

1 Article

## 2 **Fucoidan does not exert anti-tumorigenic effects on** 3 **uveal melanoma cell lines**

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### 16 **Abstract**

#### 17 Background

18 The polysaccharide Fucoidan is widely investigated as an anti-cancer agent. Here, we tested the  
19 effect of fucoidan on uveal melanoma cell lines.

#### 20 Methods

21 The effect of 100  $\mu$ M fucoidan was investigated on five cell lines (92.1, Mel270 OMM1, OMM2.3,  
22 [OMM2.53](#) and of 1  $\mu$ g/ml – 1 mg/ml in two cell lines (OMM1, OMM2.3). Cell proliferation and  
23 viability were investigated with a WST-1 assay, migration in a wound healing (scratch) assay.  
24 Vascular Endothelial Growth Factor (VEGF) was measured in ELISA. Angiogenesis was evaluated  
25 in co-cultures with endothelial cells. Cell toxicity was induced by hydrogen-peroxide. Protein  
26 expression (Akt, ERK1/2, Bcl-2, Bax) was investigated in Western blot.

#### 27 Results

28 Fucoidan increased proliferation in two and reduced it in one cell line. Migration was reduced in  
29 three cell lines. [The effect of fucoidan on VEGF was cell type and concentration](#)  
30 [dependent](#)~~Fucoidan did not change the secretion of VEGF~~. In endothelial co-culture with 92.1,  
31 fucoidan significantly increased tubular structures. Moreover, fucoidan significantly protected all  
32 tested uveal melanoma cell lines from hydrogen-peroxide induced cell death. [Under oxidative](#)  
33 [stress](#), fucoidan did not alter the expression of Bcl-2, Bax or ERK1/2, while inducing Akt  
34 expression in 92.1 cells but not in any other cell line.

#### 35 Conclusion

36 Fucoidan did not show anti-tumorigenic effects but displayed protective and pro-angiogenic  
37 properties, rendering fucoidan unsuitable as a potential new drug for the treatment of uveal  
38 melanoma.

39 **Keywords:** Fucoidan; uveal melanoma; VEGF; angiogenesis; oxidative stress

40

## 41 Introduction

42 Uveal melanoma (UM) is the most common primary tumor of the adult eye with an incidence of  
43 4-8 per million in Western countries [1]. It arises from melanocytes of the uvea, the tissue between  
44 the inner retina and the outer scleral layer of the posterior eye, including the iris, ciliary body and  
45 choroid. Most UM arise from the choroid, which provides blood supply and maintenance for the  
46 photoreceptors of the retina. The disease generally occurs in the 6<sup>th</sup> decade of life and primarily  
47 affects fair-skinned people of Caucasian descent [2]. Treatment options for UM depend on the tumor  
48 size and patient choice, but include transpupillary thermotherapy, radiation therapy (including  
49 plaque brachytherapy, proton beam- and gamma-knife radiotherapy), local tumor resection and  
50 enucleation [2]. Radiation therapy is conducted with good success for medium sized tumors,  
51 however, may result in profound vision loss due to side effects [3]. Metastases develop in up to 50%  
52 of UM patients, primarily affecting the liver. The prognosis of these patients is poor, as the current  
53 treatment options for metastatic UM are very limited [1,4]. New treatment option for this disease is  
54 currently of great interest and activity of numerous basic and clinical research teams.

55 A promising new approach in the treatment of cancer is the use of fucoidan, a sulfated  
56 polysaccharide, obtained from the cell-wall matrix of brown algae. Fucoidan contains high amounts  
57 of L-fucose, but has a highly complex structure and may differ substantially depending on different  
58 species, regional origin and even mode of extraction [5]. Fucoidan has been described in several  
59 studies to have anti-tumorigenic properties, e.g. it has been shown to be anti-proliferative and/or  
60 pro-apoptotic on several kind of tumors cells, such as colon cancer [6], hepatoma [7], urinary bladder  
61 cancer cells [8], breast cancer [9], melanoma cells [10] or prostate cancer cells [10,11]. Fucoidan also  
62 has shown anti-angiogenic properties [6,11,12,13] and is discussed as a promising anti-cancer agent  
63 [13,14]. Therefore, fucoidan might be an interesting new therapeutic compound for the treatment of  
64 UM.

65 Important parameters in tumor progression are proliferation, migration and angiogenic  
66 potential [5]. We tested the effect of fucoidan on these parameters in five different UM cell lines. One  
67 of the factors that have been discussed to be involved in the pathogenesis of UM is Vascular  
68 Endothelial Growth Factor (VEGF). VEGF has been reported in UM, ocular fluid of UM patients and  
69 UM cell lines [14,15-17]. A meta-analysis showed that VEGF expression in patients with UM was  
70 significantly higher compared to controls [17,18]. Moreover, VEGF has been elevated in patients with  
71 metastatic UM [16,17], and has been proposed to be a marker for high risk patients [18]. Fucoidan  
72 has been described to reduce VEGF expression in breast cancer cells [9] and in Lewis tumor bearing  
73 mice [19]. Therefore, we investigated the effect of fucoidan on VEGF secretion by UM cells.

74 Oxidative stress is an important factor in tumor pathology and metastasis [19,20,21] and is  
75 utilized by therapeutic compounds to destroy the tumor tissue [21,22]. In primary UM, the tumor is  
76 treated with ionizing radiation, which induces cell death via oxidative stress-mediated killing of  
77 tumor cells [3,22,23]. Therefore, we also tested the effect of fucoidan on UM cells stressed with  
78 Hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>). Fucoidan has been shown to exert its anti-tumor functions via ERK1/2,  
79 Akt [6,10,11,12], Bcl-2 and Bax [8,9,23,24,25]; all these proteins have also been implicated in the  
80 pathogenesis of UM [25-30]. Therefore, we also assessed how fucoidan affect the expression of these  
81 proteins under oxidative stress.

82

## 83 Results

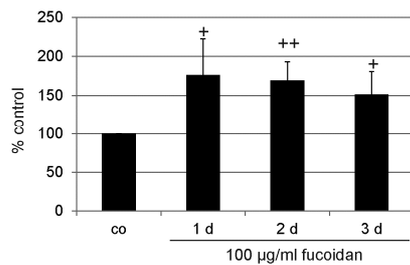
### 84 Proliferation

85 Fucoidan had a cell specific effect on cell proliferation. In 92.1 cells, fucoidan induced a  
86 significant increase in cell number after all one day ( $p < 0.05$ ), two days ( $p < 0.01$ ) and three days ( $p <$

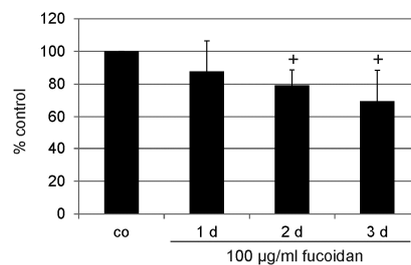
87 0.05) after incubation, while in Mel270 cells, fucoidan reduced proliferation after two and three days  
 88 (both  $p < 0.05$ ). OMM1 and OMM2.3 were not affected by fucoidan, while in OMM2.5 cells, fucoidan  
 89 increased cell number significantly after one day of incubation ( $p < 0.001$ ) (figure 1). In addition, for  
 90 OMM1 and OMM2.3, different concentrations (1  $\mu\text{g/ml}$ , 10  $\mu\text{g/ml}$ , 100  $\mu\text{g/ml}$  1  $\text{mg/ml}$ ) after 1 day of  
 91 incubation were tested. Fucoidan did not show any significant effect in either cell line and either  
 92 concentration (figure 2).

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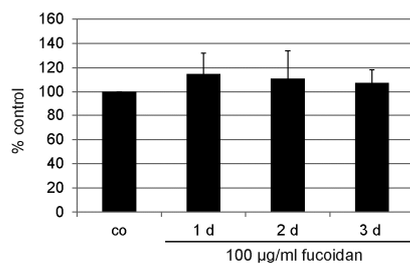
A) 92.1



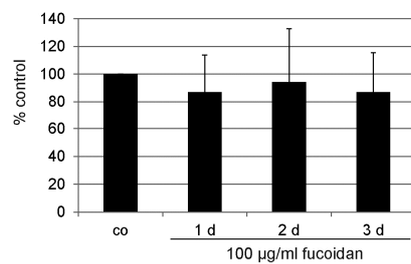
B) Mel270



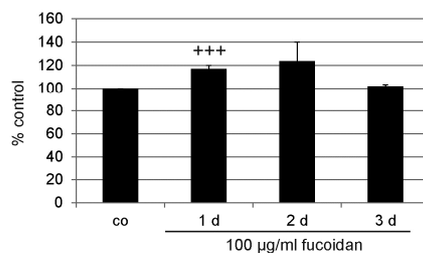
C) OMM1



D) OMM2.3

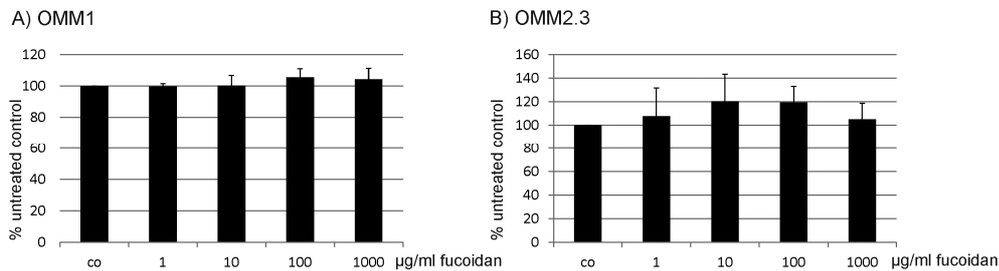


E) OMM2.5



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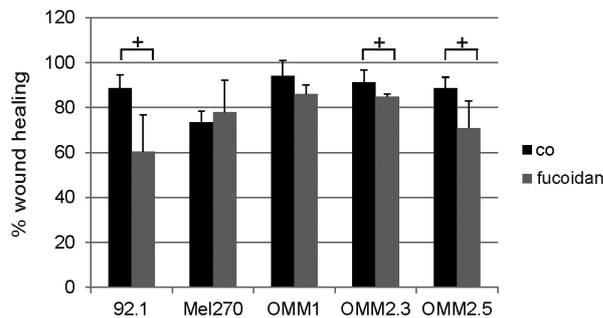
95 **Figure 1:** Proliferation (time line). Proliferation of uveal melanoma cells was tested after incubation  
 96 with fucoidan (100  $\mu\text{g/ml}$ ) for one, two, and three days in A) 92.1, B) Mel 270, C) OMM1, D) OMM2.3  
 97 and E) OMM2.5 cells. Fucoidan exhibited a cell specific effect with an acceleration of proliferation in  
 98 92.1 and OMM2.5 cells, but a decrease in Mel270 cells. Statistical significance was evaluated with  
 99 student's t-test. +  $p < 0.05$  compared to control, ++  $p < 0.01$  compared to control, +++  $p < 0.001$   
 100 compared to control. Co: control.



**Figure 2.** Proliferation (concentration). Proliferation of uveal melanoma cell lines (A) OMM1 and (B) OMM2.3 was tested after 1 day of treatment with 1 µg/ml, 10 µg/ml, 100 µg/ml or 1000 µg/ml. No significant effect on proliferation was found. Statistical significance was evaluated with student's t-test.

### Wound healing/Migration

Fucoidan induced a significant decrease in wound healing ability in 92.1 cells, OMM2.3, and OMM2.5 cells (all  $p < 0.05$ ). No significant effect was seen on Mel270 and OMM1 cells (figure 23).

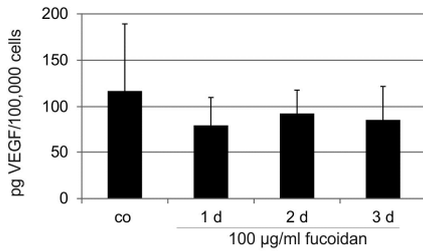


**Figure 23:** Wound healing. Wound healing ability of uveal melanoma cells was tested after incubation with fucoidan (100 µg/ml) for one day in 92.1, Mel 270, OMM1, OMM2.3 and OMM2.5 cells. Fucoidan significantly decreased wound healing in 92.1, OMM2.3 and OMM2.5 cells. Statistical significance was evaluated with student's t-test. +  $p < 0.05$  compared to control. Co = control, fucoidan.

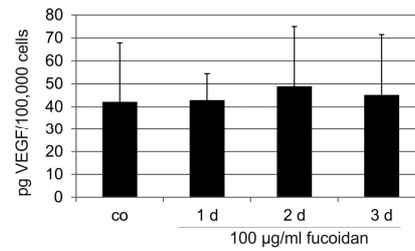
### VEGF Secretion

We have previously shown that all tested UM cell lines secrete VEGF [3432] and that this batch of fucoidan reduces VEGF in retinal pigment epithelial cells in the tested concentration [33]. Fucoidan (100 µg/ml) did not inhibit VEGF secretion in any of the UM cell lines when incubated for up to three days (figure 34). However, these results are dose and cell-line dependent. In a separate set of experiments, we investigated different concentrations of fucoidan (1 µg/ml, 10 µg/ml, 100 µg/ml 1 mg/ml) in OMM1 and OMM2.3 cells after treatment for 1 day. While for OMM2.3 cells, a slight but significant induction of VEGF could be found at 10 and 100 µg/ml, fucoidan at 1 mg/ml significantly reduced VEGF in OMM1 cells (figure 5).

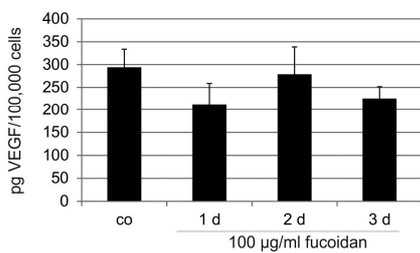
A) 92.1



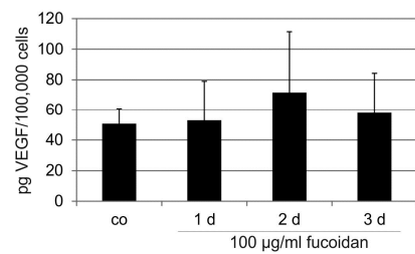
B) Mel270



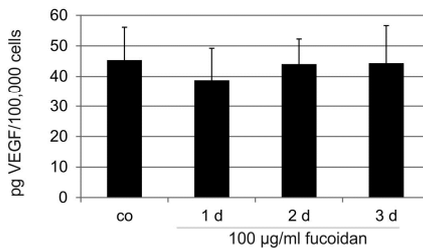
C) OMM1



D) OMM2.3



E) OMM2.5



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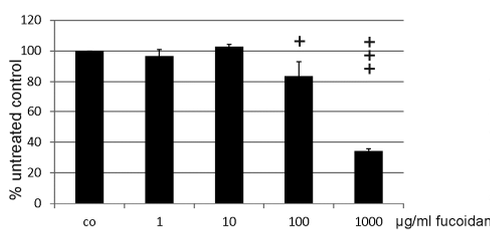
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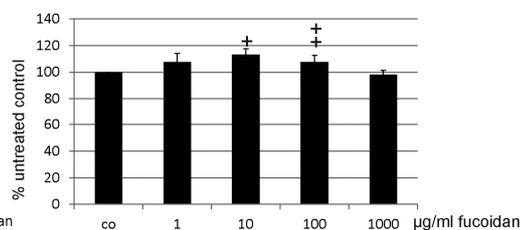
**Figure 34:** VEGF secretion (time line). Influence of fucoidan (100 µg/ml) on VEGF secretion by uveal melanoma cell lines. Treatment with fucoidan for up to three days did not show any significant influence on the secretion of VEGF in any of the cell lines tested (A) 92.1, (B) Mel270, (C) OMM1, (D) OMM2.3, (E) OMM2.5). The secretion of VEGF was determined in VEGF-VEGF-ELISA. Statistical significance was evaluated with student's t-test. Co: control.

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A) OMM1



B) OMM2.3



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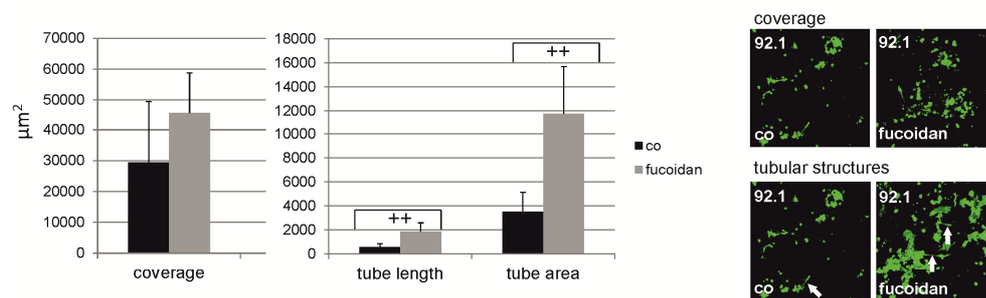
**Figure 5:** VEGF secretion (concentration). Influence of different concentrations of fucoidan (1 µg/ml – 1 mg/ml) on VEGF secretion in (A) OMM1 and (B) OMM2.3 uveal melanoma cell lines. Treatment with fucoidan displayed a dose- and cell-dependent effect with significant reduction of VEGF in

137 OMM1 (100 µg/ml, 1 mg/ml) and a slight but significant induction in OMM2.3 cells (10 µg/ml, 100  
 138 µg/ml). The secretion of VEGF was determined in VEGF-ELISA. Statistical significance was  
 139 evaluated with student's t-test. + p < 0.05, ++ p < 0.01, +++ p < 0.001. Co: control.

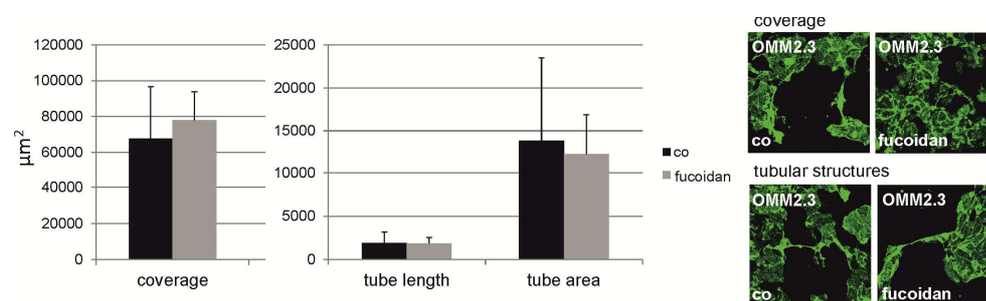
#### 140 Angiogenesis

141 Fucoidan induced an elevation of the tubular area in a co-culture of endothelial cells with 92.1  
 142 cells (p < 0.01). Similarly, fucoidan increased tubular length (p < 0.01). Fucoidan did not, however,  
 143 influence the total area of endothelial coverage in these co-cultures. No effect was seen in co-cultures  
 144 of endothelial cells with the metastatic UM cell line OMM2.3 (figure 46).

##### A) 92.1



##### B) OMM2.3

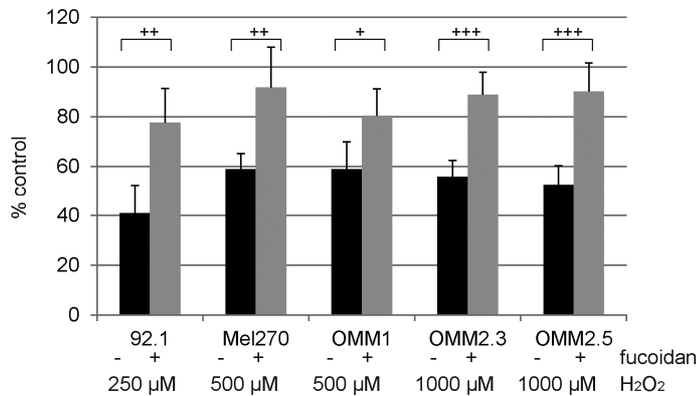


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146 **Figure 46:** Tubular structures in endothelial – uveal melanoma cell line co-culture. Uveal melanoma  
 147 cell line 92.1 and OMM2.3 were co-cultured with outgrowth endothelial cells and subjected to 100  
 148 µg/ml fucoidan. In co-cultures with endothelial cells and 92.1 cell line (A), tubular area and tubular  
 149 length were increased by fucoidan. Total coverage with endothelial cells, however, was not  
 150 influenced. In co-cultures with endothelial cells and OMM2.3 cell line, fucoidan displayed no effect  
 151 (B). Statistical significance was evaluated with student's t-test. ++ p < 0.01 compared to control. Co  
 152 = control, **fuco = fucoidan**.

#### 153 Protection

154 We have previously shown that the UM cell lines have a different susceptibility towards  
 155 H<sub>2</sub>O<sub>2</sub>-induced cell toxicity [31,32]. In all cell lines tested, fucoidan exerted a significant protection on  
 156 UM cell lines under oxidative stress. (92.1, 250 µM H<sub>2</sub>O<sub>2</sub>, p < 0.01; Mel270, 500 µM H<sub>2</sub>O<sub>2</sub>, p < 0.01;  
 157 OMM1, 500 µM H<sub>2</sub>O<sub>2</sub>, p < 0.05; OMM2.3, 1000 µM H<sub>2</sub>O<sub>2</sub>, p < 0.001; OMM2.5, 1000 µM H<sub>2</sub>O<sub>2</sub>, p <  
 158 0.001) (figure 57).



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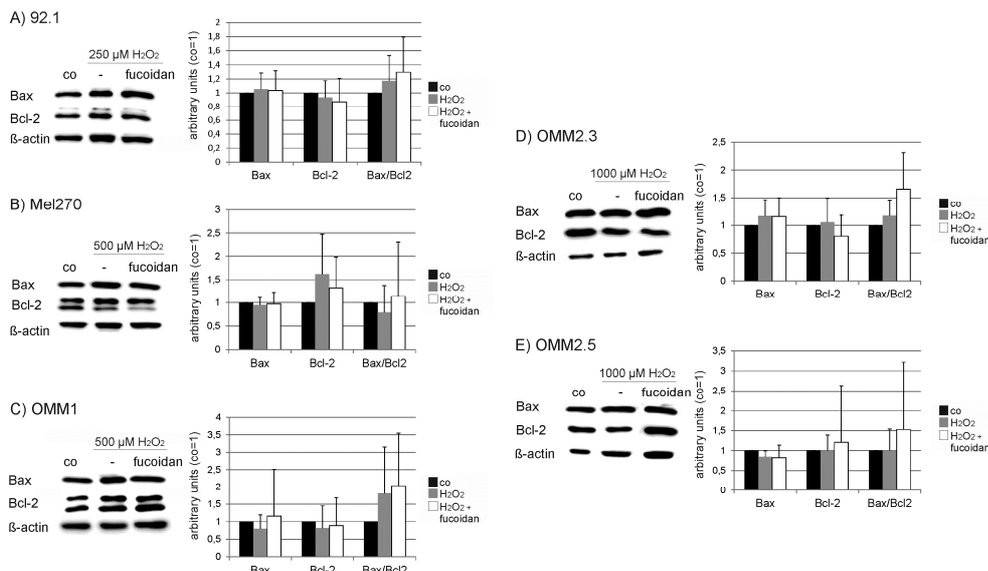
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**Figure 75:** Cell viability of uveal melanoma cell lines under oxidative stress. Uveal melanoma cell lines 92.1, Mel270, OMM1, OMM2.3, and OMM2.5 were subjected to 250  $\mu\text{M}$  (92.1), 500  $\mu\text{M}$  (Mel270 and OMM1) or 1000  $\mu\text{M}$  (OMM2.3 and OMM2.5)  $\text{H}_2\text{O}_2$ . The toxicity of these concentrations of  $\text{H}_2\text{O}_2$  in the respective cell line has been shown previously [324]. The ability of 100  $\mu\text{g/ml}$  fucoidan to protect cell viability after  $\text{H}_2\text{O}_2$  treatment was detected in WST assay. All tested substances exhibited statistically significant protection in all cell lines tested. Statistical significance was evaluated with student's t-test. +  $p < 0.05$ , ++  $p < 0.01$ , +++  $p < 0.001$ .

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**Figure 86:** Expression of Bcl-2 and Bax. Uveal melanoma cell lines (A) 92.1, (B) Mel270, (C) OMM1, (D) OMM2.3, (E) OMM2.5 were subjected to (A) 250  $\mu\text{M}$ , (B, C) 500  $\mu\text{M}$  or (D, E) 1000  $\mu\text{M}$   $\text{H}_2\text{O}_2$ . The effect of 100  $\mu\text{g/ml}$  fucoidan on the expression of Bcl-2 and Bax was investigated in Western blot. Example blots (compound) and densitometric evaluations are shown. Statistical significance was evaluated with student's t-test.

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### Protein expression

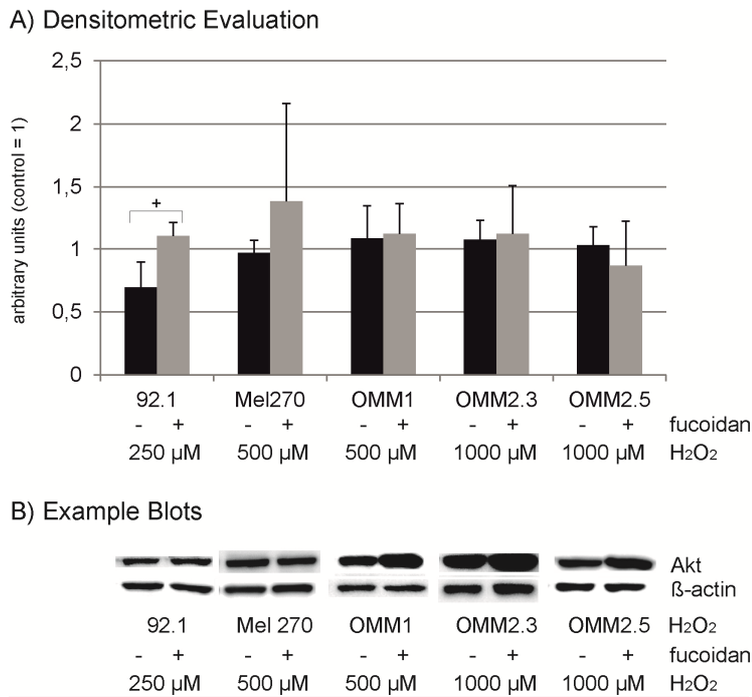
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Under oxidative stress conditions ~~Fucoidan-fucoidan~~ did not show any influence on Bcl-2 or Bax expression in any of the cell lines (figure 68). In 92.1 cell lines, fucoidan induced a significant induction of Akt expression compared to cells treated with  $\text{H}_2\text{O}_2$  alone, while it showed no

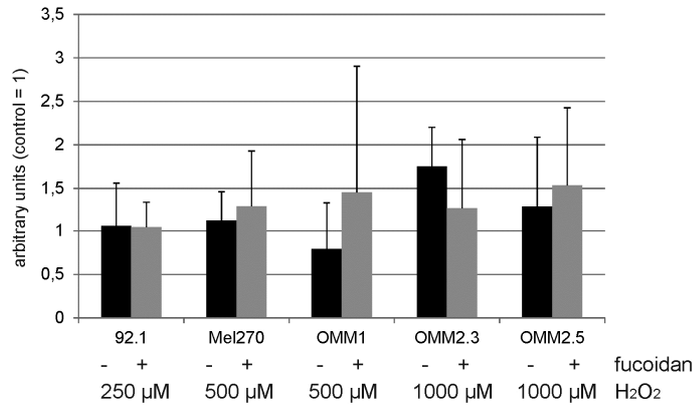
177 significant effect on the other cell lines (figure 79). Considering ERK1/2, no statistically significant  
 178 change in ERK1/2 expression or phosphorylation compared to H<sub>2</sub>O<sub>2</sub>-treated cells can be found  
 179 (figure 810).



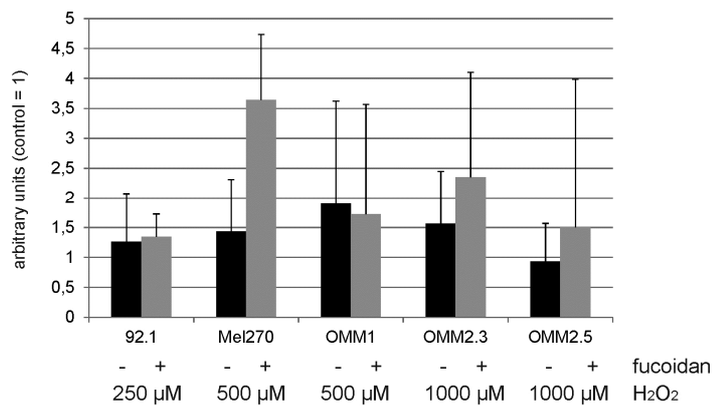
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181 **Figure 79:** Expression of Akt. Uveal melanoma cell lines 92.1, Mel270, OMM1, OMM2.3 and OMM2.5  
 182 were subjected to 250 μM (92.1), 500 μM (Mel270 and OMM1) or 1000 μM (OMM2.3 and OMM2.5)  
 183 H<sub>2</sub>O<sub>2</sub>. The effect 100 μg/ml fucoidan on the expression of Akt was investigated in Western blot.  
 184 Densitometric evaluations (A) and example blots (compound) (B) are shown. Statistical significance  
 185 was evaluated with student's t-test. + p < 0.05.

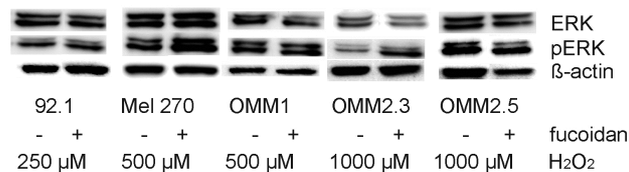
## A) Densitometric Evaluation ERK(42 kDa)



## B) Densitometric Evaluation pERK(42 kDa)



## C) Example Blots



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187 **Figure 810:** Expression and phosphorylation of ERK1/2. Uveal melanoma cell lines 92.1, Mel270,  
 188 OMM1, OMM2.3 and OMM2.5 were subjected to 250  $\mu$ M (92.1), 500  $\mu$ M (Mel270 and OMM1) or 1000  
 189  $\mu$ M (OMM2.3 and OMM2.5)  $H_2O_2$ . The effect of 100  $\mu$ g/ml fucoidan on the expression and  
 190 phosphorylation of ERK1/2 was investigated in Western blot. Densitometric evaluation of (A) ERK  
 191 and (B) pERK blots are shown for the 42 kDa isoform. (C) Example blots (compound). Statistical  
 192 significance was evaluated with student's t-test.

## 193 Discussion

194 Fucoidan has been shown to display a variety of anti-tumor effects on several types of tumors  
 195 or cancer cell lines. Here, we investigated its effect on UM, a primary malignant neoplasm of the eye.  
 196 We investigated classical parameters, such as proliferation, migration, VEGF secretion and

197 angiogenesis and additionally investigated the effect of fucoidan on H<sub>2</sub>O<sub>2</sub>-induced cell death and  
198 protein expression.

199 Anti-proliferative activity of fucoidan has been shown for several cancer cell types, such as  
200 bronchopulmonary carcinoma [342], cutaneous melanoma cells [10,353], bladder cancer cells [8],  
201 breast cancer cells [9], or B-cell lymphoma [364]. In our study, fucoidan reduced proliferation in one  
202 cell line (Mel 270), but, surprisingly, it induced proliferation in two cell lines (92.1 and OMM2.5). In  
203 OMM1 and OMM2.3 cells, both time line (100 µg/ml) and concentration (1 µg/ml – 1 mg/ml) was  
204 tested and no effect on proliferation was seen at either time point or contraction. The effect  
205 therefore is clearly cell type specific. Moreover, the pro-proliferative effect on two cell lines would be  
206 a worrisome result if fucoidan were to be used in UM patients.

207 Fucoidan decreased its wound healing ability in three (92.1; OMM2.3; OMM2.5) out of five UM  
208 cell lines. This indicates that fucoidan interferes with migration in these cell lines, especially as  
209 wound healing assay measure both proliferation and migration, and we found fucoidan to induce  
210 proliferation in 92.1 and OMM2.5 cells. Fucoidan has been shown to inhibit migration e.g. in colon,  
211 lung or bladder cancer cells [6,375,3638]. Again, the effect is cell-type dependent, and no general  
212 anti-migratory effect of fucoidan could be shown here.

213 When we tested the ability of fucoidan to reduce the availability of VEGF, no reduction of VEGF  
214 could be seen at a concentration of 100 µg/ml. This is in contrast to our findings in retinal pigment  
215 epithelial (RPE) cells [337] where we could find a significant reduction of detectable VEGF at this  
216 concentration using the same batch of fucoidan, and A similar reductive effect of VEGF expression  
217 by fucoidan has been shown for breast cancer cells [9]. It is of interest to note that the lack of effect on  
218 VEGF by fucoidan in this system cannot be solely due to the molecular properties of the fucoidan, as  
219 the exact same fucoidan has been shown to reduce VEGF secretion interactions in RPE cells [37].  
220 Therefore, the effect of fucoidan is not only determined by the molecular structure of the fucoidan  
221 [5], but also by the target cells. As higher concentrations of fucoidan did reduce VEGF in OMM1  
222 cells, possibly the concentration chosen in this experiments were too low to exert an effect. However,  
223 even in higher concentrations, the effect was cell type dependent, as OMM2.3 did not show any  
224 reduction of VEGF in any of the fucoidan concentrations tested. The pathways of fucoidan-mediated  
225 VEGF reduction have not been elucidated to date, but it has been shown that fucoidan can inhibit the  
226 activation of VEGFR-2 by preventing the binding of VEGF165 to its receptor [3839]. We have  
227 previously shown that VEGF is autoregulated via the VEGFR-2 in RPE cells [3940], and so we  
228 hypothesized that the downregulation of VEGF was mediated by interfering with the  
229 autoregulatory pathway. The cell dependent effect in ability of fucoidan to reduce concerning VEGF  
230 in the UM cells may therefore be related to the presence of due to the lack of an autoregulatory  
231 pathway of VEGF expression in the tested melanoma cells.

232 In addition, in our angiogenesis assay, fucoidan induced the outgrowth of tubular structures,  
233 both in length and area, in 92.1 cells. Even though the general interaction between 92.1 and  
234 endothelial cells were low, this result may indicate that fucoidan may facilitates angiogenesis  
235 primary UM, which would not be desirable in patient treatment. Again, this cannot simply be  
236 explained by the molecular structure of this particular fucoidan, as we have shown before that this  
237 exact fucoidan reduced angiogenic structures in RPE-endothelial cells co-cultures [3733].

238 Fucoidan displayed a significant protective effect against H<sub>2</sub>O<sub>2</sub>-induced cell death in all tested  
239 cell lines. Fucoidan has been described to protect cells against oxidative stress [40,41,42]; however, to  
240 the best of our knowledge, this has not been shown in cancer cells before. Indeed, fucoidan when  
241 given in addition with a chemotherapeutic has been shown to increase oxidative stress in breast  
242 cancer cell [4243]. Antioxidants may enhance tumor progression [4920] and oxidative stress may  
243 protect from metastasis [2021], so the protection of cancer cells against oxidative stress by fucoidan  
244 has to be taken into consideration when discussing fucoidan-derived drugs as possible new cancer  
245 agents [4314]. Our data showed that the protective effects of fucoidan are not mediated via a change  
246 in the Bcl-2/Bax expression, or via the ERK1/2 or Akt pathway. Further research needs to be  
247 conducted in order to decipher the protective pathways of these compounds.

248 Fucoidan is also under investigation to be used in combination with other chemotherapeutic  
249 drugs in order to enhance their efficacy, as seen in e.g. melanoma [44] or breast cancer cells [43],  
250 where pro-apoptotic or anti-proliferative effects of the chemotherapeutics are enhanced by fucoidan.  
251 The results found in our study cannot extrapolated towards combination treatments, however, also  
252 in combination treatments, the effect of fucoidan is cell type dependent and may reduce the efficacy  
253 of the chemotherapeutic compound [45]. Moreover, it has been suggested that the  
254 apoptosis-enhancing effects of combination therapies combining fucoidan and chemotherapy is  
255 mediated by oxidative stress-enhancement by fucoidan [43], while our data show that fucoidan  
256 protects against oxidative stress. Therefore, our data cannot give a prediction about potential  
257 combination therapies in UM, but would strongly advice for caution.

## 258 Conclusion

259 The data obtained in this study ~~strongly indicates~~ that fucoidan is not suitable as a potential  
260 treatment for UM.

## 261 Material and Methods

### 262 Cell culture of melanoma cells

263 Five established human UM cell lines were used. The cell lines 92.1 [4346] and Mel270 [474]  
264 originated from primary UM, while all OMM cell lines are of metastatic origin; OMM2.5 and  
265 OMM2.3 from liver metastases [4447] and OMM1 from a sub-cutaneous metastasis [4548]. Cell  
266 cultures were maintained in RPMI (PAA Laboratories, Cölbe, Germany), supplemented with 10%  
267 fetal calf serum (FCS) (Linaris, Dossenheim, Germany) and 1% penicillin/streptomycin (PAA).  
268 Medium was exchanged three times a week and cells were passaged after reaching confluence

### 269 Fucoidan

270 For the experiments, fucoidan from Sigma Aldrich (from Fucus vesiculosus, Sigma Aldrich,  
271 Steinheim, Germany; #F5631, [O28K3779; CAS 9072-19-9]) was used.

### 272 Proliferation

273 To determine the influence of fucoidan on proliferation, a defined number (200,000 cells) of the  
274 respective cell line was seeded on 12 well plates. Cells were stimulated with 100 µg/ml fucoidan (~~F-~~  
275 ~~vesiculosus, Sigma Aldrich, Steinheim, Germany) for up to three days for one day, respectively. In~~  
276 ~~addition, for the cell lines OMM1 and OMM2.3, a dose-response curve after 24 hours of incubation~~  
277 ~~was determined, investigating 1 µg/ml, 10 µg/ml, 100 µg/ml and 1 mg/ml fucoidan. After the~~  
278 indicated period of time, a WST- assay was conducted.

### 279 WST-assay

280 Treated cells as described above were treated with WST-1 reagent (Hoffmann-La Roche, Basel,  
281 Switzerland) for 4 hours at 37°. The cells were rocked on a shaker for 2 min, the supernatant was  
282 collected, and measured at 450 nm.

### 283 Scratch Assay

284 The scratch assay was conducted as previously described with modifications [3733]. In brief,  
285 the respective cell line was seeded in a 12-well-plate. Two wounds were scratched in the confluent  
286 cell layer with a pipette tip and the cells were washed with PBS to remove detached cells.  
287 Microscopic bright field pictures of three spots were taken (AxioCam, Zeiss, Jena, Germany).  
288 Fucoidan (100 µg/ml) was added to the wells. After 90% wound closure of the control, another  
289 picture was taken. To analyse the wound healing capability of the cells, application was conducted  
290 in duplicates and three pictures per well were taken. The gap size of the wound was measured with

291 AxioVision Rel.4.8. (Zeiss, Jena, Germany), and the percentage of coverage of the wound was  
292 evaluated. Complete coverage was defined as 100%.

### 293 VEGF-ELISA

294 The supernatant of cell cultures was collected after 100  $\mu$ M fucoidan incubation for up to three  
295 days. In addition, for the cell lines OMM1 and OMM2.3, a dose-response curve after 24 hours of  
296 incubation was determined, investigating 1  $\mu$ g/ml, 10  $\mu$ g/ml, 100  $\mu$ g/ml and 1 mg/ml fucoidan. ~~and~~  
297 VEGF-content was measured by VEGF-ELISA (R&D Systems, Wiesbaden, Germany), following the  
298 manufacturer's instructions. The range of detection of the ELISA was between 15 pg/ml and 1046  
299 pg/ml. The amount of VEGF secreted was normalized to cell number. Cell number was assessed  
300 with a trypan blue exclusion assay.

### 301 Angiogenesis Assay

302 Angiogenesis was evaluated in a direct co-culture system of UM cells and outgrowth  
303 endothelial cells.

304 The isolation of outgrowth endothelial cells from peripheral blood was conducted as described  
305 previously [46,47,49,50]. In brief, these cells were isolated from buffy coats by isolation of blood  
306 mononuclear cells. Mononuclear cells were seeded onto collagen coated 24-well plates in a density  
307 of  $5 \times 10^6$  cells/well in EGM-2 (Lonza, Basel, Switzerland) with full supplements from the kit, 5%  
308 FCS, and 1% penicillin/streptomycin. After one week, adherent cells were collected by trypsin  
309 and reseeded on collagen coated 24-well plates in a density of  $0.6 \times 10^6$  cells/well. After 2-3 weeks,  
310 colonies of endothelial cells (OEC) were harvested and further expanded over several passages  
311 using EGM-2 in a splitting ratio of 1:2.

312 Co-culture assays were performed for one primary (92.1) and one metastatic (OMM2.3)  
313 melanoma cell line. For co-cultures 100,000 cells/cm<sup>2</sup> were seeded into fibronectin coated thermanox  
314 coverslips in 24 well plates in their respective cell culture medium. On the next day outgrowth  
315 endothelial cells (OEC) were added to the cultures in a density of 100,000 cells/cm<sup>2</sup> to the respective  
316 uveal melanoma cell line and co-cultures were further maintained for 7 days in EGM-2 treated with  
317 100  $\mu$ g/ml fucoidan, respectively, or left untreated in control groups. After 7 days co-cultures were  
318 fixed with 4% paraformaldehyde and outgrowth endothelial cells were immunostained for the  
319 endothelial marker CD31. All cells are counterstained by Hoechst and pictures were taken with a  
320 confocal laser scanning microscope (Zeiss LSM 510 Meta, Jena, Germany). Angiogenesis was  
321 evaluated in comparison to untreated controls. For each group, at least 3 pictures were taken from  
322 two technical replicates. These experiments and the picture analysis were performed with  
323 endothelial cells from three different donors.

### 324 Image Analysis

325 The microscopic images were analyzed using the image processing program ImageJ Vers. 1.47  
326 and GIMP 2.8. The analysis of angiogenic structures was conducted as previously described [48,51].  
327 In brief, tube-like structures were extracted from the background by automatic segmentation after  
328 background correction. The binaries of the tube-like structures were further processed, including a  
329 final manual correction. The resulting binaries were analyzed for the area and the length of tubular  
330 structures. Additionally, the total area of fluorescence was assessed after automatic segmentation.

### 331 Cytotoxicity

332 Cells were plated on 24-well plates. H<sub>2</sub>O<sub>2</sub> (Sigma-Aldrich) was applied in order to induce  
333 oxidative stress mediated cytotoxicity. We have previously shown that uveal melanoma cell lines  
334 show a cell-line specific susceptibility to oxidative stress [34,32]. Cytotoxicity was induced by  
335 applying H<sub>2</sub>O<sub>2</sub> in the respective concentration (92.1: 250  $\mu$ M, Mel270 and OMM1: 500  $\mu$ M, OMM2.3  
336 and OMM2.5: 1000  $\mu$ M). In order to evaluate a potential protective effect of fucoidan, confluent cells

337 were treated 30 min prior to oxidative insult with 100 µg/ml fucoidan. Cell viability was assessed  
338 after 24 hours of stimulation with a WST assay.

#### 339 *Whole cell lysate*

340 After treatment of cells as indicated, whole cell lysates were prepared in an NP-40 buffer as  
341 described previously [3733]. In brief, cells were washed with PBS and NP-40 buffer (1% Nonidet®  
342 P40 Substitute, 150 mM NaCl, 50 mM Tris, pH 8.0) was added. The lysates were kept on ice for at  
343 least 30 minutes. Lysates were centrifuged at 13,000 rpm for 15 minutes and the supernatant  
344 harvested. The protein concentration of the supernatant was determined by a BioRad protein assay  
345 (BioRad, München, Germany) with bovine serum albumin (Fluka, Buchs, Switzerland) used as  
346 standard.

#### 347 *Western Blot*

348 Western blot was conducted as described previously with modifications [4952]. In brief,  
349 proteins were separated in an SDS-PAGE, using 12 % acrylamide gels. Gels were blotted on  
350 PVDF-membranes (Carl Roth GmbH, Karlsruhe, Germany) and then blocked in 4 % skim milk in  
351 Tris buffered saline with 0.1 % Tween for 1 hour at room temperature. The blot was treated with the  
352 first antibodies, beta-actin (#4967, 1:1000), Akt (#9272, 1:1000), ERK1/2 (#9102, 1:1000), p-ERK1/2  
353 (#9101, 1:1000) (all Cell-Signaling Technologies, CST, Denver, USA; all rabbit), Bax (sc-20067, 1:1000)  
354 or Bcl-2 (sc-509, 1:1000) (all Santa Cruz, Heidelberg, Germany, all mouse), respectively, in 2% skim  
355 milk in Tris buffered saline with 0.1% Tween overnight at 4°C. After washing the blot, it was  
356 incubated with appropriate secondary antibody (anti-rabbit (#7074) or anti-mouse (#7076) IgG,  
357 HRP-linked antibody (all Cell-Signaling)) in 2% skim milk in Tris-buffered saline with 0.1% Tween  
358 (Merck, Darmstadt, Germany). Following the final wash, the blot was incubated with Immobilon  
359 chemiluminescence reagent (Merck), and the signal was detected with MF-ChemiBis 1.6 (Biostep,  
360 Jahnsdorf, Germany). The density of the bands was evaluated using Total lab software (Biostep) and  
361 the signal was normalized for β-actin.

#### 362 *Statistics*

363 Statistical analysis was performed with MS-Excel. Means ± standard deviation (sd) was  
364 calculated for at least 3 independent sets of experiments. Significant differences between means  
365 were calculated by t-test. A p-value of 0.05 or less was considered significant.

#### 366 *Supplementary Materials*

367 None

368

#### 369 *Acknowledgements*

370 This study was supported by the Werner and Klara Kreitz - Foundation.

371 We would like to thank Serap Luick and Andrea Hethke for their excellent technical assistance.

372

#### 373 *Author Contributions*

374 AK, MD and JH conceived and designed the experiments. SEC and JH contributed materials

375 MD, SF, ER and HS performed the experiments. MD, AD, SF, HS, SEC, ER and AK analyzed the data.

376 AK wrote the paper, which was reviewed by all authors.

377

#### 378 *Conflicts of Interest*

379 The authors declare no conflict of interest. The Werner and Klara Kreitz - Foundation had no role in the design  
380 of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the  
381 decision to publish the results.

382

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