AGE-RELATED CHANGES IN THE CATECHOLAMINERGIC NEURONS OF THE MESOPONTINE TEGMENTUM IN THE RAT

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Immunohistochemistry and computer assisted image analysis were used to examine the age-related changes in tyrosine hydroxylase- (TH-) immunoreactivity in substantia nigra (SN), ventral tegmental area (VTA), locus ceruleus (LC) and dopamine-beta-hydroxylase- (DBH-) immunoreactivity in LC and subceruleus nuclei of the rat. The findings in 3-month-old rats were compared with 28-month-old rats. In SN TH-positive neurons were concentrated in pars compacta and to a lesser extent - in pars lateralis. In VTA the TH-positive neurons were present over the entire area. In LC the immunoreactive perikarya were densely arranged and superimposed, but in subceruleus nuclei they were less numerous and individual cells were clearly discernible. The DBH-immunoreaction distinctly demonstrated the noradrenergic LC and subceruleus neurons. The results indicate of only subtle signs of cell loss in the dopaminergic neuronal population of SN and VTA, whilst the cell loss of the noradrenergic neurons in LC and subceruleus nuclei is evident. On the other hand, considerable age-related dendritic alterations were observed in all catecholaminergic nuclei. Cross-sectional area and optical density (OD) of the TH-immunoreactive neurons in SN, VTA and LC, and of the DBH-immunoreactive neurons in LC and subceruleus nuclei were investigated. In aging the cross-sectional area decreased statistically and OD of the neurons in SN decreased with 13%. In VTA the cross-sectional area did not change its dimensions, while the OD increased with 19%. In LC and subceruleus nuclei the cross-sectional area decreased with 36% and the OD of the neurons decreased with 16%. In conclusion, the most resistant to age-related changes catecholaminergic region in the rat is the VTA, followed by the pars compacta of SN. Rodent LC is a very vulnerable region. Biomed Rev 2007; 18: 45-58.

Key words: substantia nigra, ventral tegmental area, locus ceruleus, tyrosine hydroxylase, dopamine-beta-hydroxylase
INTRODUCTION

Aging is often accompanied by behavioral changes, such as impaired memory, alterations in motor coordination, reduced sensory acuity, sleep pattern changes. Such alterations correspond to neurodegenerative disorders in man occurring with aging, such as Parkinson’s disease and Alzheimer’s disease. In the catecholaminergic nuclei of the brainstem several cellular markers have been characterized such as changes in the neuromelanin content, lipofuscin accumulation, appearance of neurofibrillary tangles, Lewy and Marinesco bodies, reduced neuronal density (necrotic and/or apoptotic cell loss), changes in neuronal arborization, increased populations of glial cells, as well as more recently described diverse neurochemical factors, associated with normal aging and neurodegenerative diseases (neurotransmitter changes, tau protein, alpha-synuclein, iron, free radicals, calcium-binding proteins, neurotrophic factors). These classic and recent markers have been discussed in numerous articles and reviews (1-30); for comprehensive reviews see (31-35).

Substantia nigra (SN) and ventral tegmental area (VTA) contain the major dopaminergic neuronal groups in the brainstem and are reliably demonstrated by immunoreaction for the enzyme tyrosine hydroxylase (TH). The TH is a rate-limiting enzyme for dopamine synthesis, therefore, immunohistochemistry for TH can be used as a marker of dopaminergic neuronal alterations in these regions, especially in the brains of lower mammals that lack neuromelanin in the catecholaminergic neurons (reviewed in 34). Immunohistochemical investigations of the TH-immunoreactive neurons during aging in humans and different animal species have so far yielded controversial results. There are data on age-related changes of density of the TH-immunoreactive neurons (11,36-43), changes of TH-activity (29,44-46), changes of TH mRNA activity (38, 47), and changes in the dopamine content of the cells (37,44,48-52). The LC and subceruleus nuclei (dorsal and ventral) attracted much attention in the recent decades and there is a considerable body of literature, reporting a decrease of the neuron number with advancing age (4,53-56). In most studies neuromelanin was used as a marker of noradrenergic neurons, and some studies have additionally used immunohistochemical staining with antibody against TH (56-59), or against dopamine-beta-hydroxylase (DBH) (58-60).

We investigated qualitatively and quantitatively the changes in the morphological appearance, cross-sectional area and optical density (OD) of the TH-immunoreactive neurons in SN, VTA, LC, and of DBH-immunoreactive neurons in LC and subceruleus nuclei during aging in rats. A preliminary report was presented (61).

We analyzed young (3-months-old) and aged (28-months-old) Wistar rats by staining their brains under standard immunohistochemical protocols (62) with antibodies directed against TH and DBH. For reconstruction of the investigated nuclei tissue sections were arranged according to the stereotaxic atlas of Paxinos and Watson (63). Subsequent image analysis was performed qualitatively and quantitatively by measuring and comparison of cross-sectional area and optical density (OD) of positive neurons. Quantitative analysis was performed on an image analysis system consisting of Jenaval photomicroscope, Hitachi videocamera, IBM computer and appropriate software. Every positive neuron was measured and the calculation was made as follows: the mean value of the area surface of the neurons in the young animals was accepted as 100%. The OD of the cell bodies was expressed on an arbitrary scale ranging from 0 to 256 (0 – absolutely black, 256 – the OD of the adjacent glass outside of the section). The results obtained were statistically processed using Student’s t-test and mean ± standard error of the area surface and OD were calculated. Statistical significance was accepted at p<0.05.

Qualitative data

In the SN of young (control) rats, the TH-immunoreactive neurons were densely arranged in pars compacta (Fig. 1a) and a lower number was found in pars lateralis, while in pars reticulata only a few immunopositive cells were observed. In the VTA the TH-immunoreactive cells were distributed over the entire area, and were condensed in the rostral linear nucleus, the paranigral nucleus, and especially – in the interfascicular nucleus (Fig. 2a, c). In the LC the TH- and DBH-immunoreactive neurons were extremely densely arranged (Fig. 3a, 4a), whilst in the dorsal and ventral subceruleus nuclei the cells were more loosely arranged and the individual immunopositive neurons were clearly outlined with Golgi silver impregnation-like quality, especially by the DBH-immunolabeling (Fig. 4a, 5a, b).

The immunohistochemical reaction for TH and DBH revealed that the reaction product was brownish, homogenous and filled the entire perikaryon of the labeled neurons. As a rule, it extended in the dendrites and axons of the catecholaminergic neurons. No reaction product was found in the neuronal nuclei.

In SN and VTA of young animals the TH-immunoreactive neurons displayed different size and shape. In SN pars com-
Figure 1. SN pars compacta. (a) Demonstration of TH-immunoreactivity in 3-month-old rat. Medial is to the right. Densely arranged labeled neurons in the medial and central portions of pars compacta; in pars reticulata (to the left) only few scattered labeled neurons are seen, x 100. (b) Detail from (a). Most of the neurons are elongated, oriented in medio-lateral direction. The neuropil contains a dense plexus of labeled processes, x 400. (c) Demonstration of TH-immunoreactivity in 28-month-old rat. Medial is to the right. Medial portion of pars compacta, close to the dorso-ventrally oriented axons of the oculomotor nerve, x 100. (d) Detail from (c). Some neurons display distorted perikarya. Many dendrites are amputated and the dendritic stumps are indicated by arrow heads, x 400.

In SN pars compacta many labeled neurons were elongated, with the long axis oriented medio-laterally (Fig. 1b). The dendrites arose usually from the two perikaryal poles but many neurons in the ventral part of pars compacta emitted also ventrally oriented dendrites that penetrated the pars reticulata. The neuropil of pars compacta exhibited a dense plexus of immunolabeled processes. In the VTA multipolar neurons predominated. Especially in regions with dense neuronal aggregations, like the rostral linear nucleus (Fig. 2b), the neuropil contained a dense network of thin dendrites and axons.

The histological changes in the TH-immunoreactive neuronal populations of SN (Fig. 1c, d), and VTA (Fig. 2c, d) in aged animals were relatively mild. In SN pars compacta only subtle signs of cell loss were encountered: 1.5–3.5% compared to the young animals. In SN many elongated neurons appeared even more “slender”, nearly fusiform, and some perikarya were distorted. Amputated dendrites, with only dendritic stumps left, were a common finding and thin secondary dendrites were few (Fig. 1d). The density of neuropil labeling diminished. In VTA (Fig. 2c, d) a limited number of the perikarya appeared shrunken and darker, most evident in the elongated neurons of nucleus paranigralis (Fig. 2d). The loss of dendrites was com-
Figure 2. VTA. (a) TH-immunoreactivity in neurons of 3-month-old rat. The abundant blood vessels indicate the midline. Between them the densely arranged immunoreactive neurons of the rostral linear nucleus are seen, x 100. (b) Detail from (a). The neurons in the rostral linear nucleus are slightly smaller than the TH-immunopositive cells of pars compacta. Most of the perikarya are rounded-multipolar. The neuropil contains a dense plexus of thin, tortuous and varicose dendrites, x 400. (c) TH-immunoreactivity in 28-month-old rat. The dense neuronal aggregation around the blood vessel is the midline, unpaired interfascicular nucleus. Lateral to it, and medial to the axons of the oculomotor nerve are the more loosely arranged neurons of the paranigral nucleus, x 100. (d) Detail from (c). Lateral border of the interfascicular nucleus and medial portion of the paranigral nucleus. Some neurons retain their dendrites but many cells display amputated dendrites close to the perikarya (arrow heads), x 400.

mon, and from certain perikarya only very short and thickened dendritic stumps arose (Fig. 2d).

The immunostaining for TH and DBH in LC and dorsal subceruleus nucleus in young animals (Fig. 3a, b, 4a) exhibited identical results. LC was sharply outlined from the pale adjacent regions. Within the nucleus the neurons were superimposed and details of the individual cells were discriminated with difficulty. Such were clearly visible only in the periphery of LC. Very characteristic for the both immunostainings was a rich dendritic radiation that penetrated the surrounding regions. The dorsal subceruleus nucleus, nominated as “subceruleus nucleus, alpha part” by Paxinos and Watson (63), contained a less dense aggregation of labeled neurons, so that the cellular details were discernible. The neuropil contained numerous labeled processes, most of them – dorsoventrally oriented dendrites (Fig. 4a). The DBH immunostaining of the ventral subceruleus nucleus in young animals (Fig. 5a, b) demonstrated large, multipolar, intensively labeled neurons with
Figure 3. LC and dorsal subceruleus nucleus. Medial is to the right. (a) TH-immunoreactivity in a young rat. The dense aggregation of the LC noradrenergic neurons is sharply outlined against the pale background. Numerous long dendrites spread over adjacent territories. Ventral to the LC are located the more loosely arranged neurons of the dorsal subceruleus nucleus, x 100. (b) Detail from (a). Ventral portion of LC. Due to the superimposition of the labeled neurons cellular details might be encountered only in the most peripheral cells. The scant neuropil contains numerous capillaries, x 400. (c) TH-immunoreactivity in an old rat. The “dendritic fan”, surrounding the LC is clearly reduced, x 100. (d) Detail from (c). Border zone between LC and the dorsal subceruleus nucleus. The perikarya of the latter emit short, stumped and thickened dendrites (arrowheads), x 400.
robust primary dendritic trunks and long secondary dendrites.
In the neuropil numerous thin, varicose axons were seen (Fig.
5b).

The histological changes in the TH- and DBH-immunoreactive
populations of the LC and subceruleus nuclei in aged ani-
mals were more pronounced than by the SN and VTA. First, in
all three structures the cell loss was immediately evident (Fig.
3c, 4b, 5c). In LC the cell loss appeared to be most pronounced
in the ventral and lateral portions of the nucleus. The number
of immunoreactive dendrites, radiating in adjacent regions,
was strongly reduced (Fig. 3c, 4b). In the dorsal subceruleus
nucleus, especially in the Paxinos’ “alpha part”, the cell loss
was drastic (compare Fig. 4a and 4b). Heavily deteriorated was
also the ventral subceruleus nucleus (Fig. 5c, d). In addition
to the prominent cell loss, the DBH-immunoreactive neurons
were heavily altered. The perikarya were clearly atrophic, and
not rarely – paler than the neurons in young animals. The rich
dendritic arborization, observed by 3-month-old animals, was
strongly reduced and represented pale, scarcely branched stem
dendrites. The amputated dendrites were a common finding,
and some of the dendritic stumps were cruelly thickened. The
fine, varicose axons practically disappeared (Fig. 5).

Quantitative analysis
The morphometric study indicates that the cross-sectional
areas of the positive neurons in SN decreased significantly in
aging (Fig. 6a), from 1157±25 v/s 829±15 μm² (p<0.001). The
cross-sectional area decreased with 28%. In the same time, the

Figure 4. LC and dorsal subceruleus nucleus. Medial is to the left. Comparison of DBH-immunoreactivity of 3-month-old rat (a)
and 28 month-old-rat (b). x100. The finding is identical with the TH-immunoreactivity shown in Figure 3. In aged rats the neuronal
cell loss is especially evident in the ventral and lateral portions of the LC as well as in the dorsal subceruleus nucleus.
Figure 5. Ventral subceruleus nucleus. Medial is to the right. This is the most ventral part of the subceruleus region, almost reaching the brain surface, often designated as the noradrenergic A5 group of Dahlstrom and Fuxe. (a) DBH-immunoreactivity in a 3-month-old rat. The robust labeled neurons are well discerned also at small magnification in the moderately dense neuronal aggregation, x 100. (b) Detail from (a). The strong immunostaining resembles a Golgi silver impregnation. Often the perikarya emit a short, thick dendritic trunk that gives rise to more slender secondary dendrites. In the neuropil a network of thin, varicose axons is seen, x 400. (c) DBH-immunoreactivity in 28-months-old rat. The drastic cell loss is evident at a first glance, x 100. (d) Detail from (c). The immunolabeled nerve cell bodies are smaller, often – paler. The dendrites are both amputated and thickened (arrow heads). The thin, varicose axons disappear, x 400.
neurons in VTA (Fig. 6b) did not changed their dimensions during aging, 516±12 versus 513±11 µm$^2$ (p>0.05). OD measurements of the TH-positive perikarya indicated that in SN OD decreased significantly with 12% (Fig. 6d), from 95±1.7 to 127±1.4 (p<0.001). In VTA the OD increased significantly in aging (Fig. 6e), from 134±1.4 to 84±1.2 (p<0.001). The OD increased with 19%. The morphometric study demonstrated that the cross-sectional area of the DBH-positive neurons in LC and subceruleus nuclei (Fig. 6c) decreased significantly with 36% in aging from 423±11.1 in 3-month-old rats to 271±7.5 sq µm (p<0.001) in old rats. OD of the DBH-immunoreactive neurons in the subceruleus nuclei (Fig. 6f) of the 3-month-old rats was 51±1.5 (p<0.001) but in aged rats it decreased with 42% to 165±1.2 (p<0.001, n=100).

**Comparison of our results with the data available**

The most characteristic neuropathologic feature of the Parkinson’s disease is the both prominent and selective neuronal cell loss of the pigmented (dopaminergic) neurons of the SN pars compacta (1,9,11,17,32-34,64,65). The observations on the age alterations of SN in humans and experimental animals are less straightforward. Mann and Yates (2,3) demonstrated that a gradual decline takes place from 60 to 90 years in all neuromelanin-containing nuclei, and especially – in SN. The number of pigmented SN neurons (2,3,9,20,66,67) was reported to progressively decrease. In 36 control cases ranged in age from 21 to 91 years, Fearnley and Lees (9) reported a decline of 4.7% per decade, and the total decrease reached 33%. There was a significant sparing of the ventrolateral and ventrointermediate neuronal groups in SN pars compacta – 15% cell loss. Interestingly, the same neuronal groups are most severely affected by idiopathic Parkinson’s disease (1,9,32,65). Faraldi et al (15) supported the hypothesis that the overload of neuromelanin in advanced aging is neurotoxic and contributes to the cell loss in SN. Bannon and Whitty (40) reported up to 75% loss of the dopamine transporter mRNA in older persons, but also noticed that the TH expression in the dopaminergic neurons was less affected. An age-related reduction in the number of SN dopamine transporter-immunoreactive neurons was established also by Ma et al (67). Chu et al (68) reported that in the human SN the number of both Nurri- and TH-immunoreactive neurons was mildly diminished already at middle age, and severely reduced in aged cases. Cabello et al (20) stated that the cell loss in the aged human SN may be compensated by hypertrophy. In contrast to the data discussed above, Kubis et al (41) provided unexpected results. They examined the human TH-immunostained neuronal population in SN, VTA and LC in 21 control subjects who died at ages 44-110 years. They found no statistically significant cell loss of TH-immunopositive neurons.

**Figure 6.** Diagram showing the aging changes in cross-sectional area in µm$^2$ of positive neurons in SN (a) and VTA (b) and OD in arbitrary units of TH-immunopositive neurons in SN (d), and VTA (e). (c) Changes in cross-sectional area and (f) staining intensity of DBH-immunoreactive neurons in subceruleus nuclei (*p<0.001).
The animal studies also provided somewhat diverse results. Irwin et al (39) examined young, intermediate-aged and old squirrel monkeys. Contrary to the obvious functional and neurochemical age-related changes (30-70% loss of dopamine in the neostriatum), the number of TH-immunoreactive cells did not significantly differ among the three age groups. Pakkenberg et al (52) examined the number of pigmented and nonpigmented SN neurons in young and old monkeys. They established that the total number of pigmented neurons was about eight times higher in old animals compared with young ones but the cell loss appeared to be minimal. Siddiqi and Peters (18) and Siddiqi et al (69) reported a SN and VTA cell loss in old rhesus monkey but, interestingly, they stated that the aging affects more significantly the small GABAergic interneurons than the large dopaminergic projection neurons. According to Gerhardt et al (70), there are only modest (15-20%) declines of SN pars compacta TH-positive cells in the course of normal aging. McCormack et al. (43) found no significant differences in the SN cell number in young, middle-aged and old squirrel monkeys, but pointed out also that there is a decrease of TH-immunoreactivity, paralleled with an increase of the neuromelanin content. Collier et al (46) reported that in rhesus monkeys the age-associated morphological changes include decline in the density of TH-positive fibers in striatum, decreased SN soma size, and optical density of TH, but no significant loss of neurons. The differences in the normal aging and neuropathology in the different neuronal groups in the human SN pars compacta is established (9; reviewed in 34) but such a delineation was reported in monkeys only very recently. Kanaan et al (29) investigated young, middle-aged and old rhesus monkeys. They found out that the TH-positive neuron numbers were inversely correlated with advancing age specifically in the ventral tier of the SN pars compacta, not in the dorsal tier or in the VTA. TH intensity decreased throughout the ventral midbrain with increasing age, an effect exacerbated in the ventral tier of SN pars compacta. Uchida et al (42) established that the TH-positive neurons in SN of old dogs appear to be well preserved but they found also cytological alterations typical for the human neuropathology. Tatton et al (36) and Greenwood et al (51) investigated the SN and LC in C57B1 mice aged 8 to 104 weeks, and found that the neuronal loss was due to neuronal death rather than loss of TH-immunoreactivity. The cytoplasmic TH was increased by 63% in 104-week-old mice in comparison to 8-week-old animals (51). McNeill et al (48,49) investigated the SN in young and aged C57B1/6N Nia mice. They found a progressive accumulation of cytoplasmic lipofuscin granules and a markedly reduced dopamine content per cell as determined by histofluorescence. Further, McNeill and Koek (50) investigated six age groups (3-30 months aged) of the same mouse strain, and reported a small decline (11%) in the total number of dopaminergic neurons of the SN with age, a decrease not reaching a statistical significance. Emerich et al (37) postulated that there are no reductions in number, area and length of TH-immunoreactive neurons in A8, A9 and A10 regions of 24-25-month-old rats. Schuligoi et al (38) suggested that the reduction in TH mRNA in the VTA and SN pars compacta in 33-month-old Sprague-Dawley rats is not due to a loss of TH mRNA expressing cells but due to a reduction in the hybridization signal per expressing cell. Himi et al (47) investigated the expression of mRNAs encoding the dopamine transporter and TH in SN of young and aged Fischer 344 rats. They found that dopamine transporter mRNA decreases by 18 months, whilst TH mRNA reduction does not occur until 24 months. In the same rat strain the dopaminergic neurons exhibited increased axonal branching between 6 and 24 months (71). De la Cruz et al (45) report a significant decrease (55%) of TH activity in rats between 12 and 24 months of age, while others established that in old mice the number of SN dopaminergic neurons was decreased by 10% in contrast to TH-immunodensity which was 24±3% higher compared to young animals (24). Recently, Cruz-Muros et al (72) compared the nigrostriatal and mesolimbic (e.g. – the VTA fiber system ascending to limbic forebrain targets) in rats and suggested that both the nigrostriatal and the mesolimbic systems are vulnerable to aging, but in contrast to what occurs in Parkinson’s disease, the mesolimbic system is more vulnerable to aging than the nigrostriatal ones.

Due to the very uneven distribution of the TH-immunoreactive neurons in VTA, we attempted a comparison of the neuronal number of labeled cells only in SN pars compacta. The results were nearly negative (a decrease of 1.5-3.5% by the old rats). Thus, our data are even lower than reported by McNeill and Koek (50), who found a decline of 11%, and declared that it does not reach a statistical significance. The observation of SN pars compacta suggested only mild atrophic changes of the TH-positive neuronal perikarya. The morphometric study, however, demonstrated that the cross sectional area of the positive neurons decreased with 28%. A decreased SN soma size without a cell loss was also reported in monkeys (46). In VTA the cross sectional area did not change. Such a discrepancy between SN and VTA dopaminergic neurons was not observed by Emerich et al (37). They declared that none of
the midbrain dopaminergic groups show area reduction of the TH-positive neurons. It is hard to explain the aging stability of the VTA neurons compared with the SN ones. Our results run counter to the data presented by Cruz-Muros et al (72), who reported findings contrary to ours – the VTA neurons are more vulnerable to aging than the SN cells. In the human brain the cell loss in the aged SN may be compensated by hypertrophy (20) but apparently this phenomenon is absent in the rodent SN. Our data suggest that the OD of TH-positive perikarya in SN decreased with 12% in aged rats. Kanaan et al (29) also found a reduction in the TH activity in SN, especially prominent in the ventral tier of pars compacta in monkeys. Strangely, we found that in VTA the OD increased in old rats with 19% - another difference between the nigrostriatal and mesolimbic neuronal population in the mesencephalon. The VTA dopaminergic neurons are not only resistant to atrophic changes but also display an increased TH activity. A significant increase of the TH activity was reported also by Greenwood et al (51) but this appears to be a compensation of unusually strong cell loss, reported by these authors. Very similar results were reported by Kim et al (24).

With few exceptions (41,73), the investigators agree that with advancing age there is a decrease in number of LC neurons in man (4,34,53-56,58-60,74,75). The data in rodents are somewhat contradictory but probably this is also due to the different strains examined. Goldman and Coleman (76) reported that in LC of Fischer 344 rats of ages from 12 to 32 months there is no neuronal cell loss with advancing age. On the other hand, Sturrock and Rao (77) found a 47% decrease in Nissl-stained LC neurons in 31-month-old mice, compared to 6- and 15-month-old mice. Shores et al (78) found a LC cell loss in Brown-Norway rats but also reported an increase of TH mRNA which may potentially increase noradrenaline synthesis in the remaining neurons. Also, Tatton et al (44) counted immediately adjacent TH-immunoreacted and Nissl-stained sections through the LC in mice at different ages, and concluded that the neuronal loss was due to neuronal death rather than loss of TH immunoreactivity. The present results indicate the occurrence of a significant cell loss in the LC of aged Wistar rats, although not as dramatic as declared by Sturrock and Rao (77). The cell loss was predominantly localized in the ventral and lateral portions of the LC, and the dendritic radiation within surrounding structures was significantly reduced. In the subceruleus nuclei the cell loss was also prominent. In two out of four cases (one of them demonstrated on Fig. 4) the “alpha part” of the dorsal nucleus subceruleus was literally erased. The cell loss in the ventral subceruleus nucleus was also massive and was accompanied by a complete degeneration of the intrinsic plexus of noradrenergic axons. We investigated the cross-sectional area and the OD mainly in the subceruleus nuclei since the superimposition of the labeled neurons in LC obscured the detailed visualization of individual neurons. According to the present results the cross-sectional area in the DBH immunoreactive neurons decreased with 36%. These data were averaged. As seen in Figures 5b and 5d, some neurons undergo even more severe atrophic changes. The OD according to the present data decreased dramatically in aging – with 42%. It is not easily detectable by the observation of the neuronal perikarya but the pale appearance of the dendrites on Figure 5d supports the validity of the OD measurement.

All catecholaminergic neuronal groups presently examined exhibited heavy alterations in the dendritic and axonal arborization by advanced aging. A common finding was the dendritic loss, accompanied by distorted and thickened dendritic stumps. This phenomenon appears to be common throughout the phylogenetic scale, since similar changes were reported also by Cruz-Sanchez et al (79). They applied detailed tracing of Golgi stained neurons in the SN of 20 human control brains ranging from 20 to 93 years of age. Cruz-Sanchez et al (79) encountered distorted profile of the cell body, loss of dendrites and dendritic spines, and swelling and beading of the dendritic branches mainly in the oldest group (70-93 years). Heavy dendritic alterations are also present in aging noncatecholaminergic neuronal populations producing nitric oxide: the cholinergic neurons of the pedunculopontine and laterodorsal tegmental nucleus (80), and the neurons of the dorsolateral column of the periaqueductal gray (81). Finally, von Bohlen und Halbach et al (82) reported a significant decrease in spine number of CA1 dendrites of the aged murine hippocampus that are clearly related to the memory and cognitive impairment. Thus, the obvious alterations of the dendritic tree appear to be a universal phenomenon for different neuronal populations, in different species.

**CONCLUSION**

There were only subtle signs of cell loss in the dopaminergic neuronal population of SN and VTA in aging, whereas the cell loss of the noradrenergic neurons in LC and subceruleus nuclei was evident. Considerable age-related dendritic alterations were observed in all catecholaminergic nuclei. In aging, the cross-sectional area decreased statistically and OD of the neurons in SN diminished with 13%. In VTA, the neuronal...
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cross-sectional area did not change its dimensions, while the OD increased with 19%. In LC and subceruleus nuclei, the cross-sectional area decreased with 36% and the OD of the neurons decreased with 16%. The most resistant to age-related changes catecholaminergic region in the rat is the VTA, followed by the pars compacta of SN, while the rodent LC appears to be a very vulnerable region.

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