Targeting focal adhesion assembly by ethoxyfagaronine prevents lymphoblastic cell adhesion to fibronectin


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Abstract. Background: Leukemic cell adhesion to proteins of the bone marrow microenvironment provides signals which control morphology, motility and cell survival. We described herein the ability of ethoxyfagaronine (etxfag), a soluble synthetic derivative of fagaronine, to prevent leukemic cell adhesion to fibronectin peptide (FN/V).

Methods: Phosphorylation of fak and pyk2 were evaluated by immunoblotting. Labelled proteins were localized by confocal microscopy. PI 3-kinase activity was evaluated by in vitro kinase assay.

Results: Subtoxic concentration of etxfag reduced L1210 cell adhesion to FN/V dependently of β1 integrin engagement. Etxfag impaired FN-dependent formation of β1 clustering without modifying β1 expression at the cell membrane. This was accompanied by a decrease of focal adhesion number, a diminution of fak and pyk2 phosphorylation at Tyr-576, Tyr-861 and Tyr-579, respectively leading to their dissociations from β1 integrin and inhibition of PI 3-kinase activity. Etxfag also induced a cell retraction accompanied by a redistribution of phosphorylated fak and pyk2 in the perinuclear region and lipid raft relocalization.

Conclusion: Through its anti-adhesive potential, etxfag, combined with conventional cytotoxic drugs could be potentially designed as a new anti-leukemic drug.

Keywords: Adhesion, ethoxyfagaronine, integrin, focal adhesion kinasas, acute leukemia

1. Introduction

Leukemias are defined as uncontrolled proliferation and enhanced invasiveness of hematopoietic cells that fail to differentiate into mature blood cells. Acute lymphoblastic leukemia (ALL) originates from a single lymphoid progenitor and is characterized by an excessive egress of leukemic blasts from bone marrow (BM) to peripheral blood. Subsequent leukemic cell invasion of extramedullary sites require cell adhesion to extracellular matrix (ECM) proteins, ECM proteolysis, basement membrane crossing and migration into the surrounding tissues [1]. Leukemic cell dissemination can be efficiently reduced by blocking matrix metalloproteinases (MMPs), endopeptidases with proteolytic activity for a large range of ECM components [2–4]. As adhesive properties of leukemic cells are responsible for leukostasis complication, the disruption of adhesion pathways might also be of therapeutic benefit [5]. Hematopoietic cell interactions with BM proteins are largely mediated by integrin adhesion receptors that consist of α and β integrin subunits [6]. Among
these receptors, α4β1 and α5β1 constitute receptors for fibronectin (FN), extensively found within the hematopoietic microenvironment [7]. FN protein levels lead to proteolytic fragments containing a RGD sequence recognized by α5β1 [8]. Another peptide FN-CH/V corresponding to the sequence WQPPRARI is a chemotactic fragment of FN[V] that promotes adhesion, spreading, and cell migration by direct interaction with α4β1 [9, 10]. High levels of α4β1 are expressed in normal and leukemic progenitors and α4β1 integrin is a modulator which alters migration and BM localization [11].

Integrin-mediated cell adhesion signaling from outside to inside the cell is mainly coordinated by the focal adhesion kinase (fak) and the proline-rich tyrosine kinase (pyk2). Once activated by phosphorylation, these non-receptor tyrosine kinases serve as docking sites for SH2 domain containing proteins such as the p85 subunit of the phosphatidylinositol 3-kinase (PI 3-K), Src family kinases and p120 Ras GAP [12, 13]. In acute myeloid leukemia, fak expression and activity are linked with enhanced blast migration, malignant or metastatic disease and poor prognosis in patients [14]. Pyk2 is highly expressed in hematopoietic and neuronal cells and its overexpression is often correlated with increased metastasis [15–18].

Despite a good cure rate of childhood ALL, many patients still develop serious acute and late complications mainly due to cancer treatment side effects. Moreover, the survival rate of ALL adults remains below 40% in most studies and treatment outcome is poor among patients who relapse on current front-line ALL regimens. New treatments are still requested to improve the cure rate and the quality of life of ALL patients. Targeting leukemic cell interactions with the surrounding microenvironment and more specifically focal adhesion assembly could represent an innovative and original approach to prevent leukemic cell dissemination. Natural substances and their designed derivatives represent powerful tools in the engineering of new anticancer drugs [19]. Ethoxyfagaronine (etxfag), a semi-synthetic derivative of fagaronine, exhibits a high DNA binding affinity but a low cytotoxicity in vitro [20, 21]. We recently reported that etxfag reduces L1210 lymphoblastic cell invasiveness through a down-regulation of plasminogen activators and MT1-MMP expression and activation [22]. We now investigate etxfag effects on L1210 cell adhesion. We describe the ability of etxfag to inhibit cell adhesion to FN/V by disrupting lipid raft organization and preventing β1 integrin clustering. This is accompanied by subsequent attenuation of fak and pyk2 phosphorylation and a decrease of PI 3-kinase activity. All together our results show that etxfag exhibits potent anti-leukemic activity and emphasize its clinical potential in combination with conventional cytotoxic drugs.

2. Materials and methods

2.1. Reagents

Ethoxyfagaronine (N-methyl-12-ethoxy-2-hydroxy-3, 8, 9-trimethoxybenzo[c]phenanthridinium chloride) was provided by Pr Olivier DUVAL (SONAS, Angers). Stock solution of ethoxyfagaronine (etxfag) was prepared at 10^{-2} M in water and stored at 4°C. All reagents for cell culture were from Gibco BRL (Invitrogen, CergyPontoise, France). The anti-mouse β1 integrin (Sc-6622; Sc-8978), anti-fak (Sc-557), anti-pyk2 (Sc-1514 and anti-β-actin (Sc-1616) and anti-vinculin (Sc-5573) antibodies were obtained from Santa Cruz Biotechnology Inc (Tebu, Le Perray en Yvelines, France). Anti-phosphotyrosine-576-fak was from USB biological (Euromedx, Souffelweyersheim, France), anti-phosphotyrosine-861-fak (#2153-1) was from Epitomics (Euromedx, Souffelweyersheim, France). Anti-phosphotyrosine-579-pyk2 was from Biosource Invitrogen ( Fisher Scientific, Illkirch, France). Peroxidase-conjugated anti-rabbit antibody (#7074) was from Cell Signaling Technology Inc. (distributed by Ozyme, Saint-Quentin-en Yvelines Cedex, France). LY303511 was from Calbiochem (distributed by WVR International, Straussbourg, France). LY294002 was purchased from Cell Signaling Technology Inc. (distributed by Ozyme, Saint-Quentin en Yvelines, France). Alexafluor 568 phalloidin, Alexafluor 488, Alexafluor 488 conjugated cholera toxin subunit B (CTxB) and Pro long mounting medium with DAPI were from Ozyme, Saint-Quentin en Yvelines, France). Extract-all was from Eurobiol (Eurobio, Les Ulis, France). Verso SYBR Green 2-Step QRT-PCR Rox kit and Absolute SYBR Green Rox mix were from Thermo Electron (Thermo Electron, Courtaboeuf,
France). The prestained protein ladder (#SM0671) used for zymography was from Fermentas (Fermentas, St Rémy Les Chevreuse, France). Anti-peroxidase-conjugated anti-goat antibody, methyl-β-cyclodextrin (MBCD), fibronectin peptide (F-3667), Phosphate Buffered Saline and all others reagents were from Sigma (Saint Quentin Fallavier, France). The protein-G-agarose beads were from Boehringer (Boehringer Mannheim, Indianapolis, IN, USA).

2.2. Cell treatment

To study the effect of etxfag on cell adhesion, $1 \times 10^{-7}$ L1210 cells were incubated for 24 h at 37°C in RPMI-1640-Glutamax containing 10% (v/v) FBS and $1 \times 10^{-7}$ M etxfag, then washed twice with serum free RPMI-1640-Glutamax. To study the effect of Mg$^{2+}$ and EDTA on adhesion, L1210 cells were incubated for 30 min under gentle stirring in serum-free RPMI-1640-Glutamax supplemented with 5 mM Mg$^{2+}$ or 5 mM EDTA and maintained in this medium throughout the experiment. Aliquots ($25 \times 10^4$ cells) were plated on 24-well plates for adhesion assays.

2.3. Adhesion assay

24-well tissue culture plates were coated with 200 $\mu$L FN/V (50 $\mu$g/mL PBS) overnight at 4°C blocked with 1% (w/v) heat-denatured BSA for 1 h at 37°C and washed once with PBS. $25 \times 10^5$ cells/800 $\mu$L RPMI 1640-Glutamax were added in each well and incubated at 37°C from 0 to 2 h. After the incubation period, non-adherent cells were discarded. Adherent cells were rinsed thrice with heated medium (37°C), fixed for 10 min with 4% (w/v) paraformaldehyde in PBS, stained for 10 min with 0.5% (w/v) violet crystal (20/80: methanol/water), washed thrice with PBS. Violet crystal was solubilized with 100 $\mu$L DMSO and absorbance was recorded at 600 nm and was strictly correlated to the cell number [23].

2.4. Western blotting

$35 \times 10^5$ cells/mL were plated on tissue culture plates (25 cm$^2$) coated with FN/V (50 $\mu$g/mL) or BSA 1% (w/v) and incubated for 2 h at 37°C. The non-adherent cells were discarded and the plates rinsed thrice with ice-cold PBS containing 50 $\mu$M Na$_2$VO$_4$. The adherent cells were lysed in 200 $\mu$L lysis buffer and cell lysates were analysed as described [22].

2.5. Immunoprecipitation

Whole cell lysates were precleared with 30 $\mu$L protein-G-agarose for 1 h at 4°C on a rotating platform. The supernatant was then transferred in a tube containing 2 $\mu$g polyclonal anti-CD29 antibodies and rotated overnight at 4°C. 35 $\mu$L protein-G-agarose beads were added to immune complexes and gently rotated for 1 h. Beads were then washed thrice with PBS containing 1% (v/v) NP-40 and bound material was recovered by boiling in Laemmli buffer for 5 min.

2.6. PI 3-kinase assay

Following etxfag incubation and adhesion to FN/V, cells were lysed as described above. The proteins were immunoprecipitated with 2 $\mu$g anti-p85 antibodies and PI 3-kinase assays were performed as described previously [24]. Quantification of PI phosphorylation detected by autoradiography was performed by densitometry and subsequent analysis using the program Image Quant (Molecular Dynamics).

2.7. Cell surface protein biotinylation

Cells were washed with ice-cold PBS containing 20 mM Na$_2$VO$_4$ and incubated with 0.5 mg/mL EZ-Link™ Sulfo-NHS-LC-Biotin, a membrane impermeable biotinylation reagent, in PBS at 4°C for 30 min. Cell surface proteins were separated as described previously and analysed by SDS-PAGE followed by western blotting [22].

2.8. RNA isolation, RT-PCR and real-time PCR

Cells (35 $\times 10^5$ cells/mL) were plated on tissue culture plates coated with FN/V (50 $\mu$g/mL) and incubated for 2 h at 37°C. Non adherent cells were discarded and the plates rinsed thrice with PBS. Adherent cells were scrapped in the same buffer and centrifuged (800 g, 5 min, 4°C). Total RNAs were extracted using TRI Reagent. 500 nanograms were subjected to reverse transcription using Verso transcriptase (Thermo Electron, Courtaboeuf, France). Primers for mouse β1, β2 and β3 integrin subunits and β actin, RPS26 were synthesized by Eurogentec (France) and their sequences
Table 1

<table>
<thead>
<tr>
<th>Genes</th>
<th>Primer sequence forward 5′−3′</th>
<th>Primer sequence reverse 5′−3′</th>
</tr>
</thead>
<tbody>
<tr>
<td>β1 integrin</td>
<td>CAA-TTG-TAG-CAG-GCG-TGG-TTG</td>
<td>GCC-CAC-ACT-GTG-CAC-GGC-ACT</td>
</tr>
<tr>
<td>β2 integrin</td>
<td>AGC-CCA-TCT-TTG-CGG-CTG-C</td>
<td>TOC-ACC-AGC-CTG-CTG-GAG-T</td>
</tr>
<tr>
<td>β3 integrin</td>
<td>AAG-AAC-GAG-GTT-ATC-CCG-GGG</td>
<td>CGG-TGA-GCC-TGT-CCT-TAA-AG</td>
</tr>
<tr>
<td>β-Actin</td>
<td>CCC-TAA-GGC-CAA-CCG-TGA-A</td>
<td>ACA-CTG-TGG-CTG-ACC-CGC-G</td>
</tr>
<tr>
<td>RPS26</td>
<td>TAG-AAG-CCG-CTG-CTG-TCA-GG</td>
<td>GGC-ACA-GCT-CAC-GCA-ATA-AT</td>
</tr>
</tbody>
</table>

were presented in Table 1. Numbers of cycles were adjusted to ensure that amplifications were in a linear range. Real-time PCR was performed as described previously [22] to assess expression of β1 integrin genes. The entire experiment from RNA extraction to real-time PCR steps was repeated four times with different sets of samples.

2.9. Cellular extraction of total lipids and GM1 titration

Cell pellets obtained from L1210 cells (400 g, 10 min, 4°C) treated in the presence or in the absence of 1 × 10⁻⁷ M etxfag for 24 h were suspended in 600 µL of distilled water. Protein concentrations were determined by Bradford assay (Bio-Rad, Marne la Vallée, France). Equal amounts of proteins were mixed with 2.4 mL of a chloroform/methanol (2 : 1 (v/v)) mixture and centrifuged (5000 g, 10 min, 4°C). Aqueous and organic phases were collected and evaporated under nitrogen and mixed with 300 µL of chloroform/methanol (1 : 5 (v/v)). A nitrocellulose membrane was then placed in a 96-well plate system using Bio-dot microfiltration system (Bio-rad). Each well was incubated with 100 µL of aqueous or organic phases from lipid extraction during 30 min. Wells were washed twice with 100 µL TBS pH 8 and saturated with 50 µL BSA 6 mg/mL during 1 h. After 2 washes with 100 µL TBS pH 8, wells were incubated with 50 µL cholera toxin B subunit coupled Alexa Fluor 488 (dilution 1/1000 in TBS pH 8) during 45 min. After 3 washes with 100 µL TBS pH 8, the nitrocellulose membrane was revealed with a microplate spectrofluorimeter Infinitt® F200 Pro (TECAN, Lyon, France).

2.10. Immunofluorescence

Cells (25 × 10⁴) were plated on glass-slides, incubated for 2 h à 37°C and fixed with 4% (v/v) paraformaldehyde for 5–10 min at 4°C. The slides were washed with PBS and saturated in PBS with 2% (v/v) BSA for 30 min. Cells were then incubated overnight at 4°C with anti β1 integrin, anti-fak or anti-vinculin antibody and with antibody against phosphorylated-Y576-fak, Y861-fak, Y579-pyk2. Slides were washed five times in PBS and incubated for 1 h with Alexafluor 488-conjugated secondary antibody diluted 1/1000 in PBS with 2% (v/v) BSA. To visualize F-actin, slides were incubated for 45 min with Alexafluor-568-conjugated phalloidin diluted 1/100 in PBS with 2% (v/v) BSA. Cells were then washed with PBS. Conditions without the first antibody were used as control. Nuclei were counterstained with DAPI. Immunofluorescence-labeled cell preparations were analysed using a Zeiss LSM 710 confocal laser scanning microscope with the 63X oil-immersion objective zoom 3X and Zeiss operating system (Carl Zeiss MicroImaging GmbH, Deutschland). Acquisitions were performed by exciting Alexafluor 488, Alexafluor 568 and DAPI dye with Argon laser, HeNe laser and chameleon IR laser tuned at 730 nm respectively. Emitted fluorescence was detected through the appropriate wavelength window. Twenty images were captured with a 0.25 µm z-step.

2.11. Images post-processing

Surfacic representations were realized using AMIRA (v5.2) software (Visage Imagin®, Berlin, Germany). To compare cell labeling, a same threshold was applied for each channel of untreated versus etxfag-treated cells. For colocalization studies, a multichannel field module was used, followed by a correlation plot treatment (sub-ranges values 15–255, gamma 0.5). A surfacic representation of the correlation plot was realized using the same threshold for untreated versus etxfag-treated cells.

2.12. Statistical analysis

Values are means ± SEM of triplicate determinations or means ± SD of duplicate determinations. The
statistical significance of differences was calculated using Student’s test. *P values referring to corresponding controls are indicated in Figure legends. **P < 0.01; ***P < 0.001; NS = non significant.

3. Results

3.1. Ethoxyfagaronine prevented L1210 cell adhesion to fibronectin via alteration of β1 integrin clustering

As cell adhesion to BM proteins regulates leukemic cell properties, we assessed adhesion of L1210 cells to fibronectin-derived peptide FN/V, described for its ability to strongly induce hematopoietic cell adhesion through interaction with VLA-4 receptor [9]. L1210 cell line is usually used in comparative in vitro tests to evaluate the anti-leukemic potential of new drugs. FN/V peptide significantly enhanced in a time-dependent manner L1210 cell adhesion as compared to BSA controls (Fig. 1a). L1210 cell adhesion to FN/V was reduced with 5 mM EDTA and enhanced with 5 mM Mg²⁺ suggesting that divalent cations could be involved in L1210 cell adhesion to FN/V (Fig. 1b). Integrins constitute one of the major classes of adhesion molecules and particularly, β1 integrin is one of the cation divalent-dependent FN/V receptors [9]. L1210 cells mostly express β1 subunit (RT-PCR and western blot analyses; Fig. 1c) and the anti-β1 blocking antibody (10 μg/mL) significantly reduced (65%) their ability to adhere to FN/V (Fig. 1d). Ethoxyfagaronine (etxfag) used at a subtoxic concentration of 1 × 10⁻⁷ M decreased the number of adhered cells to FN/V but had no effect on cell adhesion to BSA (Fig. 2a). Surprisingly, etxfag affected neither expression of β1 mRNA (qRT-PCR analyses, Fig. 2b) nor β1 protein expression in L1210 cells (western blot, Fig. 2e). To explain these results, we hypothesized that etxfag could alter β1 integrin localization at the cell membrane with consequences on β1 clustering and signaling. To address this point, cell surface proteins were biotinylated with sulfo-NHS-biotin and treated with etxfag. Western blot analyses of biotinylated proteins showed that etxfag induced cell rounding and retraction from the substratum (Figs. 3 and 4, right row). This spreading was characterized by well-organized cortex fibers, membrane ruffles, leading edge lamellipodia (Fig. 3 left row middle panel) and β1 integrin clustering at the cell periphery (Figs. 3 and 4, left row, upper panel, white arrows). Moreover, a strong colocalization of β1 integrin clusters and F-actin indicated developed cytoskeleton structures (Fig. 3, left row, bottom panel, white arrows). Imaging of vinculin (a cytoskeletal protein involved in the linkage of integrin to actin) by confocal microscopy showed localization in the extended cytoskeleton structures (Fig. 4, left row, middle panel). Moreover, β1 and fak co-localization demonstrated numerous sites corresponding to focal adhesion complexes (Fig. 4, left row bottom panel). Differential interventional contrast microscopy and confocal fluorescent imaging showed that etxfag induced cell rounding and retraction from the substratum (Figs. 3 and 4, right row). Etxfag also inhibited the FN/V-dependent β1 integrin clustering (Figs. 3 and 4, right row, upper panel) and prevented F-actin network extensions, membrane ruffles and lamellipodia (Fig. 3 right row, middle panel). Colocalization experiments of F-actin with β1 integrin supported these results (Fig. 3, right row, bottom panel) and demonstrated a diffuse distribution of β1 integrin and F-actin. To go deeper in etxfag molecular mechanisms, localization of vinculin was evaluated and appeared at the perinuclear region (Fig. 4, right row, middle panel). These results are also reinforced by the estimation of β1 and vinculin co-localization (Fig. 4, left row, bottom panel) and show clearly that etxfag induced a decreasing of these focal adhesion complexes. Interestingly, the results obtained with fak and vinculin others partners of focal adhesion complexes are identical to those obtained with β1 and vinculin (supplementary Figure 1).

3.2. Ethoxyfagaronine prevented phosphorylated fak and pyk2 association with β1 integrin

We then focused on downstream signaling of β1 integrin, particularly on fak and pyk2 kinases in FN-adhered L1210 cells, fak was phosphorylated on Tyr-576 (P(576)fak) (supplementary Figure 2a) and Tyr-861 (P(861)fak) (Fig. 5a) and associated with β1 integrin and actin distribution showed that L1210 cells treated with etxfag. Differential interventional contrast microscopy and confocal fluorescent imaging followed by surface reconstruction post-processing of β1 integrin and actin distribution showed that L1210 cells spread when adhered to FN/V (Fig. 3 left row and Fig. 4 left row). This spreading was characterized by well-organized cortex fibers, membrane ruffles, leading edge lamellipodia (Fig. 3 left row middle panel) and β1 integrin clustering at the cell periphery (Figs. 3 and 4, left row, upper panel, white arrows). Moreover, a strong colocalization of β1 integrin clusters and F-actin indicated developed cytoskeleton structures (Fig. 3, left row, bottom panel, white arrows). Imaging of vinculin (a cytoskeletal protein involved in the linkage of integrin to actin) by confocal microscopy showed localization in the extended cytoskeleton structures (Fig. 4, left row, middle panel). Moreover, β1 and fak co-localization demonstrated numerous sites corresponding to focal adhesion complexes (Fig. 4, left row bottom panel). Differential interventional contrast microscopy and confocal fluorescent imaging showed that etxfag induced cell rounding and retraction from the substratum (Figs. 3 and 4, right row). Etxfag also inhibited the FN/V-dependent β1 integrin clustering (Figs. 3 and 4, right row, upper panel) and prevented F-actin network extensions, membrane ruffles and lamellipodia (Fig. 3 right row, middle panel). Colocalization experiments of F-actin with β1 integrin supported these results (Fig. 3, right row, bottom panel) and demonstrated a diffuse distribution of β1 integrin and F-actin. To go deeper in etxfag molecular mechanisms, localization of vinculin was evaluated and appeared at the perinuclear region (Fig. 4, right row, middle panel). These results are also reinforced by the estimation of β1 and vinculin co-localization (Fig. 4, left row, bottom panel) and show clearly that etxfag induced a decreasing of these focal adhesion complexes. Interestingly, the results obtained with fak and vinculin others partners of focal adhesion complexes are identical to those obtained with β1 and vinculin (supplementary Figure 1).
integrin (Fig. 3b and S2b). Moreover, phosphorylated fak was localized at the polarized-cell periphery (Fig. 5c, left row, upper and middle panel). Colocalization studies between P(861) fak and F-actin and between P(576) fak and F-actin showed numerous clusters at the periphery of the spread cells (Fig. 5c and S2c, left row, bottom panel, white arrows). Extfag did not alter fak expression (Fig. 5a and S2a) but attenuated by 40% fak phosphorylation on Tyr-576 (S2a) and Tyr-861 (Fig. 5a). β1 integrin immunoprecipitation showed that extfag did not alter fak association with β1 but decreased the level of phosphorylated fak associated with β1, confirming the activity of extfag on fak phosphorylation only (supplementary Figure 2h and Fig. 5b). Confocal experiments revealed that extfag-induced cell retraction characterized by a reduced actin cortex and a decrease of focal adhesions (Fig. 5c and S2c, right row, upper and middle panels). Along this line, colocalization experiments demonstrated a diffuse repartition of P(861)fak and F-actin (Fig. 5c and S2c, right row, bottom panel) in L1210 cells treated with extfag.
Fig. 2. Ethoxyfagaronine (Etxfag) inhibits L1210 cell adhesion to FN/V peptide. a) $25 \times 10^5$ L1210 cells treated or not with $1 \times 10^{-7}$ M Etxfag for 24 h were seeded on 1% BSA (○) or 5 μg/cm$^2$ FN/V peptide (■) for 2 h. Cell adhesion was evaluated as described in Fig. 1a. **$P < 0.01$ versus to untreated cells. NS = non significative. b) β1 integrin transcript levels were analysed by qRT-PCR as described in section 2. Error bars represent the SEM for the average fold change of at least three independent replicates. NS = non significative. c) Whole cell extracts obtained from L1210 cells treated or not with $1 \times 10^{-7}$ M Etxfag for 24 h were analysed by 10% SDS-PAGE followed by western blotting using anti-β1 integrin antibody. A representative blot of three independent experiments was shown. d) L1210 cells were biotinylated and treated or not with $1 \times 10^{-7}$ M Etxfag for 24 h. Biotinylated proteins were analysed by 10% SDS-PAGE followed by western blotting using monoclonal anti-β1 antibody. A representative blot of three independent experiments is shown. In FN/V-adhered cells, pyk2 was also phosphorylated and co-immunoprecipitated with β1 (Fig. 6a and b). As for P-fak, P(579)pyk2 localized at the polarized cell periphery (Fig. 6c, left row, upper and middle panels). Colocalization studies between P(579) pyk2 and F-actin showed numerous clusters at the spread cell periphery (Fig. 6c, left row, bottom panel, white arrows). Etxfag reduced by 45% pyk2 phosphorylation on tyr 579 as well as association with β1 integrin (Fig. 6a and b). Moreover, P(579)pyk2 was redistributed in the perinuclear region (Fig. 6c, right row, upper and middle panels) and colocalization evaluation demonstrated a diffuse repartition between P(579)pyk2 and F-actin (Fig. 6c, right row, bottom panel).

3.3. Ethoxyfagaronine decreased PI 3-kinase activity

Since phosphorylated fak leads to the recruitment and subsequent activation of SH2 domain-containing proteins, activity of PI 3-kinase was evaluated in FN/V-adhered cells. Autoradiographic analyses of phosphorlated PI (PIP) showed that PI 3-kinase activity was lowered by 30% upon etxfag treatment (Fig. 7a). PI 3-kinase activity was partially blocked by 20 μM LY294002, a highly specific inhibitor but not by 25 μM LY303511, the cell-permeable negative control for LY294002 [25]. Moreover, 20 μM LY294002 significantly reduced L1210 cell adhesion (Fig. 7b) confirming that L1210 cell adhesion to FN/V involved PI 3-kinase activity.

3.4. Ethoxyfagaronine relocalized lipid rafts

Integrin-mediated signaling is dynamically modulated by lipid rafts. Membrane microdomains enriched in cholesterol and glycosphingolipid such as ganglioside GM1. To investigate lipid raft involvement in L1210 cell adhesion to FN/V, cells were labelled with FITC-coupled cholera toxin subunit that binds GM1 and visualized by confocal microscopy. In FN/V-adhered L1210 cells, lipid rafts appeared as large patches (Fig. 8a, left row, upper panel). Methyl-β-cyclodextrin (MβCD) that sequesters cholesterol from cell membranes, induced lipid raft disruption (Fig. 8a,
Fig. 3. Etxfag alters β1 integrin clustering and actin cytoskeleton expansion in L1210 cells. L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag during 24 h were seeded on FN/V-coated glass culture slides in serum-free medium for 2 h. Cells were fixed with 4% paraformaldehyde then incubated with anti-β1 primary antibody (1:100) and labelled with secondary antibody linked to Alexafluor 488 (green). Cells were also incubated with Alexafluor 568-conjugated phalloidin (red) to visualize F-actin distribution. Emitted fluorescence was detected through the appropriate wavelength window. Twenty images were captured with a 0.25 μm z-step. Surface reconstructions were realized from L1210 cells treated with or without etxfag (right and left row, respectively). Upper panels represent β1 integrin labelling, middle panels represent β1 integrin+F-actin labelling and bottom panels represent colocalization of β1 integrin and F-actin (gold). A representative image of three independent experiments was shown.

right row, upper panel). When used at 10 mM, a non cytotoxic dose for L1210 cells, MβCD decreased by 90% the number of cells adhering to FN/V (Fig. 8a, right row, bottom panel versus left row, bottom panels). As integrin signaling could be linked to lipid raft localization, confocal microscopy experiments were performed and showed that β1 integrin colocalized with GM1 at numerous sites (Fig. 8b, right row, upper panel, white arrows). Interestingly, etxfag induced relocalization of GM1 to the cytoplasm (Fig. 8b, left row, bottom panel versus left upper panel). Analysis of total GM1 cellular content showed that Etxfag did not modulate the level of this lipid raft component. These results reinforced the data presented above suggesting relocalization of lipid rafts without modulation of their content (Fig. 8c). Importantly, despite relocalization of lipid rafts, β1 integrin residency at the plasma membrane was not modified (Fig. 3).
Fig. 4. Etxfag alters the number of focal adhesion complexes in L1210 cells. L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag during 24 h were seeded to FN/V-coated glass culture slides in serum-free medium for 2 h. Cells were fixed with 4% paraformaldehyde then incubated with anti-$\beta$-1 primary antibody (1:1000) and labelled with secondary antibody linked to Alexafluor 488 (green). Cells were also incubated with anti-vinculin primary antibody (1:1000) and labelled with secondary antibody linked to Alexafluor 568 (red). Emitted fluorescence was detected through the appropriate wavelength window. Twenty images were captured with a 0.25 µm z-step. Surface reconstructions were realized from L1210 cells treated with or without etxfag (right and left row, respectively). Differential interferential contrast imaging of L1210 cells treated or not by etxfag was proceeded (right and left row, upper panel). Upper panels represent $\beta$-1 integrin labelling (green), middle panels represent vinculin labelling (red) and bottom panels represent colocalization between $\beta$-1 integrin and vinculin (gold). A representative image of three independent experiments is shown. Scale bar: 10 µm.
Fig. 5. Etxfag prevents fak phosphorylation on Tyr 861.  

a) Whole cell lysates prepared from L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag during 24 h were subjected to immunoblotting with anti-phosphorylated-Y861 fak and fak. A representative blot of three independent experiments was shown. Graph represents the densitometric analysis of the ratio P(861)fak/fak in control (H17040) and treated (H17039) cells. *$P<0.05$.

b) Cell lysates prepared from L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag during 24 h were immunoprecipitated with anti-$\beta$1 antibody and immunoblotted with anti-phosphorylated-Y861-fak and fak. Non specific IgGs were used as negative controls of immunoprecipitation (windows). A representative blot of three independent experiments was shown. Graph represents the densitometric analysis of the ratio P(861)fak/fak in control (H17040) and treated (H17039) cells.

c) L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag for 24 h were seeded to FN/V for 2 h. Cells were fixed with paraformaldehyde 4% and permeabilized with 0.1% Triton X-100. Blue color represents nuclear staining with DAPI. Cells were labelled with anti-phosphorylated-Y861-fak and then with a secondary antibody linked to Alexafluor 488 (green). Cells were also incubated with Alexafluor 568-conjugated phalloidin (red) to visualize F-actin distribution. Surface reconstructions were realized from L1210 cells treated with or without etxfag (right and left row, respectively). Upper panels represent P(861)fak labelling, middle panels represent P(861)fak+F-actin labelling and bottom panels represent colocalization between P(861)fak and F-actin (gold). A representative image of three independent experiments is shown.
Fig. 6. EtXfag prevents pYk2 phosphorylation and association with β1 integrin. a) Whole cell lysates prepared from L1210 cells treated or not with \(1 \times 10^{-7}\) M EtXfag during 24 h were subjected to immunoblotting with anti-phosphorylated-Y579-pyk2 and pyk2. A representative blot of three independent experiments was shown. Graph represents the densitometric analysis of the ratio P(579)pyk2/pyk2 in control (\(1\)) and treated (\(\square\)) cells. *P<0.05. b) Cell lysates prepared from L1210 cells treated or not with \(1 \times 10^{-7}\) M etxfag during 24 h were immunoprecipitated with anti-β1 antibody and immunoblotted with anti-phosphorylated-Y579-pyk2 and pyk2. Non specific IgGs were used as negative controls of immunoprecipitation (windows). A representative blot of three independent experiments was shown. Graph represents the densitometric analysis of the ratio P(579)pyk2/pyk2 in control (\(1\)) and treated (\(\square\)) cells. *P<0.05. c) L1210 cells treated or not with \(1 \times 10^{-7}\) M EtXfag for 24 h were seeded to FN/V for 2 h. Cells were fixed with paraformaldehyde 4% and permeabilized with 0.1% Triton X-100. Cells were labelled with anti-phosphorylated-Y579-pyk2 and then with a secondary antibody linked to Alexafluor 488 (green). Cells were also incubated with Alexafluor 568-conjugated phalloidin (red) to visualize F-actin distribution. Blue color represents nuclear staining with DAPI. Surface reconstructions were realized from L1210 cells treated with (right row) or without etxfag (left row). Upper panels represent P(579)pyk2 labelling, middle panels represent P(579)pyk2+F-actin labelling and bottom panels represent co-localization between P(579)pyk2 and F-actin (gold). A representative image of three independent experiments is shown.
Fig. 7. Etxfag decreased PI-3 kinase activity in FN/V-adhered L1210 cells. a) L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag for 24 h were seeded to FN/V for 2 h in a serum-free medium. Adherent cells were incubated with 25 μM LY294002 or 25 μM LY303511 for 30 min before lysis. Anti-p85 immunoprecipitates were subjected to PI 3-kinase activity determined by PI phosphorylation in the presence of [$\gamma$-32P] ATP. Phosphorylated PI (PIP) was separated by TLC and analysed by autoradiography. Quantification of PI phosphorylation was expressed as the percentage of signal obtained in FN/V-adhered cells. Results are representative of two independent experiments and values are the means ± SD. **P < 0.01 compared with control FN/V-adhered cells. NS = non significative. b) L1210 cells treated or not with either $1 \times 10^{-7}$ M etxfag for 24 h or 25 μM LY294002 or 25 μM LY303511 for 30 min were seeded to FN/V for 2 h. Non-adherent cells were discarded and adherent cells stained by violet crystal were visualized by microscopy.

4. Discussion

Progression of acute leukemias is depicted by excessive egress of leukemic blasts and cell infiltration in multiple organs through a series of processes including cell adhesion and ECM proteolysis. Adhesive interactions with the ECM proteins play a crucial role in the homing of normal hematopoietic cells within the bone marrow but can also facilitate leukemic cell lodgment leading to the persistence of residual disease [26]. These interactions can also protect leukemic cells from chemotheraphy-induced apoptosis [27]. New anti-leukemic therapeutics targeting cell interactions with the BM could be used in combination with conventional cytotoxic drugs to eradicate leukemic cells. Fagarone zanthoxylum L is a benzo[c]-phenanthridine described for its anti-leukemic capacity linked to DNA topoisomerase inhibition [28–30]. Its
Fig. 8. Lipid raft integrity is necessary for L1210 cell adhesion to FN/V peptide.

a) L1210 cells treated or not with 10 mM MβCD for 1 h were seeded to FN/V-coated glass culture slides in serum-free medium for 2 h. Blue color represents nuclear staining with DAPI. L1210 cells were fixed with 4% paraformaldehyde then incubated with Alexa Fluor 488-conjugated CTxB (green). Z-projection (Z = 0.25 μm) were realized by confocal microscopy using adequate settings for each fluorophore. Images were analysed with Amira and are representative of three independent experiments (upper panels).

b) L1210 cells treated or not with 1 × 10⁻⁷ M etxFag during 24 h were seeded onto FN/V-coated glass culture slides in serum-free medium for 2 h. Cells were fixed with 4% paraformaldehyde then incubated with a primary anti-β1 antibody (1:100) and labelled with a secondary antibody linked to Alexafluor 568 (data not shown). Cells were also incubated with Alexafluor 488-conjugated cholera toxin subunit B (CTxB). Blue color represents nuclear staining with DAPI. Surface reconstructions were realized from L1210 cells treated with (bottom panels) or without etxFag (upper panels). Surface reconstructions were proceed from GM1 labelling (left row, upper and bottom panels) and from colocalization between GM1 and β1 (right row, upper and bottom panel) (gold). A representative image of three independent experiments is shown.

c) L1210 cells were treated or not with 1 × 10⁻⁷ M EtxFag during 24 h then subjected to lipid extraction. GM1 ganglioside was used as a marker of lipid raft and its expression was assessed by using dot-blot CTxB staining followed by fluorescence spectroscopy analysis.
high toxicity and insolubility in biological fluids render fagaronine unusable in vivo [31]. In order to improve the anti-leukemic potency while reducing the toxicity of benzo[c]phenanthridines, derivatives of fagaronine like ethoxfagaronine (etxfag) have been synthesized [20, 32]. Etxfag exhibits higher solubility in aqueous solvents and we recently reported that it reduced murine leukemic L1210 cell invasiveness [22].

In order to gain insight into the mechanism of action of etxfag as an anti-leukemic agent, our present study explored the effect of etxfag treatment on cell adhesion to FN, one of the major components of the bone marrow microenvironment. We showed that subtoxic concentration of etxfag modulated integrin-dependent adhesion of L1210 cells to FN and that this effect was associated with an inhibition of integrin clustering leading to a decrease of focal adhesion complexes.

Interaction of leukemic cell with FN on stromal cells controls minimal residual disease and involves integrin [26]. The value of αβ1 as a prognostic marker may not be an efficient strategy [33]. This is reinforced by studies reporting that integrin-dependent adhesion up-regulate MMPs [34]. Integrins act as bidirectional transducers that connect ECM proteins with the cytoskeleton through focal adhesion sites recruiting scaffolding proteins. Within these sites, fak is the key kinase initiating integrin signaling pathway. Fak is aberrantly regulated in acute leukemic cells and its expression correlates with enhanced cell migration and drug resistance in vitro [12]. Pyk2, another focal adhesion tyrosine kinase, can replace fak in integrin-mediated signaling events and its overexpression is often associated with enhanced metastasis [35]. Fak and pyk2 could thereby represent new molecular therapeutic targets for leukemia but these two kinases can be regulated differently in response to the same stimulus while sharing some functions [36, 37]. The ability of pyk2 to compensate fak deficiency suggests that targeting pyk2 might be a better therapeutic approach in leukemia.

Our data indicate that fak and pyk2 are phosphorylated on tyrosine 576, 861 and 579 and associated with β1 integrin in FN-attached L1210 cells. Interestingly, phosphorylated fak and pyk2 were localized at the cell periphery as well as in actin-rich extensions of L1210 cells thus confirming their role in cell spreading and migration. Etxfag affected all these phosphorylation sites as well as pyk2 recruitment to β1 integrin. In the focal adhesion complex, pyk2 phosphorylation induces PI 3-kinase recruitment and its subsequent activation. Activated PI 3-kinase phosphorylates pyk2 which binds to β1 integrin cytoplasmatic tail [35]. Our data show that PI 3-kinase activity was enhanced in FN-adhered L1210 cells and partially reduced by etxfag. The fact that etxfag inhibits PI 3-kinase might thus explain prevention of pyk2 recruitment to β1 integrin and points the inhibitory activity of etxfag on β1 integrin signaling cascade.

Integrin signaling is controlled by several mechanisms including receptor clustering. Large lipid rafts are necessary to create a new microenvironment where focal adhesion proteins are phosphorylated thus modulating downstream signaling and integrin partitioning in lipid rafts [38]. In FN-attached L1210 cells, lipid rafts appeared as large patches and co-localized with β1 integrin. Cholesterol depletion impaired L1210 cell adhesion indicating that lipid raft integrity was required for this process. Treatment with etxfag induced a relocalization of lipid rafts without impacting the cellular distribution of β1 integrin. Modification of the membrane microenvironment could explain the loss of β1 integrin adherence properties. As a consequence, the inhibition of lipid raft clustering by etxfag might constitute the major event leading to the suppression of integrin clustering, focal adhesion contact formation and ultimately cell adhesion to FN.

In conclusion, our results depicted that etxfag displays a significant inhibitory effect on L1210 cell adhesion and spreading through the inhibition of β1 integrin signaling cascade. The observed effects have been also confirmed in two human acute leukemia cell lines (supplementary Figure 3), SHI-1 (established...
from mononuclear cells isolated from the bone marrow of a patient with AML at relapse) and NALM-6 (established from B cell precursors from the peripheral blood of a patient with ALL at relapse). By targeting several processes responsible for leukemic cell development i.e. cell dissemination and adhesion to bone marrow proteins, etxfag appears as a new potential anti-leukemic agent and further preclinical studies may provide the basis for the development of more effective anti-leukemic treatments.

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Supplementary Figure 2. Etxfag prevents fak phosphorylation on Tyr 576.
a) Whole cell lysates prepared from L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag during 24 h were subjected to immunoblotting with antibodies anti-phosphorylated-Y576 fak and fak. A representative blot of three independent experiments is shown. Graph represents the densitometric analysis of the ratio P(576)fak/fak in control (•) and treated (○) cells. *P < 0.05.
b) Cell lysates prepared from L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag during 24 h were immunoprecipitated with anti-β integrin antibody and immunoblotted with antibodies anti-phosphorylated-Y576 fak and fak. Non-specific IgGs were used as negative controls of immunoprecipitation (windows). A representative blot of three independent experiments was shown. Graph represents the densitometric analysis of the ratio P(576)fak/fak in control (•) and treated (○) cells. *P < 0.05.
c) L1210 cells treated or not with $1 \times 10^{-7}$ M etxfag for 24 h were seeded to FN/V for 2 h. Cells were fixed with paraformaldehyde 4% and permeabilized with 0.1% Triton X-100. Blue color represents nuclear staining with DAPI. Cells were labelled with antibodies anti-phosphorylated-Y576 fak and then with a secondary antibody linked to Alexafluor 488 (green). Cells were also incubated with Alexaflour 568-conjugated phalloidin (red) to visualize F-actin distribution. Surface reconstructions were realized from L1210 cells treated with etxfag (left row) or without etxfag (right row). Upper panels represent P(576)fak labelling, middle panels represent P(576)fak+F-actin labelling and bottom panels represent colocalization between P(576)fak and F-actin (gold). A representative image of three independent experiments is shown.
Supplementary Figure 3. Etxfag inhibits Nalm-6 and SHI-1 cell adhesion to FN/V peptide. 

(a) $2.5 \times 10^5$ Nalm-6 cells treated or not with $1 \times 10^{-7} \text{M etxfag}$ for 24 h were seeded on FN/V peptide ($5 \mu\text{g/cm}^2$) for 2 h in serum free condition. Non-adherent cells were then discarded and adherent cells stained by violet crystal were visualized by microscopy. 

(b) $2.5 \times 10^5$ SHI-1 cells treated or not with $1 \times 10^{-7} \text{M etxfag}$ for 24 h were seeded on FN/V peptide ($5 \mu\text{g/cm}^2$) for 2 h in serum free condition. Non-adherent cells were then discarded and adherent cells stained by violet crystal were visualized by microscopy. (Colours are visible in the online version of the article; http://dx.doi.org/10.3233/ACP-2012-0055)

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