Efficient computation via sparse coding in electrosensory neural networks
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The electric sense combines spatial aspects of vision and touch with temporal features of audition. Its accessible neural architecture shares similarities with mammalian sensory systems and allows for recordings from successive brain areas to test hypotheses about neural coding. Further, electrosensory stimuli encountered during prey capture, navigation, and communication, can be readily synthesized in the laboratory. These features enable analyses of the neural circuitry that reveal general principles of encoding and decoding, such as segregation of information into separate streams and neural response sparsification. A systems level understanding arises via linkage between cellular differentiation and network architecture, revealed by *in vitro* and *in vivo* analyses, while computational modeling reveals how single cell dynamics and connectivity shape the sparsification process.

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Introduction
Peripheral sensory neurons typically encode a wide range of environmental stimuli. Moving more centrally, this dense coding is progressively replaced by a narrowing of neural responses to a subset of features of the whole stimulus ensemble [1–5]. Such sparse coding makes for easier read-out by downstream neurons, and is more energy efficient [6]. The mechanisms underlying sparsification are still elusive, and their elucidation is a fundamental goal of systems neuroscience. In particular, there is a quest for links between cellular/molecular properties – revealed through *in vitro* recordings – and network architecture and responses – revealed through *in vivo* and anatomical studies. Ideally the process would be understood for a behaving animal. Computational modeling has revealed that sparse coding requires non-linear mechanisms in both the biophysical and network properties of sensory circuitry.

This review looks at the successive transformations of sensory input in the electrosensory system of weakly electric fish. This system has enabled major advances in understanding how differences in anatomy and cellular physiology underlie sparse coding. This is because links between *in vitro* and *in vivo* subthreshold and spiking dynamics are possible in the awake animal responding to a wide range of behaviorally relevant sensory input. The review is organized as follows. We first introduce the electrosensory system and its natural stimuli. We then review how electroreceptors can respond differentially to all these stimuli. Next, we describe the first order electrosensory target (hindbrain) emphasizing how linked cellular and architectural differentiation initiates the routing of these inputs into separate output streams. Finally, we summarize recent work on how an even sparser and more efficient representation of the stimulus ensemble is generated in the midbrain. Throughout, we highlight the dynamical and computational features of neural function that provide the substrate for the sparse code.

The electrosensory system
Gymnotiform wave-type weakly electric fish produce an electric field by repetitively discharging a specialized electric organ located in their tail (Figure 1a). This electric field is commonly referred to as the electric organ discharge (EOD). The EOD is sensed everywhere on the body as a transdermal potential difference. Perturbations of the EOD, caused by nearby objects with conductivity different than that of the surrounding water (e.g. prey, conspecifics), modify the transdermal potential difference from its baseline value in a spatially dependent fashion that is commonly referred to as the electric image (Figure 1a). Computational studies have explained the detailed spatiotemporal structure of electric images caused by small objects (e.g. prey), self-generated movement (e.g. tail bends) [7], as well as spatially diffuse images caused by conspecifics [8]. Further, they have predicted important characteristics such as spatial resolution [9].

Scattered on the skin surface are electroreceptors that monitor the transdermal potential difference. While
P-type electroreceptors respond to amplitude modulations (AMs) of the EOD, T-type electroreceptors respond instead to frequency modulations [10] while ampullary electroreceptors respond to exogenous (non-EOD) electric signals. Here we consider exclusively P-type electroreceptors and their downstream neurons, and henceforth refer to P-type electroreceptors as ‘electroreceptors’.

Electroreceptor coding
Electroreceptors are sensitive to phase-modulated voltages [21]. They are sensitive to stimuli that are too small to elicit a response from ELL’s primary sensory cortex [21]. The coding of electroreceptors is non-linear, i.e. the response of a neuron is not proportional to the input stimulus [21].

The Electrosensory lateral line lobe: Efficient circuits for feature extraction and sparse coding
Electroreceptors project to the electrosensory lateral line lobe (ELL). Each afferent trifurcates and synapses onto pyramidal cells within three structurally similar parallel segments: centromedial (CMS), centrolateral (CLS) and lateral (LS) [29–33]. Ampullary electroreceptors project to the medial segment (MS) and are not considered here (Figure 2). All three segments thus receive identical input and each forms an entire topographic map of the animal’s body surface. The output from pyramidal cells from all maps converges onto the midbrain torus semicircularis (TS) (Figure 2).

ELL circuitry: on and off center cells and columnar organization
The ELL has a characteristic layered organization and is organized in columns with all cells in a column receiving identical afferent input [34,35]. Our description will center on the main output ELL neurons, pyramidal cells, which fall into two major classes, E and I, corresponding to On and Off center cells in the visual system [36–38]. This E/I dichotomy is a major architectural feature for sparse neural coding by having E and I-cells respond selectively to input caused by conductors (e.g. prey) and dielectric (e.g. rocks) objects, respectively. E and I-cells also preferentially respond to chirps associated with same and opposite-sex interactions, respectively [39,40].

Functional heterogeneities in pyramidal cells
There are large heterogeneities amongst pyramidal cells of a given type: superficial pyramidal cells receive strong feedback, deep pyramidal cells do not [41]. Superficial cells encode input more selectively and nonlinearly than deep cells and electroreceptors [28,42,43]; they are tuned to a narrower temporal frequency range that can depend on the stimulus’ spatial extent (i.e. object vs. conspecific) (Figure 3a). Further, superficial pyramidal cells do not respond to self-generated AMs and can adapt to changes in such AMs through synaptic plasticity [10]. In contrast, like electroreceptors, deep cells responses are largely insensitive on the stimulus’ spatial extent and have broad frequency tuning curves [28,42,44]. Further, they display little or no plasticity compared to superficial cells [45]. We describe below the intrinsic and network-based mechanisms that mediate these functional heterogeneities.

Feedback determines functional heterogeneities in ELL
The ELL projects topographically to two contralateral regions: the nucleus praeminentialis (nP, rhombencephalon – feedback) and the torus semicircularis (TS, midbrain – ascending projections). A striking feature of the electrosensory system is that only the deep pyramidal cells project to nP whereas all project to TS [45]. NP mediates direct inhibitory and excitatory feedback to the ELL, as well as indirect feedback to ELL via cerebellar granule cells (Egp).

The excitatory component of the direct feedback may mediate a sensory ‘searchlight’, but this is not further considered here [32]. Further, the diffuse inhibitory component of a direct feedback of nP to ELL enables ‘categorical’ coding that is dependent on the stimulus’
Spatio-temporal content of electrosensory signals varies based on the behavioral context. (a) Images of Apteronotus leptorhynchus (brown ghost knife fish). The electric field generated by Fish 1’s electric organ discharge (EOD) varies sinusoidally in time at a constant high frequency (species range: ~650–1000 Hz) and is seen at the peak of its head negative phase. A prey distorts the EOD to create an ‘electric image’ on the skin. A conspecific (Fish 2) also generates an EOD (not shown) that interferes with fish 1’s EOD. A small dipole and amplifier placed near the skin of Fish 2 shows the electrical signal (black).
Anatomy of the electrosensory system. The image at the right is a side view of the A. leptorhynchus brain with rostral to the right. The key electrosensory structures are the hypertrophied electrosensory lateral line lobe (ELL) within the rhombencephalon and torus semicircularis (TS) within the mesencephalon. The outline of the TS is indicated by the orange ovoid, but this structure is actually located interior to the external optic tectum (TeO). A horizontal slice through the ELL is shown near the middle of the figure. Electroreceptor afferents project only to the ipsilateral ELL where they form four parallel topographic maps, each within its own ELL segment. For each map, the head of the fish is represented rostrally in ELL, while the trunk is represented caudally. The dorso-ventral axis of the fish is represented along the medio-lateral axis of each map. The four segments are: medial (MS), centromedial (CMS), centrolateral (CLS) and lateral (LS). Passive electroreceptors project strictly to MS (not further considered). Each active electroreceptor afferent trifurcates to provide identical input to the CMS, CLS and LS maps. However, as indicated in this figure, the sizes of the maps are very different with CMS > CLS > LS. All four ELL maps converge onto the midbrain TS. Thus, the TS contains a single topographic map of the entire body surface. CCb – corpus cerebellum; Tel – telencephalon.

Figure 2

Spatial extent. Indeed, conspecific-like AMs elicit ELL gamma range oscillatory bursting whereas prey-like AMs do not [46]. A combination of experimental, computational, and theoretical studies have shown that this is because this feedback system exhibits a resonance, which is excited in proportion to the degree of spatial correlation in the stimulus [46,47]. The oscillatory bursting may enhance directionally selective responses to object movement in the TS [48].

A consequence of the anatomical/molecular arrangement of the indirect feedback is that superficial cells reject global spatially redundant low frequency input that can be caused by self-movement, using a ‘teacher’ input from deep cells that is not altered by feedback [45]. Experimental [28,45] and theoretical [49] studies have revealed how this feedback reduces the system’s response to low frequencies in an adaptive manner via burst dependent LTD [45,50].

(Figure 1 Continued) from both fish as indicated in the trace at the bottom. Also shown is time varying amplitude of the signal (red). (b) When two fish with different EOD frequencies (upper traces) are located close to one another, the EODs of each fish (middle traces) show alternating regions of constructive and destructive interference, which leads to a sinusoidal amplitude modulation (SAM, red) or a beat with frequency equal to the difference of the individual fish’s EOD frequencies (lower trace) for the summed signal (black). (c) Weakly electric fish can emit communication signals known as chirps that consist of a transient increase in its EOD frequency (upper traces) when close to a conspecific. As a result, the SAM frequency increases transiently which leads to a phase reset of the beat (lower traces). (d) Electroreceptor afferents respond differentially to time varying AMs characteristic of beats and chirps. For low (<5 Hz) frequencies, the afferent’s baseline firing rate is smoothly modulated by the beat (left panel) and in excellent agreement with the response predicted from a linear system. For high (~90 Hz) frequencies, afferents display nonlinear phase locking (middle panel) in that the afferent only fires near a preferred phase of the SAM; this is quite different than what would be predicted from a linear system (green line). For chirps, the resulting phase reset of the beat is a high frequency stimulus that causes a greater transient increase in firing rate than expected from a linear prediction (green line) as seen in the PSTH. This translates to a transient synchronization at the population level.
Emergence of sparse coding in the hindbrain and further specialization in midbrain. (a) (Top) Tuning curves from example pyramidal neurons from the CMS (left), CLS (middle), LS (right), and TS (far right) under spatially localized (mimicking an object, blue) and spatially diffuse (mimicking a conspecific, red) stimulation. While CMS neurons tend to be tuned to low (∼20 Hz) frequencies under both stimulation geometries, CLS neurons tend to switch their tuning contingent on stimulation geometry. LS neurons are tuned instead to higher (∼80 Hz) frequencies. TS neurons display large heterogeneities in their tuning: some neurons are tuned to low frequencies, others display more broadband tuning while others are tuned to high frequencies. The differential tuning across the ELL segments is manifested in the responses to small chirps (bottom). Indeed, LS neurons show the strongest response while CMS shows the weakest. Some TS neurons respond selectively to chirps. (b) Left: Schematic showing an object (metal bar) moving back and forth.
Across-map differences: intrinsic and network mechanisms lead to differential frequency tuning

Pyramidal cells within all three maps display differential temporal frequency tuning [43,51]. CMS cells are tuned to lower frequencies, and LS cells to higher frequencies (Figure 3d). CLS cells can switch their tuning depending on the stimulus’ spatial extent [52]. Recent studies have shed light on the mechanisms mediating these differences in frequency tuning. Receptive fields of CMS are small, but those of LS cells are significantly larger (160–360 mm²) [34]. As such, CMS cells display better spatial resolution than LS cells. This combined with a higher spike threshold makes LS cells respond preferentially to synchronous electroreceptor input evoked by high frequency spatially diffuse stimuli (e.g. communication) (Figure 3a) [39,40,53]. By contrast, CMS cells will respond preferentially to spatially localized low frequency input (e.g. prey) (Figure 3a).

All pyramidal cells display burst firing that results from somatodendritic interactions [54–56]. A somatic action potential backpropagates to the dendrite and causes a dendritic action potential that in turn propagates back to the soma and causes a depolarizing after-potential (DAP) leading to another somatic action potential. The DAP size grows throughout the burst thereby leading to shorter interspike intervals. The burst terminates when the interspike interval is below the dendritic refractory period [57]. The functional role of burst firing in pyramidal cells is not fully understood as these bursts can sometimes code for low frequency AMs in CMS and CLS cells [44,58] although isolated spikes can also perform this function [59]. Recent studies have shown that these bursts most probably signal the presence of specific features and contain little information in their structure [44,60]. SK channels in LS E-cells cause an AHP that opposes burst firing by cancelling the DAP [61]. This AHP can be overcome during chrip stimulation that thus gives rise to burst firing in these cells [39]. Burst firing is extensively regulated by various mechanisms including neuromodulators such as serotonin [62] and acetylcholine [63,64]. Burst firing is thus most probably used under particular behavioral contexts in order to signal specific stimulus features.

In summary, ELL pyramidal cells begin segregating electroreceptor input into ethologically relevant features related to navigation, predation and communication, and self-movement. To do so, they use a combination of intrinsic and network mechanisms that take advantage partly of the fact that electroreceptors respond differentially to different stimulus classes.

Feature selectivity in the electrosensory midbrain (TS)

The separation of electrocommunication and electrolocation channels has been nearly completed in TS. Indeed, TS neurons generally respond to a much narrower range of spatiotemporal frequencies than ELL pyramidal cells [65–67]. Some are tuned to low frequencies, others to mid-range or even high frequencies (Figure 3a). In addition, some TS neurons respond selectively to communication stimuli [68] (Figure 3a).

Moreover, TS neurons also respond selectively to objects (Figure 3b). Indeed, perception of directional motion is also an important function in weakly electric fish [69] and directional selectivity has recently been shown to be an emergent property in TS [70,71] (Figure 3b) and is summarized in Figure 3c. Although ELL pyramidal cells are not directionally selective, cells with receptive fields at different positions along the fish’s body respond at different phases of the movement, creating a delay between their responses (Figure 3c). Moreover, ELL-TS synapses display differential degrees of synaptic depression along the animal’s body [71]. High-pass filtering caused by synaptic depression can counteract the aforementioned delay, but only in one direction of movement, creating a directional bias. Nonlinear integration via T-type calcium channels further increases this bias [72] (Figure 3c). Ongoing studies are focusing on the role of intrinsic and network mechanisms in the further sparsification of TS responses.

Summary

Through a combination of intrinsic and network properties, electrosensory circuitry progressively segments inputs of varying temporal and spatial scales encountered in different behavioral contexts. This results in the creation of separate neural streams for electrolocation and electrocommunication. Experimental and computational work have revealed general principles of sparse coding that can be summarized as follows:

(Figure 3 Continued) forth sinusoidally along the animal’s rostro-caudal axis. Right: Example responses from ELL (left) and TS (right) neurons to a moving object. ELL neurons strongly respond when the object is at a given position along the animal independently of movement direction. By contrast, TS neurons display directional selectivity and can respond only when the object moves in a given direction. (c) The mechanism by which TS neurons acquire directional selectivity involves combining the responses of two ELL neurons (blue and cyan) whose receptive fields are located at different positions along the animal’s body. The ELL neurons will respond at different phases of the object’s movement in a directionally independent fashion, thereby creating a time delay between their responses. However, differential filtering alters the output of one ELL neuron through high-pass filtering (dark blue) but does not affect the other (cyan), which can reduce the time delay is one direction and increase it in the other. This causes a greater overlap between the ELL neuron responses and thus a greater depolarization in the preferred movement direction and less overlap and depolarization in the other direction. The increased depolarization in the preferred direction can then activate subthreshold T-type calcium channels that then further increase the response in the head-to-tail but not in the tail-to-head direction (orange), thereby explaining the directionally selective responses in TS neurons.
1. Electroreceptors use different strategies to encode different temporal frequencies. They implement noise reduction via negative interval correlations. They respond similarly to self vs. externally generated stimuli and are not tuned to spatial frequency.

2. Within the ELL, ON and OFF center-like cells provide a first step in sparsification by segregating responses to dielectric vs. conductive objects in the environment.

3. Parallel maps in ELL receive the same electroreceptor input and segregate information according to temporal and spatial frequency content. These differences are partly due to intrinsic (e.g. SK channels) and network (e.g. receptive field) mechanisms.

4. Within each ELL map, feedback pathways are used to make some pyramidal cells reject redundant stimuli in an adaptive fashion, and make them respond selectively to stimuli caused by conspecifics. The feedback input originates from a subgroup of dense coding deep pyramidal cells with receptor-like tuning properties and is used to make another subgroup of superficial pyramidal cells code more sparsely. Further, a dendritic backpropagation-based bursting mechanism allows superficial cells to respond preferentially to specific stimuli.

5. Further sparsification occurs in midbrain TS neurons with a near complete separation of responses to electrolocation and electrocommunication stimuli. Further, TS neurons acquire directional selectivity to moving objects by nonlinearly summing differentially filtered inputs from non-directionally selective ELL neurons.

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References and recommended reading
Papers of particular interest, published within the period of review, have been highlighted as:

- of special interest
- of outstanding interest


This paper was inspired by the electroreceptor negative ISI correlations as well as by [35]. It showed that the state of the adaptation process generating negative correlations itself contains information about the sensory input. It further demonstrated that a suitably matched postsynaptic process could extract this information and therefore implement rapid very weak signal detection.


This paper is one of the few that applied Fisher information methods to estimate the physiological requirements for detection of weak signals. In this case very realistic estimates of the E cell receptive field dimensions across the ELL topographic maps were used for the computational analysis. The Fisher information analysis also incorporated detailed physiological data, and concluded that rate coding might not be sufficient for the detection of weak prey signals. This conclusion further inspired the theoretical analysis of [25].


This paper demonstrated that superficial E cells of the lateral ELL segment (LS) were exclusively sensitive to chirps occurring during same sex interactions and responded to them by emitting spike bursts. Burst firing is in general suppressed by an SK channel mediated AHP except during the chirp. This illustrates how a core burst mechanism due to back-propagating dendritic spikes can be modulated at a molecular level (SK channels) to change a neuron’s coding properties. As described in [62], more subtle regulation of sensory responsiveness can then be generated by molecular machinery that modulates the SK-mediated AHP via 5-HT receptors.


This paper demonstrated that a type of chirp used in courtship could be discriminated by the firing patterns of ELL I cells. A key point of this paper was the in vivo demonstration that cellular heterogeneity across I cells was essential for their differential response, and therefore for the ability of the I cell population to discriminate many chirps that occur during opposite sex interactions.


The first paper to explore the functional role of 5-HT in the ELL. The authors show that 5-HT promotes burst firing through downregulation of the AHP, thereby making the cells more responsive to a wide range of stimuli.


The first paper showing that the time constants of synaptic depression vary depending on the stimulus’ location along the animal’s body, thereby creating a directional bias that is exploited by midbrain electrosensory neurons.


The first study showing that subthreshold T-type can act as the necessary nonlinear integrator in order to generate directional selectivity in midbrain neurons.