Binocularly Asymmetric Crowding in Glaucoma and a Lack of Binocular Summation in Crowding

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Received: August 26, 2021
Accepted: January 5, 2022
Published: January 27, 2022

Citation: Shamsi F, Liu R, Kwon M. Binocularly asymmetric crowding in glaucoma and a lack of binocular summation in crowding. Invest Ophthalmol Vis Sci. 2022;63(1):36. https://doi.org/10.1167/iovs.63.1.36

PURPOSE. Glaucoma is associated with progressive loss of retinal ganglion cells. Here we investigated the impact of glaucomatous damage on monocular and binocular crowding in parafoveal vision. We also examined the binocular summation of crowding to see if crowding is alleviated under binocular viewing.

METHODS. The study design included 40 individuals with glaucoma and 24 age-similar normal cohorts. For each subject, the magnitude of crowding was determined by the extent of crowding zone. Crowding zone measurements were made binocularly in parafoveal vision (i.e., at 2° and 4° retinal eccentricities) visual field. For a subgroup of glaucoma subjects (n = 17), crowding zone was also measured monocularly for each eye.

RESULTS. Our results showed that, compared with normal cohorts, individuals with glaucoma exhibited significantly larger crowding—enlargement of crowding zone (an increase by 21%; P < 0.01). Moreover, we also observed a lack of binocular summation (i.e., a binocular ratio of 1): binocular crowding was determined by the better eye. Hence, our results did not provide evidence supporting binocular summation of crowding in glaucomatous vision.

CONCLUSIONS. Our findings show that crowding is exacerbated in parafoveal vision in glaucoma and binocularly asymmetric glaucoma seems to induce binocularly asymmetric crowding. Furthermore, the lack of binocular summation for crowding observed in glaucomatous vision combined with the lack of binocular summation reported in a previous study on normal healthy vision support the view that crowding may start in the early stages of visual processing, at least before the process of binocular integration takes place.

Keywords: crowding, glaucoma, binocular summation, ganglion cell damage, crowding zone

Glaucoma is a leading cause of irreversible blindness worldwide, characterized by the progressive loss of retinal ganglion cells (RGCs) and the resultant visual field defects.1 The conventional view has been that glaucoma spares central vision until the end stage and, thus, it has little impact on central visual function.2–5 However, accumulating evidence has shown that even early glaucomatous injury involves the macula, and this macular damage is more common than generally thought.6–15 For example, a number of anatomical studies8,11,12,14–16 using spectral-domain optical coherence tomography have shown that the thickness of the retinal nerve fiber layer and the RGC plus inner plexiform layer (RGC+) even in the macula, are significantly thinner in patients with early glaucoma than in healthy controls. In parallel with anatomical evidence, perceptual evidence indicates noticeable deficits in assumed-to-be central vision tasks, such as reading and object/face recognition.3,5,17–24 Given the conventional view, it is surprising that reading difficulty is a common complaint among patients with glaucoma.5,25–31

Although the exact perceptual mechanism limiting pattern recognition function in glaucoma remains unclear, converging evidence suggests that changes in visual crowding in glaucomatous vision may be one of the limiting factors.32–36 Visual crowding refers to the inability to recognize a target object in clutter37 on account of the deleterious influence of nearby items on visual recognition.38 In real life, objects rarely appear in isolation. Therefore, the ability to isolate the target item from nearby clutter plays a critical role in everyday visual function such as reading, face recognition, and visual search.39–43 Because the observers have no difficulty recognizing objects in the same retinal eccentricity in the absence of clutter, the phenomenon of crowding cannot be simply accounted for by decreased visual acuity or a loss of contrast sensitivity. Although the exact locus and mechanism of crowding remain under debate, a
Crowding grows with increasing retinal eccentricity as the receptive (perceptual) field size increases in the periphery (i.e., scale shift). Although very little crowding exists in normal central/parafoveal vision, some clinical conditions like amblyopia manifest noticeably increased foveal crowding, which correlates with reading rate. Hence, crowding, particularly in the central visual field, can be a good indicator of a person’s everyday visual function.

Thus, the current study was undertaken to examine whether crowding is indeed exacerbated in parafoveal vision of glaucomatous eyes (i.e., the central 8° visual field), the visual field relevant to daily visual function. The relationship between crowding and glaucomatous damage was investigated in two ways: a between-subjects study design comparing crowding between patients with glaucoma and age-similar normal controls and a within-subjects study design comparing crowding between the worse eyes and the better eyes of patients with glaucoma. We, thus, hypothesized that the parafoveal crowding would be significantly larger in the glaucomatous vision compared with the age-matched normal vision and in the worse eye compared with the better eye.

Furthermore, because glaucoma is often bilateral and asymmetry is common, this binocularly asymmetric in glaucomatous damage provides us with a unique opportunity to explore the binocular summation of crowding. Binocular summation refers to an increase in binocular performance over monocular performance that is often quantified as a ratio of visual performance or sensitivity of the binocular to that of the better eye. Previous studies on contrast sensitivity or visual acuity have reported binocular summation ratio of 1.4 (i.e., a 40% increase in binocular condition) or beyond in normal vision.

However, little is known about how the crowding effect is integrated between two eyes. Thus, the secondary aim of the current study was to explore the mechanism of binocular summation in crowding.

To this end, we assessed binocular crowding in both patients with glaucoma (n = 40) and in age-similar normal controls (n = 24). Crowding was assessed with a well-established method: the spatial extent of crowding (i.e., threshold spacing between the target and flankers required to yield a criterion recognition accuracy). Crowding measurements were made in parafoveal vision (i.e., retinal eccentricity of 2° or 4°). For the binocular summation analysis, both monocular and binocular crowding were measured in a subset of patients with glaucoma.

**METHODS**

A total of 68 subjects participated in the current study: 44 patients with bilateral primary open-angle glaucoma (mean age, 63.66 ± 8.91 years) and 24 age-similar normal control subjects (mean age, 60.21 ± 9.56 years). Patients with bilateral glaucoma and control subjects were recruited from the Callahan Eye Hospital Clinics at the University of Alabama at Birmingham. Patients with glaucoma whose diagnosis was confirmed through medical records met the following inclusion criteria in both eyes: (i) glaucoma-specific changes of the optic nerve or a nerve fiber layer defect. The presence of the glaucomatous optic nerve was defined by masked review of optic nerve head photos by glaucoma specialists using previously published criteria. (ii) Glaucoma-specific visual field defect: a value of Glaucoma Hemifield Test from the Humphrey Field Analyzer (HFA) must be outside normal limits. (iii) No history of other ocular or neurological disease or surgery that caused visual field loss.

Visual field tests were performed with standard automatic perimetry using SITA Standard 24-2 and 10-2 tests with an HFA (Carl Zeiss Meditec, Inc., Jena, Germany). Goldmann size III targets with a diameter of 0.43° were presented for 200 ms at one of 54 (68) test locations for 24-2 (10-2) in a grid on a white background (10 cd/m²).

The average mean deviation obtained from the HFA (24-2 test) in patients with glaucoma was −8.55 ± 9.75 dB for the right eye and −10.93 ± 8.43 dB for the left eye. According to the Hodapp–Anderson–Parrish glaucoma grading system, the majority of our patients with glaucoma were either in the early or moderate stages of glaucoma.

The mean binocular visual acuity (Early Treatment Diabetic Retinopathy charts) for patients with glaucoma was −0.01 ± 0.10 logMAR (or 20/20 Snellen equivalent). The mean monocular visual acuity was 0.09 ± 0.18 logMAR for the right eye and 0.10 ± 0.16 logMAR for the left eye. The mean binocular log contrast sensitivity (Pelli–Robson charts) was 1.69 ± 0.19.

Normal vision was defined as better than or equal to 0.2 logMAR best-corrected visual acuity in each eye with normal binocular vision (confirmed through preliminary vision tests including the Worth four dot test, Stereo Fly vision test, HFA, visual acuity, and contrast sensitivity) and with no history of ocular or neurological disease other than cataract surgery.

The mean binocular visual acuity for normal control subjects was −0.07 ± 0.08 logMAR (or 20/20 Snellen equivalent). The mean monocular visual acuity was −0.01 ± 0.08 logMAR for the right eye and −0.01 ± 0.1 logMAR for the left eye. The mean binocular log contrast sensitivity for normal controls was 1.92 ± 0.10.

All participants were native or fluent English speakers without known cognitive or neurological impairments, confirmed by the Mini Mental Status Exam (score of ≥ 25). The main experiments were conducted with binocular viewing (n = 40). This was done to assess the amount of crowding relevant to real-life visual tasks. Proper refractive correction for the viewing distance was used. For a subset of patients with glaucoma (n = 17), the experiments were also performed under a monocular viewing condition. The number of subjects with both binocular and monocular crowding measurements was 13. For the subjects with both binocular and monocular measurements, the binocular and monocular experiments were performed on different days and monocular measurements were done after the binocular measurement. The experimental protocols followed the tenets of the Declaration of Helsinki and were approved by the Internal Review Board at the University of Alabama at Birmingham. Written informed consents were obtained from all participants before the experiment after explanation of the nature of the study.

**Stimulus and Apparatus**

For the crowding task, the stimuli consisted of a target letter flanked by four tumbling Es on four cardinal sides of the target. The target letter was randomly drawn from a set of
The luminance of the display monitor was measured linear using an 8-bit look-up table in conjunction with photometric readings from a MINOLTA LS-110 Luminance Meter (Konica Minolta Inc., Tokyo, Japan).

10 Sloan letters: CDHKNORSVZ. All the letters were black on a uniform gray background (159 cd/m²) with a contrast of 99%, and a letter size of 0.8° (x-height). The fixation dot used in this experiment was a black circle in the center of the screen spanning 0.25° of the visual field. All stimuli were generated and controlled using MATLAB (version 8.3) and Psychophysics Toolbox extensions (version 3) for Windows 7, running on a PC desktop computer (model: Dell Precision Tower 5810). Stimuli were presented on a liquid crystal display monitor (model: Asus VG278HE; refresh rate: 144 Hz; resolution: 1920 × 1080, graphic card: 2 GB Nvidia Quadro K2000, subtending 60° × 34° visual angle at a viewing distance of 57 cm) with the mean luminance of the monitor at 159 cd/m². The luminance of the display monitor was made linear using an 8-bit look-up table in conjunction with photometric readings from a MINOLTA LS-110 Luminance Meter (Konica Minolta Inc., Tokyo, Japan).

Measuring the Spatial Extent of Crowding (Crowding Zone)

The crowding effect was measured by determining the spatial extent of crowding. Threshold spacing (or crowding zone) was defined as the center-to-center distance between the high-contrast target and flankers that yields a target-identification accuracy of 79.4%. Thus, threshold spacing becomes larger with increasing crowding. A subject’s threshold spacing was estimated at eight retinal locations: two retinal eccentricities (2° or 4°) and four azimuth angles around the fixation (45°, 135°, 225°, or 315°) (Fig. 1), using a three-down-one-up staircase procedure66 with a step size of 15%. The total number of staircase reversals were set to nine and the threshold spacing was determined by taking the geometric average of the last seven staircase reversals. For each retinal location, a target letter was flanked by four tumbling Es. At each block, one of the eight target locations was tested. The tested location was predetermined for each block and counterbalanced across blocks to minimize the order effect. At the beginning of each block, a small red dot was shown at the target location. Then, the subject pressed a key on the keyboard to initiate the experiment block. In each trial, a target letter with flankers was presented at the assigned location for 150 ms (approximately 22 frames = 152.7 ms). During the last trial of each block, the target letter was presented at the same location without flankers (i.e., uncrowded condition). Participants were instructed to fixate on a central dot during the stimulus presentation and to report the target letter they saw during the stimulus interval in a subsequent response interval (as shown in Fig. 1A) by clicking the mouse on the selected letter. No time limit was considered for the subjects’ responses and they responded at their own pace. Auditory feedback was given whenever the correct answer was chosen. The spacing between the letters in the response panel was fixed and the letters in the response panel were uniformly spaced in a circle with the radius of 5° eccentricity. It should be noted that the subjects did not have to maintain their central fixation during the response interval. The time interval between the offset of the stimulus and the onset of the response panel was set to 500 ms. The task procedure was the same for monocular and binocular measurements.

A chinrest was used to minimize head movements. The experimenter visually observed subjects to confirm that fixation instructions were followed. Note that the stimulus duration (150 ms) was too short to allow for any reliable saccadic eye movements (considering the fact that the average saccades usually take about 230 ms and up to 250 ms). Therefore, the subjects knew there was no advantages in moving their eye during the stimulus duration. All subjects had practice trials for both crowding and visual span tasks before data collection. A subject’s stable fixation was also monitored using a high-speed eye tracker (EyeLink 1000 Plus/Desktop mount, SR Research Ltd., Kanata, Ontario, Canada).
Data Analysis

The normality of the data was checked using the quantile-quantile plot. To examine if there are any significant differences in crowding effect between (i) retinal eccentricities and (ii) patients with glaucoma and age-matched normal cohorts, we performed an ANOVA on crowding zone – 2 (retinal eccentricity: 2° and 4°) × 2 (subject group: glaucoma and normal cohorts) repeated measures ANOVA with retinal eccentricity as a within-subject factor. To determine which specific groups differ from each other, we also performed Tukey’s HSD post hoc test. Statistical analyses were performed using MATLAB software (version (R2020b; The MathWorks Inc., Natick, MA). The ratio between the crowding zone of the individual glaucoma subjects with respect to the average crowding of the age-matched normal control group as well as the ratio of the average of the two groups were calculated at two eccentricities (2° and 4°). To compare the binocular with monocular crowding, we categorized the data into worse and better eyes based on the crowding values (i.e., more and less crowded eyes) and calculated the average crowding for the better and worse eyes and compared them with the average binocular crowding. The binocular summation ratio of crowding is defined as the ratio of binocular crowding to the crowding of the better eye. To follow the notion that the summation ratio as the ratio of binocular crowding to the crowding of the better eye crowding. To determine the crowding effect, we capitalized on the fact that the severity of glaucomatous damage tends to differ between the two eyes. Monocular crowding zone was measured monocularly at 2° and 4° eccentricities for a group of glaucoma subjects (n = 17) using the same experimental paradigm as the binocular crowding zone measurement. Crowding zone of the worse eye (i.e., the eye with more severe glaucomatous damage) was compared with that of the better eye. The worse and better eyes were determined based on the mean deviation value from the HFA 10-2 test (i.e., perimetry in the central 10° visual field), where more negative mean deviation is considered to be more severe glaucoma.

Increased Monocular Crowding in the Eyes With More Severe Glaucoma

To further confirm the effects of the glaucoma on crowding, we capitalized on the fact that the severity of glaucomatous damage tends to differ between the two eyes. Monocular crowding zone was measured monocularly at 2° and 4° eccentricities for a group of glaucoma subjects (n = 17) using the same experimental paradigm as the binocular crowding zone measurement. Crowding zone of the worse eye (i.e., the eye with more severe glaucomatous damage) was compared with that of the better eye. The worse and better eyes were determined based on the mean deviation value from the HFA 10-2 test (i.e., perimetry in the central 10° visual field), where more negative mean deviation is considered to be more severe glaucoma.

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No Evidence of Binocular Summation or Inhibition in Crowding

In the foregoing session, we showed that the binocularly asymmetric glaucomatous damage brings about corresponding binocularly asymmetric crowding. This asymmetry in the monocular crowding provided us with a unique opportunity to explore the binocular combination of crowding.
Binocularly Asymmetric Crowding in Glaucoma

A) Exemplary subjects

FIGURE 2. Binocular crowding in glaucoma vs age-similar normal controls. (A) Exemplary subjects. The extent of crowding zone at each testing location was plotted in polar coordinates for 10 individuals with glaucoma (orange) and the average of normal subjects (green).

(B) Group average results. (Left) The crowding zone averaged across testing locations ($\Theta = 45^\circ, 135^\circ, 225^\circ$, or $315^\circ$) are plotted as a function of eccentricity ($2^\circ$ and $4^\circ$): patients with glaucoma (orange dots) and normal cohorts (green dots). Each dot represents the data point from an individual subject. (Right) Mean ratio of the crowding zone of patients with glaucoma to that of normal cohorts (ratio = crowding zone$_{glaucoma}$/crowding zone$_{normal control}$) was plotted as a function of eccentricity. Individual data points are the ratio of the crowding for individual patients with glaucoma to the mean crowding of the normal group at two eccentricities. Note that ** indicates statistical significance at a $P$ value of less than 0.01.

(C) Average recognition accuracy of the uncrowded condition.

Binocular summation often refers to an increase in the binocular performance over the monocular performance. We, thus, examined a potential binocular advantage in crowding. To this end, we obtained both monocular and binocular crowding measurements in a subset of patients with glaucoma ($n = 13$). Binocular summation was quantified as the ratio of crowding of the binocular to that of the better eye (i.e., the eye with less crowding). A ratio value of greater than 1 would indicate binocular summation, whereas a value of equal to or less than 1 would indicate no binocular summation or inhibition. Note that, to follow the notion that the binocular summation ratio of greater than 1 indicates binocular summation, we considered the inverse values for calculating the summation ratio, that is, $1/$crowding. Figure 3B (i) shows the average of monocular crowding zone for the worse and better eyes as well as the binocular crowding zone. It should be noted that, here, the worse and better eyes are considered as the eyes with larger and smaller monocular crowding zones, respectively. Figure 3B (ii) plots the mean and individual data of the binocular ratio for crowding. As shown in Figure 3B (ii), the mean binocular ratio for crowding is $1$, $t_{(12)} = -0.015$, $P = 0.99$, suggesting that the binocular
A) Monocular crowding in glaucoma

i) An exemplary subject

- An exemplary subject. The extent of monocular crowding zone at each testing location is plotted in polar coordinates for the worse (solid line) and better (dashed line) eyes.

ii) Group average results

- Group average results. The extent of monocular crowding zone is compared between the worse (orange solid bounding boxes) and better (orange dotted bounding boxes) eyes for 2°, 4°, and both eccentricities. Measurements between the two eyes of a single subject are connected by a gray solid line. Note that the worse and better eyes were determined for each subject based on the mean deviation (MD) values from HFA 10-2 test. Note that * denotes statistical significance at a P value of less than 0.05.

B) Binocular summation of crowding

i) Monocular vs Binocular crowding

- Monocular versus binocular crowding. The boxplots represent the extent of monocular crowding zone for the worse and better eyes and binocular crowding zone. Note that here, the worse and better eyes refer to eyes with larger (more crowded) and smaller (less crowded) crowding zones, respectively. Each gray dot represents the data point from an individual subject.

ii) Binocular summation ratio

- Binocular summation ratio. Binocular summation ratio defined as the crowding of the binocular to that of the better eye. Note that we used the inverse values of crowding for calculating the binocular summation ratio to follow the notion that the binocular summation ratio of greater than 1 indicates the binocular summation.

**DISCUSSION**

In this study, we showed that glaucomatous damage is associated with increased crowding even in the parafoveal region corresponding with the central 8° visual field (−4° to +4°). We found a statistically significant increase in the binocular crowding in patients with glaucoma relative to age-similar normal controls.

More important, to control for the possible effect of relatively higher cognitive factors such as attention or memory lapse between the stimulus interval and the response interval and low-level sensory factors such as acuity limit or decreased contrast for a peripheral target location on the behavioral results, our study design included the uncrowded experiment as a control condition (Fig. 1A). The results showed that both patients with glaucoma and normal controls were able to recognize a “single letter” (i.e., a target without flankers) at a given retinal eccentricity with a high level of accuracy (95% and 98% for glaucoma and normal groups, respectively). This result further assured us that our subjects had no trouble recognizing the target letter at a given retinal location when presented alone. Therefore, the observed difference in crowding between glaucoma and normal cohorts is likely to be due to the crowding effect rather than decreased acuity or contrast sensitivity at a given...
target location or other high-level cognitive factors, such as a lack of attention or memory lapse. The binocular visual acuity of these patients with glaucoma (−0.01 vs. −0.07 logMAR) and log contrast sensitivity (1.70 vs. 1.92) were comparable with those of normal controls, further supporting the view that signal letter acuity or contrast sensitivity were not likely plausible explanations for our findings. It should also be noted that the time interval between the onset of the stimulus and the onset of the response panel was set to 500 ms, which is much longer than the duration that likely induces backward masking (<50 ms). We, however, cannot completely rule out the possible role of backward masking on our results. Even if backward masking had played a role, it should have affected both the glaucoma and the age-matched normal cohorts equally.

It should be noted that we compared the crowding effect between glaucomatous and normal vision under binocular viewing, because binocular visual recognition likely reflects what patients would experience in real life. However, to further investigate the linkage between the severity of glaucomatous damage on crowding, we went on to measure the monocular crowding in patients with glaucoma using within-subjects comparison. Comparing the monocular crowding between the worse and better eyes defined by the severity of glaucoma-induced damage (i.e., mean deviation values) indeed confirmed larger crowding with more severe glaucomatous damage. This, in turn, can lead to asymmetrical monocular crowding in glaucomatous vision. Here we used this opportunity to address the question of how the binocular asymmetry in glaucomatous damage can affect the way crowding effect is integrated between two eyes (i.e., summation or inhibition).

Our findings showed that the glaucomatous vision did not exhibit binocular summation of crowding as the binocular crowding was determined by the better eye. This absence of binocular summation is consistent with a previous study done by Siman-Tov et al.68

They examined target recognition performance (in d’prime) with or without distractors under both monocular and binocular viewing conditions. They found that the binocular summation was nearly absent when the target appeared in clutter for a stimulus duration of less than 240 ms. In contrast, the single target condition yielded binocular summation of about 1.4 (an 40% increase), as expected from previous findings.68,69-72 Taken together, this absence of binocular summation for crowding suggests that crowding is likely to start at least before the process of binocular combination or integration known to take place or occur in the primary visual cortex (V1).69,70 It is, however, worth mentioning that a bigger sample size may be required to confirm these results in a future study.

In normally sighted subjects, crowding has minimal impact on daily central vision tasks, because very little crowding exists in foveal or parafoveal regions. However, people with some clinical conditions such as amblyopia are known to experience considerable crowding, even in their foveal vision; their functional deficits in central vision tasks such as reading and word recognition have been shown to correlate with the increased foveal crowding.69,75-77 Because feature segmentation and integration are the core processes of visual recognition, crowding—the inability to isolate the target item from nearby distractors—is an essential bottleneck for visual recognition.69 Therefore, determining whether individuals with impaired vision experience increased crowding is an important step toward a better understanding of daily pattern vision in clinical populations.

Glaucoma is typically thought to be peripheral vision loss, with central vision being preserved by the damage until the end stage of the disease. For this reason, little attention has been paid to understanding central pattern recognition function in glaucoma. However, recent research65,66,71,74 using optical coherence tomography or retinal staining techniques has demonstrated significant structural damage even in early glaucoma. Such damage includes loss of RGCs or significant shrinkage of dendritic structure and cell body of remaining cells in the macula.

Various models have been put forward to explain the phenomenon of crowding. These models include, but are not limited to, low-level feature integration,64,65,67 mid-level visual processing such as grouping,77,78 substitution,79 summary statistics,80,81 saccade-confounded image statistics,82 and higher level attentional account.80 Although crowding is known to be a cortical phenomenon, the question regarding the exact mechanism and locus of the crowding remains a subject of debate. Despite various accounts of crowding, there is one common thread: crowding is ascribed to signals being pooled over a greater spatial extent (extensive pooling)81 either owing to bottom-up computations,35,45 such as hardwired integration fields, and/or top-down cognitive factors,90 such as a spotlight of attention.

Thus, it is reasonable to speculate that glaucomatous damage, such as loss of ganglion cells, may bring about changes in the way visual signals are integrated across space, thereby leading to changes in crowding. Numerous studies have shown alterations in spatial summation mechanisms after glaucomatous damage. Redmond et al.82 reported significant enlargement of Ricco’s area in early glaucoma with respect to healthy subjects. A similar increase in the extent of Ricco’s area was also observed by Mulholland et al.83 Ricco’s area is regarded as the spatial extent over which visual signals are integrated for the system to achieve threshold detectability and has been linked to ganglion cell density. It has been proposed that, to maintain threshold detectability in the presence of glaucomatous ganglion cell loss, the system actively compensates the loss by integrating signals over a larger area.84,85 This view is also consistent with the inverse relationship between threshold stimulus size and RGC density reported in a number of psychophysical studies.85-87 A close linkage between the sampling density of RGCs and the extent of spatial integration such as Ricco’s area or crowding zone has also been demonstrated by the work done by our group.86 Furthermore, King et al.80 provided the neural basis of changes in summation mechanisms following glaucomatous damage. They found that the size of receptive field sizes in the adult rat brain increases in response to experimentally induced ganglion cell death. The increase of receptive fields was proportional to the degree of glaucomatous damage, highlighting the close linkage between the size of signal integration zones and ganglion cell damage. It is also important to note that the macular RGC+ layer thickness is closely correlated with RGC counts,89-91 the thinner the layer gets, the more the RGCs are being lost.

We, however, acknowledge that a more quantitative relationship between the degree of crowding and severity of glaucomatous damage needs to be explored in future studies. Perhaps, a cross-sectional study with different stages of the disease progression, including preperimetric glaucoma,
will help us to further characterize the relationship between the two.

In summary, the current study shows that crowding is exacerbated in parafoveal vision in glaucoma and binocularly asymmetric glaucoma is associated with binocularly asymmetric crowding. Our findings are consistent with the view that glaucomatous damage brings about alterations in spatial pooling mechanisms. Furthermore, the absence of binocular summation for crowding observed in glaucomatous vision combined with the lack of binocular summation found in Siman-Tov et al.’s normal healthy vision support the view that crowding may start in the early stages of visual processing, at least before the process of binocular integration takes place.

Acknowledgments

Supported by NIH/NEI Grant R01 EY027857, Research to Prevent Blindness (RPB) / Lions’ Clubs International Foundation (LCIF) Low Vision Research Award, and Eyewitness Foundation of Alabama.

Disclosure: F. Shamsi, None; R. Liu, None; M. Kwon, None

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