Antiinflammatory adaptation to hypoxia through adenosine-mediated cullin-1 deneddylation

Joseph Khoury,1 Juan C. Ibla,1 Andrew S. Neish,2 and Sean P. Colgan3

1Department of Anesthesiology, Perioperative and Pain Medicine, Children’s Hospital Boston, Harvard Medical School, Boston, Massachusetts, USA. 2Department of Pathology and Laboratory Medicine, Emory University, Atlanta, Georgia, USA. 3Mucosal Inflammation Program, Division of Gastroenterology, University of Colorado Health Sciences Center, Denver, Colorado, USA.

A major adaptive pathway for hypoxia is hypoxic preconditioning (HPC), a form of endogenous protection that renders cells tolerant to severe challenges of hypoxia. We sought to define the antiinflammatory properties of HPC. cDNA microarray analysis of lung tissue from mice subjected to hypoxia or HPC identified a cluster of NF-κB-regulated genes whose expression is attenuated by HPC. Studies using an NF-κB luciferase reporter assay confirmed a significant suppression of NF-κB activation during HPC. HPC-elicited activity was concomittal, as a soluble supernatant from HPC-treated cells, and the active fraction was purified and identified as adenosine (Ado). Guided by recent studies demonstrating bacterial inhibition of NF-κB through cullin-1 (Cul-1) deneddylation, we found a dose-dependent deneddylation of Cul-1 by Ado receptor stimulation predominantly mediated by the Ado A2B receptor subtype. Further, siRNA-mediated repression of CSN5, a subunit of the COP9 signalsome responsible for deneddylation of Cul-1, partially reversed HPC-mediated inhibition of NF-κB. Cul-1 deneddylation was evident in a murine model of HPC and lost in animals lacking extracellular Ado (Cd73−/− mice). Taken together, these results demonstrate that HPC induces extracellular accumulation of Ado and suppresses NF-κB activity through deneddylation of Cul-1. These results define a molecular regulatory pathway by which Ado provides potent antiinflammatory properties.

Introduction

Active inflammatory responses are characterized by significant shifts in tissue metabolism. These changes coincide with increased energy demand and decreased availability of oxygen and can result in significant levels hypoxia (1–3). Thus, it is important to understand the interactions among microenvironmental metabolic changes (e.g., hypoxia) as they relate to the development and resolution of inflammation. Hypoxic preconditioning (HPC) represents a major hypoxia adaptive pathway; however, the molecular mechanisms of HPC remain largely unknown. Recent studies by Zhang et al. (4) have demonstrated that mice subjected to HPC show significant increases in survival when exposed to otherwise lethal levels of hypoxia. Improved lung function was suggested as a major adaptive mechanism, although the ultimate pathways responsible for this effect were not identified.

Regulated protein degradation is an essential feature of cell signaling for many adaptive processes. The proteasomal degradation of IκB proteins that inhibit NF-κB is one such example of rapid response by the cell to signal for cell growth, differentiation, apoptosis, or inflammation. The E3 SCF ubiquitin ligase specific to IκB family members, comprising SKP1, CUL1, and the F-box domain of β-TrCP, is responsible for the polyubiquitination of IκB (5). Activity of the E3 ligase requires the regulatory subunit Cul-1 to be posttranslationally modified by the ubiquitin-like protein Nedd8. Loss of the Nedd8 modification is mediated by the COP9 signalsome and results in loss of SCF activity. Deneddylated Cul-1 is incapable of ubiquitination of IκB — hence the inactivation of NF-κB (6).

Physiologic adaptation to hypoxia is an area of intense investigation. In this regard, it is widely accepted that adenosine (Ado) is a critical mediator during ischemia and hypoxia (7) and contributes to conditions as diverse as inflammation and carcinogenesis (8–12). While the source of interstitial Ado in hypoxic tissue has been the basis of much debate, it is now appreciated that inhibition of Ado kinase and the dephosphorylation of ATP and AMP by surface apyrases (e.g., CD39) and ecto-5’-nucleotidase (CD73), respectively, represent the major pathways of extracellular Ado liberation during oxygen supply imbalances (13–15). Once liberated in the extracellular space, Ado is either recycled (e.g., through dipyridamole-sensitive carriers) or interacts with cell-surface Ado receptors (10). Presently, 4 subtypes of G protein–coupled Ado receptors have been identified, designated A1, A2A, A2B, and A3, and are classified according to utilization of pertussis toxin–sensitive pathways (A1 and A3) or adenylate cyclase (A2A and A2B) (10). Here, we sought to investigate endogenous mechanisms underlying HPC adaptive responses to hypoxia, particularly as they relate to inflammation.

Results

HPC attenuates NF-κB activity via the release of a soluble factor. Initially, we subjected mice to protocols of HPC similar to those of Zhang et al. (4) (see Methods for protocol). Our results confirmed these previous findings and indicated that HPC-adapted animals survive significantly longer when challenged with severe hypoxia (fractional inspired O2 [FiO2] 5%) (data not shown). These results led us to define the underlying mechanisms of HPC protection.
To this end, we profiled mRNA transcripts from pulmonary tissue of mice subjected to normoxia, hypoxia, or HPC (see Methods for detailed protocols) using cDNA microarray analysis. This profiling approach identified HPC-dependent repression of a major cluster of NF-κB-mediated inflammatory genes (Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI30049DS1). Given the association of NF-κB activation with hypoxia (16), we pursued the antiinflammatory aspects of HPC as they relate to NF-κB.

To define the functional attributes of NF-κB inhibition by HPC, we used an NF-κB luciferase reporter. HeLa cells were transfected with the pNRE-Luc vector followed by exposure to normoxia, anoxia (0% O₂, 18 hours), or HPC followed by anoxia (see HPC protocol in Methods). These studies revealed that cells subjected to HPC displayed a significant attenuation of NF-κB activity compared with those exposed to anoxia alone (Figure 1A). Likewise, a well-established reporter gene for NF-κB activation is IkBα (17). In the course of these experiments, we examined the induction of IkBα by hypoxia and whether such induction was attenuated by HPC. As shown in Supplemental Figure 1, hypoxia was a strong stimulus for induction of IkBα mRNA in HeLa cells (10.4 ± 2.0-fold increase compared with normoxia control; P < 0.01). Parallel examination in HeLa cells subjected to HPC revealed a complete loss of hypoxia-induced IkBα (0.32 ± 0.04-fold change compared with normoxia control; P < 0.01), thus confirming our findings from the NF-κB luciferase reporter experiments. Such findings provide in vitro confirmation of our findings in vivo.

We next addressed whether HPC-induced antiinflammation resulted from the release of a transferrable factor. Here, we harvested soluble supernatants from HPC-exposed cells and examined hypoxia-mediated induction of NF-κB using pNF-κB-Luc–transfected HeLa cells. These studies established the existence of a soluble factor that, when transferred to naive cells, confers the HPC phenotype of attenuated NF-κB activation (51% ± 5.2% decrease; P < 0.05; Figure 1B). We next used standard biophysical approaches to identify the soluble factor (see Methods). As shown in Figure 1C, a soluble HPC factor of smaller than 5 kDa was evident in supernatants filtered through successively smaller molecular weight cutoffs (i.e., 100-, 30-, and 5-kDa filters). These results suggested that a small, stable molecule was responsible for the functional attenuation of NF-κB activation by HPC. Spectrometric analysis of the filtrates showed a prominent chromophore at 260 nm (data not shown), indicating the likely existence of a nucleoside/nucleotide. Further analysis of the active fraction by HPLC revealed high concentrations of Ado within the HPC-conditioned sample (Figure 2A). Moreover, Ado levels increased with successive cycles of HPC (Figure 2B). To determine the role of Ado in the HPC phenotype, we used the nonspecific Ado receptor antagonist 8-phenyl-theophylline and demonstrated a significant attenuation in inhibition of NF-κB by HPC-conditioned supernatants (Figure 2C). These findings of HPC-induced Ado are consistent with previous results implicating Ado in cytotoxic protection afforded by ischemic preconditioning, particularly in cardiac tissues (18, 19). While the source of interstitial Ado in hypoxic tissue has been the basis of much debate, it is now appreciated that inhibition of Ado kinase and the dephosphorylation of AMP by surface apyrases and ecto-5’-nucleotidase (CD73) represent the major pathways of extracellular Ado release during oxygen supply imbalances (13–15). In this regard, Ado and CD73 have been strongly implicated in preconditioning of cardiac tissue (10, 20).

Ado analogs inhibit TNF-α–mediated NF-κB activation. Mechanisms of Ado action in inflammatory processes are currently an area of intense investigation (21–23). To extend the findings of HPC-mediated antiinflammation by Ado, we invoked a series of experiments to define mechanisms of Ado action on NF-κB activity. While a number of mechanisms have been suggested to mediate the endogenous inactivation of NF-κB, each of these pathways...
converge on changes in the inducible degradation of IκB (24, 25). Thus, we determined whether Ado influences IκBα stability. For this purpose, HeLa cells were exposed to the stable Ado analog N-ethyl-carbamido-adenosine (NECA), challenged with an NF-κB activator (TNF-α), and immunoblotted for IκBα (Figure 3A). To obtain additional evidence, we examined the cytoplasmic-to-nuclear translocation of the p65 subunit NF-κB, an assay commonly used to determine NF-κB activation. Compared with exposure to TNF-α alone (Figure 3D), treatment with both NECA and TNF-α significantly attenuated TNF-α–stimulated p65 translocation (Figure 3E). No differences in p65 localization were noted between buffer alone and NECA alone (Figure 3, B and C).

Recent studies identified a mechanism of inhibition of NF-κB by probiotic bacteria (26). These studies implicated the active neddylation of Cul-1 by bacteria and its direct inactivation of NF-κB. Importantly, this cullin-1 neddylation occurs in the presence of normal IκBα phosphorylation patterns (26), such as we observed with HPC. These findings with probiotic bacteria were based on the fact that ubiquitination of phosphorylated IκB occurs through interactions with the SCF E3 ubiquitin ligase complex (consisting of Skp1, Cul-1, and β-TrCP F-box) (24). This SCF complex is active only when Cul-1 is covalently modified by the ubiquitin-like protein Nedd8, and, hence, the neddylation status of Cul-1 directly reflects NF-κB activation (27). Based on these previous studies, we examined whether Ado receptor activation influenced the neddylation of Cul-1. As shown in Figure 4A, immunoblotting of Cul-1 derived from HeLa cell lysates revealed a higher-molecular-weight band, which has been previously shown to be the neddylated form of Cul-1 (26). Densitometric analysis of these results is shown in Supplemental Figure 2A. An immunoblot directed against the Nedd8 protein revealed a pattern similar to the upper band pattern seen for Cul-1, further demonstrating the presence of Nedd8 modification on Cul-1. Indeed, an NECA concentration–dependent loss of Nedd8 was apparent by Western blot analysis (Figure 4A).

We next demonstrated that these findings are not specific for HeLa cells. Here we used Calu-3 cells, a lung cell line, for purposes...
Ado analogs (NECA) inhibit TNF-α-mediated NF-κB activation. (A) Western blot analysis of the stabilization of IκBα. While NECA stimulation (10 μM; 30 minutes) resulted in no appreciable change in IκBα levels, TNF-α (3 ng/ml; 15 minutes) induced a significant decrease in IκBα. The stabilization of IκBα was reestablished when cells were pretreated with NECA prior to TNF-α stimulation. Similarly, NECA inhibited TNF-α-mediated p65 nuclear translocation, as shown in D (arrowheads). (B) Control conditions. (C) NECA stimulation (10 μM; 30 minutes). (D) TNF-α stimulation (3 ng/ml; 15 minutes). (E) NECA pretreatment (10 μM, 30 minutes) followed by TNF-α stimulation (3 ng/ml; 15 minutes). (F) Representative Western blot of total and phospho-specific IKKβ in HeLa cells subjected to normoxia, hypoxia, and HPC.

**Figure 3**

Ado analogs (NECA) inhibit TNF-α-mediated NF-κB activation. (A) Western blot analysis of the stabilization of IκBα. While NECA stimulation (10 μM; 30 minutes) resulted in no appreciable change in IκBα levels, TNF-α (3 ng/ml; 15 minutes) induced a significant decrease in IκBα. The stabilization of IκBα was reestablished when cells were pretreated with NECA prior to TNF-α stimulation. Similarly, NECA inhibited TNF-α-mediated p65 nuclear translocation, as shown in D (arrowheads). (B) Control conditions. (C) NECA stimulation (10 μM; 30 minutes). (D) TNF-α stimulation (3 ng/ml; 15 minutes). (E) NECA pretreatment (10 μM, 30 minutes) followed by TNF-α stimulation (3 ng/ml; 15 minutes). (F) Representative Western blot of total and phospho-specific IKKβ in HeLa cells subjected to normoxia, hypoxia, and HPC.

es of comparison. As shown in Figure 4B, CaLu-3 cell Cul-1 was similarly deneddylated by NECA when probed for Ned8 by Western blotting, suggesting that this finding is not limited to HeLa cells. Ado receptor stimulation with NECA revealed a concentration-dependent deneddylation of Cul-1, with an EC₅₀ of approximately 30 nM. Densitometric analysis of these results are shown in Supplemental Figure 2B. Moreover, the general Ado receptor antagonist 8-phenyltheophylline (8-PT) potently inhibited NECA-induced deneddylation (Figure 4C).

To define Ado receptor subtype specificity, we utilized heterologous overexpression of individual Ado receptors in HeLa cells, wherein cells were cotransfected with pNF-κB-Luc and examined for inhibition by HPC. As shown in Figure 4D, the Ado receptor determining HPC-mediated inhibition of pNF-κB-Luc activity was predominantly A2B (P < 0.01) and to a lesser extent A1 receptors (P < 0.05).

siRNA-mediated repression of the CSN5 component of the COP9 signalosome inhibits Cul-1 deneddylation. Deneddylation of Cul-1 is normally accomplished through interaction with subunits of the COP9 signalosome (e.g., subunit 5 or CSN5/JAB1) (27). Here, we examined whether repression of CSN5 expression (and thereby inhibition of Cul-1 deneddylation) might influence the antiinflammatory aspects of HPC (i.e., attenuation of NF-κB activation). For these purposes, we transfected HeLa cells with siRNA directed against CSN5/JAB1 and examined Ado and HPC influences on Cul-1 deneddylation and NF-κB translocation, respectively. As shown in Figure 4E, this siRNA strategy decreased CSN5 protein expression by greater than 70% relative to mock transfection. Under these conditions, NECA-induced Cul-1 deneddylation was nearly completely abolished compared with control (Figure 4E), and CSN5-repressed cells lost their ability to precondition with respect to hypoxia-induced NF-κB activation (Figure 4F). These findings confirm our hypothesis that HPC confers an antiinflammatory phenotype through influences on CSN5-mediated neddylation of Cul-1.

HPC offers in vivo protection to mice by decreasing inflammation in lungs and by deneddylating Cul-1. In a proof of principle experiment examining whether Ado production contributes to deneddylation of Cul-1 in vivo, we compared wild-type with Cd73−/− null mice, which lack the critical enzyme for extracellular Ado formation (ecto-5′-nucleotidase or CD73) (28). Cd73−/− or wild-type mice were subjected to protocols of preconditioning followed by hypoxia and lung tissues harvested for analysis. Figure 5A shows explanted whole left lungs from mice exposed to protocols of HPC and revealed that lungs from wild-type animals subjected to hypoxia alone were large and swollen, indicative of significant hemorrhage and edema. Lungs from wild-type animals that had been preconditioned had the macroscopic appearance of normoxic controls. By contrast, lungs of the Cd73−/− mice displayed increased volume in all but the normoxic condition, indicating a nearly complete loss of HPC phenotype in Cd73−/− mice (Supplemental Figure 5). Histology of the left lung from hypoxic animals revealed notable perivascular hemorrhage located in the alveolar spaces, whereas the normoxic and preconditioned mice showed no signs of hemorrhage (Figure 5, B–G). Hemorrhage consists of not only red blood cells, but also plasma, which contributes to edema. Further, periarterial cuffing was observed (Figure 5, H–J) similar to that originally identified by Whayne and Severinghaus in acute hypoxia in rats (29). Hypoxia often leads to both inflammation and edema in the lung; therefore, we used edema formation (tissue water content) as a marker of inflammation and barrier function. To further examine this observation, we measured lung water content as an endpoint for evolving inflammation. As shown in Figure 6A, in wild-type mice, hypoxia alone led to an increase in lung water when compared with normoxia or HPC (P < 0.02). However, lungs of Cd73−/− mice had an increased level of lung water and showed no significant decrease with HPC, thereby confirming our macroscopic observations (Figure 5A and Supplemental Figure 4). Extensions of these findings revealed that mice subjected to protocols of HPC followed by hypoxia display deneddylate Cul-1 in immunoblots from protein derived from lung tissue (Figure 6B). Densitometric analysis of these results are shown in Supplemental Figure 3. Interestingly, in whole lung tissue in vivo, hypoxia increased the ratio of neddylated to unmodified Cul-1, thereby implicating the activation of NF-κB due to an increase in...
neddylation by hypoxia. Conversely, HPC resulted in a near complete loss of Cul-1 neddylation, consistent with NF-κB repression. Similarly, IκB immuno blots confirmed the increased levels of NF-κB in hypoxic lungs and the apparent decrease with HPC, whereas the lungs of the Cd73–/– mice did not show such changes. Taken together, such findings of directed deneddylation by Ado and HPC suggest an active inhibitory loop for NF-κB activation. Notably, Western blots from lung tissue from Cd73–/– mice displayed hyperneddylated Cul-1 and further showed the loss of deneddylation with HPC observed in wild-type animals. This observation thereby confirms our findings that Ado may play an important role in the stabilization of the E3 SCF ubiquitin ligase and its role in the inactivation of NF-κB.

Discussion

Inflammation can be defined as a complex set of interactions among soluble factors and cells that arise in any tissue in response to traumatic, infectious, postischemic, toxic, or autoimmune injury (30). The inflammatory process normally leads to recovery; however, if not properly phased, it could lead to tissue damage. Inflammation is commonly seen in many pulmonary disorders, including high altitude pulmonary edema (31) and hypoxia-induced pulmonary hypertension (32).

Inflammatory pathways are regulated by a limited number of transcription factors, the most important being NF-κB (33). Upon cellular stimulation by a variety of mediators, including cytokines, bacterial toxins, or oxidative stress, a signal transduction cascade is activated, leading to the phosphorylation of IκBα on Ser32 and -36 by the multimeric IKK complex (34). Phosphorylation of IκBα is followed by ubiquitination via the E3 ligase SCFβTRCP and is targeted for proteasomal degradation by the 26S proteasome (35). Once IκBα is degraded, NF-κB translocates to the nucleus and binds to the promoter regions of several proinflammatory genes, inducing their expression and thus amplifying the inflammatory response. The extent of NF-κB activation depends on diverse factors, including the variable E3 activity of the SCFβTRCP complex, which is regulated by a reversible covalent modification with the ubiquitin-like protein Nedd8 (36).
In the current study, we have assessed the role of Ado as an anti-inflammatory molecule released during HPC. Our studies revealed that cells subjected to HPC display a significant attenuation of NF-κB activation compared with those exposed to anoxia alone. A soluble mediator found to be released during HPC was identified as Ado. These findings are consistent with our findings in vivo that lungs of mice subjected to HPC prior to more severe hypoxia display no signs of perivascular hemorrhage, a precursor to NF-κB–mediated inflammation (37, 38).

A recent study by Majumdar and Aggarwal (39) identified Ado as a suppressor of NF-κB activity induced by TNF-α and concluded that it may contribute to its role in suppression of inflammation of the immune system. We further showed that Ado analogs such as NECA are capable of inhibiting NF-κB activation as measured by IκBα degradation by the potent NF-κB activator TNF-α. The exact mechanism by which Ado inhibits NF-κB activation was previously unknown. Our studies focused on one pathway involved in the regulation of NF-κB activation, the ubiquitination of IκB via the E3 ligase SCFβTRCP. Ado and NECA analogs displayed a dose-dependent deneddylation of Cul-1 (Figure 4A). It has been demonstrated that the SCF complex is active only when Cul-1 is covalently modified by the ubiquitin-like protein Nedd8; therefore, the neddylation status of Cul-1 directly reflects NF-κB activation (40). We further examined not only the neddylation status of Cul-1 but also, by direct immunoblotting against Nedd8, whether these effects were inhibited by the general Ado receptor antagonist 8-PT.

The COP9 signalosome (CSN), an 8-subunit complex that regulates protein ubiquitination, posttranscriptionally modifies the cullin subunit of E3-ubiquitin ligases by cleaving off the covalently coupled protein Nedd8 (5, 6). Of the 8 different subunits of CSN, CSN5/JAB1, which has metalloprotease activity, has been established to be important in the role of deneddylation of Cul-1. The cleavage of the Nedd8 moiety is catalyzed not by a conventional thiol protease, but rather by metalloprotease activity centered within the CSN5 subunit (5). In our studies, downregulation of CSN5/JAB1 by siRNA inhibited NECA-induced Cul-1 deneddylation and resulted in a loss of precondensation as measured by NF-κB activation (Figure 4, E and F). These findings confirm the role of Ado released during HPC in conferring an antiinflammatory phenotype through influences on CSN5-mediated neddylation of Cul-1.

The findings were tested in vivo to further elucidate the pathway by which Ado released during HPC confers antiinflammatory protection via NF-κB inhibition. Our in vivo findings are consistent with those of in vitro studies, particularly those showing that Cul-1 is deneddylated following HPC and that stabilization of IκB is maintained. Morphologically and histologically, evidence of antiinflammation was indicated by the gross size and water content of the lung as well as the protection conferred against hemorrhage. Hemorrhage has been well established to be a precursor to NF-κB–mediated inflammation (37, 38). Hemorrhage consists of not only red blood cells, but also plasma, which contributes to pulmonary edema. H&E staining did not reveal plasma infiltration in the alveolar spaces, but edema was evidenced in the wet/dry ratio. With regard to the direct role of NF-κB inhibition in this model, it is difficult to assess the exact contribution of Ado. For example, a number of drugs that have been shown to inhibit NF-κB activation (e.g., pyrrolidine dithiocarbamate) can result in tissue phenotypes and outcomes that closely resemble HPC (41). However, none of the known NF-κB inhibitors used in in vivo settings have sufficient specificity to convincingly distinguish the role of NF-κB in preconditioning. A recent study has shed new light on this topic. Frantz et al. utilized NF-κB p50 subunit knockout mice to investigate the role of NF-κB in myocardial preconditioning (42). These studies revealed a crucial role for p50
in damage resulting from myocardial infarction. Multiple levels of analysis revealed that NF-kB is central to endpoint tissue damage associated with infarction and, likewise, that inhibition of NF-kB is central to the phenotype of ischemic preconditioning. Taken together, these results strongly support our ongoing hypothesis that Ado-mediated inhibition of NF-kB promotes a protective phenotype during preconditioning.

As proof of principle, Cd73-/- mice, which lack the ectonucleotidase to convert AMP to extracellular Ado, do not precondition to hypoxia as compared with wild-type littermates. In these animals, lung water is not decreased, Cul-1 is not deneddylated, and there are no changes in iKB levels. Taken together, our results clarify the importance of Ado as an antiinflammatory molecule released during HPC and its role downstream on the inactivation of the NF-kB pathway.

Cumulatively, these results provide new insight into the mechanisms of actions of Ado and, by association, antiinflammatory aspects of lung HPC. Whether such findings are universal for tissues other than lung is not clear at present. Since individual tissues and cells are known to express different Ado receptor profiles (10), it is likely that specific tissues responses will depend on the relative density of individual Ado receptors. Studies using transient transfection assays identified primarily Ado A2B, and to a lesser extent A1 receptors, in Cul-1 deneddylation. Our previous studies have implicated A2B receptors in vascular and inflammatory protection (14, 43, 44). These observations have been made primarily through the use of available pharmacological inhibitors, as well as molecular tools for in vitro settings. More specific inhibitors as well as A2B receptor-null mice have now become available, which will allow some of these questions to be answered, and the results, at least in part, support our ongoing hypothesis that Ado liberated at sites of inflammatory hypoxia protects via A2B receptor activation. In particular, Yang et al. demonstrated that A2B receptor-null mice are prone to inflammation and that acute inflammatory endpoints (e.g., leukocyte adhesion to the vasculature) are augmented in sepsis models (45). These studies revealed the likelihood that both bone marrow and vascular A2B receptors contribute to this phenotype. Conversely, in more chronic models of inflammation, A2B receptors may be detrimental to disease progression. For instance, Sun et al. utilized the highly specific A2B receptor antagonist CVT-6883 and revealed that following systemic administration, both wild-type and Ado deaminase-deficient animals showed less pulmonary fibrosis and fewer inflammatory endpoints (46). More work will be necessary to define potential differences between acute and chronic responses mediated by the A2B receptor.

Neddylation/deneddylation responses are highly conserved and appear to be universal in all cell types examined (27). Deneddylation reactions on cullin targets via CSN-associated proteolysis is increasingly implicated as a central point for cullin-mediated E3 ubiquitination (47). Notably, other pathways for deneddylated have been reported. For example, the recent identification of the Nedd8-specific proteases NEDP1 and DEN1 (48, 49) have provided new insight into this emerging field. NEDP1/DEN1 appear to have isopeptidase activity capable of directly deneddylating cullin targets (49, 50). Determining whether Ado receptor activation might influence NEDP1/DEN1 activity should prove interesting. Overall, our findings here that Ado deneddylates Cul-1 provide new insight into the use of selective Ado receptor analogs as antiinflammatory-based therapies.

Methods

Murine preconditioning model. C57BL/6 (Charles River Laboratories) male mice of 8–10 weeks of age were used in this study. The Animal Care and Use Committee of Brigham and Women’s Hospital approved all procedures. Whole-body preconditioning was performed in a manner similar to that previously described (4). Briefly, mice were placed in humidified environmental chambers and subjected to FiO2 8% for 10 minutes followed by FiO2 21% for 10 minutes for 3 cycles. The O2 concentrations in the chambers were continuously measured by an O2 analyzer. Following preconditioning, mice were subjected to FiO2 21% for 120 minutes. Preconditioned and nonpreconditioned animals were subsequently placed in the chambers and subjected to FiO2 5% for 10 minutes. Animals were then removed from the chamber and sacrificed, and pulmonary tissue and blood were collected. Total RNA derived from lungs of mice subjected to either normoxia, hypoxia, or HPC were reverse transcribed and
cDNA was arrayed through the Massachusetts General Hospital DNA Core Facility using the PGA mouse v1.1 chip. For each experiment, 3 animals were triplicated.

**Cell culture, treatments, transfections, and in vitro preconditioning.** HeLa cells (ATCC) were maintained in DMEM plus 10% FBS at 37°C in a humidified incubator with 5% CO2 in room air. Where indicated, cells were treated with (all chemicals from Sigma-Aldrich, unless otherwise noted) NECA in DMSO at concentrations varying from 100 μM to 1 μM; Ado in H2O, 1 nM to 100 μM; 8-PT in PBS, 1 μM; or TNF-α (R&D Systems) in PBS, 3 ng/ml. Treatment time is indicated in results. Transfection of cells was carried out in 24-well plates using PolyFect reagent as directed by the manufacturer (Qiagen). Plasmids used in transfections were pNF-κB-Luc (0.4 μg/well; Clontech) or each individual Ado receptor in pCMV6 (OriGene) along with Renilla (2 ng/well). Cellular preconditioning was performed on cells that were plated on either 6- or 24-well plates following a modified in vivo protocol optimized for cells (4). HeLa cells were placed in a hypoxia chamber (Coy Laboratory Products Inc.) in preequilibrated hypoxic HBSS at 2% O2 for 45 minutes. Cells were returned to normoxic conditions (21% O2) for 20 minutes. When cells were returned to hypoxia, media was once again replaced with fresh hypoxic HBSS in order to minimize the effects of oxygen present. This protocol was followed for 3 cycles, and then, where indicated, cells were placed in an anoxia chamber (0% O2) with fresh preequilibrated anoxic HBSS for the same indicated.

**NF-κB reporter assays.** To measure transcriptional activity of NF-κB, HeLa cells in 24-well plates were transfected with pNF-κB-Luc (pNRE; Clontech) at a concentration of 0.4 μg per well along with 0.02 μg of Renilla for 24 hours and then either left at normoxia or subjected to a protocol of preconditioning (2% O2 for 45 minutes followed by 21% O2 for 20 minutes for 3 cycles; fresh hypoxic media was changed prior to the start of each hypoxic interval) and were then either treated with 3 ng/ml TNF-α or subjected to anoxia (0% O2) for an additional 6–24 hours. In experiments where supernatants were studied, supernatants were collected from an additional group of cells following intervals of HPC and placed on normoxic cells for 65 minutes (3 cycles). Following treatments, cells were rinsed in PBS, lysed in passive lysis buffer (Promega) for 15 minutes, and spun down, and 20 μl of lysates was assayed using the Dual-Glo Luciferase assay system (Promega) with the use of a luminesimeter (Turner Biosystems).

**Analysis of soluble factor.** HeLa cells plated in 6-well plates were subjected to preconditioning, and supernatants were collected following each hypoxic event and replaced with fresh hypoxic media at the start of the next cycle. Supernatants were systematically analyzed according to size (filtered through 100-, 30-, and 5-kDa filters), stability (heat, freeze/thaw cycles), and UV chromophores. In each case, the fractions were plated on HeLa cells transfected with pNF-κB-Luc in 24-well plates and assayed as described above. Further analyses of the active fraction by HPLC identified the molecule as Ado and were confirmed by coinjection with internal standards.

**Transcriptional analysis.** Real-time PCR (iCycler; Bio-Rad) was employed to examine kB activation in HeLa cells. Primer sets contained 10 pM each of the sense primer 5′-AACCTGAGCAGACTCATC-3′ and the antisense primer 5′-AGTCTCGAGGGTGTTAGCA-3′. The primer set was amplified using increasing numbers of cycles of 94°C for 1 minute, 60°C for 2 minutes, 72°C for 4 minutes, and a final extension of 72°C for 7 minutes. Samples were controlled for β-actin using the following primers: sense 5′-GGTTGCTTTAGATGCGAAG-3′; antisense 5′-ACTGGAACGTAGAAGTGACAG-3′.162 bp).

**Western blot and immunofluorescence analyses.** Whole-cell lysates were isolated from HeLa cells using reducing buffer (62.5 mM Tris pH 6.8, 2% SDS, 10% glycerol, 5% β-mercaptoethanol, 1% bromphenol blue). An equal number of cells were lysed and boiled. The samples were then resolved on 8% polyacrylamide denaturing gels and transferred to nitrocellulose (Bio-Rad). After transfer, the membranes were stained with ponceau S stain in order to verify equal loading. Antibodies used for Western blotting included rabbit polyclonal anti-κB (1:200, Zymed) and goat polyclonal anti-actin (1:1,000, Santa Cruz Biotechnology Inc.). Blots were washed, and species-matched peroxidase-conjugated secondary antibody was added. Labeled bands from washed blots were detected with enhanced chemiluminescence (Amersham Biosciences). For p65 nuclear translocation studies, HeLa cells were plated on 4-well glass slides (Nalgene Nunc International) and allowed to grow to approximately 70% confluence. Treatment with NECA and or TNF-α was then begun. Cells were fixed in 1% paraformaldehyde/PBS at 4°C for 10 minutes and permeabilized with prechilled 0.2% Triton X-100/PBS/2% BSA. Cells were incubated with rabbit anti-p65 (1:200, Rockland Immunocamicals) in 1% normal goat serum in PBS for 1 hour followed by anti-rabbit Oregon Green 514 (1:100, Molecular Probes; Invitrogen) in the same buffer for 30 minutes. Cell images were captured on a fluorescence microscope.

**Tissue water content.** Whole left lungs were harvested from wild-type or Cd73−/− mice following HPC protocols and were photographed for macroscopic assessment of inflammation. Lungs were then placed in preweighed Eppendorf tubes and placed in a SpeedVac (Thermo Scientific) set for vacuum mode at 65°C for 12 hours. Dry weight was then subtracted from the wet weight to calculate cc water per milligram tissue (51).

**Statistics.** Values are expressed as mean ± SEM. Where appropriate, groups were compared by analysis of variance (ANOVA), otherwise, comparisons between groups were conducted using 2-tailed Student’s t test. A P value of less than 0.05 was considered to be significant.

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Address correspondence to: Juan C. Ibla, Department of Anesthesiology, Perioperative and Pain Medicine, Children’s Hospital, Bader 3, 300 Longwood Avenue, Boston, Massachusetts 02115, USA. Phone: (617) 355-7737; Fax: (617) 730-0894; E-mail: juan.ibla@childrens.harvard.edu.

Joseph Khoury and Juan C. Ibla contributed equally to this work.

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