Molecular basis of glaucoma and its therapeutical analysis in Pakistan: an overview

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Abstract

The human eye is an organ of vision. It plays a prime role in life, gives us the sense of sight, and enables to understand about the world around us. Visualization and interpretation of colors, shapes and dimensions of numerous objects is made possible by eye. Inherited eye diseases comprise 1/3 of all reported human genetic disorders. This review will focus on Glaucoma which comprises a predictable visual illnesses concerning optic nerve deterioration and if remains without any cure can result in failure in eyesight. The optical nerve injury comprises deterioration of the retinal ganglion cells (RGCs). Glaucoma represents a heterogeneous group of optic neuropathies with a complex genetic basis. These neuropathies gradually reduce vision without warning and often without symptoms. Different forms of glaucoma share some common clinical manifestations that usually include specific abnormal appearance of the optic nerve head, characteristic loss of visual field and chronic painless progression. Glaucoma is a progressive optical neuropathy considered by optical disc changes, nerve fiber film break, and visual field defects. Present-day treatment preferences predominantly targeting at reducing IOP by making use of pharmaceutical means, laser treatment and surgical procedure. Developed conducts target neuroprotection with vaccines, the hang-up of NO synthesis and apoptosis. Attaining a better appreciative of the pathogenesis can support in the improvement of novel handling options and, perhaps, even a remedy for glaucoma. There are more than 1.8 million glaucoma patients in Pakistan and almost half of them have already lost their eyesight, permanently, due to delay in diagnosis and treatment. About 90% population in the country has no awareness about this disease, resultantly; more and more people are becoming permanently blind in Pakistan due to untreated glaucoma.
Keywords

Eye, glaucoma, diagnosis, treatment, Pakistan

Introduction

The term “glaucoma” covers a quantity of diverse eye disorders, all of which encompass impairment to the optic nerve (Janssen et al., 2013). One common reason is that there is too much pressure inside the eye. The situation tends to be hereditary and may not display up until later in life. The improved pressure, called intra-ocular pressure, can harm the optic nerve, which communicates images to the brain. Intra-ocular pressure is produced by a fluid termed as aqueous humor formed by the eye themselves in the compartments of the eye in the middle of the cornea and the lens (Fatt and Weissman, 2013). Generally, this liquid, called aqueous humor, runs out of the eye through a mesh-like network. If this network becomes jammed, fluid forms up, triggering glaucoma. Glaucoma typically occurs when intra-ocular pressure upsurges. This take place when the fluid stress in the eye's anterior cavity, the area in the middle of the cornea and the iris, rises.

If the aqueous humor is prevented from draining appropriately, it twitches to accumulates pressure inside the eye builds up (Gupta et al., 2014). This presses alongside the optic nerve and there is a threat that nerve cells perish. Whether the amplified intra-ocular pressure does cause loss depends on, amid other things, how well the optic nerve can struggle this pressure. Glaucoma will cause loss of vision. Readings between 10 and 21 mm Hg are considered normal. Someone who has glaucoma does not always have above-average intra-ocular pressure.

Glaucoma sufferings roughly 70 million people around the globe, of whom around 10 % are supposed to be bilaterally unsighted (Lawrence, 2014). Estimation that is put forward in 2010 nearly 60.5 million individuals were affected by glaucoma and around 8.4 million were visionless from the sickness and it is expected that by the year 2020, this amount would rise to nearby 79.6 million. Statistics gather round by the World Health Organization (WHO) indicated that glaucoma is the second foremost reason of blindness worldwide, after cataract (Bourne et al., 2016). It is expected that glaucoma affects 12 million individuals accounting for 12.8% of the republics blindness and by 2020; this is estimated to be 16 million. Glaucoma is accountable for 10% of blindness worldwide. Glaucoma denotes to a group of disorders categorized by distinctive variations to the retinal nerve fiber layer and optical nerve head ensuing in compact optical field compassion. Its massive social and cost-effective impact
can be cherished by the fact that it leftovers a principal cause of sightlessness around the globe (Quigley and Broman, 2006). Glaucoma is a progressive optical neuropathy considered by optical disc changes, nerve fiber film break, and visual field defects (Pascolini et al., 2009; Resnikoff et al., 2004). This situation is known as glaucomatous optic neuropathy (GON) causing failure of visual field and ultimately to a state of irretrievable blindness (Alguire, 1990; Quigley, 1996). Glaucoma is the second principal reason of permanent blindness worldwide, thinking to influence 60 million inhabitants (Kelliher et al., 2006; Quigley and Broman, 2006). A convincing 70 million citizens suffer from glaucoma, and Asian ethnic minorities are extra susceptible to this visual ailment (Zhong et al., 2012). Linkage examinations have recognized 23 loci (the GLC1A, GLC1L, the GLC3A-GLC3B, 2p14, 2q33-q34, 5q22.1-q32, 10p12-p13, 14q11, 14q21-q22, 17p13, 17q25, and 19q12-q14) (Gemenetzi et al., 2012; Stone et al., 1997) for dissimilar kinds of glaucoma. Conversely, merely 4 genes (MYOC/TIGR, CYP1B1, optineurin [OPTN], and WD recap domain 36 (WDR36) (Monemi et al., 2005) have been recognized so far.

Currently, the only confirmed treatment for glaucoma is to reduce intra-ocular stress (IOP) with the aim of preventing supplementary glaucomatous optic nerve smash up (Heijl et al., 2002). Whilst many patients can be controlled with medications, patient devotion and optical toxicity are key issues in the residential globe, and lifetime outlay and ease of access to medications are issues in growing regions (Lemij et al., 2015).

Glaucoma is a composite peculiarity, in utmost condition have not pursue a straightforward heritage shape, and has inconsistent penetrance, subtle advancement, and habitually a later on commencement. Linkage examination have recognized 23 loci (the GLC1A, GLC1L, the GLC3A-GLC3B, 2p14, 2q33-q34, 5q22.1-q32, 10p12-p13, 14q11, 14q21-q22, 17p13, 17q25, and 19q12-q14) (Gemenetzi et al., 2012; Stone et al., 1997) for dissimilar kinds of glaucoma. Conversely, merely 4 genes (MYOC/TIGR, CYP1B1, optineurin [OPTN], and WD recap domain 36 (WDR36) (Monemi et al., 2005) have been recognized so far.

**Classification of Glaucoma**

Glaucoma is categorized conferring for the cause (primary VS secondary), composition of the frontal compartment (open angle VS closed angle) and stage of beginning (juvenile VS adult) (Sarfarazi, 1997). Glaucoma can categorized as primary once it happen with not any identified etiology or secondary wherever an earlier damage or illness is contributing. In broad-spectrum, glaucoma can be classified into 3 main kinds:
Primary open-angle glaucoma (POAG)

“Open-angle” means that the angle where the iris meets the cornea is as wide and open as it should be. It is also called chronic glaucoma and is the most common form, accounting for at least 90% of all glaucoma cases. The most general type of glaucoma is primary open angle glaucoma (POAG), disturbing over 33 million persons (Quigley, 1996). It is produced by the slow impediment of the drainage channels, causing in improved eye pressure. Primary open-angle glaucoma (POAG), in which the iridocorneal angle and frontal eye organizations emerge ordinary in gonioscopy inspection. It is the supreme ordinary form analyzed in all populaces investigated and is particularly predominant (~4.2%) in individuals with African lineage. Genetic line of attack make to know that primary open angle glaucoma may be innate either as a general, multifaceted trait with fully developed onset or, fewer commonly, as a conventional Mendelian or monogenic sickness that inclines to have an premature onset (Wiggs, 2007). Hereditary link studies of complex relations, frequently of European lineage, have acknowledged at smallest amount twenty one loci (GLC) for Mendelian arrangements of POAG (Fan and Wiggs, 2010).

Juvenile onset of open angle glaucoma (JOAG) is an alternative of primary open angle glaucoma (POAG). JOAG is diagnosed prior before the age of 40 and universally described as an artifact of autosomal dominant inheritance (Alward et al., 1998). The manner of inheritance in POAG is inconclusive (Nemesure et al., 2001). Furthermore, greater than 90 point alterations have been recognized universally in numerous racial groups. These kinds of mutations report for three to five percent of POAG cases and a higher percentage of Juvenile-onset of open angle glaucoma cases (just about six to thirty six %) (Challa, 2008; Gemenetzi et al., 2012). Primary open angle glaucoma (44.7 million cases wide-reaching) is measured more communal than primary angle closure glaucoma (15.7 million).

Primary Congenital Glaucoma (PCG)

Primary congenital glaucoma (PCG) is the supreme recurrent babyhood glaucoma and can lead to sightlessness during neonatal or early infantile period. Primary congenital glaucoma is the common term used for a glaucoma diagnosed in infancy or early childhood and is caused by abnormal intra-ocular fluid drainage from the eye as a result of a blocked or defective trabecular meshwork (the mesh- like drainage canals in the eye) (Ghate and Wang, 2015). It may also be due to a hereditary defect or abnormal development during pregnancy. In other cases, an abnormal drainage system may be the result of some other disease in the eye which results in secondary glaucoma. In these cases, the glaucoma may be associated with recognizable iris (the colored part of the eye), corneal, or other eye problems. Molecular genetic studies conducted during the last numerous years have established that PCG is an autosomal recessive trait. At present, four chromosomal loci have been concerned in PCG on GLC3A harboring the cytochrome P4501B1 (CYP1B1).
gene (Stoilov et al., 1997). Congenital glaucoma is a type of blinding eye illness that ruthlessly affects the improvement of visual acuity amongst children, infants and teenagers. It is caused by the hindrance of the aqueous outflow by the mal-development of the frontal chamber angle and the trabecular meshwork of the eye for the period of the embryonic development phase (Maul et al., 1980). PC Glaucoma is autosomal recessive eye disarray observed in intermittent and familial cases. The Primary Congenital Glaucoma is universal in North Africa and at hand in a more harsh form than in the western world (Helmy, 2016). The pervasiveness of pediatric glaucoma in the Middle East is 1:2500, while in consanguineous Slovakian children, it is 1 out of 1250. The incidence ranges between 1 out of 10,000 to 1 out of 12,500 in western countries (Helmy, 2016).

The popularity of Primary congenital glaucoma differs permitting to the topographical position and civilization. The occurrence is 1/10,000 in Western motherlands, 1 out of 2,500 in Arabic people, and uppermost in the Gypsy inhabitants of Slovakia, wherever its prevalence is 1 in every 1,250 living births outstanding to an elevation speed of cousin marriages in these republics (Bejjani et al., 2000; Sarfarazi et al., 2003). The predominance of Primary congenital glaucoma in south Indian is estimated to be 1 in 3,300 and result 4.2 percent of infancy loss of sight (Dandona et al., 1998). 3 loci have been recognized: the GLC3A (2p21) (Sarfarazi et al., 1995), the GLC3B (1p36) (Akarsu et al., 1996) and GLC3C (14q24.3) (Stoilov et al., 2002).

**Primary angle Closure Glaucoma (PACG)**

It is caused by clogged drainage channels, causing in a sudden increase in intraocular pressure 12-21. It is also called acute glaucoma or narrow-angle glaucoma. Unlike open-angle glaucoma, angle-closure glaucoma is a result of the angle between the iris and cornea closing. It is relatively a less common form of glaucoma. Primary angle closure glaucoma (PACG) is the supreme widespread type of glaucoma universally (Kuehn et al., 2011). Nevertheless, primary angle closure glaucoma (PACG) is thought to be the most familiar reason of bilateral glaucoma sightlessness cosmopolitantly (Quigley et al., 2001). This is a sub kind of glaucoma, and can be defined as an anatomic disarray of the frontal cavity, through which the drainage position is obstructed by the frontal development of the iris (Lin et al., 1997). Primary angle closure glaucoma leftovers a main reason of unalterable blindness, predominantly in Asian countries such as China (Foster et al., 2000), Mongolia (Foster et al., 1996), Singapore (Foster et al., 2000), and India (Dandona et al., 2000) with up to 80% of the predictable 15 million citizens afflicted with PACG inhabitant in Asia (Quigley and Broman, 2006). The figure of patients with PACG is probable to rise by roughly 5 million people from 16 million over the next decade (Quigley and Broman, 2006). Quite a lot of anatomic hazard issues for the improvement of Primary angle closure glaucoma has been famous together with a narrow frontal chamber deepness and petite axial distance. Pretended replica organizations have providing the facts that the occurrence of equally amplified lens curving and a undersized zonule-iris space add to a pupil block in angle closure glaucoma (Huang and Barocas, 2004).
PACG patients have been found to have exacting anatomic biometric skin tone together with shallow anterior chambers (Lin et al., 1997), lens breadth and position (Foster et al., 1996), constricted iridio-trabecular drainage angles, dumpy axial lengths (Abu-Amero et al., 2007), and hyperopic refractive fault (Congdon et al., 1996). The infection is accompanying with grownup stage, womanly masculinity and rivalry (anterior compartment angle is thinner in Eskimos and Asians (Salmon, 1999).

Numerous studies have exposed that genetic factors take part in an chief role in the development of PACG (Alsbirk, 1975; Amerasinghe et al., 2011). Although a sum of susceptible loci and genetic factor have been investigated for PACG, the accurate genes underlying PACG have not been recognized (Cong et al., 2009).

**Most frequently glaucoma genes reported in Pakistan**

Novel CYP1B1 mutations in consanguineous Pakistani families with primary congenital glaucoma identified an innovative cyp1b1 mutation in consanguineous Pakistani families with primary congenital glaucoma. The disease-causing alterations could be identified in ~31% of PCG affected families from Pakistani population (Firasat et al., 2008). This frequency estimation of CYP1B1 mutations in our population may be imprecise because of a small number of PCG families studied. Nonetheless, it connotes the prominence of CYP1B1 in PCG pathogenesis. Homozygosity mapping in consanguineous Pakistani family revealed one 11-Mb homozygous area incorporating the CYP1B1 gene (Micheal et al., 2015b). A homozygous CYP1B1 missense mutation (p. Arg390His) was recognized in this family. Sequence analysis of CYP1B1 in 39 supplementary families revealed one known and three fresh homozygous mutations in PCG (p.Ala288Pro, p.Asp242Ala, p.Arg355* and p.Arg290Profs*37). In POAG, one novel heterozygous missense mutation (p.Asp316Val) was recognized in one family and a beforehand reported mutation (p.Glu229Lys) was identified in three families. CYP1B1 mutations are the prime cause of primary congenital glaucoma in Pakistani patients (Rauf et al., 2016). Also recounted a family from Pakistan in which participants have inherited JOAG and PCG due to a known homozygous mutation in CYP1B1 (Bashir et al., 2015).

LTBP2 is the succeeding gene concerned in PCG to date. identification of homozygous null mutations in LTBP2 as a basis of PCG in human patients (Khan, 2011). These results might have insinuations for the clinical supervision of infancy sightlessness and give new intuitions into the development of the anterior organizations of the eye, implying that LTBP2 may have an critical structural role in keeping the shape of the ciliary body and its neighboring structures.

Another homozygous null mutation in LTBP2 as a trigger of PCG reported in Pakistani as well as Gypsy patients (Ali et al., 2009). Micheal et al.,
revealed that variants in ASB10 were found to be significantly associated with sporadic POAG in the Pakistani population (Micheal et al., 2015a).

First described MYOC correlated glaucoma from Pakistan isolating as autosomal dominant peculiarity in large family detected with JOAG (Waryah et al., 2013). Identification of innovative disease producing allele in MYOC suggests hereditary heterogeneity of the populace. Association of novel disarray allele indicates genetic heterogeneity of the populace. Linkage scrutiny revealed an autosomal dominant allele at GLC1A1 locus, co-segregating with disorder phenotype in a huge consanguineous household, enrolled from inner of Sindh area. Succeeding sequencing of MYOC gene in all patients, recognized a heterozygous c.1130 CNG variation in exon 3, substituting Threonine to Arginine at codon 377 of Myocilin. Preponderance of MYOC mutations (97%) has been discovered in exon 3 of the gene, which code a functionally essential Olfactomedin domain. The data indicate that Threonine residue at 377 codon of MYOC is functionally essential and its changeover with Arginine bases hard-hitting form of glaucoma.

**Present-Day Treatment Preferences in Pakistan**

Glaucoma is a lineage of chronic disorders due to the worsening of the optic nerve; its onset is asymptomatic in early phases and leads to sightlessness if left untouched. It is the next supreme usual cause of blindness globally with an predictable worldwide occurrence of about 67 million (Consoli and Ramlogan, 2015). The causes behindhand glaucoma and the manner in which it spearheads to loss of eyesight have not been clearly acknowledged; glaucomatous loss can be decelerated down but not reversed, and no particular precautionary measures exist. To date the maximum important regions of enhancements distresses the cataloguing of different forms have the disorder as well as cleansing clinically and pharmacological involvements for very specific conclusions. Glaucoma creates a thought-provoking case analysis in that the obstinacy of a specific framing which is now solidly considered by the ophthalmology medical society as only somewhat correct at source of much misdiagnosis and fairly unsuccessful clinical reaction.

The objective of remedying glaucoma lies predominantly on checking or postponing the loss of optical field (Brubaker, 2003). Since neuronal cell fatality is irretrievable, no treatment exists once the visual field is disappeared. Though, since IOP is the principal risk source causing the loss of RGCs, the approaches of handling mostly involve lessening IOP (Brubaker, 2003). Other essential factors such as cost, suitability and well-being should also be measured (Fechtner and Realini, 2004). Undercurrent treatments for glaucoma involve medication, laser usage and operation.
With early analysis and appropriate medication and cure, glaucoma can be controlled. Conversely, sight loss causing from glaucoma cannot be renovated. At the present-day, there is no remedy. Once perceived, glaucoma usually needs continuing, long-standing care.

But beforehand the cure recognizing is important. Diagnosing glaucoma is not continually stress-free. The most principal anxiety is defending eyesight. Physician aspects at many aspects before making decisions about treatment.

**Glaucoma Diagnosing**

**Tonometry**

In tonometry, eye drops are used to emotionless the eye. Afterward a physician or technician uses a stratagem called a tonometer to measure the inner stress of the eye. A small aggregate of pressure is pertained to the eye by a minute device. The normal range for eye pressure is 12–22 mm Hg. The level of eye pressure by which glaucoma progresses is not the same for everybody and some individuals can get glaucoma even if their pressures are in the normal range.

**Ophthalmoscopy**

This investigative route aids the doctor scrutinize optic nerve for glaucoma impairment. Eye drops are used to expand the pupil so that the physician sees throughout eye with a particular lens in direction to observe the form and color of the optic nerve.

**Perimetry**

Perimetry (or a ophthalmic field assessment) creates a map of field of visualization. This analysis helps a specialist to regulate whether a individual vision has been disturbed by glaucoma.

**Gonioscopy**

This is a investigative exam that supports to regulate whether the position wherever the iris gathers the cornea is open, thin, or closed. Through the examination, eye droplets are used to distress the eye and a distinct hand-held contact lens is smoothly positioned on the eye for a few minutes.

**Pachymetry**

It measures the wideness of the cornea, the well-defined opening at the anterior of the eye. Corneal thickness has likely to effect eye pressure interpretations. Uncertainty a cornea is thicker than normal, pressure evaluations with a
This tonometer may be sophisticated. This contributes your eye physician surplus evidence for glaucoma analysis.

**Medications**

Glaucoma is characteristically cured with the use of medications that either helps the unsolidified drain healthier or lessening the quantity of fluid built by the eye. In Most incidents, prescription can securely control eye pressure for several years.

Medications encompass hindering the incursion of aqueous humor, improving the outlay of aqueous humor, shielding the optical nerves (Woodward and Gil, 2004) and biasing the osmotic pressure concerning plasma and the eyes (Kwon et al., 2009). $\alpha_2$ adrenoreceptor agonists and $\beta_1$ receptor antagonists depress IOP by impeding the inflow of aqueous humor to the eye. Timolol, which is the supreme recommended drug, and betaxolol, which has the littlest universal side effects, are together $\beta_1$ receptor blockers (Woodward and Gil, 2004). A third kind of drug that restrains the inflow of humor is carbonic anhydrase inhibitors, such as acetazolamide and dorzolamide. Such medications are regularly prepared jointly as in Cosopt (dorzolamide hydrochloride and Timolol maleate) (Fechtner and Realini, 2004). Alternative approach of decreasing IOP is by boosting the outflow of humor from the eyes over the use of muscarinic acetylcholine receptor agonists (Schwartz and Budenz, 2004; Woodward and Gil, 2004). This method is unintended, but encompasses a muscarinic acetylcholine receptor (M3)-mediated narrowing of the ciliary muscle (Woodward and Gil, 2004). The shrinkage triggers the broadening of the gaps in the trabecular meshwork. The latest class of drugs using this approach is the prostaglandin F2$\alpha$ byproducts which boost the uveoscleral outflow (Khaw et al., 2004). Bimatoprost falls underneath this classification and is considered the furthermost successful anti-glaucoma drug (Woodward and Gil, 2004).

**Laser and Operation**

A subordinate choice for cure of glaucoma is the use of laser treatment. The principal scheme encompasses “burning” holes in several spaces inside the eyes incorporating the ciliary and the pigmented trabecular meshwork cells (Schwartz and Budenz, 2004). The profits contain being noninvasive, demanding fewer patient observance and decreasing the likelihood of infectivity or hemorrhage. The IOP of most patients can cutback approximately 20-30%, but the treatment result wears off 5-10% each year. In addition with timolol, the two year IOP decreasing attainment ratio is 70%, likened with the laser isolated (44%) and timolol alone (30%) (Schwartz and Budenz, 2004).
A everyday practice of operation is trabeculectomy, which generates a secured channel permitting aqueous humor to run from the fore chamber inside the eye to sub-Tenon’s and sub conjunctival area (Khaw et al., 2004; Schwartz and Budenz, 2004). The benefits of operation involve alleviating IOP and avoiding the necessities for exacting patient passivity and constant drug expenditures (Schwartz and Budenz, 2004). Surgery is well-thought-out as the last remedy as failure of operation can outcome in instantaneous blindness due to impediments such as choroidal outflow, hypotonic maculopathy, suprachoroidal bleeding and optical nerve dowsing (Schwartz and Budenz, 2004).

**Alternative Surgical Treatments**

Looking for to moderate difficulties accompanying with established glaucoma operation, alternate surgical possibilities have been established.

- The Ex-Press tiny glaucoma shunt is used with established trabeculectomy methods to systematize the procedure and conceivably decrease the possibilities of the eye pressure becoming excessively low in the immediate post-operative stage, which is infrequently a problematic with established methodologies.

- Canaloplasty is a technique that implicates magnifying the surviving fluid outflows passageway (the Schlemm’s canal) in supplement to generating a novel fluid outflow alleyway within the eye wall.

**Gentler sorts of laser Cyclophotocoagulation**

These innovative methodologies to glaucoma operation show potential for improved protection. As with all novel practices, time and continuation conclusions are compulsory to see which medical measures will persist advantageous for relieving glaucoma patients durable.

**Forthcoming Treatment Possibilities for treatment of glaucoma in Pakistan**

Based on latest information gotten from research on the pathology of neuronal apoptosis, there are numerous practices of new treatments that can be helpful for diagnosis of glaucoma in Pakistani population. Some of them are discussed here.

**Minimally Invasive Glaucoma Surgery (MIGS)**

In this technique, contrasting conventional glaucoma operation, there is minimum maneuvering of the sclera and the conjunctiva. Whilst these practices
lessen the frequency of hitches, some quantity of efficacy is also transacted for the better protection.

- Miniaturized forms of trabeculectomy; operating tiny, microscopic sized ducts that can be injected into the eye and trench fluid from the internal of the eye to beneath the conjunctiva, these innovative procedures are aimed to create the trabeculectomy process securer.

- Complete internal or suprachoroidal shunts; by Using teeny tubes with very slight internal openings, the front of the eye is connected to the suprachoroidal space between the retina and the wall of the eye to intensify the drainage of fluid from the eye.

- Trabecular Operation; the trabecular meshwork can be detached or evaded.

**Neuro defensive Vaccines**

Subsequently resistance to elevated intra ocular Pressure is immune related, T-cell provoked neuro-protection may immunize the retinal ganglion cells from apoptosis. For instance, copolymer-1 could be used as a vaccine as it is an antigen that cross responds with a widespread array of T-cells, and can provoke a defensive immune reaction to shield retinal ganglion cells from cell mortality initiated by toxins or improved Intra ocular Pressure (Bakalash et al., 2003). A fundamental concern for planning neuro-defensive vaccines is the position of protection. Some recommended that the focus should be in the retinal ganglion cells and not in the optical nerves because in the initial glaucomatous phase cell fatality stimulated by higher intra ocular pressure that arises in the retinal ganglion cells, not in the optical nerve (Bakalash et al., 2003). R16, a peptide derivative from the retinal ganglion cell, is one paradigm of a neuro-protecting vaccine. Though R16 can instigate slight retinal ganglion cell death for those deprived of glaucoma, the advantage from this treatment extremely surpassed the mutilation from untreated glaucoma situations (Bakalash et al., 2003).

**NMDA Receptor Antagonists**

Memantine is a NMDA receptor opponent that chunks unnecessary NMDA receptor action. This medication has been accepted for medical use in Europe for the healing of Alzheimer’s disease and vascular dementia, although its efficiency in preventing glaucomatous retinal ganglion cell disintegration is still undetermined (Lipton, 2003).

**INOS-2 Inhibitors**

Subsequently the up regulation of iNOS-2 is injurious to neurons, its inhibition could have a neuro-defensive outcome. An inhibitor like amino guanidine can inhibit the mortality of around 75% of retinal ganglion cells during six months of
provoked higher intra ocular pressure and help to prevent additional damage of retinal ganglion cells (Neufeld, 2004).

**Nutritive Complements**

A thought-provoking alternate to amino guanidine is *Ginkgo biloba* extract (EGb 761). *Ginkgo biloba* is used as a nourishing complement for the decline of platelet accumulation, vasodilation and lessening of blood viscidness (Bartlett and Eperjesi, 2004). This comprises 24 percent flavonoid glycosides and 6 percent terpenoids, which might prevent toxicity and NO free radical accretion by impeding iNOS (Hirooka et al., 2004). In experimentations when intra ocular pressure is raised in rats by cautery of episcleral vessels, nurturing the rats 30 mg of EGb 761/day for about five months reduced retinal ganglion cell damage from 29.9% to 4.6% (Hirooka et al., 2004). Additional research scrutinizing the alteration of NO in reaction to EGb 761 treatment might deliver insight into its neuro-defensive method.

**Calcium Channel Blockers**

Utmost apoptotic means comprise increasing intracellular calcium stages. Flunarizine, a calcium network blocker, has been indicated to expressively increase retinal ganglion cells existence in rat and rabbit mockups by decreasing intra ocular pressure (Osborne et al., 2002). Nevertheless, the accurate method of this and further calcium channel blockers desires to be explained.

**STAT-3 Initiation**

Alternative objective is the signal transducers and activators of transcription protein 3 (STAT-3). They play an significant role in cell development and discrimination and are of significance due to the messenger RNA of this protein is upbeat regulated in rats with glaucoma (Thanos and Naskar, 2004). The initiation of STAT-3 passageway could impede apoptosis by suppressing caspase-3. One molecule that has been examined is ciliary neurotropic factor (CNTF), which is an interleukin-6 cytokine. The insertion of CNTF into rat eyes with enlarged intra ocular pressure decreases apoptosis, phosphorylates STAT-3, and downgrades the action of caspase-3 (Adamus et al., 2003). Interleukin-10 also has neuro protecting action through the STAT-3 passageway (Boyd et al., 2003).

**Caspase Inhibitors**

Inhibitors of apoptosis protein (IAP) can correspondingly decrease apoptosis by impeding Caspase. Baculoviral IAP repeat-containing protein-4 (BIRC-4) is a direct inhibitor of Caspase 3, 7 and 9. BIRC-4, transducted into the eye, can obstruct apoptosis of optical nerve axons (McKinnon et al., 2002). A pharmacological methodology for apoptosis inhibition is the usage of minocycline, which could inhibit caspase-3-induced apoptosis (Baptiste et al., 2003).
2004). It proliferate the persistence rate of retinal ganglion cells visible to the damaging results of glutamate. Moreover, it can act synergistically with MK-801, an opponent to NMDA receptors, to escalate retinal ganglion cells existence percentage.

**Heat Shock Proteins**

Geranylgeranylacetone (GGA) is an acyclic polyisoprenoid presently used in Japan as an anti-ulcer remedy. The neuro defensive effects of the medication are accelerated through the use of heat shock proteins. Specifically, Heat Shock Protein 72 looks to act as an opposing apoptotic chaperone protein that inhibits with various phases in the apoptotic passageway. Systematically, the drug is assumed to stimulate Heat Shock Factor 1 (HSF-1), a transcription element for heat shock protein, which oligomerizes in the cytosol and transfer into the nucleus once visible to stressors (Sohn et al., 2013).

**CONCLUSION**

There is no medication for glaucoma still, and vision loss is irreplaceable, consequently molecular diagnostics for predictive assessment and early interference is essential to decrease the influence of visual mutilation and eventually blindness. To accomplish this goal, the requirement is to exemplify all subtypes of glaucoma at molecular level and recognize loci/genes contributing to this ophthalmic disorder in diverse populaces.

**Abbreviations**

PCG: Primary Congenital glaucoma  
POAG: Primary Open Angle Glaucoma  
PACG: Primary angle Closure Glaucoma  
RGCs: Retinal ganglion cells  
IOP: Intra Ocular Pressure  
CYP1B: Cytochrome P450 1B1  
LTBP2: Latent Transforming Growth Factor-Beta-Binding Protein 2

**Author Contribution**

LK collected data and wrote the manuscript. MA, MQ participated in the design and editing of the manuscript. FJ, and BH edited the first draft.
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