Reduction of O-Linked N-Acetylglucosamine-Modified Assembly Protein-3 in Alzheimer’s Disease

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Abnormal protein processing and modification is associated with Alzheimer’s disease (AD) pathology. The role of phosphorylation in AD has been studied extensively because the presumed abnormal phosphorylation of tau protein is believed to play a role in the formation of paired helical filaments. Glycosylation with O-linked N-acetylglucosamine (O-GlcNAc) to serine and threonine residues is a dynamic protein modification of intracellular proteins, and it shares similar features with protein phosphorylation. In this study, O-GlcNAc glycosylation of proteins from autopsied human brains with confirmed AD and non-AD age-matched controls was examined. O-GlcNAcylation was demonstrated by labeling protein extracts with [3H]galactose in the presence of galactosyltransferase and subsequent analyses of saccharide–protein linkage products in AD brains (Ledesma et al., 1994; Griffith and Coleman, 1995). NFTs are composed largely of hyperphosphorylated tau protein (Goedert 1993 and references within). Abnormal tau phosphorylation may cause NFT formation and cytoskeletal disruption and may ultimately lead to neuronal malfunction and death. Recent observations of glycation and glycosylation products in AD brains (Ledesma et al., 1994; Griffith and Schmitz, 1995; Smith et al., 1995; Wang et al., 1996) suggest that protein modifications other than phosphorylation may also contribute to AD pathology.

Glycosylation by N-acetylglucosamine monosaccharides, O-linked to the hydroxyls of serine or threonine residues (O-GlcNAc), is a unique type of protein modification (Torres and Hart, 1984; Hart et al., 1995; Haltiwanger et al., 1997). Unlike most other types of glycosylation, the majority of O-GlcNAc-modified proteins are found within the nucleoplasm and cytoplasm of cells (Hart et al., 1995) O-GlcNAcylating enzymes. Analogous to phosphorylation reactions catalyzed by kinases and phosphatases, O-GlcNAc transferase and O-GlcNAcase are responsible for the addition and removal of O-GlcNAc, respectively (Haltiwanger et al., 1992; Dong and Hart, 1994). Virtually all O-GlcNAc proteins are also phosphoproteins (Hart et al., 1995; Haltiwanger et al., 1997), and at least for some of them, the degree of O-GlcNAcylation appears to be reciprocal to phosphorylation (Kelly et al., 1993). Thus, O-GlcNAc may play a role in regulating protein phosphorylation, or it may regulate proteins in a manner analogous to phosphorylation.

Changes in O-GlcNAc level have been observed in cells under various stimuli. Activation of T-lymphocytes by mitogens or by phorbol esters resulted in rapid changes in O-GlcNAc levels in many nuclear and cytosolic proteins (Kearse and Hart, 1991). When HT29 cells, a human colon cancer cell line, were treated with okadaic acid or microtubule-destabilizing agents, O-GlcNAc glycosylation of many proteins in crude cell extracts was altered (Chou and Omary, 1994). Furthermore, after colcemid-induced mitotic arrest, keratin proteins from HT29 cells showed a significant change in both O-GlcNAc glycosylation and phosphorylation (Chou and Omary, 1993). These results suggest that the level of O-GlcNAc, like phosphorylation, could be a widespread event in cells that respond to stimuli.

As a beginning to elucidating the possible involvement of O-GlcNAc in AD, in which abnormal phosphorylation occurs, we have been examining the O-GlcNAc level in brain tissues affected by AD. In this study, using galactosyltransferase as a probe, we compared the levels of O-GlcNAc in crude extracts of AD brains with neurologically normal age-matched controls. The overall GlcNAc level was not much different in AD brains when compared with controls except for a substantial decrease in the level of a 160 kDa O-GlcNAcylated protein in AD. We have identified this 160 kDa O-GlcNAcylated protein as clathrin assembly enzyme.
protein-3 (AP-3) and obtained quantitative information on its reduction in several brain regions of AD.

**MATERIALS AND METHODS**

Reagents. Human milk galactosyltransferase was obtained from Boehringer Mannheim (Indianapolis, IN). The enzyme was autogalactosylated by the manufacturer and shown to contain negligible amounts of GlcNAc sites (see Fig. 1). Peptide N-glycosidase F (PNGase F) was also from Boehringer Mannheim. Uridine diphosphate [3H]Galactose (8.1 Ci/mmol) and EN' HANCE were obtained from DuPont NEN (Boston MA). Ovalbumin, galactose, and disaccharide standards were from Sigma (St. Louis, MO). F1–20 antibody and CLAP3 antibodies were generously provided by Dr. Eileen M. Lafer (Center for Molecular Medicine, University of Texas Health Science Center) (Sousa et al., 1990) and Dr. James H. Keen (Department of Pharmacology, Thomas Jefferson University) (Murphy et al., 1991), respectively. Anti-AP-3 antibody (clone AP180-I) (Ahle and Ungewickell, 1986) and anti-AP-2 antibody (clone 100/2) (Ahle et al., 1988) were purchased from Sigma. Anti-clathrin heavy chain antibody (clone CHC 5.9) was purchased from ICN Biomedicals (Aurora, OH). All chemicals and reagents used were of the highest quality available.

Human brain tissues. Postmortem human brain tissues were provided by the Alzheimer's Disease Center, University of Rochester. All cases were well characterized on the basis of clinical and neuropathological criteria (Kachaturian, 1985). All cases were judged in a consensus conference of neurologists, psychiatrists, neuropsychologists, and neuropathologists on the basis of clinical history and neuropathological survey of the brain with adherence to the National Institutes of Health consensus age-adjusted criteria for definition of AD (Kachaturian, 1985). Information regarding clinical history, mental status, age, gender, the interval between death and autopsy, and pertinent pathology have all been documented in detail.

Postmortem tissues from three brain regions of both AD and age-matched controls were used for this study: (1) *frontal cortex*, including gray and white matter of neocortex at the level of the olfactory bulb; all deep structures were removed; 1-cm-thick slabs of frontal cortex from five controls and five AD cases were used for one part of the study (Table 1 provides information about each case); (2) *middle frontal gyrus*; and (3) *anterior cerebellum*. Middle frontal gyrus and anterior cerebellum from 11 individuals with AD and 10 age-matched controls were used for another part of the study. Table 2 provides information related to each case.

NFT density was determined for each case by manual counting and averaging of the five most severely affected (for AD) or random (for non-AD controls) microscopic fields at 200× total magnification (1.29 mm² viewing area).

Preparation of brain extracts. All tissues were collected directly at autopsy with postmortem delays indicated in Tables 1 and 2. The samples were immediately frozen on dry ice and then transferred to liquid nitrogen for storage. The tissues were maintained in their frozen state until processing for protein extraction.
with galactosyltransferase for 1 hr at 37°C as described above. The labeled immunoprecipitates were washed again, boiled in sample buffer containing 2% SDS and 10% glycerol, and then analyzed by one- or two-dimensional SDS-PAGE and fluorography.

**SDS-PAGE, fluorography, and immunoblot analysis.** SDS-PAGE was performed according to the method of Laemmli (1970). For fluorography, gels were stained with Coomassie blue, treated with EN3HANCE, dried, and exposed to Eastman Kodak (Rochester, NY) x-ray film at -80°C for an appropriate length of time. Some gels were silver stained by the method of Blum et al. (1987).

For immunoblot analysis, 3H-labeled samples were separated on SDS-PAGE and transferred to polyvinylidene difluoride (PVDF) membranes. The membranes were blocked with 5% nonfat dry milk in PBS containing 1% Tween 20 at 4°C for 6 hr. The membranes were then incubated with AP180-I monoclonal antibody (mAb) (1:5000), F1–20 mAb (1:1000), CLAP3 mAb (1:1000), or anti-clathrin antibody (1:500) at 4°C overnight, followed with an appropriate secondary antibody conjugated with horseradish peroxidase. The antibody was then detected with an enhanced chemiluminescence (ECL) system using luminal and H2O2. Negative controls consisted of membranes incubated in the absence of primary antibodies.

**Two-dimensional electrophoresis.** Two-dimensional electrophoresis was performed according to the method of O’Farrell (1975) by Kendrick Labs, Idaho. Isoelectric focusing was performed using 2% ampholines pH 4–8 and lyophilized IEF strips. The PVDF membrane was immobilized on a ProBlott (Molecular Dynamics, Sunnyvale, CA). The PVDF membrane was immersed in the sample for 1 hr at room temperature and then transferred to a PVDF membrane. The membranes were blocked with 5% nonfat dry milk in PBS containing 1% Tween 20 at 4°C for 6 hr. The membranes were then incubated with AP180-I monoclonal antibody (mAb) (1:5000), F1–20 mAb (1:1000), CLAP3 mAb (1:1000), or anti-clathrin antibody (1:500) at 4°C overnight, followed with an appropriate secondary antibody conjugated with horseradish peroxidase. The antibody was then detected with an enhanced chemiluminescence (ECL) system using luminal and H2O2. Negative controls consisted of membranes incubated in the absence of primary antibodies.

**Protein–saccharide linkage and saccharide structure analysis.** For PNGase F digestion, 3H-galactosylated proteins were denatured in solution containing SDS and DTT. After adding a solution containing 75 mM Tris-HCl, pH 8.6, 10 mM EDTA, 0.7% (w/v) NP-40, and protease inhibitor mixtures (in mg/ml: leupeptin, 1; antipain, 2; benzamidine, 10; aprotinin, 1; and pepstatin, 1), PNGase F (5 μl, 200 μU/μl determined by the manufacturer) was added and incubated at 37°C for 16–24 hr. An additional 5 μl of PNGase F was added and incubated for 16–24 hr further. The same digestion condition was also used for 3H-galactose-labeled ovalbumin to serve as a control. The reaction was stopped by heating for 5 min. After fractionation over a Sephadex G-50 column (1 × 60 cm), products eluted at v0 were pooled, lyophilized, and analyzed by SDS-PAGE followed by fluorography. In some cases, the bands of interest were cut from gels (using the corresponding fluorographs as guides), and the amount of radioactivity on the gel slices was determined by scintillation counting.

For β-elimination experiments, the 3H-galactosylated-labeled protein samples were first resolved in SDS-PAGE and transferred to a PVDF membrane. The membrane was sprayed with EN3HANCE and exposed to a Kodak x-ray film. Using the film as a guide, the membrane areas corresponding to the bands of interest were cut out and subjected to β-elimination by incubating the membrane strips in 0.5 ml of β-elimination buffer (1 M NaOH, 0.1 M Na2CO3) at 37°C for 24 hr. For the β-elimination reaction, membrane strips were counted. β-Elimination products in the reaction solution were neutralized with acetic acid and passed through a 3 ml Dowex 50 × 8 (hydrogen form) column. Lyophilized samples were then washed three times with methanol with 0.1% acetic acid, dried, suspended in H2O, and analyzed by the method of Laemmli (1970).

**Analysis of galactosyltransferase-labeled proteins from selected brain regions of AD.**

**Frontal cortex**

To evaluate O-GlcNAc-containing proteins in AD brains, extracts from frontal cortices of end-stage, confirmed AD, and age-matched controls were labeled with galactosyltransferase under the conditions in which all accessible terminal GlcNAc was labeled with 3H-galactose (see above). The labeled proteins were equally divided and resolved separately on the same SDS-PAGE gel to determine which proteins contain terminal GlcNAc residues for galactosylation. Half of the gel was silver-stained (Fig. 2A), and the other half was subjected to fluorography (Fig. 2B). It is apparent that many major silver-stained protein bands were

**RESULTS**

**Optimization of galactosyltransferase labeling conditions for proteins from crude lysates of human brains.**

GlcNAc β-(1–4) galactosyltransferase is a specific enzyme that adds galactose covalently to terminal N-acetylgalcosamine residues of both N- and O-linked carbohydrate structures (Wallenfels, 1979; Beyer et al., 1981; Torres and Hart, 1984). By using UDP-[3H]galactose as a sugar donor, terminal GlcNAc residues of glycoproteins accessible to the enzyme can be labeled. The [3H]galactosylated products are stable and allow for further protein–saccharide linkage analysis.

In this study, we used galactosyltransferase as a probe to examine O-GlcNAc-modified proteins in the crude lysates from autopsied human brains of confirmed AD and neurologically normal age-matched controls. The optimal amount of galactosyltransferase required was determined by incubating proteins (100 μg of total proteins) with UDP-[3H]galactose and increasing concentrations of galactosyltransferase as described (Roquemore et al., 1994; Haltiwanger and Philipsberg, 1997). The [3H]galactosylated products were then fractionated over a Sephadex G-50 column. The quantity of protein-bound [3H]galactose, eluting in the v0 of the column, was determined at each transferase concentration (Fig. 1A). The amount of label incorporated was undetectable when no enzyme was added, indicating a negligible amount of endogenous galactosyltransferase activity present in the sample. The human milk galactosyltransferase (Boehringer Mannheim) used in the study contained an insignificant amount of terminal GlcNAc sites. Because these brain proteins demonstrated saturated [3H]label incorporation at a galactosyltransferase concentration of ≤20 μU (defined by the manufacturer), all the subsequent analyses were conducted at a galactosyltransferase concentration of 20 μU.

To compare the galactosyltransferase-accessible sites on proteins in the samples quantitatively, it was necessary that the protein concentration be within the linear range of the galactosyltransferase labeling. Various amounts of proteins (25–400 μg) were labeled with UDP-[3H]galactose and 20 μU of galactosyltransferase (Fig. 1B). Between 25 and 200 μg of protein, the amount of radiolabeling was proportional to the amount of protein in the samples. The labeling continued to rise when 400 μg of protein was used but at a slower rate. Because 100 μg of protein was within the linear range of the assay, the remaining studies were conducted using 100 μg of total proteins. Figure 1C shows that maximal [3H]galactosylation occurred at 1 hr of incubation. Results shown in Figure 1 are obtained from control brain proteins. Similar profiles were seen in proteins from AD brains (data not shown).

**Analysis of galactosyltransferase-labeled proteins from selected brain regions of AD.**

To evaluate O-GlcNAc-containing proteins in AD brains, extracts from frontal cortices of end-stage, confirmed AD, and age-matched controls were labeled with galactosyltransferase under the conditions in which all accessible terminal GlcNAc was labeled with [3H]galactose (see above). The labeled proteins were equally divided and resolved separately on the same SDS-PAGE gel to determine which proteins contain terminal GlcNAc residues for galactosylation. Half of the gel was silver-stained (Fig. 2A), and the other half was subjected to fluorography (Fig. 2B). It is apparent that many major silver-stained protein bands were
Figure 1. Optimal conditions for the galactosyltransferase-mediated galactosylation of brain proteins. Labeling as a function of enzyme concentration (A), substrate concentration (B), and time (C). For each parameter, certain variables were held constant as needed: 100 μg of total proteins, 20 mU of enzyme, and 60 min. A and B were repeated three times, and C was repeated twice, and similar profiles were obtained.

Figure 2. Comparison of galactosyltransferase labeling of AD and control brain proteins. Proteins from frontal cortex extract of AD and age-matched controls were labeled with galactosyltransferase. Radiolabeled samples were then equally divided and resolved separately on the same 10% SDS-PAGE. The half of the gel visualized by silver staining (A) shows both labeled and unlabeled proteins. The other half of the gel that was subject to fluorography (B) reveals labeled proteins. Positions of molecular weight standards (kilodaltons) are indicated in the middle. Densitometric scanning of lanes 1 and 3 of B (the fluorograph) is shown in C. Peak 3, 160 kDa; peak 7, 50 kDa, as mentioned in the text.
not radiolabeled by galactosyltransferase, indicating the specificity of the enzyme. The silver-stained gel also shows that total amount of protein and overall protein pattern were similar in control and AD extracts. Figure 2 represents the results from two of the five AD and two of the five control brains examined. All five yielded similar results.

In the control tissues, a protein with a molecular size of \( \sim 160 \) kDa was labeled more intensely with \([3H]\)galactose than any other labeled protein band (Fig. 2B, lanes 1, 2, arrowhead). A striking difference between control and AD brain tissues was that the intensity of the radiolabeling for the same 160 kDa protein was decreased in AD tissues (Fig. 2B, lanes 3, 4). This considerable reduction is more evident from densitometric scanning of the fluorograph and an example of the densitometric profiles is shown in Figure 2C. By comparing the number and the height of peaks corresponding to the major labeled protein bands on the fluorograph, we did not observe a difference in number of peaks between control and AD. However, the height of the 160 kDa peak is significantly lower in AD (Fig. 2C, peak 3), confirming the visual inspection of the fluorograph. A similar reduction in the radiolabeled 160 kDa band was seen in all five AD frontal cortices examined (data not shown).

A protein with an \( \sim 50 \) kDa size was among other proteins labeled by \([3H]\)galactose (Fig. 2B, indicated with arrow). In some of the AD cases studied, the intensity of this band was increased, and the band appeared smeared. However, this finding was not seen in every AD case and did not seem to be related to the age or postmortem delay of the cases used for the study (\( n = 5 \); Table 1). Moreover, this 50 kDa protein seems unlikely to be modified by O-linked GlcNAc (see description below for Fig. 5). Therefore, it was not investigated further in this study.

Middle frontal gyrus

The above-observed reduction of glycosylated 160 kDa protein in AD was apparent in the frontal cortex, a grossly defined brain region. To confirm this observation, O-GlcNAc-modified proteins from a more limited brain area, the middle frontal gyrus, were studied. The middle frontal gyrus is among the brain regions generally spared of certain characteristic AD pathologies. It has selected aspects of the AD pathology such as diffuse plaques and gliosis but does not contain NFTs (Braak et al., 1989). We examined the O-GlcNAcylated 160 kDa protein in the anterior cerebellum from the same 11 AD and 10 control cases. Figure 4 illustrates the fluorograph results from two of these cases, and Figure 4B shows a typical densitometric scanning profile of the fluorographs. Unlike the frontal cortex (Fig. 2) and the middle frontal gyrus (Fig. 3), the 160 kDa protein was not the major galactosylated protein even in non-AD normal cerebellum (Fig. 4, A, arrowhead, B, peak 3). Furthermore, in AD cerebellum, neither overall GlcNAc level nor glycosylated 160 kDa protein are significantly affected (Fig. 4C).

Characterization of the saccharides that modify the 160 kDa protein

The above results suggested that among a number of GlcNAc-containing proteins from brain extracts, the level of the 160 kDa glycoprotein was reduced considerably and was inversely correlated to the number of NFTs. We therefore chose to focus on the 160 kDa protein and further characterized its glycosylation.

Galactosylated brain proteins were first digested with PNGase F that cleaves N-linked structures at an asparagine–GlcNAc bond (Plummer and Tarentino, 1981). Figure 5A shows that heavily labeled 160 kDa glycoprotein was unaffected by PNGase F treatment in either AD or control samples (arrowhead), suggesting it is not N-linked glycosylation. In contrast, several other labeled glycoproteins in the samples, including the 50 kDa glycoprotein (arrow), contained N-linked radiolabeled saccharides, because they were no longer visible after PNGase treatment. Longer fluorographic exposures of the gel substantiated this observation (data not shown). Ovalbumin, containing N-linked terminal GlcNAc moieties, was used as a positive control for the PNGase F digestion. Figure 5B shows the lack of proteases in the PNGase F digestion, because the amount of protein shown on Coomassie blue-stained gels was not reduced with PNGase F treatment. Counts of gel slices corresponding to ovalbumin and to the 160 kDa protein revealed that 94% of the \([3H]\)galactose-labeled ovalbumin was released by PNGase F, whereas only 4% of labeling was released from the 160 kDa glycoprotein (Fig. 5C).

To confirm that the sugar moieties that modify the 160 kDa protein were in the form of O-linked GlcNAc, the samples were subjected to alkaline-induced \( \beta \)-elimination, a common method used to release O-linked saccharides from protein backbones (Zinn et al., 1977). The \( ^3\)H-labeled 160 kDa protein resolved by SDS-PAGE was transferred to a PVDF membrane, and the corresponding membrane strip was cut out and subjected to \( \beta \)-elimination as described in Materials and Methods. The radioactivity from the 160 kDa PVDF membrane strip had essentially been eliminated at the end of \( \beta \)-elimination compared with non-\( \beta \)-eliminated sample, whereas \( ^3\)H labeling of ovalbumin remained unchanged (Fig. 5D). The \( \beta \)-elimination products released from the PVDF membrane strips were further analyzed on a Sephadex
Figure 3. A marked and significant reduction of glycosylated 160 kDa protein occurs in AD middle frontal gyrus. Proteins from middle frontal gyrus of AD and age-matched controls were analyzed with galactosyltransferase labeling. The labeled products were revealed with SDS–PAGE followed by fluorography. A, Fluorographs of labeled proteins from two representative AD and control cases. Arrowheads indicate the 160 kDa protein. B, Densitometric scanning profiles of control case 1 and AD case 1 shown in A. Peak 3 represents the 160 kDa protein band. C, Densitometric values gathered from two bands on the scanned autofluorographs, the intensity of which was relatively consistent among all AD and control samples (peaks 4, 8), were used for normalization between fluorographs. Densitometric values (optical density) for the 160 kDa 3H-galactosylated protein from all 11 cases of AD and 10 cases of non-AD controls were analyzed by Student’s t test. Data are expressed as mean ± SEM. D, Densitometric values of six major 3H-galactosylated protein bands other than the 160 kDa protein from the same cases were also analyzed by Student’s t test. None of the AD versus control comparisons were statistically significant. E, Relation between O-glycosylated 160 kDa protein and NFT density of the same tissues.
G-50 gel filtration column, revealing the 160 kDa protein as a single included volume peak (Fig. 5E). The released products by β-elimination also comigrated with a disaccharide standard, Galβ1,4GlcNAcitol in thin-layer chromatography (Fig. 5F, standard b). These analyses demonstrated that the 160 kDa protein is modified by O-linked single GlcNAc residues. Analyses of saccharide–protein linkage and saccharide structure of all three brain regions (frontal cortex, middle frontal cortex, and cerebella cortex) of both control and AD demonstrated the same results.

Identification of the 160 kDa O-GlcNAcylated protein
The unidentified 160 kDa O-GlcNAcylated protein demonstrated several characteristics: it was enriched in the soluble fraction of the brain extracts; it migrated at ~160 kDa on SDS–PAGE; it was weakly stained with Coomassie blue; and it was extensively modified by O-GlcNAc. These features appear to be shared by a known neuronal O-GlcNAcylated protein, the clathrin assembly protein AP-3 (Murphy et al., 1991). When [3H]galactose-labeled proteins from brain extracts were analyzed with immunoblotting using an AP-3-specific antibody, AP180-I mAb (Ahle and Ungewickell, 1986), a major immunoreactive protein band was revealed at 160 kDa (Fig. 6A, left panel). Fluorographing of the same blot (Fig. 6A, right panel) showed that the 160 kDa [3H]-labeled protein band was superimposed with the major AP180-I mAb-immunoreactive protein band (Fig. 6A, arrowheads). AD tissues demonstrated a similar result, although the immunoreactive signal to the antibody was much weaker.

Two other AP-3-specific monoclonal antibodies, CLAP3 and F1–20, were also used in immunoblotting of [3H]-labeled proteins.
Figure 5. The 160 kDa protein is modified by O-linked single N-acetylglucosamine residues. [3H]Galactose-labeled proteins were deglycosylated by PNGase F and resolved on 10% SDS-PAGE. A, Fluorographs demonstrate labeled proteins that are resistant to PNGase F digestion. The arrowhead indicates the 160 kDa protein, and the arrow indicates the 50 kDa protein. The migration positions of molecular weight standards (kilodaltons) are indicated. B, Coomassie blue-stained gels reveal the overall protein pattern. C, The 160 kDa band and ovalbumin band in each group were cut out and counted. D, [3H]Galactose-labeled sample proteins were separated by SDS-PAGE and transferred to a PVDF membrane. The membrane strip corresponding to the position of 160 kDa was cut out, subjected to alkaline-induced \( \beta \)-elimination, and counted. Ovalbumin was used as a control. E, G-50 Sephadex column profile of \( \beta \)-elimination products released from the PVDF membrane. F, Saccharides released by \( \beta \)-elimination were also analyzed by thin layer chromatography using a cellulose-coated plastic plate. The lanes of 160 kDa protein from AD and control were cut separately into 0.5-cm-wide strips and counted. Disaccharide standards were revealed by AgNO\(_3\) staining, and their migration positions are indicated by arrows. Closed circles, Control; open circles, AD. All above analyses were performed at least twice on proteins from all three brain regions examined from both AD and controls. These data demonstrate that the 160 kDa protein is modified by O-linked single GlcNAc residues.
Both CLAP3 mAb (Murphy et al., 1991) and F1–20 mAb (Sousa et al., 1990; Zhou et al., 1993) have been shown to react with AP-3 specifically, and both, in this study, revealed a prominent band at 160 kDa (Fig. 6B, lanes 1, 2, arrowhead). Several minor bands that migrated faster than the 160 kDa band are seen to a variable extent with different antibodies and likely represent degradation products of AP-3. AP-3 is known to be very susceptible to proteolysis (Ahle and Ungewickell, 1986; Zhou et al., 1993).

The same 3H-labeled samples were also immunoblotted with an antibody that specifically recognizes a subunit of another clathrin assembly protein, AP-2 (Ahle et al., 1988). As seen in Figure 6B (lane 3), the AP-2-specific antibody (100/2 mAb) failed to react with the 160 kDa glycoprotein. Although it is often difficult to distinguish between AP-3 and clathrin heavy chain protein on SDS-PAGE (Ahle and Ungewickell, 1986; Murphy et al., 1991; Zhou et al., 1993), the 160 kDa glycoprotein is unlikely clathrin heavy chain, because a heavy chain-specific monoclonal antibody (CHC 5.9) did not reveal an immunoreactive protein band that can be superimposed with the 3H-labeled 160 kDa protein band (Fig. 8A). The corresponding fluorograph shows a doublet-spot at 160 kDa, and pI 5.2–5.4, consistent for the known pI for AP-3 (Ahle and Ungewickell, 1986) (Fig. 8A). Western blot analysis of a duplicate gel showed that the doublet spot was specifically immunoreactive to the AP180-I mAb (Fig. 8B), supporting the evidence that the protein is AP-3. It is not known why there is a doublet showing in the fluorograph and in the blot instead of a single spot. Human AP-3 proteins may be present as two different isoforms, which has yet to be determined, or perhaps not all AP-3 molecules contain the same amount of sugar residues. The extra spot detected by Western blotting (Fig. 8B, arrow) could be a protease fragment of AP-3.

Finally, protein extracts from middle frontal gyri of all 11 AD and 10 control cases were immunoprecipitated and labeled with [3H]galactose. Figure 9A is the Coomassie blue-stained gel showing immunoprecipitates from two representative AD and control brains. Whereas IgG heavy chain was the major protein band stained with Coomassie blue, the AP-3 immunoprecipitates were not visible in most cases. In contrast, the fluorograph of the same gel demonstrated heavily glycosylated AP-3 immunoprecipitates (Fig. 9B). The intensity of 3H-labeled protein bands was markedly reduced in AD. The amount of AP180-I mAb was not a limiting factor, because there was no AP-3 immunoreactivity present in...
The 160 kDa protein is labeled with \([^{3}H]\)galactose and galactosyltransferase. A. Immunoprecipitation of brain extracts was performed using monoclonal anti-AP-3 antibody AP180-I, and the resulting immunoprecipitates were subsequently labeled with \([^{3}H]\)galactose as described in Materials and Methods. Coomassie blue staining of the immunoprecipitates revealed a barely visible band at 160 kDa (arrowhead). Fluorograph (two-day exposure) of the same Coomassie blue-stained gel is on the right, showing that the immunoprecipitated 160 kDa protein using the AP180-I mAb was heavily labeled with \([^{3}H]\)galactose. B. Immunoprecipitation was prepared using AP180-I mAb or 100/2 mAb (anti-AP-2). The resulting supernatants (S) and pellets (P) were analyzed by immunoblotting using AP180-I mAb. The 100/2 mAb was used as a control antibody because, like AP180-I mAb, it is IgG2 from mouse ascites fluid. Whereas the majority of AP-3 was present in the pellets when AP180-I mAb was used for immunoprecipitation, no AP-3 was detected in the pellets with the use of 100/2 mAb (lanes 3, 4).

**Figure 7.** Immunoprecipitated 160 kDa protein is labeled by \([^{3}H]\)galactose and galactosyltransferase. A. Immunoprecipitation of brain extracts was performed using monoclonal anti-AP-3 antibody AP180-I, and the resulting immunoprecipitates were subsequently labeled with \([^{3}H]\)galactose as detailed in Materials and Methods. Coomassie blue staining of the immunoprecipitates revealed a barely visible band at 160 kDa (arrowhead). Fluorograph (two-day exposure) of the same Coomassie blue-stained gel is on the right, showing that the immunoprecipitated 160 kDa protein using the AP180-I mAb was heavily labeled with \([^{3}H]\)galactose. B. Immunoprecipitation was prepared using AP180-I mAb or 100/2 mAb (anti-AP-2). The resulting supernatants (S) and pellets (P) were analyzed by immunoblotting using AP180-I mAb. The 100/2 mAb was used as a control antibody because, like AP180-I mAb, it is IgG2 from mouse ascites fluid. Whereas the majority of AP-3 was present in the pellets when AP180-I mAb was used for immunoprecipitation (lanes 1, 2), no AP-3 was detected in the pellets with the use of 100/2 mAb (lanes 3, 4).

**Figure 8.** Two-dimensional gel analysis of AP-3 labeled with \([^{3}H]\)galactose. Immunoprecipitates of AP-3 were labeled with \([^{3}H]\)galactose as described in Materials and Methods. The samples were then equally divided and resolved separately on duplicated 2-dimensional gels. One gel was subjected to fluorography (A), showing labeled AP-3 (10 d exposure). The other gel was analyzed with immunoblotting using the AP180-I mAb (B). The acid edge is to the left. The asterisk indicates the position of the internal standard, tropomyosin (pl, 5.2; molecular weight, 32.7 kDa). Positions of protein markers (in kilodaltons) are indicated between the panels. The AP180-I mAb-immunoreactive spot that was not \(^3\)H-galacosylated is indicated with an arrow.

the supernatant after immunoprecipitation (data not shown). Taken together, these results suggest that the O-GlcNAcylated 160 kDa protein is AP-3 and O-GlcNAcylated AP-3 is reduced significantly in selected brain regions affected by AD.

**DISCUSSION**

In this study, we detected and compared O-GlcNAc-modified proteins in brain extracts from confirmed AD and non-AD age-matched controls and made several observations. First, in normal-aged human brains, a number of proteins possess terminal O-linked GlcNAc residues. Among them, a protein with molecular mass of ~160 kDa on SDS-PAGE gels is glycosylated prominently. This distinct and reproducible glycosylation pattern is more evident in proteins from neocortical regions (frontal cortex and middle frontal gyrus) than in cerebellum. Second, in AD brains, although other GlcNAc-modified proteins show no dramatic changes when compared with controls, there is a substantial decrease in the levels of the 160 kDa glycoprotein. The reduction appears to correlate with the number of NFTs. Third, based on the results of immunoblotting, immunoprecipitation, and two-dimensional electrophoresis analysis, the 160 kDa glycoprotein is identified as AP-3, one of the clathrin assembly proteins known to contain O-GlcNAc (Murphy et al., 1994).

Galactosyltransferase has been commonly used as a specific probe to detect GlcNAc on purified proteins (Dong et al., 1993; Arnold et al., 1996), on immunoprecipitates (Chou and Omary, 1993; Haltiwanger and Philipsberg, 1997), on protein mixtures from cell subfractions (Kearse and Hart, 1991; Haltiwanger and Philipsberg, 1997). We used this galactosyltransferase-labeling method to examine O-GlcNAc-modified proteins in human post-mortem brains. In addition, we also attempted to detect O-GlcNAc-bearing proteins in brain extracts using anti-O-GlcNAc antibodies (Turner et al., 1990), as well as lectin wett germ agglutinin (WGA) (Monsigny et al., 1980). We found that neither the anti-O-GlcNAc antibodies nor the lectin WGA were suitable for studying O-GlcNAc glycosylation in tissue extracts. Both reagents gave unacceptable high background and/or were not adequately specific (data not shown). In contrast, the galactosyltransferase labeling method allowed us to obtain information on GlcNAc levels and the overall profile of GlcNAc-bearing proteins in the crude brain extracts. Further analyses of the galactosyltransferase labeled products with PNGase F digestion and alkali-induced β-elimination enabled us to demonstrate that the 160 kDa protein is glycosylated by monosaccharide GlcNAcs through O-glycosidic linkages (Fig. 5). Although this galactosylation technique is a better method for detecting O-GlcNAc (Roquemore et al., 1994; Haltiwanger et al., 1997), it detects only accessible terminal GlcNAcs and therefore may underestimate total GlcNAcs.

This study does not provide evidence of a direct link between O-GlcNAc glycosylation and the abnormal hyperphosphorylation
immunoreactive to three different AP-3 specific antibodies but can be modified by a component of NFTs in AD brains (e.g., Goedert 1993). That tau hyperphosphorylated tau is well documented as being a major 160 kDa protein, the AD brains we examined were essentially no different from the controls in the gross level of terminal GlcNAc. Hyperphosphorylated tau is well documented as being a major component of NFTs in AD brains (e.g., Goedert 1993). That tau can be modified by O-GlcNAc has been demonstrated, albeit only on purified bovine tau (Arnold et al., 1996). If there were a direct competition between phosphorylation and O-GlcNAcylation on tau proteins, as has been shown on some proteins (Kelly et al., 1993; Hart et al., 1995), then it might be anticipated that O-GlcNAc level on abnormal hyperphosphorylated tau would be different from tau with normal levels of phosphate (Hart et al., 1995; Arnold et al., 1996; Haltiwanger et al., 1997). However, we did not observe differences in GlcNAc levels on proteins with molecular mass between 40 and 60 kDa (Figs. 2, 3), the range in which various tau isoforms migrate on SDS-PAGE gels (Greenberg and Davies, 1990; Bramblett et al., 1993). Although the brain extracts used in the study did contain both total tau and hyperphosphorylated tau proteins (P. J. Yao and P. D. Coleman, unpublished data), we do not exclude the possibility that the approach used in this study may not reveal changes in individual O-GlcNAc glycoproteins unless the protein is either very abundant in the brain extracts or is heavily glycosylated.

The clearly evident and consistent change found is the significant reduction of O-GlcNAcylated 160 kDa protein in end-stage AD neocortices. The limited number of AD cases we examined suggests a negative correlation between the glycosylated 160 kDa protein and the density of NFTs. This 160 kDa glycoprotein strikingly resembles clathrin assembly protein AP-3 in biochemical characteristics (Ahle and Ungewickell, 1986; Kohtz and Puszkin, 1988; Lindner and Ungewickell, 1992; Zhou et al., 1993; Murphy et al., 1994): it is from the soluble fraction of brain extracts and is abundant in brain; it migrates with an ~160 kDa molecular mass on SDS-PAGE gels; it is poorly stained by Coomassie blue; and it is glycosylated extensively with O-GlcNAc. The identity of the 160 kDa glycoprotein as AP-3 is further supported by several lines of evidence. First, the [3H]galactose-labeled 160 kDa protein from brain extracts was immunoreactive to three different AP-3 specific antibodies but not to antibodies against other clathrin-coated vesicle proteins such as AP-2 and clathrin heavy-chain protein (Fig. 6). Second, using one of the AP-3-specific antibodies (AP180-I mAb), an immunoprecipitation product with a molecular mass of 160 kDa was the only protein band labeled with [3H]galactose (Fig. 7). Furthermore, two-dimensional SDS-PAGE demonstrated that the pI for the immunoprecipitated protein was ~5.2, in agreement with the value known for AP-3 (Ahle and Ungewickell, 1986) (Fig. 8). Finally, quantitative immunoprecipitation revealed that O-GlcNAcylated AP-3 in AD brains was markedly reduced (Fig. 9).

AP-3, a synapse-specific protein, was independently discovered by several laboratories but given different names, including pp155 (Keen and Black, 1986), AP180 (Ahle and Ungewickell, 1986), NP185 (Kohtz and Puszkin, 1988), and F1–20 (Zhou et al., 1993). It is known now that all these proteins are the same and therefore have been renamed AP-3. AP-3 is very effective in promoting the assembly of clathrin into clathrin-coated vesicles (Lindner and Ungewickell, 1992) and probably plays an important role in synaptic vesicle recycling (Maycox et al., 1992; Lafer et al., 1997). Measurement of the saccharide content indicated that 1 mol of AP-3 contained ~1 mol of GlcNAc (Murphy et al., 1994), which is ~10 times more than the amount of GlcNAc on neurofilaments (Dong et al., 1993). This is consistent with our observation that AP-3 is labeled with more [3H]galactose than any other protein in the brain extracts tested. As with other O-GlcNAc-modified proteins, the specific function of this modification on AP-3 remains unknown. In clathrin assembly assay, the interaction of AP-3 with clathrin was not affected by in vitro galactosylation of AP-3 (Murphy et al., 1994). Moreover, bacterially expressed AP-3 that lacks O-GlcNAcyl appears to be fully capable of clathrin binding and assembly (Ye and Lafer, 1995). These results suggest that GlcNAc residues may modulate AP-3 protein in other ways than its association with clathrin.

Because of its poor affinity for Coomassie blue, it is not apparent whether the observed diminution in O-GlcNAcylated AP-3 is attributable to a decrease in the level of sugar, in the level of protein itself, or both. Nonetheless, our studies with Western blotting using AP-3-specific antibodies (Fig. 6A) and additional

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**Figure 9.** Glycosylated AP-3 is reduced significantly in AD brains. Extracts from middle frontal gyri of AD and age-matched controls were immunoprecipitated using AP180-I mAb and subsequently labeled with [3H]galactose. A, Coomassie blue-stained gel; B, corresponding fluorograph. Lanes 1 and 2 are two representative controls, and lanes 4 and 5 are two AD cases. The migration positions of IgG heavy chain and AP-3 are indicated. Positions of protein markers (lane 3 in both panels) are indicated between the panels.
unpublished results clearly show the level of the AP-3 protein was reduced in AD brains. It seems unlikely that the 2.5-fold reduction in glycosylated AP-3 in AD neocortex is attributable solely to loss of neocortical neurons. In the same brain extracts used for our study, we see little, if any, parallel changes in the levels of tau proteins and of neurofilament medium and high molecular mass (Yao and Coleman, unpublished observations), indicating relatively unchanged levels of these neuronal markers. Morphological data consistent with this conclusion come from a recent study using unbiased stereological methods demonstrating a statistically nonsignificant 6% loss of neocortical neurons in AD (Reger et al., 1994). Although other recent studies have shown major neuronal loss in selected regions of AD brain (for review, see Morrison and Hof, 1997), we did not sample from any of these selectively vulnerable regions.

Because AP-3 is a synaptic protein associated with clathrin-coated vesicles (Lindner and Ungewickell, 1992; Zhou et al., 1993; Lafer et al., 1997), the decline of this glycoprotein may be part of the synapse loss that occurs in AD. In fact, electron microscopic (Scheff et al., 1990; Scheff and Price, 1993) and immuno-histochemical (Masliah et al., 1989) determinations of neocortical synapse density in AD showed a decrease of ≤50%, which is less than the loss in AP-3 found by us. This suggests that there may be a loss of clathrin-coated vesicles in AD in addition to the loss of synapses. Thus, the loss of glycosylated AP-3 may be an earlier event in the pathological cascade of synaptic loss in AD.

The apparent decline of glycosylated AP-3 protein in AD may be related to the modification by O-GlcNAc. The level of O-GlcNAc may change the susceptibility of a protein to proteases and thus modulate its stability. Han and Kudlow (1997) showed a direct correlation between reduced O-GlcNAc level on transcription factor Sp1 and increased proteolytic degradation. This possibility is particularly intriguing because AP-3 is known for its high susceptibility to proteolytic degradation (Kohtz and Puskin, 1988; Murphy et al., 1991; Lindner and Ungewickell, 1992; Zhou et al., 1993). If indeed the glycosylation state modifies the vulnerability of AP-3 to proteolysis, then hypoglycosylation may be responsible for the degradation and loss of AP-3 in AD. Insufficient AP-3 at the nerve terminals may therefore result in abnormal clathrin-coated vesicle assembly and consequently affect synaptic vesicle recycling. We are currently investigating this potential function for O-GlcNAc residues on AP-3, and we hope to gain a better understanding of the role of O-GlcNAc modification and the mechanisms of synaptic vesicle recycling in AD pathology.

REFERENCES


