Original Article

Dopaminergic innervation of the human subventricular zone: a comparison between Huntington’s chorea and Parkinson’s disease

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Abstract: The subventricular zone retains its neurogenic capacity throughout life and, as such, is often considered a potential source for endogenous repair in neurodegenerative disorders. Because dopamine is believed to stimulate adult neurogenesis, we looked for possible variations in the dopaminergic innervation of the subventricular zone between cases of Huntington’s chorea and Parkinson’s diseases. Antibodies against tyrosine hydroxylase (TH) and proliferating cell nuclear antigen (PCNA) were used as specific markers of dopaminergic axons and cell proliferating activity, respectively. The immunohistochemical approach was applied to postmortem tissue from 2 Parkinson’s disease cases, 4 Huntington’s disease cases, along with age-matched controls. The immunostaining was revealed with either diaminobenzidine or fluorescent-conjugated secondary antibodies. Optical density measurements were made along the entire dorso-ventral extent of the caudate nucleus. An intense TH+ zone was detected along the ventricular border of the caudate nucleus in Huntington’s disease cases, but not in patients with Parkinson’s disease or age-matched controls. This thin (287±38 µm) paraventricular zone was composed of numerous small and densely packed dopamine axon varicosities and overlapped the deep layers of the subventricular zone. Its immunoreactivity was 47±8% more intense than that of adjacent striatal areas. The dopamine innervation of the subventricular zone is strikingly massive in Huntington’s chorea compared to Parkinson’s disease, a finding that concurs with the marked increase in neurogenesis noted in the subventricular zone of Huntington’s disease patients. This finding suggests that dopamine plays a crucial role in mechanisms designed to compensate for the massive striatal neuronal losses that occur in Huntington’s disease.

Keywords: Stem cells, adult neurogenesis, basal ganglia, Parkinson’s disease, Huntington’s chorea, subventricular zone, neurodegenerative disorders, human striatum

Introduction

The subventricular zone (SVZ) is one of the rare brain areas where new neurons are produced throughout life [1, 2]. This germinal zone covers much of the anterior lateral ventricles, including the ventricular surface of the head of caudate nucleus, just beneath the ependymal layer. Under physiological conditions, the SVZ provides neuroblasts that migrate towards the olfactory bulb, where they differentiate into GABAergic and dopaminergic interneurons that integrate themselves into local networks [3-5]. In pathological conditions, such as stroke, demyelinating disorders and neurodegenerative diseases, the rate of adult neurogenesis can be either positively or negatively altered [6-9] and new neurons can migrate towards sites of injury [7, 10]. The close proximity of the SVZ with the striatum makes it a potential source of endogenous neurons that could be engaged in brain repair strategies for Huntington (HD) and Parkinson’s diseases (PD), two neurodegenerative disorders that are characterized by opposite motor deficits and by a marked alteration of the striatal microcircuitry. The pathological hallmark of HD is a massive atrophy of the striatum [11, 12] largely caused by degeneration of the medium spiny projection neurons. In this hereditary hyperkinetic disorder, a robust progenitor cell proliferation in the SVZ has been reported [7]. The pathological
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changes in PD most consistently affect dopaminergic neurons of the substantia nigra that innervate the striatum and the adjacent SVZ. This hypokinetic disorder is generally associated with a decrease of precursor cell proliferation, although there exist conflicting results on this issue [8, 13-15].

Animal studies have been useful to improve our understanding of the role played by various growth factors and signaling molecules in the control of adult neurogenesis. For example, investigations in rodents have led to the concept that dopamine promotes the production of new neurons in the adult SVZ [8, 16, 17]. This view is supported by the fact that experimental lesions of the nigrostriatal dopaminergic pathway in an animal model of PD decreases precursor cell proliferation in the SVZ, whereas proliferation is restored completely by selective agonists of D2-like receptors [8]. Altogether, these findings prompted us to compare the status of the dopamine innervation of the SVZ and adjacent striatal parenchyma in patients who have suffered from idiopathic PD and others affected by HD. The results pertaining to HD patients have been presented in an abbreviated form elsewhere [18], but this material has been reanalyzed here in more details and its functional significance underlined in the light of novel findings gathered in PD patients with the very same quantitative immunohistochemical procedures. Hence, the present paper provides the very first direct comparative study of the dopaminergic innervation of the human SVZ in HD and PD, two neurodegenerative disorders that affect basal ganglia microcircuitry and motor behavior quite differently.

Materials and methods

Tissue collection

The post-mortem material used in this study was gathered from the brains of 2 PD patients, 4 patients who suffered from HD, together with 5 age-matched controls (Table 1). Tissue samples were obtained from the human brain bank of the Centre de recherche de l’Institut universitaire en santé mentale de Québec, which required informed consent before donation of tissues. The Ethics Committee at Université Laval approved the brain collecting procedures, as well as the storage and handling of post-mortem human brain material that has been described previously [19].

In the present study, all PD and HD patients satisfied clinical and neuropathological criteria established for these diseases. The two PD patients were considered at stage 5 on the Hoehn and Yahr scale [20]. The duration of the disease was 19 and 11 years and the daily dose of L-Dopa taken at the time of death was 825 and 500 mg, respectively. One PD patient showed dyskinesias. Lewy bodies and neuronal loss were documented within the substantia nigra of the 2 PD brains. All 4 HD patients had a positive family history for the disease. The striatal atrophy was documented by magnetic
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resonance imaging at least 10 years before the death of the patients, who all displayed choreic movements and severe cognition impairment at the time of death. The post-mortem neuropathological examination of the brains confirmed the marked volumetric atrophy of the striatum and histological analyses of hematoxylin/eosin-stained sections showed important neuronal losses associated with a compensatory gliosis, as revealed by immunostaining for GFAP and CD68. No signs of neurofibrillary degeneration (Gallyas staining) or any other major neuropathological features were disclosed. They were characterized as either grade 3 or 4 according to the Vonsattel’s scale [12]. The duration of the disease ranged from 15 to 21 years. Individuals used as controls had no clinical or pathological signs of neurological or psychiatric diseases. Their striata did not display neuronal losses nor reactive astrocytosis.

**Immunohistochemistry**

Antibodies raised against tyrosine hydroxylase (TH) and proliferating cell nuclear antigen (PCNA) were used as specific markers of dopamine and cell proliferating activity, respectively. Free-floating brain sections were incubated with the primary antibody against TH (mouse monoclonal IgG 1:500, ImmunoStar, Hudson, WI) and PCNA (rabbit polyclonal IgG 1:50, Santa Cruz Biotechnology, Santa Cruz, CA). Biotinylated horse anti-mouse IgG (Vector Laboratories, Burlingame, CA) was used as secondary antibody for TH immunolabeling. For optical density measurements, TH immunostaining was achieved by incubating sections with 2% avidin-biotin complex (ABC standard kit; Vector Laboratories) followed by diaminobenzidine (DAB, 0.05%), to which 0.005% $\text{H}_2\text{O}_2$ was added (see 19 for details). TH and PCNA immunofluorescent staining was revealed with Alexa 488-conjugated streptavidin antibody (1:200, Invitrogen, Carlsbad, CA) and Alexa 568-conjugated goat anti-rabbit secondary antibody (1:200, Invitrogen), respectively. Once dried, sections were treated with an autofluorescence eliminator reagent (Millipore, Billerica, MA). In addition, some sections adjacent to those used for TH/PCNA visualization were stained with an antibody raised against the serotonin (5-hydroxytryptamine) transporter (SERT) to compare the patterns of dopamine and serotonin innervations of the striatum in PD and HD cases. These sections were incubated overnight with a SERT antibody (1:1000, goat polyclonal antibody; Santa Cruz Biotechnology) and immunoreactivity was revealed using DAB as the chromogen.

**Material analysis**

TH-immunostained sections from PD, HD and control brains were carefully examined using a Leica Leitz DM RB light microscope (Leica, ON, Canada). Optical density measurements of TH immunoreactivity were performed on transverse HD brain sections taken at three anteroposterior levels of the striatum (pre-commissural, commissural and post-commissural) with the anterior commissure as a mid-landmark. On each section, 6 photomicrographs were taken at 2 mm interval along the dorsoventral axis of the caudate nucleus, and at 4 mm from the ependymal layer along the mediolateral axis, using a 10X/0.30 objective. Six other photomicrographs equally spaced along the dorsoventral axis were taken at 150 µm from the ependymal layer, upon the intense TH-immunopositive (+) zone of the caudate nucleus. From these photomicrographs, luminance information was collected using the public domain “Image J” processing software from NIH (v. 1.43r). For double-immunostained sections, fluorescence signals were imaged with a confocal laser-scanning microscope LSM 700 (Zeiss, Oberkochen, Germany). The emission signals of Alexa 488 (TH) and Alexa 568 (PCNA) were assigned the green and red colors, respectively. Statistical significance of optical density measurements was assessed by using two-tailed unpaired t-test.

**Results**

Examination of TH-immunostained sections through the human striatum revealed the presence of a thin but very intense TH+ zone lying along the inner surface of the ventricular lining of the caudate nucleus in HD brains but not in PD or control brains (Figure 1). In the pre-commissural portion of the striatum, this TH+ zone covered the entire dorsoventral extent of the ventricular border of the caudate nucleus (Figure 1E), whereas it was restricted to its dorsal two-thirds at commissural and post-commissural levels. This TH+ paraventricular zone, whose thickness ranged from 150 to 400 µm, with a mean value of 287±38 µm, was composed of many small and closely packed TH+
axon varicosities among which some thin and varicose immunoreactive fibers were scattered. In sections immunostained for both TH and PCNA, the intense TH+ zone was found to lie immediately beneath the SVZ, with some overlap with the deeper layers of the SVZ. Several varicose TH+ fibers emerging from this zone were seen to penetrate the more superficial layers of the SVZ, where they arborize in a rather diffuse manner. The SVZ itself was found to be 2 to 3 times thicker in HD cases than in controls (Figure 1B, 1F). In PD, the thickness of the SVZ appears slightly thinner than in controls (Figure 1B, 1D). As expected, TH immunoreactivity was significantly lower in the striatum of PD brains than in that of controls and lower in the putamen compared with the caudate nucleus, whose immunoreactivity displayed a marked mediolateral decreasing gradient (Figure 1C). The TH immunoreactivity of the paraventricular zone located beneath the SVZ was weaker in PD than in controls and HD brains.

Optical density measurements revealed that TH immunostaining was significantly more intense in the paraventricular zone compared to the remaining sectors of the caudate nucleus in all 4 HD brains examined. Despite some inter-individual variations in the mean optical density values, the TH immunostaining in the paraventricular zone was 36% to 55% (mean value of 47±8%) higher than that of adjoining striatal areas. This increase appears to be the result of an augmentation of the number of TH+ axon terminals rather than an increase in TH immunostaining intensity of single TH+ axon terminals.

In sections immunostained for SERT, numerous labeled axons were present in and around the paraventricular zone [18]. Long, thin and varicose SERT+ axons coursed among the myelin-
Discussion

This study has revealed the existence of an intense TH+ zone lining the ventricular border of the caudate nucleus in patients with HD but not in PD or age-matched controls. Giving that TH is the rate limiting enzyme in dopamine synthesis, the presence of a multitude of densely packed TH+ axon terminals suggests the existence in that specific locus of an intense dopamine activity, which might significantly contributes to the robust progenitor cell proliferation that has been previously reported in the SVZ of HD [7]. This increase in the degree of cell proliferation leads to a marked augmentation of the size of the SVZ zone itself, which contrasts strikingly with the severe cell loss and atrophy that occurs in the adjoining portion of the striatum in this neurodegenerative disease. Interestingly, no similar intense SERT+ zone has been observed in the paraventricular region of the striatum in HD brains, nor in that of PD brains, suggesting that serotonin does not play a major role in the increase in size and activity of the SVZ noted in HD brains, nor in the relative reduction of this zone in PD. The absence of a SERT+ paraventricular zone also demonstrates that the atrophy of the caudate nucleus in HD is not sufficient to explain the increased density of TH+ axons reported here.

As in rodents, the adult SVZ in human is composed of three distinct types of cells (i.e. A, B and C) that are differentially distributed in the various layers [7]. Among these three cell types, type C cells are of particular interest because they appear to be preferentially targeted by the dopamine axons and their proliferative capacity is reportedly regulated by this neurotransmitter through D_2-like receptors [8]. Type C cells, also known as transit-amplifying progenitor cells, are located in the deeper part of the SVZ, close to the myelin layer [7], a portion of the SVZ that overlapped the intense TH+ paraventricular zone identified in the present study. In adult rodents, dopamine appears to stimulate the release of epidermal growth factor, which in turn causes C cells to become activated B (glial) cells by acting upon epidermal growth factor receptors that are coexpressed with D_/D_2 receptors on C cells [23]. Dopamine can act either through direct synaptic contact via the varicose dopamine axons that invoke the more superficial layers of the human SVZ or through indirect, volumic transmission, both modes of dopamine release having been well documented at the striatal level [24, 25].

Degeneration of the dopamine nigrostriatal pathway in PD has been associated with a reduction in progenitor cell proliferation [8, 23, 26]. In accordance with these findings, the SVZ of the PD patients examined in the present study was slightly thinner than that of controls. However, whether this relatively minor thinning of the SVZ in PD is associated with a reduction in neurogenesis is unclear since, in contrast to the results of Hoglinger and collaborators [8], van den Berge and coworkers found no difference in the number of proliferating cells in the SVZ of PD brains compared to that of controls [15]. The discrepancy between these two sets of data might simply reflect methodological differences [13] or variability in pharmacotherapy administered to PD patients, including L-Dopa that is known to positively regulate neurogenesis [26]. In contrast to PD, a clear numerical increase in the three types of SVZ cells occurs in HD brains, but much of the cell proliferation is reportedly due to a massive transformation of C cells into B cells [27]. The mechanism responsible for such a major change is unknown but, based on the intense dopamine activity that occurs in this specific portion of the human striatum, as evidenced by the intense TH+ paraventricular zone disclosed in HD brains, we hypothesise that dopamine is a key factor in this remarkable SVZ transformation. The reason for such a prominent increase in the degree of SVZ cell proliferation in HD brains is unknown, but the increase in the production of B cells could be related to the robust gliogenesis that occurs in the striatal tissue in such a pathological condition [28]. It could also represent an
attempt to produce new neurons to compensate for the massive losses of striatal projection neurons that characterize this neurodegenerative disease. If this is the case, the newly generated neurons obviously do not succeed in coping with the powerful neurodegenerative mechanisms at play in this devastating disorder.

Conclusion

The present study has revealed the existence of a thin but intense dopamine zone lying immediately beneath the ventricular surface of the caudate nucleus in HD patients, but not in PD and age-matched controls. The synaptic or volumic release of dopamine by the multitude of small and densely packed dopamine axon varicosities that compose this paraventricular zone, which overlap the deep layers of the SVZ, might play a crucial role in the striking progenitor cell proliferation noted in the SVZ of HD patients. Our findings suggest that the dopamine innervation of the SVZ is a key element in intrinsic cellular mechanisms designed to compensate for the massive striatal neuronal losses that occur in HD.

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