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Microsaccades and Blinks Trigger Illusory Rotation in the “Rotating Snakes” Illusion

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Certain repetitive arrangements of luminance gradients elicit the perception of strong illusory motion. Among them, the “Rotating Snakes Illusion” has generated a large amount of interest in the visual neurosciences, as well as in the public. Prior evidence indicates that the Rotating Snakes illusion depends critically on eye movements, yet the specific eye movement types involved and their associated neural mechanisms remain controversial. According to recent reports, slow ocular drift—a nonsaccadic type of fixational eye movement—drives the illusion, whereas microsaccades produced during attempted fixation fail to do so. Here, we asked human subjects to indicate the presence or absence of rotation during the observation of the illusion while we simultaneously recorded their eye movements with high precision. We found a strong quantitative link between microsaccade and blink production and illusory rotation. These results suggest that transient oculomotor events such as microsaccades, saccades, and blinks, rather than continuous drift, act to trigger the illusory motion in the Rotating Snakes illusion.

Introduction

The “Rotating Snakes Illusion” produces the perception of strong illusory rotation in most observers and has generated an extraordinary amount of academic and popular interest (Fig. 1) (Martinez-Conde and Macknik, 2010; Ramachandran and Rogers-Ramachandran, 2010). Here we demonstrate a direct relationship between the onsets of both microsaccades and blinks and perceived illusory rotation, showing that transient oculomotor events act to trigger the Rotating Snakes effect.

More than three decades ago, Fraser and Wilcox (1979) developed a repetitive spiral arrangement of sawtooth luminance gradients that elicited illusory motion (Fig. 1A). The “Peripheral Drift Illusion”—a variant on the Fraser-Wilcox effect—later incorporated circular rather than spiral arrangements of sawtooth luminance gradients (Fig. 1B) (Faubert and Herbert, 1999). Kitaoka and Ashida (2003) subsequently developed a new and improved version of the Peripheral Drift Illusion, with stepwise rather than sawtooth luminance profiles (Fig. 1C) and fragmented or curved edges rather than straight edges (Fig. 1D,E). The resultant Rotating Snakes are characterized by the periodic placement of colored patches of four different luminances along the circumferences of concentric circles. This layout generates the perception of rotational motion in the direction that follows the colored patches from black to blue to white to yellow to black (Fig. 1E) (Kitaoka, 2005). A grayscale version of the illusion is effective perceptually as well, and drives the responses of directionally selective visual neurons, so long as the luminance relationship between the patches is preserved (Conway et al., 2005).

The Rotating Snakes is a spatiotemporal illusion, in which both spatial layout and temporal stimulation—usually brought about by eye movements (Backus and Oruç, 2005; Conway et al., 2005; Murakami et al., 2006; Beer et al., 2008; Ashida et al., 2010)—are critical to the perception. The specific eye movements and neural mechanisms involved remain controversial, however.

One theory explaining the phenomenon is that contrast-induced latency differences in neural responses and/or neural adaptation trick the brain’s motion detectors into responding to the stationary patches as if to actual motion (Faubert and Herbert, 1999; Backus and Oruç, 2005; Conway et al., 2005; Kitaoka, 2006). For instance, latency differences between higher- and lower-contrast patches may generate equivalent neuronal responses in areas V1 and MT to those induced by actual motion (Conway et al., 2005). If this proposal is correct, any “reset” or “refresh” (Faubert and Herbert, 1999) of the retinal image—caused by microsaccades, saccades, blinks, flash, or displacement of the image—should trigger the motion signal.

A second theory is that the temporal and spatial integration of residual motion vectors—due to incorrectly compensated eye movements—causes the perception of illusory motion (Murakami et al., 2006; Beer et al., 2008; Fermüller et al., 2010). According to this proposal, slow ocular drift triggers the motion signal.

Here we presented subjects with the Rotating Snakes Illusion while we measured their oculomotor output—including microsaccades, blinks, and fixational instability/drift—in correlation to their perception (Fig. 2). Our results show a strong quantitative link between microsaccade and blink production and subsequent illusory rotation.
Materials and Methods

Subjects

Eight subjects (6 males, 2 females) with normal or corrected-to-normal vision participated in this study. Each subject participated in three sessions: one training session and two experimental sessions of ~50 min each. Six of the subjects were naive and were paid $15 per session. Experiments were performed under the guidelines of the Barrow Neurological Institute’s Institutional Review Board (protocol 04BN039) and written informed consent was obtained from each participant.

Figures 3D, 4, and 5D include data from seven subjects only—one subject was excluded due to an insufficient number of transitions toward illusory rotation (i.e., the subject produced only three transitions throughout the whole experiment). Figure 6 includes data from five subjects—three subjects were excluded because of their low blink rates (one or no blinks per trial).

Eye movements recordings and analyses

Eye position was recorded noninvasively in both eyes with a fast video-based eye movement monitor (EyeLink 1000; SR Research) at 500 samples per second (instrument noise, 0.01° RMS).

We identified blink periods as the portions of the EyeLink 1000 recorded data where the pupil information was missing. We added 200 ms before and after each period to further include the initial and final parts of the blink, where the pupil is partially occluded. We moreover removed those portions of the data corresponding to very fast decreases and increases in pupil area (>20 units per sample) plus the 200 ms before and after. Such periods are probably due to partial blinks, where the pupil is never fully occluded (thus failing to be identified as a blink by the EyeLink 1000 software) (Troncoso et al., 2008b).

We identified microsaccades automatically with an objective algorithm (for details using λ = 6, see Engbert and Kliegl, 2003). To reduce the amount of potential noise (Engbert, 2006), we analyzed only binocular microsaccades (that is, microsaccades with a minimum overlap of

Figure 3. Microsaccade characterization and perceptual reports. **A**, Distribution of microsaccade magnitudes for the subject average (n = 8 subjects). **B**, Peak velocity–magnitude relationship for all microsaccades (n = 23,610 microsaccades; n = 8 subjects). **C**, Distribution of intermicrosaccadic intervals for the subject average (n = 8 subjects). **D**, Distribution of durations of rotating (red) and stationary (blue) periods during the main experimental condition, as indicated by the subjects’ report (n = 7 subjects). Shaded red and blue areas indicate SEM across subjects. **E**, Examples of recorded eye positions in relation to perceptual reports. The gap in the lower eye-position trace, denoted by the dashed line, represents a blink. Arrows identify microsaccades. Drifts are the periods in between microsaccades (or between a blink and a microsaccade).
one data sample in both eyes) (Laubrock et al., 2005; Engbert, 2006; Rolfs et al., 2006; Troncoso et al., 2008a). We also imposed a minimum intersaccadic interval of 20 ms so that overshoot corrections were not categorized as microsaccades (Møller et al., 2002; Otero-Millan et al., 2008; Troncoso et al., 2008a). One subject produced microsaccades that were often larger than 1°; thus, we chose a maximum microsaccade magnitude threshold of 2° (Betta and Turatto, 2006; Martinez-Conde et al., 2006; Troncoso et al., 2008a). Across subjects, >85% of all microsaccades were smaller than 1°. Figure 3A plots the distribution of microsaccade magnitudes for the subject population [up to 3° (Otero-Millan et al., 2008)]. Of these, we studied further only microsaccades smaller than 2° (n = 23,610) (Troncoso et al., 2008b). Equivalent results were obtained for 1- or 3-degrees-magnitude thresholds (data not shown) (Troncoso et al., 2008b). Figure 3B shows the peak velocity–magnitude relationship for all analyzed microsaccades. Table 1 summarizes various microsaccade and perceptual parameters for individual subjects and for the subject average.

**Experimental design**

Subjects rested their head on a chin/forehead-rest 57 cm from a video monitor (Reference Calibrator V, 60 Hz refresh rate; Barco). Each experimental session included 48 trials, half of those corresponding to the main experimental condition and the other half corresponding to a control condition, as detailed below. Trials of both types were pseudorandomly interleaved.

**Main experimental condition: perception of illusory rotation in the Rotating Snakes Illusion.** Subjects fixated on a small red dot (0.25° in diameter) on the center of the monitor’s screen and pressed a key to start each trial. A simplified version of the Rotating Snakes Illusion (including only two 8°-diameter sets of concentric rings, or snake-disk) appeared on the screen upon the key press. The two disks were presented on opposite sides of the screen, with the center of each disk separated from the fixation dot by 9° (Fig. 2A). To avoid adaptation across trials, the disk pairs were randomly positioned from one trial to the next, toward one of eight chosen points around the compass (45°, 90°, 135°, 180°, 225°, 270°, 315°).

Subjects simultaneously and continuously reported whether the disks were stationary (button press) or rotating (button release) (Fig. 2B). Figure 3D illustrates the distribution of stationary and rotating periods for the subject average, and Table 1 summarizes the corresponding parameters.

After 30 s, all stimuli disappeared and the trial ended. To control for initial stimulus-onset transient effects at the start of each trial, we analyzed only those data recorded after the first button press.

**Control condition: perception of physical rotation.** Here we presented disk pairs that looked very similar to those in the main experimental condition (Rotating Snakes Illusion), but did not generate the perception of illusory motion (Fig. 2C) (Kuriki et al., 2008). These disks rotated physically (0.01 cycles per second), starting and stopping according to (i.e., replaying) the timing of the perceptual transitions that each subject had reported in the most recent trial of the main condition. All other details, including the subjects’ task (Fig. 2D), were as in the main condition.

We calculated the average latency between the transitions in physical rotation and the reported transitions (subjects’ button presses/releases;
Table 1. Microsaccade and perceptual parameters

<table>
<thead>
<tr>
<th></th>
<th>S1</th>
<th>S2</th>
<th>S3</th>
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<th>S6</th>
<th>S7</th>
<th>S8</th>
<th>Average ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of microsaccades</td>
<td>1908</td>
<td>2690</td>
<td>1538</td>
<td>2050</td>
<td>1143</td>
<td>2289</td>
<td>2907</td>
<td>1619</td>
<td>2000 ± 200</td>
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<tr>
<td>Microsaccade rate (ms/°)</td>
<td>0.8</td>
<td>1.3</td>
<td>0.6</td>
<td>0.8</td>
<td>0.48</td>
<td>0.9</td>
<td>1.1</td>
<td>0.6</td>
<td>0.8 ± 0.1</td>
</tr>
<tr>
<td>Microsaccade magnitude (deg)</td>
<td>0.39</td>
<td>0.44</td>
<td>0.5</td>
<td>0.23</td>
<td>0.6</td>
<td>0.47</td>
<td>1.0</td>
<td>0.6</td>
<td>0.5 ± 0.1</td>
</tr>
<tr>
<td>Microsaccade duration (ms)</td>
<td>4.9</td>
<td>8</td>
<td>9</td>
<td>4.3</td>
<td>12</td>
<td>8</td>
<td>10</td>
<td>6</td>
<td>8 ± 1</td>
</tr>
<tr>
<td>Microsaccade peak velocity (deg/°)</td>
<td>23</td>
<td>26</td>
<td>27</td>
<td>14</td>
<td>27</td>
<td>25</td>
<td>50</td>
<td>30</td>
<td>28 ± 4</td>
</tr>
<tr>
<td>Control condition: perception of physical rotation</td>
<td></td>
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<tr>
<td>Main experimental condition: perception of illusory rotation in the Rotating Snakes Illusion</td>
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<tr>
<td>Number of transitions to rotation (n)</td>
<td>72</td>
<td>39</td>
<td>224</td>
<td>3</td>
<td>48</td>
<td>28</td>
<td>257</td>
<td>288</td>
<td>120 ± 40</td>
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<tr>
<td>Duration of illusory rotation periods (subsequent to the first illusory period) (s)</td>
<td>1.2</td>
<td>2.0</td>
<td>1.3</td>
<td>0.8</td>
<td>3.4</td>
<td>0.8</td>
<td>0.8</td>
<td>1.7</td>
<td>1.5 ± 0.3</td>
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<td>Duration of first illusory rotation period after stimulus onset (s)</td>
<td>3.5</td>
<td>8.2</td>
<td>2.5</td>
<td>4.5</td>
<td>5.3</td>
<td>4.1</td>
<td>2.0</td>
<td>2.7</td>
<td>4.1 ± 0.7</td>
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<td>Time spent in rotation periods (%)</td>
<td>7</td>
<td>8</td>
<td>22</td>
<td>0</td>
<td>14</td>
<td>2</td>
<td>15</td>
<td>36</td>
<td>13 ± 4</td>
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<tr>
<td>Number of transitions to stationary (n)</td>
<td>69</td>
<td>38</td>
<td>214</td>
<td>3</td>
<td>45</td>
<td>25</td>
<td>253</td>
<td>268</td>
<td>110 ± 40</td>
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<td>Duration of stationary periods (s)</td>
<td>17</td>
<td>25</td>
<td>4.8</td>
<td>410</td>
<td>23</td>
<td>49</td>
<td>4.5</td>
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<td>92</td>
<td>78</td>
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<td>85</td>
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<td>Control condition: perception of physical rotation</td>
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</tr>
<tr>
<td>Reaction times to transitions to rotating (ms)</td>
<td>440</td>
<td>600</td>
<td>390</td>
<td>450</td>
<td>500</td>
<td>500</td>
<td>430</td>
<td>440</td>
<td>470 ± 20</td>
</tr>
<tr>
<td>Reaction times to transitions to stationary (ms)</td>
<td>460</td>
<td>480</td>
<td>440</td>
<td>480</td>
<td>500</td>
<td>600</td>
<td>480</td>
<td>430</td>
<td>490 ± 20</td>
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<td>Correctly identified transitions to rotating (%)</td>
<td>95</td>
<td>94</td>
<td>98</td>
<td>100</td>
<td>94</td>
<td>78</td>
<td>91</td>
<td>95</td>
<td>94 ± 2</td>
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<tr>
<td>Correctly identified transitions to stationary (%)</td>
<td>100</td>
<td>100</td>
<td>97</td>
<td>100</td>
<td>91</td>
<td>90</td>
<td>95</td>
<td>100</td>
<td>97 ± 1</td>
</tr>
</tbody>
</table>

Fig. 4A). To do this, we discarded latency values >1500 ms [1% of all reported transitions; such latencies were considered too long to be meaningful (Einhäuser et al., 2008; Troncoso et al., 2008b)]. Table 1 summarizes the corresponding parameters.

Event-triggered averages

Event-triggered averages (such as those in Fig. 4) were calculated by averaging a multisecond window of data around or after the trigger event [reported transitions (Fig. 4B, D), physical transitions (Fig. 4E), and microsaccade onset (Fig. 4C)]. Thus, the event-triggered average (Fig. 4B) represents the average percentage increase from chance in microsaccade rate at any given time during the 3 s window plotted around the reported transitions (all reported transitions are aligned at time = 0 ms). If there were <3 s of data around a trigger event (due to the edge of the recording time during that specific trial), the 3 s window was dynamically reduced to fit the available data. For the analyses where the trigger event was a transition, we avoided using the same data point multiple times by considering only the time from the previous transition until the next transition (Einhäuser et al., 2008; Troncoso et al., 2008a).

Individual subjects’ event-triggered averages concerning microsaccade rates and transition probabilities were calculated with a 200 ms sliding window that advanced one data sample (2 ms) per iteration. For each subject, calculations were made separately for each eye and then averaged between the two eyes. The population average and SEM were then calculated across subjects.

Probability of perceptual transitions triggered by microsaccade onsets

Only those microsaccades that are produced during a stationary period can trigger the transition to a period of illusory rotation. Conversely, only those microsaccades produced during a period of rotation can trigger a transition to a stationary period. Thus, to calculate the probability of transitions to and from rotating periods as a function of microsaccade onsets, we considered only microsaccades produced in the previous period. To control for chance correlations between microsaccades and transitions, we repeated the above calculations, this time replacing microsaccades with the same number of events, now randomly distributed in time, for each subject. This determined a baseline of chance correlation for each individual subject, from which we calculated the percentage increase in the probability of perceptual transitions after actual microsaccades (Fig. 4C). This analysis is comparable to the cross-correlogram shuffling often used to rule out spurious correlations between potential presynaptic and postsynaptic neurons (Alonso and Martinez, 1998; Martinez and Alonso, 2001).

ROC analysis

We used a receiver operating characteristic (ROC) analysis (Green and Swets, 1966; Britten et al., 1992; Hernández et al., 2002; Romo et al., 2002, 2004; Feierstein et al., 2006) to quantify how well microsaccade rate predicts the type of perceptual transition (i.e., toward stationary vs rotating “snakes”) in the Rotating Snakes Illusion. This analysis makes no assumptions about the underlying probability distributions (Feierstein et al., 2006). The area under the ROC curve provides a measure of the discriminability of two signals and is directly related to the overlap of the two distributions of responses that are compared (Feierstein et al., 2006). In our case, the area under the ROC curve can be interpreted as the probability that an ideal observer, given the microsaccade rate around a particular transition, will correctly determine the type of transition (toward stationary vs toward rotating percepts). An ROC area of 0.5 corresponds to completely overlapping distributions (the ideal observer cannot discriminate between the two types of transition); an area >0.5 corresponds to transitions that can be discriminated correctly from the microsaccade rate. At each point in time, we compared the microsaccade rate distributions for transitions toward rotating percepts (true-positive rate) and transitions toward stationary percepts (false-positive rate) in each subject. The ROC curve is the plot of the probability of true positives as a function of the probability of false positives for all possible criterion response levels. We performed a sliding ROC analysis (kernel width 500 ms, slid in 2 ms increments) to calculate each subject’s area under the ROC curve at each time point around the transition (Fig. 4B, D). To determine the time point at which the ideal observer became better than chance, we calculated significance, using a permutation procedure (Siegel and Castellan, 1988; Hernández et al., 2002; Romo et al., 2002, 2004; Feierstein et al., 2006) with 1000 shuffles for each subject and a criterion level value of p < 0.01.

Fixation instability analysis

We measured the variability of ocular drift following Murakami et al. (2006). First, we bandpass (1–31 Hz) filtered the horizontal eye position and calculated the instantaneous eye velocity as the rate of displacement from one data sample to the next. Then, we detected and removed microsaccades (i.e., we used a 10% velocity threshold to detect microsaccades and we ignored all data within ±26 ms around each microsaccade).
Finally, we fit the distribution of the remaining instantaneous eye velocities with a Gaussian and used its standard deviation as the index of fixation instability (Fig. 5B,E). Fixational instability results calculated using the horizontal eye position only (Murakami et al., 2006) were equivalent to those calculated using both horizontal and vertical components (data not shown).

**Within-subjects analyses**

We performed an analysis of covariance, with microsaccade magnitude (bins of 0.2°) and subject as independent variables, to calculate the regressions in Figure 5, C and D. We calculated the percentage of microsaccades followed by transitions toward rotation within 1 s of microsaccade onset for each microsaccade magnitude bin and each subject (Fig. 5C). We also calculated the average time from microsaccade onset to the next transition, whenever the next transition occurred within 1 s, for each microsaccade magnitude bin and each subject (Fig. 5D).

We calculated the index of fixation instability during the 1 s period preceding the transition toward rotation/stillness (for those periods lacking microsaccades and/or blinks) (Fig. 5E).

**Results**

**Main experimental condition: perception of illusory rotation in the Rotating Snakes Illusion**

Subjects fixated a small spot while viewing a simplified version of the Rotating Snakes Illusion (Fig. 2A). They continuously reported (via button press) whether the disks (“snakes”) were stationary (button press) or rotating (button release). Their eye movements were recorded simultaneously with high precision (for details, see Materials and Methods, above). Figure 2B describes a typical epoch during a trial.

As with other bistable stimuli paradigms (Spillmann and Kartenbach, 1992; van Dam and van Ee, 2005, 2006a,b; Martinez-Conde et al., 2006; Troncoso et al., 2008b), subjects reported that their perception oscillated between two alternating states (stillness vs rotation). We found a clear positive relationship between increased microsaccade rates and the perception of snake rotation. Before transitions to a rotation period, microsaccade rates increased (Fig. 4B, red line). Conversely, before transitions to a period of stillness, microsaccade rates decreased (Fig. 4B, blue line). The results were statistically significant (one-tailed t tests; p = 0.01), revealing a direct quantitative link between microsaccade production and the perception of illusory rotation.

To further establish the potential role of microsaccades in driving the Rotating Snakes Illusion, we determined the numbers of transitions to stillness versus rotation periods after microsaccade onsets (Fig. 4C). Microsaccade onsets were followed by increased transitions to rotation and decreased transitions to stillness periods (a 106% peak increase and a 31% peak decrease from random, respectively), suggesting a causal relationship between microsaccades and illusory rotation in the Rotating Snakes Illusion. At the very least, we cannot exclude a causal role of microsaccades in driving the Rotating Snakes Illusion, and specifically the perceptual transitions that characterize it. Equivalent results were obtained for naive versus nonnaive subjects, and for maximum microsaccade magnitude thresholds set to 1°, 2°, or 3° (data not shown) (Troncoso et al., 2008b).

**Control condition: perception of physical rotation**

We tested an alternative interpretation: that microsaccades do not drive the Rotating Snakes Illusion, but the illusion instead causes both the subjects’ reports (button press/release) and the change in microsaccade rate. If this were true, a change in microsaccade rate might precede the subjects’ reports (button press/release), as found in Figure 4B, but it would not precede the actual perceptual transitions. To test this possibility, we pseudo-randomly interleaved a control condition in which the subjects detected physical rotations for real moving stimuli that simulated the Rotating Snakes Illusion (for details, see Materials and Methods, above; Fig. 2C) (Kuriki et al., 2008). We then used the latency between physical transitions and reported transitions to estimate the timing of perception in the main illusory condition (van Dam and van Ee, 2005, 2006a,b; Troncoso et al., 2008b). Figure 2D describes a typical epoch during a trial. Subjects indicated correctly 95 ± 2% of the physical transitions with an average latency of 480 ms (SD ± 100 ms; Fig. 4A, vertical gray band). The average latency and SD values were consistent with those reported previously for various bistable stimuli (van Dam and van Ee, 2006a,b; Einhäuser et al., 2008; Troncoso et al., 2008b).

Next, we calculated the rates of microsaccades around both the reported transitions (Fig. 4D) and the physical transitions (Fig. 4E; equivalent analyses as in Fig. 4B). Microsaccade rates decreased before reported transitions to physical stillness as well
as before reported transitions to physical rotation (a 66% decrease in microsaccade rate 200 ms before the reported transition; Fig. 4D). This is contrary to our findings with illusory rotation: microsaccade rates increased before reported transitions to rotation periods and decreased before reported transitions to stationary periods in the Rotating Snakes Illusion (Fig. 4B). If perceptual rotation drove microsaccade production, then microsaccade rates should vary comparably for illusory motion (main condition; Fig. 4B) and physical motion (control condition; Fig. 4D); however, they do not. Thus, perceptual transitions in the Rotating Snakes Illusion do not drive changes in microsaccade rate.

In Figure 4E, the data from Figure 4D is realigned around the physical (rather than the reported) transitions. Microsaccade rates decreased after physical transitions, regardless of whether the change was toward rotation or stillness. This result is consistent with previous studies showing that changes in peripheral stimulation cause a transient drop in the rates of saccades and microsaccades, followed by a temporary enhancement (Engbert and Kliegl, 2003; Galfano et al., 2004; van Dam and van Ee, 2006b; Troncoso et al., 2008b). This transient course of microsaccadic inhibition may indicate a fast reflex of the oculomotor system to sudden changes in visual input (Laubrock et al., 2005).

The control condition provided us with the latency between changes in physical rotation and their associated reported transitions (Fig. 4A). This latency, when applied to the Rotating Snakes Illusion results, allowed us to estimate the timing of perceptual transitions due to changes in illusory rotation with respect to reported transitions. To accomplish this, we overlaid the same latency from Figure 4A, vertical gray band, on the data from the Rotating Snakes Illusion condition (Fig. 4B). The results showed that changes in microsaccade rate preceded the perception of the Rotating Snakes Illusion: microsaccade rates began changing (~130 ms before the perceptual transitions and 810 ms before the reported transitions in illusory rotation. Thus, microsaccades may trigger the perceptual transitions in illusory rotation, but not the other way around.

Figure 4C, moreover, shows that reported transitions to illusory rotation following microsaccades have the same approximate latency (~500 ms) as reported transitions following actual physical rotation (Fig. 4A, gray band). Thus, perceptual transitions following microsaccades are equivalent in their timing to perceptual transitions following physical rotations.

To quantify our conclusions further, we conducted a ROC analysis (for details, see Materials and Methods, above) to calculate the ability of an ideal observer to predict the type of transition (to rotation vs to stillness) based on microsaccade rates (Fig. 4B,D, insets, green trace). In the main experimental condition (Rotating Snakes Illusion), the ideal observer becomes significantly better than chance (determined by permutation analysis) ~180 ms before the estimated perceptual transitions and ~860 ms before the reported illusory transitions (Fig. 4B, inset, green trace). In the control condition (physical rotation), the ideal observer cannot distinguish between physical transitions to rotation versus to stillness from microsaccade rates alone (Fig. 4D, inset, green trace).

We also found microsaccade magnitude to be correlated to the perception of the illusion. Subjects with larger microsaccades perceived illusory rotation more frequently than those with smaller microsaccades (Fig. 5A). Fixation instability did not correlate significantly with illusory perception, suggesting that slow drift does not trigger the perception of the Rotating Snakes Illusion (Fig. 5B).

Discussion
The Rotating Snakes Illusion produces the perception of intermittent illusory rotation. Previous studies investigating the role of eye movements on the Rotating Snakes Illusion (or the related Peripheral Drift Illusion) did not examine its intermittent characteristics, however, focusing instead in the overall strength of the illusion over a given period of time (Murakami et al., 2006; Beer et al., 2008). Here we determined the onset times of intermittent rotation and stillness periods as experienced by observers, then correlated them to microsaccade and blink production.

Microsaccades generate strong neural transients in the early visual system (Martinez-Conde et al., 2000, 2002, 2009) and counteract visual fading during fixation (Martinez-Conde et al., 2006; Troncoso et al., 2008a). They are also correlated to the perception of certain types of illusory motion (Laubrock et al., 2008; Troncoso et al., 2008b). The present results suggest that
microsaccades (and other transient ocular events such as saccades and blinks) trigger the motion signal underlying the perception of the Rotating Snakes Illusion.

Origin of the motion signal and the role of microsaccades in the perception of the Rotating Snakes Illusion

The inability of motion sensor units to distinguish between temporal changes in contrast and actual motion can result in the perception of illusory motion (Heeger, 1987). Thus, the Rotating Snakes Illusion and the related Peripheral Drift Illusion may result from abrupt changes in the stimulus (i.e., due to sudden onset or displacement of the image, or to transient ocular events such as microsaccades, saccades, and blinks) leading to contrast-driven latency differences (Faubert and Herbert, 1999).

In agreement with this idea, Conway et al. (2005) showed that pairs of stimuli of different contrasts can generate motion signals (i.e., phi movement) in directionally selective neurons of areas V1 and MT.

Our results reveal a tight correlation between microsaccade and blink onsets and the time of the perception of the Rotating Snakes Illusion, supporting the hypothesis that “resetting” or “refreshing” the retinal image triggers the motion signal and the resultant illusory motion (Faubert and Herbert, 1999; Backus and Oruç, 2005; Conway et al., 2005; Kitaoka, 2006). These findings are also consistent with the known physiological and perceptual effects of microsaccades and blinks, such as the generation of neural transients in visual neurons (Gawne and Martin, 2000, 2002; Martinez-Conde et al., 2000, 2002, 2009) and the reversal of visual fading due to adaptation during fixation (Martinez-Conde et al., 2006; Troncoso et al., 2008a; McCamy et al., 2011).

Role of fixation instability in the perception of the Rotating Snakes Illusion

Previous studies did not find a correlation between microsaccade production and the perception of illusory rotation (Murakami et al., 2006; Beer et al., 2008). The discrepancy between past and present results may be due to differences in experimental design. Murakami et al. (2006) found that subjects with more fixation instability experienced the Rotating Snakes Illusion more strongly than subjects with less fixation instability. They proposed that the illusion results from a failure to compensate the image motion induced by fixational eye movements—particularly ocular drift—with the brain interpreting the residual (uncompensated) motion as actual rotation. Beer et al. (2008) similarly found that fixation instability enhanced the perception of a variant of the Peripheral Drift Illusion. These previous studies presented their stimuli for very short time intervals (i.e., ranging from a few seconds to less than a second), however (Murakami et al., 2006; Beer et al., 2008). It follows that the stimulus onsets themselves may have caused neural transients, resulting in perception of the illusion throughout the entire duration of the trial. Moreover, because microsaccades typically occur once or twice per second, they may have been absent in approximately half of the trials in previous experiments (Murakami et al., 2006). Therefore, the correlations between fixation instability and illusion strength reported in prior research cannot be disentangled from unmeasured correlations between the stimulus onsets and illness strength. Accordingly, an alternative interpretation of the previous findings is that fixation instability does not trigger the illusion, but rather modulates its strength once the illusion has been triggered by the stimulus onset itself.

To test the possibility that stimulus onsets could produce illusory percepts lasting for the entire duration of a 1–2 s trial, we compared the duration of the first illusory period after stimulus onset to the duration of subsequent illusory periods in our data (i.e., throughout our 30 s trials). The onset-driven illusory period persisted for an average of 4 s, that is, 2.5 s more than in subsequent illusory periods within the same trial. The difference was significant ($p = 0.005$, $t$ test) and consistent across individual subjects (Table 1), supporting the idea that stimulus onset is a strong contributor to the illusion and suggesting that previous studies could have conflated the effect of the transient onset with that of fixation instability. The present experiments controlled for this potential confound for the first time by analyzing only the data recorded after the first illusory rotation report (i.e., discarding the only illusory period due to the stimulus onset).

To sum up, the current experiments—by presenting long-duration trials and focusing on the illusion’s intermittent features (i.e., alternating periods of perceptual rotation vs stillness)—allowed the optimal identification of the oculomotor trigger mechanisms of the Rotating Snakes Illusion. Thus, the lack of correlation between fixation instability and illusory rotation in our data suggest that drift, by itself, cannot trigger the illusion (though it may modulate the strength of the illusion, once triggered). In support of this idea, Backus and Oruç (2005) found that saccade-like jumps in the Rotating Snakes pattern produced illusory rotation, whereas small jitter of the image was ineffective. Similarly, Tomimatsu et al. (2010) found that fast displacements of the image produced a strong illusory percept, whereas smooth motion resulted in a much weaker illusion.

The present findings are in agreement with the growing literature concerning the effects of microsaccades on illusory perception (Martinez-Conde et al., 2006, 2006; Laubrock et al., 2008; Troncoso et al., 2008b; Hsieh and Tse, 2009), particularly those concerning motion illusions (Laubrock et al., 2008; Troncoso et al., 2008b).

Conclusion

We show a strong quantitative link between microsaccade and blink production and the perception of the Rotating Snakes Illusion during fixation, suggesting that transient ocular events such as microsaccades, saccades, and blinks trigger the perception of illusory rotation. Our results support the theory that visual transients drive the illusion motion signal in the Rotating Snakes Illusion, and suggest that previous reports indicating that either fixation instability or ocular drift trigger the illusion confute the effects of stimulus onsets with the effect of drift.

References


