The Missing Piece in Glaucoma?

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Abstract

Glaucoma is defined as an optic disc neuropathy meaning the nerve fibers are being atrophied similar to the fate occurring in non-glaucomatous optic atrophies. Furthermore, the nerve fibers are always being destroyed randomly in all the non-glaucomatous optic atrophies. In contrast, the nerve fibers in glaucoma are invariably destroyed in an orderly tandem fashion, from peripheral to central, never randomly. Is glaucoma really an optic disc neuropathy in light of orderly destruction of nerve fibers in glaucoma? The current prevailing theories in glaucoma such as posterior bowing of the lamina cribrosa or cupping can’t explain the orderly destruction of nerve fibers occurring in glaucoma. In fact, there is no biological mechanism acting directly on the nerve fibers or their RGCs which could lead to their orderly destruction. Therefore, there should be some mechanical way, which could result in the orderly destruction of nerve fibers even though this mechanical scenario may have resulted from the direct biological effect of raised IOP on some important component of the optic disc. It is proposed that the border tissue of Elschnig (BT) atrophies due to chronic ischemia caused by raised IOP, and as a result, the lamina cribrosa (LC) begins sinking in the scleral canal—a mechanical problem. Due to sinking of the LC, the nerve fibers get stretched and broken starting with the most peripheral nerve fibers being closest to the edge of the scleral opening and ending with the most central nerve fibers in an orderly tandem fashion. Therefore, in view of the orderly destruction of nerve fibers, glaucoma may not be an optic disc neuropathy but an optic disc axotomy.

Keywords

Glaucoma, Normal Tension Glaucoma, Severance, Arcuate Field Defects, Disc Notching, Disc Hemorrhage, Sinking Disc, Cupping Disc, Excavated Disc, RNFL

1. Introduction

Chronic glaucoma has been a mystery ever since it was given a separate entity 160 years ago [1]. Ophthalmolo-
gists of the time found the optic discs of chronic glaucoma subjects being cupped, instead of being normally flat. It is presumed that the disc becomes cupped due to the force of raised intraocular pressure (IOP) resulting in atrophy and shrinkage of the nerve fibers. Until now, cupping of the disc and atrophy of the nerve fibers are still considered the salient features of the glaucomatous disc and thus glaucoma is defined as an optic disc neuropathy.

However, the concept of cupping disc and atrophy of the nerve fibers has failed to answer many pathognomonic features of glaucoma including predictable visual field defects. I believe the concept of cupping was given mistakenly in the 1850s, which has put us on the wrong path in glaucoma. One hundred years later, the term cup/disc ratio was introduced which gave further credence to the cupping concept but at the same time created conundrum in glaucoma diagnosis since ironically we started using the same parameter of “cupping” in describing both physiological as well as glaucomatous cupping of the disc.

It is imperative to mention the arrangement of nerve fibers in this discussion. There are three main aspects in which the nerve fibers are arranged in the retina and in the optic disc. First, the nerve fibers in the retina are arranged in layers superficial to deep. Second, the most central vision fibers originate closest to the disc, like most superficial (closest to vitreous) and exit from the most central part of the disc. In contrast, the most peripheral nerve fibers originate from the most distant retina or farthest from the optic disc, lie deepest (closest to sclera) and exit closest to the edge of the scleral opening. Figure 1. Third, the nerve fibers originating from the nasal retina proceed directly to the nasal part of the optic disc. However, the situation is different in the temporal retina because of the presence of the macular fibers. The nerve fibers originating from the nasal aspect of the macular area proceed directly to the central temporal part of the disc. The fibers originating from the temporal macular and peripheral retina have to arch above and below the macular fibers to reach the superior and inferior poles of the optic disc respectively and hence are known as the arcuate fibers.

We may differ on many aspects in glaucoma but on one issue we all have consensus: that the million or so nerve fibers densely packed in a disc are always invariably being destroyed, one by one, from peripheral to central in an orderly tandem fashion, and never randomly. If the nerve fibers are not destroyed in a predictable order sequence, the role of visual field tests in glaucoma would be meaningless.

The cupping disc/atrophy of the nerve fibers paradigm has failed to explain the orderly destruction of nerve fibers [2], which is hallmark of glaucoma. For any glaucoma theory to prevail, it must explain the orderly destruction of nerve fibers otherwise it will not be valid. In light of the orderly destruction of nerve fibers all the prevailing glaucoma theories such as the direct role of raised IOP, apoptosis, neurodegeneration [3], increased sensitivity of the disc to IOP, posterior bowing of the lamina cribrosa or cupping become wrong as none of them can explain the orderly destruction of nerve fibers occurring in glaucoma. If the nerve fibers are being destroyed
in an orderly tandem fashion in glaucoma, then we should expect the mechanism for their destruction to be orderly as well.

In fact, there is no biological mechanism which acting directly on the nerve fibers or their RGCs could result in their orderly destruction. Therefore, for the orderly destruction of nerve fibers to occur in glaucoma, there must be some mechanical way even though that mechanical scenario may have resulted from the direct biological effect of raised IOP on some very important component of the optic disc.

2. What May Be the Mechanical Scenario?

The border tissue of Elschnig (BT) keeps the LC firmly in place in the scleral opening. The LC is sinking due to atrophy of the border tissue which is solely supplied by ciliary circulation. Systemic circulatory pressure supplying the BT and IOP are opposing forces. Normally, the circulatory pressure supplying the BT should be higher than the IOP for the proper perfusion and healthy maintenance of the BT, Figure 2.

However, if this delicate situation is reversed, either due to raised IOP or if pressure supplying the BT becomes lower than the IOP due to systemic problems such as chronic hypotension, then even normal range IOP can take the upper hand. Thus, even the normal range IOP will begin compressing the circulation of the BT thereby inducing chronic ischemia and its atrophy and NTG will ensue. Therefore, it is the IOP becoming higher than the circulatory pressure of the BT resulting in both HTG and NTG.

The eyeball is supplied by dual circulation, the central retinal artery (CRA)—a high pressure system and ciliary circulation—a comparatively lower pressure system. In acute glaucoma, when IOP exceeds pressure of the CRA, it compresses circulation of the retina resulting in immediate death of neuronal tissue and optic atrophy. Furthermore, the optic atrophy resulting from acute glaucoma is a flat disc (non-excavated) as there is no sinking of the LC and thus no severance of nerve fibers taking place.

In contrast, chronic glaucoma develops when IOP becomes higher than the ciliary pressure supplying the BT. Since the ciliary pressure is a lower pressure system, even a moderate elevation of IOP can become higher than ciliary pressure and chronically compress circulation of the BT, resulting in its atrophy. As the BT becomes atrophied, the LC begins sinking and the nerve fibers get stretched and severed at the scleral edge. In summary,

![Figure 2. Relationship between ciliary pressure and IOP. Normally, ciliary circulatory pressure supplying the border tissue should be higher than IOP for healthy perfusion as in column (1). In column (2), the IOP is increased to 30 whereas the ciliary pressure remains the same at 25, this will result in high-tension glaucoma. In column (3) the ciliary pressure is decreased to 15 mm but the IOP is same, normal at 20, resulting in normal-tension glaucoma.](image)
the nerve fibers are being atrophied in a non-orderly fashion in acute glaucoma whereas, the nerve fibers are being severed in an orderly fashion in chronic glaucoma.

Due to atrophy of the BT, the LC starts sinking resulting in stretching and ultimately breakage of the prelaminar nerve fibers since one end is attached to the soma of the RGC and the other end anchored in the pores of the LC. Only the prelaminar nerve fibers can be destroyed in an orderly tandem fashion since they are still loose and have not yet fastened in bundles in the pores of the LC. Once the nerve fibers are anchored in the pores of the LC, they can’t be separated individually and thus their orderly tandem severance is not possible. Therefore, the LC may not be the site of injury in glaucoma as commonly believed.

3. Why Are the Nerve Fibers Being Destroyed in an Orderly Fashion?

The sinking of the LC and severance of the nerve fibers can explain their orderly destruction in glaucoma. As the LC sinks, the peripheral nerve fibers closest to the scleral edge are stretched and broken first, Figure 3. As a result, the next central fiber will move towards the periphery to occupy the space vacated by the preceding severed fiber and thus also get stretched and severed at the scleral edge.

In addition to the border tissue, the 360 degrees of nerve fibers also anchor the LC as roots anchor a tree. Thus, the severance of nerve fibers leads to further disc sinking. The cascade of severance of the nerve fibers and sinking disc would become self-propagated and will continue until all the nerve fibers have moved in an orderly tandem fashion to the scleral edge and get severed. This may explain the unstoppable nature of glaucoma despite maximum lowering of IOP. The severed segments undergo phagocytosis and thus will create empty spaces or excavation that we may be interpreting as cupping of the disc.

4. Do We Have Evidence of Severance of Nerve Fibers?

Progressive thinning of the RNFL in glaucoma can only be explained due to severance of nerve fibers as it is not occurring in non-glaucomatous optic atrophies. The arcuate retinal empty spaces in glaucoma are due to severance and depletion of arcuate nerve fibers and notching due to their depletion at the site of their entry in the disc. All of the 360 degrees of nerve fibers are being severed simultaneously, however the arcuate fibers being fewer in number are depleted earlier, thus producing the arcuate field defects, Figure 4 and Figure 5. Notching at the poles of the disc is the initial excavation in the disc and a confirmatory sign of glaucoma. At this stage, the pathognomonic arcuate field defects will appear on perimetry.

The histology of the end-stage glaucomatous disc is not 100% cupped LC, but an empty crater left over after the phagocytosis of severed nerve fibers, Figure 6. Splinter hemorrhages and characteristic whitish pallor of the disc are due to severance of the vasculature which is also meeting the fate of nerve fibers. In contrast, the histology of the non-glaucomatous optic atrophy such as due to multiple sclerosis, reveals the presence of nerve fibers though atrophied and shrunken, Figure 7.
Figure 4. Due to temporal sinking all the temporal fibers (Macular, superior and inferior arcuate) are being axotomized. However, the arcuate fibers being fewer in number will be depleted earlier resulting in arcuate field defects as in Figure 5.

Figure 5. Right eye: Double arcuate/ring scotoma after arcuate fibers have been severed. Arcuate fibers being fewer in number compared to macular fibers, will be depleted earlier resulting in arcuate field defects.
5. Conclusion

The severance of the nerve fibers appears to be the missing piece in glaucoma mystery. The sinking disc and severance of nerve fibers is able to corroborate with their orderly destruction, a hallmark of glaucoma. In essence, the nerve fibers along with vasculature are being severed in glaucoma. Therefore, glaucoma may not be
an optic disc neuropathy but an optic disc axotomy.

References


