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The potential therapeutic efficacy of intravenous versus subconjunctival mesenchymal stem cells on experimentally ultraviolet-induced corneal injury in adult male albino rats

W.A. Nasr El-Din et al., Ultraviolet corneal injury stem cells

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ABSTRACT

Background: The X-rays and the visible light are the main source of ultraviolet radiation (UVR). About 90% of ultraviolet B (UVB) are absorbed by the cornea which may promote corneal inflammation, oedema and damage of its epithelial layer. Bone marrow mesenchymal
stem cells (BM-MSCs) have been demonstrated to ameliorate the injured corneal tissue and accelerate its wound healing. This study aimed to compare the healing effect of intravenous (IV) versus subcutaneous (SC) BM-MSCs on the rats' corneas subjected to UVB-irradiation.

**Materials and methods:** Ten rats were used as donors for BM-MSCs and the other 40 were allocated into 4 equal groups. Group I (control group), Group II (ultraviolet irradiated group), Group III (ultraviolet irradiated + IV BM-MSCs treated group) and Group IV (ultraviolet irradiated + SC BM-MSCs treated group). Rats of all groups were euthanized after 3 weeks and the corneal specimens were processed for histopathological, immunohistochemical and electron microscopy assessment.

**Results:** ultraviolet irradiated group showed remarkable thinning of epithelial thickness, wide partial epithelial separation, and desquamation. Neovascularization of the disorganized stroma and disrupted Descemet’s membrane. The superficial and basal epithelial cells appeared irregular and separated by wide intercellular spaces and inflammatory cells. Immunohistochemical examination showed a significant decrease in PCNA immunoreaction. In contrast, minimal changes were observed in rats treated with BM-MSCs with more improvement associated with the subconjunctival administration compared to intravenous route.

**Conclusions:** Local SC injection of BM-MSCs has an amazing regenerative efficacy on the corneal injury compared to the systemic IV route.

**Key words:** ultraviolet radiation, corneal injury, mesenchymal stem cells, rats

**INTRODUCTION**

Annually, more than 1.5 million recent patients are presented with corneal blindness [28]. Because of the high cost and lack of the donors, corneal transplantations used as the
treatment of only less than 5% of cases [18]. Scarring of the cornea and thinning of its stroma are the expected fate of corneal injuries, which may result in visual impairment [40].

Ultraviolet radiation (UVR) is a type of the electromagnetic radiation that found in X-rays and visible light. Generally, UVR is divided into ultraviolet A (UVA) accounts for 90% to 95% with a wavelength 320–400 nm, and ultraviolet B (UVB) accounts for 5% to 10% with a wavelength 290–320 nm. UVB is likely more harmful than UVA as it possesses an extra energy and more capacity to destroy tissue [14, 24].

Cornea and anterior segment of the eye absorb most of UVB radiation, so the inner eye segments are preserved from the damage of radiation. Nearly 90% of UVB radiation are absorbed by the cornea which has greater sensitivity to UVB injury [33]. Exposure to toxic dose of UVB radiation may induce corneal epithelial injury, oedema, photophobia and photokeratitis [3, 15]. Corneal epithelial injury is composed of diminished viability of cells with degeneration of mitochondria and nuclei with a large number of apoptotic cells [63]. The cellular mechanism of injury may be due to antioxidant/prooxidant imbalance leading to oxidative stress [10]. Inadequately cleaved reactive oxygen species may lead to enhanced secretion of the proteolytic enzymes with significant activation of proinflammatory cytokines, causing creation of exaggerated inflammation, neovascularization and apoptosis of the cornea in addition to alterations in the optics of the cornea [8]. Furthermore, injury of the endothelium of the cornea is the consequence of exposition to toxic doses of UVB resulting in blurring of vision [9]. Many causes are related photokeratitis such as welder’s arc burns, snow blindness, lamps of metal halide, sunlamps and laboratory or germicidal ultraviolet lamps [17].

Mesenchymal stem cells (MSCs) are pluripotent undifferentiated cells that can proliferate and differentiate into various mature cells adipocytes, osteocytes and chondrocytes [57]. MSCs induce cellular differentiation leading to promotion of wound healing [51]. Furthermore, MSCs has an attractive powerful ability for self-renewal as well as restricted immunogenicity [46, 58], which encourage many researchers to study of the efficacy of MSCs to minimize inflammation and reconstruct the transparency of the cornea
after its damage [16, 44, 61]. Bone marrow mesenchymal stem cells (BM-MSCs) greatly ameliorate the injured corneal surface and accelerate its wound healing after alkali burn [42, 62].

The aim of this study was to compare the potential healing efficacy of two methods of injection of intravenous (IV) versus subcutaneous (SC) BM-MSCs on UVB-irradiated corneas in adult male albino rats.

MATERIALS AND METHODS

Animals

All experimental procedures and animal maintenance were conducted in accordance with the roles and the guidelines of the Research Ethics Committee, Menoufia University, Faculty of Medicine with the ethical approval number (ANAT- 8/2020- 1). A total of fifty Sprague-Dawley male albino rats were utilized during this study (aged 3-4 weeks and weighing 230-260 gm). They were purchased from the animal house of Faculty of Medicine, Ain Shams University, Cairo, Egypt and housed in the laboratory animal house unit of Kasr Al-Ainy, Faculty of Medicine, Cairo University, Cairo, Egypt. Two weeks before the experiment the rats were left for acclimation, kept in a monitored climate and exposed to a 12-hour light/dark cycle, with free access to food and tap water. Ten of the rats were used as donors for BM-MSCs and the other 40 were used for the experimental procedures.

Experimental design

The animals were allocated into 4 equal groups (n =10 per group):

— **Group I (control group):** which was equally divided into; subgroup A: the rats received IV injection of 0.1 ml phosphate-buffered saline (PBS). Subgroup B: the rats received SC injection of 0.1 ml PBS.
— **Group II (ultraviolet irradiated group):** the left eyeballs of the rats were exposed to a narrow beam of UVR-B for five consecutive days.
— **Group III (ultraviolet irradiated + IV BM-MSCs treated group):** one week after the last session of ultraviolet exposure, the rats received intravenous injection of 0.1 ml
PBS containing $1 \times 10^6$ MSCs in the lateral tail vein using a 25-gauge needle slowly over 1 min [41].

— **Group IV (ultraviolet irradiated + SC BM-MSCs treated group):** the rats received local anaesthesia by anaesthetic eye drops and injection of 10% chloral hydrate (4 ml/kg) then under an ophthalmic microscope, they received a single subconjunctival injection of 0.1 ml PBS containing $1 \times 10^6$ BM-MSCs one week after the last session of ultraviolet irradiation [19].

The rats of all groups were euthanized 26 days from the start of the experiment and half the eyeballs were enucleated then were submersed in 10% neutral buffered formalin solution and the other half were put in 2.5% glutaraldehyde for 24 hours. Thereafter the corneas were carefully dissected and processed for light microscopic and electron microscopic study.

**Induction of corneal injury by ultraviolet exposure**

The rats were anesthetized 10 minutes before the procedure by intraperitoneal injection of a mix of 2% Xylazine (10 mg/kg/ body weight) and 10% Ketamine (60 mg/kg/body weight). LS-1 tungsten halogen lamp with a spectral range 290–400 nm (supplying a 6.5 W output power) was provided as a pump for the source for a narrow beam of UVR-B which applied to the left cornea for 15 minutes per session [22].

**Isolation of BM-MSCs**

The donors’ rats were sacrificed by cervical dislocation and their femoral and tibial bones were harvested under sterilized condition. BM-MSCs were isolated based on the following protocols [55]. In brief, the marrow cavity was flushed with PBS (Lonza, Belgium) in a 1-ml syringe with an 18-gauge needle. After collection of the cells by centrifugation at 400 × g for 10 minutes, the BM-MSCs were resuspended at a density of 107 cells/ml in α-MEM Hank's medium. The cell pellet was suspended and plated into 25 cm$^2$ culture plates (3 ml/plate) (Sigma-Aldrich, St. Louis, MO, USA). Isolated BM-MSCs were cultured with adding 10% fetal bovine serum (FBS), 0.5% penicillin/ streptomycin and incubated at 37°C
and 5% CO2. The medium was changed on the day 3 and the non-adherent cells were discarded. The medium was changed every 48 hours.

**Passaging of BM-MSCs**

The primary cultured cells were kept in culture until reaching 80-90% confluence. Afterwards, the cells were trypsinized at 37°C by adding trypsin-EDTA (Gibco, South America) for 10 minutes. Thereafter, the action of trypsin-EDTA was stopped by addition of 5 ml of complete medium (CM). Furthermore, each cell pellet suspension was centrifugated and resuspended in fresh CM then distributed into 2-3 Petri dishes after counting the number of cells by the haemocytometer. Subculture of BM-MSCs repeated till reaching the 3rd passage.

**Viability testing**

Trypan blue exclusion assay was performed to test the cell survival in the culture before transplantation. The viable cells appeared with transparent clear cytoplasm while the dead cells had blue cytoplasm. The viability was estimated to be greater 95% through the different passages was excellent [56].

**Stem cell characterization and differentiation in vitro**

MSCs were demonstrated and examined by the inverted microscope (Leica, Wetzlar, Germany) which revealed their characteristic shape (fusiform spindle fibroblast like cells), adherence capacity and the ability for multipotent differentiation into osteocytes, adipocytes and chondrocytes. Differentiation to osteocytes was recognized by using osteocytes StemPro® osteogenesis differentiation kit (Gibco, Grand Island, NY, USA) and stained with Alizarin Red S stain (Sigma- Aldrich). The adipocytes differentiation was recognized by using adipocytes StemPro® adipogenesis differentiation kit (Gibco, Grand Island, NY, USA) and stained with Oil Red O stain (Sigma-Aldrich, St Louis, MO, USA). The chondrocytes differentiation was recognized by using chondrocytes StemPro® chondrogenesis differentiation kit (Gibco, Grand Island, NY, USA) and stained with Alcian Blue stain (Sigma- Aldrich) [7].

**Flow cytometric analysis**
For studying the phenotypic characters of BM-MSCs, the cells of the 3rd passage were detached by adding 0.25% trypsin. Thereafter, PBS was applied for washing of cells. Mesenchymal stem cell markers CD 90 and CD 105 and the hematopoietic stem cell marker CD 45 were detected in the surface the cultured cells. This was carried out by using Fluorescence-Activated Cell Sorting (FACS) (BD Biosciences, Franklin Lanes, NJ, USA) in which the cells were incubated at 4°C in the dark for 20 minutes with 2 μl fluorescein isothiocyanate (FITC)-conjugated rat monoclonal antibody against mouse CD90 and CD105 (BD Biosciences) and 5 μl phycoerythrin (PE)-conjugated rat monoclonal antibody against mouse CD45 (BD Biosciences). The samples were centrifuged at room temperature for 5 minutes at 300 × g, resuspended in 500 μl PBS then analysed by the flow cytometry [7].

**Immunofluorescence cell labelling**

According to the manufacturer's protocol, BM-MSCs at the 2nd passage were labelled with the red fluorescent dye PKH26 (sigma Aldrich). In summary, serum-free medium was utilized to wash BM-MSCs then the cells were resuspended in 1 ml of dilution buffer. An equal mixture volume of BM-MSCs suspension and the labelling solution containing 1×10⁶ PKH26 were incubated at room temperature for 5 minutes. The activity of the mixture was blocked by addition of 2 ml fetal bovine serum (FBS), finally, Dulbecco's modified Eagle's medium (DMEM)/F12 cells was applied three times for multiple cells washing, then the cells were injected intravenously into rat tail vein. The corneas were examined with a fluorescence microscopy to detect and trace the transplanted cells stained with PKH26 [38].

**Histopathological study**

The corneas were immersed in 10% neutral formalin (Sigma-Aldrich St. Louis, MO, USA) and processed for paraffin sections. 5μm thick sections were prepared and stained with Hematoxylin and eosin (H&E) stains, other sections were stained with periodic acid Schiff stain (PAS) for examination of the Descemet’s membrane [4]. All sections were examined under an Olympus DP70 light microscope (Olympus, Tokyo, Japan).

**Immunohistochemistry**
Paraffin sections were deparaffinized in xylene and hydrated. Sections were incubated with primary antibodies after stopping the reaction of peroxidase by adding 10% hydrogen peroxide. Proliferating cell nuclear antigen (PCNA) antibody is a mouse monoclonal antibody at 1:400 dilution (Sigma-Aldrich, St Louis, Missouri, USA). Antigen retrieval was applied for 30 minutes by steam heating the slides in a 1-mmol/L solution of EDTA (pH 8.0). After stopping of endogenous biotin reaction, staining was carried out by an automated immunostainer then by applying a streptavidin-biotin detection system (Dako, Glostrup, Denmark). Sections were counterstained with hematoxylin. Primary antibody (as a negative control) was changed by PBS [29].

**Transmission electron microscopy (TEM)**

Small corneal specimens were immersed for 2 hrs in 2.5 % buffered glutaraldehyde in 0.1 PBS at pH 7.4 and 4°C, then replaced by 1% osmium tetroxide in 0.1 mol/L for 1 hr at 4°C to fix the specimens. After that, specimens were dehydrated in ascending serial dilution of ethanol (50, 70, 90, 95 and four times 100%, each for 30 min) then dehydrated by acetone for 30 minutes. Finally, the fixed specimens were embedded in epoxy resin (Epoxy Embedding Medium Kit; Sigma). Semi- and ultra-thin sections were cut on ultramicrotome (RMC PT-XL PowerTome Ultramicrotome). Semithin (1μm) thick sections were prepared and stained with 1% toluidine blue and examined by using an Olympus BX61 light microscope. Ultrathin sections (70-90 nm) were cut by ultramicrotome then stained with 2% uranyl acetate as a principal stain and lead citrate as counter stain to be examined by using JEM-1400 Plus (JEOL, Japan) transmission electron microscope at the Electron Microscope Unit, Faculty of Science, Alexandria University, Alexandria, Egypt [30].

**Morphometric analysis**

All morphometric measurements were performed at a magnification of ×400 by the ImageJ® (1.50e, USA) software using at least five sections per animal, and from each section five non overlapping fields were examined at Anatomy Department, Faculty of Medicine, Menoufia University by (‘Olympus BX40, Ships to USA). All measurements were carried out by the same investigator to avoid inter-observer errors.
The H&E-stained sections were used to measure and estimate the corneal epithelial thickness then the results were recorded. The PAS-stained sections were utilized to investigate and measure the thickness of the Descemet’s membrane and the color intensity of PAS positive reaction then the means were calculated and reported. Furthermore, the mean area percent of PCNA was measured in the immune-stained sections in all studied groups.

**Statistical analysis**

The analysis of data was conducted with Statistical Package for the Social Science (SPSS) 18.0. One-way Analysis of Variance (ANOVA) followed by Tukey’s post-hoc test was performed to compare among the studied groups. P-values <0.05 were regarded as a statistically significant.

**RESULTS**

**Morphology of isolated BM-MSCs**

On day 0 post-seeding, the cultured cells appeared rounded with different size while some of them were aggregated into small groups and floated (Fig. 1A). One week after plating, MSCs were long, spindle shaped and colony forming units (Fig. 1B). On day 10, the number of cells was greatly increased and they acquired fusiform shape (Fig. 1C). Fourteen days after plating, the cell population appeared fibroblast like, and became confluent (Fig. 1D).

**Stem cell characterization and differentiation in vitro**

MSCs were differentiated to osteocytes stained with Alizarin Red S stain (Fig. 2A), differentiated to adipocytes stained with Oil Red O stain (Fig. 2B) and to chondrocytes stained with Alcian Blue stain (Fig. 2C).

**Phenotypic analysis of isolated BM-MSCs**

The surface markers of the isolated cells of the 3rd passage were analyzed by flow cytometry. Most of the cells expressed the mesenchymal stem cell markers CD 90 (99.9%) and
CD 105 (90.38%) while very few cells expressed the haematopoietic stem cell marker CD 45 (0.04%) (Fig. 3). This proved that the isolated MSCs were purified BM-MSCs.

**Immunofluorescence cell labelling**

By using Carl Zeiss LSM 710 Duo Confocal microscope, PKH26- labelled cells were traced after one week of their intravenous injection of the group III. Many labelled cells were seen scattered in the corneal tissue (Fig. 4).

**Histopathological findings**

Regarding H&E stain, the control subgroups showed a normal structure of the cornea. The stratified squamous non-keratinized epithelium was organized into five to seven cell layers resting on a uniform basement Bowman's membrane. Bowman's membrane was a thin acidophilic membrane which separated the corneal epithelium from the underlying corneal stroma. The layer of corneal stroma forming the main bulk of corneal thickness and consisted of avascular regularly well-organized bundles of parallel collagen fibers with scattered spindle-shaped fibroblast (keratocytes). Descemet’s membrane appeared as a homogenous and acidophilic thin layer below the stroma and was covered by a single layer of simple squamous Descemet’s endothelium (Fig. 5). On the other hand, the ultraviolet irradiated group showed remarkable histological alterations. The epithelium was mostly affected by wide partial separation and desquamation from the underlying stroma. Some epithelial cells showed pale vacuolated cytoplasm and deeply stained nucleus (Fig. 6A, 6B and 6C). Moreover, remarkable thinning of the corneal epithelial thickness was observed in this group compared to the control one. The collagen bundles of the stroma appeared thin and extensively disorganized with wide spaces among their fibers (Fig. 6B and 6C). Furthermore, neovascularization and mononuclear cellular infiltration were also observed in the stroma (Fig. 6B). Descemet’s membrane and its endothelial cells covering appeared highly disorganized and widely separated from the overlying stroma (Fig. 6C). Whereas, in the IV BM-MSC treated group, revealed moderate improvement of the corneal tissue in the form of restoration of organization and most of the thickness of the corneal epithelium. Vacuolated cytoplasm is still observed in some of the epithelial cells. Collagen bundles in the stroma are separated by many wide spaces. The
Descemet’s membrane and endothelium restored most of their normal features (Fig. 7A). The SC BM-MSC treated group showed marked improvement and restoration of many of the normal corneal tissue characteristics. The corneal epithelium was nearly similar to that of the control group with uniform normal thickness, rested on intact basement membrane with absence of epithelial separation or desquamation from the underlying stroma. The stroma showed regular organized collagen bundles and spindle-shaped keratocytes. The Descemet’s membrane appeared acidophilic with intact covering Descemet’s endothelium (Fig. 7B).

Regarding PAS stain, the control subgroups showed a regular Descemet’s membrane with a strong positive PAS reaction (Fig. 8A). On contrast, the ultraviolet irradiated group exhibited a thin irregular and wavy Descemet’s membrane with a weak PAS positive reaction (Fig. 8B). In IV BM-MSCs treated group the Descemet’s membrane showed partial restoration of normal appearance with a moderate positive PAS reaction (Fig. 8C). The SC BM-MSCs treated group demonstrated a regular thick Descemet’s membrane with a strong PAS positive reaction (Fig. 8D).

**Immunohistochemical results**

PCNA nuclear immunoexpression was applied to estimate corneal epithelial proliferation after MSCs injection. PCNA nuclear expression was mild in control group (Fig. 9A) and minimal in the ultraviolet irradiated group (Fig. 9B). On the other hand, IV BM-MSC treated group exhibited a moderate PCNA nuclear expression (Fig. 9C) and SC BM-MSC treated group displayed a strong PCNA nuclear expression (Fig. 9D). PCNA was expressed in the basal epithelial layer in control and ultraviolet irradiated groups, on contrast; the expression extended to the central and peripheral layers of the regenerated corneal epithelium in both IV BM-MSC and SC BM-MSC treated groups.

**Electron microscopic findings**

Ultrathin sections of the cornea of the control subgroups exhibited squamous cells in the superficial layer of the epithelial cells. Their nuclei appeared flattened and euchromatic. Intact desmosomes connected the adjacent cells together (Fig. 10A). The cells of intermediate layer showed normal nuclei with regular outline, mitochondria and rough endoplasmic
reticulum (RER) appeared normal. Junctional complex was intact (Fig. 10B). The basal layer consisted of tall columnar cells with regular euchromatic nuclei and mitochondria inside their cytoplasm. These basal cells were resting on an intact continuous basement membrane and connected to each other by many intact regular electron-dense desmosomes (Fig. 10C). The stroma showed the presence of a well-organized collagen fibers. Spindle shaped keratocytes with a euchromatic nucleus and little cytoplasm were observed in-between the collagen fibrils (Fig. 10D). A thick homogenous noncellular Descemet’s membrane was clearly demonstrated underneath the stroma and lined by endothelial cell layer. The endothelial cells possessed electron-dense flattened euchromatic nucleus and scanty cytoplasm with multiple small pinocytotic vesicles (Fig. 10E).

Ultraviolet irradiated group showed that the nucleus of squamous cells in the superficial layer of the epithelial cells appeared irregular and swollen. The cells were separated by wide intercellular spaces. Multiple Inflammatory cells were observed infiltrating the intercellular tissue (Fig. 11A). Cells of intermediate layer appeared with indented and irregular nucleus. Loss of junctional complex represented as wide intercellular spaces were clearly seen. Mitochondria showed complete lysis of their cristae. (Fig. 11B). The cells of the basal layer were disturbed, disorganized and irregular in shape. Some of these cells had an irregular nucleus while others possessed a shrunken heterochromatic pyknotic nucleus. The cells were separated by wide intercellular spaces and inflammatory cells. The basement membrane was seen irregular and interrupted (Fig. 11C). The stroma composed of irregular disturbed collagen fibrils. The keratocytes appeared irregular in shape with heterochromatic nucleus and vacuolated cytoplasm. The stroma displayed a very wide area devoid of collagen fibrils (Fig. 11D). The Descemet’s membrane appeared nonhomogeneous and distorted. The endothelial cells had an irregular nucleus with many variable sized vacuolations in the cytoplasm. The endothelial cells showed partial separation from the Descemet’s membrane (Fig. 11E).

On the other hand, the IV BM-MSCs treated group showed moderate improvement in all layers of the cornea. The squamous cells of superficial epithelial layer had an elongated euchromatic nucleus. Cells were connected together with intact continuous desmosomes;
however, some cells were still separated by intercellular spaces (Fig. 12A). Cells of intermediate layer showed irregular nucleus. Mitochondria was observed with partial lysis of their cristae. Desmosomes appeared intact. There was a slight widening in the intercellular spaces (Fig. 12B). Some cells of the basal layer had an irregular euchromatic nuclei with multiple mitochondria while others had a pyknotic nucleus. The cells were resting on a slightly irregular basement membrane. Desmosomes appeared connecting the cells with each other (Fig. 12C). The stroma showed well-organized regular collagen fibrils. Some keratocytes were irregular in shape with distorted nuclei and surrounded by a large space devoid of collagen fibrils (Fig. 12D). The Descemet’s membrane appeared thick, homogenous with a regular contour. The endothelial cells were distorted with many vacuolations in their cytoplasm (Fig. 12E).

The SC BM-MSCs treated group revealed marked improvement and restoration of the normal features in all layers of the cornea. The squamous cells in the superficial epithelial layer had a flattened elongated euchromatic nucleus. The cells were attached together by desmosomes. Narrow intercellular spaces were still seen among the cells (Fig. 13A). Cells of intermediate showed nearly normal nuclei appear with regular outline. Mitochondria and RER were normal. The junctional complex showed focal areas of slightly widening in the intercellular space while the desmosomes were intact (Fig. 13B). The cells of basal layer possessed normal contour with an euchromatic mild indented nuclei with many mitochondria resting on a regular intact continuous basement membrane. Intact desmosomes connected the epithelial cells to each other (Fig. 13C). The stroma showed well-organized regular collagen fibrils. A spindle shaped keratocytes with a euchromatic nucleus appeared in-between the collagen fibrils (Fig. 13D). The Descemet’s membrane appeared homogenous and noncellular similar to the control one and its lining endothelial cells had an elongated euchromatic nucleus (Fig. 13E).

**Morphometric assessment**

The mean thickness of the corneal epithelium showed a highly significant decrease in the ultraviolet irradiated group compared to the control one. Moreover, IV and SC injections in
the BM-MSCs groups showed a non-significant decrease compared to the control group, On the contrary showed a significant increase compared to the ultraviolet irradiated one (Table 1).

The mean color intensity of PAS positive reaction and the mean thickness of the Descemet’s membrane revealed a highly significant decrease in the ultraviolet irradiated group compared to the control one. SC BM-MSCs group showed a highly significant increase compared to the ultraviolet irradiated and IV BM-MSCs groups while a non-significant difference was recorded between SC BM-MSCs and control one (Table 2).

The mean area percent of PCNA for all groups was represented in (Table 3). There was a highly significant decrease in mean area percent of PCNA immunoexpression in the ultraviolet irradiated group compared to the control group. On the other hand, there was a significant increase in SC BM-MSCs group when compared to the other three groups.

DISCUSSION

The current study exhibited that BM-MSCs has a regenerative, anti-apoptotic and anti-inflammatory effects on the cornea exposed to ultraviolet radiation. This was assessed by histopathological, immunohistochemical and TEM evaluation. Local SC administration of BM-MSCs showed a better amelioration more than systemic IV administration on the corneal damage induced by UVB.

The cornea is avascular transparent structure, which permits the passage of light to the profound parts of the eyeball and is permanently at a high risk of exposure to a broad spectrum of radiation involving UV type [6]. Previous studies mentioned that the extent of injury resulted from UV radiation exposure is based on numerous factors, like the time of exposure and the wavelength of radiation. Wavelengths under 290 nm are totally absorbed by the epithelial layer of the cornea then pass to the deep located structures, whereas those in the range between 300–320 nm are absorbed by the cornea and crystalline stroma, resulting in injury to several structures [50].

In this study, repeated application of UV radiation in the wavelength range between 290–400 nm for successive five days leads to epithelial desquamation of the cornea. Some
epithelial cells revealed deeply stained nucleus and pale vacuolated cytoplasm. Furthermore, the thickness of corneal epithelium was markedly decreased in some focal areas. The stroma shower disturbed collagen fibers separated by wide spaces, neovascularization and inflammatory infiltration. The Descemet’s membrane seemed to be disorganized with some endothelial cells were disturbed with pale vacuolated cytoplasm. Current results were in accordance with Golu et al. [22], who concluded that repeated exposure to UV radiation with 290–400 nm wavelength resulted in necrosis of the membrane of the basal epithelium that may extend deeper to erode the Bowman’s membrane. The Bowman’s membrane was seen detached from the epithelium with fluid accumulation and damage of the junction in-between them. The Bowman’s membrane was seen thickened, wavy and interrupted, permitting lymphocytes and macrophages migration in the anterior epithelium.

Similar findings have been reported by Chen et al. [12] who conducted a study on the mice which subjected to UVB light (0.72 J/cm2 /daily), and Mahmoud et al. [43] who applied a single UVB irradiation at a dose of 1.2 J/cm2 on the cornea of the rats. The corneal epithelial thickness after UVB exposure appeared extremely thinner relative to the control group. The nuclei of the epithelial cells appeared more condensed, indicating cell death. UVB group considerably showed exfoliation of the corneal epithelium. Discontinuation or may be absence of the Bowman’s membrane was clearly seen. Furthermore, the stroma revealed disarranged collagen fibers which separated by wide spaces, keratocytes were degenerated were infiltrated by polymorphonuclear leukocytes, multiple lymphocytes, macrophages and hemorrhage [22, 43].

The current work revealed degenerated superficial epithelial cells with irregular basal cells, wide separation of corneal stroma and interruption of Descemet’s membrane by electron microscopic examination. These findings were consistent with the ultrastructural study conducted by Mahmoud et al. [43] who showed irregular nuclear membrane, condensation of chromatin, absent rough endoplasmic reticulum and mitochondrial dissolution of the epithelial cells of the cornea.
Previous studies have reported many alterations of the corneal structure were induced by UV radiation by provocation of reactive oxygen species. Exposure of the cornea once to UV-B radiation was enough to terminate endothelial cell proliferation. Marked decrease in the epithelial thickness was associated with higher doses of UV-B [10]. Exposure to UV is associated with reduction of the oxygen removal rate from the cornea and blockage of glucose transmission system inside the cornea [34, 35]. Injury of the anterior corneal epithelium and endothelium is associated with UV exposure [48]. These results are matching with data of the present study which declare the presence of detachment in the layers of anterior epithelium and the endothelium after UV radiation exposure.

In the recent years, the cellular and molecular events inducing photokeratitis have been greatly investigated [2, 32]. A variety of proinflammatory molecules including cytokines, nuclear factor-κB (NF-κB), interleukins and matrix metalloproteinases (MMPs) are involved in the disease progression [21, 37, 49, 59]. NF-κB activation was generally reported to be provoked by UVB. The transcription of several down-stream genes such as inducible nitric oxide synthase (iNOS) and cyclooxygenase-2 (COX-2) is induced by NF-κB. Both factors are the main activators of the induction of inflammatory cells [12, 23, 27, 39].

In the current work, the stroma was infiltrated by inflammatory cells. Wang et al. [60] reported that the damaged epithelial cells release interleukin 1β (IL1β) which is responsible for inflammatory cellular reaction in the stroma. IL1β is an active cytokine that is implicated in immediate inflammatory reaction resulting in activation and infiltration by neutrophils and macrophages.

In the present study, neovascularization of the corneal stroma was detected in the irradiated group. This was in accordance with Golu et al. [22] and Mureşan et al. [45] who reported the appearance of stromal neovascularization of corneal stroma of the rat following prolonged ultraviolet exposure. Excess of pro-angiogenic agents, like vascular endothelial growth factor (VEGF), MMPs and fibroblast growth factor (bFGF) and lack in antiangiogenic agents such as endostatin, angiostatin and pigment epithelium–derived factor resulted in imbalance between angiogenic and antiangiogenic factors which stimulate the avascular cornea to be vascularized [11]. VEGF were released from the basal layer epithelial cells,
endothelial cells and fibroblasts of the cornea [49]. On the other hand, Clements and Dana [13] reported that the neovascularization may occur due to inflammation.

In current study, BM-MSCs groups showed improvement in all parameters. These findings are in agreement with Harkin et al. [25] who emphasized that the healing of epithelium was improved by BM-MSCs. There are different mechanisms by which the MSCs repaired the corneal damages such as growth factors and cytokines excretions that controlled cell proliferation and signaling after corneal alkali burn. Other study found that BM-MSCs-treated damaged corneas exhibited a marked amelioration and closure of the defects in the epithelium resulting in normal well-organized layers of epithelium with no intercellular gaps relative to the untreated corneas. Furthermore, the organized stromal structure is preserved by important factors such as keratan-sulfate proteoglycans which is released by the regenerated keratocytes. They are crucial and essentially needed for the transparency of the cornea by production and maintenance of normal collagen interfibrillar spacing and the fibril width [26].

The present results are in matching with Giacomini et al. [20] and Lazarus et al. [36] who demonstrated that BM-MSCs protected and accelerated healing of the corneal epithelium and endothelium resulting from free oxygen radicals and their oxidative stress via its immunomodulatory function through inhibition of leukocytes chemotaxis and phagocytosis of macrophages and suppression of release of free radicals. Previous researches conducted by Ma et al. [42] and Oh et al. [47] stated that promoted wound healing and ameliorated regeneration of the corneal surface were improved by MSCs therapy through its anti-inflammatory properties. Ma et al. [42] hypothesized that suppression of CD45, interleukin 2 (IL-2) and MMP2 expression was produced due to the anti-inflammatory efficacy of MSCs. On the other hand, Oh et al. [47] reported that the anti-inflammatory properties of MSCs was achieved by blocking the adaptive CD4+ T cells invasion and also through suppression of CD4+ T cell-related cytokines and MMP2 expression potentially by releasing soluble agents.

In the current work, a highly significant decrease in PAS positive reaction in ultraviolet treated group compared to the control was recorded, while statistically significant increase was recorded between MSCs treated groups compared to ultraviolet treated group. These findings are consistent with Sharaf Eldin et al. [52] who registered that after exposure to alkali burn,
the Descemet’s membrane of the cornea of the rat showed a mild positive reaction to PAS stain. On contrast, Ahmed et al. [1] recorded that after exposure to alkali burn the Descemet’s membrane of the rabbit’s cornea exhibited a significant increase in the thickness of the PAS positive reaction relative to control group. They concluded that the reduction in efficiency of the barrier in addition to a drop in the pump function resulting from variation in pH of the endothelium of the cornea is the main cause of the increase in the thickness of Descemet’s membrane.

In this study, PCNA was highly expressed in basal epithelium and extended to the central and peripheral layers of the regenerated corneal epithelium in both IV and SC BM-MSCs treated groups. In agreement with these finding, Ye et al. [62] mentioned that during the healing of corneal alkali burn, the efficacy of stem cell and the proliferation of cell kinetics were achieved by using stable cell cycle-regulated protein (PCNA). Expression of PCNA reaction is an indicator that MSCs promote the proliferation of limbal stem cells. This data suggested that the MSCs moved to the damaged cornea and induced native corneal cell proliferation. The probable interpretation of this suggestion is that many limbal stem cells were promoted by the transplanted MSCs in cornea; these stimulated stem cells react to the local specific medium produced by inflammatory reaction by secreting cytokines from MSCs, acquiring the ability to proliferate and differentiate. Thus, healing of the wound could be promoted by the transplanted MSCs through activating and synchronizing with native cells.

The corneas of IV BM-MSCs treated group are still suffered from some residual histological alterations while SC BM-MSCs treated group displayed corneas with nearly normal features. These findings are consistent with Shukla et al. [53] who assessed the effect of various routes of MSCs administration in amelioration of reconstruction and maintaining the transparency of cornea after inducing corneal damage in a murine model. They reported that SC and IV injections of MSC after corneal damage are successful in dramatically reducing the inflammation and opacification in addition to repair and re-epithelialization of the corneal tissue, however; there was slightly more improvement associated with the SC administration compared to IV route. The only interpretation is that SC route evades the MSCs aggregation of ‘first-pass’ in the lungs and decreases the possibility for residing outside the target and
immunomodulatory impacts [54]. As the IV MSCs were retained in the lungs, huge numbers of cells are required to achieve the needed therapeutic effect, a consideration that decreases the possibility of using MSC as a therapy in the clinical purposes [5, 31, 64]. On the other hand, administration of SC MSCs allowed the whole transplanted cells to reach the target with bypass of the first-pass action in the lung.

CONCLUSIONS

In summary, the current study exhibited that the local SC injection of BM-MSCs has a marvelous regenerative efficacy on the corneal tissue compared to the systemic IV injection of BM-MSCs. This occurs through inhibition of neovascularization and suppression of the inflammatory reaction and degenerative changes of the epithelial cells induced by chronic ultraviolet irradiation exposure. Therefore, the usage of SC injection of BM-MSCs is recommended as a better therapeutic agent more than the IV injection in the chronic corneal injury.

REFERENCES


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**Table 1. Corneal epithelial thickness of the study groups**

<table>
<thead>
<tr>
<th>Group</th>
<th>Corneal epithelial thickness</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group (Group I)</td>
<td>41.8±0.45</td>
</tr>
<tr>
<td>Ultraviolet irradiated group (Group II)</td>
<td>24.6±0.54</td>
</tr>
<tr>
<td>IV BM-MSCs group (Group III)</td>
<td>40.2±0.44</td>
</tr>
<tr>
<td>SC BM-MSCs group (Group IV)</td>
<td>40.8±0.44</td>
</tr>
</tbody>
</table>

Values are mean±SD. \(^a^p <0.05\) vs. Group I and \(^b^p <0.05\) vs. group II. Statistical analysis was performed by ANOVA, followed by Tukey’s *post-hoc* test.
**Table 2.** Thickness of Descemet’s membrane and color intensity of PAS-positive reaction of the study groups

<table>
<thead>
<tr>
<th>Group</th>
<th>Thickness of Descemet’s membrane (µm)</th>
<th>Color intensity of PAS-positive reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group (Group I)</td>
<td>7.16 ± 0.18</td>
<td>21.36 ± 0.60</td>
</tr>
<tr>
<td>Ultraviolet irradiated group (Group II)</td>
<td>4.11 ± 0.43&lt;sup&gt;a&lt;/sup&gt;</td>
<td>11.49 ± 0.92&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>IV BM-MSCs group (Group III)</td>
<td>5.94 ± 0.46&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>17.95 ± 0.53&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
<tr>
<td>SC BM-MSCs group (Group IV)</td>
<td>7.07 ± 0.34&lt;sup&gt;b,c&lt;/sup&gt;</td>
<td>21.00 ± 0.84&lt;sup&gt;b,c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Values are mean±SD. <sup>a</sup>p<0.05 vs. group I, <sup>b</sup>p<0.05 vs. group II and <sup>c</sup>p<0.05 vs. group III.

Statistical analysis was performed by ANOVA, followed by Tukey’s post-hoc test. SD: Standard deviation; ANOVA: Analysis of variance.

**Table 3.** PCNA immunoexpression of the study groups

<table>
<thead>
<tr>
<th>Group</th>
<th>PCNA immunoexpression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group (Group I)</td>
<td>135.0±3.6</td>
</tr>
<tr>
<td>Ultraviolet irradiated group (Group II)</td>
<td>113.8±3.1&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>IV BM-MSCs group (Group III)</td>
<td>158.2±2.8&lt;sup&gt;a,b&lt;/sup&gt;</td>
</tr>
<tr>
<td>SC BM-MSCs group (Group IV)</td>
<td>176.0±2.6&lt;sup&gt;a,b,c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Values are mean±SD. <sup>a</sup>p<0.05 vs. Group I, <sup>b</sup>p<0.05 vs. group II and <sup>c</sup>p<0.05 vs. group III.

Statistical analysis was performed by ANOVA, followed by Tukey’s post-hoc test.

**Figure 1.** MSCs are characterized in culture A: On day 0 of MSCs isolation: cultured cells appear rounded with different size (arrow) while some of them are aggregated into small groups and floated (double arrow) (40x) B: On day 7, MSCs are long, spindle shaped (arrow) with colony forming units (double arrows) (40x) C: On day 10, the number of cells are greatly
increased and they acquire fusiform shape (arrow) (100x) D: On day 14, the cell population appear confluent fibroblast like cells (arrow) (200x).

**Figure 2.** A: MSCs are differentiated in culture into osteocytes and stained with Alizarin Red S stain B: MSCs are differentiated in culture into adipocytes and stained with Oil Red O stain C: MSCs are differentiated in culture into chondrocytes and stained with Alcian Blue stain.

**Figure 3.** A chart of the flow cytometric analysis of the surface antigens of MSCs: they are positive for CD90 (99.9%), CD105 (90.38%) and negative for CD45 (0.04%).

**Figure 4.** Immunofluorescence micrograph of the cornea of IV BM-MSCs treated group showing the presence of numerous PKH26 labeled cells in the corneal tissue (arrows) (scale bar = 50 μm).

**Figure 5.** A photomicrograph of a section of the cornea of the control group showing normal corneal tissue consists of a nonkeratinized stratified squamous epithelium (E) which organized into multiple layers overlying Bowman's membrane (arrow). The stroma (S) constitutes most of the thickness of the cornea and consists of avascular well-organized collagen bundles with scattered spindle-shaped keratocytes (arrowhead). Descemet's membrane and single squamous Descemet's endothelial layer (D) are seen below the stroma. (H&E X 400 – scale bar = 20 μm).

**Figure 6.** Photomicrographs of sections of the cornea of the ultraviolet irradiated group showing A. Wide partial separation and desquamation of most of the epithelial layer of the cornea from the underlying stroma (curved arrow). Some epithelial cells showing vacuolated cytoplasm (arrows). B. Thinning of the corneal epithelial thickness (E). Some epithelial cells showing vacuolated cytoplasm (arrowhead). The collagen bundles of the stroma appear thin and extensively disorganized with wide spaces between its fibers (asterisks). Neovascularization (V) and mononuclear cellular infiltration (curved arrow) are also observed in the stroma. C. Many epithelial cells are distorted with deeply stained nuclei and vacuolated
cytoplasm (arrow heads). The stroma showing thin dispersed irregular collagen fibers with wide spaces (asterisks). The Descemet’s membrane and its covering endothelial cells appearing highly disorganized and widely separated from the overlying stroma (arrows). (H&E X 400 – scale bar = 20 μm).

**Figure 7.** Photomicrographs of sections of the cornea of **A.** IV BM-MSCs treated group showing restoration of organization and most of the thickness of the corneal epithelium (E). Vacuolated cytoplasm is still observed in some of the epithelial cells (arrow heads). Collagen bundles in the stroma are separated by many wide spaces (asterisks). The Descemet’s membrane and endothelium restored most of their normal features (arrow). **B.** SC BM-MSCs treated group showing uniform organization and normal thickness of the corneal epithelium (E). The stroma showing regular organized collagen bundles (S) and spindle-shaped keratocytes (arrowhead). The Descemet’s membrane appears acidophilic with intact covering Descemet’s endothelium (D). (H&E, × 400 – scale bar = 20 μm).

**Figure 8.** Photomicrographs of sections of the cornea of the studied groups. **A.** Control group showing a regular Descemet’s membrane with a strong positive PAS reaction (thin arrow). **B.** Ultraviolet irradiated group showing a wavy, irregular Descemet’s membrane with marked decrease in its thickness and weak PAS positive reaction (thick arrow) **C.** IV BM-MSCs treated group showing a Descemet’s membrane with a moderate positive PAS reaction (arrowhead). **D.** SC BM-MSCs treated group showing a regular thick Descemet’s membrane with a strong PAS positive reaction (double arrow). (PAS; ×400 – scale bar = 20 μm).

**Figure 9.** Photomicrographs of immunostaining sections of the cornea of the studied groups. **A.** Control group showing a mild nuclear immunoreactivity of the corneal epithelial cells. **B.** Ultraviolet irradiated group exhibiting a minimal nuclear immunoreactivity of the corneal epithelial cells. **C.** IV BM-MSCs treated group demonstrating a moderate nuclear immunoreactivity of the corneal epithelial cells. **D.** SC BM-MSCs treated group showing a strong nuclear immunoreactivity of the corneal epithelial cells. (PCNA immunostaining; ×400 – scale bar = 25 μm).
**Figure 10.** Electron micrographs of the cornea of the control group showing: **A.** Squamous cells are seen in the superficial layer of the epithelial cells. The nuclei of these cells appear flattened and euchromatic (N). Intact desmosomes connecting the adjacent cells together (arrows). **B.** Cells of intermediate layer of corneal epithelium. Nuclei appear normal with regular outline (N), mitochondria (m) and RER (arrowhead) are normal. Intact junctional complex (arrow). **C.** Columnar cells in the basal layer of the epithelial cells with euchromatic nuclei (N). The cells resting on an intact continuous basement membrane (arrow) and connected to each other by many intact regular electron-dense desmosomes (arrowhead). The mitochondria are seen inside its cytoplasm (m). Note: regular collagen bundles (C) surrounding a keratocyte (K) in the stroma. **D.** The stroma showing the presence of an organized arrangement of the collagen fibers (C). A spindle shaped keratocyte (K) with a euchromatic nucleus (N) and little cytoplasm. **E.** Thick homogenous noncellular Descemet’s membrane (D) is lined by endothelial cell (E) with electron-dense flattened euchromatic nucleus (N) and scanty cytoplasm containing multiple small pinocytotic vesicles (arrow). (TEM Fig. A, C, D, E; ×17500 – Scale bar = 1 μm; Fig. B; ×8000 – Scale bar = 4 μm).

**Figure 11.** Electron micrographs of the cornea of the ultraviolet irradiated group showing: **A.** Squamous cells of the superficial layer of the epithelial cells. The nucleus (N) appears irregular and swollen. The cells are separated by wide intercellular spaces (arrows). Multiple inflammatory cells (I) are observed infiltrating the intercellular tissue. **B.** Cells of intermediate layer with indented (arrowhead) and irregular nucleus (N), loss of junctional complex appears as wide intercellular spaces (arrows). Note, mitochondria (m) can be observed with complete lysis of their cristae. **C.** Disturbed disorganized irregular shaped cells are noticed in the basal layer. Some of these cells have irregular nucleus (N) while others possess a shrunken heterochromatic pyknotic nucleus (p). The cells are separated by wide intercellular spaces (arrows) and inflammatory cells (I). The basement membrane is seen irregular and interrupted (arrow heads). Note: inflammatory cell (i) is observed in the region of stroma (S). **D.** The stroma (S) containing irregular disturbed collagen fibrils. A keratocyte (K) appears irregular in shape with heterochromatic nucleus (N) and vacuolated cytoplasm (v). The stroma
exhibiting a very wide area devoid of collagen fibrils (asterisk). **E.** Descemet’s membrane (D) appears nonhomogeneous and distorted. The endothelial cell (E) has an irregular nucleus (N) with many variable sized vacuolations in the cytoplasm (v). The endothelial cell shows partial separation from the Descemet’s membrane. Note: The stroma showing distorted irregular collagen fibrils (asterisk). (TEM Fig. A-C; ×8000 – Scale bar = 4 μm; Fig. D, E; ×17500 – Scale bar = 1 μm).

**Figure 12.** Electron micrographs of the cornea of IV BM-MSCs treated group showing: **A.** Squamous cells of superficial epithelial layer has an elongated euchromatic nucleus (N). The cells are connected together with intact continuous desmosomes (curved arrows); however, some cells are still separated by medium sized intercellular spaces (arrows). **B.** Cells of intermediate layer with irregular nucleus (N), mitochondria (m) with partial lysis of their cristae, intact normal desmosomes can be observed (arrow), slightly widening in intercellular spaces (circle). **C.** Some basal layer cells have irregular euchromatic nucleus (N) with multiple mitochondria (m) while others have pyknotic nucleus (p). The cells are observed resting on a slightly irregular basement membrane (arrow). Desmosomes (curved arrows) could be seen connecting the cells with each other. The stroma shows a regular arrangement of the collagen fibrils (S). **D.** The stroma reveals well-organized regular collagen fibrils (C). A keratocyte (K) is observed with an irregular shape, distorted nucleus and surrounded by a large space devoid of collagen fibrils (asterisk). **E.** The Descemet’s membrane (D) appears thick, regular and homogenous in contour. Endothelial cell (E) is distorted with many vacuolations in its cytoplasm (V). (TEM Fig. A-C, E; ×8000 – Scale bar = 4 μm; Fig. D; ×17500 – Scale bar = 1 μm).

**Figure 13.** Electron micrographs of the cornea of SC BM-MSCs treated group (IV) showing marked improvement in all of its layers: **A.** Squamous cells in the superficial epithelial layer have a flattened elongated euchromatic nucleus (N). The cells are attached together by desmosomes (curved arrows). Narrow intercellular spaces are still seen among the cells (arrows). **B.** Cells of intermediate layer with nuclei appear nearly normal with regular outline (N). Mitochondria (m) and RER (arrowhead) are normal. The junctional complex shows focal
area of slightly widening in the intercellular space (circle) while the desmosomes are intact (arrow). C. The cells of basal layer possess a normal contour and an euchromatic mild indented nuclei (N) with many mitochondria (m) resting on a regular intact continuous basement membrane (arrow). Intact desmosomes connecting the epithelial cells to each other (curved arrows). D. The stroma is seen with well-organized regular collagen fibrils (C). A spindle shaped keratocyte (K) is observed with an euchromatic nucleus (N). (TEM; ×8000 – Scale bar = 4 μm). E. Homogenous noncellular Descemet’s membrane (D). The lining endothelial cell (E) has an elongated euchromatic nucleus (N). Note: the regular organization of collagen fibrils in the stroma (C). (TEM Fig. A, B, D, E; ×8000 – Scale bar = 4 µm; Fig. C; ×17500 – Scale bar = 1 µm).