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Expression and characterization of a novel recombinant cytotoxin II from *Naja naja oxiana* venom: A potential treatment for breast cancer



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ABSTRACT

Breast cancer (BC) is among the leading causes of mortality from cancer in women. Many of the available anticancer drugs have various side effects. Therefore, researchers are seeking novel anticancer agents particularly from natural compounds and in this regard, snake venom is still one of the main sources of drug discovery. Previous studies showed potential anticancer effects of Cytotoxin II (CTII) from *Naja naja oxiana* against the different types of cancers. In this study, a pET-SUMO-CTII vector was transformed into SHuffle® T7 Express, an *Escherichia coli* strain, for recombinant protein expression (rCTII) and the cytotoxic effects of this protein was assessed in MCF-7 cells. The flow cytometry assay was applied to measure the apoptosis and cell cycle. Also, mRNA levels of the Bax, Bcl2, P53, caspase-3, caspase-8, caspase-9, caspase-10, matrix metalloproteinases (MMP)-3, and MMP-9 were analyzed by quantitative real-time PCR to determine the underlying cellular pathways affected by rCTII. The results of this study showed that treatment with 4 µg mL⁻¹ of rCTII enhanced apoptosis through the intrinsic and extrinsic pathways. Also, the increase of the cells' proportion in the sub-G1 phase as well as a reduction in S phase was observed. In addition, the expression of MMP-3 and MMP-9 was decreased in the treated group in comparison to the control group that may contribute to the reduced migratory ability of tumor cells. These experimental results indicate that rCTII has anti-proliferative potential, and so this protein could be a potential drug for BC therapy in combination with other drugs.

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1. Introduction

Breast cancer (BC) is the most frequent type of malignancy in women and is also the main cause of cancer mortality in females [1]. Globally, it is reported that more than one million people are diagnosed with BC every year with about 400,000 women death annually, accounting for 14% of the total cancer deaths [2,3]. The conventional treatment

methods for BC, like other cancers, include surgery, chemotherapy, and radiotherapy [4]. The main purpose of these treatments is to exterminate tumors while enhancing the survival rate of the patients. On the other hand, the emergence of therapy resistance is an important obstacle to achieving this purpose [5]. Considering the harmful side-effects presented in the classical cancer therapeutics, the search for novel active drugs for different malignancies is one of the fundamental aims of the biotechnological research and pharmaceutical industries [6]. During the last twenty years, BC therapeutic drugs have been mainly identified in the field of immunotherapy. Regardless of the significant efficacy of monoclonal antibodies e.g. trastuzumab, BC treatment strategies remain unsatisfactory [7,8]. Hence, several advanced, safer, and more efficient biotherapeutics have been developed in laboratories for treating tumors. During recent years, bispecific antibodies [9], oncolytic viruses

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[10], antibody-drug conjugates (ADCs) [11], and toxins [12] have gradually become research hotspots. In the case of toxins, several compounds from snakes or other venomous animals have largely studied and shown promising anti-cancer properties [13–16]. Snake venoms consist of various components including protein, peptide, and other bioactive mixtures, which are isolated from the snakes 'venom glands and injected by unique fangs of snakes to debilitate and digest their prey. Despite its toxicity, a variety of isolated proteins and peptides from snake venoms are functional as pharmaceutical agents with different properties [17,18].

Recent studies have shown that a group of 60-70 amino acids polypeptides, generally named cytotoxins, can kill cancer cells [19-21]. Two specific cytotoxins have identified as cytotoxin-I (CTI) and cytotoxin-II (CTII) from Naja naja oxiana that have been identified for their cytotoxic and cell membrane disturbing actions [22]. Although both cytotoxins have about 80% homology in amino acid sequences, their cytotoxic properties are different [23]. In 2014, Ebrahim et al. investigated the potential anticancer of CTII (4 μ g mL⁻¹) in the MCF-7 cell line. They indicated that CTII has potent anti-BC impacts in the MCF-7 cells via intrinsic apoptosis pathways [24]. More recently, Attarde et al. purified NN-32, a very similar toxin to CTII from Naja naja venom with significant cytotoxicity in both MCF-7 and MDA-MB-231 cell lines in a doseand time-dependent fashion [25]. In spite of all promising data about the anti-cancer effects of CTII, more researches need to identify the structure and molecular mechanisms of its action. Considering the limited availability of natural sources and large quantities that are needed for researches, the production of the recombinant form of CTII could provide enough CTII and promote the investigations. Herein, in this research, we expressed recombinant Oxus cobra CTII in SHuffle® T7 Express, an Escherichia coli (E. coli), and characterized the secondary structure. Finally, the biological activity of the recombinant CTII (rCTII) protein, as well as anti-proliferative mechanisms have been investigated in BC cell line.

2. Materials and methods

2.1. Bacterial strain and cell culture

E. coli SHuffle® T7 Express was used as the expression host for the production of the requested protein. The MCF-7 cell line was obtained from the National Cell Bank of Iran. The cells were cultured in RPMI-1640 medium, which supplied with 10% fetal bovine serum (FBS) and penicillin/streptomycin mixtures (Gibco, Carlsbad, CA, USA) and were grown at 37 °C incubator in a humidified atmosphere with 5% CO_2 level. Each experiment has been repeated three times.

2.2. Construction of pET-SUMO-CTII

The DNA sequence corresponding to the mature form of CTII peptide (*Naja oxiana*, UniProtKB: P01441) was codon-optimized for *E. coli* cell expression. This sequence was synthesized with ubiquitin-like modifier (SUMO) tag by Bioneer Inc. (Daejeon, South Korea) and cloned into pET-28a (+) vector (Invitrogen, California, United States) containing an N-terminal polyhistidine tag (His-tag). Fig. 1 illustrates the mechanisms and strategies for synthesizing the fusion gene.

2.3. Expression of SUMO-fusion proteins in E. coli

The constructed pET-SUMO-CTII vector was transformed into SHuffle® T7 Express for protein expression. A single colony of transformed bacteria was transferred into 10 mL of Luria Bertani (LB) broth medium (Merk, Germany) and spent overnight at 37 °C in a shaking incubator. Next, 1 mL of this medium was mixed with 100 mL of sterilized LB medium and 50 μ g mL⁻¹ Kanamycin. Following that, this medium kept at 37 °C and 150 rpm till they reached the exponential phase (OD_{600 nm} of 0.4–0.6), then the expression processes continued through

the addition of 1 mM Isopropyl- β -D-thiogalactopyranoside (IPTG) and kept for 4 h at 28 °C. At the end of expression time, the pellets of induced bacteria were gathered after 20 min 7000g centrifuge at 4 °C and analyzed by 12% gel SDS-PAGE.

2.4. Purification of SUMO-fused protein

The bacterial pellets were resuspended in lysis buffer, and then the suspension was sonicated five-six times on ice at 60–70% pulse for 20 s with 30 s pauses intervals in between to maintain a low temperature. Then, bacterial debris was separated after 30 min 7000g centrifuge at 4 °C. The final fusion protein was added to the affinity chromatography column packed with immobilized metal ion-affinity chromatography (IMAC) (Qiagen, Germany) as we performed in the previous study [26]. The eluted proteins were examined on SDS-PAGE and collected for further tests.

2.5. Quantification of protein

Determining the size and purity of the obtained protein was analyzed by reducing SDS-PAGE and subsequent staining with silver nitrate. The concentration of the protein was evaluated by both UV–Vis spectroscopy at 280 nm and the Bradford method using standard BSA protein (Sigma Aldrich, Germany).

2.6. Western and dot immunoblotting analysis

For western blotting, purified rCTII and BSA were applied to SDS-PAGE. Then, the proteins were transferred to the nitrocellulose membrane (NC) (Sigma Aldrich, Germany). After blocking with skimmed milk 2% (w/v) in PBS, the membrane was incubated with an anti-Histag monoclonal antibody (Biolegend, USA) diluted in 0.5% (v/v) Tween 20 in PBS buffer. Then after several times washing with PBST buffer (20 mM Tris and 150 mM NaCl, pH 7.6, 0.1% Tween-20), the blots were treated with goat anti-mouse IgG (GAM-HRP) as a secondary antibody (Biolegend, USA) (diluted 1: 3000) and ECL kit (Roche Diagnostics GmbH) was applied to signal detection. For dot blot analysis, 10 µL of purified rCTII, BSA (negative control), and anti-His-tag antibody (positive control) were blotted on NC membrane at 1 cm intervals and left to dry for 1 h at room temperature (RT). Unreacted sites on the membrane were blocked by immersing the membranes by 2% skimmed milk and incubated for 1 h at RT and washed three times with PBST. NC membranes were incubated with GAM-HRP as a secondary antibody (Biolegend, USA) (diluted 1:3000) for 1 h at RT followed by washing three times with TBST. The target proteins were detected and visualized by BCIP and NBT (Sigma, Deisenhofen, Germany).

2.7. Cleavage of SUMO fusions

After 1 mg mL⁻¹ dilution of the eluted target protein and addition of 10 units of SUMO protease, the mixture was kept for 1 h at 4 °C in a high salt buffer (50 mM Tris-HCl, pH 8.0, 0.15 M NaCl, 1 mM DTT). After incubation time, the proteins were transferred to a Ni-NTA resin column (Qiagen, Hilden, Germany) to take the rCTII. The SUMO, uncleaved-SUMO-CTII, and SUMO protease became bound to the Ni-NTA column and just the rCTII was eluted after washing with elution buffer. Imidazole was removed by dialysis with 1× PBS buffer (5.85 g L⁻¹ NaCl, 4.72 g L⁻¹ Na₂HPO₄, 2.64 g L⁻¹ NaH₂PO₄.2H₂O, pH = 7.2) for overnight at 4 °C. The final target proteins were checked on SDS-PAGE and the samples were saved at -80 °C for subsequent structural characterization with CD spectroscopy and identification of its biological activities through different molecular biology techniques.

Cloned nucleotide sequence (from 5' to 3'):



Fig. 1. Schematic representation of the expression vector pET-SUMO-CTIL rCTII was expressed as a fusion protein with the SUMO.

2.8. rCTII secondary structure analysis by CD spectroscopy

To determine the secondary structure of rCTII of SHuffle® T7 Express, a far-UV CD experiment was performed using a model 410 circular dichroism spectrometer (Aviv Biomedical Inc., Lakewood, NJ) in a standard quartz cuvette with 1 mm thickness at 20 °C. Further detailed conditions of CD spectroscopy were identical to those described by Hoffmann and colleagues [27]. For this analysis, 400 μ g mL⁻¹ of recombinant protein in 1× PBS buffer pH 7.0 was used. Spectra were adjusted for buffer additions and transformed to mean residue ellipticity according to the previous study [28]. The changes in secondary structure content were calculated with CDNN 2.1 software.

2.9. Cell proliferation assays

The half-maximal inhibitory concentration (IC50) values of rCTII on MCF7 cell line was evaluated using MTT (3-(4, 5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium bromide) assay. The cells were grown ($15*10^3$ cells/well) in 96-well plates for 24 h at 75% confluency. Then cells were treated with different concentrations of rCTII for 24 h. At the end of treatment, cells were exposed to MTT (5 mg mL⁻¹) and incubated at 37 °C for 4 h. After the removal of the supernatant, 100 µL of DMSO was added to each well and the plates were left incubated for 10 min to solubilize the formazan crystals. Absorbance for each well of plates was measured using a Sunrise microplate reader (Tecan, Groedig, Austria) at a test wavelength of 570 nm and a reference wavelength of

620 nm. The optical density (OD) was calculated as absorbance at the reference wavelength minus that of the test wavelength.

2.10. Apoptosis assay

2.10.1. Annexin V/PI assay

Apoptosis was measured by annexin V/propidium iodide (PI) assay. MCF-7 cells were seeded in six-well plates (SPL Life Science Co) at a density of 3×10^5 cells and examined in control and treated groups. After 24 h treatment with 4 µg mL⁻¹ of rCTII, the treated and untreated cells were detached with trypsin/EDTA and centrifuged at 1500g for 7 min. The cells were then treated with Annex V and PI according to the instructions of the manufacturer (Roche). For Annexin V staining, 5 µL of Annexin V, 5 µL of Propidium Iodide, and 200 µl of Binding Buffer were applied to the sample. Then they were incubated for 15 min at room temperature under the dark condition. Finally, the samples were analyzed by flow cytometry instrument (Milteny BiotecTM FACS Quant 10; Milteny Biotec, Bergisch Gladbach, Germany). The rate of apoptosis was assessed by FlowJo software (Tree Star, San Carlos, CA).

2.10.2. Cell cycle analysis

For investigating the cell cycle arrest properties of rCTII, the MCF-7 cells were pelleted 24 h after 4 μ g mL⁻¹ incubation. Following that, these cells were fixed with 50% ethanol, treated with 5 mg mL⁻¹ RNase A (Daejeon, Korea), and then used 50 μ g mL⁻¹ of PI solution for staining, and assessed by FCM system for cell cycle progression study.

2.11. Wound healing assay

The migration rate of the MCF-7 cells was assessed by using a wound-healing assay. For this analysis, approximately 3×10^5 of MCF-7 cells were cultured in the six-well plates for 24 h, and in confluency of around 90%, the wound was created by the use of the yellow pipette tip across the cell monolayer to make an open gap to mimicking the wound. Then the cell debris washed away with a serum-free medium. Wells were divided into two groups (a treated group with 4 µg mL⁻¹ of rCTII and the untreated groups). The cells were photographed under the inverted microscope and kept for the next 24 h at the incubator and permitted to migrate to the wound zone and images of the plate were collected at 0, 12, and 24 h. The migratory ability of the cells was analyzed by determining the distances between the edges of the wound.

2.12. RNA preparation, cDNA synthesis, and qRT-PCR

The expression of apoptosis and migratory genes (Bax, Bcl2, P53, caspase-3, caspase-8, caspase-9, caspase-10, MMP-3, and MMP-9) was analyzed by qRT-PCR. For this examination, in each experimental group, cells were subjected to RNA extraction using TRIzol reagent (RiboEx). After cDNA synthesis using a kit (Bio FACT, Daejeon, South Korea), the qRT-PCR was performed utilizing light cycler 96 (Roche Diagnostics, Mannheim, Germany). The data were analyzed using $2^{-\Delta\Delta CT}$ method. The sequences of each primer are shown in Table 1.

2.13. Statistical analysis

All data were presented as mean \pm standard deviation (SD). Statistical significance of differences between variables was assessed via *t*test analyses by using GraphPad Prism 6 software (San Diego, CA, USA). The *p*-value below 0.05 was deemed relevant.

3. Results

3.1. Expression and purification of His-SUMO-CTII fusion protein

To prevent intracellular aggregation of the rCTII, the protein was expressed under the T7 promoter with a small SUMO tag. SUMO was used to help in the proper folding of the protein and therefore in the production of soluble proteins. In addition, a SUMO specific protease was used to the elimination of the SUMO tag and production of N-terminal amino acids free proteins [29]. SHuffle® T7 Express cells

Table 1	
Sequence of primers used in this study	

Primers	Sequences
Bax	5'-TTTGCTTCAGGGTTTCATCCA-3'
	5'-TCTGCAGCTCCATGTTACTGTC-3'
Bcl2	5'-CTGTGGATGACTGAGTACCTG-3'
	5'-GAGACAGCCAGGAGAAATCA-3'
P-53	5'-GAGCTGAATGAGGCCTTGGA-3'
	5'-CTGAGTCAGGCCCTTCTGTCT T-3'
Caspase-3	5'-ATGGTTTGAGCCTGAGCAGA-3'
	5'-CATCCACACATACCAGTGCGTA-3'
Caspase-8	5'-CTGGTCTGAAGGCTGGTTGTT-3'
	5'-GTGACCAACTCAAGGGCTCAG-3'
Caspase-9	5'-CCTTCCTCTCTTCATCTCCTGCT-3'
	5'-TTGCTGTGAGTCCCATTGGT-3'
Caspase-10	5'-AATCTGACATGCCTGGAG-3
	5'-ACTCGGCTTCCTTGTCTAC-3'
MMP3	5'-TACTGGAGATTTGATGAGAAGAG-3'
	5'-TACAGATTCACGCTCAAGTTCC-3'
MMP9	5'-ATTCATCTTCCAAGGCCAATCC-3'
	5'-CTTGTCGCTGTCAAAGTTCG-3'
GAPDH	5'-AAGGTGAAGGTCGGAGTCAAC-3'
	5'-GGGGTCATTGATGGCAACAA-3'

were first successfully transformed with pET-28a vector containing His-SUMO-CTII sequence and then induced for expression of interest protein by 1 mM of IPTG at 28 °C. The induction analysis of SDS-PAG proved the generation of recombinant major-band protein at approximately 18.5 kDa that was compatible with the anticipated molecular weight (MW) of His-SUMO-CTII (Fig. 2). SUMO-CTII fusion protein with N-terminal His-tag was purified by affinity chromatography on Ni-NTA column. SUMO-CTII was eluted from the column and analyzed by SDS-PAGE which appeared with an Mw of ~18.5 kDa (Fig. 2).

3.2. Immunoblot analysis

The identity of purified His-SUMO-CTII was confirmed by dot and western blot analysis. For dot blot analysis, the purified His-SUMO-CTII and Anti-His Tag (positive control) were detected while BSA as negative control didn't detect (Fig. 3A). Also, for western blot analysis, as shown in Fig. 3B, His-SUMO-CTII was detected while the BSA as negative control didn't detect.

3.3. Proteolytic digestion and scale-up purification of rCTII

Purified SUMO-CTII was subjected to cleavage reaction by SUMO protease for removal of the SUMO tag. The presence of two bands of rCTII and SUMO on 12% SDS-PAGE gel confirmed the successful cleavage of the SUMO tag (Fig. 2). These results verified that the tertiary structure of SUMO was properly formed in *E. coli* and identified by SUMO protease. rCTII was purified from cleavage reaction by Ni-NTA column and then collected in the flow through. After removal of the SUMO tag, purified rCTII was resolved by 12% SDS-PAGE that showed a single band of ~6.6 kDa (Fig. 2). The data from Bradford assay has also exhibited



Fig. 2. SDS-PAGE analysis of the purification steps of rCTII. Samples from different purification steps were analyzed by 12% SDS-PAGE. Lane 1: Cell lysate insoluble fraction (centrifugation pellet) of *E. coli* SHuffle® T7 Express pET-SUMO-CTII not induced; Lane 2: Cell lysate insoluble fraction *E. coli* SHuffle® T7 Express pET-SUMO-CTII induced; Lane 3: Pre-stained Protein Ladder (PM2700 Protein Marker, SMOBIO®, Taiwan); Lane 4: purified sUMO-CTII; Lane 5: Purified rCTII after SUMO protease cleavage.



Fig. 3. A dot blot (A) and western blotting (B) analysis of His-SUMO-CTII. (A) 1: Ani-Histag antibody (positive control); 2: Purified His-SUMO-CTII; 3: BSA (negative control). (B) 1: Purified His-SUMO-CTII; 2: BSA.

that the concentration of rCTII was 400 μ g mL⁻¹ in *E. coli* SHuffle® T7 Express.

3.4. Biophysical analysis of rCTII

The CD approach aims to the rapid investigation of the secondary structure and environmentally caused structural alterations of proteins [30]. In this study, the secondary structure of rCTII was studied by CD at 25 °C and the obtained spectrum was analyzed with CDNN software (version 2.1) according to previous studies and shown in Fig. 4A [31]. The content of different secondary structures of rCTII was measured and the results included 6.106% α -helix, 35.87% antiparallel β -sheet, 5.15% parallel β -sheet, 18.7% β -turn, and 34.16% random coil. These findings revealed that the β -sheet and random coils are the dominant secondary structures of rCTII.

3.5. The effect of CTII on the visibility of MCF7 cells

CTII decreased the number of living cells in the MCF7 cell line in a dose-dependent manner. MTT result showed that the IC50 of rCTII for MCF7 was 3.66 μ g mL⁻¹ which have shown in Fig. 4B. In our study, we used 4 μ g mL⁻¹ for the evaluation of the apoptosis, cell cycle, migration, and gene expression tests.

3.6. CTII enhanced apoptosis of MCF-7 cells

3.6.1. CTII induced the early and late apoptosis of MCF-7 cells

To evaluate the potential anti-proliferative function of rCTII, we assessed the effects of this recombinant protein on apoptosis of MCF-7 cells through Annexin-V/PI staining. The CTII treated and untreated cells incubated for 24 h. In Annexin-V/PI assay, Annexin V positive (+) and PI-negative (-) cells were characterized as early apoptotic cells and double-positive Annexin V (+) and PI (+) cells were considered as late apoptotic cells. In treated MCF-7 cells after 24 h exposure with rCTII, the percentage of Annexin V+/PI+ (late apoptosis) and Annexin V+/PI- (early apoptosis) cells were statistically significant than untreated control cells (Fig. 5A and B). The summarized apoptotic data was demonstrated in Fig. 5C. In addition to this flow cytometry study, the apoptosis-related genes were assessed by Real-time PCR to further validation of the effect of rCTII on apoptosis.

3.6.2. CTII induced the arrest of the cells in the sub-G1 phase

Cell cycle analysis of the cells that incubated with $4 \ \mu g \ mL^{-1}$ of rCTII presented in Fig. 6. The results showed that there is an increasing ratio of the cells in the sub-G1 group (apoptotic cells) after treatment. The proportion of the cells in the sub-G1 phase was 0% in the control group in comparison with 35.5% in treated cells. In addition, a



Fig. 4. A: Circular dichroism (CD) analysis of purified rCTII. The spectrum was recorded with the Aviv CD spectrometer at 20 °C in a 1 mm quartz cuvette. The protein concentration was (400 μ g mL⁻¹). B: Dose-response cell viability curves of recombinant CTII in MCF7 cells. The cells were treated with different concentrations of recombinant cytotoxin-II for 24 h and at the end of the incubation time, cell viability was determined by the MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) reduction assay. The IC50 of rCTII in MCF7 cell line was 3.66 μ g mL⁻¹.

decreasing trend in the S phase in treated cells compared to the control group was observed in the study (Fig. 6).

3.7. CTII treatment reduced the cells migration ability of MCF-7 cells

A wound-healing approach was done to assess the migration rate of MCF-7 cell line in rCTII treated and untreated groups. The wound space was recorded at 0, 12, and 24 h. The results indicated that the migration of MCF-7 cells was significantly inhibited after 12 h and 24 h treatment with rCTII (Fig. 7). Moreover, for further evaluation of the migration rate of MCF-7 cells following rCTII treatment the MMP-3 and MMP-9 genes expression were evaluated to find the effect of rCTII on the migration of the cells.

3.8. CTII treatment induced the apoptosis and reduced the migrationrelated mRNA expression levels of MCF-7 cells

To evaluate the underlying pathways of apoptosis, the expression of P53, Bcl2, Bax, caspase-3, caspase-8, caspase-9, and caspase-10 was evaluated by real-time PCR in the control and rCTII treated group. Our result illustrated that this protein potentially activated both intrinsic and extrinsic pathways related to apoptosis. P53, Bax, caspase-3, caspase-8, caspase-9, caspase-10 (*p*-value: 0.0002, *p*-value <0.0001, *p*-value <0.0001 respectively) were upregulated in the treated group in comparison with control, on the other hand, Bcl2 (*p*-value <0.0001) as an anti-apoptotic factor was down-regulated in the treated group. Furthermore, the expression of MMP-3 and MMP-9 (*p*-value <0.0001 and *p*-value <0.0001



Annexin V-FITC



Fig. 5. Cell apoptosis was detected by Flow cytometry histograms of cell death assays by the PI method in MCF-7 cells. The cells were treated with CTII and then evaluated for cell cycle distribution at 24 h. Cells were categorized as necrotic (Q1), late apoptotic (Q2), viable (Q3), and early apoptotic cells (Q4). A: untreated group, B: treated group C: early and late apoptotic phase. ****p < 0.0001 were considered as significant levels compared to the untreated control group.

respectively) was assessed and showed significant down-regulation in the treated group compared with control which may indicate a reduction of migration ability of the cells in this cancer cell line. The results have shown in Fig. 8.

4. Discussion

BC is the most frequent cause of cancer death in women [24]. Like the other cancers, there are some options for BC cancer therapy. Several compounds of venomous species have generally shown anti-cancer pharmacological properties [32]. Researchers indicated cytotoxic effects of snake venom can diminish and damage tumor cells. Kim et al. showed the L-amino acid oxidase which was isolated from the venom of the king cobra, *Ophiophagus hannah* can inhibit cell proliferation in fibroblastic sarcoma, ovarian cancer, colorectal cancer, stomach cancer, and murine melanoma cell lines according to [3H] thymidine uptake approach [33]. The function of this enzyme can be linked to thymidine incorporation blockade and DNA interaction [33]. Zhang et al. extracted ACTX-6 protein, which has 98 kDa Mw that could enhance the Hela cells' apoptosis [34]. The isolated cardiotoxic-cytotoxic protein with 6.76 kDa Mw from the monocled cobra (Naja kaouthia) significantly, could inhibit human leukemic U937 and K562 cells growth in a dose- and time-dependent fashion [35]. In another research, NN-32 which was a purified venom from Indian Naja naja through ion-exchange chromatography displayed cytotoxic effects on Ehrlich ascites carcinoma (EAC) mice, which decreased the tumor size and enhanced the survival of animals [36]. The Caspian cobra (Naja naja oxiana), also known as the Central Asian cobra or Oxus cobra, is a highly venomous cobra species found in Central Asia within the Elapidae family. Naja naja oxiana venom has an anticancer role in many different types of cell lines [19-37]. Researchers showed that the IC₅₀ values of Caspian cobra venom were 26.59, 21.17, and 47.1 μ g mL⁻¹ for the cell lines HepG2, DU145, and MDCK respectively [37]. The concept of this research came back to Ebrahim et al. researches which investigated the potential anticancer of native CTII in



Fig. 6. Flow cytometric analysis of cell death. rCTII decreased the number of cells in the S phase and increased the number of cells in the sub-G1 phase (apoptotic cells). (A) Untreated group. (B) Treated group.



Fig. 7. Treated and untreated MCF-7 cells were subjected to scratch wound-healing assays. MCF-7 cells divided into 2 groups including treated and untreated. The wound space was photographed at 0, 12, and 24 h and the migration of cells to the wound sides was evaluated and statistically analyzed. Error bars indicate the mean \pm SD of independent experiments.

the MCF-7 cell line [24]. The main purpose of this study was to optimize a process for large scale production of CTII and compare this recombinant protein with the native form.

Given the broad potential usage of CTII, alternative, cost-effective, and high-yield production systems are advantageous. Active recombinant toxin preparation using heterologous gene expression systems seems to be the only method of obtaining large quantities of toxins. By using recombinant methods, we can easily express heterologous genes in pro and/or eukaryotic hosts, i.e. genes that they normally do not express. In this regard, the E. coli is one of the important hosts of recombinant protein production due to its rapid growth, simple handling, and cost-effectiveness [26]. In the present study, for the first time, we successfully transformed the pET-SUMO-CTII vector into SHuffle® T7 Express and effectively expressed rCTII in this bacterium as shown in Fig. 2. The SUMO tag can be cleaved from the fusion protein via SUMO-protease, which specifically identifies the SUMO tertiary structure, to generate the native protein without any redundant amino acids. We purified and then cleaved the rCTII and considered by using SDS-PAGE and immune blotting assays (Fig. 3). As sown in Fig. 2 the MW of the current protein was 6.6 kDa.

In spite of the many studies which were conducted on CTII, there is no previous report of CTII secondary structure. In this regard, after the recombinant production and purification of rCTII, biophysical characterization was conducted using CD spectroscopy. This technique is a powerful tool for the fast determination of the secondary structure, folding, and binding features of the proteins [40]. The content of different secondary structures of rCTII was determined via CDNN software (Fig. 4), and it contains 6.106% α -helix, 35.87% antiparallel β -sheet, 5.15% parallel β -sheet, 18.7% β -turn, and 34.16% random coil. This novel information would lead to a better understanding of CTII specifications.

Multiple physiological and biological processes involved in the appropriate cell development processes and homeostasis-pathogenesis and many diseases associated with the impaired apoptosis process. Apoptosis has an important impact on maintaining the balance among cell death and division and avoiding apoptosis contributes to excessive cell proliferation leading to various diseases, including many types of cancers. Due to minimal inflammation that occurs during apoptosis of the cells, the establishment of anticancer drugs to improve apoptosis has



Fig. 8. The mRNA expression of candidate genes. The expression of P53, Bcl2, Bax, caspase-3, caspase-8, caspase-9, and caspase-10 was assessed by real-time PCR in the Control and treated group. Our result illustrated this protein potentially activated both intrinsic as well as extrinsic pathways related to apoptosis. Error bars display the \pm SD. The independent *t*-tests assessed the *p*-values. *p* < 0.001 (***), *p* < 0.001(****).

received more interest in cancer research [41]. In the present research, we applied different techniques for the detection of molecular pathways, which involved in apoptosis in MCF-7 cell line after treatment with $4 \mu g m L^{-1}$ of rCTII. We chose this concentration based on previous research on the natural type of CTII in MCF-7 cells [24]. Annexin V/PI staining indicated a significant translocation of phosphatidylserine (PS) from the inside of cancer cells to the outside after exposure to rCTII. The rCTII increased the early apoptosis in the treated cells in comparison with control cells. In addition, the increasing rate in the late apoptosis ratio has been shown in the treated cells in comparison with controls (Fig. 5). The cell cycle progression analysis showed a reduction of S phase and increasing proportions of cells in the sub-G1 group (apoptotic cells). Our results confirmed the previous study which showed that ~88% of the cells treated with rCTII were in the sub-G1 phase [24]. In this study, our recombinant CTII showed more effectiveness in comparison to the isolated extract from the venom source in the $4 \mu g m L^{-1}$ concentration. As indicated by many researchers, CTII has potent anti-cancer effects in the MCF-7 cell line, which are induced via the intrinsic apoptosis pathways [24]. More recently, Attarde et al. purified NN-32, a very similar toxin to CTII amino acid sequences from Naja naja venom with cytotoxicity versus MCF-7 and MDA-MB-231 cells and the IC₅₀ values were 2.5 and 6.7 μ g mL⁻¹ respectively [25]. Yang et al. which isolated Cardiotoxin-3 (CTX-3), from Naja naja atra venom, showed the induced apoptosis of K562 cells occurred through up-regulating Bax and endonuclease G and downregulating of Bcl-X [42]. The other study also confirmed LAAO isolated from Agkistrodon acutus snake venom arrests cancer cells during the sub-G1 phase of the cell cycle and induces apoptosis via the Fas signaling cascade in A549 cells [25]. The process of apoptosis through the mitochondrial pathway induces loss of mitochondrial membrane potential (MMP), the release of cytochrome *c*, and activation of caspases genes. Our study showed the expression of P53, Bax, caspase-3, caspase-8, caspase-9, and caspase-10 significantly upregulated after treatment with rCTII. Although we showed the down-regulation of Bcl2 in the treated group in compression with the controls. These findings illustrate the capability of rCTII to activate the intrinsic and extrinsic pathways of apoptosis simultaneously. The previous study confirmed our finding that showed CTII extracted from native form up-regulated the caspase-9 in MCF-7 cells after treatment and their observations confirmed the possibility that apoptosis was mediated by the intrinsic pathway [24]. The other research on NKCT1 toxin from *Naja Koauthia* venom showed that after conjugation with gold nanoparticles, it shows anticancer effects in EAC cells. GNP-NKCT1 induced caspase-dependent apoptosis in EAC cells, induced a late apoptotic stage, and arrested the cells in the G0/G1 phase [24,43].

Metastasis is a multi-step mechanism that contains the movement and invasion of cancer cells and defines as a signature for malignancy. The next step in our study was the assessment of the potential effects of rCTII on the migration of MCF-7 cells. Migration was evaluated by wound healing assay, which is the complex, dynamic movement of the cells. Observation of the motility of living cells is an important tool for calculating the rate of movement to the area produced by the initial cut. The ideal condition is when the wound gap filled slowly by the cells over a prolonged time, which ensures that the investigated agent prevents the movement of cancer cells. Fig. 7 indicates the effects of rCTII on the wound healing assay. In untreated cells, the migration was very dynamic after only 12 and 24 h, but the wound's closure was completed after 24 h. By using a 4 μ g mL⁻¹ of rCTII, the motility of the MCF-7 cells was inhibited in 12 and 24 h. Furthermore, it is well understood that MMPs are essential to the process of cancer cell movement [24,37]. Therefore, after the scratch test, we investigated to understand if our rCTII had similar effects on MMP-3 and MMP-9 expression. The findings indicate that this protein down-regulated the expression of MMP-3 and MMP-9 in the breast cancer cell line at the mRNA level and confirmed our scratch test results.

5. Conclusion

In this novel in vitro study, the recombinant form of CTII successfully produced and it's bioactivity against MCF-7 cell line confirmed by different methods. The results of this study indicate that rCTII has anti-proliferative effects and can be considered as a promising candidate for breast cancer therapy. More studies particularly on animal models or through clinical trials are needed to prove our findings.

Author contributions

Afshin Derakhshani the first author of the manuscript, design the project, perform the experiment, analyzed the data, and write the initial version of the manuscript. Nicola Silvestris, Khalil HajiAsgharzadeh, and Sara Mahmoudzadeh participate in analyzing the data and completing the work. Mohammad Fereidouni, Angelo Virgilio Paradiso, and Oronzo Brunetti read the manuscript and contributing in the English editing of the manuscript, and also helping in data categorization. Deyhim Atarod analyzed the CD spectroscopy data. Behzad Baradaran and Hossein Safarpour: the corresponding authors of the manuscript, financially support the work and supervise the project and also contributing to the revising of the main text of the manuscript.

Declaration of competing interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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