Microscopic Eye Movements Compensate for Nonhomogeneous Vision within the Fovea

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Summary

Humans rely on the fovea, the small region of the retina where receptors are most densely packed, for seeing fine spatial detail. Outside the fovea, it is well established that a variety of visual functions progressively decline with eccentricity [1–5]. In contrast, little is known about how vision varies within the central fovea, as incessant microscopic eye movements prevent isolation of adjacent foveal locations [6–8].

In this study, we examined the discrimination of fine spatial patterns at different eccentricities within the fovea. To overcome the problems of previous studies, we relied on a combination of techniques. First, we developed a new procedure for localizing the line of sight, which greatly improved determination of the position of the stimulus on the retina over standard methods (Figures S1D and S1E; see Supplemental Experimental Procedures). Second, we presented stimuli under conditions of retinal stabilization [12–14]; that is, we updated localizing the line of sight, which greatly improved determination but not during normal viewing, even when the two bars were displayed at the same retinal (or monitor) location (Figure S2).

The two conditions in Figure 1D only differed for the consequences of oculomotor activity on retinal stimulation: fixational eye movements moved the stimulus on the retina during normal viewing but not under retinal stabilization (Figure 1C). Figure 2 summarizes the effects of eye movements in the normal viewing condition (see also Figure S3). Although minute, gaze shifts were not random; subjects looked preferentially toward the grating: to the left side of the monitor when the grating was displayed in the left bar and to the right side of the monitor when the grating appeared in the right bar (see examples in Figure 2A and Movie S1, Movie S2, and Movie S3). Across subjects, this difference in gaze position...
was significantly correlated with the eccentricity of the stimuli \( r = 0.75; p = 0.01 \) and was visible even with stimuli at 5° eccentricity, so that the mean horizontal gaze positions measured during presentation of the left and right gratings differed significantly at all tested eccentricity angles (Figure 2B).

Gaze shifts were primarily caused by microsaccades rather than fixational drifts. Microsaccades occurred frequently in the experiments and were clearly modulated by the task: they were more likely to occur in the periods immediately following the appearance of each grating (Figure S4C), and both their frequency (Figure S4A) and amplitude (Figure S4B) increased with the eccentricity of visual stimulation \[ rate: \text{F}(2,3) = 7.02, \text{p} = 0.03; \text{amplitude}: \text{F}(2,3) = 14.89, \text{p} = 0.005; \text{two-way ANOVA} \]. Furthermore, regardless of the eccentricity of the stimulus, microsaccades were always more likely to land close to the bar in which the grating was currently displayed than close to the other bar, while the probability of terminating on the background region far from the two bars was always low (Figure 3A). These precise movements were responsible for the average changes in gaze position of Figure 2B: when microsaccades were removed from the eye movement traces and the remaining periods of drifts were concatenated, the line of sight no longer moved away from the center of the screen (Figure 3B). Thus, in the normal condition, stimuli did not remain on the retina around their intended eccentricity angles as under retinal stabilization but moved, because of microsaccades, toward regions at lower eccentricities, where performance in the task was higher (Figure 1D).

Together, the results of Figures 1, 2, and 3 suggest that microsaccades normally compensate for nonuniform visual capabilities across the foveola: discrimination performance dropped significantly with eccentricity under retinal stabilization but not during normal viewing, when microsaccades enabled examination of the stimulus with the preferred retinal locus.

To directly assess the importance of microsaccadic gaze shifts, we examined performance in the normal (unstabilized) trials in which microsaccades did not occur. These trials were very rare, particularly during stimulation at larger eccentricities (only 4% of the trials at 15°), which by itself argues in favor of the importance of microsaccades in this task. Figure 4 compares performance with and without microsaccades for

![Figure 1. Experimental Procedure and Results](image)

(A) In a forced-choice task, subjects reported whether two gratings (11 cycles/degree tilted by ±45°) were parallel or orthogonal.

(B) Gratings appeared within two rectangular noise bars centered at the desired eccentricity \( d \). They were displayed sequentially first in the left and then in the right bar while subjects maintained fixation at the center of the display (cross).

(C) Stimuli were either displayed at fixed positions on the screen (normal) and normally moved on the retina because of fixational eye movements or at fixed locations on the retina (stabilized) and moved on the display under computer control to compensate for the subject’s eye movements.

(D) Average subject performance \( (n = 4) \) as a function of the stimulus eccentricity in the two conditions. Both proportions of correct responses (filled symbols) and \( d’ \) values (empty symbols) are shown. In each condition, asterisks mark statistically significant differences with respect to the proportions of correct responses at 5° \( (*p < 0.05; **p < 0.005; \text{two-tailed paired t test}) \). Error bars represent SEM. See also Figures S1 and S2.

![Figure 2. Gaze Location in Normal, Unstabilized Trials](image)

(A) Examples of fixational eye movements. Red and blue segments represent microsaccades and drifts, respectively.

(B) Average horizontal position of the center of gaze during presentation of each of the two gratings. In both (A) and (B), different panels show data obtained with stimuli at different eccentricities. Error bars represent SEM. Asterisks mark significant differences \( (p < 0.05; \text{two-tailed paired t test}) \). The center of gaze is defined in this study as the point on the screen projecting onto the center of the preferred retinal locus of fixation. This point was estimated by means of a preliminary calibration procedure, in which the observer maintained prolonged fixation on markers at known positions on the display. See also Figure S3 and Movies S1, S2, and S3.
two subjects who were tested extensively in order to collect sufficiently large pools of drift-only trials. For both observers, performance dropped in the absence of microsaccades: levels of discrimination at 15° eccentricity were lower than those measured in the presence of microsaccades, an effect already significant at 10° eccentricity for one subject. This reduction in performance was similar to that observed under retinal stabilization; for both subjects and at all three eccentricity angles, results in the absence of microsaccades were statistically indistinguishable from those measured with a stabilized retinal image (p > 0.21; paired two-tailed t test).

These impairments were not caused by lack of attention or the absence of the temporal transients of microsaccades. Figure 4 also compares levels of performance in the trials in which microsaccades precisely relocated gaze, enabling the retinal projections of both gratings to fall within 5° eccentricity and those in which microsaccades occurred but were not as precise. Performance was significantly higher in the trials in which both gratings were accurately fixated. Thus, precise microsaccadic relocations of gaze were necessary for ensuring approximately equal levels of performance during viewing of stimuli at different foveal eccentricities.

The results of this study show that fine spatial vision is not uniform across the central fovea but drops sharply with minute displacements from the preferred retinal locus of fixation. Although idiosyncratic differences exist, cone density is known to decrease rapidly within the fovea outside a very small region of a few squared arcminutes [15]. Thus, the decline in performance with eccentricity observed in our experiments could be the consequence of the variable distribution of cone spacing. However, our stimuli were well within the spatial frequency limits of foveal vision [16] and should have been easily discriminable at all tested eccentricities. Furthermore, subjects used microsaccades to bring the stimulus close to the preferred retinal locus of fixation, a region which not always coincides with the area of highest cone density [17]. These observations suggest that other factors, in addition to cone spacing, contributed to the eccentricity effects shown in Figures 1 and 4. An interesting possibility is that these factors are not restricted to the spatial domain but that changes in temporal sensitivity may also occur within the fovea. Fine spatial vision seems to depend on the temporal modulations resulting from ocular drift, the smooth motion of the eye that continually occurs during the intersaccadic periods [14, 18–22]. Thus, both the local spatial and temporal characteristics of neural processing may be responsible for the emergence of a preferred fixation locus, a region in which neurons optimally extract fine spatial information from temporal modulations.

These findings also provide an answer to the long-investigated and controversial question of the visual function of microsaccades [7, 8]. Microsaccades have been deemed necessary to prevent, during natural viewing, the perceptual fading experienced in the laboratory when retinal image motion is eliminated [23, 24], but the plausibility of this theory has been criticized on multiple grounds [7, 25–28]. In our experiments, image fading did not appear to play any obvious role in the production of microsaccades. Subjects did not notice any fading, not even under retinal stabilization, possibly because stimuli were flashed at high contrast and for brief periods, with an abrupt change caused by the blank interval in between the two gratings, a procedure that minimized neural adaptation. Furthermore, the temporal transients resulting from microsaccades were by themselves not sufficient to improve performance. As shown in Figure 4, discrimination was impaired when microsaccades did occur but were not precisely directed toward the stimuli. Thus, our findings cannot be explained by a mere refreshing of the image following microsaccades.

In contrast, our results show that microsaccades are critical for high-acuity judgments, as they serve, at a microscopic scale, the same explorative function as larger saccades, as long hypothesized [29]: both movements reposition the stimulus on the retina to enable its examination at a finer level of detail. These data provide an explanation for the previous observation of precisely directed microsaccades in high-acuity visuomotor tasks [30] and are consistent with recent findings highlighting the similarities between microsaccades and saccades in terms of the underlying neural substrate [31, 32], their impact on neural responses [33], and their associated perceptual consequences [34–36]. Furthermore, our results imply that fine spatial vision may be compromised when microsaccades are not precisely executed, raising the possibility for a causal link between the abnormal fixational eye movements and the reduced visual acuity that coexist in various disorders [37–44]. Further work is necessary to...
investigate this hypothesis as well as the factors responsible for the emergence of a preferred fixation locus in the fovea.

Experimental Procedures

Stimuli were rendered by means of a custom-developed system for flexible gaze-contingent display control [45] on a fast-phosphor cathode ray tube monitor (Iyama HM204DT) at a vertical refresh rate of 150 Hz. Stimuli were observed monocularly with the right eye, while the left eye was patched. The movements of the right eye were measured by means of a Generation 6 Dual Purkinje Image eye tracker (Fourward Technologies) and sampled at 1 kHz. A dental-imprint bite bar and a headrest prevented head movements. Informed consent was obtained from all the participants following the procedures approved by the Boston University Charles River Campus Institutional Review Board.

Experiments were conducted in complete darkness with visual stimulation restricted to the two rectangular bars (5’ × 20’) centered at the same absolute eccentricity angle on the horizontal meridian in the nasal and temporal hemifields. In the normal condition, eccentricity angles were measured on the screen relative to the required location of fixation (the center of the display), as it is customary in behavioral measurements of visual functions. Stimuli remained immobile on the screen and normally shifted on the retina with eye movements. In the stabilized condition, eccentricity angles were measured on the retina, relative to the center of gaze (the estimated center of the preferred retinal locus of fixation). Stimuli moved on the screen to compensate for the subject’s eye movements, so that they remained at fixed retinal locations. Subjects did not report any stimulus fading under retinal stabilization and were, in general, unable to tell whether or not a trial was stabilized. To avoid possible stabilization artifacts, performance under retinal stabilization was evaluated over the trials without microsaccades. Results did not change if the stabilized trials with microsaccades were also included in the analysis.

Supplemental Information

Supplemental Information includes Supplemental Experimental Procedures, four figures, and three movies and can be found with this article online at http://dx.doi.org/10.1016/j.cub.2013.07.007.

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