CCR6 Regulation of the Actin Cytoskeleton Orchestrates Human Beta Defensin-2- and CCL20-mediated Restitution of Colonic Epithelial Cells*  

Intestinal inflammation is exacerbated by defects in the epithelial barrier and subsequent infiltration of microbes and toxins into the underlying mucosa. Production of chemokines and antimicrobial peptides by an intact epithelium provide the first line of defense against invading organisms. In addition to its antimicrobial actions, human beta defensin-2 (HBD2) may also stimulate the migration of dendritic cells through binding the chemokine receptor CCR6. As human colonic epithelium expresses CCR6, we investigated the potential of HBD2 to stimulate intestinal epithelial migration. Using polarized human intestinal Caco2 and T84 cells and non-transformed IEC6 cells, HBD2 was equipotent to CCL20 in stimulating migration. Neutralizing antibodies confirmed HBD2 and CCL20 engagement to CCR6 were sufficient to induce epithelial cell migration. Consistent with restitution, motogenic concentrations of HBD2 and CCL20 did not induce proliferation. Stimulation with those CCR6 ligands leads to calcium mobilization and elevated active RhoA, phosphorylated myosin light chain, and F-actin accumulation. HBD2 and CCL20 were unable to stimulate migration in the presence of either Rho-kinase or phosphoinositide 3-kinase inhibitors or an intracellular calcium chelator. Together, these data indicate that chemokines up-regulated in human inflammatory disorders enhance barrier repair.

Intestinal epithelial cells actively modulate the innate immune system through regulated production of cytokines, bioactive amines, chemokines, and antimicrobial peptides (1, 2). Chemokines are important innate immune molecules that are prototypic mediators of cell migration and regulate the trafficking of leukocytes through binding G-protein-coupled chemokine receptors (3, 4). Chemokines have also been implicated in several cell biological processes, including cancer metastasis, angiogenesis, and stem cell recruitment (3, 5, 6). These chemotactant molecules can be subdivided into two distinct subsets, inducible chemokines are up-regulated by inflammatory stimuli and constitutive chemokines are minimally regulated by pro-inflammatory cytokine stimulation (4).

Defensins, like chemokines, are highly conserved key host defense molecules that participate in host defense through the direct killing of microbes (7). Unlike alpha defensins, which are produced by Paneth cells at the base of intestinal crypts, beta defensins are produced by intestinal epithelial cells. Phylogenetic studies show that beta defensins are evolutionarily conserved in mammals (7–9) and are characterized by pairing of specific cysteine residues (Cys1–Cys5, Cys2–Cys4, and Cys3–Cys4). Of the four characterized human beta defensins (HBD),2 HBD1 is constitutively expressed, whereas HBD2, HBD3, and HBD4 are inducibly expressed (10). Structurally, HBD1–4 share six conserved cysteine residues and tertiary structure that is key to their biologic activity (10). HBD2 is up-regulated in mucosal inflammatory disorders (11–13).

The current, restricted, model states that chemokines direct the trafficking of damage–provoking or damage-exacerbating immune cells to the gut mucosa (1, 14–17). This model is limited in that it ignores the physiologic contribution of chemokine signaling through their cognate receptors expressed by the cells of the intestinal epithelium. Expression of an array of chemokine receptors by human intestinal epithelial cells makes them robust targets for innate immune mediators produced in host defense responses (17–21). The studies herein support the significant ongoing expansion of the current model and indicate that chemokines up-regulated in human inflammatory disorders enhance barrier repair.

Like the homeostatic chemokine receptor CXCR4, the inducible chemokine receptor CCR6 is expressed by immature dendritic cells and circulating T cells and directs their trafficking to sites of inflammation following binding by the chemokine ligand CCL20 (22–24). CCL20 is prominently expressed by intestinal epithelial cells and up-regulated during mucosal inflammatory disorders, including the inflammatory bowel diseases (IBD) (17, 25, 26). CCR6 is constitutively expressed by the human colonic epithelium and, like its cognate ligand, is up-

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2 The abbreviations used are: HBD, human beta defensin; IBD, inflammatory bowel disease; MLC, myosin light chain; pMLC, phosphorylated myosin light chain; ROCK, Rho-kinase; TER, transepithelial resistance; TGFβ1, transforming growth factor β; PI3K, phosphoinositide 3-kinase; BAPTA-AM, 1,2-bis(2-aminophenoxy)ethane-N,N',N''-tetraacetic acid tetrasodium (acetoxyethyl ester); LPA, lysophosphatidic acid; PBS, phosphate-buffered saline; GST, glutathione S-transferase.
regulated during inflammation (17, 18, 26, 27). The conserved
tertiary structure of HBDs facilitate binding and activating
G-protein-coupled receptors, with human HBD1–4 shown to
variably regulate chemotactic migration via the chemokine
receptors CCR6 and CXCR4 (28–30).
Epithelial expression of CCR6 and production of its ligands,
HBD2 and CCL20, are markedly up-regulated in the course of
inflammatory diseases when the innate epithelial barrier is
compromised. Using epithelial cell model systems we dem-
strate for the first time that HBD2 and CCL20 stimulate
restitutive intestinal cell migration through mobilization of
intracellular calcium, activation of phosphoinositide 3-ki-
nase (PI3K), monomeric RhoGTPase, and myosin light chain
(MLC) signaling pathways. Those distinct, co-regulated path-
WAYS converge upon and regulate reorganization of the F-actin
cytoskeleton to increase epithelial sheet migration. These
results significantly expand the mechanistic role for chemok-
ines and defensins and are consistent with the notion that
HBD2 and CCL20 have dual benefits as frontline defense mol-
ecules through the concomitant killing of microbes and leuko-
cyte recruitment with activation of epithelial wound repair
mechanisms.

EXPERIMENTAL PROCEDURES

Materials—Recombinant HBD2 and human CCL20 was pur-
chased from Peprotech (Rocky Hill, NJ), and were 96 and 99%
pure as defined by the manufacturer. Pertussis toxin was pur-
chased from EMD Biosciences (La Jolla, CA). Recombinant
human CXCL12 and transforming growth factor-β1 (TGFβ1)
were purchased from R&D Systems (Minneapolis, MN). The
Rho-kinase (ROCK) inhibitor Y27632 (K_i = 140 μM) and the
specific PI3K inhibitor LY294002 (IC_50 = 1.4 μM) were pur-
chased from EMD Biosciences. BAFTA-AM and lysophospha-
tidic acid (LPA) were purchased from Sigma (St. Louis, MO).
Alexafluor-488 phalloidin was from Invitrogen. Antibodies for
phospho-myosin light chain (pMLC) and total-myosin light
chain (MLC) were obtained from Cell Signaling Technolo-
gies (Danvers, MA). Antibody to total RhoA was purchased from
Santa Cruz Biotechnology (Santa Cruz, CA). Neutralizing anti-CCR6 antibody was from R&D Systems.
Monoclonal antibody to total RhoA was purchased from
Cytoskeleton (Denver, CO).

Cell Culture—The human intestinal carcinoma cell line
Caco2 was cultured in Dulbecco’s modified Eagle’s medium
(4 g/liter glucose) supplemented with 10% (v/v) heat-inactivated
fetal bovine serum (Omega Scientific, Tarzana, CA), 2 mM
L-glutamine, and 1.5 g/liter NaHCO_3. Human T84 colonic car-
icoma cells (31) were cultured in Dulbecco’s modified Eagle’s
medium/Ham’s F-12 medium (1:1) supplemented with 5% (v/v)
newborn calf serum (Invitrogen) and 2 mM L-glutamine as
described previously (20, 21). The normal, non-transformed rat
small intestinal (IEC-6) cell line (CRL-1592) were cultured in
Dulbecco’s modified Eagle’s medium supplemented with 10% (v/v)
heat-inactivated fetal bovine serum, 2 mM L-glutamine, 1.5g/L NaHCO_3, and 0.1 unit/ml bovine insulin (Invitrogen).

IEC-6 Wounding Assay—Confluent IEC-6 cell monolayers
grown in 60-mm dishes were incubated for 24 h in serum-free
medium, wounded with a sterile razor blade and incubated in
medium alone or in the presence of defined concentrations of
HBD2 or CCL20 for 18 h at 37 °C in 5% CO_2. To assess cell
migration signaling mechanisms, monolayers were pre-treated
for 30 min with Y27632 (10 μM), LY294002 (2–50 μM), or
BAFTA-AM (10 μM) and stimulated in the presence or
absence of HBD2 and CCL20. Photomicrographs were taken
using 100X magnification at 4–5 locations per wound, and
the number of migrated cells was determined by counting
nucleated cells that crossed the wound edge.

T84 and Caco2 Wounding Assay—Polarized T84 and Caco2
cells were grown to confluence in 6-well Transwell inserts (pore
size, 0.4 μm; Corning, Danvers, MA) and transepithelial resis-
tance (TER) measured using a hand-held Millicell-ERS volt-
ohmmeter (Millipore, Billerica, MA). Cells were serum-starved
24 h and wounded with a 0.1- to 10-μl plastic pipette tip (USA
Scientific, Ocala, FL) connected to a bench top vacuum aspira-
or. In our hands, this apparatus consistently established
wounds of between 800 and 1000 μm in diameter. Medium on
wounded polarized monolayers was replaced with serum-free
medium, or serum-free medium containing HBD2 or CCL20
every 24 h throughout the duration of the experiment. CXCL12
(20 ng/ml) or TGFβ1 (5 ng/ml) were assessed as positive con-
trols. Photomicrographs were taken of the circular wounds
using the 4× objective after wounding and each day thereafter,
and the area of each wound was defined using MetaMorph soft-
ware (Molecular Devices, Downingtown, PA). The TER of
wounded monolayers was monitored immediately before
wounds were photographed.

Cell Proliferation Assay—Cell proliferation was measured
using propidium iodide staining and cell cycle analysis. Cells
were stimulated with either HBD2 or CCL20 for 4, 8, 12, and
24 h. Ten percent serum was assessed as a positive control.
Ethanol-fixed cells were stained in 50 μg/ml propidium iodide
(EMD Biosciences) and 10 μg/ml RNase A (Promega, Madison,
WI) and analyzed by flow cytometry.

F-actin Formation—To quantify cellular F-actin content,
IEC-6 cells were grown to 80% confluence and serum-starved
24 h prior to stimulation. Cells were pre-treated for 30 min with
10 μM Y27632 to assess the involvement of ROCK in HBD2-
and CCL20-mediated activation of F-actin. Cells were perme-
bilized with 1% (w/v) saponin in PBS and stained with Alex-
fluor-488 phalloidin for 20 min at 37 °C 5% CO_2. To facilitate
release of the cells from the dish the cells were incubated at
37 °C for 20 min in 50 mM EDTA/PBS. The cells were trans-
ferred to FACS tubes (BD Biosciences, San Jose, CA), washed in
PBS, fixed in 2% (w/v) paraformaldehyde/PBS, and fluores-
cence was measured using flow cytometry (BD Biosciences).

Immunoblot Analysis—IEC-6 cells were grown to 80% con-
fluence and serum-starved 24 h before stimulation with titrated
doses of HBD2 or CCL20. Cells were solubilized in hypotonic
lysis buffer (50 mM Tris-HCl, pH 7.4, 10 mM MgCl_2, and Prote-
ase Inhibitor Mixture Set III (EMD Biosciences)). Lysates were
passed through a pipette tip several times and centrifuged at
3800 rpm for 10 min at 4 °C. Protein concentrations were deter-
mained using a Bradford protein assay kit (BCA kit, Pierce), and
10 μg of protein was size-separated using reducing SDS-PAGE.
Proteins were electrophoretically transferred to polyvinylidene
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difluoride (Immobilon-P, Millipore) for immunoblot analyses as detailed previously (21).

**Fluorescence Microscopy**—IEC-6 cell were grown to 30% confluence, washed in PBS, and fixed in 4% (w/v) paraformaldehyde for 15 min. After a wash step in PBS, the cells were incubated in 1% (w/v) bovine serum albumin/PBS for 30 min, followed by an overnight incubation with a 1:50 dilution of rabbit polyclonal anti-CCR6 antibody (Santa Cruz Biotechnology) that specifically binds CCR6. Cells were washed, and cell surface CCR6 was detected by incubation with an anti-rabbit fluorescein isothiocyanate-conjugated antibody. Cells were counterstained with 4',6-diamidino-2-phenylindole and visualized using a fluorescence microscope at 200×.

**RhoA Activation**—Activated RhoA was detected using the solid-phase G-LISA™ RhoA Activation Assay Biochem Kit™ from Cytoskeleton according to the manufacturer’s instructions. Briefly, IEC-6 cells were grown to 95% confluence and serum-starved overnight. Monolayers were stimulated with optimal doses of HBD2 and CCL20, and 1 μM LPA for 5 min. The cells were solubilized, and RhoA-GTP was detected according to the manufacturer’s instructions. Data were analyzed by measuring light emission in counts per second for 0.1 s using Victor2 Wallac (PerkinElmer Life Sciences, Waltham, MA). Total RhoA and actin were detected from the same cell lysates using immunoblot analysis as described above.

**RhoGTP Immunofluorescence**—Wounded Caco2 monolayers were stimulated with 20 ng/ml HBD2 or 20 ng/ml CCL20 for 20 min. The cells were fixed with 4% (w/v) paraformaldehyde (Kodak Eastman Co., Rochester, NY). Autofluorescence was quenched with 50 mM NH4Cl in PBS, and the cells were permeabilized with 0.3% (v/v) Triton X-100 in PBS for 10 min. Cells were washed in 1% (w/v) bovine serum albumin in PBS wash buffer and blocked 30 min in 5% (w/v) bovine serum albumin/PBS and incubated with 40 μg/ml BD-GST or 40 μg/ml recombinant GST (Upstate, Charlottesville, VA) overnight at 4 °C. The cells were washed in buffer and incubated 1 h with 1 μg/ml mouse-anti-GST (Cell Signaling) or mouse IgG (Molecular Probes). Cells were then stained for F-actin using Alexafluor-595 phalloidin according to the manufacturer’s directions. Cells were visualized using confocal or fluorescence microscopy.

**Calcium Mobilization Assay**—Intracellular calcium mobilization was measured using the Fluo-4NW Assay as we defined previously (32). IEC-6 cells were plated in 96-well white walled plates (BD Biosciences, Franklin Lakes, NJ) and grown to 90% confluence. Cells were serum-starved overnight and loaded with the cell-permeant Fluo-4 AM. HBD2 and CCL20 were added at indicated concentrations, and intracellular calcium flux was measured by fluorescence spectroscopy every 5 s for 220 s (Victor2 Wallac). Background fluorescence for each well was measured for 30 s before addition of ligand, and the average background was subtracted from each value.

**Statistical Analysis**—Differences between unstimulated controls, and experimental samples were analyzed using an unpaired Student’s t test using SigmaStat (Jandel Scientific Software, San Rafael, CA).

### RESULTS

**HBD2 and CCL20 Stimulate Cellular Migration of Model Intestinal Epithelium**—To ascertain the role for inflammation-induced CCR6 ligands on enterocyte restitution, a conventional wound healing model was employed. Cells stimulated with either CCL20 (Fig. 1A) or HBD2 (Fig. 1B) migrated more than the unstimulated control monolayers and equal to the TGFβ1-positive control. Furthermore, stimulation of migration was specific for the inducible defensin HBD2, because IEC-6 monolayers treated with 20 ng/ml of the constitutively expressed defensin HBD1 did not significantly increase migration (Fig. 1C). HBD2 and CCL20 dose-dependently stimulated migration of non-transformed IEC-6 cells (Fig. 1E) consistent with previously published chemotaxis of CCR6-transfected HEK293 cells (33).

Migration in the absence of proliferation, defined as restitution, governs the early processes of barrier repair (34). Therefore, we sought to determine if cellular proliferation contributed to the migratory phenotype observed in monolayers stimulated with HBD2 and CCL20. To this end, serum-starved cells were stained with propidium iodide, and cell cycle analysis was performed. As shown in Fig. 1D, IEC-6 monolayers stimulated with either HBD2 or CCL20 did not have an increased percentage of cells in S-phase compared with unstimulated controls after 24 h. However, cells stimulated with the positive control, 10% fetal bovine serum, had a significant increase in cells undergoing DNA synthesis. The lack of proliferation observed with the migration optimal dose of 20 ng/ml was mirrored at 100 ng/ml or 1000 ng/ml of either HBD2 or CCL20 at 4, 8, 12, or 24 h (data not shown). These results indicate that HBD2 and CCL20 specifically induce restitutive migration of model intestinal epithelium.

**HBD2 and CCL20 Induce Cell Migration of Human Polarized Monolayers**—We next confirmed restitution of those migrating IEC-6 epithelial sheets using two complimentary human polarized model epithelial cell lines. For this, Caco2 and T84 cells were grown until the TER was >300 Ω·cm² or 700 Ω·cm², respectively. Cells were wounded, and closure was calculated by measuring the area of the denuded surface. Human Caco2 epithelial monolayers stimulated with either 20 ng/ml CCL20 (Fig. 2A) or 20 ng/ml HBD2 (Fig. 2B) had increased wound closure after 24 h compared with unstimulated controls. Moreover, HBD2- and CCL20-stimulated wound closure of polarized Caco2 monolayers was equal to TGFβ1 (Fig. 2).

To further strengthen the notion that inflammatory mediators regulate epithelial migration, wounded human T84 monolayers were stimulated with either HBD2 or CCL20. In agreement with our data from the IEC-6 and Caco2 model epithelia, the motogenic 20 ng/ml concentration increased wound closure above the unstimulated controls, an increase paralleled by CXCL12 (Table 1) assessed as a positive control. Consistent with these data, barrier integrity, defined as a measure of TER, demonstrated that polarized model epithelium stimulated with HBD2 or CCL20 increased resistance more rapidly than unstimulated controls (Table 1). In sum, results from three model epithelia indicate that HBD2 and CCL20 regulate intestinal barrier homeostasis.
Neutralization of CCR6 Blocks HBD2- and CCL20-stimulated IEC-6 Cell Migration—HBD2 and CCL20 evoke cellular migration of dendritic cells and neutrophils specifically by binding and activating the chemokine receptor CCR6 (35, 36). Human intestinal epithelial cells in vitro and in vivo express CCR6 (17). Therefore, we next sought to determine if HBD2 and CCL20 utilize CCR6 for cell migration. Because chemokine receptor expression in IEC-6 cells was incomplete, we first confirmed CCR6 expression in that particular model intestinal epithelial cell line. Immunoblot analysis defined expression of CCR6 in cell lysates of IEC-6 cells and Caco2 cells (Fig. 3A).

Immuno-fluorescence microscopy of non-permeabilized IEC-6 cells verified CCR6 localization to the cell surface in a pattern consistent with published reports (Fig. 3B) (37). These results indicate that IEC-6 cells express CCR6 at the cell surface where it is available to bind extracellular ligand.

To define specificity of HBD2 for CCR6 in cell migration, we used a specific neutralizing antibody to block activation of CCR6. Because CCL20 is the cognate ligand for CCR6 we also assessed the ability of the neutralizing antibody to block CCL20-mediated intestinal cell migration as a control (22). IEC-6 monolayers were preincubated with 5 μg/ml CCR6 neutralizing antibody or 5 μg/ml isotype control antibody. Wounded cells were stimulated with 20 ng/ml HBD2 or CCL20, and cellular migration was quantified. Pretreatment with the CCR6 neutralizing antibody inhibited HBD2- and CCL20-mediated cell migration, whereas the isotype control antibody did not block migration (Fig. 3C and D). Migration assays showed that pre-treatment with CCR6 neutralizing antibody or the nonspecific isotype control antibody did not affect TGFβ1 migration (Fig. 3E). These data indicate that HBD2 and CCL20 activate cellular migration specifically through CCR6.

HBD2 and CCL20 Stimulate Accumulation of F-actin, Phosphorylation of MLC, and RhoGTP—We next sought to define signaling molecules involved in HBD2- and CCL20-mediated cell migration. Previous work from our laboratory suggests chemokine receptors activate a canonical wound healing pathway consisting of RhoGTP, Rho-kinase, phospho-myosin light chain, and F-actin accumulation in model intestinal epithelium (21). Moreover, work by others has defined components of this pathway to be key regulators of migration in a variety of cell types (38–42). Therefore, we initially focused on defining the activation of these key cell migration molecules in IEC-6 cells treated with HBD2 and CCL20. Because F-actin accumulation at the leading edge of a migrating monolayer is a major hallmark of cell migration, we first quantified F-actin accumulation in IEC-6 cells using Alexafluor-488-phalloidin and flow cytometry. As shown in Fig. 4 (A and B), HBD2 and
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CCL20 stimulation increased F-actin ~25% above unstimulated cells. MLC is a regulatory subunit of myosin that upon activation by phosphorylation on Ser19 facilitates the assembly of F-actin bundles (42). Therefore, we used immunoblot analyses to show that HBD2 and CCL20 induced Ser19 phosphorylation on MLC, an upstream regulator of F-actin (Fig. 4C). We next examined RhoA as an upstream regulator of pMLC. Rho is activated in its GTP bound form and stimulates pMLC and F-actin bundling through Rho-kinase (ROCK) (40). Using an enzyme-linked immunosorbent assay-based solid-phase assay, we determined that HBD2 and CCL20 activate RhoA in IEC-6 cells (Fig. 4D, top panel). Total RhoA and actin were subsequently assessed in those same cell lysates as a loading control (Fig. 4D). These data indicate that HBD2 and CCL20 activate key molecules regulating the actin cytoskeleton in migrating epithelia.

Next, we sought to verify that inflammatory mediators CCL20 and HBD2 are specifically regulating the actin cytoskeleton through CCR6 effectors. Like all chemokine receptors, CCR6 is a G-protein-coupled receptor activated predominantly via the Ga subunit (3). In fact, CCR6-mediated chemotaxis of immune cells is potently inhibited upon blockade of Ga with pertussis toxin (33, 36). Although not previously described for epithelial migration, pertussis toxin was used to assess if Ga signaling was involved in the activation of RhoA by HBD2 and CCL20. In agreement with our data on CXCL12 (20), pretreatment of IEC-6 cells with pertussis toxin decreased RhoGTP (Fig. 4E) with the concomitant decrease in HBD2- or CCL20-stimulated migration (data not shown). Although the inhibition was not complete, the data support the notion that heterotrimeric proteins coupled to CCR6 are activated and initiate downstream effectors of the actin cytoskeleton.

To ascertain if CCR6-regulated F-actin accumulation was simply a function of epithelial sheet migration, or a more global effector of stimulated epithelium, we examined polarized Caco2 monolayers. Fluorescence microscopy was next used to determine that both active RhoGTP and F-actin bundles increasingly localize at the leading edge of wounded Caco2 monolayers stimulated with optimal concentrations of HBD2 or CCL20 (Fig. 4F). These data indicate that the mechanisms regulating enterocyte migration in human polarized, circular wound model system parallel those activated in migrating IEC-6 epithelial sheets.

ROCK Participates in HBD2- and CCL20-mediated Migration and F-actin Accumulation—ROCK is a direct downstream effector of RhoGTP and controls MLC by inactivating its regulatory phosphatase or directly catalyzing MLC phosphorylation (40). To further dissect the regulatory mechanisms in HBD2- and CCL20-mediated migration, the specific ROCK inhibitor Y27632 was used. Inhibition of ROCK abrogated IEC-6 cell migration (Fig. 5A) and F-actin accumulation stimulated by HBD2 and CCL20 (Fig. 5B). These data indicate that Rho and its immediate downstream effector ROCK participate in both F-actin accumulation and cell migration induced by HBD2 and CCL20.

**TABLE 1**

Increased wound closure and barrier integrity in T84-polarized human model epithelia stimulated with CCR6 ligands

| Stimuli  | Wound closure (%) | TER (Ω·cm²/cm²) | % day 0 |
|----------|-------------------|-----------------|---------|
| None     |                   |                 |         |
| HBD2     |                   |                 |         |
| CCL20    |                   |                 |         |
| CXCL12   |                   |                 |         |

Values are mean ± S.E. (n = 3). TER: transepithelial resistance. Cells were stimulated with an optimal 20 ng/ml concentration of HBD2, CCL20, or CXCL12, and wound closure and TER were measured as defined under “Experimental Procedures.”

* Statistical difference (p ≤ 0.05) between stimulated and unstimulated control monolayers.
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Intracellular Calcium Flux Induced by HBD2 and CCL20 Contributes to Cell Migration—Intracellular calcium flux is an established regulator of F-actin accumulation and cell migration and is stimulated by CCR6 ligands in immune cells (22, 43, 44). Therefore, we first asked if HBD2 and CCL20 regulated calcium mobilization in model intestinal epithelium. The Fluor4NW assay defined for the first time that HBD2 (Fig. 6A) and CCL20 (Fig. 6B) induce a dose-dependent intracellular calcium flux in adherent IEC-6 cells. Calcium mobilization in response to 100 ng/ml of the chemokine CXCL12 was shown as a positive control, because it is known to induce calcium flux in human intestinal cells (19).

Calcium mobilization occurs either by the opening of channels on the plasma membrane allowing extracellular calcium to enter or via release from internal stores located primarily in the endoplasmic reticulum (45). To distinguish between these two mechanisms, we first chelated extracellular calcium with 3 μM EGTA and determined that the initial calcium flux, before 100 s, was not significantly altered following addition of those ligands (Fig. 6C). However, the sustained calcium response after 100 s was impaired in cells treated with EGTA indicating influx of extracellular calcium was responsible for the persistent elevation in intracellular calcium. Next, we treated cells with the intracellular calcium chelator, BAPTA-AM, and ascertained that calcium mobilization was decreased at the 3 μM dose (data not shown) and abolished at the 10 μM dose (Fig. 6D). These data indicate that HBD2 and CCL20 induce release of intracellular calcium stores, which, in turn, stimulate a sustained influx of extracellular calcium consistent with store-operated calcium entry.

Calcium Mobilization Is a Critical Regulator of Epithelial Cell Migration—Upon influx, calcium binds to calmodulin, the kinase primarily responsible for the activation of MLC (42). Because CCR6 ligands stimulate calcium flux in leukocytes (22), we reasoned that calcium mobilization was involved in HBD2- and CCL20-directed migration of wounded epithelial cells. As shown in Fig. 7A, preincubation with 3 μM or 10 μM BAPTA-AM did not affect baseline IEC-6 migration in the wound healing assay. However, pretreatment with 30 μM or 100 μM BAPTA-AM dose-dependently blocked migration, with the latter dose abolishing cell movement. These results indicate that treatment with 10 μM BAPTA-AM does not affect constitutive migration, yet this concentration was sufficient to block HBD2- and CCL20-induced calcium mobilization (Fig. 6D). Furthermore, 10 μM BAPTA-AM did not affect TGFβ1-stimulated wound healing indicating chelation of intracellular calcium did not globally disrupt enterocyte migration signaling (Fig. 7A). In contrast to TGFβ1, 10 μM BAPTA-AM was sufficient to block HBD2- and CCL20-stimulated migration (Fig. 7B), indicating BAPTA-AM specifically interrupts migratory signaling by those CCR6 ligands. Together these data implicate calcium mobilization as a necessary step for the induction of cell migration by that G-protein-coupled receptor.
FIGURE 4. HBD2 and CCL20 regulate the actin cytoskeleton in migrating epithelial cells. A and B, IEC-6 cells were stimulated with 20 ng/ml HBD2, 20 ng/ml CCL20, or LPA (1 μg/ml), assessed as a positive control, and F-actin stained using Alexafluor-488-phalloidin. A, representative flow cytometry histogram of F-actin accumulation 15 min after stimulation. B, mean fluorescence intensity was determined and normalized to unstimulated (no stim) control values. Increased F-actin accumulation in HBD2- or CCL20-stimulated cells as a percent of control. C, increased pMLC in HBD2- or CCL20-stimulated IEC-6 cells. LPA was assessed as a positive control. Cell lysates were analyzed by immunoblot analysis. Total myosin light chain (tMLC) and F-actin were assessed as a loading control. Representative blots from four experiments are shown. D, IEC-6 cells stimulated with 20 ng/ml HBD2, 20 ng/ml CCL20, or 1 μg/ml LPA as a positive control had more activated RhoA than untreated controls. Activated RhoA was analyzed using a solid-phase assay (top panel). Total RhoA and actin were assessed by immunoblot as a loading control and representative data shown (bottom panel). E, treatment with pertussis toxin decreased HBD2- and CCL20-stimulated RhoA activation to unstimulated (no stim) levels. IEC-6 cells were pretreated with 200 ng/ml pertussis toxin (PTx), and activated RhoA was assessed. Representative immunoblots confirmed equal protein loading (bottom panel). F, increased localization of RhoGTP (green) and F-actin (red) at the leading edge of wounded Caco2 monolayers stimulated with 20 ng/ml HBD2 and CCL20 for 20 min. Data are representative of four separate wounds. Values in B, D, and E are mean ± S.E. of 3–5 experiments. Asterisks denote statistically significant difference from untreated cells (p < 0.05).
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PI3K Regulates HBD2- and CCL20-mediated Cell Migration—Having demonstrated a role for Gαi, RhoGTP and calcium in activated CCR6-regulated epithelial cell migration we next addressed the role for PI3K in epithelial cell migration. In leukocytes, chemokine receptor functions are tightly linked with PI3K signaling (46). Moreover, heterotrimeric Gαi1 protein-coupled receptor activation of PI3Kα has previously been shown to regulate the sustained influx of external calcium, a response we demonstrated in HBD2- and CCL20-stimulated IEC-6 cells (Fig. 6) (47). Although a role for activated PI3K in epithelial restitution had not previously been demonstrated, we had previously shown that inhibition with wortmannin or LY294002 potently blocked human T84 colonic epithelial cell migration (20). IEC-6 monolayers pretreated with the specific PI3K inhibitor LY294002 were wounded and stimulated with CCR6 ligands, and migration was assessed. Consistent with a role for PI3K in chemokine receptor-regulated migration, LY294002 dose-dependently inhibited restitution stimulated by 20 ng/ml HBD2 (Fig. 7C). Further, PI3K-dependent migration was mediated in part through activation of Rho (relative RhoGTP levels: HBD2 = 131.8 ± 3.5; HBD2 plus LY294002 = 97.1 ± 24.6; LY294002 = 112 ± 17.1). Based on these data, we propose that HBD2 and CCL20 signal cell migration via interrelated mechanisms consisting of calcium, PI3K, and Rho that lead to increased F-actin accumulation and localization within the migrating epithelial cells.

DISCUSSION

The single layer of epithelial cells lining the mucosal surface of the gastrointestinal tract is a critical component of the mucosal innate immune system and comprises a physical barrier between the external luminal milieu and the internal environment. The intestinal epithelium is injured on a daily basis by a variety of stimuli, including noxious luminal contents, normal digestion, inflammation, interactions with microbes, and pharmaceuticals (48). Therefore, maintenance of this essential innate immune barrier requires the ability of this single layer of cells to efficiently repair wounds and establish polarity to maintain homeostasis (49). Upon injury the intestinal epithelium undergoes a wound repair process that starts with proliferation-independent epithelial cell migration, termed restitution, into the wounded area, whereupon the migrated cells subsequently proliferate and differentiate into mature enterocytes (48–50). Pathologic intestinal inflammation is exacerbated by breakdowns in the epithelial barrier and subsequent penetration of luminal microbes and toxins into the underlying mucosa. An intact barrier, chemokine signaling, and antimicrobial peptides, provide the first line of protection against invading organisms. Together, our data are consistent with the notion that HBD2 and CCL20 are bi-functional host defense molecules that function to prevent penetration of luminal contents by directing dendritic cell trafficking or directly killing microbes and by stimulating efficient barrier repair. These findings significantly expand the model and indicate that secreted innate host defense mediators may also orchestrate epithelial wound repair to further limit entry of noxious stimuli.

Enterocyte migration induced by CCR6 ligands was demonstrated using Caco2- and T84-polarized human model epithelium. Further, both HBD2 and human CCL20 robustly stimulated cellular migration in a model of the non-transformed rat IEC-6 epithelium. It is not surprising that human ligands are functional on rat cell lines given the high degree of conservation among chemokine receptors, chemokines, and beta defensins (8, 9). Structural studies on rodent CCL20 and HBD2 suggest that HBD2 is a simplified version of CCL20 and both contain similar Asp-Leu residues considered responsible for binding CCR6 (8). Although structural studies of rat beta defensins are not available, the residues proposed to be important for HBD2 binding to CCR6 are conserved in several rat beta defensin genes (9, 51). Despite the lack of those structural studies HBD2 has been shown to bind and activate both mouse and rat cells in culture (52). This is a phenomenon shared by chemokine receptors and their ligands. For example, the human chemokine CXCCL12 can bind and activate its receptor CXCR4 on rodent cells (20, 21). Moreover, the ability of HBD2 and human CCL20 to signal in rat cells was verified herein using calcium flux assays, a classic and well established readout of chemokine receptor signaling.

Neutralizing antibodies verified CCR6 as the receptor regulating CCL20 and HBD2 intestinal migration. In additional
studies we showed that motogenic and antimicrobial doses of either CCL20 or HBD2 failed to induce a rapid proliferative response, indicating those CCR6 ligands specifically stimulate restitutive migration. In contrast, a prior report shows that sustained incubation with supraphysiologic doses of CCL20 activates proliferation in Caco2 cells after 72 h of treatment (37). These latter data suggest that CCL20 signaling may have an even broader role in epithelial barrier repair than rapid restitutive migration. We also assessed the ability of a constitutively expressed beta defensin, to stimulate wound healing and found that 20 ng/ml HBD1 was not sufficient to induce migration. Our findings are consistent with previous reports in keratinocytes and neutrophils that show HBD2 but not HBD1 induce chemotaxis and cellular migration (36, 53). These results indicate that cell migration is not a common property of all beta defensins, but specific to at least HBD2. Further studies with HBD3 and HBD4 will determine if these beta defensins can also be categorized as cell migratory.

Other investigators have recently shown that HBD2 induces epithelial migration at 1000 ng/ml in the transformed HT29 intestinal cell line (54). In marked contrast, we chose to focus our studies using 20 ng/ml of HBD2, because that concentration is within bactericidal range of the molecule (55) and approximates the concentration of HBD2 observed in human gastric juices and human bronchoalveolar lavage (56). Although the exact concentration of HBD2 in colonic mucosa remains unclear, studies confirm its presence in human colon and that it is up-regulated during inflammation and IBD (12, 13). In addition, salt concentrations are known to inhibit the anti-microbial functions of beta defensins and therefore may also affect their ability to stimulate migration (7). The 20 ng/ml concentration of HBD2 has been shown to stimulate immune cell chemotaxis, however it is below the 1000 ng/ml dose shown to be effective in dendritic cell chemotaxis (33) and may reflect the fundamental differences between epithelial and leukocyte migration (50). Alternatively, differences in beta-defensin-induced migratory responses may reflect lower salt concentrations at the gut mucosa or selective pressure of intestinal epithelium to be more sensitive to HBD2 to maintain this important immune barrier.

Intestinal permeability defects are associated with several intestinal diseases such as IBD, cancer, radiation injury, enterocolitis, and Celiac disease (57–62). Inflammatory molecules are classically thought to contribute to defects in permeability and exacerbation of disease (63, 64). However, using polarized T84 and Caco2 monolayers we showed that HBD2 and CCL20 enhanced barrier integrity of epithelium as measured by transepithelial resistance. This suggests that inflammatory molecules like HBD2 and CCL20 could be beneficial in preventing or limiting disease in individuals with gut permeability defects.

FIGURE 6. HBD2 and CCL20 induce calcium mobilization. Intracellular calcium mobilization was induced in a dose-dependent manner when IEC-6 cells were treated with titrated concentrations of HBD2 (A), or CCL20 (B), or 100 ng/ml CXCL12 as a positive control. C, calcium flux after 100 s was diminished in EGTA-treated cells. Pre-treatment with 3 μM EGTA minimally regulated the initial (<100 s) calcium flux. D, pre-treatment with the cell-permeant chelator 10 μM BAPTA-AM (BAPTA) abolished calcium mobilization stimulated by 100 ng/ml HBD2, 100 ng/ml CCL20, or the positive control 100 ng/ml CXCL12. Intracellular calcium mobilization was measured every 5 s for 220 s using a pre-loaded fluorescent indicator dye Fluo-4NW. Relative fluorescence units (RFU) were obtained using a fluorescence plate reader and background (bkgrd) was subtracted from each value. Data are representative of 3–5 independent experiments.
Likewise, these data present the possibility that dysregulation of HBD2 and CCL20 contributes to intestinal permeability defects consistent with studies showing that individuals with Crohn’s disease have impaired induction of beta defensins (2).

Despite a battery of molecules having been shown to stimulate restitution the mechanisms by which they elicit their functions are not well known. Therefore, we investigated the mechanism(s) of HBD2- and CCL20-mediated restitution in model epithelium. It is important to note that the intestinal mucosa resides in a highly complex and dynamic milieu. Restitution is dependent on several factors present in vivo that are absent from our model system, including mucin-producing goblet cells, extracellular matrix producing fibroblasts, immune cells, and luminal contents. However, the IEC6, Caco2, and T84 model systems have been successful at predicting cellular mechanisms important in restitution in vivo (65–67).

Previously our laboratory has shown that the constitutively expressed chemokine CXCL12 activates RhoGTP, and in turn its downstream effectors ROCK and phosphorylated MLC leading to the accumulation of F-actin (21). This pathway is classically associated with organization of contractile F-actin bundles, a hallmark of epithelial cell migration (39, 41). Our studies determined that HBD2 and CCL20 similarly activated RhoA, pMLC, and accumulation of F-actin, with ROCK signaling a regulator in CCR6-driven migration and F-actin accumulation. Moreover, we built upon that foundation and determined that intracellular calcium flux was involved in HBD2- and CCL20-mediated cell migration.

Calcium is an established regulator of F-actin and a well defined readout for chemokine receptor activation (44, 45). HBD2 and CCL20 similarly induced intracellular calcium mobilization in a dose-dependent manner. Elevation in intracellular calcium was maintained after chelation of external calcium, whereas mobilization of intracellular calcium measured after 100 s was decreased. Furthermore, intracellular calcium mobilization was abolished after chelation with BAPTA-AM. Together, those data suggest that intracellular calcium flux is derived primarily from internal stores of calcium (45). In support of that notion, chelation of intracellular calcium abolished HBD2- and CCL20-directed cell migration without disrupting TGFβ1-mediated migration. These data suggest that calcium chelation specifically inhibits CCR6-mediated enterocyte cell migration and does not globally disrupt epithelial cell migration. Intracellular calcium mobilization may be a mechanism of cell migration unique to chemokine receptors or other G-protein-coupled receptors.

Calcium, in conjunction with calmodulin, regulates a variety of cellular kinases, including myosin light chain kinase, primarily responsible for phosphorylating MLC (42, 68, 69). Calcium also regulates F-actin formation, although the mechanism for that increase remains poorly understood (44). Therefore, it is conceivable that calcium mobilization is involved in regulating components of the Rho-directed accumulation and reorganization of the F-actin cytoskeleton needed for epithelial cell migration.

Together, our results show HBD2 and CCL20 work through the chemokine receptor CCR6 to activate Rho and PI3K, and mobilize intracellular calcium to evoke reorganization of the actin cytoskeleton. Activation of these regulatory pathways contributes to efficient epithelial migration and mucosal barrier repair. It is important to note that other signaling molecules may also contribute to the wound-healing process. Recent studies have shown that molecules such as Rac, LIM-kinase, cofilin, and mDia (70, 71) are also important in cell migration and may also be involved in CCR6-mediated cell migration.

Mucosal wound healing is a treatment goal for individuals undergoing therapy for IBD (72). This concept was validated in recent studies that show mucosal wound healing is significantly associated with a low risk of colectomy and decreased inflammation (73). These studies strengthen the notion that the ability to rapidly heal wounds that afflict the intestinal epithelium is critical to maintaining homeostasis and prevention of disease. Therefore, factors that stimulate wound healing are of clinical importance as possible therapeutics for IBD. Overall, these migration studies suggest that chemokines and beta defensins are protective host defense molecules that function not only to
recruit immune cells and kill microbes, but also to increase the efficiency wound healing in the gut.

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