Role of Stem Cells in Management of Experimentally Induced Hypothyroidism in Albino Rat


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Abstract

Background: Hypothyroidism is a common condition with potentially devastating health consequences that spread worldwide. Stem cells (SCs) based therapy had showed promising benefits in treatment of many diseases on animal models and applied on some human clinical issues.

Aim of Study: To investigate the effect of bone marrow mesenchymal stem cells (BM-MSCs) on the biochemical and histological structures of the thyroid gland after experimentally induced hypothyroidism in adult male albino rats.

Material and Methods: Forty five adult male Sprague-Dawley albino rats were divided into 3 equal groups: Control group (Group I), Hypothyroidism induced group (Group II); received orally carbimazole 45mg/kg of rat/day for 5 days/week for 4 weeks and Induced hypothyroidism treated with BM-MSCs group (Group III); received oral dose of carbimazole 45mg/kg of rat/day for 5 days/week for 4 weeks then received single intravenous injection of 1x10^6 cells of BM-MSCs. This group was subdivided into three equal subgroups according to the time of scarification; Subgroup III6: Scarified at the end of 6th week, Subgroup III8: Scarified at the end of 8th week and Subgroup III10: Scarified at the end of 10th week.

At the end of the experiment, the hormones of thyroid function were measured and histological examination of the thyroid tissue were done. In addition to morphometric study and statistical analysis.

Results: BM-MSCs injection to the induced hypothyroidism rats showed that after 6 and 8 weeks, there were highly significant increase (p<0.01) of freeT3 and T4 values and highly significance decrease of TSH compared to the induced hypothyroidism group. On the other hand, after 10 weeks of BM-MSCs injection, there were insignificant differences (p>0.05) of free T3, T4 and TSH compared to the induced hypothyroidism group. The histological examination of the induced hypothyroidism after 6 weeks of BM-MSCs injection showed variable degree of response. However, after 8 weeks of BM-MSCs injection, there was universal improve-

Conclusion: BM-MSCs injection almost restored the function and histological structures of the thyroid gland after 8 weeks. However, this improvement was declined after 10 weeks of BM-MSCs injection. So second dose is recommended.

Key Words: Albino rats – Thyroid gland – Induced hypothyroidism – BM-MSCs.

Introduction

HYPOTHYROIDISM is a hypometabolic clinical state resulting from inadequate production of thyroid hormones for prolonged periods, or rarely, from the resistance of the peripheral tissues to the effects of thyroid hormones [1]. Carbimazole is an antithyroid medication, generally used to treat Graves' disease. Carbimazole was a prodrug of the active structure methimazole which kept the thyroid peroxidase enzyme from coupling and iodinating the tyrosine deposited on thyroglobulin [2]. The stem cells (SCs) are undifferentiated cells that can differentiate into specialized cells and could divide (through mitosis) to produce more SCs. They are present in multicellular organism. There are three types of SCs: embryonic (ESCs), fetal (FSCs) and adult (ASCs) [3]. ASCs refers to any cell which is found in a developed organism that has two properties; the ability to divide and create another cell like itself and the ability to divide and create a cell more
differenitated than itself \[4\]. ASCs is commonly used in cell based therapy because of their ability to divide or selfrenew indefinitely, and it can generate all the cell types of the organ from which they originate. Unlike for ESCs, the use of human ASCs in research and therapy is not considered to be controversial, as they are derived from adult tissue samples rather than human embryos. It can be easily obtained in large amounts from patient's own tissue (especially bone marrow and fat) and present an extremely low risk of tumorigenesis \[1\]. Mesenchymal stem cells (MSCs) are special type of ASCs. MSCs have the potential to differentiate in vitro and in vivo into multiple lineages including adipogenic, chondrogenic and osteogenic. In addition, MSCs have the capability to self-renew in order to maintain their undifferentiated state. Regardless of their tissue of origin, MSCs are characterized by the presence or absence of certain markers \[6\]. MSCs are used in treatment of Parkinson's disease and muscular dystrophies. These cells also have been used for treatment of diabetes and its complications. These cells were used in the management of infertility (both male and female) \[7\]. Bone marrow mesenchymal stem cells (BM-MSCs) are cellular component of the bone marrow stroma which contains a heterogeneous population of cells that, as a whole, contributed to hematopoi- esis. It includes adipocytes, reticular cells, macrophages, vascular endothelial cells, smooth muscle cells and MSCs \[8\].

This work aims to investigate the effect of BM-MSCs on the biochemical and histological structures of thyroid gland after experimentally induced hypothyroidism in adult male albino rats.

**Patients and Methods**

This work was conducted from 9/2016 to 9/2018 at animal house of Faculty of Medicine for Girls, Al-Azhar University, Egypt.

**A- Experiment:**

Forty five adult male Sprague-Dawley albino rats weighting about 250-3 00±50gm were used. The rats were purchased from the Egyptian Organization for Biological Products and Vaccines (Cairo, Egypt). The animals were kept in the animal house of Faculty of Medicine for Girls Al-Azhar University (Cairo, Egypt). They were housed in stainless steel cages (27X38X17cm) at 21ºC-24ºC and acclimatized in a 12h light/dark cycle. They were allowed for laboratory rat chow diet and water. Strict care and hygiene were taken to maintain normal and healthy environment for all rats all time. The study followed the International Society of Applied Ethology (ISAE) guidelines concerning animal rights. The rats were divided into three equal groups of 15 rats/each as following:

**Group I: Control group:** Without receiving anything.

**Group II: Hypothyroidism induced group:** Each rat received oral dose of carbimazole 45mg/kg of rat/day for 5 days/week for 4 weeks.

**Group III: Induced hypothyroidism treated with BM-MSCs group:** Each rat received oral dose of carbimazole 45mg/kg of rat/day for 5 days/week for 4 weeks then received single intravenous injection of 1x 10^6 cells of BM-MSCs suspended in 1 ml of phosphate buffered saline and nutritive serum in the tail vein. This group was subdivided into three equal subgroups according to time of scarification after BM-MSCs injection: Subgroup III6: Scarified at the end of 6 th week. Subgroup III8: Scarified at the end of 8 th week. Subgroup III10: Scarified at the end of 10 th week.

**B- Drugs:**

1. Carbimazole is an antithyroid medication, generally used to treat Graves’ disease. It is produced by Chemical Industries Development (CID) in the form of white, crystalline tablet that is freely soluble in water. Each tablet contains 5mg (43.8 µmol) methimazole \[9\]. The dose used in this study was 45mg/kg of rat/day for 5 days/week for 4 weeks via gastric gavage tube according to Europa et al. (2010) \[10\].

2. BM-MSCs were isolated from 10 albino rats 6-8 weeks old. The animal was put into an anesthesia chamber. Skin incisions in the lateral aspect of the thigh were done then the femurs and tibias from the hind limbs were cut off after removal of tendons and muscles. The dissected femurs and tibias were put in 70% isopropanol for a few seconds then transferred to Phosphate buffer saline (PBS). In a biosafety cabinet, the femurs and tibias were transferred to a 10cm dish containing Dulbecco’s modified Eagle’s medium (DMEM). Each bone was then held with tweezers and the two ends were cut open with a scissors. A22G needle attached to a 3ml syringe and filled with DMEM. The bone marrow was harvested by flushing the tibiae and femurs with DMEM supplemented with 10% Fetal bovine serum (FBS). The bone marrow was harvested in heparinized conical tubes. Under aseptic conditions for all cases, the harvested bone marrow cells were gently pipetted to break up cell clumps, the cells were centrifuged at 2000 rpm for 20min,
then the cell pellet was resuspended carefully onto 5ml of 60% Ficoll Hypaque separating solution (Ficoll/Paque (Pharmacia)) in sterile conical tube, and then centrifuged for 20-25min at 2000rpm at 8°C. The mononuclear cells were retrieved from the buffy coat layer by sterile Pasteur pipette and placed in 5ml sterile conical tube. The cells were washed two to three times with PBS and centrifuged at 2000rpm for 20min. Cells were counted using automated cell counter (Cell Dyn, Inc, USA). The nucleated cells were resuspended in complete culture medium supplemented with 1% penicillin-streptomycin. The cell suspension was seeded in three T25 tissue culture flasks, in a density of 100,000 cells in each flask, with 5ml culture medium. Then the cells were incubated at 37°C in humidified atmosphere of 5% CO₂ for 12-14 days as primary culture or upon formation of large colonies. When large colonies developed (80-90% confluence), cultures were washed twice with PBS. To remove the none-adherent cells, the cultures were inspected daily for the formation of adherent spindle shaped fibroblast cell colonies. The sub-culturing was carried out by 0.25% trypsin in 1ml EDTA for 5min at 37°C. After centrifugation, cells were resuspended with serum-supplemented medium and incubated in 50cm² culture flask (Falcon). The resulting cultures were referred to as first-passage cultures. On day 14, the adherent colonies of cells were trypsinized, and counted [11,12]. For identification of BM-MSCs, Flow cytometric analyses were carried out on fluorescence activated cell sorter flow cytometer. The cell surface molecules analysis was performed to verify the surface marker expressions. The cells were detached from culture dish with Trypsin/EDTA and counted. About 2X10⁵ cells of BM-MSCs were divided into three aliquots in amber-tinted 5ml centrifuge tubes and 3% rat serum was added. The cells were incubated on ice for 30 minutes, resuspended in 400 µl PBS and pelleted by centrifugation for 10 minutes at 400Xg [13]. After that the cells were stained with three fluorescent markers for 1 hour at 4°C; anti-CD29 (mesenchymal stem cell-specific marker; Biolegend- USA), anti-CD90 (mesenchymal stem cell-specific marker; Biolegend, San Diego, CA) and Isothiocyanate-conjugated rat antimouse CD34 (BD Pharmingen, Franklin Lakes, NJ, USA). After washing with PBS, the cells were resuspended in 0.5ml stain buffer (BD- Pharmingen) for flow cytometric analysis [14]. The cells showed positive staining against MSC markers CD29 (Fig. A) and CD90 (Fig. B) but negative against CD34 (Fig. C). This indicated that the enrichment of the cells was good for rat MSCs.

Figs. (A,B,C): Photographs of flow cytometer analysis to verify the markers of MSCs.

C- Collection of the specimen and preparation for examination:

At the end of the experiment, the animals were anesthetized by ether to avoid the effect of stress of manipulation on hormonal levels; blood was
collected through the retro-ocular puncture. Then
all studied animals were dissected, the skin at the
midline was pulled up and cut forward to the level
of lower lips by using a forceps and a pointed
scissor. The skin is then separated from the under-
lying muscle by pulling it gently by using a forceps
and a blunt probe, the ventral side of neck is clearly
exposed. The right and left sternomastoid form a
‘V’ shape at the base of the neck. After identification
of the muscles, retraction of the muscle was done
to expose the underlying trachea. The thyroid gland
is located immediately caudal to the larynx on
either side of the trachea. Once the trachea is
exposed, it needs to be traced upwards gently until
the thyroid gland is visible. The thyroid gland
usually appears as two small reddish oval masses,
one on each side of the trachea but the isthmus is
barely visible. The thyroid glands were excised and
prepared for histological studies by light mi-
croscope.

1- Hormonal assay: The blood samples were
left to clot for 20min and centrifuged at 3000 (rpm)
for 20min then the separated serum was kept frozen
at 20ºC till analysis of thyroid function test.
Assessment of levels of T3, T4 and TSH hormones
were done at Unite of Biochemistry and Molecular
Biology, Faculty of Medicine, Cairo University by
ELIZA test.

2- Light microscopic examination: The thyroid
glands were fixed in Bouin’s solution for 48hs.
Later, they were dehydrated in graded levels of
ethanol, cleared in xylene, and embedded in paraffin
wax for sectioning. The 4-gm thick sections were
cut, mounted on glass slides, and stained with
hematoxylin and eosiin stain to study the general
features, Masson’s trichrome stain to study the
distribution of collagen fibers, periodic acid-Schiff
(PAS) reactions to demonstrate the colloid content
and toludine blue stain to study the presence of
mast cells [15]. The images were taken by a micro-
scope (Leica) DM750 connected to a digital camera
in Anatomy Department, Faculty of Medicine for
Girls, Al-Azhar University, Cairo, Egypt.

3- Morphometric study: All studied groups were
assessed by ordinary light microscope using an
image analyzer computer system in the Pathology
Department of Dental Medicine for girls, Al-Azhar
University. It was used for measuring the epithelial
heights of the follicular cells in Hx & E stained
sections in ten non overlapped fields within a
standard measuring frame in magnification x400.
Connective tissue area percent in Masson’s tri-
chrome stained sections in ten non overlapped
fields within a standard measuring frame of known
area in magnification x200. Colloidal area percent
in PAS stained sections in ten non overlapped fields
within a standard measuring frame of known area
in magnification x200. The equipment consisted of
digital camera attached to a light microscope
and a computer system equipped with the software
Leica Quin500, England capable of performing
high aped digital image processing. The image
analyzer was calibrated automatically to convert
the measurement units (pixel) produced by the
image analyzer program into actual micrometer
units. The data was subjected to statistical analysis.

4- Statistical analysis: The data were collected
and analyzed by SPSS (statistical package for
social science) version 17.0 on IBM compatible
computer. Statistical analysis of the data obtained
were expressed as mean values and standard devi-
ations, and statistical significance was determined
by one-way analysis of variance (ANOVA) fol-
lowed by post hoc analysis using LSD test for
multiple comparison [16].

Results

Biochemical results:
Assessment of serum levels of freeT3, T4 and TSH:
The mean values of free T3, T4 of the induced
hypothyroidism group (group II) showed highly
significant decrease ($p<0.01$) as compared to the
control group (group I), while the mean value of
TSH showed highly significant increase ($p<0.01$)
as compared to the control group (group I). Treat-
ment of the induced hypothyroidism by BM-MSCs
injection showed that after 6 weeks (III6), there
were highly significance increase ($p<0.01$) of mean
values of serum level of T3 and T4 and highly
significance decrease of mean value of serum level
of TSH in compare to the induced hypothyroidism
group but still there were significant decrease
($p<0.05$) of mean values of free T3 and T4 and
significance increase of mean value of TSH when
compare with the control group. After 8 weeks
(III8), there were highly significance increase
($p<0.01$) of mean values of serum level of T3 and
T4 and highly significance decrease of mean value
of serum level of TSH in compare to the induced
hypothyroidism group, but there were insignifcant
difference ($p>0.05$) of mean values of free T3, T4
and TSH in compare to the control group. On the
other hand, after 10 weeks from injection of BM-
MSCs (III10), there were insignificance differences
($p>0.05$) of mean values of serum levels of T3, T4
and TSH in compare to the induced hypothyroidism
group but still there were highly significant decrease
($p<0.01$) of mean values of free T3 and T4 and
highly significance increase of mean value of TSH
when compare with the control group (group I) (Table 1 and Graphs 1,2,3).

**Microscopic results:**

Microanatomy of the thyroid gland of the adult male albino rats of the control group: Examination of serial cross sections of the thyroid gland showed that the thyroid gland was covered by thin connective tissue capsule which sent thin ill-defined septa divided the gland into small lobules. The lobules contained many rounded or oval follicles of variable sizes. The peripheral follicles were relatively larger than the central follicles. The follicles had luminae filled with homogenous acidophilic substance called colloid. The colloid of peripheral follicles had peripheral vacuolations (Fig. 1a). Each follicle was surrounded by a thin basement membrane and lined by follicular cells (thyrocytes) arranged in a single layer of either cuboidal epithelium with rounded central nuclei or columnar epithelium with basal rounded nuclei (Fig. 2a). The cytoplasm of the follicular cells contained secretory granules in their apices. There were prominent cytoplasmic villi on the apices of the follicular cells extending into the lumen (Fig. 3a). There was another type of cells which were small oval and had dark stained nuclei compared with the follicular cells. These cells were called parafollicular cells or C cells. They were apparent few in number and situated between the follicular cells on the basement membrane. The follicles were separated from each other by scanty interstitial connective tissue which contained blood capillaries. Also, there were aggregations of multiple polygonal cells with central rounded nuclei called solid cell nest inbetween the follicles (Figs. 2a,3a). Masson Trichrome stain, showed the normal distribution of collagen fibers in the capsule, the connective tissue septa and in the wall of interstitial blood capillaries (Fig. 4a). There was a strong PAS positive reaction in the basement membrane of the follicles and it appeared more prominent in the colloid (Fig. 5a). Toluidine blue stain showed few mast cells which were oval or spindle shaped with intense metachromasia masking their nuclei. These cells were present either near the follicular cells (perifollicular mast cell) or in the stroma between the follicles (stromal mast cell) (Fig. 6a).

Microanatomy of the thyroid gland of adult male albino rats of experimentally induced hypothyroidism group: Examination of serial cross sections of the thyroid gland showed that the thyroid gland was covered by thin loose connective tissue capsule which sent septa divided the gland into small lobules. The lobules contained many follicles of variable sizes and shapes. The large follicles were located mainly at the periphery. The lumen of almost all follicles contained little amount of pale colloid with large peripheral vacuolations. Few follicles had no colloid (Fig. 1b). Each follicle was surrounded by thin basement membrane. The follicles were lined by either flat follicular cells with dark flat nuclei or cubical cells with dark central rounded nuclei (Fig. 2b). Most of these cells showed cytoplasmic vacuolations or became ballooned (Fig. 3b). The lumen of some follicles contained exfoliated cells with densely stained nuclei. There were also few degenerated follicles filled with pyknotic cells due to sloughing of follicular cells. The parafollicular cells were difficult to differentiate. There was an apparent widening in the interstitial spaces between the follicles and the follicles are fenestrated by large blood capillaries (Figs. 1b,2b). Masson Trichrome stain, showed loose collagen fibers deposition in the capsule, the septa and in the wall of interstitial blood capillaries (Fig. 4b). There were moderate positive PAS reaction of follicular basement membrane and in the colloid when present (Fig. 5b). By toluidine blue stain, there were many small perifollicular and stromal mast cells mainly around the blood capillaries (Fig. 6b). Furthermore, the morphometric results clarified that there was highly significant decrease ($p<0.01$) of the mean heights of follicular cells, area % of collagen fibers and area % of colloid in group II (Induced hypothyroidism) compared to the control group (Table 2 and Graphs 4,5,6).

1- **Subgroup III6:** Examination of serial cross sections of the thyroid gland showed variable degree of response from mild to moderate to good response. The capsule was slightly thick and sent ill defined connective tissue septa divided the gland into small lobules. The follicles appeared compact. Most follicles were filled with colloid with some peripheral vacuolations. The interstitial space was nearly absent and still contained dilated and congested blood capillaries (Figs. 1c,2c). In mild response area, the thyroid follicles were lined by cubical cells with rounded central nuclei in some parts and lined by flat cells with flat nuclei in other parts. The cytoplasm of many follicular cells had apparently few dark secretory granules near their apices. Few follicular cells had cytoplasmic villi. There were many parafollicular cells situated on basement membrane (Fig. 3c). In moderate response
area, the majority of the follicular cells were cuboidal with large rounded central nuclei. The cytoplasm of follicular cells had dark secretory granules near their apices and the cytoplasmic villi were more prominent. There were many parafollicular cells situated between the follicular cells on the basement membrane. Some thyroid follicles had honey comb appearance of its colloid (Fig. 3d). In good response area, the follicular cells were either cuboidal with large rounded central nuclei or columnar with large rounded basal nuclei similar to the control group. The cytoplasm of follicular cells had dark secretory granules near their apices and the cytoplasmic villi were more prominent. The parafollicular cells were situated between the follicular cells on the basement membrane (Fig. 3e). Solid cell nests appeared between the follicles when compared to the hypothyroidism induced group which was absent (Fig. 3c). Masson Trichrome stain, showed moderate collagen fibers deposition in the connective tissue septa and the wall of interstitial blood capillaries (Fig. 4c). There were variable degrees of reaction in the same field rang from strong, moderate and weak PAS positive reaction in the colloid and in the basement membrane of the follicles (Fig. 5c). By Toludine stain, there was apparent increase in number of perifollicular and stromal mast cells with increased areas of metachromasia (Fig. 6c). Furthermore, the morphometric results showed that there was highly significant increase ($p<0.01$) of the mean heights of follicular cells, area % of collagen fibers and area % of colloid of the induced hypothyroidism treated after 6 weeks of BM-MSCs injection (group III8) in compared to the induced hypothyroidism group but still there was highly significant decrease ($p<0.01$) of the mean heights of follicular cells in compared to the control group (group I). On the other hand, there was highly significant increase ($p<0.01$) of the mean values of area % of collagen fibers and area % of colloid of the induced hypothyroidism treated after 8 weeks of BM-MSCs injection (group III8) in compared with the control group (group I). Also, there was insignificant difference ($p>0.05$) of the mean values of area % of colloid of the induced hypothyroidism after 8 weeks of BM-MSCs injection (group III8) in compared to the control group (group I) (Table 2 and Graphs 4,5,6).

2- Subgroup III8: Examination of serial cross sections of the thyroid gland of adult male rats of the induced hypothyroidism after 8 weeks of BM-MSCs intravenous injection showed universal improvement of all examined fields more than the previous subgroup. The capsule was slightly thick. There were more compact thyroid follicles with variable size and shape (Fig. 1d). The luminae of all follicles were filled with a homogenous acidophilic colloid with few peripheral vacuolations. The interstitial spaces were nearly absent and there were more SCN between the follicles. Many follicles were lined by a single layer of columnar epithelium with apparently large rounded basal nuclei. Some follicular cells were cuboidal with apparently large rounded central nuclei (Fig. 3d). The cytoplasm was dense acidophilic and filled with secretory granules. The villi were more prominent. There were many parafollicular cells which situated within the basement membrane (Fig. 3f). Masson trichrome stain showed fine collagen fibers deposition in the capsule and the connective tissue septa (Fig. 4d). Strong PAS positive reaction was present in the basement membrane of the follicles and in the colloid (Fig. 5d). By Toludine stain, there were many perifollicular and stromal mast cells which appeared large in size with increased areas of metachromasia (Fig. 6d). Furthermore, the morphometric results showed that there was highly significant increase ($p<0.01$) of the mean heights of follicular cells, area % of collagen fibers and area % of colloid of the induced hypothyroidism after 8 weeks of BM-MSCs injection (group III8) in compared to the induced hypothyroidism group but still there was highly significant decrease ($p<0.01$) of the mean heights of follicular cells in compared to the control group (group I). On the other hand, there was highly significant increase ($p<0.01$) of the mean values of area % of collagen fibers and area % of colloid of the induced hypothyroidism after 8 weeks of BM-MSCs injection (group III8) in compared to the control group (group I) (Table 2 and Graphs 4,5,6).

3- Subgroup III10: Examination of serial cross sections of the thyroid gland of adult male albino rats of the induced hypothyroidism after 10 weeks of BM-MSCs intravenous injection showed marked regression in response to SCs treatment as compared to previous two subgroups. The thyroid gland was covered by thin loose connective tissue capsule. It had variable size thyroid follicles. Some follicles lost their normal architecture. Many follicles were filled with colloid but some follicles had central vacuolations of their colloid (Figs. 1e,2e). Many thyroid follicles were lined by flat follicular cells with small dark flat nuclei and few follicles were lined by cubical cells with small dark rounded nuclei. The cytoplasm of many follicular cells showed vacuolations. Moreover, some cells had ballooned cytoplasm with compressed flat nuclei. In addition, some follicles had disrupted basement membrane. The parafollicular cells were few and difficult to differentiate. The interstitial spaces became wide again. However by screening exam-
In the cross section, there was absence of SCN (Figs. 2e, 3g). Masson trichrome stain showed mild collagen fibers deposition in the capsule and the connective tissue septa (Fig. 4e). Strong PAS positive reaction was present in the basement membrane of the follicles and in the colloid (Fig. 5e). By Toluidine blue stain, there were small scanty mast cells in the stroma with less areas of metachromasia than the previous subgroup (Figs. 6e, 6d).

Furthermore, the morphometric results showed that there was insignificant difference \( (p>0.05) \) of the mean heights of follicular cells, area % of collagen fibers and area % of colloid of the induced hypothyroidism treated after 10 weeks of BM-MSCs injection (group III10) in compared to the induced hypothyroidism group. In contrast, there was highly significant decrease \( (p<0.01) \) of the mean heights of follicular cells, area % of collagen fibers and area % of colloid of the induced hypothyroidism treated after 10 weeks of BM-MSCs injection (group III10) compared to the control group (group I) (Table 2 and Graphs 4, 5, 6).

Table (1): Mean values and standard deviations of the serum level of free T3, T4 and TSH of all experimental groups.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Serum Level of free T3</th>
<th>Serum Level of free T4</th>
<th>Serum Level of TSH</th>
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<tr>
<td></td>
<td>Mean±SD</td>
<td>Mean±SD</td>
<td>Mean±SD</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>Range</td>
<td>Range</td>
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<tr>
<td>Control group (I)</td>
<td>74.38±1.09 (72.6–76.4)</td>
<td>3.40±0.43 (2.8–4.1)</td>
<td>0.09±0.00 (0.09–0.1)</td>
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<tr>
<td>Induced hypothyroidism group (II)</td>
<td>49.64±1.58 (47.6–51.3)</td>
<td>1.86±0.45 (1.2–2.5)</td>
<td>0.30±0.04 (0.24–0.36)</td>
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<td>Induced hypothyroidism treated by Stem cells after 6weeks subgroup (III6)</td>
<td>71.55±1.63 (68.7–73.9)</td>
<td>2.96±0.15 (2.6–3.09)</td>
<td>0.15±0.03 (0.1–0.19)</td>
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<td>Induced hypothyroidism treated by Stem cells after 8 weeks subgroup (III8)</td>
<td>74.64±1.04 (72.9–76.1)</td>
<td>3.27±0.45 (2.6–3.9)</td>
<td>0.11±0.02 (0.09–0.14)</td>
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<tr>
<td>Induced hypothyroidism treated by Stem cells after 10 weeks subgroup (III10)</td>
<td>60.23±1.26 (58.3–62.1)</td>
<td>1.83±0.25 (1.4–2.05)</td>
<td>0.31±0.04 (0.25–0.37)</td>
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</table>

Graph (1): Mean values of serum level of T3 level.

Graph (2): Mean values of serum level of T4 level.

Graph (3): Mean values of serum level of TSH level.

- \( (p\text{-value}>0.05 \text{ for T3, T4 and TSH}) = \) Non significant.
- \( (p\text{-value}<0.05 \text{ for T3, T4 and TSH}) = \) Significant when compared with control group (*) and Significant when compared with induced hypothyroidism group (#).
- \( (p\text{-value}<0.01 \text{ for T3, T4 and TSH}) = \) Highly significant when compared with control group (**) and highly significant when compared with induced hypothyroidism group (##).
Table (2): It shows the means and standard deviations of the heights of follicular cells, area % of collagen fibers and area % of colloid of all experimental groups.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Heights of follicular cells</th>
<th>Area % of collagen fibers</th>
<th>Area % of colloid</th>
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<tbody>
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<td></td>
<td>Mean±SD Range</td>
<td>Mean±SD Range</td>
<td>Mean±SD Range</td>
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<tr>
<td>Control group (I)</td>
<td>15.82±4.01 (8.4–23.75)</td>
<td>8.42±2.46 (4.25–12.39)</td>
<td>23.90±7.44 (15.35–36.01)</td>
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<tr>
<td>Induced hypothyroidism group (II)</td>
<td>5.42±1.71 (2.1–8.6)</td>
<td>3.20±1.53 (1.23–6.17)</td>
<td>3.97±1.86 (0.73–6.8)</td>
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<td>Induced hypothyroidism treated by Stem cells after 6 weeks subgroup (III6)</td>
<td>10.68±2.58 (6.18–16.2)</td>
<td>10.07±3.14 (5.37–14.99)</td>
<td>15.75±4.60 (9.43–23.77)</td>
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<tr>
<td>Induced hypothyroidism treated by Stem cells after 8 weeks subgroup (III8)</td>
<td>13.25±2.82 (8.55–18.01)</td>
<td>12.97±5.52 (4.72–19.26)</td>
<td>19.52±4.47 (12.38–27.42)</td>
</tr>
<tr>
<td>Induced hypothyroidism treated by Stem cells after 10 weeks subgroup (III10)</td>
<td>4.58±1.18 (2.1–6.7)</td>
<td>5.99±2.28 (2.66–10.73)</td>
<td>12.97±4.32 (5.8–20.62)</td>
</tr>
</tbody>
</table>

Graph (4): Mean values of level of Heights of follicular cells.

Graph (5): Mean values of area % of collagen fibers.

Graph (6): Mean values of area % of colloid.

- (p-value >0.05 for heights, area % of collagen and area % of colloid) = Non significant.
- (p-value <0.05 for heights, area % of collagen and area % of colloid) = Significant when compared with control group (*) and Significant when compared with induced hypothyroidism group (#).
- (p-value <0.01 for heights, area % of collagen and area % of colloid) = Highly significant when compared with control group (**) and highly significant when compared with induced hypothyroidism group (##).
Fig. (1): Photomicrographs of cross sections of the thyroid glands of the adult male albino rats of all studied groups showed that:

Fig. (1a) control: The thyroid gland is surrounded by connective tissue capsule (C) which sends septa (S) dividing it into lobules; each of which contains many follicles (Fo) filled with colloid (Co). Notice that the peripheral follicles are relatively large and their colloid show peripheral vaculations (arrow).

Fig (1b): Induced hypothyroidism: The capsule is thin and loose (C) and the lumen of most of the follicles contain little amount of pale colloid with peripheral vacuolations (Co).

Fig. (1c): Induced hypothyroidism after 6 weeks of BM-MSCs injection: The capsule is slightly thick (C). Notice that almost all follicles filled with colloid (Co) except some peripheral follicles have small peripheral vaculations (arrow).

Fig. (1d): Induced hypothyroidism after 8 weeks of BM-MSCs injection: The capsule is slightly thick (C). Notice that all the follicles are more compact and filled with colloid (Co).

Fig. (1e): Induced hypothyroidism after 10 weeks of BM-MSCs injection: The capsule is thin and loose (C). Notice that almost all follicles filled with colloid (Co) and the interstitial space is wide (Sp). (Hx. &E. x100).

Fig. (2): Photomicrographs of cross sections of the thyroid glands of the adult male albino rats of all studied groups showed that:

Fig. (2a) control: The follicles are surrounded by thin basement membrane (arrow) and lined by follicular cells. The follicular cells are either cuboidal with central rounded nuclei (f) or columnar with basal rounded nucleus (F). The luminae of the follicles are filled with homogenous acidophilic colloid (Co). The parafollicular cells are small oval and have dark stained nuclei and situated between the follicular cells on the basement membrane (P).

Fig (2b) Induced hypothyroidism: The follicles are lined by flat (arrow) or cubical (double arrow) follicular cells on thin basement membrane. Notice that one follicle has exfoliated follicular cells with densely stained nuclei in the lumen (zigzag arrow). Also there is one degenerated follicle filled with pyknotic cells (star). Most of the follicles have small amount of pale colloid with peripheral vacuolations (Co). Notice also that the interstitial spaces (Sp) between the follicles are wide and contain dilated blood capillaries (V).

Fig. (2c) Induced hypothyroidism after 6 weeks of BM-MSCs injection: The follicles are lined by cuboidal follicular cells with large rounded central nuclei (f). There are many parafollicular cells (P). Notice that almost all the follicles are filled with colloid (Co). The interstitial spaces appear narrow with presence of dilated blood capillaries (V).

Fig. (2d) Induced hypothyroidism after 8 weeks of BM-MSCs injection: Many follicles are lined by columnar follicular cells with basal nuclei (F) while some thyroid follicles are lined by cuboidal follicular cells with large rounded central nuclei (f). There are many parafollicular cells (P). Notice that all the follicles are filled with colloid (Co). Notice that the interstitial space (Sp) is very narrow. Notice also the presence of SCN between follicles (SCN).

Fig. (2e) Induced hypothyroidism after 10 weeks of BM-MSCs injection: Many follicles are lined by flat follicular cells with flat nuclei (arrow). All the follicles are filled with colloid (Co) but some follicles have central vacuolation (Va). One follicle loses its normal architecture (arrow head) and some follicles have disrupted basement membrane (zigzag arrow). Notice that the interstitial space (Sp) is wide. (Hx. &E. x400)
Fig. (3): Photomicrographs of cross sections of the thyroid glands of the adult male albino rats of all studied groups showed that:

Fig. (3a): control: The follicle is surrounded by basement membrane (arrow) and lined by follicular cells (F). It lumen is filled with colloid (Co). Notice that the follicular cells have prominent cytoplasmic villi on their apices extending into the lumen (zigzag arrow). The cytoplasm of these cells contained secretory granules (G) in their apices. Notice also that the parafollicular cells (P) are situated between the follicular cells on the basement membrane.

Fig. (3b): Induced hypothyroidism: The follicles are lined by flat follicular cells (arrow). Notice that most of these cells show cytoplasmic vacuolations (zigzag arrow). Some follicular cells are ballooned (B) with dark compressed basal nuclei. The parafollicular cells are difficult to differentiate. The follicles have little amount of pale colloid with large peripheral vacuolations (Co).

Fig. (3c): The induced hypothyroidism after 6 weeks of BM-MSC injection: In mild response area, the thyroid follicle is lined by cubical cells with round central nuclei (arrow) in some parts and lined by flat cells with flat nuclei in other parts (double arrow). The cytoplasm of follicular cells have few dark secretory granules near their apices (G) and few cytoplasmic villi are prominent (arrow head). The follicles are filled with colloid with peripheral vacuolations (Co).

Fig. (3d): In moderate response area, the thyroid follicles are lined by cuboidal to columnar cells with large vesicular rounded central nuclei (F) and laying on thin basement membrane (arrow). The cytoplasm of follicular cells have dark secretory granules near their apices (G) and the cytoplasmic villi are prominent (arrow head). The parafollicular cells (P) are situated between the follicular cells on the basement membrane. The thyroid follicle has honey comb appearance of its colloid (Co).

Fig. (3e): In good response area, some thyroid follicles are lined by cuboidal cells with large vesicular rounded central nuclei (f) and laying on thin basement membrane (zigzag arrow). Some follicles are lined by columnar cells with large vesicular rounded basal nuclei (F). The cytoplasm of follicular cells have dark secretory granules near their apices (G) and the cytoplasmic villi are prominent (arrow head). The parafollicular cells (P) are situated between the follicular cells on the basement membrane. The follicles are filled with colloid (Co).

Fig. (3f): Induced hypothyroidism after 8 weeks of BM-MSC injection: Some thyroid follicles are lined by cuboidal cells with central nuclei (f) while many follicles are lined by columnar cells with basal nuclei (F). The cytoplasmic villi are prominent (arrow head) and the cytoplasm is filled with condensed dark secretory granules (G). There are many parafollicular cells (P). Almost all follicles are filled with colloid (Co).

Fig. (3g): Induced hypothyroidism after 10 weeks of BM-MSC injection: The thyroid follicles are lined by flat follicular cells with small dark flat nuclei (arrow). Notice the ballooning of cytoplasm of many follicular cells (B). In addition there are some follicular cells have vacuolated cytoplasm (arrow head). There are separation of contact between the disturbed basement membrane and the follicular cells (zigzag arrow). There are central vacuolation (Va) of the colloid. Notice also that the interstitial space (Sp) return wide. (Hx. &E. x1000).
Fig. (4): Photomicrographs of cross sections of the thyroid glands of the adult male albino rats of all studied groups showed that:

Fig. (4a): control: Normal distribution of the collagen fibers in the capsule (C), the connective tissue septa (S) and in the wall of interstitial blood capillaries (V).

Fig. (4b,e): Induced hypothyroidism and induced hypothyroidism after 10 weeks of BM-MSC injection showing few loose collagen fibers deposition in the connective tissue capsule (C), septa (S) and the wall of blood capillaries (V) in the interstitial tissue between the follicles.

Fig. (4c,d): Induced hypothyroidism after 6 and 8 weeks of BM-MSC injection showing moderate deposition of collagen fibers in the capsule (C), the connective tissue septa (S) and the wall of blood vessels (V) in the interstitial tissue. (Masson Trichrome x200).

Fig. (5): Photomicrographs of cross sections of the thyroid glands of the adult male rats of all studied groups showed that:

Fig. (5a,d) control and Induced hypothyroidism after 8 weeks of BM-MSC injection showing strong PAS positive reaction in the basement membrane (arrow) of the follicles and it appears more prominent in the colloid (Co).

Fig. (5b,e): Induced hypothyroidism and Induced hypothyroidism after 10 weeks of BM-MSC injection: There is moderate PAS positive reaction of follicular basement membrane (arrow) and in the colloid (Co) when present.

Fig. (5c): Induced hypothyroidism after 6 weeks of BM-MSC injection: There are variable degrees of reaction from strong (I), moderate (II) and weak (III) in the colloid (Co) and the basement membrane (arrow) of the follicles. (PAS x100).
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Fig. (6): Photomicrographs of cross sections of the thyroid glands of the adult male albino rats of all studied groups showed that:

Fig. (6a): control: Few mast cells which either oval or spindle shaped cells with intense metachromasia masking their nuclei. They are either present near the thyrocytes, perifollicular mast cell (Mf) or present in the stroma, stromal mast cell (Ms).

Fig. (6b): Induced hypothyroidism: There are many apparently small perifollicular mast cells (Mf) and stromal mast cells (Ms) mainly present around the blood capillaries (V).

Fig. (6c, 6d): Induced hypothyroidism after 6 and 8 weeks of BM-MSC injection: There are many perifollicular mast cells (Mf) and stromal mast cell (Ms) with increase areas of metachromasia.

Fig. (6e): Induced hypothyroidism after 10 weeks of BM-MSC injection: There are small scanty mast cells in the stroma (Ms) with less areas of metachromasia. (TB x400).

Discussion

Biochemical study of the experimentally induced hypothyroidism group (group II) proved evidences of hypothyroidism whereas the serum levels of free T3 and T4 were low with highly significance decrease in compare to the control group. Also, the serum level of TSH was high with highly significance increase in compare to the control group. The serum levels of the thyroid hormones seemed to reflect the disturbed histological features in the thyroid gland. Similar suggestion was reported by Zaidi et al., (2004) [17]. Milosevic et al., (2004) [18] clarified that the thyroid gland activity was regulated by hypothalamic pituitary thyroid axis, including negative feedback loop. The authors pointed to TSH as a major growth factor for the thyroid. The thyroid gland under high TSH underwent enlargement, neovascularisation and morphological alterations of the thyrocytes.

In the present study, examination of serial cross sections of the thyroid gland of adult male albino rats of the induced hypothyroidism group showed that the thyroid gland was covered by loose connective tissue capsule. The follicles were lined by either flat follicular cells with dark flat nuclei or cubical cells with dark central rounded nuclei. Most of these cells showed cytoplasmic vacuolations or became ballooned. The lumen of almost all follicles contained small amount of pale colloid with peripheral vacuolation and few follicles had no colloid. These finding were suggested to be due to hypoactivity of the thyroid gland with fluid accumulation and glandular over stimulation by high TSH which confirmed statistically by highly significance decrease of the heights of follicular cells in compare to the control group. This suggestion went hand in hand with Serakides et al., (1999) [19] and Elkalawy et al., (2013) [20]. Furthermore, Rubin and Strayer (2008) [21] explained that hydropic swelling resulted from impairment of cellular volume regulation with accumulation of sodium in the cell led to an increase in water content to maintain isosmotic conditions and consequently caused cell swelling.

In the current study, the lumen of some follicles contained exfoliated cells with densely stained nuclei. There were few degenerated follicles filled with pyknotic cells due to sloughing of follicular cells in addition to the presence of widening in the interstitial spaces between the follicles. These finding might be due to loss of cell support and hydropic degeneration of the interstitial tissue with the presence of interstitial oedema. Another explanation was reported by El Bekry and Tawfik (2014) [22] who mentioned that collapse of some follicles or necrosis of the other follicles leaving empty spaces lead to widening in the interstitial spaces.

Masson trichrome stained thyroid sections of the induced hypothyroidism group showed that there was an apparent widening in the interstitial...
spaces with loose thin collagen fiber deposition in the connective tissue capsule, septa and the wall of blood capillaries which confirmed statistically by highly significance decrease of area % of collagen fibers in compare to the control group. Similar observation was reported by Abdel-Dayem and Elgendy (2009) [23]. PAS stained thyroid showed irregular shaped follicles that exhibited moderate positive PAS reaction of follicular basement membrane and in the colloid when present which confirmed statistically by highly significance decrease of area % of colloid in compare to the control group. The referred cause might be related to increased demand to thyroid hormones which stored in thyroid colloid and the empty thyroid luminae might be due to more extraction of stored thyroid colloid to meet the hypothyroidism state. The findings of the present study were similar to Rubin and Strayer (2008) [21] and El Bekry and Tawfik (2014) [22]. In toludine blue stained thyroid sections of this group, there were many mast cells in the interstitial tissue mainly around the blood capillaries. This increment may be due to tissue compensatory mechanism to overcome the effect of hypothyroidism whereas the mast cells were a corner stone in the thyroid repair, it has the ability to act as angiogenesis and folliculogenesis. These suggestions were in agreement with Abd-El Aty and Badr El-Din (2015) [24].

In the present study, examination of the thyroid gland of the adult male albino rats of induced hypothyroidism after 6 weeks of intravenous injection of BM-MSCs showed variable degrees of improving. However, the thyroid gland became nearly similar to the control group. The serum levels of free T3 and T4 showed highly significance increase and the serum level of TSH was highly significance decrease in compare to the induced hypothyroidism group. While the serum levels of free T3 and T4 still showed highly significance decrease and the serum level of TSH was highly significance increase in compare to the control group. While after 8 weeks, the serum levels of free T3 and T4 were highly significance increase and serum level of TSH was highly significance decrease in compare to the induced hypothyroidism group. The serum levels of free T3, T4 and TSH were insignificance difference in compare to the control group. These results confirmed that BM-MSCs could compensate the induced defect. In histological examination after 6 weeks of injection of BM-MSCs, the capsule was slightly thick. The follicular cells had different shapes, some of them were cuboidal with large rounded central nuclei, some were columnar with large rounded basal nuclei and others were cubical with rounded central nuclei with intense vascularisation in between the follicles. These finding confirmed statistically by highly significant increase of the mean heights of follicular cells in compare to the induced hypothyroidism group but still there was highly significant decrease of the mean heights of follicular cells in compare to the control group. Many thyroid follicles were filled with colloid with peripheral vacuolations and this might indicate active thyroid follicles. These finding confirmed that the thyroid gland partially restored its normal histological pattern. The explanation of this partial restoration referred to the ability of the BM-MSCs to differentiate to thyrocytes in the presence of acceptable level of thyroid hormones and stoppage of administration of the trigger (carbimazole). Similar observation was reported by Salinas et al., (2007) [25]. The authors added that the active thyroid follicle extracted the stored thyroid colloid from the lumen and converted it into active thyroid hormones and the scalloped pale edge of the colloid indicated where the colloid had been removed from the follicle lumen. After 8 weeks from BM-MSCs injection, the histology of thyroid gland became apparently similar to the control group. Many follicles were lined by a single layer of columnar epithelium with apparently large rounded basal nuclei and some follicular cells were cuboidal with apparently large rounded central nuclei which confirmed statistically by insignificance difference of the heights of follicular cells in compare to the control group and highly significant increase of the mean heights of follicular cells in compare to the induced hypothyroidism group. Similar study was performed by Mikhailov et al., (2012) [26] who found that the injection of BM-MSCs contributes in follicular epithelium renewal.

The improvement in thyroid gland function and structures after 8 weeks from BM-MSCs injection appeared to be due to the migration of injected BM-MSCs to thyroid gland. Tao et al., (2016) [27] confirmed that as they found that stem cells migrated into the target organ and differentiated into a variety of cells, including vascular endothelial cells, thus revealing that BM-MSCs might contribute to regeneration of target organ by enhancing angiogenesis. Another explanation to the improvement with BM-MSCs was their paracrine effect. This was confirmed by Hu et al., (2017) [28] who reported that BM-MSCs paracrine activity had an anti-inflammatory effect and an antiapoptotic effect on intervertebral disc degeneration and this was mediated by nuclear factor-KB and mitochondrial apoptotic pathways in annulus fibrosus cells. Zuo et al., (2019) [29] added that a new specific secretory vesicles which involved in the paracrine effects of
BM-MSCs are called exosomes. It plays important role in repairing, improved safety and reduced immune rejection.

In this group, there were many large SCN appeared between the follicles when compared to the hypothyroidism induced group which was nearly absent. This might be due to its ability to rearrange as new follicles. Thomas et al., (2008) [30] reported that the main cells of solid cell nests of the human thyroid gland have some SCs properties, including a capacity for self renewal and an ability to differentiate into more than one cell type.

In Masson trichrome stained thyroid sections after 6 and 8 weeks of BM-MSCs injection, there were moderate collagen fiber deposition in the capsule, the connective tissue septa and the wall of blood vessels in the interstitial tissue between the follicles which confirmed statistically by highly significance increase of area % of collagen fibers in compare to the induced hypothyroidism group. While there was insignificance difference of area % of collagen fibers in subgroup III6 in compare to the control group and highly significance increase in subgroup III8 in compare to the control group. Isacksona et al., (2013) [31] reported that BM-MSCs increased vascularization, increased cellularity and increase collagen content. It is suggested that BM-MSCs produced these effects through two suggested mechanisms through paracrine signaling mechanisms and differentiation into resident cells.

In PAS stained thyroid sections after 6 weeks of BM-MSCs injection, there were variable degrees of reaction from strong, moderate and weak in the colloid and in the basement membrane of the follicles which confirmed statistically by highly significance increase of area % of colloid in compare to the induced hypothyroidism group but still highly significance decrease of area % of colloid in compare to the control group. On the other hand, in PAS stained thyroid sections after 8 of BM-MSCs injection, there were strong reaction in the colloid and in the basement membrane of the follicles which confirmed statistically by highly significance increase of area % of colloid in compare to the induced hypothyroidism group and insignificance difference in compare to the control group. These findings seemed to be similar to Abdul-Hamid and Salah (2013) [32] and Kassab and El-Aasr (2018) [33].

Regarding the toluidine blue stained thyroid section, there were many mast cells; perifollicular and stromal mast cell with increased areas of metachromasia present around blood vessels which indicate its role in regeneration of the follicles. Toda et al., (2001) [34] demonstrated that in thyroid regeneration, the number of mast cells increased near blood capillaries. They were closely associated with connective tissue remodeling and localized at the thyroid tissue regenerative site where both thyroid folliculogenesis and angiogenesis took place.

In the current study, examination of rats of induced hypothyroidism after 10 weeks of BM-MSCs injection showed progressive downward response to SCs treatment as compared to the previous subgroups. The serum levels of free T3, T4 and TSH were insignificance difference in compare to the induced hypothyroidism group. On the other hand, the serum levels of free T3, T4 were highly significance decrease and the serum level of TSH was highly significance increase in compare to the control group. In histological examination, the capsule was thin and loose. The basement membrane of some follicles was disrupted and few follicles lost its normal architecture. The follicular cells became either flat with small dark flat nuclei or cubical cells with small dark rounded nuclei which confirmed statistically by highly significance decrease of height of follicular cells in compare to the control group and insignificance difference in compare to the induced hypothyroidism group. Some follicular cells had cytoplasmic vacuolations and few cells were ballooned with compressed flat nuclei. The parafollicular cells were few and difficult to differentiate. The luminae of many follicles filled completely with colloid but some had central vacuolation of its colloid. The interstitial space was wide and showed dilated and congested blood capillaries. The cells nest had disappeared. Masson trichrome stain, showed mild collagen fibers deposition in the capsule and the connective tissue septa which confirmed statistically by highly significance decrease of area % of collagen fibers in compare to the control group and insignificance difference in compare to the induced hypothyroidism group. Strong PAS positive reaction was present in the basement membrane of the follicles and in the colloid but statistically there was highly significance decrease of area % of colloid in compare to the control group but highly significance increase in compare to the induced hypothyroidism group. By Toludine blue stain, there were small scanty mast cells in the stroma with less areas of metachromasia than the previous subgroups. These finding indicated that BM-MSCs failed to maintain their action in regeneration and folliculogenesis. A lot of causes could be contributed in this condition which mentioned by Wolf (2009) [35] who
reported that SCs graft failure might be due to an inadequate numbers of transplanted SCs or the failure of adequate number of these cells to survive. The authors added that the barriers to engraftment included immunologic destruction, infectious agents, drug toxicity or a poor bone marrow microenvironment. Herberts et al., (2011) [36] mentioned that the risk factor of use of SCs might be the potential high number of cells needed for the beneficial effect. It is generally unknown how many cells are needed, however, given the very low rate of retention and possible low cell survival, large number of cells may be required for obtaining maximal clinical benefit. In the same time, injection of concentrated cells into tissue may have unwanted effects. Hao et al., (2018) [37] studied the effects of multiple intravenous infusions of BM-MSCs in treatment of experimental type 2 diabetes rats. They said that single dose of BM-MSCs infusion ameliorated hyperglycemia and relieved the drug induced pancreatic damage but these effects only maintained for 2-3 weeks. Also single dose injection failed to restore normoglycemia in diabetic animals so they did multiple injections of BM-MSCs with 2 weeks interval between each dose. They found that multiple BM-MSCs injection reversed the hyperglycemia and restored the damaged pancreatic tissue.

Conclusion and recommendations:

It could be concluded that injection of single dose of BM-MSCs partially restored the function and histological structures of the thyroid gland of male adult albino rats after experimentally induction of hypothyroidism. This improvement was clearly observed after 8 weeks from BM-MSCs injection. However, progressive downward response occurred after 10 weeks. So, another dose of BM-MSCs is recommended.

References

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Role of Stem Cells in Management of Experimentally Induced Hypothyroidism


دور الخلايا الجذعية في علاج قصور الغدة الدرقية المستحدثة

الخلاصة: قصور الغدة الدرقية في حالة شائعة نوء عواقب صحية مدوية ورائدة تنتشر في جميع أنحاء العالم. وقد أظهر العلاج بالخلايا الجذعية فوائد واعدة في علاج العديد من الأمراض في النماذج الحيوانية وتطبيقها على بعض الحالات البشرية.

الهدف من العمل: دراسة تأثير الخلايا الجذعية النخاع العظمي على تركيب البروبيكيميائي والسيجي للغدة الدرقية المستحدثة تجريبياً في ذكور الجرذان البيضاء.

المواد والطريقة: تم تقسيم خمسة وأربعون من ذكور الجرذان إلى 2 مجموعات متساوية: المجموعة الضابطة (المجموعة الأولى)، مجموعة قصور الغدة الدرقية (المجموعة الثانية) تتلقى كريبتيموزول بالفم 5 مجم/كم من الجرذ/يوم لمدة 5 أيام/اسبوع لمدة 4 أسابيع ومجموعة قصور الغدة الدرقية المستحدثة تجريبياً ومعالج بالخلايا الجذعية للنخاع العظمي (المجموعة الثالثة)، جرعة بالفم من كريبتيموزول بالفم 5 مجم/كم من الجرذ/يوم لمدة 5 أيام/اسبوع لمدة 4 أسابيع ثم تلقى حقنة في الوريد من الخلايا الجذعية للنخاع العظمي. تم تقسيم هذه المجموعة إلى ثلاث مجموعات فرعية متساوية وفقاً لوقت التنبغ، المجموعة الفرعية: IIII تنبغ في نهاية الأسبوع السادس، المجموعة الفرعية: IIII تنبغ في نهاية الأسبوع الثامن، والفئة الفرعية: IIII تنبغ في نهاية الأسبوع العاشر. في نهاية التجربة، تم قياس هرمونات وظيفة الغدة الدرقية.

وتم إجراء الفحص السيرجي لقياس الغدة الدرقية بالإضافة إلى دراسة قياسية وتحليل إحصائي.

النتائج: حقن الخلايا الجذعية للنخاع العظمي لفترات قصور الغدة الدرقية أظهر أن بعد 8 أسابيع كانت هناك زيادة عالية من هرمون الثيروكسين وانخفاض كبير في قيمة هرمون محظ الغدة الدرقية في مقارنة مع الغدة الدرقية الناجمة عنها. من ناحية أخرى، بعد 10 أسابيع من حقن الخلايا الجذعية للنخاع العظمي كانت هناك اختلافات طفيفة في هرمون الثيروكسين وهرمون محظ الغدة الدرقية مقارنة مع مجموعة الخلايا الجذعية المستحدثة تجريبياً. أظهر الفحص السيرجي لقياس الغدة الدرقية بعد 8 أسابيع من حقن الخلايا الجذعية للنخاع العظمي تحسن كل фактор من الاستجابة ومع ذلك، بعد 8 أسابيع من حقن الخلايا الجذعية للنخاع العظمي، كان هناك تحسن عام في جميع الحقول التي تم فحصها، والتي أكدت إحصائياً عن طريق تحسين الخلايا، ونسبة الاياف الكولاجين ونسبة المادة الرغبية. من ناحية أخرى، بعد 10 أسابيع حقن الخلايا الجذعية للنخاع العظمي، كان هناك إصدار ملحوظ في جميع الجوانب التي تم فحصها.

الخلاصة: حقن الخلايا الجذعية للنخاع العظمي أستعاد تقريباً الوظيفة والبنية النسيجية للغدة الدرقية بعد 8 أسابيع. ومع ذلك انخفض هذا التحسن بعد 10 أسابيع من حقن الخلايا الجذعية للنخاع العظمي، إذا يوجد بالجرعة الثانية.