Phase 1B study of the safety and tolerability of the mineralocorticoid fludrocortisone acetate in patients with geographical atrophy

Thomas Hong,¹ Andrew Chang,¹,² Ted Maddess,³ Jan Provis,⁴,⁵ Philip Penfold⁶,⁷

ABSTRACT

Objective To evaluate the safety and tolerability of a mineralocorticoid, in a single-dose intravitreal (IVT) injection of 1 mg/0.1 mL and 2 mg/0.1 mL fludrocortisone acetate (FCA) in subjects with geographical atrophy (GA) secondary to age-related macular degeneration.

Methods and Analysis This phase 1b study was a two-part dose-escalation prospective study. Part 1 involved a single participant treated with 1 mg/0.1 mL and monitored up to 28 days before being reviewed by a safety review committee. Two subsequent participants were then dosed with the same dose. Part 2 involved a single participant dosed with 2 mg/0.1 mL and monitored up to 28 days when a further five participants were dosed. All participants were followed up for 6 months after baseline. A full ophthalmic assessment was performed at study visits which included GA area, best-corrected visual acuity (BCV), low-luminance BCV (LL-BCV) and intraocular pressure (IOP). Adverse events (AEs) were reported from the first dose of FCA until the end-of-study visit.

Results There were no serious AEs (ocular or systemic) observed with IVT FCA at either 1 mg/0.1 mL or 2 mg/0.1 mL among nine participants. There was no evidence of increased IOP or cataract development. Neither BCV or LL-BCV changed significantly in the study-eye over the follow-up period (p=0.28 and 0.38, respectively). Mean GA area increased in the study (0.5 mm², p=0.003) and fellow-eyes (0.62 mm², p=0.02) over 6 months. Differences between eyes were not significant (p=0.64), and at the lower end of population norms.

Conclusion IVT FCA is clinically safe and well tolerated and did not increase IOP.

INTRODUCTION

Steroids are commonly used to treat retinal disease such as diabetic macular oedema and retinal vein occlusions as either sole or combination therapy. These steroids have mostly been glucocorticoids which have been shown to have side effects including a raised intraocular pressure (IOP) and lens opacification.

Fludrocortisone acetate (FCA) (9α-fluoro-11β, 17α, 21-trihydroxy-4-pregnene-3,20-dione acetate, FCA) is a synthetic steroid possessing a potent mineralocorticoid effect and high glucocorticoid activity, and so has anti-inflammatory and antiexudative properties. FCA is a mineralocorticoid receptor and glucocorticoid receptor agonist that binds to cytoplasmic receptors, translocates to the nucleus and subsequently initiates the transcription of glucocorticoid-responsive genes such as lipocortins to inhibit phospholipase A2. This prevents the release of arachidonic acid, a precursor to prostaglandins and leukotrienes, both important mediators in the proinflammatory response mechanism. In addition, this agent exerts its mineralocorticoid effect on the distal tubules and...
collecting ducts of the kidney by inducing permease, an enzyme that regulates Na+ permeability in cells, thereby enhancing Na+ reabsorption and water retention as well as increasing K+, H+ excretion.

Previous preclinical studies have indicated that FCA showed neuroprotective properties in mice, with a preservation of the outer nuclear layer in the retina of mice with induced photo-oxidative damage compared with triamcinolone acetonide and control eyes. Furthermore, the a-wave and b-wave response on electroretinography was higher than comparator arms suggesting preservation of photoreceptors. Furthermore, a study in rabbits found that intravitreal (IVT) FCA did not increase IOP.

Age-related macular degeneration (AMD) is the leading cause of severe vision loss in people over the age of 65 in Western countries. In the USA, about 1.75 million people have the advanced forms of AMD. The early signs of AMD (drusen and pigmentary changes) are common in individuals over age 65 and precede the advanced forms, which are visually devastating. The advanced forms of AMD are classified into either choroidal neovascularisation (CNV) (wet or exudative) or geographical atrophy (GA) (dry).

GA is a disease characterised by thinning and loss of the retinal pigment epithelium (RPE), and concurrent atrophy of photoreceptors and choriocapillaris. Clinically, GA is characterised by islands of dead retinal cells in the back of the eye that gradually expand. Although GA can result in significant visual function deficits in reading, night vision, and dark adaptation, and produce dense, irreversible scotomas in the visual field, the initial decline in VA may be relatively limited if the fovea is spared. When the fovea is involved, GA quickly causes blindness. GA is responsible for approximately 20% of all legal cases of blindness in North America with increasing incidence and prevalence owing to a higher life expectancy.

AMD is a highly complex disease that is affected by multiple factors, such as ageing, genetic predisposition, environmental elements, oxidative stress and inflammatory effects. Smoking, age, alcohol consumption, diet and obesity are important risk factors related to oxidative stress. High body mass index, cardiovascular disease, hypertension and a variety of dietary patterns are risk factors less consistently. Several single-nucleotide polymorphisms that confer increased or decreased risk of inflammation have been identified. They include the well-recognised complements factor H, CX3CR1, Toll-like receptor 3 (TLR3), TLR4 and interleukin 8 (IL-8).

Although AMD is not a classic inflammatory disease, inflammatory cells have an important role in AMD pathogenesis and progression. Evidence has also suggested that some infectious agents are associated with AMD. Interleukin 6 (IL-6) has also been found to be upregulated in neovascular AMD and GA, furthermore it has been linked to GA progression.

The purpose of this study was to assess the safety and tolerability of a single IVT dose of FCA among patients with GA over a 6-month period.

METHODS
Study design
This single-centre, phase 1b prospective, open-labelled, single-dose, dose-escalation clinical trial was conducted on nine participants enrolled at a single site between August 2019 and April 2021 at an eye clinic in Sydney, Australia. All patients were followed up for 6 months after baseline. Ethics approval was obtained before commencement and the trial was listed on the Australian and New Zealand Clinical Trial Registry (accessible via www.anzctr.org.au; ANZCTR no. 12618001308280). An independent data and safety committee provided oversight of the clinical trial. This study adhered to the tenets of the Declaration of Helsinki. Informed consent was obtained from all participants before enrolment into the study.

Study medication
FCA was formulated for IVT administration as a powder solution for injection to ensure long-term stability similar to other corticosteroids. Vials contained 10 mg of FCA powder and were reconstituted with sterile sodium chloride solution (0.9%) according to the appropriate dosage prior to injection.

Study population
Inclusion criteria referring the study eye were as follows: diagnosis of GA secondary to AMD confirmed using fundus autofluorescence (FAF) imaging, GA area between ≥1.9 and ≤17 mm² (1-disc and 7-disc areas, respectively), best-corrected visual acuity (BCVA) of 24 letters or better using Early Treatment Diabetic Retinopathy Study (ETDRS) charts.

Exclusion criteria were as follows: GA due to causes other than AMD such as Stargardt disease, cone rod dystrophy or toxic maculopathies like plaqueenil maculopathy, spherical equivalent of the refractive error demonstrating >6 dioptres of myopia or an axial length of ≥26 mm, evidence or history of exudative (wet) AMD including evidence of RPE rips or evidence of neovascularisation anywhere in the retina based on fluorescein angiogram in either eye within 12 months, retinal disease likely to confound visual performance or be affected by intraocular steroid, intraocular surgery (including lens replacement surgery) within 3 months prior to dosing, aphakia or absence of the posterior capsule, previous violation of the posterior capsule is also excluded unless it occurred as a result of yttrium aluminium garnet laser posterior capsulotomy in association with prior posterior chamber intraocular lens implantation and at least 60 days prior to day 0, glaucoma or family history of glaucoma, any contraindication of IVT injection including current ocular or periocular infection, history of uveitis.
of IVT injection within 12 months.

If both eyes met the criteria, the eye with the best VA at the screening visit was designated as the study eye.

**Study protocol**
A full ophthalmic assessment was performed at each study visit which included GA area assessed through FAF imaging, BCVA, lower-luminance BCVA (LL-BCVA) and IOP. Participants were assessed at screening, baseline, day 1, day 7, day 14, day 28, day 60, day 90 and day 150 (end of study). Blood and urine samples were collected for safety analysis at screening, baseline, day 7, day 28 and day 150.

Part 1 of the study involved a single participant treated with 1 mg/0.1 mL FCA to assess safety and tolerability. This participant was followed up for 28 days before the results were reviewed by an independent data safety monitoring committee (DSMB). Subsequent to approval by the DSMB, a further two participants were treated with 1 mg/0.1 mL FCA and followed up for a further 28 days after treatment before commencement of part 2.

Part 2 involved a single dose of 2 mg/0.1 mL of FCA in a single participant and followed up for 28 days like part 1. The DSMB reviewed the results prior to enrolment of the remaining five participants.

**Outcome measures**
Fundus autofluorescence was captured using Heidelberg Spectralis (Heidelberg Engineering, Heidelberg, Germany). The assessment of GA size and progression was performed using FAF was performed by two graders (TH and AC) in a blinded manner using Heidelberg region finder software V.2.6.2.0 (figure 1), a semiautomated programme used to quantify atrophic areas. Baseline FAF images were defined and used to assess subsequent visits using the region finder software. In cases where there was a discrepancy of more than 20% between the two graders, a third grader (AC) evaluated the images. Areas of peripapillary atrophy were not included in the measurements. GA was defined as well-demarcated regions of hypofluorescence on FAF from the absence of the RPE layer over the neurosensory retina.

BCVA was assessed at every study visit at 4 m using an ETDRS chart following subjective refraction. LL-BCVA was assessed similarly with a neutral density lens. IOP was assessed with Goldmann applanation tonometry. Adverse events (AEs) were reported from the first dose of FCA in the first patient until the last patient last visit.

**Statistical analysis**
Statistical analysis was performed using SPSS software V.24.0 (SPSS) and are primarily descriptive. Summaries of safety data are presented in the results. Descriptive statistics (mean, SD, median, minimum and maximum) are calculated for summaries of continuous data. Paired t-tests were performed to assess change from baseline and two sampled t-tests were performed to compare change between eyes. As this was a phase 1B study, there was no formal sample size calculation.

Safety data, including vital signs, clinical safety labs and AEs, will be summarised. AEs were coded using the medical dictionary for regulatory activities, and data will be summarised by system organ class and preferred term.

![Figure 1](example.png)  
**Figure 1**  Example of atrophy as measured by Heidelberg region finder software (A) baseline multicolour, (B) baseline fluorescein angiogram, (C) baseline autofluorescence, (D) baseline region finder, (E) month 1, (F) month 2, (G) month 3 and (H) month 6.
RESULTS

Nine participants were enrolled in the study, baseline characteristics are presented in table 1. The mean age of participants was 79.7±6.2 years and 5 of 9 were female. Mean baseline BCVA and LL-BCVA were 53.1±10.0 and 39.3±11.2 letters, respectively. Mean baseline area of GA was 9.50±5.7 mm².

The initial pilot participant dosed with 1 mg/mL FCA experienced a loss of 22 letters (≥15 letters) at day 28, however this was deemed most likely unrelated to the IP by the DSMB as vision improved (+9 letters) on subsequent visit (day 60) and there were no signs of any other AEs or safety concerns.

One participant dosed with 2 mg/mL FCA experienced a nasal subconjunctival haemorrhage after day 90 in the study eye that resolved after 1 week. No other AEs were noted throughout the study.

No significant increases in IOP (≥10 mm Hg) were observed among any patients throughout the study. The mean change in IOP was −0.25 mm Hg (p=0.75) at day 150 compared with baseline (table 2). There were no differences in progression rate among study eyes dosed with 1 mg or 2 mg FCA (p=0.41).

Progression of GA area

The mean area of GA increased in both the study (0.5 mm², p=0.003) and fellow (0.62 mm², p=0.02) eyes over the duration of the study. This translates to a mean reduction in progression rate of 19.9% over the duration of the study. The change in area was not significant between eyes (p=0.64) (table 2). There were no differences in progression rate among study eyes dosed with 1 mg or 2 mg FCA (p=0.41).

DISCUSSION

This study shows that IVT FCA is clinically safe and well tolerated among this cohort of patients with GA secondary to AMD. Raised IOP is a typical side effect of intraocular steroids and was not observed in this study. Increased lens opacity is another typical side effect, however, no participants required cataract surgery during the study. Furthermore, no systemic AE was observed during this does escalation study.

A recent meta-analysis of 23 studies reported the natural progression of GA lesions in untreated eyes to be 1.66 mm²/year. Results from the Proxima A and B clinical trials reported a growth rate of 2.09 and 1.90 mm²/year, respectively, over the first 12 months. Another study by Schmitz-Valckenberg et al. reported a change of 0.88 mm² over 6 months. Our study findings indicate that GA progression was lower in both the treated (0.5 mm²) and fellow eyes (0.62 mm²) over the study period, suggesting a possible treatment effect of FCA. Both the GATHER one and Filly studies have reported a 20%–30% reduction the rate of GA progression over 12 months, our findings report a similar rate of 19.9%.

Previous studies assessing the efficacy of complement inhibitors in reducing GA progression have found mild to moderate rates of conversion of GA to CNV. The GATHER one study (Zimura) reported CNV developing in 3.5% of study eyes and 2.7% of controls over a 12-month period. The Filly study assess the safety and efficacy of Pegcetaplan and reported conversion to CNV of 20.9% in the monthly arm, 8.9% in the bimonthly arm and 1.2% in the control arm over 12 months. No eyes developed CNV during the study period. No participants developed CNV during the study.

Table 1 Baseline characteristics

| Baseline characteristic | N=9 |
|-------------------------|-----|
| Age (years) mean±SD     | 79.73±6.19 |
| Female                  | 55.5% |
| Right eye               | 55.5% |
| Pseudophakic            | 55.5% |
| BCVA (letters) mean±SD  | 53.11±10.01 |
| LLVA (letters) mean±SD  | 39.33±11.21 |
| IOP Mean±SD             | 13.33±3.08 |
| GA area (mm²) mean±SD   | 9.53±5.69 |

BCVA, best-corrected visual acuity; GA, geographical atrophy; IOP, intraocular pressure; LLVA, low-luminance visual acuity.

Table 2 Changes in characteristics from baseline

| Variable           | Study eye | Fellow eye | P value | P value between eyes |
|--------------------|-----------|------------|---------|----------------------|
| BCVA (letters)     | −2.63±7.01| 4.88±8.37  | 0.28    | 0.69                 |
| LLVA (letters)     | 3.25±9.23 | 4.5±11.49  | 0.38    | 0.72                 |
| IOP (mm Hg)        | −0.25±4.06| −0.625±4.24| 0.75    | 0.69                 |
| GA area (mm²)      | 0.5±5.69  | 0.62±4.49  | 0.003   | 0.02                 |

BCVA, best-corrected visual acuity; GA, geographical atrophy; IOP, intraocular pressure; LLVA, low-luminance visual acuity.
in this study, however, participants were only followed-up for 150 days as part of the study.

LLVA is a known predictor of visual loss in patients with GA, where it can detect a change in central function earlier than standard VA assessments.23–27 Our findings show that LLVA did not deteriorate over the study period may be considered promising, and worthy of a more detailed study to be conducted in a larger cohort.

Previous studies assessing IVT glucocorticoid treatments have found improvements in the function of the blood retinal barrier.28–30 These findings are consistent with preclinical investigations into the use of FCA in vitro, and in vivo using a mouse model of AMD.3 In those experiments FCA was found to have potent anti-inflammatory effects and to be neuroprotective in the AMD model.

Clinical studies have previously reported previously that monocular treatments can affect fellow eyes,30–32 meaning that comparison of measures between treated and fellow eyes cannot serve as valid controls, and that population norms are the most useful comparators. Given that both the treated and fellow eyes showed a slower progression rate compared with previous reports, there may also be a similar sympathetic response in the fellow untreated eye.

The limitations of this study are the small sample size and the absence of formal lens grading, both of which were considered non-essential to the primary objective of this safety study. Another limitation of the study is its short duration. The strength of this study is its prospective design.

In summary, the data show that IVT FCA is clinically safe and well-tolerated among this cohort. Treatment with FCA did not increase IOP, further longer-term studies with larger sample size and multidose regimens may aid in assessing the efficacy of FCA in reducing GA progression.

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Patient consent for publication Consent obtained directly from patient(s)

Ethics approval This study involves human participants and was approved by Bellberry (2018-08-709). Participants gave informed consent to participate in the study before taking part.

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