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CRISPRi-based screens in iAssembloids to elucidate neuron-glia interactions

by
Emmy Li

DISSERTATION

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DOCTOR OF PHILOSOPHY

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in the

GRADUATE DIVISION

of the

UNIVERSITY OF CALIFORNIA, SAN FRANCISCO

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By

Emmy Li

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Contributions to Presented Work

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CRISPRi-based screens in iAssembloids to elucidate neuron-glia interactions

Emmy Li

Abstract

The sheer complexity of the brain has complicated our ability to understand its cellular mechanisms in health and disease. Genome-wide association studies have uncovered genetic variants associated with specific neurological phenotypes and diseases. In addition, single-cell transcriptomics have provided molecular descriptions of specific brain cell types and the changes they undergo during disease. Although these approaches provide a giant leap forward towards understanding how genetic variation can lead to functional changes in the brain, they do not establish molecular mechanisms. To address this need, we developed a 3D co-culture system termed iAssembloids (induced multi-lineage assembloids) that enables the rapid generation of homogenous neuron-glia spheroids. We characterize these iAssembloids with immunohistochemistry and single-cell transcriptomics and combine them with large-scale CRISPRi-based screens. In our first application, we ask how glial and neuronal cells interact to control neuronal death and survival. Our CRISPRi-based screens identified that $GSK3\beta$ inhibits the protective NRF2-mediated oxidative stress response in the presence of reactive oxygen species elicited by high neuronal activity, which was not previously found in 2D monoculture neuron screens. We also apply the platform to investigate the role of APOE- $\epsilon 4$, a risk variant for Alzheimer's Disease, in its effect on neuronal survival. This platform expands the toolbox for the unbiased identification of mechanisms of cell-cell interactions in brain health and disease.

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Chapter 1 : Introduction

The human brain contains over 86 billion neurons and approximately the same number of non-neuronal cells¹, which are all intricately interconnected in the brain. One of the most challenging problems in biology is to disentangle the complexity of the brain and to understand how brain function is altered in disease. Genetic and genomics approaches are uncovering how genetic variation and mutations can drive cellular and molecular changes in the brain. For example, single-cell transcriptomics have revealed the diversity of cell types and cell states in neuronal and glial cells at the molecular level², while genome-wide association studies (GWAS) have identified genetic variants associated with specific neurological phenotypes and diseases³⁻⁵. However, these approaches do not elucidate the molecular mechanisms, whether cell autonomous or cell-non autonomous, by which genes control brain function and disease.

To enable the scalable functional characterization of gene function in brain cell types, we previously developed CRISPR-based functional genomics screens in human iPSC-derived neurons^{6,7}, microglia⁸, and astrocytes⁹. However, these screens were conducted in monoculture systems and are thus unable to capture mechanisms driven by glia-neuron interactions. Multiple systems have been developed in hiPSC organoid-based culture systems to link specific genetic perturbations to functions¹⁰⁻¹². A limitation of organoid-based models is the lengthy and labor-intensive process to generate them, and remaining challenges in heterogeneity. To address these challenges, we developed a 3D co-culture system we call iAssembloids (induced multi-lineage assembloids) that enables the rapid generation of homogenous neuron-glia spheroids. iAssembloids

enable specific control over the origin of cell types of interest, for instance, to combine cells from different genetic backgrounds, enabling the investigation of non-cell autonomous mechanisms. This allows for the investigation of cell-type specific roles of disease-associated genetic variants.

Here, we characterize these iAssembloids and combine them with large-scale CRISPRi-based screens to model how glial and neuronal cells can interact to modulate neuronal survival. We also apply the platform to investigate the role of APOE- ϵ 4, a major risk variant for Alzheimer's Disease, in its effect on neuronal survival. Our platform aims to expand our existing functional genomics toolbox to enable a simple, highly scalable method for screening genetic modifiers for neuronal survival in a glial context. Thus, iAssembloids in combination with CRISPRi-based functional genomics will enable the unbiased dissection of mechanisms underlying neuron-glia interactions in health and disease.

Chapter 2 : Development of 3D neuron-astrocyte-microglia co-culture system (iAssembloids)

Neurons, astrocytes, and microglia integrate into three dimensional cultures (iAssembloids)

To generate a scalable three-dimensional neuron-astrocyte-microglia co-culture system that can be used for CRISPRi-based genetic screening approaches, we selected differentiation protocols for hiPSC-derived neurons, astrocytes and microglia that are scalable and rapid. We used the dox-inducible NGN2 system for neuron differentiation^{13,14} and our recently developed six transcription factor (6TF) induction protocol for microglia differentiation⁸. For astrocyte differentiation, we used either of two different protocols, one that is serum-based¹⁵ and one that is serum-free¹⁶. For iAssembloid generation, we seeded differentiated astrocytes together with pre-differentiated i3Neurons (day 0) at a ratio of 3 neurons to 1 astrocyte into AggreWell™ 800 plates. After one week, we added microglia at a third of the number of astrocytes (**Figure 1**). These cultures self-assemble into spheroids within the wells, are uniform in size (200-400 microns; ~10,000 cells) and retain their shape over time (**Figure 1**).

To visualize the three cell types and their interactions, we seeded iAssembloids (induced multi-lineage assembloids) with hiPSC-derived neurons expressing BFP, astrocytes expressing GFP and microglia expressing mScarlet along with cells expressing no fluorescent proteins. Neurons, astrocytes, and microglia were imaged in the iAssembloids at 14 days post-culture in the AggreWell™ 800 plate. Astrocytes developed a stellate morphology (**Figure 1**). Neurons developed processes that traversed the iAssembloids

(Figure 1). To confirm that all three cell types maintained their cellular identity in iAssembloids throughout differentiation, we stained the iAssembloids for neuronal and glial markers. Neurons within the culture expressed neuronal cell markers NEUN and TUJ1 **(Figure 1)**. Microglia, stained with IBA1, fell into two categories: some were amoeboid and migrated towards the middle of the iAssembloid; the others (denoted by arrows) sent projections through the iAssembloid **(Figure 1)**. Astrocytes stained positively for S100 β **(Figure 1)**. Taken together, these data show that the three cell types have integrated into the iAssembloids and develop morphologies characteristic of more mature neurons and glia than those grown in monoculture.

Neurons cultured in iAssembloids are more active than neurons in 2D monoculture

We performed single-nucleus RNA sequencing to further characterize each cell type in the iAssembloids. Cells within iAssembloids expressed appropriate cell type specific markers (neurons: RBFOX3, MAPT; astrocytes: SLC1A3; microglia: SPI1, CSF1R) and maintained their identity throughout differentiation **(Figure 2.1)**. We then compared gene expression profiles of neurons cultured within iAssembloids to monocultured neurons that were harvested at the same time point (day 14 post-differentiation)¹⁷. Using EnrichR, we found genes related to axon guidance and synaptic transmission were more highly expressed in neurons cultured in iAssembloids compared to monoculture **(Figure 2.1)**. More specifically, we found that glutamate receptor subunits such as GRIA1 and GRIA2 as well as ion-channel subunits (CACNA1C, KCNQ1, KCNIP4) were more highly expressed in neurons cultured in iAssembloids **(Figure 2.1)**. Therefore, we hypothesized that neurons cultured in iAssembloids may have higher levels of neuronal activity than

neurons in monoculture. We therefore assessed the neurons' electrophysiological activity by performing multi-electrode array analysis in the two culture conditions (**Figure 2.1**). After two weeks in culture, we found that neurons in iAssembloids had tenfold greater number of spikes on average (**Figure 2.1**). Taken together, these results suggest that neurons cultured in iAssembloids are more electrophysiologically active than those cultured in monoculture.

Astrocytes in iAssembloids express genes important for supporting neuronal function

We found that astrocytes cultured in iAssembloids developed transcriptional profiles that did not exist in 2D monocultured astrocytes in used in previous CRISPRi-based screens⁹ (**Figure 2.2**). The clusters that overlapped between the two conditions were either in TOP2A+ proliferative clusters or clusters without definitive markers. As astrocytes within iAssembloids matured, astrocytes had transcriptional profiles more strongly associated with supporting neuronal function based on GO annotation of the upregulated genes. These functions include BDNF signaling (JUN, BTG1, PEG10, TUBB), axon guidance (ROBO2, NRXN1, NCAM1), as well as synaptic transmission (APP, GRID2, NRG3) (**Figure 2.2**). Single-nucleus RNA sequencing also revealed that some astrocytes expressed higher levels of genes associated with glycolysis (PFKFB4, PDK1, ENO1) as well as the unfolded protein response (HSPA5, DDIT3, HERPUD1). None of the iAssembloid-derived astrocytes expressed markers of reactive astrocytes induced by the microglial cytokines IL1a+TNF+C1q stimulation such as IL6, IL1B, VCAM1, CXCL2, or CXCL10^{9,18}, suggesting that astrocytes in iAssembloids do not develop the same transcriptional profile as astrocytes activated by the three cytokines. These results

suggest that astrocytes develop homeostatic signatures in iAssembloids that are not represented in monoculture.

Microglia populations unique to iAssembloids express major histocompatibility complex genes

To characterize transcriptional changes that occur in microglia cultured in iAssembloids, we compared single-nucleus RNA-sequencing data for microglia in iAssembloids to those for monocultured iPSC-derived microglia⁸ (**Figure 2.3**). We found that 95% of the microglia in day 14 iAssembloids (iMicroglia age 13 days, 5 days in iAssembloid culture) fell into clusters also seen in monocultured microglia. By 28 days in iAssembloid culture, 79% of microglia fell into unique clusters. These clusters, which we named “iAssembloid SPP1+,” “iAssembloid chemokine,” and “iAssembloid proliferative,” have profiles like the monocultured microglia SPP1+, chemokine, and proliferative clusters, but are characterized by expression of MHC class II markers such as HLA-DRA and HLA-DRB1 (**Figure 2.3**), which were either not expressed or not captured by microglia in the monoculture dataset. Components of MHC class II pathway have been shown to be upregulated in microglia present in neurodegenerative disease models^{19,20}. Therefore, iAssembloids provide a model system to study this disease-relevant state of human microglia.

Chapter 3 : CRISPRi-based screens identify iAssembloid-specific factors in neuronal survival

As neuronal death is a key hallmark of neurodegenerative disease, we wanted to probe which pathways can make neurons vulnerable and which can protect neurons from neuronal death. While we previously conducted CRISPRi-based modifier screens for neuronal survival in 2D monocultured neurons^{6,7}, the more physiological context of our iAssembloid model motivated us to conduct similar screens in iAssembloids. To perform the screen, hiPSCs expressing the dox-inducible transcription factor, NGN2, and dCas9-KRAB CRISPRi machinery were transduced with 19,000 sgRNA targeting over 2,900 genes, including kinases and druggable targets (H1 library)²¹, as well as neurodegeneration-related genes from GWAS (neurodegeneration library). To determine genes essential for neuronal survival, we induced neuronal differentiation, incorporated these cells into iAssembloids and compared sgRNA frequencies in the starting population (1 day in culture) and later timepoints (14 and 28 days in culture) to uncover genes controlling neuronal survival (**Figure 3.1**). Many of the hits fell into categories related to metabolism, neuronal activity, cell adhesion, lipid and lipoprotein function, immune response, stress response, and endocytosis (**Figure 3.1**). We confirmed that our screen was highly reproducible across two different time points (**Figure 3.2**) and in iAssembloids containing astrocytes generated from two different protocols (**Figure 3.3**).

We compared our findings to those of screens previously performed in neuron monocultures in our lab^{6,7}. The screen using the H1 library were compared to Tian et al., 2019, while the screen using the neurodegeneration library was compared to the genome-

wide screen from Tian et al., 2021. Pearson correlation of gene scores from survival screens of 2D neurons versus iAssembloids show that iAssembloid screens conducted at two different timepoints were not as highly correlated with the original 2D neuron screen (Fig. S3C). Essential genes for neuronal survival identified in monoculture, such as HMGCR, HMGCS1 and ATP6V1C1 were also found in our iAssembloid screens (**Figure 3.1**). Out of these hits, we wanted to select genes of interest that were iAssembloid-specific and had the same phenotype at day 14 and day 28 in the iAssembloids (**Figure 3.1**). These hits represent knockdowns that yielded highly reproducible phenotypes whose survival may be uniquely affected by the presence of glia and higher neuronal activity in the iAssembloids. In total, we curated a list of 68 genes that fit the criteria for further validation.

Focused validation screens assess effect of cell type and media composition on neuronal survival in iAssembloids

We conducted a focused validation screen targeting the 68 curated genes from the primary screen. Comparing the focused validation screen conducted with the same conditions as the primary screen, we found that the results were highly correlated ($r=0.9$) (**Figure 3.3**). As media composition is altered when microglia are seeded, we assessed whether changing media composition affected neuronal survival in the iAssembloids. In addition, we tested BrainPhysTM media as it has been shown to increase neural electrical activity in culture²². Through hierarchical clustering, we found that screens conducted with differing media compositions were more closely related to each other, except for the screen conducted in BrainPhysTM media. We also compared iAssembloids with neurons

co-cultured with microglia and 3D neuronal monocultures and found that with 3D neuron monoculture clustered more closely to the screen conducted in BrainPhys™ media (Fig. S4B, Table S4). To understand the downstream effects on neuron function after perturbation of these genes, we selected hits that maintained a phenotypic difference between monoculture and iAssembloid culture for a focused screen with single-cell transcriptomic readout.

CROP-seq (CRISPR droplet sequencing) suggests that GSK3B (Glycogen Synthase Kinase 3 Beta) influences neuronal NRF2 activity in the iAssembloid system

To assess transcriptomic changes resulting from genetic perturbations and to uncover pathways that are differentially regulated between iAssembloid and monocultures, we performed CROP-seq (CRISPR droplet sequencing)²³. After transduction with a pooled sgRNA library and neuronal differentiation, we enriched the neuronal population using magnetic activated cell sorting (MACS) and then we performed single cell sequencing on the enriched neuronal population (**Figure 4**) from iAssembloids and 2D monoculture.

We found that knockdown of several genes, including GSK3B, TTN, LMAN1 and players in the JNK MAPK pathway drove up expression of different targets of the transcription factors NRF2 (Nuclear factor erythroid 2-related factor 2) and STAT3 (**Figure 4**). We were interested in the relationship between GSK3 β and NRF2. GSK3 β is hypothesized to be highly relevant in a variety of neurodegenerative diseases^{24,25}. GSK3 β is a kinase that has been proposed to regulate many different cellular functions, ranging from cellular differentiation to apoptosis²⁶. In Alzheimer's disease, especially, GSK3 β has been shown to phosphorylate the protein Tau^{27,28}. Phosphorylation of Tau has been suggested to drive

Tau aggregation, leading to neuronal death and the progression of disease²⁹ while NRF2 regulates genes in response to oxidative stress³⁰. When comparing the effect of GSK3B knockdown on neurons in iAssembloids (**Figure 4**) and in 2D monoculture (**Figure 4**), we found differential regulation of NRF2 targets such as some metallothioneins^{31,32} such as MT2A, MT1E, and other genes identified through ChEA Transcription Factor Targets³³ for NRF2 such as RORB, HS3ST5, and MEIS1 only in iAssembloids. We therefore concluded that GSK3 β may influence NRF2 activity selectively in the iAssembloid system.

Chapter 4 : Knockdown of GSK3B protects neurons from oxidative stress related death

Knockdown of GSK3B promotes NRF2 activity by promoting NRF2 nuclear localization

To investigate the mechanism underlying the effect of GSK3B knockdown on NRF2, we first confirmed that GSK3 β protein levels were reduced by the sgRNAs used in the CROP-seq screen in iAssembloids (Fig. 5A). We next validated the differential effect of GSK3B knockdown on neuronal survival in different contexts. We co-cultured cells expressing non-targeting control sgRNAs (GFP+) with cells expressing sgRNAs targeting GSK3B (BFP+) at equal ratios and examined changes in those ratios at pre-differentiation and 14 days post-culture. Strikingly, neuronal survival in iAssembloids and in 3D neuronal culture is greatly improved after GSK3B knockdown (**Figure 5.1**), whereas there was no effect of GSK3B KD in 2D-monocultured neurons. Thus, the beneficial effect of GSK3B knockdown was dependent on 3D culture, but not on the presence of glia.

Upon induction of oxidative stress, NRF2 translocates to the nucleus and activates transcription of genes protective against oxidative stress^{30,34,35}. To characterize how NRF2 activity may be modified by GSK3B knockdown, we sectioned iAssembloids into 20-micron sections and determined the levels of nuclear NRF2. We found that knockdown of GSK3B resulted in higher levels of NRF2 in the nucleus (**Figure 5.1**), consistent with the induction of its target genes (**Figure 5.1**). We then asked whether the knockdown of GSK3B itself increased oxidative stress, resulting in the translocation of NRF2 to the nucleus. To test this, we assessed reactive oxygen species (ROS) levels by using the dye, CellROX™ Orange. Knockdown of GSK3B did not influence the level of ROS in

neurons in 2D or 3D culture (**Figure 5.1**). This suggests that knockdown of GSK3B does not drive oxidative stress and that the effect of GSK3B knockdown is downstream of oxidative stress.

GSK3 β activity has been reported to be activated in response to ROS to inhibit NRF2 by preventing its translocation to the nucleus in a model for acute kidney injury³⁶. We hypothesized that GSK3 β activity prevented neurons from mounting an appropriate response to oxidative stress using a similar mechanism. To test this hypothesis, we first assessed ROS levels in 2D as well as 3D neurons and found that ROS levels in 3D neurons were consistently higher than in 2D (**Figure 5.1, Figure 5.2**). We then assessed differences in GSK3 β expression and activity in 2D neurons versus 3D neurons and iAssembloids. We found that although GSK3B expression did not differ between culture systems, GSK3 β activity (denoted by the loss of the inhibitory phosphorylation site at serine 9), was 3-4 times higher in 3D culture versus 2D neurons (**Figure 5.1, Figure 5.2**). We asked if oxidative stress could induce this change in GSK3 β activity, and indeed, by adding rotenone to 2D neurons, which increases ROS levels (**Figure 5.2**), we found that the activity of GSK3 β increased (**Figure 5.1**). This suggests that GSK3 β activity prevents the NRF2-mediated protective response to oxidative stress, and that knockdown of GSK3 β confers a beneficial effect for cell survival by increasing the NRF2 response. We then asked whether knockdown of GSK3B could ameliorate downstream effects of increased ROS, such as lipid peroxidation⁷. Indeed, we found that knocking down GSK3B led to a decrease in the level of peroxidated lipids in neurons (**Figure 5.1, Figure 5.2**). Peroxidated lipids can drive neuronal death via ferroptosis^{7,37}. Therefore, we

hypothesized that the neuronal death occurring in 3D culture was due to this accumulation of peroxidated lipids driving ferroptosis. We found that the ferroptosis inhibitor ferrostatin trended towards improved neuronal survival in 3D neurons (**Figure 5.1**) suggesting neurons are dying in part via ferroptosis.

Since neuronal activity is much higher in iAssembloids compared to 2D-cultured neurons (**Figure 2.1**), and in 3D-cultured neurons compared to 2D-cultured neurons (**Figure 5.1**), we hypothesized that neuronal activity could be driving higher levels of ROS and thus ferroptosis induced cell death. We found that ROS levels of neurons in iAssembloids which contains glia are lower when compared to those of 3D neuronal monocultures (**Figure 5.2**). Interestingly, neuronal activity is much higher in 3D-cultured neurons compared to iAssembloids, which contain glial cells, suggesting that glial cells may prevent neuronal hyperexcitability, such as through glutamate-buffering or through ROS clearing activity of astrocytes³⁸⁻⁴⁰. Previous studies have shown that high levels of neuronal activity lead to a generation of ROS, which can drive excitotoxicity^{41,42}. To test whether neuronal activity drives the increase in ROS in our culture system, we added tetrodotoxin (TTX), a drug that blocks voltage-gated sodium channels and thus neuronal firing. We found that TTX treatment decreased ROS levels (**Figure 5.1**) and levels of peroxidated lipids (**Figure 5.1**). We also found that blocking neuronal activity improved neuronal survival in 3D neurons (**Figure 5.1**).

Excess neuronal activity can drive GSK3 β activity and prevent NRF2 from enacting protective functions

Based on the data above, we hypothesize that in 2D-monocultured neurons, there is a low level of neuronal activity and ROS levels remain low, thus providing no mechanism driving neuronal death via ferroptosis (**Figure 6**). However, in 3D monocultured neurons, higher levels of neuronal activity trigger a build-up in ROS, and both processes increase GSK3 β activity. This increase in GSK3 β activity blocks NRF2 from protecting neurons from oxidative stress, ultimately leading to neuronal death via ferroptosis (**Figure 6**). In GSK3 β knockdown neurons, the ability for GSK3 β to block NRF2 from protecting neurons from neuronal death is abrogated, enabling the protective NRF2 mediated oxidative stress response that improves neuronal survival (**Figure 6**). The presence of glial cells in iAssembloids partially reduces neuronal activity and ROS in neurons, thereby reducing, but not abrogating GSK3 β activity and its suppression of protective NRF2 activity (**Figure 6**).

Chapter 5 : CRISPRi-based screens in APOE3 and APOE4 astrocyte-neuron 3D co-cultures

CRISPRi screens in iAssembloids uncover cell non-autonomous effects of astrocytes expressing different Apolipoprotein E (APOE) variants on neuronal health

We next applied the iAssembloid system to specifically study the effects of glial cells on neurons in a neurodegenerative disease relevant context. We were particularly interested in the genetic risk factor for Alzheimer's Disease, APOE- ϵ 4. APOE- ϵ 4 is a variant of the APOE gene and carriers of the APOE- ϵ 4 allele are much more likely to develop Alzheimer's Disease^{43,44}. APOE- ϵ 3 allele is considered "neutral." APOE is mainly expressed by glial cells such as astrocytes, although stress can induce its expression in neurons^{45,46}. However, how expression of these APOE variants in glial cells may affect neuronal survival is currently unknown.

To assess this, we designed a CRISPRi-based screen in which we transduced ~16,000 sgRNAs targeting ~2,000 genes in the "druggable genome" (H1 library)²¹ into hiPSCs expressing dox-inducible NGN2 and CRISPRi machinery. These neurons are homozygous for the APOE- ϵ 3 allele. We then differentiated these cells into neurons and co-cultured them with either APOE- ϵ 3 homozygous or APOE- ϵ 4 homozygous astrocytes for 14 days. We then sequenced the sgRNAs from the surviving neurons to determine the sgRNA frequencies in the starting versus surviving population in both conditions. By comparing the hits from the APOE- ϵ 3 and the APOE- ϵ 4 screen, we can uncover

pathways in neurons that differentially control their survival in the presence of APOE- ϵ 4 and the APOE- ϵ 3 astrocytes (**Figure 7**).

We conducted this screen in two different cell line background pairs: one where the neurons and astrocytes were not isogenic with astrocytes generated from the TCW 1E33 and 1E44 background⁴⁷, and neurons from WTC11 background⁶ (**Figure 7**), and another where the neurons and astrocytes were isogenic with cells from the iNDI project⁴⁸ (**Figure 7**). We found that hits obtained from both screens were similar, with pathways that were related to insulin signaling, MAPK pathway, metabolism, immune response, microtubule regulation, and synaptic transmission were most consistently different between the APOE- ϵ 3 and APOE- ϵ 4 conditions (**Figure 7**). Pathways such as insulin signaling, the MAPK pathway and metabolism are all intimately interlinked. All in all, we hypothesize that APOE- ϵ 4 expressing astrocytes may sensitize neurons to neuronal death due to alteration in one or many of these pathways.

Chapter 6 : Discussion

In this study, scalable 3D co-culture model, which we call iAssembloids, in combination with CRISPRi-based functional genomics screens to uncover how genes regulate neuron and glia function and interactions in a scalable manner. iAssembloids are fast to generate and homogeneous, which is ideal for CRISPR-based screens and other systems requiring high throughput screening (e.g., compound screening). Relative to monoculture, neurons, and glia in iAssembloids adopt more mature states. This enabled us to uncover a pathway controlling the survival of highly active neurons, which would have been difficult to do in 2D monoculture neurons. iAssembloids allow precise control of glial cell genetic backgrounds, allowing causal assessment of those genetic backgrounds on neuronal survival. We demonstrate this application to examine the effect of glia-expressed APOE- ϵ 4, a variant of the gene APOE known to increase risk of late-onset Alzheimer's Disease^{43,44}, on neuronal survival.

GSK3 β as a player in oxidative stress in response to neuronal activity

Many studies have reported a link between seizure activity and dementia in advanced stages⁴⁹⁻⁵¹. However, developing unprovoked seizures later in life have also been associated with early-stage dementia and have been posited as one of the first signs of neurodegenerative disease⁵². Our model shows that neurons in iAssembloids are more electrophysiologically active than in monoculture, thus allowing us to assess the pathways that are affected by neuronal activity in neuronal survival.

Our CRISPRi-based screens uncovered GSK3 β as a factor controlling the response to neuronal activity and oxidative stress. In Alzheimer's Disease, GSK3 β plays a major

role in phosphorylating Tau, which is thought to contribute to Tau aggregation^{28,53}. It is also commonly found to be hyperactivated in brains of Alzheimer's Disease patients^{24,54}. Here, we unravel a secondary role for GSK3 β that may contribute to its role in neurodegeneration. In this case, neuronal activity, and an increase in reactive oxygen species (ROS) can drive GSK3 β activity. GSK3 β blocks NRF2 from mounting a protective response to oxidative stress. We also find that the addition of glial cells can be protective against the generation of reactive oxygen species, which is consistent with previously described roles of glia in response to neuronal activity and oxidative stress^{38,39}. Therefore, we hypothesize that in disease, the combined effects of aberrant neuronal activity driving increases in ROS and GSK3 β activity as well loss of glial cell homeostasis in protecting neurons against ROS can drive neurodegeneration. Our findings suggest that there may be some beneficial effects of GSK3 β inhibition in neurons that enables the activation of a protective response against oxidative stress in excitotoxic conditions. GSK3 β is a kinase that has many downstream targets, making it difficult to target as a therapeutic target⁵⁵. However, more specific therapeutic strategies to boost the oxidative stress response could recapitulate some beneficial aspects of GSK3 β inhibition, without off-target effects.

APOE-ε3 vs APOE-ε4 screen suggest differences in insulin signaling, MAPK signaling, and regulation of microtubules

In our screen comparing APOE-ε3 astrocyte-neuron co-cultures to APOE-ε4 co-cultures, we found many differential targets relating to insulin signaling and the MAPK pathway, which is closely linked to insulin signaling⁵⁶. This data supports previous experiments performed in mice⁵⁷. Future studies will focus on validating these hits and identifying how pathways that may drive differential sensitivity to neuronal death in APOE-ε4 vs APOE-ε3 cultures may be interlinked.

Limitations of this study

Our model does not recapitulate all aspects of an in vivo brain environment. It is a simplified model that does not mimic the vast amount of cellular heterogeneity found inside a human brain. For instance, iAssembloids do not contain vasculature or other glial cell types, such as oligodendrocytes. Due to the nature of the NGN2-induced neurons we use, we also cannot accurately model aspects of brain development or tissue architecture. However, this platform does enable mixing and matching neuronal and glia subtypes, pinpointing the specific contributions to a phenotype by a specific glial subtype. It also enables the examination of non-cell autonomous contributions of specific genetic variants on neuronal function, such as APOE-ε4. The model can be expanded upon by integrating vasculature as previously shown by others^{58,59} and can be applied to study the glial contributions of other genetic risk factors.

Neuronal survival is highly relevant phenotype for neurodegenerative diseases.

However, it is a late phenotype that is the result of many preceding functional changes a

neuron may experience. Future studies will investigate the functional effects of genetic perturbations such as GSK3 β on more complex phenotypes such as neuronal activity, states of glial cells, and intercellular connections. Beyond these studies, this model could provide an additional platform to assess the effectiveness of small molecules on neuronal phenotypes relating to neuronal activity. In the future, this model could also provide a unique platform to examine cell-cell interactions in the context of other neurological diseases where neuron-glia interactions play an important role, such as other neuropsychiatric and neurodegenerative disorders.

Chapter 7 : Acknowledgements and author contributions

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Author contributions

E.L. and M. Kampmann contributed to the study's overall conception, design, and interpretation and wrote the manuscript and created the figures with input from the other authors. E.L. optimized the iAssembloid model with helpful discussions from M.Koontz and E.M.U. E.L. characterized the iAssembloid model, performed and analyzed the CRISPRi-based screens (including survival screens and CROP-seq), and performed mechanistic follow-up on relevant targets with guidance from M. Kampmann. N.D. and M. Kampmann. selected genes for and cloned the neurodegeneration library. M. Koontz supplied serum-free astrocytes for CRISPRi-based screens. C.B. and E.L. generated data from multi-well multi-electrode arrays in iAssembloids 3D co-cultures and S.C.B. generated corresponding data in 2D monoculture neurons. A.J.S. performed western blots in rotenone induced 2D monocultured neurons. I.V.L.R. and E.L. generated APOE3 and APOE4 KOLF2.1 astrocytes used in this study. O.M.T. generated 6TF hiPSC cell line expressing mScarlet.

Declarations of interest

M. Kampmann is an inventor on US Patent 11,254,933 related to CRISPRi and CRISPRa screening, serves on the Scientific Advisory Boards of Engine Biosciences, Casma Therapeutics, Cajal Neuroscience and Alector, and is an advisor to Modulo Bio and Recursion Therapeutics.

Chapter 8 : Methods

hiPSC Maintenance and Culture

We cultured hiPSCs in the WTC11⁶⁰ and KOLF2.1⁴⁸ backgrounds on plates coated with Matrigel (Corning, Cat. No. 356231) at a concentration of 1:100 in Knockout DMEM (Gibco/Thermo Fisher Scientific, Cat. No. 10829-018). For plating, hiPSCs were seeded with 10 μ M Rock Inhibitor (Y-27632, Biotechne/Tocris Cat. No. 1254) diluted in GibcoTM StemFlexTM Medium (Gibco/Thermo Fisher Scientific, Cat. No. A3349401). Media (no RI) was exchanged when colonies reached ~20 cells in size. hiPSCs were maintained with media changes every day. When hiPSCs reached ~80% confluence, hiPSCs were coated with either GibcoTM Versene Solution (0.48 mM) (Gibco/Thermo Fisher Scientific, Cat. No. 15040066) for routine passaging or StemProTM AccutaseTM Cell Dissociation Reagent (Gibco/Thermo Fisher Scientific, Cat. No. A1110501) for differentiation and incubated at 37 °C for 5-7 minutes. After dissociation, cells were collected and rinsed with 1X DPBS (MilliporeSigma, Cat. No. D8537). Cells were centrifuged at 250g for 5 minutes and replated.

hiPSC derived Neurons

hiPSC-derived neurons were differentiated using methods previously described^{6,7}. Briefly, we dissociated hiPSCs containing doxycycline-inducible NGN2 in the AAVS1 locus and pC13N-dCas9-BFP-KRAB in the CLYBL locus and re-plated cells on Matrigel coated plates with Pre-differentiation medium containing Knockout DMEM/F12 (Gibco/Thermo Fisher Scientific, Cat. No. 12660-012), 1X GlutaMAX Supplement (Gibco/Thermo Fisher Scientific, Cat. No. 35050-061), 1X MEM Non-Essential Amino

Acids (Gibco/Thermo Fisher Scientific, Cat. No. 11140-050), 1X N2 Supplement (Gibco/Thermo Fisher Scientific, Cat. No. 17502-048), 10uM ROCK inhibitor (Y-27632, Biotechne/Tocris, Cat. No. 1254), 1 µg/mL mouse laminin (Thermo Fisher Scientific, 23017-015), 10 ng/mL BDNF (PeproTech, Cat. No. 450-02B) and 10 ng/mL NT3 (PeproTech, Cat. No. 450-03B) with 2µg/mL doxycycline (Takara Bio, Cat. No. 631311) to induce expression of NGN2. Half of the media was exchanged over the next 2 days until hiPSC-derived neurons were ready for iAssembloid generation (Day 0). For monocultures, pre-differentiated neurons were plated onto PDL-coated plates (Corning, Cat. No. 356469) and maintained in differentiation media: Base media: 50% DMEM/F12 (Gibco/Thermo Fisher Scientific, Cat. No. 11320-033) and 50% Neurobasal-A (Gibco/Thermo Fisher Scientific, Cat. No. 10888-022) with supplements: 1X MEM Non-Essential Amino Acids, 0.5X GlutaMAX Supplement, 0.5X N2 Supplement, 0.5X B27 Supplement (Gibco/Thermo Fisher Scientific, Cat. No. 17504-044), 10 ng/mL NT-3, 10 ng/mL BDNF and 1 µg/mL Mouse Laminin.

hiPSC derived Astrocytes: Serum Free Protocol

Astrocytes were generated as described previously^{16,61,62}. WTC11 iPSCs were cultured and expanded in six-well plates using Essential 8 media (Gibco/Thermo Fisher Scientific, Cat. No. A1517001). Once iPSCs were confluent, cells were enzymatically dissociated into a single cell suspension using Accutase (Gibco/Thermo Fisher Scientific, Cat. No. A1110501). The resulting single cell suspension was transferred into T-75 flasks in ~10 ml of neural induction media (50% DMEM/F12 (Gibco/Thermo Fisher Scientific, Cat. No. 11320-033) and 50% Neurobasal-A (Gibco/Thermo Fisher Scientific,

Cat. No. 10888-022) with supplements: 1X MEM Non-Essential Amino Acids (Gibco/Thermo Fisher Scientific, 11140050), 1X GlutaMAX Supplement (Gibco/Thermo Fisher Scientific, Cat. No. 35050-061), 1X N2 Supplement (Gibco/Thermo Fisher Scientific, Cat. No. 17502-048), 0.5X B27 Supplement (Gibco/Thermo Fisher Scientific, Cat. No. 17504-044), Anti-Anti (Gibco/Thermo Fisher Scientific, Cat. No.15240062) to allow iPSCs to aggregate into small spheroids in suspension. To facilitate neural induction, dual SMAD inhibitors DMH1 (Tocris, Cat. No. 73634) and SB-431542 (StemCell Technologies, Cat. No. 72234) are added to the media. Once rosette formation is observed, spheroids are transferred to culture dishes coated with Matrigel (Corning, Cat. No. CB40230C) and allowed to attach and distributed across culture dish surfaces. Once sufficiently distributed, neuroepithelial rosettes are mechanically removed and isolated. Neuroepithelial spheroids are maintained in suspension in neural media (DMEM-F12, N2/B27 supplements, Heparin (Stemcell Tech, Cat. No. 07980), Anti-Anti) plus mitogenic growth factors EGF (Peprotech, Cat. No. AF-100-15) and FGF β (Peprotech, Cat. No. 100-18B). Spheroid cultures are maintained in suspension and triturated/dissociated regularly to maintain size and structural homogeneity of neuroepithelial spheroids. After ~180 days of culture maintenance, neuroepithelial progenitor cells have differentiated into unipotent populations of glial progenitors.

hiPSC derived Astrocytes: Serum Based Protocol

Astrocytes were differentiated as described previously⁴⁷. hiPSCs were cultured on Matrigel and in StemFlex media supplemented with 10 μ M RI. hiPSCs were then differentiated to neural precursor cells by dual SMAD inhibition using 0.1 mM

LDN193189 (Biotechne/Tocris Cat. No. 6053) and 10 mM SB431542 (Biotechne/Tocris, Cat. No. 1614) in N2/B27 media inhibition in embryoid bodies (EB) media (DMEM/F12 + 1x N2 + 1x B27-VA). EBs were cultured in AggreWell™800 (StemCell Technologies, Cat. No. 34815) plates rinsed with Anti-Adherence solution (StemCell Technologies, Cat. No. 07010) for 7 days and then plated onto a Matrigel coated plate. Neural rosette formation occurs at day 14 where rosettes were selected for with STEMdiff™ Neural Rosette Selection Reagent (StemCell Technologies, Cat. No. 05832). NPCs were expanded in NPC media which contained DMEM/F12 + 1x N2 + 1x B27-VA supplemented with FGF2 (Biotechne, Cat. No. 233-FB-010). Using Magnetic Activated Cell Sorting (MACS). NPCs were enriched by sorting against neural crest cells (Miltenyi Biotech, Cat. No. 130-097-127) and for CD133+ cells (Miltenyi Biotech, Cat. No. 130-097-049) using LD (Miltenyi Biotech, Cat. No. 130-042-901) and LS (Miltenyi Biotech, Cat. No. 130-042-401) columns respectively. Magnet was 3D printed using previously published protocol⁶³. by NPCs were maintained in NPC media (were validated with immunostaining for SOX2 (Rabbit anti-SOX2, Abcam, Cat. No. ab97959) and Nestin (rabbit anti-Nestin, Abcam, Cat. No. ab92391). Dissociated NPCs were transferred to astrocyte media (ScienCell, Cat. No. 1801) and cultured for 30 days on Matrigel until they stained positively for S100B (mouse anti-S100B, MilliporeSigma, Cat. No. S2532) and NFIA (MilliporeSigma, Cat. No. HPA008884).

hiPSC derived Microglia

Microglia were differentiated as previously described. Briefly, hiPSCs expressing six inducible transcription factors (MAFB, CEBP α , IRF, PU1, CEBP β , IRF5) were grown in

Essential 8™ Basal Medium (Gibco/Thermo Fisher Scientific, Cat. No. A15169-01), 10 µM ROCK inhibitor, and 2 µg/ml Doxycycline on Matrigel and 10 cm Poly-D-Lysine coated plates at a density of 1.5 million cells per dish. After 2 days, cells were grown in a differentiation media consisting of Advanced DMEM/F12 (Gibco/Thermo Fisher Scientific, Cat. No. 35050-061), 1X GlutaMAX, 2ug/ml doxycycline, 100 ng/mL Human IL34 (Peprotech; Cat. No. 200-34) and 10 ng/mL Human GM-CSF (Peprotech; Cat. No. 300-03). Two days later, media was exchanged for iTF-Microglia media consisting of Advanced DMEM/F12, 1X GlutaMAX, 2 µg/mL doxycycline, 100 ng/mL Human IL-34, 10 ng/mL Human GM-CSF, 50 ng/mL Human M-CSF (Peprotech; Cat. No. 300-25) and 50 ng/mL Human TGFB1 (Peprotech; Cat. No. 100-21C). On day 8, cells were dissociated with TypLE express (Gibco/Thermo Fisher Scientific, Cat. No. 12605-028) and seeded into iAssembloids.

Generation of iAssembloids

AggreWell™800 plates were first rinsed with Anti-Adherence rinsing solution and then washed with DMEM/F12. Neurons (2 million) and astrocytes (0.66 million) were seeded at a 3:1 ratio within each well in Astrocyte Media (0.5X B27 + 0.5X N2 + 1X GlutaMAX in DMEM/F12). Plates were centrifuged at 100g for 3 minutes until cells settled at the bottom. Half of the media were exchanged with fresh media every day. Microglia were seeded a week after initial Neuron-Astrocyte assembloid formation at one-third the number of astrocytes (0.2 million). Half the media was exchanged with iTF-Microglia media and doxycycline (2 µg/mL). Subsequently, half the media is then removed every

other day and replaced with Astrocyte media supplemented with iTF-Microglia media cytokines.

Tissue clearing for imaging cells expressing fluorescent proteins in iAssembloids

To obtain images of cells within the iAssembloids, hiPSC derived neurons expressing BFP, astrocytes expressing membrane GFP and microglia expressing mScarlet were seeded into iAssembloids with cells not expressing any fluorescent proteins at a 1:10 ratio. iAssembloids were then cleared with the ClearT2 protocol⁶⁴. In brief, iAssembloids were first fixed with 4% paraformaldehyde (16% stock, diluted 1:4 in DPBS, Electron Microscopy Sciences, Cat. No. 15710) and then incubated in a 25% formamide (MilliporeSigma, Cat. No. F9037) /10% DPBS solution. Then, iAssembloids were incubated in a 50% formamide / 20% DPBS solution and imaged with a confocal microscope.

Immunohistochemistry

iAssembloids were fixed with 4% paraformaldehyde in DPBS for 20 minutes on a shaker. iAssembloid were then washed with 0.33% Triton X-100 in DPBS 3 times for 5-10 minutes each time. Following washing, iAssembloids were blocked with 10% Normal Goat Serum + 1% BSA (blocking buffer) for an hour at room temperature. iAssembloids were incubated in antibodies diluted in blocking buffer overnight at 4°C. Following incubation, iAssembloids were washed with 0.33% Triton X-100 in DPBS 3 times for 5-10 minutes and incubated in secondary antibody diluted in blocking buffer for 1-2 hours at room temperature. iAssembloids were mounted and imaged on a confocal

microscope. iAssembloids for NRF2 images were sectioned into 20 μm sections.

Primary antibodies used for this study were as follows: rabbit anti-Iba1 (Wako, Cat. No. 019-19741), mouse anti-NeuN, clone A60 (MilliporeSigma, Cat. No. MAB377), mouse anti-S100 (β -Subunit) (MilliporeSigma, Cat. No. S2532), chicken anti-Tuj1 (Aves Labs, Cat. No. TUJ-0020), rabbit anti-NRF2 (Abcam, Cat. No. ab62352).

Nuclear isolation from iAssembloids for single-nuclei sequencing

Nuclear isolation was performed as previously described^{65,66}. First, ~300 iAssembloids were dounce homogenized in 5 ml of lysis buffer (0.25 M sucrose, 25 mM KCl, 5 mM MgCl_2 , 20 mM Tricine-KOH, pH 7.8, 1 mM DL-Dithiothreitol (DTT) (MilliporeSigma, Cat. No. D0632-1G), 0.15 mM Spermine tetrahydrochlorine (MilliporeSigma, Cat. No. S1141-1G), 0.5 mM Spermidine trihydrochloride (MilliporeSigma, Cat. No. S2501-1G), 1X protease inhibitor (MilliporeSigma, Cat. No. 4693159001), and RNase Inhibitor (Promega, Cat. No. N2615) with 10 strokes. Following initial dounce homogenization, IGEPAL CA-630 (MilliporeSigma, Cat. No. I8896-50ML) was added to a final concentration of 0.3% and the sample was homogenized with 5 more strokes. Cells were then filtered through a 40 μm cell filter, and Optiprep was added to the cell solution for a final concentration of 25% Optiprep (MilliporeSigma, Cat. No. D1556-250ML). This solution was layered onto a 30%/40% Optiprep gradient and centrifuged at 10,000g for 18 minutes using a SW41-Ti rotor. The nuclei were collected at the 30%/40% Optiprep interface and quantified using the Countess FL Automated Cell Counter (ThermoFisher Scientific/Invitrogen, Cat. No. AMQAX2000). Droplet-based nuclear capture and library preparation were performed using the Chromium Single Cell Gene Expression

workflow. Nuclei were brought to a concentration of 1000 nuclei/ μ L in 30% Optiprep solution before loading according to manufacturer's specifications. 10,000 cells per sample were targeted for sequencing. Sample preparation and library generation were performed using the v3.1 10X 3' single cell RNA sequencing library kit (10X Genomics, Cat. No.1000147). cDNA fragment analysis was performed using the Agilent 4200 TapeStation System with the D5000 HS Kit (Agilent, Cat. Nos. 5067-5592, 5067-5593, 5067-5594). Sequencing parameters and quality control were performed as described by The Tabula Muris Consortium⁶⁷.

Multi-well Multielectrode Array (Axion): iAssembloids

6 (Axion Biosystems, Cat. No. M384-tMEA-6B and 24 (Axion Biosystems, Cat. No. M384-tMEA-24W) well Cytoview MEA plates were coated with poly-L-ornithine (0.01%), fibronectin (1 μ g/mL) and laminin (5 μ g/mL) to facilitate iAssembloid adherence. iAssembloids were positioned into the well over the electrodes using a microscope. iAssembloids were equilibrated in BrainPhys™ Neuronal Medium (StemCell Technologies, Cat. No. 05790) 24 hours before plating to facilitate comparison to 2D monocultured neurons. Measurements were taken using the Maestro Edge multiwell microelectrode array (MEA) and Impedance system (Axion Biosystems) at 15 minute intervals with the Maestro MEA platform software (Axion Biosystems) with the "Neural" default setting.

Multi-well Multielectrode Array (Axion): 2D monoculture

24 well Cytoview MEA plates (Axion Biosystems, Cat. No. M384-tMEA-24W) were coated with 0.1% PEI (MilliporeSigma, Cat. No. 03880) dissolved in 1X Borate Buffer (10mM Boric Acid, 2.5mM Sodium Tetraborate, 7.5mM NaCl, adjusted to pH 8.4) overnight at 37°C, washed with water, then dried at room temperature. Wells were then coated with BrainPhys™ media and 15 µg/mL Laminin. Cells were then seeded at a density of 200,000 cells per well on day 0 of differentiation. Measurements were taken using the Maestro Edge multiwell microelectrode array (MEA) and Impedance system (Axion Biosystems) at 3 x 5-minute intervals with the Maestro MEA platform software (Axion Biosystems) with the “Neural” default setting after 14 days in culture.

CRISPRi-based pooled screens

The H1 CRISPRi library was packed into lentivirus as previously described⁶. Briefly, H1 plasmid library with the top 5 sgRNA per gene²¹ were transfected into HEK293s using the TransIT®-Lenti Transfection Reagent (Mirus, Cat. No. MIR 6606) along with third generation lentiviral packaging mix. Viral particles were harvested by removing the supernatant and filtering it through a 45 nm syringe filter. Virus was concentrated using cold Lentivirus Precipitation Solution (Alstem; Cat. No. VC100) was added to this filtered solution at a 1:4 ratio. The mixture was then centrifuged at 1500g for 30 minutes at 4°C. The supernatant was then removed, and the virus was resuspended in StemFlex. CRISPRi-NGN2 expressing iPSCs were then transduced at a MOI < 0.7. Cells expressing sgRNAs were then selected for with 2 µg/mL puromycin (Gibco/Thermo Fisher Scientific, Cat. No. A1113803) for 2 days as cells were expanded for screens.

After expansion, cells were pre-differentiated and seeded into iAssembloids for screens. At days 1, 14 and 28 post-assembly of the iAssembloids, iAssembloid were harvested using a wide-orifice pipette tip and washed with DPBS, after which samples were frozen for later sample preparation. Genomic DNA was extracted with the NucleoSpin® Blood XL (Macherey Nagel, Cat. No. 740950.10) and samples were prepared for sequencing on an Illumina NextSeq 500 based on previously described protocols^{68,69}

Neurodegeneration library gene selection and cloning

sgRNAs targeting risk genes for neurodegenerative diseases were selected from GWAS and other published studies. Protospacer sequences and gene targets are listed in Table S2. This small, focused library consists of 5 sgRNA targeting 376 genes. Pooled sgRNA libraries were generated using the same protocol as previously described^{69,70}. Briefly, an oligonucleotide pool that encoded the library was synthesized by Agilent, amplified via PCR and then cloned into the pLG15 backbone⁶⁹. HiPSCs expressing CRISPRi-NGN2 machinery were then transduced with the sgRNA library at an MOI < 0.3. Cells were then differentiated and utilized for screens in iAssembloids.

Focused library secondary screens

146 sgRNAs, including 136 targeting sgRNAs (2 sgRNAs per gene) and 10 non-targeting sgRNAs were cloned into the pMK1334 CROP-seq vector⁶. sgRNA oligos were synthesized (top and bottom strands; IDT), annealed, and pooled in equal amounts (Table S4). hiPSCs expressing CRISPRi-NGN2 machinery were then transduced with the sgRNA library at an MOI < 0.3 and differentiated. Timepoints at 1

day after seeding into AggreWell™ 800 plates and 14 days after seeding into AggreWell™ 800 plates were taken. The following conditions were then tested: 2D monoculture, standard iAssembloids, 3D neurons and microglia, 3D neurons and astrocytes, and 3D neurons by themselves. Different media conditions were also tested including the normal condition (Standard w/cytokines), without supplementing any microglia media (Astrocyte w/cytokines), adding in microglia with microglia base media, but with no cytokines (Standard), adding in only astrocyte media with no cytokines (Astrocyte) and with BrainPhys™. Half the media was exchanged every day. Genes with consistent phenotypes across conditions were selected for CROP-seq.

CROP-seq

Pooled sgRNA library (Table S4) consisting of 2 sgRNAs per targeted gene and 4 non-targeting controls was selected from focused library secondary screens. Libraries were cloned and transduced as described with the same method above in focused library secondary screen with an MOI < 0.30.

Sample preparation for CROP-seq

iAssembloids and 2D monocultured neurons were dissociated with the Papain (Worthington; Code: PAP2; Cat. No.LK003178) diluted in DMEM/F12 at 20U/mL Papain solution was warmed to 37°C and 500 µL was added to iAssembloids and incubated for 5 minutes. iAssembloids were triturated 15 times and filtered through a 22 µm cell filter. After filtering, neurons were selected for using the anti-PSA-NCAM microbeads kit (Miltenyi Biotec, Cat. No. 130-092-981), similarly to previously described methods⁷¹.

Cells were centrifuged at 300g for 10 minutes and resuspended in 60 μ L of PBS + 0.5% BSA (MilliporeSigma, Cat. No. A7979). To this suspension, 20 μ L of Anti-PSA-NCAM microbeads were added. The mixture was incubated at 4°C for 15 minutes. Cells were spun down at 300g for 10 minutes and washed with PBS + 0.5% BSA. MACS was performed according to manufacturer's specifications. After selection, purity was evaluated with flow cytometry. Sample preparation and library generation were performed using the v3.1 10X 3' single cell RNA sequencing library kit. cDNA fragment analysis was performed using the Agilent 4200 TapeStation System. A portion of the cDNA was utilized to enrich for sgRNA sequences as previously described⁶. Sequencing parameters and quality control were performed as described by The Tabula Muris Consortium⁶⁷.

Flow cytometry: Cell Staining

Cells from iAssembloids and 2D neurons were dissociated with papain as described above and stained according to manufacturer's protocols. Cells were washed three times with PBS after dissociation and incubated in DMEM/F12 with Liperfluor at 5 μ M (Dojindo Molecular Technologies Inc., Cat. No. L24810) for 30 min or CellROX™ Orange Reagent at 2.5 μ M (ThermoFisher Scientific/Invitrogen, Cat. No. C10443). Cells were then spun down and washed 2 more time with DPBS before analysis on the flow cytometer. Flow cytometry analysis was conducted with FlowJo v10.8.1.

Flow cytometry: GSK3B vs non-targeting guide survival assay

Neurons expressing two different GSK3B targeting guides and BFP were seeded at a 1:1 ratio with neurons expressing two different non-targeting guides expressing GFP. Seeding ratios were confirmed with flow cytometry before starting culture. After 14 days in culture in either iAssembloids or in monoculture, iAssembloids and neurons were dissociated with papain as described above and the number of cells expressing BFP (GSK3B knockdown) vs GFP (non-targeting guide) were quantified using FlowJo v10.8.1.

Quantification of viability using trypan blue

Cells from iAssembloids and 2D neurons were dissociated with papain as described above and stained at 1:1 ratio with trypan blue solution (0.04%) (ThermoFisher Scientific/Gibco, Cat. No., 5250061). The cell and trypan blue mixture were then loaded into the Countess™ Cell counter. Two counts were taken for each sample and averaged. Average percent viability was recorded based on percent of trypan blue negative cells. Tetrodotoxin (Biotechne/Tocris, Cat. No. 1069) was used at a final concentration of 1.5 μ M for a duration of 1 week, and ferrostatin 1 (Biotechne/Tocris, Cat. No. 5180) at 10 μ M for a duration of 1 week.

Western Blot

Approximately equal numbers of cells for both 2D monoculture, 3D monoculture and iAssembloids were lysed in 4X NuPAGE™ LDS Sample Buffer (ThermoFisher Scientific/Invitrogen, Cat. No., NP0007) + 10% 2-Mercaptoethanol, boiled for 10 minutes

at 98 °C and then diluted 1:1 with DPBS. Samples were then loaded at equal volume into NuPAGE 4–12% Bis-Tris gels (ThermoFisher Scientific/Invitrogen, Cat. No. NP0336BOX). Gels were then transferred onto nitrocellulose membranes in Towbin buffer (25mM Tris base, 192 mM glycine, 20% (v/v) methanol) and blocked with 4% BSA in TBST (1X TBS + 0.1% Tween 20) followed by overnight incubation with primary antibodies at 4°C. Membranes were then washed 4X with TBST for 5 minutes and incubated with secondary antibodies at room temperature for 45 minutes. Membranes were imaged on the Odyssey Fc Imaging System (LI-COR, Cat. No. 2800) using Image Studio (v5.2) software. Primary antibodies used were mouse anti-GAPDH (Santa Cruz Biotechnology, Cat. No. sc-47724), rabbit anti-GSK3 β (Cell Signaling, Cat. No. 12456S), and rabbit-anti S9 GSK3 β (Cell Signaling, Cat. No. 9336S). Secondary antibodies were IRDye 680RD goat anti-mouse IgG (1:10,000 dilution, LI-COR, cat. no. 926-68070) and IRDye 800CW goat anti-rabbit IgG (1:10,000 dilution, LI-COR, cat. no. 926-32211).

Data Analysis: snRNA sequencing

Data processing was performed using CellRanger v4.0. Demultiplexing was performed using CellRanger mkfastq, a 10x-aware wrapper for bcl2fastq. Due to the large number of introns resulting from snRNA-seq, a pre-mRNA GTF was generated using the cellranger--mkref function. Briefly, GTF annotation rows were extracted based on the feature “transcript” of the original, pre-built GRCh38 GTF file. The feature types were then replaced from “transcript” to “exon.” Then cellranger--mkref was run to build a pre-mRNA GTF file. Alignment, filtering, barcode counting, and UMI counting was performed using cellranger count, though which gene expression matrices were

generated. Outputs from cellranger were then loaded into Seurat^{72,73}, where nuclei with less than 500 UMI or greater than 15000 UMIs were filtered out. Doublets were then removed with DoubletFinder⁷⁴. Using the NormalizeData() function in Seurat, raw counts per cell were normalized to the total expression in the cell, scaled to 10,000 transcripts per nucleus, and then log transformed. After normalization, 2000 of the most highly variable genes were identified using the FindVariableFeatures() function. The data were scaled such that the mean expression of each gene across all the cells is equal to 0 and the variance equal to 1, so that highly expressed genes do not dominate downstream analyses. Principal component analysis was performed using RunPCA() with default parameters and the first 10 PCAs were considered for clustering. Clustering was performed with the FindNeighbors() and FindClusters() functions at a resolution of 0.5. Uniform Manifold Approximation and Projection (UMAP) was calculated using the runUMAP() function in Seurat. Cell identities were then defined based on expression of cell type specific markers. Differentially expressed genes were identified by running FindMarkers() using the Wilcoxon-rank sum test.

Datasets from other publications were integrated using Seurat's pipeline for mapping and annotating query datasets⁷². Briefly, data were preprocessed as above and a list of "anchor" genes between datasets were found using the function FindIntegrationAnchors(). The data is then integrated using the function IntegrateData().

Pathway enrichment analysis for snRNAseq

Pathway enrichment analysis for snRNAseq for neurons, astrocytes, and microglia were performed using EnrichR to classify assigned pathways for specific clusters⁷⁵⁻⁷⁷.

CRISPRi-based screens

Pooled screens were analyzed as previously described⁶, with the MAGeCK-iNC pipeline. First, sequencing reads were aligned to a reference file using Bowtie v0.99 to determine sgRNA frequencies per sample. Further analysis was then performed using the MAGeCK-inc software. Gene scores were calculated for each gene, which is defined as phenotype score multiplied by the $-\log_{10}(\text{P value})$. Hit genes were defined as genes with a false discovery rate < 0.05 . Volcano plots were generated with ggplot2 and heatmaps were generated with the pheatmap function in R.

Pathway enrichment analysis for CRISPRi-based screens

GO term enrichment analysis was performed using The Database for Annotation, Visualization and Integrated Discovery (DAVID)^{78,79} with the library gene list as the background list and hits from the screens as the genes of interest.

Multiwell Multielectrode array (MEA) analysis

MEA analysis was performed with the default Axion's Integrated Studio (AxIS) software. The default settings were used for spike detection with the Adaptive Threshold Crossing method. and network burst detection setting was applied. Spike files were then imported

into the Axion neural metrics software where a 120 s snapshot of the total datafile was taken, and raster plots were generated.

CROP-seq differentially expressed genes analysis

sgRNA mapping was performed a previously published program and sgRNA assignment was performed using the program quantifyTranscriptome.R (<https://github.com/powellgenomicslab/CROP-seq>). Guides were only assigned if they were 1) the only guide found in the cell, 2) have more reads than three times the sum of other reads assigned to the other guides in the cell and the guides target a different gene, or 3) the guides for a single gene have 3x the sum of reads assigned to other guides in the cell. This information was then integrated as metadata for each sample as a Seurat object. UMAPs were generated with the same method as described above for snRNA-sequencing.

Differentially expressed genes were found using DESeq2 by pseudo bulking cells with the same sgRNA sequences by adapting previously described methods⁸⁰. Cells with targeting guides versus non-targeting guides with individual sgRNA sequences counting as replicates. The P value cut off was 0.05. All differentially expressed genes were identified and compared log2FC for these genes within the cells with targeting guides. Differentially expressed genes with a log2FC magnitude greater than 2 in at least one condition (one gene target) was selected for heatmap visualization. Genes were manually annotated with some guidance from EnrichR. Differentially expressed genes for GSK3B was then independently examined and plotted using ggplot2.

Image analysis for NRF2 localization in iAssembloids

Three different iAssembloid sections for GSK3B KD and non-targeting control were taken and analyzed after staining with the protocol described above. Images were taken with a Leica SoRA confocal microscope. Nuclei were recognized as nuclear localized BFP+ (blue) objects while NRF2+ cells were recognized in a separate channel (red+). Nuclei were then masked onto the NRF2 stain to determine the location of the nuclei in relationship to NRF2 levels and then the intensity of the objects were then quantified using the MeasureObjectIntensity function.

Data availability

snRNA, CROPseq and CRISPRi-based screen datasets generated in this study will be available on NCBI GEO. CRISPRi-based screens will be uploaded to CRISPRbrain (<https://www.crisprbrain.org/>). Protocols for iAssembloid culture, dissociation, and nuclear dissociation for snRNAseq will be uploaded to protocols.io.

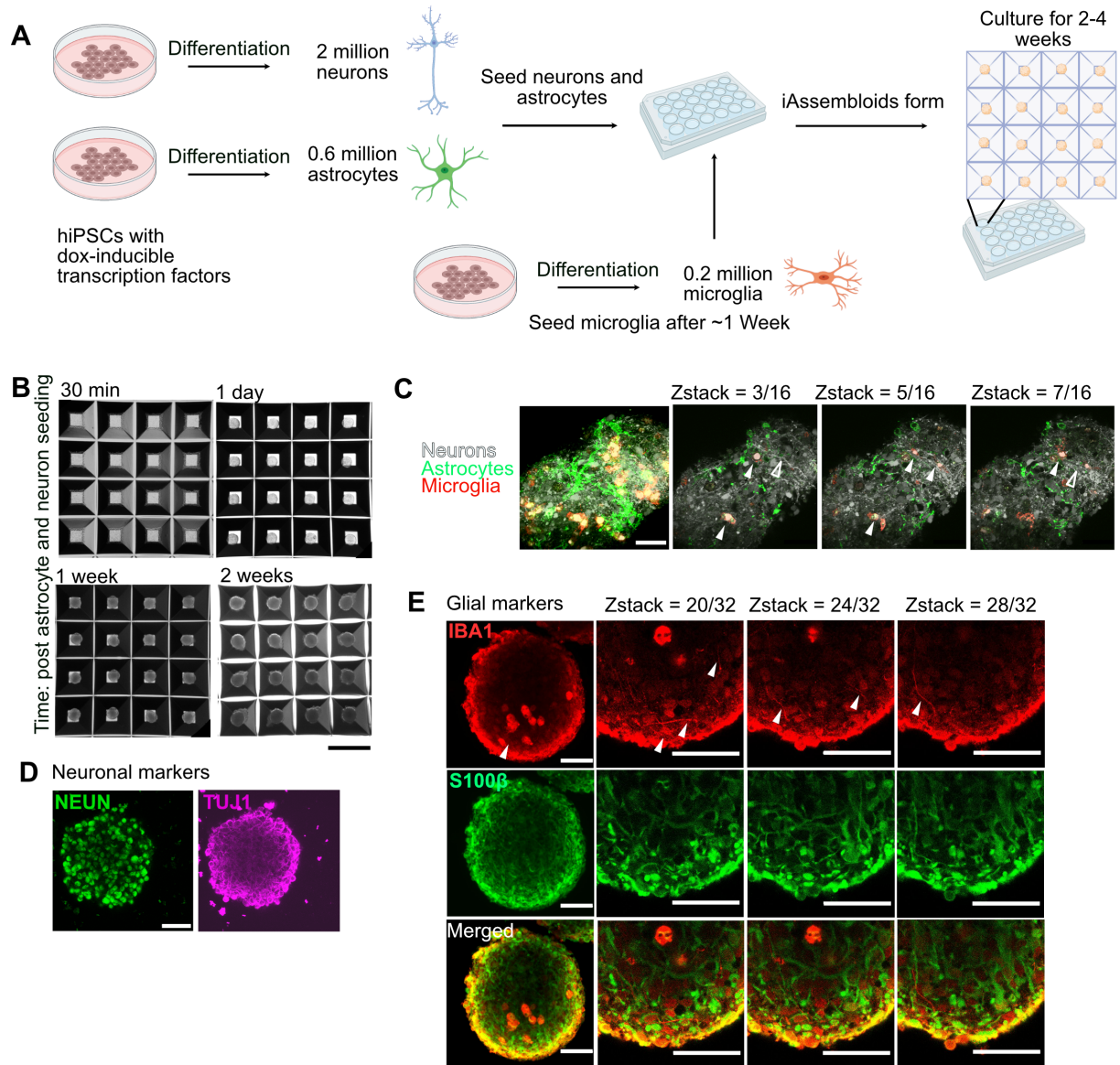


Figure 1. Integrating hiPSC-derived neurons, astrocytes, and microglia into three-dimensional cultures (iAssembloids)

(A) hiPSCs expressing CRISPRi machinery and inducible NGN2 are pre-differentiated and seeded with hiPSC-derived astrocytes at a 3:1 neurons:astrocyte ratio. After 1 week, hiPSC-derived microglia are seeded at one-third of the number of astrocytes.

(B) Brightfield images of iAssembloids 30 min, 1 day, 1 week and 2 weeks post seeding. Scale bar = 800 μ m.

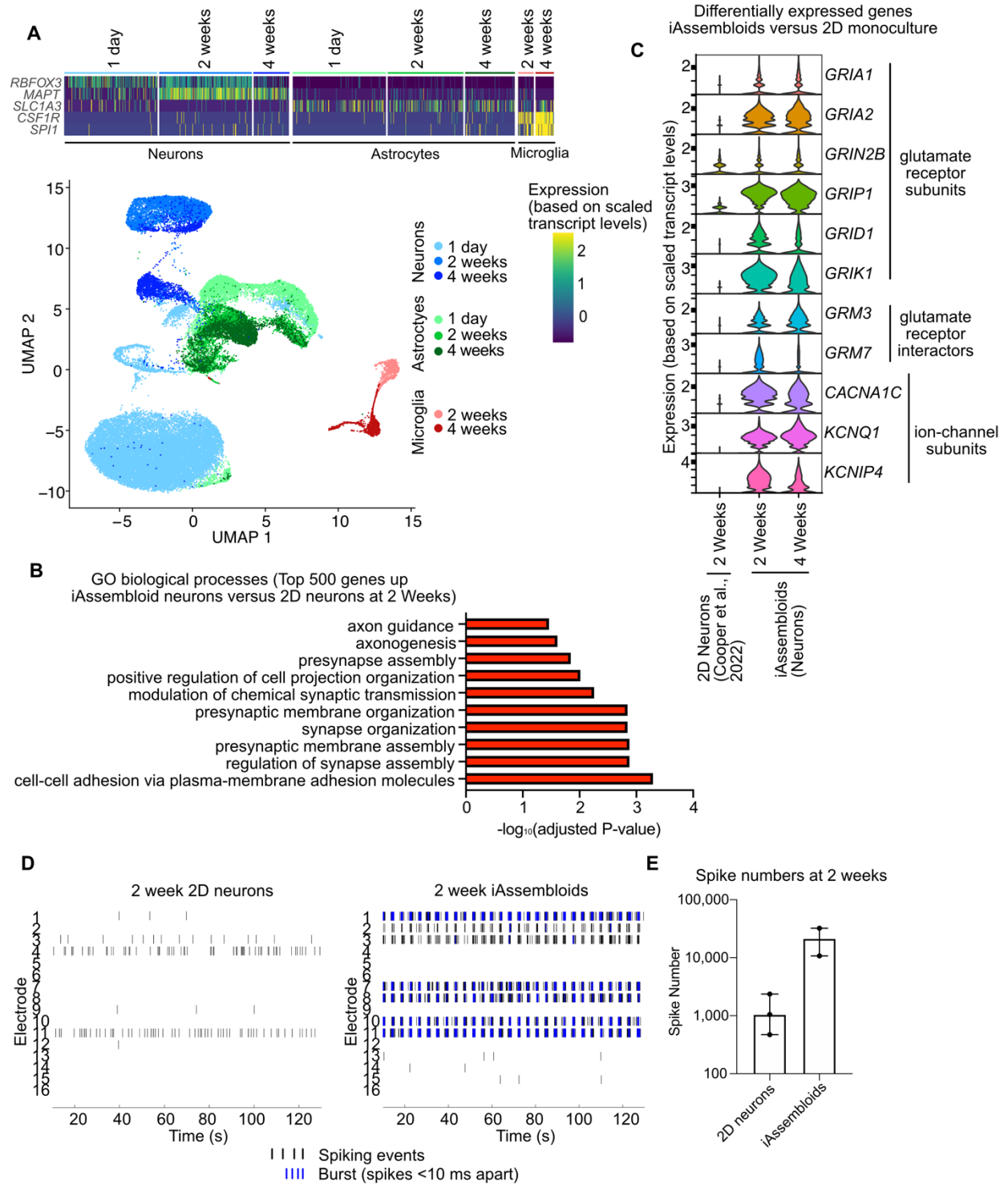
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(C) Neurons (gray), astrocytes (green) and microglia (red) expressing different fluorescent proteins were seeded to form iAssembloids. Left image: maximum-intensity projection, other images: individual images from the horizontal sample images (z-stack) generated from confocal microscopy. Arrows denote neurons co-localized with microglia (closed arrowheads) as well as neuronal extensions across the culture (open arrowheads). Scale bar = 50 μm . Images were taken 14 days post seeding into AggreWell 800 plates.

(D) Maximum intensity projections of iAssembloids stained with antibodies against neuronal markers NEUN and TUJ1. Scale bar = 50 μm . Images were taken 14 days post seeding into AggreWell™ 800 plates.

(E) Maximum intensity projections of iAssembloids stained with antibodies against the microglial marker IBA1 and the astrocyte marker S100 β . Scale bars = 50 μm . Images were taken 14 days post seeding into AggreWell™ 800 plates. Arrows denote microglia projections.



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Figure 2.1: Neurons cultured in iAssembloids are more functionally mature than monocultured neurons

(A) Single-nucleus RNA sequencing was performed on iAssembloids in culture for 1 day, 2 weeks and 4 weeks. In total, 43,182 nuclei passed quality control and are represented in this dataset. Cell types were assigned to clusters using cell type-specific markers such as RBFOX3 and MAPT for neurons, SLC1A3 (GLAST) for Astrocytes, and CSF1R and SPI1 for microglia. The heatmap is subsampled at 5,000 cells for visualization. UMAP is labeled by cell type and sample.

(B) Most significant Gene Ontology Biological Processes enriched in the 500 top genes expressed more highly in neurons in iAssembloids vs. 2D monocultured neurons. Adjusted P values were calculated using the Benjamini-Hochberg method for correction for multiple hypothesis testing.

(C) Analysis of selected differentially expressed genes comparing iAssembloid and monocultured neuronal culture systems shows that expression of glutamate receptor subunits, glutamate receptor interactors and various ion channel subunits is significantly higher in iAssembloids versus monoculture¹⁷.

(D) Example raster plot from multi-electrode array (MEA, Axion) analysis of spikes of neuronal activity in monocultured neurons versus iAssembloids.

(E) Bar graphs represent cumulative spike data over a 15-minute time span from 3 independent wells of 2D monoculture neurons and 2 independent wells of iAssembloids cultured on a MEA plate. Bars represent the median and the error bars represent the 95% confidence interval.

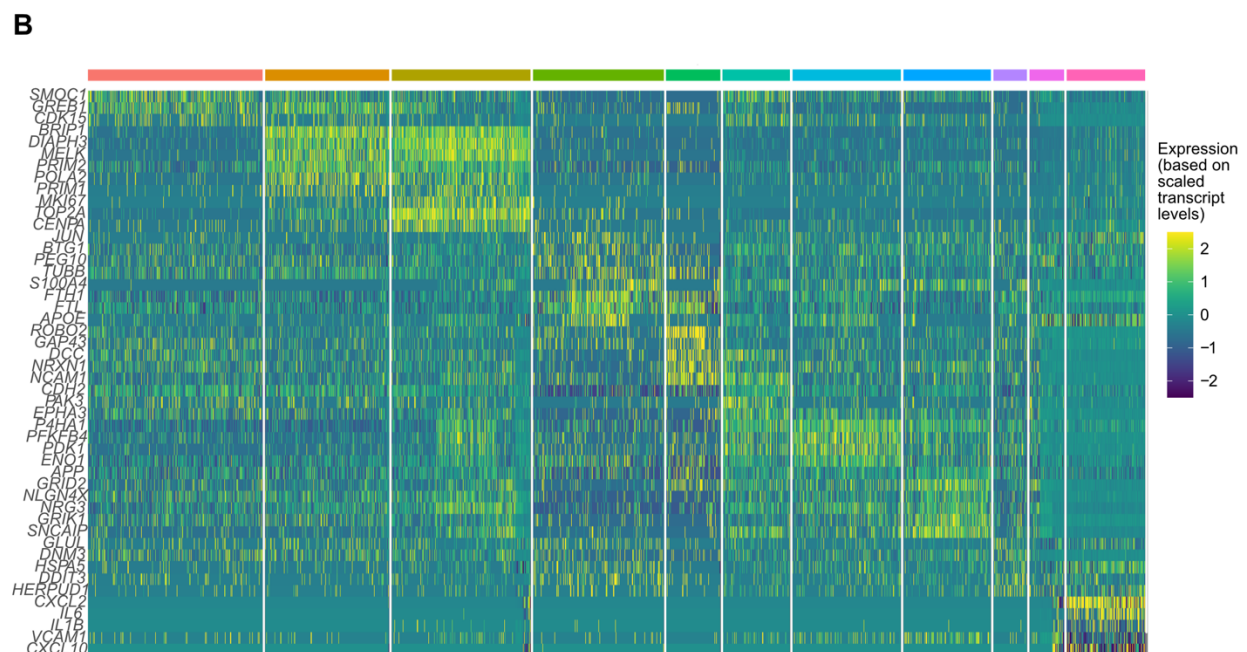
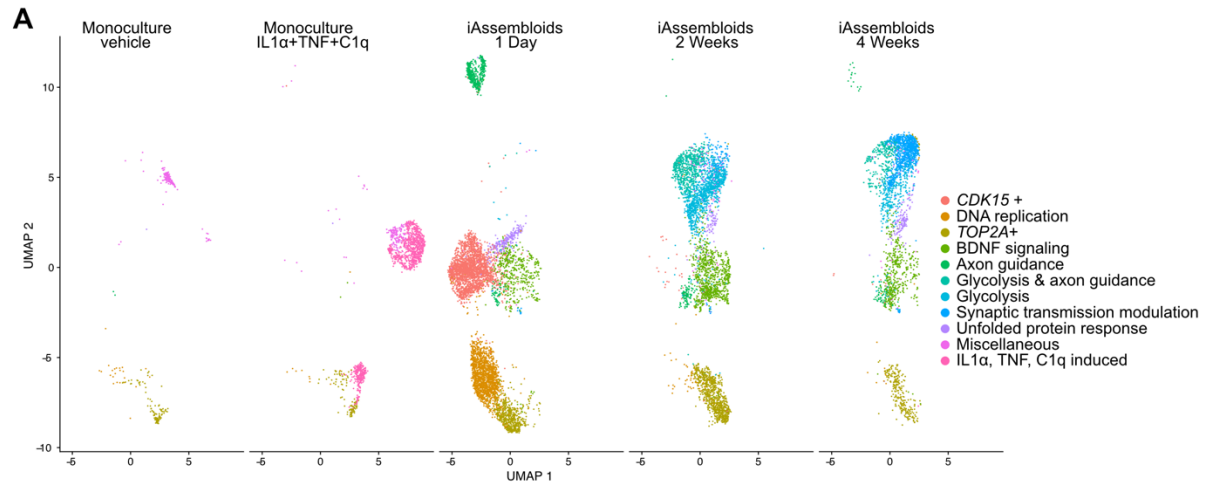


Figure 2.2: Astrocytes in iAssembloids express genes associated with neuronal support

(A) Cells identified as astrocytes from snRNA-seq in iAssembloids (this study) were integrated with data from single cell sequencing of monocultured astrocytes (Leng et al., 2022) using Seurat's data integration pipeline. A UMAP was generated and split by origin of sample and time point. Cluster names were defined by running defined markers through EnrichR. Clusters include a CDK15+ cluster, cells undergoing DNA replication, an immature astrocyte cluster (TOP2A+), cells expressing genes related to BDNF signaling, axon guidance, glycolysis, synaptic transmission modulation, the unfolded protein response, and a cluster that had no specific markers (miscellaneous). The final cluster (IL1 α , TNF, C1q induced) is based on previous studies for cytokine-induced markers⁹.

(B) Heatmap generated based on markers defined in A.

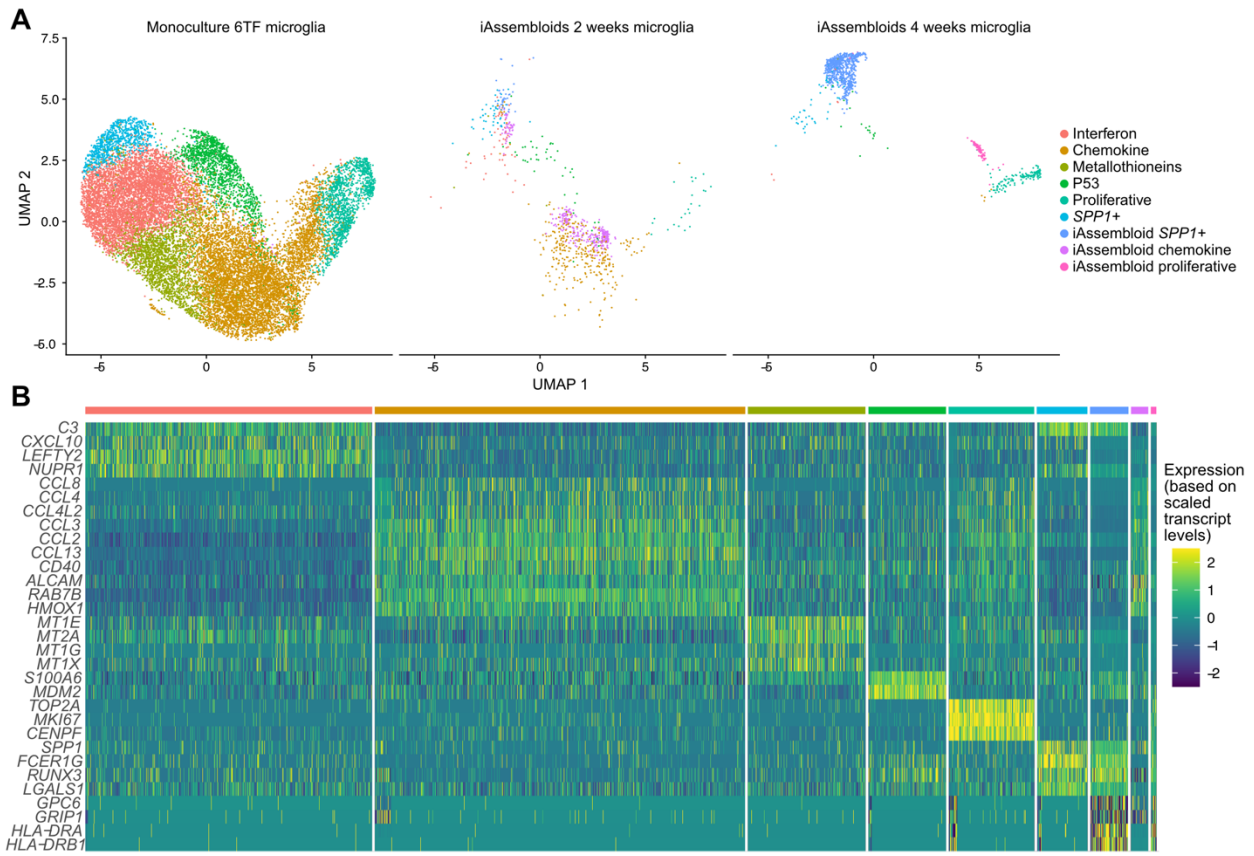


Figure 2.3. Microglia within iAssembloids express MHC Class II members

(A) Cells identified as microglia from snRNA-seq in iAssembloids were integrated with single-cell sequencing of microglia⁸ using Seurat's data integration pipeline. A UMAP was generated and split by origin of sample and the time point. Clusters include those that fall into the interferon, chemokine, metallothioneins, P54, proliferative and SPP1+ cluster. iAssembloid microglia specific clusters (iAssembloid SPP1+, iAssembloid chemokine, and iAssembloid proliferative are also highlighted). Microglia mainly map to existing clusters. However, cells in iAssembloids uniquely express MHC Class II members such as HLA-DRA and HLA-DRB1.

(B) Heatmap was generated based on markers defined in A.

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Figure 3.1. CRISPRi-based screens in iAssembloids reveals genes affecting neuronal survival not found in screens in 2D monoculture.

(A) Schematic of screen design.

(B) Volcano plot of knockdown phenotypes from the survival screen from day 14 iAssembloids. Hit genes (FDR < 0.05) are labeled in either light blue (knockdown decreases survival) or light pink (knockdown increases survival). Selected hit genes were color-coded based on curated functional categories.

(C) Scatterplot of gene scores from our previous survival screen in 2D monocultured neurons⁶ (x-axis) compared to the screen in iAssembloid (y-axis). Both screens used the H1 sgRNA library targeting the “druggable genome.” Selected genes with consistent phenotypes in both screens are labeled in black. Other genes of interest are color-coded based on functional category as in panel B.

(D) Heatmap representing gene scores for neuronal survival from screens in 2D monoculture screens^{6,7}, and from iAssembloids (this study, screens were conducted with the H1 sgRNA library and an sgRNA library targeting neurodegeneration-related genes). Neurodegeneration library hits were compared to gene scores from a genome-wide 2D monoculture screen⁷. Genes are grouped by functional categories as in panels B and C. Genes selected for secondary screens are highlighted by yellow boxes on the left.

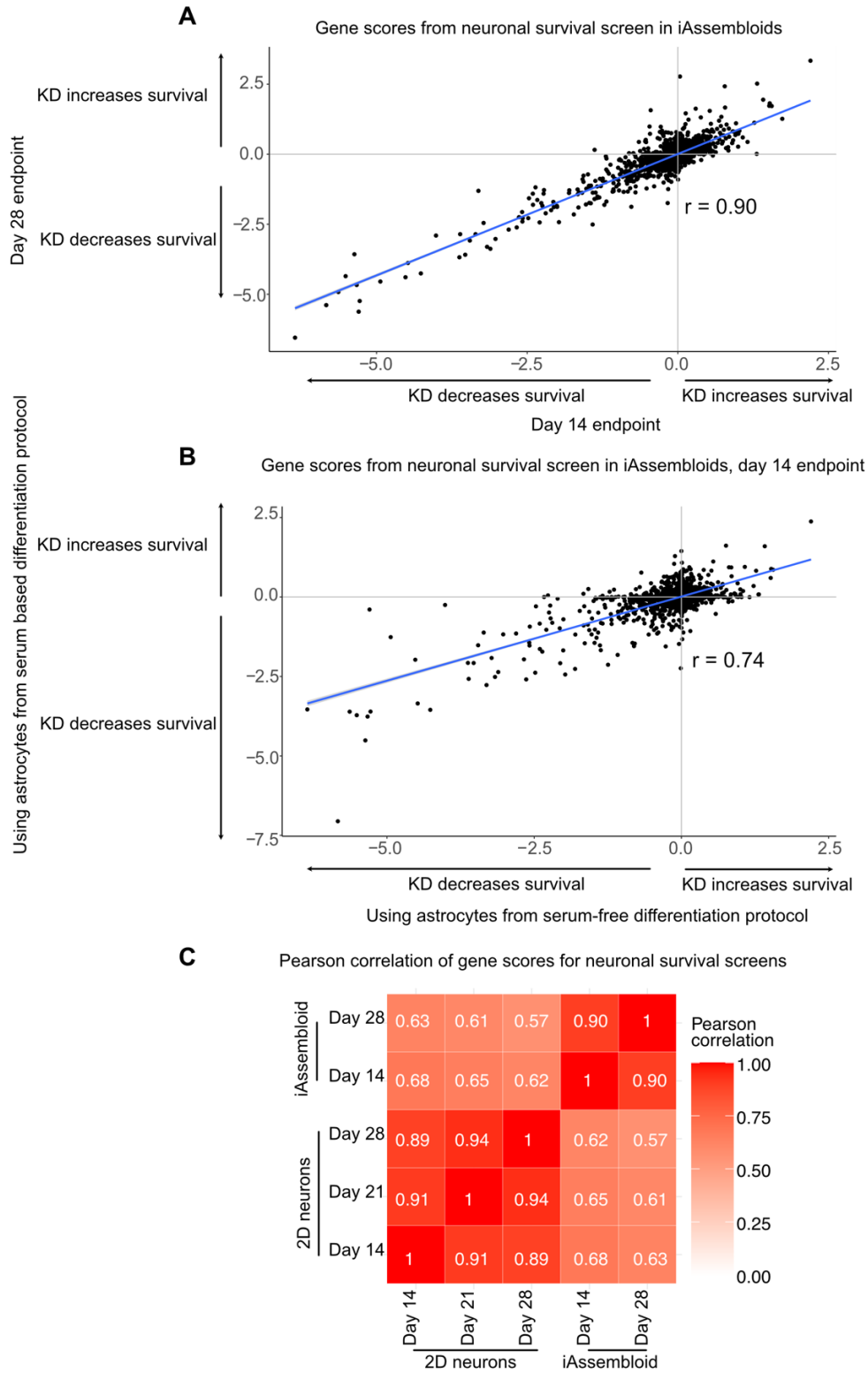


Figure 3.2. CRISPRi-based functional genomics screens in iAssembloids are reproducible

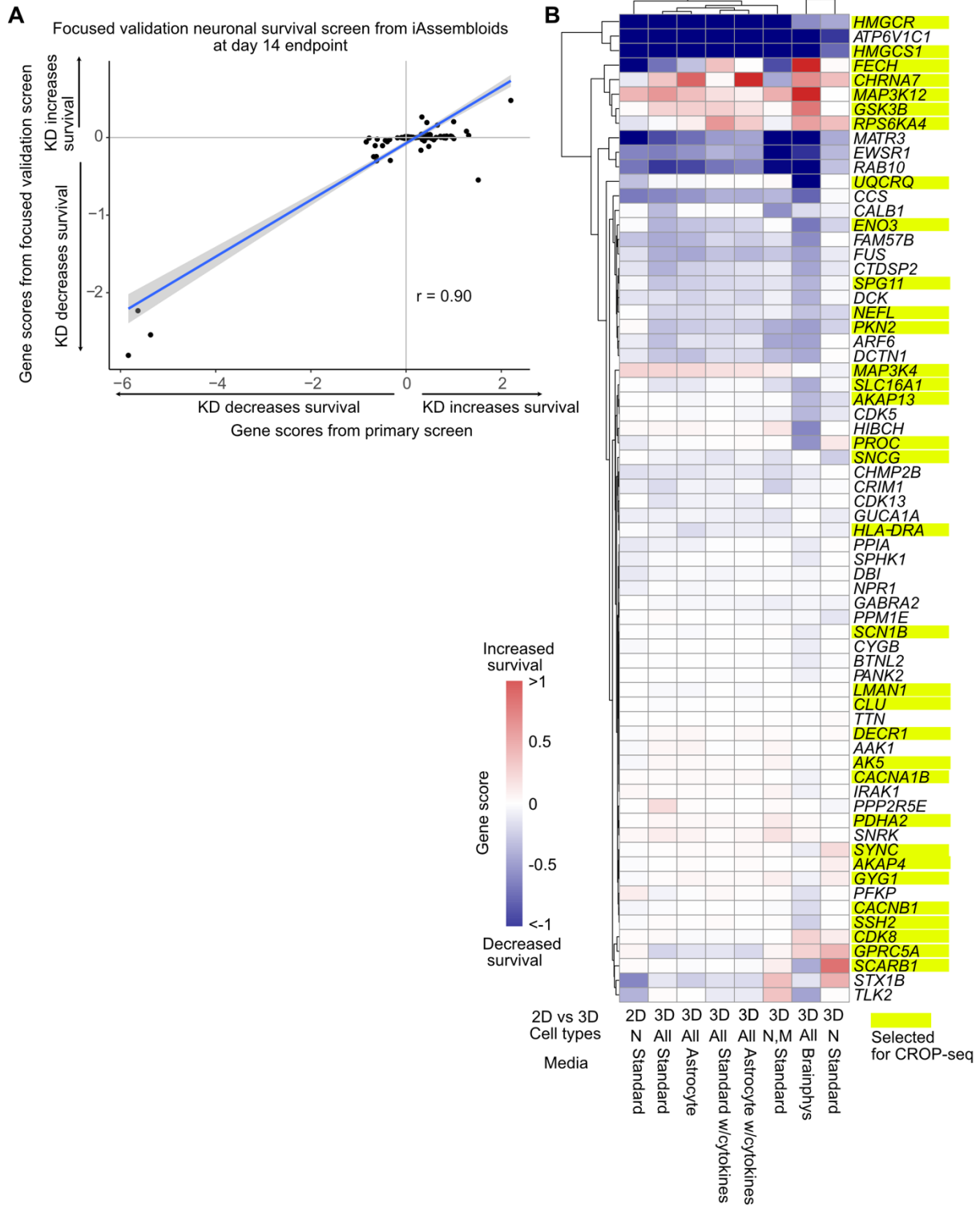
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(A) Scatterplot comparing CRISPRi-based neuronal survival screens at 14 days vs. 28 days post seeding into AggreWell plates.

(B) Scatterplot comparing CRISPRi-based neuronal survival screens using astrocytes from serum-based differentiation protocol^{15,47} vs. the serum-free differentiation protocol^{16,61}.

(C) Correlation heatmap of CRISPRi-based neuronal survival screens in the context of 2D monocultures⁶ vs. iAssembloids (this study).



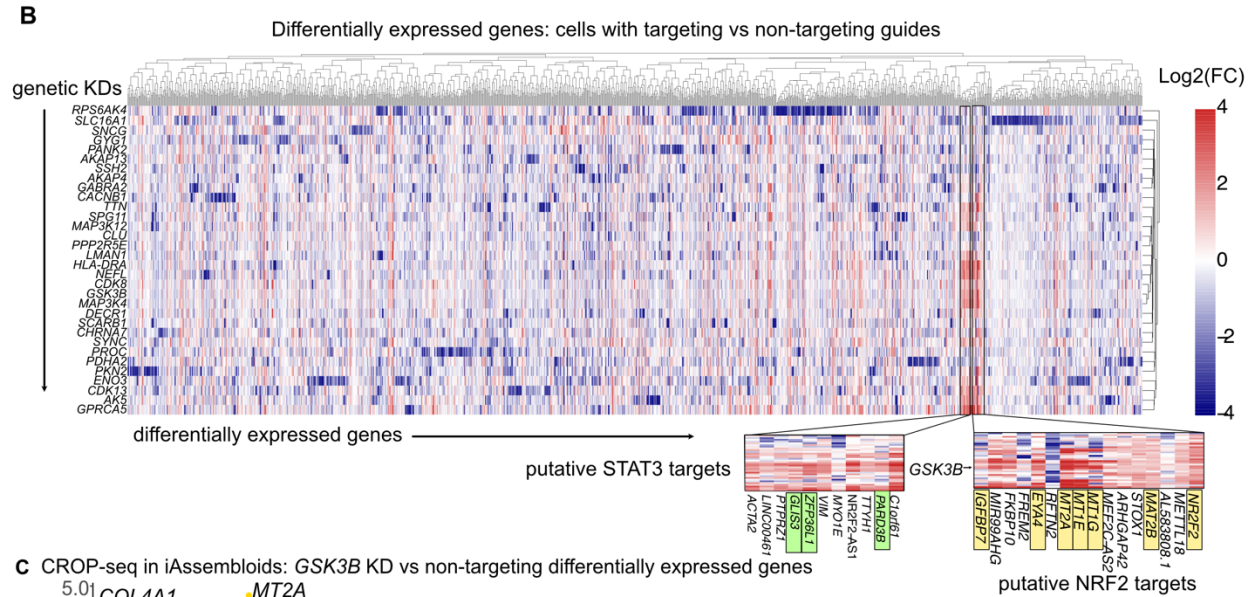
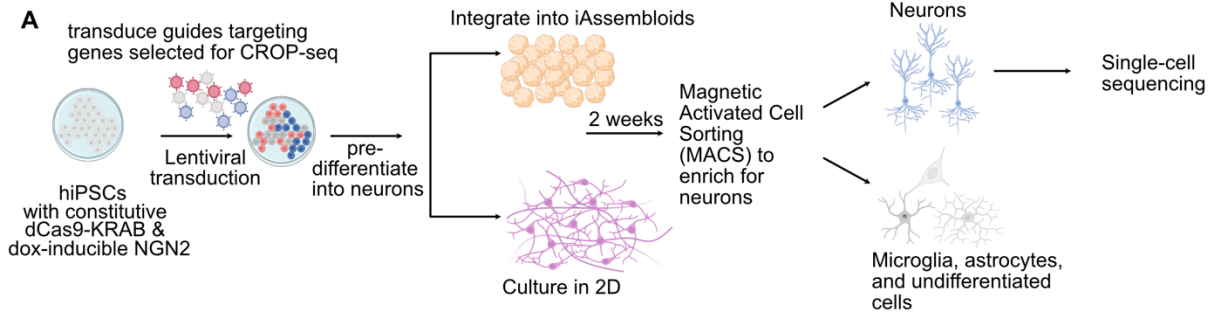
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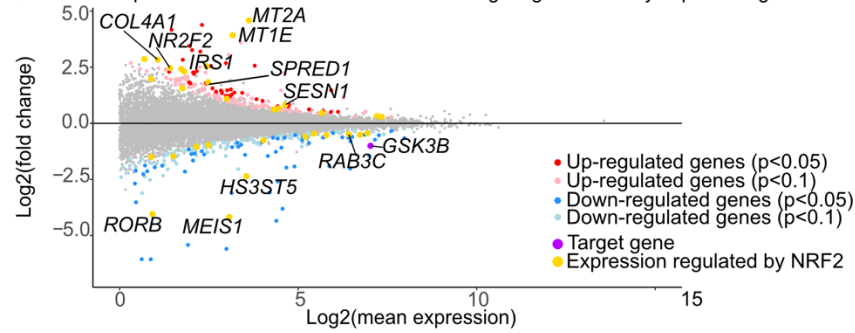
Figure 3.3 Focused validation screens for top hits from the primary screen

(A) Scatterplot of gene scores from the primary iAssembloid screen hits vs the secondary validation screen. Phenotypes from the secondary screen have a narrower dynamic range but are highly correlated with the primary screen ($r = 0.9$).

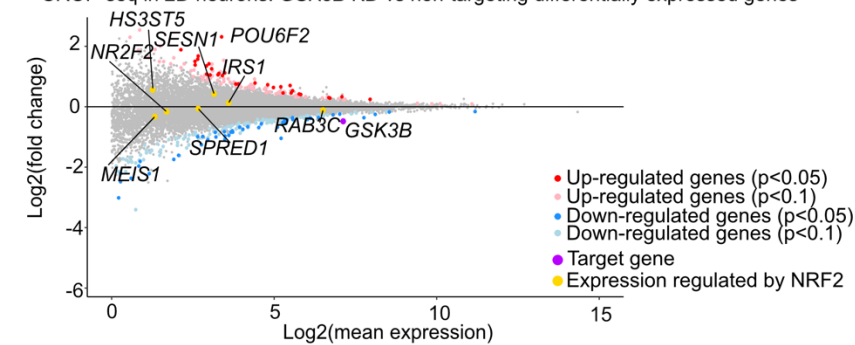
(B) Results from secondary screens comparing iAssembloid culture in various media: normal media (composed of 0.5X astrocyte media and 0.5X microglia base media), normal media plus cytokines (our iAssembloid culture condition), astrocyte media, and astrocytes media plus cytokines, as well as 3D cultures with different cell type compositions: neurons only (N) and neurons plus microglia (N,M).



C CROP-seq in iAssembloids: GSK3B KD vs non-targeting differentially expressed genes



D CROP-seq in 2D neurons: GSK3B KD vs non-targeting differentially expressed genes



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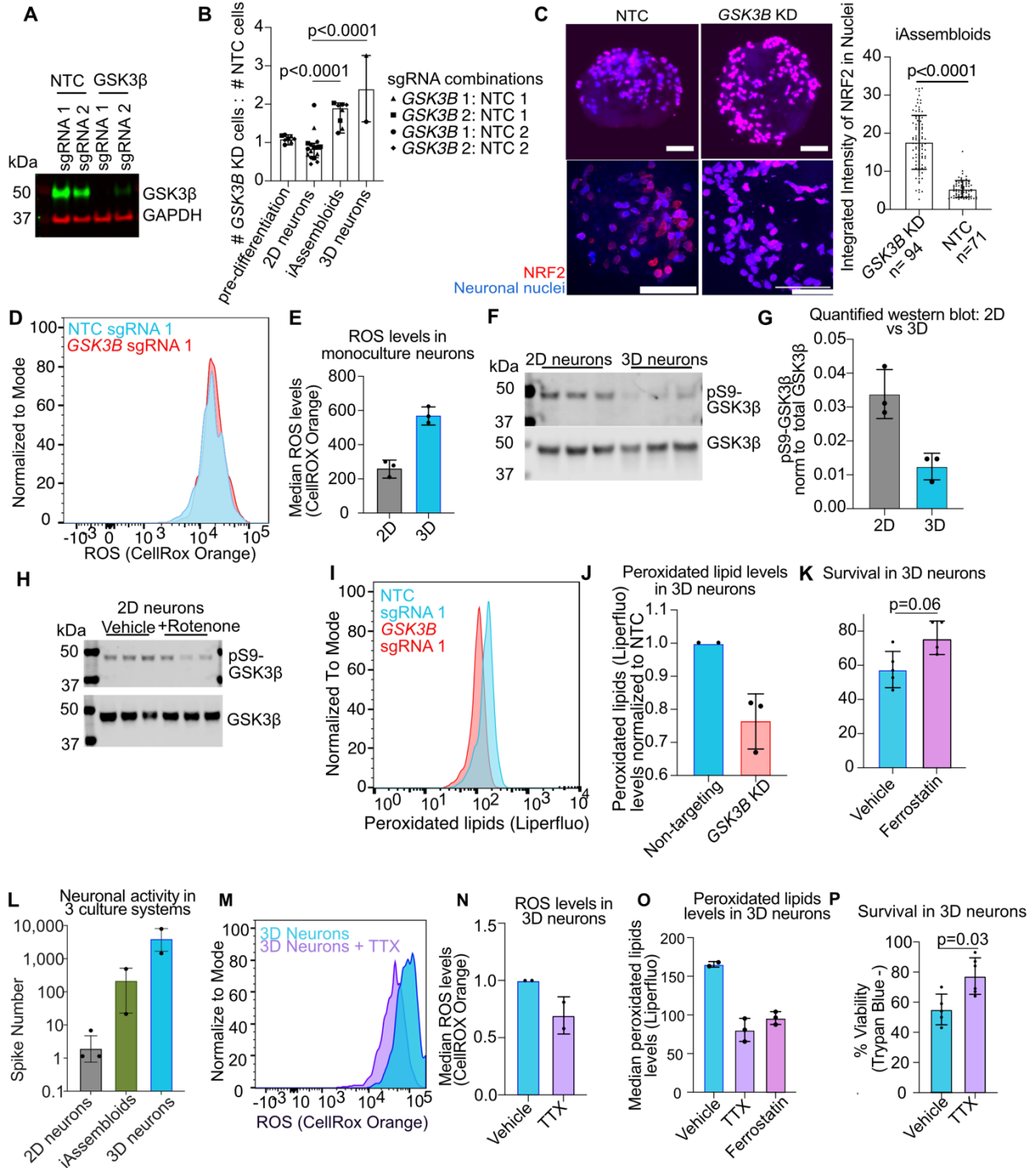
Figure 4. CROP-seq screen reveals that GSK3B knockdown induces neuronal expression of NRF2 target genes in iAssembloids, but not 2D monoculture

(A) Schematic of CROP-seq experimental design.

(B) Heatmap of combined significantly expressed genes ($p < 0.05$) from CROP-seq in neurons cultured in iAssembloids. Rows represent genes that were knocked down in the cells, columns are differentially expressed genes in cells with knockdowns compared to controls. Highlighted are upregulated genes that are putative NRF2 targets (yellow) and STAT3 targets (green).

(C) Differentially expressed genes comparing GSK3B knockdown versus cells containing non-targeting controls in iAssembloids were determined. Red dots represent upregulated genes below the $p < 0.05$ cutoff whereas pink dots represent genes below the $p < 0.1$ cutoff. Dark blue represents downregulated genes meeting $p < 0.05$ cutoff whereas light blue represents downregulated genes below the $p < 0.1$ cutoff. The target gene, GSK3B, is highlighted in purple and genes that have been putatively shown to be regulated by NRF2 are highlighted in yellow. P values were determined using the Wald test.

(D) Differentially expressed genes comparing GSK3B knockdown versus cells containing non-targeting controls in 2D neuronal monoculture were determined as specified in C.



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Figure 5.1 GSK3 β activity prevents NRF2 from protecting neurons from oxidative stress

(A) Western blot validation of GSK3 β knockdown in 3D Neurons. NTC: non-targeting control sgRNA. 3D cultures generated from NTC cells from results for 2 independent non-targeting control (NTC) sgRNAs and 2 independent sgRNAs targeting GSK3B are shown.

(B) Validation of the effect of GSK3 β knockdown on cell survival over 14 days comparing the ratio of surviving cells expressing a non-targeting control sgRNA and GFP versus the proportion of cells expressing an sgRNA targeting GSK3 β and BFP. Experiment was performed with two different NTC sgRNAs as well as two GSK3 β sgRNAs with a total of four different combinations of NTC and GSK3 β sgRNAs. Medians and 95% confidence intervals are represented. ANOVA followed by Tukey's multiple comparison test was conducted to determine significance.

(C) Immunohistochemistry staining for NRF2 (red) in iAssembloids. Neuronal nuclei express BFP (blue). Scale bars = 50 μ m. Images taken at 14 days post seeding of iAssembloids into AggreWell™ 800 plates. Nuclei were identified using BFP in images and the integrated intensity for NRF2 within nuclei was quantified. Images from three different iAssembloids expressing NTC sgRNA 1 (n = 71 cells) and GSK3 β KD sgRNA 1 (n = 94 cells) were taken. Mean with standard deviation is represented. P values were determined using Student's t-test.

(D) Levels of reactive oxygen species (ROS, via CellROX™ orange staining) determined via flow cytometry of neurons expressing a non-targeting control sgRNA or an sgRNA knocking down GSK3 β in 3D monoculture. NTC sgRNA 1 and GSK3 β KD sgRNA 1 were used in this experiment. Neurons with those sgRNAs were co-cultured and sgRNA identity was assigned based on expression of a BFP (GSK3 β sgRNA 1) or mClover (NTC sgRNA 1) marker.

(E) ROS levels (CellROX™ orange staining) comparing 2D monoculture neurons versus 3D monoculture neurons. Each dot represents an independent culture well. Bars represent means of the median fluorescence level and error bars represent standard deviation.

(F, G) Western blot of GSK3 β with the inhibitory phosphorylation at serine 9 (pS9-GSK3 β) and total GSK3 β levels in 2D monoculture neurons versus 3D monoculture neurons. F, western blot image. G, quantification of western blot. Bars represent means of phospho-S9 GSK3B normalized to total GSK3B levels. Error bars represent standard deviation.

(H) Western blot of pS9-GSK3 β levels in 2D monocultured neurons after induction of oxidative stress with rotenone (200 nM for 24 hours). 3 independent wells are shown for each the vehicle and rotenone conditions.

(I, J) Peroxidized lipid levels of GSK3 β KD cells as compared to NTC. Cells were stained with Liperflu and median fluorescence levels were measured. Levels were then normalized to the NTC. H, example trace. I, quantification of 3 replicates (3 cell culture wells), bars represent mean, error bars represent standard deviation.

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(K) Viability of 3D neurons treated with vehicle or with 10 μM ferrostatin. Neurons were treated starting at one week in 3D culture. Cells were stained with trypan blue and the percentage of trypan blue negative cells were obtained after 14 days in 3D culture (total 7 day treatment). Individual points represent one well of 3D neuronal cultures. Bars represent mean, error bars represent standard deviation.

(L) Neuronal activity of 2D neurons, iAssembloids and 3D neurons at day 14 was characterized using multielectrode arrays (MEA). Dots represent the number of spikes from individual MEA wells. Bars represent mean, error bars represent standard deviation.

(M, N) Reactive oxygen species (ROS) levels of 3D neurons treated with tetrodotoxin (1.5 μM , duration of 1 week). Neurons are stained with CellROXTM orange and fluorescence levels were read out using flow cytometry. M, example trace. N. median fluorescence was normalized to the fluorescence level of the vehicle condition to account for different levels of background fluorescence across experiments. Dots represent individual wells. Error bars represent standard deviation.

(O) Peroxidated lipid levels (Liperfluo stain) in 3D monocultured neurons treated with vehicle, tetrodotoxin (1.5 μM , duration of 1 week), and ferrostatin (10 μM , duration of 1 week). Bars represent mean, error bars represent standard deviation. Dots represent cells from 3 independent wells.

(P) Viability of 3D monocultured neurons after 14 days in culture with and without tetrodotoxin (TTX, 1.5 μM , duration of 1 week) treatment. Cells were stained with trypan blue, and the percentage of trypan blue negative cells were obtained. Bars represent mean, error bars represent standard deviation. Dots represent cells from independent wells.

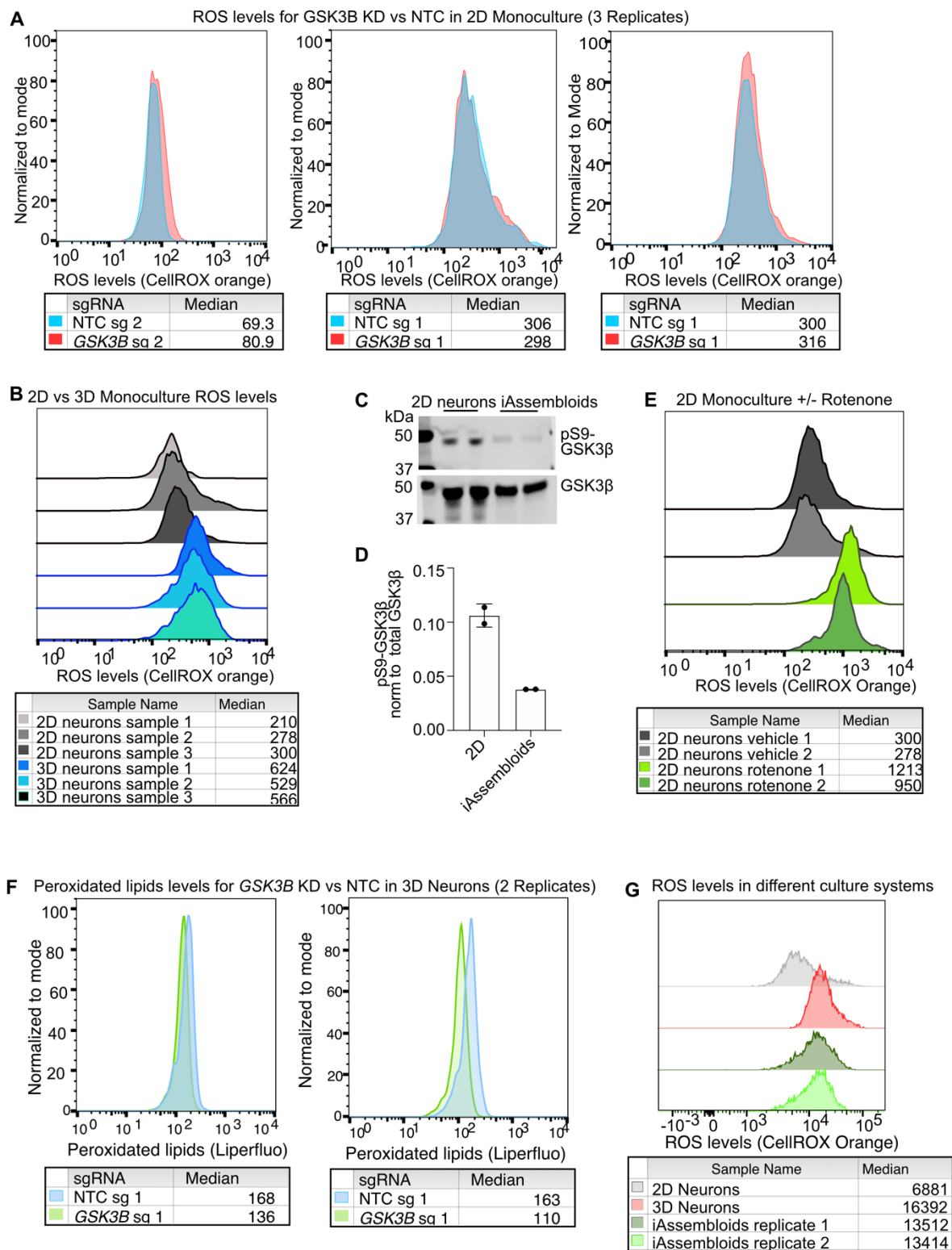


Figure 5.2 Representative flow cytometry results for neuronal phenotypes
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(A) Three replicates of 2D monoculture neurons with and without GSK3 β knockdown stained with CellROXTM orange.

(B) Replicates of 2D vs 3D monocultured neurons stained for ROS levels by CellROXTM orange.

(C, D) Western blot for phospho-S9 GSK3 β and total GSK3 β . Two independent culture wells are shown for each 2D monocultured neurons and iAssembloids. C, blot image. D, quantification of C.

(E) Replicates of 2D monocultured neurons with and without rotenone treatment stained for ROS levels by CellROXTM orange.

(F) Replicates of 3D cultured neurons with and without GSK3 β knockdown stained with Liperfluo to detect peroxidated lipid levels

(G) Neurons from 3 different culture systems (2D monoculture, 3D monoculture, iAssembloids) stained for ROS levels by CellROXTM orange.

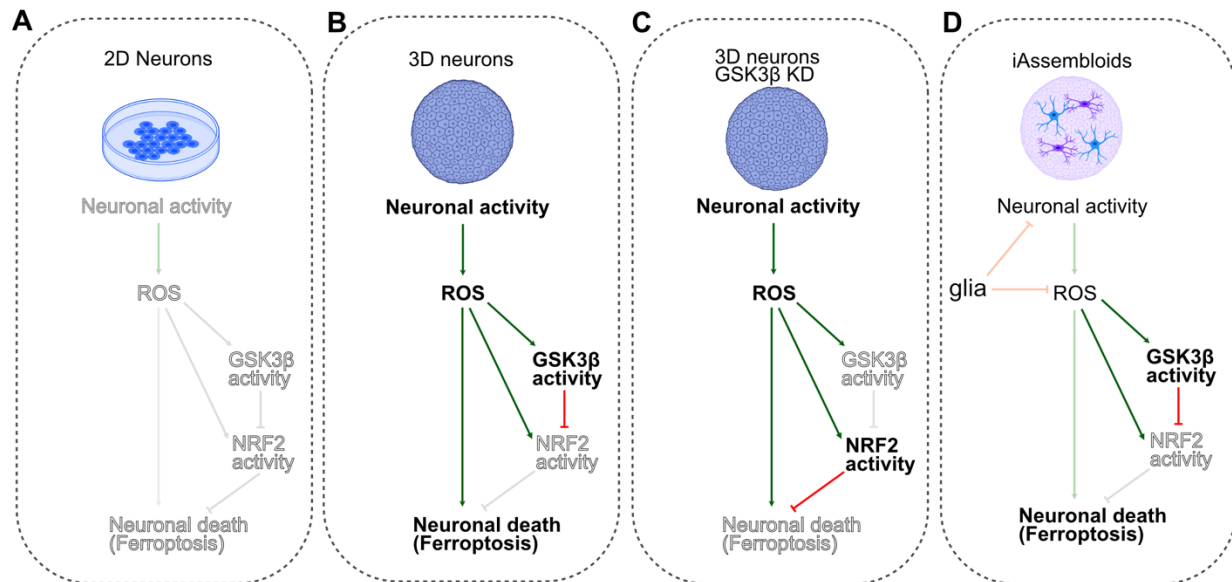


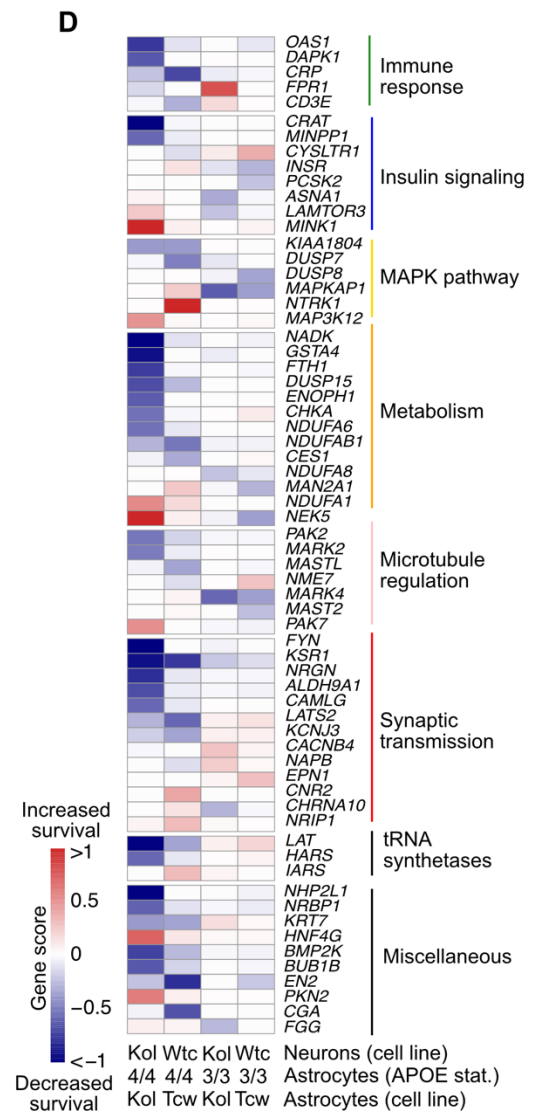
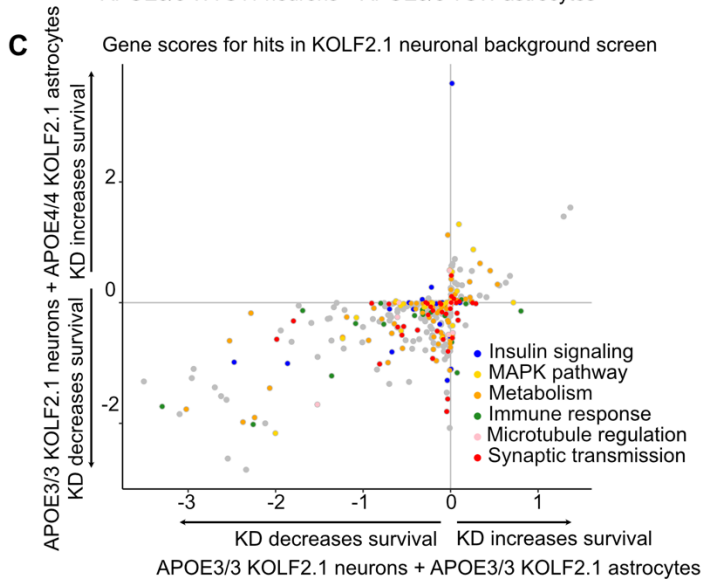
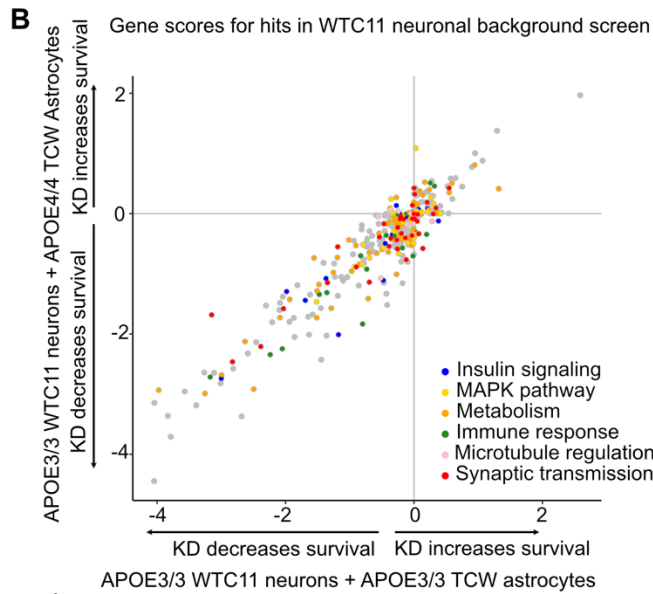
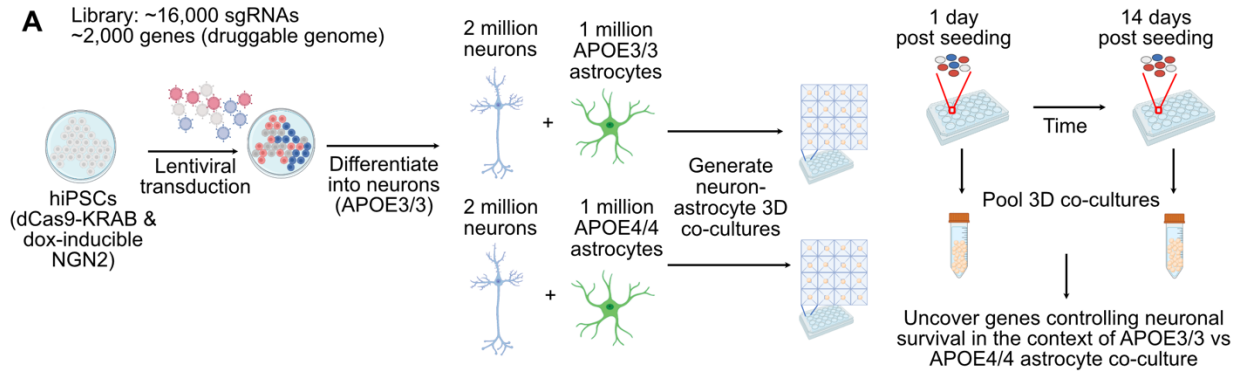
Figure 6. Proposed model of GSK3 β function in oxidative stress and neuronal death

(A) In 2D monocultured neurons, there is not sufficient neuronal activity to drive oxidative stress. Therefore, GSK3 β activity is low and neuronal death is not triggered.

(B) In 3D monocultured neurons, neuronal activity is high. This triggers an increase in ROS and oxidative stress and leads to an increase in GSK3 β activity. Increase of GSK3 β activity blocks NRF2 from translocating to nucleus to mount a protective oxidative stress response. In the absence of the protective response, neurons undergo ferroptosis.

(C) In 3D monocultured neurons with GSK3B knockdown, neuronal activity and oxidative stress is triggered, but in the absence of GSK3 β , NRF2 can translocate to the nucleus, preventing ferroptosis.

(D) In iAssembloids, the addition of glial cells reduces neuronal activity and ROS, possibly by regulation of glutamate levels via glutamate reuptake, which could lessen GSK3 β activity and protect cells against ferroptosis.



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Figure 7. CRISPRi-based screen for neuronal survival in APOE- ϵ 3 (APOE3/3) versus APOE4- ϵ 4 (APOE4/4) astrocyte 3D co-cultures

(A) Schematic of experimental design. Screens for neuronal survival were conducting in APOE3/3 neurons with dCas9-KRAB and dox-inducible CRISPRi that were 3D co-cultured with APOE3/3 vs APOE4/4 astrocytes.

(B) Scatterplot of gene scores from screens using APOE3/3 (x-axis) compared to APOE4/4 (y-axis) astrocytes (TCW 1E33-C, TCW 1E44-C) and APOE3/3 neurons (WTC11). Hits (FDR < 0.05) from screens were included in the scatterplot. Genes belonging to selected functional categories were annotated in different colors; other genes are shown in gray.

(C) Screen as described in B, but both neurons and astrocytes were in the KOLF2.1 background.

(D) Heatmap of hit genes that were consistent between the two screens (WTC11 and KOLF2.1) and had the strongest difference in gene score between APOE3/3 and APOE4/4 astrocyte co-cultures were selected and annotated.

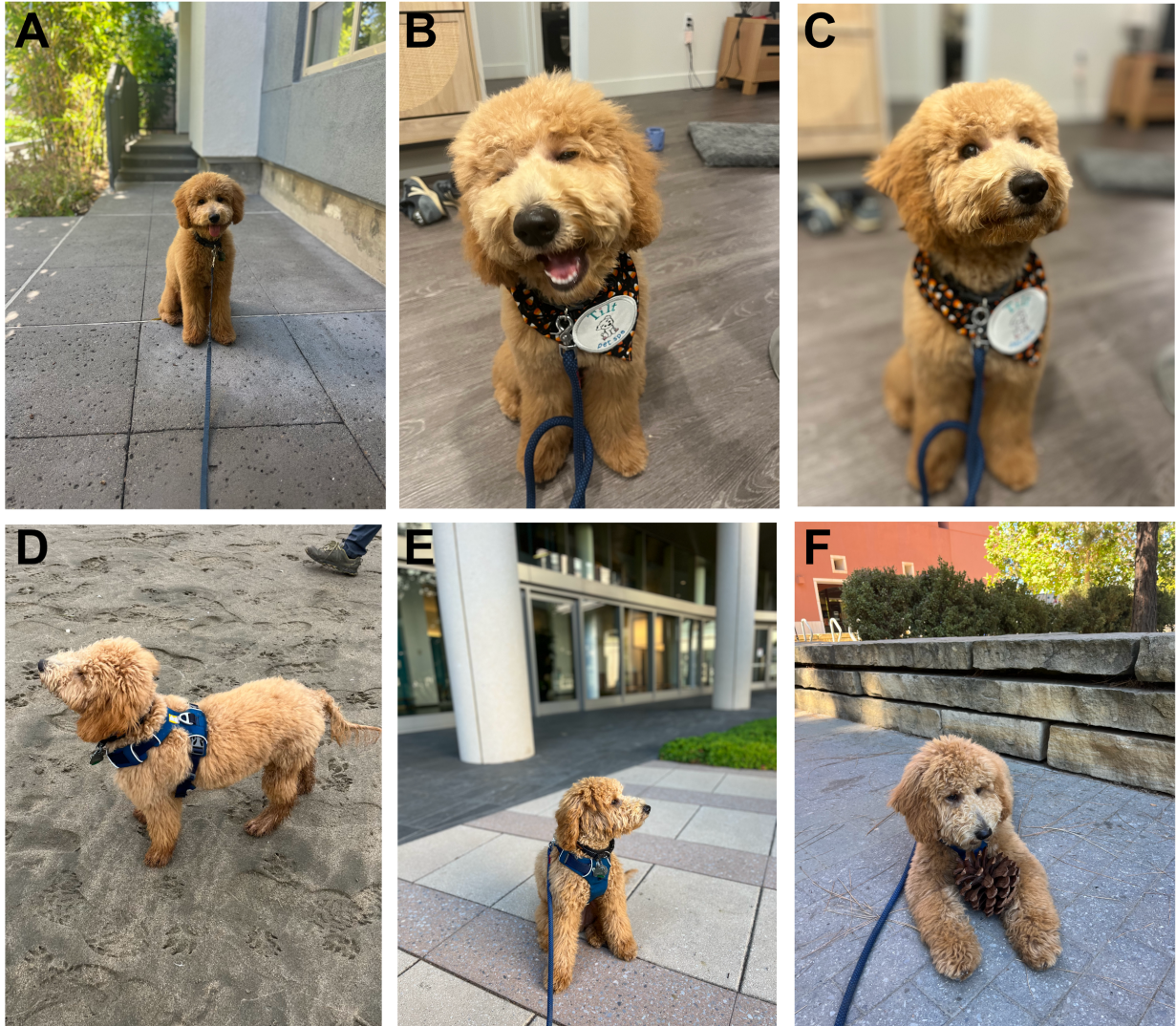


Figure 8. Charlie, the best boy.

(A) Charlie getting ready for a walk.

(B, C) Charlie after his first groom. He looks very dapper.

(D) Charlie sniffing ocean breeze. He is at Fort Funston.

(E, F) Charlie waiting for me outside Sandler for me to finish tissue culture. Such a good, patient boy.

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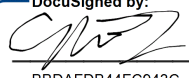
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