Modelling brain-wide neuronal morphology via rooted Cayley trees

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Neuronal morphology is an essential element for brain activity and function. We take advantage of current availability of brain-wide neuron digital reconstructions of the Pyramidal cells from a mouse brain, and analyze several emergent features of brain-wide neuronal morphology. We observe that axonal trees are self-affine while dendritic trees are self-similar. We also show that tree size appear to be random, independent of the number of dendrites within single neurons. Moreover, we consider inhomogeneous branching model which stochastically generates rooted 3-Cayley trees for the brain-wide neuron topology. Based on estimated order-dependent branching probability from actual axonal and dendritic trees, our inhomogeneous model quantitatively captures a number of topological features including size and shape of both axons and dendrites. This sheds lights on a universal mechanism behind the topological formation of brain-wide axonal and dendritic trees.

Neurons, the primary components of central nervous system are electrically excitable cells that receive, process, and transmit information through electrical and chemical signals between each other. Digital reconstructions of neurons provide information for quantitative measurements of neuronal morphology. With digital reconstructions, many studies have highlighted the importance of neuronal morphology in its biological function. Mainen and Sejnowski¹ have shown a causal relationship between dendritic structures and intrinsic firing patterns observed from in vitro electrical recordings for a wide variety of cell types. Vetter et al. have shown that branching pattern strongly affected the propagation of action potentials which links information processing at different regions of the dendritic tree². Ferrante et al. have shown that even subtle membrane readjustments at branch points could drastically alter the ability of synaptic input to generate and propagate the action potentials³. It has also been shown that the input-output response function of neuron’s dendritic arbor grows with the tree size⁴,⁵. More recently, Yi et al. have demonstrated a crucial role of neuronal morphology in determining field-induced neural response⁶. These results, among others, have contributed to a now widespread acceptance that neuronal morphology plays a critical role in its activity and function.

The formation of neurons through branching is driven by complex interactions of intracellular and extracellular signaling cascades which are proving difficult to be completely understood by molecular biology alone. Mathematical or computational modelling instead provides an alternative and complementary approach to uncover mechanism underlying neuronal morphology. Simulators L-Neuron⁷,⁸ and NeuGen⁹ were developed to create virtual neuronal structures in silico by means of iteratively sampling experimental statistical distributions of shape parameters (including e.g. branch diameter, length, ect). In contrast to L-Neuron or NeuGen simulators which require a large number of experimental neuron samples to obtain reliable distributions of shape parameters, Van Ooyen et al. used models in which the morphology of a single dendritic tree was represented in a highly abstract manner¹⁰–¹³ where growing neurons were modeled as expanding, circular neuritic fields. Based on such phenomenological dendritic growth models, NETMORPH was developed to simulate 3D neuronal networks from the perspective of individual growth cone¹⁴, using simple rules for neurite branching at each terminal segment. Other modelling approaches with simple rules to account for the spatial embedding of tree structures, include optimal rewiring and particle-based diffusion limited aggregation approach: the optimal rewiring approach generates branching geometry by minimizing the wiring length and the path-length to

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root between branch points and synapses in dendrite trees\(^{15,16}\), the particle-based diffusion limited aggregation approach provides model-based measures to estimate “diffusive” shape of neuronal tree-like structures\(^{17,18}\).

However, analysis and modelling of neuron morphology to the best of our knowledge so far are largely based on neuron digital reconstructions from certain regions or layers of a brain (e.g.\(^{7,14}\)). Moreover, a majority of these neuronal morphology analysis and modelling concentrate on neuron dendrites, with only a few studies of axonal branching structure in certain layers recently (e.g.\(^{19,20}\)). Knowledge about how dendrites and axons branch in an entire brain is still limited. Recently, breakthroughs in imaging\(^{21,22}\) and molecular labeling\(^{23,24}\) techniques have provided tools to trace and digital reconstruct the almost complete morphology of neuronal populations at a single-axon resolution through a whole brain\(^{25,26}\); this offers the opportunity to quantify brain-wide dendritic and axonal branching morphology. In this manuscript, we take advantage of current availability of brain-wide neuron digital reconstructions of pyramidal cells from a mouse brain\(^{25}\) and analyze the complete axonal and dendritic branching morphology. In particular we show that axons are topologically self-affine whereas dendrites are topologically self-similar, and topological structures of both axon and dendrites are far away from symmetry and appear to be random. We also develop a self-organized probabilistic model for the entire axonal and dendritic branching structures. In contrast to the growth models\(^{13}\) which require a simulation/growth time to create virtual trees of finite size, we use the generation of rooted 3-Cayley trees (where non-terminal nodes are linked to 3 neighbors; in neuron trees, 3 linkages of a node represent one mother branch and two daughter branches) via a probabilistic process to create virtual trees of infinite size, which are used to quantify several statistical properties of neuronal morphology including distributions of neuron size and topological width/length.

Results

Brain-wide axonal and dendritic morphology quantifications. In this section, we analyse neuron morphology from 35 brain-wide digital reconstructions of pyramidal neurons in a mouse brain\(^{25}\). A typical neuron consists of a cell body (soma), dendrites, and an axon. Here we analyze axon and dendrite tree morphology in aspects of size, asymmetry, and shape of trees as well as correlations among axons and dendrites in single neurons.

Topological structure of axonal and dendritic arbors. We first focus on the topological structure of dendritic and axonal branching from soma to neuron terminals, despite the spatial position of each branch. Figure 1 illustrates an example of a single brain-wide neuronal morphology and its topological branching structure. We separate individual neurons into one axon and several dendrites for analysis; see Fig. 1 as an example. From 35 actual neuron reconstructions, we find that on average there are 6.8 ± 0.56 number of dendrites besides one axon. Neurons have been widely viewed as binary trees (e.g.\(^{27}\)), here we first examine this by looking into the degree (i.e. the number of branches each node links to) of each node. Clearly, each terminal is connected to one branching node (i.e. degree 0) and branching nodes are mainly linked with three branches in both axons and dendrites - with only <1% of branching nodes in axons and 2% in dendrites linking to more than 3 branches; those could be due to limited spatial resolution in imaging which leads to possible errors in digital reconstructions. Overall, this suggests that brain-wide neurons are well approximated as binary trees.

We quantify the number of branching nodes (which is equivalent to the number of terminals minus one) for both axonal and dendritic trees, and refer as the topological size. On average, we find that axons are of size 224.1 ± 20.89\(^{(n = 35)}\), whereas dendrites are significantly smaller, of size 8.29 ± 0.72\(^{(n = 239)}\), i.e. about 1/27 of the axonal tree size. We also classify the branching nodes into three types\(^{28}\): B-type if both child branches themselves bifurcate; M-type if only one child branch bifurcate; S-type if branching with two terminals. By such...
classification, we find that axons have smaller proportion of S-type nodes whereas dendrite trees have larger proportion of S-type branching nodes; see Fig. 2(a).

We characterize the topological shape of neuron trees by measuring its topological width and length as illustrated in Fig. 1(c). To do this, we first give an order (denoted as \( k \)) for each node as its topological distance to the soma, i.e. the number of branches in its path to the soma. The topological length (denoted as \( k_{\text{max}} \)) of a tree is then defined as the maximal order among nodes. For the topological width, we calculate the number of branching nodes (denoted as \( j_k \)) for each order \( k \) and define the topological width (denoted as \( j_{\text{max}} \)) of a tree as the maximum among all \( j_k, k = 1, \ldots, k_{\text{max}} \). This is similar to Sholl analysis\(^{26}\) when topological distance is applied. In the population of our 35 actual brain-wide neuron reconstructions, we observe that on average \( k_{\text{max}} \propto N^\lambda, j_{\text{max}} \propto N^\lambda \) and the scaling exponents are different between axons (\( \tau = 0.503 \pm 0.035, \lambda = 0.533 \pm 0.036 \); see Fig. 2(b)). Note that in axons, exponents \( \tau \) and \( \lambda \) are significantly different (F test; \( p = 0.012 \)) where in dendrites there is no significant difference between the exponents (F-test \( p = 0.97 \)); this suggests that in aspects of topological shape, dendrites are approximately self-similar (i.e. \( \tau \approx \lambda \)) whereas axons are self-affine (i.e. \( \tau \sim \lambda \)).

Next, we quantify morphological asymmetry of neuron trees. Following the definition in\(^{31}, \) tree asymmetry \( A \) describes the average of local partition asymmetry \( A_{sp}(r, s) = \frac{|r - s|}{r + s} \) for all branching nodes (here \( r, s \) are the number of terminals in two subtrees from a branching node) except for those with partition (1, 1) (i.e. branching to two terminals). This tree asymmetry reflects the relative balance of branching within a tree based on the distribution of terminals between two subtrees bifurcated from a node. This tree index range forms from 0 (symmetry) to 1 (asymmetry). The axon and dendrite from brain-wide neuron digital reconstructions show no significant difference (student \( t \)-test \( p = 0.274 \)) on tree asymmetry with \( A = 0.791 \pm 0.006 \) (median \( n = 35 \)) for axonal trees and \( A = 0.743 \pm 0.019 \) (median \( n = 211 \)) for dendritic trees and both are far away to symmetry. These measured partition asymmetry values are close to reported values in http://neuromorpho.org/. Another asymmetry measurement - excess partition asymmetry introduced by Samsonovich and Ascoli\(^{32}\) considers the difference between partition asymmetry actually measured at one branch and the average of partition asymmetry computed for the same branch after all possible shuffling of the granddaughter branches. This excess partition asymmetry is useful to test the randomness in branching or the existence of control/regulation process within neurons. Interestingly, in contrast to ‘regulation mechanism’ suggested in\(^{32}\) based on systematically positive excess partition asymmetry \( E_p \) measured from pyramidal cells on some slice of a brain\(^{33}\), we show that our brain-wide axonal and dendritic trees appear to have almost no excess partition asymmetry with \( E_p = 0.012 \pm 0.005 \) (median \( n = 35 \)) (t-test to zero mean with \( p = 0.013 \)) for axons and \( E_p = 0.004 \pm 0.006 \) (median \( n = 151 \)) (t-test to zero mean with \( p = 0.054 \)) for dendrites. This suggests branching in both axonal and dendritic trees appears to be random.

**Geometric size of axons and dendrites.** Regarding geometric size of neuron trees, we measure the total branch length \( L \) of a tree (i.e. the sum of all branch length in a tree). On average, axons have a total branch length of \( 5.97 \times 10^4 \pm 0.55 \times 10^3 \mu m \) (median \( 5.86 \times 10^3 \mu m \)) and dendrites have a total branch length \( 9.6 \times 10^4 \pm 0.99 \times 10^3 \mu m \) (median \( 5.6 \times 10^3 \mu m \)). Moreover, we observe that the total length \( L \) strongly correlates with tree size \( N \) (Pearson \( r = 0.68, p < 0.0001 \) for axons and \( r = 0.91, p < 0.0001 \) for dendrites) and increases approximately linear as the tree size \( N \) for both axons and dendrites; see Fig. 3 top panels. The mean total length is expected to be the mean branch length (say \( m \)) multiplied by the number of branches \( 2N + 1 \) for a given tree size \( N \). The data show a best linear fit to the function \( L = m(2N + 1) \) with a mean branch length \( m = 123.2 \mu m \) for axons and \( m = 59.9 \mu m \) for dendrites. Such strong correlations and similar linear slopes between axon length and its tree size were also observed in vitro axons of human neurons across all layers of medial temporal cortex, though total length of axons in human brains is much larger than that in mouse\(^{34}\); this suggests a potential universal mechanism controlling branch length cross species. We also note that in our brain-wide neurons, though the total branch length approximately grows linearly with the number of branches, the branch length is not uniform.

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**Figure 2.** Tree branching nodes and topological shape. (a) shows proportion of each type of branching nodes in axons (\( n = 35 \)) and dendrites (\( n = 239 \)) from 35 actual brain-wide neuron reconstructions. (b) shows topological length and width and their scaling with tree size as \( k_{\text{max}} \sim N^{\lambda} \) and \( j_{\text{max}} \sim N^{\tau} \); each dot represent the \( k_{\text{max}} \) (black) or \( j_{\text{max}} \) (grey) averaged over trees with the same tree size \( N \). Exponents \( \lambda \) and \( \tau \) are indicated in each panel for both axons and dendrites. They show significant difference in axons (F-test; \( p = 0.012 \)) and no significant difference in dendrites (F-test; \( p = 0.97 \)).
along orders; see Fig. 3 bottom panels for both axons and dendrites. In particular, the branch length of axons is larger at intermediate orders than those at smaller or larger branch orders.

**Correlation among axons and dendrites in single neurons.** It has been suggested that in one given portion of a neuron, dendritic morphology may be under intrinsic homeostatic control which regulates tree size fluctuations systematically by counterbalancing the remaining dendrites in the same cell\(^3\). We test such a control mechanism in axons and dendrites from our actual brain-wide neurons. If trees of a neuron were mainly up(down)-regulated by common factors, then neurons with larger axons in terms of tree size would have larger dendrites; if axons and dendrites were mutually regulated by competition, then neurons with larger axons would have smaller dendrites. Based on calculated tree size of axons, average tree size among dendrites and the number of dendrites for each of 36 neurons, we perform Pearson correlation analysis among these quantities. Interestingly, we find no significant correlation between axon tree size and average dendrite tree size (\(p = 0.98\)), between axon tree size and number of dendrites (\(p = 0.23\)), or between average dendrite tree size and the number of trees (\(p = 0.061\)). These suggest that axon tree size, dendrite tree size and dendrite number in single neurons are likely to be independent. Moreover, if tree size is regulated within single neurons, then tree shuffling would lead to an increase in the variance of average tree size in the neuron population; if tree size is random among populations, then shuffling would not lead to significant change on the variance\(^3\). To further test such randomness, we consider neurons with the same number of dendrites. Then we randomly shuffle the dendrites in these neurons and calculate the standard deviation (std) of the average dendrite tree size among single neurons. Figure 4(c) show that 35.5% of 1000 random shuffling gives higher std of the average dendrite tree size than the std from actual neurons, suggesting tree size of dendrites within single neurons is random.

**Modelling topology of axonal and dendritic trees.** As indicated by above statistical analysis, the tree size of dendrites and axons appear to be random independent of the number of dendrites, indicating that the formation of brain-wide neuron trees is likely to be a stochastic process. In this section, we use stochastic generation of rooted z-Cayley trees\(^3\) to model the topological structure of axonal and dendritic arbors. A z-Cayley tree is a tree where non-terminals have z-neighbours. Here to model neuron trees, we use \(z = 3\) as non-terminal nodes in neuron trees are shown to mainly have 3 branches (3 neighboring nodes): one in previous order and two in the next order. In the generation of a rooted 3-Cayley tree, we start with an arbitrary node as the root (which represents the soma in a neuron tree), and assign an link from this root connecting to another node (viewed as

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**Figure 3.** Tree geometric length. Top two panels show total branch length \(L\) (measured as the sum of all branch length) as a function of topological tree size \(N\); lines give the best fitting to the function \(L(N) = m(2N + 1)\) with \(m = 123.2 \mu m\) for axons and \(m = 59.9 \mu m\) for dendrites. Bottom two panels show the average branch length (mean ± sem is shown) at each order \(k\) for axons (left) and dendrites (right).
the 1-th order node); the generation of tree proceeds from this 1-th order node, by (1) bifurcating to two daughter branches (linking to 2 neighbouring nodes in the next order) and (2) randomly selecting its neighboring nodes in the next order with certain probability; if one node in the k−th order is selected, then it bifurcate into two daughter branches to next order k + 1 and random selection on its two neighbours applies independently; if a neighbouring node is not selected, then this neighboring node is considered as a terminal (which could not bifurcate into the next order). The procedure continues until no more nodes are selected to bifurcate and eventually generates a rooted 3-Cayley tree.

We first investigate the homogeneous branching model where all nodes are selected with the equal probability p. In this case it is well known that beyond a critical probability \( p_c = 1/2 \), a positive probability exists that an infinite tree is generated, whereas below this threshold, the probability for such infinite spanning is zero and a finite tree is generated. Moreover, the mean tree size of generated finite trees increases with branching probability as

\[
\langle N \rangle = \frac{1}{1 - 2p}, \quad \text{if } p < p_c = 1/2.
\]

In fact if we denote T by the mean number of branches in one branch from the 1-th order node, then by statistical equivalence between sub-branches \( T = p(1 + 2T) \), which gives \( T = p/(1 - 2p) \), and the mean tree size \( \langle N \rangle = 1 + 2T \) where 1 corresponds to the 1-th order branching node.

To model the axonal and dendritic trees, we estimate the constant branching probability \( p \) from Eq. (1) using measured average tree size, which gives \( p = 0.498 \) for axons and \( p = 0.44 \) for dendrites. Indeed, the model with estimated branching probability produce similar mean tree size as the actual data; see Fig. 5 top two panels. We then test this simple homogeneous model for other quantities described in the above section, such as proportions of 3 types of branching nodes (we take the B-type branching node as an example for the comparison), topological shape indexes \( (j_{\text{max}} \text{ and } k_{\text{max}}) \), and tree asymmetry A. Note that the model takes independent probability for each node and thus no excess asymmetry is expected. Figure 5 top panels show the relative difference of the modelling results to actual data on these quantities; in particular, we see that proportion of B-type branching node and topological width \( (j_{\text{max}}) \) from homogeneous model from both axons and dendrites are far different (>20% relative difference) to measurements from actual data. Moreover, distributions on tree size \( (N) \), tree topological length \( (k_{\text{max}}) \) and width \( (j_{\text{max}}) \) as shown in Fig. 5 bottom panels for both axons and dendrites are significantly different to corresponding distributions from actual axonal and dendritic trees \( (p < 0.0001 \text{ for both axons and dendrites}) \). We also notice that distributions from the homogeneous model illustrate higher frequency at lower values and larger variation on these quantities for both axons and dendrites, indicating that it has high probability to generate smaller tree than the expected mean. We thus conclude that homogeneous model is not sufficient to model the topology of neuron trees.

We next consider an inhomogeneous model where branching probability differs between nodes. In particular we investigate order dependent branching probability \( p_k \); such dependence was considered in the growth model and the computational model. Figure 6 indeed shows that the branching probability does depend on the order in particular at low orders. In the growth model for dendritic topology, branching probability is assumed to be decay exponentially to 0 as the order \( k \rightarrow \infty \). However, the observed branching frequencies for both axons and dendrites in our brain-wide neurons do not fit well to exponential decay with zero plateau, instead, the data exhibit steady branching frequencies at high orders \( k \) and statistically the data fit better to exponential decay with non-zero plateau in the form of \( p_k = b \exp(-ak) + c \) (F-test; \( p < 0.001 \) for both axons and dendrites). In the best fitting curve, as \( k \rightarrow \infty \), the branching probability \( p_k \rightarrow c < p_c = 0.5 \) (the critical probability); this ensures a finite
tree generation in our model\textsuperscript{35}. The plateau $c$ also reflects the almost constant branching probability at large orders where subtrees branch in an approximately “homogeneous” manner.

With the best fitting branching probability in the form of exponential decay to a non-zero plateau, the inhomogeneous model leads to mean tree sizes $\langle N \rangle \approx 214.3$ and 7.1 for axonal and dendritic trees respectively, which are close to measured values from actual data. We also test this inhomogeneous model on other topological quantities and show in Fig. 5 that the model not only recaptures the mean tree topological measurements (with relative difference $<20\%$ to measurement from actual neurons), but also agrees well on their distributions in particular the tree size and topological shapes $k_{\text{max}}$ and $j_{\text{max}}$ (two-sample Kolmogorov-Smirnov test with $p \geq 0.05$, except for test on $j_{\text{max}}$ of dendrite where $p = 0.01$). We next examine the scaling of topological shapes in relation to tree size. Figure 7 shows that the conditional averages of tree length and width scale with the tree size $N$ as $\langle |l| \rangle \sim \lambda N$ and $\langle |w| \rangle \sim \tau N$ with exponents $\lambda = 0.339 \pm 0.007$, $\tau = 0.754 \pm 0.009$ (significant difference between $\lambda$ and $\tau$; F-test; $p < 0.0001$) for axons and $\lambda = 0.631 \pm 0.048$, $\tau = 0.522 \pm 0.044$ (no significant difference between $\lambda$ and $\tau$; F-test; $p = 0.339$) for dendrites. These exponents from simulations are close to those estimated from actual neurons, showing that as similar as in actual neurons, simulated virtual axons are self-affine while simulated virtual dendritic trees are self-similar. These suggest that inhomogeneous model with a simple order dependent rule on branching probability is sufficient to statistically capture a number of topological features of both axonal and dendritic morphology in a quantitative manner.

**Statistical properties of the probability model.** In this section, we explore how topological characteristics of artificial trees generated from the inhomogeneous model with branching probability $p_k = \min \left\{ \exp \left(-ak\right) + c, 1 \right\}$, $\forall k \geq 2$ vary with parameters. Note that we take a min operation in $p_k$ here, considering a probability cannot be larger than 1. Also note that, if we denote $p_1$ as the branching probability for order $k = 1$, then $p_1 \equiv 1$, as the first branching is initially selected in the model.
Simply, the mean tree size which can be calculated as the total number of branching nodes from all orders, and read as

\[
\langle N \rangle = \sum_{k=1}^{\infty} k \left( 1 + \sum_{m=2}^{\infty} \prod_{n=m}^{k-1} P_n \right)
\]

(2)

For the homogeneous model \( p_m = p \), it reduces to Eq. (1). For the inhomogeneous model, this estimation agrees well with simulated data; see Fig. 8 top panels. As expected, tree size \( N \), tree length \( k_{\text{max}} \) and width \( j_{\text{max}} \) increase with decreasing decay rate \( a \) or increasing \( b, c \). Moreover, we note that for a similar variation of parameters, the decay rate \( a \) allows to give a broad range of values on topological quantities in particular the tree size \( N \). Furthermore, from the simulated data, we observe that on average tree length \( k_{\text{max}} \) is larger than tree width \( j_{\text{max}} \) when tree size is small, whereas when tree size is big (e.g. see simulations with small \( a \) on the top left panel in Fig. 8) tree length \( k_{\text{max}} \) is smaller than tree width \( j_{\text{max}} \) on average. This is further confirmed in Fig. 8 bottom panel, which illustrates changes of conditional average of tree length \( k_{\text{max}} \) and width \( j_{\text{max}} \) for various tree size \( N \). Note that the model has three parameters \( a, b \) and \( c \) and different combinations of the parameters could give the same tree size \( N \) depending on the branching probability expression on \( p_m \). However, from Fig. 8 bottom panel, we see that for the same tree size \( N \), the...
corresponding mean tree length and width from different combinations of parameters are similar. This suggests that the tree shape largely depends on the tree size regardless of the branching probability pattern along the orders in the model. Moreover, Fig. 8 bottom panel shows that for small magnitude of tree size \(N\), the mean tree length and width grow with the tree size \(N\) in a similar scale, i.e. \(\langle | \rangle \approx \langle | \rangle \lambda \tau N_{\text{max}}\); dendrite trees are particularly of this case. Whereas for a large magnitude of tree size \(N\), the mean tree width grow faster than tree length when increasing tree size \(N\), i.e. \(\lambda < \tau\); axons are particularly of this case. An intersection between tree length and tree width in relation with tree size occurs at around \(N = 400\); for tree size much larger than this, the mean tree width is larger than mean tree length (i.e. \(\langle | \rangle > \langle | \rangle\)). These could be used as predictions for topological tree length and width when knowing tree size.

**Conclusion and Discussion**

In this manuscript we examine several neuronal morphological features and develop an inhomogeneous model to generate finite realistic virtual axonal and dendritic tree structures. By analyzing topological shape (length and width) in relation with tree size, we find that axons exhibit an self-affine pattern while dendrites are self-similar. Moreover, analyses on excess asymmetry and dendrite shuffling, suggest that tree size appear to be random; this supports the idea of a probability model for neuron tree structures. Using inhomogeneous branching probability, our probability model generates finite virtual trees of statistically similar to brain-wide Pyramidal neurons in a number of features (including axonal and dendritic tree size and shape). In contrast to published simulation tools\(^7\text{–}^9,\text{13}\), this modelling approach provides a tool with simple rules in a self-organized manner to generate virtual topological neuron structures of finite size. We remark here that models using different order dependent branching probability functions (e.g. polynomial decay, modelling data not shown) which fit the measured branching frequency as shown in Fig. 6, are also able to recapture measured data. Mechanisms underlying the decay pattern of branching probability remains unclear; however order dependent branching frequency alone is thus sufficient to capture the tree topology.

Our modelling of brain-wide neuron branching topology can be extended to take geometric information of branches into account, e.g. branch length and branch angles as studied in\(^34,\text{36}\). Note that a branch in a real neuron is not a straight line between two branching points, but exhibit complicated curvature structures; see Fig. 1(a). Including curvature structures would thus be important for a complete and better understanding of mechanism underlying brain-wide neuron morphology and its relation to specific functions at a single neuron scale. One possibility for such extending could be the inclusion of a stochastic process for the outgrowth direction of each branch as considered in the NETMORPH\(^14\). Single neurons can expand towards different regions in the brain and different areas in the brain are specialized for different functions; e.g. left hemisphere of the brain is dedicated to language while the right hemisphere is involved in more creative activities such as drawing. Thus morphological difference between different regions in a single neuron scale is likely to link at with different region-specified...
functions[37–39]. Indeed, neuron morphology and function in different layers of brains have been studied[40,41]. However, we are still far away to understand neuron morphology-function relationship at a single neuron scale.

As a theoretical approach in studying morphological role of neurons in their function, our brain-wide modeling of neuron branching could be useful to study the enhancement of dynamic range[26], by creating virtual realistic neuron topologies. Traditional Cayley trees (where terminals are of the same order) considered as an excitable medium for input signal propagation has been recently used to investigate dynamic range[42]. With our modelling approach for realistic neuron tree topology, it would be interesting to explore statistical properties of neuron trees in relation with dynamic ranges of neurons. In particular, it has been suggested that larger tree size could give larger dynamic ranges[42]; this can be tested with realistic virtual neuron trees using our modelling approach instead of traditional Cayley trees. Moreover, using more realistic virtual neuron trees also allow investigating the impact of tree asymmetry as well as tree shape in dynamic range.

Methods
In this manuscript, we use 35 neuron digital reconstructions of pyramidal cells for analysis. Pyramidal neurons digital reconstructions are provided by authors in[25]. Briefly, neuron digital reconstructions were obtained as follows. The brain is from C57BL/6J mouse line, and its pyramid neurons are sparsely labelled with Adeno-associated virus (AAV). Fluorescence micro-optical sectioning tomography microscopy (fMOST) was used to image these labelled neurons which span different brain regions or even the whole brain. From the imaging dataset, GTree method was used to reconstruct these labelled neurons, all of which contained axonal and dendrite tree morphology. The animal experiments were approved by the Institutional Animal Ethics Committee of Huazhong University of Science and Technology, and all experiments were performed in accordance with relevant guidelines and regulations.

The digital reconstruction of each brain-wide neuron is stored in a morphological file (in “SWC” format) which includes position information of each traced point and its linkage points. For analysis, each empirically neuron is decomposed into one axonal tree and several dendritic trees; each tree stems from the same soma. The axon part is identified as the largest component (in terms of the number of branching points) among all subcomponents and the rest are considered as dendrites.

A subcomponent of digital reconstruction stemmed from the soma is excluded for analysis if there is no branching points in this subcomponent; this in fact corresponds to a tree of size \( N = 0 \). Note that there is a small proportion of branching points which bifurcate to more than 2 branches; this could possibly due to errors in digital reconstruction. We manually adjust this by splitting such branching points so that all branching points bifurcate to 2 branches.

Data Availability
The data that support the findings of this study are available from the corresponding author on request.

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Author Contributions
Y.H., C.L. and T.Q. performed the analysis, C.L. and Y.H. performed modelling, and Y.Z. and C.L. conceived and designed the project. All authors reviewed the manuscript.

Additional Information
Competing Interests: The authors declare no competing interests.

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