Auricular cartilage versus donor sclera as a wrapping of hydroxyapatite orbital implants

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Abstract

• AIM: To retrospectively compare postoperative outcomes after primary enucleation and placement of a hydroxyapatite (HA) implant without wrapping, wrapped with auricular cartilage or donor sclera.

• METHODS: Medical records of patients presented as intraocular tumor or severe ocular injury were identified from the electronic medical record system. Cases underwent enucleation and HA orbital implantation were enrolled in this study and were divided into 3 groups according to the wrapping material of HA implant. Cases with autogenous cartilage caps were enrolled in group A (n=11), with donor sclera caps in group B (n=12), and without any wrapping material in group C (n=9). Follow-ups were set at 1, 2wk, 1, 3, 6, and 12mo after surgery.

• RESULTS: Altogether 32 cases finished the follow-up and were enrolled in this study. Three cases (27.27%) in group A, 4 cases (33.33%) in group B, and 4 cases (44.44%) in group C developed one complication each after surgery. In group A, no HA exposure occurred, but conjunctival inclusion cyst occurred in one and severe conjunctive chemosis in two cases. In group B, one HA exposure occurred, conjunctival inclusion cysts occurred in one, severe conjunctive chemosis occurred in one, and conjunctival granuloma occurred in one case. In group C, one HA exposure occurred, severe conjunctive chemosis occurred in two cases, and conjunctival granuloma occurred in one case. The case of exposure of none-wrapped implant was noted in the first 6mo after placement of the orbital implant. The case of exposure of donor sclera-wrapped implant was noted at the 12mo after placement of the orbital implant. Both exposure cases were treated successfully with conservative treatment.

• CONCLUSION: With low incidence of implant exposure and mild complications, auricular cartilage can be a good choice of alternative wrapping material of orbit implant with satisfied outcome.

• KEYWORDS: hydroxyapatite orbital implants; wrapping; auricular cartilage; donor sclera; complication

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INTRODUCTION

The cosmetic results of simple enucleation are always poor because of the immobile staring appearance and tendency of lower lid sagging which would lead to a deep hollow below the brow. This outcome can be considerably improved by placing an orbital implant to fill the volume of the lost eye[1].

The hydroxyapatite (HA) porous implant was the most widely used type of orbital implant due to its good biocompatible (minimal tissue inflammation) and low resorption (rapid host tissue ingrowth)[2-3]. However, the surface of porous implants may be abrasive to the orbital soft tissue above, which would potentially cause tissue erosion with implant exposure. Thus this porous implants would increase the risk of orbital implant exposure[4-5]. Wrapping the implant could physically add another layer of tissue between the implant and overlying conjunctiva, thus to decrease the incidence of implant exposure and provide a surface for muscle attachment as well[6].

Donor sclera has been used widely as the wrapping material of HA implants[7-8]. Although donor sclera kept unchanged in most cases, late thinning or entirely disappearing was happened in some cases[9]. This late thinning or disappearing of the sclera over unorganized implants could cause spontaneous rupture of the conjunctiva with exposure of the unorganized implant[9]. Also, it has been proved that current screening tests may miss an infected blood donor 1:1 800 000 for HIV, 1:1 200 000 for hepatitis C virus, 1:200 000 for hepatitis B virus and Creutzfeldt-Jakob disease (CJD)[10-12]. What’s more, donor sclera which needs to be preserved in the eye bank, may be unavailable sometimes especially in case of an emergency.
This possibly happening of late thinning, infectious disease spreading and occasional unavailability of donor sclera has prompted the use of other alternative wrapping materials. Auricular cartilage is the autogenous material which shares various characteristics with donor sclera\[13-14\]. It has been used for lots of oculoplastic procedures including as a substitute of tarsal plate for reconstruction of full-thickness eyelid defect, as a spacer for upper and lower eyelid retraction correction, and as a support of nasal tip in rhinoplasty\[14-16\]. This study aimed to compare postoperative outcomes after primary enucleation and placement of the HA implant without any wrapping, with auricular cartilage cap or donor sclera cap.

**SUBJECTS AND METHODS**

**Ethical Approval** This retrospective study followed the tenets of the Declaration of Helsinki and was approved by the Ethics Committee of the Eye Hospital of Wenzhou Medical University (Medical Ethics Committee, Wenzhou Medical University, Wenzhou, Zhejiang Province, China). Informed consent was obtained from all the patients. Patients were eligible for inclusion if they had been diagnosed with intraocular tumor or severe ocular injurychronic dacryocystitis and elected to undergo surgical treatment enucleation and HA orbital implantation. Patients were excluded when they were <18 years old or >65 years old, had lid malposition, unable to tolerate general anesthesia, with systemic autoimmune disease, needing radiotherapy or chemotherapy after surgery, and follow-up period <12 mo.

We record the age, gender, eye, the cause of enucleation, size of implant, and the surgical type (autogenous ear cartilage cap, donor sclera cap, without wrapping material) of all patients. The complications, its occurrence time, subsequent therapy, and prognosis were recorded as well. A total of 32 patients meet our criteria. These cases were divided into 3 groups (groups A, B, C) according to the wrapping material of HA implant. Cases with autogenous ear cartilage caps were enrolled in group A, with donor sclera caps were enrolled in group B, without any wrapping material were enrolled in group C.

All of the procedures were conducted under general anesthesia by a single surgeon (Chen X). A piece of autogenous ear cartilage graft, 15×15 mm\(^2\) in size, harvested from the conchal lateral wall through a post-auricular incision of the right ear and then be preserved in a mixture of gentamicin and normal saline (Figure 1A, 1B). Alcohol-preserved sclera provided by Eye Bank of Eye Hospital of Wenzhou Medical University was used in group B cases. After 360 degree conjunctival peritomy, 4 rectus were isolated and suspended with 6-0 vicryl suture [Alcon (China) Ophthalmic Product]. The inferior and superior oblique were cut after clipping. Then the transection of the optic nerve was performed followed by application of pressure for hemostasis. A sizing ball was used to evaluate the residual intraconal volume, and implant size was chosen to allow tension-free closure of the anterior ocular tissue. The HA implant (Bio-Eye; Integrated Orbital Implant) was placed in the cone after soaking in a mixture of gentamicin (80 000 units), hexadecadrol (1 mL:5 mg) and normal saline for 5 min (Figure 1C). Autogenous ear cartilage (group A; Figure 1D) or donor sclera (group B) was then sutured to wrap the frontal part of the implant. Muscles were sutured to attach at the recti normal anatomical insertion sites with 6-0 vicryl sutures [Alcon (China) Ophthalmic Product; Figure 1E]. The rectus were imbricated anterior to the HA implant in group C cases as Guthoff RF’s method. The superior and inferior rectus muscles, as well as the medial and lateral rectus muscles, were respectively knotted together forming a joint-like structure between the anterior part of the HA implant and the Tenon’s capsule\[17\]. Tenon’s tissue was then closed with multiple interrupted 6-0 vicryl sutures and conjunctiva was closed with a continuous 6-0 vicryl sutures (Figure 1F). A conformer was placed in conjunctival sac with antibiotic ointment. Pressure bandaging was applied for 3 d after surgery. Postoperative care included administering methylprednisolone (20 mg/kg/d) for 3 d and ceftriaxone (1.0 g/d) for 3 d. Topical tobramycin dexamethasone (Alcon, ophthalmic ointment) was prescribed for 1 mo after removing the bandage.

Follow-up period was set at 1, 2 wk, 1, 3, 6, and 12 mo after surgery. At each follow-up, complications including the existing of implant exposure, conjunctival inclusion cysts, conjunctive granuloma, and severe conjunctive chemosis were recorded. For cases with intraocular tumor before surgery, orbital magnetic resonance imaging (MRI) was performed to evaluate the possibility of residual tumor. Statistical analyses were performed with SPSS version 19.0. The demographic data of 3 groups were compared using the one-way ANOVA or Chi test or Fish’s test. Results were considered significant at \(P<0.05\).

**RESULTS**

In total, 32 cases finished the follow-up and were enrolled in this study, including 11, 12, and 9 cases in group A, group B and group C, respectively. The clinical characteristic of them were list in Table 1.

No statistically significant differences were identified in patients age (\(P_{1-2}=0.908\), \(P_{1-3}=0.573\), \(P_{2-3}=0.659\)), sex (\(P_{1-2}=0.628\), \(P_{1-3}=0.670\), \(P_{2-3}=0.659\)), cause of enucleation (\(P_{1-2}=0.590\), \(P_{1-3}=1.0\), \(P_{2-3}=1.0\)), and the size of implant (\(P_{1-2}=0.573\), \(P_{1-3}=0.907\), \(P_{2-3}=0.515\)) among 3 groups.

Altogether 3 cases (27.27\%) in group A developed complications after surgery, 4 cases (33.33\%) in group B, and 4 cases (33.33\%) in group C.
in group C. All cases with complications developed only one complication.
No HA exposure (0) occurred in group A (Figure 2). One case (8.33%) developed HA exposure (1×1 mm$^2$) in group B at 1y follow-up. It is a 65-year-old man who had never taken out the ocular prosthesis after surgery. The exposure developed at 12mo after surgery and was recovered with conservative treatment. One case (11.11%) developed HA exposure in group C at 6mo follow-up. It is a 31-year-old man and was treated successfully with conservative treatment as well.

Of the 11 patients in group A, three cases (27.27%) developed mild complications, including one conjunctival inclusion cysts and two severe conjunctive chemosis. Of the 12 patients in group B, three cases (25%) developed mild complications, including one conjunctival inclusion cysts, one severe conjunctive chemosis, and one conjunctive granuloma. Of the nine patients in group C, three cases (33.33%) developed mild complications, including one conjunctive granulation and two severe conjunctive chemosis (Table 2).

At 12mo follow up, the MRI showed good shape and position of autogenous ear cartilage in cases with intraocular tumor before surgery in group A (Figure 3).

DISCUSSION
In this study, we report our experience using auricular cartilage or human donor sclera to cover enucleation implants. This is the very first time using auricular cartilage as the wrapping
material of implant. The incidence of exposure in patients receiving auricular cartilage cap, donor sclera cap and without any wrapping material were 0, 8.33%, and 11.11% respectively. The incidence rate of mild complications, including conjunctival inclusion cysts, severe conjunctive chemosis, and conjunctive granuloma, were 27.27%, 25%, and 33.33% in patients with auricular cartilage cap, donor sclera cap, and without wrapping cap. Although no statistically significant difference was found among 3 groups (mainly due to small sample size), the incidence rate of exposure was lowest in cases with auricular cartilage cap, followed by cases with donor sclera cap. The incidence of mild complications was lower in patients with wrapping material, either auricular cartilage or donor sclera.  

The cosmetic results of simply enucleation are always poor. Therefore placing an orbital implant would be the necessary procedure in most cases to improve the appearance. However, implant exposure is the main complication after it, which could lead to poor prognosis. Rates of HA implants were reported to range from 9.6% to 28%[3,18-19]. The factors of development of exposure could mainly be categorized into 3 groups: implant related, patient related, and surgery related[20]. Thus, factors such as poor surgical technique, excessively large size of implant, and implant infection play important roles in implant exposure[21].  

In this study, we excluded patients <18 years old or >65 years old, with systemic autoimmune disease or needing radiotherapy or chemotherapy after surgery, thus to remove patient-related factor. All surgical procedure was performed by a single experienced surgeon (Chen X). A sizing ball was used during surgical procedure to evaluate the residual intraconal volume, and implant size was chosen to allow tension-free closure of the anterior ocular tissue. This procedure could prevent the usage of excessively large implant. Also, the averages of the implant size were 21.64±0.67, 21.50±0.52, and 21.67±0.50 mm in 3 groups respectively. No statistically significant differences were found among 3 groups. The factors related to poor surgical technique and excessively large size of implant were excluded as well.

Wrapping the implant was believed to significantly decrease the incidence of implant exposure by adding another layer of tissue between the implant and overlying conjunctiva[22]. We found higher exposure rate in cases with no wrapping material than cases with wrapping cap. Besides, for cosmetic purpose, it is essential not only to correct the tissue defect of these cases by inserting an orbital implant, but also to ensure the movement of ocular prosthesis. The wrapping material could provide a surface for muscle attachment which would ensure the movement of ocular prosthesis[23].  

Donor sclera is the most widely used wrapping material[7-8]. However, because of the potential spreading risk of Creutzfeldt-Jakob disease (CJD)[10], human immunodeficiency virus (HIV)[11-12], hepatitis B virus and hepatitis C virus[11], possibly occurring of late thinning of the sclera[9], and occasional unavailability of donor sclera has raise the need of other alternative materials for wrapping. These materials includes autogenous materials like fascia lata, animal-derived tissue like bovine pericardium, and synthetic materials like polyglactin 910 mesh and Mersilene mesh[6,7,11].  

Bovine pericardium as an animal-derived tissue, was initially believed to be a viable alternative to donor sclera as the wrapping material[7-8]. However, Char[22] found an even higher rate of implant exposure with bovine pericardium than donor sclera, especially at the early-stage after surgery. Another histopathologic study, which was performed in an animal model, showed that HA implants with bovine pericardium cap would had later and less extensive fibrovascularization than implants with sclera cap. They also found that bovine

### Table 2 The clinical data of patients with complications in three groups

<table>
<thead>
<tr>
<th>Cases No.</th>
<th>Complications</th>
<th>Gender</th>
<th>Age (y)</th>
<th>Cause of enucleation</th>
<th>Implant size (mm)</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group A</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Conjunctival inclusion cysts</td>
<td>M</td>
<td>41</td>
<td>Severe ocular injury</td>
<td>22</td>
<td>2wk</td>
</tr>
<tr>
<td>2</td>
<td>Severe conjunctive chemosis</td>
<td>F</td>
<td>58</td>
<td>Severe ocular injury</td>
<td>22</td>
<td>1wk</td>
</tr>
<tr>
<td>3</td>
<td>Severe conjunctive chemosis</td>
<td>F</td>
<td>63</td>
<td>Intraocular tumor</td>
<td>21</td>
<td>1wk</td>
</tr>
<tr>
<td>Group B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Implant exposure</td>
<td>M</td>
<td>65</td>
<td>Severe ocular injury</td>
<td>22</td>
<td>12mo</td>
</tr>
<tr>
<td>2</td>
<td>Conjunctival inclusion cysts</td>
<td>M</td>
<td>64</td>
<td>Severe ocular injury</td>
<td>21</td>
<td>1wk</td>
</tr>
<tr>
<td>3</td>
<td>Severe conjunctive chemosis</td>
<td>F</td>
<td>59</td>
<td>Severe ocular injury</td>
<td>21</td>
<td>1wk</td>
</tr>
<tr>
<td>4</td>
<td>Conjunctive granuloma</td>
<td>M</td>
<td>26</td>
<td>Severe ocular injury</td>
<td>22</td>
<td>1mo</td>
</tr>
<tr>
<td>Group C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Implant exposure</td>
<td>M</td>
<td>31</td>
<td>Severe ocular injury</td>
<td>22</td>
<td>6mo</td>
</tr>
<tr>
<td>2</td>
<td>Severe conjunctive chemosis</td>
<td>M</td>
<td>41</td>
<td>Severe ocular injury</td>
<td>21</td>
<td>1wk</td>
</tr>
<tr>
<td>3</td>
<td>Severe conjunctive chemosis</td>
<td>F</td>
<td>63</td>
<td>Severe ocular injury</td>
<td>21</td>
<td>1wk</td>
</tr>
<tr>
<td>4</td>
<td>Conjunctive granuloma</td>
<td>F</td>
<td>43</td>
<td>Severe ocular injury</td>
<td>22</td>
<td>2wk</td>
</tr>
</tbody>
</table>
The rub of ocular prosthesis has suitable flexibility and rigidity which could resist through a post-auricular incision. Also auricular cartilage has a spherical surface and fits well to a bulbar surface. This modification could avoid the implant exposure at the front part without inhibiting valcular ingrowth at the back part.

Mild complications, including conjunctival inclusion cysts, severe conjunctival chemosis and conjunctive granuloma, were found in this study as well. Lower incidence rate was found in cases with wrapping cap. This seems to indicate that the wrapping material wouldn’t increase the incidence of prolonged postoperative inflammation. Two conjunctival inclusion cysts were found in this study. Severe ocular injury was the cause of enucleation of both cases. The disorder of ocular tissue before surgery would lead to incomplete tissue isolation during the surgical procedure. This may be the reason of the happening of conjunctival inclusion cysts.

In conclusion, auricular cartilage can be a good choice of alternative wrapping material of orbit implant with satisfied outcome. Based on the absence of possible disease transmission, easy obtained, good biological activity and resistant of rubbing, nonabsorbable and lower incidence of late shrinking or thinning, auricular cartilage is now our favorite wrapping material for orbital implants. However, there are still weaknesses in this study, including small sample and short follow-up time. A large-sampled, longer follow-up, controlled perspective study is needed to further confirm the safety and efficacy of auricular cartilage as the implant wrapping material.

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