A Novel Response of Cancer Cells to Radiation Involves Autophagy and Formation of Acidic Vesicles

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Abstract

The mechanisms underlying neoplastic epithelial cell killing by ionizing radiation are largely unknown. We discovered a novel response to radiation manifested by autophagy and the development of acidic vesicular organelles (AVO). Acidification of AVO was mediated by the vacuolar H^+-ATPase. Staining with the lysosomotropic agent acridine orange enabled us to quantify AVO accumulation and to demonstrate their time- and dose-dependent appearance. The appearance of AVO occurred in the presence of the pan-caspase inhibitor z-Val-Ala-Asp(OMe)-fluoromethyl ketone, but was inhibited by 3-methyladenine, an inhibitor of autophagy. The accretion of AVO in surviving progenies of irradiated cells, and the increased incidence of clonogenic death after inhibition of vacuolar H^+-ATPase suggest that formation of acidic organelles represents a novel defense mechanism against radiation damage.

Introduction

The cellular and molecular processes involved in the response of neoplastic epithelial cells to radiation are largely unknown. Whereas in some cell types, particularly cells of reticuloendothelial origin, death after irradiation is preceded by apoptotic changes, apoptosis plays little or no role in the killing of epithelial neoplastic cells by radiation (1–3). Epithelial cells do not undergo apoptosis after irradiation and are likely to respond with a different sequence of programmed cytoplasmic and nuclear events. Several investigators have proposed two types of programmed cell death (4, 5). Type I programmed cell death, or apoptosis, is mediated by a cascade of cysteine aspartases (caspases) and factors released by the mitochondria (6), and has typical morphological and biochemical characteristics such as chromatin marginalization and condensation, early nuclear collapse, and nucleosomal ladder formation (5). In contrast, type II programmed cell death is marked morphologically by increased autophagy and early destruction of the cytoplasm that occurs either without nuclear collapse or precedes it (5). Type II programmed cell death has been documented mainly in the Lepidoptera during metamorphosis and during involution of the rat mammary gland (4, 5), but it has rarely been associated with stress-inducing stimuli (7, 8). Unfortunately, methods for quantification of type II programmed cell death are lacking, and the molecular mechanisms that regulate it have not been defined. The work presented here characterizes and quantifies a novel form of response to radiation in cancer cells that is reminiscent of type II programmed cell death. This response is dominated by the appearance and accumulation of AVO. Interference with the acidification of these AVO results in increased radiosensitivity and thus identifies a new target for modulating the radiation response of cancer cells.

Materials and Methods

Cell Culture. MCF-7 (human breast adenocarcinoma), LoVo (human colon adenocarcinoma), and LNCaP (human prostate carcinoma) were obtained from the American Type Culture Collection. Cells were maintained as described previously (9). Irradiation was carried out 48 h post-plating (time 0) at 25°C using a Cs-137 irradiator (Shepherd Mark-I, model 68, SN643) at a dose-rate of 243 cGy/min. Bovine aortic endothelial cells were obtained from Dr. Haimovitz-Friedman, Memorial Sloan Kettering Cancer Center, New York) and were grown and treated with H2O2 as described previously (10). PMA (Alexis Biochemicals Corporation, San Diego, CA) was dissolved in DMSO and added to cells for the duration of 1 h. Final DMSO concentration was 0.03%. The cells were then rinsed with warm growth medium before being returned to the incubator for an additional 48 h.

Supravit Cell-staining with Acidine Orange. Cell staining was performed according to published procedures (12–14). Acridine orange (Polysciences, Warrington, PA) was added at a final concentration of 1 μg/ml for a period of 15 min. Bafilomycin A1 (Sigma Chemical Co., St. Louis, MO) was dissolved in DMSO and added to the cells 30 min before addition of acridine orange. LysoSensor Blue DND-167 (Molecular Probes, Eugene, OR) was added for 8 min at a final concentration of 10 μM. Pictures were obtained with a fluorescence microscope (Olympus BH-2 RFCA) equipped with a mercury 100-W lamp, 490-nm band-pass blue excitation filters, a 500-nm dichroic mirror, and a 515-nm long-pass barrier filter. Images of control and irradiated cells were recorded on Kodak Elite II 100 ASA film for color slides by 4-s exposure.

Determination of Mean Red:Green Fluorescence Ratio in Acridine Orange-stained Cells Using Flow Cytometry. In acridine orange-stained cells, the cytoplasm and nucleous fluoresce bright green and dim red, whereas acidic compartments fluoresce bright red (13, 14). The intensity of the red fluorescence is proportional to the degree of acidity and/or the volume of the cellular acidic compartment (14). Therefore, by comparing the mean red:green fluorescence ratio within different cell populations, we could measure a change in the degree of acidity and/or the fractional volume of their cellular acidic compartment. Cells were stained with acridine orange for 17 min, removed from the plate with trypsin-EDTA, and collected in phenol red-free growth medium. Green (510–530 nm) and red (>650 nm) fluorescence emission from 10^6 cells illuminated with blue (488 nm) excitation light was measured with a FACSCalibur from Becton Dickinson (San Jose, CA) using CellQuest software. The red:green fluorescence ratio for individual cells was calculated using FlowJo software (TREE STAR, Inc., San Carlos, CA). To control for the possible effect of trypsinization on the measured red:green fluorescence ratio, we compared the ratios obtained by flow cytometry with those obtained with a Laser Scanning Microscope (LSM510; Zeiss). Stained cells, grown on coverglass, were illuminated with a 488-nm argon laser beam. The red (>650 nm):green (505–545 nm) fluorescence ratio of an entire image was obtained.

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using software LSM 510 version 2.01 SP2. These measurements yielded similar results to those obtained with flow cytometry. All determinations of red:green fluorescence ratio reported here were therefore obtained via flow cytometry.

**Electron Microscopy.** Cell processing for electron microscopy and staining with DAMP (Molecular Probes, Eugene, OR) was done according to published procedures (15, 16). The fraction of the cytoplasmic volume occupied by AVO (the fractional volume of AVO) was quantified from electron micrographs according to Dunn (16) and Lenk et al. (17). Digital images of the micrographs were obtained with an Epson ES-1200S flat bed scanner with Adobe Photoshop version 5. The fractional volume was calculated with Image Pro Plus version 3 and expressed as a percentage of total cytoplasmic volume.

**Detection of Nucleosomal Fragmentation of Genomic DNA.** DNA extraction and electrophoresis on agarose gel was carried out according to Bose et al. (18). DNA preparation and resolution with pulse field gel electrophoresis was conducted as described by Gilles et al. (19) using the CHEF Mapper (Bio-Rad, Richmond, CA). DNA strand breaks were assayed by the TUNEL method and analyzed by flow cytometry (10).

**Gel Electrophoresis and Western Blotting.** Cells were scraped and collected in PBS containing protease inhibitors (Complete and pepstatin A; Boehringer Mannheim) and lysed in 2% SDS by heating at 95°C. Protein content was determined with bicinchoninic acid reagent (Pierce). PAGE and immunoblotting were performed (20) using anti-LAMP-1 antibodies (Hybri-doma Bank, Department of Biological Sciences, Iowa City, IA).

**Immunocytochemistry.** Cells were fixed with 3% paraformaldehyde, permeabilized with 1% Triton X-100, and stained with anti-LAMP-1 and Texas Red conjugated antimouse IgG (Jackson ImmunoResearch Laboratories, Inc., Jackson, IL).

**Surviving Fraction.** Cells were plated in growth medium at a density of 30 cells/cm² and irradiated 22 h later with 2 and 3 Gy. Cells were irradiated at room temperature in a Cs-137 Irradiator (Shepherd Mark-I, Model 68, SN 643) at a rate of 2.5 Gy/min. Six days later, 90–95% of the grown colonies possessed >50 cells. For determination of their red:green ratio, colonies were processed as described above. For determination of surviving fractions, cells were stained with crystal violet and colonies containing ≥50 cells were counted with a dissecting microscope. The surviving fraction was defined as the ratio between the number of surviving colonies in irradiated culture and in unirradiated culture, and it was calculated at each dose level (21).

**Results and Discussion**

Our experiments indicated that human breast cancer cells are sensitive to standard doses of radiation. After irradiation with 6 Gy, only 0.1% of the cells remained clonogenic (22). Nonetheless, the cells did not show any of the biochemical and morphological changes that are associated with apoptosis up to 4 days after irradiation with 10 Gy. At 4 days after irradiation, 30% of the cells were already dead and did not exclude trypan blue. Still, nucleosomal ladder formation and a positive TUNEL reaction could not be demonstrated (Fig. 1). Furthermore, electron microscopy did not reveal the morphological changes that are typical of apoptosis, i.e., chromatin margination and condensation (Fig. 2). Instead, DNA damage was manifested by micronuclei formation (9) and nondiscrete DNA degradation (Fig. 1). Our findings and the absence of apoptotic markers after irradiation of malignant epithelial cells reported by others (1, 2) led us to search for type II programmed cell death-related cellular events, such as changes in the cellular acidic compartments.

For detecting of the acidic compartment, we used the lysosomotropic agent acridine orange, a weak base that moves freely across the vesicles fluoresced bright red, whereas the cytoplasm and the nucleus showed dominant green fluorescence (Fig. 3Ab). In contrast, the majority of unirradiated cells exhibited mainly green fluorescence with minimal red fluorescence (Fig. 3Aa). In numerous studies, demonstration of vacuolar H⁺-ATPase-dependent acidification of cellular organelles, as well as its involvement in different cellular processes, was achieved by using its specific inhibitor bafilomycin A1 (23, 24). Similarly, by addition of the inhibitor to MCF-7 cells, we were able to demonstrate that acidification of AVO is mediated by the vacuolar H⁺-ATPase (Fig. 3A, b and c; Ref. 23). Preincubation of the cells with 300 μM of the weak amine chloroquine also inhibited acridine orange accumulation in AVO (data not shown).

To measure the radiation-induced increase in fractional volume and/or acidity of AVO, we determined the mean red:green fluorescence ratio in control and irradiated cells. At 24 h after irradiation, 92% of the unirradiated controls were tightly distributed around their...
mean red:green fluorescence ratio (Fig. 3A). Only 8% had a red:green fluorescence ratio that increased asymptotically above 2 (the highest value in the descending limb of the histogram). On the other hand, 55% of the irradiated cells had a red:green ratio that was above 2, and the mean value of red:green fluorescence ratio was 2.3-fold higher than in controls (Fig. 3B). Bafilomycin A1 decreased the mean red:green fluorescence ratio in unirradiated cells and also inhibited its radiation-induced increase. In the presence of bafilomycin A1, the mean red:green fluorescence ratio was similar in both unirradiated and irradiated cells, indicating that the radiation-induced increase in this ratio is attributable to the development of AVO, rather than other possible changes in the molecular composition of the irradiated cells (Fig. 3B, c and d). AVO appearance was dependent upon the radiation dose and increased with the time after irradiation (Fig. 4).

To differentiate further between the acidic compartments of control and irradiated cells, we used an additional lysosomotropic agent, LysoSensor Blue DND-167. The fluorescence of LysoSensor Blue is pH-dependent and increases as pH decreases. In unirradiated cells, LysoSensor Blue hardly showed any fluorescence, whereas irradiated cells fluoresced bright blue (Fig. 3A, e and f), indicating that the pH of the acidic compartments in irradiated cells is indeed lower than that of unirradiated cells. AVO appearance was associated with increased levels of the lysosomal membrane protein LAMP-1, as evident from Western blot analysis and immunocytochemistry (Fig. 5A and B).

Fig. 2. Ultrastructure of AVO formed in irradiated cells. A, a, control unirradiated cells 48 h postirradiation time (Time 0); b, cells irradiated with 10 Gy, 48 h postirradiation. The arrow points to newly formed AVO. Bar (a–b), 4 μm. B, a–c, newly formed vesicular organelles in cells irradiated as above. The arrows point to the part-rough, part-smooth membrane cisterna (a), to vesicles fusing with membrane cisternae (b), to lamellar structures (c and d), and to residual digested material (e). Bar (a–e), 0.6 μm. C, concentration of the lysosomotropic agent DAMP in AVO (AVO-EM) demonstrated by immunogold histochemistry. Cells were stained with DAMP 24 h postirradiation with 10 Gy and processed for viewing as described in “Materials and Methods.” The arrow points to the gold particles over the AVO (a). Cells were incubated with 0.5 μM bafilomycin A1 (b) or with 300 μM chloroquine (c) before the addition of DAMP. Bar (a–c), 0.6 μm. D, distribution of unirradiated and irradiated cell populations according to the fraction of the cytoplasmic volume occupied by AVO-EM (fractional volume). Fractional volume of AVO-EM was calculated as described in “Materials and Methods.”

Fig. 3. A, detection of radiation-induced appearance of AVO by vital staining with lysosomotropic agents. Acridine orange: a and c, unirradiated cells; b and d, 30 h after irradiation with 10 Gy. c and d, cells were incubated with 200 nM bafilomycin A1 for 30 min before the addition of acridine orange. LysoSensor Blue DND-167: e, unirradiated cells; f, cells 30 h after exposure to 10 Gy. Bar, 18 μm. B, determination of mean red:green fluorescence ratio in acridine orange-stained cells using flow cytometry. The mean red:green fluorescence ratio in irradiated and control unirradiated cells was determined as described in “Materials and Methods.” a and c, unirradiated cells; b and d, 24 h after irradiation with 10 Gy; c and d, unirradiated and irradiated cells preincubated with 500 nM bafilomycin A1 30 min before the addition of acridine orange.
The increase in the red:green fluorescence ratio could be modulated by PMA. The mean red:green fluorescence ratio increased by a factor of 1.7 ± 0.3 (n = 3) 48 h after stimulation with 30 nM PMA. This result suggests that protein kinase C may be involved in the increased red:green fluorescence ratio after irradiation.

The increase in the mean red:green fluorescence ratio after irradiation was also observed in two other cancer cell lines. Forty-eight h after irradiation with 10 Gy, the mean red:green fluorescence ratio increased in prostate cancer (LNCaP) and in colon adenocarcinoma (LoVo) cells by 1.6 ± 0.1 (n = 3) and 2 ± 0.2 (n = 3) -fold, respectively. As in MCF-7 cells, the increase in the mean red:green fluorescence ratio in LoVo and LNCaP cells was associated with the appearance of red fluorescent AVO.

Parallel investigations with electron microscopy confirmed the radiation-induced formation of a new acidic compartment (Fig. 2, A and B). These subcellular AVO were composed of core vesicles with granular, vesicular, or lamellar content. The core vesicles were often surrounded by and intertwined with smooth or part smooth/part rough membrane cisternae that were found to fuse with smooth vesicles of unknown origin (Fig. 2B). The diameter of these organelles ranged from 0.5–2.5 μm and was comparable with the diameter of the largest red fluorescent AVO in irradiated cells. Because fluorescent AVO may consist of a heterogeneous population of AVO, we termed the ones characterized by electron microscopy “AVO-EM.” AVO-EM were found to be acidic by virtue of their ability to concentrate the lysosomotropic agent DAMP (Fig. 2C). By 48 h postirradiation with 10 Gy, the average fractional volume of AVO-EM in the population was 16 ± 0.1% (Fig. 2D), whereas, the average fractional volume in unirradiated cells was 0.91 ± 0.01%. The emergence of AVO-EM during the first 48 h postirradiation with 2–10 Gy was dose- and time-dependent (data not shown).

During autophagy, portions of the cytoplasm and subcellular organelles are sequestered by the endoplasmic reticulum, resulting in vesicular bodies that are bound by double-membrane cisternae (25). The association of the core vesicles with membrane cisternae in AVO-EM bears morphological similarities to autophagous bodies. We therefore examined the effect of 3-methyladenine, an inhibitor of autophagy (25, 26), on AVO formation. 3-Methyladenine at a final concentration of 5 mM decreased the red:green ratio at 48 h postirradiation with 10 Gy from 1.77 ± 0.01 to 1.15 ± 0.01 (n = 3). Electron microscopy analysis demonstrated a parallel reduction in the fraction of cells containing AVO-EM from 94% to 22%. The effect of 3-methyladenine on irradiated cells suggests that the formation of AVO after irradiation may share similar pathways with processes that regulate autophagy.

It is important to note that in addition to ionizing irradiation, other death-inducing agents such as tumor necrosis factor and staurosporin kill MCF-7 without producing typical apoptotic changes (27). It has recently been reported that the lack of apoptotic response to tumor necrosis factor results from the absence of caspase-3 in these cells (27). The absence of caspase-3 may well explain the lack of apoptotic response to ionizing irradiation. Nonetheless, the emergence of AVO...
in the presence of the pan-caspase inhibitor z-VAD-fmk at concentrations ranging from 50–154 μM (Table 1) suggests that the programmed events that lead to AVO formation are not related to apoptosis.

Cells that survive radiation may continue to divide and form colonies, although their DNA might have sustained damage (28). We found that the progenies of irradiated cells contain an increased level of AVO (Table 2). This led us to postulate that the emergence of acidic compartments protects the cells against radiation damage. In fact, experiments with bafilomycin A1 showed that inhibition of vacuolar H⁺-ATPase, the enzyme that mediates AVO acidification, augmented DNA degradation and decreased survival after irradiation (Fig. 6; Table 2). Addition of bafilomycin A1 2 days after irradiation with 10 Gy, for a period of 24 h, dramatically increased DNA cleavage into large fragments (20–1000 kb). Also, addition of bafilomycin A1 for 24 h after irradiation with 2 and 3 Gy reduced the surviving fraction by 30–40% without significantly affecting the survival of unirradiated cells.

Increased autophagy, the hallmark of programmed cell death type II, is thought to lead to cell death via destruction of the cytoplasm. Still, the lysosomal compartment has been linked to cellular defense mechanisms such as protection against infectious agents (29). Recently, acidic compartments have been associated with drug resistance of breast cancer cell lines (30), and in yeast autophagy is required for cell survival during starvation (31). Similarly, our results suggest that accumulation of AVO after irradiation is modulated by cellular defense mechanisms. These AVO may protect the cells by preventing cytoplasmic acidification, by providing catabolites required for repair processes, and/or by containing toxic molecules. Our experiments show that moderate formation of AVO in surviving colonies provides long-term protection against low-radiation damage. However, continuous accretion of AVO after high levels of damage may offset their protective effect, leading to replacement of the normal cytoplasm and possibly to necrosis and cell death. Therefore, inhibition of AVO formation or function may serve as a tool to increase cell death after low-radiation damage and facilitate cell-kill after high-radiation damage. Modulation of AVO function may prove useful for increasing the therapeutic ratio of radiation treatment of epithelial cancers.

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