

Decision-making with multiple alternatives

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Simple perceptual tasks have laid the groundwork for understanding the neurobiology of decision-making. Here, we examined this foundation to explain how decision-making circuitry adjusts in the face of a more difficult task. We measured behavioral and physiological responses of monkeys on a two- and four-choice direction-discrimination decision task. For both tasks, firing rates in the lateral intraparietal area appeared to reflect the accumulation of evidence for or against each choice. Evidence accumulation began at a lower firing rate for the four-choice task, but reached a common level by the end of the decision process. The larger excursion suggests that the subjects required more evidence before making a choice. Furthermore, on both tasks, we observed a time-dependent rise in firing rates that may impose a deadline for deciding. These physiological observations constitute an effective strategy for handling increased task difficulty. The differences appear to explain subjects' accuracy and reaction times.

Organisms face decisions of varying complexity. In simple decisions, perceptual observations allow an animal to choose between action and inaction, or between two alternative actions. These are simple instances of complex cognitive processes, which may require additional information from the environment or from memory. The ability to delay a response to consider incoming information is a hallmark of higher brain function.

Decisions between two choices in a perceptual-motion task^{1–3} demonstrate mechanisms of decision-making. In the task, humans or monkeys reported the net direction of motion in a patch of moving random dots. In the reaction-time version², subjects communicated their decision with a saccade when they were ready. This version identifies the period when subjects are accumulating evidence for a decision, but have not yet committed to an alternative. How rapidly evidence accumulates depends on the strength of motion. The process ends when the evidence reaches a threshold or bound corresponding to one alternative. These 'bounded accumulation of evidence' models encompass multiple mechanisms that have been proposed to explain choice and decision time^{1,4,5}.

Several observations are consistent with the idea that evidence accumulates to a bound. First, a formal model of bounded accumulation accounts quantitatively for subjects' speed and accuracy^{3,6}. Second, neuronal firing rates in the lateral intraparietal area (LIP) are consistent with evidence accumulation^{2,7–9}. Specifically, when stimulus motion favors the target in the response field of an LIP neuron, firing rates progressively build up as monkeys form their decisions; the rate of buildup is proportional to motion strength. Finally, in the reaction-time version, firing rates are similar at decision end when the monkey selects a target in the neuron's response field, for all motion strengths and reaction times. The stereotyped firing rate may reflect a bound that is common to both easy and difficult motion strengths.

Because two-alternative choice tasks are simple, they may offer limited insight into decision-making in general; organisms regularly face decisions with multiple alternatives. Here, we compare responses on a four-choice decision task with a two-choice task. Our results argue that the bounded accumulation framework can be extended to explain more complex decisions, and they begin to reveal how decision-making circuitry adjusts to increasingly difficult decisions.

RESULTS

Behavior

Two monkeys were trained on a two-choice and a four-choice motion-discrimination task (**Fig. 1**). We measured both the accuracy and speed of their choices. In both tasks, accuracy was nearly perfect at high motion strengths, but fell toward chance levels with lower motion strengths (**Fig. 1d**). The performance with strong motion indicates that the monkeys understood the relationship between stimulus direction and choice targets. Therefore, the more frequent errors at low motion strengths on the four-choice task may be the result of a failure to discriminate the direction of motion, rather than confusion about action selection.

Like accuracy, decision speed also depended on both motion strength and the number of choices. Reaction times were longer on the four-choice task¹⁰. These differences were largest at lower motion strengths, but were significant at all motion strengths ($P < 0.01$; **Fig. 1f**).

We measured responses on an additional condition with two targets, spaced 90° apart (**Fig. 1c**). This configuration (90° control) uses a subset of the targets and motion directions in the four-choice task. Accuracy on this task emulated the standard two-choice task (**Fig. 1e**), albeit with longer reaction times, which fell between those on the standard two- and four-choice conditions (**Fig. 1g**).

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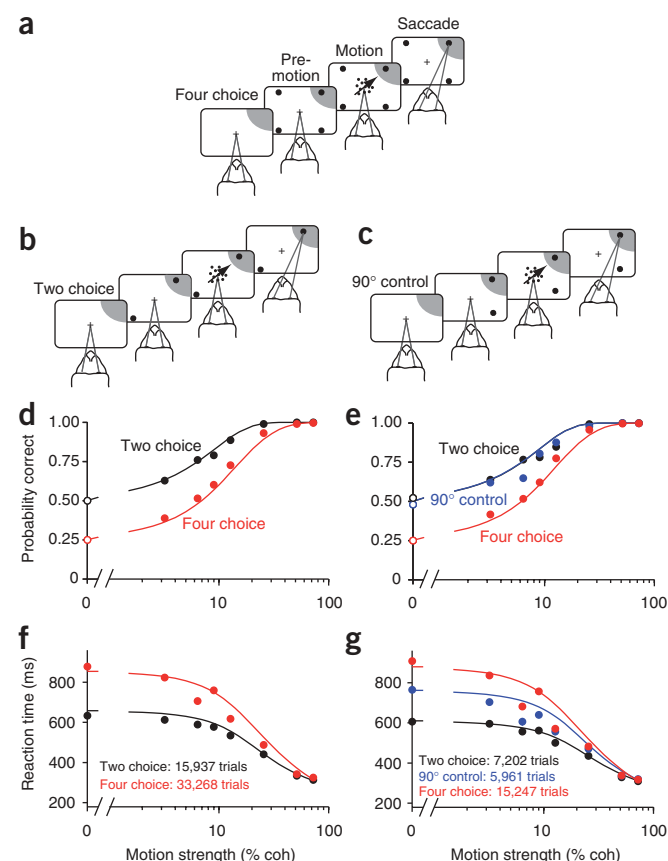


Figure 1 Task and performance. (a–c) Sequence of events on two- and four-choice direction-discrimination tasks. The monkey fixates a central point until the random dot motion appears and then indicates its decision by making a saccadic eye movement to a choice target. The motion is in one of two or four directions (trials randomly interleaved in a 1:2 ratio). A liquid reward is given for choosing the target along the axis of random dot motion, or it is given with probability 1/2 or 1/4 when the motion strength is zero. Random intervals (truncated exponential distributions) separate fixation, appearance of choice targets and motion onset. The random-dot motion is extinguished when the monkey initiates a saccade to one of the choice targets. One of the choice targets is in the response field of an LIP neuron recorded during the task (shading). The directions were 90° apart in the four-choice task (a). The directions were 180° apart in the two-choice task (b). One direction is toward the target in the neuron's response field (T_{in}). The 90° control task is shown in c. (d–g) Speed and accuracy of decisions. Smooth curves in all panels are fits to the bounded diffusion model. The fits were performed separately for d and f and for e and g. Psychometric functions are shown in d. The probability of a correct choice is plotted as a function of motion strength. All experiments contribute to these graphs. At 0% motion strength, choices were rewarded randomly (open symbols). Psychometric functions for the 29 experiments that included the 90° control are shown in e. Chronometric functions are shown in f. Mean reaction time for correct trials is plotted as a function of motion strength. Each point reflects correct responses from all experiments. Error bars for s.e.m. are smaller than the symbols. Chronometric functions for the 29 experiments that included the 90° control are shown in g.

to determine whether physiological differences on the two tasks can account for the behavioral differences that we observed (Fig. 1).

Comparison of responses before motion

Responses before the stimulus reflect the state of decision-making circuitry before the brain begins accumulating evidence. Target appearance was the first cue indicating whether the discrimination would involve two or four directions. Between target appearance and motion onset, there was a clear difference in firing rates between the two- and four-target conditions (Fig. 3). In a single neuron (Fig. 3a), responses were 11.9 ± 4.0 spikes s^{-1} lower on the four-choice task ($P < 0.003$). The reduced firing rate on the four-choice task was evident across the population of neurons tested (Fig. 3b,c). It was subtly apparent in the transient target onset response and then was prominent until motion onset (mean difference = 16.1 ± 1.6 spikes s^{-1} , $P < 10^{-5}$; Fig. 3c). This effect was statistically significant in both monkeys (Monkey I: mean difference = 15.5 ± 1.8 spikes s^{-1} , $P < 10^{-5}$; Monkey S: mean difference = 18.8 ± 2.6 spikes s^{-1} , $P < 10^{-4}$).

Firing rate was also reduced in the 90° control task, although much less than in the four-choice task (responses were 3.7 ± 2.1 spikes s^{-1} lower on the 90° control task than on the two-choice task, $P < 0.05$; Fig. 3d). We conclude that the smaller angular separation between the targets contributes to the reduction in activity seen in the four-choice task in this epoch, but that the reduction is largely explained by the number of choices.

Responses early in the motion epoch

The reduced firing rate on the four-choice task persisted after motion onset. Recall that the motion was presented outside the neuron's response field, and its earliest effect on LIP was a dip in the firing rate (Fig. 2a). During this dip, firing rates were 9.17 ± 1.05 spikes s^{-1} lower for the four-choice than for the two-choice task ($P < 10^{-5}$; Supplementary Fig. 3 online). Thus this epoch retains much of the difference in firing rate seen before motion onset.

Following the dip, there was a time-dependent rise in the firing rate ('buildup'). The rate of this buildup offers insight into the conversion of sensory information into a decision variable: that is, the form of the

Physiological responses on the four-choice task

The 70 neurons recorded in our experiment had spatially selective persistent activity on delayed and memory-guided eye-movement tasks (Supplementary Fig. 1 online). Multi- and single-unit responses with these characteristics are common in the ventral portion of LIP targeted in these experiments¹¹. In the motion-discrimination task, one of the choice targets (T_{in}), was centered in the neuron's response field. The other choice targets (T_{out} and two orthogonal targets, T_{90}) were evenly spaced around a central fixation point (see Methods). On any one trial, the direction of motion was toward one of the targets.

Figure 2 introduces the pattern of activity seen over the course of the trial on the four-choice task. Target appearance caused a large, transient increase in the firing rate (Fig. 2a,c), followed by the establishment of a baseline firing rate as the monkey awaited the onset of the random dot motion. Shortly after the onset, the responses underwent a brief dip, followed by a gradual rise in the firing rate. The dip, observed in other studies^{2,12–14}, seems to mark the beginning of evidence accumulation. After the dip, the rate of increase in the firing rate depended on motion strength. Near the saccade, however, this neural correlate of motion strength^{1,2,15} vanished (Fig. 2a,c). The stereotyped firing rate at the end of the decision was clearer when responses were grouped by reaction time (four choice, Fig. 2b,d; two choice, Supplementary Fig. 2 online), instead of by motion strength. Analysis of variability across different reaction-time groups provided quantitative support for a common firing rate close to the end of the trial (Supplementary Fig. 2).

This pattern of LIP activity in the four-choice task resembles the process underlying two-choice decisions^{1,2,16}. It is broadly consistent with a bounded accumulation of evidence among competing response mechanisms. Next, we compared responses on two- and four-choice before the motion, during motion viewing and just before the saccade

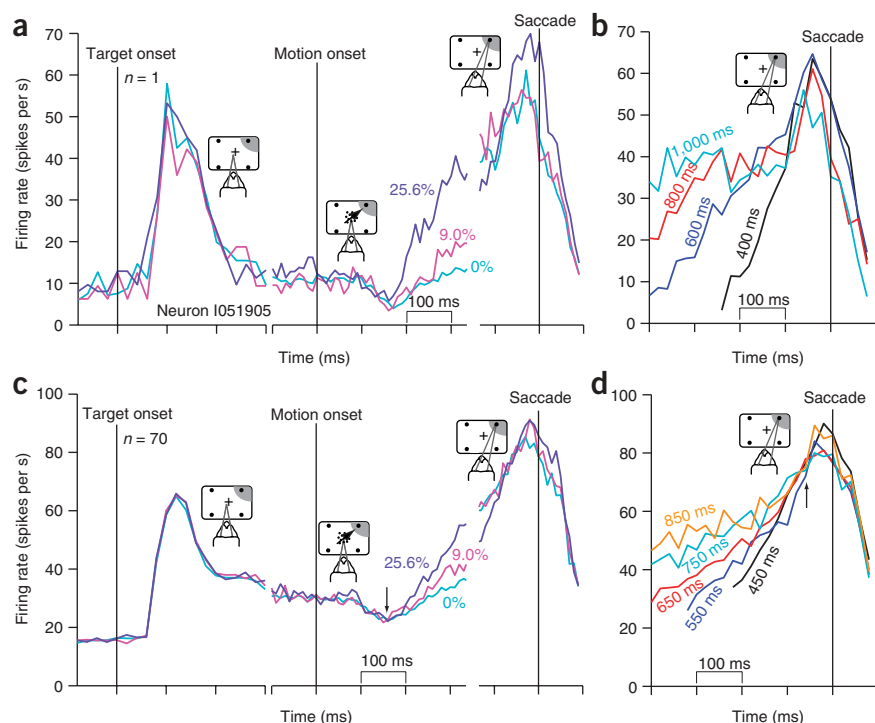


Figure 2 Responses of LIP neurons on the four-choice task are consistent with bounded accumulation. Firing rates are aligned to key events in the course of a trial, which are marked by vertical lines. Motion was either random (0% coherence) or in the T_{in} direction. **(a)** Average firing rates from one neuron. Left, responses aligned to the onset of the choice targets. Middle, responses aligned to the onset of stimulus motion. Right, responses aligned to saccade initiation. For saccade-aligned responses, only T_{in} choices are shown. Traces were smoothed with a 30-ms exponential filter. **(b)** Responses reflect termination of the decision. Responses are grouped by reaction time at the values indicated (± 25 ms). The averages are aligned to saccade initiation and exclude neural activity in the first 200 ms of motion onset. Only correct T_{in} choices were included and, for clarity, every other reaction-time group is not displayed. **(c,d)** Population average responses ($n = 70$ neurons). Same conventions as in **a** and **b**, except that no smoothing was performed; firing rates were computed in 20-ms nonoverlapping bins. Arrow in **c** indicates the dip in firing rate seen shortly after motion onset. Arrow in **d** indicates the time when responses appeared to coalesce, approximately 60 ms before the saccade.

evidence that underlies the choice and decision time. Four factors affect buildup: decision outcome, motion strength, the number of choices present and the passage of time (Figs. 4 and 5).

The effect of choice is most conspicuous late in the decision process (for example, Fig. 5a–d). Our sample was screened for spatially selective responses on delayed eye-movement task, so all neurons are expected to indicate the decision outcome. We were interested in the change in firing rate accompanying decision formation. Early on, decision outcome has only a weak effect on the neural responses. We therefore analyzed groups of trials with the same motion strength and direction, regardless of the monkey's eventual choice (our conclusions also hold if trials are grouped by motion strength and choice; Supplementary Fig. 4 online).

To quantify how sensory information is converted into decision evidence, we estimated firing-rate buildup at each motion strength (buildup rate = slope of a line fit to the shaded portion of the response;

Fig. 4a–d). Buildup rates scaled approximately linearly as a function of motion strength for motion toward and away from the neuron's response field. This relationship was similar on the two- and four-choice tasks. For motion toward T_{in} , these slopes differed by only 0.17 ± 0.45 spikes s^{-2} per unit change in motion strength (that is, per 1% random dot coherence; %coh $^{-1}$ from here on) ($P = 0.71$; Fig. 4f). For motion toward T_{out} , increasing motion strength suppressed buildup rate slightly more in the two-choice condition, but the difference was not reliable (difference = 0.50 ± 0.31 spikes s^{-2} %coh $^{-1}$; $P = 0.11$, see Methods, equation (3)). These trends were also apparent in single

Figure 3 Neural responses in the pre-motion epoch are larger on the two-choice task. **(a)** Average firing rate from a single neuron during the pre-motion epoch when two or four choice targets were displayed. Vertical black line indicates the onset of the choice targets. Insets are a schematic of the target configurations used in this experiment. One target is in the neuron's response field (shading). **(b)** Population average response. The same conventions are used as in **a**, except that the traces are average firing rates from 70 neurons. Insets illustrate that one target is in the response field of the neuron; the location of this response field varies from neuron to neuron. **(c)** Comparison of firing rates from individual neurons on the two- and four-choice tasks. Responses were measured from 200 to 300 ms after choice target onset. The green circle marks the neuron shown in **a**. Points for three neurons with high background firing rates are omitted from the plot to facilitate an appropriate scale for the remaining points ((227, 174), (132, 99) and (144, 127)). Error bars are s.e.m. Histogram shows the firing-rate differences for all 70 neurons. Shading indicates significance ($P < 0.05$). **(d)** Comparison of firing rates from individual neurons on the two-choice and the 90° control tasks. The same conventions are used as in **c**. Two neurons were omitted from the scatter plot ((131, 138) and (227, 226)).

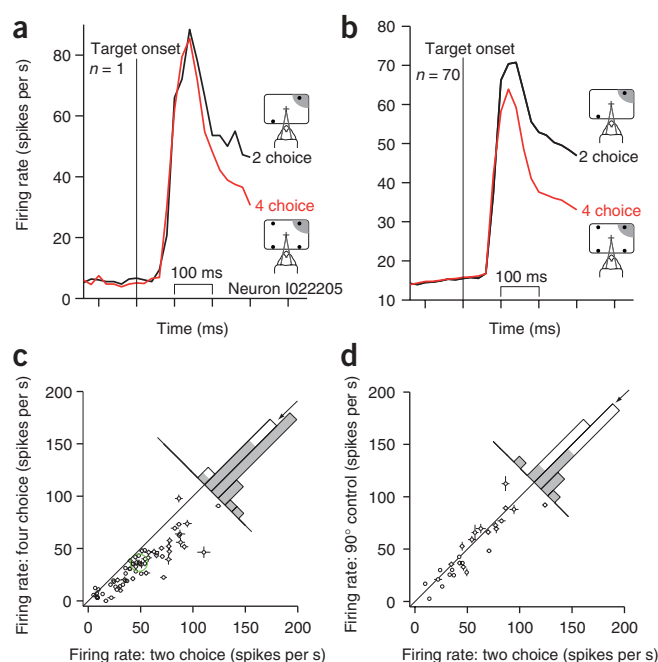
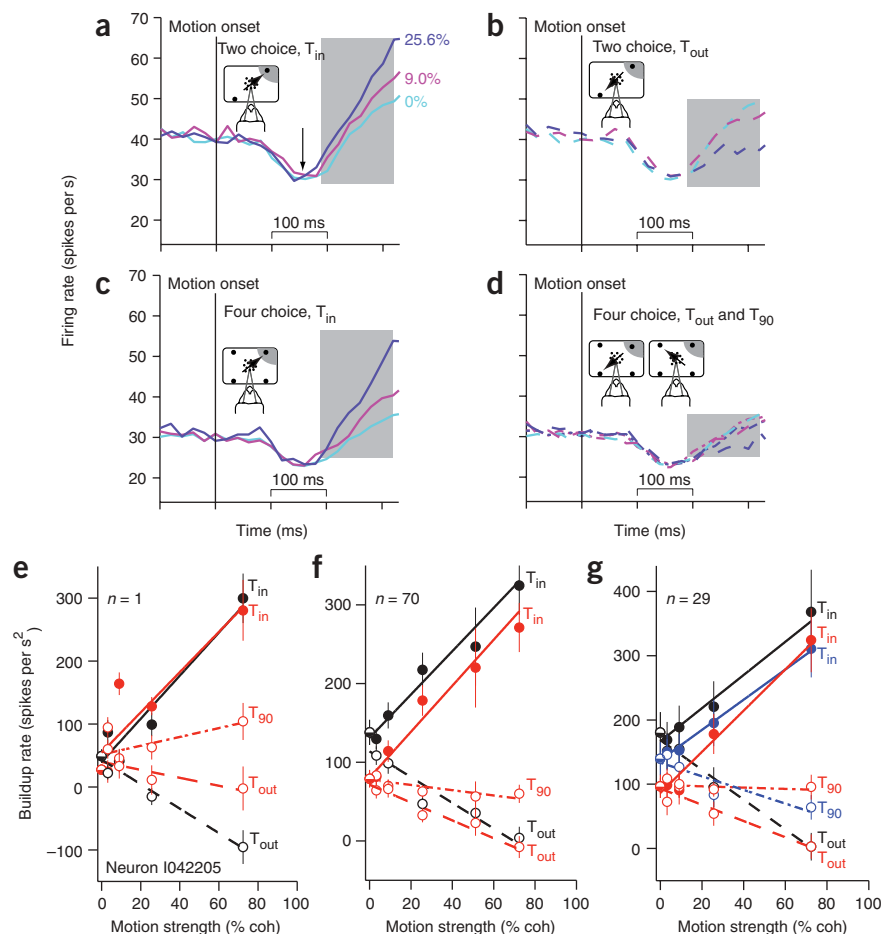


Figure 4 Neural responses during motion viewing depend on difficulty. (**a–d**) Population average firing rates ($n = 70$ neurons) for three motion strengths. Trials are grouped on the basis of the direction of motion and the number of choices (insets). Correct and incorrect trials are included in these averages. The traces for 0% coherent motion are identical in **a** and **b** and in **c** and **d**. Only three motion strengths are shown for clarity. Shaded rectangle indicates the epoch used to estimate the buildup rates (190–320 ms after the onset of stimulus motion). Two-choice trials where the motion direction was toward T_{in} are shown in **a** and trials where the motion direction was toward T_{out} are shown in **b**. Arrow indicates the stereotyped firing rate ‘dip’ that occurs after motion onset. Four-choice trials where the motion direction was toward T_{in} (**c**) or toward an orthogonally positioned (T_{90}) target (dot-dash line) are shown. The five traces in **d** are largely superimposed. The single cyan trace is for 0% coherent motion averaged across all choices (T_{in} , T_{out} and T_{90} ; same as cyan trace in **c**). (**e–g**) Effect of motion strength on buildup rates. A single neuron example is given in **e**. Buildup rates were estimated for each motion strength; these buildup rates (\pm s.e.m.) are plotted as a function of motion strength. The slope of the fitted line estimates the effect of a unit change in motion strength on the buildup rate (black, two choice; red, four choice). Five motion strengths were tested for this neuron (Methods). Buildup rates were calculated in individual neurons and then averaged across the population ($n = 70$ neurons) before fitting the line (**f**). Error bars are s.e.m. of buildup rates across neurons. Population analyses for the 29 neurons tested with the 90° control condition in addition to two- and four-choice trials are shown in **g**. Blue lines correspond to the 90° control condition; error bars are s.e.m. of buildup rates across neurons. Points corresponding to 51.2% motion strength are not included on this plot because this motion strength was tested in only three neurons.



neurons (Supplementary Fig. 5 online). We concluded that LIP registers evidence along the T_{in} – T_{out} axis similarly for two- and four-choice tasks. Thus, differences in behavior are not explained by differences in the mapping of motion information onto a change in LIP firing rate.

This analysis also suggests that LIP is only weakly affected by motion in directions orthogonal to the T_{in} – T_{out} axis on the four-choice task. Stronger motion reduced the firing rate slightly (slope of T_{90} trace = -0.33 ± 0.09 spikes $s^{-2} \%coh^{-1}$, $P < 10^{-3}$; Fig. 4f). Although either of the T_{90} directions might provide weak positive evidence for a T_{in} choice in any one experiment (for example, Fig. 4e), the net effect of orthogonal motion was, on average, weak negative evidence.

There is also a prominent buildup in firing rate that does not depend on motion strength or direction and is also not explained by the monkey's choice, as seen in 0% motion strength trials, which favor all directions equally (Fig. 4a–d). Although the monkeys distributed their choices with nearly equal frequency to all (two or four) choice targets, firing rates increased as a function of time. The rate of this buildup was considerably larger for two- than for four-choice (the y-intercept for two-choice was 50.7 ± 8.2 spikes s^{-2} greater than the y-intercept for four-choice, $P < 10^{-5}$; Fig. 4e,f; see Discussion and Supplementary Fig. 6 online). This time-dependent rise is not explained by random fluctuations in motion energy, as the increase was evident when trials ending in T_{in} and T_{out} choices were averaged together (as in Fig. 4a–d). We interpret this rise as a reflection of the cost associated with the passage

of time, corresponding to the psychological sense of urgency, $u(t)$ (refs. 17,18). The positive buildup rates at all motion strengths and for all directions effectively impose a deadline on the decision process, a deadline that is imposed earlier when there are just two choices.

The 90° control task ($n = 29$) allowed us to ascertain whether the differences in LIP responses on the two- and four-choice tasks are explained by differences in the number of choices *per se* or the difference in angle between directions of motion. Recall that before motion onset, activity on the 90° control trials was slightly reduced in comparison with the standard two-choice task. This reduction became more modest throughout the dip during the beginning of motion (two-choice responses were 2.66 ± 2.0 spikes s^{-1} higher than 90° responses, $P = 0.10$), followed by a buildup in firing rates leading to the decision.

The effect of motion strength on these buildup rates was similar to two-choice responses for motion toward T_{in} (for the population, slopes differed by 0.23 ± 0.32 spikes $s^{-2} \%coh^{-1}$, $P = 0.47$; Fig. 4g). For motion toward T_{90} , however, a 1% increase in motion strength decreased buildup rate. This suggests that motion toward T_{90} constitutes weak evidence against the T_{in} choice. It also raises the possibility that the near absence of an effect of orthogonal motion on the four-choice task is an artifact that results from averaging the two orthogonal directions. Alternatively, the T_{90} direction may be more likely to contribute negatively to T_{in} choices without a T_{out} alternative.

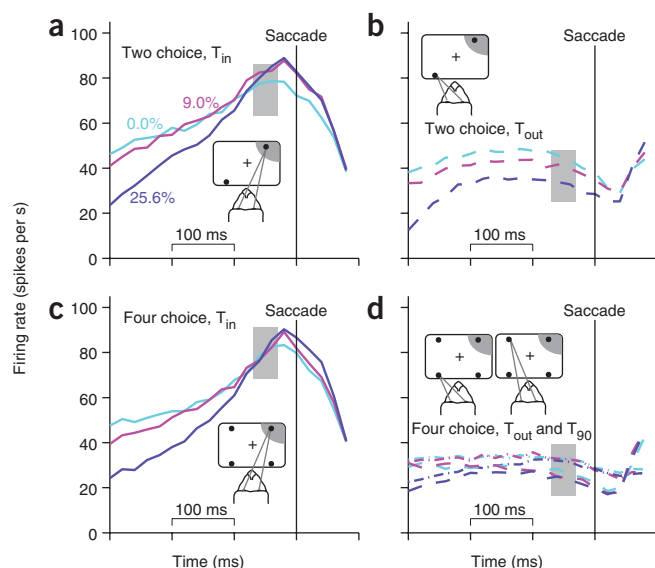


Figure 5 Neural responses just preceding the eye-movement responses. Population average firing rates ($n = 70$ neurons) are shown aligned to the initiation of saccades (vertical line). Trials are grouped on the basis of the direction of the saccade with respect to the response field of the neuron (insets). Only three motion strengths are shown for clarity. (a) T_{in} choices in the two-choice task. (b) T_{out} choices in the two-choice task. (c) T_{in} choices in the four-choice task. (d) T_{out} (dashed) and T_{90} (dot-dash) choices in the four-choice task.

Firing-rate excursion

Firing rates at the beginning and end of the motion-viewing period suggest that LIP neurons undergo a larger change in firing rate during decisions among four possible alternatives. When the monkey chose the target in the neuron's response field, firing rates for the two- and four-choice task ultimately reached a similar value near the time of the saccade (Fig. 2d and Supplementary Fig. 2). However, the initial response when the motion began was lower for the four-choice task than for the two-choice task (Fig. 3). We confirmed this by estimating the 'firing-rate excursion' from each neuron in the population (Fig. 6).

The combined difference in firing rates at the beginning and end of motion viewing was 6.11 ± 0.21 spikes s^{-1} larger for the four-choice task than for the two-choice task (see Methods, equation (2)). This difference, present in individual neurons (for example, Fig. 6a), was highly significant across the population ($P < 0.0005$; Fig. 6b,c) and in both monkeys individually (Monkey I: excursion difference = 4.7 ± 0.27 spikes s^{-1} , $P < 0.02$; Monkey S: difference = 11.9 ± 1.0 spikes s^{-1} , $P < 0.005$). The magnitude of this difference depended on which interval we used to estimate firing rates at the end of the trial, but the effect was statistically significant for a wide range (Supplementary Fig. 2). These estimates were obtained using only correct trials, but we obtained a similar estimate when we included all of the trials made to the T_{in} target.

The difference in excursion is attributed to the number of choices. The firing-rate excursion on the 90° control task was nearly identical to that seen on the standard two-choice task (mean difference = 0.13 ± 0.37 spikes s^{-1} , $P = 0.53$; Fig. 6d). The change in excursion in the four-choice task was also confirmed in this subset of neurons (mean difference = 4.2 ± 2.3 spikes s^{-1} larger for four choice than for two choice, $P < 0.04$). Thus, the angular separation between choice targets is, by itself, insufficient to cause an increase in firing-rate excursion of the magnitude seen on the four-choice task.

Relationship between LIP activity and behavior

The preceding analyses exposed the similarities and differences in how LIP activity reflects a bounded accumulation of evidence during two- and four-choice decisions. We next attempted to relate these

As before, some of the buildup in firing rate on the 90° control task is not the result of an accumulation of motion evidence, as it is present even on the trials with 0% motion strength. The magnitude of this urgency signal, $u(t)$, fell between those measured on the two- and four-choice tasks (Table 1 and Supplementary Fig. 6).

Responses late in the motion epoch

On a variety of two-choice decision tasks, a threshold crossing is associated with termination of the decision^{1,2,19}. We found evidence for a threshold on the four-choice task as well (Fig. 2d). We asked whether this threshold differs for two- and four-choice tasks.

Firing rates just preceding T_{in} choices were similar for two- and four-choice trials (Fig. 5a,c; responses in shaded region were on average only 3.1 ± 1.3 spikes s^{-1} higher for four choice than for two choice, $P < 0.03$). Although significant, this difference amounts to only 3.7% of the firing rate in this epoch. Furthermore, unlike the buildup rates at the beginning of the decision, the firing rates at the end of decisions for T_{in} did not vary as a function of motion strength ($P > 0.57$ for two and four choice) and were the same for correct and error trials ($P > 0.78$ for two and four choice). We conclude that the termination of decisions is associated with a fixed firing rate when the monkey chooses the target in the response field.

Firing rates just preceding T_{out} and T_{90} choices were harder to interpret. Firing rates tended to be lower on the four-choice task (difference = 9.81 ± 4.08 spikes s^{-1} , $P < 0.02$). Because firing rates were already lower at the beginning of these trials, the persistence of this difference is only notable in contrast with the T_{in} choices. What remains unclear is whether the firing rates achieve a stereotyped level before the saccade or whether they reflect the motion strength and reaction time. We observed a weak dependence of firing rate on motion strength, but the effects were not statistically reliable ($P > 0.07$). In the context of bounded accumulation, a weak relationship between motion strength and firing rate raises the possibility that the losing decision process (or processes) conveys information about the degree of difficulty²⁰.

Table 1 Model parameters

	Estimated from neural recordings		Fitted parameters		
	Bound (spikes s^{-1})	Urgency ($\tau_{1/2}$, u_∞) (ms, spikes s^{-1})	Nondecision time (T_D) (ms)	K (spikes s^{-2} %coh $^{-1}$)	σ (spikes $s^{-1.5}$)
$n = 70$					
2 choice	48.6	133.2, 34.7	205.2 ± 1.0	222.5 ± 2.5	35.2 ± 0.15
4 choice	54.7	343.2, 39.0	141.3 ± 0.3	165.0 ± 0.2	27.3 ± 0.001
$n = 29$					
2 choice	55.9	224.4, 60.2	191.0 ± 2.6	189.5 ± 3.9	26.1 ± 0.28
4 choice	60.0	402.3, 49.4	134.8 ± 0.04	177.0 ± 0.3	26.5 ± 0.01
90° control	55.7	265.7, 49.2	169.5 ± 1.0	361.5 ± 1.4	31.2 ± 0.05

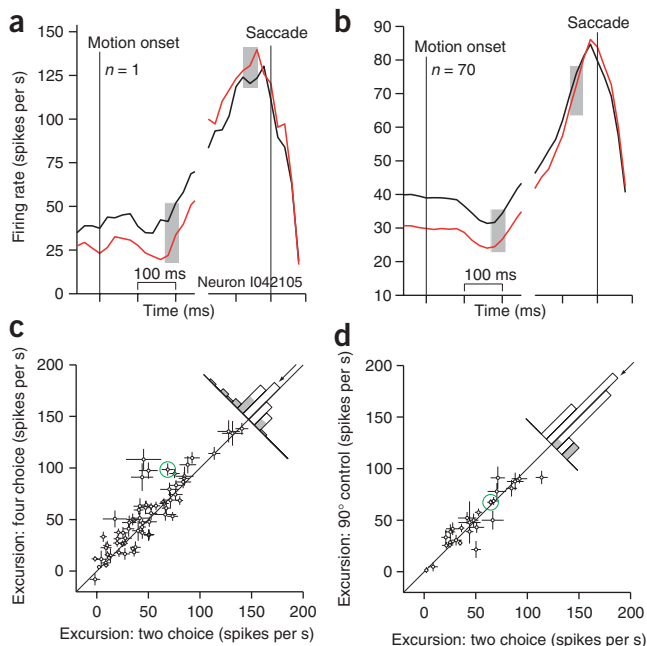


Figure 6 Firing-rate excursion is larger on the four-choice task. **(a)** Firing rates on two- and four-choice trials at the beginning and end of the motion-viewing period for a single example neuron. Responses are aligned to motion onset (left) and saccade initiation (right). Black and red traces indicate responses on the two- and four-choice tasks, respectively. Traces are smoothed with a 30-ms exponential filter. Excursion is the difference between firing rates in the shaded regions. All responses leading to correct T_{in} choices and with reaction time > 450 ms were used in this analysis. Firing-rate excursion was 29.7 ± 8.4 spikes s^{-1} larger for four choice than for two choice. **(b)** As in **a**, except that traces reflect the average firing rate from 70 neurons and no smoothing was performed; firing rates were computed in 20-ms nonoverlapping bins. **(c)** Comparison of firing rate excursion on two- and four-choice tasks. Points are estimates of the firing-rate excursion from single neurons. One point (170, 159) was omitted from the scatter plot to facilitate scaling of the remaining points. Error bars show standard error of the excursion (equation (2)) and are occasionally obscured by the points. Green circle marks the example neuron in **a**. Histogram depicts the differences in excursion on two- and four-choice tasks. Arrow indicates the mean. Only correct responses to T_{in} targets were used for this analysis. Gray shading indicates individual neurons with significant differences ($P < 0.05$). **(d)** Comparison of firing rate excursion on two-choice and 90° control tasks. The same conventions were used as in **b**. The same neuron (170, 159) was omitted from the scatter plot.

observations to the behavioral measurements of choice and reaction time. We have already noted that firing rates build up faster in association with strong motion and fast reaction time. Moreover, this buildup begins at a fixed time after the onset of motion and signals the end of decision (for T_{in}) by achieving a critical level. Here, we provide two additional analyses in support of this mechanism.

First, we examined the relationship between buildup rate and reaction time for single trials that terminated with a T_{in} choice (Fig. 7). For each neuron, the single-trial estimates of buildup rate and the measured reaction time were detrended to remove the effect of motion strength (see Methods) and combined in a scatter plot (Fig. 7a,b). Trial-to-trial variability in buildup rate was inversely correlated with trial-to-trial variability in reaction time, despite the noisy estimate of buildup rate and large variation in reaction time. This observation was typical of the data (two choice: mean correlation coefficient = -0.22 ± 0.02 , $P < 10^{-5}$, $n = 70$ neurons; four choice: mean correlation coefficient = -0.26 ± 0.02 , $P < 10^{-5}$, $n = 69$; Fig. 7c,d).

Next, we attempted to reconcile LIP activity with the speed and accuracy of the

monkeys' decisions (Fig. 1d–g). The excursion of LIP firing rates throughout the decision was larger on the four-choice task (Fig. 6). The change in LIP buildup rates produced by a change in motion strength toward T_{in} was similar for the two- and four-choice tasks (Fig. 4). The rise in firing rate that depends on time, but not on motion strength or direction ($u(t)$, urgency), was larger in the two-choice task (Supplementary Fig. 6). All of these observations are consistent with longer reaction times on the four-choice task; it takes longer for LIP firing rates to undergo the larger excursion. These observations have less intuitive effects on accuracy. For example, the larger excursion and less pronounced time-varying increase would lead to better accuracy on the four-choice task were it not for the larger number of choices.

Our model implements a race between two and four choice mechanisms to determine which target will be selected (Supplementary Fig. 6). Each mechanism accumulates momentary evidence in

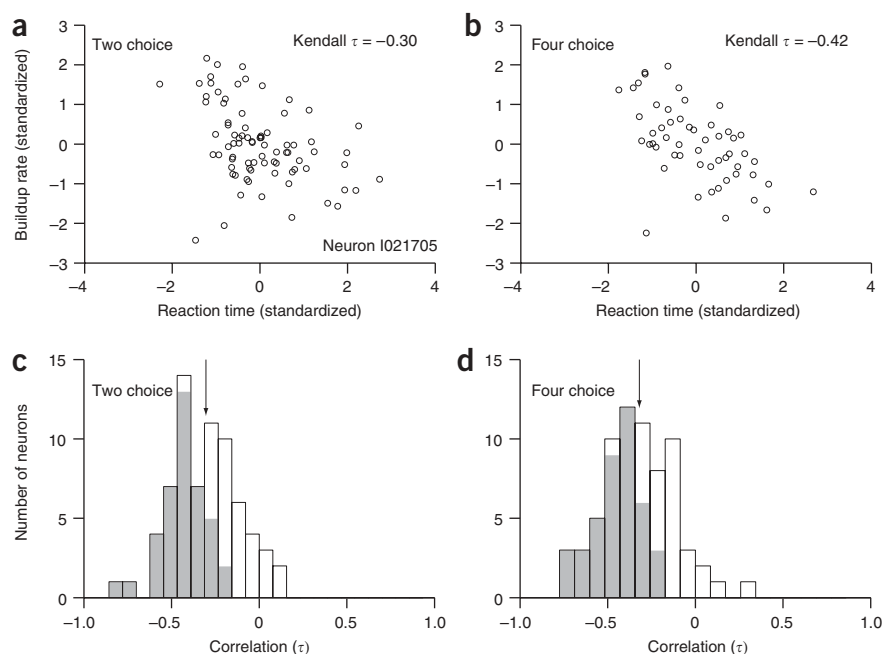


Figure 7 Neural responses and reaction times are inversely correlated on single trials. **(a)** Trial-by-trial correlation between reaction time and buildup rate for one representative neuron (Kendall, $\tau = -0.30$, $P < 10^{-4}$). Values are expressed in units of s.d. from mean. This detrending was performed for each motion strength using all correct T_{in} choices and all T_{in} choices for 0% coherence. **(b)** As in **a**, but for four-choice trials ($\tau = -0.42$, $P < 10^{-4}$). **(c,d)** Distribution of correlation coefficients (Kendall, τ) for each neuron in the dataset for the two-choice **(c)** and four-choice tasks **(d)**. Gray shading indicates individual neurons with significant correlation ($P < 0.05$). Arrow indicates the mean.

favor of the choice target in its response field. A decision is reached when one of the accumulators reaches a bound. We allowed the monkeys' physiology to constrain bound height and $u(t)$. Three additional parameters were adjusted to fit the choice and reaction-time data (see Methods and **Table 1**). We are cautious about interpreting the values of the fitted parameters, as we have fixed the most important model parameters using the LIP measurements. The main point is that they are reasonably similar for the two- and four-choice tasks.

The model provides an adequate account of both the reaction time and accuracy seen on the two- and four-choice tasks (**Fig. 1d–g**). This fit supports the hypothesis that the framework of bounded accumulation can be extended to explain decisions with more than two alternatives. Moreover, the physiological differences on the two- and four-choice tasks are of the right size to explain the behavioral differences that we observed. The model also accounts for the longer reaction times that we observed on error trials for both the two- and four-choice task (**Supplementary Fig. 7** online). The model, especially the hybrid of fixed and free terms, is not unique, but underscores the consistency of physiological and behavioral measurements in the framework of bounded accumulation.

DISCUSSION

Simple experimental models that build on knowledge about sensory and motor functions (reviewed in ref. 1) have the downside that they rely on mechanisms that may be inadequate for more complex decisions. Relative to two-choice decisions^{2,8,9,21}, our four-choice task doubles the number of alternatives and halves the angular separation between directions. Although this angle remains large compared with the fine discriminations that humans and monkeys can make^{22–24}, it did increase difficulty slightly, as shown by 90° separation in a two-choice task. Longer viewing durations were required for the same accuracy (**Fig. 1f,g**).

To achieve an acceptable reward rate, we used a mixture of motion strengths that favored easier conditions, compared with previous choice–reaction time experiments (for example, see ref. 2). The set was identical in all conditions, so the two-choice task was easy, probably explaining the faster reaction times that we measured. The monkeys could afford to terminate choices using a more lax criterion, what we interpret as a shorter excursion to a termination bound, without increasing the average probability of error.

Many models have been proposed to explain the coordination of reaction time and choice accuracy in simple two-choice tasks^{4,18,25–27}. These models share a common feature: a termination bound is applied to some form of evidence representation, termed the decision variable. A putative neural correlate of this operation is evident in area LIP; for all reaction times (and motion strengths), the firing rates of LIP neurons reached a stereotyped level shortly before the monkey initiates a saccade to indicate its choice for T_{in} (**Fig. 2b,d** and **Supplementary Fig. 2**). Our findings suggest that this same bounded accumulation framework applies to decisions among four choices. This hypothesis has been suggested by theorists^{5,28,29}, but has not been tested until now.

A major difference between two- and four-choice responses was seen in the firing rate at the beginning of the decision process (**Fig. 3**). This effect resembles the inverse relationship between the number of possible eye movement directions and firing rates in the superior colliculus preceding a saccade instruction³⁰. We think that the reduction in firing rate on a four-choice response is unlikely to be the result of a low-level factor, such as the presence of a target in the neuron's inhibitory surround. This possibility was refuted in the superior

colliculus by dissociating uncertainty from the number of targets. We tried to place the monkey in a two-choice task while providing four targets (two were irrelevant), but this manipulation introduced unacceptable biases in subsequent testing. Nonetheless, we doubt that surround suppression explains the firing-rate reduction. First, no inhibitory surround is seen in LIP response fields^{31,32}. Second, responses to the T_{90} target on our screening saccade task were not lower than responses to the T_{out} target, as would be expected if they were in an inhibitory surround (**Supplementary Fig. 1**). Third, in two experiments, we observed a reduction in firing rate on the four-choice task even when the response fields were large enough to encompass the orthogonal targets in their excitatory regions (data not shown).

In our four-choice task, this lower firing rate at the decision's beginning conferred a higher threshold for terminating the decision. Three experimental findings support this conclusion. First, the firing rate at the decision's end was similar in the two- and four-choice configurations. Thus, the lower starting point implies a larger excursion in firing rate from the start to the end of the decision. Second, the rate of buildup in LIP that can be attributed to motion strength was similar on the two- and four-choice configurations for motion toward T_{in} (**Fig. 4**) and for T_{in} choices (**Supplementary Fig. 4**). This rules out the possibility that the difference in excursion was compensated for by a change in the scaling of accumulated evidence into units of LIP firing rate. Third, the urgency was stronger in two- than in four-choice conditions. This amplifies the effective difference in excursion between the two- and four-choice conditions because it effectively shrinks the excursion; less of the range of LIP firing rates between start and end is used to represent the accumulated evidence.

In the framework of bounded accumulation, a larger excursion improves signal to noise at decision termination, or equivalently, a reduction in uncertainty, at the expense of decision time. When human subjects carry out the two-choice motion task, they appear to adjust the bound (or excursion) when they trade off speed and accuracy³. When the number of alternatives is increased, all else being equal, the level of uncertainty is initially increased. Thus, it is not surprising that the brain would accrue more evidence before terminating the decision. The cost is decision time, but this is tempered by the implementation of a deadline that is imposed by the urgency signal.

We identified this urgency signal as the time-dependent rise in firing rate that is seen on all trials regardless of motion strength and direction. For example, when the evidence is neutral (0% motion strength), assuming no choice bias, the bounded accumulation model approximates an unbiased diffusion of evidence: a rising variance, but no change in the mean evidence. The urgency signal is the difference between the observed firing rates and this expectation from unbiased diffusion. It is an essential feature of the model used to explain choice and reaction time in these experiments. A time-dependent rise of firing rate in LIP and elsewhere has been interpreted as a representation of elapsed time in the form of a hazard rate or anticipation function^{12,33–35}. In the framework of bounded accumulation¹⁷, it causes decisions to terminate as time elapses, regardless of the evidence.

In the four-choice task, time is costly. Long decision times would improve performance above chance (25% correct), but not for the 0% motion-strength trials and not by much for the next weakest motion strength. It is therefore sensible to place time limits on the decision process. Similar reasoning applies to the two-choice configuration. Because chance represents 50% correct, there is less impetus to complete the trial at weak motion strengths, which probably explains the weaker urgency signal seen previously in two-choice experiments^{2,36}. Here, however, our easy set of motion strengths rendered fast decisions less costly in the two-choice task.

One question that remains is how signals from direction-selective neurons in the middle temporal area are combined into momentary evidence. Observations from the standard two-choice task suggest that LIP neurons reflect the difference in firing rates between neurons tuned to the two opposing directions of motion²¹. This oppositional relationship was demonstrated directly by stimulating neurons in the middle temporal area and measuring the effect on choice and reaction time. Buildup rates in LIP reflect, at least in part, this subtraction; a change in motion strength in the direction favoring T_{in} leads to a proportional rise in buildup rate in LIP, and an opposite change in motion strength leads to a comparable decline. Our results suggest that the computation that LIP performs on the motion signals that it receives from the middle temporal area (and elsewhere) is similar in the two- and four-choice tasks. In the latter, the contribution of the two nonopposing directions (T_{90}) was small (Fig. 4f), but we doubt that this will generalize to more directions and choices. Our experiment is not definitive, in part because the angle between alternatives is large compared with the range of angles that primates can discriminate^{22–24}.

Our study suggests that some of the principles gleaned from simple binary decisions probably extend to decisions with more than two choices. Specifically, we observed a neural correlate of evidence accumulation in structures that represent the choice. In addition, the observed bound on the accumulation ties the content of a decision with the time required to make it together in a single mechanism. This idea also receives support from a recent study in human psychophysics³⁷. It remains to be seen whether the framework will extend to decisions among more than four choices^{28,38,39} or to decisions among a continuous array of choices, in which a parameter can take on any value, like the heading of a compass⁴⁰. It may seem that, according to the bounded accumulation framework, decisions among continuous choices would require an infinite number of accumulators. This framework may simply not be able to account for decisions among continuous choices, and may need to be replaced with a more suitable model^{27,41}. Alternatively, decisions among continuous choices might be explainable using a finite number of accumulators. The resolution of cognitive and motor systems may not require that the brain represent an infinite array of possibilities with precision; the degree of uncertainty reduction need not exceed that of the systems that use the estimate. If so, a bounded accumulation framework with a finite number of accumulators may be sufficient, even for decisions among continuous choices.

METHODS

Behavioral tasks. Stimuli were shown on a CRT monitor with a refresh rate of 99 Hz positioned 59 cm from the monkeys. Stimuli were generated using Matlab 5.2 (Mathworks) and the Psychophysics toolbox⁴².

For the random dot motion task, trials were similar to those reported elsewhere^{2,8,36} (Fig. 1a–c). Following successful fixation ($\pm 1.5^\circ$) of a central white target, two or four highly visible, red peripheral choice targets appeared (diameter = 0.5°). Choice targets were 180° apart (two-choice task) or 90° apart (four-choice task and 90° control task) and all choice targets were equally eccentric (range = $4\text{--}15^\circ$, mean = 10.5°). Two- and four-choice trials were randomly interleaved. The dynamic random dots were displayed in a 5° circular aperture centered on the fixation point. We used a variable coherence random dot display^{2,8,43} (dot density = 16.7 dots per deg^2 per s). Coherently moving dots were displaced to produce 6° s^{-1} motion. For the first 20 neurons examined, six motion strengths were used (0%, 3.2%, 6.4%, 25.6%, 51.2% and 76.8%). For the remaining 50 neurons, only five motion strengths (0%, 3.2%, 9.0%, 25.6% and 72.4%) were used, so as to maximize the number of trials per condition collected. For graphs, combined data from the highest motion strengths are shown at 72.4% coherence.

The motion stimulus was extinguished when the monkey's gaze exited the fixation window. Saccades ending in a window (range, $\pm 1.5^\circ$ to $\pm 3.5^\circ$

depending on eccentricity) around the choice target corresponding to the correct direction of stimulus motion were rewarded with juice or water. Otherwise, no reward was given and the correct choice target briefly doubled in size to provide feedback. For Monkey S, a delay of 1,000 ms was imposed after incorrect trials before the next trial began to discourage fast choices.

We also used overlap and memory saccade tasks. In the overlap saccade task, a peripheral target was placed in the visual field, while the monkeys maintained fixation of a central spot during a delay period. The delay period was random, sampled from a truncated exponential distribution. When the central spot was extinguished, the monkeys made a saccade to the choice target. In the memory saccade task, the peripheral target was flashed only briefly and the monkeys made a saccade to its remembered location. Memory and overlap saccade tasks were carried out before the discrimination task to verify the spatial selectivity of the neuron; trials using the overlap saccade task were usually randomly interleaved with motion discrimination trials to test for stability of the response field and the neural responses (Supplementary Fig. 1).

Electrophysiology. Two monkeys (I, 51 neurons; S, 19 neurons) were prepared for chronic single-neuron recording using standard surgical procedures, as described previously^{2,8}. All surgical and experimental procedures were in accordance with the US National Institutes of Health Guide for the Care and Use of Laboratory Animals and were approved by the University of Washington Animal Care Committee.

Electrodes (Alpha Omega) were introduced into area LIP at each daily recording session. Neural responses were amplified conventionally, and waveforms corresponding to different neurons were sorted online using the Plexon Sort Client (Plexon). The Plexon Offline Sorter was used after each experiment to confirm that all recorded spikes came from single well-isolated neurons, determined by waveform shape and the presence of a refractory period.

Neurons were selected according to anatomical and physiological criteria. Anatomical landmarks were provided by magnetic resonance images of the monkeys' brains that were compared to cortical partitioning schemes using Caret software⁴⁴. These observations were confirmed by physiological observations of white matter, gray matter and lumen crossings. Neurons included in the study all had spatially selective responses during the delay on the overlap and memory saccade tasks⁴⁵ (Supplementary Fig. 1).

For 70 neurons, we recorded responses on an interleaved block of two- and four-choice trials. For 29 of these neurons, responses were collected on a second block of trials that contained 90° control trials interleaved with either two- or four-choice trials.

Data analysis. We measured firing rates in several time windows defined with respect to the task epochs: before motion, early motion and before saccades. For the pre-motion epoch, firing rates were obtained 200–300 ms after the onset of the choice targets. An early window (170 to 210 ms after motion onset) and a late window (40–80 ms before saccade initiation) provided estimates of the firing rates at the beginning and at the end of the decision process (Figs. 3 and 6). These points were established from an analysis of firing-rate variance associated with the different motion stimuli and reaction times (Supplementary Fig. 2). For both the two- and four-choice task, firing rate variance remained at a stable, low value until 190 ms after the onset of stimulus motion, after which it increased markedly, reflecting the divergence of responses to different stimuli. We used a 40-ms window centered at this point, but the estimate of the firing rate was nearly identical for a variety of intervals. The end of the evidence accumulation was taken to be the point at which the variance was lowest just before the saccade (Supplementary Fig. 2), as in previous studies². We used a 40-ms time window centered on the point corresponding to minimal variance. For analyses of firing rates around the saccade, the first 200 ms of the response to the dot motion were removed to exclude the stereotyped dip that was time-locked to the onset of the random dot display (Fig. 2c). For the analysis of firing-rate excursion and firing rate at the end of the decision, trials with reaction times shorter than 450 ms were excluded.

A wide variety of interval definitions yielded nearly identical results, with one exception. Estimates of firing rate at the end of evidence accumulation were sensitive to the interval used because the firing rates changed markedly near the

time of the saccade. However, the effect of choice number on firing-rate excursion that we report here was present for a range of time intervals, centered up to 130 ms before the saccade (Supplementary Fig. 2). Two- and four-choice firing-rate excursions were more similar for intervals centered at longer time points before the saccade, but these earlier time points are unlikely to correspond to the end of evidence accumulation^{2,46}.

To estimate the buildup in firing rate during decision formation, we first constructed a peri-stimulus time histogram (10-ms bins) for each neuron from responses aligned to the onset of random dot motion, excluding the epoch beginning 60 ms before saccade initiation. The buildup rate is the slope of the line fit to a peri-stimulus time histogram in the epoch from 190–320 ms after motion onset using regression. We used an early epoch to include as many trials as possible in the averages, minimize attrition of portions of trials from the averages (preceding the saccade) and minimize the potentially confounding effect of choice on the stimulus-dependent buildup rate. Analyses employing different time intervals affected the reported values of buildup rate only slightly; the largest effect was on buildup rates corresponding to the highest motion strength, owing mainly to the attrition of trials with short reaction times. Data from the 3.2% and 6.4% coherences were combined, as were data from the 72.4% and 76.8% coherences so that population averages would not be skewed by values associated with motion strengths that were only used on a minority of days.

On the four-choice task, the buildup rates for orthogonal motion were generated from combined responses to the two orthogonal motion directions used. We obtained similar results whether buildup rates were from individual neurons (as described above) or were computed from the average population response (data not shown).

Population analyses that group trials with a common motion direction (relative to T_{in}), as well as the estimate of $u(t)$, could be affected by a choice bias. We therefore analyzed the distribution of choices made on the 0% coherence motion trials. No reliable bias was found. For two-choice trials, the numbers of T_{in} and T_{out} choices were 1,452 versus 1,387 ($P = 0.22$, binomial distribution, $H_0: p = 0.5$). For four-choice trials, the numbers of T_{in} , T_{out} , T_{+90} and T_{-90} choices were 1,461, 1,421, 1,387 and 1,426 ($P = 0.65$, χ^2 test, H_0 : equal proportions).

To estimate the relationship between buildup rate during decision formation and reaction time, we analyzed single trials (Fig. 7), using weak motion strengths (0–25.6% coherence) leading to correct T_{in} choices (all T_{in} choices for 0% coh). We estimated the firing rate on each trial by convolving the spike train with an alpha-like function $(1 - e^{-t/d}) \cdot e^{-t/d}$, where $d = 25$ ms and $g = 1$ ms. The buildup rate is the slope of the best fitting line to the smoothed firing rate in the epoch from 190 ms after motion onset to 120 ms before saccade initiation. The buildup rates and the associated reaction times were standardized for each motion strength (difference from the mean in units of sample s.d.). The standardized buildup rates and reaction times were combined for all coherences to estimate a correlation coefficient (Kendall, τ) for each neuron. We required that at least three spikes be present in a trial to estimate buildup rate and that at least five trials be present in a given neuron to estimate the correlation coefficient. One neuron was excluded on the four-choice task for failing to meet these criteria. After estimating the correlation coefficient for each neuron, we tested whether the mean of the population was significantly less than 0 (t -test).

We used regression analyses to estimate the effect of the number of choices on the LIP firing rates⁷. Unless otherwise stated, all fitting uses maximum likelihood under Gaussian assumptions for noise (that is, weighted least-squares regression with known variance); standard errors of parameter estimates were obtained by inverting the Hessian matrix of derivatives of the log likelihood with respect to the parameters, evaluated at the maximum likelihood solution. For these regression analyses, tests of significance were determined by t -statistics computed from the estimated coefficients and their standard errors. For many analyses, single-neuron measurements were gathered in a frequency histogram (Figs. 3c,d and 6c,d; see also Supplementary Figs. 1, 4, and 5), on which we used a t -test (effectively a paired t -test on the associated scatter plots). For all analyses, we also used maximum likelihood to generate population means and associated t -statistics. These two methods sometimes yielded slightly different estimates, but the direction and significance of all of the results reported here were always in agreement.

For simple comparisons of firing rates in a specified epoch, the regression is effectively a t -test,

$$y = \beta_0 + \beta_1 I_n, \quad (1)$$

where $I_n = 0$ or 1 for two and four choice, respectively, and y is the spike rate on individual trials. β_i are fitted coefficients. The null hypothesis is that choice number does not affect the firing rate ($H_0: \beta_1 = 0$).

To compare firing-rate excursion for two versus four choice, we fit the model:

$$y = \beta_0 + \beta_1 I_n + \beta_2 I_T + \beta_3 I_n I_T \quad (2)$$

where the β_i are fitted coefficients, y is the spike rate measured in the beginning and late epochs (defined above) and the I_x are indicator variables, $I_n = 0$ or 1 for two and four choice, respectively, and $I_T = 0$ or 1 for early and late epochs, respectively. The coefficients estimate the following parameters: β_0 is the firing rate for the two-choice condition in the early epoch, $\beta_0 + \beta_1$ is the firing rate in the four-choice condition in the early epoch, $\beta_0 + \beta_2$ is the firing rate in the two-choice condition in the late epoch and $\beta_0 + \beta_1 + \beta_2 + \beta_3$ is the firing rate in the four-choice condition in the late epoch. β_2 furnishes an estimate of the firing-rate excursion in the two choice and β_3 estimates the difference in excursion on the four-choice task. The null hypothesis that the number of choices does not affect the excursion is $H_0: \beta_3 = 0$.

To compare responses between the two- and four-choice tasks during the motion epoch, we computed the relationship between motion strength and buildup rate using a weighted regression,

$$y = \beta_0 + \beta_1 I_n + \beta_2 C + \beta_3 I_n C \quad (3)$$

where y is the buildup rate either averaged across all the neurons in the population (Fig. 4f,g) or for a single neuron (Fig. 4e and Supplementary Fig. 5) for each motion strength, I_n is an indicator for the number of targets (as above) and C is motion strength (% coherence). The buildup rates associated with 0% coherence motion are furnished by β_0 and $\beta_0 + \beta_1$ for two- and four-choice tasks; β_2 and $\beta_2 + \beta_3$ provide estimates for the effect of a change in motion strength on the buildup rates. The null hypothesis is that the number of choices does not affect this relationship ($H_0: \beta_3 = 0$).

Eye movements. Eye position was sampled at 1 kHz using the scleral search-coil method⁴⁷, as described previously². To identify differences in the oculomotor responses on two-choice versus four-choice trials, we compared peak saccade velocity, saccade amplitude and saccade error on each condition. Some of these measures were affected by the number of choices^{48,49} (Supplementary Fig. 8 online). To test whether these factors explained (as potential confounders) the effect of choice number on LIP responses, we incorporated the three significant parameters into the regression described above (equation (1) and (2)):

$$y = \beta_0 + \beta_1 I_n + \alpha_1 S + \alpha_2 \dot{S} + \alpha_3 E \quad (4)$$

$$y = \beta_0 + \beta_1 I_n + \beta_2 I_T + \beta_3 I_n I_T + \alpha_1 S + \alpha_2 \dot{S} + \alpha_3 E$$

where y is the spike rate measured on each trial either at the beginning or at the end of the decision for all trials in all cells (that is, each trial contributes two measures), α_i are additional fitted coefficients, and S , \dot{S} and E are amplitude, peak velocity and endpoint error, respectively. The null hypotheses were re-evaluated with these extra terms in the regression. The null hypothesis, that the number of choices did not affect the excursion, was again rejected both at the population level ($P < 0.01$) and in all but one of the individual cells where the effect of firing-rate excursion was significant. In this cell, firing-rate excursion was found to be longer for two choice than for four choice, an exception to the trend that we ordinarily observed.

Diffusion model. We fit behavioral data using an accumulator model (Table 1 and Supplementary Fig. 6). An accumulator corresponding to each target integrates the momentary evidence toward a decision bound. We assume the evidence is a linear combination of middle temporal area activity for motion toward the targets. The linear weights were taken from the cosine of the angle difference between the directions²². The bound height and the motion-independent signal $u(t)$ were obtained from the neural responses to the two- and four-choice tasks, as explained below. Three free parameters

remained to be fit: K , a scaling parameter that converts motion strength to momentary evidence (that is, the buildup rate if $u(t)$ were absent), σ , the s.d. of this momentary evidence (diffusion coefficient), and T_0 , the mean nondesideration time.

The fit maximized the likelihood of the observed choices and reaction times for each dataset (two or four choice; the model performed only slightly worse when T_0 was constrained to be the same value for two- and four-choice conditions). These likelihoods were calculated directly using established numerical solutions⁵⁰ to the partial differential equations (Fokker-Planck) describing diffusion to a nonstationary bound, $B - \hat{u}(t)$, where B is the initial bound height and $\hat{u}(t)$ is an estimate of the urgency signals (see below). For the two-choice task, the race between two choice mechanisms is equivalent to one-dimensional diffusion with two symmetric, but collapsing, bounds. For four choice, the four races reduce to a race between two of the two-choice diffusion processes. The model with urgency explains the longer reaction times on error trials³⁶ (Supplementary Fig. 7).

The urgency signals, $u(t)$, were estimated from neural responses to 0% coherent motion. We first averaged responses to T_{in} and T_{out} trials separately. A grand average of these averages should be flat if LIP reflects only accumulated evidence. $u(t)$ is the time-varying signal that would need to be subtracted to produce this flat response (Supplementary Fig. 6). The urgency signals for each condition were parameterized separately with a hyperbolic function,

$$\hat{u}(t) = u_{\infty} \frac{t}{t + \tau_{1/2}},$$

where u_{∞} is the maximum and $\tau_{1/2}$ is the time to reach 50% of the maximum. The fitted values are listed in Table 1 for each condition. In the model, this parameterized function was used to add a time-varying signal to the accumulated evidence. Parameters were optimized separately for behavioral responses taken in conjunction with all the recorded neurons (Table 1 and Fig. 1d,e) and for those taken in conjunction with the 29 recording sessions when the 90° control was also presented (Table 1 and Fig. 1f,g).

Note: Supplementary information is available on the Nature Neuroscience website.

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- Gold, J.I. & Shadlen, M.N. The neural basis of decision making. *Annu. Rev. Neurosci.* **30**, 535–574 (2007).
- Roitman, J.D. & Shadlen, M.N. Response of neurons in the lateral intraparietal area during a combined visual discrimination reaction time task. *J. Neurosci.* **22**, 9475–9489 (2002).
- Palmer, J., Huk, A.C. & Shadlen, M.N. The effect of stimulus strength on the speed and accuracy of a perceptual decision. *J. Vis.* **5**, 376–404 (2005).
- Link, S.W. & Heath, R.A. A sequential theory of psychological discrimination. *Psychometrika* **40**, 77–105 (1975).
- Laming, D.R.J. *Information Theory of Choice-Reaction Times* (Wiley, New York, 1968).
- Mazurek, M.E., Roitman, J.D., Ditterich, J. & Shadlen, M.N. A role for neural integrators in perceptual decision making. *Cereb. Cortex* **13**, 1257–1269 (2003).
- Shadlen, M.N. & Newsome, W.T. Neural basis of a perceptual decision in the parietal cortex (area LIP) of the rhesus monkey. *J. Neurophysiol.* **86**, 1916–1936 (2001).
- Hanks, T.D., Ditterich, J. & Shadlen, M.N. Microstimulation of macaque area LIP affects decision-making in a motion discrimination task. *Nat. Neurosci.* **9**, 682–689 (2006).
- Huk, A.C. & Shadlen, M.N. Neural activity in macaque parietal cortex reflects temporal integration of visual motion signals during perceptual decision making. *J. Neurosci.* **25**, 10420–10436 (2005).
- Hick, W.E. On the rate of gain of information. *Q. J. Exp. Psychol.* **4**, 11–26 (1952).
- Lewis, J.W. & Van Essen, D.C. Corticocortical connections of visual, sensorimotor and multimodal processing areas in the parietal lobe of the macaque monkey. *J. Comp. Neurol.* **428**, 112–137 (2000).
- Leon, M.I. & Shadlen, M.N. Representation of time by neurons in the posterior parietal cortex of the macaque. *Neuron* **38**, 317–327 (2003).
- Sato, T. & Schall, J.D. Pre-excitatory pause in frontal eye field responses. *Exp. Brain Res.* **139**, 53–58 (2001).
- Kiani, R., Hanks, T.D. & Shadlen, M.N. Bounded integration in parietal cortex underlies decisions even when viewing duration is dictated by the environment. *J. Neurosci.* **28**, 3017–3029 (2008).
- Cook, E.P. & Maunsell, J.H. Dynamics of neuronal responses in macaque MT and VIP during motion detection. *Nat. Neurosci.* **5**, 985–994 (2002).
- Ratcliff, R. *et al.* Dual diffusion model for single-cell recording data from the superior colliculus in a brightness-discrimination task. *J. Neurophysiol.* **97**, 1756–1774 (2007).
- Reddi, B.A. & Carpenter, R.H. The influence of urgency on decision time. *Nat. Neurosci.* **3**, 827–830 (2000).
- Reddi, B.A., Asrress, K.N. & Carpenter, R.H. Accuracy, information, and response time in a saccadic decision task. *J. Neurophysiol.* **90**, 3538–35346 (2003).
- Hanes, D.P. & Schall, J.D. Neural control of voluntary movement initiation. *Science* **274**, 427–430 (1996).
- Vickers, D. *Decision Processes in Visual Perception* (Academic Press, London, 1979).
- Ditterich, J., Mazurek, M.E. & Shadlen, M.N. Microstimulation of visual cortex affects the speed of perceptual decisions. *Nat. Neurosci.* **6**, 891–898 (2003).
- Jazayeri, M. & Movshon, J.A. A new perceptual illusion reveals mechanisms of sensory decoding. *Nature* **446**, 912–915 (2007).
- Purushothaman, G. & Bradley, D.C. Neural population code for fine perceptual decisions in area MT. *Nat. Neurosci.* **8**, 99–106 (2005).
- Churchland, A.K. *et al.* Directional anisotropies reveal a functional segregation of visual motion processing for perception and action. *Neuron* **37**, 1001–1011 (2003).
- Ratcliff, R. & Rouder, J.N. Modeling response times for two-choice decisions. *Psychol. Sci.* **9**, 347–356 (1998).
- Luce, R.D. *Response Times: Their Role in Inferring Elementary Mental Organization* (Oxford University Press, New York, 1986).
- Lo, C.C. & Wang, X.J. Cortico-basal ganglia circuit mechanism for a decision threshold in reaction time tasks. *Nat. Neurosci.* **9**, 956–963 (2006).
- Usher, M. & McClelland, J.L. The time course of perceptual choice: the leaky, competing accumulator model. *Psychol. Rev.* **108**, 550–592 (2001).
- McMillen, T. & Holmes, P. The dynamics of choice among multiple alternatives. *J. Math. Psychol.* **50**, 30–57 (2006).
- Basso, M.A. & Wurtz, R.H. Modulation of neuronal activity in superior colliculus by changes in target probability. *J. Neurosci.* **18**, 7519–7534 (1998).
- Platt, M.L. & Glimcher, P.W. Response fields of intraparietal neurons quantified with multiple saccadic targets. *Exp. Brain Res.* **121**, 65–75 (1998).
- Ben Hamed, S., Duhamel, J.R., Bremmer, F. & Graf, W. Visual receptive field modulation in the lateral intraparietal area during attentive fixation and free gaze. *Cereb. Cortex* **12**, 234–245 (2002).
- Janssen, P. & Shadlen, M.N. A representation of the hazard rate of elapsed time in macaque area LIP. *Nat. Neurosci.* **8**, 234–241 (2005).
- Brody, C.D., Hernandez, A., Zainos, A. & Romo, R. Timing and neural encoding of somatosensory parametric working memory in macaque prefrontal cortex. *Cereb. Cortex* **13**, 1196–1207 (2003).
- Maimon, G. & Assad, J.A. A cognitive signal for the proactive timing of action in macaque LIP. *Nat. Neurosci.* **9**, 948–955 (2006).
- Ditterich, J. Stochastic models of decisions about motion direction: behavior and physiology. *Neural Netw.* **19**, 981–1012 (2006).
- Niwa, M. & Ditterich, J. Perceptual decisions between multiple directions of visual motion. *J. Neurosci.* **28**, 4435–4445 (2008).
- Salzman, C.D. & Newsome, W.T. Neural mechanisms for forming a perceptual decision. *Science* **264**, 231–237 (1994).
- Bogacz, R., Usher, M., Zhang, J. & McClelland, J.L. Extending a biologically inspired model of choice: multi-alternatives, nonlinearity and value-based multidimensional choice. *Philos. Trans. R Soc. Lond. B Biol. Sci.* **362**, 1655–1670 (2007).
- Nichols, M.J. & Newsome, W.T. Middle temporal visual area microstimulation influences verbal judgments of motion direction. *J. Neurosci.* **22**, 9530–9540 (2002).
- Ma, W.J., Beck, J.M., Latham, P.E. & Pouget, A. Bayesian inference with probabilistic population codes. *Nat. Neurosci.* **9**, 1432–1438 (2006).
- Brainard, D.H. The psychophysics toolbox. *Spat. Vis.* **10**, 433–436 (1997).
- Britten, K.H., Shadlen, M.N., Newsome, W.T. & Movshon, J.A. The analysis of visual motion: a comparison of neuronal and psychophysical performance. *J. Neurosci.* **12**, 4745–4765 (1992).
- Van Essen, D.C. Surface-based approaches to spatial localization and registration in primate cerebral cortex. *Neuroimage* **23** Suppl 1: S97–107 (2004).
- Gnadt, J.W. & Andersen, R.A. Memory related motor planning activity in posterior parietal cortex of macaque. *Exp. Brain Res.* **70**, 216–220 (1988).
- Thier, P. & Andersen, R.A. Electrical microstimulation suggests two different forms of representation of head-centered space in the intraparietal sulcus of rhesus monkeys. *Proc. Natl. Acad. Sci. USA* **93**, 4962–4967 (1996).
- Judge, S.J., Richmond, B.J. & Chu, F.C. Implantation of magnetic search coils for measurement of eye position: an improved method. *Vision Res.* **20**, 535–538 (1980).
- Arai, K., McPeck, R.M. & Keller, E.L. Properties of saccadic responses in monkey when multiple competing visual stimuli are present. *J. Neurophysiol.* **91**, 890–900 (2004).
- Walker, R., Deubel, H., Schneider, W.X. & Findlay, J.M. Effect of remote distractors on saccade programming: evidence for an extended fixation zone. *J. Neurophysiol.* **78**, 1108–1119 (1997).
- Risken, H. *The Fokker-Planck Equation: Methods of Solutions and Applications* (Springer, New York, 1989).

Corrigendum: Decision-making with multiple alternatives

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In the version of this article initially published, the panels in Figure 1 were mislabeled. The correct legend should be ‘Psychometric functions for the 29 experiments that included the 90° control are shown in e. Chronometric functions are shown in f’. This error has been corrected in the HTML and PDF versions of the article.