BASIC SCIENCE

Sustained dasatinib treatment prevents early fibrotic changes following ocular trauma

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Abstract

Purpose Posterior ocular trauma and the subsequent fibrotic retinal complication termed proliferative vitreoretinopathy (PVR) are leading causes of blindness in children and young adults. A previous study suggested that changes occurring within the first month post-trauma can lead to development of PVR later. The aim of this study was to examine the effect of dasatinib, a tyrosine kinase inhibitor clinically used to treat chronic myeloid leukemia, on fibrotic changes occurring within the first month following ocular trauma.

Methods A previously established swine ocular trauma model that mimics both contusion and penetrating injuries was used. Dasatinib was administered on days 4 and 18 post-trauma via intravitreal injection of either bolus solution or suspension of a sustained release system incorporated in biodegradable poly (lactic-co-glycolic acid) (PLGA) nanoparticles. Animals were followed up to day 32, and the development of traction full-thickness fold in the posterior retina was assessed.

Results A full-thickness retinal fold extending from the wound site developed in 3 out of 4 control eyes injected with PLGA nanoparticles alone at 1 month. Administration of dasatinib solution had little preventative effect with 6 out of 7 eyes developing a fold. In contrast, dasatinib-incorporated PLGA nanoparticle injection significantly reduced the incidence of fold to 1 out of 10 eyes.

Conclusions Injection of dasatinib-incorporated PLGA significantly reduced early fibrotic retinal changes which eventually lead to PVR following posterior ocular trauma. Thus, our sustained dasatinib release system can potentially be used to both prevent and/or broaden the surgical treatment window for PVR.

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Introduction

Ocular trauma is a leading cause of visual impairment. In particular, penetrating and perforating trauma involving the posterior segment has a high incidence of blindness [1]. Proliferative vitreoretinopathy (PVR) is the main complication leading to loss of visual acuity with incidence ranging from 10 to 50% in penetrating/perforating injuries [2, 3]. Early studies using animals, as well as clinical observation of patients, demonstrated that the vitreous plays a key role in the development of post-traumatic PVR [4, 5]; thus, vitrectomy is the current standard of care following posterior ocular trauma and associated retinal detachment as well as non-clearing vitreous hemorrhage. However, there is still a debate on the suitable timing of vitrectomy and some patients may not have access to operating facilities immediately. In a rabbit penetrating trauma model, delaying vitrectomy for 4–6 weeks had a higher risk of developing PVR when compared to vitrectomy performed within a week following injury [6]. Furthermore, vitrectomy conducted after 28 days post-injury was identified as a high risk factor for developing advanced PVR (Grade C) in patients [2], and many retinal surgeons recommend surgical intervention within 2 weeks of injury. This suggests that changes occurring within this 4-week period can lead to development of PVR at a later time point. Adjunctive therapy has been suggested to reduce the incidence of post-traumatic PVR but only a handful of agents have been tested [7–9], and currently no therapeutic agent is used routinely for this purpose.

In this study, we tested dasatinib, a tyrosine kinase inhibitor used for the treatment of chronic myeloid leukemia, as a potential adjunctive therapy to prevent fibrotic changes occurring within 4 weeks following posterior ocular trauma. Dasatinib prevents traction retinal detachment in a swine PVR model that mimics rhegmatogenous retinal detachment [10], and in vitro studies using cultured retinal pigment epithelial cells and Müller glia cells showed that dasatinib targets matrix contraction, a characteristic cellular function associated in PVR [10, 11]. We created a sustained release system for dasatinib by incorporation into poly (lactic-co-glycolic acid) (PLGA) nanoparticles (Das-PLGA) [12]. In this study, using a swine model of posterior ocular trauma [13], we examined the effect of Das-PLGA on post-injury fibrosis. We observed that intravitreal injection of Das-PLGA prevented fibrotic retinal changes within the initial 4-week period following posterior ocular trauma, thus providing a window for surgical correction of this retinal complication if it subsequently developed.

Materials and methods

Production and characterization of PLGA and dasatinib-incorporated PLGA particles

Production of PLGA and Das-PLGA particles by spray drying method as well as characterization of particle size and release profile has been described in detail elsewhere [12]. The size of particles used for the study had an average diameter of 0.79 μm, and in vitro release profile is shown in Fig. 1.

Full-field electroretinography measurement to determine potential adverse effects of Das-PLGA particles

PLGA is biocompatible and its intraocular use is generally well tolerated [14]. A recent study has, however, demonstrated endophthalmitis following intravitreal injection of PLGA...
particles [15]. Thus, full-field electroretinography (ERG) was utilized to assess potential retinal toxicity of PLGA and Das-PLGA injected intravitreally into swine eyes. The use of animals was approved by the University of Louisville Institutional Animal Care and Use Committee and adhered to the ARVO statement on the Use of Animals in Ophthalmic and Vision Research. Full-field ERG, following the ISCEV standard, was performed on swine as previously described using a UTAS ERG system with a BigShot Ganzfeld Stimulator (LKC Technologies Inc., Gaithersburg, MD) [10]. Intravitreal injection of PLGA (without dasatinib) or Das-PLGA was performed immediately after the ERG measurement (day 0, 0.5 mg particles/eye). Particle injection was repeated on day 14 but at half the amount (0.25 mg particles/eye) to reduce any possibility of adverse effects. Full-field ERG measurements were repeated on day 28 to assess the effect of particles on retinal function.

**Induction of ocular trauma in swine**

Combined ocular contusion/penetrating injuries were induced based on a previously published protocol [13] with some modification. For sedation and induction of anesthesia, a combination of atropine, ketamine, butorphanol, and dexmedetomidine was administered IM, and anesthesia was maintained by isoflurane following intubation. After preparation of a conjunctival flap, contusion injury was produced nasal to the vortex veins, and 6 mm posterior to the limbus with a BB pellet. Subsequently, a fornix-based triangular scleral flap consisting of an incision 6 mm long (3 mm length on each side) penetrating sclera, choroid, and retina was created at the site of impact. Approximately 0.7 ml of prolapsed vitreous was trimmed and removed by Weck sponges, and the wound closed by suturing the apex of the triangular flap, as well as the center of each 3 mm incision. Finally, 0.7 ml of autologous blood was injected into the vitreous through the pars plana.

Bilateral surgery was approved by IACUC based on the following reasons. Published data showed, and confirmed by a preliminary study, that the outcome of retinal changes within the experimental period are limited to development of full-thickness fold in the inferior retina between the wound site and optic disc. As the visual streak of swine is superior to the optic disc, the effect of retinal fold to vision of animals was considered to be minor. Furthermore, haziness caused by blood injection was temporary, and did not cause blinding of animals that prevented basic functions such as eating or socializing. Nonetheless, animals were monitored daily, and carefully examined upon showing signs of pain or distress.

**Drug administration and evaluation of fibrotic contraction**

Post-operative ocular examination involving slit lamp and indirect ophthalmoscopy, and/or B-scan ultrasound, were performed on days 4, 18, and 32. Following ocular examination on day 4, 0.1 ml of (1) PLGA particle suspension (control, without dasatinib, 0.5 mg total particles), (2) dasatinib solution (12 μM stock concentration), or (3) dasatinib-incorporated PLGA particle suspension (Das-PLGA, 0.5 mg total particles) were injected into the vitreous. Injection of 0.1 ml PLGA particle suspension (with or without dasatinib, containing 0.25 mg particles) or dasatinib solution (12 μM stock concentration) was repeated on day 18 after ocular examination. PLGA particles were halved for the second injection to avoid the possibility of vitritis previously reported by others [14]. Animals were euthanized following the final examination (day 32), and eyecups created from unfixed enucleated eyes. Both observations from indirect ophthalmoscopy (prior to euthanasia) and examination of enucleated eyecups were used to determine development of a full-thickness fold and/or retinal detachment posterior to the wound site.

**Histological procedure**

Following examination, eyecups were fixed overnight with 4% paraformaldehyde, dehydrated through a graded series of ethanol followed by xylene, and embedded in paraffin. Paraffin sections were stained for Hematoxylin and eosin (HE) (Surgipath H&E kit, Leica Biosystems, Buffalo Grove, IL), Masson’s trichrome (MT) (Artisan Masson’s trichrome staining kit, Agilent, Santa Clara, CA), and elastic van Gieson (EVG) stain using a Leica automated system (Leica Biosystems). Immunohistochemical staining for tenascin-C (TNC) and alpha-smooth muscle actin (aSMA) were performed using rabbit monoclonal antibody clone EPR4219 (Abcam, Cambridge, MA) and mouse monoclonal antibody
clone ASM-1 (Leica Biosystems), respectively, and BOND polymer refine red detection system (Leica Biosystems).

Statistical analyses

Statistical analyses were performed using Prism version 6 software. Student’s t test was used to compare ERG b-wave amplitude and latency time prior to and after PLGA particle injection. Fisher’s exact test was used to compare difference in number of eyes that developed full-thickness retinal folds among different treatment groups.

Results

ERG response in eyes injected with PLGA or Das-PLGA particles

Injection with either PLGA or Das-PLGA particles (n = 6 each) did not significantly affect full-field ERG responses. Average b-wave amplitudes, as well as latency, in dark-adapted 0.01 and 3 cd s m⁻² (rod and rod/cone mix, respectively) and light-adapted 3 cd s m⁻² (cone response) ERG responses measured on day 28, after injections on day 0 and 14, were not significantly different than baseline (on day 0, Fig. 2). Since there was no evidence of an adverse effect, PLGA and Das-PLGA particles were tested in our ocular trauma model.

Early fibrotic changes following posterior ocular trauma, and effect of treatment

A full-thickness posterior retinal fold (Fig. 3a) developed in 3 of 4 control eyes injected with PLGA particles (Table 1). Retinal folds were observed at day 18 and remained until final examination on day 32 (Fig. 3a).

Injection of dasatinib solution (Das-solution) every other week did not prevent retinal fibrotic changes with 6 of 7 eyes developing a full-thickness posterior retinal fold (Fig. 3b, Table 1). In contrast, intravitreal Das-PLGA significantly reduced the incidence of retinal fibrosis with only 1 of 10 eyes developing a full-thickness retinal fold (Table 1). Most eyes injected with Das-PLGA had no detectable change in the posterior retina (Fig. 3c).
One eye developed signs of sterile endophthalmitis following injection of Das-PLGA. This eye was treated with intra-vitreal dexamethasone, and removed from the study before its completion.

Histological analyses of the wound site

Past studies have shown that the fibrotic proliferation at the entrance site in penetrating injuries results in vitreoretinal traction [4, 5, 16]. H&E, Masson’s trichrome (MT), and Elastic van Gieson (EVG) staining all showed inward extension of a fibroproliferative scar from the wound site (Fig. 4). MT and EVG staining confirmed strong expression of collagens (blue for MT, red for VEG) in scar tissue (Fig. 4). Incidence of scar formation on the retinal surface matched PVR grade in most eyes injected with PLGA or Das-solution (Table 2). Despite the lack of full-thickness retinal folds in 9 of 10 eyes injected with Das-PLGA, scar formation was also observed on the retinal surface in 6 out of 9 eyes (Table 2). We could not identify the wound site in one of the Das-PLGA eye due to damage to the tissue during processing, and thus, this eye was omitted from histological analyses.

Expression of tenascin-C (TNC) (Fig. 5), an extracellular matrix protein upregulated in various fibrotic diseases/complications, was detected by immunohistochemistry in 45% of all eyes used in the study (Table 2). Control showed noticeable expression in 2/4 eyes, and treatment with Das-solution (3/7 eyes) or Das-PLGA (4/9 eyes) had no noticeable effect on TNC expression (Table 2). Expression of aSMA (Fig. 6), a marker for myofibroblasts, matched the PVR grade in most eyes injected with PLGA (control) or Das-solution (Table 2). Eyes injected with Das-PLGA showed a trend for reduced expression of aSMA but the difference did not reach statistical significance (Table 2).

Discussion

PVR is the major complication following posterior ocular trauma that leads to visual disability and frequently blindness. The incidence of post-traumatic PVR depends on a variety of factors such as type of injury (i.e. blunt, penetrating, perforating or rupture), presence of intraocular hemorrhage, retinal tear or retinal detachment [2, 3]. Ryan and colleagues established several animal models to study penetrating trauma [13, 17, 18], and together with observations from clinical specimens [5], concluded that fibroproliferative changes following penetrating eye injuries cause vitreous traction resulting in retinal folds and/or detachment. In this study, we adopted the swine model established by Gregory and Ryan that involves both contusion and penetrating injury, and displays fibrotic changes within 4 weeks post-injury [13]. While we shifted the injury site posteriorly from the pars plana, as in the original study, to include damage to the retina, this difference in the position of the entrance wound did not alter the outcome. Both the current study, as well as the original study [13], resulted in comparable localized, full-thickness posterior retinal folds. Furthermore, the reported time course of retinal

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**Table 1**

| n number | Ctrl (PLGA) | Das-solution | Das-PLGA |
|----------|-------------|--------------|----------|
| 4        | 7           | 10           |
| Full-thickness fold | Yes | 3 (75%) | 6 (86%) | 1 (10%)* |
| No       | 1 (25%) | 1 (14%) | 9 (90%) |

*Significantly different (p<0.05) from control (PLGA)

**Table 2**

| | Ctrl (PLGA) | Das-solution | Das-PLGA |
|---|-------------|--------------|----------|
| Retinal scar formation | 3/4 (75%) | 7/7 (100%) | 6/9 (67%) |
| Tenascin-C positive scar | 2/4 (50%) | 3/7 (43%) | 4/9 (44%) |
| Alpha-SMA positive scar | 3/4 (75%) | 6/7 (86%) | 3/9 (33%) |
fold development (> 8 days) [13] and the incidence of retinal folds at 2 weeks (7 out of 9 eyes) [19] of the original studies were similar to the current study. Despite penetrating injury to the retina/RPE, extensive retinal detachment did not develop in the current study. This may be due to efficient pumping function of swine RPE cells, as evidenced by reattachment of the retina within 3 days of an artificially created total retinal detachment [20], and a viscous vitreous in young swine compared to man.

Injection of Das-PLGA prevented post-traumatic fibrotic changes in the current study. Dasatinib was originally developed as a dual inhibitor of Src family kinases (SFKs) and Abl kinase [21]. However, it has subsequently been shown to inhibit a host of tyrosine kinases including PDGF receptor kinase and c-kit [22]. Dasatinib has been shown to suppress fibrotic changes in various animal models of tissue fibrosis [23–27]. Within the eye, we have previously demonstrated that dasatinib inhibits traction retinal detachment in a swine PVR model involving RPE cell injection [10]. Prevention of RPE cell epithelial-mesenchymal transition, proliferation, and migration by dasatinib was due to inhibition of SFKs [10]. Furthermore, dasatinib significantly reduced matrix contraction by cultured Müller glia cells via targeting SFKs and PYK2, both of which play a key role in phosphorylation of focal adhesion proteins [28]. While similar molecular mechanisms may be playing key roles in the prevention of fibrotic changes observed in the current study, further investigation is required for confirmation.
In the current study, bolus injection of dasatinib solution failed to inhibit the development of a retinal traction fold. This is in contrast to our previous study in which traction retinal detachment was prevented in a model that mimics PVR following rhegmatogenous retinal detachment by bolus injection of dasatinib solution. However, we injected dasatinib solution twice-a-week in that study [10], whereas in the current study it was injected every other week. Retention of small molecules such as dasatinib in the vitreous cavity is known to be relatively short, and it is likely that injection performed every other week failed to maintain an effective concentration of the drug. The fact that Das-PLGA, which releases dasatinib up to 14 days [12], was effective in preventing development of traction retinal folds in our current model strongly implies that sustained presence of the drug is required. Maintaining effective concentration is an important factor for prevention of PVR with other drugs as well. Recent studies using methotrexate for the prevention of recurrent PVR demonstrate that, while a single injection at the time of corrective surgery failed to show statistically significant difference in retinal re-detachment rate compared to control [29], no re-detachment was observed in two separate pilot studies when drug presence was sustained in eyes by repeated intraocular injection [30, 31].

PLGA is well established as a biocompatible and biodegradable polymer [14]. Multiple studies have tested PLGA for intravitreal drug delivery, and most studies have demonstrated it to be well tolerated. Our data agrees with such studies with full-field ERG not detecting any significant difference in baseline retinal function after PLGA or Das-PLGA injection. In the current study, however, we did lose one eye to endophthalmitis following Das-PLGA injection. A recent study demonstrated the effect of shape of PLGA particles on the development of intraocular uveitis, with rod-shaped PLGA implants being innocuous compared to PLGA microparticles that induced endophthalmitis [15]. Furthermore, PLGA nanoparticles that were about half the diameter of particles used in this study were superior to PLGA microparticles in intraocular delivery of EGF receptor tyrosine kinase inhibitor AG1478 [32]. Therefore, further testing and optimization of the shape and size of PLGA for intravitreal drug delivery may be warranted.

In summary, intravitreal injection of Das-PLGA significantly inhibited development of traction retinal folds, which could eventually lead to severe PVR, in our pig model of posterior intraocular traumatic injury. Vitrectomy is used to repair posterior ocular trauma but its optimal timing remains a source of debate [33, 34]. Prevention or even significant delay of early fibrotic changes will allow more treatment options, as well as broadening the timing of surgical intervention. Our data suggests that adjunctive therapy with Das-PLGA shortly after posterior segment traumatic wounds will minimize early fibrotic changes and that sustained dasatinib release systems merit further investigation to prevent the loss of vision associated with PVR.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Ethics approval Animal work has been approved by the University of Louisville IACUC.

Consent to participate Not applicable

Consent to publication Not applicable

Code availability Not applicable

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References

1. Kuhn F, Morris R, Witherspoon CD, Mann L (2006) Epidemiology of blinding trauma in the United States Eye Injury Registry.
2. Fung K, Hu Y, Wang C, Shen L, Pang X, Jiang Y, Nie H, Wang Z, Ma Z (2013) Risk factors, anatomical, and visual outcomes of injured eyes with proliferative vitreoretinopathy: eye injury vitreoretinal study. Retina. https://doi.org/10.1097/IAE.0b013e3182525469
3. Cardillo JA, Stout JT, LaBree L, Azen SP, Omphroy L, Cui JZ, Kimura H, Hinton DR, Ryan SJ (1997) Post-traumatic proliferative vitreoretinopathy. The epidemiologic profile, onset, risk factors, and visual outcome. Ophthalmology 104:1166–1173
4. Cleary PE, Ryan SJ (1981) Vitreectomy in penetrating eye injury. Results of a controlled trial of vitreectomy in an experimental posterior penetrating eye injury in the rhesus monkey. Arch Ophthalmol 99:287–292. https://doi.org/10.1001/archopht.1981.03930010289014
5. Winthrop SR, Cleary PE, Minckler DS, Ryan SJ (1980) Penetrating eye injuries: a histopathological review. Br J Ophthalmol 64:809–817. https://doi.org/10.1136/bjo.64.11.809
6. Tolentino FL, Liu HS, Freeman HM, Natchiar G (1979) Vitreectomy in penetrating ocular trauma: an experimental study using rabbits. Ann Ophthalmol 11:1763–1771
7. Cardillo JA, Farah ME, Mitre J, Morales PH, Costa RA, Melo LA, Kuppermann B, Jorge R, Ashton P (2004) An intravitreal biodegradable sustained release naproxen and 5-fluorouracil system for the treatment of experimental post-traumatic proliferative vitreoretinopathy. Br J Ophthalmol 88:1201–1205. https://doi.org/10.1136/bjo.2003.039917
8. Khanna B, Guha R, Sivaprasad U, Nandi S, Basak SK, Konar A, Hazra S (2017) Pirfenidone inhibits post-traumatic proliferative vitreoretinopathy. Eye (London, England) 31:1317–1328. https://doi.org/10.1038/eye.2017.21
9. Nassar K, Luke J, Luke M, Abd El-Nabi E, Soliman M, Rohrbach M, Grisanti S (2011) The novel use of decorin in prevention of the development of proliferative vitreoretinopathy (PVR). Graefes Arch Clin Exp Ophthalmol 249:1649–1660. https://doi.org/10.1007/s00417-011-1730-9
10. Umazume K, Liu L, Scott PA, Fernandez de Castro JP, McDonald K, Kaplan HJ, Tamiya S (2013) Inhibition of PVR with a tyrosine kinase inhibitor, Dasatinib, in the swine. Invest Ophthalmol Vis Sci 54:1150. https://doi.org/10.1167/iovs.12-9768
11. Tsukahara R, Umazume K, McDonald K, Kaplan HJ, Tamiya S (2017) Focal adhesion kinase family is involved in matrix response. Invest Ophthalmol Vis Sci 58:4274–4285. https://doi.org/10.1167/iovs.16-21334
12. Gregor Z, Ryan SJ (1982) Combined posterior contusion and penetrating injury in the pig eye. II. Histological features. Br J Ophthalmol 66:799–804
13. Cleary PE, Ryan SJ (1979) Method of production and natural history of experimental posterior penetrating eye injury in the rhesus monkey. Am J Ophthalmol 88:212–220. https://doi.org/10.1016/0002-9394(79)90468-9
14. Umazume K, Ryan SJ (1979) Experimental posterior penetrating eye injury in the rabbit. I. Method of production and natural history. Br J Ophthalmol 63:306–311
15. Gregor Z, Ryan SJ (1983) Combined posterior contusion and penetrating injury in the pig eye. III. A controlled treatment trial of vitrectomy. Br J Ophthalmol 67:282–285
16. Umazume K, Barak Y, McDonald K, Liu L, Kaplan HJ, Tamiya S (2012) Proliferative vitreoretinopathy in the swine—a new model. Invest Ophthalmol Vis Sci 53:4941–4916. https://doi.org/10.1167/iovs.12-9768
17. Lombardo LJ, Lee FY, Chen P, Norris D, Barrish JC, Behnia K, Castaneda S, Cornelius LA, Das J, Doweyko AM, Fairchild C, Hunt JT, Inigo I, Johnston K, Kamath A, Kan D, Klei H, Marathe P, Pang S, Peterson R, Pitt S, Schieven GL, Schmidt RJ, Tokarski J, Wen ML, Wityak J, Borzilleri RM (2004) Discovery of N-(2-chloro-6-methyl- phenyl)-2-(4-(4-hydroxyethyl)-piperazin-1-yl)-2-methylpyrimidin-4-yl amino)theiazole-5-carboxamide (BMS-548525), a dual Src/Abl kinase inhibitor with potent antitumor activity in preclinical assays. J Med Chem 47:6658–6661. https://doi.org/10.1021/jm049486a
18. Hanschel O, Rix U, Superti-Furga G (2008) Target spectrum of the BCR-ABL inhibitors imatinib, nilotinib and dasatinib. Leuk Lymphoma 49:615–619. https://doi.org/10.1080/10428190801896103
19. Balasubramanian S, Pleasant DL, Kasigianes H, Quinones L, Zhang Y, Sundararaj KP, Roche S, O’Connor R, Bradshaw AD, Kuppuswamy D (2015) Dasatinib attenuates pressure overload-induced cardiac fibrosis in a murine transverse aortic constriction model. PLoS One 10:e0140273. https://doi.org/10.1371/journal.pone.0140273
20. Cruz FF, Horta LF, Maia Lde A, Lopes-Pacheco M, da Silva AB, Morales MM, Goncalves-de-Albuquerque CF, Takiya CM, de Castro-Faria-Neto HC, Rocco PR (2016) Dasatinib reduces lung inflammation and fibrosis in acute experimental silicosis. PLoS One 11:e0147005. https://doi.org/10.1371/journal.pone.0147005
21. Kanemaru R, Takahashi F, Kato M, Mitsuishi Y, Hayakawa D, Yagishita S, Goto T, Koinuma Y, Hayakawa D, Yagishita S, Ko R, Sato T, Harada N, Koda Y, Naruwyida F, Sasaki S, Niwa SI, Takahashi K (2018) Dasatinib suppresses TGF-beta-mediated epithelial-mesenchymal transition in alveolar epithelial cells and inhibits pulmonary fibrosis. Lung 196:531–541. https://doi.org/10.1007/s00438-018-0134-6
22. Yilmaz O, Ozayt F, Kayalar O (2015) Dasatinib attenuated bleomycin-induced pulmonary fibrosis in mice. Growth Factors 33:366–375. https://doi.org/10.1080/08977194.2015.1109511
23. Zeng XP, Wang LJ, Guo HL, He L, Bi YW, Xu ZL, Li ZS, Hu LH (2019) Dasatinib ameliorates chronic pancreatitis induced by caerulein via anti-fibrotic and anti-inflammatory mechanism. Pharmacol Res 147:104357. https://doi.org/10.1016/j.phrs.2019.104357
24. Tsukahara R, Umazume K, McDonald K, Kaplan HJ, Tamiya S (2017) Focal adhesion kinase family is involved in matrix
contraction by transdifferentiated Muller cells. Exp Eye Res 164: 90–94. https://doi.org/10.1016/j.exer.2017.08.010

29. Falavarjani KG, Hadavandkhani A, Parvaresh MM, Modarres M, Naseripour M, Alemzadeh SA (2020) Intra-silicone oil injection of methotrexate in retinal reattachment surgery for proliferative vitreoretinopathy. Ocul Immunol Inflamm 28:513–516. https://doi.org/10.1080/09273948.2019.1597894

30. Nourinia R, Borna F, Rahimi A, Jabbarpoor Bonyadi MH, Amizadeh Y, Daneshtalab A, Kheiri B, Ahmadieh H (2019) Repeated injection of methotrexate into silicone oil-filled eyes for grade C proliferative vitreoretinopathy: a pilot study. Ophthalmol J Int d’ophthalomol Int J Ophthalmol 242:113–117. https://doi.org/10.1159/000500271

31. Benner JD, Dao D, Butler JW, Hamill KJ (2019) Intravitreal methotrexate for the treatment of proliferative vitreoretinopathy. BMJ Open Ophthalmol 4:e000293. https://doi.org/10.1136/bmjophth-2019-000293

32. Robinson R, Viviano SR, Criscione JM, Williams CA, Jun L, Tsai JC, Lavik EB (2011) Nanospheres delivering the EGFR TKI AG1478 promote optic nerve regeneration: the role of size for intraocular drug delivery. ACS Nano 5:4392–4400. https://doi.org/10.1021/nn103146p

33. Aylward GW (2008) Vitreous management in penetrating trauma: primary repair and secondary intervention. Eye (London, England) 22:1366–1369. https://doi.org/10.1038/eye.2008.74

34. Han L, Jia J, Fan Y, Yang L, Yue Z, Zhang W, Liu F, Kang H, Huo T, Han S, Shen H, Tian G, Su X (2019) The vitrectomy timing individualization system for ocular trauma (VTISOT). Sci Rep 9: 12612. https://doi.org/10.1038/s41598-019-48862-2

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