Sensory and central contributions to motor pattern generation in a spiking, neuro-mechanical model of the salamander spinal cord

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Abstract

This study introduces a novel neuromechanical model employing a detailed spiking neural network to explore the role of axial proprioceptive sensory feedback in salamander locomotion. Unlike previous studies that often oversimplified the dynamics of the locomotor networks, our model includes detailed simulations of the classes of neurons that are considered responsible for generating movement patterns. The locomotor circuits, modeled as a spiking neural network of adaptive leaky integrate-and-fire neurons, are coupled to a three-dimensional mechanical model of a salamander with realistic physical parameters and simulated muscles. In open-loop simulations (i.e., without sensory feedback) the model accurately replicates locomotor patterns observed in-vitro and in-vivo for swimming and trotting gaits. Additionally, a modular architecture of the descending reticulospinal (RS) drive to the central pattern generation (CPG) network, allows to accurately control the activation, frequency and phase relationship of the different sections of the limb and axial circuits. In closed-loop simulations (i.e. with the inclusion of axial proprioceptive sensory feedback), systematic evaluations reveal that intermediate values of feedback strength significantly enhance the locomotor efficiency and robustness to disturbances during swimming. Specifically, our results show that sensory feedback increases the tail beat frequency and reduces the intersegmental phase lag, contributing to more coordinated and faster movement patterns. Moreover, the presence of feedback expanded the stability region of the closed-loop swimming network, enhancing tolerance to a wider range of external stimulations, internal parameters' modulation and noise levels. This study provides new insights into the complex interplay between central and peripheral pattern generation mechanisms, offering potential strategies for developing advanced biomimetic robots. Additionally, this study underscores the critical role of detailed, biologically-realistic neural networks to improve our understanding of vertebrate locomotion.

Author summary

In this paper, we developed a computational model to investigate how salamanders move, both while swimming and walking. Unlike previous studies that often oversimplified the dynamics of these complex neural networks, our model includes detailed simulations of the classes of neurons that are considered responsible for generating movement patterns. The locomotor circuits, modeled as a spiking neural network, are coupled to a three-dimensional mechanical model of a salamander with realistic physical parameters and simulated muscles. The neural model integrates axial proprioceptive sensory feedback from the body's movements to modulate the locomotor gaits. Our simulations suggest that this sensory feedback plays a major role in controlling the rhythm and coordination of movements. This has implications for understanding not only how salamanders move but also provides insights into the evolution of locomotion in vertebrates. By investigating how central and sensory mechanisms interact to produce efficient and adaptable movement, our work contributes to the broader field of neuroscience and robotics, offering potential strategies for designing more effective biomimetic robots.

Introduction

Animal locomotion depends on the coordinated interplay between body mechanics and the neural circuits in the brain, spinal cord, and sensory systems. Decades of experiments and modeling studies have established that specialised spinal circuits called central pattern generators (CPGs) are capable of generating rhythmic outputs underlying motor skills, such as walking and swimming, in the absence of rhythmic inputs (for a review, see [1]). The numerous types of CPG circuits and their respective locomotion patterns were described across multiple vertebrate species, such as swimming in the lamprey [2–5], zebrafish [6–8] and tadpole [9,10], walking and swimming in the salamander [11,12], as well as walking/running in the mouse [13–15] and cat [16,17].

The basic repertoire of motor outputs and several anatomical and genetic properties of CPG, brain stem and sensory circuits appear to be largely conserved across vertebrate species [1, 12, 18–22]. However, evolution also gave rise to specialized CPG and sensorimotor circuits that allowed new species to adapt to their habitat and to meet the specific needs of each species. A general view is that CPG circuits have been genetically hard coded during evolution to generate the central "stereotypical" motor programs which allow actions to be executed, while sensory feedback loops fine tune these programs and contribute to select new motor programs [23]. However, several studies suggest that sensory feedback can have a bigger role than expected, by contributing to the generation of rhythmic motor outputs in different spinal body parts and to their coordination, especially in large and slow animals [24–28]. Thus it has become increasingly clear that understanding animal locomotion requires studying the interaction between the central and sensory feedback systems in closed loop.

Recent advances in molecular genomics and optogenetics - mostly in the mouse and the zebrafish - combined with traditional electrophysiological and anatomical experiments have identified anatomically and functionally distinct reticulospinal and spinal CPG neuronal populations, which are shared across these and other vertebrate species [5,22,29]. Reticulospinal neurons (RS) in the brain stem relay motor commands from the mesencephalic locomotor region (MLR) to the spinal interneurons to trigger/terminate a locomotion episode, control the steering, frequency and speed of locomotion and change the expressed locomotor pattern [30–36]. Similarly, in the salamander, low level stimulation in the MLR evokes slow walking, while high level stimulation evokes swimming at higher movement frequencies [37]. Genetically g

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identified classes of CPG interneurons are responsible for the coordination of different body parts during locomotion [22]. For example, in the zebrafish and tadpoles, chains of commissural glycinergic V0d neurons and ipsilateral glutamatergic CHx10 positive V2a (called dINs in tadpoles) neurons are responsible for the generation of the alternating left-right rhythm and forward wave propagation during swimming. More complex circuits consisting of multiple cell types are responsible for the coordination between limb CPGs during quadrupedal gaits in the mouse [15, 38–40].

In addition to brainstem and CPG circuits, spinal cord circuits are equipped with sensory feedback loops originating from muscles, cutaneous afferents and intraspinal sensory neurons that modify the CPG locomotor patterns according to the interaction with the external world [23]. Proprioceptive (curvature) sensory neurons (PS) in the lamprey and the zebrafish detect stretch and stretch rate within the internal region of the spinal cord [41–44]). Interestingly, while PS neurons in the lamprey (also called edge cells) are made up of distinct excitatory ipsilateral and inhibitory commissural populations [41,45], PS neurons in the zebrafish are exclusively commissural and inhibitory and connect preferentially to V2a interneurons [44]. In both animals PS neurons were shown to increase the tail beat frequency [44,46,47] and decrease the intersegmental phase lag [44,47]. This observation suggests that different feedback topologies may allow to implement common functions to achieve the generation of an efficient swimming behavior [48].

Following a spinal cord transection, lampreys and eels can still generate coordinated swimming movements below the lesion site despite the loss of descending signals [46, 49, 50]. This suggests that sensory feedback can replace the missing RS excitability and CPG coupling to allow for the generation of rhythmic and coordinated outputs. This hypothesis was successfully tested in computational and robotic studies of the lamprey's swimming CPG network with proprioceptive [51] and/or lateral hydrodynamic force [52] sensing. Unlike lampreys and eels, salamanders do not display coordinated swimming immediately after spinal cord injuries [53, 54]. This is likely due to a lower level of excitability of the spinal circuits below the lesion, that consequently require the regrowth of descending signals to restore locomotion [53, 55]. Nonetheless, salamanders can regenerate their spinal cord and display swimming within 6-8 weeks after the lesion. Interestingly, sensory stimulation was shown to significantly enhance the recovery capability in spinally-transected eels [56].

During terrestrial locomotion in the cat, limb proprioceptive sensory feedback appears to have a role at different phases of the locomotor cycle, from initiating, blocking, entraining and resetting the rhythm in various types of afferents (for a review, see [23]). In the mouse, it regulates the rhythm of the alternating extensor-flexor muscle activity at the individual limb joints level and the intra-limb coordination [57].

Salamanders are unique animals to study how CPGs and sensory feedback circuits have evolved from fishes to mammals, and to explore the relative contribution of CPG and sensorimotor circuits during aquatic and terrestrial locomotion. Like other amphibians, salamanders appeared on Earth around 360 million years ago, sometime between the arrival of bony and jawless fishes (i.e. lamprey) and that of mammals [22, 58, 59]. Salamanders are close extant representatives of the first tetrapods that evolved from water to land. They swim in water moving the tail in an undulatory fashion reminiscent to lamprey's swimming with a characteristic rostrocaudally propagating traveling wave of motoneuron activity in alternation between the left and right sides, and walk on land using lateral sequence walking or trotting gaits ressembling those described in mammals [15,60–63]. Salamanders are also interesting animals for their extraordinary regenerating abilities. Indeed they are the only tetrapods that can restore spinal circuits and their function after spinal cord lesions [54,55,64,65]. Some species of salamanders can regrow missing limbs [66] and

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Fig 1. Neural model. A) shows the schematics of the locomotor network organization. The axial network is comprised of two chains of oscillators (i.e., the hemisegments), divided into 40 equivalent segments (only a fraction is shown). The axial segments are spatially organized in trunk and tail sub-networks. In correspondence of the two girdles (green oscillator symbols) there are the equivalent segments for the limb networks. Each segment projects to a pair of antagonist muscle cells. The axial network receives proprioceptive sensory feedback (PS) based on the local curvature of the model. The limb network does not receive sensory feedback. \mathbf{B}) shows the organization of the RS population. The axial reticulospinal neurons (RS) are divided into 6 sub modules (3 for the trunk, 3 for the tail) projecting to the axial CPG neurons according to overlapping trapezoidal distributions, shown as dotted lines. The limb RS neurons are divided into 4 modules, each projecting to a separate limb segment. The limb segments project to the axial CPG network with local descending distributions. C) shows the organization of each equivalent segment of the CPG network. The structure comprises excitatory (EN) and inhibitory (IN) CPG neurons, motoneurons (MN), excitatory (PS_{ex}) and inhibitory (PS_{in}) proprioceptive sensory neurons and muscle cells (MC). Only a fraction of the neurons and their connections originating from the right hemisegment are shown. The connections from the left hemisegment follow the same organization principle. The range of the ascending and descending projections is displayed in terms of equivalent segments (seg). Note that the projections from EN and IN populations are dominantly descending. **D**) shows the limb-to-body connectivity pattern. Each flexor (i.e. protractor) hemisegment of a limb oscillator excites the contralateral axial hemisegmens segments and inhibits the ipsilateral axial hemisegments. \mathbf{E}) shows the inter-limb connectivity scheme. Each limb oscillator sends inhibitory projections from the flexor side to the flexor side of the commissural and ipsilateral limbs. The commissural connections (thick lines) are stronger than the ipsilateral projections (dotted lines) to ensure the left-rial ternation between opposed for limbs and hindlimbs.

even parts of the brain tissue [67]. Investigating how salamanders re-configure their nervous system to restore motor functions could help us better understand the motor recovery from spinal cord injury also in mammals [68, 69].

Previous simulation studies on salamander locomotor networks have for the most part used high levels of abstraction (e.g., abstract oscillators) for the modeling of the CPG dynamics [70–73]. These models allow one to investigate of the core principles

underlying the coordination between axial segments and they are suitable for neuromechanical simulations [71, 74] and robotics studies [62, 73]. On the other hand, abstract models do not allow the investigation of the core rhythmogenic components of real neural networks. Similarly, the effect of sensory feedback on these models can only be studied through abstract mathematical rules that are difficult to compare to experimental findings [75, 76]. In the lamprey, many studies have focused on bridging this gap with detailed bio-physical models (e.g. Hodgkin–Huxley neurons [77]) of the locomotor networks [4,78–81]. Taking inspiration from the lamprey, similar efforts have been done also for the salamander locomotor circuits [82]. Unfortunately, these models 100 are not suitable for an extensive use in neuromechanical simulations due to their 101 excessive computational cost. Additionally, the high-dimensionality of these models 102 calls for an amount of data for parameter instantiation that is not currently available 103 for most animal models. For these reasons, several efforts have been made to use an 104 intermediate level of abstraction to study the salamander locomotor networks [83,84]. 105 In these models, the circuits are simulated as networks of spiking neurons modeled with 106 reduced two-dimensional formulations including membrane potential and an adaptation 107 variable (e.g. adaptive integrate-and-fire [85], Izhikevich [86], Fitzhugh-Nagumo [87]). 108 These models allow to grasp the core rhythmogenic mechanisms of the locomotor 109 networks while maintaining a low computational complexity [88]. Moreover, these 110 formulations enable the creation of closed-loop neuromechanical simulations, in which 111 sensory feedback is integrated into the networks in a biologically-realistic manner [12]. 112 In [83], this approach was used to investigate the role of axial and limb proprioceptive 113 feedback during walking gaits in salamanders. Interestingly, this work showed that 114 sensory feedback played a significant role in obtaining a coordinated lateral sequence 115 walking gait, as well as transitions from walking to trotting. On the other hand, the 116 work focused solely on ground locomotion and it did not investigate swimming, nor the 117 effect of modulating the strength and/or topology of the axial PS feedback. 118

Indeed, despite the advancements in simulating the salamander's locomotor circuits at different levels of abstraction, a full understanding of the relative contribution of CPG and axial PS feedback has so far remained elusive. In the present work, we aim to investigate this problem with a neuromechanical model of the salamander's locomotor network (see Fig 1) based on adaptive leaky integrate-and-fire spiking neurons (i.e., with an intermediate level of abstraction) [85]. The model was built upon previous studies [82, 89, 90] and it was refined to include new findings from salamander and other species. Additionally, the spiking neural network was coupled to a simulated salamander body with realistic physical parameters in order to provide axial PS feedback to the network and to evaluate the resulting locomotor performance (see Fig 2). The main questions addressed by the paper can be summed up as

- 1. How can a spiking neural network replicate the salamander locomotor circuits to generate both swimming and walking patterns with a unified architecture?
- 2. How does axial PS feedback influence the performance of the model in terms of frequency, intersegmental phase lag, forward speed and locomotor efficiency?
- 3. What is the effect of varying the topology of axial PS feedback on the rhythmic motor patterns generated by the model?
- 4. How does the feedback affect the model's robustness to noise and its ability to recover stable locomotor rhythms in the presence of disturbances?
- 5. Can axial PS feedback extend the functional range of the CPG network by 138 enhancing its tolerance to the variability of the external drive and/or the internal 139 parameters? 140

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The spinal neural network consists of spiking neurons of excitatory and inhibitory 141 types that encompass different classes of neuronal populations [12]. These neuronal 142 classes, which were genetically identified in the mouse and zebrafish [1, 22], could indeed 143 be present in salamanders based on recent comparative genome analysis and 144 physiological results [12,91]. The topology of the synaptic connectivity was constrained 145 using indirect evidence from neuronal and kinematic recordings of *Pleurodeles waltl.* and 146 by taking inspiration from that of the lamprey and zebrafish [12, 22, 82]. The 3D 147 mechanical salamander model (see Fig 2) consists of multiple connected rigid links that 148 faithfully resemble the body of the real salamander and it is actuated by virtual 149 muscles [92]. This actuation allows the model to move in simulated water or a flat 150 terrain to reproduce swimming and walking, respectively. This model is simulated in 151 the MuJoCo physics engine [93] through the FARMS framework [92]. Physical and 152 muscles parameters were optimised to match the swimming kinematics of *Pleurodeles* 153 waltl salamanders using a novel system design based on the transfer function of the 154 linearized muscle model response (more details in S3 File). For validation, the 155 kinematics reported from *Pleurodeles waltl* in [62,94] was replicated and matched (see 156 S4 File). The axial PS neurons received input from the local curvature of the simulated 157 salamander body. Their activation lead to direct mono-synaptic effects in the axial 158 CPG and MN neurons, thus directly affecting the whole locomotor network. 159

In summary, we proposed a new spiking neuromechanical model of the salamander 160 swimming and walking networks to investigate the contributions of CPG and axial 161 stretch feedback in the locomotor circuits. In open loop (i.e., when simulating the 162 spiking neural network in isolation) the model could replicate observations from 163 previous experimental and simulation studies [95,96]. The rhythmogenic principle of the 164 network was investigated in simulation and it was shown to be strongly dependent on 165 commissural inhibition according to an escape-from-inhibition mechanism [97,98]. In 166 closed loop, the spiking network was coupled to a simulated salamander body to receive 167 axial PS feedback. In this configuration, we found that the effect of stretch feedback 168 was largely conserved across different biologically-realistic topologies. Indeed, all the 169 considered PS connectivities led to an increase of the frequency and a decrease of the 170 intersegmental phase lag of the CPG oscillations. Additionally, we proposed a 171 mechanistic explanation of the reported sensory feedback effect. Interestingly, the 172 presence of sensory feedback largely expanded the stability region of the closed-loop 173 network with respect to the level of external stimulation, noise and strength of 174 commissural inhibition. Additionally, intermediate values stretch feedback strength 175 boosted the salamander's frequency, speed and locomotion efficiency with respect to the 176 pure open loop paradigm, while maintaining a flexible control over frequency and phase 177 lag. On the other hand, extreme levels of sensory feedback strength led to a loss in 178 controllability of the network (in particular, a reduced range of frequencies, phase lags 179 and speeds) and a switch to feedback-based rhythm generation. Overall, the proposed 180 model helped in shining a light on the interplay between central and peripheral pattern 181 generation in the salamander locomotor networks. 182

Results and discussion

Open loop swimming

Swimming network

The locomotor network underlying swimming behaviors can be dynamically activated through the targeted stimulation of the RS populations projecting to the axial CPG network (see Fig 3A). Upon stimulation, the RS convey a descending excitatory signal that initiates rhythmic motor patterns. The ensuing activity within the axial CPGs 189

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Fig 2. Mechanical model. A) shows the exchange of information between the neural and mechanical model. The neural model sends torque commands to the joints and reads back the corresponding joint angles. B) shows the active joints (in red) acting about the horizontal plane and the passive joints (in blue) acting about the sagittal plane. \mathbf{C}) shows the active degrees of freedom of each limb. The shoulder and hip joints are spherical joints allowing rotations about the three axes. The elbow and knee joints are hinge joints that allows flexion/extension. \mathbf{D}) displays the equivalent scheme of an Ekeberg muscle model. It is a rotational spring-damper system with the addition of a variable stiffness term $\beta(M_L + M_R)\theta^*$. The active term of the model acts an external torque $\alpha(M_L - M_R)$. **E**) shows the equivalent scheme for the mechano-electrical transduction principle. The threshold angle θ_{TH} indicates the critical angle for which the PS neurons become active. The feedback synaptic weight ω_{PS} modulates the strength W (i.e., the slope) of the PS response once θ_{TH} is crossed. F) displays the control diagram for the mechano-electrical and electro-mechanical transduction. The axial CPG receives curvature information through the PS population. The activity of the CPG is translated into torque information through the Ekeberg muscle model via motoneurons (MN) and muscle cells (MC).

manifests as a left-right alternating travelling wave of neuronal activity which closely resembles the motor patterns observed both in-vitro and in-vivo in salamanders (see Fig 3A1). The swimming activity generated by the open-loop network when linked to the mechanical model is displayed in S2 Video.

The frequency (f_{neur}) of the generated motor patterns can be precisely modulated by adjusting the level of excitatory drive directed toward the RS neurons (see Fig 3A2). Moreover, the total wave lag (TWL), see Eq 3 in S1 File) of the network can be modulated by adjusting the relative drive to the rostral or caudal components of the network. Specifically, augmenting the drive to the rostral RS tends to increase the TWL (see Fig 3A2). Conversely, reducing the drive to these neurons (or increasing the drive to caudal-projecting ones) results in a reduction of the TWL. This adjustment of the delay in phase synchronization supports the theoretical basis proposed by the trailing oscillator hypothesis [99]. It emphasizes the intricate control over the rhythm



Fig 3. Open loop swimming network. A) shows the swimming pattern emerging from three different levels of stimulation to the axial RS. The level of stimulation provided to the 10 RS modules (6 axial, 4 limbs) is shown on top of each plot. The raster plots represent the spike times of excitatory neurons (EN) and reticulospinal neurons (RS). For the axial network (AX), only the activity of the left side is displayed. For the limb network (LB), only the activity of the flexor side is displayed. Limbs are ordered as left forelimb (LF), right forelimb (RF), left hindlimb (LH), right hindlimb (RH), from top to bottom. A1 shows the pattern obtained with a constant drive of 5.0pA. The neural activity corresponds to $f_{neur} = 2.1Hz$ and TWL = 1.04. A2 shows the response to a higher constant drive of 5.7pA. The neural activity corresponds to $f_{neur} = 3.1Hz$ and TWL = 1.27. A3 shows the effect of driving the network with a gradient of excitation ranging from 5.7pA (to the rostral RS neurons) to 5.0pA (to the tail RS neurons). The neural activity corresponds to $f_{neur} = 3.1Hz$ and TWL = 1.83. **B**) shows the coordination patterns achieved for different levels of stimulation to the trunk and tail regions. B1 shows the activity corresponding to a stimulation of 5.7pA of the trunk RS. The trunk network is rhythmically active, but the activity does not spread to the tail network. B2 shows the activity corresponding to a stimulation of 5.7pA of the tail RS. The tail network is rhythmically active, but the activity does not spread to the trunk network. B3 shows the activity corresponding to a different level stimulation of the trunk (5.0pA) and tail (6.2pA) RS neurons. Both regions are rhythmically active and the tail network oscillates at a higher frequency with a 2:1 ratio with respect to the trunk network.

and coordination enabled by the modular RS drive [100].

Interestingly, the trunk and tail networks could also be independently controlled thanks to the modular RS organization and the discontinuity within the CPG connections in correspondence of the hip girdle. A targeted stimulation of the trunk RS neurons leads to the generation of a travelling wave of activity in the trunk CPG

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neurons, while the tail network remains silent (see Fig 3B1). Similarly, stimulating the tail RS neurons leads to a rhythmic activity isolated to the tail network (see Fig 3B2). The trunk and tail networks could also be simultaneously activated while displaying non-synchronized oscillations. In Fig 3B3, an increased drive to the tail RS population leads to higher-frequency oscillations within the tail CPG network with a 2:1 ratio with respect to the trunk section. The independent control of the trunk and tail sections resonates with experimental findings from the salamander locomotor networks [101].

Overall, the ability to differentially excite distinct axial regions through modulated 215 RS drive represents a straightforward yet potent mechanism to control the axial CPGs. 216 This level of control underscores the integral role that the reticulospinal pathways could 217 have in tuning the spatiotemporal dynamics of locomotor networks, facilitating adaptive 218 motor outputs in response to varied environmental demands. Nonetheless, unless stated 219 otherwise, the following analyses on the swimming network imposed a constant level of 220 stimulation amplitude to all the axial RS populations in order to better highlight the 221 key parameters affecting the model performance in open and closed loop. 222

The network shows a triphasic response to the stimulation level

The response of the open-loop network was studied across varying stimulation values targeting the axial RS neurons. The stimulation amplitude was varied in the range [4.0, 8.0]pA for 20 different instances of the network, with different random seeds. The results are reported in terms of mean and standard deviation of the corresponding computed metrics.

The analysis revealed a triphasic response of the network within the range of 229 considered stimulation amplitudes (see Fig 4A). Notably, when the stimulation 230 amplitude was lower than the rheobase current of the RS neurons (equal to 4.0pA on 231 average), the CPG network did not receive any descending drive and thus remained 232 silent. For stimulation values close to the rheobase current, the network displayed 233 aperiodic patterns and disorganized oscillations. This resulted in low PTCC values (see 234 Eq 4 in S1 File), indicating diminished network rhythmicity (see Fig 4B, top panel). For 235 intermediate stimulation values, the network manifested traveling waves of coordinated 236 activity alongside a distinct left-right alternation (see Fig 4B, middle panel), analogous 237 to the motor patterns observed during swimming [12]. This phase represents a 238 functional regime wherein the network accurately replicates rhythmic motor outputs 239 associated with swimming. At higher stimulation values, the network deviated from 240 oscillatory behavior, with both hemisegments exhibiting sustained tonic activity (see 241 Fig 4B, bottom panel). In this phase, the combined action of commissural inhibition 242 and neuronal adaptation was insufficient to counteract the persistent excitation induced 243 by the elevated background drive, resulting in a loss of rhythmic oscillations across the 244 hemisegments. Note that, in this regime, the network loses the capability to produce 245 stable rhythms (i.e.; $PTCC \approx 0$) and the corresponding f_{neur} and TWL metrics are 246 therefore unreliable. 247

Within the oscillatory regime delineated by intermediate stimulation levels, a linear 248 relationship emerged between the stimulation drive and the frequency of oscillations (see 249 Fig 4A, middle panel). Additionally, the frequency of oscillations varied in the range 250 [1.0, 4.0] Hz, aligning with empirical findings from salamander studies involving MLR 251 stimulation [37]. Conversely, the total wave lag (TWL), see Eq. 3 in S1 File), initially 252 increased with rising drive, plateauing around a value of 1.2 (see Fig 4A, bottom panel). 253 This TWL value was in the lower range of those reported for the swimming kinematics 254 in the Ambystoma Mexicanum (ranging from 1.2 to 2.1 in [102]). Conversely, the TWL255 for the epaxial EMG activity during swimming in *Pleurodeles waltl* was markedly lower 256 (0.5 in [60], 0.74 in [96]). Notably, these experiments were carried out on behaving 257 animals naturally receiving sensory feedback during locomotion. Interestingly, isolated 258

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Fig 4. The triphasic response to the stimulation amplitude to the axial RS **neurons.** A) displays the effect of the stimulation amplitude provided to the axial RS neurons, on the open loop swimming network. The stimulation was varied in the range [4.0, 8.0]pA. From top to bottom, the plots show the relationship between the level of stimulation and the periodicity (PTCC), frequency (f_{neur}) and total wave lag (TWL)of the neural signals, respectively. In each plot, the thick line represents the average metric computed across 20 different instances of the network. The shaded area denotes one standard deviation of distance from the mean. Note that, for $PTCC \approx 0$, the network loses its capability to produce stable rhythms and the values of f_{neur} and TWL are therefore unreliable. **B**) shows the response of the network for three different drive amplitudes to the axial RS neurons (equal to 4.25pA, 5.75pA and 7.5pA, respectively). The raster plots represent the spike times of excitatory neurons (EN) and reticulospinal neurons (RS). For the axial network (AX), only the activity of the left side is displayed. For the limb network (LB), only the activity of the flexor side is displayed. For low stimulation values (blue dot), the network displays unpatterned activations. For intermediate stimulation values (orange star), the network displays travelling waves of left-right activity. For high stimulation values (red square), the network displays tonic activity.

Pleurodeles waltl spinal cord preparations were shown to display much higher TWL values when stimulated with NMDA baths (up to 2.5 in [95]). As it will be shown in the following sections, the introduction of sensory feedback can lower the TWL values to match the experimental findings. This result also provides a possible explanation for the mismatch between isolated and intact spinal cord recordings.

The observed drive-frequency relationship parallels the findings from prior simulation studies conducted at higher abstraction levels, with coupled nonlinear

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oscillator models [70]. This consistency strongly suggests that for the salamander the assumptions of models at higher levels of abstraction are well justified through a chain of models, ranging from detailed bio-physical models (e.g., [82]) through formal spiking models (e.g., [90] and present model), all the way to more abstract controllers [70, 73]. Unless stated otherwise, the following sections impose a stimulation amplitude of 5.7pA to the axial RS neurons. With the selected drive, the network reliably generates stable oscillations with an average frequency of 3.1Hz and a total wave lag of 1.27BL/cycle.

The open loop network is inherently robust to external disturbances

The swimming network was simulated with different noise levels to investigate its robustness (see Fig 5A). The noise amplitude varied in the range [0.0, 10.0]mV and each simulation was carried out for 20 instances of the network, with different random seeds. It was found that the axial CPG network could tolerate high levels of noise without significantly altering the resulting pattern (see Fig 5B). Indeed, despite a high noise level of 7mV, the network could generate a coordinated swimming pattern that resembled the one without external disturbance from Fig 4C.

This robustness is likely due to the diversity in the neuronal parameters of the CPG populations, as suggested by [4]. The diverse neuronal parameters may provide a form of network heterogeneity that helps buffer against the disruptions caused by noise, ensuring functional outputs are maintained under various conditions. However, there is a limit to this tolerance. When noise levels were exceedingly high (i.e., greater than 8mV), a noticeable decline in the rhythmic pattern was observed, leading to a cessation of oscillatory activity and, consequently, low values of PTCC (see Fig 5A).

The findings from this simulation highlight the axial CPG network's robust nature in handling noise and also outline the boundaries of this robustness.

The network is highly sensitive to a modulation of the commissural inhibition strength

The analysis pointed out the significant role of commissural inhibitory connections in modulating rhythmicity (PTCC), total wave lag (TWL), and frequency (f_{neur}) of the network's locomotor output (see Fig 5C). The figure reports the distribution of the total order sensitivity indices computed across 20 instances of the network, with different random seeds. The indices were computed for all the neuronal and synaptic parameters with respect to the periodicity, total wave lag and frequency of the neural activity. Indeed, for all the considered metrics, the synaptic weight of inhibitory connections (Δg_{glyc}) resulted in the highest total-order sensitivity index among all neuronal and synaptic parameters. The result was particularly evident for f_{neur} , where Δg_{glyc} showed a very strong modulatory effect. The nature of the modulation of f_{neur} , TWLand PTCC, in terms of increase or decrease of the corresponding metrics, is discussed in detail in the next section.

This analysis sheds light on the critical role of commissural inhibitory connections as major modulators of network rhythmicity and locomotor tempo. The results support the escape-from-inhibition hypothesis [97,98], where rhythmic locomotor patterns arise from a balance of excitatory and inhibitory interactions within the network.

The commissural inhibition governs the rhythmogenesis, frequency and intersegmental phase lag of the swimming patterns

In the previous section, we discussed the critical role of the CPG commissural inhibition in generating patterned locomotor activity. It notably affects the frequency, phase lag, and periodicity of the network, making it crucial to understand the range of inhibition

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Fig 5. The network's inherent robustness to noise and sensitivity analysis. **A**) displays the response of the swimming network to different values of noise level, which was varied in the range [0.0, 10.0]mV. The thick line indicates the average PTCC value over 20 instances of the network. The shaded area around the curve indicates the region within one standard deviation of distance from the mean. \mathbf{B}) shows the response of the network for a noise level of 7mV. The raster plots represent the spike times of excitatory neurons (EN) and reticulospinal neurons (RS). For the axial network (AX), only the activity of the left side is displayed. For the limb network (LB), only the activity of the flexor side is displayed. C) shows the results of the sensitivity analysis performed on the open loop network. The x-axis lists the neuronal and synaptic parameters changed during the analysis. The y-axis shows the distribution of the corresponding total-order sensitivity indices for 30 different instances of the network. The three plots show the sensitivity indices with respect to the periodicity (PTCC), total wave lag (TWL) and frequency (f_{neur}) , respectively. It can be noted that the weight of inhibitory connections Δg_{qlyc} is the parameter with the highest influence on the considered metrics.

strengths that allow rhythm generation. We conducted an analysis by varying the strength of commissural inhibition between hemisegments and studying the corresponding periodicity (PTCC), frequency (f_{neur}) and total wave lag (TWL) values displayed by the network (see Fig 6A). The commissural inhibition strength weight was varied in the range [0.0, 4.0]. The curves display the average PTCC (in blue), f_{neur} (in red) and TWL (in green) values across 20 instances of the network. The shaded areas highlight the metrics values within one standard deviation from the average. The figure reveals three distinct regions based on inhibition strength (see Fig 6B). At low inhibition strength, the descending drive overpowers the commissural inhibition, leading to bilateral tonic activation in the network (see Fig 6B1). At high inhibition strength, neuronal adaptation is not enough to allow the contralateral side to escape from the inhibition, resulting in a 'winner takes all' configuration where one side becomes

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tonically active while the opposite side remains silent (see Fig 6B3). At intermediate ³²⁵ inhibition strength, the network exhibits stable oscillations (i.e. PTCC > 1.0), as ³²⁶ indicated by the plateau in the value of PTCC (see Fig 6B2). Interestingly, in the ³²⁷ oscillatory regime the frequency of the oscillations decreased monotonically with ³²⁸ increasing values of inhibition strength. Conversely, the TWL showed an initial increase ³²⁹ up to $TWL \approx 1.3BL/cycle$ followed by a sharp decrease to $TWL \approx 0.0BL/cycle$ (i.e. a ³³⁰ standing wave of activity). ³³¹



Fig 6. Commissural inhibition governs rhythm generation, frequency and intersegmental phase lag. A) shows the dependency of periodicity (PTCC, in blue), frequency $(f_{neur}, in red)$ and total wave lag (TWL, in green) on the strength of the CPG commissural inhibition. The inhibition strength was varied in the range [0.0, 4.0]. The thick lines represent the mean across 20 instances of the network. The shaded areas represent one standard deviation of difference from the mean. The values of f_{neur} and TWL are shown only for combinations where the average PTCC was greater than 1.0 (dotted line). **B**) shows the three regimes displayed by the network depending on the level of commissural inhibition strength. Raster plots represent the spike times of excitatory (EN) and reticulospinal (RS) neurons of the axial (AX) and limbs (LB) sub-networks. For each population, the upper half represents the neurons on the left side (L), the lower half represents the neurons on the right side (R). For low inhibition values (B1) both sides of the network are tonically active. For intermediate inhibition values (B2) the network displays rhythmic left-right alternating waves of activation. For high inhibition values (B3) one side of the network is tonically active causing the other one to remain silent ("winner takes all" configuration).

Open loop walking

Walking network

Walking behaviors can be obtained in the simulated network via the activation of the RS modules projecting to the axial and limb CPGs (see Fig 7A1). With a stimulation value of 5.0pA and 4.0pA to the limbs and axial RS populations, respectively, the four equivalent segments displayed rhythmic activation with a frequency of 0.8Hz (see Fig 7A3). Notably, the obtained frequency is lower than the ones obtained for swimming, in accordance with experimental observations [60, 62]. Also, the stimulation amplitude used to achieve this stepping pattern was markedly lower than the one used for swimming, matching experiments involving MLR stimulation in salamanders [37]. The coordination between the limb oscillators is obtained through the inter-limb connectivity. The connectivity enforces the alternation of contralateral limbs via inhibitory commissural inter-limb projections. Additionally, ipsilateral limbs also display anti-phase relationship due to the presence of inhibitory ipsilateral inter-limb connections. These two conditions enforce the simultaneous activation of diagonally opposed limbs. Additionally, given the lower intrinsic frequency of the limb oscillators and their connections to the axial network, the oscillations of the axial CPGs are slowed down and they synchronize to the limb activity. Therefore, the axial network is entrained by the limb oscillators and it displays a phase shift in correspondence with the hip girdle. This pattern of activation resembles the one displayed by salamanders during trotting [62, 73].

Interestingly, the trunk network displayed an average intersegmental phase lag (IPL, see Eq 2) in S1 File) of 0.33%, corresponding to a (quasi) standing wave of activity. Contrarily, the tail network displayed an IPL of 1.62%, corresponding to a travelling wave. This observation contrasts with experimental results from behaving salamanders, where the tail segments also displayed a quasi-synchronous behavior [60, 103]. On the other hand, fictive locomotion experiments showed that the isolated tail network displays travelling waves of activation [101]. Previous simulation studies proposed long-range limb-to-body connections to achieve a standing wave in the tail network [70]. On the other hand, these connectivity patterns were subsequently ruled out based on the observation that limb oscillations could coexist with swimming-like patterns of axial activation during underwater stepping [73, 104]. In this context, sensory feedback could provide a mechanism to synchronize the activity of the trunk and tail oscillators while maintaining the local (and flexible) connections between the limbs and the body.

Similarly to the swimming network, increasing the stimulation to the limbs RS populations to 5.5pA (while maintaining the axial RS drive of 4.0pA) lead to an increase in the oscillation frequency to 1.3Hz (see Fig 7A2), while maintaining the synchronization between diagonally opposed limbs (see Fig 7A4). In principle, different connectivity patterns between the limbs could be achieved by modulating the inter-limb connectivity and/or the descending drive from the RS neurons [39, 105, 106]. Nonetheless, the study of additional walking patterns was outside of the scope of this work and it was not investigated further.

Analogously to the swimming network, the differential stimulation to the trunk and tail RS populations could be exploited to further increase the flexibility of the model. Silencing the drive to the tail RS neurons led to oscillations in the trunk and limb segments that did not spread to the tail CPG section (see Fig 7B1). This pattern corresponds to stepping with a passive tail, a behavior that is observed in behaving salamanders [73, 104]. Contrarily, increasing the stimulation to the tail RS populations could lead to a "double-bursting" in the tail section (see Fig 7B2). This pattern was observed in-vivo in salamanders walking on slippery surfaces [60, 107]. Finally, the activation of the limb CPGs did not necessarily result in a 180* phase shift between the

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trunk and tail oscillations. Indeed, increasing the drive to all the axial RS neurons allowed the axial network to escape the phase-locking imposed by the limbs (see Fig 7B3). In this case, the network displayed rhythmic limb activity coexisting with a full travelling wave of axial activity, analogously to the patterns observed during underwater stepping in salamanders [73,96]. As previously discussed, the coexistence of a travelling wave and limb activity is possible thanks to the local nature of the connections between the limbs and the axis.



Fig 7. Open loop walking network. A) shows the walking pattern emerging from different levels of stimulation to the axial and limb RS. The level of stimulation provided to the 10 RS modules (6 axial, 4 limbs) is shown on top of each plot. The raster plots represent the spike times of every neuron of the spinal cord model. For the axial network (AX), only the activity of the left side is displayed. For the limb network (LB), only the activity of the flexor side is displayed. Limbs are ordered as left forelimb (LF), right forelimb (RF), left hindlimb (LH), right hindlimb (RH), from top to bottom. A1 shows the pattern obtained with a constant drive to the limbs and axial RS with an amplitude of 5.0pA and 4.0pA, respectively. The neural activity corresponds to $f_{neur} = 0.8Hz$. A2 shows the response to a higher constant drive of 5.5pA to the limb RS. The neural activity corresponds to $f_{neur} = 1.3Hz$. A3 and A4 show the limb activity recorded during the experiments in A1 and A2, respectively. \mathbf{B}) shows the coordination patterns achieved when modulating the drive to the axial RS modules. B1 shows the response of the network from A)2 when the tail RS modules do not receive any drive. The tail network remains silent. B2 shows the response of the network from A)2 when the tail RS modules receive an increased drive of 6.5pA. The tail network oscillates at a higher frequency with a 2:1 ratio with respect to the trunk network. B3 shows the response of the network with an axial RS drive of 6.0pA and a limbs RS drive of 4.5pA. The axial network displays a travelling wave of activity despite the rhythmic limb activation.

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Walking in the neuromechanical model

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When linked to the neuromechanical model, the network could reproduce a biologically-realistic trotting gait (see S3 Video). The generated walking had a frequency of 0.9Hz and a stride length of 0.43SVL/cycle, which resembles kinematics data from salamanders [62]. On the other hand, the use of a single equivalent segment for each limb limits the capability of the model to replicate the complex inter-joint coordination patterns displayed during ground locomotion. Additionally, the necessary speed-dependent change in the amplitude of the angles of axial and limbs joints could not be reproduced with constant motor output gains (see G_{MC} in Eq 10). Finally, the current network design does not allow to modulate the duty cycle of the flexion-extension phases of each limb, which were forced to 50% of the period. Overall, these limitations lead to difficulty in controlling the mechanical stability of the model during ground locomotion. For this reason, the current work focused primarily on investigating the effect of axial proprioceptive feedback on the swimming locomotor networks of the salamander. Future studies will improve on the aforementioned limitations to better understand the role that sensory feedback plays during walking behaviors.

Closed loop network

The effect of axial sensory feedback is largely consistent across different feedback topologies

Stretch feedback serves the purpose of responding to a high level of bending on one side of the body with an activation of the muscles that control the contracted side. This mechanism is achieved in animals through the excitation of the locomotor circuits on the stretched side and/or through the inhibition of the locomotor circuits on the contracting side.

Different animals were shown to display different patterns of effects and connectivities between the stretch-sensitive neurons and the locomotor network. In the lamprey, both ipsilateral excitatory neurons and inhibitory commissural neurons were reported [42, 43, 45]. These sensory neurons, called Edge Cells, were shown to lead to an increase in the frequency and a decrease in the intersegmental phase lag of the generated axial oscillations [47]. In larval zebrafish, two classes of mechanosensory neurons were found to have an active role during locomotion [108]. The first class includes ipsilaterally-projecting Glutamatergic Rohon-Beard (RB) neurons. These neurons are sensitive to touch stimuli and they were shown to degenerate over the first few weeks of development [109]. The second class includes GABAergic cerebrospinal fluid-contacting (CSF-c) neurons. These neurons are sensitive to body bending and many other chemical and neuronal cues [110, 111]. Interestingly, CSF-c neurons were shown to respond to a compression (rather than stretching) of the spinal cord by inhibiting the ipsilateral CPGs and motor neurons [110]. In the adult zebrafish, the intraspinal lateral proprioceptor organ (ILP) is the only reported class of intraspinal stretch-sensitive neuron. The ILP organs were shown to have commissural inhibitory projections to pre-motor interneurons [44]. Similarly to the Edge Cells in the lamprey, these neurons were shown to increase the frequency and decrease the intersegmental phase lag of the axial CPG activity [44].

Interestingly, the described sensory feedback connections typically present an asymmetry between caudally-oriented connections and rostrally-oriented connections. Both in the zebrafish and in the lamprey, stretch-sensitive organs were reported to connect preferentially to neurons located more rostrally in the spinal cord [41,44]. This variety of solutions found in animals raises the question about what is the effect of sensory feedback connectivity on the locomotor networks and whether there are different effects related to the different types of feedback.

For this reason, we studied the effect of feedback topology by varying the type of feedback type (ipsilateral excitation or contralateral inhibition) and the range of ascending and/or descending connections to the locomotor circuits (see Fig 8A). The analysis included four distinct connectivity schemes, namely ascending ipsilateral excitation (EX-UP), descending ipsilateral excitation (EX-DW), ascending contralateral inhibition (IN-UP) and descending contralateral inhibition (IN-UP) and descending contralateral inhibition (IN-DW). The range of the connections (PS range) was varied in the range [0.0, 2.5]cm (i.e., between 0 and 10 equivalent segments). Similarly, the PS synaptic weight (ω_{PS}) was varied in the range [0, 5].

Studying the effect on the frequency (f_{neur}) it was observed that increasing the feedback weight generally led to an increase of the frequency of oscillations (see Fig 8B). The effect was stronger for ascending (UP) connections with respect to descending connections (DW). Moreover, the effect was also stronger for excitatory projections (EX) with respect to inhibitory projections (IN). With respect to the length of the projections, a higher range also corresponded to a higher oscillation frequency. The same effect was observed for the IN-DW combination. The relationship was more nonlinear for IN-UP projections, although the higher frequency was still observed in the network with the highest PS weight and range. Overall the trend observed for the frequency was largely conserved across all the considered connection types. In the limit case $(PS_{range} = 2.5mm, \omega_{PS} = 5)$, all the topologies lead to an increase of f_{neur} compared to the open-loop case. The relative increase was equal to +41% (EX-UP), +36% (EX-DW), +36% (IN-UP) and +22% (IN-DW), respectively. This observation suggests that the result is a general consequence of introducing stretch feedback in the locomotor circuits. Notably, increasing the strength of the inhibitory PS connections had an opposite effect on f_{neur} with respect to increasing the commissural CPG inhibition strength (previously discussed in Fig 4A), suggesting that the two connection types play different roles in the locomotor circuits. A mechanistic explanation of this phenomenon is provided in the next section.

Studying the effect on the total wave lag (TWL, see Eq 3 in S1 File) it was observed that, irrespective of the feedback topology, long range connections with high values of feedback weight lead to a decrease of TWL (see Fig 8B). Descending connections had a markedly higher effect, leading to lower values TWL that correspond to quasi-standing waves of neural activity (i.e., $TWL \approx 0$). Additionally, for descending connections, TWL decreased monotonically both for an increase of the feedback weight and for an increase of the feedback range. Conversely, ascending connections with low or intermediate values of feedback weight, almost did not affect the value of TWL with respect to the open loop one. A decrease in the TWL value was only observed for long range ascending connections with high value of feedback weight. In the limit case $(PS_{range} = 2.5mm, \omega_{PS} = 5)$, all the topologies lead to a large decrease of TWL compared to the open-loop case. The relative decrease was equal to -73% (EX-UP), -108% (EX-DW), -59% (IN-UP) and -115% (IN-DW), respectively. Notably, the reduction in the TWL expressed by the network is consistent with the observation that in-vitro intersegmental phase lags are distributed over a larger range of values with respect to in-vivo preparations [96]. A mechanistic explanation of this phenomenon is provided in the next section.

The results of the analyses showed the same trends when changing the rheobase angle (θ_{RH} , see Eq 14) from 10% to 50% of the joints angular excursions (see Fig S1 from S5 File). In particular, the effect on the frequency of the oscillations was significantly weaker and it only led to a small increase of f_{neur} (up to 14%) regardless of the feedback topology. Conversely, the effect on the total wave lag was comparable to

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Fig 8. Frequency and phase lag modulation during swimming with different sensory feedback topologies and strengths. A) shows the different PS topology patterns studied in the model. The connectivity schemes are ascending ipsilateral excitation (EX-UP), descending ipsilateral excitation (EX-DW), ascending contralateral inhibition (IN-UP) and descending contralateral inhibition (IN-DW). The range of the connections (PS range) was varied in the range [0.0, 2.5]cm (i.e., between 0 and 10 equivalent segments). Similarly, the PS synaptic weight (ω_{PS}) was varied in the range [0, 0, 5.0]. **B**) shows the modulation of frequency (f_{neur}) observed for the different feedback topologies, represented in the four quadrants. In each quadrant, the contour plot shows the dependency of the frequency (f_{neur}) on PS range and ω_{PS} . On top, the projection of the relationship in the $f_{neur} - \omega_{PS}$ plane is displayed to better highlight the asymptotical values reached by f_{neur} . The different curves represent the frequency expressed by the networks with different connection ranges, displayed by the legend. The thick lines represent the mean metric values across 20 instances of the network. The shaded areas represent one standard deviation of difference from the mean. C) follows the same organization as the previous panel to represent the effect of feedback topology on the total wave lag (TWL).

the one described for θ_{RH} set to 10% of the joint angles amplitudes. Indeed, in the limit case, the TWL was reduced by -58% (EX-UP), -98% (EX-DW), -53% (IN-UP) and -95% (IN-DW), in agreement with the previous analysis. Overall, the results show that f_{neur} is strongly dependent both on the critical angle of activation θ_{RH} of the sensory neurons and on the weight of their synaptic connections ω_{PS} . On the other hand, the total wave lag was primarily affected by ω_{PS} irrespective of the choice of θ_{RH} .

While in the extreme scenarios (i.e. high ω_{PS} and PS_{range}) we found a high degree of invariance between the different connection types, there were notable differences in the PS effects for the intermediate cases. Indeed, the range of descending connections affected the total wave lag to a much larger extent with respect to ascending

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connections. On the other hand, the frequency followed a similar trend irrespective of the considered topology. This observation could provide an explanation for the asymmetry found in the connectivity of sensory feedback organs of swimming animals between ascending and descending projections.

Notably, this analysis investigated only a fraction of the possible feedback topologies, 505 namely the ones involving a single connection type (ipsilateral EX or commissural IN) 506 projecting continuously to rostrally or caudally-located target neurons. The effect of 507 feedback could vary when considering multiple coexisting PS types (e.g. EX-UP and 508 IN-UP), discrete connectivity schemes (e.g. targeting segments at a specific distance as 509 in [52]) or different target populations (e.g. PS to PS connections, as reported in [44]). 510 Given the absence of anatomical data regarding the connectivity of the sensory neurons 511 within the spinal cord of salamanders, the analyses in the next sections were performed 512 using symmetrical ascending and descending PS projections and a range of 5 segments, 513 a conservative hypothesis that is consistent with the range observed in the zebrafish and 514 lamprey [44, 45]. 515

The neuromechanical principles of frequency and phase-lag control by stretch feedback during swimming

In Fig 8 numerical simulations demonstrated that increasing the stretch feedback w_{PS} led to a global increase in the frequency and to a decrease in the phase lag of the oscillatory traveling waves. In this section, we provide some mechanistic arguments for these behaviors.

During locomotion the body bends on one side following the activation of the CPG neurons on the ipsilateral (same) side (see Fig 9A1). After some delay, the curvature crosses the angle θ_{TH} , which in turn activates PS neurons on the opposite side, according to the mechano-electrical transduction relationship (see Eq 13). During the time interval when the body curvature is higher than θ_{TH} , PS_{IN} neurons inhibit the CPGs on the opposite side, while PS_{EX} excite the CPGs on the same side. The former interaction facilitates the burst termination of the (active) opposite-sided CPGs, while the latter facilitates the burst initiation of the (inactive) same-sided CPGs. This mechanism causes an anticipated switch in the escape-from-inhibition mechanism of the CPGs, in accordance with experimental results on lampreys [43, 112]. The effect is analogous to the action of a bilinear torsional spring that acts to restore the straight position of the body once a critical angle is crossed (see Fig 9A2). The activation point of the spring and its elastic constant are proportional to the rheobase angle θ_{TH} and slope W_{PS} of the PS's mechanotransduction relationship, respectively (see Fig 9A3). This theoretical analogy confirms the increase in the swimming frequency with increasing $w_{PS} \sim W_{PS}$ and/or G_{PS} predicted numerically (see Fig 8 and text).

In terms of the impact on the total wave lag, one can assess the influence of global connections from PS_{in} to CPG, as illustrated in Figure 9B. Note that the identical line of reasoning is equally applicable to PS_{ex} connections, with the effects being interchanged for ascending and descending connections (see Fig S2 from S5 File).

Without sensory feedback, the CPG network produces an oscillatory traveling wave 542 propagating with positive wave lag (see Fig 9A, similar to the wave shown numerically 543 in Fig 3). When sensory feedback is turned on, it generates an imbalanced longitudinal 544 inhibition that destabilizes the wave propagation (see Fig 9B2 and B4). In the case of 545 ascending PS_{in} , rostral CPGs (marked by red in Fig 9B2) receive additional inhibition 546 from the PS_{in} . Since the CPGs oscillate due to escape from inhibition, the additional 547 PS_{in} inhibition will delay the activation of these rostral segments in the subsequent 548 cycles, reducing the overall wave lag. The asymmetry in the inhibition between rostral 549 and caudal segments makes this configuration unstable. 550

The wave will therefore reduce the wave lag up to reaching a stable state without

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Fig 9. The mechanism of frequency and phase lag control by stretch feedback. A)1 shows a sketch of a sinusoidal-like body curvature during anguilliform swimming. A2 shows populations of CPG and PS neurons at the location of maximal curvatures and their connections (active population have filled colors). The activity of PS neurons in correspondence of large curvatures (above θ_{PS}) acts as an equivalent spring that induces an increase of the swimming frequency. A3 shows the mechanotransduction relationship of PS neurons. B) represents the stereotypical CPG activities on the left side during swimming in a brief time window (black=active/firing, white=inactive) sorted according to the CPG positions (top=rostral, bottom=caudal). The sub-panel B1 shows the activities of CPGs with no feedback. The sub-panels B2-B3 and B3-B4 show subsequent time-windows at starting and final stages of activities from the time when inhibitory proprioceptive sensory feedback (PS_{in}) is turned on. B2 and B3 shows the network activity with ascending PS_{in} connections. B2 shows the network activity with descending PS_{in} connections. In each panel, green shows PS_{in} activities from the right side, brown the overlap of CPG and PS_{in} activities on opposite sides, and red the inhibition from PS_{in} to the CPGs on the subsequent cycle. The system evolves to minimize the interference between PS_{in} activity and CPG activity (i.e., the red area). Yellow arrows indicate the delay shift in the CPG activation caused by PS_{IN} inhibition.

imbalanced PS inhibition (i.e. when the wave lag is sufficiently low that PS_{IN} do not interact with the CPGs at their activation times, see Fig 9B3 and B4). In this configuration the network has minimized the intersection area between the PS activity and the CPG activity (i.e., the red area).

In the case of descending PS_{in} the more caudal CPGs receive an additional PS_{in} inhibition (marked by red in Fig 9B2), delaying their activation and increasing the overall wave lag. However, any increase of wave lag will cause even more rostral CPGs to be unbalanced. The wave therefore will be unstable until it breaks and reaches 0 or negative wave lag.

Interestingly, this observation explains why, in simulations, ascending projections (either excitatory or inhibitory) led to positive TWL values while descending projections caused an inversion of the wave propagation direction (see Fig 8), and why both excitatory and inhibitory unidirectional (ascending or descending) connections

from PS cells converge to similar values of TWL in the limit for high feedback strengths (see Fig 8).

Overall, the frequency increase can be explained as a local effect by the stretch sensors to the CPGs in correspondence to regions of large curvature. Conversely, the decrease in total wave lag can be explained as a global effect derived from the length of the sensory projections and the natural tendency of the network to produce rostro-caudal waves of activation.

Intermediate levels of sensory feedback improve the swimming performance 572

The effect of drive on the closed-loop network was studied by varying the value of the feedback connections weight ω_{PS} . The stimulation amplitude was varied in the range [3.0, 9.0]pA. The feedback weight was varied in the range [0.0, 5.0]. For each combination of the parameters, 20 instances of the network were simulated with different random seeds and the results are reported in terms of mean and standard deviation of the computed metrics (see Fig 10). The figure was obtained for a rheobase angle (θ_{RH} , see Eq 14) of 10%. The same analysis was performed for $\theta_{RH} = 50\%$ and the corresponding results are reported in Fig S3 from S5 File.

When studying the effect on the network's periodicity (PTCC, see Fig 10A), it can be noted that increasing the value of ω_{PS} results in an expansion of the stable oscillations region with respect to the range of the input stimulation. Indeed, the range of stimulation values for which PTCC > 1 increased by 50% when ω_{PS} was set to 1.0 and it reached up to a 90% increase for $\omega_{PS} = 5$. For low values of stimulation, sensory feedback allows to stabilize the spurious activations displayed by the network, giving rise to a more patterned locomotor output and a higher PTCC (see Fig 10A2). For high values of stimulation, sensory feedback acts as an additional commissural inhibition pathway, delaying the emergence of bilateral tonic activity and, consequently, also the drop in PTCC values. Overall, the mean PTCC value across all stimulation amplitudes increased monotonically with ω_{PS} (see Fig 10A3, black dots), underlining the beneficial effect of sensory feedback with respect to the stability of the oscillations. Interestingly, the same trends were observed for $\theta_{RH} = 50\%$ (see Fig S3A). In this case, however, the region of stable oscillations was expanded to a lower extent (+29% for $\omega_{PS} = 1, +41\%$ for $\omega_{PS} = 5$). This is due to the incapability of the network to reach the higher rheobase angle, thus preventing the sensory feedback from exerting its beneficial action on the locomotor rhythm.

Regarding the frequency (f_{neur}) , as observed in the previous sections, increasing ω_{PS} monotonically increases the frequency of the oscillations independently from the stimulation amplitude (see Fig 10B). Interestingly, the frequency does not increase indefinitely. Instead, f_{neur} settles at a specific value of 5.23Hz, acting as a feedback-related resonance frequency of the neuromechanical system. A similar saturation effect was observed was observed for $\theta_{RH} = 50\%$ (see Fig S3B). In this case, however, the higher rheobase angle led to a lower resonant frequency of 4.20Hz.

Notably, the range of variation of f_{neur} decreased monotonically with ω_{PS} , (see Fig 10B3, green dots). This observation highlights the trade-off between robustness and flexibility of the locomotor circuits. A network dominated by sensory feedback oscillates over a wider range of input drives but it also loses the capability to modulate the frequency of the generated patterns.

The described trade-off is even more evident when studying the effect of ω_{PS} on the total wave lag (TWL) with varying values of stimulation (see Fig 10C). Indeed, for $\omega_{PS} > 2$ the network only displays quasi-standing waves of activation independently of the drive (see Fig 10C2,3). This further emphasizes the necessity for intermediate feedback weights that do not lead to a sensory-dominated network.

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Fig 10. Intermediate feedback strengths improve the performance of the closed loop swimming network. A) shows the effect of the stimulation amplitude to the axial RS neurons and of the feedback weight (ω_{PS}) on the periodicity (PTCC) of the generated activity. The stimulation amplitude was uniform across all the axial RS modules and it was varied in the range [3.0, 9.0]pA. The feedback weight was varied in the range [0.0, 5.0]. The displayed data is averaged across 20 instances of the network. In A2, the curves differ by their corresponding value of ω_{PS} , reported in the bottom legend. The shaded areas represent one standard deviation of difference from the mean. A3 shows the minimum (in blue), maximum (in red) average (in black) and range (in green) for the PTCC values across all the considered stimulation values. The range values are reported on the y-axis on the right. Panels (B-E) follow the same organization to display the effect on the oscillation frequency (f_{neur}) , total wave lag (TWL), forward velocity (V_{fwd}) and cost of transport (COT). Combinations leading to PTCC values lower than 1 were not displayed.

Interestingly, the highest range of variation of TWL was observed for $\omega_{PS} = 0.7$ (see Fig 10C3, green dots). Similarly, for $\theta_{RH} = 50\%$, the highest range of variation was observed for $\omega_{PS} = 0.5$ (see Fig S3C). These results suggest that an intermediate level of sensory feedback could increase the network's flexibility in phase lag modulation.

With respect to the speed (V_{fwd}) , it can be seen that increasing ω_{PS} also increases 619 the speed of locomotion (see Fig 10D). This effect is primarily due to the augmented 620 oscillation frequency discussed above. Concurrently, intermediate levels of ω_{PS} also lead 621 to an increase in the average stride length (see Fig S4C from S5 File). Indeed, both the 622 stride length and the speed display a local maximum for $\omega_{PS} \approx 1$ and a stimulation 623 amplitude of 6.5pA (see Fig 10D1 and S4A). This outcome suggests that the model 624 produces an optimal forward propulsion when there is a balance between the sensory 625 feedback signal and the activity of the CPG network. Moreover, similarly to the 626 previous metrics, for $\omega_{PS} > 2$ we observe a decrease in the speed and stride length of the 627 generated locomotion. Concurrently, also the range of variation of the two metrics drops 628 significantly for high values of feedback weight (see Fig 10D3 and S4C). Overall, these 629 observations further show that a feedback-dominated network does not produce efficient 630 swimming. The discussed effects were also observed when changing the rheobase angle 631 to 50% (see Fig S3D), increasing the reliability and of the drawn conclusions. 632

Finally, with respect to the cost of transport, we observe a complex non-linear relationship with respect to the weight of the sensory feedback (see Fig 10E1). Interestingly, there appears to be a region of minimal cost of transport in correspondence of $\omega_{PS} \approx 0.7$. Indeed, the average COT across all stimulation values displays a global minimum in correspondence of such feedback weight (see Fig 10E3, black dots). In that region, the simulated salamander swims more efficiently with respect to the open loop network as well as with respect to the closed loop networks with higher values of feedback weight.

Interestingly, there is a mismatch between the ω_{PS} values that led to an optimal speed and to an optimal cost of transport. Indeed, robotics experiments proved that the kinematics necessary for efficient swimming differs from the one that optimises the speed of locomotion [113]. This observation suggests that a modulation of the sensory feedback weight could serve as a strategy for a transition between different swimming modes (e.g., cruising and escaping)

Overall, the results show that the best swimming performance is obtained with a balanced combination of open-loop and feedback-based rhythm generation. A purely open-loop network is more flexible in terms of frequency modulation but it lacks in terms of robustness and swimming performance. A feedback-dominated network robustly oscillates at high frequencies but is not controllable through the descending drive. Finally, balanced combinations of the two rhythm generation strategies lead to optimal swimming networks. The swimming activity generated by the closed-loop network when linked to the mechanical model is displayed in S4 Video.

Sensory feedback extends the region of noise rejection

The influence of noise on locomotor patterns was examined in the closed-loop network with varying levels of ω_{PS} (see Fig 11A,B). The noise level was varied in the range [5.0, 15.0]mV. The feedback weight was varied in the range [0.0, 3.0]. For each combination of the parameters, 20 instances of the network were simulated with different random seeds and the results are reported in terms of mean and standard deviation of the computed network periodicity (PTCC).

The figure shows that sensory feedback extends the stability region of the swimming network. Indeed, for $\omega_{PS} = 0.5$, the network could tolerate noise levels 13% higher than the open loop configuration while maintaining PTCC values higher than 1. Furthermore, the improvement scaled with ω_{PS} itself (+44% for $\omega_{PS} = 1.0$, +81% for $\omega_{PS} = 1.5$). This finding supports the idea of sensory feedback serving as a redundant rhythmogenic mechanism that boosts the robustness of the locomotor networks, as mentioned in [52].

To further showcase the positive effect of sensory feedback in reducing noise interference during swimming, a simulation was carried out starting with an initially 633

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Fig 11. Effect of noise on the closed loop swimming network. A) shows the effect of noise amplitude on the rhythmicity of the network (PTCC) with different levels of sensory feedback weight (ω_{PS}). The noise level was varied in the range [5.0, 15.0]mV. The feedback weight was varied in the range [0.0, 3.0]. The different curves represent the average PTCC values across 20 instances of the network for different levels of ω_{PS} (shown in the legend). The shaded areas represent one standard deviation of difference from the mean. **B**) displays the noise-feedback relationship in 2D. For a noise level of 9mV, increasing ω_{PS} from 0 (red dot) to 0.7 (green point) leads to a switch from irregular to organized rhythmic activity. C) shows the switch effect with a noise level of 7mV. The raster plot represents the spike times of excitatory neurons (EN), reticulospinal neurons (RS), motoneurons (MN) and proprioceptive sensory neurons (PS). Only the activity of the left side of the network is displayed. At time=2.5s, the sensory feedback weight is changed from $\omega_{PS} = 0$ (Feedback OFF) to $\omega_{PS} = 0.7$ (Feedback ON), restoring a left-right alternating rhythmic network activity. **D**) shows the muscle cells activity during the simulation in \mathbf{E} . \mathbf{E}) shows the axial joint angles during the simulation in E. Each angle is shown in the range $[-25^{\circ}, +25^{\circ}]$. F) shows the center of mass trajectory during the simulation in E. Activating feedback restored forward swimming.

unstable open-loop network (see Fig 11C). In this scenario, the noise amplitude was set to 9mV, which disrupted the swimming locomotor pattern. At time 2.5s, sensory feedback was activated with $\omega_{PS} = 0.7$. This activation revived the activity of the axial CPG network, leading to rhythmic activation of the muscle cells (see Fig 11E) and, subsequently, oscillations in the axial joints (see Fig 11D), restoring the swimming behavior. The activity switch had a transient time of approximately two seconds. During this time interval, small oscillations generated in correspondence with the head and tail joints lead to the activation of their respective PS neuronal pools. This activation ultimately spread to the entire network, resulting in a full wave of left-right alternating activity (see Fig 11D). The resulting locomotion was a forward swimming gait with kinematics that closely resembled the one produced in the absence of noise (see S5 Video).

Interestingly, the same switch mechanism could be achieved also for higher noise levels, provided that ω_{PS} was set to a sufficiently large value. In Fig S5 from S5 File, the feedback weight was set to 2.0 with an initial noise amplitude of 14mV. Despite the higher noise level, the emergence of synchronized oscillations was faster than in the previous case. On the other hand, the high ω_{PS} resulted in a quasi-standing wave of activity and, consequently, in inefficient swimming kinematics (see S6 Video).

It is important to note that sensory feedback could restore the rhythmic activity only in cases where the noisy network was capable of generating joints oscillations that could reach the threshold for the activation of the PS neurons (i.e., the rheobase angle θ_{RH} , see Eq 14). Indeed, when θ_{RH} was set to 30%, the expansion in the noise rejection region was significantly lower (see Fig S6 from S5 File).

Overall, sensory feedback and fluid-body dynamics were shown to improve the robustness of the swimming locomotor networks. Future biological experiments could explore this aspect using semi-intact preparations where dorsal roots are kept intact and the caudal joints are allowed to move, similarly to [37, 114, 115]. These experiments could add noise to the locomotor circuits by means of external current injection and evaluate the different responses of in-vitro and semi-intact preparations. This investigation could offer a more comprehensive understanding of the interaction between sensory feedback and rhythmic locomotor activity in salamanders.

Sensory feedback expands the rhythm-generating capabilities of the network

To better understand how sensory feedback enhances the rhythmogenic properties of the network, we examined the effect of varying inhibition strength with different ω_{PS} values (see Fig 12). The commissural inhibition strength was varied in the range [0.0, 6.0]. The feedback weight was varied in the range [0.0, 5.0]. For each combination of the parameters, 20 instances of the network were simulated with different random seeds and the results are reported in terms of mean and standard deviation of the computed metrics.

It was observed that as ω_{PS} increased, the region of stable oscillations expanded, both at low and high inhibition strengths. At higher inhibition levels, sensory feedback delayed the onset of the 'winner takes all' configuration by providing an extra burst termination mechanism for the active side, allowing the opposite side to escape ongoing inhibition (see Fig 12A, B). This is consistent with observations from models of the lamprey locomotor networks [1,43]. Interestingly, for high values of ω_{PS} the 'winner takes all' configuration completely disappeared and the network displayed oscillations for the entire range of studied commissural inhibition strengths.

At lower inhibition strength levels, sensory feedback helped channel the unpatterned oscillations of the open-loop network, reinstating an ordered left-right alternation pattern (see Fig 12C,D). Interestingly, this is in accordance with previous modeling studies on *C. Elegans* where coordinated swimming activity could be achieved in absence of central pattern generation from the axial circuits [116,117]. Overall these two effects lead to an expansion of the stable oscillation region by 63% for $\omega_{PS} = 1$. For



Fig 12. Sensory feedback extends the rhythmogenic capabilities of the network and reduces the sensitivity to internal parameters. A) shows the dependency of the periodicity (PTCC) on the inhibition strength and on the feedback weight (ω_{PS}) for the swimming network. The commissural inhibition strength was varied in the range [0.0, 6.0]. The feedback weight was varied in the range [0.0, 5.0]. The displayed values are averaged across 20 instances of the network. B) shows the inhibition-feedback relationship in 1D. The different curves represent the average PTCC values for different levels of ω_{PS} (shown in the legend). The shaded areas represent one standard deviation of difference from the mean. Panels C and D follow the same organization as the previous panels to highlight the inhibition-feedback relationship for ω_{PS} values in the range [0.0, 6.0]. E) displays the distribution of the total-order sensitivity indices computed for the different network parameters (listed on the y-axis and legend) with respect to the PTCC, total wave lag (TWL), forward velocity (V_{fwd}) and cost of transport (COT). The colors are associated to different values of ω_{PS} , reported in the legend.

very high feedback values (i.e., $\omega_{PS} > 2$), the region of stable oscillations was wider than the considered interval of inhibition strengths.

Interestingly, when the rheobase angle θ_{RH} (see Eq 14) was changed to 50%, the network's periodicity displayed a similar dependency on the inhibition strength and feedback weight (see Fig S7 from S5 File). In particular, the 'winner-takes-all' configuration was again deleted for high ω_{PS} values thanks to the feedback-mediated burst termination mechanism (see Fig S7A,B). On the other hand, for low levels of commissural inhibition strength, the effect was sensibly weaker (see Fig S7C,D). This is due to the incapability of the network to produce bending angles capable of reaching the higher θ_{RH} when both sides of the spinal cord are tonically active. Nonetheless, the stable oscillation region was expanded by 16% for $\omega_{PS} = 1$.

The findings highlight the duality in rhythm generation mechanisms, where both central pattern generation and sensory feedback play essential roles [118]. Particularly, the model shows how intermediate levels of sensory feedback not only preserve but enhance the network's capability to generate rhythmic locomotor patterns, emphasizing the need for a balanced integration of central and peripheral inputs for optimal locomotor performance.

Future biological experiments could further investigate the results of the current work using semi-intact preparations, where the caudal part of the body is left free to move and the strength of commissural inhibition is adjusted either pharmacologically or optogenetically. Through such experimental setups, we could gain a deeper understanding of the modulatory impact of sensory feedback on rhythmogenic network properties under different inhibition strengths.

Sensory feedback reduces the sensitivity to the neuronal and synaptic parameters of the network

The sensitivity analysis was repeated in closed loop control of the mechanical model with different values of ω_{PS} (see Fig 12E). The varied parameters included the same neuronal parameters studied for the open loop network. Additionally, the value of ω_{PS} was also modified by the analysis around its nominal value. The studied nominal values for ω_{PS} ranged from 0 (no sensory feedback case) to 1.5 (high sensory feedback case). A separate sensitivity analysis was carried out for each nominal ω_{PS} value. Each panel of Fig 12E shows the distribution of the total-order sensitivity indices computed for the different neuronal and mechanical parameters with respect to the correspondent metric. The distributions were obtained from 20 instances of the network simulated with different random seeds.

Studying the effect on the network's periodicity (PTCC), it can be seen that increasing ω_{PS} led to a decrease of the total-order sensitivity indices for all the remaining parameters. This result could indicate that the network becomes less dependent on a specific parameter tuning to generate stable oscillations. On the other hand, for $\omega_{PS} > 0.5$, the sensitivity index for the feedback weight itself tends to decrease. Indeed, for high ω_{PS} values, the effect of sensory feedback reaches a saturation and the network is not affected by small variations around its nominal value. Interestingly, the commissural inhibition weight (Δg_{glyc}) was consistently the parameter with the highest total-order sensitivity index. This indicates that, regardless of ω_{PS} , the network's periodicity is still primarily affected by changes in the inhibition strength.

The described trends are even more evident for the sensitivity indices computed with respect to the total wave lag (TWL, see Eq 3 in S1 File). Also in this case we observe a large decrease in the sensitivity indices of all the parameters with increasing values of ω_{PS} . Similarly to the previous case, the sensitivity index for the feedback weight reaches a maximum for $\omega_{PS} = 0.5$ and then drops for higher values. Indeed, higher

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values of ω_{PS} lead to a quasi-standing wave of axial activation (see Fig 10C) independently of small variation of the feedback weight value.

Finally, similar observations can be made for the sensitivity indices of the studied parameters with respect to the frequency (f_{neur}) , forward speed (V_{fwd}) and cost of transport (COT) of the generated activity. Overall, it was shown that intermediate values of ω_{PS} reduce the dependency of the locomotor circuits on a precise selection of the neuronal and synaptic parameters. On the other hand, excessive values of sensory feedback strength dominate the network activity and reduce the capability to actively control its behavior.

Limitations of the model

The current network architecture serves as a simplified model for the locomotor circuits of the salamanders. The biological motor circuits likely express different classes of interneurons (e.g., V2a, V0d, V0v [1, 12]) that were not explicitly modeled in this network iteration. The advancements in genetic identification of the neuronal types within the salamander spinal cord will allow to refine the number and identity of neuronal classes of the simulated network. Additionally, this identification will allow to gather class-specific electrophysiological data that will help characterizing the neuronal and synaptic properties of the network [1].

With respect to the limbs network, the current model only accounted for a single equivalent segment to simulate the rhythmic limb activity during locomotion. On the other hand, several studies revealed that distinct pools of interneurons govern the activity of each joint within the limbs [119, 120]. Future iterations of the network should account for this limitation in order to investigate the emergence of intra-limb coordination in salamanders. Additionally, the inter-limb connectivity of the current model was designed to generate coordination between diagonally opposed limbs, as observed during trotting gaits [62]. Different interlimb connectivity schemes could allow to simulate a richer variety of patterns (e.g., lateral and diagonal sequence walks) expressed by salamanders during ground locomotion [73].

Given the limitations of the walking network in the current model, this study focused on the effect of axial proprioceptive sensory feedback during swimming. Future studies will evaluate whether the beneficial effects of sensory feedback identified in the current work still exist during walking. More generally, the effect of the body dynamics (e.g. body stiffness and damping) and the environment dynamics (change of viscosity, etc.) should also be investigated to get a deeper understanding of the interplay between central pattern generation, mechanical properties and sensory feedback [116, 121, 122].

Finally, the proposed model could also be used to investigate the effects of other sensory modalities on the locomotor circuits of the salamander. In particular, the effect of exteroceptive sensory feedback from the hydrodynamic forces acting on the body during swimming could be analysed to extend the results from previous robotics studies [52]. Similarly, the walking network could receive pressure feedback from the ground reaction forces acting on the body and the limbs. Interestingly, the sensory feedback from ground interaction forces might explain the generation of S-shaped standing waves during walking [123]. Additionally, this sensory modality would allow to investigate the emergence of interlimb coordination through sensory feedback [124, 125].

Conclusions

In this study, we have developed a novel neuromechanical model to investigate the interplay between CPGs and proprioceptive feedback in governing the locomotor behaviors of salamanders. Our findings underline the intricate balance between central

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control and sensory feedback in producing adaptable and robust locomotor patterns in a bio-inspired network architecture.

Our simulations demonstrate that the rhythmic locomotor outputs, essential for 823 swimming and walking, can be dynamically modulated by varying the excitatory drive 824 to reticulospinal neurons, underscoring the significance of central neuronal mechanisms. 825 The model effectively replicates the physiological propagation of motor patterns seen in 826 natural salamander locomotion, highlighting the role of CPGs in generating basic motor 827 rhythms [1, 12, 95]. Additionally, the open-loop network responds to increasing levels of 828 external stimulation similarly to previous simulation studies, suggesting that the model 829 assumptions are well justified through a chain of controllers, ranging from detailed 830 bio-physical networks (e.g., [82]) through formal spiking models (e.g., [90] and present 831 model), all the way to more abstract controllers [70, 73]. 832

Incorporating sensory feedback into our model revealed its pivotal role in refining locomotor efficiency and adaptability [12]. The feedback mechanisms, modeled on different topologies observed in nature [42–44], not only increased the tail beat frequency and speed during swimming but also enhanced the network's robustness against noise and against badly-tuned CPG networks [52, 126]. This suggests a broader role of sensory feedback in locomotor control than previously appreciated, extending beyond mere fine-tuning to actively shaping the motor output and allowing it to work in a larger range of conditions than open-loop networks.

Our analyses indicate that sensory feedback contributes significantly to the locomotor network's robustness, enabling it to withstand high levels of noise and variability in neuronal parameters [52]. For intermediate values of sensory feedback strength, this robustness is achieved without compromising the flexibility of the system. Indeed, in this case the network retains the ability to adjust its locomotor patterns in response to changes in external drive and sensory input, echoing the adaptive nature of biological systems. Conversely, high values of feedback weight lead to uncontrollable networks where frequency and phase lags could not be modulated through descending commands.

The study elucidates a duality in rhythm generation mechanisms, where both central pattern generation and sensory-driven feedback play essential roles [118, 127]. Particularly, the model shows how intermediate levels of sensory feedback not only preserve but enhance the network's capability to generate rhythmic locomotor patterns, emphasizing the need for a balanced integration of central and peripheral inputs for optimal locomotor performance.

Our findings have significant implications for the design of bio-inspired robotic systems, suggesting that incorporating models of proprioceptive feedback can substantially improve the adaptability and robustness of locomotor control in bio-inspired robots [52, 73, 83]. Furthermore, the insights gained from this study contribute to our understanding of the evolutionary adaptations in vertebrate locomotion, offering a computational perspective on how sensory feedback mechanisms might have evolved to complement central control strategies across different species.

This work opens avenues for future research to explore the effects of more complex 863 sensory feedback mechanisms, including those involving other modalities such as tactile 864 and visual inputs [1,128]. Notably, the current paper mainly focused on the analysis of 865 the swimming locomotor networks. In the future, the investigation should be expanded 866 to include the walking network and the effect that proprioceptive (e.g., shoulder and hip 867 flexion/extension) and exteroceptive (e.g., ground reaction forces acting on the limbs) 868 feedback modalities have on the resulting locomotor activity. Additionally, experimental 869 validation of the model's predictions in biological systems, particularly through 870 manipulation of sensory feedback pathways, could provide deeper insights into the 871 neuro-mechanical underpinnings of locomotion [12, 44]. 872

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In conclusion, our model advances the understanding of the multifaceted control 873 mechanisms underlying salamander locomotion, bridging the gap between simplistic 874 representations and the complex realities of biological locomotion. By highlighting the 875 synergistic roles of CPGs and proprioceptive feedback, this work lays the groundwork 876 for future explorations into the neuronal basis of movement in vertebrates, with 877 potential applications in robotics, rehabilitation, and the study of locomotor evolution. 878

Methods

Neuronal model

In this model, neurons are modeled as adaptive integrate-and-fire (hereafter referred to 881 as IF) neurons [129], which are essentially formal spiking neurons, augmented with an 882 adaptation variable as 883

$$\tau_{memb}\frac{du}{dt} = -(u - E_{rest}) - R_{memb}\omega + R_{memb}I_{syn} + R_{memb}I_{ext}(t)$$
(1)
$$\tau_{\omega}\frac{d\omega}{dt} = -\omega$$
(2)

In this context, u represents the membrane potential, E_{rest} signifies the neuronal resting potential, R_{memb} is the input membrane resistance, I_{syn} represents the total 885 synaptic current, I_{ext} is the external input current, and ω denotes the adaptation variable. The neuronal model integrates inputs and, upon reaching the firing threshold 887 E_{th} , the spike time of a neuron is documented followed by a reset of the membrane 888 potential to the reset voltage E_{rest} . Each spike occurrence increments the adaptation 889 variables by a value of $\Delta \omega$. Moreover, post-spike, the neuron is held at the resting 890 membrane potential for a neuron type-dependent refractory duration t_{refr} . 891

The total synaptic current is given by the contribution of different synaptic conductances g_{syn} and their respective synaptic reverse potential E_{syn} according to

$$I_{syn} = \sum_{syn} g_{syn}(u - E_{syn}) \tag{3}$$

The conductance g_{syn} for both inhibitory and excitatory synaptic currents sees an 894 increment by Δg_{syn} with every incoming spike [130]. Additionally, there is a constant 895 synaptic transmission delay of 2ms between the spike event and the corresponding g_{syn} 896 modification. The temporal evolution of synaptic conductances is dictated by 897

$$\tau_{syn} \frac{dg_{syn}}{dt} = -g_{syn} \tag{4}$$

The excitatory postsynaptic potentials (EPSPs) embody a swift alpha-amino-3-hydroxy-5-methylisoxazole-4-propionic acid (AMPA)-like component alongside a slowly diminishing N-methyl-D-aspartic acid (NMDA)-like component, taking cues from lamprey findings [131] and salamander modeling endeavors [82,90]. Conversely, the inhibitory postsynaptic potentials (IPSPs) act on a single glycinergic (GLYC)-like component.

When noise was included in the network, it was implemented as an Ornstein-Uhlenbeck random process [132, 133] added to the membrane voltage of the neurons as a current

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$$I_{noise} = \frac{\sigma_{noise}}{R_{memb}} \sqrt{\frac{2}{\tau_{memb}}} \xi(t) \tag{5}$$

where $\xi(t)$ is a zero-mean Gaussian white noise process with standard deviation σ_{noise} denoting the strength of the disturbance.

Muscle cells model

Muscle cells (MC) represent an additional non-spiking cell population added to the model in order to transform the discrete spiking trains coming from the simulated neurons into a continuous output signal which is then used to compute the output torque. They are described by

$$\tau_{mc} \frac{dM_{side_i}}{dt} = -M_{side_i} + RI_{syn_{mc}} \tag{6}$$

$$I_{syn_{mc}} = \sum_{syn_{mc}} g_{syn_{mc}} (M_{side_i} - E_{syn_{mc}})$$
⁽⁷⁾

Where, similarly to Eq 1, τ_{mc} is the membrane time constant, M_{side_i} is the membrane potential ($side \in L, R$), $I_{syn_{mc}}$ is the total synaptic current, $g_{syn_{mc}}$ is the synaptic conductance and $E_{syn_{mc}}$ is the synaptic reverse potential. Effectively, the muscle cells are modeled as firing rate neurons [85] that act as low-pass filters of the input activity. Their input is normalized in the range [0.0, 1.0] and can be interpreted as the degree of activation of the motor neurons in the corresponding network region.

Network model

The salamander network comprises 5 different classes of neurons, namely excitatory neurons (EN), inhibitory neurons (IN), motoneurons (MN), reticulospinal neurons (RS) and propriosensory neurons (PS). The populations differ for their neuronal and synaptic properties, as well as for their connectivity patterns with other neurons. The values of the neuronal and synaptic properties of the network are reported in S2 File.

Figure 1 depicts conceptual schematics for the network model of the salamander spinal circuitry. Neurons of the axial network are arranged uniformly in a continuous two-dimensional sheet without explicit segmental boundaries. The network is 10cm long, in accordance with the typical length of *Pleurodeles waltl*'s spinal cord [134]. Each neuron is assigned a coordinate (x_{neur}) specifying its position along the spine starting from the rostral part. Additionally, the axial network is divided in two hemicords built according to the same organization principles. For this reason, each neuron is also assigned a second coordinate (y_{neur}) indicating its position on the left or right hemicord. The coordinate system introduced for the network organization allows for the definition of position-dependent connections between neurons in order to target specific regions of the spinal cord as well as the most proximal neurons for each candidate. Connection densities and weights in the salamander spinal cord are largely unknown and have been treated as open parameters.

Since the salamander spinal cord exhibits 40 vertebrae and ventral roots, equally spaced consecutive sections of the network can be viewed to represent a spinal segment (henceforth termed an equivalent segment), with a length of 2.5mm. Each equivalent segment accounts for 60 EN and 60 IN, representing the CPG component of the network. The number of CPG neurons was derived by [90, 120]. Additionally, each equivalent segment comprises 10 MN and 10 PS units. The number of motoneurons was

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inspired from [135, 136]. The number of proprioceptive sensory neurons was inspired by studies in the lamprey and zebrafish [44, 45]. Finally, each equivalent segment includes 2 MC units whose role is to integrate the spiking activity of the motoneurons. Note that the classes and numbers of neurons in the salamander locomotor networks is largely unknown and it could only be estimated in the present work. The model could be further refined with future advancements in imaging and genetic identification of the salamander spinal interneurons [12].

Figure 1C depicts a schematic of the neuronal organization of each equivalent segment. We follow the convention of denoting each equivalent segment by two oscillator symbols since they can generate antiphase activity. However, a half-center (in our case a hemi-segment) cannot oscillate by itself in the current model due to the absence of rhythm-generating pacemaker currents. Interestingly, in-vitro experimental results showed that abolishing the persistent sodium current via bath application of riluzole in surgically-isolated spinal segments did not affect the frequency and rhythmicity of the corresponding ventral root oscillations [95]. This result indicates that pacemaker currents may not be necessary for rhythm generation at a segmental level in salamanders.

Separate RS populations of 100 excitatory neurons each project to the rostral portion (Rost), the mid-trunk (Midt), the end of the mid-trunk (Endt), the pelvic girdle area (Pelv), the tail (Tail), and the caudal end (Caud) regions of the axial network, with distinct connectivity patterns (cf. Figure 1B, dashed lines indicating incoming descending connection density from RS nuclei). The connection probabilities of the RS populations are given by overlapping trapezoidal distributions. The parameters of the distributions are listed in Table S5 of S2 File. The overlapping of the RS projections is reduced between the end of the mid-trunk and the pelvic girdle area (see Figure 1B). This reduced connectivity permits to effectively decouple the control of the trunk and tail regions, thus allowing the two parts to be independently activated and modulated, consistently with experimental findings [60, 101, 107].

The connectivity between neurons within the axial network is obtained by means of asymmetric exponential distributions centered in correspondence with the pre-synaptic neuron. The probability distribution as a function of the pre-synaptic (x_{pre}) and post-synaptic (x_{post}) neurons positions is defined as

$$P(x_{pre}, x_{post}) = \begin{cases} Ae^{-\frac{(x_{pre} - x_{post})^2}{4\sigma_{up}^2}}, \text{ if } x_{pre} >= x_{post} \\ Ae^{-\frac{(x_{pre} - x_{post})^2}{4\sigma_{dw}^2}}, \text{ if } x_{pre} < x_{post} \end{cases}$$
(8)

Where A specifies the maximum connection probability, while the parameters σ_{up} and σ_{dw} determine how wide the probability distributions are for ascending (i.e. rostrally oriented) and descending (i.e. caudally oriented) connections respectively. The parameters of the connections for the axial network are listed in Table S6 of S2 File.

Within the axial CPG network, the connections are biased towards more caudally located neurons, as suggested by physiological and modeling studies in the lamprey [4,81,137,138] (see 1C). Conversely, the axial PS and MN populations project symmetrically to the neighbouring neurons. The MN neurons target specifically the closest muscle cells, thus providing an estimate of the activity of the local axial region. In the axial network, the PS neurons target the closest EN, IN and MN neurons both ipsilaterally and contralaterally. Previous experimental results from lamprey and zebrafish highlighted an asymmetry of the PS projections in favour of dominantly ascending connections [41,44,45]. For this reason, the effect of different PS topologies was also studied and it is discussed in the following sections.

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Similarly to the RS connectivity, the range of the connections between the end of the 991 mid-trunk and the pelvic girdle area is reduced to 0.75mm (i.e. one third of an 992 equivalent segment). The discontinuity allows the trunk and tail sub-networks to 993 independently generate waves of activation that do not propagate to the other section 994 when not excited from its corresponding RS modules (more details in the next sections), 995 in accordance to experimental findings [101, 107]. 996

In addition to the axial network, four sub networks are modeled to simulate the activities of the limbs. Two limb pairs are placed in correspondence of the first axial segment (X = 0.0mm) and of the first pelvis segment (X = 37.5mm) respectively (see Fig 1B). Separate oscillatory networks for each limb are implemented as equivalent 1000 segments, analogously to the axial network. Note that this is a simplification that does 1001 not take into account the possible presence of separate oscillatory pools for each distinct 1002 degree of freedom [119, 120]. Also, the current model does not take into account the 1003 presence of a pattern formation layer, which is generally accepted in other animal 1004 models such as cats and mice [139–142]. The two half centers of each limb oscillator 1005 represent the flexor and extensor components respectively. Additionally, the limb 1006 segments project to the pelvic and scapular region of the axis with an exponential 1007 probability (see Eq 8) with A = 0.1, $\sigma_{up} = 0.0mm$, $\sigma_{dw} = 5.0mm$ (see Fig 1C). The 1008 connections involve the EN from the flexor component of each limb and the 1009 caudally-oriented ipsilateral EN and IN of the axial network. 1010

The synaptic connections between the four limb CPGs were designed to resemble the 1011 activation patterns displayed by salamanders during trotting gait [94] (see Fig 1D). The 1012 inter-limb projections were drawn between the IN populations of the flexor side of the 1013 limbs segments. The connections targeted ipsilateral limbs (i.e., limbs on different 1014 girdles located on the same side of the body) and commissural limbs (i.e., limbs on the 1015 same girdle located on opposite sides of the body). The two connection types ensured a 1016 synchronous activation of the diagonally opposed limbs segments. The density of 1017 inter-limb connections was lower than the one within the limb hemisegments in order to 1018 avoid the disruption of their rhythm generation properties. In animals, the inter-limb 1019 connectivity may be subject to modulation depending on the expressed gait 1020 pattern [39, 105, 106]. A full analysis of the possible inter-limb connectivity schemes and 1021 their implication in inter-limb coordination is outside the scope of the current work. 1022

The synaptic weights were adjusted manually by modulating the corresponding 1023 Δg_{sun} parameter (see Eq 4) and are listed in Table S3 of S2 File. The weights for the 1024 PS populations are additionally modulated by an adimensional gain ω_{PS} (see Fig 2G,H) 1025 according to 1026

$$\Delta g_{syn}^* = \Delta g_{syn} \omega_{PS} \tag{9}$$

Where $syn \in \{ampa, nmda, glyc\}$ and Δg_{syn} is the nominal value listed for the PS 1027 populations in Table S3 of S2 File. The role of ω_{PS} is to represent the relative strength 1028 between PS connections and intra-CPG connections. When ω_{PS} is set to 1, the 1029 PS-originating synapses have the same strength as the connections between EN and IN 1030 populations within the locomotor circuits. More details about the role of ω_{PS} are 1031 discussed in the next sections. 1032

Mechanical model

The mechanical model consists of a salamander body, inclusive of axial and limb joints, 1034 that was previously introduced in [92]. The axial configuration is a chain of 9 links (see 1035 Fig 2A), interconnected by 8 active joints (i.e., driven by muscle models) functioning in 1036 the horizontal plane (see Fig 2B, red arrows). Moreover, the axial structure 1037

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incorporates 6 passive joints positioned at the mid trunk (2 joints) and tail (4 joints) 1038 sections (see Fig 2B, blue arrows). These passive joints are activated solely by the 1039 passive elements of the muscle model, they operate in the sagittal plane and they enable 1040 the belly and tail to engage with the ground. The model also comprises four limbs, with 1041 each limb accounting for 4 degrees of freedom (DoF). These encompass 1042 flexion/extension, abduction/adduction, and internal/external rotation of the shoulder 1043 (or hip) along with flexion/extension of the elbow (or knee, see Fig 2C). The 1044 geometrical and dynamical properties of the model were derived from *Pleurodeles waltl* 1045 biological data. More details about its design are given in [92]. For simplicity, the 1046 density of the model was considered constant across the length of the body and equal to 1047 $981.0kq/m^3$. This value renders the model positively buoyant in water, thus allowing to 1048 study swimming behaviors during surface-level aquatic locomotion. In the future, 1049 further refinements of the model could aim at varying the density of the segments based 1050 on the differential amount of muscles, bones and other tissues. The cross-section of the 1051 model varies with its axial position, resulting in a heavier head and trunk section 1052 compared to the lightweight tail section. 1053

Electro-mechanical transduction

The activity of muscle cells is transduced in output torques for the mechanical model using the muscle model proposed by Ekeberg [143] (see Fig 2D). For each joint *i*, the model receives in input the flexor (M_{L_i}) and extensor (M_{R_i}) activations from the muscle cells (see Eq 6), and the current joint angle (θ_i) and speed $(\dot{\theta_i})$ to compute the resulting output torque (τ_i) via:

$$\tau_i = \alpha_i M_{diff_i} + \beta_i (\gamma_i + M_{sum_i})(\theta_i - \theta_{0_i}) + \delta_i \dot{\theta}_i) \tag{10}$$

$$M_{diff_i} = G_{MC_i}(M_{L_i} - M_{R_i}) \tag{11}$$

$$M_{sum_i} = G_{MC_i} (M_{L_i} + M_{R_i})$$
 (12)

Where we α_i is the active gain, $\beta_i \gamma_i$ is the passive stiffness, $\beta_i M_{sum_i}$ is the active 1060 stiffness, δ_i is the damping coefficient, θ_{0_i} is the resting angle position. The flexor and 1061 extensor activations are scaled by a factor G_{MC_i} that is varied across different joints 1062 based on the salamander kinematics (more details in the following sections). The full 1063 electro-mechanical pipeline is displayed in Fig 2F. The segmental activity information 1064 from the axial CPG flows through the motoneurons and it is temporally and spatially 1065 integrated by the muscle cells. The muscle cells consequently provide the activation 1066 signal for the Ekeberg muscles that, in turn, output the total torque generated by the 1067 corresponding joint. Effectively, the muscle model is a spring-mass-damper system with 1068 variable stiffness which results in a non-linear input-output relationship (see Fig 2D). 1069 This implementation is a simplified version of the widely used Hill-type muscle 1070 model [144] that, while being more numerically tractable, still comprises active and 1071 passive components of muscle activations. 1072

In the axial joints, the muscle parameters were chosen in order to operate in a target dynamical regime in terms of resonance frequency, damping factor and zero-frequency gain of the Ekeberg muscles. A detailed description of the muscle parameters optimization is reported in S3 File.

During swimming, the resting angles of the limb joints were set in order to fold them along the body axis [94]. During walking, since a single equivalent segment was used for each limb, delayed copies of the same MC signals were used to control its multiple degrees of freedom to provide the periodic limb movements necessary for ground locomotion. In order to achieve that, the period of each limb's oscillations was estimated online with a threshold-crossing strategy. The threshold for the membrane

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potential of the muscle cells was set to the same value for all the limbs ($M_{side_i} = 0.25$, 1083 see Eq 6). The times of threshold crossing were registered and the corresponding limb 1084 period was estimated as the difference between two successive threshold times. The 1085 different degrees of freedom were then driven by a delayed version of the same MC 1086 signals. Similarly to [74], the delays chosen for the DoF were applied as a fraction of the 1087 estimated period. The delay fraction was set to 0% for the shoulder 1088 abduction/adduction degree of freedom, which thus acted as the leading joint. 1089 Conversely, the delay was set to 25% for the shoulder flexion/extension, shoulder 1090 internal/external rotation and elbow flexion/extension to ensure ground clearance 1091 during the swing phase. 1092

Mechano-electrical transduction

The proprioceptive feedback neurons residing in the stretched side of the body receive 1094 an input that is proportional to the angles of the mechanical joints according to: 1095

$$I_{PS} = max(0, G_{PS} \cdot \theta) \tag{13}$$

Where G_{PS} is the gain for the mechano-electrical transduction (see Fig 2F). The 1096 angle input provided to each sensory neuron is obtained from the linear interpolation 1097 between the two closest mechanical joints, thus assuming a continuous curvature 1098 gradient along the body. The value of the sensory feedback gain for each segment was 1099 chosen based on the rheobase current of the sensory neurons $(I_{RH} = 50pA)$ in order to 1100 select the critical angle at which they start to fire during locomotion. The gains were 1101 chosen so that the critical angle scales proportionally with respect to the reference 1102 amplitude of the corresponding joint oscillations (Θ_i , derived from [62]), according to 1103

$$\theta_{RH} = \frac{I_{RH}}{G_{PS}\Theta} \cdot 100\% \tag{14}$$

Where θ_{RH} denotes the rheobase angle (expressed as a percentage of Θ_i), and it was 1104 equal for all the segments of the network. This was done according to the hypothesis 1105 that the stiffer body joints will be more sensitive to stretch with respect to the flexible 1106 tail joints. The scaling also ensures that the activation of the sensory neurons happens 1107 in the same phase of the oscillation regardless of their amplitude. Unless stated 1108 otherwise, the rheobase angle for each segment was set to 10%. Several analyses were 1109 also carried out with different θ_{RH} values (e.g., 30% and 50%) and the corresponding 1110 figures can be found in the supplementary material. Future biological studies could 1111 investigate whether salamanders present a gradient in the degree of excitability of the 1112 sensory neurons along the spine and, consequently, what is the critical spine curvature 1113 that leads to their activation. 1114

Overall, the effect of sensory feedback is mainly modulated by two parameters, namely the G_{PS} gain and ω_{PS} (see Fig 2E,F). The former affects the rheobase angle θ_{RH} . The latter modulates the influence that an activation of PS has on the other neurons. In an analogy with proportional control, the two parameters effectively represent the threshold value (θ_{TH}) and slope (W) of the overall sensory signal provided to the locomotor network.

Hydrodynamic model

When submerged in water, the body is subjected to buoyancy and hydrodynamic forces. $_{1122}$ Buoyancy forces are applied to each link according to its submerged volume V_{UW} $_{1123}$

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$$\mathbf{F}_{\mathbf{b}_{i}} = V_{UW_{i}}\rho g \tag{15}$$

Where ρ is the water density and g is the gravity acceleration $(g = 9.81m/s^2)$. Since salamanders typically swim in high Reynolds number regimes, we opted for a simplified hydrodynamic model composed of inertial drag forces, analogously to [143,145]. For each link, a speed dependent drag force (F_d) is applied in each dimension (i) according to:

$$\mathbf{F}_{\mathrm{d}_{i}} = 2\rho C_{d_{i}} A_{i} V_{i}^{2} \tag{16}$$

Where A is the link surface, c_{d_i} is the drag coefficient in the i-th dimension, V_i is the 1129 current speed of the joint in the i-th dimension. The drag coefficients were chosen in 1130 order to match the mechanical metrics computed from swimming salamanders. To 1131 achieve this, the muscle cells gains G_{MC_i} (see Eq 10) were adjusted so that the angular 1132 excursions displayed by the joints would match the ones reported in [62]. Successively, 1133 the drag coefficients were hand-tuned to achieve a good level of agreement between the 1134 experimental and simulation metrics. A detailed description of the muscle cells factors 1135 selection and resulting swimming metrics is reported in S4 File. The selected 1136 coefficients allowed to closely match the experimental results in terms of axial 1137 displacements and speed (see Fig S1 from S4 File). An animation displaying the 1138 generated swimming activity can be seen in S1 Video 1139

Neuromechanical simulation

The neural network was simulated with the Brian2 library [146, 147] using the Python 1141 programming language. The tool allows to define custom neuronal and synaptic models, 1142 as well as to define connection probability distributions dependent on neurons' type, 1143 position and properties. Overall, the model is comprised of for 7'408 neurons and 1144 approximately 100'000 synaptic connections. Unless stated otherwise, the duration of 1145 each simulation was set to 10s. The network dynamics were integrated with forward 1146 Euler integration method and with a timestep of 1ms. Using Brian2 allows to convert 1147 the neural network into Cython code, boosting the simulation performance. 1148

The mechanical model was simulated in synchrony with the neural model with the MuJoCo physics engine [93] via the FARMS simulation framework [92]. The tool provides a platform for the definition and simulation of neuromechanical simulations, as well as for the incorporation of hydrodynamic and muscle models.

The two simulations exchanged data via queue communication among their respective threads. The integration step of the mechanical model was set to 1ms (the same as the neural model) and data exchange was carried out at each timestep.

The data from the neuromechanical simulations were used to compute the 1156 corresponding neural and mechanical metrics to evaluate the performance of the model. 1157 The details about the signal processing pipeline can be found in S1 File. 1158

Sensitivity analysis

Sensitivity analysis is the study of how the uncertainty in the output of a system can be apportioned to different sources of uncertainty in its inputs. Studying the sensitivity of the network performance metrics (e.g., periodicity, frequency, phase lag) with respect to the network parameters helps identify the key variables influencing the system's behavior, offering insights into the importance of various design decisions. The sensitivity of each input is represented by a numeric value, called the sensitivity index.

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Total-order indices measure the contribution to the output variance caused by a model 1166 input, including higher-order interactions. 1167

Simulations were performed using SALib [148, 149], an open-source library written in 1168 Python for performing sensitivity analyses. The parameters were chosen using Saltelli's 1169 extension of the Sobol' sequence, a quasi-random low-discrepancy sequence used to 1170 generate uniform samples of parameter space [150, 151]. The neuronal and synaptic 1171 parameters of the CPG network were varied for the analysis. In particular, the neuronal 1172 parameters included τ_{memb} , R_{memb} , τ_{ω} , $\Delta \omega$ of the EN and IN populations. Similarly, 1173 the synaptic parameters included the weights Δg_{ampa} , Δg_{nmda} , Δg_{glyc} of the EN and 1174 IN projections. Finally, when the sensitivity analysis was carried out in the full 1175 closed-loop network, the sensory feedback parameters G_{PS} and ω_{PS} were also modified. 1176 During the analysis, each of the aforementioned parameters was sampled in a range 1177 between 0.95 and 1.05 times its nominal value. 1178

Overall, each sensitivity analysis involved the simulation of 704 combinations of parameter values. Additionally, the analysis was repeated for 30 instances of the network, built with different random seeds. For each iteration, the total-order sensitivity indices were computed with respect to each neural and mechanical metric.

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SI	upporting information	1183
S 1	File. Signal processing pipeline. See S1_signal_processing.pdf	1184 1185
S 2	File. neuronal and synaptic properties of the network. See S2_neuronal_synaptic_properties.pdf	1186 1187
$\mathbf{S3}$	File. Optimization of the muscle parameters for the mechanical model. See S3_muscle_properties_optimization.pdf	1188 1189
S4 sw	File. Matching of the salamander kinematics during anguilliform rimming. See S4_matching_swimming_kinematics.pdf	1190 1191 1192
$\mathbf{S5}$	File. Supplementary figures for the analyses described in the results. See S5_supplementary_results.pdf	1193 1194
$\mathbf{S1}$	Video. Swimming salamander matching the experimental kinematics.	1195

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See S1_video_matching	g_kinematics_swimming	g.mp4		119

- S2 Video. The generated swimming activity without sensory feedback. 1197 See S2_video_open_loop_swimming.mp4 1198
- S3 Video. The generated trotting activity without sensory feedback. 1199 See S3_video_open_loop_trotting.mp4 1200
- S4 Video. The generated swimming activity with sensory feedback. 1201 See S4_video_closed_loop_swimming.mp4 1202

S5 Video.	Sensory feedback restoring the swimming activity in a noisy	120
network.		120
See S5_vie	leo_restoring_swimming.mp4	120

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S6 Video. Sensory feedback restoring the swimming activity in a very noisy network.

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See S6_video_restoring_swimming_high_noise.mp4

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