Adaptation to inflammatory acidity through neutrophil-derived adenosine regulation of SLC26A3

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Acute intestinal inflammation includes the early accumulation of neutrophils (PMN). Based on recent evidence that PMN infiltration “imprints” changes in the local tissue environment through local oxygen depletion and the release of adenine nucleotides, we hypothesized that the interaction between transmigrating PMN and intestinal epithelial cells (IECs) results in inflammatory acidification of the tissue. Using newly developed tools, we revealed that active PMN transepithelial migration (TEM) significantly acidifies the local microenvironment, a decrease of nearly 2 pH units. Using unbiased approaches, we sought to define acid-adaptive pathways elicited by PMN TEM. Given the significant amount of adenosine (Ado) generated during PMN TEM, we profiled the influence of Ado on IECs gene expression by microarray and identified the induction of SLC26A3, the major apical Cl−/HCO3− exchanger in IECs. Utilizing loss- and gain-of-function approaches, as well as murine and human colonoids, we demonstrate that Ado-induced SLC26A3 promotes an adaptive IECs phenotype that buffers local pH during active inflammation. Extending these studies, chronic murine colitis models were used to demonstrate that SLC26A3 expression rebounds during chronic DSS-induced inflammation. In conclusion, Ado signaling during PMN TEM induces an adaptive tissue response to inflammatory acidification through the induction of SLC26A3 expression, thereby promoting pH homeostasis.

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INTRODUCTION

Transmigration of neutrophils (PMN, polymorphonuclear leukocytes) to regions of injury is a hallmark of active intestinal inflammation. Without efficient PMN clearance at sites of infiltration, PMN can accumulate and contribute to bystander tissue damage in diseases such as ulcerative colitis (UC) and Crohn’s disease (CD). PMN infiltration is accompanied by energy-demanding processes such as migration, phagocytosis, and generation of NADPH oxidase burst and are believed to shift the metabolism of inflamed tissues.1,2 For example, recent studies have demonstrated that PMN transepithelial migration (TEM) rapidly depletes microenvironmental O2, stabilizing hypoxia-inducible factor. This results in an increase in anaerobic glycolysis and induction of multiple targets along the nucleotide metabolism pathway.3,4 It has been well established that shifts in glycolysis results in the production of lactic acid, promoting tissue acidification.5–7 In severe UC it has been observed that colonic pH can drop to values less than 5.6,7 PMN appear to be sensitive to changes in extracellular pH.8 Decreases in extracellular pH (pH 6.5–7) have been shown to stimulate H2O2 production, inhibit apoptosis, and extend the functional lifespan of PMN.9 The major mechanism for regulating tissue pH is through SLC26A3, a transmembrane glycoprotein that transports Cl− ions across the cell membrane in exchange for HCO3−.10 Despite the significance of extracellular acidification, relatively little is known about the mechanisms involved in maintaining tissue pH and how the tissue adapts to changes in extracellular pH during intestinal inflammation.

During TEM, PMN release significant amount of adenine nucleotides, including ATP, ADP, and dinucleotide polyphosphates (DNP). Through ecto-nucleoside triphosphate diphosphohydrolase (NTPDase), ecto-5′-nucleotidase (CD73), and extracellular nucleoside pyrophosphatase/phosphodiesterases (ENPPs), intestinal epithelial cells (IECs) rapidly metabolize ATP, ADP, and DNP into adenosine (Ado).11–13 This PMN-derived Ado promotes barrier function in endothelial and epithelial cells of the lung and intestine.16,21 Furthermore, activation of the intestinal epithelial A2B Ado receptor (Adora2B) promotes wound healing and improves epithelial barrier during acute colitis.17,18 It remains unclear to what extent Ado signaling impacts tissue homeostasis during acute inflammation and how such changes influence disease outcome.

In the present work, we define mechanisms and outcomes of Ado signaling in maintaining cellular homeostasis during PMN TEM. Guided by global mRNA profiling to ascertain the transcriptional impact of Ado on IECs, we hypothesized that Ado signaling is an integral component for adapting to and maintaining extracellular pH balance. Validation studies demonstrated that...
Ado-induced SLC26A3 expression is CREB/CAMP dependent and increased expression of SLC26A3 significantly limited the shift in extracellular pH following PMN TEM. Moreover, in examining SLC26A3 expression at multiple time points during inflammation in murine colitis models, we show that SLC26A3 is lost in a time-dependent manner during active inflammation, remains decreased during resolution, and ultimately rebounds during chronic inflammation.

RESULTS

Extracellular acidification during PMN transepithelial migration

Tissue acidification is commonly observed during active inflammation. In an effort to understand mechanisms and adaptation to tissue acidification, we developed a model to monitor extracellular pH in real-time during PMN TEM (Fig. 1a). Briefly, T84 IECs were grown to confluence on the underside of 3.0 µm pore permeable supports. A chemotactic agent (1 µM fMLP) was applied to the apical surface (bottom chamber), and 1 x 10⁶ PMN were applied to the basolateral surface (top chamber) to allow for transmigration in the physiologically-relevant basolateral-to-apical direction. Extracellular pH was monitored using a HydroDish and a SDR system which utilizes a fluorescent based pH sensor positioned directly under the T84 IECs. PMN TEM induced a significant decrease in extracellular pH when compared to IECs alone (Fig. 1b, p < 0.01). To determine if physical contact between activated PMN and the IECs was required for extracellular acidification, T84 IECs were grown to confluence on 0.4 µm pore permeable supports. When grown on the smaller pore inserts there was no significant difference in extracellular pH when comparing IECs alone to IECs in the presence of activated PMN (Fig. 1c). As PMN produce a significant amount of extracellular adenine nucleotides during TEM, we extended the study to determine the contribution of Ado signaling in this extracellular acidification. There was no significant difference in extracellular pH when comparing IECs treated with 100 µM Ado to untreated IECs (Fig. 1d, p = not significant). Such findings suggest that Ado-mediated signaling does not drive extracellular acidification. When PMN, in the absence of IECs, where added to the inserts there was no decrease in extracellular pH; however, PMN in the presence of fMLP resulted in a significant decrease in pH (Supplementary Fig. 1A, p < 0.0001). The overall decrease in pH observed in the activated PMN, −0.718 ± 0.0090, was not significantly different then the decrease in pH observed in IECs alone, −0.758 ± 0.0087. When analyzing the change in pH from non-PMN exposed controls, we see that there is no significant difference in the change in pH when comparing -fMLP, -PMN, Ado exposed, or +PMN/fMLP on 0.4 µM pore inserts (Fig. 1e, p = 0.28 ± 0.14, -0.39 ± 0.024, -0.23 ± 0.06, and −0.13 ± 0.14, respectively). The change in pH from controls in these four conditions was significantly less than the −1.06 ± 0.070 change in pH observed during active TEM (Fig. 1e, p > 0.001). Finally, the addition of fMLP to PMN alone decreased the extracellular pH by 0.66 ± 0.0090, which is significantly less than the 1.06 ± 0.070 pH decrease seen in IECs + PMN/fMLP (p < 0.005). In summary, it is shown that inflammatory acidification requires the TEM of activated PMN.

Extracellular acidification by epithelial-derived lactate

It has been observed that lactate is elevated at sites of inflammation, including UC and CD. To determine the impact of lactate on extracellular acidification we measured secreted lactate levels during the course of PMN TEM. As seen in Fig. 1f extracellular lactate significantly increased at 3 h and remained elevated in the presence of activated PMN, 209 µM ± 10.17, 317 µM ± 49.52, and 244 µM ± 29.84 for 3, 4, and 5 h, respectively (p < 0.0001). There was no significant increase in extracellular lactate when T84 IECs were exposed to PMN or fMLP alone when compared to control T84 IECs. To determine the source of lactate, we measured secreted lactate from PMN and PMN + fMLP in the absence of T84 IECs. PMN, regardless of fMLP exposure, produced at most 12.04 ± 2.29 µM lactate (Supplementary Fig. 1B). This amount of lactate is significantly less than the 50.28 ± 1.41 µM observed in control T84 IECs, p < 0.001. This indicates that lactate is being produced by the IECs and not PMN during TEM. When T84 IECs were grown on 0.4 µM pore inserts, there was no increase in extracellular lactate following PMN TEM when compared to control T84 IECs (Fig. 1f). These observations indicate that physical PMN/IECs interaction is required for both lactate production and extracellular acidification.

To determine the role of lactate in PMN TEM induced extracellular acidification, we monitored changes in pH with increasing concentrations of lactate (Table 1). From the extrapolated standard curve (y = 7.294 – 0.001256x – 1.461e^-x), with a R squared of 0.9827, we calculated the predicted pH based on lactate concentrations. Based on these calculations, the observed pH in both IECs control and PMN exposed groups was significantly lower than the pH predicted based on observed lactate levels alone (Table 1). These findings indicate that lactic acid production plays a role in extracellular acidification, but is not exclusively responsible for the observed acidification during PMN TEM. This is supported by the observations of lactate independent extracellular acidification in tumors.

Epithelial gene expression in response to Ado TEM has been shown to transcriptionally “imprint” gene expression patterns onto epithelia through localized oxygen depletion and the stabilization of HIF. Given the propensity of PMN to elevate tissue Ado levels during TEM, we sought to gain insight into mechanisms of adaptation to inflammatory acidity mediated by PMN-derived Ado. To do this, we profiled gene expression in T84 IECs exposed to Ado (100 µM) for periods of 3 or 6 h. As shown in Fig. 2a, a principal component analysis revealed prominent separation between treatment groups and tight clustering within individual treatment groups (p < 0.0001). The top induced and repressed genes are highlighted in the heatmap displayed in Fig. 2b. Of the top hits from this screen, NR4A2 has previously been characterized as an Ado-responsive gene, thereby providing a level of validation for this approach. Likewise, SIK1 has been shown to respond to elevations in cAMP in other cell types. Of significant interest within this Ado-mediated profiling was the highest induced gene SLC26A3, (also called down-regulated in adenoma or DRA), the major Cl⁻/HCO₃⁻ exchanger expressed on the apical IECs surface. To validate these gene targets, T84 IECs were independently exposed to Ado (100 µM, 6 h). As shown in Fig. 2c, transcript levels for NR4A2, SIK1, and SLC26A3 were examined by qPCR and revealed that each was significantly increased by exposure to Ado (15.21 ± 0.61, 15.62 ± 1.03, and 7.28 ± 0.74-fold increase compared to vehicle, respectively, each p < 0.001).

Given the magnitude of PMN-derived Ado observed during PMN TEM and the role of SLC26A3 as a Cl⁻/HCO₃⁻ exchanger, we pursued molecular analysis of the expression and regulation of SLC26A3 by Ado. To further validate SLC26A3 as an Ado responsive gene, T84 IECs were treated with Ado (100 µM) and RNA was collected at 0, 3, 6, 16, and 24 h. As seen in Fig. 2d, transcript levels of SLC26A3 increased at 3 and 6 h following exposure to Ado (40.2 ± 0.42 and 8.25 ± 0.76-fold increase, p < 0.01) and returned to baseline by 16 h. This qPCR analysis was confirmed with protein expression where Ado exposure resulted in a 3.85 ± 0.78-fold increase in SLC26A3 expression with 24 h treatment with 100 µM Ado (Fig. 2e). To rule out cell line specificity, these results were recapitulated in T84, Caco2, and A549 epithelial cells using a luciferase-based promoter assay. In response to 3 h treatment with Ado (100 µM), SLC26A3 promoter activity was increased in all cell lines tested (Fig. 2f, each p < 0.01). Thus, SLC26A3 regulation by Ado appears to be a generalizable response in epithelial cells.
Increased SLC26A3 expression following PMN TEM
To determine SLC26A3 expression following PMN TEM, we examined SLC26A3 expression in T84 IECs 24 h after PMN TEM. 1 × 10⁶ PMN were migrated across 3.0 μm pore inserts in HBSS + HEPES at 37 °C, after 24 h the inserts were collected and washed with PBS to remove PMN. Protein from four inserts was pooled for each replicate and SLC26A3 expression was examined by western blot. As seen in Fig. 2g, there was a 3.79 ± 0.35 fold increase in SLC26A3 expression in T84 monolayers exposed to activated PMN when compared to T84 monolayers exposed to non-activated PMN.
PMN. The level of SLC26A3 expression in PMN TEM T84 IECs is similar to SLC26A3 expression in T84 IECs treated with 100 µM Ado for 24 h (3.79 ± 0.35 and 2.98 ± 0.18 respectively. In addition to analyzing SLC26A3 expression via western blot, SLC26A3 expression was also determined via immunofluorescence (Fig. 2h).

SLC26A3 expression and localization appeared similar between Ado treated T84 IECs and T84 IECs following TEM with activated PMN.

Ado-induced SLC26A3 expression is dependent on CAMP/CREB

The Ado A2B receptor has been shown to be the predominant Ado receptor on T84 IECs. To confirm this observation, we examined the relative expression of Ado A2A and A2B receptor mRNA. Ado A2B receptor mRNA is expressed 100-fold higher than Ado A2A mRNA in T84 IECs (Fig. 3a, p < 0.0001). To determine the signaling cascade of Adora2B on SLC26A3 expression, T84 IECs were treated with a combination of the Adora2B inhibitor, PSB-603 (1 µM), and Ado (100 µM) and examined for induction of SLC26A3 by protein expression after 24 h (Fig. 3b). When analyzed by densitometry, Adora2B-inhibited cells displayed a slight increase in SLC26A3 expression following Ado treatment when analyzed by densitometry (1.58 ± 0.07 and 1.59 ± 0.17, respectively) (Fig. 3b). We speculate that the slight elevation in SLC26A3 expression following PSB-603 treatment is a result of an off-target effect or that PSB-603 is a partial agonist in T84 IECs. The influence of Adora2B inhibition was also assessed using a commercially available SLC26A3 luciferase promoter. In both T84 and Caco2 IECs containing the SLC26A3 luciferase reporter construct, there was no increase in native luminescence following Ado treatment in cells pre-treated for 1 h with 1 µM PSB-603 when compared to vehicle controls (Supplementary Fig. 2A).

It has been reported previously that forskolin is able to stimulate SLC26A3 expression and activity in a CAMP dependent fashion, we extended these studies to determine the signaling pathway(s) involved in SLC26A3 regulation by Ado. Utilizing pharmacological inhibition, we targeted three significant points within the Ado signaling pathway: namely adenylyl cyclase, CREB, and Erk. We quantified SLC26A3 via mRNA expression in T84 IECs pretreated for 1 h with selective inhibitors followed by a 3 h treatment with Ado (100 µM). Ado induced SLC26A3 expression was significantly inhibited in T84 IECs treated with both the adenylyl cyclase inhibitor and CREB inhibitor; whereas, Erk inhibited T84 IECs had a 13-fold increase in SLC26A3 mRNA expression following treatment with Ado (Fig. 3c). As shown in Fig. 3d and Supplementary Fig. 2B, the transcripational response was validated by protein expression via western blot. T84 IECs pretreated with 1 µM CREB and 10 µM adenylyl cyclase inhibitors had no observable increase in Ado induced SLC26A3 expression at 24 h when compared to cells treated with the inhibitors alone. Cells pretreated with 30 µM Erk inhibitor had a similar increase in SLC26A3 when compared to vehicle control cells treated with Ado (3-fold). It was noted that Erk inhibition alone had a small influence on SLC26A3 expression and given the extensive cross-talk between MAPK/Erk and CAMP signaling we speculate this increase in SLC26A3 is an off-target effect of the inhibitor.

Independent analysis of the SLC26A3 promoter sequence revealed five strong CREB binding sites (Fig. 3e). In order to determine the relative contribution of individual CREB sites in the regulation of SLC26A3, each CREB site was mutated independently. Sanger sequencing was utilized to confirm that each CREB site had four nucleotides within the CREB binding site mutated, with the exception of CREB5 which had two nucleotides altered (Supplementary Fig. 3B). Both T84 and Caco2 IECs were transfected with the SLC26A3 reporter and baseline SLC26A3 expression was determined. As seen in Fig. 3f, in T84 IECs SLC26A3 expression was decreased in promoters containing mutations in CREB sites 2, 3, and 4. In Caco2 IECs, only cells with promoters containing a mutation in the CREB3 site had decreased SLC26A3 expression. To further examine which of the CREB sites are involved in Ado signaling, we examined SLC26A3 expression following Ado treatment (100 µM) in each of the CREB mutants. In both T84 and Caco2 IECs, mutations to the CREB3 site resulted in no increased SLC26A3 expression following Ado treatment; all other CREB mutants displayed a 2–3-fold increase in luminescence when compared to controls (Fig. 3g; Supplementary Fig. 3C).

SLC26A3 promotes fluid homeostasis

To further understand the role of SLC26A3 in Ado signaling, SLC26A3 was knocked down (KD) using CRISPR approaches or overexpressed (OE) with transduction of a full-length open reading frame (ORF) construct in T84 IECs. SLC26A3 was decreased by 73 ± 1.3% in the non-clonal KD T84 IECs and increased by 2.50 ± 0.28-fold in the OE T84 IECs (Fig. 4a, b). The KD and OE status of the T84 IECs can also be seen using immunofluorescence (Fig. 4c). As seen in the Ado exposed T84 IECs (Fig. 2h), SLC26A3 expression in the OE T84 IECs appears to be predominantly apical, but there is also basal lateral expression as well. Additionally, we validated the increased fluid secretion phenotype observed in SLC26A3 knockout mice in our SLC26A3 KD cell line by monitoring Ado and forskolin associated fluid flux. SLC26A3 KD cells showed significant increases in fluid transport from the basal to apical compartment following both Ado (100 µM) and forskolin (10 µM) treatment when compared to both vector controls (Supplementary Fig. 4,A,B p < 0.0001). Interestingly, the Ado associated fluid flux observed in SLC26A3 OE cells was not only decreased, but there was a net flux of 14.44 ± 3.85 µL/cm² from the apical to basal compartment (p < 0.001).

To examine the effect of SLC26A3 expression on chloride secretion the peak short-circuit current (Isc) was calculated following treatment with Ado. When the basal surface was exposed to Ado, decreased expression of SLC26A3 did not alter peak Isc when compared to vector controls. However, SLC26A3 OE cells had a significant increase in peak Isc with a 1.86 ± 0.085-fold change compared to vector control, when treated with Ado (30 µM) (Supplementary Fig. 4C). Next, Ado was applied to the apical side of the inserts and the Isc was measured. As seen in the basally treated cells, SLC26A3 OE cells had an increase in peak Isc. Interestingly, SLC26A3 KD cells also had a significant increase in peak Isc.
Fig. 2  Microarray analysis of adenosine-treated intestinal epithelial cells. a Principal component analysis (PCA) of the Ado microarray. b Top genes induced and repressed by 3 or 6 h 100 μM Ado treatment (n = 3). c Validation of induced expression of NR4A2, SIK1, and SLC26A3 by 6 h 100 μM Ado treatment. Data expressed as fold change compared to respective controls (n = 3). d Time course of SLC26A3 transcript expression following treatment with 100 μM Ado (n = 3). e Representative western blot and densitometry analysis of SLC26A3 expression following 24 h 100 μM Ado treatment (n = 3). f Luciferase promoter assay of induced expression of SLC26A3 in T84, Caco2, and A459 by 3 h 100 μM Ado treatment (n = 3). g Western blot and densitometry of SLC26A3 24 h post PMN TEM (n = 3). h Representative confocal images of DAPI and SLC26A3 expression in T84 IECs following PMN TEM. b-f n = number of independent experiments performed, separate passages of cells were used for each experiment. The data for each experiment was pooled and expressed as the mean ± SEM and p-value determined by T-test. *p < 0.01, **p < 0.001, ***p < 0.0001.
peak Isc when compared in vector controls treated with 30 μM Ado on the apical surface (Supplementary Fig. 4D, p < 0.001). When comparing the OE and KD cells, the OE cells have a larger Isc response to apical application of 30 µM Ado than the KD cells (p < 0.001).

SLC26A3 maintains pH homeostasis during PMN transmigration

Based on SLC26A3’s role as a Cl⁻/bicarbonate transporter and regulator of colonic pH, we next determined the influence of SLC26A3 expression on pH regulation during PMN TEM. As previously described, PMN were transmigrated across confluent...
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Adaptation to infiltrating neutrophils in SLC26A3 by 6 h 100 μM luciferase promoter assay (expression of both SLC26A3 and NR4A2 in the normal human IECs). We then examined SLC26A3 expression in human tissues using qPCR. As seen in Fig. 5a, b, forskolin induced detectable fluid flux (reflected as an increase in intracellular pH) at 6 and 24 h. Ado-induced fluid accumulation was detectable at 24 h. Given this response, we examined SLC26A3 expression in the murine colonoids by qPCR. Similar to responses in transformed cell lines, Ado (100 μM, 3 h) induced a significant increase in SLC26A3 in murine colonoids compared to vehicle controls (Fig. 5c, p < 0.01). Extending these studies, we exposed human derived colonoids to 100 μM Ado for 3 h and examined SLC26A3 and NR4A2 expression by qPCR. As seen in Fig. 5d Ado significantly induced the relative expression of both SLC26A3 and NR4A2 in the normal human colonoids (4.43 ± 1.08 and 8.49 ± 2.17, p < 0.005 and p < 0.0001). As a positive control for CREB/CAMP response, human colonoids were treated with 10 μM forskolin for 3 h and qPCR was used to measure the expression of several CREB responsive genes, SLC26A3, cFOS, NR4A2, and NR4A3.31,34 There was a significant increase in all genes tested following treatment with forskolin (Supplementary Fig. 4F, p < 0.001).

SLC26A3 dysregulation in inflammatory bowel disease

We then examined SLC26A3 expression in human tissues using biopsies from CD, active UC, inactive UC patients, and healthy controls. Immunostaining for SLC26A3 was done to examine SLC26A3 expression (Fig. 5e). In healthy colons, there was a high degree of apical IECs expression of SLC26A3. This pattern was aberrant (Fig. 5e, red arrows) and SLC26A3 expression was significantly repressed in both active and inactive UC and CD when compared to healthy controls (Fig. 5f, p < 0.01). Despite an overall decrease in SLC26A3 expression in UC patients, we observed regions of the colon that had similar expression levels and localization as healthy controls (Fig. 5e, f). This relatively "normal" SLC26A3 expression was seen in all four active UC and four inactive UC patients. These regions of increased SLC26A3 expression were absent in the CD patients.

SLC26A3 expression in murine models of colitis

It has been well established that SLC26A3 expression is repressed in DSS induced colitis. However, it has not been well documented at what point during the development of colitis that SLC26A3 expression is lost. To determine the kinetics of SLC26A3 loss, we treated mice with 3% DSS and collected cohorts at days 1, 3, 5, and 7. As seen in Fig. 6a, SLC26A3 expression, as determined by the fluorescent ratio of SLC26A3:DAPI, begins to diminish after 3 days of DSS treatment and continues to decrease at days 5 and 7 when compared to control mice (0.45 ± 0.040, 0.3589 ± 0.038, 0.25 ± 0.053, and 0.82 ± 0.015 respectively, p-value < 0.02 for all comparisons). The staining pattern of SLC26A3 begins to change starting at day 3 (Fig. 6b, red arrow), the expression of SLC26A3 begins to move from being expressed exclusively on the apical surface to being both apically expressed and expressed in the cytoplasm. Additionally, SLC26A3 can be seen being expressed in the crypt of the villus, which is not seen in control mice.

To examine SLC26A3 expression in the context of chronic inflammation, we utilized a chronic DSS murine model.33 Mice were treated with 3% DSS for 7 days followed by 3 weeks of water before undergoing a second and third round of DSS treatment. Cohorts of mice were collected after the first, second, and third round of DSS. When epithelial SLC26A3 expression was examined in the context of chronic DSS we observed expression increasing after each subsequent round of DSS treatment (Fig. 6c). We see the relative expression of SLC26A3, as measured by SLC26A3 fluorescent intensity normalized to DAPI intensity, significantly lower after the first round of DSS when compared to controls (0.27 ± 0.024 and 0.84 ± 0.063, p < 0.0001). After the second round of DSS, SLC26A3 expression increases over 3-fold when compared to SLC26A3 expression after round 1 (0.61 ± 0.04 and 0.27 ± 0.024 respectively, p < 0.0004). SLC26A3 expression continues to increase following the third round of DSS (1.016 ± 0.10). SLC26A3 expression after the third round of DSS is similar to expression seen in control mice, however, the expression pattern seen after the third round of DSS is very different. As seen in Fig. 6d, there are regions of the colon with SLC26A3 expression patterns similar to controls and regions with diffuse cytoplasmic staining. In addition, SLC26A3 staining is observed in the crypts and in transverse sections, which is not observed in control mice (Fig. 6d, red arrows).

DISCUSSION

An understanding of the metabolic responses to inflammation is an area of significant interest. Recent studies, for example, have implicated shifts in tissue metabolism as important clues that determine the overall outcomes of inflammatory responses.36 Of particular interest in the mucosa is the adaptation to "inflammatory hypoxia", wherein oxygen depletion generated by the PMN
Fig. 4 Impact of SLC26A3 on cellular homeostasis. a, b Representative western blot and densitometry analysis of SLC26A3 expression in CRISPR-mediated lentiviral transduced cells or Orf-mediated lentiviral transduced cells (n = 3). c Confocal images of SLC26A3 expression in vector controls, OE, and KD T84 IECs. Main image is a condensed z-stack and inserts depict the orthogonal view of the stacked image. d, e Analysis of extracellular pH following PMN transmigration over the course of 400 min in KD and OE T84 IECs (n = 6–8). f Analysis of extracellular pH following PMN transmigration over the course of 400 min in T84 IECs pretreated with 100 μM Ado for 24 h. g Change in pH between PMN exposed and non-PMN exposed IECs Vector, KD, and OE cells treated with 0, 10, or 100 μM Ado (n = 3). h Change in intracellular pH between T0 and T120 min in vector control, KD, and OE cells treated with 0, 10, or 100 μM Ado (n = 3). n = number of independent experiments performed, separate passages of cells were used for each experiment. The data for each experiment was pooled and expressed as the mean ± SEM and p-value determined by ANOVA or T-test, *p < 0.01, **p < 0.001.
Fig. 5  SLC26A3 expression in non-transformed cells and active disease.  

**a** Brightfield images of respective colonoids following 6 and 24 h 100 μM Ado or 10 μM forskolin treatment. **b** The size of C57BL/6 derived intestinal colonoids treated with 100 μM Ado or 10 μM forskolin were quantified by measuring the pixels/colonoid at 6 and 24 h (n = 3). **c** Induced SLC26A3 expression by 3 h 100 μM Ado treatment in mouse colonic colonoids via RT-PCR (n = 3). **d** Induced SLC26A3 and NR4A2 expression by 3 h 100 μM Ado treatment in human colonoids via RT-PCR (n = 3). **e** Representative confocal images depicting localization and expression of SLC26A3 (Green) and DAPI (Blue) in control, Crohn’s, active UC, and inactive UC colons. **f** Quantification of epithelial specific SLC26A3 expression via fluorescent intensity ratio between SLC26A3 and DAPI in control, Crohn’s, active UC, and inactive UC colons. Data expressed as both the average of each experiment (right) and individual values (left) (n = 4). n = number of independent experiments performed, separate passages of cells were used for each experiment. The data for each experiment was pooled and expressed as the mean ± SEM and p-value determined by ANOVA or T-test, *p < 0.01, **p < 0.001, ***p < 0.0001.
Fig. 6  
SLC26A3 expression in DSS induced colitis.  

a Quantification of epithelial specific SLC26A3 expression via fluorescent intensity ratio between SLC26A3 and DAPI in control and DSS treated mice. Data expressed as both the average of each experiment (left) and individual values (right) (n = 4).  

b Representative confocal images depicting localization and expression of SLC26A3 (green) and DAPI (blue) in mice treated with 3% DSS or H2O for 1, 3, 5, and 7 days.  

c Quantification of epithelial specific SLC26A3 expression via fluorescent intensity ratio between SLC26A3 and DAPI in control and chronic DSS treated mice. Data expressed as both the average of each experiment (left) and individual values (right) (n = 4).  

d Representative confocal images depicting localization and expression of SLC26A3 (green) and DAPI (blue) in control mice and mice treated with 1, 2, or 3 rounds of 3% DSS.  

n = number of independent experiments performed, separate passages of cells were used for each experiment. The data for each experiment was pooled and expressed as the mean ± SEM and p-value determined by ANOVA or T-test, *p < 0.01, **p < 0.001, ***p < 0.0001.
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SLC26A3 plays an adaptive role in our inflammatory in vitro model. Given the multiple health issues associated with SLC26A3-null mice (e.g., high Cl− content diarrhea, growth retardation and volume depletion), we elected to extend our analysis to define SLC26A3 expression in human inflammatory bowel disease patients. We confirmed previous findings that SLC26A3 expression is decreased in active UC patients, and also show decreased expression in individuals with inactive UC and in the colon of CD patients. While UC and CD are related but distinct diseases, we do not know the nature of these differences in SLC26A3 expression. Likely they reflect some differences in the tissue microenvironment (e.g., cytokines, microflora, other inflammatory mediators), studies that will require significant work to understand.

Using immunohistochemical approaches, we not only confirmed that SLC26A3 expression was decreased in the DSS murine model of mucosal inflammation, but also expanded upon these observations to show the time course of SLC26A3 loss in DSS colitis as well as expression patterns in chronic murine models. The increase in SLC26A3 expression observed in chronic DSS colitis supports our observations that SLC26A3 expression is an adaptive phenotype in response to inflammatory acidity. While limited, however, there is evidence which suggests the PMN infiltration increases with each subsequent round of DSS, supporting the idea that PMN influence the expression of SLC26A3. We speculate that this phenotype represents an adaptive phenotype wherein tissues respond to inflammatory acidity by adapting over longer periods of inflammation (e.g., changes in the inflammatory mediator milieu changes that favor expression of SLC26A3). Our observations made in vitro that PMN TEM significantly induces SLC26A3 expression also supports the possibility that PMN-associated nucleotide signaling could be relevant in the regulation of SLC26A3 in the in vivo setting. This premise is supported by our previous observation that PMN accumulation is strongly associated with HIF stabilization and HIF target gene induction (e.g., GLUT1) in patients with active IBD. In this regard, it is notable that a number of gene products along the adenine nucleotide metabolism and signaling cascade are direct transcriptional targets of HIF. Thus, it is possible that HIF could contribute fundamentally to the regulation of SLC26A3 by PMN-derived nucleotides.

Taken together, these results provide new molecular insight into the role of PMN signaling during active inflammation. Our results highlight, for the first time, an Ado-mediated adaptive tissue response that buffers inflammatory acidification. SLC26A3 promotes an adaptive tissue response, thereby limiting the shifts in pH following PMN TEM. Like previous work highlighting HIF signaling, the present observations implicate transcriptional imprinting of Ado signaling in which transmigration provides mucosal memory after the clearance of PMN and elicits functional resolution responses important for tissue homeostasis.

METHODS

PMN isolation and stimulation

Human neutrophils were isolated from whole venous blood of healthy volunteers as described in detail previously. Briefly, whole venous blood was collected in syringes containing anticoagulant (K2EDTA at 1.8 mg/ml blood). Blood was gently layered over double-density Histopaque gradients (1119/1077) and centrifuged at 700 x g in a swinging bucket rotor...
centrifuge for 30 min without brake. The resulting granulocyte layer was collected and residual red blood cells lysed. PMN were washed with ice-cold HBSS (w/out CaCl₂ or MgCl₂) and used within 2 h of isolation.

Cell culture

T84 (#CCL-248; American Type Culture Collection (ATCC), Manassas, VA, USA), Caco2 (#HTB-37; ATCC), and A549 (#CCL-185; ATCC) human epithelial cell lines were obtained from ATCC and were cultured in 95% air with 5% CO₂ at 37 °C according to instructions provided by ATCC. Where indicated, cells were cultured on 3.0 µm or 0.4 µm pore polyester transwell inserts for 10–14 days to obtain confluent cell monolayers as measured by transepithelial resistance (CoStar, Cambridge, MA, USA). Ado (100 µM) and forskolin (10 µM) from Sigma-Aldrich (St. Louis, MO, USA) were added to sterile filtered HBSS or DMEM, respectively (Sigma-Aldrich). ERK activation inhibitor Peptide I (30 µM), CREB inhibitor (1 µM), 666-15, adenylyl cyclase inhibitor (10 µM), 2’,5’-dideoxyadenosine, and adenosine A2B receptor inhibitor (10 µM), PSB-603, from Sigma-Aldrich were added to sterile DMEM (Sigma-Aldrich). Mouse colonoids created from mouse IECs were isolated from PSB-603, from Sigma-Aldrich were added to sterile DMSO (Sigma-Aldrich) interfaced with an SDR SensorDish Reader (PreSens Precision Sensing) were utilized. To measure real-time changes in extracellular pH HydroDish HD24 (PreSens Precision Sensing, Regensburg, Germany) and a SDR SensorDish Reader (PreSens Precision Sensing) were utilized. Briefer, T84 IECs were grown to confluence on the underside of Transwell 3.0 µm or 0.4 µm pore permeable supports were loaded with 1 µM BCECF-am (Thermo Fisher Scientific) in HBSS with HCO₃⁻ for 60 min at 37 °C. After washing in HBSS with HEPES, the cells were treated with Ado (10 or 100 µM) or forskolin (10 µM). Using a Synergy H1 microplate reader (Biotek) the dual-excitation ratio with λ₁ = 490 nm and λ₂ = 535 nm was measured every minute for 2 h at 37 °C under atmospheric CO₂ levels. pH values were calculated based on values obtained from a standard curve generated each run per standard protocol provided by Thermo Fisher Scientific. For each experiment we averaged the data collected from four technical replicates.

To measure real-time changes in extracellular pH HydroDish HD24 (PreSens Precision Sensing, Regensburg, Germany) and a SDR SensorDish Reader (PreSens Precision Sensing) were utilized. Briefer, T84 IECs were grown to confluence on the underside of Transwell 3.0 µm or 0.4 µm pore permeable supports. The inserts were placed into HBSS with HCO₃⁻ and without HEPES. 1 µM FMLP was added to the apical (bottom) chamber immediately prior to the addition of 1 x 10⁶ PMN to the basolateral side (top) and the pH was monitored every minute for 400 min at 37 °C under atmospheric CO₂ levels. The extracellular pH was also determined with PMN alone and T84 treated with 100 µM Ado on the basolateral surface. For each experiment we averaged the data collected from four technical replicates.

Animal studies

C57BL/6 mice (The Jackson Laboratory) were bred in-house. All experiments were handled according to institutional protocols approved by the institutional committee for animal use. Gender, age, and weight-matched mice were used in DSS studies. DSS (40,000–50,000 MW; MP Biomedicals, Colon, OH, USA) was added to drinking water (3%), followed by monitoring for weight loss over 7 days. Cohorts of 4 mice were collected on day 1, 3, 5, and 7. For chronic DSS, 3% DSS was added to drinking water, followed by monitoring for weight loss over 7 days. DSS was replaced with water for 21 days before adding 3% DSS to the drinking water for another 7 days. This was repeated for a third round. Cohorts of 4 mice were collected after the first, second, and third round of DSS. Post-mortem colon samples were harvested by blunt dissection and the distal 2 cm collected and fixed in 10% buffered-formalin (Sigma-Aldrich) prior to paraffin imbedding and staining for immunofluorescence. Four mice were used for each experimental group.

Human samples

Slide mounted paraffin embedded tissue samples from healthy controls, active UC, inactive UC, and CD, identified by a pathologist, were obtained from the University of Colorado Biorepository (IRB# 14-2012). There were four patients in each group.

Microarray and analysis

T84 IECs grown on inserts and the net fluid movement was measured as described previously with minor modifications following application of 100 µM Ado or 10 µM forskolin to the apical compartment. In brief, the apical solution of confluent T84 IECs monolayers grown on 0.33 cm² permeable supports was replaced with 60 µl of media and layered with 60 µl of warm, sterile mineral oil to minimize evaporation as previously described. After 24 h, the apical solution was collected, centrifuged at 10,000 x g, quantified with a calibrated pipette and weighed on a balance (Sartorius, Bohemia, NY, USA). For each experiment we averaged the data collected from four technical replicates.
arrays were washed and stained in a GeneChip Fluidics Station 450 (Thermo Fisher Scientific) and scanned in a GeneChip Scanner 3000 (Thermo Fisher Scientific). Each condition was performed in triplicate. The data were analyzed as gene expression (fold change) at 3 and 6 h over expression in untreated cells. Principal component analysis was performed with XLSTAT (Addinsoft, Paris, France).

Measurement of lactic acid production
To measure lactic acid produced during PMN TEM a commercially available colorimetric l-lactate assay kit was utilized (AAT Bioquest, Sunnyvale, CA, USA). 1 × 10^6 PMN were migrated across T84 monolayers grown on 0.4 or 3.0 μm pore inserts. Supernatant from the apical surface was collected at various time points during PMN TEM. For each experiment we averaged the data collected from four technical replicates.

Transcriptional analysis
TRizol reagent (Invitrogen) was used to isolate RNA from T84 IECs. RNAeasy Plus (Qiagen) was used to isolate RNA from mouse colon tissue. cDNA was reverse transcribed using the iScript cDNA Synthesis Kit (Bio-Rad, Hercules, CA, USA). PCR analysis was performed using SYBR Green (Applied Biosystems, Carlsbad, CA, USA). Primer sequences used are identified in Supplementary Fig. 1. CREB site mutations identified in the SLC26A3 promoter were mutated using QuikChange Lightening Site-Directed Mutagenesis Kit (Agilent Technology, Santa Clara, CA, USA). Primer sequences were confirmed by Sanger sequencing. Purified plasmid was sent to Quintarabio (San Francisco, CA, USA) and sequenced using the primer 5'-CTAGCAAATAGGCTGTCCC-3'.

Protein analysis and immunofluorescence
Whole cell lysates were extracted into RIPA buffer with protease inhibitor (Roche). Lysates were spun at max RPM for 45 min at 4 °C. Supernatant was collected and quantified for normalization using Pierce BCA protein assay kit (Thermo Fisher Scientific). Then 4× Laemmli sample buffer (Bio-Rad) with 2-mercaptoethanol was added. Samples were not boiled and 20 μg lysate per well was used. Western blotting of these lysates was performed using rabbit polyclonal anti-SLC26A3 ab244452 (Abcam) followed by goat anti-rabbit secondary Ab (Invitrogen) and counter stained with ProLong Gold Antifade with DAPI (Thermo Fisher Scientific).

Luciferase reporter assay
T84, Caco2, and A549 cells at ~60–70% confluency were transfected with 200–400 ng of SLC26A3 Renilla luciferase (Switchgear genomics, Carlsbad, CA, USA) and 20–100 ng control Firefly luciferase plasmids (Promega, Madison, WI, USA). Transfection was performed using Lipofectamine 3000 reagent (Invitrogen) according to manufactures instructions. After 24 h, triplicate wells were treated with vehicle, Ado (100 μM), A2B inhibitor (PSB-603, 1 μM), CREB inhibitor (666-15, 1 μM), 2,5′-dideoxyadenosine (5 μM), or Erk inhibitor (30 μM) for 2 h. Cell lysis was performed with Passive Lysis Buffer (Promega) and assayed using a Dual-Luciferase Reporter Assay (Promega) on a GloMax Multidetection System (Promega). SLC26A3 Renilla signal was normalized to the control Firefly signal to account for transfection efficiency.

The five CREB sites identified in the SLC26A3 promoter were mutated using QuikChange Lightening Site-Directed Mutagenesis Kit (Agilent Technology, Santa Clara, CA, USA). Primer sequences were used are identified in Supplementary Fig. 1. CREB site mutations confirmed by Sanger sequencing. Purified plasmid was sent to Quintarabio (San Francisco, CA, USA) and sequenced using the primer 5'-CTAGCAAATAGGCTGTCCC-3'.
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