Functionally distinct Purkinje cell types show temporal precision in encoding locomotion

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Purkinje cells, the principal neurons of cerebellar computations, are believed to comprise a uniform neuronal population of cells, each with similar functional properties. Here, we show an undiscovered heterogeneity of adult zebrafish Purkinje cells, revealing the existence of anatomically and functionally distinct cell types. Dual patch-clamp recordings showed that the cerebellar circuit contains all Purkinje cell types that cross-communicate extensively using chemical and electrical synapses. Further activation of spinal central pattern generators (CPGs) revealed unique phase-locked activity from each Purkinje cell type during the locomotor cycle. Thus, we show intricately organized Purkinje cell networks in the adult zebrafish cerebellum that encode the locomotion rhythm differentially, and we suggest that these organizational properties may also apply to other cerebellar functions.

Purkinje cells | locomotion | central pattern generator | cerebellum | zebrafish

The vertebrate cerebellum has powerful computational abilities (1–3) and, thus, is involved in numerous diverse motor and nonmotor functions (4–14). Traditionally, the cerebellar cortex has been thought of as containing many uniform microcircuit modules (15) that use the same computational model, influenced only by variations in inputs (sensory, motor, or cognitive). Purkinje cells, the principal cells behind cerebellar computation, were thus presumed identical. However, recent evidence emerged demonstrating functional and molecular differences between Purkinje cells (16, 17), urging the need for meticulous evaluation of any structural and functional variability existing within the Purkinje cell population.

Purkinje cells are regularly active during locomotion (10, 18–24), yet their firing activity varies significantly across the locomotor cycle (10, 18, 19). The degree to which such functional inconsistency reflects differences in Purkinje cell properties and connectivity as the cells regulate locomotion remains still unclear.

We investigated the organization of Purkinje cell population in detail and probed their contribution to locomotion using anatomical, electrophysiological, and behavioral approaches in adult zebrafish. We discovered diversity in the Purkinje cell population and found that Purkinje cells are organized into at least four functionally and morphologically distinct groups, each firing during a particular phase of the locomotor/swim cycle. Our results reveal the architectural complexity of the adult zebrafish cerebellum, and this organizational scheme might be relevant to other cerebellar functions.

Results

Purkinje Cell Synaptic Output Affects Locomotor Performance In Vivo.

Purkinje cells, the principal neurons of the cerebellar computations, play a key role in the dynamic formation of internal models of the cerebellum (25); thus, they exert a powerful influence in the generation and execution of smooth motor behaviors (26). As a first step toward understanding how Purkinje cell activity is related to the locomotor behavior of the adult zebrafish in vivo, we used a transgenic animal line Tg(aldoca:BoTx-GFP) that expresses botulinum neurotoxin under a specific Purkinje cell promoter (aldoca; SI Appendix, Fig. S1 A and B) (27). Botulinum neurotoxin silences synaptic transmission by blocking neurotransmitter release mediated by vesicular soluble N-ethylmaleimide-sensitive factor attachment protein receptor (SNARE; SI Appendix, Fig. S1A) (28). We examined the effect of Purkinje cell silencing on adult zebrafish by recording and analyzing locomotor behavior during spontaneous swimming episodes using the open field test (SI Appendix, Fig. S1C and Materials and Methods). While the locomotion itself was not affected as the animals could perform movements during the task, our analysis revealed that animals with silenced Purkinje cell output generated an erratic form of body displacements (SI Appendix, Fig. S1D) and produced significantly slower speed swimming episodes compared to the control animals (SI Appendix, Fig. S1 E and F). Further analysis of locomotion revealed that the BoTx animals were more immobile than the controls, which eventually affected the overall distance they traveled (SI Appendix, Fig. S1G). Together, our results highlight the critical influence of Purkinje cell outputs in the generation of a smooth and well-coordinated locomotor behavior.

Identification of Purkinje Cell Diversity. We evaluated the organizational scheme of Purkinje cell populations in adult zebrafish using a transgenic line in which an enhancer element derived from zebrafish carbonic anhydrase 8 (ca8; refs. 29, 30) drives enhanced green fluorescent protein (eGFP) expression exclusively in all Purkinje cells (PV+, Fig. L4 and SI Appendix, Fig. 2A).

Significance

Purkinje cells, the key cells for cerebellar computations, are thought to function similarly throughout the cerebellum and to comprise a uniform neuronal population. Here, we challenge this notion by providing detailed evidence that the Purkinje cells are organized into distinct types, each exhibiting specific activity patterns that are associated with a particular phase of the zebrafish swim cycle during locomotion. Connectivity experiments suggest that all the identified Purkinje cell types participate in orchestrating the cerebellar circuit ensembles, which are, therefore, highly heterogeneous.

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We focused on the corpus cerebelli (CCe), the largest and most accessible area of the adult cerebellum, where the cellular organization is highly conserved between fish and mammals (31). In the mammalian cerebellum, distinct functional Purkinje cell populations were identified by ZebrinII expression (16, 32), while in zebrafish ZebrinII is expressed in all Purkinje cells (ref. 33 and Fig. 1A). Yet, systematic analysis of Purkinje cell soma size revealed a large variability (Fig. 1B) with no obvious topographic organization within the CCe (Fig. 1B), indicating the possibility that distinct Purkinje cell subpopulations may exist in the cerebellar circuitry.

Next, we asked if the physiological properties of the Purkinje cells varied in relation to the cell soma size. We performed whole-cell patch-clamp recordings using a recently developed ex vivo preparation of the intact adult zebrafish brain (SI Appendix, Fig. S3A and Materials and Methods). As seen before in larval zebrafish (22, 23), the adult Purkinje cells displayed high spontaneous activity comprising simple and complex spikes (SI Appendix, Fig. S3B). This spontaneous activity hindered the assessment of the firing pattern properties of the Purkinje cells. Thus, we applied a bias hyperpolarization current to silence their spontaneous activity, and we found that this intervention did not affect their firing pattern (SI Appendix, Fig. S3C). We found that all recorded adult Purkinje cells displayed three similar properties: 1) strong hyperpolarizing current injection caused an immediate sag potential, 2) small depolarizing currents induced sodium-based spikes, and 3) large depolarizing current injections produced calcium-based spikes (SI Appendix, Fig. S3 D–G). Besides these common properties, we surprisingly observed that repetitive firing varied significantly between the Purkinje cells in response to increasing steps of depolarizing current (Fig. 1C). Using the elicited responses to the current injection steps, we grouped the Purkinje cells into four broad categories. Type I displayed strong spike frequency adaptation (strong adapting); type II discharged several action potentials with pronounced spike frequency adaptation (adapting); type III fired tonically during depolarizing current pulses without spike frequency adaptation (not-adapting); and type IV displayed bursting firing properties (bursting). Unbiased random recordings of several Purkinje cells (n = 124) throughout the whole CCe revealed the differential representation of the four Purkinje cell types (Fig. 1D–F). More interestingly, the soma size varied significantly between the different Purkinje cell types (Fig. 1D–F). We hypothesized that the soma size could influence the spontaneous activity comprising simple and complex spikes (Fig. 1D–F). Next, we questioned whether each Purkinje cell type relates to a specific soma size. Our analysis showed a broad association between firing type and soma size (Fig. 1F) that also reflected associated differences in their input resistance (Fig. 1G). Further analysis revealed specific cellular properties distinctive to each of the four adult Purkinje cell type categories (Fig. 1H and SI Appendix, Fig. S4). We found that the input resistance, the rheobase for the sodium spikes, the sodium spike frequency, and the calcium spike frequency were significantly different between all Purkinje cell type categories (Fig. 1H and SI Appendix, Fig. S4). The existence of distinct Purkinje cell types was also confirmed by 2D). We observed that all of the Purkinje cells, irrespective of type, had common morphological properties such as spineless proximal dendrites, distal dendrites with spines, and numerous axonal collaterals extended in large distances from the soma (Fig. 1J, Insets and i–ii). Unlike the strict two-dimensional dendritic spread of mammalian Purkinje cells, adult zebrafish Purkinje cells extend in three dimensions (Fig. 1L and Movie S1). Yet dendritic morphology clearly varied between the different Purkinje cell types (Fig. 1K), exhibiting distinct Sholl profiles (Fig. 1M). Types I and IV have more confined dendritic fields than types II and III, which have extended dendrites covering larger areas within the adult CCe (Fig. 1K–O). These data indicate a link between the physiological properties and morphologies of the Purkinje cell types, suggesting that each type may receive and process different inputs. Altogether, our results argue against the notion of a uniform Purkinje cell population and, instead, suggest a diverse population organized into morphologically and physiologically distinct types that may perform specialized functions. Yet, these different Purkinje cell types are topographically not segregated.

**Connectivity Patterns between Purkinje Cell Types.** We found that numerous Purkinje cell axonal collaterals exist and extend within the adult zebrafish CCe, including the Purkinje cell layer (Fig. 1J, Inset and iii and Movie S1). Previous characterization of axonal collaterals confirmed Purkinje cell interconnectivity (35, 36) and proposed its potential participation in generating prolonged responses (37) and synchronized firing (38) to control the activity of their targets (39). We tested whether adult zebrafish Purkinje cells are interconnected and whether a particular connectivity pattern exists between the distinct types by performing dual whole-cell patch-clamp recordings of identified adult zebrafish Purkinje cells (Fig. 2A). We observed that single and train of action potentials in one Purkinje cell could induce vigorous but small-amplitude monosynaptic GABAergic inhibitory postsynaptic potentials (IPSPs) in other Purkinje cells (Fig. 2B and SI Appendix, Fig. S5A). Yet, the Purkinje cell interconnectivity was high (70–80%) and showed no preference in relation to the type (Fig. 2C). Moreover, we tested if any differences in the connectivity strength (IPSP amplitude) relates to connectivity between Purkinje cells of the same or different type, yet we did not detect any difference (Fig. 2C). It also became apparent that in all pairs (connected and not-connected), there was no significant difference in relation to the distance of the recorded cells with the type that is categorized (Fig. 2C). Next, we asked whether the connectivity between the Purkinje cell types exhibited any directionality preferences (SI Appendix, Fig. S5B). Most connections (~80%) were bidirectional (Fig. 2D). Our morphological reconstruction experiments showed Purkinje-to-Purkinje cell dye coupling occurring following intracellular neurobiotin injection into a single cell (Fig. 2E, black arrows), implying the presence of gap junctions (electrical synapses). We also confirmed the existence of connexin 35/36 puncta on Purkinje cell somata (Fig. 2F). Bidirectional electrical coupling between Purkinje cells was confirmed using electrophysiology (Fig. 2G), but occurred only between pairs of the same Purkinje cell type that were also chemically connected (Fig. 2G). Such coupling may be essential for synchronizing cerebellar network activity as reported before (40, 41). Collectively, our results portray communication between the four Purkinje cell types and imply that all types identified here participate in the organization of the cerebellar circuit.
Fig. 1. Diverse cellular, firing, and morphological properties of the adult zebrafish Purkinje cells. (A) Specific expression of eGFP in all Purkinje cells of brains from the Tg(Ca8:eGFP) line. All eGFP-expressing Purkinje cells (green) are ZebrinII and PV positive (magenta). (B) Quantification and intermingling distribution of the Purkinje cells soma size. (C) The Purkinje cells display distinct firing patterns. Black trace shows the response at the rheobase. (D) Differential representation of the Purkinje cell types in the adult zebrafish cerebellum. (E) Purkinje cell types in the CCe lack topographic organization. Axes represent the normalized distances of the midline, lateral, rostral, and caudal edges of the CCe. (F) Purkinje cell types have significantly different soma sizes (P < 0.0001, one-way ANOVA/Tukey's post hoc test). (G) Correlation between soma size and input resistance (Rin) segregates the different Purkinje cell types. (H) Normalized mean values of the electrical properties observed for the Purkinje cell types that are detailed described in SI Appendix, Fig. S4. Normalizations were performed for each property to the highest obtained value. (I, Left) t-SNE plot depicting clusters of the Purkinje cells based on the electrophysiological properties of the dataset as in H and SI Appendix, Fig. S4. (I, Right) Fan (polar) shaped hierarchical clustering dendrogram of the electrophysiological data yielded similar results to t-SNE. All cells and data colored by assigned cell type. (J) Reconstructed neurobiotin-filled Purkinje cell showing the common morphological features for all Purkinje cell types. (i) Distal dendrites with spines. (ii) Proximal spineless dendrites. (iii) Axonal collaterals. (K) Reconstructed neurobiotin-filled Purkinje cells are showing the morphology of each type. (L) Dorsal view of the cerebellum (CCe) showing the three-dimensional intermingled distribution of the distinct types of Purkinje dendrites. (M) Sholl analysis of dendritic complexity with increasing radial distance from the soma center for each Purkinje cell type. The solid lines represent the mean of the intersections while the shades refer to the SEM. (N) Analysis of dendritic trees on each Purkinje cell type. (O) Plot illustrating the Purkinje cell soma and the rostro-caudal distribution of the dendrites obtained from the dorsal view reconstructions. The lines represent the maximum and minimum rostro-caudal position. All Purkinje cells are aligned based on their soma (solid circles). CCe, corpus cerebelli; PV, Parvalbumin; RMP, resting membrane potential; Rin, input resistance. Data are presented as box plots or violin plots showing the median with 25/75 percentile (box and line) and minimum–maximum (whiskers). ***P < 0.001; ****P < 0.0001; ns, not significant. For detailed statistics, see SI Appendix, Table S1.
Purkinje Cell Types Show Temporally Precise Activity during Locomotion. Understanding the structure–function relationships behind specific behaviors is a key goal in neuroscience. Purkinje cells are regularly active during locomotion, shaping and coordinating commands with the locomotor cycle, but their activity proved very variable when executing locomotion (10, 18–20, 24), suggesting a dynamic, regulated, computational performance throughout the locomotor episode (10).

Motivated by our previous findings, we explored the possibility that each Purkinje cell type could encode locomotion differently. We performed whole-cell patch-clamp recordings on individual Purkinje cells while recording motor nerve activity of the ipsilateral central pattern generator (CPG) in an ex vivo preparation (ref. 42 and Fig. 3A). We induced fictive locomotion by electrically stimulating (10 pulses, 1 Hz) the descending axons from the brainstem and assessed individual Purkinje cell activity relative to locomotor burst activity (SI Appendix, Fig. S6A). As seen before in other species, we observed large variability in Purkinje cell activity during fictive locomotion in zebrafish (SI Appendix, Fig. S6B and refs. 10, 19, 24). Surprisingly, we also detected Purkinje cells that did not discharge during the ongoing swim episode (SI Appendix, Fig. S6B, top trace).

Next, we assessed the relationship between Purkinje cell type categories and activity patterns during swimming (Fig. 3B). We found that of the four types identified, only types II, III, and IV were active during locomotion, while type I cells generated no action potentials (Fig. 3B and C). Also, we observed a significant membrane depolarization in types II and III during locomotion (Fig. 3D), suggesting the presence of an underlying network excitation during the execution of the motor program. Analysis of the Purkinje activity as a function of swimming frequency showed that the different types of Purkinje cells are deployed at all different locomotor frequencies obtained (Fig. 3E).

We further studied the activity of the cell types with respect to the swim cycle phase revealed an unexpected Purkinje type relative phase preference: type II (adaptating) cells exhibited phase-locked firing at peak CPG activity while types III and IV (not adapting and bursting) discharged only during the inhibitory phase (Fig. 3F). Moreover, Purkinje cells of type III discharged just before or after activation of the ipsilateral CPG, while type IV cells were active throughout the midcycle inhibitory phase (Fig. 3F). One remaining question concerned the reason why type I cells were inactive during ongoing locomotion and escape response (SI Appendix, Fig. S7A). Voltage-clamp recordings revealed that, during the locomotor episode, type I cells receive significantly less excitatory input currents excitatory postsynaptic currents (EPSCs) (SI Appendix, Fig. S7B) that differ in frequency and amplitude from those they receive after completing locomotion (SI Appendix, Fig. S7C). Interestingly, we also observed that all type I cells discharge several seconds (~9) after fictive locomotion ends (SI Appendix, Fig. S7D) using both sodium- and calcium-based spikes (SI Appendix, Fig. S7E). Collectively, these data show that the diversity identified in the Purkinje cell population could be meaningful for encoding and shaping different aspects of the locomotor behaviors.

Discussion

We discovered structural, physiological, and functional diversity in the adult zebrafish Purkinje cell population of the CCE (SI Appendix, Fig. S8). Distinct Purkinje cell types were identified and demonstrated extensive within- and between-type communication, indicating a functional specialization to encode locomotion (SI Appendix, Fig. S8). Our results portray an architectural model of adult cerebellar microcircuits and imply that different types of Purkinje cells are necessary for dividing the labor of the heavy computational duties of the cerebellum.
The Purkinje cell population is well characterized in fish and mammals (3, 10, 22–24, 27, 30, 33, 35, 36, 43, 44), but little is known of its intrapopulation differences (45–47). Purkinje cell compartmentalization was previously observed through ZebrinII expression (32). Yet only recently did evidence of a disparity in Purkinje cell groups emerge, as shown by significant differences in activity between cells of varying ZebrinII identity (16), implying the possible existence of distinct developmental programs that direct the specification of Purkinje cells with type and area identities. However, our results reveal a nonontopographic organization of the Purkinje cell types in the zebrafish cerebellar circuit. The lack of topography in conjunction with the fact that all of the Purkinje cells in zebrafish are ZebrinII positive (33) could imply that the mammalian Purkinje cell compartmentalization developed later in the evolution. We cannot rule out the possibility that the Purkinje cell diversity can also reflect the age of the neurons, although in zebrafish the vast majority of the Purkinje cells are generated early during development, within the first week (2–7 dpf), and their number remains relatively stable afterward (30). By that time, the Purkinje cells shape their dendritic morphologies and establish their connections (27). Therefore, our data, in conjunction with previous studies, could also suggest the possibility that the Purkinje cell diversity in zebrafish could already arise early during the development. Recent single-sequence analysis provided further evidence, showing that different Purkinje cell types indeed exist in the mammalian cerebellum (17, 48). Further evaluation of the diversification schemes for zebrafish Purkinje cell development is, therefore, critical.

What is the importance of multiple types of Purkinje cells to exist in the cerebellar microcircuit ensemble? Do they have distinct functions? Our results resoundingly imply that they do and support the importance of Purkinje cell diversity in the cerebellar microcircuit and the distinct contributory function of each cell type. Each of the distinct Purkinje cell types we identified possessed specialized functions for encoding different phases of the locomotor cycle; only the Purkinje cells of type I did not fire during swimming and escape but did so after. Given that the type I Purkinje cells discharge exclusively after the completion of the swimming and actively receive less input, they could well contribute to encoding the overall locomotor episode. One intriguing question that remains is to identify the Purkinje cell types that exert a powerful influence in motor learning. Future studies that selectively probe the physiological activities of the different Purkinje cell types will provide further insight into the type-specific role in new motor skill acquisition and information flow within the cerebellar microcircuitry.

Purkinje cells are critical for generating cerebellar output functions and often demonstrate coordinated synchronous activity (21, 38, 39, 49). We showed that extensive communication between Purkinje cells depends mainly on inhibitory GABAergic chemical neurotransmission that is occasionally supplemented by electrical synapses (gap junctions). Mutual inhibition and electrical synapses can mediate synchronous coordinated activity within neuronal networks (50). The broad chemical interconnections between the adult zebrafish Purkinje cells could account for the formation of the numerous function-dependent Purkinje cell ensembles that control the execution of motor behaviors.

Fig. 3. Adult zebrafish Purkinje cell type-specific activity during locomotion. (A) Ex vivo setup of the brain-spinal cord allows simultaneous recordings of Purkinje cells and ipsilateral motor nerves. Stimulating descending inputs elicit a swimming episode. (B) Sample recordings from each Purkinje cell type during locomotion. (C) Purkinje cell types II, III, and IV discharged during locomotion; type I did not fire (P = 0.0003, one-way ANOVA/Tukey’s post hoc test). (D) Significant changes to more depolarized membrane potential were observed in type II and type III Purkinje cells during locomotion (P < 0.0001, one-way ANOVA/Tukey’s post hoc test). (E) Graph showing the activity of the different Purkinje cells as a function of the instantaneous swimming burst frequency. Individual data points represent the instantaneous swimming frequencies of all swimming cycles where the respective Purkinje cell produced at least one action potential. (F) Purkinje cells show type-specific phase-locked activity during the swim cycle. (Top) The activity of six individual Purkinje cells from each type. (Middle and Bottom) Population data from all of the Purkinje cells recorded from each type, followed by a frequency distribution graph. The gray trace is a representative motor nerve recording showing the duration of the swim cycle. 0 defines the pick of the motor nerve activity. Data are presented as box plots showing the median with 25/75 percentile (box and line) and minimum–maximum (whiskers). **P < 0.01; ***P < 0.001; ****P < 0.0001. For detailed statistics, see SI Appendix, Table S1.
while the electrical interconnection could establish their synchronization (21). Thus, Purkinje-to-Purkinje cell type connectivity is particularly interesting as it can shape several parameters of target cell activity, such as timing, gain, tuning, rhythmicity, and synchrony (51).

The cerebellum is widely recognized for its critical role in locomotion (4, 10, 11, 18–24). Nevertheless, recent studies imply its role in processing nonmotor functions such as perception, cognition, emotion, reward, and social behavior (5–9, 12–14). Cell-type-specific vulnerability (55) and the surrounding head tissue were carefully used to expose the cerebellum, and the preparation was transferred to a recording chamber that was continuously perfused with an extracellular solution containing 135.2 mM NaCl, 2.9 mM KCl, 1.2 mM MgCl₂, 2.1 mM CaCl₂, 10 mM Hepes, and 10 mM glucose at pH 7.8 (adjusted with NaOH) and an osmolality of 290 mOsm. For whole-cell intracellular recordings of Purkinje cells in voltage- and current-clamp mode, electrodes (resistance ~15 MΩ) were pulled from borosilicate glass (outer diameter, 1.5 mm; inner diameter, 0.87 mm; Hilgenberg) on a micropipette puller (model P-97, Sutter Instruments) and filled with an intracellular solution containing 120 mM K-gluconate, 5 mM KCl, 10 mM Hepes, 4 mM Mg₂ATP, 0.3 mM Na₃GTP, and 10 mM Na-phosphocreatine at pH 7.4 (adjusted with KOH) and osmolality of 275 mOsm. GFP-positive Purkinje cells were visualized with a fluorescent microscope (Leica; Luigs & Neumann), equipped with a CCD camera (Lumenera), and were then explicitly targeted. Intracellular patch-clamp electrodes were advanced to the neurons using a micromanipulator (Luigs & Neumann) while applying constant positive pressure. Intracellular signals were amplified with a MultiClamp 700B intracellular amplifier (Molecular Devices). All Purkinje cells were clamped at −70 mV or 0 mV throughout the voltage-clamp recordings. All experiments were performed at room temperature (23 °C). The resting membrane potential (RMP) was measured in the absence of any bias current. All other electrical properties quantified after the application of a bias hyperpolarization current to eliminate the spontaneous activity of the Purkinje cells. For dual whole-cell recordings of Purkinje cells, two patch-clamp electrodes were advanced from opposite directions into the cerebellum. Multiple short-duration suprathreshold current pulses were used to stimulate the presynaptic Purkinje cell and record IPSPs in the postsynaptic Purkinje cell (52). All dual whole-cell recordings were presented as net IPSPs from the motor nerves running through the intermyotomal clefts at the tail, where the musculature was left untouched. Activation of the locomotion was induced by extracellular stimulation (using a train of 10 pulses: 1 Hz) applied via a glass pipette placed at the junction between the brain and the spinal cord. To elicit escape, the stimulation electrode delivered three strong pulses (50 Hz) in the same region as for the swimming. All recordings were made from ipsilateral located Purkinje cells and motor nerves.

Electrophysiological Recordings. Adult zebrafish were cold-anesthetized in a slush of a frozen extracellular solution containing MS-222. The scalp and surrounding head tissue were carefully used to expose the cerebellum, and the preparation was transferred to a recording chamber that was continuously perfused with an extracellular solution containing 135.2 mM NaCl, 2.9 mM KCl, 1.2 mM MgCl₂, 2.1 mM CaCl₂, 10 mM Hepes, and 10 mM glucose at pH 7.8 (adjusted with NaOH) and an osmolality of 290 mOsm. For whole-cell intracellular recordings of Purkinje cells in voltage- and current-clamp mode, electrodes (resistance ~15 MΩ) were pulled from borosilicate glass (outer diameter, 1.5 mm; inner diameter, 0.87 mm; Hilgenberg) on a micropipette puller (model P-97, Sutter Instruments) and filled with an intracellular solution containing 120 mM K-gluconate, 5 mM KCl, 10 mM Hepes, 4 mM Mg₂ATP, 0.3 mM Na₃GTP, and 10 mM Na-phosphocreatine at pH 7.4 (adjusted with KOH) and osmolality of 275 mOsm. GFP-positive Purkinje cells were visualized with a fluorescent microscope (Leica; Luigs & Neumann), equipped with a CCD camera (Lumenera), and were then explicitly targeted. Intracellular patch-clamp electrodes were advanced to the neurons using a micromanipulator (Luigs & Neumann) while applying constant positive pressure. Intracellular signals were amplified with a MultiClamp 700B intracellular amplifier (Molecular Devices). All Purkinje cells were clamped at −70 mV or 0 mV throughout the voltage-clamp recordings. All experiments were performed at room temperature (23 °C). The resting membrane potential (RMP) was measured in the absence of any bias current. All other electrical properties quantified after the application of a bias hyperpolarization current to eliminate the spontaneous activity of the Purkinje cells. For dual whole-cell recordings of Purkinje cells, two patch-clamp electrodes were advanced from opposite directions into the cerebellum. Multiple short-duration suprathreshold current pulses were used to stimulate the presynaptic Purkinje cell and record IPSPs in the postsynaptic Purkinje cell (52). All dual whole-cell recordings were presented as net IPSPs from the motor nerves running through the intermyotomal clefts at the tail, where the musculature was left untouched. Activation of the locomotion was induced by extracellular stimulation (using a train of 10 pulses: 1 Hz) applied via a glass pipette placed at the junction between the brain and the spinal cord. To elicit escape, the stimulation electrode delivered three strong pulses (50 Hz) in the same region as for the swimming. All recordings were made from ipsilateral located Purkinje cells and motor nerves.

Morphological Reconstructions. In some experiments during intracellular whole-cell recordings, Purkinje cells were passively filled with neurobiotin tracer (0.5–1%, Vector Labs, SP-1120) to reveal their morphological characteristics post hoc. After the electrophysiological evaluation of the Purkinje cell properties, the brain was kept for an additional 15–20 min in the recording chamber to allow the complete diffusion of the tracer. The brain was then removed from the recording chamber, thoroughly washed with PBS and fixed in 4% PFA overnight at 4 °C. Tissue was then incubated with the following: 7.4 mM KCl, 1.2 mM MgCl₂, 2.1 mM CaCl₂, 10 mM Hepes, and 10 mM glucose during fictive locomotion, we modified the adult zebrafish ex vivo preparation previously described (42). Briefly, zebrafish were cold anesthetized in a slush of a frozen extracellular solution, then the scalp and adjacent tissue were gently removed to expose the brain and the cerebellum. The skin was gently removed to reveal all axial musculature. The epiaxial musculature was carefully removed up to the caudal end of the dorsal fin, leaving the musculature of the tail region intact. Extracellular recordings were performed from the motor nerves running through the intermyotomal clefts at the tail, where the musculature was left untouched. Activation of the locomotion was induced by extracellular stimulation (using a train of 10 pulses: 1 Hz) applied via a glass pipette placed at the junction between the brain and the spinal cord. To elicit escape, the stimulation electrode delivered three strong pulses (50 Hz) in the same region as for the swimming. All recordings were made from ipsilateral located Purkinje cells and motor nerves.

Behavioral Experiments. For the open field test, each animal was placed in 140-mm Petri dishes and allowed to swim freely. Animals were recorded for 5 min after 1 min of adaptation at 30 frames per second with a digital camera. Their swimming behavior was analyzed by using MTrack2 ImageJ plugin and

**Materials and Methods**

**Animal Model.** All animals were raised and kept in a core zebrafish facility at the Karolinska Institute in accordance with established practices. Adult zebrafish (Danio rerio; n = 294 animals; 8–10 wk old; length: 15–20 mm; weight: 0.04–0.06 g; both sexes) WT (AB/Tübingen), and transgenic Tg(CaBE:EGFP, 29,30) and Tg(aldoa:BoTx-GFP) lines were used. No selection criteria and biasing procedures were used to allocate zebrafish to any experimental group. The local Animal Research Ethical Committee approved all experimental protocols, Stockholm (Ethical permit no. 9248–2017), and were implemented following European Union guidelines for the care and use of laboratory animals (2010/63/EU). All efforts were made to utilize only the minimum number of experimental animals necessary to obtain reliable scientific data.

**Immunohistochemistry.** All animals were deeply anesthetized with tricaine methane sulfonate (MS-222, Sigma-Aldrich, E10521). The brains were then extracted and fixed in 4% paraformaldehyde (PFA) and 5% saturated picric acid (Sigma-Aldrich, P6744) in phosphate buffered saline (PBS) (0.01 M; pH: 7.4–7.8) overnight at 4 °C. The brains were then stored in PBS and fixed in 4% PFA overnight at 4 °C. Tissue was then incubated with the following: 7.4 mM KCl, 1.2 mM MgCl₂, 2.1 mM CaCl₂, 10 mM Hepes, and 10 mM glucose during fictive locomotion, we modified the adult zebrafish ex vivo preparation previously described (42). Briefly, zebrafish were cold anesthetized in a slush of a frozen extracellular solution, then the scalp and adjacent tissue were gently removed to expose the brain and the cerebellum. The skin was gently removed to reveal all axial musculature. The epiaxial musculature was carefully removed up to the caudal end of the dorsal fin, leaving the musculature of the tail region intact. Extracellular recordings were performed from the motor nerves running through the intermyotomal clefts at the tail, where the musculature was left untouched. Activation of the locomotion was induced by extracellular stimulation (using a train of 10 pulses: 1 Hz) applied via a glass pipette placed at the junction between the brain and the spinal cord. To elicit escape, the stimulation electrode delivered three strong pulses (50 Hz) in the same region as for the swimming. All recordings were made from ipsilateral located Purkinje cells and motor nerves.

**Fictive Locomotion.** For the evaluation of the activity of the Purkinje cells during fictive locomotion, we modified the adult zebrafish ex vivo prepara-

**Behavioral Experiments.** For the open field test, each animal was placed in 140-mm Petri dishes and allowed to swim freely. Animals were recorded for 5 min after 1 min of adaptation at 30 frames per second with a digital camera. Their swimming behavior was analyzed by using MTrack2 ImageJ plugin and

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employing the t-SNE (34) using the R package "Rtsne" with the following settings: pca_scale = T, perplexity = 5, max_iter = 2000. Hierarchical clustering across all cell types was performed using the 13 cellular properties, with the priori assumption that specific types of neurons existed in our sample. Using R, we first normalized and scaled the data. Subsequently, we calculated the distance matrix with the method "euclidean" and performed hierarchical clustering (hclust) with the "average" algorithm.

**Statistics.** The significance of differences between the means in experimental groups and conditions was analyzed using parametric tests such as the two-tailed unpaired or paired Student’s t-test, one-way ANOVA (ordinary) followed by post hoc Tukey’s test, or two-way ANOVA (repeat measures) followed by Sidak’s comparison test, using Prism (GraphPad Software Inc.). The q values indicated in all figures are as follows: *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. All data are presented as mean ± SEM or as box plots and violin plots showing the median, 25th, and 75th percentile (box and line), and minimal and maximal values (whiskers). Finally, the n values indicate the final number of validated animals per group, cells, or events that were evaluated.

**Data and Code Availability.** Further information and requests for data, resources, and reagents should be directed to and will be fulfilled by K.A. Raw data and R scripts used in this study for dimensionality reduction and clustering of the Purkinje cells are available at https://github.com/steveniagianicolellozebrafish and http://dx.doi.org/10.7939/r2zz7xvkw/z.

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**Dimensionality Reduction and Clustering.** We reduced the dimensionality of the cellular parameters obtained from the recorded cells (n = 64 neurons) by

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