Neuronal calcium spikes enable vector inversion in the Drosophila brain

Itzel G. Ishida¹, Sachin Sethi¹, Thomas L. Mohren¹, L.F. Abbott², and Gaby Maimon¹,*

¹Laboratory of Integrative Brain Function and Howard Hughes Medical Institute, The Rockefeller University, New York NY, USA
²Mortimer B. Zuckerman Mind Brain Behavior Institute, Department of Neuroscience, Columbia University, New York NY, USA

*Correspondence: maimon@rockefeller.edu (G.M.)
Summary

A typical neuron signals to downstream cells when it is depolarized and firing sodium spikes. Some neurons, however, also fire calcium spikes when hyperpolarized. The function of such bidirectional signaling remains unclear in most circuits. Here we show how a neuron class that participates in vector computation in the fly central complex employs hyperpolarization-elicited calcium spikes to invert two-dimensional mathematical vectors. When cells switch from firing sodium to calcium spikes, this leads to a ~180° realignment between the vector encoded in the neuronal population and the fly’s internal heading signal, thus inverting the vector. We show that the calcium spikes rely on the T-type calcium channel Ca-α1T, and argue, via analytical and experimental approaches, that these spikes enable vector computations in portions of angular space that would otherwise be inaccessible. These results reveal a seamless interaction between molecular, cellular and circuit properties for implementing vector math in the brain.
Introduction

Neural circuitry that implements vector mathematics has been characterized in the insect central complex\textsuperscript{1-4}. The output of one such circuit in \textit{Drosophila} is a calcium signal—evident across a genetically defined neuronal population—that tracks the fly’s traveling direction referenced to external cues, that is, in allocentric coordinates\textsuperscript{1,5}. This signal is constructed from the activity of four presynaptic neuronal populations, each of which can be modeled as encoding a two-dimensional mathematical vector\textsuperscript{1}. The lengths of these neuronally encoded vectors are controlled by the fly’s traveling direction referenced to its body axis—i.e., the egocentric traveling direction—and their angles are controlled by the fly’s allocentric heading angle. A downstream neuronal population functionally sums the four vectors to build an estimate of the fly’s allocentric traveling direction\textsuperscript{1}.

Formally, this computation could be implemented by summing only two vectors—akin to \(x\) and \(y\) components in Cartesian geometry—provided that each component vector could point in either the positive or negative direction along its respective axis. The traveling-direction circuit uses four vectors because the neuronal populations encoding each vector can only signal to downstream cells when their constituent neurons are depolarized. This rectification means that only one projection direction along an axis can be signaled and thus two vectors are needed to cover both the positive and negative directions.

We wondered whether a single neuronal population in the \textit{Drosophila} central complex might be able to encode a vector that is invertible, that is, a vector that can point in either of two directions offset by 180°. We found that a specific class of neurons, PFNa cells, can indeed do this by virtue of single PFNa cells expressing two classes of action potentials: sodium spikes induced by membrane depolarization and calcium spikes induced by membrane hyperpolarization. Sodium spikes allow the PFNa population to signal vectors pointing along one axis direction and calcium spikes allow the same population to signal vectors pointing along the inverse direction. We further found that the hyperpolarization-elicited calcium spikes in PFNa cells are mediated by T-type calcium channels, which are known for producing analogous calcium spikes and burst firing in mammalian thalamocortical circuits\textsuperscript{6,7}. Finally, we found that neurons monosynaptically downstream of PFNa cells can perform a vector-addition-like read out of the two PFNa populations when both vectors are optogenetically made to have equal lengths and point in either the sodium-spike encoded directions or the inverse, calcium-spike encoded directions. These results—supported by a model rooted in detailed anatomy and physiology—reveal a mechanism by which a central-brain circuit can circumvent canonical rectification constraints on neuronal signaling to implement invertible, two-dimensional vectors.
Results

How neuronal populations could encode invertible, two-dimensional vectors

Spatial vector computations in the fly central complex (Figure 1A) are anchored to a common, internally generated sense of heading signaled by EPG cells\(^8,9\). Individual EPG cells innervate one wedge of the circular ellipsoid body with their dendrites and one glomerulus of the linear protocerebral bridge with their axons, with the full population of EPG neurons tiling these two structures. The fly’s heading is indicated by a spatially localized calcium signal, or bump, across the EPG population. This bump rotates around the ellipsoid body as the fly turns, with its position around the ellipsoid body, or phase, tracking the fly’s heading. There are two copies of the ellipsoid body bump present in the EPG axons within the protocerebral bridge (Figure 1B). In total, the EPG system thus consists of three neural rings that represent angular space, one closed ring in the ellipsoid body and two open rings in the protocerebral bridge (Figure 1B).

In the bridge, EPG cells provide synaptic input to dozens of downstream neurons, including several classes of PFN cells\(^4\). Whereas the two EPG calcium bumps in the bridge are often narrow in shape, the downstream pair of bumps in each class of PFN cells—one in the left bridge and one in the right bridge—are broader, conforming well to sinusoidal functions across the bridge\(^1,4\) (Figure 1B). A sinusoidal reformattting between EPG and PFN bumps makes sense in light of extensive inputs that PFN cells receive from a set of bridge interneurons called \(\Delta7\) cells, whose anatomy positions them to act as sinusoidal spatial-convolutional filters\(^4,10\). The resulting sinusoidal activity profile of each PFN population appears to encode a two-dimensional vector in a so-called phasor representation. The amplitude of the sinusoid encodes the vector’s length and its peak position, or phase, encodes the vector’s angle\(^1-3,11\).

The phases of the PFN bumps in the left and right bridge are both yoked to the EPG signal and thus, in the bridge, the two PFN-encoded vectors both point in the fly’s heading direction. However, when PFN populations project from the bridge to the fan-shaped body, they do so with a precise anatomical offset that is equivalent to rotating the right-bridge vector \(\sim45^\circ\) clockwise and the left-bridge vector \(\sim45^\circ\) counterclockwise\(^1,4\) (Figures 1C and 1D). The two PFN vectors thus develop a \(\sim90^\circ\) offset between them in the fan-shaped body. PFN-recipient neurons in the fan-shaped body perform a vector sum across one or more such vector pairs. They do so by adding, column by column, the sinusoidal activity patterns across their input PFN populations\(^1\), which produces a sinusoidal activity pattern in the recipient neuronal population whose amplitude and phase match the length and angle of the vector sum of the input vectors.
Adding a pair of vectors with a 90° offset allows the output vector to point anywhere between them, that is, within a 90° sector of angular space. Other parts of angular space can only be covered by including a second vector pair\(^1,3,5\) or by allowing the vectors within the one pair to invert their direction. We thus hypothesized that there might exist situations in which PFN-encoded vectors could invert, such that each vector could align with either the positive or negative direction of its particular axis. Physiologically, when a vector needs to point in the positive direction, one would observe the standard alignment between the peak of the PFN sinusoid and the EPG bump driving it in the bridge (Figures 1B and 1D, solid lines). Conversely, when the vector needs to point in the inverse direction, there would be a \(\sim 180°\) offset between the PFN and EPG signals in the bridge (Figures 1B and 1D, dotted lines). A \(\sim 180°\) offset of a phasor is equivalent to inverting the direction of the encoded vector.

Population calcium signals in PFNa neurons signal invertible vectors

Anatomical considerations\(^4\) suggested to us that PFNa cells—a specific subtype of PFN neuron—might perform vector operations with only two vectors, where each vector is invertible. To study vector computation in PFNa neurons, we placed head-fixed flies on an air-cushioned ball and allowed them to navigate in a simple virtual environment as we imaged neural activity with a two-photon microscope (Figure 2A). The flies’ left/right turns controlled the position of a bright vertical bar on a panoramic visual display, such that the bar’s angular position tracked the fly’s heading in the virtual world akin to a distant cue in the real world, like the sun. Because PFNa neurons have been previously shown to be responsive to airflow stimuli\(^12\), they seemed poised to transform the direction of airflow from egocentric to allocentric coordinates\(^12\). Thus, we also surrounded the flies with a ring of 36 static tubes that allowed us to deliver airflow stimuli from various directions (Methods, Figure 2A).

One complication is that airflow can induce the EPG heading bump (i.e., the fly’s sense of allocentric heading) to rotate under certain circumstances\(^13\). We verified that with our protocol—brief pulses of airflow delivered in open loop from randomized directions—the EPG heading bump did not measurably rotate in response to each puff (Figure S1). In other words, the fly seemed to interpret each air puff as a disturbance arriving from a different allocentric direction rather than its body having turned in the context of a static wind direction. We could thus study how PFNa neurons signal the direction of each puff.

We co-imaged calcium in the axon terminals of EPG cells and in the dendrites of PFNa cells in the protocerebral bridge as flies walked with a closed-loop bar and experienced air puffs from various directions (Methods). The EPG bumps in the bridge were consistently measurable during these experiments. The PFNa bumps, on the other hand, were often dim, but typically became clearer during
puffs (Figures 2B and 2C). Notably, when we puffed air on a single fly from the left, the peak of the left-bridge PFNa bump was aligned to the peak of the EPG signal, but the peak of the right-bridge PFNa bump was offset from the EPG peak by ~180° (Figure 2B, dotted box). When we puffed air on the right side of the same fly, the opposite relationship was evident; the left-bridge PFNa bump was now antiphase to the EPG peak (Figure 2C, dotted box). These findings stand in contrast to what is observed in the traveling direction system, where the activity bumps of all four PFN populations always align with the EPG peak in the bridge.\textsuperscript{1,5}

For each imaging frame, we computationally shifted the EPG bumps to the same position in the bridge and rotated the PFNa bumps by this same, EPG-determined, angle. When we plotted the mean across all flies of these EPG-phase-aligned bumps (Figures 2D and S2A), we observed that the PFNa bumps in both the left and right bridge had a sinusoidal shape, consistent with these signals encoding 2D vectors (Figure S2H). Additionally, we found that, as in the example fly, the PFNa bumps could exist both in phase and antiphase with the EPG heading signal, contingent on the egocentric airflow direction (Figures 2D, 2E, and S2A). With air puffs from the front, both the left and the right PFNa bumps had their peaks aligned with the EPG peaks. With air from the side, the PFNa sinusoid contralateral to the stimulated side of the body had its phase offset by ~180° relative to the EPG bump (Figures 2D and 2E). With air from behind, both PFNa sinusoids expressed an antiphase relationship to the EPG bump (Figures 2D and 2E). Thus, both PFNa sinusoids in the bridge invert, or equivalently shift their phase by ~180°, when air puffs arrive from the back and one or the other sinusoid inverts when air comes from the side.

PFNa cells project axons from the protocerebral bridge to the fan-shaped body (Figure 1C) with an anatomical offset that shifts the peak of each PFNa sinusoid—and thus the angle of the encoded vector—by ~45° clockwise for right-bridge PFNa cells and ~45° counterclockwise for left-bridge PFNa cells (Figure 1D, solid lines). When a PFNa sinusoid is phase inverted relative to the EPG bump in the bridge, this adds an additional 180° to the encoded phasor’s angle, rotating these vectors from ±45° to ±135° orientations (Figure 1D, dotted lines). Thus, the anatomy and physiology together point to a system in which two PFNa phasors signal the projection of the airflow vector onto two orthogonal axes, and that when this projection requires one or both PFNa vectors to point in the negative direction along their respective axis they can do so. In this way, a two-vector system can signal air puffs both from the front and from the rear.

Whereas the above logic explains the direction of the PFNa vectors, it does not reveal information about their length. PFN vector length is reflected in the amplitude of the PFN phasors in the bridge and also in the PFN calcium signals in the noduli, a set of paired structures just ventral to the fan-shaped...
body (Figure 1A). The entire set of left-bridge PFNa cells innervate the right nodulus, and vice versa (Figure 1C), with the intensity of left and right noduli tracking the amplitude of the right-bridge and left-bridge PFNa vectors, respectively. Thus, if our vector inversion model is correct then as the air direction rotates 360° around the fly, one should observe two angles, offset by 180°, at which the nodulus activity of a PFNa population is high — once when the air angle aligns with the positive direction of each vector’s projection axis and once when the air angle aligns with the negative direction. Remarkably, we observed two peaks in the PFNa calcium signal in the nodulus as a function of the airflow angle around the fly, for both the left- and right-bridge PFNa cells (Figures 2F, S2B, and S2C). In previously studied PFN cells that encode vectors that do not invert, noduli tuning curves express only a single peak1,5.

The entire array of left-bridge PFNa cells receives a common input from a single LNOa cell in the right nodulus and vice versa for right-bridge PFNa cells4. These two LNOa cells are the conduits for airflow information to PFNa cells4,12. Intriguingly, when we measured the calcium response of LNOa cells, we observed tuning curves to the direction of airflow with only a single peak (Figures S2D and S2E). How is it that PFNa cells show two calcium response peaks in the nodulus as a function of the airflow direction when their inputs should cause them to maximally depolarize at only one preferred angle? Moreover, what is the mechanism that allows the PFNa sinusoids in the bridge to become phase inverted relative to the EPG signal when airflow arrives from certain directions and not others? We reasoned that electrophysiological measurements from PFNa cells might provide insight into these questions.

PFNa neurons exhibit two types of spikes poised to underlie signaling of phase-aligned and phase-inverted vectors

We performed whole-cell patch-clamp recordings from PFNa neurons in head-fixed flies navigating with a closed-loop bar while they also received air puffs from various directions (Figures 3A and 3B). Consistent with past reports12, we observed strong responses from PFNa cells to air puffs, both in their membrane potential (V_m) and in their spike rate (Figure 3B, air-puff periods; note that sodium spikes in PFNa cells have a very small amplitude when recorded at the soma and are hardly visible at the scale used in Figure 3B). The V_m of PFNa cells was not only sensitive to air puffs, but also to changes in the fly’s heading, as signaled by changes in the angular position of the closed-loop bar (Figure 3B, vertical dotted line). Consistent with the example trace, in a population of PFNa cells we observed strong tuning at the level of the mean V_m to both the fly’s heading as well as the direction of airflow (Figures 3C, 3E, S3A, S3B, S3D, and S3E). The V_m tuning curves for both variables conformed well to sinusoidal
functions (Figure S2I), as would be expected from neurons that participate in encoding vectors via a phasor representation.

Conjunctive tuning to allocentric heading and the egocentric airflow direction is what one would expect from a set of PFN neurons that aim to transform the airflow direction experienced by a fly from egocentric to allocentric coordinates. However, these results were initially confusing because they revealed PFNa cells to have Vm tuning curves to airflow with only a single peak, at approximately ±45° (Figure 3E), whereas the same cells showed double-peaked tuning curves to the same stimuli when measuring calcium in the noduli (Figure 2F). If a left-bridge PFNa cell’s Vm is most hyperpolarized when an air puff arrives from the fly’s back right (i.e. ~100-150°, Figure 3E), how is it that we observe elevated calcium for air puffs at these angles in the population to which this cell belongs?

In the course of performing these experiments, we noticed that when PFNa neurons were strongly hyperpolarized, their membrane potential showed large periodic oscillations (Figure 3B). The oscillations had a periodicity of ~2-6 Hz (Figures S4A and S4B, right panels), which is a frequency range previously reported to be enhanced in the Vm of PFNa cells12. Using power in the 2-6 Hz band as a quantitative measure of the oscillation strength, we noted robust Vm oscillations whenever the membrane potential of PFNa cells was sufficiently hyperpolarized, independently of whether the hyperpolarization was caused by the fly changing its heading toward a non-preferred direction (Figure 3F, right) or by an air puff from a non-preferred angle (Figure 3G, right). Similarly, PFNa cells expressed canonical sodium spikes when their Vm was sufficiently depolarized, independently of whether this depolarization was due to the fly’s change in heading or an air puff (Figures 3F and 3G, left).

To evaluate how PFNa neurons respond to combinations of heading and airflow direction, we plotted two-dimensional tuning curves (i.e. heatmaps) of both PFNa canonical spikes and oscillation strength as a function of these two variables (Figures 3H and 3I). We observed the strongest oscillations in response to stimuli that induce the largest membrane hyperpolarization, that is, anti-preferred heading angles and ±120° airflow angles (Figures 3H and 3I, right panels). The airflow directions that elicited Vm oscillations matched those that yielded phase inversions of the PFNa sinusoids in the protocerebral bridge in relation to the EPG phase (Figure 2E, 3H and 3I). Conversely, the airflow directions that elicited the strongest canonical spikes in PFNa cells were those that yielded phase alignments between PFNa and EPG bumps in the bridge (Figures 2E, 3H and 3I). A switch in signaling from depolarization-driven canonical spikes to hyperpolarization-driven oscillations is thus poised to underlie vector inversions in the PFNa system. These electrophysiological findings also explain how the LNOa inputs to PFNa cells, which express only a single calcium peak across airflow
directions, can drive a two-peaked calcium response in PFNa cells. One calcium peak was likely due to PFNa cells being depolarized and firing canonical sodium spikes. The other peak was likely due to PFNa cells being sufficiently hyperpolarized to express oscillations, which could also lead to elevated calcium.

**Hyperpolarization-elicited oscillations in PFNa cells are mediated by Ca-α1T calcium channels**

To test whether the second calcium peak in the PFNa noduli tuning curves stemmed from oscillatory activity, we wished to better understand the mechanism of oscillatory Vm dynamics in PFNa cells. We reasoned that if artificially hyperpolarizing a single PFNa cell could reliably trigger oscillations, this would suggest a role for intrinsic membrane conductances in the phenomenon. Consistent with this notion, we were able to routinely induce oscillations in PFNa neurons by injecting hyperpolarizing current into PFNa cells (Figures S4C, S4D, and S4G).

The oscillations in the Vm of PFNa neurons reminded us of the burst firing mode of mammalian thalamocortical neurons, where low-voltage activated, T-type calcium conductances are key to the production of rhythmic spiking\(^6,7\) (Figure 4A). Specifically, hyperpolarization of thalamic neurons relieves T-type channels from inactivation, enabling them to produce regenerative calcium spikes\(^6,7\). The *Drosophila* genome encodes a single T-type calcium channel, Ca-α1T, and a recently published data set revealed that PFNa cells express the *Ca-α1T* transcript at a 35-fold higher level than other cell types analyzed in that study\(^14\) (Figure 4B). In addition, we observed strong immunohistochemical signal from a GFP-tagged knock-in allele of Ca-α1T\(^15\), *GFP::Ca-α1T*, in the third layer of the noduli and the ventral layers of the fan-shaped body, which are regions innervated by PFNa neurites (Figure 4C). These observations suggested that a Ca-α1T-mediated conductance may endow PFNa neurons with the ability to oscillate when hyperpolarized.

We knocked down the transcript levels of Ca-α1T in PFNa neurons using RNAi\(^16\) and recorded PFNa Vm while presenting open-loop air puffs to flies navigating a virtual environment. Unlike in control flies, PFNa neurons with Ca-α1T-knockdown rarely expressed any Vm oscillations to airflow stimuli arriving from behind (Figures 4D, 4G, S3G and S3H) or when we hyperpolarized the cells with current injection (Figures S4E, S4F, S4H, and S4I). PFNa neurons with Ca-α1T-knockdown also showed diminished secondary calcium peaks, specifically to airflow stimuli arriving from behind, in their calcium tuning curves measured in the noduli (Figures 4E, 4F, 4H, and 4I). Together, these results demonstrate a critical role for T-type channels in generating the Vm oscillations of PFNa neurons, and thus we will refer to these oscillations as calcium spikes hereafter. These results also implicate the Vm oscillations in generating the calcium signals associated with air puffs from behind, which underlie the
PFNa system’s ability to invert its encoded vectors.

**A qualitative model for how PFNa neurons can invert their encoded vector**

The findings up to this point allow us to propose a conceptual model for how a population of PFNa neurons can encode a vector whose direction is invertible. Each population of PFNa cells expresses a sinusoidally shaped $V_m$ signal across a set of anatomical columns (e.g., across eight glomeruli on one side of the protocerebral bridge or across the columns of the fan-shaped body) (Figure 5A). Two (possibly overlapping) thresholds exist in the system, a sodium-spike threshold in the depolarizing direction and a calcium-spike threshold in the hyperpolarizing direction (Figure 5A). Calcium rises when the $V_m$ of a PFNa cell deviates from these thresholds in either direction. Calcium influx occurs with sodium spikes because the membrane is depolarized enough to presumably activate high-voltage activated (HVA) calcium channels. With calcium spikes, calcium influx occurs via the T-type channels directly, or, potentially also indirectly via the activation of HVA calcium channels with sodium spikes that sometimes ride on top of the calcium spikes, a point we return to later. In either case, a calcium bump that encodes a vector is induced across the neuronal population (Figure 5B, middle and top row).

LNOa input to the PFNa population in the nodulus can uniformly depolarize (Figure 5B, left) or hyperpolarize (Figure 5B, right) the PFNa population as a function of the air puff direction. When the PFNa population is uniformly depolarized, the system expresses a sodium-spike induced calcium bump that is aligned with the EPG bump (set to be $0^\circ$ in Figure 5B, left). When the PFNa population is uniformly hyperpolarized, cells $180^\circ$ away from the EPG peak are maximally hyperpolarized and the system will generate a T-type spike induced calcium bump whose peak is $180^\circ$ offset from the EPG bump, thus implementing vector inversion (Figure 5B, right). In this way, an LNOa input signal with a single tuning peak can induce two peaks in the calcium tuning curves of PFNa cells—one peak due to sodium spikes and the other due to calcium spikes—while also inverting the vector encoded by the PFNa population across the two spike modes.

**A quadratic model for computing the allocentric direction of airflow**

Motivated by the above concept for how vector sign inversion could work, we aimed to develop a formalism for the computation implemented in the PFNa network. The totality of our experimental results led us to a mathematically concise model for how the egocentric-to-allocentric transformation of airflow direction is accomplished.
The model’s overarching framework is as follows. The PFNa bumps or phasors in the protocerebral bridge are either in phase or ~180° out of phase with the EPG heading angle, thus instantiating invertible vectors. PFNa cells send axons from the bridge to the fan-shaped body with an offset that rotates the left- and right-bridge phasors by ±45°, giving rise to a pair of orthogonal vectors in the fan-shaped body, which form a basis for the allocentric direction of airflow (Figure 1D). The allocentric airflow direction can be estimated by summing the two vectors. Air puffs from the front lengthen the ±45° vectors. Air puffs from the rear invert the vectors such that they shift to pointing at ±135° (Figure 1D), allowing the system to represent directions that have negative projections along the ±45° axes. We present a detailed analysis of these vector angles in the Supplementary Text and Figure S5.

To formalize our model quantitatively, we first focused on our electrophysiological data. The \( V_{m} \) of PFNa neurons appeared to be sinusoidally modulated both by the egocentric direction of airflow and by the allocentric heading angle (Figures 3C and 3E). We therefore fit the \( V_{m} \) responses of PFNa neurons (Figure 5C, left), over the full range of heading and airflow directions, to a sum of two sinusoids, with one sinusoid representing the airflow response and the other representing the heading response (Figure 5C, right). The resultant, three-parameter fits explain 91% and 94% percent of the variance of the 216 data points in the \( V_{m} \) two-dimensional tuning-curve from the left- and right-bridge PFNa populations, respectively (Methods). The quality of these fits support the notion that the heading and airflow inputs combine additively.

To fit the spiking responses of the PFNa cells (Figures 3H and 3I), we made use of the fact that both the sodium-spike rate and the calcium-oscillation strength showed a quadratic dependence on \( V_{m} \) (Figure 5D). Using the sum-of-sinusoids model for \( V_{m} \), followed by a squaring operation, we fit both the sodium-spike rate and the calcium oscillation strength across all airflow and heading angles (Figure 5E). Specifically, we modeled the sodium-spike rate response for egocentric airflow angle \( \theta \) and relative heading angle \( H \) as \((b_{H} \cos(H) + b_{W} \cos(W \pm 45°))^2\), where \( b_{H} \) and \( b_{W} \) are fitted constants, \([\_+]\) indicates rectification, and \( \pm \) refers to left/right PFNa neurons. We modeled the oscillation strength similarly except with reverse rectification for negative values. The model accounts for 87% of the variance for spike rate and 73% for oscillation strength, averaged across left and right PFNa neurons. The model fit the patch-clamp data extremely well for 11 of the 12 air-puff angles tested and thus independently of whether cells were emitting sodium or calcium spikes (Figure 5E). One exception was that air puffs from directly behind the fly yielded a low rate of sodium spikes from PFNa neurons, which our model did not predict (Figures 3H and 3I, left panels). This anomalous measurement is unlikely to reflect a technical artifact of rear airflow not arriving to the fly (Figures S2F and S2G) and thus future work will be needed to explain the observation.
It is not unusual to model firing rates as a quadratic function of $V_m$ above some firing threshold but, in the case of PFNa neurons, the existence of calcium spikes introduces the possibility of a full, unrectified squaring operation in terms of the effective output of PFNa neurons (Methods). If we assume that this is the case, the spiking and oscillatory outputs can be combined, yielding a response proportional to the single expression $(b_H \cos(H) + b_W \cos(W \pm 45^\circ))^2$, similar to what is written above, but without the rectification. If we shift the term $\cos(H)$ by $\pm45^\circ$ to model the left and right PFNa responses in the fan-shaped body, we can write the combined output of a matched pair of PFNa neurons to a downstream target as $[(b_H \cos(H - 45^\circ) + b_W \cos(W + 45^\circ))^2 + (b_H \cos(H + 45^\circ) + b_W \cos(W - 45^\circ))]^2$. Using trigonometric identities, this expression is equal to $b_H^2 + b_W^2 + 2b_H b_W \cos(W + H)$ (Methods). The last term in this expression describes the tuning of the downstream output as being a function of $W + H$, the airflow angle in allocentric coordinates, and the full collection of these outputs across the columns of the fan-shaped body is a phasor that represents the allocentric airflow direction (Methods).

Finally, we asked whether we could sum the experimentally measured responses of PFNa neurons, rather than use the above equations, to obtain the allocentric airflow direction. To do this, we used PFNa calcium signals recorded in the protocerebral bridge. Because such signals include calcium due to ordinary spikes as well as calcium spikes, which can have different physiological consequences, we needed to introduce a factor that converts the calcium-spike signal into an equivalent sodium spike signal. Using a factor of 0.2 (Methods), we computed the allocentric airflow direction by summing the left- and right-bridge PFNa phasors appropriately shifted and weighted by this factor and then we determined the phase (or equivalently maximum) of the resulting signal. This algorithm generated an output signal that accurately tracked the actual airflow direction (Figure 5F).

**FC3 cells read out the EPG-phase-aligned vectors during air puffs**

Our modeling work argues that the PFNa vectors can, in principle, be read out to track the allocentric direction of air puffs. We wished to understand if neurons that are monosynaptically downstream of PFNa cells actually perform such a read-out operation. PFNa neurons synapse onto ten classes of columnar cells in the fan-shaped body (Figure S6A). We chose to focus on FC3 neurons (Figure 6A) because their characteristic anatomy aided us in genetically targeting them with high specificity, and also because the majority of the heading inputs to FC3 cells are mono- or disynaptically received from PFNa cells, rather than from other PFN types (Figure S6A).
We imaged calcium in FC3 and EPG neurons as head-fixed flies navigated a virtual environment and received air puff stimuli from 12 directions. Akin to EPG cells in the ellipsoid body, FC3 cells express a single bump of calcium activity whose left/right position, or phase, in the fan-shaped body changes over time (Figure S6B). Prior to the first air pulse and during inter-air-puff intervals, the phase of the FC3 bump was not consistently aligned to the EPG phase (Figure S6B, purple and black curves). During air-puff stimuli, the FC3 bump amplitude increased and its phase systematically realigned in reference to the EPG phase (Figure S6C). Specifically, in an analysis of 7 flies we found that the FC3 phase deviated from the EPG phase in linear proportion to the egocentric angle of the air puff for puffs delivered within ±55° of the midline. For air puffs delivered more peripherally than ∼55° from the midline (i.e., puffs from the sides and rear), the FC3 phase did not systematically deviate beyond 55-60° (Figures S6D and S6E). With air puffs directly from the back, we did not observe a consistent difference between the EPG and FC3 phases (Figures S6D and S6E). Given the anatomy and physiology of the PFNa system described above, these data are consistent with the PFNa front vectors, encoded by sodium spikes, being able to reposition the FC3 bump in the fan-shaped body. However, the inverted, rear-facing vectors, encoded by PFNa calcium spikes, did not appear to influence the FC3 bump in these experiments because the FC3 phase did not consistently deviate more than ±55° from the midline when flies were presented with air puffs from the side and back. (See Figure S5 and Supplementary Text for why the front vectors, alone, could enable deviations of up to ±55° rather than just up to ±45°.)

**FC3 cells read out the EPG-phase-aligned and EPG-phase-inverted PFNa vectors during optogenetic stimulation of PFNa cells**

One reason for why the FC3 bump did not accurately signal the allocentric angle of air puffs from the back could be that flies need to be in a specific behavioral state for PFNa calcium spikes to impact downstream circuitry. If so, different tasks or stimuli might ultimately reveal cases in which the inverted vectors induce downstream effects. Because we do not yet know the nature of such putative tasks and/or stimuli, we reasoned that optogenetic perturbations of the PFNa population might serve as another fruitful approach for testing whether FC3 neurons can sum both the phase-aligned and phase-inverted PFNa signals.

According to our model, uniform depolarization of both PFNa populations should extend the two EPG-phase aligned vectors. If the activity of FC3 neurons reflects the sum of these two vectors, the phase of the FC3 bump should align with the EPG phase during such PFNa depolarization. To test this hypothesis, we optogenetically depolarized PFNa cells while simultaneously imaging the FC3 and EPG
calcium bumps (Figures 6A-6C). To depolarize PFNa cells, we expressed the cation-permeable channelrhodopsin CsChrimson in them and to image the FC3 and EPG bumps, we expressed jGCaMP8s in these cells (Figures 6B and 6C). With the same 920 nm illumination light, we could simultaneously stimulate PFNa axons in the fan-shaped body, and image calcium in FC3 neurons in the fan-shaped body and EPG neurons in the ellipsoid body (Figure 6B). We found that with optogenetic depolarization of PFNa cells, the FC3 and EPG phases became more aligned (Figures 6D-6G), consistent with our model. This effect was clear both during airflow (Figure 6E) for which we observed muted deviations of the FC3 phase from the EPG phase for air puffs from the sides and rear—and during moments when no airflow was presented (Figures 6F and 6G).

We also performed the converse experiment, optogenetically hyperpolarizing PFNa cells while simultaneously imaging the FC3 and EPG calcium bumps (Figures 6H-6L). Uniform hyperpolarization of both PFNa populations should induce calcium spikes in PFNa cells and thus extend the two EPG phase-inverted vectors (Figure 6H). If the inverted vectors can impact downstream physiology, the phase of the FC3 bump should show a 180° offset relative to the EPG bump during such a perturbation. To hyperpolarize PFNa cells, we expressed the chloride-selective channelrhodopsin GtACR19 in them. Otherwise, the experiment we performed was identical to the CsChrimson experiment just described. We found that with PFNa cells hyperpolarized, the FC3 bump was consistently antiphase to the EPG bump (Figures 6I-6L), supporting our model and providing evidence for the ability of the calcium-spike-associated vectors to impact downstream physiology. This effect was clear both during air puffs (Figure 6J) and during moments when no air puffs were presented (Figures 6K and 6L).

Eliminating the ability of neurons to express T-type calcium spikes should eliminate phase-inversion of the FC3 bump relative to the EPG bump with PFNa hyperpolarization. We repeated the experiment in Figure 6H in Ca-a1T-null flies (Ca-a1T Δ135), which are expected to lack functional T-type calcium channels (Figure 6M). Consistent with our model, the FC3 bump no longer appeared phase inverted relative to the EPG bump in these experiments (Figures 6N-6Q). These results support the hypothesis that PFNa calcium spikes can induce postsynaptic FC3 activity, with the phase of the FC3 bump reflecting a vector sum of the two phase-inverted vectors expressed in the PFNa populations. We verified that optogenetic hyperpolarization of PFNa neurons induces T-type calcium spikes in non-mutant flies (Figure S4K). In the Discussion, we consider why these spikes might have led to fast transmitter release in the optogenetic experiments but not in the air-puff experiments. Taken together, our optogenetic experiments demonstrate that both EPG phase-aligned and EPG phase-inverted vectors in PFNa cells can induce vector-addition-like signaling in downstream cells.
Discussion

What is the function of PFNa cells?

PFNa cells receive inputs related to the fly’s allocentric heading angle in the protocerebral bridge and inputs related to the egocentric direction of airflow in the noduli (Figure 1C). Using these inputs, two PFNa populations—one originating in the left bridge and the other in the right bridge—generate two calcium signals that are sinusoidally modulated across space, which function as invertible vectors (Figures 2 and 3). By summing the two invertible vectors signaled by the two PFNa populations, the allocentric direction of airflow (or, potentially, other directional stimuli that activate PFNa cells) can be calculated (Figure 5F). In situations where flies are standing still, or walking slowly, the direction of airflow sensed by the body directly reflects the direction of wind in the external world. The direction of wind is of broad importance to navigating insects²¹-²⁹, and thus this circuit has the potential to generate a signal³⁰ that can guide many homing and food-orienting behaviors that rely on wind assessments. Indeed, impairing PFNa physiology has been shown to impact airflow related orienting in tethered, flying flies¹².

The airflow tuning of PFNa cells likely originates from indirect inputs that these cells receive from the antennae⁴,³¹,³². Airflow tuning makes sense because each antenna houses a directional airflow sensor called the Johnston’s organ³³,³⁴. Beyond airflow, the Drosophila antennae also house sensors for temperature³⁵,³⁶, sound³⁷,³⁸, and the angle of the fly’s body in relation to gravity³⁹,⁴⁰. If PFNa neurons were to receive inputs related to these other variables, they could transform angles associated with those modalities into allocentric coordinates as well. It is also possible that PFNa cells function to transform an “angle of current relevance” into allocentric coordinates and that this angle transcends any specific sensory modality. In this view, our air puff stimuli simply emphasized one direction around the fly as being important in the moment, thus engaging the PFNa network. Regardless of the ultimate function(s) of PFNa cells, our work reveals that these neurons can implement an egocentric-to.allocentric coordinate transformation by instantiating invertible vectors.

A common allocentric reference frame in the fan-shaped body

The egocentric-to-allocentric coordinate transformation of the air puff direction implemented by PFNa cells is analogous to the coordinate transformation implemented by other PFN neurons in relation to the fly’s traveling direction¹,⁵. The fan-shaped body thus seems to house multiple calcium bumps that track navigationally relevant variables in a common allocentric reference frame. The positions of bumps along the left-right axis of the fan-shaped body map to allocentric angles in the outside world (i.e.,
Having multiple variables, or calcium bumps, expressed in a common allocentric reference frame is likely to facilitate downstream navigational calculations. By storing a trace of a given bump’s position in the fan-shaped body, for example, a fly could store a memory of an angular direction of relevance, thus allowing the fly to use variables dissociated from its current orientation or sensory experience. There are seven additional PFN cell types in the Drosophila central complex that have yet to be studied physiologically. An important next step is to identify the variables that these other PFN neurons place within this common reference frame.

**Synaptic transmission and T-type calcium spikes**

Classic synaptic release relies on calcium entry through high-voltage-activated calcium channels, which are found very close to the synaptic-vesicle fusion machinery. Because low-voltage activated calcium channels, like T-type channels, do not typically reside immediately adjacent to active zones, calcium entry through these channels is not commonly thought to drive vesicle fusion. The air-puff responses of FC3 neurons, which are monosynaptic recipients of PFNa input, generally support this dichotomous view of calcium signaling at PFNa synapses. That is, the phase of the FC3 bump quantitatively tracked the allocentric direction of air puffs when these arrived to the front of the fly and were thus signaled by presynaptic sodium spikes; however, the phase of the FC3 bump did not properly track the allocentric direction of air puffs when these arrived to the rear of the fly and were thus signaled by presynaptic calcium spikes (Figure S6E). With open-loop air puffs, it seemed that only the sodium-spike-mediated PFNa vectors were able to influence the phase of the downstream FC3 cells.

In contrast to open-loop air puffs, the FC3 phase was consistently offset by ~180° from the EPG phase—thus signaling the rearmost allocentric angle—when we induced calcium spikes via optogenetics in both the left and right bridge PFNa populations (Figure 6K). Notably, we often observed sodium action potentials expressed at the peak of calcium spikes when we injected hyperpolarizing current into PFNa cells with a patch pipette (Figure S4J), but we did not typically observe such sodium spikes with air puff-induced hyperpolarization of PFNa neurons. We thus speculate that we reliably triggered sodium action potentials at the peak of each calcium spike with two-photon optogenetic stimulation of PFNa cells. With rear air puffs, on the other hand, we may not have triggered any sodium spikes on top of the calcium spikes, and the calcium spikes acting alone may not have been able to induce synaptic transmission. One idea is that a specific modulator state was not engaged during our open-loop air puffs, which muted the expression of sodium spikes and thus synaptic transmission. Optogenetic stimulation, on the other hand, may have been sufficiently strong to bypass the need for such a modulatory input.
channels can evoke is limited, our optogenetic stimulation is likely to have induce a physiologically relevant state.) Sodium spikes riding on T-type calcium spikes have been observed in other types of *Drosophila* neurons\(^4^6\).

Because sodium spikes do not always ride on top of calcium spikes in PFNa neurons, and because calcium spikes have very different timescales and potential downstream effects than sodium spikes, it is important to note that our results to date are agnostic as to whether the central complex can combine one sodium-spike (phase-aligned) phasor (say, in the left bridge) with one calcium-spike (phase-inverted) phasor (say, in the right bridge) to perform a vector sum. Our quantitative model argues that this sort of interaction has the potential to generate an accurate allocentric angle as its output—with appropriate weighing of the two signals—but it is possible that the two modes of signaling do not interact in the real brain. Future work will be needed to resolve this matter.

Beyond eliciting fast neurotransmitter release from clear synaptic vesicles, the wide (~200 ms) calcium spikes elicited by T-type calcium channels are also well suited for promoting peptide release from dense core vesicles\(^4^7^-^4^9\). The release of neuropeptides into the fan-shaped body might not be immediately apparent at the level of postsynaptic calcium. Rather, neuropeptides may elicit downstream molecular processes that are sinusoidally modulated in their intensity across the left/right extent of the fan-shaped body, representing a vector memory in the system that can alter navigation-related computations at a later timepoint.

**Vector integration and T-type calcium spikes**

T-type calcium spikes in PFNa neurons might also serve intracellular signaling roles, rather than, or in addition to, synaptic transmission-related roles. For example, calcium-calmodulin dependent protein kinase II (CaMKII) is particularly sensitive to calcium oscillations at the 2-6 Hz frequency range\(^5^0\), which matches the calcium spike rate of PFNa neurons. CaMKII might therefore integrate the information available in calcium spikes over time. Because the calcium spike amplitude (i.e., the power of the 2-6 Hz \(V_m\) oscillation) and thus calcium influx at 2-6 Hz, varies sinusoidally across the left/right extent of the fan-shaped body, a CaMKII integral of this process—expressed across the array of PFNa axon terminals in the fan-shaped body—could represent a vector that grows in amplitude with each T-type spike across the population. Such an integrated vector could indicate for how long airflow has arrived at the fly’s rear, for example, which might be useful for driving orienting behaviors. Past work had hypothesized that PFN neurons might function as vector integrators\(^3\) and the calcium spikes described here provide a potential mechanism for this idea to be implemented. Calcium potentials could also mark
synapses as being eligible for plasticity\(^5^1\) (e.g. via CaMKII phosphorylation\(^5^2\)), creating a sinusoidally modulated vector-trace signal across the fan-shaped body in this manner as well.

**Neural computation and T-type calcium spikes**

Beyond the fly central complex, broad calcium spikes have been observed in the giant motor axons of the jellyfish *Aglantha* and in neurons of the mammalian inferior olive\(^5^3\). Delta rhythms in thalamocortical networks, which rely on T-type calcium channels\(^5^4,5^5\), are a famous correlate of sleep\(^7\). Hippocampal and cortical networks express oscillatory dynamics in the delta and theta range during navigational tasks, and the functions of these oscillations are still being studied\(^5^6–5^8\). Our work shows that T-type calcium channels serve a quantitatively precise vector-computing function in the central complex of *Drosophila*. Similarly explicit, real-time computational functions for calcium spikes likely await discovery in other neurobiological systems as well.
References


Figure 1. A framework for implementing vector inversions in the *Drosophila* central complex. (A) Schematic of the *Drosophila* brain and the central complex. (B) EPG neurons tile the ellipsoid body with their dendrites and the protocerebral bridge with their axons. In the ellipsoid body, the EPG dendrites express a single bump of calcium activity whose position around the structure tracks the fly’s heading. Due to anatomy, one copy of the heading bump exists in the EPG axons of the left protocerebral bridge, and a second copy exists in the right bridge (top, black dots and curves). The two copies of the EPG bump slide along the bridge when the fly’s heading changes. EPG neurons drive activity in varying downstream columnar cells whose dendrites tile the bridge, including PFNa neurons (middle, red, and blue traces). The shape of the two PFNa bumps is expected to be sinusoidal, with their phase (i.e., their position in the bridge) and amplitude representing the angle and length of two vectors, in a so-called phasor representation [1,3,5]. If the two peaks of the PFNa sinusoids could invert (i.e. rotate 180°), then this would be equivalent to inverting the vector they encode (bottom, red and blue dotted traces). (C) Left- and right-bridge PFNa neurons project to the fan-shaped body with anatomical offsets that introduce, approximately, ±45° rotations in the angles of the two vectors they encode. Those PFNa neurons that receive EPG (heading) input in the left bridge also receive LNOa input in the right nodulus, i.e., on the opposite side of the midline. The nodulus inputs to PFNa neurons are tuned to the direction of airflow experienced by the fly [12]. (D) Graphical representation of how the two sinusoidal activity bumps in the PFNa neurons of the protocerebral bridge are transmitted to the fan-shaped body. In the bridge, the left- and right-bridge PFNa sinusoids are phase aligned to the EPG signal on each side of the bridge (red and blue solid curves at the top) and these sinusoids are offset by ~±45°, due to the PFNa anatomy, when they get transmitted to the fan-shaped body (red and blue solid curves inside the fan-shaped body schematic). As a result, the two vectors that these PFNa phasors represent are offset by ~±45° in the fan-shaped body (red and blue solid vectors in diagram at the bottom). If either PFNa sinusoid in the bridge were to invert its phase (red and blue dotted curves at the top) then this would invert the direction of the encoded vector (red and blue dotted vectors in diagram at the bottom).
Figure 2. PFNa neurons express activity bumps that can be both in-phase and antiphase relative to the EPG bumps in the protocerebral bridge. See next page for figure legend.
Figure 2. PFNa neurons express activity bumps that can be both in-phase and antiphase relative to the EPG bumps in the protocerebral bridge. (A) Imaging neural activity in PFNa neurons in head-fixed flies walking in a virtual reality environment. The flies receive visual feedback in closed loop on their orientation in the environment (blue bar) alongside air puffs, in open loop, from varying directions (see main text and Methods for details). PFNa neurons cross the midline in their anatomical projections from the protocerebral bridge to the fan-shaped-body/nodulus; we always depict right-bridge/left nodulus PFNa neurons in blue and left-bridge/right nodulus neurons in red. (B) Simultaneous calcium imaging of EPG and PFNa neurons in the protocerebral bridge. Gray heatmap: jRGECO1a signal from EPG neurons. Red and blue heatmap: GCaMP7f signal from PFNa neurons. The estimated phases of the EPG, left-bridge PFNa, and right-bridge PFNa activity bumps are shown in the third column. The EPG phase on each side of the bridge is shown with a black trace. The PFNa phases are depicted as dots whose opacity scales with the length of the population vector average (PVA) for the relevant bump (i.e., with the size of the bump); data points below PVA strength of 0.15 are not displayed. Air puff time period (4 s) and direction (-90°, from the fly’s left) is indicated. (C) Same as B, but for an air-puff stimulus from 90° (from the fly’s right). (D) Population-averaged, phase-nulled EPG bumps (gray traces) (Methods). We rotated the PFNa bumps by the same amount as the EPG bumps on each frame, averaged the resultant signals, and plotted the population-averaged PFNa curves over the EPG curves (red and blue traces). We show overlaid curves for four of the twelve air puff angles tested (0°, −90°, +90°, and 180°). (E) Absolute difference between the phase of the left- and right-bridge PFNa bumps as a function of the airflow direction. 0° corresponds to perfect alignment of the PFNa phase and the EPG phase on either side of the protocerebral bridge. Thin lines show the averaged data from single flies. Bold lines show the population averages. (F) Mean PFNa activity in the noduli as a function of the airflow direction. Thin lines show the averaged data from single flies. Bold lines show the population averages.
Figure 3. PFNa neurons fire canonical spikes when depolarized and express oscillations when hyperpolarized, with the oscillations signaling airflow stimuli from the egocentric rear. See next page for figure legend.
Figure 3. PFNa neurons fire canonical spikes when depolarized and express oscillations when hyperpolarized, with the oscillations signaling airflow stimuli from the egocentric rear. (A) Experimental setup. (B) Example trace showing the membrane potential ($V_m$) of a right-bridge PFNa neuron. Air puff moments indicated. The angular position of the closed-loop bar on the visual display, which tracks the fly's heading, is shown underneath. We highlight a moment of a large heading change (dotted line) and a moment showing oscillations in the cell's $V_m$. (C) Tuning of PFNa $V_m$ to the fly's heading, as estimated by the angular position of the closed-loop bar on the visual display. All tuning curves have been phase-aligned to have their peak at 0° (Methods). Left-bridge PFNa neurons: red. Right-bridge PFNa neurons: blue. Thin lines: single fly averages. Thick lines: population averages. We indicate the mean $V_m$ at baseline on the right (~63 mV). (D) Schematic of PFNa neurons, showing our color conventions. (E) Airflow direction tuning curves of PFNa neurons. These tuning curves have not been phase-nulled. Thin lines: single fly averages. Thick lines: population averages. (F) Example traces from the same PFNa neuron shown in in panel B. We isolated two moments in time where the fly turned on the ball and thus changed heading (dotted lines). In one moment, this heading change caused the cell to depolarize and fire sodium spikes (left). In the other moment, the fly's turn caused the cell to hyperpolarize and express large $V_m$ oscillations (right). (G) Example traces from the same PFNa neuron shown in in panels B and F. We isolated two moments where the fly experienced airflow stimuli (gray box) that strongly altered the PFNa cell's $V_m$. With one airflow stimulus, the cell depolarized and fired sodium spikes (left). With the other airflow stimulus, the cell hyperpolarized and expressed large $V_m$ oscillations. (H) Conjunctive tuning of left-bridge PFNa-cell activity to the direction of airflow and heading. Each two-dimensional heatmap shows the population averaged tuning of 10 cells to the two variables. Heading tuning data from individual cells were rotated to make 0° the preferred heading angle, prior to averaging across flies. The left panel shows the tuning of the canonical, sodium-spike signal across the cell population, using the power in the 2-6 Hz band of the $V_m$ as a proxy for oscillation strength (Methods). (I) Same as panel E, but for the right-bridge PFNa neurons.
Figure 4. The T-type calcium channel Ca-α1T mediates oscillations in PFNa neurons and thus the ability of these cells to signal rear airflow stimuli. See next page for figure legend.
Figure 4. The T-type calcium channel Ca-α1T mediates oscillations in PFNa neurons and thus the ability of these cells to signal rear airflow stimuli. (A) Schematic of how T-type calcium channels might contribute to noncanonical calcium spikes, based on the physiology of cat thalamocortical neurons\(^7\). T-type channels inactivate at depolarized \(V_m\) and recover from inactivation upon membrane hyperpolarization. After sufficient time at a hyperpolarized \(V_m\), a large enough pool of channels is relieved from inactivation, enabling the cell to fire a calcium spike (two shown). In some systems, at the top of each T-type spike, the cell will fire a barrage of sodium action potentials. (B) Relative abundance of the RNA transcripts encoding the Ca-α1T channel in a published dataset of 67 cell types in the adult \textit{Drosophila} brain\(^{14}\). The relative abundance of the Ca-α1T transcript in PFNa neurons is indicated (arrow). (C) Immunohistochemically-amplified GFP signal in the brain of an example fly expressing a GFP-tagged Ca-α1T channel (i.e., a GFP::Ca-α1T fusion protein)\(^{13}\). The same staining pattern was observed in two additional brains (data not shown). A neuropil stain (bruchpilot) is shown in magenta and GFP fluorescence is shown in white. (D) 2-6 Hz power (left panel) and canonical spike rate (right panel) as a function of membrane potential in PFNa neurons recorded in flies from the empty-RNAi control genotype (the genetic background used to create the TRiP RNAi libraries\(^{18}\), but where eGFP was expressed in PFNa neurons). Responses of right- and left-bridge PFNa neurons were pooled. Thin lines: single fly averages. Thick lines: population averages. (E) Temporal profile of jGCaMP7f responses in the noduli from PFNa neurons. Shown is the population averaged signal from flies expressing GCaMP7f and an empty-RNAi construct in PFNa cells. The time course of the calcium signal at two airflow angles (-120° and 120°) is shown. The thick lines represent the population-mean z-score value and the error regions indicate s.e.m. (F) Population-averaged noduli tuning curves of PFNa jGCaMP7f responses. We averaged the z-scored calcium signal value between 2 and 4 seconds after the onset of airflow. Thick lines represent the population-averaged z-score calcium signal. Error bars: s.e.m. (G) Same as in panel F, but in PFNa cells carrying the construct TRiP:HMS01948\(^{15}\), which allows for expression (under UAS control) of a double-stranded RNA that targets Ca-α1T transcripts for degradation (Ca-α1T RNAi). (H) Same as in panel E, but in PFNa cells expressing jGCaMP7f and TRiP:HMS01948 (Ca-α1T RNAi). (I) Same as in panel F, but in PFNa cells expressing jGCaMP7f and TRiP:HMS01948 (Ca-α1T RNAi).
Figure 5. A qualitative and quantitative model for vector computation with invertible vectors in PFNa neurons. See next page for figure legend.
Figure 5. A qualitative and quantitative model for vector computation with invertible vectors in PFNa neurons. (A) Schematic of the population activity pattern of a set of eight central complex neurons, arrayed across a structure (e.g., one side of the protocerebral bridge or the fan-shaped body). A sinusoidally modulated pattern of activity across the population is reflected in the $V_m$. We assume that the population has two spike thresholds. One, canonical, sodium spike threshold and one, non-canonical, calcium-spike threshold. (B) If an external input uniformly depolarizes the population, the sinusoidal $V_m$ signal of the population uniformly rises (left column, bottom). The further above the sodium-spike threshold that a neuron’s $V_m$ is, the stronger it will fire sodium spikes and the more calcium will enter the cell. Thus, with uniform depolarization of the population, one expects a sinusoidal, or nearly sinusoidal, calcium signal across the population with a peak at $0^\circ$ (left column, middle). If we assume that the sinusoidally modulated calcium signal encodes a vector, then the vector would have an angle of $0^\circ$ (left column, top). Conversely, if an external input uniformly hyperpolarizes the population, then the sinusoidal $V_m$ signal across the population will drop (right column, bottom). The further below the calcium-spike threshold a neuron’s $V_m$ is, the stronger its T-type spike signal will be and the more calcium will enter the cell. Thus, with uniform hyperpolarization of the population, one expects a sinusoidal, or nearly sinusoidal, calcium signal across the population with a peak at $180^\circ$ (right column, middle). If we assume that the sinusoidally-modulated calcium signal encodes a vector, then the vector would have an angle of $180^\circ$, which is phase-inverted relative to the original case (left column, top). (C) Comparison of the airflow-nulled and heading-nulled normalized $V_m$ responses of the PFNa neurons (left panel) and the fit to a sum of two sinusoids representing heading and airflow, respectively (right panel, see Methods). The 2-D histograms show data across 18 bins of heading directions (x axis) and the 12 tested airflow directions (y axis). To align data for all cells, we nulled the heading by shifting the data to center each cell’s preferred direction at $0^\circ$, and we nulled the airflow responses by first mirroring the data for the left-bridge cells, and then applying a $-45^\circ$ shift to the pooled data for both hemispheres. (D) Sodium and calcium spiking responses of the PFNa neurons as a function of membrane potential, mean across 20 cells. Each curve has been normalized to a maximum value of 1 and a minimum value of 0 to highlight the relationship between them despite the difference of their individual units (spikes/s and 2-6 Hz power, respectively). The responses were plotted as a function of the normalized membrane potential, which is simply the membrane potential minus the mV value for the minimum value of the heading tuning curve (Methods). The two normalized curves for spiking rate and 2-6 Hz power were summed (gray line) and a quadratic fit was performed on these values (red line). (E) Comparison of the airflow and heading-nulled spiking and oscillation responses of the PFNa neurons (left panel) and the prediction of the quadratic model (right panel). The data panel on the left shows normalized and pooled values for the data in panels 3H and 3I, pooled across brain hemispheres and nulled both by preferred airflow and by preferred heading direction. (F) Predicted output direction of the PFNa neurons using the measured, summed calcium responses in Figure S2A (open purple circles). Because the phase-aligned and phase-inverted PFNa signals have different calcium sources and thus different amplitudes in terms of their jGCaMP7f responses, we scaled the phase-inverted calcium responses by a uniform factor of 0.2 to generate this graph.
Figure 6. FC3 neurons can functionally sum two sodium-spike or two calcium-spike encoded vectors in the PFNa populations.

(A) Anatomy of the PFNa and FC3 neurons. PFNa neurons shown at left synapse onto the FC3 neurons. A single FC3 neuron is shown on the right. (B) Schematic of the experimental setup. For all the experiments in this figure, we imaged the EPG neurons in the ellipsoid body and the PFNa neurons at the axonal terminals in layer 2 of the fan-shaped body. The bump in the FC3 neurons is aligned across layers 2 (dendrites) and 5 (axonal terminals) of the fan-shaped body (data not shown); in these experiments, we report the FC3 axonal signals in layer 5 because they were easier to segment accurately. (C) Schematic of the expected effect of depolarizing PFNa neurons with CsChrimson on the sinusoidal signals that PFNa populations express and the vectors that these population signals encode. The sinusoids shown schematize the population activity pattern of PFNa neurons in the protocerebral bridge, expected effect of depolarizing PFNa neurons with CsChrimson on the sinusoidal signals that PFNa populations express and the vectors that these population signals encode. The sinusoids shown schematize the population activity pattern of PFNa neurons in the protocerebral bridge.

(D) Example EPG and FC3 bumps in the context of optogenetic depolarization of PFNa neurons. The values shown here correspond to the average phase difference over the last two seconds of the airflow stimulus. Error bars represent s.e.m. The three Gal4 lines used to drive effector expression are written above the panel. (E) Same as in panel D, but for optogenetic hyperpolarization of PFNa neurons via GtACR1.

(F) Probability distributions of the EPG-FC3 phase difference in the context of optogenetic depolarization of PFNa neurons. The data shown here are taken from time periods with no airflow. Thin lines: single fly averages. Thick lines: population averages. The three Gal4 lines used to drive effector expression are written above the panel. (G) Same as in panel C, but for optogenetic hyperpolarization of the PFNa neurons via GtACR1.

(H) Circular difference of the average density of EPG-FC3 phase difference curves in shown in panel K. Error bars: s.e.m. The three Gal4 lines used to drive effector expression are written above the panel. (I, J, K) Same as in panels D, E, and F, but for optogenetic hyperpolarization of PFNa neurons via GtACR1. (L) Circular difference of the average density of EPG-FC3 phase difference curves in shown in panel K. Error bars: s.e.m. (M) Same as in panel H, but in the context of flies harboring a null mutation of Ca-α1T (Ca-α1T<sup>Ca,T</sup>). (N, O, P, Q) Same as in panels I, J, K, L but in the context of flies harboring a null mutation of Ca-α1T (Ca-α1T<sup>Ca,T</sup>).
Figure S1. EPG neurons track the fly’s heading equally well in the presence and absence of open-loop airflow pulses. See next page for figure legend.
Figure S1. EPG neurons track the fly’s heading equally well in the presence and absence of open-loop airflow pulses. (A) Example trace of EPG activity in the ellipsoid body in the closed-loop visual and open-loop airflow experiment. The ellipsoid body is split and unfolded as to display the donut as a single 16-wedge long array over time (gray color mesh), the color corresponds to the intensity of activity along each sector at every imaging frame. The position of the EPG activity peak is the population vector average estimate along the ellipsoid body, called the EPG phase (shown in black lines). The position of the visual stimulus on the LED screen is shown as a gray line. (B) Correspondence between the EPG phase and the bar position as a function of the 12 different air puff directions. The arbitrary offset between the EPG phase and bar position was zeroed at the beginning of each airflow trial, to highlight perturbations of the EPG phase-to-bar correspondence during the stimulus period. Average of 9 flies. (C) Average values of the last two seconds of airflow stimulation for the data shown in B. 9 flies. (D) Correspondence between the EPG phase position to the visual stimulus position on the screen in an example fly. The visual stimulus is never off: in the “airflow off” condition (left panel), only the closed-loop visual stimulus is present. In the “airflow on” condition (right panel), both the closed-loop visual stimulus and the open-loop airflow are present. The circular correlation coefficient $\rho$ is noted. (E) Circular correlation between the EPG phase and the bar position for 9 flies, with airflow on and airflow off. Gray dots show the “airflow off” condition, blue dots show the “airflow on” condition, and thin lines pair values for individual flies. The arrow highlights the data points corresponding to the example fly in D. (F) EPG phase velocity as a function of the visual stimulus velocity in an example fly. The unity line is shown in dotted black. Pearson’s R is noted. (G) Correlation between the EPG phase velocity and the visual stimulus velocity in 9 flies. Gray dots show the “airflow off” condition, blue dots show the “airflow on” condition, and thin lines pair values for individual flies. The arrow highlights the data points corresponding to the example fly in F. (H) Quantification of the arbitrary offset between the EPG phase and the visual stimulus for an example fly. The circular difference between the EPG phase and the bar position on the screen was divided into 36 bins of 10°, and the bin with the highest counts (labeled by the red line) was selected as the offset for the airflow on and airflow off conditions for each fly. (I) Summary of the arbitrary offset between the EPG phase and the bar position on the screen for 9 flies. Each point in this plot represents the mean difference between the offset-nulled EPG phase and the bar position over the last 2 s of the airflow stimulus. Gray dots show the “airflow off” condition, blue dots show the “airflow on” condition, and thin lines pair values for individual flies. The arrow highlights the data points corresponding to the example fly in H.
Figure S2. Tuning curves to airflow stimuli in the PFNa system. See next page for figure legend.
Figure S2. Tuning curves to airflow stimuli in the PFNa system. (A) Average data for the PFNa neurons’ phase alignment (red and blue traces) relative to the EPG heading neurons (black trace); we show the full series of airflow angles tested. We nulled the EPG phases on both sides of the bridge independently and at every imaging frame, corresponding to one time point. The EPG phase estimate at each time point was aligned to the middle glomeruli in a virtual protocerebral bridge. We then shifted the PFNa phase using the EPG phase estimate to determine its peak relative to the EPG neurons’ phase. (B) Time course of mean PFNa population activity, as measured via calcium signals, in the noduli as a function of airflow direction. The left nodulus is shown in blue, and the right nodulus is shown in red. The egocentric airflow direction is shown above each panel, and the time of airflow stimulus presentation is flanked by the dashed gray lines. (C) Tuning curve of PFNa-derived calcium signals in the noduli to egocentric airflow direction. Each point in this plot represents the mean value of the noduli z-score over the last 2 s of the airflow stimulus presentation in B. Note that the tuning curves for PFNa population activity in the noduli are double-phased. (D) Time course of mean LNOa population activity in the noduli as a function of airflow direction. The LNOa neurons are one of the numerically dominant synaptic inputs to the PFNa neurons in the noduli. The left nodulus is shown in blue, and the right nodulus is shown in red. The egocentric airflow direction is shown above each panel, and the time of airflow stimulus presentation is flanked by the dashed gray lines. (E) Tuning curve of PFNa-derived calcium signals in the noduli to egocentric airflow direction. Each point in this plot represents the mean value of the noduli z-score over the last 2 s of the airflow stimulus presentation in D. Note that the tuning curves for LNOa calcium have a single peak. (F) Comparison of the phase-nulled calcium signals of PFNa neurons in the absence of airflow (left panel) and when airflow is blown from directly behind the fly (180°, right panel). The horizontal dotted line marks the highest mean calcium responses in the no airflow condition, to aid comparison of the values in the two panels; note that both sinusoids peak at values above the line in the 180° airflow condition. (G) Reproduction of data from panels B and D (left two panels showing GCaMP signals), plus GFlamp170 signals in PFNa neurons measured at the noduli in response to airflow presented from the egocentric rearmost angle (180°). GFlamp1 is a cAMP reporter, but displays non-negligible cGMP binding as well. Note that whereas PFNa calcium shows minimal change during the stimulus presentation, the LNOa neurons are responsive. In addition, GFlamp1 signal rises within the PFNa neurons themselves shortly after the stimulus onset. (H) Sinusoidal function fits to the average phase-nulled PFNa calcium bumps elicited by frontal airflow (left panel) and airflow from the rear (right panel). The fit is shown as a dotted black line; the data is replotted from panel A, using the same color conventions. (I) Sinusoidal function fits to the average baseline-subtracted Vm, as a function of heading direction and airflow. The fit is shown as a dotted black line; the data is replotted from Figure 3C and 3E. Data from left-bridge-innervating PFNa neurons is shown in red, whereas data from right-bridge-innervating neurons is shown in blue.
Figure S3. Raw and normalized $V_m$ tuning curves from PFNa neurons. See next page for figure legend.
Figure S3. Raw and normalized Vm tuning curves from PFNa neurons. (A) Tuning of raw PFNa Vm to the fly’s heading, as estimated by the angular position of the closed-loop bar on the visual display. All tuning curves have been phase-aligned to have their peak at 0° (Methods). Left-bridge PFNa neurons: red. Right-bridge PFNa neurons: blue. Thin lines: single fly averages. Thick lines: population averages. Data from 20 cells (10 cells per hemisphere). (B) Raw Vm airflow direction tuning curves of PFNa neurons. These tuning curves have not been shifted. Thin lines: single fly averages. Thick lines: population averages. Data from 20 cells (10 cells per hemisphere). (C) 2-6 Hz power (left panel) and canonical spike rate (right panel) as a function of raw Vm in PFNa neurons. Responses of right- and left-bridge PFNa neurons were pooled. Thin lines: single fly averages. Thick lines: population averages. Mean points in which less than 50% of the curves contributed to the bin are displayed in gray. Data from 20 cells. (D) Same as in A, but showing the baseline-subtracted Vm instead of the raw Vm. The baseline was defined as the minimum value of the heading tuning curve (see Methods). (E) Same as in B, but showing the difference in Vm pre- and post-airflow stimulus instead of the raw Vm. (F) Same as in C, but showing the baseline-subtracted Vm instead of the raw Vm. The baseline was defined as the minimum value of the heading tuning curve (see Methods). Mean points in which less than 50% of the curves contributed to the bin are displayed in gray. (G) Conjunctive tuning of left- and right-bridge PFNa cells to the direction of airflow and heading. These recordings were made from flies of the genetic background used to create the TRiP RNAi libraries ("empty RNAi control"), but where eGFP was expressed in PFNa neurons. (H) Same as in panel G, but in PFNa cells carrying the construct TRiP:HMS01948, which allows for expression (under UAS control) of a double-stranded RNA that targets Ca-α1T transcripts for degradation (Ca-α1T RNAi). (I) Same as in C, but recording from PFNa cells in flies of the empty RNAi control genotype. (J) Same as in C, but recording from PFNa cells carrying the construct TRiP:HMS01948 (Ca-α1T RNAi).
Figure S4. Features of the Vm of PFNa neurons, with and without knockdown of Ca-α1T channels. See next page for figure legend.
Figure S4. Features of the Vm of PFNa neurons, with and without knockdown of Ca-α1T channels. (A) Power spectral density (PSD) of the Vm of PFNa neurons during moments where the fly’s heading was aligned to the cell’s preferred direction (left panel) and moments where it was aligned to the null direction (right panel). Each panel shows a 60° bin of heading data (e.g. the left panel shows the PSD for the Vm of PFNa neurons when the fly’s heading was within ±30° relative to each cell’s preferred heading). The gray box indicates the 2-6 Hz band. Values from 20 cells. (B) Same as in A, but for depolarizing stimuli (airflow from 30° for right-bridge-innervating PFNa neurons and -30° for left-bridge-innervating PFNa neurons) and hyperpolarizing stimuli (airflow from -120° for right-bridge-innervating PFNa neurons and 120° for left-bridge-innervating PFNa neurons). Values from 20 cells. (C) Schematic of the current injection experiment. Each experiment consisted of a family of 15 current steps spaced 5 pA apart, starting at -40 pA and ending at 30 pA. We pooled cells innervating both brain hemispheres in this figure. (D) Representative families of current injections for 2 out of 11 different PFNa cells. This genotype is SS02255>UAS-2xeGFP, where no intentional perturbations were inserted. To the extent that it was possible, the recordings were preferentially acquired while the fly was quiescent. This was done to minimize depolarizing input from self-motion or membrane potential changes due to the visual stimulus entering or exiting the cell’s receptive field. (E) Same as in D, but in empty RNAi control genotype flies. (F) Same as in E, but in PFNa neurons expressing TRiP:HMS01948, which targets the Ca-α1T transcript (Ca-α1T RNAi). (G) 2-6 Hz power (left panel, dark yellow) and canonical spike responses (right panel, blue) as a function of membrane potential in PFNa neurons in SS02255>UAS-2xeGFP flies, where no intentional perturbations were inserted. The responses of right- and left-bridge innervating neurons were pooled. Thin lines represent individual cells, thick lines represent the average responses of 11 cells. (H) Same as in G, but in empty RNAi control genotype flies. 8 cells. (I) Same as in (H), but in PFNa cells expressing 2xeGFP and TRiP:HMS01948, which targets the Ca-α1T transcript. 10 cells. (J) Three example traces of current injection steps that yielded sodium spiking at the peak of a calcium spike. The red insets are magnified epochs highlighting the sodium spikes (blue dots). (Note that the cell bodies of Drosophila neurons connect to the cell arbors through a thin neurite that considerably filters voltage signals from the processes. Spikes in the PFNa neuron recordings are hard to visualize in raw voltage traces but are easily detected by filtering and thresholding the membrane potential.) (K) Two example traces of PFNa neurons showing calcium spiking in response to GtACR1-mediated hyperpolarization. We stimulated the fly using a 565 nm LED. The light intensity was measured to be 21 µW/mm² prior to inserting a neutral density filter in the light path to reduce the magnitude of optogenetic stimulation.
Figure S5. Analysis of the vector-axis directions and peak-vector length angles in PFNa neurons. (A) Angular indexing of the protocerebral bridge, with different schemes for the EPG neurons and the Δ7 neurons. These angles were experimentally defined in Lyu et al 2022 and we adopt them in this study. Adjacent glomeruli are generally spaced 45° apart in both cell types, but the indexing values of the Δ7 neurons are shifted to those from the EPG neurons by 1/4th of a glomerulus (11.25°). Note that the EPG neurons do not innervate the outermost two glomeruli (1 and 18); these angles were inferred given the bridge innervation pattern of the PEN (and PFN) neurons. (B) Connectivity matrix of the PFNa neurons onto the FC3 neurons, averaged over neuron instances that innervate the same bridge glomerulus or fan-shaped body column. Every fan-shaped body column innervated by the FC3 neurons receives input from both the left- and the right-bridge PFNa neurons. (C) Angle difference between left-bridge and right-bridge PFNa neuron inputs onto the FC3 neurons. Each circle corresponds to the values calculated for individual FC3 neurons, using the bridge indexing of either EPG alone, Δ7 alone, or a weighted combination. (D) Anatomically defined angle between the left and right PFNa vectors. 110.6° corresponds to the value calculated in C via connectome analysis. (E) Same as in D, but assuming pure EPG or pure Δ7 input. (F) We fit the heading-and-airflow 2-D heatmaps for spiking responses using the quadratic model in Figure 5. The best-fit angles for the preferred airflow directions are shown as the angles for each of the four vectors. (G) Overlay of the angles of the projection axes (purple) and the angles of the maximum vector lengths (green) as estimated in panels D and F.
Figure S6. Assessment of the ability of FC3 neurons to signal the allocentric direction of air puffs. See next page for figure legend.
Figure S6. Assessment of the ability of FC3 neurons to signal the allocentric direction of air puffs. (A) Connectivity matrix showing all of the identified PFN neurons in the Drosophila hemibrain connectome, and how they output onto the 10 types of columnar cell classes postsynaptic to PFNa neurons: the FC1A, FC1B, FC1C, FC1D, FC1E, FC1F, FC3, FR1, hΔC, and hΔJ cell classes. The FR1, hΔC, and hΔJ neurons receive synaptic input from additional types of PFN cells, potentially introducing confounds into our analysis of the PFNa neurons’ synaptic output; we thus chose not to study these cell classes. The FC1 and FC3 neurons, in contrast, receive essentially all of their PFN-related synaptic input from PFNa neurons. The FC1 neurons are the largest recipients of monosynaptic PFNa neuron input, but the six subtypes (FC1A-E) are challenging to parse using light microscopy. Therefore, we focused on the FC3 neurons, which have an identifiable anatomy, as our model postsynaptic neuron class. The FC3 neurons also receive heavy synaptic input from the FC1 neurons (not shown in this graph). (B) Example trace of an experiment where we simultaneously imaged the EPG and FC3 neurons. The EPG bump was imaged in the ellipsoid body, whereas the FC3 were imaged in the fan-shaped body, pictured here is the FC3 activity in layer 5 of the FB. The phase of the two population signals is overlaid in the third row, and the position of the visual stimulus on the screen is shown on the fourth row. (C) Example airflow responses of the EPG and FC3 neurons at the 12 airflow angles tested. The airflow-on period is flanked by the dotted lines, and the content and structure of the panels is otherwise the same as in B. (D) Time course of the difference of the FC3 phase position relative to the EPG phase during periods of airflow stimulation. The airflow-on period is flanked by the dotted lines, and the red dotted line denotes the expected magnitude of the phase difference if the PFNa system were operating as a four-vector set with equal amplitudes. The thin lines show individual flies, the thick lines show the average values across 7 flies. (E) Difference of the FC3 phase position relative to the EPG phase in the context of airflow stimulation like the one shown in D. The values shown here correspond to the average phase difference over the last two seconds of the airflow stimulus. The gray dots show individual flies, the black dots show the average values across 7 flies. The anatomically-defined expected bounds of the front vectors (see Supplementary Text and Figure S5) are noted and marked with dotted black lines; the expected phase differences if the front and back vectors contributed equally are noted in red.
Methods

Fly husbandry

Unless indicated otherwise, flies were reared in standard cornmeal-agar-molasses food in a 12h/12h light cycle incubator set to 25 °C. Progenies from crosses were transferred into fresh vials on the day of eclosion and housed in an incubator for 2-7 days before being affixed to a physiology platform for calcium imaging. For electrophysiological experiments, we used 4-7 day old flies.

Flies being crossed for optogenetic experiments were raised in cornmeal-agar-molasses containing vials, wrapped in aluminum foil to minimize light exposure during development. On the day of eclosion, newly hatched flies were transferred into cornmeal-agar-molasses containing vials, supplemented with 0.4 mM all-trans retinal (Sigma Aldrich). These vials were wrapped in aluminum foil for 2-5 days, until flies were affixed to a physiology platform for imaging experiments.

Fly stocks

Genotypes for each experiment are listed in Table 1. Stock sources are listed in Table 2.

We used existing Gal4 and LexA driver lines\(^{60–64}\) to target transgene expression to central-complex neurons. We used the 12E04-LexA driver line to target FC3 neurons, and we used the 27F02-LexA, 60D05-LexA, and 60D05-Gal4 lines to target EPG neurons. In addition to targeting EPG neurons, we found that the 60D05-LexA and 60D05-Gal4 lines also target unidentified cells that innervate layers 2 and 5 of the fan-shaped body, which have airflow responses (data not shown). Thus, whenever we used 60D05-Gal4 or 60D05-LexA to image EPG neurons in conjunction with either PFNa or FC3 neurons, we expressed the red calcium indicator jRGECO1a\(^{65}\) to avoid uncertainty as to the cellular origin of fluorescence signals in the fan-shaped body. In optogenetic experiments—where it was not possible for us to work with two different calcium indicators—we used 27F02-LexA to drive syt-jGCaMP8s (based on jGCaMP8s\(^{18}\) and described below) expression in EPG cells because this driver line does not target airflow-responsive fan-shaped body cells. For all other imaging experiments, we used jGCaMP7f\(^{66}\) instead.

We targeted PFNa neurons using four different driver lines: the split-Gal4 SS02255\(^{67}\), 30E10-Gal4, 41H07-Gal4, and VT056655-Gal4. From inspection of the publicly available multi-color Flip-out images by Janelia Research Campus\(^{68}\) as well as from our own immunohistochemistry data, we believe that PFNa neurons are the only PFN cells targeted in these four driver lines. We found that using 30E10-Gal4 to drive UAS-GtACR1 expression was lethal at the pupal stage, and thus we used 41H07-Gal4
and VT056655-Gal4 for the experiments in Figure 6. We targeted the LNOa neurons using split-Gal4 SS04743267.

We designed the syt-jGCaMP8s construct by linking the Drosophila synaptotagmin-1 coding sequences and jGCaMP8s (Addgene Plasmid #162380)69 using a GSGSGS linker, with the Syt1 sequence at the C terminus. We then placed this construct into pJFRC19-13xLexAop2 backbone (Addgene Plasmid #26224)60, replacing myrGFP with syt-jGCaMP8s. The plasmid was synthesized by GenScript and inserted in the VK00022 landing site (BDSC Stock #9740) using PhiC31-based integration, performed by BestGene.

10xUAS-GFlamp1 was created using an attB-site carrying plasmid gifted to us by Yulong Li’s research group70. We inserted the plasmid at the VK00005 integration site (BDSC #9725) with PhiC31-based integration, performed by BestGene.

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<td>We cold-anesthetized and mounted adult female flies to a custom stage, which allows for head-fixed behavior simultaneous with neural imaging as described previously. In brief, we attached the dorsal tip of the head and the anterior tip of the thorax to a form-fitting hole in the stage using a blue-light-activated glue (Bondic). After being thus attached, the posterior edge of the head capsule can be dissected for physiological measurements from the brain. For calcium imaging experiments that did not require optogenetic activation of neurons, we allowed the flies to recover for ≥ 2 hours under low levels of ambient light after being mounted. For experiments with imaging and optogenetics (Figure 6), we allowed the flies to recover for ≥ 2 hours inside a dark cardboard box. For electrophysiology experiments, we allowed the flies to recover for 2-4 hours inside a dark cardboard box. We used slightly different head pitch angles, depending on the brain structure which we needed to access. The angle between the front vertical drop of the thorax and back of the fly's head in experiments that involved co-imaging the ellipsoid body and the fan-shaped body was ~60° (as in refs. 1, 74). This same angle was</td>
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closer to ~45° for experiments in which we performed imaging of the protocerebral bridge and the noduli, or electrophysiology from PFNa somas.

**Extracellular saline composition and delivery**

For both imaging and electrophysiology experiments, we exposed the dorsal surface of the fly’s brain by cutting a rectangular window in the head capsule using a 30-gauge syringe (BD PrecisionGlide). We perfused the brain with an artificial extracellular saline solution bubbled with carbogen (95% CO2 / 5% O2). The composition of the saline solution, in mM, was 103 NaCl, 3 KCl, 5 N-Tris(hydroxymethyl)methyl-2-aminoethanesulfonic acid, 10 trehalose, 10 glucose, 2 sucrose, 26 NaHCO3, 1 NaH2PO4, 1.5 CaCl2, and 4 MgCl2. All chemicals were sourced from Sigma Aldrich. The solution’s osmolarity was measured to be ~280 mOsm, and after carbogen bubbling, the solution’s pH was close to 7.3. The saline was delivered to the brain using a gravity-fed perfusion system. Using a Peltier device (SC-20, Warner Instruments) regulated by a closed-loop temperature controller (CL-100, Warner Instruments), we set the saline’s temperature, measured in the bath, to 22°C for calcium imaging experiments and 25°C for electrophysiology experiments.

**Two-photon calcium imaging**

Calcium imaging data were acquired using an Ultima IV two-photon microscope (Bruker) powered with a Chameleon Ultra II Ti:Sapphire tunable laser (Coherent). In experiments where GCaMP fluorescence was imaged alone, the laser wavelength was set to 925 nm. In experiments where we simultaneously imaged GCaMP and jRGECO1a (Figures 2 and S2), the Chameleon laser wavelength was set to 1000 nm and supplemental excitation of the jRGECO1a calcium sensor was provided by a coaxial second laser set to 1070 nm (Fidelity-2, Coherent). The power of the Chameleon laser, as measured at the back aperture of the objective, was 20-50 mW for all experiments. Emitted light was collected through a 40x/0.8 NA objective (LUMPLFLN 40XW, model 1-U2M587, Olympus), split by a 575 nm dichroic mirror (575dcxr, Chroma), and collected by a pair of GaAsP photomultiplier tubes (H7422P-40, Hamamatsu). The green channel, capturing the jGCaMP7f or jGCaMP8s emission signal, was filtered through a 490-560 nm bandpass filter (525/70m-2p, Chroma). In dual imaging experiments, the red channel (containing the jRGECO1a signal) was filtered through two stacked 590-650 nm bandpass filters (620/60m-2p, Chroma). The objective was mounted on a Piezo device (525800-400, Bruker), which allowed for rapid scanning along the z axis. We imaged exclusively in galvo-galvo mode and used the Piezo device to acquire imaging volumes consisting of 3 to 6 z-planes scanned at 128x128 pixel resolution. The volumetric scanning rates ranged from 3 Hz—for experiments with a large region of
interest (ROI), such as when we co-imaged the fan-shaped body and ellipsoid body (e.g., Figures 6 and S6) — to 8 Hz when we employed very small ROIs, like when imaging the noduli (e.g., Figures 2 and S2). The dwell time for each pixel ranged from 3.6 to 4.8 µs. These settings minimized fluorophore bleaching while still providing adequate fluorescence signals.

Optogenetic stimulation

We used the two-photon laser to perform simultaneous two-photon imaging and GalACR1\textsuperscript{19} or CsChrimson\textsuperscript{17} activation (Figure 6), similar to previous approaches\textsuperscript{1,41,76}. In these experiments, the laser power was increased from 20 to 50 mW across the six imaging planes at increasing depth, following an exponential trajectory. The 20-mW plane covered the posterior end of the fan-shaped body, and the 50-mW plane covered the anterior end of the ellipsoid body. Such an exponential power increase had several advantages. First, by having the power in the fan-shaped body be relatively low, this limited photobleaching of jGCaMP8s in the fan-shaped body. Second, the low laser intensity in the fan-shaped body should help to lead to moderate, hopefully physiological, excitation levels of optogenetic reagents in PFNa terminals, rather than overactivation. Third, the high intensity in the ellipsoid body helped to increase the quality of the signal from this deep structure. We imaged volumes at ~3.5 Hz in optogenetic experiments. Approximately three planes were dedicated to imaging the fan-shaped body and approximately three planes were dedicated to imaging the ellipsoid body. Because we were optogenetically activating the terminals of PFN neurons in the fan-shaped body, we thus activated PFN cells at ~3.5 Hz and with a ~50% duty cycle.

Electrophysiology

We cold-anesthetized flies and affixed them to a custom stage as described above and previously\textsuperscript{73}. We opened a small cuticular window over the central complex using a 30-gauge syringe (BD) and removed the underlying fat and tracheal tissue, with fine forceps, to expose the brain. We illuminated the fly using an 850 nm LED (M850F2, Thorlabs) coupled to a 400 µm wide fiber optic cable (M28L01, Thorlabs) that was focused onto the fly with lenses (MAP10100100-A, Thorlabs). We visualized GFP-expressing cell bodies via standard epifluorescence, except that we used a custom GFP emission filter, which passed 510-560 nm and >800 nm (Chroma). This filter allowed us to visualize green and infrared while also cutting out the red fluorescence of Alexa-568, which we often included in our pipette for anatomical fills.

We pressure ejected 0.5% collagenase IV (Worthington) in extracellular saline from a pipette with a 4-6 µm tip positioned over the neural lamella and perineurial sheath, which weakened and ultimately
breached these layers. We raised the bath temperature to ~30°C during this desheathing process, which took < 5 min., to activate the collagenase. Following rupture of the perineurial sheath, we lowered the bath temperature to 21°C for ≥ 5 min. and increased the flow rate of the perfusion to ensure that the collagenase was fully washed out. We then performed additional manual clearing of tissue with the micropipette containing extracellular saline until individual somas of interests we exposed. We held the temperature of the bath between 24°C - 28°C for the remainder of the experiment.

We used borosilicate glass capillaries (BF150-86-7.5, Sutter Instruments), pulled on a P-1000 puller (Sutter Instruments) and fire polished with a microforge (MF2-LS2, Narishige) to produce pipettes with 6-12 MΩ resistance and ~1 μm tip openings. Pipettes were filled with intracellular saline whose composition, in mM, was 140 potassium aspartate, 1 KCl, 10 HEPES, 1 EGTA, 0.5 Na3GTP, 4 MgATP, 13 biocytin hydrazide, and 0.02 Alexa Fluor 568 hydrazide (A10437, ThermoFisher Scientific). The pH of the intracellular saline solutions was adjusted to 7.3 with KOH and its osmolarity was adjusted to 265 mOsm with water.

We visualized somas for targeting with a patch pipette using a Kinetix sCMOS camera (Teledyne Photometrics) mounted on an upright epi-fluorescence microscope (Slicescope, Scientifica) with a 40×/0.80 NA water immersion objective (LUMPLFLN 40XW, Olympus). We took care to only record from GFP expressing cells. Electrophysiological signals were amplified and low pass-filtered at 10 kHz using a MultiClamp 700B amplifier (Molecular Devices). We streamed all voltage and current signals continuously at 10kHz with a Digidata 1440A input-output board (Molecular Devices). All data presented are Vm measurements in current clamp mode. We injected a small amount of current (-1 to -3 pA) during Vm recordings, to partially address the depolarizing effect of the seal conductance on small cells, with high input resistance. Out of 41 PFNa cell recordings, across all genotypes, in which we were able to present the entirety of the airflow-direction protocol, we excluded one cell from analysis because the fly did not walk enough to allow us to estimate a heading tuning curve. We excluded two additional cells because their baseline membrane potentials were above our threshold Vm for analysis, which was ~40 mV before liquid-liquid junction potential correction, or, equivalently, ~53 mV after correction. Vm measurements reported in the paper are corrected for a ~13 mV liquid-liquid junction potential.

Closed-loop visual environment

We presented visual stimuli on a panoramic LED array spanning 270° in azimuth and 81° in height, with ~1.875° pixel resolution. The arena consisted of blue LEDs (BM-10B88MD, Betlux Electronics),
covered by 7 sheets of blue gels (Tokyo Blue, Rosco) to reduce detection of light from the display by the microscope’s photomultiplier tubes. In patch-clamp electrophysiology experiments, we used the same LED display with 6 sheets of blue gels, 3 sheets of Clear-Shield film (Less EMF A1210-24) (that were electrically grounded) and one layer of stainless steel wire cloth (McMaster-Carr 85385T89) to minimize optical reflections between opposite sides of the display. In all experiments, we typically presented on the LED arena a 6-pixel (11°) wide bright blue vertical bar against a dark background. The bar rotated in closed loop with the fly’s yaw turns on an air-supported ball, thus simulating the movements of a distant landmark that can be used for orienting. The rotations of the air-supported ball along the pitch, yaw and roll axes was detected via image analysis, using a custom-modified version of the open-source software FicTrac 2.0. We modified the FicTrac 2.0 Python code to implement it within the Robot Operating System (ROS) platform, running it at 50 Hz during electrophysiology experiments and 80 Hz during imaging experiments. We illuminated the ball using 850 nm LEDs (OSRAM Platinum DRAGON, SFH 4235) guided by optical fibers (02-535, Edmund Optics). We imaged the ball using a Chameleon3 camera (CM3-U3-13Y3M, Teledyne FLIR) with an InfiniStix lens (144100, Infinity Photo-Optical). The lens was equipped with an 850/50 bandpass filter (84-778, Edmund Optics).

**Design and construction of the airflow delivery device**

We designed an airflow delivery device based on a previously published apparatus. The previous apparatus employed a single airflow outlet pointing at the fly, which could rotate to any arbitrary angle around the yaw axis. We modified this device to ensure that nothing visual changed in the field of view of the fly when the airflow direction needed to be altered. Our goal was to make it possible to present separable visual and airflow stimuli to the fly; with the original device, whenever one would change the airflow direction, the rotating nozzle necessarily created a concomitant moving visual stimulus. To solve these potential stimuli confounds, the new apparatus in this study separated the rotating outlet from the airflow delivery part, which took the form of a static disc surrounding the fly. The rotating outlet in the new device is a circular assembly consisting of a stepper motor driving a rotating nozzle. The nozzle delivers air, whose flow rate is regulated by a mass flow controller, to a subset of 36 circularly arrayed airflow channels. Depending on the stepper motor’s position, different airflow channels are engaged. Each of these 36 airflow channels was connected to a matching airflow channel in the airflow delivery disc that surrounded the fly via flexible plastic tubing, this is the part of the device featured in the figure schematics throughout the paper. The static disc consisted of 36 tubular airflow channels, spaced evenly at 10° increments, with each channel pointing at the fly from a different angle around the yaw axis.
axis. The connecting plastic tubes attached to the outer rim of the disc were gathered using soft Velcro tape and funneled away from the fly’s field of view. In summary, a motor-controlled rotating outlet induced airflow into a distributor manifold connected to the tubes, which passed through specific airflow channels in the disc and ultimately hit the fly’s body from a specific direction around the yaw axis.

The airflow disc surrounding the fly was sufficiently thin in the vertical dimension that the fly could still see the vertical blue bar on the LED display. Thus, we could change the airflow direction and visual experience of the fly completely independently. The ~1 m length of the plastic tubes does not significantly delay an airflow pulse from hitting the fly because air pressure changes can be considered as being transmitted instantaneously from one end of the tube to the other, for our purposes. Specifically, assuming a meter-long traveling distance, a airflow pulse from the rotating outlet would take ~2.9 ms to travel across the tube, governed by the speed of sound.

We should note that, in its current incarnation, this device has two main disadvantages. First, it is not well suited for rapidly switching between different odorants, because unlike with air pressure changes, odor molecules need to travel down the entire length of each plastic tube before reaching the fly; this process would take much longer than a few milliseconds. Second, the airflow direction is discretized by the presence of the 36 static tubes, introducing variation in airflow speed depending on the phase of the rotating outlet relative to the holes in the first static ring. Specifically, we found that when we made the nozzle opening exactly the diameter of one airflow channel, the air speed arriving at the fly varied up to ~50% depending on whether the nozzle was perfectly aligned with the channel or centered on the plastic midline between two channels, for example (data not shown). To combat this effect, we made the nozzle opening span two airflow channels, i.e., we made it 20° wide, which kept the proportion of the nozzle engaged with open air (within a channel) versus plastic more consistent independent of the nozzle’s position around the ring. This design feature minimized pressure changes that were contingent on the exact angle of airflow being delivered to the fly, while still maintaining a reasonable directional accuracy for the air stream.

3D printing, part processing, and apparatus assembly

All custom parts were fabricated via 3D printing using VisiJet M3 Crystal material on a Projet MJP 3600 series 3D printer (3D Systems) at 30 μm resolution. We cleaned the printed parts using a multi-step procedure. First, the parts were incubated in an oven set to 65° C to melt and remove most of the wax support material. Afterward, the parts were sonicated in a mineral oil bath for at least one hour to dislodge the finer wax coating still attached to the plastic. The mineral oil residue was then removed by passing a jet of compressed air through all of the holes in the part. Afterwards, the parts were washed
using water and dish soap (Dawn), rinsed with distilled water, and dried with compressed air. Each of
the holes in the static airflow channels were manually tapped and fitted with a 1/16” brass hose barb
(Clippard 12841) and the barbs on both parts (the airflow disc and the discretizing manifold on top of
the rotating nozzle) were connected using soft plastic tubing (Tygon E-3603 via McMaster Carr,
5155T12). The rotating outlet component of the device was attached to an aluminum post and mounted
on the same air table as the microscope. The airflow disc was mounted to the device that held the air-
supported ball, which helped to ensure that the disc’s center was precisely located at the position in
which the fly stood. Additionally, the airflow disc featured notches that nested the fly plates and ensured
correct alignment of the fly’s body relative to the airflow disc during each experiment.

Airflow calibration

We calibrated the device by measuring the air speed at the center of the airflow disc with a hot wire
anemometer (Climomaster Anemometer 6501-CE, Kanomax) fitted with an omnidirectional spherical
probe (6543-2G, Kanomax). Using custom components, especially printed for alignment purposes, we
positioned the spherical probe in the precise position that a fly would occupy during an experiment. We
then manually rotated the nozzle so as to align it with the frontmost (0°) airflow channel. At this zero
position, the 20° nozzle thus fully spanned the central airflow channel as well as ~5° each of the
channels on either side of the frontmost channel. The airflow angles used in this paper were all
multiples of 10° from this zero position, meaning that for all airflow stimuli we expect there to have been
the same phase relationship between the nozzle opening and the airflow channels downstream of the
nozzle. This fact makes it more likely that the air speed that the fly experienced was consistent across
different delivery angles (because we noted changes in air speed that could occurred as a function of
the phase relationship of the nozzle and the downstream, discrete, set of tubes). Indeed, we
experimentally verified that the air speed was consistent at the position of the fly, with the exact air puff
angles used in the paper, using the anemometer. We also validated the directionality of the air puffs
delivered to the fly by visualizing CO₂ gas (dry ice) with a laser sheet. We performed the dry ice
visualization in a separate, but nominally identical, assembly.

Simultaneous control of the airflow and bar stimuli

We used analog voltages—generated by the microcontroller that controlled the visual pattern on the
LED display⁷⁷—to control the angular position of the airflow stimulus as well as the mass flow controller
that turned the airflow on and off. To control the angle of the airflow stimulus, the analog voltage
controlled the position of a stepper motor that drove the rotating air outlet to new set points. To control
the air speed, a voltage signal controlled the opening and closing of 2-SLPM mass flow controller (Alicat Scientific). The approach of having the LED-arena’s microcontroller also drive the airflow stimuli made onset and offset latencies of the air puffs more reliable and also aided temporal alignment between the visual and air-puff stimuli.

Experimental protocol for presenting open-loop airflow and closed-loop visual stimuli

To solidify the visually mapping between the position of the blue bar on the LED display and the EPG bump position in the brain, all electrophysiology experiments and most imaging experiments began with a period of 5-10 minutes of the fly interacting with the bright blue bar in closed loop. This procedure was particularly important for electrophysiological data collection because we could not directly estimate the EPG phase in these experiments, and thus any changes in the mapping between the EPG phase and the visual cue would have manifested as multi-peaked heading tuning curves from single neurons. We could be less strict on this matter in two-photon imaging experiments because we were directly monitoring the EPG neuron population and could always estimate its phase in the brain, independent of the bar’s position on the LED arena. Following the acclimation period for establishing a robust EPG visual mapping, we began to present air puffs in open loop. We presented air puffs from twelve angles around the fly, in $30^\circ$ increments, to fully and equally cover $360^\circ$ of azimuthal space. The $0^\circ$ direction, where air arrived from directly in front of the fly, was always included, which anchored the eleven other angles presented. Each open-loop air pulse lasted for 4 s with a 5 s inter-pulse interval. The air-puff angles were pseudorandomized in each block of twelve trials. We collected 3-5 blocks of trials per fly during imaging experiments, and 10 blocks during electrophysiology experiments. We used an airflow speed of 20 cm/s for all puffs. In this study, $0^\circ$ represents the direction directly in front of the fly, and $180^\circ$ represents the direction directly behind the fly. Negative angles represent air arriving to the fly’s egocentric left side, and positive angles other than $0^\circ$ (front) and $180^\circ$ (back) are to the egocentric right of the fly.

Synchronized acquisition of behavior and physiology data

We recorded all experiment-related signals in the form of voltages using a Digidata 1440A I/O board and Axoscope software (Molecular Devices). The air-supported ball’s yaw, pitch, and roll angles, the control signals to and from the airflow stepper motor, the Alicat flow meter output, the angular position of the blue bar the LED display, the internal triggers of the FicTrac ball-tracking camera, and custom signals to identify trials epochs, were recorded at 10 kHz in all experiments. Signals specific to either calcium imaging or electrophysiology, such as the membrane potential, injected current, or triggers for
two-photon imaging frame acquisition, were recorded in addition when applicable. Behavioral, electrophysiological, and calcium imaging signals updated at different rates and we used either the FicTrac camera’s triggers, or the two-photon imaging frame triggers, for careful temporal alignments to behavioral and brain-imaging signals, as required.

**Calcium imaging data analysis**

*Data processing*

We registered raw time series of fluorescence images using either the rigid motion correction algorithm in the CaImAn software suite\(^1\) or a custom algorithm described previously\(^7,8\). Both registration approaches produced qualitatively similar results. When available, we used the tdTomato or jRGECO1a image for registration, otherwise, we used the jGCCaMP7f signal directly. After registration, we drew regions of interest (ROIs) manually, in each z-plane separately, using custom software\(^4\) written in Python 3.8. We drew ROIs on fluorescence images averaged across an entire recording session, and our assignment of glomerular boundaries was aided by simultaneously viewing autocorrelation images using data from the entire session. When analyzing data in the protocerebral bridge, individual glomerulus identities were assigned by adhering to the following established anatomical principles\(^8\): (1) the entire protocerebral bridge should be composed of 18 distinct glomeruli, (2) PFN neurons do not innervate the pair of glomeruli closest to the midline, and (3) EPG neurons do not innervate the pair of glomeruli furthest from the midline. Out of a total of 12 collected bridge co-imaging datasets, we did not analyze 4 recordings in which more than one glomerulus was not visible, or whose identity could not be clearly established. When analyzing the ellipsoid body, we divided the structure into 16 evenly spaced radial sectors, whose boundaries and center were defined across every plane, as previously described\(^1,7,8\). Following previous conventions\(^1,7,8\), and to establish a consistent anatomical correspondence with the bridge and fan-shaped body, wedge 1 and wedge 16 of the ellipsoid body were defined, respectively, as the wedges immediately to the left and immediately to the right of the vertical bisector line at the bottom of the torus, when viewed from the posterior side of the head. When analyzing the fan-shaped body, we defined columns as follows. We first drew an outline of the entire structure. Then, the left and right edges of the structure were marked with angled lines. The vertex point at which the extension of these two angled lines met defined the entire angular extent of the fan-shaped body, which we then subdivided into 16 equally-spaced columns\(^1\). Column 1 of the fan-shaped body, which corresponds functionally to wedge 1 of the ellipsoid body, was defined as the leftmost fan-shaped body column when viewing the structure from the posterior side of the head. Noduli outlines were manually drawn and assigned as either left or right, using the same posterior view.
Fluorescence signal normalization

For each imaging volume and structure being imaged, we averaged together the signal from ROIs across z-planes, if they belonged to the same glomerulus, wedge, column or nodulus side, thus generating a final, unidimensional array of fluorescence intensity values for each structure at each time point. Each of these unidimensional arrays, corresponding to individual sector ROIs, were concatenated into a matrix where each data column corresponded to one sector and each row corresponded to a time point. Thus, the matrices of fluorescence values were composed of 18 data columns for the 18 glomeruli in the protocerebral bridge, 16 data columns for the 16 wedges of the ellipsoid body, 16 data columns for the 16 anatomical columns of the fan-shaped body, and 2 data columns for the 2 sides of the noduli. We report all imaging data from the ellipsoid body and fan-shaped body as \( \Delta F/F \), calculated as \( (F - F_{min}) / (F_{min}) \). In this equation, \( F \) corresponds to the raw fluorescence intensity measured in a given sector ROI (i.e. in each wedge or column) and \( F_{min} \) corresponds to the 5th percentile of \( F \) values observed in that same sector ROI over the whole recording. When analyzing calcium imaging data from the protocerebral bridge, we normalized each glomerulus ROI to its own maximum and minimum \( F \) values because some glomeruli were much brighter or dimmer than others likely due to the expression levels of the fluorophore. Thus, the values of \( \Delta F/F_{max}-F_0 \) that we report here for the protocerebral bridge correspond to \( (F - F_{min}) / (F_{max} - F_{min}) \), where \( F \) corresponds to the raw fluorescence signal in sector ROI, and \( F_{min} \) and \( F_{max} \) represent the 5th and 95th percentile of the \( F \) values from that same sector ROI over the entire recording. When analyzing the data from the noduli, we performed a z-score normalization over the raw fluorescence values, calculated separately for each of the left and right nodulus sector ROIs.

Aligning fluorescence readings with behavioral measurements

We matched behavioral (80 Hz) and imaging (4-8 Hz) measurements in the following manner. We determined the epochs corresponding to entire volumetric scan cycles using the voltage triggers output by the two-photon microscope, and we averaged the fluorescence values corresponding to each ROI across all z-slices containing them in the cycle. We averaged behavioral-related measurements, such as the visual stimulus’ position on the screen, over the time corresponding to an entire volumetric scan cycle as well. The behavioral signals are thus averaged over epochs of ~125-250 ms (corresponding to the imaging period given our scanning rates of 4-8 Hz).
Phase extraction

Columnar neurons in the central complex often express spatially localized calcium signals that are referred to as bumps of activity. Because space around the fly, 0°-360°, physically maps to positions in central complex structures, the position of the calcium bumps within a structure has an angular interpretation and thus is often referred to as the bump’s phase. We extracted the phase of calcium bumps in the ellipsoid body and fan-shaped body by taking a population vector average, as described previously\textsuperscript{8,78,83}. We defined the sectors of the ellipsoid body and fan-shaped body to have their numerical values (1 to 16) match the known anatomical and functional correspondence across these two structures. Thus, when phase of bumps in these two structures matched, this meant that the angular signal carried by the two neuronal populations were functionally aligned in angular space, and phase differences across the ellipsoid body and fan-shaped body could be calculated through a simple angular subtraction as described in the Data processing subsection above. When imaging EPG neurons in the protocerebral bridge, the phase of the EPG heading estimate was extracted as previously described\textsuperscript{78}. In brief, we took a Fourier transform of the sector ROI signal over the 16 glomeruli. The phase of the EPG bump in a given time point was then defined as the phase of the Fourier component with a wavelength of 8 glomeruli. This approach assumes that an invariable 8-glomerulus spacing exists between the peaks of the two EPG bumps in the bridge, and that, together, these two bumps encode a single, shared, heading angle estimate, which has proven to be a robust assumption.

Phase nulling

We phase-aligned EPG and PFN bumps in the protocerebral bridge using an algorithm described previously\textsuperscript{78}. In brief, the sector ROI arrays from the protocerebral bridge for EPG and PFNa cells were first, separately, interpolated to 1/10th of a glomerulus resolution using a cubic spline. We then circularly rotated the vector at every time point so that the EPG phase was at the same position along the x axis. This EPG-determined rotation was then applied to the PFNa interpolated vector signal, allowing us to visualize the phase of the PFNa bumps in reference to that in the EPG system. It is only by employing this phase nulling analysis that we analyzed the phase of the PFNa bumps presented in this study. We found that we could not easily define the phase of the PFNa bumps otherwise, because they were not stably periodic nor always visible in calcium experiments.
Electrophysiology data analysis

Data processing

We processed current-clamp recordings to extract each cell’s membrane potential (Vm), sodium-spike rate and 2-6 Hz power (due to calcium spiking). To make analysis of Vm convenient, we downsampled the raw 10kHz Vm traces (corrected for the liquid-liquid junction potential) to 1000 Hz. Afterwards, we median-filtered those traces using a kernel size of 40 ms to remove spikes. We detected sodium spike times by first bandpass filtering the raw 10kHz Vm traces in the range of 150-1000 Hz using an 8-pole Bessel filter. In this way, we selected for higher frequency events such as sodium spikes which are characterized by fast changes in the Vm, and we selected against lower frequency events such as subthreshold synaptic potentials. Second, we identified discrete events in the bandpass filtered trace based on crossing of a high threshold. Finally, we rejected all events with an inter-event interval of less than 6.66 ms. We labeled all events detected this way as sodium spikes, and visually inspected a subset of the events to ensure appropriate spike detection. We converted sodium-spike times to spike rate by convolving spike times with a sliding 1-second-wide Hann window. Sodium-spike amplitudes were often small—only a few millivolts or less—in these recordings, which is typical of recordings from somata of neurons in invertebrates. As a result, any spike-detection algorithm used is likely to have resulted in calling some non-spike events as spikes and to not have detect some number of genuine sodium spikes emitted by the cells. We do not view a potential, low rate of Type I and Type II errors as a substantive concern, however, for two reasons. First, our results depend on an overall estimate of spike rate and not precise spike times; some low level of spike misdetections might raise or lower the estimate rate over time, but this should not drastically impact the shape of the relevant tuning curves for our model. Second, empirically, the spike detection algorithm that we employed detected putative sodium spikes only extremely rarely when PFNa cells were hyperpolarized (e.g. Figures 3F-3G), arguing that we were not typically detecting subthreshold events, erroneously, as sodium action potentials.

We quantified calcium spikes by extracting membrane potential oscillation strength in the 2-6 Hz range. To extract oscillation strength, we downsampled raw Vm traces to 100 Hz and performed a fast Fourier transform on the traces (specgram function in Matplotlib; window size = 4 s; window step size = 20 ms). We averaged the calculated power of Vm frequencies in the range of 2-6 Hz and used this value as our metric for estimating the strength of T-type oscillations in PFNa neurons. To quantify the oscillation strength during 2 s periods of current injection, we first conditioned the current-injection Vm traces as follows. We defined two 600-ms time segments. One segment started 550 ms before the start of the current injection pulse and the second segment started 50 ms before the end of the current
injection pulse. We then replaced the actual \( V_m \) sample points in both segments with a linear interpolation between the first and last sample point within the time segment. We then took the fast Fourier transform of the \( V_m \) traces and quantified the power in the 2-6 Hz band, using a window size of 2 s and window step size of 0.01 s. The linear interpolation of the \( V_m \) trace that we performed at the onset and offset of the current pulse prevented artifactual frequencies from appearing to have high power in the Fourier transform analysis simply due to the large \( V_m \) step-like changes that necessarily occur at the start and end of the current injection pulse.

To align behavioral readings with electrophysiological measurements, the sample points from the FicTrac camera were upsampled to 1000 Hz, using linear interpolation, to match the analyzed membrane potential signal, which was downsampled to 1000 Hz as described above.

**Estimation of baseline \( V_m \)**

PFN\( a \) neurons express large calcium spikes, alongside large synaptic potentials, even when very hyperpolarized, which makes defining a resting \( V_m \) difficult for these cells. Thus, instead of estimating a resting \( V_m \) for PFNa cells, we instead defined the baseline \( V_m \) as the minimal value in the heading tuning curve calculated for each cell (mean: -64 mV, standard deviation: 4.8 mV, range: 19.9 mV). This baseline \( V_m \) value was typically stable during our recordings, which usually lasted 45 to 60 minutes. Whereas the baseline membrane potential varied substantively across cells (Figures S3A and S3B), we found that the peak-to-minimum membrane potential amplitude of the heading tuning curve (mean: 13.7 mV, standard deviation: 2.9 mV), and the sodium spike rate at the minimum value of the heading tuning curve (mean: 0 spikes/s, standard deviation: 0.1 spikes/s) was more consistent across cells. Given this observation, membrane potential data was combined across cells by subtracting the baseline membrane potential from each cell. We compare tuning curves with and without normalization in Figure S3. Spike rate data were combined across cells without any such normalization.

**Estimation of heading, airflow direction and conjunctive tuning curves**

For each PFNa cell, we generated \( V_m \), spike rate, and oscillation strength tuning curves as a function of heading and airflow angle. We generated single-variable tuning curves against these two variables, and we also generated conjunctive, two-variable (heat map) tuning curves against the two variables. For heading tuning curves, we binned the time series of bar positions on the LED display into 10° bins, and we averaged the \( V_m \), spike rate or oscillation strength in each bin. We required that a bin included at least 2 s worth of sample points for its average to be calculated. We also required that the fly not be
standing (forward speed > 0.5 mm/s) for sample points to be included in the tuning curve. We calculated the preferred heading direction of a given cell’s tuning curve by finding the angular shift of the curve that produced the highest correlation between the actual tuning curve and a normalized cosine function. We calculated tuning curves to air puffs by averaging the $V_m$, spike rate, or oscillation strength in the 10 air-puff trials for each of 12 air puff directions presented in a 2 s window, starting 0.5 s after the onset of the air puff. To estimate conjunctive tuning to heading and airflow, we split the data into 18 (20° heading) x 12 (45° airflow direction) bins. We required that at least 0.5 s of data populate each bin in order to calculate a mean value. After calculating the conjunctive tuning heat-map for each cell separately, we generated combined heat maps across cells after centering each single-cell heat map on the preferred heading direction of a given cell.

Model construction

Membrane potential data over 12 egocentric airflow directions, $W$, and 18 relative heading directions, $H$ (the heading direction relative to the preferred heading direction for a particular neuron), averaged over the recorded PFNa left and right neurons, were fit using the form $V_m = a_o + a_r \cos(H) + a_w \cos(W \pm \pi/4)$, with free parameters $a_o$, $a_r$, and $a_w$, and the ± applied to left or right neurons, respectively. This 3-parameter fit of 216 data points for each case (left/right) explained 93.7% (left) and 90.9% (right) of the variance of the data.

Firing rates for sodium spikes were fit to the equation $r = ([b_r \cos(H) + b_w \cos(W \pm \pi/4)]_+)^2$, with $b_r$ and $b_w$ free parameters and $[x]_+ = x$ for $x > 0$ and 0 otherwise. This two-parameter fit of 216 data points (as above) explained 85.9% and 85.7% of the variance for the left/right PFNa neurons, respectively. We fit the oscillation power for these data to $p = c_o + ([c_r \cos(H) + c_w \cos(W \pm \pi/4)]_-)^2$, with $c_o$, $c_w$, and $c_r$ free parameters and $[x]_- = x$ for $x < 0$ and 0 otherwise. This three-parameter fit of 216 data points explained 71.4% and 77.1% of the variance for the left/right PFNa neurons.

For the mathematical model of the full response discussed in the text, we assumed that the plus and minus rectifications discussed in the previous paragraph could be combined, when modeling the output of a PFNa neuron across all angles. This means that we assumed that the proportionality constant relating output due to calcium spikes to output due to sodium spikes is such that the total output can be expressed as $(b_r \cos(H) + b_w \cos(W \pm \pi/4))^2$, with no rectification as a consequence of summing sodium and calcium spike contribution. Summing the left and right contributions and
accounting for the ±45° anatomical shifts of the PFNa projections from the protocerebral bridge to the fan-shaped body gives the total signal from this similarly tuned pair as

\[
\left( b_h \cos \left( H - \frac{\pi}{4} \right) + b_w \cos \left( W + \frac{\pi}{4} \right) \right)^2 + \left( b_h \cos \left( H + \frac{\pi}{4} \right) + b_w \cos \left( W - \frac{\pi}{4} \right) \right)^2
\]

\begin{align*}
= & \ b_h^2 \left( \cos \left( H - \frac{\pi}{4} \right)^2 + \cos \left( H + \frac{\pi}{4} \right)^2 \right) + b_w^2 \left( \cos \left( W + \frac{\pi}{4} \right)^2 + \cos \left( W - \frac{\pi}{4} \right)^2 \right) \\
& + 2b_h b_w \left( \cos \left( H - \frac{\pi}{4} \right) \cos \left( W + \frac{\pi}{4} \right) + \cos \left( H + \frac{\pi}{4} \right) \cos \left( W - \frac{\pi}{4} \right) \right).
\end{align*}

Because \( \cos(x - \pi/4) = \sin(x + \pi/4) \), and the sum of squared sines and cosines is 1, the first line on the right of the equal sign above equals \( b_h^2 + b_w^2 \). Using the sum-of-angles identity, \( \cos(H \pm \pi/4) \cos(W \pm \pi/4) = (\cos(W + H) + \cos(W - H \pm \pi/2))/2 \), and noting that \( \cos(x + \pi/2) + \cos(x - \pi/2) = 0 \), the last expression above becomes \( 2b_h b_w \cos(W + H) \). Putting this all together, we find that the total output of this pair of PFNa neurons is \( b_h^2 + b_w^2 + 2b_h b_w \cos(W + H) \), as given in the text. The output across the full population of PFNa neurons, parameterized by their preferred heading angle \( \theta \) is then a phasor representing a vector of unit length pointing in the allocentric direction of the airflow.

**Immunohistochemistry**

We dissected adult fly brains and fixed them in 2% paraformaldehyde for 55 minutes in 24-well crystallization plates (Cryschem M Plate, Hampton Research). We washed the brains 3x for 20 minutes each in phosphate-buffered saline containing 0.5% Triton-X (PBST), then blocked them with 5% normal goat serum (NGS, sourced from Gibco). For immunostaining, we used a primary antibody solution consisting of 1:1000 chicken anti-GFP (600-901-215, Rockland Immunochemicals), 1:30 mouse anti-bruchpilot (nc82, Developmental Studies Hybridoma Bank), and 5% NGS diluted in PBST. We nutated the brain samples in primary antibody solution for 4 h at room temperature, followed by an overnight rocking incubation at 4°C. Afterwards, we washed the brains 3x for 20 minutes each in PBST, then incubated them in a secondary antibody solution, composed of 1:800 goat anti-chicken Alexa Fluor 488 (A11039, ThermoFisher Scientific), 1:200 goat anti-mouse Alexa Fluor 594 (A11032, ThermoFisher Scientific), and 5% NGS in PBST. We nutated the brains for 4 h at room temperature, then incubated them overnight at 4°C. Afterwards, we washed the brains 3x for 20 minutes each in PBST, performed a final wash in phosphate-buffered saline, and mounted the brains onto glass slides in 8 µl of FocusClear (CelExplorer) with the posterior side up (i.e. the posterior end of the brain faced the coverslip). We imaged the mounted brains using an upright Zeiss LSM 780 confocal microscope fitted with a 20x 0.8NA air objective (Plan-Apochromat 20x/0.8, Zeiss). Each stack of images consisted of 80-120 optical
sections spaced ~1 µm apart.

**Analysis of open-source RNA sequencing dataset**

The data in Figure 4B were obtained from an open-source RNA sequencing dataset by Davis et al, 2020\(^{14}\), available in the NIH’s Gene Expression Omnibus (GEO)\(^{84,85}\) under accession number GSE116969. The specific dataset shown here is GSE116969_dataTable4.genes_x_cells_TPM.coding_genes_QCpass, which reports the cell type-mean abundance of protein coding genes in quality control-passed samples. No additional processing was performed on the reported abundances.

**Analysis of open-source hemibrain connectome dataset**

All electron microscopy-based analysis in this study was performed using the hemibrain connectome\(^{59}\) dataset v1.2.1 and neuPrint-python\(^{86}\) v0.4.15. Skeleton renderings were performed using the NAVis python library v1.3.1.

**Statistics**

Details on statistical testing, as well as exact p-values, are presented below in Table 3. Throughout the paper, we used a circular mean function implemented after Fisher and Lee, 1983\(^{87}\). We performed t-tests using the scipy.stats package\(^{88}\) and we performed Watson-Williams tests using the pycircstat package, a Python implementation of CircStat\(^{89}\), a toolbox for directional statistics in MATLAB.

<table>
<thead>
<tr>
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<td>S4H, S4I</td>
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<td>Samples have equal means</td>
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Code and data availability

Analysis code will be released on GitHub at the time of final publication of this manuscript. Until such point, data analysis scripts will be made available upon request.

Acknowledgements

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Author contributions

I.G.I. and G.M. conceived of the project. I.G.I. performed the imaging and optogenetic experiments and the associated analyses. S.S. performed the electrophysiological experiments and the associated analyses. I.G.I., S.S., L.F.A. and G.M. jointly interpreted the data and decided on new experiments. L.F.A. developed and implemented the formal models. T.L.M. and I.G.I. developed the airflow device. I.G.I., L.F.A., and G.M. wrote the paper with input from S.S. and T.L.M.
Supplementary Text

When we fit our model to $V_m$, spiking, and oscillation data, we used fixed angle offsets of ±45° for the airflow tuning curves. However, we also fit these data allowing these offset angles to be free parameters. When we did this for the $V_m$ data, we obtained angles of -41.9° and 41.3°. Although these angles differ from the ±45° assumed in our model, the improvement in the fits obtained by introducing these extra free parameters was small, only 0.02% additional variance explained for the left-bridge and right-bridge PFNa neurons. Allowing the angles to be free parameters for fitting the spiking and oscillation data gave angles of -40.0°, 39.3°, -60.4°, and -51.9° (Figure S5F) for the left-bridge and right-bridge spiking data and left-bridge and right-bridge oscillation data, respectively. This allowed the fits to explain an additional 0.8%, 1.0%, 7.3% and 1.2% of the variance in the data. These angles are close to, but slightly offset from, the ±45° expected from a perfectly orthogonal system.

The deviations from orthogonality suggested by the above analysis make quantitative predictions about the nature of heading tuning in the PFNa neurons if a vector sum process aims to accurately signal the allocentric airflow direction. For example, if a PFNa cell were to express its maximal airflow response at +40°—i.e., 5° closer to the fly’s midline than the +45° orthogonal prediction—its response to heading in the fan-shaped body should be shifted by +50° (relative to its response in the protocerebral bridge), i.e., 5° further from the midline, for the vector sum process to be accurate. Put another way, if a PFNa-encoded vector expresses its maximal vector length when airflow arrives from +40°, then the axis that this vector points along should be +50° for the vector sum process to be accurate. With non-orthogonal axes for a pair of basis vectors, the angle of each axis and the airflow angle at which each basis vector has its maximal projection to that axis are no longer the same angle.

To determine the angular shift in the PFNa projection from the bridge to the fan-shaped body—which determines the axis along which each PFNa vector points—we used the hemibrain connectome. In the bridge, we assigned an angle to each synapse that a PFNa cell receives from an EPG or $\Delta 7$ cell, based on standard assumptions of how heading is signaled in the EPG/$\Delta 7$ system (Methods, Figure S5A). We averaged all the synaptic angles onto a given PFNa cell to assign each PFNa cell an overall heading angle for which it codes. We then analyzed the pattern of PFNa synapses onto a specific recipient cell class in the fan-shaped body, FC3 neurons. We found that the mean angular deviation between the left- and right-bridge PFNa cells that co-innervate FC3 cells in a given fan-shaped body column is 90.5° if EPG cells were considered to be the only drivers of PFNa activity and 113° if $\Delta 7$ cells were considered to be the only drivers (Figures S5A, S5B, S5C, and S5E). Because EPG and $\Delta 7$ cells express 11% and 89% of the synapses onto PFNa cells, respectively, their circular weighted average predicts a deviation of 110.6° between the left and right PFNa vectors as
they influence a given FC3 column (Figures S5C and S5D). This anatomical calculation thus argues that the projection axes for the two PFNa vectors are offset by 55.4° to the left and right of the midline (Figure S5D). As mentioned, the fact that this angle is bigger than 45° is expected given that the peak vector lengths, assessed with physiology, were measured to be closer to the midline than 45°. Specifically, our physiological measurements of 39° and 40° predicted projection axes of ±50-51°, which are in reasonable alignment with the 55° estimate extracted from the connectome. The fact that the FC3 phase deviated by no more than ±55° from the EPG phase in response to air puffs (Figure S6E) is consistent with the possibility of the anatomical projection axes of the two PFNa vectors being separated by ~110°, as the connectome analysis suggests. This offset angle would limit the maximal deviation of any downstream bump that sums the two front vectors.

Together, our angular measurements across the anatomy and physiology of the PFNa neurons suggest that its vector system could function in a non-orthogonal manner (Figure S5G). However, given the rather modest improvements provided by the fitted angular variables mentioned in the first paragraph above, we chose to use the canonical angles of ±45° for our model.