Continual monitoring of intraocular pressure: effect of central venous pressure, respiration, and eye movements on continual recordings of intraocular pressure in the rabbit, dog, and man

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SUMMARY A new method has been devised for continual monitoring of intraocular pressure by radiotelemetry. The use of this instrument for monitoring intraocular pressure in a variety of ophthalmic conditions is described.

Diurnal variation in intraocular pressure (IOP) has been recognised for many years. A practical method for noninvasive monitoring of IOP would be important for the diagnosis and management of several forms of glaucoma, particularly the low tension group (Phelps et al., 1974). Intraoperative and postoperative monitoring may be useful for early warning of acute IOP elevation. Such a system will also provide valuable information on the effects of normal physiological processes on intraocular pressure.

Previous attempts at developing such systems have been reviewed elsewhere (Cooper et al., 1979a). In this laboratory we have developed a method of noninvasive passive radiotelemetry of IOP. This communication reports the development of the system up to the present and illustrates its use in conditions designed to simulate clinical situations.

Materials and methods

Previously described miniature guard ring applanating transensors (AT) (Cooper and Beale, 1977) were mounted in acrylic or Sauflon haptic elements individually designed for the human, rabbit, or dog. The human AT holder was modelled from a standard acrylic haptic whose central optic was removed. The AT was mounted in the lower part of the scleral haptic so that it applanated the inferior sclera under the lower lid. The whole haptic ring was placed in the conjunctival fornix (Fig. 1). The rabbit and dog ATs were mounted in simple bilobed haptics, which rested in the upper conjunctival fornix of the rabbit or were sutured to the sclera in the dog eye.

Intraocular pressure was monitored with an automatic continual frequency monitor (ACFM) invented by one of us (DGB). This miniaturised monitor was fixed to the head by elastic bands,
The aerial was glued to the surface of the lower lid, overlying the AT, with silicone rubber adhesive (Dow Corning) (Fig. 1). The ACFM induces electromagnetic oscillations in the AT. The frequency of the oscillation is varied continually over the range of the resonant frequencies of the AT. The resonant frequency (Rf) of the AT is directly proportional to IOP. The ACFM monitors these changes in Rf and transmits the resonant frequency to a remote radio receiver, whose output is converted to an analogue signal recorded on a standard single channel chart recorder or processed by a Hewlett-Packard 9825 A computer and subsequently recorded on a 4-channel plotter (HP 9872 A). The tracing obtained is thus in the form of a continual variation of Rf, expressed in kilohertz (kHz), which faithfully follows IOP fluctuations.

Recordings of IOP were made in the anaesthetised dog and rabbit and in one conscious person.

**General Anaesthesia**

The effects of suxamethonium, the Valsalva manoeuvre, and Pco₂ variations on IOP were studied in the anaesthetised dog. The dog was intubated and maintained either under spontaneous respiration or intermittent positive pressure ventilation (IPPV). Central venous pressure (CVP) was recorded in this animal with a Sanborn pressure transducer.

In the rabbits IOP recordings were made with the animals maintained by intermittent intravenous injection of pentobarbitone sodium into an ear vein. Studies were made of the ability of the monitor to follow rapid variations in IOP due to pulse, respiration, and variations in depth of anaesthesia. Depth of anaesthesia was monitored by the lash reflex. The effect of digital pressure on the eye was recorded in this animal.

**The Conscious Human**

In man transducer stability and ability to follow rapid changes in eye IOP with respiration, eye movements, and lid closure were investigated. The response of the monitor was tested against intermittent recordings of IOP with a Mackay-Margtonometer (Berkeley Electronics).

**Results**

**IOP Monitoring under Anaesthesia**

The Valsalva manoeuvre was simulated by manual compression of the canine chest during the inhalation stroke of the ventilator. This resulted in a small rise in IOP (Fig. 2). Mean CVP was shown to rise from the initial level of zero to mean pressure of 10 mmHg (Fig. 3).

Injection of suxamethonium intravenously in the dog caused a dramatic rise in IOP, which lasted approximately 40 seconds, and settled to a level below baseline. The level to which IOP rose was more than 30 mmHg, however, as can be seen from the tracing; the range of the recorder was not high enough to record the maximum peak of IOP (Fig. 4). The CVP recording showed a similar rise in mean venous pressure from zero to approximately 15 mmHg, with a gradual decline to a mean CVP of approximately 5 mmHg (Fig. 5).

Variations in Pco₂ were accompanied by minimal changes in IOP. Correlation between Pco₂ and IOP was poor, whereas correlation between IOP and pH was better.

In the rabbit recordings of IOP showed high frequency excursions of IOP which were synchronous with the pulse. Well-defined low-frequency excursions of IOP due to respiration were observed.
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The IOP fell to just below baseline following this pressure on the eye.

Monitoring of IOP in the conscious human

Recording of the effect of heartbeat and respiration is shown in Fig. 9. As in the rabbit recordings, a high-frequency ripple is superimposed on the main recording, which was synchronous with pulse. The lower frequency variation was due to respiration. The effect of forced inspiration and expiration on IOP is shown in Fig. 10. Inspiration is accompanied by a rapid fall in IOP, with the converse occurring in all recordings. An example of such a recording is shown in Fig. 6.

If the depth of anaesthesia in the rabbit was allowed to decrease, there was invariably a rapid rise in intraocular pressure, as the rabbit regained lash reflex, and tone of eyelids and extraocular muscles. An example of a sudden rise in the pressure is shown in Fig. 7. The rapid excursions are due to respiration. With further injection of pentobarbitone and loss of lash reflex the IOP fell to approximately 10 mmHg. During the period of lightened anaesthesia the rabbits developed eye rolling movements and eventually blepharospasm, due to the presence of the haptic.

The effect of digital pressure through the eyelids in the rabbit is dramatically shown in Fig. 8. High spikes in intraocular pressure are seen towards the end of the recording, which were due to the application of a Mackay-Marg tonometer in order to correlate the monitor output with applanation pressures. The IOP fell to just below baseline following this pressure on the eye.

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on expiration. The effect of heartbeat is well shown. Inspiration and breath holding cause an initial reduction in intraocular pressure, followed by a gradual rise as the glottis is closed off (Fig. 11). During this manoeuvre there was always shown to be a short rise in IOP prior to the glottis being closed off and the chest wall relaxing.

Maximal gaze from right to left was accompanied by a rise and fall of IOP. In Fig. 12 the response of

Fig. 8 Ocular hypertension induced by digital compression of the eye of the anaesthetised rabbit. The high spikes are due to intermittent Mackay-Marg tonometry

Fig. 9 Continual recording of IOP by radiotelemetry in the conscious human. Note the excursions due to pulse and respiration

Fig. 10 Continual recording of IOP in the conscious human: the effect of deep breathing on IOP. In = respiration, out = expiration. Deep inspiration (in) decreases resonant frequency by approximately 270 kHz. Expiration (out) causes a rise of approximately 300 kHz. The rise and fall in IOP cannot be accurately estimated in mmHg without extensive calibration studies
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Fig. 11 The effect of breathholding in the conscious human. IOP decreases on inspiration, and increases to baseline on closure of the glottis.

the IOP monitor to right gaze (R) return to the primary position (A) and then full left gaze (L) followed by a repeat of the manoeuvre is shown. These IOP variations are about 10 mmHg.

Blepharospasm is thought to cause a rise in intraocular pressure, which was demonstrated in the rabbit experiments. In man closure of the eyelids resulted in a sustained rise in intraocular pressure (Fig. 13). The IOP rose approximately 7 mmHg in this experiment. It will be noted at the beginning of this recording that there are multiple high spikes in the IOP recording. These are blink artefacts, the amplitude of which do not fully reflect IOP, but are due to rapid variations in inductive coupling between the AT and the aerial, as shown by oscillographic monitoring.

The response of the monitor to IOP, when tested against intermittent recordings with the Mackay-Marg tonometer, correlated well with the tonometry readings for up to 2 hours of continual recording. There was then a decline in Rf. We were interested in the effect of continual application of the Mackay-Marg tonometer to the cornea for purposes of calibration of the instrument for individual eyes. The results of these experiments are reported elsewhere (Cooper et al., 1979b). Fig. 14 illustrates the Mackay-Marg tracing above and the monitor output below. The monitor output faithfully reproduces the rise in IOP induced by the tonometer application. The superimposed fluctuations due to heartbeat, respiration, and tremor of the hand holding the tonometer are reproduced in both tracings.

Fig. 12 The effect of maximal eye movements on IOP in the human: R=right gaze, A=primary position, L=left gaze. Subsequent changes in IOP are due to similar alternating gaze movements ending in a rapid movement to the left, reverting finally to the primary position. True change in IOP cannot be accurately assessed in mmHg. Some of the rise in IOP may be due to distortion of the sclera by the acrylic haptic

Fig. 13 Continual recording of IOP in the conscious human by radiotelemetry. Closure of the eyelids causes a rise in IOP. The rapid spikes are blink artefacts (see text)

Discussion

It is well known that IOP is normally maintained at a constant level with only minor diurnal variations. There is some compensation for rapid variations of IOP by the viscoelastic properties of the globe and by efflux of blood from the choroid. However, rapid rises of IOP cannot be fully accommodated by either mechanism. We simulated several conditions that are known to cause acute ocular hypertension during eye surgery. We have demonstrated 2 mechanisms for suxamethonium induced ocular hypertension, namely, contraction
have to be improved in order to reduce water vapour permeability for long-term stability. Mechanical problems with the AT and its coupling to the sclera have to be overcome in order to improve accuracy enough to make the instrument capable of quantitative assessment of IOP over at least 24 hours.

The final system, at present being completed, consists of the ACFM reported here, which will transmit IOP information to a portable processing unit, which can sample and record pressure once every 2 minutes for a period of several seconds, over 24 hours. The result of each IOP sample will be recorded on the digital memory board.

Information will be read out from the memory board by the computer and recorded on a disc or plotter. Later a microprocessor and simple chart recorder will be substituted for the computer. The size of the portable processing unit will be such that it can be carried on a waist belt, a shoulder strap, in a handbag, or stored on a bedside table at night. There will be no wired connection between the monitor and the portable processor/memory board. The patient will therefore be unencumbered apart from the requirement to carry the processor around with him or have it within 3 to 6 feet (2 to 3 m) from the ACFM, which is placed on the head or can be included on a spectacle frame.

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References


ADDED IN PROOF. After submission of this paper for publication a reference was found to direct recording of intraocular pressure in man. It reports similar observations to those observed in man by our monitor (Coleman, DJ, Trokel, S. Arch Ophthalmol 1969; 82: 637-40.