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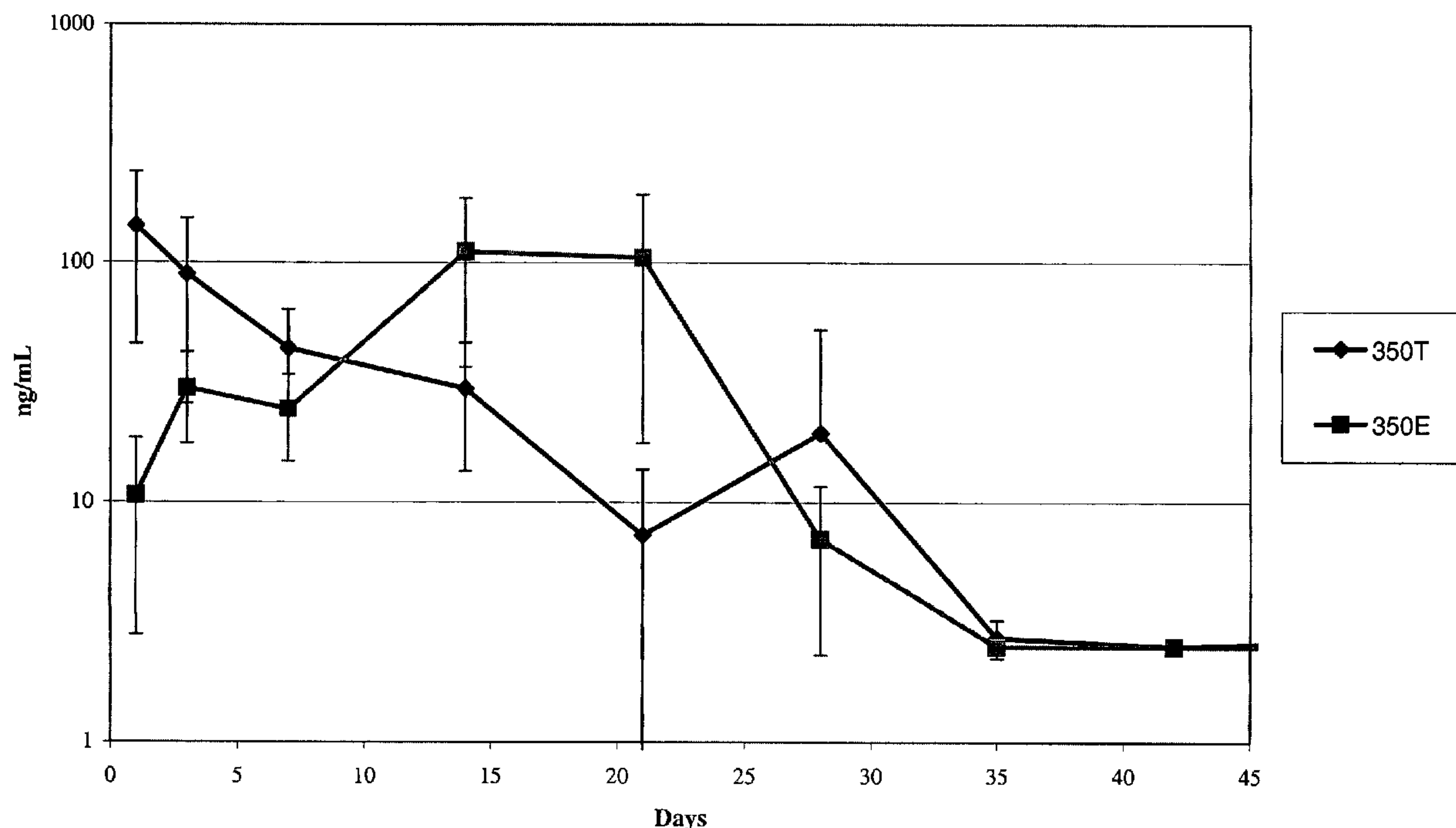
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(57) Abrégé/Abstract:

The invention provides biodegradable implants size for implantation in an ocular region and methods for treating medical conditions of the eye. The implants are formed from a mixture of hydrophilic end and hydrophobic end PLGA, and deliver active agents into an ocular region without a high burst release. Preferably the implant is made by a double extrusion process. Preferably at least 75% of the particles of the active agent have a diameter of less than 10 or 20 microns.

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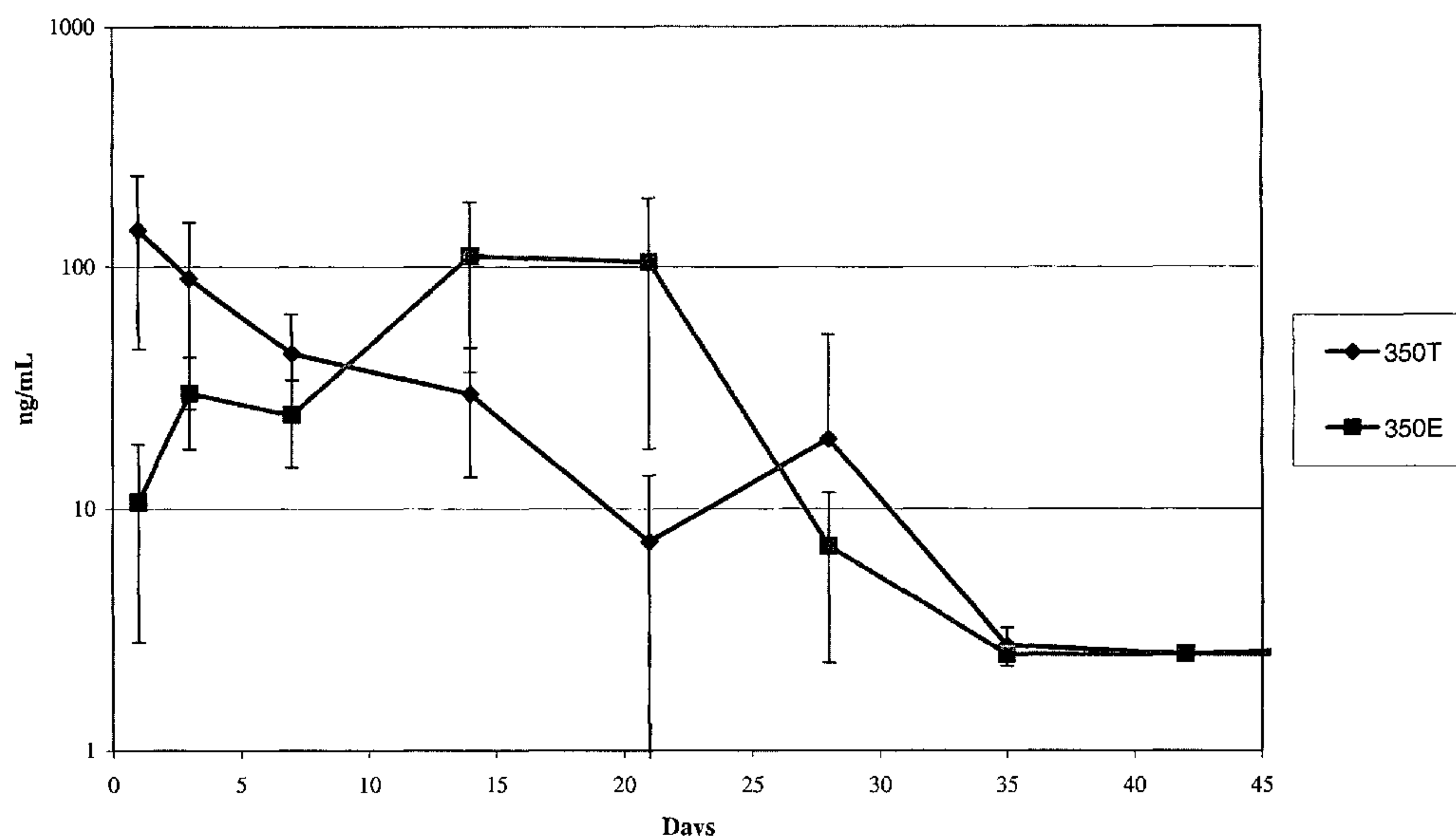
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OCULAR IMPLANT MADE BY A DOUBLE EXTRUSION PROCESS

by

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BACKGROUND

This invention relates to implants and methods for treating an ocular condition. In particular the present invention relates to implants and methods for treating an ocular condition by implanting into an ocular region or site a bioerodible implant comprising an active agent and a bioerodible polymer matrix, wherein the
15 implant is made by a double extrusion process. The bioerodible implants of this invention have varying and extended release rates to provide for improved kinetics of release of one or more active (therapeutic) agents over time.

An ocular condition can include a disease, ailment or condition which affects or involves the eye or one of the parts or regions of the eye. Broadly speaking the
20 eye includes the eyeball and the tissues and fluids which constitute the eyeball, the periocular muscles (such as the oblique and rectus muscles) and the portion of the optic nerve which is within or adjacent to the eyeball. An anterior ocular condition is a disease, ailment or condition which affects or which involves an anterior (i.e. front of the eye) ocular region or site, such as a periocular muscle, an eye lid or an eye ball
25 tissue or fluid which is located anterior to the posterior wall of the lens capsule or ciliary muscles. Thus, an anterior ocular condition primarily affects or involves, the

conjunctiva, the cornea, the conjunctiva, the anterior chamber, the iris, the posterior chamber (behind the retina but in front of the posterior wall of the lens capsule), the lens or the lens capsule and blood vessels and nerve which vascularize or innervate an anterior ocular region or site. A posterior ocular condition is a disease, ailment or
5 condition which primarily affects or involves a posterior ocular region or site such as choroid or sclera (in a position posterior to a plane through the posterior wall of the lens capsule), vitreous, vitreous chamber, retina, optic nerve (i.e. the optic disc), and blood vessels and nerves which vascularize or innervate a posterior ocular region or site.

10 Thus, a posterior ocular condition can include a disease, ailment or condition, such as for example, macular degeneration (such as non-exudative age related macular degeneration and exudative age related macular degeneration); choroidal neovascularization; acute macular neuroretinopathy; macular edema (such as cystoid macular edema and diabetic macular edema); Behcet's disease, retinal disorders,
15 diabetic retinopathy (including proliferative diabetic retinopathy); retinal arterial occlusive disease; central retinal vein occlusion; uveitic retinal disease; retinal detachment; ocular trauma which affects a posterior ocular site or location; a posterior ocular condition caused by or influenced by an ocular laser treatment; posterior ocular conditions caused by or influenced by a photodynamic therapy;
20 photocoagulation; radiation retinopathy; epiretinal membrane disorders; branch retinal vein occlusion; anterior ischemic optic neuropathy; non-retinopathy diabetic retinal dysfunction, retinitis pigmentosa and glaucoma. Glaucoma can be considered a posterior ocular condition because the therapeutic goal is to prevent the loss of or reduce the occurrence of loss of vision due to damage to or loss of retinal cells or
25 optic nerve cells (i.e. neuroprotection).

An anterior ocular condition can include a disease, ailment or condition, such as for example, aphakia; pseudophakia; astigmatism; blepharospasm; cataract; conjunctival diseases; conjunctivitis; corneal diseases; corneal ulcer; dry eye syndromes; eyelid diseases; lacrimal apparatus diseases; lacrimal duct obstruction;

myopia; presbyopia; pupil disorders; refractive disorders and strabismus. Glaucoma can also be considered to be an anterior ocular condition because a clinical goal of glaucoma treatment can be to reduce a hypertension of aqueous fluid in the anterior chamber of the eye (i.e. reduce intraocular pressure).

5 The present invention is concerned with and directed to an implant and methods for the treatment of an ocular condition, such as an anterior ocular condition or a posterior ocular condition or to an ocular condition which can be characterized as both an anterior ocular condition and a posterior ocular condition.

 Therapeutic compounds useful for the treatment of an ocular condition can
10 include active agents with, for example, an anti-neoplastic, anti-angiogenesis, kinase inhibition, anticholinergic, anti-adrenergic and/or anti-inflammatory activity.

 Macular degeneration, such as age related macular degeneration ("AMD") is a leading cause of blindness in the world. It is estimated that thirteen million Americans have evidence of macular degeneration. Macular degeneration results in a
15 break down the macula, the light-sensitive part of the retina responsible for the sharp, direct vision needed to read or drive. Central vision is especially affected. Macular degeneration is diagnosed as either dry (atrophic) or wet (exudative). The dry form of macular degeneration is more common than the wet form of macular degeneration, with about 90% of AMD patients being diagnosed with dry AMD. The wet form of
20 the disease usually leads to more serious vision loss. Macular degeneration can produce a slow or sudden painless loss of vision. The cause of macular degeneration is not clear. The dry form of AMD may result from the aging and thinning of macular tissues, depositing of pigment in the macula, or a combination of the two processes. With wet AMD, new blood vessels grow beneath the retina and leak blood
25 and fluid. This leakage causes retinal cells to die and creates blind spots in central vision.

Macular edema ("ME") can result in a swelling of the macula. The edema is caused by fluid leaking from retinal blood vessels. Blood leaks out of the weak vessel walls into a very small area of the macula which is rich in cones, the nerve endings that detect color and from which daytime vision depends. Blurring then occurs in the middle or just to the side of the central visual field. Visual loss can progress over a period of months. Retinal blood vessel obstruction, eye inflammation, and age-related macular degeneration have all been associated with macular edema. The macula may also be affected by swelling following cataract extraction. Symptoms of ME include blurred central vision, distorted vision, vision tinted pink and light sensitivity. Causes of ME can include retinal vein occlusion, macular degeneration, diabetic macular leakage, eye inflammation, idiopathic central serous chorioretinopathy, anterior or posterior uveitis, pars planitis, retinitis pigmentosa, radiation retinopathy, posterior vitreous detachment, epiretinal membrane formation, idiopathic juxtafoveal retinal telangiectasia, Nd:YAG capsulotomy or iridotomy. Some patients with ME may have a history of use of topical epinephrine or prostaglandin analogs for glaucoma. The first line of treatment for ME is typically anti-inflammatory drops topically applied.

Macular edema is a non-specific response of the retina to a variety of insults. It is associated with a number of diseases, including uveitis, retinal vascular abnormalities (diabetic retinopathy and retinal vein occlusive disease), a sequelae of cataract surgery (post-cataract cystoid macular oedema), macular epiretinal membranes, and inherited or acquired retinal degeneration. Macular edema involves the breakdown of the inner blood retinal barrier at the level of the capillary endothelium, resulting in abnormal retinal vascular permeability and leakage into the adjacent retinal tissues. The macula becomes thickened due to fluid accumulation resulting in significant disturbances in visual acuity (Ahmed I, Ai E. Macular disorders: cystoid macular oedema. In: Yanoff M, Duker JS, eds. Ophthalmology. London: Mosby; 1999:34; Dick J, Jampol LM, Haller JA. Macular edema. In: Ryan S, Schachat AP, eds. Retina. 3rd ed. St. Louis, MO: CV Mosby; 2001, v2, Section 2 chap 57:967-979).

Macular edema may occur in diseases causing cumulative injury over many years, such as diabetic retinopathy, or as a result of more acute events, such as central retinal vein occlusion or branch retinal vein occlusion.

In some cases macular edema resolves spontaneously or with short-term treatment. Therapeutic choices for macular oedema depend on the cause and severity of the condition. Currently there are no approved pharmacological therapies for macular edema. Focal/grid laser photocoagulation has been shown to be efficacious in the prevention of moderate visual loss for macular oedema due to diabetic retinopathy (Akduman L, Olk RS. The early treatment diabetic retinopathy study. In: Kertes PS, Conway MD, eds. Clinical trials in ophthalmology: a summary and practice guide. Baltimore, MD: Lippincott Williams & Wilkins; 1998:15-35; Frank RN. Etiologic mechanisms in diabetic retinopathy. In: Ryan S, Schachat AP, eds. Retina. 3rd ed. St. Louis, MO: CV Mosby; 2001, v2, Section 2, chap 71:1259-1294). Argon laser photocoagulation increased the likelihood of vision improvement in patients with macular oedema due to branch retinal vein occlusion (BRVO) (Orth D. The branch vein occlusion study. In: Kertes P, Conway M, eds. Clinical trials in ophthalmology: a summary and practice guide. Baltimore, MD: Lippincott Williams & Wilkins; 1998:113-127; Fekrat S, Finkelstein D. The Central Vein Occlusion Study. In: Kertes PS, Conway MD, eds. Clinical trials in ophthalmology: a summary and practice guide. Baltimore, MD: Lippincott Williams & Wilkins; 1998:129-143), but not in patients with macular oedema due to central retinal vein occlusion (CRVO) (Fekrat and Finkelstein 1998, *supra*; Clarkson JG. Central retinal vein occlusion. In: Ryan S, Schachat AP, eds. Retina. 3rd ed. St. Louis, MO: CV Mosby; 2001, v2, chap 75:1368 -1375). For CRVO, there are no known effective therapies.

An anti-inflammatory (i.e. immunosuppressive) agent can be used for the treatment of an ocular condition, such as a posterior ocular condition, which involves inflammation, such as an uveitis or macula edema. Thus, topical or oral glucocorticoids have been used to treat uveitis. A major problem with topical and oral drug administration is the inability of the drug to achieve an adequate (i.e.

therapeutic) intraocular concentration. See e.g. Bloch-Michel E. (1992). *Opening address: intermediate uveitis*, In *Intermediate Uveitis*, Dev. Ophthalmol, W.R.F. Böke et al. editors., Basel: Karger, 23:1-2; Pinar, V., et al. (1997). *Intraocular inflammation and uveitis*" In *Basic and Clinical Science Course. Section 9* (1997-
5 1998) San Francisco: American Academy of Ophthalmology, pp. 57-80, 102-103, 152-156; Böke, W. (1992). *Clinical picture of intermediate uveitis*, In *Intermediate Uveitis*, Dev. Ophthalmol. W.R.F. Böke et al. editors., Basel: Karger, 23:20-7; and Cheng C-K et al. (1995). *Intravitreal sustained-release dexamethasone device in the treatment of experimental uveitis*, Invest. Ophthalmol. Vis. Sci. 36:442-53.

10 Systemic glucocorticoid administration can be used alone or in addition to topical glucocorticoids for the treatment of uveitis. However, prolonged exposure to high plasma concentrations (administration of 1 mg/kg/day for 2-3 weeks) of steroid is often necessary so that therapeutic levels can be achieved in the eye.

Unfortunatley, these high drug plasma levels commonly lead to systemic side
15 effects such as hypertension, hyperglycemia, increased susceptibility to infection, peptic ulcers, psychosis, and other complications. Cheng C-K et al. (1995). *Intravitreal sustained-release dexamethasone device in the treatment of experimental uveitis*, Invest. Ophthalmol. Vis. Sci. 36:442-53; Schwartz, B. (1966). *The response of ocular pressure to corticosteroids*, Ophthalmol. Clin. North Am. 6:929-
20 89; Skalka, H.W. et al. (1980). *Effect of corticosteroids on cataract formation*, Arch Ophthalmol 98:1773-7; and Renfro, L. et al. (1992). *Ocular effects of topical and systemic steroids*, Dermatologic Clinics 10:505-12.

Additionally, delivery to the eye of a therapeutic amount of an active agent can be difficult, if not impossible, for drugs with short plasma half-lives since the
25 exposure of the drug to intraocular tissues is limited. Therefore, a more efficient way of delivering a drug to treat a posterior ocular condition is to place the drug directly in the eye, such as directly into the vitreous. Maurice, D.M. (1983).

Micropharmaceutics of the eye, Ocular Inflammation Ther. 1:97-102; Lee, V.H.L. et

al. (1989). *Drug delivery to the posterior segment* Chapter 25 In Retina. T.E. Ogden and A.P. Schachar eds., St. Louis: CV Mosby, Vol. 1, pp. 483-98; and Olsen, T.W. et al. (1995). *Human scleral permeability: effects of age, cryotherapy, transscleral diode laser, and surgical thinning*, Invest. Ophthalmol. Vis. Sci. 36:1893-1903.

Techniques such as intravitreal injection of a drug have shown promising results, but due to the short intraocular half-life of active agent, such as glucocorticoids (approximately 3 hours), intravitreal injections must be frequently repeated to maintain a therapeutic drug level. In turn, this repetitive process increases the potential for side effects such as retinal detachment, endophthalmitis, and cataracts. Maurice, D.M. (1983). *Micropharmaceutics of the eye*, Ocular Inflammation Ther. 1:97-102; Olsen, T.W. et al. (1995). *Human scleral permeability: effects of age, cryotherapy, transscleral diode laser, and surgical thinning*, Invest. Ophthalmol. Vis. Sci. 36:1893-1903; and Kwak, H.W. and D'Amico, D. J. (1992). *Evaluation of the retinal toxicity and pharmacokinetics of dexamethasone after intravitreal injection*, Arch. Ophthalmol. 110:259-66.

Additionally, topical, systemic, and periocular glucocorticoid treatment must be monitored closely due to toxicity and the long-term side effects associated with chronic systemic drug exposure sequelae. Rao, N.A. et al. (1997). *Intraocular inflammation and uveitis*, In Basic and Clinical Science Course. Section 9 (1997-1998) San Francisco: American Academy of Ophthalmology, pp. 57-80, 102-103, 152-156; Schwartz, B. (1966). *The response of ocular pressure to corticosteroids*, Ophthalmol Clin North Am 6:929-89; Skalka, H.W. and Pichal, J.T. (1980). *Effect of corticosteroids on cataract formation*, Arch Ophthalmol 98:1773-7; Renfro, L and Snow, J.S. (1992). *Ocular effects of topical and systemic steroids*, Dermatologic Clinics 10:505-12; Bodor, N. et al. (1992). *A comparison of intraocular pressure elevating activity of loteprednol etabonate and dexamethasone in rabbits*, Current Eye Research 11:525-30.

U.S. patent 6,217,895 discusses a method of administering a corticosteroid to the posterior segment of the eye, but does not disclose a bioerodible implant.

U.S. patent 5,501,856 discloses controlled release pharmaceutical preparations for intraocular implants to be applied to the interior of the eye after a
5 surgical operation for disorders in retina/vitreous body or for glaucoma.

U.S. patent 5,869,079 discloses combinations of hydrophilic and hydrophobic entities in a biodegradable sustained release implant, and describes a polylactic acid polyglycolic acid (PLGA) copolymer implant comprising dexamethasone. As shown by *in vitro* testing of the drug release kinetics, the 100-120 μ g 50/50
10 PLGA/dexamethasone implant disclosed did not show appreciable drug release until the beginning of the fourth week, unless a release enhancer, such as HPMC was added to the formulation.

U.S. patent No. 5,824,072 discloses implants for introduction into a suprachoroidal space or an avascular region of the eye, and describes a
15 methylcellulose (i.e. non-biodegradable) implant comprising dexamethasone. WO 9513765 discloses implants comprising active agents for introduction into a suprachoroidal or an avascular region of an eye for therapeutic purposes.

U.S. patent 4,997,652 and 5,164,188 disclose biodegradable ocular implants comprising microencapsulated drugs, and describes implanting microcapsules
20 comprising hydrocortisone succinate into the posterior segment of the eye.

U.S. patent 5,164,188 discloses encapsulated agents for introduction into the suprachoroid of the eye, and describes placing microcapsules and plaques comprising hydrocortisone into the pars plana. U.S. Patent Nos. 5,443,505 and 5,766,242 discloses implants comprising active agents for introduction into a suprachoroidal
25 space or an avascular region of the eye, and describes placing microcapsules and plaques comprising hydrocortisone into the pars plana.

Zhou *et al.* disclose a multiple-drug implant comprising 5-fluorouridine, triamcinolone, and human recombinant tissue plasminogen activator for intraocular management of proliferative vitreoretinopathy (PVR). Zhou, T, *et al.* (1998). *Development of a multiple-drug delivery implant for intraocular management of proliferative vitreoretinopathy*, Journal of Controlled Release 55: 281-295.

U.S. patent 6,046,187 discusses methods and compositions for modulating local anesthetic by administering one or more glucocorticosteroid agents before, simultaneously with or after the administration of a local anesthetic at a site in a patient.

10 U.S. patent 3,986,510 discusses ocular inserts having one or more inner reservoirs of a drug formulation confined within a bioerodible drug release rate controlling material of a shape adapted for insertion and retention in the "sac of the eye," which is indicated as being bounded by the surfaces of the bulbar conjunctiva of the sclera of the eyeball and the palpebral conjunctiva of the eyelid, or for placement
15 over the corneal section of the eye.

U.S. patent 6,369,116 discusses an implant with a release modifier inserted in a scleral flap.

EP 0 654256 discusses use of a scleral plug after surgery on a vitreous body, for plugging an incision.

20 U.S. patent 4,863,457 discusses the use of a bioerodible implant to prevent failure of glaucoma filtration surgery by positioning the implant either in the subconjunctival region between the conjunctival membrane overlying it and the sclera beneath it or within the sclera itself within a partial thickness sclera flap.

EP 488 401 discusses intraocular implants, made of certain polylactic acids, to be applied to the interior of the eye after a surgical operation for disorders of the retina/vitreous body or for glaucoma.

EP 430539 discusses use of a bioerodible implant which is inserted in the
5 suprachoroid.

U.S. patent 6,726,918 discusses implants for treating inflammation mediated conditions of the eye.

Significantly, it is known that PLGA co-polymer formulations of a bioerodible polymer comprising an active agent typically release the active agent
10 with a characteristic sigmoidal release profile (as viewed as time vs percent of total active agent released), that is after a relatively long initial lag period (the first release phase) when little if any active agent is released, there is a high positive slope period when most of the active agent is released (the second release phase) followed by another near horizontal (third) release phase, when the drug release reaches a plateau.

15 One of the alternatives to intravitreal injection to administer drugs is the placement of biodegradable implants under the sclera or into the subconjunctival or suprachoroidal space, as described in U.S. 4,863,457 to Lee; WO 95/13765 to Wong et al.; WO 00/37056 to Wong et al.; EP 430,539 to Wong; in Gould et al., *Can. J. Ophthalmol.* 29(4):168-171 (1994); and in Apel et al., *Curr. Eye Res.* 14:659-667
20 (1995).

Furthermore, the controlled release of drugs from polylactide/polyglycolide (PLGA) copolymers into the vitreous has been disclosed, e.g., in U.S. 5,501,856 to Ohtori et al. and EP 654,256 to Ogura.

Recent experimental work has demonstrated that uncapped PLGA degrades
25 faster than capped (end-capped) PLGA (Park et al., *J. Control. Rel.* 55:181-191

(1998); Tracy et al., *Biomaterials* 20:1057-1062 (1999); and Jong et al., *Polymer* 42:2795-2802 (2001). Accordingly, implants containing mixtures of uncapped and capped PLGA have been formed to modulate drug release. For example, U.S. 6,217,911 to Vaughn et al. ('911) and U.S. 6,309,669 to Setterstrom et al. ('669) disclose the delivery of drugs from a blend of uncapped and capped PLGA copolymer to curtail initial burst release of the drugs. In the '911 patent, the composition delivers non-steroidal anti-inflammatory drugs from PLGA microspheres made by a solvent extraction process or PLGA microcapsules prepared by a solvent evaporation process over a duration of 24 hours to 2 months. In the '669 patent, the composition delivers various pharmaceuticals from PLGA microcapsules over a duration of 1-100 days. The PLGA microspheres or microcapsules are administered orally or as an aqueous injectable formulation. As mentioned above, there is poor partitioning of drug into the eye with oral administration. Furthermore, use of an aqueous injectable drug composition (for injecting into the eye) should be avoided since the eye is a closed space (limited volume) with intraocular pressure ranges that are strictly maintained. Administration of an injectable may increase intraocular volume to a point where intraocular pressures would then become pathologic.

Potent corticosteroids such as dexamethasone suppress inflammation by inhibiting edema, fibrin deposition, capillary leakage and phagocytic migration, all key features of the inflammatory response. Corticosteroids prevent the release of prostaglandins, some of which have been identified as mediators of cystoid macular oedema (Leopold IH. Nonsteroidal and steroidal anti-inflammatory agents. In: Sears M, Tarkkanen A, eds. *Surgical pharmacology of the eye*. New York, NY: Raven Press; 1985:83-133; Tennant JL. Cystoid maculopathy: 125 prostaglandins in ophthalmology. In: Emery JM, ed. *Current concepts in cataract surgery: selected proceedings of the fifth biennial cataract surgical congress, Section 3*. St. Louis, MO: CV Mosby; 1978:360-362). Additionally, corticosteroids including dexamethasone have been shown to inhibit the expression of vascular endothelial growth factor (VEGF), a cytokine which is a potent promoter of vascular permeability (Nauck M,

Karakiulakis G, Perruchoud AP, Papakonstantinou E, Roth M. Corticosteroids inhibit the expression of the vascular endothelial growth factor gene in human vascular smooth muscle cells. *Eur J Pharmacol* 1998;341:309-315).

The use of dexamethasone to date, by conventional routes of administration, has yielded limited success in treating retinal disorders, including macular oedema, largely due to the inability to deliver and maintain adequate quantities of the drug to the posterior segment without resultant toxicity. After topical administration of dexamethasone, only about 1% reaches the anterior segment, and only a fraction of that amount moves into the posterior segment (Lee VHL, Pince KJ, Frambach DA, Martini B. Drug delivery to the posterior segment. In: Ogden TE, Schachat AP, eds. *Retina*. St. Louis, MO: CV Mosby, 1989, chap 25:483-498). Although intravitreal injections of dexamethasone have been used, the exposure to the drug is very brief as the half-life of the drug within the eye is approximately 3 hours (Peyman GA, Herbst R. Bacterial endophthalmitis. *Arch Ophthalmol* 1974;91:416-418). Periocular and posterior sub-Tenon's injections of dexamethasone also have a short term treatment effect (Riordan-Eva P, Lightman S. Orbital floor steroid injections in the treatment of uveitis. *Eye* 1994;8 (Pt 1):66-69; Jennings T, Rusin M, Tessler H, Cunha-Vaz J. Posterior sub-Tenon's injections of corticosteroids in uveitis patients with cystoid macular edema. *Jpn J Ophthalmol* 1988;32:385-391).

Adverse reactions listed for conventional ophthalmic dexamethasone preparations include: ocular hypertension, glaucoma, posterior subcapsular cataract formation, and secondary ocular infection from pathogens including herpes simplex (Lee et al, 1989 *supra*; Skalka HW, Prchal JT. Effect of corticosteroids on cataract formation. *Arch Ophthalmol* 1980;98:1773-1777; Renfro L, Snow JS. Ocular effects of topical and systemic steroids. *Dermatol Clin* 1992;10(3):505-512; Physician's Desk Reference, 2003). Systemic doses are associated with additional hazardous side-effects including hypertension, hyperglycemias, increased susceptibility to infection, and peptic ulcers (Physician's Desk Reference, 2003).

By delivering a drug directly into the vitreous cavity, blood eye barriers can be circumvented and intraocular therapeutic levels can be achieved with minimal risk of systemic toxicity (Lee et al, 1989 *supra*). This route of administration typically results in a short half-life unless the drug can be delivered using a formulation
5 capable of providing sustained release.

Consequently, a biodegradable implant for delivering a therapeutic agent to an ocular region may provide significant medical benefit for patients afflicted with a medical condition of the eye.

DRAWINGS

10 Figure 1 shows the *in vivo* concentration of dexamethasone in the vitreous of rabbit eyes over a 42 day period after implantation of compressed and extruded biodegradable implants containing 350 µg dexamethasone into the posterior segment of rabbit eyes.

Figure 2 shows the *in vivo* cumulative percentage release of dexamethasone in
15 the vitreous of rabbit eyes over a 42 day period after implantation of compressed and extruded biodegradable implants containing 350 µg dexamethasone and 700 µg dexamethasone into the posterior segment of rabbit eyes.

Figure 3 shows the *in vivo* concentration of dexamethasone in the aqueous humor of rabbit eyes over a 42 day period after implantation of compressed and
20 extruded biodegradable implants containing 350 µg dexamethasone into the posterior segment of rabbit eyes.

Figure 4 shows the *in vivo* concentration of dexamethasone in the plasma (from a rabbit blood sample) over a 42 day period after implantation of compressed and extruded biodegradable implants containing 350 µg dexamethasone into the
25 posterior segment of rabbit eyes.

Figure 5 shows the *in vivo* concentration of dexamethasone in the vitreous of rabbit eyes over a 42 day period after implantation of compressed and extruded biodegradable implants containing 700 µg dexamethasone into the posterior segment of rabbit eyes.

5 Figure 6 shows the *in vivo* concentration of dexamethasone in the aqueous humor of rabbit eyes over a 42 day period after implantation of compressed and extruded biodegradable implants containing 700 µg dexamethasone into the posterior segment of rabbit eyes.

10 Figure 7 shows the *in vivo* concentration of dexamethasone in the plasma (from a rabbit blood sample) over a 42 day period after implantation of compressed and extruded biodegradable implants containing 700 µg dexamethasone into the posterior segment of rabbit eyes.

15 Figure 8 shows the *in vivo* concentration of dexamethasone in the vitreous of rabbit eyes over a 42 day period after implantation of compressed and extruded biodegradable implants containing 350 µg dexamethasone and 700 µg dexamethasone into the posterior segment of rabbit eyes.

20 Figure 9 shows the *in vitro* total cumulative percentage release of dexamethasone into a saline solution at 37°C from 60/40 w/w dexamethasone/PLGA implants having a weight ratio of 40:0 hydrophobic end to hydrophilic end PLGA (312-140-2), weight ratio of 30:10 hydrophobic end to hydrophilic end PLGA (312-140-4), weight ratio of 20:20 hydrophobic end to hydrophilic end PLGA (312-140-3), and weight ratio of 0:40 hydrophobic end to hydrophilic end PLGA (312-140-1).

25 Figure 10 compares the *in vitro* cumulative percentage release of dexamethasone into a saline solution at 37°C for six lots of extruded implants having 60% by weight dexamethasone, 30% by weight hydrophilic end PLGA, and 10% by weight hydrophobic end PLGA.

Figure 11 is a flow chart illustrating manufacturing processes for tablet, single and double extrusion methods for making an ocular implant within the scope of the present invention.

Figure 12 is a graph which shows the cumulative amount of dexamethasone released *in vitro* over time for an ocular implant made by either tableting or a single extrusion processes.

Figure 13 are scanning electromicrographs (SEM) pictures of DEX PS DDS implants made by a tableting process and by a single extrusion process.

Figure 14 shows two graphs of batch to batch vs within batch variability of %LC (% of total dexamethasone) for implants made from either unmilled or milled PLGAs.

Figure 15 is a graph showing in vitro release of dexamethasone from DEX PS DDS implants made by either a single extrusion or by a double extrusion process.

Figure 16 is a flow chart illustrating a double extrusion manufacturing processes for making an ocular implant within the scope of the present invention.

Figure 17 provides a cut-away side view of an applicator to implant an ocular implant within the scope of the present invention.

SUMMARY

Definitions

The following terms as used herein have the following meanings:

“About” means approximately or nearly and in the context of a numerical value or range set forth herein means $\pm 10\%$ of the numerical value or range recited or claimed.

5 “Active agent” and “drug” are used interchangeably and refer to any substance used to treat an ocular condition.

10 “Bioerodible polymer” means a polymer which degrades *in vivo*, and wherein erosion of the polymer over time is required to achieve the active agent release kinetics according to the present invention. Thus, hydrogels such as methylcellulose which act to release drug through polymer swelling are specifically excluded from the term “bioerodible (or biodegradable) polymer”. The words “bioerodible” and “biodegradable” are synonymous and are used interchangeably herein.

15 “Concentration equivalent to dexamethasone”, or “dexamethasone equivalent” means a concentration of an active agent, such as a steroidal anti-inflammatory agent, necessary to have approximately the same efficacy *in vivo* as a particular dose of dexamethasone. For example, hydrocortisone is approximately twenty five fold less potent than dexamethasone, and thus a 25 mg dose of hydrocortisone would be equivalent to a 1 mg dose of dexamethasone. One of ordinary skill in the art would be able to determine the concentration equivalent to dexamethasone for a particular steroidal anti-inflammatory agent from one of several standard tests known in the art.

20 Relative potencies of selected corticosteroids may be found, for example, in Gilman, A.G., *et al.*, eds. (1990). Goodman and Gilman’s: The Pharmacological Basis of Therapeutics. 8th Edition, Pergamon Press: New York, p.1447.

25 “Cumulative release profile” means to the cumulative total percent of an active agent released from an implant into an ocular region or site *in vivo* over time or into a specific release medium *in vitro* over time.

"Glaucoma" means primary, secondary and/or congenital glaucoma. Primary glaucoma can include open angle and closed angle glaucoma. Secondary glaucoma can occur as a complication of a variety of other conditions, such as injury, inflammation, vascular disease and diabetes.

- 5 "Inflammation-mediated" in relation to an ocular condition means any condition of the eye which can benefit from treatment with an anti-inflammatory agent, and is meant to include, but is not limited to, uveitis, macular edema, acute macular degeneration, retinal detachment, ocular tumors, fungal or viral infections, multifocal choroiditis, diabetic uveitis, proliferative vitreoretinopathy (PVR),
10 sympathetic ophthalmia, Vogt Koyanagi-Harada (VKH) syndrome, histoplasmosis, and uveal diffusion.

"Injury" or "damage" are interchangeable and refer to the cellular and morphological manifestations and symptoms resulting from an inflammatory-mediated condition, such as, for example, inflammation.

- 15 "Measured under infinite sink conditions *in vitro*," means assays to measure drug release *in vitro*, wherein the experiment is designed such that the drug concentration in the receptor medium never exceeds 5% of saturation. Examples of suitable assays may be found, for example, in USP 23; NF 18 (1995) pp. 1790-1798.

- "Ocular condition" means a disease, ailment or condition which affects or
20 involves the eye or one or the parts or regions of the eye, such as a retinal disease. The eye includes the eyeball and the tissues and fluids which constitute the eyeball, the periocular muscles (such as the oblique and rectus muscles) and the portion of the optic nerve which is within or adjacent to the eyeball. "Ocular condition" is synonymous with "medical condition of the eye"

"Plurality" means two or more.

"Posterior ocular condition" means a disease, ailment or condition which affects or involves a posterior ocular region or site such as choroid or sclera (in a position posterior to a plane through the posterior wall of the lens capsule), vitreous, vitreous chamber, retina, optic nerve (i.e. the optic disc), and blood vessels and nerve which vascularize or innervate a posterior ocular region or site.

"Steroidal anti-inflammatory agent" and "glucocorticoid" are used interchangeably herein, and are meant to include steroidal agents, compounds or drugs which reduce inflammation when administered at a therapeutically effective level.

"Substantially" in relation to the release profile or the release characteristic of an active agent from a bioerodible implant as in the phrase "substantially continuous rate" of the active agent release rate from the implant means, that the rate of release (i.e. amount of active agent released/unit of time) does not vary by more than 100%, and preferably does not vary by more than 50%, over the period of time selected (i.e. a number of days). "Substantially" in relation to the blending, mixing or dispersing of an active agent in a polymer, as in the phrase "substantially homogeneously dispersed" means that there are no or essentially no particles (i.e. aggregations) of active agent in such a homogenous dispersal.

"Suitable for insertion (or implantation) in (or into) an ocular region or site" with regard to an implant, means an implant which has a size (dimensions) such that it can be inserted or implanted without causing excessive tissue damage and without unduly physically interfering with the existing vision of the patient into which the implant is implanted or inserted.

"Therapeutic levels" or "therapeutic amount" means an amount or a concentration of an active agent that has been locally delivered to an ocular region

that is appropriate to safely treat an ocular condition so as to reduce or prevent a symptom of an ocular condition.

The meaning of abbreviations used herein is explained below:

Term	Meaning
1^{H} -NMR	Proton nuclear magnetic resonance
ABS	Poly acrylonitrile butadiene styrene
ACC	Anterior chamber cell
ALT	Alanine aminotransferase
API	Active pharmaceutical ingredient
AVC	Anterior vitreous cells
BCVA	Best-corrected visual acuity
BI	Boehringer Ingelheim
BRVO	Branch retinal vein occlusion
BSE	Bovine Spongiform Encephalopathy
BVOS	Branch Vein Occlusion Study
B/N	Batch number
°C	Degrees Centigrade
CA	California
CAS	Chemical abstract services
CF	Count fingers
CFU	Colony forming unit
cGMP	Current Good Manufacturing Practice
CI	Confidence interval
CIB	Clinical Investigator's Brochure
CO ₂	Carbon dioxide
COEX	Co-extruded
CRVO	Central retinal vein occlusion
CVOS	Central Vein Occlusion Study
DDS	Drug delivery system
DEX	Dexamethasone
DEX PS DDS	Dexamethasone posterior segment drug delivery system (implant)
DEX PS DDS Applicator	Dexamethasone posterior segment drug delivery

Term	Meaning
system	system (medicinal product)
DME	Diabetic macular oedema
EMA	European medicine evaluation agency
ETDRS	Early Treatment of Diabetic Retinopathy Study
EU	Endotoxin unit
°F	Degrees Fahrenheit
G	Gram
GLP	Good Laboratory Practice
GRB	Geographical BSE (Bovine Spongiform Encephalopathies) risk
H ₂ O	Water
HDPE	High density polyethylene
HPLC	High performance liquid chromatography
IEC	Independent Ethics Committee
IMPD	Investigational medicinal product dossier
INN	International Non-proprietary Name
IOP	Intraocular pressure
IPC	In process control
IR	Infrared
IRB	Institutional Review Board
ISO	International standard organisation
Kg	Kilogram
kGy	Kilo Grey
LAF	Laminar Air Flow
LAL	Limulus Amebocytes Lisat
LC	Label Claim
LOCF	Last observation carried forward
LS	Label strength
ME	Macular oedema
µg	Microgram
Mg	Milligram
µJ	Microjoules
mL	Millilitre(s)
Mm	Millimetre(s)

Term	Meaning
mmHg	Millimeters of mercury
mol	Mole
n or N	Number
n/a	Not applicable
ND	Not detected
Ng	Nanogram(s)
NSAID	Nonsteroidal anti-inflammatory drug
NT	Not tested
OCT	Optical Coherence Tomography
PDE	Permitted daily exposure
PET	Polyethylene terephthalate
pH	Hydrogen potential
Ph. Eur.	European Pharmacopoeia
PK	Pharmacokinetics
pKa	Acid dissociation constant
PLGA, PLG	Poly (D,L-lactide-co-glycolide).
PME	Persistent macular edema
ppm	Part per million
PS	Posterior segment
PVR	Proliferative vitreoretinopathy
RH	Relative humidity
SAE	Serious adverse event
SD	Standard deviation
SEM	Scanning electron microscope
TSE	Transmissible spongiform encephalopathy
USA	United States of America
USP	United States Pharmacopoeia
UV	Ultra violet
VEGF	Vascular endothelial growth factor
WPE	Ultrahigh molecular weight polyethylene

Our invention encompasses a bioerodible implant for treating a medical condition of the eye comprising an active agent dispersed within a biodegradable polymer matrix, wherein at least about 75% of the particles of the active agent have a

diameter of less than about $10\mu\text{m}$. Preferably, at least about 99% of the particles have a diameter of less than about $20\mu\text{m}$.

The active agent can be selected from the group consisting of ace-inhibitors, endogenous cytokines, agents that influence basement membrane, agents that
5 influence the growth of endothelial cells, adrenergic agonists or blockers, cholinergic agonists or blockers, aldose reductase inhibitors, analgesics, anesthetics, antiallergics, anti-inflammatory agents, steroids (such as a steroidal anti-inflammatory agent), antihypertensives, pressors, antibacterials, antivirals, antifungals, antiprotozoals, anti-infective agents, antitumor agents, antimetabolites, and antiangiogenic agents. Thus,
10 the active agent can be cortisone, dexamethasone, fluocinolone, hydrocortisone, methylprednisolone, prednisolone, prednisone, triamcinolone, and any derivative thereof.

The bioerodible implant is sized for implantation in an ocular region. The ocular region can be any one or more of the anterior chamber, the posterior chamber,
15 the vitreous cavity, the choroid, the suprachoroidal space, the conjunctiva, the subconjunctival space, the episcleral space, the intracorneal space, the epicorneal space, the sclera, the pars plana, surgically-induced avascular regions, the macula, and the retina.

An alternate embodiment of the bioerodible implant can comprise a steroid
20 active agent dispersed within a biodegradable polymer matrix, wherein at least about 75% of the particles of the active agent have a diameter of less than about $20\mu\text{m}$.

Our present invention also encompasses a method for making a bioerodible implant for treating a medical condition of the eye, the method comprising a plurality of extrusions of a biodegradable polymer. This method can also comprise the step of
25 milling the biodegradable polymer prior to the extrusion. The biodegradable polymer can be a poly(lactic-co-glycolic)acid (PLGA) copolymer. The ratio of lactic to glycolic acid monomers in the polymer can be about 50/50 weight percentage.

Additionally, the PLGA copolymer can be about 20 to about 90 weight percent of the bioerodible implant. Alternately, the PLGA copolymer can be about 40 percent by weight of the bioerodible implant.

5 A detailed method for making a bioerodible implant for treating a medical condition of the eye can have the steps of: (a) milling a biodegradable polymer; (b) blending the milled biodegradable polymer and particles of an active agent, to thereby obtain a blended mixture of the milled biodegradable polymer and the particles of the active agent, wherein at least about 75% of the particles of the active agent have a diameter of less than about 20 μ m; (c) carrying out a first extrusion of
10 the blended mixture, to thereby obtain a first extrusion product; (d) pelletizing the first extrusion product, and; (e) carrying out a second extrusion of the pelletized first extrusion product, thereby obtaining a bioerodible implant for treating a medical condition of the eye. Our invention also includes a bioerodible implant for treating a medical condition of the eye made by this detailed method.

15 DESCRIPTION

The present invention provides biodegradable ocular implants and methods for treating medical conditions of the eye. Usually, the implants are formed to be monolithic, i.e., the particles of active agent are distributed throughout the biodegradable polymer matrix. Furthermore, the implants are formed to release an
20 active agent into an ocular region of the eye over various time periods. The active agent may be release over a time period including, but is not limited to, approximately six months, approximately three months, approximately one month, or less than one month.

Biodegradable Implants For Treating Medical Conditions of the Eye

25 The implants of the invention include an active agent dispersed within a biodegradable polymer. The implant compositions typically vary according to the preferred drug release profile, the particular active agent used, the condition being

treated, and the medical history of the patient. Active agents that may be used include, but are not limited to, ace-inhibitors, endogenous cytokines, agents that influence basement membrane, agents that influence the growth of endothelial cells, adrenergic agonists or blockers, cholinergic agonists or blockers, aldose reductase
 5 inhibitors, analgesics, anesthetics, antiallergics, anti-inflammatory agents, antihypertensives, pressors, antibacterials, antivirals, antifungals, antiprotozoals, anti-infectives, antitumor agents, antimetabolites, and antiangiogenic agents.

In one variation the active agent is methotrexate. In another variation, the active agent is retinoic acid. In a preferred variation, the anti-inflammatory agent is a
 10 nonsteroidal anti-inflammatory agent. Nonsteroidal anti-inflammatory agents that may be used include, but are not limited to, aspirin, diclofenac, flurbiprofen, ibuprofen, ketorolac, naproxen, and suprofen. In a more preferred variation, the anti-inflammatory agent is a steroidal anti-inflammatory agent.

Steroidal Anti-Inflammatory Agents

15 The steroidal anti-inflammatory agents that may be used in the ocular implants include, but are not limited to, 21-acetoxypregnenolone, alclometasone, algestone, amcinonide, beclomethasone, betamethasone, budesonide, chloroprednisone, clobetasol, clobetasone, clocortolone, cloprednol, corticosterone, cortisone, cortivazol, deflazacort, desonide, desoximetasone, dexamethasone,
 20 diflorasone, diflucortolone, difluprednate, enoxolone, fluazacort, flucloronide, flumethasone, flunisolide, fluocinolone acetonide, fluocinonide, fluocortin butyl, fluocortolone, fluorometholone, fluperolone acetate, fluprednidene acetate, fluprednisolone, flurandrenolide, fluticasone propionate, formocortal, halcinonide, halobetasol propionate, halometasone, halopredone acetate, hydrocortamate,
 25 hydrocortisone, loteprednol etabonate, mazipredone, medrysone, meprednisone, methylprednisolone, mometasone furoate, paramethasone, prednicarbate, prednisolone, prednisolone 25-diethylamino-acetate, prednisolone sodium phosphate, prednisone, prednival, prednylidene, rimexolone, tixocortol, triamcinolone,

triamcinolone acetonide, triamcinolone benetonide, triamcinolone hexacetonide, and any of their derivatives.

In one variation, cortisone, dexamethasone, fluocinolone, hydrocortisone, methylprednisolone, prednisolone, prednisone, and triamcinolone, and their
5 derivatives, are preferred steroidal anti-inflammatory agents. In another preferred variation, the steroidal anti-inflammatory agent is dexamethasone. In another variation, the biodegradable implant includes a combination of two or more steroidal anti-inflammatory agents.

The steroidal anti-inflammatory agent may constitute from about 10% to
10 about 90% by weight of the implant. In one variation, the agent is from about 40% to about 80% by weight of the implant. In a preferred variation, the agent comprises about 60% by weight of the implant.

The Biodegradable Polymer Matrix

In one variation, the active agent may be homogeneously dispersed in the
15 biodegradable polymer matrix of the implants. The selection of the biodegradable polymer matrix to be employed will vary with the desired release kinetics, patient tolerance, the nature of the disease to be treated, and the like. Polymer characteristics that are considered include, but are not limited to, the biocompatibility and biodegradability at the site of implantation, compatibility with the active agent of
20 interest, and processing temperatures. The biodegradable polymer matrix usually comprises at least about 10, at least about 20, at least about 30, at least about 40, at least about 50, at least about 60, at least about 70, at least about 80, or at least about 90 weight percent of the implant. In one variation, the biodegradable polymer matrix comprises about 40% by weight of the implant.

25 Biodegradable polymer matrices which may be employed include, but are not limited to, polymers made of monomers such as organic esters or ethers, which when degraded result in physiologically acceptable degradation products. Anhydrides,

amides, orthoesters, or the like, by themselves or in combination with other monomers, may also be used. The polymers are generally condensation polymers. The polymers may be crosslinked or non-crosslinked. If crosslinked, they are usually not more than lightly crosslinked, and are less than 5% crosslinked, usually less than 1% crosslinked.

For the most part, besides carbon and hydrogen, the polymers will include oxygen and nitrogen, particularly oxygen. The oxygen may be present as oxy, e.g., hydroxy or ether, carbonyl, e.g., non-oxo-carbonyl, such as carboxylic acid ester, and the like. The nitrogen may be present as amide, cyano, and amino. An exemplary list of biodegradable polymers that may be used are described in Heller, Biodegradable Polymers in Controlled Drug Delivery, In: "CRC Critical Reviews in Therapeutic Drug Carrier Systems", Vol. 1. CRC Press, Boca Raton, FL (1987).

Of particular interest are polymers of hydroxyaliphatic carboxylic acids, either homo- or copolymers, and polysaccharides. Included among the polyesters of interest are homo- or copolymers of D-lactic acid, L-lactic acid, racemic lactic acid, glycolic acid, caprolactone, and combinations thereof. Copolymers of glycolic and lactic acid are of particular interest, where the rate of biodegradation is controlled by the ratio of glycolic to lactic acid. The percent of each monomer in poly(lactic-co-glycolic)acid (PLGA) copolymer may be 0-100%, about 15-85%, about 25-75%, or about 35-65%. In a preferred variation, a 50/50 PLGA copolymer is used. More preferably, a random copolymer of 50/50 PLGA is used.

Biodegradable polymer matrices that include mixtures of hydrophilic and hydrophobic ended PLGA may also be employed, and are useful in modulating polymer matrix degradation rates. Hydrophobic ended (also referred to as capped or end-capped) PLGA has an ester linkage hydrophobic in nature at the polymer terminus. Typical hydrophobic end groups include, but are not limited to alkyl esters and aromatic esters. Hydrophilic ended (also referred to as uncapped) PLGA has an end group hydrophilic in nature at the polymer terminus. PLGA with a hydrophilic

end groups at the polymer terminus degrades faster than hydrophobic ended PLGA because it takes up water and undergoes hydrolysis at a faster rate (Tracy et al., *Biomaterials* 20:1057-1062 (1999)). Examples of suitable hydrophilic end groups that may be incorporated to enhance hydrolysis include, but are not limited to, carboxyl, hydroxyl, and polyethylene glycol. The specific end group will typically result from the initiator employed in the polymerization process. For example, if the initiator is water or carboxylic acid, the resulting end groups will be carboxyl and hydroxyl. Similarly, if the initiator is a monofunctional alcohol, the resulting end groups will be ester or hydroxyl.

10 The implants may be formed from all hydrophilic end PLGA or all hydrophobic end PLGA. In general, however, the ratio of hydrophilic end to hydrophobic end PLGA in the biodegradable polymer matrices of this invention range from about 10:1 to about 1:10 by weight. For example, the ratio may be 3:1, 2:1, or 1:1 by weight. In a preferred variation, an implant with a ratio of hydrophilic
15 end to hydrophobic end PLGA of 3:1 w/w is used.

Additional Agents

Other agents may be employed in the formulation for a variety of purposes. For example, buffering agents and preservatives may be employed. Preservatives which may be used include, but are not limited to, sodium bisulfite, sodium bisulfate, sodium thiosulfate, benzalkonium chloride, chlorobutanol, thimerosal, phenylmercuric acetate, phenylmercuric nitrate, methylparaben, polyvinyl alcohol and phenylethyl alcohol. Examples of buffering agents that may be employed include, but are not limited to, sodium carbonate, sodium borate, sodium phosphate, sodium acetate, sodium bicarbonate, and the like, as approved by the FDA for the
20 desired route of administration. Electrolytes such as sodium chloride and potassium chloride may also be included in the formulation.

The biodegradable ocular implants may also include additional hydrophilic or hydrophobic compounds that accelerate or retard release of the active agent.

Furthermore, the inventors believe that because hydrophilic end PLGA has a higher degradation rate than hydrophobic end PLGA due to its ability to take up water more readily, increasing the amount of hydrophilic end PLGA in the implant polymer matrix will result in faster dissolution rates. Figure 9 shows that the time from
5 implantation to significant release of active agent (lag time) increases with decreasing amounts of hydrophilic end PLGA in the ocular implant. In Figure 9, the lag time for implants having 0% hydrophilic end PLGA (40% w/w hydrophobic end) was shown to be about 21 days. In comparison, a significant reduction in lag time was seen with implants having 10% w/w and 20% w/w hydrophilic end PLGA.

10 Release Kinetics

The inventors believe the implants of the invention are formulated with particles of an active agent dispersed within a biodegradable polymer matrix. Without being bound by theory, the inventors believe that release of the active agent is achieved by erosion of the biodegradable polymer matrix and by diffusion of the
15 particulate agent into an ocular fluid, e.g., the vitreous, with subsequent dissolution of the polymer matrix and release of the active agent. The inventors believe that the factors that influence the release kinetics include such characteristics as the size of the active agent particles, the solubility of the active agent, the ratio of active agent to polymer(s), the method of manufacture, the surface area exposed, and the erosion rate
20 of the polymer(s). The release kinetics achieved by this form of active agent release are different than that achieved through formulations which release active agents through polymer swelling, such as with crosslinked hydrogels. In that case, the active agent is not released through polymer erosion, but through polymer swelling, which releases agent as liquid diffuses through the pathways exposed.

25 The inventors believe that the release rate of the active agent depends at least in part on the rate of degradation of the polymer backbone component or components making up the biodegradable polymer matrix. For example, condensation polymers may be degraded by hydrolysis (among other mechanisms) and therefore any change in the composition of the implant that enhances water uptake by the implant will

likely increase the rate of hydrolysis, thereby increasing the rate of polymer degradation and erosion, and thus increasing the rate of active agent release.

The release kinetics of the implants of the invention are dependent in part on the surface area of the implants. A larger surface area exposes more polymer and active agent to ocular fluid, causing faster erosion of the polymer matrix and dissolution of the active agent particles in the fluid. The size and shape of the implant may also be used to control the rate of release, period of treatment, and active agent concentration at the site of implantation. At equal active agent loads, larger implants will deliver a proportionately larger dose, but depending on the surface to mass ratio, may possess a slower release rate. For implantation in an ocular region, the total weight of the implant preferably ranges, e.g., from about 100-5000 μg , usually from about 500-1500 μg . In one variation, the total weight of the implant is about 600 μg . In another variation, the total weight of the implant is about 1200 μg .

The bioerodible implants are typically solid, and may be formed as particles, sheets, patches, plaques, films, discs, fibers, rods, and the like, or may be of any size or shape compatible with the selected site of implantation, as long as the implants have the desired release kinetics and deliver an amount of active agent that is therapeutic for the intended medical condition of the eye. The upper limit for the implant size will be determined by factors such as the desired release kinetics, toleration for the implant at the site of implantation, size limitations on insertion, and ease of handling. For example, the vitreous chamber is able to accommodate relatively large rod-shaped implants, generally having diameters of about 0.05 mm to 3 mm and a length of about 0.5 to about 10 mm. In one variation, the rods have diameters of about 0.1 mm to about 1 mm. In another variation, the rods have diameters of about 0.3 mm to about 0.75 mm. In yet a further variation, other implants having variable geometries but approximately similar volumes may also be used.

As previously discussed, the release of an active agent from a biodegradable polymer matrix may also be modulated by varying the ratio of hydrophilic end PLGA to hydrophobic end PLGA in the matrix. Release rates may be further manipulated by the method used to manufacture the implant. For instance, as illustrated in

5 Examples 4-7, extruded 60/40 w/w dexamethasone/PLGA implants having a ratio of hydrophilic end and hydrophobic end PLGA of 3:1, compared to compressed tablet implants, demonstrate a different drug release profile and concentration of agent in the vitreous over about a one month period. Overall, a lower burst of agent release and a more consistent level of agent in the vitreous is demonstrated with the extruded

10 implants.

As shown in Figure 2 and Examples 4 and 5, a higher initial burst of active agent release occurs on day one after implantation with the 350 µg dexamethasone compressed tablet implant (350T) in comparison to the 350 µg dexamethasone extruded implant (350E). A higher initial burst of active agent release also occurs

15 with the 700 µg dexamethasone compressed implant (700T) in comparison to the 700 µg dexamethasone extruded implant (700E) on day 1, as shown in Figure 2 and Examples 6 and 7.

The proportions of active agent, biodegradable polymer matrix, and any other additives may be empirically determined by formulating several implants with

20 varying proportions and determining the release profile *in vitro* or *in vivo*. A USP approved method for dissolution or release test can be used to measure the rate of release *in vitro* (USP 24; NF 19 (2000) pp. 1941-1951). For example, a weighed sample of the implant is added to a measured volume of a solution containing 0.9% NaCl in water, where the solution volume will be such that the active agent

25 concentration after release is less than 20% of saturation. The mixture is maintained at 37°C and stirred or shaken slowly to maintain the implants in suspension. The release of the dissolved active agent as a function of time may then be followed by various methods known in the art, such as spectrophotometrically, HPLC, mass

spectroscopy, and the like, until the solution concentration becomes constant or until greater than 90% of the active agent has been released.

In one variation, the extruded implants described herewith (ratio of hydrophilic end PLGA to hydrophobic end PLGA of 3:1) may have *in vivo* cumulative percentage release profiles with the following described characteristics, as shown in Figure 2, where the release profiles are for release of the active agent *in vivo* after implantation of the implants into the vitreous of rabbit eyes. The volume of rabbit eyes is approximately 60-70% of human eyes.

At day one after implantation, the percentage *in vivo* cumulative release may be between about 0% and about 15%, and more usually between about 0% and about 10%. At day one after implantation, the percentage *in vivo* cumulative release may be less than about 15%, and more usually less than about 10%.

At day three after implantation, the percentage *in vivo* cumulative release may be between about 0% and about 20%, and more usually between about 5% and about 15%. At day three after implantation, the percentage *in vivo* cumulative release may be less than about 20%, and more usually less than about 15%.

At day seven after implantation, the percentage *in vivo* cumulative release may be between about 0% and about 35%, more usually between about 5% and about 30%, and more usually still between about 10% and about 25%. At day seven after implantation, the percentage *in vivo* cumulative release may be greater than about 2%, more usually greater than about 5%, and more usually still greater than about 10%.

At day fourteen after implantation, the percentage *in vivo* cumulative release may be between about 20% and about 60%, more usually between about 25% and about 55%, and more usually still between about 30% and about 50%. At day fourteen after implantation, the percentage *in vivo* cumulative release may be greater

than about 20%, more usually greater than about 25%, and more usually still greater than about 30%.

At day twenty-one after implantation, the percentage *in vivo* cumulative release may be between about 55% and about 95%, more usually between about 60% and about 90%, and more usually still between about 65% and about 85%. At day twenty-one after implantation, the percentage *in vivo* cumulative release may be greater than about 55% , more usually greater than about 60% , and more usually still greater than about 65%.

At day twenty-eight after implantation, the percentage *in vivo* cumulative release may be between about 80% and about 100%, more usually between about 85% and about 100%, and more usually still between about 90% and about 100%. At day twenty-eight after implantation, the percentage *in vivo* cumulative release may be greater than about 80%, more usually greater than about 85%, and more usually still greater than about 90%.

At day thirty-five after implantation, the percentage *in vivo* cumulative release may be between about 95% and about 100%, and more usually between about 97% and about 100%. At day thirty-five after implantation, the percentage *in vivo* cumulative release may be greater than about 95%, and more usually greater than about 97%.

In one variation, the percentage *in vivo* cumulative release has the following characteristics: one day after implantation it is less than about 15%; three days after implantation it is less than about 20%; seven days after implantation it is greater than about 5%; fourteen days after implantation it is greater than about 25%; twenty-one days after implantation it is greater than about 60%; and twenty-eight days after implantation it is greater than about 80%. In another variation, the percentage *in vivo* cumulative release has the following characteristics: one day after implantation it is less than about 10%; three days after implantation it is less than about 15%;

seven days after implantation it is greater than about 10%; fourteen days after implantation it is greater than about 30%; twenty-one days after implantation it is greater than about 65%; twenty-eight days after implantation it is greater than about 85%.

- 5 In yet another variation, the extruded implants described in this patent may have *in vitro* cumulative percentage release profiles in saline solution at 37°C with the following characteristics, as further described below, and as shown in Figure 10.

- The percentage *in vitro* cumulative release at day one may be between about 0% and about 5%, and more usually between about 0% and about 3%. The
10 percentage *in vitro* cumulative release at day one may be less than about 5%, and more usually less than about 3%.

- The percentage *in vitro* cumulative release at day four may be between about 0% and about 7%, and more usually between about 0% and about 5%. The
15 percentage *in vitro* cumulative release at day four may be less than about 7%, and more usually less than about 5%.

 The percentage *in vitro* cumulative release at day seven may be between about 1% and about 10%, and more usually between about 2% and about 8%. The percentage *in vitro* cumulative release at day seven may be greater than about 1%, and more usually greater than about 2%.

- 20 The percentage *in vitro* cumulative release at day 14 may be between about 25% and about 65%, more usually between about 30% and about 60%, and more usually still between about 35% and about 55%. The percentage *in vitro* cumulative release at day 14 may be greater than about 25%, more usually greater than about 30%, and more usually still greater than about 35%.

The percentage *in vitro* cumulative release at day 21 may be between about 60% and about 100%, more usually between about 65% and about 95%, and more usually still between about 70% and about 90%. The percentage *in vitro* cumulative release at day 21 may be greater than about 60%, more usually greater than about 65%, and more usually still greater than about 70%.

The percentage *in vitro* cumulative release at day 28 may be between about 75% and about 100%, more usually between about 80% and about 100%, and more usually still between about 85% and about 95%. The percentage *in vitro* cumulative release at day 28 may be greater than about 75%, more usually greater than about 80%, and more usually still greater than about 85%.

The percentage *in vitro* cumulative release at day 35 may be between about 85% and about 100%, more usually between about 90% and about 100%, and more usually still between about 95% and about 100%. The percentage *in vitro* cumulative release at day 35 may be greater than about 85%, more usually greater than about 90%, and more usually still greater than about 95%.

In one variation, the percentage *in vitro* cumulative release has the following characteristics: after one day it is less than about 1%; after four days it is less than about 7%; after seven days it is greater than about 2%; after 14 days it is greater than about 30%; after 21 days it is greater than about 65%; after 28 days it is greater than about 80%; and after 35 days it is greater than about 90%. In another variation, the percentage *in vitro* cumulative release has the following characteristics: after one day it is less than about 3%; after four days it is less than about 5%; after seven days it is greater than about 2%; after 14 days it is greater than about 35%; after 21 days it is greater than about 70%; after 28 days it is greater than about 85%; and after 35 days it is greater than about 90%.

Besides showing a lower burst effect for the extruded implants, Figures 2 and 10 also demonstrate that after 28 days *in vivo* in rabbit eyes, or *in vitro* in a saline

solution at 37°C, respectively, almost all of the active agent has been released from the implants. Furthermore, Figures 2 and 10 show that the active agent release profiles for the extruded implants *in vivo* (from the time of implantation) and *in vitro* (from the time of placement into a saline solution at 37°C) are substantially similar and follow approximately a sigmoidal curve, releasing substantially all of the active agent over 28 days. From day one to approximately day 17, the curves show approximately an upward curvature (i.e., the derivative of the curve increases as time increases), and from approximately day 17 onwards the curves show approximately a downward curvature (i.e., the derivative of the curve decreases as time increases).

10 In contrast, the plots shown in Figure 2 for the 350 µg and 700 µg dexamethasone compressed tablet implants exhibit a higher initial burst of agent release generally followed by a gradual increase in release. Furthermore, as shown in Figures 1 and 5, implantation of a compressed implant results in different concentrations of active agent in the vitreous at various time points from implants that have been extruded. For example, as shown in Figures 1 and 5, with extruded implants there is a gradual increase, plateau, and gradual decrease in intravitreal agent concentrations. In contrast, for compressed tablet implants, there is a higher initial active agent release followed by an approximately constant decrease over time. Consequently, the intravitreal concentration curve for extruded implants results in more sustained levels of active agent in the ocular region.

In addition to the previously described implants releasing substantially all of the therapeutic agent within 35 days, by varying implant components including, but not limited to, the composition of the biodegradable polymer matrix, implants may also be formulated to release a therapeutic agent for any desirable duration of time, for example, for about one week, for about two weeks, for about three weeks, for about four weeks, for about five weeks, for about six weeks, for about seven weeks, for about eight weeks, for about nine weeks, for about ten weeks, for about eleven weeks, for about twelve weeks, or for more than 12 weeks.

Another important feature of the extruded implants is that different concentration levels of active agent may be established in the vitreous using different doses of the active agent. As illustrated in Figure 8, the concentration of agent in the vitreous is significantly larger with the 700 μ g dexamethasone extruded implant than
5 with the 350 μ g dexamethasone extruded implant. Different active agent concentrations are not demonstrated with the compressed tablet implant. Thus, by using an extruded implant, it is possible to more easily control the concentration of active agent in the vitreous. In particular, specific dose-response relationships may be established since the implants can be sized to deliver a predetermined amount of
10 active agent.

Applications

Examples of medical conditions of the eye which may be treated by the implants and methods of the invention include, but are not limited to, uveitis, macular edema, macular degeneration, retinal detachment, ocular tumors, fungal or viral
15 infections, multifocal choroiditis, diabetic retinopathy, proliferative vitreoretinopathy (PVR), sympathetic ophthalmia, Vogt Koyanagi-Harada (VKH) syndrome, histoplasmosis, uveal diffusion, and vascular occlusion. In one variation, the implants are particularly useful in treating such medical conditions as uveitis, macular edema, vascular occlusive conditions, proliferative vitreoretinopathy (PVR), and
20 various other retinopathies.

Method of Implantation

The biodegradable implants may be inserted into the eye by a variety of methods, including placement by forceps, by trocar, or by other types of applicators, after making an incision in the sclera. In some instances, a trocar or applicator may
25 be used without creating an incision. In a preferred variation, a hand held applicator is used to insert one or more biodegradable implants into the eye. The hand held applicator typically comprises an 18-30 GA stainless steel needle, a lever, an actuator, and a plunger.

The method of implantation generally first involves accessing the target area within the ocular region with the needle. Once within the target area, e.g., the vitreous cavity, the lever on the hand held device is depressed to cause the actuator to drive the plunger forward. As the plunger moves forward, it pushes the implant into
5 the target area.

Extrusion Methods

The use of extrusion methods allows for large-scale manufacture of implants and results in implants with a homogeneous dispersion of the drug within the polymer matrix. When using extrusion methods, the polymers and active agents that are
10 chosen are stable at temperatures required for manufacturing, usually at least about 50°C. Extrusion methods use temperatures of about 25°C to about 150°C, more preferably about 60°C to about 130°C.

Different extrusion methods may yield implants with different characteristics, including but not limited to the homogeneity of the dispersion of the active agent
15 within the polymer matrix. For example, using a piston extruder, a single screw extruder, and a twin screw extruder will generally produce implants with progressively more homogeneous dispersion of the active. When using one extrusion method, extrusion parameters such as temperature, extrusion speed, die geometry, and die surface finish will have an effect on the release profile of the implants
20 produced.

In one variation of producing implants by extrusion methods, the drug and polymer are first mixed at room temperature and then heated to a temperature range of about 60°C to about 150°C, more usually to about 130°C for a time period of about 0 to about 1 hour, more usually from about 0 to about 30 minutes, more usually
25 still from about 5 minutes to about 15 minutes, and most usually for about 10 minutes. The implants are then extruded at a temperature of between about 60°C to about 130°C, preferably at a temperature of between about 75°C and 110°C, and more preferably at a temperature of about 90°C.

In a preferred extrusion method, the powder blend of active agent and PLGA is added to a single or twin screw extruder preset at a temperature of about 80°C to about 130°C, and directly extruded as a filament or rod with minimal residence time in the extruder. The extruded filament or rod is then cut into small implants having the loading dose of active agent appropriate to treat the medical condition of its intended use.

DEX PS DDS

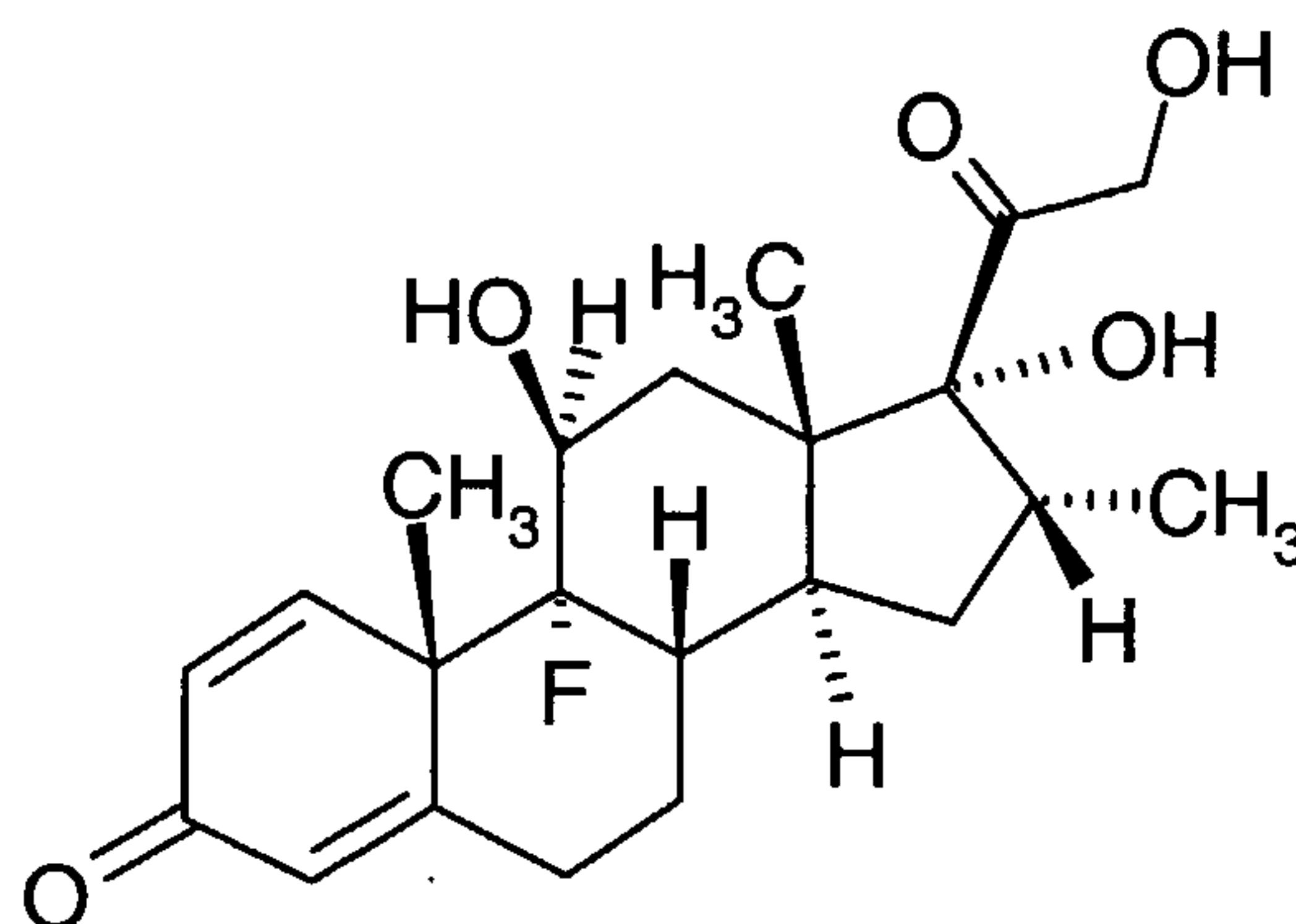
The present invention is based upon the discovery of an intraocular drug delivery system which can address many of the problems associated with conventional therapies for the treatment of ocular conditions, such as posterior segment inflammation, including fluctuating drug levels, short intraocular half-life, and prolonged systemic exposure to high levels of corticosteroids. The intraocular drug delivery system of the present invention encompasses use of dexamethasone as the active pharmaceutical agent, in which case the intraocular drug delivery system of the present invention can be referred to as a Dexamethasone Posterior Segment Drug Delivery System (DEX PS DDS). The DEX PS DDS is intended for placement into the posterior segment by a pars plana injection, a familiar method of administration for ophthalmologists. The DEX PS DDS can be comprised of a biodegradable copolymer, poly(lactic glycolic) acid (PLGA), containing micronised dexamethasone. The DEX PS DDS can release dexamethasone, providing a total dose of approximately 350 or 700 µg over approximately 35 days. In comparison, other routes of administration (topical, periocular, systemic and standard intravitreal injections) require much higher daily doses to deliver equivalent levels of dexamethasone to the posterior segment while also exposing non-target organs to corticosteroids. Topical administration of 2 drops of dexamethasone ophthalmic suspension 0.1% four times daily to both eyes is equivalent to almost 500 µg per day. Systemic doses may be as high as 1,000 µg/kg/day (Pinar V. Intermediate uveitis. Massachusetts Eye & Ear Infirmary Immunology Service 1998; Weisbecker CA, Fraunfelder FT, Naidoff M, Tipperman R, eds. 1999 Physicians' Desk Reference for Ophthalmology, 27th ed. Montvale, NJ: Medical

Economics Company, 1998; 7-8, 278-279). With the DEX PS DDS, substantially lower daily doses of dexamethasone can be administered directly to the posterior segment compared to the doses needed with conventional topical, systemic, or intravitreal therapies, thereby minimizing potential side effects. While releasing
5 dexamethasone, the polymer can gradually degrade completely over time so there is no need to remove the DEX PS DDS after its placement into the posterior segment of a patient eye.

To facilitate delivery of DEX PS DDS into the posterior segment of the eye, an applicator has been designed to deliver the DEX PS DDS directly into the
10 vitreous. The DDS Applicator allows placement of the DEX PS DDS into the posterior segment through a small hollow gauge needle, thereby decreasing the morbidity associated with surgery and pars plana injection at vitrectomy. The extruded DEX PS DDS is placed in the Applicator during the manufacturing of the sterile finished drug product. The DEX PS DDS Applicator System can be a single-
15 use only device.

700 µg and 350 µg Dexamethasone Posterior Segment Drug Delivery System (DEX PS DDS Applicator System) can be used in the treatment of, for example, patients with macular oedema following central retinal vein occlusion or branch retinal vein occlusion.

20 Dexamethasone can be obtained from Aventis Pharma, Montvale, New Jersey, U.S.A. The chemical name of dexamethasone is pregna-1,4-diene-3,20-dione-9-fluoro-11,17,21-trihydroxy-16-methyl-, (11 $\square$ 16 $\square$), and its chemical structure can be represented diagrammatically as follows:



Other characteristics of dexamethasone are:

	Molecular Formula:	$C_{22}H_{29}FO_5$
5	Molecular Weight:	392.47
	Chirality / Stereochemistry:	Dexamethasone has 8 chiral centres and is optically active
	Description:	White or almost white, crystalline powder
	pH and pKa:	Dexamethasone has no ionisable groups
10	Melting Point:	253°C to 255°C
	Solubility:	Water: practically insoluble Ethanol: Sparingly soluble Methylene chloride: slightly soluble

Further information on the physical and chemical properties of
 15 dexamethasone is summarised in the current European Pharmacopoeia (Ph. Eur.).

An embodiment of our invention can be referred to as a DEX PS DDS. DEX PS DDS is an implant (a drug delivery system or DDS) for intravitreal (i.e. posterior segment, or PS) use, comprised of dexamethasone (i.e. DEX) (drug substance) and a

polymer matrix of 50:50 poly (D,L-lactide-co-glycolide) PLGA, constituted of two grades of PLGA (50:50 PLGA ester and 50:50 PLGA acid). See Table 1 for details. This biodegradable drug delivery system is designed to release the drug substance into the posterior segment of the eye over a 35-day period. DEX PS DDS can be
 5 implanted into the vitreous humour of the eye using an applicator system..

Two dose levels, one containing 350 μ g and one containing 700 μ g of dexamethasone, have been evaluated in a clinical trial. Both dose levels have the same formulation as detailed in Table 2. They are prepared using the same bulk and double extrusion process, but cut to different lengths to obtain the appropriate dosage
 10 strength.

Table 1: Qualitative composition of a sample DEX PS DDS

Component	Quality Standard	Function
Dexamethasone	Ph. Eur.	Active ingredient
50:50 PLGA ester	Allergan, Inc.	Biodegradable extended release polymer matrix
50:50 PLGA acid	Allergan, Inc.	Biodegradable extended release polymer matrix

Table 2: Quantitative Composition of a sample DEX PS DDS (manufacturing batch formula)

Component	Formula number	350 μ g	700 μ g	Representative 80 g Batch Quantity
		9635X	9632X	
Dexamethasone		350 μ g (60%)	700 μ g (60%)	48 grams
50:50 PLGA ester (hydrophobic)		58 μ g (10%)	116 μ g (10%)	8 grams
50:50 PLGA acid (hydrophilic)		175 μ g (30%)	350 μ g (30%)	24 grams

15

The drug substance used in the DEX PS DDS is dexamethasone micronised.

DEX PS DDS can contain two excipients (i.e. non-active ingredients) which can be present as two different grades of the same biodegradable polymer 50:50 Poly (D,L lactide-co-glycolide) (PLGA), which can be supplied by Boehringer Ingelheim: 50:50 PLGA ester and 50:50 PLGA acid.

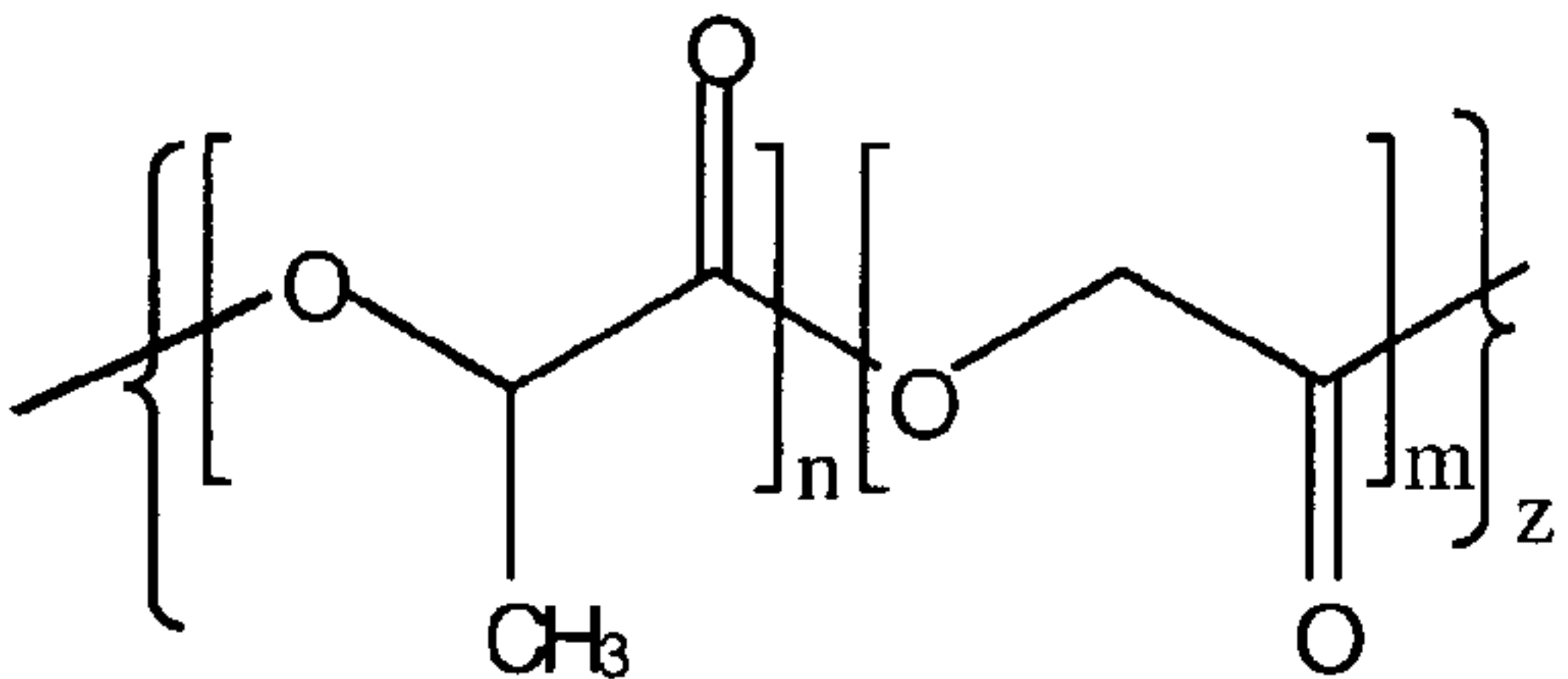
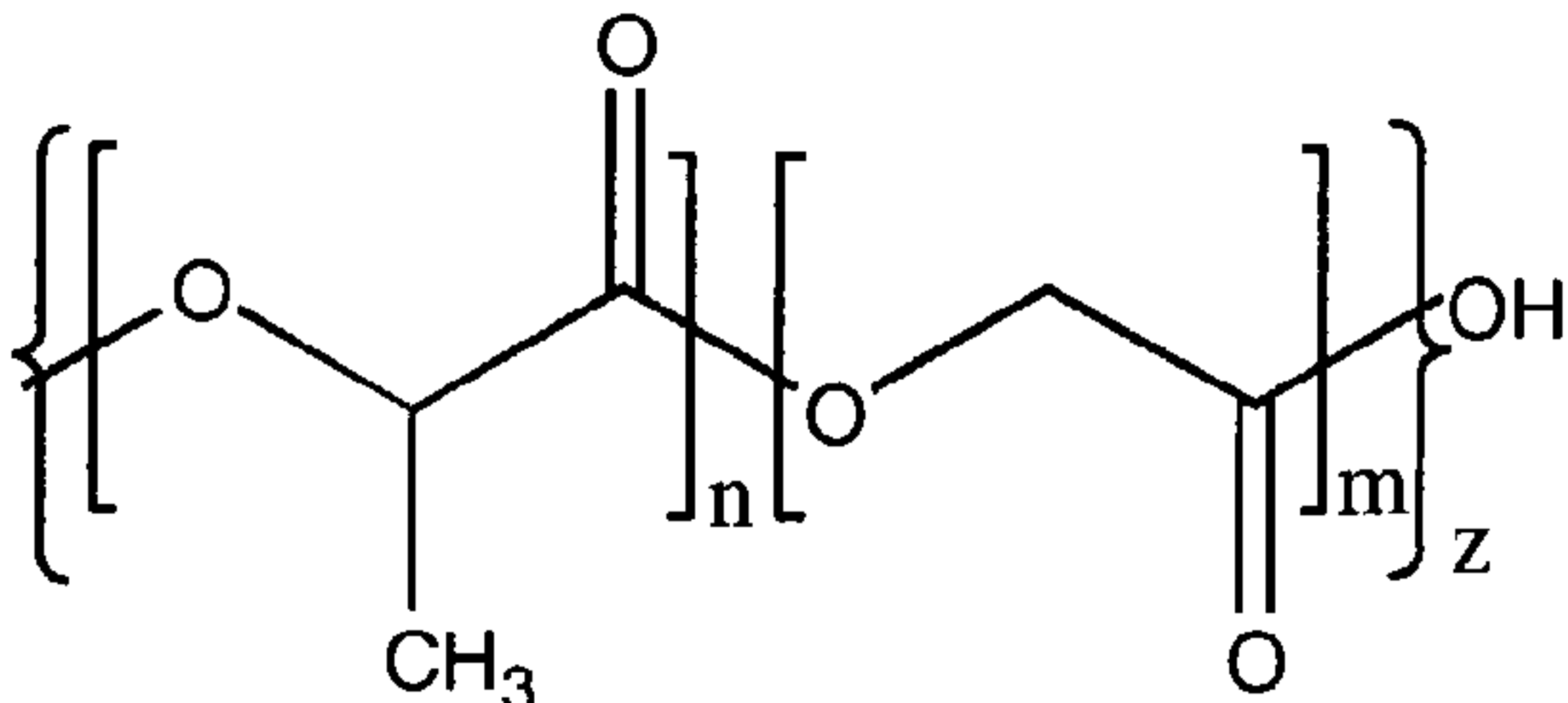
- 5 Poly D,L lactide-co-glycolide has been used for more than 15 years in parenteral products and is a main component of absorbable sutures. A list of some of the medical products commercially available is supplied in Table 3.

Table 3 List of commercial medical products containing PLGA

Name	Manufacturer	Drug Substance	Dosage form	Mode of administration
Vicryl®	Ethicon	Suture used in ocular surgery		
Enantone®	Tadeca	Leuprorelin	Microsphere suspension	Injection (SC or IM)
Prostap®	Wyeth	Leuprorelin acetate	Microsphere suspension	Injection (SC or IM)
Bigonist®	Aventis	Buserelin	Implant	Injection (SC)
Somatuline®	Beaufour Ipsen Pharma	Lanreotide acetate	Microparticle suspension	Injection (IM)
Sandostatin®	Novartis	Octreotide acetate	Microsphere suspension	Injection (IM)
Zoladex®	Astra Zeneca	Goserilin acetate	Implant	Injection (SC)
Risperdal consta®	Janssen-Cilag	Risperidone	Microparticle suspension	Injection (IM)
Decapeptyl®	Ipsen	Triptorelin		Injection (IM)
Gonapeptyl Depot®	Ferring Pharmaceutical	Triptorelin acetate	Microparticle suspension	Injection (SC or IM)

PLGA exists in different grades depending on the ratio of lactide to glycolide and polymer chain ending. All PLGAs degrade via backbone hydrolysis (bulk erosion), and the degradation products, lactic acid and glycolic acid, are ultimately metabolised by the body into CO₂ and H₂O. The two PLGAs combination as presented in Table 2 was chosen in order to obtain a drug substance release over a 35-day period. General properties of the chosen PLGAs are presented in Table 4.

Table 4 General properties of PLGAs

	50:50 PLGA ester	50:50 PLGA acid
Common Names	Resomer RG 502, PLG, PLGA, Poly (lactic-glycolic) acid, 50:50 Poly (D,L-lactide-co-glycolide), Polylactic/Polyglycolic acid, Polyglactin 910	Resomer RG 502H, PLG acid end, PLGA acid end, 50:50 Poly (D,L-lactide-co-glycolide) acid end
Structure	 Where: $n = m$ n = number of lactide repeating units m = number of glycolide repeating units z = overall number of lactide-co-glycolide repeating units	 Where: $n = m$ n = number of lactide repeating units m = number of glycolide repeating units z = overall number of lactide-co-glycolide repeating units
CAS Number	34346-01-5	26780-50-7
Empirical Formula	$[(C_3H_4O_2)_x \cdot (C_2H_2O_2)_y]CH_3$, $x:y=50:50$	$[(C_3H_4O_2)_x \cdot (C_2H_2O_2)_y]OH$, $x:y=50:50$
Description	white to off white powder	white to near white powder

DEX PS DDS was designed to release dexamethasone in the posterior segment of the eye over an extended period of 35 days. This extended release is achieved by including dexamethasone in a biodegradable polymer matrix. The polymer chosen is 50:50 PLGA. The rate of release is mainly linked to the rate of degradation of the PLGA, depending on several factors such as molecular weight and weight distribution, lactide to glycolide ratio, polymeric chain endings, etc. The mechanism for the degradation of PLGA is a hydrolysis triggered by the presence of body fluids i.e. vitreous humour in the case of DEX PS DDS.

Early formulations contained only one grade of PLGA (50/50 ratio with ester end) custom synthesised. Subsequently, it was discovered that the "acid end" form of PLGA designated 50:50 PLGA acid, combined with the 50:50 PLGA ester (equivalent to the initial PLGA), produced the desired drug release profile. "Acid end" PLGA is slightly more hydrophilic and therefore degrades faster in water. Both polymer backbones are identical, but the polymerisation process used to produce acid end PLGA involves a different chain termination agent leading to carboxylic moieties at the end of the polymer chains. During biodegradation of the implant, the degradation products are the same for both polymers, i.e. lactic acid and glycolic acid. Details of the formulation proposed can be found above. In addition, the stability of DEX PS DDS was evaluated.

20 EXAMPLES

The following examples serve to more fully describe the manner of using the above-described invention. It is understood that these examples in no way serve to limit the scope of this invention, but rather are presented for illustrative purposes.

25 EXAMPLE 1

Manufacture of Compressed Tablet Implants

Micronized dexamethasone (Pharmacia, Peapack, NJ) and micronized hydrophobic end 50/50 PLGA (Birmingham Polymers, Inc., Birmingham, AL) were accurately weighed and placed in a stainless steel mixing vessel. The vessel was

sealed, placed on a Turbula mixer and mixed at a prescribed intensity, e.g., 96 rpm, and time, e.g., 15 minutes. The resulting powder blend was loaded one unit dose at a time into a single-cavity tablet press. The press was activated at a pre-set pressure, e.g., 25 psi, and duration, e.g., 6 seconds, and the tablet was formed and ejected from the press at room temperature. The ratio of dexamethasone to PLGA was 70/30 w/w for all compressed tablet implants.

EXAMPLE 2

Manufacture of Extruded Implants

Micronized dexamethasone (Pharmacia, Peapack, NJ) and unmiconized PLGA were accurately weighed and placed in a stainless steel mixing vessel. The vessel was sealed, placed on a Turbula mixer and mixed at a prescribed intensity, e.g., 96 rpm, and time, e.g., 10-15 minutes. The unmiconized PLGA composition comprised a 30/10 w/w mixture of hydrophilic end PLGA (Boehringer Ingelheim, Wallingford, CT) and hydrophobic end PLGA (Boehringer Ingelheim, Wallingford, CT). The resulting powder blend was fed into a DACA Microcompounder-Extruder (DACA, Goleta, CA) and subjected to a pre-set temperature, e.g., 115°C, and screw speed, e.g., 12 rpm. The filament was extruded into a guide mechanism and cut into exact lengths that corresponded to the designated implant weight. The ratio of dexamethasone to total PLGA (hydrophilic and hydrophobic end) was 60/40 w/w for all extruded implants.

EXAMPLE 3

Method for Placing Implants Into the Vitreous

Implants were placed into the posterior segment of the right eye of New Zealand White Rabbits by incising the conjunctiva and sclera between the 10 and 12 o'clock positions with a 20-gauge microvitrectomy (MVR) blade. Fifty to 100 µL of vitreous humor was removed with a 1-cc syringe fitted with a 27-gauge needle. A sterile trocar, preloaded with the appropriate implant (drug delivery system, DDS), was inserted 5 mm through the sclerotomy, and then retracted with the push wire in

place, leaving the implant in the posterior segment. Sclerae and conjunctivae were then closed using a 7-0 Vicryl suture.

EXAMPLE 4

5 In vivo Release of Dexamethasone From 350 μ g Dexamethasone Compressed Tablet 10 Implants

Example 4 demonstrates the high initial release but generally lower intravitreal concentration of dexamethasone from compressed tablet implants as compared to extruded implants. The 350 μ g compressed tablet implant (350T) was placed in the right eye of New Zealand White Rabbits as described in Example 3. Vitreous samples were taken periodically and assayed by LC/MS/MS to determine *in vivo* dexamethasone delivery performance. As seen in Figure 1, dexamethasone reached detectable mean intravitreal concentrations from day 1 (142.20 ng/ml) through day 35 (2.72 ng/ml), and the intravitreal concentration of dexamethasone gradually decreased over time.

In addition to the vitreous samples, aqueous humor and plasma samples were also taken. The 350T showed a gradual decrease in aqueous humor dexamethasone concentrations over time, exhibiting a detectable mean dexamethasone aqueous humor concentration at day 1 (14.88 ng/ml) through day 21 (3.07 ng/ml), as demonstrated in Figure 3. The levels of dexamethasone in the aqueous humor strongly correlated with the levels of dexamethasone in the vitreous humor, but at a much lower level (approximately 10-fold lower). Figure 4 shows that only trace amounts of dexamethasone was found in the plasma.

25 EXAMPLE 5

In vivo Release of Dexamethasone From 350 μ g Dexamethasone Extruded Implants

Example 5 demonstrates the lower initial release and generally more sustained intravitreal concentration of dexamethasone from extruded implants. The 350 μ g

extruded implant (350E) was placed in the right eye of New Zealand White Rabbits as described in Example 3. Vitreous samples were taken periodically and assayed by LC/MS/MS to determine *in vivo* dexamethasone delivery performance. Referring to Figure 1, 350E showed detectable mean vitreous humor concentrations on day 1 (10.66 ng/ml) through day 28 (6.99 ng/ml). The 350T implant had statistically significant higher dexamethasone concentrations on day 1 ($p=0.037$) while the 350E had a statistically significant higher dexamethasone level on day 21 ($p=0.041$).

In addition to the vitreous samples, aqueous humor and plasma samples were also taken. In Figure 3, the 350E showed detectable mean dexamethasone aqueous humor concentrations at day 1 (6.67 ng/ml) through day 42 (2.58 ng/ml) with the exception of day 35 in which the values were below the quantification limit. On the whole, the levels of dexamethasone in the aqueous strongly correlated with the levels of dexamethasone in the vitreous humor, but at a much lower level (approximately 10-fold lower). Figure 4 demonstrates that only a trace amount of dexamethasone was found in the plasma.

EXAMPLE 6

In vivo Release of Dexamethasone From 700 μ g Dexamethasone Compressed Tablet Implants

Example 6 also shows the high initial release and generally lower intravitreal concentration of dexamethasone from compressed tablet implants. The 700 μ g compressed tablet dosage form (700T) was placed in the right eye of New Zealand White Rabbits as described in Example 3. Vitreous samples were taken periodically and assayed by LC/MS/MS to determine *in vivo* dexamethasone delivery performance. As seen in Figure 5, the 700T reached detectable mean dexamethasone vitreous humor concentrations at day 1 (198.56 ng/ml) through day 42 (2.89 ng/ml), and a gradual decrease in the intravitreal dexamethasone concentration over time.

In addition to the vitreous samples, aqueous humor and plasma samples were also obtained. As seen in Figure 6, the 700T exhibited a gradual decrease in aqueous

humor dexamethasone concentrations over time, and reached detectable mean dexamethasone aqueous humor concentrations at day 1 (25.90 ng/ml) through day 42 (2.64 ng/ml) with the exception of day 35 in which the values were below the quantification limit. The levels of dexamethasone in the aqueous humor strongly correlated with the levels of dexamethasone in the vitreous humor, but at a much lower level (approximately 10-fold lower). Figure 7 demonstrates that only a trace amount of dexamethasone was found in the plasma.

EXAMPLE 7

In vivo Release of Dexamethasone From 700 μ g Dexamethasone Extruded Implants

Example 7 also illustrates the lower initial release and generally higher intravitreal concentration of dexamethasone from extruded implants. The 700 μ g extruded implant (700E) was placed in the right eye of New Zealand White Rabbits as described in Example 3. Vitreous samples were taken periodically and assayed by LC/MS/MS to determine *in vivo* dexamethasone delivery performance. As seen in Figure 5, the 700E had a mean detectable vitreous humor concentration of dexamethasone from day 1 (52.63 ng/ml) through day 28 (119.70 ng/ml).

In addition to the vitreous samples, aqueous humor and plasma samples were also taken. As seen in Figure 6, the 700E reached a detectable mean aqueous humor concentration on day 1 (5.04 ng/ml) through day 28 (5.93 ng/ml). The levels of dexamethasone in the aqueous strongly correlated with the levels of dexamethasone in the vitreous humor, but at a much lower level (approximately 10-fold lower). Figure 7 demonstrates that only a trace amount of dexamethasone was found in the plasma.

EXAMPLE 8

Extrusion Methods for Making an Implant

1. DEX PS DDS implants were made by a tableting process, by a single extrusion process and by a double extrusion process.

The excipients (polymers) used for the DEX PS DDS implant made were two grades of 50:50 Poly (D,L lactide co glycolide) ester end and acid end. Both excipients were a pharmaceutical grade of non-compendial material.

5 The preferred specifications of three batches of both 50:50 Poly PLGA ester used to make implants are shown in Table A. The preferred specifications of three batches of 50:50 Poly PLGA acid use to make implants are shown in Table B.

Table A Preferred Specifications for 50:50 PLGA ester

Tests	Specifications	1001933	1004907	1004925
Appearance: Colour and shape	White to off white	Off white	Off white	Off white
Odour	Odourless to almost odourless	almost odourless	almost odourless	almost odourless
Identification	^1H -NMR spectra conforms to reference	conforms	conforms	conforms
Polymer composition				
DL-Lactide units	48 to 52%	51	51	51
Glycolide units	52 to 48%	49	49	49
Inherent viscosity	0.16 to 0.24 dl/g	0.24	0.19	0.19
Water	□0.5%	Conforms	Conforms	Conforms
Residual monomers				
DL-lactide	□0.5%	Conforms	Conforms	Conforms
Glycolide	□0.5%	Conforms	Conforms	Conforms
Residual solvents				
Acetone	□0.1%	Conforms	Conforms	Conforms
Toluene	□0.089%	Conforms	Conforms	Conforms
Total	□0.1%	Conforms	Conforms	Conforms
Tin	□100 ppm	30	31	35
Heavy metals	□10 ppm	Conforms	Conforms	Conforms
Sulphated ashes	□0.1%	Conforms	Conforms	Conforms

Table B Preferred Specifications for 50:50 PLGA acid

Test	Specification	1006825	1008386	1009848
Appearance: Colour and shape	White to nearly white	White	White	Off white
Odour	Odourless to nearly odourless	Odourless	Odourless	Almost odourless
Identification	¹ H-NMR spectra conforms to reference	Conforms	Conforms	Conforms
Polymer composition				
DL-Lactide units	48 to 52%	51	51	51
Glycolide units	52 to 48%	49	49	49
Inherent viscosity	0.16 to 0.24 dl/g	0.19	0.19	0.19
Water	□0.5%	Conforms	Conforms	Conforms
Residual monomers				
DL-lactide	□0.5%	Conforms	Conforms	Conforms
Glycolide	□0.5%	Conforms	Conforms	Conforms
Residual solvents				
Acetone	□0.1%	Conforms	Conforms	Conforms
Toluene	□0.089%	Conforms	Conforms	Conforms
Tin	□200ppm	149	83	141
Heavy metals	□10 ppm	Conforms	Conforms	Conforms
Sulphated ashes	□0.1%	Conforms	Conforms	Conforms
Acid number	□6.5 mg _{KOH} /g	11	9	12

Preferred specifications of Polymers use to make Implants

Polymer composition: It was determined that the ratio of lactide to glycolide is essential for the kinetic of degradation of the polymer and hence the dexamethasone release profile of the implant. It was controlled in a 48% to 52% (wt %) range to ensure consistency of active agent release.

Inherent viscosity: the inherent viscosity is essential for the kinetics of degradation of the polymer and hence the dexamethasone release profile of the implant. It is a measure of the size of the polymer backbone and the size distribution

(i.e. molecular weight and weight distribution). It was controlled in a 0.16 to 0.24 dl/g range to ensure consistency of release.

Water: The moisture content of the polymer influences its stability during the shelf life and is a facilitating factor for the biodegradation of the polymer matrix. It was controlled below 0.5% to ensure stability of the excipients as well as the drug substance (dexamethasone) and to ensure consistency of the (dexamethasone) release profile.

Residual monomers: residual monomers indicate the completion of the synthesis of the polymer and was controlled below 0.5 wt. %.

Residual solvents:

- Acetone was controlled to below 0.1 wt. %
- Toluene was controlled to be kept below 0.0890 wt. %.

Acid number: the acid number measures the number of chain ends in the PLGA acid polymer. The number of acid polymer endings facilitates the ingress of moisture upon injection of the implant and influences the release profile of the implant. It was controlled to be higher than 6.5 mg KOH/g to ensure consistency of release profile.

Preferred Dexamethasone Characteristics

The particle size and particle size distribution of the dexamethasone is regarded as a critical parameter for the homogeneity of the DEX PS DDS. A preferred dexamethasone particle size distribution has at least 75% of the particles of dexamethasone smaller than (i.e. diameter less than) 10 μ m. A more preferred dexamethasone particle size distribution has at least 99% of the particles of dexamethasone smaller than (i.e. diameter less than) 20 μ m. We found that use of such small particles of dexamethasone in the implant provides for a more uniform distribution of the active agent in the implant (i.e. no clumping) which leads to a

more uniform release of the active agent from the implant upon implantation of the implant.

In addition to all the Ph. Eur. tests for dexamethasone, additional tests were performed on the dexamethasone using a particle size analyser and an additional
5 analytical method, so as to ensure that the dexamethasone used in the DEX PS DDS had the preferred or the more preferred particle size and particle size distribution.

In our invention dexamethasone a particle size and particle size distribution is an important factor because homogeneity of the dexamethasone affects release characteristics.

10 It is additionally preferred that the dexamethasone used in the present invention comprise $\square$ 1% of total impurities, including $\square$ 0.50% of dexamethasone acetate, $\square$ 0.25% of bethamethasone, $\square$ 0.25% of 3 keto delta 4 derivative and $\square$ 0.10% of any other impurity.

A representative formula for a typical 80 g manufacturing batch (used to make
15 an implant manufactured by the tableting, single extrusion or double extrusion process) is provided in Table 2. For the 350 μ g and 700 μ g dosages, the bulk manufacturing and terminal sterilisation processes are identical.

A flow diagram of the three different manufacturing processes is shown by Figure 11.

20 2. A single extrusion process was used to made an implant. In a continuous extrusion, single extrusion manufacturing process, the micronised dexamethasone and un-micronised polymer was blended, before being loaded into a twin-screw compound extruder, and then subjected to a set temperature and screw speed. The filament was extruded into a guide mechanism and cut into exact lengths that
25 correspond to the correct DEX PS DDS weight. This continuous extrusion process

was more controllable and more predictable than the tableting process. This is illustrated in the *in vitro* release profiles of the DEX PS DDS as show in Figure 12.

Four lots of 700 μ g DEX PS DDS, two manufactured by the tableting process and two by the single extrusion process were studied. With the single extrusion
5 process the only difference between the two doses is that the 350 μ g dose filament is cut from the same extrudate (same formulation) as 700 μ g dose filament but is half as long. At 5 time points, over a 28-day period, 12 DEX PS DDS units from each lot were tested. The standard deviations for the mean dexamethasone release rates were found larger for the two tabletted lots than for the two extruded lots. A three-fold
10 reduction in standard deviations across the release profile was observed with the extruded versus the tabletted product. In addition, the initial burst release is reduced with implants manufactured by a single extrusion process, as compared to implants made by a tableting process.

These results were confirmed in a GLP *in vivo* pharmacokinetics study in
15 rabbits comparing the release of dexamethasone from the tabletted and the extruded DEX PS DDS. It was shown that the tabletted and the single extruded DEX PS DDS release the same amount of dexamethasone over the same period, providing approximately a 35-day delivery.

To further characterise and compare the DEX PS DDS manufactured by the
20 tableting and single extrusion processes, scanning electron microscope (SEM) photographs were taken to assess physical appearance. Figure 13 shows that the single extruded DEX PS DDS is more uniform than was the tabletted implant. It was found that not only is this gives a more consistent *in vitro* release profile from the single extruded implant, but also with its increased resistance to crushing. Using a
25 texture analyser it was shown that a 3-fold increase in force (1200 g compared to 400g) was required to crush a single extruded implant compared to a tabletted one. This demonstrates that the extruded product is more able to withstand handling.

Additionally, it was determined that the DEX PS DDS made by single extrusion and by double extrusion processes is stable during a minimum of 12 months (and for as long as 18-24 months) when stored at 25°C/60%RH and a minimum of 6 months at 40°C/75%RH. Stability was determined based upon dexamethasone
5 potency, dexamethasone impurities (acid, ketone, aldehyde and total impurities), moisture content, applicator actuation force, implant fracture force/fracture energy and *in vitro* dissolution dexamethasone release profile and sterility..

3. The inventors improved the single extrusion process by (1) micronising the polymers prior to blending and (2) adding a second extrusion after pelletisation of the
10 first extruded filament. When both 50:50 PLGA acid and 50:50 PLGA ester were micronised an acceptable DEX PS DDS homogeneity was obtained. Homogeneity promotes a more even and regular dissolution of the polymer and release of the dexamethazone active agent. The PLGAs were milled using an air jet process. Figure 14 presents batch-to-batch versus in-batch variability from batches made of
15 milled (i.e. micronised) and un-milled (i.e. unmiconized) PLGAs. It clearly demonstrates that the double extrusion process allows better control, especially where in-batch variability was reduced from a 94.7%LC to 107.0%LC range (unmilled PLGAs) to a 98.9%LC to 101.5%LC range (milled PLGAs. "LC" means label claim (a regulatory term), that is the amount of dexamethasone present in the implant (350
20 µg or 700µg), as measured by various *in vitro* assays, such as by HPLC.

Single and double extrusion processes were compared. As shown by Figure
15 implants made by a double extrusion process had released about 60% of the dexamethasone by day 14, while the single extrusion implants had released about 40% of its dexamethasone load by day 14, although total dexamethasone released
25 was comparable by day 21. Therefore, where more release of dexamethasone is desired sooner, the double extrusion process is a preferred process for making the DEX PS DDS. A double extrusion process also provides for a higher yield of the

desired filament implant, i.e. with a uniform distribution of dexamethasone throughout the implant polymer.

A detailed manufacturing schematic flow diagram for the double extrusion implant is provided by Figure 16. The major equipment used in the manufacture of
5 DEX PS DDS is listed in Table C.

Table C Major Equipment Used in the Manufacture of DEX PS DDS

Step	Purpose	Equipment Description
1	Milling both PLGAs	Jet Mill
2	Powder blending	Shaker
3	First extrusion	Extruder and Force Feeder, Puller Assembly and Filament Cutter
4	Pelletising	Stainless steel ball and bottle Shaker
5	Second extrusion	Extruder and Force Feeder, Puller Assembly and Filament Cutter
6	Automated DDS cutting and inspection procedure	Guillotine Cutter and Vision Inspection System
7-8	Applicator Assembly	Applicator loading fixture and Heat sealer

4. The specifics of the double extrusion process used are as follows

(a) Milling of PLGAs (Resomers RG502 and RG502H)

10 30 grams of RG502 (50:50 PLGA ester) were milled using the Jet-Mill (a vibratory feeder) at milling pressures of 60 psi, 80 psi and 80 psi for the pusher nozzle, grinding nozzle, and grinding nozzle, respectively. Next, 60 grams of RG502H were milled using the Jet-Mill at milling pressure of 20 psi, 40 psi and 40 psi for the pusher nozzle, grinding nozzle, and grinding nozzle, respectively. The
15 mean particle size of both RG502 and RG502H was measured using a TSI 3225 Aerosizer DSP Particle Size Analyzer. Preferably, both milled polymers must have a mean particle size of no greater than 20 um.

(b) Blending of PLGAs and Dexamethasone

48 grams of dexamethasone, 24 grams of milled RG502H and 8 grams of milled RG502 were blended using the Turbula Shaker set at 96 RPM for 60 minutes.

(c) First Extrusion

- 5 (1) All 80 grams of the blended dexamethasone/RG502H/RG502 mixture was added to the hopper of a Haake Twin Screw Extruder. The Haake extruder was turned on and set the following parameters:

Barrel Temperature: 105 degrees C.

Nozzle Temperature: 102 degrees C.

10 Screw Speed: 120 RPM

Feed Rate Setting: 250

Guide Plate Temperature: 50 - 55 degrees C.

Circulating water bath: 10 degrees C.

- (2) Filament were collected. The first filament comes out about 15-25
15 minutes after the addition of the powder blend. Discard the first 5 minute of extruded filaments. Collecting the remaining filaments until exhaustion of extrudates; this normally takes 3-5 hours.

(d) Pelletization

- The filaments from step 3 above were pelletized using the Turbula Shaker and
20 one 19 mm stainless steel ball set at 96 RPM for 5 minutes.

(e) Second Extrusion

- (1) All pellets were added into the same hopper and the Haake extruder was turned on.

The following parameters were set on the Haake extruder:

25 Barrel Temperature: 107 degrees C.

Nozzle temperature: 90 degrees C.

Screw speed: 100 RPM

Guide Plate Temperature: 60-65 degrees C.

Circulation water bath: 10 degrees C.

(2) All extruded filaments were collected until exhaustion of extrudates. This normally takes about 3 hours.

(f) Processing of Bulk Filament to Dosage strengths – 350 μ g or 700 μ g

5 DEX PS DDS was prepared as 350 μ g or 700 μ g dosage forms by cutting the filaments to the appropriate length.

(g) Insertion of DEX PS DDS into the Applicator

The DEX PS DDS was inserted into the Applicator System during the applicator assembly process. All operations took place in a Class 10 000 clean room.

10 (h) Packaging of DEX PS DDS Applicator System

The assembled DEX PS DDS Applicator System was placed into a foil pouch containing a small bag of desiccant and heat-sealed. Samples for pre-sterilisation bioburden testing were taken prior to step 9.

(i) Gamma Radiation Sterilization of DEX PS DDS Applicator System

15 The sealed foil pouches containing the finished DEX PS DDS Applicator System and a small desiccant bag were placed into a cardboard box and the box sealed. Terminal sterilisation of these product containing boxes was accomplished by exposure to a dose within the range of 25-40 kGy of gamma-radiation. Samples from each batch were tested for sterility according to Ph. Eur. and USP requirement.

20 (j) Labelling of DEX PS DDS Applicator

The single and double extruded implants had the preferred characteristics shown by Tables D and E, respectively.

Table D In Process Controls results for the first extrusion

Batch Number		03J001	03H004	03M001
Batch size		80g	80g	80g
Parameter	Specifications			
Filament density	0.85 to 1.14 g/cm ³	1.03	1.01	1.04
Uniformity	85.0 to 115.0% ⁽¹⁾	99.3	100.5	98.7
Potency	97.0 to 103.0% label strength	100.1	100.0	99.8
Degradation products	□ 1.5% total	0.2	0.2	0.2
	□ 0.75% acid	ND	ND	ND
	□ 0.75% ketone	□ 0.08	□ 0.10	□ 0.13
	□ 0.75% aldehyde	□ 0.15	□ 0.10	□ 0.12

⁽¹⁾ Percentage of target weight**Table E In Process Control results for the second extrusion**

Batch number		03J001	03H004	03M001
Batch size		80g	80g	80g
Parameter	Specifications			
Appearance	White to off white	pass	pass	pass
Filament density	1.10 to 1.30 g/cm ³	1.18	1.13	1.19
Diameter	□ 80% within 0.0175 to 0.0185 inch	100	100	100
Fracture force	□ 2g	9.88	9.39	9.52
Fracture energy	□ 0.9μJ	5.88	4.54	4.64
Moisture	□ 1.0%	0.4	0.4	0.4
Foreign particulate	No visible foreign materials	Pass	Pass	Pass
Insoluble mater (for information only)	Particle count			
	Diameter □ 10μm	17	26	2.6
	Diameter □ 25μm	0.5	1	0
Dexamethasone identity	Positive for dexamethasone	positive	positive	positive
Potency	95.0 to 105.0 % label strength	98.5	101.2	99.9
Degradation products	□ 2% total	1.1	0.6	1.0
	□ 0.5% acid	ND	ND	ND
	□ 1.0% ketone	0.4	0.2	0.4
	□ 1.0% aldehyde	0.7	0.4	0.5
Dexamethasone release	See Table 2.1.P.5.1-1	Pass	Pass	Pass

Batch number		03J001	03H004	03M001
Batch size		80g	80g	80g
Parameter	Specifications			
Uniformity	85.0 – 115.0% Label Strength (LS) Stage 1 (n=10): If one unit is outside the range and between 75% and 125% LS or RSD $\leq$ 6.0%, test 20 more units. Stage 2 (n=20): pass if no more than 1 unit is outside the range, and is between 75% and 125% LS, and the RSD $\leq$ 7.8%.	97.0% all values within range	97.1% all values within range	98.0% all values within range

Table F sets forth further preferred specifications for both the DEX PS DDS implant and the applicator.

Table F Preferred specifications

Attribute		Specifications
Implant appearance		White to off-white, rod shaped Drug Delivery System (DDS), essentially free of foreign matter.
Fracture	Force	Minimum 2.0 g
	Energy	Minimum 0.85 μ joule
Moisture content		No more than 1%
Foreign particulates		No visible foreign material
Insoluble matter		Record particle count for information only (diameter $\leq$ 10 μ m and $\leq$ 25 μ m)
Dexamethasone identity		Positive for dexamethasone
Dexamethasone potency		90.0 to 110.0%LC
Impurities		Dexamethasone acid not greater than 0.5% HPLC area Dexamethasone ketone not greater than 1.0% HPLC area Dexamethasone aldehyde not greater than 1.0% HPLC area Total degradation not greater than 2% HPLC area
Weight Range		700 μg dose: 1.050 mg to 1.284 mg (1.167 mg +/- 10%) 350 μg dose: 0.525 mg to 0.642 mg (0.583 mg +/- 10%)
Content uniformity		85% to 115% Label Claim

Attribute	Specifications
<i>In vitro</i> Dissolution test (% of total amount of dexamethasone released)	Ranges: 24 hours: not greater than 10.0% 7 days: not greater than 30.0% 14 days: 25.0% to 85.0% 21 days: not less than 50%
Applicator Actuation force required	No more than 5.0 lbs

Implants and Applicators made as set forth above were found to be within the parameters of the preferred specifications.

Preferred Applicator A preferred applicator to use to implant the DEX PS DDS is shown in international patent publication WO 2004/026106, published April 1, 2004. The applicator was designed to facilitate the insertion of the implant in the posterior segment of the eye. The implant is housed in the needle of the applicator. The applicator is designed to fit comfortably into the hand of the physician, and to allow for single-handed operation. It is similar in size to retinal forceps, measuring 165 mm in length by 13 mm in width. Figure 17 provides a cut-away side view of the applicator illustrating the typical functions and positions of all the elements.

As the lever is depressed, it applies a force on the linkage, which collapses and moves the plunger forward into the needle, pushing the DEX PS DDS into the posterior chamber of the eye. Once the DEX PS DDS is delivered, the lever then latches within the Applicator housing to signal use and prevent any reuse. The needle used is a 22-gauge thin-wall hypodermic needle. A silicone o-ring, is placed into a slot in the needle to retain the DEX PS DDS within the needle and remains outside the eye, in contact with the conjunctiva. To ensure that air is not introduced into the eye, the applicator has been designed to vent. A small gap between the DEX PS DDS and inner needle wall allows air to move back through and out of the needle as the DEX PS DDS is being delivered. The small size of this gap prevents fluid from flowing out of the eye through the needle. The components of the Applicator that may

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contact the patient during use are the plunger, needle and o-ring. The plunger and needle are manufactured from materials of known biocompatibility, and with a history of human use. Biocompatibility of the o-ring was evaluated through cytotoxicity testing.

- 5 The applicator is packed with desiccant in a pouch designed to protect the implant from humidity. The packaged implant in the applicator is then sterilised by gamma irradiation. The pouch also ensures that the product remains sterile during the shelf life.

- 10 The DEX PS DDS is terminally sterilised by gamma irradiation, in its applicator as presented packed in the foil pouch, using a 25 to 40 kGy dose. Terminal sterilisation process steam sterilisation (autoclaving) is not used because the polymers used for the controlled release are extremely sensitive to moisture and heat and degrade even with non-compendial low temperature sterilisation cycles.

- 15 The DEX PS DDS Applicator System is a sterile, single use applicator intended to deliver one DEX PS DDS. The DEX PS DDS is loaded into the needle of the Applicator during the assembly process. It is then packaged in a foil pouch with desiccant and terminally sterilised by gamma irradiation.

- 20 Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and

25

modifications may be made thereto without departing from the spirit and scope of the appended claims.

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REPLACEMENT SHEET

PCT/US2005/026500
Allergan, Inc.

Docket 17659-CIP (OCU)
April 29, 2006

Claims

1. A method for making a bioerodible implant for treating an ocular condition, the method comprising the steps of:
 - (a) milling a biodegradable polymer;
 - (b) blending the milled biodegradable polymer and particles of an active agent, to thereby obtain a blended mixture of the milled biodegradable polymer and the particles of the active agent, wherein at least about 75% of the particles of the active agent have a diameter of less than about 20 μ m;
 - (c) carrying out a first extrusion of the blended mixture, to thereby obtain a first extrusion product;
 - (d) pelletizing the first extrusion product; and
 - (e) carrying out a second extrusion of the pelletized first extrusion product, thereby obtaining a bioerodible implant for an ocular condition.
2. The method of claim 1 wherein at least about 99% of the particles of the active ingredient have a diameter of less than about 20 μ m.
3. The method of claim 1 wherein the active agent is selected from the group consisting of ace-inhibitors, endogenous cytokines, agents that influence basement membrane, agents that influence the growth of endothelial cells, adrenergic agonists or blockers, cholinergic agonists or blockers, aldose reductase inhibitors, analgesics, anesthetics, antiallergics, anti-inflammatory agents, steroids, antihypertensives, pressors, antibacterials, antivirals, antifungals, antiprotozoals, anti-infective agents, antitumor agents, antimetabolites, and antiangiogenic agents.
4. The method of claim 1 wherein the active agent comprises an anti-inflammatory agent or any derivative thereof.

REPLACEMENT SHEET

5. The method of claim 1 wherein the active agent comprises a steroidal anti-inflammatory agent or any derivative thereof.
6. The method of claim 4 wherein the active agent is selected from the group consisting of cortisone, dexamethasone, fluocinolone, hydrocortisone, methylprednisolone, prednisolone, prednisone, triamcinolone, and any derivative thereof.
7. The method of claim 1 wherein the active agent comprises dexamethasone.
8. The method of claim 1 wherein the bioerodable polymer comprises poly(lactic-co-glycolic)acid (PLGA) copolymer.
9. The method of claim 8 wherein the ratio of lactic to glycolic acid monomers is about 50/50 weight percentage.
10. The method of claim 8 wherein the PLGA copolymer is about 20 to about 90 weight percent of the bioerodable implant.
11. The method of claim 10 wherein the PLGA copolymer is about 40% by weight of the bioerodable implant.
12. The method of claim 1 wherein the implant is sized for implantation in an ocular region.
13. The method of claim 12 wherein the ocular region is selected from the group consisting of the anterior chamber, the posterior chamber, the vitreous cavity, the choroids, the suprachoroidal space, the conjunctiva, the subconjunctival space, the episcleral space, the intracorneal space, the epicorneal space, the sclera, the pars plana, surgically-induced avascular regions, the macula, and the retina.
14. The method of claim 13 wherein the ocular region is the vitreous cavity.
15. A bioerodible implant for treating an ocular condition made by the method of claim 1.

Figure 1

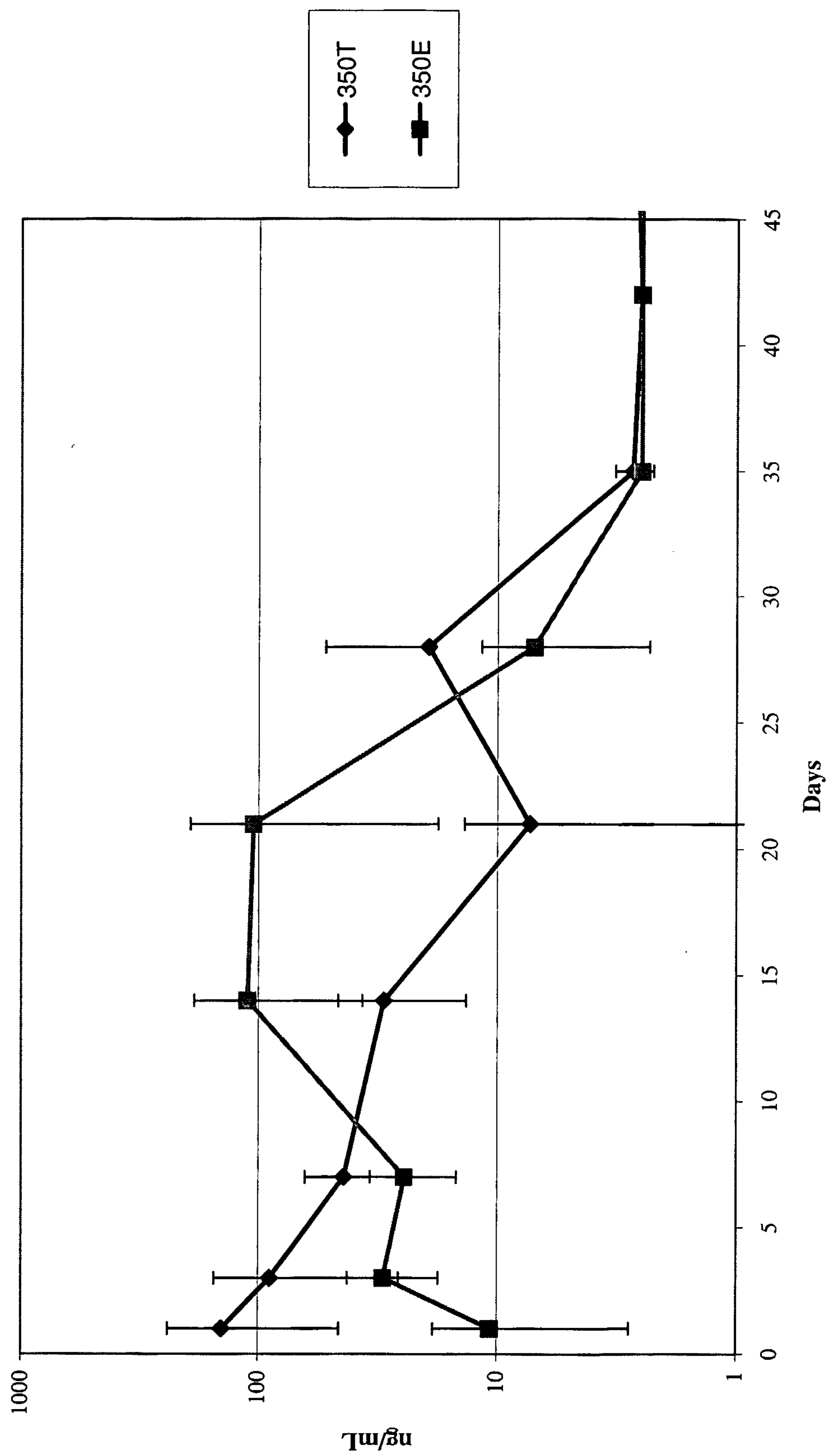


Figure 2

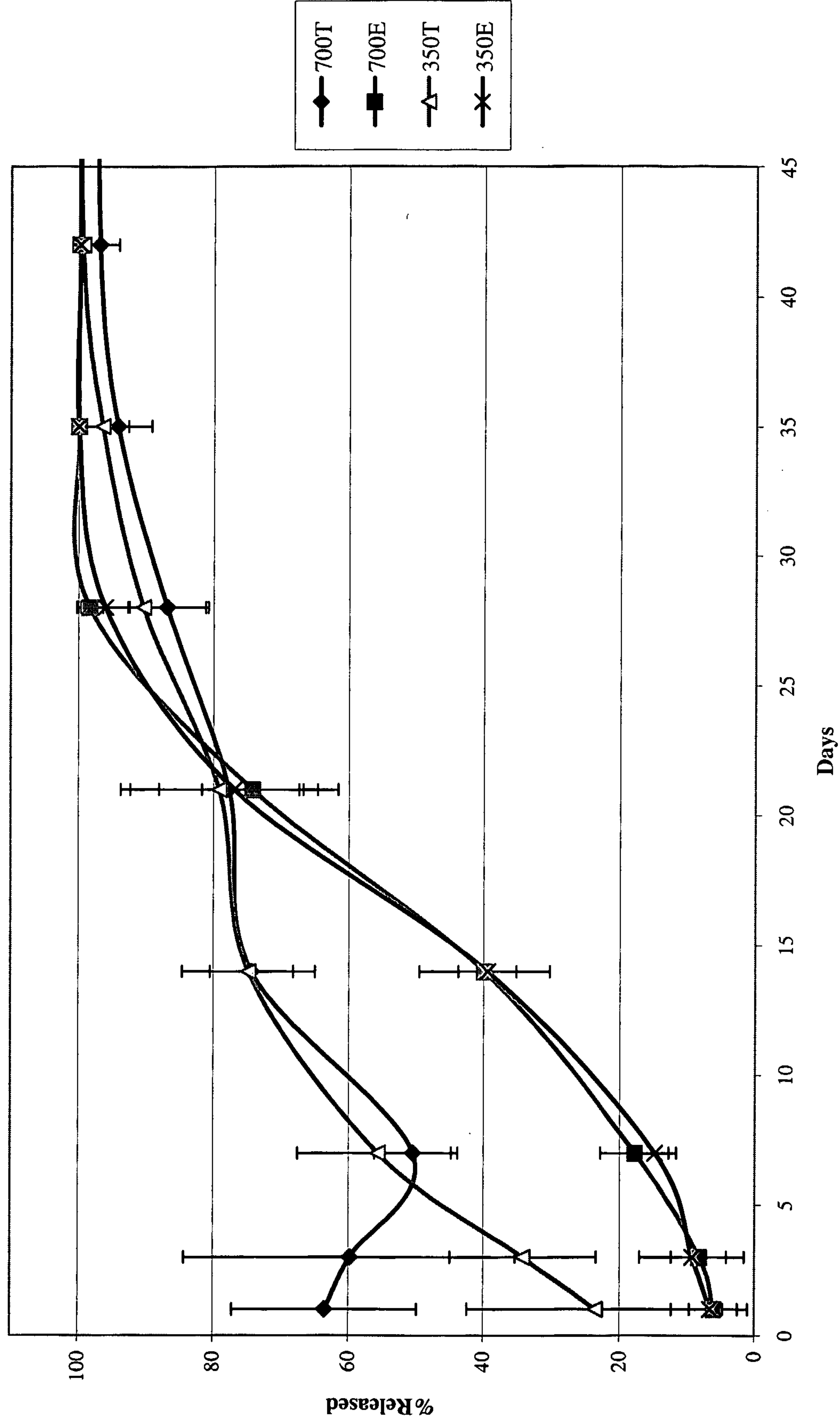


Figure 3

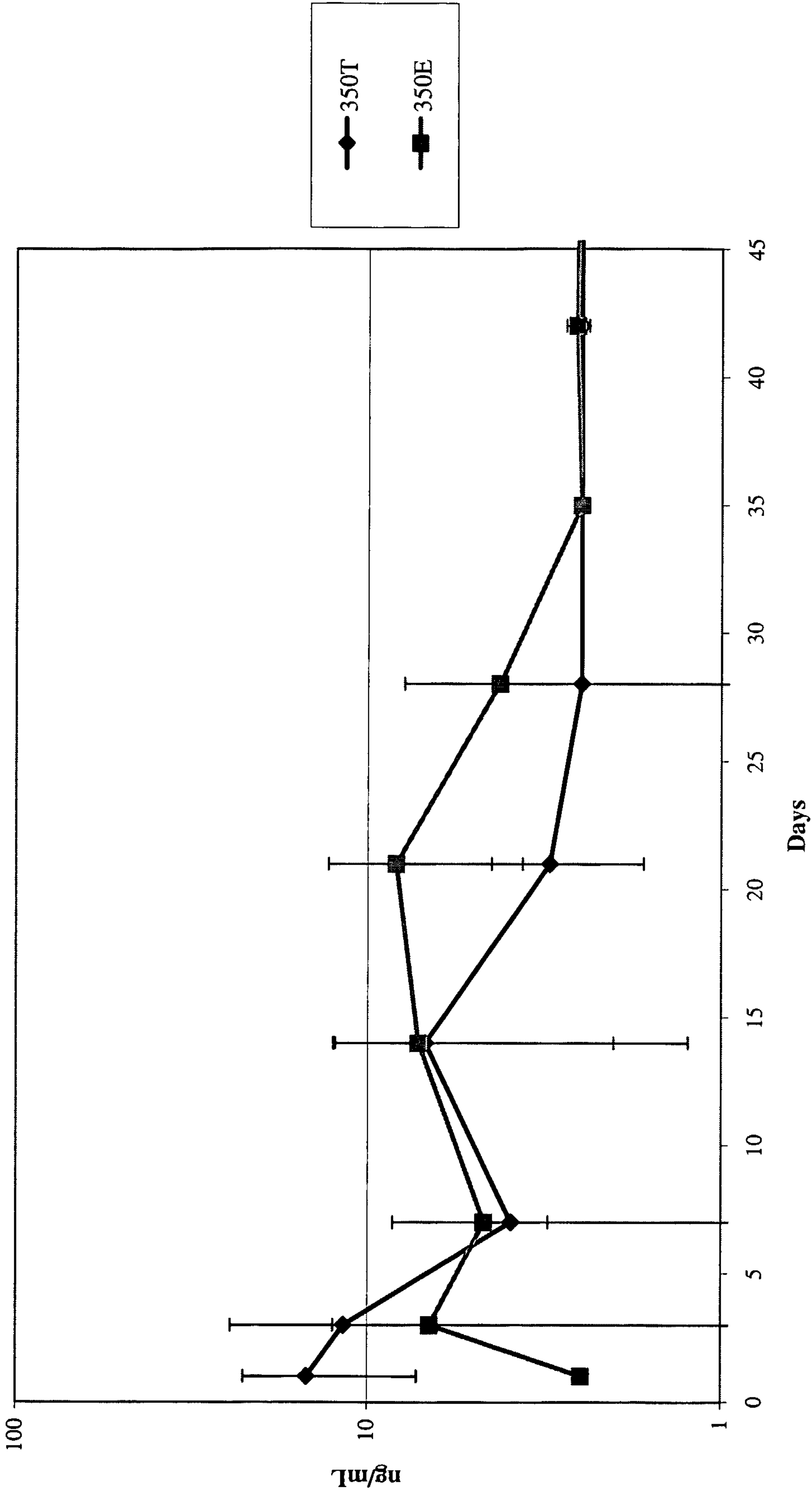
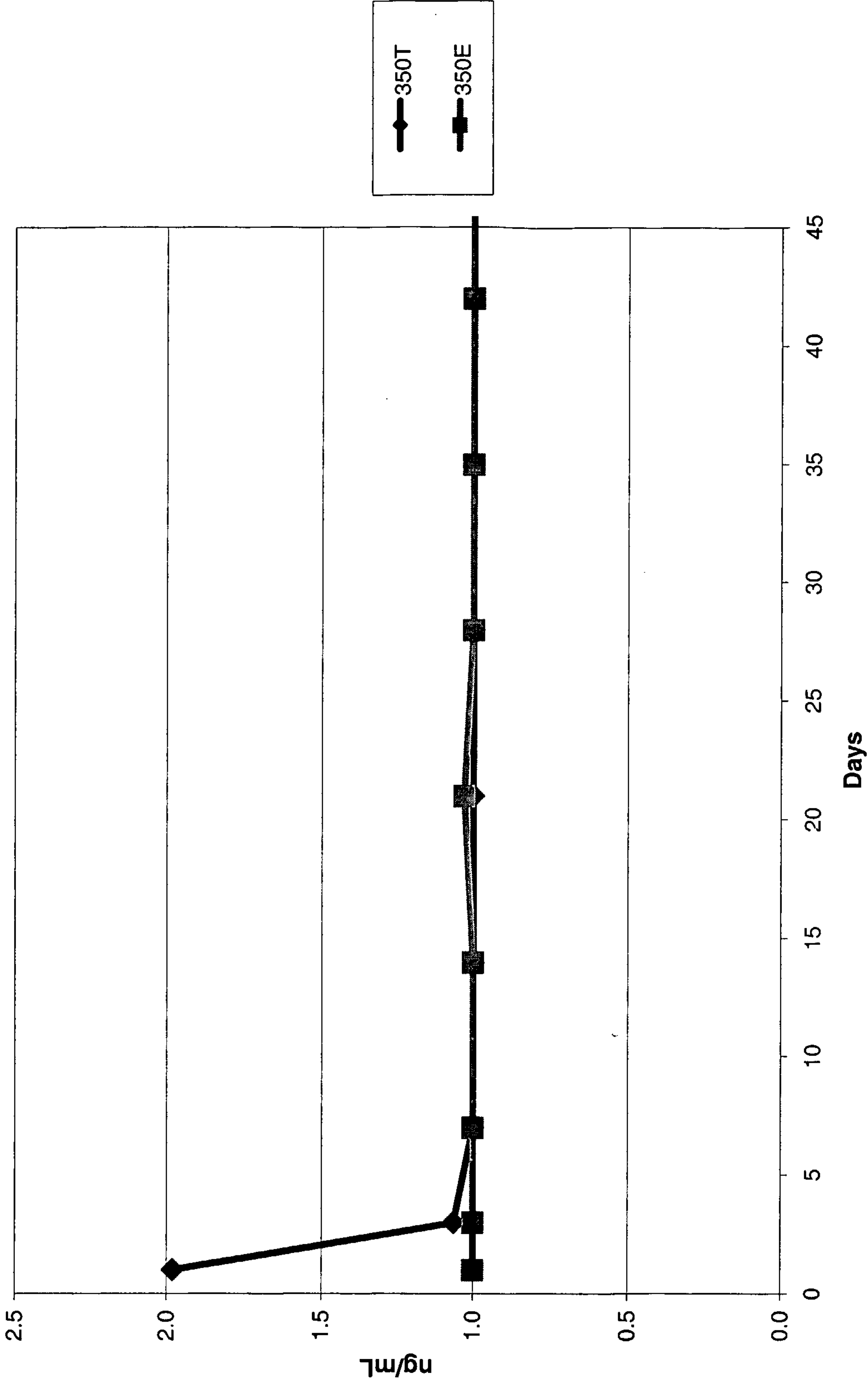


Figure 4



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Figure 5

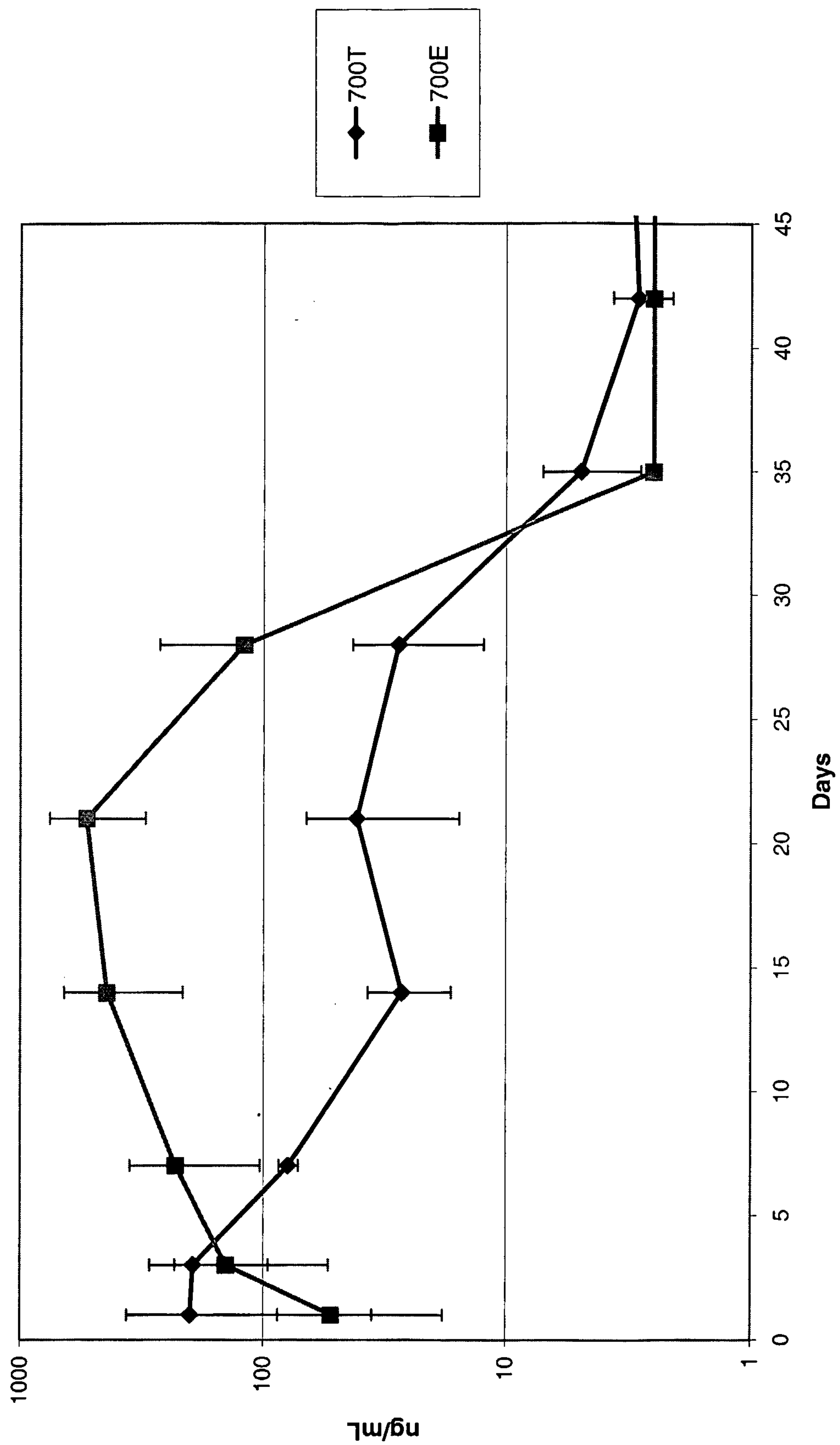
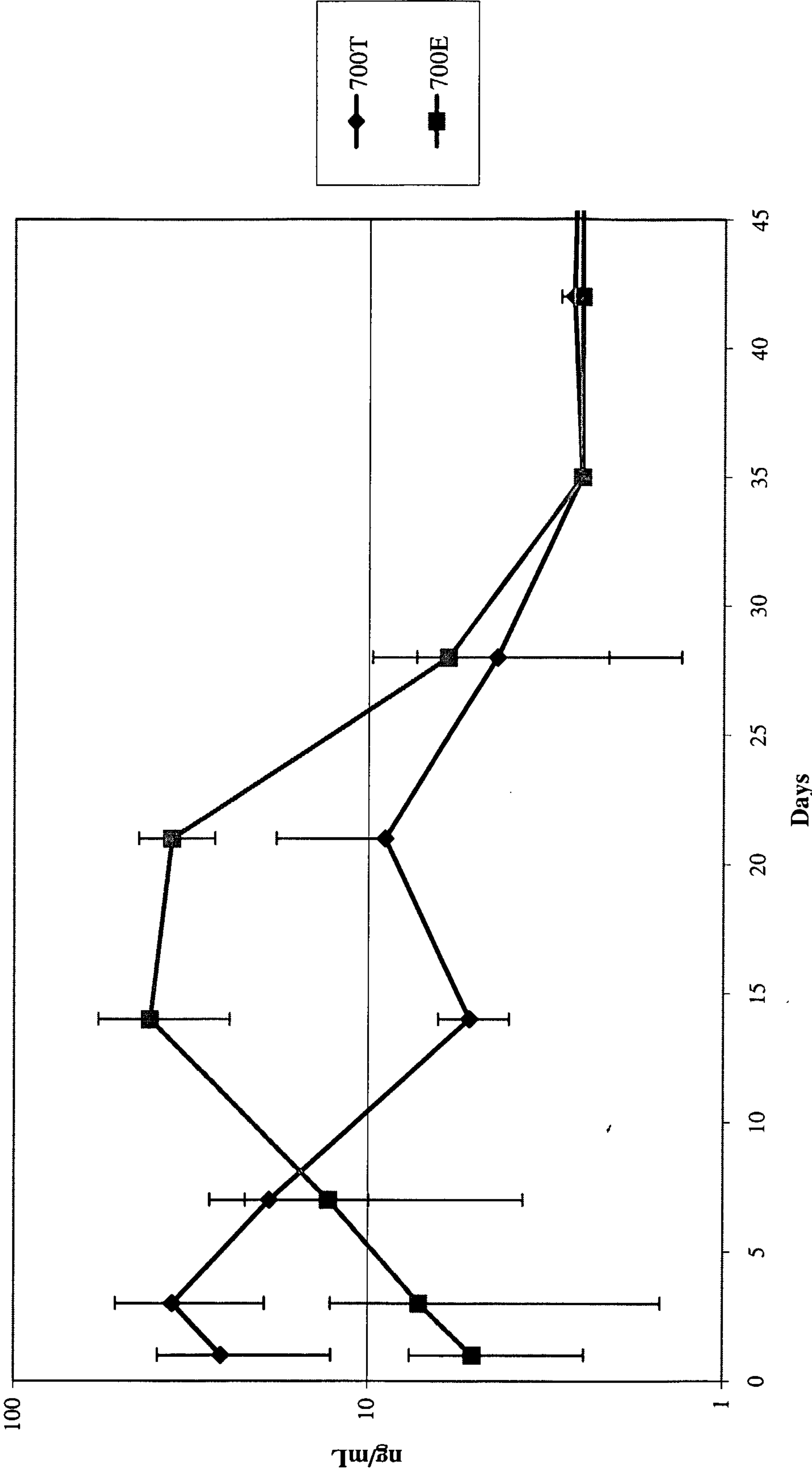
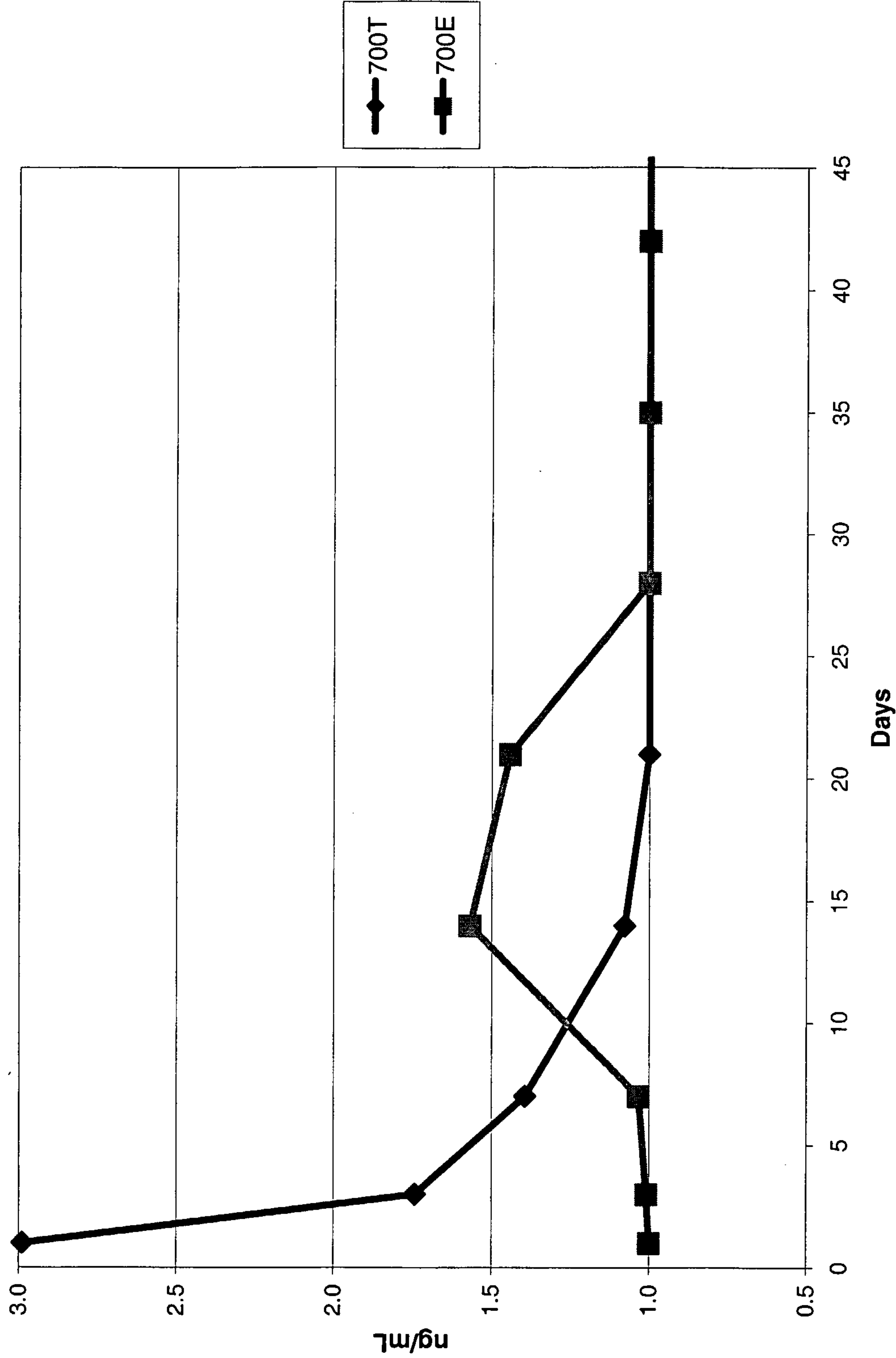


Figure 6



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Figure 7



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Figure 8

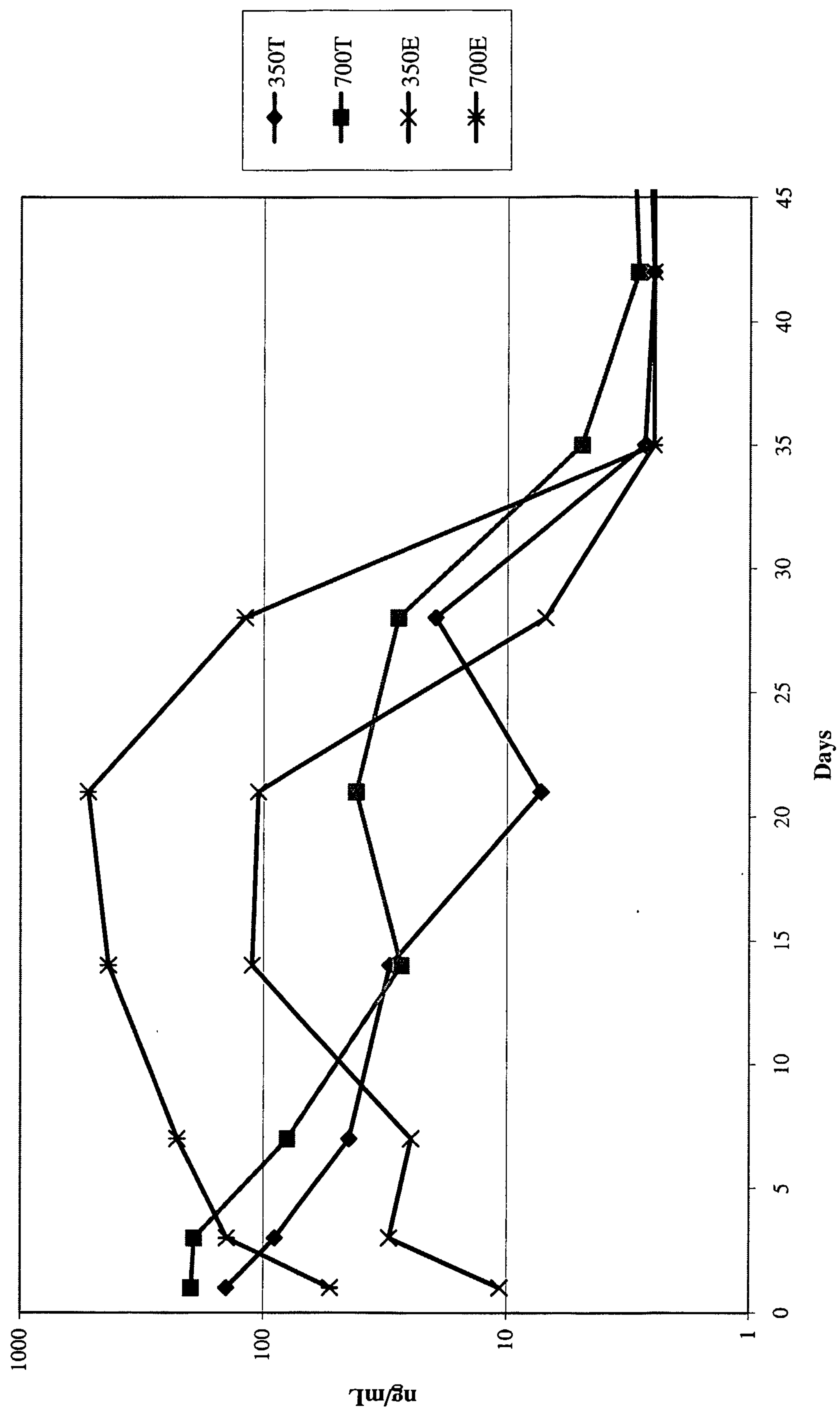


Figure 9

Release Profile Comparison for Lots 312140-1 (40% RG502H), 312140-2 (40% RG502), 312140-3 (20% RG502H, 20% RG502) and 312140-4 (10% RG502H, 30% RG502). All Formulations are 60% Dexamethasone

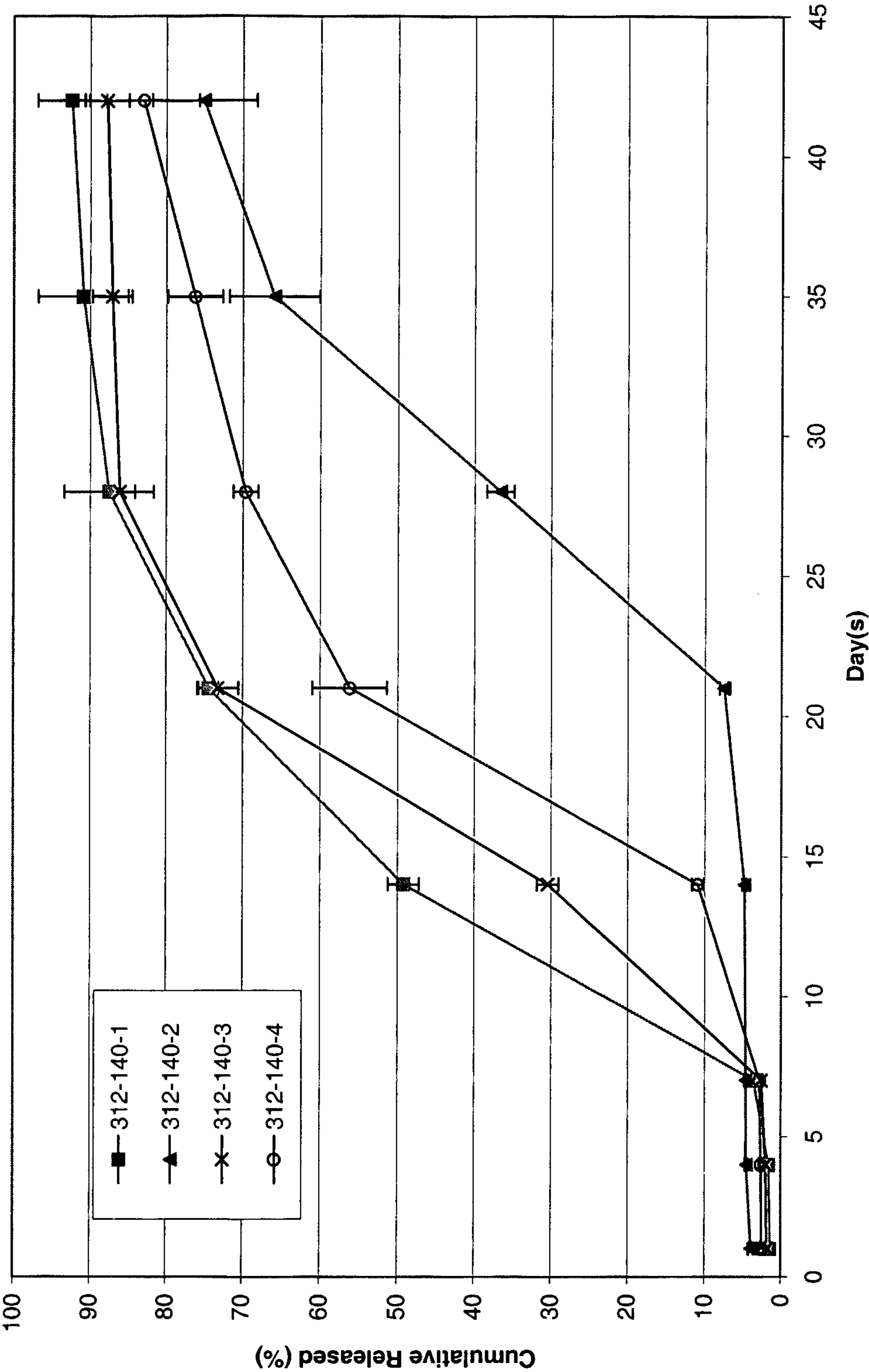


FIGURE 10

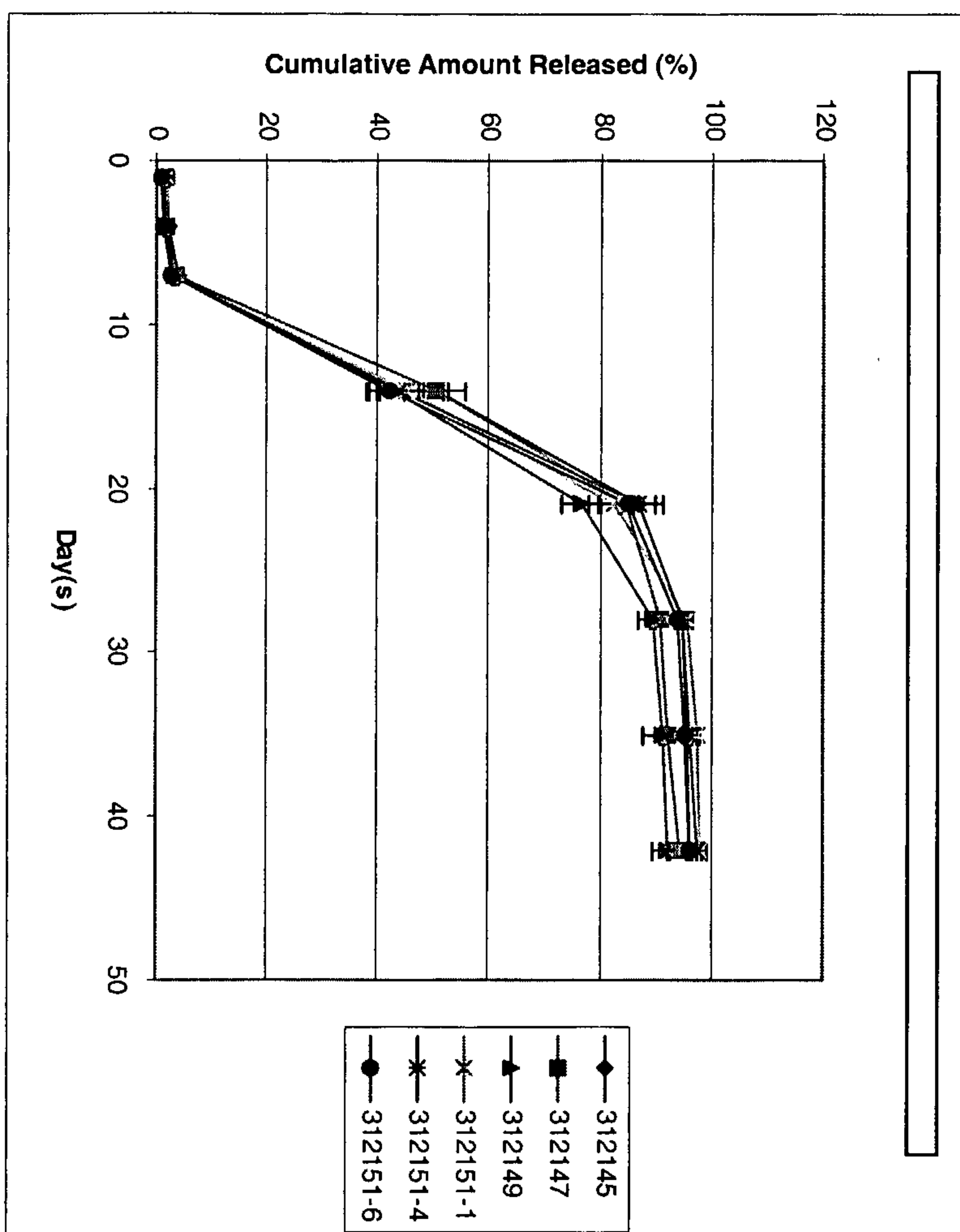


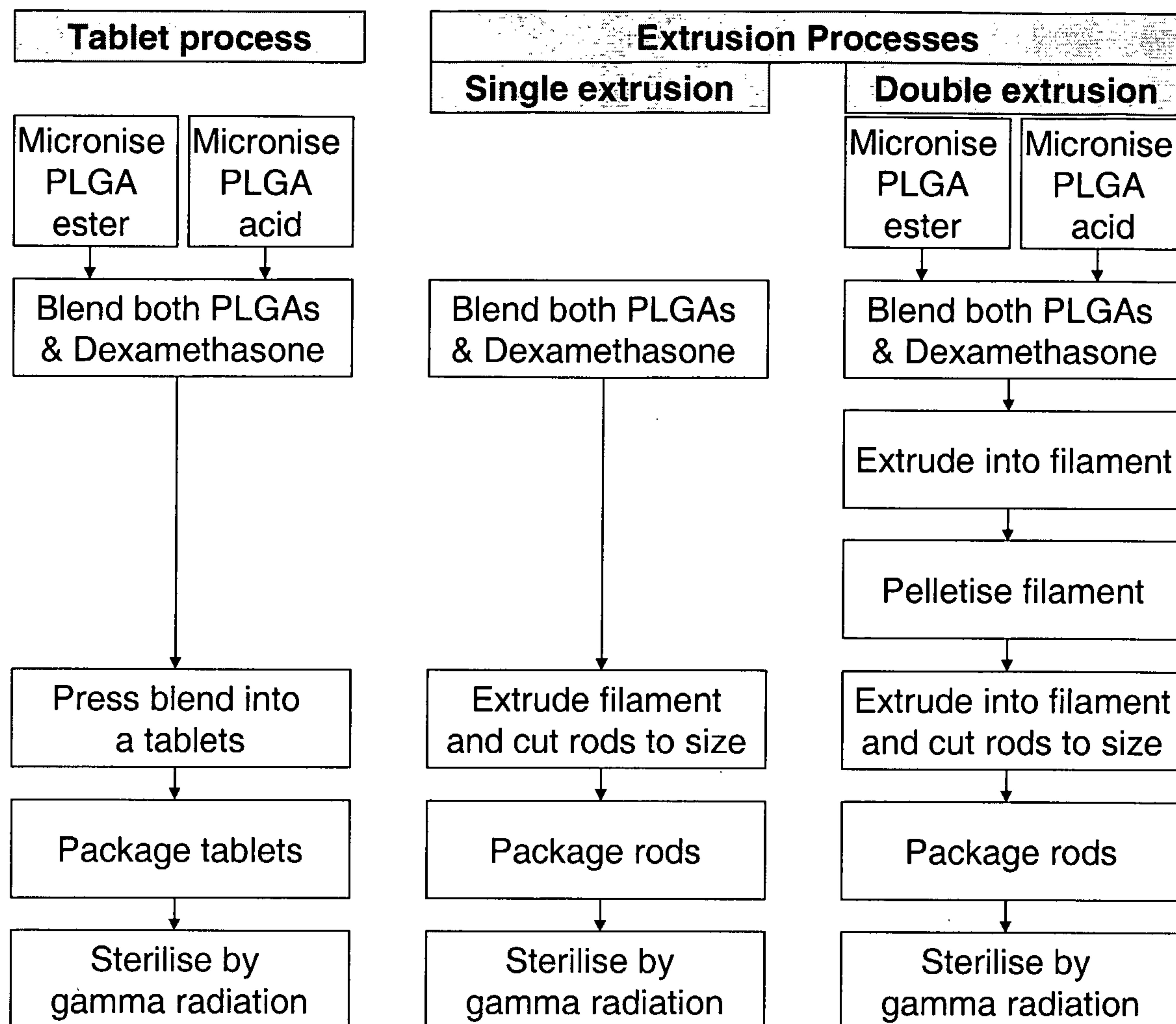
Figure 11 **Processes used for the manufacture of the DEX PS DDS**

Figure 12 *In vitro* dexamethasone release rate comparison of two Lots of tableted and single extruded DEX PS DDS

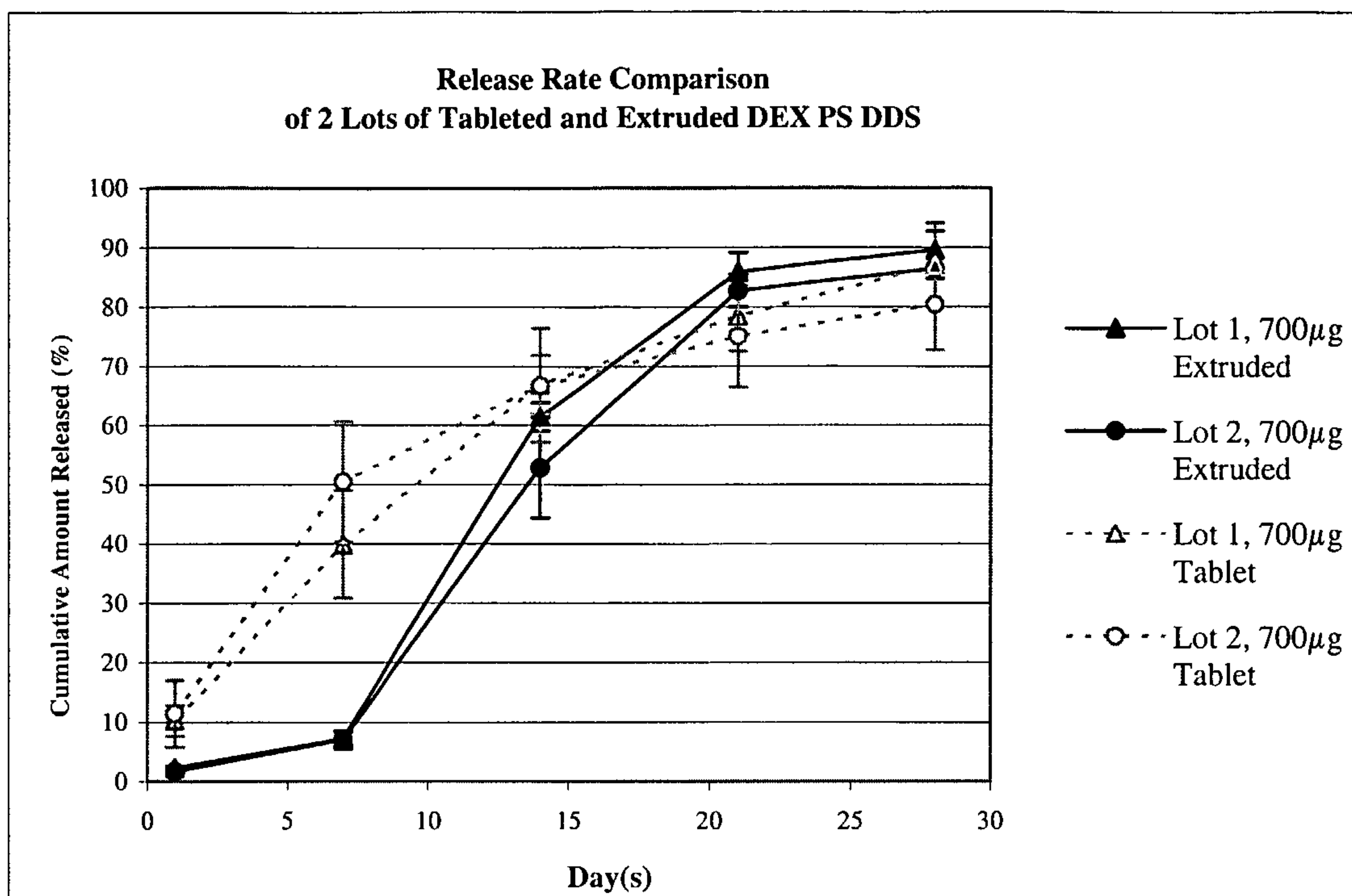


Figure 13. SEM pictures of tabletted and extruded DEX PS DDS

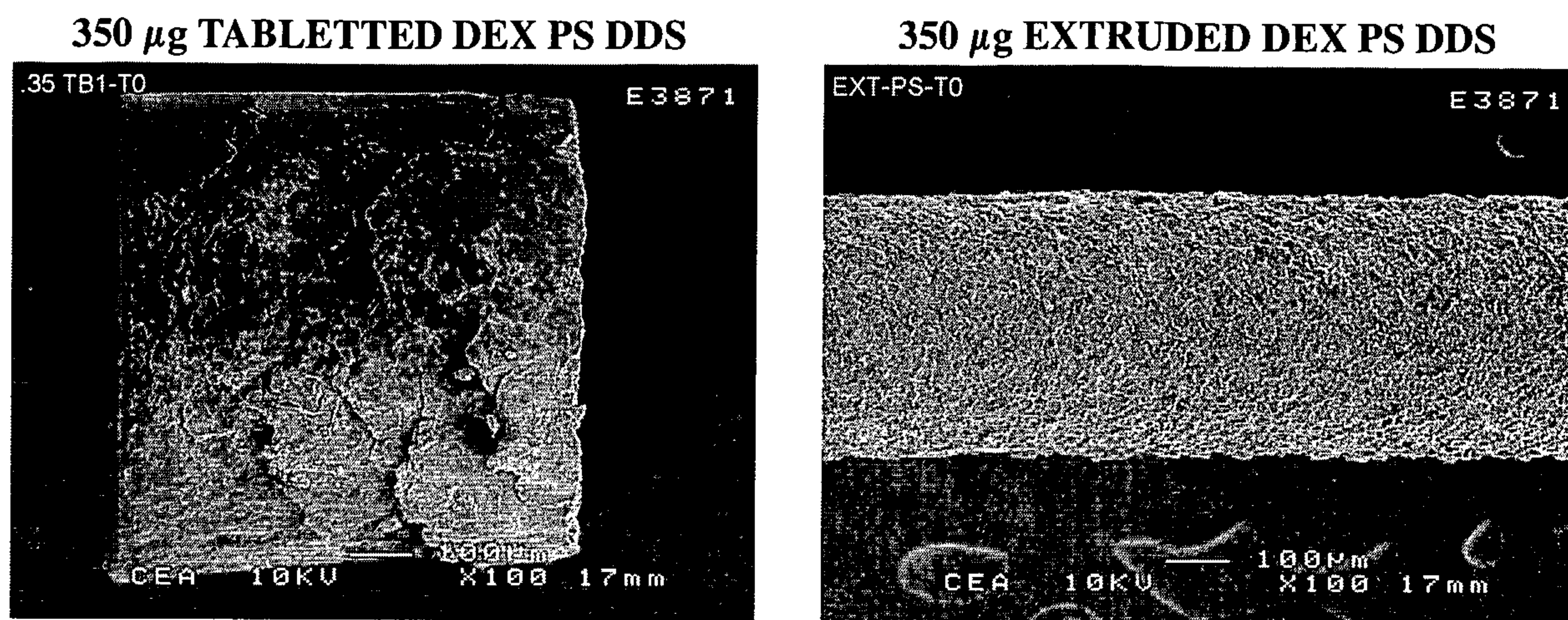


Figure 14 **Content uniformity in DEX PS DDS after extrusion from batches made of milled and un-milled PLGAs**

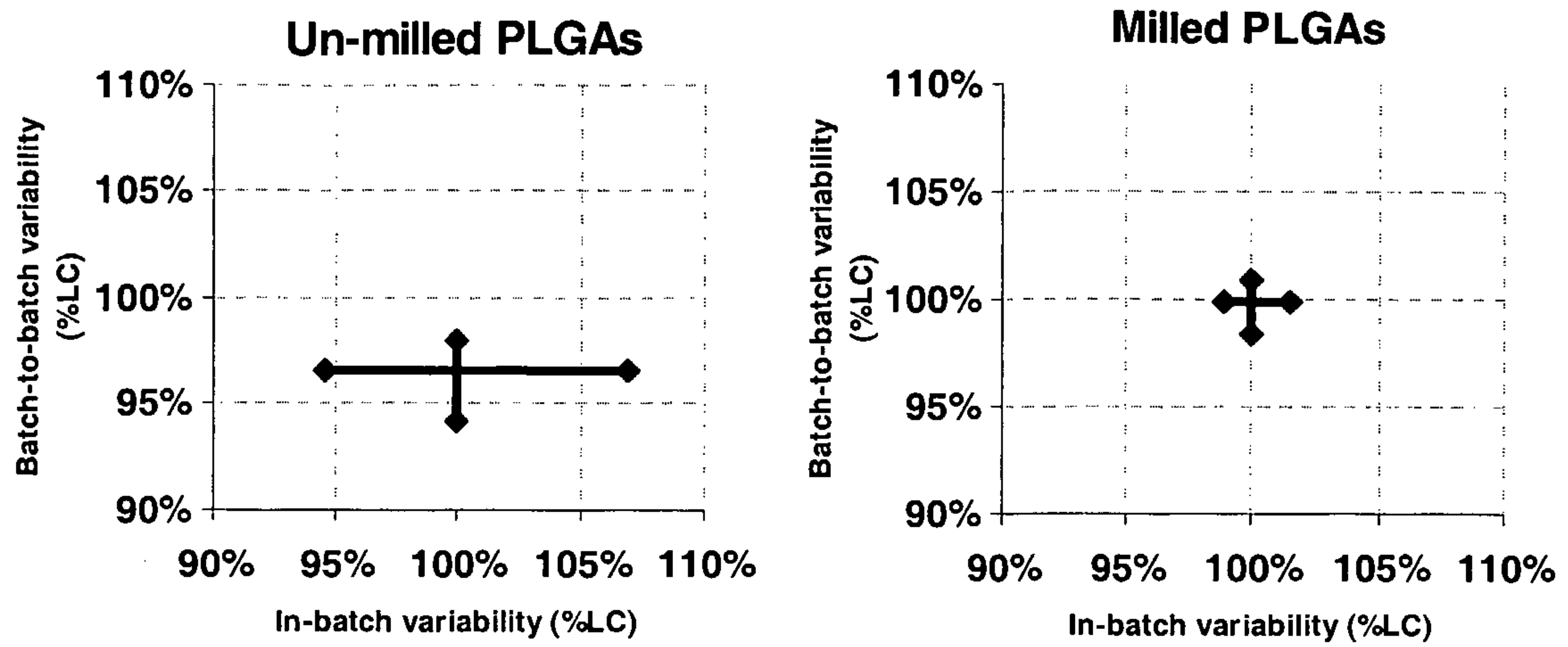


Figure 15

In vitro Release profile of DEX PS DDS manufactured by single and double extrusion.

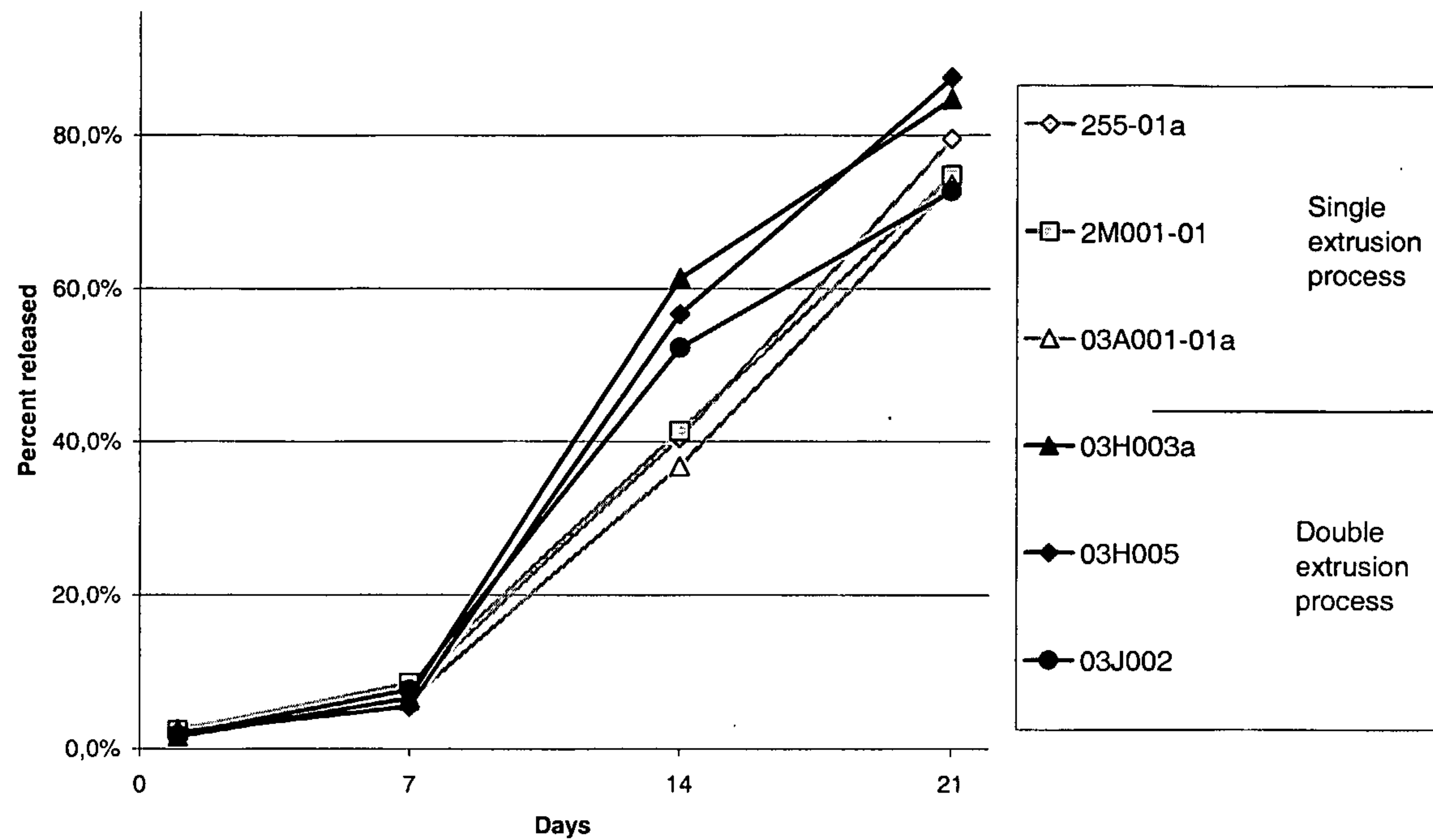
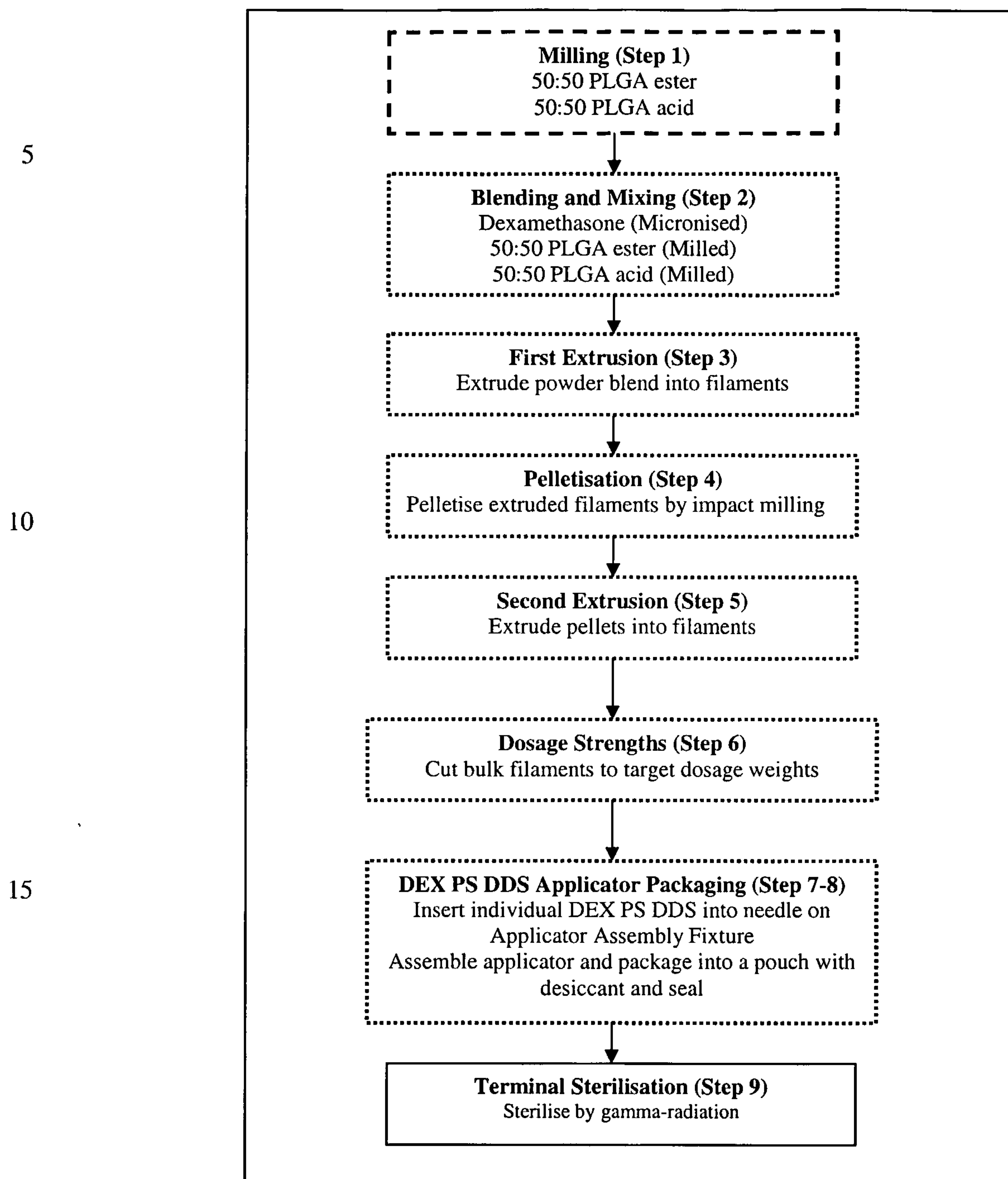


Figure 16

Flow diagram of double extrusion process for the manufacture of the DEX PS DDS

Legend

Step in clean room: cabinet in class 10 000 Class 1 30 Not in clean room

Figure 17 **Cut-Away View of the DEX PS DDS Applicator System**

