Neuronal Cell Death in Alzheimer’s Disease Correlates with apoE Uptake and Intracellular Aβ Stabilization

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Abstract

The brains of individuals with Alzheimer’s disease (AD) are characterized by extracellular deposition of β-amyloid protein (Aβ), intracellular neurofibrillary tangles, and loss of neurons. To study molecular markers associated with dying cells in the AD brain, in situ DNA labeling techniques were used to visualize cells with DNA fragmentation. We observed that intracellular accumulation of apolipoprotein E (apoE) is correlated with the detection of intracellular Aβ-like immunoreactivity within the same cytoplasmic granules, suggesting that uptake of lipids may have stabilized the hydrophobic Aβ protein within the cell. These apoE-containing neurons also exhibit high expression of a cell surface receptor, gp330, which is known to bind apoE. Cells containing significant nuclear DNA fragmentation express numerous neuronal cell ghosts. Based on our in situ analysis of nuclear DNA fragmentation, we conclude that neuronal cell death likely occurs before the extracellular deposition of Aβ in AD brains. (J. Clin. Invest. 1997, 100:310–320.) Key words: Alzheimer’s disease • β-amyloid protein • apolipoprotein E • glycoprotein 330 • apoptotic cell death

Introduction

A defining histopathological feature of Alzheimer’s disease (AD)1 is the presence of diffuse and neuritic plaques localized to specific brain regions, particularly the temporal lobe (1). The principal constituent of these extracellular deposits is the 39–43 amino acid β-amyloid peptide (Aβ) (2, 3), which is derived by proteolytic processing from the β-amyloid precursor protein (APP). The shorter Aβ fragments, particularly Aβ1–40, appear to be the predominant soluble species, while Aβ1–42 is the form initially deposited in diffuse plaques (4). The generation of Aβ from APP has been shown to be a normal processing event (5–7), and Aβ can be detected in the cerebrospinal fluid and plasma of both normal and AD patients (6–8). The biochemical process leading to Aβ accumulation in the diseased brain, however, is unknown.

The neurological deficits that occur in AD likely stem from neuronal cell dysfunction and death (9–11). The factors and circumstances that induce the dysfunction and death of neurons in the AD brain remain to be elucidated. Although there has been some controversy as to whether plaque accumulation correlates with dementia, recent evidence indicates that the Aβ load in AD brains can be correlated with the severity of dementia (12). In vitro and in vivo investigations using transgenic mice indicate that the Aβ peptide is toxic to neurons, and that cell death proceeds via an apoptotic pathway (13–16). Nevertheless, many questions regarding the pathobiological effects of Aβ in the human brain remain unresolved (5–7). For instance, is Aβ neurotoxic in its natural environment, and if so how? Where and how is Aβ generated, and how does it assume an extracellular location? Does Aβ exert its function after extracellular deposition?

We have used in situ DNA labeling techniques (17, 18) in an attempt to identify pathological markers that are associated with damaged cells in the AD brain. Our results indicate that there is substantial detection of damaged cells in the AD brain after terminal deoxynucleotidyl transferase–biotin dUTP nick-end labeling (TUNEL) staining, a finding that correlates well with previous studies (19–21). Importantly, we find that many of these damaged neuronal cells contain Aβ-like immunoreactivity and apolipoprotein E (apoE) immunoreactivity. Although apoE immunoreactivity has been demonstrated previously in neurons (22, 23), we believe our results are the first to document it in damaged cells. In addition, a receptor (gp330) that is known to bind apoE is also found selectively elevated in these damaged cells. The sum of these findings suggests that gp330 receptor–mediated uptake of apoE may result in the intracellular stabilization of Aβ, and may contribute to the demise of the cell.

Methods

Tissues and histology. Tissues from neuropathologically defined AD cases, which fulfilled the requirements of the Consortium to Establish a Registry for the Diagnosis of Alzheimer’s Disease (24), were used. Comparable brain regions from AD and control cases were matched for age and postmortem delay as closely as possible, and were processed in parallel for all staining procedures. Sections of paraffin-embedded tissues were cut at 5-μm thickness, and were stained with hematoxylin and eosin (H&E), Bielschowsky’s silver stain, thioflavin,
periodic acid Schiff (PAS), or processed for TUNEL or immunohistochemistry (15, 16).

**TUNEL.** Several enzymatic methods have been used to detect apoptosis at the single-cell level by capitalizing on the fragmented nature of the DNA to add labeled or conjugated nucleotides to the termini. These techniques either involve tailing DNA by terminal deoxynucleotidyl transferase (TdT), referred to as the TUNEL method (17), or nick-translation with DNA polymerase I or the Klenow fragment, referred to as in situ nick-translation (25). The tailing method has been demonstrated to be superior to nick-translation for the detection of apoptotic cells; in contrast, necrotic cells in which the DNA is randomly fragmented were labeled preferentially by nick-translation (26). Although caution must be applied in interpreting the TUNEL assay (18), this assay in combination with other morphologic or biochemical criteria could allow for discrimination of apoptotic cells from necrotic cells.

Paraffin sections were dewaxed, rehydrated, and digested with proteinase K (20 μg/ml). Endogenous peroxidase was quenched by treatment in 3% hydrogen peroxide. Sections were then rinsed in water and labeled at 37°C for 30 min with TdT (0.5 U/μl) in a cocktail consisting of 25 mM Tris, pH 6.6, 200 mM potassium cacodylate, 1 mM CoCl₂, 0.25 mg/ml bovine serum albumin, and 2 μM biotin-dUTP. Sections were rinsed and incubated with avidin and biotinylated horseradish peroxidase (Vector Laboratories Inc., Burlingame, CA), rinsed in water, and then stained with either dianaminobenzidine or with True Blue (Kirkegaard and Perry Laboratories, Gaithersburg, MD).

**Immunohistochemical staining.** For immunohistochemical analysis of Aβ, the mouse monoclonal antibody 4G8, which is directed against an epitope consisting of amino acids 17–24 of Aβ and recognizes diffuse and senile plaques in AD brains (27), or an antibody to amino acids 1–28 of Aβ (15) was used. For apoE staining, the monoclonal antibody 3H1 and a rabbit polyclonal antibody (28) were used. Characterization of the rabbit polyclonal antibody R239 to gp330 (29), monoclonal antibody 8G1 (30), polyclonal antibody R777 to LDL receptor–related protein (LRP) (31), monoclonal antibody C7 to LDL receptor (LDLR) (32), and rabbit polyclonal antibody R2623 to VLDL receptor (VLDLR) (33) have been described previously. Paraffin sections were dewaxed and rehydrated. Sections were then rinsed in PBS, blocked for 1 h with 2% blocking serum, and incubated overnight with the appropriate antibody. This was followed by a brief rinse, incubation with a secondary antibody, PBS rinse, and incubation with avidin and biotinylated horseradish peroxidase (Vector Laboratories). All immunostained sections were developed with dianaminobenzidine substrate.

**Results**

**Analysis of cellular DNA fragmentation in AD brain.** We have analyzed brain specimens from AD patients (who were at various stages in the progression of disease) based on clinical history, total brain weight at autopsy, and extracellular Aβ deposition on immunohistopathological analysis. Without exception, brain tissue sections from all seven AD patients showed extensive cellular DNA fragmentation as revealed by TUNEL. In contrast, sparse or no TUNEL-positive cells were detected in five of six brains from control individuals (Table I). The exceptional control sample in which TUNEL-positive cells were detected was from a 101-yr-old man who had a clinical history of depression. Since results obtained with the TUNEL assay are a reflection of the extent of cellular DNA damage, and by inference a measure of cell death, our results indicate that there are considerable numbers of damaged or dying cells in the AD brain tissue specimens compared to control samples. Since adjoining regions of the brain frequently have markedly different numbers of TUNEL-positive cells, attempts to quantitate the overall extent of cell death in each of the patients have proven difficult, and may not be particularly informative. This finding correlates well with several prior studies (19–21).

Regardless of the individual, all regions of the brain with extensive TUNEL staining invariably showed surrounding extracellular Aβ immunoreactivity. Characteristic differences, however, were observed for most individuals. A representative sample shown in Fig. 1 a, shows the deep cortical layers of the superior temporal gyrus of AD patient 1250, in which TUNEL-positive cells of varying intensities are interspersed among numerous TUNEL-negative cells. It is important to note the different intensities of TUNEL staining, which likely reflect various levels of cellular DNA fragmentation. Furthermore, in accord with only a minority of the cells containing detectable DNA damage, weak extracellular Aβ immunoreactivity with the appearance of diffuse plaques was localized to this brain region (Fig. 1 b). The lack of mature neuritic plaques in this area suggests that this brain region may have only recently been affected pathologically. In contrast, the superior temporal gyrus from AD patient 1173 showed subregional specificity, where an area with most cells being intensely TUNEL-positive was found abutting an area with much fewer and less intense TUNEL-positive nuclei (Fig. 1 c). Analysis of a serial section with anti-Aβ antibody indicated that the area with the large number of TUNEL-positive cells also contained numerous diffuse plaques, whereas the adjoining area with fewer TUNEL-positive cells had barely any detectable extracellular Aβ immunoreactivity (Fig. 1 d).

Not surprisingly, the region of the brain with the most extensive TUNEL-positive cells varied from individual to individual. For instance, the inferior temporal cortex of AD patient 1112 showed a uniformity of intense TUNEL-positive cells (Fig. 1 e) and a corresponding increase in Aβ immunoreactivity (Fig. 1 f). A serial section stained by H&E was unremarkable except for a detectable change in the nuclear morphology of most cells (Fig. 1 g). Bielschowsky silver stain revealed early indications of dystrophic neurites in association with the diffuse Aβ deposits (Fig. 1 h). In addition, we also observed occa-

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**Table I. Clinical and Pathological Characteristics of Case Material**

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IF, inferior frontal gyrus; HC, hippocampus; MFG, middle frontal gyrus; MTG, middle temporal gyrus; SP, superior parietal; STG, superior temporal gyrus; TP, temporal pole; C, control.
sional cells with intracellular neurofibrillary tangles that were also TUNEL-positive. Notably, since most of the AD patients exhibited regional cellular DNA damage, the lack of TUNEL-positive cells in regions where Aβ immunoreactivity was not detected provided an internal control for the absence of cell death resulting from postmortem changes.

AD patient 1002 was exceptional among the cases studied, in that intense Aβ immunoreactivity (Fig. 1 j) in association with dystrophic neurites (Fig. 1 l), was found throughout the gray matter, regardless of the brain region analyzed. A representative histological section from this patient is shown. Interestingly, although the majority of cells in the superficial layers of the superior temporal gyrus showed evidence of DNA fragmentation, they were weakly detectable by TUNEL. Since H&E staining revealed that the brain parenchyma was spongy, possibly reflecting the dropout of cells, and cell nuclei were all highly condensed and pyknotic (34), this suggests that these cells have progressed to a terminal stage of the cell death pathway (Fig. 1 k). Furthermore, this finding indicates that the TUNEL method cannot be used to detect effectively dead cells in which the cellular DNA has been totally degraded.

Representing brain samples from five of the six controls, the temporal pole of sample 532 showed not even a single TUNEL-positive cell (Fig. 1 m), had no extracellular Aβ deposition (Fig. 1 n), revealed no changes in cell morphology (Fig. 1 o), and had no detectable neuritic abnormalities (Fig. 1 p). Significantly, postmortem changes that might have resulted from the delay in processing of the brain samples did not yield any background staining after the TUNEL method. Occasional TUNEL-positive cells were found scattered throughout sample 960 (from a 101-yr-old man suffering from clinical depression but with no known history of AD), but extracellular Aβ deposition was not detected anywhere (data not shown). This finding may be representative of normal aging rather than AD.

Intracellular Aβ correlates with cell death. To investigate the relationship between extracellular Aβ deposition and cell death, tissue sections from the AD brains were analyzed for small foci of TUNEL-positive cells, which we posited represent the earliest evidence of cell death, to determine whether extracellular Aβ immunostaining invariably was associated with them. Since many of the AD patients had extensive plaques confined to focal subregions of the brain, it was relatively easy to locate small clusters of TUNEL-positive cells in areas that otherwise appeared histologically normal. In each of those areas containing small foci of cells with DNA damage, not only did we not detect any extracellular Aβ deposition, but rather we observed intracellular Aβ-like immunoreactivity. A representative sample (Fig. 2 a) shows the subiculum of AD patient 1,112, where small focal areas of perhaps 50–100 cells are observed, some of which were intensely TUNEL-positive, and others that were only weakly labeled. Aβ immunostaining revealed no evidence of extracellular Aβ plaques in the immediate vicinity, but showed intracellular Aβ-like immunoreactivity of varying extent in most of the cells within the cluster (Fig. 2 b). The 4G8 monoclonal antibody used for this analysis is directed against an epitope consisting of amino acids 17–24 of Aβ, and recognizes diffuse and senile plaques in AD brains, although it cannot be ruled out that it may recognize APP as well (27). Detection of thioflavin S-reactive deposits (Fig. 2 h) within cells with intracellular Aβ immunoreactivity (Fig. 2 f), however, suggests accumulation of the Aβ protein in these neurons. While we cannot exclude the possibility that the intracellular Aβ reactivity might result from uptake from the extracellular compartment, our failure to detect Aβ plaques in the vicinity may argue against this suggestion. It has to be pointed out that the various methods used, such as TUNEL, immunohistochemistry, and thioflavin staining, have different intrinsic sensitivities, and attempts to draw conclusions from such a study are dependent upon the assumption that changes detected by each method can be compared directly.

Although the TUNEL staining was predominantly intranuclear, the Aβ immunostaining was exclusively cytoplasmic. The gradation of TUNEL and Aβ reactivities may suggest that cell death in these areas was restricted to cells with intracellular Aβ-like immunoreactivity, and that cells with a higher level of Aβ accumulation were more prone to undergo cell death. Such findings were not restricted to a specific subtype of neurons, based on their morphological appearance and geographical location. In the CA3 region of the hippocampus of AD patient 1240, focal groups of neurons had acquired a spindleshaped morphology, showed intense TUNEL reactivity (Fig. 2 c), and demonstrated cytoplasmic accumulation of Aβ immunoreactivity (Fig. 2 d). Again, no extracellular Aβ deposition could be detected in the vicinity, although we cannot exclude the possibility that the TUNEL staining observed in the cell body stems from long-distance neuritic damage incurred in an area enriched with extracellular Aβ.

The presence of intracellular Aβ-like immunostaining appeared to occur without detectable extracellular Aβ plaques, although the presence of extracellular Aβ deposits predicted the coexistence of intracellular Aβ. Every brain region from every AD patient with either diffuse or neuritic plaques contained intermingling cells with intracellular Aβ-like immunoreactivity, albeit at different cellular densities and staining intensities (Fig. 1, b, d, f, and j). Intracellular Aβ-like immunoreactivity was detected not just with the mouse monoclonal antibody (4G8), but also with a rabbit polyclonal antibody against Aβ1-28 (15), and the immunostaining could be competed out using the immunizing Aβ peptide (data not shown). Interestingly, somewhat similar but much less intense intracellular immunostaining was also observed in some of the control brains (Fig. 1 n). Unlike that observed in AD patients, however, only few scattered cells that were not organized into defined foci were stained. More importantly, they were never correlated with TUNEL staining, possibly because the level of Aβ was below threshold for inducing cellular damage. Control brain sample 960 was exceptional in that an increase in the

Figure 1. Detection of cell death in the brain of AD patients. Analysis of paraffin-embedded brain sections (6 μm) from different AD patients (a–l) and from a representative control individual (m–p) revealed a good correlation between the presence of TUNEL-positive cells and extracellular deposition of the Aβ protein. Samples a and b are serial sections from the superior temporal gyrus (STG) of patient 1250; c and d, STG of patient 1173; e–h, the inferior temporal cortex of patient 1112; i–l, STG of patient 1002; and m–p, the temporal pole of a non-AD control. Sections are stained by the TUNEL method (a, c, e, i, m), for Aβ with antibody 4G8 (b, d, f, j, n), by H&E (g, k, o), or by Bielschowsky’s silver stain (h, l, p). ×50.

gp330-mediated apoE Uptake and Intracellular Aβ Stabilization
Figure 2. Evidence for extracellular deposition of A\(\beta\) upon cell death. Brain sections from AD patients were stained by the TUNEL method (a, c, e), for A\(\beta\) with antibody 4G8 (b, d, i–k), by H&E (g), by thioflavin S (h), or by Bielschowsky’s silver stain (l). Samples a and b are serial sections from the subiculum of patient 1112; c and d, the CA3 region of the hippocampus of patient 1240; e–h, the lateral geniculate body of patient 1240; i–j, the STG of patient 1173; k and l the STG of patient 1002. a–d, ×100; e–h, ×200; i–l, ×100.

Figure 3. Evidence for apoE uptake by dying neurons. Serial brain sections from an AD patient were stained by PAS (a), for apoE with antibody 3H1 (b), for A\(\beta\) with antibody 4G8 (c), for LRP with antibody 8G1 (d), for gp330 with antibody 1H2 (e and g), by the TUNEL method using True Blue for visualization (f), successively for gp330 with antibody 1H2 using 3,3′-diaminobenzidine, and by the TUNEL method using True Blue (h), or by Bielschowsky’s silver stain (i). ×100.
gp330-mediated apoE Uptake and Intracellular Aβ Stabilization
number of Aβ-reactive cells was correlated with TUNEL-positive cells.

Accumulating evidence indicates that apoptosis may play an important role in cell loss in AD (14, 15). Since the TUNEL assay can label cells that are dying by either apoptosis or necrosis (18), other criteria are required to define better the mechanism of cell death. Apart from TUNEL reactivity, we also observed a common series of morphological changes in many, but not all, of the TUNEL-positive cells in the AD brains that is more suggestive of cell death by apoptosis. Specifically, a high magnification view of the lateral geniculate body of patient 1240 showed cells at different stages of the apoptotic pathway (Fig. 2, e–h). Stage 1 cells were completely TUNEL-negative with no detectable DNA fragmentation within their nuclei, and no detectable intracellular Aβ reactivity. These cells were considered normal neurons. Stage 2 cells had somewhat enlarged nuclei with variable amounts of TUNEL staining due to the different extent of DNA fragmentation and marked expansion of the cytoplasmic compartment that had abundant Aβ-like immunoreactivity and thioflavin S-reactive deposits. We classified these as early-stage apoptotic cells that have not previously been identified morphologically. Stage 3 cells had TUNEL-positive nuclei that were somewhat condensed, and had lost their open chromatin morphology, as well as a cytoplasmic region that showed compact Aβ immunostaining with increased basophilia and TUNEL-staining, suggesting nuclear leakage. These were intermediate-stage apoptotic cells. Stage 4 cells frequently had a crescent to panhandle shape where the collapsed TUNEL-positive nuclei appeared in the process of being extruded from one end of the cell. These were considered late-stage apoptotic cells.

**apoE uptake correlates with intracellular Aβ accumulation.**

In describing these stages, it appears that DNA fragmentation was preceded by a marked expansion of the cytoplasmic compartment of the cell that accommodated the accumulation of not just thioflavin S-reactive, Aβ-related material, but also coarse lipid granules, as defined by staining with either H&E or PAS (Fig. 3 a). The PAS staining may be consistent with the accumulation of lipofuscin, a neuronal pigment for which biochemistry remains undefined (35). These granules were found invariably in the cytoplasm of neurons in each of the AD patients during every step of the apoptotic process (stages 2–4).

Since apoE isoforms are a major risk determinant for AD (18–21), and recognizing that apoE is the predominant lipid carrier in the central nervous system (36), we hypothesized that the PAS-positive, lipofuscin-like material in the neurons might contain apoE. To investigate this possibility, we performed immunohistochemical staining with both a monoclonal (3H1) and a polyclonal antibody (28). We found that the PAS-positive cells contain varying levels of apoE-immunoreactivity, regardless of the antibody used (Fig. 3 b). Interestingly, apoE has also been found to colocalize with Aβ within the same cytoplasmic granules (Fig. 3 b and c). Since lipid-containing apoE has been shown to bind Aβ (37–40), and the two have been detected as a complex in AD brains (41), it is possible that the intracellular accumulation of lipids serves to stabilize the hydrophobic Aβ protein (or the proteolytic precursors from which it is derived).

As apoE is believed to be produced by glial cells in the CNS (42, 43), its accumulation in neurons is likely the result of receptor-mediated uptake (22, 23). Accordingly, we analyzed AD brains to determine if we could identify the receptor. Four members of the LDLR gene family have been shown to bind and internalize apoE in cultured cells; they are LDLR (44, 45), VLDLR (46), LRP (47, 48), and gp330 (49, 50). We speculated that the receptor that is not normally activated but is inducible, is a likely candidate that is responsible for initiating a cascade of events leading to AD.

Using antibodies specific to each of the receptors, we investigated their expression profile in neurons from different brain regions. Both LDLR and VLDLR were detected indiscriminately at a very low level in both control and AD brains (data not shown). Similarly, LRP, while more easily detected, also did not discriminate between regions of the brain that did or did not contain AD lesions; in a serial section containing cells with varying levels of apoE and Aβ, LRP was found uniformly on the cell membrane and processes of every neuron (Fig. 3 d), with either polyclonal antibody R777 (31) or monoclonal antibody 8G1 (30). On the other hand, gp330 was found in abundance almost exclusively on cells in the pathologically affected areas (Fig. 3 e), with virtual sparing in normal regions except for an occasional cell with weak immunoreactivity. The immunostaining could be detected with either polyclonal antibody R239 (29) or monoclonal antibody 1H2 (51). The immunostaining within the involved areas highlighted the cell membrane and its extensions, consistent with the location of a cell surface receptor (Fig. 3 g). Remarkably, the level of gp330 varied markedly between cells (Fig. 3 e), reflecting perhaps both the differing levels of intracellular apoE (Fig. 3 h) and the varying extent of DNA fragmentation (Fig. 3 f). Indeed, in a double-labeling experiment, we have been able to demonstrate that cells with the highest gp330 expression have the highest intranuclear TUNEL reactivity (Fig. 3 h). Of the four known receptors, only activation of gp330 correlated with DNA fragmentation and, possibly, the induction of apoptosis.

Interestingly, only an occasional cell with fragmented DNA was observed to show evidence of neurofibrillary tangles (Fig. 3 i). Since the tangles were evaluated by Bielschowsky’s silver stain, and not by immunohistochemical staining, it is possible that their numbers are slightly underrepresented. It did, however, appear that when present, these structures were found invariably around the periphery of the cell surrounding the collection of apoE- and Aβ-containing granules.

If the dying neurons contain Aβ-related material, it is possible that these molecules may be released to the surrounding parenchyma sometime during the apoptotic process, thereby giving rise to the extracellular Aβ deposits. This is suggested by our findings with a transgenic mouse model, in which intracellular expression of Aβ leads to neuronal cell death and formation of diffuse extracellular Aβ deposits (16). To explore this possibility, we analyzed regions of the brain from AD patients with marginal extracellular Aβ immunostaining. In the least involved regions, we observed diffuse plaques that were not much larger than the size of a neuron. For example, in the superior temporal gyrus of AD patient 1173, anti-Aβ immunostaining revealed numerous small halolike structures intermingled among neurons with intense intracellular Aβ reactivity (Fig. 2 i). Within the halos were frequently either an unstained cell ghost, or a darkly stained structure that resembled a collapsed cell. This finding suggested the diffusion of Aβ away from dying neurons. In an adjoining region where extracellular Aβ immunostaining was more extensive, even larger but still well-defined halos were observed (Fig. 2 j). This latter observation suggested that the Aβ protein became insoluble in the

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extracellular environment shortly after its release from the dying cell. In fact, these early structures were observed in every AD patient analyzed, and were found interspersed among advanced plaques.

We observed that many of the larger anti-Aβ immunostained structures from the superior temporal gyrus of AD patient 1002 had variable numbers of cell ghosts within them (Fig. 2 k). Surrounding these large Aβ deposits were invariably smaller Aβ structures of different sizes. This observation is consistent with large neuritic plaques being derived from multiple neurons either dying simultaneously or in succession. The extent of coalescence of individual structures determines the ultimate size of the neuritic plaque. It is anticipated that such plaques will continue to enlarge with increased severity of cell death. Bielschowsky silver stain confirmed the frequent presence of multiple darkly stained cores within the same large neuritic plaque (Fig. 2 l). The detection of dystrophic neurites within these plaques may suggest secondary involvement of neighboring neuritic and synaptic structures as well.

Discussion

Our findings indicate that neurons in the AD brain are particularly susceptible to accumulating DNA damage, and consequently to degenerate and die. Although TUNEL is not a definitive marker of apoptosis, it still allows identification of dying cells in situ. The number of damaged cells identified by TUNEL in the AD brain may appear unexpectedly high, but our results are comparable to those reported by others (19–21). It must be emphasized that the time scale for the demise of apoptotic cells has not been characterized in the adult brain, especially in response to chronic pathological stimuli. Potentially, these cells may acquire DNA damage, yet persist for some time before their eventual removal; this may, in fact, be reflected by the varying intensities of TUNEL staining that have been observed between individual cells within a region. Alternatively, it is possible that the extensive number of TUNEL-positive cells is part of a coordinated response that may have contributed to the death of the patient. In addition, since these damaged cells occur in focal areas of the AD brain and are not widely distributed, it indicates that cellular DNA fragmentation is a consequence of the disease process, and is not likely due to postmortem effects.

In this present study, morphological and biochemical endpoints were examined in defining some of the molecular events associated with cellular DNA fragmentation in the AD brain. Not only did we find that damaged cells detected by the TUNEL assay were generally localized in regions that contained extracellular Aβ deposition, but we also found that the extent of TUNEL staining could be correlated with the detection of intracellular Aβ-like immunoreactivity. Lassmann et al. (20) also reported that TUNEL-positive cells were positioned in brain regions containing extracellular Aβ deposition. In this regard, although there has been some debate as to whether amyloid deposition correlates with the degree of dementia (10, 52), recent evidence indicates that the severity of dementia can be correlated with the Aβ load (12). This does not, however, imply that other factors such as cytoskeletal abnormalities are not also involved in the disease process (53).

Although the site of Aβ generation in vivo is unresolved, there is increasing evidence for its derivation or accumulation occurring intracellularly. The possibility that Aβ accumulation occurs within cells, however, is controversial and difficult to verify in histological sections since Aβ is derived from APP, and any Aβ-specific antibodies may concomitantly recognize various proteolytic fragments of APP as well. Nevertheless, analysis of various cell lines and animal models suggest that Aβ may occur within neurons: (a) the Swedish mutation in the APP gene results in increased Aβ accumulation in cultured cells (54); (b) Aβ can be directly immunoprecipitated from cell extracts of a neuronal-like cell line, NT2N (55); (c) the form of Aβ deposited in plaques (Aβ1–42) is resistant to intracellular degradation, but not to Aβ1–40 (56); (d) using transient transfection assays, intracellular expression of amyloidogenic fragments results in formation of amyloid-like fibrils that suffice to kill cells (57); (e) nonfibrillar Aβ peptide exists intracellularly within neurons before the appearance of extracellular deposits of Aβ in aged monkey brains (58); (f) intracellular Aβ immunoreactivity has been documented within dog brains (59); (g) rabbits fed a high cholesterol diet accumulate Aβ immunoreactivity within neurons (60); (h) several reports indicate that Aβ is associated with intracellular neurofibrillary tangles (43, 61, 62).

Based on our findings, the following sequence of events represents a plausible mechanism to account for extracellular Aβ deposition and cell death. The ability of neurons to synthesize Aβ, which is a normal processing event, appears insufficient to assure intracellular Aβ accumulation; other cofactors are required. The accumulation of Aβ in neurons is facilitated by the presence of apoE, which may provide a suitable lipidic environment to stabilize the hydrophobic Aβ. The uptake of apoE from the surrounding environment appears to be dependent upon the neuronal expression of gp330, a member of the LDL receptor family, and on the apoE variant secreted by neighboring astrocytes as to whether it can efficiently bind gp330 (apoE3 and apoE4 can bind, but apoE2 cannot). Stabilization of Aβ inside the neurons above a threshold level may trigger cell death. The death of neurons would result in the release of Aβ, possibly in a complex with apoE, which over time would evolve into neuritic plaques. While our findings suggest that intracellular accumulation of Aβ may suffice to induce neuronal cell death which then leads to the extracellular deposition of Aβ, it is also likely that the neuritic injury and glial activation resulting from the formation of senile plaques induce a second level of cell killing (13, 14, 63); in that case, the extent of neuronal cell death can be amplified greatly.

The precise mechanism by which neurons die in the AD brain still remains to be determined. Defining the molecular mechanism may eventually contribute to the design of therapeutic interventions. It is likely, however, that both apoptotic and necrotic mechanisms will be involved in the neuronal cell death that occurs in AD brains. Accumulating evidence suggests that apoptosis is likely to be an important factor in AD pathology. In this regard, a role for Aβ in inducing apoptosis in cell culture and in transgenic mice has been documented (14–16); however, whether the concentration of Aβ in the AD brain is adequate to induce directly apoptotic cell death is unclear. Furthermore, other gene products linked to familial early-onset AD, such as presenilin-2, have also been implicated as playing a role in apoptosis (64, 65).

The product of another AD-linked gene may also play a role in apoptosis through its interaction with Aβ. The APOE genotype exerts a profound influence on the susceptibility of the individual to AD (37, 66–68). Strittmatter et al. (37) have proposed that apoE functions as an isofrom-specific stabilizer

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of Aβ. The three major isoforms of the apoE protein, designated E2, E3, E4, are products of three alleles at a single gene locus (36). The predominant E3 molecule is considered the parental form, while E2 (Arg158→Cys) and E4 (Cys112→Arg) are considered variants. Specifically, the e4 allele increases the risk and lowers the age of onset of AD, while the e2 allele appears to decrease the risk and increase the age of onset. Indeed, Ohm et al. (69) have suggested that intraneuronal apoE appears a decade earlier as a function of the e4 allele. Our suggestion that neuronal uptake of apoE is involved in Aβ-induced cell death in the AD brain may explain why E4 exacerbates and E2 protects against the development of AD, since one known biochemical difference of the AD apoE variants is that E4 can efficiently bind the LDL receptor while E2 cannot (70). It is relevant to note that the ligand-binding domain of LDLR and gp330 are highly conserved (50). That activation of gp330 expression may be involved in the neurodegenerative process is intriguing. There have already been reports of apoE uptake into endosomes (71, 72) and studies on intracellular trafficking of apoE complexes (73). Additionally, there has been suggestion of Aβ enhancing the uptake of apoE (74).

Pathological chaperones have been hypothesized to mediate amyloid formation (75, 76) and apoE has been shown to induce Aβ aggregation and filament formation in cell-free systems (39, 40). Furthermore, Aβ–apoE complexes have been isolated from AD brains (41). These observations are consistent with our findings that Aβ and apoE colocalize in neurons, and our suggestion that association between the two within the cell may suffice to cause injury and trigger the apoptotic pathway. After cell death, Aβ and apoE may be released as a complex (37, 41, 77). We have observed that a key difference between a normal individual and an AD patient is the number of apoE-containing cells within a specific region of the brain. In control individuals, these cells are scattered, and the apparent level of apoE does not appear sufficient to cause cell death. In AD patients, these cells are found in clusters, and show both varying levels of intracellular apoE and different extent of unclear DNA fragmentation. A highly related receptor, LRP, has been shown recently to be able to internalize APP, and to mediate its degradation (78). LRP and gp330 bind many ligands in common, and it is possible that gp330 may internalize APP as well. If so, the increased expression of gp330 would lead to not only increased apoE uptake, but also increased degradation of APP, thereby generating the pathological situation requisite for AD.

Acknowledgments

We thank Dr. H.M. Wisniewski and Dr. K.S. Kim for the 4G8 antibody, R. Milne for the 3H1 antibody, and Ms. Luisa Ruiz for assistance in the preparation of the manuscript.

We acknowledge the support of U.S. Public Health Service grant AG 05146, the Alzheimer’s Association, and the Charles A. Dana Foundation.

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