Histological Effect of Cytarabine on Liver and Buccal Mucosa in Mice

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Aims: This study aimed to investigate the histopathological effect of cytarabine drug on liver and buccal mucosa in mice. Materials and Methods: Sixteen male albino mice were randomly assigned to two experimental groups and housed as eight animals/cage as a group. Control Group: was given daily I.P distilled water for 5 days. Cytarabine Group: was given daily Cytarabine at dose (100mg/kg I.P) for 5 days. At the end of experiment, all animals were sacrificed and liver in addition to the buccal mucosa were excised and placed in 10% buffered formalin solution for histological preparation and evaluation. Results: histopathological study of cytarabine treatment group showed multiple changes in liver like defused vacuolar swelling with abnormal hepatic cords patterns, congested sinusoids and multiple foci of apoptotic cells while in buccal mucosa sections revealed a severely shrinkaged and atrophied appearance of mucus-salivary glands, vacuolation of the stratified squamous epithelium and interstitial edema. Conclusions: this study concluded the direct cytoxicity of cytarabine to liver and buccal mucosa. So caution should be taken when administrating the drug to patient with liver or salivary glands dysfunction.

Key words: Cytarabine, Buccal mucosa, Mice, chemotherapy.

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INTRODUCTION
Cancer is considered as a serious worldwide health problem, being the second leading reason of global spontaneous mortalities after cardiovascular diseases. The reason behind developing rapidly may be mainly due to environmental carcinogens and unhealthy lifestyles (1). The mode of treatment depends on the type, site, and grade of cancer, the step of the disease and the overall health of the patient (2).

Chemotherapeutic agents are the compounds that used in cancer treatment and they are varying in structure and mechanism of action. Cytarabine or Cytosine arabinoside (Cytosar®) is a pyrimidine nucleoside-based chemotherapeutic agent. Cytarabine is called cytosine arabinose because it combines a cytosine base with an arabinose sugar (3). It is a cell-phase-specific chemotherapeutic drug, primarily act during the S phase when cells are undergoing DNA synthesis (4). It causes extensive chromosomal destruction through induction of chromatoid aberrations (5). Thus Rapidly dividing cells, which need DNA replication for mitosis, are the mostly affected (3).

This drug, used to treat acute myeloid leukemia (AML), acute lymphocytic leukemia (ALL), chronic myelogenous leukemia (CML), and non-Hodgkin's lymphoma (6). Also used as an antiviral agent against herpes simplex virus and human cytomegalovirus (7).

It is administered through continuous intravenous infusion or injection, intrathecally or subcutaneously then after three phosphorylation steps cytarabine is transformed to cytarabine triphosphate (8). The majority of the drug is metabolized in the kidney, liver and gastrointestinal tract into its inactive metabolite and large amount of the administered dose is eliminated by renal excretion within one day (9). It is Common side effects include vomiting, liver function disturbances, diarrhea, bone marrow suppression, oral mucosal ulceration, rash, and bleeding (10).

Aims of study: This study aims to investigate the side effect of cytarabine drug on liver and buccal mucosa in mice through the histopathological examination.

MATERIALS AND METHODS
The drugs injection was done at the pharmacology laboratory of Department of dental basic sciences at college of dentistry / university of Mosul / Iraq from 1/9/2019 to 20/1/2020.

Laboratory animals
Sixteen healthy adult male albino mice weighing (25-30) gm obtained from the Animal House of Experimental Research Unit, College of veterinary medicine, University of Mosul, Iraq. They housed in rodent plastic cage with
steel wire mesh covers at (22 ±2) °C, 12hr light/12 hr. dark cycle and received standard laboratory diet and water. Animals permitted to acclimate one week before experiment. This study was done in accordance with the guidelines of the institutional animal research ethics committee / college of dentistry/ University of Mosul.

**Experimental Design**

Sixteen male albino mice were randomly assigned to two experimental groups and housed as eight animals per cage, and each group was treated as the following: **Group A:** served as normal control group and was given daily intraperitoneal injection of distilled water at dose (5ml/kg) for 5 days. **Group B:** was given daily intraperitoneal injection of Cytarabine at dose (100mg/kg) (11) for five days. At the end of experiment, all animals of each groups were sacrificed and liver in addition to the buccal mucosa were excised and preserved in 10% buffered formalin for histological investigation.

**Tissue preparation for histopathological study**

Liver and buccal mucosa of each mice were prepared for histological study by preserving them in solution of 10% neutral buffered formalin for 24 hours in order to be fixed, after that the tissues were dehydrated by using a gradual concatenations of ethyl alcohol (70%-100%) for a period of (30minutes) for each concentration. Then the samples were cleared in 2 separated xylene changes before Passing them in 2 stages of paraffin wax at 57 degree temperature for impregnation , then embedding with paraffin in blocks for sectioning. Following that the samples were cross sectioned at 5 μm thickness, later on it stained by hematoxylin and eosin (HE) stain to examine the histological changes by means of light microscope (12).

**RESULTS**

Histopathological alterations of liver sections in mice in control group sections showed normal cellular and structural details including normal hepatic cord cells around central vein and normal portal areas Figure (1) where as in cytarabine treated group there was defused vacuolar swelling with abnormal hepatic cords patterns and more sever at centrio-lobular Zones also there was multiple foci of apoptotic cells, congested sinusoids and minimal inflammatory response were noticed. Figures (2, 3)
Figure (1): Photomicrograph of mouse liver section from control group showing normal cellular and architectural details of the central vein (→) and hepatic cords (→). Staining H&E. Magnification 200 X.

Figure (2): Photomicrograph of mouse liver section from cytarabine treated group showing moderately congested sinuses (→) acute centro-lobular hepatic cell swelling with hepatic cords dysregulation (→). Staining H&E. Magnification 145 X.
In concern with histopathological modifications in buccal mucosa in control group showed normal squamous epithelial lining, normal mucus – salivary glands, normal ducts opening normal vasculature and under lining tissue. Figures (4, 5)

Figure (3): Photomicrograph of mouse liver section from Cytarabine treated group showing acute vacuolar cell swelling (→) different stages of Apoptotic necrosis of hepatic cells (→). Staining H&E. Magnification 256 X.

Figure (4): Photomicrograph of mouse buccal mucosa section from control group showing normal epithelium (→) normal salivary glands (→) normal mucous glands (→) and normal ducts (→). Staining H&E. Magnification 115 X.
Figure (5): Photomicrograph of mouse buccal mucosa section from control group showing normal epithelium (→) normal mucous glands (→) and normal lobular ducts (→). Staining H&E. Magnification 256 X.

While in cytarabine treated group revealed a severe shrinkage and atrophied appearance of mucus-salivary glands, zymogen granules depletion in the cytoplasm of serous cells of the glands, reduction of diameter of inter lobular and lobular salivary ducts, vacuolation f the stratified squamous epithelium and interstitial edema. Figures (6, 7)

Figure (6): Photomicrograph of mouse buccal mucosa section from Cytarabine treated group showing shrinkage and atrophy of mucous glands (→) interstitial sub mucosal edema (→) empty lobular ducts (→). Staining H&E. Magnification 115 X.
DISCUSSIONS

Histology of liver sections in cytarabine group showed defused vacuolar swelling with abnormal hepatic cords patterns also there was congested sinusoids and multiple foci of apoptotic cells and minimal inflammatory response were noticed. This current result explained by oxidative damage that leads to mitochondrial DNA damage (mtDNA) and those changes related to DNA fragmentation and apoptosis initiation\(^{(13)}\).

This result in agreement with previous study of histological changes of cytarabine in tissue sections of liver and shown dissimilar levels of hepatic apoptosis and cell necrosis which can lead to incomplete or disappearance of the nucleus and cell membrane. Apoptotic cells will separately scatter in tissues with concentration of the nuclear chromatin and spallation of the nucleus\(^{(14)}\).

As well as similar to result obtained by other study of the histopathological effects of cisplatin, doxorubicin and 5-fluouracil on the liver of rats and mentioned that there is ultrastructural abnormalities in the liver including marked disruption of hepatic cords and dilated blood sinusoids, inflammatory infiltration, periportal fibrosis, hyperplasia and many hepatocytes showed karyomegaly and pyknotic nuclei representing apoptosis\(^{(15)}\).

Regarding histopathological result of current study in buccal mucosa showed severe shrinkage and atrophied appearance of mucus-
salivary glands, zymogen granules depletion in the cytoplasm of serous cells of the glands, reduction of diameter of interlobular and lobular salivary ducts, vacuolation of the stratified squamous epithelium and interstitial edema. This result may be explained through the fact that Cytarabine is determined as most recorded mucotoxic agents that damage the entire gastrointestinal tract, from oral cavity to anus (16). In general chemotherapy are drugs that having a direct cytotoxic activity, so any normal tissue from the body is possibly vulnerable and can be affected, earlier or later and due to the lack of high selectivity of the antineoplastic drug on tumor tissue that can’t distinguish between cancer cells and cells of rapidly dividing normal tissue, such as digestive epithelial tissue as a result it can cause structural or functional changes in the digestive tract (17). Most of chemotherapeutic agent like cytarabine acts through direct DNA damage then cytokines enter the circulation and activate an inflammatory cascade. Both intrinsic and extrinsic apoptotic pathways are up regulated and mucosal integrity is damaged by inflammatory infiltrates and tight junction interruption (18). Additionally, Primary mucosal cell injuries resulting from oxidative stress lead to the genes expression as early response and activation of DNA transcription factor as well as chemotherapy affect microflora diversity and load; this imbalance responsible for opportunistic infections or the reactivation of latent viruses (19). This current result in line with previous study of pathophysiology behind xerostomia and reduced function of the salivary glands during and after chemotherapy and mentioned that Antimetabolites chemotherapy (5-fluorouracil, methotrexate and cytarabine) cause nuclear degeneration, Vacuolization, ductal dilatation, inflammation, cyst formation and reduced salivary flow rate (20). Also this result similar to result obtained by other study regarding the effect of methotrexate on parotid gland and mentioned that most of the serous acini had irregular outlines and was widely separated; ductal cells were faintly stained with variable sized vacuoles displacing the nuclei more peripherally and marked hemorrhage between acinar cells and the cytoplasm (21). As well as in consistence with another study regarding the effect of methotrexate on rabbits submandibular salivary gland and reported that there was a glandular degeneration detected by marked vacuolations in acinar and ductal cells and complete replacement of some acinar cells by large vacuoles (22).

**CONCLUSIONS**

Although cytarabine is the highly effective in treatment of cancer but it is associated with direct cytotoxicity to organs and tissues such as liver and buccal mucosa so caution should be
taken when giving the drug to patient with liver
disease or failure and salivary glands
dysfunction.

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