Oxidative stress-induced cell cycle blockage and a protease-independent programmed cell death in microaerophilic *Giardia lamblia*

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**Abstract:** *Giardia lamblia* is a microaerophilic human gastrointestinal parasite and considered as an early-diverged eukaryote. *In vitro* oxidative stress generation plays a significant role in cell cycle progression and cell death of this parasite. In the present study hydrogen peroxide, metronidazole, and a modified growth medium without cysteine and ascorbic acid have been chosen as oxidative stress-inducing agents. Cell cycle progression has been found to be regulated by different types of oxidative stresses. Apoptosis is not an established pathway in *Giardia*, which is devoid of ideal mitochondria, but in the present investigation, apoptosis-like programmed cell death has been found by the experiments like Annexin V-FITC assay, DNA fragmentation pattern, etc. On the contrary, Caspase-9 assay, which confirms the caspase-mediated apoptotic pathway, has been found to be negative in all the stress conditions. Protease inhibitor assay confirmed that, even in absence of any proteases, programmed cell death does occur in this primitive eukaryote. All these results signify a novel pathway of programmed suicidal death in *Giardia lamblia* under oxidative stress. This is the first demonstration of protease-independent programmed cell death regulation in *Giardia* exclusive for its own specialties.

**Keywords:** *Giardia lamblia*, oxidative stress, reactive oxygen species, programmed cell death, apoptosis, early branching eukaryotes

**Introduction**

Living aerobic organisms, from prokaryotes to complex eukaryotes, have developed elaborate sequences of adaptive mechanisms to maintain oxygen homeostasis and equilibrium.¹ Any deviation from homeostasis, or physiological change in oxygen pressure, is recognized as an exposure to oxidative stress.² *Giardia lamblia* is an intestinal protozoan parasite causing malabsorption and diarrhea (giardiasis) in human. It is an early diverging eukaryote with unicellular, binucleated, flagellated structure, without ideal mitochondria, golgi complex or peroxisomes.³,⁴ Measurement of O₂ consumption as a function of dissolved O₂ indicates that at low levels (0–50 μM), the organism is capable of scavenging O₂ (apparent *Kₘ* for O₂ 6.4 μM for the trophozoite). On the contrary, in human gut cell lining, *Giardia* resides in a higher oxygen tension where the O₂ concentration has been measured at 60 μM.⁵ This fact has become more interesting as the conventional enzymes for detoxifying reactive oxygen species (ROS) (*viz.* superoxide dismutase [SOD], catalase, peroxidase, glutathione reductase) are not found in *Giardia*.⁶ Change in the oxygen environment has played the key role in the evolution of aerobic organisms from anaerobes. In case of *Giardia*, it is also evident that change in oxygen pressure has achieved an important role in its survival strategy as well as the evolution and adaptation of this parasite.

In the present study, three conditions have been chosen to generate oxidative stress in the trophozoites *in vitro*. First, hydrogen peroxide (H₂O₂) is a very well known
chemical reagent that can generate free oxygen radicals very easily. Secondly, metronidazole is a commonly used drug against giardiasis in the most developing countries. Its mode of action involves free nitro radical generation within the organism. Third one is a modified medium that is devoid of cysteine and ascorbic acid. Cysteine has been found to protect *Giardia* trophozoites from thiol-blocking reactants, indicating a role of reducing agent for protection of crucial thiol groups. Ascorbic acid also protects the trophozoites under high pO₂. These three conditions would give us an idea about the effects of different modes of oxidative stresses on *Giardia* cells. Cell cycle analysis using flow cytometer has been carried out to check the effects of oxidative stress on survival of parasitic cells. The changes in whole cell morphology have been studied in each case using transmission electron microscopy (TEM). As mitochondria regulates the caspase-dependent apoptotic cell death pathway in higher eukaryotes and no ideal mitochondria has been reported in *Giardia* cells, it was our endeavor to identify whether a programmed cell death (PCD) is possible in this early divergent eukaryote. Several cell death assays have been performed to verify whether the mode of death is apoptotic or necrotic. Protease inhibitor assay has also been performed to check if other proteases like serine/cysteine proteases, calpains, fragmentin/granenzymes, etc are playing any role in parasitic death as they are reported in higher eukaryotes.

Oxygen concentration always plays a vital role in the survival strategy and commitment of death from lower groups of organisms to higher eukaryotes. With the changes in global oxygen concentration from one era to another along thousands of years, lives on earth have evolved themselves from obligatory anaerobes to obligatory aerobic system via several facultative and mutual adaptations. Currently, the detail mechanism is not clear on how these severe changes occurred gradually with time. Studies on oxidative stress regulation in *Giardia* and similar early branching eukaryotes (EBE) may shed some new ideas about these changes.

**Materials and methods**

**Giardia culture maintenance**

*Giardia lamblia* Portland I was maintained axenically in TYIS-33 medium at 35.5 °C and sub-cultured at each 48 h.

**Standardization of the conditions generating oxidative stress**

Dose and time kinetics of the three different oxidative stress-generating conditions (as described above) have been standardized following the IC₅₀ values as reported previously. Finally from the standardized data, 0.1 μM H₂O₂ and 1 μg/ml metronidazole have been administered in all the experiments mentioned below.

**Cell fixation and staining for cell cycle analysis in flow cytometry**

Two × 10⁶ cells were harvested from each culture sets (control and three experimental sets) and the cell pellet were taken by centrifugation at 2000 rpm for 10 min. Cells were washed with phosphate buffered saline (PBS) and treated with methanol for fixation and then in 1% Triton X for membrane permeabilization. RNase treatment was carried out prior to propidium iodide (PI) staining of the cells. It was incubated on ice and then used for cell cycle analysis using flow cytometer (FC 500; Beckman Coulter, Miami, FL, USA).

**Cell death assays**

**Cell morphology study by electron microscopy**

Control and all sets of experimental samples were harvested (∼10⁶ trophozoites in each case), prefixed with 3% gluteraldehyde in 0.1 M sodium cacodylate buffer (pH 7.4) and post fixed in 1% osmium tetroxide in 0.1 M sodium cacodylate buffer (pH 7.4). The post-fixed cells were block stained in 1% uranyl acetate and then treated with graded series of ethanol and subsequently embedded in agar 100 resin (Agar Aids, Stansted, UK). Ultra thin sections were cut in a Leica Ultracut UCT ultramicrotome and stained with uranyl acetate and lead citrate. Sections were examined in a Tecnai 12 Bio-twin transmission electron microscope (FEI, Hillsboro, OR, USA).

**Annexin V-FITC assay**

This experiment was performed using Annexin V-FITC Kit (Beckman Coulter), following the manufacturer’s protocol. Briefly, cell samples (Control cell as well as oxidative stressed cells) were washed with chilled PBS and cell pellet was resuspended in 1 × binding buffer and kept on ice. Annexin V-FITC solution was added and incubated on ice in dark. Cell preparation was analyzed using confocal microscope (LSM 510, Meta; Carl Zeiss, Thornwood, NY, USA).

**FITC-PI staining**

For this experiment the cells were similarly treated with Annexin V-FITC solution (Beckman Coulter) as per manufacturer’s protocol as mentioned above. Along with that, PI solution was also added and incubated on ice in the dark. This cell preparation was analyzed for its immunofluorescence using confocal microscope and flow cytometer.
DNA fragmentation
Permeabilized cells (as prepared above) were treated with PI to observe the DNA damage in nucleus under confocal microscope.

DNA laddering (degradation)
Control cells and stressed cells were harvested and resuspended in digestion buffer (10 mM EDTA, 50 mM Tris, 0.5% SDS Sarcosine, pH = 8.0) containing 0.5 mg/ml proteinase K. Samples were incubated at 37 °C for 1 h after adding DNase-free RNase (0.1 mg/ml). After phenol–chloroform treatment, salt precipitation and 70% ethanol wash, pellet was air dried and resuspended in autoclaved triple distilled water and checked in a 1% agarose gel stained with ethidium bromide.

Caspase-9 assay (Colorimetric)
This assay was performed using Caspase-9 Assay Kit (Calbiochem®, San Diego, CA, USA) following manufacturer’s protocol. Briefly, all the control and stressed cell pellets were resuspended in cell lysis buffer provided with the kit and incubated on ice for 10 mins. It was then centrifuged and the supernatant (cytosolic extract) was collected in a tube, kept on ice. After measuring protein concentration it was dissolved in 50 μl cell lysis buffer. Then reaction buffer with 10 mM DTT was added to each sample. Finally LEHD-pNA substrate (200 μM final concentration) was added, incubated at 37 °C and samples were read at 400–405 nm.

Protease inhibitor assay
Mixture of several protease inhibitors (a broad range protease inhibitor; Roche, Nutley, NJ, USA) were added to the culture and incubated for 1 h. Small water soluble molecules of protease inhibitors were absorbed by the cells along with medium. Then the cells were treated with oxidative stress producing agents. Annexin V-FITC assay and DNA-laddering assay were repeated to check if programmed cell death occurred in absence of proteases.

Results and discussion
Cell cycle analysis in flow cytometry
In case of Giardia, normal trophozoites of G1 phase contain 4 N chromosome.12 Hence, in control set, maximum numbers of cells lie in G0/G1 phase (Figure 1i). In case of metronidazole, as observed in flow cytometer, the cell cycle was arrested near S-phase (Figure 1ii). Metronidazole is known to exert its toxic effect through unstable intermediate products (nitro and nitroso-free radicals).14 The reactive products also interact with DNA, causing DNA damage, strand breakages, and cross-linking in the nucleic acid and the release of nucleotide phosphates. Hence in our study, the growth arrest of trophozoites near S-phase of cell cycle is due to a similar action of metronidazole. It probably inhibits the DNA segregation and hence prevents the completion of cell division. Peroxides have been found to induce a G1 checkpoint response (Figure 1iii). It suppresses S phase entry by inhibiting cyclin E/Cdk2 activity in a dose-dependent manner.13 In this experiment, we have also found that the maximum number of cells lie in sub-G1 stage. It might also be an indication of apoptotic cell death due to peroxide application. For cystein-ascorbate deprived modified medium, most of the cells move to apoptotic death, but a significant amount has a tendency to increase their DNA content (Figure 1iv).

![Figure 1](image-url) Cell cycle analysis of control and treated cells. D, E, and F are showing sub-G1*, G1 and G2/M phase in the trophozoite cell cycle. Each unit distance in the X-axis denotes 2N DNA content of the trophozoites. Normal Giardia trophozoite contains 4N genomic DNA content in G1 phase. i) In the control set, maximum no. of trophozoites are showing 4N DNA content with few sub G1 population indicating apoptotic cell death. ii) Metronidazole treated cells stop their cell cycle propagation just at synthesis phase and commit apoptotic type of cell death. Cells lie in between D to late E region. iii) Effect of H2O2 on Giardia trophozoites is very prompt. All the trophozoites undergo apoptotic type of death and lie sharply in the D region, sharply within sub G1 stage. iv) Cells treated with modified medium show a group of trophozoites in D region with sub-G1 phase, but few cells are moving beyond S phase towards G2, indicating a tendency to encystation in stressed condition.

Notes: *Sub-G1 implies that the DNA content is less than usual G1 stage (4N in case of Giardia). In the dead cells when nuclei start degradation then another peak arises in flow cytometer just left of the original G1 peak, indicating that the DNA content is decreasing.
Cell death assays

Cell morphology study by electron microscopy
Oxidative stress induced cells have shown apoptotic features like membrane blebbing and increased vacuolation in comparison to control cells under transmission electron microscopy (Figure 2).

Annexin V-FITC assay
Externalization of phosphatidyl serine (PS) is one of the indicators of apoptosis in the eukaryotic cells. Fluorescein isothiocyanate (FITC)-tagged annexin can bind with the PS when it comes outside the membrane and the fluorescence signal can be measured in a fluorescent microscope. A characteristic green ring formation can be observed in the membrane of an ideal apoptotic cell. In our study we have used this assay to check the mode of death in stress-induced cells. Normal live cells do not show any ring formation whereas in case of all the experimental sets treated with different oxidative stress producing agents revealed a clear ring formation in their outer membrane part (Figures 3A, B, D, and F) confirming the mode of cell death to be apoptotic.

FITC-PI staining
Cells stained with FITC and PI can be targeted for observing their viability and mode of death in a flow cytometer. The gated region in the flow cytometer denotes the type of cells according to their location in the four coordinates. Live cells do not take the stains and lie in the left hand bottom panel whereas apoptotic cell population taking FITC stain lies in right hand bottom panel. All the secondary necrotic cells taking the PI stain lie in the right hand upper panel. In our experiment, control live cells did not take any type of stains and consequently lied in the left hand bottom panel (Figure 4A). All the experimental sets showing cell populations in the right hand bottom panel indicate the intake of FITC stain specific for apoptotic death (Figures 4B, C, and D).

DNA degradation
Cell nucleus takes the PI stain as this dye intercalates in the DNA. In normal live cells, the two nucleus of Giardia can be easily identified after PI staining (Figure 5A). DNA degradation is another hallmark for apoptotic type of cell death. Hence, after the onset of apoptosis the genetic material gets dispersed throughout the cell and dispersed red patches of the dye can be visualized under fluorescent microscope. In all the experimental sets treated with different stress-producing agents, DNA degradation has been observed (Figures 5B, D, and F).

DNA laddering (fragmentation)
Cells undergoing apoptosis show characteristic morphological and biochemical features. The biochemical marker

Figure 2 Electron microscopic view of control and treated cells. A) Live Giardia cell with two nuclei and flagella. B) Live Giardia (mainly two nuclei are focused). C) Giardia cells treated with H$_2$O$_2$: Nuclear membrane blebbing due to PCD. D) Giardia cells treated with metronidazole: Increased vacuolation and outer membrane blebbing due to PCD. E and F) Giardia cells treated with modified medium: Increased vacuolation and nuclear membrane and outer membrane blebbing due to PCD. G) Change in cell morphology after treated with modified medium.

Abbreviations: N, nucleus; P, plasma membrane.
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**Figure 3** Annexin V-FITC assay. A) normal live Giardia cells, B) Giardia cells treated with H$_2$O$_2$ for 2 h. C) Giardia cells treated with protease inhibitor cocktail and then further treated with H$_2$O$_2$ for 2 h. D) Giardia cells treated with metronidazole for 4 h. E) Giardia cells treated with protease inhibitor cocktail and then further treated with metronidazole for 4 h. F) Giardia cells grown in modified medium deprived of Cys-ascorbate for 6 h. G) Giardia cells grown in modified medium deprived of Cys-ascorbate with protease inhibitor cocktail. Panels B to G clearly show ring formation due to apoptosis.

**Figure 4** Flow cytometric analysis of FITC-PI stained cells. A) Control live cells lying in the left hand bottom quadrant (B3) as the cell population has not taken any of the two dyes, FITC or PI. B) H$_2$O$_2$ treated cells after one hour takes FITC and lie in the right hand bottom quadrant (B4) indicating apoptotic type of death. Similarly, in C) and D) cells treated with metronidazole and modified medium respectively, show their mode of death in apoptotic way.
for apoptosis is the degradation of the genomic DNA, an irreversible event for the programmed cell death and occurs before changes in plasma membrane permeability. DNA fragmentation (mainly in lower molecular weight region) is the characteristic for apoptosis in some lower eukaryotes. Distinct DNA laddering is absent in these organisms. Compared with the control one, all the test samples are showing sheared fragmented DNA in low molecular weight region (Figure 6).

Colorimetric caspase-9 assay

Higher eukaryotes having developed apoptotic machinery undergo caspase dependent pathway to carry out apoptosis. Caspase 9 initiates the signaling pathway. It is expected that in a programmed cell death, colorimetric caspase-9 assay will produce a positive result confirming caspase-dependent apoptotic pathway. In our study, a negative result was obtained (Table 1) indicating that the programmed cell death in this parasite is devoid of any caspase-signaling pathway.

Protease inhibitor assay

In some cases where caspase is not activated in programmed cell death, other proteases such as serine/cysteine proteases, calpeins, or granenzymes, etc get involved in the pathway. So, protease inhibitor assay was performed to check whether other proteases except caspase control the programmed cell death in *Giardia*. The cells were treated

![Figure 5 Degradation of nuclei due to programmed cell death in *Giardia*. A) normal *Giardia* cells stained. B) *Giardia* cells treated with H₂O₂ for 2 h and PI stained. C) *Giardia* cells treated with protease inhibitor cocktail and then further treated with H₂O₂ for 2 h and PI stained. D) *Giardia* cells treated with metronidazole for 4 h and PI stained. E) *Giardia* cells treated with protease inhibitor cocktail and then further treated with metronidazole for 4 h and PI stained. F) *Giardia* cells grown in modified medium deprived of Cys-ascorbate for 4 h and PI stained. G) *Giardia* cells grown in modified medium deprived of Cys-ascorbate with protease inhibitor cocktail and PI stained. Panels B to G clearly show degradation of nuclear content. The phase images in all seven pictures are shown in the right side of the individual pictures.](image-url)
observed. The DNA marker ranges from 1–10 kb.

with a large-scale protease inhibitor cocktail before applying the stress-producing agents. Surprisingly, it was found that in absence of proteases, the trophozoites still could undergo PCD. The result has been verified by Annexin V assay (Figures 3C, E, and G) and DNA degradation assay (Figures 5C, E, and G). Characteristic green fluorescent ring formation and scattered PI staining were obtained in each set, respectively.

A similar type of report has been obtained in the case of *Trichomonas vaginalis*. PI has shown typical DNA degradation in the dead cells. But it has been found that caspase or other proteases are not responsible for this type of death. Cells, after incubating with protease inhibitors when under stress, still committed death in programmed pathway. This was confirmed by Annexin V test and DNA fragmentation pattern, as before. Two cases in *Leishmania* and *Blastocystis* have been reported where also the PCD is independent of caspase pathway or other proteases. It is possible that these EBE including our model organism *G. lamblia* have undergone PCD using a novel pathway.

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**References**


