HIF-1–dependent repression of equilibrative nucleoside transporter (ENT) in hypoxia

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Extracellular adenosine (Ado) has been implicated as central signaling molecule during conditions of limited oxygen availability (hypoxia), regulating physiologic outcomes as diverse as vascular leak, leukocyte activation, and accumulation. Presently, the molecular mechanisms that elevate extracellular Ado during hypoxia are unclear. In the present study, we pursued the hypothesis that diminished uptake of Ado effectively enhances extracellular Ado signaling. Initial studies indicated that the half-life of Ado was increased by as much as fivefold after exposure of endothelia to hypoxia. Examination of expressional levels of the equilibrative nucleoside transporter (ENT) 1 and ENT2 revealed a transcriptionally dependent decrease in mRNA, protein, and function in endothelia and epithelia. Examination of the ENT1 promoter identified a hypoxia inducible factor 1 (HIF-1)–dependent repression of ENT1 during hypoxia. Using in vitro and in vivo models of Ado signaling, we revealed that decreased Ado uptake promotes vascular barrier and dampens neutrophil tissue accumulation during hypoxia. Moreover, epithelial Hif1a/H9251 mutant animals displayed increased epithelial ENT1 expression. Together, these results identify transcriptional repression of ENT as an innate mechanism to elevate extracellular Ado during hypoxia.

Limited oxygen availability (hypoxia) is central to the pathogenesis of many cardiovascular, infectious, and inflammatory diseases. Pathophysiologic changes related to hypoxia include changes in tissue permeability, accumulation of inflammatory cells, and transcriptional induction of proinflammatory cytokines (1, 2). Recent studies have also identified hypoxia-elicited factors that counterregulate such proinflammatory circuit, thereby functioning as endogenous antiinflammatories (3). Central to this latter pathway is extracellular adenosine (Ado), which through activation of surface-expressed receptors on a variety of cells, dampens ongoing inflammation (2, 4–6) and promotes wound healing (7). At present, the exact metabolic steps for generation of extracellular Ado in hypoxia are not well characterized, but likely involve increased enzymatic phosphohydrolysis from precursor adenine nucleotides (ATP, ADP, and AMP). For instance, we recently demonstrated that hypoxia coordinates both transcriptional and metabolic control of the surface ectonucleotidases CD39 and CD73 (8–10), and thereby amplifies extracellular accumulation of Ado. Additional mechanisms also exist to amplify Ado signaling during hypoxia, including coordinate changes at the Ado receptor level. For instance, the vascular endothelial Ado receptor subtype AdoRA 2B is selectively induced by hypoxia and such increases in receptor density are associated with increased vascular barrier responses to Ado (8).

Once generated into the extracellular milieu, Ado is rapidly cleared through passive or active uptake by nucleoside transporters, termed equilibrative nucleoside transporters (ENTs) and concentrative nucleoside transporters, respectively, expressed on a variety of cell types (11, 12). Recent studies suggest that the predominant functional nucleoside transporters in the vascular endothelium are ENT1 and ENT2,
Published December 5, 2005

with a minimal contribution by centrience nucleoside transporters (13), wherein ENT1 and 2 are bidirectional transporters functioning as diffusion-limited channels for transmembrane nucleoside flux. Previous studies have suggested that vascular Ado transport during hypoxia is predominantly inward (14), thereby terminating extracellular Ado signaling. However, more recent studies show that the expression of ENT1 may be transcriptionally regulated by hypoxia (15, 16), thereby functioning to fine tune extracellular levels of Ado.

Therefore, we examined the influence of hypoxia on endothelial and epithelial Ado transport. Results from these studies revealed that ENT1 and ENT2 gene expression and function are attenuated by hypoxia, and that this regulatory circuit maps to hypoxia inducible factor 1 (HIF-1)–mediated repression of ENT expression. These studies provide new molecular insight into endogenous mechanisms of tissue protection during hypoxia.

RESULTS
Increased extracellular Ado half-life in posthypoxic endothelia

Recent studies suggest that hypoxia promotes a cellular phenotype that supports high capacity for rapid nucleotide phosphohydrolysis and enhanced Ado signaling (8, 9). As an extension of these studies, we addressed whether mechanisms of extracellular Ado clearance were different in posthypoxic endothelial cells. Initial studies indicated that hypoxia significantly decreased endothelial uptake of exogenous Ado (Fig. 1 a), thereby increasing Ado half-life. Indeed, these studies indicated that human microvascular endothelial cells (HMEC)-1 subjected to hypoxia (pO2 20 torr, 48 h) and examined for their capacity to transport extracellular Ado (AdoEC, 50 μM final concentration), had significantly attenuated ability to transport Ado (AdoEC 19.2 ± 0.6 μM at 2 h) compared with normoxic controls (AdoEC 1.3 ± 1.1 μM at 2 h, P < 0.01). Control experiments performed at 4°C or in the presence of the ENT-inhibitor dipyridamole (10 μM) revealed significant attenuation of Ado uptake (AdoEC 40 ± 1.6 and 33 ± 2.1 μM concentrations, respectively; P < 0.001), indicating that endothelial Ado uptake requires metabolism and depends on dipyridamole-sensitive carriers. Subsequent studies addressing the hypoxia time course (Fig. 1 b) and dose response (Fig. 1 c) revealed a time-dependent (P < 0.01) and dose-dependent (P < 0.05) decrease in transport of AdoEC.

To confirm that the observed decrease in AdoEC reflects transport and to rule out other potential mechanisms of Ado loss (e.g., metabolism by the ecto-Ado deaminase (17), we determined the kinetics of 14C-AdoEC transmembrane transport (Fig. 1 d). These studies confirmed our studies with native Ado and revealed that 14C-AdoEC transport was 63 ± 6% decreased in posthypoxic endothelia relative to normoxia (P < 0.001). In parallel, 14C-AdoEC transport at 4°C or in the presence of dipyridamole revealed significant attenuation of endothelial uptake (P < 0.001). Together, these data suggest an additional mechanism to elevate AdoEC involves diminished active uptake through endothelial membrane transporters.

Hypoxia rapidly represses expression of endothelial and epithelial ENT1 and ENT2 mRNA

Prompted by these results, recent findings that ENT1/2 predominate in the vascular endothelium (13) and that hypoxia down-regulates dipyridamole-sensitive Ado transporters (15, 16), we examined the influence of hypoxia on expression of ENT1 and ENT2 mRNA. Initial insight gained by microarray analysis revealed an 85 ± 5 and 78 ± 4% inhibition of ENT1 and ENT2 mRNA expression, respectively (18 h hypoxia, pO2 20 torr). As shown in Fig. 2 a, real-time PCR analysis confirmed our microarray results and revealed a time-dependent loss of both ENT1 (95 ± 8% loss at 24 h, P < 0.01) and ENT2 (81 ± 7% loss at 24 h of hypoxia, P <
In addition, we found that expression levels of ENT1 mRNA in HMEC-1 were approximately fourfold higher compared with ENT2 mRNA.

Similar results of ENT1 and ENT2 repression were found when human saphenous vein tissues were subjected to hypoxia, indicating that such findings are not limited to cultured endothelium (Fig. 2 b, P < 0.001). Additionally, ENT1 and ENT2 expression were decreased in T84 cells and intestinal epithelial cell line (Fig. 2 c, P < 0.001), further suggesting that these findings are not limited to endothelial cells.
We extended these mRNA findings to examine ENT1 protein expression in cultured HMEC-1. As shown in Fig. 2 d, Western blot analysis of lysates derived from HMEC-1 subjected to hypoxia (pO2 20 torr for 6–36 h) revealed a dramatic loss of ENT1 with increasing time in hypoxia. Protein repression was maximal at 36 h time periods of hypoxia (no additional loss was evident at 48 h).

**ENT1 repression mediates attenuated endothelial Ado uptake during hypoxia**

We next determined whether inhibition of ENT gene expression was required for the observed functional differences in Ado uptake between posthypoxic and postnormoxic endothelia. Here, we used siRNA-mediated repression in HMEC-1 of ENT1 or ENT2 for gaining insight into relative contributions of these individual gene products on AdoEC transport. As a first step, we generated siRNA double-stranded oligoribonucleotides specific for ENT1 and ENT2 and profiled ENT expression by PCR. (Fig. 3, a and b). As control siRNA, we used a single-stranded forward oligoribonucleotide, which did not influence ENT1 or ENT2 mRNA expression (Fig. 3 c), suggesting that the aforementioned findings of decreased ENT1 or ENT2 expression (Fig. 3, a and b) were specific for targeted gene knock-down by the chosen siRNA molecules. As additional control, we used double-stranded siRNA directed against the 5′-ectonucleotidase (CD73), an endothelial surface enzyme that facilitates conversion of Ado monophosphate to Ado. As shown in Fig. 3 (d and e), siRNA repression of CD73 did not influence expression levels of ENT1 or ENT2. After showing that siRNA-mediated repression of ENT1 and ENT2 was specific and highly efficient using the aforementioned conditions (81 ± 8% inhibition of ENT1 mRNA expression, P < 0.01, and 79 ± 10% inhibition of ENT2, P < 0.01), we measured endothelial Ado uptake using 14C-Ado. As shown in Fig. 3 f, whereas 14C-Ado uptake was not influenced by siRNA repression with control siRNA (5 ± 1% decrease compared with mock-loaded cells, p-values were not significant) or siRNA directed against ENT2 (14 ± 7% decrease compared with mock-loaded cells, p-values were not significant), Ado uptake was decreased after siRNA-mediated repression of ENT1 (76 ± 7% decrease compared with mock-loaded cells, P < 0.01) or simultaneous ENT1 and ENT2 repression (81 ± 5% decrease compared with mock-loaded cells, P < 0.01). Together, these results implicate ENT1 as the prevailing mechanism for the observed changes in endothelial Ado uptake.

**Decreased endothelial Ado uptake promotes endothelial barrier function in vitro**

We next determined whether hypoxia-mediated repression of ENT1 might enhance signaling mediated by AdoEC. Here, we compared Ado responses of normoxic and posthypoxic endothelia using a previously described in vitro model of endothelial barrier (8, 18). As reported previously (8), we observed that barrier function (i.e., flux 70-kD FITC-dextran) in response to Ado was enhanced to a greater extent in posthypoxic HMEC-1 than in normoxic HMEC-1 (Ado EC50’s of ∼30 μM and ∼1 μM for normoxia and posthypoxia, respectively; Fig. 4 a). To define the contribution of hypoxia-mediated repression of ENT1/2, we repeated this experiment in the presence of the ENT inhibitor dipyridamole (10 μM). As shown in Fig. 4 b, in the presence of dipyridamole, the dose response to Ado became indistinguishable between hypoxia and normoxia (Ado EC50 of ∼0.1 μM for both), implicating hypoxia-mediated repression of ENT1/2 in functional Ado responses.

We next defined the contribution of ENT1 or ENT2 to such changes in Ado-elicted barrier responses of the endothelium. To do this, we used the aforementioned siRNA repression approach. As shown in Fig. 4 c, siRNA repression...
of ENT1 was associated with an increase of Ado elicited barrier responses (P < 0.01). In contrast, loss of ENT2 was not associated with a measurable change in endothelial Ado response (Fig. 4 d). These data reveal that inhibition of Ado uptake, either via hypoxia inhibition of ENT gene expression, pharmacological intervention, or siRNA repression of ENT1, represents an adaptive mechanism to elevate extracellular Ado levels, and thereby prolong signaling mediated by this nucleoside.

**HIF-1α–dependent repression of ENT1 promoter**

In an attempt to gain specific insight into the molecular mechanisms of ENT1 repression by hypoxia, we profiled promoter activity in HMEC-1 transfected with luciferase promoter constructs expressing varying length 5’–truncations (225, 282, 344, and 756 bp relative to the major transcription start site). Highest basal expression (i.e., normoxia) was observed in the 756-bp construct (15 ± 2.1-fold over background) and graded decreases in baseline activity were observed with increasing 5’-truncation (6.8 ± 1.7, 3 ± 1.1, and 5 ± 0.6-fold over background for 344-, 282-, and 225-bp truncations, respectively). Comparison of promoter activity in HMEC-1 exposed to hypoxia or normoxia revealed that, whereas other constructs were expressed at or near the level of the normoxia control, expression of the 756-bp fragment was repressed by 59 ± 6% compared with normoxia controls (Fig. 5 a, P < 0.01). Such analysis indicates that the region spanning −344 to −756 of the ENT1 promoter confers hypoxia-repressive activity.

In the course of our experiments, we identified two potential HIF binding sites oriented on the antisense strand of the ENT1 gene promoter (DNA consensus motif 3’–CCGTG–5’ and 3’–CCGTG–5’ located at positions from −653 to −657 and −717 to −721, respectively, relative to the major transcription start site). Because HIF can mediate transcriptional induction or repression (19, 20), we determined if either of these putative HIF–1α–binding sites conferred hypoxia repression of ENT1 in HMEC-1. First, we examined the kinetics of HIF-1α expression in our HMEC-1 cultures. As shown in Fig. 5 b, Western blot analysis of HIF-1α revealed minimal protein in hypoxia, and time-dependent induction over a 24-h period of hypoxia. Such kinetics are consistent with the loss of ENT1 protein during hypoxia (Fig. 2 d). Second, HIF–1α–binding site mutations (Δ657 and Δ721) were introduced in the 756-bp construct, and as shown in Fig. 5 c, mutation of the −717 to −721 site, but not the −653 to −657 site, reversed hypoxia repression of this promoter construct, suggesting that this region of the promoter, at least in part, confers hypoxia repressor activity. Third, we determined whether this region of the ENT1 promoter binds HIF-1α. For these purposes, we used chromatin immunoprecipitation (ChIP) to analyze HIF-1α binding in live cells. As shown in Fig. 5 d, ChIP analysis of nuclei derived from HMEC-1 cells revealed a prominent band of 192 bp in hypoxic but not normoxic samples. No bands were evident in control IgG immunopre-
cipitates, and input samples (preimmunoprecipitation) revealed the predictable 192-bp band under conditions of both hypoxia and normoxia. Such results indicate that hypoxia induces HIF-1\(\alpha\) binding to the distal 192-bp region of the ENT1 promoter. Together, these results provide strong evidence for a functional hypoxia repressor activity, mediated by HIF-1, in the distal 5\' region the ENT1 promoter.

Role of HIF-1\(\alpha\) in epithelial ENT1 regulation

We next pursued the finding that epithelial ENT expression, like that in endothelia, is repressed by hypoxia (Fig. 2 c). To probe the role of HIF-1\(\alpha\) in an epithelial model, we generated a T84 cell line expressing oxygen-stable HIF-1\(\alpha\) (T84\(-\alpha\)ODD) via lentiviral transduction (Fig. 6). As shown in Fig. 6 a, using this lentiviral vector-expressing GFP, >90% of T84 cells were stably transduced by this method. Analysis of the T84\(-\alpha\)ODD by Western blot (Fig. 6 b) and by hypoxia-responsive element (HRE) luciferase (Fig. 6 c) revealed increased HIF function and activity in both normoxia and hypoxia, relative to the control T84-GFP cell line. The further increase of HIF function and activity with hypoxia most likely reflects the presence of oxygen-stable and oxygen-unstable HIF-1\(\alpha\).

Using the T84\(-\alpha\)ODD cell line, we addressed whether ENT1 expression was differentially regulated. As shown in Fig. 7 a, ENT1 mRNA expression in normoxia was decreased by >80% in the T84\(-\alpha\)ODD relative to the control T84-GFP cell line (\(P < 0.01\)). Likewise, enhanced repression of ENT1 was observed in T84\(-\alpha\)ODD compared with control T84-GFP cells at 6, 12, and 24 h of hypoxia (\(P < 0.025\) for each). Additionally, comparison of promoter activity in T84\(-\alpha\)ODD and T84-GFP cell lines exposed to hypoxia or normoxia revealed enhanced repression of the 756-
In vivo analysis of ENT function on the vasculature

As proof of principle for these in vitro and ex vivo findings of hypoxia-induced ENT repression, we extended these findings to an in vivo model. We and others have shown that hypoxia promotes vascular leak and tissue PMN accumulation in vivo (1, 8, 10, 22–24). Thus, we pursued pharmacological inhibition of Ado uptake in a previously described murine hypoxia model (8, 10, 22).

As first step, we administered the ENT inhibitor dipyridamole (10 mg/kg i.p. and 10 mg/kg s.c.) or PBS and subjected animals to either normoxia or hypoxia (8% O₂ and 92% N₂) for 4 h. Plasma levels of dipyridamole (determined by HPLC) ranged between 25- and 110-nm concentrations during the course of the experiment (Fig. 8 a), consistent with previous dosing studies with dipyridamole and dipyridamole analogues (25). In previous in vivo hypoxia experiments, we observed the lungs as particularly prone to developing hypoxia-induced vascular leak and pulmonary edema (10). Therefore, we assessed pulmonary edema by determining lung water content (wet/dry ratio). As shown in Fig. 8 b, hypoxia increased lung water content from 3.81+/−0.42 mg/mg dry tissue to 5.83+/−0.54 (P<0.05) in PBS treated mice. In contrast, lung water was decreased in dipyridamole-treated animals by 16+/−2% under normoxic conditions, and by 21+/−3% (P<0.025) under hypoxic conditions (P<0.05). These results suggest that pharmacological inhibition of Ado uptake is associated with a decrease in lung water.

To assess vascular barrier function of other organs, we administered the albumin marker Evan’s blue before hypoxia. Comparative analysis in normoxia revealed decreased Evan’s blue concentrations in all examined organs (colon, lung, liver, and kidney) of dipyridamole-treated mice compared with PBS controls. Comparison of vascular permeability in dipyridamole or PBS-treated mice subjected to normobaric hypoxia (8% O₂ and 92% N₂ for 4 h) revealed significantly increased vascular leak in all animals, however to smaller degree in dipyridamole-treated animals by 16+/−2% under normoxic conditions, and by 21+/−3% (P<0.025) under hypoxic conditions (P<0.05). These findings support our hypothesis that decreased Ado uptake by the endothelium is a protective cellular strategy to dampen changes in vascular barrier function, both at baseline and during hypoxia-induced barrier dysfunction.

We next examined whether a reduction of Ado uptake at the vascular surface may also function as antiinflammatory response during hypoxia, particularly given the antiinflammatory properties of Ado (4, 6). To test this hypothesis, we screened multiple mucosal organs (colon, lung, liver, and kidney) for PMN accumulation after administration of dipyridamole (Fig. 8 d). This analysis revealed decreased myeloperoxidase (MPO) activity in all organs of dipyridamole-treated animals. Consistent with previous studies (26), MPO activity...
levels were significantly increased after hypoxia exposure in all animals, however to a much lesser extend in all organs of dipyridamole-treated animals compared with PBS controls (P < 0.05 for all organs). These in vivo findings support the hypothesis that Ado uptake via dipyridamole-sensitive carriers is a critical control point in the regulation of PMN accumulation within normoxic and hypoxic tissues and suggest ENT down-regulation as innate antiinflammatory adaptation.

It is known that the two nucleoside transporters ENT1 and ENT2 differ in their sensitivity to inhibition by the inosine analogue NBTI (27, 28). NBTI is a highly potent inhibitor of ENT1 (IC$_{50}$ = 0.4–8 nM), whereas it is only a moderate inhibitor of ENT2 (IC$_{50}$ = 3 μM). Based on pharmacokinetic studies of NBTI in mice, we used an NBTI dose that would result in estimated plasma levels between 1 and 10 nM (29). To pursue this, we administered NBTI (1 mg/kg i.p.) or vehicle (DMSO) and subjected animals to either normoxia or normobaric hypoxia. As shown in Fig. 9 a, hypoxia significantly increased lung water content in vehicle-treated animals (P < 0.025). In contrast, lung water was decreased in NBTI-treated animals under normoxic or hypoxic conditions. Measurement of vascular barrier function revealed significantly increased vascular leak with hypoxia in all animals, however to a smaller degree in NBTI treated animals (P < 0.05 for all organs; Fig. 9 b). Finally, we also screened multiple mucosal organs for PMN accumulation (Fig. 9 c). Consis-

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**Figure 9.** Influence of S-(4-Nitrobenzyl)-6-thioinosine (NBTI) on pulmonary edema, vascular permeability, and PMN accumulation in vivo. BL/6/129 mice were administered NBTI (1 mg/kg i.p. in DMSO) or vehicle alone, and exposed to normoxia or hypoxia for 4 h. (a) Assessment of lung water content in normoxia (black bars) and hypoxia (gray bars) after dipyridamole or vehicle treatment. Data are expressed as mean ± SD mg H$_2$O/mg dry tissue, and are pooled from four animals per condition. *, difference between hypoxia and normoxia (P < 0.025); †, difference between dipyridamole treatment and vehicle control (P < 0.025). (b) To assess vascular barrier function, animals were administered intravenous Evan’s blue solution (0.2 ml of 0.5% in PBS) before normoxia/hypoxia exposure. Animals were killed and indicated organs were harvested. Evan’s blue concentrations were quantified as described in Materials and methods. Data are expressed as mean ± SD Evan’s blue OD/50 mg wet tissue and are pooled from four to six animals per condition. *, differences between NBTI/vehicle treatment groups (P < 0.025); †, differences between normoxia/hypoxia exposure (P < 0.01). (c) Organ assessment of PMN accumulation by MPO measurements in the indicated organs after 4 h of normoxia/hypoxia exposure (*P < 0.025 compared with vehicle; †P < 0.025 compared with normoxia).

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**Figure 10.** ENT1 expression patterns in conditional Hif1α mutant tissue. Real-time PCR analysis of murine epithelial ENT1 transcript in conditional Hif1α mutant and littermate control animals subjected to normoxia or hypoxia. Data were calculated relative to β-actin and are expressed as relative ENT1 mRNA ± SD, where transcript levels in control animals were normalized to 1. Results are derived from three animals in each condition.
tent with the aforementioned results with dipyridamole, we found decreased MPO activity in all organs of NBTI-treated animals after hypoxia exposure (P < 0.05 for all organs). These in vivo results suggest that inhibition of ENT1 contributes to the in vivo resuscitation of vascular leak syndrome by nucleoside transport inhibitors.

Role of HIF in epithelial ENT1 expression in vivo

We extended these findings of HIF-1-mediated repression of ENT1 expression into a genetic in vivo model. Here, we examined baseline expression and the influence of hypoxia on mENT1 mRNA levels in intestinal epithelia derived from conditional Hif1α knockout mice, in which intestinal epithelia from these mice lack detectable Hif1α expression in >70% of cells (3). As shown in Fig. 10, ENT1 levels in mice expressing wild-type Hif1α showed a normal pattern of hypoxia-associated ENT repression (68 ± 3% decrease after hypoxia for 6, P < 0.025). Consistent with our hypothesis that HIF-1 transcriptionally represses ENT1, real-time PCR analysis revealed a 37 ± 6.8-fold increase in intestinal epithelial ENT1 expression in Hif1α mutant animals (Fig. 10), relative to their littermate controls (P < 0.01). Exposure of HIF1α mutants to hypoxia decreased ENT1 expression (P < 0.05), but to a far lesser extent than in wild-type animals. These latter findings likely reflect the remaining ~30% wild-type Hif1α expression in these mice (3). Together, such findings support our in vitro findings and indicate the likelihood that HIF-1 directly regulates murine ENT1 expression.

DISCUSSION

Ado exerts paracrine and autocrine functions on most cell types. Pathophysiologic conditions of hypoxia/ischemia result in numerous adenine nucleotide metabolic changes, and targets for extracellular Ado signaling is now an area of much interest. In the present studies, we explored the mechanisms of extracellular Ado accumulation during hypoxia. For these studies, we used vascular endothelial and mucosal epithelial cells, which both lie anatomically positioned to function as determinants of the inflammatory response. Our results revealed that hypoxia increases extracellular Ado half-life. This response is mechanistically determined, at least in part, by HIF-1α-regulated ENT1 repression.

The results from the present study are consistent with previous findings that hypoxia inhibits intracellular metabolism of Ado to AMP. Decking et al. showed, by measuring Ado and AMP concentrations in the coronary sinus and coronary arteries of isolated guinea pig hearts, that hypoxia is associated with a functional inhibition of the Ado kinase (30). The authors propose that hypoxia-associated inhibition of the Ado kinase leads to increased intracellular Ado levels. It is known that during hypoxia, Ado flux is predominantly directed from the extracellular to the intracellular space (14). Therefore, two cellular strategies with regard to Ado transport may play a role to elevate extracellular Ado levels during hypoxia. First, as described by Decking et al., increased intracellular Ado levels due to inhibition of Ado kinase-dependent metabolism of Ado to AMP would decrease the transcellular Ado gradient, thereby decreasing flux through bidirectional ENTs and, thus, elevating extracellular Ado levels during hypoxia (30). Second, transcriptional repression of ENTs, as we demonstrate in the present work, decreases overall equilibrative Ado transport capacities, thereby decreasing intracellularly directed Ado transport. It should be noted that we have not observed a direct change in ENT function by hypoxia per se. Rather, our studies addressed function as it related to expression levels of ENT. From this standpoint, it is likely that inhibition of adenine nucleotide metabolism (e.g., inhibition of Ado kinase) could serve as an immediate response to hypoxia (within minutes), whereas our proposed transcriptional mechanism serves as a more adaptive phenotype (within hours). Therefore it is reasonable that both mechanisms could function during hypoxia and, from this perspective, contribute synergistically to elevate extracellular Ado during hypoxia.

Consistent with recent work in cardiac myocytes (16), the present studies in vascular endothelia and mucosal epithelia identified transcriptional repression of ENT as a mechanism for such decreased Ado uptake during hypoxia. Given the temporal and robust hypoxia response observed in the repression of ENT1, a candidate regulator was HIF-1α, a heterodimeric transcription factor whose activation is dependent on stabilization of an O2-dependent degradation domain (31). It is recently appreciated that HIF can function as both a transcriptional activator as well as a repressor (19). Therefore, a search of the cloned ENT1 gene promoter identified two potential HIF-1α-binding DNA consensus motifs located in the distal 5′ region of the ENT1 promoter. However, the existence of a HIF-1α–binding consensus is not evidence for HIF-1α–mediated response; instead, the HRE is defined as a cis-acting transcriptional regulatory sequence located within 5′-flanking, 3′-flanking, or intervening sequences of target genes (32, 33). Therefore, two strategies were used to define the role of HIF-1 in repression of ENT1. First, mutational analysis of ENT1 promoter constructs defined a functional HRE located in the distal 5′ region of the ENT1 promoter (−717 to −721 relative to the major transcription start site). Second, ChIP analysis confirmed HIF-1α binding to the ENT1 promoter. At present, we do not know if this binding is direct or indirect, and we do not know the nature of HIF-mediated transcription. While previous work with peroxisome proliferator-activated receptor α (PPAR-α) gene implicated potential repressor activity with HIF-1α binding sites oriented on the antisense strand (20), implying some degree of transcriptional directionality, there is not direct evidence for such a mechanism. Moreover, while a recent study comparing transcriptional responses between hypoxia and constitutively active HIF1α identified a large cohort of transcriptionally repressed genes (19), no unique patterns of HRE expression were noted. Thus, more work will be necessary to define the nature of HIF-mediated repression.
The present study also identified ENT1 as a potential physiologic target in vivo. Indeed, using the ENT inhibitor dipyridamole, we demonstrated that blockade of ENT1 prevents vascular leakage associated with hypoxia, presumably through accumulation of vascular Ado. Several lines of evidence support the presumption that elevated Ado attenuates vascular leakage. First, Nozik-Grayck determined that dipyridamole effectively attenuates edema in an isolated perfused lung model of hyperoxia (34). Second, in murine models of hypoxia, decreased vascular Ado is associated with substantially increased vascular leakage (8, 10, 22). In particular, these observations have been made in animals lacking the ability to metabolize adenosine nucleotide precursors into Ado, including cd39 and cd73-null animals (8, 10, 22). Third, the administration of Ado analogues in the form of Ado receptor A2a or A2b agonists, as well as the nonspecific analogue 5′- (N-ethylcarboxamido)-Ado (NECA), protect the vasculature during hypoxia-induced leakage (10). This hypothesis was also tested in a conditional Hif1α mutant mouse, a line that has revealed a protective role for murine HIF-1α in a mucosal inflammation model (3). Using this same mouse line, we confirmed our hypothesis that HIF transcriptionally represses ENT1. Indeed, our findings revealed that colonic epithelial cells lacking Hif1α express increased ENT1 transcript. Of particular interest, analysis of unmanipulated Hif1α mutant mice (i.e., normoxia condition) revealed substantial increases in ENT1. Such findings suggest, as previously speculated (3, 9), that HIF may regulate homeostatic gene expression (i.e., “physiologic hypoxia”) in tissues with normally low pO2 values (e.g., colon) (35).

These results identify ENT repression as a mechanism for attenuating both inflammatory responses and vascular leakage during hypoxia. Interaction of HIF–1 with the ENT1 promoter region is central to the observed inhibition of ENT1 gene expression. Extensions of these findings will determine whether ENT regulation via HIF–1 might function as a pathway for development of therapies for disorders involving vascular leak syndromes or excessive inflammatory responses.

**MATERIALS AND METHODS**

**Cell culture.** HMEC-1 were a gift from F. Candal (Centers for Disease Control, Atlanta, GA) (36) and were cultured as described previously (37). For preparation of experimental HMEC-1 monolayers, confluent endothelial cells were seeded at ~105 cells/cm2 onto either permeable polycarbonate inserts or 100 mm Petri dishes. Endothelial cell purity was assessed by phase microscopic “cobblestone” appearance and uptake of fluorescent acetylated low-density lipoprotein. Where indicated, T84 cells were cultured as described previously (9, 38).

**Measurement of Ado in supernatants.** HMEC-1 were grown to full confluency and exposed as indicated to normoxia or hypoxia. To measure endothelial Ado uptake, 10 nCi of 8-14C-Ado (Sigma-Aldrich) was added to the supernatant, in addition to a 50 μM concentration of “cold” Ado. After indicated time periods, cells were washed with HBSS and lysed in cold H2O. Assessment of intracellular Ado uptake was done by measuring the radioactivity within the lysates by directly diluting the lysate into the scintillation mixture (Ultima Gold HFL–LSC–Cocktail; Packard) and counting decays per minute (Liquid Scintillation Counter Packard 1600).

**Transcriptional analysis.** Conventional PCR was used to verify endothelial ENT1 and ENT2 mRNA regulation, as described previously (38). The PCR reaction contained 1 μM each of the sense primer 5′-AAGGCACCCTGTTCTGTC-3′ and the antisense primer 5′-CTTGGGCT-TGGAGAAC-3′ for ENT1, and 5′-CCTCCATACCCCTCTC-CACC-3′ and 5′-GAGAGAGGGGATTTGCGTC-3′ for ENT2, respectively. The primer set was amplified using increasing numbers of cycles of 94°C for 1 min, 58°C for 2 min, 72°C for 4 min, and a final extension of 72°C for 7 min. The PCR transcripts were visualized on a 1.5% agarose gel containing 5 μg/ml of ethidium bromide. Human β-actin (sense primer, 5′-TGACGGGGTCACCCACACTGTGCCCATCTA-3′ and antisense primer, 5′-CTAGAACATTTCGGGACATGAGG-3′) in identical reactions was used to control for the starting template.

In subsets of experiments, the transcriptional profile of endothelial cells subjected to normobaric hypoxia (12 h) was compared in RNA from control or hypoxic endothelia using quantitative genechip expression arrays (Affymetrix, Inc.) as described before (9). Where indicated, mRNA was also quantified by real-time PCR (iCycler; Bio-Rad Laboratories Inc.) as described previously (9). The primer sets contained 1 μM sense and 1 μM antisense containing SYBR green I (GE Healthcare) in the reaction mixture. Transcript levels and fold change in mRNA were determined as described previously (39). In addition, real-time PCR was performed from RNA isolations of human saphenous vein after ex vivo exposure to hypoxia. After approval by the institutional review board, saphenous vein material was obtained from patients undergoing aorta-coronary bypass surgery. Equal portions of dissected vein tissue were subjected to hypoxia (pO2 20 torr) immediately after the operation for different time points (0, 2, 8, and 24 h). After hypoxia exposure, total RNA was isolated from the tissues and real-time PCR was performed as described above.

**Macromolecule paracellular permeability assay.** Using a modification of methods previously described (18), HMEC-1 were grown on polycarbonate permeable supports (0.4-μm pore, 6.5-mm diam; Costar Corp.) and studied 7–10 d after seeding (2–5 d after confluence). Inserts were placed in HBSS-containing wells (0.9 ml), and HBSS (alone or with indicated concentrations of Ado, with and without 10 μM concentration of dipyridamole) was added to inserts (100 μl). At the start of the assay (t = 0), FITC–labeled dextran 70 kD (concentration 3.5 μM) was added to fluid within the insert. The size of FITC–dextran, 70 kD, approximates that of human albumin, both of which have been used in similar endothelial paracellular permeability models (18). Fluid from opposing well (reservoir) was sampled (50 μl) over 60 min (t = 20, 40, and 60 min). Fluorescence intensity of each sample was measured (excitation, 485 nm; emission, 530 nm; Cytofluor 2300; Millipore Corp., Waters Chromatography) and FITC–dextran concentrations were determined from standard curves generated by serial dilution of FITC–dextran. Paracellular flux was calculated by linear regression of sample fluorescence.

**Immunoblotting experiments.** HMEC-1 were grown to confluency on 100-mm dishes and exposed to indicated experimental conditions. The PN80 and an UVD 170 detector on a reverse-phase column (Grom-Sil 120-ODS-ST-5a; 150 × 3 mm Grom) using a mobile phase gradient from 0 to 25% acetonitrile/0.3 mM KH2PO4 (pH 5) in 5 min. Ultraviolet absorption spectra were measured at 260 nm with a retention time for Ado of 2.5 min.

**Measurement of Ado uptake.** HMEC-1 were grown to full confluency and exposed as indicated to normoxia or hypoxia. To measure endothelial Ado uptake, 10 nCi of 8-14C-Ado (Sigma-Aldrich) was added to the supernatant, in addition to a 50 μM concentration of “cold” Ado. After indicated time periods, cells were washed with HBSS and lysed in cold H2O. Assessment of intracellular Ado uptake was done by measuring the radioactivity within the lysates by directly diluting the lysate into the scintillation mixture (Ultima Gold HFL–LSC–Cocktail; Packard) and counting decays per minute (Liquid Scintillation Counter Packard 1600).
monolayers were lysed for 10 min in 300 μl lysis buffer (150 mM NaCl, 25 mM Tris, pH 8.0, 5 mM EDTA, 2% Triton X-100), and 10% mammalian tissue protease inhibitor cocktail (Sigma-Aldrich), scraped and collected into microfuge tubes. After spinning at 14,000 g to remove cell debris, the pellet was discarded. Proteins were solubilized in nonreducing Laemmli sample buffer and heated to 90°C for 5 min. Samples were resolved on a 12% polyacrylamide gel and transferred to nitrocellulose membranes. The membranes were blocked for 1 h at room temperature in PBS supplemented with 0.2% Tween 20 (PBS-T) and 4% BSA. The membranes were incubated in 5 μg/ml polyclonal rabbit anti-ENT1 in PBS-T for 1 h at room temperature, followed by 10 min washes in PBS-T. The membranes were incubated in 1:3,000 goat anti-rabbit IgG (ICN Biomedicals/Cappel), and conjugated to horseradish peroxidase for 1 h at room temperature. The wash was repeated and proteins were detected by enhanced chemiluminescence. To control for protein loading, blots were stripped and reprobed for β-actin using a mouse monoclonal anti-human β-actin antibody (Abcam Inc.). In subsets of experiments, identical conditions were used to examine HIF-1α protein levels in nuclear lysates derived from HMEC-1. For these purposes, 1 μg/ml of monoclonal mouse anti-HIF-1α (Novus Biologicals) was used.

**ENT1 and ENT2 suppression with RNA interference.** HMEC-1 were either grown on inserts or in 60-mm Petri dishes. Different sets of siRNA directed against human ENT1 and ENT2 were designed using standard molecular tools, synthesized by Dharmacon, and tested with RT-PCR for their efficiency to suppress ENT1 and ENT2 expression at different concentrations. Highest efficiencies were found with 5′-GGGGCUACUGAGUUAUUCCAGdTdT-3′ (sense strand) and 5′-GGUGAAUAUCAGUGGGCCGdTdT-3′ (antisense strand) for ENT1 and 5′-CUCUUCUCAACGGAACUAACTdTdT-3′ (sense strand) and 5′-UUAGGUCCUGUGAGAGAGdTdT-3′ (antisense strand) for ENT2. As non-specific control, the sense strand of the ENT2 oligoribonucleotide was used under identical conditions. As an additional control, knockdown with a set of siRNA directed against human 5′-eucotinucleotide (CD73) was performed using the sense sequence 5′-CUAUCAUCUGGUUCACCGGUATdTdT-3′ (sense strand) and 5′-UACACGGUGUACACGAGAUdTdT-3′ (antisense strand). HMEC-1 loading was accomplished using standard conditions of Fugene6 (Roche Diagnostics Corp.), when the cells reached 40–60% confluence. After 48 h of loading, RNA was isolated as described before or functional assays (flux assays, Ado uptake) were performed.

**ENT1 reporter assays.** Plasmids expressing sequence corresponding to full-length ENT1 (~756 to +1), or the following 5′ truncations: ENT1-344 (bp – 334 to +1), ENT1-282 (bp – 282 to +1), ENT1-225 (bp – 225 to +1), have been previously characterized (unpublished data). As a control for hypoxia, cells were transfected with a PGL3-based HRE plasmid containing four tandem HIF-1 enhancer sequences from the 3′-region of the erythropoietin gene (40). HMEC-1 were cotransfected with constitutively expressed Renilla plasmids using standard methods of overnight transfection using polyfect transfection reagent (QIAGEN). T84 cells were transfected using solution T and a modified form of electroporation (Amaza Inc.). After transfection, cells were subjected to hypoxia or normoxia for 48 h. Luciferase activity was assessed (Turner Designs) using a luciferase assay kit (Promega). All firefly luciferase activity was normalized with respect to the constitutively expressed Renilla luciferase reporter gene.

In subsets of experiments, HIF-1–binding site mutations were introduced in ENT1-756 plasmids using the GeneEditor in vitro site-directed mutagenesis system (Promega). In brief, mutations encoding two nucleotide mutations ENT1 HIF-1 binding sites at positions –721 and –657, respectively, as described previously. All mutations were confirmed by sequencing using pG3L-based primers. Hypoxia inducibility in transient transfectants using such mutated luciferase constructs was exactly as described in the previous paragraph.

**ChiP assay.** ChiP assays were performed using HMEC-1 subjected to normoxia or hypoxia. In brief, 2 × 10⁶ cells were fixed with 1% paraformaldehyde for 10 min. Cross-linking was stopped by the addition of 125 mM glycine, and chromatin derived from isolated nuclei was sheared using a FS50 micro-tip cell sonicator (Fisher Scientific). After centrifugation, supernatants containing sheared chromatin were incubated for 4 h with 5 μg of anti-HIF-1α antibody (Novus Biologicals) or control IgG. Protein A sepharose was added and the incubation continued overnight at 4°C. Immunocomplexes were washed extensively and eluted from the protein A sepharose. The supernatants were transferred to a new tube, and 1 μg/ml of RNase was added and incubated for 5 h at 67°C. Samples were frozen at –80°C and 60 μg/ml proteinase K was added and incubated for 2 h at 45°C. Next, samples were diluted with TE containing 10 μg/ml of RNA following by one extraction with phenol/chloroform and one extraction with chloroform. DNA was precipitated from the samples, washed, dried, and resuspended in 30 μl of autoclaved water. 2 μl of sample was used for each PCR reaction. The sequences of the ENT1 promoter–specific primers spanning the putative HIF-1–binding region were as follows: sense, 5′-GTG-TCATGTGCACATCTCTGCTGGC-3′ and antisense, 5′-CCTGTCCG-CTTCCCCCTTCTAAG-3′. The size of the amplified product resulting from the use of this primer pair was 192 bp.

**Lentiviral vector design, production, and transduction in T84 cells.** The HIV-1 lentiviral vector used was based on a vector previously described in detail (41). In short, the ΔODD variant of HIF-1α (containing a proline→alanine substitution at position 564; the ODD mutant HIF-1α plasmid was provided by H. Franklin Bunn [Harvard Medical School, Boston, MA]) (21) was cloned into the Bam HI/ClaI sites of the lentiviral vector. Virus was produced by first transfecting 293T cells with 3.2 μg of vector, 4.0 μg of the packaging plasmid, and 0.4 μg of REV, TAR, and VSV-G and collecting and filtering the cell supernatant through a 0.45-μm filter. Viral titers were determined by FACS analysis and/or quantitative Southern blot analysis. T84 cells were plated at a density of 2 × 10⁶ cells/10cm² dish 24 h before transduction. These cells were transduced using 2 × 10⁶ infectious units/ml (MOI = 10) of viral supernatant along with 5 μg/ml protamine sulfate for 3 h at 37°C. Cells were fed and propagated in standard T84 media.

**In vivo hypoxia model.** C57BL/6/129 svj wildtype mice were matched according to sex, age, and weight. Total organ vascular permeability was quantified by intravascular administration of Evan’s blue as described previously (42). Animals were exposed to normobaric and normoxic exposure, the animals were killed and the colon, kidney, liver, and lungs were harvested. Organ Evan’s blue concentrations were quantified after formaldehyde extraction (55°C for 2 h) by measuring absorbances at 610 nm with subtraction of reference absorption at 450 nm. Pulmonary edema was assessed in additional experiments. For this purpose, lungs of the animals (n = 6) were collected, weighed, and dried by speed-vac (Eppendorf Vacufuge). Weight differences before and after drying were used to calculate lung water content. For the purpose of quantifying vascular permeability, 0.2 c.c. of Evan’s blue (0.5% in PBS) were injected intravenously. Animals were exposed to normobaric hypoxia (8% O₂, 92% N₂) or room temperature air for 4 h (n = 6 animals per condition). After hypoxia/normoxia exposure, the animals were killed and the colon, kidney, liver, and lungs were harvested. Organ Evan’s blue concentrations were quantified after formaldehyde extraction (55°C for 2 h) by measuring absorbances at 610 nm with subtraction of reference absorption at 450 nm. Pulmonary edema was assessed in additional experiments. For this purpose, lungs of the animals (n = 6) were collected, weighed, and dried by speed-vac (Eppendorf Vacufuge). Weight differences before and after drying were used to calculate lung water content. For the purpose of quantifying PMN tissue concentrations, the animals were exposed to normobaric hypoxia (8% O₂, 92% N₂) or room temperature air for 4 h (n = 4–6 animals per condition). After hypoxia/normoxia exposure, the animals were killed, organs were harvested, and the PMN marker MPO was quantified as previously described (43). Plasma levels of dipyridamole were determined by HPLC with a pump P680 and a Hitachi Fluorescence Detector L-7480 on a reverse-phase column (Genom-Sil 120-ODS-ST-5μ; 150 × 3 mm Grom) using a mobile phase gradient from 0 to 33% acetonitrile/0.3 mM KH₂PO₄, (pH 5) in 10 min. Fluorescence was measured at 404 nm excitation and 480 nm emission wavelength, with a retention time for dipyridamole of 8 min.

In subsets of experiments, colonic mucosal scrapings (enriched in epithelial cells) were obtained from 6–8-wk-old conditional Hif1α mutant mice or littermate controls, as described before (3). Scrapings were homogenized in RNAlater (QIAGEN) using a 22-gauge syringe (Becton Dickin-
Data analysis. Data were compared by two-factor analysis of variance, or by Student’s t test where appropriate. Values are expressed as the mean ± SD from at least three separate experiments.

The authors wish to thank Dr. P. Allen for advice on lentiviral construction and A. Mayer, S. Zug, M. Faigle, and M. Hochgutbrot for technical assistance.

The authors have no conflicting financial interests.

Submitted: 20 January 2005
Accepted: 21 October 2005

REFERENCES


