PMF, Cesium and Rubidium Nanoparticles Induce Apoptosis in A549 Cells


Abstract—Cancer becomes one of the leading cause of death in many countries over the world. Fourier-transform infrared (FTIR) spectra of human lung cancer cells (A549) treated with PMF (natural product extracted from PM 701) for different time intervals were examined. Second derivative and difference method were taken in comparison studies. Cesium (Cs) and Rubidium (Rb) nanoparticles in PMF were detected by Energy Dispersive X-ray attached to Scanning Electron Microscope SEM-EDX. Characteristic changes in protein secondary structure, lipid profile and changes in the intensities of DNA bands were identified in treated A549 cells spectra. A characteristic internucleosomal ladder of DNA fragmentation was also observed after 30 min of treatment. Moreover, the pH values were significantly increases upon treatment due to the presence of Cs and Rb nanoparticles in the PMF fraction. These results support the previous findings that PMF is selective anticancer agent and can produce apoptosis to A549 cells.

Keywords—Apoptosis, FTIR spectroscopy, pH therapy, Scanning Electron Microscope- Energy Dispersive X-ray (SEM-EDX).

I. INTRODUCTION

NANO TECHNOLOGY has considerable promise for the detection, staging and treatment of cancer [1]. The past two decades have witnessed rapid advances in the ability to

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structure matter at the nanoscale with sufficient degree of control over the material size, shape, composition, and morphology [2]. The combination of the unique properties with the appropriate size has motivated the introduction of nanostructure into biology [3]-[6] Cells and their constituent organelles lie on the sub-micron to micron size scale. Further, proteins and macromolecules found throughout the cell are on the nanometer size scale. [7]. Thus nanoparticles ranging from a few to a hundred nanometers in size become ideal as labels and probes for incorporation into biological systems[5],[6]. Despite significant investment and research, cancer is still responsible for 25% of all deaths in developed countries[8]. There is a pressing need for more sensitive, selective and cost-effective methods for detecting and treating cancer.

The presence of natural occurring nanoparticles that can produce apoptosis in cancer cells would be of great help. PMF, the active fraction separated from PM 701 [(natural product previously proved to be selective anticancer agent) [9]-[15] would give a promising therapeutic criterion in treating cancer. The effect of PMF on lung cancer cells A549 will be monitored by Fourier-transform infrared (FTIR) spectroscopy.

FTIR spectroscopy has become a useful analytical tool in biomedical science in the past decade, e.g., for the characterization of microorganisms[16], isolated cells or cell lines[17-20], body fluids and tissues [21]-[26] The ability to detect drug action, disease or dysfunction rapidly has obvious benefits, including early intervention of therapeutic strategies, hopefully in a prognostic fashion, significant reduction in mortality and morbidity, and the freeing up of much needed economic resources within health care systems [27]. With these highly sensitive techniques, the frequency and the intensity of light in the resulting spectrum provides biochemical information regarding the molecular composition, structure and interaction in cells and tissues.

Apoptosis, as a pre-programmed physiological mode of cell death, plays an important role in the pathogenesis and progression of cancer. Understanding of the basic mechanisms that underlie apoptosis will point to potentially new targets of therapeutic treatment of diseases that show an imbalance between cell proliferation and cell loss. As an active process, apoptosis involves biochemical changes on three essential cellular components, DNA, protein and lipid. There are three steps in apoptosis, the initiation phase triggered by a stimulus received by the cell, the decision phase during which the cell commits itself to live or to die, and the degradation phase
when the cells acquire the morphological and biochemical hallmarks of apoptosis [28]. Clearly, it would be desirable to detect apoptosis at an early stage, i.e. before phase that entails the visible changes of apoptosis.

In this study we will explore the potential of PMF as a selective anticancer agent which produces apoptosis in lung cancer cells (A549) by using FTIR spectroscopy technique and to bring to light the role of Cesium (Cs) and Rubidium (Rb) nanoparticles - found in PMF- that thought to be one of many other reasons responsible of the selectivity of PMF fraction.

II. MATERIALS AND METHODS

A. Media

The following commercially available media were prepared according to published literature, these include: Ordinary media, Minimal essential medium (MEM) (10%FCS); MEM is a rich, multipurpose medium that was used for cultivation of human lung cancer cells (A549), Phosphate Buffer Saline (PBS) is a Phosphate-Buffered physiological Saline solution. Calcium and Magnesium free Solution Trypsin. [29] Examined media: PMF (extracted from PM 701) is a natural product, easily available, cheap, sterile, and non-toxic according to our chemical and microbiological testing. PMF added to the ordinary media with ratio 2.5 μg : 1 ml media

Human Lung Cancer Cells line: Human Lung Cancer Cells, non- small cell carcinoma (A549) was obtained from cell strain from (ATCC) American Type Cultural Collection, available in the cell bank of Tissue Culture Unit, King Fahd Medical Research Center (KAU, Jeddah).

In vitro proliferation of cells
1. Human Lung Cancer Cells (A549) were suspended in culture medium MEM.
2. The cells were dispensed in 3X (6wells plate), 1x10^5/ml culture medium MEM.
3. Each group of cells was incubated 24 hrs in suitable media and apoptosis was induced by incubating cells in PMF at different time points.

B. Infrared spectroscopy

At various time points, the cells treated with PMF were harvested and washed twice (by centrifugation for 3 min at 300 g). The cell pellet was kept at -80°C and freeze dried proier to IR measurments. The lyopholized samples - from three different separated experiments - were dispersed in potassium bromide (KBr) discs by mixing them gently in an agate mortar and with pestle to obtain homogenous mixture as described in [14, 30]. The mixture then pressed in a die at 5 metric tons force for 3 s, creating a 1.1 cm diameter transparent disc with imbedded sample. The FTIR spectra were recorded in absorbance form using Shimadzu FTIR-8400 s spectrophotometer with continuous nitrogen purge. The spectra were obtained in the wavenumber range of 4000-400 cm\(^{-1}\) with an average of 20 scans to increase the signal to noise ratio and at spectral resolution of 4 cm\(^{-1}\). All the samples were baseline corrected and normalized to amide II band by using IR Solution software. The parameters studied were nucleic acid, proteins and lipids. Second derivative and difference method measurements were taken in comparison studies.

C. Scanning Electron Microscopy (SEM)

Detecting Cs and Rb nanoparticles in PMF:

For SEM studies the PMF were suspended in distilled water, then treated in an ultrasonic bath (BRANSON, 1510) about 20 min. A small drop of this suspension placed on the double side carbon tape on Al- Stub and dried in air. The specimens were analyzed - without gold coating- by using energy dispersive analyzer unit (EX-23000BU) which attached to the scanning electron microscope (JSM-6360LA, JEOL, Tokyo, Japan). The microscope was operated at an accelerating voltage of 20 kV. Quantitative method ZAF, and characterization method as pure.

D. Cs and Rb level measurements

Cs and Rb levels were measured in The Ministry of Petroleum-The Egyptian Mineral Resources Authority- Central Laboratories Sector by using Inductively Coupled Plasma (ICP) the results were multiplied by the dilution factor 20.1.

E. DNA fragmentation and pH measurements

DNA was extracted according to the method followed in [23] by electrophoresis in agarose gel. The genomic DNA was run in a 1.5% agarose gel for 20min.

The pH values of the A549 cell's media were measured immediately by [Jenway 3200] at the end of each treating time. These measurements were repeated for three different experiments.

III. RESULTS AND DISCUSSION

A. IR spectral features and assignments

The addition of PMF to A549 cells after 5, 15, 30 min and 2, 24hr of treatment caused a series of changes in the spectra of these cell lines (Fig.1). The absorption profiles are typical of tissue infrared spectra, which are dominated by protein, lipids and DNA. (Table I) represents the most absorption IR spectral peaks and the proposed biomolecular assignments. Fig.1a. shows a noticeable decrease in the intensities of proteins (amide I & amide II), nucleic acids band, and an increase in the intensity of the weak shoulder located at 1737
Fig. 1 Raw FTIR spectra of A549 cells treated with PMF at different times (a) shows the amide I band shift and the increase in the intensity of the band located at 1735 cm⁻¹ (C=O ester group) in range 1760-1400 cm⁻¹. (b) shows the decrease in DNA peaks at the onset of apoptosis (5-15min) and the increase in 970 cm⁻¹ peak after 30 min of PMF treatment. (c) shows the band shift in CH₂ bands.

Table I

<table>
<thead>
<tr>
<th>Wavenumber (cm⁻¹)</th>
<th>Proposed biomolecular assignment</th>
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<tbody>
<tr>
<td>2963</td>
<td>C-H stretch (asymmetric) of CH₃ in fatty acids, lipids, proteins.</td>
</tr>
<tr>
<td>2926</td>
<td>C-H stretch (asymmetric) of CH₂ in fatty acids, lipids, proteins.</td>
</tr>
<tr>
<td>2873</td>
<td>C-H stretch (symmetric) of CH₃ in fatty acids, lipids, proteins.</td>
</tr>
<tr>
<td>2853</td>
<td>C-H stretch (symmetric) of CH₂ in fatty acids, lipids, proteins.</td>
</tr>
<tr>
<td>1737</td>
<td>Ester C=O stretching of phospholipids.</td>
</tr>
<tr>
<td>1652</td>
<td>Amide I of α-helix protein structures (C=O) stretch.</td>
</tr>
<tr>
<td>1542</td>
<td>Amide II (N-H) bend</td>
</tr>
<tr>
<td>1460</td>
<td>CH₃ antisymmetric bend</td>
</tr>
<tr>
<td>1430-1360</td>
<td>COO⁻ stretch</td>
</tr>
<tr>
<td>1240</td>
<td>PO₃⁻ asymmetric stretch of phosphodiester group in nucleic acids and phospholipids</td>
</tr>
<tr>
<td>1168</td>
<td>C-O, C-C stretch, C-O-H, C-O-C deformation of carbohydrate threonine, tyrosine in cell proteins</td>
</tr>
<tr>
<td>1085</td>
<td>PO₃⁻ symmetric stretch of phosphodiester group in nucleic acids and phospholipids</td>
</tr>
<tr>
<td>975</td>
<td>PO₄²⁻ stretch</td>
</tr>
</tbody>
</table>

Spectral assignment taken from references [16]-[26].
Early apoptotic cells appear in the cell cycle distribution as cells with a hypodiploid DNA. This alteration in DNA content results from degradation of cellular DNA by activation of endogenous endonucleases during apoptosis [36]. Cells in the pre-Go/G1 phase (M4) were therefore defined as apoptotic cells. These results are also consistent with the finding that at the early stage of apoptosis, initiation phase is triggered by a stimulus received by the cell, this can be explained by the first disappearance of nucleic acid bands at 975 and 923 cm\(^{-1}\) at 5 and 15 min upon PMF treatment. A549 cells treated up to 30 min with PMF showed clear absorbance bands of DNA at 975 and 923 cm\(^{-1}\) which may referred to the degradation of DNA. Reference [41] reported a large increase in the DNA signals of apoptotic cells. In apoptosis DNA is degraded into oligonucleotides with a few hundreds of base pairs that subsequently diffuse out of the nucleus. This observation suggests that the DNA, although highly condensed in the nucleus, has a low absorption, yet its degradation products in late apoptosis are detected by IR spectroscopy.

Fig. 2 represents the raw IR spectra of A549 cells treated with PMF from 5 min up to 24 hr without normalization. The figure shows a dramatic decrease in bands intensities over the range 1800-900 cm\(^{-1}\). The same results were obtained from normal human fetal lung fibroblasts IMR-90 in G1, S-, and G2/M-phase of the cycle [39]. Hoi-Ying et al., 2000 showed that the IR spectra of these three phases are clearly different. During the S-phase the DNA was undergoing replication and they observed that the absorption in the DNA/RNA spectral region increased relative to the G1-phase spectra. When a G2/M-phase cell was measured they observed a large increase in the overall absorbance (like the uppermost spectrum in Fig. 2. This may have been a result of more material in the cell or because of the thickness may have been different in the M-phase.

B. Second Derivative Analysis

Since apoptosis involves the modification of existing proteins, as well as the synthesis of new proteins [35, 39] we analyzed the changes of protein structure after PMF treatment in more detail. Fig. 3 shows only the mean spectra of control and PMF-treated cells for 30 sec - for clarity reason - in the region of the amide I (1600-1700 cm\(^{-1}\)) and amide II (1500-1600 cm\(^{-1}\)) bands. Spectra are shown as second derivative, a method commonly used for narrowing broad IR bands for better visualization. The spectrum of a dying cell due to treatment shows significant shifts in the amide I peak compared to the spectrum of a living A549 cell. The changes in the secondary structure of the proteins appeared after 15 min of treatment. The dying cell shows two characteristic spectral signatures indicative of death [36, 41]. First, the centroids of the protein amide I and II peaks shift to lower energy, indicating a change in the overall protein conformational states within the cell. Second; the appearance of a peak at around 1743 cm\(^{-1}\) [39]. These structural changes in the cellular proteins could be due to a different distribution of proteins during apoptosis or to denaturation of the existing proteins. The lipids in the plasma membrane are composed mainly of phospholipids that determine membrane stability, fluidity and membrane enzymatic activity [40]. Thus, monitoring lipid absorbance in the treated samples is an important for detecting apoptosis. The peak at 1737 cm\(^{-1}\) - which is associated with the non-hydrogen bonded ester carbonyl C=O - splits into two bands in the second derivative analysis (Fig 3.) centered at 1743 and 1725 cm\(^{-1}\). the reason is that the 1743 cm\(^{-1}\) peak is significantly more intense than the 1725 cm\(^{-1}\) peak implies that the C=O ester carbonyl groups of lipids in the cell are becoming predominantly non-hydrogen bonded, which would be in agreement with oxidative damage having occurred.
Apoptosis is associated with, among other factors, increased oxidative damage [42, 43]. The positive intensity at 1740cm$^{-1}$ suggest an increase in the C=O ester components in the plasma membranes following PMF treatment.

C. Difference Method

In order to further assess and identify these membrane changes, typical IR marker bands for lipids were evaluated by creating a difference spectrum in the range of 2800-3050 cm$^{-1}$ Fig. 4(a).

The two methyl bands at 2870 and 2954 cm$^{-1}$, which originate from the symmetric and asymmetric stretching vibrations of the acyl chain CH$_3$ groups, are negative in the IR 'difference spectrum' in contrast, the two methylene bands at 2851 and 2931 cm$^{-1}$, attributed to the symmetric and asymmetric stretching vibrations of acyl chain CH$_2$ groups are positive, indicating that PMF treatment increases the lipid content in the plasma membranes of A549 cells.

The characteristic CH$_3$ group stretching vibration bands at 2870 & 2954 cm$^{-1}$ show a negative band in the difference spectrum. These results may reflect structural changes of phospholipids rather than relative changes of lipid in the treated cells [34], [35]. PMF contains mainly peptides and amino acids [44]. Thus, PMF may contact with A549 cell membrane and may form some sort of an ion channel or pore in the cell membrane thereby, enhancing the ion permeability of the membrane and destroying the cell i.e. induce apoptosis. References [45]-[47] suggested that the peptide could perturb membrane functions responsible for osmotic balance.

The mode of action of PMF with the membrane phospholipids bilayers could possibly be due to the aggregation of peptides on the surface of the membrane and then inserts into the bilayer forming a so-called barrel-stave pore. Another possibility is that the accumulation of peptide on the membrane surface forms a destabilizing carpet, leading to a local disintegration of the bilayer. A third possibility is that peptide chains aggregate on the membrane surface, possibly forming a carpet, partially submerge, and then blend with lipids so that peptide chains and lipid head groups together line the wall of a toroidal pore [35], [44]-[48].

The present and our previous study [14] extend these findings and suggest that PMF may alter the conformation of membrane proteins such as nucleoside transporters, ion channels and proof the formation of pores.

The Transmission Electron microscope (TEM) Photograph (Fig. 5) was taken after 5 sec only from adding PMF to the media "unpublished paper" [49]. The arrow in the Fig. pointed to the entrance of nanoparticles to inside the nucleus via temporary created artificial pore (T) in the nuclear envelope. The nanoparticle interpenetrated the nuclear envelope and the underneath euchromatin. The normal nuclear pore was also seen in other place, whereas no underneath chromatin prevent the normally entrance and exit of molecules.

![Difference spectra between control A549 cells and that treated with PMF for 5 min. The figure shows negative DNA bands (930cm$^{-1}$, 1085cm$^{-1}$ and 1240cm$^{-1}$), negative band of proteins (1652cm$^{-1}$) and positive band of lipids (1735cm$^{-1}$).](image)
PM701 on cancer cells started immediately after 5-6min since adding the examined substrate. They continued that the cancer cells incubated in PM701 showed that the substrate attacks the cell's nuclei, which is indicated by the appearance of pale rings around the nucleus of the lung cancer cell after 30 min of incubation. This leads to the degradation of the cells, which could not be reversed to recover the cells by re-growing the cells in ordinary media again.

**D. Electrophoresis and pH measurements**

We extracted DNA from A549 control and the treated cells to confirm the finding that PMF induces apoptosis and analyzed the DNA fragmentations using gel electrophoresis. A characteristic internucleosomal ladder of DNA fragmentation was observed in A549 cells treated with PMF for 30 min (Fig. 6). These results are in good consistent with the above results obtained from infrared spectroscopy observations.

The pH values of the A549 cells were measured as described earlier. The relationship between the time of treatment and the pH values were shown in Fig. 7. For all times of treatment, the pH values increased significantly after adding PMF to the media of A549 cells, the maximum value was obtained at 2hr of treatment. Our results consistent with the previous reported results describing the pH value of cancer cells and the using of high pH therapy in cancer patients [48], [51], [52]. The cancer cells contain high amounts of hydrogen ions rendering them acidic and they also contain higher Na$^+$ levels than found in normal cells. If Cs$^-$ or Rb$^-$ enters the cancer cells, their pH increases. At a pH of 7.6 the cancer cell division will stop, and at a pH of 8.0 to 8.5 the life span of cancer cells is considerably shortened to only hours [51]. Cs, Rb, and the other elements were detected in PMF by using SEM-EDX (Fig. 8) (Table II). The presence of Cs or Rb in the fluids adjacent to the tumor cells is believed to raise the pH of the cancer cells where cell mitoses will then cease resulting in reduction of life span of the cancer cell [52].

The mechanism of action of Cs in cancer has been little studied. That both Cs$^-$ and Rb$^-$ can specifically enter cancer cells and embryonic cells, but not normal adult cells, [51].
PMF contain 70.9128, 8.04 ppm Rb and Cs respectively and thus may explain the selectivity action of this new anticancer agent (PMF). It is clear from (Table II) that PMF contain also silver (Ag) and gold (Au) nanoparticles [49]. Noble metal, especially Au, nanoparticles have immense potential for cancer diagnosis and therapy [7].

IV. CONCLUSION

To conclude, we can state that PMF can induce apoptosis to A549 cells. The dominant protein secondary structure shifts from α-helix structure to unordered, indicating an altered protein profile in the apoptotic cells. In addition the total cellular lipid content increases starting from the first 5 to 15 min up to 24 hr of treatment while the amount of DNA decreases dramatically during treatment with PMF as observed in difference spectrum.

It should mention here that condensation of chromatins at the early stage of apoptosis was observed after 5 to 15 min of PMF treatment in the FTIR spectra indicated by the disappearance of the DNA absorbance bands. On the other hand, fragmentation of chromatins after 30 min of PMF treatment was evident by both FTIR spectroscopy and formation of DNA Ladder in the electrophoresis analysis. This study indicates that PMF contains Cs and Rb nanoparticles which proved to have a role in attacking cancer cells by elevating the pH. Thus, PMF as a natural product gives promising hope in cancer therapy with its prosperity of different potentially variable nanoparticles. These nanoparticles nowadays are all represent the new trend in treating cancer.

REFERENCES


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