Radiobiological Study of Retinal Microvessel Proliferation in Diabetic-like Rat Model

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Abstract: Progressive evolution of retinal vascular in diabetic retinopathy leads to blindness in up to 8,000 patients yearly. The major purpose of this investigation was to determine proton radiation dose response of the eye's retinal vasculature in the hypergalactosemia induced rat model of diabetic-like retinopathy and gain insight of possible role of proton radiotherapy in controlling diabetic retinopathy. A single dose range of proton radiation (8, 14, and 20 Gy) was delivered to one eye of each rat at 4 months following induction of hypergalactosemia. The opposite eye of each rat, which was not irradiated, showed normal progression of retinopathy. Stereologic techniques were used to quantify tissue parameters in situ in a retinal digest preparation that allowed unobstructed access to the vasculature. 15 months following 50% galactose diet, characteristic histopathological lesions of retinopathy such as capillary endothelial cell proliferation, capillary closure, capillary microaneurysms, pericyte loss developed in non-irradiated eyes. The endothelial cell densities for rat receiving 50% galactose diet were significantly higher than that of control (p<0.05). Proton irradiation inhibited significant endothelial cell proliferation at dose from 14Gy to 20Gy (p<0.05), yet not diminished pericyte loss at current dose schedule. These findings indicated beneficial effects of proton radiation on hypergalactosemia induced diabetic-like retinopathy. Our study should have an impact on further studies to optimize radiation treatment strategies for diabetic retinopathy.

Keywords: Proton radiation, retinal microvessel, diabetic-like model, proliferation, stereology.

INTRODUCTION

Overview

Proliferative vascular disorders of the eye are a major contributor to blindness in both diabetic retinopathy and subfoveal neovascular membrane (SNFM) formation [1-3]. Both diseases are likely to increase in the coming two decades as the "baby boom generation" enters its 60s and 70s. Diabetic retinopathy shows a progressive evolution of retinal vascular changes that leads to blindness [4].

Published reports of LLU clinical Phase I/II dose escalation study (efficacy and safety) for therapy of subfoveal neovascular membrane formation (SFNM) report arrest of the progression of the visual loss and control of the membrane formation in 90% of 300 patients with an average three year follow-up [5-7]. Thus, proton radiotherapy has become a promising, innovative alternative strategy for controlling vision loss due to SFNM.

Microvessel Changes in Response to Irradiation

The dose dependent loss of microvessel length and cell density with vessel dilatation characterized the radiation changes in microvessels of swine dermis and in patients [8-11]. A dose and time dependent loss of endothelial cells, but not of pericytes was also noted [12-15]. Vessel stranding occurred, but not tissue necrosis.

Progression of Diabetic-Like Retinopathy

Among the changes considered to document this proliferation are the presence and increasing numbers of microaneurysms, hypercellular shunts, increased number of microvessels, and increased number of endothelial cells focally in the central retina [16-19]. These are logical associations but cellular proliferation has not been documented as such. The formations of pericyte ghosts, loss of pericytes and endothelial cells with the formation of many acellular vessel strands were documented during the first 7 months following galactose feeding [20-24].

Although galactosemic rats are not diabetic [25] -- the insulin levels are physiological and glucose continues to be metabolized -- they are an excellent preclinical animal model of diabetic-like retinopathy. The etiologic agent is galactitol which is not metabolized and increases to high levels in the blood. Galactation of proteins is produced in cells resulting in deposition of glycans (galactan) in basement membranes. Galactitol is also osmotically active producing cataracts and rupture of pericytes [26]. Fluorescein angiography (in dogs) [27, 28] and retinal digests document a symptom producing increase in number of vessels (branching), formation of microaneurysms, vessel tortuosity and shunt formation by 16-24 months [29-31]. The microaneurysms and cellular shunts occur, but they do not occur uniformly in all animals or in time or throughout the whole retina. The highest density of proliferative changes was found in the posterior retina surrounding the optic nerve from several studies [32].

The objective of the investigation was to determine the proton radiation dose response of the eye's retinal vasculature and gain insights into the radiobiologic and therapeutic
mechanism of action of proton radiation. Our preclinical model is hypergalactosemia induced diabetic-like retinopathy in laboratory rats. The morphological changes produced are similar to those reported for diabetic retinopathy in humans. This model was selected as the bioassay for diabetic retinopathy because it would enable us to anticipate the possible effects in humans.

Stereologic techniques were used to quantify tissue parameters \textit{in situ} in the retinal digest preparation that allowed unobstructed access to the vasculature. Endothelial cell and pericyte cell number were determined; vessel density, microvessel profile area and length were measured.

**MATERIALS AND METHODS**

**Animals**

A total of 90 weanling male Sprague-Dawley rats, each weighing about 50 grams, were purchased locally and maintained, three per cage, under constant ambient temperature of 65° F with a twelve-hour day/night cycle. Water and a diet with 50% content of galactose or 50% starch were available \textit{ad libitum}. Rats’ body weights were monitored at monthly intervals and the animals were observed daily for signs of radiation-related toxicity. Rapid CO2 asphyxiation was performed in compliance with the NIH Guide for the Care and Use of Laboratory Animals.

**Proton Irradiation**

Irradiation was performed 4 month following 50% galactose or 50% starch diet. Each eye was irradiated with a modulated and shaped 100-MeV proton beam, using dose schedules of 0, 8, 14 and 20Gy. The dose was chosen from our previous experience and clinically dose schedule [33]. Rats were positioned in the path of the beam using a cardboard holder after being anesthetized with 0.3 cc Ketamine:Xylazine mixture (5:2) \textit{via} intraperitoneal injection. Field localization is achieved using a 0.8mm-diameter light field projected through the beam nozzle to the (iso)center. Protons were delivered in 300-millisecond pulses every 2.2 seconds, yielding an average dose rate of 7 Gy/min.

**Retinal Digest Preparation**

The rats were euthanasia 15 month following proton irradiation (19 month following the diet). The retina was removed from eyes and fixed at room temperature for at least 4 days by immersing the whole eye in 4% w/v paraformaldehyde in 50mmol/l sodium/potassium phosphate buffer with 4% sucrose at pH 7.2 using Laver’s detailed approach [34]. The entire retina was mounted on the microscope slide.

**Stereology**

The dose response of the retinal digest microvasculature was determined using stereologic analysis. Digest was used for stereologic enumeration on an Olympus Computer Assisted Stereological Toolbox system (The Olympus C.A.S.T.-Grid system, Denmark). In order to have an equal probability that any part of the retina was sampled, the retinal digest perimeter was drawn. Then, using the stereological software grid system stepping function, the surface area was divided into a designated number of equal area “pixels” or steps. The number was designated by determining the number of steps required to permit a fixed number of counts or intersections to be made. Generally, 200 steps (counting areas) are sufficient. A series of point, line, and area probes calculations was used to determine area (points), length (line probes) and cell enumeration (area probes or counting frames of known area) endothelial cell and pericyte density.

**Endothelial Cells and Pericytes**

The endothelial cells are distinguished from pericytes using criteria described in the pilot study [35]. The endothelial cell faced the lumen; the pericytes were found outside the basement membrane. They were smaller and more deeply stained than endothelial cells.

**Statistical Methods**

An 80% power analysis indicated that for a definitive statistically conformal protocol at least 6 animals were used for statistical comparison in each irradiated group and in the unirradiated group. A statistical comparison between parameter means can be made at control, 8Gy, 14Gy and 20Gy. Results were subjected to statistical analyses including one-way analysis of variance (ANOVA) and Tukey’s HSD (honesty significant difference) test using SigmaStat™ version 3.5 software (SPSS Inc., Chicago, IL). A p value of < 0.05 was selected to indicate significant differences among groups.

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**Fig. (1).** Hypergalactosemic-induced microaneuysms in rat retinas from rats fed 50% galactose for 19 months (15 months after radiation) (Original magnifications 40X). A: Cylindrical type, this microaneurysm occurred near the middle of the capillary plexus. B: Irregular type, this microaneurysm occurred in close association with the artery.
RESULTS

Microaneurysm

Microaneurysm formation (Fig. 1) was observed around 15 to 19 months in galactose-fed rats. Different types of microaneurysms have been noticed in our study. Cylindrical or irregular type with vessel dilation was shown in Fig. (1A) and Fig. (1B) respectively. Fewer microaneurysms were seen in galactose-fed rat received irradiation. A proliferation of endothelial cells and focal dilation of capillaries resulted in microaneurysms. By contrast, other capillaries which were adjacent, maintained a normal diameter or lost their endothelia cell and pericytes, and became acellular, non-functional stranding. Vessel proliferations in 50% galactose-fed rat were shown in central retina surrounding optic disc (Figs. 2A, 2B).

Microvessel Morphology Changes

Changes in the retinal vasculature following radiation and the 50% galactose diet are shown in Fig. (3). In the control retina (Fig. 3A), the normal capillaries had a uniform caliber and smooth contour. Two populations of cells were abundant in the vessel wall: endothelial cells and pericytes. Retina from galactose fed rats showed irregular capillary contour with stranding. (Fig. 3B) Vessel walls were hypocellular in pericytes. Staining variations were noted between the control and galactose retinal vessel. The variations were attributed to variations in the thickness of basement membrane. Changes in contours may be explained from loss of pericytes.

Microvessel Cell Population

The retinal vessel digest provided excellent two-dimensional visualization of the microvasculature of the rat retina. Following irradiation, the total retinal pericyte counts in rats receiving 50% galactose diet were significantly lower than the counts for the 50% starch diet (p<0.05) following 14 and 20 Gy of proton irradiation at 15 months (Fig. 4). The slope of the pericyte cell population regression line from rats receiving 50% galactose was greater than that for 50% starch diet. At 15 months, the endothelial cell density for non-irradiated control rat receiving 50% galactose diet was significantly higher than that starch diet (p<0.05). Endothelial
cell densities in galactose fed non-irradiation control rats were significantly high than that of rats fed with galactose diet receiving 14 and 20Gy of irradiation (p<0.05) (Fig. 5). Dose dependent cell loss was seen at galactose-fed rats receiving 8 to 20Gy of proton radiation.

Fig. (4). Total pericytes in retina of age matched control eyes and eyes received irradiation 15 months earlier of starch-fed and galactose –fed rats. At 15 months following irradiation, pericytes in galactose –fed retina were significantly lower than starch-fed rat at all dose (0, 8, 14 and 20Gy) (p<0.05).

Fig. (5). Endothelial cell densities in retina of age matched control eyes and eyes received irradiation 15 months earlier of starch-fed and galactose –fed rats. a Endothelial cell density in galactose fed non-irradiation control rats significantly high than that of starch fed rats. b Endothelial cell density in galactose fed non-irradiation control rats significantly high than that of rats fed with galactose diet receiving 14 and 20Gy of irradiation (p<0.05).

Microvessel Length

The vessel length densities were not significantly different among groups fed with 50% galactose diet or rats fed 50% starch diet following 8Gy, 14Gy and 20Gy of proton irradiation (Fig. 6). The tendency of dose dependent vessel loss was shown in galactose rats following irradiation.

Retina Area

The total retinal area increased with age in starch–fed rats, ranging from 42 to 79 mm² at 1 to 15 months following irradiation. Retinal growth was arrested in rats fed with galactose (Table 1). From 6 to 15 months, the difference in retinal area between rats fed with galactose and starch was significant (P<0.05).

Fig. (6). Microvessel lengths density in the 50% starch-fed or galactose-fed retina 15 months following irradiation at each dose schedule.

DISCUSSION

Our main objectives of this lab were to study and quantify the architectural and population changes in microvessels after radiation. Our previous emphasis was directed to the quantification of normal, quiescent microvessels and their population in the rat retina model. A progressive time and dose-dependent endothelial cell loss were seen over 15 to 24 months following proton irradiation at 8, 14, 20 and 28Gy dose schedules. No proliferation was documented over two year observation period. Rat retinal vasculature was proved to be a slow responding, non-proliferative tissue [35]. It raised question of radiation dose response of proliferative microvasculature. The main focus of the present study was to quantify proton dose response of proliferative retinal vasculature in the hypergalactosemia induced rat model of diabetic-like retinopathy.

The hypergalactosemia induced diabetic-like retinopathy produces morphological changes similar to those reported for man. It was selected as the bioassay because it would permit the investigators to anticipate what the possible role would be in irradiating man. The hypergalactosemia induced rat model was also selected because it permitted defining a generic model that could give insight into the response of proliferating microvasculature.

While it is convenient and useful to characterize and combine the morphological changes into a descriptive sequence, it remains an oversimplification. Certainly, the initial cell loss, vessel stranding (degenerative phase) ending in a designated distorted vasculature does not confirm that a hypoxic environment is created. It is, however, attractive to associate these changes as etiologic for the subsequent “regenerative” or proliferative phase characterized by the formation of microaneurysms, cellular shunts, vessel dilation, increased number and length of microvessels and endothelial cells. While these changes may represent proliferation, it is not known if the process results from angiogenesis or a vasculogenesis resulting from circulating angioblasts (stem cells). Equally important, there is no explanation for the
The close proximity of these changes to large vessels or for the unequal distribution of change in the retina.

The study was well designed and the morphologic parameters could be quantified. This permitted distinguishing the total cellularity in the retina and the change of area involved. Also, it was possible to distinguish a difference between a rapid cell loss of 2-4% of cells per day of an acutely responding proliferation from a slowly proliferating population losing 0.1% of cells per day. These parameters allowed a decision to be made about the efficacy of the irradiation.

In this present study, at four months, endothelial cells and capillary density due to galactose diet were increased compared to that of starch diet. Following irradiation, dose response for galactose–altered retina is significantly greater than starch-fed control retina. It was suggested that proliferative stimuli were active at that time.

Subsequent to selective pericyte loss, the other important histopathological lesions are: endothelial cell proliferation, microaneurysms, and capillary closure [36]. Capillary proliferation was seen in our study, but the total endothelial cell and pericytes count from rats receiving 50% galactose diet were decreased compared to their counterparts on 50% starch diet. This could be explained that endothelial cell and capillary proliferation are accompanied by selective endothelial cell and capillary loss in surrounding area. When total number of cells in the retina was counted, increase in cell number was not shown (Fig. 7).

CONCLUSION

1. Galactose diets suppress weight gain in weanlings and arrest the retina growth.
2. Selective pericyte loss is one of the histopathological lesions that are essentially unique to diabetic-like retinopathy. Proton irradiation at current dose does not prevent retinal pericyte loss.
3. Microaneurysms and focal endothelial cell proliferation were seen in galactose fed weanling rats. Relatively fewer microaneurysms were shown in irradiated retina from rats received 50% galactose diet for 4 months before radiation.
4. Irradiation alleviated or inhibited vessel proliferation in hypergalactosemia induced rat retina with dose dependence.

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REFERENCES


