Modeling neurodegenerative diseases with cerebral organoids and other three-dimensional culture systems: focus on Alzheimer’s disease

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Abstract
Many neurodegenerative diseases (NDs) such as Alzheimer’s disease, Parkinson’s disease, frontotemporal dementia, amyotrophic lateral sclerosis and Huntington’s disease, are characterized by the progressive accumulation of abnormal proteinaceous assemblies in specific cell types and regions of the brain, leading to cellular dysfunction and brain damage. Although animal- and in vitro-based studies of NDs have provided the field with an extensive understanding of some of the mechanisms underlying these diseases, findings from these studies have not yielded substantial progress in identifying treatment options for patient populations. This necessitates the development of complementary model systems that are better suited to recapitulate human-specific features of ND pathogenesis. Three-dimensional (3D) culture systems, such as cerebral organoids generated from human induced pluripotent stem cells, hold significant potential to model NDs in a complex, tissue-like environment. In this review, we discuss the advantages of 3D culture systems and 3D modeling of NDs, especially AD and FTD. We also provide an overview of the challenges and limitations of the current 3D culture systems. Finally, we propose a few potential future directions in applying state-of-the-art technologies in 3D culture systems to understand the mechanisms of NDs and to accelerate drug discovery.

Keywords Alzheimer’s disease · Neurodegenerative diseases · hiPSCs · 3D culture · cerebral organoids · tau pathology

Introduction
Alzheimer’s disease (AD), Parkinson’s disease (PD), Frontotemporal dementia (FTD), Amyotrophic lateral sclerosis (ALS) and Huntington disease (HD) are devastating neurodegenerative diseases (NDs) that are prevalent worldwide, particularly in the elderly [1–3]. Although NDs are heterogeneous in their clinical manifestations and mechanisms of cellular vulnerabilities, they share two important common features: 1) an increase in prevalence of disease with age, and 2) the aggregation of cytosolic or nuclear proteins within vulnerable cells and brain regions [3–6]. Protein aggregates such as beta-amyloid (Aβ) plaques, neurofibrillary tangles (NFTs) and aggregates of α-synuclein, TAR DNA-binding protein (TDP)-43 and polyglutamine protein are prevalent in these NDs. Furthermore, studies with human brain tissue, animal models and 2D cell cultures have shown that NDs share some common pathological mechanisms that underlie their development and progression, including a defective protein homeostasis network, dysfunctional mitochondrial homeostasis, propagation and seeding of protein aggregates, synaptic toxicity and network dysfunction as well as maladaptive innate immune responses [3, 6, 7].
In this review, we discuss the limitations of animal and 2D cell culture models that 3D culture systems such as reproducible hiPSC-derived 3D cerebral organoids (COs) may address. Specifically, we discuss the potential for COs to model crucial features of AD, including Aβ and tau pathology, oxidative stress and mitochondrial dysfunction, excitotoxicity, neuroinflammation and neuronal network dysfunction. We then examine the limitations as well as the technical progress being made in CO generation to overcome these challenges. Finally, we discuss a few potential future directions in applying state-of-the-art technologies in COs to further improve the efficiency of CO modeling and to understand the underlying disease mechanisms and affected biological pathways and changes in COs in response to various therapeutic drugs (Fig. 1).

COs provide an innovative model of the human brain to understand neurodegenerative disease mechanisms

Although animal models and 2D neuronal cultures have advanced our mechanistic understanding of neurodegeneration, these models have been unable to recapitulate the complexity of human diseases [1, 8]. Thus, the lack of biologically relevant model systems hinder progress in discerning disease mechanisms and the development of therapeutic interventions. COs have diverse progenitor and differentiated cell populations that have a spatial organization resembling the fetal neocortex [9–11]. These cell populations are electrophysiologically active displaying periods of coordinated firing, indicating the presence of neuronal networks [12–14]. Compared to studies with analogous monolayer cultures, COs have gene expression profiles that more closely resemble in vivo conditions [15, 16]. In addition, CO models permit the study of human-specific mechanisms or processes that cannot be easily tested or investigated with animal models. For instance, the developing human brain is comprised of a diversity of progenitor types, including outer radial glia with basal restricted processes that are not represented during mouse corticogenesis [17]. Therefore, model organisms are limited in their ability to recapitulate some disease processes, such as microcephaly [18]. Further, comparative studies of rodent and human mature cortex have demonstrated species divergence in respect to the identities and proportions of neuronal and glial cell types, morphologies, and transcriptomes [19]. These differences suggest that the current understanding of brain pathophysiology, including neurodegenerative disease mechanisms, may be expanded using human-derived models [20].

Despite the advantages of COs, several challenges remain for their use in modeling NDs. Principally, COs exhibit variable morphological and cellular developmental features, which presents challenges in analyzing pathological phenotypes. CO cell type diversity and relative representation greatly vary by protocol type and culture length [21, 22]. However, recent efforts to enhance the reproducibility of COs led to various improved protocols. These protocols yield remarkably similar cell type composition in up to 95% of cultures [12, 23]. Variability between individual organoids is comparable to the variability in individual human brains which warrants further study [23]. In addition, progenitor zone structures can vary in size and organization among organoids, which may ultimately impact relative cell type presence. To address these challenges, advancements in bioreactor designs and differentiation protocols have increased CO reproducibility [11, 12, 23, 24].

3D modeling of AD

Human COs and other 3D cell culture systems exhibit key neuropathological features of NDs as shown in Fig. 2 and Table 1. Here we focus on 3D modeling of AD-associated neuropathology, oxidative stress and mitochondrial dysfunction, excitotoxicity, neuroinflammation as well as synaptic and network dysfunction.

Aβ pathology

AD is characterized by the presence of extracellular Aβ plaques and intracellular NFTs. Familial AD (FAD) is associated with early onset of AD symptoms and is caused by variants in the presenilin-1 (PSEN1), presenilin-2 (PSEN2), or amyloid precursor protein (APP) genes, whereas sporadic AD (SAD) is typically associated with multiple genetic polymorphisms and a later onset of symptoms [25]. APP is the precursor of Aβ, therefore understanding its processing is important in designing therapeutics that reduce Aβ levels. Sequential APP cleavage with α and γ-secretase (a complex composed of four components: PSEN1, nicastrin, anterior pharynx defective 1 (APH-1) and presenilin enhancer 2 (PSEN-2)) leads to a soluble sAPPα fragment under normal conditions [26, 27]. However, APP cleavage via β and γ-secretase results in a sAPPβ fragment and Aβ peptide, the primary component of amyloid plaques in AD [26, 27]. Several human clinical trials for AD have focused on inhibition of β and γ-secretase to decrease Aβ production [28–34]. Although these trials showed a decrease in plasma and CSF Aβ levels, they have failed when it comes to establishing safety and efficacy. These trials have been discontinued based on increased incidence of skin cancer and worsening cognitive scores [29–34].

3D cell culture systems overexpressing one or more FAD variants have been used to understand AD pathology [35–37]. Either a thin-layer (100-200 μm) or a thick-layer (~4 mm) 3D culture with Matrigel matrix was developed by introducing viral vectors with FAD variants (Swedish and London
APPSL and PSEN1Δ9 variants) into neural progenitor cells (ReN) [36]. High Aβ42/40 ratios were detected with the presence of SDS-resistant oligomeric Aβ [37] resulting in robust Aβ aggregate deposits (6-8 weeks) [35–37]. 3D cultures with variants (I45F and I47F) in the APP transmembrane domain (TMD), but without the overexpression of the PSEN1Δ9 variant, showed either higher (I45F) or lower (I47F) Aβ42/40 ratio compared to wild-type (WT) control cultures [37]. Similar to human clinical trials discussed above, application of β and γ-secretase inhibitors in these 3D cultures showed a significant decrease in Aβ pathology [35]. Recent studies indicate that these enzymes have multiple substrates and have several important physiological functions outside the brain including muscle spindle formation [38–40]. Failure of prior clinical trials can potentially be overcome by testing therapies aimed at brain-specific inhibition of β and γ-secretase to minimize off-target effects. CO model provides an excellent platform to initially validate such therapies.

3D cultures have also been developed with modified hydrogel-based scaffolds [41, 42], such as StarPEG-heparin hydrogel that promotes proliferation of neural stem cells [41]. Upon Aβ42 treatment, the spontaneous electrical activity of these cultures was disrupted, resulting in dystrophic axons, matrix stiffness and decrease in synaptic plasticity [41].

![Fig. 1 Three-dimensional (3D) modeling of neurodegenerative diseases. The overview of limitations, current progress and future direction in using human induced pluripotent stem cells (hiPSCs)-derived 3D cultures/ cerebral organoids (COs) to model Alzheimer’s disease and other neurodegenerative diseases (created with BioRender.com).]
| Disease                               | Origin/Cell line | Model | Pathology                                                                                                                                                                                                 | Reference |
|---------------------------------------|------------------|-------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------|
| Parkinson’s Disease (PD)              | Healthy hNESCs   | 3D microfluidic bioreactor - matrigel embedded hNESCs with medium flow on one/either side of chamber | Mature neurons with bipolar morphology, larger neurites, branching, interconnectivity and synchronized electrical activity. Only 19% of neurons express tyrosine hydroxylase indicative of dopaminergic neurons | Moreno et al., 2015 [183] |
|                                       | Healthy hPSCs    | Seeding of dissociated 2D hPSCs grown into mebiol gel | Development of floorplate-derived substantia nigra-specific midbrain dopaminergic neurons with high tyrosine hydroxylase levels. Electrophysiologically active | Adil et al, 2017 [184] |
|                                       | hNESCs with LRRK2 G2019S variant | Midbrain organoids | Clusters of dopaminergic neurons with high neurite myelination. Presence of astrocytes and oligodendrocytes | Monzel et al, 2017 [185] |
|                                       | hiPSCs with LRRK2 G2019S variant | Neuroepithelial spheres | Transcriptomic profiles show alterations in synaptic transmission (synaptic vesicle trafficking), axon guidance, toll-like receptor signaling pathway and TGF-β signaling pathway | Son et al, 2017 [186] |
|                                       | PD hiPSCs with Q456X and I368N variants in PINK1 | Cerebral organoids | Decrease in dopaminergic neurons, neuronal loss. Decrease in differentiation efficiency with increase in proliferation. Altered electrical activity with decrease in mitophagy. HP-β-CD treatment increase differentiation efficiency | Jarazo et al, 2019 [187] |
|                                       | hiPSCs with LRRK2 G2019S variant | Midbrain organoids | α-synuclein aggregate pathology with defective clearance. Increase in TXNIP levels involved in LRRK2-associated PD | Kim et al, 2019 [188] |
|                                       | hiPSCs derived midbrain floor plate NPCs with LRRK2 G2019S variant | Midbrain organoids | Decrease in the number and complexity of dopaminergic neurons. Increase in FOXA2 levels indicating neurodevelopmental defect | Smits et al, 2019 [189] |
|                                       | hiPSCs from two idiopathic PD patients | Midbrain organoids | Defective neurodevelopment with dopaminergic neuronal loss and reduction in organoid growth. Low levels of midbrain floorplate progenitors FOXA2 and LMX1A and high levels of PTX3 gene involved in dopaminergic neurodevelopment and survival | Chlebanowska et al, 2020 [190] |
|                                       | Single cell dissociated from hESCs | Midbrain organoids | Presence of dopaminergic neurons and glial cells with dopamine and neuromelanin-like granules. MPTP treatment leads to astrocyte mediated dopaminergic neuronal loss. Electrophysiological activity | Kwak et al, 2020 [191] |
| Frontotemporal Dementia (FTD)         | hiPSCs with tau P301L and tau P301LΔp35KI | Cerebral organoids | Higher p25/p35 protein ratios. Decrease in p25, pTau-T181, S202 and total tau in P301LΔ p35KI COs. Inhibition of p25/Cdk5 reduces tau phosphorylation and pathology and increases synaptic density | Seo et al, 2017 [59] |
|                                       | hiPSCs with R406W variant | Cortical organoids | Tau hyperphosphorylation and mislocalization to dendrites. Mitochondrial destabilization and axonal transport disruption. Decrease in phosphorylation by GSK3β, Rho-associated protein kinase (RhoK) and protein kinase A (PKA) at T181, S404, S409 sites | Nakamura et al, 2019 [192] |
| Amyotrophic Lateral Sclerosis (ALS)   | Human chorion derived mesenchymal stem cells | 3D culture - cross-linked electrospun gelatin scaffolds | Shh and RA induction upregulate motor neuron proteins such as islet-1, choline acetyl transferase, nestin, NEFH and transcription factor Hb-9 responsible for neuronal differentiation. Decrease in NEFH after a week | Faghihi et al, 2016 [193] |
| Motor Neuron Disease (MND)            | hiPSCs derived spinal motor neuron spheroid | Microdevice-chamber for MN spheroid, axon fascicle microchannel and chamber for axon terminals | Strong electrical activity and axonal staining. Hydrogen peroxide treatment mimics neurodegeneration resulting in physical changes of the axon fascicle such as reduced/damaged microtubules, changes in directionality, texture, and elasticity | Kawada et al, 2017 [194] |
|                                       | NSCs and skeletal myoblast from sALS hiPSCs | Microdevice-MN spheroid, muscle tissue, and neurite elongation chamber | Low muscle contractions with increase in motor neuron degradation and apoptosis. Bosutinib and rapamycin treatment leads to increase in autophagy and TDP-43 degradation | Osaki et al, 2018 [195] |
Treatment with IL-4 suppressed this kynurenine aminotransferase-mediated Aβ toxicity [41]. Another model used a silk fibroin porous scaffold with intercalated hydrogel to culture neurons derived from a SAD patient [42]. This culture demonstrated a decrease in spontaneous electrical activity and an increase in Aβ42/40 ratio [42].

In contrast, AD-like CO models are designed to grow in a self-organizing manner [43]. FAD hiPSCs with either APP duplication or variants in PSEN1 (M146I and A264E) and PSEN2 (N141I) have been used to generate such models. These COs exhibited increases in intracellular and extracellular Aβ aggregates consistent with the neuropathology of AD [43, 44] (published as a preprint in bioRxiv). These characteristics have also been observed in Down Syndrome COs [45, 46]. Patients with Down Syndrome typically present AD pathology very early in life and experience a rapid advancement of the disease progress [47]. This has been attributed to the overexpression of APP on chromosome 21 [47]. Several studies have shown that Down Syndrome and AD-like COs have high levels of diffuse and fibrillar extracellular Aβ deposits as well as increased cell death [43, 45, 46]. As expected, a dramatic decrease in Aβ aggregation was observed when Down Syndrome and AD-like COs were treated with β and γ-secretase inhibitors [43, 45].

Although overexpression of mutant APP, PSEN1 and PSEN2 genes in COs readily models FAD pathology, these variants restrict modeling of SAD pathology. However, healthy COs treated with Aftin-5, an Aβ-42 inducer, showed time-dependent increases in both Aβ40 and Aβ42 levels, leading to the formation of Aβ oligomers [48]. This model is able to demonstrate physiological interactions, colocalization and parallel expression of APP/Aβ and their high-affinity receptor, cellular prion protein (PrPc), during early phases, indicating a potential role for these proteins in neurodevelopment [48]. This type of chemical induction shows that environmental and other sporadic causes of AD can also be studied in vitro in CO models.

**Tau pathology**

Tauopathies are a spectrum of NDs characterized by abnormal tau hyperphosphorylation and aggregation in the brain. Despite variants identified in the MAPT gene resulting in the formation of paired helical filaments and NFTs in FTD, these variants have not been identified in other tauopathies including AD [37]. Different tauopathies are characterized by the presence of unique, conformational tau strains capable of propagating via anatomically connected brain regions [6]. Here, we review 3D cultures and COs that have been used to model tau hyperphosphorylation, aggregation and its mislocalization, especially in AD and FTD.
In AD-like 3D cultures, high Aβ42/40 ratios correlate with high levels of pTau (S396/A40, T202/S204, T181, S199/S202/T205), indicating that the Aβ42/40 ratio drives tau pathology [35–37]. Similarly, 3D cultures with variants in the APP transmembrane region showed either high (I45F variant) or low pTau levels (I47F variant) correlating with Aβ42/40 ratios [37]. Aβ42 treatment in starPEG-heparin hydrogel-based 3D culture and a silk fibroin-based 3D culture of SAD hiPSC-derived neurons also showed an increase in pTau (T231) [41] (S199) [42] levels with Gallyas- and Thioflavin S-positive NFTs [41]. Treatment of these cultures with β and γ-secretase inhibitors caused a decrease in both Aβ and tau pathology [35].

Apart from 3D cultures, COs that overexpress FAD variants exhibit increased levels of pTau-S396, pTau-T181 and Thioflavin S-positive NFTs [43]. Treatment of the AD-like COs with β and γ-secretase inhibitors reduces Aβ and tau hyperphosphorylation, reiterating that the accumulation of toxic Aβ drives tau pathology [43, 45]. COs have also been used to explore post-translational modifications that may affect the interaction of tau with the molecular chaperones that are responsible for its degradation [49]. For instance, histone deacetylase 6 (HDAC6) is responsible for the deacetylation of α-tubulin and tau [50]. HDAC6 levels are elevated in AD and may be involved in the degradation of tau based on HDAC6 involvement in the ubiquitin-proteasome system and autophagy pathway [50]. As expected, decreased levels of acetylated α-tubulin and elevated levels of HDAC6 were observed in AD-like COs with pathological tau aggregates compared to control COs [49]. CKD-504, a blood-brain barrier (BBB) penetrating HDAC6 inhibitor, was tested on COs generated from hiPSCs with FAD variants. Total tau and pTau levels were decreased in AD-like COs treated with CKD-504, NFT-like deposits (pTau at S202/T205, S396/S404) [46]. Treatment of the AD-like COs with β and γ-secretase inhibitors reduces Aβ and tau hyperphosphorylation, reiterating that the accumulation of toxic Aβ drives tau pathology [43, 45].
indicating that this inhibitor has the potential to rescue tau pathology [49]. CKD-504 reduced tau levels in the presence of a lysosomal inhibitor but did not degrade tau in the presence of a proteasomal inhibitor, indicating that tau is degraded via a proteasomal pathway. CKD-504 also promoted interactions between tau and chaperones by increasing their acetylation, which further led to proteasomal degradation of tau [49].

In another CO model, EBV-based plasmids were used to introduce long-term expression of WT tau or mutant P301S tau in hiPSCs. This novel plasmid was shown to be stable for more than 30 passages and did not affect the differentiation of the COs [51]. Unlike overexpression of genes via retroviral approaches, these vectors do not integrate into the genome but remain extrachromosomal and maintain low copy numbers. Although hyperphosphorylated tau (S202/T205) is observed, this model does not exhibit tau aggregation, which is likely due to the requirement of a longer time for disease development [51].

**FTD**

FTD is the second leading cause of early-onset dementia and is characterized by a predominant tau, TDP-43 or Fused-in-Sarcoma pathology [5, 6]. Here, we focus on 3D cultures and COs derived from FTD patient-derived hiPSCs with MAPT variants that lead to hyperphosphorylation and aggregation of tau.

Alternative splicing of human MAPT in the adult brain results in six tau isoforms [52, 53]. Due to the lack of an aging phenotype in hiPSC-derived cultures, expressing all six adult tau isoforms is a challenge [15, 54–56]. This has been partially overcome in a 3D culture model with Matrigel-coated alginate capsules [57]. When BrainPhys medium was used with this model, cortical neurons, astrocytes and oligodendrocytes developed by 15 weeks of culture, whereas they were not all present when standard medium was used even at 25 weeks of culture [57]. All six tau isoforms were expressed when BrainPhys media was used as compared to only five isoforms with standard media. Despite using BrainPhys media, the 2N3R and 2N4R isoforms in this model account for only 1% and 0.9% of their prevalence in the adult human brain, respectively [57]. Further studies are required to demonstrate similar ratios of tau isoforms as are expressed in the adult human brain [52, 53]. Another Matrigel-based 3D culture model was characterized by the presence of 3R tau and the absence of 4R tau [55]. It was demonstrated that 3R tau was less prone to tau aggregation [52] and did not show any pathology when seeded with truncated tau K18 fibrils [55]. However, when mutant tau P301L was overexpressed in the culture, seeding with K18 fibrils resulted in NFT-like inclusions evidenced by the presence of pTau (S202/T205) [55].

Apart from Matrigel-based 3D cultures, COs have also been used to explore the mechanisms underlying FTD [58, 59]. In a recent study, COs derived from FTD hiPSCs with mutant R406W MAPT were dissociated into cortical neurons [58]. To better understand the progression of tau pathology and its phosphorylation, the three major kinases responsible for tau phosphorylation, glycogen synthase kinase 3 β (GSK3β), Rho-associated protein kinase, and protein kinase A were investigated. R406W variant cortical neurons showed a decrease in tau phosphorylation by GSK3β, Rho-associated Protein Kinase and Protein Kinase A at the T181, S404 and S409 sites of tau, respectively. In particular, inhibition of GSK3β led to an overall increase in tau phosphorylation [58]. Phosphorylation of β-catenin, another substrate of GSK3β, remained unchanged as did phosphatase activity, which can also regulate tau phosphorylation. These results indicate that GSK3β is mainly responsible for the reduced phosphorylation of tau in the R406W variant [58]. Although mutant tau hyperphosphorylation is increased in mouse models and in humans at late stages of the disease, this study indicates that the reduced phosphorylation of tau observed initially is possibly due to the conformational change of tau caused by the variant [58]. Tau mislocalization to the dendrites, mitochondrial destabilization, and disruption of axonal transport are also observed in this model [58]. Although tau hyperphosphorylation occurs ten days after dissociation, further work is required to investigate cellular phenotypes at later time points to further understand the role of the R406W variant in tau pathology and neurodegeneration [58].

To further understand the mechanisms underlying FTD, the role of p25/cyclin-dependent kinase 5 (CDK5) in tau accumulation and the effect of p25 generation on hyperphosphorylated tau has been examined with tau P301L and P301L:Δp35KI isogenic control COs [59]. Cdk5 plays a role in neuronal migration, synapse development as well as synaptic plasticity, and is associated with the regulatory activator p35. High levels of p35 are regulated by either proteasomal degradation or truncation to a soluble 25 kDa form (p25) by calpain [60]. While P301L mutant organoids showed higher p25/p35 protein ratios, P301L: Δp35KI organoids showed significant reductions of p35 and pTau at T181/S205 sites. This finding indicates that p25/Cdk5 may lead to tau hyperphosphorylation that further increases p25 levels and Cdk5 hyperactivation, forming a feed-forward loop [59]. This model demonstrated that inhibition of p25/Cdk5 successfully reduces tau phosphorylation and the pathology associated with FTD [59], which is consistent with previous reports [61]. Further studies are required to understand if kinases like GSK3β are involved in p25/Cdk5-mediated tau hyperphosphorylation. GSK3β inhibitors, such as Tideglusib, have been explored in human clinical trials not only for FTD [62, 63], but also for AD [64, 65]. Although safety was established, Tideglusib did not produce a significant difference in cognitive scores between treatment and placebo groups. Since these clinical trials were carried out in a small group (n=406) and for a short duration (n=26 weeks) [65],...
further dose dependent studies are required in both 3D models and in humans to understand its potential as a therapeutic for AD.

Other pathologies

Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) causes severe upper respiratory tract infection and has also been shown to cause neurological symptoms [66]. Two recent studies have explored the neurological effects of this virus in COs [67, 68] (published as preprint in bioRxiv). Day 15 and Day 60 COs were exposed to SARS-CoV-2 [67]. Relatively mature neurons in day 60 COs exhibited substantial SARS-CoV-2 infection in cortical neurons within two days of exposure compared to Day 15 COs [67]. Similar viral load between two- and four-day post infection in COs indicated that the virus did not actively replicate in the COs [67]. Interestingly, the infected COs also showed an increase in early neurodegeneration-like phenotype as pTau (T231) was missorted from the axon to the soma and was colocalized with SARS-CoV-2 [67]. Furthermore, the virus-mediated early tau pathology in COs was shown to be associated with apparent neuronal death [67].

In the second study, SARS-CoV2 infection in healthy nine-week-old COs displayed a significant increase in viral titers from 24 to 96 hours post infection [68]. With an increase in viral titer, an increase in neuronal loss surrounding the infected regions was observed. Single-cell RNA-Seq (scRNA-Seq) of infected COs indicated that genes related to cell division, organelle fission and metabolic processes were enriched following infection [68]. These data also indicated that while SARS-CoV-2-positive cells exhibited upregulation of genes related to viral transcription and hyperoxia, the surrounding SARS-CoV-2-negative cells also displayed upregulated genes related to hypoxia and cell death [68]. This indicates that SARS-CoV-2 has the ability to manipulate host metabolic processes, which may elicit features of neurodegeneration. Thus, these SARS-CoV-2 CO models might be used to study the neuropathology associated with such viruses, and mechanisms underlying accumulation and mis-sorting of tau in different tauopathies including SAD.

Another alternative is to use Traumatic brain injury (TBI) models to study the role of tau in neurodegeneration since it is considered a major risk factor for NDs [69]. A recent study published as a preprint in bioRxiv describes a TBI CO model [70]. In this study, a high intensity ultrasound with a frequency of 510 kHz was applied to healthy COs in 10 ms bursts for a duration of 2 min to mimic TBI [70]. These COs showed an increase in astrogliosis accompanied by neurodegeneration after the injury [70]. Total RNA-Seq identified changes in genes related to glycolysis, AMPK signaling and protein homeostasis [70]. There was also a significant increase in pTau (T231), pTDP-43 (S409/410) and intracellular calcium levels [70]. Previous studies in FTD and other NDs have identified dysregulation of nuclear pore proteins which can form pathological aggregates with tau and TDP-43 [71]. Similar disruption of the nuclear pore protein, Nup98, was observed in this TBI CO model [70]. Thus, this model can be used to study ND mechanisms involving nuclear pore protein dysregulation, TDP-43, and tau phosphorylation.

Oxidative stress and mitochondrial dysfunction

Generation of reactive oxygen species (ROS) and mitochondrial dysfunction are early features of AD that lead to an increase in cell toxicity and neuronal loss [72]. These signatures have been readily demonstrated in comparative proteomic analyses of AD neuro-spheroids and post-mortem AD brain tissue [73]. This is an important reflection of the potential of similar in vitro systems to recapitulate and investigate key protein signatures that underlie AD pathogenesis [73].

Pitirysin metallopeptidase 1 (PITRM1) is responsible for the processing and degradation of mitochondrial precursor proteins [74]. PITRM1 dysfunction causes improper degradation and over-accumulation of Aβ in the mitochondria, resulting in mitochondrial dysfunction and neurodegeneration [74]. Since PITRM1 deficiency can cause embryonic lethality in mice, mechanisms underlying Aβ accumulation were investigated using human PITRM1 knockout COs (published as a preprint in bioRxiv) [75]. Increases in APP, Aβ40, Aβ42, the Aβ42/Aβ40 ratio, plaque-like amyloid aggregates and pTau-S202/T205 levels were detected in PITRM1 KO COs compared to controls. In addition, increases in ubiquitinated proteins, mitochondrial unfolded protein response (UPR) transcripts and upregulation of mitochondrial proteases were detected in PITRM1 KO COs compared to control COs. These findings indicate that PITRM1 is essential for maintaining mitochondrial protein homeostasis by degrading target sequences like Aβ and possibly tau [75].

Mitochondrial UPR inhibition via integrated stress response inhibitor (ISRIB) resulted in increases of APP, Aβ42/Aβ40 and pTau in PITRM1 KO COs, suggesting a decrease in mitochondrial clearance [75]. Treatment with the NAD+ precursor, nicotinamide mononucleotide, which can improve mitochondrial function and clearance [76], rescued the Aβ and tau pathology in PITRM1 KO COs [75]. This indicates that mitochondrial UPR activation and mitochondrial clearance may play an important protective role in attenuating AD-like phenotypes induced by PITRM1 deficiency. This CO model provides a link between mitochondrial dysfunction and neurodegeneration in AD.

Abnormal endosome recycling and morphology is another related characteristic feature of AD [77, 78]. Endosomes are involved in the formation and engulfment of autophagic
substrates around damaged mitochondria during Parkin-mediated mitophagy [79, 80]. COs with FAD variants show abnormal endosome morphology and a significant increase in large endosomes (1-2 μm) compared to control organoids [43]. Uptake of labelled transferrin via clathrin-mediated endocytosis can be used to assess the functionality of endosomes. There is a significant increase in the size of transferrin-containing endosomes in AD-like COs indicating abnormal endosome recycling [43]. Thus, COs can be employed to investigate the mechanisms underlying mitochondrial dysfunction and cell death in NDs, especially AD.

**Excitotoxicity**

Certain neuronal subpopulations are selectively susceptible to protein aggregation and neuronal loss in AD and other NDs [4]. COs with a PSEN1 M146V variant exhibit an imbalance in excitatory and inhibitory neurons [81]. COs derived from mouse iPSCs also show an increase in glutamatergic neurons [82]. Analyses of COs derived from AD hiPSCs (with different variants) are needed to detect changes in cell-type composition and neurotransmitter properties. Further analysis of AD-like and control COs with scRNA-Seq and complementary immunohistochemistry will be especially critical to determine the gene signatures responsible for selective cellular vulnerability and excitotoxicity in AD.

**Neuroinflammation**

Accumulation of aggregated Aβ and hyperphosphorylated tau can trigger an immune response, mediated by the release of pro-inflammatory cytokines, contributing to the progression of AD [72]. Despite the lack of microglia, neuroinflammation has been observed in COs with a PSEN1 M146V variant [81]. Elevated gene expression of proinflammatory cytokines such as IL-6 and TNF-α has been observed in AD-like COs compared to controls [81]. Increased LDH activity was also detected, indicating increased cytotoxicity and neuronal loss [81]. Treatment of AD-like COs with molecules such as DAPT (a γ-secretase inhibitor), heparin (to compete with heparan sulfate proteoglycan-Aβ42 binding), and heparinase III (to digest heparin sulfate proteoglycan) showed anti-inflammatory effects with reduced toxicity and decreased Aβ levels [81]. While DAPT increased cell survival in AD-like COs, no change was observed with heparin or heparinase III treatment, which may be due to their large size and inefficient diffusion [81].

As detailed below, COs lack microglia, so one strategy to overcome this lack of microglia in 3D models has been addressed using an engineered microfluidic platform [83]. This approach involves the use of chambers for microglia to model the interactions of AD neurons, astrocytes, and microglia. The neurons and astrocytes in the microfluidic platform showed higher differentiation potential and stronger functional electrical activity compared to 2D cultures [83]. This model allows the observation of microglia recruitment, release of cytokines (IL8, TNF-α and IFN-γ), chemokines (CCL2, CXCL10 and CX3CL1) and the neuroinflammatory processes involved in neurodegeneration [83]. Primary human microglia transformed with simian virus 40 have been used in this model [83], and further studies are required with hiPSC-derived microglia and human primary microglia to replicate these findings.

**Neural circuit dysfunction**

Synaptic dysfunction and associated network abnormalities occurring before overt neurodegeneration are implicated as an early pathological event in AD and are believed to contribute to the hyperexcitability in AD as well as other NDs [84–88]. To understand the mechanisms underlying neuroelectric deficits in AD, COs were derived from AD hiPSCs with either a PSEN1 variant (M146V) or the APP Swedish variant [89]. Electrical activity was measured using microelectrode array (MEA) in both 2D cultures and COs, and AD-like COs showed enhanced excitability with shortened neurites compared to control COs [89]. In 2D neuronal culture, increases in the frequency and amplitude of spontaneous excitatory post synaptic currents and a decrease in inhibitory signals in AD-like neurons have also been shown to contribute to network imbalance [89]. This model recapitulates aspects of early AD pathophysiology with changes observed in neuronal morphology and synaptic transmission contributing to aberrant electrical activity, i.e. hyperexcitability [89]. Further studies are required to assess changes in excitatory and inhibitory neuronal subpopulations, synaptic activity, and circuitry function in AD-like COs.

**3D microenvironment abnormality**

The BBB is composed of pericytes, astrocytes and endothelial cells (bECs), which form functional tight junctions to separate brain tissue from peripheral blood and provide a defense against disease-causing pathogens and toxins that may be present in peripheral blood [90]. This highly selective semipermeable membrane becomes dysfunctional due to endothelial and pericyte degeneration in AD [90]. An AD microfluidic platform derived from a ReN cell line with FAD variants has been used to study BBB dysfunction [91]. Increased BBB permeability and decreased tight junctions and adherent proteins were observed in this AD-like model which could be reversed with the drug Etodolac [91].
The importance of the mechanical environment in mimicking AD pathology has been further explored by investigating P21 activated kinases (PAKs) in 3D culture [92]. PAKs convert mechanical stimuli such as the mechanical properties of a 3D matrix to chemical stimuli in a cell [93]. PAKs translocate from the cytosol to the cell membrane in hippocampal neurons treated with Aβ oligomers [93]. Cofilin, an actin-regulatory protein, has a reciprocal relationship with both pPAK and Drebrin, an actin-stabilizing protein [93]. PAKs have thus been implicated in AD, especially in cytoskeleton dynamics and neuronal morphology [93] which has been investigated with a 3D self-assembling peptide (SAP) matrix [92]. As expected, pPAK and Drebrin levels were significantly higher in 3D compared to 2D cultures, with relatively lower levels of Cofilin [92]. Higher co-localization of F-Actin and Drebrin was observed in 3D cultures, however, it could not be recapitulated in 2D cultures [92]. These differences could be due to the SAP matrix that provides a 3D scaffold recapitulating the native brain tissue environment. The effects of Aβ oligomer treatment on the PAK-cofilin signaling pathway has also been studied in 2D and 3D cultures [92]. While no difference has been observed between control and Aβ treatment in 2D cultures, there was a depletion of activated PAK, a redistribution of pPAK to the submembranous region, and the presence of fragmental pPAK in 3D cultures [92]. This study shows that compared to 2D cultures, the 3D culture environment offers a more suitable mechanical environment that recapitulates the AD phenotype and serves as a more relevant in-vitro model for studying the disease.

Changes in the extracellular matrix (ECM) have also been investigated using COs generated from hiPSCs with the PSEN1 M146V variant [81]. Altered levels of ECM molecules like Syndecan-3 and matrix metalloproteinases have been observed in AD-like COs, indicating that changes in ECM composition and 3D environment remodeling is possibly associated with the AD phenotype [81].

**Challenges/Limitations**

Despite the many advantages associated with modeling NDs using COs, several challenges still exist. This section details the major drawbacks associated with COs, including the lack of aging, cell heterogeneity, vasculature, mature synaptic connection, and brain networks. We also discuss how these challenges are being addressed and point out areas that warrant further study.

**Aging**

One current challenge is that hiPSC-derived neuronal cultures do not express all six tau isoforms expressed in the adult brain. These neurons express high levels of embryonic 3R tau and low levels of the 4R tau isoforms [15, 54, 55, 94]. Although this issue has been overcome in a 3D culture with both 3R and 4R tau expression [57], this model does not mimic the ratios of human adult tau isoform expressed in vivo. COs also show fetal neocortex properties making it difficult to model NDs [10, 15, 95, 96]. Additionally, there is a loss of aging-associated markers such as DNA damage and mitochondrial stress in COs [97]. This can potentially be overcome by using the SARS-CoV2 virus, or ZIKV virus, which elicit neurodegenerative phenotypes in COs that mimic age-related deficits as seen in AD [5, 67, 98, 99]. These viruses may therefore be used to model pathological features associated with neurodegeneration. Alternatively, drugs like Aftin-5, rotenone and hypoxic culture conditions can be used to induce Aβ and other aging disease related pathologies in COs [48, 100, 101].

Telomerase manipulation or overexpression of progerin [97, 102] can also be used to model aging. Progeria is caused by a variant in the Lamin A/C gene, which causes accelerated premature aging [103]. Progerin overexpression in hiPSC-derived PD neurons results in the accumulation of ROS, shorter dendrites, DNA damage and loss of tyrosine hydroxylase expression [97]. Telomere shortening is another tool that has proven effective in inducing aging and disease-specific phenotypes such as increase in mitochondrial stress and reduction in the number of dendrites [102]. While artificial aging might not replicate all the real aging phenotypes in human NDs, it provides a more relevant platform for modeling AD and other NDs.

**Microglia and oligodendrocytes**

Occurrence of different cell types in COs such as neural progenitor cells, radial glia, astrocytes, and oligodendrocytes largely depend on the culturing protocol. Longer culture times increase the complexity and diversity of cell types in COs [95]. While cell types within COs are derived from the neuroectodermal lineage, other brain-resident cell types such as microglia are derived from the mesodermal lineage, which accounts for the absence of microglia in COs. Although modifying the differentiation protocol results in innate microglia, there is also an increase in organoid variability with differences in microglial distribution between organoids [104]. The lack of microglia and reduced astrocytes in COs can be overcome by transplantation of human astrocytes and microglia derived from hiPSCs [105–108] into COs, which enhances the frequency of synchronized burst activity and CO development. This type of co-culture is stable long-term and provides a platform to model neuroinflammation and neuron-glia interactions. Methods have also been recently developed to increase the proportion and maturation of oligodendrocytes in COs. In a recent study, COs were exposed to a combination of small molecules and growth factors at defined time points to modulate SHH and WNT signaling and also promote
Oligodendrocyte development and maturation [109]. Additional methods to increase the percentage of oligodendrocytes in COs involve using growth factors such as IGF-1 and PDGF-AA and pro-myelination drugs [110].

Although scRNA-Seq of COs shows the presence of broad classes of cell types, the culture environment may induce ER stress, which in turn affects cellular subtype differentiation [21]. Generation of a heterogenous cellular population in COs involves the use of neural induction and patterning cues along a 3D printed biocompatible scaffold, capable of providing a suitable microenvironment for CO development [81, 92, 111]. One of the major challenges in the field is optimization of differentiation protocols to increase cellular heterogeneity, while reducing the stress and variability associated with organoid culture. One strategy to overcome this challenge is by using a 3D acoustofluidic device to model neuroinflammation in AD [112]. Piezoelectric transducers in the device can manipulate the neurospheroids to model Aβ toxicity, microglia recruitment and activation in AD [112]. Lack of microglia and astrocytes and the inability to model brain regions specifically affected by AD can be overcome by fusing two organoids to form a fusion organoid or assembloid [96, 113]. This can be used to model the inter-neuron migration and neuroinflammation underlying AD.

Vasculature

COs do not have inherent functional vasculature, which may lead to the formation of necrotic cores as their central most region does not receive sufficient nutrients and oxygen supply [43, 111, 114, 115]. Despite the use of miniaturized, multi-well spinning bioreactors which increase diffusion of nutrients and oxygen within COs, oversized COs still lack uniformity and display a necrotic core [116]. This limits the size of organoids that can be grown in vitro without functional vasculature.

COs co-cultured with endothelial cells show vascularization on the exterior but no vascularization in the core of the organoid [117]. Human E26 transforming sequence variant 2 (ETV2) is a transcription factor responsible for the development of vasculature in embryos [118]. Ectopic expression of human ETV2 in COs results in the formation of fetal neocortex-like vasculature [119]. Another alternative is an organoid graft transplanted into a mouse brain, which becomes vascularized with the development of blood vessels, integration of microglia, neuronal maturation and axonal growth extending from the organoid graft into the host [120]. However, in another study, a similar organoid graft into a mouse brain (n=1) did not survive [117], indicating the need for additional studies to assess the viability and development of vasculature in such grafts.

One important aspect of brain vasculature is the formation of the BBB [90]. Several microfluidic devices successfully model the BBB and overcome the lack of vasculature in COs. Despite changes in the microenvironment and BBB permeability, this BBB model lacks astrocytes and pericytes which play an important role in the function of the BBB [91]. To overcome this issue, some models have incorporated rat astrocytes [121, 122] and hiPSC-derived pericytes [123] with endothelial cells, astrocytes, and neurons to model the BBB. These BBB models show in vivo BBB characteristics with high transendothelial electrical resistance and expression of tight junction proteins [121–126]. Despite these advances, the BBB platform and differentiation protocols still require optimization to develop functional vasculature in COs.

Neural circuits

COs fail to establish mature neural networks that can model NDs. COs transition from initial irregular activity to synchronous oscillatory patterns. As the COs develop inhibitory neurons, there is complex spatiotemporal variation in network activity [127]. However, the electrical activity observed in COs is very similar to preterm human EEGs [127], indicating the lack of maturity.

Applications and Future Direction

COs offer several advantages over animal models and 2D cultures, especially in terms of studying human-specific disease initiating events, ease of genetic and environmental manipulation, and applicability to high throughput assays for drug discovery. In this section, we discuss applications of COs and explore avenues that need further research to better understand NDs.

Multi-omics Analysis

Multi-omics studies leverage collective insights from transcriptomics, proteomics, and metabolomics. Total and single-cell multi-omics analyses have been useful in resolving distinct cellular architectures and functions of heterogeneous populations in COs. They can further aid in the elucidation of the relationship between observed phenotypes in NDs and their underlying disease mechanisms. Such analyses may also accelerate advances in the fields of personalized diagnostics and therapeutics.

Transcriptomics

The transcriptomic identities of COs and other 3D culture model systems have been extensively studied [12, 21–23, 95, 127]. Cumulatively, these reports demonstrate that the transcriptome can be readily interrogated to understand the
etiology of NDs and evaluate treatment responses at the RNA level.

Several total RNA sequencing (RNA-Seq) analyses have been performed to validate 3D ND models and discern their disease characteristics. For instance, the transcriptome of a 3D starPEG-Heparin-hydrogel-based culture of human fetal primary astrocytes exhibit several in-vivo signatures such as enrichment of transcripts related to mature neuronal physiology as well as synapse and axon development when compared to an equivalent 2D model [41]. In the same study, Aβ42 treatment in primary and induced 3D cultures resulted in impaired plasticity and network formation, though via distinct molecular mechanisms [41]. These findings underscore the important considerations in experimental design, such as the limitations in molecular programming that induced 3D models may have compared to analogous primary or self-organizing CO models.

Total RNA-Seq analyses have also been carried out in APOE3 and APOE4 COs. A recent transcriptomic study implicated the loss of transcriptional repression by Repressor Element 1-Silencing Transcription factor (REST) in the pathogenicity of APOE4 with 3D SAD models [128]. In comparison to APOE3 COs, APOE4 COs exhibit accelerated neuronal differentiation at the mRNA level, as evidenced by their enrichment of genes associated with nervous system development, synaptic transmission, and neurogenesis [128]. Interestingly, APOE4 COs also exhibit increased total tau expression and pTau-Ser202 [128]. These altered transcriptomic phenotypes appear to be, in part, a reflection of increased activity of REST targets. This mechanism extends to 3D SAD models and provides a more physiologically relevant platform to appreciate the potential importance of transcriptional regulators such as REST in the etiology of NDs.

While total RNA-Seq studies allow for the identification of large-scale changes in transcriptomic profiles, the advent of scRNA-Seq has made it possible to appreciate transcriptionally distinct and diverse cell populations [22, 23, 129]. Given that the human brain and COs are intrinsically heterogenous structures, the resolution of specific cell types at the transcriptomic level is important. An expansive scRNA-Seq study of over 80 thousand single cells from nearly 50 AD patient prefrontal cortex samples revealed that AD has high cell-type specificity in early pathogenesis that later manifest as global stress response [130]. There are also sex-specific trends in AD indicated by transcriptomic alterations in particular cell types [130], which may lend an important insight to the selection of iPSC donors for AD CO models. Novel disease-specific deficits in neuronal differentiation, neurodevelopment and neurotransmission have been identified via scRNA-Seq in CO models for bipolar disorder, schizophrenia and autism spectrum disorder (ASD) [131–133]. Although these studies may not be directly generalizable to neurodegenerative conditions, they exemplify an ideal strategy to detect disease-specific transcriptomic changes at the cellular level, which may be of great benefit to further our understanding of ND mechanisms.

The ability to detect disease- and cell-type-specific molecular signatures of NDs with scRNA-Seq can be of additional value in spatial transcriptomic applications. Spatial transcriptomics methods provide an important morphological context to reconstruct the physiologic organization of expression patterns [134]. It has been used to refine inter-glial activity implicated in the Aβ plaque accumulations of AD pathogenesis present in rodent models [135]. This application of spatial transcriptomics is particularly interesting as it permits the specific resolution of pathogenic features in the vicinity of plaque accumulations. If adapted for use in COs, this method holds great potential to study the human-specific features of AD pathogenesis as they progress with relative spatial organization.

**Proteomics**

Proteomics provides a functional map of all the proteins within a cell or tissue. Proteomic analysis of 3D neuro-spheroids derived from five AD patients exhibited differences in the expression of proteins related to axon growth, mitochondrial function, and antioxidant defense compared to controls [73]. Analysis of AD post-mortem brain tissue also revealed differences in proteins related to axonal injury and transport, oxidative stress, and inflammation [73]. The overlap in the differentially expressed proteins between AD postmortem brain tissue and the 3D AD neuro-spheroids indicates that 3D models can recapitulate at least some of the AD-like phenotypes [73]. Another study involved proteomic analysis of the CA1 and subiculum regions of human postmortem AD brains (n=40) [136]. Analysis revealed an increase in proteins related to astrocyte, ECM, cytoskeleton and synaptic environment over the Braak stages [136]. A similar study of the olfactory bulb in human postmortem AD brains (n=4-5/group) revealed impaired neuritogenesis, synaptic plasticity and protein degradation [137]. Further analysis of the different Braak stages also revealed cytoskeletal dysregulation followed by mitochondrial impairment and unstable RNA transcripts [137]. Such proteomic analyses of postmortem human brains and 3D ND models shed light on the early neurodegenerative mechanisms aiding in the identification of potential biomarker and therapeutic candidates.

Proteomics has also been used to assess drug efficacy by measuring changes in the proteome after drug treatment. For example, one study demonstrated that the treatment of BACE1 and γ-secretase inhibitors significantly reduced Aβ42 levels in four out of five 3D AD neuro-spheroids [138]. Proteomic analysis of these neuro-spheroids with lower Aβ42 levels revealed a 30-40% reduction of APP protein...
(BACE1 substrate) and clathrin (APP endocytosis) compared to the AD neuro-spheroid with high Aβ42 levels [138].

In another study, the proteomic changes in Down Syndrome COs was measured before and after treatment with harmine, a β-carboline alkaloid and an inhibitor of chromosome 21 encoded protein kinase (dual specificity tyrosine-phosphorylation-regulated kinase 1A, DYRK1A) [139]. The Down Syndrome organoids exhibited dysregulation of neurogenesis, axon guidance and extracellular matrix remodeling. After the treatment with harmine, there was a partial restoration of key regulators of cortical development, transcription factors and normalization of ECM proteins [139].

Another proteomics study analyzed COs treated with 5-methoxy-N, N-dimethyltryptamine, a traditional psychedelic drug associated with cognitive gain and structural changes related to attention in the brain [140, 141]. Differential expression of proteins associated with long-term potentiation, dendritic spine formation, microtubule dynamics and cytoskeletal reorganization was observed [141]. There was also a decrease in cell death, suggesting that the drug confers a protective effect [141]. These results indicate the potential of proteomics in identifying subtle differences in protein expression after drug treatment, which could help in elucidating the mechanistic changes induced by drug treatment and could pave the way for personalized medicine in the future.

Metabolomics

Metabolomics is the study of the metabolome, which generally refers to all small molecules from a biological sample with a molecular weight of less than 1.5 kDa. It provides a snapshot of the interaction between genetic and environmental factors. Since metabolites are the end products of the cellular regulatory processes, metabolomics is regarded as the omics field most closely related to phenotype [142–144]. Therefore, it holds great potential for the discovery of ND-relevant biomarkers and therapeutic targets. Although to date, there have been no reports of metabolomics profiles in COs generated from patients with NDs, metabolomics has been applied to AD patient samples with some success. In one such study, the authors utilized untargeted metabolomics of cerebrospinal fluid (CSF) from control subjects or subjects with stable mild cognitive impairment (MCI), MCI that progressed into AD, or AD and identified 17 biomarkers capable of predicting the development of AD with an accuracy of 98.7% [145]. This and other metabolomics studies [146–150] of AD patient samples not only provide great potential for developing methods for early diagnosis in AD, but also provide a potential mechanism to validate AD-like COs and other ND organoid models to ensure that they recapitulate the phenotypes seen in humans.

Metabolomics studies on organoid models of other diseases demonstrate the promise of the method for discovery in NDs. The CLN3Q352X variant was identified in a juvenile form of neuronal ceroid lipofuscinosis (JNCL), a lysosomal storage disorder that causes pediatric neurodegeneration. Decreases in metabolites like creatinine and gamma-aminobutyric acid (GABA) were observed in CLN3Q352X organoids compared to isogenic control organoids [151]. Transcriptomics and metabolomics data showed that the CLN3Q352X variant caused neurodevelopmental changes and neurodegeneration in JNCL [151].

Metabolic profiles have also been generated from healthy 3D neurospheres at 2, 4 and 8 weeks of differentiation [100]. Changes in the metabolic profile are consistent with the different stages of differentiation [100]. After the treatment of 5 μM rotenone, which interferes with the electron transport chain in mitochondria, analysis of the metabolites confirmed mitochondrial dysfunction, disruption of gene regulation by peroxisome proliferation and defective nervous system development [100]. Thus, metabolomics offers great potential for assessing drug efficacy, elucidating underlying disease mechanisms and discovery of clinically relevant biomarkers in COs modeling NDs.

Propagation and spread of pathogenic protein aggregates

Protein misfolding, aggregation, and cell-to-cell propagation of these protein aggregates is believed to play important roles in the pathogenesis of many NDs. These processes have been modeled in animals, cell culture, and human postmortem brain tissue [6, 7, 152]. To our knowledge, COs have not been used to model the propagation and spread of Aβ and pTau. However, COs have been used to model propagation in Creutzfeldt-Jakob Disease (CJD), a type of prion disease in which there is abnormal buildup of prion protein, which propagates from cell to cell in an infectious manner causing dementia [153]. In this study, COs were inoculated with two different conformational CJD subtypes MV1 or MV2 (Methionine-Valine type 1 and 2) [153]. MV2 subtype showed higher seeding activity than the MV1 subtype. Even though seeding caused by the initial inoculum was cleared, a resurgence of prion seeding activity was detected in the COs indicating de-novo propagation and spread [153]. Modeling prion propagation in COs provides encouraging evidence that the propagation and spread of other conformational proteins such as pTau and Aβ isolated from patients with AD and FTD may be possible in COs as well.

Although literature on propagation and spread in COs is still lacking, 2D cultured neurons derived from hiPSCs have been used to investigate the spread of intermediate protein strains in AD. Tau repeat domain (tau-RD) with P301L and V337M (tau-RD-LM) variants introduced into hiPSC-derived...
neurons have shown AD-like aggregation of tau. Tau seeds extracted from this culture have shown the ability to recapitulate and induce tau aggregation in a non-transduced neuronal culture [154]. This indicates that the extracted pathological tau seeds can cause tau aggregation via cell-to-cell propagation and similar studies should be replicated in COs.

Aβ and tau oligomers are intermediate protein species that are toxic in AD [155]. Mechanisms underlying the pathogenicity and spread of these species have been investigated in 2D cultures. In one study, neurons derived from healthy hiPSCs were treated with either full-length tau monomers or oligomers. While there was no significant difference between the monomer treatment and control, neurons treated with tau oligomers recruited endogenous tau and showed an increase in pTau (T231, S396/404) aggregation, resulting in progressive neurite degeneration and neuronal loss [156]. Tau oligomer treatment has also shown a reduction in synaptic integrity, neurotransmitter release, and calcium signaling [156], which is in agreement with the toxicity induced by tau oligomers in primary neurons and AD-like animals [155, 157–159].

Another study analyzed the propagation of AD brain-derived oligomeric Aβ (oAβ) via exosomes in neurons differentiated from the SHSY5Y neuroblastoma cell line and the neural stem cell line (AF22) derived from healthy hiPSCs [160]. Neurons incubated with oAβ-rich exosomes (donor cells) and co-cultured with recipient neurons showed the presence of oAβ in both neuronal types [160]. When secondary neurons were further co-cultured with untreated SHSY5Y neurons, they also showed the presence of oAβ, indicating that unexposed neurons take up and propagate intact exosomes rich in oAβ [160]. Inhibition of exosome secretion by siRNA significantly reduced oAβ transfer between neurons. A 97.7% reduction in exosome uptake by the recipient cells was observed when neurons were treated with Dynasore (a dynamin inhibitor) [160]. This indicates that dynamin is likely involved in the propagation and spread of oAβ [160]. These results need to be replicated in COs since they provide a more complex environment to study the propagation of protein aggregates [8]. Other mechanisms underlying the propagation of aggregated proteins such as receptor mediated, clathrin dependent and independent endocytosis [161] also need to be explored in COs.

**Neuronal and regional cellular vulnerability**

Neuronal sub-populations show differential vulnerability to different pathological protein aggregates found in NDs, which cumulatively aggregate and thereby affect distinct regions of the brain [3, 4, 6]. Mechanisms underlying this ‘selective vulnerability’ are important to understanding disease etiology and designing targeted therapeutic strategies. The APOE gene encodes for the apolipoprotein E protein and the E4 variant is the strongest genetic risk factor for AD [162]. Reduction in Aβ was observed in APP duplication COs treated with APOE3-carrying microglia-like cells compared to the treatment with APOE4-carrying microglia-like cells. This indicates that the APOE4 variant hinders the ability of microglia to clear Aβ in AD [78]. Despite high levels of astrocytes by six months, APOE4 COs still have high levels of Aβ and significantly lower levels of full-length APOE protein, indicating impaired Aβ clearance by astrocytes compared to APOE3 COs [78]. APOE4 COs also exhibit increased Aβ aggregates and pTau (S202/T205) levels compared to APOE3 COs. Although AD-like pathology is delayed in APOE4 COs (six months) compared to FAD COs (two months) [43], these results illustrate that the APOE4 variant alone is sufficient to make the COs vulnerable to AD-like pathology [78].

To further understand and observe if APOE4 effects can be reversed, hiPSCs with the APOE4 allele were gene edited to APOE3 via CRISPR/Cas9 and used to generate COs. The gene-edited COs showed a non-significant decrease in Aβ42, Aβ40 levels and early endosomes possibly pointing toward additional factors in the hiPSC line that may be responsible for AD-like pathology [78]. Higher APOE levels correlated with an increase in Aβ uptake by astrocytes and microglia [78]. Similar to 2D neurons, Aβ accumulation was reduced in APOE3 (CRISPR/Cas9 edited) organoids compared to APOE4 COs, indicating that AD-like pathology can be reversed by gene editing APOE4 to APOE3 [78].

Although APOE4 COs are more susceptible to AD-like pathology, the affected neuronal subpopulations have been explored only in 2D cultured neurons [163]. Cholinergic neurons derived from AD patients with the APOE3/4 genotype (AD-E3/E4), FAD and control hiPSCs are all electrophysiologically active [163]. However, compared to control neurons, AD-E3/E4 neurons show a significant increase in the ratio of Aβ42/Aβ40 and altered responses to γ-secretase inhibitors. When exposed to low levels of ionomycin (calcium ionophore), AD-E3/E4 neurons show a 50% reduction in viability compared to controls [163]. AD-E3/E4 neurons also display increased cytoplasmic calcium and excitotoxic death [163], indicating that these neurons are more susceptible to glutamate induced excitotoxicity, which is consistent with the increased risk of AD associated with the E4 allele [162].

Another study analyzed neurons derived from hiPSCs with APOE3/3 and APOE4/4 genotypes [164]. APOE4/4 neurons produced more fragmented APOE with higher Aβ and pTau levels (S396/S404, S202/T205) compared to APOE3/3 neurons [164]. The APOE4/4 genotype affected only GABAergic neurons, making them more susceptible to degeneration compared to glutamatergic and dopaminergic neurons [164]. When the APOE4/4 line was genetically edited to APOE3/3 or treated with a structure corrector (such as PH002), the neurons showed higher levels of full-length APOE with decreased Aβ and pTau levels [164], rescuing AD-like phenotypes.
Interestingly, Aβ levels did not change in APOE knockout (KO) neurons treated with purified recombinant APOE3 or APOE4, indicating that Aβ aggregation is dependent on endogenous levels of APOE [164]. Human astrocyte-derived APOE4 also does not alter Aβ and pTau levels in APOE KO neurons, indicating that the gain of toxic effects is probably neuron specific [164]. More investigations are required to assess the selective neuronal vulnerability to different Aβ and tau species in AD.

Neurons derived from hiPSCs with the FAD APP London variant showed different APP processing depending on the differentiation fate of these neurons [165]. For example, an increase in pathogenic Aβ42 production in rostrally directed neurons compared to caudally directed neurons has been observed. In addition, the α- and β-secretases and Anterior Pharynx Defective 1, a component of γ-secretase, were differentially expressed between different neuronal fates thereby differentially affecting APP processing [165]. The pTau (S202/T205) and total tau levels were also elevated in rostral neurons compared to neurons directed to caudal fates [165]. Control neurons without a FAD variant exposed to soluble AD-brain extracts also show increased tau phosphorylation in rostrally directed neurons, while tau phosphorylation remains unaffected in neurons directed to caudal fates [165]. This model illustrates that neuronal subtypes in different brain regions respond differently to the APP variant in terms of APP processing and tau homeostasis [165, 166], which may partially dictate the pattern of selective regional and neuronal cell vulnerability in human AD.

In another study, accumulation of tau in neurons led to increased vulnerability to oxidative stress. Cortical neurons derived from FTD patients with the MAPT heterozygous variant A152T showed time-dependent increases in pTau levels (S202/T205, T231/S235, S396/S404, S396) and decreases in the solubility of tau protein [94]. These neurons also showed an increase in ubiquitinated proteins and markers of autophagy, indicating a disruption in protein degradation and clearance via Ubiquitin Proteasome System and the autophagy-lysosome-system [94]. Treatments of A152T neurons with different compounds (mitochondrial electron transport chain inhibitors, excitotoxins, or proteotoxins) showed selective neuronal loss [94], suggesting that pTau accumulation in A152T neurons is associated with increased stress vulnerability. Using CRISPR/Cas9 genomic engineering, A152T neurons were modified to disrupt the MAPT gene resulting in a downregulation of tau expression [94]. When these neurons lacking MAPT were treated with the same set of compounds, there was a rescue of stress vulnerability with an increase in viability similar to healthy control neurons [94]. This model demonstrates that accumulation of tau makes neurons more vulnerable to stress leading to neurodegeneration. Different factors such as APOE genotype variants and drug treatment should be studied in both 2D and 3D cultures, especially COs, to further understand the selective neuronal/regional vulnerability associated with AD.

**Neural circuits**

COs lack proper spatial information important for establishing functional neural circuits as observed in vivo [16]. Fusion of regionally specified COs would provide a more physiologically relevant platform to study neuronal circuits. For example, the dorsal/ventral forebrain spheroid model creates a gradient of trophic factors that encourages interneuron migration, resulting in diverse and complex 3D brain regions with functional neuronal networks [113].

Medial ganglionic eminence organoids (MGEOs) and COs have been fused to generate functional neuronal networks to study interneuron migration [96]. Individual neurons in COs and MGEOs showed calcium surges resulting in synchronized neuronal activity [96]. Following treatment with bicuculline, a GABA receptor antagonist, these assembloids show an increase in calcium surge synchronization [96]. This indicates that GABAergic inhibition increases neural activity in both COs and MGEOs [96].

Human cortical spheroids (hCS) and subpallium spheroids (hSS) have also been fused to study interneuron migration, and changes in electrical activity have been found before and after the fusion [113]. Neurons that migrated from the hSS to the hCS display a doubled maximum action potential generation rate compared to non-migrated cells and neurons in unfused hSS [113]. Whole-cell voltage clamp recordings of migrated neurons also display excitatory and inhibitory postsynaptic currents (EPSCs and IPSCs) [113]. However, these neurons receive more EPSCs than IPSCs with a 3-fold increase in synaptic input. EPSCs have also been observed upon electrical simulation indicating the establishment of a functional neural network after assembly of hCS and hSS [113].

Alternative models have been established that fuse regionally specified COs or use distinct classes of neurons within a microfluidics compartment to reconstruct neuronal circuitry [16, 167]. For example, in vitro models that reconstruct cortico-striatal and cortico-cortico circuits have been developed to understand molecular mechanisms underlying Huntington’s disease and ASD, respectively [16, 167]. In some cases, MEA has been utilized to assess localized electrical activities within these models. However, a current challenge relates to correlating acquired electrical activities with the identity of neuronal subtypes. Advances in genetically encoded calcium and voltage indicators [168, 169] can potentially provide a solution for correlating electrical activities acquired with MEA with distinct neuronal subpopulations.

Improvements in design and culturing protocol can also aid in establishing mature neural circuits. An example is the use of fetal brain-derived ECM, rich in chondroitin sulfate, hyaluronan and heparin sulfate proteoglycan, which contain...
understand the role of these and other risk factors in the progression of AD. It should be noted that CRISPR-Cas9 genomic editing has not only been used to generate isogenic controls [45, 54, 75, 89] but also used as a therapeutic strategy to modulate or revert accumulation of Aβ [78]. This technology can also be used to design a high-throughput screen [182] for genes associated with Aβ and tau aggregation, which will shed more light on the underlying mechanisms of AD.

**Biomarker identification and drug screening**

Compared to 2D cultures, COs reproduce human conditions like AD more closely [35, 36, 45, 138]. Single cell transcriptomics of patient derived COs have shown considerable promise in identifying novel biomarker candidates [15, 132, 171–173]. β and γ-secretase inhibitors decrease Aβ and pTau accumulation in COs [35, 36, 45, 138]. Treatments with other compounds like heparin, heparinase III, DAPT [81], IL-4 [41] and nicotinamide mononucleotide [75] also show a reduction in Aβ induced toxicity and reversal of other AD-like pathologies such as pTau and Aβ plaque-like accumulation. Further studies are required to investigate if treatment and removal of toxic Aβ or tau species prior to the onset of toxic protein aggregation, altered synaptic function and cognitive deficits, can prevent, and/or delay the development of AD-like pathology in COs.

COs also provide greater translatability to clinical trials and thus accelerate the drug discovery process. These trials either target clearance of Aβ and tau or focus on immunotherapy involving active or passive immunization. Several of them have failed in Phase 2/3 clinical trials due to lack of efficacy or toxicity issues [29–34, 174–177]. The development of a human choroid plexus organoid [178] may accelerate the drug discovery process since it is involved in the production of CSF which regulates movement of nutrients/drugs into the brain [179]. Strategies could be devised to employ choroid plexus organoids in high-throughput testing of all FDA-approved drugs for their ability to provide therapeutic benefit in AD. Based on bioavailability, safety, efficacy, and clearance of these drugs in the organoids, they can potentially be quickly repurposed as AD therapeutics or prescribed for off-label use in treating AD.

Although most of the current CO models are based on FAD variants (Table 1), it is also important to use the CO platform to model AD-associated risk factors such as diabetes, hypertension, ischemia, TBI, smoking, and other environmental factors by modifying culture conditions [180]. For example, exposure of COs to heme results in brain injury leading to extensive structural disorganization, increases in proinflammatory chemokines and neurodegeneration [181]. This can be used to model and study the mechanisms underlying TBI in COs. These novel CO models will thus help the field to understand the role of these and other risk factors in the development and progression of AD.

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**Compliance with ethical standards**

**Conflicts of Interest**

The authors declare they have no conflict of interest.

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