**The Role of Type IV Collagen in Developing Lens in Mouse Fetuses**

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**Abstract**

**Objective(s)**
Extracellular matrix (ECM) and basement membrane (BM) play important roles in many developmental processes during development and after birth. Among the components of the BM, collagen fibers specially type IV are the most important parts. The aim of this study was to determine the time when collagen type IV appears in the BM of lens structure during mouse embryonic development.

**Materials and Methods**
In this experimental study, 22 female Balb/C mice were randomly selected and were kept under normal condition, finding vaginal plug was assumed as day zero of pregnancy. From embryonic day 10 to 20, all specimens were sacrificed by cervical dislocation and their heads were fixed, serially sectioned and immunohistochemistry study for tracing collagen type IV in lens were carried out.

**Results**
Our data revealed that collagen type IV appeared at the early stage of gestation day 12 in BM of anterior epithelial lens cells and the amount of this protein gradually increased until days 15-17 in ECM and posterior capsule epithelium. After this period, severe reaction was not observed in any part of the lens.

**Conclusion**
These findings establish the important role of collagen IV in developing optic cup and any changes during critical period of pregnancy may be result in severe visual system defect

**Keywords:** Basement membrane, Collagen IV, Lens capsul

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Introduction
Lenses are transparent and flexible components of the optic system which develop along with the other parts of the eyeball. In addition, not only do they direct light into the environment of the visual system, but they also have the potential for accommodation. The presence of elastic components in early stages of lens development facilitates the accommodation of the lens capsule to physical characteristics such as strength, thinness, and recoil (1). The previous studies show that collagen type IV is a major protein of the lens structure consisting of subunits \( \alpha_1-\alpha_6 \). The chains \( \alpha_3, \alpha_4, \alpha_5 \) are cross-linked with each other during lens development and provide the strength required to accommodate the lens capsule (2). Hence, considering that sufficient appearance of collagen guarantees proper function of the lens (3), one can conclude that in the process of natural development the lenses are more flexible in newborns than adults. This causes the accommodation and transmission of light to be more facilitative in newborns than grown-ups who have missed their accommodation or suffer from diseases such as cataract (4).

Researches have shown that defects in collagen type IV expression can cause congenital defects of the optic system such as Alport syndrome (5, 6). In this case, although complex of abnormalities such as macula defect or detachment of the retina appears, vision defects such as conical anterior and posterior in lens capsule are caused (7). The absence of collagen subunits \( \alpha_3, \alpha_4, \alpha_5 \) causes not only deformation of lens but also fragility and rupture of the capsule (8). This is one of the most important processes related to Alport syndrome (9). Based on different studies, various types of proteins are comprised in lens capsule structure such as entactin/nidogen, laminin, heparin sulfate, glycoproteins, and collagen type XVIII. Although we should not ignore the structural role of lens proteins, collagen type IV is the most abundant type of collagen in lens and plays a critical role in lens development (10). It consists of triple helix structure, known as protomers, each of which forms tropocollagen (11). The helix consists of six subunits; each is expressed by a separate gene (12). Mutation in any of the genes can result in defect of collagen synthesis and bring about a change in components of eye structures. Therefore, it is necessary to investigate its formation and structural molecule during lens development and its association with ocular defects. The present study has investigated the collagen type IV distribution and expression patterns during lens development.

Materials and Methods
Twenty two pregnant female Balb/c mice of 35 g body weight were obtained from animal house of Mashhad University of Medical Sciences, the environmental conditions were 23-25 °C, relative humidity 50-55%, 12-hr light-dark cycle (light on at 6.00 am). Female mice were mated with males of the same strains (two females and one male) overnight and examined the next morning for vaginal plug. Females with sperm-positive smears or vaginal plug were designated as day zero of pregnancy. Two pregnant mice from gestational days E10 to E20 were anesthetized with chloroform and perfused transcardially with formaldehyde 10% and the head of fetuses were removed and fixed for 24 hr at room temperature in the same fixative. The head of fetuses routinely processed and embedded in paraffin. Then eight-micron sagittal sections were carried out serially. Interval sections were one from each ten. Sections were stained with cresyl violet or incubated with monoclonal antibody (anti-collagen IV/ Dako Co.).

Immunohistochemistry study
The Avidin-Biotin peroxidase procedure was used for immunohistochemistry study. Sections deparaffinized, rehydrated and washed twice for 5 min in 0.05 Tris buffer containing 1.5% NaCl, pH 7.4. For blocking nonspecific antibody, sections were preincubated in 0.3% Triton X-100 in TB-NaCl followed by 5% goat serum for 2-3 hr. Then sections were reacted for 12-24 hr at 4 °C with primary antibody (anti-collagen IV) diluted 1:250 in TB-NaCl with 0.3% Triton and 2% serum. Tissues were washed...
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with TB-NaCl for three times, each for 10 min and incubated for 2 hr in biotinylated goat anti-rabbit IgG (1:400 in TB-NaCl). After three further rinses, each for 1 hr, endogenous peroxidase activity was blocked by incubating in 0.03% H2O2 in methanol for 30 min. Tissues were incubated for 2 hr in 1:100 avidin-biotinylated horseradish peroxidase complex. Then they were washed three times, each time for 30 min in TB-NaCl and finally reacted with 0.03% solution of 3,3-diaminobenzidine tetrahydrochloride containing 0.03% H2O2 for 10-15 min. Tissues were washed and were lightly counterstained with hematoxylin. Subsequently, they were washed, air-dried, dehydrated, and then mounted in PBS glycerol. Photographs were taken by a bh2 Olympus light microscope and collagen IV reaction in lens components in each of embryonic days was evaluated.

Statistical analysis
The data were analyzed by using SPSS software and Kruskal Wallis and Mann-Whitney tests. P-values<0.05 was considered as significant.

Results
Our data indicated that primary lens was visible in anterior capsule epithelium of optic cup as an undifferentiated cellular mass on day 11 of gestation (Figure 1a). On day 11, although anterior part of the lens is formed, collagen indicated no reaction in basement membrane (Figure 1b). On day 12, most of the primordial lens cells were degenerated and replaced with lens matrix. Based on our finding, collagen IV appeared at the early stage of embryonic day 12 during critical period of the developing lens. In addition, this study indicated high levels of collagen IV expression at the BM of anterior epithelial cells of the lens during early development of mouse lens (E-15 to E-17), while anterior epithelial capsule was growing. After that, the structure of BM was completed in this area and collagen IV became stable after this period. From day 14, intensity of collagen expression in lens significantly increased compared to embryonic day 11. On day 15 of gestation, the reaction of anterior epithelium remarkably increased and expanded into the lens nucleus but decreased in this region and there was weak reaction in posterior lens capsule (Figure 1c and 1d). On day 17 of gestation, collagen fibers were expanded into adjacent lens nucleus (table 1). In this case, although posterior lens capsule was labeled with antibody against collagen, narrow band of posterior capsule epithelium indicated no reaction except for the remains of primordial cells in this region (Figure 1e and 1f).

![Figure 1](image-url)
Table 1. Lens components collagen IV reaction during embryonic development

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<th>Day embryo</th>
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<th>Lens nucleus</th>
<th>Anterior pole</th>
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This gradation was scored ranging from negative to 4 positive in conformity with the severity of reaction from negative, weak, moderate, strong and very strong.

Discussion

Basement membranes are specialized extracellular matrices consisting of tissuespecific organizations of multiple matrix molecules and serve as structural barriers as well as substrates for cellular interactions. Collagen fibers specially type IV are the most important part of this area. Our immunohistochemistry studies showed that there is no reaction of collagen type IV until day 12 of gestation. Another investigation also revealed that first collagen IV expression in mouse was detectable in lens capsule as early as embryonic day 11.5 (13, 14). The appearance of first signals of collagen type IV in basement membrane of the anterior capsule represents an important role in lens development because anterior capsule consists of specialized basement membrane to which epithelial cells bind (15). As the structural compositions of the lens are completed, the specific role of collagen becomes distinguished in this part of visual system. Since the marginal zone of capsule binds to zonules which connects capsule to ciliary body using stable and thin fibers (16), the strength force diffuses to the lens surface. So compositions such as collagen are required to provide stability and flexibility (17). Besides serving structural role, collagen type IV may transmit signals for natural eye development (18). In other words, subunits of collagen (IV) α₁, α₆ in pre- and postnatal stages showed that first collagen is formed in anterior and posterior capsule epithelium and then is expanded to the other parts of lens until day 16.5 of gestation (22).

The results of this study showed that density of collagen in posterior capsule epithelium is lower in contrast to other parts of the lens except for the patchy distribution of embryonic cells observed in this region. Although these cells disappear gradually to provide transparency of the lens in early stages of postnatal, the collagen causes induction effects on apoptosis. However, weak reaction of collagen in posterior capsule epithelium didn’t accelerate the apoptosis in embryonic period. In diseases such as Alport syndrome changes in collagen expression result in ocular defects such as cataract. These changes can result in a dense collagen in the aged and diabetic individuals as well as cataract.
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References