Effect of Methotrexate on AgNOR Count in Liver of Wistar Rats

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ABSTRACT

A study was undertaken to evaluate the Nucleolar organizing regions (NOR) count and histopathological changes in rat's liver treated with methotrexate (MTX). Male Wistar rats aged 4 months, maintained in our institution were used in the present study. Animals were injected intraperitoneally with 3 different doses of methotrexate on alternative days for one week and the animals were sacrificed on the seventh day by using ether. Paraffin slides were prepared and histopathology as well as AgNOR studies were done. In the histopathological study, liver showed disruption of normal hepatic plates, vaculation of the hepatocytes, nuclear variability and enlarged blood vessels. In AgNOR study, data showed an overall decrease in the total count (numerical) from normal to 12 mg dose MTX. The total count includes small, medium and large sized AgNORs. A statistically high significant difference was found from normal to 12 mg dose MTX as well as between the doses. In conclusion, the genotoxicity in rats is well documented with high dose of methotrexate (12 mg/kg) compared to low dose (8 mg/kg). Methotrexate treatment reduces the transcription rate and ploidy. Methotrexate reduces the cell proliferation. Thus it reduces the count.

KEYWORDS: comma separated keywords

MATERIALS AND METHODS

Study design. The rats were divided in to four groups. Each group contains 6 animals. The animals were injected intraperitoneally with three different doses (8 mg, 10 mg and 12 mg) of methotrexate in alternative days for one week and the animals were sacrificed on the seventh day by using ether.

Sample size. Male Wistar rats aged 4 months, maintained in our institution were used in the present study. Animals were maintained under controlled conditions of light, temperature and humidity in an air conditioned animal house. The average weights of the animal were 200gm.

Tissue processing. The animals were deeply anesthetized with ether and fixed in a dissection board and its chest cavity is opened to expose the heart. About 15 ml of 0.9% saline was perfused through left ventricle at the rate of 1ml / minute. This is followed by perfusion of 10% formalin about 250 ml for adult rats at the same flow rate. The organs were removed and kept in 10% formalin for 48 hrs (post fixation). Paraffin blocks were made in an embedding bath. Sections of 3-5 microns thickness were cut from the blocks using rotary microtome. Sections were mounted on air dried gelatinized slides.
The tissues were de-paraffinized in xylene and transferred to water through descending grade of alcohol (absolute, 90 %, 70 %, 50 % alcohol). Sections were washed in tap water and stained with Ehrlich’s Haematoxyline for 5 minutes and washed in tap water. Then stained with 1 % of eosin for 30 seconds and dehydrated with ascending grades of alcohol, cleared in xylene and mounted in Canada Balsam and observed under microscope for assessing the histological state of the tissue.

**AgNOR STUDY**

**Preparation of AgNO3 solution**

To prepare 50% of silver nitrate solution 5 gm of silver nitrate powder was dissolved in 10 ml of distilled water and filtered through Whatman filter paper 1 in dark. The solution was stored in dark at 4°C, and used within one week.

**Preparation of Gelatin**

1 gm of powdered gelatin was dissolved in 49 ml of distilled water and 1 ml of formic acid. To dissolve faster, mixture was kept in hot water bath at 60°C for at least 10 minutes. The solution was stored in dark at 4°C and used within one week.

**Silver Nitrate staining**

A rapid staining and de-staining method developed in our lab [7] was used throughout the study. 4 drops of silver nitrate solution and 2 drops of gelatin solution were added on to the slide. During the staining process precautions were taken to avoid over heating of the slides. Over stained slides were flooded with either 50% alcohol (absolute, 90 %, 70 %, 50 % alcohol)

**De-staining method**

Over stained slides were flooded with either 50% alcohol (absolute, 90 %, 70 %, 50 % alcohol) or hydrogen peroxide for one second or 25% Hydrogen peroxide for 15 seconds. The process could be monitored under the microscope, and did not alter cellular architecture, composition and even staining ability. The dissolved silver was washed from the sections under running water.

**Inclusion and exclusion criteria**

Standard protocols were followed for recording number and quantitation of size and shape of dots [8].

**AgNOR count**

Areas with minimal cell overlap and no artifact were demarcated for counting. Weak or dark stained slides were not evaluated. The s appeared as black dots within the orange colored nuclear background. Dots were defined as discrete homogeneous silver precipitates with well defined edges. Overlapped dots with well defined edges were counted as greater than one when these appeared on viewing through different focal planes. Dots lying in a group with indistinguishable boundaries were treated as one dot. Dots outside the nucleolus were not considered. Observations were completed within a week of staining due to fading of stain on prolonged storage. Counting was performed, using oil immersion at 400X.

**Size**

Size were classified in to three groups on the basis of their diameter (Small<=1 μm, Medium>1 and≤3μm and Large > 3 μm).

**Shape**

On the basis of the shape, s was classified in to regular (with round or oval well-defined margin) and irregular (with irregular serrated margin) dots. In each category size (maximum diameter) of all dots were documented.

**Tools used**

Tools were measured using ocular micrometer (calibrated with stage micrometer).

**Statistical methods used**

Data on mean count; size wise distribution of dots was tested using one way ANOVA. Since the test of homogeneity of variance showed high significance between different groups, data were reanalyzed using square root transformation. For each dose and distribution, mean count, standard deviation, f-ratio and p-value were computed.

**RESULTS**

The histopathological study of methotrexate as well as the dose wise expression of s in liver was considered in the present study. Data were collected and analyzed. Liver showed disruption of normal hepatic plates, vaculation of the hepatocytes, nuclear variability and enlarged blood vessels.

Data showed an overall decrease in the total AgNOR count (numerical) from normal to 12 mg dose MTX. The total count includes small, medium and large sized AgNORs. A statistically high significant difference was found from AgNORmal to 12 mg dose MTX as well as between the doses.

Large sized AgNORs (> 3μm) (Figes 1 and 5)- Data showed an overall numerical decrease in the large sized AgNORs from control to 12 mg dose MTX. A statistically significant difference was found between control and the different doses. But there was no statistical difference between 8 mg dose vs 10 and 12 mg doses of MTX as well as between 10 and 12 mg doses of MTX.

Medium sized AgNORs (> 1-3 μm) (Figes 1 and 5)- Medium sized AgORs showed a statistically high difference between control and 12 mg dose MTX. However there was a numerical decrease between control and 10 mg dose MTX and an increase thereafter.
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Fig 1. Size wise distribution of AgNORs in liver - Total count and size wise distribution of s in control and drug treated groups. Each value represents mean of ± S.D from 6 animals/group. Significant values are represented as, control vs 8 mg, 10 mg and 12 mg - <0.05 = a, <0.01 = b and <0.001 = c. Between 8 mg vs 10 mg and 12 mg - <0.05 = d and <0.001 = f. Between 10 mg and 12 mg - <0.01 = h and < 0.001= i. (One way ANOVA and Banferroni post test)

Fig 2. Shape wise distribution of AgNORs in liver - Shape wise distribution of s in control and drug treated groups. Each value represents mean of ± S.D from 6 animals/group. Significant values are represented as, control vs 8 mg, 10 mg and 12 mg - <0.05 = a and <0.001 = c. Between 8 mg vs 10 mg and 12 mg - <0.001 = f. Between 10 mg and 12 mg - < 0.001= i.

Fig 3. Photograph showing Liver of 12 mg MTX treated rat. H & E stain (20 x).

Fig 4 Photograph showing liver of control rat. H & E stain (40 x).

Fig 5. A – Regular shaped B – Large sized C – Small sized D – Medium sized AgNOR

Small sized AgNORs (< 1 µm) (Figes 1and 5) Irregular shaped AgNORs (Figes 2 and 6)

Small sized AgNORs showed a numerical increase from control to 10 mg dose MTX and there after a decrease between 10 and 12 mg doses of MTX. The data showed a highly statistically significant difference from control to the higher doses as well as between the doses.

Regular shaped AgNORs (Figes 2 and 6)

Data showed a numerical increase in the values from control to 12 mg dose MTX. And it showed a highly to give rise to malabsorption syndrome. It inhibits the enzyme dihydrofolate reductase (DHFR) which is required for DNA synthesis and cell division [9].

DISCUSSION

Methotrexate (MTX) is an antimetabolite widely used in cancer chemotherapy, which can cause intestinal mucosal injury. The antimitotic effect of MTX is known to give rise to malabsorption syndrome. It inhibits the enzyme dihydrofolate reductase (DHFR) which is required for DNA synthesis and cell division [9].
In the present study liver showed disruption of normal hepatic plates, vacuolation of the hepatocytes, nuclear variability and enlarged blood vessels (Fig 3). MTX is a novel xenobiogenic inducer of rat liver and intestinal sulfotransferase. Intestinal inductions were found to be much greater than those found in liver, and MTX is the first antifolate apoptosis-inducing drug to show induction of intestinal sulfotransferase [10]. Methotrexate therapy is associated with liver damage, both acute (notably after high dose) and more seriously in chronic cases (generally after long term administration) [11-13]. Hepatic fibrosis and cirrhosis may develop without signs of hepatotoxicity and have led to eventual death. Reversible evaluation of hepatic enzymes may occur in some patients [14]. Administration of methotrexate increased mortality in patients with primary biliary cirrhosis [15].

Agynorphic nuclear organizer regions were widely used in clinics, more than 500 papers on relevant field have been published in China, including the application in different kinds of malignant tumors, such as stomach cancer, lung cancer, liver cancer, colon cancer, and breast cancer. In differential diagnosis of malignant lymphoma and benign lymph node diseases, is more valuable [16]. Studies suggest potential usefulness of AgNOR parameters, such as mean count and area of distribution within the nucleus, in diagnosis and prognosis of various neoplasms [17-23], especially oral squamous cell carcinoma [18, 21, 24, 25]. Mean AgNOR counts can be useful in diagnosing histopathological features 18) and have been shown to correlate with clinical outcomes of various cancers [24, 26, 27, 28]. Chia et al analysed the AgNOR pattern in urinary cytology in samples from 70 patients and concluded that a combined qualitative and quantitative AgNOR analysis can be useful in the differential diagnosis of urinary cytology [19]. Jay et al emphasized that since mean AgNOR count of a lesion is an easily reproducible measure, it would serve as a more objective criterion than hematoxylin and eosin (H&E) staining-based assessment for diagnosing epithelial dysplasia [29]. Further, AgNORs are sound replicatory markers and H&E criteria are subjective, it is conceivable that the H&E-based 'gold standard' misclassifies some of these lesions.

In conclusion, the genotoxicity in rats is well documented with high dose of methotrexate (12 mg/kg) compared to low dose (8 mg/kg). Methotrexate treatment reduces the transcription rate and ploidy. It also reduces the cell proliferation. Thus it reduces the AgNOR count. Higher the dose lesser the AgNOR count and the irregularity of AgNORs.

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