

1 **Cigarette Smoke Reduces the Efficacy of Cisplatin in Head and Neck**  
2 **Cancer Cells – Role of ABCG2**

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27           **Abstract**

28           Background/Aim: We evaluated the influence of smoking on HNSCC which are in their  
29 majority tobacco-driven as tobacco smoke is expected to influence the expression of ABCG2-  
30 transporters involved in MDR. The aim was to evaluate the effect of cigarette smoke on the  
31 expression of the ABCG2-transporter in representative HNSCC cell lines, the influence of CSC  
32 on ABCG2 and its expression levels in HNSCC.

33           Materials and methods: HNSCC cell lines were treated with cigarette smoke condensate  
34 (CSC) and basal and induced ABCG2 expression was examined. The impact of CSC on cellular  
35 viability/proliferation during cytotoxic drug treatment was also evaluated. ABCG2 expression  
36 levels in HNSCC were correlated with the smoking history of patients.

37           Results: HNSCC cells showed low basal ABCG2 expression. CSC treatment resulted in  
38 increased ABCG2 expression threefold and in resistance to cisplatin. Tumor samples of never  
39 smokers showed significantly higher ABCG2 expression compared to ever smokers. ABCG2  
40 expression correlated with pack years of cigarette consumption.

41           Conclusion: Tobacco consumption is linked to an inducible and increased ABCG2 protein  
42 expression and has an impact on drug resistance.

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46 Head and neck squamous cell carcinomas (HNSCC) comprise two major, distinct cancer  
47 entities referred to as “classic” tobacco-driven tumors and cancers that result from persistent  
48 infections with human papillomaviruses (HPV) predominantly arising in the oropharynx (1, 2).  
49 “Classical” HNSCC are induced by a multitude of chemical carcinogens contained in tobacco  
50 smoke, which lead to genetic alterations, accumulation of mutations, and ultimately malignant  
51 transformation of epithelial cells (3). In HNSCC caused by persistent HPV infections, viral  
52 oncoproteins induce proteolytic degradation and inactivation of the prominent tumor  
53 suppressors p53 and pRb in host cells, thus preserving the replicating potential of infected cells  
54 and ensuring viral reproduction (4). Accordingly, genetic alterations and mutations are found to  
55 a lesser extent in HPV-positive HNSCC (5).

56 In 1993 Browman and colleagues reported that patients with head and neck cancer who  
57 continue to smoke during radiation therapy have lower rates of response and survival than  
58 patients who stopped smoking (6). In a study of more than 1800 HNSCC patients who received  
59 either postoperative or definitive (chemo-) radiotherapy, smoking during treatment led to  
60 significantly lower local control and survival rates (7). Gillison and co-workers showed that the  
61 risk of progression or death increases by 1% per pack-year or 2% per year of smoking in  
62 patients with oropharyngeal cancer treated with radiotherapy and chemoradiation (8).

63 According to the evidence reviewed in the 2014 Surgeon General’s Report, continued  
64 smoking by cancer patients represents a major cause of increased overall mortality, cancer-  
65 specific mortality and risk of second primary carcinomas. This is the result of increased tumor  
66 aggressiveness and/or altered response to treatment (9).

67 As reviewed by Warren et al. numerous molecular changes were found in cancer cells  
68 exposed to cigarette smoke. It was shown to increase ligand-dependent activation and  
69 autophosphorylation of epidermal growth factor receptor leading to increased proliferation. It  
70 modifies the regulation of cell cycle and senescence, and activates the expression of vascular  
71 endothelial growth factor and matrix metalloproteinases. Long-term exposure of breast cancer  
72 cells to cigarette smoke enhanced their invasiveness and metastatic potential (10).

73           Approximately 30% of primary and 70% of recurrent HNSCC show resistance to cytotoxic  
74 drugs (11). ATP-binding cassette (ABC) transporters play a critical role in the development of  
75 this resistance. ABC transporters are transmembrane proteins that extrude many different,  
76 structurally divergent substrates. ABCG2, also known as breast cancer resistance protein  
77 (BRCP), was initially cloned from drug resistant breast cancer cells and was later detected on  
78 cancer cells from several solid tumors (12, 13). ABCG2 expressing head and neck cancer cells  
79 were shown to be resistant to cisplatin (14).

80           The term *multidrug resistance* (MDR) originates from the discovery of ABC proteins on  
81 cancer cells resistant to various cytotoxic drugs. MDR is inducible. Malignant cells that  
82 developed resistance to a single anticancer drug often show cross-resistance to other drugs.  
83 ABC transporters are part of a general defense against xenobiotics in living organisms and  
84 highly conserved in evolution. In the human body, ABC transporters are particularly expressed  
85 in tissues that operate as interfaces between the organism and the environment(15).

86           Being chronically exposed to cigarette smoke, a complex mixture of numerous chemicals  
87 including more than 60 proven carcinogens, it seems reasonable, that epithelial cells in the  
88 upper aerodigestive tract learn to defend themselves from these harmful compounds. ABCG2  
89 extrudes the major tobacco carcinogen benzo(a)pyren and its conjugates and might play a  
90 critical role in defense against mutagenesis and carcinogenesis in the upper aerodigestive tract  
91 (16, 17). Tobacco smoke-induced ABCG2 expression in epithelial cells could be a cause of  
92 resistance to anticancer drugs after malignant transformation. Cigarette smoke has been shown  
93 to increase ABCG2 expression in esophageal, lung and head and neck cancer cells (18, 19).

94           The aim of the study was to evaluate the effect of cigarette smoke condensate (CSC) on  
95 ABCG2 expression on cancer cells, to demonstrate adverse effects of cigarette smoke during  
96 anticancer treatment *in vitro* and to assess the prevalence of ABCG2 expression in HNSCC. We  
97 treated HNSCC cell lines (FaDu and PiCa) with CSC and examined basal and induced ABCG2  
98 expression. We then evaluated the impact of CSC on cellular viability and proliferation during  
99 cytotoxic drug treatment. We also detected ABCG2 by immunohistochemical staining in tissue  
100 samples of HNSCC and correlated expression levels with smoking history of patients.

101 Furthermore, we analyzed a selection of cases of *the cancer genome atlas* (TCGA) for ABCG2-  
102 mRNA in correlation to smoking history.

103

## 104 **Materials and Methods**

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### 106 ***Fluorescence-activated cell sorting***

107 All FACS measurements were performed on an LSRII cytometer, and the data were  
108 analyzed with FACSDiva Software. ABCG2 expression was determined using the BV421  
109 Mouse Anti-Human CD338 antibody and the isotype control BV421 Mouse IgG2b, κ.

110 Cell lines were cultured in DMEM, supplemented with 10% fetal bovine serum (FBS) and  
111 1% penicillin-streptomycin. CSC was added at a final concentration of 10 µg/ml for 24 h to 48 h.  
112 At the indicated time points, cells were harvested and stained with either ABCG2-specific or  
113 isotype control antibodies.

114 The cell line FaDu was obtained from *Leibniz Institute German Collection of*  
115 *Microorganisms and Cell Culture* (DSMZ). The identity of all cell lines was validated by STR-  
116 typing. The cell line PiCa was generated in-house from surgical specimens (20).

117 ABCG2 expression levels were evaluated as ratio to untreated controls.

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### 119 ***Cell culture***

120 PiCa and FaDu cells were kept in a humidified incubator at 37.5°C and 5% CO<sub>2</sub> (20, 21).  
121 To generate spheroids, 1x10<sup>3</sup> cells were seeded on a 96 well ultra-low attachment plates and  
122 cultured for 1 week.

123

### 124 ***Chemicals***

125 Cigarette smoke condensate (CSC) was obtained from *Chemisches und*  
126 *Veterinäruntersuchungsamt*, (Sigmaringen, Germany). Total smoke of 40 filter cigarettes was  
127 passed through a filter, which was then cut into small pieces, placed in a 50 ml tube and  
128 covered with 30 ml DMSO. After 24 h, 20 ml DMSO were added, the solution was filtered and  
129 diluted until a concentration of 10 mg/ml CSC was reached. Dose-response incubations were  
130 performed using six different concentrations of CSC (10, 20, 30, 50, 70 & 100 µg/ml), cisplatin

131 (1, 5, 10, 25, 50 & 100  $\mu$ M) and 5-fluoruracil (5-FU; 1, 10, 30, 50, 100 & 1000  $\mu$ M) in monolayer  
132 cultures.

133

#### 134 ***Determination of resistance to chemotherapy***

135 Being kept in DMEM, cells were co-incubated with CSC and cisplatin or 5-FU,  
136 respectively, for 24 h. Cultures were subsequently washed using phosphate-buffered saline and  
137 re-suspended in DMEM/BEGM. We further incubated cells with cisplatin or 5-FU for 24 h to  
138 secure that cytotoxic drugs were present during a complete doubling time. Cell cultures were  
139 washed again in PBS and dissociated using StemPro Accutase Cell Dissociation reagent.  
140 Single cells were suspended in DMEM and transferred to a 96 well standard plate for adhesion  
141 during 24 h. Cellular viability was determined using the colorimetric WST-8 cell viability kit  
142 according to the manufacturer's instructions. Cellular proliferation was evaluated by the  
143 colorimetric cell proliferation BrdU-kit.

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#### 145 ***Immunostaining***

146 Snap-frozen tissue samples were processed to 4 $\mu$ m consecutive sections. The ABCG2-  
147 specific antibody was used for immunohistochemical detection of antigens by using the avidin-  
148 biotin-peroxidase method (22). Staining intensity was assessed in a semiquantitative manner  
149 (Figure 1).

150

#### 151 ***ABCG2 expression in TCGA HNSCC patients***

152 To characterize the mRNA expression of the ABCG2 gene in the TCGA HNSCC cohort,  
153 RNAseq gene quantification data of primary tumors were downloaded from the Genomic Data  
154 Commons GDC data portal. Curated clinical data of appropriate patients were downloaded from  
155 CBioportal. The following subgroups were extracted from the TCGA HNSCC cohort: 19 patients  
156 with smoking history status "current", 23 "former smoking" patients and 19 "never smoking"  
157 patients. Differences in ABCG2 expression in the three groups was examined in all patients with  
158 known HPV status, in HPV-positive only and HPV-negative only.

159

160 ***Statistical analysis***

161 All results were tested regarding normal distribution using the Kolmogorov-Smirnov test.  
162 For normally distributed variables, we applied the student's t-test, whereas the Wilcoxon- and  
163 Kruskal-Wallis test we used for non-normally distributed parameters. Associations between  
164 ordinal and nominal scaled factors were evaluated using the Chi-square test.

165  $p$ -Values  $<0.05$  were considered statistically significant. To analyze the results of the  
166 incubation experiments, multiple testing between cell lines and treatment groups was  
167 necessary. Because of multiple testing within the groups, the  $p$ -level was set to 0.005 according  
168 to the Bonferoni correction.

169

## 170 **Results**

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### 172 **ABCG2 expression in HNSCC cell lines**

173 Two HPV-negative HNSCC cell lines (PiCa, FaDu) were assessed for ABCG2 expression  
174 levels. All cell lines were either kept untreated or treated with CSC (10 µg/ml) for 24 h and 48 h.  
175 Expression levels are given as the difference between the ABCG2 and isotype control  
176 antibodies. Basal expression levels were low for Pica and FaDu. PiCa were produced from a  
177 laryngeal carcinoma of a current smoker and, thus, represent tobacco-driven HNSCC (23).  
178 Basal ABCG2 expression was low in this cell line with a significant 2-fold induction after  
179 incubation with CSC for 24 h ( $p<0.001$ , t-test) and a 3-fold induction after incubation with CSC  
180 for 48 h (both  $p<0.001$ , t-test). This behavior of HPV-negative HNSCC was confirmed by  
181 analysis of FaDu cells, which also showed a significant 3-fold induction of ABCG2 after  
182 incubation with CSC for 48 h (figure 2,  $p<0.001$  both 24 h and 48 h, t-test). In both cell lines, the  
183 increase of the ABCG2 signal from 2-fold to 3-fold between 24h and 48h of incubation with CSC  
184 was significant (PiCa  $p=0.003$ , FaDu  $p=0.002$ , t-test).

185

### 186 **Impact of cigarette smoke condensate (CSC), cisplatin and 5-FU on cellular** 187 **viability and proliferation**

188 To exclude major cytotoxicity caused by CSC we chose a concentration of 20 µg/ml, as  
189 the viability of PiCa monolayer cultures continuously decreased from 94.5% (10 µg/ml) to 38.3%  
190 (100 µg/ml) after 24 h of incubation. In order to assess if CSC influences tumor cell treatment,  
191 PiCa and FaDu HNSCC cell lines were treated with CSC and cisplatin or 5-FU, respectively, in  
192 the indicated combinations. CSC had no significant influence on the viability and proliferation of  
193 PiCa [Figure 3,  $p=0.654$ , Wilcoxon signed-rank test (WSRT)] and FaDu cultures (Figure 3,  
194  $p=0.021$ , WSRT). In PiCa cells, 20 µg/ml CSC significantly decreased proliferation ( $p=0.002$ ,  
195 WSRT).

196 As seen in Figure 4, incubation with cisplatin led to a decrease of spheroid size in both  
197 cell lines. It also significantly reduced viability and proliferation compared to untreated controls  
198 in both cell lines (Figure 3; all  $p < 0.001$ , WSRT). It was more cytotoxic in PiCa compared to  
199 FaDu cells ( $p < 0.001$ , Mann-Whitney-U test). No such difference was seen regarding  
200 proliferation ( $p = 0.015$ , Mann-Whitney-U test).

201 5-FU also significantly decreased viability and proliferation of both cell lines compared to  
202 untreated controls (Figure 3; all  $p < 0.001$ , WSRT). PiCa cells were more sensitive to 5-FU as  
203 compared to FaDu cells (Figure 3;  $p < 0.001$  Mann-Whitney-U test). No difference between the  
204 cell lines were observed regarding the effect of 5-FU on proliferation ( $p = 0.550$ , Mann-Whitney-U  
205 test).

206 Incubation of PiCa cells with CSC significantly enhanced resistance to cisplatin. Cisplatin  
207 reduced viable cell counts to 46.9%. Incubation with CSC and cisplatin caused a decrease to  
208 only 52.9% ( $p < 0.001$ , WSRT). This is also reflected by the BrdU test-results. Cisplatin  
209 decreased the proportion of proliferating cells to 40.3%. Incubation with CSC and cisplatin  
210 resulted in 59.0% proliferation rate cells ( $p < 0.001$ , WSRT).

211 In FaDu cells CSC also significantly enhanced resistance to cisplatin as the proportion of  
212 proliferation cells decreased to 49.5% after incubation with cisplatin compared to a decrease to  
213 only 58.6% after incubation with cisplatin and CSC ( $p < 0.001$ , WSRT). However, no significant  
214 effect on the viable cell count of FaDu cells could be shown as cisplatin reduced the viable cell  
215 count to 55.3% and incubation with CSC and cisplatin caused a decrease to 50.7% ( $p = 0.040$ ,  
216 WSRT).

217 CSC had no influence on the cytotoxic effects of 5-FU in both cell lines ( $p = 0.012$  and  
218  $p = 0.478$ , WSRT). It had also no effect on 5-FU-induced reduction of proliferation ( PiCa cells:  
219  $p = 0.351$ , WSRT; FaDu cells:  $p = 0.014$ , WSRT).

220

## 221 **ABCG2 expression in oropharyngeal squamous cell carcinomas**

### 222 **Study population**

223 Table I summarizes the epidemiologic data of the cohort. Consistent with the etiology, we  
224 found fitting smoking histories of patients with p16-negative carcinomas HNSCC. At the time of  
225 diagnosis, 100% of p16-negative patients were formerly or currently exposed to cigarette  
226 smoke. In this group, 64.3% were current smokers. Former and current p16-negative smokers  
227 consumed 50.6 pack-years on average, whereas smokers of the p16-positive control group  
228 consumed only 8.9 pack-years ( $p<0.001$ , Mann-Whitney-U-test).

### 229 **ABCG2 immunohistochemistry**

230 ABCG2 expression was assessed by immunohistochemistry staining of kryosections of  
231 tumor samples (exemplified in Figure 5). Semi-quantitative evaluation of ABCG2 expression in  
232 30 tissue samples of oropharyngeal squamous cell carcinomas revealed an association  
233 between ABCG2 expression and smoking history of patients. Never smokers (HPV-positive  
234 control group) showed significantly lower expression levels than ever smokers ( $p=0.019$ , Chi-  
235 Square test; Figure 1). Moreover, ABCG2 expression correlated with tobacco consumption as  
236 measured in pack-years (Table I;  $p=0.025$ , Chi-Square test).

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### 238 **ABCG2 gene expression in TCGA HNSCC patients**

239 No differential expression of the *ABCG2* gene could be shown between current, former  
240 and never smokers or HPV-negative or -positive HNSCC.

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## 251           **Discussion**

252           Three-dimensional spheroid cultures of cells were used for the evaluation of  
253 chemoresistance. as spheroids were shown to maintain morphology, polarity and cell-to-cell  
254 interactions. Cancer cell spheroids contain heterogeneous cell populations with regard to  
255 metabolic activity, a major determinant of response to cytotoxic drugs (20, 21).

256           Continued smoking after being diagnosed with cancer has been recently put into focus  
257 because of its detrimental effects on several clinical parameters including cancer-specific  
258 mortality, more severe treatment-related toxicities and poorer response to therapy (10).  
259 According to available literature, continued smoking leads to more recurrences and second  
260 primary tumors in patients suffering from tobacco-driven cancers (24). Bjarnason and  
261 colleagues have shown that head and neck cancer patients who received radiotherapy and did  
262 not stop smoking during treatment, had significantly higher grades of mucositis when they were  
263 treated in the afternoon, after they smoked cigarettes, compared to patients who were treated in  
264 the morning (25). With respect to response to therapy, resistance to anticancer drugs plays a  
265 significant role in longterm survival. Cytotoxic pressure imposed by therapeutic agents on  
266 cancer cells is a persistent stimulus for the cells to adapt and to develop mechanisms of  
267 resistance. Part of this resistance is provided by members of the ABC transporter family. As  
268 efflux pumps, they are involved in the development of multidrug resistance, in which an  
269 individual tumor acquires cross-resistance to many different compounds that can be chemically  
270 and functionally unrelated (26).

271           Tobacco-associated head and neck carcinogenesis is commonly seen as the result of  
272 heavy smoking. It is likely that epithelial cells in the upper aerodigestive tract establish defense  
273 mechanisms against the multitude of tobacco-contained chemicals over time, potentially by  
274 increasing expression and/or function of efflux carriers like ABCG2, given that tobacco  
275 carcinogens are substrates of ABCG2 (16, 27). Cigarette smoke has been shown to rapidly  
276 induce ABCG2 expression in various cell lines, e.g. lung and esophageal cancer cells followed  
277 by a rapid decline after the stimulus was removed (18). In head and neck cancer cells, CSC has

278 been shown to cause expression of ABCG2 and membrane localization of the transporter within  
279 24 h after exposure (19).

280 Drug resistance could partly be the result of metabolic imprinting during carcinogenesis as  
281 cells maintain their preparedness to defend themselves against xenobiotics. Accordingly,  
282 increased ABCG2 expression has been shown to be a common feature in HNSCC cells  
283 resistant to cisplatin (28-30).

284 Both, PiCa und FaDu cells had low basal ABCG2 expression. Incubation of PiCa and  
285 FaDu cells with cigarette smoke resulted in an increased resistance to cisplatin. CSC led to a 3-  
286 fold increase of ABCG2 expression in PiCa and FaDu cells. FaDu cells showed significantly  
287 greater resistance to 5-FU. CSC did not modify 5-FU resistance in both cell types. This  
288 observation could be explained by the greater affinity of ABCG2 for cisplatin, particularly  
289 sulfated conjugates of cisplatin (31, 32). Cisplatin, but not 5-FU, is highly reactive towards  
290 glutathione, building sulfated complexes (28, 30).

291 These results are in agreement with our hypothesis that resistance to cytotoxic drugs is a  
292 consequence of a permanent xenobiotic impact of tobacco-contained carcinogens on the upper  
293 aerodigestive tract mucosa. It has also previously been shown that HPV-positive HNSCC and  
294 cervical cancer cell lines possess an intrinsic resistance to chemotherapy, independent of  
295 tobacco-related carcinogenesis (33, 34).

296 Considering the distinct etiologies of tobacco- and HPV-related HNSCC, tobacco-exposed  
297 PiCa and FaDu cells may have been primed to respond to cigarette smoke with increased  
298 ABCG2 expression and/or membrane localization to fend off carcinogens (27). The reported low  
299 inducibility of ABCG2 expression in HPV-positive HNSCC cells could be the result of HPV-  
300 specific carcinogenesis and may rather reflect stem-cell-like properties and increased  
301 proliferative capacity (35, 36).

302 Immunohistochemical analysis showed that ABCG2 is expressed more frequently in  
303 carcinomas of ever smokers compared to never smokers. Expression levels correlated with  
304 lifetime tobacco consumption as measured in pack years. These results indicated an  
305 association between tobacco consumption and ABCG2 expression on the protein level in PiCa

306 cells. Although p16-positive carcinomas occurred in patients who smoked significantly less,  
307 these tumors did not differ in ABCG2 expression from p16-negative cancers in all subgroups.  
308 We found no such differences in ABCG2 expression on the mRNA level.

309         There are several limitations of the present study that need to be discussed. Firstly, the  
310 study lacks a HPV-positive cell line. We were not able to produce spheroids of HPV-positive cell  
311 lines, in part due to a comparably slow growth. Nevertheless, we believe that three-dimensional  
312 cell cultures are an indispensable prerequisite for experiments with cytotoxic drugs. Secondly,  
313 regarding the tumors used for immunostaining, p16 was used as a surrogate marker for HPV-  
314 association. The presence of HPV was not confirmed by in-situ hybridization or PCR because  
315 our study focusses on the impact of tobacco consumption and p16 is a sufficient surrogate  
316 marker according to the 8<sup>th</sup> edition of the TNM staging system.

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## Summary

334        There is growing evidence that ABCG2 plays a critical role in drug resistance of head and  
335 neck cancer cells and that smoking influences its expression. Our results confirm these  
336 observations. Although PiCa and FaDu cells showed low basal ABCG2 expression, after  
337 stimulation with CSC both cell lines readily increased ABCG2 expression threefold and became  
338 resistant to cisplatin. ABCG2 immunostaining revealed significantly higher ABCG2 expression in  
339 HNSCC of ever compared to never smokers. Carcinomas of patients who smoked >10 pack  
340 years were significantly more often ABCG2-positive. No such correlations were found on the  
341 mRNA-level by TCGA analysis.

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343        Conflicts of Interest: none

344

345        Authors' Contributions: Data acquisition, data analysis, preparation and revision of  
346 manuscript

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348

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351

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478 **Figure/Table legends**

479

480 Table I: Clinical data in dependence of ABCG2 immunostaining

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482 Figure 1: Semiquantitative evaluation of ABCG2 staining intensity in oropharyngeal carcinomas (n=30) in  
483 relation to smoking.

484

485 Figure 2: Expression of ABCG2 in PiCa and FaDu cells before (control) and 24h/48h after incubation with  
486 CSC as a ratio to control.

487

488 Figure 3: Cytotoxic (WST8 assay) and anti-proliferative (BrdU assay) effect of cisplatin and 5-FU with and  
489 without co-incubation with cigarette smoke condensate (CSC) in spheroid cultures of PiCa and FaDu cells  
490 (bars indicate 95% confidence interval).

491

492 Figure 4: Spheroid cultures of a) PiCa and FaDu cells, untreated control and b) PiCa and  
493 FaDu cells, after incubation with cisplatin for 48 h.

494

495 Figure 5: Immunohistochemical staining of ABCG2 in an

496 a) ABCG2-negative carcinoma of base of tongue.

497 b) ABCG2-positive tonsillar carcinoma (+/+++).