Assessment of Cytogenetic Instability and Gene Transcription of Chronic Myelogenous Leukemia Cells Exposed to Non-thermal Plasma

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Abstract

Background: This study investigates the effect of Atmospheric Pressure Plasma Jet (APPJ) on cytogenetic instability and transcription of genes associated with propagation of chronic myelogenous leukemia (CML).

Material and methods: Blood samples were collected from six patients with CML and compared with control group. The first three cases were in accelerated phase (AP-CML) resistant to imatinib and nilotinib drugs. The others partially responded and returned to chronic phase (CP-CML). Triple blood cultures for each case were exposed to single dose of plasma, with treatment times of 40, 80, and 120 sec. Mitotic phases, indices and abnormalities, as well transcription levels of P53, caspase-3 and Bcl-2 genes were detected before and after plasma jet exposure.

Results and conclusion: The data showed that the exposure to APPJ caused remission of the mitotic indices and give a significant increase in the arresting of chromosome instabilities by induction of different mitotic checkpoints and express of sticky and condensed chromosomes which ended by apoptosis and necrosis processes. Additionally, the same events detected for the transcriptional levels of P53, caspase-3 and Bcl-2 genes.

Keywords: Plasma jet; Mitotic abnormalities; P53; Caspases-3; B-cl2; CML

Introduction

Next to solid, liquid and gaseous state, physical plasma is referred to as the fourth state of matter. Plasma is generated by the input of sufficient energy (beams, thermal or electric field energy) to the gas that atoms collide with each other and knock their electrons off. It is defined as an ionized gas with quasi-neutral characteristics [1]. Plasma has a complex composition. It consists of ions, electrons, exited and neutral atoms, free radicals (ROS and RNS), ultraviolet (UV), thermal and infrared radiation, electric fields and molecules [2]. Therapeutic application of plasma at or in the human body is a challenge both for medicine and plasma physics [3].

Non-thermal atmospheric pressure plasma has drawn more and more attention worldwide in the biomedical sector over the last two decades. Plasma effects on mammalian cells are of basic interest in vitro tests to characterize plasma-cell interactions should include the basic cellular parameters like morphology, viability or proliferation and the cellular responses like influence on DNA or cellular proteins [4]. Moreover, the plasma acted at the cellular level to remove diseased tissue without inflammation and damage, to suppress infectious and to modulate the viability (apoptosis/necrosis) of tumoral cells [5].

CML is a prototypical stem cell malignancy with a natural course of progression from an initial CP to AP and blast crisis (BC). Transformation to BC is associated with acquisition of additional chromosomal aberrations beyond the underlying t (9:22) chromosomal translocation that characterizes CML [6].

Nilotinib is pharmacological related to imatinib mesylate and dasatinib, these drugs are inhibitors of Bcr-Abl tyrosine kinase. Imatinib resistance can be defined as lack of complete hematologic response in patients with CP-CML, or as a failure to return to CP for patients with CML in AP or BC. Drug resistance is associated with reactivation of Bcr-Abl signal transduction [7].

Chromosomal aberrations can occur spontaneously or after exposure to genotoxic exposures and play an important role in cancer pathogenesis [8]. Acquisition of chromosomal aberrations is a major feature of disease progression in myeloproliferative and myelodysplastic disorders and acute myeloid leukemia [6]. Ectopic repair between DNA sequences in different chromosomes may result in translocations and dicentric chromosomes [9].

As well, for the last 30 years, chromosome aberration assay and detection of unstable aberrations, dicentrics and acentric fragments, have been used for the estimation of genome damage caused by physical and chemical clastogens [10].

Dicentric chromosomes have been identified as instigators of the genome instability associated with cancer, but this instability is often resolved by one of a number of different secondary events. These include centromere inactivation, inversion, and intercentromeric deletion. Deletion or excision of one of the centromers may be a significant occurrence in myeloid malignancy and other malignancies but has not previously been widely recognized. The failure to identify centromere deletion in cancer cells until recently can be partly explained by the standard approaches to routine diagnostic cancer genome analysis, which do not identify centromeres in the context of chromosome organization [11].

Mitotic index is defined as the ratio between the numbers of cells in a population undergoing mitosis to the number of cells not undergoing

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mitosis. The purpose of the mitotic index is to measure cellular proliferation. The mitotic index is an important prognostic factor predicting both overall survival and response to chemotherapy in most types of cancer [12].

The cell cycle control system is arrested when cells detect DNA damage. Meanwhile, the premature onset of mitosis leads to another type of cell death that is mitotic catastrophe [13]. Mitotic catastrophe occurs either during or after mitosis [14]. A new type of cell death that takes place at metaphase is defined as metaphase fragmentation where condensed chromosomes lose viability and are progressively degraded [15]. Chromosome fragmentation does not exhibit the typical oligosomal DNA degradation of apoptosis [16]. Cell death is the consequence of inappropriate cell development with different paths. So, it was proposed that hyperplasia could result from decreased apoptosis rather than increased mitosis [17].

The P53 tumor suppressor limits cellular proliferation by inducing cell cycle arrest and apoptosis in response to cellular stresses such as DNA damage, hypoxia, and oncogene activation, many apoptosis-related genes that are transcriptionally regulated by P53 have been identified. In mammals, apoptosis can be initiated by three different pathways: (1) the extrinsic pathways, which can be triggered by ligation of death receptors and subsequent Caspase-8 activation; (2) the intrinsic pathway, which is initiated by cellular stress followed by activation of caspase-9; or (3) the granzyme B pathway, where the cytotoxic cell protease granzyme B pathway is delivered to sensitive target cells. Each of these pathways converges to a common execution phase of apoptosis that requires proteolytic activation of caspase-3 and/or -7 from their inactive zymogens [18].

Because of its fundamental importance, programmed cell death is a highly regulated pathway. One important set of proteins involved in the regulation of apoptosis in the Bcl-2 family. Over 25 family members have been identified [19]. These can be broadly divided into two classes: those that inhibit apoptosis and those that promote apoptosis.

Due to their importance in keeping cancer cells alive, Bcl-XL and Bcl-2 are considered relevant targets for cancer chemotherapy. In fact, Bcl-2 anti-sense nucleotides are currently being tested in clinical trials for treatment of cancer [20].

Our study aimed to investigate the potency of APPJ exposure to perturb genetic and cell cycle progression leading to a programmed death of the CML cells which resistant or partially responded to chemotherapy.

Materials and Methods

Character of exposure machine

Non-thermal plasmas are frequently called “on-equilibrium” plasmas because they are characterized by a large difference in the temperature of the electrons relative to the ions and neutrals. Since the electrons are characterized by a large difference in the temperature of the electrons relative to the ions and neutrals. cold plasma devices have recently attracted significant attention [21]. The most important devices for generating atmospheric pressure non-thermal plasmas can be considered: APPJ [22], plasma needle [23], plasma pencil [24], miniature pulsed glow-discharge torch [25], one atmosphere uniform glow-discharge plasma [26], resistive barrier discharge [27] and dielectric barrier discharge [28].

A schematic of the pulsed APPJ discharge and of the experimental setup is shown in Figure 1. The gas is fed through an annular region between the two metal electrodes 15 cm in length. The inner electrode is 5 mm in diameter and is powered with pulsed high voltage power supply, while the grounded outer electrode is separated from the inner electrode by a gap of 2-3 mm. A rotating spark gap is made by rotating the ground electrode on a rotating plane.

The APPJ device operates using 10-20 kV power supply with a gap between two electrodes is 2-3 mm under atmospheric pressure [22]. The electrical discharge inside the reactor of APPJ was induced by a pulsed high voltage power supply. The discharge power was adjusted by the pulse voltage with a pulse frequency of 167 kHz.

Chemicals

The chemicals of the blood culture were purchased from Gibco-BRL, USA. Heat-inactivated fetal calf serum (FCS) from Sigma/Aldrich chemical Co, St Louis, USA.

Blood sampling

Blood samples obtained from six volunteers of CML patients, the first three of them were in AP-CML, and they were imatinib resistant (TR-case), and the others were in CP-CML, and they were partially responding to the imatinib treatment (T-case). The diagnosis was based on clinical examination and laboratory evaluation, which were carried out by the consultant medical staff.

Further, healthy subjects matched with patient groups for ethnic background, sex, smoking and age were also investigated as control group.

All the cases were clinically diagnosed with a medium duration 1-2 years at the accelerated phase. We chose three cases were responded to nilotinib and returned to the CP through one year but the others three cases were not responding through the same period of treatment and still persisted in AP. One patient from this group evolved to death.

The mean value of WBCs count for the T-group at the beginning of treatment was 80,000 cells/mm3 and within 8 to 12 months through the nilotinib period reached to 8,000 cells/mm3. In addition the Bcr-abl was decreased through the same period of the treatment with ratio (2.03:0.27) prior to sampling, whereas the TR-group recorded mean value of total leukocytes count at the beginning of the treatment was 118,000 cells/mm3 and within one year of nilotinib treatment, the mean value was 90,000 cells/mm3 (Bcr-abl ratio was 3.84: 2.03) accordingly they received interferon therapy at last three months prior to sample collection.

All subjects were gave an informed consent for participation in this study. The donors were selected according to current International Programme on Chemical Safety (IPCS) guidelines for the monitoring of genotoxic effects of carcinogens in human [29].

Venous blood were collected under sterile conditions in heparinized
vacationer tube \((v = 5 \text{ ml})\), Becton Dickinson, USA) containing Lithium heparin as anticoagulant.

**Experimental Design**

The blood from each volunteer was divided into four groups for each case (TR and T cases). One was non-exposed and the other three were applied directly to APP at a distance of 3 cm from the blood surface. In each group, three time period groups were selected for the study (40, 80 and 120 sec.). The first four groups of TR-case were as a following: TR, TR-D1 (duration for 40 sec.), TR-D2 (duration for 80 sec.) and TR-D3 (duration for 120 sec.) groups. The other four groups of T-case were as a following: T, T-D1, T-D2, and T-D3 groups. All previous groups were matched with healthy subjects group, which represent the negative control (C-group).

The blood incubated at 37°C before initiation of cultures then cytogenetic analysis and gene transcriptions done for each group before and after exposure to APP.

**Blood culture**

Triple blood cultures were set up for 48 hours for each sample according to the protocol described by Evans and O’Riodran [30].

**Chromosomal aberration test**

The cells were cultured for 48 hours, blocked in metaphase, fixed, stained with Giemsa, and scored; at least 50 cells per slide were examined under 100X magnification for mitotic analysis. The number of total cells in the mitotic division was scored and the percentage of cell division was calculated (MI), percentage of mitotic phases, percentage and type of abnormalities in each mitotic phases. The most common abnormalities were pictured by microphotography. The cytogenetic analysis technique was done according to protocol recommended by [16,31].

**P53 and Bcl2 Genes transcription and caspase-3 levels**

Analysis of gene transcriptions carried out using by ELISA kit according to the following kits:

1. P53 tumor protein (P53/Tp53), purchased by CUSABIO Company, Cat No.: CSB-E08336r.

**Statistical analysis**

Data was presented as distribution analysis, percentages, means ± SE and analyzed using two ways analysis of variance "F" test according to Abramowitz and Stegum [32], the level for statistical significance was p<0.05.

**Results**

Table 1 showed that the mitotic index in TR group had a significant increase as 5.4 folds when compared with that of C-group. Whereas, exposed groups (TR-D1, TR-D2 and TR-D3) mitotic indices showed significant decreases when compared with that of the non-exposed group (TR). TR-D2 group scored the lowest value in mitotic index when compared with the other exposure doses. As well, mitotic index in TR-D3 group showed a significant decrease when compared with that of TR-D1.

On the other hand, a significant decrease in number of cells entered prophase in CML patients by 2.2 folds than that of C-group. However, exposed groups showed no significant differences in the frequency of prophase when compared with that of non-exposed group (Figure 2B).

A significant decrease in metaphase frequencies (normal mitotic pattern) was observed in TR-D2 and TR-D3 groups when compared to that of the other groups. The lowest number of metaphases was detected in TR-D3 group (Figure 2A).

TR-D1 and TR-D2 were recorded significant decrease in anaphase and telophase cells when compared with that of TR-group and C-group. In addition, a significant difference was scored in TR-D3 in comparing with that of TR-group.

As regard to mitotic abnormalities, induction of sticky cells was significantly increased in exposed groups when compared with TR- and C-group. The values of the frequencies of the sticky metaphases were scored with gradually increments from TR-D1 to TR-D3 (Figures 2C-2E). Moreover, condensed chromosome metaphases were detected with low frequency in the control group comparing with the other estimated groups (Figures 2D-2I). Whereas, the exposed groups (TR-D1, TR-D2 and TR-D3) showed nearly or more than two folds increments in the condensed chromosome metaphases than that of non-exposed group (TR-group).

Figure 3 illustrated that the total number of metaphases in the normal and abnormal patterns were more arrested in the exposed groups than that of control and TR groups.

Table 2 showed that the T-group had a significant increase in mitotic index when compared with the C-group and no significant decrease was observed when compared with all exposed groups (T-D2 group had the lowest value of mitotic index).

The statistical differences between the means of the percentages of

<table>
<thead>
<tr>
<th>Group</th>
<th>Mitotic index</th>
<th>Mitotic Phase</th>
<th>Mitotic Abnormalities</th>
<th>Condensed Chromosome Metaphases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Prophase</td>
<td>Metaphase (Normal Pattern)</td>
<td>Anaphase&amp; Telophase</td>
</tr>
<tr>
<td>C-group</td>
<td>8.5 ± 0.76</td>
<td>17.33 ± 0.5</td>
<td>63.33 ± 1.93</td>
<td>17.0 ± 2.22</td>
</tr>
<tr>
<td>TR-group</td>
<td>45.60 ± 2.28a</td>
<td>7.75 ± 1.14a</td>
<td>65.81 ± 2.47</td>
<td>18.36 ± 1.32</td>
</tr>
<tr>
<td>TR-D1 group</td>
<td>34.13 ± 1.14ab</td>
<td>6.33 ± 0.67a</td>
<td>61.0 ± 1.00</td>
<td>10.11 ± 0.68ab</td>
</tr>
<tr>
<td>TR-D2 group</td>
<td>19.67 ± 0.88abc</td>
<td>8.30 ± 1.43a</td>
<td>53.0 ± 1.53abc</td>
<td>9.85 ± 1.17abc</td>
</tr>
<tr>
<td>TR-D3 group</td>
<td>25.97 ± 1.11abc</td>
<td>7.04 ± 0.58a</td>
<td>40.26 ± 2.67abc</td>
<td>11.37 ± 1.89a</td>
</tr>
</tbody>
</table>

*P < 0.05
a. P-value Significant when compared with C-group.
b. P-value Significant when compared with TR group.
c. P-value Significant when compared with TR-D1 group.
d. P-value Significant when compared with TR-D2 group.

Table 1: Mitotic index, phase and abnormalities in exposed and non-exposed TR-and control groups (Means ± SE)
prophase of all groups were not significant except that of T-D1 group when compared with C and T-groups. Metaphases data recorded significant decreases in T-D1, T-D2 and T-D3 groups when compared with C and T-groups. T-D2 group evidenced the lowest rate in metaphase frequencies. In addition, anaphase and telophase frequencies stated no significant differences between T-D1 and T-D2 groups and C-group. Contrary, significant differences were observed between T-D1, T-D2 and T-D3 groups and T-group.

The means of percentages of mitotic abnormalities were indicated in T-group as the following: significant increase of sticky and condensed chromosome metaphases in exposed and non-exposed groups when compared with C-group. In addition, there are significant increases of sticky cells in T-D1, T-D2 and T-D3 groups more than T-group; particularly T-D2 group had the highest frequency of sticky cells. On the other hand, for condensed chromosome metaphases the exposed groups were significantly less than T-group except T-D3 group, which recorded non-significant value higher than T-group.

The total number of normal and abnormal patterns of the metaphases in the non-exposed and exposed groups was more than that of the control group. Whereas, T-D2 group scored the lowest values in the two metaphases patterns (Figure 4).

As regard to P53 and Bcl-2 genes transcriptional levels, Table 3 showed that TR-group had high significant incidences when compared with control group. P53 level scored significant increase in the exposed groups when compared with TR group. The remarkable increment was observed in TR-D2 group more than the other exposed groups. In the meantime, the transcriptional levels of Bcl-2 genes were significantly decreased in exposed groups when compared with TR-group. However, TR-D3 group revealed higher level in this gene expression comparing with that of control, TR-D1 and TR-D2 groups.

As for caspase-3, its levels were detected with lower values in exposed and non-exposed groups comparing with that of the control group. Whereas, its values were increased in TR-D1 and TR-D2 groups more than that of TR-group. While, TR-D3 was recorded with slightly decreased value comparing with TR-group.

Table 4 showed that the non-exposed T-group and exposed groups showed significant increase in level of p35 when compared with that of control group. Moreover, P53 levels were remarkably increased in TD-2 and T-D3 when compared with that of non-exposed group. The data of caspase-3 was showed non-significant increase in T-D1 and T-D2 when compared with T-group but T-D3 insignificantly decreased. Moreover, the caspase-3 levels in the exposed groups were non-significantly different when compared with control group.
Table 4: Transcriptional levels of P53, Caspase-3 and BCL-2 genes in exposed and non-exposed T- and control group

<table>
<thead>
<tr>
<th>Group</th>
<th>Mitotic index</th>
<th>Mitotic Phase (Normal Pattern)</th>
<th>Anaphase &amp; Telophase</th>
<th>Sticky Chromosome Metaphases</th>
<th>Condensed Chromosome Metaphases</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-group</td>
<td>8.5 ± 0.76</td>
<td>17.33 ± 0.51</td>
<td>63.33 ± 1.93</td>
<td>17.0 ± 2.22</td>
<td>0.56 ± 0.56</td>
</tr>
<tr>
<td>T-group</td>
<td>15.67 ± 1.20</td>
<td>16.0 ± 1.53</td>
<td>56.67 ± 1.45</td>
<td>5.67 ± 0.33</td>
<td>4.33 ± 0.33</td>
</tr>
<tr>
<td>T-D1 group</td>
<td>13.33 ± 1.20</td>
<td>10.0 ± 0.58</td>
<td>49.67 ± 0.33</td>
<td>10.11 ± 0.68</td>
<td>15.33 ± 0.88</td>
</tr>
<tr>
<td>T-D2 group</td>
<td>9.33 ± 0.33</td>
<td>16.0 ± 0.58</td>
<td>41.0 ± 0.58</td>
<td>18.0 ± 1.16</td>
<td>18.0 ± 0.58</td>
</tr>
<tr>
<td>T-D3 group</td>
<td>10.0 ± 0.58</td>
<td>13.33 ± 0.88</td>
<td>44.0 ± 0.58</td>
<td>8.0 ± 0.58</td>
<td>15.33 ± 0.33</td>
</tr>
</tbody>
</table>

* P < 0.05
B: P-value Significant when compared with T-group.
C: P-value Significant when compared with T-D1 group.
D: P-value Significant when compared with T-D2 group.

For T-D2 group, there is notable decrement of Bcl-2 gene transcription level when compared with T-group rather than the other exposed groups, which recorded non-significant decrement.

Discussion

Our somatic cells are born by mitosis and almost all will die by apoptosis, a physiological process of cellular suicide. Cancers can occur when this balance is disturbed, either by an increase in cell proliferation or a decrease in cell death. The goal of cancer therapy is to promote the death of cancer cells without causing too much damage to normal cells [33].

Imatinib mesylate (Gleevec, Novartis Pharma, New Jersey, USA) is a first-generation tyrosine kinase inhibitor (TKI) that was approved for frontline therapy in patients with (CML) by the US Food and Drug Administration (FDA) in 2002. It is dosed at 400 mg daily in patients with CML in the CP-CML and 600 mg daily in patients with -AP-CML and for those in the blast phase (CML-BP). Imatinib mesylate is a BCR-ABL-targeted therapy and considered the standard of care in CML management [34].

Despite the positive results obtained in previous studies, approximately 33% of patients with CML treated with imatinib do not achieve a complete cytogenetic response, while others have drug resistance or cannot tolerate drug-related toxicities [35].

On the other hand, BCR-ABL kinase domain mutations may confer more specific resistance to nilotinib which will predominantly affect response rates, the presence of additional chromosomal aberrations may reflect genetic instability and, therefore, intrinsic aggressiveness of the disease which will be less amenable to subsequent alternative treatments and thus negatively affect overall survival [36].

Sensenig et al. [37] stated that apoptosis, or programmed cell death is a critical element of cellular self-regulation, which is a vital function in multi-cellular organisms allowing for appropriate growth, development, and death at the necessary times. The non-functioning of a tumor-suppressor gene or the over expression of an anti-apoptotic protein are both important pathways in cancer development.

Plasma is an ionized gas composed of charged particles (electrons, ions), electronically excited atoms and molecules, radicals, and UV photons. Plasma treatment exposes cells or tissue surface to active short-
and long-lived neutral atoms and molecules, including ozone (O$_3$, NO, OH radicals), and singlet oxygen (O$_2$), and a significant flux of charged particles, including both electrons and positive and negative ions like super oxide radicals [38].

Mitosis process follows five stages and is terminated by cytokinesis. The cells should pass through two checkpoints before entering into mitosis. First is the G2 checkpoint in response to DNA damage, [39], followed by the antephase checkpoint that is activated to prevent mitotic defects in response to a number of stresses, changes in chromatin structure and spindle damage (Figures 2F and 2G) before the cells reach the point of 'no-return' [40]. The antephase checkpoint which was first introduced by Bullough and Johnson [41] exists between the G2 and early prophase [42].

Our findings showed that the treatment resistant group (TR) exhibited a significant increase rate of mitotic index more than control group (5.4 folds). After APPJ exposure with the three doses (TR-D1, TR-D2 and TR-D3), the mitotic indices decreased significantly remarkably with TR-D2 (Table 1). At the same current, the treatment responded group (T) was significantly increased in the rate of mitotic index more than that of control group (1.8 folds). However, there is non-significant decrease of the mitotic indices rates between T-group before and after plasma exposure. T-D2 group scored the lowest value of mitotic index comparing with the other exposure doses (Table 2).

This depression in mitotic indices which occurred after exposure to APPJ may be attributed to the check points which enforce the correct sequence of events after DNA or spindle damage this mechanisms arrest cell cycle progression in response to damage. DNA damage sent negative signals to a series of protein kinases, which arrest the cycle at DNA damage checkpoint [43].

Light microscopy cells were well preserved at all time intervals studied. General observations on nuclear morphology and mitotic phases are summarized in Tables 1 and 2. All mitotic phases were represented in dividing cells, providing a basis for comparison with treated samples. Mitotic phase required chromosome condensation and a visible nuclear membrane. Metaphase required near linear alignment of chromatids on one or more equatorial planes and no visible nuclear membrane. Anaphase was scored when chromatids were condensed, were segregated into two or more groups, and were not arranged in linear arrays. All anaphase nuclei lacking this feature were scored as abnormal. Telophases required complete segregation of chromatid clusters, early dispersion of chromatin material, and evidence of nuclear membrane reformation with or without evidence of cytokinesis [44].

In the present study, prophase scored in control group were significantly increased more than that of both non-exposed and exposed groups, but the frequencies of metaphases, anaphases and telophases in the TR-group were non-significantly increased than that of C-group.

Metaphases with normal pattern recorded in TR-D1, TR-D2 and TR-D3 groups were decreased than that of TR-group. On the other hand, the total frequencies of the metaphases with normal and abnormal patterns were lower than that of TR-group (Figures 2 and 3). After APPJ exposure, Anaphase & telophase were expressed lower frequencies in exposed groups than that of non-exposed and control groups.

The accumulation of normal and abnormal patterns of metaphases after plasma exposure may be discussed in view of the inability of the cells to perform metaphase checkpoint, by stabilizing activity of mitotic cyclin dependent kinases and preventing cells to exit from mitosis [45]. Thus, the presented data confirm that both entry into and exit from mitosis is blocked in treated cells suggesting that plasma exposure may interfere with the balance between cyclin condensation required for participation into mitosis and its ubiquitous destruction by anaphase promoting complex [46].

Another interpretation presented by Shah and Cleveland [47] in their study were most microtubule- damaging agents that inhibit normal spindle formation, either by increasing microtubule stability or by depolymerization, conduces cells to arrest at metaphase-to-anaphase transition by the action of the spindle-attachment checkpoints.

In contrary, the presented data (Table 2 and Figures 2 and 4) for the treated cases which respond to the chemotherapy but with severe depletion in the total leukocytic counts. The exposure to the APPJ improves the mitotic index and the ratio between the different phases. Therefore, there are not significant differences between exposed and non-exposed groups and control group for most percentages of the phase's data.

The most effective role of the exposure of CML cells to the plasma through its division is the expression of abnormal pattern of metaphases (sticky and condensed chromosome metaphases), which take its paths to apoptosis and necrosis. Our findings as presented in Table 1. pointed to highly significant increment in the number of metaphases with sticky chromosomes in the exposed groups (TR-D1, TR-D2 and TR-D3 groups) comparing with non-exposed (TR- group) and control groups. Moreover, the same event occurred for the appearance of condensed chromosome metaphases after APPJ exposure.

Table 1 showed that there are highly increment for sticky and condensed chromosome metaphases in the exposed group (T-D1, T-D2 and T-D3 groups) comparing with non exposed and control groups but this increment were less than their data that investigated for the exposed TR-group.

Sticky chromosomes or metaphases and condensed chromosomes which be detected in this study, after APPJ exposure, considered the main observed abnormalities due to cell cycle and spindle check points. Babich et al. [48] reported that metaphases with sticky chromosomes lose their normal appearance and appear to have a sticky "surface" which causes chromosome agglomeration, possibly due to effects on chromatin and chromosome organization.

Moreover, Erenpreisa et al. [49] stated that the increase of mitotic index in response to heat shock represented mostly an arrest in mitosis. The morphology and the increased ratio between metaphases and anaphases metaphase telophases evidenced that this was arrest in metaphase, thus at a spindle checkpoint. A large portion of this delay looked like pyknotic metaphases, which shared morphological features with apoptosis. They contained sticky chromosomes, which tended to clump, a property dependent on DNA strand breaks.

The cytogenetic findings in our work interpreted and concomitant with data of the apoptic gene transcription and caspases activation, which detected in CML cells before and after APPJ exposure. Tables 3 and 4 illustrated that the increments of transcriptional level of P53 in TR- and T-groups comparing with control group were inadequate for controlling the mitotic abnormalities and apoptic expression. After exposure to plasma, the data showed notable increases in P53 expression especially in the exposed TR-group and chiefly for 80 and 120 sec. periods.

Banin et al. [50] stated that plasma-jet treatment increased the level of phospho-p53. The tumor suppressor protein p53 plays a major role in the cellular response to DNA damage and other genomic aberrations. In response to cellular stresses including DNA damage, phosphorylation-mediated activation of p53 plays important roles in cell survival. Activation of p53 can lead to cell cycle checkpoint arrest and apoptosis by activating transcription of many downstream target genes.
The levels of caspases-3 in T and TR-groups were lower than that of the control group other than; there are gradually increasing in these levels after APPJ exposure, except for TR-D3 and T-D3 groups.

Caspases have been divided into two groups: initiators and effectors, first group of initiator (or apical) caspases (caspase-2, -8, -9, -10, and probably, -11). The effectors (or downstream) caspases are able to directly degrade multiple substrates including the structural and regulatory proteins in the cell nucleus, cytoplasm, and cytoskeleton [51]. Caspase-3, -6, and -7 are three effectors caspases highly homologous to each other [52]. Their final functions are also similar or in executing apoptosis. In apoptotic cells, caspase-3 is the main executioner as it can be activated through both extrinsic and intrinsic signaling pathway, but it cannot be cleaved caspase-2 [53]. So, activation of the effector caspase-3 occurs as a last step of the apoptotic caspases cascade and is therefore considered as a late apoptotic signal. Plasma treatment induced activation of caspase-3 in response to DNA damage [54].

The transcriptional levels of Bcl-2 in TR- and T-groups were significantly and non-significantly higher than that of the control group respectively. But, there are gradually decreasing in these levels after APPJ exposure, except for TR-D3 and T-D3 groups.

Bcl-2 has not only been localized to the outer mitochondrial membrane but also to the nuclear envelope and the endoplasmic reticulum membrane (ER). In the ER, it regulates calcium storage, whose intracellular levels have been shown to affect apoptosis. ER-associated Bcl-2 is able to protect from apoptosis induced by various triggers. Beyond Bcl-2, Bcl-XL also interacts with pro-apoptotic members like BAX and BAK thought their BH3 domains [55]. It is possible that the anti-apoptotic action of Bcl-2 and Bcl-XL is converted to a pro-apoptotic one when these proteins are cleaved by caspases after initiation of apoptosis [56].

The mechanism of action of cold plasma on cancer cells is related to generation of reactive oxygen species with possible induction of apoptosis pathway, and the cancer cells are more susceptible to the effects of cold atmospheric plasma because a greater percentage of cells are in S-phase of the cell cycle [57]. When leukemia cells were treated with the cold plasma plume, the scientists found something unexpected: The cells did not die right away. Rather, right after experiment in the zero th hour, the cells are still okay. But after four to eight hours, they start dying “Said Laroussi” [58].

In summary, we investigated the feasibility of APPJ treatment for CML therapy and examined the mechanism by which plasma induces anti-proliferative properties and apoptosis. This effect occurred through not only the decreasing of mitotic index and increased metaphases arrest, but also through the increment of p53 transcription levels, activated caspase-3 and downstream of Bcl-2.

The presented study is the third part of our work on CML investigation and treatment by APPJ exposure. In addition, the period time of 80 sec. was the most effective and save dose for attendant with the advice of Ahmed et al. [59-61].

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Declaration of interest

The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

References


