A new microphysiological system shows hypoxia primes human ISCs for interleukin-dependent rescue of stem cell activity

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1 ABSTRACT

- Background & Aims: Hypoxia in the intestinal epithelium can be caused by acute ischemic events or conditions like Inflammatory Bowel Disease (IBD) where immune cell infiltration produces 'inflammatory hypoxia', a chronic condition that starves the mucosa of oxygen. Epithelial regeneration after ischemia and IBD suggests intestinal stem cells (ISCs) are highly tolerant to acute and chronic hypoxia; however,
- 6 the impact of acute and chronic hypoxia on human ISC (hISC) properties have not been reported. Here
- 7 we present a new microphysiological system (MPS) to investigate how hypoxia affects hISCs isolated
- 8 from healthy human tissues. We then test the hypothesis that some inflammation-associated
- 9 interleukins protect hISCs during prolonged hypoxia.
- Methods: hISCs were exposed to <1.0% oxygen in the MPS for 6-, 24-, 48- & 72hrs. Viability, HIF1α
 response, transcriptomics, cell cycle dynamics, and hISC response to cytokines were evaluated.
- Results: The novel MPS enables precise, real-time control and monitoring of oxygen levels at the cell surface. Under hypoxia, hISCs remain viable until 72hrs and exhibit peak HIF1α at 24hrs. hISCs lose stem cell activity at 24hrs that recovers at 48hrs of hypoxia. Hypoxia increases the proportion of hISCs in G1 and regulates hISC capacity to respond to multiple inflammatory signals. Hypoxia induces hISCs to upregulate many interleukin receptors and hISCs demonstrate hypoxia-dependent cell cycle regulation and increased organoid forming efficiency when treated with specific interleukins
- **Conclusions**: Hypoxia primes hISCs to respond differently to interleukins than hISCs in normoxia through a transcriptional response. hISCs slow cell cycle progression and increase hISC activity when treated with hypoxia and specific interleukins. These findings have important implications for epithelial regeneration in the gut during inflammatory events.

22 KEYWORDS

23 Inflammatory Hypoxia; Intestinal stem cells; Stem Cell Priming; Oxygen Sensor; Cytokines

24 ABBREVIATIONS

25 Inflammatory Bowel Disease (IBD) Intestinal stem cell (ISC) 26 27 Human intestinal stem cell (hISC) 28 Microphysiological system (MPS) integrated Phosphorescent Oxygen Biosensor (iPOB) 29 Near Infrared (NIR) 30 31 2-Dimensional (2D) Occludin (OCLN) 32 organoid forming efficiency (OFE) 33 CellRaft[™] Arrays (CRAs) 34 35 Principal Component Analysis (PCA) Principal Component (PC) 36 Gene Set Enrichment Analysis (GSEA) 37 38 interleukin (IL) 39 Differential gene expression (DGE) endoplasmic reticulum (ER) 40 unfolded-protein response (UPR) 41 42 Innate Lymphoid Cells Type 2 (ILC2s) 43 dithiothreitol (DTT) 44 Expansion Media (EM) polymethylmethacrylate (PMMA) 45 dichloromethane (DCM) 46 poly(2-hydroxyethyl methacrylate) (pHEMA) 47 48 Fluorescence-activated cell sorting (FACS) 49 RNA integrity number (RIN) integrated fluidic circuits (IFCs) 50

51 **INTRODUCTION**

52 Cellular respiration in complex multi-layered tissues relies on a constant supply of oxygen to 53 maintain healthy physiologic states. To this end, the intestinal epithelium maintains physiologic oxygen tension through a highly vascularized tissue that is perfused by a vast network of vessels that terminate 54 55 in capillary beds where oxygen is released to adjacent cells in a local microenvironment.¹ In the crypt-56 villus epithelial architecture, cells can experience marked differences in physiologic oxygen 57 concentrations where crypt-based cells experience higher oxygen levels compared to cells at the villus tips.² Along the crypt-villus axis, a steep oxygen gradient is present wherein microenvironments that 58 are separated by just ~100 cell distances experience a ~10-fold lower oxygen concentration.³ While 59 lower oxygen concentrations within the gradient are tolerated as normal, sudden or dramatic changes 60 in the magnitude and duration of oxygen loss can lead to pathological hypoxia resulting in cell 61 dysfunction and/or death.4 62

63 Most studies that evaluate the impact of tissue oxygen deprivation do so in the context of 64 ischemia-reperfusion injury. This is characterized by an initial ischemic event such as bowel strangulation, aneurysm, or organ harvest for transplantation which induces hypoxia. Subsequent 65 reperfusion of blood flow, and thus oxygen, induces a separate type of oxidative injury caused by free 66 radicals and reactive oxygen species.^{5,6} Hypoxic events can be categorized as acute, which are 67 typically transient events that last on the order of minutes to hours, and chronic hypoxic events that last 68 69 much longer, for at least 24 hours or more.⁷ Inflammatory hypoxia, a type of chronic hypoxia, is caused by the presence of large numbers of immune cells in the submucosal compartment depleting local 70 oxygen supplies.^{4,8,9} Following epithelial injury, a massive influx of immune cells is recruited to prevent 71 pathogens and other luminal contents from causing further damage. This massive influx of immune 72 cells consumes the local oxygen supply to the epithelium^{4,10-13} and can cause vasoconstriction, which 73 further starves the epithelial cells of oxygen.^{3,14,15} Hypoxia due to inflammation can exist in isolated 74 75 microenvironments that might initially go unnoticed and eventually resolve themselves. However, 76 longer periods of inflammation, such as in Inflammatory Bowel Disease (IBD) may compromise the epithelium and normal repair mechanisms ultimately resulting in, or contributing to, mucosal erosions 77 78 and ulcerations.^{8,12} Consistent with this concept, inflammatory responses have been shown to produce 79 localized and chronic hypoxic episodes in the intestinal stem cell (ISC) zone, potentially leading to 80 impaired ISC survival, renewal, and differentiation, ultimately impeding mucosal repair and producing ISC death.^{6,15,16} 81

82 While numerous studies focus on the impact of hypoxia on repair mechanisms involving restitution in differentiated entrocytes^{6,17-21} few studies investigate the impact of hypoxia on ISC 83 properties and ISC-dependent repair mechanisms²², especially in human tissues. Traditionally, human 84 cancer cell lines are used to study acute and chronic hypoxia at the cellular level but transformed cells 85 86 exhibit abnormal proliferation and cell death dynamics and they do not properly differentiate, thus they are poor models of human ISCs (hISCs).²³ Studies in porcine small intestines show that the so-called 87 88 'reserve' ISC (rISC) exhibits high tolerance to hypoxic episodes of up to 4 hours of hypoxia/ischemia^{15,22}, but these studies do not evaluate the impact of chronic hypoxia on hISCs. A 89 highly physiologically relevant human model to study acute ischemia was developed²⁴ and while the 90 91 study primarily characterizes the differentiated epithelium under hypoxia, a lack of apoptosis in the crypt 92 observed after 45 min of ischemia indicates hISCs are highly resistant to short hypoxic episodes.²⁰ 93 Despite the important observations and advances outlined in these in vivo studies, all have been 94 performed either in the context of ischemia-reperfusion injury, precluding the evaluation of hypoxic 95 injury uncoupled from reperfusion injury, or performed under short-duration ischemic conditions (<4 hours), and thus interpretations are limited to acute rather than chronic hypoxia injury. 96

97 Investigating the impact of hypoxia on hISCs has been historically limited by the inability to 98 indefinitely culture hISCs, the lack of high-resolution in vitro platforms where oxygen levels can be tuned 99 and accurately monitored in real time, and ethical considerations of human research that preclude in 100 vivo experimentation. In this study, we engineered a microphysiological system (MPS) that delivers multiple durations and magnitudes of hypoxia to primary hISCs. The MPS allows for precise remote 101 102 monitoring and control of oxygen concentrations in real-time at the cell surface interface. We use the 103 MPS to create controlled durations of acute and chronic hypoxic episodes and evaluate dynamic transcriptomic changes within hISCs under these conditions. We focused our evaluation on chronic 104 105 hypoxic events related to hISC-immune cell cross talk through interleukin signaling that might occur during inflammatory hypoxia. Our findings have broad implications for understanding the role in which 106 inflammatory hypoxia can impact hISC behavior in gross clinical presentation of ischemia-reperfusion 107 108 injury and focal ischemic events that occur in Inflammatory Bowel Disease-related hypoxia.

109 **RESULTS**

110 **A new microscopy-compatible microphysiological system enables precise regulation and real-**111 *time monitoring of oxygen levels at the cell-media interface.*

To address the limitations of sensitivity, accuracy and scale in existing environmental culture systems that create a hypoxic environment, we developed a novel MPS the size of a standard microscope slide where oxygen tension could be tailored to precise levels. The MPS design consists of 2 major structural components, 1) the 3D-printed light- and air-tight container required for phosphorescence detection and hypoxia induction in the closed system and 2) a clear-bottomed acrylic five-well tissue culture plate with each well containing a hydrogel scaffold that facilitates hISC expansion, which all sits inside of the sealed chamber²⁵ (**Fig. 1A,B**, see methods).

119 The MPS was designed as a closed system in which ambient air flowed over each well via 120 continuous delivery of mixed gases. A biocompatible integrated Phosphorescent Oxygen Biosensor (iPOB) was placed on the hydrogel scaffold surface to provide real-time oxygen measurements 121 specifically at the cell surface interface (Fig. 1C)²⁶. Oxygen levels were quantified by the iPOB using 122 Near Infrared (NIR) phosphorescence lifetime fluorimetry which measures the oxygen-dependent 123 124 decrease in phosphorescence lifetime to determine local oxygen concentration (Fig. 1C,D)²⁷. The 125 iPOBs are reusable, highly sensitive, and capable of detecting a broad range of oxygen levels created by an off-chip gas mixer²⁸ (Fig. 1D, see methods). 126

The MPS design facilitates easy access to refresh media, add compounds, and perform a variety of downstream assays including microscopy, transcriptomic analysis, and immunostaining. The MPS is fabricated from relatively inexpensive commercially available materials and can be sterilized and reused many times. Notably, multiple MPSs can be connected to the same gas source to accommodate complex experimental designs that require many perturbations and technical replicates (see methods). Together, these demonstrate an ideal model system for evaluating the effect of hypoxia on hISCs.

133 The MPS supports culture of hISC/progenitor epithelial monolayers

We recently developed methods that support indefinite 2-Dimensional (2D) expansion of proliferative human intestinal stem cells from primary small intestine and colon²⁵. Under normoxic conditions (i.e., 37°C, 5% CO₂, and remainder atmospheric air), hISC monolayers can be cultured on a defined hydrogel and media formulation that supports hISC expansion while repressing differentiation^{25,29}. This method was used to expand primary jejunal hISCs isolated from small intestinal

crypts of an organ transplant donor and then transferred to the MPS. Initial characterization of the
 proliferative hISC monolayer was performed in the MPS under normoxic conditions by immunostaining.
 The monolayers demonstrated DNA synthesis as measured by S-phase marker EdU, expression of
 hISC/progenitor cell marker SOX9^{30,31}, and epithelial cell tight junction protein Occludin (OCLN) (Fig.
 These results demonstrate the MPS is biocompatible with primary hISCs that are consistent with
 proliferating hISC/progenitor cell populations.

145 Next, the baseline functional stemness of proliferative hISCs cultured on the MPS under normoxia was evaluated. hISCs were cultured on the MPS as proliferative monolayers under 146 147 atmospheric O₂ levels, dissociated to single cells and plated in Matrigel[™] on a CellRaft[™] Array to assess organoid forming efficiency (OFE), a measure for in vivo hISC activity³²⁻³⁴. Single cells in the 148 microwells were identified, quantified, and putative hISCs were individually tracked over a 6-day culture 149 period to determine whether an organoid formed. The data demonstrate ~4% of single cells cultured in 150 151 the MPS under hISC expansion conditions generated an organoid at each time point measured (2, 4, and 6 days after plating) (Fig. 1F), which is consistent with OFE of FACS-isolated hISCs from a Lgr5-152 153 EGFP expressing reporter gene mouse and also consistent with human organoid-derived hISCs cultured for the same period of time in synthetic matrices^{34,35}. These findings demonstrate that hISCs 154 cultured as 2D monolayers on the MPS maintain hISC properties over time, and at similar ratios to 155 156 those observed in vivo and in 3D organoid systems.

157 To define the speed at which severe hypoxia could be achieved and maintained in the MPS, an iPOB was added to the epithelial cell surface, and real-time oxygen levels at the cell-media interface 158 were measured for 24 hours (Fig. 1G, inset). Oxygen concentrations of <1% (below 10 µM) are 159 160 considered severely hypoxic but not anoxic³⁶. The data show the MPS produced a rapid hypoxic environment of 0.3% O₂ (3 µM) within ~30 minutes, which was constantly maintained during the 24-161 162 hour experiment (Fig. 1G). Interestingly, when proliferative hISC monolayers were cultured in normoxic conditions as a control, cellular respiration reduced oxygen levels at the cell-media interface from 18% 163 (180 µM) to 3.5% (35 µM) in the MPS. These data demonstrate that physiologic O₂ metabolism of 164 proliferative hISCs produce a steady-state flux of O₂ in just 30-minutes that is 5-fold less than 165 166 atmospheric oxygen levels.

167 Human ISC monolayer cultures demonstrate peak HIF1 α response at 24 hours of severe 168 hypoxia with no loss of stem cell activity after 48 hours of severe hypoxia

The HIF1 α transcription factor¹³ is a master regulator of the hypoxic response conserved across 169 170 all tissues and species.^{14,37} Under normoxia, HIF1 α is constitutively expressed but is immediately targeted for degradation¹⁴. By contrast, under hypoxia, HIF1 α degradation is inhibited through well-171 established mechanisms, resulting in rapid accumulation and regulation of a broad range of 172 173 downstream target genes that regulate cell survival, proliferation, metabolism, and cell migration^{13,38}. Immunostaining for HIF1 α in hISCs under durations of hypoxia shows a significant accumulation of 174 175 HIF1 α only at 24 hours of severe hypoxia (Fig. 2A,B). No significant change in HIF1 α was observed at 6 hours of hypoxia compared to normoxia, which may indicate tolerance of proliferative hISCs to 176 shorter hypoxic episodes. Lack of HIF1 α accumulation at 48 hours of hypoxia is likely due to known 177 dampening of the HIF-response that occurs after cells re-equilibrate to tolerate new and sustained low 178 O_2 levels.⁴ Interestingly, hISCs cultured under normoxia showed a significant increase in HIF1 α levels 179 180 24 hours after media change when compared to the 6 hour normoxic samples. This minor hypoxic response is consistent with previous observations that cells in submersion cultures experience a 181 182 localized hypoxic environment due to the slower rate of diffusion of O₂ through medium relative to the

rate of consumption in cellular metabolism.³⁹ Together, these findings established a timeframe for further investigation into the mechanisms by which hISCs resist death and preserve functional stemness and viability in response to HIF1 α accumulation in hISCs during short-, medium-, and longterm hypoxia.

To confirm whether proliferative hISCs in the MPS were experiencing a HIF-response at 0.3% 187 188 O₂, qPCR was performed to detect expression of canonical HIF1 α target genes that are typically upregulated during hypoxia. After 24 hours of hypoxia, HIF1 α demonstrated little transcriptional change 189 190 as expected since its activity is mediated by post-transcriptional stabilization (Fig 2C)^{40,41}. By contrast, the set of classic HIF1α target genes (VEGF, SCL2A1/GLUT1, BCL2, BNIP3) were all significantly 191 upregulated compared to normoxic controls (Fig. 2C)⁴²⁻⁴⁴. These findings demonstrate that an O₂ 192 193 tension of 0.3% (3 µM) for 24 hours is sufficient to activate a canonical hypoxia response inside 194 proliferative hISCs.

Next, we sought to explore the extent to which hISCs can survive severe hypoxia. hISCs were exposed to severe hypoxia (<1% O₂ as measured by the iPOB) in the MPS for 6, 24, 48, or 72 hours and viability was evaluated by staining for early apoptosis marker, Annexin V (**Fig. 2D**).⁴⁵ Flow cytometric quantification of Annexin V negative cells demonstrates ~90% viability through 48 hours of severe hypoxia. However, at 72 hours of severe hypoxia, there was a significant decrease in viability as measured by Annexin V-negative staining, from ~90% viable in normoxic controls to ~70% viable after 72 hours of hypoxia.

202 To determine how longer episodes of hypoxia would impact hISC activity, monolayers were 203 exposed to 24 hours or 48 hours of hypoxia or normoxia as a control in the MPS. At 24 and 48 hours of hypoxia, hISCs demonstrate persistent expression of the proliferation marker, KI67, albeit nuclear 204 205 KI67 levels are reduced at 48 hours of hypoxia (Fig 2E). To quantify hISC activity after these durations 206 of hypoxia, monolayers were subjected to 24 or 48 hours hypoxia, dissociated to single cells, and then applied to CellRaft[™] Arrays (CRAs) for high-throughput accurate guantification of clonal OFE.³⁴ CRAs 207 208 enable up to ~10,000 hISCs to be plated in microwells on a small device and individual hISCs can be evaluated for OFE over time^{34,46}. After 24 hours of hypoxia OFE decreased ~2.5-fold (Fig. 2F). 209 210 Interestingly, when hISCs were exposed to a longer 48-hour hypoxia duration, the OFE was 211 indistinguishable from normoxia controls suggesting hypoxia-induced survival mechanisms were 212 invoked in hISCs after 24 hours of hypoxia (Fig. 2G).

213 To evaluate the time-dependent gene expression changes occurring in response to hypoxia and 214 HIF1a accumulation, RNA-sequencing was performed on proliferative hISCs cultured under normoxia 215 and severe hypoxia for 6, 24, and 48 hours (Fig. 2H). Principal Component Analysis (PCA) demonstrated high agreement between technical replicates (n = 3) for each timepoint and revealed 216 217 global transcriptomic changes in hISCs between normoxic and hypoxic samples. The 6-hour and 48-218 hour normoxic samples clustered closely on the first Principal Component (PC), whereas the 24-hour 219 normoxic samples deviated from the other normoxic samples on the first PC. Interestingly, the 48-hour 220 normoxic samples deviated from the 6-hour normoxic samples on the second axis, which is consistent 221 with our previous data that hISCs mount a transient HIF-response at 24 hours following media change 222 that resolves after 48 hours in culture (Fig 2A,B). These results suggest that normoxic hISCs 223 experience a minor hypoxic event following media change that results in a subtle, but meaningful, shift 224 in gene expression profile in line with a hypoxic response and that following this shift, the hISCs return 225 to a slightly altered gene expression profile.

226 For hypoxic samples, the 6-hour samples clustered closest to the 24-hour normoxic samples, 227 which is consistent with the previous observation that the 24-hour normoxic hISCs were undergoing a 228 minor HIF-response. By contrast, the 24-hour and 48-hour hypoxic samples clustered together but 229 deviated significantly from all other samples on the first PC, indicating the onset of a strong HIF-230 response in hISCs at 24 hours of severe hypoxia that persists through 48 hours of hypoxia. The 231 transcriptomic shift at 24 hours of hypoxia is aligned with the robust HIF1 α accumulation and significant 232 decrease in OFE after 24 hours of hypoxia, reinforcing the concept that hISCs tolerate shorter periods 233 of hypoxia through at least 6 hours but require transcriptional changes at 24 hours to regain organoid 234 forming activity at 48 hours (Fig. 2F,G).

235 To determine if durations of severe hypoxia promoted differentiation, transcriptomic analysis was performed to evaluate the expression of hallmark lineage markers for hISCs and main classes of 236 differentiated cells (Fig 2I).47,48 Consistent with decreased OFE at 24 hours of hypoxia, hISCs show 237 238 decreased expression of stem and progenitor marker genes. Interestingly, this decreased expression 239 continues through 48 hours of hypoxia, even though OFE is restored to normoxic levels, which is consistent with the changes seen in our PCA (Fig 2H). The data show very low expression and no 240 241 appreciable trends in hypoxia-associated changes of genetic markers for Absorptive Enterocytes (ALPI, 242 SI) and secretory lineages, Paneth- (DEFA5), Enteroendocrine- (CHGA, CHGB), Tuft- (PTGS1), 243 Goblet- (MUC2), and BEST4 cells (BEST4) compared to markers of proliferative hISCs and progenitor 244 cells. These data indicate that hypoxia alone does not promote differentiation of hISCs.

Hypoxia primes an hISC transcriptional state to respond to extrinsic interleukin signals and intrinsic downstream mediators of inflammation.

247 To investigate the potential molecular and physiological context associated with these time-248 dependent transcriptomic changes, Gene Set Enrichment Analysis (GSEA) (Fig. 3A) was used to 249 evaluate the comprehensive gene expression changes observed in our RNA sequencing data. GSEA 250 demonstrates that the hypoxia-related gene set is highly enriched at all timepoints compared to controls, providing confidence in the pathway-based analysis tools^{49,50} GSEA findings indicate hISC 251 252 monolayers regulate a subset of pathways consistently across all hypoxia durations while other 253 pathways are regulated in a dynamic manner. Specifically, transcriptomic analyses on hISC monolayers revealed enriched gene sets for inflammatory signaling through specific interleukin-254 mediated pathways, (e.g., Inflammatory Response, IL6 JAK STAT3 Signaling, IL2 STAT5 Signaling, 255 and INF α/γ Responses) (Fig. 3A). 256

257 Within the gene set for 'Hallmark Inflammatory Response' ~4.5% of the genes (8/184) were interleukin (IL) receptors and all demonstrated some form of hypoxia-dependent regulation, thus we 258 259 focused our analysis on all IL receptors^{4,10,14} with the goal of revealing how hypoxia-related inflammatory conditions might extrinsically signal to hISCs (Fig. 3B). Interleukins and their cognate 260 receptors have well established roles in regulating stem cell proliferation in diverse tissue types⁵¹⁻⁶⁰. Of 261 45 IL receptors, 91% had detectable expression across all durations of hypoxia. Differential gene 262 263 expression (DGE) between normoxic and hypoxic environments revealed a subset of 13 IL receptors 264 (IL1R1, IL1R2, IL1RAP, IL1RAPL1, IL2RG, IL4R, IL11RA, IL13RA1, IL17RA, IL17RC, IL17RE, IL18R1, 265 IL20RA) that were regulated by hypoxia in a time-dependent fashion, potentially indicating specific roles 266 during early (6 hours), mid (24 hours), and late (48 hours) durations of severe hypoxia.

To compare hypoxia-dependent changes in IL receptor expression, we looked at fold change differences in gene expression for the 45 IL receptor genes observed in our dataset. IL1RAPL1, IL1R2,

269 IL10RB, IL22RA1, IL6R, IL11RA, IL17RE, IL1RN, IL2RG, IL4R, IL17RA, and IL17RB were considered early responders, as all showed hypoxia-dependent up- or down-regulated expression at 6 hours (Fig 270 271 4A). Of the early responders IL1RN, IL6R, IL10RB, IL17RB, and IL22RA1 were considered early 272 persistent responders as their hypoxia-dependent expression changes were consistent through 24 and 273 48 hours of hypoxia (Fig 4A-C). IL1RAP, IL1R1, IL18R1, and IL20RA were considered mid-responders 274 as their regulation patterns changed beginning at 24 hours of hypoxia. Of these, only IL1RAP had 275 significant change in regulation through 48 hours. IL13RA1 and IL17RC were considered late 276 responders as their hypoxia-dependent gene expression patterns were only observed following 48 277 hours of hypoxia. IL1R2 and IL11RA were considered dynamic responders as their expression patterns 278 changed at 6 and 48 hours of hypoxia. The significant trends were verified by qPCR for the subset of 279 IL receptors that were consistently upregulated (IL6R, IL10RB, IL22RA1) or downregulated (IL17RB) 280 (Fig. 4D). To confirm that hISCs expressed the IL receptors in vivo and in the MPS culture system, we 281 compared IL receptor expression from our recent single cell transcriptomic atlas of the human intestine 282 to single cell transcriptomic dataset for hISCs cultured as monolayers in media that supports hISC 283 expansion (Fig. 4E,F).^{48,61} The data show that all members of the subset of hypoxia-regulated IL 284 receptors have measurable expression in hISCs both in vivo and in vitro. These results show hISC 285 undergo time-dependent changes in IL receptor expression during acute and chronic hypoxic events 286 that primes hISCs to respond to extrinsic inflammatory signals.

287 Hypoxia increases proportion of hISCs residing in G1 phase

288 As a baseline, in the absence of interleukins, we determined the impact of hypoxia on regulating 289 proliferation and cell cycle phases (Fig 5A). hISC monolayers were pulsed with EdU for the last 2 hours 290 of culture to quantify cells in S-phase during normoxia or hypoxia, then co-stained for general cell-cycle marker KI67 (Fig 5B). Individual nuclei were segmented using Cellpose and the percent of cells in S-291 phase (EdU⁺) or in any cell cycle phase (KI67⁺) were quantified.^{62,63} Confocal images were used to 292 train the algorithm for precise nuclear segmentation (Bisbenzimide+). Approximately 150,000 nuclei 293 294 were automatically identified (i.e., segmented) across all experimental conditions (Fig 5B).62,63 After 295 segmentation, KI67 and EdU channels were overlaid onto the automatically segmented nuclei to 296 quantify the mean fluorescence intensity and percent of cells positive for each cell cycle marker on a 297 per-nuclei and per-colony basis. The number of cells in mitosis (M-phase) were evaluated by counting 298 condensed chromosomes that were stained by KI67 and morphologically identifiable (Fig 5C).

299 To determine whether hypoxia caused a specific arrest or delay in any particular cell cycle phase, 300 we leveraged a recent study which shows that KI67 expression levels can reliably predict cell cycle 301 phases.⁶⁴ KI67 levels can serve as a marker for cell cycle progression and exit, with the lowest levels 302 of KI67 in G1/G0 phase, intermediate levels during S-phase, and highest levels in G2/M phase⁶⁴. We 303 evaluated the levels of KI67 in hISCs for segmented nuclei from ~38,000 cells and found that severe hypoxia significantly decreased KI67 under severe hypoxia (Fig. 5D). Hypoxic hISC cultures showed 304 approximately 67.2% of cells undergoing cell cycling (KI67⁺), a significant decrease compared normoxic 305 conditions where 78.9% of cells reside in the cell cycle (Fig 5E).⁶⁴ More granular analysis revealed 306 307 reductions in the number of cells in S-phase (EdU⁺) and M-phase (mitotic bodies) (Fig 5E). The lower 308 levels of KI67, combined with the significant decrease in S- and M-phase cells, suggests that 24 hours of severe hypoxia likely causes either G1 arrest or lengthening of G1. Transcriptomic data does not 309 310 show increases in differentiated lineage markers (Fig. 2I) indicating that hypoxia is not pushing hISCs to differentiate as they exit the cell cycle. Together, these findings demonstrate that over 24 hours of 311 hypoxia, hISCs negatively regulate progression through the cell cycle and that hypoxia likely promotes 312 313 G1 dormancy in hISCs.

314 Hypoxia alters hISC cell cycle regulation by specific interleukins

We next explored whether the hypoxia-regulated IL receptors could impact hISC cell cycle when 315 316 in the presence of their cognate interleukins: $IL1\alpha$, $IL1\beta$, IL2, IL4, IL6, IL10, IL13, $IL17^{A/F}$ (heterodimer). IL22, and IL25. hISC monolayers were treated with ILs under normoxia or hypoxia for 24 hours, to 317 simulate ILs that might be present during an chronic inflammatory response, before fixing, staining, and 318 319 imaging^{8,12,65-68}. No ILs caused significant impacts on the total proportion of proliferating cells (KI67⁺) 320 after 24 hours of hypoxia; however, at 24 hours of normoxia or hypoxia, hISCs shift cell cycle phase dynamics in response to interleukin signals (Fig 6A-F). IL1a increases the proportion of hISCs in S-321 322 phase in normoxia, but this effect is largely abrogated in hypoxic hISCs (Fig 6A). IL2, IL4, and IL25 323 significantly increase the percentage of hISCs in M-phase only in hypoxia, with no change noted in normoxia (Fig 6B-E); whereas IL22 affects normoxic hISCs, but not hypoxic (Fig 6F). IL6, IL10, IL13, 324 and IL17^{A/F} had no effects on cell cycle phase dynamics (Fig. 7A-D). The IL1 α -dependent decrease in 325 S-phase cells with no increase in M-phase hISCs suggests that IL1 α further promotes G1 arrest or 326 elongation under hypoxia, thereby reducing observable S-phase cells. IL2, IL4, and IL25-dependent 327 increases in M-phase hISCs with no observable decrease in KI67⁺ cells suggest slow progression of 328 hISCs through M-phase under hypoxic conditions. These indicate that hypoxia primes hISCs to respond 329 330 to ILs differently then hISCs in normoxia

331 Hypoxia affects hISC activity in response to interleukin treatment

332 While the previous experiments were designed to focus on the impact of hypoxia-333 dependent IL receptor priming on proliferation dynamics, we next sought to probe whether this IL 334 receptor priming would regulate stem cell activity in the presence of cognate interleukins. hISC monolayers were exposed to 24 hours of normoxia or severe hypoxia, then dissociated to single hISCs 335 336 and plated in conventional 3D organoid assays to evaluate OFE of clonal hISCs in the absence or 337 presences of ILs (Fig 8A). Similar to OFE assays in the CellRaft arrays (Fig 2F), we found that hypoxia induced a decrease in OFE at 24 hours (Fig. 8B,C). IL6, IL10, and IL13 (Fig 8D-F) increased OFE in 338 339 a hypoxia-independent manner, while IL1β, IL2, IL4, and IL25 demonstrated hypoxia-dependent increases in OFE (Fig 8G-J), thereby rescuing the negative impacts of hypoxia on hISC. No effects on 340 OFE were observed for IL1 α , IL17^{A/F}, and IL22 (**Fig 8K-M**). Together, these findings show 24 hours of 341 342 hypoxia primes hISCs to respond to cognate ILs differently in hypoxia than in normoxia by altering cell 343 cycle phase dynamics and in some cases enhancing hISC activity.

344 **DISCUSSION**

345 Human health conditions that cause loss of oxygen to the intestinal epithelium result in rapid loss of the differentiated cells, requiring quick regeneration from the stem cell pools located at the base 346 347 of the crypts^{6,69,70}. Evaluating the impact of hypoxia on hISCs in vivo is challenging due to ethical considerations and the inability to precisely control many experimental variables. While in vitro systems 348 349 are available to regulate ambient oxygen levels, these platforms are limited in that oxygen levels are monitored in large volume incubators and not localized to the cell-medium interface. Here we developed 350 351 a flexible and oxygen-tunable MPS that addresses many limitations of conventional systems and represents a next-generation platform to study hypoxic injury across a broad range of cell and tissue 352 353 types. While not explored in this study, the flexible design and small scale of the MPS will enable co-354 culture of hISC with other cell types, such as immune cells, which are key players in hypoxia-related 355 intestinal conditions.

356 Murine intestinal epithelial monolayers were shown to experience repeated bouts of hypoxia during conventional cell culture methods.³⁹ Upon media changes, oxygen levels were at saturation but 357 the levels substantially reduced in media over time, ostensibly by cellular metabolism, thereby creating 358 a hypoxic environment and inducing a HIF1 α response.³⁹ These observations indicate that oxygen 359 diffusion from ambient incubator gas mixes was insufficient to compensate for the oxygen metabolism 360 of mouse intestinal epithelial cells submerged in culture media.³⁹ Using our newly developed MPS that 361 supports 2D cultures of hISCs, we show similar findings in that fresh media at the cell surface interface 362 is saturated at about ~12% O₂ from atmospheric gasses and that hISCs metabolize O₂ levels in the 363 364 media from $\sim 12\%$ to $\sim 3\%$ in approximately 3 hours. At that point, O₂ levels stabilize but are sufficiently 365 low to induce a small but appreciable hypoxia-induced HIF1 α response after 24 hours of submersion culture. Our findings challenge general assumptions regarding oxygen availability following regular 366 367 medium changes in many conventional cell culture systems. More specifically, our data suggest that 368 any cells cultured in submersion culture can experience unappreciated HIF1 α -dependent 369 transcriptional changes following media changes, confounding interpretations.

370 Our findings show that hISCs are highly resilient to severe hypoxia for up to 48 hours. There 371 was a decrease in functional hISC activity at 24 hours of hypoxia, but this resolved at 48 hours with no significant difference in OFE compared to normoxic controls. Several studies in rodent and human 372 intestines highlight the role of endoplasmic reticulum (ER) stress and the unfolded-protein response 373 (UPR) during acute hypoxic episodes.^{39,71,72} The UPR is considered to mediate cell death by activating 374 375 the intrinsic apoptotic cascade; however, several more contemporary reports in the context of cancer 376 cell lines indicate that unresolved ER-stress, which can occur during long durations of hypoxia⁷³, can induce necroptosis.⁷³⁻⁷⁵ We found no clear evidence of increased apoptosis during 48 hours of severe 377 378 hypoxia, and while necroptosis pathways were not evaluated, there was no decrease in hISC activity 379 at 48 hours of hypoxia, strongly supporting that other cell death pathways are being repressed in hISCs 380 through 48 hours of hypoxia.

381 Transcriptomic profiling of hISCs during chronic and severe hypoxia revealed enrichment of regulated genes related to inflammation and inflammation responses. Among these genes we noticed 382 383 many cytokine/interleukin receptors were significantly regulated in a hypoxia-dependent manner. While 384 the interleukin responses mediated through these receptors impact many different aspects of cell 385 biology, we focused our analysis on how hypoxia-induced changes in hISC expression of IL receptors, 386 could impact cell cycle dynamics and stem cell activity through stimulation using their cognate ILs. In 387 the absence of ILs, the baseline hISC response to 24 hours of hypoxia was a significant decrease in the overall number of cycling cells (KI67⁺) with notable reductions in numbers of cells in each phase of 388 389 the cell cycle. While the number of cycling cells was reduced at 24 hours of hypoxia based on KI67, 390 the lack of differentiated cell markers and absence of reduced OFE of hISCs at 48 hours suggests hISCs do not exit cell cycle or differentiate as the result of severe hypoxia but rather enter a 'safe mode' 391 392 characterized by a reversable dormant state, reminiscent of a reserve ISC state⁷⁶. Further genetic and 393 molecular analysis will be necessary to reveal these mechanisms.

394

395 During inflammatory hypoxia, local immune cells express a complex mixture of ILs near hISCs.⁸ In our model, hypoxia primed hISCs to respond to subsets of ILs by regulating cognate IL receptors. 396 397 Stimulation of hypoxia-regulated IL receptors by their cognate ILs produced notable hypoxia-dependent increases in G2/M-phase cells for IL2, IL4, and IL25 that generally rescued the baseline hypoxia-398 dependent loss of G2/M-phase cells. Less notable, IL1 α caused a modest but significant reduction in 399 400 S-phase cells. There is a collection of studies describing how IL2 and IL4 promote proliferation in immune cells^{53,56,77,78}, and studies in colon and pancreatic cancer cells demonstrate that IL4 enhances 401 proliferation^{79,80}, but we found no studies describing the role of these ILs on mouse or human ISCs. 402 IL25 is expressed by tuft cells during parasitic worm infections.⁸¹⁻⁸⁴ Secreted IL25 interacts with the 403 cognate heterodimer receptors, IL17RA and IL17RB⁸⁵, on Innate Lymphoid Cells Type 2 (ILC2s), which 404 signal back through IL-4 and IL-13 to the epithelium to aid worm clearance.81,83,84,86 IL17RA/B 405 expression has not previously been reported in hISCs, suggesting a novel role for IL25 signaling in 406 407 hISCs.

Under 24 hours of hypoxia, hISCs demonstrate less active progression through the cell cycle 408 and lower organoid forming efficiency. hISCs show increased stem cell activity in response to IL6, IL10, 409 410 and IL13, regardless of hypoxic conditioning. On the other hand, hypoxia induces hISCs to respond to IL1β, IL2, IL4, and IL25 by increasing stem cell activity in a hypoxia-dependent fashion. IL2 and IL4 411 promote cell survival effects by stimulating the Akt-pathway⁸⁷, and although there are fewer clear links 412 413 between IL25 (IL17RA/B) and cell survival, the IL17RA/B receptors can stimulate NF-kB activation^{88,89}, which has cell survival effects under some stress stimuli.⁹⁰ While hypoxia alone can stimulate NF-kB 414 activation through HIF1 $\alpha^{8,91}$, IL25 could amplify NF-kB activation to increase hISC survival during 415 severe and prolonged hypoxic episodes. Interestingly, IL2, IL4, and IL25 stimulated increases in G2/M 416 417 in hISCs under hypoxia, with no change seen in hISCs under normoxia. G2/M arrest or delay by these three cytokines might protect cells from undergoing checkpoint-induced cell death as the completion of 418 cytokinesis could be compromised by the lack of energetic and metabolic resources under severe 419 420 hypoxia.⁹² While many ILs have cell cycle effects on hISCs under normoxia, we see that many of these 421 effects change and produce novel responses that are only seen in the context of hypoxia. As such, it 422 appears hypoxia-primed hISCs likely respond differently to immune stimuli than normoxic hISCs, laying 423 the foundation for studies into unique, hypoxia-dependent genetic and biochemical mechanisms, which preserve and restore hISC activity during and after hypoxic injury. 424

425 We acknowledge that deconstruction of the complex microenvironment of the in vivo human stem cell niche limits some physiological interpretations. However, second generation studies using 426 427 our MPS can build on the findings of this study to test complex mixtures of hypoxia-related cytokines, 428 interactions of multiple cell types, and tunable O_2 concentrations to more accurately model and study 429 hISC behaviors under inflammatory hypoxia, more acute forms of oxygen deprivation, and clinically relevant reperfusion. Previous iterations of the MPS have shown the ability of the MPS to dynamically 430 431 'reperfuse' hypoxic cultures with O₂, enabling studies of other modes cell death, such as ferroptosis, which likely have a role in reperfusion injury due to reactive oxygen species production^{26,93,94}, or 432 433 pyroptosis, which are relevant to inflammasome-mediated cell death and are complex to model as they involve microbes and immune cell components.^{95,96} As such the MPS is currently being optimized for 434 435 co-culture of hISCs, immune cells, and anaerobic commensal or pathological microbiota with the goal of creating a more physiologically relevant environment to explore many facets that impact hISC 436 437 function and behavior.

438 MATERIALS AND METHODS

439 **Primary human crypt isolation and intestinal epithelial stem cell culture**

440 A surgical specimen of human small intestine (jejunum) was obtained from a donor at UNC Hospitals with consent of the patient (under the approved protocol UNC IRB #14-2013). Villi and crypts 441 were detached from the specimen by incubation in a chelating buffer for 75 min at 20°C followed by 442 443 vigorous shaking in a 50 mL conical tube. The chelating buffer was composed of EDTA (2 mM), dithiothreitol (DTT, 0.5 mM, freshly added), Na₂HPO₄ (5.6 mM), KH₂PO₄ (8.0 mM), NaCl (96.2 mM), 444 KCI (1.6 mM), sucrose (43.4 mM), D-sorbitol (54.9 mM), pH 1/4 7.4.97 Released crypts were expanded 445 as a monolayer on a neutralized collagen hydrogel as described previously.²⁵ Briefly, crypts were 446 placed on the top of 1.0 mg/mL collagen hydrogels (1 ml into each well of 6-well plate (T1006; Denville, 447 Holliston, MA)) at a density of 5,000 crypts/ well and overlaid with 4 ml of Expansion Media (EM) 448 449 containing 10 mmol/L Y-27632 (S1049; SelleckChem). EM contains a mixture of advanced Dulbecco's modified Eagle medium/F12 medium (12634010; ThermoFisher) and Wnt-3A, R-spondin 3, noggin 450 (WRN) conditioned medium (WRN medium prepared in lab from L-WRN cells (CRL- 3276; ATCC) 451 following a published protocol⁹⁸ at a volumetric ratio of 1:1, and supplemented with GlutaMAX 452 (35050061; ThermoFisher), B27 supplement without vitamin A (12587010; ThermoFisher), 10 mM 453 HEPES (15630-080; ThermoFisher), 1.25 mM N-acetyl cysteine (194603; MP Bio, Santa Ana, CA), 10 454 mM nicotinamide (N0636; Sigma-Aldrich), 50 ng/mL epidermal growth factor (315-09; Peprotech), 2.0 455 nM gastrin (AS-64149; Anaspec), 10 nM prostaglandin E2 (14010; Cayman Chemicals), 3.0 µM 456 SB202190 (S1077; Selleckchem), 100 U/mL penicillin-streptomycin (15140122; ThermoFisher), and 457 458 50 mg/mL primocin (ant-pm-1; InvivoGen, San Diego, CA). EM was used to expand the epithelial cell 459 numbers as monolayers or organoids. Y-27632 was present only in the first 48 hours of cell culture and was not added to subsequent media changes. The medium was changed every 48 hours. When the 460 cell coverage was greater than 80% (typically 5 to 7 days), the epithelium was dissociated to fragments 461 462 to seed onto the intestinal MPS.

463 **Fabrication of primary human intestinal microphysiological system (MPS)**

The human intestinal MPS was fabricated from polymethylmethacrylate (PMMA) and 464 photocured resin (Formlabs, Inc.). PMMA provided an optically transparent material with a low oxygen 465 diffusion coefficient that could be sterilized and reused.⁹⁹ The photocured resin provided a 3D printable 466 bottom frame to support the PMMA device and simultaneously house the optical reader for oxygen 467 measurements. The cell culture chamber and gas exchange channels were fabricated from 5.8 mm 468 469 and 1.5 mm-thick PMMA sheets (44352, 44292; US Plastics). The microfluidic culture region was composed of 5 rectangular wells, with dimensions of 11.6 mm by 7.0 mm each. The rectangular gas 470 channel on top of the wells was 70 mm by 45 mm by 1.5 mm. The bottom and top pieces of PMMA 471 472 were laser cut from a 1.5 mm-thick PMMA sheet, while the middle piece, for the cell culture wells, was laser cut from a 5.8 mm-thick PMMA sheet. Briefly, to remove dust and burr material with minimal 473 cracking, each PMMA surface was quickly wiped with 100% IPA solution and air-dried. All 3 pieces 474 475 were pressed together between two sheets of brass using a pneumatic heat press (SwingPress 10-0403; Across International). Annealing was performed with the heat press set at 100°C and 300 psi for 476 2 hours, then the bonded device was cooled for 3 hours at room temperature. A rubber gasket was 477 478 laser cut and placed on top of the PMMA layer to seal the device, prior to bolting together. The device was tested for leaks using red and blue dyed water and cracks were sealed by application of 479 480 dichloromethane (DCM) to the seams.

In the completed device, barbed connectors screwed into the photopolymer resin top lid provide gas flow into and out of the device. A rubber gasket seals gas flow into and out of the PMMA gas exchange channel. The gas exchange channel frame is bonded to the PMMA culture chamber and PMMA base support. A photopolymer resin bottom frame supports the entire PMMA device, and screws attach the top lid to the bottom frame to seal the MPS closed. The optics front-end connects via a µUSB-to-HDMI cord to a phosphorescence-lifetime fluorimeter detector and interrogates the middle culture well, where the iPOB is located.

A neutralized collagen hydrogel (2.0 mg/ml) was cast into each well of a prefabricated PMMA device and allowed to polymerize at 37°C for 1 hour. 250 μ L of 1X DPBS was overlaid on the polymerized hydrogel and allowed to pre-swell for at least 5 hours at room temperature. DPBS was removed and the hydrogel was rinsed three times before overlaying with 250 μ L of EM containing hISCs that had been mechanically dissociated by pipetting 10 times until cells were ~1-10 cell-clumps. Once the hISCs formed a confluent monolayer, an iPOB was added to the epithelium to measure oxygen.

494 **Real-time monitoring and control of oxygen concentration in 3D culture**

495 Oxygen concentration in the system was continuously measured with an integrated 496 phosphorescent oxygen sensor (iPOB) using near infrared (NIR) phosphorescence lifetime fluorimetry. 497 The iPOB is composed of porous poly(2-hydroxyethyl methacrylate) (pHEMA) gel functionalized with palladium-benzoporphyrin derivatives (Pd-BPD) that respond to local oxygen concentrations via 498 phosphorescence quenching.²⁷ The photoluminescence excitation and detection wavelengths are 630 499 500 nm and 800 nm, respectively. The iPOB (Profusa, Inc., San Francisco, CA) has been manufactured in a variety of sizes, but for all experiments a disk-shaped, 5 mm diameter and 0.5 mm thickness, iPOB 501 502 was used. The optics front-end of the NIR phosphorescence lifetime fluoroscope (Profusa, Inc., San Francisco, CA) was inserted into the support frame below the MPS. The oxygen concentration was 503 504 controlled via a PMMA gas-mixing microfluidic chip with tubing connecting outlets from the gas-mixing 505 chip to the inlets of each MPS (Fig 9A,B). The gas-mixing chip was previously used to generate 8 concentrations of mixed gas, ranging from less than 3 µM of oxygen to 180 µM. (Fig 9C,D)²⁸ Gas flow 506 to the gas-mixing chip was regulated using an air flow control valve (62005K313; McMaster-Carr) and 507 508 monitored with a mass flow meter with digital output (GFMS-010061; Aalborg GFM) (Fig 9B, Left). Hydrated mixed gas exiting the gas-mixing chip was introduced into individual MPS at a rate of 5 509 mL/min to prevent media evaporation. Within 30 minutes, the gas mixture equilibrated with the local 510 MPS environment to generate the desired oxygen concentration at the hydrogel surface-media 511 interface where oxygen was measured using the iPOB (Fig. 1D). The intestinal epithelium inside MPS 512 513 was cultured in the mixed gas environment and compared to intestinal epithelium statically cultured in 514 a normobaric incubator with an oxygen environment of 186 μ M.

515 *Immunocytochemistry*

To assess the impact of various durations and magnitudes of hypoxia on hISC proliferation, 516 517 primary human intestinal epithelium and single cells isolated from each MPS chamber were stained for Ki-67. For Ki-67 staining of intestinal epithelium following hypoxia, the intestinal MPS was opened, and 518 519 media was removed from each chamber. Intestinal epithelium on top of the collagen hydrogels were 520 fixed with 4% PFA for 15 minutes. After fixation, samples were rinsed once with 1X PBS and overlaid with PBS. Samples were permeabilized for 15 minutes with 0.5% Triton-X 100 in 1X PBS. Samples 521 522 were blocked for 30 minutes with a 3% BSA solution. After blocking, samples were stained for 523 proliferation marker Ki-67 Alexa Fluor 647 (1:250 dilution in 3% BSA, Cat. No. 652407; BioLegend) for

1 hour and nuclei counterstain bisbenzimide (1:1000 dilution in 1X dPBS, Cat. No. 1155; Millipore Sigma) for 5 minutes at room temperature. To look at cell-cell contacts, tight junction protein Occludin (Cat. No. 13409-1-AP; Proteintech) was added for 1 hour, followed by incubator with the secondary antibody Cy3 (Cat. No. C2306; Sigma) for 2 hours. After staining, the gels were overlaid with 1X dPBS and stored at 4°C until imaging. All samples were imaged with a Olympus (Waltham, MA) IX81 microscope, using Metamorph Basic (Molecular Devices, San Jose, CA) software, and image analysis was performed with ImageJ.¹⁰⁰

531 Organoid forming efficiency assay and Fluorescence-activated cell sorting (FACS) of live cells

532 Following MPS culture, human intestinal epithelium was dissociated into single cells, sorted 533 using a flow cytometer, and suspended in Matrigel[®] on a quad CellRaft[™] Array (CRA) platform (Cell Microsystems, Durham, NC).³⁴ Briefly, each sample was retrieved and placed in a separate conical. 534 500 U/mL Collagenase IV (17104019; Gibco) was added to breakdown the scaffold and incubated for 535 10 minutes at 37°C. After centrifugation, the cell pellet was rinsed twice in DPBS and re-suspended in 536 537 150 μL of 0.5 mmol/L EDTA in DPBS with 10 mmol/L Y-27632 and incubated for 5 minutes at 37°C. The fragments were further dispersed by triturating 30 times using a 200 µL pipet tip. After 538 539 centrifugation, the cell pellet was re-suspended in 500 µL TrypLE Express (12605-036; Gibco) with 10 mmol/L Y-27632 and incubated for 5 minutes at 37°C. The cell suspension was gently triturated 7 times 540 541 using a 28.5-gauge insulin needle to further dissociate. 500 µL of EM was added to guench the reaction, 542 and cells were pelleted and rinsed once in EM. After pelleting again, cells were re-suspended in 200 µL of EM containing 1% FBS. Immediately before FACS, APC Annexin V (1:200, 640941; BioLegend) 543 was added to cells for live/dead discrimination. After staining, cells were rinsed in EM and filtered 544 through 0.4 µm FACS tube top filter (352235; Corning). 5,000 Annexin V- live cells were isolated via 545 FACS and re-suspended in 250 µL of Growth Factor Reduced Matrigel[®] (354230; Corning). Cell-gel 546 suspensions were plated in each chamber of the quad CRA. To cover the CRA, each array was 547 centrifuged at 200 x g for 5 minutes. CRAs were polymerized for 20 minutes at 37°C in an incubator 548 and then overlaid with EM containing 10 mmol/L Y-27632. Media was replaced every 2 days. To 549 determine OFE, the CRA was scanned, and the total number of organoids was counted on day 2, day 550 4, and day 6. The OFE (%) was calculated as the total number of organoids created in one CRA 551 552 chamber, divided by 5,000 cells and multiplied by 100. FACS and flow cytometry were performed using a SH800Z Cell Sorter (Sony Biotechnology, San Jose, CA). 553

554 Flow cytometry

Cells collected from normoxic and hypoxic MPS were dissociated following the protocol above 555 up to the step of addition of TrypLE Express. Following trituration with the insulin needle and guenching 556 with the EM, cells were pelleted and re-suspended in 100 µL of 1X DPBS. Approximately 50,000 cells 557 from each chamber were fixed by adding 400 µL of 4% PFA solution while being constantly vortexed 558 to prevent cell aggregation. After fixation, single cells were pelleted and re-suspended in 0.3% Triton-559 X for 15 minutes to permeabilize the cell membrane. Single cells were pelleted and re-suspended in a 560 staining solution of DMEM containing 1% FBS and anti-Ki-67 Alex Fluor 647 (1:250 dilution in 3% BSA, 561 562 Cat. No. 652407; BioLegend) for 1 hour on ice. After staining, cells were rinsed in PBS and filtered through 0.4 µm FACS tube top filter (352235; Corning). Ki-67-positive single cells were quantified by 563 564 flow cytometry.

565 **qRT-PCR**

To assess the expression of genes that are responsive to hypoxia, human intestinal epithelial 566 567 samples from each MPS were collected for gRT-PCR analysis. Briefly, cells attached to collagen hydrogels were lysed in 200 µL of RNA Lysis buffer (AM1931; ThermoFisher). Total RNA was extracted 568 using RNAqueous-Micro Total RNA Isolation Kit (AM1931; ThermoFisher) according to manufacturer's 569 protocols. cDNA was generated from ~2 ng of total RNA from each sample using iScript Reverse 570 Transcription Supermix for gRT-PCR (170-8891; BioRad) according to manufacturer's protocols. cDNA 571 was diluted 1:20 and 1 µL was used for qRT-PCR using the following Tagman probes: 572 573 HIF1a:Hs00153153 m1, BNIP3: Hs00969289 m1, SLC2A1 (GLUT1): Hs00892681 m1, VEGFA: Hs00900055_m1, IL6R: Hs01075664_m1, IL10RB: Hs00175123_m1, IL17RB: Hs00218889 m1, 574 IL22RA1: Hs00222035 m1 (Applied Biosystems) and SsoAdvanced Universal Probes Supermix 575 (1725281; BioRad) according to manufacturer's protocols. gRT-PCR was carried out in a StepOnePlus 576 577 Real Time PCR System (Applied Biosystems). For each sample and experiment, triplicates were made and normalized to 18S mRNA levels. Fold change was expressed relative to normoxic controls using 578 579 $\Delta\Delta$ CT analysis.¹⁰¹ All statistics for gene expression were generated using a Student's t-test. In all 580 statistical analysis, p < 0.05 was considered significant.

581 Bulk RNA-sequencing

To investigate the dynamic response of hISC to hypoxia at the whole transcriptomic level, we 582 performed RNA-seq on human intestinal epithelium exposed to 6 hours, 24 hours, or 48 hours of 583 584 hypoxia inside intestinal MPS, along with normoxic control intestinal MPS which were cultured for each respective time point with no hypoxia exposure. RNA samples were collected from each intestinal MPS, 585 and Total RNA was extracted using RNAqueous-Micro Total RNA Isolation Kit (AM1931; 586 ThermoFisher) according to manufacturer's protocols and stored at -80° C. To assess RNA guality prior 587 588 to submission for sequencing, an RNA integrity number (RIN) was measured using the Agilent 2100 Bioanalyzer. After confirmation that each sample had a RIN of at least 8, integrated fluidic circuits (IFCs) 589 590 for gene expression and genotyping analysis were prepared using the Advanta[™] RNA-Seq NGS Library Prep Kit for the Fluidigm Juno™. 591

592 Informatics

593 Gene level expression was obtained through pseudo alignment of reads to human genome GRCh38 using Kallisto.¹⁰² Principal component and sample correlation analysis were done with 594 Bioconductor packages Biobase, cluster and gvalue.¹⁰³ Expression values for plotting were obtained 595 by TMM normalization across all samples using EdgeR package and differential expression between 596 normoxic and hypoxic samples at each time point was calculated from raw counts with DESeq2.^{104,105} 597 598 GSEAs were performed using all hallmark gene sets on n = 3 samples from each duration of hypoxia (6-, 24-, and 48-hours) to its corresponding normoxic control separately. These comparisons were 599 ranked by NES.106,107 600

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615 **AUTHOR CONTRIBUTIONS**

- K.R.R. designed the study and conducted the experiments, analyzed data, and wrote the manuscript.
- R.J.B. conducted experiments, analyzed RNA sequencing results, and wrote the manuscript
- 618 M.J.C. conducted experiments and analyzed RNA sequencing results.
- J.B. conducted experiments, helped analyze data, and reviewed the manuscript
- 520 J.L. conducted experiments and analyzed data
- 521 J.M.T. conducted experiments
- 622 C.M.H. helped analyze data
- 623 K.A.B. helped analyze data
- S24 S.J. conducted experiments
- 525 V.A.P. worked with K.R.R. to build and integrate all components of the microphysiological systems.
- 626 M.Y. designed, fabricated, and validated the gas mixing chip using COMSOL Multiphysics and 627 experimental methods.
- M.D. and S.T.M. conceived the study and wrote the manuscript.
- A.Z. Contributed experimental design and informatics analysis
- A.T.B. Provided intellectual contributions on the topics of mechanisms of ischemia/reperfusion and hypoxic injury

632 CONFLICT OF INTEREST

533 STM has a financial interest in Altis Biosystems, Inc., Durham, NC. The remaining authors declare no 534 competing related financial interests or conflict of interest at the time of the conduct of this study.

635 **FIGURE CAPTIONS**

Figure 1. Development of a tunable hypoxic microphysiological system (MPS) with integrated oxygen sensors and cultured primary human intestinal epithelium.

A) Exploded view of MPS compartments. B) (left) Top view photograph of a standard microscope glass slide (75 x 25 mm) and dime, for reference, placed above the MPS with iPOB (black arrow) inside well 3 and red and blue dyes in wells 1 and 5 and 2 and 4, respectively. (middle) 1 well brightfield image of monolayer of cells. (right) Inset from 1 well brightfield image shows magnified colonies in hISC monolayer. C) (Top) Side view photograph of MPS with middle well containing the green iPOB. Schematic of iPOB integrated into hydrogel to measure oxygen at the cell layer inside the cell culture

644 well. (Bottom-left) Schematic of porphyrin-based luminescence. In high oxygen, luminescence from the porphyrin is guenched by energy transfer to oxygen, resulting in a decrease in phosphorescence 645 646 lifetime. In the absence of oxygen, porphyrin molecules are excited by LED from the detector and 647 phosphoresce with increased lifetime. **D**) Plots of 8 oxygen concentration versus time (7.5 hours) measurements from the iPOB inside the MPS, with 8 different mixed gas inputs, show generation of 8 648 649 established oxygen environments. E) (Top-left) Brightfield image of human intestinal epithelial stem 650 and progenitor cells grown inside a MPS well to confluence. Fluorescent images show tight junction structures between epithelial cells marked by the protein Occludin (magenta, bottom-right), stem cells 651 652 marked by SOX9 (magenta, top-right, and proliferative cells marked by EdU (red, bottom). Nuclei are marked by Bisbenzimide+ staining in grey or blue. F) Organoid forming efficiency (%) measured over 653 6 days from single cells isolated from MPS after 5 days in culture. G) Oxygen concentration tracked 654 655 using the iPOB inside a MPS containing human intestinal epithelium for 24 hours with normoxic and 656 hypoxic culture environments.

Figure 2. Primary human intestinal epithelium cultured under hypoxia inside microphysiological system shows dynamic response to different durations of hypoxia

659 A) Representative images showing colonies of human intestinal stem cells cultured on collagen scaffolds as 2D monolayers showing dynamic changes in HIF1 α IF staining over different durations of 660 normoxia or acute hypoxia. B) Mean pixel intensity of HIF1 α staining in A over different durations of 661 normoxia or hypoxia. Technical replicates represent mean pixel intensities of all nuclei in 3 individual 662 colonies. **C**) RT-qPCR results of human intestinal epithelium for downstream HIF1 α targets (BNIP3, 663 GLUT1, VEGFA) after 24 hours of normoxic or hypoxic environment exposure. **D**) Flow cytometry 664 results from single cells isolated after 6, 24, 48, and 72 hours from normoxic and hypoxic MPS and 665 stained for Annexin V(-) to quantify cell viability cells (%)E) Immunostaining of hISC monolayer colonies 666 for proliferation marker KI67 following 24 or 48 hours of hypoxia F) Functional stemness as measured 667 668 by Organoid forming efficiency (OFE %) for 24-hour normoxic and hypoxic samples after 6-days of 3D Matrigel[™]-embedded culture. G) Functional stemness for 48-hour normoxic and hypoxic samples after 669 6-day of 3D Matrigel[™]-embedded culture. H) Principal component analysis of RNA-seq results showing 670 671 clusters of replicate samples from each time point (6, 24, 48 hours) and condition (normoxia or hypoxia). Bulk RNA-sequencing samples were sequenced from a range of 39 to 93 million reads I) Heatmap 672 373 showing mean CPM-normalized and log-transformed expression of stem/progenitor markers or 674 differentiated lineage markers from hypoxia bulk RNA sequencing in 2H.

Figure 3. Characterizing the transcriptomic response to different durations of hypoxia by primary human intestinal epithelium

A) Hallmark GSEA results for differentially expressed genes at each time point of hypoxia shown compared to normoxic controls. DEGs from RNAseq from each timepoint from **2H** were compared using GSEA and ranked by Normalized Enrichment Score. Gene sets related to hypoxia or inflammatory signaling are highlighted in pink. **B**) Heatmaps of gene expression from bulk RNAseq data from **2H** for all genes from the Hallmark Inflammatory Response GSEA gene expressed in the bulk data set. Interleukin receptors from the Hallmark Inflammatory Response gene set that are expressed in our dataset are highlighted in pink.

Figure 4. Hypoxia-induced changes in hISC gene expression for interleukin receptors

685 A-C). Fold change comparison of expressed IL receptor genes from each duration of hypoxia in 2H is shown. Differentially expressed genes that are upregulated in hypoxia compared to their normoxic 686 687 counter parts are shown above 1 and genes downregulated in hypoxia are shown below 1. Pink dots 688 signify genes that are significantly upregulated in hypoxia at a given timepoint and blue dots represent genes that are significantly downregulated in hypoxia at a given time point. Grey dots represent IL 689 690 receptor genes that are not significantly differentially expressed at a given time point. Text lists genes 691 represented by the colored dots sorted by descending fold change. D) gRT-PCR measured after 24 hours of normoxia or hypoxia for IL receptor genes that are consistently up- or down-regulated at either 692 693 6, 24, or 48 hours of hypoxia. E) Dot plot showing DEG interleukin receptors from (A-C) and hISC and proliferation marker genes from hISC and TA cells from scRNAseq data of primary human intestinal 694 epithelium. F) Dot plot showing DEG interleukin receptors from (A-C) and hISC and proliferation marker 695 696 genes from hISC and TA cells from scRNAseq data of in vitro primary hISC and progenitors.

Figure 5. Severe hypoxia increases proportion of hISCs residing in G1 phase

698 A) Experimental schematic to assess impact of hypoxia and interleukins on cell cycle progression in human intestinal epithelial cells. B) Example of raw confocal imaging data showing Bisbenzimide+ 699 700 nuclei (top), Cellpose segmented nuclei mask, (second from top), EdU (second from bottom), and KI67 (bottom). C) Example of mitotic bodies found in KI67 channel. D) Histogram showing distribution of 701 702 normalized KI67 mean pixel intensities across Normoxic and Hypoxic conditions without interleukin treatments. Table shows key descriptive statistics that highlight statistically significant differences in 703 distributions. **** p<0.0001 by Kolgorov-Smirnov test of cumulative distributions. E) Percent of KI67+, 704 705 EdU⁺, and Mitotic Bodies counted per colony in normoxia and hypoxia conditions without treatment from interleukins. Dots represent per-colony averages of single-cell mean-pixel intensity values from 706 707 separate normoxic or hypoxic colonies.

Figure 6. Severe hypoxia primes hISCs to respond differently to regulation of cell cycle by interleukins

710 A) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL1 α . **B**) Quantification of cell cycle progression metrics from **5E** for hISCs 711 712 treated with 24 hours of normoxia or hypoxia with or without IL1 β . **C**) Quantification of cell cycle progression metrics from **5E** for hISCs treated with 24 hours of normoxia or hypoxia with or without IL2. 713 714 D) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia 715 or hypoxia with or without IL4. E) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL25. F) Quantification of cell cycle 716 717 progression metrics from 54E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL22. 718

719 Figure 7. Interleukins with no significant effect on hISCs regardless of oxygen condition

A) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL6. B) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL10. C) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL13. D) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL13. D) Quantification of cell cycle progression metrics from 5E for hISCs treated with 24 hours of normoxia or hypoxia with or without IL17^{A/F}.

Figure 8. hISCs demonstrate hypoxia-dependent rescue of organoid forming efficiency when treated with specific interleukins

728 A) Experimental schematic to assess impact of hypoxia, then reoxygenation and interleukins on functional stem cell activity. B) Impact of hypoxia and reoxygenation on functional stem cell activity 729 730 without treatment of interleukins. C) Representative images of organoids from conditions in B,D-M. Images have been pseudocolored to highlight outer membranes of organoids. D) Impact of hypoxia 731 732 and reoxygenation plus IL6 from on functional stem cell activity. E) Impact of hypoxia and reoxygenation plus IL10 from on functional stem cell activity. F) Impact of hypoxia and reoxygenation plus IL13 from 733 734 on functional stem cell activity. G) Impact of hypoxia and reoxygenation plus IL1ß from on functional stem cell activity. H) Impact of hypoxia and reoxygenation plus IL2 from on functional stem cell activity. 735 I) Impact of hypoxia and reoxygenation plus IL4 from on functional stem cell activity. J) Impact of 736 737 hypoxia and reoxygenation plus IL25 from on functional stem cell activity. K) Impact of hypoxia and 738 reoxygenation plus $IL1\alpha$ from on functional stem cell activity. L) Impact of hypoxia and reoxygenation plus IL17^{A/F} from on functional stem cell activity. M) Impact of hypoxia and reoxygenation plus IL22 739 from on functional stem cell activity. 740

Figure 9. Gas mixing chip integrated into microphysiological system to generate 8 different oxygen environments.

743 A) Schematic of complete microphysiological system design with mixed gases in tanks leading into flow meters. Flow meters connect into humidification 15 mL conical tube filled with sterile PBS. Humidified 744 gases travel into each input of the gas mixer chip, and 8 concentrations of gases exit individual outlets 745 connect into separate microphysiological systems (devices). Device oxygen concentration is 746 to monitored using detectors that send information to a laptop located beside the incubator. B) Images 747 taken of each component of the system. C) COMSOL Multiphysics results for the simulated gas 748 749 mixtures from each outlet of the gas mixing chip. D) Image of the gas mixing chip and measured output 750 oxygen concentrations recorded inside individual devices.

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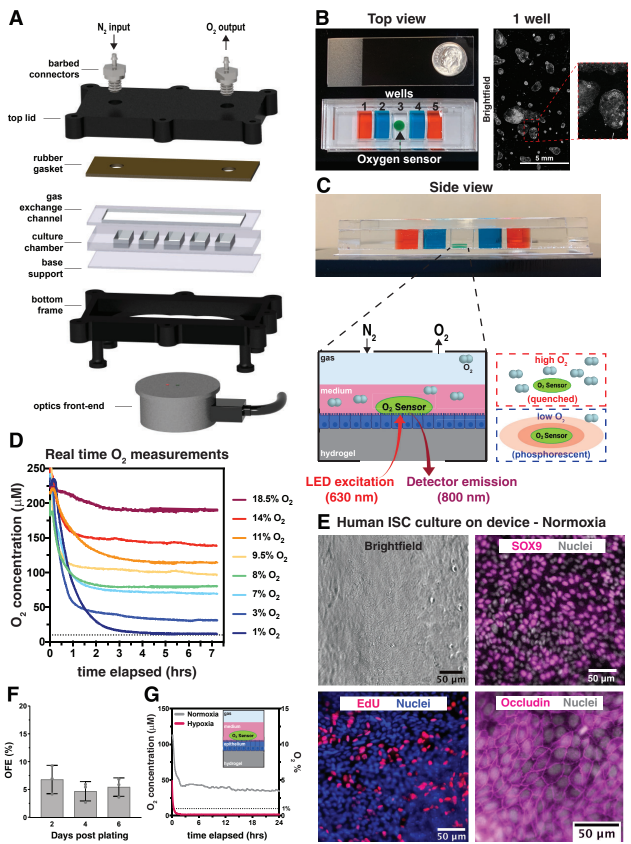
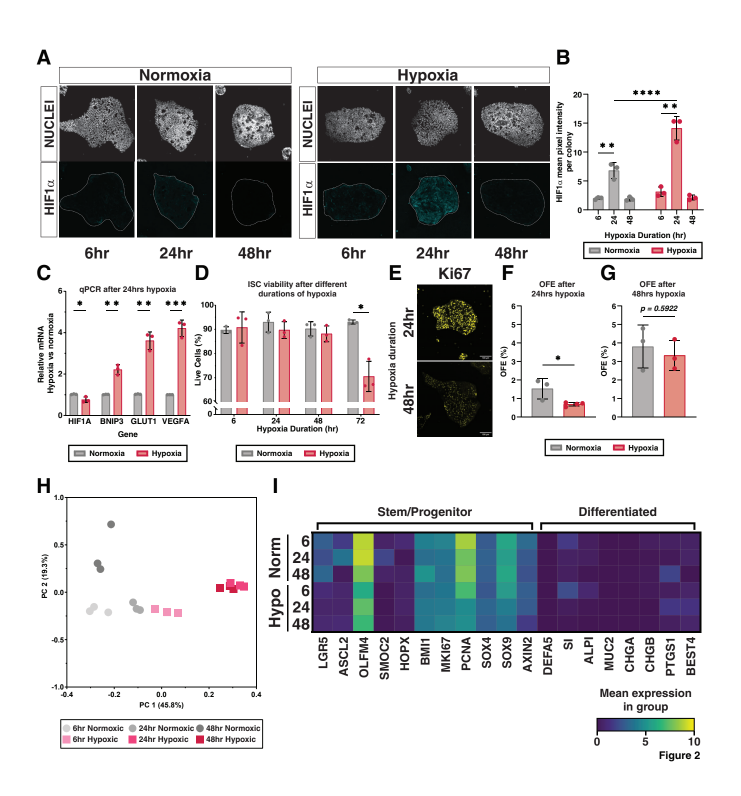
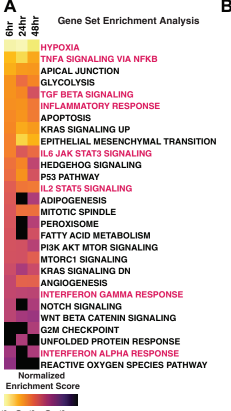


Figure 1







B Hallmark Inflammatory Response

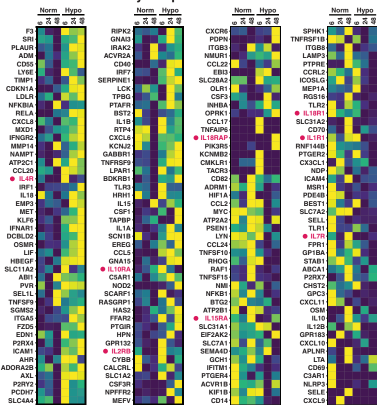


Figure 3

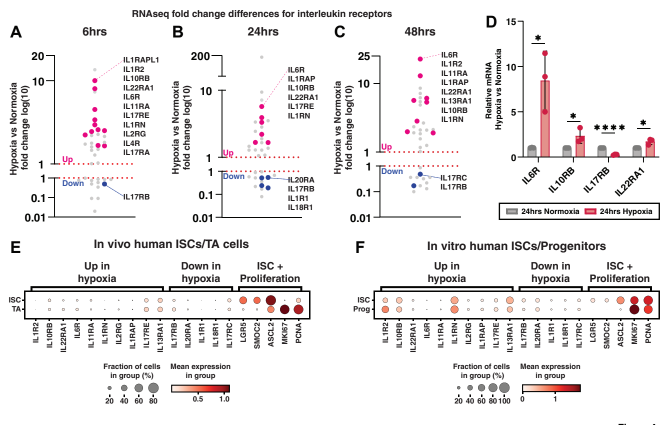
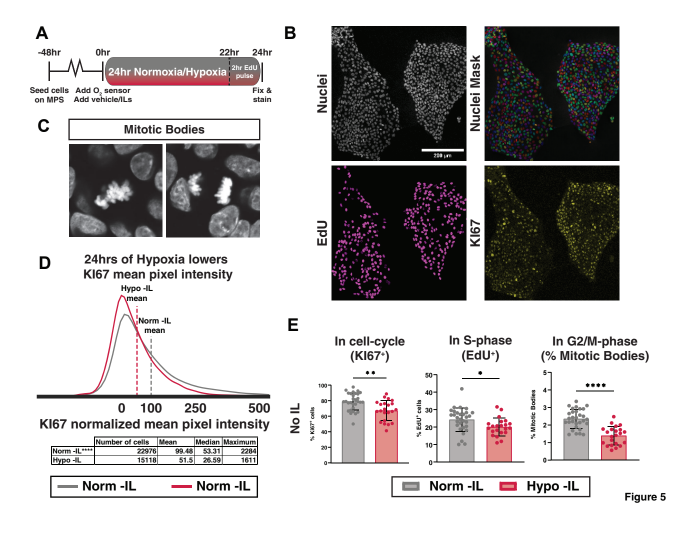


Figure 4



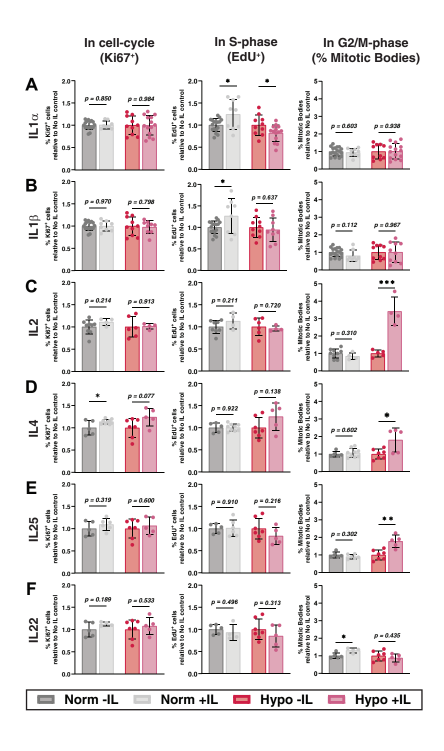


Figure 6

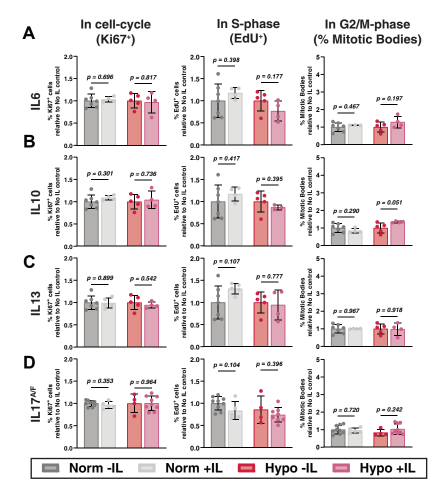


Figure 7

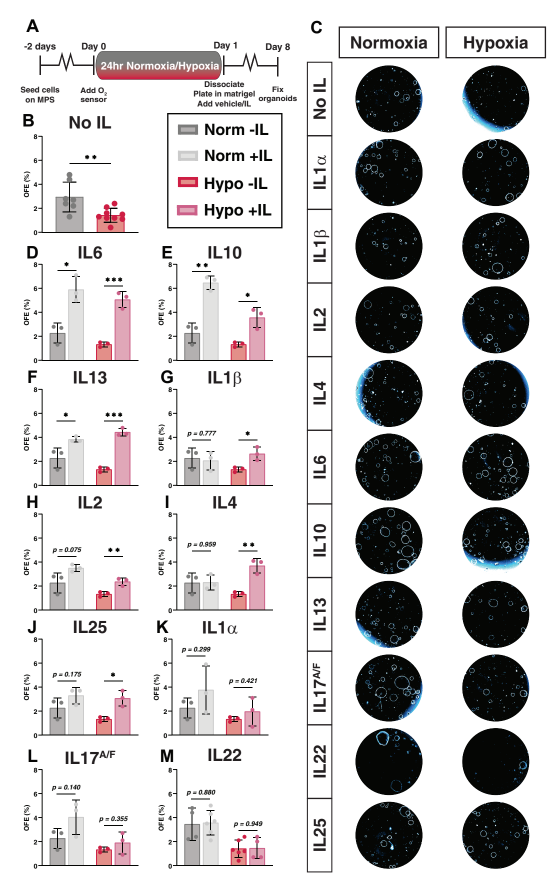
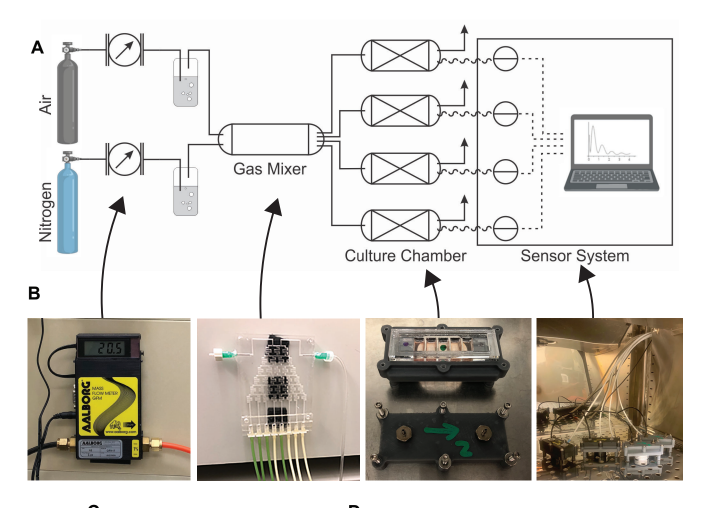


Figure 8



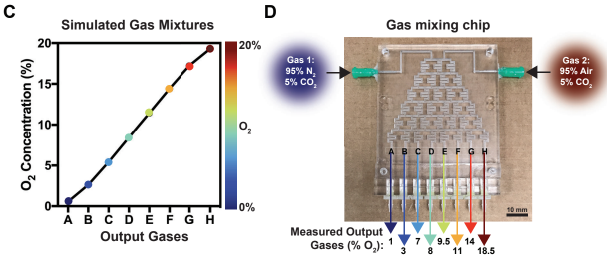


Figure 9