New Dacron tissue colonisable keratoprosthesis: clinical experience

S Pintucci, F Pintucci, M Cecconi, S Caiazza

Abstract
Background—Keratoprostheses (KPs) are made of an optical cylinder integrated with a supporting element which conditions their biocompatibility. A new KP with a Dacron tissue colonisable support has been designed in order to reduce significantly the complication rate.

Methods—This new KP was implanted into 20 eyes of 20 patients with bilateral corneal blindness unsuited to a corneal implant. The follow up ranged from 24 to 96 months.

Results—All 20 patients had some improved visual acuity with 13 retaining this improvement for more than 2 years.

Conclusion—These favourable results may be indicative of the reliability of the new KP.

(Br J Ophthalmol 1995; 79: 825-829)

A keratoprosthesis (KP) implantation is the last resort in severe corneal blindness not amenable to a corneal transplant.1 Although early visual recovery may be good, the percentage of severe complications with KPs is extremely high.1-2

KPs are generally made up of a polymethylmethacrylate (PMMA) optical cylinder,3-4 focusing the images on a functioning retina, integral with a supporting element fixing the KP to the eye.

The most important problem, which arises with the implantation of a KP, is obtaining a perfect biological, mechanical, and functional anchorage of the supporting element to eye tissues. However, the main complications are related to the eye’s capability of extruding biomaterials only mechanically anchored but not biologically integrated, such as rigid KP supporting elements made of metal, plastic, ceramic,1-6 as well as hetero-, auto-, and homotissues (dentin, cartilage, bone).7-9 On the basis of previous studies, it can be stated that an ideal supporting element should fix the device without biological and mechanical adverse effects.8-10,11

With the aim of avoiding the complications caused by interactions between the supporting element and the eye tissues, a new KP with a biointegratable supporting element12 was developed by Pintucci in 197913 and the implantation technique is being constantly improved.14-16

In this new device, a Dacron supporting element12 is fixed, according to a personal technique (international patent pending), to a medical grade PMMA optical cylinder.

Previous experiments13 performed with human fibroblasts cultured in the presence of untreated Dacron tissue showed normal cells proliferating and adhering to the Dacron filaments. Dacron tissue implanted for 30 days under the conjunctiva of rabbit eyes, appeared to be colonised by a vascularised connective tissue filling any space within the Dacron weft.13-16 Moreover, specimens of Dacron tissue, placed under the lower lid skin for a period ranging from 20 to 40 days appeared to be tridimensionally colonised by autologous connective tissue (Figs 1 and 2).

On the basis of these results, we decided to implant our new KP in 20 eyes with bilateral corneal blindness not suitable for a corneal transplant.

Materials and methods
The KP is made of medical grade PMMA and of Sauvage filamentous Dacron fabric (nominal thickness 0.6 mm, mean water porosity 1·600 ml).

The weft of the Dacron tissue supporting element is fixed to a PMMA optical cylinder 5·4 mm long and 3·5 mm wide. The weft of the Dacron tissue is shown in Figure 3 and the assembled device and its nominal size are shown in Figures 4 and 5, respectively.

For inclusion in this study, bilateral corneal blindness not suitable for corneal transplant was required. All procedures were first time KP implants in a single eye with a severely vascularised cornea. All patients gave informed consent.

A preoperative diagnosis of ocular pemphigoid was made in 12 patients, three eyes had...
trachoma, two eyes had recurrent severe herpes keratitis, and three eyes had alkali chemical burns. The diagnoses were made on the basis of clinical guidelines without relying on histopathological or immunological testing. Previous cataract surgery at the time of the initial KP operation was recorded in five patients.

Preoperative examination included vision testing, slit-lamp biomicroscopy, ultrasonography (A and B scans), evaluation of intraocular pressure by whichever means possible, electroretinogram and visual evoked potentials, Schirmer 1 test, and basic lacrimation test. Conjunctiva and lid examination is very important because reconstructive plastic surgery is often necessary before the KP implant, as in cases of trichiasis and lagophthalmos, etc.

The age of the patients at surgery ranged from 25 to 86 years with a mean of 57.9; there were 10 men and 10 women. All patients were in the chronic, non-inflammatory phase of the disease. All patients had cicatricial entropion with marked shortening (greater than 50%) of conjunctival fornices, symblephara, and a variable amount of trichiasis. A history of glaucoma was obtained in one patient.

TECHNIQUE

The surgical technique of implanting a KP consists of two stages. In the first stage after eyedrop anaesthesia, the centre of the cornea is marked with gentian violet. General anaesthesia with nasal intubation is preferred.

In order to colonise the KP, a 15 mm incision of the skin is performed in the inferior orbitopalpebral sulcus, the orbicular muscle fibres are spread apart to the orbitopalpebral septum and the KP is introduced upside down with the optical cylinder vertical in the pocket. The orbicular muscle is sutured with 8-0 Dexon and the skin with 6-0 black silk.

If trichiasis, entropion, or symblepharon are present, palpebral and conjunctival plastic procedures must be performed. In dry eyes the lacrimal puncta are closed with diathermy. To expose the eye, two lid traction sutures are applied. The corneal epithelium is completely removed and a 10-0 Nylon suture is applied on the mark at the corneal centre for future reference.

A free buccal mucosal graft is dissected bearing in mind that once removed from the mouth, the mucosa shortens by one third in diameter. The buccal mucosa must also cover possible defects in the palpebral and bulbar conjunctiva.

Excessive submucosal fat is dissected away with fine scissors. The mucosa is washed and kept in a balanced salt solution-gentamicin solution. The oral mucosa is applied on the cornea whose epithelium is removed and sutured with Dexon 8-0. All blood clots are carefully removed. Antibiotic ointment is instilled and the lids closed. The dressing and a protective plastic shield are applied.

The second stage is performed after 2 months. Eye hypotony is obtained and general anaesthesia is performed. The colonised KP is removed from the lower lid. Under the operating microscope the excessive connective tissue that covers the optical cylinder is removed, the fixation of the optical part is tested, and the colonisation of the Dacron tissue is checked. The cornea is partly exposed dissecting the oral mucosal graft from the temporal-superior sector to the centre.

Once the cornea is exposed the KP optical cylinder site is stained with a 4 mm circular optical zone marker for corneal refractive surgery. With a diamond knife three partial thickness radial incisions are performed in the 2, 6, and 10 o’clock meridians. The cornea is trephined with Franceschetti trephine, then the radial incisions are completed. The iris is drawn down in the 6 o’clock direction and, with lateral movements, is extirpated, followed by cryoextraction of the lens.

The KP optical cylinder is positioned and the colonised Dacron tissue and the radial incisions are sutured. The oral mucosa is sutured to cover the KP and trephined to allow the passage of the anterior optical part.
In dry eyes, before the KP implantation in the eye, in order to protect the oral mucosa, the length of the palpebral fissure is reduced with a Blaschcovicus lateral and a Strefl nasal tarsorrhaphy, leaving a central aperture for the optical cylinder.

Antibiotic ointment is instilled, and a protective plastic shield applied.

Results
We reviewed the charts of the 20 eyes of patients who underwent a Pintucci’s trans-mucous KP implantation from January 1987 to December 1991. Follow up after surgery ranged from 24 to 96 months with a mean follow up of 58 months.

On preoperative examination the best corrected visual acuity was perception of light. A history of preoperative glaucoma was obtained in one patient.

In the follow up the appearance of the eye must be checked frequently. Follow up was done every week for the first 5 months after the KP was implanted, and thereafter once a month. Cases were reviewed with reference to the length of time the KP remained in situ, types and rates of complications, best postoperative aided visual acuity, and length of time vision was maintained.

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Y=year of implantation; Age=age when implanted; S=sex; PP=primary pathology; P=pemphigoid, T=trachoma, HK=herpes keratitis, AB=alkali burn; VA=postoperative visual acuity; R=postoperative correction; F-U=months follow up; GN=oral mucosal graft necrosis; AM=anterior membrane; RM=retroprosthetic membrane; RD=retinal detachment; CD=choroidal detachment; EN=endophthalmitis; KE=keratoprosthesis extrusion.
Discussion

A keratoprosthesis implantation is the last resort to regain useful visual acuity in severe corneal blindness caused by trauma, chemical burns, infections, trachoma, ocular pemphigoid, Stevens-Johnson syndrome as well as severe dry eyes, repeated graft failures, etc. and when a corneal transplant cannot be attempted or has repeatedly failed.

None of the patients had a bullous keratopathy, which we consider a poor indication for a KP implantation because it is amenable to keratoplasty.

KPs basically consist of an optical cylinder, penetrating the cornea and focusing the images on the retina, integral with a supporting element fixing the device to the eye.

Although early visual recovery may be good severe complications, such as extrusion, are extremely high in spite of the number of devices and surgical techniques described in literature.

Some problems with the long term performance of KPs have not yet been solved and, unfortunately, worldwide research in this field is very limited.1 2 Since 1789, several KPs have been developed and from the clinical results of a very large number of different implants it can be recognised that PMMA has been the most widely used material for optical cylinders and it has not been reported as a cause of failure.1 3 4 On the contrary, most complications are due to rigid supporting elements which are simply sutured and/or screwed to the cornea. In these devices the mechanical stress leads to inflammation with tissue melting and to the formation of empty spaces at the implant-eye interface. As a consequence, aqueous humour leakage, infections, and reparative epithelial and connective tissue proliferation may occur, with subsequent encapsulation, formation of retroprosthetic membranes, and KP extrusion.4 8 11 17

Many biological materials, autologous (fascia lata, periosteaum, conjunctiva, Tenon's capsule, cornea, labial mucus, cartilage, eyelid skin) and homologous (cornea and sclera) have been used for covering rigid supporting flanges1 4 6 8; however, in 1979 we developed a new KP with a Dacron tissue supporting element with the aim of reducing significantly the complication rate. Since 1954, Dacron fabrics have been used successfully in permanent cardiovascular prostheses which were colonised by host tissues.1 2

Previous in vitro experiments performed with human lung fibroblasts cultured in the presence of untreated Dacron tissue showed normal cells proliferating and adhering to the Dacron filaments. The same tissue implanted for 30 days under the conjunctiva of rabbit eyes, appeared to be colonised by a vascularised connective filling any space within the Dacron weft.13-16 Moreover, specimens of Dacron tissue, placed under the lower lid skin for a period ranging from 20 to 40 days before implantation on the eye and observed with light microscopy and scanning electron microscopy, appeared to be tridimensionally colonised by autologous connective tissue (Figs 1, 2).

After extensive in vitro and in vivo testing of different kinds of Dacron tissues, the Sauvage filamentous Dacron fabric was chosen for its good colonisability and mechanical performance. The main characteristics of this Dacron tissue are: it is soft and pliable (thus preventing aseptic corneal necrosis by mechanical pressure); chemically inert; not subject to resorption; does not activate the complement (to some extent); can be autoclaved; can be easily cut in the desired shape; and can be sutured.12

Human fibroblasts in vitro proliferate, adhering to the Dacron filaments without cellular damage.13 In vivo the Dacron tissue is colonised by a vascularised connective tissue migrating from the surrounding tissues, filling completely the free spaces among the filaments. It must be stressed that reparative epithelial proliferation along the optical cylinder is stopped owing to contact inhibition17 and to lack of empty spaces which have been filled by the neoformed connective tissue.

Results were gauged by measuring the visual acuity, the length of time improved vision is retained, and the time the KP remains in situ. Visual acuity itself is not necessarily a criterion of success or failure, as good guiding vision may be achieved without central vision. Complications must be overcome by careful follow up and prompt action.

It is difficult to predict which cases will be successful. Patients thought to have a very poor prognosis may do well. In general patients with good tear secretion do better than patients with dry eyes.

Contrary to what has been suggested by some authors,3 10 18 19 we do not graft the supporting flange in the corneal stroma thickness but we place it on the corneal surface. In this way the colonised Dacron tissue plays a trophic and a mechanical role. It is important to underline that extremely thin and vascularised corneas may be treated successfully in this way too.

The oral mucosa employed to cover the colonised Dacron flange shows best results probably because it is more vascularised and has a faster cellular turnover than skin, conjunctiva,20 and other tissues.

It must be stressed that the colonised Dacron tissue KP implant can be considered and behaves as an autotransplant placed between the cornea and the oral mucosal graft. It heals, becoming biologically and mechanically fully integrated with the surrounding tissues, and also acts as a barrier to microbial contamination. It is, in fact, well known that biomaterials, as well as traumatised tissues, may be a pabulum for bacteria leading to infections typically associated with prosthetic devices.21 22

Infections occur in all kinds of implanted devices and are related to the biomaterials...
New Dacron tissue colonisable keratoprosthesis: clinical experience

which lower the natural defences of the organism. In addition they promote a preferential adhesive colonisation\textsuperscript{21} complicated by enhanced antibiotic resistance owing to the modifications of the saprophytic behaviour of bacteria.\textsuperscript{21-24} Inflammation, aseptic bacteria.\textsuperscript{21-24} Infiltration of modifications of the which lower the natural defences of the organism to predispose to implant failure.

We think that this new device may be considered as a real step on the way to overcoming the apparently inseparable difficulties represented by KP mechanical anchorage and biointegrability.

20 Kozarsky AM, Knight SH, Waring GO. Clinical results with a ceramic keratoprosthesis placed through the eyelid. Ophthalmology 1987; 94: 904–11.
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