Neuron Death in the Substantia Nigra of Weaver Mouse Occurs Late in Development and Is Not Apoptotic

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Weaver is a spontaneous mutation in mice characterized by the postnatal loss of external granule cells in the cerebellum and dopaminergic neurons of the midbrain, especially in the substantia nigra. We have shown previously that natural cell death with the morphology of apoptosis occurs in the substantia nigra of normal rodents during postnatal development. We therefore sought to determine whether the loss of dopaminergic neurons in homozygous weaver mice occurs during the period of natural cell death in the substantia nigra and whether it has the morphology of apoptosis. We have found, using a silver stain technique, that although apoptotic cell death does occur early postnatally in homozygous weaver substantia nigra, it also does so with equal magnitude in wild-type and heterozygous weaver littermates. Unique to homozygous weavers is the occurrence of degenerating neurons in the nigra that are not apoptotic. These degenerating neurons are observed at postnatal day 7, and they are most abundant on postnatal days 24–25. The nonapoptotic nature of this cell death is confirmed by negative in situ end labeling of nuclear DNA fragmentation and by ultrastructural analysis. Ultrastructural studies reveal irregular chromatin aggregates in the nucleus, as well as marked cytoplasmic changes, including the formation of vacuoles and distinctive stacks of dilated cisternae of endoplasmic reticulum. We interpret these changes as indicative of either a variant morphology of programmed cell death or a pathological degenerative process mediated by an as yet unknown mechanism related to the recently described mutation in the GIRK2 potassium channel.

Key words: weaver; apoptosis; programmed cell death; substantia nigra; potassium channel; development; ultrastructure

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earliest abnormalities observed in the wv/wv nigrostriatal system (Roffler-Tarlov et al., 1990), and that such an abnormality may impair the maintenance of target-derived support, we hypothesized that the wv mutation may be associated with augmented apoptotic cell death in SN. We have found, however, that although apoptosis does occur in postnatal wv/wv SN, it occurs with equal magnitude in the +/+wv and +/+ mice. Unique to wv/wv, SN is the later occurrence of neuronal degeneration with a nonapoptotic morphology.

MATERIALS AND METHODS

Animals: All mice were offspring of either +/+wv or +/+ breeding pairs supplied by the Jackson Laboratory (Bar Harbor, ME) and bred and maintained in the laboratory colony (SKR-T, SH). All mice were on a C57BL/6JLe-Am × CBA/CaImF hybrid background. Identification of phenotype as +/+wv, +/+v, or wv/wv was based on examination of behavior and postmortem examination of cerebellum, as described previously (Roffler-Tarlov and Turey, 1982). The day of birth was defined as P0.

Silver staining: To demonstrate cell death and to characterize its morphology, consecutive sections from the SN and striatum were silver-stained (Gallyas et al., 1980). At selected postnatal ages, mice were anesthetized with Nembutal and then perfused through the left ventricle with 0.9% saline at 4°C anesthetized with Nembutal and then perfused through the left ventricle with 0.9% saline at 4°C, followed by perfusion with 4% paraformaldehyde in 0.1 M phosphate buffer, pH 7.2, for 20 min. Brains were carefully removed from the skull and post-fixed in the same fixative for at least 1 week. Brains were then cryoprotected by immersion in 20% sucrose/4% paraformaldehyde/0.1 M phosphate buffer overnight. Blocks were rapidly frozen in 2-methylbutane chilled on dry ice and sectioned in a cryostat at 30 µm. Representative sections through the entire SN and striatum were obtained and processed for silver staining. Sections were maintained in serial order and process free-floating in custom-made plastic grids with nylon mesh bottoms. Sections were collected into cold fixative, washed 3 times in distilled water, and then immersed in pretreating solution (equal volumes 9% NaOH and 1.2% NaNO3) for 5 min twice. They were then immersed in impregnating solution (60 ml of 9% NaOH; 40 ml of 16% NH4NO3; 0.5 ml of 50% AgNO3) for 10 min. Sections were then washed 3 times in washing solution (1 ml of 1.2% NaNO3, added to 100 ml of a solution containing 5 gm anhydrous Na2CO3, 300 ml of 95% ethanol, brought to 1 l with distilled water) followed by immersion in developing solution (1 ml of 1.2% NH4NO3 and 100 ml of a solution consisting of 0.5 gm citric acid in 15 ml of 37% formalin, 100 ml of 95% ethanol, 700 ml of water brought to pH 5.8–6.1 with 9% NaOH, and finally brought to 1 l with water). Sections were kept in developing solution for ≥1 min. Sections were then processed to subbed slides, dried, and immersed in 0.5% acetic acid 3 times for 10 min each. Sections were then dehydrated through alcohols, cleared in xylene, and coverslipped under Permount.

Quantitative morphological analysis of silver-stained sections: At P7, all available SN silver-stained sections were analyzed (mean number of sections per animal = 133.3). For older mice, the SN was divided into three caudorostral regions: (1) a caudal region containing the medial lenticulus and the interpeduncular nucleus; (2) a central region containing the medial terminal (MT) nucleus of the accessory optic tract; and (3) a region anterior to MT, containing a well defined SNpc. Each of these regions was sampled with 0.9% saline at 4°C, followed by perfusion with 4% paraformaldehyde in 0.1 M phosphate buffer and post-fixed for 1 week. Sections through the SN were cut on a Vibratome at 100 µm thick and collected into 0.1 M Sorensen’s buffer and post-fixed for 1 h. Sections through the SN were cut on a vibratome at 100 µm and collected into 0.1 M Sorensen’s buffer. Sections were stained in 1% OsO4/0.1 M Sorensen’s for 60 min, and then washed. After dehydration in ascending concentrations of alcohols, sections were embedded in Epoxy. A plastic section was thus obtained. The sections were then trimmed to include the medial portion of the SNpc, mounted on an Epon stub, and cut at 10 µm for examination under phase contrast. Sections of interest were then reembedded, thin-sectioned, and examined with a JEOL 1200EX electron microscope.
Figure 1. **Top.** A, An example of an apoptotic profile, demonstrated by silver staining, in the SNpc of a \( w^v/w^v \) mouse at P7 (arrow). Characteristic of apoptosis are the two darkly silver-impregnated, rounded chromatin clumps. The chromatin clumps are surrounded by a moderately silver-impregnated cellular profile. The dark silver impregnation of the apoptotic profile contrasts with the golden hue of adjacent, normal neurons. B, An example of an apoptotic profile in the SNpc of a \(+/w^v\) mouse at P14 (arrow). This particular profile is somewhat smaller than the profile shown in A, as it appears to be undergoing some breakup (there is scattered adjacent silver-stained material), but it is otherwise similar in its basic features: there are darkly impregnated, rounded chromatin clumps surrounded by a less intensely silver-impregnated, rounded cellular profile. Scale bar, 10 \( \mu m \).

Figure 3. **Bottom.** An example of a silver-stained, nonapoptotic neuronal profile demonstrated in the SNpc of a \( w^v/w^v \) mouse at P14. This silver-impregnated profile contrasts with the apoptotic profiles demonstrated in Figure 1 in several respects. There are no distinct chromatin clumps; rather, the entire nucleus is darkly impregnated. Rather than being rounded and faintly silver-impregnated, the surrounding cytoplasm retains a polygonal, neuronal appearance and is heavily silver-impregnated. The appearance of long and heavily impregnated neuronal processes, with bifurcations, would be distinctly unusual for an apoptotic neuronal profile, and yet was observed universally in these cells. Scale bar, 10 \( \mu m \).
profiles at P14 in the +/+ group; at no other time was there an apparent genotype effect. We cannot discount the possibility that the apparent difference at P14 was attributable to other sources of variability related to tissue processing. At no time was there a significant difference between +/wv and wv/wv, and there was no significant interaction between age and genotype. ANOVA did reveal a highly robust effect of age ($p < 0.0001$) on the prevalence of apoptotic cells, with more cells being present at P14 ($p < 0.05$, Newman–Keuls post hoc analysis). In addition, the adult group differed from all others in the very small number of apoptotic profiles ($p < 0.05$, post hoc). Both of these findings are similar to what we have reported previously for the time course of apoptotic cell death in rat SN (Janec and Burke, 1993). We conclude that the magnitude and time course of apoptotic cell death postnatally in SN in wv/wv is not significantly different from +/wv and +/+ littermate controls.

We identified a small but significant difference between wv/wv and littermates in the size of apoptotic profiles demonstrated by silver staining. The mean area of these profiles was $34.3 \pm 0.9 \, \mu m^2$ in wv/wv ($n = 88$ profiles), whereas in +/wv and +/+ it was $30.1 \pm 0.6$ ($n = 88$) and $31.5 \pm 0.7$ ($n = 88$), respectively. This small increase in the size of the wv/wv profiles would not influence the analysis performed in Figure 2, which was based on a stereological assessment. If a traditional Abercrombie correction (Abercrombie, 1946) were applied to the data shown in Figure 2, the only change in the analysis would be that +/+ would not be less than wv/wv and +/wv, and +/wv would be only marginally ($p < 0.04$) greater than +/+ and wv/wv. This analysis underscores our interpretation that the small differences in counts among the genotypes at P14 are unlikely to be relevant to the wv/wv phenotype.

Examination of silver-impregnated sections revealed that darkly impregnated neuronal profiles, predominantly in SNpc (Fig. 3), were unique to wv/wv mice at later postnatal ages. These impregnated profiles could be distinguished from apoptotic profiles in several respects. (1) The entire nucleus was darkly impregnated without the appearance of characteristic discrete, rounded chromatin clumps. (2) The cytoplasm was not shrunk and rounded, as is typical for apoptosis, but retained a polygonal, neuronal appearance, often with clear visualization of dendrites, including primary and secondary bifurcations. (3) The cytoplasm typically contained a dense, punctate silver impregnation. The time course of appearance of these degenerating neurons differed from that of apoptotic cell death, as shown in Figure 4. A few of these cells were apparent in wv/wv at P7. A maximum number was observed at P24–P25. By adulthood, very few were present. In +/+ animals, these cells were never observed. In +/wv animals, few darkly impregnated neuronal profiles were observed in SN at P24–P25 and in adulthood.

The regional distribution of these nonapoptotic cells was examined in four wv/wv mice at P24–P25. These nonapoptotic degenerating cells were most numerous in the medial region of the SNpc (the “dense zone”), but they were also found throughout the medial-to-lateral extent of the SNpc, and in the pars lateralis. In the rostrocaudal dimension, 50.5% of the nonapoptotic cells were located in the rostral-most SN planes. Another 37.8% were located in the caudal-most planes, whereas only 11.8% were located in the central region. Although we did not count the number of TH-positive cells in these regions, and therefore did not normalize counts of dying cells for the number of TH-positive neurons present, the number of dying cells appeared to parallel the density of TH-positive neurons in these regions. The vast majority (99%)
of degenerating neurons was localized in the SNpc; only 1% were observed in SN pars reticulata (SNpr). Those in SNpr were found exclusively in caudal SN planes where TH-positive dopaminergic neurons of the ventral tier are known to reside. In the midbrain, nonapoptotic cells were not localized exclusively in the SN; a number were also observed in the ventral tegmental area. Other nuclei of the basal ganglia with important anatomical relationships with the SN, including the striatum and the globus pallidus, were examined in wv/wv animals at different postnatal ages, but nonapoptotic degenerating cells were not found.

To further characterize the nature of the cell death in wv/wv SN at P24–P25, we performed an in situ 3′-end labeling procedure to identify the presence of fragmented nuclear DNA, a characteristic of apoptosis. As shown in Figure 5, and consistent with our assessment of the morphology revealed by silver stain, sections throughout the weavers’ midbrains were without free 3′ ends, indicating a nonapoptotic form of cell death. This negative result was obtained in spite of our ability to demonstrate positive 3′-end labeling in wv/wv cerebellum, where apoptosis does occur (Harrison and Roffler-Tarlov, 1995).

To attempt to define the dopaminergic phenotype of dying cells in SNpc, we performed TH immunostaining with Nissl counterstain. We have shown previously that this procedure successfully detects apoptotic cell death in these neurons, both during natural cell death (our unpublished observation) and induced cell death (Macaya et al., 1994; Oo et al., 1995). Presumably, therefore, the apoptotic death process is rapid enough to permit demonstration of retained cytoplasmic phenotypic markers. Accordingly, 2600

Figure 4. Quantitative analysis of the number of silver-stained nonapoptotic profiles in wv/wv and littermate controls at representative developmental ages. The same animals analyzed for apoptotic profiles, shown in Figure 2, have been analyzed here. Nonapoptotic profiles first appeared in wv/wv at P7 and increased in number to become most numerous at P24–P25. These nonapoptotic profiles appeared almost exclusively in wv/wv. None was observed in +/+ animals, and only a few were observed in +/wv at P24–P25 and in adulthood.

TH-positive neurons were examined on 14 sections obtained from the midbrain of a single P24 wv/wv mouse. No TH-positive apoptotic cells were observed. In regions known to contain degenerating cells, identified on silver-stained adjacent sections, abnormal Nissl staining was observed among TH-negative neurons of the SNpc (Fig. 6). These cells contained pleomorphic, basophilic chromatin aggregates, and a vacuolated “foamy” appearance of the cytoplasm. Thus, by this histological approach, as with end labeling, we were unable to demonstrate apoptotic cell death in SNpc, but again observed morphological features of nonapoptotic degeneration.

**Electron microscopy**

The nonapoptotic morphology of the neuron death in wv/wv at P24 was further confirmed by analysis at an ultrastructural level. Examples of degenerating cells observed in wv/wv are shown at the light microscope level in Figure 7 and at the ultrastructural level in Figure 8, A and B. Morphological alterations are observed in both the nucleus and the cytoplasm. The nuclei show an overall increase in electron density, the nuclear membrane is deeply invaginated, and there is dilatation of the perinuclear cisternal space (Fig. 8A,B). Moreover, the nuclear chromatin forms abnormal, irregularly shaped aggregates. These aggregates differ from those typically observed in apoptosis in that they are neither apposed to the nuclear membrane nor rounded with sharply demarcated edges (Sloviter et al., 1993; Macaya et al., 1994) (see Fig. 9C). The cytoplasm of these cells also shows an overall increase in electron density. In some of the cells, exemplified by the profile in Figure 8A, there were numerous, aggregated stacks of endoplasmic reticulum (ER). The cisternae of the ER were dilated, and the regions between cisternae were packed with aggregates of ribosomes (Fig. 8A, inset). Other cells, as shown in Figure 8B, displayed extensive vacuole formation. Some of the vacuoles were continuous with a dilated perinuclear space. As shown in Figure 8B (inset), the vacuoles contained a loosely aggregated matrix, and they were surrounded by densely packed ribosomes. These abnormal features are readily contrasted with adjacent normal cells in wv/wv SN (Fig. 9A) and normal cells in +/+ SN (Fig. 9B). These features can also be contrasted with an instance of apoptotic cell death in a +/+ animal at P24 (Fig. 9C). This cell shows features characteristic of the late stages of apoptotic cell death: formation of rounded, well demarcated, electron-dense chromatin clumps and an increase in the electron density of the cytoplasm, with relative preservation of mitochondria. Cells with the abnormal ultrastructural features shown in Figure 8, A and B, were not observed in +/+ animals.

We rarely observed abnormal cells in wv/wv mice, which we believe may represent early stages of the degenerative process illustrated in Figure 8, A and B. Such cells showed milder, but similar abnormalities. The cell shown in Figure 10 contains abnormal, irregular chromatin clumps in the center of the nucleus and adjacent to the nuclear membrane. There also is a slight increase in the electron density of the cytoplasm. Within the cytoplasm, rare vacuoles are seen, as well as aggregates of ER, associated with dense accumulations of ribosomes.

The neuronal nature of the degenerating cells in wv/wv midbrains could be confirmed in some instances by the identification of synaptic densities, as shown in Figure 11.

**DISCUSSION**

The morphological appearance of apoptosis is so often observed in the course of programmed cell death that many investigators
Figure 5.  Top. In situ end labeling in a wv/wv mouse. A, TH immunostain of a P24 wv/wv mouse, demonstrating the medial portion of the SNpc. B, In situ end labeling of a region comparable to that shown in A, in a P24 wv/wv mouse. There is an absence of blue X-gal staining over the SNpc, indicating a negative reaction for free 3' ends. Scale bar, 100 μm.

Figure 6.  Bottom. TH immunostain and Nissl counterstain of an SN section from a P24 wv/wv mouse. A normal-appearing, peroxidase-positive (brown) neuron is indicated (solid arrow). Note that the nucleus of this cell contains two stained heterochromatin clumps, characteristic of normal mouse cells (Moser et al., 1975). Adjacent to the TH-positive neurons is a TH-negative cell (open arrow) with a vacuolated, foamy-appearing cytoplasm, and an abnormal nucleus containing multiple, intensely basophilic, pleiomorphic chromatin aggregates of varying size. These aggregates differ from those observed in apoptosis in that they are greater in number, more diverse in their shape (rather than being simply rounded), and they are smaller. The chromatin aggregates in this micrograph can be compared with those typical for apoptosis shown at the same magnification in Figure 1.
use the terms synonymously. However, we will use the term apoptosis in its more restricted sense, to refer to one particular morphology of programmed cell death. Therefore, by interpreting the morphology of neuron degeneration in the SN to be “non-apoptotic,” we do not mean to imply that this is not a form of programmed cell death; this remains a possibility, as discussed below. Our use of the term apoptosis to refer strictly to a morphological concept is based on the original use of the term (Kerr et al., 1972) and recognition of the fact that other, nonapoptotic morphologies are observed in programmed cell death (Clarke, 1990; Schwartz et al., 1993).

The strictest morphological criteria for identification of apoptosis in brain depend on ultrastructural observations. Distinctive features are observed both in the nucleus and cytoplasm, but those in the nucleus occur earlier and are most distinctive. The most characteristic alteration of nuclear morphology is the formation of sharply delineated, homogeneous, electron-dense chromatin masses (Kerr et al., 1995). The mere presence of intranuclear chromatin aggregates is not sufficient to identify apoptosis; irregular aggregates with poorly defined edges have been clearly identified in necrotic cell death (Olney, 1969; Kerr et al., 1995). The cytoplasmic features of apoptosis are also distinctive. Early, there is an overall condensation of the cell with an increase in its electron density, with preservation of intracellular organelles (such as mitochondria) and both nuclear and cellular membranes (Kerr et al., 1995). Although not as definitive as ultrastructure, several methods at the light microscope level can be highly suggestive (Clarke and Oppenheim, 1995). Nissl stains have been used to identify the distinct, round chromatin clumps characteristic of apoptosis, and in many instances the ultrastructural features have been confirmed (Cunningham, 1982; Williams and Rakic, 1988; Ferrer et al., 1990; Sloviter et al., 1993; Macaya et al., 1994). Suppressed silver-staining methods also have been used to identify intensely argyrophilic, distinct, round chromatin clumps (Yamamoto et al., 1986; Janec and Burke, 1993; Macaya et al., 1994; Mitchell et al., 1994), and confirmed by ultrastructural analysis. Labeling of free 3′ ends (Gavrieli et al., 1992) can also be highly suggestive of apoptotic cell death at the light microscope level, if it is used with strict attention to the morphological appearance of apoptosis (Oo et al., 1995).

We have used all of these approaches to identify the character of the neuron death in the SN of wv/wv mice. Silver staining clearly demonstrated typical apoptotic morphology early postnatally in the wv/wv SN, as anticipated. However, identical profiles were also observed within the SN of the +/+/ and +/-wv mice, and quantitative analysis of these profiles failed to reveal a consistent difference between wv/wv and the other genotypes. In all three genotypes, a highly significant effect of postnatal age on the prevalence of apoptosis was observed, with many more profiles observed at P14 and very few in adulthood. This temporal pattern...

Figure 7. Numerous dark, osmium-stained abnormal cells are observed in wv/wv SN. A, An osmium-stained plastic-embedded 10 μm section of wv/wv SNpc at P24, shown in phase contrast. Shown is the medial portion of SNpc (dorsal is indicated by an up arrow, lateral by a left arrow, lower right corner). Compared with the same region of +/- SNpc shown in B, there are fewer neurons in wv/wv. In wv/wv, there are numerous dark, osmium-stained degenerating neurons (indicated by black arrows within white arrows). The neurons labeled with the numbers 1 and 2 are shown at higher magnification in Figure 8. Each of the dark neurons in wv/wv is surrounded by a bright rim of high illumination in phase. B, SNpc of +/- at P24. Rare cells in SNpc were stained (black arrow within white arrow). A representative example is shown in Figure 9C. Scale bar, 100 μm.
Figure 8. Ultrastructural appearance of degenerating cells in \textit{wv/wv} SNpc at P24. \textit{A}, This cell is labeled with the number 1 in Figure 7. The nucleus has become invaginated and multilobular. There has been an increase in the intramembranous space of the nuclear membrane. Within the nucleus, there are multiple aggregates of electron-dense chromatin material, which are irregular in size and shape. In the cytoplasm, there are vacuoles and layered structures consisting of dilated cisternae of ER alternating with regions densely packed with ribosomes. An example of these stacks is enclosed within the \textit{white box} and shown at higher power in the \textit{inset}. Within the \textit{inset}, a single representative example of a mitochondrion is marked with an \textit{arrow}. In general, mitochondria were of normal size and shape, but with increased electron density and poorly defined cristae. In no case were mitochondria dilated. \textit{B}, This cell is labeled with the number 2 in Figure 7. The nuclear changes are like those observed in cell 1. The predominant cytoplasmic change in cell 2 is the formation of vacuoles. They contained an irregular, sparse electron-dense matrix and were bounded by numerous ribosomes. An example is enclosed in the \textit{white box} and shown at higher magnification in the \textit{inset}. Scale bars: 2 \textmu m in \textit{A} and \textit{B}; 200 nm in the \textit{insets}. 
Figure 9. Ultrastructure of normal cells in wv/wv and +/+ mice, and a single apoptotic cell in a +/+ mouse. A, Ultrastructure of a normal neuron in SNpc of wv/wv. This normal-appearing cell contrasts with the pathological features observed in degenerating cells in wv/wv, as shown in Figure 8. The nucleus shows multiple, small clumps of heterochromatin, which is characteristic of mouse cells (Moser et al., 1975). Within the cytoplasm, normal-appearing rough ER and mitochondria are observed. B, Ultrastructure of a normal neuron in SNpc of +/+ mouse. C, Abnormal, dense cell in SNpc of a +/+ mouse. This cell shows an overall increase in electron density. The nucleus contains a large, rounded, and distinctly bounded chromatin clump. Numerous intact mitochondria are observed in the cytoplasm. We consider these morphological features to be typical of late apoptotic cell death. Thus, this cell confirms the presence of apoptotic profiles observed on silver staining in P24 +/+ mice. Scale bar, 2 μm in all panels. Note lower magnification in C.
is quite similar to what we had previously observed for rat, with one peak in the prevalence of apoptotic cells at P14 (Janec and Burke, 1993). Thus, \( w^v/w^v \) does not appear to be associated with a change in the magnitude or timing of a natural cell death event in the SN.

Silver impregnation identified degenerating neurons in \( w^v/w^v \) SN that were not apoptotic by the light microscopic or ultrastructural criteria outlined above. The cellular pattern of silver impregnation observed in \( w^v/w^v \) SN (that of uniform, heavy impregnation of the nucleus and extensive impregnation of the cytoplasm) has been observed in a wide variety of models of induced neuronal degeneration, and does not implicate a single, specific mechanism. We have observed this pattern after direct injection of the neurotoxin 6-hydroxydopamine into the SN (Jeon et al., 1995), and it has been observed in the SN after treatment of mice with the selective dopamine neurotoxin \( N \)-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) (Jackson-Lewis et al., 1995). The ultrastructural features observed in the \( w^v/w^v \) cells do not correspond to the features described for these neurotoxins. 6-hydroxydopamine induces a watery appearance of both the cytoplasm and the nucleus (Ichitan et al., 1994). Although MPTP induces cytoplasmic vacuoles, it does not induce nuclear chromatin aggregates (Mizukawa et al., 1990). MPTP induces mitochondrial dilatation, which does not occur in \( w^v/w^v \) cells. The changes in \( w^v/w^v \) also do not resemble those induced acutely by excitotoxins, which consist of cellular swelling and bloating of mitochondria (Olney, 1969).

Although electron-dense “dark” neurons have been demonstrated to be artifacts attributable to poor fixation (Cammermeyer, 1961), they do not demonstrate the ER stacks we have observed. The dark neuron phenomenon usually reveals a bland, electron-dense appearance, whereas in the \( w^v/w^v \) SN, we observe a spectrum of morphological changes, suggesting an evolving pathological process. Furthermore, morphology such as that observed in \( w^v/w^v \) SN was not seen in \( +/+ \) controls.

The possibility that the changes we have observed in the \( w^v/w^v \) neurons may represent a late stage in the evolution of the apoptotic process seems unlikely, because in brain, apoptotic cell death is a brief and asynchronous event, so that, at any time, its full morphological spectrum should be evident. Apoptotic profiles in the mice we examined were most abundant at P14, whereas the nonapoptotic morphology was most prevalent at P24–P25. In addition, the morphological features that we observed in the
wv/wv SN have not been described in other paradigms of apoptosis.

One possible explanation for the morphology that we have observed in wv/wv SN is that it represents an alternate, nonapoptotic form of developmental cell death. Clark has pointed out that a variety of morphologies can be identified during developmental cell death (Clark, 1990). His Type 1 category is characterized by nuclear chromatin condensation and corresponds to apoptosis. His Type 3b (“cytoplasmic”) resembles the morphology we have observed in wv/wv SN in a number of respects: (1) there is dilatation of the ER and the intranuclear membrane space; (2) some of the vacuoles appear continuous with ER; and (3) there is an abundance of ribosomes. However, in Type 3b, the nuclear changes, especially early in the process, are minimal, whereas in wv/wv we observe clear nuclear chromatin abnormalities even at early stages. Therefore, if this morphology represents an example of Type 3b developmental cell death, it must be considered a variant. If this morphology is a variant form of augmented developmental cell death, we must consider why this variant is observed in wv/wv SN, rather than simply an augmented number of cells with an apoptotic morphology. In some developmental settings, the morphology of cell death is related to the maturity of the cell at the time of death (Pilar and Landmesser, 1976; Cunningham, 1982), and maturity may be a factor here. The difference in morphology of the cell death between SN and cerebellum (where it is apoptotic) (Smeyne and Goldowitz, 1989; Migheli et al., 1995; Wullner et al., 1995) may also be related to the maturity of the cells when they die in SN. In SN, how the possibility of augmented developmental cell death relates to the recently described mutation in the GIRK2 potassium channel in weavers (Patil et al., 1995) is unknown.

An alternative interpretation of our results is that cell death in SN represents a pathological morphology attributable to cell injury mediated by the GIRK2 mutation. Recent functional studies of the effect of the weaver mutation on the GIRK2 channel have suggested that it leads to a loss of selectivity for the K+ ion (Slesinger et al., 1996) and, in cerebellar granule cells, this leads to a constitutive, depolarizing inward Na+ current (Kofuji et al., 1996). If the mutation results in an aberrant sodium current that depolarizes the cell, this may lead to a chronic, excitotoxic form of neuron death that may differ from that described for acute excitotoxicity (Olney, 1969). Whether the cell death in wv/wv SN is augmented developmental cell death, as considered above, or strictly a pathological degenerative process, it may in either case involve the mechanisms of programmed cell death, defined in the broadest sense as a genetically regulated death process.

There is an apparent discrepancy between our observation that the largest number of nonapoptotic profiles appeared at P24–P25 and the previous observation that most of the TH-positive neurons have been lost by P21 in wv/wv animals (Roffler-Tarlov et al., 1996). One possible explanation for this discrepancy is that the TH-positive neurons are dysfunctional and have lost their expression of TH before the onset of degeneration, as identified by silver staining and ultrastructure. In favor of this possibility is our observation that abnormal cells could be demonstrated on Nissl staining and ultrastructure. In favor of this possibility is our observation that abnormal cells could be demonstrated on Nissl staining and ultrastructure. In favor of this possibility is our observation that abnormal cells could be demonstrated on Nissl staining and ultrastructure. In favor of this possibility is our observation that abnormal cells could be demonstrated on Nissl staining and ultrastructure. In favor of this possibility is our observation that abnormal cells could be demonstrated on Nissl staining and ultrastructure. In favor of this possibility is our observation that abnormal cells could be demonstrated on Nissl staining and ultrastructure.

Whatever the pathogenesis of dopamine neuron degeneration in wv/wv, it is a late developmental event. At this developmental time, these neurons are postmitotic, postmigratory, and they probably are also beyond the period of target dependence, based on target ablation studies in developing rat (our unpublished observations). The degeneration in SN may be related to the late development of a system property, such as the arrival of synapticity active afferent input. In rat, the developmental period showing the greatest rate of increase in number of synaptic profiles per cell in the SNpc is P15–P30 (Lauder and Bloom, 1975), which corresponds to the period of maximal cell death in SNpc in wv/wv.

REFERENCES


