Stem Cell Ophthalmology Treatment Study (SCOTS) for retinal and optic nerve diseases: a preliminary report

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Abstract
In this report, we present the results of a single patient with optic neuropathy treated within the Stem Cell Ophthalmology Treatment Study (SCOTS). SCOTS is an Institutional Review Board approved clinical trial and is the largest ophthalmology stem cell study registered at the National Institutes of Health to date- www.clinicaltrials.gov Identifier NCT 01920867. SCOTS utilizes autologous bone marrow-derived stem cells in the treatment of optic nerve and retinal diseases. Pre- and post-treatment comprehensive eye exams were independently performed at the Wilmer Eye Institute at the Johns Hopkins Hospital, USA. A 27 year old female patient had lost vision approximately 5 years prior to enrollment in SCOTS. Pre-treatment best-corrected visual acuity at the Wilmer Eye Institute was 20/800 Right Eye (OD) and 20/4,000 Left Eye (OS). Four months following treatment in SCOTS, the central visual acuity had improved to 20/100 OD and 20/40 OS.

Key Words: stem cells; optic nerve; optic neuropathy; ophthalmology; bone marrow-derived stem cells; blindness; visual loss


Introduction
Optic nerve disease is complicated and there are a number of pathophysiologic mechanisms that can lead to retinal ganglion cell impairment or death. Compression, ischemia, inflammation, trauma and hereditary neuronal and mitochondrial diseases are among the causes of optic nerve dysfunction and visual loss. There is substantial support for the use of bone marrow-derived stem cells (BMSCs) in neurologic and ophthalmic diseases as outlined by Ng et al. (2014). Transplanted mesenchymal stem cells (MSCs) have shown significant neuroprotective effects in preclinical models of glaucoma as suggested by Li et al. (2009). After intravitreal transplantation in a rat model of glaucoma, MSCs survived for at least 5 weeks within the vitreous body and some of them migrated into the host retina as reported by Johnson et al. (2010). MSCs from the trabecular meshwork and conjunctivae have produced photoreceptor-like cells in vitro as per Nadri et al. (2013). BMSC transplantation can increase ganglion cell survival in animal models of transient ischemia. Increased survival has also been demonstrated in an ocular hypertension model. These observations are in agreement with Yu et al. (2006).

BMSCs include hematopoietic stem cells and MSCs. MSCs were originally identified in the bone marrow and are also known as marrow stromal cells. These are the adult population of stromal progenitor cells of mesodermal origin that have high proliferative and differentiation capacities and can provide neurons and glial cells. They have shown neuroprotective and immunomodulatory effects and have an important role in the treatment of nervous tissue disease including retinal and optic nerve diseases through brain-derived neurotrophic factor as reported by Wilkins et al. (2009).

Methodology of SCOTS
SCOTS, the Stem Cell Ophthalmology Treatment Study, is the largest ophthalmology stem cell study registered at the National Institutes of Health: Identifier NCT Number 01920867. SCOTS is an open label, non-randomized, efficacy study. There is no placebo or sham arm. All patients enrolled in the study meet eligibility criteria and receive active treatment. Bone marrow aspirated from the posterior iliac crest is separated to provide BMSCs within the stem cell concentrate.

Inclusion criteria for SCOTS include that patients should
- Have objective, documented damage to the retina or optic nerve unlikely to improve OR.
- Have objective, documented damage to the retina or optic nerve that is progressive.
- AND have less than or equal to 20/40 best corrected central visual acuity in one or both eyes AND/OR an abnormal
visual field in one or both eyes.

- At least 3 months post-surgical treatment, intend to treat any ophthalmologic disease and are stable.
- If under current medical therapy (pharmacologic treatment), a retinal or optic nerve disease be considered stable on that treatment and unlikely to have visual function improvement (for example, glaucoma with intraocular pressure stable on topical medications but visual field damage).
- Have the potential for improvement with BMSC treatment and are at minimal risk of any potential harm from the procedure.
- Be 18 years of age or older.
- Be medically stable and able to be medically cleared by their primary care physician or a licensed primary care practitioner for the procedure. Medical clearance means that in the estimation of the primary care practitioner, the patient can reasonably be expected to undergo the procedure without significant medical risk to health.

Exclusion criteria include:

- Patients who are not capable of an adequate ophthalmologic examination or evaluation to document the pathology.
- Patients who are not capable or not willing to undergo follow up eye exams with the principle investigator or their ophthalmologist or optometrist as outlined in the protocol.
- Patients who are not capable of providing informed consent.
- Patients who have significant health risk.

There are three arms of SCOTS with the type of treatment chosen based on the degree of visual loss, etiology of visual loss, associated risk factors for the treatment arms and the patient’s medical risk status. Bilateral treatment is provided assuming both eyes meet eligibility requirements. As these are autologous stem cells, no immunosuppression is required. An FDA cleared class II medical device is used to separate the bone marrow aspirate into a stem cell concentrate. This concentrate has averaged 1.2 billion total nucleated cells including MSCs in approximately 14–15 cm³ of concentrate. Retrobulbar injection consists of 3 cm³ of concentrate, subtenons injection 1 cm³ of concentrate, intravitreal injection 0.05 cm³ of concentrate, subretinal injection approximately 0.1 cm³ of concentrate, intra-optic nerve injection approximately 0.2 cm³ of concentrate, and intravenous injection the remainder of the concentrate.

Arm 1 consists of retrobulbar and subtenons injection, followed by intravenous infusion, of stem cell concentrate. Patients with ophthalmic conditions which preclude safe or effective utilization of intravitreal injection of concentrate, such as the presence of silicon oil, may be offered Arm 1 if meeting inclusion criteria. Arm 2 consists of retrobulbar, subtenon and intravitreal administration, followed by intravenous infusion, of concentrate. Patients meeting inclusion criteria with visual acuity between 20/40 and 20/200 in one or both eyes and/or visual loss may be offered Arm 2. Arm 3 is reserved for retinal and optic nerve patients with severe visual loss meaning acuity of 20/200 or worse in at least one eye. Typically patients admitted to Arm 3 have much poorer vision. Arm 3 consists of the better-seeing eye receiving the same treatment as Arm 1 or more typically, Arm 2, and the eye with more extensive visual acuity loss receiving a core pars plana vitrectomy with subretinal or intra-optic nerve injection of concentrate followed by intravenous infusion of stem cells. Monocular patients are not eligible for Arm 3. Follow up is required at 1, 3, 6 and 12 months post treatment with reporting of the eye exam results to the principal investigator and study director.

The SCOTS procedure is patient funded and performed under general anesthesia. Treatment is provided in a fully licensed ambulatory surgical center in Coconut Creek, Florida, USA. All human investigations were performed with informed consent. The study was reviewed and approved by our Institutional Review Board. The study was registered at the National Institutes of Health.

Clinical History and Course

The patient is a 27 year old, right-handed female who had normal vision until June 2009, when she developed decreased vision in each eye and pain with eye movement. Over the next several weeks she developed severe, bilateral visual loss. She initially demonstrated swelling of the right optic nerve and ultimately developed significant bilateral temporal pallor with centrocecal scotomas. Differential diagnosis included neuromyelitis optica spectrum disorder and Leber’s hereditary optic neuropathy. There was a history of accidental carbon monoxide poisoning from a leaking boiler 16 months prior to the loss of vision, but given the lengthy intervening period, this was thought not to be the etiology of the present visual loss. The patient was evaluated for neuromyelitis optica or Devic’s syndrome and was negative for neuromyelitis optica immunoglobulin G (NMO-IgG) on two separate tests performed at The Johns Hopkins Hospital. Testing for Leber’s hereditary optic neuropathy was negative. On fluid attenuation inversion recovery (FLAIR) imaging, the MRI showed mild middle cerebellar peduncle hyperintensity. She received treatment with methylprednisolone but the vision continued to decline. Additional brain MRIs were normal without demyelinating lesions. The final diagnosis was idiopathic bilateral optic neuritis leading to bilateral optic neuropathy.

Dr. Malkin examined the patient on January 21, 2010. The past medical history was significant for a history of asthma, migraines, and memory loss secondary to carbon monoxide exposure in the past. The best corrected visual acuity was 5/400 right eye (OD) equivalent to 20/1,600 with eccentric fixation, and 5/700 left eye (OS) equivalent to 20/2,800. Contrast sensitivity was 1.00 log units.

On March 21, 2013, the patient’s visual acuity was 5/225 OD, equivalent to 20/900 with eccentric fixation, and 5/300 OS equivalent to 20/1,200 with eccentric fixation. Contrast sensitivity was moderately reduced to 1.28 log units.

On January 16, 2014, the visual acuity was 5/200 OD and 5/400 OS. Contrast sensitivity was 1.32 log units. Biomicroscopic examination was unremarkable and fundus
examination revealed 3° pallor of the optic nerve bilaterally (OU).

Neuro-ophthalmology evaluation at Wilmer Eye Institute on February 5, 2014, demonstrated a best corrected visual acuity of 5/200 OD with eccentric fixation and 1/200 OS with eccentric fixation. The pupils were equal, round and slowly reactive to light in both eyes (OU). The extraocular movements were unaffected. The biomicroscopic examina-
tion was unremarkable OU. The fundus examination was significant for 3+ optic nerve pallor OU.

The patient was accepted into SCOTS in early February 2014, and underwent the preoperative assessment and informed consent with Dr. Jeffrey Weiss on March 3, 2014. Preoperative testing demonstrated a visual acuity of Count Fingers at 1 foot OD and Count Fingers at 2 feet OS. The biomicroscopic examination was unremarkable. Fundus examination was significant for pale optic nerves OU. Humphrey automated visual field testing demonstrated significant loss OU (Figures 1, 2).

She was approved for Arm 3 of SCOTS because her vision

Figure 2 Preoperative visual field of the left eye.
was 20/200 or less in at least one eye. This consists of retrobulbar injection, subtenon injection and intravitreal injection of BMSC for the right eye (OD), and vitrectomy and direct intra-optic nerve injection of BMSC for the left eye (OS), followed by intravenous infusion. She underwent the procedure uneventfully on March 4, 2014. On March 7, the vision was Count Fingers 5 feet/ Pinhole, Count Fingers 6 feet OD, and Count Fingers 2 feet/ Pinhole, Count Fingers 3 feet OS.

Side effects included some initial tearing and conjunctival ecchymosis. There were no complications.

Three months postoperatively, the patient reported that she was able to "find spaces in the scotomas". She was examined by Dr. Malkin on August 5, 2014. On that date, the patient’s visual acuity was 5/300 OD and 20/40–2 OS. The pupils were equal and reactive to light. The optic nerve was graded as 3+ pallor OD and 2+ pallor OS.
On August 21, 2014, approximately 6 months following the SCOTS procedure, she again underwent neuro-ophthalmology evaluation at the Wilmer Eye Institute. On examination, the uncorrected visual acuity was 20/100 OD and 20/40 OS. She was now Jaeger 4 (J-4) OS (near vision). Humphrey automated visual field testing (Figures 3, 4) performed at the Retina Associates of South Florida demonstrated post-operative improvement OU compared to the pre-operative studies. On March 11, 2015, 12 months following the procedure, the uncorrected visual acuity was 20/100/
JNW designed the study, performed the research, collected the data, interpreted the data, wrote and revised the paper. SL designed the study, collected the data, interpreted the data, wrote and revised the paper. AM collected the data. All authors approved the final version of this paper.

Conflicts of interest: None declared.

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