



# Single neurons in the thalamus and subthalamic nucleus process cardiac and respiratory signals in humans

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Visceral signals are constantly processed by our central nervous system, enable homeostatic regulation, and influence perception, emotion, and cognition. While visceral processes at the cortical level have been extensively studied using non-invasive imaging techniques, very few studies have investigated how this information is processed at the single neuron level, both in humans and animals. Subcortical regions, relaying signals from peripheral interoceptors to cortical structures, are particularly understudied and how visceral information is processed in thalamic and subthalamic structures remains largely unknown. Here, we took advantage of intraoperative microelectrode recordings in patients undergoing surgery for deep brain stimulation (DBS) to investigate the activity of single neurons related to cardiac and respiratory functions in three subcortical regions: ventral intermedius nucleus (Vim) and ventral caudalis nucleus (Vc) of the thalamus, and subthalamic nucleus (STN). We report that the activity of a large portion of the recorded neurons (about 70%) was modulated by either the heartbeat, the cardiac inter-beat interval, or the respiration. These cardiac and respiratory response patterns varied largely across neurons both in terms of timing and their kind of modulation. A substantial proportion of these visceral neurons (30%) was responsive to more than one of the tested signals, underlining specialization and integration of cardiac and respiratory signals in STN and thalamic neurons. By extensively describing single unit activity related to cardiorespiratory function in thalamic and subthalamic neurons, our results highlight the major role of these subcortical regions in the processing of visceral signals.

visceral processing | interoception | single neurons | thalamus | STN

Our brain continuously receives signals originating from visceral organs providing a moment-by-moment sense of the physiological condition of the body. Such monitoring of bodily organs ensures the stability of the organism through regulation of homeostatic reflexes and adaptive behavior (1, 2). There has been a recent upsurge of interest in the understanding of how information from internal organs is integrated in the human brain, because, beyond its function in homeostasis, visceral processing and the perception of the physiological condition of the body (i.e., interoception) are increasingly recognized to impact cognitive, social, and affective processes (3–9) and to be a prominent feature of several psychiatric disorders (10–13).

Based on animal studies relying on electrical stimulation, chemical activation, pharmacological interventions, and various tracing methods, it is commonly acknowledged that visceral inputs reach the central nervous system through the vagal nerve or the spinothalamic tract projecting to the nucleus tractus solitarius, the parabrachial nucleus, and periaqueductal gray matter (14–17). Importantly, chest movement or pulsatility of blood vessels are also detected by proprioceptive or tactile peripheral receptors and are conveyed in the dorsal column-medial lemniscus pathway. Visceral information is subsequently transmitted to the hypothalamus, the amygdala, and the thalamus, from which it reaches primary sites of the viscerosensory cortex such as the insula and the anterior cingulate cortex (5). Specifically, the ventral posterior complex of the thalamus has been put forward as the main relay of visceral information traveling to the cortex in most animal models (18–21). Importantly, electrophysiological studies in animals have shown single-unit responses to thermal (22), gustatory (22, 23), gastrointestinal (24, 25), respiratory (26, 27), and cardiac (28) activity located in ventral posterior medial (VPM) (22, 23), ventral posterior lateral (VPL) (24, 25), and/or posterior part of the ventro-lateral nucleus (VLp) of the thalamus. Similarly, STN neurons in the cat have been reported to tonically fire during specific respiratory phases (27) and, when stimulated, to induce strong cardiorespiratory responses (29, 30).

In humans, visceral processing has been characterized in different cortical and limbic regions, while the interoceptive ascending pathways linking the periphery to higher-order regions of the brain have been relatively neglected (14). One extensively studied signal is

## Significance

Internal bodily signals, like cardio-respiratory ones, are constantly transmitted from the body to the brain ensuring self-regulation of the organism. Subcortical brain regions are particularly important for this body-brain communication, yet their processing of internal bodily signals in humans is largely unknown. Studying the activity of single neurons in three subcortical regions in humans (two thalamic nuclei and the subthalamic nucleus), we found that a large portion of neurons was modulated by either the heartbeat, the respiration, or the cardiac cycle duration, while the prevalence of those signals was much reduced in cortical control regions. Our study shows how vital cardio-respiratory signals are widely processed in these subcortical regions, expanding our understanding of their role in body-brain communication.

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The authors declare no competing interest.

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the heartbeat evoked potential (HEP)—obtained by averaging surface EEG signals time-locked to heartbeats—which has been associated with widely distributed cortical sources over frontal, central, and parietal regions (see ref. 31 for a recent review). HEP has been linked with cardiac function (32, 33), but also with various cognitive and affective processes (7, 34–41). Although less investigated than cardiac signals, respiratory signals have also been shown to modulate brain oscillations at rest in widespread brain networks (42, 43). Similarly to cardiac signals, respiration has also been shown to influence a range of cognitive and motor processes (44–51).

A powerful approach to characterize how visceral information is encoded in the human brain is to use invasive recordings that directly measure local neural activity. Recently, such studies have profited from the high spatial and temporal resolution of intracranial electroencephalography (iEEG) and local field potentials to reveal specific heart-related processes in the primary somatosensory cortex (52), but also the insula, operculum, amygdala, and fronto-temporal cortex (37). Slow (~0.2 Hz) respiration-entrained oscillatory activity has also been reported in the human limbic system (51, 53). However, at the single neuron level, to date only two studies have been able to describe a relation between cardiac signals (i.e., duration of the cardiac cycle) and neural firing rates, reporting evidence for cardiac coding in the human anterior cingulate cortex and medial temporal lobe (54, 55). For respiration, while prior intracranial studies in humans have shown local neural activity linked to breathing (56, 57), to date, no study has, to the best of our knowledge, investigated the link between respiration and single neuron activity in humans.

However, major brainstem and midbrain centers are involved in the processing and regulation of cardiac and respiratory signals for homeostasis and control of adaptive behavior (14). While cortical processing of visceral signals has thus been elucidated by the listed studies, a fundamental gap in our understanding of visceral processing in the human brain remains. How are visceral signals processed at subcortical centers that project to the above-mentioned cortical structures? The neural properties of subcortical relays, such as the thalamic and subthalamic nuclei, are particularly understudied, likely due to the methodological challenges in accessing and analyzing these signals in humans. Rare human neuroimaging experiments (i.e., fMRI) and neurostimulation protocols have suggested an involvement of the mediodorsal thalamus, the periaqueductal gray matter, the STN, the hypothalamus in autonomic system regulation (58, 59), and of different subregions of the thalamus in respiratory control (60, 61). Moreover, electrical stimulation of the ventral portion of the human thalamus (59) and STN (62) modulates baroreceptors sensitivity, increases heart rate and blood pressure while changes in single neuron activity related to cardio-vascular modulation have been reported in the ventro-postero-lateral nucleus of the thalamus (63) and STN (64).

An invaluable opportunity to investigate the subcortical functional aspects of visceral responses in humans is provided by intraoperative recordings of single neurons during the implantation procedure of electrodes for deep brain stimulation (DBS). During DBS surgeries, intraoperative microelectrode recordings (MER) provide the opportunity to record from thalamic and subthalamic nuclei in awake humans (65). However, an important challenge in DBS recordings—especially when investigating signals time-locked to the heart—is the presence of pulsatility artifacts (PA) resulting from the pulsating motion of the brain and nearby pulsating blood vessels. Such movements lead to changes in the recorded spike waveform across the cardiac cycle that can affect spike detection and bias neuron classification. To our knowledge, only one study, so far, investigated single cell activity related to

cardio-vascular processing in the human thalamus (63). While representing an important and pioneering report, Oppenheimer et al. limited their investigation to the relationship of thalamic neuronal activity with respect to the pulse signal (measured via a pulse-oximeter on the toe; as no ECG was available for the analyzed dataset). Furthermore, single units and their stability were only assessed by visual inspection, which may lead to multi-unit clusters and result in an underestimation of the number of modulated units.

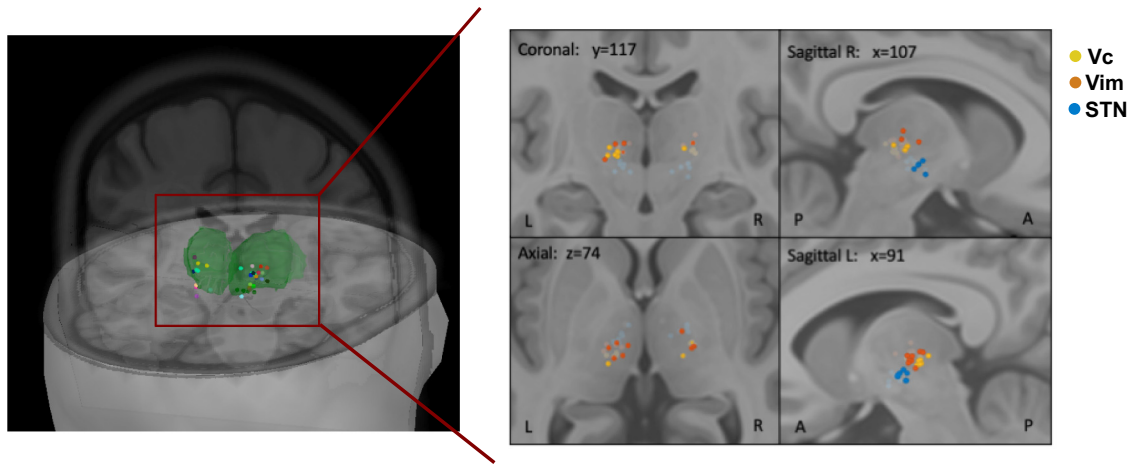
Here, we combine MER with simultaneous physiological signal acquisition (heart rate, HRV, breathing rate, breathing variability) to investigate whether single neurons in the thalamus and STN process basic signals of heartbeat and respiration. To overcome the challenges of intraoperative MER recordings faced by Oppenheimer et al., we applied latest single-unit analysis methodology, using unsupervised parametric spike sorting algorithms (66, 67) and a recently developed method for the analysis of cardiac-motion features in the recorded spikes (68). Following preprocessing of the extracellular potentials, we identified three different visceral signals in thalamic and subthalamic neurons, linked to 1) the heartbeat, 2) the duration of the cardiac cycle, and 3) the respiratory cycle. The activity of a large portion of the recorded neurons (about 70%) was modulated by at least one of these visceral signals, with a substantial proportion of those neurons (30%) encoding multiple signals. Our results provide functional characterization of the activity of thalamic and subthalamic single neuron populations in humans encoding cardiorespiratory activity, highlighting the role of these subcortical regions in interoceptive processing.

## Results

**Microelectrode Recordings and Identification of Neurons.** We recorded data during DBS implantation surgery in a total of 23 patients. Implantation site of all recordings was based solely on medical indication, that is Vim target for essential tremor and STN target for Parkinson disease. We acquired data from three different subcortical regions: the subthalamic nucleus (STN), the ventral intermedialis (Vim) thalamic nucleus, and ventral caudal (Vc) thalamic nucleus. Fig. 1 shows the MNI normalized coordinates of the recording location across the 23 surgeries (63 recording sessions in total).

After the initial spike detection and sorting, we identified 127 putative single units, of which 27 were in STN, 45 in Vc, and 55 in Vim (mean number of units per recording session = 2.0, SD: 1.1). We then applied a modified version of a recently proposed method (68) to identify and remove neurons that are contaminated by cardiac-related movement artifacts in their EAPs (*SI Appendix, Supporting Methods and Fig. S1*). Following this procedure, 73 neurons were considered for further analysis (16 in STN, 24 in Vc, and 33 in Vim). The number of neurons detected in each area per patient is reported in *SI Appendix, Table S1*.

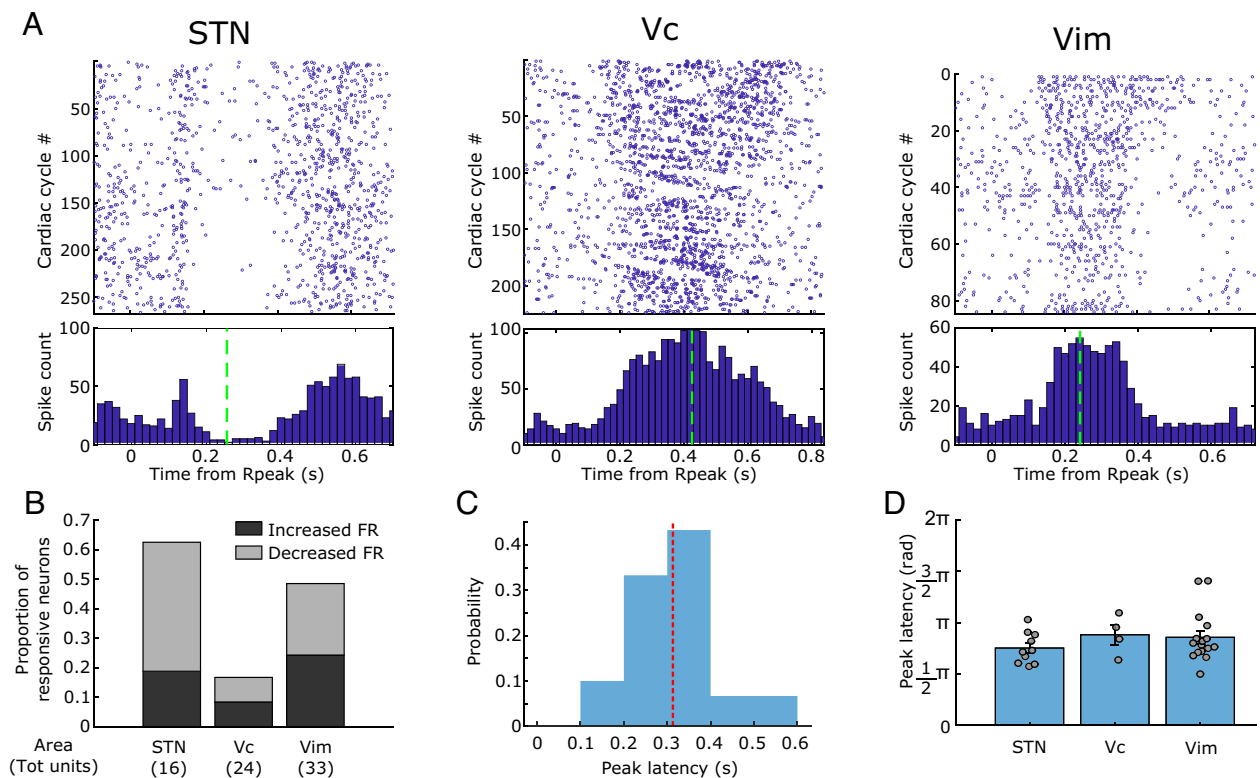
**Neural Responses to Heartbeats.** Modulation of the neural activity in response to the heartbeat was assessed by analyzing the changes in neural firing rates along the cardiac cycle for each of the 73 neurons (following exclusion of neurons with artifacts) by computing a non-uniformity test and the coherence between spikes and cardiac phase (see *Methods* for further details). This analysis led to the identification of 30 neurons (41%) with a significant modulation of their firing activity along the cardiac cycle. In a post hoc analysis, we tested whether the proportion of responding neurons differed between recordings sites and found a significantly larger proportion of neurons responded to the heartbeat in both STN (10 units, 63%) and Vim (16 units, 48%) compared to Vc



**Fig. 1.** Recording sites. Location of recording electrodes across the 23 surgeries color-coded by patient (*Left*) and color-coded by identified region for Vc (yellow), Vim (orange), and STN (blue) (*Right*). Green volume represents the thalamus. Labeling of recording sites was obtained intraoperatively based on each patient's brain scan, coordinates were then converted to MNI space for visualization and are projected here on the MNI/ICBM152 human brain template (69). Note that for some sessions, location dots overlap as different recordings were performed in the same location. Abbreviations used: ventral caudal thalamic nucleus (Vc), ventral intermedialis thalamic nucleus (Vim), subthalamic nucleus (STN). *Left* and *Right* panels generated using Brainstorm (70).

(4 units, 17%) [z-test on proportions, STN vs. Vc:  $z(38) = 3.1$ ,  $P = 0.002$ ; Vim vs. Vc:  $z(55) = 2.7$ ,  $P = 0.006$ ]. Some exemplary responsive units across STN, Vim, and Vc are shown in Fig. 2*A*. Neurons presented different response patterns along the cardiac cycle. For example, the STN neuron in Fig. 2*A*, *Left* panel shows a clear decrease in firing activity following the R-peak, with the average number of spikes approaching zero during the central part

of the cardiac cycle. Conversely, the illustrative Vc and Vim neurons shown in the central and *Right* panel of Fig. 2*A* show a different pattern, that is, a marked increase in spiking activity following the R-peak. Across the 30 responsive units, 13 increased their firing rate following the R-peak (43%) and 17 decreased it (57%). No significant difference in the proportion of increasing vs. decreasing responses was observed [z-test,  $z(28) = 1.02$ ,  $P = 0.3$ ]. In addition,



**Fig. 2.** Neural responses to heartbeats. (*A*) Examples of neurons responding to the heartbeat with firing rate modulations in the three different target regions (STN, Vc, and Vim) recorded from three different patients. The *Top* panel shows the raster plot for the neural spikes aligned to the R-peak across the different cardiac cycles. The bottom plot shows the cumulative histogram of spikes across cardiac cycles binned in 10 ms bins. Green dotted lines represent the peak latency (time with maximal modulation of the response). The maximum value for the x-axis is equal to the median IBI for the recording session. Sample recordings and EAP waveform modulation test for these exemplary units are shown in *SI Appendix, Figs. S4.1–S4.3*. (*B*) Proportion of neurons showing a significant increase (dark gray) or decrease (light gray) of firing rates along the cardiac cycle in the three regions. The total number of neurons tested in each region is shown in parentheses. (*C*) Distribution of peak latencies for the 30 responsive neurons identified. Average latency was 313 ms (SD = 100 ms). Of note, 77% of responses had a peak latency between 200 and 400 ms. (*D*) Comparison of peak latencies normalized by IBI (in rad) across areas. Bars represent the mean ( $\pm$ SEM). Circles represent single peak latencies measured. No significant difference in peak latency was observed across regions [ANOVA test,  $F(2, 27) = 0.89$ ,  $P = 0.42$ ].

when we repeated our analysis by time-locking the spikes to the end of the cardiac cycle (i.e., looking backward from the R-peak), we found that all the 30 neurons still exhibited a significant response. Furthermore, we verified that the identified heartbeat responses could not be explained by spikes near the detection threshold (*SI Appendix, Supplementary Results and Fig. S5*).

Single Fig. 2*B* shows the proportion of significant responsive neurons identified in the three regions color-coded per type of response (increased or decreased firing). Although, the STN was found to present the largest proportion of decreasing responses (70%), compared to the two thalamic nuclei (50% both in Vc and Vim), the differences in proportion were not significant [ $z$ -test on STN vs. thalamus (Vc + Vim),  $z(28) = 1.04$ ,  $P = 0.29$ ].

For each response, the peak latency (represented by the green dotted lines in Fig. 2*A*) from the ECG's R-peak was determined from the instantaneous firing rates as the time where the modulation of activity (increase or decrease) was maximal. Distribution of all peak latencies for the 30 responsive neurons is shown in Fig. 2*C*. The average peak latency across responses was 313 ms (SD = 100 ms), with 77% of peak latencies (23 out of 30) falling in the interval between 200 and 400 ms. In terms of angular latency (in rad), the mean latency was 2.5 rad (SD = 0.6 rad). Because of the variability in IBIs across patients, peak latencies across regions were compared in terms of their angular values (in rad) that correct for differences in IBI observed across recordings. No significant difference was observed across regions according to an ANOVA test [ $F(2, 27) = 0.89$ ,  $P = 0.42$ ] (Fig. 2*D*). Furthermore, no significant difference in peak latencies was observed between increasing and decreasing responses [unpaired  $t$  tests:  $t(28) = 0.53$ ,  $P = 0.6$ ].

These data show that signals about the cardiac cycle are widely encoded in thalamic and subthalamic neurons (41% of units). These signals manifest as a rhythmic modulation (increase/decrease) of spiking activity during specific portions of the cycle, with the peak modulation on average around 300 ms (post R-Peak), roughly corresponding to the end of the first half of the cardiac cycle.

### Relationship between Firing Rates and Inter-Beat Interval (IBI).

The heart rate variability (HRV)—the fluctuation in the time intervals between consecutive heartbeats—is an important physiological marker, in cardiac physiology, of heart–brain interactions, and a useful tool to assess sympathetic and parasympathetic influences on disease states (71). We therefore asked whether human thalamic and sub-thalamic neurons process IBI-related signals. For each neuron, we tested the relationship between spiking activity and IBI by means of a Pearson correlation between the average firing rates during each cardiac cycle and the corresponding cycle duration (IBI). Fig. 3*A1* shows examples of neurons whose firing rate was significantly correlated with the IBI. The corresponding raster plots for the units are shown in Fig. 3*A2*. Overall, we found significant correlation for 29% of the recorded units (21 out of 73,  $t$ -statistic corrected for multiple comparisons), with nine units exhibiting a positive correlation (higher firing rates for longer IBIs) and 12 units exhibiting a negative correlation (lower firing rates for longer IBIs) (Fig. 3*B*). The probability of positively/negatively correlated units across the three recorded regions is shown in Fig. 3*C*. These data show that spontaneous spiking activity in the recorded regions is linked to IBI for almost one third of recorded neurons, and that longer IBIs can be linked to either positive or negative changes in firing rates.

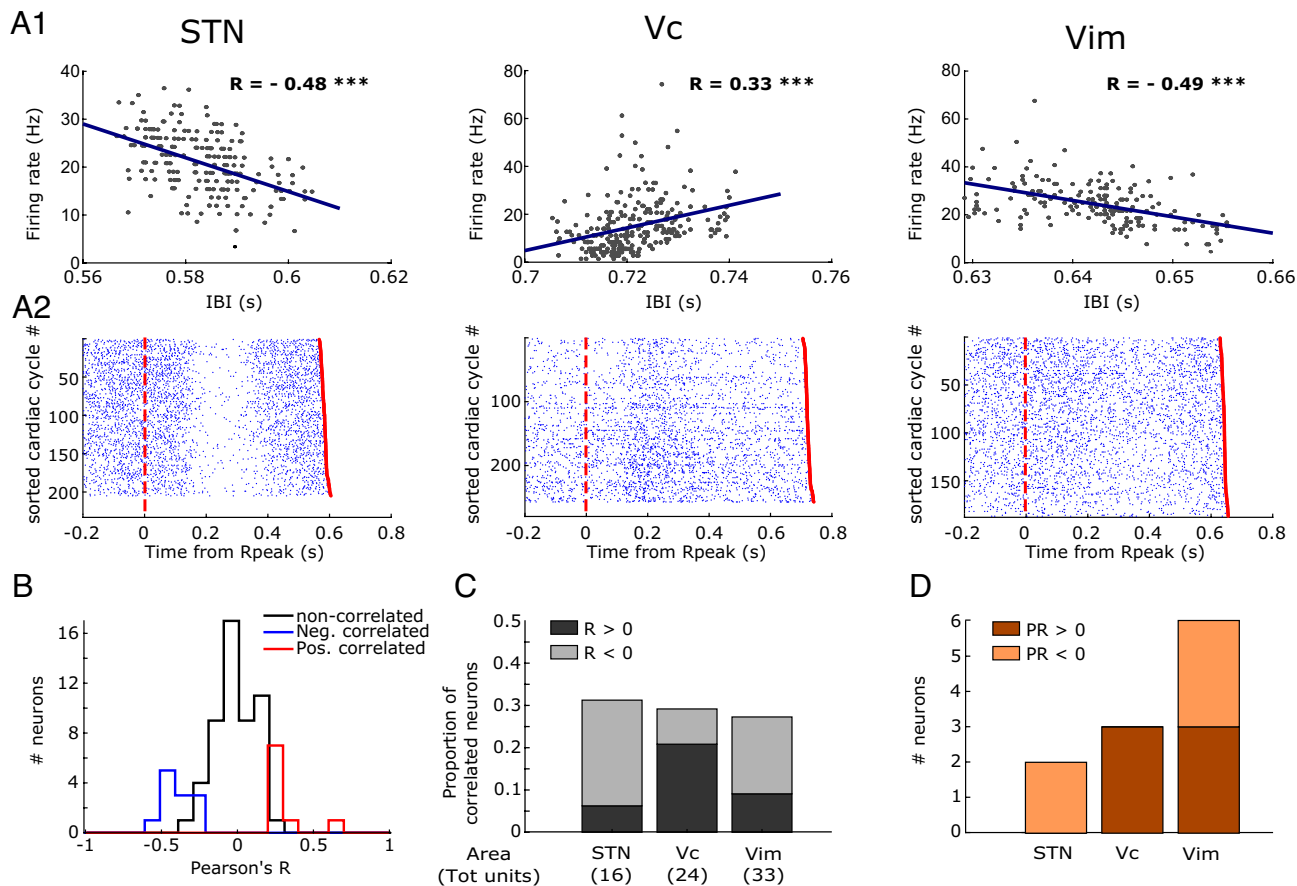
Given that the heart–brain communication is bidirectional and can be mediated by several pathways, we next tried to characterize the directionality of the observed relationship. We used phase-response (PR) analysis at the cycle-by-cycle level, as recently proposed by Kim et al. (55), to identify neurons for which the spiking activity elicited changes in the duration of the cardiac cycle. PR

measures how much the IBI is shortened or lengthened in response to a neuron's spiking activity. Among the units exhibiting a significant correlation between firing rate and IBI (21), we found that 11 also showed a significant PR of IBI to the spiking activity. Specifically, six units had positive PR (lengthening of the cardiac cycle following spikes) and five units had a negative PR (shortening of the cardiac cycle following spikes). As expected, the sign of PR was always matching the sign of Pearson's  $R$  measured from the correlation.

Altogether, these results demonstrate that information about the duration of the cardiac cycle is encoded in the activity of about a third of the present thalamic and subthalamic neurons. Moreover, the relationship between firing rate and IBI was characterized either by a positive or negative correlation in STN, Vim, and Vc. In a few neurons, this relationship extended to more than one cardiac cycle (*SI Appendix, Supplementary Results and Fig. S6*).

**Neural Responses to Respiratory Phase.** Apart from the widely studied heart-related processes, interoception also embraces other systems, including input from the respiratory system. Respiratory-associated thalamic activity has been reported in cats (26), and the existence of respiratory signals in the human thalamus and STN would provide evidence for their broader implication in interoceptive processing. Furthermore, since respiratory signals have a much slower frequency than cardiac ones, their neural correlates are less prone to be affected by pulse artifacts. Therefore, we decided to capitalize on respiratory signals by analyzing the recorded neural activity in relationship to the respiratory cycle. Respiratory signals were extracted from the ECG traces using an ECG-derived respiration (EDR) algorithm (*SI Appendix, Supporting Methods and Fig. S7A*). We could successfully extract the EDR for all the recording session containing the identified neurons. Given that systematic coupling between cardiac and respiratory signals exists (e.g. respiratory sinus arrhythmia), neurons showing a significant EAP modulation along the cardiac cycle were also excluded from all respiration-related analyses. Furthermore, we verified that none of the considered neurons had a significant EAP waveform modulation along the respiratory cycle either (*SI Appendix, Supporting Methods*). Therefore, all 73 neurons were considered for the following analysis. For each neuron, we assessed whether the spiking probability was non-uniform along the respiratory cycle, that is, if the neuron fired preferentially during a specific respiratory phase (preferred phase). The existence of a preferred phase was assessed via surrogate population testing that intrinsically corrects for eventual differences in the distribution of phases along the recording. Some examples of neurons that fired preferentially for a specific phase of the respiratory cycle are shown in Fig. 4*A1* and *A2*. Overall, we identified 22 neurons (31%) whose activity was modulated by the respiratory phase (7 in STN, 5 in Vc, and 10 in Vim). The proportion of respiration-modulated neurons in each region is shown in Fig. 4*B*. The distribution of preferred phases for those neurons is shown in Fig. 4*C*. While we found that a slightly larger number of neurons (13 out of 22) fired preferentially during the inspiration ( $\pi < \phi < 2\pi$ ), no significant phase preference overall was identified [Rayleigh test for non-uniformity,  $z(21) = 0.37$ ,  $P = 0.69$ ]. Additionally, the relationship between firing rates and duration of the respiratory cycle was tested similarly to what was done for the IBI (previous section). To this end we tested the correlation between average firing rate and the time interval from the beginning of one expiration to the beginning of the following one. None of the recorded units showed a significant correlation between the two measures.

These results show that information about the respiratory cycle is encoded in about one third of the thalamic and subthalamic



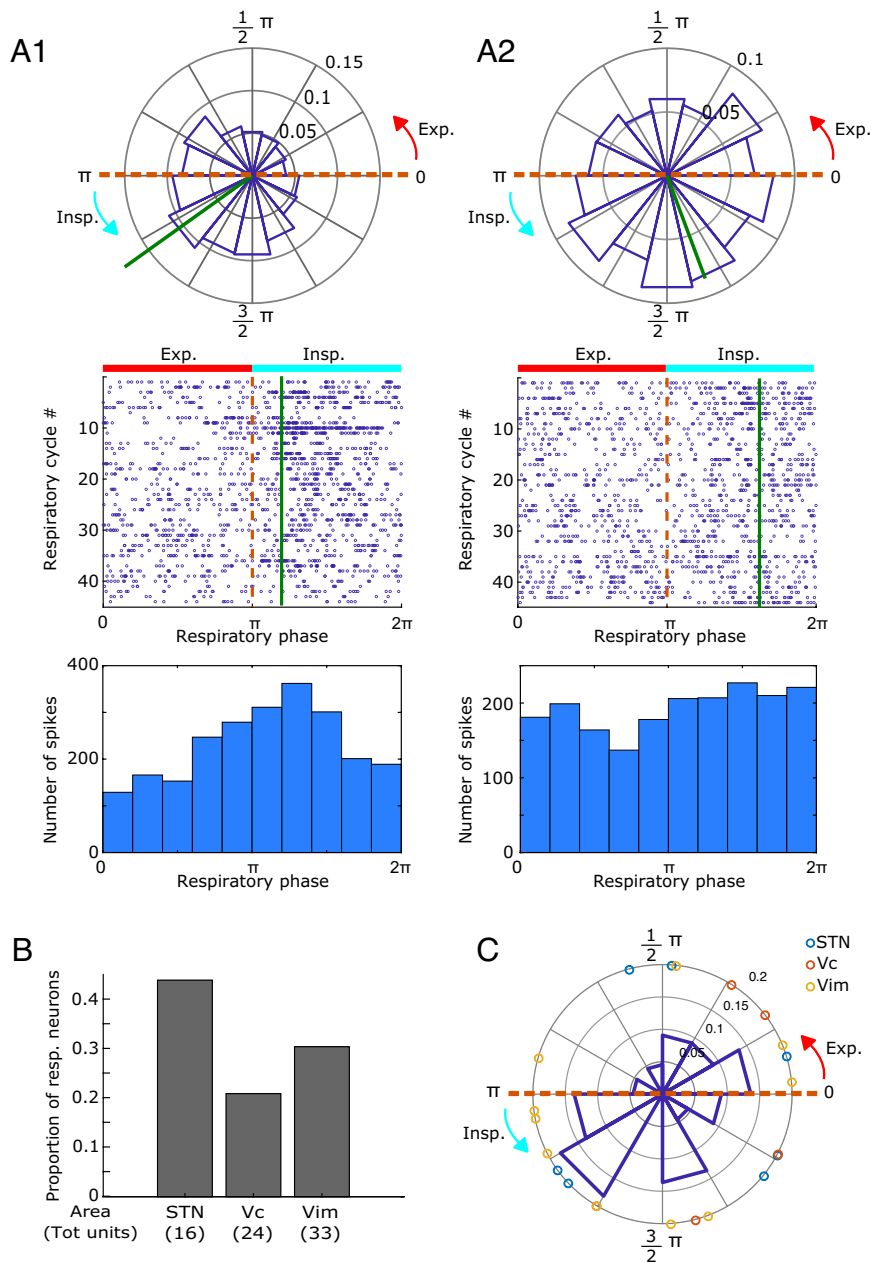
**Fig. 3.** Neural activity covaries with Inter-Beat Interval. (A1) Correlation between the duration of the cardiac cycle (IBI) and the firing rate across cycles in three exemplary neurons recorded from the three different target regions (STN, Vc, and Vim). Pearson's R is reported for each plot. Asterisks mark correlation significance (\*\*\* indicate  $P < 0.001$ ). (A2) Raster plots from the three exemplary neurons of the neural spikes aligned to the R-peak across the different cardiac cycles where cycles were sorted according to their IBI. The red dotted line represents the beginning of the cardiac cycle (R-peak) while the red dot at the end of each trial represents the following R peak. (B) Distributions of Pearson's R values for the correlation between IBI and firing rates separated in un-correlated (black), and significantly correlated (blue for  $R < 0$ , red for  $R > 0$ ). (C) Proportion of recorded neurons exhibiting a significant positive (dark gray) or negative (light gray) correlation between IBI and firing rate in the three regions. The total number of neurons tested in each region is shown in parentheses. (D) Number of recorded neurons in the three regions for which a significant phase-response (PR) of IBI to neural activity was detected. Bars are colored according to the sign of the detected PR (dark orange for positive, light orange for negative).

neurons. While cortical neural excitability is known to vary over the respiratory cycle (72), our findings cannot be simply explained by such a generalized increase of neural activity during a specific respiratory phase, because different neurons that we studied here exhibit different preferred phases for firing. Instead, the variability of preferred phases is more likely connected to the activity of respiratory neurons in the brainstem whose activation is observed in different phases of either inspiration or expiration (73).

**Overlap of Cardiorespiratory Modalities.** Given that we identified three different interoceptive signals in our dataset (heartbeat, cardiac IBI, and respiration), we next asked whether the same responsive neurons encode these different types of signals or whether there are rather separate populations of neurons encoding each type of signal. When looking for neurons responding to each of the different signal types (Fig. 5A) we found that most of them (35 out of 53 responsive units, 66%) responded to only a single type of interoceptive signal. From the remaining 18 neurons, 16 (30% of the total responsive units) responded to two types of stimuli, whereas only very few (2, 4%) responded to heartbeat, IBI, and respiration cycle. For example, the 21 neurons responding to the IBI were in the large majority distinct from those responding to the heartbeat (with only seven exhibiting both heartbeat and IBI modulation), and distinct from those responding to respiration (with only six exhibiting both heartbeat and

respiration modulation). Furthermore, among the seven neurons responding to both the heartbeat and the IBI, the directionalities of these modulations (increase/decrease of firing rates and positive/negative correlation) only matched in four units, indicating that the two modulations are not univocally affecting each other and therefore are not necessarily related. The proportion of neurons that responded to each modality or to multiple modalities (multi-responsive visceral neurons) across the three recorded regions is shown in Fig. 5B. There was no clear bias in the distribution of multi-responsive visceral neurons across regions. Furthermore, no relationship was found between respiratory sinus arrhythmia, reflecting coupling between respiratory and cardiac signals, and multimodality of the responses (SI Appendix, Supplementary Results and Fig. S8).

Additionally, we performed the same analyses as carried out for subthalamic and thalamic responses in the present manuscript on a previously published dataset (55), consisting of depth microelectrode recordings from several cortical regions in epileptic patients undergoing seizure monitoring (see SI Appendix, Supplementary Results, Comparison Regions for further details). Critically, we found that none of the isolated units responded to the heartbeat, only two units (3.8%) responded to the respiratory phase, and nine units (about 10%) exhibited a correlation between firing rates and IBI, underscoring the less prominent role of cortical regions in processing cardiorespiratory signals.



**Fig. 4.** Neural activity is modulated by respiration. (A1) Exemplary neuron in the STN whose spiking activity was significantly non-uniform along the respiratory cycle (surrogate test on Rayleigh statistic for non-uniformity,  $P = 10^{-4}$ ). *Top:* Circular histogram of spike density across the respiratory phase. The green line represents the mean vector of the distribution, with its angular direction being the preferred spiking phase for the neuron ( $\phi = 3.8$  rad). *Middle:* Raster plot for the neural spikes as a function of the respiratory phase across the different respiratory cycles. *Bottom:* Cumulative histogram of spikes across respiratory cycles binned in  $\pi/5$  rad bins. (A2) Exemplary neuron in the Vim whose spiking activity was significantly non-uniform along the respiratory cycle (surrogate test on Rayleigh statistic for non-uniformity,  $P = 2 \times 10^{-3}$ ). Conventions are as in panel (A1). Preferred spiking phase  $\phi = 5.1$  rad. (B) Proportion of neurons whose activity was significantly modulated by the respiratory phase in the three recorded regions. The total number of neurons tested in each region is shown in parentheses. (C) Distribution of preferred spiking phases for the 22 significantly modulated neurons. Circles indicate individual values (color-coded by region). No significant phase preference overall was identified [Rayleigh test for non-uniformity,  $z(21) = 0.37$ ,  $P = 0.69$ ].

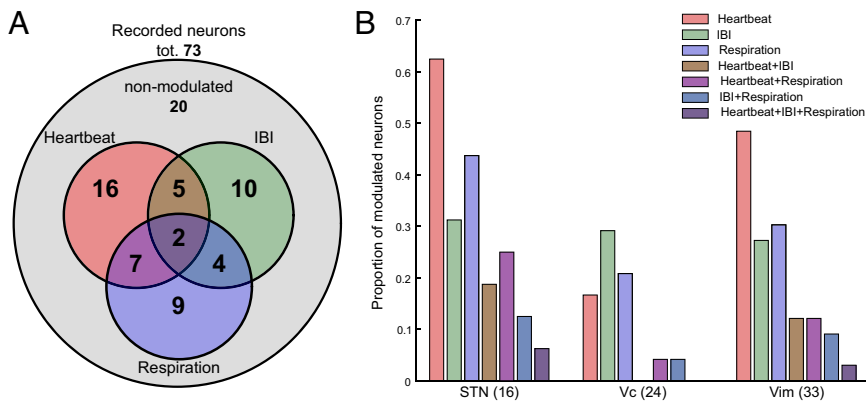
Overall, these data show that visceral signals are processed in the thalamus and STN by partially overlapping neural populations. The finding that diverse signals can be encoded by the same neuron underlies the interaction between different visceral modalities. Finally, these results are further reassurance that these cardiac and respiratory signals in thalamic and subthalamic neurons are not the effect of recording artifacts since the observed signals are independent from one another and not the result of the same underlying input.

## Discussion

By acquiring intraoperative MER with simultaneous monitoring of physiological signals, we report that the activity of single neurons in the thalamus and STN encodes visceral signals related to cardiac and respiratory function. In particular, we find that 53 out of the 73 single neurons included in the present analysis respond to at least one of the three interoceptive signals that we analyzed, that is around 40% responded to heartbeat, 30% to IBI, and 30% to the respiratory cycle. While a large proportion of these neurons

were significantly modulated by cardiorespiratory activity, their patterns show variable response properties. We observed increases and decreases in firing rate, peaks of modulation during different phases of the respiratory and cardiac cycle, as well as positive or negative correlations between firing rate and the duration of the cardiac cycle. Finally, while most of the present neurons responded to a single type of signal, some units were found to encode two, or even all three tested interoceptive signals.

The present data significantly extend earlier preliminary evidence of neural responses to visceral inputs in the thalamus in human and non-human primates (25, 63, 74) by describing extensive neural responses to heartbeats, IBI, and respiratory cycle in the human thalamus. The thalamus is a major diencephalic structure bidirectionally relaying information between brainstems and cortical regions. Apart from olfaction, every sensory system transits to a thalamic nucleus that receives, processes, and sends information to a specific cortical area. For instance, the ventral posterolateral nucleus [equivalent to Vc according to the nomenclature of Hirai and Jones (75)] receives medial lemniscal fibers from cuneate and



**Fig. 5.** Overlap of cardiac and respiratory neurons. (A) Venn diagram showing the number of neurons among the recorded ones that responded to one or more interoceptive signals. Red circle: units responsive to the heartbeat; Green circle: units with firing rate correlated with IBI; Blue circle: units responsive to the respiratory phase. (B) Proportion of different interoceptive neurons in each of the three regions color-coded by type of response. The total number of neurons tested in each region is shown in parentheses.

gracile nuclei conveying somatosensory information to the somatosensory cortices. Similarly, visual inputs from the retina transit to the lateral geniculate nucleus before being further processed in the visual cortex. Thalamic neurons respond to a variety of stimuli, including tactile stimuli (skin mechanoreceptors) in the Vc (76, 77) and retinal inputs in the dorsal lateral geniculate nucleus (78), and are driven by proprioceptive information (passive and active joint movement) in the Vim (79, 80).

In addition to the thalamic nuclei Vim and Vc, we also recorded single cells in the STN—the most ventral part of the diencephalon situated between the thalamus and the midbrain which receives inputs from motor areas of the cerebral cortex, projects to the substantia nigra, and is reciprocally connected with the globus pallidus (81). STN is classically considered as a motor structure given its important position in the basal ganglia circuitry and its prominent function in the execution and control of voluntary movements. Accordingly, neurons in the human STN have mainly been reported to be activated during different phases of sensorimotor tasks such as action selection, movement, and in response to stop signals (82, 83). Our description of neuronal responses in STN to cardiac and respiratory inputs is thus at first sight surprising. However, the basal ganglia circuitry is also known to be involved in the control of autonomic function during respiration (independently of voluntary breathing; 84), and single neurons in the STN of cats have been reported to tonically fire in specific respiratory phases (27). There is also evidence that STN is involved in cardiorespiratory control (85). Thus, midbrain stimulation (locomotor region in decorticated cats, corresponding to STN in humans)—induces strong cardiorespiratory responses (29, 30). Moreover, changes in heart rate and blood pressure modulate spike activity in human STN (64) and STN-DBS modulates baroreceptors sensitivity (a key regulator of cardiovascular homeostasis) (62) heart rate and blood pressure (59). Extending this former evidence, here we report neurons in the human STN responding specifically to heartbeats, IBI, and respiratory cycle. The neural responses observed in STN were similar to those observed in the thalamus, both in terms of proportion of units modulated and in terms of variability of response patterns.

**Neural Responses to Heartbeats.** In the present work, we found that the heartbeat is widely encoded in the human thalamus and the STN, that is about 40% of the neurons are significantly modulated by the cardiac cycle, with larger proportions in STN and Vim, compared to Vc (only 17% of Vc units exhibited modulation). The large proportion of neurons responding to the cardiac cycle seems reasonable given the significant proportions of cardiac-related activity observed in the hippocampus and amygdala of epileptic patients (~20%) (54) and of other visceral information encoded in monkey thalamic cells (85%) (25). However, a previous study in

humans (63) only reported a very small percentage of thalamic units (<1%) exhibiting phasic modulation of firing rates related to cardiovascular activity. While representing an important and pioneering report, this study had several important technical limitations. First, neuronal spikes were only assessed by visual inspection; lack of spike sorting could have led to an underestimation of the single units recorded (because of multi-unit detection) and therefore of the units modulated. In addition, no ECG was recorded, and the authors only considered as responsive those neurons that showed a peak of activity aligned to the systolic peak pressure (measured via a pulse-oximeter on the toe) and at least 50% larger than the background firing rate. This differs from the present study where we used feature-based spike sorting and analyzed cyclical modulations (i.e., increase and decrease of firing rate) of neural activity along the entire cardiac cycle without setting an a priori threshold for response, potentially explaining the present proportion of cardiac units. Furthermore, Oppenheimer et al. performed their recordings only in Vc, which showed the lowest number of responses to the heartbeat in our dataset (17%) with only two units showing an increase in firing rate following the R-peak. Finally, Oppenheimer et al. did not perform spike clustering and only one unit would have been detected in our dataset without this important step, which is consistent with this previous work.

The cardiac neurons we observed responded to the heartbeat by either decreasing (~60% of responding units) or increasing (~40% of responding units) their firing rate following the R peak. Decreases in neural firing rate (suppressive response) in response to external stimuli have been widely observed in other brain regions, such as the primary visual cortex and the inferior temporal cortex (86). Decreasing responses in visual cortices for non-preferred stimuli are thought to be a mechanism that enhances tuning selectivity (87). Accordingly, we propose that the increase/decrease of firing rate following R-peak may represent tuning to different periods of the cardiac cycle and that, likely, the entire cardiac cycle is tracked by thalamic and subthalamic neuronal populations. Alternatively, it is known that a multitude of systems and feedback loops exist between the different visceral brainstem and midbrain nodes and such inputs may also explain the simultaneous presence of both increasing and decreasing firing rate. We note that any conclusions about the directionality of the observed responses is difficult, considering that firing rate modulations were found when time-locking neurons spikes both to the R-peak and to the end of the cardiac cycle. This is likely the consequence of the extremely small variability of the IBI intervals in our recordings (*SI Appendix, Fig. S2*), possibly due to monitored cardio-respiratory function (anesthesia during surgery) or to the movement disorder itself (88).

Concerning timing of activity, the peak of modulation for the neural activity for 77% of all units processing the heartbeat was found to fall between 200 ms and 400 ms post R-Peak, with a

significant heterogeneity of latencies (SD = 100 ms). This is consistent with animal data of subcortical visceral processing (89). This timing suggests that for many of the identified neurons, the modulation could potentially be linked to baroreceptor-related activations (given that the timing of blood ejection corresponds to ~300 ms after R-peak). This time window is also in line with the latency of HEP modulations observed with both iEEG in insular (37) and somatosensory recordings (52), and with scalp EEG (32). However, the significant variability of observed latencies and response duration indicates that the effects we observed could originate from different pathways and are not limited to baroreceptor-related activation.

The large number of modulated neurons and their variable patterns of response underscore the importance and the complexity of the interplay between cardiac signals and brain function. Several processes and different information pass through STN and thalamus and could underlie the observed neural responses to heartbeats since different kinds of inputs reach the brain at each heartbeat. These different inputs include discharges from baroreceptors in the heart wall and blood vessels as well as signals from mechanoreceptors in the chest wall carrying information about the heart's contraction timing and strength (90). Additional signals include proprioceptive or tactile peripheral receptors which detect the pulsatility of blood vessels (91–93). Moreover, evidence also exists for astrocyte monitoring of cerebral perfusion and vasculo-neuronal coupling directly in the CNS (94, 95). However, even higher levels of available oxygen could have increased the likelihood of spontaneous firing in the recorded neurons during the systole. Considering that we observed both increases and decreases in firing rates in different timing and given the sharpness of change in firing rate observed in many neurons, we argue that the multiple modulations we observed are most likely due to different mechanisms.

To summarize, we show that different phases of the cardiac cycle are encoded in a large proportion of thalamus and STN neurons. The timing of many neural responses observed is compatible with baroreceptor activation but the variability in latency and direction of the modulations indicates that multiple heartbeat-related signals are processed in these subcortical structures.

**IBI-Related Signals.** The IBI and its variability are important physiological parameters, as beat-to-beat monitoring and control of heart rate are fundamental to regulate cardiac function and ensure homeostasis in response to internal and external environmental changes (71). This precise regulation is maintained through local reflex and feedback loops involving afferent information related to cardiac function, cardiac frequency, and blood pressure from various peripheral receptors (96) that is relayed to subcortical and cortical structures. Descending projections to parasympathetic and sympathetic relays, in turn, regulate heart rate (chronotropic control), conduction (dromotropic control), and myocardial contractility (ionotropic control) (97, 98). However, very little is known about how single neuron activity in humans relates to the IBI. Two single-cell studies in humans investigated this relationship at the cortical level and have reported a significant correlation between IBIs and the firing rates of 10% of anterior cingulate cortex neurons, and 20% of medial temporal lobe neurons (54, 55). In the present study, when investigating the relationship between neural activity and the duration of the cardiac cycle, we found that the IBI is encoded in the activity of a substantial portion of thalamic and subthalamic neurons (about 30%), by showing that the firing rates of these neurons correlated with the IBI length. This is in line with evidence from animal research where the activity of 33% of somatosensory neurons in the cat thalamus was found to correlate with the IBI (28). The present data are a report of such evidence in

the human thalamus and STN and show that all three tested regions are involved in neural IBI monitoring.

Importantly, we observed both positive (~40%) and negative (~60%) correlations, meaning that during longer cardiac cycles some neurons increased their firing rate, while others decreased it. Considering the bidirectionality and the complexity of homeostatic IBI control, these different responses most likely reflect different regulatory mechanisms. For instance, they could reflect the monitoring and processing of various signals (heart rate, cardiac function, and blood pressure) to control the heart rate. Conversely, heart rate information not only influences chronotropic feedback control but also ionotropic and dromotropic parameters. We also note that the presence of both positive and negative correlations between firing rates and IBI rules out the trivial hypothesis that fluctuations in central oxygen/CO<sub>2</sub> associated with the cardiac cycle duration might underlie the IBI effects. When investigating the directionality of the observed modulation using phase response analysis, we found that only in 50% of the IBI-correlated units the firing rate had a significant effect on the duration of the cardiac cycle (lengthening or shortening), suggesting that we observed an efferent signal in only half of the units. Using the same approach for neurons in the medial temporal lobe and anterior cingulate cortex, Kim et al. (55) reported a significant phase response for all units that exhibited a significant IBI correlation. However, none of the units they identified showed a significant modulation of activity in response to the heartbeat itself, while we found that 7 out of 21 correlated units responded to the cardiac cycle.

To summarize, we observed an important number of neurons with firing rates that positively or negatively correlated with IBI in both thalamic nuclei and the STN. The differences in proportions of directional and multi-modal signals observed here in comparison with earlier reports at higher cortical structures, underscore the central role of the thalamus and STN in the regulatory feedback loops involved in the control of heart function.

**Methodological Considerations.** A challenge in evaluating neural activity relative to the cardiac cycle is the presence of heart cycle-related artifacts in the recorded signals (52). Extensive work on cardiac movement of the brain, using imaging approaches (e.g., refs. 99 and 100), has highlighted the importance of carefully considering this issue when studying neural responses to the heart. MER is known to be affected by the cardioballistic effect caused by pulsating vessels near the recording electrode and observed in the waxing and waning of the extracellular signal (101, 102). This effect can affect spike detection and sorting and generate spurious modulation of the neural activity (68). Recent work has proposed an interesting approach of post hoc motion correction for both LFP and single unit using high-density recording probes in cortical regions (103). To overcome this major issue, given the limitations of intraoperative MER (acute single channel recordings), we opted for an approach of detecting and excluding neurons showing signs of motion. We applied methodology derived from recent work on heartbeat-related modulation of EAPs (68) to identify neurons showing cyclical modulation of EAPs with the cardiac cycle (a sign that the neuron was possibly affected by the cardioballistic artifact) and excluded them from our sample. Accordingly, we included only neurons without significant EAP shape modulation along the cardiac cycle in further analyses (73 out of the 127 recorded units). This conservative approach aimed at ensuring that spurious modulations of the neural activity were not falsely counted as interoceptive responses. Importantly, we observe that the neurons detected here encoded different modalities (heartbeat, respiration and/or IBI), showing that the observed effects cannot relate to the cardioballistic artifact. This is further confirmed in the present data by showing various



types of responses, such as the increase or decrease in firing rates, the positive and negative correlations between firing rate and IBI, as well as the positive or negative PR, and the encoding of different phases of the respiratory cycle. Collectively, these findings exclude that the present changes in neural firing in STN, Vc, and Vim were merely caused by the cardioballistic artifact.

**Respiratory Signals.** Analyzing the thalamic and sub-thalamic responses with respect to the cyclic respiratory signal, we were able to identify 22 neurons (~30%) the firing activity of which was modulated by the phase of the respiratory cycle. We also report that the firings were centered around different phases of the respiratory cycle for the different neurons, with some neurons firing more likely during inspiration and others preferring expiration. To the best of our knowledge, the existence of respiratory signals in thalamic and subthalamic neurons has not been reported in humans before, but confirms previous animal data, reported from the thalamus and STN of the cat (26, 27), with about 15% of units found to correlate with the level of the respiratory drive. Corollary discharges from the brainstem respiratory neurons to the sensory cortex could be at the basis of the thalamic signals we observed; as the pre-Bötzinger complex projects directly to several subcortical structures, including the thalamus (104). In particular, the finding that the preferred respiratory phases vary across neurons is reminiscent of respiratory neurons in the brainstem whose activation is linked to different phases of either inspiration or expiration, controlling the generation of respiratory rhythms (73, 105). Alternatively, some of the observed modulations could also be linked to lung movement relayed by the vagus nerve, to mechanoreceptors in the chest wall whose projections appear to play a role in shaping respiratory sensations (106, 107), or to chest expansions-contractions relayed by somatosensory pathways. Finally, basal ganglia and thalamus activity have also been common findings in neuroimaging studies of volitional breathing as well as in investigating complex behavioral breathing acts, such as breathing during speech, while singing or exercising (12, 42, 61, 108, 109).

We observed neurons modulated by respiratory phase in all three recorded regions, without clear differences in either proportion of responsive units or in phase preference across the three recording sites. Accordingly, recent evidence from mice has shown widespread neural responses to the breathing phase in several brain regions, including the sensory and midline thalamus (110). The breathing signal has been hypothesized to act as an oscillatory pacemaker improving functional coordination between brain regions by synchronizing neural activity (110, 111). The present data show that thalamic and STN neurons may be part of this larger network of neurons involved in the coordination of brain activity. Concerning overall methodology, we also note that, given the much slower frequency of the respiratory cycle, these signals should not be affected by eventual cardiac-related artifacts or signals. Furthermore, since different neurons showed different phase preferences the present data do not appear to be linked to physiological changes during a specific phase of respiration. Additionally, a good portion of the respiration-entrained units (~40%) was found to only encode respiratory signals (no response to the heartbeat or to the IBI), thus providing further evidence that this effect is not the result of indirect cardiac modulation.

To summarize, our results show that information about the respiratory cycle is encoded in about one third of the thalamic and subthalamic neurons. The preferred spiking phases varied across neurons, suggesting that breathing-related signals in thalamic nuclei and STN may function as oscillatory pacemaker improving functional coordination between brain regions by synchronizing neural activity.

**Thalamic and Subthalamic Nuclei Encode Visceral Signals.** In this study, we identified neurons in the thalamus and STN encoding different visceral information that is related to cardiac and respiratory signals. The modulation of human thalamic neuronal activity with visceral signals is compatible with several anatomical and functional studies in animals that have demonstrated widespread connectivity between the thalamus and important visceral and interoceptive centers such as the insular cortex (18, 112, 113). Similarly, projections from STN to key structures involved in visceral processing such as the anterior cingulate cortex, the somatosensory cortex, and the insular cortex have been extensively described in rodents and primates (114–123).

Most of the neurons we identified (66%) responded to a single type of signal (either heartbeat, IBI, or respiration), and these responsive units were localized in all three investigated subcortical regions of the thalamus and STN, with no clear bias in their distribution across regions. The remaining 34% of the units were found to encode two (30%), or only very rarely all three measured modalities (4%). This is evidence of multimodal visceral neurons in the human thalamus and STN. This is consistent with data from neurons in the thalamus of non-human primates, which have shown to widely encode visceral signals (more than 80% of units responded to gastro-intestinal or bladder-related signals), with about 70% of those units encoding multimodal signals (somatic and multivisceral) (25). The present evidence for multimodal visceral information at the single neuron level and at the level of each tested subcortical region suggests that, in addition to their function as a relay to efferent and afferent information to cortical regions, the thalamus and the STN also integrate different visceral inputs and outputs. Importantly, we note that the functions of the heart and lung that we analyzed show physiologically close interactions, as illustrated for instance during inspiration-induced cardio-acceleration [referred to as respiratory sinus arrhythmia (124)]. Together with previous evidence of widespread visceral signals encoded in subcortical regions, our findings underscore the interplay between different visceral signals and between cardiac and respiratory function in particular. These present data also reveal that visceral cardiorespiratory signals are processed in all three subcortical regions, at the level of single neurons. Many of the observed neurons, encoded more than one signal at the same time, highlighting the continuous interaction between diverse visceral information at the subcortical level, essential for successful homeostasis and behavioral control. Furthermore, when analyzing responses from different cortical control regions, we found no responses to heartbeats, minimal response to respiration, and a reduced prevalence of units responding to IBI (*SI Appendix, Supplementary Results, Comparison Regions*). These results highlight the specificity of our main findings, indicating that cardio-respiratory signals are not ubiquitously and equally encoded across the brain, and underscore the possibly more prominent role of the thalamus and STN in the processing of cardio-respiratory information. More studies in the future should be conducted, both in animals and humans, to further characterize how these vital signals are differentially encoded across brain regions.

Overall, the present results demonstrate the direct functional involvement of thalamic and subthalamic single neurons in the processing of visceral signals in humans. The large number of detected units, the richness, and diversity of signals encoded are in line with the idea that these regions could be important relays of the diverse types of visceral information reaching the brain and could be furthermore involved in regulatory feedback loops that control visceral functions. Our findings can help to better characterize the functional pathway of visceral signal processing through the subcortical regions which is of importance also in several medical specializations (cardiology, pulmonology, neurology, and psychiatry), but also psychological

research. For example, the importance of the relationship between the nervous autonomic system and cardiovascular conditions, such as lethal arrhythmia, has become increasingly clear in the past decades (125), and there is increasing evidence of the role of the thalamus in autonomic regulation and disturbance (126). Visceral processing is also recognized to affect cognition and to be an important feature of some psychiatric disorders (4, 12). Direct and indirect measures of visceral signals and their neural correlates could in the future become important biomarkers of the state of the autonomic system, and our methodologies could support the exploration of those signals at the level of single cells. The development of chronically recording DBS implants and of novel closed-loop brain stimulation interfaces also represents an ideal scenario where our investigations could find both scientific and therapeutic application.

## Limitations

While we tried to address several methodological issues, our study is still limited by the nature of MER. First of all, we are aware that all our subjects are patients suffering from either PD or essential tremor, conditions that are known to affect neural computation in subcortical regions and possibly physiological parameters (88). Therefore, it is possible that these diseases affected the present responses. Furthermore, it is important to note that different subcortical regions were targeted depending on disease (PD-STN; essential tremor-thalamus). The recording of neural responses in other subcortical regions involved in interoceptive processing (e.g. nucleus of the solitary tract) was not possible in the present study, and protocols in animals investigating visceral responses along the interoceptive pathways will certainly be invaluable to disentangle the impact of the thalamus and STN. While different regions were recorded in the different patient populations, we note, however, that we observed only minor differences in the distribution of responsive units and direction of IBI correlation, while we found no difference in the latency of responses to the heartbeat and preference of respiratory phase across the different regions and hence observed similar findings whether recordings were carried out in patients with essential tremor or PD. The lack of statistically significant differences across regions could be partially explained by the variability of neural responses to interoceptive signals within each region and by the relatively small number of responsive units within each region (we may be underpowered in statistically assessing the characteristics and proportions of responses across regions). The very small IBI variability observed in our dataset may also be masking some neural responses to IBI making them undetectable. In addition, while patients are awake during MER, residual effects of anesthesia on cardio-respiratory activity cannot be completely excluded despite the 30' washout period. Monitoring of several physiological parameters, such as direct recording of air flow, and experimentally increased variability of cardio-respiratory functions will be important for future investigation but seems difficult to achieve under the current OR practice for DBS implantation. Future studies may also capitalize on direct manipulations (e.g., changes in respiratory depth, holding of breath, and Valsalva maneuver), to investigate how the neural activity in thalamic and subthalamic nuclei is affected by these changes and by baroreceptor responses. Finally, further investigations should try to disentangle to possible underlying mechanisms of the various neuronal responses to the visceral activity we observed.

## Methods

**Patients and Recordings.** We acquired intraoperative microelectrode recordings (MER) from 23 patients [nine females, mean age: 64 y (SD: 6.0 y), body mass index range 20 to 35 kg/m<sup>2</sup>] undergoing surgical implantation of DBS

electrodes in the subthalamic nucleus and thalamus for treatment of refractory Parkinson disease (N = 9) or essential tremor (N = 14), respectively. Patients with untreated depression or severe anxiety were not recruited. Recordings were performed at Ohio State University Medical Center between March 2017 and July 2018. All patients signed informed consent to participate in the study before surgery. Experimental protocol was approved by the Ohio State University Medical Center Institutional Review Board (Columbus, OH). We report data from 63 recording chunks with at least one neuron (22 from STN, 18 from Vc, 23 from Vim), the average duration of those recording blocks was 148 s (SD: 43.3 s, range 52.8 s to 273.4 s), with an average of 202 cardiac cycles recorded (SD = 71, range 60 to 373). Electrophysiological recordings were acquired with a sampling rate of 48 kHz using the FHC 4000 LP (Lead Positioner) Neuromodulation Targeting System (manufactured by FHC, Inc.). Simultaneous acquisition of electrocardiogram (ECG) was performed by means of two surface electrodes placed on the patient's chest and a reference electrode placed on the right ankle. ECG was initially sampled at 48 kHz. Respiratory signals were extracted offline from the ECG traces (*SI Appendix, Supporting Methods and Fig. S7A*). Patients were awake during the recordings; light sedation was administered at the beginning of surgery, but interrupted at least 30' before MERs. Further details about the MER procedure are reported in *SI Appendix, Supporting Methods*.

**Signal Processing and Spike Extraction.** All recorded signals were processed offline using Matlab. R-peaks were detected from the ECG signal with the HEPLAB package (127) using the "ecglib fast" algorithm for peak detections (128). Detected R-peaks were then inspected manually to exclude spurious events. Aberrant cardiac cycles with IBI < 0.5 or IBI > 1.5 (corresponding, respectively to a cardiac rate > 120 bpm and < 40 bpm) were excluded from all further computations. The heart rate variability was assessed from the SD of the IBIs for all beats (SDRR, equivalent to SDNN). The distribution of IBIs, heart rates, and SDRRs across recording chunks is shown in *SI Appendix, Fig. S2*. Electrophysiological signals were processed using the latest implementation of Wave-clus (66) to extract and sort single neuron activity. Signal was initially high-pass filtered (above 300 Hz), single spike activity was then extracted using a detection threshold equal to five times the estimated SD of the background noise ( $\sigma_N$ ). Spikes from different putative neurons were sorted using the automatic implementation of Wave-clus. Initial spike sorting was then reviewed manually to exclude noise clusters and optimize cluster separation. Assessment of spike sorting quality was then performed as detailed in *SI Appendix, Supporting Methods*.

**Neural Responses to Heartbeat.** Modulation of the neural firing rates in response to the heartbeat was tested for each neuron as follows. For each detected R-peak, we considered all neural spikes falling in a time interval around the time of the R-peak. The time interval ranged from -100 ms to the median value of the cardiac cycle duration (median IBI) for the given recording. Spike times were realigned relatively to the time of the R-peak and binned into 10 equally spaced bins. The time bins were then converted into radians to represent the angular phase along the cardiac cycle (with 0 ms corresponding to 0 rad, and the maximum time corresponding to  $2\pi$  rad). We then computed the z-statistic for the non-uniformity of circular data (Rayleigh's test) on the binned spikes and compared it to a null distribution of 1,000 surrogate statistics obtained by jittering the R-peaks (between -500 ms and +500 ms) while preserving the IBI and variability of cardiac cycles (7, 37). In order to ensure a robust estimation of modulation across trials, we additionally measured the spike-field coherence between the neural spikes and the phase of the cardiac cycle (averaged in the frequency band < 20 Hz). For a neuron to have a significant modulation of firing rates in response to the heartbeat we asked that the z-statistic for non-uniformity was larger than 95% of the surrogates ( $P < 0.05$ ) and that the coherence between spikes and cardiac phase was larger than the coherence significance level (129):

$$c = \sqrt{1 - \alpha^{1/(N-1)}},$$

with  $\alpha$  alpha equal to 0.05 significance level corrected for multiple comparisons (Bonferroni correction) and  $N$  being the number of cardiac cycles used to measure coherence. Additionally, we repeated the response analysis but this time time-locking the spikes to the end of the cardiac cycle. If the neural response was disrupted when time-locked to the end of the cycle, this would disambiguate the directionality of the effect (from heart to neuron or vice versa).

**Relationship between Spike Firing and Inter-Beat Intervals.** We assessed the relationship between neural firing and duration of the cardiac cycle as follows. For each cardiac cycle, we measured the interbeat interval (IBI) which corresponds to the time between one ECG R-peak and the following one. For each neuron, the average firing rate during each cardiac cycle (FRc, measured in Hz) was calculated as the number of spikes fired during the cycle divided by the corresponding IBI. Correlation between FRc and IBI was calculated using Pearson's correlation. Statistical significance of the correlation was assessed from t-distributions and corrected for multiple comparisons (Bonferroni correction).

**Neural Responses to Respiration.** The phase of the respiratory signal was extracted as the angular component of the Hilbert transform of the HDR signals. For each neural spike, the corresponding respiratory phase (spike-phase) was extracted as the instantaneous respiratory phase at the time of the spike. Non-uniformity of the spike density phase distribution was tested using Rayleigh's test for circular data. Significance of the modulation was assessed by comparing the z-statistic obtained with a surrogate statistic (10,000 surrogates) obtained by jittering the spike times (between  $-2.5$  s and  $+2.5$  s). Note that in this case a large interval for jittering was chosen in order to break time dependency between the spikes and the slow respiratory rhythm.

**Data, Materials, and Software Availability.** Codes for analysis were custom made in Matlab and the following toolboxes were used: EAP waveforms were

tested using partially the code from Mosher et al. (68); R-peaks were detected with the HEPLAB package (127); RRest toolbox was used for ECG-derived respiration (<https://peterhcharlton.github.io/RRest>) (130). Analysis scripts and data supporting our findings will be available in a public repository following publication (<https://osf.io/m86se>) (131).

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