

Current Biology

Locomotion and Task Demands Differentially Modulate Thalamic Audiovisual Processing during Active Search

Highlights

- Auditory thalamic activity is significantly suppressed during movement
- Visual thalamic activity is subtly enhanced, only at high running speeds
- Behavioral relevance modulates activity in visual—but not auditory—thalamus
- Locomotor state and task relevance can be used to improve neural decoding accuracy

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In Brief

Williamson et al. recorded from the auditory and visual thalamus of mice engaged in an audiovisual search task. They find a double dissociation between task relevance and movement, highlighting a role for modulation of thalamic responses by internal state and suggesting key differences in modulatory circuitry between auditory and visual pathways.



Locomotion and Task Demands Differentially Modulate Thalamic Audiovisual Processing during Active Search

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SUMMARY

Active search is a ubiquitous goal-driven behavior wherein organisms purposefully investigate the sensory environment to locate a target object. During active search, brain circuits analyze a stream of sensory information from the external environment, adjusting for internal signals related to self-generated movement or “top-down” weighting of anticipated target and distractor properties. Sensory responses in the cortex can be modulated by internal state [1–9], though the extent and form of modulation arising in the cortex de novo versus an inheritance from subcortical stations is not clear [4, 8–12]. We addressed this question by simultaneously recording from auditory and visual regions of the thalamus (MG and LG, respectively) while mice used dynamic auditory or visual feedback to search for a hidden target within an annular track. Locomotion was associated with strongly suppressed responses and reduced decoding accuracy in MG but a subtle increase in LG spiking. Because stimuli in one modality provided critical information about target location while the other served as a distractor, we could also estimate the importance of task relevance in both thalamic subdivisions. In contrast to the effects of locomotion, we found that LG responses were reduced overall yet decoded stimuli more accurately when vision was behaviorally relevant, whereas task relevance had little effect on MG responses. This double dissociation between the influences of task relevance and movement in MG and LG highlights a role for extrasensory modulation in the thalamus but also suggests key differences in the organization of modulatory circuitry between the auditory and visual pathways.

RESULTS

Mice Can Use Dynamic Audiovisual Feedback to Search for Hidden Rewards

We adapted a closed-loop sensory foraging task [13–15] to explore how behavioral state modulates thalamic sensory pro-

cessing. The task required mice to locate a hidden target region within an annular track to obtain a water reward. Every 0.5 s, mice were presented with pairs of visual flashes or acoustic chirps. The temporal interval separating individual flashes or chirps within the pair switched from long to short as mice moved into the visual or auditory target regions, respectively (Figure 1A). Water reward was contingent upon remaining within the target region for one modality, but not the other, with the choice of modality assigned randomly to each mouse. This arrangement ensured that a change in the inter-pulse interval was task relevant (TR) for one modality but was a task-irrelevant (TI) distraction for the other. After a period of behavioral shaping (see Supplemental Experimental Procedures), a 32-channel silicon probe was implanted into the thalamus at an orientation that enabled simultaneous recordings from the medial geniculate body and dorsal lateral geniculate nucleus of the thalamus (MG and LG, respectively; Figure 1B). This approach allowed us to record from each thalamic subdivision during periods of movement or rest and to contrast responses in mice where vision was TR and audition was TI (Figure 1C) versus mice trained with the opposite stimulus-reward contingency (Figure 1D).

This type of closed-loop active search behavior eschews the rigid structure of conventional psychophysical tasks in favor of an ethologically relevant foraging behavior wherein mice are free to modulate their search speed and movement trajectories according to real-time changes in sensory feedback [16, 17]. On some trials, mice quickly doubled back into the TR target region after crossing through it (Figure 2A), whereas on others, mice circled clockwise and counter-clockwise around the entire track several times before settling on the TR target region (Figure 2B). To test whether search behavior was under stimulus control, we measured the probability that mice would remain within a target region long enough to trip the reward for the TR modality versus pause within the TI target region for an equivalent length of time (TR: visual [$n = 3$]; TR: auditory [$n = 3$]). When initially introduced to the behavioral task, mice were as likely to pause within the TR target region as they were the TI target or a randomly selected region (Figure 2C, left; ANOVA; main effect for task relevance; $F_{2,2} < 0.7$; $p > 0.5$ for both groups). With additional weeks of training, mice exhibited a significant choice bias, such that the probability of pausing within the TR target region was significantly greater than the probability of selecting either the target for the TI modality or a randomly selected region (Figure 2C, right; ANOVA; main effect for task relevance; $F_{2,2} > 15.0$; $p < 0.02$ for both groups). As further evidence that

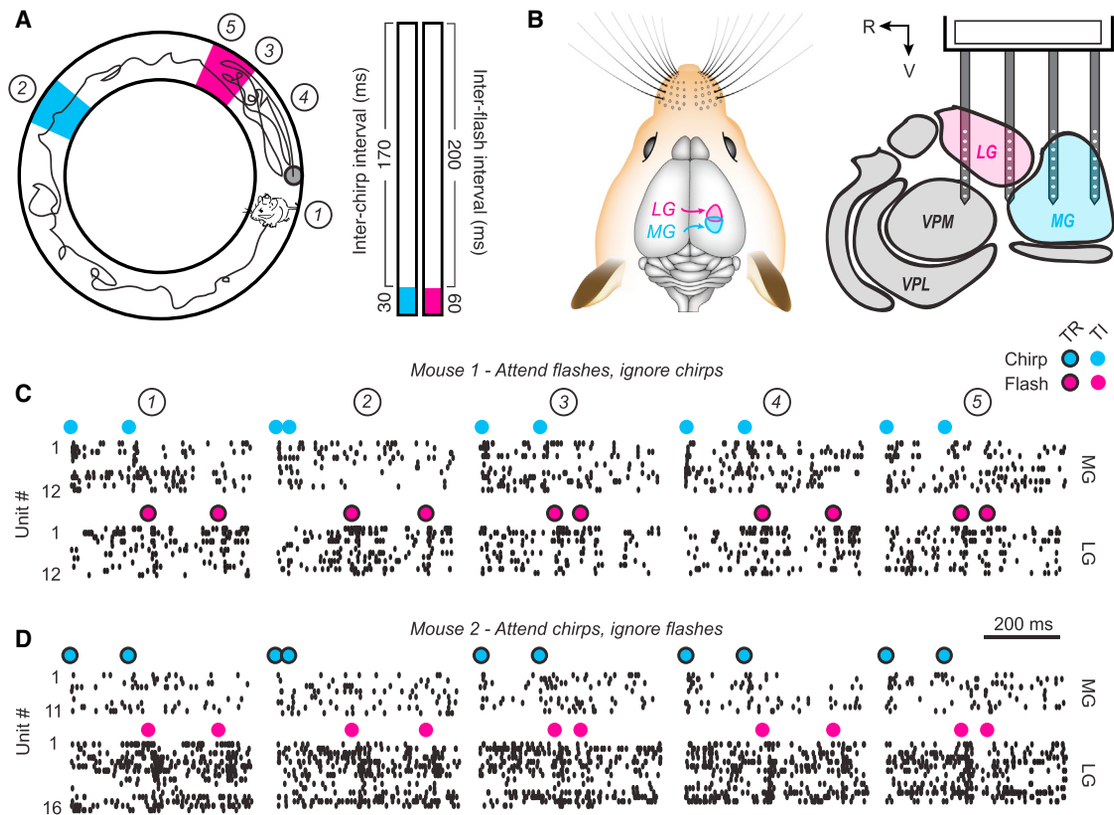


Figure 1. Recording from Auditory and Visual Subdivisions of the Mouse Thalamus during an Active Search Task

(A) Illustration of a mouse's movement path (thin black line) within the annular track during a single behavioral trial. Cyan and magenta areas correspond to short-interval target regions for auditory and visual pulse pairs, respectively. Gray circle represents the water reward spout. Circled numbers correspond to unit recordings below.

(B) The medial geniculate body (cyan; MG) and dorsal lateral geniculate nucleus (magenta; LG) schematized from a dorsal (left) and sagittal (right) perspective. The positioning of the 32-channel probe is shown on the right. R, rostral; V, ventral.

(C and D) Rastergrams compiled from ensembles of 11–16 simultaneously recorded MG (top rows) and LG (bottom rows) units from one mouse trained to associate changes in the visual inter-flash interval with reward (C) and another mouse trained to associate changes in the auditory inter-chirp interval with reward (D). TR and TI denote whether the corresponding modality provides the sole cue to identify the hidden target (TR) or is a distractor (TI). Rastergrams are drawn from five 0.5-s epochs recorded at positions approximately corresponding to the numbered locations in (A).

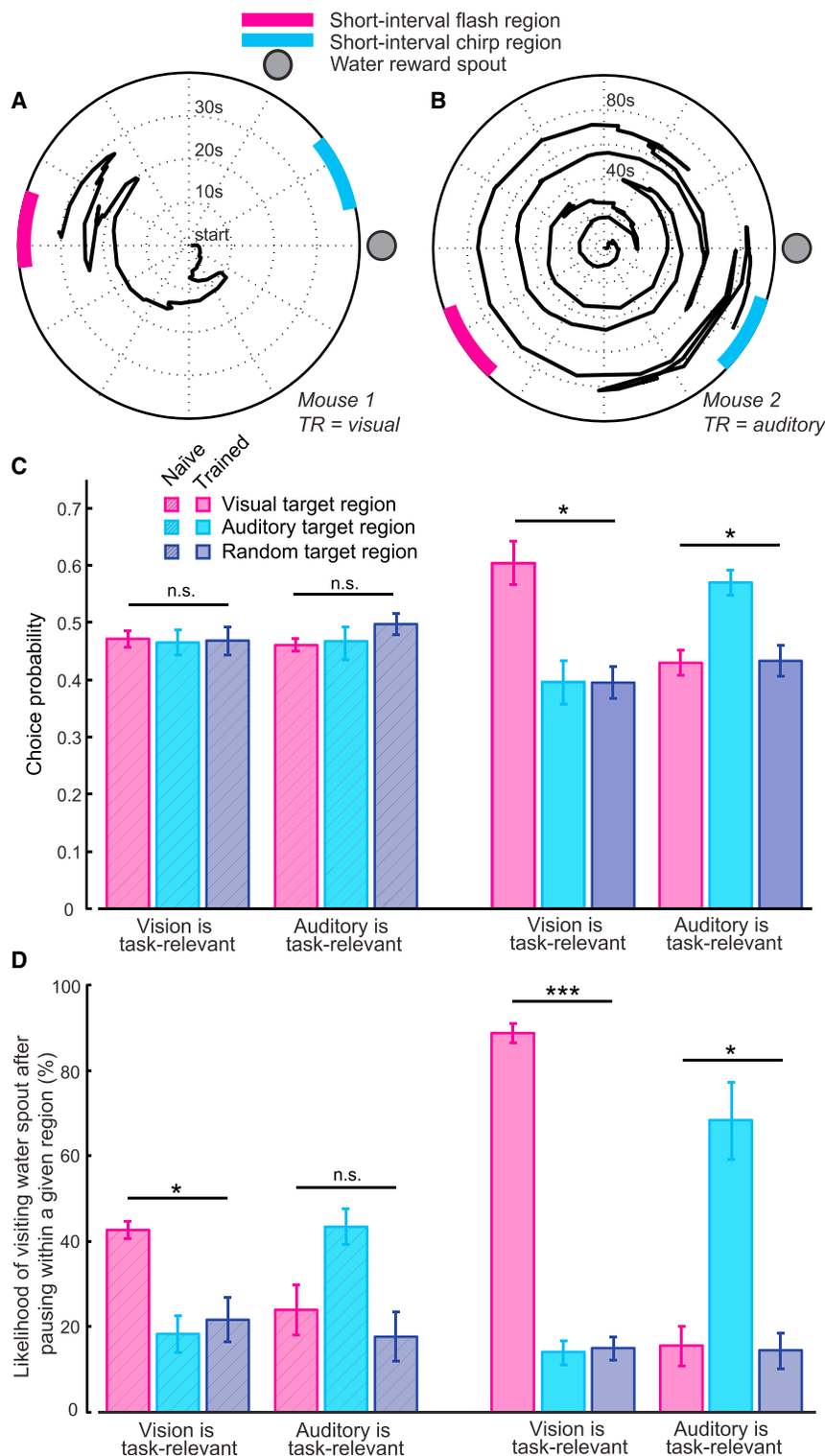
behavior was under stimulus control, we observed that checking the water spout for reward availability was far more likely after pausing within the TR target region than after pausing within the TI target or a random position in well-trained mice (Figure 2D, right; ANOVA; main effect for task relevance; $F_{2,2} > 20.0$; $p < 0.01$ for both groups).

A Double Dissociation in Thalamic Response Modulation by Internal State

The set of behaviors that naturally occur during this task provided us with a means to estimate the influence of internal modulatory signals such as either locomotion or task demands on LG and MG unit activity. Running increases the gain on visually evoked responses in the visual cortex [5, 18–20] but attenuates sound-evoked responses in the auditory cortex [8, 9, 12]. Subcortical antecedents for movement-related response modulation are less clear. Initial characterization of locomotion effects reported no change in sensory-evoked LG or MG responses during periods of movement versus rest [5, 8], though subsequent studies suggest that a comparatively subtle degree of gain and

attenuation may be occurring in the LG and MG, respectively [9, 12, 21].

To clarify the effect of locomotion on thalamic responses, we contrasted LG and MG spike rates during periods of movement and rest. We found that sound-evoked firing rates in MG were reduced in 94.1% of all recorded units ($n = 474$) by an average of 15.1% during movement, whereas spontaneous activity was not affected (bootstrapped ANOVA; main effect for locomotion: evoked, $F_{1,46} = 52.36$, $p < 1 \times 10^{-20}$; spontaneous, $F_{1,46} = 8.15$, $p = 0.17$; Figure 3A). Significant movement-related suppression was noted in recordings from both putative dorsal and ventral subdivisions of the MG (Figures S1A and S1B). By contrast, locomotion was not associated with significant changes in either visually evoked or spontaneous firing rates in the LG (mean change = 2.1% increase; $n = 518$ units; bootstrapped ANOVA; main effect for locomotion: evoked, $F_{1,73} = 3.17$, $p = 0.07$; spontaneous, $F_{1,73} = 2.23$, $p = 0.82$; Figure 3A). Similar movement-related changes in LG and MG firing rates were noted on trials when mice were disengaged from the behavioral task, suggesting that the locomotion effects described here



are consistent with previous descriptions of movement-related modulation reported in head-fixed mice that are not engaged in an explicit task (Figures S1C and S1D) [5, 9, 12, 18–21].

Moving and stationary are but two discrete states along a locomotion continuum. By quantifying the change in firing rate across the full range of observed movement speeds, we noted a mono-

In the present study, all mice operated on the same bottom-up statistics of chirp and flash pairs, but they learned that the temporal interval of one modality provided the sole cue about reward availability whereas the other was an uninformative distraction. Thus, simultaneous LG and MG recordings could be made from mice that regarded vision as the TR modality and audition

Figure 2. Mice Learn to Recognize Changes in Temporal Interval within the Task-Relevant Stimulus Modality

(A and B) Polar plots illustrate the mouse's position on the annular track as a function of time (radial axis, in seconds) relative to visual (magenta) and auditory (cyan) target locations. Each plot depicts a single, representative trial from a mouse where vision was the TR modality (A) and another mouse where auditory was the TR modality (B).

(C) The operational definition of behavioral choice is satisfied when the mouse remained within a single target region for at least 1.5–1.75 s. The probability that mice would initially choose the target region linked to reward or the TI target region is plotted for visual-TR ($n = 3$) and for auditory-TR mice ($n = 3$). Bias for pausing within the TR target region is evident in trained mice, but not in naive mice (hatched).

(D) Mice develop a selective bias for returning to the water spout after pausing within the TR target region. Behavioral data for the naive condition are drawn from the first 20% of trials (C) or the first 100 trials (D). Values reflect mean \pm SEM. Asterisks denote statistically significant differences ($p < 0.05$) with a two-way ANOVA.

tonic increase in sound-evoked suppression with running speed in MG. Surprisingly, LG firing rates were modestly but significantly increased at high running speeds (ANOVA; main effect for running speed: MG, $F_{16,473} = 29.01$, $p = 5.32 \times 10^{-86}$; LG, $F_{16,517} = 3.36$, $p = 6.11 \times 10^{-6}$; Figure 3B). Movement velocities associated with substantive LG firing rate enhancements were uncommon (running speeds that cause $>5\%$ suppression occurred in only 17% of all observations; Figure 3C). Thus, as a first approximation, our findings confirm that movement was not associated with changes in LG firing rate [5], though modest increases were noted at uncommonly high running speeds, in keeping with subsequent findings [21]. By contrast, sound-evoked spiking in the MG is suppressed across the full spectrum of locomotion, reaching levels as high as 30% at the highest running speeds. An analysis of movement-related modulation on single-unit firing rates in LG and MG yielded a nearly identical pattern of results to multi-unit recordings (Figure S1D).

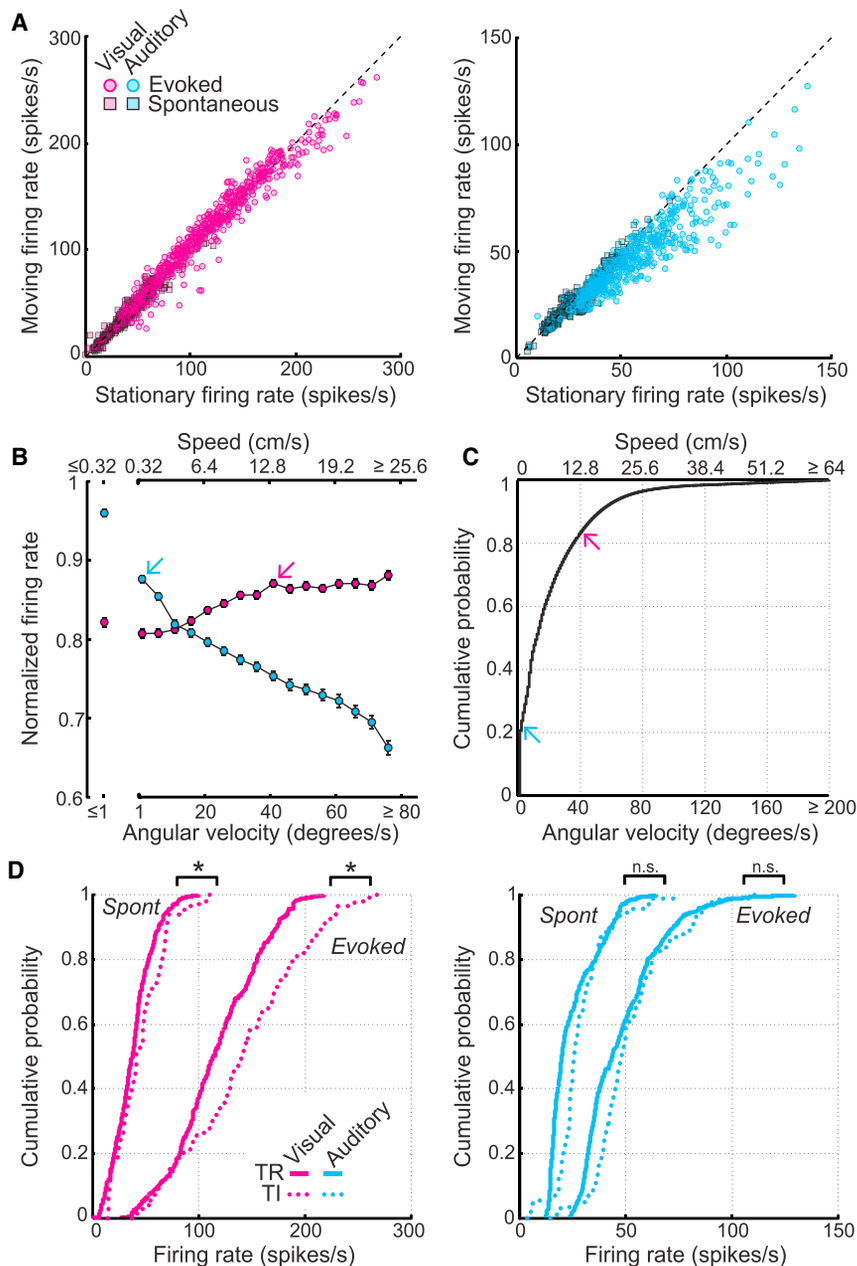


Figure 3. Movement Is Associated with Robust Firing Rate Modulation in MG, but Not LG; Task Relevance Modulates Firing Rates in LG, but Not MG

(A) Scatterplots present the firing rate for all recorded units in LG (left column) and MG (right column). Firing rates were obtained from stimulus-evoked (circles) or spontaneous (squares) PSTH epochs. Each data point is the mean response to both pulses in a given pair for a single recording site averaged across a single behavioral session.

(B) Evoked firing rate as a function of running speed in LG (magenta) and MG (cyan). Firing rates were normalized to the maximum firing rate for each unit. Values reflect mean \pm SEM.

(C) Cumulative fraction of times the animals spent at each particular speed. Arrows depict the lowest running speed associated with firing rate modulation $\geq 5\%$ (B) and the probability that the corresponding running speed occurs (C).

(D) Cumulative firing rate distributions are organized according to task relevance rather than locomotion status for sensory-evoked and spontaneous responses. Asterisks denote statistically significant differences ($p < 0.05$) with a bootstrapped ANOVA.

ditions; Figure 3D, right). However, LG firing rates were suppressed by an average of 17.5% when vision was TR compared to TI, with significant reductions evident in both evoked and spontaneous firing rates (TR, $n = 367$ units; TI, $n = 151$ units; evoked, $F_{1,73} = 16.14$, $p = 0.01$; spontaneous, $F_{1,73} = 12.51$, $p = 0.04$; Figure 3D, left). Thus, as predicted, LG responses were modulated by behavioral relevance whereas MG responses were not. However, the direction of modulation was unexpected in that LG activity was suppressed when vision was TR, not enhanced.

Ensemble Decoding of Stimulus Identity Recapitulates Firing Rate Modulation

These findings highlight a striking double dissociation in modulation of thalamic response by internal states. Locomotion suppressed sound-evoked responses in the MG but weakly enhanced responses in LG;

as the TI modality and another set of mice with reversed TR and TI contingencies. Even though LG units were relatively unaffected by locomotion, changes in LG activity have been reported in the context of other internal state variables such as spatial attention [10, 11, 22]. By contrast, MG firing has been described as comparatively refractory to modulation by “top-down” cognitive signals relating to task demands [4, 23]. Thus, our a priori hypothesis was that firing rates would be increased in the LG—but not MG—when the corresponding modality was TR.

Consistent with this hypothesis, we found that MG firing rates were not significantly changed by the behavioral utility of sound (firing rates were increased by a mean 4.3% in TR versus TI units; bootstrapped ANOVA; $F_{1,46} < 1.65$; $p \geq 0.1$ for evoked and spontaneous contrasts between TR [$n = 382$] and TI [$n = 92$] con-

ditions; Figure 3D, right). However, LG firing rates were suppressed by an average of 17.5% when vision was TR compared to TI, with significant reductions evident in both evoked and spontaneous firing rates (TR, $n = 367$ units; TI, $n = 151$ units; evoked, $F_{1,73} = 16.14$, $p = 0.01$; spontaneous, $F_{1,73} = 12.51$, $p = 0.04$; Figure 3D, left). Thus, as predicted, LG responses were modulated by behavioral relevance whereas MG responses were not. However, the direction of modulation was unexpected in that LG activity was suppressed when vision was TR, not enhanced.

As a first step, we determined the optimal PSTH bin size for decoding visual and auditory pulse timing. With very small bin

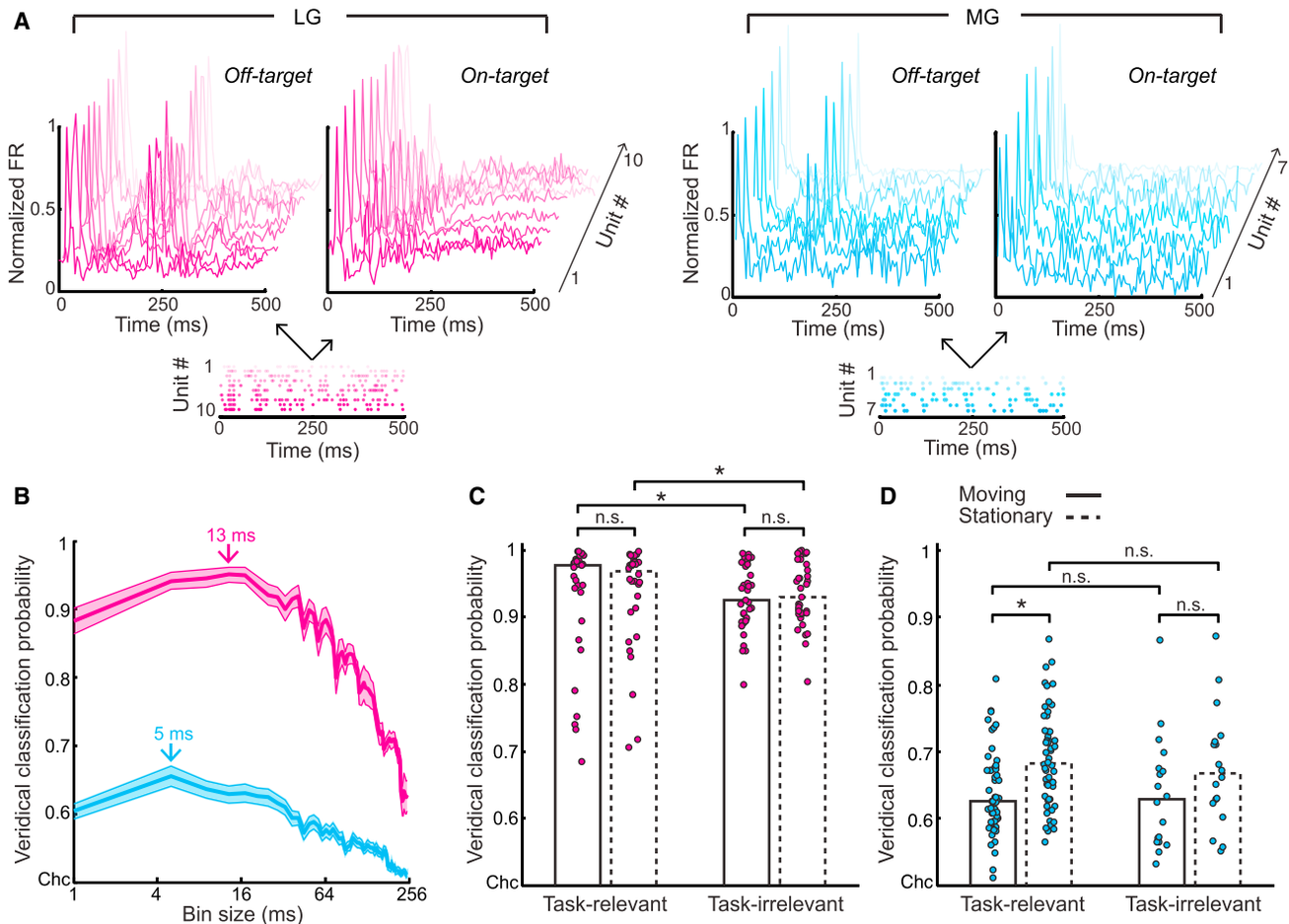


Figure 4. Modulation of Firing Rate by Locomotion and Task Relevance Underlies Differences in Stimulus Decoding Accuracy

(A) Classification of inter-pulse interval based on thalamic ensemble activity recorded during a single behavioral “moment.” Shown here are examples of simultaneously recorded LG ($n = 10$) and MG ($n = 7$) unit ensembles from a mouse in the visual TR condition. PSTH templates for each unit are averaged from a subset of moments where the mouse was in a long interval non-target area (left) or was inside the short interval target (right). The inter-pulse interval of a given chirp or flash pair was decoded by calculating the shorter Euclidean distance separating a single behavioral moment (rastergrams, bottom) from the target and non-target averaged templates. Euclidean distance between a single trial and each template in the mathematical model is proportional to the length of each corresponding arrow. PSTH templates were averaged from the same number of moments for all four conditions (off-target versus on-target and moving versus stationary) in each brain region (MG and LG) for each mouse, using subsampling (see [Supplemental Experimental Procedures](#)). Half of all moments were used to create the corresponding templates, and the other half were used individually for single-moment classification.

(B) Median classification accuracy across all conditions and behavioral sessions as a function of PSTH bin size. Arrows indicate the optimal bin size for MG (cyan) and LG (magenta). Shaded regions = 95% confidence interval; solid lines, median; chance (Chc) classification = 50% accuracy.

(C and D) Inter-pulse interval classification accuracy as a function of task relevance and locomotion for LG (C) and MG (D) ensembles. Each data point represents the mean decoding accuracy from a single behavioral session. Bar height represents the sample median. Asterisks denote statistically significant differences based on permutation tests corrected for multiple comparisons.

sizes (e.g., 1 ms) the internal jitter of spike times degraded the representation of each pulse within the pair. Similarly, larger bin sizes degraded signal-to-noise ratios by incorporating an increasing proportion of spikes that do not directly relate to pulse timing (Figure 4B). Our analysis suggested that pulse rate could be optimally decoded by temporally integrating spikes over a 13-ms window in LG and a 5-ms window in MG, which agrees closely with behavioral inter-pulse interval discrimination threshold values following direct activation of the central visual or auditory cortex, respectively [24].

We then used these optimized bin sizes to contrast differences in inter-pulse interval decoding accuracy within a daily

behavioral session as a function of locomotion state and stimulus task relevance. For the most part, differences in decoding accuracy recapitulated the double dissociation in firing rates, although the sign of firing rate change (increase or decrease) was not directly linked to classification accuracy. In the LG, movement had no effect on classification accuracy (permutation test: TR versus TI [$n = 32$ and 35 behavioral sessions, respectively]; $p > 0.8$ for moving versus stationary for TR and TI conditions; Figure 4C). Although LG firing rates were reduced when vision was TR, the spiking patterns supported a more-accurate classification of visual flash interval (permutation test; $p = 0.01$ for TR versus TI for both moving and stationary

conditions; Figure 4C). Conversely, classification accuracy in MG was not affected by task relevance (permutation test; $p > 0.5$ for both TR versus TI [$n = 54$ and 18 behavioral sessions, respectively] in moving and stationary conditions; Figure 4D), but the reduced firing rates during movement were associated with lower accuracy in decoding the chirp interval. This difference reached statistical significance in the TR condition, but not in the TI condition (permutation test; $p = 0.0001$ and $p = 0.15$, respectively).

DISCUSSION

We recorded from auditory and visual subdivisions of the thalamus as mice used closed-loop audiovisual feedback to navigate around an annular track in search of a hidden target. The design of the task enabled us to measure whether thalamic sensory responses were modulated by non-sensory signals related to internal state. Previous studies in head-fixed mice placed atop a movable platform demonstrated that locomotion augments visually evoked responses in the visual cortex [5, 20, 25] but attenuates sound-evoked activity in auditory cortex [8, 9, 12]. Evidence on the subcortical origins of these effects are mixed with some studies reporting no evidence of locomotion effects in LG or MG [5, 8] and more-recent reports suggesting that locomotion can impart a comparatively subtle augmentation or suppression in the visual and auditory thalamus, respectively [9, 12, 21]. Our data feature simultaneous recordings from both LG and MG and are unequivocal on this point: movement is associated with a subtle increase in LG responses only at uncommonly high running speeds and has no effect on temporal interval decoding, whereas sound-evoked MG responses are strongly suppressed and decode stimulus identity less accurately across a broad range of running speeds. Importantly, movement-related modulation of thalamic activity in either subdivision is less robust than what has been reported in primary sensory cortex, underscoring the likely involvement of additional intracortical circuits that mediate more-extensive response modulation in downstream processing [8, 9, 12, 20, 21].

At the level of the cortex, sensory traces have been largely reformatted into an abstraction of the source signal [26]. This spatiotemporal pattern of spikes continues to encode relevant features of the stimulus but is also powerfully modulated by non-sensory factors such as emotion, learning, attention, and motor planning [3, 6, 7, 27–31]. Whereas the neural circuitry for extrasensory modulation is exquisite and comparatively well understood in sensory neocortex [9, 12, 20, 25, 32–36], additional modulatory networks in the thalamus [37] or midbrain [38] could provide a means for rapid and flexible adjustments of subcortical auditory [23, 39–41] and visual [10, 42, 43] processing depending on task demands, attentional load, and learning. Here, for example, we report reduced visually evoked responses in LG that nevertheless more accurately encode inter-flash interval when visual inputs are relevant to solving the behavioral task compared to when they are a distraction. The extent to which internal state modulation of subcortical responses is mediated through subcortical modulatory networks versus descending corticofugal modulation remains a promising area for further investigation.

EXPERIMENTAL PROCEDURES

Audiovisual Search Task

All procedures were approved by the Animal Care and Use Committee at Massachusetts Eye and Ear Infirmary and followed the guidelines established by the NIH for the care and use of laboratory animals. Six male C57/BL6 mice, aged 6–8 weeks, were maintained above 80% of their pre-training body weight. During training, mice received their daily water allowance through the behavioral task with additional supplements as needed.

Detailed information on the behavioral task, chronic thalamic recordings, and data analysis can be found in the [Supplemental Experimental Methods](#).

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, one figure, and one table and can be found with this article online at <http://dx.doi.org/10.1016/j.cub.2015.05.045>.

AUTHOR CONTRIBUTIONS

All authors contributed to experimental design and manuscript editing. R.S.W. collected and analyzed all data. D.B.P. and R.S.W. wrote the manuscript. K.E.H., R.S.W., and D.B.P. developed hardware and software control for the behavioral neurophysiology experiment.

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