Title: Thalamic state influences timing and feature selectivity in the thalamocortical circuit

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**Summary**

Thalamic neurons dynamically encode sensory information in a state-dependent manner as a mechanism for gating information flow to the cortex. Here, we investigated the role of thalamic state on precise feature selectivity in the thalamocortical circuit of the rat vibrissa pathway. In thalamic neurons, tonic spike triggered averaging revealed clear feature selectivity, while the feature selectivity associated with burst spikes could not be recovered with this approach. These thalamic state dependent changes propagated to cortex such that the cortical feature selectivity was diminished during the optogenetically hyperpolarized (burst biased) thalamic condition. Further analysis revealed that the perceived loss of feature selectivity was likely not due to a true loss of stimulus selectivity but instead to changes in the precision of the temporal spiking in burst firing modes. Therefore, alterations to thalamic state enable a dynamic interplay between spike timing and spike rate that shapes stimulus encoding in the thalamocortical circuit.
Introduction

Sensory thalamus plays a critical role in gating information flow from our sensors in the periphery to sensory cortex, ultimately shaping how we perceive the world. Importantly, thalamic gating properties are not static, but instead vary dynamically through a range of modulatory mechanisms, including local membrane and synaptic properties (Wolfart et al., 2005), stimulus history (Whitmire et al., 2016), and neuromodulatory inputs from brainstem and cortex (Castro-alamancos, 2002; Mease et al., 2014). Although arising from different mechanisms, these modulatory inputs have the net effect of altering the baseline membrane polarization level in the thalamus, which we refer to here as “thalamic state”, which plays an important role in determining the encoding properties of the thalamic neurons that serve as primary inputs to sensory cortex. Perhaps most prominently, modulation of the baseline membrane potential in thalamic neurons enables distinct tonic and burst firing modes due to the selective engagement of low threshold calcium channels during prolonged hyperpolarization (Suzuki and Rogawski, 1989). In addition to their roles in thalamocortical oscillations (Steriade et al., 1993), it has long been posited that these two firing modes could be a mechanism for dynamically controlling information processing (Sherman 2001). At the thalamocortical synapse, spontaneous burst spikes are more effective at driving cortical spiking (Swadlow and Gusev, 2001) and evoke larger cortical depolarizations (Bruno and Sakmann, 2006) than tonic spikes. It has been proposed that both burst and tonic spikes carry stimulus information (Reinagel et al., 1999), but the relationship between burst and tonic firing in representing temporal stimulus information in tactile encoding remains unclear.

In the visual pathway, the role of thalamic state in sensory processing has identified distinct sensory selectivity associated with burst versus tonic firing (Alitto et al., 2005; Denning and Reinagel, 2005; Lesica and Stanley, 2004; Reinagel et al., 1999; Wang et al., 2007). The burst response can be reliably elicited across trials in response to sensory stimulation (Lesica and Stanley, 2004; Martinez-Conde et al., 2002; Wang et al., 2007) and the feature selectivity of the bursts are characterized by a prolonged inhibitory stimulus before the depolarizing input that occurs immediately prior to the spike onset. This demonstrated that bursting activity is not
just an indicator of state, but it is also repeatably evoked by the sensory stimulation and is present in the awake animal (Guido and Weyand, 1995). However, how these results extend to other sensory modalities, such as touch, and the implication for downstream sensory encoding remains unknown.

Here, we quantified the role of thalamic state on temporal feature selectivity in the thalamocortical circuit of the rodent whisker pathway by implementing a class of models that maps sensory inputs to observed neuronal activity. Specifically, we characterized neurons using a two-stage, linear-nonlinear framework, the first stage of which represents the sensory feature selectivity, and the second stage of which represents the overall sensitivity of the input-output relationship. This characterization was performed for neurons recorded extracellularly both in the ventro posterior-medial (VPm) thalamus and in layer 4 of primary somatosensory cortex (S1) in the fentanyl-anesthetized rat. For thalamic neurons, we found that tonic spiking was associated with clear whisker-stimulus feature selectivity consistent with previous findings (Petersen et al., 2008). However, analysis of burst firing suggested a lack of feature selectivity, a finding which was further confirmed using optogenetic hyperpolarization of VPm to switch the thalamus into a burst state. An assessment of the temporal precision of the sensory-driven thalamic firing identified an increase in the timing jitter of burst spikes relative to tonic spikes that could underlie the differences in the recoverability of the feature selectivity. In the cortical neurons, we found that when thalamus was dominated by tonic firing, the cortical neurons exhibited similar feature selectivity as observed in VPm. However, when the thalamus was optogenetically hyperpolarized to push the local thalamic population further into a burst firing mode, this had a detrimental effect on the spike time precision of the downstream cortical neurons as well as the feature selectivity reflected by the model. Given the sensitivity of the thalamocortical circuit to precise timing of thalamic projection neurons, the results here suggest an important relationship between thalamic state, or membrane polarization, and the dynamic regulation of timing fidelity across thalamic firing modes that could have profound implications for coding during varying behavioral conditions.
**Methods**

**Experimental Procedures**

**Acute Surgery:** All procedures were approved by the Georgia Institute of Technology Institutional Animal Care and Use Committee and were in agreement with guidelines established by the National Institutes of Health. 19 female albino rats (Sprague-Dawley, 250-300g) were anesthetized intravenously using a fentanyl cocktail (fentanyl (5 µg/kg), midazolam (2 mg/kg), dexmedetomidine (150 µg/kg)). A craniotomy was performed over VPm (2-4 mm caudal to bregma, 1.5-3.5 mm lateral to the midline), and in a subset of animals, a second craniotomy was performed over S1 (1-3 mm caudal to bregma, 4.5-6 mm lateral to the midline). At the termination of the experiment, the animal was euthanized with an overdose of sodium pentobarbital (euthasol, 0.5 mL at 390 mg/mL). All optogenetically transfected animals that underwent cortical probe recordings were perfused and their brains were imaged for verification of opsin location and cortical probe location.

**Electrophysiology:** Tungsten microelectrodes were lowered into the thalamus (depth: 4.5-6 mm) using either a micropositioner (Kopf, Luigs-Neumann). Multi-electrode probes (A1x32-10mm-50-177, NeuroNexus) were lowered perpendicular to S1 (45° relative to vertical; depth: 2 mm). The topographic location of the electrode was identified through manual stimulation of the whisker pad. Upon identification of the primary whisker for the recorded unit(s), the primary whisker was threaded into the galvo motor to permit stimulation of a single whisker.

**Sensory Stimulus:** Mechanical whisker stimulation was delivered using a precisely controlled galvo motor (Cambridge Technologies, custom Matlab software). The mechanical stimulus applied to the whisker in the rostral-caudal direction consisted of sensory white noise (low pass filtered at 200 Hz, standard deviation of the noise was 0.6° or 223°/s). Feedback from the whisker stimulator were used for further spike triggered analysis across all units (down sampled to 4.88 kHz).
**Optogenetics surgeries:** All surgical procedures followed sterile protocol. A small craniotomy was made above VPm (3 mm lateral, 3 mm caudal to bregma). A 10 µL syringe (Neuros Syringe, Hamilton, Inc) filled with the virus (rAAV5-CamKIIa-Jaws-KGC-GFP-ER2 or rAAV5-CamKIIa-eNpHR3.0-EYFP, UNC Viral Vector Core Services) was lowered to depth of 5.2 mm before injecting 1 µL of virus at a rate of 0.2 µL/min (iSi system, Stoelting). The syringe remained in place for five minutes after the injection was complete to allow the virus to diffuse. Opsin expression was fully realized at 2-3 weeks post-surgery.

**Optogenetic Stimulus:** Optical manipulation was administered with a controlled pulse of light through a custom optrode consisting of an optical fiber (200µm diameter; Thorlabs) and an electrode (Tungsten microelectrode; FHC) that was lowered into the VPm. Upon identifying a whisker sensitive cell, light sensitivity was assessed by the post-inhibitory rebound spiking response using a train of 250 millisecond light pulses (λ = 590 or 617nm for Halorhodopsin and Jaws, respectively). The whisker was then stimulated without (baseline) and with (hyperpolarized) light provided directly to the thalamus (50 mW/mm²). Optogenetic stimulus conditions (light on/hyperpolarized, light off/baseline) were interleaved to avoid long-term adaptation effects.

**Analytical Methods:** Spike sorting for single channel recordings was performed online and validated offline using Waveclus (Quiroga et al., 2004). Spike sorting for multichannel electrodes was performed offline using the KlustaKwik software suite (Rossant et al., 2015). Isolation of the unit was confirmed by the waveform amplitude (absolute and relative to the background noise >3) and the interspike-interval distributions (VPm: mean of 0.22%, S1: mean of 0.38% of spikes in absolute refractory period of 1ms).

Feature selectivity was estimated for each recorded unit using the spike triggered average (STA) (Schwartz et al., 2006).

\[
STA = \frac{1}{N} \sum_{j} s(t_j)
\]
Where $N$ is the number of spikes and $s$ is the stimulus segment in a window surrounding each spike (-30 to +5 ms, spike-triggered ensemble, STE). The burst and tonic triggered averages were computed from burst and tonic spikes, respectively. The baseline/hyperpolarized condition triggered averages were computed from all spikes in a given stimulus condition. The bootstrap estimate of the confidence intervals on the spike triggered average was computed as the +/- 2 standard deviation of this shuffled STA distribution across 500 repetitions (Schwartz et al., 2006). Note that we implemented multiple techniques of estimating the feature selectivity of the neurons including spike triggered covariance, generalized linear models, and nonlinear-input models (McFarland et al., 2013). The results were qualitatively consistent across all methods employed, so we chose to use spike triggered average throughout the manuscript due to its simplicity.

The signal-to-noise ratio of the recovered STA was quantified as the peak-to-peak amplitude of the STA within 10 milliseconds of the spike (where the significant filter activity is contained) divided by the peak-to-peak amplitude of the STA from 30 to 20 milliseconds before the spike (where there is no expected filter information). An SNR value of 1 means the amplitude of the STA near the spike time is not different from the amplitude of the noise fluctuations. Therefore, any units with an SNR value less than 2 were excluded from further analysis.

To make comparisons of the feature selectivity across the population of recorded neurons, we computed a principle component analysis of the recovered STA (Estebanez et al., 2012). The first two principle components accounted for the majority of the variance (71.8% VPm, 78.4% S1).

The non-linearity ($P(y|spike)$) was estimated as the ratio of the probability of spike-triggered stimuli ($P(y|spike)$) to the probability of any stimulus segment in the stimulus ($P(y)$) multiplied by the mean firing rate of the neuron($P(spike)$) (Schwartz et al., 2006):

$$p(spike|y) = p(spike) \frac{p(y|spike)}{p(y)}$$
Where $y$ is defined as the stimulus ($s$) convolved with the feature selectivity of the unit (STA) (Lesica et al., 2007), referred to as filtered stimulus. For all conditions, the STA was defined as the baseline or tonic spike triggered average. Throughout the manuscript, we separate the firing rate ($p(spike)$) from the shape of the non-linearity ($p(y|spike)/p(y)$) to avoid confounding differences in firing rate with differences in tuning.

The precision in the noise evoked firing was estimated for each spike classification (tonic, burst, baseline, hyperpolarized). The precision was defined as:

$$precision = \sum_{n=-1}^{+1} \tau_{jitter}(n)$$

Or the number of spikes with $\tau_{jitter}$ values of +/- 1 ms duration normalized by the total number of spikes. $\tau_{jitter}$ is defined for each spike as the temporal lag ($t_{lag}$) for the peak correlation between the STA and the stimulus segment surrounding that spike (-30 ms to +5 ms).

$$\tau_{jitter} = \arg \max_{t_{lag}} \text{correlation} \left( \text{STA}, s(t_j) \right)$$

The $\tau_{jitter}$ distribution was normalized by the total number of spikes in each condition (tonic, burst, baseline, hyperpolarized).

All pairwise statistical comparisons were computed using a Wilcoxon signed rank test unless otherwise noted.
Results

Feature selectivity is conserved across the thalamus and cortex

We recorded thalamic and cortical extracellular spiking activity in response to sensory white noise stimulation of a single whisker in the vibrissa pathway of the fentanyl-anesthetized rat (Figure 1A, see Methods). We estimated the feature selectivity for each unit as the spike triggered average (STA), which captures the features of the sensory stimulus that tended to precede spiking, and the static, point nonlinearity, which captures the translation into suprathreshold spiking activity (Figure 1B; see methods). Although this quantification was performed on longer unique noise segments, we also recorded the response to short (4-10 second) frozen white noise segments to examine the response across trials. Figure 1C shows an example recording from a simultaneously recorded pair of neurons in topographically aligned regions of the thalamus (left column, ventral posteromedial nucleus, VPm) and cortex (right column, primary somatosensory cortex, S1) in response to the repeated presentation of a single frozen white noise segment (top of each column). Across trials, the repeatability of the response to the noise stimulus is apparent in the raster plot, with clear vertical patterns across trials. The STA was computed for the thalamic and cortical unit for stimulus segments from -30 milliseconds prior to the spike to +5 milliseconds afterwards at a 0.2ms resolution (Figure 1C, bottom. The black dotted line indicates the time of the spike). The VPm STA shows clear feature selectivity in the 10-15 milliseconds prior to the thalamic spike as evidenced by the large amplitude of the STA relative to the shuffled case (grey confidence intervals). Beyond 15 milliseconds prior to the spike, the VPm STA is essentially flat and within the confidence bounds on the shuffled process. This suggests that, on average, the thalamic unit is only sensitive to the stimulus occurring in the previous 10-15 milliseconds. The S1 unit also displays feature selectivity as evidenced by the shape and amplitude of the S1 STA immediately prior to the cortical spike relative to the shuffled case. Although the VPm STA is nearly ten times as large in amplitude as the S1 STA, the similarity in the temporal dynamics can be visualized by shifting the VPm STA by 2 milliseconds relative to the S1 STA (Figure
While this simple comparison provides an interesting observation for a single pair of topographically aligned neurons, we also made comparisons of the feature selectivity across
the population of recorded STA for thalamus and cortex. First, we visualized the shape of the
spike triggered average for a sample of example thalamic (Figure 1D left; greyscale) and
cortical (Figure 1D right; greyscale) units. These STA filters cannot be simply averaged
together to give an estimate of the population filter due to differences in the phase and
directionality of the recovered STA across different recorded units. Instead, we performed a
principle component analysis on the set of recovered thalamic and cortical STA filters across
recordings to identify salient filter properties that generalized (Figure 1E). The first two
principle components for the spike triggered averages of both thalamus and cortex explain the
majority of the variance for the set of recovered filters, similar to what has been seen previously
for cortex (Estebanez et al., 2012). Furthermore, a simple time shift of 2.5 milliseconds for the
VPm principle components relative to the S1 principle components (Figure 1F, dashed lines)
demonstrates the similarity in the STA subspace spanned by these principle components. It
seems that despite not necessarily being recorded simultaneously or even in the same animal,
there is a high degree of overlap in the low dimensional subspace of feature selectivity for
thalamocortical neurons in the whisker pathway.

Tonic and burst spike feature selectivity in thalamus

Inherent in the spike triggered analysis, however, is an assumption that the average
filter is representative of the sensory stimulus preceding all spikes (Stanley, 2002). Yet
neurons in the thalamus are well known for exhibiting two fundamentally different types of
firing: tonic spiking and burst firing mediated through T-type calcium channels (Suzuki and
Rogawski, 1989). Burst spikes were classified here from the extracellular recordings as two
or more spikes with an inter-spike interval of less than four milliseconds with the first spike in
the burst preceded by 100 milliseconds of silence (Figure 2A, see methods). Using this
classification, we asked if or how the feature selectivity of an individual thalamic unit changes as a function of the spiking mechanism in the whisker pathway.

**Figure 2: Feature selectivity of burst and tonic spikes in thalamic neurons.**

A. Burst definition. B. STA from thalamic unit presented in Figure 1 estimated from all spikes (black solid line, n = 44105 spikes), tonic spikes (tSTA; grey solid line, n = 36558 spikes), a subsample of tonic spikes (tSTA; dashed line, n = 2363 of 36558 spikes) all burst spikes (bSTA; red solid line, n = 7547 spikes), and the first spike in the burst (bSTA; red dashed line, n = 2363 spikes). C. STA\textsubscript{SNR} across recorded population (n = 30 units). Black dots depict the STA\textsubscript{SNR} for all bSTA computed from all burst spikes while red circles indicate the STA\textsubscript{SNR} for bSTA computed from only the first spike in the burst (p = 1e-6). Arrow depicts example unit in A. D. Example unit nonlinearity. E. Average non-linearity across all units. F. Burst and tonic firing rate (p(spike)) across recorded population (p = 2.6e-5). G. Spike triggered average using different spike classifications.
In the thalamic recordings, tonic and burst spikes were interspersed throughout most of the recordings. For the example thalamic unit presented in Figure 1B, we computed the spike triggered average from all spikes (STA), the tonic spike triggered average from only tonic spikes (tSTA), and the burst spike triggered average from only spikes that are classified as being part of a burst (bSTA) (Figure 2B). The tSTA (grey) closely resembles the STA computed from all spikes (black) while the bSTA (red) is significantly degraded as evidenced by the flat shape of the filter. To compare the difference between burst and tonic feature selectivity across thalamic units, we quantified the signal-to-noise ratio of the STA (STA_{SNR}, see methods). Across all thalamic units, the SNR_{STA} was higher for tonic spikes (tSTA_{SNR}) than for burst spikes (bSTA_{SNR}) (Figure 2C).

Given the estimated feature selectivity, we can compute the static non-linearity, or the input-output function, which provides a mapping between this filtered stimulus ($y$) and the spiking response of the neuron ($p(\text{spike}|y)$) by taking the ratio of the $p(y|\text{spike})$ to the $p(y)$ (Figure 1B, see methods). Here, we used the tSTA as the filter for all spiking conditions when estimating the non-linearity. The probability of the filtered stimulus ($p(y)$) remains unchanged when the filter is held constant. Therefore, any change in the non-linearity is then only due to changes in the probability of the filtered stimulus given that a spike occurred ($p(y|\text{spike})$), or the spike triggered ensemble. Because the slope of the static non-linearity is determined by the separation between the spike triggered ensemble and the Gaussian distributed white noise, as the spike triggered ensemble distribution becomes more selective (i.e. the mean moves away from the filtered stimulus distribution), the separability of the distributions increases, and the slope of the non-linearity also increases. Intuitively, this means that the shape of the non-linearity gives an estimate of the separability of the spike triggered ensemble and the stimulus distribution, or how strongly tuned a neuron is for that particular feature, given by the STA. A steeper slope in the non-linearity suggests a stronger tuning than a shallower slope. Therefore, we also assessed the spiking nonlinearity as a function of the spike classification. In this example unit, we found that the tonic spikes were well tuned to the STA,
as evidenced by the steep slope of the non-linearity while the burst spikes were not well tuned
to the STA, as evidenced by the relatively flat non-linearity (Figure 2D). This trend was
consistent across units where the burst spikes showed reduced tuning to the STA as
compared to tonic spikes as assessed by the slope of the spiking nonlinearity (Figure 2E).
Here, we have separated the difference in the slope of the non-linearity from the difference in
the prevalence of burst and tonic spikes (p(spike)), which is markedly higher for tonic spikes
than for burst spikes (Figure 2F). Furthermore, we tested alternative burst spike classifications
and quantified the implication for the STA (Figure 2G, top). Across spiking classifications,
increased periods of silence prior to the spike (t_{silence}) led to decreased STA_{SNR} while bursts of
spikes (t_{sil} < 4 or <10) had consistently lower STA_{SNR} relative to tonic spikes (t_{sil} > 20) (Figure 2G,
example unit in middle, population data in bottom). Therefore, our data do not provide
evidence to support a difference in feature selectivity for tonic and burst spiking in this
pathway, but instead suggests a reduction in stimulus selectivity in burst spiking within this
analytic framework.

Thalamic state dependent feature selectivity

The previous analysis was conducted by presenting sensory white-noise stimuli and
parsing measured thalamic spiking activity into tonic and burst classes, while these classes of
spiking were intermingled throughout the recordings. However, the thalamus was in tonic firing
mode, with relatively low burst firing rates (Figure 2F). Here, we used optogenetic
hyperpolarization of the thalamic neurons not to silence the thalamic neurons, but instead to
shift the thalamus into a burst firing mode during sensory white noise stimulation (Figure 3A).
Using this optogenetic manipulation, we asked whether the optogenetically manipulated firing
mode (baseline and hyperpolarized conditions) of the thalamus impacts feature selectivity and
how this relates to the classified burst/tonic modes.
Here, we recorded the thalamic response to sensory white noise with and without the presence of a light stimulus (hyperpolarized and baseline conditions, respectively) to shift the firing mode of the thalamus towards burst firing (baseline burst ratio = 0.16 ± 0.15, hyperpolarized burst ratio = 0.36 ± 0.27, n = 10 units). For an example unit, we have plotted Figure 3: Optogenetic Manipulation of Thalamic State. A. Experimental paradigm. Scale bar: 1 mm. B. Example thalamic response to frozen white noise segments without (baseline) and with (hyperpolarized) optogenetic stimulation. C. Spike triggered average computed as a function of spike classification (burst, tonic) or thalamic state (baseline, hyperpolarized). D. Example unit tonic/burst and baseline/hyperpolarized spike nonlinearity. E. STA_{SNR} across recorded population (n = 11 units). Arrow depicts example unit in B. F. Average non-linearity across all units. G. Baseline and hyperpolarized firing rate (p(spike)) across recorded population (p>0.05).
the spiking response to a frozen white noise segment without optogenetic stimulation (Figure 3B, baseline condition) and with optogenetic stimulation (Figure 3B, hyperpolarized condition). We have pseudocolored the tonic spikes grey and the burst spikes red to qualitatively visualize the thalamic firing mode (Figure 3B). In the baseline condition, the response is primarily tonic as evidenced by the grey raster plots (Figure 3B, Baseline, BR = 0.10). In the hyperpolarized condition (optically stimulated), the firing mode is biased towards a burst encoding scheme, as evidenced by the prevalence of red burst spikes (Figure 3B, Hyperpolarized, BR = 0.67). The tonic STA showed pronounced feature selectivity for this unit while the burst STA did not (Figure 3C top), consistent with the earlier findings (Figure 2). In the optogenetically manipulated states, the baseline STA has prominent feature selectivity while the hyperpolarized condition is much smaller in amplitude (Figure 3C bottom). Qualitatively, we can see that the STA from hyperpolarized condition reflected the STA obtained from the burst spiking in the previous analysis.

The similarity between the burst spike response and hyperpolarized condition can also be seen in this example nonlinearity where the burst and hyperpolarized nonlinearities are effectively flat while the tonic spikes and baseline condition show obvious tuning (Figure 3D). Across units, we found an overall reduction in the STA_{SNR} for the hyperpolarized condition relative to the baseline condition (Figure 3E, p = 0.037). We also found that the tuning of the nonlinearity was lower for the hyperpolarized condition relative to the baseline condition as reflected in the overall gain/slope (Figure 3F). Importantly, the baseline and hyperpolarized conditions both contain burst and tonic spikes. Instead of completely separating the firing modes into all burst spikes or all tonic spikes, we have optogenetically altered the spiking probabilities such that the baseline condition has more tonic spikes and the hyperpolarized condition has more burst spikes while maintaining similar numbers of spikes (Figure 3G). The similarities between the STA and the NL properties of the burst and hyperpolarized state as well as the tonic and baseline state suggest that there was no discernable difference for the
estimation of feature selectivity when assessed based on the state of the thalamus at the time
of the stimulus (hyperpolarized/baseline) versus the spike type classification (burst/tonic).

Temporal precision of thalamic firing modes

Given the difference between the recovered estimates of burst/hyperpolarized and
tonic/baseline feature selectivity, we implemented a series of computational controls to identify
any potential shortcomings of the methodologies that could underlie these results. The first
issue we considered was whether or not the burst spike feature selectivity was unrecoverable
due to the effect of subsequent spikes in the burst. If the timing of spikes within a burst is not
repeatable and structured, the presence of these additional spikes will serve to destroy the
temporal structure in the feature selectivity as revealed by the spike triggered analysis. When
the bSTA was computed from only the first spike in each burst (Figure 2B, red-dashed line),
there was no apparent feature selectivity for this example unit. This can also be visualized
across units in the STASNR where the bSTASNR is plotted when computed from all burst spikes
(black dot) and when computed from the first spike in each burst (red circle, Figure 2C).
Therefore, including all spikes in a burst (or not) does not strongly impact the ability to estimate
the feature selectivity from the STA.

The second issue we considered was the overall difference in spike rates. Spike
triggered analyses require a large number of spikes to effectively estimate the underlying
selectivity. The proportion of spikes classified as bursts was lower than the spikes classified
as tonic (Figure 2F) as quantified by the burst and tonic firing rate. Therefore, it was possible
that we could not recover an STA for the burst spike condition due to the reduced number of
burst spikes relative to tonic spikes. In an example unit, we computed the tSTA using only a
subset of the spikes (n = 2362 of 36558 spikes corresponding to n = 2363 bursts with n =
7547 burst spikes) and found that the linear filter was essentially identical to the tSTA (Figure
2B, grey dashed line). We computed this for all thalamic units and again found that the burst-
count matched tSTA was also significantly larger than the bSTA. Furthermore, there was no
statistically significant difference in the firing rate between the baseline and hyperpolarized optogenetic conditions (Figure 3G), but still the difference in the STA persisted. This suggests that simple spike counts alone were insufficient to explain the difference in the tonic/baseline STA and the burst/hyperpolarized STA.

The third issue we considered was the inherent assumption that the feature selectivity for each unit could be recovered as the STA. It was possible that the burst STA was not recoverable because the burst firing mode was better estimated by a symmetric nonlinearity and therefore the filter could only be recovered using spike triggered covariance (STC) techniques. We therefore computed the STC for all recorded thalamic units and compared this for each spiking condition. Although the dataset was more limited because the number of units with a significant STC filter was lower than those with a significant STA filter (n = 13 units with STC filter compared to n = 30 units with STA filter), the same trends regarding the reduction in the amplitude of the filter (STC_{SNR}) and the slope of the symmetric nonlinearity persisted (data not shown). Therefore, this suggests that the method of extracting the feature selectivity (STA compared to STC) was insufficient to explain the inability to estimate the feature selectivity in the hyperpolarized/burst spiking conditions.

The fourth assumption made throughout the analysis was that burst spikes are actually driven by sensory stimuli such that there is a recoverable burst spike feature selectivity. The alternative explanation would be that burst spikes are not feature selective and instead occur randomly due to intrinsic or other non-sensory processes. To assess this, we quantified the trial-to-trial repeatability for bursts in response to frozen white noise segments. As can be seen in Figure 3B, the qualitative assessment of temporally aligned bursts in response to the frozen white noise segment suggests that the bursts are driven by the sensory stimulus in a repeatable way. For units with a sufficient number of repeated trials, we computed the reliability of the burst spiking as the correlation between the peristimulus time histogram of even and odd trials in response to the frozen white noise segment. We found that all units showed greater reliability than what is expected based on just the temporal correlations in the
burst spiking (shuffle control, $p = 0.002, n = 10$ thalamic units). This suggests that the bursts are not randomly generated or due entirely to a non-stimulus related phenomenon.

From these controls, we propose that the difference in the spike triggered encoding properties could not be attributed to differences in the overall spike rates, the temporal properties of the spikes within the burst, or the mechanism of filter estimation. Instead, we propose that the burst spikes are driven by the sensory stimulus and have an underlying feature selectivity, but that this cannot be recovered using spike triggered techniques due to the reduced temporal precision of burst spiking relative to tonic spikes.

Recovering an STA relies on precise temporal spiking relative to the sensory stimulus. To simulate degradation of the spike timing precision, we added independent samples of normally distributed temporal jitter of varying amplitudes (standard deviation of the jitter distribution) to each tonic spike for an example unit and computed the STA (Figure 4A). Across units, we quantified the degradation of the STA as the jittered-STA$\text{SNR}$ normalized by the tSTA$\text{SNR}$ (0 ms jitter). The jittered-STA$\text{SNR}$ (black) is within the band expected for the bSTA$\text{SNR}$ with the addition of 4 milliseconds of jitter to the spike times (red shaded, Figure 4A, right). We propose that the effects of temporal jitter are particularly evident for whisker selectivity, presumably due to the short temporal duration of the filters (approximately 10-15 milliseconds in duration, Figure 1F).

Given the marked effects of jitter on the ability to recover the STA, we investigated the variability in the spike timing relative to the noise stimulus (Figure 4B). For this example unit, we have identified a segment in the noise stimulus that closely resembles the tonic STA for this unit and elicits a reliable spiking response (Figure 4B, top; stimulus – black, tSTA – grey dashed). The vertical dashed line indicates the spike time for the spike triggered average ($t_0$) with the grey bar indicating the duration of the STA. If there was no variability in the neural spiking, the raster plots would all be perfectly aligned to $t_0$ because the similarity between the stimulus and the STA would predict a spiking response at that time point. However, the timing of evoked neural responses is always variable to some extent and this can be visualized for
this example response segment as the temporal variability of the spike times surrounding this
stimulus feature in the noise stimulus (Figure 4B, as indicated by the grey stimulus bars that
extend from the first spike response to this particular sensory feature). For this example
snapshot, it is also apparent that the burst spikes in the hyperpolarized condition show greater
temporal variability than the tonic spikes in the baseline condition.

To quantify this jitter across all spikes, we developed a \( \tau_{\text{jitter}} \) metric that determines the
time lag of the peak correlation between the STA and the stimulus segment (\( s(t_i) \)) surrounding
each spike (Figure 4C). Intuitively, this is a correlative method to identify the time lag between
when we predict a spike is most likely to occur based on the STA and the stimulus (peak
correlation) and when the spike actually occurred. For this analysis, we treated the tSTA as
the true feature selectivity of the neuron across all spiking conditions because we could not
recover a reliable estimate of the bSTA.

We computed \( \tau_{\text{jitter}} \) for each spike and plotted \( \tau_{\text{jitter}} \) distributions for each spike condition
(tonic, burst, baseline, hyperpolarized). If a neuron was infinitely precise such that when the
stimulus matched the spike triggered average, the neuron fired a spike without delay, this
distribution would be represented by a delta function at \( \tau_{\text{jitter}} \) equals zero. As the variability of
the timing increases, the width of this distribution will also increase. For the tonic and baseline
condition spikes, we found a clear peak in \( \tau_{\text{jitter}} \) values at \( \tau_{\text{jitter}} \) equals zero (Figure 4C, grey,
black). For the burst and hyperpolarized condition spikes, we observe little-to-no peak in the
\( \tau_{\text{jitter}} \) metric at zero (Figure 4C, red, yellow). We computed the \( \tau_{\text{jitter}} \) distribution across all
thalamic units and found that the tonic and baseline spikes had higher peaks at \( \tau_{\text{jitter}} \) equals
zero than the burst and hyperpolarized spike conditions (Figure 4D). We quantified this
statistically by computing a precision metric (Figure 4C) that computes the proportion of spikes
within ±1 millisecond of \( \tau_{\text{jitter}} \) equals zero (Figure 4E). The tonic and baseline spike condition
were more precise than burst and hyperpolarized conditions.
Figure 4: Thalamic timing variability in the response to the sensory white noise. A. Effect of increased jitter on thalamic STA (schematic: left, example: center). Across units, the normalized amplitude of the jittered STA (jitteredSTA_{SNR}/tSTA_{SNR}) was plotted across jitter intensities (black, right). The normalized amplitude of the burst STA (bSTA_{SNR}/tSTA_{SNR}) is shown in red (mean ± s.d.). B. In this example, the same stimulus (black; scale bar: 0.1°) was presented with and without optogenetic hyperpolarization (data from example unit presented in Figure 3 with associated STA (grey dotted line; scale bar: 0.025°)). C. Example unit τ_{jitter} distributions for burst, tonic, baseline, and hyperpolarized conditions. D. Average τ_{jitter} distributions for burst/tonic spikes (n = 31 units) and baseline/hyperpolarized condition (n = 11 units). (mean +/- sem). E. Precision for burst/tonic spikes (n = 31 units, p = 1e-6) and baseline/hyperpolarized condition (n = 11 units). (mean +/- sem, p=1e-3).
These data suggest that tonic spikes showed greater temporal precision in response to the sensory white noise than burst spikes and that this could underlie the difference in the recoverability of the feature selectivity in the thalamus between firing modes. It is well established that the timing of sensory inputs is particularly important in the thalamocortical circuit such that changes in thalamic spike timing could have large impacts on the downstream representation of sensory information in the cortex. Next, we investigated how these changes in temporal precision in optogenetically modulated thalamic states impact cortical encoding properties.

Optogenetic modulation of thalamic firing modes directly impacts cortical representation of sensory information.

Cortical neurons that receive direct thalamic input are integrating information over a population of thalamocortical neurons that can be operating in different firing modes. This makes it difficult to determine the impact of a single burst from a single neuron on information representation in the pathway. Instead, we used the optogenetic manipulation of thalamic state as presented in Figure 3 to bias the activity of the thalamic population towards burst firing (hyperpolarized condition) and investigated the effects on the cortex. Here, we transfected the thalamus with a hyperpolarizing opsin and lowered an optic fiber into the thalamus while recording the cortical activity extracellularly (Figure 5A) in response to sensory white noise with and without optogenetic manipulation of the thalamus (hyperpolarized VPm, baseline VPm).
For an example unit, we have plotted the cortical STA in the baseline and hyperpolarized VPm conditions (Figure 5B). Here, the amplitude of the cortical STA was smaller when the thalamus is hyperpolarized compared to when it is not (Figure 5B). This cortical unit also shows a reduced tuning to the STA when the thalamus was hyperpolarized (Figure 5C). Across the population of recorded cortical neurons, we saw the same effect seen in this example neuron of a reduced STA$_{SNR}$ when the VPm was hyperpolarized compared to

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**Figure 5: Optogenetic modulation of thalamic firing modes directly impacts cortical representation of sensory information.**

A. Experimental paradigm. Scale bar: 1 mm. B. Example STA for a cortical unit recorded during optogenetic manipulation of the thalamus. C. Example cortical nonlinearity as a function of thalamic state. D. STA$_{SNR}$ across cortical units (n = 32 units. p = 1e-6). E. Spiking nonlinearity across cortical units. F. Cortical firing rate (p = 0.31). G. Example cortical unit τ$_{jitter}$ distributions for baseline and hyperpolarized thalamic conditions. H. Average τ$_{jitter}$ distributions for baseline/hyperpolarized thalamic condition (mean +/- sem). I. Precision for cortical spikes during baseline/hyperpolarized thalamic condition (mean +/- sem, p=8e-6).
when it was not (Figure 5D) and a reduction in the tuning across all cortical units as quantified by the spiking nonlinearity (Figure 5E). These findings mirror what was seen for thalamic neurons when comparing the baseline and the optogenetically manipulated conditions demonstrating that the changes in thalamic encoding properties are propagated to cortex.

Interestingly, there was no significant difference in the noise-evoked firing rate in the cortex as a function of the VPm condition (Figure 5F). This suggests that it was not overall spike counts influencing the cortical feature selectivity. Instead, we propose the temporal jitter in the thalamic spiking patterns propagated to cortex. We investigated the temporal precision of the cortical spiking in response to the sensory white noise using the same methodology employed for the thalamus. As we saw for the thalamus, the cortical spikes from this example unit also showed greater temporal precision in the baseline VPm condition compared to the hyperpolarized VPm condition (Figure 5G) as evidenced by the peak in the $\tau_{\text{jitter}}$ distribution around $\tau_{\text{jitter}}$ equals zero. This effect was consistent across the population of recorded cortical units (Figure 5H) and showed significant differences in the precision of the cortical firing (Figure 5I). This suggests that the temporal jitter present in the thalamus is transmitted to the cortex where it also impacts the representation of sensory information.
Discussion

The highly interconnected thalamocortical pathway dynamically gates information flow between the periphery and higher cortical centers dependent on many factors including behavioral state (Niell and Stryker, 2010), task demands (Atiani et al., 2009), and sensory adaptation state (Whitmire et al., 2016). Although there have been extensive investigations into the cortical state-dependent processing of the thalamocortical circuit, we know surprisingly little about how this information is processed in a thalamic state-dependent manner. Here, using a combination of optogenetic stimulation and electrophysiological recording techniques, we have performed a series of experiments modulating the state of the thalamus (through constant optogenetic hyperpolarization) and quantified the effects on encoding in the thalamocortical circuit. Using this technique, we have coarse control of the firing mode in thalamus without altering the processing occurring from the whisker to thalamus, enabling us to decouple the changes in thalamic firing mode on thalamocortical processing from changes occurring in subthalamic processing. We found that, unlike the visual pathway, the feature selectivity of burst spikes in the vibrissa pathway could not be recovered using spike triggered techniques due to increased burst spike timing variability relative to tonic spike timing. Recordings from barrel cortex during optogenetic manipulation of thalamic state demonstrated a shift in the temporal precision of the cortical spiking that also led to a degradation of the recovered feature selectivity. This suggests that bursts in the whisker pathway are less precise than tonic spikes during ongoing sensory stimulation and that this loss of temporal precision is propagated to cortex.

These results could be interpreted as consistent with the view that bursts are not representing detailed stimulus information. However, there is evidence that bursts may convey more information than the presence or absence of a burst through inter-burst spike timing and the number of spikes per burst (Mease et al., 2017), suggesting a role of temporally precise burst firing in information representation. Furthermore, thalamic bursting can be temporally precise within and across neurons in response to high intensity whisker stimuli (Whitmire et
Instead, we propose that the temporal precision of the thalamic firing is a function of both the state of the thalamus and the intensity of the sensory stimulus. It has previously been shown that the temporal precision of thalamic encoding increases with the intensity of the sensory stimulus (Desbordes et al., 2008; Whitmire et al., 2016) while here we have shown that the temporal precision of the thalamic firing decreases with sustained hyperpolarization. These two competing factors would enable the burst firing mode to encode high amplitude stimuli in a temporally precise fashion while low amplitude stimuli, such as the sensory white noise presented here, would not be able to overcome the variability present in the burst state.

There are multiple mechanisms that could underlie the reduced temporal precision in the burst firing mode including variability introduced by the slow dynamics of the calcium depolarization, increased variability in the time to reach threshold due to the prolonged hyperpolarization of the baseline polarization, as well as potential changes in the integration properties of the thalamic neurons. Furthermore, these mechanisms could occur independently such that the variability across neurons is uncorrelated or these mechanisms could be coordinated in some way to enable correlated variability across the thalamic population. Both coordinated and uncoordinated jitter would have a detrimental effect on the ability to recover the STA because either the spike timing would no longer be locked to the stimulus itself or the input to the cortex would be temporally imprecise. However, coordinated jitter would maintain the information about the stimulus while uncoordinated jitter would degrade the recoverability of the stimulus features. Future work should investigate the jitter in the burst spiking across units to determine whether or not the variability in the spike timing is coordinated across thalamic units in this context.

Given the importance of thalamic spike timing precision within and across neurons in transmitting information downstream to cortex (Bruno and Sakmann, 2006; Wang et al., 2010), alterations to thalamic state can shape multiple properties of the spiking inputs to cortex. Manipulation of the thalamic state can lead to changes in the stimulus evoked cortical dynamics (Whitmire et al., 2017) and spatiotemporal cortical activation (Borden et al., 2017).
We have primarily considered thalamic state-dependent encoding as a feedforward representation from thalamus to cortex, but the highly interconnected thalamocortical circuitry is a dynamic interaction that shapes coding properties in both feedforward and feedback manner. Changes in thalamic activity impact cortical activity which then provides feedback to thalamus to further alter activity (Crandall et al., 2015; Mease et al., 2014; Poulet et al., 2012; Reinhold et al., 2015; Wimmer et al., 2015). It is possible for thalamus to influence cortical state and for the cortex to influence thalamic state, but how this plays out during natural behaviors is not yet known and must be decoupled using more sophisticated techniques such as closed-loop control of neural activity (Bolus et al., 2018; Newman et al., 2015). The ability to shift the temporal precision of the thalamic spike timing through changes to the thalamic state, or the baseline membrane potential, provides a biophysical mechanism for the thalamus to gate information flow to cortex. Furthermore, this mechanism could be under both feedforward and feedback control. This sets the stage for a dynamic interaction between thalamic and cortical states to drive highly interactive patterns of neural activity.
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