Visual performance in giant cell arteritis (temporal arteritis) after 1 year of therapy

Mark J Kupersmith, Regina Langer, Hal Mitnick, Robert Spiera, Harry Spiera, Marjorie Richmond, Stephen Paget

Abstract

Aims—To determine if patients with giant cell arteritis (GCA) treated with corticosteroids develop delayed visual loss or drug related ocular complications.

Methods—In a multicentre prospective study patients with GCA (using precise diagnostic criteria) had ophthalmic evaluations at predetermined intervals up to 1 year. The dose of corticosteroid was determined by treating physicians, often outside the study, with the daily dose reduced to the equivalent of 30–40 mg of prednisone within 5 weeks. Subsequently, treatment guidelines suggested that the dose be reduced as tolerated or the patient was withdrawn from steroids in a period not less than 6 months.

Results—At presentation, of the 22 patients enrolled, seven patients had nine eyes with ischaemic injury. Four eyes had improved visual acuity by two lines or more within 1 month of starting corticosteroids. No patients developed late visual loss as the steroid dose was reduced. At 1 year the visual acuity, contrast sensitivity, colour vision, and threshold perimetry were not significantly different from the 4–5 week determinations. At 1 year, there were no significant cataractous or glaucomatous changes. At 2 months, there was no difference in systemic complications between patients who received conventional dose (60–80 mg per day) or very high doses (200–1000 mg per day) of corticosteroids at the start or early in the course.

Conclusions—Patients with GCA related visual loss can improve with treatment. Corticosteroids with starting doses of 60–1000 mg per day, with reduction to daily doses of 40–50 mg per day given for 4–6 weeks, and gradual dose reduction thereafter, as clinically permitted, did not result in delayed visual loss. There were no significant drug related ophthalmic complications.

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Giant cell arteritis (GCA) is a systemic illness which affects men and women older than 55 years of age (more often older than 65) which can cause severe visual loss in affected individuals unless treated early. The inflammation of the ophthalmic arterial branches leads to ischaemic disease of the orbit with profound optic neuropathy, retinal and choroidal infarction, and less commonly, ocular movement limitation.1,2 Corticosteroid treatment must be started as soon as the diagnosis is suspected since in 10% to 33% of untreated patients, the second eye will lose vision within 1 day to 3 weeks.3 Unfortunately, the required long term (6–24 months) treatment with corticosteroids often leads to severe systemic complications such as osteoporosis, gastric ulcer, diabetes, high blood pressure, immunosuppression, depression, weight gain,4,5 and ocular complications such as cataract and glaucoma.4,6 Preliminary studies have suggested that an anti-inflammatory drug, methotrexate may be beneficial for GCA because its use permits lower doses of corticosteroids to be administered with concomitant reduction in complications.7 However, one prospective study using low dose (7.5 mg/week) methotrexate showed no benefit on GCA or polymyalgia rheumatica.8

We established a clinical trial at three university medical centre rheumatology and neuroophthalmology services to compare conventional corticosteroid therapy with a combination of methotrexate and corticosteroid treatment to address the question of disease control and rate of drug related complications. Difficulties with recruitment precluded sufficient numbers to answer the question whether methotrexate is beneficial but the prospective collection of data has provided some results with respect to the following questions:

(1) Do daily pharmacological doses of corticosteroids given for 1 year cause significant cataract development or elevation of intraocular pressure in the elderly? Can the total cumulative corticosteroid dose be correlated with any ophthalmic complication?

(2) Can corticosteroid therapy improve the vision in an eye with ischaemic injury from GCA?

(3) Does a short term (<7 days) initial daily dose of 200–1000 mg of corticosteroids cause more systemic complications than an initial dose of 40–60 mg?

(4) Do patients with GCA experience visual loss vision at 1 year while on small doses or no corticosteroid?

(5) Do the eyes of patients with GCA have a poorer vision at 1 year compared with the visual performance after 4–6 weeks of treatment?

Methods

Twenty two patients aged over 55 years with GCA who met criteria modified from the American College of Rheumatologists14 (Table 1)
Table 1 Criteria for diagnosis of giant cell arteritis (GCA)

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<th>Criteria</th>
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<td>(1) Patients must have symptoms suggestive of GCA and a temporal artery biopsy (performed within 1 week of steroid initiation), with signs of panarteritis including chronic inflammatory cells, disruption of the internal elastic lamina, with or without giant cells, except for (2) or (3).</td>
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<tr>
<td>(2) Patients with acute anterior or posterior ischaemic optic neuropathy with symptoms suggestive of GCA (polymyalgia rheumatica, temporal fossa headache, scalp tenderness, loss of appetite, jaw claudication, tongue claudication, weight loss, night sweats), and ESR &gt;60 mm in the first hour (without other cause for elevated ESR—that is, blood dyscrasia, lymphoma, tuberculosis, renal failure) even if negative temporal artery biopsy.</td>
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<tr>
<td>(3) Patients with symptoms suggestive of GCA and pulseless disease with ESR 60 mm in the first hour.</td>
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for diagnosis, did not have factors that excluded them, and gave consent were entered into a randomised, prospective, placebo controlled study within 5 weeks of establishing the diagnosis. Patients were excluded if they had additional visual or systemic illnesses that contraindicated the use of corticosteroids or methotrexate or could complicate or bias the study (in fact no patients meeting criteria and giving consent were excluded).

Patients with symptoms or signs of ophthalmic involvement were examined by MJY immediately before or after the onset of visual disturbance. Baseline and all subsequent evaluations (1, 3, 6, and 12 months) included full ophthalmological evaluation with best corrected visual acuity testing using Bailey-Lovie logMAR chart under standard conditions, contrast sensitivity testing using Pelli–Robson charts under standard conditions,15 colour vision testing with Ishihara plates, applanation tonometry, slit lamp biomicroscopy after miosis, determination of relative afferent pupillary defect, dilated fundus examination with stereoscopic measurement of the horizontal and vertical cup/disc ratio, and visual field testing with the Humphrey 24-2 strategy. For each eye, the visual acuity was expressed as decimal (20/20 = 1.0, finger counting = 0.012, hand movement = 0.006, light perception = 0.001, no light perception = 0), the lowest contrast seen in log units of contrast, the visual field threshold as mean deviation (MD) and pattern standard deviation (PSD) <5%. The visual field data were also scored using the advanced glaucoma intervention study protocol (AGIS score).16

The degree of opacification in the lens was measured using a modification of LOCSII.17 18 Following pupillary dilatation >5 mm, with tropicamide 1%, the lens was examined and the degree of opacification in the nucleus, cortex, and posterior subcapsular zones was graded. Opacities were defined as being present only if they were visible against the red reflex, otherwise the grade was 0. “Aggregate” opacification was derived by mentally summing the area of the opacities from non-contiguous locations in one zone and determining if the sum was as large as an entire quadrant (grade 2) or just less than the quadrant, or a small, minor cluster (grade 1a). A separate grade was recorded for the cortical and the posterior subcapsular zones. Nuclear opacification was determined using the slit beam slightly wider than the pupil and simply graded as clear or unclear.

All patients received daily corticosteroid therapy (range 40–1000 mg) after initial diagnosis. The starting dose was left to the clinical judgment of the referring treating physician. Over 4–6 weeks the daily steroid dose was reduced to the equivalence of 30–40 mg of prednisone at which time each patient was randomised to receive either methotrexate or placebo in addition to the steroids. Since not all the rheumatologists clinically managing the patients were study investigators, the following regimen for the prednisone therapy was only suggested:

*The dose of prednisone was to be gradually decreased by 10 mg/week until 40 mg/week is reached by the end of the first month, then by 5 mg/week until 20 mg/week is attained at the end of the second month. Subsequent tapering should be by 2.5 mg per week until the drug is completely suspended. The prednisone taper should be halted or the dose increased by 5–10 mg/day or more in order to control the symptoms or signs if a relapse occurs. A relapse is defined as recurrence of symptoms of GCA after definite improvement by prednisone, followed by symptom normalisation upon resumption or increase of the prednisone dose. An increase in the erythrocyte sedimentation rate (ESR) alone is not used as the indication for increasing the steroid dose.*

After 4–6 weeks of corticosteroids, patients were randomised to either the addition of oral placebo or methotrexate of 10 mg per week. At 1 year, if a remission was achieved, the methotrexate dose was to be reduced by 2.5 mg/month until it is completely withdrawn.

All patients received 1500 mg/day of calcium carbonate and 400 IU of vitamin D to prevent osteoporosis, folic acid 1 mg orally each day, a H2 blocking agent or carafate 1 g daily to prevent gastrointestinal ulceration. No patient received other anti-inflammatory drugs, sulfa drugs, or other immunosuppressive agents. When necessary the appropriate agent was added to control blood pressure, blood glucose, or prevent worsening osteoporosis.

**DATA ANALYSIS**

In the patients with visual loss, the visual acuity at presentation was compared with the visual acuity at entry into the study which was after 4–6 weeks of corticosteroid therapy.

The mean and standard deviations were determined for all the visual measures at baseline and at 1 year and Student’s test was used to determine significance. For each, visual variable values at baseline were subtracted from the 1 year values. Linear regression analysis was used to demonstrate correlation between total cumulative steroid dose and change in lens, total cumulative steroid dose and change in contrast sensitivity, change in contrast sensitivity and change in MD and PSD, and change in contrast and change in lens. Eyes with synthetic intraocular lens were omitted from lens related analyses (six eyes of four patients). Two eyes of one patient with...
known chronic open angle glaucoma were included in all analyses. One eye (not clinically affected by GCA) of one patient with a large macular scar from childhood was eliminated from all analyses of visual performance. Owing to patient non-compliance, there was no repeat visual field data at 1 year for her two eyes.

Results

Seven men and 15 women with a mean age 73 years (SD 7 years) were enrolled. The mean ESR was 70 mm in the first hour (SD 39 mm). Histopathological examination revealed a positive temporal artery biopsy in 19 patients and a negative result in three patients. Owing to the nature of referrals to the study by both a neuro-ophthalmologist and rheumatologists and the various modes of clinical presentation, the initial steroid dose was not uniform and patients received daily doses of corticosteroids of 40–1000 mg. The initial daily corticosteroid dose was prednisone 40 mg in two patients, 60 mg in 14 patients, 80 mg in one patient, 80–120 mg in one patient, 200 mg in three patients, and one patient received methylprednisolone 1000 mg per day. Two patients subsequently were treated with methylprednisolone 1000 mg per day after failing with daily prednisone 60 mg and 100 mg.

Seven patients had nine eyes with permanent visual loss (seven eyes with ischaemic optic neuropathy and two with central retinal artery occlusion). Two patients developed visual loss after prednisone was begun; one had both eyes affected after 3 days of 60 mg and the second had one eye affected after 80 mg for 2 days and 120 mg for 2 days. These two patients and a third patient with acute ischaemic optic neuropathy were treated with intravenous methylprednisolone 1000 mg per day. Three other patients with subacute visual loss, untreated for 1 week before diagnosis, were treated with prednisone 200 mg/day. One additional patient with a mild ischaemic optic neuropathy (20/25 acuity and an altitudinal field defect) received 80 mg of prednisone daily. No patient developed visual loss or worsened after 5 days of therapy.

At the entry examination, of the 44 eyes, the intraocular pressure was elevated above 22 mm Hg in four eyes, two of which were known to have chronic open angle glaucoma (in one patient without GCA caused visual loss). The eyes in this patient had the same intraocular pressure elevation after starting steroids as before. The other two eyes (in the one patient with bilateral visual loss) had pressures of 25 and 26 mm Hg after steroids (not before), which normalised on prophylactic topical β blocker ophthalmic drops. Neither of the latter two eyes had any worsening of the visual field or increase in the cup to disc ratio over 1 year.

Compared with the pre-entry evaluation of patients with visual loss, the visual acuity improved two lines or more in four eyes within 1 month of starting treatment (Fig 1). In one patient, who received intravenous methylprednisolone 1000 mg per day for 3 days, the worse of her two affected eyes improved from finger counting (0.012) to 20/800 (0.025), and the second eye improved from 0.286 to 0.317 (not two lines). One patient treated with intravenous methylprednisolone 1000 mg per day for 5 days had an improvement in one eye from no light perception with an amaurotic pupil at presentation to 0.03 at 1 month which was maintained at 1 year. Two patients who received oral prednisone 200 mg per day for 5 days each had one eye improve from 0.05 to 0.2 and 0.012 to 0.2. A fifth patient, originally thought to have non-arteritic anterior ischaemic optic neuropathy because of the mild visual acuity loss and altitudinal defect, was treated with prednisone 80 mg per day for the first week and improved the visual acuity from 0.8 to 1.0 without a significant change in the visual field.

In general, most eyes of patients, regardless of arteritis related visual loss at the baseline entry examination, did not have a significant change in visual acuity, contrast threshold, mean deviation, or pattern standard deviation of threshold perimetry, AGIS score, or colour vision from baseline to the 1 year evaluation (Table 2, Figs 2 and 3). However, 13 eyes of nine patients were considered normal by AGIS at 1 year in contrast with only six eyes in five patients at baseline. For MD, 18 eyes were improved (mean change 4.7 (SD 2.6)), 12 eyes were worse (mean change −3.3 (2.0)) and 11 were unchanged. For PSD, 12 eyes were improved (mean change −3.4 (1.6)), 10 eyes were worse (mean change 2.8 (1.2)), and 19 eyes were unchanged. There was no correlation with change in the contrast threshold or the change in MD or PSD (Table 3).

Changes in the cortical zone or the posterior subcapsular zone occurred in 15 of the 38 eyes with their own lens. Cortical zone changes of one grade and two grades were noted in six and two eyes, respectively. Posterior subcapsular changes of one grade were noted in 11 eyes and none had two grade changes. There was no correlation with the change in lens and change in contrast threshold, MD, or PSD (Table 3).
Table 3  Comparison of change in visual performance measures from baseline to 1 year for 44 eyes*

<table>
<thead>
<tr>
<th>Measure</th>
<th>r2</th>
<th>p</th>
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<tbody>
<tr>
<td>Change in contrast and mean deviation</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>Change contrast and pattern standard deviation</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>Change contrast and cortical zone</td>
<td>0.02</td>
<td></td>
</tr>
<tr>
<td>Change contrast and posterior subcapsular zone</td>
<td>0.08</td>
<td></td>
</tr>
<tr>
<td>Change in cortical zone and cumulative steroid dose</td>
<td>0.008</td>
<td></td>
</tr>
<tr>
<td>Change in posterior subcapsular zone and cumulative steroid dose</td>
<td>0.02</td>
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*All eyes except as noted in methods.

Discussion

As described in earlier studies, patients with GCA associated visual loss can recover with corticosteroid treatment. In our patients, the improvement in acuity was noted within weeks of starting therapy and never after 1 month. One patient with acute (less than 2 hours) loss of vision and no light perception in an eye with an amaurotic pupil improved after receiving methylprednisolone 1000 mg daily. Of the four eyes that improved two lines or more on the visual acuity chart, only one received methylprednisolone 1000 mg/day. The others were treated with oral prednisone 200 mg daily.

All measures of visual performance after 1 year of therapy in patients with giant cell arteritis were fairly stable when compared with the measurements taken at approximately 4–5 weeks after initiating corticosteroid therapy (entry point into treatment trial). The more sensitive measures of vision, contrast sensitivity and threshold perimetry, showed no significant pattern of change and, in fact, in many eyes the improvement or worsening on one test was not paralleled by an improved result on another visual performance measure. However, because of fixation losses and false negative and positive errors, the results of threshold perimetry were probably less reliable than the other visual studies.

Patients can be given corticosteroids 200–1000 mg per day for a week without severe long term ocular or short term unmanageable systemic complications. Prednisone 40–80 mg daily for the next 3–5 weeks followed by a judicious dose reduction over months also appeared to be tolerated relatively well. In contrast, the high rate of complications in other studies probably resulted from protracted use...
of high dose steroids. The fact that in our study gradual reduction to 40 mg daily after 4–5 weeks was not associated with the onset of new visual loss suggests it is probably unnecessary to maintain patients on larger doses of prednisone for months. We are in the process of determining whether initial treatment with a short course of corticosteroids 200–1000 mg per day causes more late systemic complications such as hypertension, diabetes, osteoporosis, and pathological bone fractures than initial daily doses of 40–80 mg. Although it appears that megadose (1000 mg) intravenous or 200 mg orally can be taken by the elderly without more complications than conventional lower dose corticosteroid, the optimal dose to possibly improve vision or prevent visual loss in the second eye has not been established by this or any other study. Anecdotal cases of visual loss developing after 1 week of up to prednisone 200 mg daily which reversed with 1000 mg daily have been described. Since treating patients with high doses of corticosteroid for the first 1–2 weeks seems unlikely to increase the incidence of complications, until it is established that lower doses are as effective, higher dose (200–1000 mg per day) therapy seems warranted in patients with visual loss and maybe even in those who have not experienced neurological or ophthalmological symptoms. There appear to be few reasons not to use 200 mg or more per day in cases with acute visual loss or in cases that are worsening while on conventional moderately high dose oral steroids (60–80 mg). Afterwards, unless there is a definite clinical flare up of GCA, the dose of corticosteroid should be reduced to the equivalent of 40 mg of prednisone within 4–6 weeks. Lowering the daily steroid dose thereafter as clinically tolerated appears to be prudent in that it appeared to be the reason for the reduced number of permanent or long term complications in this study.

We had two patients who developed loss of vision while receiving prednisone 60 mg and 80–120 mg per day, with both showing deterioration in less than 1 week into treatment. Numerous reports have documented the deterioration in less than 1 week into treatment in this study. No cases that are worsening while on conventional lower dose corticosteroid, the optimal dose to possibly improve vision or prevent visual loss in the second eye has not been established by this or any other study. Anecdotal cases of visual loss developing after 1 week of up to prednisone 200 mg daily which reversed with 1000 mg daily have been described. Since treating patients with high doses of corticosteroid for the first 1–2 weeks seems unlikely to increase the incidence of complications, until it is established that lower doses are as effective, higher dose (200–1000 mg per day) therapy seems warranted in patients with visual loss and maybe even in those who have not experienced neurological or ophthalmological symptoms. There appear to be few reasons not to use 200 mg or more per day in cases with acute visual loss or in cases that are worsening while on conventional moderately high dose oral steroids (60–80 mg). Afterwards, unless there is a definite clinical flare up of GCA, the dose of corticosteroid should be reduced to the equivalent of 40 mg of prednisone within 4–6 weeks. Lowering the daily steroid dose thereafter as clinically tolerated appears to be prudent in that it appeared to be the reason for the reduced number of permanent or long term complications in this study.

In contrast, delayed visual loss following treatment for months with 20–30 mg per day of prednisone after initial therapy using 60–80 mg per day is rare. We had no patients who developed ischaemic visual loss after 1 or more months of treatment. Although repeat temporal artery biopsy 6 months to up to 9 years after corticosteroid treatment, may show continued or recurrent inflammatory changes, the risk of visual loss in these patients is unknown. Because the population affected by GCA is also at risk for non-arteritic vascular occlusive disease, reports of central retinal artery occlusion or anterior ischaemic optic neuropathy occurring many months to years after what would be considered adequate treatment may not actually represent recurrence of arteritis. There are numerous reports which contain cases of ischaemic visual loss which appear to be non-arteritic. Unfortunately, some articles have poor documentation of the visual system findings that developed after months of treatment so no definite conclusion can be drawn from many of these reports. We could find three cases with delayed GCA visual loss, two following 2 months and one after 5 months of therapy, but none after 1 or more years of treatment with doses of steroids considered adequate for GCA by most North American rheumatologists.

Lastly, the results in this small study differ from the results of earlier larger studies of longer duration that found significant ocular complications of chronic corticosteroid use. Glaucoma has been reported to develop in approximately 8% of patients on long term systemic therapy. There is no clear correlation with dose or duration of therapy. The new onset of elevation of intraocular pressure occurred in 4% of patients (one case) without the development of glaucomatous cupping. Cataract formation, usually bilateral, is associated with dose and duration of the systemic corticosteroid therapy. The opacification typically begins in the posterior subcapsular region and eventually spreads to the cortex. After 1 year of systemic corticosteroids in our patients older than 55 years of age, significant cataractous changes or visual loss from lens opacification, determined by a reduction in visual acuity or contrast sensitivity, or worsening of the mean deviation of threshold perimetry, were not found. This was not unexpected since the prevalence of cataractous change, particularly posterior subcapsular opacification, appears to increase with duration longer than 1 year, typically longer than 5 years, of systemic corticosteroid use.

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