Endoscopic visualisation of the human nasolacrimal system: an experimental study

A D Singh, A Singh, I Whitmore, E Taylor

Abstract
Orthograde and retrograde endoscopy of the upper and lower nasolacrimal system was performed using two prototype ultrathin (0.5 mm and 1.1 mm diameter) fibrescopes on four cadaver heads. Appearances were verified by subsequent dissection. The procedure, which we term 'dacryocystoscopy' is described. With modifications this technique may have clinical applications in the treatment of nasolacrimal disorders.

Thorpe' was the first to describe the ophthalmic application of an endoscope in 1934 for the removal of intravitreal non-magnetic foreign bodies. With the advent of fibroptics, flexible and fine calibre endoscopes could be designed and Norris' adapted a 1.7 mm fiberoptic endoscope for both intracocular and orbital surgery. More recently a new generation of electronic videendoscopes has become commercially available for gastrointestinal endoscopy. Instead of fibroptics these endoscopes are based on a charge-coupling device (CCD). The image from the tip of the endoscope is focused on a CCD that transmits the image electronically via a video system to the television monitor. As the image is stored digitally this offers potential capability of image enhancement and computer analysis. A new ophthalmic electronic videendoscope system for intraocular surgery with a 20 gauge (0.89 mm diameter) probe has been designed. This helps to visualise the pars plana, ciliary body, and the posterior surface of the iris.

Endoscopy of the nasolacrimal system has been reported previously with rigid and semirigid endoscopes. Ashenhurst and Hurwitz developed a prototype lacrimal endoscope (canaliculoscope) and used it to visualise the canaliculus and lacrimal sac. Following a recent report on salivary gland endoscopy using 0.8 mm ultrathin diameter fibroscope (endoscope based on fibreoptics) we decided to investigate the feasibility of endoscopic visualisation of the entire nasolacrimal system using prototype fibrescopes on cadavers and to confirm the appearances by dissection.

Material and methods
Four embalmed cadaveric heads were made available at the Department of Anatomy, Queen Mary and Westfield College. The upper nasolacrimal system was studied in two specimens. To study the lower system (lacrimal sac and nasolacrimal duct) sagittal sections of the head (two) were used. The nasal septum and the inferior turbinate were removed to visualise the opening of the nasolacrimal duct into the inferior meatus.

Two prototype ultrathin fibrescopes PF-5 and XTUF-11 (Olympus Co Ltd, Tokyo) were used. The PF-5 fibrescope has an outer diameter of 0.5 mm, is forward viewing, with a depth of field of 3 to 50 mm (Fig 1). The XTUF-11 fibrescope has an outer diameter of 1.1 mm and has an instrument channel 0.1 mm wide. The depth of field is similar to the PF-5 fibrescope. Both instruments have a 60 degree field of view. CLV-10 OES was used as the light source. The fibrescope was attached to a compact colour video camera (Olympus OTV-F2) which measures 17 mm x 48 mm and weighs 20 g. This produces high resolution images even in low light conditions. This was coupled to a portable CCTV unit (Olympus PVS-1) combining a 9 inch monitor and Video8 VTR; this provided a display of the magnified images during the length of the procedure. The magnification from the fibrescope to the PVS-1 monitor is by a factor of 1.7. A continuous recording was made and at the end of each procedure the tapes were analysed and selected images were printed.

UPPER NASOLACRIMAL SYSTEM (ORTHOGRADE ENDOSCOPY)
After dilatation of the punctum, graduated silver probes ('0' to '3' John Weiss) were inserted to dilate the upper canaliculus. After flushing the nasolacrimal system with normal saline using a syringing canula, healon was injected to keep the structures dilated so as to facilitate introduction

Figure 1 The PF-5 fibroscope.

Figure 2 The positions of the fibroscope tip in the following figures.
of the fibrescopes. The PF-5 fibrescope could be introduced easily into the canaliculus up to the level of the lacrimal sac. Greater dilatation with a size '3' silver probe was required before the thicker XTUF-11 fibrescope could be introduced. Neither of the fibrescopes could be manipulated to enter the upper opening of the nasolacrimal duct owing to the rigidity of the embalmed cadaveric tissues and extreme flexibility of the fibrescopes. Both fibrescopes provided adequate visualisation of the upper nasolacrimal system.

LOWER NASOLACRIMAL SYSTEM (RETROGRADE ENDOSCOPY)
Using the sagittal sections of the head and after removing the inferior turbinate, the lower nasolacrimal duct opening was identified by syringing through the canaliculus. Through the lower opening the nasolacrimal duct was gently probed before the fibrescopes were introduced. Again both types of fibrescopes could be passed easily through the nasolacrimal duct into the sac up to the level of the fundus of the lacrimal sac. Both fibrescopes provided good quality pictures allowing us to identify distinctly various structures. A small piece of steel wire (0·1 mm diameter) with a gentle curve introduced into the 0·1 mm instrument channel of the XTUF-11 fibrescope gave it sufficient rigidity and curvature which facilitated exploration and visualisation of the inner aspect of the lacrimal sac. Following visualisation the lacrimal sac was dissected to reveal the features of its inner surface.

Results
Both prototype ultrathin fibrescopes PF-5 and XTUF-11 provided good quality magnified images which could be viewed constantly on the TV monitor and recorded simultaneously.

Fibrescope PF-5 provided superior quality images (Figs 5, 6, 9) to the fibrescope XTUF-11 (Figs 3, 4, 7). There was no other difference in the optical aspects of the image. The magnification is inversely proportional to the distance between the tip of the fibrescope and the object to be viewed. The image is further magnified 1·7× between the fibrescope and PVS-1 monitor.

The endoscopic findings are reported for each nasolacrimal system separately. Figure 2 shows the various positions of the fibrescope tip in the nasolacrimal system in relation to Figures 3–11.

PUNCTUM (FIG 3)
Though this is strictly not an endoscopic picture it is included for the sake of completion. The papilla appeared as a ring of bright light which is surmounted by a dark appearing puncta (post dilatation).

CANALICULUS (FIG 4)
The 'tunnelling' effect is obvious. The canaliculus appeared as a dark hole. The obliquity of the fibrescope tip in relation to the canalicular wall gave it a sloping appearance and this also explains

\[Fig 5a\]
\[Fig 5b\]
Figure 5 (a) Endoscopic view of the fundus of the lacrimal sac. (b) m=mucosal fold and f=fundus.

\[Fig 6a\]
\[Fig 6b\]
Figure 6 (a) Endoscopic view of the side walls of the sac. (b) w=wall and f=fundus.

\[Fig 7a\]
\[Fig 7b\]
Figure 7 (a) The opening of the canaliculi into the lacrimal sac. (b) c=canalicular openings.
Endoscopic visualisation of the human nasolacrimal system: an experimental study

Figure 8 (a) Appearance after dissection confirms separate openings of upper and lower canaliculi into the lacrimal sac. (b) l=lid margin, a=anterior lacrimal crest, c=canalicular openings, and e=edge of the reflected anterior wall of the lacrimal sac.

![Image](image_url)

**Figure 8a**

**Figure 8b**

the apparent eccentric location of the canalicular lumen.

**LACRIMAL SAC (A) (FIG 5)**

In this position the tip is very close to the fundus of the lacrimal sac. The central circular brighter zone corresponds to the inner concavity of the fundus. The two bright spots probably represent mucosal folds.

**LACRIMAL SAC (B) (FIG 6)**

As the tip was withdrawn slightly the appearance changed completely. The central brighter zone was replaced by a central dark zone surrounded by a bright area. This was more obvious in the upper half of the picture. As the tip moved away from the fundus the light being reflected by it reduced (and hence the bright to dark transition) and the light reflected from the side walls of the sac came into the view (outer bright area).

**LACRIMAL SAC (C) (FIG 7 AND FIG 9)**

With gentle manipulation of the tip it was possible to visualise the inner openings of the canaliculi. The upper and lower canaliculi opened separately into the lacrimal sac in both cases. On subsequent dissection this appearance was clearly confirmed in one case (Fig 8). However in the other case dissection presented an appearance of partial separation (Fig 10).

**NASOLACRIMAL DUCT (FIG 11)**

An oval appearance of the upper end of nasolacrimal duct was identified. The bright spot near the opening is either an artefact or a mucosal fold.

**Discussion**

The two prototype fibrescopes PF-5 and XTUF-11 investigated in this study provided clear and reproducible images of the entire nasolacrimal system. Our findings demonstrate progress from previous attempts at endoscopic visualisation to include orthograde and retrograde endoscopy. This has enabled the visualisation of the inner openings of the canaliculi and of the nasolacrimal duct origin. In addition endoscopic appearances have been confirmed by dissection.

The fibrescopes used were thinner than the silicon tubing (1.19 mm) suggested by Crawford" for intubation of the lacrimal system. The outer diameters of the fibrescopes are compared with the size of silver probes (Fig 12). The fine calibre of these fibrescopes makes them inherently flexible. Because of this the fibrescopes could not be passed into the upper nasolacrimal opening during orthograde endoscopy and the inferior turbinate had to be removed to facilitate retrograde endoscopy. Though the quality of images generated by the fibrescope XTUF-11 were inferior to those of fibrescope PF-5, it may still be the preferred endoscope for two reasons. Firstly, greater thickness (1.1 mm compared
coupled to a 300 μm quartz fibreoptic has been used to create an intranasal dacyrocystorhinostomy fistula in a patient undergoing endonasal laser dacryocystorhinostomy. This is claimed to provide good haemostasis with reduced morbidity. Gonnering et al. have reported excellent results in 18 patients undergoing transnasal laser assisted lacrimal procedures. They performed laser rhinostomy with carbon dioxide and potassium titanyl phosphate laser delivered through a 300–600 μm fibre under video endoscopic visualisation. In addition chromium-sensitised, and thulium and holmium doped YAG laser (THC:YAG laser) coupled to a 480 μm optic probe can successfully create a limbal sclerostomy. The technique of probing used in children with epiphora is a ‘blind’ procedure. Bends in the course of the lower tear duct have been shown to exist and canalicular stenosis following probing for congenital nasolacrimal duct obstruction is a well known complication. Using the technique described by us, the upper nasolacrimal duct opening may be visualised and the block in the lower nasolacrimal duct treated under direct observation.

This technique, termed dacyrocystoscopy, is still experimental. However, further studies on patients are planned to assess its full potential.

We are grateful to Mr Andrew Ekins of KeyMed Ltd, Southend, UK, for providing the prototype fibrescopes and the recording equipment. We thank Mr J J Kanski, FRCS, for his critical reading of the manuscript. The authors wish to declare that they have no commercial interest in any of the instruments/products used in this study.


Table 1: Comparison of outer diameters (in mm) of the fibrescopes and the commonly used Bowman’s lacrimal probes. (Data have been kindly provided by KeyMed Ltd, Southend, UK and John Weisz and Son Ltd, Milton Keynes, UK.)

<table>
<thead>
<tr>
<th>Diameter (mm)</th>
<th>PF-5</th>
<th>XTUF-11</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>00</td>
<td>00</td>
</tr>
<tr>
<td>1.0–1.1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>0.89</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>0.98</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.0–2.22</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 10 (a) Partial separation of canalicular openings as seen on dissection. (b) l= lid margin, c= canalicular opening, s=sac lumen, and a= anterior lacrimal crest.

Figure 11a (a) The view of the nasolacrimal duct origin. (b) w= lacrimal sac, n= nasolacrimal duct opening, and a= artefact.

Figure 12 Comparison of outer diameters (in mm, magnified 10×) of the fibrescopes and the commonly used Bowman’s lacrimal probes. (Data have been kindly provided by KeyMed Ltd, Southend, UK and John Weisz and Son Ltd, Milton Keynes, UK.)
Endoscopic visualisation of the human nasolacrimal system: an experimental study