characterized prior to transplantation. Safety studies during the expansion phase minimize risks of tumorigenicity and transmission of infectious vectors. Several refinements of these techniques, with the recent understanding of molecular events of developmental and regenerative neurogenesis, will optimize these processes.

Unlike the host residual nigrostriatal neurons, regenerated dopaminergic neurons have not been exposed to years of chronic oxidative stress or other predisposing factors leading to neurodegeneration. In this study, 15% of differentiated neurons were characterized in vitro as functional dopaminergic neurons, with evidence of dopamine synthesis and secretion. This reflects a significant increase over fetal mesencephalic tissue, containing an estimated 5% of viable dopaminergic neurons [24]. Until intrinsic neurogenesis can be triggered in vivo in humans and new cells can be induced to migrate, differentiate, and integrate spontaneously into damaged circuits, therapeutic neuroregeneration will require direct delivery of differentiated neurons into specific intracerebral targets.

The exact survival rate and possible integration of implanted differentiated neural stem cells in this patient remain unknown. The clinical improvement observed here is disproportionate to the number of surviving dopamine neurons predicted from our cell survival studies in animals. On the other hand, the contribution of GABAergic cells to the clinical outcome and tremor control is probably important. GABAergic cells play a major role by their local effect on striatal D1 receptors [38], known to be dysfunctional in animal models of Parkinson's disease [39]. Alternatively, implanted GABAergic cells may enhance the direct inhibitory output from the striatum to the overactive motor output of the basal ganglia [40, 3]. These findings challenge the pure dopaminergic deficit dogma of Parkinson's studies in both animal and clinical trials, and suggest a chronic striatal deficit or dysfunction of striatal GABAergic neurons in addition to severe dopaminergic cellular loss in human Parkinson's disease.

The therapeutic roles of undifferentiated neural stem cells and neurotrophic factors contained in the injected cell suspension remain unknown. The previously implanted thalamic stimulator had been kept “off” for four years post-operatively and did not contribute to the clinical evaluation or outcome.

The functional imaging studies showed a significant increase in dopamine uptake within the transplanted putamen at three-months post-transplantation. In contrast, other striatal regions of interest, including the non-transplanted contralateral putamen, remained abnormally low, consistent with Parkinson's disease. This increase can represent presynaptic captation of the marker and be correlated with the survival and functionality of transplanted dopaminergic cells [41, 42]. At one year post-implantation, the dopamine uptake in the injected putamen remained at 33.2% above baseline, suggesting dopaminergic neuron survival. The clinical outcome at one year, however, is disproportionate to these results, contrary to findings in patients receiving fetal tissue [43, 6]. We also noted contralateral putaminal increase in the fluorodopa uptake of 28%. Bilateral clinical improvement has previously been documented in certain cases receiving unilateral fetal transplantation [44, 45], but was not necessarily reflected by functional imaging studies. Separately, sequential grafting has been shown to produce contralateral increase of fluorodopa uptake in the caudate nucleus that did not receive the implant by 20% [46]. Others have noted focal extra-striatal increase in the fluorodopa uptake in early Parkinson's disease patients and this was interpreted as compensatory
In our study, the contralateral putaminal increase at one year remains unclear and could also be interpreted as compensatory.

The continued clinical improvement over time could be explained by the non-dopaminergic interaction of the implanted cell suspensions within the host striatum. This therapeutic effect persisted over three years and is much greater than placebo effect demonstrated at one year in another transplantation study [48]. A word of caution about the significance of these results: first, the severe asymmetrical presentation of this case and improvement after treatment magnifies the overall clinical outcome and second, this single case requires additional clinical studies to substantiate our conclusions. The clinical regression at year four and five post-operatively indicates bilateral progression of the disease. Further refinement of the dosage, dose escalation, and a larger clinical trial is currently being planned to assure reproducibility of these findings. This next clinical trial will consist of a prospective randomized controlled study and will compare best medical treatment to autologous neural stem cell therapy for advanced Parkinson’s disease. Selection criteria based on severity of the disease and genetic background will be among several other aspects that will be analyzed to provide statistically significant data after primary objectives are met.

CONCLUSIONS

Neural stem cells can be isolated from the human adult cerebral cortex, expanded in vitro using epigenetic factors, induced to differentiate into dopaminergic, GABAergic and other types of mature neurons, and selectively delivered back to striatal targets without immunosuppressant. Because of their biocompatibility, safety and potential integration into the host striatum, autologous neural stem cell-derived differentiated neurons represent an alternative to current cell therapy aimed at the restoration of the nigro-striatal circuitry in Parkinson’s disease.

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