Selective Toxicity and Angiogenic Inhibition by Euglenophycin: A Role in Cancer Therapy?

Paul V Zimba*, Philip Ordner and Danielle B Gutierrez

Department of Life Sciences, Center for Coastal Studies, Texas A&M University Corpus Christi, Texas, USA

Abstract

A recently discovered euglenoid algal toxin, euglenophycin, is known to have anticancer properties. Euglenophycin is closely related to the alkaloid toxin solenopsin A, originally isolated from fire ants, that is an angiogenic inhibitor. Angiogenesis, the formation of new blood vessels, is required for the development of cancers. Three different human leukemia cell cultures (K562, THP-1, and Jurkat), a murine endothelial (SVR), and epithelial (IEC-6) cell lines were treated with euglenophycin at varying concentrations (0 - 100 μg/ml). After a 48 hour exposure, decreased production of Vascular Endothelial Growth Factor (VEGF) and Angiopoietin 2 (Ang-2) was observed using an Enzyme-Linked Immunosorbent Assay (ELISA). LC50: 48 hour for all cell lines was less than 50 μg/ml. Viability of cell cultures showed a dosage dependent decrease as compared to control using (3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium (MTS) assay and trypan blue exclusion. Morphological traits of SVR cells determined by phase contrast microscopy and scanning electron microscopy supported toxicity and viability findings. These data strongly suggest that euglenophycin possesses angiogenic inhibition properties, including metabolic retardation of VEGF and Ang-2 production and inhibition of cell proliferation against human leukemia and murine endothelial and epithelial cells. Future animal model research is required to assess euglenophycin’s potential as a new therapeutic anticancer treatment.

Keywords: Angiogenesis; Anticancer activity; Euglenophycin; Tumors

Introduction

There are more than 100 different types of cancers that affect humans [1]. Cancers are generally classified via their tissue type of origin. Leukemia, originating from blood producing tissue or bone marrow, is one of the main types of cancer [2]. In people with leukemia, abnormal White Blood Cells (WBC) are produced from the bone marrow and released into the peripheral blood stream [2]. Unlike normal WBC, leukemias exhibit a higher rate of proliferation resulting in an accumulation of dysfunctional cells. Over time leukemia cells can crowd out normal red and white blood cells producing condition of anemia and hypoxia, which can stimulate the production of Vascular Endothelial Growth Factor (VEGF) [3], an important factor in angiogenesis.

Angiogenesis is the process of forming new blood vessels and is essential for tumor formation [4,5]. Paracrine signaling is the major method of cellular communication in the angiogenic process [6]. Endothelial cells that line the interior of all blood vessels are a source of numerous factors (stimulating/inhibiting) and corresponding target receptors that are involved in the angiogenic process including: VEGF-A, VEGF Receptor-2 (VEGFR-2), Angiopoietin-2 (Ang-2), and Tyrosine Kinase with Immunoglobulin-like and Endothelial Growth Factor-like Domains Receptor-2 (TIE2) [3,7,8]. It is widely accepted that cancerous cell masses have a critical size of ~1 mm² and need new vasculature to grow larger [9,10]. Angiogenesis requires endothelial cell proliferation / migration and VEGF for success [11].

In mammals, the VEGF family consists of several members of signaling compounds (VEGF-A,-B,-C,-D,-E, and placenta growth factor). VEGF-A is a potent angiogenic signaling protein that regulates the differentiation, migration, proliferation, permeability, and survival of endothelial cells [12-14]. VEGF165 is the most abundantly expressed of the six isoforms of VEGF-A found in humans [11]. Of the three subtypes of cell surface receptor tyrosine kinases (VEGFR-1,-2, and -3), VEGFR-1 and VEGFR-2 bind to VEGF-A. Angiogenesis is thought to be mediated by VEGF-2 binding and activation [11]. Within this study, VEGF165 will be examined and referred to as VEGF-A for brevity.

Historically cancer angiogenesis, which includes both types of angiogenesis, has been associated with solid tumors; however, newer evidence suggests that angiogenesis is also involved in leukemia formation and proliferation. Like growing tumor masses, leukemia cells depend on angiogenesis in the bone marrow for increased nutrient inflow and cellular waste removal [3].

Alkaloids from fresh water microorganisms, such as the toxin euglenophycin isolated from Euglena sanguinea Ehrenberg, may provide promising options for future novel cancer treatments [15]. This euglenoid species was the dominant algal species present in commercial freshwater ponds during a fish kill event [16], and chemical structure and various biological activities were later reported [15]. Euglenophycin has anticancer activity against HT-29 and HCT-116 cell lines derived from human colon cancer [15]. Interestingly, euglenophycin shares some degree of chemical structure similarity with the alkaloid solenopsin A, a potent angiogenic inhibitor [17]. Solenopsin A is a naturally occurring alkaloid isolated from the fire ant, Solenopsis invicta. Using the SVR cell proliferation bioassay, Arbiser et al., [17] found that solenopsin A can...
prevent the activation of angiogenesis through the Phosphatidylinositol-3-Kinase Pathway (PI3K).

Hypothesis

This study examines cellular effects of euglenophycin exposure on human leukemias (THP-1, Jurkat, and K562), a modified murine endothelial cell line (SVR), and a murine non-cancerous epithelial cell line (IEC-6). We hypothesize that euglenophycin exposure to these cell lines will induce a significant change in the concentration of growth factor(s), the number of viable cells, cellular protrusions, and cellular ultrastructure compared to control when treated for 48 hours. We further hypothesis that euglenophycin can serve as an effective treatment for certain cancers.

Methods

Euglenophycin purification method

Euglenophycin was purified from the single celled euglenoid algae *E. sanguinea* Ehrenberg. At least seven strains of *E. sanguinea* have been identified as toxin producers (Zimba, pers. comm.). Cultures were grown in 20 L - 40 L batches in polycarbonate culture vessels using AF6 media [15]. Culture conditions were 20°C, 12:12 L:D cycle. The culture was filtered using a 7 μm mesh netting to collect a pellet of algal cells, then centrifuged and frozen at -80°C. The cell pellet was thawed and sonicated with methanol to extract toxin as previously described [15]. Euglenophycin was purified through mass-directed fractionation using an Agilent 1200 series HPLC-MS as previously described [15]. The purified euglenophycin was freeze dried (Labconco Freezone 4.5 L) and stored at -80°C. The toxin was reconstituted in Dimethyl Sulfoxide (DMSO) at a final treatment concentration not exceeding 0.1% total DMSO for cell assays.

Cell cultures and media conditions

Human leukemia cell lines Jurkat (clone E6-1), K-562, and THP-1 were purchased from ATCC (Manassas, VA). Jurkat (clone E6-1) was established from a 14 year old male with acute T cell leukemia [18]. K-562 was established from a 53 year old female with chronic myelogenous leukemia [19]. THP-1 was established from a 1 year old male with acute monocytic leukemia [20]. All leukemia cell cultures were maintained in RPMI 1640 medium supplemented with 10% Fetal Bovine Serum (FBS), penicillin (100 U/ml), and streptomycin (100 μg/ml).

Murine endothelial cell line SVR (SVRN 1 ras) strain C57BL/6 was purchased from ATCC. The SVV cells were established from the pancreatic islets of Langerhans endothelium of adult mice. SVR cells were maintained in high-glucose Dulbecco Modified Eagle’s Medium (DMEM) containing 4 mM L-glutamine, and supplemented with 10% Fetal Bovine Serum (FBS), penicillin (100 U/ml), and streptomycin (100 μg/ml).

Murine epithelial cell line IEC-6 was purchased from ATCC. IEC-6 cells were established from non-cancerous rat small intestine epithelium which synthesizes fibronectin and collagen, indicative of normally functioning epithelial cells [21]. IEC-6 were maintained in DMEM containing 4 mM L-glutamine, 1.5 g/L sodium bicarbonate, 4.5 g/L glucose and supplemented with 90% 0.1 U/ml bovine insulin, 10% Fetal Bovine Serum (FBS), penicillin (100 U/ml), and streptomycin (100 μg/ml).

All cell lines were incubated in T25 and T75 cell culture vented-capped flasks at 37°C and 5% CO2. Subculturing occurred two times a week as recommended. Cell integrity was monitored using a phase contrast inverted microscope for morphological assessment; trypan blue stain was used to assess cellular viability/mortality using a compound light microscope. Cell lines were grown for a minimum of one week before experimentation to ensure consistency of exponential cellular growth conditions. All experiments were performed on cell lines between passages 7 - 15 to prevent use of cells undergoing genetic drift or mutation.

Cell counts and viability

An improved Neubauer Bright-Line Hemacytometer was used with trypan blue stain to count viable and dead cells on a compound light microscope. Endothelial and epithelial adherent cell lines were washed with PBS then trypsinized using the Trypsin-EDTA Solution for Endothelial Cells 1X (Sigma, St. Louis, MO). When cells detached from the culturing flask, FBS in fresh RPMI 1640 media was used to inactivate the trypsin. Equal parts of homogenized cell-media mixture and trypan blue stain were combined for 2 - 5 minutes, and then cells were enumerated. For each treatment, counts were performed in triplicate and averaged.

Additionally, the 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxy methyl phenyl)-2-(4-sulfophenyl)-2H-tetrazolium/ phenazinemethosulfate (MTS/PMS) assay called CellTiter 96 AQueous Non-Radioactive Cell Proliferation Assay (Promega, Madison, WI) was performed as a biochemical method to detect viable respiring cells. After one hour of incubation a standard curve was created for each cell line, by plotting absorbance versus cell density. The normalization was necessary to account for differing metabolic activity between cell types and to establish lower and upper assay limits.

Euglenophycin treatment assays

At 50% - 70% confluency, cells were counted and seeded in a 24 well plate with 10,000 cells per well in 1 - 2 ml fresh media. Cells were grown overnight, exposed to 0, 3, 6, 9, 25, or 100 μg/ml of euglenophycin in triplicate for 48 hours, and viable cells were counted based on respiratory activity measured by the MTS assay. For these experimental conditions were used to minimize nutrient depletion during the 48 hour incubation period.

Light microscopy and scanning electron microscopy

Cells were counted using a Zeiss AxioLab compound light microscope. A Nikon TMS inverted microscope with phase contrast was used to observe all cell lines during maintenance and pre/post-experimentation. Phase contrast pictures of murine IEC-6 and SVR cells were taken with a mounted High Tech Computer (HTC) 5.0 megapixel digital camera on the inverted microscope at 400x magnification. Cells were in 24 well plates or T25 culture flask with appropriate growth medium when imaged. Images of controls and treatments were taken to compare morphological differences.

Cell surface morphology of the SVR endothelial cell line SEM was visualized using a modified protocol [22]. Glutaraldehyde fixative (50% EM grade) and 0.2 M Sorenson phosphate buffer pH 7.2 were obtained as a Karnovsky’s Fixative kit (Electron Microscopy Sciences). Glutaraldehyde and osmium tetroxide (dry; Electron Microscopy Sciences) were diluted with phosphate buffer to obtain 2% and 1% working solutions respectively.

Euglenophycin treatments were performed as described above with 12 mm diameter glass coverslips (Neuvitro Corporation) added to the well bottom. After 48 hours of exposure, media was aspirated,
and coverslips were removed and placed in a glass petri dish containing 2% glutaraldehyde for 30 minutes to 1 hour at room temperature. Coverslips were carefully washed (three times) with phosphate buffer, then placed in a glass petri dish containing 1% osmium tetroxide for 30 minutes to 1 hour at room temperature. Coverslips were washed (three times) with phosphate buffer then placed in a dehydration gradient of 25%, 50%, 75%, 90%, and 100% ethanol for 5 - 10 minutes at each gradient step. After dehydration, coverslips were immediately dried using an Eimatech (Ashford, England) KS80 Critical Point Dryer, sputter coated with gold-palladium, and viewed using the JOEL NeoScope SEM.

SVR angiogenesis proliferation bioassay

The SVR angiogenesis proliferation bioassay was used to test for angiogenic inhibition of euglenophycin against endothelial cells as previously described [17]. The SVR cell line has been extensively used to test inhibition of angiogenesis [17,23-25]. SVR cells were treated with euglenophycin as described above. Carbazole was used as a positive control as it is a known angiogenic inhibitor [26]. After a 24 - 48 hour incubation period, media was aspirated and examined for non-attached dead cells. Wells were again trypsinized and cells counted after trypan blue staining using a hemocytometer. The SVR bioassay was repeated three times.

Enzyme-Linked Immunosorbent Assays (ELISA)

VEGF and Ang-2 concentrations were assayed using a human VEGF ELISA kit (Thermo Scientific, Waltham, MA) and a human Ang-2 ELISA kit (Invitrogen) in Jurkat, K-562, THP-1, and murine SVR cell lines. Using this assay, the minimum detectable concentrations of VEGF and Ang-2 are > 8.0 pg/ml and > 6.0 pg/ml, respectively. All treatments were performed in triplicate or greater according to manufacturer protocols. After a 48 hour incubation period absorbance values were obtained using a thermo Scientific Multiskan Spectrum plate reader. A standard curve was generated from plotting the average absorbance for each standard versus the corresponding concentration. Blank wells containing only growth medium provided a baseline control.

Statistical approaches

The statistics software package JMP 9.0.2 (SAS Institute) was used for statistical analysis of experimental data. Data were examined for normalcy using the Shapiro-Wilk test (goodness-of-fit test) in addition to viewing normal quantile plots, histograms, and box plots. Data sets were transformed (log+1) if not distributed normally. One-way Analysis of Variance (ANOVA) was conducted to test significant differences among euglenophycin treatment means. Treatments having significant differences were compared using Tukey-Kramer HSD (α = 0.05).

A 48 hour LC50 probit value was manually calculated. The logEC50 values of euglenophycin treatment concentrations were calculated (n = 3). Percent mortality was calculated by the formula [(dead cells) / (live + dead cells)]*100. Abbott's formula [(Treatment_dead-Control_dead) / (100– Control_dead)]*100 was used to obtain corrected percent mortality values. Corrected percent mortality was converted to probit values manually using Finney's table [27]. Probit values versus logEC50 of concentrations were graphed and a regression line was fitted. LC50 value was determined by solving for a Probit value (Y-value) of 5 (or 50%) then taking the inverse log of each concentration.

Results and Discussion

Cell viability

Human leukemia cell lines exhibited a dosage dependent response to euglenophycin concentration, with viable cells decreasing as euglenophycin concentration increased. Euglenophycin affected K562, THP-1, and Jurkat viable human leukemia cell numbers similarly, with 16 - 43 μg/ml euglenophycin treatment for 48 hours reducing cell numbers by 50% and 25 μg/ml euglenophycin causing over an 8-fold decrease in viability compared to control values. One hundred percent mortality occurred at 100 μg/ml. Significant differences in cell survival were evident from ANOVA analysis (Figure 1).

Figure 1: VEGF concentration determined by ELISA after 48 hour euglenophycin exposure in K562 (A), THP-1 (B), Jurkat (C), and (D) SVR cell lines. In all treatments, n = 3, standard deviation as indicated by error bars. Treatment groups having different letters are significantly different from control (α = 0.05). Letters above bars indicate Tukey-Kramer HSD post-hoc results (P = 0.05).

In this study, SVR modified endothelial cells were selected for euglenophycin treatment as a known assay for screening the anti-angiogenic potential of euglenophycin, a compound similar to solenopsin-A [17]. Euglenophycin inhibited the growth of ras-transformed SVR cells in a dosage dependent manner (Figure 2). Over 50% mortality was measured at 16 μg/ml euglenophycin with 100% SVR mortality at 100 μg/ml. Similarly, solenopsin-A reduced SVR cell number by 50% at 3 μg/ml as previously described [17]. Solenopsin-A is known to inhibit angiogenesis through the PI3K pathway [17] which suggests that euglenophycin may inhibit SVR cell proliferation. Arbiser et al. [17] found that only solenopsin-A was
effective in inhibiting angiogenesis while other analogs of solenopsin with shorter carbon chains were not.

Euglenophycin LC₅₀ in K562, THP-1, Jurkat, SVR, and IEC-6 cell cultures after 48 hour exposure ranged from 16-43 µg/ml. K562 leukemia had the highest cellular growth in the control treatment (Figure 3) which may explain the higher LC₅₀ value of 43 µg/ml - higher division rates would compensate for cell mortality. The non-cancerous cell line IEC-6 had the 2nd highest LC₅₀ of 40 µg/ml while the SVR endothelial cell line had the lowest at 16 µg/ml. These results suggest that SVR ras-transformed cells are more sensitive than IEC-6 cells to euglenophycin for growth inhibition, indicating euglenophycin’s potential as an anti-angiogenic therapeutic.

Morphological assessment via phase contrast and Scanning Electron Microscopy (SEM) provided further insight into euglenophycin’s toxicity on a cellular level. In this study, cellular protrusions are referred to as pseudopodia, as distinction between pseudopodia, filopodia, and lamellipodia were not made (exact location and distribution of cellular actin filaments and microtubules were not examined). Cell morphology was significantly affected by euglenophycin treatment. With increasing euglenophycin concentration, cell-to-cell contacts decreased and the pseudopodia stretched bi-directionally; at the highest concentrations, cellular rounding was observed (Figure 4). This suggests that euglenophycin alters endothelial surface communications, which require cell surface contact either through physical or ligand / receptor interactions. Euglenophycin’s exposure to SVR endothelial cells appears to immobilize them by inhibiting cellular protrusions. Endothelial migration is an essential component of angiogenesis which is orchestrated by cellular signals that are associated with cytoskeleton remodeling such as the PI3K pathway [28].

Via SEM it was apparent that microvilli were present on the squamous non-treated cells but decreased with higher concentrations of euglenophycin treatment. At dosages >25 µg/ml, the sphere-like cell surfaces appeared blebbled or furrowed and not entirely smooth. SVR microvilli and pseudopodia may be acting as antennae on the cellular protrusions for the endothelial cells much like an endothelial tip cell [29,30]. Similar observations were made in a study of spontaneous hypertensive rat endothelial cells in which numerous microvilli were present on the apical surface [31]. Lamalice et al., [28] also states that endothelial cell migration is partially driven by chemotaxis, in which VEGF and angiopoietin gradients are detected by membrane protrusions (filopodia).

Effect of euglenophycin on VEGF and ANG-2 concentration

Leukemia, as with other cancer, proliferates within complex microenvironments in which cells have increased levels of growth factors such as VEGF and Ang-2 [3,7,14]. These growth factors are important steps in the angiogenic process of forming new blood vessels. Most cells involved in the angiogenic process are non-cancerous; however, they assist in forming new vasculature supporting uncontrolled cancer growth. Figure 2 shows VEGF concentration for human K562, THP-1, Jurkat, and murine SVR cell cultures after a 48 hour euglenophycin exposure period. With a 6 µg/ml euglenophycin treatment, all leukemias had an 11% - 44% decrease in VEGF concentration. SVR cells experienced a 22% decrease as compared to control. At concentrations of 9-100 µg/ml, all cells showed a reduction in VEGF concentration after toxin exposure; however, the response was not dosage dependent and some treatment values were not significantly different from each other.

All leukemia cells exhibited a similar Ang-2 response to euglenophycin. Non-treated leukemia cells had Ang-2 concentrations from 653 pg/ml to 1721 pg/ml (Figure 3). K562 cells experienced a 58% decrease in Ang-2 concentration as compared to control after a 6 µg/ml euglenophycin treatment. At 100 µg/ml euglenophycin, all leukemia cell lines had Ang-2 levels of 78 pg/ml – 238 pg/ml. Significant differences in Ang-2 concentration were evident from ANOVA analysis for the three cell lines (Figure 3).

Interestingly, all leukemias showed detectable concentrations of VEGF and Ang-2 for 100 µg/ml euglenophycin treatments while each well had no detectable viable cells present. One explanation is the half-life of VEGF is 6-8 hours under hypoxic conditions [32]. The leukemia cells could have produced the VEGFs then died. Another possible explanation is that not all the background absorbance was removed prior to measuring ELISA absorbance results.

Qualitative analysis of VEGF and Ang-2 provides insight into euglenophycin’s effect on SVR cells (Figure 3). A reduction in VEGF concentration occurred with increased euglenophycin concentration, which also caused SVR endothelial cells to have fewer cellular protrusions and microvilli.

Murine SVR ultrastructure was examined by SEM after euglenophycin exposure (0, 3, 6, 9, 25, and 100 µg/ml dosage) for 48 hours (Figure 5). Pseudopodia loss occurs with increasing concentrations of euglenophycin. Microvilli were present on the squamous non-treated cells but decreased with higher concentrations of euglenophycin. SVR cells dosed with euglenophycin (0, 3 and 6 µg/ml concentration) had loss of ultrastructure as euglenophycin concentrations increase, which
was more apparent when comparing control to 25 μg/ml euglenophycin treated endothelial cells. At 25 μg/ml euglenophycin, the SVR cell surface appeared smooth and without microvilli. Similar to phase contrast observations, a noticeable rounding of cells and pseudopodia loss was observed at higher euglenophycin concentrations. At 100 μg/ml, spherical shaped cells had no defined ultrastructure, such as pseudopodia and microvilli, as compared to control. The sphere-like cell surfaces appear blebbed or furrowed and not entirely smooth.

A cell's surface area is reduced when the number of cellular protrusions and microvilli decrease, which presumably causes a concomitant reduction in the number of cell surface receptors, such as VEGFR-2 and Ang-2. Reducing the paracrine signaling of VEGF between endothelial cells would assist in the inhibition of the angiogenic process through immobilization of endothelial cells.

In summary, euglenophycin can reduce the number of viable leukemia cells and reduce leukemic metabolic activity in vitro. Having a reduced number of viable leukemia cells, which produce less pro-angiogenic growth factors, is desirable in cancer therapies. Euglenophycin's ability to inhibit SVR growth and its similar chemical structure to solenopsin-A suggest that future studies could help determine if euglenophycin acts to inhibit angiogenesis through the PI3K pathway. Evidence herein of euglenophycin's ability to inhibit angiogenesis in vitro should stimulate future research in an animal model.

Conflicts of Interest
The authors declare no conflict of interest.

Acknowledgement
This research was in part supported by the Center for Coastal Studies, Morris J. Lichtenstein Medical Foundation, and NSF. J Cancer Biol Treat 3: 008.

References
1. National Institutes of Health, What Is Cancer? National Cancer Institute, Maryland, USA.
2. National Institutes of Health, What You Need To Know About Leukemia. National Cancer Institute, Maryland, USA.


