Co-development of central and peripheral neurons with trunk mesendoderm in human elongating multi-lineage organized gastruloids

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Stem cell technologies including self-assembling 3D tissue models provide access to early human neurodevelopment and fundamental insights into neuropathologies. Gastruloid models have not been used to investigate co-developing central and peripheral neuronal systems with trunk mesendoderm which we achieve here in elongating multi-lineage organized (EMLO) gastruloids. We evaluate EMLOs over a forty-day period, applying immunofluorescence of multi-lineage and functional biomarkers, including day 16 single-cell RNA-Seq, and evaluation of ectodermal and non-ectodermal neural crest cells (NCCs). We identify NCCs that differentiate to form peripheral neurons integrated with an upstream spinal cord region after day 8. This follows initial EMLO polarization events that coordinate with endoderm differentiation and primitive gut tube formation during multicellular spatial reorganization. This combined human central-peripheral nervous system model of early organogenesis highlights developmental events of mesendoderm and neuromuscular trunk regions and enables systemic studies of tissue interactions and innervation of neuromuscular, enteric and cardiac relevance.
three-dimensional in vitro tissue models called organoids have revealed the astonishing innate capacity of stem cells to self-organize into complex cytoarchitectures resembling in vivo states. Directed single-organoid models of anatomical endpoints of interest, such as regions of the brain, spinal cord (SC), gastrointestinal tract, and heart, are rapidly becoming indispensable to biomedical research for modeling physiology, development, disease, aging, and toxicity. Dual-fated trunk organoids have been applied to model the neuromuscular circuit, demonstrating an ability to recapitulate several key aspects such as interneurons, motor neurons, and the neuromuscular junction (NMJ) with functional output. However, beyond organogenesis, the study of multi-tissue interactions in human models will benefit from an ability to better recapitulate the morphogenetic complexity of embryogenesis. As such, a shift towards new organoid models, deemed gastruloids, which more accurately reflect early mammalian development in a multi-lineage, embryo-like context has emerged. Pioneered primarily using mouse embryonic stem cells (mESCs), and in one study using human embryonic stem cells (hESCs) for anterior-posterior organization, gastruloids have been applied in vitro to demonstrate symmetry-breaking and axial elongation events. By small molecule interference with germ-layer primitive gut tube is a central element along with the SC endoderm and mesoderm that is then populated by NCCs. The formation and elongation, and the corresponding neuronal patterning events in that compartment. We characterize EMLOs demonstrating the presence of the mu opioid receptor (MOR) in EMLOs and the potential utility of EMLO models in pharmacological studies with agents impacting sensory neuronal activity versus spinal motor neurons that lack the receptor. Our unique EMLO approach with human stem cells is an advanced heterogeneous 3D culture platform for NCC analysis, trunk endoderm innervation, and neuromuscular studies.

Results

Reproducible generation of EMLO gastruloids from African American, Hispanic-Latino, and Asian hiPSC lines. To increase stem cell studies that reflect ethnic diversity (ED) in the United States, we analyzed EMLO gastruloids using previously derived hiPSC lines from our laboratory. Nine ED-hiPSC lines derived from the fibroblasts of three donors of self-designated African American (F3.5.2, F3.2.2, F3.3.1), Hispanic-Latino (H3.3.1, H3.1.1, H3.4.1), and Asian (A2.1.1, A2.2.1, A2.2.2) ethnicities were compared. EMLOs were generated by modeling a dual-fated neuromuscular organoid (NMO) protocol (Supplementary Table 2). These modifications allowed retention of primitive mesendoderm and ectodermal phenotypes during EMLO formation that has not been previously investigated. In brief, intact 2D stem cell colonies were pretreated with CHIR and fibroblast growth factor (FGF2) in N2B27 induction medium for 2 days. After pretreatment, we generated highly uniform 3D starting aggregates (~100 μm diameter, ~300–400 cells per aggregate) by dissociation and immediate transition to low adhesion shaking cultures, bypassing static 96-well plates. FGF2, hepatocyte growth factor (HGF), and insulin-like growth factor-1 (IGF-1) growth factors were included in N2B27 at the time of spontaneous aggregation. Aggregates were split 1:2 after 24 h and maintained to day 4, at which point aggregates were expanded to 100 mm dishes and maintained in N2B27 for 2 days.

We monitored morphogenetic properties over time by phase-contrast microscopy and whole-mount immunofluorescence (IF) of tissue-cleared samples. SOX2 and GATA6 are two essential transcription factors involved in myriad early developmental polarization events, wherein the individual deletion of each gene is embryonic lethal. In EMLOs, we observed polarized gene expression of SOX2 versus GATA6 as soon as 24 h after aggregation (Fig. 1d). A prominent GATA6 domain emerged by day 3, opposite the SOX2 domain as previously observed in human gastruloids with single bilateral protrusions extending out of the SOX2 domain. Although the outer cell layer was predominately GATA6+, the inner invaginated layer co-expressed GATA6 and SOX2. This mixed factor protrusion continued to elongate between days 4 and 16 (Fig. 1e, day 13), a critical EMLO elongation period, and ultimately produced a laminated tube-like structure that retained SOX2/GATA6 double-positivity (Fig. 1f). In general, although there is overlap during initial EMLO polarization events (Supplementary Fig. 2e), the SOX2 domain matures to have primarily neural SC identity that is posterior, and the GATA6 domain matures to have primarily mesoderm and endoderm identity (mesoderm–endoderm, ME), and is anterior. We use the terminology "domain" when referring to early biomarkers, and "compartment" when referring to the tissue structure as a whole. To exclude any contribution of aggregate fusion to EMLO gross morphological complexity, we tracked individual aggregates and monitored elongation (Supplementary Fig. 1g). By this approach, we also observed the emergence of a dilated vesicular structure in the anterior EMLO
pole of ME that extended in length and in size to day 13 (Fig. 1e), after which this region resembled amorphous sacs lined by GATA6+ cells (Supplementary Fig. 1h).

To evaluate reproducibility of EMLOs, we measured gross size parameters and assessed formation efficiency using the morphologically distinct hourglass shape in all nine ED-hiPSC lines (Fig. 1g–i). The major axis length (L) was measured between the most distant points of the posterior and anterior poles, while the minor axis after elongation was defined as the width (W) of the ME compartment near the SC exit point (Fig. 1g). We categorized day 13 EMLOs as elongated long, elongated short, or spherical/ovoid based on aspect ratio in N = 3 repeated formation experiments (Fig. 1h). Replicate ED-hiPSC lines had more similar elongation efficiencies, with greater differences seen between
coherents. Among all lines, H3.3.1 had the highest efficiency of elongation (52.8% elongated long, 20.8% elongated short, 73.6% total). Given similar EMLO formation efficiencies within replicate lines, we chose one line from each cohort to investigate further that were F3.5.2, H3.3.1, and A2.1.1. Representative lines were previously extensively characterized genomically, transcriptomi- cally, functionally, and by protein biomarker in multi-lineage differentiation platforms. All three lines form teratomas in nude mice. Aspect ratios in EMLOs were measured over time (Supplementary Fig. 3). As well, EMLO elongation was not observed in cerebral organoids generated from anterior NSCs, or in suspension neurospheres that used caudal scNSCs as starting material (Supplementary Fig. 3). As well, EMLO elongation was prevented by addition of 1 μM retinoic acid starting on day 2, presumably owing to premature conversion to neuroectoderm (Supplementary Fig. 3a, c). Culture conditions applied to promote or inhibit EMLO formation and elongation are summarized (Supplementary Table 2).

Self-organization of a primitive gut tube-like structure follows initial polarization in EMLOs. Although a 3-day pretreatment of hiPSCs seeded as single cells yields homogenous adherent cultures of SOX2/Bra neuromesodermal progenitors (NMPs) that benefit NMO formation (Supplementary Fig. 1c–f),3, our shorter induction period yielded a less-restricted mesoderm-like cellular starting material characterized by expression of SOX2, Bra, and the definitive endoderm marker FOXA2/HNF-3β (Supplementary Fig. 2a–d). By IF of day 3 early EMLOs (Supplementary Fig. 2e), we observed compartmentalized expression of transcription factors that orchestrate multi-lineage gene regulatory networks in development. Similarly to gastruloids, EMLO morphological asymmetry and elongation were preceded by asymmetric polarization of cellular pools within aggregates. Cells expressing definitive endoderm markers SOX17 and FOXA2 were detected within the SOX2 domain as previously observed.5,6 Bra+ cells that precede mesoderm were localized to the tip of the posterior pole by day 3. Expression of CDX2, master upstream regulator of caudal Hox genes, occurred opposite the GATA6 domain. Together, these data suggest that multiple germ layers co-emerge in early EMLOs and contribute to downstream morphology and reproducibility. Early cytological complexity was evident and nearly indistinguishable in all three representative lines.

We performed scRNAseq analysis on EMLOs at day 16 in formation using the H3.3.1 line (n = 15,576 cells) (Fig. 2a) and applied 10x Genomics Loupe analysis software. Eighteen clusters were produced and were annotated based on canonical gene biomarkers. Gene lists used for annotation are provided as a separate data file. At this time point, several important clusters were identified including gut endoderm (cluster 8), mixed mesoderm (cluster 6), floor plate-like and roof plate-like signaling hubs (clusters 7 and 15, respectively), neural tube and SC-like regions (clusters 4, 5, 9, 10), NCCs (cluster 14), ectoderm (cluster 16), differentiating neural progenitor populations (clusters 1–3, 11, 12), and a small vascular/endothelial cluster (cluster 18). A robust, self-organizing tube-like structure spanned the SC and ME compartments in ~100% of EMLOs that underwent elongation. This structure did not occur in the non-elongated spherical/ovoid subset. To determine the identity of this tube structure, we investigated individual transcripts within clusters and performed additional imaging analysis (Fig. 2b–j). This architectural feature was visible by phase-contrast microscopy (Fig. 2c). By Z-stack 3D reconstruction (Fig. 2d), we show that the GATA6 expressing tube is indeed a continuous, cylinder-like structure with a central lumen containing mitotic figures at the apical aspect (see also Supplementary Fig. 4, Supplementary Movie 1).

We determined that the structure has gut tube epithelial identity by coupling canonical gene expression in cluster 8 for definitive endoderm (EPCAM, CDH1/E-cadherin, FOXA2, GATA6, SOX17, CDX2) (Fig. 2b) with IF of FOXA2 (Fig. 2g) and E-Cadherin (Fig. 2h). E-Cadherin was expressed in the tube in both SC and ME compartments indicating definitive endoderm (Fig. 2h). GATA6 was strictly excluded from SC, though highly expressed both in the palisading tube epithelium and the surrounding, loose mesenchyme-like tissue. These two regions in ME were separated by acellular space indicating basement membrane. Similarly, FOXA2 and GATA6 were found to colocalize in cluster 8. Based on these data, we classified this structure as a putative primitive gut tube, hereon simply referred to as the gut tube. Gut tube identity was supported by low level SOX2 co-expression with high GATA6.20,21 To determine the approximate relative proportion of neural versus non-neural lineages, we quantified the areas of SOX2 versus GATA6 domains in EMLOs over time in maximally projected images from complete Z-stacks (Fig. 2e). SOX2 basal expression in the gut tube was excluded for the purpose of these measurements. We also measured the gut tube major and minor axes over time (Fig. 2f), which correlated with total EMLO elongation and aspect ratio (Fig. 1i) in support of a critical elongation period from day 4 to 16. A FOXA2+ gut tube was evident by day 8 (Supplementary Fig. 2f), though became increasingly morphologically complex in close spatial relation to TUJ1 neuronal fibers (Fig. 2g–i, Supplementary Movie 2) and late-stage day 40 EMLOs had meandering gut tube morphologies (Fig. 2i). Using Boolean filter rules of day 16 scRNAseq data, we identified distinct gastrointestinal cell populations in cluster 8 that are intestinal stem cell...
(LGR5/SOX2), esophagus–trachea junction (NKX2-1/SOX2), Paneth cell (LYZ/SOX2), and enterocyte (VIL1/SOX2). A qualitative summary of EMLO biomarker distribution analyzed by IF is provided (Supplementary Table 3).

EMLO gastruloids have default dorsal-intermediate SC identity that can be ventralized. We corroborated EMLO trunk identity by scRNAseq data, identifying an SC-like region in clusters 4, 9, 10, floor plate- and roof plate-like signaling hubs, and a suite of Hox genes that are known to be absent in anterior neuroectoderm. EMLOs exhibited a distinct lack of anterior neuroectodermal transcripts OTX2, FOXG1, and TBR2 (Supplementary Fig. 5) and these proteins were not observed by IF. To evaluate cell and lineage identity in the EMLO neural SC compartment, we
performed IF of neurons and neural crest cells, or NCCs (Fig. 3). By IF of β-III-tubulin (TUJ1), a neuron-specific β-tubulin isotype, we observed a dense population of neurons in SC that colocalized with TFP2α, indicating NCC lineages and/or developing GABAergic interneuron progenitors. A small subset of neurons projected into the ME compartment, shown within GATA4+ tissue at day 13 (Fig. 3a top). Notably, GATA4 was excluded from the gut tube but highly expressed in the surrounding mesenchyme. At day 16, we demonstrated that the projecting neurons enveloped the GATA6+ gut wall as well as close association with neuronal fibers in ME, indicating a migratory population of NCC-derived neurons or fate-biased non-neuronal NCCs that migrate along or in proximity to neuronal fibers. The number of TFP2α+ nuclei in ME increased over time, whereas TUJ1 signal increased in both SC and ME. By day 16, TFP2α+ neurons self-organized into a single ganglionic structure at the base of the gut tube. The neuroanatomical term ganglion is used in the literature to describe neural structures in organisms that range from invertebrate to mammals, and is used here similarly to mean a peripheral structure that contains a collection of neuronal progenitors, types of neurons, and neurotransmission phenotypes. We further applied scRNAseq and biomarker imaging to distinguish neural progenitors, types of neurons, and modes of neurotransmission (Fig. 4). We distinguished neural progenitors (TU/BD3 OR MAP2; 6700 cells) from committed neurons (TU/BD3 AND MAP2; 3030 cells). By this filter, cluster 10 contained the highest proportion of these neuron types and was expanded for visualization. Together, these data suggest that EMLos have default dorsal-intermediate SC identity, and that populations of neural progenitors and subtypes can be further manipulated by exogenous signaling interventions.

**EMLOs specialize to generate motor, sensory and autonomic neurons, and neurotransmission phenotypes.** We further applied scRNAseq and biomarker imaging to distinguish neural progenitors, types of neurons, and modes of neurotransmission (Fig. 4). We distinguished neural progenitors (TU/BD3 OR MAP2; 6700 cells) from committed neurons (TU/BD3 AND MAP2; 3030 cells). By this filter, cluster 10 contained the highest proportion of these neuron types and was expanded for visualization. Together, these data suggest that EMLos have default dorsal-intermediate SC identity, and that populations of neural progenitors and subtypes can be further manipulated by exogenous signaling interventions.

**Neuronal cytoarchitectures are established in close relationship to the gut tube.** Given the close physical relationship between neurons and the gut tube in EMLos, we sought to further elaborate and quantify this interaction. In parallel with an increase in the diameter of the SC compartment over time (Fig. 5a), we
measured a corresponding increase in TUJ1 density in ME from days 13 to 22 (Fig. 5b). Normalized TUJ1 fluorescence was computed from complete Z-stack volumes of this isolated region as a proxy for TUJ1 density. At day 16, the early formation of peripheral ganglia was observed in addition to a neuronal bottleneck effect at the transition zone between SC and ME (Fig. 5c), and using 3D reconstruction of high magnification Z-stacks, it is clear that neuronal fibers in ME completely envelop the gut tube, but are largely excluded from the lumen and the basal lamina (Fig. 5d; Supplementary Movie 3). In a subset of EMLOs, scant TUJ1 fibers were detected within the lumen. By comparison of maximally projected Z-stacks to individual Z-slices in EMLOs at day 22, we demonstrated the retained envelopment of the gut tube and the primary large peripheral ganglion, in addition to the establishment of smaller satellite ganglia distributed throughout the ME in support of advancing neural pattern complexity (Fig. 5e). By 3D reconstruction of the EMLO exterior, an undulating GATA4+ ME surface interpenetrated by neuronal fibers and ganglia was evident. We refer to the anterior pole of the gut tube as the anterior intestinal portal-like region, or AIP.
Gut tube and mesoderm contribute to EMLO elongation and neuronal pattern formation. To determine whether mesoderm and endoderm have a role in EMLO elongation, we performed a pharmacological inhibition experiment (Fig. 6a, b). We introduced dual SMAD inhibition by small molecules LDN 193189 (200 nM) and SB 431542 (10 μM) to inhibit mesoderm and endoderm progression during the elongation period. LDN and SB were added at day 2 in EMLO formation, a time point near the establishment of sub-aggregate polarized domains, and maintained until day 10. EMLO morphology was quantified at day 22 versus DMSO controls. At this time point, a significant proportion of EMLOs exhibited truncated ME compartments in all three lines (Fig. 6b). N = 3 repeat experiments were performed. Notably, SC cytoarchitecture that is characterized by the emergence of neurons at the base of rosettes and a bottleneck effect in the transition zone was not affected by this intervention. However, we did not detect peripheral ganglionic structures in truncated aggregates. These results suggest that EMLO elongation may occur due to ME-driven processes as opposed to NMP-driven processes of axial elongation. Although EMLOs were formed from cells co-expressing SOX2, Bra, and FOXA2 (Supplementary Fig. 2c, d), NMPs were not detected at the day 16 time point by scRNAseq, suggesting that NMPs are transitory and may precede this stage. These multiple features distinguish EMLOs from all other previously described gastruloid models. To follow-up on motor neuron transcriptional programs observed in the scRNAseq cluster 8 (gut tube epithelium), we performed additional sub-cluster analysis (Fig. 6c). Interestingly, a subset of cells expressing epithelial transcripts FOXA2 and EPCAM also expressed neuronal TUBB3 (243 cells). A further subset of these FOXA2/EPCAM/TUBB3 cells expressed neuronal MAP2 (128 cells), and then the additional endodermal marker SOX17 (16 cells). We performed FOXA2 and TUJ1/TUBB3 IF on day 40 EMLOs and identified a subset of FOXA2 cells in the epithelium that also strongly expressed TUJ1 with neuronal morphology (Fig. 6d). It was recently shown in vivo in mice that enteric neurons have dual embryonic origins from both neural crest and endoderm. Our result may represent a similar phenomenon, and if so, EMLOs would constitute the only human model system currently available to further interrogate this developmental process.

Neural crest cells position to form neuronal networks along the gut tube. Gastruloids and NMOs have been shown to generate numerous cell types from more than one germ layer including NCCs. In vivo, competing transcriptional programs are first co-activated in NCCs, which are then lineage-biased in migrating progenitors. We interrogated the diversity of cell types that arise in EMLOs and their relationship to the gut tube by scRNAseq and IF (Figs. 7, 8), focusing our analysis here on NCCs. TFAP2a and SOX10 are canonical markers of NCCs. We identified cluster 14 as the primary NCC cluster based on these and other canonical biomarkers (Figs. 7, 8, Supplementary Fig. 6). At day 6, SOX10 and TFAP2a largely overlapped by IF but were seen to partially diverge by day 20, where subpopulations of SOX10+ cells could be distinguished based on TFAP2a expression in the expanded cluster 14 (Fig. 7a, b). Similarly, a subpopulation of TFAP2a+ cells could be distinguished based on TUBB3 expression indicating both neuronal and non-neuronal NCC lineages (Fig. 7c), again validated both by scRNAseq and IF. A small subpopulation of EMLOs lacked the SC compartment (Fig. 5d). In these elongated ME-only aggregates, the gut tube was present (ISL1+), but the NCC biomarkers SOX10 and TFAP2a were distinctly lacking, suggesting that NCCs can populate the ME over time to produce neurons. However, a separate population of NCCs that can arise in ME given additional time or additional signaling manipulations is also supported by our data (Fig. 6d).

To assess epithelial–mesenchymal transition (EMT) that is thought to be a prerequisite for NCC migration, we performed IF of EMT biomarker, vimentin, in the context of extracellular matrix components Type I and Type IV collagen (Supplementary Fig. 6a). EMLOs exhibited a demarcated zone of EMT at the base of SC where bottlenecking of SC neurons occurs. By IF staining of PAX7 and PAX3, we validated that EMLO SC identity is largely that of dorsal-intermediate SC (PAX7), whereas PAX3 that marks NCCs originating in the SC were distributed throughout the ME (Supplementary Fig. 6b). As an internal control, non-elongated aggregates had dense, uniform PAX3 distribution. SOX10+ and TFAP2a+ cells were always observed in SC, interpenetrating between neuroectodermal rosettes, concentrated at the base of SC with only a subset observed in ME (Supplementary Fig. 6c).

N-cadherin, absent from the gut tube epithelium proper in EMLOs, was concentrated in SC but also adjacent to the distal anterior aspect of the gut tube, and the IF signal at this region increased as the number of cells populating ME and the ISL1+ ganglion increased (Fig. 7e–h). The proportion of EMLOs that exhibited a prominent peripheral ganglionic structure at day 22 was...
was quantified in the three representative lines (Fig. 7i), indicating high reproducibility in patterning.

**Linking NCCs in EMLOs to in vivo events by scRNAseq analysis.** Soldatov et al. recently described an updated model for NCCs in mouse based on extensive spatiotemporal sequencing analysis (Fig. 8b, d). In this model, NCC ontogeny is categorized into four stages that are: (1) pre-EMT (neural tube roof plate), (2) delamination/EMT, (3) fate-biased migration of progenitors, and (4) lineage commitment to sensory and autonomic peripheral neural cells and mesenchymal cell types. We applied this model that is based on sequential binary cell fate decisions to further characterize the NCCs in EMLOs and performed imaging of peripherin to detect peripheral neuronal axons in the ME compartment of day 40 EMLOs (Fig. 8c, e). A suite of genes implicated in pre-EMT NCCs located in the dorsal neural tube (GDF7, OLIG3, LMX1A, ZIC3) were seen expressed in cluster 15 that has a roof plate phenotype and clusters 4 and 9 that also have a neural tube/SC transcriptional phenotypes. By investigating the distribution of cells from pre-EMT through delamination/EMT (PAK3, HAPLN1, PDGFRα, VIM) to fate-biased migration (ITGA4, HOXB2, MCAM, NGFR), we observed more localization to cluster 14, providing indirect evidence of migratory NCC populations. Sets of genes for lineage commitment to sensory neurons, autonomic neurons, peripheral glial cells, and mesenchyme were also detected, thereby satisfying all four elements of the model. The mesenchymal identity of the ME compartment was also identified as splanchnic mesoderm (FOXF1, GATA4, GATA6, VIM, PDGFRα) that is known to surround the epithelial structures in the trunk (Supplementary Fig. 7). Reciprocal signaling interactions between splanchnic mesenchyme and the gut tube are known to contribute to cardiogenesis and myogenesis. We identified transcript and

![Fig. 4 Quantification of neuron specialization and neurotransmission phenotypes in EMLOs. a Distribution of neural progenitors (TUBB3 OR MAP2, blue) and neurons (TUBB3 AND MAP2, orange) in day 16 H3.3.1 EMLOs by scRNAseq (left). Canonical neuronal differentiation genes are shown (log2Exp) (right). b Proportion of specialized neurons generally classified as motor (MNIX1/TUBB3, blue), sensory (POU4F1/TUBB3, orange), and autonomic (PHOX2B/ASCL1, purple) at day 16. Cluster 10 is expanded. c Proportion of neurotransmission phenotypes generally classified as excitatory (blue), inhibitory (orange), serotonergic (purple), dopaminergic (blue), and cholinergic (green) at day 16. d Inhibitory (GPHN in cyan, GAD65&67 in magenta) and excitatory (PSD-95 in cyan, VGlut1 in magenta) neurotransmission biomarkers in SC at day 40 (N = 2 repeat experiments). Individual scale bars provided.](https://doi.org/10.1038/s41467-021-23294-7)
EMLOs differentiate functional spinal neurons that express the MOR and have opioid-responsive firing. There are few studies using stem cell models of opioid mechanisms yet opioids have important biomedical relevance. We applied EMLOs to calcium imaging studies to investigate the effect of MOR modulation on firing activity in various spinal neuron subtypes (Fig. 9). On day 22, EMLO staining for the MOR (or OPRM1) identified populations of OPRM1+ neurons and highlighted the apical aspect of neural rosettes (Fig. 9a, b top). OPRM1 signal was also distributed along the length of the gut tube lumen (Fig. 9a bottom). GABA-synthesizing enzymes GAD65&67 were expressed in SC in congruence with dorsal-intermediate SC identity and GABAergic neurotransmission that is required for the central antinociceptive effects of exogenous opioids (Fig. 9b bottom)29,30. We dissociated H3.3.1 EMLOs at day 22 using a neural tissue dissociation kit and formed 2D adherent neuronal networks (Fig. 9d). In parallel, we derived putative spinal sensory and motor neurons as positive and negative controls for MOR expression, respectively. Transcription factors BRN3A (sensory) and Nkx-6.1 (motor) were used to validate these neurons (Fig. 9d). 2D adherent neuronal differentiation was initiated on the same day as EMLO formation. Cultures were maintained for ten days after seeding in BrainPhys medium supplemented with BDNF, GDNF, and dibutyryl-cyclic AMP. On day 32, we performed successive calcium imaging experiments using Fluo-4 AM (Fig. 9e, f). Baseline calcium transients were first recorded in BrainPhys medium followed by 1 μM addition of the selective MOR agonist DAMGO in the same culture. Calcium imaging was repeated, followed by the addition of 10 μM of naloxone hydrochloride to outcompete DAMGO at MOR. The percentage of neuronal somata with multiple transients above the threshold over the duration of recordings (1.5 min) was quantified. In EMLO-derived cultures, the percentage of neurons
with multiple transients was significantly reduced from 31 ± 5% (Supplementary Movie 4) to 13 ± 3% (Supplementary Movie 5) (mean ± s.e.m.). After naloxone addition, the percentage of neurons increased to 30 ± 5%. A similar result was observed in spinal sensory neurons, decreasing from 37 ± 4% to 10 ± 2%, and then increasing to 33 ± 7%. Spinal motor neurons did not show a statistically significant difference between conditions, although a slight percentage increase was observed after DAMGO addition. These data support the use of EMLOs as an early developmental model that may be informative to pharmacological studies with agents impacting many of the organs derived from the primitive structures that self-organize in the EMLO system.

**Discussion**

In mammals, the breadth of development occurs in the post-implantation embryo, thereby limiting studies of early informative events required for a comprehensive understanding of human development and disease. Gastruloids offer an ability to study the co-development of interconnected systems and organogenesis...

**Fig. 6 Gut tube formation events impact neuronal patterning.** a–b Dual SMAD small molecule inhibitors LDN 193189 (LDN) and SB 431542 (SB) added before the EMLO elongation phase opposes mesenchymal and endodermal elongation a, and was quantified in b in representative ED-iPSC lines F3.5.2 (**p < 0.0001, t = 4.518, df = 61), H3.3.1 (**p < 0.0001, t = 7.293, df = 61), A2.1.1 (**p < 0.0001, t = 6.723, df = 61) by unpaired two-tailed t test. n = number of EMLOs counted (day 22). N = 3 biological replicates performed with similar results. Eight to 45 fields were analyzed per condition and data were quantified per field. Violin plot statistics are as follows: F3.5.2 (DMSO: max = 40, min = 0, median = 15.48, q1 = 0, q3 = 25.89; LDN + SB: max = 100, min = 0, median = 15.48, q1 = 25, q3 = 66.25); H3.3.1 (DMSO: max = 50, min = 0, median = 0, q1 = 0, q3 = 11.90; LDN + SB: max = 100, min = 0, median = 33.33, q1 = 18.33, q3 = 58.57); A.2.1.1 (DMSO: max = 48.86, min = 0, median = 13.39, q1 = 0, q3 = 20; LDN + SB: max = 100, min = 0, median = 60, q1 = 32.05, q3 = 73.21). c Identification of competing neuronal versus endodermal transcriptional programs in cluster 8 in day 16 H3.3.1 EMLOs by scRNAseq. Canonical endoderm markers FOXA2 and EPCAM were co-expressed with TUBB3 (243 cells). A subset of FOXA2/EPCAM/TUBB3 triple-positive cells (blue) also expressed MAP2 (128 cells, orange) and then further expressed SOX17 (16 cells, green). Cluster 8 is expanded. This phenomenon was previously described in vivo for enteric nervous system development24. d IF of FOXA2 and TUJ1 in day 40 H3.3.1 EMLOs. High magnification Z-slices depict the emergence of FOXA2 + neurons in support of the dual embryonic origin model of the mammalian ENS. FOXA2 and TUJ1 staining on day 40 H3.3.1 EMLOs was performed in N = 3 biological replicate experiments with similar results. Individual scale bars provided. Data reported as (mean ± s.e.m.).
that is not currently addressable by single-organoid methods. To date, gastruloid studies have primarily applied mouse ESCs to dissect polarized developmental processes with spatiotemporal accuracy. To the best of our knowledge, no study using three germ-layer gastruloids has been used to investigate neurons specifically in mouse or human models, and, in particular, the neuronal trunk interactions with other primordial endoderm organ structures such as the primitive gut tube and enteric...
nervous system. The human multi-system EMLO gastruloid is therefore expected to be broadly useful for evaluating neurodevelopmental events impacting early CNS–PNS, endoderm, and neuromuscular co-emergent processes.

We developed the EMLO protocol as a tool for studying co-development of CNS and PNS neurons of the trunk. This approach modifies an existing protocol for neuromuscular trunk organoids (NMOs) that lack endoderm. To accurately initiate...
Fig. 8 NCC trajectories mimic in vivo events in EMLOs. a Cluster 14 scRNAseq annotation as NCC by combined expression of SOX10, FOXD3, and S100B. b Schematic representation of neural tube cross-section depicts NCC in vivo developmental steps: pre-EMT (1), delamination/EMT (2), fate-biased lineage migration (3), and lineage commitment (4) to establish autonomic (yellow), sensory (blue), and mesenchymal (purple) anatomic derivatives. DRG (dorsal root ganglia), PVG (prevertebral ganglia), Ao (aorta). c Peripheral neuron biomarker peripherin (red) in mesoderm-endoderm (ME) compartment ($N = 2$ repeat experiments). Individual scale bar is shown. d Updated NCC model by Soldatov et al. depicts developmental events described in b and the implicated genes. e Transcripts from the NCC processes labeled 1–4 as in b, d by scRNAseq in day 16 EMLOs (log2Exp). We infer that, in general, cells progress from the roof plate (cluster 15) and spinal cord (clusters 4, 9) as they undergo EMT to cluster 14 and express transcripts associated with fate-biased migration. Lineage-committed transcriptional programs are also detected for sensory and autonomic neurons, glia, and mesenchyme.

Developmental polarization events, the gastruloid literature emphasizes the importance of attention to cell number and paracrine signaling5–7, similar to the in vivo gastrula state. The NMO protocol applies 15–30× more cells for initial formation (~4500–9000 cells). Our strategy uses reduced initial cell numbers to achieve the desired length scale and more uniform exposure to external mitogens/morphogens for successful multi-lineage specification. NMP cellular starting material is less restricted by shortening the 2D induction with CHIR and FGF (Supplementary Fig. 1, 2). As well, we transition immediately to rotating culture to introduce additional mechanical cues at the critical early polarization stage. An important advance in our method is the ability to transition from a small multi-germ-layer gastruloid to larger organoid-like integrated structures with segregated compartments linked by neurons and the gut tube that follow developmental principles. This has not been achieved in any other gastruloid study, and is distinct from NMOs that were specifically designed to lack endoderm.

Perhaps, the most remarkable feature that we demonstrate in EMLO gastruloids is the widespread marriage of developing neurons with the self-organizing primitive gut tube, reflective of the enteric nervous system (Figs. 2 and 5). Our data suggest a model (Fig. 10) in which the primitive gut tube acts as a morphological organizer of neuronal fibers. The term “organizer” as we use it here26 refers to regions of an embryo or gastruloid/organoid that can induce other cell fates and direct the pattern formation of other lineages. The temporal events in EMLO formation and high reproducibility are consistent with the gut tube acting as an early self-organizing event. In this regard, it is a bioenergetically prominent feature in EMLOs, and is inundated with neurons that are more compact around this structure versus elsewhere in the EMLO ME compartment (Supplementary Movie 3). Neuronal growth in tangential patterns is typically directed by gradient-driven responses, and we observe the consistent extension of multiple neuronal fibers around the gut tube and neuronal bodies that accumulate as ganglia. The stereotypic, dynamic transformation into neuronal assemblies with peripheral ganglionic structures that occurs in these EMLOs is reproducible. Based on our analysis, we favor the view that this represents a PNS-like region that is, in part, patterned by NCCs (Figs. 7, 8).

Ectodermal NCC ontogeny in development continues to be an exciting new area of discovery. In a recent comprehensive spatiotemporal sequencing study in mouse53, Soldatov et al. described a model for NCC ontogeny that differs from the classical model (Fig. 8). In this model, competing transcriptional programs are established in precursors even prior to EMT and are successively lineage-biased during migration to ultimately achieve a variety of differentiation landscapes and cell types in the periphery. The human EMLO data are consistent with this murine model (Figs. 7 and 8). By fixed time-series imaging and scRNAseq analysis, we infer that ectodermal NCCs originating in the SC compartment are able to populate the ME, effectively as proposed in Soldatov et al.25, by migrating and differentiating throughout the EMLO. Similar to in vivo events, early NCCs co-express lineage-specific transcriptional programs, but commit over time to diverge (Figs. 7 and 8). We summarize these points on ectodermal NCCs in a proposed EMLO model figure (Fig. 10). We have inferred ectodermal NCC migration in this study from analysis of multiple biomarkers in fully intact EMLOs and ME-only subpopulations, as well as the use of scRNAseq for detailed RNA expression profiling. Full validation of migration, however, will require the demonstration of live cell continuous motility. A truncated subset of EMLOs containing ME-only gut tube and mesenchyme does elongate, but lacks NCC biomarkers and TUJ1. These data are consistent with either migration of NCCs from SC into ME, or some contribution from the SC compartment for ME presence of non-ectodermal NCCs.

In the developing CNS, NMPs act at the major bifurcation point in generating the SC and somites54. In vitro, NMP-derived trunk NCCs26,32 have been demonstrated to differentiate into a variety of relevant PNS cell types, including autonomic sympathetic neurons33,34 and cells of the enteric nervous system35, as well as spinal motor neurons and interneurons of the CNS16. NMPs in EMLOs may constitute a transitory stage during the maturation of SC and ME compartments. Although EMLOs are formed with cell starting material co-expressing SOX2 and Bra that is characteristic of NMPs (Supplementary Fig. 2c–e), we no longer detect this population by these biomarkers at day 16 with scRNAseq. We favor the view that the SC region is patterned by NMPs, and that subsequent to initial early-stage polarization events, EMLO elongation is driven by mesenchymal and endodermal processes rather than by NMP-driven axial elongation. This is supported by the pharmacological inhibition of dual SMAD signaling pathways in developing EMLOs. All other gastrointestinal models require sustained NMPs with elongation, whereas, here we observe elongation via endoderm and mesoderm signaling.

Autonomic and enteric nervous systems form integrating neural circuitry with communicating nodes called ganglia in which neuronal cell bodies reside. In EMLOs, highly reproducible ganglionic structures emerge at the base of the gut tube and are comprised of peripheral neurons with NCC-derived expression profiles. By scRNAseq of day 16 EMLOs, we identified TUBB3+ neurons co-expressing ASCL1 and PHOX2B, indicating the presence of developing autonomic neurons. Future investigations into the emergence and progressive distribution and organization of autonomic neurons using techniques to simultaneously apply scRNAseq with cell organization as the EMLOs mature will be informative, including the identification of non-ectodermal derived enteric neurons24,36. This, in addition to the ectodermal origin of NCCs, supports the possibility of two separate neural lineages for the enteric nervous system34. As well, it was recently demonstrated in murine models that cranial ectodermal NCCs undergo an in vivo reprogramming event wherein pluripotency genes OCT4 and Nanog are reactivated prior to non-ectodermal NCC lineage commitment37, further emphasizing the complexity of NCC and trunk neuronal ontogeny. The EMLO model recapitulates dual NCC ectodermal and endodermal embryonic origins of neurons contributing to the human enteric nervous system that was recently described in mice (Fig. 6c, d)24. On day
Fig. 9 Modeling mu opioid receptor modulation in EMLO-derived neuronal cultures. a TUJ1 (cyan) and OPRM1 (magenta) IF in day 22 H3.3.1 EMLO. High magnification images provided (bottom-left) with multi-dimensional view of gut tube (bottom-right). b Top: OPRM1 expression in apical aspect of neural rosettes and basal neurons. Bottom: TFAP2α expression at the basal aspect of GABAergic rosettes. c Fluo-4 AM in dissociated EMLO cultures. High magnification image with example soma ROI (right). d 2D adherent spinal sensory neurons (top, TUJ1 in blue, BRN3A in magenta) and spinal motor neurons (bottom, TUJ1 in green, Nkx-6.1 in red) used as controls for opioid-responsive firing. IF of MOR was performed in N = 3 separate H3.3.1 EMLO formation experiments with similar results. e Examples of calcium transients quantified in dissociated EMLO neuronal soma using Fluo-4 AM. Shown are no calcium transients (none), one transient (single), or multiple transients (multiple) within a 1.5 min acquisition window. f Histogram of percent of neurons with multiple calcium transients. EMLO-dissociated cultures were compared with spinal sensory neurons (positive control) and spinal motor neurons (negative control). Baseline firing in BrainPhys (blue) was compared to addition of 1 μM DAMGO (tan), followed by 10 μM naloxone in the same cultures (light blue). n = number of neurons quantified. Statistical p values are as follows from left-to-right on the plot: **p = 0.0031, ***p = 0.002, n.s. p = 0.766; ***p = 0.001, *p = 0.014, n.s. p = 0.972; n.s. p = 0.593, n.s. p = 0.64, n.s. p = 0.94). Violin plot statistics are as follows: EMLO-derived (BrainPhys: max = 47, min = 13, median = 29, q1 = 29, q3 = 39; BrainPhys + DAMGO: max = 32, min = 7, median = 11, q1 = 9, q3 = 15; BrainPhys + DAMGO + naloxone: max = 47, min = 32, median = 27, q1 = 21, q3 = 39); spinal sensory (BrainPhys: max = 48, min = 27, median = 37, q1 = 28, q3 = 46; BrainPhys + DAMGO: max = 16, min = 6, median = 10, q1 = 6, q3 = 14; BrainPhys + DAMGO + naloxone: max = 59, min = 7, median = 36, q1 = 19, q3 = 44); spinal motor (BrainPhys: max = 35, min = 15, median = 32; BrainPhys + DAMGO: max = 38, min = 23, median = 31, q1 = 25, q3 = 35; BrainPhys + DAMGO + naloxone: max = 38, min = 14, median = 29, q1 = 22, q3 = 34). Data reported as (mean ± s.e.m.). Individual scale bars provided.
16 scRNAseq data, we identified a subpopulation of FOXA2/EPCAM/SOX17 endodermal cells that also co-express neuronal TUBB3 and MAP2 transcripts. It is worth noting that a large number of developmental regulators are now known to be co-involved in both nervous system and gastrointestinal development such as MNX1/HB9, ISL1, CDX2, SOX2, PAX6, FOXA1, FOXA2, NKX-6.1, ALDH1A2, SHH, WNT5A, BMP4, and Hox genes to name several. Human EMLOs may therefore constitute the only current human system capable of enabling the further investigation of ectodermal and non-ectodermal neuronal origins of the enteric nervous system for validation.

Human model systems that might be useful towards understanding the early neurodevelopmental impacts of pharmacologic agents, such as opioids used biomedically, are limited. Very few studies in the literature apply human stem cell models to opioid investigations, and virtually none that include organoids or opioids used biomedically, are limited. Very few studies have analyzed the early neurodevelopmental impacts of pharmacologic agents.

We chose to evaluate the MOR and opioid biomimetic compounds owing to their relevance to the trunk and SC development, although their importance encompasses broader CNS neuronal plasticity and biomedial uses. This analysis is limited by the use of EMLO and dissociated neurons to allow specific neuronal maturation for quantitative comparison of motor and sensory neurons using calcium imaging (Fig. 9). It is proof-of-principle of the potential of EMLOs for modeling neurogenic changes of clinical and pharmacological relevance in regard to integrated CNS, PNS, and other organ systems that have self-organizing precursors in EMLOs. In future studies, the ability to take full advantage of developmental insights from EMLOs or other gastruloid, organoid, and assembloid models is expected as the technology for detailed analysis of 3D structures continues to advance. The rapid pace of signaling events, the resolution needed to capture those events in thick samples, and the need for quantification supports the mixed analysis of both dissociated and intact organoids. The expression of MOR receptors in the membrane in EMLOs and our 2D analysis have not previously been described in the field since human stem cell models are not yet well-established for opioid studies.

In this work, we have not yet explored all optimized outcomes for EMLOs. For example, in earlier work, we showed that different ED-iPSC lines can achieve uniform high-efficiency cardiomyocyte differentiation and contractility when we optimize the CHIR 99021 concentration. This was beneficial when analyzing clonal replicate lines derived from the same donor (e.g., H3.3.1 vs. H3.1.1). The need for optimization was similarly observed by other researchers, such as in the anteroposterior gastruloid study using hESCs, and for line-specific differences to generate NMOs. In this current work, we apply a single protocol to nine lines that successfully generate EMLOs. At last, we were unable to identify commercially available gut tube-specific inhibitors. We recognize that dual SMAD inhibition can inhibit both mesoderm and endoderm, and therefore enhance commitment to neuroectodermal lineages, and so future work will be necessary to more fully elucidate the role of the gut tube as a developmental organizer using specific gut inhibitors.

Together, EMLOs constitute an important bridge to more systemic tissue interactions and innervation studies together with other gastruloid and assembloid models (Fig. 10). The human EMLO system enables the continued dissection of early neurodevelopment and, importantly, continued differentiation within an embryo-like, multi-lineage context. We are also exploring EMLOs in regard to skeletal muscle differentiation and optimization for heart field progression (Supplementary Figs. 8–9). The human spine is integral to CNS–PNS functions and EMLOs are expected to also provide an ability to investigate mechanisms for impaired human CNS neuronal regeneration with respect to other gastruloid, organoid, and assembloid models is expected as the technology for detailed analysis of 3D structures continues to advance. The rapid pace of signaling events, the resolution needed to capture those events in thick samples, and the need for quantification supports the mixed analysis of both dissociated and intact organoids. The expression of MOR receptors in the membrane in EMLOs and our 2D analysis have not previously been described in the field since human stem cell models are not yet well-established for opioid studies.

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insights in modeling organogenesis, systems interconnectivity, and innervation in a multi-lineage platform.

Methods

Human ED-hiPSC lines and culture conditions. Nine ED-hiPSC lines were previously generated in a Paluh laboratory study under an approved ISSCR protocol using commercial de-identified fibroblast cell lines from three individuals obtained from Coriell11 and were extensively characterized by our laboratory 12, 15, 16. Coriell cell lines were obtained under informed consent according to the company website. Catalog numbers are as follows: GM22268 (F3; African American male, foreskin, 1 day); AG08498 (A-2; Asian male, foreskin, 1 day); GM22186 (H3; Hispanic-Latino male, foreskin, 1 day), hiPSC lines were validated for normal karyotype. Details of ED-hiPSC characterization are provided in Supplementary Table 1. hiPSCs were maintained in pluri potency medium mTeSR Plus (STEMCELL Technologies) supplemented with 1× penicillin-streptomycin (P-S) on hESC-qualified Matrigel (1:100 dilution; Corning) at 37°C, 5% CO2. Cells were passaged 1:6 in six-well plates every 4–7 days using Gentle Cell Dissociation Reagent and cryopreserved in mTeSR according to manufacturer’s instructions (STEMCELL Technologies). All lines were expanded and stored at a low passage number.

EMLO gastruloid formation and signal manipulation. Our gastruloid protocol begins with the pretreatment of 2D adherent hiPSC colonies maintained in mTeSR Plus medium. At ~60% confluence, mTeSR Plus was replaced with the N2B27 basal medium supplemented with 3 μM CHIR 99021 (Tocris Bioscience) and 40 ng/ml basic fibroblast growth factor bFGF/FGF2 (R&D Systems). N2B27 basal medium: 1:1 Dulbecco’s Modified Eagle Medium (DMEM)/F-12/neurobasal plus medium, 2% (v/v) B27 Plus supplement, 1% (v/v) GlutaMAX, 1× non-essential amino acids, 1× P-S. Colonies were maintained in pretreatment medium for 2 days, with fresh culture medium replenished every day. On the day of dissociation (day 0), wherein migratory cells become visible at the colony edge, cells were dissociated with 1× accutase/HBSS (Hank’s Balanced Salt Solution) (Ca-Mg free) at 37°C for 5 min followed by manual trituration with a P-1000 pipette. Six-well plates were pretreated with anti-adhering reagent solution (STEMCELL Technologies) for 5 min to reduce cell-to-cell adhesion. Cells were suspended in N2B27 supplemented with 10 ng/ml FGF2, 2 ng/ml HGF. On day 3, the entire volume of medium was replenished and 1× FGF2, 40 ng/ml FGF8 (R&D Systems), 2 μM SB431542, 100 nM LDN 193189, 0.36 U/ml heparin15,16 were added during a 24 h period. Spinal sensory and spinal motor neuron differentiation began the edge of day 10 and EMLOs were quantified at day 22. EMLOs were dissociated to single cells using a cold activated protease method previously described24,28. EMLO single-cell dissociation by cold activated protease for scRNAseq.

Phase contrast imaging, whole-mount IF, and tissue clearing. For phase contrast microscopy, samples were imaged at room temperature directly in culture plates. For IF, samples were acquired with a Zeiss Invertedkopf 40 C (Zeiss, Apochromat, 10x/0.25 Ph1 A-Plan, and 20x/0.30 Ph1 LD A-Plan, 40x/0.50 Ph2 A-Plan) mounted with an Olympus DP22 color camera and cellSens acquisition software. Whole-mount IF preparation was performed as previously described and detailed as follows28. EMLOs were pooled on the day of fixation, rinsed once with 1× phosphate-buffered saline (PBS), and fixed in 10% neutral-buffered formalin solution at 4°C for 2 h. EMLOs were rinsed three times in 1× PB for 5 min. Samples were then permeabilized by three successive incubations in 0.2% Triton X-100 in 1× PBS (PBST) for 20 min at 4°C, and blocked overnight in 1% bovine serum albumin (BSA) in PBST. The next day, samples were distributed to 12-well plates in 1 ml blocking solution per well. Primary antibodies were added to requisite dilutions (see Supplementary Information). Plates were left rocking at 4°C for 48–72 h, then rinsed three times in blocking solution followed by three times in PBST for 5 min at room temperature in 2 ml centrifuge tubes. Species-matched secondary antibodies were incubated 1:1,000 with additional NuclBlue fixed cell stain (Invitrogen) directly in 2 ml centrifuge tubes, rocking overnight at 4°C. For triple-antibody stains, goat anti-mouse or goat anti-rabbit Cy5 secondary antibodies were added the next day after removal of donkey anti-goat AlexaFluor secondary antibody by additional wash steps. All excess secondary antibodies were ultimately removed by two wash steps in blocking solution followed by two wash steps in PBST. Stained and rinsed EMLO samples were equilibrated in 0.1 M phosphate buffer (PB: 0.025 M Na2HPO4, 0.075 M NaH2PO4, pH 7.4) by three successive incubations of 5 min at room temperature. EMLOs were then post-fixed in 10% neutral-buffered formalin for 20 min at 4°C, and rinsed three more times in 0.02 M PB. To clear samples, PB was aspirated and replaced with 200 μl of 88% H2O2 in PBST solution (w/v) on 0/2/4/6/8/10 days. Samples were washed and dried at 4°C for 24–48 h and then mounted on glass slides, sealed in clear nail polish.

Primary antibodies are listed with company, catalog number, and RRID in Supplementary Data 1. The following antibody dilutions in 1% BSA (1× PB) were used for the corresponding antibodies: anti-TFAP2A (4 μg/ml); anti-SSEA4 (4 μg/ml); anti-SOX2 (mouse, 10 μg/ml); anti-SOX17 (5 μg/ml); anti-β-III-tubulin (rabbit, 1:2,000); anti-β-III-tubulin (mouse, 1:1,000); anti-MES132 cocktail (1:1,000); anti-Peripherin (1:200); CD2H/N-cadherin (1:200); anti-ZO-1 (1:1,000); anti-OPRM1 (1:500); anti-GAD65/67/81 (1:1,000); anti-TLX3 (1:1,000); anti-LBX1 (2 μg/ml); anti-PAX2 (5 μg/ml); anti-PAX3 (5 μg/ml); anti-PAX7 (10 μg/ml); anti-LX19 (2 μg/ml); anti-CHX10 (1:500); anti-NKX6.1 (0.5 μg/ml); anti-BRNSA (1:1,000); anti-BMP2/4 (5 μg/ml); anti-desmin (5 μg/ml); anti-Mycosin Heavy Chain (10 μg/ml); anti-cardiac troponin-T (MAb18874). All secondary antibodies were incubated at 1:1,000 in the same buffer.

Images were collected on a Leica confocal TCS SP5 II system in conjunction with Leica Application Suite Advanced Fluorescence software (v2.7.3.9723). The SP5 II system was equipped with 10×/0.30 HCX PL FLUOTAR air, 20×/0.70 HC PL APO CS air or immersion, and 40×/1.25 HCX PL APO immersion objective lenses. Complete or partial Z-stacks were acquired at ~2 μm separation distance.

EMLO single-cell dissociation by cold activated protease for scRNAseq. Day 16 EMLOs were dissociated to single cells using a cold activated protease method previously described28 and essential details are provided here. Dissociation solution 2% Triton X-100 in 1× PBS (Briefer/24 h) containing 100 μM CHIR 99021, 125 μM DAPT (Tocris Bioscience), 1 μM SB431542, 100 nM LDN 193189, 0.36 U/ml heparin15,16 was added during a 24 h period. Cells were washed and pelleted by centrifugation at 1,200 × g for 5 min. The supernatant was completely aspirated. Cells were resuspended in 1× PBS/10% FBS and counted, centrifuged once more, aspirated, and resuspended in 10× cold dissociation solution (BioLife Solutions) that was filtered through a 40 μm cell strainer and then added to a 1.8 ml Nun cryo-storage tube. Cells were frozen at −80 °C and shipped overnight on dry ice to University of Buffalo Genomics and Bioinformatics Core.

Single-cell sequencing and cluster annotation. Samples received by the University of Buffalo Genomics and Bioinformatics Core were immediately stored at −80 °C. Single-cell DNA (BS5 in PBST) was drawn and loaded on a 10× Genomics Chromium platform using the NextGem V3.1 single-cell reagent kit.
The data were reviewed for quality to ensure sufficient sequencing quality reports, fastq (V4.0.0), which associates each transcript to its cell-of-origin and produces sample was demultiplexed using 10× Genomics Cell Ranger analysis software sequencing on the NovaSeq 6000 (28 × 91). Upon sequencing run completion, the KAPA universal qPCR kit. Samples were diluted to 250 pM and denatured for of the libraries was determined by the Agilent Fragment Analyzer and the Roche used to generate Illumina-ready double-stranded DNA libraries. The concentration Genomics Loupe Browser (V5.0.0). Clusters were annotated based on canonical provide another comprehensive list.

EMLO dissociation for 2D neuronal cultures and IF. Day 22 EMLOs (~25 count) were pooled and dissociated using the Neural Tissue Dissociation Kit (P, palpation) according to manufacturer’s instructions (Miltenyi Biotec) and similarly to a previous report. Samples were manually triturated with a P-1000 pipette. Dissociation was confirmed by visual inspection. Spinal sensory and spinal motor neurons used as controls for calcium imaging were derived as detailed above. On day 22 in differentiation, adherent cultures were detached with 1:1 accutase in HBSS and seed into chambered wells of matrigel-coated glassware. At the same time, day 22 dissociated EMLOs were seeded similarly. Parallel cultures were maintained in BrainPhys supplemented with N2, B27, and 10 ng/ml BDNF, 10 ng/ml GDNF, 1 μM ROCK inhibitor. One-half volume of media was replaced every 3–4 days. For IF of adherent spinal sensory and motor neurons, day 32 adherent cultures were rinsed once in 1× PBS and fixed with 10% neutral-buffered formalin for 30 min at 37°C. Samples were rinsed again in 1× PBS, permeabilized for 5 min in 0.1% Triton X-100, and blocked for 30 min in 1% BSA fraction V (1× PBS) at room temperature. Primary antibodies were applied in 1 ml fresh blocking buffer and incubated at 4°C overnight. Samples were rinsed thoroughly in 1× PBS before applying immunoglobulin- and species-matched secondary antibodies for 1 h in the dark with NucBlue fixed cell stain (4°C). Cells were imaged directly in chambered glassware wells. Wide-field fluorescence microscopy was performed using a Zeiss Axios Observer.Z1 inverted fluorescence microscope (×20/0.8 air and ×63/1.4 Oil Plan-Apochromat DIC objectives). Images were acquired using a Hamamatsu ORCA ER CCD camera and Zeiss AxiosvisionRel software (ver. 4.8.2). For adherent cultures, 10–20-slice Z-stacks were gathered at 1 μm separation distance and compressed using the extended focus feature. If necessary, images were adjusted linearly for brightness in Keynote or ImageJ.

MOR modulation and calcium imaging. On the day of live cell calcium imaging, 30 μg Fluo-4 AM cell-permeant dye (Invitrogen) was diluted in 10 μl of 20% pluronic acid in Fluoro-4 AM was diluted 1:1000 in BrainPhys and added to chambered glass wells after rinsing cells once in HBSS. Samples were incubated at 37°C for 30 min. After incubation, Fluoro-4 AM medium was removed, washed once in HBSS and fresh BrainPhys without phenol red was added. Samples were imaged directly on the Zeiss Axios Observer.Z1 system. Time-lapse series were acquired using a 248 nm LED at 200 ms intervals for 1.5–3 min duration. After imaging baseline activity in BrainPhys, 1 μM DAMGO (MOR agonist, 10 mM stock in water; Abcam) was added directly to chamber wells and imaged again. The competitive opioid receptor antagonist naloxone hydrochloride was then added at 10 μM (10 mM stock in water; Abcam). EMLO-dissociated cultures, spinal sensory, and spinal motor neuron samples were imaged in suc- cision to control for time of Fluoro-4 AM cell loading. Time-lapse series were acquired in ImageJ.

Neuromuscular organoids, cerebral organoids, neurospheres, and small molecule manipulation. The protocol for previously reported polarized NMOs was validated using ED-hiPSC lines (Supplementary Fig. 1). In brief, hiPSC colonies were dissociated with accutase and seeded on matrigel as single cells at 100,000/cm² in N2B27 supplemented with 3 μM CHIR, 40 ng/ml FGF2. The medium was changed every day for 3 days. At day –3, uniform NMPs were dissociated with accutase and forced to aggregate in 96-well plates (9–10,000 cells/ well). At the time of dissociation, the culture medium was changed to N2B27 supplemented with 10 ng/ml FGF2, 2 ng/ml HGF, 2 ng/ml IGF-1 (100 μl/well). In all, 50 μM ROCK inhibitor was used at time of aggregation. One-half volume of medium was replaced with 100 μl N2B27 supplemented with 2 ng/ml HGF and IGF-1–24 h later (n = 20). After day 1, aggregates were maintained in N2B27 only. Day 10 organoids were transferred to a 60 mm dish and moved to an orbital shaker (70 rpm). To test the effect of RA on EMLO elongation, 1 μM RA was added to the shaking culture medium on day 2 of EMLO formation and maintained to day 13. In a separate experiment, we converted 2D adherent cultures that were pretreated for three days with 3 μM CHIR 99021 and 40 ng/ml FGF2 to spinal cord neural stem cells (scNSCs) by addition of 100 nM RA and 200 nM of the potent Hedgehog agonist Hh-Ag1.5. After 10 days, neurospheres were generated from adherent scNSCs on the orbital shaker at 75 rpm and were monitored for elongation.

Cerebral organoids were formed using a modified protocol from Trujillo et al. combined with the STEMCELL Technologies STEMdiff Cerebral Organoid and Maturation kit. In brief, ~70% confluent hiPSC colonies maintained in mTeSR Plus were dissociated with Accutase diluted 1:1 in HBSS and transferred to 6-well plates treated with anti-adherence rinsing solution (STEMCELL Technologies) at ~4 × 10⁶ cells/well. Cells were allowed to aggregate for 1 h before sequential quality reports, fastq files, and per-cell-per-transcript count matrices. The data were reviewed for quality to ensure sufficient mean-reads per cell (64,538), a high fraction of reads in cell (80.6%), and confident mapping rates to the Genome (91.6%). Data were then analyzed using the 10x Genomics Loupe Browser (V5.0.0). Clusters were annotated based on canonical biomarkers, provided here as a supplemental file (Source data file). Rossi et al.10 provide another comprehensive list.

Human stem cells ethics statement. No new cell lines were generated in this study. The ED-hiPSC lines used are published and were previously generated in a Paluh laboratory study under an approved ISSCR protocol using commercial de-identified hiPSC cell lines of three individuals obtained from Coriell.11 The Coriell cell lines were obtained under informed consent according to the company website.

Statistical analysis and reproducibility. The detailed protocol for EMLO for- mation is publicly available.10 Raw data were compiled in Microsoft Excel (v16.16.27) and exported to GraphPad Prism (v9.0.1) for plotting and statistical analysis. Data were reported as (mean ± s.e.m) and analyzed using an unpaired two- tailed t test. EMLO cell nuclei were manually counted using ImageJ to quantify IF data using Z-slices at least 20 μm apart to avoid double counting. **p < 0.001, **p < 0.01, *p < 0.05, n.s. not significant (α = 0.05). Power analysis was not performed. Detailed information for each experiment is provided in Results and Figure Legends. Key resources including primary antibodies, chemicals, and other reagents, software, equipment, and commercial kits are provided (Supplementary Data 1), along with the Source date file and scRNAseq gene annotation lists (Supplementary Data 2).

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The authors declare that all data supporting the findings of this study are available within the article and its Supplementary Information files or from the corresponding author upon reasonable request. The raw single-cell sequencing data have been deposited in the Gene Expression Omnibus (GEO) database under accession code: GSE166603. Figures for this manuscript were made in Keynote (v10.3.5) and Adobe Illustrator Creative Commons 2020. Data plots were generated using GraphPad Prism 9. Source data are provided with this paper.

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Author contributions

J.P. and Z.O. conceived of the project and experimental design and analyzed data and wrote the manuscript. Z.O. performed EMLO formation and characterization experiments, expanded hiPSC lines for their differentiation and EMLO formation, and compiled figures and illustrations.

Competing interests

The authors declare no competing interests.

Additional information

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