Cleavage of the Nuclear Matrix Protein NuMA during Apoptosis

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INTRODUCTION

Apoptosis or programmed cell death is an active intrinsic process through which cells kill themselves [1]. It is a fundamental feature of animal cells essential for regulating cell populations [2]. The main distinctive morphological features of apoptosis are cell rounding and shrinking, condensation and fragmentation of the nucleus, cytoplasmic blebbing, and disassembly of the dead cell through budding of apoptotic bodies [3, 4].

The downstream effectors of apoptosis in nuclei may be ubiquitous since apoptotic cell-free extracts induce apoptosis in nuclei from a wide range of cultured cells. Nonnucleolar nuclear matrix-associated proteins, including a 126-kDa polypeptide [5] and the 13-kDa component of the U1snRNP particle [6], relocate within the nucleus during apoptosis. Nuclear matrix proteins topoisomerase I and II and lamins [7, 8] are cleaved during apoptosis. NuMA, a ubiquitous nuclear matrix protein which relocates to the spindle poles during mitosis [9, 10], is also proteolytically processed during apoptosis [11, 12] indicating that it could be a structural target for a death protease. Here we analyze the position of the cleavage site and assess whether this cleavage is affected by protease inhibitors.

METHODS

Apoptosis. Apoptosis was induced in HeLaSS6 and BHK cells grown in DME medium plus 10% fetal calf serum (i) by hydroxyurea (HU, final concentration 2.5 mM, Sigma) in medium for 5 h, Stau- sorporine (St, 200 ng/ml, Sigma) was then added for a further 9 to 12 h, or (ii) by tumor necrosis factor α (TNFα, 3200 units/ml, Signal Transduction Laboratories) and cycloheximide (10 μg/ml, Sigma) for 12 h.

Mutant NuMA cDNA constructs. Full-length (FL) and the new d1 and d2 NuMA cDNAs (Fig. 1) were derived from pCMV5 NuMA expression vectors [13]. Amino acid residues 7 to 1502 were deleted by digesting the pCMV5 vector containing NuMA cDNA with Smal and religating the appropriate digest products. The FL NuMA site-directed mutant was obtained using the ExSite PCR-based site-directed mutagenesis kit (Stratagene). Ligated and mutated regions were checked by sequencing.

Transfection. BHK cells were transiently transfected using the modified calcium phosphate precipitation technique [14]. Cells were allowed to recover from transfection in fresh medium for 6 h before apoptotic induction.

Immunofluorescence microscopy. Immunofluorescence was by standard procedures using monoclonal NuMA SPN-3 and SPN-7 antibodies [9] as undiluted hybridoma supernatants. To detect cell death in situ, apoptotic cells were labeled with dUTP–FITC for 1 h with the TUNEL–FITC kit (Boehringer Mannheim) prior to NuMA staining. DNA was stained with Hoechst 33342.

Gel electrophoresis and immunoblotting. Cells from a single confluent 60-mm-diameter petri dish were scraped into 100 μl double-strength sample buffer and boiled at 100°C. SDS–polyacrylamide gel electrophoresis and immunoblotting on nitrocellulose were performed using standard procedures. Immunostaining of blots was performed using undiluted hybridoma supernatant for the SPN-3 and SPN-7 antibodies and peroxidase-labeled anti-mouse antibodies (Dako, Hamburg) diluted 1:5000 in the ECL technique (Amersham).

Use of protease inhibitors. HeLa cells were treated with 2.5 mM HU for 9 h. Then staurosporine and protease inhibitors were added. Final concentrations were 0.1 mM for TLCK and TPCK (Sigma), 20 μM for ICE inhibitor II (Ac-YVAD-cmk, Bachem), and 10 μM for E-64 (Sigma).

RESULTS

Induction of Apoptosis

We induced apoptosis in HeLa cells using a 5-h incubation in hydroxyurea containing medium followed by a 9-h treatment with the protein-kinase inhibitor staurosporine. The HU treatment blocked the cells in interphase (Fig. 2A), while the subsequent staurosporine treatment induced a drastic change in morphology, giving rise to small round cells which protruded from the substratum but did not detach (Fig. 2B [15]). Hoechst staining of such cells showed patches of highly condensed chromatin located predominantly at the nuclear margin (Figs. 2E and 2G). These changes in cell

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shape and chromatin state and distribution correspond to changes normally seen during apoptosis [3, 4]. The apoptotic state was further demonstrated using an in situ TUNEL (terminal deoxynucleotidyltransferase-mediated dUTP nick end labeling) assay. TUNEL detected all cells showing condensation of the chromatin but did not stain cells showing a normal Hoechst staining (Figs. 2G and 2H). The long stability of the treated cells (more than 24 h), the absence of cell blebbing, the persistence of the nuclear lamina, and the absence of a typical DNA ladder reflecting internucleosomal fragmentation (although such a ladder is detected in HL60 cells under the same conditions) suggested an incomplete apoptotic state, which could not be reversed by growth in normal medium without drugs.

NuMA Location Changes during Apoptosis

In untreated cells, NuMA is found throughout the nucleoplasm, with the exception of the nucleolus, in a distribution which approximates that of chromatin (Figs. 2C and 2D). In apoptotic cells, NuMA was clearly excluded from both the condensed chromatin masses and from the nucleoli (Figs. 2E and 2F).

Proteolysis of NuMA during Apoptosis

To see whether the change in location of NuMA during apoptosis could be linked to specific proteolytic cleavage, we immunoblotted the whole cell extracts (Fig. 3A). In control HeLa cells, SPN-3 antibody revealed a 240-kDa band corresponding to the FL NuMA protein (lane 1). In cells treated with HU + St, SPN-3 antibody detected both the FL protein and a smaller 190-kDa fragment (lane 2). The 190-kDa form was not

FIG. 1. Schematic representation of expressed recombinant wild-type (FL) or mutant (d1, d2) NuMA polypeptides. 216, head; 255 - 267, coiled-coil rod domain; 1701, tail; 1600 - 1700, nuclear localization signal (NLS). Expected molecular weights are given in parentheses. The SPN-3 and SPN-7 epitopes are given under the FL sequence. The amino acid numbering system is from Yang and Snyder [18].

FIG. 2. (A, B) Phase-contrast microscopy of live HeLa cells. Note the absence of round mitotic cells after 5-h treatment with 2.5 mM hydroxyurea (A). After treatment with staurosporine for an additional 9 h many cells are rounded (B). (C - F) Hoechst stain (C, E) and immunofluorescence with the NuMA SPN-3 antibody (D, F) of control HeLa cells (C, D) and after HU + St treatment (E, F). Note the overlap between the Hoechst and the NuMA stain in control cells and the complementarity in treated cells. (G, H) Hoechst (G) and TUNEL (H) labeling of the same HU + St-treated cells. Note the colocalization of the Hoechst and TUNEL stain in the top two cells, and the lack of TUNEL signal in the lower cell which has normal Hoechst staining. Magnification: A and B, ×135; C - F, ×1500; G and H, ×1000.
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1502 introduced to increase the polypeptide separation in the immunoblotting experiment. Since the SPN-3 epitope is deleted in the d1 and d2 constructs, the blot was stained with SPN-7 antibody which recognizes an epitope between NuMA residues 1600 and 1700 (Fig. 1 [17]). Figure 3B shows clearly that d1NuMA was cleaved upon apoptosis, while d2NuMA retained its initial molecular weight. Thus the apoptotic cleavage site is located in the NuMA tail domain between residues 1701 and 1725, i.e., in the sequence 1701QVATDALKS-REPQAKPQLDLS1725 [18]. Because the DA in this sequence is a potential ICE protease site [19], a point mutation in FLNuMA was constructed. However, changing D to G at position 1705 did not prevent cleavage under the conditions used in Fig. 3A.

Inhibition of Apoptosis and NuMA Cleavage through Proteolytic Inhibitors

Proteolytic inhibitors such as TPCK, TLCK, E-64, and ICE inhibitor II were tested for effects on apoptosis induction and on NuMA cleavage in HeLa cells. Only TPCK affected the kinetics of morphological changes. Cell retraction preceding cell rounding, usually detected 30 min after staurosporine addition, was delayed until 5 h in the presence of TPCK. TPCK was also the only protease inhibitor which prevented proteolytic cleavage of NuMA after 8 h of staurosporine treatment.

Discussion

Our data suggest that proteolytic cleavage of NuMA, a nuclear matrix component of interphase cells, may be important for the apoptotic process. In normal interphase cells, NuMA and DNA, as judged by Hoechst 33242 staining, are found in an overlapping distribution, whereas in apoptotic cells they have a strikingly complementary localization (Figs. 2B–2E). Our observation of an 190-kDa apoptotic NuMA fragment (Fig. 3) is consistent with data of Weaver et al. [11] who observed a 200-kDa NuMA fragment after induction of apoptosis in thymocytes by dexamethasone, and of Hsu and Yeh [12] who detected a 180-kDa NuMA fragment which reacted with the SPN-3 antibody (data not shown). Thus the 190-kDa form results from cleavage of FL recombinant NuMA.

The epitope of the SPN-3 antibody is located in NuMA residues 255–267 (Fig. 1 [17]). Since the epitope is retained in the 190-kDa fragment, (cf. Fig. 3A, lane 5), this fragment could be generated either by a single cleavage in the C-terminal tail domain or by a cleavage in the tail domain combined with a second cleavage in the N-terminal 255 residues. Therefore, the HU + St treatment was repeated on BHK cells transfected with d1 and d2 NuMA human cDNA mutants shown in Fig. 1. d1 is truncated at residue 1725 and d2 is truncated at residue 1700 in the tail domain. Both d1 and d2NuMA have an internal deletion of residues 7 to 1502 introduced to increase the polypeptide separation in the immunoblotting experiment. Since the SPN-3 epitope is deleted in the d1 and d2 constructs, the blot was stained with SPN-7 antibody which recognizes an epitope between NuMA residues 1600 and 1700 (Fig. 1 [17]). Figure 3B shows clearly that d1NuMA was cleaved upon apoptosis, while d2NuMA retained its initial molecular weight. Thus the apoptotic cleavage site is located in the NuMA tail domain between residues 1701 and 1725, i.e., in the sequence 1701QVATDALKS-REPQAKPQLDLS1725 [18]. Because the DA in this sequence is a potential ICE protease site [19], a point mutation in FLNuMA was constructed. However, changing D to G at position 1705 did not prevent cleavage under the conditions used in Fig. 3A.

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that the stable fragment may retain the full amino terminal end. The specific proteolytic cleavage of NuMA results in the release of the C-terminal tail domain for which multiple functions have been documented [10, 13]. This region also contains the nuclear location signal [13].

A variety of proteolytic enzymes with different substrate specificities are thought to be involved in apoptosis [4, 20]. Thus far our results provide no evidence that NuMA is a substrate for the interleukin-1-β-converting enzyme (ICE) which may be involved in initiating the active phase of apoptosis [4]. Thus changing amino acid residues DA into GA at positions 1705–1706 did not inhibit NuMA cleavage nor did the specific ICE inhibitor II prevent cleavage. Of the various protease inhibitors tested only the chymotrypsin inhibitor TPCK was effective. Interestingly TPCK delayed both the morphological changes and NuMA cleavage.

Induction of apoptosis results in the cleavage of NuMA and in the dissociation of the 190-kDa fragment from chromatin probably because of the loss of the tail sequence. Two models of nuclear structure in which NuMA plays a key role can be envisaged. First, NuMA binds to defined DNA sequences called matrix attachment regions (MARs) in vitro [21]. MARs are viewed as the likely starting points of DNA fragmentation [22], and thus proteolytic cleavage of NuMA might expose DNA to fragmentation and induce chromatin reorganization. Alternatively, if NuMA is a key component of a putative nuclear matrix, cleavage of NuMA may be sufficient to cause the matrix to collapse and the DNA to rearrange.

REFERENCES