Synergy of novel coumarin derivatives and tamoxifen in blocking growth and inducing apoptosis of breast cancer cells

Lulzime Ballazhi1,2, Faik Imeri3, Aleksandar Dimovski1, Ahmed Jashari4, Emil Popovski5, Pranvera Breznica-Selmani1,6, Bozana Mikhova7, Gerald Dräger8, Edita Alili-Idrizi2, Kristina Mladenovska1*

1Faculty of Pharmacy, University “Ss Cyril and Methodius”, Mother Theresa 47, 1000 Skopje, Macedonia
2Faculty of Medical Science, State University of Tetovo, Pashe Deralla bb, 1200 Tetovo, Macedonia
3Institute of Pharmacology, University of Bern, Friedbühlstrasse 49, CH-3010 Bern, Switzerland
4Faculty of Natural Sciences & Mathematics, State University of Tetovo, Pashe Deralla bb, 1200 Tetovo, Macedonia
5Institute of Chemistry, Faculty of Natural Sciences & Mathematics, University “Ss. Cyril and Methodius”, PO Box 162, 1000 Skopje, Macedonia
6Faculty of Medicine, University “Hasan Pristina”, Mother Theresa 10 000, Pristina, Kosovo
7Institute of Organic Chemistry with Centre of Phytochemistry, Bulgarian Academy of Sciences, Acad. G. Bonchev 9, 1113 Sofia, Bulgaria
8Institute of Organic Chemistry, Leibniz Universität Hannover, D-30167, Hannover, Germany

Abstract

Possible synergistic effect of tamoxifen (2 μM) and hydrazinyldiene-chroman-2,4-diones (10-100 μM) was examined with an aim to create more effective treatment for ER+ breast cancer. Anti-breast cancer effect has been evaluated on the proliferation of MCF-7 breast adenocarcinoma cells using MTT and alamarBlue assays. Cell viability was evaluated after 48h-treatment and the IC50 of the coumarin derivatives were determined. The apoptotic effect was evaluated by detection of PARP cleavage and reduced activity of the survival kinase Akt. The results demonstrated dose-dependent activity, with a percent of growth inhibition after combination treatment being significantly higher (53% to 79%, 10 μM and 100 μM, respectively) than the one in the cell lines treated with tamoxifen (29% to 37%) and the synthesized coumarin derivatives alone (11% to 68%, 10 μM and 100 μM, respectively). The IC50 of the synthesized compounds significantly decreased in synergy with tamoxifen (33% to 51%). Coumarin derivative having thiazole moiety with additional methyl groups attached to the carbons at positions 5 and 4 in the thiazole ring showed to be the most potent, with IC50 20 μM when administered alone and 10 μM in synergy with tamoxifen. The levels of phospho-Thr308 Akt were down-regulated by the combination treatment, pointing to tyrosine kinase phosphorylation inhibition. In conclusion, the novel coumarin derivatives enhance the activity of tamoxifen and this combination may be suitable for prevention of ER+ breast cancer or development of related compounds. Further studies are needed to elucidate precisely the type of receptor involved in the activity and the mechanism of action.

Keywords: hydrazinyldiene-chroman-2,4-diones, tamoxifen, breast cancer, MCF-7 cells, antiproliferative effect

* tel: 00389 2 3126 032; fax: 00389 2 3123 054
  e-mail: krml@ff.ukim.edu.mk
Introduction

The balance between proliferation and cell death, mainly apoptosis, is crucial in determining the overall growth and regression of tumor. For this reason, traditional cytotoxic agents, such as DNA intercalating agents, DNA cross-linking agents, topoisomerase inhibitors, cytoskeleton disrupting agents and antimetabolites have usually been targeted to damage the aberrantly dividing cancer cells by interrupting the cell division process (Hengartner, 2000; Johnstone et al., 2002; Mehlen and Puisieux, 2006). However, rapidly growing nonmalignant cells are also significantly affected, which in turn, limits efficacy and probably increases the risk of drug resistance, particularly when toxicities lead to less than optimal drug dosing. Drug resistance, the signal transductions within the process of oncogenesis, cancer cell survival and metastasis are complex, and for survival, cancer cells often rely on redundant signaling pathways when the original pathway is blocked. Therefore, to achieve the best outcome for the cancer patients, often several anti-cancer agents that inhibit multiple targets or redundant pathways are combined in the treatment regimen (Li et al., 2014; LoRusso et al., 2012). Treatment combinations, including combination of nonspecific small molecule chemotherapeutic agents have been successfully applied to various cancer types, including (metastatic) breast cancer (Lee and Nan, 2012).

Breast cancer is still the most frequent malignancy cancer in women and principal cause of cancer death for females worldwide (Jemal et al., 2009; Ferlay et al., 2010). It is estimated that app. 80% of the breast tumors are estrogen receptor positive (ER+) and their growth is stimulated by estrogens (Hertz et al., 2009). Despite tremendous progress in their treatment during past decades, severe toxic side-effects and de novo and acquired resistance to the existing therapies present a major clinical problem urging for design and development of novel drugs and treatment combinations.

The coumarin (benzopyran-2-one) ring system, present in natural products, has intrigued medicinal chemists for decades to explore the natural coumarins or their synthetic analogs for their applicability as anticancer drugs due to the antiproliferative and cytotoxic effect on malignant cells and usefulness in prevention of recurrence and metastases of malignant cells (Sadiu et al., 2012; Goel et al., 2009; Jacquot et al., 2001). Of particular interest in breast cancer chemotherapy, some coumarins and their active metabolite 7-hydroxycoumarin analogs have shown sulfatase and aromatase inhibitory activities. Coumarin based selective estrogen receptor modulators and coumarin-estrogen conjugates have also been described as potential anti-breast cancer agents (Musa et al., 2008).

In search for novel coumarin based antibreast cancer drugs, we have synthesized compounds that combine the coumarin core and five membered heterocycles (isoxazoles and thiazoles) in hydrazinylidiene-chroman-2,4-di-ones (Jashari et al., 2013; 2014; Ballazhi et al., accepted for publication). Isoxazole and thiazole substituents were chosen as important pharmacophores in agents with antiproliferative and tumor vascular-disrupting activity (Gulsoy and Guzeldemirci, 2007; Poma et al., 2007). Eight compounds were synthesized, showing to have antiproliferative effects and induce apoptosis on breast cancer cells MCF7 and MDA-MB-231 (Jashari et al., 2014) and bone and lung metastatic cell lines from breast cancer, SCP1833 and SCP4175, respectively (Ballazhi et al., accepted for publication). Three of the eight compounds tested, having thiazole moiety, without or with additional methyl group(s) attached to the carbon(s) at the position(s) 5 and/or 4 in the thiazole ring showed to be the most potent, requiring further evaluation.

Based on these findings, the aim of this study was to test their antiproliferative effects on MCF-7 cell lines in synergy with tamoxifen (Tam). Tam induces growth arrest and apoptosis in breast cancer cells through the inhibition of estrogen binding to the ERs, improving the clinical outcome of patients with both early and advanced breast cancer. Also, at high concentrations (≥10 μM), it has been shown to mediate apoptosis in ER-negative cancer cells (Higgins et al., 2011; Fisher et al., 2005; Gelmann, 1996; Sutherland et al., 1982). However, about half of patients with advanced ER+ disease immediately fail to respond to Tam, and the disease progresses to a resistant phenotype. Mechanisms by which breast cancer cells become resistant may include changes in the expression of ERα or ERβ, alterations in co-regulatory proteins, and the influences of cellular kinase signal transduction pathways (Kruh, 2003; Dorsses et al., 2001). Significantly higher concentration-dependent reduction in cell viability and increase in apoptosis, confirmed by PARP cleavage and reduced activity of the survival kinase Akt, was observed with combination of Tam and coumarin derivatives in comparison with the compounds alone.

Materials and Methods

Cell culturing

The human breast cancer cell line MCF-7 (MCF-7 from American Type Culture Collection) was maintained in Dulbecco’s modified eagle’s medium (DMEM, Invitrogen, Switzerland) containing 10% fetal bovine serum (FBS, Invitrogen, Switzerland) supplemented with 300 μg mL⁻¹ glutamax (Invitrogen, Switzerland), 100 μg mL⁻¹ benzylpenicillin (Invitrogen, Switzerland) and 100 units mL⁻¹ streptomycin (Invitrogen, Switzerland). MCF-7 cells were incubated at 37 °C in an atmosphere containing 5% CO₂. For experiments, cells from exponentially growing culture were used.

MTT assay

Cytotoxic effects of Tam (T5648 Sigma, USA), 4-hydroxycoumarin (4-HC, as a reference, Merck KGaA, Ger-
Synergy of novel coumarin derivatives and tamoxifen in blocking growth and inducing apoptosis of breast cancer cells

many) and the three coumarin derivatives synthesized in our laboratory (1c, 1d, 1e; Fig. 1) on MCF-7 cells were determined by 3[4, 5-dimethylthiazol-2-yl]-2, 5-diphenyltetrazolium bromide (MTT, Sigma-Aldrich, USA) assay.

In short, 100 µL of the growth medium DMEM was poured in each well of a 96-well plate and seeded with 5000 MCF-7 cells per well. Cells were allowed to attach overnight and then 4-HC, 1c, 1d and 1e in increasing concentrations (10, 20, 50, 100 µM) and Tam (2µM) were added to respective wells. After 48 hours of incubation at 37 °C, 5% CO₂ and relative humidity 95%, 20 µL of MTT reagent (5 mg mL⁻¹) was added to each well. After additional incubation within 4 hours, 100 µL of MTT-solvent (4 mM HCl, 0.1% Nonidet P-40 (in isopropanol), AppliChem, Germany) was added to each well to solubilize the MTT crystals. The plates were covered with tinfoil and the cells were agitated on orbital shaker for 15 min. Then, the absorbance was read at 570 nm in a microplate reader (SpectraMax M2 Fluorimeter, BucherBiotec Inc, USA). All experiments were performed at least 3 times, with 4 wells for each concentration of the tested agents. The control cells were grown under the same conditions without adding the test compounds. Cell survival was calculated using the following formula:

$$\% \text{ of cell viability} = \frac{\text{mean experimental absorbance}}{\text{mean control absorbance}} \times 100\%$$

**AlamarBlue assay**

AlamarBlue assay was performed in 96-well microtiter plates. In each well, 100 µL of the culture medium with 4000 cells was added. After 24 hours, the medium was removed from all wells and replaced by fresh medium (150 µL) containing the compounds. After 48 hours of incubation, the plates were inspected under an inverted microscope to assure growth of the controls and sterile conditions. Afterwards, alamarBlue reagent® (10 µL) (Invitrogen, Basel, Switzerland) was added to each well and the plates were incubated for another 2 to 3 hours. Then, the plates were read with a SpectraMax M2 microplate reader (SpectraMax M2 Fluorimeter, BucherBiotec Inc, USA) at excitation wavelength of 540 nm and emission wavelength of 570 nm. The percent of viability was expressed as fluorescence counts in the presence of test compound as a percentage of that in the vehicle control.

**Statistical analysis**

The 50% inhibitory concentration (IC₅₀) was determined as the anticancer drug concentration causing 50% reduction in cell viability and calculated from the viability curves by linear interpolation between the values immediately above and below the 50% inhibition using the Bliss’s software (Bliss Co, Castro Valley, CA 94552, USA).

The results were presented as mean ± SD and the statistical analysis was performed using one-way analysis of variance (ANOVA) followed by a Bonferroni’s post hoc test for multiple comparisons (GraphPad InStat version 3.00 for Windows NT, GraphPad Software, San Diego, CA, USA).

**Results**

**Cytotoxic and anti-proliferative effects of Tam, 4-HC and coumarin derivatives on breast cancer cells**

The effects of Tam, 4-HC and coumarin derivatives (1c, 1d and 1e) on cell viability and proliferation, alone or in combination, evaluated by MTT and alamarBlue assays, are presented in Fig. 2A and 2B, respectively. Differences in cell viability and proliferation were observed when data obtained from the two assays were compared, with the per-
The results showed that only synthesized coumarin derivatives, but not 4-HC induced anti-proliferative effect in MCF-7 cell lines. Even when 4-HC was administered with Tam, the number of viable cells remained high, from app. 67% (100 µM, MTT assay) to 100% (10 µM, alamarBlue assay). 4-HC alone induced no apoptosis when concentrations equal to or lower than 20 µM were applied.

The percentage of growth inhibition after treatment with the corresponding hydrazinylidiene-chroman-2,4-diones was dramatically higher than the one in the untreated control cells, especially when concentrations higher than 10 µM were applied in synergy with Tam. Namely,
as MTT and alamarBlue assays showed, when the synthesized agents were administered in combination with Tam, the percent of growth inhibition ranged from app. 51% to app. 79% (MTT assay) and from app. 12% to 65% (alamarBlue assay), with increase in the concentration of the agents from 10 to 100 µM. Administration of Tam alone decreased the number of viable cells to lower extent, from 37% (MTT assay) to 29% (alamarBlue assay), expressed as a percent from control (100%). Similarly, when synthesized compounds were administered alone, with the lowest concentration, the percent of viable cells was higher in comparison with Tam alone, from 89% to 83% (MTT and alamarBlue assays) vs. app. 63% (MTT assay) and 71% (alamarBlue assay) for Tam, respectively. With increase in the concentration of the synthesized coumarin derivatives, from 50 to 100 µM, when they solely administered, the percent of viable cells was lower than the one when only Tam was administered, app. 32% and 45% (1d, 100 µM, MTT and alamarBlue assay, respectively). Synergy of the synthesized agents with Tam resulted in average 30.3±7.6% (MTT assay) to 21.0±3.5% (alamarBlue assay) higher decrease in viable cells in comparison with the coumarin derivatives alone, when administered in concentration of 100 µM. When concentration of 10 µM was applied, decrease in non-apoptotic cells was 43.3±2.5% (MTT assay) to 11.1±4.7% higher in comparison with the coumarin derivatives alone.

As Table 1 shows, the IC\textsubscript{50} of the synthesized compounds significantly decreased when they administered in synergy with Tam, from 33% (1c, alamarBlue assay) to 51% (1c, MTT assay). The coumarin derivative having thiazole moiety, with additional methyl groups attached to the carbons at the positions 5 and 4 in the thiazole ring (1e) showed to be the most potent, with IC\textsubscript{50} around 20 µM and 10 µM when administered alone and in synergy with Tam, respectively (Table 1).

Table 1. Cytotoxicity of 4-hydroxycoumarin and its synthesized derivatives against MCF-7 breast cancer cell lines determined by MTT and alamarBlue assays\textsuperscript{a}

<table>
<thead>
<tr>
<th>Coumarin derivatives</th>
<th>IC\textsubscript{50} (µM)</th>
<th>Compound+Tam (2 µM)</th>
<th>IC\textsubscript{50} (µM)</th>
<th>Compound+Tam (2 µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MTT assay</td>
<td></td>
<td>AlamarBlue assay</td>
<td></td>
</tr>
<tr>
<td>4-HC</td>
<td>&gt; 100</td>
<td>&gt; 100</td>
<td>&gt; 100</td>
<td>&gt; 100</td>
</tr>
<tr>
<td>1c</td>
<td>19.76±3.95</td>
<td>9.72±3.91</td>
<td>29.60±2.19</td>
<td>19.76±5.52</td>
</tr>
<tr>
<td>1d</td>
<td>19.64±5.40</td>
<td>10.05±4.58</td>
<td>29.24±1.60</td>
<td>16.60±7.87</td>
</tr>
<tr>
<td>1e</td>
<td>19.94±1.90</td>
<td>9.80±3.45</td>
<td>19.91±5.98</td>
<td>10.33±2.00</td>
</tr>
</tbody>
</table>

\textsuperscript{a}Values are means ± SD from at least three independent experiments.

![Western blot analysis of cleaved PARP and phospho-Akt](image)

**Fig. 3.** Western blot analysis of cleaved PARP and phospho-Akt after incubation of MCF-7 cells with the vehicle (Co), Tam and Tam+novel coumarin derivatives for 48 h. Western blot analysis was performed by using antibodies against PARP (upper panels), phospho-Thr308-Akt (middle panels) and the house-keeping protein GAPDH (lower panels). The data represent three identical experiments.
Apoptotic effects of Tam and coumarin derivatives on breast cancer cells

We investigated whether the combination of Tam and 4-HC or coumarin derivatives synergistically affected apoptosis of MCF-7 cells. PARP (poly ADP ribose polymerase) cleavage which is a marker of apoptosis was determined by Western blot analysis. Fig. 3 shows that in MCF-7 cells, all concentrations of 1c, 1d and 1e in combination with Tam led to a reduced protein expression of 116 kDa full length PARP with a concentration-dependent appearance of a cleavage product at 86 kDa. In parallel, the levels of phospho-Thr508 Akt were downregulated by combination of Tam and 1c, 1d and 1e when compared to control and to combination of Tam and 4-HC treated cells. Akt is a well known protein kinase involved in cell growth and survival and reduced activity often correlates with reduced proliferation and increased apoptosis. The level of phospho-Thr508 Akt is well accepted to reflect Akt activity.

Discussion

Tam, a selective ER modulator is a partial agonist of ER, blocking the proliferative effects of estrogen via this receptor. Since its introduction in cancer therapy, it has become the standard treatment option for hormone-responsive breast cancer patients (Bush, 2007; Goldhirsch et al., 2009; Fisher et al., 1998; 2005). It was shown that the Tam effect can be reversed by addition of 17β-estradiol and that Tam has no effect on the cell cycle kinetics of the receptor-negative MDA-MB-231 cells, confirming that the antiestrogen effect is mediated through the ER (Osborne et al., 1982). This was also confirmed in the study of Gelmann (1996) and Mandlekar and Kong (2001), in which more apoptosis was induced in ER+ MCF-7 breast cancer cells compared to the MDA-MB-231 cells. Several hormone receptor-independent effects have been also described for Tam, leading to apoptosis when higher concentrations are applied (Fisher and Redmond, 1992; Mandlekar and Kong, 2001).

Despite the relative safety and significant anti-neoplastic activities of Tam, most initially responsive breast tumors develop resistance to this drug. Even though an improved understanding, resistance to anti-estrogen therapy remains a significant clinical problem. However, combination therapies of Tam with other drugs that aimed at the plastic activities of Tam with other drugs that aimed at the regulation of caspase-9 or cleavage of poly (ADP-ribose) polymerase has a primary role in the acute death response of the MCF-7 cells. Accordingly, an inhibitor of mitochondrial permeability transition but not the caspase inhibitor could be able to protect MCF-7 cells against Tam.

In this study, we have demonstrated that the antiestrogen Tam, used in concentration of 2 μM inhibits proliferation of ER+ MCF-7 cells. These results, are similar to those reported by Lippman et al. (1976) and Sutherland et al. (1982) who found that, at concentrations of Tam observed in women treated for breast cancer (<1.0 μM), the antiestrogen slowed proliferation but was not lethal to MCF-7 cells associated with release of mitochondrial cytochrome c, a decrease of mitochondrial membrane potential and an increase in production of reactive oxygen species (Kallio et al., 2005). This suggests that disruption of mitochondrial function but not immediate activation of caspase-9 or cleavage of poly (ADP-ribose) polymerase has a primary role in the acute death response of the MCF-7 cells. Accordingly, an inhibitor of mitochondrial permeability transition but not the caspase inhibitor could be able to protect MCF-7 cells against Tam.

In our previous studies, we have shown that three of the eight newly synthesized coumarin derivatives, having thiazole moiety without or with additional methyl group(s)
attached to the carbon(s) at the position(s) 5 and/or 4 in the thiazole ring were effective in inhibiting the growth of different cancer cells, including MCF-7 breast cancer cells, in a dose- and time-dependent manner (Jashari et al., 2014). Similar results were obtained when estrogenic activity on MCF-7 breast carcinoma cells of compounds prepared by addition of 2-aminothiophenol to substituted 4-HC derivatives was evaluated. Among the compounds tested, 6,12-dihydro-3-methoxy-1-benzopyran[3,4-b][1,4]benzothiazin-6-one and 6,12-dihydro-3-hydroxy-1-benzopyran[3,4-b][1,4]benzothiazin-6-one exhibited an ER-dependent proliferation and a high binding affinity to ER, but a moderate capacity to activate the transcription of a reporter gene (Jacquot et al., 2001). The pyrone-substituted coumarin, namely warfarin, isoflavone genistein and 6,7-dihydroxycoumarin known as esculetin were also examined, under similar assay procedures, using the MCF-7 cell lines. Proliferation assays yielded that the MCF-7 cells tested were quite sensitive to the effects of all three compounds, with potency of growth inhibition by genistein being greater than the one of esculetin which, in turn was greater than that of warfarin. Genistein has exceptionally interesting, multidirectional therapeutic properties and the biological activity of this substance, as of all isoflavones, is conditioned by the location of the phenyl ring near the third carbon of the benzo-γ-pyrone. It has been shown that inhibition of cancer cell proliferation was attributed to inhibition of several key enzymes, especially tyrosine kinase, which plays a critical role in cell proliferation and transformation, and is also associated with oncogene expression in breast cancer. The investigation of how esculetin, genistein and warfarin affect the signal transduction cascade and cell cycle progression in MCF-7 cells confirmed inhibition of tyrosine phosphorylation, with warfarin being a less potent inhibitor of cell proliferation and metabolic activity than esculetin and genistein, partly due to the fact that it shows no significant tyrosine kinase phosphorylation inhibition in comparison to the other compounds (Lacy and O'Kendy, 2004). In addition, in vitro studies showed that genistein also exhibited a synergistic additive effect when cancer cells were exposed to a combination treatment with Tam, confirming to have both estrogenic and anti-estrogenic effects (Tanos et al., 2002; Luczkiewicz et al., 2003).

All these findings implicated that the effects of Tam may be enhanced in combination with our coumarin derivatives, which was confirmed by a significant additive cytotoxic effect in the MCF-7 cell lines. Non-significant differences in cell viability and proliferation, observed when the data obtained from the two assays were compared, can be explained by the different sensitivity to the effects of the compounds on cellular metabolism of the two assays. The MTT assay is based on the reduction of tetrazolium salt such as MTT into formazan crystals by living cells, which determines mitochondrial activity. It relies on the activity of just one group of mitochondrial enzymes to predict adverse effects on metabolism, and in doing so may underestimate metabolic effects, which was confirmed for interferons in the past (Jabbar et al., 1989). Upon entering the cells, the active ingredient of alamarBlue®, resazurin, is continuously reduced to resofurin which is further reduced to hydroresorufin. It is still unknown how this reduction occurs, intracellularly via enzyme activity or in the medium as a chemical reaction, although the reduced fluorescent form of alamarBlue was found in the cytoplasm of living and cells nucleus of dead cells (O'Brien et al., 2000). Our data are in contradiction with the previous study (Jabbar et al., 1989) and also with the results of Hamid et al. (2004) who reported slightly higher sensitivity for alamarBlue assay than the MTT assay for most of the tested drugs, out of 117 in total. However, it must be emphasized that the results of Hamid et al. (2004) were obtained from tests in human hepatoma cell line HepG2, at a single point screen. When the drugs were re-tested in both assays for reconfirmation of cytotoxicity and determination of the IC₅₀ values, except for daunorubicin, the IC₅₀ values were comparable in both assays, pointing that both assays provide useful information to identify in vitro cytotoxic drugs at early stages of drug candidate selection. There is also possibility different results to be caused by different level of induction and/or inhibition of the metabolic enzymes responsible for transformation of cell toxicity end points, as was demonstrated using dicumarol (Hamid et al., 2004). This could be one of the reasons why combination treatment of Tam and 4-HC results in higher cell viability in comparison with Tam alone (Fig 2A and B). Namely, there are literature data that report on inhibition of microsomal mixed function oxidase by 4-HC (Deckert et al., 1972) and this could decrease Tam activity knowing that its antiestrogenic effect has been attributed to its metabolism to an active 4-hydroxy derivative and the avid binding of the active metabolite to the estrogen receptor. Biotransformation by P450 forms are expressed extrahepatically, in the breast and endometrium, and may be particularly important in determining tissue-specific effects of Tam (Crewe et al., 2002).

In addition, Tam has different partial agonist-antagonist activities in different tissues and the differences may be related in part to the milieu of ER coactivators and corepressors in these tissues. If a cell or tissue requires only activating factor 1 to interact with transcription factors at the promoter, Tam is agonistic. However, if a cell type requires activating factors 1 and 2 of the estrogen receptor to be functioning concurrently, Tam is antagonistic (Shou et al., 2004). There is a probability that the newly synthesized coumarine derivatives and 4-HC at higher concentrations (50-100 μM) prevent activation of ER and co-activators such as AIB1 in MCF-7 cells, reduce the recruitment of co-activator complexes and enhance recruitment of corepressor complexes to Tam-bound Er on gene promoters, thus potentiating Tam’s estrogen antagonist effects on gene expression and tumor growth. However, this should be confirmed in further studies in which MCF-7 breast can-
cer cells, which express high levels of A1B1, will be treated with Tam alone and in combination with these coumarine derivatives. In addition, effects of these newly synthesized coumarine derivatives on mitochondrial permeability transition as well as enzyme induction/inhibition effects should be evaluated.

Our data demonstrated that Tam and 4-HC derivatives inhibited MCF-7 cells proliferation by inducing apoptosis. The enhanced apoptosis may account for the synergistic inhibition of the combination treatment. The levels of phospho-Thr548 Akt were down-regulated by combination of Tam and the coumarin derivatives, pointing to tyrosine kinase phosphorylation inhibition.

**Conclusion**

The results of this study indicate that the combination of Tam and hydrazinylidine-chroman-2,4-diones having thiazole moiety without or with additional methyl group(s) attached to the carbon(s) at the position(s) 5 and/or 4 in the thiazole ring can effectively inhibit cell proliferation and induce apoptotic pathway. They may be considered as new candidates that could contribute to the development of a large chemical library or related compounds by a combinatorial synthesis approach. Further studies are needed to elucidate with precision the type of receptor involved in the activity and their mechanism of action, including binding mode.

**References**


Synergy of novel coumarin derivatives and tamoxifen in blocking growth and inducing apoptosis of breast cancer cells

Резиме

Синергистички ефект на нови кумарински деривати и тамоксифен во блокирање на растот и предизвикување апоптоза на клетки од рак на дојка

Лулзиме Балажи1,2, Фаик Имери3, Александар Димовски1, Ахмед Јашари4, Емил Поповски5, Пранвера Брезнца-Селмани6, Божана Микхова7, Гералд Дрегер8, Едита Алили-Идризи2, Кристина Младеновска1*

1Фармацевтски факултет, Универзитет “Св. Кирил и Методиј”, бул. Мајка Тереза 47, 1000 Скопје, Македонија
2Факултет за медицински науки, Државен универзитет во Тетово, ул. Паше Дерала бб, 1200, Тетово, Македонија
3Институти за хемија, Факултет за природни науки и математика, Универзитет “Св. Кирил и Методиј”, PO 162, 1000 Скопје, Македонија
4Институт за органска хемија со центар за фитохемија, Бугарска академија за науки, Акад. Г. Бончев 9, 1113 Софија, Бугарија

Ключни зборови: хидразинилдиен-хроман-2,4-диони, тамоксифен, рак на дојка, MCF-7 клетки, антипролиферативен ефект.

Во овој труд е испитуван потенцијалниот синергистички ефект на тамоксифен (2 μM) и хидразинилдиен-хроман-2,4-диони (10-100 μM) со цел да се дизјанира поефективен третман за ER+ клетки на рак на дојка. Антиканцер ефектите беа оценувани во однос на пролиферацијата на MCF-7 клетки со користење на MTT и аlamаргесиите. Виталноста на клетките беше следена после третман од 48 часа, после што беа одредувани IC50 на кумаринските деривати. Апоптотичниот ефект беше оценет со детекција на раскинувањето на поли ADP рибоза полимераза и преку намалувањето на активноста на киназа на преживување Akt. Резултатите покажаа дозно-зависна активност, при што процентот на инхибиција на клеточен раст после комбиниранот третман беше значајно повисок (53% до 79%, 10 μM и 100 μM, соодветно) од соодветниот процент во клеточните линии третирани само со тамоксифен (29% до 37%) и синтетизираним кумарински деривати (11% до 68%, 10 μM и 100 μM, соодветно). IC50 на синтетизираним клеточните линии значајно се намали во синергија со тамоксифен (33% до 51%). Кумаринските дериват со тиазолна структура и дополнителни метил групи на јаглеродите во позиција 5 и 4 во тиазолното јадро прикажа најголем потенцијал, со IC50 20 μM, применет сам, и 10 μM, во синергија со тамоксифен. Со комбиниранот третман, нивоата на фосфо-Thr308 Akt беа надолно регулирани, упатувајќи на инхибиција на фосфорилацијата на тирозин киназа. Како заклучок, новите кумарински деривати ја зголемуваат активноста на тамоксифен и овој комбиниран третман може да биде соодветен за превенција на ER+ рак на дојка или за развој на сродни соединија со комбинаториска синтеза. Потребни се понатамошни студии со кои прецизно ќе се оцени типот на рецепторите одговорен за активноста и механизмот на дејство.