Physiologic Hypoxia and Oxygen Homeostasis in the Healthy Intestine

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Abstract

In recent years, the intestinal mucosa has proven to be an intriguing organ to study tissue oxygenation. The highly vascularized lamina propria juxtaposed to an anaerobic lumen containing trillions of metabolically active microbes results in one of the most austere tissue microenvironments in the body. Studies to date have determined that a healthy mucosa contains a steep oxygen gradient along the length of the intestine and from the lumen to the serosa. Advances in technology have allowed multiple independent measures and indicate that in the healthy mucosa of the small and large intestine, the lumen apposed epithelia experience pO₂ conditions of less than 10 mmHg, so called physiologic hypoxia. This unique physiology results from a combination of factors including counter-current exchange blood flow, fluctuating oxygen demands, epithelial metabolism, and oxygen diffusion into the lumen. Such conditions result in the activation of a number of hypoxia-related signaling processes, including stabilization of the transcription factor hypoxia-inducible factor (HIF). Here, we review the principles of mucosal oxygen delivery, metabolism, and endpoint functional responses that result from this unique oxygenation profile.
Introduction

Studies of the mucosa have provided important insight into metabolic demands associated with normal tissue function. Central to all metabolic processes is the availability, and in some cases the unavailability of molecular oxygen. The gastrointestinal (GI) tract, for example, is characterized by a particularly unique oxygenation profile, experiencing profound fluctuations in blood perfusion on regular intervals throughout the day (26). Even at baseline, epithelial cells lining the mucosa exist in a relatively low oxygen tension environment, herein described as ‘physiologic hypoxia’. Countercurrent oxygen exchange mechanisms in the small intestine have revealed that oxygen from arterial blood supply diffuses to adjacent venules, along the crypt villus axis, resulting in graded levels of low oxygen (124). A steep oxygen gradient has also been documented in more distal, colonic regions of the GI tract, spanning from the anaerobic lumen, across the epithelium to the richly vascularized sub-epithelial mucosa (1). Given the high-energy requirement of the gut and the integral role of the epithelium in maintaining intestinal homeostasis, it is not surprising that these cells have evolved a number of coping mechanisms for this relatively austere metabolic environment. Here, we will discuss how such localized differences in oxygenation contribute fundamentally to the function of the healthy mammalian intestine.

Oxygen Landscape of the Intestine

A steep oxygen gradient exists within the human intestinal tract. Breathable air at sea level contains a partial O$_2$ pressure (pO$_2$) of ~145 mmHg (approximately 21% O$_2$). Measurements of the healthy lung alveolus have revealed a pO$_2$ of 100–110 mmHg (119). By stark contrast, the most luminal aspect of the healthy colon exists at a pO$_2$ below 10 mmHg (1, 68, 69). Such differences reflect a combination of O$_2$ sources, local metabolism and the anatomy of blood flow (Figure 1). The pO$_2$ drops precipitously along the radial axis from the intestinal submucosa to the lumen, which is home to trillions of anaerobic microbes. Over the last fifty years, significant progress has been made toward describing oxygenation at this interface; there are now numerous methods and tools
to measure pO$_2$ in the gut of mammals and in cell culture systems. Results from these experiments provide direct support for the oxygen gradient along the radial axis of the gut and are summarized in Table 1.

The Clark-type electrode was one of the initial tools that enabled measurement of tissue oxygen tension. Placement of the probe onto tissue, such as the mucosal surface, generates a signal proportional to the flux of oxygen molecules to the cathode. In 1965, the oxygen tension in the lumen of the small intestine of the domestic duck was measured using this method. Single-point oxygenation measurements found the pO$_2$ be less than 0.5 mmHg (29). Subsequent adopters of this technology have characterized the oxygen concentration in the small intestine and colon to be heterogeneous with pO$_2$ that ranges from the <1 to >30 mmHg, respectively (125). Despite the probe’s sensitivity and ability to provide real-time oxygen ionization information, it was invasive, sampled a limited area, and had a low signal-to-noise ratio due to oxygen leakage.

A more recent method, termed electron paramagnetic resonance (EPR) oximetry, is an imaging technique that enables oxygen to be quantified across larger areas in the colonic lumen repeatedly and non-invasively. It requires ingestion of activated charcoal that acts as a spin probe, though any paramagnetic material that interacts with oxygen such as nitroxides, lithium phthalocyanine, or India ink can be utilized (47). Oxygen concentrations within the intestinal lumen are proportional to the decay of the spin polarization when subjected to an external magnetic field. Using this approach, luminal pO$_2$ was shown to decrease along the longitudinal gut axis; the duodenum, ascending colon and sigmoid colon were determined to be 32, 11, and 3 mmHg, respectively (56). One concern with this method is its limited spatial resolution. Another concern is the discrepant results compared with the Clark electrode. For example, EPR measurements of the distal colon found pO$_2$ of 3 mmHg, while those done with the Clark electrode were <0.5 mmHg (56, 81). Nevertheless, despite their individual limitations, Clark electrode and EPR have enabled insight into intestinal oxygen concentrations along the longitudinal axis of the mammalian gut.
More recently, a specialized intraluminal probe that uses phosphorescence quenching has enabled very accurate intraluminal pO2 measurement (2). Tissue oxygenation is quantified by exciting the probe with a pulse of light from an optical fiber and measuring the phosphorescence decay as molecular oxygen quenches phosphorescence. This probe has advantages over those used for EPR oximetry because it cannot be endocytosed and therefore remains within the lumen and is minimally impacted by the viscous luminal contents. This surface pO2 of mouse cecal tissue was determined to be ~40 mmHg with this technique and the cecal lumen to be <1 mmHg. Further, microbiome analysis demonstrated that this radial oxygen gradient facilitates oxygen-tolerant organisms near the mucosa. The microbes inhabiting the gut reflect the local pO2 and, hence, it is not surprising that strict anaerobes thrive in this environment given the exceedingly low pO2 measured within the lumen.

The discovery that 2-nitroimidazoles form adducts in hypoxic cells has also enabled the study of oxygen gradients in mammalian intestinal tissues both in vitro and in vivo (138). In their oxidized form, these compounds are taken up by living cells and readily excreted. However, when the pO2 is less than 10 mmHg, these compounds are reduced which enables them to form adducts with thiol groups in proteins leading to retention in hypoxic cells. Further, pimonidazole HCl, a derivative of the 2-nitroimidazole, has aided in validating EPR pO2 measurements (40). Pimonidazole staining could even be used to predict pO2 electrode measurements (100). It is neither dependent on redox enzymes nor changed by the NADH and NADPH levels (5). This technology, coupled with immunostaining, has been used to visually reflect the oxygenation of mouse tumors (6). Furthermore, it has been used to visualize the steep oxygen gradient between the gut lumen and submucosa (Figure 2), a phenomenon sometimes referred to as “physiologic hypoxia” (26). In fact, it is a more stable marker than staining for HIF-1α since it is retained in chronically hypoxic cells (48) and such physiologic hypoxia can be reversed by oxygenation of the colonic lumen (e.g. using oxygenated perfluorodecalin) (57). These nitroimidazole dyes have also been used to image inflammatory lesions and revealed that mucosal lesions are profoundly hypoxic
or even anoxic, similar to that seen in some large tumors, and penetrate deep into the mucosal tissue. It is likely that there are multiple contributing factors (i.e. vasculitis, vasoconstriction, edema, increased O₂ consumption) that predispose the inflamed intestinal epithelia to decreased oxygen delivery and hypoxia (68). These 2-nitroimidazole compounds have shown significant clinical utility, for example in tumor imaging and in the identification of stroke regions within the brain (132). As opposed to other mucosal imaging techniques, these compounds are superior: they only image viable tissue and are inactive within apoptotic or necrotic regions (77). Studies are underway to use these compounds as adjunct radiosensitizers for enhancing chemotherapy targeting (50).

Pimonidazole is now used widely as a hypoxia marker in both research and clinical studies of both normal and disease tissues (64, 120, 144).

Technologies to accurately measure oxygen consumption in cultured cells have also developed rapidly over the past several years. These technologies have enabled studies that monitor local oxygen concentrations in real-time and during modeled conditions that mimic the mammalian microenvironment (e.g. leukocyte interactions with intestinal epithelial cells) (16). The SDR Oxodish system and the Seahorse XF extracellular flux analyzer have both been recently developed to precisely monitor oxygen consumption and are valid alternatives to the Clark type electrodes, which were tedious and required significant expertise to operate (16, 110). The Oxodish uses a fluorescent dye embedded in a sensor spot at the bottom of a plate that is dependent on local pO₂. This technology enables real-time data acquisition and continuous monitoring, but it samples only a small area near the sensor and requires indirect calculation of oxygen consumption rates.

The Seahorse XF extracellular flux analyzer uses optical sensors to measure proton and oxygen in an isolated volume of media. The XF technology combines sophisticated electro-optical instrument with “smart plastic” cartridges that allows for real-time measurements of cellular bioenergetics in a non-invasive and multi-well microplate format. Mitochondrial respiration, indicated by oxygen consumption rates, and lactic acid production from glycolysis, are indicated by changes
in the extracellular acidification rate and can be measured in real-time. This instrument enables characterization of *in vitro* metabolic activity and determination of oxygen consumption rates, glycolysis, ATP production and respiratory capacity (38).

These technologies have allowed basic investigations into intestinal epithelial physiology such as measuring oxygen consumption in response to microbial derived substrates (74). The ever growing number of tools available to quantify intestinal pO\textsubscript{2} has painted a much more complete picture of oxygen homeostasis in the epithelial interface. No doubt, further studies and technologic improvements will lead to a better understanding of intestinal physiology and crosstalk with the microbiota.

**Perfusion of the Intestinal Mucosa**

Oxygenation of the intestinal epithelium depends on the balance of oxygen delivery, consumption and diffusion into the lumen. Oxygen reaches the epithelium by way of three vessels: the celiac, superior and inferior mesenteric arteries, which supply the digestive organs including the small intestine and colon. The three vessels, in the unfed state, receive 20-25% of cardiac output (93). Small arterial branches penetrate the intestinal muscularis and coalesce into a submucosal arterial plexus that forms a vascular layer throughout the length of the intestines. Despite its mass, the muscularis receives only a fraction of the blood flow with up to 80% being directed to the mucosal layer (21, 24). In the small intestine, 60% of mucosal blood flow is distributed to the villi and 40% to the crypts (93). Under fasting conditions only a fraction of mucosal capillaries are utilized (124). However, following a meal, capillaries are recruited as intestinal blood flow can increase up to 200% (93).

The small and large intestine differ in structure and function. The small intestine is optimized for digestion and absorption and has villi that amplify surface area. Villi necessitate a unique microcirculation to maintain perfusion. Each villus contains one or two arterioles 10-15 µm in diameter, which travel toward the villous tip where it forms a hairpin capillary structure. The blood
travels up these vessels and then back in the opposite direction toward a single venule. Within the villi, afferent and efferent vessels are separated by a distance of only 20 µm (86). Interestingly, this arrangement allows for a countercurrent shunt in which oxygen carried into the villus is able to diffuse across to the venule without being transported through the vascular circuit bound to red cells; this reduces the oxygen content of blood delivered to the villous tip (52) (Figure 1). Direct evidence for the existence of a countercurrent oxygen shunt was obtained when investigators placed an oximeter in the venous outflow of an intestinal segment (67). Upstream precapillary access was obtained which enabled injection of oxygen saturated blood and blood with impaired oxygen carrying capacity (methemoglobin), which allowed red cell transit time to be measured. Following injection, the appearance of a peak in post capillary oxygen was detected earlier than the time it took for red blood cells to circulate, indicating the presence of extravascular oxygen shunt in the villous circulation. This countercurrent shunt has also been demonstrated in humans. With the consent of surgical patients, intraoperative arterial injection of a noble gas was observed to enter venous blood more quickly than possible had it remained within the vasculature, implying that diffusible gasses such as oxygen are able to short-cut the vasculature at the villous tip (52). Countercurrent shunt explains why partial occlusion of the superior mesenteric artery causes animals to develop mucosal lesions that could be prevented with application of oxygenated saline into the lumen. Despite auto-regulatory responses that preserve overall blood flow in this model, red cell transit time is prolonged 3-5 fold, which is thought to increase the fraction of oxygen diffused through the countercurrent shunt and result in lower oxygen delivery to the villus tip (51, 86).

Neural regulation of intestinal microvasculature is coordinated by extrinsic and enteric innervation. Sympathetic input controls vasoconstriction, which originates from the celiac and mesenteric ganglia, and acts primarily on mucosal arterioles (137). This arrangement is part of the coordinated acute stress response that diverts blood to the brain and skeletal muscle during physical activity or stressful times (104). The primary stimulus for neural mediated vasodilation is mechanical input, mediated by intrinsic enteric neurons. This has the effect of increasing blood flow
in response to luminal signals. However, during periods of low sympathetic activity, metabolic
regulation of mucosal vasodilation appears to play a greater role in regulating mucosal blood flow
than does neural input (137).

Metabolic regulation of intestinal blood flow occurs even during the fasting state. The small
intestine is presented with 8.5 liters of fluid per day including its own secretions. It absorbs 6 liters
and the remainder is presented to the colon, which absorbs all but ~100 ml that is lost in stool (13).
Much of this fluid transport is mediated by sodium absorption and is driven by the basolateral Na/K-
ATPase (11). Continuous ion and fluid transport has a profound impact on intestinal blood flow. This
was illustrated in a rat model where intestinal blood flow of fasting animals was reduced by over
40% following replacement of intraluminal sodium chloride solution with isotonic mannitol that
impaired ion transport (11). This function was attributed to nitric oxide (NO)-mediated vasodilation,
as pharmacologic inhibition NO synthase produced results of similar magnitude (11). Supporting
this mechanism, it has been shown that increasing the osmolarity of submucosal lymph caused a
dose-dependent increase in arteriole dilation that was dependent on NO (128). NO is a gaseous
signaling molecule with autocrine and paracrine actions. It acts by diffusing into the vascular
smooth muscle and induces vasodilation by activating soluble guanylate cyclase leading to the
formation of cGMP, though recent evidence indicates NO metabolites are also active (136). This
ultimately promotes calcium reuptake, loss of myosin phosphorylation, and relaxation of smooth
muscle. In non-inflamed states, most NO in the intestinal mucosa is synthesized in endothelial cells
by endothelial nitric oxide synthase (eNOS) using oxygen and L-arginine as substrates. Though it
requires oxygen, eNOS functions well in a low pO2 environment with a Km of 6.3 uM/L for oxygen
(pO2 <1 mmHg) (108). The lifespan of NO is inversely correlated to the local pO2, therefore NO
persists much longer in low oxygen environments (134). NO is normally inactivated by oxidized
cytochrome C, but this activity is impaired in low oxygen conditions (102) and may explain why NO
formed in the low pO2 environment of the intestinal mucosa could have an exaggerated lifespan and
action. Cellular pO2 influences eNOS expression in endothelial cells thereby linking local pO2 with
NO mediated perfusion (28, 58, 99). Intraluminal bacteria may also contribute to NO production near the intestinal epithelium (37). In contrast with eukaryotic production of NO, which relies on arginine, bacteria are able to reduce nitrogen oxides that are used as electron acceptors in anaerobic environments. However, there is currently no evidence that NO derived from the microbiota influences intestinal blood flow (126).

Postprandial hyperemia is the marked increase in blood flow to the small intestine stimulated by intraluminal nutrients, particularly lipids and carbohydrates (25). Mesenteric artery flow increases by 28-132% (15, 41, 42, 139, 140) and corresponds with sequential perfusion of the duodenum, jejunum, and then ileum (93). In healthy subjects, splanchnic oxygen uptake increases by 40-64% after consuming a meal of 860-1000 kcal (53, 87). Blood flow associated with postprandial hyperemia is not distributed equally, it is preferentially directed to the mucosa in response to multiple signals. NO plays an important role in hyperemia associated with intraluminal glucose (10) and postprandial hyperemia can be prevented by inhibition of nitric oxide synthase (94). Interestingly, the role of NO in postprandial hyperemia is dependent on adenosine signaling. Adenosine is a purine nucleoside that mediates vasoconstriction or vasodilation depending on concentration, tissue, and receptor subtype (91). Adenosine is a potent vasodilator in the intestine and this response is mediated by A1 or A2B receptors (91, 92). ATP intensive epithelial processes, such as glucose absorption, are believed to stimulate flux of adenosine into the circulation. Infusion of adenosine into the canine superior mesenteric artery resulted in a 2.5 fold increase in intestinal blood flow (49) and adenosine receptor antagonism prevented food induced hyperemia (118) and the appearance of NO metabolites in the portal circulation (92). Use of intravital microscopy to observe the influence of NOS inhibition and adenosine receptor inhibition on postprandial hyperemia demonstrated the linkage between these pathways in responding to acute perfusion demands (91). Additional mediators such as prostaglandins likely also contribute to postprandial hyperemia, but their role is less well defined (23).
In contrast to the small intestine, where absorption of ingested nutrients is a major
determinant of blood flow, postprandial hyperemia has not been observed in the colon (12, 44).
Signals for changes in colonic blood flow are derived from short chain fatty acids (SCFA).
Physiologic concentrations of these compounds can exceed 100 mM in the proximal colon, (14) and
it is estimated that they provide as much as 10% of human energy requirements and up to 70% in
ruminant animals (8). In the colon, SCFA are an important stimulus for perfusion. Mortensen, et al.
provide the best human evidence for this when they utilized resected colonic segments to
determine the response of resistance vessels to varying concentrations of SCFA (98). Treatment
with SCFA individually or in combination, but not glutamine, produced dose-dependent vasodilation
starting at concentrations as low as 3 mM (98). It is unclear what concentration of SCFA the
resistance vessels are exposed to \textit{in vivo}. SCFA, comprised primarily of butyrate, acetate, and
propionate, are present at high concentrations within the lumen. Butyrate is the only SCFA
extensively metabolized at the epithelium while significant amounts of acetate and propionate to
enter the portal circulation. SCFA concentrations in portal blood have been measured at autopsy to
be 0.375 mM (30). However, this has been diluted by blood returning from the proximal GI tract,
small intestine, spleen, and pancreas, which are not sites of SCFA production, so the actual
concentration in the colonic vasculature would most certainly be higher. Support for the homeostatic
role of SCFA stimulated colonic perfusion is provided by a study that instilled SCFA into the human
rectum to promote postsurgical healing (97). This intervention demonstrated a marked increase in
mucosal blood flow following 10-14 days of SCFA instillation into the human rectum, although it is
not clear if this is the result of resistance vessel dilation or increased metabolic activity. The
mechanism for SCFA induced vasodilation appears to be independent of NO (31). Recent studies
reveal a role for the short chain fatty acid receptor GPR41/FFAR3 in systemic blood pressure
regulation (97, 106). While colonic blood flow in response to extracellular SCFA receptor stimulation
has not been studied, it represents a possible mechanism to explain this phenomenon.
Epithelial Oxygen Consumption

The intestinal epithelium has diverse roles including secretion, absorption, and immunity. An incredible amount of energy is invested to harvest energy from food. The thermic effect of food (i.e. the energy expended to digest and absorb food) has been quantified to be approximately 10% of the total daily energy expenditure in humans (107). To support these activities, gastrointestinal oxygen consumption increases disproportionally to gastrointestinal blood flow (22). One study reported that 79% of ATP consumed by the Na⁺/K⁺-ATPase pump was derived from oxidative phosphorylation (34) and inhibition of basal sodium absorption in the ileum of fasting rats reduced oxygen consumption by nearly half (11). Moreover, the rate of oxygen consumption by the human colon, ~8 μM/hr/cm², is even greater than the values reported for the rat (17, 114, 115). In one study, notable for its use of full thickness healthy colonic tissue obtained during surgery, ex vivo treatment with ouabain, an inhibitor of the Na⁺/K⁺-ATPase pump, decreased whole tissue oxygen consumption by 26% (17). The small intestine and colon are dependent on ATP derived from oxidative phosphorylation and epithelial O₂ consumption is an important determinant of oxygen balance at the interface between host and environment.

It is well established that butyrate, an end product of anaerobic bacterial metabolism, is the preferred energy source of the colonic epithelium, even over circulating energy sources (109) to the extent that very little of it escapes into the portal circulation (30). Butyrate metabolism has a direct bearing on epithelial oxygen consumption. Intestinal epithelial cell lines stimulated with butyrate exhibit an increased and sustained oxygen consumption rate that results in depletion of environmental oxygen relative to glucose control (74). Antibiotic depletion of the microbiota was shown to increase the pO₂ of the colonic epithelium as indicated by an oxygen sensitive probe. This treatment mirrored germ free mice, which also have higher pO₂ of the colonic epithelium compared with controls (74). Restoring luminal butyrate in antibiotic treated mice reconstituted the physiologic low pO₂ of the colonic epithelium and hypoxia dependent signaling (74). Interestingly, Donohoe et al. showed that compared with conventionalized mice, the colonocytes of germ-free mice are ATP
deficient and the provision of butyrate can reverse this energy deficit by restoring oxidative respiration (35). Thus, given the voracious epithelial oxygen consumption in the presence of butyrate, it is possible that the prolific production of SCFA in the cecum could, at least in part, explain the proximal to distal pO2 gradient along the longitudinal gut axis (56).

**Luminal Oxygen Diffusion**

The intestinal epithelium is a single cell layer with a surface area that approximates 300 m². This cell layer is positioned between the low pO2 lumen and highly vascular lamina propria. Electron microscopy indicates that the average distance separating the base of the epithelial cells to the fenestrae of the capillaries is narrower in the colon compared with the ileum (1.04 vs 1.94 μm) (79). This arrangement permits oxygen to freely diffuse into the lumen. Establishment of the oxygen gradient depends on prior microbial colonization, which has been noted to occur sequentially with oxygen tolerant organisms being established before strict anaerobes (18, 117). Mucosal samples from human ileostomy sites show facultative anaerobes dominate locations that are normally colonized by strict anaerobes in the native bowel (55). Beyond this, there is now direct evidence that oxygen diffusion from the vasculature influences the mucosal microbiota. Albennberg, et al. showed that exposure of mice to hyperbaric oxygen (100% O2 at 2 atmospheres of pressure) for 4 days reduced the frequency of *Anaerostipes*, an obligate anaerobe and caused complex alterations of 28 other species (2). Importantly, intraluminal oxygen returned toward baseline quickly after the animals returned to room air. This suggests that microbial utilization of oxygen by aerotolerant organisms near the mucosa was active in driving the gradient and in facilitating anaerobic organisms deeper within the lumen. This arrangement of non-anaerobic organisms at the outer radial axis of the intestinal lumen was observed when the microbiome of rectal biopsies and paired stool samples from healthy individuals were analyzed (2). Taken together, these studies indicate that mucosal associated organisms in the lumen actively consume host-derived oxygen and
simultaneously contribute to the steep gradient across the epithelium and enable anaerobic organisms to thrive within the deep luminal space.

Hypoxia Inducible Factor

The hypoxia-inducible factor (HIF) is a global regulator of oxygen homeostasis and facilitates both oxygen delivery and adaptation of oxygen deprivation in numerous cell types, including intestinal epithelial cells (122). HIF is a member of the Per-ARNT-Sim (PAS) domain family of basic helix-loop-helix transcription factors (143). HIFα is degraded in the presence of oxygen and is stabilized when oxygen is limited; HIF-1β is expressed in abundance and forms a heterodimer with the HIFα subunit to regulate gene transcription (62, 123). HIF-2α is an isoform of HIF-1α that is regulated and functions in a similar manner, with some notable differences in target gene specificity (83).

Under adequate oxygen conditions, prolyl hydroxylase enzymes (PHDs) hydroxylate the α subunits of HIF, which enables binding to the von Hippel-Lindau (VHL) protein (65). PHDs use oxygen as a substrate for hydroxylation of HIF and are inhibited under hypoxic conditions. In this reaction, oxygen is inserted into the prolyl residue and into the co-substrate α-ketoglutarate, which splits it to CO₂ and succinate. All three PHD isoforms are expressed on intestinal epithelium and loss of PHD domains have been implicated in detrimental phenotypes. These include loss of exercise performance with PHD1 homozygous knockout, enhanced tumor angiogenesis with PHD2 heterozygous knockout, and decreased neuronal apoptosis, abnormal sympathoadrenal system development and reduced blood pressure with PHD3 homozygous knockout (26).

Binding of hydroxylated HIFα to VHL leads to HIFα ubiquitination, which targets the protein for subsequent degradation by the proteasome. There are physiologic and chemical mechanisms to inhibit HIF activation. Factor inhibiting HIF-1 (FIH-1) blocks HIF transactivation by hydroxylating an asparaginyl residue and blocking association of HIFα with the p300 coactivator protein (88). HIFα is also stabilized by inhibitors of hydroxylases, including dimethyloxalylglycine (DMOG), a competitive
agonist of α-ketoglutarate. Other classes of HIF stabilizers include iron chelators, PHD active site inhibitors, Cullin-2 deneddylators and Fe^{2+} substitutes (36).

Original studies indicated that HIF is stabilized in a graded fashion with decreasing oxygen concentrations (63). Hundreds of genes are positively and negatively regulated in response to hypoxia in a HIF-dependent manner. Binding of HIF to target gene promoters, as determined by EMSA or ChIP, has been observed for a large number of these genes. Many genes respond to hypoxia in a HIF-independent manner. This could reflect indirect regulation by HIF, for example by transcriptional repressors and microRNAs, or regulation by other pathways (78, 96, 146).

Interestingly, only 40% of HIF-1 binding sites are within 2.5 kb of the transcriptional start site (120).

Epithelial Responses to HIF

An expanding body of literature points to HIF as the key mediator of intestinal epithelial adaptation to its low pO_2 microenvironment. HIF coordinates transcriptional responses that directly influence the determinants of oxygen homeostasis including perfusion, metabolism, and barrier maintenance (Figure 3). HIF is a fundamental regulator of whole-body oxygen delivery through regulation of critical genes such as erythropoietin and vascular endothelial growth factor that support production and distribution of red blood cells (61). The intestinal epithelium also contributes to whole-body oxygen distribution through its role in iron absorption, which supports erythropoiesis.

Here too, HIF has emerged as a key regulator. Intestinal epithelial HIF, specifically HIF-2, targets include the gene encoding divalent metal transporter 1, which mediates uptake of Fe^{2+} from the lumen (90), and ferroportin, which mediates basolateral iron efflux from epithelial cells (133). Hepcidin is a circulating protein produced in the liver that prevents iron efflux by binding ferroportin, inducing internalization and degradation. Hepcidin has recently been shown to be negatively regulated by hypoxia in a process mediated by both HIF-1 (105) and platelet derived growth factor BB (127). HIF is also involved in production of adenosine, which as discussed above, plays a fundamental role in regulating perfusion of the intestinal mucosa. This is achieved by the membrane
bound proteins CD39 and CD73 which enzymatically convert ATP/ADP to AMP and AMP to
adenosine, respectively (4). Importantly, CD39 and CD73 expression is regulated by HIF-1α,
providing yet another link between HIF and oxygen delivery (116, 131).

HIF also influences metabolic adaptation of the intestinal epithelium to its low pO2
environment. The small intestine is subject to pronounced pO2 fluctuations associated with periodic
ingestion of nutrients and appears flexible in its use of metabolic substrates utilizing both glucose
and glutamine to varying degrees (75, 141). In contrast, the colonic epithelium consistently favors
oxidative metabolism of butyrate (17, 34). Notably in the colon, the insatiable metabolism of
butyrate depletes local O2 resulting in HIF stabilization and transcriptional activation of HIF target
genes (74). HIF target genes that influence metabolism classically include those involved in
glycolysis (33, 89) as well as pyruvate dehydrogenase kinase (76). This kinase inactivates pyruvate
dehydrogenase thereby preventing glucose-derived pyruvate from entering the TCA cycle as acetyl-
CoA. Butyrate in its role as a HDAC inhibitor also stimulates PDK1 expression (9). This enables
acetyl-CoA derived from β-oxidation of butyrate to enter the TCA cycle. Adding to this, MCT4, a
butyrate transporter is induced by HIF-1α, facilitating further butyrate uptake from the lumen (71,
135). These observations help explain how >70% of oxygen consumed by the human colonic
epithelium is attributed to butyrate oxidation (109).

The contribution of HIF to the intestinal barrier is multifaceted and includes classical and
non-classical components of the epithelial barrier (72). Through cytoskeletal anchoring, tight
junctions form the backbone of the epithelial barrier and help to maintain polarity of epithelial cells
by preventing lipid diffusion between apical and basolateral membranes (59). Claudins are integral
membrane proteins responsible for selective permeability of tight junctions. Recently, claudin-1
(CLDN1) was identified to explain an aberrant junctional morphology of HIF-1β deficient intestinal
epithelial cell lines (112). This work showed that HIF maintains CLDN1 expression through binding
HRE sequences in the gene promoter. The reintroduction of CLDN1 into HIF-1β KD cells restored
barrier function and morphologic abnormalities (112). Adherens junctions are just basal to tight
junctions. These structures are critical components of the apical junction complex and anchor to the perijunctional cytoskeleton, which includes a circumferential ring of actin and myosin. Notably, creatine kinase enzymes (CKM and CKB) as well as the creatine transporter (SLC6A8) were shown to be positively regulated by HIF-2α through interactions with HRE elements in their promoter (46). Moreover, cytosolic CKB co-localizes with adherens junctions and plays an important role in supplying energy at junctional sites for tasks such as tight junction assembly, maintenance and restitution (46).

Given its location between the vasculature and lumen, xenobiotic clearance is an important function of the intestinal epithelium. P-glycoprotein, also called multidrug resistance protein 1, has broad substrate specificity and is a primary effector of xenobiotic transport into the lumen. P-glycoprotein is transcriptionally regulated by HIF-1, thereby providing a distinct example of HIF mediated barrier augmentation (27). In conjunction with mucus secreting goblet cells, the intestinal epithelium extends its barrier apically through formation of the mucus later. Mucus is a complex mixture of glycoproteins that allows delivery of nutrients to the epithelium while preventing exposure to potentially damaging substances and organisms. At least 10 distinct gel-forming and surface mucins are secreted by the intestinal epithelium (82). The mucus layer consists of an adherent layer that is normally devoid of bacteria and a thicker superficial layer that is many times the diameter of the epithelium. Diameter of the intestinal mucus layer has been measured to range from 123-480 µm in the small intestine (15-29 µm firmly adherent) to 642 -830 µm in the colon (101-116 µm firmly adherent) in thickness (7, 130). HIF regulates several components of the mucus layer that are secreted by intestinal epithelial cells. First, MUC3 is a HIF-1α target whose product, mucin-3, co-localizes with intestinal trefoil factor, another barrier-protective molecule characterized by robust trefoil domains (84, 85). Interestingly, intestinal trefoil factor itself is positively regulated by HIF-1α (43). One reason the mucus layer is such an effective microbial barrier is that it functions as a reservoir for secreted antimicrobial peptides (3). Defensins are a prominent class of antimicrobial peptides which are cationic, cysteine-rich, and possess broad antimicrobial activity (45, 103).
Human β defensin-1 (hBD1) is notable within the intestinal epithelium because it is constitutively secreted whereas others are only induced by inflammatory mediators (54, 101). Constitutive expression of hBD1 was shown to depend on basal HIF-1α signaling in multiple intestinal epithelial cell lines and hBD1 expression correlated with other HIF target genes in human tissues (73). Another distinguishing feature of hBD1 is that the full spectrum of its antimicrobial activity is only revealed when its disulfide bonds are reduced (121). Reduction of the hBD1 disulfide bonds is accomplished by thioredoxin, which co-localizes with hBD1 in the colonic mucus; oxidation of hBD1 is prevented by the low pO2 environment of the lumen (60). Considered as a whole, HIF signaling coordinates the transcription of manifold barrier protective genes that maintain the structure and function of the intestinal epithelium in low pO2 environments (Figure 2, 3).

HIF is known to interact with other O₂ responsive signaling pathways that are critical for epithelial homeostasis. For example, hypoxia is known to influence pathways such as the AMP-activated protein kinase (AMPK) pathway, which is activated when PHD activity is limited during hypoxia (145). Further, AMPK was shown to be necessary, though not sufficient for the transcriptional regulation of HIF-1α (80). The X-box binding protein (XBP1) transcription factor provides another example. XBP1 is a regulator of the unfolded protein response (70). Intestinal epithelial XBP1 is critical for maintaining paneth and goblet cells numbers and preventing colitis (70). XBP1 has been shown to protect cells from hypoxia induced apoptosis (111), possibly by facilitating expression of HIF-1α targets through recruitment of RNA polymerase II (19). Another factor with a homeostatic role in the intestinal epithelium is NF-κB. The physiologic role for NF-κB is illustrated by intestinal epithelial specific deletion, which reveals its part in immune homeostasis (147) and expression of antimicrobial peptides and anti-apoptotic genes (129). Here too, the low pO₂ epithelial environment is critical. IKKβ, which mediates NF-κB repression, is itself regulated by oxygen-dependent PHD1 such that basal NF-κB activity is maintained in oxygen limiting environments (32). Finally, the Wnt/β-catenin pathway is linked to oxygenation. The homeostatic role of β-catenin in the intestinal epithelium was exemplified when inducible loss of this signal
caused terminal differentiation of intestinal stem cells resulting in loss of crypt structure and impaired IEC proliferation (39). HIF acts in a yin-yang manner to balance β-catenin signaling by exerting opposing pressures with HIF-1α negatively (66) and HIF-2α positively regulating this pathway (20).

Conclusions

Differences in baseline O₂ tension in mucosal tissues and the profound shifts in energy demand during normal physiologic functions of the intestine provide a unique opportunity to understand tissue metabolism in health and disease. Results from in vitro and in vivo model systems have provided keen insight toward a better understanding of homeostatic physiology. Of particular recent interest is the interplay between tissue oxygenation and the microbiota, many of which culminate on HIF-target pathways that are strongly associated with tissue barrier function and metabolic pathways fundamental to normal intestinal function. Ongoing studies to better define localized metabolomic signatures hold promise in elucidating the interplay of multiple pathways relevant to health and disease.
References:


## Table 1

### Measurements of Intestinal Mucosa Oxygenation

<table>
<thead>
<tr>
<th>Location</th>
<th>Organ</th>
<th>Method</th>
<th>$pO_2$ (mmHg)</th>
<th>Species</th>
<th>Reference</th>
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<tbody>
<tr>
<td>Serosa</td>
<td>Ileum</td>
<td>Electrode</td>
<td>52</td>
<td>Pig</td>
<td>(95)</td>
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<tr>
<td>Serosa</td>
<td>Terminal Ileum</td>
<td>Electrode</td>
<td>34</td>
<td>Human</td>
<td>(125)</td>
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<tr>
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<td>Cecum</td>
<td>Electrode</td>
<td>30</td>
<td>Human</td>
<td>(125)</td>
</tr>
<tr>
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<td>Sigmoid Colon</td>
<td>Electrode</td>
<td>39</td>
<td>Human</td>
<td>(125)</td>
</tr>
<tr>
<td>Mucosa</td>
<td>Colon</td>
<td>Pimonidazole</td>
<td>&lt;10</td>
<td>Mouse</td>
<td>(68)</td>
</tr>
<tr>
<td>Tissue</td>
<td>Cecum</td>
<td>OxyphorMicro Probe</td>
<td>40</td>
<td>Mouse</td>
<td>(2)</td>
</tr>
<tr>
<td>Lumen</td>
<td>Rectum</td>
<td>Electrode</td>
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<td>(81)</td>
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<tr>
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<td>Duodenum</td>
<td>EPR</td>
<td>32</td>
<td>Mouse</td>
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<tr>
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<td>Mouse</td>
<td>(2)</td>
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<tr>
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<td>Mouse</td>
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<tr>
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<td>(56)</td>
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<tr>
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<td>Colon</td>
<td>Electrode</td>
<td>1-30</td>
<td>Human</td>
<td>(142)</td>
</tr>
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</table>
Figure Legends:

Figure 1: Comparison of basal hypoxia in intestinal tissue and other organs and counter-current blood flow in the healthy intestinal mucosa. A: organs from healthy HIF luciferase reporter mice enables visualization of basal tissue hypoxia compared with other organs (113). B: a model of blood flow dynamics in the healthy intestinal mucosa. Counter-current blood flow reduces local pO2 along the crypt-villus axis and results in low pO2 at the villus tip.

Figure 2: Physiologic hypoxia in the colonic epithelium mirrors localization of human β defensin-1 in human colonic biopsies. A: “physiologic hypoxia”. Colonic mucosa of healthy mice show small amounts of pimonidazole, nitroimidazole adduct, along the luminal aspect of the colon (red) suggestive of physiologic hypoxia in the normal colon. B: immunofluorescence staining of hBD-1 (green) in human colonic biopsies reveal localization within epithelium. The staining pattern is similar to pimonidazole with the greatest intensity is along the luminal aspect. DAPI (blue) was used to visualize nuclei in both panels.

Figure 3: Oxygen homeostasis and physiologic regulation of intestinal epithelial function. In addition to the influence of counter-current blood flow (see Figure 1), microbial-derived short-chain fatty acids (e.g. butyrate) stimulate epithelial metabolism and deplete intracellular O2 to the extent that HIF-1 is stabilized. Transcriptional HIF responses in the normal colon include the physiologic regulation of genes important for butyrate transport (MCT1), xenobiotic clearance (P-glycoprotein), adenosine metabolism (CD39, CD73) epithelial barrier function (MUC3, ITF, CLDN1), energy metabolism (CKM/CKB, SLC6A8), antimicrobial defense (hBD1) and iron absorption (DMT1, ferroportin, hepcidin). TJ = tight junction, AJ = adherens junction.
FIGURE 1

A

B

Villus

\[
pO_2 < 10
\]

\[
pO_2 \sim 40
\]

\[
pO_2 \sim 85
\]

Artery
Vein

Spleen
Liver
Heart
Colon
Ileum
Cecum

Color Bar
Min = 7.1915e+05
Max = 6.0343e+06