Review

Intravitreal Injectable Hydrogels for Sustained Drug Delivery in Glaucoma Treatment and Therapy

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Abstract: Glaucoma is extensively treated with topical eye drops containing drugs. However, the retention time of the loaded drugs and the in vivo bioavailability of the drugs are highly influenced before reaching the targeted area sufficiently, due to physiological and anatomical barriers of the eye, such as rapid nasolacrimal drainage. Poor intraocular penetration and frequent administration may also cause ocular cytotoxicity. A novel approach to overcome these drawbacks is the use of injectable hydrogels administered intravitreally for sustained drug delivery to the target site. These injectable hydrogels are used as nanocarriers to intimately interact with specific diseased ocular tissues to increase the therapeutic efficacy and drug bioavailability of the anti-glaucomic drugs. The human eye is very delicate, and is sensitive to contact with any foreign body material. However, natural biopolymers are non-reactive, biocompatible, biodegradable, and lack immunogenic and inflammatory responses to the host whenever they are incorporated in drug delivery systems. These favorable biomaterial properties have made them widely applicable in biomedical applications, with minimal adversity. This review highlights the importance of using natural biopolymer-based intravitreal hydrogel drug delivery systems for glaucoma treatment over conventional methods.

Keywords: glaucoma; natural biopolymers; intravitreal injectable hydrogel; drug delivery systems

1. Introduction

Reports of the World Health Organization (WHO) indicate that nearly 80 million people have glaucoma globally, and around half of the people with the disease are unaware that they have it. This number is expected to rise to 112 million individuals by 2040 [1]. Glaucoma has become the second most example cause of irreparable visual impairment around the world. It is a multifactorial, imperceptible, and gradual neurodegenerative disease that affects the optic nerve [2].

In human physiology, the eye is a delicate and complex organ that is specialized for detecting and converting light stimuli into meaningful information that is structured into different sections—namely, the anterior and posterior sections [3]. The anterior chamber located in between the cornea and the iris is filled with aqueous humor made by the ciliary body [4]. Fluid from the anterior chamber flows out through the pupil and then reaches the eye’s drainage system, as well as the trabecular meshwork and a network of canals [5,6]. The vitreous chamber situated between the lens and the back of the eye contains a thick, gel-like fluid called vitreous humor or vitreous gel [7,8]. Proper drainage of this fluid helps the eye to keep internal pressure at a normal level, which is an active and continuous process that is needed for the health of the eye.

In the human eye, the normal level of intraocular pressure (IOP)—as indicated in Figure 1—can be regulated by the balance between how much fluid is made and how much...
drains out of the eye in a given time [9]. In most types of glaucoma, the eye’s drainage system becomes clogged, so the intraocular fluid cannot drain [10].

![Figure 1. Physiology of a human eye with normal IOP.](image)

The physiological optic cycle establishes the drainage flow through and out of the eye, and if any partial or total drainage obstruction occurs, elevation of the IOP may cause adverse effects on the retinal ganglion cells (RGCs) and RGC axons, causing glaucoma. The course of glaucomic progression is chronic and perdurable for a long time—usually asymptomatic, but gradually impairing the peripheral visual field before optimal damage [11].

The main causes of increased and abnormal elevation in IOP driving primary glaucoma remain undiscovered [12]; however, the causes of increased IOP leading to secondary glaucoma have been identified. Abnormally high IOP may be caused by advanced cataracts, inflammation, high or elevated blood pressure, optic tumors, diabetes, myopia, or hyperthyroidism [13]. Genetic and ethnic background, age, lifestyle, and narrowing retinal nerve fibers are risk factors that have been associated with glaucoma [14]. The disease has also been suspected to arise as a side effect of the prolonged use of corticosteroids [15].

Currently, there are various glaucoma treatments on the market that are designed to lower the IOP by lessening the production of aqueous humor or expanding non-trabecular fluid humor drainage with surgical- (i.e., implants and therapeutic surgeries) and pharmacological-based methodologies. Pharmaceutical therapeutics for the management of glaucoma through topical eye drops, ointments, and oral medications reduce elevated IOP; however, there are limitations, including low patient compliance and insufficient bioavailability [16]. To elevate bioavailability and improve patient compliance, advanced drug delivery mechanisms such as liposomes, microneedles, niosomes, dendrimers, ocular inserts, nanoparticles, and injectable hydrogels should be used [17].

To overcome the limitations of pharmaceutical therapeutics, several methods have been explored, including the use of in situ hydrogels, which potentially serve as delivery vehicles for nutrients, oxygen, and drugs to the targeted area [18].

Hydrogels play a significant role in upgrading the remedial adequacy of anti-glaucomic drugs, and are relevant in glaucoma treatment because of their reliable drug delivery applicability [19]. Polymers crosslinked with hydrophilic drugs can retain more than 90% water within the mesh of their porous network structure, thus aiding in the encapsulation of hydrophilic drugs [20]. Therefore, localized delivery of drug-loaded hydrogels can be achieved relatively more easily and less invasively than by implantation, and can reach the tissues that are difficult for conventional delivery methods to reach [21].

### 2. Glaucoma

Glaucoma is classified primarily according to the severity and different causes of the ailment. The most predominant types are primary open-angle glaucoma (POAG) and
primary acute angle-closure glaucoma (PACG) [22]. Gradual blockage of the eyes drainage channel—resulting in increased IOP, as shown in Figure 2—is the cause of POAG [23].

![Figure 2. Open-angle glaucoma (chronic).](image)

Primarily, PACG is caused by elevation of the intraocular aqueous humor outflow at the closure angle, caused by mechanical occlusion of the iris tissue [24,25], or blockage of the Schlemm canal, as shown in Figure 3. Uveitis glaucoma, pigmentary glaucoma, and normal-tension glaucoma (NTG) are classified as secondary types of glaucoma [26].

![Figure 3. Blockage of Schlemm canal drainage.](image)

3. Ocular Barriers

The eye’s anatomy and physiological structure comprise ocular barriers that are profound for defending its inner components from unfamiliar substances [27]. These ocular barriers include the blood–ocular barrier, tear film, conjunctiva, cornea, blood–aqueous barrier, and blood–retina barrier, which control the uptake of liquids. The anatomical barriers comprise conjunctiva, sclera, and the cornea, retina–blood–aqueous barrier, and blood–retina barrier. Secondly, all ocular mechanisms are protected by active physiological clearing systems. The systems are nasolacrimal drainage and pre-corneal tear secretion for the elimination of irritants, the blinking reflex, conjunctival blood flow, efflux transport, and choroid, which shield the eye from the effects of destructive drugs [28].

The principal objective of ophthalmological therapy is to bypass the defensive hindrances of the eye without affecting the surrounding tissue. These barriers forestall the transition, withholding, and bioavailability of some ophthalmic medications by restricting ocular drug permeability to the foremost segments of the eye [29].
4. Current Therapies for Glaucoma

Ordinarily, the noninvasive methodologies incorporate topical eye drops, ointments, and oral drugs, while surgical nanotechnology has enabled glaucoma drainage through inserts, laser treatment procedures, and trabeculectomy. A schematic summary of current drug delivery methods and formulations is presented in Figure 4. However, these methodologies have drawbacks [30].

![Figure 4. Schematic summary of drug delivery systems for glaucoma treatments.](image)

Despite their efficiency in lowering the IOP, these therapeutic approaches have some important adverse effects. For example, drawbacks associated with oral medication in the treatment of glaucoma include conjunctival hyperemia in certain formulations. One of the most common treatments among oral medications is topical eye drops. However, this treatment is subject to low patient compliance and low bioavailability [31]. Alternatively, laser and surgical methodologies such as inserts and trabeculectomy have been found to be effective. However, they also carry adverse limitations, such as swelling, soreness, dryness of the cornea, and post-surgical complications [32].

In light of this, searching for novel targets to treat IOP and play a neuroprotective role could be taken as an advancement in the treatment of glaucoma by decreasing the side effects of the currently available drugs. To counteract these hindrances, targeted drug delivery systems that simplify ophthalmological treatments of glaucoma for an extended duration after the administration in the anterior and posterior parts of the eye have been designed to be sustainable [33]. To elevate bioavailability or alleviate chronic visual impairments, advanced drug delivery mechanisms should be used. Liposomes, microneedles, niosomes, dendrimers, ocular inserts, nanoparticles, and in situ hydrogels assume a significant role in upgrading the remedial adequacy of the anti-glaucomic drugs [34].

5. Constraints of Current Glaucoma Drug Delivery Treatment

The drug delivery of anti-glaucoma medications and therapies is challenging because of the presence of ocular barriers, which result in low bioavailability of the active ingredient within the drug [9]. Essential challenges in the administration of ocular medications through conventional strategies incorporate the lack of patient training for the technique in terms of medication, consistency, adherence, and diligence. Each treatment approach has its limitations, and there are severe side effects of some applications.

5.1. Eye Drops and Eye Ointments

Eye drops are a fundamental type of topical administration due to their ease of application, favorable cost, and good patient consistency [35]. The typical retention time of eye drops administered topically in the pre-corneal tear film is about 1 min. This retention time is the only time presented for drug permeation through the cornea to reach the
aqueous humor. Due to obstruction of the cornea and pre-corneal components, under 5% of completely directed medications permeate to the aqueous humor. The corneal epithelium, which contains different desmosomes and tight intersections, forestalls the permeation of particles bigger than 500 Da, barring them from infiltrating the cornea [36].

Subsequently, 80% of the conveyed drug is unable to enter the cornea, and might be absorbed into the veins of the conjunctiva. Just under 10% of the given medication is uptaken into the ocular system, and roughly 1–7% of that arrives at the target site—particularly in the aqueous humor [37]. However, the most widely prescribed anti-glaucoma medications—the PG analogs—do not suffer from these same limitations, and can be effectively administered once daily, with good effects in many patients. Some drugs can infiltrate the cornea, and are immediately separated through the trabecular meshwork. In the trabecular meshwork, most ocular drugs possess a half-life slightly less than 2 h, which is a hindrance for drug molecules in reaching the targeted tissue. Regularly utilized eye drops and ointments have little corneal penetrability, and are thus restricted to treatment in the external fragment of the eye. Subsequently, to achieve the ideal dosage using this approach, eye drops should be controlled with high-recurrence dosing regimens. Consequently, using eye ointments to attain the required dosage to the posterior chambers may cause harm to the ocular cells. Patient movement while administering the eye drops and regular medication bring about poor patient compliance, and making sure to take a daily dose of the optical drug might be a challenge for patients as well [38].

5.2. Trabeculectomy

Trabeculectomy is frequently associated with visual hypotony, which is a post-surgical complication due to an overabundance in the filtration of fluid humor after a medical procedure. Nonetheless, surgical procedures are restricted to treatments—for example, corneal transplant, glaucoma treatment, or removal of the vitreous humor. Surgical drainage inserts are utilized in glaucoma treatment when IOP-decreasing medications cease to work. A common postoperative complication after implantation of these devices is the development of fibrosis around the implants [39,40].

5.3. Laser Treatment

For patients who are unable to endure the administration of other forms of medication, or for whom the therapeutic drug alone has not been satisfactory, laser treatment is an alternative. Laser treatment is becoming more common—particularly in Europe—for the treatment of glaucoma.

Subliminal trans-scleral cyclophotocoagulation (SL-TSCPC) is one of the alternative therapies to decrease IOP safely and efficiently. However, there are few studies regarding SL-TSCPC using a Supra 810 laser machine, and limited data regarding its effectiveness in moderate-severity glaucoma that still has good preservation of vision. SL-TSCPC is a safe and alternative method of lowering IOP in moderate-to-advanced glaucoma over 6 months of follow-up. As it has a good safety profile and repeatability, it is a good treatment option for patients with uncontrolled glaucoma. The parallel effect while using laser treatment is usually temporary, and may cause swelling, soreness, dryness of the cornea, and/or risk of corneal scratching by the laser [41].

5.4. Oral Medication

As it is rare for oral medications to be administered to glaucoma patients, when applied, the potential side effects include metallic taste, depletion of potassium, and development of kidney stones [42].

6. Current Pharmaceutical Interventions for the Treatment of Glaucoma

A number of medications are currently in use for the treatment of glaucoma. Typically, medications are intended to decrease elevated IOP and prevent loss of optic nerve fibers. Generally, drugs used in the treatment of glaucoma are classified by their active ingredient.
These include prostaglandin (PG) analogs, β-blockers, α-adrenergic agonists, carbonic anhydrase inhibitors, and rho-kinase inhibitors. Combination drugs are also available for patients who require more than one type of medication. With various technological advancements, some drugs of various classes—including carbonic anhydrase inhibitors, PG analogs, β-blockers, miotics, α-adrenergic agonists, and hyperosmotics—have been developed. These drugs are responsible for treating glaucomatous complexes either by increasing aqueous humor drainage from the eye or by reducing aqueous humor production [43].

6.1. Beta-Adrenergic Blockers

Timolol maleate (TM) is the primary line of medication in the treatment of glaucoma [44], belonging to the class of β-adrenergic blockers. Even after the appearance of the most recent medications, such as PG analogs and α-2 agonists, TM remains the best option because of its cost-effectiveness. Long-lasting treatment with skin drops is typically needed in the treatment of glaucoma. Consequently, a decrease in dosing recurrence can improve tolerance consistency and treatment. Having low blood pressure, fatigue, and a low pulse rate are the side effects of the medication. β-blockers can also be a reason for shortness of breath in people who have a history of asthma or other respiratory disorders, and can alter cardiac activity by decreasing the amount of blood the heart pumps out, which may reduce the pulse rate and/or slow down the heart’s response rate during rare side effects of exercise, including reduced libido and depression [45,46].

6.2. PG Analogs

PG analogs are another course of visual hypotensive medications produced for the treatment of POAG. Latanoprost and unoprostone are medications that lower the IOP specifically by increasing the uveoscleral drainage. The standard dosage of PG that reduces IOP by 30% in glaucomatous patients is 50 µg/mL, applied topically once per day. Additional therapeutic impact is achieved when PGs are used with other glaucoma treatments. Potential side effects include eye color change, eyelash growth, droopy eyelids, darkening of eyelid skin, sunken eyes, stinging, eye redness, and itching [47].

6.3. Alpha-Adrenergic Agonists

Alpha-adrenergic agonists are usually applied after ocular laser therapy to decrease the aqueous humor secretion and to control the adverse increases in IOP and episcleral venous pressure. Unfortunately, they can induce ocular irritation and dry eyes, along with systemic side effects involving the central nervous system, and are therefore usually not recommended for long-term therapy [43].

6.4. Carbonic Anhydrase Inhibitors

Carbonic anhydrase inhibitors are sulfonamide derivatives that decrease ocular pressure by lessening the production of intraocular fluid, thus reducing the formation of aqueous humor and inhibiting the activity of carbonic anhydrase in the ciliary process of the eye, consequently decreasing the IOP. They are available and administered in the form of eye drops and as pills. Systemic use of carbonic anhydrase inhibitors reduces the IOP by approximately 40%. Utilizing these medications causes a quick impact on the therapeutic treatment of acute angle-closure glaucoma. Loss of strength of the hands and feet, along with tingling, upset stomach, mental fuzziness, memory problems, depression, kidney stones, and frequent urination, are among the side effects of the pill form of these medications. Side effects of the eye drops include stinging, burning, and other forms of eye discomfort [48].

6.5. Miotic Agents

Treatments using miotic agents primarily decrease pressure in the eye by increasing the drainage of intraocular liquid through the trabecular meshwork. Miotics work by con-
striction of the ciliary muscle, fixing the trabecular meshwork and permitting an expanded surge of fluid through the customary pathways. Miosis results from the activity of these medications on the pupillary sphincter. Patients who use these medications complain of dim vision, especially at night or in darkened areas such as movie theaters. This is due to constriction of the pupil. Miotics increase the drainage of intraocular fluid by making the pupil smaller, thereby increasing the flow of intraocular fluid from the eye [49].

6.6. Hyperosmotic Agents

Hyperosmotic agents essentially lower the IOP by causing an osmotic inclination between ocular fluid and blood. Systemic adverse effects include nausea, vomiting, headache, increased thirst, chills, fever, confusion or disorientation, electrolyte imbalances, and urinary retention [50].

7. Natural Polymer-Based Hydrogels as Drug Delivery Vehicles for Glaucoma Therapy

Biopolymers have been extensively investigated in a number of medical fields, including tissue engineering and drug delivery. This is largely due to the fact that they are biodegradable within the body, and do not induce an inflammatory reaction [51]. A summary of some polymers used in anti-glaucoma drug delivery systems can be seen in Section 7.9. Polynucleotides such as nucleic acids (DNA and RNA), proteins such as polypeptides, and polyesters derived from both plants and animals are also used [52].

When compared to synthetic polymers, naturally occurring biopolymers and their derivatives have acquired preference, and have a comprehensive range of applications in pharmaceutical as well as biomedical research. Natural biopolymers are preferred for medical applications due to their biodegradability, biostability, biocompatibility, and non-toxicity [53]. Additionally, natural polymers have the advantage of being readily available, economically friendly, and ecologically friendly. Hydrogels derived from natural polymers exhibit high potential as drug delivery systems for biomaterials to treat ocular impairments [54,55].

The current market is brimming with numerous formulations and applications of biopolymers that are intended to treat glaucoma. Every one of the present modes and applications of drug delivery utilizes a particular biopolymer. The advantages and disadvantages of various natural biopolymers are tabulated in Section 7.9.

7.1. Silk Fibroin

*Bombyx mori* silk is a natural biopolymer obtained from arthropods and lepidopteran insects, particularly silkworms and some spider groups, that produce silk fibers at large. Due to their remarkable mechanophysical and biological properties, silk fibers have attracted the interest of researchers [56], for biomedical and pharmaceutical applications. Silk fibroin is an essential biopolymer used in biomedical applications due to its adaptable properties, with a natural physiology that makes it preferable in the study of tissue reconstruction in age-related ocular disease [57]. Silk fibroin is a fibrous protein that exhibits favorable biocompatibility, bioresorbability, low immunogenicity, and hydrophilicity, promoting its increasing consideration in hydrogel design. It is also rich in $\beta$-sheet structures, owing to hydrophobic domains that influence its biodegradability rate [58], as well as its cytological compatibility [59]. Silk fibroin proteins have been used for ocular therapies such as wound healing [60], ocular drug delivery [61], and ocular prostheses [62].

7.2. Chitosan

Linear-structured chitosan is a natural biopolymer composed of an acetylated unit of N-acetyl-D-glucosamine and $\beta$-(1→4)-linked D-glucosamine, a deacetylated unit. It is prepared by treating chitin shrimp shells and various crustacean shells with sodium hydroxide [63].

Due to poly-oxy salt formation, chitosan exhibits basic properties different from those of other polysaccharides [64]. As with other polymers, chitosan can also form hydrogels,
films, and particles that can be used for biomedical applications in terms of drug delivery units, tissue engineering, cell culturing, and platforms for cancer diagnosis. Its low toxicity, high biocompatibility, and easy degradability in a natural environment which makes it suitable as natural extracellular matrices [65].

According to surveys carried out by several researchers, the major constraint when working with injectable hydrogel preparations is regulating the time of gelation [66]. However, a chitosan-based formulation of injectable hydrogel was developed that regulates the time of gelation [67].

To be used as a biomaterial, chitosan has important properties that mimic the extracellular matrices of cells, tissues, and organs. Chitosan is prepared and used either in a dried form or in the form of gels, depending on the temperature used and the amount of water present in the structure, which impart properties of flexibility [68].

According to the study conducted by Franca, J.R. et al. [69], chitosan can be widely applied in the treatment of glaucoma-induced intraocular pressure, acting as a basis for controlled drug delivery in the eye. This is because chitosan is polycationic by its very nature, allowing interaction with the polyanionic surface through hydrogen bonding of the ocular mucosa. Chitosan has several biological properties that make it an attractive material for use in ocular formulations [70]. Chitosan has inherent antimicrobial and mucoadhesive properties [71], as well as low toxicity, biodegradability, biocompatibility, and hemocompatibility [72]. Chitosan can disrupt epithelial tight junctions, thus acting as a permeability enhancer [71].

7.3. Alginic Acid

Brown algae are the main source of the naturally derived polysaccharide alginic acid (Alg), with the molecular formula \((\text{C}_6\text{H}_{11}\text{NO}_6)_n\). The molecular structure and composition of alginic acid consist of L-guluronic acid and D-mannuronic acid structures connected with alpha-1,4 bonds [73]. As a result of the carboxyl group attached to the C-5 carbon as a chain, it exhibits an acidic nature, and with properties such as high hydrophilicity, the capacity for gelation, and pH-dependent viscoelasticity. Furthermore, biocompatibility and biodegradability are some of the physiological properties that make it suitable for use as films and gels developed for medical and food applications [74]. Alginic acid has a biodegradable and biocompatible nature that is favorable for researchers; therefore, its use has been encouraged in ocular treatments [75]. Ocular delivery therapeutics are a current trend in ophthalmology, and alginates have been employed to play an imperative role because of their biocompatibility and immunogenicity [76].

7.4. Pullulan

Pullulan is a non-ionic polysaccharide extracted from the fermentation of black yeast (Aureobasidium pullulan), and is used broadly in biomedical applications because of its less immunogenic reaction, along with its non-toxic, non-mutagenic, and non-carcinogenic nature [77]. It is utilized in the targeted delivery of drug mechanisms, tissue engineering therapy, and wound-healing activities. Pullulan responds to external stimuli so that it can be used to design hydrogels, which can be used to deliver drugs, nutrients, and (any) other molecules to a targeted area of the host [78]. The biological properties of pullulan include high water retention, biocompatibility, cytocompatibility, protective activity against microbes and biodegradation, and tissue-regenerative characteristics [79].

7.5. Hyaluronic Acid

A biopolymer regularly found and extracted from the human body, applications of hyaluronic acid as injectable hydrogels have been researched for ocular drug delivery systems, since they can be designed as both stimulus-responsive and static [80]. Anionic hyaluronic acid is incapable of gelation without additive molecules. Hence, hydrogels produced using hyaluronic acid depend on chemical modifications. Egbo et al. formulated two hyaluronic acid gel systems embedded with infliximab for the treatment of blinding
infections influencing the elderly population [81]. Hyaluronic acid has been applied in ocular therapeutics because of its favorable biological characteristics, such as biocompatibility, biodegradability, and non-immunogenicity [82]. Due to hyaluronic acid’s biological safety, it has various ophthalmology-related applications, such as treatment for dry eyes, intravitreal drug delivery, and use in contact lenses [82].

The main objective in the development of ophthalmic drug treatment is to extend the therapeutic extent of medications, particularly proteins and antibodies [83].

7.6. Dextran

Dextran methacrylate and cyclodextrin–dextran are a few examples of dextran hydrogels used in ocular drug delivery [84]. Properties such as stiffness, mechanical strength, and solidness can be adjusted by regulating the monomer in the gel, subsequently improving their significance in drug delivery. Yao et al. [85], designed a drug delivery system for effective in vivo drug release of bevacizumab from a hyaluronic acid/dextran-situated in situ hydrogel for 6 months after intravitreal infusion in hare eyes. The in vivo drug release efficiency results indicated that bevacizumab was delivered at a therapeutically relevant concentration by means of a controlled release mechanism within the vitreous humor [86]. Dextran has been found to exhibit great biocompatibility and low cytotoxicity. Additionally, it has hydrophilic domains, which promote its biodegradability in water and other organic solvents. This biological feature enables its applicability in blended forms with bioactive agents of hydrophobic polymers [87].

7.7. Methylcellulose

Derived from cellulose, hydroxypropyl methylcellulose (HMPC) is widely used in the pharmaceutical industry because of its solvency in water, rheological properties, and transparency [88]. A group of researchers designed a trans-scleral antisense oligonucleotide-loaded gel for the delivery of drug-loaded macromolecules using methylcellulose and ι-carrageenan dispersions [89]. Periocular injection of the gel resulted in impressive choroid and sclera bioavailability in comparison to the injection of an oligonucleotide solution alone. Methylcellulose has been incorporated into ocular inserts of three types: soluble, insoluble, and bio-erodible, [90]. Methylcellulose has low reactivity with cells. Additionally, interest has been shown in mixing it with biologically active materials such as cytokines and/or the extracellular matrix to control the organization or functions of the cells [91].

7.8. Gelatin

Gelatin is a collagen-derived biopolymer normally found in scleral and corneal stroma, and its structural networks make it an attractive natural complex for research applications. El-Feky et al. [92] developed an oxidized sucrose-crosslinked gelatin–chitosan hydrogel with the end goal of TM drug conveyance for the treatment and control of ocular hypertension [93]. In vivo and in vitro discoveries indicated that the formulated system maintained favorable release efficacy of the active ingredient, in contrast to the regular eye drops [94]. Gelatin has favorable biological characteristics such as low antigenicity, biocompatibility, and biodegradability, and promotes cell proliferation; therefore, it is widely researched in ophthalmologic therapeutics [95].

7.9. Collagen

Collagen is biocompatible, biodegradable, and non-toxic for living organisms [96]. Type 1 collagen is an essential biopolymer that has been utilized in hydrogels for tissue engineering applications [97]. Wong et al. [41] designed an injectable composite comprising collagen and alginate for retinal treatment through a drug delivery system loaded with an ocular drug. A summary of polymers used in anti-glaucoma drug delivery systems discussed in Table 1. Table 2 presents some natural biopolymers used in ophthalmic injectable hydrogels. Intravitreally infused gels exhibited adequacy in rodents with deteriorating
retinae and photoreceptor apoptosis. Twofold portion infusions into a similar eye yielded greater results without sacrificing the gel’s feasibility [98].

Table 1. Summary of some polymers used in anti-glaucoma drug delivery systems.

| Polymers                              | Delivery System | Drug Used                  | Feature                                                                 | Reference |
|---------------------------------------|-----------------|----------------------------|-------------------------------------------------------------------------|-----------|
| Silk fibroin                          | Nanoparticles   | TM                         | TM caused a sustained and prolonged reduction in IOP without adverse effects on the physiology of the eye compared to conventional free drug use. | [99]      |
| Hydroxyethyl chitosan                 | Hydrogel        | Heparin                    | The heparin-loaded hydroxyethyl chitosan hydrogel was able to sustain and improve the reduction in the IOP after GFS for protracted periods of time. Clear inflammatory responses and results were not seen in the eye during the trial’s timeframe. | [100]     |
| Gelatin-g-poly(N-isopropylacrylamide) | Hydrogel        | Pilocarpine                | Pilocarpine-loaded gelatin hydrogels were designed by grafting with carboxylic end-capped poly(N-isopropylacrylamide) for anti-glaucoma treatment by intracameral administration. | [101]     |
| Poly(lactic-co-glycolic acid) (PLGA)  | Nanoparticles   | Dexamethasone and melatonin | A dual-loaded melatonin and dexamethasone poly(lactic-co-glycolic acid) nanoparticle system was designed as an anti-glaucoma treatment option. The in vitro release of the loaded drug from the nanoparticles revealed a supported delivery profile for the two medications, with no signs of burst discharge. Development of a novel fluorescein delivery system that is applied topically in dry nanofibrous form and gels in situ immediately after administration guaranteed a solid match to the eye structure by the designed nanofibers, which were molded into conforming geometries. Prolongation of the ocular drugs’ residence time was achieved. | [102]     |
| Gellan gum/pullulan                   | Nanofibers, in situ gel | Fluorescein sodium | Dorzolamide-loaded ocular inserts were effective in glaucoma treatment. The ophthalmologic drug embedded in the polymeric matrix displayed a 3-h drug release efficiency, and released 75% of the loaded drug. This study proposed that loading and delivering TM onto alginic–chitosan nanoparticles may be a suitable drug delivery approach for controlled delivery of TM through the cornea. | [103]     |
| Chitosan/hydroxyethyl cellulose       | Ocular inserts   | Dorzolamide                | Dorzolamide-loaded ocular inserts were effective in glaucoma treatment. The ophthalmologic drug embedded in the polymeric matrix displayed a 3-h drug release efficiency, and released 75% of the loaded drug. This study proposed that loading and delivering TM onto alginic–chitosan nanoparticles may be a suitable drug delivery approach for controlled delivery of TM through the cornea. | [104]     |
| Alginate–chitosan                     | Nanoparticles/ nanogels | TM                        | Dorzolamide-loaded ocular inserts were effective in glaucoma treatment. The ophthalmologic drug embedded in the polymeric matrix displayed a 3-h drug release efficiency, and released 75% of the loaded drug. This study proposed that loading and delivering TM onto alginic–chitosan nanoparticles may be a suitable drug delivery approach for controlled delivery of TM through the cornea. | [99]      |

Table 2. Natural biopolymers used in ophthalmic injectable hydrogels.

| Natural Biopolymer | Gelation | Strengths                                      | Drawbacks                                                   | Reference |
|--------------------|----------|-----------------------------------------------|-------------------------------------------------------------|-----------|
| Silk fibroin       | Ionic crosslinking, hydrophobic interactions | Easily modified                                              | Low mechanical strength                                     | [53]      |
| Chitosan           | Chemical crosslinking, pH gelation          | Simple to adjust                                              | Low dissolvability at neutral pH                            | [54]      |
| Alginate           | Chemical gelation, ionic crosslinking       | Favorable mechanical properties, rapid gelation              | Poor cytoadhesion                                           | [55]      |
| Gelatin            | Chemical crosslinking                        | Hydrophilic, various responses available                      | Susceptible to degradation, poor mechanical properties      | [94]      |
| Pullulan           | Chemical crosslinking                        | Easily dissolvable                                           | Undesirable swelling properties and mechanical properties   | [77]      |
| Methylcellulose    | Hydrophobic, chemical, physical             | Easy modification of physiochemical properties               | Uncontrollable degradation, poor cell adhesion, poor mechanical properties | [74,89]   |
| Dextran            | Chemical crosslinking, physical crosslinking | Simple crosslinking, large capacity, hydrophilic, controlled drug release | Prone to causing in vivo side effects                        | [71,90]   |
8. Design of Hybrid Hydrogels for Injectable Drug Delivery in the Treatment of Glaucoma

The development of hybrid hydrogels should be conducted in a sterile environment that is free from the creation of an overabundance of gas, protons, heat, or substances dangerous to living beings. The gel should be shaped under physiological states of temperature and ionic strength, light, and chemical gelation, in a controllable manner [105,106].

A hydrogel ought to have the capacity to conform to various applications, and should have the option to be administered via a slight needle (i.e., 30-gauge or thinner) in the confined space of the eye. Other fundamental variables to be considered are the mode of crosslinking, the solvents used in biopolymer dissolution, the solvent and chemical molecular weight and concentration, and the crosslinking time period [107,108].

8.1. Physicochemical, Pharmacokinetic, and Pharmacodynamic Properties of Ophthalmic Hydrogels

To design this intravitreal injectable hydrogel system, there should be strict parameters adhered to, such as viscoelasticity, viscosity, drug release efficiency, sustainability, etc. [109,110].

8.1.1. Drug Release Efficiency

Hydrogels developed for targeted drug delivery should have the ability to encapsulate a highly concentrated drug with a sustained release profile from the crosslinked hydrogel, so that an initial burst release is inevitable [111]. A high local concentration of the active pharmaceutical ingredient is retained over a significant stretch of time by means of a suitable release mechanism controlled by swelling, diffusion, or chemical/environmental stimuli [112].

Covalently crosslinked hydrogels have been utilized in the development of ocular drug delivery systems. These hydrogels remain in situ if applied topically to the lacrimal canal; however, if administered intravitreously, they tend to display swelling mechanisms. The drug release efficiency of embedded drugs can be modulated to some degree by variance in biopolymer concentration and molecular weight, adjustment of crosslinking density, and alteration of the degradation rate. Modulating these parameters may also alter other parameters of the drug–complex system, such as biocompatibility, mechanical properties, and the stability of the active ingredient [113].

8.1.2. Biocompatibility

Biocompatibility testing provides initial screening of whether or not the components of the desired biomaterial may cause adverse effects when interacting with the human body. These biocompatibility tests may include sensitization assays, carcinogenesis, hemocompatibility, genotoxicity, and systemic toxicity [111]. To design an injectable hydrogel, biocompatibility with cells, bodily fluids, and tissues should be considered in light of the fact that the hydrogel should maintain cell differentiation without causing cytotoxicity or adverse inflammatory responses in the host organism. Subunits of most biopolymers derived from natural sources are similar to organic extracellular matrix (ECM), making them more biocompatible than manmade polymers [114].
8.1.3. Biodegradability

Naturally derived biopolymers with the ability to degrade to naturally occurring byproducts without causing adverse inflammatory and immune responses are preferred for use [115]. The hydrogel ought to be biodegraded, bio-eroded [116], and expelled from the host at a rate that is relative to the pace of the tissue/organ development, so as to make space for cell multiplication [117].

8.1.4. Porosity

Exceptionally interconnected and profoundly coordinated permeable networks that exhibit micro- or nanoscale pores are preferred in hydrogel design and development [118]. The partial permeability facilitates active ingredient release of the therapeutic drug from the hydrogel, and supports cell nutrient movement for cell growth [119].

8.1.5. Viscosity

Low hydrogel viscosity is a crucial factor when designing these biomaterials, because the hydrogel should permit a homogeneous distribution of the active ingredient prior to complex gelation [120].

8.1.6. Mechanical Strength

A hydrogel ought to offer mechanical support and guide cell differentiation [121,122]. Mechanical strength is significant because the biomaterial must be able to withstand its biochemical structure and shape to overcome unavoidable natural forces that come with the eye’s physiology [123].

8.1.7. Swelling Properties

A favorable property of hydrogels is their capacity to grow in contact with a thermodynamically viable solvent [124,125]. At the point when a hydrogel in its underlying state is in contact with dissolvable particles, the latter assaults the hydrogel’s surface and infiltrates into the polymeric organization [126]. Solvent molecules penetrate into the polymeric network due to charge repulsion between polymer chains, causing an increase in the polymer volume due to liquid uptake [127].

8.1.8. Rheology

Rheology instruments can be utilized to gauge antecedent arrangement attributes, e.g., yield pressure, which have direct importance for clinicians’ utilization of the materials [110,128]. Practically speaking, the three most applicable rheological boundaries are the simplicity of infusion (shear reaction), time for position (recovery time), and maintenance of the hydrogel’s forerunner arrangement at the deformity site (yield pressure) [129].

8.1.9. Opacity and Transparency

Hydrogels that bio-mimic the high opacity and transparency of the natural ocular humor remain materials of preference [130,131].

9. Intravitreal Administration of Injectable Drug-Loaded Hydrogels to The Eye

Ophthalmic applications of intravitreal injectable drug-loaded hydrogels for glaucomic treatment to the posterior section of the eye have been shown to overcome ocular barriers and effectively treat glaucoma [132,133]. This novel drug delivery system has properties of good adherence and viscosity, and achieves the objective of retention and therapeutic treatment of ocular diseases in the most remote areas of the eye.

To counteract ocular barriers and defense mechanisms, scientists have developed intravitreal injectable hydrogel-based drug delivery mechanisms that enable retention on the surface of the eye and in the posterior segments of the eye for an extended duration after their administration, where conventional therapies have not successfully and sustainably achieved the same targets [134]. Administration of intravitreal injectable hydrogels results
in an acceptable retinal bioavailability, since the drug is directly injected into the targeted area [135]. An advantage of administering intravitreal injectable hydrogels is that they are biodegradable, so they are exceptionally alluring as intravitreal delivery systems because they are degraded by the body over a favorable timeframe that relies upon the particular polymeric framework utilized [136].

The installation of this type of delivery system is just one methodology. For example, the drug-loaded polymeric complex will be required once as an injection to the target site, whereafter the polymer will undergo optimal biodegradation, bio-erosion, and elimination from the body, without creating any possible incendiary side effects. Intravitreal injectable hydrogels alleviate the problem faced while using solid implants, because injectable hydrogels can hold water. In addition, their easy dissolution enables them to encapsulate drugs to be released in the local area. The administration of intravitreal hydrogels as drug delivery systems increases the bioavailability of the active drug at the required target site by overcoming ocular barriers [137].

Due to their biocompatibility and suboptimal associated inflammatory response, intravitreal hydrogels today have become a potential solution to current treatment complications, especially for preventable neural retinal diseases such as macular degeneration and glaucoma. There are several treatments that are used for such therapy. However, many of them exhibit various problems and limitations. In order to be able to solve these drawbacks, the use of injectable hydrogels as drug delivery materials can improve the success of the therapy [138].

Non-expulsion of a depleted non-biodegradable embed may aggravate the visual tissue. Thus, it is better if it is eliminated after the medication is depleted. There is no requirement for expulsion after the medical procedure. This implies that there will be a lesser danger of difficulties related to the significant visual medical procedure. Trends in systems have seen in situ gel frameworks become an exploration hotspot—particularly for improvements in responsive hydrogels, such as ion-sensitive hydrogels, thermo-sensitive hydrogels, and pH-sensitive hydrogel [139].

Alternative Injection Locations

Intravitreal injection is one of the current methods of pharmaceutical delivery mechanisms performed as often as monthly, which can result in resistance [116]. Currently, there are alternative intraocular injections with respect to the location and drug being delivered that might be more useful [140]. These alternatives include sub-conjunctival injection and parabulbar injection, which take place underneath the conjunctiva for trans-scleral delivery [141]. Practically, a sustained in vivo drug release for up to 4 weeks was shown following sub-conjunctival injection of dorzolamide-loaded polymer disks [142]. Dorzolamide is a carbonic anhydrase inhibitor, and it works by decreasing the pressure in the eye [143].

Another alternative is sub-retinal injection, which is delivered beneath the retina to the sub-retinal fluid. Sub-retinal injection has more direct effects on the targeted cells in the sub-retinal space, providing a new therapeutic method for vitreoretinal diseases—especially when gene therapy and/or cell therapy is involved [144]. Via this effective technique, clinicians can administer drugs such as anti-VEGF [145] and steroids [146], among others, directly into the back of the eye to increase the drug concentration in the vitreous humor and the retina.

10. Pharmacokinetics of Intravitreal Hydrogel Drug Release

With this approach, the drug-loaded polymer complex is administered intravitreously at the targeted posterior segment of the eye via a minimally invasive procedure. The injectable systems utilize an anti-glaucoma drug-loaded polymer drug delivery system [36]. The polymer derived from ethylene-vinyl acetic acid is known by its non-degradable nature, which prompts a resistant reaction because of the extended presence of an unfamiliar body. The pace of drug release of medications from these frameworks is variable [147].
The water solvency of the medication influences its adequacy, on the grounds that hydrophilic medications cooperate ineffectively with biodegradable polymers due to their hydrophobic nature. The water solubility of some drugs has a direct influence on the drug release efficacy due to their hydrophilic nature [148].

Once the anti-glaucoma drug-loaded hydrogel has been injected into the target, the drug molecules are distributed by Fickian diffusion from the vitreous humor to the surrounding ocular tissues within reach and the target retinal sites. The vitreous body is not a hindering factor of drug diffusion of some soluble proteins; however, it limits the mobility of various drug delivery systems, such as drug-loaded nanoparticles. The vitreous humor facilitates an increased rate of low-molecular-weight drug diffusion, since the mesh size in the vitreous network has been estimated to be ~500 nm. Drugs administered intravitreally normally have a diameter of 10 nm and below, e.g., ranibizumab (4.1 nm) and bevacizumab (6.5 nm). Distribution of therapeutic drugs and the biodegradation of self-assembled polymers and nanoparticulate transporters have previously been confirmed [149].

Intravitreal injection of the biopolymer–drug complex into the sub-conjunctival space and posterior target site can lead to prolonged delivery over weeks or months, compared with simple topical application, which would last at most of a few hours or days [54]. Biodegradable and non-biodegradable polymers have been investigated for their application as injectable hydrogels for ocular treatment and therapy. Poly(ethylene-co-vinyl acetate)—a non-degradable polymer—exhibits extended drug delivery efficiency for a wide range of active drug ingredients, but unfortunately exhibits poor biodegradability, and the continued retention of implanted foreign body material causes immune responses. On the other hand, biodegradable polymers such as silk fibroin, chitosan, and alginate may be the best alternatives for intravitreal injection, as they are more suitably biodegradable [150].

11. Discussion

In this review, various therapeutic anti-glaucometic drug delivery systems were investigated, and the most promising ones were highlighted. This survey elaborates the importance of using natural biopolymer-based injectable hydrogels rather than other therapeutic drug delivery systems utilizing synthetic materials, because of the former’s biodegradability, biocompatibility, and non-immunogenic properties.

Numerous studies have proven that natural biopolymers rarely cause adverse inflammatory and immune responses when in contact with ocular tissue, as opposed to synthetic polymers. The therapeutic injectability of these drug-loaded polymer complexes has successfully overcome anatomical and physiological ocular barriers, thus increasing bioavailability and therapeutic efficacy. Unlike other therapeutic drug delivery systems that have failed to permeate through the ocular barriers, injectable hydrogels have the potential for prolonged sustainability in the delivery of ocular drugs. The potential improvement in patient compliance and persistence for optimal outcomes with the help of these systems is unprecedented.

12. Conclusions

The development of intravitreal injectable hydrogel drug delivery systems is a promising approach for the treatment of ophthalmological diseases—particularly glaucoma. Ocular treatment remains challenging for scientists because of physiological ocular barriers to any foreign substances. Thus far, using intravitreal injectable hydrogels for the treatment of ocular disease has facilitated the delivery of drugs to the targeted area at the desired dosage, with improved properties of penetration, bioavailability, and extended retention time for the release of the drugs. All of this has been achieved by encapsulating drugs into hydrogels made from naturally obtained biopolymers. Therefore, the assessment presented in this review indicates that hydrogels made from natural biopolymers have the ability to overcome the limitations of conventional ocular treatment, and could hence become potential sources and suitable matrices with excellent biocompatibility, acting as useful vehicles for the delivery of drugs. Researchers should devote more attention to
the production and application of intravitreal hydrogels made from natural polymers to deliver drugs to targeted areas for the treatment of glaucoma.

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References
1. Bro, T.; Wickström, K.; Lindén, C. The future is old—Patients with topical ocular hypotensive treatment in the Nordic region between 2008 and 2017 with projections for 2040. Acta Ophthalmol. 2021, 99, e1442–e1448. [CrossRef] [PubMed]
2. Sánchez, G.M.; del Pozo, C.E.; Medina, J.R.; Naude, J.; Solorzano, A.B. Numerical simulation of the aqueous humor flow in the eye drainage system; a healthy and pathological condition comparison. Mod. Eng. Phys. 2020, 83, 82–92. [CrossRef] [PubMed]
3. Cholkar, K.; Dasari, S.R.; Pal, D.; Mitra, A.K. Eye: Anatomy, physiology and barriers to drug delivery. In Ocular Transporters and Receptors; Elsevier: Amsterdam, The Netherlands, 2013; pp. 1–36.
4. Modarreszadeh, S.; Abouali, O.; Ghaffarieh, A.; Ahmadi, G. Physiology of aqueous humor dynamic in the anterior chamber due to rapid eye movement. Physiol. Behav. 2014, 135, 112–118. [CrossRef] [PubMed]
5. Downie, L.E.; Bandlitz, S.; Bergmanson, J.P.; Craig, J.P.; Dutta, D.; Maldonado-Codina, C.; Ngo, W.; Siddireddy, J.S.; Wolfsohn, J.S. BCLA CLEAR—Anatomy and physiology of the anterior eye. Contact Lens Anterior Eye 2021, 44, 132–156. [CrossRef] [PubMed]
6. Buffault, J.; Labbé, A.; Hamard, P.; Brignole-Baudouin, F.; Baudouin, C. The trabecular meshwork: Structure, function and clinical implications. A review of the literature. J. Fr. Ophthalmol. 2020, 43, e217–e230. [CrossRef]
7. Silva, A.F.; Pimenta, F.; Alves, M.A.; Oliveira, M.S. Flow dynamics of vitreous humour during saccadic eye movements. J. Mech. Behav. Biomed. Mater. 2020, 110, 103860. [CrossRef]
8. Yadav, K.S.; Rajpurohit, R.; Sharma, S. Glaucoma: Current treatment and impact of advanced drug delivery systems. Life Sci. 2019, 221, 362–376. [CrossRef]
9. El Hofty, N.M.; Azim, E.A.A.; Hathout, M.A.; Elkheshen, S.A. Glaucoma: Management and Future Perspectives for Nanotechnology-Based Treatment Modalities. Eur. J. Pharm. Sci. 2020, 158, 105648. [CrossRef]
10. Varela-Fernández, R.; Díaz-Tomé, V.; Luaces-Rodriguez, A.; Conde-Penedo, A.; García-Otero, X.; Luzardo-Álvarez, A.; Fernández-Ferreiro, A.; Otero-Espinar, F.J. Drug Delivery to the Posterior Segment of the Eye: Biopharmaceutical and Pharmacokinetic Considerations. Pharmaceutics 2020, 12, 269. [CrossRef]
11. Baudouin, C.; Kolko, M.; Melik-Parsadiantz, S.; Messmer, E.M. Inflammation in Glaucoma: From the back to the front of the eye, and beyond. Prog. Retin. Eye Res. 2020, 83, 100916. [CrossRef]
12. Wu, A.; Khawaja, A.P.; Pasquale, L.R.; Stein, J.D. A review of systemic medications that may modulate the risk of glaucoma. Eye 2020, 34, 12–28. [CrossRef]
13. Belay, D.B.; Derseh, M.; Damtie, D.; Shiferaw, Y.A.; Adigeh, S.C. Longitudinal analysis of intraocular pressure and its associated risk factors of glaucoma patients using Bayesian linear mixed model: A data from Felege Hiwot Hospital, Ethiopia. Sci. Afr. 2020, 8, 362–376. [CrossRef]
14. Yang, X.-L.; Van Der Merwe, Y.; Sims, J.; Parra, C.; Ho, L.C.; Schuman, J.S.; Wollstein, G.; Lathrop, K.L.; Chan, K.C. Age-related Changes in Eye, Brain and Visuomotor Behavior in the DBA/2J Mouse Model of Chronic Glaucoma. Sci. Rep. 2018, 8, 4643. [CrossRef] [PubMed]
15. Haeck, I.M.; Rouwen, T.J.; Mik, L.T.-D.; de Bruin-Weller, M.S.; Bruijnzeel-Koomen, C.A. Topical corticosteroids in atopic dermatitis and the risk of glaucoma and cataracts. J. Am. Acad. Dermatol. 2011, 64, 275–281. [CrossRef] [PubMed]
16. Guglielmi, P.; Carradori, S.; Campestrile, C.; Poco, G. Novel therapies for glaucoma: A patent review (2013–2019). Expert Opin. Ther. Patents 2019, 29, 769–780. [CrossRef]
17. Singh, M.; Dev, D.; Prasad, D. A Recent Overview: In Situ Gel Smart Carriers for Ocular Drug Delivery. J. Drug Deliv. Ther. 2021, 11, 195–205. [CrossRef]
18. Bahram, M.; Mohseni, N.; Moghtader, M. An Introduction to Hydrogels and Some Recent Applications. In Emerging Concepts in Analysis and Applications of Hydrogels; IntechOpen: London, UK, 2016. [CrossRef]
19. Xi, L.; Wang, T.; Zhao, F.; Zheng, Q.; Li, X.; Luo, J.; Liu, J.; Quan, D.; Ge, J. Evaluation of an Injectable Thermosensitive Hydrogel As Drug Delivery Implant for Ocular Glaucoma Surgery. PLoS ONE 2014, 9, e100632. [CrossRef]
20. Wang, K.; Han, Z. Injectable hydrogels for ophthalmic applications. J. Control. Release 2017, 268, 212–224. [CrossRef]
Cordeiro, F.; Tian, K.; Shibata-Germanos, S.; Pahlitzsch, M. Current perspective of neuroprotection and glaucoma. *Clin. Ophthalmol.* 2015, 9, 2109–2118. [CrossRef]

Pakravan, M.; Yazdani, S.; Javadi, M.-A.; Amini, H.; Behroozi, Z.; Ziae, H.; Katibeh, M.; Solaimanizad, R.; Ghahari, E.; Yaseri, M. A Population-based Survey of the Prevalence and Types of Glaucoma in Central Iran. *Ophthalmology* 2013, 120, 1977–1984. [CrossRef]

Bertaud, S.; Aragno, V.; Baudouin, C.; Labbé, A. Le glaucome primitif à angle ouvertPrimary open-angle glaucoma. *Rev. Méd. Interne* 2018, 40, 445–452. [CrossRef] [PubMed]

Artiero-Castro, A.; Rodriguez-Jimenez, F.J.; Jendelova, P.; VanderWall, K.B.; Meyer, J.S.; Erceg, S. Glaucoma as a Neurodegenerative Disease Caused by Intrinsic Vulnerability Factors. *Prog. Neurobiol.* 2020, 193, 101817. [CrossRef] [PubMed]

Sun, X.; Dai, Y.; Chen, Y.; Yu, D.-Y.; Cringle, S.J.; Chen, J.; Kong, X.; Wang, X.; Jiang, C. Primary angle closure glaucoma: We know what and we don’t know. *Prog. Retin. Eye Res.* 2017, 57, 26–45. [CrossRef]

Kesav, N.; Palestine, A.G.; Kahook, M.Y.; Pantcheva, M.B. Current management of uveitis-associated ocular hypertension and glaucoma. *Surv. Ophthalmol.* 2019, 65, 597–407. [CrossRef] [PubMed]

J边gagam, D.R.; Wu, L.; Lowe, T.L. Nanoparticles for drug delivery to the anterior segment of the eye. *Adv. Drug Deliv. Rev.* 2017, 122, 31–64. [CrossRef]

Ribeiro, A.M.; Figueiras, A.; Veiga, F. Improvements in Topical Ocular Drug Delivery Systems: Hydrogels and Contact Lenses. *J. Pharm. Pharm. Sci.* 2015, 18, 683–695. [CrossRef]

Kompella, U.B.; Hartman, R.R.; Patil, M.A. Extraocular, periocular, and intraocular routes for sustained drug delivery for glaucoma. *Prog. Retin. Eye Res.* 2020, 82, 100901. [CrossRef]

Anjali, S.; Rameshwar, D.; Shivani, D.; Ranjit, S. Hydrogels in ophthalmic drug delivery system—A mini review. *Asian Pac. J. Health Sci.* 2018, 5, 96–104. [CrossRef]

Wadhwa, A.; Jhadav, C.; Yadav, K.S. Bimatoprost: Promising novel drug delivery systems in treatment of glaucoma. *J. Drug Deliv. Sci. Technol.* 2022, 69, 103156. [CrossRef]

Kumar, V.; Abu Zaalan, K.A.; Bezzabotnov, A.I.; Dushina, G.N.; Shradqa, A.S.S.; Rustamova, Z.S.; Frolow, M.A. Bleb-Independent Glaucoma Surgery to Activate the Uveolympathic Route of Non-Trabecular Aqueous Humor Outflow: Short-Term Clinical and OCT Results. *Vision* 2022, 6, 4. [CrossRef]

Yu, S.; Wang, Q.-M.; Wang, X.; Liu, D.; Zhang, W.; Ye, T.; Yang, X.; Pan, W. Liposome incorporated ion sensitive in situ gels for ophthalmic delivery of timolol maleate. *Int. J. Pharm.* 2015, 480, 128–136. [CrossRef] [PubMed]

Occhiutto, M.L.; Maranhão, R.C.; Costa, V.P.; Konstas, A.G. Nanotechnology for Medical and Surgical Glaucoma Therapy—A Review. *Adv. Ther.* 2020, 37, 155–199. [CrossRef] [PubMed]

Tabak, S.; Schreiber-Avisar, S.; Beit-Yannai, M. Influence of Anti-Glaucoma Drugs on Uptake of Extracellular Vesicles by Trabecular Meshwork Cells. *Int. J. Nanomed.* 2021, 16, 1067–1081. [CrossRef] [PubMed]

Ilochonwu, B.C.; Urtti, A.; Hennink, W.E.; Vermond, T. Intravitreal hydrogels for sustained release of therapeutic proteins. *J. Control. Release* 2020, 326, 419–441. [CrossRef] [PubMed]

Waduthanthri, K.D.; He, Y.; Montemagno, C.; Cetinel, S. An injectable peptide hydrogel for reconstruction of the human trabecular meshwork. *Acta Biomater.* 2019, 100, 244–254. [CrossRef]

Jayanetti, V.; Sandhu, S.; Lusthaus, J.A. The Latest Drugs in Development That Reduce Intraocular Pressure in Ocular Hypertension and Glaucoma. *J. Exp. Pharmacol.* 2012, 683–695. [CrossRef]

Artero-Castro, A.; Rodriguez-Jimenez, F.J.; Jendelova, P.; VanderWall, K.B.; Meyer, J.S.; Erceg, S. Glaucoma as a Neurodegenerative Disease Caused by Intrinsic Vulnerability Factors. *Prog. Neurobiol.* 2020, 193, 101817. [CrossRef] [PubMed]

Arino, S.; Miyake, S.; Iwasaki, K.; Gozawa, M.; Matsumura, T.; Takamura, Y.; Inatani, M. Randomised Clinical Trial for Glaucoma Surgery to Activate the Uveolymphatic Route of Non-Trabecular Aqueous Humor Outflow: Short-Term Clinical and OCT Results. *Vision* 2022, 6, 4. [CrossRef]

Lee, D.A.; Higginbotham, E.J. Glaucoma and its treatment: A review. *Am. J. Health Syst. Pharm.* 2005, 62, 691–699. [CrossRef]

Miller, P.E.; Eaton, J.S. Medical anti-glaucoma therapy: Beyond the drop. *Vet. Ophthalmol.* 2020, 24, 2–15. [CrossRef] [PubMed]

Huang, J.; Peng, T.; Li, Y.; Zhan, Z.; Zeng, Y.; Huang, Y.; Pan, X.; Wu, C.-Y.; Wu, C. Ocular Cubosome Drug Delivery System for Timolol Maleate: Preparation, Characterization, Cytotoxicity, Ex Vivo, and In Vivo Evaluation. *AAPS PharmSciTech* 2017, 18, 2919–2926. [CrossRef] [PubMed]

Quiroz-Mercado, H.; Ivri, E.; Gonzalez-Salinas, R.; Kourtis, I.C.; Gilbert, J.; Pérez-Vázquez, J.F.; Blumenkranz, M.; Jiménez-Román, J.; Marcellino, G. Clinical Evaluation of a Novel Electromechanical Topical Ocular Drug Delivery System: Two Phase 1 Proof of Concept Studies. *Clin. Ophthalmol.* 2020, 14, 139–147. [CrossRef] [PubMed]

Nocentini, A.; Ceruso, M.; Bua, S.; Lomelino, C.L.; Andring, J.T.; McKenna, R.; Lanzi, C.; Sgambellone, S.; Pecori, R.; Matucci, R.; et al. Discovery of β-Adrenergic Receptors Blocker–Carbonic Anhydrase Inhibitor Hybrids for Multitargeted Antiglaucoma Therapy. *J. Med. Chem.* 2018, 61, 5380–5394. [CrossRef] [PubMed]
47. Holló, G.; Katsanos, A.; Boboridis, K.G.; Irkeç, M.; Konstas, A.G.P. Preservative-Free Prostaglandin Analogs and Prostaglandin/Timolol Fixed Combinations in the Treatment of Glaucoma: Efficacy, Safety and Potential Advantages. Drugs 2017, 78, 39–64. [CrossRef]

48. Luthscha, J.; Goldberg, I. Current management of glaucoma. Med. J. Aust. 2019, 210, 180–187. [CrossRef]

49. Holló, G.; Katsanos, A.; Boboridis, K.G.; Irkeç, M.; Konstas, A.G.P. Preservative-Free Prostaglandin Analogs and Prostaglandin/Timolol Fixed Combinations in the Treatment of Glaucoma: Efficacy, Safety and Potential Advantages. Drugs 2017, 78, 39–64. [CrossRef]

50. Figus, M.; Agnifili, L.; Lanzini, M.; Brescia, L.; Sartini, F.; Mastropasqua, L.; Posarelli, C. Topical preservative-free ophthalmic treatments: An unmet clinical need. Expert Opin. Drug Deliv. 2020, 18, 655–672. [CrossRef]

51. Xu, E.; Campaillone, O.H.; Ye, X.; Jin, Z.; Liu, D.; BeMiller, J.N. Advances in conversion of natural biopolymers: A reactive extrusion (REX)—enzyme-combined strategy for starch/protein-based food processing. Trends Food Sci. Technol. 2020, 99, 167–180. [CrossRef]

52. Samadian, H.; Maleki, H.; Fathollahi, A.; Salehi, M.; Gholizadeh, S.; Derakhshankhah, H.; Allahyari, Z.; Jaymand, M. Naturally occurring biological macromolecules-based hydrogels: Potential biomaterials for peripheral nerve regeneration. Int. J. Biol. Macromol. 2020, 154, 795–817. [CrossRef]

53. Meng, L.; Shao, C.; Cui, C.; Xu, F.; Lei, J.; Yang, J. Autonomous Self-Healing Silk Fibroin Injectable Hydrogels Formed via Surfactant-Free Hydrophobic Association. ACS Appl. Mater. Interfaces 2020, 12, 1628–1639. [CrossRef] [PubMed]

54. Silva, M.M.; Calado, R.; Marto, J.; Betencourt, A.; Almeida, A.J.; Gonçalves, L.M.D. Chitosan Nanoparticles as a Mucoadhesive Drug Delivery System for Ocular Administration. Mar. Drugs 2017, 15, 370. [CrossRef] [PubMed]

55. Zhang, W.; Zhao, X. Alginate hydrogel dressings for advanced wound management. Int. J. Biol. Macromol. 2020, 162, 1414–1428. [CrossRef] [PubMed]

56. Bhattacharjee, P.; Fernández-Pérez, J.; Ahearn, M. Potential for combined delivery of riboflavin and all-trans retinoic acid, from silk fibroin for corneal bioengineering. Mater. Sci. Eng. C 2019, 105, 110093. [CrossRef]

57. Applegate, M.B.; Partlow, B.P.; Coburn, J.; Marelli, B.; Pirie, C.; Pineda, R.; Kaplan, D.L.; Omenetto, F.G. Photocrosslinking of silk fibroin for corneal bioengineering. Mater. Sci. Eng. C 2018, 86, 105–11031. [CrossRef]

58. Galam, N.; Tulay, P.; Adali, T. The Wonders of Silk Fibroin Biomaterials in the Treatment of Breast Cancer. Crit. Rev. Eukaryot. Gene Expr. 2018, 28, 129–134. [CrossRef]

59. Galam, N.; Tulay, P.; Adali, T. In Vitro MCF-7 Cells Apoptosis Analysis of Carboplatin Loaded Silk Fibroin Particles. Molecules 2020, 25, 1110. [CrossRef]

60. Abdel-Naby, W.; Cole, B.; Liu, A.; Liu, J.; Wan, P.; Guaiquil, V.H.; Schreiner, R.; Infanger, D.; Lawrence, B.D.; Rosenblatt, M.I. Silk-derived protein enhances corneal epithelial migration, adhesion, and proliferation. Investig. Ophthalmol. Vis. Sci. 2017, 58, 1425–1433. [CrossRef]

61. highlight, C.B.; Rodell, C.B.; Burdick, J.A. Direct 3D Printing of Shear-Thinning Hydrogels into Self-Healing Hydrogels. Adv. Mater. 2015, 27, 5075–5079. [CrossRef] [PubMed]

62. Wang, H.; Shi, J.; Wang, Y.; Yin, Y.; Wang, L.; Liu, J.; Liu, Z.; Duan, C.; Zhu, P.; Wang, C. Promotion of cardiac differentiation of brown adipose derived stem cells by chitosan hydrogel for repair after myocardial infarction. Biomaterials 2014, 35, 3986–3998. [CrossRef]

63. Che, I.S.; Park, C.G.; Huh, B.K.; Cho, M.O.; Khatun, Z.; Li, Z.; Kang, S.-W.; Bin Choy, Y.; Huh, K.M. Thermosensitive hexaoyl glycol chitosan-based ocular delivery system for glaucoma therapy. Acta Biomater. 2016, 39, 124–132. [CrossRef]

64. Skwarczynska, A.L.; Binias, D.; Maniukiewicz, W.; Modrzewińska, Z.; Douglas, T.E. The mineralization effect on chitosan hydrogel structure containing collagen and alkaline phosphatase. J. Mol. Struct. 2019, 1187, 86–97. [CrossRef]

65. Franca, J.R.;oureaux, G.; Fuscaldi, L.L.; Ribero, T.G.; Castilho, R.O.; Yoshiida, M.I.; Cardoso, V.N.; Fernandes, S.; Nogueira, J.C.; et al. Chitosan/hydroxyethyl cellulose inserts for sustained-release of dorzolamide for glaucoma treatment: In vitro and in vivo evaluation. Int. J. Pharm. 2019, 570, 118662. [CrossRef] [PubMed]

66. Zamboulis, A.; Nanaki, S.; Michailidou, G.; Koumentakou, I.; Lazaridou, M.; Ainali, N.M.; Xanthopoulou, E.; Bikiaris, D.N. Chitosan and its Derivatives for Ocular Delivery Formulations: Recent Advances and Developments. Polymers 2020, 12, 1519. [CrossRef]
72. Popa, L.; Ghica, M.V.; Dinu-Pirvu, C.E.; Irimia, T. Chitosan: A good candidate for sustained release ocular drug delivery systems. In *Chitin-Chitosan—Myriad Functionalities in Science and Technology*; InTech: London UK, 2018; pp. 283–310.

73. Dekamin, M.G.; Karimi, Z.; Latifidoost, Z.; Ilkhanizadeh, S.; Daemi, H.; Naimi-Jamal, M.R.; Barikani, M. Algicin acid: A mild and renewable bifunctional heterogeneous biopolymeric organocatalyst for efficient and facile synthesis of polyhydroquinolines. *Int. J. Biol. Macromol.* 2018, 108, 1273–1280. [CrossRef]

74. Matsumoto, Y.; Ishii, D.; Iwata, T. Synthesis and characterization of algicin acid ester derivatives. *Carbohydr. Polym.* 2017, 171, 229–235. [CrossRef] [PubMed]

75. Karmakar, S.; Manna, S.; Kabiraj, S.; Jana, S. Recent progress in alginate-based carriers for ocular targeting of therapeutics. *Food Hydrocoll. Health 2022*, 2, 100071. [CrossRef]

76. Reddy, S.G. [CrossRef]

77. Panyamao, P.; Ruksiriwanich, W.; Sirisa-Ard, P.; Charumanee, S. Injectable Thermosensitive Chitosan/Pullulan-Based Hydrogels with Improved Mechanical Properties and Swelling Capacity. *Polymers* 2020, 12, 2514. [CrossRef] [PubMed]

78. Saeaeh, K.; Thummarungsan, N.; Paradee, N.; Choeichom, P.; Phasuksom, K.; Lerdwijitjarud, W.; Sirivat, A. Soft and highly responsive multi-walled carbon nanotube/pullulan hydrogel composites as electroactive materials. *Eur. Polym. J.* 2019, 120, 109231. [CrossRef]

79. Coltell, M.-B.; Danti, S.; De Clerck, K.; Lazzeri, A.; Morganti, P. Pullulan for Advanced Sustainable Body- and Skin-Contact Applications. *J. Funct. Biomater.* 2020, 11, 2020. [CrossRef]

80. Larrañeta, E.; Henry, M.; Irwin, N.J.; Trotter, J.; Perminova, A.A.; Donnelly, R. Synthesis and characterization of hyaluronic acid hydrogels crosslinked using a solvent-free process for potential biomedical applications. *Carbohydr. Polym.* 2018, 181, 1194–1205. [CrossRef]

81. Egbü, R.; Brocchini, S.; Khaw, P.T.; Awuwad, S. Antibody loaded collapsible hyaluronic acid hydrogels for intraocular delivery. *Eur. J. Pharm. Biopharm.* 2018, 124, 95–103. [CrossRef]

82. Chang, W.-H.; Liu, P.-Y.; Lin, M.-H.; Lu, C.-J.; Chou, H.-Y.; Nian, C.-Y.; Jiang, Y.-T.; Hsu, Y.-H. Applications of Hyaluronic Acid in Ophthalmology and Contact Lenses. *Molecules* 2021, 26, 2485. [CrossRef]

83. Cheng, Y.-H.; Ko, Y.-C.; Chang, Y.-F.; Huang, S.-H.; Liu, C.-L. Thermosensitive chitosan-gelatin-based hydrogel containing curcumin-loaded nanoparticles and latanoprost as a dual-drug delivery system for glaucoma treatment. *Exp. Eye Res.* 2019, 179, 179–187. [CrossRef]

84. Campos, F.D.; Cassimiro, D.L.; Crespi, M.S.; Almeida, A.E.; Gremião, M.P. Preparation and characterisation of Dextran-70 hydrogel for controlled release of praziquantel. *Braz. J. Pharm. Sci.* 2013, 49, 75–83. [CrossRef]

85. Yao, Y.; Saw, P.E.; Nie, Y.; Wang, P.; Jiang, L.; Ye, X.; Chen, J.; Ding, T.; Xu, L.; Yao, H.; et al. Multifunctional sharp pH-responsive nanoparticles for targeted drug delivery and effective breast cancer therapy. *J. Mater. Chem. B* 2018, 7, 576–585. [CrossRef] [PubMed]

86. Kilic Bektas, C.; Burcu, A.; Gedikoglu, G.; Telek, H.H.; Ornek, F.; Hasirci, V. Methacrylated gelatin hydrogels as corneal stroma substitutes: In vivo study. *J. Biomater. Sci. Polym. Ed.* 2019, 30, 1803–1821. [CrossRef]

87. Silva, S.S.; Fernandes, E.M.; Pina, S.; Silva-Correia, J.; Vieira, S.; Oliveira, J.M.; Reis, R.L. 2.11 Polymers of biological origin. *Polymers* 2017, 9, 27–49. [CrossRef]

88. Bonetti, L.; De Nardo, L.; Faré, S. Thermo-Responsive Methylcellulose Hydrogels: From Design to Applications as Smart Biomaterials. *Tissue Eng. Part B Rev.* 2021, 27, 486–513. [CrossRef]

89. Thrimawithana, T.; Young, S.; Bunt, C.; Green, C.; Alany, R. In-vitro and in-vivo evaluation of carrageenan/methylcellulose polymeric systems for transscleral delivery of macromolecules. *Eur. J. Pharm. Sci.* 2011, 44, 399–409. [CrossRef]

90. Gupta, B.; Mishra, V.; Ghurat, S.; Momin, M.; Omri, A. Cellulosic Polymers for Enhancing Drug Bioavailability in Ocular Drug Delivery Systems. *Pharmaceuticals* 2021, 14, 1201. [CrossRef]

91. Kojima, N.; Tao, F.; Mihara, H.; Aoki, S. Methods for Engineering of Multicellular Spheroids to Reconstitute the Liver Tissue. In *Stem Cells and Cancer in Hepatology*; Academic Press: Cambridge, MA, USA, 2018; pp. 145–158. [CrossRef]

92. El-Feky, G.S.; Zayed, G.; Elshaier, Y.; Alsharif, F.M. Chitosan-Gelatin Hydrogel Crosslinked with Oxidized Sucrose for the Ocular Delivery of Timolol Maleate. *J. Pharm. Sci.* 2016, 107, 3098–3104. [CrossRef]

93. Wong, F.S.Y.; Tsang, K.K.; Chan, B.; Yao, K.M.; Lo, A.C.Y. Injectable cell-encapsulating composite alginate-collagen platform with inducible termination switch for safer ocular drug delivery. *Biomaterials* 2019, 201, 53–67. [CrossRef]

94. Sun, J.; Lei, Y.; Dai, Z.; Liu, X.; Huang, T.; Wu, J.; Xu, Z.P.; Sun, X. Sustained Release of Brimonidine from a New Composite Drug Delivery System for Treatment of Glaucoma. *ACS Appl. Mater. Interfaces* 2017, 9, 7990–7999. [CrossRef]

95. Gaspar-Pintilescu, A.; Stefan, L.M.; Antón, E.D.; Berger, D.; Matei, C.; Negreanu-Pirjol, T.; Moldovan, L. Physicochemical and Biological Properties of Gelatin Extracted from Marine Snail Rapana venosa. *Mar. Drugs* 2019, 17, 589. [CrossRef]

96. He, Z.; Xiong, L. Evaluation of Biological Properties of Collagen/Hyaluronic Acid Composite Scaffolds. *Polym. Polym. Compos.* 2013, 21, 457–462. [CrossRef]

97. Jain, E.; Hill, L.; Canning, E.; Sell, S.A.; Zustiak, S.P. Control of gelation, degradation and physical properties of polyethylene glycol hydrogels through the chemical and physical identity of the crosslinker. *J. Mater. Chem. B* 2017, 5, 2679–2691. [CrossRef] [PubMed]

98. Peroglio, M.; Grad, S.; Mortisen, D.; Sprecher, C.M.; Illien-Junger, S.; Alini, M.; Eglin, D. Injectable thermoreversible hyaluronan-based hydrogels for nucleus pulposus cell encapsulation. *Eur. Spine J.* 2011, 20, 839–849. [CrossRef] [PubMed]
126. Martin, N.; Youssef, G. Dynamic properties of hydrogels and fiber-reinforced hydrogels. J. Mech. Behav. Biomed. Mater. 2018, 85, 194–200. [CrossRef] [PubMed]

127. Kiyotake, E.A.; Douglas, A.W.; Thomas, E.E.; Nimmo, S.L.; Detamore, M.S. Development and quantitative characterization of the precursor rheology of hyaluronic acid hydrogels for bioprinting. Acta Biomater. 2019, 95, 176–187. [CrossRef] [PubMed]

128. Leone, G.; ConsDMI, S.; PardiNM, A.; BonechIC, C.; Tamasaki, G.; DonatiaM, A.; LamponiaM, S.; RossiaM, C.; Magnani, A. Enriched Gellan Gum hydrogel as visco-supplement. Carbohydr. Polym. 2020, 227, 115–347.

129. Luo, K.; Upadhyay, K.; Subhash, G.; Spearet, D.E. Transient-State Rheological Behavior of Poly(ethylene glycol) Diacrylate Hydrogels at High Shear Strain Rates. Macromolecules 2019, 52, 5860–5871. [CrossRef]

130. Gyles, D.A.; Castro, L.D.; Silva, J.O.C.; Costa, R.M.R. A review of the designs and prominent biomedical advances of natural and synthetic hydrogel formulations. Eur. Polym. J. 2017, 88, 373–392. [CrossRef]

131. Sun, W.; Chen, G.; Wang, F.; Qin, Y.; Wang, Z.; Nie, J.; Ma, G. Polyelectrolyte-complex multilayer membrane with gradient porous structure based on natural polymers for wound care. Carbohydr. Polym. 2018, 181, 183–190. [CrossRef]

132. Polat, T.G.; Duman, O.; Tünc, S. Preparation and characterization of environmentally friendly agar/x-carrageenan/montmorillonite nanocomposite hydrogels. Colloids Surf. A Physicochem. Eng. Asp. 2020, 602, 124987. [CrossRef]

133. Nguyen, D.D.; Luo, L.; Lai, J. Dendritic Effects of Injectable Biodegradable Thermogels on Pharmacotherapy of Inflammatory Glaucoma-Associated Degradation of Extracellular Matrix. Adv. Healthc. Mater. 2019, 8, e1900702. [CrossRef]

134. Mealy, J.E.; Chung, J.J.; Jeong, H.; Issadore, D.; Lee, D.; Athuri, P.; Burdick, J.A. Injectable Granular Hydrogels with Multifunctional Properties for Biomedical Applications. Adv. Mater. 2018, 30, 1705912. [CrossRef]

135. Vernon, R.B.; Gooden, M.D.; Preisinger, A.; Gebe, J.A. Controlled release of monoclonal antibodies from poly-l-lysine-coated alginate spheres within a scaffolded implant mitigates autoimmune responses to transplanted islets and limits systemic antibody toxicity. Mater. Sci. Eng. C 2018, 93, 390–398. [CrossRef]

136. Zarembsinski, T.L.; Doty, N.J.; Erickson, I.E.; Srinivas, R.; Wirostko, B.M.; Tew, W.P. Thiolated hyaluronan-based hydrogels crosslinked using oxidized glutathione: An injectable matrix designed for ophthalmic applications. Acta Biomater. 2014, 10, 94–103. [CrossRef] [PubMed]

137. Anwary, M.; Kumar, P.; du Toit, L.C.; Choonara, Y.E.; Pillay, V. Polymeric, injectable, intravitreal hydrogel devices for posterior segment applications and interventions. Artif. Cells Nanomed. Biotechnol. 2018, 46, 1074–1081. [CrossRef] [PubMed]

138. Fernández-Colino, A.; Quinteros, D.; Allemoni, D.A.; Girotti, A.; Palma, S.D.; Arias, F.J. Self-Assembling Elastin-Like Hydrogels for Timolol Delivery: Development of an Ophthalmic Formulation Against Glaucoma. Mol. Pharm. 2017, 14, 4498–4508. [CrossRef] [PubMed]

139. Bergers, G.; Hanahan, D. Modes of resistance to anti-angiogenic therapy. Nat. Cancer 2008, 8, 592–603. [CrossRef] [PubMed]

140. Gooch, N.; Burr, R.M.; Holt, D.J.; Gale, B.; Ambati, B. Design and In Vitro Biocompatibility of a Novel Ocular Drug Delivery Device. J. Funct. Biomater. 2013, 4, 14–26. [CrossRef]

141. Papadopoulos, Z. Recent Developments in the Treatment of Wet Age-related Macular Degeneration. Curr. Med. Sci. 2020, 40, 851–857. [CrossRef]

142. Natu, M.V.; Gaspar, M.N.; Ribeiro, C.F.; Cabrita, A.; de Sousa, H.C.; Gil, M.H. In vitro and in vivo evaluation of an intraocular implant for glaucoma treatment. Int. J. Pharm. 2011, 415, 73–82. [CrossRef] [PubMed]

143. Tambe, S.; Jain, D.; Amin, P. Simultaneous determination of dorzolamide and timolol by first-order derivative UV spectroscopy in simulated biological fluid for in vitro drug release testing. Spectrochim. Acta Part A Mol. Biomol. Spectrosc. 2021, 255, 119682. [CrossRef]

144. Sommersperger, M.; Weiss, J.; Nasserı, M.A.; Gehlbach, P.L.; Jordachita, I.; Navab, N. Real-time tool to layer distance estimation for robotic subretinal injection using intraoperative 4D OCT. Biomed. Opt. Express 2021, 12, 1089–100. [CrossRef]

145. Hwang, D.D.-J.; Kim, Y.W.; Woo, S.J.; Park, K.H. Comparison of Systemic Adverse Events Associated with Intravitreal Anti-VEGF Injection: Ranibizumab versus Bevacizumab. J. Korean Med. Sci. 2012, 27, 1580–1585. [CrossRef]

146. Jonas, J.; Rensch, F. Intravitreal Steroid Slow-Release Device Replacing Repeated Intravitreal Triamcinolone Injections for Sympathetic Ophthalmia. Eur. J. Ophthalmol. 2018, 18, 834–836. [CrossRef] [PubMed]

147. Nasef, M.M.; Aly, A.A.; Saidi, H.; Ahmad, A. Optimization of reaction parameters of radiation induced grafting of 1-vinylimidazole onto poly(ethylene-co-tetrafluoroethylene) using response surface method. Radiat. Phys. Chem. 2011, 80, 1222–1227. [CrossRef] [PubMed]

148. Zeng, Y.; Chen, J.; Li, Y.; Huang, J.; Huang, Z.; Huang, Y.; Wu, C. Thermo-sensitive gel in glaucoma therapy for enhanced bioavailability: In vitro characterization, in vivo pharmacokinetics and pharmacodynamics study. Life Sci. 2018, 212, 80–86. [CrossRef] [PubMed]

149. Al-Kinani, A.A.; Zidan, G.; Elsaid, N.; Seyfoddin, A.; Alani, A.W.G.; Alany, R.G. Ophthalmic gels: Past, present and future. Adv. Drug Deliv. Rev. 2018, 126, 113–126. [CrossRef] [PubMed]

150. Voss, K.; Falke, K.; Bernsdorf, A.; Grabow, N.; Kastner, C.; Sternberg, K.; Hokvakimyan, M. Development of a novel injectable drug delivery system for subconjunctival glaucoma treatment. J. Control. Release 2015, 214, 1–11. [CrossRef]