Treatment Options for the Patients with Diabetic Retinopathy

Nataliia Sych1,*, Mariya Klunnyk1, Iryna Matiyashchuk1, Mariya Demchuk1, Olena Ivankova1, Andriy Sinelnyk1, Khrystyna Sorochynska2 and Marina Skalozub3

Abstract

Objective: to elaborate the methods of treatment for the patients with diabetic retinopathy (DR) by means of both conventional treatment (medical or surgical) and preparations with extracted fetal stem cells (FSCs) in suspensions containing stem cells of fetal liver, brain and fetal eye.

Materials and methods: A comparative study was conducted on 15 patients with diabetic retinopathy including 7 women and 8 men aged from 27 to 61 years. All patients were administered conventional treatment, laser coagulation and, only over 6 months following the latter, FSCs suspensions containing cells separated from human embryonic tissues of fetal liver, brain and fetal eye were used for treatment of the patients under study.

Results: Authors proved effectiveness and safety of treatment using FSCs in patients suffering from DR. Based on the study we recorded improvements of glycemic and lipids profile as well as overall ophthalmology status among the patients.

Conclusion: FSCs use in complex treatment of patients with DR stabilizes disease compensation, can improve laboratory values and ophthalmology status.

Keywords

Diabetes mellitus; Diabetic retinopathy; Fetal stem cells

Introduction

Diabetes mellitus (DM) is considered to be a significant medical and social problem. A number of patients suffering from DM tend to annually increase and in accordance with data of the WHO over 1 million of patients (which constitute around 2% of population) have been affected in Ukraine [1]. Nowadays, diabetic retinopathy (DR) – a damage of the vessels of eye globe retina remains one of the most frequent and prognostically unfavorable signs in DM patients which often can result in severe loss of vision, blindness and invalidity among the patients. Vision complications are observed in 85% of patients suffering from type 1 DM (T1DM) during 20 years and over. Immediately after type 2 DM (T2DM) diagnosis in people of middle and elderly age in over 50% of likely cases the patients revealed damage of vessels responsible for eyes blood supply [2].

The main causes of DR occurrence are immunology, metabolic, hormonal, blood rheological, hypoxic, genetic, and the other factors which cause lesions of walls in capillaries of eye retina and disturbances of vessels permeability. Swelling of macular zone is the principal cause of eyesight impairment in cases of DR disease. Microangiopathy along with occlusion of capillaries consists in DR pathogenesis [3-6]. Altogether, such disturbances lead to leakage in micro vessels and cause disruption of blood-retina barrier resulting in retinal hemorrhage, exudates and edema, as well as occurrence of edema in macular area. Subsequently, microvascular occlusion and ischemia contribute to “cotton wool” maculae, changes in capillaries, in particular, resulting in development of arteriovenous shunts and neovascularization. Elevation of vascular endothelial growth factor (VEGF) can be regarded as one of the principal angiogenic causes which are implicated within pathogenesis of DR [7,8].

Among the factors of risk for DR disease development one can emphasize the following: persistent control of blood pressure ranges (<130/80 mmHg), constant glycaemia control (HbA1C <7%), undeviating assessment of lipid levels (total cholesterol <4.0 mmol/L, low-density lipoproteins (LDL cholesterol) <2.0 mmol/L, high-density lipoproteins (HDL cholesterol) >1.0 mmol/L) and levels of triglycerides <1.7 mmol/L.

Persistent hyperglycaemia can result in loss of endothelial cells, dysfunction in vascular smooth muscle cells as well as pericytal regions leading to hypoxia. Such changes are remarkable for both T2DM (insulin non-dependent) and T1DM (insulin-dependent). In particular, endothelial cells and pericytial capacity to self-renewal can be affected in case of DM; their turnover potential is subsequently disrupted [3] after which acellular capillaries become non-perfused, and adjacent retina appears to be hypoxic. Such hypoxia environment upregulates vascular endothelial growth factor (VEGF), which contributes to higher vascular permeability resulting in diabetic macular edema [4,5] and, eventually, visual function loss occurs [6] as well as neurodegeneration, including neuronal apoptosis and reactivity of glial cells [9].

At present one can emphasize two main directions (medical and operative) among the existing methods of treatment for DR. Despite of enormous possibilities of therapy for retina by means of laser coagulation in patients suffering from DR, there are still some limitations of this therapy mode as it can lead to a different extent of complications. To absolute contraindications for laser coagulation in patients with DR we can refer large areas of occlusion of capillaries; in particular - within a central zone of ocular fundus, as well as intensive neovascularization, marked glial cells proliferation, vitreomacular traction, preretinal and vitreal hemorrhages. Among complications of laser coagulation in DR, intense swelling of eye macule; hemorrhages to the retina and vitreous body as well as exudative retinal detachments most frequently occur. Though laser coagulation remains the main method of DR treatment for today it does not affect the pathogenetic mechanisms of this disease, but it is rather used only on a local level with the aim of stabilization of pathology process and inhibiting further progressive loss of vision acuity [10].

Collaborators proved effectiveness and safety of treatment for DM
by use of FSCs [11]. So far, no therapy has been established yet that could support regeneration of the damaged vessels of retina resulting from long-term hyperglycemia. Treatment using stem cells might become a feasible option for a purpose of both neurovascular damage prevention as well as promoting retinal damage regeneration, which is supported by the evidence from the recent studies by use of several types of stem cells.

Contemporary approach has been recently developed for treatment of early and moderate stage of DR [12-15]; that is specifically based on capacity of mesenchymal stem cells (MSCs) to production of neurotrophic and neuro-protective factors including a potential of endothelial progenitor cells (EPCs) for repairing the vessels, or likely ability of adipose stromal cells (ASCs) to assure both of the above mentioned functions.

Material and Methods

Study was performed for 15 patients suffering from DR including 7 women and 8 men with their mean age between 27 to 61 years. All patients were given conventional treatment, laser coagulation and, over 6 months following the latter, FSCs preparations with inclusion of separated stem cells were administered for the patients in suspensions containing stem cells, specifically cells of fetal liver, brain and fetal eye.

DR diagnosis was established by means of imaging analysis and with consideration of past history of DM. All patients were tested by way of the following blood laboratory parameters as glycosylated hemoglobin (HbA1c), C-peptide, insulin and fasting blood glucose ranges, including lipidogram profile. The patients have been suffering from T2DM during a period of 3-15 years; average disease duration constituted around 7 ± 3.12 years. From among 15 patients a diagnosis of non-proliferative DR was confirmed in 8 individuals (16 eyes), whereas in 4 patients (8 eyes) – we recorded pre-proliferative stages of DR and 3 of the patients (6 eyes) under study reported proliferative stages of DR disease.

Ocular organs examination in all patients was performed following a generally accepted method. In this respect a complex of measures for diagnosis consisted in: optical coherence tomography (OCT) study, automated perimetry for macula threshold, as well as measurement of macular zone thickness was performed. The patients were examined at baseline, over 3-4 months and following 6-8 months after beginning of FSCs therapy.

Stem cells procedure

All preparations are produced as cryopreserved suspensions of separated stem cells: one of suspensions containing stem cells of fetal liver; the second one – stem cells of fetal brain and the third suspension with inclusion of cells of fetal eye. For this purpose, EmCell collaborators received FSCs material, specifically separated from 8 weeks gestation human fetal liver and fetal brain, following strict aseptic and antiseptic rules in surgery premises. The fetuses were examined at baseline, over 3-4 months and following 6-8 months after beginning of FSCs therapy.

Stem cell procedure was undertaken by transplantation of cryopreserved FSCs in the form of suspension following conventional premedication with infusion of diphenylhydramine 30 mg and prednisolone 15 mg on treatment day 1. Subsequently, all patients were administered a specially prepared solution during the treatment day 2. Patients suffering from DR were performed drip-feed intravenous infusion of fetal liver cells in a volume of 3.2 ± 0.52 mL containing nucleated stem cells ranging from 1.0 to 54 × 10⁶ / mL as well as the levels of progenitor cells CD34+ were 11 ± 2.6 × 10⁵ /mL and amount of CFU constituting from 0.72 ± 0.34 × 10⁶ /mL. During the second day the patients received treatment using suspensions with cells extracted from fetal brain and administered subcutaneously in a volume from 3.2 ± 1.3 mL and suspensions containing fetal eye stem cells which were injected into retrobulbar tissue in amount of 0.5 ± 0.2 mL.

To control overall health, prior to fetal stem cell treatment (FSCT), over 3 and 9 months after therapy general instrumental investigations as well as laboratory tests were performed for the patients. All patients signed their informed consent agreement before they could enter our procedure with FSCs administration.

Our study was conducted according to the ethical guidelines and regulations on good clinical practice, in conformity with Helsinki declaration as well as the ethical standards of practice by use of human fetal tissues samples, approved by the Ministry of Health of Ukraine.

The principal aim of the study was improvement of methods of treatment for the patients with DR by means of a combination with classical therapy using medicines or surgery approach along with transplantation of FSCs in suspensions with separated cells of fetal liver, fetal brain and stem cells of fetal eye.

Statistical analysis

In a process of this investigation doctors applied a complex of statistic programs Statistica 6.0 (StatSoft, Tulsa, Oklahoma, USA). Samplings for all the results obtained were checked-up for a normal distribution by means of Kolmogorov-Smirnov test. With account for the data under study having different laws of distribution, mean difference and sample standard deviation were applied for a purpose of presenting specific characteristics. For calculations we used the methods for parametric (Student’s t-criterion with normal distribution) and non-parametric assessments: U - Mann-Whitney rank test (for independent sample), W - Wilcoxon signed rank (for dependent sample) following a distribution different from normal. Fischer’s exact test we applied to control a significant difference by all qualitative characteristic features. In all clinical cases such such a difference was regarded as significant when it reached the level of
In summary, based on the results presented in Table 1 we can observe significant improvement of lipids profile by 6 months following FSCs therapy. Significant lowering of total cholesterol was recorded over 3 months after FSCT (5.9 mmol/L versus 6.4 mmol/L), \( p < 0.05 \). By 6 months following FSCs therapy significant increase in HDL levels was demonstrable (1.42 mmol/L) compared to the levels baseline (1.18 mmol/L), \( p < 0.05 \) as well as significant decrease of the LDL ranges over 6 months after treatment if compared with the values baseline (3.5 mmol/L and 5.2 mmol/L respectively, \( p < 0.05 \)). The same changes were characteristic for the Atherogenic index (AI) (2.5 and 4.4, respectively, \( p < 0.05 \)).

In accordance with data of OCT, 53.33% of patients revealed higher thickness in the layer of retinal neural fibers throughout a peripapillary zone of both the right and left eye. Thickness of the right eye became higher by 5-7% on average, whereas the left eye thickness increased by 4-6%.

We observed insignificant increase in the mean thickness of macular retinal zone in both eyes (by 8-10 µm) in 60% of patients. According to the results of automated perimetry for Macula Threshold in 73.33% of patients we recorded increased light perception by macular cells both in the right eye and in the left, whereas a reduction of relative scotoma (almost disappearance) was noteworthy in both eyes.

Nowadays, scientists proved few effective methods of therapy for DR focused on prevention, which include tight glycemic and cholesterol control promoting a sustained protection from disease ongoing progression [16-19]. If likely disease progresses to sight-threatening macular edema or proliferative diabetic retinopathy (PDR), laser photocoagulation helps reaching disease regression by means of peripheral retina destruction and thus, reduced demand in oxygen [20-23].

Figure 1: HbA1c dynamics before and after FSCs treatment in DR patients.
Retinal vessels degeneration is considered to be a criterion for earlier DR stage, which corresponds to non-proliferative diabetic retinopathy (NPDR). In the diabetic environment characterized by elevated levels of reactive oxygen species (ROS), vascular progenitors switch to producing pathologic cytokines such as tumor necrosis factor (TNF)-α, IL-8, resulting in higher expression of pathologic inducible nitric oxide synthase (iNOS) instead of endothelial nitric oxide synthase (eNOS) [24]. As a result, diabetic EPCs have reduced bioavailable nitric oxide (NO) due to either decreased activity of eNOS or increased generation of ROS via upregulated NADPH oxidase [25]. NO-mediated signaling events are significant for mobilization of EPCs from the bone marrow and homing in the ischemia zones [26,27]. As shown by the Grant’s group, the function of diabetic EPCs can be partially restored by increased expression of eNOS, either by using NO donors, or by reducing NADPH oxidase-dependent ROS production [28,29]. In addition, increased ROS and reduced bioavailable NO, as well as several other molecular alterations have been detected in dysfunctional diabetic EPCs, including decreased cathepsin L activity [30] and elevated expression of thrombospondin-1 [31].

In regard to proliferative diabetic retinopathy disease at advanced stages, or PDR, contemporary studies prove that high amounts of bone marrow-derived EPCs constitute a major factor in the development of such serious complications as pathologic neovascularization of ischemic tissues. Although increased numbers of circulating CD34+CD45–endothelial colony-forming cells (ECFCs) were found in patients with PDR compared to the control group [32], the same cells could be defective in capacity to migrate towards SDF-1, incorporate into and form vascular tubes. Such results suggest that even though the ECFCs from PDR patients are mobilized into the circulation, they are unable to properly migrate and repair damaged vascular endothelium [32].

Despite such interpretations, cell-based treatment could be presented as effective alternative strategy for the new approaches to treatment of end-stage DR similar to the other ischemic retinopathies. Stem cell therapy is targeted at early and intermediate stages of vascular degeneration by promoting vessels repair, contributing to reversed ischemia, reducing hypoxic or inflammatory signaling, as well as preventing progression to advanced and eyesight-threatening stages of DR [33]. Likely approach is proven to be successful if autologous progenitors are modified to function properly. In recent years, a number of methods have been developed to reverse defects of EPCs in diabetic patients, including improved mobilization for the EPCs.

Table 1: Dynamics of blood lipid levels among the DR patients before and following FSCs therapy.

<table>
<thead>
<tr>
<th>Values</th>
<th>Before treatment</th>
<th>Over 3-4 months</th>
<th>Over 6-8 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cholesterol, mmol/L</td>
<td>6.4 ± 0.02</td>
<td>5.9 ± 0.04*</td>
<td>5.2 ± 0.9**</td>
</tr>
<tr>
<td>Triglycerides, mmol/L</td>
<td>0.35 ± 0.05</td>
<td>0.42 ± 0.05</td>
<td>0.51 ± 0.05</td>
</tr>
<tr>
<td>HDL Cholesterol, mmol/L</td>
<td>1.18 ± 0.02</td>
<td>1.21 ± 0.04</td>
<td>1.42 ± 0.05**</td>
</tr>
<tr>
<td>LDL Cholesterol, mmol/L</td>
<td>5.2 ± 0.12</td>
<td>4.7 ± 0.13</td>
<td>3.5 ± 0.06**</td>
</tr>
<tr>
<td>Atherogenic index (Al)</td>
<td>4.4 ± 0.11</td>
<td>3.8 ± 0.11</td>
<td>2.5 ± 0.07**</td>
</tr>
</tbody>
</table>

* p<0.05 before treatment and following 3 months after FSCs therapy
** p<0.05 before treatment and following 6 months after FSCs therapy
and cells homing with granulocyte colony-stimulating factor (G-CSF) [34] and stromal derived factor (SDF-1) [35], or use of an NO donor to alleviate defects of SDF-1-mediated migration [36]. Subsequently, some data confirm that diabetic EPCs dysfunction could be improved or corrected by treatment using peroxisome proliferator-activated receptor (PPAR)-δ and -γ agonists GW501516 [37], rosiglitazone [38], or atorvastatin. Additionally, it was found that the levels of transforming growth factor (TGF)-β1 were significantly higher both in the EPCs and in the serum of type 2 diabetic patients. TGF-β1 inhibition in CD34+ cells enhanced overall cells survival, NO release, and in vivo vascular reparative ability, suggesting that this approach can be used for improved potential of dysfunctional diabetic CD34+ cells to regenerate cells by means of autologous therapy [36].

If one considers diabetic eye disease at large, stem cells might contribute to future treatment not only for retinal vasculature, but also for diabetic corneal disruptions known as diabetic keratopathy. Severity of likely changes, e.g. of neuropathy, can be correlated with corresponding severity of retinopathy [37]. A significant decrease in the expression of a number of putative stem cell markers in the compartment of corneoinimal epithelial stem cells has been recently established [38]. Since corneal epithelium could be renewed using limbal stem cells, this explains clinically a delay which could be observed at healing for diabetic wounds, for example, after epithelial debridement for vitrectomy [39] or following refractive surgery [40]. Adenoviral gene therapy with overexpression of c-met proto-oncogene and/or silencing of matrix metalloproteinase-10 and cathepsin F promotes stabilized epithelial wound healing due to stem cell marker expression in human organ–cultured diabetic cornneas [38-41]. It is essential that gene therapy by means of limbal stem cell niche could only produce the same normalization of expression for stem cell marker and healing of lesions [42]. In the future, gene therapy or replacement of ailing stem cells by the cultured normal cells similar to those made from iPSC [43] could become an alternative therapy option for alleviating diabetic corneal disease.

To summary everything above, fetal stem cells can facilitate new trends for retarding DR progression and, simultaneously, alleviating the symptoms of DR disease. Nowadays, their use is supported by an evidence of stem cells effect at early stages of this disease; long before the onset of PDR. Some of these cells by way of secreting special growth factors can serve for neuroprotection inside of diabetic retina. New strategies to normalize functions of diabetic progenitors offer ways to use them for autologous therapy. Endothelial progenitor cells are used for prevention and/or repair for capillary closure and will facilitate reduction in pericyte dropout. Established technologies by administration of ESCs and iPSC can also help generating bankable and renewable sources of stem cells being capable for differentiation in order to raise regeneration of retinal cells in a diabetic patient.

Conclusion

Use of fetal stem cells in combined treatment of the patients with DR can be characterized as safe and effective method contributing to compensation of carbohydrates and lipids metabolism among the patients.

The following significant clinical effects of FSCs have been observed:

1. Syndrome of early post-infusion effects.
2. Advantages for overall physical and emotional states of the patients were recorded in 86.7% of cases.
3. Advantages in regard to blood glucose ranges.
   Gradual reduction in fasting blood glucose levels, postprandial glycemia and HbA1c have been initially demonstrable among the patients as early as by 3-4 months following FSCs administration; whereas the same values shown a significant lowering only by 6-8 months after FSCs administration.
4. Hypolipidemia therapeutic effects. We have recorded a significant improvement in parameters of lipids profile over 6-8 months after treatment administered.
5. Ophthalmology status among the patients. In conformity with results of OCT enhanced thickness of retinal neural fibers in peripapillary zone of the eye was characteristic.

We also observed insignificant increasing in moderate thickness of macular region of eyes retina bilaterally (by 8-10 µm) in among 60% of patients. Due to the results of automated perimeter for Macula Threshold, increased light perception by macular cells in both eyes was observed in 73.33% of individuals.

References


Author Affiliations

1Clinical Department, Cell Therapy Center EmCell, Kyiv, Ukraine
2Stem Cells Bank, Cell Therapy Center EmCell, Kyiv City, Ukraine
3Laboratory and Biotechnology Department, Cell Therapy Center EmCell, Kyiv City, Ukraine

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