Evaluating Gamma-H2AX Expression as a Biomarker of DNA Damage after X-ray in Angiography Patients

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ABSTRACT

Objective: Coronary heart disease (CHD) is one of the most common diseases. Coronary angiography (CAG) is an important apparatus used to diagnose and treat this disease. Since angiography is performed through exposure to ionizing radiation, it can cause harmful effects induced by double-stranded breaks in DNA which is potentially life-threatening damage. The aim of the present study is to investigate phosphorylation of Histone H2AX in the location of double-stranded breaks in peripheral blood lymphocytes as an indication of biological effects of radiation on angiography.

Materials and Methods: This method is based on the phosphorylation measurement of Histone (gamma-H2AX or γ-H2AX) levels on serine 139 after the formation of DNA double-strand break. 5 cc of blood samples from 24 patients undergoing angiography were taken pre- and post-radiation. Blood lymphocytes were extracted, fixed and stained with specific γ-H2AX antibodies. Finally, the percentage of phosphorylation of Histone H2AX as an indicator of double-strand break was measured by a cytometry technique.

Results: An increase was observed in all patients’ percentage of phosphorylated Histone H2AX (double-stranded breaks DNA) after radiation (20.15 ± 14.18) compared to pre-exposure time (1.52 ± 0.34). Also, the mean of DNA double-strand break is shown in a linear correlation with DAP.

Discussion: Although induction of DNA double-strand breaks was associated with the radiation dose in patients, the effect of individual factors such as radio-sensitivity and regenerative capacity should not be ignored. In the future, if we are able to measure DNA damage response in every angiography patient, we will use it as a biomarker for the patient dose; this will promote public health.

Conclusion: Using flow cytometers readings done automatically is possible to detect γ-H2AX in the number of blood cells, therefore, the use of this technique could play a significant role in monitoring patients.

Keywords
X-ray, Angiography, Double-strand Break DNA, γ-H2AX, DNA Damage

Introduction

Coronary Heart Diseases (CHD) are considered to be among the most prevalent and lethal diseases of today’s world. Coronary angiography (CAG) has become an important apparatus in the diagnosis and treatment of CHD. The widespread use of CAG has raised concerns among physicians and patients about the harmful effects of radiation exposure [1]. The findings of this study will help better understand the risks associated with radiation exposure, so that clinicians
and patients can make informed decisions about this method, and also it helps determine whether additional strategies are needed to protect patients against radiation exposure [2]. Ionizing Radiation (IR) can cause damage which is usually related to the radiation dose [3]; this damage can, in turn, lead to DNA damage such as base damage, cross-links, Single-Strand Breaks (SSBs) and Double-Strand Breaks (DSBs) [4].

In comparison with SSBs which leave one strand intact to guide the repair, DSBs demand a more difficult repair process. With regard to ionizing radiation, SSB levels are 20-40 times higher than the levels of DSBs do [5], thus, DSBs signal activation usually leads to cell-death and permanent growth arrest [6].

Clinical signs related to DSB-inducing agents vary greatly among people [7]. Therefore, the same radiation dose could lead to severe complications in one patient. However, there is no trace of them in another patient [5].

Since phosphorylated Histone H2AX proteins (gamma-H2AX or γ-H2AX) depend on the approximate number of DSBs, it is believed that γ-H2AX is a marker [8, 9], thus, this marker is able to determine the number of DSBs by using a suitable antibody [10]. Therefore, if we were able to measure the DNA damage responses in each patient, we could use this signal as a guide for our radiation dose [5]. The aim of this paper is to study the phosphorylation of Histone H2AX at the site of DSBs in peripheral blood lymphocytes as a marker for the biological effects of radiation on patients undergoing angiography.

Materials and Methods

An experimental study was carried out in 24 patients. Blood samples were taken before and after 10 minutes angiography in order to evaluate DSBs based on H2AX phosphorylation. Our study exclusion criteria include; the history of Lymphoma or Leukaemia, radiotherapy or chemotherapy during the past six months and X-rays within the past two days.

24 patients were enrolled in this study; 12 men and 12 women (50-65 years old). All 24 patients were under treatment for acute coronary syndrome (ACS).

The patients’ DSB responses were monitored by determining the phosphorylated Histone H2AX in the lymphocytes through flow-cytometry technique. Blood samples were collected after taking a written informed consent and poured into separate tubes containing an anticoagulant substance (EDTA). Lymphocytes were separated and then the analysis was done with slight changes to the protocol. 3ml of blood was diluted in PBS with a 1:1 ratio and added to a sterile Falcon tube containing additional Lymphodex with a ratio of 2:1, then, centrifuged for 20 minutes at 3000 rpm.

The lymphocyte layer was then separated and after benign washed three times with PBS and centrifuged for five minutes at 2500 rpm. It was diluted in 2 ml PBS, then added to a 2 ml microtube and centrifuged for five minutes at 1200 rpm. Ultimately, the lymphocytes were diluted in 1ml PBS, and after a cell count to measure the percentage of phosphorylated Histone H2AX, the solution was ready to be used by a BD Cytofix/Cytoperm fixation/permeabilization kit.

A total of 1 × 10^6 cell/ml was taken from the sample containing lymphocytes and the rest of the test was performed based on Cytofix/ Cytoperm fixation/permeabilization kit (Cat number: 554714) instructions. The cells were diluted in 250 µl BD Cytofix/Cytoperm solution and incubated in the dark at 4°C for 20 minutes. Next, they were washed with 250 µl Buffer 1x BD perm/wash twice. Afterwards, the cells were diluted in a 500 µl BD perm/wash Buffer 1x solution, and 15 µl of antibody Alexa Fluor 488 mouse anti-H2AX (ps139) was added to 3 × 10^4 cell/ml. At that point, the solution was incubated in the dark at 4°C for 30 minutes and washed with 250 µl BD perm/wash Buffer 1x twice. Finally, the cells were diluted in PBS, and the flow cytometric analysis was done while it was using a flow cytomo-
It is worth mentioning that four micro-tubes were used in each test (unstained, isotype, before and after).

**Statistical Analysis**

Data were analyzed via SPSS Statistics 16.0. Continuous variables were expressed as means ± SD. The mean percentages of phosphorylated Histone H2AX before radiation were compared with after it by using Wilcoxon test. The effect of gender was evaluated with Mann-Whitney test. Spearman’s correlations (rho, 95% confidence intervals, p value) were used to evaluate associations among variables. In order to do regression analysis, Stepwise regression was the method of choice.

**Ethical Considerations**

All patients signed informed consent and the study was approved by the Ethics Committees in Shiraz University of Medical Sciences.

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**Figure 1 A:** Samples flow cytometry dot plot from two patients before Angiography

**Figure 1 B:** Samples flow cytometry dot plot from two patients after Angiography
Results

All variables can be seen as mean ± Std. Deviation (SD) in Table 1.

In order to evaluate the normal distribution of variables, Kolmogorov-Smirnov Test was used. Due to the lack of normality of variables, non-parametric tests were used.

A total of 45% of patients (11 of 24) had a 4.6% increase in phosphorylation. At least, there was one phosphorylation H2AX after exposure (Figures 2A and 2B).

Wilcoxon test showed that the effect of radiation on the percentage of phosphorylated Histone H2AX was effective, so that an increase was observed in the percentage of phosphorylated Histone H2AX after radiation (14.184 ± 20.151) compared to pre-exposure time (0.3 ± 1.522) even when to use very small doses 57mGy (534μGym2) and (p-value<0.001)(The median range(Interquartile) after radiation was 15.90 (18.80) and 1.53 (0.51) after exposure (Figure3).

Average changes in phosphorylation of any DNA damage marker were lower in patients exposed to <270 mSv of radiation 7.3% (4.62% to 10.34%) compared with exposed to > 270 mSv of radiation, 28.21% (12.37% to 54.05%).

In order to evaluate the effects of gender on the percentage of phosphorylated Histone H2AX after radiation in patients angiography, Mann-Whitney test results demonstrated no significant difference between the gender of the samples after radiation in the percentage of phosphorylated Histone H2AX (p-value=0.18).

At least changes were observed in men 21.87% (mean dose 760.86), while for the women it was 15.384% (mean dose 492.76 (Figures 3).

The relation among variables was examined by Spearman’s correlation test. Spearman’s correlation test was used to investigate the effects of different variables (the blood radiation dose, Age, DAP (dose area product) and time) on the percentage of phosphorylated Histone H2AX after radiation in angiography patients.

According to Table 2, a significant relation was observed among variables (dose, time and DAP) and the percentage of phosphorylated Histone H2AX after radiation (p<0.001). Accordingly, DNA damage was also associated with radiation dose (Figure 4). However, no significant relation was observed with regard to age variable

**Table 1: Clinical and angiography parameters of patients.**

| Clinical Parameter       | Mean  | Std. Deviation |
|-------------------------|-------|----------------|
| Age                     | 58.166(50-65) | 5.096          |
| Sex                     |       |                |
| Male                    | 12    |                |
| Female                  | 12    |                |

| Scan Parameters          | Mean       | Std. Deviation |
|--------------------------|------------|----------------|
| DAP*, μGy.m²              | 4300.525   | 4923.231       |
| blood radiation dose, mSv| 626.817    | 758.722        |
| Time, min                | 8.538      | 10.854         |

*DAP = dose area product

Patients tested (n=24)
this indicates an extreme collinearity between three variables of DAP, time and dose. These three variables have the same weight in determining the percentage of phosphorylated Histone H2AX after radiation. Thus, for a better understanding of the correlation between these variables, three regression models have been applied (Table 3).

At first, DAP was considered with sex, age, and percentage of phosphorylated Histone H2AX before radiation variables. In the second the model, the dose was evaluated with sex, age, and the percentage of phosphorylated Histone H2AX before radiation variables. In the third model, time was investigated with sex, age, and the percentage of phosphorylated Histone H2AX before radiation variables.

In the first and second models, where only DAP and Dose with the three variables of sex, age, and the percentage of phosphorylated Histone H2AX before radiation were available, only DAP and Dose were significant. Whereas in a third model that time with the three variables of sex, age, and percentage of

Figure 2: DNA Damage Detected After Angiography (A) Scatter plot graph of levels of phosphorylated H2AX, at before and after radiation from CAG. (B) Quantitative assessment of protein biomarkers of DNA damage in patients undergoing CAG.
phosphorylated Histone H2AX the before radiation was available, not only time but also sex became significant.

With regards to the average of the percentage of phosphorylated Histone H2AX after radiation in men (23.33±14.59) and women (16.96±13.62), which was different? We found that time variable was a better variable to determine the percentage of phosphorylated Histone H2AX after radiation in comparison with the variables of DAP and Dose. This makes sex variable significant.

Discussion

After the discovery of X-ray, its adverse effects on biological tissues were manifested [11]. Meanwhile, nowadays it is hard to imagine the medical sciences without the use of X-rays [12].

Following radiation, DNA damage occurs in cells and among damage resulted from ionizing radiation, DSB is the most significant one that can lead to cancer [8, 13]. Due to its predictability, DSB is considered as a useful measure in researches [5].

After inducing DSB, phosphorylation of a few thousand Histone H2AX molecules would be the first cellular responses [14]. DNA is phosphorylated in a matter of minutes, both caused by SSBs and DSBs [15].

In the repair process of damaged DNA, H2AX is a key factor that accumulates at the damage site; which is a starting point for other repair mechanisms [14]. After phosphorylation of H2AX, γ-H2AX can be seen by various techniques [16].

In this study, flow cytometry was used to measure changes in the percentage of phosphorylated Histone H2AX; then, we evaluated the association between changes and the ionizing radiation used in our patients.

Our results revealed a significant increase in the percentage of phosphorylated Histone H2AX among patients following X-ray exposure.

Using flow cytometry, Macphaila et al. observed the formed γ-H2AX an hour after 20cGy of radiation. The maximum level of
γ-H2AX formation was observed between 15-30 minutes after being exposed to radiation, which was consistent with our results [17].

Kuefner et al. evaluated the induction and repair of DSB in peripheral blood lymphocytes among 19 patients (23-88 years of age), using an immunofluorescence microscopy. A number of DSBs (0.03-1.50 per cell) were observed, they observed increases even in small doses (at least 338 micro-Gy*m) [18].

In another study, Kuefner et al. evaluated DSB as a marker for the biological effects of radiation among 31 patients undergoing angiography in different body parts by using a fluorescence microscopy. After fluoroscopy, they obtained values of 0.01-1.50 DSBs/cell. There was a linear correlation between γ-H2AX foci levels and DAP (abdomen: R²=0.96; pelvis/legs: R²=0.71) [19].

As it can be seen in Table 2, results from Correlation-Spearman indicated that the induced DSBs reduced based on patients’ ages. Rothkamm and Lobrich measured DSBs in a population of cells using Comet assay; they observed a significant variation in γ-H2AX levels which did not correlate with the mea-

**Table 2: Correlation-Spearman.**

| variables | percentage of phosphorylated Histone H2AX after radiation | Dose | Time | DAP | Age |
|-----------|----------------------------------------------------------|------|------|-----|-----|
| percentage of phosphorylated Histone H2AX after radiation | 1 | - | - | - | - |
| Dose | 0.997** | 1 | - | - | - |
| Time | 0.970** | 0.975** | 1 | - | - |
| DAP | 0.996** | 0.998** | 0.969** | 1 | - |
| Age | -0.044 | -0.052 | -0.016 | -0.049 | 1 |

** p=<0.001

**Figure 4:** Scatterplot graph of the correlation analysis between DNA damage estimate and radiation dose (n = 24).
asured DNA levels and patient’s ages. This could have been caused by differences in the number of γ-H2AX formed at a given DNA damage level indicating differences in individuals related to efficiency by which cells convert DSBs to γ-H2AX foci [5].

Vandevoorde et al. examined the blood samples of 51 children undergoing thoracic and abdominal CT scans; they observed a drastic increase in DSBs between the range of 2-10 mGy doses, their linear regression analysis revealed that the number of induced DSBs per mGy had decreased with age [20]. These results are in line with our findings; however, they cannot explain the age dependency as to its relation with different radio sensitivities or different rates of DSB repair [21].

Since the rate of DSB repair varies among different patients, the rate of damage would vary as well.

Radiation damages are not strictly related to radiation dose and duration but are related to personal factors such as individual repair capacity, radio sensitivity and the use of contrast agents.

Iodinated contrast agents can intensify X-ray-induced DNA damages; this could be caused by decreased repair. Another theory is that these effects are caused by the increased photoelectric absorption in Iodine atoms, as well as continuous exposure to nearby lymphocytes [22].

Moreover, another study found a significant correlation between a number of DSBs and blood density levels in the heart and large vessels at the scan site [23].

Due to different blood volumes in different parts of the body, and different numbers of lymphocytes being exposed to radiation, cardiac catheterization causes more damages in comparison to procedures such as abdominal interventions and pelvic angiography, circulatory pathway and the cerebrum [18, 19].

Importantly, in this study we recruited patients who had a wide range of radiation dose to determine its effects on DNA damage. Overall, results are in accordance with our hypothesis that higher doses lead to more DNA damage.

Radiation disrupts DNA processes and causes DNA breaks [24]. Our study also revealed DNA damage in lymphocytes of patients undergoing CAG. γ-H2AX is proved to be a sensitive indicator for DSBs induced by ionizing radiation [25]. γ-H2AX shows a significant stability following IR and can be used as a radiosensitive marker [26]. There is a variety of methods in detecting DSBs such as Comet assay, neutral elution and pulse field electrophoresis (2-D gel electrophoresis); however, evaluation via γ-H2AX is a more accurate method in comparison with other methods [4]. This method sensitivity has turned it into a suitable technique for measuring DSB levels.

In this study, we used γ-H2AX to evaluate DNA damage caused during angiographic procedures. Although the use of an immunofluorescence microscopy is currently the most sensitive method for detecting γ-H2AX foci, it is quite difficult to use it in clinical research. On the other hand, flow cytometry can provide us with a simple detection of γ-H2AX in a large number of cells [27].

Numerous reports have indicated a convenient link between γ-H2AX detected through flow cytometry with the number of DNA breaks, cell deaths and radio sensitivity [17, 28, 29].

**Conclusion**

Despite the fact that CAG is a clinical tool
in the management of cardiovascular patients, physicians and patients’ awareness of DNA damage after CAG is required for dose reduction strategies.

Therefore, based on our study findings we recommend the use of flow cytometry to determine the percentage of phosphorylated Histone H2AX while monitoring patients undergoing angiographic procedures in order to promote health throughout the society.

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Conflict of Interest
The authors declare no conflict of interest.

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