Agarikon.1 and Agarikon Plus Affect Cell Cycle and Induce Apoptosis in Human Tumor Cell Lines

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# INTRODUCTION

- Continuation of the study on 6 blended mushroom products and 3 single extracts on 4 human tumor cell lines (Durgo, Jakopovich 2013)
- Neutral Red and MTT proliferation assays demonstrate that blended extracts cause increased tumor cell membrane and mitochondria damage
- This study concentrates on the mechanisms; effects on the cell cycle and inducing apoptosis

Medicinal mushroom metabolites can interfere and modulate all processes related to the 8 steps of carcinogenesis (Petrova 2012):

- inflammation
- cancer cell proliferation
- adhesion
- apoptosis
- angiogenesis
- gene expression
- invasiveness
- metastasis

Apoptosis - programmed cell death - is a crucial tumor suppression mechanism

 eliminates cells with extensive DNA damage (potentially leading to cancer)

 differentiation - cell growth – apoptosis balance

## **Purpose of the Study**

Investigate:

- A. proliferation (cell cycle disturbance)
- B. induction of apoptosis
- medicinal mushroom extract blends
   Agarikon Plus and Agarikon.1
- concentration response
- 24 and 48 hour response

– 2 human tumor cell lines: H460 (lung carcinoma) and Caco-2 (colon carcinoma)

 Camptothecin, referent antitumor compound; cytotoxic dose 10 µM used

# **MATERIALS AND METHODS**

#### **Cell lines**

- H460 (large cell lung carcinoma)
- Caco-2 (colorectal adenocarcinoma)

#### Tested extract blends

- Agarikon.1 tablets
- Agarikon Plus

#### **Experimental methods**

- Proliferation Assay by MTT
- Cell Cycle Analysis by flow cytometry
- Annexin V Assay for Apoptosis Induction Detection
- Western Blot Analysis

# **TESTED PRODUCTS**

- Agarikon.1 tablets (AG.1)
- proprietary mushroom extract blend from Dr Myko San company
- Ganoderma lucidum, Lentinus edodes, Grifola frondosa, Pleurotus ostreatus, Agaricus brasiliensis
- registered med. mushroom supplement
- recommended treatment dose: ~0.1 g/kg bodyweight per day of soluble polysaccharides



- Agarikon Plus extract blend (AG+)
- proprietary mushroom extract blend from DMS
- 10 medicinal mushroom species (inc. G. lucidum, L. edodes, G. frondosa, P. ostreatus, A. brasiliensis)



- in liquid form
- Recommended treatment dose: ~0.16g/kg BW per day of soluble polysaccharides

# **1 Proliferation Assay**

- Cells cultured as monolayers, plated in parallel on day 0, at 3 x 10<sup>3</sup> cells/well (H460) and 7 x 10<sup>3</sup> cells/well (Caco-2), depending on doubling times
- AG.1 and AG+ added at 0.001, 0.01, 0.1, 1 and 10 mg/ml concentrations (stock solution for both 40 mg/ml, and 4 x 10<sup>-3</sup> M/DMSO for camptothecin)
- We used MTT assay to evaluate cell growth rate after 72 hours (absorbance was measured at 570 nm)

# 2 Cell Cycle Analysis

- seeded at 1x10<sup>5</sup> cells/well (H460) and 2x10<sup>5</sup> cells/well (Caco-2), depending on the doubling times
- After 24 hours, AG.1 and AG+ applied at concentrations 0.1 mg/ml and 1 mg/ml; camptothecin (10 μM) for positive control
- After the incubation period, cells were trypsinized, washed with Phosphate Buffer Saline (PBS); stained with propidium iodide (PI) and analyzed on FACScalibur flow cytometer
- Ratio of cells in each cell cycle phase was determined by analyzing the DNA histograms using ModFit LTTM software

# **3 Annexin V Assay for Apoptosis Induction Detection**

- same concentrations used; 0.1 and 10 mg/ml
- cell populations were gated into regions corresponding to live, early apoptotic and late apoptotic/necrotic cells

Annexin V	ΡΙ	Cell Region
-	-	Live cells
+	-	Early apoptotic
+	+	Late apoptotic /necrotic

# **4 Western Blot Analysis**

- mushroom extracts (0.1 and 1 mg/ml) were added to well plates after 24 hours
- total proteins were measured using BCA Protein Assay Reagent, separated by SDSpolyacrylamid gel electrophoresis and transferred to PVDF membrane → probing with anticaspase 3, anti-p53, and anti-p21 primary antibodies
- equal loading confirmed using anti-tubulin primary antibody

### RESULTS

#### **Proliferation Assay**

- Agarikon Plus (strong effect at 10 mg/ml, GI50 ≈ 2-3 mg/ml) and Agarikon.1 inhibit the growth of both tumor cell lines
- H460 cells are more resistant to Agarikon.1 (approaching GI50 above 10 mg/ml mass concentration)

GI <sub>50</sub> ª (mg/ml)				
Test agent	Cell lines			
	Caco-2	NCI-H460		
Agarikon Plus	1.9 ± 0.1	3.4 ± 1		
Agarikon.1	1.6 ± 0.3	≥10		
<sup>a</sup> GI <sub>50</sub> ; growth inhibition 50 - the concentration that causes 50% growth inhibition				

 Concentration-response curves showing growth inhibition of H460 and Caco-2 cell in vitro after 72 hours after adding Agarikon.1, Agarikon Plus, and camptothecin.



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# **Cell Cycle Analysis**

- Distribution of H460 cells by cell cycle phase: G0/G1, S, G2/M; and sub G1 (dead/apoptotic cells)
- treated with Agarikon Plus, Agarikon.1 (at 0.1 and 10 mg/ml), camptothecin at 10 µM.
- measured by flow cytometry at 24 and 48 hours

## H460 - 24h



no significant influence on cell cycle
moderate increase in apoptotic/dead cells for AG+ at higher concentrations

#### H460 - 48h



 AG+ and AG.1 (1mg/ml) induce accumulation of cells in G1, reduction in S, increase in sub G1 (apoptotic/necrotic)

Caco-2 - 24h



AG+ (1mg/ml): reduced G1, increased S
no significant sub G1 influence

#### Caco-2 - 48h



- AG+, AG.1 (1 mg/ml) reduced G1, increased S phase
- no significant sub G1 influence

# Apoptosis Induction Detection by Annexin V Assay

 Ratio of H460 cells in early or late apoptosis, obtained by co-staining with FITC-labeled annexin V and propidium iodide (PI) and analyzed by flow cytometry.



 AG+ (1 mg/ml): moderate increase in early apoptotic cells

### H460 - 48 h



 AG+ (1 mg/ml): larger increase in late apoptotic/necrotic cells

#### Caco-2 - 24h



 no significant influence, AG+ (1mg/ml) moderate early apoptotic cell increase Caco-2 - 48h



 AG+ (1 mg/ml): moderate increase in induced early apoptosis

# Apoptosis induction detection by caspase-3 cleavage assessment

Appearance of the 17-kDa subunit (caspase-3 p17) – a major cleaved product of the 32-kDa zymogen procaspase-3 - confirms caspase-3 activation which marks the induction of apoptosis.



# The effect of AG+ and AG.1 on cleavage of procaspase-3 in H460 cells (24; 48 hours)

- A. AG+ at 0.1 mg/ml(lanes 1;6), 1 mg/ml (lanes 2;7).
- B. AG.1 at o.1 mg/ml (lanes 3;8), 1 mg/ml (lanes 4;9)
- C. Camptothecin at 10µM (lanes 5;10)



# The effect of AG+ and AG.1 on cleavage of procaspase-3 in Caco-2 cells (24; 48 hours)

- A. AG+ at 0.1 mg/ml (lanes 1;6), 1 mg/ml (lanes 2;7).
- B. AG.1 at o.1 mg/ml (lanes 3;8), 1 mg/ml (lanes 4;9)
- C. Camptothecin at 10µM (lanes 5;10)



# Influence of AG+ and AG.1 on p53 and p21 protein expression

 Agarikon.1, in both concentrations, induced mild p53 protein expression in H460 after 24 hours (A; lanes 3 and 4)





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 Both Agarikon Plus (lane 2) and Agarikon.1 (lane 4) at the concentration of 1 mg/ml induced a moderate increase in the expression of both p53 and p21 after 24 hours in H460  In Caco-2 cells (B), only minor upregulation of p21 protein expression is detected after 48 hour treatment.



# CONCLUSIONS

- Agarikon Plus and Agarikon.1 possess antiproliferative, mainly cytostatic activity, on H460 and Caco-2 cells, in the concentration range 1-10 mg/ml
- Both induce cell cycle perturbations, by delaying the progress through the G1 and S phase
- This points to disturbances occurring before or during DNA replication (confirmed by increase in both p53 and p21 protein expression)

# **CONCLUSIONS (II)**

- Although a modest induction of early (after 24 hours) and late (48 hours) apoptosis was noticed by annexin V test, no processing (cleavage) of caspase-3 was detected
- More-pronounced antiproliferative activity (MTT) of tested agents towards Caco-2 line at maximal concentration (10 mg/ml) points to a non-specific cytotoxic effect

## Acknowledgements

Dr. Marijeta Kralj Dr. Ana-Matea Mikecin Dr. Damir Kralj

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#### Thank you for your attention!

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#### Percentage of Growth definition

The percentage of growth (PG) expressions:

1. For 
$$(A_t - A_0) \ge 0$$
:  
PG=100\* $(A_t - A_0)/(A_c - A_0)$   
2. For  $(A_t - A_0) < 0$ :  
PG = 100\* $(A_t - A_0) / A_0$ 

A<sub>0</sub> ... avg. absorbance before exposure
 A<sub>t</sub> ... avg. absorbance after test (72 h)
 A<sub>c</sub> ... avg. absorbance after 72 h, no
 exposure to test compound

## **Camptothecin Proliferation Assay**

#### Camptothecin

