A novel role for synaptic acetylcholinesterase as an apoptotic deoxyribonuclease

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In addition to terminating neurotransmission by hydrolyzing acetylcholine, synaptic acetylcholinesterase (AChES) has been found to have a pro-apoptotic role. However, the underlying mechanism has rarely been investigated. Here, we report a nuclear translocation-dependent role for AChES as an apoptotic deoxyribonuclease (DNase). AChES polypeptide binds to and cleaves naked DNA at physiological pH in a Ca2+–Mg2+-dependent manner. It also cleaves chromosomal DNA both in pre-fixed and in apoptotic cells. In the presence of a pan-caspase inhibitor, the cleavage still occurred after nuclear translocation of AChES, implying that AChES-DNase acts in a CAD- and EndoG-independent manner. AChE gene knockout impairs apoptotic DNA cleavage; this impairment is rescued by overexpression of the wild-type but not (aa 32–138)-deleted AChES. Furthermore, in comparison with the nuclear-localized wild-type AChES, (aa 32–138)-deleted AChES loses the capacity to initiate apoptosis. These observations confirm that AChES mediates apoptosis via its DNase activity.

Keywords: synaptic acetylcholinesterase; nuclear translocation; DNA-binding protein; DNA cleavage; deoxyribonuclease; apoptosis

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Introduction

Synaptic acetylcholinesterase (AChES), similar to the other two variants of AChE (erythrocytic AChE and read-through AChE (AChER)), belongs to the type-B carboxylesterase/lipase family. The three different isoforms are encoded by a single gene, AChE, but because of alternative splicing at the 3′ region of acetylcholinesterase messenger RNA, the three variants differ in their carboxy-terminal sequences [1]. Erythrocytic AChE is expressed primarily in erythroid tissues, where it associates with membranes via the phosphoinositide moieties added posttranslationally. AChER is expressed in embryonic and tumor cells, and is thought to be involved in the stress response and, possibly, inflammation [2]. AChES is the major form of acetylcholinesterase found in brain, muscle, and other tissues. The classical role of AChES is to terminate neurotransmission by hydrolyzing acetylcholine at cholinergic synapses and neuromuscular junctions [3]. In addition, the enzyme has an important role in apoptosis of various types of cells. Studies in vitro and in vivo, have shown that AChES is upregulated in response to various apoptotic stimuli and that apoptosis is attenuated by knockdown of its expression either by antisense RNA, small interfering RNA, or by heterozygous deletion of the AChE gene [4–6]. These results demonstrate a pro-apoptotic function of AChES, although the underlying mechanism remains to be elucidated.

Apoptosis is essential for many biological processes [7]. Distinct deoxyribonucleases (DNases) participate in apoptosis by catalyzing the hydrolytic cleavage of phosphodiester linkages in the DNA backbone. DNase I is the first enzyme to be recognized in mammalian cells to cleave nuclear DNA during apoptosis [8, 9].
However, DNase I-deficient JA3 cells are still capable of undergoing DNA fragmentation in response to treatment with an anti-Fas antibody [9]. This effect can be attributed to other DNases, such as the 40-kDa DNA fragmentation factor (CAD/DFF40) and endonuclease G (EndoG).

Figure 1 Synaptic acetylcholinesterase (AChE<sub>S</sub>) initiates apoptosis in the nuclear compartment. (a) Time-lapse images showing the alteration of the distribution of the indicated proteins during apoptosis. HeLa cells were transfected with the plasmids expressing the indicated fusion proteins. After 24 h, cells were exposed to 100 μM H<sub>2</sub>O<sub>2</sub>, and time-lapse images were captured using a Leica AS MDW live cell image acquisition system. See also Supplementary Video S1a. (b) Western blot analysis of fusion protein expression. HeLa cells were transfected with the empty plasmid as a control or the indicated fusion protein expression plasmids. After 24 h, cells were harvested. Whole-cell extracts were prepared, resolved by SDS–polyacrylamide gel electrophoresis, and blotted with anti-GFP antibody (Abmart, M20004). (c) Confocal images showing deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling (TUNEL) staining in HeLa cells transfected with the plasmids expressing the indicated proteins. At 48 h after transfection, TUNEL assays were performed. Cells expressing green fluorescence protein (GFP) alone were used as the empty plasmid control, and those expressing histone H2B were used as the negative control. Scale bars, 15 μm. In the detailed image, the cells are indicated by an arrowhead (right panel; ×2 magnification). (d) DNA ladder assay for apoptosis triggered by the indicated proteins overexpressed in HeLa cells for 48 h. (e) Images showing the GFP-positive HeLa cells. At 18 h after transfection, GFP-positive HeLa cells were sorted by fluorescence-activated cell sorting. At 72 h after sorting, the images were taken using an inverted fluorescence microscope. Scale bar, 30 μm. (f) The viability curves of GFP-positive HeLa cells overexpressing the indicated proteins after sorting. The relative number of cells at the indicated time points was determined by the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazoliumbromide (MTT) assay, with measurement of optical density (OD) at 570 nm. The data are presented as the mean ± s.d. of triplicate samples from one experiment that is a representative of three independent experiments. NLS, nuclear localization signal.
CAD/DFF40, the main effector involved in the apoptotic degradation of nuclear DNA into oligonucleosomal fragments, is a caspase-3-dependent DNase. After activated caspase-3-specific cleavage of its inhibitor ICAD/DFF45, CAD/DFF40 is released from its heterodimeric complex and enters the nucleus to cleave DNA by introducing double-stranded breaks; in the absence of activated caspase-3, CAD/DFF40 was inactive and confined to the cytoplasm by binding with ICAD/DFF45 [10, 11].

EndoG, which is another caspase-dependent apoptotic DNase, is localized in the mitochondrion. After caspase activation in response to apoptotic insults, EndoG is released from mitochondria and translocates to the nucleus where it causes DNA degradation [12]. In DFF45-deficient cells, nuclear-translocated EndoG contributes to the residue of nucleosomal DNA fragments [12].

In the process of stepwise DNA degradation, CAD/DFF40 and EndoG function at the early stage [10, 13], whereas DNase II functions at the later stage and is required for complete degradation of the fragments [14]. Among these well-known DNases, CAD/DFF40 is considered to account for the majority of the nuclease activity responsible for chromosomal DNA fragmentation [15, 16]. Although multiple DNases involved in chromatin DNA degradation have been reported, further DNases remain to be identified. Here, we report that AChE<sub>S</sub> mediates cell apoptosis by acting as a CAD- and EndoG-independent DNase.

Results

The nucleus is the optimal subcellular compartment for the pro-apoptotic function of AChE<sub>S</sub>

The subcellular localization of a protein is closely related to its function. Therefore, we first investigated alterations in the distribution of AChE<sub>S</sub> protein during apoptosis. When treated with hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>, one of the reactive oxygen species inducing oxidative stress), the cells underwent apoptosis, as shown by caspase-3 activation (Supplementary Figure S1A). In addition, upregulated AChE protein expression and its nuclear translocation were observed.
These data further support those obtained in our previous studies [4, 5]. The AChE-specific antibody used in our study recognizes the common peptide fragments of all the three AChE variants. To investigate whether the variant AChES translocates into the nucleus in response to apoptotic stimuli, we constructed the pEGFP–AChES plasmid, encoding the AChES protein fused with green fluorescence protein (GFP) at its C terminus. Time-lapse imaging showed that AChES nuclear translocation closely accompanied the morphological changes during apoptosis induced by H2O2 (100 μM) in HeLa cells, suggesting that AChES has a pro-apoptotic role in the nucleus (Figure 1a; Supplementary Video S1a). To further confirm this suggestion, we constructed the pEGFP–NLS–AChES plasmid, encoding the AChES protein fused with a nuclear localization signal (NLS) at its N terminus and GFP at its C terminus (Figure 1b). Overexpression of NLS–AChES–GFP stimulated DNA breakage, which was examined by deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling (TUNEL) (Figure 1c). In contrast, this effect was not stimulated by the overexpression of AChES–GFP without the NLS (Figures 1c and d). Furthermore, chromosomal DNA cleavage was detected as a DNA ladder pattern, a biochemical characteristic of apoptosis, by gel electrophoresis (Figure 1d), thus, demonstrating that the cell death induced by NLS–AChES overexpression occurred through apoptosis. Subsequently, 3-(4,5-dimethyl thiazol-2-yl)-2,5-diphenyltetrazoliumbromide (MTT) assays showed that although the GFP and AChES–GFP fusion protein hardly affected cell survival, NLS–AChES–GFP stimulated gradual apoptosis (Figures 1e and f). These results demonstrate that AChES promotes apoptosis only after translocation into the nucleus. The nucleus is the optimal subcellular compartment for the pro-apoptotic function of AChES.

Figure 2 Interaction between synaptic acetylcholinesterase (AChES) polypeptide with plasmid DNA. (a) Silver staining and western blot showing the purified human AChES polypeptide, hAChE-T547. The anti-AChE antibody provided by Dr Palmer Taylor was used at a dilution of 1:1000. (b) Cholinesterase activity of hAChE-T547 examined by the Ellman assay, with measurement of optical density (OD) 405 nm. Data shown represent one of three independent experiments. Values represent the mean ± s.d. of triplicate samples. (c) The interaction between hAChE-T547 and pEGFP-c1 plasmid DNA examined by Biacore T100. DNase I and bovine serum albumin (BSA) were used as positive and negative controls, respectively. Data shown represent one of three independent experiments.

(Supplementary Figure S1). These data further support those obtained in our previous studies [4, 5]. The AChE-specific antibody used in our study recognizes the common peptide fragments of all the three AChE variants. To investigate whether the variant AChES translocates into the nucleus in response to apoptotic stimuli, we constructed the pEGFP–AChES plasmid, encoding the AChES protein fused with green fluorescence protein (GFP) at its C terminus. Time-lapse imaging showed that AChES nuclear translocation closely accompanied the morphological changes during apoptosis induced by H2O2 (100 μM) in HeLa cells, suggesting that AChES has a pro-apoptotic role in the nucleus (Figure 1a; Supplementary Video S1a). To further confirm this suggestion, we constructed the pEGFP–NLS–AChES plasmid, encoding the AChES protein fused with a nuclear localization signal (NLS) at its N terminus and GFP at its C terminus (Figure 1b). Overexpression of NLS–AChES–GFP stimulated DNA breakage, which was examined by deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling (TUNEL) (Figure 1c). In contrast, this effect was not stimulated by the overexpression of AChES–GFP without the NLS (Figures 1c and d). Furthermore, chromosomal DNA cleavage was detected as a DNA ladder pattern, a biochemical characteristic of apoptosis, by gel electrophoresis (Figure 1d), thus, demonstrating that the cell death induced by NLS–AChES overexpression occurred through apoptosis. Subsequently, 3-(4,5-dimethyl thiazol-2-yl)-2,5-diphenyltetrazoliumbromide (MTT) assays showed that although the GFP and AChES–GFP fusion protein hardly affected cell survival, NLS–AChES–GFP stimulated gradual apoptosis (Figures 1e and f). These results demonstrate that AChES promotes apoptosis only after translocation into the nucleus. The nucleus is the optimal subcellular compartment for the pro-apoptotic function of AChES.
The purified AChEs polypeptide binds to plasmid DNA in vitro

The intriguing question of why nucleus-localized AChEs triggers apoptosis remains to be answered.
AChE₈ acts as an apoptotic DNase

with chromatin DNA cleavage (such as CAD/DFF40 and EndoG) [11, 18]. We aimed to determine whether AChE₈ hydrolyzes DNA.

First, the DNA-binding capacity of AChE₈ was examined. The amino-acid (aa) 1–574 region is shared by all three variants of human AChE, and aa 36–574 is...
the putative region used to investigate the acetylcholine esterase activity of AChE and its crystalline structure [19]. Therefore, human AChES aa 32–578 (hAChE-T547) was overexpressed and purified (Figure 2a). As expected, the Ellman assay showed that hAChE-T547 exhibited acetylcholine-hydrolyzing activity (Figure 2b). Importantly, similar to DNase I, hAChE-T547 bound plasmid DNA, although bovine serum albumin (Sigma) did not (Figure 2c). Accurate measurement of the dissociation constant (Kd) was rendered impractical because the plasmid solutions in the mobile phase became too ‘sticky’ to be injected onto the chip at concentrations higher than 1.88 × 10^{-6} M. Nevertheless, the data indicated the binding of hAChE-T547 to DNA, which prompted us to determine the capacity of AChE to digest DNA.

**The purified human AChES polypeptide cleaves naked DNA at physiological pH in a Ca\(^{2+}\)-Mg\(^{2+}\)-dependent manner**

After incubation with pEGFP-c1 plasmid DNA in hydrolysis buffer (5.0 mM Tris-HCl, pH 7.5; 2.5 mM CaCl\(_2\); 5.0 mM MgCl\(_2\)) at 37 °C for 6 h, hAChE-T547 converted the supercoiled DNA to the nicked and linear forms in a dose- and time-dependent manner (Figures 3a and b). The cleavage pattern of plasmid DNA was the same as that resulting from cleavage by CAD/DFF40 [20], EndoG [21], DNase I [22], and tDCR-1 (a functional analog of DFF40 in Caenorhabditis elegans) [10]. As expected, the amount of the products correlated directly with the amount of the substrate plasmids (Figure 3c). Furthermore, the optimal pH for cleavage by the peptide was found to be 7.5 (Figure 3d), similar to those of CAD/DFF40 [23], EndoG [21], DNase I [24], DNase γ [25], and tDCR-1 [10].

Apoptotic DNases are usually activated by divalent metal ions [26]. hACh-E-T547 showed very weak DNA cleavage activity in the presence of Mg\(^{2+}\) (1.25–5 mM) alone, and the cleavage was hardly detected in the presence of Ca\(^{2+}\) (1.25–5 mM) only (Figure 3e). However, the combination of Mg\(^{2+}\) and Ca\(^{2+}\) showed an obvious co-activating effect on the enzyme, which was further confirmed by the observation that either the Ca\(^{2+}\)-chelator ethylene glycol tetraacetic acid or the versatile chelating agent EDTA markedly inhibited the cleavage (Figure 3e). These data indicated a Ca\(^{2+}\)-Mg\(^{2+}\)-dependent DNA cleavage activity of AChES.

Human AChE-T547 was overexpressed in HEK 293S stable cells, secreted into and purified from the cell culture medium. It is noteworthy that DNase I is also a secreted protein detected in most body fluids, including serum [27, 28]. In spite of this, G-actin, a specific inhibitor of DNase I, inhibited DNA cleavage activity of DNase I but not that of hAChE-T547 (Figure 3f). Thus, the DNA degradation was not induced by contaminating DNase I.

Despite this, AChES-DNase activity was relatively low compared with that of DNase I (Figures 3e and f), which is ubiquitously expressed in mammalian tissues. However, the enzyme (Sigma) used in our work was obtained from bovine pancreas, the most efficient DNase, which mainly is a digestive enzyme. In view of this, it is not surprising that AChES-DNase activity is much lower than that of DNase I. To further determine whether it is normal for an apoptotic DNase to show such low activity as that of AChES-DNase, we compared the DNA degradation capacity of AChE-T547 and CAD. The purified human CAD polypeptide (Arg87–Lys323; predicted molecular mass 31.9 kDa), was purchased from USCN Life Science, China. Silver-stained gels showed a band with a molecular weight between 26 and 33 kDa, which was recognized by CAD antibody (Millipore, ab16926) (Figure 3g). More importantly, because the concentrations at which the two enzymes cleaved DNA were of the same order of magnitude, demonstrating that AChES-DNase activity was comparable to that of CAD (Figure 3h). Similar to AChE-T547 and CAD, the other apoptotic DNases, CRN-4/RNase T [29], CYP-13/CYPE [30], and CPS-6/EndoG [12], degraded DNA at micromolar concentrations. Although the activity of these enzymes is much lower than that of DNase I, this is normal and sufficient for effective DNase function during apoptosis.

When the peptide was incubated with other plasmids of various sizes, similar cleavage patterns were observed (Supplementary Figure S2A). In addition, hAChE-T547 degraded naked genomic DNA (Supplementary Figures S2B–E), which produced a smear pattern similar to that produced by the digestion of linearized plasmid (Supplementary Figure S2F).

**The purified mouse AChES polypeptide cleaves naked DNA**

To determine whether AChES derived from other species possess DNA cleavage activity, mouse AChES aa 32–579 (mAChE-T548), which was provided by Dr Palmer Taylor (Department of Pharmacology, University of California, San Diego, USA), and purified as described previously [31, 32] (Supplementary Figure S3A, lanes 2 and 2′). Similar to hAChE-T547, mAChE-T548 cleaved both plasmid and naked genomic DNA efficiently (Supplementary Figures S3B–E).
These data indicated the DNase activity of mouse AChE.

The N terminus of AChE, but not its cholinesterase active center, is responsible for its DNA cleavage activity

We investigated whether the cholinesterase active center of AChE also contributes to its DNase activity by using AChE inhibitors (AChEIs) (10 μM huperzine A, 1 μM tacrine and 13 μM donepezil). The acetylcholinesterase activity of mAChE-T548 was markedly inhibited by AChEIs (Supplementary Figure S4A), but its DNA cleavage activity was not (Supplementary Figure S4B). These data suggested that the functional domain responsible for the DNase activity of AChE is distinct from that for its cholinesterase activity. To further confirm this suggestion, AChE residues S234, E365 and H478 of the catalytic triad contributing to its cholinesterase activity were all mutated to alanine (A) to generate pEGFP–NLS–AChE (S234A, E365A, H478A). This construct expresses a mutant AChE fusion protein with a NLS at its N terminus and GFP at its C terminus (NLS–mtAChE–GFP) (Supplementary Figure S4C). As expected, acetylcholine hydrolysis activity was completely abolished in the mtAChE, verifying that the catalytic triad is necessary for its hydrolyzing acetylcholine (Supplementary Figure S4D). If the cholinesterase active center also contributes to its DNA cleavage activity, the nucleus-localized mtAChE would lose the capacity to initiate apoptosis. However, it still stimulated cell apoptosis to a similar degree to that induced by wild-type (wt) AChE (Supplementary Figures S4E–G). These results demonstrate that the catalytic triad of AChE is indispensable for its cholinesterase activity, but is irrelevant to its DNase activity.

To verify that DNA cleavage in the cell-free hydrolysis system was caused by the AChE polypeptide itself but not by other DNase contaminants, we mapped the functional domain responsible for the DNase activity of AChE by screening the fragments with the ability to initiate apoptosis. Therefore, plasmids pEGFP–NLS–tAChE were constructed, which respectively encode the truncated AChE (tAChE) forms, including AChE aa 2–247, aa 2–191, aa 2–138, and aa 2–72 (Figure 4a). The tAChE forms were fused with NLS at the N terminus and GFP at the C terminus. NLS–tAChE247, -191, and -138 showed the same strong apoptosis-inducing capacity as the NLS–(wt-AChE), whereas NLS–tAChE-72 did not (Figures 4b and c). These data confirmed that the pro-apoptotic fragment of AChE is localized within aa 2–138, and aa 72–138 is indispensable.

Next, we investigated whether AChE aa 2–138 possesses DNA cleavage activity. The 31 amino-acid residues at its N terminus form a signal peptide for the translocation of precursor AChE into the lumen of the endoplasmic reticulum [1, 33]. Furthermore, neither hAChE-T547 nor mAChE-T548 contains the signal peptide, but both cleave DNA efficiently. Therefore, we hypothesized that AChE aa 1–31 is irrelevant to the DNA cleavage activity. The polypeptides, human AChE aa 32–138 (hAChE-T107) and aa 32–72 (hAChE-T41), were synthesized and purified by
Ketai BioTech (Shanghai, China) (Supplementary Figures S5A and B). Purified synthetic peptides often contain a small amount of intermediate products composed of partially protected sequences carrying protecting groups and truncated sequences missing one or two N-terminal residues [34]. Analysis of peak fractions by high-performance liquid chromatography is a common method used to identify the purity of chemically synthesized peptides [34]. High-performance liquid chromatography analysis resulted in one major peak revealing 93.9% purity of the product hAChE-T107 and 96.1% purity of the hAChE-T41 (Supplementary Figure S5A and B). The peptides were dissolved in the hydrolysis buffer with or without Mg²⁺ and (or) Ca²⁺ and folded slowly at 4 °C. hAChE-T107 digested both plasmid and naked genomic DNA (Figures 4d–f; Supplementary Figures S5C–D). Furthermore, the peptide degraded the linearized plasmid DNA into smears similar to those produced by the digestion of naked genomic DNA (Supplementary Figure S5E). DNA cleavage by the peptide at pH 7.5 was slightly more efficient than that at any other pH (Figure 4g). Moreover, Mg²⁺ and Ca²⁺ showed a synergistic effect on the enzyme activity (Figure 4h). These data are consistent with those obtained by incubation of DNA with the purified hAChE-T547 (Figure 3). However, hAChE-T41 (10 μM) showed no DNA cleavage activity (Figures 4d and e).

These data suggest that aa 32–138 is the functional domain responsible for AChE₅-DNase activity, which is consistent with the observation that AChE₅ aa 2–138 possesses the pro-apoptotic capacity.

Both the purified and the synthesized AChE₅ peptides degrade chromosomal DNA in pre-fixed cells

In addition to naked DNA, chromosomal DNA was digested by AChE₅, as evidenced by positive TUNEL staining. The average intensity of TUNEL staining in MEFs was quantified by Leica Confocal Software. Fifty cells were randomly selected and analyzed in each experimental group in c. Values represent the mean ± s.d. **P < 0.01, ***P < 0.001, #P > 0.05 vs AChE−/− MEFs without virus infection. Two-tailed Student’s t-test. The data shown are representative of two independent experiments. DAPI, 4',6-diamidino-2-phenylindole; GFP, green fluorescence protein.
staining in the pre-fixed and permeabilized HeLa cells after incubation with hAChE-T547 (0.2 μM) and hAChE-T107 (5 μM) at 37 °C for 6 h. In contrast, bovine serum albumin (0.2 μM) or hAChE-T41 (5 μM) failed to do so (Figure 5a). Digestion of DNA by endogenous DNases in nuclei is dependent on the accessibility of DNA; however, endogenous DNases cannot access chromosomal DNA in normal living cells. For this reason, the endogenous DNases, including EndoG and CAD/DFF40, can no longer ‘actively’ perform biochemical reactions in pre-fixed cells, and thus, the resulting digestion must be the result of the direct action of the AChES polypeptide that we have supplied. In this case, AChES initiated the chromosomal DNA cleavage independent of the function of endogenous DNase, including CAD/DFF40 and EndoG.

TUNEL assays are commonly used to detect DNA fragmentation and strand breakage by the generation of free 3′-hydroxyl-terminal ends [10, 35]. Therefore, positive TUNEL staining caused by hAChE-T547 and hAChE-T107 in pre-fixed cells demonstrates that AChES degrades chromosomal DNA, generating 3′-hydroxyl DNA breaks. The DNA fragmentation capacity of AChES was further confirmed by DNA ladder formation (Figure 5b). The purified hAChE-T547, rather than the synthesized hAChE-T107, was used in this assay because the DNA cleavage activity of...
hAChE-T107 was much weaker than that of hAChE-T547. This difference might be attributed to the non-optimal folding of the synthesized peptide.

**Nuclear-localized AChEs degrades DNA independently of CAD/DFF40 and EndoG**

To detect the functional relationships between NLS–AChEs and the other apoptotic DNases, and more importantly, with caspase-activated CAD/DFF40 and EndoG, we pre-inhibited caspase activation using the pan-caspase inhibitor Z-VAD-FMK (150 μM) 1 h prior to overexpressing NLS–AChEs in HeLa cells. As expected, caspase activation was blocked (Supplementary Figure S6A). In the absence of activated caspase-3, CAD/DFF40 is bound by its chaperone ICAD/DFF45 and fails to have a role in apoptotic DNA cleavage [13, 26]. Besides, EndoG, another caspase-dependent apoptotic DNase, was retained outside the nucleus by the inhibitor and was unable to gain access to the nuclear DNA (Supplementary Figure S6B). However, DNA cleavage was still stimulated by NLS–AChEs (Supplementary Figures S6B and C). These data imply that nuclear-translocated AChEs functions as an apoptotic DNase in a CAD- and EndoG-independent manner. This is consistent with the observation that AChEs polypeptides cleave chromosomal DNA in pre-fixed cells where CAD/DFF40 [36] and EndoG are sequestered in the cytoplasm (Figure 5a).

**AChEs degrades apoptotic DNA via its functional domain aa 32–138**

The capacity of AChEs to cleave DNA during apoptosis was further confirmed by the function of endogenous AChEs. During apoptosis induced by mitomycin C (MMC) (40 μM) in mouse embryonic fibroblasts (MEFs) (Figure 6a), DNA cleavage in MEFs with or without the wt AChE gene (AChE+/+) MEFs, AChE Δ MEFs) (Figure 6b) was detected by TUNEL staining. In comparison with AChE+/+ MEFs, AChE gene knockout significantly impaired DNA cleavage (Figure 6c, upper panel). Furthermore, overexpression of the wt-AChEs rescued the impaired DNA cleavage, whereas (aa 32–138)-deleted AChEs (AChES Δ (aa 32–138)) did not (Figure 6c, lower panel, and Figure 6d; Supplementary Figure S7). Together with the data showing DNA cleavage activity of AChEs aa 32–138 in vitro, these results confirm that AChEs performs an apoptotic DNase function via its aa 32–138 domain, although the mechanism underlying this activity requires further investigation.

**AChES prompts apoptosis via its DNA cleavage domain**

Consistent with the impaired DNA cleavage in apoptotic AChE Δ MEFs, AChE gene knockout also significantly attenuated drug-induced apoptosis (Figure 7a), thus confirming the pro-apoptotic role of endogenous AChEs. To determine whether AChEs prompts apoptosis via its DNA cleavage domain, we attempted to establish stable cell lines. In accordance with the apoptosis-inducing effects of nuclear-localized AChEs (Figures 1c–f), we were unable to establish the NLS–AChEs–GFP-overexpressing stable cell line because the cells could not survive, and certainly, the transfected cells lose tumorigenicity in BALB/c nude mice. In contrast, the stable cell line expressing NLS–AChEs Δ (aa 32–138)–GFP was established successfully (Supplementary Figure S8) and showed a similar rate of tumor development compared with those stably expressing GFP (Figures 7b–d). These data reveal that the DNA cleavage domain aa 32–138 is also the pro-apoptotic domain of AChEs, through which AChEs participates in apoptosis. The obviously lowered rate of tumor formation by AChEs–GFP-overexpressing cells (Figure 7d) can be attributed to the fact that overexpression of AChEs slows down cell growth [37].

**Discussion**

AChEs has emerged as an important contributor to apoptosis in various types of cells [2–6]. However, it is inexplicable that cholinergic neurons with high basal levels of AChEs protein show long-term growth and normal morphology [3]. In the light of the finding that AChEs is a bifunctional enzyme with acetylcholine hydrolysis and DNA cleavage domains, a nuclear translocation-dependent role for AChEs may shed light on this question. Despite abundant AChEs expression in cholinergic neurons, under normal conditions the protein is localized outside of the nucleus and is inaccessible to the chromosomal DNA. Consequently, AChEs is unable to act as a DNase, which makes it understandable that the neurons with high basal AChE levels survive normally. In this case, AChEs might perform its canonical function to terminate neurotransmission by hydrolysis of ACh. However, in response to Aβ stress, AChEs translocates into the nucleus and DNA cleavage occurs (Supplementary Figure S9). This translocation event might act as a critical switch of the canonical function of AChE as a cholinesterase to a noncanonical function as a DNase.

The optimum pH for enzyme activity depends on the environment in which the enzyme normally works.
In the stepwise degradation of DNA in apoptotic mammalian cells, DFF40 and EndoG act at the early stages to initiate DNA breakage [27]. Consistent with this, these enzymes show maximum DNA cleavage activity at pH 7.5, near to the physiological pH 7.4 [23]. In contrast, DNase II, required for metabolizing residual DNA in dying cells, acts at a later stage when acidification occurs [10, 38] and its optimal pH is 5.0–6.0 [38, 39]. We proposed that the optimum pH for an apoptotic DNase is closely correlated with the stage of apoptosis at which it functions. The optimal pH 7.5 for AChES suggests that the enzyme acts in early stage apoptosis. This hypothesis was supported by the observation of (NLS–AChES)-induced CAD/DFF40- and EndoG-independent 3'-hydroxyl DNA cleavage after overexpression in normal living cells without treatment with any other stimuli.

The requirement for divalent cations for DNA cleavage is a general feature of apoptotic DNases. Both DFF40 [23] and EndoG [21] are Mg²⁺-endonucleases, requiring Mg²⁺ and are not costimulated by Ca²⁺. However, DNase I [13] requires both Ca²⁺ and Mg²⁺ for DNA hydrolysis. One Ca²⁺ stabilizes the functional DNase I structure. The presence of Mg²⁺ in close proximity to the catalytic pocket of DNase I reinforces the idea of a cation-assisted hydrolytic mechanism [13].

Our study demonstrates that AChES shows characteristic Ca²⁺–Mg²⁺-dependent DNase activity, similar to that of DNase I. DNA cleavage in the (hAChE-T547)-containing reaction system was not caused by DNase I contamination, because G-actin effectively inhibited the DNA cleavage by DNase I but did not inhibit that by hAChE-T547. Overexpression of the nuclear-localized wt rather than aa 32–138-deleted AChES rescued the attenuated DNA cleavage in apoptotic AChE⁻/⁻ MEFS. Together with the demonstration of the DNase activity of the synthesized AChES aa 32–138, these results further confirm that AChES performs its DNase function via its aa 32–138 domain.

The mechanism by which AChE acts as a common and crucial component in the induction of apoptosis has rarely been investigated. It has been reported that the cytoplasm is the subcellular compartment in which AChE mediates apoptosis, where it participates in apoptosome formation by interaction with caveolin-1, and subsequently with cytochrome c and protease-activating factor-1 (Apaf-1) [40]. Another report documents that N-terminally extended AChES induces apoptosis via the structure of its cholinesterase active center, whereas the cholinesterase activity itself is irrelevant to the induction of apoptosis [41]. N-terminally extended AChES is located in the membrane with the catalytic domain positioned toward the extracellular space, where it might act as a ligand-activated receptor to mediate intracellular signaling in response to extracellular cues [41]. In this study, AChES was found to promote apoptosis by its DNase function. The functional domain was found residing within AChES aa 32–138, losing cholinesterase activity, thus indicating no direct necessity for cholinesterase function in pro-cytotoxic DNase action.

The lack of involvement of cholinesterase function in pro-cytotoxic DNase activity sheds light on the following question. Some AChEIs, such as huperzine A, tacrine, and donepezil, have the ability to partially inhibit cell apoptosis caused by some insults [4–6, 42–44], whereas they do not affect DNA cleavage by AChES. AChEIs protect cells against apoptosis via different mechanisms, depending on the nature of the toxic insult [45]. Studies also indicate that AChEIs impair the apoptosis of neurons by modulating gene expression, including downregulation of pro-apoptotic p53, c-jun, and bax, and upregulation of anti-apoptotic Bcl-2 [46, 47]. Alternatively, it has been suggested that the neuroprotective effect of donepezil is mediated via direct binding to an allosteric site on the nicotinic acetylcholine receptor (nAChR) [45, 48]. This apoptosis-inhibitory effect is independent of the blockage of AChE [45, 48]. Moreover, some AChEIs fail to protect cells from apoptosis induced by certain insults [45]. Thus it is clear that AChEIs inhibit apoptosis in a cholinesterase activity-dependent or -independent manner. Taken together, this explains why some AChEIs exert anticytotoxic effects without inhibition of its DNase activity.

The findings of this study may help to elucidate the mechanisms underlying neuron loss during Alzheimer’s disease (AD) progression. It has been found that AChES is expressed abundantly in normal hippocampus, whereas AChE is rarely expressed [49]. The hippocampus is one of the most vulnerable regions to apoptotic stimuli during development of AD [50]. The results of this study indicate that high-level AChES expression confers apoptotic susceptibility on neurons, which is strongly supported by other studies. AChES transgenic mice exhibited increased neural apoptosis in hippocampi and the mice show impaired acquisition and retention of knowledge, whereas AChE transgenic mice did not [51]. In this study, Aβ (a type of toxin found in the AD brain) was found to be deposited in a brain section prepared from a 1-year-old B6C3-Tg (APPswe, PSEN1dE9)85Dbo/J transgenic mouse (a
mouse model of AD), and AChE was found to be translocated into the nuclei (Supplementary Figures S9A and B). In addition, in response to Aβ-induced neurotoxicity, primary hippocampal neurons showed nuclear translocation of AChES and chromosomal DNA cleavage (Supplementary Figure S9C). Together with the observation that AChES exerts a DNA-hyrolsis function after translocation into the nuclei, these data suggest that nuclear translocation and subsequent cleavage of chromosomal DNA is one of the functions of AChES in neuron loss during AD progression, although this speculation requires further investigation.

In summary, this study demonstrates that AChES performs a vital function as a DNase in apoptosis. The stepwise events, including upregulated expression, nuclear translocation, subsequent binding with and digestion of chromosomal DNA, constitute the mechanism by which AChES mediates cell apoptosis. The region comprising aa 32–138 is the indispensable domain conferring apoptotic DNase activity on AChES. However, the mechanism by which AChES is activated during apoptosis and how AChES and other DNases are coordinated and recruited into apoptotic machinery remain to be determined. Nevertheless, this work elucidates a novel role of AChES, and indicates the potential for the development of novel drugs targeting the DNase activity of AChES for the treatment of neurodegenerative diseases, such as AD.

Materials and Methods

Animals
Heterozygous AChE gene knockout (AChE+/−) mice (stock number: 005987; strain name: 129-Achtm1Loc/J) were purchased from the Jackson Laboratory, Bar Harbor, ME, USA. They were bred, and AChE-deficient embryo mice were identified as described previously [3, 44]. B6C3-Tg (APPswe, PSEN1dE9) 85Dbo/Mjjax transgenic mice (stock number: 004462) were also purchased from the Jackson Laboratory. The following primers were used for identification of the transgenic mice: 5′-CTTCTTTGTGACTATGTGGACAGTGACTGATGTCGG-3′, reverse 5′-GTGGATAACCCCGTCCCCCAGCTAGACC-3′; APPswe: forward primer 5′-GACTGACCACTCGACAGGTTCTG-3′, reverse 5′-CTTGTAAGTTGAGTTCTCATATCCG-3′. Four-week-old female BALB/c mice and male Sprague-Dawley rats (250–300 g) were purchased from Shanghai SLAC Laboratory Animal Co. Ltd (Shanghai, China). All animals were housed under standard conditions of 12 h light/12 h dark cycles with free access to food and water. The experimental protocols were approved by the Institutional Animal Ethics Committee of the Shanghai Institutes for Biological Sciences.

Cell culture, transfection, apoptosis induction, and generation of stable cell lines
HeLa and 293 T cells were obtained from the Shanghai Cell Resource Center, Chinese Academy of Science. HEK-293 S cell lines lacking N-acetylglucoasaminyltransferase 1 activity (GnTI−/− HEK 293S) were provided by Dr Palmer Taylor (Department of Pharmacology, University of California). All cells were cultured in Dulbecco’s modified Eagle’s medium (Invitrogen, Carlsbad, CA, USA) supplemented with 10% fetal calf serum (Tianhang Biological Technology Co. Zhejiang, China). Primary MEFs were isolated from AChE−/− or AChE+/− E13 129 mouse embryos as described previously [52] and cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum. Primary hippocampal neurons were isolated from the Sprague-Dawley rats as described previously [53] and cultured in Neurobasal-A medium supplemented with GIBCO B-27 (Invitrogen). All cells were cultured at 37 °C in a humidified atmosphere of 95% air and 5% CO2. FuGENE HD Transfection Reagent (Roche Diagnostics, Mannheim, Germany) was used for transfection of plasmids into HeLa cells, according to the manufacturer’s instructions. For infection of MEFs with the lentiviral system (System Biosciences, Mountain View, CA, USA), medium from 293T cells co-transfected with pCMV-delta-8.2, pCMV-VSV-G, and pCDH-CMV-MCS-EF1-Puro-GFP/AChES-GFP/AChES A (aa 32–138)–GFP using lipofectamine 2000 (Invitrogen) was collected, centrifuged at 2 500 g for 10 min and filtered (0.45 μm pore size) at 48 h post transfection. The supernatant was applied to MEF cells for 48 h followed by treatment with 4 μM MMC (Sigma-Aldrich, St Louis, MO, USA) for 36 h. Cell apoptosis was detected by the TUNEL assay. For generation of HeLa cell lines stably expressing GFP or AChES–GFP or NLS–AChES A (aa 32–138)–GFP, transiently transfected cells were grown in 1 mg/ml G418 (Sigma-Aldrich) for 7 days after transfection. Pooled populations of G418-resistant cells were obtained and then continuously cultured in 200 μg/ml G418-containing culture medium. After 4 weeks, GFP-positive cells were further sorted using a FACSort flow cytometer (Becton Dickinson, CA, USA), cultured in 200 μg/ml G418-containing culture medium, and used for propagation.

Preparation of polypeptide hAChE-T547
GnTI-293 S cells stably expressing human AChE aa 32–578 (hAChE-T547) were cultured in Dulbecco’s modified Eagle’s medium culture medium with 10% fetal calf serum and 2 μg/ml puromycin. Two days before protein purification, the culture medium was replaced by serum-free medium (UltraCULTURE; Biowhittaker, Walkersville, MD, USA) with 1% l-glutamine. Human AChE-T547 was purified according to modified protocols, as described previously [54, 55].

The dynamics of DNA–protein interactions
The real-time kinetics of the interactions between hAChE-T547 and pEGFP-c1 plasmid DNA were examined using a BIACORE T100 system (GE Healthcare Biacore; Piscataway, NJ, USA). The protonated hAChE-T547 polypeptide, bovine serum albumin (irrelative control protein), and DNase I (positive control protein) (Sigma-Aldrich) were immobilized on the...
activated sensor chips (Series S Sensor Chip CM5) (GE Healthcare Biacore). The pEGFP-c1 plasmids were diluted to 1.88 × 10⁻⁴ M in a running buffer (10 mM HEPES, pH 7.5, 2.5 mM CaCl₂, 5 mM MgCl₂) ((Sigma-Aldrich) and injected over the sensor chip surface at 20 μl/min at 37 °C to generate ~130 response units on the surface of hAChE-T547 peptides. The plasmids were then further diluted in running buffer to the concentrations indicated in Figure 2c, and injected at 37 °C at a flow rate of 20 μl/min for 150 s. Surface regeneration was achieved using a 2-min injection of the running buffer at 100, 30, and 20 μl/min. Plasmid concentrations were analyzed in duplicate, and any background signal generated by the running buffer was subtracted. The data were analyzed using the Biacore T100 evaluation software (GE Healthcare Biacore).

Plasmid cleavage assays
AChE polypeptides were incubated with plasmid DNA in a cell-free hydrolysis system (5 mM Tris-HCl pH 7.5, 2.5 mM CaCl₂, 5 mM MgCl₂) for 6–12 h at 37 °C. The hydrolysis products were subjected to 1% agarose (Invitrogen) gel electrophoresis. The gel was stained with ethidium bromide and visualized using a Tanon 2500 gel imaging system (Bio-tanon, Shanghai, China).

TdT-mediated dUTP nick end labeling (TUNEL) assays
The TMR red in situ cell death detection kit was purchased from Roche, Basel, Switzerland. TUNEL assays were performed according to the manufacturer’s instructions. The transfected HeLa cells grown on coverslips in 24-well plates were fixed with 4% paraformaldehyde for 15 min and permeabilized with 0.5% Triton X-100 for 10 min at room temperature (RT), followed by incubation with 18 μl labeling solution plus 2 μl enzyme solution at 37 °C for 1 h. Cell nuclei were stained with 0.1 μg/ml 4',6-diamidino-2-phenylindole (Sigma-Aldrich) at RT for 5 min. The labeled cells were then washed, transferred onto glass slides, and observed by laser scanning confocal microscopy.

Assessment of acetylcholinesterase activity
Acetylcholinesterase activity was examined using a modified Ellman method as described previously [44]. Collected cells were resuspended in potassium phosphate buffer (pH 7.4) containing 0.5% Tween 20 and 1 mM NaCl, sonicated at 4 °C using an ultrasonoton generator, and centrifuged at 10 000 g at 4 °C for 10 min to get rid of cells/debris. The supernatant was incubated with 25 μM iso-OMPA in sodium phosphate (pH 8.0) containing 0.315 mM 5,5'-dithio-bis(2-nitrobenzoic acid) (DTNB) at 37 °C for 30 min. After addition of acetylthiocholine iodide (final concentration, 5 mM), optical density values at 405 nm were measured spectrophotometrically every 5 min in a 96-well microtiter plate at 37 °C.

Preparation of a double-stranded DNA oligonucleotide of NLS
The single-stranded NLS oligonucleotides were synthesized by Sangon Biotech, Shanghai, China. Forward: 5'-gatct ATGCGCAAGAGAAAGCGTAAGGTCCCAAAAGAAGAA GCGTAAAGGTa-3; reverse: 5'-agctAACCCTACGTTCTG TCTTTGGAACCTTACGTTCTCTTTTTCGATa-3'.

The single-stranded NLS oligonucleotides were dissolved in sterile distilled water to a concentration of 50 μM. The complementary single strands were added to the annealing buffer (10 mM Tris, pH 7.5–8.0, 50 mM NaCl, 1 mM EDTA) in a 1.5-ml tube to a final concentration of 22.5 μM and incubated in boiling water for 5 min. The oligonucleotides were allowed to cool slowly to RT.

Plasmid constructs
pEGFP–AChES was generated as previously described [3]. For construction of pEGFP–NLS–AChES, the double-stranded NLS DNA with sticky ends (Bg/II–NLS–HindIII) was inserted between the Bg/II and HindIII sites in pEGFP–AChES. For construction of pEGFP–NLS–AChES (S32A, E365A, H478A), site-directed mutagenesis was performed with the primary template plasmid pEGFP–NLS–AChES. The following three pairs of primers were used in sequence: forward, 5'-CTGTTTTGGGAGGCCGCCGGAGCCGGC-3' and reverse, 5'-GGGGCTCCCCGCCCCTCCACCCCAACAGC-3' for introducing a S234A change; forward, 5'-GTGTGGTGAAG GATGCGGGCTGCTTATTTTC-3' and reverse, 5'-AGAAA ATACAGGACCCCATCCTCCACACAC-3' for generation of an E365A change. Finally, forward, 5'-GATGGGTTGTC CGCGCGGCCTAGAGATC-3' and reverse, 5'-GATCTCG TAGCCCGGGGCTACCGACCCATC-3' for introducing a H478A change. For construction of pEGFP–NLS–tAChES encoding tAChES, the tAChES complementary DNAs (cDNAs) encoding the aa 2–1250 gel imaging system (Bio-tanon, Shanghai, China).
Time-lapse fluorescence microscopy imaging

HeLa cells grown in q3.5 cm-dishes were co-transfected with AChE γ-GFP and histone H2b-RFP or with tubulin-GFP and histone H2b-RFP. After 24 h, cells were exposed to 100 μM H2O2. The changes in cell morphology and the distribution of AChE γ-GFP were then monitored under a Leica AS MDW live cell image acquisition system (Leica Microsystems, Wetzlar, Germany). Representative cells were photographed at 2 min intervals for 290 min.

Cell sorting by flow cytometry

At 18 h after transfection, cells were harvested, centrifuged at 800 g for 10 min, resuspended in cell culture medium (1 × 10^7 cells/ml), and filtered through a 40-μm nylon mesh (BD Falcon, Bedford, ME, USA). GFP-positive cells were then sorted with a Becton Dickinson FACSort flow cytometer (excitation at 488 nm).

MTT assay of cell viability

Cell viability is commonly measured using MTT assays [44]. MTT (Sigma-Aldrich, Shanghai, China) was dissolved in phosphate-buffered saline (1 ×, pH 7.2–7.4) to give a final concentration of 5 mg/ml. MTT solution (20 μl) was added to each well of a 96-well plate containing 100 μl culture medium and then incubated at 37 °C for 4 h. The formazan crystals were dissolved in 100 μl dimethyl sulfoxide. Finally, optical density values at 570 nm were measured by using a Multiscan MC3 microplate reader (Thermo Labsystems, Vantaa, Finland).

Western blot analysis

Western blot analysis was performed as described previously [3]. The following primary antibodies were used: mouse anti-GFP-tag (7G9) mAb (Abmart, Shanghai, China, 1:10,000), rabbit polyclonal anti-AChE antibody (1:1000) (Dr Palmer Taylor’s laboratory). The secondary antibodies were goat anti-mouse-HRP (Santa Cruz, sc-2030, 1:5000) and goat anti-rabbit-HRP (Santa Cruz, sc-2030, 1:5000), respectively.

AChE cytochemical staining

Frozen sections of wt B6C3 and B6C3/Tg(APPswe, PSEN1ΔE9)85Db/1 mouse brains were prepared using standard procedures. AChE cytochemical staining was performed as described previously [3]. The specimens were incubated in 15 ml of 0.1 M sodium phosphate (pH 6.0) containing 10 mg of acetylthiocholine iodide, 1 ml of 0.1 M sodium citrate solution, 2 ml of 30 mM copper sulfate solution, and 2 ml of 5 mM potassium ferricyanide solution at RT for 4–8 h. Thereafter, the slides were incubated in Harris’ hematoxylin solution for another 30 s. The samples were then dehydrated with ethanol and sealed in neutral balsam. AChE staining was observed under a phase-contrast microscope.

DNA ladder assay

HeLa cells (5 × 10^6) were harvested by trypsin digestion at 48 h after transfection, centrifuged at 15,000 g for 20 min at RT, and the supernatant was discarded. In the experiment examining chromosomal DNA fragmented by AChE polypeptide, HeLa cells (2.5 × 10^6) were harvested, fixed with 4% paraformaldehyde for 15 min, permeabilized twice with 0.5% Triton X-100 (15 min per incubation) at RT, and then incubated with 21 μM hAChE-T547 overnight at 37 °C. Subsequently, DNA was extracted using the phenol/chloroform method. The samples were subjected to 1.5% agarose gel electrophoresis, stained with ethidium bromide, and visualized using the Tanon 2500 gel imaging system.

Immunofluorescence assays

The assay was performed as described previously [3]. Briefly, cells grown on the coverslips in a 24-well plate were washed with 0.01 M phosphate-buffered saline (153.8 mM NaCl, 11.2 mM Na2HPO4, 12H2O, 2.6 mM NaH2PO4·2H2O, pH 7.2–7.4) and fixed with 4% paraformaldehyde for 15 min at RT. Following permeabilization in 0.5% Triton X-100 for 10 min, cells were washed three times with 0.01 M phosphate-buffered saline and then incubated with blocking buffer (5% normal goat serum) for 20 min at RT. Cells were incubated with primary antibody at 4 °C overnight, washed three times and incubated with the secondary antibody for 30 min at 37 °C in the dark, followed by 0.1 μg/ml 4',6-diamidino-2-phenylindole (Sigma-Aldrich) staining at RT for 5 min. The labeled cells were then washed, transferred onto glass slides, and observed under a laser scanning confocal microscope (Leica). The following primary antibodies were used: cleaved caspase-3 (Asp175) rabbit polyclonal antibody (Cell Signaling, Shanghai, China, #9661, 1:100) and rabbit polyclonal anti-EndoG antibody (Abcam, Shanghai, China, ab9647, 1:100). The corresponding secondary antibodies were Cy3-AfFlPure goat anti-rabbit IgG (Jackson Immunoresearch Laboratories, West Grove, PA, USA, 111–165–045, 1:500) and Alexa Fluor 647 goat anti-rabbit IgG (Molecular Probes, Eugene, Oregon, USA, A-21244, 1:500), respectively. The images were taken under a laser scanning confocal microscope (Leica, Wetzlar, Hessen, Germany).

Congo red staining

Congo red (Sigma-Aldrich) staining for Aβ amyloid was performed as described previously [56].

Statistical analysis

Data were expressed as mean ± s.d. The significance of differences between two groups was analyzed using two-tailed Student’s t-tests. P-values < 0.05 were considered to indicate statistical significance.

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