Lycopene treatment stalls the onset of experimentally induced hepatocellular carcinoma: a radioisotopic, physiological and biochemical analysis

Nisha Bhatia¹, Baljinder Singh², Ashwani Koul¹

¹Department of Biophysics, Panjab University, Chandigarh 160014, India.
²Department of Nuclear Medicine, Post Graduate Institute of Medical Sciences and Research (PGIMER), Chandigarh 160012, India.

Correspondence to: Prof. Ashwani Koul, Department of Biophysics, Basic Medical Sciences (BMS) Block II, South Campus, Sec-25, Panjab University, Chandigarh 160014, India. E-mail: drashwanikoul@yahoo.co.in; ashwanik@pu.ac.in


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Abstract

Aim: The present study was aimed to determine the modulatory role of lycopene enriched tomato extract (LycT) during initiation of N-nitrosodiethylamine (NDEA) induced hepatocellular carcinoma (HCC).

Methods: Female Balb/c mice were divided into 4 groups: control, NDEA (200 mg NDEA/kg b.wt, cumulative dose), LycT (5 mg/kg b.wt, thrice a week) and LycT + NDEA. LycT administration was commenced 2 weeks prior to NDEA administration in LycT + NDEA group.

Results: NDEA treatment caused histopathological alterations in hepatic tissue and was associated with enhanced serum levels of inflammatory markers, i.e., tumor necrosis factor-alpha, interleukin (IL)-6 and IL-β. NDEA treatment also induced functional alterations in liver as evident by slow ⁹⁹m⁴Tc-mebrofenin hepatic excretion. LycT administration to NDEA mice showed improved hepatic functional status as demonstrated by normal ⁹⁹m⁴Tc-mebrofenin excretion. NDEA treatment also caused alterations in the hematological parameters such as hemoglobin, red blood cells, platelets and total leucocyte counts. A significant increase in plasma lipid peroxidation and decrease in reduced glutathione levels with alterations in various enzymatic antioxidants were observed upon NDEA treatment. LycT pre-treatment aided in boosting the antioxidant defense system and ameliorated the inflammatory and hematological alterations.

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Conclusion: As evident by improved functional, hematological and biochemical markers, it may be inferred that LycT has the potential to delay HCC initiation.

Keywords: Lycopene, ⁹⁹mTc-mebrofenin, hepatocellular carcinoma, hematology, oxidative stress, inflammation

INTRODUCTION

Environmental factors including exposure to chemical pollutants due to growing industrialization and adoption of unhealthy lifestyle choices like physical inactivity, imbalanced dietary regimen and smoking play a major role in the etiology of various ailments including cancer. The existing structural embodiment of our environment and surroundings makes impossible to evade the exposure from its pernicious clutches. Among various pollutants, N-nitrosodimethylamine (NDEA) is a potent environmental carcinogen found in air, soil, water and food[1-3]. Metabolic activation of NDEA through hepatic biotransformation enzymes renders liver as a target organ for carcinogenesis. Hepatocellular carcinoma (HCC) is figured to be the fifth most common tumors of liver across the globe and accounts for 2.5% increase of death rates every year[4]. Extremely high incidence rate, poor prognosis and asymptomatic behavior associated with HCC makes its diagnosis restrictable at initial stages. Furthermore, angiogenesis and malignant nature of tumorous tissue may also reduce the life expectancy of HCC patients at later stages.

Early HCC diagnosis has become a priority in improving the survival among cancer patients. Despite the availability of various modalities for HCC treatment such as surgical resection, radiation and chemotherapy, the outcome of patient remains dismal. Additionally, various non-invasive modalities such as ultrasound, computed tomography, and magnetic resonance imaging are in use for early monitoring, diagnosis and stratification of HCC[5]. But these provide only anatomical information not the functional status of the tumorous tissue. Therefore, the development of non-invasive diagnostic techniques and identification of tumor specific serological markers linked to early stages of carcinogenesis would be of great clinical significance in the early management of HCC patients. This may be accomplished by ⁹⁹mTc-mebrofenin hepatobiliary functional test that mainly relies on functional perturbations in the tissue as functional alterations precede anatomical alterations[6]. The main rationale behind the identification of candidate biomarkers relevant for cancer risk is to find a link between biological alterations in nonspecific tissues, such as blood, and the occurrence of similar events in specific tissues involved in the carcinogenesis. Moreover, blood acts as a pathological reflector of the systemic status of an animal exposed to carcinogen[7].

Hematological parameters are surrogate markers playing a key role in diagnosing the extent of damage to blood thus acting as a prognostic indicator of HCC patients[8]. NDEA is also known to induce chronic inflammatory responses characterized by the upregulation of pro-inflammatory cytokines and recruitment of innate immune cells to liver tissue. Chronic inflammation further predisposes hepatic tissue to the development of HCC[9]. Free radicals and reactive oxygen species (ROS) generation during NDEA metabolism further causes the oxidation of major cellular biomolecules thus may augment an oxidative stress which may be postulated as a major contributor in the genesis of cancer[10]. Various reports support the relationship between the hepatic antioxidant system and development of hepatocarcinogenesis[11,12]. However, only few studies are available to evaluate the role of blood antioxidants in early diagnosis of HCC[13,14]. These observations emphasize the need for urgent implementation of efficient strategies to curb this disease. From the past few decades, preventive control approaches using the natural products derived from common dietary sources have been the main focus of scientific research to impede the induction of carcinogenesis[15].

Lycopene extracted from red tomatoes has found its widespread use in natural medicine because of its highest antioxidant and radical scavenging activity[16-18]. Consumption of lycopene enriched tomato extract
has been revealed to be effective in alleviating cancer progression due to its increased bioavailability and synergistic effects of its multiple phytochemicals\cite{19,20}. The ameliorative potential of lycopene enriched extract has been found in patients of oesophageal cancer\cite{21}. The consistently reduced risk of chronic diseases associated with increased consumption of lycopene enriched products provides a strong foundation for its use as a potent chemopreventive agent against liver cancer.

Our earlier studies have reported the delay in progression of hepatic cancer upon lycopene enriched tomato extract (LycT) consumption which was revealed by reduced histopathological alterations, improved survival rate, reduced tumor incidence and burden\cite{3,18}. This was also evident through modulation in the expression of apoptosis and cell proliferation associated genes which further interferes in the progression of tumor cells\cite{18,22}. We have also found that LycT consumption aided in up regulating the detoxification system, reducing chromosomal aberrations and modulating physiochemical characteristics of hepatocellular membrane\cite{12}. Recently, in our laboratory, the role of LycT in inhibiting multiple dysregulated pathways including hypoxia, angiogenesis and metastasis has also been delineated. The study suggested that it does so by attenuating the expression of hypoxia inducible factor-alpha, vascular endothelial growth factor, cluster of differentiation 31, matrix metalloproteinases (MMP)-2 and MMP-9\cite{23}. Moreover, the modulation of hepatic tumor marker [alpha fetoprotein (AFP)] and hepatic functional markers by LycT was also demonstrated\cite{23}.

Thus the current scientific scenario has prompted us to study HCC during its early stages of development by analyzing a panel of hematological, inflammatory and blood antioxidant markers whose dysfunction may be related to critical events in hepatic cancer progression and their intervention with LycT. The assessment of these markers in blood on a regular basis along with AFP and liver function markers may allow earlier HCC detection. In addition, the physiological perturbations occurring in the hepatic tissue during carcinogenesis was also assessed using $^{99m}$Tc-mebrofenin hepatobiliary functional test.

**METHODS**

**Animal model for development of HCC**

Female Balb/c mice (25-30 g) procured from the Central Animal House facility of Panjab University, Chandigarh (India) were provided standard animal pellet diet (Ashirwad Industries, Kharar, Punjab, India) and drinking water ad libitum. The animal house was maintained at a controlled temperature of 21 ºC ± 1 ºC and humidity of 50%-60% with a 12-h dark and light cycle. All the experimental studies were performed in accordance with the Indian National Science Academy Guidelines for the use and care of experimental animals and were initially approved by the Institutional Animal Ethics Committee (IAEC), Panjab University, Chandigarh (IAEC/284-295 at Sr. No. 47). The mice were acclimatized to the experimental conditions for duration of 1 week prior to the commencement of various treatments. LycT was extracted from red tomatoes using hexane/acetone/ethanol as an extraction medium as described by Gupta \textit{et al.}\cite{3}. The content of lycopene in the extract was estimated using UV-VIS spectrophotometer as described earlier\cite{3}. LycT in the upper hexane layer showed the presence of three characteristic peaks, i.e., at 444, 470 and 503 nm. Lycopene quantification was performed at 503 nm as to avoid the interferences from other carotenoids including β-carotene, lutein, neoxanthin, etc.\cite{24,25}. The average lycopene content was approximately 14 mg/kg tomato\cite{3}.

Female Balb/c mice were randomly segregated into 4 groups. Animals of group 1 (control) were given vehicle (olive oil) treatment orally thrice a week. Animals of group 2 (NDEA) and group 4 (LycT + NDEA) received an intraperitoneal injection of NDEA at a cumulative dose rate of 200 mg/kg body weight for a total duration of 8 weeks. Group 3 (LycT) and group 4 animals were administered LycT in olive oil orally at a dose rate of 5 mg/kg body weight thrice a week for 10 weeks. Oral administration of LycT was commenced 2 weeks prior to NDEA treatment and continued until the termination of experimental period in LycT + NDEA group.
Assessment of hepatobiliary function
At the end of 10th week, $^{99m}$Tc-mebrofenin hepatobiliary functional test was performed to assess hepatocellular function, biliary obstruction and to quantify hepatic extraction fraction (HEF). 185-200 MBq of $^{99m}$Tc-sodium pertechnetate prepared in normal saline was mixed with mebrofenin according to the instructions provided by the manufacturer (BRIT, India). The scintillator counter was calibrated at 140 KeV with a window setting of ± 20% using $^{99m}$Tc as a radioactive source. Mice were then positioned over the scintillation counter immediately after the intravenous administration of $^{99m}$Tc-mebrofenin with liver and mediastinum in the field of view. Liver activity and blood pool activity was monitored as a function of time and then used to measure the percentage of activity retained by the hepatic tissue (hepatic retention). The time required for maximum uptake of mebrofenin ($T_{peak}$) as well as the time at which the activity reduces to its half ($T_{1/2 \, peak}$) was also calculated for the hepatic and cardiac tissues.

Assessment of hematological parameters

Collection of whole blood samples
Blood samples from mice of different groups were collected through an ocular vein in sterilized eppendorf containing anticoagulant at the end of treatment period. Blood samples were mixed properly and processed for the estimation of various hematological parameters.

Hemoglobin
The hemoglobin (Hb) content was estimated in whole blood by cyanmethemoglobin method as given by Dacie and Lewis[26]. The estimation is based on the oxidation of hemoglobin to cyanmethemoglobin in presence of potassium ferricyanide. The intensity of red colored complex thus formed was measured spectrophotometrically at 540 nm against a Drabkin's solution and expressed as g/dL.

Red blood cells
Total red blood cell (RBC) counts were measured in non-coagulated whole blood as per the method described by Dacie and Lewis[26]. Hayem's fluid is an isotonic solution consisting of sodium sulphate, sodium chloride and mercuric chloride. The measurement is based on the dilution of blood samples with Hayem's fluid and then counting of the RBCs in four corners and one central square of a Neubauer's chamber. The RBC counts were further expressed as counts × 10$^6$/mm$^3$.

Total leucocyte count
The counting of total leucocytes (TLC) was performed in whole blood according to the method of Dacie and Lewis[26] using Turk's fluid. The glacial acetic acid in this fluid causes the destruction of RBCs while the gentian violet helps in the staining of white blood cells (WBCs) nuclei makes them visible under the microscope. Counting of cells was done in four corner WBC squares and expressed as counts × 10$^3$/mm$^3$.

Platelet count
The counting of blood platelets was done in a hemocytometer using 1% ammonium oxalate as a platelet diluting fluid[24]. In this, oxalate induces the complete hemolysis of RBCs and preservation of platelets. The number of platelets were counted in whole blood in all the central RBC squares under a microscope and expressed as counts × 10$^5$/mm$^3$.

Differential leucocyte count
Differential leucocyte counting was performed to compute the presence and number of different type of leucocytes in blood according to the method of Dacie and Lewis[26]. The percentage (%) counts of neutrophils and lymphocytes were determined by observing a blood smear under a microscope.
Assessment of inflammatory markers
The quantitative estimation of various inflammatory markers including tumor necrosis factor (TNF)-α, interleukin (IL)-1β and IL-6 was carried out in serum using a commercially available kit by solid phase enzyme linked immunosorbent assay according to the instruction provided by the manufacturer (RayBiotech, Inc., USA).

Assessment of antioxidant defense system
Preparation of blood plasma
At the end of various treatments, blood was withdrawn from the retro-orbital plexus of a mouse eye in an eppendorf containing EDTA as an anticoagulant. This was followed by the centrifugation of blood samples at 3000 rpm for 15 min at 4 °C. The supernatant (plasma) thus obtained was used for the estimation of various oxidative stress markers.

Lipid peroxidation
Lipid peroxidation (LPO) levels were measured in plasma as per the method described by Trush et al.\textsuperscript{[27]}. It is based on the reaction of malondialdehyde (MDA) and thiobarbituric acid (TBA) to form pink colored MDA-TBA complex which has its maximum absorption intensity at 532 nm. The amount of chromophore thus obtained was measured as an index of lipid peroxidation using an extinction coefficient of $1.56 \times 10^5$ M$^{-1}$cm$^{-1}$ and expressed as nanomoles of MDA-TBA chromophore formed/min/mg protein.

Reduced glutathione
The plasma levels of reduced glutathione (GSH) were estimated as a total non-protein sulphydryl compound according to the method of Moron et al.\textsuperscript{[28]}. It involves the reduction of a 5,5'-dithiobis-2-nitrobenzoic acid by the -SH group of reduced GSH to produce a yellow colored 2-nitro-5-mercaptobenzoic acid. The optical density of the compound thus produced was measured spectrophotometrically at 412 nm and expressed as nanomoles of GSH/mg protein.

Glutathione-S-transferase
Plasma activity of glutathione-S-transferase (GST) was estimated using a method of Habig et al.\textsuperscript{[29]}. GST aids in the coupling of GSH with a substrate, i.e., 1-chloro-2, 4 dinitrobenzene (CDNB). The absorbance of chromophore thus formed was read at 340 nm and described as micromoles of GSH-CDNB conjugates formed/min/mg protein using an extinction coefficient of 9.6 mM$^{-1}$cm$^{-1}$.

GSH peroxidase
Plasma GSH-peroxidase (Px) activity was assayed as per the method given by Paglia and Valentine\textsuperscript{[30]}. It catalyzes the production of GSSG from GSH with the simultaneous oxidation of NADPH. The change in optical density was read at 340 nm based on an extinction coefficient of 6.22 mM$^{-1}$cm$^{-1}$ and expressed as nanomoles of NADPH consumed/min/mg protein.

Glutathione reductase
Plasma glutathione reductase (GR) activity was estimated according to the method of Williams and Arscott\textsuperscript{[31]}. GR causes the reduction of GSSG to GSH using NADPH as a reducing agent with the simultaneous conversion of FAD to FADH$^\text{+}$. The change in optical density was monitored at 340 nm and calculated as nanomoles of NADPH consumed/min/mg protein using an extinction coefficient of 6.22 mM$^{-1}$cm$^{-1}$.

Catalase
Catalase (CAT) activity was determined in plasma using the method of Luck\textsuperscript{[32]}. CAT assists in the breakdown of hydrogen peroxide to produce water and molecular oxygen. The activity was assayed at 240 nm and measured as international units (IU)/mg protein based on the extinction coefficient of 0.0394 mM$^{-1}$cm$^{-1}$.
Superoxide dismutase
Plasma superoxide dismutase (SOD) activity was assayed according to the method of Kono[33] based on the ability of SOD to inhibit the reduction of nitroblue tetrazolium (NBT) mediated by superoxide radicals, which were produced by photo-oxidation of hydroxylamine hydrochloride. One unit of SOD activity is the amount of enzyme required to cause 50% inhibition in optical density. The rate of reduction of NBT complex was measured at 560 nm and described as IU/mg protein.

Assessment of histopathological alterations
After the completion of the treatment period, mice were euthanized by decapitation and liver tissues were removed carefully followed by the immediate fixation of tissue in neutral formalin. Fixed liver tissue samples were embedded, sectioned and then stained using hematoxylin and eosin staining and examined for the microscopic alterations.

Statistical analysis
Data was expressed as mean ± standard deviation (SD). Statistical significance was analyzed using one-way analysis of variance followed by least significant difference (LSD) post hoc test.

RESULTS
Effect of LycT and/or NDEA on 99mTc-mebrofenin hepatobiliary functional test
After intravenous administration of 99mTc-mebrofenin, NDEA induced a significant delay in the hepatocyte uptake, retention and excretion of 99mTc-mebrofenin in comparison to control, LycT and LycT + NDEA mice [Figure 1A-D]. The hepatic extraction fraction at the end of 60 min of 99mTc-mebrofenin injection was observed to be around 8.9%, 61.9%, 11.5% and 17.8% in case of control, NDEA , LycT and LycT + NDEA group respectively [Figure 1E]. Hepatic T_peak value was observed to be around 5 min in both control and LycT animals [Figure 2A]. However, NDEA administration led to a significant delay in attaining maximum activity thus showing a T_peak value of around 10 min while animals of LycT + NDEA group showed T_peak value of around 7 min [Figure 2A]. The hepatic T_1/2_peak value in case of control and LycT animals was observed to be around 7-8 min while it was around 22 min in case of LycT + NDEA group [Figure 2B]. In contrast, NDEA animals did not show any T_1/2_peak value which may be due to the extremely slow excretion rate of 99mTc-mebrofenin.

Effect of LycT and/or NDEA on hematological parameters
Hb
A significant decline in Hb levels was observed in mice of NDEA group when compared to control and LycT (P ≤ 0.001) groups [Table 1]. Likewise, animals of LycT + NDEA group showed a decrease in Hb levels in comparison to control and LycT (P ≤ 0.05) mice. However, a significant elevation in Hb levels was observed in mice that received LycT along with NDEA treatment when compared to NDEA (P ≤ 0.001) alone injected mice [Table 1]. No significant change in Hb levels was noticed between control mice and LycT administered mice.

RBC
NDEA administration caused a significant decrease in RBC counts in mice of NDEA group when compared to control and LycT (P ≤ 0.001) groups [Table 1]. Also, a decrease in RBC counts was observed in mice of LycT + NDEA group when compared to control and LycT (P ≤ 0.01) groups. But the increase in RBC counts was observed when NDEA mice were pre-treated with LycT when compared to NDEA (P ≤ 0.01) intoxicated mice [Table 1]. The counts of RBC did not differ significantly between mice of control and LycT groups.

TLC
TLCs were found to be enhanced in mice of NDEA group when compared to control and LycT (P ≤ 0.001) groups [Table 1]. In addition, mice of LycT + NDEA groups also showed elevated total leucocyte counts


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when compared to control and LycT ($P \leq 0.01$) groups. However, the mice that received LycT in addition to NDEA treatment showed a reduction in TLC when compared to NDEA ($P \leq 0.01$) mice [Table 1]. Moreover, no statistical difference in the blood TLC was found in LycT per se group and control group.

Platelets
The blood platelet counts were seen to be significantly decreased in NDEA mice as compared to control and LycT ($P \leq 0.001$) mice [Table 1]. Likewise, a significant reduction in platelets counts was observed in LycT + NDEA group when compared to control ($P \leq 0.05$) and LycT ($P \leq 0.01$) group. In contrast, supplementation of NDEA mice with LycT induced a significant increase in platelet counts when compared to NDEA ($P \leq 0.001$) mice. LycT pretreatment to mice did not induce any alterations in platelets counts in comparison to normal control mice.

Neutrophils
NDEA administration led to a significant enhancement in the neutrophil counts when compared to control and LycT ($P \leq 0.001$) mice [Table 1]. Pretreatment of LycT to NDEA exposed mice also induced a marked increase in neutrophil counts in comparison to LycT ($P \leq 0.01$) mice and remained unaltered when compared...
to control mice. A significant decrease in these counts was observed in LycT + NDEA group when compared to NDEA (\(P \leq 0.05\)) group. No statistical alterations in the neutrophil counts were observed in LycT per se group and control group.

**Lymphocytes**

NDEA treatment exhibited a marked decline in blood lymphocyte counts when compared to control and LycT (\(P \leq 0.001\)) mice. However, no change in the lymphocyte counts was observed in LycT + NDEA group when compared to control and LycT groups [Table 1]. In contrast, LycT supplementation to NDEA exposed mice induced a significant enhancement in lymphocyte counts when compared to NDEA (\(P \leq 0.01\)) intoxicated mice. No change was observed in the blood lymphocyte counts in LycT group when compared to control group.

**Effect of LycT and/or NDEA on serum inflammatory markers**

The levels of serum TNF-\(\alpha\), IL-1\(\beta\) and IL-6 were found to be elevated in mice exposed to NDEA when compared to control and LycT (\(P \leq 0.001\)) mice. LycT supplementation to NDEA animals also induced a significant enhancement in the levels of TNF-\(\alpha\) and IL-1\(\beta\) while IL-6 levels remained unaltered as compared to control and LycT (\(P \leq 0.001\)) mice [Figure 3A-C]. On the contrary, animals of LycT + NDEA group revealed a marked reduction in the levels of these inflammatory cytokines in comparison to NDEA (\(P \leq 0.001\)) afflicted group. No significant alterations in their levels were noticed between mice treated with LycT and control mice.

**Effect of LycT and/or NDEA on plasma enzymatic and non-enzymatic antioxidants**

**LPO**

NDEA treatment significantly raised the plasma LPO levels when compared to control and LycT (\(P \leq 0.001\)) mice [Table 2]. Further, a significant elevation in plasma LPO levels was also observed upon LycT
supplementation to NDEA animals when compared to control and LycT \((P \leq 0.001)\) animals. However, the levels of LPO came down to the baseline levels in LycT + NDEA group in comparison to NDEA \((P \leq 0.001)\) group. No statistical difference in the plasma LPO levels was found in LycT per se group and control group.

**Reduced GSH**

The levels of plasma GSH were found to be declined in NDEA and LycT + NDEA mice as compared to control and LycT \((P \leq 0.001)\) mice [Table 2]. However, LycT pre-treatment to tumor bearing mice caused a significant enhancement in their levels when compared to NDEA \((P \leq 0.001)\) mice. No significant alterations were observed in plasma GSH levels of LycT per se group and control group.

**Table 2: Effect of LycT and/or NDEA on plasma antioxidant defense system in different treated groups**

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>NDEA</th>
<th>LycT</th>
<th>LycT + NDEA</th>
</tr>
</thead>
<tbody>
<tr>
<td>LPO (nmol of MDA-TBA chromophore formed/mg protein)</td>
<td>0.02 ± 0.002</td>
<td>0.08 ± 0.003 (^a)</td>
<td>0.03 ± 0.002 (^b)</td>
<td>0.05 ± 0.005 (^{a,b,c})</td>
</tr>
<tr>
<td>GSH (nmol of GSH/mg protein)</td>
<td>7.60 ± 0.23</td>
<td>4.11 ± 0.45 (^a)</td>
<td>7.29 ± 0.20 (^b)</td>
<td>6.25 ± 0.59 (^{a,b,c})</td>
</tr>
<tr>
<td>GR (nmol of NADPH oxidized/min/mg protein)</td>
<td>1.06 ± 0.08</td>
<td>0.66 ± 0.06 (^a)</td>
<td>1.01 ± 0.04 (^b)</td>
<td>0.88 ± 0.04 (^{a,b,c})</td>
</tr>
<tr>
<td>GSH-Px (nmol of NADPH oxidized/min/mg protein)</td>
<td>0.67 ± 0.08</td>
<td>1.06 ± 0.09 (^a)</td>
<td>0.68 ± 0.06 (^b)</td>
<td>0.81 ± 0.05 (^{a,b,c})</td>
</tr>
<tr>
<td>GST (μmol GSH-CDNB conjugates/min/ mg protein)</td>
<td>0.40 ± 0.04</td>
<td>0.55 ± 0.02 (^a)</td>
<td>0.38 ± 0.04 (^b)</td>
<td>0.46 ± 0.03 (^{a,b,c})</td>
</tr>
<tr>
<td>SOD (IU/mg protein)</td>
<td>0.10 ± 0.008</td>
<td>0.18 ± 0.010 (^a)</td>
<td>0.11 ± 0.007 (^b)</td>
<td>0.14 ± 0.014 (^{a,b,c})</td>
</tr>
<tr>
<td>CAT (μmol/min/mg protein)</td>
<td>0.61 ± 0.01</td>
<td>1.10 ± 0.10 (^a)</td>
<td>0.58 ± 0.03 (^b)</td>
<td>0.84 ± 0.02 (^{a,b,c})</td>
</tr>
</tbody>
</table>

Values are expressed as mean ± SD \((n = 5)\) and analyzed by one-way analysis of variance followed by post hoc test. \(^a\) \(P \leq 0.001\) and \(^{a,b}\) \(P \leq 0.01\), significant as compared to control group; \(^b\) \(P \leq 0.001\) and \(^{a,b,c}\) \(P \leq 0.01\), significant as compared to NDEA group; \(^c\) \(P \leq 0.001\), significant as compared to LycT group. NDEA: N-nitrosodiethylamine; LycT: lycopene enriched tomato extract; LPO: lipid peroxidation; GSH: glutathione; GR: glutathione reductase; GSH-Px: GSH-peroxidase; GST: glutathione-S-transferase; SOD: superoxide dismutase; CAT: catalase; MDA-TBA: malondialdehyde-thiobarbituric acid; NADPH: nicotinamide adenine dinucleotide phosphate; CDNB: 1-chloro-2, 4 dinitrobenzene

![Graph A](image1.png)

**Figure 3:** Effect of LycT and/or NDEA on serum inflammatory markers in different treatment groups. Values are expressed as mean ± SD \((n = 5)\) and analyzed by one-way analysis of variance followed by post hoc test. \(^a\) \(P \leq 0.001\), significant as compared to control group; \(^b\) \(P \leq 0.001\), significant as compared to NDEA group; \(^c\) \(P \leq 0.001\), significant as compared to LycT group. NDEA: N-nitrosodiethylamine; LycT: lycopene enriched tomato extract; TNF: tumor necrosis factor; IL: interleukin
GST
NDEA administration induced a significant increase in plasma GST activity when compared to control and LycT \((P \leq 0.001)\) animals [Table 2]. Animals of LycT + NDEA group also showed a significant increase in GST activity in comparison to control \((P \leq 0.01)\) and LycT \((P \leq 0.001)\) animals. On the contrary, mice that received LycT in addition to NDEA showed a significant decrease in plasma GST activity as compared to NDEA \((P \leq 0.001)\) alone administered group. No alterations were observed in plasma GST activity of LycT group when compared to control group.

GSH-Px
NDEA treated mice exhibited a significant increase in plasma GSH-Px activity when compared to control and LycT \((P \leq 0.001)\) mice [Table 2]. Likewise, plasma GSH-Px activity was raised in LycT + NDEA group when compared to control and LycT \((P \leq 0.01)\) group. However, administration of LycT to NDEA group of animals induced a significant reduction in the GSH-Px activity when compared to NDEA \((P \leq 0.001)\) treated mice. Plasma GSH-Px activity did not differ significantly between the control and LycT group of animals.

GR
NDEA exposure exhibited a significant decline in plasma GR activity in comparison with control and LycT \((P \leq 0.001)\) animals [Table 2]. Similarly, a significant decrease in GR activity was also observed in LycT + NDEA group when compared to control \((P \leq 0.001)\) and LycT \((P \leq 0.01)\) group. In contrast, there was a marked increase in GR activity on LycT supplementation to NDEA afflicted animals when compared to NDEA \((P \leq 0.001)\) alone group. No significant alteration in the plasma activity of GR was observed between the control and LycT group.

CAT
A significant increase in plasma CAT activity was observed in NDEA and LycT + NDEA group of animals when compared to control and LycT \((P \leq 0.001)\) mice [Table 2]. In contrast, pretreatment of NDEA exposed animals with LycT induced a significant decrease in CAT activity when compared to NDEA \((P \leq 0.001)\) animals. Plasma CAT activity remained unaltered in mice treated with LycT as compared to control mice.

SOD
The administration of NDEA caused a significant increase in plasma SOD activity in animals of both NDEA and LycT + NDEA group when compared to control and LycT \((P \leq 0.001)\) mice [Table 2]. On the contrary, a significant decrease in SOD activity was observed in LycT + NDEA group when compared to NDEA \((P \leq 0.001)\) afflicted group. There was no significant alteration in SOD activity between mice treated with LycT and normal control mice.

Effect of LycT and/or NDEA on histopathological alterations
Liver sections from control and LycT mice exhibited normal histooarchitecture [Figure 4A and C]. Hexagonal hepatic lobules containing central vein in the middle and portal triad at the periphery were visible. Liver acinus was divided into three zones: zone 1, zone 2 and zone 3. Liver sections from NDEA group showed the presence of high grade dysplasia characterized by architectural and nuclear atypia, differential cytoplasmic staining with diminished sinusoidal spaces and fatty accumulation. No stromal invasion was visible in mice of NDEA group. Liver sections obtained from animals of LycT + NDEA group exhibited mild hepatocellular damage with no vascular invasion [Figure 4B and D; Table 3].

DISCUSSION
Increased oxidative stress and altered redox status during carcinogenesis accentuate the need for developing efficient strategies in curtailing the cancer development. This may be accomplished via use of LycT as an exogenous antioxidant to maintain the redox balance and homeostasis. The role of LycT in delaying
the process of hepatocarcinogenesis has already been studied in terms of diminished histopathological alterations, reduced mortality and induction of apoptosis[3,18]. Moreover, the anti-angiogenic and anti-metastatic potential of lycopene was also investigated recently in our laboratory[23]. Periodical assessment of serological markers at early stages could serve as an important parameter to evaluate the onset of hepatic pathology. Thus the present piece of work was planned to gain insight into the identification of candidate biomarkers linked to early stages of hepatocarcinogenesis and their amelioration by LycT. In addition, the physiological status of the liver was also assessed using non-invasive ⁹⁹ᵐTc-mebrofenin hepatobiliary functional test.

It helps in the evaluation of hepatocellular function, biliary obstruction thus providing the functional or physiological status of liver. In the present study, an appearance of maximum activity of ⁹⁹ᵐTc-mebrofenin in blood pool within 2-3 min in all the groups showed that NDEA exposure does not induce any effect on cardiac tissue. Being a hepatocarcinogen, NDEA induces major pathophysiological alterations in the hepatic tissue as observed by the changes in HEF, percentage counts in the liver at different time intervals,
Control and LycT animals showed the normal uptake, efficient hepatic extraction and rapid excretion from the liver. However, a significant deviation from the normal pattern of radioactivity as observed in NDEA animals confirmed the physiological alterations in hepatic tissue. The delay in hepatic uptake in NDEA animals may be due to severe hepatocellular dysfunction. Neyt et al. [34] have also reported the delayed uptake due to the dysfunction of various Oatp transporters located on hepatocytes. The retention of radiotracer activity in the liver of NDEA mice up to 60 min showed marked impairment in the excretion of the activity. This may be due to the biliary obstruction induced by NDEA which was also evidenced by the inability to calculate T_{1/2} value in case of NDEA animals. We have previously reported the delay in hepatic excretion in the case of DMBA induced hepatotoxicity [35]. Joseph et al. [36] observed the involvement of inflammation in delaying the hepatic excretion of 99mTc-mebrofenin. Similarly, LycT administration to NDEA insulted animals also showed delayed uptake but the clearance of the activity at 60 min showed the protective effect of LycT against hepatocarcinogenesis. Deshpande et al. [37] observed the increased clearance of 99mTc-mebrofenin upon administration of dietary turmeric extract to rats exposed to D-galactosamine HCl. Our laboratory also observed the ameliorative effect of Azadirachta indica against DMBA induced hepatotoxicity by efficient clearance of mebrofenin [35].

During tumor progression, cells generally demand more oxygen than is available for its growth, which results in the creation of hypoxia. Continued hypoxia leads to the adaptation of various genomic and proteomic alterations and results in the aggressive and malignant tumor phenotype. This may further causes reduced oxygen transport throughout the body due to the alterations in various hematological markers. NDEA exposed mice showed a marked decline in the levels of Hb, RBC, platelets and lymphocytes as compared to control mice. In addition to this, a significant enhancement in the neutrophils and WBC counts were observed upon NDEA exposure. The reduction in Hb and RBC suggested the occurrence of anemia in tumor bearing mice. This may be due to the increased oxidative stress induced by excessive ROS which causes the oxidative destruction of mature erythrocytes or inhibiting its production. This can also be evidenced by a decline in GSH and elevation in LPO levels on NDEA treatment [38]. A marked enhancement in WBC counts may reflect the activation of an immune system to fight against invading particles [39]. This was further confirmed by the release of various cytokines by activated kupffer cells and accumulation of neutrophils in hepatic cells as discussed above. Histopathological examination also supported the infiltration of leukocytes in hepatic tissue upon NDEA administration. The diminished platelets count may apparently be due to the decreased production of thrombopoietin hormone by damaged liver cells. The decrease in lymphocytes and enhanced neutrophil counts might suggest the decrease in efficiency of an immune system to cope up with the triggered inflammatory cascade [40]. Similar observations were also noticed by Farooq et al. [41] and Gangar et al. [42], who also observed the alterations in various hematological parameters in patients suffering from gastric and forestomach carcinoma respectively. LycT treatment to NDEA insulted mice tends to restore the levels of these markers which might be attributed to decreased hypoxia and reduction in tumor growth by an enhancement of apoptosis [18,23]. Several other researchers have also supported the restoration of blood parameters upon lycopene treatment thus showing its anti-inflammatory potential [43,44].

Inflammatory cytokines also play a major role in the progression of cancer. Exposure of liver tissue to certain hepatotoxicanits induces the release of pro-inflammatory cytokines by kupffer cells which can further aggravate the tumor progression by triggering of inflammatory cascade [45]. The current findings revealed a marked increase in serum levels of various inflammatory markers, i.e., TNF-α, IL-1β and IL-6 in tumor bearing mice. Overexpression of these cytokines may provide proliferating signals to the mutated hepatocytes through the secretion of angiogenesis and metastasis markers. These findings were in concordance with the report of Abdel-Hamid et al. [46]. Supplementation of NDEA mice with LycT modulated the serum levels of these cytokines by inhibiting their production and induction of apoptosis thus showing its anti-inflammatory effect. Literature also supported the amelioration of these inflammatory markers upon administration of lycopene [47-49].
NDEA treatment also exhibited enhanced plasma LPO and reduced GSH levels in comparison to control and LycT animals. The increase in LPO may be ascribed to the excessive formation of ROS and their diffusion into the blood by an oxidative deterioration of membrane lipids. Declined GSH levels might be due to the impairment of antioxidant defense system and increased consumption of GSH by the detoxification system. Literature also supported the alterations in these oxidative stress markers with the progression of cancer\cite{50,51}. Interestingly, the reversal of their levels by LycT pretreatment may probably be due to the neutralization of free radicals and enhancement of xenobiotic detoxification by LycT. The amelioration of oxidative stress by consumption of tomato enriched diet has also been reported by Dogukan et al.\cite{52} and Gupta et al.\cite{12}.

The enhancement in plasma activities of SOD, CAT and GSH-Px were also observed upon NDEA administration. An elevated SOD activity causes the excessive production of deleterious H2O2 by the dismutation of superoxide anions which may further be counterbalanced by excess CAT and GSH-Px. However, complete neutralization of H2O2 may not occur due to the failure of defense system by sufficient lipid oxidation which further increases the chances of DNA damage, thus, contributing to a growth advantage to the tumorous cell\cite{53}. Our results are in agreement with other reports who found similar alterations in activities of these enzymes in cancer patients\cite{54,55}. Reduction in the activities of these enzymes by LycT supplementation to tumor bearing mice showing the antioxidant capability of LycT to scavenge the free radical formation thus mitigating intracellular oxidative damage. The present results are in harmony with the findings of Ural\cite{56} and Ibrahim\cite{44} who also reported the diminished enzymatic activities on LycT supplementation.

The increase in GST activity in NDEA intoxicated mice may lead to the excessive utilization of this enzyme in the detoxification in response to metabolic induction in the tumor cells. These results are in concordance with the observation of Sadik et al.\cite{57}, who also reported the correlation between the increased GST levels and carcinogenesis. In contrast, Li et al.\cite{58} has observed the decrease in blood GST levels during the development of HCC. Suppression of GST activity upon LycT supplementation to NDEA afflicted mice indicated the protective efficacy of LycT against the induction of oxidative stress. Sadik et al.\cite{57} has also reported the restoration of GST activity upon consumption of a diet rich in fruits and vegetables. The increased activity of GSH-Px and decreased levels of GSH may lead to the accumulation of oxidized glutathione (GSSG) which further cannot be converted to GSH due to the reduction in GR activity upon NDEA treatment. Maffei et al.\cite{59} has also reported the drop in plasma GR activity in patients suffering from colorectal cancer. Pretreatment with LycT to NDEA mice attenuated the decrease in GR activity probably by radical scavenging potential and increase in GSH level thus maintaining the oxido-reductive balance. Lycopene enriched tomato extract ameliorates the oxidative stress by maintaining the integrity of cellular membrane thus preserving the antioxidants levels in the liver cells\cite{12,59}. The activities of various enzymatic and non-enzymatic antioxidants did not show any alterations between control and LycT mice.

It is evident that LycT supplementation modulated the inflammatory and hematological markers, boosted the antioxidant system and improved the functional status of hepatic tissue in tumor bearing mice. Data from the present study and previously published studies reiterate the potential of LycT in delaying the initiation of HCC which may have significant implications in its overall chemopreventive potential.

DECLARATIONS

Authors’ contributions
Study design: Koul A, Singh B
Data analysis and manuscript preparation: Koul A, Singh B, Bhatia N
Experimental studies: Bhatia N
Literature search: Koul A, Singh B, Bhatia N
Manuscript review: Koul A, Singh B, Bhatia N
Data source and availability
The data presented is original and obtained in our laboratory. It is available with the authors and can be made available if required.

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Conflicts of interest
The authors declare that they have no conflicts of interest.

Patient consent
Not applicable.

Ethics approval
All the experimental studies were performed in accordance with the Indian National Science Academy Guidelines for the use and care of experimental animals and were initially approved by the Institutional Animal Ethics Committee (IAEC), Panjab University, Chandigarh (IAEC/284-295 at Sr. No. 47).

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REFERENCES


