Cytotoxic and Apoptotic Effects of Urtica dioica Agglutinin on AGS Cells

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Introduction

Urtica dioica, conventionally stinging nettle, in plant kingdom it is a species of the Urticaceae family. U. dioica is a perennial flowering herb with rhizomes, ovate leaves covered with bristly stinging hairs and have fleshy, drooping, serrated, roughly heart-shaped leaves [1]. Additionally, Urtica prefers wet, rich soil and tends to grow in large patches. The plant is distributed diversified regions of the world. It favours illuminated areas and nutrient rich soils to inhabit. U. dioica is endemic in the Europe and Asia [2]. It is outspreaded to North and South Africa, India, China, New Zealand, Australia, North and South America and endemic throughout Europe in particularly. It is very popular in Black Sea region of Turkey and another part of Turkey. All are considered therapeutically interchangeable. The seeds, rhizomes and leaves of U. dioica have been used in alternative medicine as a secondary remedy source in treatments of illnesses [3,4]. U. dioica is a small (8.5 kDa) monomeric protein agglutinin (UDA), at different concentrations (1, 5, 10, 20 µg/ml) on AGS cell line by carrying out WST-1 (cell proliferation), LDH (cytotoxicity), xCELLigence (real-time cell counter) and TUNEL (apoptosis) assays, respectively. According to our results different concentrations of UDA have anti-proliferative and apoptotic effects on AGS cell line after 48 hours. The RTCA assay indicated that lectin exhibited cytotoxicity, with an IC50 value 4.96µg/ml. Therefore, our current information regarding UDA confirms that UDA may be used in cancer therapeutic strategies as a primary cure in the future.

Keywords: Urtica dioica; Urtica dioica agglutinin; AGS; Cytotoxicity, Genotoxicity

Material and Methods

All products used for cell culture (Dulbecco’s Modified Eagle’s Medium (DMEM), fetal bovine serum (FBS, -20°C), phosphate buffer saline (PBS), L-glutamine and penicillin-streptomycin (PS) were purchased from Biochrom AG (Mannheim, Germany, +4°C). Cell Proliferation Reagent (WST-1) kit, LDH cytotoxicity detection, TUNEL kits were purchased from Roche (Mannheim, Germany, -20°C). Pure and FITC conjugated Urtica dioica agglutinin from stinging nettle, EY Laboratories, (San Matea, CA, USA, -20°C) Cell death induced by the binding of lectin was examined in a human gastric adenocarcinoma cell line (AGS cells) in vitro. All other chemicals were of reagent grade.

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LDH activity assay
Detection according to the manufacturer instructions, Cytotoxicity Assay Kit (LDH; Roche) is used to measure the amount of lactate dehydrogenase (LDH) released by dead cells. The conversion of tetrazolium salt into a red formazan dye product was measured calorimetrically. For this assay, a positive control (PC), leading to 100% cytotoxicity by lysis the cells, was included in the assay. As a positive control (i.e., decreasing viability) 2% Triton X-100 solution in the assay medium, as proposed by the manufacturer was used. As a negative control (equivalent to normal viability) the cells were cultured without any substance in full medium. After preincubation of the cells, before addition of the test compounds at concentrations (UDA) 1, 5, 10, 20 µg/ml, the growth medium was exchanged from medium containing 5% FBS to medium containing only 1% FBS. For testing the released LDH activity, 100 µl of culture medium was transferred to a new 96-well plate at 24th and 48th hour. The reaction solutions of 100 µl from the kit, containing the detection dye and the catalyst, were added to the wells and kept in the dark at room temperature for 30 min at 490 nm with 690 nm as a reference wavelength in an enzyme linked immunosorbent assay (ELISA) microplate reader (BioTek-Power Wave, Winooski, USA). A culture medium without cells was used as a blank. As for the other assays, background values from wells without cells were subtracted and average values for the six times calculated. Means ±SD values are represented in Figure 1.

Cytotoxicity (%) = (experimental value – negative control)/ (positive control-negative control) x100

Proliferation assay
AGS cell line was seeded in triplicate at 1x10⁴ cells/well to the 96 well plates in final volume of 100 µl/well culture medium in a humidified atmosphere (37°C, 5% CO₂) for 24 h before the treatment. After that cell lines were exposed to the determined concentrations of Urtica dioica agglutinin (UDA) 1, 5, 10, 20 µg/ml. Cell proliferations were evaluated for AGS cell line at 24h and 48 h were determined with the WST-1 cell proliferation assay (Roche). Cell Proliferation Reagent WST-1 was added in a 10 µl/well volume. Cells were incubated for 4h in a humidified atmosphere (37°C, 5% CO₂). As a negative control (equivalent to normal viability) the cells were cultured without any substance in full medium. The samples absorbance was measured at 420 nm through ELISA reader.

x CELLeigence system for analyzing the cell proliferation
AGS cell lines was grown and expanded in tissue-culture flasks. After reaching 75% confluence, the cell lines were washed with PBS, afterwards detached from the flasks by a short time treatment with trypsin/EDTA. Subsequently, 100 µl of cell culture media at room temperature was added into each well of E-plate 96. After this the E-plate 96 was connected to the system and checked in the cell culture incubator for proper electrical-contacts and the background impedance was measured during 24s. The cell line was resuspended in cell culture medium and adjusted to 10,000 cells/ml at the same time. 100 µl of each cell suspension was added to the 50 µl medium containing wells on E-plate 96, in order to determine the optimum cell concentration. After 30 minutes of incubation at room temperature, E-plate 96 was placed into the cell culture incubator. Finally, adhesion, growth and proliferation of the cells was monitored every 15 min for a period of up to 24h via the incorporated sensor electrode arrays of the E-Plate 96. The electrical impedance was measured by the RTCA-integrated software of the xCELLigence system (Roche) as a dimension less parameter termed CI. The proliferation, attachment and spreading of the cells was monitored for each 15 minutes by the xCELLigence system. Approximately 24h after seeding the medium in the well each was discarded in order to get rid of the dead cells. When the cells were in the log growth phase, the cells were exposed to 100 µl fresh DMEM containing the following substance: UDA (1, 5, 10, 20 µg/ml). DMEM and 10% FBS were used as negative control group. All experiments were run for 48h. Cell index (CI) is derived to represent cell status based on the measured relative change in electrical impedance that occurs in the presence and absence of cells in the wells Impedance is measured at 3 different frequencies (10, 25 or 50kHz) and a specific time.

TUNEL assay
DNA fragmentation is an indicator for detecting the late apoptosis. TUNEL (Roche) is an assay that is composed of an enzyme solution and labeling solution which detects the nicks (single strand breaks) in DNA and binds to free 3'-OH ends. Adding of dUTPs to 3'-OH ends by terminal deoxynucleotidyl transferase cause labeling of DNA. Fluorescently labeled ends were detected by using confocal microscopy (Leica).AGS was used in this technique. The cell line was grown for TUNEL assay. AGS cells were seeded at 1x10⁴ cells/well to the 96 well plates in final volume of 100 µl/well culture medium in a humidified atmosphere (37°C, 5% CO₂) for 24h before the treatment. Then the next day the highest concentration of UDA, 20 µM, was chosen for all of the cells. The cells were treated with 20 µg/ml UDA for 48h. Mitomycin C (10 µM) was used for the positive control. As a negative control (equivalent to normal viability) the cells were cultured without any substance in full medium. After the incubation period, the cells were washed twice with phosphate buffer solution (PBS), then fixed with 5% paraformaldehyde for 60 minutes and washed again twice with PBST (PBS+0.1% Triton-X-100) on the shaker. 0.1% Triton X were treated with 200 µl with cells on the ice for 10 minutes. After that the cells were washed twice with PBST on shaker. TUNEL reaction mixture was added 50 µl and incubated at 37°C in a humidified atmosphere with 0.5% CO₂ for an hour. After that the cells washed with PBST. And then DAPI (4',6-diamidino-2-phenylindole) is a fluorescent dye was applied on the cells for 15 minutes then washed once with PBST and once with distilled H₂O. The results were analyzed by fluorescence microscopy.
Binding on cell surface

AGS cancer cell line was passaged right after reaching desirable density in flasks. Passaged cells were seeded in 96 well-plate for each plate to be 10,000 cells. Plate was incubated for 24 h to provide the cell growth and attachment. After 24 h diluted fluorescent labeled lectin (EY Laboratories, USA) was applied on cell culture at 20 µg/ml UDA concentration. After 24 h of incubation, cells were washed with buffer solution three times. Cells were examined with or without fixation under fluorescent microscope. Appropriate filter was used and FITC absorption was 492 nm and emission rate was 517 nm.

Statistical analysis

The statistical analysis of WST-1 and LDH assay values Mann–Whitney U-test was used. Also for TUNEL assay values Chi-square (Χ²) test was used. Statistical differences between time and dosage were analyzed. A value of P less than 0.05 was accepted as statistically significant. Results were expressed as mean ± SE. For these procedures, SPSS 11.5 version for Windows (SPSS Inc, Chicago, Illinois, USA) was used. Cell Index calculations were obtained using the RTCA-integrated software of the xCELLigence system. The RTCA software performs a curve-fitting of selected “sigmoidal dose–response equation” to the experimental data points and calculates logarithmic half maximal inhibitory concentration (IC₅₀) values at a given time point based on log of concentration producing 50% reduction of CI value relative to solvent control CI value (100%) in our results.

Results

Effects of urticadioica agglutinin on AGS Cells

Throughout 24th and 48th hours process of LDH assay, there was a significant effect of U. dioica agglutinin on AGS cells in which amount of LDH has increased. Namely, AGS cell which are exposed to LDH assay were ruptured. In other words, UDA has an effect on lipid peroxidation, either. On the other hand, WST-1 results showed that number of cells has decreased in all concentrations along with the treatment (p<0.05).

In another aspect, xCELLigence pointed out that as the concentration rises up, cell number goes down during both 24 and 48 hours. Also, there is clear difference between control and other concentrations (Figure 2A). The sigmoidal dose response curves of lectin tested in RTCA are shown in Figure 2B. All curves have an r² greater than or equal to 0.99.

Considering TUNEL assay, positive control group and 20 µg/ml concentration are statistically meaningful over negative control (p<0.05) (Figure 3). Hence, U. dioica agglutinin for 20 µg/ml concentration along with 48th hours stated that cells ended up undergoing apoptosis. All indications together revealed that each assay compatible with the others. Particularly, 20µg/ml concentration was the most effective one in terms of cytotoxicity and apoptosis among all concentrations (Figure 4) (p<0.05).

In addition that, Figure 5 represents the 20 µg/ml concentration of FITC-UDA bound to carbohydrate residues in AGS cell membrane.

Discussion

A heterogeneous group of glyco-proteins having in common the unique ability to recognize and bind specific sugar residues is called to be plant lectins [11]. It is also noteworthy that lectins derived from non-immune origin to have numerous links a wide range of pathological complicacies, most notably cancer [16]. Among these mistletoe lectins (MLs), Ricin, Concanavalin A (ConA) and Polygonatum cyrtonema lectin (PCL) can lead to cancer cell programmed death via targeting apoptotic pathways [17]. Moreover, ConA and PCL can also result in cancer cell programmed death. Possible anti-cancer therapeutic implications of plant lectins used as simple tumor recognition tools to differentiate malignant tumors from benign and the degree of glycosylation associated with metastasis [18]. Besides, other lectins such as ConA and PCL can result in cell death after internalization or binding certain sugar-containing receptors on the surface of cancer cells [19]. Plant lectins are thought to be decreasing expression of anti-apoptotic nuclear factor-kB (NF-kB) [18]. Very recently it has been communicated that french bean hemagglutinin has been shown to induce breast cancer MCF-7 cell apoptosis [20]. Subsequently, lectins were found to bear remarkable inhibitory effects on the growth of
MCF-7 and hepatoma HepG2 cells [21]. WGA treated cell at different concentrations (0.5, 1, 2.5, 5 or 10 µM) for 24 h reduced cell viability of H3B, JAr and ROS cell lines [22]. In an effort to detect novel strategies in bladder cancer therapy, the potential and the applicability of different plant lectins was investigated using 5,637 cells as a model for human urinary carcinoma [23]. Specifically, the UDA is not blood group-specific and is specifically inhibited by N-acetyl glucosamine oligomers [11]. Moreover, UDA induces HuIFN-gamma in human lymphocytes at concentrations comparable to those of other inducers [11]. UDA is known having the ability to bind itself to the sugar residues on the cell surface [15].

UDA is first to be researched on cancer cell lines. In the light of previous studies, as a lectin form we thought that it would induce the inhibition of tumor growth in the way of the lectins in previous and recent studies. Accordingly, since the carbohydrate pattern of UDA is reflected by high abundance of N-acetyl-D-glucosamine, we guessed that UDA would attach itself to cells through high carbohydrate content of cell membrane. In another saying, high binding rates, binding specificity would be advantageous for carbohydrate-mediated targeting in carcinoma therapeutics. In an effort to develop novel strategies for cancer therapy, at first the interaction of different concentrations of UDA was chosen. On the basis of former researches, we made a decision to pick the highest concentration as 20 µg/ml.

We looked into cytotoxic, proliferative and apoptotic effects of UDA on AGS cell lines. This study points out that throughout 24 and 48 h of periods of time UDA decreased cell proliferation of AGS.
cell line at discrete concentrations. Number of viable cells was less in comparison to positive control group. In similar way, amount of lactate dehydrogenase enzyme increased along with both periods of time of 24 and 48 hours except for 20 µg/ml concentration. Namely we can interpret that manual assays conformed eachother at the time and dose dependent manner. According to xCELLigence results as the concentration goes up in density biological status of cells like cell number and viability decrease. At this point we thought that there might have been a complicacy or an experimental error between 10 µg/ml and 20 µg/ml concentrations of AGS cell line during the process. Yet, in general we can conclude that for AGS cells as the concentration ascends, number of viable cells escalates. Last but not least, TUNEL results provided us percentage of DNA fragmentation on AGS cells at 20 µg/ml concentration. With the help of DNA fragmentation percent we became able to say that apoptosis is statistically significant. Which collectively means that UDA especially at the highest concentration likely to have cytotoxic effect on AGS cell line. Our study indicates that cell proliferation reduced number of viable cells at the time basis and increasing concentrations. Finally, binding ability of UDA was searched in this study, either. And FITC Conjugated Lectin provided us UDA has the feature of binding itself to sugar moieties on the cell surfaces. It can be said that the surface binding tendency of UDA have been shown.

In literature there is no study about anti-proliferative effects of UDA on AGS cell line. In short, UDA has anti-proliferative and apoptotic effects on AGS cell line. UDA decreases cell proliferation and increases apoptosis in AGS cell line.

**Conclusion**

In this research, we evaluated the updated research of one of the good representative plant lectins that may lead to programmed cell death via targeting apoptotic and cytotoxic pathways in different types of cancer cells. In conclusion, UDA at different concentrations has anti-proliferative and apoptotic effects on AGS cell line. Moreover lowest concentration of UDA, affects AGS cells in cytotoxic manner. Also, this study has the unique feature of being studied with FITCH-labeled lectin.
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