Retrobulbar pressures measured during surgical decompression of the orbit

A Jan Otto, Leo Koornneef, Maarten Ph Mourits, Loes Deen-van Leeuwen

Abstract

**Aims/background**—In Graves’ ophthalmopathy the increase in volume of intraocular muscles and fat will cause elevated intraorbital pressure. In order to investigate the pressure levels involved, intraorbital pressure, or retrobulbar pressure (RBP) was measured continuously in orbits of patients with Graves’ ophthalmopathy during surgical decompression.

**Methods**—Retrobulbar pressure was measured before and during surgical decompression using an intraorbitally applied pressure transducer.

**Results**—In eight patients with dysthyroid optic neuropathy (DON) RBPs between 17 and 40 mm Hg were recorded. At the end of the surgical procedure the mean RBP was reduced from 28.7 mm Hg to 18.7 mm Hg, the decrease ranging from 8 to 12 mm Hg, which showed a high correlation with the starting pressures (p<0.001). In two cases without DON, pressures were 11 and 9 mm Hg. Forces exerted by spatula manipulation usually resulted in a RBP level of more than 70 mm Hg.

**Conclusions**—This study shows that RBPs are markedly elevated in Graves' ophthalmopathy and that surgical decompression can result in a significant reduction in the intraorbital pressure. Optic nerve dysfunction in Graves' ophthalmopathy may not be caused exclusively by the direct pressure of swollen extraocular muscles upon the optic nerve, but also by a raised RBP. It is hypothesised that the damage inflicted upon the optic nerve can be caused in consequence by RBP induced incarceration of the nerve, compressed by surrounding periosteal lined orbital fat bulging posteriorly into the entrance of the optic canal.

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A discrepancy between the volume of the bony orbit and its content is seen in Graves’ ophthalmomopathy and will probably lead to exophthalmos and visual loss. Protrusion of the eye with displacement of the eyelids and the orbital septum only offer a limited additional space anteriorly due to the rigidity of the connective tissue in septal and suspending ligaments. A relation between intraorbital volume increase and intraorbital or retrobulbar pressure (RBP) has been described earlier in both clinical and experimental settings.1–5

A volume induced increase in orbital pressure has always been considered to be the cause of protrusion (and its clinical consequences), but is usually referred to by ill defined terms such as ‘orbital tissue consistency’, ‘orbital tension’, or ‘orbital rigidity’. Data on ‘tensions’ or ‘orbital compliance’ can be estimated indirectly by measuring the posterior displacement of the globe as the result of a certain force exerted upon the eye by an orbitotonometer or orbitotonograph.6,14 Surgical ‘decompression’, by removing parts of the bony orbital wall, is an intervention which leads to favourable results in cases with optic neuropathy, corneal exposure, or cosmetic complaints.10–21

Earlier studies, investigating volume induced RBP alterations in monkey orbits,22 have shown that reproducible RBP measurements can be performed with a micro pressure transducer introduced intraorbitally. In monkey orbits no differences could be established in pressure levels exploring various sites (intracranially) in the ‘steady state’ (4–9 mm Hg) or after artificially increasing RBP up to 40 mm Hg by means of retrobulbar injection. Since similar experiments cannot be performed in healthy human orbits, we examined RBPs in the orbit of patients with Graves’ ophthalmopathy. Elevated RBPs were noted in patients with dysthyroid optic neuropathy (DON), which could be markedly reduced by surgical intervention.

**Patients and methods**

Nine patients (age range 48–63; four women, five men) with Graves’ ophthalmopathy, in whom surgical intervention was performed, participated in this study. Exophthalmometer readings of the left and right eye showed differences that ranged between 2.5 and 7 mm. Bilateral decompression was performed in six of these cases. Informed consent was obtained from the patients and the protocol of this study was approved by the medical ethics committee of the Faculty of Medicine of the University of Amsterdam.

During surgery, which consisted of a three wall coronal orbital decompression,25–26 RBPs were recorded in 10 orbits of nine patients. Direct RBP measurements were performed by placing a micro pressure transducer catheter (4 F Honeywell MTCR) in the orbit, behind the globe, and connection to a Princeton Applied Research (PAR) amplifier with a chart recorder. The transducer was inserted retrobulbarly through a trocar needle, and was sutured to the skin after withdrawal of the
trocar. In order to investigate the force/pressure relation during surgery the force exerted upon intraorbital tissues by the surgical spatula was measured. The spatula was mounted with strain gauges (stretch sensitive elements) on both sides for registration of the exerted force. All patients were under general anaesthesia during surgical intervention.

Results

Eight patients with bilateral and one with unilateral Graves' ophthalmopathy underwent surgical orbital decompression because of progressive loss of vision or progressive loss of visual field.

In these eight patients showing DON, RBP was measured in one orbit before, during, and at the end of the operation. Two more orbits (the contralateral orbit of patient no 3 and the orbit of patient no 9) were decompressed for rehabilitative reasons.

In the eight orbits operated for DON the RBP values, obtained before surgery, ranged between 17 (patient no 3L) and 40 mm Hg (patient nos 7 and 8) (mean 28.7 mm Hg). Pre- and postoperative intraorbital pressure levels showed a decrease of the RBPs at the end of surgical decompression averaging 10.0 mm Hg (range 9–12 mm Hg) (Fig 1). Before surgery the two orbits without DON showed much lower pressure levels (8.8 mm Hg (patient no 9) and 11.0 mm Hg (patient no 3R)}. Because the transducer was transferred from the right to the left orbit in case 3 no further recordings could be made. After decompression the RBP of patient no 9 did not change significantly.

RBP was measured before and after turning down the bicoronal scalp flap and no significant changes in the intraorbital pressure were noted during this surgical manoeuvre. Lowering of the RBP started as soon as incisions in the periorbit were made.

The evoked RBPs exceeded 70 mm Hg as a result of a mean exerted force of about 275 g by an orbital spatula (Fig 2).

Discussion

Assuming a normal 'steady state' RBP of 3.0–4.5 mm Hg, the experiments performed in the study described here show that all investigated patients with bilateral Graves' ophthalmopathy had a significantly increased tissue pressure in the orbit. Although no data are available for orbital pressures in healthy humans the normal steady state RBP has been calculated to be 3.0–4.5 based on theoretical calculations obtained by assuming a muscle force of 30 g applied to the globe.27–28 No muscle relaxants were administered which could have an effect on the tone of the recti eye muscles and thus on RBP (Norcuron was used instead of succinylcholine or Flaxedil). We postulate that the physiological homeostatic RBP regulatory mechanism, usually active after retrobulbar injection during intraorbital volume increase by balloon filling,18 possibly in the early phase of Graves' ophthalmopathy or tumour growth, may no longer be functioning.

At the end of surgical decompression the RBP already showed a marked reduction in all patients with functional visual impairment. The pressure decrease is highly correlated with the RBPs measured at the start of surgery (Fig 3). Although not necessarily related, most patients lost their sensation of pressure in and around the orbit in the weeks following surgery. We feel that perhaps in the orbit a new pressure equilibrium is re-established, although the mechanism of pressure regulation is not known. In our experience it mostly takes...
It has been found that compression of the vagus nerve in rabbits at 30 mm Hg for 8 hours, induces inhibition of fast and slow anterograde as well as retrograde axonal transport.32-35 Consequently, we frequently observed improvement of the visual function after surgical decompression, even without removal of the bone fragments bordering the 'crowded' region. Among orbital surgeons it is not a standard procedure to remove part of the optic canal in orbital decompression for dysthyroid optic neuropathy. Furthermore, we consider it unlikely that a direct or diffuse pressure rise will be the only cause of impairment of the nerve function and came to the conclusion that secondary incarceration might also be involved. When RBP has reached higher levels than the intracranial pressure (which remains normal), optic nerve encroachment may occur near the optic foremen, caused by orbital tissues, bulging backward inside their lining periosteal membranes (Fig 4). Cadaver studies should be performed to prove the hypothesis mentioned above. In fact, in many coronal computed tomography scans of the orbital apex in Graves' ophthalmopathy a persistent narrow clear ring of low density (probably bulging adipose tissue) can be observed around the optic nerve.36 In this respect, an analogy may be opportune. The optic nerve compression could be comparable with medulla incarceration in the foremen magnum, seen with increased intracranial pressure. It may be concluded from our limited material, that ophthalmopathy with DON is associated with much higher pressures than without DON. Moreover it is evident that an RBP decrease can already be established at the end of the decompression surgery.

The force of 250 to 300 g against the intraorbital tissues exerted intraoperatively by the normally handled spatula resulted in transient RBP levels of over 70 mm Hg. At this pressure, dangerous impairment of the arterial vascularisation of the optic nerve and retina might ensue, especially in cases with pre-existing vascular disorders.

Future studies using less invasive techniques to measure intraorbital pressure may allow an improvement in the timing of decompression surgery to prevent optic neuropathy in these patients.

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