Arginine stimulates intestinal cell migration through a focal adhesion kinase dependent mechanism

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Background: L-Arginine is a nutritional supplement that may be useful for promoting intestinal repair. Arginine is metabolised by the oxidative deiminase pathway to form nitric oxide (NO) and by the arginase pathway to yield ornithine and polyamines.

Aims: To determine if arginine stimulates restitution via activation of NO synthesis and/or polyamine synthesis.

Methods: We determined the effects of arginine on cultured intestinal cell migration, NO production, polyamine levels, and activation of focal adhesion kinase, a key mediator of cell migration.

Results: Arginine increased the rate of cell migration in a dose dependent biphasic manner, and was additive with bovine serum concentrate (BSC). Arginine and an NO donor activated focal adhesion kinase (a tyrosine kinase which localises to cell matrix contacts and mediates β1 integrin signalling) after wounding. Arginine stimulated cell migration was dependent on focal adhesion kinase (FAK) signalling, as demonstrated using adenovirus mediated transfection with a kinase negative mutant of FAK. Arginine stimulated migration was dependent on NO production and was blocked by NO synthase inhibitors. Arginine dependent migration required synthesis of polyamines but elevating extracellular arginine concentration above 0.4 mM did not enhance cellular polyamine levels.

Conclusions: These results showed that L-arginine stimulates cell migration through NO and FAK dependent pathways and that combination therapy with arginine and BSC may enhance intestinal restitution via separate and convergent pathways.

After acute mucosal injury, rapid restoration of epithelial continuity depends on migration of uninjured epithelial cells to cover denuded sections of basement membrane. Enterocyte migration is a mitosis independent process that is completed within 6–12 hours. Cell adhesion and migration are mediated by key cytoskeletal and signalling proteins that are organised in lamellipodial extensions. At these focal adhesions, numerous proteins colocalise, including p120 focal adhesion kinase (FAK), talin, alpha-actinin, vinculin, paxillin, p130Cas, and p60c-src. Several of these proteins are phosphorylated and activated by FAK.1–4

Arginine (ARG) has been reported to lessen intestinal damage in animal models of necrotising enterocolitis.5–7 ARG is metabolised by two major pathways in enterocytes: conversion by arginase to ornithine, the precursor of polyamines, and conversion by nitric oxide synthase (NOS) to nitric oxide (NO) and citrulline. Polyamines are polycations that are required for normal formation of lamellipodia and stress fibres during cell migration. NO is a lipophilic free radical that stimulates mucus secretion and water absorption, produces smooth muscle relaxation, and regulates bowel permeability.1–5

In the current studies, we determined the effect of ARG on enterocyte migration through FAK phosphorylation after wounding of cultured cells6 as a first step in intestinal repair. Bovine serum concentrate (BSC) was used as a positive control. BSC has been shown by our group to facilitate intestinal villous regrowth and improve bowel permeability in experimental cryptosporidial diarrhoea,7 and serum activates ornithine decarboxylase (ODC) in intestinal cells.8 BSC is inexpensive and approved by the United States Department of Agriculture for use as a dietary supplement (Immunolin or NutraGammex; Proliant Inc., Ames, Iowa, USA) in health food stores. BSC contains active IgG, IgM, and IgA, transforming growth factor β (TGF-β), and insulin-like growth factor 1 (IGF-1).

METHODS

Chemicals

Acrylamide and bisacrylamide were from National Diagnostics (Atlanta, Georgia, USA). Protease and phosphatase inhibitors (aprotinin, leupeptin, bestatin, 4-nitrophosphoryl phosphate, pepstatin, diithiothreitol, and NP-40) were from Boehringer Mannheim (Indianapolis, Indiana, USA). Herbimycin was from Gibco BRL (Gaithersburg, Maryland, USA). Tyrphostins (AG213 and 216) were from Professor Alex Levitski (Hebrew University, Jerusalem, Israel). BSC was obtained from Proliant Inc. (Ames, Iowa, USA). BSC powder contains approximately 80% protein, of which 60% is albumin and 25% is immunoglobulin G (IgG). The manufacturer has measured significant levels of IGF-1 (6000 ng/g protein) and TGF-β1 (90 ng/g) in BSC. All other chemicals, including 3,3-bis(aminomethyl)-1-hydroxy-2-oxo-1-triazene (Deta-NONOate), were obtained from Sigma (St Louis, Missouri, USA).

Abbreviations: ARG, L-arginine; BSC, bovine serum concentrate; DFMO, difluoromethyl-ornithine; Deta-NONOate, 3,3-bis(aminomethyl)-1-hydroxy-2-oxo-1-triazene; DMEM, Dulbecco’s modified Eagle’s medium; FAK, focal adhesion kinase; FBS, fetal bovine serum; iNOS, inducible nitric oxide synthase; IGF-1, insulin-like growth factor 1; L-N6(1-iminoethyl)lysine; NEC, necrotising enterocolitis; NMMA, L-N6(1-iminoethyl)lysine; NOS, nitric oxide synthase; ODC, ornithine decarboxylase; PBS, phosphate buffered saline; PUT, putrescine; SDS-PAGE, sodium dodecyl sulphate-polyacrylamide gel electrophoresis; TGF-β, transforming growth factor β.
Antibodies

Mouse monoclonal antibody IgG1 to FAK (clone 4.47) was obtained from Upstate Biotechnology (Lake Placid, New Jersey, USA). Mouse monoclonal antipolyarginine (PY-20) and rabbit polyclonal anti-nitrergic oxide synthase II (anti-NOS II) antibodies were obtained from Transduction Laboratories (Lexington, Kentucky, USA).

Cells

We selected IPEC-J2 cells derived from newborn piglet jejunal because of their differentiated characteristics, and Cdx2 transformed IEC-6 cells because of a more differentiated phenotype, including a fourfold increased rate of cell migration.18 19 IPEC-J2 cells were obtained from H B erschneider (North Carolina State University College of Veterinary Medicine, Raleigh, North Carolina, USA). IPEC-J2 cells were grown in Dulbecco’s modified Eagle’s medium (DMEM)/F12 medium with 5% serum, split weekly, and were studied at passages 28–56. The Cdx2 transformed rat crypt cell line IEC-610 10 was obtained from Dr J-Y Wang (University of Maryland, Baltimore, USA). Cdx2 transformed IEC-6 cells were cultured in DMEM with isopropyl-β-D-thiogalactopyranoside (4 mM), which served as the inducer for Cdx2 directed by the LacSwitch system (Stratagene, La Jolla, California, USA), for four weeks prior to the experiments and were studied at passages 5–20.

Migration assay

Cells were plated in six well Costar (Corning, New York, USA) plates. After reaching confluence, cells were serum starved overnight in DMEM to achieve quiescence. DMEM contains 0.4 mM ARG. In studies of Cdx2 transformed IEC-6 cells, cells were transferred to acidic free media for six hours (BME with Earle’s salts; Atlanta Biologicals, Norcross, Georgia, USA). Mitomycin C (2 μg/ml) was added. Preliminary studies showed that mitomycin C fully inhibited IPEC-J2 cell proliferation at 24 hours. Treatments were added 15 minutes before razor injuring, and inhibitors (L-N-monomethyl arginine (NMMA)20 or difluoromethylornithine (DFMO), an irreversible inhibitor of ODC) were added 15 minutes after wounding. Cell migration, and washing, 107 cells were treated with 10% goat serum and then incubated with rabbit anti-NOS antibodies (Transduction Laboratories) (1:200) for one hour at 4°C. After several rinses with PBS, cells were further incubated for 45 minutes with fluorescein conjugated secondary antibody (goat anti-rabbit) in PBS.

Phosphotyrosine analysis

FAK immunoprecipitation and immunoblotting for FAK and phosphorylase were done according to previously published protocols.23 Briefly, cells were “starved” of serum and amino acids in Earle’s balanced salts solution for four hours. After wounding, cell migration, and washing, 106 cells were scraped in lysis buffer (150 mM NaCl, 0.1% NaN3, 50 mM Tris HCl (pH 7.6), 2 mM Na orthovanadate, 2 μg/ml aprotinin, 2 μg/ml leupeptin, 2 mg/ml phenyl methyl sulphonyl fluoride, 0.1% Triton X-100, and 0.1% Na deoxycholate) at 4°C. Cell lysates were then clarified by centrifugation for 10 minutes at 15 000 g and lystate volumes were normalised for protein content. The supernatants were incubated with anti-FAK antibody (Upstate Biotechnology) at 4°C for a minimum of 90 minutes. Rabbit antimouse IgG (Jackson ImmunoResearch, West Grove, Pennsylvania, USA) bound to protein A sepharose was then added and incubated with mixing for a minimum of 90 minutes at 4°C. Beads were sedimented and washed extensively with lysis buffer. Proteins were then released for sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) and blot analysis by boiling in Laemmli sample buffer with 1 mM Na orthovanadate for three minutes. Samples were electrotransferred on 8% SDS-PAGE and transferred to nitrocellulose. After blocking with 1% bovine serum albumin in Tris buffered saline with 0.05% Tween-20 for one hour, membranes were incubated with horseradish peroxidase conjugated PY20 (ICN Biochemicals Inc., Costa Mesa, California, USA) for one hour and developed using enhanced chemiluminescence (Amersham Corp., Arlington Heights, Illinois, USA). After stripping of the antiphosphotyrosine antibody with 100 mM β-mercaptoethanol and 0.2% SDS in 62.5 mM Tris-HCl, pH 6.7, the membrane was washed and then blocked with 5% dry milk in Tris buffered saline with 0.05% Tween-20 for one hour. Anti-FAK immunoblotting was then accomplished using the anti-FAK antibody, horseradish peroxidase conjugated antimouse IgG secondary antibody, and enhanced chemiluminescence.

Adenoviral transfection of IEC-6 cells

Ad5FAK-CD contains a dominant negative version of FAK lacking the N terminus and the kinase domain.24 Ad5Luc contains the luciferase reporter gene 5’ of the adenovirus promoter and was used as a control virus. IEC-6 cells, after a 24 hour period of culture in DMEM with 0.5% serum, were infected with Ad5FAK-CD or Ad5Luc at a multiplicity of infection of 500 for 12 hours in DMEM with 10% fetal bovine serum (FBS). The infection medium was changed to fresh 0.2% FBS medium, and cells were incubated for an additional 24 hours. The efficiency of viral transfection was assessed using an antibody directed to the carboxy terminus of FAK (antibody 5158, produced by L Romer). Western blots of Ad5FAK-CD transfected cells revealed a dense band at a molecular weight of 42–44 kDa, indicating expression of FAK-CD. There was no FAK immunoreactive band at a molecular weight of 45 kDa in control cells.

Statistics

Results in the text and figures are expressed as mean (SEM). Statistical significance of differences (p<0.05) between mean values was assessed with one way ANOVA and a post hoc Tukey’s test. For analysis of additivity of ARG and BSC, we used a regression model with surface area of cell migration as the response and variables ARG, BSC, and
ARG increased cell migration
IPEC-J2 and Cdx2 cell migration was enhanced by ARG after razor wounding in a dose dependent manner (fig 1). Maximal stimulation of migration by ARG was seen with 4 mM ARG, producing a 1.67-fold increase in surface area covered. Arginine stimulated migration as effectively as any other agonist tested, including 100 ng/ml epidermal growth factor (47% enhancement); 1% fetal calf serum (55% enhancement), and prostacyclin (1 μM) (25% enhancement). Significant stimulation (p < 0.05) was observed at an ARG concentration as low as 2 mM. Maximal effect was at 4 mM, with diminishing results above 20 mM ARG. Similar stimulatory effects of ARG on migration rate were observed with untransformed IEC-6 cells (data not shown).

ARG was compared with five other amino acids with respect to stimulation of cell migration. Glutamate, leucine, phenylalanine, proline, and citrulline did not significantly stimulate migration. L-Glutamine produced 46% enhancement and was the only amino acid comparable to ARG. ARG in these comparative studies with other amino acids stimulated migration 1.7-fold (p < 0.05) (n = 3).

ARG stimulated migration is associated with increased tyrosine phosphorylation of FAK
To determine if increased tyrosine phosphorylation is required for ARG stimulated intestinal cell migration, cells

![Figure 1](image1.png)

**Figure 1** (A) Enhancement of IPEC-J2 cell and Cdx2-IEC-6 cell migration by arginine (ARG): dose response. Cells were serum starved overnight and then changed to Dulbecco’s modified Eagle’s medium (DMEM) with different concentrations of ARG (with no serum) one hour prior to wounding. Cells were incubated in DMEM, and additional ARG was added after wounding (DMEM without supplemental ARG contains 0.4 mM ARG). After migration, cells were fixed in 2% formaldehyde in phosphate-buffered saline. Surface area covered by migrating cells was measured 18 hours later for IPEC-J2 cells and six hours later for Cdx2 transformed IEC-6 cells. Computer-assisted morphometry allowed quantification of the area covered in front of a 1 mm wound at 40× magnification. Shown are means (SEM) of four experiments, with triplicate measurements for each experiment. Data were normalised for control cells with 0 mM ARG added to media. (Surface area of migration for IPEC-J2 cells with 0 mM added ARG was 132 ± 7 μm²/μm²/24 hours and for Cdx2 transformed IEC-6 cells was 48 ± 5 μm²/μm²/6 hours.

Assuming linear rates of migration after wounding, our data indicate that Cdx2 transformed IEC-6 cells migrate 1.5-fold faster than IPEC-J2 cells. p < 0.05 compared with control cells.

![Figure 2](image2.png)

**Figure 2** Relationship between tyrosine phosphorylation, phosphorylation of focal adhesion kinase (FAK), and intestinal cell migration. (A) Tyrosine kinase inhibitors inhibited cell migration. Tyrosine kinase inhibitors were added to IPEC-J2 cells 30 minutes before wounding with a razor (AG216 denotes tyrphostin). Surface area of migration was measured as described in figure 1, 24 hours after wounding; n = 3. *p < 0.02 compared with I-arginine (ARG) treated cells. For control (CT)-+AG216 treated cells, surface area migrated was marginally less than in CT cells (p = 0.082). (B) ARG enhanced phosphorylation of FAK: effect of ARG dose and time after wounding. Cdx2 cells were starved with Dulbecco’s modified Eagle’s medium (DMEM) containing no serum and ARG overnight. Then, one hour before wounding, different doses of ARG were added to ARG-free DMEM. Cells were wounded and harvested at 1, 4, 8 hours after wounding. They were immunoprecipitated (IP) with anti-FAK antibody and immunoblotted using either antiphosphotyrosine antibody (PY-20) or anti-FAK (to control for equal loading). Densitometry analysis revealed that relative to that of control unwounded cells, FAK phosphorylation increased in wounded monolayers by twofold at one and four hours. Furthermore, ARG stimulated phosphorylation of FAK by threefold at one hour and by sixfold at four hours. Deto-NONOate and BSC treated cells had identical phosphorylation as control wounded cells at one hour and 2.5-fold increased phosphorylation of FAK at four hours.

ARG+ BSC interaction. The software program used was SigmaStat (Jandel Scientific, San Rafael, California, USA).
were wounded in the presence or absence of an inhibitor of tyrosine kinases, tyrphostin (AG216). Cells were grown on plastic because adhesion to exogenous matrix proteins increases the activation of FAK. Cdx2 transformed IEC-6 cells treated with AG216 migrated at a rate that was ~80% of normal (p = 0.08, n = 3). Cell monolayers were wounded and incubated with control medium or ARG+AG216. Response to ARG was inhibited by addition of AG216 to 75% of that of ARG treated cells (p<0.05) (fig 2). IPEC-J2 cellular response to ARG was blocked by two other tyrosine kinase inhibitors (herbimycin A 10 μM; tyrphostin 47 200 μM, data not shown).

The effect of ARG on tyrosine phosphorylation of FAK was determined by incubating cells with different concentrations of ARG (0–4 mM) after razor wounding. FAK was immunoprecipitated from lysates of each treatment group, and FAK tyrosine phosphorylation was evaluated by western blotting. ARG treatment after wounding increased tyrosine phosphorylation of FAK at 1–4 hours post-wounding. The maximal effect was seen with 2 mM ARG, which increased the phosphotyrosine content of FAK 1.75 (0.2)-fold (p = 0.038, n = 3) at one hour after wounding (fig 2B). Tyrphostin AG216 markedly reduced ARG induced changes in FAK phosphorylation (data not shown). Enhancement of FAK activity by ARG persisted at four and eight hours (fig 2B, 2C).

The NO donor Deta-NONOate also stimulated FAK phosphorylation by 2.5-fold at four hours post-wounding (n = 3, fig 2D).

Role of NO in ARG stimulated migration
We determined if ARG treatment enhanced nitrite levels in the media as an index of NO production. Nitrite levels were measured at 24 hours after injury. The “baseline” concentration of nitrite in the growth medium of normal unwounded IPEC-J2 cells was 6.5 (0.5) μM at 24 hours after plating. Multiple wounding resulted in an increase to 20 (1.4) μM nitrite after 24 hours. Incubating cells with 4 mM ARG more than doubled NO production to 47 (3.7) μM nitrite (p<0.05 v control media containing 0.4 mM ARG) but BSC had no stimulatory effect on nitrite level (24 (1.2) μM) (n = 4). ARG (fig 3A) and Deta-NONOate (at 5–10 μM) increased nitrite levels.

Figure 3  (A) L-Arginine (ARG) dose dependently increased nitrite (NO) levels in media at 24 hours. Values are means (SEM), n = 4. **p < 0.01 compared with control wounded cells. (B) Cell wounding enhanced the level of inducible nitric oxide synthase (iNOS). Cdx 2 transformed IEC-6 cells were wounded multiple times with a plastic pipette tip and washed. At specified time intervals, cells were harvested and 100 μg protein were electrophoresed and blotted with polyclonal anti-NOS II antibody. Densitometric analysis revealed maximal activation of iNOS by four hours. Results in relative densitometric values were: non-wounded (NW) 1 ×; wounded 0 min 1.7 ×; wounded 30 minutes 4.7 ×; wounded four hours 5 ×; and wounded 24 hours 3.3 ×. (C) Cell wounding enhanced cytoplasmic expression of iNOS at the leading edge. Immunocytochemical staining of Cdx-2 transformed IEC-6 cells was perfomed on unwounded and wounded cells. Weak cytoplasmic fluorescence in unwounded cells was transformed to bright cytoplasmic fluorescence which was most evident in cells at the leading edge. Cells were stained and fixed at time points after wounding, as indicated.
levels dose dependently. (Measurement of nitrite levels with Deta-NONOate might not be accurate because the acidic conditions of the Griess reaction induce NO release from NO donors.)

Tissue injury is known to produce an increase in the level of iNOS. To determine if cell injury induced expression of iNOS, cells were multiply wounded and harvested at different times after injury (fig 3B). The results indicated that iNOS was expressed 1.8-fold at the time of wounding, 2.3-fold at 30 minutes after wounding, with a peak of 2.8-fold at four hours, and reduced activation thereafter (fig 3B). Immunofluorescence detection of iNOS in IPEC-J2 cells (fig 3C) revealed a low baseline in unwounded resting monolayers and in freshly wounded cells. A dramatic increase in iNOS was seen in cells at the wound edge at 1–4 hours after wounding.

NMMA18 is an ARG analogue that competitively inhibits NOS.15 We studied the effect of increasing concentrations of NMMA on 2 mM ARG stimulated cell migration (fig 4A). NMMA reduced ARG stimulated IPEC-J2 cell migration in a dose dependent manner, with complete inhibition at 2 mM NMMA. Individually given to IPEC-J2 cells in standard media, NMMA did not inhibit the basal rate of migration (data not shown, n = 4). NMMA completely blocked NO synthesis from ARG in IPEC-J2 cells. NMMA fully inhibited ARG stimulated nitrite production at 24 hours (wounded control cells = 10.4 µM nitrite; wounded ± ARG (4 mM) = 25.7 µM nitrite; wounded ± ARG ± NMMA (4 mM) = 8.2 µM nitrite; wounded ± NMMA = 4.6 µM nitrite).

Similar to NMMA, the specific inhibitor of iNOS, l-N6(1-iminoethyl)lysine (l-Nil), inhibited ARG stimulated migration at 5 µM (data not shown). These findings indicate that NO production is essential for ARG stimulated migration. To further support this relationship, we demonstrated that the NO donor Deta-NONOate individually stimulated IPEC-J2 cell migration by 1.5-fold (data not shown). NMMA inhibition of ARG stimulated migration could be ‘‘rescued’’ by administration of the NO donor Deta-NONOate (50 µM; fig 4B). NMMA did not significantly affect BSC stimulated cell migration (n = 3, data not shown).

**Additive effects of ARG+BSC on enterocyte migration**

BSC at 5 mg/ml stimulated cell migration 1.5 fold (p < 0.05), an effect comparable with that of 4 mM ARG. We could not achieve saturation because of limiting solubility (5 mg/ml) although stimulation with 5 mg/ml was not significantly greater than that with 0.1 mg/ml. Figure 5 shows a photomicrograph of migrating cells treated with ARG (4 mM), BSC (0.1 mg/ml), or combinations of these treatments. ARG plus BSC (‘‘A+B’’) increased IPEC-J2 cell migration significantly more effectively than ARG or BSC alone. In three experiments of 3–4 observations each, the adjusted mean response of control cells was 60 (5) µm²/µm; of ARG treated cells 88 (7) µm²/µm; of BSC treated cells 107 (6) µm²/µm; and of ARG+BSC treated cells 128 (6) µm²/µm wound (p < 0.001 for ARG+BSC compared with either ARG or BSC).

**ARG stimulated enterocyte migration is blocked by ODC inhibition**

ARG is a precursor of ornithine via the action of arginase(s).26–27 Ornithine derived polyamines (putrescine (PUT), spermidine, and spermine) are essential for intestinal cell migration.28–30 We postulated that inhibition of ODC, the rate controlling enzyme in polyamine biosynthesis, would abolish stimulation of migration by ARG. The ODC inhibitor DFMO was used at 5 mM, a concentration that was previously shown to maximally inhibit ODC in the IPEC-J2 cell line.15 DFMO by itself did not inhibit basal levels of migration (113 (6) µm/mm v 106 (6) µm/mm in control cells). Failure of DFMO to fully block migration is consistent with the observation that after four days of DFMO, cellular spermine levels remained 50% of normal.15 Figure 6A shows that ARG stimulated migration was blocked by DFMO, and that PUT rescued this DFMO effect.

High pressure liquid chromatography was used to determine if ARG treatment for 24 hours enhanced IPEC-J2 cell polyamine levels. Figure 6B shows that addition of ARG did not enhance polyamine levels. Serum induces ODC in intestinal cells,16 BSC 100 mg/100 ml produced threefold increases in the levels of PUT, spermidine, and spermine (p < 0.05). These levels were higher compared with those of freshly isolated piglet enterocytes31 but were very similar to those previously reported for cultured Cdx2 transformed IEC-6 cells.15

In our additional studies, high dose PUT (0.1 mM) alone did not significantly stimulate IPEC-J2 cell migration (data not shown, n = 3). These results suggest that despite a constitutive role of polyamines, the efficacy of ARG cannot be attributed to enhanced polyamine synthesis.
Requirement for FAK in ARG stimulated enterocyte migration: transfection studies
To determine if FAK phosphorylation is required for intestinal cell migration and the response to ARG, Cdx-2 transformed IEC-6 cells were transfected with a dominant negative construct of FAK mediated by adenoviral gene transfer.24 Adenovirus promoter linked to luciferase was used as a control for cells with intact FAK activity. Preliminary studies showed that control cell infection for 12 hours led to a steady rise in luciferase expression (n = 3). Therefore, transfections were performed for 12 hours. FAK-CD transfection reduced basal cellular migration by 33% compared with control virus and also fully inhibited the cell migration response to ARG and BSC. In contrast, control transfection with adeno-luciferase had no inhibitory effect on the ARG response, with cells migrating at the control rate after treatment with either ARG or BSC (p ≤0.05) (fig 7).

DISCUSSION
Nitric oxide in cell migration
One of the most important metabolites of arginine is NO which plays an important role in regulating intestinal barrier function.13 In newborn pigs, pretreatment with intravenous ARG greatly attenuated necrotising enterocolitis (NEC) produced by intraluminal acidified casein31 and improved intestinal recovery after ischaemic damage in rats by an NOS dependent mechanism.9 10 Our finding that NO stimulates post-injury intestinal epithelial cell migration is consistent with several studies in other cell types. ARG, NO donors, and NO second messengers enhanced endothelial cell migration.11 12 NO was reported to inhibit cell adhesion to matrix (and activation of FAK) in renal mesangial cells, vascular smooth muscle cells, and endothelial cells.13 15 21 In normal renal epithelial monolayers, after wounding there is a biphasic release of NO during cell migration with peaks at 1–2 minutes and again after 2–3 hours. This NO release is accompanied by iNOS expression at the wound edge.11 32 It has been suggested that increased NO facilitates detachment and scalar movement, as opposed to vectorial movement.32

FAK as a signalling molecule in intestinal cell migration
Previous studies demonstrated a central role for FAK in cell migration in mesenchymal cells, such as human umbilical vein endothelial cells. Tyrosine phosphorylation of FAK correlated well with FAK activation and motility while a tyrosine kinase inhibitor blocked migration and FAK activation.3 31 Overexpression of FAK increased cell migration.32 In research focusing on intestinal cells, Caco-2 cell adhesion to laminin or type 1 collagen was shown to require tyrosine phosphorylation.3 In research focusing on intestinal cells, Caco-2 cell adhesion to laminin or type 1 collagen was shown to require tyrosine phosphorylation.3

Our studies support a primary role for FAK in ARG stimulated small intestinal cell migration. ARG increased...
Focusing on the key points:

- ALA (4 mM) stimulation was blocked by DFMO (1 mM) was added 24 hours before the treatment of ARG
- Relationship between polyamine synthesis and cell migration
- Intracellular polyamine concentration was determined 24 hours after wounding
- Polyamines (PUT, spermidine, and spermine) are central to the restitution process in intestinal tissues
- Polyamines are constitutively required for the stimulatory effect of ARG. However, polyamines can stimulate cell matrix adhesion, as proposed in fig 8. Santos et al showed that polyamine deficiency inhibited cell attachment to plastic, laminin, fibronectin, collagen IV, and Matrigel by different extents. Polyamines were essential for normal expression of the integrin subunit α2 but not for expression of the α1 subunit. Thus polyamines participate in cell attachment and expression of the integrin α2β1, a putative receptor for collagen and laminin.
- We have shown in this study that increasing ARG levels above the basal medium concentration of 0.4 mM did not lead to increased polyamine synthesis.
increase polyamine synthesis, consistent with studies in cultured smooth muscle cells and endothelial cells.65–67 These findings may indicate that arginase rather than arginine levels limit the production of polyamines in cultured intestinal cells, as recently reported for cultured endothelial cells.68

Clinical implications of ARG in paediatric intestinal disease

ARG metabolism is of major importance to neonatal nutrition.49 ARG serum levels dwindle below normal levels in preterm infants that later develop NEC, and these levels are in the range of 0.04–0.06 mM, well below postprandial luminal levels (see above) and below the maximal levels to stimulate intestinal cell migration shown in these studies.65–67 A recent prospective controlled trial of ARG versus non-essential amino acid therapy for preterm infants showed that ARG supplementation significantly reduced the incidence of NEC. Our investigations have identified the cellular mechanisms of a restitution enhancing effect of ARG. Our studies show that the magnitude of the stimulated cell migration equals that of growth factors and prostaglandins. Further defining the role of ARG in restitution may provide a rationale for administering exogenous free ARG and BSC (or components of BSC) in treatment solutions for patients with intestinal injury.

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