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Faster processing of moving compared with flashed bars in awake macaque V1 provides a neural correlate of the flash lag illusion

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INTRODUCTION

Moving objects in nature typically follow smooth, predictable trajectories, potentially enabling the brain to minimize or compensate for motion processing delays. The flash lag illusion has kindled much interest among neuroscientists as it is thought to provide a window into the neural mechanisms of localizing moving objects. In this illusion, observers report that a moving bar is located ahead of an aligned flash (Fig. 1A) (Mackay 1958; Nijhawan 1994). Although this phenomenon has been studied extensively at the behavioral level, its underlying neural mechanisms are poorly understood.

In an initial attempt to explain this illusion, it was posited that the brain extrapolates the position of moving stimuli to an extent that compensates its own processing delays (Nijhawan 1994). Since then, many alternative models have been proposed. These diverse models (Fig. 1, B–D), reviewed extensively elsewhere (Eagleman and Sejnowski 2007; Öngen et al. 2004), have pointed toward equally diverse neural mechanisms that range from simple bottom-up explanations such as latency differences to high-level top-down mechanisms such as attention and feedback (Bachmann and Pöder 2001; Baldo and Klein 1995; Brenner and Smeets 2000; Eagleman and Sejnowski 2007; Krekelberg and Lappe 2000; Patel et al. 2000; Purushothaman et al. 1998; Sheth et al. 2000; Whitney and Murakami 1998). For example, the differential latency model (Purushothaman et al. 1998; Whitney and Murakami 1998) maintains that moving stimuli are processed faster compared with flashed stimuli, leading to the perception of flashes temporally coinciding with a moving bar further along its trajectory. Alternatively, the motion-biasing model (Eagleman and Sejnowski 2007; Rao et al. 2001) argues that motion signals after the detection of a flash event affect position representation and judgments, such that observers report a misalignment between a flash and a moving stimulus. There has also been a recent attempt to subsume all these models into a single mechanism.
Most models of the flash lag illusion are formulated at the psychophysical level. At this level of abstraction, the three most prominent models (Fig. 1, B–D) of the illusion differ in their prediction for the relative latencies of flashed and moving stimuli (Fig. 2). In this context, the latency or “representation delay” refers to the time interval between stimulus appearance at a particular location in the physical world and the emergence of neural activity corresponding to the reported perception of the stimulus location. The spatial extrapolation model, as it posits full compensation for neural delays (Nijhawan 1994), would predict zero representation delay for motion; the differential latency model predicts shorter latency for motion (Patel et al. 2000; Purushothaman et al. 1998; Whitney and Murakami 1998); and the postdiction model assumes equal latency for flash and moving stimuli (Eagleman and Sejnowski 2007; Rao et al. 2001). The models do not specify in which part of the brain one would observe such predicted latency differences of stimulus representations. Here, to systematically investigate the neural mechanisms of the flash lag illusion and to test predictions of the psychophysical models, we measured the latencies or representation delays of flashed and moving stimuli in primary visual cortex (V1) of awake, fixating macaques. This allows us to determine the contribution of early visual processing toward the illusion. It should be noted that, at the level of V1, the term latency or representation delay refers to the time interval between stimulus appearance at a particular location in the physical world and the time of stimulus-evoked activity in V1 at which a decoder or a downstream processing region can obtain the best estimate of the stimulus location (see MATERIALS AND METHODS).

The few physiological studies that have explored the neural mechanisms of the illusion found a shorter latency for motion signals compared with flashes in the rabbit and salamander retina (Berry et al. 1999), cat lateral geniculate nucleus (LGN) (Orban et al. 1985), and cat V1 (Jancke et al. 2004b), providing evidence for a bottom-up latency difference between flashes and moving stimuli. Although these studies provide valuable hints at plausible neural mechanisms of the flash lag illusion,
they were done either in vitro or in anesthetized animals, and it is unknown whether the animals used in these studies actually perceive the illusion.

We previously showed that, similar to humans, macaque monkeys perceive the flash lag illusion (Subramanian et al. 2013). Hence, we performed the physiological experiments in awake macaque monkeys, which allowed us to directly test the predictions of different models of the illusion at the level of V1 neural representation of flash and moving stimuli (Fig. 2). Specifically, we estimated the latency of the stimulus representations by two different methods, one based on multiunit response peak times and the other based on probabilistic decoding of simultaneously recorded single-unit and multiunit population activity. Crucially, we measured the dependence of latency on different stimulus parameters—speed, luminance, and direction of motion—to test whether the resulting changes in neural responses accounted for the corresponding changes in perception. Under all these manipulations, neural latency differences between flash and motion in V1 explained a large part of the psychophysically measured perceived spatial offsets. Thus our results show that even at the very first cortical visual information processing stage a neural correlate of the illusion can be observed, providing mechanistic constraints on the models of the flash lag illusion.

MATERIALS AND METHODS

Subjects. Four male macaque (Macaca mulatta) monkeys (A, CH, CL, and L) weighing 8, 9, 12, and 9.5 kg, respectively, and aged 10, 8, 8, and 8 yr, respectively, were used in the physiological experiments. A cranial head post and a scleral search coil were implanted in each monkey with standard aseptic surgical procedures. All animal procedures were approved by the Institutional Animal Care and Use Committee of Baylor College of Medicine and followed National Institutes of Health regulations. Two of the authors [M. Subramanian (subject MS) and S. S. Patel (subject SP)] participated in psychophysical experiments following procedures approved by the Institutional Review Board of Baylor College of Medicine.

Electrophysiological recording and data processing. We used chronically implanted tetrode arrays for recording neural activity from monkeys A, CL, and CH as described previously (Ecker et al. 2010; Tolias et al. 2007). Briefly, in each monkey we chronically implanted arrays of 24 tetrodes on the left hemisphere over the operculum in area V1. The tetrodes were custom built from nichrome or platinum-iridium wires. We implanted a 96-electrode microelectrode array (“Utah” array; Blackrock Microsystems, Salt Lake City, UT) over area V1 on the right hemisphere in monkey L. For both tetrode arrays and the Utah array, the neural signals were preamplified at the head stage by unity gain preamplifiers (HS-27; Neuralynx, Bozeman, MT). These signals were then digitized with 24-bit analog data acquisition cards with 30 dB onboard gain (PXI-4498; National Instruments, Austin, TX) and sampled at 32 kHz. Broadband signals (0.5 Hz to 16 kHz) were continuously recorded with custom-built LabVIEW software for the duration of the experiment. For tetrode array data, the spike detection and spike sorting methods have been described previously (Ecker et al. 2014; Tolias et al. 2007). For the Utah array, spikes were detected from individual electrodes according to the same procedure. In this study, the term “multiunit” refers to the set of all the spikes detected from a single tetrode or a single electrode (Utah array).
Behavioral task. Visual stimuli were presented in a dark room with dedicated graphics workstations using Psychophysics Toolbox 3 (Brainard 1997; Kleiner et al. 2007; Pelli 1997). For all experiments with monkeys A, CH, and CL we presented stimuli on CRT monitors (model: Sgi C220 Flat Diamondtron; display size: 22° × 16° from a distance of 100 cm; resolution: 1,600 × 1,200 pixels; refresh rate: 100 Hz). For monkey L, we presented stimuli on an LCD monitor (Samsung model S23A950D; refresh rate: 120 Hz; monitor resolution: 1,920 × 1,080 pixels, subtending visual angles of 29° × 16° from a viewing distance of 100 cm). We gamma-corrected the monitors to achieve a linear luminance response profile. The monitor background luminance was 6.1 cd/m² (monkeys CL and A), 9.5 cd/m² (monkey CH), or 0.04 cd/m² (monkey L). The monkeys sat in a custom primate chair at 100 or 107 cm from the stimulus display monitor. Eye positions were continuously monitored online with a scleral search coil for monkeys A, CH, and CL and with a custom-built video tracker (frame rate: 250 Hz) for monkey L. Eye position signals were also saved for off-line analysis. Each trial (see Fig. 4A) began with a brief sound that instructed the monkeys to start fixating at a red dot (0.12–0.14°) within a circular window of radius 0.5–0.6° of visual angle. After the monkeys fixated for 300 ms, we presented different visual stimuli. The monkeys fixated for an additional 300 ms after the stimulus offset. For successfully completing the trials, the monkeys received a juice or water reward. The next trial began after an intertrial time of 1,500 ms.

Receptive field mapping. We mapped the spatiotemporal receptive fields with a white noise random dot stimulus. On a gray background, we presented black and white squares (0.11–0.14° side) on a rectangular grid covering the receptive field of all recorded neurons. The squares were presented one at a time for three video frames (25–30 ms) in a pseudorandom sequence for 1,200–2,000 ms. The sequence consisted of many iterations, in each of which every grid location was visited exactly once in a random order, thus balancing the number of times each location was visited over the course of the experiment. The monkeys performed 242 trials (standard deviation (SD) 56) in a session that lasted for ~20 min. Since primate V1 contains many complex cells and we were interested primarily in the location of the receptive fields, we performed reverse correlation ignoring the sign of the stimulus (i.e., both black and white were treated as positive). We assessed the quality of the receptive field estimation by the following heuristic method. We first averaged the receptive field maps obtained at lags ranging from 40 to 100 ms, resulting in a single spatial kernel for each multiunit. We fitted the spatial kernel with a two-dimensional Gaussian and computed the percentage of variance explained (across pixels) by the model. For all analyses in this study, we included multiunits for which the model explained >75% of the variance. From the model fitting, we also extracted receptive field centers and outlines. For illustration we outlined receptive fields by the elliptical contour at two SDs from the center.

Speed manipulation experiment. Monkeys A, CH, and CL were used in this experiment. Moving and flashed vertical bars of identical luminance and size (0.28 × 1.7°) were used as visual stimuli. The bar luminance was either 23 cd/m² (monkeys A and CL) or 37 cd/m² (monkey CH). We defined a stimulus presentation center for each monkey as the average of the receptive field centers (ARFC of the neurons we recorded from; the mean eccentricity of this location was 1.5° (SD 0.11) (azimuth: 0.87° (SD 0.3), elevation: 1.2° (SD 0.3)). In each stimulus period, only a flash or a moving bar was presented. We presented flashes for one video frame (10 ms). Since we recorded from many neurons simultaneously, to stimulate all the recorded neurons we presented flashes at five to seven locations around the ARFC (see Fig. 4B). These locations were abutting each other without any overlap. The trajectory length of the moving bar was 4.6° or 5.4°. The midpoint of the moving bar’s trajectory was at the ARFC. The moving bar translated horizontally from left to right or from right to left at one of three speeds: 7, 14, or 28/°s (range: 6.9–7.4, 13.8–14.7, and 27.5–29.5/°s, respectively). All stimulus conditions were presented with equal probability. In each trial (Fig. 4A), we chose more than one stimulus condition randomly (2 flashes and 1 moving stimulus, for example) and presented them one after the other with an interstimulus period of 300 ms; this allowed us to use the monkeys’ fixating period efficiently and present multiple stimulus conditions within each trial. During the stimulus period of ~1,800 ms, we presented 4 stimuli (SD 1). In a session, we repeated each stimulus condition 426 times (SD 216). The monkeys performed 1,597 trials (SD 718) per session. Each session lasted for 3 h (SD 1).

Luminance manipulation experiment. Monkey L was used in this experiment. The stimulus presentation followed the same overall design as the speed manipulation experiment (see above), with the following exceptions. The size of the bar was 0.15 × 1.8°. Moving and flashed bars with luminance values of 0.24, 0.82, 9.4, and 48 cd/m² were presented in each session. Flashes were presented at one of nine abutting locations with the ARFC at an eccentricity of 0.92° (SD 0.07) (azimuth: −0.46° (SD 0) elevation: 0.79° (SD 0.08)). The trajectory length of the moving bar was 8.7°. The moving bar translated horizontally from left to right or from right to left at 18/°s. In the stimulus period of each trial, we presented 5 stimuli (SD 1). Each stimulus period was repeated 120 times (SD 46). The monkey performed 1,128 trials (SD 432) per session, with each session lasting 2 h (SD 1). It should be noted that to fit all luminance conditions within the recording duration, we did not test multiple speeds. Instead we chose a speed (18/°s) that was intermediate between speeds of 7 and 28/°s that were used in the speed manipulation experiment. We also reduced the width of the bar to roughly half (0.15°) that of the bar used in the speed manipulation experiment (0.28°) so that when the bar moves the footprints of the bars in the trajectory are contiguous without overlap or leaving a gap between adjacent instantaneous positions. The flash duration (8.3 ms) is also shorter than that used for speed manipulation (10 ms) because we had to use to test very low luminance levels we had to set the background luminance to the lowest possible value; at that setting (but not at the background used in the speed manipulation experiment) when the bar moved on the CRT monitor, it left behind a trail of phosphorescence that was obvious to a human observer. Such trailing luminance was not observed on the LCD monitor.

Control experiment. Monkeys A and CL were used in this experiment. Stimuli were presented as outlined for the speed manipulation experiment. However, in addition to presenting flashed and moving bars separately as above, we also interleaved additional stimulus conditions in which we presented the flash and moving bars together in two arrangements, A1 and A2 (Fig. 1A). In arrangement A1, we presented a flash inside the receptive fields and the moving bar below the flash but outside the receptive fields. To mimic the psychophysical experiment of the flash lag illusion, in arrangement A1 when the instantaneous position of the moving bar hit the azimuth of the ARFC, a flash was presented at one of five to seven horizontal spatial offsets (0°, ±0.27°, ±0.55°, ±0.82°). We assigned a negative sign to the offsets if the flash appeared ahead of the moving bar along the motion direction and a positive sign if the flash appeared behind the moving bar. In arrangement A2, the vertical positions of the flash and moving bar in arrangement A1 were interchanged. The moving bar translated at a speed of 14/°s. The vertical center-to-center distance between the flash and the moving bar was 2.1°. With the bar height being 1.7°, the edge-to-edge gap between the bar and the flash was 0.4°. In each trial, we presented 3 (SD 1) stimulus conditions. Each stimulus condition was repeated 159 times (SD 81). The monkeys completed 1,930 trials (SD 742) per session, with each session lasting 3 h (SD 1).

Electrophysiological data set. For the entire study, we recorded neural data from a total of 1,457 multiunits (monkey A: 288, CH: 191, CL: 306, L: 672) over 62 sessions (A: 12, CH: 23, CL: 20, L: 7) in an average period of 6 wk from each monkey (A: 4, CH: 12, CL: 6, L: 2). For the flash, relative to the prestimulus fixation period, a majority

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[1,038 (71%); A: 247, CH: 180, CL: 276, L: 335] of the multiunits showed significantly enhanced responses measured over a window of 30–130 ms after the flash onset. A minority [44 (3%); A: 2, CH: 11, CL: 20, L: 11] of the multiunits showed flash-evoked suppression. For analyses, we included a subset of the multiunits [915 (63%); A: 237, CH: 166, CL: 256, L: 256] that showed enhanced flash-evoked responses and passed the receptive field-based selection criterion [955 (66%); A: 247, CH: 176, CL: 271, L: 261; see Receptive field mapping]. After the above selections, one multiunit from monkey A was excluded from the analyses in Figs. 7 and 10 because its receptive field center was outside the flashed region. For the speed manipulation experiment, a total of 163 (A: 57, CH: 58, CL: 50) single units were isolated, out of which 44% (total: 71; A: 32, CH: 12, CL: 27) met the selection criteria described above. For population decoding we chose all the single units from monkey CL since it had the most well-isolated units [median contamination measure (Tolias et al. 2007) (interquartile range): CL: 0.039 (0.005, 0.086), CH: 0.076 (0.048, 0.117), A: 0.092 (0.015, 0.142)]. Neural data associated with this study can be found at https://datadryad.org/resource/doi:10.5061/dryad.md2n929.

Response peak delay as neural representation delays for flash and moving stimuli. For the moving stimuli, assuming a receptive field-based labeled-line code for position in V1, the latency of peak activity of a neuron closely approximates the representation delay. This is because whenever there is a bar moving in the visual field, a population activity hill representing some moving bar position is simultaneously present in V1 (Fig. 2), except during the motion onset and offset. We assume that any subsequent visual area decoding moving bar position based on V1 activity would assign the instantaneous position of the bar center to the position encoded by the neurons whose activities maximally contribute to the peak of the hill. This would imply that the time at which a given neuron fires maximally is also the time at which the moving population hill activity is centered over this neuron’s topographic location in V1. Under this reasoning, the response peak latency would correspond to the latency of the V1 representation of the moving bar’s instantaneous position. For the flash, the situation is different because when a flash is presented in the visual field a population activity hill starts to develop only after a delay. The hill then rises and falls over time without any change in the position of the peak of the hill. It is currently unknown at what point in time the activity hill fully represents the flash location. To be consistent with the method of latency computation of motion, we chose to compute peak response latency for flash as well.

Estimation of flash response peak latency. For each flash condition, we first aligned the spike times of a given stimulus presentation to the flash onset time. We then computed mean firing rates across all stimulus presentations of a given condition after binning the spikes at half the monitor refresh period (4.2 or 5 ms). In each session, multiple flashes were presented, covering the receptive field of a given multiunit. We sought to find the mean firing rate response profile to a flash that was horizontally aligned with the center of the receptive field. However, there might not be any flash that was presented perfectly over the receptive field center since we did not optimize the flash locations for any particular neuron. In such cases, the mean firing rate profile that corresponds to a flash at the receptive field center was obtained by linearly interpolating the mean firing rate profiles of the flash locations left and right of the receptive field center. The mean firing rate response starting 150 ms before and ending 300 ms after the flash onset was then normalized (z scored) to have zero mean and unit variance. After z scoring, the responses of all multiunits under a given condition were averaged and smoothed with a Gaussian kernel with a SD of 10 ms. Peak response latencies were then computed from these averages. The responses of individual single units and multiunits to flashed and moving bars were sometimes multimodal. Since we had a much larger multiunit data set compared with single units, we chose to extract the latencies from responses obtained across multiunits. This procedure turned out to be more robust than extracting latency for each unit (for a description of how we estimated confidence intervals on the latencies, see Statistical analysis).

Estimation of motion response peak latency. For each motion condition, we aligned the spike times of a given presentation to the time at which the moving bar hit the center of the receptive field (i.e., the response time is set to 0 when the moving bar’s instantaneous position matched the receptive field center). Since the moving bar occupied discrete positions along the trajectory that did not necessarily coincide with the receptive field center, we linearly interpolated the trajectory time points to obtain the time at which the trajectory crossed the receptive field center. We then computed mean firing rate across all presentations of a given condition after binning the spikes at half the monitor refresh period (4.2 or 5 ms). The mean firing rate response starting 150 ms before and ending 300 ms after the zero time point was then normalized (z scored) to have zero mean and unit variance. This normalized response was then averaged across multiunits. After this step, we followed the same procedure as for the flash responses described above and computed response peak latencies for each stimulus condition. The latencies were then averaged across the two motion directions.

Latency estimation in control experiment. In the control experiment, we computed response peak latencies for flashes from arrangement A1 and for moving bars from arrangement A2 (see Control experiment). To compute flash response latency for a given spatial offset, we first selected multiunits whose receptive field centers were within the spatial extent of the presented flash. Response peak latency was then extracted from this set of multiunits as described in Estimation of flash response peak latency. To compute the motion response latency for any spatial offset, we first selected multiunits whose receptive field centers were within the spatial extent of the moving bar when it hit the ARFC. Since the flashes were presented at different horizontal locations when the moving bar hit the ARFC, the same set of multiunits were used for extracting latencies under different spatial offsets. Motion response peak latencies were then computed as described in Estimation of motion response peak latency. Note that we chose to include a spatial offset for analysis only if there were >10 multiunits for that condition. With this criterion, only the three spatial offsets around the ARFC qualified.

Statistical analysis. All the statistical analyses on the neural data were done by bootstrapping (Efron and Tibshirani 1994). From the response (averaged over multiunits) peak latencies of flash and moving bar under various conditions, we computed the following test statistics: latency difference between flash and moving bar (Fig. 7B, Figs. 12 and 13, F); slope of the trend in the latencies (Fig. 7B, Figs. 12 and 13, F); latency differences and perceived spatial offset equivalents when changing speed (Fig. 7, C and D, Fig. 10, E and F, Figs. 12 and 13, G and H) and luminance (Fig. 9, C, F, and G, Fig. 10, H and I, Fig. 14, G–I); and for the control experiments latency differences across multiple spatial offsets (Fig. 11B) and latency differences for stimuli presented in isolation vs. in combined condition (flash and moving bar presented together) (Fig. 11C). To obtain significance levels and confidence intervals on these test statistics, we repeated the entire procedure that generated a test statistic 2,000 times, each time with a different random set of multiunits obtained by resampling with replacement. Since the electrodes were implanted chronically, individual recordings from different days may not represent independent samples. To ensure that we used only independent samples for bootstrapping, we removed zero sampled electrode identities and included all units obtained from sampled electrodes. This procedure estimates the unit-to-unit variability without being confounded by dependent samples due to chronic recordings. From this bootstrap distribution, we computed the 95% percentile confidence intervals, which are reported as error bars. We defined the significance level (P value) as \( P = 2 \min(q, 1 - q) \), where \( q \) is the percentile of zero under the bootstrap distribution (this analysis assumes that the bootstrap distribution is an appropriate measure of the variability under the null hypothesis).
Human psychophysics task. Two human subjects (MS and SP) performed the standard flash lag psychophysical experiment as described previously (Subramaniyan et al. 2013). The subjects sat in a dark room with their heads stabilized by a chin rest. After the subjects dark-adapted their eyes for 5 min, the stimulus presentation began. The subjects were simply instructed to stay fixated at the fixation spot during stimulus presentation; their eye movements were not tracked. In any given trial, we presented a flash below another bar that moved from left to right; the gap between the bottom edge of the moving bar and the top edge of the flashed bar was 0.3°, and the bars had identical luminance. We used seven different horizontal offsets between the flash and moving bar centers. The offset values ranged from around -6.3° to 2.4° in steps of around 1.5°. We used a constant flash location and created the spatial offsets by choosing the time of flash relative to the instantaneous position of the moving bar. To be comparable to the physiological experiments with monkey L, we made sure that at zero spatial offset the average position of the flash and moving bar centers matched the ARFC used for monkey L. In each session, we randomly interleaved four bar luminance values. These luminance values, bar dimensions, monitor background luminance, and speed of the moving bar were identical to those used in the luminance modulation experiment with monkey L, although here a longer motion trajectory of 18° was used. Using a keyboard, the subjects reported whether the leading edge of the moving bar was on the right or left side of the flash at the moment the flash appeared. The subjects completed a total of seven sessions (MS: 5, SP: 2). In most sessions, we presented a total of 28 stimulus conditions (7 offsets × 4 luminance values × 1 motion direction × 1 speed). Each condition was repeated 20 times, giving ~560 trials per session. Each session lasted for an average 23 min.

Estimation of perceived spatial offset. To quantify the perceived spatial offset, we first converted the subjects’ responses into a probability of reporting that the moving bar was ahead of the flash. Then we fitted a logistic function to these probabilities as a function of spatial offsets, using psignifik3.0 toolbox (Fründ et al. 2011; Wichmann and Hill 2001a, 2001b). In the toolbox, we chose the constrained maximum likelihood method for parameter estimation and parametric bootstrapping for estimation of confidence intervals for parameters. We constrained the upper and lower asymptotes of the psychometric function to be equal with the prior distribution being a uniform distribution on the interval [0, 0.1]. We defined the perceived spatial offset as the point of subjective equality, that is, the veridical spatial offset at which subjects reported that the moving bar was ahead or behind the flashed bar with equal probability. To examine how the perceived spatial offset changed with luminance, we pooled the responses across sessions for each bar luminance before fitting the psychometric function. To perform statistical tests, however, we fitted the psychometric function for each session separately and computed perceived spatial offset.

Statistical analysis of psychophysical data. For all statistical tests on psychophysical data, linear mixed models were constructed in the statistical software PASW-18, with the following common settings: subjects were treated as random effects and perceived spatial offset as dependent variable. Specifically, the slope of the trend of the perceived spatial offset as a function of bar luminance (Fig. 9E) was tested for significance using the bar luminance as a covariate with the session start times set to indicate repeated measures. To test the effect of motion condition (foveopetal vs. foveofugal) and speed on the perceived lag (Fig. 10, B and D), speed was used as a covariate and motion condition as a factor, with the combination of motion condition and session start times set to indicate repeated measures.

Probabilistic population decoding. The decoding method used here was chosen for its simplicity and its suitability for our experimental conditions abstracting away from neuronal implementation level details. Our goal was to decode the stimulus position presented to the animal from the single-unit or multunit population activity based on the framework of probabilistic population coding (Dayan and Abbott 2005; Ma et al. 2006; Zhang et al. 1998). We took advantage of the fact that the motion stimulus we used was essentially a sequence of flashes. Hence, to decode the moving bar location, we first modeled the spatial encoding by measuring the population activity for the different flashed locations of the bar. Then, when the moving bar was presented, we decoded its instantaneous position by identifying the bar stimulus that was most likely, given the population activity at that instant. It should be noted that in our experiments only part of the motion trajectory overlaps with the space covered by the flashes. Since the spatial encoding is based on flash responses, we restricted the motion decoding to the region of the trajectory overlapping the flash locations. The decoding method is formalized as follows.

A flashed stimulus (Fig. 3A) evokes neural activity that extends over time, outlasting the presence of the stimulus (~10 ms) on the monitor (Fig. 3B). The poststimulus period can be split into a sequence of contiguous time bins (of width Δt). We assume that, conditioned on the stimulus, the spiking responses are independent across both time and neurons. That is, activity (R) in any given time bin depends only on the stimulus location (S) and the elapsed time since stimulus onset (e). Under this assumption, the neurons spike according to independent inhomogeneous Poisson distribution, with a time- and neuron-dependent mean spike count parameter (λ(S,e)) (Fig. 3C). This produces the following probability distribution for neural activity (R) in a single time bin of width Δt:

![Fig. 3. Probabilistic population decoding. A: outlines of receptive fields (gray circles) of simultaneously recorded multiunits (n = 21) from a single representative session from monkey CL. Gray rectangles show the outlines of different flashes (labeled as F1, 2, 3, 4, 5) presented one at a time. B: single-trial raster plot of spiking responses (dark vertical bars) of all multiunits in A to a flash (F2). The spike counts within the thin gray vertical box (of width Δt = 10 ms) forms the activity vector (R) used in the decoding procedure. C: mean spike count across trials in 10-ms consecutive time bins for all multiunits under each flash condition indicated at top. An example value for λ(S,e) parameter (at bottom left). The y-axis scale bar for the traces is shown at bottom right. D: single-trial raster plot of spiking responses (dark vertical bars) of all multiunits in A to a bar moving from left to right at a speed of 7°/s. Thin gray vertical box as in B. Black horizontal bar on x-axis marks the time period the moving bar spent within the flashed region shown in A. E: graphical model of population activity. The population neural response at a given time bin (R) is governed by the stimulus (S) and the time elapsed (e) since stimulus onset.](https://www.jn.org/content/jn/110/7/2435/F3.large.jpg)
where $S$ is stimulus (bar) at one of $M$ possible locations (for example, see gray rectangles in Fig. 3A), $e$ is time elapsed since stimulus onset, $N$ is the number of neurons simultaneously recorded, $n_i$ is spike count in a given time bin of width $\Delta t$ for neuron $i$, $\lambda_i(S,e)$ is mean spike count of neuron $i$ in a time bin of width $\Delta t$ after a delay of $e$ from stimulus $S$ onset at one of the $M$ possible locations, and $\mathbf{R}$ is population activity [spike count vector $(n_1, n_2, \ldots, n_N)$] in a single time bin.

Note that, for the flash-evoked neural activity in any given time bin, the experimenter knows which flash stimulus caused the activity and how much time has elapsed since the stimulus onset (Fig. 3B). However, these two parameters are unknown from the brain’s perspective. In the case of the moving stimulus for which a priori we do not know the response latency, even the experimenter cannot know which stimulus location causes neural activity in a given time bin (Fig. 3D). This is due to the moving bar changing its location in every time bin, leading to essentially multiple stimulus locations driving the neural activity in different time bins. As the experimenter cannot know which stimulus location caused the activity, he/she also cannot know how much time has elapsed since the onset of the stimulus (at a given location) driving the activity. For these reasons, in our decoding of flashed and moving stimuli we treat the stimulus location and time elapsed since stimulus onset/arrival at a given location as random variables that follow a uniform distribution with flat priors.

Note that the response at a single time bin for a moving stimulus is likely driven by multiple stimulus (bar) locations (spatiotemporal integration). However, to decode this activity, we are using an encoding model in which population activity at each time arises from single stimulus (flash) locations. Hence, in our decoding procedure we are only approximating the spatiotemporal integration involved in generating population activity during motion. This leads to a graphical model (Fig. 3E) that, in combination with Eq. 1, can be used to decode the stimulus position from the neural activity. Decoding this way in small time bins (~10 ms) implies that a rate code is used by the brain for computing stimulus position. To compute the probability of a stimulus given the population activity in a single time bin, we first derive a joint distribution based on the model in Fig. 3E.

\[
p(S, e, \mathbf{R}) = p(S)p(e)p(R|S, e)
\]

We assumed that $S$ and $e$ follow a uniform distribution (range of $S$: horizontal extent of flashed region, range of $e$: 10 to ~175 ms) and hence $p(S)$ and $p(e)$ are constants (flat priors). We can then marginalize the above joint distribution over the elapsed time $e$ to compute the probability of a stimulus location given the population activity $\mathbf{R}$ in any arbitrary time bin:

\[
p(S, \mathbf{R}) = \frac{p(S, e, \mathbf{R})}{p(\mathbf{R})} = \frac{p(S)p(e)p(R|S, e)}{p(\mathbf{R})}
\]

\[
p(S|\mathbf{R}) = \sum_e p(S, e|\mathbf{R}) = p(S)\sum_e p(e|\mathbf{R})\frac{p(R|S, e)}{p(R)}
\]

As $p(S)$ and $p(e)$ are assumed to be constants for all values of $S$ and $e$, respectively, they can be absorbed along with $p(R)$ into the normalization constant $Z(\mathbf{R})$, simplifying Eq. 4 as

\[
p(S|\mathbf{R}) = \frac{1}{Z(\mathbf{R})}\sum_e p(S, e|\mathbf{R})
\]

where $Z(\mathbf{R})$ can be computed with the following normalization constraint:

\[
\sum_{j=1}^{M} p(S|\mathbf{R}) = 1
\]

where the subscript $j$ indexes the possible positions of the bar stimulus in visual space. Since the decoding was restricted to the space occupied by the flashes, the above constraint (Eq. 6) is justified, as the decoded position should be within the flashed space. By using the above constraint, we avoided computing $p(R)$ explicitly, as done by previous studies in similar decoding problems (Sanger 1996; Zhang et al. 1998). Note that for monkeys $A$ and $CH$, although seven flashes were presented in the task, we only included the central five flashes in the analysis, as the flashes at the periphery did not have sufficient receptive field coverage.

Decoding was done trial by trial for each recording session with neurons recorded simultaneously. The same number of trials was used for all stimulus conditions within a session. In each trial, from stimulus onset, we stepped forward in small contiguous (nonoverlapping) time bins ($\Delta t$ = monitor refresh period, 8.3 ms for monkey $L$, 10 ms for others) and computed the posterior probability of each of the possible stimulus positions ($M$ in total) given the population activity at that time bin. Hence, at every time instant, for a given test stimulus (flash or moving bar), we get an $M$-element vector of posterior probabilities that sum up to 1. For all stimulus conditions, the posterior probability was assigned to the end of the time bins. For example, the probability computed in the [0, 10) ms time bin was assigned to $t = 10$. This ensures that probability is causally related to the population activity. Also note that for the speed of 7/s, only every fifth moving bar center matched flash centers (Fig. 5A). Hence when computing $\lambda_i(S,e)$, we interpolated the mean firing rate from the flash centers to all positions ($M$ in total) that the moving bar center occupied (white dots in Figs. 12–14, D). For simplicity, the same interpolated $\lambda_i(S,e)$ was used for all speeds. For the luminance modulation experiment, a similar interpolation procedure was done and the decoding of bar stimuli of a given luminance was based on encoding obtained from responses of flashes of matching luminance.

For the marginalization in Eq. 4, we chose a time window that covers the flash-evoked responses of all recorded neurons for all monkeys. Based on visual inspection of the neural responses, this window was set to 10–175 ms for monkey $L$ (to allow for longer response latencies at low-luminance conditions, see Fig. 8) and 10–150 ms for all other monkeys (see Fig. 6). The results and conclusions based on decoding are not sensitive to the exact values of the above time windows. For example, shortening the above windows to 20–130 ms and 20–100 ms, respectively, does not change the results and conclusions presented. However, including some time bins in which the population activity is at the baseline level minimizes the “edge effect” where the decoder, when decoding baseline-level activity, assigns a relatively higher probability to stimulus locations at the periphery (“edge”) of the flashed region (see the decoding in the 0–50 ms window in Fig. 14B). This effect arises because the edge regions often have relatively poor receptive field coverage in our data set (see first and last gray rectangles in Fig. 14A). When a bar stimulus is presented here, it evokes a population response that is similar to the baseline activity (see Fig. 3C, stimulus F5). Consider a decoder that does not include any baseline-level time bins in the marginalization time window in Eq. 5. When this decoder decodes baseline-level activity from any stimulus condition, it will assign a higher posterior probability to the edge regions (edge effect), as the bar stimuli at the interior locations are unlikely to evoke such poor baseline-level activity. Instead, including some baseline bins (e.g., bins with $e = 10–30$ ms) in the marginalization time window minimizes this effect. This is because these bins contribute appreciably to the likelihood term inside the summation operation in Eq. 5. Hence, for a given S, the total likelihood summed over $e$ $[\sum_e p(S|\mathbf{R}, e)]$ will be higher than when not including these bins. Moreover, as the baseline activity is similar for all bar locations ($S$), the large likelihood
contribution will also be similar for all $S$. The result of such an overall increase in the total likelihood is that, after normalization in Eq. 5 [division by $\mathcal{Z}(\mathbf{R})$], the posterior probabilities of the $M$ locations become similar when there is no stimulus-evoked activity, thereby minimizing the edge effect.

Cross-validation. The decoding was done on individual trials. It should be noted that in the above model we learned the spatial encoding from the population response to flashes. Hence, when we decoded flash stimuli, to prevent overfitting we kept aside a given trial for testing and used the remaining trials to train the model (i.e., compute the $\lambda$’s). This was then repeated for all available trials. For decoding the motion stimulus, however, the separation into training and testing trials was unnecessary because the trials used for training (flash trials) were different from the trials in which testing was done (motion trials).

Computing latencies from probabilistic decoding. The posterior probabilities of bar locations computed as described above were first averaged across trials and then across sessions for a given monkey. For estimating the decoding latency for flashes, we first averaged the probability values corresponding to positions within the horizontal spatial extent (see white horizontal bars in Figs. 12–14, $B$) of a flash stimulus. This was repeated for each flash condition, and then the probabilities were averaged across the flashes (Figs. 12–14, $C$) and smoothed with a Gaussian kernel with a standard deviation of ~5 ms. We then computed the latency of the peak of this averaged posterior probability as a measure of the latency of flash stimulus representation (Figs. 12–14, $F$). For motion latency, first we aligned (centered) the probability vector computed at each time bin to the moving bar’s instantaneous position. We included time bins starting from the time the moving bar entered the flashed zone until ~120 ms or less after the bar exited the flashed region of space to account for latency of responses. The aligned vectors were then averaged (Fig. 12–14, $E$) and smoothed with a Gaussian kernel with a standard deviation of 0.1°; the distance between the peak of the posterior probability and the origin was taken as the spatial lag of the moving bar representation. The time delay (latency) corresponding to this spatial lag was then calculated by dividing the spatial lag by the speed of the moving bar. To obtain significance levels and confidence intervals on test statistics based on latencies computed as above, we repeated the entire decoding procedure that generated a test statistic 2,000 times, each time with a different random set of single units or multiunits within each session, by resampling with replacement.

RESULTS

We assessed whether there are differences in the representation delays (latencies) of moving and flashed stimuli and whether this could account for the perceived spatial misalignment (offset) in the flash lag illusion. To this end, we recorded neural activity from V1 while the monkeys were shown either a flashed or a moving bar in a passive fixation task (Fig. 4A; see Fig. 4. Fixation task and stimuli. $A$: monkeys fixated their gaze at a red circular dot (shown here in black) at the center of the monitor within a fixation radius of 0.6°. After they maintained fixation for 300 ms, a single randomly chosen bright flash or moving stimulus was presented in a gray or dark background. The stimulus offset was followed by a 300-ms period in which no stimulus was presented except for the fixation spot. Then a randomly chosen flash or moving bar was presented again. With the monkeys maintaining fixation, this cycle continued until at most 1,800 ms elapsed, after which they obtained a squirt of juice as reward. The next trial started after an intertrial period of 1,500 ms. $B$: a flash (black bar) was presented at one of 5–7 adjoining locations (gray rectangles) tiling the receptive fields (gray circles) of all recorded neurons. $C$: the moving bar (black bar) had the same size as the flash and moved from left to right or from right to left. Dots denote the positions of the bar center along the entire trajectory as the bar moved from left to right at a speed of 7°/s. In $B$ and $C$, the coordinate (0° azimuth, 0° elevation) marks the center of fixation, and the bars and receptive field outlines are drawn to scale. Note that the gray circles show the outlines of only a subset of the recorded neurons. $D$: markers show median of receptive field centers of monkeys CH, CL, A, and L. Horizontal and vertical error bars indicate 95% percentile limits of azimuth and elevation, respectively, of receptive field centers. Isoeccentricity lines are shown in gray.
We performed two experiments: In the first, we varied the direction of motion and speed of the moving bar (7, 14, or 28°/s) while keeping the moving and flash stimuli at a fixed luminance (23 or 36 cd/m²). In the second, we kept the speed constant and manipulated the luminance of the flash and the moving bar. In both cases, we measured the effect of the manipulation on the latency difference between the moving and the flashed bar and compared it to the psychophysical results from monkeys (Subramaniyan et al. 2013) and humans (Murakami 2001; Purushothaman et al. 1998; Subramaniyan et al. 2013; Whitney et al. 2000; Wojtach et al. 2008).

For experiment 1, we recorded from 523 multiunits in three animals with chronically implanted tetrode arrays. For experiment 2, we collected responses of 256 multiunits in one animal, using a 96-channel Utah array. After an initial receptive field mapping session, the main task began. We presented bright bars on a gray (experiment 1) or dark (experiment 2) background. In each trial either a flash or a moving bar was shown. Since we recorded from many neurons simultaneously, the flash locations were not optimized for any particular neuron. Instead, in each recording session, flashes were shown at five to seven fixed locations covering the receptive fields of all the recorded neurons (Fig. 4B). The moving bar swept across the receptive fields horizontally at a constant speed from left to right or from right to left with equal probability (Fig. 4C). For experiments 1 and 2, the receptive fields of units were in the right and left hemisphere, respectively (Fig. 4D). To test the predictions of different models of flash lag illusion, we estimated the stimulus representation delays of flashed and moving bars in V1 with two different approaches. The first method was based on the neuronal responses recorded on individual recording sites, and the second was based on decoding simultaneously recorded single-unit and multunit population activity. Specifically, we tested the dependence of the latency difference between flashed and moving stimuli on speed, luminance, and direction of motion.

**Dependence of latency difference on bar speed.** We asked whether the latency difference between the responses to flashed and moving bars depends on the speed of the moving bars. To this end, we recorded neural activity when a flash or a moving bar was presented and estimated response peak latencies, using the receptive field center as a reference location (Fig. 5A). We then asked how long the neuron takes to reach its peak firing rate for a bar that is flashed at this location and for the same bar at the same location when it is part of a motion trajectory (Fig. 5A). For both stimuli, the time of response peak with respect to the time at which the bar appears (flash, Fig. 5B) or arrives (motion, Fig. 5C) at the same reference location was taken as their respective representation delays (Fig. 5D). The assumptions behind using response peaks for computing representation delays are described in Materials and Methods. Note that when the speed increases, the time required for the bar to arrive at the reference location decreases (Fig. 5D). However, this difference in bar arrival times does not add to motion latencies, as we measured latency after all stimuli arrive at a common reference location. The center of the receptive field is operationally defined as the region that elicits maximal response. Assuming a receptive field-based labeled-line code common for flashed and moving stimuli, if both stimuli are processed with the same delay, then when either the moving or the flashed bar is at the receptive field center they should both elicit their respective maximal response with the same delay. In other words, when the moving bar arrives at the receptive field center one would expect a peak response to occur with the same delay, because at that instant the moving bar is indistinguishable from a flash. In contrast, we find that the response peak for all three moving stimulus conditions occurs earlier compared with that of the flash (Fig. 5D). In addition, as the speed increases, the response peak latency also increases and approaches that of the flash. These observations suggest that a moving stimulus is processed differently from a flashed one and is represented earlier in time in a speed-dependent manner compared with a flash in the same location.

To estimate latency at the population level, we chose to first average the responses across the multiunits and then compute response peak latency from this average rather than vice versa. This was done because some multiunit responses had multiple response peaks, making it unclear as to which peak should be considered for latency estimation, and in experiment 2 the individual unit responses were too weak (Fig. 8) at the lowest luminance values to reliably find the response peak. Averaging the responses over the multiunits first enabled us to robustly estimate latency and to apply a single procedure uniformly across all stimulus conditions.

Across our sample of multiunits from each monkey (Fig. 6), the peak response latencies for the motion condition at all three speeds were shorter compared with those for flashes (Fig. 7, A and B; for each monkey, P < 0.0005, Bonferroni corrected, bootstrap test; see Materials and Methods). As the speed increased, the latency of the motion response approached that of the flash (Fig. 7B). Therefore, the latency difference between flash and motion decreased as the speed increased (Fig. 7C; P < 0.0005, bootstrap test) but remained greater than zero (P < 0.0005, Bonferroni corrected, bootstrap test).

This effect is consistent with the speed dependence of the magnitude of the perceived spatial offset observed in the psychophysical data collected in macaques (Subramaniyan et al. 2013). In our electrophysiological experiments, we manipulated the speed and measured the representation latencies of flash and moving bar rather than the perceived spatial offset, which cannot be computed directly from the neural responses since it is a subjectively perceived quantity. In the psychophysical test, the subjects report the relative spatial offset between the flash and moving bar rather than how far the moving bar lags behind its own veridical location. Hence in computing the neural equivalent of perceived spatial offset, the latency of moving bar alone cannot be used—it is the difference (L̄d) in the representation delays of flash (L̄f) and moving (L̄m) that is needed. The neural equivalent of perceived spatial offset (X) was then computed by multiplying the speed (v) by the latency difference, i.e.,

\[ X(v) = v \cdot L̄d = v \cdot (L̄f - L̄m) \]  \hspace{1cm} (7)

Although the latency difference decreased with speed, the perceived spatial offset equivalent increased with speed (Fig. 7D; P < 0.0005, bootstrap test). This counterintuitive effect can be explained by noting that the latency difference is not a constant but varies with speed (Fig. 7C). Hence,

\[ X(v) = v \cdot [L̄f - L̄m(v)] \] \hspace{1cm} (8)
Fig. 5. Single-neuron responses to flash and moving bar and estimation of response peak latencies. A: illustration (drawn to scale) of the bar stimulus (rectangle) in visual space. Red dotted circle shows the 2-standard deviation outline of the neuron’s (from monkey A) receptive field (RF), with the RF center marked by the asterisk. In all panels, the vertical gray dashed line refers to the azimuth (1.11°) of RF center. In the flash condition, the bar is presented for 1 video frame as depicted. For moving conditions, the bar center occupies sequential positions marked by black dots (6.8°/s), small red circles (13.7°/s), or large blue circles (27.3°/s); the bar shown is the instantaneous moving bar position that matches the flash. Triangles indicate the starting position of the moving bar. B, left: raster plot showing neural responses to the flash shown in A, aligned to stimulus onset time. Each dot denotes a spike, and each row is a trial (only a subset of trials is shown). Right: mean firing rate plot for flash. C, left: raster plots of responses for the bar moving from left to right at speeds indicated on y-axes. Response times are aligned to motion trajectory onset time marked by triangles. Gray horizontal bars mark the time needed to traverse the horizontal spatial extent of the receptive field outline shown in A. Right: mean firing rate plots corresponding to the respective raster plots shown on left. In all subpanels of C, the time at which the moving bar center crosses the receptive field center is marked by an asterisk. D: mean firing rate responses to all stimuli. Flash response is aligned to flash onset time. Moving bar responses are aligned to the time (asterisks in C) at which the moving bar center crosses the receptive field center. The latency of response peaks for flash and moving bars is computed from this plot.
Fig. 6. Trial-averaged responses of multiunits in the speed manipulation experiment in monkeys CH, CL, and A. Columns of panels represent monkey subjects indicated at top. Rows of panels represent stimulus conditions indicated on left. In each image, each row, ordered in ascending order of recording day, represents response of a multiunit. Vertical white line marks the time the stimulus hits the receptive field center. Horizontal white line separates the 2 motion direction conditions: L→R, motion from left to right; R→L, motion from right to left. Color range is clipped at 95th percentile of responses grouped from all conditions.
Differentiating both sides with respect to speed,
\[
\frac{dX(v)}{dv} = L_f - \left[ v \cdot \frac{dL_m(v)}{dv} + L_m(v) \right]
\]  
(9)

From Eq. 9, for the perceived spatial offset to increase with speed, i.e., for \( \frac{dX(v)}{dv} > 0 \),
\[
L_f - \left[ v \cdot \frac{dL_m(v)}{dv} + L_m(v) \right] > 0
\]
or
\[
L_f > \left[ v \cdot \frac{dL_m(v)}{dv} + L_m(v) \right]
\]  
(10)

Hence, as long as Eq. 10 is satisfied, the perceived spatial offset will increase with speed even if motion latency increases \( \frac{dL_m(v)}{dv} > 0 \) (our data) or decreases \( \frac{dL_m(v)}{dv} < 0 \) with speed.

The simplest case arises when motion latency does not change with speed \( \frac{dL_m(v)}{dv} = 0 \), that is,
\[
L_f = L_m
\]
\[
L_f - L_m = c
\]
where \( c \) is a constant.

Therefore,
\[
X(v) = v \cdot c
\]  
(11)

Hence Eq. 11 shows that the perceived spatial offset linearly increases with speed as long as \( L_f > L_m \). This assumption of constant motion latency is commonly made in the psychophysical literature of flash lag illusion. Consistent with this assumption, some psychophysical studies have shown that the perceived spatial offset increases with speed (see DISCUSSION). However, as indicated by Eq. 10, this assumption is not necessary for explaining the increase in perceived spatial offset with speed. Our results demonstrate that despite an increase in motion latency with speed (Fig. 7B), the neural equivalent of perceived spatial offset increases with speed (Fig. 7D). In summary, when \( L_m \) does not vary with speed, the spatial offset equivalent will increase with speed as long as \( L_f > L_m \). On the other hand, when \( L_m \) changes with speed, the condition \( L_f > L_m \) alone is not sufficient. For example, for certain values of \( \frac{dL_m(v)}{dv} \) the condition in Eq. 10 may not hold and consequently the spatial offset equivalent will not increase with speed even when \( L_f > L_m \). Hence to observe a neural correlate (increasing spatial offset equivalent with speed), the empirically measured latencies need to satisfy the condition in Eq. 10.

The increase of the perceived spatial offset equivalent with speed is consistent with our psychophysical results (Subramaniyan et al. 2013) from two other monkeys of the same species (Fig. 7E) and with human psychophysical studies (see DISCUSSION). Together, these results show that in V1, irrespective of the speed, the moving bar latency is not fully compensated (zero latency) as would be predicted by the spatial extrapolation model and that the latencies of flash and moving bar are not equal as would be predicted by the motion-biasing model. On the other hand, our results are consistent with the differential latency model.

Dependence of latency difference on bar luminance. The latency difference between moving and flashed bars may also depend on bar luminance. To test this, in the second experiment, we fixed the speed of the moving bar at 18°/s and presented flashes and moving bars with luminance values of 0.2, 0.8, 9, and 48 cd/m² (Fig. 8). We found that the motion response occurred earlier in time relative to the flash response (Fig. 9A). For all luminance values tested, the motion response peak latency was lower than that of the flash (Fig. 9B; \( P < 0.0005 \), Bonferroni corrected, bootstrap test). For both the flash and moving bar, the peak response latencies decreased as the luminance increased, although they decreased differently (Fig. 9B). Accordingly, the latency difference decreased as the luminance increased (Fig. 9C; \( P < 0.0005 \), bootstrap test).
To compare physiological and psychophysical data, we again converted the latency differences into perceived spatial offset equivalent by multiplying the latency differences by speed (Eq. 7). The perceived spatial offset equivalent decreased with luminance (Fig. 9C, \( P < 0.0005 \), bootstrap test). Although we currently do not have psychophysical data on the luminance dependence of the flash lag effect in monkeys, we showed previously that monkeys perceive the illusion similar to humans (Subramaniyan et al. 2013). We therefore measured perceived spatial offsets from two human subjects, using the same luminance and stimulus parameters used for the monkey physiology. Indeed, the perceived spatial offset decreased with luminance in both observers [Fig. 9, \( D \) and \( E \); \( F(1, 24) = 14.6, P = 0.001 \); linear mixed model], in good agreement with the physiological results.

In the above analysis, we computed latency difference data between flash and moving bar with identical luminance and showed that they correlate well with human psychophysical data. Given that we presented each luminance condition in isolation, it is possible to compute the latency difference between a flash and a moving bar having different luminance values. In human psychophysics, when the flash luminance is fixed at a very low detectability level the perceived spatial offset increases with the moving bar luminance (Ögmen et al. 2004; Purushothaman et al. 1998). To see whether this is also evident in our neural data, we used the latency of the flash
condition with the lowest luminance to compute latency difference at all moving bar luminance conditions. Interestingly, qualitatively similar to the human psychophysical results, we found that the perceived spatial offset equivalent increased ($P < 0.0005$, bootstrap test) with the moving bar luminance (Fig. 9F). An even more interesting psychophysical result is obtained when the moving bar luminance is fixed at a very low detectability level and the flash luminance is varied. For a sufficiently high flash luminance, the flash lag effect is reversed where humans perceive the flash to be in front of the moving bar (flash lead effect) (Ögmen et al. 2004; Purushothaman et al. 1998). We again found a qualitatively similar result in our neural data (Fig. 3G), where the perceived spatial offset equivalent decreased ($P < 0.0005$, bootstrap test), changing from being positive (flash lag) to negative (flash lead) as the flash luminance level was increased, correlating well with the human psychophysical results.

Dependence of latency difference on motion direction. In addition to speed and luminance, the direction of motion has also been shown to affect the perceived spatial offset. Humans report a larger spatial offset for motion toward fovea (foveopetal; Fig. 10A) than motion away from fovea (foveofugal) (Kanai et al. 2004; Mateeff et al. 1991; Shi and Nijhawan 2008). We reproduced this finding in our stimulus paradigm, where humans reported a higher spatial offset for foveopetal motion direction in a speed-dependent manner [Fig. 10B; significant speed effect: $F(1, 93.2) = 14.8, P < 0.001$; nonsignificant motion condition effect: $F(1, 75.8) = 2.56, P = 0.11$; significant speed $\times$ motion condition interaction: $F(1, 79.2) = 10.4, P < 0.01$]. Surprisingly, in the monkeys this motion effect was reversed under the same stimulus conditions (Fig. 10, C and D; significant main effects and interaction: speed: $F(1, 64) = 27.3, P < 0.001$; motion condition: $F(1, 67.6) = 12, P = 0.001$; speed $\times$ motion condition: $F(1, 64.8) = 6.6, P = 0.013$). Correlating with this, the neural response latencies were lower (Fig. 10E) and the perceived spatial offset equivalents were higher (Fig. 10, F and G) for the foveofugal condition in two of the three monkeys (latency and perceived spatial offset equivalent: $P < 0.0005$ for CH and CL and $P > 0.05$ for A; Bonferroni corrected for multiple speeds, bootstrap test). It should be noted that in the neural data from all three monkeys ($CH, CL,$ and $A$) the receptive fields were in the right hemifield. Consequently, foveopetal condition is inseparable from motion from right to left visual hemifield, and the neural effect we observed may reflect the later rather than the former condition. However, this is less likely for the following reason. In monkey $L$, for which we varied stimulus luminance, the receptive fields were in the hemifield opposite to that of the above other three monkeys (Fig. 4D). This led to foveopetal condition being coupled with motion from left to
right hemifield. Despite this, we observed the same effect found in the other data set (CH, CL, and A), i.e., the latencies were lower (Fig. 10H) and the perceived spatial offset equivalents were higher (Fig. 10I; $P < 0.0005$, Bonferroni corrected, bootstrap test) for foveofugal condition under all luminance values tested, suggesting that in monkeys motion away from fovea produces a larger flash lag effect. The internal consistency between psychophysical and neural data within the monkey species strongly suggests that latency difference can explain a species-specific aspect of the flash lag illusion.

**Simultaneous presentation of flashed and moving stimuli.** In summary, our physiological data from speed and luminance manipulation are in good agreement with psychophysical results and the predictions of the differential latency model of the flash lag effect. One potential caveat is that in our physiology experiments we presented the flashes and moving bars in isolation. However, to generate the flash lag illusion the flashed and the moving bar are presented simultaneously with perfect alignment. It is thus conceivable that if we had presented the flash and the moving bar together the results might have been different. To rule out this possibility, we conducted a control experiment in which we presented the flash and moving bar together at different spatial offsets, including a zero-offset condition in which the flashed and the moving bar were in alignment. This allowed us to determine whether there is a change in latency as a function of spatial offset for simultaneously displayed stimuli.

We presented the flashes and moving bars simultaneously (“combined” condition) in two different arrangements. We presented flashes at the receptive fields and the moving bar together at different spatial offsets, including a zero-offset condition in which the flashed and the moving bar were in isolation.
between the combined and single conditions (obtained where stimuli were presented in isolation (“single”)
offsets in the combined condition and compared them to those
condition a second stimulus affected response latencies, we
motion inside the receptive fields. To test whether in the combined
stimulus outside the classical receptive field.

These results suggest that in awake fixating
population activity is diminished, we averaged the motion latencies
across the two motion directions.

A probabilistic Bayesian decoder (see MATERIALS AND
METHODS) was used to estimate the representation delays of
the stimuli based on simultaneously recorded single-unit or
multiunit population activity. We assumed that the neurons
spike as inhomogeneous Poisson processes that are condi-
tionally independent given the stimulus and used a decoder
tained on flashes to decode moving stimuli. It is well
established that population activity in V1 at a given time is
influenced by the location of the bar stimulus and signal
conduction and processing delays. This notion is captured in
the forward probabilistic model of population activity in Fig.
3. Based on this formalism, a joint distribution of stimulus
location, population activity, and response delay was obtained
(Eq. 2) from which a posterior probability estimate (Eq. 5) of
a stimulus position can be obtained from the population activity
at any given time. On the basis of the encoding that was
learned from the flash-evoked responses, we decoded the
position of the moving bar under different speeds and lumin-
ance values. For decoding flashes, we used trials that were not
used for encoding, to prevent overfitting. For the luminance
modulation experiment, the decoding of bar stimuli of a given
luminance was based on encoding obtained from responses to
flashes of matching luminance.

The probability of the stimulus position given population activity at different times was computed trial by trial with
simultaneously recorded single units or multiunits (Figs. 12–
14, Ax). The resulting position estimates were first averaged
across trials and then across sessions (Figs. 12–14, B and D).
The latency of the peak of the posterior probability (Figs.
12–14, C) was taken as the representation delay of the flashes.
For the moving bars, first we computed the distance (spatial
lag) between the most probable stimulus location and the
instantaneous location of the moving bar. Toward this, the trial
and session-averaged posterior probabilities (rows in Figs.
12–14, D) were aligned (centered) to the instantaneous hori-
zontal positions of the moving bar center (white dots in Figs.
12–14, D). For each speed and direction, the aligned probabil-
ities were averaged across the instantaneous positions of the
motion trajectory (Figs. 12–14, E). The distance between the
peak of this aligned probability and the origin gives the spatial
lag of the most probable stimulus location. Note that we did not
aligning individual neuronal responses to stimulus location in
their receptive field centers. However, it is possible that neu-
ronal representation delays based on population coding may
lead to different conclusions. Hence we proceeded to check
whether we could reproduce the main results of the study
presented in Figs. 7 and 9 with probabilistic population decod-
ing that does not use any response alignment to receptive field
centers to compute representation delays. Rather, the moving
bar position is decoded based on the population response. It
should be noted that this approach was restricted to the results
presented in Figs. 7 and 9 and not used for results in Figs. 10
and 11 because there was an insufficient number of neurons for
reliable decoding. We also pooled the two motion directions to
obtain a robust estimate of motion latency especially at high
speeds, where the moving bar traverses the decoded space very
quickly, giving much fewer trajectory positions to obtain a
reliable latency estimate. Similarly, to improve the position
decoding under the lower-luminance conditions where the
neural activity is diminished, we averaged the motion latencies
right) at five to seven different spatial offsets in a gray
background. For analysis, we chose the central three offset
conditions that had a sufficient number of multiunits (see
MATERIALS AND METHODS). We then computed the flash response
peak latencies from the first arrangement and the motion
response peak latencies from the second. The latency differ-
ence was not significantly different among the three spatial
offsets (P > 0.76, bootstrap test). In the same recording
sessions, we also presented flashes and moving bars in isolation
inside the receptive fields. To test whether in the combined
condition a second stimulus affected response latencies, we
pooled the latency difference data across monkeys and spatial
offsets in the combined condition and compared them to those
obtained where stimuli were presented in isolation (“single”
condition; Fig. 11, B and C). We found no significant differ-
ce between the combined and single conditions (P > 0.99,
bootstrap test). These results suggest that in awake fixating
macaques the latencies of the flash or moving bar representa-
tion in V1 are not influenced by the presence of a second bar
stimulus outside the classical receptive field.

Population decoding of flashed and moving bars. The
conclusions reached so far were based on latencies estimated by

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intend to decode the motion speed; hence we treated it as a known quantity. The latency of the moving bar representation was then computed by dividing the spatial lag by speed.

As reported in Fig. 7, B–D, in all three monkeys, based on multiunit population decoding, as speed increased the motion latency increased (Fig. 12F; $P < 0.0005$, bootstrap test), latency difference decreased (Fig. 12G; $P < 0.0005$, bootstrap test), and the perceived spatial offset equivalent increased (Fig. 12H; $P < 0.0005$, bootstrap test). From one of the monkeys (CL), we were able to isolate a sufficiently large number of single units, so we were able to verify that the results held true for single well-isolated neurons (Fig. 13, F–H) as well.

For the luminance modulation experiment, we decoded stimulus position for flashes (Fig. 14, B and C) and moving bars (Fig. 14, D and E) as described above. Again, as found above in Fig. 9, the multiunit population decoding showed that for all luminance values tested the latency of moving bar was less than that of flashes (Fig. 14F; $P < 0.0005$, Bonferroni corrected), latency difference and perceived spatial offset equivalent decreased with luminance (Fig. 14G; $P < 0.0005$, bootstrap test). Similarly, the perceived spatial offset equivalent increased with moving bar luminance when flash luminance was fixed at the lowest value tested (Fig. 14H; $P < 0.0005$, bootstrap test). When the moving bar luminance was fixed at the lowest value tested, the perceived spatial offset equivalent decreased (Fig. 14I; $P < 0.0005$, bootstrap test), changing from being positive (flash lag) to negative (flash lead) as the flash luminance level was increased. These results suggest that our conclusions on speed and luminance dependence of latencies and perceived spatial offset equivalents based on individual multiunit responses are consistent with those obtained by population decoding.

**DISCUSSION**

Our results show that moving stimuli are processed faster than flashed stimuli in awake macaque V1. In particular, the
latency difference between the neural representations of the two stimuli depends on luminance and speed in a way that resembles the perceptual effects of these manipulations in both monkeys (Subramaniyan et al. 2013) and humans (Krekelberg and Lappe 1999; Murakami 2001; Nijhawan 1994; Ögmen et al. 2004; Patel et al. 2000; Purushothaman et al. 1998; Subramaniyan et al. 2013).

Both precortical and cortical mechanisms likely contribute to the observed faster motion processing. These mechanisms potentially include motion-induced dynamic shift in the receptive field location and faster conduction/processing of motion signals. Our data cannot distinguish between these two possibilities since both will give rise to a shift in motion response relative to flash response. Motion-induced receptive field shifts have not been reported in the precortical stages in macaques. If found, it would suggest that the labeled line code is not static but more dynamic and will depend on properties of the stimuli.

However, there is some evidence for shorter latency of motion signals in the precortical stage—the LGN. In anesthetized cats, it was found that in the different types of LGN cells the response peak latency for a moving bar was shorter compared with that of a flashed bar (Orban et al. 1985). Future studies are needed to confirm these findings in monkeys to locate the mechanisms underlying the flash lag effect. Cortical processing such as gain control similar to that described in the retina (Berry et al. 1999) and motion-related feedback signals may contribute to dynamic shift in the receptive field location toward the motion direction. For example, a recent study (Ni et al. 2014) found that V1 receptive fields in fixating macaques shifted by ~10% (0.1°) on average in the direction that accounted for the size-distance illusion. Such receptive field shifts, if induced by motion, can readily explain part of the faster motion processing. Another study that addressed a different illusion called flash-jump illusion also found that V4
neuronal receptive fields shift when the color of one of the bars of an apparent motion sequence changes abruptly (Sundberg et al. 2006). Given that a color change was necessary for such a shift, the implications of their study for the neural mechanisms of flash lag illusion remains unclear.

Faster cortical motion processing could also be achieved by the spreading of subthreshold activity through lateral connections from the currently activated cortical region into the region activated in the future. This spread may facilitate responses by bringing the membrane potential of the target neurons closer to threshold. As a result, those neurons will reach their peak firing earlier, resulting in shorter motion latency. The influence of such subthreshold activity has already been reported in cat V1 in the context of line-motion illusion, where the spread of subthreshold activity initiated by one stimulus facilitates the response to a subsequently presented stimulus (Jancke et al. 2004a). In light of this mechanism, it could also be expected that the slower motion would exhibit shorter latency through this mechanism compared with the faster one, as there would be more time for the subthreshold activity to spread farther for the slower compared with the faster motion, potentially explaining the speed dependence of motion latency we observed.
We found that the moving bar response peak latency increased with speed. Consistent with our results, conversion of the direction-averaged spatial lag data reported by Jancke et al. (2004b) (Fig. 6 in their study) into latency also revealed a similar trend in the speed dependence of motion peak latency. Our data show that latency difference between flash and motion condition decreased with speed. This is in sharp contrast to the constant latency difference that most psychophysical studies assume when interpreting the effect of speed in perceived spatial offset (Krekelberg and Lappe 1999; Murakami 2001; Nijhawan 1994; Whitney et al. 2000). Equivalent latency difference computed from the perceived spatial offsets from a recent psychophysical study (Wojtach et al. 2008), however, clearly decreases with speed (Fig. 15), similar to our findings. The discrepancy among the psychophysical studies can be reconciled by noting that Wojtach et al. (2008) used a wide range of speeds (up to 50°/s), whereas the previous studies used a narrow speed range (up to ~15°/s), which missed the full trend of the speed effect.

We found that the perceived spatial offset equivalent depended on speed and luminance (Fig. 7D, Fig. 9C, Figs. 12 and 13, H, and Fig. 14G), in line with psychophysical results (Fig. 7E and Fig. 9E). The magnitude of the perceived offset computed from the population decoding method appeared to be closer to the behaviorally measured values than the values computed based on individual multiunit activity. Interpreting our data conservatively, we think that the perceived spatial offset equivalents we measured in V1 are likely to be smaller than the behaviorally measured values for the following reasons. 1) We measured neural responses from the very first cortical processing stage, and the physiological effect may get larger as the information is processed further in the higher cortical areas; 2) the smaller receptive field sizes in V1 may potentially limit the extent to which receptive field shifts can occur in order to reduce motion stimulus representation delays; and 3) the monkeys we recorded from did not perform the task, and making a relative position judgment may lead to a larger physiological effect. Moreover, we may have also underestimated the discrepancy between the behaviorally measured perceived spatial offset and its neural equivalent because we presented flashes randomly in multiple locations (5–7) for physiology, whereas for psychophysics the flash was presented at one (Fig. 9E) or two (Fig. 7E, Fig. 10, B–D) fixed locations. Given that predictability of flashes is known to reduce the flash lag effect (Baldo et al. 2002; Brenner and Smeets 2000; Krekelberg et al. 2000; Vreven and Verghese 2005), it is possible that psychophysical measurement of the lag could have been higher if the flashes were equally unpredictable, as in our physiology experiments.

It should be noted that the human psychophysical data were collected from two nonnaive subjects whose bias could have an effect on the observed data. We think that this is less likely because our results build upon previously well-established psychophysical results on luminance manipulation (Lappe and Krekelberg 1998; Öğmen et al. 2004; Purushothaman et al. 1998) and are well in accordance with what would be predicted from them. Nevertheless, further experiments from naive subjects would be essential to confirm our human psychophysical results.

In our luminance manipulation experiment, we kept the background luminance near zero and changed only the bar luminance. This stimulus configuration, although suitable for mimicking flash lag psychophysical experiments, is not readily comparable to previous psychophysical studies in V1 that examined luminance or contrast effect on latency using different stimulus configurations (Carandini and Heeger 1994; Gawne et al. 1996; Maunsell and Gibson 1992; Oram 2010; Reich et al. 2001). Despite these stimulus differences, similar to the above.
flashed stimuli are misaligned. However, in our main experience, the relative position judgment to decide whether the moving and flashed objects are equal, in contrast to what we observed only when the subjects are asked to make an explicit prediction that a neural correlate of the motion biasing would be unclear is whether and exactly when V1 spatial representations of the brain this “biasing” process is implemented. Also, it is unclear exactly in which parts of the brain this “biasing” process is implemented. Although species difference could be partly responsible for this, further investigations are needed to fully understand the sources of the discrepancy.

Our data provide two independent lines of evidence supporting the differential latency model (Ögmen et al. 2004; Patel et al. 2000; Purushothaman et al. 1998; Whitney and Murakami 1998), which predicts a shorter time needed for representing moving stimuli. First, as predicted, the perceived spatial offset equivalent computed directly from the latency difference increased with the speed of the moving bar. Second, the luminance dependence of the flash and motion representation delays (Fig. 9B) is also consistent with the key predictions of the differential latency model (Ögmen et al. 2004; Patel et al. 2000; Purushothaman et al. 1998), namely, that for a fixed low flash luminance the perceived spatial offset should increase with moving bar luminance and for a fixed low moving stimulus luminance progressively increasing the flash luminance should change the flash lag to a flash lead effect. Our neural data support both predictions (Fig. 9, F and G). In addition, latency differences (Fig. 9C) also explained the trend in the luminance modulation of perceived spatial offset using identical luminance for flash and moving bar that we showed in humans (Fig. 9E) for the first time.

According to the motion-biasing model (Eagleman and Sejnowski 2007; Rao et al. 2001), the latencies of flash and moving bar representations are equal, in contrast to what we find in our data. In addition, the illusion arises because “when the brain is triggered to make an instantaneous position judgment, motion signals that stream in over ~80 ms after the triggering event (e.g., a flash) will bias the localization” (Eagleman and Sejnowski 2007). It is unclear exactly in which parts of the brain this “biasing” process is implemented. Also unclear is whether and exactly when V1 spatial representations are altered by this “biasing.” Moreover, this model would predict that a neural correlate of the motion biasing would be observed only when the subjects are asked to make an explicit relative position judgment to decide whether the moving and flashed stimuli are misaligned. However, in our main experiments, only a flash or a moving bar was presented in isolation, and the animals used in our study were neither trained to make any relative position judgment nor trained in any other task like the present task; we still found a neural correlate of the illusion in V1. First, these results suggest that reporting relative position judgment is not necessary for observing a neural correlate of the flash lag illusion in visual area V1. Second, they argue against the current version of the motion-biasing model that involves only higher cognitive functions (Eagleman and Sejnowski 2007) and suggest that low-level mechanisms underlying the observed latency differences need to be taken into account.

While there is substantial evidence against the spatial extrapolation model at the psychophysical level (Baldo and Klein 1995; Brenner and Smeets 2000; Eagleman and Sejnowski 2000; Lappe and Krekelberg 1998; Purushothaman et al. 1998; Whitney and Murakami 1998), it is possible that spatial extrapolation could be happening at the level of V1. Given that any spatial extrapolation would manifest as a reduction in latency as measured by our method, a full delay compensation as predicted by the model would result in zero response peak latency for the moving bar. However, this was not the case, as we found significant delays for the moving bar at all speed and luminance conditions tested. Nevertheless, spatial extrapolation might still hold true in other brain regions or for other sensory systems as shown for auditory motion (Witten et al. 2006).

Irrespective of the model of the flash lag illusion, if the motion representation/perception delays are not ultimately reduced to zero, moving objects will always be mislocalized. Our results suggest that the overall shorter motion latency compared with flashes helps to reduce this mislocalization. Given our results that motion response latencies also change with speed and luminance, how would organisms cope with this in behaviors that require accurate localization of moving objects? One simple and viable solution would be calibration of the sensorimotor integration system. For example, to accurately hit the ball in a baseball game, players spend numerous hours in learning (calibrating) to swing the bat at the correct time, taking the speed of the ball into account. Hence, the nervous system could in principle learn to respond appropriately to a given moving stimulus condition.

We focused our study on V1, where both flash and motion signals first arrive in the cortex. We showed that moving objects are processed faster in a speed-, direction of motion-, and luminance-dependent way compared with suddenly appearing static stimuli. These provide a neural correlate of the flash lag illusion. In this visual area, our data are fully consistent with the predictions of the differential latency model. While the motion-biasing model cannot explain our results, this in itself is not evidence against the model in its entirety. It is possible that the monkeys need to perform the task for the mechanisms proposed by the model to be activated. Visual signals leaving V1 reach a multitude of cortical areas. It is yet to be seen whether the differential latency theory would hold in these other areas. Hence further combined behavioral and physiological studies in V1 and subsequent processing stages in the brain are essential to generate additional constraints to narrow down the models.

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DISCLOSURES
No conflicts of interest, financial or otherwise, are declared by the authors.

AUTHOR CONTRIBUTIONS
M.S., A.S.E., S.S.P., M.B., and A.T. conceived and designed research; M.S., A.S.E., S.S.P., R.J.C., and P.B. performed experiments; M.S., A.S.E., and R.J.C. analyzed data; M.S., A.S.E., S.S.P., R.J.C., M.B., X.P., P.B., and A.T. interpreted results of experiments; M.S. prepared figures; M.S. drafted manuscript; M.S., A.S.E., S.S.P., R.J.C., M.B., X.P., P.B., and A.T. edited and revised manuscript; M.S., A.S.E., S.S.P., R.J.C., M.B., X.P., P.B., and A.T. approved final version of manuscript.

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