

Brief period of 'blindness' is essential for vision

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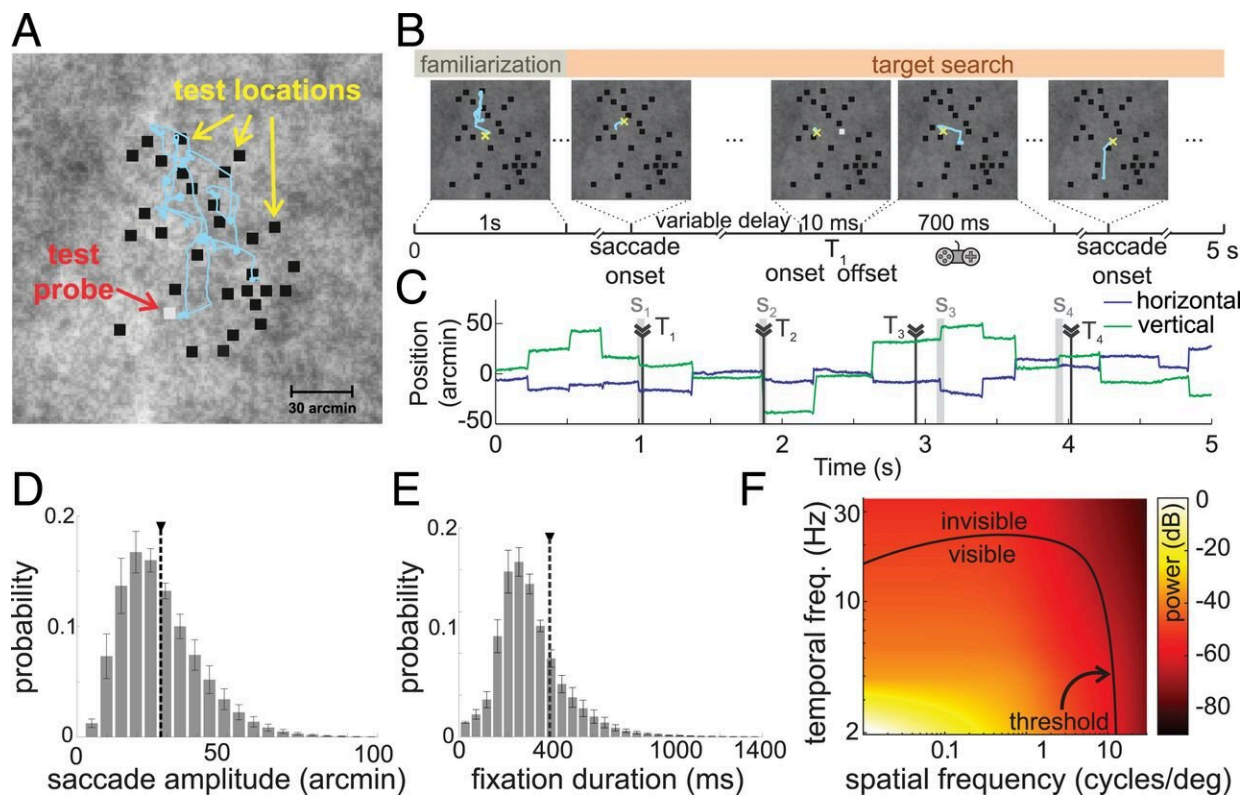


Fig. 1. A virtual grooming task. Observers were instructed to search for fleas hiding within the animal’s fur (a naturalistic noise field). (A) Thirty dots (5’5’ width) were distributed at random test locations across the central 2°2° region of the display. Subjects were told that a few of these dots were fleas and would distinguish themselves from the remaining dust particles by occasionally jumping (a 10-ms contrast pulse; the probe). The subject’s goal was to “catch” each flea as soon as it jumped by pressing a button on a joypad. (B–D) Example of a trial. (B) Following an initial familiarization period (1 s), the onset of a saccade triggered, with variable delay, a probe (TkTk) at one of the test

locations. Both location and timing were selected in real time according to the observer's eye movements to test performance at various positions in the fovea and lags relative to saccades. The yellow cross and cyan segments represents the center of gaze and eye movements, respectively. (C) Gaze position during the course of the trial. Each probe was associated with the closest saccade (SkSk in C). Only probes with no more than one saccade within a $\pm 200 \pm 200$ -ms window were selected for data analysis. (D and E) Characteristics of eye movements. Shown are average distributions of saccade amplitude (D) and intersaccadic intervals (E) across $N = 6$ observers. Error bars represent SEM. Vertical dashed lines mark the means of the distributions. (F) Power spectrum of the luminance flow delivered to the retina by the recorded saccades. The black line marks contrast sensitivity thresholds in humans [data from Kelly (33)]. The small saccades recorded in this experiment yield visual signals well within the ranges of spatiotemporal sensitivity. Credit: DOI: 10.1073/pnas.2101259118

Fixational eye movements are tiny movements of the eye—so small we humans aren't even aware of them. Yet they play a large role in our ability to see letters, numbers, and objects at a distance.

In a new paper published in *Proceedings of the National Academy of Sciences*, researchers at the University of Rochester, including Michele Rucci, a professor of brain and cognitive sciences, and Janis Intoy, a postdoctoral research associate in Rucci's lab, further cement the evidence for the important role of these tiny movements. By studying how a type of fixational eye movement called a microsaccade affects the foveola, a small region at the center of the retina, the researchers provide important foundational information that can lead to improved treatments and therapies for [vision impairments](#).

A tiny, but essential, region of the eye

Although the foveola is tiny, it is essential for seeing fine details and

conducting everyday tasks such as searching for a friend in a crowd or reading distant road signs while driving. Because the region is so small, however, we need to constantly shift our gaze to allow the foveola to get a full view of the world, similar to rotating a telescope to get a full view of a scene. Unlike when we might rotate a telescope, however, our eyes make most of these gaze shifts, especially the smallest ones, on their own, often beneath our awareness. But the gaze shifts are critical for vision; says Intoy, "How well we see at any given moment is tightly linked to how and when we shift our gaze."

The researchers focused on microsaccades, tiny rapid gaze shifts that frequently occur when we're examining fine details. It's long been known that vision is transiently impaired during larger gaze shifts, such as those we are aware of making, for instance looking back and forth between two computer screens. This phenomenon of transiently impaired vision is known as saccadic suppression. Until now, however, it was unknown whether a suppression also occurs during microsaccades and whether that would affect visibility in the foveola.

Says Rucci: "In our lab we have the high-resolution tools to study vision at this small scale, whereas other research has historically focused on the peripheral regions of the eye, where such precision and accuracy are not required."

A period of blindness

The researchers recorded microsaccades in human observers who were engaged in a computer task— searching on the screen for "fleas" jumping in a patch of "fur," a task that resembles social grooming in primates.

What the researchers found was surprising.

Immediately before and immediately after participants' gaze shifted, the participants could not see the fleas, even when they were looking directly at them.

"We observed that microsaccades are accompanied by brief periods of visual suppression during which we are essentially blind," Intoy says.

However, the researchers found that vision recovered rapidly at the center of the gaze and continued to improve, so that vision was overall transiently enhanced in this region after the saccade.

"Our results show that the very center of gaze undergoes drastic and rapid modulations every time we redirect our [gaze](#)," Intoy says. "This brief loss of [vision](#) likely occurs so that we do not see the image of the world shifting around whenever we move our eyes. By suppressing perception during saccades, our visual system is able to create a stable percept."

Future research will determine more about this phenomenon and how humans control eye movements to balance the saccadic suppression with the visual enhancement that follows.

More information: Janis Intoy et al, Fast and nonuniform dynamics of perisaccadic vision in the central fovea, *Proceedings of the National Academy of Sciences* (2021). [DOI: 10.1073/pnas.2101259118](https://doi.org/10.1073/pnas.2101259118)

Provided by University of Rochester

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