The Efficacy of an Acrylic Intraocular Lens Surface Modified with Polyethylene Glycol in Posterior Capsular Opacification

To investigate if the surface modification of intraocular lens (IOL) is efficient in the prevention of posterior capsular opacification (PCO), the acrylic surface of intraocular lens (Acrysof®) was polymerized with polyethylene glycol (PEG-IOL). The human lens epithelial cells (1 x 10⁶ cells/mL) were inoculated on PEG grafted or unmodified acrylic lenses for the control. The adherent cells on each IOL surface were trypsinized and counted. The every PEG-IOL was implanted in 20 New Zealand rabbits after removal of crystalline lens. The formations of PCO were checked serially through retroilluminated digital photography, and the severity scores were calculated using POCOman®. The cell adherence patterns on each IOL were examined by scanning electron microscopy. As a result, the mean number of adherent cells of PEG-IOL (3.2 ± 1.1 x 10⁶) tended to be smaller than that of the acrylic controls (3.6 ± 1.9 x 10⁶) without a statistical significance (p=0.73). However, the mean severity of PCO formation in PEG-IOL was significantly lower than that in the control during the third to sixth weeks after surgery. Scanning electron microscopy revealed that the more patch-like cells were found firmly attached to the IOL surface in control than in the PEG-IOL. Conclusively, PEG polymerization to the acrylic IOL would possibly lessen the formation of PCO after cataract removal.

Key Words: Acrylates; Cataract; Lenses Intraocular; Polyethylene Glycols

INTRODUCTION

Despite the rapid progress made in modern cataract surgery, posterior capsular opacification (PCO) still remains as one of the most common postoperative complications. Usually the opacification is minimal and confined to the periphery and is not of great concern. However, in some severe cases, it impairs vision and can make previous cataract removal worthless. Thus, additional intervention is often necessary for visual recovery. The most well known method to overcome PCO is neodymium:YAG capsulotomy, which is straightforward and immediately effective, but perhaps the best way of dealing with PCO is to prevent its occurrence. As PCO is thought to be formed by the migration and metaplasia of lens epithelial cells (LECs), it may be beneficial to remove intraoperatively as much of the cortical remnant and LEC nests as possible, though admittedly there is some dispute in this regard (1-3).

Many theories have been put forward to preclude the formation of posterior capsular opacities, and attempts to modify interactions between lens capsules and intraocular lens (IOL) surfaces are abundant. Recently lens design, especially the configuration of the optic edge have been found to be most effective at preventing PCO, as the edge itself could act as a mechanical barrier against LEC migration and could induce capsular bending (4-6). Many attempts have also been made to identify an ideal lens material and gradually it has been accepted that hydrophobic acrylic plastic offers the best means of reducing PCO and postoperative inflammation (7). Moreover, the lens surface may also play a primary role in LEC adhesion, and thus, development programs have been initiated to produce less adhesive surfaces. However, few results are available due to the tardy progress made in the synthesis of biocompatible materials (8-11). In this study, we investigated the effect of surface modification with polyethylene glycol (PEG); a type of hydrophilic polymer that is known to be effective at reducing protein deposition and cell adhesion. In our previous studies, graft polymerization of PEG to polymethylene methacrylate (PMMA) was found to be effective at reducing keratocyte adhesion, and therefore, we further investigated the efficacy of a PEG graft on an acrylic lens with a square edge, which is presumed to be the best design and materials in terms of reducing PCO (12).
MATERIALS AND METHODS
Preparation of a PEG grafted acrylic IOL (PEG-IOL)

Acrylic lenses with a square edge of 6 mm optic size (SA 60AT®, Alcon) were used, and graft polymerization was done as described in our previous study (12). Briefly, the lens surface was first treated with oxygen plasma, which was generated under low pressure (70 torr) at 100 Watt, and then it was incubated in PEG solution at 60°C for 24 hr.

In vitro human lens epithelial cell adhesion assay

IOLs were sterilized with ethylene oxide gas and washed with phosphate buffered saline (PBS) before use. The two haptics were severed and the remaining optic portions were placed at each bottom of 96-well culture plate. As many as $1 \times 10^4$ immortalized human lens epithelial cells (B-3, ATCC, U.S.A.) were layered onto the optics and incubated for 24 hr at 37°C in 5% CO2. The medium was then removed and IOL surfaces were washed twice carefully with PBS to remove non-adherent cells. The 0.25% trypsin (200 μL) was then added and incubated for 10 min at 37°C to harvest adherent cells. Cells were then placed in the mixture of 400 μL of PBS and 400 μL of minimal essential medium (MEM) containing 20% fetal bovine serum, and finally centrifuged at 1,500 rpm for 5 min. Pellets were resuspended in MEM (200 μL), stained with trypan blue and counted in a hemocytometer (Superior/Marienfeld, D-97965, Bad Mergentheim, Germany). A total of nine samples were tested per group. To verify statistical significances, we used the Mann-Whitney U test.

In vivo PCO formation assay

In accord with the principles of ARVO Statements on the Use of Animals in Ophthalmic and Vision Research, 20 PEG grafted acrylic lenses (PEG-IOL) and 20 control uncoated lenses (SA 60AT®) were implanted randomly into either eye of male New Zealand white rabbits weighing from 2.5-3.0 kg. Briefly, animals were anesthetized with an intramuscular injection of ketamine hydrochloride (100 mg/kg) (Yuhan, Seoul, Korea) and xylazine (5 mg/kg) (Rompun®, Bayer, Leverkusen, Germany). Pupils were dilated with tropicamide (Mydriacyl®, Alcon, Fortworth, U.S.A.) 3 times at 10-min intervals prior to surgery. Operations were conducted under sterile condition using a Zeiss surgical microscope by one surgeon. The anterior chamber was entered through a 2.75 mm sized superior self sealing clear corneal incision, and side entry was made 90 degrees apart from the initial incision using a 15° standard angle blade. Using a visosurgical device (Viscoat®, Alcon), a 4.5 to 5.0 mm round, well centered continuous curvilinear capsulotomy was made with a capsulorhexis forceps, such that the anterior capsular margin covered the whole optic margin completely. A complete hydrodissection followed by hydrodelineation was performed and the remaining lens material was removed by phacoemulsification (Premier®, Storz, Germany). The capsular bag was fully inflated with viscoelastics, and a random IOL was inserted using an injector (Monarch®, Alcon, Korea). Then IOL was positioned in the desired central position with 360 degree anterior capsular coverage. After removal of the remaining whole viscoelastics, the wound was closed with one 10-0 monofilament nylon suture in all cases. Because the rabbit cornea is relatively thin, it was sometimes difficult to make a complete self sealing clear corneal incision. Postoperatively a mixture of dexamethasone, neomycin, and polymixin B sulfate ointment (Maxitrol®, Alcon, Fortworth, U.S.A.) was applied 3 times daily for 7 days. Complications were evaluated at weekly and standardized retro-illuminated photographs were taken through a dilated pupil until the 8th postoperative week by the same examiner using a light source with the same size and angle.

Fig. 1. The severity grades of posterior capsular opacification with the POCOman software. A retroilluminated picture (A) is transformed into a black and white picture (B). The margin of the anterior capsule should be marked then the grid is allocated automatically. Using the standard photographs (grade 1-3), the severity score is calculated with the formula as follows: severity score=3*Red [grade3]+2*Yellow [grade2]+1*Blue [grade1]/total area.
To nullify the effects of the anterior capsule in the formation of PCO, we excluded those rabbits with implanted lens optic margins not covered completely by the anterior capsule or extruding into the anterior chamber. The severity of PCO was graded using semi-objective calculation software (POCOman) from St. Thomas’s Hospital and the King’s College London imaging process group (Fig. 1). Results were compared using the nonparametric rank summation technique and the Mann-Whitney U test, and significance was accepted at the 5% confidence level. At the end of our study, 8 weeks after implantation, all animals were sacrificed and implanted IOLs were carefully removed. The IOLs were again fixed in glutaraldehyde (2.5%), washed with 0.1 M/L PBS, dehydrated in graded ethanol, critical-point-dried in liquid CO₂ (CPD 7501), and coated with gold (Giko-IB-3) for scanning electron microscopy (JSM-300, Joel, Tokyo, Japan). Adherent cells were counted in 5 or more fields at 200× and statistical significances between study groups were determined using the Student’s t-test.

RESULTS

In vitro human lens epithelial cell adhesion assay

The mean number of adherent LEC’s harvested from each lens surface was $3.6 \pm 1.9 \times 10^3$ for the control and $3.2 \pm 1.1 \times 10^3$ for the PEG-IOL, which was not significant ($p=0.73$, Fig. 2).

In vivo PCO formation assay

No intraoperative complications developed, e.g., capsular tear. However, during follow-up, some parts of IOLs were
found to have extruded outside the capsular bag in 7 (4 in the control, 3 in the PEG-IOL group), and thus complete anterior capsular envelopment over the optic was impossible. These rabbits were excluded from the study. Minimal anterior chamber reactions and mild corneal edema were resolved during the first week. The formation of PCO was evident from the third week after surgery and grew weekly until the last follow up. In the PEG-IOL group, PCO began to take on a fine fibrotic appearance rather than a conglomerated cellular proliferation like the Elschnig’s pearl in the control group, and became a similar figure later on (Fig. 3A, B). The PCO formation was significantly lower during the third to sixth weeks in the PEG-IOL group (Table 1, Fig. 3C, Mann-Whitney- \( U \) test). However, during the final two weeks (weeks 7 and 8), no significant differences were found, and some rabbits showed marked LEC proliferation starting from the optic-haptic junction, regardless of group (Fig. 4). Scanning electron microscopic examinations of IOL surfaces revealed fewer adherent LECs on PEG-IOLs. Mean numbers were 26.4 ± 13.7 for the control and 16.2 ± 11.2 for the PEG-IOL, which were significantly different (\( p = 0.041 \), Mann-Whitney- \( U \) test). Regarding the appearance, more cells were found to be firmly attached to control lens surfaces with a patch-like figure (Fig. 5).

### DISCUSSION

The formation of PCO is believed to be a multifactorial process. The surgical technique, lens material, and configuration including optic edge design and haptic angulations are thought to play important roles in PCO formation, though patient condition and the postoperative administration of drugs may also have an influence (4, 6, 13, 14). However, there are many different opinions concerning the pathogenesis of PCO (2, 3). In general, the optic material is not regard-

![Table 1. The changes of severity score calculated from POCO-man](image)

|                | Control (n=16) | PEG (n=17) | \( p \)-value |
|----------------|---------------|------------|---------------|
| 1 week         | 0.136         | 0.136      | 1             |
| 2 week         | 0.38          | 0.304      | 0.548         |
| 3 week         | 0.588         | 0.322      | 0.008*        |
| 4 week         | 0.794         | 0.472      | 0.096*        |
| 5 week         | 0.944         | 0.468      | 0.016*        |
| 6 week         | 1.086         | 0.706      | 0.032*        |
| 7 week         | 1.668         | 1.152      | 0.222         |
| 8 week         | 1.854         | 1.448      | 0.151         |

\( * \), \( p < 0.05 \) by Mann Whitney-\( U \) test.

At all time periods, the posterior capsular opacification in PEG-IOL tended to be less severe compared to the control, but the statistical significance was lost again 7th week of examination.

![Fig. 4. An enormous proliferation of lens epithelial cells through the optic-haptic junction. (A) 4th week. Few cells were seen except small cell buds at the optic-haptic junction (arrowhead). (B) 6th week. The cell nests were rapidly growing and threatening the visual axis. (C) 8th week. The visual axis was completely obliterated by enormously proliferating cells.](image)
ed so important as optic configuration in the pathogenesis of PCO, although the adhesion of various inflammatory cells and fibronectin is clearly optic material and surface-dependent (15, 16). Thus, surface modifications, including heparin incorporation, have been examined, and some reports have shown favorable results under selective conditions (8-11). We undertook this study to investigate the effect of surface modifying acrylic lenses with PEG, which is a high molecular weight hydrophilic polymer and is tethered on the surface, and thus the remainder of the molecule is free to move and can prevent the deposition of protein materials and inflammatory cells (17-19). In spite of some debates, hydrophobic acrylic lenses are known to have an affinity for fibronectin, and thus, we consider that PEG grafting might have a beneficial effect (20). Moreover, although acrylic lenses are better at preventing PCO than other type of IOL, PCO formation can also be influenced profoundly by other physical characteristics of IOL, and therefore, in the present study we compared the effect of PEG surface grafting on optic materials of same design (7).

In our previous study, PEG-grafted PMMA had a markedly lower contact angle and reduced cellular adhesion (12). However, in the present study, we found no significant differences between the two study groups in vitro, though a tendency for cells to adhere less was observed for the PEG-acrylate. We consider this is due to the characteristics of materials that compose the IOL. Since acrylates are soft and their hardness can change markedly with the sudden drop in temperature during washing, their adhesiveness might also be changed accordingly.

When PCO formations were compared, it was found to be significantly less severe in the PEG-IOL group than the in control group from the third week postoperatively, when sufficient amount of time passed for the cells to migrate and proliferate themselves. Moreover, the opacification appeared as a fine fibrosis in the PEG-IOL group, suggesting reduced visual impairment for the PEG-IOL. However, this PCO lowering effect did not last for long, showing no differences from seventh week of examination. And some rabbits in both groups revealed enormous LEC proliferation starting from the optic-haptic junction, and soon covering throughout the optic surface. The acrylic lens used in our study is a single piece and considered to be better at stretching the posterior capsule because it exerts even forces on the posterior capsule after insertion into the capsular bag, and at the same time avoids early posterior capsular striae, which is believed to be the route of early LEC migration (21, 22). However, the junction between optic and haptic is large and there is no definite barrier, and thus, it can act as a gate for LEC migration. In this study, LECs entered the junction at around 1 month and spread very rapidly to the whole optic surface (Fig. 3). The amount and rapidity of this migration were so great that after six weeks our results might have lost statistical significance. This demonstrates that such modifications of IOL surface could work during the early phase of PCO formation and that the lens configuration might have a greater effect after a certain time period, as PCO formation is a multifactorial process.

The limitation of our study is that it involved a relatively small number, and thus, a further study using a larger sample number is needed to confirm the findings of this study. Another limitation is that as we used only one type of IOL, our results cannot be generalized to other lenses with different contours. So it is also needed to study various lens materials and configurations to elucidate their efficacies in terms of preventing PCO formation. Although the formation of PCO is multifactorial and it seems to be impossible to consider its all causative factors in one study, the inflammation induced by the surgery is sure to have significant influence. To investigate its role, more frequent examinations in early postoperative phases are also needed.

In conclusion, the surface modification of acrylic IOL with PEG may reduce the formation of PCO during the early stage.
However, the multifactorial nature of PCO means that we should always consider that other contributors are likely to be important.

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