Introduction

Gastric (peptic) ulcer is a major gastrointestinal disorder with high morbidity and mortality [1]. It occurs due to an imbalance among aggressive factors (such as acid-pepsin secretion), mucosal protection factors (like mucus and bicarbonate secretion), and the process of regeneration after cellular injury [2].

The most common causes of gastric ulcer are the use of non-steroidal anti-inflammatory drugs (NSAIDs), and tobacco smoking [3]. Gastric ulcers occur in 4% of the population mostly in elderly [4].

Individuals with gastric ulcer are at risk of developing complications such as gastric hemorrhage, perforation, and obstruction [5]. Globally about 15 patient among every 15,000 die each year from gastric ulcer complications [6, 7]. Several drugs have been used to treat gastric ulcer; how-
ever, the literature contains many reports of the association between long-term use of proton pump inhibitors and the adverse side effects such as bacterial overgrowth within the gastric mucosa and community acquired pneumonia [3]. Providing gastric protection by preventing NSAID-induced peptic ulcer disease is still considered a difficult issue [1, 8].

Gastric mucosal defense mechanisms include local defense mechanisms by the mucus bicarbonate barrier [9], and surface epithelial cells [10]. Another local defense mechanism is continuous cell renewal by mucosal progenitor cells to replace injured or aged surface epithelial cells completely within 3 to 7 days; and in glandular cells, it takes months [11]. Additional local defense mechanisms include alkaline tide, mucosal microcirculation [12], and continuous generation of prostaglandin E2 (PGE2) and prostaglandin I2 (PGI2) by the mucosa [13]. Neurohormonal regulation of the gastric mucosal defense is another important line of defense in which central corticotropin-releasing factor signaling pathways contribute to the endocrine and visceral responses to stress [14].

Gastric ulcers result from tissue sloughing initiated by mucosal ischemia, free radical formation and hindered nutrient access. They are caused by vascular and microvascular injury such as thrombi, constriction, or other occlusions [15]. Ulcer healing is an active and complicated process that involves re-epithelialization, granulation tissue formation, and angiogenesis. It requires the interaction of a variety of tissues and cellular systems as well as a number of growth factors as vascular endothelial growth factor (VEGF) and cytokines [16].

Mesenchymal stem cells (MSCs) are undifferentiated multipotent cells which are able to proliferate, self-renew, and differentiate into multiple cell lineages [17, 18]. Because of their unique capabilities as immunomodulatory, multipotent and fast proliferating cells, MSCs can be used for a wide range of treatments [3]. A cascade of events occurs when MSCs enter the microenvironment of injured tissues. Many factors including toxins of infectious agents, inflammatory stimuli, hypoxia and cytokines such as TNF-α, interleukin (IL)-1, interferon-γ stimulate MSCs to release many growth factors, such as EGF, FGF, PDGF, TGF-b, VEGF, HGF, insulin growth factor-1, angiopoietin-1, keratinocyte growth factor, and stromal cell derived factor-1 [19]. These growth factors enhance the development of fibroblasts, endothelial cells and tissue progenitor cells, which implement tissue replacement and repair [20]. It has been reported that trophic factors produced by MSCs can be used to achieve therapeutic effects to inflammatory diseases, whereas in damaged tissue engrafted MSCs can act in harmony with immune cells, stromal cells, endothelial cells and tissue progenitor cells to enhance tissue repair [21, 22].

In our study, we examined the role of MSCs in treatment of the gastric ulcer.

Materials and Methods

Experimental animals

A total number of 36 male albino rats (aged 8 weeks, weighing 150–200 g) were purchased from the Animal House of Assiut University. The animals were maintained under controlled laboratory conditions for humidity, temperature, and light /dark cycle and fed normal rat’ chow and allowed free access to water. All animal procedures used were approved by the Institutional Ethics Committee of Assiut University (17200549).

Drugs and chemicals

Indomethacin ampules (EL–NILE Company, Cairo, Egypt). VEGF antibody (Cat. NO NBP2-45235; Novus Biologicals, Centennial, CO, USA) and Proliferating cell nuclear antigen antibody (PCNA; Cat log #YPA1100; Biospes, Chongqing, China).

Isolation and culture of mesenchymal stem cells

Adipose tissue specimens from lipectomy procedures were obtained aseptically from the Plastic Surgery Department, Assiut University Hospital. The tissues were dissected into very small pieces; digested using collagenase solution, and transferred to a shaking water bath where they were incubated at 37°C for 1 hour. The solution was centrifuged and washed, sieved through nylon mesh, and cultured in complete media. On the second day, the cultured cells were examined using an inverted microscope. The cells underwent two passages before transplantation into the rats [23].

Experimental design

The animals were divided into 4 main groups. Group I, group II, group III , and Group IV. Group III and IV are divided into two subgroups. Each main and sub group contains 6 rats.

Group I (control group): the rats received isotonic saline. Group II (ulcer group): the rats fasted for 24 hours, with free
water intake, and then were given indomethacin 200 mg/kg orally [24]. After 4 hours, they were sacrificed [25]. Group III (autohealing groups) subdivided into Group IIIa and Group IIIb. Group IIIa: after ulcer induction, the rats were left to heal spontaneously without any intervention, they were dissected after the 1st day. Group IIIb: after ulcer induction, the rats were left to heal spontaneously without any intervention except for isotonic saline injection then dissected after the 4th day. Group IV (stem cells-treated groups) subdivided into Group IVa and Group IVb. Group IVa: after ulcer induction the rats were injected intraperitoneally with a single dose of MSCs (1×10⁶ cell) [26]. They were dissected after the 1st day. Group IVb: after ulcer induction the rats were injected intraperitoneally [27], with a single dose of MSCs (1×10⁶ cell) and were dissected after the 4th day. Normal animals received MSCs in a single dose (1×10⁶ cell) cells. After scarification and dissection after the 1st and 4th days, the stomachs were subjected to RNA extraction and subsequent steps taken to detect human leukocyte antigen (HLA) in the specimens.

**Macroscopic examination**

At the end of the experiment, the dissected stomachs were photographed using the dissecting microscope at the Histology Department to document the number and size of the ulcers and allow calculation of the ulcer index for the different groups. Ulcer index: The ulcer severity can be evaluated according to the scale using a modification of the score created by Kunchandy et al. [28] as follows: 0, Normal gray colored stomach; 0.5, Pink to red coloration of stomach; 1, Spot ulcer; 1.5, Hemorrhagic streak; 2, Number of ulcers less than 5; 3, Number of ulcer more than or equal to 5; 4, Ulcer with bleeding and 5; Perforation of gastric/duodenal wall. The ulcer index was calculated by adding the total number of ulcers plus the severity of each ulcer.

**Light microscopic examination**

*By hematoxylin and eosin stain*

Stomach sections from the six groups were fixed in 10% formalin, embedded in paraffin, cut into sections, 5 µm thick, and stained with Hematoxylin & Eosin (H&E) to assess the extent of injury (gastric ulcer) and detect evidence for the extent of healing [29].

**Immunohistochemical preparation for VEGF and PCNA antibodies**

Paraffin embedded sections were used for immunostaining. Briefly, hydrogen peroxidase was added to the sections for 10 minutes. Antigen retrieval was done by adding citrate buffer to the slides which were placed in a microwave for 9 minutes. Primary antibodies were added (1:200; VEGF), (1:100; PCNA) for the Group 1, 2, IIIa, IIIb, IVa, and IVb. The slides were incubated with antibodies at 4°C over night, followed by secondary antibody (Asnf Medical Company, Cairo, Egypt) for 15 minutes. DAB buffer was mixed with chromogen and added to the slides for 10 minutes. Slides were stained by H&E stain [30], mounted, covered and examined by light microscope.

**Flow cytometry analysis for mesenchymal stem cells**

Isolated adipose derived MSCs were analyzed using a fluorescence-activated cell sorting (FACS) cell analyzer at Southern Egypt Institute of Tumors, Assiut University [31]. The cells were harvested and incubated for 30 minutes on ice with 1% bovine serum albumin containing the primary antibodies directed against: CD90, CD45, CD73, CD271, and CD34 (Santa Cruz Biotechnology, Santa Cruz, CA, USA), after which the cells were washed and incubated for another 30 minutes away on ice from light with Alexa Fluor 647 (Santa Cruz Biotechnology). The cells were washed again.

**Gene expression of HLA in control & stem cells treated groups**

RNA extraction was carried out from stomach of MSCs injected groups (Group I and IV) on the 1st and 4th days using an RNase extraction kit according to manufacturer’s directions. All polymerase chain reaction (PCR) reagents were obtained from Applied Biosystems (Waltham, MA, USA) and examined using RT2SYBR® (Precision PLUS Master Mix premixed with SYBR green). Complementary DNA (cDNA) was synthesized from RNA using reverse transcriptase with random primers. The reaction mixture was then subjected to PCR amplification with specific forward and reverse oligonucleotide primers for 40 cycles. The samples were run on 1% agarose gel and photographed under UV light. Regarding qRT-PCR for quantitative expression of HLA; an aliquot of 1,000 ng of the total RNA from the sample was used for complementary cDNA synthesis by reverse transcription using a high capacity CDNA Reverse Transcriptase kit (CAT #4368814; Applied Biosystems). The cDNA was
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subsequently amplified with the SYBR Green I PCR Master Kit (CAT #K0251; Fermentas, Waltham, MA, USA) using the Step One instrument (Applied Biosystems), and the Precision PLUS Master Mix premixed with SYBR green. Samples were prepared for reverse transcription with the precision Nano Script 2 Reverse Transcription kit. The reactions were done using a thermal cycler (MJ Research, Inc., Waltham, MA, USA). The thermal profile consisted of 2 minutes at 50°C, 10 minutes at 95°C, followed by 40 cycles of 15 seconds at 95°C, and 1 minute at 60°C. Gene expression was normalized with the house keeping gene (GADPH) and reported as fold change in expressions. Primers used were as follows: HLA-DQB-10201: Forward primer GTGCGTCT TGTGACAGAAG, Reverse primer: GCAAGGT CGT GCGGAGCT (Size 205 bp). Glyceraldehyde 3 phosphatedehydrogenase (GAPDH):F: AAC TTT GGC ATT GTG GAA GG R: ACA CAT TGG GGG TAG GAA CA [32].

Morphometric studies
The numbers of ulcers were calculated grossly in all groups to assess the ulcer index.

Computerized image analyzer system software (Leica Q 500 MCO; Leica, Wetzlar, Germany) connected to a camera attached to a Leica universal microscope at the Histology Department, Faculty of Medicine, Assiut University was used. Six non overlapping fields in 5 randomly chosen PCNA immunostained sections from three different rats from Groups I, II, IIIa, IIIb, IVa, and IVb were used.

Statistical analysis
Statistical analysis was performed using the GraphPad Prism® software package, version 7 (GraphPad Prism 7 Software Inc., San Diego, CA, USA). Analysis of variance (ANOVA) followed by Tukey–Kramer multiple comparisons test was done for ulcer index and PCNA while unpaired Student’s t-test was used for statistical analysis of HLA expression. Data were considered significant at P<0.05. Data were expressed as mean±standard error of mean.

Results
The macroscopic examination
The opened dissected stomachs of rats from Group I showed that the mucosa of stomach the near esophagus (fore stomach) is whitish in color and the mucosa near the small intestine (glandular stomach) is greyish pink in color with prominent rugae (Fig. 1A). In Group II, The glandular stomachs showed multiple a dark brown mucosal lesion of different sizes, shapes and number (ranged from 8 to 9 with mean of 8.0) (Fig. 1B). The glandular stomachs of Group IIIa revealed multiple dark brown lesions (range, 5–7; mean, 7.75) (Fig. 1C). In Group IIIb, the glandular stomachs showed a few dark brown lesions (range, 1–2; mean, 3.5) (Fig. 1D). As regard of Group IVa, the glandular stomachs showed fewer lesions (range, 2–3; mean, 3.5) than were observed on the 1st day in Group III (Fig. 1E). While in Group IVb, The glandular stomachs showed very small dark brown lesions (mean, 0.875) which was decreased compared with observation with the Group III on the 4th day (Fig. 1F). Significant difference from the Group I at P<0.05, significant difference from the Group II at P<0.05; histogram 1 (Fig. 1G).

Hematoxylin and eosin stain
The sections of the glandular stomach of Group I showed that the fundic mucosa has normal intact surface epithelium with gastric pits. The surface epithelium and gastric pits are lined with simple columnar epithelial cells containing oval nuclei and pale vacuolated acidophilic cytoplasm (Fig. 2 B). Fundic glands with regular narrow tubules occupying the whole thickness of the lamina propria were observed. They comprised an isthmus, neck, and base. In addition, the muscularis mucosa, submucosa and muscularis externa all presented a normal appearance (Fig. 2A). The lamina propria contained connective tissue (C.T.) cells and fibers (Fig. 2B). The fundic glands are lined by mucous neck cells, parietal cells and chief cells. The mucous neck cells occupied the upper portion of the glands (the neck) and were lodged in between the parietal cells (Fig. 2B). They were columnar cells with pale basophilic cytoplasm and oval nuclei. The parietal cells occupied the middle and lower part of the glands. They were rounded or pyramidal in shape with rounded nuclei and deep acidophilic cytoplasm (Fig. 2B, C). The chief cells predominated in the lower part of the glands and were columnar in shape with rounded nuclei and cytoplasm showed basal basophilia and apical acidophilia. The muscularis mucosa appeared to comprise two layers of smooth muscle cells; inner circular and outer longitudinal cells with flat nuclei and acidophilic cytoplasm (Fig. 2C). The submucosa appeared to be composed of C.T. and blood capillaries (Fig. 2C). In Group II, the H&E stained sections showed variable degrees of mucosal lesions comprising partial loss of the mucosa, shedding of the whole thickness of the epithelium.
and disorganized fundic glandular cells with preservation of some glands (as the chief cells) in the lower portion of the mucosa (Fig. 2A, B). The surrounding fundic mucosa showed normal structure (Fig. 2A). Dilated blood capillaries were detected in the lamina propria (Fig. 2A, C). Widening of spaces of the submucosal C.T. components was also observed (Fig. 2A). Some of the surface epithelial cells showed densely stained nuclei with distorted cell shape (Fig. 2B). The glandular cells were also distorted in shape and arrangement and their nuclei were densely stained (Fig. 2B). The parietal cells lost their normal arrangement and appeared vacuolated and ballooned. Some of the parietal cells demonstrated deeply acidophilic cytoplasm and densely stained nuclei, while others preserved their normal histological structure (Fig. 2B, C). Mononuclear cellular infiltration appeared in the lamina propria (Fig. 2C). The H&E stained sections of glandular stomach mucosa of Group IIIa showed minimal regeneration features (Fig. 2A). Partial loss of fundic mucosa and invasion of the ulcer with granulation tissue (fibroblasts and blood capillaries) were detected (Fig. 2A). The architecture of the mucosa remained disorganized with poorly differentiated fundic glands (Fig. 2C). Monocellular infiltration was detected in both the lamina propria (Fig. 2B, C). The H&E stained sections of Group IIIb revealed disorganized epithelial and glandular cells (Fig. 2). In addition, heavy cellular infiltration of lamina propria and submucosa with granulation tissue was detected (Figs. 2A, B). Lymphocytic infiltration was observed (Fig. 2B). Infrequent parietal cells appeared

Fig. 1. Macroscopic photographs of gastric mucosa from rats in the six groups showing: (A) the glandular portion (Gl) with stomach rugae and fore-stomach (F) (Group I). (B–E) Showing numerous hemorrhagic lesions in the glandular mucosa (H) (Group II); (C) Group IIIa; (D) Group IIIa; (E) Group IVa; (F) Group IVb. (G) Statistical analysis of the ulcer index. *Significant difference from the control group (Group I) at P<0.05; †Significant difference from the ulcer group (Group II) at P<0.05.
Fig. 2. H&E stained sections of the glandular portion showing (Group I) surface epithelium (arrowhead) and gastric pits (arrow) (A), high magnification (B, C). Group II (A) the ulcer region (double arrow) with disfigured appearance and partial shedding of the surface epithelium (E) and upper portion of the gland and NO. (B, C) High magnification; the middle and lower part of fundic glands showing dense pyknotic nuclei (tailed arrow), cellular infiltration (INF), and nuclei of connective tissue (CT) cells (wavy arrow). Group IIIa (A, B) the glandular portion showing partial loss of the mucosa in the ulcer region (double arrow) with apparent normal mucosa (NO), cellular infiltration of the submucosa (INF) and dilated blood capillaries (BC). Disorganized fundic glands (G) with granulation tissue (thick arrow), high magnification (C). Group IIIb (A) the fundic mucosa in the ulcer region (double arrow), granulation tissue (thick arrow) with disfigured epithelial and glandular cells and heavy cellular infiltration (INF) of mucosa and submucosa (S). (B, C) High magnification shows infrequent parietal cells (P). Group IVa (A, B) fundic mucosa in the ulcer region (double arrow) with disorganized mucosa, granulation tissue (thick arrow) and cellular infiltration (INF) of submucosa. (C) High magnification. Group IVb (A) the glandular parts showing complete restoration of the epithelium (arrow head), gastric pits (arrow) and fundic glands (G), the muscularis mucosa (curved arrow) and muscularis externa (ME) appear normal. (B, C) Parietal (P) cells, mucous neck (M) cells and chief cells (C) appear normal, the parietal cells (P) and chief cells (C) appear normal. Scale bar for all photomicrographs: 25 µm.
ballooned with vacuolated cytoplasm and irregular dense nuclei. The other types of glandular cells were difficulty distinguished (Fig. 2C). Dilated blood capillaries were detected (Fig. 2B). In Group IVa, the H&E stained sections of glandular mucosa showed healed areas composed of granulation tissue. There were some fundic glandular cells that had begun to regenerate but the glandular architecture was still not well organized. In addition, there was heavy cellular infiltration (neutrophils, monocytes and lymphocytes) of the lamina propria (Fig. 2). The sections of glandular mucosa of Group IVb showed that was complete restoration of the epithelium and fundic glands (parietal, mucous neck and chief cells) with minimal cellular infiltration and granulation tissue (Fig. 2).

**Immunohistochemical results**

**Proliferating cell nuclear antigen**

The immunostained sections of Group I showed characteristic expression of PCNA in the epithelium, more apparent in the neck region, in the form of dark brown colored nuclei (Fig. 3A). In Group II, the immune-stained sections showed weak expression of PCNA in the ulcer region (Fig. 3B). In Group IIIa, The immune-stained sections showed weak
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expression of PCNA (Fig. 3C). As regards Group IIIb, the immunostained sections showed weak expression of PCNA (Fig. 3D). While in Group IVa, the immunostained sections showed weak expression of PCNA (Fig. 3E). In Group IVb, the immunostained sections showed strong expression for PCNA (Fig. 3F) in the epithelium and neck region of the fundic glands. Statistical analysis: significant difference from the Group I at \( P < 0.05 \). Significant difference from the Group II at \( P < 0.05 \), histogram 2 (Fig. 3G).

Vascular endothelial growth factor

The immunostained sections of Group I showed that the VEGF expression was revealed as the dark brown color of the cytoplasm of the epithelial and the endothelial lining cells of blood capillaries of the lamina propria and fundic glands (Fig. 4A). The immune-stained sections showed weak expression in Group II (Fig. 4B), Group IIIa (Fig. 4C), Group IIIb (Fig. 4D), and Group IVa (Fig. 4E). While in Group IVb, the immunostained sections showed strong positive expression in the epithelium and endothelial lining of blood capillaries of the lamina propria (Fig. 4F).
Fig. 6. The flow cytometric analysis of the isolated mesenchymal stem cells (MSCs) showing (A) control group (Isotope); (B): MSCs are positive to CD90 antibody; (C) MSCs are positive to CD271 antibody; (D) MSCs are positive to CD73 antibody; and (E) MSCs are negative to CD45 antibody.

Fig. 7. Images Gel electrophoresis analysis after PCR showing (A) GABDH expression on the 1st and 4th days; human leukocyte antigen (HLA) expression was down regulated in the control group on both the 1st and the 4th days. (B) GABDH expression in the stem cells-treated group on the 1st 4th days. (C) HLA expression was upregulated in the Stem cells treated group on the 1st and 4th days. (D) Statistical analysis (histogram 3). *Significant difference of the control group vs. the Stem cells treated group at P<0.05.
Morphology of isolated MSCs

Examination of the cultured adipose-derived MSCs on the 1st day after isolation showed that while many cells were floating, a few cells were attached. These attached cells were fibroblast-like and appeared spindle shape with vesicular nuclei (Fig. 5A). The cells reached ~80% confluence on the 7th day (Fig. 5B).

Characterization of MSCs by flow cytometry

FACS flow cytometry analysis of isolated MSCs revealed that the cells were negative for CD45 (Fig. 6D), and positive for CD90, CD271, and CD73 (Fig. 6A–C).

Reverse transcription polymerase chain reaction

Gel electrophoresis performed following conventional PCR showed higher expression of HLA in the Group IV (Fig. 7C) than in the Group I (Fig. 7A). Expression of HLA was normalized to GADPH levels in the two groups (Fig. 7A, B).

Real time PCR results

The RT-PCR results showed that the expression of HLA gene in the Group IV was higher than that in the Group I. Statistical analysis: Significant difference of the Group I vs. Group IV at \( P < 0.05 \) (Fig. 7D).

Discussion

Gastric ulceration is a benign lesion on the mucosal epithelium when exposed to excess acid and aggressive pepsin activity [33]. Adipose-derived MSCs represent a very promising line of treatment in many diseases. The use of MSCs is based on their potential capacity to repair damaged tissues and inhibit inflammation and fibrosis [34, 35].

In this study, indomethacin was used for ulcer induction in adult male albino rats. Upon gross examination of stomach tissues, the results of our study showed that administration of indomethacin resulted in multiple hemorrhagic lesions along with a significant increase (\( P < 0.05 \)) in gastric ulcer index in the ulcer group (Group II) when compared with the control group (Group I). This was in agreement with results from previous studies [36, 37]. This is explained by indomethacin comprises polar lipids with a high affinity for the lipophlic areas of cell membranes, where their polar groups trigger membrane disruption, with a loss of structural phospholipids and membrane proteins. This leads in turn to reduce hydrophobicity of the mucosal coat adherent to the mucosal cell surface which facilitates the entry of harmful water-soluble agents and also alter membrane fluidity that plays a key role in the development of the gastric mucosal lesions induced by indomethacin [38]. It was reported that indomethacin possesses much more ability to cause gastric injury than other NSAIDs; as it inhibits both cyclooxygenases (COX-1 and COX-2) which are enzymes responsible for the formation of prostanooids including thromboxane and prostaglandins from arachidonic acid [38, 39]. This will result in suppression of endogenous biosynthesis of prostaglandin with subsequent inhibition of gastro-duodenal bicarbonate secretion; interference with the mucosal barrier and interruption of the mucosal blood flow in animals [24].

The results of our macroscopic examinations in the Group III showed that the number and severity of ulcer differed on successive days following indomethacin administration. On the 1st day, the number of ulcers was higher than the 4th day as indicated by the ulcer index. Also, the rats in this group were found to be in bad general physical condition; four rats in this group died on the 4th day. This may be due to the occurrence of complications of healing as infection which delay the healing process. The isolated MSCs have the characteristics of stem cells according to the Mesenchymal and Tissue Stem Cell Committee of the International Society of Cellular Therapy. Results of the macroscopic examination of tissue from the Group IV, showed that the number and severity of ulcers decreased from the 1st to the 4th day as indicated by the ulcer index. Unlike the rats in the Group III, we found that the rats in this group demonstrated normal general condition at the 4th day; this could be due to enhanced healing of the ulcer in Group IV.

Regarding the microscopic results, our study revealed distorted appearance of the fundic mucosa of the Group II, which took the form of desquamated surface epithelium and upper portion of the fundic gland. We also observed submucosal edema marked infiltration with inflammatory cells and congested blood vessels. Separation of cells from one another and loss of the normal architecture of the mucosa were frequently observed. This may be due to a loss of integrity in the intercellular junctions of the surface epithelial and glandular cells. These results are supported by the observations of Werther [40] and Abd-Elmenm et al. [1], who demonstrated that the tight junctions between surface epithelial cells constitute a barrier that prevents back diffusion of acid and pepsin. Additionally, generation of free oxygen radicals and especially lipid peroxide plays a key role in the progression
of the gastric mucosal lesions caused by indomethacin. The
changes observed in the Group II, in this study are due to the
permeability disorders of the membranes caused by reactive
oxygen species (ROS), which are believed to be responsible
for damage to glandular and epithelial cells, which is due to
enhancement of water and electrolyte transport into the cells
[41-43].

The microscopic results in the Group III showed several
different histological features that appeared on the 1st day,
including the formation of granulation tissue (fibroblasts and
blood capillaries). In addition, normal architecture in the
fundic mucosa was lost and cellular infiltration occurred in
both the mucosa and submucosa. These conditions continued
until the 4th day without significant change. Giant cells
were found that could indicate heavy debris accumulation
resulting from a severe inflammatory process. All these findings
could be due to bad healing process. The formation of
granulation tissue observed in the Group III could be due to
angiogenesis, which is a feature of ulcer healing. One of the
angiogenic growth factors that stimulate the process of angio-
genesis is VEGF [44], a naturally secreted glycoprotein that
acts as growth factor for endothelial cells and it is produced
by several tissues when oxygen tension is decreased [45].

The microscopic examination of the Group IV revealed
that the ulcer healing started on the 1st day in the form of
granulation tissue formation, disorganized architecture of the
fundic mucosa and cellular infiltration. By the 4th day,
we found that cellular infiltration, edema and granulation
tissue had decreased, which indicate improved healing of the
ulcer. This could be explained by that MSCs produce a range
of factors implicated in immune modulating effects, such as
indoleamine 2,3-dioxygenase, IL-6, IL-10, PGE2, transforming
growth factor-β1, nitric oxide, heme oxygenase-1, and
HLA-G5 [46]. They also suppress NK cells, induce polymor-
phonuclear cell and cytotoxic T cell apoptosis, and promote
the generation of regulatory T cells [47]. In short, they inhibit
innate immunity by blocking the differentiation and matu-
rature of monocytes to dendritic cells, inhibiting NK cells,
and inducing neutrophil apoptosis [48].

In addition, regeneration of the surface epithelium and
glandular cells in the mucous neck cells, chief cells and para-
telial cells was observed on the 4th day. This could be due
to ability of MSCs to repair tissues, stimulate angiogenesis
and prevent fibrosis through their multilineage differentiation
capacity. However, recent reports have demonstrated that
most of the biological effects of MSCs are mediated by
paracrine mechanisms involving the secretion of cytokines,
chemokines, and growth factors [49, 50]. In addition, the bi-
ological effects of MSC also may be exerted by the induction
and stimulation of endogenous host progenitor cells which
improves the regenerative process [51]. In the results of our
study, we noticed a decreased incidence of inflammation as
cellular infiltration, edema and destructed cells in Group IV,
which could be due to the anti-inflammatory effect of MSCs.
Recent studies have posited that MSCs may have either an
anti-inflammatory or a pro-inflammatory role, depending
upon their interaction with other cell types and/or soluble
factors [52, 53].

Our results showed that PCNA-positive cells expression
in the Group III were weak, with no obvious difference in
the results between the 1st and 4th day. In the Group IV, we
noticed strong expression observed in the on the 4th day. We
also noticed reepithelization of the tissue surrounding the
ulcer which is normally occurring feature of healing process.
Our morphometrical results showed that PCNA-positive cell
expression which indicates a greater proliferation of cells
during healing process was statistically higher in the Group
IV than in the Group III. The distribution of PCNA-positive
cells within the gastric mucosa occurred specifically in the
neck region of the fundic glands where the stem regenerating
cells are found. Proliferation was also detected at the ulcer
margin due to proliferation of the surrounding tissue to re-
place damaged cells in the ulcer. These could be due to the
ability of MSCs to induce proliferation of cells at the ulcer
base and at the margin to replace a variety of damaged cells.
It has been reported that the MSCs induce cell proliferation
by secreting growth factors such as hepatocyte growth factor
(HGF) [54] and epidermal growth factor (EGF) [3], both of
which are mandatory for cell proliferation. HGF is a potent
stimulus for cell proliferation in the gastric epithelium. As an
established factor, it may be involved in angiogenesis during
ulcer healing [54, 55]. It has been found that MSC adminis-
tration stimulates liver regeneration in normal mice, and re-
stores this response in obese mice after partial hepatectomy.
The therapeutic effects were related to an in vivo stimulation
of hepatocyte proliferation as indicated by the expression of
PCNA [56].

In addition, hippocampus cell proliferation and neuro-
logical function was markedly enhanced in a previous study,
this underscores MSCs’ potential to promote cellular prolifer-
ation by accelerating the expression of PCNA after ischemia/
reperfusion injury [57]. Other investigators reported that ac-
acceleration in the production of HGF by the MSCs in the early period through paracrine effects, supporting the effective treatment strategy in liver regeneration [58].

In our results, we found that VEGF expression in the Group III was weak, with no obvious difference in the results between the 1st and 4th day. This could be due to insufficient amount of secreted VEGF due to the underlying defects in the healing process as infection or fibrosis which can hinder secretion of angiogenic growth factor and hence the angiogenesis. The results from the Group IV revealed strong expression of the antibody in the cytoplasm of the endothelial cells in the lamina propria of the fundic mucosa. We found no obvious difference in VEGF expression on the 1st day. A clear difference was detected on the 4th day on which the expression of the antibody was strong positive. No obvious difference was observed in VEGF expression on the 1st day between the Group III and the Group IV. This could be explained by the ability of the MSCs to secret more VEGF in the ulcer area or it might stimulates the cells which secret VEGF to increase its secretion.

Our flow cytometry results showed that MSCs are adherent, highly proliferating non-hematopoietic cells that express the surface markers CD90, CD217, and CD73, and do not express CD34 and CD45 as demonstrated by flow cytometry analysis, these results were in agreement with those analyses [54, 59].

RT-PCR studies revealed that while the HLA-gene was over expressed in the Group IV on 1st and the 4th days, it was expressed in very low levels in the control animals on both days. This means that it was increased (up regulated) in the Group IV compared to the Group I which is with the homing theory. This provides evidence that the enhancement of ulcer healing has been due to injected Stem cells and also reveals that MSCs remain viable during the process of ulcer healing. Also, HLA-up regulation on the 4th day after stem cell injection demonstrated that the cell were in place and still viable at this time allowing them to contribute to the healing process. Statistical analysis, revealed no significant difference in HLA gene expression between the numbers of MSCs that migrated on the 1st day and those that migrated on the 4th day. This could be because the severe inflammatory process that occurred after ulcer induction was still present at the 1st day, and caused the MSCs to migrate to the site of inflammation. On the 4th day, the MSCs were found at the site of the ulcer in nearly the same number which indicates that MSCs were viable and continued to secrete the growth factors that accelerated the healing process. There was a statistically significant difference between the Group IV and Group I.

Resident MSCs movement seems to be directed by cytokines and/or chemokines that become upregulated under conditions of inflammation with subsequent release of MSCs into the circulation and down-regulation of the adhesion molecules that retain MSCs in their niche [58]. The chemokines CXCL9, CXCL16, CCL20, and CCL25 significantly enhanced migration of MSCs across the endothelial cells. It was also found that following adhesion and trans endothelial migration of MSCs, gene expression was altered(disturbed) and chemokine receptors (CXCR3, CXCR6, CCR6, and CCR9) were down-regulated (knocked down) [60].

From the above-mentioned findings, we conclude that adipose-derived MSCs accelerate healing of gastric ulcer in rats. They accelerate cell proliferation to repair the damage caused by the ulcer and enhance angiogenesis which keeps the site of ulcer well oxygenated. This guard against infection which results in defective and complicated of wound healing as fibrosis and the development of chronic gastric ulcers.

**ORCID**

Safaa A. Hassan: https://orcid.org/0000-0002-2902-3228
Amal Taha Abou Elghait: https://orcid.org/0000-0002-5173-6320
Zainab S. Abdelqader: https://orcid.org/0000-0003-3275-7780
Fatma Y. Meligy: https://orcid.org/0000-0002-6372-8731

**Author Contributions**

Conceptualization: SAH, ATAE, ZSA, FYM. Data acquisition: SAH, ATAE, ZSA, FYM. Data analysis or interpretation: SAH, ATAE, ZSA, FYM. Drafting of the manuscript: SAH, ATAE, ZSA, FYM. Critical revision of the manuscript: SAH, ATAE, ZSA, FYM. Approval of the final version of the manuscript: all authors.

**Conflicts of Interest**

No potential conflict of interest relevant to this article was reported.
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