Flexible degrees of connectivity under synaptic weight constraints

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Abstract
Biological neural networks face homeostatic and resource constraints that restrict the allowed configurations of connection weights. If a constraint is tight it defines a very small solution space, and the size of these constraint spaces determines their potential overlap with the solutions for computational tasks. We study the geometry of the solution spaces for constraints on neurons’ total synaptic weight and on individual synaptic weights, characterizing the connection degrees (numbers of partners) that maximize the size of these solution spaces. We then hypothesize that the size of constraints’ solution spaces could serve as a cost function governing neural circuit development. We develop analytical approximations and bounds for the model evidence of the maximum entropy degree distributions under these cost functions. We test these on a published electron microscopic connectome of an associative learning center in the fly brain, finding evidence for a developmental progression in circuit structure.

1 Introduction
Learning in neural networks is the process of finding network structures that minimize a cost function. The capacity of a network (the number of computations it can successfully learn) depends on a number of factors. For associative memories, the capacity depends on the structure of the inputs [3], the learning rule [6], and constraints on the connectivity [1]. In biological neural networks, the cost function, learning rule, and structure of input activity are often unknown. Biological circuit structure is shaped by developmental programs and slow structural plasticity that construct a scaffold for and stabilize learning and memory on faster timescales [5]. Motivated by this, we consider the hypothesis that developmental programs that structure circuits might aim for flexibility: to allow many different weight configurations under fixed constraints.

The total strength of synaptic connections between two neurons is limited by the amount of receptor and neurotransmitter available and the size of the synapse [4]. Pyramidal neurons of mammalian cortex and hippocampus undergo synaptic scaling, regulating their total synaptic input strengths to regulate postsynaptic activity levels [7]. Motivated by these observations, we consider simple models of resource limitations and homeostatic constraints on total and individual synaptic weights. We examine how the size of the solution space for these constraints depends on the number of connections (the degree) and compute the optimally flexible degrees under different constraints on total and individual connection strengths. We then develop the maximum entropy degree distributions under these constraints. We derive the Laplace approximation for the evidence of these degree distribution models. Finally, we apply these models to a recently characterized connectome of a learning and memory center of the larval Drosophila melanogaster [2], asking which constraints best explain the degree distributions of neurons at different developmental stages. We find that overall, a homeostatically fixed net weight best predicts the degree distributions of Kenyon cell inputs and outputs. The most mature Kenyon cells, however, are better explained by a simple binomial random wiring model, suggesting a developmental progression in the cost functions governing mushroom body wiring.

2 Geometry of constraint spaces

We consider a simple model of synaptic interactions where a neuron has \( K \) synaptic partners and the total strength of projection \( i \) is \( J_i \). \( K \) is the synaptic degree. We assume that the existence of a connection is determined separately from its strength, and model the synaptic weights as continuous variables. A particular configuration of synaptic weights occupies a point in the \( K \)-dimensional synaptic weight space.

Given the degree \( K \), a constraint on synaptic weights defines a solution space. The size of that solution space measures the flexibility of the constraint. Importantly, we non-dimensionalize the synaptic weights so that the size of these constraint spaces are dimensionless quantities. For any \( K \), the size of the constraint space approximates the number of \( K \)-dimensional synaptic weight configurations allowed.

2.1 Flexibility under bounded net synaptic weight

We begin by considering an upper bound on the net synaptic weight, so that

\[
\sum_{i=1}^{K} J_i \leq \bar{J} K^p
\]

This bound could be interpreted multiple ways, for example as a presynaptic limit due to the number of vesicles currently available before more are manufactured or a postsynaptic limit due to the amount of dendritic tree available for synaptic inputs. Scaling the summed synaptic weight as \( K^p \) corresponds to scaling the individual synaptic weights as \( K^{p-1} \). If every synaptic weight has an order \( 1/K \) strength, the sum of the synaptic weights would be order 1 and \( p = 0 \). If every synaptic weight has an order 1 strength, the summed weight is order \( K \) and \( p = 1 \). If synaptic weights have balanced \((1/\sqrt{K})\) scaling \( [8] \), then the summed weight would have \( p = 1/2 \).

With \( K \) synaptic partners, the solution space for Eq. [1] is the volume under a \( K-1 \) simplex (Fig. [1]). Thus, for the bounded weight constraint the number of weight configurations is proportional to the volume of the \( K-1 \) dimensional simplex, \( V(K, \bar{J}) = (\bar{J} K^p)^{K-1}/K! \) (Fig. [1]). We can also view this as a count of available configurations if we divide the maximum synaptic weight \( \bar{J} \) into \( N \) increments of measurable synaptic weight changes \( \Delta J \), and measure synaptic weights relative to \( \Delta J \). In the continuum limit \( N \to \infty \), \( \Delta J \to 0 \) with \( J \) fixed, the volume under the simplex approximates the number of synaptic weight configurations. We call the synaptic degree that maximizes the volume under the simplex the optimal degree, \( K^* \). We computed this optimal degree. It is approximately linearly related to the total synaptic weight:

\[
(K^*)^p \bar{J} = \left( K^* + \frac{1}{2} \right) \exp (-p) + \mathcal{O} (1/K^*)
\]

with a slope that depends on \( p \) (Fig. [1]). We can see from Eq. [2] that if \( p = 1 \), we obtain the condition \( \bar{J} = 1/e \) (to leading order). So if \( p = 1 \) and \( \bar{J} = 1/e \), the volume is approximately independent of \( K \). If \( p = 1 \) and \( \bar{J} < 1/e \), the volume decreases monotonically and vice versa.

2.2 Flexibility under fixed net synaptic weights

Motivated by the observation that different types of neuron regulate their total synaptic weights \([7]\), we also consider a simple model of homeostatic synaptic scaling: \( \sum_{j=1}^{K} J_j = K^p \bar{J} \). The fixed net weight constraint defines the same simplices as the bounded net weight, but requires synaptic weights to live on their surfaces instead of the volumes under them (Fig. [1]). The size of this space of allowed weights is given by the surface area of the \( K-1 \) simplex, \( A(K, \bar{J}, p) = (K^*)^p \sqrt{\bar{J}/(K-1)} \). The surface area of the simplex increases with the net excitatory weight, but for \( \bar{J} \geq 1 \) it has a maximum at positive \( K \) (Fig. [1]).

The optimal degrees obey:

\[
(K^*)^p \bar{J} = (K^* + p - 1) \exp (-p) + \mathcal{O} (1/K^*)
\]

revealing an approximately linear relationship, similar to the constraint on the maximum possible synaptic weight (Eq. [2]). As for the bounded net weight, we can see from Eq. [3] that if \( p = 1 \), we obtain the condition \( \bar{J} = 1/e \) (to leading order). So if \( p = 1 \) and \( \bar{J} = 1/e \), the surface area is approximately independent of \( K \).

If \( p = 1 \) and \( \bar{J} < 1/e \), the area decreases monotonically and vice versa.
2.3 Flexibility under individual connection strength bounds

We consider a simple model for resource limitations at individual connections: $J_j \leq \bar{W} K^{p-1}$. The scaling with $K^{p-1}$ here ensures that the sum of $K$ synaptic weights scales as $K^p$, as for the previous constraints. The volume of the hypercube, $C = (\bar{W} K^{p-1})^K$ measures the size of the solution space. If $p = 1$ here (individual synaptic weights do not scale with $K$) then the volume of the cube only decreases with $K$ for $\bar{W} < 1$ and increases with $K$ for $\bar{W} > 1$. If $p < 1$, however, the volume exhibits a maximum at positive $K$ (Fig. 1h). At those maxima,

$$\bar{W}(K^*)^{p-1} = \exp(1 - p)$$

In contrast to the constraints on the total connection strength, the upper limit for each connection strength is independent of $K$. In all these cases, the value of the constraint at the optimal degree decreases with $p$.

3 Maximum entropy degree distributions under connectivity constraints

We postulated above that the entropy (log volume) of solution spaces provides a cost function for neural connectivity. Under this cost function, the maximum entropy degree distributions are proportional to the size of the solution space:

$$p(K | \bar{J}, p) = |S_K| / Z_S$$

where $|S_K|$ is the size of the solution set $S_K$. For large $K^\text{max}$, we compute the normalization constant as $Z_S(\bar{J}, p) = \sum_{K=1}^{\infty} |S_K|$. These provide predictions for neural degree distributions. To test them we turned to an electron microscopic reconstruction of connectivity of Kenyon cells in the larval Drosophila melanogaster’s mushroom bodies, a center of learning and memory in the fly brain [2]. These data include the number of synapses for each connection. To map these anatomical measurements onto our theory, we assume that the synapse counts are proportional to the physiological synaptic weights (for constraints on...
the net synaptic weight, $\bar{K}^p = \alpha\bar{S}$). We computed the Laplace approximation for the model evidence under each of the constraints discussed above, marginalizing out the scale factors relating the anatomical measurements to the modeled synaptic weights. We also computed the model evidence for a binomial random wiring model.

The models with bounded individual synaptic weights provided the poorest explanations for KC connectivity degrees (Fig. 2b, c green). For single-claw KCs, the binomial wiring model had the highest evidence (Fig. 2b, c black; log likelihood ratio at least 1.57 for binomial vs fixed or bounded net weights for in-degrees, at least 0.78 for out-degrees). For young and multi-claw KCs, the fixed net weight had the highest evidence (Fig. 2b, c orange; log likelihood ratio at least 70.46 for fixed net weight vs bounded or binomial models on young KC in-degrees; at least 9.92 for multi-claw KC in-degrees; at least 48.17 for fixed net weight vs binomial on young KC out-degrees; at least 0.41 for fixed vs bounded net weight on young KC out-degrees; at least 201.8 for fixed net weight vs binomial on multi-claw KC out-degrees; at least 20.18 for fixed vs bounded net weight on multi-claw KC out-degrees). This suggests that less mature KCs have connectivity governed by a homeostatically regulated total input and output strength and as KCs mature, other factors come to dominate their wiring.

KCs can be morphologically classified by the structure of their dendrites. Immature KCs have smooth dendrites, while more mature KCs’ dendrites exhibit claws around input axons. Single-claw KCs are more mature than multi-claw KCs [2]. For models with a fixed net weight, the normalization constant was not tractable; because of this we computed bounds for the evidence (Fig. 2b, c orange shows the lower bounds).

The models with bounded individual synaptic weights provided the poorest explanations for KC connectivity degrees (Fig. 2b, c green). For single-claw KCs, the binomial wiring model had the highest evidence (Fig. 2b, c black; log likelihood ratio at least 1.57 for binomial vs fixed or bounded net weights for in-degrees, at least 0.78 for out-degrees). For young and multi-claw KCs, the fixed net weight had the highest evidence (Fig. 2b, c orange; log likelihood ratio at least 70.46 for fixed net weight vs bounded or binomial models on young KC in-degrees; at least 9.92 for multi-claw KC in-degrees; at least 48.17 for fixed net weight vs binomial on young KC out-degrees; at least 0.41 for fixed vs bounded net weight on young KC out-degrees; at least 201.8 for fixed net weight vs binomial on multi-claw KC out-degrees; at least 20.18 for fixed vs bounded net weight on multi-claw KC out-degrees). This suggests that less mature KCs have connectivity governed by a homeostatically regulated total input and output strength and as KCs mature, other factors come to dominate their wiring.

References


