Lycopene prevents sugar-induced morphological changes and modulates antioxidant status of human lens epithelial cells

Ipseeta Mohanty, Sujata Joshi, Deepa Trivedi, Sushma Srivastava and S. K. Gupta*

Department of Pharmacology, All India Institute of Medical Sciences, Ansari Nagar, New Delhi - 110029, India

(Received 28 July 2001 – Revised 15 April 2002 – Accepted 2 May 2002)

Cataract is a multifactorial disease. Osmotic stress, together with weakened antioxidant defence mechanisms, is attributed to the changes observed in human diabetic cataract. Epidemiological studies provide evidence that nutritional antioxidants slow down the progression of cataract. The usefulness of lycopene, a dietary carotenoid, in the pathogenesis of human cataracts has not been studied so far. Since the epithelium is the metabolic unit of the lens, the effect of lycopene on galactose-induced morphological changes and antioxidant status of human lens epithelial cells (HLEC) in culture was evaluated in the present study. HLEC of fresh cadaver eyes obtained from an eye bank were cultured in medium supplemented with fetal calf serum (200 ml/l). On confluency, the cells were subcultured in medium containing either 30 mM-d-galactose or 30 mM-d-galactose + lycopene (5, 10 or 20 μM) for 72 h. The cells were observed under the phase-contrast microscope and transmission electron microscope for any morphological changes and then harvested for the estimation of various biochemical variables. Malondialdehyde, glutathione and antioxidant enzymes were significantly altered in the control as compared with the normal cultures. Vacuolization was also observed in the presence of galactose. Addition of lycopene confers significant protection against these changes in HLEC.

Cataract: Human lens epithelial cells: Lycopene

Cataract is the leading cause of blindness worldwide. It is a multifactorial disease and free radicals as well as oxidative stress have been implicated in its pathogenesis (Spector, 1995). A number of studies indicate that in most cataracts the pathological process is triggered by an initial damage to the epithelial cell membrane by the reactive oxygen species generated in situ. The reactive oxygen species generated may result in damage to the lens DNA, oxidation of protein, oxidation of glutathione, lipid peroxidation, disulfide bond formation, protein unfolding with sulfhydryl group exposure, protein aggregation, protein insolubilization and loss of membrane transport function due to oxidation of membrane component such as Na⁺,K⁺-ATPase pump. The lens epithelial cells are the most metabolically active part of the lens and antioxidant enzymes are concentrated mainly in the epithelium (Bermbach et al. 1991).

Osmotic stress in the diabetic cataract has been suggested to be the initiating mechanism of sugar-induced cataract (Collier & Small, 1993). A link between the osmotic and oxidative stress has been established through demonstration of weakened antioxidant defence potential of the system subjected to high sugar concentration (Miyoshi et al. 1999; Obrosova et al. 1999). It has been suggested that auto-oxidation of monosaccharides, leading to the formation of H₂O₂ and free radicals, induces oxidative stress under diabetic conditions (Wolff & Dean, 1985). However, these studies are mainly confined to either isolated animal lenses maintained in organ culture in a high-sugar environment or to diabetic cataract models. Furthermore, certain antioxidants are shown to prevent sugar-induced cataract-type changes in the lens. Lycopene is the major carotenoid present in the diet and provides the familiar red colour of tomato products. Studies suggest that lycopene is a more potent scavenger of singlet oxygen than other major dietary carotenoids (Di Mascio et al. 1989). Protective associations for cervical intraepithelial neoplasia, prostate, gastric and lung cancers are already documented by virtue of its antioxidant property (Clinton, 1998). Carotenoids have been found to exhibit prevention against naphthalene-induced cataract in rats (Jacques & Chylack, 1991). Recently, studies on experimental rats have demonstrated that incorporation of lycopene

**Abbreviations:** CAT, catalase; GSHPx, glutathione peroxidase; GST, glutathione S-transferase; HLEC, human lens epithelial cells; MDA, malondialdehyde.

*Corresponding author:* Professor S. K. Gupta, fax + 91 11 686 2663, email skgupta@hotmail.com
into the diet retards the onset and progression of sugar-induced cataract (Pollock et al. 1996, 1999). However, the role of lycopene in sugar-induced cataracts in human subjects has not been investigated so far.

The present study has been undertaken on human lens epithelial cells (HLEC) with the aim of investigating the effect of galactose on the antioxidant status of these cells subjected to hypergalactosaemic conditions in culture. Furthermore, the potential of lycopene in protecting against the adverse effect of high galactose on HLEC morphology and biochemistry was also evaluated.

Materials and methods

Dulbecco’s modified Eagle’s medium was obtained from Hi Media Laboratories (Mumbai, India). Galactose was procured from Sigma Chemicals (St Louis, USA) and lycopene from Jagspal Pharmaceuticals (New Delhi, India). Gentamicin was obtained from Fulford (Mumbai, India) and betadine from Vasistha Pharmaceuticals (New Delhi, India). Fetal calf serum was from Biological Industries (Hyderabad, India). All other reagents were of analytical grade.

Donor eyes were obtained from the eye bank at the Dr Rajendra Prasad Center for Ophthalmic Sciences, All India Institute of Medical Sciences, New Delhi, India. Eyes were donated to the eye bank for use in corneal transplantation. Eyes were brought in a moist chamber to the laboratory within 8 h of the death of the donor. In all, fifty-six eyes with normal transparent lenses from donors of age range 40–50 years were used for the present study.

Dissection and monolayer culture

The method of Reddy et al. (1988) for culture of HLEC was followed. The eyes were washed with gentamicin (50 ml/l) and betadine (10 ml/l) solution. The cornea and iris were completely removed with the help of corneal scissors and the lens was exposed. An incision was made at the peripheral margin of the lens. The capsule flap was cut into four pieces. Each explant was spread out on the lateral wall of the culture flask (Falcon flask, volume 50 ml, 25 mm²) with the anterior epithelial layer facing upwards. Dulbecco’s modified Eagle’s medium (4 ml) supplemented with fetal calf serum (200 ml/l) and antibiotics (100 µg streptomycin and 100 IU penicillin/ml) was poured in the flask without disturbing the adhered capsule explant. In order to do this, the flask was first set on its end in the CO₂ incubator so that the medium was located at the base of the prepared flask. The rising water vapour protects the tissue from drying, forming a wet chamber. After 1 h the flask was laid down so that the medium covered the epithelium, which was now firmly attached to the surface of the flask. The HLEC explants were incubated at 37°C in a 5% CO₂ atmosphere and the medium was changed twice per week.

Subculture of human lens epithelial cells

HLEC were observed for growth under the microscope. Upon observing confluency, the cells were subcultured. For this procedure, the medium was discarded and the confluent cells were treated with 1 ml trypsin (0.5 mg/l) – EDTA (0.2 mg/l) for 8 min and gently dissociated by pipetting with the addition of fresh medium (Reddy et al. 1991). Following dissociation, the cell number in each flask was determined using a haemocytometer after staining with Trypan Blue (4 mg/l) under a phase-contrast microscope (magnification ×10). HLEC (10⁶ cells/ml) were transferred from primary culture into subsequent passages to maintain HLEC in culture.

For the present study, after trypsinization HLEC were distributed into different culture flasks (10⁶ cells/flask) to serve as normal, control and test group samples. In the normal group flasks, HLEC were incubated with Dulbecco’s modified Eagle’s medium + fetal calf serum (200 ml/l). In the control group flasks, 30 mM-galactose was added to the medium. The culture medium in the test group was supplemented with 30 mM-galactose and 5, 10 or 20 µM-lycopen. During incubation, the lycopene-enriched medium was changed every 6 h. The stock solution of lycopene (1 mM) was prepared in absolute alcohol by vigorous shaking and was clear to the naked eye. The final concentration of alcohol in the culture medium was not more than 0.1 ml/l in all the three test subgroups of lycopene.

Flasks for all the groups were maintained in duplicate and incubated for 72 h. After completion of the incubation period, one set of HLEC from normal, control and test group flasks were observed for any morphological changes under a light microscope (Nikon, Tokyo, Japan) and then processed for transmission electron microscopy (Phillips 410 LS; CA, USA). The other set was utilized for the estimation of biochemical variables.

Electron microscopy

The media in the flasks to be processed for transmission electron microscopy study was discarded. Medium (4 ml) containing an equal volume of fixative (20 ml glutaraldehyde/l 0.1 M-PBS) was added to the flask and kept undisturbed for 1 h at room temperature. The HLEC were then scrapped out using a Falcon scraper (NJ, USA) and centrifuged at 8000g (Remi Instruments, Mumbai, India) for 15 min. For studying these HLEC under transmission electron microscopy, a standard protocol for fixation, dehydration, sectioning and viewing, as described by Eguchi & Okada (1971), was followed.

Stability studies

The stability of microcrystalline aqueous solution of lycopene (1 mM) in Dulbecco’s modified Eagle’s medium was tested by the spectrophotometric method of Stahl et al. (1993). Briefly, 1 ml medium was drawn at 0, 3 and 6 h and to 1 vol. lycopene microcrystalline solution 1 vol. 2-propanol was added and vigorously vortexed for 15 s. Subsequently, 3 vol. n-hexane–dichloromethane (5: 1, v/v) containing 1.2 mM-butylated hydroxytoluene was added and mixed by vigorous vortexing for 30 s. The mixture was then centrifuged briefly for phase separation and the organic phase was read at 472 nm. Freshly prepared lycopene standard was also run at 0, 3 and 6 h. The concentration of lycopene in the medium was calculated using
molar absorption 1.85 × 10^5 optical density units. The concentration of lycopene at 0 h in the medium was considered as 100%.

**Biochemical analyses**

The culture media in the other set of flasks were discarded and HLEC were washed with PBS. The cells were then removed by scraping with a Falcon scraper (NJ, USA), collected in an Eppendorf tube and centrifuged (Remi Instruments) at 8000 g for 10 min. HLEC homogenate (10^9/100 μl) was prepared in 50 mM-phosphate buffer, pH 7, under cold conditions. A sample of this homogenate (100 μl) was used for the estimation of malondialdehyde (MDA). A sample of homogenate was centrifuged (Remi Instruments) at 8000 g (MDA). A sample of homogenate was centrifuged (Remi Instruments) at 8000 g for 15 min at 4°C and the supernatant fraction obtained was used for the estimation of glutathione.

**Estimation of glutathione.** Glutathione estimation was done by the method described by Moron et al. (1979). A protein-free supernatant fraction was obtained by addition of equal volume of TCA (100 g/l) to the HLEC homogenate and centrifuged (Remi Instruments) at 8000 g for 30 min. A sample of this supernatant fraction (0-1 ml) was used for the estimation of glutathione. The volume of the reaction mixture was made up to 1 ml with 0.3 M-Na_2HPO_4 (pH 8) and colour was produced by the addition of 1.25 ml 0.6 M-5,5′-dithiobis(2-nitrobenzoic acid) prepared in trisodium citrate (10 ml/l). The absorbance of the resulting yellow colour was recorded within 10 min at 412 nm on a Beckman’s spectrophotometer (Beckman Instruments, Minnesota, MN, USA). A parallel standard was also run to calculate the amount of glutathione in HLEC.

**Estimation of malondialdehyde.** MDA estimation was done in HLEC homogenate by the method described by Ohkawa et al. (1979). To 0-1 ml homogenate, 0-2 ml SDS (81.0 g/l), 1.5 ml acetic acid (200.0 g/l) and 1.5 ml thiobarbituric acid (8.1 g/l) were added. The mixture was heated for 30 min at 95°C in a temperature-controlled waterbath. After cooling, 5 ml n-butanol pyridine (15:1 v/v) was added to it. The mixture was centrifuged (Remi Instruments) at 8000 g for 10 min. Absorbance of the organic layer was read spectrophotometrically at excitation wavelength 532 nm and emission wavelength 515 nm. 1,1,3,3-tetraethoxypropane was used as standard to obtain a curve for the calculation of unknown MDA in the samples.

**Estimation of antioxidant enzymes.** For the estimation of antioxidant enzymes, HLEC homogenate (10^9/100 μl) was prepared under cold conditions in 50 mM-phosphate buffer, pH 7, and centrifuged at 8000 g (Remi Instruments) for 15 min at 4°C and the supernatant fraction was used for the estimation of glutathione S-transferase (GST), glutathione peroxidase (GSHPx) and catalase (CAT). The effect of 10 μM-lycopene was evaluated on the activity of these antioxidant enzymes.

GST activity was measured by the method of Hebig (1974) at 250°C. The conjugation of GST with 1-chloro-2,4-dinitrobenzene, a hydrophilic substrate, was observed spectrophotometrically at 340 nm. Briefly, the reaction mixture contained 1.7 ml 100 mM-phosphate buffer (pH 6.5), 0.1 ml 30 mM-1-chloro-2,4-dinitrobenzene, 0.1 ml 30 mM glutathione and 0.05 ml HLEC supernatant fraction. The change in absorbance was recorded at 340 nm at 30 s intervals for 3 min. One unit GST is defined as the amount of enzyme required to conjugate 1 μmol 1-chloro-2,4-dinitrobenzene with glutathione/min.

CAT was estimated according to the method of Aebi (1974). For the estimation of CAT H_2O_2 was used as substrate. The assay was conducted at 25°C. The reaction mixture consisted of 1 ml 50 mM-phosphate buffer (pH 7) and 0.05 ml HLEC supernatant fraction. The change in absorbance at 240 nm was followed at 5 s intervals for 30 s, after the addition of 0-1 ml 30 mM H_2O_2 against the sample blank. One unit of CAT activity represents the amount of enzyme required to decompose 1 μmol H_2O_2/min.

GSHPx activity was measured by the method of Paglia & Valentine (1967). GSHPx activity was monitored at 25°C. Briefly, to 0.8 ml 50 mM-phosphate buffer 0.05 ml each of 75 mM-EDTA, 25 mM-sodium azide, 250 mM-glutathione, 7.5 mM-NADPH and 0.05 ml HLEC supernatant fraction were added. Then, 6 units glutathione reductase were added to this reaction mixture. Blank was set and the reaction was then started by the addition of 0.05 ml H_2O_2 (0.075 mm). The change in absorbance/min was recorded at 340 nm for 2 min at intervals of 15 s. GSHPx activity was calculated from the extinction coefficient of NADPH. One unit GSHPx activity is defined as amount of enzyme required to utilize 1 nmol NADPH/min at 37°C.

**Protein estimation.** The method of Lowry et al. (1951) was followed for the estimation of protein levels in HLEC samples. The amount of protein was estimated per million cells.

**Statistical analysis**

The data were presented as mean values and standard deviations. The number of samples in each group was six. Unpaired Student’s t test was applied to compare the results of the normal and the control group and probability value P<0.05 was regarded as statistically significant. ANOVA with post-hoc analysis (Bonferroni multiple range test) was applied to compare control and test groups.

**Results**

**Microscopic studies**

HLEC in primary culture took on an average 10–12 d to attain confluency. Normal HLEC appeared hexagonal in shape, had prominent nuclei and smooth cell margins under the phase-contrast microscope. The nucleus was mostly central and the cytoplasm uniformly distributed (Fig. 1).

Electron microscopy of normal HLEC showed demarcated cell membranes, cytoplasm and nuclei (Fig. 2). HLEC showed intracellular cytoplasmic vacuoles at 72 h incubation in the presence of galactose (Fig. 3). The mean value for the size of the vacuole observed in the presence of galactose at 72 h was 0.34 (SD 0.09) μm. Its volume and area were 0.021 (SD 0.003) μm^3 and 0.37 (SD 0.08) μm^2 respectively. Inclusion of 10 μM-lycopene to the culture medium containing 30 mM-galactose decreased the size
of vacuoles significantly \( (P<0.01, \text{Fig. 4}) \). The size, area and volume of the vacule in the presence of lycopene were 0.27 (SD 0.08) \( \mu \text{m} \), 0.23 (SD 0.05) \( \mu \text{m}^2 \) and 0.01 (SD 0.00) \( \mu \text{m}^3 \) respectively.

**Stability studies**

A significant \( (P<0.05) \) decrease in lycopene concentration was observed at 3 and 6 h as compared with 0 h. The levels were reduced by 24.3 and 41.2\% at 3 and 6 h respectively.

**Effect of lycopene on glutathione and malonaldehyde levels**

A significant fall (55.7\%, \( P<0.05 \)) in glutathione level was observed in the presence of 30 mM-galactose (control) in comparison with the normal group. The basal glutathione levels measured in the HLEC was 87.99 \( \mu \text{g/mg protein} \) whereas the glutathione content of the HLEC in the control group was found to be 38.91 \( \mu \text{g/mg protein} \). No significant effect on glutathione level was seen with 5 \( \mu \text{M}-\text{lycopene as compared with the control group. A positive modulation of glutathione levels was observed with 10} \mu \text{M as well as 20} \mu \text{M-lycopene.} \)

**Fig. 1.** Normal human lens epithelium cells under light microscope (magnification \( \times 10 \)). For details of procedures, see p. 348.

**Fig. 2.** Transmission electron micrograph of normal human lens epithelium cells (magnification \( \times 15000 \)). For details of procedures, see p. 348.
Fig. 3. Transmission electron micrograph of human lens epithelium cells incubated for 72 h in growth medium supplemented with 30 mM-galactose (magnification × 15,000). Intact cell and nuclear membrane with condensation of chromatin. ↑, Vacuolization observed; |—|, 0.4 μm. For details of procedures, see p. 348.

Fig. 4. Transmission electron micrograph of human lens epithelium cells incubated for 72 h in growth medium supplemented with 30 mM-galactose and 10 μM-lycopene (magnification × 15,000). ↑, Vacuoles decrease in number and size; |—|, 0.4 μm. For details of procedures, see p. 348.
Fig. 5. Effect of lycopene on glutathione levels in human lens epithelial cells. Normal cells were cultured in Dulbecco’s modified Eagle’s medium: control, normal + sugar; test, normal + sugar + lycopene. □, Normal; ■, control; ▬, test; (5 μM); □, test (10 μM); ▬, test (20 μM). For details of procedures, see p. 348. Values are means for six samples with standard deviations represented by vertical bars. Mean value was significantly different from normal value: *P < 0.05. Mean values were significantly different from control value: **P < 0.01. Mean values were significantly different from test (5 μM) value: †P < 0.05.

μM-lycopene. However, the difference between the two groups was not significant. The glutathione levels increased to 65.30 and 67.20 μg/mg protein respectively (Fig. 5).

The basal MDA level in HLEC was found to be 1.4 nmol/mg protein. On comparing the MDA levels in the control and the treatment groups, it was observed that incubation in the presence of galactose produced a significant increase in lipid peroxidation in the control group as evaluated by the MDA values (2.6 nmol/mg protein). However, in the presence of 10 and 20 μM lycopene, lipid peroxidation was significantly reduced and malondialdehyde levels were found to be 1.90 and 1.82 nmol/mg protein respectively in the treated group as compared with control (Fig. 6).

**Effect of lycopene on antioxidant enzymes**

The effect of 10 μM-lycopene on different antioxidant enzymes (GST, GSHPx and CAT) is presented in Table 1. It was observed that in the control group the activities of GST and GSHPx enzymes were significantly reduced (P < 0.01) as compared with the normal group. In contrast to GSHPx and GST, a threefold increase in CAT activity was observed in the control group. Lycopene (10 μM) significantly increased GST and GSHPx activity by 88 and 86% respectively as compared with the control group. In addition, lycopene decreased CAT activity significantly (52%, P < 0.05)

**Discussion**

The HLEC are the metabolic unit of the lens and are responsible for maintaining its homeostasis and transparency. Being the most anterior part of the lens, it is the primary site of external insult that ultimately leads to cataract. Therefore, recent efforts are being directed to maintain HLEC in culture (Reddy et al. 1988). The establishment of lens epithelial cultures is of great importance, not only for the study of lens differentiation, but also to investigate the aetiology of cataract and to serve as a useful model for screening various anti-cataract agents.

Cataract is a multifactorial disease associated with several risk factors, diabetes being one of the most important risk factors (Kador & Konoshita, 1984). Epidemiological studies suggest that nutritional antioxidants such as carotenoids delay the onset and progression of cataract (Jacques & Chylack, 1991). Hence, interest in the anti-cataract potential of carotenoids is growing. Lycopene, a
carotenoid that provides the familiar red colour to tomato products, is one of the major carotenoids present in the diet. Lutein and zeaxanthin are the carotenoids that have already been detected in the human lens (Yeum et al. 1999). No report showing the presence of lycopene in the human lens is known to auto-oxidize in the presence of trace metal ions (Jasmina et al. 1994). Formation of vacuoles is presumably due to the osmotic effect of intercellular accumulation of dulcitol. Supplementation with 10 μM-lycopene reduced the number of vacuoles. Present findings are consistent with those reported by Lin et al. (1991), who also observed vacuolization in HLEC incubated in the presence of galactose.

Glutathione plays a very important role in various biochemical processes including lipid peroxidation and it provides a defence mechanism for tissue against the reactive oxygen species. Glutathione also forms one of the substrates for the two antioxidant enzymes GSHPx-1 and GST. The level of glutathione decreased significantly (P < 0.05) in HLEC in the presence of galactose. The levels were significantly (P < 0.01) maintained in the presence of 10 and 20 μM-lycopene. The fall in glutathione levels could be attributed to the excessive diversion of NADPH, a co-substrate for aldose reductase enzyme, towards polyol synthesis and its consequent unavailability for glutathione reductase, leading to an overload on the cell antioxidant mechanisms. An increase in lipid peroxidation product in the presence of galactose clearly suggests membrane damage and involvement of reactive oxygen species in sugar cataracts. High levels of sugars are also known to auto-oxidize in the presence of trace metal ions and generate reactive oxygen species and dicarbonyl sugar derivatives (Wolff & Dean, 1985). Significant (P < 0.01) inhibition of lipid peroxidation and intact membrane by 10 and 20 μM-lycopene was observed. However, the exact mechanism by which lycopene prevents lipid peroxidation is not clear. Lycopene, due to its poor solubility and perhaps in the form of microcrystalline solution, might stick to the outside of the cell and prevent exposure of the lens epithelial cells to reactive oxygen species. However, the exact site of action of lycopene could not be established as the intracellular levels of lycopene were not measured.

A fall in enzyme activities of GSHPx-1 and GST in the control group was observed which might be correlated to decreased availability of its substrate, glutathione. However, lycopene treatment prevented the decrease in glutathione level and positively modulates GSHPx-1 and GST activities.

CAT plays an important role in lens defence mechanisms against the harmful effects of oxygen free-radicals in biological systems (Spector, 1995). The most interesting finding of the present study is the threefold increase in activity of CAT within 72 h in the presence of galactose in incubation medium and its down-regulation by lycopene. In the control group, enhanced CAT activity clearly suggests overproduction of H2O2 in the presence of galactose. On addition of lycopene to the culture medium, it was observed that CAT activity decreased significantly (P < 0.05). Decreased activity of CAT in the presence of lycopene could be explained either through inhibition of autooxidation of galactose itself leading to reduced generation of H2O2, or by the increased utilization of H2O2 by enzymes GSHPx-1 and GST. Increased GSHPx-1 and GST activities in presence of lycopene could probably be a reflection of the same processes. It is also possible that like pyruvate, a physiological antioxidant, lycopene may act through the peroxidative decarboxylation pathway decomposing H2O2 into non-toxic acetic acid, water and CO2 (Varma et al. 1990).

The results obtained from the present study indicate that lycopene offers protection to the lens by modulating various antioxidant variables and could be a potential anti-cataract agent. The present study is consistent with epidemiological evidence of the protective role of lycopene against oxidative damage in the epithelium cortex of the human lens (Lyle et al. 1999; Yeum et al. 1999). Although lycopene is quite unstable in an aqueous medium, adequate levels were found to be present at 3 h (0.76 mm) and 6 h (0.59 mm). Though lycopene levels at 6 h were significantly less than at 0 h (P < 0.05), it appears from the results of the present study that lycopene offers beneficial effects even at this low concentration. However, further studies on lycopene regarding its pharmacokinetics in the human eye should be contemplated for a better understanding of the mechanism and site of action of lycopene.

Acknowledgements

This work was supported by grants from Department of Biotechnology (DBT), India.
References


