Eye Movements Reveal Spatiotemporal Dynamics of Active Sensing and Planning in Navigation

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Abstract

Goal-oriented navigation is widely understood to depend critically upon internal maps. Although this may be the case in many settings, humans tend to rely on vision in complex, unfamiliar environments. To study the nature of gaze during visually-guided navigation, we tasked humans to navigate to transiently visible goals in virtual mazes of varying complexity, observing that they took near-optimal trajectories in all arenas. By analyzing subjects’ eye movements, we gained insights into the principles of active sensing and planning. The spatial distribution of gaze revealed that environmental complexity mediated a striking trade-off in the extent to which attention was directed towards two complimentary aspects of the world model: reward location and task-relevant transitions. The temporal evolution of gaze revealed rapid, sequential prospection of the future path, evocative of neural ‘preplay’. These findings suggest that the spatiotemporal characteristics of gaze during navigation are significantly shaped by the unique cognitive computations underlying real-world, sequential decision making.
Introduction

Active sensing, the cognitively motivated process of gathering information from the surroundings, and planning, the evaluation of prospective future actions, are both thought to play a critical role in sequential decision making [1, 2]. Humans and animals have been shown to perform near-optimal active sensing during binary decision making tasks [3, 4] and visual search tasks [5–7]. Likewise, two-step choice tasks have revealed quantitative evidence that humans are capable of flexible planning [8–10]. However, the principles of active sensing and planning uncovered using simplified, discrete engagements may not generalize to complex behaviors like navigation in which our cognitive capacity is seriously limited by resource constraints. How can we study active sensing and planning during unstructured, naturalistic sequential decision-making ventures such as navigation?

Theoretical work suggests that information acquisition and navigational planning can be achieved through active inference – orienting the sensory apparatus to reduce uncertainty about task variables in the service of decision making [2]. Humans are fortuitously equipped with a highly evolved visual system to perform goal-oriented inference. By swiftly parsing a large, complex scene on a millisecond timescale, the eyes actively interrogate and efficiently gather information from different regions in space to facilitate complex computations [11, 12]. At the same time, eye movements are influenced by the contents of internal deliberation and the prioritization of goals in real-time [13–16]. Thus, eye tracking lends itself as a valuable tool for investigating both active sensing and planning [13, 17–19].

Over the past few decades, research on eye movements has led to a growing consensus that the oculomotor system has evolved to prioritize top-down, cognitive guidance over image salience [20–22]. During routine activities such as making tea, we tend to foveate specifically upon objects relevant to the task being performed (e.g. boiling water) while ignoring salient distractors [21, 23]. The consensus from research on eye movements is that gaze elucidates how humans use active sensing to mitigate uncertainty in a goal-oriented manner. Therefore, we hypothesize that in the context of navigation, gaze will be directed towards most informative regions of space depending on the specific relationship between the participant’s position and their goal. A candidate framework to formalize this hypothesis is Reinforcement Learning (RL), whereby the goal of behavior is cast in terms of maximizing total long-term reward [24]. For example, this framework has been previously used to provide a principled account of why neuronal responses in the hippocampal formation depend upon behavioral policies and environmental geometries [25, 26]. Incidentally, RL provides a formal interpretation of active sensing, which can be understood as optimizing exploration for the purpose of improving knowledge about the environment, allowing for better planning and ultimately greater long-term reward [13]. Here, we use this framework to quantitatively predict the precise spatial locations toward which eye movements should be directed, given the objective of navigating to a known destination.

Unlike active sensing, which manifests as overt behavior, planning is thought to be more covert [1]. From a normative standpoint, successful planning entails evaluating and comparing all possible paths leading to the goal. Researchers have proposed that certain neural codes, such as hierarchical representations, would support more efficient navigational planning by exploiting structural redundancies in the environment [27, 28]. While technological limitations have precluded the study of neural codes that underlie planning during naturalistic navigation in humans, there is evidence for predictive sequential neural activations during sequence learning and visual motion viewing tasks [29–31]. This is reminiscent of ‘preplay’, a well-documented phenomenon in rodents during navigation [32–35]. The mechanism by which humans perform navigational planning may similarly involve simulating sequences of actions that chart out potential trajectories. As recent evidence shows that eye movements reflect dynamics of internal beliefs during sensorimotor tasks [36], we hypothesize that participants’ gaze dynamics would rapidly reflect sequential trajectory simulation, and thus reveal the strategies by which humans plan during navigation.

To test both hypotheses, we designed a virtual reality navigation task in which both cognitive processes could be investigated using human eye movements. Participants were asked to navigate to transiently visible targets using a joystick in unfamiliar arenas of varying degrees of complexity. We found that human subjects balanced foveating the hidden reward location with viewing highly task-consequential regions of
space both prior to and during active navigation, and that environmental complexity mediated a trade-off between the two modes of information sampling. The experiment also revealed that subjects' eyes indeed rapidly traced the trajectories which they subsequently embarked upon, with such sweeps being more prevalent in complex environments. Furthermore, participants seemed to decompose convoluted trajectories by focusing their gaze on one turn at a time until they reached their goal. Taken together, these results suggest that during naturalistic navigation, the spatiotemporal dynamics of gaze are significantly shaped by active sensing and planning, thus reflecting the unique cognitive demands of real-world sequential decision making.
Results

Humans use vision to efficiently navigate to hidden goals in virtual arenas

To study human eye movements during naturalistic navigation, we designed a virtual reality (VR) task in which subjects navigated to hidden goals in hexagonal arenas. As we desired to elicit the most naturally occurring eye movements, we used a head-mounted VR system with a built-in eye tracker to provide a full immersion navigation experience with few artificial constraints. Subjects freely rotated in a swivel chair and used an analog joystick to control their forward and backward motion along the direction in which they were facing (Figure 1a — left). The environment was viewed from a first-person perspective through an HTC Vive Pro headset with a wide field of view, and several eye movement parameters were recorded using built-in software.
Facilitating quantitative analyses, we designed arenas with a hidden underlying triangular tessellation, where each triangular unit (covering 0.67% of the total area) constituted a state in a discrete state space (Figure S1b). A fraction of the edges of the tessellation was chosen to be impassable barriers, defined as obstacles. Subjects could take actions to achieve transitions between adjacent states which were not separated by obstacles. As subjects were free to rotate and/or translate, the space of possible actions was continuous such that subjects did not report knowledge about the tessellation. Furthermore, subjects experienced a relatively high vantage point and were able to gaze over the tops of all of the obstacles (Figure 1a—right). On each trial, subjects were tasked to collect a reward by navigating to a random goal location drawn uniformly from all states in the arena. The goal was a realistic banana which the subjects had to locate and foveate in order to unlock the joystick. The banana disappeared 200 ms after foveation, and participants were instructed to press a button when they believed that they have arrived at the remembered goal location. Then, feedback was immediately displayed on the screen, showing subjects that they had received either two points for stopping within the goal state, one point for stopping in a state neighboring the goal state, or zero points for stopping in any other state. While subjects viewed the feedback, a new goal for the next trial was spawned without breaking the continuity of the task. In separate blocks, subjects navigated to fifty goals in each of five different arenas (Figure 1h). All five arenas were designed by defining the obstacle configurations such that the arenas varied in the average path length between two states, as quantified by the average state closeness centrality (Methods — Eq 2 Figure 1c). Lower centralities correspond to more complex arenas. One of the blocks involved an open arena that contained only a few obstacles at the perimeter, such that on most trials, subjects could travel in straight lines to all goal locations (Figure 1f — leftmost). On the other extreme was a maze arena in which most pairs of states were connected by only one viable path (Figure 1f — rightmost).

To quantify behavioral performance, we first computed the optimal trajectory for each trial using dynamic programming (Figure S1b). This technique uses two pieces of information — the goal location (reward function) and the obstacle configuration (transition structure) — to calculate an optimal value function over all states such that the value of each state is equal to the (negative) length of the shortest path between that state and the goal state (Figure 1d). The optimal policy requires that subjects select actions to climb the value function along the direction of steepest ascent, which would naturally bring them to the goal state while minimizing the total distance traveled. Figure 1c shows optimal (dashed) as well as behavioral (colored) trajectories from an example trial in each arena. Behavioral path lengths were computed by integrating changes in the subjects’ position in each trial. Subjects took near-optimal paths (i.e. optimal to within the width of the reward zone) on most trials (Figure 1f), scoring 74 ± 5% of the points across all arenas and stopping within the reward zone on 86 ± 4% of all trials (Figure S1f). However, participants occasionally took a suboptimal route (Figure 1f — second from right), or sometimes misremembered the goal location and took a near-optimal trajectory to a different location (Figure 1f — rightmost). The latter type constituted a majority (68 ± 13%) of the unrewarded trials — most of the unrewarded trajectories would...
have been near-optimal if the subjects’ stopping location was deemed to be the goal state (Figure S1H — left), which suggests that remembering the goal location was not straightforward. We quantified the degree of optimality by computing the ratio of observed vs. optimal path lengths to the subjects’ stopping location. Across all trials (rewarded and unrewarded), this ratio was close to unity (median = 1.06, interquartile range = 0.14), suggesting that subjects were able to navigate efficiently in all arenas (Figure 1j, gray). This was the case even in unrewarded trials, where the observed path lengths were within 3% of the optimal path length to the subjects’ stopping location (Figure S1I — right). Navigational performance was near-optimal from the beginning, such that there was no visible improvement with experience (Figure S1B). However, the fraction of rewarded trials decreased with increasing arena complexity ($r = 0.70$, $p = 9.3 \times 10^{-8}$), suggesting that the ability to remember the goal location is compromised in challenging environments (Figures 1I, light green).

In order to understand how subjects tackled the computational demands of the task, it is critical to break down each trial into three main epochs: search — when subjects sought to locate the goal, pre-movement — when subjects surveyed their route prior to utilizing the joystick, and movement — when subjects actively navigated to the remembered goal location (Figure 1H). On some trials, participants did not end the trial via button press immediately after stopping, but this post-movement period constituted a negligible proportion of the total trial time.

We compared the fraction of time spent in different epochs. Although subjects spent a major portion of each trial navigating to the target, the relative duration of other epochs was not negligible (mean fraction ± SD — search: 0.26±0.03, pre-movement: 0.11±0.04, movement: 0.63±0.05; Figure S1). There was considerable variability across subjects in the fraction of time spent in the pre-movement phase (coefficient of variation (CV) — search: 0.11, pre-movement: 0.38, movement: 0.07), although this did not translate to a significant difference in navigational precision (Figure S1I). One possible explanation is that some participants were simply more efficient planners or were more skilled at planning on the move. Because we are interested in principles that are conserved across subjects, we pooled subjects for all subsequent analyses. While the duration of the search epoch was similar across arenas, the movement epoch duration increased drastically with increasing arena complexity (Figure 1I). This was understandable as the more complex arenas posed, on average, longer trajectories and more winding paths by virtue of their lower centrality. Notably, the pre-movement duration was also higher in more complex arenas, reflecting the subject’s commitment to meet the increased planning demands in those arenas (Figures 1I, S1H). Nonetheless, the relative pre-movement duration did not correlate with the probability of reward, and was similar for rewarded and unrewarded trials (Figures S1H). This suggests that the participants’ performance is limited by their success in remembering the reward location, rather than in meeting planning demands. Overall, these results suggest that in the presence of unambiguous visual information, humans are capable of adapting their behavior to efficiently solve navigation problems in relatively complex, unfamiliar environments.

A computational analysis supports that human eye movements are task relevant

Aiming to gain insights from subjects’ eye movements, we begin by examining the spatial distribution of gaze positions during different trial epochs (Figure 2A). Within each trial, the spatial spread of the gaze position was much larger during visual search than during the other epochs (mean spread ± SD — search: 6.3±2.4 m, pre-movement: 1.9±0.6 m, movement: 3.0±0.6 m; Figure 2D — left). This pattern was reversed when examining the spatial spread across trials (mean spread ± SD — search: 5.4±2.4 m, pre-movement: 6.7±1.2 m, movement: 6.5±0.6 m; Figure 2D — right). This suggests that subjects’ eye movements during pre-movement and movement were chiefly dictated by trial-to-trial fluctuations in task demands.

How did the task demands constrain human eye movements? Studies have shown that reward circuitry tends to orient the eyes toward the most valuable locations in space [38, 39]. Moreover, when the goal is hidden, it has been argued that fixating the hidden reward zone may allow for the oculomotor circuitry to carry the burden of remembering the latent goal location [38, 40]. Consistent with this, subjects spent a large fraction of time looking at the reward zone (pre-movement: 67±3%, movement: 53±6%). However, this
fraction decreased with arena complexity (Figures 2c — top) resulting in a larger mean distance between the gaze and the goal in more complex arenas (Figure 2c — bottom). This effect could not be attributed to subjects forgetting the goal location in more complex arenas, as we found a similar trend when analyzing gaze in relation to the eventual stopping location (believed goal location; Figure S2). A more plausible explanation is that constantly looking at the goal may not enable subjects to efficiently learn the task-relevant transition structure of the environment, as the transition structure is both more instrumental to solving the task and harder to comprehend in more challenging arenas. If central vision is attracted to the remembered goal location only when planning demands are low, this tendency should become more prevalent as subjects approach the target. Indeed, subjects spend significantly more time looking at the goal when there is a straight path to the goal than when the obstacle configuration requires that they make at least one turn prior to arriving upon such a straight path (Figure 2d). As mentioned earlier, computing the optimal trajectory requires precisely knowing both the reward function as well as the transition structure. While examining the proximity of gaze to goal allows for the study of the extent to which eye movements are dedicated to encoding the reward function, how may we assess the effectiveness with which subjects interrogate the transition structure of the environment to solve the task of navigating from point A to point B? This is the paramount challenge that we address in the remainder of this section by constructing a theoretical measure to guide data analysis.

In the case that a subject has a precise model of the transition structure of the environment, they would
be theoretically capable of planning trajectories to the remembered goal location without vision. However, in this experiment, the arena configurations were unfamiliar to the subjects, such that they would be quite uncertain about the transition structure. The finding that subjects achieved near-optimal performance on even the first few trials in each arena (Figure S1b) indicates that humans are capable of using vision to rapidly reduce their uncertainty about the aspects of the model needed to solve the task. Motivating the quantitative characterization of the task relevance of visual samples, we show an illustration of the consequence of mistaken beliefs about the passability of specific transitions on the subjective value function. Transition toggling can be defined as the act of removing an obstacle between two states if an obstacle was previously present, or adding an obstacle at that location if one was previously absent. In alignment with intuition, some transitions are more important to veridically represent (Figure 3a — middle), as toggling them results in a dramatic change to the value function (which is essential to computing the optimal set of actions to reach the goal), while toggling some other transitions causes a relatively minimal change (Figure 3a — right).
We leveraged this insight and defined a metric to quantify the task-relevance of each transition by computing the magnitude of the change in value of the subject’s current state, for a given goal location, if the status of the transition was toggled:

$$\Omega_k(s_0, s_G) = [V(s_0|T_k = 1) - V(s_0|T_k = 0)]^2$$  \hspace{1cm} (1)

where $\Omega_k(s_0, s_G)$ denotes the relevance of the $k^{th}$ transition for navigating from state $s_0$ to the goal state $s_G$, $T_k$ denotes the status of that transition (1 if it is passable and 0 if it is an obstacle), and $V(s_0|T_k = 1)$ denotes the value of state $s_0$ computed with respect to the goal state $s_G$ by setting $T_k$ to 1. It turns out that this measure of relevance is directly related to the magnitude of expected change in subjective value of the current state when looking at the $k^{th}$ transition, provided that the transitions are stationary and the subject’s uncertainty is uniform across transitions (Supplementary Notes). Thus, maximally relevant transitions identified by Equation (1) are precisely those which may engender the greatest changes of mind about the current return (i.e. utility) of the current state. In the supplementary notes, we point to a generalization of this relevance measure for settings in which the transition structure is stochastic (e.g. in volatile environments) and the subjective uncertainty is heterogeneous (i.e. the subject is more certain about some transitions than others).

We found that the transitions with the highest relevance on each trial strikingly correspond to bottleneck transitions that bridge clusters of interconnected states (Figures 3a). We quantified this effect using betweenness centrality, a metric describing the degree to which a state controls the traffic flowing through the arena (see Methods). The mean relevance of a transition was strongly correlated with the betweenness centrality averaged across two states on either side of the transition (3a — left). At the same time, the relevance was also high for obstacles that precluded a straight path to the goal (Figure 3a — center), as well as for transitions along the optimal trajectory (3a — right). By defining relevance of transitions according to Equation (1) we can thus capture multiple task-relevant attributes in a succinct manner. Theoretically investigating whether looking at task-relevant transitions improves navigational efficiency, we simulated artificial agents performing the same task that we imposed upon our human subjects. One agent (“perfect”) had a veridical subjective model of the environment, and thus was capable of computing the optimal trajectory (Figure S4). Its antithesis (“blind”) had an incorrect subjective model where half of the obstacle positions were ‘misremembered’ (toggled), but this agent was incapable of using vision to correct their model prior to taking actions according to their subjectively computed value functions. Performance at these two extremes was compared against the performance of three agents that were allocated a fixed budget of ‘saccades’ to rectify their incorrect models. These agents either randomly interrogated transitions (“random”), preferentially sampled transitions along the direction connecting the agent’s starting location to the goal location (“goalward”), or chose the most task-relevant transitions as defined by the relevance metric in Eq (1)”smart”).

While all three agents showed an improvement over the “blind” agent, the agent with knowledge about the most task-relevant transitions resulted in much shorter average path lengths than agents looking at transitions along the general direction of the goal or looking at random transitions (mean path length ± SE

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**Figure 3:** Simulations validate the utility of precisely knowing the status of theoretically important transitions. A. Value functions corresponding to an arbitrary goal location (closed circle) in an example arena (left) and in the arenas resulting from blocking either a bottleneck transition (center) or a transition that was not a bottleneck (right). B. Theoretical relevance of all transitions (circles) for the example arena for four different pairs of start (open black circle) and goal (closed black circle) states. C. Left: Across all possible start and end locations, the mean normalized relevance of non-obstacle transitions (across all possible start and goal state pairings) was positively correlated with betweenness centrality values. Middle: The mean normalized relevance of obstacles was negatively correlated with the eccentricity of each obstacle from the straight line connecting the current state to the goal state. Right: Transitions that fell on the optimal trajectory had greater relevance than those that fell outside of it. D. Simulation results of an agent instantiated with a perfect transition model (orange), and four agents with imperfect transition models, three of whom were endowed with the ability to correct their model according to different rules (see text). Those three agents were allowed to make eight ‘saccades’, each of which could update one transition. Left: Cumulative distributions (CDFs) of the path lengths of various agents (100 trials each from 25 different arenas; see Methods). Right: Median of trial-averaged path lengths across all simulated arenas; data points denote trial-averaged path lengths in individual arenas. E. Results of simulations similar to D but with a variable budget of ‘saccades’. Each line denotes the average path length (across arenas) of one agent as a function of the number of ‘saccades’. For each trial, path lengths of different agents were normalized by the optimal path length before trial-averaging. Error bars denote ±1 SEM.
Moreover, the performance of the smart sampling agent quickly approached optimality as the number of sampled transitions increased (Figure 3b). The rate of performance improvement was substantially slower for the goalward and random samplers (linear rather than exponential). These results were robust to the precise algorithm used to compute the value function in Eq [1]. In particular, the successor representation (SR) has been proposed as a computationally efficient, biologically plausible alternative to pure model-based algorithms like value iteration for responding to changing goal locations [8, 26]. We found that estimating the task-relevance of transitions using values implied by SR resulted in a similar performance improvement (Figure S5). Nevertheless, we emphasize that our objective was to use the relevance metric simply as a means to probe whether humans preferentially looked at task-relevant transitions. Understanding how the brain might compute such metrics is outside the scope of this study.

To apply the above theoretical insights to the assessment of subjects’ eye movements, we quantified the usefulness of subjects’ eye position on each frame as the relevance of the transition closest to the point of gaze, normalized by that of the most relevant transition in the entire arena for that trial. This resulted in frame-by-frame gaze relevance values that ranged between zero (least relevant) and one (most relevant). Then, we constructed a distribution of shuffled relevances by analyzing gaze with respect to a random goal location. Figure 4a shows the resulting cumulative distributions across trials for the average subject during the three epochs in an example arena. As expected, the relevance of subjects’ gaze was not significantly different from chance during the search epoch, as the subject had not yet determined the goal location. However, relevance values were significantly greater than chance both during pre-movement and movement (median relevance and interquartile range for the most complex arena, pre-movement — true: 0.21{0.06}, shuffled: 0.06{0.04}; movement — true: 0.21{0.07}, shuffled: 0.11{0.05}). Results were qualitatively similar in other arenas (Figure S6 Table S1).

To concisely describe subjects’ tendency to orient their gaze toward relevant transitions in a scale-free manner, we constructed receiver operating characteristic (ROC) curves by plotting the cumulative probability of shuffled gaze relevances against the cumulative probability of true relevances (Figure 4a — rightmost). An area under the ROC curve (AUC) greater (less) than 0.5 would indicate that the gaze relevance was significantly above (below) what is expected from a random gaze strategy. Across all arenas, the AUC was highest during the pre-movement epoch (Figures 4b; mean AUC ± SD — search: 0.52±0.03, pre-movement: 0.76±0.04, movement: 0.71±0.06). This suggests that subjects were most likely to attend to relevant transitions when contemplating potential actions before embarking upon the trajectory.

As the most relevant transitions can sometimes be found near the goal (e.g. Figure 3b — left), it is natural to wonder whether our evaluation of gaze relevance was confounded by the observation that subjects spent a considerable amount of time looking at the goal location (Figure 2c). Therefore, we first quantified the tendency to look at the goal location in a manner analogous to the analysis of gaze relevance (Figures 4a-b) by computing the area under the ROC curves (AUC) constructed using the true vs. shuffled distributions of the fraction of the duration spent foveating the goal in each epoch (Figure 4c-d). Across all arenas, AUCs were high during the pre-movement and movement epochs, confirming that there was a strong tendency for subjects to look at the goal location (Figure 4e). When we excluded gaze positions that fell within the reward zone while computing relevance, we found that the degree to which subjects looked at task-relevant transitions outside of the reward zone increased with arena complexity: the tendency to look at relevant transitions was greater in more complex arenas, falling to chance for the easiest arena (Figure 4f; Pearson’s r — pre-movement: -0.95, movement: -0.87). In contrast, the tendency to look at the goal location followed the opposite trend: it was greater in easier arenas (Figure 4f; Pearson’s r — pre-movement: 0.95, movement: 0.88). These analyses reveal a striking trade-off in the allocation of gaze between encoding the reward function and transition structure that closely mirrors the cognitive requirements of this task. This trade-off is not simply a consequence of directing gaze more/less often at the reward, as such temporal statistics are preserved while shuffling. Instead, it points to a strategy of directing attention away from the reward and towards task-relevant transitions in complex arenas. This compromise allowed subjects to dedicate more time to surveying the task-relevant structure in complex environments and likely underlies their ability to take near-optimal paths in all environments, albeit at the cost of an increased tendency to
Figure 4: Eye movements reveal a cognitive trade-off between reward and transition encoding. A. Left: Cumulative distribution (computed by pooling trials from all subjects) of average log (normalized) relevance values (pink) and the corresponding shuffled distribution (gray) during search (left), pre-movement (center), and movement (right) epochs (data for the most complex arena is shown). Shaded regions denote 95% confidence bounds computed using Greenwood’s Formula. Rightmost: ROC curves characterizing the gaze relevance during the three epochs. B. Area under the ROC curves (AUC) for different epochs, colored according to the color scheme in A. C-D: Similar plots as A-B, but for the distributions of the log fraction of the duration in each epoch spent gazing within two meters of the eventual stopping position (which was assumed to be the subjects’ believed goal location). E. AUC values of gaze relevance computed for the distributions of trial-averaged relevances, after excluding fixations within the reward zone, during the pre-movement (orange) and movement (blue) epochs. F. Similar to E, but showing the AUC values of gaze durations within the reward zone. All error bars were computed using bootstrapping.

The temporal evolution of gaze includes distinct periods of sequential prospection

So far, we have shown that the spatial distribution of eye movements adapts to trial-by-trial fluctuations in task demands induced by changing the goal location and/or the environment. However, planning and executing optimal actions in this task requires dynamic cognitive computations within each trial. To gain insights into this process, we examined the temporal dynamics of gaze. Figure 5a shows a participant’s gaze in an example trial which has been broken down into nine epochs (pre-movement: I-VI, movement: VII-IX) for illustrative purposes (see Video S1 for more examples). The participant initially foveated the goal location (epoch I), and their gaze subsequently traced a trajectory backwards from the goal state towards their starting position (II) roughly along a path which they subsequently traversed on that trial (dotted line). This sequential gaze pattern was repeated shortly thereafter (IV), interspersed by periods of non-sequential eye movements (III and V). Just before embarking on their trajectory, the gaze traced the trajectory, now in the forward direction until the end of the first turn (VI). Upon reaching the first turning point in their trajectory (VII), they executed a similar pattern of sequential gaze from their current position toward the goal (VIII), tracing out the path which they navigated thereafter (IX). We refer to the sequential eye movements along the future trajectory in the backwards and forwards direction as backward sweeps and forward sweeps, respectively. During such sweeps, subjects seemed to rapidly navigate their future paths with their eyes, and all subjects exhibited sweeping eye movements without being explicitly instructed to plan their trajectories prior to navigating. The fraction of time that subjects looked near the trajectories which they subsequently embarked upon increased with arena difficulty (Figure S7a). To algorithmically detect periods of sweeps, gaze positions on each trial were projected onto the trajectory taken by the participant by locating the positions along the trajectory closest to the point of gaze on each frame (Methods). On each frame, the

forget the precise goal location in complex environments (Figure 1b).
geodesic length of the trajectory up until the point of the gaze projection was divided by the total trajectory length, and this ratio was defined as the "fraction of trajectory". We used the increase/decrease of this variable to automatically determine the start and end times of periods when the gaze traveled sequentially along the trajectory in the forward/backward directions (sweeps) for longer than chance (Figure 5b; see Methods).

Figure 5: Gaze traveled forwards and backwards along the intended trajectory. A. Spatial locations of gaze positions (the arrow of relative time within each window increases from violet to orange) and subject positions (violet to blue) during individual time windows demarcated in the bottom panel. Panels in the bottom row correspond to time periods corresponding to sweeps. The subject’s trajectory from the starting location (open black circle) to the goal (closed black circle) is denoted by a black dashed line. B. Time-series of the points on the trajectory that were closest to the subject’s gaze on each frame, expressed in percentiles (0: start of trajectory, 1: end of trajectory) during one example trial. Only frames during which the gaze position fell within 2 m of the trajectory are plotted. The gray trace shows the movement velocity of the subject during this trial. Red and green shaded regions highlight time windows during which the sweep classification algorithm detected backward and forward sweeps, respectively. In this trial, there were two backward sweeps before movement, and one forward sweep each before and during movement. C. Across all subjects, the fraction of time spent sweeping in the forward and backward directions within each epoch reveals an antiparallel effect: more time was spent sweeping forwards during movement than during pre-movement (top), whereas more time was spent sweeping backwards during pre-movement than during movement (bottom). Error bars denote ±1 SEM.

The complexity of the transition structure exerted a strong influence on the probability of sweeping: the fraction of trials in which this phenomenon occurred was significantly correlated with arena centrality scores (Pearson’s r = -0.95, p = 0.01; Figures S7b). This suggests that sweeping eye movements could be integral
to trajectory planning. It turns out that aside from gazing upon the target or the trajectory on each trial, roughly 20% of eye movements were made to other locations in space (Figure S7c), but this percentage did not vary across arenas nor epochs. Most notably, on average, backward sweeps occupied a greater fraction of time during pre-movement than during movement, but forward sweeps predominantly occurred during movement (backward sweeps — pre-movement: 6.0 ± 1.2%, movement: 2.0 ± 0.2%; forward sweeps — pre-movement: 5.5 ± 0.6%, movement: 10.2 ± 1.5%; Figure 5b). This suggests that the initial planning is primarily carried out by sweeping backwards from the goal. The mean speed of backward sweeps was greater than the speed of forward sweeps across all arenas (backward sweeps: 11.3 ± 0.9 m/s, forward sweeps: 6.7 ± 0.2 m/s; Figure S7d). Notably, sweep velocities were 3-4 times greater than the maximum movement velocity (2.26 m/s) and more than fivefold greater than the average subject velocity during the movement epoch (1.47 ± 0.08). This is reminiscent of the hippocampal replay or preplay of trajectories through space, as such sequential neural events are also known to be compressed in time (around 2-20x the speed of neural sequence activation during navigation) [30, 41]. Both sweep speeds and durations slightly increased with arena complexity (Figures S7f). An explanation for this finding requires first recognizing that peripheral vision processing must lead the control of central vision to allow for sequential eye movements to trace a viable path [42, 43]. In more complex arenas such as the maze where the search tree is narrow and deep, the obstacle configuration is more structured and presents numerous constraints, and thus path tracing computations might occur more quickly. However, due to the lengthier trajectories in those arenas, the gaze must cover greater distances, resulting in sweeps which last longer. Another property of sweeps is that they were comprised of increasingly more saccades as arenas became complex, and saccade rates were higher during sweeps than at other times after goal detection (Figure S7g). This suggests that either visual processing during sweeps was expedited compared to average, or sweeps resulted from eye movements which followed a pre-planned saccade sequence.

If the first sweep on a trial occurred during pre-movement, the direction of the sweep was more likely to be backwards, while if the first sweep occurred during movement, it was more likely to be in the forwards direction (Figure S7h). The latency between goal detection and the first sweep increased with arena difficulty (Figure S7g), suggesting that sweep initiation is preceded by brief processing of the arena, and more complex arenas elicited longer processing. While the sequential nature of eye movements could constitute a swift and efficient way to perform instrumental sampling, we emphasize that task-relevant eye movements were not necessarily sequential. When we reanalyzed the spatial distribution of gaze positions by removing periods of gazing upon the trajectory on each trial, the resulting relevance values remained significantly greater than chance (Figure S7h).

What task conditions promote sequential eye movements? To find out, we computed the probability that the subjects engaged in sweeping behavior as a function of time and position, during the pre-movement and movement epochs respectively, and focusing on the dominant type of sweep during those periods (backward and forward sweeps respectively; Figure 5). During pre-movement, we found that the probability of sweeping gradually increased over time, suggesting that backward sweeps during the initial stages of planning are separated from the time of target foveation by a brief pause, during which subjects may be gathering some preliminary information about the environment (Figure 6 — left). During movement, on the other hand, it turns out that the probability of sweeping is heavily influenced by whether subjects are executing a turn in their trajectory. Obstacles often preclude a straight path to the remembered goal location, and thus participants typically find themselves making multiple turns while actively navigating. Consequently, a trajectory may be broken down into a series of straight segments separated by brief periods of elevated angular velocity. We isolated such periods by applying a threshold on angular velocity, designating the periods of turns as subgoals, and aligned the subject’s position in all trials with respect to subgoals. The likelihood of sweeping the trajectory in the forward direction tended to spike precisely when subjects reached a subgoal (Figure 6 — right). There was a concomitant decrease in the average distance of the point of gaze from the goal location in a step-like manner with each subgoal achieved (Figure 6 — right). In contrast to backward sweeps, which were made predominantly to the most proximal subgoal prior to navigating (Figure 6 — left), forward sweeps that occurred during movement were not regularly directed toward one particular location. Instead, in a strikingly precise and stereotyped manner, subjects appeared to lock their gaze upon the upcoming subgoal when rounding each bend in the trajectory (Figure 6 — right). This suggests that
subject likely represented their plan by decomposing it into a series of subgoals, focusing on one subgoal at a time until they reached the final goal location.

To summarize, we found subjects made sequential eye movements sweeping forward and/or backward along the intended trajectory, and the likelihood of sweeping increased with environmental complexity. During the pre-movement phase, participants typically traced the trajectory backwards from the goal to the first subgoal (Figure 6d — orange). While moving through the arena, they tended to lock their gaze upon the upcoming subgoal until they reached it, thereafter sweeping their gaze forward to the next subgoal (Figure 6d — blue). Via eye movements, navigators could perform active sensing and construct better-informed plans to refine their policies such that the sequence of actions that they choose would most efficiently lead to rewards (Figure 6e).
Discussion

In this study, we highlight the critical role of eye movements for model-based computations that support navigation in complex environments. We found that humans took near-optimal trajectories in all environments, but spent more time planning prior to navigation in more complex environments. The spatial distribution of gaze was largely concentrated at the hidden goal location in the simplest environment, but increasingly skewed towards the task-relevant structure of the environment as the complexity increased. In the temporal domain, subjects often traced trajectories to and from the goal rapidly with their eyes (sweeping), and subsequently concentrated on one subgoal at a time until they reached the goal. In summary, we found evidence that the neural circuitry governing the oculomotor system optimally schedules and allocates resources to tackle the diverse cognitive demands of navigation, producing efficient eye movements through space and time.

Eye movements provide a natural means for researchers to understand active sensing strategies [44, 45]. Past studies using simple decision making tasks probed whether active sensing, specifically via eye movements, reduces uncertainty about the state of the environment (such as whether a change in an image has occurred) [3, 4, 46]. But in sequential decision making tasks such as navigation, there is added uncertainty about the causal structure (model) of the environment – i.e. task contingencies [2]. Common paradigms for goal-oriented navigation occlude large portions of the environment from view, usually in the interest of distinguishing model-based strategies from conditioned responses [47–49]. Oclusions eliminate the possibility of gathering information about the structure of the environment using active sensing. By removing such constraints, we allowed subjects to acquire a model of the environment without physically navigating through it, which yielded new insights about how active sensing mitigates model uncertainty. In particular, we found that the gaze is distributed between the two components of the model required to plan a path — the transition function, which describes the relationship between states, and the reward function, which describes the relationship between the states and the reward — with the distribution skewed in favor of the former in more complex environments. This suggests a context-dependent mechanism which dictates the dynamic arbitration between competing controllers of the oculomotor system that seek information about complementary aspects of the task. Neurally, this could be implemented by circuits that have executive control of voluntary eye movements. Candidate substrates include the dorsolateral prefrontal cortex, which is known to be critical for contextual information processing and memory-guided saccades [50–52], and the anterior cingulate cortex, which is known to be involved in evaluating alternative strategies [53, 54].

Eye movements are also influenced by many other neural circuits. Hippocampal projections to higher oculomotor controllers (e.g. supplemental eye fields through the orbitofrontal cortex) may guide the embodiment of simulation and memory through eye movements [55–57]. The contents of gaze have also previously been found to influence activity in the hippocampus [58–64] and entorhinal cortex [65–67]. Mentally simulating sequences of actions is a theorized function of hippocampal replay and preplay [31–35, 68]. Several recent studies have also supported that replay serves to consolidate memory and generalize information about rewards [69–71]. Shortly after fixating on the goal, subjects’ gaze often swept backwards along their future trajectory, evocative of both preplay and depth-first tree search, a model-based algorithm for path discovery [72]. Past studies have demonstrated that humans can smoothly trace paths through novel 2D mazes, suggesting that peripheral vision processing leads saccade generation [43, 73], and we suspect that a similar mechanism could underlie the sweeping eye movements in our task. To better understand the precise neural mechanisms underlying the gaze patterns we observed, it would be enlightening to examine the direction of information flow between the oculomotor circuitry and brain regions with strong spatial representations during this task.

Furthermore, the tendency of humans to break larger problems into smaller, more tractable subtasks has been previously established in domains outside of navigation [27, 65–67, 74]. However, theoretical insights on clustered representations of space have not been empirically validated in the context of navigation [75, 76], primarily due to the difficulty in distinguishing between flat and hierarchical representations from behavior alone. As a first step towards this direction, our observation that subjects’ locus of concentra-
tion was often locally oriented towards navigating to the end of each path segment suggests that subjects viewed turns as subgoals of the overall plan.

We used the RL framework to formulate the task-relevance of an eye movement by assessing the extent to which incorrect knowledge about the navigability of specific locations would affect navigation efficiency. Our approach was agnostic to the temporal ordering of relevant eye movements. Given that previous work demonstrated evidence for the planning of multiple saccades during simple tasks like visual search [7], incorporating chronology into the theory of active sensing in sequential decision-making tasks presents an excellent opportunity for future studies. One complementary approach to the study of active sensing involves the information theoretic framework [13]. A recently proposed framework of active inference unifies planning and information seeking by integrating the RL framework, which describes exploiting rewards for their extrinsic value, and the information theoretic framework, which describes exploring new information for its epistemic value [2]. This framework inspires future research that can unify the various contributing influences on eye movements.

We hope that the study of active sensing and planning during navigation will eventually generalize to understanding how humans accomplish a variety of sequential decision-making tasks. A major goal in the study of neuroscience is to elucidate the principles of biological computations which allow humans to effortlessly exceed the capabilities of machines. Such computations allow animals to learn environmental contingencies and flexibly achieve goals in the face of uncertainty. However, one of the main barriers to the rigorous study of active, goal-oriented behaviors is the complexity in estimating the subject’s prior knowledge, intentions, and internal deliberations which lead to the actions that they take. Luckily, eye movements reveal a wealth of information about ongoing cognitive processes during tasks as complex and naturalistic as spatial navigation.
Methods

Experimental Model and Subject Details. Nine human subjects (all >18 years old, six males) participated in the experiments. All but two subjects (S6 and S9) were unaware of the purpose of the study. Four of the subjects, including S6 and S9, were exposed to the study earlier than the rest of the subjects, and part of the official dataset for two of these subjects (S4 and S8) was collected two months prior to the rest of data collection as a safety precaution during the COVID-19 pandemic. Six additional human subject recruits (all >18 years old, two males) were disqualified due to experiencing motion sickness while in the VR environment. All experimental procedures were approved by the Institutional Review Board at New York University and all subjects signed an approved consent form.

Stimulus. Subjects were seated on a swivel chair with 360° of freedom in physical rotation and navigated in a full-immersion hexagonal virtual arena with several obstacles. The stimulus was rendered at a frame rate of 90 Hz using the Unity game engine v2019.3.0a7 (programmed in C#) and was viewed through an HTC VIVE Pro virtual reality headset. The subjective vantage point (height of the point between the subjects’ eyes with respect to the ground plane) was 1.72 meters. The subject had a field of view of 110.1° of visual angle. Forward and backward translation was enabled via a continuous control CTI Electronics M20U9T-N82 joystick with a maximum speed of 2.26 m/s. Subjects executed angular rotations inside the arena by turning their head, while the joystick input enabled translation in the direction in which the subject’s head was facing. Obstacles and arena boundaries appeared as gray, rectangular slabs of concrete. The ground plane was grassy, and the area outside of the arena consisted of a mountainous background. Peaks were visible above the outer boundary of the arena to provide crude orientation landmarks. Clear blue skies with a single light source appeared overhead.

State space geometry. The arena was a rectangular hexagon enclosing an area of approximately 260 m² of navigable space. For ease of simulation and data analyses, the arena was imparted with a hidden triangular tessellation (deltille) composed of 6n² equilateral triangles where n determines the state space granularity. We chose n = 5, resulting in triangles with a side length of 2 meters, each of which constituted a state in the discrete state space (Figure S1a). The arena contained several obstacles in the form of unjumpable obstacles (0.4 meters high) located along the edges between certain triangles (states). Obstacle locations were predetermined offline using MATLAB by either randomly selecting a chosen number of edges of the tessellation or by using a graphical user interface (GUI) to manually select edges of the tessellation; these locations were loaded into Unity. Outer boundary walls of height 2.5 m enclosed the arena. We chose five arenas spanning a large range in average state closeness centrality \( \langle C(s) \rangle \) (Eq 2), where \( C(s) \) is defined as the inverse average path length \( d \) from state \( s \) to every other state \( s' \) (\( N \) states in total). On average, arenas with lower centrality will impose a greater path length between two given states, making them more complex to navigate. The order of arenas presented to each subject was randomly permuted but not entirely counterbalanced due to the large number of permutations (Table S2).

\[
C(s) = \frac{N - 1}{\sum_{s'} d(s', s)}
\]  

Eye tracking. At the beginning of each block of trials, subjects calibrated the VIVE Pro eye tracker using inbuilt Tobii software which prompted subjects to foveate several points tiling a 2D plane in the VR environment. Both eyes were tracked, and the subject’s point of foveation (x-y coordinates), object of foveation (ground, obstacles, boundaries, etc.), eye openness, and other variables of interest were recorded on each frame using the inbuilt software. Sipatchin et. al. (2020) reported that during free head movements, point-of-gaze measurements using the VIVE Pro eye tracker has a spread of 1.15° ± 0.69° (SE) [77]. This means that when the subject fixates a point on the ground five meters away, the 95% confidence interval (CI) for the measurement error in the reported gaze location would be 0-23 cm (roughly one-tenth of the length of one transition or obstacle) and 0-67 cm (one-third of a transition length) for points fifteen meters away. While machine precision was not factored into the analyses, the fraction of eye positions that may have been misclassified due to hardware and software limitations is likely very tiny. Furthermore, Sipatchin et. al.
reported that the system latency was 58.1 ms. While there is reason to suspect that the subject’s position was recorded with a similar latency of around five frames, even if the gaze data lagged the position data, the subject would only have moved 13 cm if they were translating at the maximum possible velocity over this interval.

**Behavioral task.** At the beginning of each trial, a target in the form of a realistic banana from the Unity Asset store appeared hovering 0.4 meters over a state randomly drawn from a uniform distribution over all possible states. The joystick input was disabled until the subject foveated the target, but the subject was free to scan the environment by rotating in the swivel chair during the visual search period. Two hundred milliseconds after target foveation, the banana disappeared and subjects were tasked with navigating to the remembered target location without time constraints. Subjects were not given instructions on what strategy to use to complete the task. After reaching the target, subjects pressed a button on the joystick to indicate that they have completed the trial. Alternatively, they could press another button to indicate that they wished to skip the trial. Feedback was displayed immediately after pressing either button (see section below). Skipping trials was discouraged except when subjects did not remember seeing the target before it disappeared, and these trials were recorded and excluded from the analyses (< 1%).

**Reward.** If subjects stopped within the triangular state which contained the target, they were rewarded with two points. If they stopped in a state sharing a border with the target state, they were rewarded with one point. After the subject’s button press, the number of points earned on the current trial was displayed for one second at the center of the screen. The message displayed was ‘You earned p points!’; the font color was blue if $p = 1$ or $p = 2$, and red if $p = 0$. On skipped trials, the screen displayed ‘You passed the trial’ in red. In each experimental session, after familiarizing themselves with the movement controls by completing ten trials in a simplistic six-compartment arena (granularity, $n = 1$), subjects completed one block of fifty trials in each of five arenas (Figure 1b). At the end of each block, a blue message stating ‘You have completed all trials!’ prompted them to prepare for the next block. Session durations were determined by the subject’s speed and the length of the breaks that they needed from the virtual environment, ranging from 1.5-2 hours. Subjects were paid $0.02/point for a maximum of 5 arenas x 50 trials/arena x 2 points/trial x $0.02/point = $10, in addition to a base pay of $10/hour for their time (the average payment was $27.72).

**RL formulation.** Navigation can be formulated as a Markov Decision Process (MDP) described by the tuple $< S, A, P, R, \gamma >$ whose elements denote, respectively, a finite state space $S$, a finite action space $A$, a state transition distribution $P$, a reward function $R$, and a temporal discount factor $\gamma$ that captures the relative preference of distal over proximal rewards [78]. Given that an agent is in state $s \in S$, the agent may execute an action $a \in A$ in order to bring about a change in state $s \rightarrow s'$ with probability $P(s'|s, a)$ and harvest a reward $R(s, a)$. To relate this formalism to the structure of the arena, it is instructive to consider the possibility of traversal from state $s$ to any state $s'$ in a single time step as described by the adjacency matrix $T$: $T(s, s') = 1$ if there exists an available action which would bring about the change in state $s \rightarrow s'$ with a non-zero probability, and $T(s, s') = 0$ otherwise. By definition, $T(s, s') = 0$ if there is an obstacle between $s$ and $s'$. Thus, the arena structure is fully encapsulated in the adjacency matrix.

In the case that an agent is tasked with navigating to a goal location $s_G$ where the agent would receive a reward, the reward function $R(s, a) > 0$ if and only if the action $a$ allows for the transition $s \rightarrow s_G$ in one time step, and $R(s, a) = 0$ otherwise. Given this formulation, we may compute the optimal policy $\pi^*(a|s)$, which describes the actions that an agent should take from each state in order to reach the target state in the fewest possible number of time steps. The optimal policy may be derived by computing optimal state values $V^*(s)$, defined as the expected future rewards to be earned when an agent begins in state $s$ and acts in accordance with the policy $\pi^*$. The optimal value function can be computed by solving the Bellman Equation (Eq 3) via dynamic programming methods such as value iteration, an algorithm that involves iteratively unrolling the recursion in this equation [79]. The optimal policy is given by the argument $a$ that maximizes the right-hand side of Eq. 3. Intuitively, following the optimal policy requires that agents take actions to ascend the value function where the value gradient is most steep (Figure 1b).
We incorporated twelve possible degrees of freedom in the action space, such that one-step transitions could result in relocating to a state that is 0°, 30°, 60°, . . . , 300°, or 330° with respect to the previous state. However, the center-to-center distances between states for a given transition depends on the angle of transition. Specifically, as shown in Figure [ST]b, if a step in the 0° direction requires translating 1 m, then a step in the 30°, 120°, 180°, 240°, and 300° directions would also require translating 1 m, but a step in the 30°, 150°, and 270° directions would require translating 2\sqrt{3}/3 m, and a step in the 90°, 210°, and 330° directions would require translating \sqrt{3}/3 m. Therefore, in Eq 3, \( R(s,a) = -1, -2\sqrt{3}/3, \) or \( -\sqrt{3}/3, \) depending on the step size required in taking an action \( a. \) The value of the goal state \( s_G \) was set to zero on each iteration. Value functions were computed for each goal location, and the relative value of states describes the relative minimum number of time steps required to reach \( s_G \) from each state. The lower the value of a state, the greater the geodesic separation between the state and the goal state. We set \( \gamma = 1 \) during all simulations and performed 100 iterations before calculating optimal trajectory lengths from an initial state \( s_i \) to the target state \( s_G, \) as this number of iterations allowed for value iteration to converge.

Relevance computation. To compute the relevance \( \Omega_k(s_0,s_G) \) of the \( k \)th transition to the task of navigating from a specific initial state \( s_0 \) to a specific goal \( s_G, \) we calculated the absolute change induced in the optimal value of the initial state after toggling the navigability of that transition by changing the corresponding element in the adjacency matrix from 1 to 0 or from 0 to 1 (Eq 4). For the simulations described below, we also tested a non-myopic, path-dependent metric \( \Omega_k(s_0,s_G;\pi^*) \) defined as the sum of squared differences induced in the values of all states along the optimal path (Eq 5). Furthermore, we tested the robustness of the measure to the precise algorithm used to compute state values by computing value functions using the successor representation (SR) algorithm, which caches future state occupancy probabilities learned with a specific policy [26]. As we used a random walk policy, we computed the matrix of probabilities \( M \) analytically by temporally abstracting a one-step transition matrix \( T: M = (I - \gamma T)^{-1}. \) The cached probabilities can then be combined with a one-hot reward vector \( R(s) = \mathbb{1}(s = s_G) \) to yield state values \( V = MR. \) We set the temporal discount factor \( \gamma = 1 \) and integrated over 100 time steps.

\[
\Omega_k(s_0,s_G) = [V(s_0|T_k = 1) - V(s_0|T_k = 0)]^2
\]

\[
\Omega_k(s_0,s_G;\pi^*) = \sum_{s=s_0}^{s_G} [V(s|T_k = 1) - V(s|T_k = 0)]^2
\]

Relation to bottlenecks. In order to assess whether the relevance metric is predictive of the degree to which transitions are bottlenecks in the environment, we correlated normalized relevance values (averaged across all target locations and normalized via dividing by the maximum relevance value across all transitions for each target location) with the average betweenness centrality \( G \) of the two states on either side of a transition (Eq 6). Betweenness centrality essentially calculates the degree to which a state controls the traffic flowing through the arena. \( \sigma_{ij} \) represents the number of shortest paths between states \( i \) and \( j, \) and \( \sigma_{ij}(s) \) represents the number of such paths which pass through state \( s. \) For this analysis, transitions within 1 m from the goal state were excluded due to their chance of having spuriously high relevance values.

\[
G(s) = \sum_{a \neq s \neq b} \frac{\sigma_{ij}(s)}{\sigma_{ij}}
\]

Simulations. Behavior of three qualitatively different artificial agents with different planning capacities was simulated. All agents were initialized with a noisy model of the environment. Representational noise was simulated by toggling 50% of randomly selected unavailable transitions from \( T(s,s') = 0 \) to 1, and the equivalent number of randomly selected available transitions from \( T(s,s') = 1 \) to 0. This is analogous to the agents misplacing obstacles in their memories, or equivalently, a subjective-objective model mismatch induced by volatility in the environment. The blind agent was unable to correct its model during a planning
period. On each trial, eight transitions (out of 210 available) were drawn for each sighted agent; the agent’s model was compared with the true arena structure at these transitions and, if applicable, corrected prior to navigation. Samples were drawn uniformly from all possible transitions (without replacement) for the random exploration agent. For the goalward looking agent, the probability of drawing a transition was determined by a circular normal (von Mises) distribution with \( \mu = \theta_G \) (where \( \theta_G \) is the angle of the goal w.r.t the agent’s heading), \( \sigma = 1 \), and concentration parameter \( \kappa = 5 \). In contrast, the directed sampling agent gathered information specifically about the eight transitions that were calculated to be most relevant for that trial. After the model updates, if any, the agents’ subjective value functions were recomputed, and agents took actions according to the resulting policies. When an agent encountered a situation in which no action was subjectively available, they attempted a random action. In the case that a new action is discovered, the agents temporarily updated \( T(s, s') \) from 0 to 1 for that action. Conversely, in the case that an agent attempted to take an action but discovered that it was not actually feasible, they temporarily updated their subjective models to account for the transition block which they had just learned about. In both cases, value functions were recomputed using the updated model. Simulations were conducted with 25 arenas of granularity \( n = 3 \) (state space size = 54 for computational tractability) and 100 trials per arena. Furthermore, we tested the agents’ performance using a range of gaze samples evenly spaced between 2 and 14 foveations.

**Data processing.** In order to identify moving and non-moving epochs within each trial, movement onset and offset times were detected by applying a moving average filter of window size 5 frames on the absolute value of the joystick input function. When the smoothed joystick input exceeded the threshold of 0.2 m/s (approx. 10% of the maximum velocity), the subject was deemed to be moving, and when the input fell below this threshold for the last time on each trial, the subject was deemed to have stopped moving. Subjects’ relative planning time was defined as the ratio of pre-movement time to the total trial duration, minus the search period (which was roughly constant across arenas). Prior to any eye movement analyses, blinks were filtered from the eye movements by detecting when the fraction of the pupil visible dipped below 0.8. The spread in the \( (x, y) \) gaze positions within trials was calculated as the expectation of variance, \( \mathbb{E}_n[\sqrt{\text{Var}_t[x] + \text{Var}_t[y]}] \), where \( \text{Var}_t[\cdot] \) denotes the variance across time \( t \) within a trial and \( \mathbb{E}_n[\cdot] \) denotes expectation across trials denoted by \( n \). The spread across trials was calculated as the variance of the expectation, \( \sqrt{\text{Var}_n[\mathbb{E}_t[x]] + \text{Var}_n[\mathbb{E}_t[y]]} \), where \( \mathbb{E}_t[\cdot] \) denotes expectation across time and \( \text{Var}_n[\cdot] \) denotes the variance across trials.

For Figures 1f, 1g, S1d, S1e, and S1g, the first trial of each run was removed from the analyses due to an occasional rapid teleportation of the subject to a random starting location associated with the software starting up. While there were a few instances where more than one run occurred per block due to subjects adjusting the headset, at least 51 trials were actually collected during each block such that most blocks consisted of 50 trials when the first trial of each run was omitted. For analyses such as epoch duration, gaze distribution, relevance, sweep detection, and subgoal detection, the first trial was not discarded since the teleportation only affected the recorded path length, but as the teleportation was virtually instantaneous, the new starting locations on such trials could be used for analyses which do not depend upon the path length variable.

**Relevance estimation.** Prior to estimating the task-relevance of the subjects’ gaze positions at each time point, the closest transition \( k \) to the subject’s point of gaze was identified and the effect of toggling the transition on the value function was computed as \( \Omega_k(s(t), s_G) \). In order to construct a null distribution of relevance values, we paired the eye movements on each trial with the goal location for a random trial, given the subject's position in the current trial. This shuffled average is not task-specific, and therefore may be compared with the true \( \Omega \) values to probe whether the spatial distribution of gaze positions was sensitive to the goal location on each trial. Similarly, the shuffled fraction of time looking at the goal was computed with a goal state randomly chosen from all states.

**Sweep classification.** Forward and backward eye movements (sweeps) along the intended trajectory were classified by first calculating the point \( (x, y) \) on the trajectory closest to the location of gaze in each
frame. For each trial, the fraction of the total trajectory length corresponding to each point was stored as a variable \( f \), and periods when \( f(t) \) consecutively ascended or descended were identified. For each period, we determined \( m \), an integer whose magnitude denoted the sequence length and whose sign denoted the sequence direction (+/- for ascending/descending sequences). We then constructed a null distribution \( p(m) \) describing the chance-level frequency of \( m \) by selecting 20 random trials and recomputing \( f \) based on the subject’s trajectories on those trials. Sequential eye movements of length \( m \) where the CDF of \( p(m) \) was less than \( \alpha/2 \) or greater than \( 1 - \alpha/2 \) were classified as backward and forward sweeps, respectively. The significance threshold \( \alpha \) was chosen to be 0.02. Compensating for noise in the gaze position, we applied a median filter of length 20 frames to both the true and shuffled \( f \) functions. During post-processing, sweeps in the same direction that were separated by less than 25 frames were merged, and sweeps for which the gaze fell outside of 2 meters from the intended trajectory on >30% of the frames pertaining to the sweep were eliminated. Sweeps were required to be at least 25 frames in length. To remove periods of fixation, the minimum variance in \( f(t) \) values for all time points corresponding to the sweep was required to be 0.001. Finally, sweeps which did not cover at least 20% of the total trajectory length were removed from the analyses. This algorithm allowed for the automated detection of sequential eye movements pertaining to the prospective evaluation of trajectories which subjects subsequently took.

**Saccade detection.** Saccade times were classified to be the times at which eye movement speeds \( v \) crossed a threshold of 50°/s from below, where speeds were computed using Eq 7, where \( x, y, \) and \( z \) correspond to the coordinates of the point of gaze (averaged across both eyes), and \( \alpha \) and \( \beta \) respectively correspond to the lateral and vertical displacement of the pupil in degrees.

\[
\alpha(t) = \tan^{-1}\left(\frac{x(t)}{\sqrt{y^2(t) + z^2(t)}}\right), \quad \beta(t) = \tan^{-1}\left(\frac{z(t)}{\sqrt{y^2(t) + x^2(t)}}\right), \quad v(t) = \sqrt{\left(\frac{d}{dt}\alpha(t)\right)^2 + \left(\frac{d}{dt}\beta(t)\right)^2}
\]

(7)

**Subgoal analysis.** Turns in the subjects’ trajectories (defined as subgoals) were isolated by applying a threshold of 60 deg/s on their angular velocity (smoothed with a median filter; window size = 8 frames). The first and last frames for periods of elevated angular velocity were recorded. For the purposes of the analysis in Figure 6, all trials (for all subjects in all arenas) were broken down into periods of turns vs. periods of navigating straight segments. Starting from the stopping location, these periods were independently interpolated to fit an arbitrarily defined common timeline of 25 time points per turn and 100 time points per straight segment. For example, if a trial had two turns, then eye movement variables from the last turn to the stopping location were interpolated to the points -100 to -1. The last turn was represented by -125 to -101, and the second last turn was represented by -250 to -226. The segment between the two turns was represented by -225 to -226. Finally, the segment between the subject’s starting position and the first turn was represented by -350 to -251. Note that as a consequence, the number of trials for which there were (for example) more than four turns in the trajectory was substantially fewer than the number of trials for which there were one or no turns, such that the quantity of raw data contributing to each normalized position value in Figure 6 increases from left to right.
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References


Figure S1: A. Arenas were regular hexagons with a side length of 10 m, and a triangular tessellation with a unit length of 2 m. Two points were rewarded if participants reached the goal state (green), and one point was rewarded if participants reached a state neighboring the goal state (light green). B. To incorporate twelve degrees of freedom in translation, value functions were computed by dynamic programming whereby the cost of actions scaled in accordance with the center-to-center distance between states $s$ and $s'$ (pertaining to the transition which results from taking action $a$). C. Top: Across all subjects and all trials, the probability of being awarded two points (green) increased with arena centrality, while the probability of being awarded one point (light green) is relatively constant across all arenas. Gray denotes the probability of not being rewarded. Middle: The total fraction of points earned increased as a function of arena centrality. Bottom: Distance between the stopping location and goal in rewarded (green) and unrewarded (red) trials. Error bars denote ±1 SEM. D. Left: Across all arenas (colored according to the arena coloring scheme introduced in Figure 1b), the path lengths observed in unrewarded trials were close to the optimal trajectory lengths between the starting state and the state at which subjects stopped on these trials, suggesting that unrewarded trials were predominantly caused by subjects forgetting the precise location of the target. Right: The ratio of observed to optimal path lengths (to the subjects’ stopping location on unrewarded trials) was close to unity in all arenas. E. Performance was stable across each block, as measured by the average probability of being rewarded on each trial (green), as well as the average ratio between the empirical and optimal path lengths (gray). F. Distribution of epoch durations across all subjects and all trials. Pre-movement occupied a greater fraction of the total trial time for more complex arenas. G. Some subjects spent lesser time deliberating before movement, but this did not impact task performance. Relative pre-movement duration was defined as the average ratio of the duration of the pre-movement epoch to the duration of the entire trial after goal detection. The average proportion of time that subjects spent making prospective eye movements prior to using the joystick did not correlate with their average path lengths across all arenas (gray), nor with the overall probability of them being rewarded (green). H. The relative pre-movement duration did not differ between rewarded and unrewarded trials (after matching the mean trial duration of the two groups, separately for each arena), except for the two easiest arenas, where planning demands are low. This suggests that failure to obtain rewards is not mere due to poor planning. Error bars denote ±1 SEM.
Figure S2: Gaze is increasingly concentrated around the goal/stopping location for easier arenas. The believed goal location was assumed to be the subjects’ stopping position, and the point of gaze was visibly more concentrated around the stop location than around the true goal location (especially in the most complex arena). The effects of working memory on gaze were more apparent for easier, more open arenas. Each panel depicts eye movements on a random subset of trials (3 trials x 9 subjects) in each arena. The origin (0,0) denotes the goal location or the stopping location. Raw gaze positions relative to these points are depicted during the pre-movement and movement epochs. Axis limits are ±15 m for all panels.
Figure S3: A. The average value (calculated using dynamic programming) of the states upon which subjects gaze increased as the subject approached the remembered goal location. Specifically, the greater the number of turns remaining in the trajectory (where turns were classified using a threshold applied on angular velocity), the lower the value of the states upon which the subjects looked, partly due to a lower probability of gazing upon or near the goal state (see Figure 6). Values shown here are negative due to the formulation of value as a function of the path length from a state to the target state (the value of the target state is zero). B. During search, subjects spend a greater fraction of time foveating the arena borders (purple) than during the active navigation phase (which consists of both the pre-movement and movement epochs). During navigation (both before and during movement), subjects spend more time foveating the ground (green). While there appears to be a trend in the fraction of time foveating obstacles (orange) vs. arena centrality, this is explained by a higher obstacle density in the more complex arenas. C. Across all subjects and all trials, the probability of gazing upon each obstacle remains relatively constant across all arenas during each epoch. All error bars denote ±1 SEM.
Figure S4: Example simulated trajectories, as well as the gaze samples (red dots, if applicable), taken by each agent. The configuration of the arena reflects the agent’s subjective model at the end of all eye movements. Note that the subjective model of the “smart” agent was still quite mismatched with the true world model after eight eye movements, but the visual samples allowed for the correction of the model at crucial locations such that the trajectory of the “smart” agent was closer to optimal than that of the other agents.
Figure S5: Simulations reveal that foveating ‘relevant transitions’ reduces path length. Results were robust to the precise algorithm (value iteration vs. successor representation) as well as the degree of temporal abstraction (current state vs. optimal trajectory) used to estimate the relevance of transitions. Plots similar to Figure 3c and 3d are shown for relevance values calculated with A value iteration, current state, B value iteration, entire trajectory, C successor representation, current state, and D successor representation, entire trajectory.
**Figure S6:** **A-E:** Breakdown of Figure 4a for arenas 1 (most complex) through 5 (least complex).

**Video S1:** Six representative trials in which subjects exhibited sweeping eye movements. (Top) Aerial view of the arena with the subject’s dynamically evolving position (lilac) and gaze (green). The target is represented as a black circle. (Bottom) Time-evolving version of the plot described in Figure 5b. The video speed is veridical, and the search epoch was omitted from each trial.
Figure S7: A. The fraction of time that subjects spent gazing within 2 m from the trajectory that they took on each trial (excluding points of gaze within 2 m from the goal location) decreased with arena centrality. B. The fraction of trials with sweeps was lower for less complex arenas. Error bars for A and B denote ±1 SEM across subjects. C. Eye movements on each trial was decomposed into fixations within 2 m of the target (green), fixations within 2 m of the subject's trajectory (excluding the target) during sweeps vs. outside of sweeps, and fixations outside of sweeps that were neither made to the target nor trajectory. Subjects viewed the hidden target location more in easier arenas, and gazed upon the rest of the trajectory more in difficult arenas. The fraction of time which was spent looking elsewhere was relatively constant across arenas and epochs. D. Left: Across all subjects and all trials, the speed of backward sweeps (bottom plot) was greater than the speed of forward sweeps (top plot) for all arenas. Before movement (orange), the speed of forward sweeps was faster than that during movement (blue) in the two most complex arenas. Right: Forward sweeps before movement, as well as backwards sweeps before and during movement, were relatively constant in duration across different arenas. However, forward sweeps during movement were longer in the two most complex arenas. E. Left: The average number of saccades per sweep was lower for easier arenas, and was highest during forward sweeps while the subject was actively moving. Right: The saccade rate was higher during sweeps than outside of sweeps. F. The direction of the first sweep was more likely to be backwards if it occurred prior to movement (left), and forwards if it occurred during movement (right). While the fraction of trials with a sweep occurring prior to movement increased as a function of arena difficulty, the fraction of trials for which the first sweep occurred during movement was rather constant for all but the easiest arena. G. Left: The average delay between goal detection and the first sweep increased with arena difficulty. Right: The cumulative distribution of the delays across all subjects and all trials. H. Left: ROC curves constructed as described in Figure 4a (rightmost) for the distributions of true vs. shuffled average relevance values for each trial (pooled across all arenas, all subjects, and all trials), with periods of sweeping eye movements removed, reveals that during the pre-movement (orange) and movement (blue) epochs, non-sequential eye movements are still directed towards task-relevant locations. AUC plots were constructed with sweeps removed, as described in Figure 4b. The AUC values remain well above chance during the pre-movement (orange) and movement (blue) epochs. Right: The same analysis was performed with gaze positions falling within 2 m of the subject’s trajectory on each trial removed, revealing that the remaining visual samples were still made to relevant locations in space.
### Table S1: Median true relevance values (with interquartile range (IQR)) for each arena ($x10^{-3}$).

<table>
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<tr>
<th>Epoch</th>
<th>Arena 1</th>
<th>Arena 2</th>
<th>Arena 3</th>
<th>Arena 4</th>
<th>Arena 5</th>
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<tbody>
<tr>
<td>search</td>
<td>3.3 (6)</td>
<td>5.2 (5)</td>
<td>0 (3)</td>
<td>0 (4)</td>
<td>0 (2)</td>
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<tr>
<td>pre-movement</td>
<td>135 (10)</td>
<td>45 (11)</td>
<td>51 (11)</td>
<td>22 (11)</td>
<td>67 (17)</td>
</tr>
<tr>
<td>movement</td>
<td>184 (8)</td>
<td>36 (10)</td>
<td>38 (9)</td>
<td>3.5 (10)</td>
<td>0 (10)</td>
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### Table S2: The order of arena presentation was randomized across subjects.

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Relevance derivation: In this section, we derive a general measure to quantify the relevance of transitions with respect to the task of navigating between two given states. The following derivation focuses on the general setting when external noise (stochastic transitions) is present and internal noise (model uncertainty) is inhomogeneous. Let $T_k$ denote the status of the $k^{th}$ stochastic transition (1 or 0) and $p_k$ be the parameter of the true probability distribution (p.d.) of that transition such that $P(T_k = 1) = p_k$ and $P(T_k = 0) = 1 - p_k$. Let $\hat{p}_k$ be the parameter of the subjective probability distribution of the transition, i.e., the agent thinks that $P(T_k = 1) = \hat{p}_k$ and $P(T_k = 0) = 1 - \hat{p}_k$. Given a particular goal state, let $V^k_s$ denote the value of the agent's current state $s$ evaluated using the true transition status $T_k$ such that $V^k_s = V_s(T_k = 1)$ if $T_k = 1$ and $V^k_s = V_s(T_k = 0)$ if $T_k = 0$. Let $\hat{V}^k_s$ denote the expectation of the value of state $s$ evaluated using the subjective transition p.d. of the $k^{th}$ transition such that $\hat{V}^k_s = \hat{p}_k V_s(T_k = 1) + (1 - \hat{p}_k) V_s(T_k = 0)$. Since looking at a transition will dramatically reduce the uncertainty about the status of that transition, this can impact the subjective value of the current state, provided that transition is critical to the task at hand. For instance, discovering a subway line linking your neighborhood and downtown will increase the value of your neighborhood if your workplace is located downtown, but will have no impact if your workplace is located crosstown. Therefore, we define relevance ($\Omega_k$) of the $k^{th}$ transition as the expectation of the (log) change in subjective value about the current state $s$ induced by looking at that transition. Then, we have:

$$\Omega_k = \mathbb{E}[\log(|V^k_s - \hat{V}^k_s|)]_{p_k} \text{ where } \mathbb{E}[\cdot|p_k] \text{ denotes expectation taken w.r.t the true p.d.}$$

$$= \mathbb{E}[\log(|V^k_s - \hat{p}_k V_s(T_k = 1) - (1 - \hat{p}_k) V_s(T_k = 0)|)]$$

$$= p_k \log(|V_s(T_k = 1) - \hat{p}_k V_s(T_k = 1) - (1 - \hat{p}_k) V_s(T_k = 0)|) + (1 - p_k) \log(|V_s(T_k = 0) - \hat{p}_k V_s(T_k = 1) - (1 - \hat{p}_k) V_s(T_k = 0)|)$$

$$= p_k \log((1 - \hat{p}_k)|\Delta V|) + (1 - p_k) \log(|\hat{p}_k|\Delta V|) \text{ where } \Delta V = V_s(T_k = 1) - V_s(T_k = 0)$$

$$= p_k \log(1 - \hat{p}_k) + (1 - p_k) \log(\hat{p}_k) + \log(|\Delta V|)$$

$$= (p_k - 1 + 1) \log(1 - \hat{p}_k) - p_k \log(\hat{p}_k) + \log(|\Delta V|)$$

$$= -(1 - p_k) \log(1 - \hat{p}_k) - p_k \log(\hat{p}_k) + \log(1 - \hat{p}_k) + \log(|\Delta V|)$$

$$= H(p_k, \hat{p}_k) + \log(\hat{p}_k (1 - \hat{p}_k)) + \log(|\Delta V|)$$

where $H(X, Y)$ denotes the cross entropy between $X$ and $Y$

$$= H(p_k) + D_{KL}(p_k || \hat{p}_k) + \log(\hat{p}_k (1 - \hat{p}_k)) + \log(|\Delta V|)$$

where $H(X)$ denotes the entropy of $X$

$$= H(p_k) + D_{KL}(p_k || \hat{p}_k) + \log(Var(\hat{T}_k)) + \frac{1}{2} \log(|\Delta V|^2)$$

where $\hat{T}_k$ denotes the subjective knowledge about the status of the $k^{th}$ transition

Observe that $\Omega_k$ is comprised of four factors: (I) $H(p_k)$, the entropy of $p_k$, which captures transition volatility, (II) $D_{KL}(p_k || \hat{p}_k)$, the Kullback-Leibler divergence between true and subjective p.d., which captures the degree of mismatch between the subjective and true transition models, (III) $\log(Var(\hat{T}_k))$, the log variance of the subjective status of the transition, which captures the agent's uncertainty, and (IV) $\log(|\Delta V|^2) = \log(|V_s(T_k = 1) - V_s(T_k = 0)|^2)$, the log change in the value of the current state induced by changing the transition status, which captures the sensitivity of the value function to the transition. The first and third terms suggest that an agent should prioritize looking at transitions with high volatility and high subjective uncertainty. The second term suggests that it is best to look at transitions whose subjective status is known to be wrong. Although this is mathematically correct, agents would not know the true model to begin with and therefore cannot direct their attention at such transitions. Therefore, if external and internal noise are homogeneous, the best strategy would be to look at transitions which the value function is highly sensitive to, as postulated by Equation 1 in the main text. Note that in deriving $\Omega_k$, we have neglected the contribution of model mismatches that may exist at other transitions. This approximation will be valid if the model mismatch is small, and the solution works well in practice as demonstrated by the simulations (Figure 3).