Amygdalin Influences Bladder Cancer Cell Adhesion and Invasion In Vitro

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Abstract

The cyanogenic diglucoside amygdalin, derived from Rosaceae kernels, is employed by many patients as an alternative anti-cancer treatment. However, whether amygdalin indeed acts as an anti-tumor agent is not clear. Metastasis blocking properties of amygdalin on bladder cancer cell lines was, therefore, investigated. Amygdalin (10 mg/ml) was applied to UMUC-3, TCCSUP or RT112 bladder cancer cells for 24 h or for 2 weeks. Tumor cell adhesion to vascular endothelium or to immobilized collagen as well as tumor cell migration was examined. Effects of drug treatment on integrin α and β subtypes, on integrin-linked kinase (ILK) and total and activated focal adhesion kinase (FAK) were also determined. Integrin knock-down was carried out to evaluate integrin influence on migration and adhesion. A 24 h or 2 week amygdalin application distinctly reduced tumor cell adhesion and migration of UMUC-3 and RT112 cells. TCCSUP adhesion was also reduced, but migration was elevated under amygdalin. Integrin subtype expression was significantly and specifically altered by amygdalin depending on the cell line. ILK was moderately, and activated FAK strongly, lost in all tumor cell lines in the presence of amygdalin. Knock down of β1 integrin caused a significant decrease in both adhesion and migration of UMUC-3 cells, but a significant increase in TCCSUP adhesion. Knock down of β4 integrin caused a significant decrease in migration of RT112 cells. Since the different actions of amygdalin on the different cell lines was mirrored by β1 or β4 knock down, it is postulated that amygdalin influences adhesion and migratory properties of bladder cancer cells by modulating β1 or β4 integrin expression. The amygdalin induced increase in TCCSUP migratory behavior indicates that any anti-tumor benefits from amygdalin (seen with the other two cell lines) may depend upon the cancer cell type.

Introduction

The use of complementary and alternative medicine (CAM) has steadily increased over the past decades. CAM includes non-conventional therapy such as homeopathy, vitamin therapy, phytotherapy, and traditional Chinese medicine, acupuncture and yoga [1]. The consumption of natural products is most wide spread. Up to 80% of cancer patients in the United States [2], and more than 50% of cancer patients in Europe use CAM together with or in place of conventional therapy [3]. Dissatisfaction with conventional treatment and reduction of chemotherapeutic side effects are the most commonly given reasons for the use of CAM [4,5].  

In contrast to the wide spread use of natural compounds, information about their therapeutic effectivity is sparse. The discrepancy between use and factual benefit is particularly apparent with the cyanogenic diglucoside amygdalin (D-mandelonitrile-β-gentiobioside), present in the kernels of fruits from Rosaceae species such as Prunus persica (peach), Prunus armeniaca (apricot) and Prunus amygdalus amara (bitter almond). Amygdalin was first isolated in 1873. Since the 1920s, amygdalin has been orally applied to treat cancer patients in the United States. In the 1950s, an intravenous form of amygdalin was synthesized and patented as laetrile [6]. Although laetrile is chemically different from amygdalin, the terms are used interchangeably, making interpretation of clinical data difficult. The present report exclusively refers to “amygdalin”.

Amygdalin was one of the most popular, non-conventional, anti-cancer treatments in the 1970s and by 1978, 70,000 US cancer patients had used amygdalin [7]. Still, evidence based research on amygdalin was and is sparse and its benefit controversial. Proponents consider amygdalin a natural cancer cure, whereas opponents warn that amygdalin is ineffective and even toxic. Randomized clinical trials and follow-up studies have
never been carried out. A clinical study sponsored by the National Cancer Institute 30 years ago did not reveal signs of tumor regression [8], whereas a retrospective analysis of 67 tumor patients taking amygdalin reported 2 complete and 4 partial responses [9]. Ambivalence has also been reflected in case reports, where amygdalin was ineffective in five and effective in four cases [6].

The present study was designed to evaluate whether amygdalin alters metastatic tumor cell progression in vitro since invasion and metastasis are critical steps in malignant tumor progression and the main cause of treatment failure. Therefore, interfering with the tumor cell invasion cascade might be an innovative option to counteract metastatic tumor dissemination. Employing a panel of bladder cancer cell lines, the efficacy of amygdalin to block tumor-matrix and tumor endothelial interaction was evaluated. Additionally, the capability of amygdalin to prevent motile spreading was assessed. A cohort of adhesion molecules is involved in the complex process of tumor cell dissemination. Since adhesion receptors of the integrin α and β family are closely involved in tumor cell binding and transendothelial penetration, these were the objects of investigation. The membranous integrin receptor expression profile, as well as the intracellular protein content of each subtype, was compared in amygdalin treated and non-treated cells. siRNA knock down studies were also carried out to investigate those parameters altered by amygdalin, which may have clinical relevance. The in vitro data presented here point to significant adhesion and invasion blocking effects of amygdalin, probably induced by altering β1 or β4 integrin expression.

**Materials and Methods**

**Cell culture**

RT112, UMUC-3 (ATCC/LGC Promochem GmbH, Wesel, Germany) and TCCSUP (DSMZ, Braunschweig, Germany) bladder carcinoma cells were grown and subcultured in RPMI 1640, 10% fetal calf serum (FCS), 20 mM HEPES-buffer, 1% glutamax and 1% penicillin/streptomycin (all: Gibco/Invitrogen; Karlsruhe, Germany). Subcultures from passages 7–24 were selected for experimental use. Human endothelial cells (HUVEC) were isolated from human umbilical veins and harvested by enzymatic treatment with dispase (Gibco/Invitrogen). HUVEC were grown in Medium 199 (M199; Biozol, Munich, Germany), supplemented with 10% FCS, 10% pooled human serum, 20 μg/ml endothelial cell growth factor (Boehringer, Mannheim, Germany), 0.1% heparin, 100 ng/ml gentamycin and 20 mM HEPES-buffer (pH 7.4). Subcultures from passages 2–6 were selected for experimental use. HUVEC were used in the study. The institutional ethics committee of the Goethe-University Hospital, Frankfurt, Germany, approved the investigation and waived the need for consent, since HUVEC were used anonymously for in vitro assays with no link to patient data.

**Amygdalin treatment**

Amygdalin from apricot kernels (Sigma-Aldrich, Taukirchen, Germany) was freshly dissolved in tumor cell culture medium and then added to tumor cells at a concentration of 10 ng/ml for either 24 h or for 2 weeks [10] to evaluate acute versus chronic treatment. Controls received tumor cell culture medium alone. 0.5×10⁶ tumor cells/well were added to HUVEC monolayers for 0.5, 1 and 2 h. Mean adherent tumor cells from five fields was calculated and depicted as percentage of the 100% control (dotted line). One representative of six experiments is shown. *indicates significant difference to controls. doi:10.1371/journal.pone.0110244.g001

**Figure 1. Adhesion of UMUC-3, TCCSUP and RT112 bladder cancer cells to HUVEC.** Tumor cells were treated with 10 ng/ml amygdalin for either 24 h or for 2 weeks. Controls received cell culture medium alone. 0.5×10⁶ tumor cells/well were added to HUVEC monolayers for 0.5, 1 and 2 h. Mean adherent tumor cells from five fields was calculated and depicted as percentage of the 100% control (dotted line). One representative of six experiments is shown. *indicates significant difference to controls.

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**Tumor cell adhesion**

To analyze tumor cell adhesion, HUVEC were transferred to 6-well multiplates (Sarstedt, Nümbrecht, Germany) in complete HUVEC-medium. When confluent, RT112, UMUC-3 or TCCSUP cells were detached from the culture flasks by accutase
treatment (PAA Laboratories, Colbe, Germany) and 0.5 × 10^6 cells were then added to the HUVEC monolayer for 30, 60 or 120 min. Subsequently, non-adherent tumor cells were washed off using warmed (37°C) Medium 199. The remaining cells were fixed with 1% glutaraldehyde. Adherent tumor cells were counted in five different fields of a defined size (3 × 0.25 mm^2) using a phase contrast microscope and the mean cellular adhesion rate was calculated.

Attachment to immobilized collagen

6-well plates were coated with collagen G (extracted from calveskin, consisting of 90% collagen type I and 10% collagen type III; Biochrom, Berlin, Germany; diluted to 400 μg/ml in PBS) overnight. Plastic dishes served as the background control. Plates were washed with 1% BSA (bovine serum albumin) in PBS to block nonspecific cell adhesion. 0.5 × 10^6 tumor cells were then added to each well and left for 60 min incubation. Subsequently, non-adherent tumor cells were washed off, the remaining adherent cells were fixed with 1% glutaraldehyde and counted microscopically. The mean cellular adhesion rate, defined by adherent cells to coated well—adherent cells to background, was calculated from five different observation fields.

Measurement of tumor cell migration

Serum induced chemotactic movement was examined using 6-well Transwell chambers (Greiner, Frickenhausen, Germany) with 8-μm pores. 0.5 × 10^6 RT112, UMUC-3 or TCCSUP cells/ml were placed in the upper chamber in serum-free medium, either free of amygdalin (terminated “amygdalin A”) or containing amygdalin (terminated “amygdalin B”). Serum free medium in the upper chamber and 10% serum in the lower chamber provided the serum gradient necessary for tumor cell migration in this model. After 20 h incubation, the upper surface of the Transwell membrane was gently wiped with a cotton swab to remove cells, which had not migrated. Cells which had moved towards the serum gradient to the lower surface of the membrane were stained using hematoxylin and counted microscopically. The mean migration rate was calculated from five different observation fields.

Integrin surface expression

Tumor cells were washed in blocking solution (PBS, 0.5% BSA) and then incubated for 60 min at 4°C with phycoerythrin (PE)-conjugated monoclonal antibodies directed against the following integrin subtypes: Anti-α1 (IgG1; clone SR84), anti-α2 (IgG2a; clone 12F1–H6), anti-α3 (IgG1; clone C3II.1), anti-α4 (IgG1; clone 9F10), anti-α5 (IgG1; clone GoH3), anti-β1 (IgG1 clone MAR4), anti-β3 (IgG1 clone VI-PL2) or anti-β4 (IgG2a; clone 439–9B; all: BD Pharmingen, Heidelberg, Germany). Integrin expression of tumor cells was then measured using a FACscan (BD Biosciences, Heidelberg; FL-2H (log) channel histogram analysis; 1 × 10^4 cells/scan) and expressed as mean fluorescence units. A mouse IgG1-PE (MOPC-21) or IgG2a-PE (G155–178; all: BD Biosciences) was used as an isotype control.

Western blotting

To investigate integrin content, tumor cell lysates were applied to a 7% polyacrylamide gel and electrophoresed for 90 min at 100 V. The protein was then transferred to nitrocellulose...
membranes. After blocking with non-fat dry milk for 1 h, the membranes were incubated overnight with the monoclonal antibodies listed above. Additionally, integrin-related signaling was explored by anti-integrin-linked kinase (ILK; clone 3, dilution 1:1000), anti-focal adhesion kinase (FAK; clone 77, dilution 1:1000) and anti-phospho-specific FAK (pY397; clone 18, dilution 1:1000) antibodies (all: BD Biosciences). HRP-conjugated goat-anti-mouse IgG (Upstate Biotechnology, Lake Placid, NY, USA; dilution 1:5000) served as the secondary antibody. The membranes were briefly incubated with ECL detection reagent (ECL™, Amersham/GE Healthcare, München, Germany) to visualize the proteins and then analyzed by the Fusion FX7 system (Peqlab, Erlangen, Germany).

b-actin (1:1000; Sigma, Taufkirchen, Germany) served as the internal control.

Gimp 2.8 software was used to perform pixel density analysis of the protein bands. The ratio of protein intensity/b-actin intensity was calculated, and expressed in percentage, related to controls set to 100%.

siRNA knock down studies

Tumor cells (3×10^5/6-well) were transfected with small interfering RNA (siRNA) directed against integrin b1 (2 μM, target sequence: AAAAGTCTTGGAAACAGTCTG, HS_ITGB1_5, Qiagen, Hilden, Germany) or integrin b4 (2 μM, target sequence: GTGGATGAGTTCCGGAATAAA; Hs_ITGB4_5, Qiagen) with a siRNA/transfection reagent (HiPerFect Transfection Reagent; Qiagen) ratio of 1:6. Non-treated cells and cells treated with 5 nM control siRNA (All stars negative control siRNA; Qiagen) served as controls. Subsequently, tumor cell adhesion to immobilized collagen as well as tumor cell migration was analyzed as indicated above.

Statistics

All experiments were performed 3–6 times. Statistical significance was determined by the Wilcoxon–Mann-Whitney-U-test. Differences were considered statistically significant at a p value less than 0.05.

Results

Amygdalin diminishes tumor-endothelium and tumor matrix interaction

Amygdalin significantly reduced attachment of all three bladder cancer cell lines to HUVEC when compared to untreated cells (fig. 1). Cell adhesion of TCCSUP and RT112 was more strongly altered by amygdalin than that of UMUC-3 cells. No difference between short-term (24 h) and long-term (2 weeks) amygdalin treatment was apparent. The binding capacity of UMUC-3, TCCSUP and RT112 cells to immobilized collagen was also significantly down-regulated, compared to controls (fig. 2). Extending the treatment period from 24 h to 2 weeks did not further increase the blocking potential of amygdalin. No sign of toxicity due to amygdalin was detected by the trypan blue exclusion test.

Amygdalin alters the migratory behavior of tumor cells

Exposing the tumor cells to amygdalin for 24 h did not change their migratory activity (fig. 3). However, after a 2 week pretreatment period the number of UMUC-3 and RT112 cells beneath the Transwell chamber membrane was reduced, compared to the untreated control cells. The inhibitory effect in
Figure 4. FACS analysis of integrin α and β subtype expression on UMUC-3 cells. The left panel depicts integrin expression as histogram plots with a dotted line indicating background fluorescence and a solid line indicating specific fluorescence in untreated cells. The right panel shows integrin subtype expression after 24 h and 2 weeks amygdalin exposure, compared to controls set at 100%. n.c. = not calculated. * indicates significant difference to controls.
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Figure 5. FACS analysis of integrin α and β subtype expression on TCCSUP cells. The left panel depicts integrin expression as histogram plots with a dotted line indicating background fluorescence and a solid line indicating specific fluorescence. The right panel shows integrin subtype expression after 24 h and 2 weeks amygdalin exposure, compared to controls set at 100%. n.c. = not calculated. * indicates significant difference to controls.
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and \( b \) were all suppressed by amygdalin (fig. 6, right). The effects on for 24 h or 2 weeks, whereas \( a \) characterized by a high after 24 h.

The integrin subtypes \( \alpha 2, \alpha 3, \alpha 5, \alpha 6, \beta 1 \) and \( \beta 3 \) were strongly expressed on UMUC-3 cells, \( \alpha 4 \) was very moderately expressed and \( \alpha 1 \) and \( \beta 4 \) were not expressed (fig. 4, left). Amygdalin elevated \( \alpha 3 \) but reduced \( \alpha 5, \alpha 6, \beta 1 \) and \( \beta 3 \), independent from exposure time. No amygdalin induced modification was noted in \( \alpha 4 \) integrin. The \( \alpha 2 \) receptor was down-regulated after 24 h but up-regulated after 2 weeks amygdalin exposure (fig. 4, right). TCCSUP cells distinctly expressed the \( \alpha 2, \alpha 3, \alpha 5, \alpha 6, \beta 1 \) and \( \beta 4 \) integrin members (fig. 5, left). The \( \beta 3 \) type was moderately present on the cell surface. Both \( \alpha 1 \) and \( \alpha 4 \) subtypes were not detectable. Amygdalin led to a significant increase in integrin \( \alpha 5, \alpha 6, \beta 1 \) and \( \beta 4 \) on TCCSUP, whereby the 2 weeks application induced stronger effects than the 24 h incubation (fig. 5, right). The \( \alpha 2 \) and \( \beta 3 \) subtypes were enhanced after 2 weeks, but not after 24 h. \( \alpha 3 \) was not altered by amygdalin. RT112 cells were characterized by a high \( \alpha 2, \alpha 3, \alpha 6, \beta 1 \) and \( \beta 3 \) expression level (fig. 6, left). Integrin \( \alpha 5 \) was moderately expressed, and \( \beta 3 \) was only slightly elevated over background. Integrins \( \alpha 1 \) and \( \alpha 4 \) were not expressed on RT112 cells. The integrins \( \alpha 3, \alpha 6, \beta 3 \) and \( \beta 4 \) were all suppressed by amygdalin (fig. 6, right). The effects on \( \alpha 5 \) and \( \beta 3 \) did not depend on whether amygdalin had been applied for 24 h or 2 weeks, whereas \( \alpha 6 \) and \( \beta 4 \) were diminished to a greater extent after 2 weeks, compared to 24 h. \( \alpha 2 \) distinctly increased after 2 weeks but not after 24 h, and \( \alpha 5 \) and \( \beta 1 \) remained unchanged by amygdalin.

**Modifications of integrin proteins by amygdalin**

Alterations of the integrin protein content induced by amygdalin are shown in figure 7 (left) and quantification is expressed as percentage difference between control tumor cells and tumor cells treated with amygdalin (right). In UMUC-3 cells, \( \alpha 6 \) and \( \beta 3 \) were suppressed by both 24 h and 2 weeks amygdalin application, whereas \( \alpha 2 \) and \( \beta 1 \) were up-regulated. An amygdalin induced increase in \( \alpha 5 \) was apparent but this effect was restricted to the 2 week amygdalin application. \( \alpha 3 \) integrin was slightly enhanced over controls after 24 h but not after 2 weeks. The integrins \( \alpha 1, \alpha 4 \) and \( \beta 4 \) were not detectable by western blotting. The integrin related signaling proteins ILK and pFAK were diminished after 2 weeks amygdalin application. Similar action was exerted on TCCSUP cells, since \( \alpha 2, \alpha 5 \) and \( \beta 1 \) increased and \( \beta 3 \) decreased under amygdalin (2 weeks >24 h). In contrast to UMUC-3, \( \alpha 6 \) was enhanced after 24 h but reduced after 2 weeks. \( \beta 4 \), not expressed in UMUC-3, was down-regulated by amygdalin (2 weeks >24 h). The 2 week amygdalin application led to a loss of pFAK and slightly diminished ILK. Evaluation of RT112 revealed increased \( \alpha 2 \) integrin caused by 24 h or 2 week amygdalin treatment. \( \alpha 6 \) and \( \beta 4 \) integrins were down-regulated with 2 week >24 h. There was also a slight increase of \( \alpha 3 \) after 24 h but not after 2 weeks, a phenomenon also seen in UMUC-3 cells. Opposed to UMUC-3 and TCCSUP, the \( \alpha 3 \) subtype in RT112 was only moderately detectable in the control cells and was further diminished following drug treatment. Amygdalin additionally influenced integrin related signaling in RT112, evidenced by diminished FAK and pFAK.
Amygdalin distinctly altered the integrin expression profile of all three bladder cancer cell lines. To investigate whether integrin modifications are relevant to metastatic progression, knock down studies were carried out with the β-integrin members serving as representatives. Since β1 (but not β3 and β4) was highly expressed on control UMUC-3 and TCCSUP and significantly diminished by amygdalin, β1 was knocked down in these cells and adhesion and migration experiments repeated (fig. 8). The β1 integrin was also highly expressed on RT112 but not modified by amygdalin, in contrast to β4, which fulfilled both criteria, i.e. high initial expression and significant modulation by amygdalin. Therefore, β4 was knocked-down in the RT112 cell line before subjection to the adhesion and migration assay (fig. 8, lower right). Loss of β1 was accompanied by a significant reduction in UMUC-3 binding (fig. 8) and migration (fig. 9). On the other hand, TCCSUP binding to collagen was enhanced (fig. 8), whereas TCCSUP migration was not influenced by β1 knock down (fig. 9). Down-regulating the β4 integrin in RT112 cells did not alter the adhesion properties (fig. 8) but massively blocked migration (fig. 9).

**Discussion**

Since tumor cell interaction with the vascular endothelium is necessary for tumor cells to leave the blood stream for establishment at secondary sites, interfering with this process is crucial to hindering metastasis. The present report shows that amygdalin significantly inhibits attachment of bladder cancer cells to endothelial cells. Since amygdalin reduces the number of tumor cells, fewer cells may crawl beneath the endothelial layer to...
establish contact with matrix proteins to metastasize. The next step in metastasis involves the collagen matrix. Interaction of tumor cells with collagen must not only occur to escape the primary tumor but also to permit invasive spreading into the target organ, once the tumor cells have traversed the endothelial blood barrier [11]. In the presence of amygdalin, the tumor cells used in this investigation lost their capacity to bind to immobilized collagen. Therefore, amygdalin may slow metastatic progression by preventing mechanical contacts of circulating tumor cells to the vessel wall and the sub-endothelial matrix. Tumor dissemination, however, is not restricted to binding. The cells must also detach from matrix proteins to invade tissue. Amygdalin influenced the migration capacity of all the bladder cancer cell lines after 2 weeks, but not after 24 h, indicating that long-term treatment may be necessary to alter tumor cell motility.

Amygdalin’s action on the different tumor cell lines was not identical. Migration of UMUC-3 and RT112 was blocked, whereas the number of migrating TCCSUP cells increased with amygdalin. This means that although amygdalin diminishes the attachment rate in all cancer cell lines, there is a risk that chronic amygdalin exposure to a few remaining cells of a particular tumor subtype like TCCSUP may result in increased locomotive activity. Whether due to an acquired or intrinsic resistance or due to the development of undesired feedback loops remains unclear, but this does indicate that not all bladder cancer patients may profit equally well from amygdalin. In line with this speculation, it has recently been shown that resistance development is accompanied by a functional switch of integrin receptors, driving tumor cells to high motility [12].

Chen et al. recently speculated that tumor cells with a high migration speed are much more likely to metastasize than those with a low migration speed [13]. This, however, could not be confirmed by another investigation in which the number of migrating tumor cells and the distance and speed of migration did not correlate with the malignant potential of the tumor cells [14]. Rather, the direction of cell motility indicated the malignant

**Figure 8. Influence of β1 (UMUC-3, TCCSUP) or β4 (RT112) knock-down on bladder cancer cell adhesion to collagen.** Tumor cells were transfected with integrin β1 or β4 siRNA. Non-treated cells (control) and cells treated with scrambled siRNA (siRNA control) served as controls. Efficacy of receptor knockdown was evaluated by western blotting (lower right). One representative of six experiments is shown. *indicates significant difference to the controls.

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potential of the cancer cells [14]. To gain further understanding, animal studies have just been initiated to explore in vivo tumor progression under amygdalin treatment.

The role of integrins in forming cell-cell and cell-matrix bonds necessary for adhesion, extravasation and migration has been broached [15,16], whereby the β1 integrin subunit was shown to play a central role in the bladder cancer cell line T24. However, not all bladder cancer cell lines are characterized by the same integrin pattern. In the present investigation amygdalin modified adhesion and migration and altered integrin expression profiles differently in different cell lines. Integrin β4 was not expressed on UMUC-3 but on RT112 and TCCSUP, whereas β3 was only marginally detectable on RT112 but strongly detectable in UMUC-3 and TCCSUP cells. An investigation involving the influence of valproic acid on bladder cell adhesion to collagen [17], has also revealed modification of the integrin expression profile depending on the cell line employed. Other investigators have reported differing integrin subfamilies in UMUC, T24, J82, RT-4, 253J and Hu456 cells [18,19]. Each cell line may, therefore, possess a characteristic receptor set and drug treatment may influence integrin subfamilies differently.

Investigations on prostate cancer cells have revealed that the initial integrin profile of a particular tumor clone may determine its molecular response to drug treatment [20]. Indeed, amygdalin influenced the integrin composition of the evaluated bladder tumor cells differently. In the UMUC-3 cells β1 and β3 surface expression was diminished by amygdalin, but enhanced in TCCSUP cells. In RT112 cells β4 instead of β1 was altered by amygdalin. Intracellular and membrane integrin levels were also differently affected by amygdalin in UMUC-3 and TCCSUP cells. In UMUC-3 cells intracellular β1 integrin was elevated and surface expression was diminished, indicating that amygdalin induces translocation of β1 integrin away from the surface membrane. In TCCSUP cells β1 integrin expression was increased both intracellularly and on the surface membrane. Amygdalin induced a decrease in intracellular β3 integrin in both UMUC-3 and TCCSUP cells. Surface expression, however, was not the same, with UMUC-3 cells showing an amygdalin induced decrease in β3 integrin and TCCSUP cells showing an increase.

Figure 9. Knock-down of β1 (UMUC-3, TCCSUP) or β4 (RT112) integrin alters chemotaxis. Tumor cells were transfected with integrin β1 or β4 siRNA or scrambled siRNA (siRNA control). Controls remained untreated (control). Values are shown as migrated cells per 0.25 mm². One representative of six experiments is shown. * indicates significant difference to the untreated control. Insert taken from figure 8. doi:10.1371/journal.pone.0110244.g009
The main translocation in TCCSUP cells from the cytoplasm to the surface membrane seems, therefore, to be directed at β3 integrin.

Removal of certain integrin subtypes from the cell surface is not the only mechanism regulating cancer cell adhesion. Integrin trafficking between intracellular compartments and the cell surface has been shown to be a necessary prerequisite to receptor removal at the base of cell protrusions. Recycling integrin back to the leading cell edge supports adhesion [21]. A preclinical study on renal cancer cells has demonstrated that both integrin receptor up- and down-regulation can drive cancer cells towards malignancy. Consequently, therapeutic intervention aimed at ‘re-translocating’ certain integrin molecules might provide an option to prevent malignancy [22].

The relevance of integrins for adhesion and migration was demonstrated by knock-down studies on β integrins with either high initial surface expression or strong amygdalin induced alteration. These criteria were fulfilled for β1 in UMC-3 and TCCSUP cells and for β4 in RT112 cells. Suppression of β1 correlated well with reduced binding and migration activity of UMC-3, which accords to in vitro studies with T24 and 5637 cells [23,24]. Therefore, loss of β1 might be a mechanism whereby amygdalin slows UMC-3 tumor dissemination. TCCSUP cells, however, behaved differently under β1 blockade. Binding events to collagen actually increased, indicating that this receptor in these cells blocks cell-cell or cell-matrix contacts. Differential integrin guided adhesive behavior of different tumor sublines has been previously observed. Blocking the α2 subunit has been shown to inhibit HCV29 bladder cancer cell attachment to the matrix proteins laminin and fibronectin but has an opposite effect on T24 and Hu456 cell adhesion. Similarly, blocking the α5 integrin inhibited HCV29 and BC3726 cell-matrix interaction, whereas binding of the bladder cancer cell lines T24 and Hu456 was enhanced [25].

Since migration in TCCSUP is not modified by β1 integrin, this receptor cannot account for the increased locomotive activity of these cells in the presence of amygdalin. It is possible that α2 and/or β3 integrin may be responsible for their increased motility since both were distinctly elevated when TCCSUP cells were treated with amygdalin for 2 weeks, but not for 24 h, matching the motile behavior of TCCSUP becoming elevated after 2 weeks but not after 24 h.

In RT112 cells a different action of amygdalin may be assumed. β4 integrin was shown to modulate migration but not adhesion, whereby loss of β4 significantly correlated with reduced chemotaxis. Presumably, amygdalin weakens RT112 migratory capacity by diminishing β4 expression.

Aside from the cell-type dependent action on the integrin receptors, amygdalin homogeneously diminished expression of ILK (minor) and activation of FAK (major) in all bladder tumor cell lines used in this investigation. In vitro and in vivo, as well as clinical studies, have implicated ILK over-expression in invasive bladder cancer playing an important role in epithelial-to-mesenchymal transition [26,27]. Likewise, the integrin related protein FAK is critical for promoting urothelial cancer cell motility [28], with a relationship between cell adhesion force and FAK activation level [29]. Pharmacologic suppression of FAK has been shown to prevent cell migration in vitro [30] and inhibit metastasis in a murine bladder cancer model [31]. Down-regulating ILK and FAK by amygdalin might therefore be a pivotal step in counteracting tumor dissemination.

In conclusion, amygdalin’s mode of action varies, depending on the initial tumor cell’s integrin composition. Accumulation or loss of specific integrin members as well as integrin translocation and turnover may contribute to a less aggressive tumor phenotype in the presence of amygdalin. However, conclusions drawn from in vitro investigations cannot be carried over into the clinical setting. Animal studies are underway to evaluate the potential of amygdalin in vivo. The migratory behavior of amygdalin treated TCCSUP cells in vivo will receive particular attention, since amygdalin induced in vitro migration.

Author Contributions
Conceived and designed the experiments: AH RAB. Performed the experiments: JM JR EJ SK. Analyzed the data: IT KN JP. Wrote the paper: KN RAB. Revising and drafting the work: RAB AH KN JP.

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