A Study of the Vascular Network of the Iris Using Flat Preparation

Yumi Song, MD, Yoon Joong Song, MD, Myung Kyoo Ko, MD
Department of Ophthalmology, Hanyang University Hospital, Seoul, Korea

Purpose: This study was performed to examine the vascular network of the human iris using flat preparation.

Methods: The ciliary body-iris structures were separated from human eyeballs, and a portion of the irises were treated with trypsin to remove the pigment granules. These iris tissues were unfolded and placed onto glass slides using flat preparation, and the vascular network of each iris was examined by fluorescein microscopy. The ciliary body-iris structures separated from the remaining eyes were stained with hematoxylin-eosin without trypsin treatment and were examined by light microscopy.

Results: The long posterior ciliary artery formed several branches before entering the iris root, and such branches formed the major arterial circle of the iris with diverse diameters in the vicinity of the iris root and the ciliary process. In the pupillary margin, the iris vasculature network formed a cone shape and then formed an arcade by connecting to adjacent vasculatures. In the vicinity of the collarette, the iris vasculature network formed the minor arterial circle of the iris with diverse diameters perpendicular to the arcade of the iris network located in the pupillary margin. In the pupillary margin, the capillaries were somewhat thick and connected to the irregular traveling iris vein.

Conclusions: The above findings explain the human iris vascular network and provide a theoretical basis for the sectoral filling of the iris vasculature seen in fluorescein iris angiography.

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Key Words: Flat preparation, Fluorescein microscopy, Human iris, Vascular network

The iris is connected to the sphincter and dilator muscles of the eye, which automatically change the size of the pupil depending on the amount of light available to the eye and the position of the eye. In addition, the iris is in the most anterior portion of the uvea and contains abundant vascular networks. It forms a delicate and movable diaphragm between the anterior and the posterior chambers.

The iris has abundant vasculature, and disruption of the blood-aqueous barrier induces iritis. Melanocytes and fibroblasts are present in the iris stroma, and the pigmented epithelium is on the posterior side of the iris. In fluorescein iris angiography, fluorescein dye cannot penetrate the iris vessel wall because of the blood-iris barrier; nonetheless, in the ciliary body fluorescein dye penetrates the ciliary vessel wall. The leakage of fluorescein dye cannot be detected in the iris vessel of a normal eye under the age of 40 years. However, there is often fluorescein leakage at the pupillary margins in people older than 40 years, especially after 50 years of age.

Iris vasculature has abundant blood flow and plays an important role in iritis. Thus, studies on the iris vascular network have been previously conducted. However, most anatomical studies on the human iris vascular network have been performed in vivo with fluorescein iris angiography. Other studies have been conducted in vitro and were performed with scanning electron microscopy after the injection of resin into the human iris vascular network.

This study was conducted in order to assess the human iris vascular network as closely as possible to its normal state. Applying a flat preparation, the iris vascular network was examined by fluorescent and light microscopies.

Materials and Methods

Human cadaver eyes were obtained from neonates (under one year of age), which have relatively low amounts of pigment granules in the uveas. No external malformation were found on gross examination. Each eyeball was physically divided into two parts at the equator, and the crystalline lens was removed. In the anterior portion of the eyeball, the ciliary body was carefully removed from the sclera by dissection with microforceps. Then the iris was separated from the ciliary body at the iris root (Fig. 1).

Pigment granules of the uvea make microscopic examination of the iris vascular network difficult. To remove the pigment granules from the dissected irises, the specimens were treated with trypsin using the method of Kuwabara and Cogan. The tissues were treated with 0.1 M Tris
buffer (pH 7.8) containing 3% trypsin (1 : 250) and were placed in a 36°C incubator for 30 minutes. After incubation, the pigment epithelium and melanocytes were partially removed from the iris surface with a fine brush.

The partially depigmented irises were spread onto glass slides and dried at room temperature. Dehemoglobinized red blood cells within the blood vessels of the iris show pseudofluorescence; thus, it is possible to identify the shape of the blood vessels under fluorescent microscopy. The slides in this study were examined under a fluorescent microscope.

**Fig. 1.** Light micrograph of the human iris vasculature: the iris was separated from the ciliary process using flat preparation (×40).

**Fig. 2.** Light micrographs of the human iris vasculature. (A) The branch (arrowheads) of the long posterior ciliary artery (arrow) enters the iris through the ciliary body (×100). (B) The major arterial circle of the iris (arrows) was observed around the iris root, and it ran toward the pupillary margin (flat preparation) (×100).

**Fig. 3.** Fluorescent micrograph of the human iris vasculature: the vascular network of the iris consists of arteries, veins, and capillaries of the iris root and pupillary margin (flat preparation) (×100).

**Fig. 4.** Fluorescent micrograph of the human iris vasculature: large sized vessels (arrow), probably veins, and relatively small sized vessels, the major circle of the iris (arrowheads), were observed around the iris root (flat preparation) (×200).
Fig. 5. Fluorescent micrograph of the human iris vasculature: the major arterial circle of the iris (arrows) around the ciliary process is composed of large vessels, which are perpendicular to the iris vessels (flat preparation) (×200).

Fig. 6. Fluorescent micrographs of the human iris vasculature: the vascular network of the pupillary margin forms the arcade, where capillaries continue to veins. The minor arterial circle of the iris (arrows) runs perpendicularly to the arcade (arrowheads) of the vascular network of the pupillary margin (flat preparation) (×400).

Fig. 7. Fluorescent micrographs of the human iris vasculature: the arcade of the vascular network of the pupillary margin forms an anastomosis (arrow) with adjacent vessels (flat preparation) (×400).

Fig. 8. Fluorescent micrographs of the human iris vasculature: the vascular network and minor arterial circle of the iris were observed between the iris root and the pupillary margin. The iris vessel showed an irregular course and got larger as it approached the iris root (flat preparation) (×100).

with a fluorescein isothiocyanate filter (450 to 500 nm).

The irises of the remaining eyeballs which were not treated with trypsin solution were spread onto glass slides, dried at room temperature, and stained with hematoxylin-eosin. Then the slides were examined under a light microscope.

Results

The ciliary body and iris were dissected from the sclera and were separated at the iris root (Fig. 1).

The long posterior ciliary artery (LPCA) branches in the
Fluorescent micrograph of the human iris vasculature: the montage of vascular networks of the iris and the minor arterial circle of the iris. The blood vessels of the minor arterial circle of the iris did not form a complete circle.

Discussion

In this study, the ciliary body and the iris were separated during preparation of the iris. Therefore, the contribution of the anterior ciliary artery (ACA) to the formation of the major arterial circle of iris could not be assessed. However, it was observed that the LPCA that travels from the ciliary body to the pupil was branched into a V shape and formed the major arterial circle of the iris near the iris root.

The ACA flows mostly to the anterior uvea. It travels in parallel to the rectus muscles, penetrates the sclera, and forms intramuscular loops by generating many branches in the external layer of the ciliary muscles. It also forms the major arterial circle of the iris together with the LPCA, which is the major blood supplier to the iris and the ciliary body.

Hayreh et al. have reported that the temporal branches of the ACA and LPCA have less influence on the blood supply to the temporal portion of the iris than do the nasal, superior, and inferior portions, based on fluorescein iris angiography taken after strabismus surgery. Funk and Rohen have reported that the ACA and LPCA are connected in almost all areas of the human postmortem eyeball. Nevertheless, the results of vascular casting performed on these blood vessels selectively showed that most of the area of the major arterial circle of the iris was supplied from a branch of the LPCA.

In the current study, the iris artery was observed to travel from the major arterial circle of the iris toward the pupillary margin with a regular interval and diameter. Conversely, Shimizu and Ujiie have reported that the iris artery has a regular diameter and interval, and that it shows a difference in shape from the blood vessels of the ciliary processes. The veins and capillaries of the ciliary processes are large and irregularly dilated and therefore have a sinusoid appearance. The minor arterial circle of the iris was located in the pupillary constrictor muscle and was observed to travel perpendicularly to the arcade of the vascular network of the iris with diverse diameters. This minor arterial circle of the iris is formed by a branch from the iris artery traveling from the iris root to the pupil. The branches from the minor arterial circle of the iris travel to the pupillary margin and form the capillary arcade. However, according to the Shimizu and Ujiie, blood vessels comprising the minor arterial circle of the iris were detected intermittently and reached only a few hundred micrometers, and thus, did not form a complete circle. In our study, the minor arterial circle of the iris was also
observed to not form a continuous circle.

The iris vein ran on a path similar to that of the artery, but as it got closer to the iris root, the vein’s diameter became larger and thus could be distinguished from the iris artery.

In fluorescein iris angiography, during the arterial phase, fluorescein first fills the nasal side, and then fills the temporal side. A possible explanation of this finding was reported by Amalric and Rebiere who explained that the preferential filling was due to the distribution of blood vessels being most dense in the superior-nasal sector. In addition, Hayreh and Scott showed that blood vessels were evenly distributed in all areas, but in some eyeballs the blood vessels might be more densely distributed in the nasal side.

Fluorescein iris angiography often shows sectoral filling patterns. Hayreh and Scott reported that such sectoral filling is physiological and is most probably due to the fact that the various supply arteries to the iris arise at different points along the ophthalmic artery and show a large amount of variation from eye to eye.

In our study, the distribution of blood vessels was observed to be similar in each area of the iris. Several branches of the LPCA showed a V-shape prior to entering the iris root, and they formed the major arterial circle of the iris. Based on these findings, it is likely that sectoral filling in fluorescein angiography is due to the difference in time that it takes for fluorescein to fill all of the vessels. First, the iris arteries that continue to the branches of the two LPCA are filled. After that, fluorescein fills the major arterial circle of the iris and the connected iris arteries. Such a finding is in disagreement with the observations of Amalric and Rebiere, and Hayreh and Scott. In addition, the variation in diameter of the major arterial circles of the iris may explain the time difference in fluorescein filling from the major arterial circle of the iris to the iris artery.

In our study, iris capillaries were observed distinctly in each area and respect to the ciliary process. The abundant arteriovascular network in the pupillary constrictor muscle implies active nutritional support and metabolism in this area and is considered to be associated with hippus.

Because our study used flat preparation, as opposed to vascular casting which allows observation of blood vessels in three dimensions, the distribution of blood vessels could not be evaluated by distinguishing the anterior and posterior parts.

In vascular casting, resin is injected into the blood vessels, the remaining tissues are enzymatically removed, and the frame is examined three-dimensionally with a scanning electron microscope. In our study, the shapes of the iris blood vessels were examined without removing the vascular tissues. Observation of the iris blood vessels under microscopy is made difficult by pigment epithelium and pigment granules of melanocytes. We treated the tissues with trypsin, an established method for removal of retinal pigments. Importantly, since dehemoglobinized red blood cells show pseudofluorescence, we were able to use fluorescein microscopy to observe the iris vascular network, providing a new view of this system.

Through anatomical observation of the iris vascular network, this study provides a theoretical basis for the sectoral filling of fluorescein in iris angiography.

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