Gene transfer by viral vectors into blood vessels in a rat model of retinopathy of prematurity

Itay Chowers, Eyal Banin, Yitzchak Hemo, Rinat Porat, Haya Falk, Eli Keshet, Jacob Pe’er, Amos Panet

Abstract

Aims—To test the feasibility of gene transfer into hyaloid blood vessels and into preretinal neovascularisation in a rat model of retinopathy of prematurity (ROP), using different viral vectors.

Methods—Newborn rats were exposed to alternating hypoxic and hyperoxic conditions in order to induce ocular neovascularisation (ROP rats). Adenovirus, herpes simplex, vaccinia, and retroviral (MuLV based) vectors, all carrying the β galactosidase (β-gal) gene, were injected intravitreally on postnatal day 18 (P18). Two sets of controls were also examined: P18 ROP rats injected with saline and P18 rats that were raised in room air before the viral vectors or saline were injected. Two days after injection, the rats were killed, eyes enucleated, and β-gal expression was examined by X-gal staining in whole mounts and in histological sections.

Results—Intravitreal injection of the adenovirus and vaccinia vectors yielded marked β-gal expression in hyaloid blood vessels in the rat ROP model. Retinal expression of β-gal with these vectors was limited almost exclusively to the vicinity of the injection site. Injection of herpes simplex yielded a punctuate pattern of β-gal expression in the retina but not in blood vessels. No significant β-gal expression occurred in rat eyes injected with the retroviral vector.

Conclusions—Adenovirus is an efficient vector for gene transfer into blood vessels in an animal model of ROP. This may be a first step towards utilising gene transfer as a tool for modulating ocular neovascularisation for experimental and therapeutic purposes.

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In animal models, modulation of retinal neovascularisation can be achieved by administration of various agonists or antagonists to angiogenesis such as vascular endothelial growth factor (VEGF), growth hormone, and others. Additional novel investigative methods applied in the study of ocular angiogenesis are the use of transgenic animals and gene knockout techniques. For example, vascular endothelial growth factor (VEGF) receptors, and fibroblast growth factor (FGF) knockout mice, as well as transgenic mice that overexpress VEGF or FGF have been extensively studied. These animal models have greatly increased our knowledge of the importance of factors such as VEGF or FGF in retinal neovascularisation. However, the establishment of such lines of mice is quite difficult, and this approach cannot be directly used as a therapeutic modality in humans.

An alternative method to study the effects of a gene product is to deliver and express the gene in the target tissue by a viral vector. This approach has been used extensively in the study of ocular diseases, especially retinal degenerations, in animal models. To the best of our knowledge, gene delivery to ocular blood vessels has not been previously described.

The purpose of the present study was to examine the feasibility of gene transfer by four different viral vectors into hyaloid and preretinal blood vessels in the rat ROP model. Furthermore, ocular tissue tropism of these viral vectors was investigated.

Methods

RAT ROP MODEL

Animals were treated according to the ARVO statement for the use of animals in ophthalmic and vision research. Abnormal ocular neovascularisation was induced in newborn rats as described by Penn et al (“rat ROP model”). Postnatal day 1 (P1) Sabra rats and their mothers were raised for 14 days in alternating, 24 hour cycle, hypoxic (10% oxygen) and hyperoxic (70% oxygen) environments. At P14 rats were removed from the incubator to room air. Intravitreal injections were performed 4 days later (P18). P18 Sabra rats that were raised in room air served as controls.

VIRAL VECTORS

Four different viral vectors were used:

(1) Adenoviral vector containing the β galactosidase (β-gal) reporter gene under the control of the cytomegalovirus (CMV) promoter (Ad5CMVlacZ). This adenovirus vector is E1A defective. The replication defective virus was propagated in 293 cells.
(2) The herpes simplex virus type I (HSV) vector contained the reporter gene β-gal (tkL-TRZ1) under control of the murine leukaemia virus (MuLV) LTR promoter inserted at the thymidine kinase (tk) gene locus. The virus is replication competent in dividing cells and it was grown on CV1 monkey cells.

(3) The vaccinia expression vector, encoding the β-gal gene under control of the vaccinia early promoter (vSC9). The virus is replication competent and was propagated in HeLa cells.

All the DNA viruses were titrated (cell forming units, CFU) in parallel on HeLa cells using the X-gal staining technique.

(4) The β-gal encoding recombinant retrovirus (pCLMGF-LacZ) was constructed using the pLXSN vector with the β-gal gene under control of the MuLV LTR promoter. The transfer vector was packaged by co-transfection with the pCL-Eco packaging construct in 293 cells.

Virus supernatants were collected after 2 days and the virus titre (CFU) was determined using NIH-3T3 cells and X-gal staining.

INTRAVITREAL INJECTIONS

At P18 rats were anaesthetised in an ether chamber. Borosilicate glass pipettes (1.2 mm external diameter, 0.69 mm internal diameter, Sutter Instruments Co, Navato, CA, USA) were pulled to form 50 µm calibre tips by an electrode puller (P-97, Sutter Instruments Co, CA, USA) and were connected to a microinjection unit (PLI-100, Medical Systems Corp, Greenvale, NY, USA) for intravitreal injections. All injections were performed under magnification, using a binocular microscope and a micromanipulator.

A volume of 1 µl containing the same cell forming units (10^5 CFU) of one of the viral vectors (adenovirus, herpes simplex, or vaccinia) or 10^4 CFU of the retroviral vector were injected intravitreally to 10 ROP rats via the pars plana in one eye. Two control groups were studied. The first included 10 P18 ROP rats in whom saline was injected instead of the viral vectors. The second control group included 25 P18 rats that were raised in room air. The four viral vectors and saline were each injected intravitreally in five such rats.

Figure 1  Macroscopic photographs of eyes enucleated from normal and ROP P20 rats after intravitreal injection of β-gal carrying viral vectors. ROP rats underwent intravitreal injection at P18 with adenovirus vector (10^5 infectious units) (A) (original magnification ×3), saline (C) (original magnification ×3), or herpes vector (10^5 infectious units) (D) (original magnification ×5). Normal P18 rats were injected with adenovirus vector (10^5 infectious units) (B) (original magnification ×4). ROP and normal rats were also injected with vaccinia and retroviral vector (not shown). After enucleation, the cornea, iris, and lens were removed and the eye cup was fixed in 0.2% glutaraldehyde, 2% formaldehyde, 2 mM MgCl2, 0.05M NaPO4 buffer (pH 7.4), followed by overnight X-gal staining. Adenovirus injected eye (A) shows extensive β-gal expression (blue stain) in the hyaloid system; notice the paucity of retinal staining. Normal P20 rat eye injected with adenovirus vector shows X-gal staining of the remnants of the hyaloid system as well as areas of retinal staining (B). Herpes injected eye shows punctate staining of the retina (D). Control, saline injected ROP rat eye shows no staining of the hyaloid or retina (C).
X-GAL STAINING
At P20, rats were killed, eyes were enucleated, and the cornea and lens removed. The remaining eye cups were fixed in 0.2% glutaraldehyde, 2% formaldehyde, 2 mM MgCl₂, 0.05M NaPO₄ buffer (pH 7.4), washed three times in 2 mM MgCl₂, 0.02% NP40, 0.05M NaPO₄ buffer, and incubated overnight at 37°C with 5 mM potassium ferrocyanide, 2 mM MgCl₂, and 20 mg/ml X-gal (5-bromo-4-chloro-3-indolyl β-D-galactopyranoside) in 0.05M NaPO₄ buffer. Eye cups were then either fixed in formalin, embedded in paraflin, cut into 4 µm sections and stained with haematoxylin and eosin or, alternatively, the retina was separated and a whole mount preparation was examined under a microscope.

Results
Rats that were maintained until postnatal day 14 in alternating hypoxic and hyperoxic conditions manifested at P20 the ocular blood vessel abnormalities previously described in this animal model of ROP. Marked congestion and tortuosity of the retinal and iris vessels were observed along with preretinal haemorrhages and persistence of the hyaloid system. Preretinal vessels (“neovascular tufts”) emerging from the retinal circulation were detected in histological sections in two eyes. In rats maintained in room air, the normal postnatal process of hyaloid system regression and retinal vascular development was observed. At P20, only remnants of the hyaloid system could be seen.

The most extensive expression of β-gal in blood vessels occurred following intravitreal injection of the adenovirus vector (Ad5CMVlacZ) to ROP rats. There was marked X-gal staining of the hyaloid system in all 10 animals of this group (Fig 1A). Macroscopically, the whole hyaloid system seemed to express β-gal, resembling large blue “tree trunks” inserting into the optic disc (Fig 1A). Microscopically, histological sections and whole mount preparations showed β-gal expression in the walls of hyaloid blood vessels (Fig 2A and B). By contrast, retinal blood vessels did not show significant levels of β-gal expression. Histological sections revealed preretinal “neovascular tufts” emerging from retinal blood vessels in two of the 10 ROP rats that were injected with adenovirus. In both cases, β-gal expression was seen in these preretinal tufts. Interestingly, the staining seems to stop abruptly at the point at which the vessels originate from the retina. The blood vessels within the retina in this area showed no β-gal expression (Fig 2C). In general, retinal expression of β-gal (as opposed to the hyaloid system) was almost exclusively limited to the area of injection (at the pars plana). Here, neuroretinal elements other than blood vessels, such as the inner nuclear layer and ganglion cells, stained as well. Histologically, retinal structure and cellular components were well preserved in all injected animals.

Vaccinia (vSC9) injection yielded β-gal expression only in segments of the hyaloid system (as opposed to staining of the whole hyaloid system in the adenovirus injected ROP rats) in three of the 10 ROP rats but not in the five normal P18 rats; retinal staining was again mainly limited to the injection site. The HSV vector (tkLTRZ1) expressed β-gal almost...
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In this study, the feasibility of gene delivery to
rats that were injected with saline showed no
X-gal staining or inflammation (Fig 1C).
Interestingly, in these normal rats, gene deliv-
ery into the retina was more extensive as com-
pared with the ROP rats (Fig 1B).
Additional ocular structures that showed
β-gal expression after injection of adenovirus,
vaccinia, or herpes viral vectors were the ciliary
body, pigment epithelium of the iris, and the
corneal endothelium. These three vectors, but
not the retrovirus, caused a severe vitreal and
anterior chamber inflammatory reaction that
was first noted approximately 48 hours follow-
ing injection. Eyes of ROP rats and normal P18
rats that were injected with saline showed no
X-gal staining or inflammation (Fig 1C).

Discussion
In this study, the feasibility of gene delivery to
hyaloid and preretinal blood vessels in a rat
model of ROP was tested. Our results show
that adenovirus vector can efficiently deliver
genes into hyaloid blood vessels in the rat ROP
model. Although the constitutive CMV pro-
motor was used to control gene expression in
this vector, a surprising degree of specificity of
β-gal expression was observed in hyaloid blood
vessels. In the retina, adenov mediated β-gal
expression was limited almost exclusively to
the injection site.

Several factors may contribute to the rela-
tively limited adenovirus vector single cycle
infection of the retina. It is possible that the
inner limiting membrane and the posterior vit-
reous face act as a barrier, physically blocking
infection of the retina itself. Indeed, other
investigators have shown that when retinal
expression is the goal, subretinal injection of
the viral vectors is preferable to intravitreal
injection.25 In human vascular retinopathies,
splitting of the posterior cortical vitreous, or
posterior vitreochorisis, is a common finding
that is usually manifested as two dense vitreous
membranes.25 26 We speculate that a similar
condition may exist in the rat ROP model,
adding another obstacle (apart from the inner
limiting membrane of the retina) between the
vitreous cavity and the retina. Our observation
of more extensive retinal expression of the
trans-gene in the normal rat retina compared
with the ROP rat retina seems to support the
possibility of such preferential accessibility.

Another factor influencing retinal trans-gene
expression is viral vector multiplicity of infec-
tion. Higher concentrations of the vector
would perhaps deliver genes more efficiently to
the retina, and it is possible that the multiplicity
of infection used (which we tried to keep
low in order to decrease levels of inflammation)
contributed to the apparent specificity of
expression in the hyaloid.

Infection by a viral vector such as adenovirus
may damage the cell even without expressing
the trans-gene. Therefore, the relatively tissue
specific gene delivery demonstrated in our
study is encouraging, since when gene delivery
into hyaloid or preretinal blood vessels is the
goal, prevention of infection of the retina is
desirable.

The other viral vectors tested in this study
showed either partial (vaccinia) or no (retro-
virus and HSV) gene transfer to the hyaloid
vessels. The differing pattern of reporter gene
expression by the various vectors is probably
due to differences in their ability to infect spe-
cific tissues, since the promoters driving the
reporter gene are constitutive and do not con-
fer tissue specificity. It should be noted that the
four viral vectors tested have different replica-
cation cycles and different promoters to drive
trans-gene transcription. Nevertheless, when
β-gal enzyme activity was measured by the
ONPG colorimetric test in extracts of a cell
line, mouse NIH 3T3, infected at the same
viral multiplicity, expression levels were very
similar for the three DNA viruses and only
fivefold lower per cell for the retroviral vector
(unpublished data). Therefore, vector tissue
tropism appears to be related to early steps of
infection which may explain the differences in
pattern of expression between the vectors. For
instance, HSV receptors may be missing in vit-
real blood vessels, preventing infection of the
endothelium by this vector.25 26 The retroviral
vector and to some extent the HSV vector
require cell proliferation in the target tissue in
order to achieve infection. The lack of
endothelial cell proliferation in the hyaloid ves-
sels at P18 might be the critical factor contrib-
uting to the failure of gene delivery by these
two vectors. Thus, the results presented in this
report most probably reflect tissue tropism of
the four viral vectors in the eye, and are prob-
ably not due to differences in trans-gene
promoter activity among the viruses.

As opposed to the persistence of the hyaloid
system that was markedly enhanced in our
ROP rats, only two preretinal neovascular
tufts were observed. Therefore, although both ex-
pressed β-gal, conclusions regarding efficacy of
gene delivery to such preretinal tufts cannot be
drawn directly. However, the hyaloid system,
being a vascular system in the vitreous cavity,
simulates some aspects of neovascularisation
on the disc and pre-retinal neovascularisation.
The hyaloid is also intriguing from a develop-
mental aspect, affording a model in which not
only the process of blood vessel growth but also
that of normal vessel regression can be studied.
The hyaloid system, which in humans regresses
before birth, is normally present in the first few
weeks of life in the rat.27 This system is
composed of non-fenestrated capillaries as well
as larger blood vessels, all surrounded by peri-
cytes.28 VEGF serves as a survival factor for the
hyaloid system,29 and in the rat, hyaloid
capillaries continue to show sprouting after
birth.30 This depends on the postnatal age of
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Specific expression of genes in vascular endothelial cells is possible by using vectors in which the genes are under the regulation of an endothelium specific promoter. In the eye, genes such as the 67 kD laminin receptor, that are expressed preferentially in proliferating and not in quiescent retinal blood vessels, have been identified, and their nucleotide sequence has been determined. In the future, gene delivery and specific expression in the endothelium of proliferating ocular vessels can be attempted by keeping the delivered gene under the regulation of an endothelium specific promoter. Such a method can be used experimentally to assess the effect of different genes on blood vessels.

In addition, by using the strategy of suicide gene delivery, ocular neovascular tissue could be preferentially targeted and destroyed. For example, new preretinal (or choroidal) blood vessels may be infected with a vector carrying the herpes simplex thymidine kinase under the control of a promoter that activates genes only in proliferating endothelium. Gene delivery will be followed by treatment with ganciclovir, thereby activating and specifically affecting proliferating blood vessels. Such suicide gene delivery in the eye (into tissues other than blood vessels) has been used to inhibit RPE cell growth in vitro and to treat experimental proliferative vitreoretinopathy in the rabbit.

In conclusion, our study demonstrates the feasibility of gene transfer by viral vectors into hyaloid blood vessels in an animal model of ischemia-induced vitreoretinopathy. Retinal specificity of expression was observed, perhaps because of preferential accessibility of some of the viral vectors into these hyaloid blood vessels. Based on these results, and considering the available techniques in gene targeting, the concept of modulating ocular neovascularisation by gene delivery should be tested.

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