Data-driven analysis of motor activity implicates 5-HT2A neurons in backward locomotion of larval Drosophila

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Rhythmic animal behaviors are regulated in part by neural circuits called the central pattern generators (CPGs). Classifying neural population activities correlated with body movements and identifying the associated component neurons are critical steps in understanding CPGs. Previous methods that classify neural dynamics obtained by dimension reduction algorithms often require manual optimization which could be laborious and preparation-specific. Here, we present a simpler and more flexible method that is based on the pre-trained convolutional neural network model VGG-16 and unsupervised learning, and successfully classifies the fictive motor patterns in Drosophila larvae under various imaging conditions. We also used voxel-wise correlation mapping to identify neurons associated with motor patterns. By applying these methods to neurons targeted by 5-HT2A-GAL4, which we generated by the CRISPR/Cas9-system, we identified two classes of interneurons, termed Seta and Leta, which are specifically active during backward but not forward fictive locomotion. Optogenetic activation of Seta and Leta neurons increased backward locomotion. Conversely, thermogenetic inhibition of 5-HT2A-GAL4 neurons or application of a 5-HT2 antagonist decreased backward locomotion induced by noxious light stimuli. This study establishes an accelerated pipeline for activity profiling and cell identification in larval Drosophila and implicates the serotonergic system in the modulation of backward locomotion.

The neural circuits generating rhythmic behaviors such as walking and breathing are called the central pattern generators (CPGs)1-3. Since rhythmic behaviors of invertebrates and vertebrates share many features, studies on the CPGs in one animal species are expected to serve as a model for those in other species4. Identification of neurons involved in CPGs is the first important step in understanding how the rhythmic behavior is generated and regulated by the neural circuits. Such analyses have often been performed in the isolated central nervous system (CNS) since it is known that CPGs can produce fictive motor patterns that resemble the actual behavior patterns without any sensory feedback5-7.

Recent advances in imaging technology such as spinning-disc and light-sheet microscopy enabled recording of neural activity in large regions in the brain, paving new ways to investigating CPG circuits. In animals with relatively small CNS such as C.elegans, larval Drosophila and larval zebrafish, it is now possible to image the entire brain or even the whole animal in real time8-10. While the technological advances are now enabling one to record the activity of a majority of neurons in the nervous system in these animals, it remains challenging to extract useful information from the large data-sets obtained by the recording. In the case of CPG studies, for instance, one may want to determine the time windows in which specific motor activity takes place and then to identify the neurons that show activity related to the initiation, duration, and termination of the motor pattern. Previous studies used methods such as principal component analysis (PCA), independent component analysis (ICA), singular-value decomposition (SVD) and k-means clustering to quantitatively classify neural activity patterns8,11,12. Although these methods are powerful and have been used widely, they often require manual

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optimizations which are labor-intensive and confined to specific datasets. Therefore, development of alternative methods which can handle various types of activity data more efficiently is greatly anticipated.

The larval *Drosophila* is one of the most powerful model systems for studying neural circuits related to rhythmic behaviors since its CNS is numerically simple (containing ~10,000 neurons) and amenable to various genetic manipulations. Especially, imaging fictive motor patterns in the isolated CNS with genetically encoded Ca2⁺ indicators is well established. An isolated CNS can generate fictive motor outputs such as coordinated propagation of motor activity along the body axis, which resembles forward and backward locomotion of the animal, and left-right asymmetric bursts in anterior neuromeres which likely correspond to turning. Whole-animal functional imaging in embryos just before hatching confirmed that the propagating activity and asymmetric bursts occur during forward/backward locomotion and turning, respectively. An isolated CNS also generates symmetric and synchronous bursting activity in the anterior-most and posterior-most segments, which often but not always occur just prior to the initiation of backward and forward fictive locomotion, respectively. While corresponding larval motor outputs of the bursting activity is not clear, bursts in posterior-most segments may be related to movement of the gut and tail which is known to occur prior to forward locomotion. Previous studies have shown that subsets of interneurons show activity correlated with these fictive motor outputs and play roles in the regulation of larval movements, such as segmental activity propagation, left-right symmetric coordination and differential recruitment of motor pools.

In this study, we present a new methodology for classifying neural activity patterns in larval *Drosophila* that utilizes a convolutional neural network (CNN) and unsupervised learning. This method successfully classified forward and backward waves as well as synchronous activities in anterior- and posterior-most neuromeres from large activity data derived from different sub-populations of central neurons. We also identified cells associated with the classified motor activity patterns by voxel-wise correlation mapping. We then applied this method to a population of neurons targeted by 5-HT2A-GAL4, which we generated by CRISPR/Cas9-mediated gene knock-in, and successfully identified two classes of interneurons that are specifically active during fictive backward locomotion. Further functional analyses implicated these 5-HT2A neurons and 5-HT neurons in the regulation of backward locomotion. Our study provides an accelerated pipeline that efficiently identifies neurons correlated with motor patterns from wide-volumetric functional imaging data.

**Results**

**Classification of fictive motor patterns by deep convolutional feature and unsupervised clustering.** In this study, we aimed to identify motor patterns automatically in a data-driven manner. The key idea was to create x-t images in which x-axis represents the activity of multiple ROIs at each time point. Since the motor activity pattern in the x-t image is similar to images such as handwritten digits, we expected that CNN, known to give superior performance in image classification tasks, may be used to efficiently categorize the motor patterns. We, therefore, tested if a CNN model, VGG-16 pre-trained on the ImageNet dataset, which is widely used in transfer learning, can classify neural activities from the calcium imaging data. We used *Rra-GAL4* to express the genetically encoded calcium indicator GCaMP6f in aCC and RP2 motoneurons (MNs) in third instar larvae and imaged fluorescence changes associated with neuronal activity in the isolated CNS using spinning-disc confocal microscopy. As previously reported, the MNs showed left-right symmetric wave-like activity patterns reflecting forward and backward locomotion (hereafter called FW and BW, respectively) and bilateral asymmetric activities associated with turning. We used neural activity data from 18 regions of interest (ROIs) corresponding to both sides of the neuromeres T2-A7 in 12 animals for the analyses (Fig. 1a). Since our initial goal was to identify motor patterns corresponding to FW and BW, which are bilaterally symmetric, we initially combined the activity data in the left and right side and extracted activity patterns associated with propagating waves, and only later used the information from both sides to study the symmetry of the motor patterns identified. Hence, we compressed the 18-dimension (18-d) time series to 9-d time series using the maximum value of each segment (Figs 1b and 2a) and analyzed the data using the CNN model (Figs 1c and 1a).

We searched for the layer which gives the best match for FW and BW by cluster analyses of extracted features from each layer of the CNN model with hierarchical agglomerative clustering (HAC, Ward’s method) (Fig. 5b,c). By consulting the original activity pattern, we manually determined the clusters representing FW and BW, and found that the intermediate Conv4_3 layer gives the best performance in classifying the motor patterns from the neural activity data (Figs 1c,d and S1). In addition to activities reflecting FW and BW, the Conv4_3 layer also succeeded in identifying two additional motor-related activities described previously. One is left-right symmetric or asymmetric activation in anterior neuromeres (hereafter called AT). The other is synchronous and bilateral activation in posterior-most neuromeres (called PT) and is known to often occur before forward waves. It was difficult to correlate these activities to motor patterns in intact larvae. While some of the AT activities likely correspond to turning, we were unable to clearly distinguish among turning-like and other AT activities. Furthermore, the classification performance decreased when asymmetric activities were taken into consideration (i.e., using 18 ROIs). From these reasons, we decided to focus on classifying activity patterns without considering asymmetric parameters in this study and took the maximum value in each segment so that we can detect activities occurring in either side of the animal. We evaluated the efficacy of the identification by comparing with the ground truth data obtained manually (~10% of dataset) and found that our methodology yields on average, 75–100% correctly classified hits and 0–6.7% false alarms for each of the four motor patterns, FW, BW, AT and PT (Figs 2a–d and S1d). It is noteworthy that the CNN layer followed by HAC analyses succeeded in extracting all four major motor-related activities identified by human inspection (Fig. 1d). In the following analyses, we also annotated the time windows with no activity as a quiescent state (QS) and those showing activities other than the four motor patterns as unlabeled (UL).

We characterized the frequency and bilateral symmetry of the four motor patterns identified (Fig. 2c). The frequency of bursting activities in anterior and posterior neuromeres (AT and PT) and wave-like activity (FW
Intensities of neuronal activation were found to be higher during BW and FW compared to AT and PT. We also confirmed that AT but not other motor patterns shows strongly asymmetric activity as previously reported\(^1\). It has been known that transition between different motor patterns in behaving larvae is biased\(^3\).\(^4\). We, therefore, asked whether the transition probabilities of fictive behavior patterns are also biased and found it is indeed the case (Fig. 2d). Strongest transition probabilities were seen in the transition from BW to AT and that from FW to PT (Fig. 2d, bottom). Since the transition probabilities from AT to BW and from PT to FW (directly or via QS) were also high, the probability of the occurrence of the same motor pattern was high after BW and FW (i.e., BW-AT-BW, FW-PT-FW) (Fig. 2d, top). While direct transition between AT and PT frequently occurred (0.15–0.25 transition probabilities), direct transition between BW and FW did not occur. These tendencies (e.g., AT occurs before BW) were also observed in the average image of each activity (Fig. 2e). This is consistent with the fact that larvae often perform forward or backward locomotion consecutively. Thus, our methodology successfully classifies motor patterns and allows efficient quantification of a large-scale activity data without laborious human labeling processes.

Voxel-wise mapping of whole neurons. In the above experiment, GCaMP6f was expressed in a small number of identified MNs in each segment. We next asked whether the semi-automated classification method of fictive motor patterns can be applied to imaging data from a larger number of cells and, if so, whether we can use the classification method to identify previously uncharacterized interneurons that show activity pattern(s) correlated with specific fictive motor pattern(s). For this purpose, we first used a pan-neuronal GAL4 driver, elav-GAL4, to express GCaMP6f in all neurons. As with the above experiment, we analyzed the activity extracted from 18 ROIs representing all hemisegments in three larvae and succeeded in extracting the four motor patterns (Figs 3a,b and S2a). Thus, our methodology is also applicable to pan-neuronal activities. We next tried to identify voxels correlated with each of the fictive motor patterns by using Pearson’s correlation coefficient (Pearsons’ r, Fig. 3c-c”). Since some voxels showed correlation with multiple motor patterns, we also determined the motor pattern which showed the highest correlation with the signal in each voxel (Dominant motor pattern map, Fig. S1b–d). Voxels correlated with AT (>0.1 Pearson’s r) were found in large quantities in anterior neuromeres (T1 to A3, Fig. 3c), whereas those correlated with BW were seen in a complementary manner in posterior neuromeres (A4 to A7, Fig. 3c”). AT was often seen prior to BW and these two motor patterns together
appear to correspond to the backward locomotion of the larvae. Similarly, voxels correlated with FW were largely found in middle neuromeres (A1 to A5, Fig. 3c''), whereas those correlated with PT were seen in a complementary manner in posterior neuromeres (A6 to A8, Fig. 3c'''). PT was often seen prior to FW and these two motor patterns together likely correspond to the backward locomotion of the larvae. These voxels were seen in similar manners across individuals (Fig. S2b–d) and likely correspond to segmentally repeated neurons showing propagating activity (FW and BW) or bursting activities spanning several segments (AT and PT). We also identified voxels that were separated from the segmentally clustered ones described above and showed unique morphologies and thus likely belong to the same cell(s). These included structures in a dorso-medial region in anterior neuromeres which showed correlation with BW (Fig. 3c'', arrow) and a pair of longitudinal axons in dorso-lateral neuropile, likely those of ascending/descending neurons, which showed correlated activity with PT (Fig. 3c'''', arrow). By placing a ROI in each of these structures (Fig. 3c'',c'''), we confirmed that they are active specifically in BW or PT (Fig. 3d).

Voxel-wise mapping of putative 5-HT2A-receptor expressing cells identified Seta and Leta as backward-specific neurons. The correlation mapping of motor patterns described above enabled us to
identify neuronal activities specifically associated with each of the four motor patterns. However, since all neurons expressed GCaMP6f, it was difficult to identify the cells that showed these activities. To overcome these limitations, we next tried to use GAL4 lines that drive expression in subsets of neurons. In particular, we were interested in neurons expressing neuromodulator (NM) receptors because of their potential roles in behavioral control. We therefore used the CRISPR/Cas9 method to generate new GAL4 lines that recapitulate the expression pattern of each NM receptor. In these lines, the 2A-GAL4 gene was inserted in frame with each NM receptor immediately in front of the stop codon, allowing GAL4 to be co-expressed with the NM receptor in the same spatiotemporal pattern (Fig. 4a). We generated GAL4 knock-in lines for 12 NM receptors (serotonin-receptors, 5-HT1A, 5-HT1B, 5-HT2A, 5-HT2B, 5-HT7; octopamine-receptors, Octβ2R, Octβ3R, OamB; tyramine-receptor, TyrR2; dopamine-receptors, Dop1R1, Dop2R, DopEcR) and studied the expression of each GAL4 line in larvae using UAS-mCD8-GFP as a reporter. Except for 5-HT2A-GAL4, all of the NM receptor GAL4 lines drove GFP expression in large subsets of neurons in the larval CNS (Fig. S3). In contrast, 5-HT2A-GAL4 drove GFP expression in small subsets of neurons which can be individually identified (Fig. 4b). We performed calcium imaging of the cells expressing 5-HT2A-GAL4, analyzed the data in a data-driven manner as described above (as a reference for motor activity, we also expressed GCaMP6f in aCC and RP2 MNs) and generated a dominant motor pattern map (Fig. 4c,c'). By this, we identified voxels which were strongly active during BW but not FW locomotion (Figs 4c" and S4d). As will be described below, we found these voxels belong to two pairs of interneurons in the

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**Figure 3.** Motor pattern extraction from activity data of whole neurons. (a) An example of extracted motor patterns (bottom) from the activity data derived from pan-neuronally expressed GCaMP6f (top, grayscale image) (b) Mean of fictive motor pattern derived from 3 samples aligned with the end point of each label. Average duration of each motor pattern, AT: 1.8 ± 0.1 s, BW: 2.4 ± 0.1 s, FW: 1.7 ± 0.1 s, PT: 1.6 ± 0.1 s, QS: 2.3 ± 0.4 s (mean ± standard deviation). Note that unnormalized ΔF/F is shown since normalization obscured the motor pattern. (c) Motor pattern correlation mapping. Voxels with >0.1 Pearson’s r for each motor pattern, AT (c’), BW (c’), FW (c”) and PT (c’’), are shown in colors. c shows the 18 ROIs used for the analyses (visual cues). Image baselines are derived from maximum GCaMP signal. (d) Neural activity in the structures identified by the correlation mapping. The average activity in the structures identified as specific to BW (top) and PT (middle) during each motor pattern (bottom). Scale bar 100 μm.
Figure 4. Identification of Seta and Leta, backward-specific interneurons expressing 5-HT2A-GAL4. (a) A cartoon showing the design of GAL4 knock-in. The GAL4-coding sequence along with the 2A peptides was inserted by the CRISPR/Cas9 method to the target NM receptor gene locus, just downstream of the C-terminus of the NM coding sequence. dsRed flanked by loxP sites was used as a marker for successful insertion. (b) Expression of 5-HT2A-GAL4 visualized with mCD8-GFP. (c–c”) Motor pattern mapping in 5-HT2A-GAL4- and RRa-GAL4-targeted cells. (c) 18 ROIs representing each hemisegment used for the analyses (visual cues). (c’) Dominant motor pattern mapping identified voxels specific to BW. Motor patterns are colored as in Fig. 3. Two ROIs (labeled as Seta and Leta) are selected for the ROI correlation mapping in (c’). (c”) ROI correlation mapping reveals the outline of Seta and Leta. Pearson’s r between each voxel and a ROI in Seta and Leta shown in (c) was calculated. The mapping results for Seta (red) and Leta (green) and GCaMP signals (blue) are merged. Axons of Seta and Leta are revealed in yellow (red + blue) and green/blue color (green + blue) and are found to be present in the CI and DM tracts (Fas2 coordinate) respectively (arrows). (d) Mean activity of Seta (top, red) and Leta (middle, green) (derived of three samples) aligned with motor activities during AT, BW, FW and PT (bottom, grayscale image). Colored bars represent the duration of each motor pattern. (AT:3.6±0.2 s, BW:5.2±0.2 s, FW:3.4±0.4 s, PT:3.0±0.1 s, QS:8.4±0.9 s, mean±standard deviation). (e) Maximum activity level (fluorescence changes, ΔF/F) of Seta and Leta during each motor patterns. While both Seta and Leta are active during BW, only Seta is active during AT and PT. ***p < 0.001,
ANOVA with post-hoc Tukey HSD test, compared to QS. (f’-f”) ROI correlation mapping in 5-HT2A-GAL4-targeted cells. Unlike in c, ROI mapping was applied only to 5-HT2A-GAL4-targeted cells. The morphology of Seta (f”f”) and Leta (f’f”) can be seen more clearly. Scale bar 100 μm.

VNC, named as Seta (Small-η) and Leta (Large-η) after their characteristic morphologies (see below). Although Seta and Leta neurons were both active during BW (Fig. 4d), temporal profiles of the activity of these neurons differed: while Seta showed a single peak of activity during BW, Leta showed double peaks. Furthermore, Seta but not Leta showed weak activity during AT and PT (Fig. 4e).

We next studied the morphology of Seta and Leta neurons. Since neurites of 5-HT2A-GAL4-positive neurons extensively overlap, it was impossible to determine the morphology of individual neurons when markers such as mCD8-GFP were expressed in all GAL4-positive cells. We therefore first tried to reveal the morphology of Seta and Leta based on their activity by calculating the Pearson’s r. As shown in Figs 4c”f’-f” and S4, this method identified voxels which are synchronously active and thus likely belong to each of the neurons, revealing the outline of their morphology. We then searched for these neurons in cells individually labeled by single-color and multi-color flip-out techniques32 and studied their morphology more in detail. The Seta neuron was found to be located in a ventro-medial region of T1 neuromere and extended its axon and dendrites in the dorsal neuropile (Fig. 5a,e). The axon of the neuron crossed the midline and then extended posteriorly along the CI tract (Fas2-coordinate)30 to A1 neuromere. The dendrites extended posteriorly near the DM tract to T3 in the ipsilateral side. The Leta neuron was located in a medio-lateral region of T3 neuromere and extended its axon and dendrites in the dorsal neuropile (Fig. 5b,d). The axon of Leta neurons also crossed the midline and then bifurcated to extend posteriorly toward A6 neuromere and anteriorly toward T1 segment (Fig. 5b,d). The dendrites of Leta neurons innervated a medial region of the dorsal neuropil in T1–A3 neuromeres. Both Seta and Leta neurons were weakly stained with anti-Choline acetyltransferase (ChAT) antibodies (Fig. 5e,f). As will be described below, Seta and Leta also expressed other markers of cholinergic neurons, Cha-GAL80 and Cha.3.3-GAL80. Thus, these neurons are likely cholinergic.

Optogenetic activation of Seta and Leta neurons increases backward locomotion. Since Seta and Leta are specifically active during backward but not forward fictive locomotion (Fig. 4d,e), they are potentially involved in the induction of backward locomotion (Fig. 6a). We therefore next asked if forced activation of these neurons is sufficient to induce backward peristalsis. We first studied the effects of activating all 5-HT2A-GAL4-positive cells during locomotion. We expressed a red-shifted channelrhodopsin, CsChrimson44 and optogenetically activated these neurons in freely moving larvae. Light application significantly increased backward peristalsis (Fig. 6b), indicating that activation of the 5-HT2A neurons promotes backward locomotion. To narrow down candidate neurons responsible for the optogenetic induction of backward locomotion, we used intersectional methods. tsh-GAL80 and cha.3.3-GAL80 or cha-GAL80 suppress the function of GAL4 in a majority of neurons in the VNC and cholinergic neurons, respectively. Accordingly, expression of CsChrimson driven by 5-HT2A-GAL4 was abolished in Seta and Leta neurons when tsh-GAL80, cha.3.3-GAL80 or cha-GAL80 was combined with the GAL4 (Fig. S5). In these larvae, optogenetic induction of backward locomotion was significantly reduced (Fig. 6b). These results implicate cholinergic 5-HT2A neurons in the VNC, including Seta and Leta, in the induction of backward locomotion. To further narrow down the expression of CsChrimson, we next used the flip-out technique to generate clones of cells expressing CsChrimson among the 5-HT2A-GAL4-positive cells. We optimized the heat-shock condition so that less than ten neurons in the VNC expressed CsChrimson. We stimulated larvae with light and recorded the behavioral responses of 180 larvae upon optogenetic stimulation. The larvae were then dissected and the neurons expressing CsChrimson were identified in individual larvae. We observed a strong correlation between the expression of CsChrimson in Seta or Leta neurons and induction of backward locomotion (Fig. 6c–e). In contrast, such correlation was not seen for other 5-HT2A neurons in the VNC (Fig. S6). These results indicate that activation of Seta or Leta promotes backward locomotion in the larvae.

Larval serotonergic system is required for backward locomotion. We next tested if 5-HT2A-GAL4 neurons including Seta and Leta are required for the induction of backward peristalsis. We induced backward locomotion in free moving larvae by shining blue light, which is an aversive stimulus36. Thermogenically inhibiting 5-HT2A-GAL4 cells using temperature-sensitive Shibire36 significantly decreased the induction of backward peristalsis (Fig. 7a) compared to control larvae, indicating that 5-HT2A cells are required for proper induction of backward locomotion. Since 5-HT2A expressing neurons are likely to be in part regulated by 5-HT, we next asked if 5-HT releasing neurons are also involved in the induction of backward locomotion. We used the same light-induced behavioral assay and found that thermogenetic inhibition of 5-HT neurons with Shibire36 (whose expression driven by trh-GAL4, which specifically targets 5-HT neurons)37 significantly reduced the rate of backward locomotion (Fig. 7b). These results suggest the possibility that 5-HT neurons induce backward locomotion by modulating 5-HT2A neurons, including Seta and Leta, via the 5-HT2A receptor. To test this, we next studied the effect of Ketanserin, a specific antagonist for the 5-HT2 receptor38 on fictive backward locomotion. A recent study reported that Ketanserin has no effect on 5-HT2B39, the only other 5-HT2 receptor in Drosophila, suggesting that the drug is potentially specific to 5-HT2A receptors. We used our methodology described above to quantify the occurrence of backward locomotion before and after the addition of Ketanserin. Frequency of motor patterns related to backward locomotion, AT and BW, were significantly reduced upon application of the drug (Figs 7c and see S7a,b for details). In contrast, there was no change in the frequency of FW. These results render further support for the notion that 5-HT modulates backward locomotion via the 5-HT2A receptor and neurons. We also studied a loss-of-function mutation of 5-HT2A (5-HT2A50)40 but found no effect on backward
locomotion (Fig. 7b). This could be due to the fact that the mutation is partial loss-of-function\(^40\) and/or compensations during development. Finally, we studied the activity of 5-HT neurons during fictive locomotion. Since they are involved in the regulation of backward locomotion, they may be selectively active during the relevant motor patterns. We expressed GCaMP6f in 5-HT neurons using \(trh\)-GAL4 as a driver and studied their activity during fictive locomotion in isolated CNSs (Fig. 7d). We again used our methodology to classify motor patterns seen in the activity of 5-HT cells. As when GCaMP6f was expressed in motor neurons or all neurons, the four motor patterns were extracted (Fig. 7e,f). This indicates that 5-HT neurons in the VNC show activities related to all fictive motor patterns, including the segmental propagation corresponding to forward and backward locomotion. However, the intensity of the activity was much higher during motor patterns related to backward locomotion, AT and BW, compared to those related to forward locomotion, PT and FW (Fig. 7g). These results suggest that 5-HT is released preferentially during backward locomotion.

Discussion
Classifying activity patterns is crucial in studying large-scale calcium imaging data. Previous classifying methods often require manual optimization thereby making data analysis laborious and less flexible. Recently, rapid development in CNN has produced remarkable results in various fields including biomedical research\(^41\). Given the fact that human can perceive the neuronal activity pattern as images resembling patterns of MNIST handwritten digits\(^44\), we thought it would be possible to distinguish neuronal activity patterns using a CNN pre-trained by image classification. As expected, we found that motor activity patterns could be classified by the deep convolutional feature extracted from Conv4_3 of VGG-16 model and HAC. The VGG-16 model\(^28\) was used since it has a relatively simple architecture among the contemporary CNN models\(^25,42\) and has been broadly used for transfer learning. Since manual optimization is minimized, our method is less laborious compared to previous methods and could easily be adopted to imaging data obtained under various imaging conditions. Our method is also more objective since the unsupervised clustering reduces human interpretation. Furthermore, while our method was only applied to the larval preparation in this study, similar strategies would be useful for classifying activity data in other systems. Although our method was only performed as offline processing to fit HAC on the data and label its clusters, it may be performed as online processing since HAC can predict motor patterns in unseen data once it is fitted during the offline processing. Moreover, the performance of our method may be improved by introducing...
supervised learning such as fully connected neural networks and random forest classifier (transfer learning). Our methodology will greatly accelerate future analyses of fictive locomotion in *Drosophila* larvae.

The methodology successfully extracted all of the four major motor patterns, FW, BW, AT and PT, which have previously been detected by human inspection in an isolated nerve cord13. Furthermore, it did so in activity data of different cells: two pairs of MNs in each neuromere (aCC and RP2, Fig. 2), all neurons (Fig. 3) and small subsets of interneurons (5-HT neurons, Fig. 7), the data of which were acquired under various time resolution (180–420 ms per volume, Figs 2–4 and 7). It also works with small sample size (3–12 samples, Figs 2–4 and 7). Thus, our method appears to be robust and works with various numbers and types of neurons and under different imaging conditions.

We analyzed transition probabilities between the four motor patterns and found that they are biased as observed in intact larvae. This is consistent with that the behavioral decision of the intact larva is biased30,31,43 and suggests that the bias partly derives from the intrinsic properties of the CPGs. However, while free moving larvae on a substrate rarely show backward locomotion, backward waves occur frequently in the isolated CNS. The Markov chain found in the isolated CNS may be more closely related to larval behaviors within the food44 or when sensory input is genetically removed45, where backward locomotion is more frequently seen.

We also used correlation mapping to identify voxels (Figs 3a–c, 4c,d and 7e) and cells (Fig. 4d,e) showing correlated activity with each of the motor patterns classified by VGG-16 and HAC. Taken together, these methodologies establish a pipeline that efficiently extracts motor patterns and the associated cells from the calcium imaging data. We first applied the pipeline to activity data derived from all neurons (*elav>*GCaMP6f) and succeeded in identifying cellular structures which are preferentially active during PT and BW. Those correlated with PT are longitudinal axons that run across the VNC (Fig. 3c, e). Since the voxel-wise correlation mapping was limited by imaging resolution and therefore did not allow detection of the cell body, it is currently unknown whether the axons belong to ascending or descending neurons. Those correlated with BW are neurites spanning a few anterior neuromeres. To our knowledge, neurons with such morphologies have not been identified to be associated with PT or BW. Future identification of sparse GAL4 lines targeting these neurons will enable functional analyses of the roles of these neurons in motor regulation.

Although we could identify some cellular structures specific to PT and BW, most of the voxels which were preferentially active for each of the motor patterns were intermingled with each other and thus were difficult to be

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**Figure 6.** Optogenetic activation of the Seta or Leta neuron increases backward peristalsis. (a) Montage of a representative backward peristalsis of the 3rd instar larva. Scale bar 1 mm. (b) Optogenetic activation of 5-HT2A-GAL4 targeted cells with or without *ish*-GAL80, *cha3.3-GAL80* or *cha-GAL80*. Quantification of the number of backward peristalses (left, boxplot with scatter) and backward score (right, violinplot with mean and SEM) in each genotype is shown. n = 10, 9, 22, 11, 8, 13, 9, 10 (from top to bottom). *p < 0.05, **p < 0.01, ***p < 0.001, ANOVA with post-hoc Tukey HSD test. (c–e) Optogenetic activation of small subsets of 5-HT2A-GAL4 targeted cells. (c) Quantification of backward locomotion as in b in larvae with or without CsChrimson expression in Seta or Leta. *p < 0.05, **p < 0.01, Mann-Whitney U test. p = 0.016 (waves), 0.010 (score) for Seta (n = 13 (+), 167 (−)); p = 0.015 (waves), 0.015 (score) for Leta (n = 9 (+), 171 (−)); p = 0.005 (waves), 0.005 (score) for Seta or/and Leta (n = 19 (+), 161 (−)). (d,e) Examples of CsChrimson expression. A larva with expression in a Seta neuron (d) and Leta neuron (d). The number of backward peristalsis induced and backward score is also shown. Scale bar 100 μm.
individually identified. We therefore next applied the pipeline to neurons targeted by sparse GAL4 lines. As candidates, we focused on interneurons that express NM receptors since NMs are known to regulate various animal behaviors such as locomotion, foraging, feeding, aggression and courtship. Since antibodies are not available, previous studies used GAL4 lines (either enhancer-trap insertions or driven by cis-elements) which are expected to mimic the endogenous gene regulation to study the expression of some of the NM-receptors. However, for most of the NM-receptors, GAL4 lines have not been available and their expression pattern was therefore completely unknown. In this study, we systematically generated GAL4 lines targeting most of the NM receptors in Drosophila using CRISPR/Cas9-mediated gene knock-in. Since GAL4 was inserted in frame with the endogenous proteins, the GAL4 lines are expected to reproduce the expression of the NM-receptors in a more precise manner than previously reported GAL4 lines. We found that most of the NM-receptors, GAL4 lines have not been available and their expression pattern was therefore completely unknown. In this study, we systematically generated GAL4 lines targeting most of the NM receptors in Drosophila using CRISPR/Cas9-mediated gene knock-in. Since GAL4 was inserted in frame with the endogenous proteins, the GAL4 lines are expected to reproduce the expression of the NM-receptors in a more precise manner than previously reported GAL4 lines. We found that most of the NM-receptor GAL4 lines drive expression in large subsets of neurons in the CNS, sensory neurons and muscles (Fig. S3). However, one of the lines 5-HT2A-GAL4 showed sparse expression and application of the pipeline to this GAL4 line allowed us to identify two interneurons termed Seta and Leta correlated with backward locomotion. Subsequent optogenetic experiments showed that activation of these neurons increases the frequency of fictive backward locomotion (Fig. 6).

Furthermore, thermogenetic experiments showed that backward locomotion of the larvae induced by aversive stimuli is reduced when 5-HT2A-GAL4-targeted neurons are inhibited (Fig. 7a). The involvement of 5-HT2A neurons in the regulation of backward locomotion was further supported by the effects of Ketanserin, antagonist putatively specific to 5-HT2A receptor, on fictive backward locomotion (Fig. 7b). While these loss-of-function
experiments do not specifically target Seta and Leta, the results of optogenetic activation and the fact that these are the only cells we detected as active during BW strongly suggest that these neurons are required for proper execution of backward locomotion.

We also studied the role of neurons expressing the ligand of the 5-HT2A receptor, Serotonin (5-HT). 5-HT is a well-known NM that plays essential roles in the regulation or modulation of animal behaviors. For instance, 5-HT has been implicated in the regulation of escape behavior and crawling in the mollusk *Tritonia*<sup>50</sup>, dwelling, slowing and transition from crawling to swimming in the nematode *Caenorhabditis elegans*<sup>51–53</sup> and nociception and turning in larval *Drosophila*<sup>35,54</sup>. We found that thermogenetic inhibition of 5-HT neurons reduced the frequency of backward locomotion in the larvae, as when 5-HT2A cells are inhibited. We also showed that 5-HT neurons are more strongly activated during backward fictive locomotion than during forward locomotion. Furthermore, effects of Ketanserin indicate that 5-HT regulate backward locomotion in part via 5-HT2A cells. Taken together, our results strongly suggest that 5-HT modulates backward locomotion by acting on Seta and Leta neurons via the 5-HT2A receptor.

It is clear that analysis of neuronal activity at the system level is crucial for understanding the circuit underpinnings of CPGs<sup>54,55</sup>. Accordingly, improved methodologies and pipelines that link large-scale activity data to motor patterns and associated component neurons would greatly facilitate future studies of CPGs. The methodology and pipeline we developed in this study are based on open source and can be easily handled. They allow rapid and robust classification of motor patterns under various imaging conditions. We showed the utility of the pipeline by applying it to 5-HT2A-GAL4 neurons and successfully identifying interneurons involved in the regulation of backward locomotion. In the future, application of the pipeline to other GAL4 lines would enable systematic search of interneurons involved in motor regulation. This work lays the foundation for the quantitative analysis of neuronal activity in larval *Drosophila* and shows how state-of-art machine learning technique can be applied to biological research.

**Methods**

**Fly strains.** The following fly strains were used: yw, 10xUAS-IVS-mCD8::GFP<sup>56</sup>, UAS-GCaMP6f<sup>57</sup>, cha-GAL80<sup>58</sup>, cha3-GAL80<sup>58</sup>, RRA-GAL4<sup>58</sup>, tsh-GAL80<sup>59</sup>, UAS-CsChrimson-mVenus<sup>34</sup>, UAS-Shibire<sup>36</sup>, elav-GAL4<sup>40</sup>, UAS-FRT-stop-FRT-CsChrimson-mVenus<sup>20</sup>, UAS-MCGF042<sup>94</sup>, w<sup>1</sup>; P(FRT(w<sup>1</sup>)2AP(neoFRT)82B PBac( GAL4D,EYFP)5-HT2A<sup>YFP</sup>)<sup>20,00052</sup> (Bloomington Drosophila Stock Center (BDSC) stock No. 19367) was used as an allele of 5-HT2A<sup>5-HT2A</sup> (FRT<sup>10,00038</sup>)<sup>90</sup>, w<sup>1,118</sup>; P(FRT(w<sup>1</sup>))2AP( neoFRT)82B (BDSC stock No. 8218) was used as a control for the 5-HT2A mutant. Flies were raised on conventional cornmeal agar medium at 25 °C except for those expressing Shibire<sup>46</sup> which were raised at 18 °C.

**Generation of NM receptor GAL4 lines.** GAL4 knock-ins were generated by targeted transgene insertion using the CRISPR/Cas9 system (Kondo and Ueda 2013). A transgene encoding the 2A peptide and the GAL4 transcription factor was inserted in front of the stop codon of each NM receptor gene. The translated product, which is a fusion protein of the NM receptor, the 2A peptide and GAL4, is cleaved at the 2A peptide so that both the NM receptor and GAL4 are expressed in the cell. Lines were generated for the following genes: CG9652 (Dop1R1), CG33517 (Dop2R), CG18314 (DopEcR), CG15113 (5-HT1B), CG1056 (5-HT2A), CG2796 (5-HT2B), CG120732 (5-HT7), CG33976 (Octh2R), CG42244 (Octh3R), CG3856 (Oamb), CG16766 (TyrRII).

**Optogenetic activation.** Hatched larvae were raised on regular food at 25 °C until second instar (~72 hours after egg laying; hAEL) and transferred to an agar plate with yeast paste containing 1 mM All-trans retinal 24 hours before optogenetic experiments. All optogenetic experiments were done at 25 °C under dim light. Third instar larvae (~96 hAEL) were washed in deionized water to remove residual yeast paste and acclimated on an agar plate before optogenetic experiments. Larval locomotion on the agar plate was recorded using a stereo-microscope (SZX16 or MVX10; Olympus) and a CCD camera (XCD-V60, Sony) at 15 frames per second. Free-moving larvae were stimulated with red light (660 nm, LED, Thorlabs) at 1 mW/mm<sup>2</sup> for 20 seconds four times with 20 seconds interval. Number of backward peristalses was counted manually. We quantified backward locomotion in two ways: “Backward wave” is total number of backward peristalsis in four stimulations, “backward score” is defined as 0 (no backward peristalsis) or 0.25 (one and more than one backward peristalsis) per stimulus so that the maximum score is 1 for four stimuli in one trial. Backward waves and backward score measure how strong the phenotype is and how constant it is, respectively. We used progeny larvae from the cross between yw and a UAS effector line as a control.

**Stochastic optogenetic activation of 5-HT2A-GAL4 targeted neurons using Flp-out technique.** Female 20XUAS -> dFRT -> CsChrimson::mVenus (attP18), pBPhsFlp2::P:ext (attP3) flies were crossed with 5-HT2A-GAL4 males. Eggs were collected for 24 hours on an apple-juice agar plate with yeast paste. The eggs were raised at 25 °C for 24 hours. Then, heat shock (37 °C) was applied for 45 min to express Flippase. After that, the larvae were transferred to another plate, on which 1 mM All-trans- Retinal-containing yeast paste was spread, and raised at 25 °C for 48–72 hours. Optogenetic experiments were done as described above. 180 animals were tested, and the expression patterns of CsChrimson::mVenus in all larvae were examined by immunostaining. Based on the morphology, we labeled a subset of the neurons: Seta, Leta, TLN (Thoracic Long-axon Neuron), TUN (Thoracic U-shaped Neuron), and TBN (Thoracic Bump Neuron) (Fig. S6). We selected these neurons for the following reasons: located in thoracic neuromeres, not segmentally repeated and easily to identify by their characteristic morphology.

**Temperature shift experiments.** Parental flies and eggs were raised at 22 °C for Shibire<sup>46</sup> experiments. For blocking neural activity, larvae were put on an agar plate held at 32 °C on a heat plate (Thermo plate, Tokai...
Δ min-max transform (Δ indicates how much calcium signal is rising. The value of F/F of ROIs from aCC motor neurons becomes extremely high for unknown reasons. For image conversion, anti-mouse Cy5, goat anti-rabbit Alexa Fluor 488 (Life Technologies). Signaling Technology), mouse anti-ChAT (4B1, 1:5, DSHB). The following secondary antibodies were used: goat (Af2020, 1:1000, Frontier Institute), mouse anti-Fas2 (1D4, 1:10, DSHB), rabbit anti-HA (C29F4, 1:1500, Cell Technology). The larval CNS was placed dorsal-side up on a MAS-coated slide glass (cell adhesive slide glass, Hit, Japan). To induce photo-avoidance backward locomotion, we stimulated free-moving larvae with blue light (488 nm, mercury lamp) at 0.5 mW/mm². Locomotion of larvae was recorded and analyzed as described in the optogenetic activation section. Progeny larvae from yw a crossed between UAS effector line and trh-GAL4 or 5-HT2A-GAL4 were used as a control for 5-HT2A > Shibireα or trh > Shibireα, respectively.

**Immunohistochemistry.** Wandering third instar larvae were dissected in phosphate buffered saline (PBS) and fixed with 4% formaldehyde in PBS for 30 min at room temperature. Fixed larvae were washed with 0.2% Triton X-100 in PBS (hereafter PBT) and blocked with 5% normal goat serum (NGS). Blocked larvae were incubated at 4°C with the primary antibodies over 12 hours. The larvae were then washed with PBT and incubated at 4°C with the secondary antibodies over 12 hours. We used confocal scanning microscope (Fluoview FV1000, Olympus) with a water immersion objective 20x and 60x lenses for fluorescence imaging. The frame size of the images varied from 1024 × 768 pixels to 1600 × 1200 pixels. The images were processed and analyzed using Fiji (ImageJ) and Imaris (Bitplane) software. The following primary antibodies were used: rabbit anti-GFP (AF2020, 1:1000, Frontier Institute), mouse anti-Fas2 (1D4, 1:10, DSHB), rabbit anti-HA (C29F4, 1:1500, Cell Signaling Technology), mouse anti-ChAT (4B1, 1:5, DSHB). The following secondary antibodies were used: goat anti-mouse Cy5, goat anti-rabbit Alexa Fluor 488 (Life Technologies).

**Calcium Imaging.** We selected wandering third instar larvae (96–120hAEL) and isolated the CNS from the bodywall. The larval CNS was placed dorsal-side up on a MAS-coated slide glass (cell adhesive slide glass, Matsunami glass, JAPAN) and covered with TES buffer (TES 5 mM, NaCl 135 mM, KCl 5 mM, CaCl₂ 2 mM, MgCl₂ 4 mM, Sucrose 36 mM). Imaging was performed with a spinning-disk confocal unit (CSU21, Yokogawa) and EMCCD camera (iXon, Andor) installed on Axioskop2 FS microscope (Zeiss). 20x water-immersion objective lens was used. Imaging conditions for each genotype are described in Table 1.

**Preprocessing for feature extraction.** Changes in fluorescence intensity (F(t)) within 18 ROIs, which roughly correspond to hemineuromeres T2 to A7 at both sides, were measured. For each time point t (=0, 1, 2… and each ROI, we defined the baseline (F₈norm(t)) as the mean value of F(t – 16) to F(t – 1). Considering that action potential occurs when the GCaMP6f signal is rising8, and to minimize the effects of fluorescence degeneration, changes in fluorescence (ΔF/F) were defined as (F(t)/F₈norm(t) – 1) * H(F(t)/F₈norm(t) – 1) where H(x) is the Heaviside function so that the value of F(t)/F₈norm(t) less than 0 was treated as 0. Therefore, ΔF/F we defined indicates how much calcium signal is rising. The value of ΔF/F of ROIs from aCC motor neurons (R Ra > GCaMP6f; RRa, 5-HT2A > GCaMP6f) more than 2 was treated as 2 for value fitting because it occasionally becomes extremely high for unknown reasons. For image conversion, ΔF/F is scaled to between 0 to 1 using min-max transform (ΔF₈norm) and multiplied by 255 and then converted to 8 bit integer. 18 ROIs are compressed to 9 ROIs using the maximum value among the values of left and right side and the 9-dimensional values were used for the following analyses.

**Feature extraction and clustering.** We defined window size as 8 time frames (from t – 4 to t + 3) so that each window has 9 ΔF₈norm of ROIs and 8 time frames. Each window was transformed to a 72 pixel (resized from 9 ROIs) × 72 pixel (resized from 8 time frames) × 3 channel image where all channels have the same values. The deep convolutional features of the images were extracted from intermediate layer (Conv4_3) of ImageNet pre-trained VGG-16 model28. The features were compressed using global average pooling and clustered by Ward’s method. Table 2. An example of data flow of sample with 2048 time frames.
hierarchical agglomerative clustering (Ward's method, Table 2). Labeling of the clusters was performed by comparing average images of the first 20–30 clusters. The images showing high intensity in T2–A4 and A4–A7 are categorized as AT and PT, respectively. The images showing intensity flow from the posterior to anterior and from the anterior to posterior are categorized as FW and BW, respectively. The images showing nearly no intensity change are categorized as QS. We counted consecutive frames of the same label as one activity. The codes and datasets with ground truth used in Figs 1 and 2 are available (https://github.com/jeonghyuk/behavior-class.git).

**Correlation mapping.** Registration of images was performed by ImageJ (NIH) and the Template matching plugin. For $RRa > GCaMP6f, 5-HT2A, RRe > GCaMP6f and trh > GCaMP6f$, z stacked images were used for registration over time. For $elav > GCaMP6f$, each z plane was used for registration. Changes in fluorescence ($ΔF/F$) were defined as described above. We defined a binary label vector as (1 for label, 0 for not label). For voxel-wise behavior correlation mapping (behavior mapping), Pearson’s $r$ of $ΔF/F$ of voxels and the binary label vector were calculated. For voxel-wise ROI correlation mapping (ROI mapping), Pearson’s $r$ of $ΔF/F$ of voxels and $ΔF/F$ of ROIs were calculated. F(t) of a ROI is derived from the sum of F(t) of voxels whose Pearson’s correlation coefficient is 0.05 or higher.

**Bath application.** Ketanserin (+)-tartrate salt (D006-10MG) was purchased from MERCK and stored at 4 °C. Ketanserin (10 mM concentration) was solved in DMSO just before the experiment and stored at 4 °C. Experiments were performed with final concentration of Ketanserin in the bath to be 100 μM. At this concentration, the effects of 5-HT on larval heart rate is known to be completely blocked.[6]

**Statistical analysis.** We used Mann-Whitney $U$ test and one-way analysis of variance (ANOVA) followed by Tukey's HSD test for multiple comparisons. Statistical significance is denoted by asterisks: $***p < 0.001; **p < 0.01; *p < 0.05; n.s., not significant. All statistical tests were performed using Python. The results are stated as mean ± s.d. or mean ± SEM.

**Data availability.** The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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**Author Contributions**

J.P., H.K. and A.N. conceived and designed the experiments. J.P. performed the experiments and analyzed the data. S.K., H.T. and H.K. contributed reagents. J.P., H.K. and A.N. wrote the main manuscript. All authors reviewed the manuscript.

**Additional Information**

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