Fast tuning of posture control by visual feedback underlies gaze stabilization in walking *Drosophila*

**Highlights**
- Exploratory walking in *Drosophila* is configured to maximize gaze stability
- Gaze stability is jeopardized by inevitable rotations induced by postural control
- Visual feedback supports gaze stability by tuning down postural reflexes
- Self-generated visual motion limits rotations without promoting compensatory turns

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**In brief**
A balance between mechanical stability and movement flexibility is necessary to achieve behavioral goals despite noisy sensorimotor systems. Cruz et al. combine virtual reality with quantitative analysis of behavior to describe an unexplored role of visual feedback in orchestrating the interplay between posture and gaze stability.
Fast tuning of posture control by visual feedback underlies gaze stabilization in walking Drosophila

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SUMMARY

Locomotion requires a balance between mechanical stability and movement flexibility to achieve behavioral goals despite noisy neuromuscular systems, but rarely is it considered how this balance is orchestrated. We combined virtual reality tools with quantitative analysis of behavior to examine how Drosophila uses self-generated visual information (reafferent visual feedback) to control gaze during exploratory walking. We found that flies execute distinct motor programs coordinated across the body to maximize gaze stability. However, the presence of inherent variability in leg placement relative to the body jeopardizes fine control of gaze due to posture-stabilizing adjustments that lead to unintended changes in course direction. Surprisingly, whereas visual feedback is dispensable for head-body coordination, we found that self-generated visual signals tune postural reflexes to rapidly prevent turns rather than to promote compensatory rotations, a long-standing idea for visually guided course control. Together, these findings support a model in which visual feedback orchestrates the interplay between posture and gaze stability in a manner that is both goal dependent and motor-context specific.

INTRODUCTION

Every step we take is defined by an interplay between behavioral goals, the reasons why movement is executed in the first place, and mechanical stability mediated by postural reflexes, which keep our body balanced as we move through complex and unpredictable environments. This interplay requires the brain to estimate self-generated movements to determine whether the motor programs at work are executed as intended and to correct them otherwise. In addition, postural reflexes ought to be tuned flexibly within some permissive range if behavioral goals are to succeed. The brain estimates self-motion by integrating internal motor-related activity with mechanical and visual information. Yet how the brain uses this internal estimate to balance behavioral goals with postural stability requirements is poorly understood.

Visual animals move their eyes and head in a structured manner following a combination of fast gaze shifts (saccades) and fixations that maintain gaze stable. In vertebrates, gaze saccades and fixations are controlled by distinct motor programs. Saccades are initiated centrally by a common command to rotate the eyes and neck together, whereas eye fixations are controlled by compensatory eye rotations initiated by proprioceptive or vestibular signals in the cervical-ocular and vestibular-ocular reflexes. Likewise, in walking insects, saccades are characterized by syndirectional yaw-rotations of head and body, whereas gaze stabilization during roll and pitch movements of the body is based on compensatory head rotations dependent on proprioceptive systems. These structured eye and/or head rotations are complemented with body movements that control course and further reduce motion blur. At the same time, noise within sensorimotor systems and uncertainties in the interaction between limbs and the environment can cause postural perturbations. Such perturbations lead to rapid (stepwise) readjustments in body posture that may affect gaze stability goals. Thus, the trade-off between gaze stability for accurate visual and spatial perception and posture control during exploratory walking provides an opportunity to examine the role of self-motion estimation on tuning such trade-off.

While vision is the primary beneficiary of gaze stabilization, it is also an important contributor to its control, complementing non-visual stabilization mechanisms. In both vertebrates and invertebrates, experiments in which visual motion stimuli are presented uncoupled from the animal’s behavior induce directed rotations of the eyes, or the head and body known as the optokinetic responses (OKRs), or optomotor responses (OMRs), respectively. Several models have proposed that OKRs and OMRs operate via a feedback control system in which visual error signals are minimized by compensatory turns, thereby reducing gaze and course deviations, although it remains unclear how OMRs coordinates movement across the body for gaze stability. Moreover, much less is known about how vision controls gaze stability in the absence of external visual perturbations, when visual input is self-generated (visual feedback). In insects, the sufficiency of OMR mechanisms to guarantee gaze stability under self-generated conditions has been challenged. OMRs depend on visual velocity signals that must be integrated to maintain the orientation of head and body fixed. Indeed, OMRs are modeled by a proportional-integral feedback control system. The integral aspect of the controller is prone to...
Here, we examine how *Drosophila melanogaster* uses visual feedback to control gaze stability during exploratory walking. We first established an experimental paradigm in which the fly displays regular gaze stabilization behavior and in which we control and manipulate visual feedback precisely (Figure 1). By combining virtual reality tools with quantitative analyses of movement, we found that flies execute distinct motor programs coordinated across the body to maximize gaze stability. Visual feedback is critical for gaze stabilization without affecting the accompanying head-body coordination. Instead, visual feedback prevents rotations induced by postural reflexes rather than promoting compensatory turns. Altogether, our findings suggest that visual feedback orchestrates the interplay between behavioral goals—gaze stability—and postural stability requirements in an exploring fly.

### RESULTS

An experimental paradigm to study the role of visual feedback in exploratory walking

To study the role of visual feedback in gaze control, we sought to establish a paradigm in which three conditions were met. First, the fly should display frequent walking paths with stable gaze. Second, movements across the body should be tracked in freely walking individuals. Third, visual feedback should be precisely manipulated. To this end, we developed FlyVRena, a virtual reality system for flies walking freely in a circular arena (Figures 1B and 1C). A heated wall prevents flies from following the perimeter of the arena (Figure 1D). To track movements of the head, body, and legs, we recorded high-magnification videos of the walking fly using a galvo-based high-resolution system coupled to positional information from a low-magnification tracking system (latency <16 ms). Visual stimuli were projected onto the arena’s floor in closed loop with the position of the fly (maximum latency of 40 ms). To measure the fly’s ability to detect FlyVRena’s stimuli, we compared behavior under natural versus reversed gain (RG) conditions (Figures S1A and S1B). In RG a leftward turn of the fly results in a leftward instead of a rightward rotation of the visual world, which induces continuous turning or circling behavior (Figure S1C). We define visual influence as the probability of circling under natural versus RG, a metric of the specific effect a virtual world has on the behavior of an individual fly (Figure S1D). Thus, FlyVRena allowed us to examine how the fly uses specific visual feedback to control gaze.

To examine the potential contribution of yaw visual feedback on gaze stabilization, we decouple the rotational and translational components of the fly’s visual feedback (Figure 1E; Video S1). Using the low-magnification system, the position of the visual stimuli was updated with the fly’s position. This configuration minimized translational visual feedback and facilitated keeping the angular size of objects on the fly’s retina constant during walking. Thus, in our experimental paradigm, flies explore a visually controlled virtual world where we can track movement across the body at high resolution and precisely manipulate self-generated visual information.

### Exploratory walking paths reveal two distinct motor contexts

To elucidate the organization of body movement underlying the exploratory paths (Figure 2A; Video S1), we obtained an
unbiased description of critical body kinematics linked to the motor programs at play\textsuperscript{57} (Figure S2A). The continuous character of locomotion poses a challenge for an unbiased segmentation of body kinematics. Therefore, we implemented an unsupervised segmentation of multivariate time series using a principled adaptive approach\textsuperscript{58} (Figure S2A; STAR Methods). Briefly, we model local dynamics in variable time windows with first-order linear models, using the forward and angular velocities and respective accelerations as inputs. In line with previous work,\textsuperscript{57} body kinematics were segmented into a set of windows ranging from 180 to 500 ms (Figure 2B). To group similar dynamics, we ran pairwise comparisons across all the segments to construct a dissimilarity matrix. Subsequently, we applied hierarchical clustering followed by a principled merging method via a logistic regression model (5-fold cross-validated binary classification, separability >99%; Figure 2A; STAR Methods). Applying this method on our dataset (84 flies, 441 min of recordings) led to 10 clusters representing body kinematics during continuous walking that were organized into three main branches (Figure 2C; for other behaviors, see Figure S2B). Two of these branches contained
kinematics with prominent angular velocity profiles accompanied by forward velocity profiles displaying decelerations followed by accelerations (Figure 2D, cyan, magenta, dark violet, and dark green). In addition, these branches contained clusters with prominent angular bias and high forward velocity profiles (Figure 2D, black, blue, orange, and spring green). By contrast, the third branch contained clusters that on average lacked prominent angular velocity profiles and were accompanied by high and steady forward velocity profiles (Figure 2D, dodger blue and lime clusters). Thus, body kinematics can be organized into regular sets of dynamics across different timescales revealing characteristic properties of the fly’s exploratory walking.

To examine the temporal organization of clusters, we identified common transition patterns between them (Figure 2E). We performed a singular-value decomposition (SVD) on the one-step probability transition matrix between clusters and obtained a ranked description of the most relevant transitions (transition modes) (Figure 2E; STAR Methods). Because the first transition mode contains mainly steady-state transitions (Figure S2D; STAR Methods), we focused our analysis on the subsequent modes. The highest contribution to the second mode corresponds to transitions between angular peak-like rotations and reductions in the angular velocity (Figure 2E, right). The third and fourth mode’s highest contributors include transitions from the steady forward velocity clusters to either the same clusters or to initiations of peak-like rotations, respectively (Figure 2E, right). Altogether, these findings indicated the presence of two main distinctive body motor programs, characterized either by angular spike-like events resembling body saccades (angular peak plus relaxation) that are executed in about 56% of the time (14,978 segments), or by segments with steady high forward velocity (forward runs) executed in about 44% of the total time spent walking (11,744 segments). Thus, the organization of the body kinematics during exploration resembles a fixation-saccade strategy in which the fly separates turning from forward displacements.

The structure of exploratory walking is configured to maximize gaze stability

A fixation-saccade strategy orchestrates specific motor programs across the body to maintain gaze stable. We examined movement coordination between the head and body with FlyVRena’s high-magnification system (Figure 3A; Video S2). Because the computational load of the unsupervised analysis grows exponentially with data size, we leveraged the information acquired from this analysis and developed a faster, supervised identification of body saccades (Figures 3A, 3B, S3, S4A, and S4B; STAR Methods). During body saccades, the head and body moved initially in sync regardless of light conditions (8 ms resolution; Figure 3C). At the peak of the body saccade, the head initiated a counter-rotation (Figures 3D and S4C) that helped to stabilize gaze as the body completed its rotation. Unlike in vertebrates, the head’s contribution to gaze shifts was minor relative to that of the body (Figure 3E), perhaps due to the head’s limited movement range (<20°) and the body’s fast maneuverability. By contrast, and like the vertebrate cervical-ocular reflex, during forward runs gaze stability is increased by opposite rotations of the head and body, with the head following the body by 66 ms regardless of light conditions (Figures 3F and 3G). Together, these findings reveal a structural organization of exploration that is independent of visual conditions and that is configured to increase gaze stability while restricting the duration of gaze shifts.

Visual feedback rapidly controls course stability during forward runs

The limited contribution of the head relative to the body on gaze control (Figure 3E) suggests that gaze stability increases with course stability. To examine course stability, we calculated path straightness per forward run (Figures 3A and 4A; STAR Methods). Consistent with gaze stabilization, walking paths were straighter in light versus darkness or with weak visual feedback (Figures 4A–4D and S1; see also Robie et al.65). The increase in path straightness was associated with a decrease in deviations from a stable body orientation (body angular deviations) (Figure 4B). If the fly uses the projected stimuli for course control, then the stronger the visual influence of the stimuli, the straighter the forward runs should be. To examine this prediction, we designed visual worlds with variable influence on the behavior of individual flies (Figures 4C and S1). We found that although size or density of dots influenced behavior non-monotonically across the population, visual influence and walking performance were always strongly correlated, i.e., the larger the visual influence, the straighter the forward run (Figure 4E). This finding indicates that self-generated visual signals reduce the magnitude of body rotations and straighten forward runs when motor programs across the body are coordinated to maintain gaze stable.

To test how fast visual feedback affects course control, we calculated path straightness as a function of the duration of forward runs across visual worlds (Figure 4F). Forward runs lasting only 300 ms were straighter under strong versus weak visual feedback. Moreover, short, body kinematics (~180 ms) within clusters linked to forward runs displayed reduced body angular deviations under strong visual feedback (Figure 4G). Altogether, these observations suggest that visual feedback controls course stability at a timescale close to the step cycle of the walking fly.51

Visual feedback prevents pairwise interlimb correlations underlying postural adjustments

To understand how visual feedback controls path straightness at fast timescales, we first examined the relationship between interlimb coordination and course control. To characterize interlimb coordination, we used machine learning tools to track each leg’s tip position in two dimensions (Figure 5A; Video S3; STAR Methods). We labeled forward runs across visual conditions based on the cumulative body angular deviations normalized by the traveled distance (path deviation; Figure 5B). Regardless of the magnitude and direction of the path deviation, on average, flies walked with tripod gait (Figures 5C and S5A), characteristic of high-speed walking (Figures 2 and S2; see also Mendes et al.61 and Wosnitza et al.64). Thus, path deviations are not caused by changes in gait structure; instead, they may be caused by changes in leg kinematics.

To examine whether variations in single-leg kinematics affect body angular deviations, we extracted the location orthogonal (lateral) and parallel (longitudinal) to the body axis of the anterior
extreme position (AEP) and posterior extreme position (PEP) of each leg relative to the body. The AEP and PEP can vary up to 0.3 body lengths (BLs), irrespective of the magnitude and direction of path deviations (Figure 5D). Interestingly, the lateral AEP of front legs changed systematically with path deviations (Figure 5D). This effect was clearer at a single-step level, where correlations between lateral leg placement and body angular deviations were more prominent for front than middle legs, and it was not observed in hind legs (Figure S5B). These findings indicate a specific relationship between each leg’s placement and body deviations, strongly suggesting that variability in limb kinematics causes self-paced path deviations during forward runs.

Notably, the lateral placement of a front leg in a single step (FLx) is not sufficient to explain the body angular deviations. If an initial displacement is not followed by a syndirectional FLx from the contralateral leg, body angular deviations don’t occur (Figure 5F). By contrast, if the initial displacement is followed by a correlated FLx, the magnitude of the correlation directly relates to the magnitude of the body angular deviation (Figure 5F). Changes in FLx over a step can displace the fly’s center of stability away from its center of mass (Figure 5G), thereby momentarily decreasing posture stability. To recover stability, posture control systems could either adjust lateral foot placement in the next step, like in humans, or dynamically engage viscoelastic properties of the musculoskeletal system, as observed in the cockroach. Our data are consistent with the first possibility. Initial FLx displacements were typically followed by contralateral FLx displacements in the same direction (Figures 5G and S5E). These front-leg pairwise correlations are strongly correlated with body angular deviations regardless of visual conditions (Figure SSD). However, the degree of the interlimb correlation was stronger in darkness and gradually reduced with stronger visual feedback (Figures S5E and S5F). Consistently, posture stability recovery was more robust in darkness than when visual feedback was available (Figures 5H and S5G). These findings show that posture control induces body angular deviations, thereby shifting gaze when the fly intends to keep it stable (Figure 3). Our data show that visual feedback controls gaze stability by tuning down rapid postural adjustments.
External visual perturbations induce body rotations with specific interlimb coordination

Next, we asked whether visual perturbations leading to body rotations induced similar interlimb coordination. We induced visual perturbations either by OMR visual stimuli (Figure S6A) or by RG (Figure S1A). During OMR, flies turn at high forward velocity or with saccades (data not shown) in the direction of the stimulus rotation (Figure S6A). In these biased forward runs, the body angular deviations were correlated with the front legs’ lateral AEP and anti-correlated with the hind legs’ lateral AEP (Figure S6B). Under RG, flies circle with near-perfect tripod gait (Figures S1 and S6C-S6E; Video S4). Like the OMR, the
body angular deviations were correlated with the front legs’ lateral AEP and anti-correlated with the hind legs’ lateral AEP (Figure S6F). By contrast, the longitudinal AEP was unaffected under RG (Figure S6G). The consistent anti-correlation between the hind legs’ lateral AEP and the body angular deviations during open- and closed-loop visual perturbations was never observed in natural conditions (Figures 5 and S6B, static condition). Furthermore, we found that head-body coordination under RG differs from the anti-correlation found under natural conditions (Figure S6I). Altogether, our data suggest that, regardless of the gait structure (Figures S6D, S6E, and S6H), a different movement pattern across the body emerge under visual perturbations.

Motion-sensitive circuits are crucial for the rapid tuning of posture by visual feedback

To determine the specific visual circuits tuning down postural adjustments, we silenced the activity of T4/T5 cells, the first population of neurons with direction-selective visual-motion responses, via the selective expression of the inward-rectifier potassium channel Kir2.1 (Figure 6A). These experimental flies displayed characteristic exploratory paths (Video S4), suggesting that visual motion information is not required for the fixation-saccade structure of exploratory walking (Figures 6B and 6C, left).

Silencing the activity of T4/T5 cells did not affect the dynamics of saccades (Figures 6D and 7D). However, walking paths during forward runs under strong visual feedback were markedly less straight in experimental flies compared with controls, resembling walking in darkness. This effect was not explained by an impairment in head body coordination (Figures 6B and 6C, right; Figures S7D and S7E). Consistent with this observation, we found that front-leg pairwise correlations were more prominent in experimental than in control flies (Figure 6E). This finding strongly suggests that pathways postsynaptic to T4/T5
cells mediate the tuning of postural reflexes to maintain walking straight and gaze stable.

**DISCUSSION**

Gaze stability is fundamental for visual and spatial perception. Here, we demonstrate that exploratory walking *Drosophila* attempt to keep gaze stable by moving forward in fixed directions while coordinating head and body rotations in antiphase (Figures 2 and 3). However, maintaining gaze stable during walking is not trivial because of internal and external sources of noise. We identified one component of this noise, the variability of leg placement, that momentarily decreases posture stability and triggers rapid limb adjustments that induce changes in course direction, thereby impairing gaze stabilization (Figure 5). We further show that the fly uses visual feedback to stabilize gaze at the expense of posture (Figures 4 and 5). Mechanistically, self-generated visual motion cues prevent rather than promote compensatory body rotations (Figures 5 and 6), a long-standing idea for visually guided course control (Figure 7A). Moreover, external visual disturbances induced a different leg movement pattern from the one observed under natural conditions. Taken together, these findings establish that self- and externally generated visual information engages different mechanisms to control goal-directed and visually guided locomotion. Overall, our work suggests a critical role for visual feedback in orchestrating the interplay between posture and gaze stability in a manner that is both goal dependent and motor-context specific.

**Active, self-paced visual control of walking**

In flies, course control is thought to depend on compensatory systems. The proposed models rely on a combination of directional and non-directional components, which together act via compensatory turns until an internal error signal is minimized. In flight, the directional (optomotor) component depends on a feedback system with integrative components. Several lines of evidence show that these compensatory systems cannot fully explain our findings during walking. First, the fly’s course control critically depends on visual motion pathways, in contrast to position-based oriented behaviors. Furthermore, while flies follow discrete high-contrast edges (Figure S7C), they seem not to fixate on single dots (Figure S7A). Second, we found no evidence for the presence of actively generated small-amplitude turns, suggested to constitute an operant trial-and-error strategy for the optomotor system under closed-loop conditions. Third, visual feedback does not generate compensatory turns, but rather prevents body rotations. Fourth, while optomotor mechanisms could support gaze stability in flying insects, the gain of the OMR, in our case on average 0.56, is too low to guarantee straight walking.

**Figure 6. Motion-sensitive circuits are crucial for visual tuning of postural reflexes**

(A) Left: schematic of the fly optic system highlighting direction-selective and motion-sensitive T4/T5 cells. Right: selective expression of Kir2.1:GFP in T4/T5 cells (green).

(B) Walking path straightness of controls (light/dark blue) and experimental exemplar flies (orange) in light (top) and dark (bottom) conditions. Visual environment: 10° dots.

(C) Visual influence (left) and path straightness (right) across visual environments. Color code same as in (B). Grand mean ±SEM, Wilcoxon rank-sum test: *p < 0.05; **p < 0.01; ***p < 0.005.

(D) Dynamics of the first principal component of body saccades in control (blue) and experimental flies (orange). Grand mean ±SEM.

(E) Front-leg lateral displacement (ΔFLx_{i+1}) versus the contralateral leg displacement in the preceding step (ΔFLx_i) in controls (blue) and experimental flies (orange). Grand mean ±SEM. See also Figures S6, S7, and Video S4.
Reversing the gain of the virtual reality system or presenting an OMR stimulus induced additional components of limb placement absent under natural conditions, which likely contribute to compensatory turns. Altogether, we propose that parallel mechanisms exist for course control that depend on self- versus externally generated visual motion. While external perturbations, such as a gust of wind or asymmetries in the locomotor apparatus induced by acute limb damage, rely on optomotor mechanisms for proper course control, the fast action of visual feedback reported here is critical in the context of self-paced variability during walking, where every step is prone to multiple sources of noise.

Neural mechanisms involved in visual tuning of posture control

It takes \(\sim 35\) ms or half the stance period\(^{51,62}\) (Figure S5C) for visual motion signals to excite T4/T5 postsynaptic neurons (T. Fujisawa, personal communication). Thus, to control leg placement, visual feedback could act upon the swinging leg if descending signals to the ventral nerve cord (VNC) travel fast with minimal delay. T4/T5 cells connect to visual projection neurons,\(^{79-81}\) many of which receive extra-retinal\(^{82,83}\) information that further accelerates visual processing.\(^{84,85}\) In addition, visual projection neurons project to regions densely innervated by descending neurons.\(^{86}\) In insects, descending neurons carry task-specific signals to modulate reactive forces, interlimb coordination, and leg placement.\(^{87,88}\) At the same time, local circuits within the VNC perform stepwise adjustments to control posture in the presence of internally and externally generated noise.\(^{33-36}\) Here, we show that posture reflexes respond to variability in leg placement via spatial interlimb correlations, which cause small changes in body orientation while recovering posture stability (Figure 5). Although the mechanisms underlying these spatial correlations remain unclear, we speculate that information about leg landing position in contralateral pairs contributes to postural adjustments, likely via the activity of interneurons integrating information across segments within the VNC\(^{38}\) and whose gain can be gated by visual feedback (Figure 7B). Alternatively, visual descending pathways may act on a step-by-step manner on leg sensory-motor systems\(^{89}\) (Figure 7C). Very little is known about the identity and mechanisms by which neurons within the VNC modulate leg placement or how visual descending circuits interact with such population of interneurons. The findings reported here constitute a possible framework to examine these important questions in the context of active vision and its role in goal-directed walking.

Anyone who tried to walk while blindfolded knows that vision is fundamental for course control. There is evidence supporting a role for vision in human leg placement,\(^{46}\) suggesting that a similar mechanism to the one we demonstrate in flies may underlie course control in humans.\(^{67}\) Investigating the role of visual feedback under naturalistic conditions in a system with a tractable number of neurons and a comprehensive array of anatomic, genetic, and physiological tools will produce a conceptual framework to elucidate the interplay between visual feedback and posture control during locomotion suitable for comparative studies across species and scenarios.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.cub.2021.08.041.

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AUTHOR CONTRIBUTIONS
Conceptualization, T.C. and M.E.C.; methodology, T.L.C. and S.M.P.; investigation, T.L.C. and S.M.P.; software, T.L.C.; formal analysis, T.L.C. and S.M.P.; supervision, M.E.C.; writing, T.L.C. and M.E.C., with input from S.P.M.

DECLARATION OF INTERESTS
M.E.C. is an advisory board member at Current Biology.

INCLUSION AND DIVERSITY

One or more of the authors of this paper self-identifies as a member of the LGBTQ+ community.

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# STAR METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Experimental models: Organisms/strains** | | |
| *Drosophila: Top Banana wild type* | Michael Dickinson | N/A |
| *Drosophila: w118; Otd-nls:FLP0; +* | Kenta Asahina | N/A |
| *Drosophila: w118; +; pJFRC56-10xUAS > myristomyo3-SV40 > eGFPKir2.1* | Gerald Rubin | N/A |
| *Drosophila: w118; R59E08-p65ADzP;R42F06ZpGdbd* | Michael Reiser | N/A |
| **Reagents** | | |
| Vectashield | Vector Laboratories | H-1200 |
| Normal Goat Serum, | Life Technologies | PCN5000 |
| PFA | Electron Microscopy Sciences | 15710 |
| **Antibodies** | | |
| Mouse monoclonal anti-nc82 | DSHB | RRID: AB_2314866 |
| Rabbit polyclonal anti-GFP | Thermo Fisher Scientific | Cat#A6455; AB_221570 |
| Alexa Fluor 633 conjugated goat anti-mouse | Thermo Fisher Scientific | Cat#A21050; RRID: AB_2535718 |
| Alexa Fluor 488 conjugated goat anti-rabbit | Thermo Fisher Scientific | Cat#A11008; RRID: AB_143165 |
| **Software and algorithms** | | |
| FlyVRena | This paper | github, https://github.com/ChiappeLab |
| Code for behavior analysis | This paper | github, https://github.com/ChiappeLab |
| MATLAB | The MathWorks, Inc. | N/A |
| Python version 3.7 | Python Software Foundation | N/A |
| Microsoft Visual Studio | Microsoft Corporation | N/A |
| **Deposited data** | | |
| Raw and Pre-processed Data | This paper | Mendeley, https://doi.org/10.17632/tvw5k488b6p.2 |
| **Equipment** | | |
| Infrared Cameras | IDS | IDS UI-3240CP-NIR/IDS UI-3360CP-NIR-GL |
| Infrared LEDs | Osram | 850nm SFH 4235-Z |
| LED current power supply | TENMA | 72-10480 |
| LED projector | Texas Instrument | DLP Lightcrafter 4500 |
| Telecentric lens | Telecentric Lens | 220mm WD CompactTL |
| Galvo mirrors | Thorlabs | GVS012/M - 2D |

## RESOURCE AVAILABILITY

### Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Eugenia Chiappe (eugenia.chiappe@neuro.fchampalimaud.org).

### Materials availability
This study did not generate new unique reagents.

### Data and code availability
All data reported in this paper can be found at Mendeley data: https://doi.org/10.17632/tvw.
All original codes used for behavior analysis are available at https://github.com/ChiappeLab.
Any additional information required to reanalyze the data reported in this work is available from the Lead Contact upon request, Eugenia Chiappe (eugenia.chiappe@neuro.fchampalimaud.org).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Fly Strains
*Drosophila melanogaster* were reared on a standard fly medium and kept on a 12 hr light/12 hr dark cycle. Experiments were performed with 2-to-4-day old flies with clipped wings from eclosion. Although at the beginning of this study we examined both females and males, under our experimental conditions we found that males display more robust and continuous walking behavior compared to females. Thus, in this study, we focused our analysis on male flies. We used the recently derived “Top Banana” WT line (TP, a gift from the Dickinson lab) because they engaged in exploration more readily than other WT lines (Figures 1, 2, and 3, comparisons across WT lines are not shown). Experiments were performed uniformly throughout the day since the variation of the time of the experiment could not explain relevant behavior variability (data not shown). An intersectional approach designed with Otd-nls:FLPo90 restricted the effector expression to the population of neurons of interest. The complete set of transgenic flies and their specific sources are listed in the Key resources table.

METHOD DETAILS

Immunostaining
Isolated brains were fixed for 30 min at room temperature in 4% paraformaldehyde in PBS, rinsed in PBST (PBS, 0.5% Triton X-100 and 10 mg/ml BSA), and blocked in PBST + 10% NGS for 15 min. Brains were incubated in primary antibodies (1:25 mouse ncc82 and 1:1000 rabbit antibody to GFP) at 4°C for three days. After several washes in PBST, brains were incubated with secondary antibodies (1:500 goat-anti-rabbit: Alexa Fluor 488 and 1:500 goat-anti-mouse: Alexa Fluor 633) for three days at 4°C. Brains were mounted in Vectorshield, and confocal images were acquired with a Zeiss LSM710 scope with a 40× oil-immersion or 25× multi-immersion objective lens.

Virtual Reality setup for exploratory walking
Single flies walked freely in a 90mm circular arena with walls heated up to 42°C by an insulated nichrome wire (Pelican Wire P2128N60TFEW). In order to prevent walking on the ceiling, the arena was covered with a glass plate pre-treated with Sigmacote (Sigma). The fly was video recorded using a near infrared camera (IDS UI-3240CP-NIR) at a frame rate of 60Hz and resolution 900x900 pixels. The camera has attached a 2x expander (Computar EX2C), a wide field lens (Computar 5mm 1:1.4 ½), and a filter against visible light (Thorlabs AC254-100-A-ML). The illumination was set beneath the arena by IR LEDs (850nm SFH 4235-Z), powered by a current power supply (TENMA 72-10480). The centroid position and orientation of the fly were determined via real time tracking. The tracking algorithm first performed a background subtraction using an image of an empty arena, followed by the application of a pixel intensity threshold. From the distribution of pixels within the contour of the thresholded image, the 2D centroid and main orientation were estimated. The final estimate of the position and orientation of the fly was given by a Kalman filter, which combined the recorded data with prior knowledge about the system to statistically minimize the difference between the real and estimated values.

Visual stimuli were projected onto a rear projection material (Da-Lite High Contrast DA-Tex), attached to the floor of the arena. We used a small LED projector (DLP Lightcrafter 4500, Texas Instruments) at a frame rate of 60Hz and a pixel size of 160px/mm. A custom-made software (FlyVRena, https://github.com/tlcruz/FlyVRena) generated the virtual worlds. FlyVRena implements the real time tracking of the position and orientation of the fly. FlyVRena uses computer game development libraries to render virtual objects and update their behavior accordingly (closed loop). The delay between the fly movement and the update of the world is about 40ms (as measured using a photodiode, Vishay BPW21R), which is generated mainly from the lag of the projection system. Virtual worlds were 2D environments with sets of textured square-based random dots (Figures 1, 2, 3, 4, 5, and 6 and S1–S7; Video S1) or textured square-based bars (“windmill environment”; Figure S7). The models of the textured squares were designed in Blender (https://www.blender.org), the textures were generated in MATLAB and Adobe Illustrator, and rendered using the FlyVRena software.

Following FlyMad, high-resolution images of the fly were recorded at 120fps and 1024x544 pixel resolution with a near infrared camera (IDS UI-3360CP-NIR-GL) and a telecentric lens (Edmund Optics 1x, 220mm WD CompactTL Telecentric Lens). The camera pointed to a pair of galvo mirrors (Thorlabs GVS012/M - 2D Large Beam (10mm) Diameter Galvo System) whose orientation was controlled by the FlyVRena software based on the instantaneous position of the fly to maintain the fly centered in the high-resolution frame (16ms lag; Video S2).

The size of our experimental arena (> 50BL) together with its aversive walls, circumvented the fly’s innate centrifobicism and promoted longer excursions throughout the arena (Figure 1D). Under these conditions, we found that flies organize their behavior in two distinctive phases with characteristics that resemble what has been described as saccades in walking flies and fixation-saccade in patrolling and exploratory flying flies. Under our experimental conditions, the fly may be structuring its behavior similarly while searching for an exit or a cool spot within the aversive landscape generated by the hot walls. Therefore, we refer to the behavior as exploratory walking.

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Data pre-processing for free walking assays

Post experiment, the change in the position of the fly was transformed to forward and side velocities, whereas the change of its orientation was transformed to angular velocity. Jump events of the fly were detected by a threshold in the fly acceleration (200mm/s²) and eliminated from subsequent analysis. Heading direction was inferred based on the persistence of the forward velocity, and if averaged forward speed between two jumps was negative, then orientation was rotated by 180° and velocities were re-calculated. All velocity signals within a window of 166ms centered at a jump were set to zero. The speeds were smoothed with a lowess algorithm with a window of 100ms. Activity was defined as translational speed greater than 0.5mm/s, or angular speed greater than 20°/s for at least 166ms. On the other hand, inactivity bouts smaller than 333ms were considered activity. Walking bouts were a subgroup of activity bouts in which a sequence of movement lasted for at least 333ms.

Unsupervised segmentation of movement dynamics

After pre-processing the data (84 flies, total number of walking segments = 64467, total time of spent walking = 441 minutes, percentage of time spent walking = 54 ± 17%), the description of the kinematics of the walking behavior consisted of time series of the angular and forward velocities, and their respective derivatives (accelerations). Under our experimental conditions, side velocities were minimal during forward runs, and highly coupled to angular velocity during turns. We used an adaptive methodology proposed in Costa et al.⁵⁸ to segment these input time series into locally linear dynamical systems of varying length. Briefly, in an iterative manner, the algorithm searches for “dynamical breaks,” that is, points at which there would be a need to switch between one model’s dynamics and another to best fit the data among arbitrarily defined set of candidate windows \(W_k\), \(i = 1 \ldots N\). A window \(W_i\) is modeled by locally fitting a first order Autoregressive Process (AR) defined as follows

\[
\begin{align*}
    x_t &= \bar{c} + A x_{t-1} + \eta t
\end{align*}
\]

where the set of parameters \(\theta = (\bar{c}, A, \Sigma)\) are estimated by least-square regression, with the coupling matrix \(A\), the covariance matrix of the error, \(\Sigma\), and the intercept, \(\bar{c}\). We denote \(W_k\) a window of length \(W(k)\) samples, and \(W_{k+1}\) as a window of length \(W(k + 1)\) samples, with \(\theta_k\) and \(\theta_{k+1}\) their respective models. A dynamical break is found if \(\theta_k\) cannot model the dynamics in \(W_{k+1}\), in which case \(W_k\) is chunked and the search for dynamical breaks starts again. Otherwise, a comparison with a new pair of candidate windows starts, i.e \(W_{k+1}\) and \(W_{k+2}\). The algorithm finishes when it chunks all the input time series. For our analysis, we considered a minimum candidate window of 10 frames at 60 Hz (~167ms) which corresponds to about 2 steps under fast tripod walking and a lower boundary for consistent movement in freely walking flies.⁵⁷ Thus, the choice of 10 frames as a starting candidate window gives us confidence to be able to capture most chunks of consistent dynamics in the dataset.

An analysis of the timescales of walking behavior (Figure 2B) revealed that the distribution of the duration of continuous locomotion-related dynamics decayed quickly after 350ms. Given that, our final analysis considered candidate windows from 167ms to 350ms (10-21 samples). Furthermore, we set the percentile of rejection under the null hypothesis that \(\theta_k\) models \(W_{k+1}\) to 98.5, and the number of surrogates to construct the null hypothesis distribution to 1000. Varying these parameters did not affect the overall distribution of segments (data not shown).

Clustering of movement dynamics

After segmenting the multivariate time series, we obtained 64467 segments of linear dynamics. A dissimilarity matrix is constructed by performing a pairwise comparison between segments. Given two segments, A and B and their corresponding models \(\theta_A\) and \(\theta_B\), their distance (or dissimilarity) is computed by asking how likely the model of the concatenation of segments A and B, defined as \(\theta_C\), can model the individual segments. Equation 2 shows the measure of distance, where \(\lambda\) is a log-likelihood function. The output of this procedure is a symmetric, positive-defined squared matrix of 64467 rows (Figure S2A).

\[
    d_{ab} = \lambda_{ac} + \lambda_{bc}
\]
Analysis of transitions between clusters

To examine how the behavior discretized in clusters unfolds in time, we compute a one-step probability transition matrix between continuous locomotion-related clusters. To account for the underlying structure of behavioral transitions, we performed a singular-value decomposition (SVD) on the one-step probability transition matrix. This factorization of the matrix allowed us to get a ranked description of the most important transitions between clusters. The SVD factorization of a one-step transition matrix decomposes it as a summation of transition modes. Therefore, the transition matrix $A$ with $m$ states can be expressed as follows:

$$ A = U S V^T = \sum_{i=1}^{m} s_i U_i V_i^T $$

where the outer product of each left and right singular vectors $U_i V_i^T$, together with the associated singular value, generates a rank-1 matrix that here we refer as a transition mode. The larger the associated singular value, the slower the timescale of the system is reflected in the transition mode. The leading singular value contains information about long timescales—the steady-state of the system, where there is little predictability of the current state, and therefore, it reflects the averaged probability of transitioning into any other behavioral state. Because the larger clusters are the ones containing no angular bias (on averaged, “lime” and “dodge blue”), the first mode is characterized mainly by transitions from any clusters to these high forward velocity clusters (Figure S2E). Thus, we focused our analysis on the subsequent modes. The second transition mode accounts for local transitions between spike-like rotations followed by relaxations, in both directions. The third and fourth modes organized transitions from forward segments to either initiation of a rotation or continuation of a high-forward velocity segments. This analysis shows the existence of two main different motor contexts in which the fly transitions on one-step basis composed by fast angular rotations resembling body saccades (spikes plus relaxations) and high-forward velocity segments.

Identification of spikes in the fly’s angular velocity

Spike-like dynamics described by the combination of angular velocity spikes and relaxation clusters corresponded to fast and stereotyped rotations of the fly (Figure 2). To quickly classify events as spike-like, we took advantage of this stereotypy and used a continuous wavelet transform strategy. For each walking bout, the continuous wavelet transform of the angular velocity was computed using Gaussian wavelets, and a signal with power at a range of 10-15Hz was extracted (frequency signal). Note that while the selection of the frequency band slightly affects the spike detection, it is not a determinant factor in subsequent analysis (Figure S3B). Next, the local maxima of the frequency signal, and of the absolute value of the angular velocity were calculated. If these local maxima coincide between the two sets of local maxima, these were labeled as putative spike turns (Figure S3A). Putative spike turns were filtered first to remove small local maxima (< 200°/s) that did not disrupt the forward velocity of the fly (variance of forward speed < 3mm/s), and then to roughly match the signal with a template obtained via PCA on a subset of pronounced peak aligned spike-like events (the first principal component explained ~90% of the shape variance in a 500ms window and was used as the template; Figures 6D and S4A). Putative spike turns in which the square distance between the scaled shape and the template was smaller than a cutoff of 0.15 were labeled as angular spike events (Figure S3A). The free parameters of the classifier did not affect the results over a wide range of possible parameters (Figure S3B).

To identify forward walking segments (“forward runs”), angular spike events were removed from the walking bout. The remaining segments longer than 333ms, and with an average forward velocity greater than 6mm/s were defined as forward runs throughout this study.

Local curvature, path straightness and visual influence

Local curvature is calculated with a running window of 333ms centered on each point of the fly’s path. A 333ms window centered on each point of a forward segment was used to calculate the distance between the central data point and a line defined by the edges of the window (instantaneous deviation from an ideal straight path). The path straightness of a forward run was defined as the sum of traveled distances within the window divided by the sum of deviations. Path straightness per fly is the weighted average of all path straightness per forward run, with weights given by the total distance walked in each forward run (Figures 4 and 6).

Under natural gain conditions, when a fly rotates to the left, the world rotates to the right. In contrast, under the reversed gain (RG) condition, when the fly rotates to the left, the world rotates left too. The RG condition induces a persistent rotation of the fly, a behavior known as circling (Video S4). Visual influence was defined as the difference between the probabilities of circling in RG versus natural gain conditions (Figure S1). Under the dark condition we artificially divided the dataset into the same trial structure as in the random dot stimulus experiment and calculated visual influence in the same way.

Due to the delay between the estimate of the fly’s position and orientation and the update of the visual environment (~40ms), when the fly stopped walking under the reversed gain condition, the visual environment continued rotating with the previous angular velocity of the fly for three consecutive frames. This brief “open-loop” segment was sufficient to induce a head optomotor response (Figure S7). Typically, after the stimulus stops moving, the head remains with an offset position driven by the stimulation until the fly initiates a behavior.

Head tracking

High-resolution images were filtered with a Gaussian filter (width = 16pixels), and background subtracted. Pixels that belonged to the fly were estimated with a combination of edge detector operations, and morphological transformations until a connected object with
dimensions like the fly was detected. The centroid and orientation of the object were used to translate and rotate the frame to keep a vertical fly at the center of the frame. Next, a window around the head was cropped from the frame. The top of the thorax of the fly was masked out in the cropped frame, and the head was segmented using a combination of edge detector operations and morphological transformations. Head rotation was calculated via cross correlation with a template upright head. The head tracking error is the squared error between the template and the rotated head. The head angle trace was smoothed using a lowess algorithm with a window of 83ms. Frames in which the head tracking error was high (top 2.5% of the frames) were removed from the original time series.

As an alternative strategy, we used DeepLabCut to estimate the rotation of the head with respect to the body of the fly. We trained a neural network with a subset of images where the fly was in the upright position (see above) with four labels to track: Left Eye, Right Eye, Neck, and Thorax ending. After applying the tracking network to the full dataset, we transformed the positions of the labeled points into head angles. No significant differences were observed when comparing the two methods (Video S2).

**Leg position extraction**

High resolution video frames were background subtracted. Pixels that belonged to the fly were estimated with a combination of edge detector operations, and morphological transformations until a connected object with dimensions like the fly was detected. The centroid and orientation of the object were used to translate and rotate the frame to keep an upright vertical fly at the center of the frame. Next, a cropped window of 400x400px around the fly was stored in a new aligned video. We used DeepLabCut to estimate the location of the following features from the high resolution aligned videos: top of the head, left eye, right eye, neck, tip of the thorax, tip of the abdomen, tip of left front leg, tip of right front leg, tip of the left middle leg, tip of right middle leg, tip of left hind leg and tip of right hind leg. These features were thresholded based on the accuracy of estimation parameters and aligned to the low-resolution preprocessed data.

Inter-leg phase was calculated by dividing the time delay in the cross covariance between leg position signals and the period as measured in the autocovariance of those signals. Swing-stance transitions for each leg were calculated by detecting the local maxima (swing to stance -PEP) and local minima (stance to swing-AEP) in the leg position aligned to the body orientation. Landing position is taken as the transition from swing to stance, and a step is defined by the swing of a single tripod.

**Single step analysis**

For each individual leg, we used the AEP and PEP of the leg position signal to identify swing, stance, landing and lift-off. We calculated the deviations in body angle (same as path deviations) as the integrated angular velocity between consecutive lift-offs. We correlate body angular deviations with the leg lateral and longitudinal displacement between lift-off and landing, the swing and stance time between those same consecutive lift-offs (Figures 5, S5, and S6). Leg placement correlation was calculated by multiplying leg lateral placement in consecutive steps (Figures 5, 6, and S6).

**High quality steps**

Only steps where we can identify stance and swing times larger than 16ms and smaller than 125ms, and swing amplitudes smaller than 0.75BL are considered for analysis to remove tracking errors that can affect the landing position identification. High quality steps include 68% of all detected steps.

**Center of stability calculation**

We calculated the landing positions of all the legs and considered for analysis only the steps where all the legs fulfilled the previous established criteria for step quality. We considered legs belonging to a triangle when they land within 16ms of each other. We consider the center of stability as the geometric center of the triangle defined by the landing positions of the legs. (Figure 5F). The reduction in posture stability was calculated by the distance in the orthogonal direction to the fly orientation between the centroid of the fly and the center of stability in a particular step (Figure 5F).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

We performed a two-sided Wilcoxon signed-rank test for paired groups, two-sided Wilcoxon rank-sum test for comparisons between two independent groups and t test for the correlation analysis. In addition, for statistics in the cross-covariance analysis we performed a resampling, bootstrapping based method where new velocity signals were created by shuffling the time of different walking bouts, distorting the signal without affecting its general statistics. The new shuffled signal undergoes the same cross covariance analysis, and the procedure is repeated 1000 times to obtain the shuffled distributions in Figures 3E and 3F.