The Effect of Indocyanine Green Antimicrobial Photothermal/Photodynamic Therapy on the Expression of BCL-2 and BAX Messenger RNA Levels in Human Gingival Fibroblast Cells

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Abstract

Background: Antimicrobial photothermal/photodynamic therapy (PTT/PDT) with indocyanine green (ICG) is an adjuvant therapeutic approach in the treatment of periodontitis. To explore whether PTT/PDT with ICG causes cell death by apoptosis in human gingival fibroblast (HGF) cells, BAX and BCL-2 genes expression as key events for apoptosis were evaluated in this study.

Materials and methods: HGF cells were treated with: 1) different concentrations (500–2000 µg/mL) of ICG alone, 2) Diode laser irradiation alone with a fluency of 39.06 J/cm²; 3) PTT/PDT combined different concentrations (500–2000 µg/mL) of ICG with an 808 nm diode laser with a fluency of 39.06 J/cm², and 4) controls (untreated cells). After that, BAX and BCL-2 messenger RNA levels were evaluated by real-time quantitative reverse transcription PCR.

Results: PTT/PDT with 500 µg/mL of ICG caused significant increases in the expression of the BAX gene, with an 8.5-fold increase, which was approximately 7- and 8.5-fold higher than PTT/PDT with ICG for 1500 and 2000 µg/mL of ICG, respectively, indicating induction of apoptosis in HGF cells. ICG (in different test concentrations), diode laser, and PTT/PDT with ICG (1500 and 2000 µg/mL of ICG) treatment displayed insignificant increases in expression levels of BAX (all p>0.05). Our experiments showed an insignificant increase (1.1–1.6-fold) in the expression of BCL-2 after ICG, diode laser, and PTT/PDT with ICG treatment (all p>0.05).

Conclusions: This study suggests that various concentration of ICG can be the diverse expression of BAX responses to PTT/PDT on HGF cells.

Keywords: antimicrobial photothermal/photodynamic therapy, apoptosis, human gingival fibroblast cells, gene expression, indocyanine green, periodontitis, qRT-PCR

BACKGROUND

Periodontitis is a multi-factorial inflammatory illness of the tooth-supporting structures and is associated with subgingival bacterial biofilm formation; it is one of the major reasons of tooth loss. Recently, the pathogenesis of periodontitis diseases has been explained by the alteration of normal apoptosis regulation. Apoptosis is a constitutive regulated...
physiological process essential for the maintenance of cellular homeostasis in multicellular organisms that can be induced or inhibited by different stimuli.\(^2\)\(^4\) It is also clear that *Aggregatibacter actinomycetemcomitans* and *Porphyromonas gingivalis* as periopathogen bacteria can induce significant apoptosis in monocytes and primary human gingival epithelial cells, respectively.\(^5\)\(^6\) Microbial by-products including hydrogen sulfide and proteases can induce apoptosis in gingival epithelial cells and human gingival fibroblasts in the oral cavity, which may cause periodontal diseases.\(^7\)\(^8\)

There are different treatment strategies for periodontitis, including surgical and non-surgical treatment. Antimicrobial photothermal/photodynamic therapy (PTT/PDT) is an adjuvant non-surgical approach used in the treatment of periodontal diseases.\(^9\)\(^10\) In most studies, antimicrobial PTT/PDT is a beneficial adjunct for scaling and root planing (SRP), especially in chronic periodontitis treatment due to its ability to eliminate bacterial biofilms.\(^11\)\(^13\) Combining antimicrobial PTT/PDT and SRP is beneficial, especially in lesions with untoward anatomic conditions.\(^14\) Antimicrobial PTT/PDT as monotherapy shows advantages in cytokine modulation and is used as a non-invasive therapeutic method for the treatment of residual pockets and periodontal disease.\(^15\) In antimicrobial PDT (aPDT), electron transfer in excited photosensitizers (PSs) can be done in two ways: Type I photoreaction which produces radicals and/or radical ions from substrates including highly reactive hydroxyl radicals (HO•) and Type II photoreaction which produces singlet oxygen (\(^1\)O\(_2\)).\(^16\)\(^17\)

The efficiency of antimicrobial PTT/PDT is multifactorial and varies by the concentration and type of PS, and irradiation time and energy dose of the light. Different factors make PS ideal for aPDT, including high interaction with microbial cells and the optical window for sufficient tissue penetration of light.\(^18\) More tissue penetration of 810 nm wavelength, as a near-infrared (NIR) diode laser system, then 635 nm and 660 nm, as the most common wavelength used in aPDT, has been addressed in recent studies.\(^18\)\(^19\) Among the PSs that are active in this spectrum (i.e., 808 nm) can be referred to as indocyanine green (ICG). ICG is a member of the amphiphilic-tricarbocyanine dye family with the absorption peak around 808±5 nm, which is thought to be an effective PS against Gram-negative and Gram-positive bacteria. Although photo-activated ICG is known to yield \(^1\)O\(_2\), it has been shown that photothermal (PTT) destruction of microbial cells was the dominant reaction.\(^20\)\(^21\) ICG has also received Food and Drug Administration (FDA) approval to be used in clinical settings.\(^19\)

The success of ICG antimicrobial PTT/PDT as a periodontal disease treatment not only is dependent on the effective decontamination of microorganisms but also via avoiding an unacceptable degree of harm to healthy periodontal tissue due to induction of apoptosis.\(^22\)\(^23\) BAX (Bcl-2-associated X protein) and BCL-2 (B-cell lymphoma 2) are two members of the BCL-2 family that play an important role in apoptotic cell death. BAX (pro-apoptotic) has been shown to promote cytochrome c release, which in turn leads to activation of caspases and degradation of specific protein substrates, whereas BCL-2 as an antiapoptotic protein blocks BAX-induced cytochrome c release and caspase activation, which can result in induction and inhibition of apoptotic cell death, respectively.\(^24\) Thus, it has been suggested that the relative amounts of BAX and BCL-2 regulate the outcome of a living cell.\(^25\) Because apoptosis has an extensive role in periodontal diseases, knowledge about the apoptosis response to ICG antimicrobial PTT/PDT in human gingival fibroblast (HGF) cells is necessary for its therapeutic intervention at different checkpoints. In this study, we determined the apoptosis response to PTT/PDT with ICG in HGF cells in vitro via evaluation of the expression of the apoptosis-related genes BAX and BCL-2 in HGF cells exposed to PTT/PDT with ICG. In this study, we hypothesize that there is a considerably significant difference between the expression of BAX and BCL-2 in HGF cells exposed to photo-activated ICG in different concentration. Under the null hypothesis, this difference is insignificant.

**MATERIALS AND METHODS**

**Photosensitizer**

A stock solution from ICG (Serva, Heidelberg), at concentrations of 4000 µg/mL was made by dissolving the ICG powder in a sterile balanced salt solution (BSS) at pH 7.4. The ICG solution was sterilized using a 0.22 µm syringe filter, and stored at 4°C in the dark before use.\(^24\)

**Light source**

An 808 nm diode laser (Konftec, DX82, Taiwan) with power output 250 mW was employed in this study. The output power density of the diode laser was determined by a power meter (Laser Point s.r.l., Milan, Italy).

**Cell culture**

Human gingival fibroblast (HGF; IBRC C10459) cells were purchased from the Iranian Biological Resource Center (Tehran, Iran). The cells were grown in nutrient medium composed of Dulbecco’s Modified Eagle’s Medium (MDME; Gibco, USA) supplemented with 10% fetal bovine serum (FBS; Sigma, USA), 1% penicillin-streptomycin solution (10,000 Unit/mL penicillin and 10 mg/mL streptomycin), 2 mM L-glutamine and 100 µg/mL of amphotericin B. Cells were incubated at 37°C in a humidified atmosphere of air (95%) and 5% CO\(_2\). Every 3 days the medium was changed. When the cultures reached 80% confluence, the cells were detached from the flasks by using 0.25% trypsin–0.02% ethylene diamine tetraacetic acid (EDTA) solution. For experiments requiring seeding of HGF cells in microplates,
in the third passages, a density of 2×10⁶ HGF cells/well were seeded in flat-bottom 96-well cell culture microplates (Greiner Bio-One, Germany).²⁵

Design of study

To determine the apoptosis response to an 808 nm diode laser, ICG and PTT/PDT with ICG in HGF cells in vitro via evaluation of apoptosis-related genes BAX and BCL-2 expression in treated HGF cells, test groups consisted of HGF cells subjected to:

1) ICG alone (in three concentrations 2000, 1500, and 500 µg/mL; L− P−; L and P stand for "light" and "photosensitizer," respectively),
2) Diode laser alone with fluency of 39.06 J/cm² (L+ P−)
3) PTT/PDT with ICG; combined ICG (in three concentrations 2000, 1500, and 500 µg/mL) with 808 nm diode laser treatment with a fluency of 39.06 J/cm² (L+ P+)
4) Control group (no exposure to either diode laser irradiation or ICG; L− P−)

PTT/PDT procedure

The HGF cells (100 µL; at a final density of 2×10⁶ cells per well) were placed in each well of a sterile flat-bottom 96-well cell culture microplate. After HGF cells were grown until full confluence, the ICG (100 µL) was added for groups L− P− and L+ P+; whereas sterile DMEM (100 µL) was added for groups L− P− and L+ P−. In the pre-irradiation time phase, the microplates were incubated for 5 min in a cell culture incubator. The microplate wells contents of groups L+ P+ and L+ P− were exposed to 808 nm diode laser for 60 s (39.06 J/cm²) in continuous mode at room temperature (25±2°C). The tip diameter was 7 mm and used in non-initiated mode. The probe of the laser was fixed 1 mm above the top surface of microplate by a microscope stand the total energy was 15 J. The microplate wells around the test wells of the microplates were filled with Indian ink to inhibit light emission to neighboring wells.¹⁴,¹⁵ To avoid beam reflection from the table top during PTT/PDT with ICG, a sheet of black paper were used under the plates. The microplate wells in the control group did not receive any treatment. Next, cells in the microplate wells were scraped with a sterile tip of a pipette and transferred into 1.5-mL Eppendorf tubes, and three volumes of RNAlater® (Thermo Fisher Scientific, US) were then added to stabilize the RNA of HGF cells. The RNA of HGF cell suspensions were then extracted immediately.²⁵

RNA extraction

Collected HGF cells were centrifuged (13000 g×10 min) and the pellet was washed twice with phosphate-buffered saline (PBS) (500 µL; pH 7.4). After the final centrifugation, the pellet of HGF cells was subjected to RNA extraction. For this purpose, the Hybri-aidR™ Total RNA Purification Kit (GeneAll Biotechnology, Seoul, Korea) was used following the manufacturer’s recommended protocol.

Assays for RNA integrity

Agarose gel electrophoresis was used to assess the integrity of the total extracted RNA. An aliquot of the RNA sample was run on a denaturing agarose gel stained with ethidium bromide (EB). The 18S and 28S ribosomal RNA (rRNA) bands were visualized by EB staining.

Quantitation and evaluation of purity of extracted RNA

Ultraviolet (UV) spectroscopy using the NanoDrop® spectrophotometer (Thermo Fisher Scientific, US) at 260 nm and by measuring absorbance ratios at 260 vs 280 nm was used for quantification and evaluation of the purity of extracted RNA, respectively. Alternatively, to determine the purity of extracted RNA, we performed PCR using DNAasel treated RNA as a template. PCR amplification was performed in a final volume of 25 µL containing 2 µL of templates, 9 µL of PCR master mix (Amplicon, Denmark), and 2.5 pM of each GAPDH forward and reverse primers. PCR was carried out in a thermal cycler apparatus (Pqstar; PqLab, Germany) with an initial denaturation step (5 min at 95°C), followed by 35 cycles including denaturation (1 min at 95°C), annealing (30 s at 59°C), and extension (30 s at 72°C), with a final extension step (5 min at 72°C). The results were analyzed by 2% agarose gel electrophoresis in intact RNA with sharp, clear 28S- and 18S rRNA bands (Fig. 1). As shown in Figure 1, the 28S RNA band was approximately twice as intense as the 18S rRNA band. The 2:1 ratio of band intensity of 28S rRNA: 18S rRNA was an accepted indication of RNA Integrity. The results of this study showed that the RNA extracted from HGF cells ranged from 314.5

![Figure 1. A 2% agarose gel electrophoresis of total RNA extraction from HGF cells. Lanes 1 and 3 represent samples collected before and after PTT/PDT with ICG, respectively. Lane 2 represents the DNA marker 1 kb. Five microliters of RNA were loaded for each sample.](image-url)
ng/μL to 560 ng/μL with a mean yield of 412.5 ng/mL (Fig. 2). On purity assessment of RNA extraction, we found that all samples were within the optimal range of ~2.0. The amplification of GAPDH by PCR yielded no bands in DNase I treated extracted RNA, indicating the purity of DNase I-treated RNA as described in Section 2.3.2. As shown in Fig. 3, the amplification of GAPDH by PCR yielded a band in cDNA as the template, indicating the successful reverse transcription of total extracted RNA into a cDNA.

**Primers**

All primers used in this study i.e., GAPDH forward primer, 5′-GGG TCT CTC CTC CTC CTG TT-3′ and GAPDH reverse primer, 5′-ACG ACC AAA TCC GTT GAC TCC-3′; BAX forward primer, 5′-TTC TGA CGG CAA CTT CAA CTG-3′ and BAX reverse primer, 5′-AGG AAG TCC AAT GTC CAG CC-3′; and BCL-2 forward primer, 5′-AGG CTG GGA TGC CTT TGT GG-3′ and BCL-2 reverse primer, 5′-GGG CAG GCA TGT TGA CTT CAC-3′ were optimized to an equal melting temperature of 59°C. BAX primers were used from a previous study and GAPDH, as well as BCL-2 primers, were designed via gene runner software version 3.05 (Hastings Software Inc. Hastings, NY, USA). To ensure the specificity of GAPDH and BCL-2 primers, their sequences were searched against GenBank sequences with the BLAST program.

**Determination of specificity of primers**

To determine the specificity of the primers, the generation of a single amplicon of the correct size from each prim-
er was evaluated on agarose gel electrophoresis. For this assessment, cDNA was subjected to PCR amplification as mentioned above (Section 2.3.2). Alternatively, Real-Time PCR methodologies utilizing SYBR Green dye (qRT-PCR Master Mix; Takara, Japan) was used to analyze melt curves to confirm the specificity of the amplified product. For this purpose, a default melting program was run on a real-time PCR machine (Bioer Technology, Hangzhou, China) at the end of the cycling program. The amplification of GAPDH, BAX, and BCL-2 by PCR yielded no false-positive bands in negative controls and unspecific bands in test samples (Fig. 4). The agarose gels of the amplified product revealed single bands corresponding to the predicted amplicon length. As shown in Fig. 5, dissociation melting curves of GAPDH, BAX, and BCL-2 amplicons with a single peak demonstrates the specificity of the primer pairs.

**Quantitative Real-time PCR**

For the real-time reaction, a master mix was prepared as follows: 7 µL water, 10 µL SYBR green PCR Master Mix, 1 µL cDNA as a PCR template, and 2 µL volume of primers. A two-step experimental run protocol was used: 1) initial denaturation program (5 min at 95°C) and 2) 35 cycles of 15 s at 95°C, 15 s at 59°C, and 15 s at 72°C. Each PCR reaction was completed in triplicate. In this study, GAPDH used as a housekeeping gene for normalizing of test RNA expression.

**Statistical analysis**

The data are presented as mean ± standard deviation (SD) from three experiments. Fold differences in RNA expression were determined by the $2^{-\Delta\Delta C_t}$ method. Changes in RNA expression greater than or equal to twofold were considered significant.

**RESULTS**

To examine whether the expression of BAX and BCL-2 genes changed after diode laser (L+ P−), ICG (L− P+), or PTT/PDT with ICG (L+ P+) treatment, we compared the relative gene-specific messenger RNA quantities of HGF cells under-treated versus control (untreated; L− P−). The relative change in transcript expression levels of the BAX and BCL-2 genes among the treated HGF cells are shown as mean ± SD relative change over basal levels (Fig. 6).

Among treated HGF cells, PTT/PDT with ICG in 500 µg/mL of ICG with a fluency of 39.06 J/cm$^2$ revealed a significant increase in the expression of the BAX gene, with a 8.5-fold rise, which was approximately 7- and 8.5-fold higher than PTT/PDT with ICG in 1500 and 2000 µg/mL of ICG with a fluency of 39.06 J/cm$^2$, respectively (Fig. 6). According to Fig. 6, PTT/PDT with ICG treatment using 1500 and 2000 µg/mL at fluency of 39.06 J/cm$^2$ displayed an insignificant increase in expression levels of the BAX and BCL-2 genes compared with untreated HGF cells (control group; all $p>0.05$). As Figure 6 reveals, our PTT/
PTT/PDT with ICG experiments showed that an insignificant increase (1.1-fold) in the expression of the BCL-2 gene was observed after exposure to ICG (500 µg/mL) plus diode laser at fluency of 39.06 J/cm² (p>0.05) compared with the control group.

As shown in Fig. 6, the expression of the BAX and BCL-2 genes were upregulated 1.4- and 1.3-fold, respectively, following diode laser irradiation alone with a fluency of 39.06 J/cm² compared with the control group (both p>0.05). There was no remarkable difference in expression of the BAX and BCL-2 genes following ICG treatment at concentrations of 500–2000 µg/mL (all relative fold change < 2; all p>0.05).

**DISCUSSION**

The PTT/PDT is an adjuvant treatment approach and has gained approval for the treatment of various diseases such as local malignant and nonmalignant infectious diseases. Numerical previous studies have also indicated the potential ICG antimicrobial role of PTT/PDT; studies in *Staphylococcus aureus, Acinetobacter baumannii, A. actinomycetemcomitans*, and *P. gingivalis* demonstrated that aPDT reduced microbial pathogenicity, suggesting that aPDT is an adjuvant treatment approach in microbial-associated diseases such as periodontitis. Several factors including the concentration of PS are involved in the success of aPDT. PS concentration also plays an important role in cell cytotoxicity. To further investigate the potential associations of the effect of the ICG concentration in PTT/PDT on HGF cells and the BCL-2/BAX signaling pathway, the present study analyzed the effect of PTT/PDT on HGF cells based on different concentration of ICG.

ICG as a new PS is used in several therapeutic fields such as ocular tumor therapy, management of retinal disease, as well as treatment of local infections and inflammatory disease, owing to its outstanding achievements and clinically brilliant properties that are sufficiently suitable for treatment. However, ICG has a low cytotoxicity, some studies have shown its toxic effects when used locally. Ho et al. exposed ARPE-19 (human retinal pigment epitheli-
The power output of the current irradiation setting was conducted with the average laser power density of 250 mW and the duration of exposure of 39.06 J/cm², which were much lower than those which had a cytotoxic effect as a result of detectable temperature increase in cells or tissues within 60 s after administration.

In agreement with the study of Pourhajibagher et al.44 the current study demonstrated that PTT/PDT with ICG in 500 μg/mL of ICG increased expression of the proapoptotic gene, BAX, in HGF cells. In this case, in line with the other studies,46-49 it suggests cytochrome c-dependent apoptosis of the HGF cell line occurred through downregulation of the BCL2/BAX ratio, which may be stimulating mitochondrial depolarization and resulting in apoptosis. Previous studies support our findings that a reduction in BCL-2 levels inhibits cell proliferation and promotes apoptosis.50 It has revealed that BCL-2 is predominantly localized in the mitochondria. The major biological function of homodimers of BAX is to promote apoptosis, while homodimers of BCL-2 inhibit apoptosis. When a BAX/BCL-2 heterodimer is formed, the anti-apoptotic function of BCL-2 is inhibited by BAX, resulting in the promotion of apoptosis.51

It has been observed that mRNA synthesis and translational control are important in the induction of p53 protein levels in response to apoptotic cues. p53 mRNA contains the internal ribosome entry site (IRES) element, which aids in protecting the mRNA from microRNAs induced silencing. IRES prepares for a selective translation of the specific mRNAs in periods of attenuation of cap-dependent translation by bypassing the requirement for the factors that are subject to inhibition during apoptosis. Therefore, mRNA translational regulation via the IRES is the main mechanism by which p53 can respond to the different apoptotic triggers the cell is exposed to.52

It has been revealed that singlet molecular oxygen (1O₂) which produced under the photodynamic action of PSs is a potent inducer of apoptosis. Cleavage of caspase-3 and its target proteins such as lamin B and poly (ADP ribose) polymerase, a signature event of apoptosis, can be induced by the singlet oxygen.53

Along with the recent study that showed PTT/PDT with ICG has been demonstrated to be non-cytotoxic against HGF cells, in 1000–2000 μg/mL of ICG with fluency of 39.06 J/cm², 44 the current study demonstrated that PTT/PDT with ICG does not significantly increase BAX gene expression in HGF cells at or above a 1500 μg/mL concentration of ICG with an irradiation time of 60 s. The data collected here demonstrated that PTT/PDT with ICG at high concentrations of ICG inhibited the BCL-2/BAX signaling pathway. These data are supported by a previous study that revealed PTT/PDT with ICG at 1000 μg/mL concentrations of ICG inhibited this signaling pathway.25

Pourhajibagher et al.44 in their recent investigations observed a reduction in the viability of cells when exposed to a light source alone. Nowak et al.37 found that light exposure can produce thermal injury in RPE cells, whereas exposure to light for longer periods can lead to photochemical damage of treated cells. Contrary to the above studies, according to our results no significant increase in the expression of BAX and BCL-2 genes was observed in the treatment group with only a diode laser when compared to controls.

This is the first study that shows the phototoxicity mechanism of ICG during irradiation by a laser. It has been shown that the major mechanism after ICG excitation by diode laser is the transformation of 88% of energy to heat inside the ICG molecules for PTT. The residual energy can generate singlet oxygen for PDT. So, the mechanism of action is more PTT rather than photodynamic action.54,55

We hypothesize that photo-activated ICG can be hazardous to HGF cells due to its photothermal activity and on a smaller scale 1O₂-mediated DNA and membrane damage, which possibly leads to induction of apoptosis and ultimately cell death. Therefore, PTT/PDT with ICG as an adjuvant treatment should avoid using a low concentration of ICG (below 1000 μg/mL) and as much as possible, the intervention location should be kept dry, which can prevent dilution of ICG. In the present study, we found that ICG in lower concentrations had a photo-enhancing effect on the expression levels of BAX in HGF cells. High concentrations (1500 and 2000 μg/mL) of ICG, that did not elicit a thermal cytotoxic effect, also achieved insignificant increases in photo-enhancing effect in expression levels of BAX after irradiation. Although lower-temperature light-induced hyperthermia (<42°C) is identified to be synergistic with PDT56, the power intensity used in PDT treatment in this study was restricted explicitly in order to avoid significant tissue heating so that the photochemical reactions were dominant and the thermal effects did not alter the effective dose administered. Thus, the concentration of ICG should be a primary concern for the management of local infection and human health. Therefore, the optimum concentration of ICG for PTT/PDT should be at least 1000 μg/mL with a 60 s irradiation time. Further studies are needed to confirm our data and functional studies are necessary to establish a possible causal relationship between PTT/PDT with ICG and apoptosis.

CONCLUSIONS

Our results demonstrated that the PTT/PDT with ICG at low concentration of ICG had a positive effect on BAX gene expression, resulting in induction of apoptosis in HGF cells induced by the PTT/PDT procedure. The concentration of ICG applied should be considered as a crucial point for PTT/PDT as an adjuvant therapeutic approach. It is recommended to use ICG not only the best ICG antimicrobial effect in PTT/PDT, but also less effect on the induction of apoptosis in human cells with a concentration at or above 1000 μg/mL.
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Conflict of Interest

The authors declare that there is no conflict of interests regarding the publication of this paper.

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Влияние антимикробной фототермической / фотодинамической терапии индоцианиновым зелёным на экспрессию уровней BCL-2 и BAX в матричной РНК в клетках фибробластов десны человека

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Резюме
Введение: Антимикробная фототермическая / фотодинамическая терапия (ФТТ/ФДТ) с индоцианиновым зелёным (ИЦЗ) является адъювантным терапевтическим подходом в лечении периодонтита. Чтобы выяснить, вызывает ли ИЦЗ -индукция клеток в результате апоптоза в клетках фибробластов десны человека (КФДЧ), в этом исследовании оценивали экспрессию генов BCL-2 и BAX в качестве ключевых событий для апоптоза.

Материалы и методы: Клетки КФДЧ обрабатывали 1) различными концентрациями (500–2000 µg/mL) только ИЦЗ, 2) облучением диодным лазером с плотностью 39,06 J/cm²; 3) ФТТ/ФДТ, объединённые в различных концентрациях (500–2000 500–2000 µg/mL) ИЦЗ с диодным лазером 808 нм с плотностью 39,06 J/cm², и 4) контроли (необработанные клетки). Затем уровни BCL-2 и BAX в мРНК измеряли количественной ПЦР с обратной транскриптазой в реальном времени.

Результаты: ФТТ/ФДТ с 500 µg/mL ИЦЗ привели к значительному увеличению экспрессии гена BAX в 8,5 раз, что было примерно в 7 и 8,5 раз выше, чем соответственно ФТТ/ФДТ с 1500 и 2000 µg/mL ИЦЗ, который является индикатором индукции апоптоза в клетках КФДЧ. Обработка ИЦЗ (в различных тестовых концентрациях), диодным лазером и ФТТ/ФДТ с ИЦЗ (1500 и 2000 µg/mL ИЦЗ) показала значительное увеличение уровней экспрессии I – V (все p> 0,05). Наш эксперимент показал значительное увеличение (в 1,1–1,6 раза) экспрессии BCL-2 после лечения с помощью ИЦЗ , диодного лазера и ФТТ/ФДТ с ИЦЗ (все p> 0,05).

Выводы: Это исследование показало, что различные концентрации ИЦЗ могут быть разнообразной экспрессией ответов BAX на ФТТ/ФДТ клеток КФДЧ.

Ключевые слова
антимикробная фототермическая / фотодинамическая терапия, апоптоз, клетки фибробластов десны человека, экспрессия генов, индоцианиновый зелёный, периодонтит, qRT-PCR