Apoptotic Effect of co-treatment with HS-1200 and Cisplatin on SCC25 Human Tongue Squamous Cell Carcinoma Cell Line

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Bile acids are polar derivatives of cholesterol essential for the absorption of dietary lipids and regulate the transcription of genes that control cholesterol homeostasis. Recently it have been identified the synthetic chenodeoxycholic acid (CDCA) derivatives HS-1200 and cisplatin showed apoptosis-inducing activity on various cancer cells in vitro. This study was undertaken to investigate the synergistic apoptotic effect of co-treatment with HS-1200 and cisplatin on human tongue squamous cell carcinoma cells (SCC25 cells).

To investigate whether the co-treatment with HS-1200 and cisplatin compared to each single treatment efficiently reduces the viability of SCC25 cells, MTT assay was conducted. The induction and augmentation of apoptosis were confirmed by DNA electrophoresis, Hoechst staining and an analysis DNA hypoploidy. Western blot analysis and immunofluorescent staining were also performed to evaluate the expression levels and the translocation of apoptosis-related proteins following this co-treatment. Furthermore, proteasome activity and mitochondrial membrane potential (MMP) change were also assayed.

In this study, co-treatment with HS-1200 and cisplatin on SCC25 cells showed several lines of apoptotic manifestation such as nuclear condensations, DNA fragmentation, reduction of MMP and proteasome activity, the increase of Bax and the decrease of Bcl-2, decrease of DNA content, the release of cytochrome c into cytosol, translocation of AIF and DFF40 (CAD) onto nuclei, and activation of caspase-9, caspase-7, caspase-3, PARP and DFF45 (ICAD) whereas each single treated SCC25 cells did not show these patterns.

Although the single treatment of 25 μM HS-1200 and 4 μg/ml cisplatin for 24 h did not induce apoptosis, the co-treatment of these reagents prominently induced apoptosis. Therefore our data provide the possibility that the combination therapy with HS-1200 and cisplatin could be considered as a novel therapeutic strategy for human squamous cell carcinoma.

Key words: Apoptosis, Bile acid, HS-1200, Cisplatin, Human tongue squamous carcinoma
I. INTRODUCTION

Cells undergoing apoptosis usually develop characteristic changes, including nuclear condensation and degradation of DNA into oligonucleosomal fragments.\(^1\) Apoptotic cell death is thought to result ultimately from the proteolytic actions of caspase \(^2\) and alterations in mitochondrial function play a key role in the regulation of apoptosis.\(^3\) Moreover, the proteasome system has been shown to be implicated as a negative or positive mediator of apoptosis. The proteasome pathway is known to work mostly in upstream of mitochondrial alterations and caspase activation.\(^4\)

Carcinoma of the oral cavity, especially oral squamous cell carcinoma (OSCC), are one of the most leading causes of cancer related death and affect nearly 500,000 patients annually world-wide. And OSCC is one of the most malignancies that remain incurable with current therapies.\(^5\) OSCC patients are treated by classical modalities of treatment consisting of surgery, radiotherapy, and/or chemotherapy. But OSCC still shows noticeable mortality rates, therefore new therapeutic approaches have been investigated and the most promising one is obtained from natural agents with known anti-cancer effects.

Bile acids are polar derivatives of cholesterol essential for the absorption of dietary lipids and regulate the transcription of genes that control cholesterol homeostasis. Different bile acids exhibit distinct biological effects. Importantly, natural bile salts were reported to inhibit cell proliferation and induce apoptosis in various cells.\(^6\) Several ursodeoxycholic acid (UDCA) and chenodeoxycholic acid (CDCA) derivatives developed by Im et al.\(^8\) and it have been reported that they had apoptosis-inducing effect in various cancer cells\(^9,10\).

Cisplatin is one of the most potent anticancer agents showing significant clinical activity against a variety of solid tumors belonging to a class of platinum containing anti-cancer compounds.\(^11\) It is a representative of anticancer drug used to treat certain types of head and neck cancer, cervical carcinoma, lung cancer, neurologic cancers, and a wide variety of other cancers. However, its resistance remains a significant barrier to the survival of cancer patient.

Recent studies have demonstrated that the co-treatment of an antitumor agent having the synergistic anti-cancer effect can be one of the potential therapeutic strategies reducing the extent and severity of cancer treatment-related toxicity. To date, combination treatment of HS-1200 and cisplatin has systemic account of synergistic apoptotic effects on human squamous carcinoma cell line is still scarce. Here we have investigated the synergistic apoptotic effect of co-treatment with a chenodeoxycholic acid (CDCA) derivative, HS-1200 and an anticancer drug, cisplatin on human oral squamous carcinoma (SCC25) cell line.

II. MATERIALS AND METHODS

1. Reagents

The synthetic bile acid derivative, HS-1200 was kindly provided by Professor Young-Hyun Yoo (Department of Anatomy, College of Medicine, Dong-A University, Busan, Korea). The following reagents were obtained commercially: Mouse monoclonal anti-human caspase-3, caspase-7, caspase-9, poly(ADP-ribose) polymerase (PARP), cytochrome c, apoptosis-inducing factor (AIF) antibodies, and rabbit polyclonal anti-human DFF40 (CAD), DFF45 (ICAD), β-actin antibodies, and FITC-conjugated goat anti-mouse and anti-rabbit IgGs were from Santa Cruz Biotechnology (Santa Cruz, CA, USA); HRP-conjugated sheep anti-mouse and anti-rabbit IgGs were from Amersham GE Healthcare (Little Chalfont, UK). 5,5',6,6'-tetrachloro-1,1',3,3'-tetraethylbenzimidazol carbocyanine iodide (JC-1) was from Molecular Probes (Eugene, OR, USA). ApoScreenTM Annexin V-FITC Apoptosis Kit was from Beckman Coulter (Fullerton, CA, USA). Suc-LLVY-AMC was from Calbiochem (Darmstadt, Germany). Dulbecco’s modified Eagle’s medium (DMEM) and FBS were
from Gibco (Gaithersburg, MD, USA). Dimethyl sulfoxide (DMSO), eugenol, cisplatin, Hoechst 33342, RNase A, proteinase K, aprotinin, leupeptin, PMSF, thiazolyl blue tetrazolium bromide and propidium iodide (PI) were from Sigma (St. Louis, MO, USA); SuperSignal West Pico enhanced chemiluminescence Western blotting detection reagent was from Pierce (Rockford, IL, USA).

2. Cell culture

The SCC25 human oral squamous carcinoma cell line was purchased from ATCC (Rockville, USA). Cells were maintained at 37°C with 5% CO₂ in air atmosphere in DMEM/F12 with 4 mM L-glutamine, 1.5 g/l sodium bicarbonate, 45 g/l glucose and 1.0 mM sodium pyruvate supplemented with 10% FBS.

3. Treatment of HS-1200 and cisplatin for co-treatment

Stock solutions of HS-1200 (100 mM) and cisplatin (25 mg/ml) made by dissolving them in DMSO were kept frozen at -20°C until use. Twenty four hours after SCC25 cells were subcultured, the original medium was removed. The cells were washed with PBS and then incubated in the same fresh medium. Since single treatment of 25 μM HS-1200 and 4 μg/ml cisplatin for 24 h showed slight induction of cell death in MTT assay, this single concentration was utilized for further assessment of apoptosis for co-treatment. The concentration of DMSO used in this study had no effect on SCC25 cell proliferation in my preliminary studies.

4. MTT assay

SCC25 Cells were placed in a 96-well plate and were incubated for 24 h. Then they were treated with 25 μM HS-1200 and 4 μg/ml cisplatin for 24 h each and co-treatment with HS-1200/cisplatin were performed in the same condition for 24 h. After cells were treated with 500 μg/ml of thiazolyl blue tetrazolium bromide (MTT solution), they were incubated at 37°C with 5% CO₂ for 4 h. And then the medium was aspirated and formed formazan crystals were dissolved in DMSO. Cell viability was measured by an ELISA reader (Tecan, Männedorf, Switzerland) at 570 nm excitatory emission wavelength.

5. Hoechst staining

Cells were harvested and cell suspension was centrifuged onto a clean, fat-free glass slide with a cytocentrifuge. The samples were stained in 4 μg/ml Hoechst 33342 for 30 min at 37°C and fixed for 10 min with 4% paraformaldehyde.

6. DNA electrophoresis

2 x 10⁶ cells were resuspended in 1.5 ml of lysis buffer [10 mM Tris (pH 7.5), 10 mM EDTA (pH 8.0), 10 mM NaCl and 0.5% SDS] into which proteinase K (200 μg/ml) was added. After samples were incubated overnight at 4°C, 200 μl of ice cold 5 M NaCl was added and the supernatant containing fragmented DNA was collected after centrifugation. The DNA was then precipitated overnight at -20°C in 50% isopropanol and RNase A–treated for 1 h at 37°C. The DNA from 10⁶ cells (15 μl) was equally loaded on each lane of 2% agarose gels in Tris-acetic acid/EDTA buffer containing 0.5 μg/ml ethidium bromide at 50 mA for 1.5 h.

7. Proteasome activity

Cells were collected and lysed in proteasome buffer [10 mM Tris-HCl, pH 7.5, 1 mM EDTA, 2 mM ATP, 20% glycerol, and 4 mM dithiothreitol (DTT)], sonicated, and then centrifuged at 13,000 g at 4°C for 10 min. The supernatant (20 μg of protein) was incubated with proteasome activity buffer [0.05 M Tris-HCl, pH 8.0, 0.5 mM EDTA, 50 μM Suc–LLVY–AMC] for 1 h at 37°C. The fluorescence intensity of each solution was measured by a modular fluorimetric system (Spex Edison, NJ,
USA) at 380 nm excitatory and 460 nm emission wavelengths. All readings were standardized based on the fluorescence intensity of an equal volume of free AMC solution (50 μM).

8. Western blot analysis

Cells (2 x 10⁶) were washed twice in ice-cold PBS, resuspended in 200 μl ice-cold solubilizing buffer (300 mM NaCl, 50 mM Tris-Cl (pH 7.6), 0.5% Triton X-100, 2 mM PMSF, 2 μg/ml aprotinin and 2 μg/ml leupeptin) and incubated at 4°C for 30 min. The lysates were centrifuged at 14,000 revolutions per min for 15 min at 4°C. Protein concentrations of cell lysates were determined with Bradford protein assay (Bio-Rad, Richmond, CA, USA) and 50 μg of proteins were loaded onto 7.5-15% SDS/PAGE. The gels were transferred to Nitrocellulose membrane (Amersham GE Healthcare, Little Chalfont, UK) and reacted with each antibody. Immunostaining with antibodies was performed using SuperSignal West Pico enhanced chemiluminescence substrate and detected with Alpha Imager HP (Alpha Innotech, Santa Clara, USA).

9. Immunofluorescent staining

Cells were cytocentrifuged and fixed for 10 min in 4% paraformaldehyde, incubated with each primary antibody for 1 h, washed 3 times for 5 min, and then incubated with FITC-conjugated secondary antibody for 1 h at room temperature. Cells were mounted with PBS. Fluorescent images were observed and analyzed under Zeiss LSM 700 laser-scanning confocal microscope (Göttingen, Germany).

10. Assay of mitochondrial membrane potential (MMP)

JC-1 was added directly to the cell culture medium (1 μM final concentration) and incubated for 15 min. The medium was then replaced with PBS. Flow cytometry to measure MMP was performed on a CYTOMICS FC500 flow cytometry (Beckman Coulter, Brea, CA, USA). Data were acquired and analyzed using CXP software version 2.2.

11. Quantification of DNA hypoploidy by flow cytometry

After treatment for 24 h, cells were harvested by trypsinization and ice cold 95% ethanol with 0.5% Tween 20 was added to the cell suspensions to a final concentration of 70% ethanol. Fixed cells were pelleted, and washed in 1% BSA-PBS solution. Cells were resuspended in 1 ml PBS containing 20 μg/ml RNase A, incubated at 4°C for 30 min, washed once with BSA-PBS, and resuspended in PI solution (10 μg/ml). After cells were incubated at 4°C for 5 min in the dark, DNA content were measured on a CYTOMICS FC500 flow cytometry system (Beckman Coulter, FL, CA, USA) and data was analyzed using the Multicycle software which allowed a simultaneous estimation of cell-cycle parameters and apoptosis.

III. RESULTS


To investigate whether the co-treatment with HS-1200 and cisplatin reduced the viability of SCC25 cells, MTT assay was conducted. Single treatment of HS-1200 at 25 μM or cisplatin at 4 μg/ml for 24 h reduced the viability of SCC25 cells slightly (HS-1200, 94.6%; cisplatin, 93.4%). Co-treatment with HS-1200 and cisplatin significantly reduced the to 45.2% cell viability vs single treatment (Fig. 1).

2. Co-treatment with HS-1200 and cisplatin augmented the nuclear condensation in SCC25 cells.

To study a nuclear condensation induced by the
co-treatment, Hoechst staining by which a hallmark of apoptosis was conducted. Hoechst staining showed a nuclear condensation by the single treatment of HS-1200 and cisplatin. Co-treatment of HS-1200 and cisplatin remarkably reduced the proteasome activity, compared to the example of the single treatment (Fig. 4).

5. Augmentation of apoptosis by the co-treatment with HS-1200 and cisplatin was demonstrated by reduction of mitochondrial membrane potential (MMP) in SCC25 cells.

Single treatment with either HS-1200 or cisplatin reduced slightly the MMP compared to control group. The co-treatment with HS-1200 and cisplatin result in the remarkable loss of MMP (Fig. 5).

6. Augmentation of apoptosis by the co-treatment with HS-1200 and cisplatin was demonstrated by the decrease in a DNA content of SCC25 cells.

The flow cytometry result showed that the co-treatment with HS-1200 and cisplatin significantly increased apoptotic cells with DNA hypoploidy compared to the example of single treatment (Fig. 6).
Fig. 3. Co-treatment with HS-1200 and cisplatin efficiently showed DNA fragmentation in SCC25 cells. DNA fragmentation analysis was determined by the agarose gel electrophoresis. Whereas single treated cells showed no DNA fragmentation, co-treated cells showed DNA degradation characteristic of apoptosis with a ladder pattern of DNA fragments. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)

Fig. 4. Co-treatment with HS-1200 and cisplatin significantly showed the reduction of proteasome activity in SCC25 cells compared to the single treatment. Three independent assays were performed. Values are means ± SD of triplicates of each experiment. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)

Fig. 5. Co-treatment with HS-1200 and cisplatin showed remarkably the loss of MMP (△ψm) compared to the single treatment. MMP was measured by JC-1 with flow cytometry. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)

7. Co-treatment with HS-1200 and cisplatin caused the translocation of AIF from mitochondria onto the nucleus.

The confocal microscopy showed that AIF was located at mitochondria in the single treatment of HS-1200 or cisplatin whereas AIF was evidently translocated onto nuclei in the co-treatment (Fig. 7).

8. Co-treatment with HS-1200 and cisplatin caused the release of cytochrome c from mitochondria into the cytosol.

The confocal microscopy showed that cytochrome c was located at mitochondria in the single treatment with either HS-1200 or cisplatin whereas cytochrome c was evidently released into the cytosol by the co-treatment (Fig. 8).
Apoptotic Effect of co-treatment with HS-1200 and Cisplatin on SCC25 Human Tongue Squamous Cell Carcinoma Cell Line

Fig. 6. The kinetic analysis of the effect of co-treatment on the SCC25 cell cycle progression and the induction of apoptosis. Co-treatment showed remarkable increase of apoptotic cells with DNA hypoploidy compared to the single treatment. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)

Fig. 7. The confocal microscopy showed that AIF was evidently translocated onto nuclei in SCC25 cells when co-treated with HS-1200 and cisplatin. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)

9. Augmentation of apoptosis by co-treatment with HS-1200 and cisplatin was demonstrated by Western blot assay.

The expression level of anti-apoptotic protein, Bcl-2 decreased and the expression level of apoptotic protein, Bax increased by the co-treatment (Fig. 9). The co-treatment with HS-1200 and cisplatin induced the degradation of caspase-9, caspase-3, caspase-7 and PARP, and produced the processed DFF45 (ICAD) 11 kDa, PARP 85 kDa and caspase-3 17 kDa cleaved products (Fig. 10A and Fig. 11).

Fig. 8. The confocal microscopy showed that cytochrome c was evidently released to the cytosol in SCC25 cells that were co-treated with HS-1200 and cisplatin. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)
Fig. 9. Western blot analysis showing that co-treatment with HS-1200 and cisplatin induced up-regulation of the Bax and down-regulation of the Bcl-2. The levels of β-actin were used as an internal standard for quantifying Bcl-2 and Bax expression. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)

Co-treatment with HS-1200 and cisplatin caused the translocation of DFF40 (CAD) from cytosol onto the nuclei.

The confocal microscopy showed that DFF40 (CAD) was located at cytosol by the single treatment with either HS-1200 or cisplatin whereas DFF40 (CAD) was evidently translocated into the nuclei by the co-treatment (Fig. 10B).

IV. DISCUSSION

Bile acids are steroid acids that is produced in mammalians liver and stored in the gall bladder. The function of bile acid is to facilitate the formation of micelles, which promotes processing of dietary fat. Several reports demonstrated that natural bile salts, ursodeoxycholic acid (UDCA) and chenodeoxycholic acid (CDCA) were inhibit cell proliferation and induce apoptosis in various cells. Also, it has been reported that the synthetic CDCA derivatives
Apoptotic Effect of co-treatment with HS-1200 and cisplatin on SCC25 Human Tongue Squamous Cell Carcinoma Cell Line

![Western blot analysis showing the co-treatment with HS-1200 and cisplatin in SCC25 cells remarkably induced Caspase-9, Caspase-3, Caspase-7 and PARP degradations and produced the processed Caspase-3 17 kDa, and PARP 85 kDa cleaved products. β-actin, a loading control. (HS, cells treated with 25 μM HS-1200 for 24 h; cis, cells treated with 4 μg/ml cisplatin for 24 h; HS+cis, cells treated with 25 μM HS-1200 plus 4 μg/ml cisplatin for 24 h)](image)

induce the antiproliferative efficacy in various cancer cells by inducing apoptosis. Those studies demonstrated the decrease of proteasome activity, mitochondrial events, and nuclear condensation in synthetic CDCA derivatives-induced apoptosis. In addition, it has been demonstrated that a synthetic CDCA derivative, HS-1200 shows the strongest apoptosis-inducing effect among the synthetic CDCA derivatives.

Since from the 1970s, the neutral, square planar, coordination complex cis-diaminedichloroplatinum (II) (cisplatin) has become more and more popular in the treatment of cancer. Cisplatin, which is a well-known DNA-damaging agent and one of the most active cancer treatment agents available, is widely used for the treatment of many malignancies, including testicular, ovarian, bladder, cervical, head and neck, and small-cell and non-small-cell lung cancers. Unfortunately, however, cisplatin causes many untoward side effects. It has been reported that cisplatin-induced cell death is involved in apoptotic- and cell cycle-regulating pathways. Moreover, the synergistic antitumor effect of the combination treatment with cisplatin and antitumor agents or natural products have also been demonstrated.

If the co-treatment with a synthetic CDCA derivative, HS-1200 and a representative of anticancer drug, cisplatin which has severe toxicity and resistance in the chemotherapy, shows a synergistic antitumor effect, it could be a more fundamental therapeutic strategy for cancer chemotherapy.

Proteasome is a fundamental non-lysosomal tool that cells use to process or degrade a variety of short-living proteins. It was reported that proteolysis mediated by ubiquitin-proteasome system has been reported to be implicated in the regulation of apoptosis. The proteasome pathway is mostly known to work at the upstream of the mitochondrial alterations and caspase activation. In this study, the co-treatment with HS-1200 and cisplatin on SCC25 cells causes the significant reduction of proteasome activity compared to the single treatment.

Mitochondria plays an important role in the induction of the mitochondrial permeability transition and also plays a key part in the regulation of apoptosis. Outer mitochondrial membrane becomes permeable to intermembrane space proteins such as cytochrome c and AIF (apoptosis inducing factor) during apoptosis. Release of cytochrome c and disruption of mitochondrial membrane potential (MMP) are known as features in apoptosis triggered by proteasome inhibition. On induction of apoptosis, AIF translocates to the nucleus, resulting in chromatin condensation and large-scale DNA fragmentation. This study evidently showed that...
co-treatment with HS-1200 and cisplatin on SCC25 cells results in remarkable decrease of MMP, increase and decrease of Bax and Bcl-2, the release of cytochrome c into cytosol and translocation of AIF onto nuclei whereas the single treatment does not show these patterns.

A common final event of apoptosis is the nuclear condensation, which is controlled by caspases, DFF, and PARP. Caspases play an essential role during apoptotic death. Once activated, the effector caspases (caspase-3, caspase-6 or caspase-7) are responsible for the proteolytic cleavage of a broad spectrum of cellular targets, ultimately leading to cell death. The known cellular substrates include structural components (such as actin and nuclear lamin), inhibitors of deoxyribonuclease (such as DFF-45 or ICAD) and DNA repair proteins. In apoptotic cells, activation of DFF40 (CAD), also a substrate of caspase-3, occurs with the cleavage of DFF45 (ICAD). Once DFF40 is activated and released from the complex of DFF45 and DFF40, it can translocate to the nucleus and then degrade chromosomal DNA and produce DNA fragmentation. This study demonstrated that co-treatment with HS-1200 and cisplatin on SCC25 cells results in the degradation and the cleavage of caspase-9, caspase-3, caspase-7, PARP and DFF45 (ICAD), and also the translocation of DFF40 (CAD) onto nuclei whereas the single treatment does not.

Taken collectively, combination therapy with HS-1200 and cisplatin could be considered, in the future, as an alternative therapeutic strategy for human oral squamous carcinoma. Its clinical application awaits further extensive studies.

REFERENCES

Apoptotic Effect of co-treatment with HS-1200 and Cisplatin on SCC25 Human Tongue Squamous Cell Carcinoma Cell Line


국문초록

**HS-1200과 cisplatin의 병용처리가 사람구강암세포에 미치는 세포자멸사 효과에 대한 연구**

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담즙산은 지방의 흡수와 콜레스테롤의 항성성 조절하는 유전자 전사에 관여하는 필수 콜레스테롤의 생성물이다. 담즙산 함성유도체인 HS-1200이 여러 가지 암세포에서 세포자멸시(apoptosis)를 유도한다는 것이 알려져 있다. 본 연구는 사람구강편평세포암종세포(SCC25 cells)에서 담즙산 함성유도체인 HS-1200과 대표적인 항암제인 cisplatin의 병용처리 후 세포자멸시 증가효과가 있는지 알아보기 위해 수행하였다.

HS-1200과 cisplatin의 병용처리가 단독처리에 비해서 효과적인 세포생존율 감소가 있는지 확인하기 위해서 MTT법을 시행하였고, 세포자멸시의 유도와 증가를 알기 위해서는 DNA 전기영동법, Hoechst 염색법, DNA hypoploidy법을 사용하였다. 그리고 세포자멸시에 관계하는 단백질의 발현 변화와 세포내에서의 이동을 밝혀내기 위해서 Western blot 분석과 면역형광 염색법을 수행하였다. 더 나아가서 proteasome 활성도와 세립체막 전위 변화를 측정하였다.

본 연구에서는 HS-1200과 cisplatin을 병용처리한 SCC25 세포에서 핵의 농축, DNA분절, MMP와 proteasome 활성도의 감소, Bax의 증가와 Bcl-2의 감소, DNA방의 감소, cytochrome c의 세포질로의 유리, AIF와 DFF40(CAD)의 핵으로의 이동, caspase-9, caspase-7, caspase-3, PARP 그리고 DFF45(ICAD)의 활성화와 같은 다양한 세포자멸시 증거를 보였다. 반면에 상기 물질들에 단독처리 된 SCC25 세포에서는 세포자멸시 현상이 거의 없었다.

24시간 동안 25 μM의 HS-1200, 4 μg/ml의 cisplatin 을 각기 단독처리 한 결과에서는 세포자멸시를 거의 유도하지 못했으나, 병용처리한 결과에는 아주 탁월하고 명확한 세포자멸식의 유도를 보였다. 그러므로 본 실험결과는 HS-1200과 cisplatin의 병용요법이 사람구강편평세포암종 환자에게 새로운 치료전략으로서의 가능성을 보여준다고 생각한다.

**Key words**: 세포자멸사, 담즙산, HS-1200, cisplatin, 사람구강편평세포암종