Functional Characterization of Three-Dimensional Cortical Cultures for In Vitro Modeling of Brain Networks

Millimeter-scale Ca++ imaging

Neuronal activity correlation matrix

Functional neural network analysis

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HIGHLIGHTS

3D biomimetic in vitro cultures engineered to study functional neural networks

Millimeter-scale network analysis using GCaMP6f expression and widefield microscopy

Connected, unspecialized networks form in the developing 3D neural culture

Network properties change upon pharmacological perturbations of neurotransmission
Functional Characterization of Three-Dimensional Cortical Cultures for In Vitro Modeling of Brain Networks

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SUMMARY

Three-dimensional (3D) in vitro cultures recapitulate key features of the brain including morphology, cell-cell and cell-extracellular matrix interactions, gradients of factors, and mechanical properties. However, there remains a need for experimental and computational tools to investigate network functions in these 3D models. To address this need, we present an experimental system based on 3D scaffold-based cortical neuron cultures in which we expressed the genetically encoded calcium indicator GCaMP6f to record neuronal activity at the millimeter-scale. Functional neural network descriptors were computed with graph-theory-based network analysis methods, showing the formation of functional networks at 3 weeks of culture. Changes to the functional network properties upon perturbations to glutamatergic neurotransmission or GABAergic neurotransmission were quantitatively characterized. The results illustrate the applicability of our 3D experimental system for the study of brain network development, function, and disruption in a biomimetic microenvironment.

INTRODUCTION

Three-dimensional (3D) neural cultures recapitulate the cell-cell and cell-extracellular matrix (ECM) interactions of the native brain tissue more accurately than conventional 2D cultures (Zhuang et al., 2018). Because they express in vivo-like gene, protein and biomarker expression patterns of development, diseases, and network firing, 3D tissue cultures are useful systems to study brain-like functions at the cellular and tissue level (Lovett et al., 2020). In the brain, adhesion contacts and axonal pathfinding are isotropically distributed in a 3D microenvironment, with neurons synapsing with their immediate neighbors as well as distant neurons in all three dimensions to form extensive neural networks. These interactions are not recapitulated in 2D cultures, where neuronal connections are restricted unnaturally to the x-y plane (Ulloa Severino et al., 2016).

Several 3D culture platforms have been developed to model brain tissues in the laboratory. These platforms generally utilize self-assembly and self-organization principles (e.g., spheroids and organoids) (Birey et al., 2017; Dingle et al., 2015; Lancaster et al., 2013; Quadrato et al., 2017), with or without biomaterial support (e.g., hydrogels or scaffolds) (Frega et al., 2014; Tang-Schomer et al., 2014; Ulloa Severino et al., 2016). One example is our versatile composite scaffold consisting of silk and ECM components for 3D in vitro culture. These systems accommodate the long-term growth and function of rodent primary neurons as well as human neurons derived from induced pluripotent stem cells (iPSCs), induced neural stem cells (iNSCs), or other cell types isolated from patient brain or tumor explants (Cairns et al., 2016; Cantley et al., 2018; Lovett et al., 2020; Rouleau et al., 2020; Sood et al., 2016, 2019; Tang-Schomer et al., 2014).

One of the key challenges with using 3D in vitro cultures is the functional analysis of the engineered brain-like circuitry. Among the neural network analysis technologies, multi-electrode arrays (MEA) and optical-based approaches such as calcium (Ca++) and voltage imaging provide investigators detailed information on the spatial-temporal aspects of brain network development and function of rodent and human neurons in vitro (Badura et al., 2014; Poli et al., 2015; Trujillo et al., 2019). Brain network architecture is a hierarchical entity at multiple spatial scales ranging from connections between synapses to connections between brain regions (Bassett et al., 2018; Betzel and Bassett, 2017). Although MEA offer excellent temporal resolution,
cells needs to be in proximity to the electrodes to be detected, and the spatial resolution is physically fixed to the number and spacing of electrodes (typically in the tens to hundreds of μm) (Bourke et al., 2015; Obien et al., 2018). In contrast to MEA, optical-imaging-based approaches using dyes or genetically encoded reporter constructs have the spatial resolution as high as the microscope equipment allows (typically sub-μm) and have the potential to record contribution of all cells, enabling researchers to conduct analysis at different spatial scales. An optical approach can also be more widely used than MEA due to MEA’s requirement of dedicated equipment. Optical analyses of neural networks generally focus on cellular activity patterns at a fraction of 3D space (100–500 μm field width at a single x-y plane) due to the technical limitations of imaging equipment (Bosi et al., 2015; Gu et al., 2016; Huang et al., 2014; Sakaguchi et al., 2019; Ulloa Severino et al., 2016), with few exceptions (Dana et al., 2014; Lu et al., 2020; Palazzolo et al., 2017; Wu et al., 2020). There is a demand for tools to evaluate functional networks at larger, millimeter-scale in 3D cultures.

The objective of the present study was to develop an experimental 3D in vitro system that addressed this need through the incorporation of bioengineering principles, genetically encoded calcium indicators (GECIs), widefield microscopy image acquisition, and computational analysis. The output of this experimental system is a set of mathematical parameters describing the functional neural networks that investigators can use to quantitatively compare networks formed under different conditions. Primary mouse cortical neurons were cultured in 3D scaffolds and virally infected with genetic constructs expressing the calcium reporter GCAMP6f (Chen et al., 2013) to monitor activity. Network analysis showed functionally connected neural networks in the 3D cortical cultures. We demonstrated the utility of this experimental system with pharmacological perturbation of glutamatergic excitatory inputs (treatment with D-2-amino-5-phosphonovalerate, AP5; or 2,3-Dioxo-6-nitro-1,2,3,4-tetrahydrobenzo[f]quinoxaline-7-sulfonamide, NBQX) or γ-aminobutyric acid (GABA) inhibitory inputs (treatment with bicuculline or picrotoxin) and presented quantitative outcomes in the network descriptors. Bicuculline and picrotoxin increased neuronal activity, NBQX greatly reduced neuronal activity and impaired the functional network, and AP5 reduced neuronal activity to an even greater degree than NBQX. Interestingly, between the two GABA receptor antagonists, only bicuculline resulted in changes in the functional network properties at the scale we analyzed. Collectively, these data demonstrate the applicability of our 3D in vitro system for understanding the links between neurotransmission and network connectivity and for further investigation of the function and disruption of neural networks in a pathophysiological context.

RESULTS

3D In Vitro Biomimetic Cortical Culture System to Study Neural Network Structure and Function

To investigate neural network formation in 3D microenvironments, we adapted our previously published in vitro composite (silk and collagen) scaffold culture system that mimics the softness of native brain tissue and allows for growth and interaction of neurons and glial cells in three dimensions (Chwalek et al., 2015; Tang-Schomer et al., 2014). Additional elements incorporated into the system in this study were (1) expression of a GECI, GCaMP6f (Chen et al., 2013), as a reporter of neuronal activity, (2) widefield epifluorescence microscopy to record activity at micrometer- and millimeter-scales, and (3) computational tools adapted from published algorithms to investigate functional connectivity of the neural network (Muldoon et al., 2016; Newman, 2006; Onnela et al., 2005; Patel et al., 2015). The scaffolds were fabricated following established protocols, with dimensions optimized for high density cell seeding and for imaging the neural network in one single field of view (scaffold diameter = 3 mm, height = 1.5 mm, volume = 10.6 mm³ and pore size = 425–500 μm) (Figures 1 and S1). Two million cells were seeded per scaffold (Figure S1). The seeding efficiency was approximately 60%–70%, determined by amount of DNA in the 3D culture relative to the number and spacing of electrodes (typically in the tens to hundreds of μm) (Liaudanskaya et al., 2020) and detailed in Transparent Methods). We observed an increase in neurite density over the course of development (Figure 2C). To provide further evidence of structural network formation, quantitative real-time PCR (qRT-PCR) was used to analyze the expression of genes known to change during neural development. As anticipated, we observed a downregulation in mRNA expression...
levels of the neural progenitor marker (*Pax6*) and the immature neuron marker (*Dcx*) (Figure 2D) (Francis et al., 1999; Manuel et al., 2015). We detected a significant upregulation of mRNA encoding proteins with roles in presynaptic vesicle trafficking (*Syn1*), postsynaptic scaffolding (*Shank3*), glutamatergic neurotransmission (*Slc17a7, Gria1, Grin1*), and GABAergic neurotransmission (*GAD2, Gabra1*) (Figure 2D). The increased gene expression of the *Slc3a1* excitatory amino acid transporter together with immunofluorescence imaging of the astrocyte marker glial fibrillary acidic protein (GFAP) indicated the presence of astrocytes in the 3D cultures (Figure 3). Taken together, the data suggest neuronal maturation and synapse formation in the 3D culture model (Ben-Ari, 2001).

**3D Cultures Display Spontaneous Neuronal Activities Recorded with a Genetically Encoded Calcium Indicator**

For functional analysis of network activity in our 3D biomimetic cortical culture, a GCaMP6f (Chen et al., 2013) genetic construct was expressed in neurons for recording temporal calcium (Ca++) concentration changes, called Ca++ transients. The cultures were infected at 1 day in vitro (DIV) with adeno-associated virus serotype 1 (AAV1) driving expression in neurons of GCaMP6f under the control of the synapsin 1 promoter (Figure S1). A standard widefield fluorescence microscope with a 4× objective (field size = 4.15×3.51 mm²) was used to record the activity patterns from projection of the 3D culture (Figure 4A). Imaging was done for 1-min duration at 5 Hz, the temporal resolution that was sufficient to detect Ca++ events in a time-lapse series (Badura et al., 2014).

Neuronal activity was recorded at 2 and 3 weeks of culture, concurrent with the time frame in which rodent neurons grown in conventional 2D culture demonstrate increased activity (Chiappalone et al., 2006). At 2 and 3 weeks of culture, Ca++ transients were observed in individual neurons that synchronized within small, local clusters of neurons (size 30–100 μm) as well as larger clusters (>1 mm) (Figure 4A, Video S2). To perform quantitative analysis of the neural networks, the image areas were segmented into equal-size hexagonal regions of interest (ROIs) in a honeycomb configuration that covered the complete projection of the 3D culture sample (Figure 4B). We chose an ROI size of side-to-side distance of 500 μm for long-distance analysis based on previous studies, suggesting cortical neuron axon length *in vitro* (Virlogeux et al., 2018). It should be noted that the ROI size is user-defined, and computational analyses presented below were dependent on this initial ROI determination. Once an ROI mask was applied, fluorescence signal intensity from the projected image in each ROI was averaged to produce a time-series trace (Figure 4C). For each of the ROIs, Ca++ event detection was performed with the open-source software FluoroSNNAP (Patel et al., 2015). Spontaneous Ca++ events were present at both 2 and 3 weeks of culture in the majority (range 89–100%) of ROIs (Figures 4D, 4E, S3A, and S3B). The frequency of Ca++ events increased from 2 to 3 weeks (7.1 ± 0.7 events/min at 2 weeks, 12.9 ± 0.8 events/min at 3 weeks; mean ± SEM) (Figure 4F), correlating with the increased expression of markers of synaptic neurotransmission (Figure 2D). The Ca++ transients were abolished after tetrodotoxin (TTX) treatment, reflecting the action-potential-dependent nature of the neuronal activities (Figure 4D).

**Pharmacological Perturbations in Excitatory and Inhibitory Neurotransmission Alters Neuronal Activity of 3D Cultures**

To validate the responsiveness of our experimental system to network perturbations, a pharmacological strategy was employed to disrupt excitatory or inhibitory synaptic transmissions. The 3D cultures were treated with bicuculline (BIC, a competitive antagonist of inhibitory GABA_A receptors), picrotoxin (PTX,
a non-competitive antagonist of inhibitory GABA$_A$ and GABA$_P$ receptors), NBQX (a competitive antagonist of excitatory AMPA receptors), or AP5 (a competitive antagonist of excitatory NMDA receptors). Experiments were conducted on 3-week cultures because of their higher baseline activity (Figure 4F).

Spontaneous neuronal activities of each 3D culture sample were recorded and analyzed pre- (“baseline”) and post-treatment (Figures 5 and S3, Videos S3, S4, and S5, and Table S1 for detailed numbers and p values). Following treatments with BIC or PTX, Ca$^{++}$ events were present in 100% ROIs. Treatment with NBQX and AP5 significantly reduced the number of ROIs containing Ca$^{++}$ events (post-NBQX, 57 ± 11.8%; post-AP5, 30 ± 10.2%; mean ± SEM). In the AP5-treatment group, 5 of 11 samples (n = 4 experiments) had complete loss of neuronal activity (Figure 5B). Treatment with equivalent concentrations of DMSO, used to dissolve BIC and PTX, had no measurable effects on all the parameters reported in this study (Figure S4).

Both GABA receptor blockers resulted in an increase in Ca$^{++}$ event frequency (BIC group: baseline 10.9 ± 1.1, post-treatment 15.4 ± 0.7; PTX group: baseline 12.3 ± 1.2, post-treatment 17.8 ± 1.1 events/min; mean ± SEM) (Figure 5C). AMPA receptor blocker NBQX reduced the neuronal activity (baseline 10.5 ± 1.0, post-NBQX 2.7 ± 0.9 events/min). Treatment with the NMDA blocker AP5 resulted in greater reduction than NBQX (baseline 11.5 ± 1.6, post-treatment 0.4 ± 0.1 events/min) (Figure 5C). Together, these data show the presence of inhibitory and excitatory transmissions in the 3D tissue cultures and the differential responses of neurons to selective antagonists.

Figure 2. Structural Neural Network Development of the 3D In Vitro Biomimetic Cortical Culture

(A) Representative 3D projections of confocal z stack images of RFP-expressing neurons at 1, 2, and 3 weeks. Scale bar, 20 µm.

(B) Representative 3D rendering of side (x-z) view of RFP-expressing neurons at 3 weeks.

(C) Neurite density in the 3D cortical culture (mean ± SEM). n = 6-8 samples from three independent experiments.

(D) mRNA expression over time (mean ± SEM) of proteins associated with neural progenitors (Pax6), immature neurons (Dcx, doublecortin), presynaptic vesicles (Syn1, synapsin 1), postsynaptic scaffolding protein (Shank3), glutamatergic transmission (Slc17a7, vesicular glutamate transporter 1; Gria1, AMPA receptor subunit 1;Grin1, NMDA receptor subunit 1), and GABAergic transmission (GAD2, glutamate decarboxylase 2; Gabra1, GABA$_A$ receptor subunit $\alpha_1$). Expressions were normalized to housekeeping gene 18s. n = 7–11 samples from three to four independent experiments. For (C) and (D), ANOVA and post-hoc Tukey tests were used for statistical significance among groups. *p < 0.05; **p < 0.001. See also Video S1. See also Table S2.
Synchronization of neuronal activity in local clusters (5–50 neurons) has been observed in vivo during early development, with the level of synchronization changing in a brain-region-dependent manner as neural circuits become more complex (Garaschuk et al., 2000). For in vitro studies of networks, a numerical index is used to describe the degree of global synchronization. Here, the global synchronization index was computed by comparing the timing of events in all of the ROIs in each individual 3D culture using FluoroSNAP software (Patel et al., 2015). An average index of 0.29 ± 0.02 (mean ± SEM) was observed for the 3-week culture baseline conditions. No changes were observed with BIC or PTX treatments (BIC, 0.26 ± 0.04; PTX, 0.29 ± 0.05). In contrast, application of NBQX reduced the global synchronization index (0.14 ± 0.04), suggesting a reduction of global connectivity (Figure 5D). Global synchronization index was not computed for the AP5 group due to lack of neuronal activity.

**Functional Neural Network Characterization Using Graph Theory Models: Functional Connectivity, Clustering, Path Length, and Community Structure**

A fundamental goal of the field of network neuroscience is to understand brain networks as a complex web of interactions between functional units (Bassett et al., 2018; Lynn and Bassett, 2019). Graph theory is utilized extensively as a mathematical approach to analyze neural network functionality. To demonstrate the applicability of our 3D experimental system for evaluating functional neural networks, we applied previously published graph-theory-based network analyses to evaluate key attributes of the neural networks formed in our 3D cortical cultures at baseline and post-drug treatment conditions (Muldoon et al., 2016; Newman, 2006; Onnela et al., 2005). The basic principle of graph theory is to represent the network as a graph with each user-defined neural unit as a “node” and the connection between two nodes as an “edge” (Figure 6A). In this study, each of the 37 hexagonal ROIs is the equivalent of a node. The functional connectivity (edge) is defined as the statistical similarity of activity patterns of the two nodes, regardless of physical locations or the underlying structural connections.

The functional connectivity between node pairs in the 3D culture was assessed using cross-correlation analysis (Figure 6B). The cross-correlation coefficient of the paired nodes was used as the edge weight, a value to indicate the level of connectivity. These data were then used to compute average clustering coefficient, average path length, community structure, and node degree, which are each described in greater detail below (Figures 6C–6G and see Table S1 for detailed numbers and p values). Average edge weight of all node pairs at baseline and post-BIC, PTX, and NBQX treatments was calculated, but network analysis was not conducted for the AP5 group due to low level of activity. Average edge weight was significantly decreased by the AMPA receptor antagonist NBQX, suggesting a reduction in overall functional connectivity (Figure 6C). Interestingly, between the two GABA receptor antagonists, BIC treatment increased the average edge weight, whereas no significant changes were observed with PTX treatment. These results suggest that BIC, but not PTX, strengthened the overall functional connectivity of the 3D culture.

The average clustering coefficient, a measure of the intensity of functional connectivity among triplets of nodes (detailed in Transparent Methods), was evaluated for our cultures. A network with a high average clustering coefficient is thought to be better at local information integration and more robust to disruption...
In our cultures, an increase in the average clustering coefficient was observed with BIC treatment, whereas no significant changes were observed with PTX treatment. NBQX treatment resulted in a reduction in the average clustering coefficient (Figure 6D). This suggests that acute modulation of GABA and NMDA neurotransmission may alter information integration in our 3D neural networks (Bullmore and Sporns, 2009).

We similarly investigated path length as a descriptor of neural network functionality in our 3D cultures. Inversely proportional to edge weight, the path length is the shortest distance to traverse from one node to another node in the network. In classical network studies, a short average path length suggests that information can be efficiently shared across the network (Muldoon et al., 2016). However, it is important to mention the disagreements with this interpretation in the field (Fornito et al., 2016), and here we use path length as an indication of statistical dependence and not necessarily to characterize information routes. Average path length was calculated from all node pairs and normalized to the number of active nodes (detailed in Transparent Methods). BIC treatment reduced the average path length, whereas PTX had no significant effect (Figure 6E). Because average path length becomes artificially short when only a small portion of nodes remain active (Figure 6B), this descriptor was not calculated for the NMDA-treated cultures.

Many complex networks have community structures in which the network divides naturally into communities, or modules, containing densely interconnected nodes (Newman, 2006). It is hypothesized that community structures play an important role in functional specialization. Examining the community structures can thus provide insights to the specialization of the network (Bullmore and Sporns, 2009). The measurement that describes the strength of segregation of different parts of network into modules is called the

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**Figure 4. Spontaneous Neuronal Activity in the 3D In Vitro Biomimetic Cortical Culture**

(A) Representative fluorescence time-lapse image stacks of 3-week-old 3D cultures expressing GCaMP6f. Images are pseudo-colored to highlight Ca\(^{2+}\) transients. Scale bar, 500 µm.

(B) Mask with 37 hexagonal ROIs (dimensions: area = 0.22 mm\(^2\), side to side and center-to-center distances = 500 µm) is applied to the time-lapse image stack.

(C) Extraction of time-series trace of average fluorescence intensity from the projection of 3D culture in each ROI.

(D) Representative traces of spontaneous Ca\(^{2+}\) transients from individual ROIs at 2 weeks, 3 weeks, and 3 weeks post-tetrodotoxin (TTX, 40 µM) treatment. Fluorescence changes are normalized to the basal fluorescence intensity (dF/F\(_0\)).

(E) Percentage of all ROIs containing Ca\(^{2+}\) event(s) (mean ± SEM). Each data point represents a unique 3D tissue culture sample.

(F) Ca\(^{2+}\) event frequency at 2 and 3 weeks (mean ± SEM). Each data point represents average frequency of non-zero event ROIs of a sample. For (E) and (F), n = 23 samples from 3 to 4 matched experiments for 2-week and 3-week samples. An unpaired t test was used for statistical significance. **p < 0.001. See also Figure S5 and Video S2.
modularity (detailed in Transparent Methods). A higher modularity indicates that the network is more strongly divided and the nodes are more interconnected within modules than connected to nodes in other modules. At baseline conditions after 3 weeks of culture, two to three modules formed in the 3D neural network (Figure 6F, top panel) with low modularity (Figure 6F, bottom panel), suggesting a relatively weak community structure. BIC treatment reduced the modularity, whereas PTX had no significant effects. In the NBQX-treated 3D cultures, two different outcomes of community structures were observed. In cultures with fewer than 50% remaining active nodes (3 of 9 cultures), a single module with modularity of 0 was observed, suggesting that the network was deprived of a community structure after losing activity in the majority of nodes. In contrast, in cultures with 75%–100% active nodes, the modularity increased (6 of 9 cultures), suggesting that the network was more strongly divided (Figure 6F, bottom panel). Possible explanations to the variation among samples could be attributed to potential differences in expression level of AMPA receptors and relative level of AMPA receptor- and NMDA receptor-mediated glutamatergic neurotransmissions in each culture at this stage.

Another characteristic of complex networks is the presence of hubs, which facilitate efficient communication by shortening path lengths and bridging modules in a network (Barabási, 2009; Bullmore and Sporns, 2009). Hubs are nodes with a node degree (sum of edge weights touching the node) that greatly exceeds the average. The presence of hub nodes in a network can be inferred by the presence of a non-Gaussian distribution with a long right tail in a frequency distribution plot of node degrees (Schroeter et al., 2015). We computed the node degree for all of the individual nodes and graphed the frequency distribution. The absence of a long right tail in the frequency plots, combined with a normal distribution, indicate the absence of hub nodes under baseline conditions. Neither BIC nor PTX treatments resulted in the emergence of hub nodes. NBQX treatments resulted in a left shift of the node degree distribution because of the observed reduction in functional connectivity of the network, with 49% of the nodes having a node degree of zero (i.e. no remaining functional connectivity) (Figure 6G).

The above analyses were based on ROIs of side-to-side distances of 500 μm. To demonstrate that this model can be used at different spatial scales with other ROI sizes, we repeated the network analyses with smaller ROIs with side-to-side distance of 250 μm on BIC and PTX experiments (Figure S5). Our analyses showed that modifying ROI size from 500 μm to 250 μm changed the raw values of the network descriptors. It did not affect the pre- and post-treatment trends for synchronization, average edge weight, average path length, and modularity. At this scale of analysis, the increase of average clustering coefficient from pre- to post-PTX became statistically significant. This emphasizes that the network analysis is dependent on the initial determination of ROI, which should be chosen based on the question of interest.

In summary, the network analysis indicated that millimeter-sized, functional neural networks developed in the 3D biomimetic cortical cultures. The networks responded to pharmacological perturbations in
neurotransmission with measurable changes in the network descriptors. The average edge weight and average clustering coefficient increased with BIC treatment and decreased with NBQX treatment. The average path length decreased with BIC treatment. The networks at 3 weeks of culture had a weak community structure (modularity) and were reduced by BIC treatment. This, combined with the lack of hubs, suggested that unspecialized networks developed in our 3D cultures. These results demonstrate the utility of this 3D culture system for the study of functional neural networks in vitro.

DISCUSSION

Functional connection of a neural network is an important field of study for brain function and disorders. Conventional 2D culture with both rodent and human iPSC-derived neurons have been used to investigate neural networks using MEA and optical assays. For example, the correlation patterns of 2D rat cortical neuron cultures over the time frame of 7–35 DIV were evaluated with MEA of 60 electrodes with 200-μm spacing (Chiappalone et al., 2006). The emergency of hub structures in 2D mouse hippocampal neuron cultures were assessed with MEA of 64 electrodes with 200-μm spacing (Schroeter et al., 2015). In the past, network analysis has primarily been done with in-house, customized computer algorithms. FluoroSNNAP software, initially intended for 2D cellular scale analysis, was developed to facilitate the computational analysis of neural networks.
analysis in the field (Patel et al., 2015). Since that time, 3D in vitro cultures have become increasingly popular to investigate the development, functions, and disorders of brain networks as they better represent key features of the brain tissue (Lovett et al., 2020; Zhuang et al., 2018). One of the first studies demonstrating network activities in months-old human cortical organoids using MEA (200-μm spacing) and observing changes in synchronized network activities over time was reported (Trujillo et al., 2019). With the increasing popularity of 3D cultures, the field will benefit from tools designed for examining neural networks in greater detail in 3D.

Among the techniques for functional recordings and analysis, optical approaches have the advantages of contact-free, high and flexible spatial resolution, and cell-type-specific targeting (Badura et al., 2014). Here we present an in vitro experimental system that allows for interrogation of functional networks that form in a 3D biomimetic microenvironment, with graph-theory-based network analysis that had not been widely utilized by the 3D culture field. The use of a GECI driven by a neuron-specific promotior in combination with observation using widefield microscopy allows for imaging of spatiotemporal patterns in neuronal activity at the millimeter-scale (Figure 1). To describe the functional neural network characteristics and topological properties of our 3D biomimetic cortical culture, we computed several mathematical descriptors (edge weight, clustering coefficient, path length, number of modules, modularity, and node degree). Ca⁺⁺ imaging has been widely used in tissues, both in vivo and ex vivo, and in other forms of 3D cultures, including organoids. With user-optimized ROI size and Ca⁺⁺ event detection parameters according to the culture’s neuronal activities, our network analysis methods can be readily adapted to a wide range of other culture models. We compiled MATLAB algorithms into a user-friendly code and made this available to researchers (https://github.com/yutingdingle/network-analysis) so that it can be applied to other in vitro culture models. Additional network descriptors (e.g., shortcuts, core-periphery) (Lynn and Bassett, 2019) can potentially be quantified by plugging in other available open source codes and software (Hassan et al., 2015; Rubinov and Sporns, 2010).

In our 3D experimental system, dense structural networks and spontaneous neuronal activities were detected within 2 weeks of culture. In the neonatal rodent brain, AMPA receptors are initially “silent,” and glutamatergic synaptic transmission is purely driven by NMDA receptors. In contrast, activation of AMPA receptor is required for the opening of NMDA receptors in the adult brain (Ben-Ari et al., 1997). At 3 weeks of our 3D culture, a large portion of NMDA receptor activity (blocked by AP5) was independent of AMPA receptors (blocked by NBQX). NMDA receptors were also observed to drive a higher percentage of excitatory activities than AMPA receptors. The glutamatergic neurotransmission of the 3D cultures thus appears to fall within the developmental stage of immature brains at the postnatal stage. In addition, we noticed variations among the 3D cultures’ susceptibility to NMDA and AMPA receptors blockers, with one hypothesis being that differences in the expression levels of AMPA and NMDA receptors and the functionality of AMPA receptors among individual 3D samples during neural network formation is the underlying basis for this observation.

Functional networks were established by 3 weeks in the 3D cortical culture. The lack of hubs and weak community structures observed in the networks at this time point can be interpreted as an indication of an un-specialized network (Bullmore and Sporns, 2009; Lynn and Bassett, 2019; Schroeter et al., 2015). It is plausible that our 3D mouse cortical network may develop more specialized features with extended culture time and/or application of external stimuli in the form of optogenetic actuators (such as channelrhodopsins) and/or biochemical stimuli, such as brain-derived neurotrophic factor (BDNF) (Afshar Saber et al., 2018; Mishchenko et al., 2019).

A functional, balanced neural network relies on the highly regulated integration of the flow of information from both excitatory and inhibitory inputs. We demonstrated that this experimental system can model disruptions in excitatory and inhibitory neurotransmissions and network dysfunctions. Blocking excitatory AMPA receptors with NBQX reduced neuronal activity and disrupted the functional neural network with re-ductions in edge weight, clustering coefficient, and community structure. Blocking inhibitory GABA receptors with antagonists BIC and PTX increased neuronal activity (indicated by the increase of Ca⁺⁺ event frequency). Interestingly, only BIC treatment resulted in significant changes to functional network descriptors such as increased average edge weight, increased average clustering coefficient, decreased average path length, and decreased modularity, at our chosen spatial scale of analysis. In-depth examination of GABA receptor subtype expressions and distributions are needed to further address the mechanistic differences...
in network responses. However, possible explanations for these discrepancies include the differences in targets and mechanisms-of-action of BIC and PTX (i.e., competitive GABA<sub>A</sub> receptor antagonist versus non-competitive, allosteric antagonist of both GABA<sub>A</sub> and GABA<sub>P</sub>, respectively). Considering that the brain undergoes critical changes of GABA, NMDA, and AMPA receptor isoforms and functions during development in vivo, our 3D culture system may provide a useful in vitro tool to further examine how changes in receptor expressions shape functional network formation.

The inherent flexibility of our bioengineering approach allows for expression of other genetically encoded constructs to induce (e.g., optogenetic tools) and record cell and network activities (e.g., voltage sensitive and glutamate reporters) (Afshar Saber et al., 2018; Mutoh et al., 2012). Furthermore, high-speed recording of large 3D networks with cellular resolution may be possible using light-sheet confocal microscopy. Another distinct advantage of our scaffold-based bioengineering approach is the possibility of tailoring the organization, compartmentalization, and mechanical properties of the 3D culture to better match that of the native brain. For instance, specific cell-types (e.g., astrocytes, microglia, oligodendrocytes, or other neural subtypes), as well as their ratio and density, can be customized in our 3D cultures to mimic white and gray matter structure (Tang-Schomer et al., 2014), to create specific cellular interactions (e.g., neural circuits), and/or to study interactions between cells and varied ECM compositions (Sood et al., 2016). We hypothesize that systematic testing of these input parameters will advance the bioengineering of 3D in vitro biomimetic models to more closely mimic the network density and functional connectivity of the brain.

The 3D in vitro biomimetic culture approach would be particularly useful to study the many human brain disorders that are thought to be caused by dysfunction in the structure and function of synapses and neural circuits (Busche and Konnerth, 2016; Cantley et al., 2018; Centeno et al., 2018; Watanabe and Rees, 2016). For instance, we recently observed Alzheimer-disease-related changes in our long-lived 3D human, iPSC-derived neuron-astrocyte co-cultures grown in silk scaffolds (Rouleau et al., 2020). This suggests that our 3D experimental system has utility as a model for long-term studies to the onset and progression of neurodegenerative and neurodevelopmental disorders. In summary, our interdisciplinary experimental system consisting of a 3D in vitro biomimetic neural culture, genetically encoded neuronal activity reporter, widefield imaging acquisition, and graph-theory-based network analysis adds a valuable new tool for investigating development, function, and disorders of brain networks.

Limitations of the Study

The brain is organized into circuits in which diverse types of neurons connect over multiple spatial scales in a 3D space, ranging from synaptic connections to connections between brain regions. In the present study, we utilized readily accessible standard widefield microscopy to conduct millimeter-scale analysis of 3D mouse cortical cultures at the expense of resolution and the analysis in the third dimension. In comparison to 2D cultures, where all the information flow within the field of view is captured, our platform integrated the 3D information onto a projection and information flow in the Z-dimension is lost. In order to capture Ca<sup>2+</sup> transients with high spatial resolution and at multilayers at the necessary speed for single-cell and/or 3D analysis, advanced microscopy technologies such as high-speed light-sheet confocal or two-photon microscopy will be crucial. These functional neural networks change dynamically during one’s lifespan. Our functional network characterization was conducted in mouse cultures at a 3-week time point. Future studies could incorporate human iPSC-derived cultures to investigate time-dependent development of neural network functions and the biology of brain disease.

Resource Availability

Lead Contact

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Materials Availability

This study did not generate new unique reagents.

Data and Code Availability

The code generated during this study is available at https://github.com/yutingdingle/network-analysis.
METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101434.

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Conceptualization, Y.L.D., T.J.F.N., and D.L.K.; Methodology, Y.L.D., V.L., and T.J.F.N.; Software, Y.L.D., V.L., C.M., and I.G.; Formal Analysis, Y.L.D., V.L., and L.T.F.; Investigation, Y.L.D., V.L., L.T.F., and K.C.B.; Writing—Original Draft, Y.L.D. and T.J.F.N.; Writing—Review & Editing, I.G., T.J.F.N., and D.L.K; Visualization, Y.L.D.; Supervision, T.J.F.N. and D.L.K.; Funding Acquisition, D.L.K.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Functional Characterization

of Three-Dimensional Cortical Cultures

for In Vitro Modeling of Brain Networks

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Supplementary Information

Transparent Methods

Silk fibroin scaffold fabrication

A 6% (w/v) silk fibroin solution was used to prepare silk fibroin porous sponge scaffolds (pore size 425 – 500 μm) as previously described (Chwalek et al., 2015; Tang-Schomer et al., 2014). Briefly, Bombyx mori cocoons (Tajima Shoji Co., Ltd, Japan) were cut into small pieces and boiled in 0.02 M sodium carbonate (MilliporeSigma #57795) solution for 30 min to remove sericin. Dry silk fibroin fibers were dissolved in 9.3 M lithium bromide (LiBr, MilliporeSigma #213225) solution at 60°C for 4 hrs. The solution was dialyzed against water in dialysis tubing (Fisher #21-152-9, molecular weight cutoff 3,500 Da) for 2 days with six 4 L water changes over 48 hours to remove LiBr. Undissolved material was removed by centrifuging twice at 12,700 rcf at 4°C for 30 min followed by passing through a cell strainer with 100-µm pore size (Corning #431752). The resulting silk fibroin solution was adjusted to 6% (w/v) with water. Sodium chloride (NaCl) was sieved to 425 – 500 µm particle size. Thirty mL of 6% silk fibroin solution was poured into a 10-cm petri dish, and 60 g of NaCl was poured evenly into the solution. The silk fibroin was left at room temperature for 2 days and incubated at 60°C for 1 hr to allow the silk fibroin to form insoluble β-sheet crystalline structure. The resulting material was freed from the petri dish and rinsed in 4 L water for 2 days with 6 water exchanges to remove the NaCl particles and leave behind the resulting pores. Silk scaffolds for the biomimetic cortical cultures were cut into 3-mm diameter and 1.5-mm height cylinders using a biopsy punch (Integra #12-460-406) and a razor blade.

3D cortical cultures and AAV infections

All 3D cortical cultures were maintained, unless otherwise noted, in Neuro Medium: Neurobasal Medium (ThermoFisher #21103) or Neurobasal Medium minus phenol red (ThermoFisher #12348) supplemented with 2% B27 (ThermoFisher #17504), 1% Antibiotic-Antimycotic (ThermoFisher #15240), and 2 mM GlutaMax (ThermoFisher #35050).

Scaffolds were placed in water and sterilized by autoclaving. Prior to cell seeding, scaffolds were coated overnight with poly-D-lysine (PDL; 0.1 mg/mL in water, MilliporeSigma #P6407) and laminin (LN; 10 µg/mL, MilliporeSigma #MFCD00081739). Typically, in a 12-well plate we coat 20-30 sponges in 2 mL coating solution. On the day of cell seeding, the scaffolds were washed twice with PBS (10-min incubation), then incubated at least for one hour with Neuro Medium and at least for one hour in Neuro Medium containing 5% (v/v) fetal bovine serum (FBS). Excess medium was aspirated immediately before cell seeding.

Brain cortices were harvested from E16 C57BL/6 mouse embryos (Charles River Laboratories) and digested with 0.25% Trypsin (Thermo Fisher #325200) and 0.3 mg/mL DNase (MilliporeSigma #10104159001). All animal procedures were approved by Tufts University Institutional Animal Care and Use Committee (IACUC, Protocol #M2018-06). Neurons were resuspended in Neuro Medium containing 5% FBS at the density of 2×10^6 cells per 5 µL. PDL-LN-coated scaffolds aspirated to dry and transferred to a 96-well plate (one scaffold per well). A cell suspension of 5 µL was pipetted directly onto the scaffold, and the seeded scaffolds were incubated in 37°C for 30 min to allow for cell adhesion, followed by addition of 200 µL Neuro Medium + 5% FBS. The day of cell seeding was referred as 0 days-in-vitro (DIV). The next day (1 DIV), cultures were transferred to 500 µL of medium in 48-well plates. For neurite density image analysis, 3D cultures were infected on 1 DIV with AAV-hSyn1-TurboRFP (pENN.AAV.hSyn1-TurboRFP.WPRE.RBG, Addgene #105552-AAV1, titer 2.7 x 10^{13} GC/mL, a gift from Dr. James M. Wilson, unpublished) diluted in Neuro Medium at 1:1000. 3D cultures for DNA quantification, RNA analysis, and immunostaining were transferred to fresh 500 µL Neuro Medium without AAV. For Ca^{2+} imaging experiments, 3D cultures were infected with AAV-hSyn1-GCaMP6f (AAV-hSyn1-GCaMP6f-P2A-nls-dTomato, Addgene, #51085-AAV1, titer 2 x 10^{13} GC/mL, a gift from Dr. Jonathan Ting, unpublished) diluted in Neuro Medium at 1:1000. The 3D cultures where then incubated overnight at 37°C.
On 2 DIV, 3D cultures were transferred to 96-well plates, and 10 µL of 3 mg/mL collagen type I solution (Corning #354236) was pipetted onto each scaffold followed by incubation at 37°C for 30 min to allow for collagen gelation. Each 3D culture was then transferred to a 24-well containing 2 mL Neuro Medium. For cultures intended for Ca++ imaging (AAV-hSyn1-GCaMP6f infected), glass-bottom plates (Cellvis #P24-1.5P) and phenol red-free Neuro Medium were used. Half-media changes were carried out every 3-4 days.

Assessment of 3D neurite density

3D cultures infected with AAV-hSyn1-TurboRFP were chemically fixed on at 1, 2, and 3 wk with 4% paraformaldehyde/4% sucrose in PBS for 1 hr. After three PBS washes (30-min each), the 3D cultures were imaged with a Leica SP8 confocal microscope with a 40X objective (Leica HC PL APO 40X/1.10 W CORR CS2), x,y,z-dimensions= 291 x 291 x 210 µm, 0.42 µm per z-step) with excitation laser of 552 µm. 3D movie (Movie S1) was rendered with maximum projections in LAS X software (Leica). Z-stack images were analyzed using a custom MATLAB (MathWorks) script (Liaudanskaya et al., 2020). First, Otsu’s thresholding was used to remove weak background fluorescence. Then, connected objects were identified using the bwlabel MATLAB function, and objects with eccentricity lower than 0.9 were removed, as they represented signals emanating from silk and cell bodies. The remaining voxels were grouped together into objects if neighboring edges or faces were in contact using the 3D bwlabel MATLAB function. The smallest ellipsoid was drawn around each object, and the major axis length of each individual ellipsoid was measured in voxel increments. The sum of all such voxels was calculated for each z-stack image to represent the volume occupied by neurites. The neurite density was presented as the percent of neurite voxel number to total voxel number per z stack.

Immunofluorescence analysis of 3D cultures

After chemical fixation of 3D cultures, the cells were permeabilized and blocked for 1 hr in blocking solution (4% (v/v) normal goat serum (Jackson Immuno Research #005-000-121) and 0.2% (v/v) Triton-X (MilliporeSigma #T8787) in PBS). Rabbit anti-GFAP primary antibody (1:2000, MilliporeSigma #G9269) and Alexa 488 goat-anti-rabbit secondary antibody (1:1000, ThermoFisher #A11034) were diluted in blocking buffer. Primary antibody incubation was done at 4°C overnight, followed by three 30-min PBS washes. Secondary antibody incubation was done in blocking solution at room temperature for 2 hrs, followed by DAPI staining (1:1000 in 0.2% Triton-X, Thermo Fisher #D1306) and three 30-min PBS washes. Confocal z stack images were acquired using a Leica SP8 confocal microscope with a 25X objective (Leica HC FLUOTAR L 25x/0.95 W VISIR) and with excitation lasers at 405 and 488 µm.

Gene expression analysis of 3D cultures

On 1, 14, 21 DIV, 3D culture samples were collected in 600 - 700 µL ice-cold Buffer RLT (Qiagen #79216) with 1% β-mercaptoethanol (MilliporeSigma #M3148) and stored in -80°C until RNA isolation. On the day of mRNA isolation, samples were thawed and homogenized, first by manually grinding into fine pieces with a homogenizer pestle (Fisher #12-141-364), followed by homogenization for 20 sec with a motorized tissue grinder (Fisher #12-1413-61), and then passed through a QIAshredder column (Qiagen #79656). RNA was isolated using the Qiagen RNeasy Mini Kit (Qiagen #79656) according to the manufacturer’s protocol, with additional in-column removal of genomic DNA with DNase (Qiagen # 79254). The resulting RNA was eluted with 25-30 µL of water and the resulting concentrations measured using a NanoDrop Spectrophotometer (ThermoFisher). cDNA was synthesized using iScript gDNA Clear cDNA Synthesis Kit (Bio-Rad #1725035) according to the manufacturer’s protocol.

Relative gene expressions were quantified using the Taqman quantitative real-time PCR (qRT-PCR) assay and analyzed with the QuantStudio 5 RT-PCR System (Thermo Fisher). Each qRT-PCR reaction contained
10 ng cDNA, 10 µL Taqman Gene Expression Master Mix (ThermoFisher #4369016), and 1 µL Taqman assay. See Table 2 for the list of Taqman assays. All expressions were referenced to housekeeping gene Rn18s and normalized to 1 DIV. ΔΔCt method was used to calculated relative expression levels.

DNA Quantification

Relative DNA amount was quantified using Quant-iT PicoGreen dsDNA Assay Kit (ThermoFisher #P7589). Cortical cell suspensions (1 and 2 x 10⁶ cells) were centrifuged at 500 rpm for 5 min. Supernatant was removed, and cell pellets were lysed with 700 µL Buffer RLT with 1% β-mercaptoethanol and vortex. 3D cultures were lysed as described in the Gene expression analysis of 3D cultures section above. 5 µL samples or DNA standards were diluted with 95 µL Buffer RLT with 1% β-mercaptoethanol. Assays were conducted according to the manufacturer’s protocol. Raw fluorescence values were used to estimate number of cells in the 3D cultures.

Ca²⁺ imaging

A half-medium change was done 1 or 2 days prior to the day of imaging. Ca²⁺ imaging was performed on AAV-hSyn1-GCaMP6f infected 3D cultures with a Nikon Eclipse Ti-2 inverted fluorescence microscope with a 4X objective (Nikon CFI Plan Fluor DL 4x na 0.13 wd 16.5mm Objective). During imaging, plates were kept at 37°C and 5% CO₂ in a stage-top incubator (Bioscience Tools #TC-MWP). Medium was removed to keep the 3D cultures from floating during image acquisition, leaving approximately 500 µL of medium to support the culture during imaging. The 3D culture was centered in the field of view and all time-lapse images were acquired with a FITC filter and the following settings: 3x3 binning, 50-ms exposure, 200-ms intervals (i.e. 5 Hz), and 1-min duration. At 2 weeks (14 DIV) only spontaneous baseline Ca²⁺ activities were recorded. At 3 weeks (20-22 DIV), a 1-minute baseline image analysis was first performed. Picrotoxin (50 µM, Abcam #ab120315) and bicuculline (10 µM, Abcam #ab120107) were then applied for 7 min to the sample, after which recording was repeated for another minute. NBQX (5 µM Tocris 1044; and AP5 (50 µM, Sigma A8054) were applied for 22 min before the second recording. Picrotoxin and bicuculline were first dissolved in DMSO to make 2000X stock solutions. A 5 µL of a 100X concentrated solution, diluted in Neuro Medium, of each of the drugs was added to the cultures. Equivalent concentration of DMSO (1:2000) was added to a separate set of 3D to control for the addition of DMSO in picrotoxin and bicuculline. The time-lapse images were converted to TIFF stacks in NIS-Elements software (Nikon).

Neuronal activity and global synchronization analysis of 3D neural networks

Region of interest (ROI) assignment

A mask in the shape of a honeycomb pattern of 37 hexagonal ROIs (side-to-side distance = 500 µm, center-to-center distance = 500 µm) was first designed on Adobe Illustrator to cover the circular projection of the 3D scaffold and converted to binary in Adobe Photoshop or NIS-Elements (Nikon, Micro Video Instruments, Inc). Each ROI was defined as a node for all of the following analyses.

Ca²⁺ events detection, % active ROI, and Ca²⁺ event frequency calculation

Ca²⁺ event detection and event frequency calculation were conducted using an open source MATLAB-based software FluoroSNNAP (Patel et al., 2015). The time-lapse TIFF stack and ROI mask were loaded in FluoroSNNAP, and the Segmentation GUI was used to create a segmentation MATLAB. The Single-Cell Ca²⁺ Transient Kinetics module was used for Ca²⁺ event detection and to calculate total events for each individual ROI in a 3D tissue sample. The input parameters used for baseline fluorescence were F₀ = 10.
The input parameters for Ca\textsuperscript{++} event detection were template-based, threshold = 0.85, and minimum dF/F amplitude = 0.01. Descriptions of input parameters are detailed in (Patel et al., 2015). Samples with poor cell seeding efficiency (< 50% active ROI at baseline) were excluded.

% Active ROI and Ca\textsuperscript{++} event frequency of each 3D tissue was calculated as:

\[
\% \text{Active ROI} = \% \frac{\text{Number of ROIs containing at least one Ca}^{++}\text{event}}{N} \quad \text{Eq. 1}
\]

\[
\text{Ca}^{++}\text{event frequency}_{\text{Single ROI}} = \frac{\text{Number of events}}{1 \text{ min}} \quad \text{Eq. 2}
\]

\[
\text{Ca}^{++}\text{event frequency}_{\text{Tissue}} = \frac{1}{N} \sum \text{Ca}^{++}\text{event frequency}_{\text{Single ROI}} \quad \text{Eq. 3}
\]

where \(N\) is the total number of nodes, which is 37 in this study.

**Global synchronization Index**

Global synchronization index was calculated using FluoroSNNAP’s *Synchronization* module with input parameters of method = Phase, #times to perform surrogate resampling = 20, minimum size of synchronization cluster = 3. The mathematical descriptions are detailed in (Patel et al., 2015). Briefly, this method separates the time-varying fluorescence signals into its amplitude and phase components and compares the similarity of timing of events (phase) of all the ROIs.

**Functional connectivity and graph theory-based network analysis of 3D neural networks**

**Cross-correlation analysis**

Cross-correlation analyses were done using FluoroSNNAP’s *Estimate Functional Connectivity* module, with number of resampling = 100 and significance level = 0.5. The raw cross-correlation coefficient matrices of all node (ROI) pairs (i, j, whereas i≠j) were accessed in the output MATLAB file (located in the file ‘processed analysis.mat/FC/C’), extracted, and saved as a numeric matrix MATLAB file. This file is then loaded into a single, user-friendly, MATLAB code, which is a compilation of the individual codes used to compute the graph theory-based network parameters.

**Average edge weight calculation**

Network analyses in our study were based on a weighted network (i.e. edges values, not binary). The cross-correlation coefficients between paired nodes i and j were used as edge weights in the graph. The average edge weight for each 3D culture sample was calculated by averaging all the edge weights.

**Average clustering coefficient and path length**

The local clustering coefficient for each node \(c_i\), in a weighted graph was computed as described by Onnela et al, using their published MATLAB script (Onnela et al., 2005). Briefly, clustering coefficient in a weighted graph reflects the intensity of triangle networks formed by triplets of nodes is calculated as follows:
\[ c_i = \frac{1}{k_i(k_i-1)} \sum_{j,k} \frac{3(w_{ij}w_{jk}w_{ki})}{\text{max}(w)} \]  \hspace{1cm} \text{Eq. 4}

where \( k_i \) is the number edges connected to node \( j \) (36 in our case) and \( w \) is the edge weight. The average clustering coefficient for a network \( C \) is the average of all \( c_i \):

\[ C = \frac{1}{N} \sum c_i \]  \hspace{1cm} \text{Eq. 5}

where \( N \) is the total number of nodes.

**Average path length**

Path length for the 3D culture network was computed as described by Muldoon et al., 2016 using their published MATLAB script (Muldoon et al., 2016). Briefly, the shortest distance \( d \) between node \( i \) to node \( j \) is inversely proportional to edge weight \( (w) \) and is defined as:

\[ d_{ij} = \frac{1}{w_{ij}} \]  \hspace{1cm} \text{Eq 6.}

The average path length (PL) for a network \( PL_N \), with number of nodes \( N \) is calculated as:

\[ PL_{i,j} = \frac{1}{N(N-1)} \sum_{i \neq j} d_{ij} \]  \hspace{1cm} \text{Eq 7.}

**Number of modules and modularity**

Module assignment of nodes and modularity were computed as described in Newman, 2006 using their published MATLAB script (Newman, 2006). Briefly, modularity is the sum of edges falling into a module minus the sum of expected edges in an equivalent random network. Module assignment is based on maximizing modularity of the network.

**Node degree and determination of hubs**

To calculate the node degree for each node, the sum of edge weights incident to the node was calculated. The frequency distribution (%) of individual weight node degree in all samples under the same condition was calculated with bin size = 0.05 using GraphPad Prism.

\[ \text{Weighted node degree}_i = \frac{1}{A} \sum_{i \neq j} w_{ij} \]  \hspace{1cm} \text{Eq. 8}

To test if the frequency distribution of weighted node degree followed a normal distribution, a Kolmogorov-Smirnov normality test was performed.
Statistical analysis

Statistical tests were performed using GraphPad Prism. Statistical significance in changes in neurite density (Fig. 2C) and mRNA expression (Fig. 2D and 3A) were tested with ANOVA with post-hoc Tukey tests. For changes in neuronal activity (Fig. 4E, F) between different cultures at 2 and 3 weeks, unpaired t-tests were used. Differences in neuronal activity (Fig. 5B-D) and network properties (Fig. 6C-F) between baseline and post-treatments of each individual 3D tissue were determined with paired t-tests. A p-value of 0.05 was used for significance level.
Supplementary Figures

Figure S1. Schematic of the 3D *in vitro* cortical culture set up for cellular and network activity recording. Related to Figure 1.
Figure S2. Relative cell numbers in 3D culture. DNA quantity in cortical cell pellets (1 and 2 million) and in 3D cultures were analyzed with PicoGreen DNA Quantification Kit. n = 3-5 samples. Related to Figure 1.
Figure S3. Example traces of Ca** transients from three independent 3D *in vitro* biomimetic cortical cultures at (A) 2 weeks and (B) 3 week at baseline and post-drug treatments (3 wks only) of (C) bicuculline (BCC, 10 µM), (D) picrotoxin (PTX, 50 µM) (E) AP5 (50 µM), and (F) NBQX (5 µM). Each trace represents the average fluorescence intensity over time normalized to its basal fluorescence level (dF/F₀) from an individual hexagonal ROI (side-to-side distance 500 µm). Related to Figures 4-5.
Figure S4. Control experiments with DMSO treatment, at the equivalent concentration (0.5%) as for BCC and PTX treatments, did not induce changes in (A) Ca\(^{++}\) event frequency, (B) global synchronization index, (C) average edge weight, (D) modularity, (E) average clustering coefficient, and (F) average path length. Grey data points and lines indicate baseline and post-treatment measurements of the same 3D tissue culture samples, and color lines show Mean ± SEM. n = 7 from two independent experiments. Related to Figure 5.
Figure S5. Network analysis with 500 µm vs 250 µm ROIs. (A) Time-lapse image stacks were applied with 127 hexagonal ROIs with side-to-side distance of 250 µm. (B) Representative cross-correlation matrix of neuronal activities in each ROI at baseline condition. (C) Network descriptors of global synchronization index, average edge weight, average clustering coefficient, average path length, and modularity, calculated based on using 500 µm or 250 µm ROI mask. Data are individual values and Mean ± SD (n = 12 samples each group). Paired t-tests were used for statistical significance comparing the same condition with 500 µm or 250 µm ROI and for comparing baseline and post-treatment of the same sample with same ROI size. *p < 0.05. Related to Figure 6.
| Drug     | Bicuculline | Picrotoxin | NBQX   | AP5   |
|----------|-------------|------------|--------|-------|
| Concentration | 10 µM      | 50 µM      | 5 µM   | 50 µM |
| Target and mechanism of action | Competitive GABA<sub>A</sub> receptor antagonist | Non-competitive GABA<sub>A</sub> and GABA<sub>P</sub> receptors antagonist | Competitive AMPA receptor antagonist | Competitive NMDA receptor antagonist |
| Sample Number | 12          | 12         | 9      | 11    |
| Condition | Baseline | Post-Treatment | Baseline | Post-Treatment | Baseline | Post-Treatment |
| % ROI | Min | 94.59 | 100 | 89.18 | 100 | 94.59 | 16.22 | 89.19 | 0 |
|         | Median | 100 | 100 | 100 | 100 | 98.65 | 48.65 | 100 | 32.43 |
|         | Max | 100 | 100 | 100 | 100 | 100 | 100 | 100 | 100 |
| Unpaired t-test | p = 0.3388 | p = 0.3388 | p = 0.0068 | p < 0.0001 |
| Ca<sup>2+</sup> Event Frequency (events/min) (Mean ± SEM) | 10.88 ± 1.10 | 15.41 ± 0.72 | 12.30 ± 1.10 | 17.80 ± 1.08 | 10.47 ± 1.05 | 2.68 ± 0.94 | 11.48 ± 1.61 | 0.35 ± 0.12 |
| paired t-test | p = 0.0002 | p = 0.0013 | p < 0.0001 | p < 0.0001 |
| Global Synchronization Index (Mean ± SEM) | 0.269 ± 0.031 | 0.256 ± 0.037 | 0.312 ± 0.038 | 0.293 ± 0.045 | 0.268 ± 0.027 | 0.142 ± 0.035 | n.a. |
| paired t-test | p = 0.4105 | p = 0.6981 | p = 0.0273 | |
| Average Edge Weight (Mean ± SEM) | 0.360 ± 0.018 | 0.414 ± 0.021 | 0.402 ± 0.023 | 0.433 ± 0.025 | 0.394 ± 0.032 | 0.118 ± 0.051 | n.a. |
| paired t-test | p = 0.0134 | p = 0.147 | p = 0.009 | |
| Average Clustering Coefficient (Mean ± SEM) | 0.331 ± 0.025 | 0.419 ± 0.023 | 0.380 ± 0.024 | 0.433 ± 0.025 | 0.358 ± 0.036 | 0.117 ± 0.043 | n.a. |
| paired t-test | p = 0.0057 | p = 0.0711 | p = 0.0005 | |
| Average Path Length (Mean ± SEM) | 3.01 ± 0.168 | 2.56 ± 0.13 | 2.49 ± 0.09 | 2.37 ± 0.08 | n.a. | n.a. |
| paired t-test | p = 0.0331 | p = 0.2740 | n.a. | |
| n. of Modules (n. of samples) | with 3 modules | 4 | 3 | 3 | 2 | 2 | 0 |
|         | with 2 modules | 8 | 9 | 9 | 10 | 7 | 6 | n.a. |
|         | with 1 module | 0 | 0 | 0 | 0 | 0 | 3 | |
| Modularity (Mean ± SEM) | 0.102 ± 0.012 | 0.067 ± 0.006 | 0.090 ± 0.010 | 0.070 ± 0.011 | 0.083 ± 0.018 | 0.083 ± 0.184 (N. Mod > 1) | n.a. |
| paired t-test | p = 0.0173 | p = 0.2043 | p = 0.0202 (N. Module >1) | |

**Table S1.** Results summary of neuronal activity measurements and functional network descriptors of the 3D *in vitro* biomimetic cultures at baseline and post-bicuculline, picrotoxin, NBQX, and AP5 treatments at 3 weeks. Related to Figures 4-6.
| Gene   | Encoded Protein                                         | ThermoFisher Taqman Assay ID     |
|--------|--------------------------------------------------------|----------------------------------|
| *Dcx*  | Doublecortin                                           | Mm00438400_m1                    |
| *Gabra1* | Gamma-aminobutyric acid type A receptor alpha1 subunit | Mm00439046_m1                    |
| *Gad2* | Glutamate Decarboxylase 2                              | Mm00484623_m1                    |
| *Gria1* | Glutamate Ionotropic Receptor AMPA Type Subunit 1      | Mm00433753_m1                    |
| *Grin1* | Glutamate Ionotropic Receptor NMDA Type Subunit 1      | Mm00433790_m1                    |
| *Pax6* | Paired Box Protein Pax-6                               | Mm00443081_m1                    |
| *Rn18s* | 18s Ribosomal RNA, 45s Ribosomal RNA                    | Mm03928990_g1                    |
| *Shank3* | SH3/ankyrin domain gene 3                              | Mm00498775_m1                    |
| *Slc17a7* | Vesicular glutamate transporter 1                       | Mm00812886_m1                    |
| *Slc1a3* | Excitatory amino acid transporter 1                     | Mm00600697_m1                    |
| *Syn1*  | Synapsin I                                             | Mm00449772_m1                    |

**Table S2.** Taqman assays used in qRT-PCR in this study. Related to Figure 2.
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