

Neural assemblies uncovered by generative modeling explain whole-brain activity statistics and reflect structural connectivity

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Abstract

Patterns of endogenous activity in the brain reflect a stochastic exploration of the neuronal state space that is constrained by the underlying assembly organization of neurons. Yet it remains to be shown that this interplay between neurons and their assembly dynamics indeed suffices to generate whole-brain data statistics. Here we recorded the activity from ~40,000 neurons simultaneously in zebrafish larvae, and show that a data-driven generative model of neuron-assembly interactions can accurately reproduce the mean activity and pairwise correlation statistics of their spontaneous activity. This model, the compositional Restricted Boltzmann Machine (cRBM), unveils ~200 neural assemblies, which compose neurophysiological circuits and whose various combinations form successive brain states. We then performed *in silico* perturbation experiments to determine the interregional functional connectivity, which is conserved across individual animals and correlates well with structural connectivity. Our results showcase how cRBMs can capture the coarse-grained organization of the zebrafish brain. Notably, this generative model can readily be deployed to parse neural data obtained by other large-scale recording techniques.

Keywords: Light-Sheet Microscopy, Zebrafish, Neural Assembly, Maximum Entropy, Restricted Boltzmann Machine, Functional Connectivity

1 Introduction

The brain is a highly connected network, organized across multiple scales, from local circuits involving just a few neurons to extended networks spanning multiple brain regions (White et al., 1986; Song et al., 2005; Kunst et al., 2019). Concurrent with this spatial organization, brain activity exhibits correlated firing among large groups of neurons, often referred to as neural assemblies (Harris, 2005). This assembly organization of brain dynamics has been observed in, e.g., auditory cortex (Bathellier et al., 2012), motor cortex (Narayanan et al., 2005), prefrontal cortex (Tavoni et al., 2017), hippocampus (Lin et al., 2005), retina (Shlens et al., 2009), and zebrafish optic tectum (Romano et al., 2015; Mölter et al., 2018; Diana et al., 2019; Triplett et al., 2020). These neural assemblies are thought to form elementary computational units and subserve essential cognitive functions such as short-term memory, sensorimotor computation or decision-making (Hebb, 1949; Gerstein et al., 1989; Harris, 2005; Buzsáki, 2010; Harris, 2012; Palm et al., 2014; Eichenbaum, 2018). Despite the

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prevalence of these assemblies across the nervous system and their role in neural computation, it remains an open challenge to extract the assembly organization of a full brain and to show that the assembly activity state, derived from that of the neurons, is sufficient to account for the collective neural dynamics.

The need to address this challenge is catalyzed by technological advances in light-sheet microscopy, enabling the simultaneous recording of the majority of neurons in the zebrafish brain at single-cell resolution *in vivo* (Panier et al., 2013; Ahrens et al., 2013; Wolf et al., 2015, 2017; Migault et al., 2018; Vanwalleghem et al., 2018). This neural recording technique opens up new avenues for constructing near-complete models of neural activity, and in particular its assembly organization. Recent attempts have been made to identify assemblies using either clustering (Panier et al., 2013; Triplett et al., 2018; Chen et al., 2018; Mölter et al., 2018; Bartoszek et al., 2021), dimensionality reduction approaches (Romano et al., 2015; Mu et al., 2019) or latent variable models (Diana et al., 2019; Triplett et al., 2020), albeit often limited to single brain regions. However, these methods do not explicitly assess to what extent the inferred assemblies could give rise to the observed neural data statistics, which is a crucial property of physiologically meaningful assemblies (Harris, 2005). Here, we address this challenge by developing a generative model of neural activity that is explicitly constrained by the assembly organization, thereby quantifying if assemblies indeed suffice to produce the observed neural data statistics.

Specifically, we formalize neural assemblies using a bipartite network of two connected layers representing the neuronal and the assembly activity, respectively. Together with the maximum entropy principle (Jaynes, 1957; Bialek, 2012), this architecture defines the Restrictive Boltzmann Machine (RBM) model (Hinton & Salakhutdinov, 2006). Here we use an extension to the classical RBM definition termed compositional RBM (cRBM) that we have recently introduced (Tubiana & Monasson, 2017; Tubiana et al., 2019a) and which brings multiple advances to assembly-based network modeling: (1) The maximum entropy principle ensures that neural assemblies are inferred solely from the data statistics. (2) The generative nature of the model, through alternate data sampling of the neuronal and assembly layers, can be leveraged to evaluate its capacity to replicate the empirical data statistics, such as the pairwise co-activation probabilities of all neuron pairs. (3) The cRBM steers the assembly organization to the so-called compositional phase where a small number of assemblies are active at any point in time, making the resulting model highly interpretable as we have shown previously for protein sequence analysis (Tubiana et al., 2019b).

Here, we have successfully trained cRBMs to brain-scale, neuron-level recordings of spontaneous activity in larval zebrafish containing 41000 neurons on average (Panier et al., 2013; Wolf et al., 2017; Migault et al., 2018). This represents an increase of ~ 2 orders of magnitude in number of neurons with respect to previously reported RBM implementations (Köster et al., 2014; Gardella et al., 2017; Volpi et al., 2020), attained through significant algorithmic and computational enhancements. We found that all cells could be grouped into 100-200 partially overlapping assemblies, which are anatomically localized and together span the entire brain, and accurately replicate the first and second order statistics of the neural activity. These assemblies were found to carry more predictive power than a fully connected model which has orders of magnitude more parameters, validating that assemblies underpin collective neural dynamics. Further, the probabilistic nature of our model allowed us to compute a functional connectivity matrix by quantifying the effect of activity perturbations *in silico*. This assembly-based functional connectivity is well-conserved across individual fish and consistent with anatomical connectivity at the mesoscale (Kunst et al., 2019).

In summary, we present an assembly decomposition spanning the zebrafish brain, which accurately accounts for its activity statistics. Our cRBM model provides a widely applicable tool to the community to construct low-dimensional data representations that are defined by the statistics of the data, in particular for very high-dimensional systems. Its generative capability further allows to produce new (synthetic) activity patterns that are amenable to direct *in silico* perturbation and ablation studies.

2 Results

2.1 Compositional RBMs construct Hidden Units by grouping neurons into assemblies

Spontaneous neural activity was recorded from 8 zebrafish larvae aged 5-7 days post fertilization expressing the GCaMP6s or GCaMP6f calcium reporters using light-sheet microscopy (Panier et al., 2013; Wolf et al., 2017; Migault et al., 2018). Each data set contained the activity of a large fraction of the neurons in the brain (40709 \pm 13854; mean \pm standard deviation), which, after cell segmentation, were registered onto the ZBrain atlas (Randlett et al., 2015) and mapzebrain atlas (Kunst et al., 2019). Individual neuronal fluorescence traces were deconvolved to binarized spike trains using blind sparse deconvolution (Tubiana et al., 2020). This data acquisition process is depicted in Figure 1A.

We trained compositional Restricted Boltzmann Machine (cRBM) models to capture the activity statistics of these neural recordings. cRBMs are maximum entropy models, i.e., the maximally unconstrained solution that fits model-specific data statistics (Hinton & Salakhutdinov, 2006; Tubiana & Monasson, 2017; Gardella et al., 2019), and critically extend the classical RBM formulation. Its architecture consists of a bipartite graph where the high-dimensional layer of neurons \mathbf{v} (named ‘visible units’ in RBM terminology) is connected to the low-dimensional layer of latent components, termed Hidden Units (HUs) \mathbf{h} . Their interaction is characterized by a weight matrix \mathbf{W} that is regularized to be sparse. The collection of neurons that have non-zero interactions with a particular HU, noted h_μ (i.e., with $|w_{i,\mu}| > 0$), define its corresponding neural assembly μ (Figure 1B). This weight matrix, together with the neuron weight vector \mathbf{g} and HU potential \mathcal{U} , defines the transformation from the binarized neural activity $\mathbf{v}(t)$ to the continuous HU activity $\mathbf{h}(t)$ (Figure 1B). Figure 1C shows all recorded neurons of a zebrafish brain, color-labeled according to their strongest-connecting HU, illustrating that cRBM-inferred assemblies are typically densely localized in space and together span the entire brain.

Beyond its architecture (Figure 2A), the model is defined by the probability function $P(\mathbf{v}, \mathbf{h})$ of any data configuration (\mathbf{v}, \mathbf{h}) (see Methods for details):

$$P(\mathbf{v}, \mathbf{h}) = \frac{1}{Z} \exp(-E(\mathbf{v}, \mathbf{h})) \quad (1)$$

Where Z is the partition function that normalizes Equation 1 and E is the following energy function:

$$E(\mathbf{v}, \mathbf{h}) = - \sum_i g_i v_i + \sum_\mu \mathcal{U}_\mu(h_\mu) - \sum_{i,\mu} w_{i,\mu} v_i h_\mu \quad (2)$$

HU activity \mathbf{h} is obtained by sampling from the conditional probability function $P(\mathbf{h}|\mathbf{v})$:

$$P(\mathbf{h}|\mathbf{v}) = \prod_{\mu=1}^M P(h_\mu|\mathbf{v}) \propto \prod_{\mu=1}^M \exp\left(-\mathcal{U}_\mu(h_\mu) + h_\mu \cdot \sum_i w_{i,\mu} v_i\right) \quad (3)$$

Conversely, neural activity is obtained from HU activity through:

$$P(\mathbf{v}|\mathbf{h}) = \prod_{i=1}^N P(v_i|\mathbf{h}) \propto \prod_{i=1}^N \exp\left(g_i v_i + v_i \cdot \sum_\mu w_{i,\mu} h_\mu\right) \quad (4)$$

Equations 3 and 4 mathematically reflect the dual relationship between neural and assembly states: the Hidden Units \mathbf{h} drive ‘visible’ neural activity \mathbf{v} , expressed as $P(\mathbf{v}|\mathbf{h})$, while the stochastic assembly activity \mathbf{h} itself is defined as a function of the activity of the neurons: $P(\mathbf{h}|\mathbf{v})$. Importantly, the model does not include direct connections between neurons, hence neural correlations $\langle v_i v_j \rangle$ can arise solely from shared assemblies. Moreover, this bipartite architecture ensures that the conditional distributions factorize, leading to a sampling procedure where all neurons or all HUs can be sampled in parallel. The cRBM leverages this property to efficiently generate new data by Monte Carlo sampling alternately from $P(\mathbf{h}|\mathbf{v})$ and $P(\mathbf{v}|\mathbf{h})$ (Figure 2B).

The cRBM differs from the classical RBM formulation (Hinton & Salakhutdinov, 2006) through the introduction of double Rectified Linear Unit (dReLU) potentials \mathcal{U}_μ , weight sparsity regularization and normalized HU activity (further detailed in Methods). We have previously demonstrated in theory and application (Tubiana & Monasson, 2017; Tubiana et al., 2019a,b) that this new formulation steers the model into the so-called compositional phase, which makes the latent representation highly interpretable. This phase occurs when a limited number m of HUs co-activate such that $1 \ll m \ll M$ where M is the total number of HUs. Thus, each visible configuration is mapped to a specific combination of activated HUs. This contrasts with the ferromagnetic phase ($m \sim 1$) where each HU encodes one specific activity pattern, thus severely limiting the possible number of encoded patterns, or the spin-glass phase ($m \sim M$) where all HUs activate simultaneously, yielding a very complex assembly patchwork (Tubiana & Monasson, 2017). Therefore, the compositional phase can provide the right level of granularity for a meaningful interpretation of the cRBM neural assemblies by decomposing the overall activity as a time-dependent co-activation of different assemblies of interpretable size and extent.

2.2 Trained cRBMs accurately replicate data statistics

cRBM models are trained to maximize the $P(\mathbf{v}, \mathbf{h})$ log-likelihood of the zebrafish data recordings, which is achieved by matching the model-generated statistics $\langle v_i \rangle$, $\langle h_\mu \rangle$ and $\langle v_i h_\mu \rangle$ (the mean neuronal activity, mean HU activity and their correlations, respectively) to the empirical data statistics (Equation 14). In order to optimize the two free parameters of the cRBM model - the sparsity regularization parameter λ and the total number of HUs M - we assessed the cRBM performance for a grid of (λ, M) -values for one data set (fish #3). This analysis yielded an optimum for $\lambda = 0.02$ and $M = 200$ (Fig S1). These values were subsequently used for all recordings, where M was scaled with the number of neurons N .

We trained cRBMs on 70% of the recording length, and compared the statistics of model-generated data to the withheld test data set (the remaining 30% of recording, see Methods for details). After convergence, the cRBM generated data that replicated the training statistics accurately, with normalized Root Mean Square Error (nRMSE) values of $\text{nRMSE}_{\langle v_i \rangle} = 0.11$, $\text{nRMSE}_{\langle h_\mu \rangle} = 0.15$ and $\text{nRMSE}_{\langle v_i h_\mu \rangle} = 0.09$ (Figures 2C-E). Here, nRMSE is normalized such that 1 corresponds to shuffled data statistics and 0 corresponds to the best possible RMSE, i.e., between train and test data.

We further evaluated cRBM performance to assess its ability to capture data statistics that the cRBM was not explicitly trained to replicate: the pairwise correlations between neurons $\langle v_i v_j \rangle$ and the pairwise correlations between HUs $\langle h_\mu h_\nu \rangle$. We found that these statistics were also accurately replicated by model-generated data, with $\text{nRMSE}_{\langle v_i v_j \rangle} = -0.09$ (meaning that the model slightly outperformed the train-test data difference) and $\text{nRMSE}_{\langle h_\mu h_\nu \rangle} = 0.17$ (Figures 2F, G). The fact that cRBM also accurately replicated neural correlations $\langle v_i v_j \rangle$ (Figure 2F) is of particular relevance, since this indicates that (1) the assumption that neural correlations can be explained by their shared assemblies is justified and (2) cRBMs may provide an efficient mean to model neural interactions of such large systems ($N \sim 10^4$) where directly modeling all N^2 interactions would be computationally infeasible or not sufficiently constrained by the available data.

Next, we assessed the reconstruction quality after neural data was compressed by the cRBM low-dimensional bottleneck. This is important to prevent trivial, undesired solutions like $w_{i,\mu} = 0 \forall i, \mu$ which would directly lead to $\langle h_\mu \rangle_{P(\mathbf{v}, \mathbf{h})} = \langle h_\mu \rangle_{\text{data}} = 0$ (potentially because of strong sparsity regularization). Figure 2H shows the distribution of cRBM reconstruction quality of all neurons (in purple), quantified by the normalized log-likelihood (nLLH) such that 0 corresponds to an independent model ($P(v_i(t)) = \langle v_i \rangle$) and 1 corresponds to perfect reconstruction (non-normalized LLH = 0). For comparison, we also reconstructed the neural activity using a fully connected Generalized Linear Model (GLM, see Methods and Figures S2, 2H, blue). The cRBM nLLH distribution is significantly greater than the GLM nLLH distribution (one-sided Mann Whitney U test, $P < 10^{-42}$), with medians $\text{LLH}_{\text{cRBM}} = 0.24$ and $\text{LLH}_{\text{GLM}} = 0.20$. Hence, projecting the neural data onto the low-dimensional representation of the HUs does not compromise the ability to explain the neural activity. In fact, reconstruction quality of the cRBM slightly outperforms the GLM, possibly due to the suppression of noise in the cRBM estimate. The optimal ($\lambda = 0.02$, $M = 200$) choice of free parameters was selected by cross-validating the median of the cRBM reconstruction quality, together with the normalized RMSE of the 5 previously described statistics

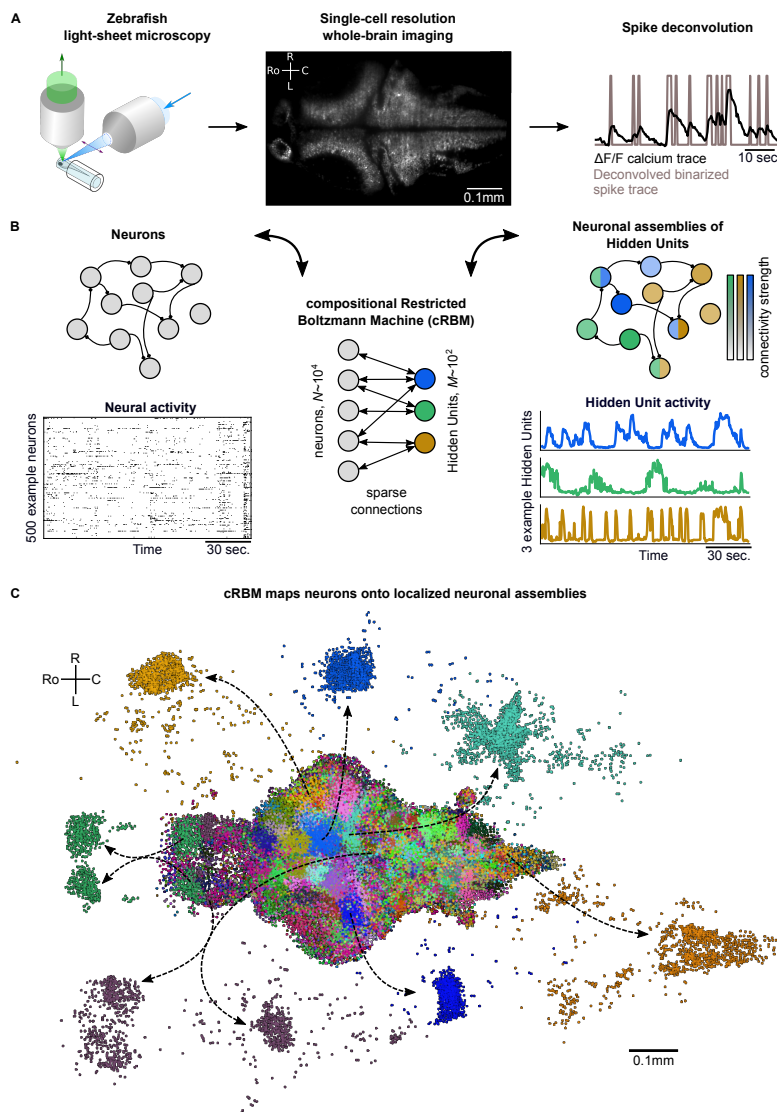


Figure 1: cRBMs construct Hidden Units by grouping neurons into assemblies

(A) The neural activity of zebrafish larvae was imaged using light-sheet microscopy (left), which resulted in brain-scale, single-cell resolution data sets (middle, microscopy image of a single plane shown for fish #1). Calcium activity $\Delta F/F$ was deconvolved to binarized spike traces for each segmented cell (right, example neuron).

(B) cRBM sparsely connects neurons (left) to Hidden Units (HUs, right). The neurons that connect to a HU are defined to be its corresponding assembly (depicted by color labeling, right panel). Data sets typically consist of $N \sim 10^4$ neurons and $M \sim 10^2$ HUs. The activity of 500 randomly chosen example neurons (raster plot, left) and HUs 99, 26, 115 (activity traces, right) of the same time excerpt is shown. HU activity is continuous and is determined by transforming the neural activity of its assembly.

(C) The neural assemblies of an example data set (fish #3) are shown by coloring each neuron according to its strongest-connecting HU. 7 assemblies are highlighted (starting rostrally at the green forebrain assembly, going clockwise: HU 177, 187, 7, 156, 124, 64, 178), by showing their neurons with a connection $|w| \geq 0.1$. See Figure 3 for more anatomical details of assemblies.

R: Right, L: Left, Ro: Rostral, C: Caudal.

(Figure S1).

145 Lastly, we confirmed that the cRBM indeed resides in the compositional phase, characterized by $1 \ll m(t) \ll M$ where $m(t)$ is the number of HUs active at time point t (Figure S3A). This property is a consequence of the sparse weight matrix \mathbf{W} , indicated by its heavy-tail log-distribution (Figure 2I, purple). The compositional

phase is the norm for the presently estimated cRBMs, evidenced by the distribution of median $m(t)$ values for all recordings (average $\frac{\text{median}(m)}{M}$ is 0.26, see Figure S3B). Importantly, the sparse weight matrix does not automatically imply that only a small subset of neurons is connected to the cRBM hidden layer. We validated this by observing that more neurons strongly connect to the hidden layer than expected by shuffling the weight matrix (Figure 2J).

Sparsity facilitated that each assembly only connects to a handful of anatomical regions, as we quantified by calculating the overlap between cRBM assemblies and anatomical regions (Figure S4). We found that cRBM assemblies connect to a median of 3 regions (interquartile range: 2 to 6 regions). Importantly, the cRBM has no information about the locations of neurons during training, so the localization to a limited set of anatomical areas that we observe is extracted from the neural co-activation properties alone. For comparison, Principal Component Analysis (PCA), a commonly used non-sparse dimensionality reduction method that shares the cRBM architecture, naturally converged to a non-sparse weight matrix (Figure 2I, yellow), with fewer connected neurons than expected by shuffling its weight matrix (Figure 2K). This led to unspecific assemblies that are difficult to interpret by anatomy (Figure S4). As a result, sparsity, a cRBM property shared with some other dimensionality reduction techniques, is crucial to interpret the assemblies by anatomy as we demonstrate in the next section. Other clustering or dimensionality reduction methods, such as k-means, PCA and non-negative matrix factorization, have been used previously to cluster neurons in the zebrafish brain (Chen et al., 2018; Mu et al., 2019; Marques et al., 2020). However, because these methods cannot generate artificial neural data using their inferred assemblies, their quality cannot be quantitatively assessed as we have done for the cRBM (but see Tubiana et al. (2019a) for other comparisons).

2.3 cRBM assemblies compose functional circuits and anatomical structures

Above, we have shown that cRBMs converge to sparse weight matrix solutions. This property enables us to visualize the neural assemblies as the collection of significantly connected neurons to an HU. Neurons from a given neural assembly display concerted dynamics, and so one may expect their spatial organization to reflect the neuroanatomy and functional organization of the brain. We here highlight a selection of salient examples of neural assemblies, illustrating that assemblies match well with anatomical structures and functional circuits, while the complete set of neural assemblies is presented in Supplementary Video 1. In particular, we identified assemblies that together compose a neural circuit, are neurotransmitter-specific, encompass a long-range pathway, or can be identified by anatomy. The examples shown here are from a single fish (#3), but results from other fish were comparable.

First, we identified six assemblies that together span the hindbrain circuit that drives eye and tail movements (Dunn et al., 2016; Wolf et al., 2017; Chen et al., 2018). We find two neural assemblies in rhombomere 2 which align with the anterior rhombencephalic turning region (ARTR, (Ahrens et al., 2013; Dunn et al., 2016; Wolf et al., 2017), Figure 3A-B). Each assembly primarily comprises neurons of either the left or right side of the ARTR, but also includes a small subset of contralateral neurons with weights of opposite sign in line with the established mutual inhibition between both subpopulations. Two other symmetric assemblies (Figure 3C, D) together encompass the oculomotor nucleus (nIII) and the contralateral abducens nucleus (nVI, in rhombomere 6), two regions engaged in ocular saccades (Ma et al., 2014) and under the control of the ARTR (Wolf et al., 2017). Additionally, we observed two symmetric assemblies (Figure 3E, F) in the posterior hindbrain (in rhombomere 7), in a region known to drive unilateral tail movements (Chen et al., 2018; Marques et al., 2020) and whose antiphasic activation is also controlled by the ARTR activity (Dunn et al., 2016).

Next, we observed assemblies that correspond to particular neurotransmitter expressions in the ZBrain atlas (Randlett et al., 2015), such as the excitatory Vglut2 (Figure 3G) and inhibitory Gad1b (Figure 3H) neurotransmitters. These assemblies consist of multiple dense loci that sparsely populate the entire brain, confirming that cRBMs are able to capture a large morphological diversity of neural assemblies. Figure 3I depicts another sparse, brain-wide assembly that encompasses the pallium, habenula (Hb) and interpeduncular nucleus (IPN), and thus captures the Hb-IPN pathway that connects to other regions such as the pallium (Beretta et al., 2012;

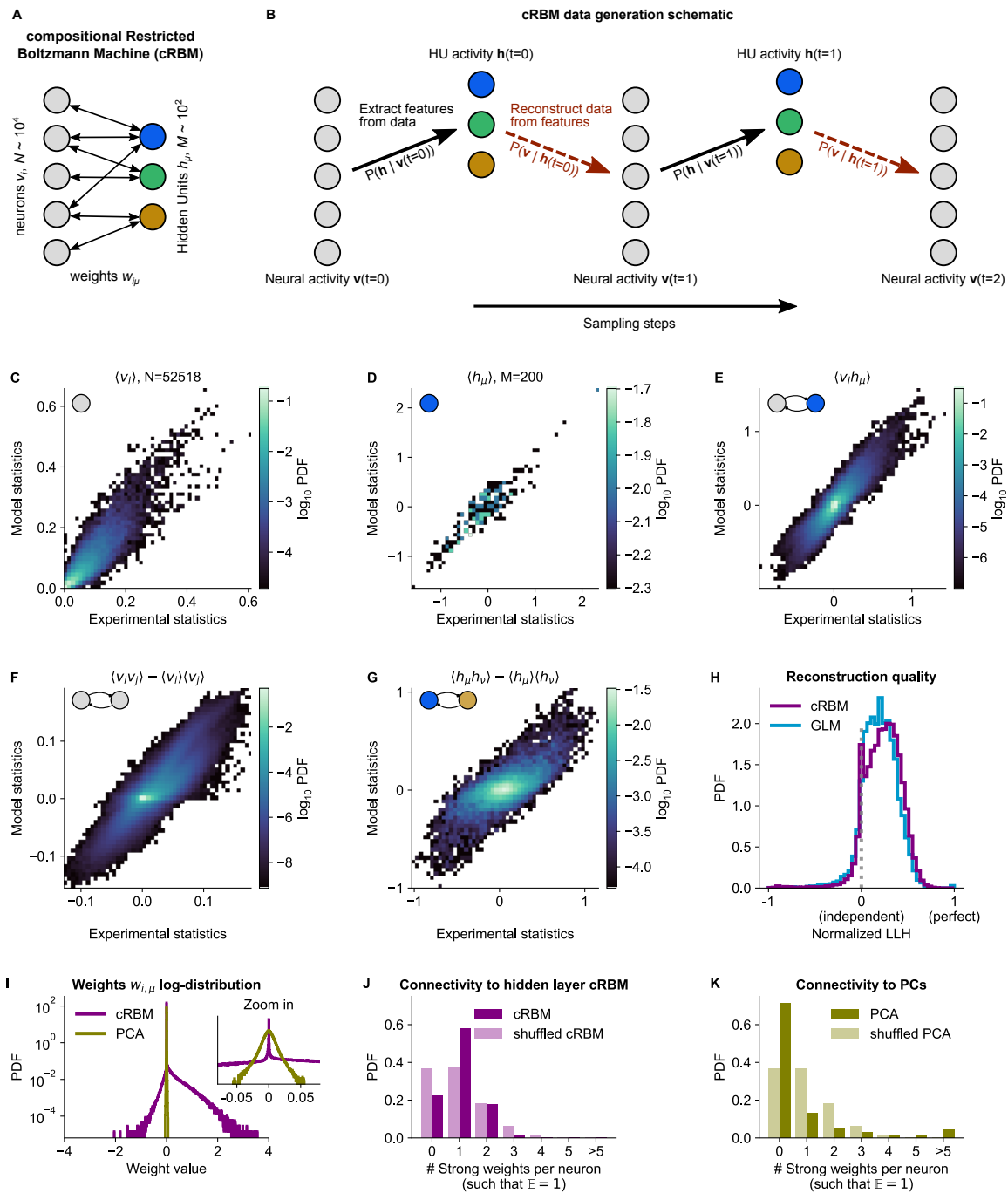


Figure 2: (Legend on next page)

Bartoszek et al., 2021).

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Larger nuclei or circuits were often composed of a small number of distinct neural assemblies with some overlap. For example, the cerebellum was decomposed into multiple, bilateral assemblies (Figure 3J) whereas neurons in the torus semicircularis were grouped per brain hemisphere (Figure 3K). As a last example, the optic

Figure 2: **cRBM is optimized to accurately replicate data statistics**

(A) Schematic of the cRBM architecture, with neurons v_i on the left, HUs h_μ on the right, connected by weights $w_{i\mu}$. (B) Schematic depicting how cRBMs generate new data. The HU activity $\mathbf{h}(t)$ is sampled from the visible unit (i.e., neuron) configuration $\mathbf{v}(t)$, after which the new visible unit configuration $\mathbf{v}(t+1)$ is sampled and so forth.

(C) cRBM-predicted and experimental mean neural activity $\langle v_i \rangle$ were highly correlated (Pearson correlation $r_p = 0.91$, $P < 10^{-307}$) and had low error (nRMSE $_{\langle v_i \rangle} = 0.11$, normalized Root Mean Square Error, see Methods). Data displayed as 2D probability density function (PDF), scaled logarithmically (base 10).

(D) cRBM-predicted and experimental mean Hidden Unit (HU) activity $\langle h_\mu \rangle$ also correlated very strongly ($r_p = 0.93$, $P < 10^{-86}$) and had low nRMSE $_{\langle h_\mu \rangle} = 0.15$ (other details as in C).

(E) cRBM-predicted and experimental average pairwise neuron-HU interactions $\langle v_i h_\mu \rangle$ correlated strongly ($r_p = 0.74$, $P < 10^{-307}$) and had a low error (nRMSE $_{\langle v_i h_\mu \rangle} = 0.09$).

(F) cRBM-predicted and experimental average pairwise neuron-neuron interactions $\langle v_i v_j \rangle$ correlated well ($r_p = 0.58$, $P < 10^{-307}$) and had a low error (nRMSE $_{\langle v_i v_j \rangle} = -0.09$, where the negative nRMSE value means that cRBM-predictions match the test data slightly better than the train data). Pairwise interactions were corrected for naive correlations due to their mean activity by subtracting $\langle v_i \rangle \langle v_j \rangle$.

(G) cRBM-predicted and experimental average pairwise HU-HU interactions $\langle h_\mu h_\nu \rangle$ correlated strongly ($r_p = 0.73$, $P < 10^{-307}$) and had a low error (nRMSE $_{\langle h_\mu h_\nu \rangle} = 0.17$).

(H) The low-dimensional cRBM bottleneck reconstructs most neurons above chance level (purple), quantified by the normalized log-likelihood (nLLH) between neural test data v_i and the reconstruction after being transformed to HU activity (see Methods). Median normalized nLLH $_{\text{cRBM}} = 0.24$. Reconstruction quality was also determined for a fully connected Generalized Linear Model (GLM) that attempted to reconstruct the activity of a neuron v_i using all other neurons \mathbf{v}_{-i} (see Methods). The distribution of 5000 randomly chosen neurons is shown (blue), with median nLLH $_{\text{GLM}} = 0.20$. The cRBM distribution is stochastically greater than the GLM distribution (one-sided Mann Whitney U test, $P < 10^{-42}$).

(I) cRBM (purple) had a sparse weight distribution, but exhibited a greater proportion of large weights $w_{i\mu}$ than PCA (yellow), both for positive and negative weights, displayed in log-probability.

(J) Distribution of above-threshold absolute weights $|w_{i\mu}|$ per neuron v_i (dark purple), indicating that more neurons strongly connect to the cRBM hidden layer than expected by shuffling the weight matrix of the same cRBM (light purple). The threshold Θ was set such that the expected number of above-threshold weights per neuron $\mathbb{E}(\#\mathbf{w}_i > \Theta) = 1$.

(K) Corresponding distribution as in (J) for PCA (dark yellow) and its shuffled weight matrix (light yellow), indicating a predominance of small weights in PCA for most neurons v_i .

All panels of this figure show the data statistics of the cRBM with parameters $M = 200$ and $\lambda = 0.02$ (best choice after cross-validation, see Figure S1) of example fish #3, comparing the experimental test data test and model-generated data after cRBM training converged.

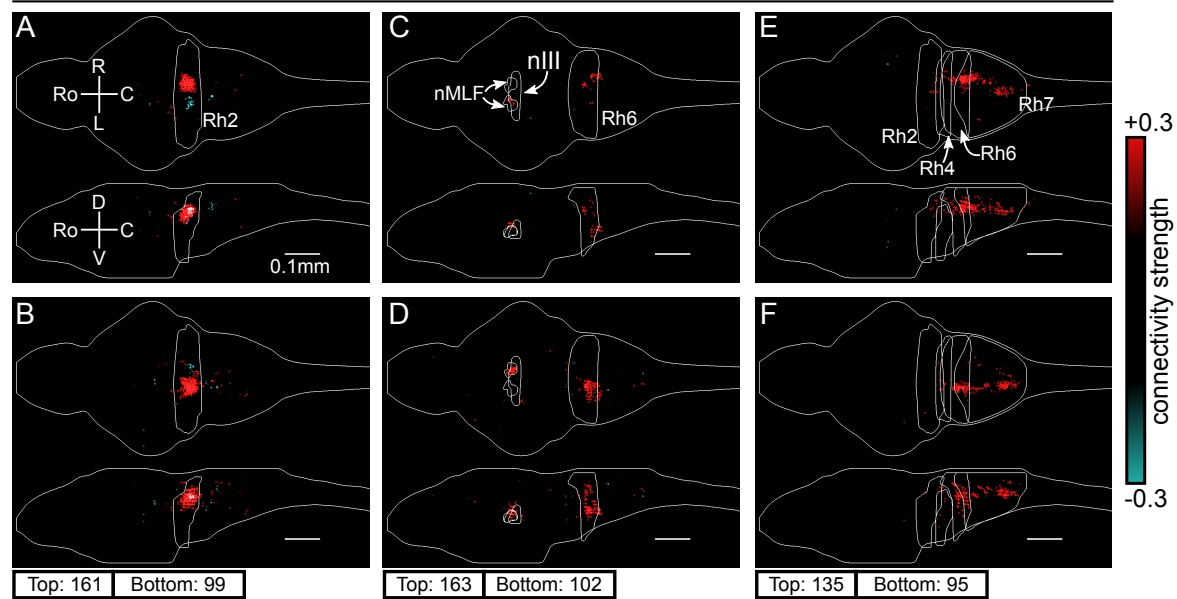
tectum was composed of a larger set of approximately 18 neural assemblies, which spatially tiled the volume of the optic tectum (Figure 3L). This particular organization is suggestive of spatially localized interactions within the optic tectum, and aligns with the morphology of previously inferred assemblies in this specific region (Romano et al., 2015; Diana et al., 2019; Triplett et al., 2020). However, Figure 3 altogether demonstrates that the typical assembly morphology of the optic tectum identified by our and these previous analyses does not readily generalize to other brain regions, where a large range of different assembly morphologies compose neural circuits.

Overall, the clear alignment of cRBM-based neural assemblies with anatomical regions and circuits suggests that cRBMs are able to identify anatomical structures from dynamical activity alone, which enables them to break down the overall activity into parts that are interpretable by physiologists in the context of previous, more local studies.

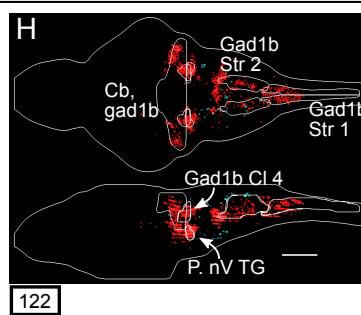
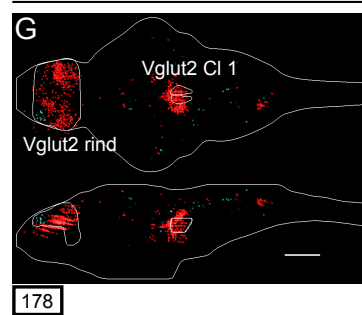
2.4 HU dynamics cluster into groups and display slower dynamics than neurons

HU activity, defined as the expected value of $P(\mathbf{h}|\mathbf{v})$ (Equation 9), exhibits a rich variety of dynamical patterns (Figure 4A). HUs can activate very transiently, slowly modulate their activity, or display periods of active and inactive states of comparable duration. Figure 4B highlights a few HU activity traces that illustrate this diversity of HU dynamics. The top three panels of Figure 4B show the dynamics of the assemblies of Figure 3A-F which encompass the ARTR hindbrain circuit that controls saccadic eye movements and directional tail flips. HUs 99 and 161 drive the left and right ARTR and display antiphasic activity with long dwell times of ~ 15 s, in accordance with previous studies (Ahrens et al., 2013; Dunn et al., 2016; Wolf et al., 2017). HU 102 and 163 correspond to the oculomotor neurons in the nuclei nIII and nVI that together drive the horizontal saccades. Their temporal dynamics are locked to that of the ARTR units in line with the previously identified role of ARTR as a pacemaker for the eye saccades (Wolf et al., 2017). HUs 95 and 135, which drive directional tail

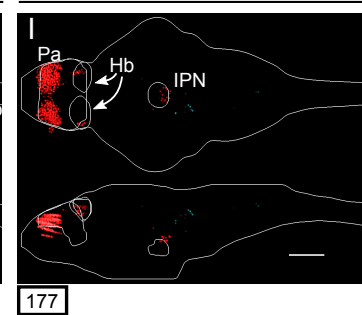
Neural circuit composed of assemblies



Neurotransmitter-specific assemblies



Pathway assembly



Groups of assemblies identified by anatomy

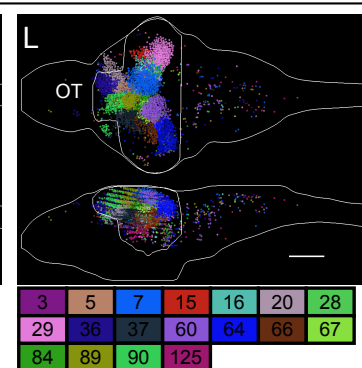
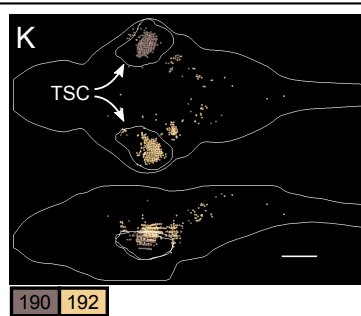
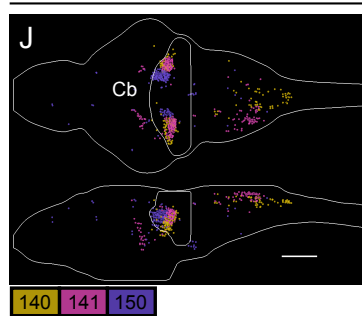


Figure 3: (Legend on next page)

flips, display transient activations that only occur when the ipsilateral ARTR-associated HU is active. This is consistent with the previous finding that the ARTR alternating activation pattern sets the orientation of successive tail flips accordingly (Dunn et al., 2016). The fourth panel shows the traces of the brain-wide assemblies of

Figure 3: **cRBM assemblies compose functional circuits and anatomical structures**

(A-I) Individual example assemblies μ are shown by coloring each neuron i with its connectivity weight value $w_{i,\mu}$ (see color bar at the right hand side). The assembly index μ is stated at the bottom of each panel. The orientation and scale are given in panel A (Ro: rostral, C: caudal, R: right, L: left, D: dorsal, V: ventral). Anatomical regions of interest, defined by the ZBrain Atlas (Randlett et al., 2015), are shown in each panel (Rh: rhombomere, nMLF: nucleus of the medial longitudinal fascicle; nIII: oculomotor nucleus nIII, Cl: cluster; Str: stripe, P. nV TG: Posterior cluster of nV trigeminal motoneurons; Pa: pallium; Hb: habenula; IPN: interpeduncular nucleus). (J-L) Groups of example assemblies that lie in the same anatomical region are shown for cerebellum (Cb), torus semicircularis (TSC) and optic tectum (OT). Neurons i were defined to be in an assembly μ when $|w_{i,\mu}| > 0.15$, and colored accordingly. If neurons were in multiple assemblies shown, they were colored according to their strongest-connecting assembly.

Figures 3G, I, displaying slow tonic modulation of their activity. Finally, the bottom panel, which corresponds to the collective dynamics of assembly 122 (Figure 3H), comprises short transient activity that likely corresponds to fictive swimming events.

Some HUs regularly co-activate, leading to strong correlations between different HUs. This is quantified by their Pearson correlation matrix shown in Figure 4C (top), which reveals clusters of correlated HUs. These were grouped using hierarchical clustering (Figure 4C, bottom), and we then manually identified their main anatomical location (top labels). We further observed that HU activity is bimodal, as evidenced by the distribution of all HU activity traces in Figure 4D. This bimodality can emerge because the dReLU potentials \mathcal{U}_μ (Equation 13) can learn to take different shapes, including a double-well potential that leads to bimodal dynamics (see Methods). This allows us to effectively describe HU activity as a two-state system, where $h_\mu(t) > 0$ increases the probability to spike ($P(v_i(t) = 1)$) for its positively connected neurons, and $h_\mu(t) < 0$ decreases their probability to spike. The binarized neuron activity is also a two-state system (spiking or not spiking), which enabled us to compare the time constants of neuron and HU state changes, quantified by the median time between successive onsets of activity. We find that HUs, which represent the concerted dynamics of neuronal assemblies, operate on a slower time scale than individual neurons (Figure 4E). This observation aligns with the expected difference between cellular and circuit-level time scales.

2.5 cRBM embodies functional connectivity that is strongly correlated across individuals

The probabilistic nature of cRBMs uniquely enables *in silico* perturbation experiments to estimate the functional connection J_{ij} between pairs of neurons, where J_{ij} is quantified by directly perturbing the activity of neuron j and observing the change in probability to spike of neuron i . We first defined the generic, symmetric functional connection J_{ij} using $P(v_i|v_j, v_{k \neq i,j})$ (Equation 15) and then used $P(\mathbf{v})$ (Equation 12) to derive the cRBM-specific J_{ij} (Equation 17, see Methods). Using this definition of J_{ij} , we constructed a full neuron-to-neuron effective connectivity matrix for each zebrafish recording. We then asked whether this cRBM-inferred connectivity matrix was robust across individuals. For this purpose we calculated the functional connections between anatomical regions, given by the assemblies that occupy each region, because neuronal identities can vary across individual specimen. For this purpose we considered anatomical regions as defined by the mapze-brain atlas (Kunst et al., 2019) for which a regional-scale structural connectivity matrix exists to which we will compare our functional connectivity matrix.

We aggregated neurons using the L_1 norm for each pair of anatomical regions to determine the functional connection between regions (see Methods). This led to a symmetrical functional connectivity matrix for each animal, three of which are shown in Figure 5A-C (where non-imaged regions are left blank, and all 8 animals are shown in Figure S5). The strength of functional connections is distributed approximately log-normal (Figure 5D), similar to the distribution of structural region-to-region connections (Kunst et al., 2019). To quantify the similarity between individual fish, we computed the Pearson correlation between each pair of fish. Functional connectivity matrices correlate strongly across individuals, with an average Pearson correlation of 0.69 (Figures 5E and F).

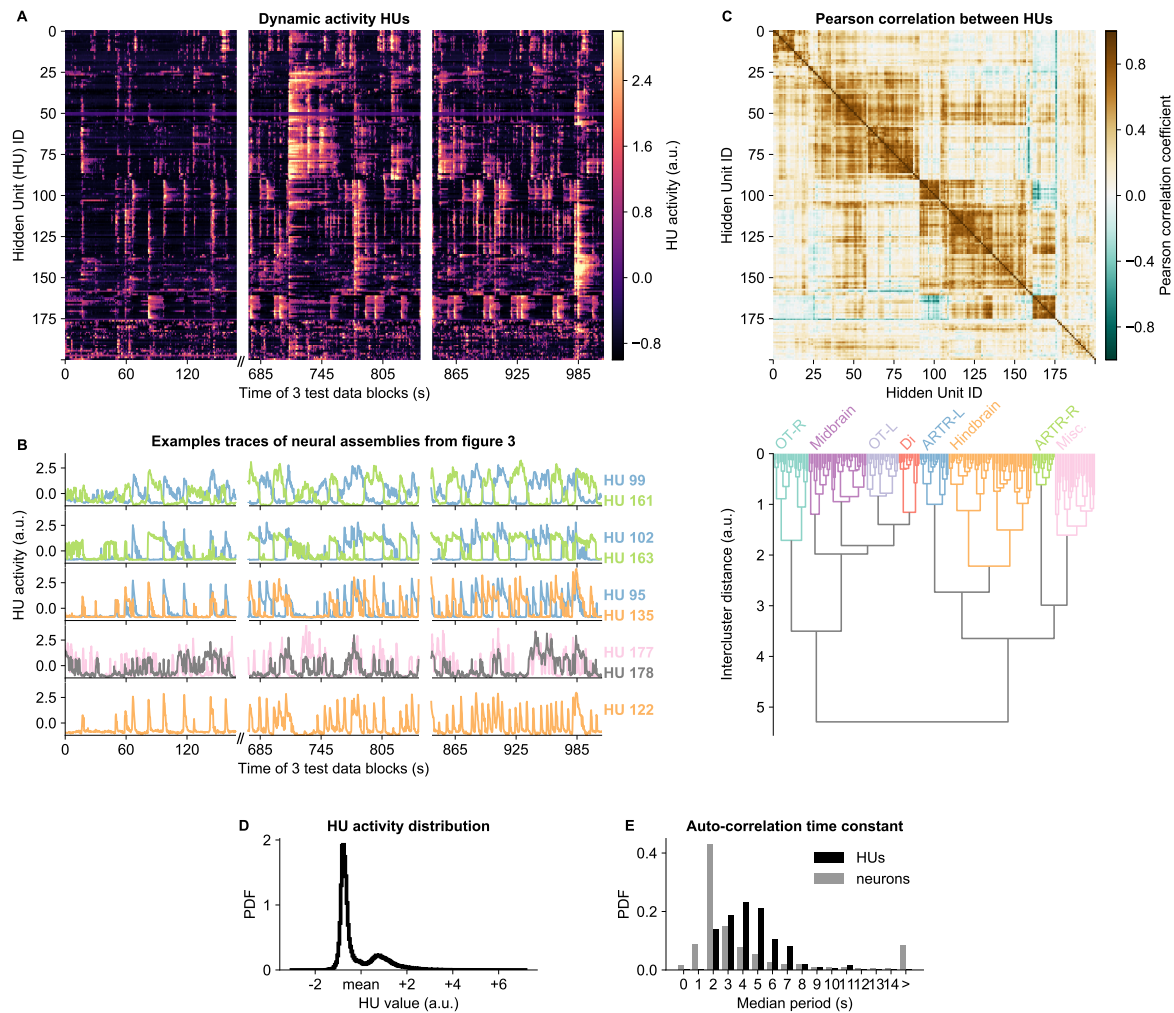


Figure 4: HU dynamics are bimodal and activate slower than neurons

(A) HU dynamics are diverse and are partially shared across HUs. The bimodality transition point of each HU was determined and subtracted individually, such that positive values correspond to HU activation (see Methods). The test data consisted of 3 blocks, with a discontinuity in time between the first and second block (Methods).

(B) Highlighted example traces from panel A. HU indices are denoted on the right of each trace, colored according to their cluster from panel D. The corresponding cellular assemblies of these HU are shown in Figure 3A-I.

(C) Top: Pearson correlation matrix of the dynamic activity of panel A. Bottom: Hierarchical clustering of the Pearson correlation matrix. Clusters (as defined by the colors) were annotated manually. This sorting of HUs is maintained throughout the manuscript. OT: Optic Tectum, Di: Diencephalon, ARTR: anterior rhombencephalic turning region, Misc.: Miscellaneous, L: Left, R: Right.

(D) The distribution of all HU activity values of panel A shows that HU activity is bimodal and sparsely activated (because the positive peak is smaller than the negative peak). PDF: Probability Density Function.

(E) Distribution of the time constants of HUs (black) and neurons (grey). Time constants are defined as the median oscillation period, for both HUs and neurons. An HU oscillation is defined as a consecutive negative and positive activity interval. A neuron oscillation is defined as a consecutive interspike-interval and spike-interval (which can last for multiple time steps, for example see Figure 1A). The time constant distribution of HUs is greater than the neuron distribution (Mann Whitney U test, $P < 10^{-16}$).

We conclude that similar functional circuits spontaneously activate across individuals, despite the limited duration of neural recordings (~25 minutes), which can be identified across fish using independently estimated cRBMs. In the next section we aggregate these individual matrices to a general functional connectivity matrix for comparison with the zebrafish structural connectivity matrix.

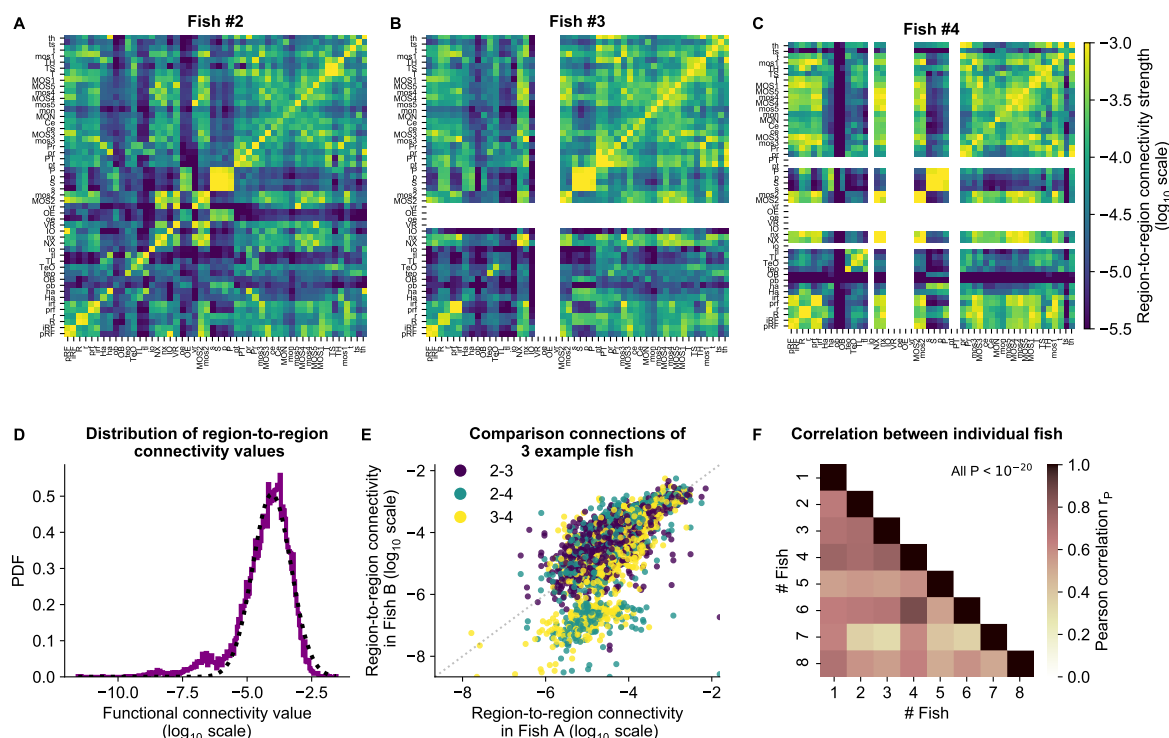


Figure 5: cRBM gives rise to functional connectivity that is strongly correlated across individuals

(A) The functional connectivity matrix between anatomical regions of the mapzebrain atlas (Kunst et al., 2019) of example fish #2 is shown. Functional connections between two anatomical regions were determined by the similarity of the HUs to which neurons from both regions connect to (Methods). Mapzebrain atlas regions with less than 5 imaged neurons were excluded, yielding $N_{\text{MAP}} = 50$ regions in total. See Table S1 for region name abbreviations. The matrix is shown in log₁₀ scale, because functional connections are distributed approximately log-normal (see panel D).

(B) Equivalent figure for example fish #3 (example fish of prior figures).

(C) Equivalent figure for example fish #4. Panels A-C share the same log₁₀ color scale (right).

(D) Functional connections are distributed approximately log-normal. (Mutual information with a log-normal fit (black dashed line) is 3.83, while the mutual information with a normal fit is 0.13). All connections of all 8 fish are shown, in log₁₀ scale (purple).

(E) Functional connections of different fish correlate well, exemplified by the 3 example fish of panels A-C. All non-zero functional connections (x-axis and y-axis) are shown, in log₁₀ scale. Pearson correlation r_P between pairs: $r_P(\#2, \#3) = 0.73$, $r_P(\#2, \#4) = 0.73$, $r_P(\#3, \#4) = 0.78$. All correlation P values $< 10^{-20}$ (two-sided t-test).

(F) Pearson correlations r_P of region-to-region functional connections between all pairs of 8 fish. For each pair, regions with less than 5 neurons in either fish were excluded. All P values $< 10^{-20}$ (two-sided t-test), and average correlation value is 0.69.

2.6 cRBM-inferred functional connectivity reflects structural connectivity

In the previous section we have determined the functional connections between anatomical regions using the cRBM assembly organization. Although functional connectivity stems from the structural (i.e., biophysical) connections between neurons, it can reflect correlations that arise through indirect network interactions (Bassett & Sporns, 2017; Das & Fiete, 2020). Using recently published structural connectivity data of the zebrafish brain (Kunst et al., 2019), we are now able to quantify the overlap between a structurally defined connectivity matrix and our functional connectivity matrix estimated through neural dynamics. Kunst et al. (2019) determined a zebrafish structural connectivity matrix between 72 anatomical regions using structural imaging data from thousands of individually Green Fluorescent Protein (GFP)-labeled neurons from multiple animals. We slightly extended this matrix by using the most recent data, filtering indirect connections and accounting for the resulting sampling bias (Figure 6A, regions that were not imaged in our light-sheet microscopy experiments were excluded). Next, we aggregated the functional connectivity matrices of all our calcium imaging recordings to one grand average functional connectivity matrix (Figure 6B).

For comparison, we also calculated the connectivity matrices defined by either covariance or Pearson correlation (Figure S6). The cRBM functional connectivity spans a larger range of values than either of these methods, leading to a more fine-grained connectivity matrix akin to the structural connectivity map (Figure 6B). This greater visual resemblance was statistically confirmed by calculating the Spearman correlation between structural and functional connectivity, which is greater for cRBM ($r_s = 0.39$, Figure 6C), than for covariance-based connectivity ($r_s = 0.18$, Figure S6 left) or correlation-based connectivity ($r_s = 0.26$, Figure S6 right). Hence, using recordings of ~ 25 minutes on average, cRBMs were able to identify functional connections that resemble the anatomical connectivity between brain regions. Strong or weak functional connections are predictive of present or absent structural connections respectively (Figure 6D), and could thus potentially be used for inference in systems where the structural connectivity pattern is unknown.

3 Discussion

We have developed a cRBM model that accurately replicated the data statistics of brain-scale zebrafish recordings, thereby forming neural assemblies that spanned the entire brain. The objective of our study was threefold: first, to show that the cRBM model can be applied to high-dimensional data, such as whole-brain recordings, second, to prove that an assembly-based model is sufficient to generate whole-brain neural data statistics, and third, to describe the physiological properties of the assembly organization in the zebrafish brain and use it to create a functional connectivity map. We have shown that, after convergence, the cRBM-generated data not only replicated the data statistics that it was constrained to fit, but also extrapolated to fit the pairwise correlation statistics of neurons and HUs, leading to a better reconstruction of neural data than a fully connected GLM (Figure 2). These results thereby quantify how neural assemblies play a major role in determining the collective dynamics of the brain. To achieve this, cRBMs formed sparsely localized assemblies that spanned the entire brain, facilitating their biological interpretation (Figure 3, S4, 4). Further, the probabilistic nature of the cRBM model allowed us to create a mesoscale functional connectivity map that was largely conserved across individual fish and correlated well with structural connectivity (Figures 5, 6).

The maximum entropy principle underlying the cRBM definition has been a popular method for inferring pairwise effective connections between neurons or assemblies of co-activating cells (Schneidman et al., 2006; Tavoni et al., 2017; Ferrari et al., 2017; Meshulam et al., 2017; Posani et al., 2018; Chen et al., 2019). However, its computational cost has limited this pairwise connectivity analysis to typically $\sim 10^2$ neurons. The two-layer cRBM model that we used here alleviates this burden, because the large number of neuron-to-neuron connections are no longer explicitly optimized, which enables a fast data sampling procedure (Figure 2B). However, we have shown that these connections are still estimated indirectly with high accuracy via the assemblies they connect to (Figure 2F). We have thus shown that the cRBM is able to infer the $\frac{1}{2}N^2 \approx 10^9$ (symmetric) pairwise connections through its assembly structure, a feat that is computationally infeasible for many other methods.

Previously, we have extensively compared cRBM performance to other dimensionality reduction techniques, including Principal Component Analysis (PCA), Independent Component Analysis (ICA), Variational Autoencoders (VAEs) and their sparse variants, using protein sequence data as a benchmark (Tubiana et al., 2019a). Briefly put, we showed that PCA and ICA could not accurately model the system due to their deterministic nature, putting too much emphasis on low-probability high-variance states, while VAEs were unable to capture all features of data due to the unrealistic assumption of independent, Gaussian-distributed latent variables. Additionally, while PCA has previously been successful in describing zebrafish neural dynamics in terms of their main covariances modes (Ahrens et al., 2012; Marques et al., 2020), we show here that it is not appropriate for assembly extraction due to the absence of both a compositional and stochastic nature (Figures 2, S4). Furthermore, we have shown that the generative component of cRBM models is essential for quantitatively assessing that the assembly organization is sufficient for reproducing neural statistics (Figure 2), moving beyond deterministic clustering analyses such as k-means (Panier et al., 2013; Chen et al., 2018), similarity graph clustering

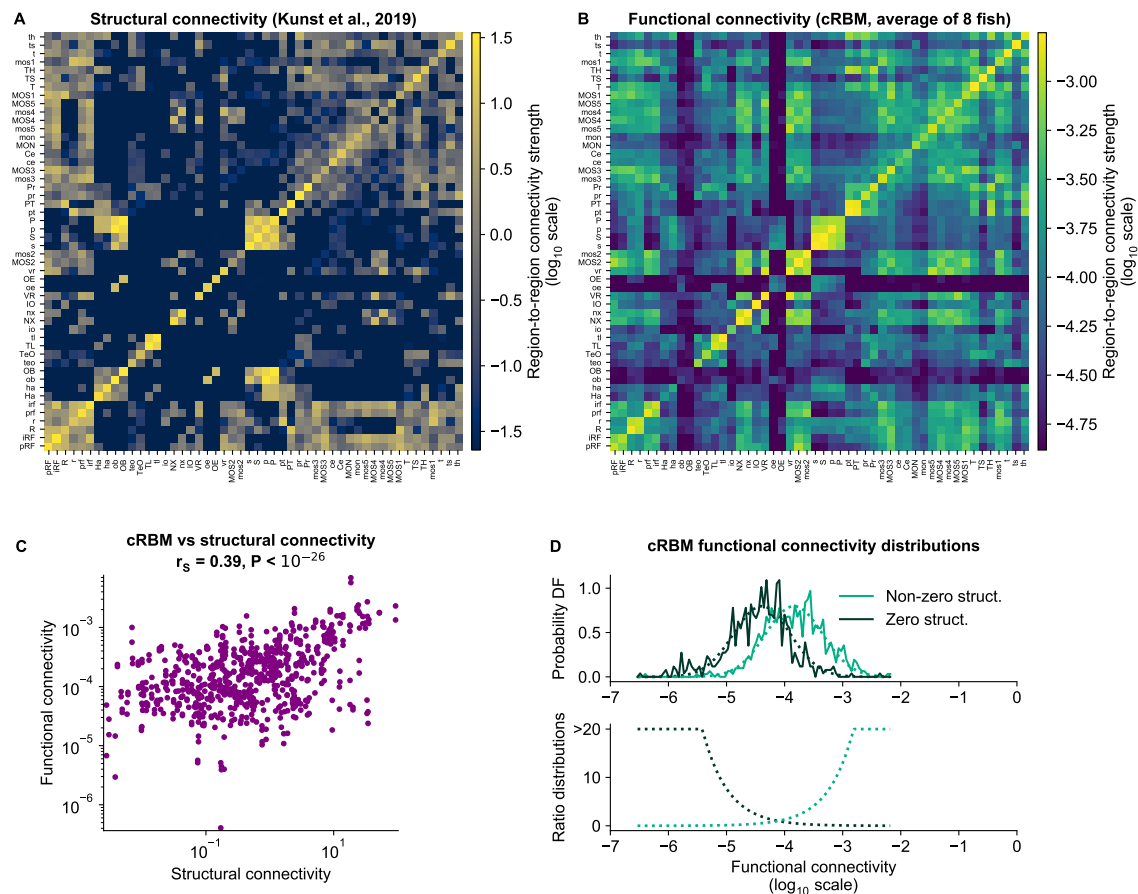


Figure 6: cRBM-inferred functional connectivity reflects structural connectivity

(A) Structural connectivity matrix is shown in \log_{10} scale, adapted from Kunst et al. (2019). Regions that were not imaged in our experiments were excluded (such that $N_{\text{MAP}} = 50$ out of 72 regions remain). Regions (x-axis and y-axis) were sorted according to Kunst et al. (2019). Additional structural data was added and the normalization procedure was updated to include within-region connectivity (see Methods). See Table S1 for region name abbreviations.

(B) Average functional connectivity matrix is shown in \log_{10} scale, as determined by averaging the cRBM functional connectivity matrices of all 8 fish (see Methods). The same regions (x-axis and y-axis) are shown as in panel A.

(C) The average functional and structural connectivity of panels A and B correlate well, with Spearman correlation $r_s = 0.39$ ($P < 10^{-20}$, two-sided t-test). Each data point corresponds to one region-to-region pair. Data points for which the structural connection was exactly 0 were excluded (see panel D for their analysis).

(D) The distribution of average functional connections of region pairs with non-zero structural connections is greater than functional connections corresponding to region pairs without structural connections ($P < 10^{-15}$, two-sided Kolmogorov-Smirnov test). The bottom panel shows the evidence for inferring either non-zero or zero structural connections, defined as the fraction between the PDFs of the top panel (fitted Gaussian distributions were used for denoising).

(Mölter et al., 2018) or non-negative matrix factorization (Mu et al., 2019).

After having quantitatively validated the resultant assemblies, we moved to discussing the biological implications of our findings. Previous studies of the zebrafish optic tectum have identified neural assemblies that were spatially organized into single dense clusters of cells (Romano et al., 2015; Diana et al., 2019; Triplett et al., 2020). We have replicated these findings by observing the distinct organization of ball-shaped assemblies in the optic tectum (Figure 3L). However, our data extends to many other anatomical regions in the brain, where

we found that assemblies can be much more dispersed, albeit still locally dense, consisting of multiple clusters of neurons (Figure 3).

cRBM allowed us to compute the effective, functional connections between each pair of neurons, aggregated to functional connections between each pair of regions, by perturbing neural activity *in silico*. Importantly, we found that this region-scale connectivity is well-conserved across specimen. This observation is non-trivial because each recording only lasted ~25 minutes, which represents a short trajectory across accessible brain states. It suggests that, although each individual brain may be unique at the neuronal scale, the functional organization could be highly stereotyped at a sufficiently coarse-grained level.

It would be naive to assume that these functional connections equate biophysical, structural connections (Das & Fiete, 2020). Both represent different, yet interdependent aspects of the brain organization. Indeed, we found that structural connectivity is well-correlated to functional connectivity, confirming that functional links are tied to the structural blueprint of brain connectivity (Figure 6). Furthermore, strong (weak) functional connections are predictive of present (absent) structural connections between brain regions, though intermediate values are ambiguous.

It will be crucial to synergistically merge structural and dynamic information of the brain to truly comprehend brain-wide functioning (Bargmann & Marder, 2013; Kopell et al., 2014). Small brain organisms are becoming an essential means to this end, providing access to a relatively large fraction of cells (Ahrens & Engert, 2015). To generate new scientific insights it is thus essential to develop analytical methods that can scale with the rapidly growing size of both structural and dynamic data (Helmstaedter, 2015; Ahrens, 2019). In this study we have established that the cRBM can model high-dimensional data accurately, and that its application to zebrafish recordings was crucial to unveil their brain-scale assembly organization. In future studies, cRBMs could be used to generate artificial data whose statistics replicate those of the zebrafish brain. This could be used for further *in silico* ablation and perturbation studies with strong physiological footing, crucial for developing hypotheses for future experimental work (Jazayeri & Afraz, 2017; Das & Fiete, 2020). Lastly, the application of cRBMs is not specific to calcium imaging data, and can therefore be readily applied to high-dimensional neural data obtained by other recording techniques.

4 Acknowledgements

T.L.v.d.P. had an Erasmus+ fellowship (European Union) and acknowledges support from the Biotechnology and Biological Sciences Research Council (BBSRC, grant No. BB/M011224/1). J.T. acknowledges support from the Edmond J. Safra Center for Bioinformatics at Tel Aviv University and from the Human Frontier Science Program (cross-disciplinary postdoctoral fellowship LT001058/2019-C). G.M. was funded by a PhD fellowship from the Doctoral School in Physics, Ile de France (EDPIF). G.L.G. had a PhD fellowship from the Systems Biology Network of Sorbonne Université. B.E. and T.L.v.d.P. were supported by an NWO-VIDI Grant.

Funding sources: European Research Council (ERC) under the European Union's Horizon 2020 research and innovation program grant agreement number 715980. Human Frontier Science Program (RGP0060/2017). The French Research National Agency under grant No. ANR-16-CE16-0017. Dutch Institute for Scientific Research NWO (Nederlandse Organisatie voor Wetenschappelijk Onderzoek) grant No. 016.VIDI.189.052.

We thank the IBPS fish facility staff for the fish maintenance, in particular Stéphane Tronche and Alex Bois. We are grateful to Carounagarane Dore for his contribution to the design of the experimental setup. We thank Misha Ahrens for providing the GCaMP lines. We are also grateful to Rémi Monasson for very fruitful discussions and his comments on the manuscript.

5 Author Contributions

T.L.v.d.P., J.T., V.B., B.E. and G.D. conceived the project, analyzed data and wrote the manuscript. T.L.v.d.P. and J.T. developed the cRBM code, performed the various analyses and wrote the original draft. G.L.G. and G.M. carried out the functional neural recording experiments and associated data pre-processing. M.K. and H.B. produced the structural connectivity data. V.B., B.E. and G.D. supervised the work. All authors contributed to the writing and editing of the manuscript.

6 Declaration of Interests

The authors declare no competing interests.

7 Methods

7.0.1 Data and Code Availability

The cRBM model has been developed in Python 3.7 and is available at: <https://github.com/jertubiana/PGM>
An extensive example notebook that implements this model is provided [here](#).

Calcium imaging data pre-processing was performed in MATLAB (Mathworks) using previously published protocols and software (Panier et al., 2013; Wolf et al., 2017; Migault et al., 2018; Tubiana et al., 2020). The functional data recordings, the trained cRBM models and the structural connectivity matrix will be made publicly available after publication.

Figures of neural assemblies or neurons (Figure 1, 3) were made using the Fishualizer, which is a 4D (space + time) data visualization software package that we have previously published (Migault et al., 2018), available at https://bitbucket.org/benglitz/fishualizer_public
Minor updates were implemented to tailor the Fishualizer for viewing assemblies, which can be found at https://bitbucket.org/benglitz/fishualizer_public/src/assembly_viewer

All other data analysis and visualization was performed in Python 3.7 using standard packages (numpy (Harris et al., 2020), scipy (Virtanen et al., 2020), scikit-learn (Pedregosa et al., 2011), matplotlib (Hunter, 2007), pandas (McKinney et al., 2010), seaborn (Waskom, 2021), h5py). The corresponding code is available from the authors on request.

7.0.2 Zebrafish larvae

Experiments were conducted on nacre mutants, aged 5-7 days post-fertilization (dpf). Larvae were reared in Petri dishes at 28°C in embryo medium (E3) on a 14/10h light/dark cycle, and were fed powdered nursery food every day from 6 dpf. They were expressing either the calcium reporter GCaMP6s (fish 1-4, 6 and 8) or GCaMP6f (fish 5 and 7) under the control of the nearly pan-neuronal promoter *elavl3* expressed in the nucleus *Tg(elavl3:H2B-GCaMP6)*. Both lines were provided by Misha Ahrens and published by Vladimirov et al. (2014) (H2B-GCaMP6s) and Quirin et al. (2016) (H2B-GCaMP6f). Experiments were approved by Le Comité d'Éthique pour l'Experimentation Animale Charles Darwin C2EA-05 (02601.01).

7.0.3 Light-sheet microscopy of zebrafish larvae

Spontaneous neural activity (i.e., in the absence of sensory stimulation) was recorded in larval zebrafish using light-sheet microscopy, which acquires brain-scale scans by imaging multiple z-planes sequentially (Panier et al., 2013; Wolf et al., 2017; Migault et al., 2018). Larvae were placed in 2% low melting point agarose (Sigma-Aldrich), drawn tail-first into a glass capillary tube with 1mm inner diameter via a piston and placed in chamber filled with E3 in the microscope. Recordings were of length 1514 ± 238 seconds (mean \pm standard

deviation), with a brain volume imaging frequency of 3.9 ± 0.8 Hz.

The following imaging pre-processing steps were performed offline using MATLAB, in line with previously reported protocols (Panier et al., 2013; Migault et al., 2018). Automated cell segmentation was performed using a watershed algorithm (Panier et al., 2013; Migault et al., 2018) and fluorescence values of pixels belonging to the same neuron was averaged to obtain cell measurements. The fluorescence intensity values F were normalized to $\Delta F/F = (F - \langle F \rangle) / (\langle F \rangle - F_0)$ where $\langle F \rangle$ is the baseline signal per neuron and F_0 is the overall background intensity (Migault et al., 2018). The $\Delta F/F$ activity of different imaging planes was subsequently temporally aligned using interpolation (because of the time delay between imaging planes) (Migault et al., 2018) and deconvolved to binarized spike traces using Blind Sparse Deconvolution (BSD) (Tubiana et al., 2020). BSD estimates the most likely binary spike trace by minimizing the L_2 norm of the difference between the estimated spike trace convolved with an exponential kernel and the ground-truth calcium data, using L_1 sparsity regularization and online hyperparameter optimization. Calcium kernel time constants used for deconvolution were inferred using BSD on the spontaneous activity of three different fish (approximately 5000 neurons per fish, recorded at 10Hz, previously reported by Migault et al. (2018)). For the GCaMP6s line, we used a rise time of 0.2s and a decay time of 3.55s; for the GCaMP6f line, we used 0.15s and 1.6s, respectively.

Brain activity was recorded of 15 animals in total. Of these recordings, 1 was discarded because of poor image quality and 6 were discarded because neurons were inactive (defined by less than 0.02 spikes/(neurons \times time points)), hence leaving 8 data sets for further analysis. The recorded brains were then registered onto the ZBrain Atlas (Randlett et al., 2015) and the mapzebrain atlas (Kunst et al., 2019) for anatomical labeling of neurons (Migault et al., 2018). The ZBrain Atlas was used in figures 1-4 because of its detailed region descriptions (outlining 294 regions in total). However, we also registered our data to the mapzebrain atlas (72 regions in total) in order to compare our results with the structural connectivity matrix which was defined for this atlas only (Kunst et al., 2019). Only neurons that were registered to at least 1 ZBrain region were used for analysis (to filter imaging artefacts). This resulted in 40709 ± 13854 neurons per recording (mean \pm standard deviation, minimum = 23446, maximum = 65517).

7.0.4 Maximum Entropy Principle

Here we provide in brief the general derivation of the class of maximum entropy probabilistic models. Restricted Boltzmann Machines are an instance of this model, which is detailed in the following sections. The maximum entropy principle is used to create probabilistic models $P(x)$ (where x denotes one data configuration sample) that replicate particular data statistics f_k , but are otherwise most random, and therefore least assumptive, by maximizing their entropy $H = -\sum_x P(x) \log(P(x))$ (Gardella et al., 2019). The goal of the model is to match its model statistics $\langle f_k \rangle_{\text{model}} = \sum_x P(x) f_k(x)$ to the empirical data statistics $\langle f_k \rangle_{\text{data}} = F_k$. This is done using Lagrange multipliers Λ_k :

$$\tilde{H} = -\sum_x P(x) \log(P(x)) - \sum_k \Lambda_k \left(\sum_x P(x) f_k(x) - F_k \right) \quad (5)$$

which yields, when \tilde{H} is maximized with respect to $P(x)$, the Boltzmann distribution (see, e.g., Bialek (2012) for a full derivation):

$$P(x) = \frac{1}{Z} \exp \left(-\ln 2 \sum_k \Lambda_k f_k(x) \right) = \frac{1}{Z} \exp(-E(x)) \quad (6)$$

where $E(x)$ is defined as the resulting energy function. Importantly, the data dependency (F_k) disappears when going from Equation 5 to Equation 6. Hence, the maximum entropy principle only defines the shape of the distribution $P(x)$, but not its specific parameters Λ_k (Bialek, 2012). In the case of RBM, these are then optimized using maximum likelihood estimation, as detailed in the sections below.

7.0.4.1 Motivation for choice of statistics

The derivation above describes the general maximum entropy model for a set of statistics $\{f_k\}$. The objective of this study is to extract the assembly structure from neural data, therefore creating two layers: a visible (neural data) layer $\mathbf{v} = (v_1, v_2, \dots, v_N)$ and a hidden (latent) layer $\mathbf{h} = (h_1, h_2, \dots, h_M)$. The model should capture the mean activity of each neuron $\langle v_i \rangle$, their pairwise correlations $\langle v_i v_j \rangle$, the neuron-HU interactions $\langle v_i h_\mu \rangle$ and a function of h_μ . The latter is determined by the potential \mathcal{U} , which we set to be a double Rectified Linear Unit (dReLU), as motivated in the following sections. Fitting all N^2 pairwise interactions $\langle v_i v_j \rangle$ is computationally infeasible, but under the cell assembly hypothesis we assume that this should not be necessary because collective neural behavior is expected to be explained by membership to similar assemblies via $\langle v_i h_\mu \rangle$, and can therefore be excluded. We later show that pairwise correlations are indeed optimized implicitly (Figure 2). All other statistics are included and therefore explicitly optimized, also see Equation 14.

7.0.5 Restricted Boltzmann Machines

A Restricted Boltzmann Machine (RBM) is an undirected graphical model defined on a bipartite graph (Smolensky, 1986; Hinton, 2002; Hinton & Salakhutdinov, 2006), see Figure 2A. RBMs are constituted by two layers of random variables, neurons \mathbf{v} and Hidden Units (HUs) \mathbf{h} , which are coupled by a weight matrix \mathbf{W} . There are no direct couplings between pairs of units within the same layer. Here, each visible unit v_i corresponds to a single recorded neuron with binary (spike-deconvolved) activity ($v_i(t) \in \{0, 1\}$). Each Hidden Unit (HU) h_μ corresponds to the (weighted) activity of its neural assembly and is chosen to be real-valued. The joint probability distribution $P(\mathbf{v}, \mathbf{h})$ writes (Hinton & Salakhutdinov, 2006; Tubiana & Monasson, 2017):

$$P(\mathbf{v}, \mathbf{h}) = \frac{1}{Z} \exp(-E(\mathbf{v}, \mathbf{h})) = \frac{1}{Z} \exp\left(\sum_i g_i v_i - \sum_\mu \mathcal{U}_\mu(h_\mu) + \sum_{i,\mu} w_{i,\mu} v_i h_\mu\right) \quad (7)$$

Where E is the energy function and $Z = \sum_{\mathbf{v}} \int_{\mathbf{h}} d\mathbf{v} d\mathbf{h} \cdot \exp(-E(\mathbf{v}, \mathbf{h}))$ is the partition function. The weights g_i and potentials \mathcal{U}_μ control the activity level of the visible units and the marginal distributions of the HUs respectively, and the weights $w_{i,\mu}$ couple the two layers.

7.0.5.1 From data to features

Given a visible layer configuration \mathbf{v} , a HU h_μ receives the input $I_\mu(\mathbf{v}) = \sum_i w_{i,\mu} v_i \equiv \mathbf{w}_\mu^T \mathbf{v}$ and, owing to the bipartite architecture, the conditional distribution $P(\mathbf{h}|\mathbf{v})$ factorizes as:

$$P(\mathbf{h}|\mathbf{v}) = \prod_\mu P(h_\mu|\mathbf{v}) = \prod_\mu \exp\left(-\mathcal{U}_\mu(h_\mu) + h_\mu I_\mu(\mathbf{v}) - \Gamma_\mu(I)\right) \quad (8)$$

Where $\Gamma_\mu(I) = \log\left(\int_{\mathbf{h}} d\mathbf{h} \cdot \exp(-\mathcal{U}_\mu(h) + hI)\right)$ is the cumulant generating function associated to the potential \mathcal{U}_μ that normalizes Equation 8 (Tubiana et al., 2019b). The average activity of HU h_μ associated to a visible configuration \mathbf{v} is given by a linear-nonlinear transformation (as defined by the properties of the cumulant generating function):

$$\langle h_\mu | \mathbf{v} \rangle = \frac{\partial \Gamma_\mu(I_\mu(\mathbf{v}))}{\partial I} = \Gamma'_\mu(\mathbf{w}_\mu^T \mathbf{v}) \quad (9)$$

Throughout the manuscript, we use this definition to compute HU activity $h_\mu(t) = \langle h_\mu | \mathbf{v}(t) \rangle$ (e.g., in Figure 4).

7.0.5.2 From features to data

Conversely, given a hidden layer configuration \mathbf{h} , a visible unit v_i receives the input $I_i(\mathbf{h}) = \sum_\mu w_{i,\mu} h_\mu \equiv \mathbf{w}_i^T \mathbf{h}$ and the conditional distribution factorizes as:

$$P(\mathbf{v}|\mathbf{h}) = \prod_i P(v_i|\mathbf{h}) \propto \prod_i \exp\left((g_i + I_i(\mathbf{h}))v_i\right) \quad (10)$$

and the average sampled v_i activity is given by:

$$\langle v_i | \mathbf{h} \rangle = \sigma(\mathbf{w}_i^T \mathbf{h} + g_i) \quad (11)$$

Where $\sigma(x) = 1/(1 + e^{-x})$ is the logistic function. Hence, a sampled visible layer configuration \mathbf{v} is obtained by a weighted combination of the HU activity followed by Bernoulli sampling. RBMs are generative models, in the sense that they can generate new, artificial data using Equations 8 and 10. Figure 2B illustrates this Markov Chain Monte Carlo (MCMC) process, by recursively sampling from $P(\mathbf{h}|\mathbf{v})$ and $P(\mathbf{v}|\mathbf{h})$, which converges at equilibrium to $P(\mathbf{v}, \mathbf{h})$.

7.0.5.3 Marginal distributions

The marginal distribution $P(\mathbf{v})$ has a closed-form expression because of the factorized conditional distribution of Equation 9 (Tubiana et al., 2019a,b):

$$P(\mathbf{v}) = \int \prod_{\mu=1}^M dh_{\mu} \cdot P(\mathbf{v}, \mathbf{h}) = \frac{1}{Z} \exp \left(\sum_{i=1}^N g_i v_i + \sum_{\mu=1}^M \Gamma_{\mu}(I_{\mu}(\mathbf{v})) \right) \quad (12)$$

For a quadratic potential $\mathcal{U}_{\mu}(h) = \frac{\gamma_{\mu} h_{\mu}^2}{2} + \theta_{\mu} h_{\mu}$, the cumulant generating function would also be quadratic and $P(\mathbf{v})$ would reduce to a Hopfield model, i.e., a pairwise model with an interaction matrix $J_{ij} = \sum_{\mu} \frac{w_{i\mu} w_{j\mu}}{\gamma_{\mu}}$ (Tubiana et al., 2019a). Otherwise, Γ_{μ} is not quadratic, yielding high-order effective interaction terms between visible units and allowing RBMs to express more complex distributions. Importantly, the number of parameters remains limited, controlled by M and does not scale as N^2 (unlike pairwise models).

7.0.5.4 Choice of HU potential

The choice of HU potential determines three related properties: the HU conditional distribution $P(\mathbf{h}|\mathbf{v})$, the transfer function of the HUs and the parametric form of the marginal distribution $P(\mathbf{v})$. Hereafter we use the double-Rectified Linear Unit (dReLU) potential:

$$\mathcal{U}_{\mu}(h) = \frac{1}{2} \gamma_{\mu,+} h_+^2 + \frac{1}{2} \gamma_{\mu,-} h_-^2 + \theta_{\mu,+} h_+ + \theta_{\mu,-} h_-, \quad \text{where } h_+ = \max(h, 0), \quad h_- = \min(h, 0) \quad (13)$$

Varying the parameters $\{\gamma_{\mu,+}, \gamma_{\mu,-}, \theta_{\mu,+}, \theta_{\mu,-}\}$ allows the potential to take a variety of shapes, including quadratic potentials ($\gamma_{\mu,+} = \gamma_{\mu,-}$, $\theta_{\mu,+} = \theta_{\mu,-}$) and double-well potentials (Tubiana et al., 2019b). The associated cumulant generating function $\Gamma(I)$ is non-quadratic in general, and depending on the parameters, the transfer function can be linear, ReLU-like (asymmetric slope and thresholding) or logistic-like (strong local slopes for binarizing inputs). Closed-form expressions of Γ are detailed in (Tubiana et al., 2019b,a), and its derivatives are also detailed in (Tubiana, 2018, p49-50). Note that the dReLU potential \mathcal{U}_{μ} and distribution $P(\mathbf{v})$ are invariant to the sign swap transformation $\gamma_{\mu,+}, \theta_{\mu,+} \iff \gamma_{\mu,-}, \theta_{\mu,-}$ and $w_{i\mu} \iff -w_{i\mu} \forall i, \mu$ (leading to $h_{\mu} \iff -h_{\mu}$). For visual clarity, we perform this sign swap transformation after training on all HUs with predominantly negative weights (defined by $\sum_i w_{i,\mu} < 0$). Subsequently all HUs are positively activated if the group of neurons to which it connects is strongly active.

7.0.5.5 RBM training

The RBM is trained by maximizing the average log-likelihood of the empirical data configurations $\mathcal{L} = \langle \log P(\mathbf{v}) \rangle_{\text{data}}$, using stochastic gradient descent methods. The gradient update steps are derived by calculating the derivative

of \mathcal{L} , using Equation 12, with respect to the model parameters (Tubiana et al., 2019a):

$$\begin{aligned}\frac{\partial \mathcal{L}}{\partial g_i} &= \langle v_i \rangle_{\text{data}} - \langle v_i \rangle_{\text{model}} \\ \frac{\partial \mathcal{L}}{\partial w_{i\mu}} &= \langle v_i h_\mu \rangle_{\text{data}} - \langle v_i h_\mu \rangle_{\text{model}} \\ \frac{\partial \mathcal{L}}{\partial \theta_{\mu,\pm}} &= -\langle h_\mu^\pm \rangle_{\text{data}} + \langle h_\mu^\pm \rangle_{\text{model}} \\ \frac{\partial \mathcal{L}}{\partial \gamma_{\mu,\pm}} &= -\frac{1}{2} \langle h_\mu^{\pm 2} \rangle_{\text{data}} + \frac{1}{2} \langle h_\mu^{\pm 2} \rangle_{\text{model}}\end{aligned}\tag{14}$$

Each gradient of \mathcal{L} is thus the difference between a data statistic $\langle f_k \rangle_{\text{data}}$ and a model statistic $\langle f_k \rangle_{\text{model}}$. Hence the model learns to match these statistics to the training data. Importantly, model statistics $\langle f_k \rangle_{\text{model}}$ cannot be evaluated exactly due to the exponentially large number of data configurations (e.g., 2^N visible configurations). Therefore they are approximated by computing the statistics of model-generated data using the MCMC sampling scheme defined with Equations 8 and 10. MCMC sampling of a Boltzmann distribution in such high-dimensional space is in general very challenging owing to the exponentially long time to reach equilibrium. We use the persistent contrastive divergence approximation (Tieleman, 2008) and discuss its validity below.

7.0.6 Compositional Restricted Boltzmann Machine

In the previous sections we have described the general properties of RBMs. We now motivate the specific RBM model choices that we have implemented, such as the dReLU potential and sparsity regularization, by discussing their impact on the properties of RBM-generated data.

Directed graphical models, e.g., PCA, ICA, sparse dictionaries or variational autoencoders, prescribe a priori statistical constraints for their data representations, such as orthogonality/independence or specific marginal distributions such as Gaussian/sparse distributions. In contrast, the statistical properties of the representation of the data learned by RBMs are unknown a priori by construction (because of the maximum entropy principle). Instead, they emerge from the structure of the weight matrix, the potentials and the recursive back-and-forth sampling procedure described above. We have therefore previously studied the properties of typical samples of RBM with random weights as a function of the visible and hidden unit potentials and properties of the weight matrix using statistical mechanics tools (Tubiana & Monasson, 2017; Tubiana et al., 2019a). We have identified the three following typical behaviors, or phases.

In the ferromagnetic phase, a typical sample from $P(\mathbf{v}, \mathbf{h})$ has a single strongly activated HU ($m(t) \sim 1$, where $m(t)$ is the number of activated HUs at time t), whereas the others are not or merely weakly activated. The corresponding active visible units v_i are defined by the weight vector \mathbf{w}_{μ^*} associated to the active HU h_{μ^*} (see Equation 10).

In the spin-glass phase, a typical sample does not have any relatively strongly activated HUs, but instead many moderately activated ones ($m(t) \sim M$). They interfere in a complex fashion to produce different visible unit configurations and there is no clear correspondence between the weight matrix and a typical data configuration.

Finally, in the compositional phase, a typical sample from $P(\mathbf{v}, \mathbf{h})$ has a small number of strongly activated HUs ($1 \ll m(t) \ll M$) whereas the others are weak or silent. Their weights are linearly combined through Equation 10 to produce the corresponding visible layer configuration. The compositional phase is desirable because, firstly, there exists a simple link between the weight matrix and typical data configurations (they are obtained by combining a few weights), which facilitates interpretation of biological systems (Tubiana et al., 2019b). Secondly, the corresponding neural activity distribution is rich, as different choices of HU subsets yield a combinatorial diversity of visible layer configurations. Moreover, the modular nature of the compositional

phase facilitates the assembly organization of neural dynamics, as motivated in the Introduction.

A set of sufficient conditions for the emergence of the compositional phase are (Tubiana & Monasson, 2017):

1. The HUs are unbounded and real-valued with a ReLU-like transfer function.
2. The weight matrix \mathbf{W} is sparse.
3. The columns \mathbf{w}_μ of the weight matrix have similar norm. (If a weight column associated to one HU is much larger than the others, visible configurations are solely aligned to it according to Equation 10.)

The first condition is satisfied by the dReLU potential (but not by quadratic potentials or binary-valued HUs). The second condition is enforced in practice by adding a L_1 sparse penalty term $\lambda \cdot \sum_{i\mu} |w_{i\mu}|$ to the log-likelihood cost function. In our experiments, the optimal sparsity parameter λ was determined to be $\lambda = 0.02$ by cross-validation (Figure S1). The final condition is achieved by enforcing that $\text{Var}(h_\mu) = 1$ and $\langle h_\mu \rangle \sim 0 \forall \mu$. This is done by an appropriate reparameterization of the HU potential of Equation 13 and a batch-norm-like procedure, described in detail in (Tubiana, 2018). This normalization promotes homogeneity among HU importance, preventing some units from being disconnected or others from dominating. In addition, ensuring that $h_\mu = O(1)$ irrespective of the visible layer size (as opposed to e.g., $\frac{1}{2}(\gamma_+ + \gamma_-) = 1$ which yields $h_\mu \sim \sqrt{N}$) avoids the problem of ill-conditioned Hessians that was previously described by Hinton (2012).

To emphasise the departure from the classic RBM formulation in this study, we name our model compositional RBM (cRBM).

7.0.7 Algorithmic Implementation

In the previous sections we have described the cRBM model in full mathematical detail. The corresponding algorithmic implementation was adapted from (Tubiana et al., 2019b). In addition, we have made several major implementation and algorithmic changes to accommodate the large data size of the zebrafish neural recordings. We provide the code open-source, and describe the code improvements and hyperparameter settings in this section. The following improvements were made, leading to a substantial reduction of computation time:

- Python 3 and numba (Lam et al., 2015) were used to compile custom functions, enabling SIMD vectorization and multicore parallelism.
- The sampling of $P(h_\mu | I_\mu)$ and evaluating its cumulant generating function Γ_μ and various moments requires repeated and costly evaluation of error functions erf and related functions (Tubiana, 2018, p49-50). Fast numerical approximations of these functions were implemented based on (Abramowitz et al., 1988, p299).
- The number of memory allocation operations was minimized.
- The optimization algorithm was changed from stochastic gradient ascent to RMSprop (i.e., ADAM without momentum) with learning rate $5 \cdot 10^{-4}$ to $5 \cdot 10^{-3}$, $\beta_1 = 0$, $\beta_2 = 0.999$, $\epsilon = 10^{-6}$, see Kingma & Ba (2014) for a definition of the parameters. Compared to the original stochastic gradient ascent, the adaptive learning rates of RMSprop/ADAM yield larger updates for the weights attached to neurons with very sparse activity, resulting in substantially faster convergence.

7.0.7.1 Hyperparameter settings

The following hyperparameters were used in the experiments of this manuscript:

- Number of hidden unit M : 200. This value was determined by cross-validation (Figure S1) on one data set (example fish #3). Because this cross-validation procedure was computationally expensive, the same value was used for all other data sets, except for 3 data sets which used $M = 100$ because their $N \approx \frac{1}{2}N_{\#3}$.
- Sparse regularization penalty λ : 0.02 (determined by cross-validation).

- Batch size: 100, 200 or 400. Larger batch sizes yield longer training time but more stable training; batch size was increased if training failed to converge.
- Number of Monte Carlo chains: 100.
- Number of gradient updates: $2 \cdot 10^5$
- Number of Monte Carlo steps between each gradient update: 15.
- Initial learning rate η : between $5 \cdot 10^{-4}$ and $5 \cdot 10^{-3}$. We used $5 \cdot 10^{-3}$ by default and if weight divergence was observed, the learning was reinitialized with a reduced learning rate. This occurred notably for high- M and low- λ models during the cross-validation procedure of Figure S1.
- Learning rate annealing scheme: the learning rate geometrically decayed during training, starting after 25% of the gradient update steps, from its initial value η to a final value of 10^{-5} .
- Number of training data samples: 70% of frames of each recording (= 4086 training data samples on average), see section 7.0.10.1 for details.

cRBM models of these zebrafish data sets could be estimated in approximately 8-12 hours using 16 CPU threads (Intel Xeon Phi processor). The (λ, M) -cross-validation was therefore completed in three weeks using two desktop computers.

7.0.8 Validity of the Persistent Contrastive Divergence algorithm

Training RBM requires extensive MCMC sampling which is notoriously difficult for high-dimensional data sets. We resolve this by using Persistent Contrastive Divergence (PCD) to approximate the gradients (Tieleman, 2008). In this section we discuss why this worked to successfully converge, despite the very large data size.

The typical number of Monte Carlo steps required to transition from one energy minimum to another through an energy barrier ΔE follows the Arrhenius law, scaling as $e^{\Delta E}$. In the thermodynamic limit ($N \rightarrow \infty$), ΔE scales as the system size N multiplied by the typical energy required to flip a single visible unit, corresponding here to the inputs received from the hidden layer I . In contrast, for PCD only a limited number of MC steps (here, 15) are applied between each gradient update. Three factors explain why reasonably successful convergence was achieved in the trainings presented here.

Firstly, the use of the L_1 regularization limits the magnitude of the weights and therefore limits the input scale I . Secondly, in the compositional phase, the energy barriers do not scale as the full system size N but rather as the size of one assembly pN where p is the fraction of non-zero weights (Tubiana & Monasson, 2017). Indeed, transitioning from one energy minimum, characterized by a subset of strongly activated HUs, to another minimum, characterized by another set of strongly activated HUs, is done by gradually morphing the first set into the second (Roussel et al., 2021). Compared to a direct transition, such a path is favored because the intermediate states are thermodynamically stable and energy barriers are smaller as each HU flip has an energy cost $\sim pN$. Lastly, throughout PCD training, MCMC sampling is not performed at thermal equilibrium and the model updates of the parameters of the distribution promote mixing (Tieleman & Hinton, 2009). This is seen from Equation 14: the log-likelihood gradient is the difference between the gradient of the energy averaged over the empirical data and the energy averaged over MCMC samples. Ascending the gradient amounts to pushing down the energy of data configurations and pushing up the energy of MCMC samples, thereby promoting mixing of the Markov chains.

Overall, combining small learning rates (and large number of gradient updates), large regularization, large number of Markov Chains and Monte Carlo steps has allowed convergence to be reached for the majority of cRBM training sessions.

7.0.9 Functional Connectivity Inference

7.0.9.1 Effective connectivity matrix

In this section we present a derivation of the effective coupling matrix between neurons from the marginal distribution $P(\mathbf{v})$ using cRBMs. This is achieved by perturbing the activity of each neuron individually and quantifying the effect on other neurons. We first define the local coupling J_{ij} between two neurons v_i and v_j for a generic probability distribution $P(v_1, v_2, \dots, v_N)$, given a data configuration \mathbf{v} :

$$J_{ij}(\mathbf{v}) = \log \left(\frac{P(v_i = 1 | v_1, \dots, v_{i-1}, v_{i+1}, v_j = 1, \dots, v_N)}{P(v_i = 1 | v_1, \dots, v_{i-1}, v_{i+1}, v_j = 0, \dots, v_N)} \right) - \log \left(\frac{P(v_i = 0 | v_1, \dots, v_{i-1}, v_{i+1}, v_j = 1, \dots, v_N)}{P(v_i = 0 | v_1, \dots, v_{i-1}, v_{i+1}, v_j = 0, \dots, v_N)} \right) \quad (15)$$

In other words, J_{ij} is defined as the impact of the state of neuron j on neuron i in the context of activity pattern \mathbf{v} . Hence, the effective connectivity matrix \mathbf{J} mathematically defines the functional connections, which can only be done using a probabilistic model $P(\mathbf{v})$. A positive (negative) coupling J_{ij} indicates correlated (anti-correlated) collective behavior of neurons i, j . This effective coupling value is symmetric (because of Bayes' rule): $J_{ij}(\mathbf{v}) = J_{ji}(\mathbf{v})$. For context, note that $J_{ij}(\mathbf{v})$ is uniformly zero for an independent model of the form $P(v_1, \dots, v_N) = \prod_i P_i(v_i)$, and that for a maximum entropy pairwise (Ising) model, with $P(v_1, \dots, v_N) = \frac{1}{Z} \exp \left(\sum_i g_i v_i + \sum_{i < j} J_{ij}^{\text{ising}} v_i v_j \right)$, the $J_{ij}(\mathbf{v})$ matrix exactly identifies with the coupling matrix J_{ij}^{ising} , and does not depend on the data configuration \mathbf{v} (so $J_{ij}(\mathbf{v}) = J_{ij}$).

However, in general, and for RBMs in particular, $J_{ij}(\mathbf{v})$ depends on the data set \mathbf{v} , and an overall coupling matrix can be derived by taking its average over all data configurations:

$$J_{ij} = \langle J_{ij}(\mathbf{v}) \rangle_{\text{data}} \quad (16)$$

Although Equation 16 has a closed-form solution for RBMs (by inserting Equation 12), a naive evaluation requires $O(N^3 MT)$ operations where T is the number of data samples. However, a fast and intuitive approximation can be derived by performing a second order Taylor expansion of $\Gamma_\mu(I_\mu)$:

$$J_{ij} = \sum_{\mu=1}^M w_{i\mu} w_{j\mu} \langle \Gamma''_\mu(\mathbf{v}) \rangle_{\text{data}} = \sum_{\mu=1}^M w_{i\mu} w_{j\mu} \langle \text{Var}(h_\mu | \mathbf{v}) \rangle_{\text{data}} \quad (17)$$

Equation 17 is exact for quadratic potential and in general justified as the contribution of neurons i, j is small compared to the scale of variation of Γ_μ , $O(\sqrt{pN})$ where p is the fraction of non-zero couplings. In conclusion, we have mathematically derived the effective coupling between any two neurons i and j . Intuitively, two neurons i, j are effectively connected if they are connected to the same HUs (Equation 17).

7.0.9.2 From inter-neuron to inter-region connectivity

In the above section we have derived the inter-neuronal connectivity matrix \mathbf{J} . This matrix is then aggregated to an inter-regional connectivity matrix \mathbf{J}^R by taking the L_1 norm of the corresponding \mathbf{J} matrix block (i.e., $J_{km}^R = \sum_{i \in R_k, j \in R_m} |J_{ij}| / (N_{R_k} \cdot N_{R_m})$, where R_k is the set of neurons in region k).

Next, to derive the average connectivity matrix across multiple recordings, we used a weighted average of the individual recordings, with a region-pair specific weight equal to the length of the recording multiplied by the sum of the number of neurons in both regions (also see Section 7.0.16). Compared to a naive average, this weighted average accounts for the variable number of neurons per region between recordings.

7.0.9.3 Training cRBM models for connectivity estimates

Constructing the functional connectivity matrix of a cRBM does not require test data, but just the estimated weight matrix \mathbf{W} (as explained above). Therefore we trained new cRBMs using the entire recordings (100% of

data) to fully use the information available. cRBM training is stochastic, and to mitigate the possible variability that could arise we trained 5 cRBMs for each recording. Then, to assess convergence, we selected all cRBMs with $0.01 < \text{std}(\mathbf{w}) < 0.1$, where std denotes standard deviation, for further functional connectivity analysis (yielding 23 cRBMs for 8 data sets in total). Connectivity estimates of multiple cRBM models per data sets were averaged.

7.0.9.4 Connectivity inference baselines

We considered four additional connectivity inference baseline methods:

- The covariance matrix.
- The Pearson correlation matrix.
- The sparse inverse covariance matrix inferred by graphical LASSO (Friedman et al., 2008) (as implemented in scikit-learn with default settings (Pedregosa et al., 2011)). Graphical LASSO is an efficient method for inference of large scale connectivity. Unfortunately, the implementation available failed to converge in reasonable time due to the high dimensionality of the data.
- The Ising model with pseudo-likelihood maximization (PLM) inference (Ravikumar et al., 2010).

Results obtained with the covariance and correlation matrices are presented in Figure S6. The connectivity matrices obtained by the PLM Ising model (not shown) correctly identified the diagonal entries of the region-region matrix, but not the off-diagonal coefficients and had a weaker correlation with the structural connectivity matrix than the covariance and correlation matrices ($r_S = 0.06$ using 4 fish).

7.0.10 Optimizing the free parameters of cRBM

We set the free parameters λ (sparsity regularization parameter) and M (number of HUs) by cross-validating a large range of (λ, M) values for one data set (fish #3). This was done by training cRBMs on 70% of the data, and evaluating model performance on the remaining test data, as detailed below. The resulting optimal values could then be used for all data sets (where M was scaled with the number of neurons N). Importantly, the (λ, M) parameters implicitly tune the average assembly size. Increasing the number of HUs and/or increasing the regularization strength decreases the average number of neurons per assembly (Tubiana et al., 2019a). Intuitively, assemblies that are too small do not have the capacity to capture high-order correlations, while assemblies that are too large would fail to account for local co-activations. Hence, the (M, λ) -cross-validation effectively identifies the optimal assembly sizes that fit the data statistics.

7.0.10.1 Train / test data split

We split up one recording (fish #3) into training data (70% of recording) and withheld test data (30% of recording) for the free parameter (λ, M) optimization procedure. This enabled us to assess whether the cRBMs learned to model the data statistics (as described in the main text, Figures 2 and S1), while ensuring that the cRBMs had not overfit to the specific training data configurations. Importantly, this assumes that the test data comes from the same statistical distribution as the training data (while consisting of different data configurations). To ensure this, we split up the recording of example fish #3 (used for parameter optimization) in training and test splits as follows (before training the cRBMs): We divided the recording of length T in 10 chronological segments of equal length (so that segment 1 has time points $\{t \in [1, \frac{T}{10}]\}$ et cetera), with the rationale that by maintaining temporal order within each segment we would later be able to conduct dynamic activity analysis. This yielded $\binom{10}{3} = 120$ possible training/test splits of the neural data. We then evaluated the statistical similarity between the training and test split of each combination, by assessing the mean neural activity $\langle v_i \rangle$ and pairwise neural correlations $\langle v_i v_j \rangle - \langle v_i \rangle \langle v_j \rangle$ statistics. We quantified the similarity between training and test statistics by calculating the Root Mean Square Error ($\text{RMSE}(\mathbf{x}_1, \mathbf{x}_2) = \sqrt{\frac{1}{N_x} \sum_{n=1}^{N_x} (x_1(n) - x_2(n))^2}$). The most similar split is defined by the lowest RMSE, but to show that cRBM are not dependent on picking the best possible split, but rather on

avoiding the bad splits, we then chose to use the split with the 10th-percentile ranking RMSE. We hope that this aids future studies, where a potentially high number of possible splits prevents researchers from evaluating all possible splits, but a good split may nevertheless be found efficiently.

7.0.10.2 Assessment of data statistics

Please note that the loss function, the log-likelihood, is computationally intractable and therefore cannot be readily used to monitor convergence or goodness-of-fit after training (Fischer & Igel, 2012). Moreover, approximations of the log-likelihood based on annealed importance sampling were found to be unreliable due to the large system size. Instead, we assessed the following quantities.

Firstly, we evaluated 3 statistics that cRBMs are trained to optimize: the mean activity of neurons $\langle v_i \rangle$, the mean activity of HUs $\langle h_\mu \rangle$ and their pairwise interactions $\langle v_i h_\mu \rangle$. Additionally, second order statistics of pairwise neuron-neuron interactions $\langle v_i v_j \rangle$, HU-HU interactions $\langle h_\mu h_\nu \rangle$ and the reconstruction quality were evaluated, which the cRBM was not constrained to fit. Data statistics were calculated on withheld test data (30% of recording). Monitoring HU single and pairwise statistics $\langle h_\mu \rangle$ and $\langle h_\mu h_\nu \rangle$ served two purposes: (i) validation of model convergence and (ii) assessing whether correlations between assemblies can be captured by this bipartite model (i.e., without direct couplings between hidden units or an additional hidden layer). Model statistics cannot be calculated exactly, because that would require one to sample all possible states $P(\mathbf{v}, \mathbf{h})$, and were therefore approximated by evaluating cRBM-generated data. Here, 300 Monte Carlo chains were each initiated on random training data configurations and 50 configurations were sampled consecutively for each chain, with 20 sampling steps between saved configurations, after a burn-in period of 100 effective sampling configurations.

The $\langle v_i h_\mu \rangle$ statistic (Figure 2C) was corrected for the sparsity regularization, by adding the sparsity regularization parameter λ to $\langle v_i h_\mu \rangle$: $\langle v_i h_\mu \rangle_{\text{model}} = \langle v_i h_\mu \rangle_{\text{model-generated data}} + \lambda \cdot \text{sign}(w_{i,\mu})$. Furthermore, (v_i, h_μ) pairs with exactly $w_{i,\mu} = 0$ were excluded from analysis (5% of total for optimal cRBM in Figure 2C).

The pairwise neuron-neuron and HU-HU statistics ($\langle v_i v_j \rangle$, $\langle h_\mu h_\nu \rangle$) were corrected for their (trivially) expected correlation due to their mean activities (by subtraction of $\langle v_i \rangle \langle v_j \rangle$ and $\langle h_\mu \rangle \langle h_\nu \rangle$ respectively), so that only true correlations were assessed.

Goodness of fit was quantified by computing the normalized Root Mean Square Error (nRMSE) for each statistic (shown in Figure S1). The RMSE between two vectors $\mathbf{x}_1, \mathbf{x}_2$ of length N_x is defined as $\text{RMSE} = \sqrt{\frac{1}{N_x} \sum_{n=1}^{N_x} (x_1(n) - x_2(n))^2}$. Ordinary RMSE was normalized so that different statistics could be compared, where 1 corresponds to $\text{nRMSE}_{\text{shuffled}}$, where both data and model statistics were randomly shuffled, and 0 corresponds to $\text{nRMSE}_{\text{optimal}}$ which is the RMSE between the training data and test data (by $\text{nRMSE} = 1 - \frac{\text{RMSE}_{\text{ordinary}} - \text{RMSE}_{\text{shuffled}}}{\text{RMSE}_{\text{optimal}} - \text{RMSE}_{\text{shuffled}}}$).

Additionally, we assessed the reconstruction quality of the test data. Here, the log-likelihood (LLH) between the test data \mathbf{v} and its reconstruction $\mathbb{E}(\mathbf{v}_{\text{recon}}) = \mathbb{E}(\mathbf{v} | \mathbb{E}(\mathbf{h} | \mathbf{v})) \in [0, 1]$ was computed. Because $v_i \in \{0, 1\}$, the LLH is defined as

$$\text{LLH}(v_i, \mathbb{E}(\mathbf{v}_{\text{recon}})) = \frac{1}{T} \sum_{t=1}^T \log \left(\mathbb{E}(\mathbf{v}_{\text{recon}})(t) * v_i(t) + (1 - \mathbb{E}(\mathbf{v}_{\text{recon}})(t)) * (1 - v_i(t)) \right) \quad (18)$$

The resulting LLH was normalized (nLLH) such that 0 corresponds to an independent model (i.e., fitting neural activity with $\mathbb{E}(\mathbf{v}_{\text{recon}})(t) = \langle v_i \rangle \forall t$) and 1 to optimal performance (which is $\text{LLH}_{\text{optimal}} = 0$), by $\text{nLLH} = \frac{\text{LLH}_{\text{ordinary}} - \text{LLH}_{\text{independent}}}{-\text{LLH}_{\text{independent}}}$.

7.0.10.3 Generalized Linear Model

We used logistic regression, a Generalized Linear Model (GLM), to quantify the reconstruction quality of a fully connected model (i.e., with neuron-to-neuron connections, see Figure S2A). Logistic regression makes a probabilistic binary prediction (Bishop, 2006), hence allowing direct comparison to the probabilistic estimates of neural activity by the cRBM. In logistic regression, for a neuron $v_i(t)$ at time t , the activity of all other neurons $\mathbf{v}_{-i}(t)$ at time t was used to predict $\hat{v}_i(t) = P(v_i(t) = 1) = \frac{1}{1 + \exp(-\mathbf{w}_i \cdot \mathbf{v}_{-i}(t))}$ where \mathbf{w}_i is the estimated weight vector. This was implemented with scikit-learn (Pedregosa et al., 2011), using L_2 regularization. L_2 regularization was favored over L_1 as it typically yields higher reconstruction performance; in the related context of protein contact map prediction, L_2 -regularized GLMs also better reconstructed contacts than L_1 -regularized GLMs (Morcos et al., 2011). The parameter λ_{GLM} was optimized to $\lambda_{\text{GLM}} = 1000$ using cross-validation (Figure S2B). This is a computationally intensive model to compute because of the large number of regressor neurons $N - 1$. Therefore we performed the cross-validation of Figure S2B on 2% of all neurons (=1050 neurons) and computed the final distribution of Figure 2H on 10% of all neurons (=5252 neurons). GLMs were trained on the same train data as cRBMs, and evaluated on the same withheld test data as cRBMs (as described above).

7.0.11 Regional occupancy

We determined the anatomical region labels of each neuron by registering our recordings to the ZBrain Atlas (as described previously). This yields a matrix L of size $N_{\text{ZBA}} \times N$, which elements are $l_{r,i} = 1$ if neuron i is embedded in region r and 0 if it is not. A cRBM neural assembly of HU μ is defined by its weight vector \mathbf{w}_μ (of size N). Because cRBMs converge to sparse solutions, most of the weight elements will be very close to 0. To determine which anatomical regions are occupied by the assembly neurons with significantly nonzero weights, we computed the dot product between the weight vector \mathbf{w}_μ and matrix L , leading to a weighted region label vector (of size N_{ZBA}) for each HU. The matrix of all M weighted region label vectors is shown in Figure S4A for cRBM and Figure S4B for PCA.

The effective number of anatomical regions that one cRBM/PCA assembly is embedded in was then calculated using the Participation Ratio (PR) of each HU/Principal Axis. PRs are used to estimate the effective number of nonzero elements in a vector, without using a threshold (Tubiana & Monasson, 2017). The PR of a vector $\mathbf{x} = (x_1, \dots, x_n)$ is defined by:

$$\text{PR}(\mathbf{x}) = \frac{\left(\sum_{i=1}^n x_i^2\right)^2}{\sum_{i=1}^n x_i^4} \quad (19)$$

PR varies from $\frac{1}{n}$ when only 1 element of \mathbf{x} is nonzero and n when all elements are equal. We therefore estimated the effective number of regions by multiplying PR of the weighted region label vectors with the total number of regions N_{ZBA} in Figure S4C.

7.0.12 Time constant calculation

The dReLU potential \mathcal{U}_μ of equation 13 can learn to take a variety of shapes, including a double-well potential (Tubiana et al., 2019a). HUs generally converged to this shape, giving rise to bimodal HU activity distributions (Figure 4D). We determined the positions of the two peaks per HU using Gaussian Mixture Models fitted with two Gaussians. The bimodality transition point was then defined as the average between the two peaks (which was approximately 0 for most HUs). To calculate the time constant of state changes between the two activity modes, we subtracted the bimodality transition point from each HU activity h_μ individually. For clarity, all dynamic activity traces shown (e.g., Figure 4) are thus bimodality transition point subtracted. The time constant of an activity trace was then defined as the period of a (two-state) oscillation. A HU oscillation is defined as a consecutive negative and positive activity interval (because the bimodality now occurs at 0). A neuron oscillation is defined as a consecutive interspike-interval and spike-interval (which can last for multiple time steps, for example see Figure 1A, right panel).

7.0.13 Sorting of HUs

HUs were sorted by hierarchical clustering of the Pearson correlation matrix of their dynamic activity (Figure 4B). Hierarchical clustering was performed using the Ward variance minimization algorithm that defines

the distance between clusters (Virtanen et al., 2020). This sorting of HUs (and thus assemblies) is maintained throughout the manuscript for the sake of consistency.

7.0.14 Validating that the cRBM is in the compositional phase

To validate that the cRBMs converged to the compositional phase (see section 7.0.6, compositional RBM formulation), we calculated the effective number of active HUs per data configuration (i.e., time step) $m(t) = \text{PR}(\mathbf{h}_+(t)) \cdot M$ where PR is the participation ratio (equation 19), M the number of HUs and $h_+ = h - h_{\text{inactive}}$, where h_{inactive} is the inactive peak as calculated with the Gaussian Mixture Models (see section 7.0.12), because PR assumes that inactive elements are approximately zero (Tubiana et al., 2019a). A cRBM is said to be in the compositional phase if $1 \ll \text{median}(m) \ll M$, which is true for all cRBMs (Figure S3).

7.0.15 Extensions of the structural connectivity matrix

The inter-region structural connectivity matrix was derived from the single cell zebrafish brain atlas (Kunst et al., 2019). We used the post-publication updated data set from (Kunst et al., 2019) (timestamp: 28 October 2019). The data set consists of $N = 3098$ neurons, each characterized by the 3D coordinates of the soma center and of its neurites; there is no distinction between dendrites and axons. The brain is subdivided into $R = 72$ regions and each neuron is duplicated by left/right hemisphere symmetry. We aim to estimate $c_{r,r'}$, the average strength of the connection between two neurons belonging to regions $r, r' \in [1, R]$. For each neuron $n \in [1, N]$, we determine, using region masks, the region $r(n)$ where its soma is located and the cumulative length of the intersection between all its neurites and each region $\ell_n(r)$. Under the assumptions that (i) the linear density of dendritic spines / axon presynaptic boutons is constant and (ii) the volumetric density of neurons is identical throughout regions, $L_n(r)$ is proportional to the volume V_r of region r times the average (bidirectional) connection strength between neuron n and any neuron of region r . Aggregating over all neurons and symmetrizing, we obtain the following estimator for $c_{r,r'}$:

$$c_{r,r'} = \text{Symmetrized} \left\{ \frac{\sum_{n=1}^N \delta_{r(n),r} \times \ell_n(r')}{V_{r'} \times \sum_{n=1}^N \delta_{r(n),r}} \right\} \quad (20)$$

where $\delta_{r(n),r} = 1$ if neuron n has its soma in region r and 0 if not. Using the same notations, the formula previously used in (Kunst et al., 2019) is:

$$c_{r,r'} = \left\{ \frac{\sum_{n=1}^N \ell_n(r) + \ell_n(r')}{N(V_{r'} + V_r)} \right\} \quad (21)$$

Equation 20 differs from Equation 21 in three aspects:

1. It discriminates between direct and indirect connections. Previously, a structural connection between region r and region r' was established if a neuron had neurites with either tips or its soma within both regions. This may however result in indirect connections between r and r' , in cases where the neuron soma resides in another region r'' . Here, we only account for direct connections, resulting in an overall slightly sparser connectivity matrix.
2. It is well-defined along the diagonal, i.e., for intra-region connections, whereas in Equation 21, each neurite would be counted as a self-connection.
3. The denominator corrects for non-uniform sampling of the *traced* neurons throughout regions. Note that this issue only arose in the post-publication data set as non-uniform sampling was used to fill missing entries of the matrix.

7.0.16 Specimen averaging of connectivity matrices

The number of neurons in a particular brain region can vary across recordings from different specimen. Since the entries of the connectivity matrix are expected to be more accurate for well-sampled regions, we

computed the weighted average of region-to-region connections $c_{r,r'}$ as follows:

$$\begin{aligned} \langle c_{r,r'} \rangle &= \frac{\sum_{\text{Fish } F} c_{r,r'}^F \cdot w_{r,r'}^F}{\sum_{\text{Fish } F} w_{r,r'}^F} \\ w_{r,r'}^F &= T^F \frac{(N_{R_r}^F + N_{R_{r'}}^F)}{2} \end{aligned} \quad (22)$$

Where T^F is the recording length and $N_{R_r}^F$ is the number of neurons in region r of fish F that were recorded.

820 **7.0.17 Correlation analysis of connectivity matrices**

Pearson correlation was used to assess the similarity between cRBM functional connectivity matrices of different individual animals (Figure 5). Spearman correlation was used to compare structural connectivity versus functional connectivity (Figure 6), because these two metrics do not necessarily scale linearly. All correlation analyses, and the Kilmogorov-Smirnov test of Figure S6C, performed on symmetric matrices excluded one
825 off-diagonal triangle (of symmetrical values) to avoid duplicates.

8 Supplementary Information

Abbreviation	Full name	Abbreviation	Full name
th	Thalamus left	s	Subpallium left
ts	Torus semicircularis left	mos2	MO stripe 2 left
t	Tegmentum left	MOS2	MO stripe 2 right
mos1	MO stripe 1 left	vr	Vagal region left
TH	Thalamus right	OE	Olfactory epithelium right
TS	Torus semicircularis right	oe	Olfactory epithelium left
T	Tegmentum right	VR	Vagal region right
MOS1	MO stripe 1 right	IO	Inferior olive right
MOS5	MO stripe 5 right	nx	Vagus motor neurons left
mos4	MO stripe 4 left	NX	Vagus motor neurons right
MOS4	MO stripe 4 right	io	Inferior olive left
mos5	MO stripe 5 left	tl	Torus longitudinalis left
mon	Medial octavolateral nucleus left	TL	Torus longitudinalis right
MON	Medial octavolateral nucleus right	TeO	Tectum right
Ce	Cerebellum right	teo	Tectum left
ce	Cerebellum left	OB	Olfactory bulb right
MOS3	MO stripe 3 right	ob	Olfactory bulb left
mos3	MO stripe 3 left	ha	Habenula left
Pr	Preteectum right	Ha	Habenula right
pr	Preteectum left	irf	Intermediate reticular formation left
PT	Posterior tuberculum right	prf	Posterior reticular formation left
pt	Posterior tuberculum left	r	Raphe nucleus left
P	Pallium right	R	Raphe nucleus right
p	Pallium left	iRF	Intermediate reticular formation right
S	Subpallium right	pRF	Posterior reticular formation right

Table S1: Table of abbreviations of mapzebrain atlas region names (used for interregional connectivity analyses).

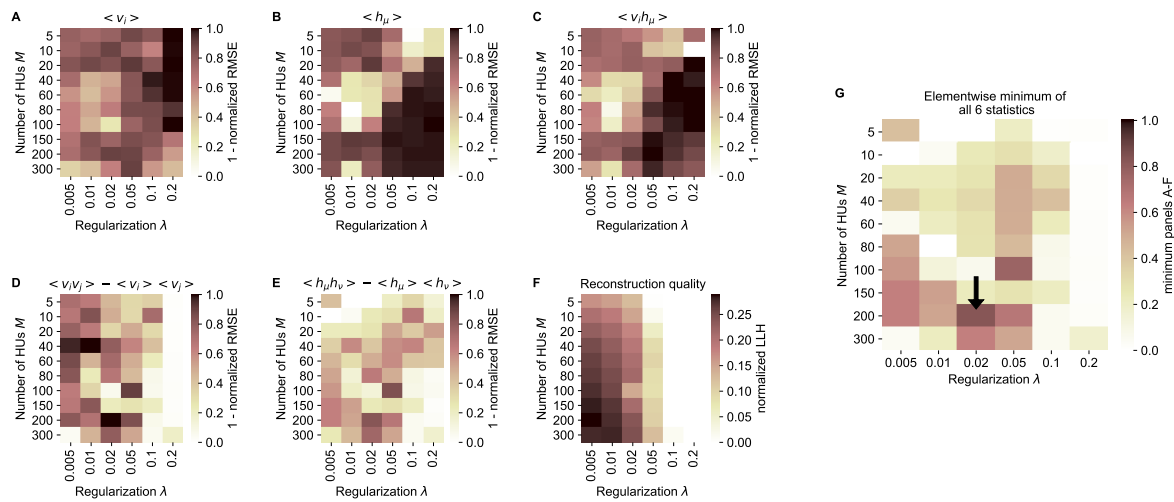


Figure S1: cRBM free parameter optimization by cross-validation

(A) Model performance, as quantified by the normalized RMSE (nRMSE, see Methods) of the mean neural activity $\langle v_i \rangle$, as a function of the sparsity regularization parameter λ (x-axis) and the number of HUs M (y-axis). cRBM models were evaluated after training converged on withheld testing data. $1 - \text{nRMSE}$ values are shown to enable optimization by maximization in panel G. Values below 0 were set to 0 for panels A-F.
 (B) Equivalent figure for the mean HU activity $\langle h_\mu \rangle$.
 (C) Equivalent figure for average pairwise neuron-HU interactions $\langle v_i h_\mu \rangle$.
 (D) Equivalent figure for average pairwise neuron-neuron interactions $\langle v_i v_j \rangle$.
 (E) Equivalent figure for average pairwise HU-HU interactions $\langle h_\mu h_\nu \rangle$.
 (F) Equivalent figure for the reconstruction quality of the low-dimensional cRBM bottleneck, quantified by the median normalized LLH.
 (G) The cRBM free parameters were optimized by maximizing the element-wise minimum of panels A-F. Using the element-wise minimum to compare the six statistics ensures that the model performs well on all aspects. First, in order to compare panels A-E with panel F, the values of panel F were scaled to 1 by dividing all elements of panel F by its maximum value ($= 0.29$). Next, for each (λ, M) combination the minimum value was determined from all 6 evaluation criteria. These are shown in this panel, with the black arrow indicating the resulting maximum of 0.83 at $(\lambda = 0.02, M = 200)$.

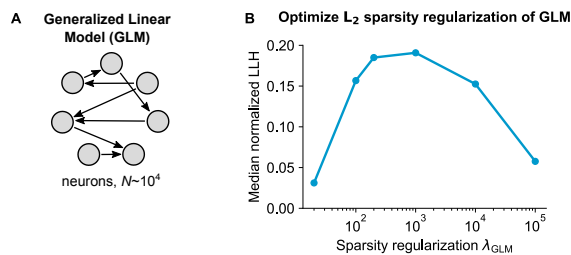


Figure S2: Generalized Linear Model (GLM) parameter optimization

(A) Schematic of the pairwise GLM model. Connections were estimated with logistic regression and L_2 sparsity regularization. (B) The sparsity regularization parameter λ_{GLM} was optimized using cross-validation of 6 parameter configurations, using 1000 randomly sampled neurons.

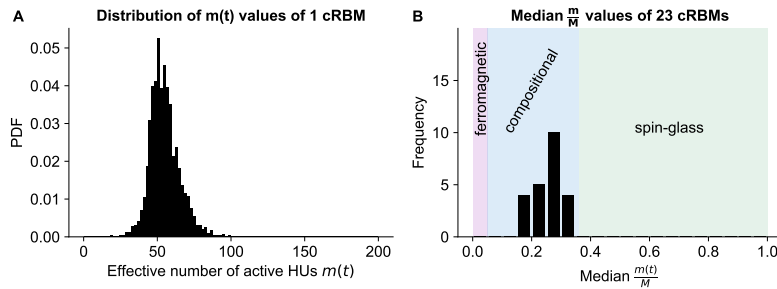


Figure S3: cRBMs are in the compositional phase after convergence

(A) Distribution of $m(t)$, the number of effective active HUs per time point t , calculated on the test data of example fish #3. $m(t)$ is defined as $m(t) = \text{PR}(\mathbf{h}_+(t)) \cdot M$, where PR is the Participation Ratio (Equation 19). Median $m(t) = 0.27 \cdot M = 54$.

(B) The distribution of median $\frac{m(t)}{M}$ values of the entire recordings of all cRBMs used for the connectivity analyses. The average $\frac{\text{median}(m)}{M}$ across all cRBMs is 0.26. The compositional phase is characterized by $1 \ll m(t) \ll M$ which occurs for all cRBMs. The three phases are indicated. Here, the ferromagnetic phase upper bound was manually set at 5%, and the spin-glass phase lower bound was determined by computing the PR of normally distributed activity (with mean and standard deviation of the test data of example fish #3 of panel A).

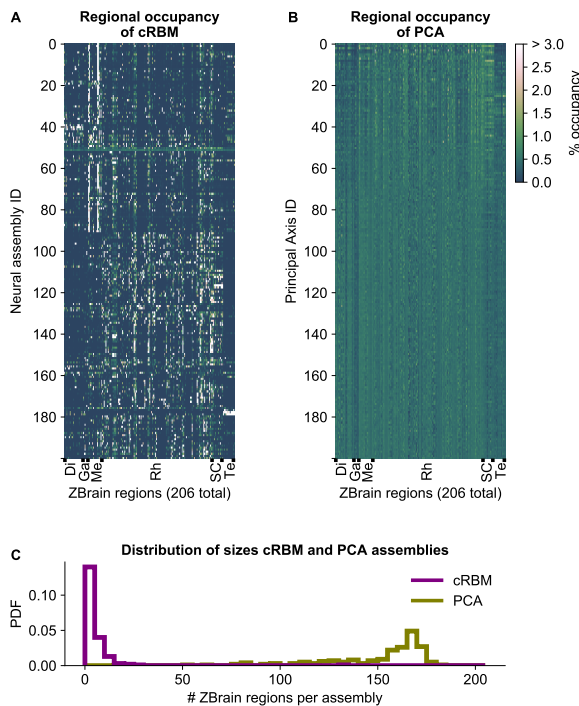


Figure S4: cRBM assemblies are sparse and spatially localized

(A) For each neural assembly (y-axis), the occupancy of ZBrain Atlas (Randlett et al., 2015) brain regions (x-axis, sorted alphabetically) was determined, by quantifying the number of anatomical regions that each HU connects to (i.e., the overlap between cRBM assemblies and anatomical regions, see Methods). ZBrain atlas regions that were not imaged were excluded, yielding $N_{\text{ZBA}} = 206$ regions in total. Occupancy was defined as the dot product between the binary ZBrain-label matrix (size $N_{\text{ZBA}} \times N$, where 1 indicates that a neuron is embedded in an anatomical region, and 0 vice versa) and the cRBM weight matrix (size $N \times M$), and was normalized to 100% for each assembly to account for different assembly sizes. Assemblies were sorted by their dynamics (Figure 4B, Methods). ZBrain Atlas regions are abbreviated by: Diencephalon (Di), Ganglia (Ga), Mesencephalon (Me), Rhombencephalon (Rh), Spinal Cord (SC) and Telencephalon (Te).

(B) Equivalent figure for Principal Axes of PCA (i.e., the PCA Eigenvectors). Principal Axes were sorted by Eigenvalue. Panels A and B share the same color scale (right).

(C) Distributions of the effective number of ZBrain Atlas regions per assembly (Principal Axis) of the occupancy metric of panel A (B) for cRBM (PCA) respectively. The effective number of regions was determined by calculating the Participation Ratio per assembly (Principal Axis), multiplied with the total number of regions N_{ZBA} (see Methods). cRBM assemblies occupy a median of 3 regions (interquartile range: 2 to 6 regions), while PCA assemblies occupy a median of 160 regions (interquartile range: 140 to 168 regions).

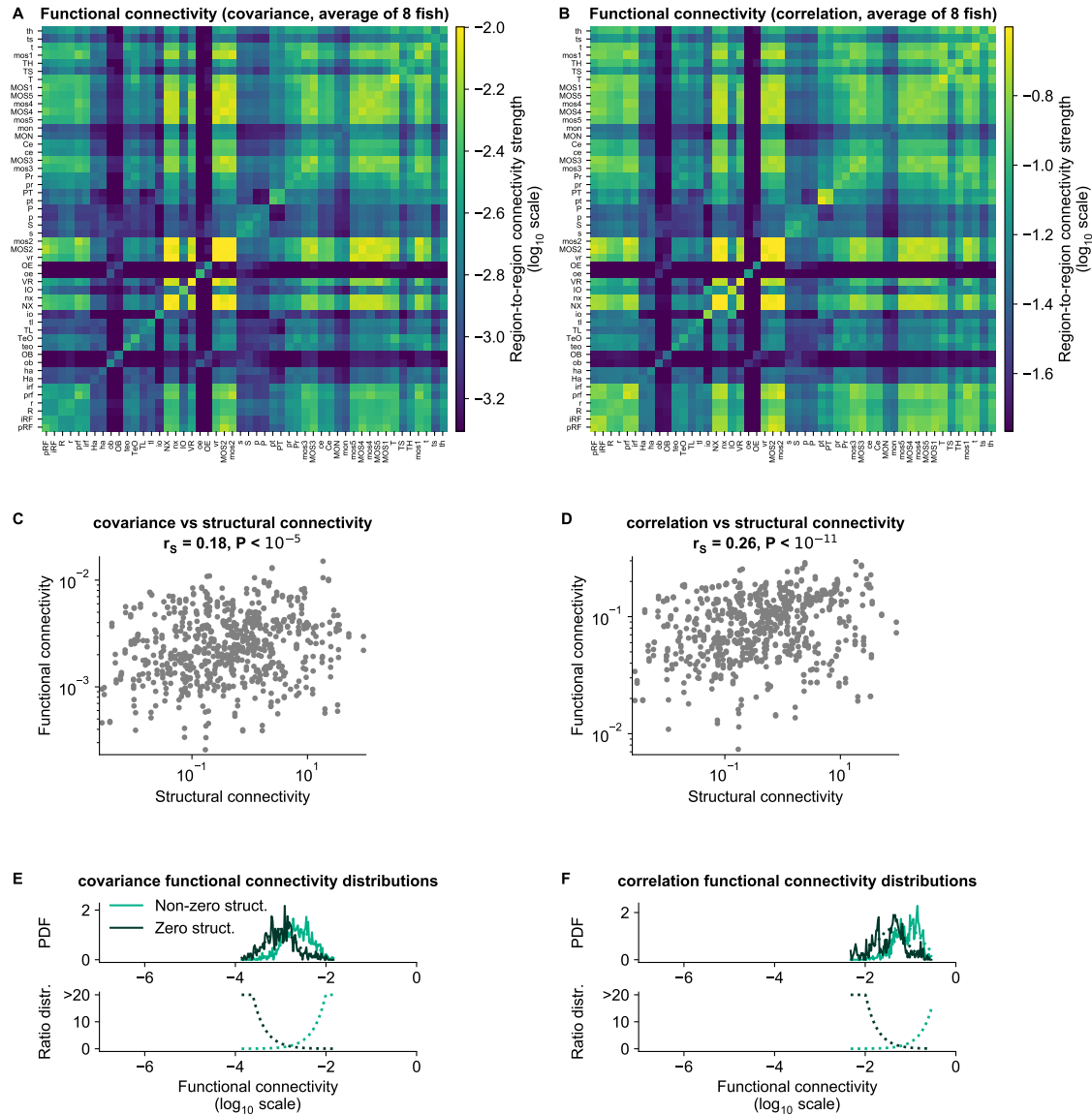


Figure S6: cRBM functional connectivity compared to baseline methods

(A) Average functional connectivity matrix of the covariance baseline method (see Methods) in \log_{10} scale.
 (B) Average functional connectivity matrix of the Pearson correlations baseline method (see Methods) in \log_{10} scale.
 (C, D) Equivalent figure to Figure 6C for functional connections as determined by the covariance (C) and correlations (D) between neurons (see Methods). The Spearman correlation coefficient between functional and structural connections for covariance is $r_s = 0.18$ and for correlation $r_s = 0.26$.
 (E-F) Equivalent figure to Figure 6D for functional connections as determined by the covariance (E) and correlations (F) between neurons. Non-zero and zero distributions are significantly different (all P values $P < 10^{-15}$, two-sided Kolmogorov-Smirnov tests).

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