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1 **An *in vitro* investigation on the cytotoxic and nuclear receptor transcriptional**
2 **activity of the mycotoxins fumonisin B1 and beauvericin.**

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9 **Abstract**

10 Fumonisin B1 (FB1) and beauvericin (BEA) are secondary metabolites of filamentous
11 fungi, which under appropriate temperature and humidity conditions may develop on
12 various foods and feeds. To date few studies have been performed to evaluate the
13 toxicological and endocrine disrupting effects of FB1 and BEA. The present study
14 makes use of various *in vitro* bioassays including; oestrogen, androgen, progestagen and
15 glucocorticoid reporter gene assays (RGAs) for the study of nuclear receptor
16 transcriptional activity, the thiazolyl blue tetrazolium bromide (MTT) assay to monitor
17 cytotoxicity and high content analysis (HCA) for the detection of pre-lethal toxicity in
18 the RGA and Caco-2 human colon adenocarcinoma cells.

19 At the receptor level, 0.001-10 μM BEA or FB1 did not induce any agonist responses in
20 the RGAs. However at non-cytotoxic concentrations, an antagonistic effect was
21 exhibited by FB1 on the androgen nuclear receptor transcriptional activity at 10 μM and
22 BEA on the progestagen and glucocorticoid receptors at 1 μM . MTT analysis showed
23 no decrease in cell viability at any concentration of FB1, whereas BEA showed a
24 significant decrease in viability at 10 μM . HCA analysis confirmed that the reduction in
25 the progestagen receptor transcriptional activity at 1 μM BEA was not due to pre-lethal
26 toxicity. In addition, BEA (10 μM) induced significant toxicity in both the TM-Luc
27 (progestagen responsive) and Caco-2 cells.

28 **Keywords: Mycotoxin, Beauvericin, Fumonisin B1, Reporter gene assay, High**
29 **Content Analysis.**

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30 **1. Introduction**

31 Mycotoxins are secondary metabolites of filamentous fungi, which under
32 appropriate temperature and humidity conditions may develop on various foods and
33 feeds. They are mainly produced by fungi belonging to the genera *Aspergillus*,
34 *Penicillium*, *Fusarium*, *Alternaria* and *Claviceps* (Fung et al., 2004). *Fusarium* species
35 are contaminants of wheat, maize, and other grains worldwide, capable of producing
36 high levels of fumonisin mycotoxins. Fumonisin B1 (FB1) is the most prevalent of the
37 fumonisins, accounting for approximately 70% of total fumonisins (Martins et al.,
38 2012). Studies have also highlighted that *Fusarium* species can co-produce other
39 mycotoxins such as Beauvericin (BEA) simultaneously (Dombrink-Kurtzman, 2003).

40 Total fumonisin concentrations in feed materials have been reported to vary from a
41 few $\mu\text{g}/\text{kg}$ to tens of mg/kg (EFSA, 2005). Dietary fumonisin estimates, by the Food
42 and Agriculture Organization of the United Nations and World Health Organization
43 (FAO/WHO, 2001), indicate exposure levels ranging from 0.02-0.2 $\mu\text{g}/\text{kg}$ in body
44 weight (b.w.)/day, thus remaining below the Tolerable Daily Intake (TDI) of 2 $\mu\text{g}/\text{kg}$
45 b.w./day as set in Europe by the Scientific Committee on Food (SCF, 2003).
46 Nevertheless, a wide range of animal diseases and pathophysiological effects such as
47 leukoencephalomalacia, porcine pulmonary oedema, liver and kidney toxicity and liver
48 cancer, as well as human oesophageal carcinoma are associated with FB1 ingestion
49 (Harrison et al., 1990; Kellerman et al., 1990; Gelderblom et al., 1997; Hussein et al.,
50 2001). While the molecular mechanism of FB1 toxicity is poorly understood, it appears
51 to be related to the deregulation of sphingolipid metabolism (Merrill et al., 2001).

52 BEA is predominantly found in cereal grains such as wheat, maize and rice
53 (Serrano et al., 2012) as well as other matrices such as nuts and dried fruits (Tolosa et
54 al., 2013). The mean dietary exposure to BEA varies from a minimum of 0.003 $\mu\text{g}/\text{kg}$
55 b.w./day to a maximum of 0.050 $\mu\text{g}/\text{kg}$ b.w./day (EFSA, 2014). However, the Panel on
56 Contaminants in the Food Chain (CONTAM) concluded that there was insufficient data
57 to establish a TDI or/and an acute reference dose (ARfD) for BEA in humans (EFSA,
58 2014). BEA possesses a wide range of biological activities. These substances are known
59 as ionophores, forming a complex with essential cations (Ca^{2+} , Na^{+} , K^{+}), which
60 increases ion permeability of biological membranes, therefore potentially affecting ionic
61 homeostasis (Chen et al., 2006). Many mycotoxins such as ochratoxin A, patulin,
62 alternariol and zearalenone have been found to possess endocrine disrupting capabilities
63 (Frizzell et al., 2011, 2013a, 2013b and 2014).

64 Endocrine disruptors (EDs) include both natural and man-made substances that
65 may interfere with the body's endocrine system by acting like endogenous hormones
66 and inducing adverse developmental, reproductive, neurological and immune effects
67 (IPCS, 2002). A few studies suggest that FB1 may act as a potential ED (Collins et al.,
68 1998; Gbore et al., 2009). While there is not enough data to confirm that FB1 is a
69 developmental or reproductive toxicant in animals or humans, Collins et al., (1998)
70 reported that FB1 was toxic to maternal rats and the foetus at 15 mg/kg of feed
71 consumption. In addition, Gbore (2009) reported that FB1 affected fertility in pigs by
72 causing a delay in sexual maturity and poor sperm production and quality. There are no
73 *in vivo* toxicological studies available on reproduction and developmental toxicity,
74 neurotoxicity or carcinogenicity for BEA. However, it has been shown to be absorbed
75 and rapidly metabolised to a range of uncharacterised metabolites as detected in the
76 eggs of laying hens and several tissues of turkeys and broilers (Jestoi, 2008).

77 *In vitro* bioassays may be used to investigate the toxicity and endocrine disrupting
78 potential of compounds (Connolly et al., 2011). The emerging technology, High
79 Content Analysis (HCA) is a highly powerful multi-parameter bio-analytical based tool
80 incorporating fluorescent microscopy with automated *in vitro* cell analysis software.
81 HCA provides assays with high sensitivity and specificity for pre-lethal cytotoxicity and
82 multiple biological endpoints for use as a high throughput-screening tool to monitor the
83 cytotoxicity, endocrine disruption and biological effects of compounds on exposed cells
84 (Clarke et al., 2015).

85 In this study, we have investigated the endocrine disrupting and cytotoxic
86 potential of FB1 and BEA using various *in vitro* bioassays. Reporter gene assays
87 (RGAs) utilising human mammary gland cells with natural steroid hormone receptors
88 for oestrogens, androgens, progestagens and glucocorticoids (Willemsen et al., 2004)
89 are employed for the identification of endocrine disruption at the level of nuclear
90 receptor transcriptional activity. HCA is used to detect early cytotoxicity, via multiple
91 markers in the progestagen responsive (TM-Luc) cell line exposed to 0.001-10 μ M
92 BEA, to ensure that a reduction in transcriptional activation of endocrine receptors is
93 not correlated with pre-lethal toxicity. HCA is also used to assess cytotoxicity in colon
94 adenocarcinoma (Caco-2) cells because the ingestion of food contaminated with FB1
95 and BEA is the main exposure route for animals and humans.

96
97

98 **2. Materials and methods**

99 **2.1 Reagents**

100 Methanol, thiazolyl blue tetrazolium bromide (MTT), FB1, BEA and the steroid
101 hormones 17 β -estradiol, testosterone, progesterone and hydrocortisone were obtained
102 from Sigma–Aldrich (Poole, Dorset, UK). Cell culture reagents were obtained from Life
103 Technologies (Paisley, UK). Multiparameter cytotoxicity 2 multiplex kit (8400202)
104 containing mitochondrial probe and cell membrane permeability dye was supplied by
105 Thermo Scientific, UK. Stock solutions of FB1 and BEA were prepared in methanol
106 and stored at -20°C. FB1 and BEA were dissolved in methanol at a final concentration
107 of 0.5% (v/v) in media for the RGAs, MTT assays and HCA.

108

109 **2.2 Cell culture**

110 All cells were routinely cultured in 75 cm² tissue culture flasks (Nunc, Roskilde,
111 Denmark) at 37 ° with 5% CO₂ and 95% humidity.

112 Four RGA cell lines were previously developed by the transformation of human
113 mammary gland cells with the luciferase gene under the control of a steroid hormone
114 inducible promoter (Willemsen et al., 2004). The MMV-Luc cell is specific for the
115 detection of oestrogens, TARM-Luc for androgens and progestagens, TM-Luc for
116 progestagens and TGRM-Luc for glucocorticoids and progestagens. The RGA cells
117 were routinely grown in cell culture medium containing Dulbecco's Modified Eagle
118 Medium (DMEM), 10% foetal bovine serum (FBS) and 1% penicillin streptomycin. As
119 phenol red is a weak oestrogen, DMEM without phenol red was used when culturing the
120 MMV-Luc cells. Cells were transferred prior to RGA analysis into assay media, which
121 was composed of DMEM and 10% hormone depleted serum.

122 The Caco-2 cell line (ATCC HTB-37) was routinely grown in DMEM medium,
123 10% FBS and 1% penicillin streptomycin.

124

125 **2.3 Reporter gene assay (RGA).**

126 RGAs were carried out as previously described by Frizzell et al. (2011). Briefly,
127 cells were seeded at a concentration of 4×10^5 cells/ml, 100 μ l/well, into white walled
128 96 well plates with clear flat bottoms (Greiner Bio-One, Germany). The cells were
129 incubated for 24 h and then exposed to BEA and FB1 (0.001, 0.01, 0.1, 1, 10 μ M) for
130 the agonist test. The positive control used with each cell line was as follows: 1.35 ng/ml
131 17 β -estradiol (MMV-Luc cells), 14.5 ng/ml testosterone (TARM-Luc cells), 157 ng/ml

132 progesterone (TM-Luc cells) and 181 ng/ml hydrocortisone (TGRM-Luc cells). A
133 solvent control 0.5% (v/v) methanol in media was also added to each plate. Antagonist
134 tests were carried out by incubating BEA and FB1 (0.001, 0.01, 0.1, 1, 10 μ M) with the
135 relevant positive control for each cell line. The cells were incubated for 48 h, after
136 which, the media was discarded and the cells washed once with phosphate buffered
137 saline (PBS). The cells were lysed with 30 μ l cell culture lysis buffer (Promega,
138 Southampton, UK) and then 100 μ l luciferase (Promega, Southampton, UK) injected
139 into each well and the response measured using the Mithras Multimode Reader
140 (Berthold, Other, Germany). The response of the cells to the various compounds was
141 measured and compared with the solvent control.

142

143 **2.4 Cell viability assay**

144 The MTT assay, based on the ability of viable cells to metabolize the yellow
145 tetrazolium salt to a blue formazan product by the mitochondria, was performed in
146 parallel to the RGA assays to monitor for cytotoxic effects of the mycotoxins and their
147 concentrations tested.

148 Briefly, the cells were exposed exactly as for the RGAs but in clear flat
149 bottomed 96 well plates (Nunc, Roskilde, Denmark). Following removal of the media,
150 50 μ L of MTT solution (2 mg/ml stock in PBS diluted 1:2.5 in assay media) was added
151 to each well and incubated for 4 h. The supernatant was removed and 200 μ L/well of
152 DMSO added to dissolve the formazan crystals. The absorbance was measured at
153 570nm and a reference absorbance of 630nm using an automatic plate reader (Tecan,
154 Safire, USA). Cell viability was calculated as a percentage absorbance of the sample
155 when compared to the absorbance of the solvent control (0.5% (v/v) methanol in
156 media).

157

158 **2.5 HCA multi-parameter assay**

159 HCA is a rapid and robust technology which can determine multiple cytotoxic
160 effects, including early (pre-lethal) as well as late-stage occurrences of cytotoxicity
161 simultaneously. The cytotoxicity of BEA and FB1 was assessed on Caco-2 cells as an
162 effective indicator of toxicity to the human gut. The TM-Luc cell line was also
163 investigated by HCA to confirm whether pre-lethal toxicity was inducing the antagonist
164 response observed at 1 μ M.

165 Briefly, cells were seeded at a concentration of 2×10^4 cells/ml, 100 μ l/well, into
166 96 well plates (Nunc, Roskilde, Denmark). The cells were incubated for 24 h and then
167 exposed to (0.001, 0.01, 0.1, 1, 10 μ M) of BEA (TM-Luc cells for 48 h) and BEA or
168 FB1 (Caco-2 cells for 24 and 48 h).

169 Cellomics® HCA reagent series multi-parameter cytotoxicity dyes were utilised.
170 Mitochondrial membrane potential dye was prepared by adding 117 μ l of anhydrous
171 DMSO to make a 1 mM stock. Permeability dye was used as provided in the
172 multiparameter cytotoxicity 2 multiplex kit (8400202). The live cell staining solution
173 was prepared by adding 2.1 μ l permeability dye to 6 ml of complete media that had
174 been preheated to 37°C, and then 21 μ l of mitochondrial membrane potential (final
175 concentration 3.5 mM). Nuclear stain solution was prepared by adding 5.5 μ l Hoechst
176 33342 dye to 11 ml 1X Wash Buffer.

177 After incubation, 50 μ l of live cell staining solution was added to each well.
178 Cells were incubated in the dark at 37°C and 5% CO₂ for 30 min. The staining solution
179 was aspirated and 100 μ l of 10% formalin solution (fixation solution) added. The cells
180 were incubated for 20 min at room temperature before discarding the fixation solution
181 and washing the cells with 100 μ l of PBS. Nuclear staining solution (100 μ l) was then
182 added, and the cells incubated for 10 min at room temperature protected from light. The
183 cells were then washed twice and the wells filled with 100 μ l of PBS. Cell number
184 (CN), nuclear area (NA), nuclear intensity (NI), plasma membrane permeability (PMP),
185 mitochondrial membrane potential (MMP) and mitochondrial mass (MM) were
186 measured using the CellInsight™ NXT High Content Screening platform (Thermo
187 Fisher Scientific, UK).

188

189 **2.6 Statistical analysis**

190 Assay exposures were carried out in triplicate wells and in three independent
191 experiments. Results were expressed as the mean \pm standard error of the mean (SEM) of
192 the triplicate exposures. For the RGAs, data was analysed using Microsoft Excel and
193 Graphpad PRISM software (San Diego, CA). A one way analysis of variance
194 (ANOVA) and Dunnett's multiple comparison test was used to determine significant
195 differences between the treatments and the corresponding controls in the RGAs, MTT
196 assays and HCA. The mean concentrations were tested for significant difference at the
197 95% confidence level. A *p* value of < 0.05 was considered statistically significant, $p = \leq$
198 0.05 (*), ≤ 0.01 (**) and ≤ 0.001 (***)).

199

200 **3. Results**

201 **3.1. Cell viability**

202 The MTT assay was used to determine the viability of the RGA cells following
203 exposure to FB1 or BEA (0.001-10 μM). No cytotoxicity was observed in any of the
204 RGA cell lines exposed to 0.001-10 μM FB1 (Fig.1) or 0.001-1 μM BEA. However, at
205 10 μM BEA, a decrease in cell viability for all RGA cell lines was observed ($p \leq 0.001$)
206 (Fig. 1).

207

208 **3.2. Reporter gene assays**

209 Neither FB1 nor BEA (0.001-10 μM) exhibited an agonist response in any of the
210 four RGA cell lines (data not shown). However FB1, at the highest concentration tested
211 (10 μM), exhibited an antagonistic effect ($p \leq 0.05$) on the androgen nuclear receptor
212 transcriptional activity (Fig. 2b). No antagonist effects were observed in the
213 progestagen, glucocorticoid or oestrogen RGAs (Fig. 2a, c and d). BEA, at the highest
214 concentration tested (10 μM), exhibited a strong antagonistic response ($p \leq 0.001$) in the
215 oestrogen, androgen, progestagen and glucocorticoid RGAs (Fig. 3a-d). However, the
216 MTT assay results indicate that this response is due to the cytotoxicity of BEA at 10 μM
217 on all of the RGA cell lines. Antagonistic effects on nuclear receptor transcriptional
218 activity in the progestagen ($p \leq 0.05$) and glucocorticoid ($p \leq 0.01$) RGAs were also
219 observed at non-toxic concentrations of 1 μM BEA (Fig. 3c and d). Considering that
220 BEA is cytotoxic to all of the RGA cell lines at 10 μM , it is possible that the antagonism
221 observed at 1 μM BEA is not a true response and instead may be due to pre-lethal
222 toxicity being initiated within the cells. The validity of this response was further
223 explored by HCA in the progestagen responsive, TM-Luc cell line.

224

225 **3.3 High Content Analysis (HCA).**

226 In the TM-Luc (progestagen responsive) cell line, BEA (10 μM) was not
227 possible to analyse due to lethal cytotoxic effects. BEA (1 μM) did not show any
228 significant differences when compared to the control. Therefore, no pre-lethal toxicity
229 was observed at 1 μM BEA, confirming that the antagonism observed in the progestagen
230 RGA was a true response (Fig. 4).

231 Exposure of Caco-2 cells to 0.001-10 μM FB1 or BEA revealed that 1 μM BEA
232 caused a significant ($p \leq 0.01$) decrease in the CN (Fig. 5). Nevertheless, 10 μM BEA
233 was not possible to analyse due to lethal cytotoxic effects on the Caco-2 cells.

234

235 **4. Discussion**

236 The MTT assay confirmed that FB1 (0.1 -10 μM) was not cytotoxic to any of the
237 four RGA cell lines. This value is consistent with other publications, Meca et al., (2010)
238 showed that exposure of Vero cells (monkey kidney) to 0-100 μM FB1 for 24 h
239 decreased cellular viability to 60 % at 100 μM when compared to the control. In
240 addition, Wan et al., (2013) did not observed a reduction of viability from 0 to 20 μM
241 FB1 in IPEC-J2 (porcine jejunal epithelial) cell line after 48 h of exposure.

242 BEA reduced cell viability at a concentration of 10 μM in all of the RGA and
243 Caco-2 cell lines. BEA (1 μM) also decreased viability in the Caco-2 cell line upon 48 h
244 exposure. This data is consistent with previous studies whereby 24 and 48 h 0-30 μM
245 BEA exposure of Caco-2 cells decreased viability to 80% and 87% respectively and
246 HT-29 (human colon adenocarcinoma) cells presented a decrease of 85% at 24 h and
247 90% at 48 h (Prosperini et al., 2012). Similar results were obtained by Calo et al.
248 (2004) with two human cell lines of myeloid origin (U-937 and HL-60 cells) and Ferrer
249 et al. (2009) who investigated 0-100 μM BEA exposure on Chinese hamster ovary cells
250 (CHO-K1). They observed a decline in viability at a concentration of 10 μM or higher
251 after 24 h.

252 The application of HCA in toxicity studies is based on the parallel analysis of
253 multiple markers for cytotoxicity, which allows early reversible and late irreversible
254 effects to be distinguished, and thus provides a more detailed analysis of compound-
255 induced toxicity (Ramirez et al. 2010; Tolosa et al., 2015). In this context, HCA can
256 identify gross toxicity and pre-lethal toxicity, whereby exposed cells are not dead but
257 are becoming unhealthy. While traditional end-point toxicity assays such as MTT can
258 identify gross toxicity, they cannot do so for pre-lethal toxicity.

259 In the current study, an antagonist response was observed in the progesterone
260 responsive TM-Luc cell line after exposure to 1 μM BEA. While the MTT assay was
261 able to confirm cytotoxicity via BEA exposure at 10 μM but not at 1 μM , the potential
262 for pre-lethal toxicity being responsible for the perceived antagonist response was
263 considered. Consequently, HCA analysis was utilised to confirm the absence of pre-

264 lethal toxicity and thus confirm the validity of the progesterone receptor antagonist
265 response.

266 The Caco-2 cell line is a well-recognised human gut cell model (Sambuy et al.,
267 2004) and as such is suited to investigating the toxic effects of food contaminants. HCA
268 analysis confirmed that FB1 was not cytotoxic at any of the concentrations tested on the
269 Caco-2 cell line. However, BEA exhibited cytotoxicity at 1 μ M on the Caco-2 cell line.
270 Furthermore, in this study was observed a slight decrease in MMP at 1 μ M BEA.
271 According to Jow et al. (2004), Ca²⁺-dependent pathway by BEA involves cell death,
272 in which it induced an increase in intracellular [Ca²⁺] that leads to a combination of
273 cellular apoptosis and necrosis responses. Moreover, Tonshin et al., (2010) in isolated
274 mitochondria BEA induced a loss of MMP where K⁺ inflow into the mitochondrial
275 matrix and uncoupling of oxidative phosphorylation, followed by induction of
276 apoptosis. In addition, Prosperini et al., (2013) investigated that Caco-2 cells exhibit
277 mitochondrial dysfunction leading a stable depolarized state of MMP and cell death
278 after exposure of 1.5 and 3 μ M BEA. Low BEA concentrations might be reached due to
279 food consumption and based on tissue accumulation (Jestoi et al., 2007). Moreover,
280 with regard to food intake, BEA might increase the absorption of commonly co-
281 occurring mycotoxins probably leading to higher toxicity. Thus, exposure to low BEA
282 concentrations activates diverse cellular stress response and protection systems
283 (Mallebrera et al., 2014). This indicates that continuous exposure to BEA might lead to
284 alter the intestinal epithelial barrier (Dornetshuber et al., 2009).

285 Antagonism of the androgen receptor in the TARM-Luc cell line was observed
286 following exposure to 10 μ M FB1. A reduction in the transcriptional activity of the
287 androgen, glucocorticoid, oestrogen and progesterone receptor was correlated to the
288 cytotoxic effects of BEA at 10 μ M rather than true antagonism. An antagonistic
289 response was also observed in the TGRM-Luc (glucocorticoid) and TM-Luc
290 (progesterone) cell lines following exposure to 1 μ M BEA. HCA established that no
291 pre-lethal toxicity was evident in the TM-Luc cell line at 1 μ M BEA and thus the
292 reduction in progesterone receptor transcriptional activity was confirmed as a true
293 antagonist response. To the authors' knowledge, this is the first study investigating the
294 endocrine disrupting effects of FB1 and BEA at the level of nuclear receptor activity.
295 The actions of progesterone, glucocorticoid and androgen are mediated by its receptor.
296 In the target cell, progesterone, glucocorticoid and androgen produce a change in

297 conformation of its receptors that is associated with transforming receptors from a non-
298 DNA binding form to one that will bind to DNA (Spitz et al., 2003). This
299 transformation is go with a loss of associated heat shock proteins and dimerization. The
300 activated receptors dimers then binds to specific DNA sequences within the promotor
301 region of progesterone, glucocorticoid and androgen responsive genes. Antagonist
302 impair the ability of receptors to interact with coactivators allowing the recruitment of
303 corepressors (Liu et al., 2002). The antagonist activity of an antihormone may depend
304 on the cell or tissue type. In addition, these transformations in the structure and function
305 of the receptor results in numerous endocrine disorders. Many antagonists of
306 progesterone receptor display antiproliferative effects in the endometrium by
307 suppressing follicular development and blocking the LH flood. Moreover, progesterone
308 antagonists are potent antiglucocorticoid agents (Neulen et al., 1996). GR signalling is
309 required for homeostatic control of pyramidal neurons. Thus, GR hormone influence
310 memory, mood, and neuronal survival (Savory et al., 2001) Therefore, inhibition of the
311 GR may affect the peripheral glucose metabolism, the stress response, and the
312 regulation of the hypothalamic pituitary axis (Honer et al., 2003; Deroche-Gamonet et
313 al., 2003). The regulatory steroidal sex hormones role in developmental processes such
314 as sex determination and differentiation is of particular interest with regard to endocrine
315 disruption (Kelce et al., 1995; 1997). Androgens, through interaction with the androgen
316 receptor, play decisive roles in sexual differentiation of the male reproductive tract,
317 accessory reproductive organs, and other tissues during fetal development. They also
318 influence male pubertal maturation and the maintenance of secondary sex characteristics
319 in adults. (Wilson et al., 2001)

320 This *in vitro* investigation has demonstrated the potential for FB1 and BEA to
321 modulate the endocrine system by antagonism of nuclear transcriptional activity as
322 observed for BEA (1 μ M) on the glucocorticoid and progesterone receptor and FB1 (10
323 μ M) on the androgen receptor. HCA has also proven to be an added value cytotoxic
324 assessment tool in establishing pre-lethal toxicity in exposed cells and confirming
325 antagonistic responses. In addition, while FB1 did not show any significant cytotoxic
326 effects on mammalian gut cells, BEA did at a concentration of 1 μ M. Further
327 investigation is needed to investigate the risk of BEA and FB1 exposure in humans and
328 animals.

329

330 **Conflict of interest**

331 The authors declare that there are no conflicts of interest.

332

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337

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482

483

484 **Legends of Figures:**

485

486 **Fig.1** Viability of the RGA cell lines a) MMV-Luc b) TARM-Luc c) TM-Luc and d)
487 TGRM-Luc following exposure to 0.001-10 μ M of FB1 and BEA for 48 h and
488 compared to the solvent control, as determined in the MTT assay. Values are means \pm
489 SEM for the three separate experiments (n=3), $p \leq 0.001$ (***).

490

491 **Fig.2** Results of RGA antagonistic test following co-exposure of the positive control
492 with FB1 (0.001-10 μ M) in the a) MMV-Luc (oestrogen responsive), b) TARM-Luc
493 (androgen responsive), c) TM-Luc (progestagen responsive) and d) TGRM-Luc
494 (glucocorticoid responsive) RGA cells. Responses measured are compared to the
495 solvent and the positive control (1.36 ng/ml 17 β -estradiol, 14.5 ng/ml testosterone, 157
496 ng/ml progesterone and 181 ng/ml cortisol, respectively). Results are expressed as the
497 mean percentage response \pm SEM for the three separate experiments (n=3), $p \leq 0.05$ (*).

498

499 **Fig.3** Results of RGA antagonistic test following co-exposure of the positive control
500 with BEA (0.001-10 μ M) in the a) MMV-Luc (estrogen responsive), b) TARM-Luc
501 (androgen responsive), c) TM-Luc (progestagen responsive) and d) TGRM-Luc
502 (glucocorticoid responsive) RGA cells. Responses measured are compared to the
503 solvent and relevant positive controls (1.36 ng/ml 17 β -estradiol, 14.5 ng/ml
504 testosterone, 157 ng/ml progesterone and 181 ng/ml cortisol, respectively). Responses
505 are expressed as the mean percentage response \pm SEM for the three separate
506 experiments (n=3), $p \leq 0.05$ (*), ≤ 0.01 (**), ≤ 0.001 (***).

507

508 **Fig.4** Quantification of the cytotoxic effects of 0.001-1 μ M BEA in the progestagen
509 responsive TM-Luc cells as measured by HCA. a) cell number (CN) b) nuclear area
510 (NA), c) nuclear intensity (NI), d) plasma membrane permeability (PMP), e)
511 mitochondrial membrane potential (MMP) and f) mitochondrial mass (MM). Data are
512 expressed as mean values \pm SEM for the three separate experiments (n=3). $p \leq 0.05$ (*)
513 and $p \leq 0.01$ (**) indicate significant differences from the solvent control.

514

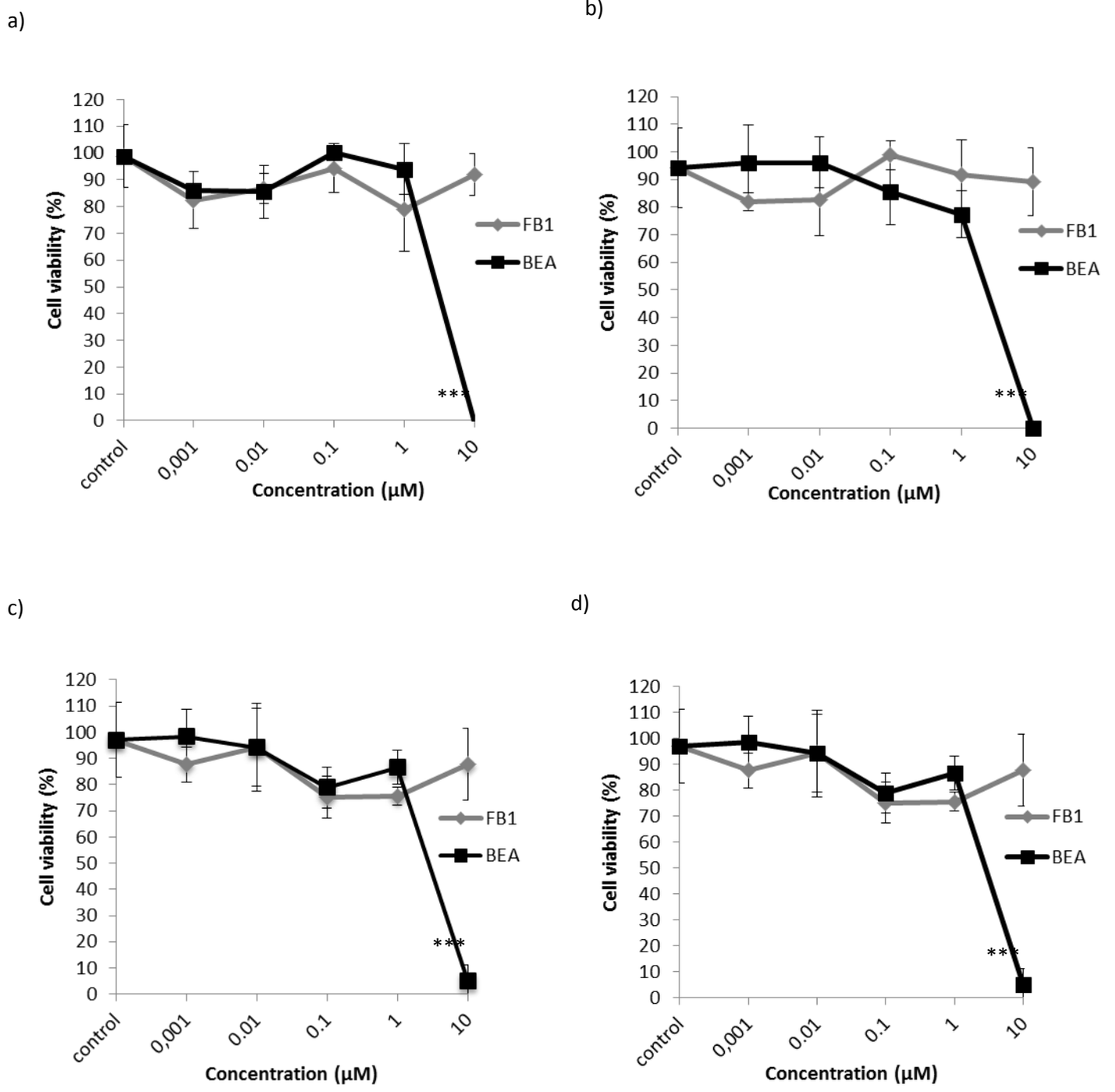
515 **Fig.5** Quantification of the cytotoxic effects of 0.001-10 μ M FB1 and BEA in the gut

516 derived Caco-2 cells after 48 h exposure as measured by HCA. a) cell number (CN) b)
517 nuclear area (NA), c) nuclear intensity (NI), d) plasma membrane permeability (PMP),
518 e) mitochondrial membrane potential (MMP) and f) mitochondrial mass (MM). Data are
519 expressed as mean values \pm SEM for the three separate experiments (n=3). $p \leq 0.001$
520 (***)indicate significant differences from the solvent control.

521

522

Fig.1



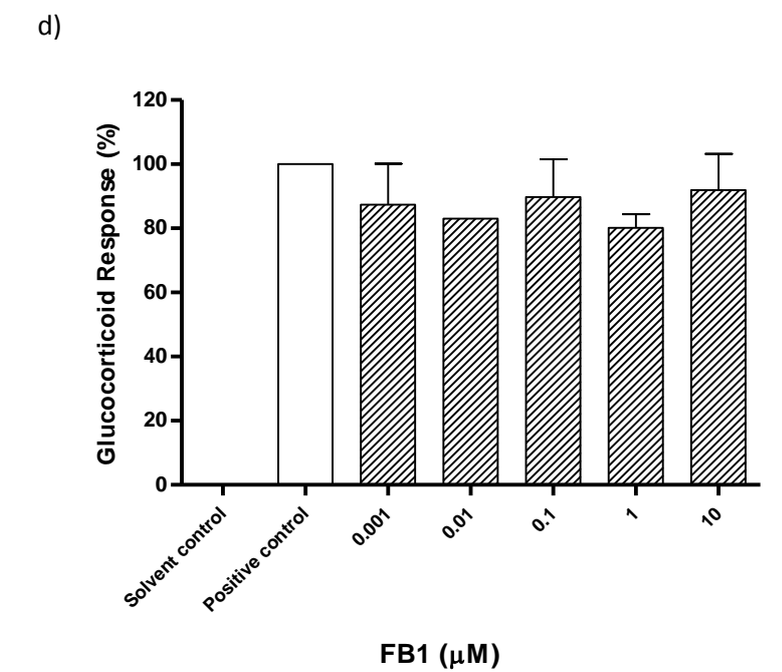
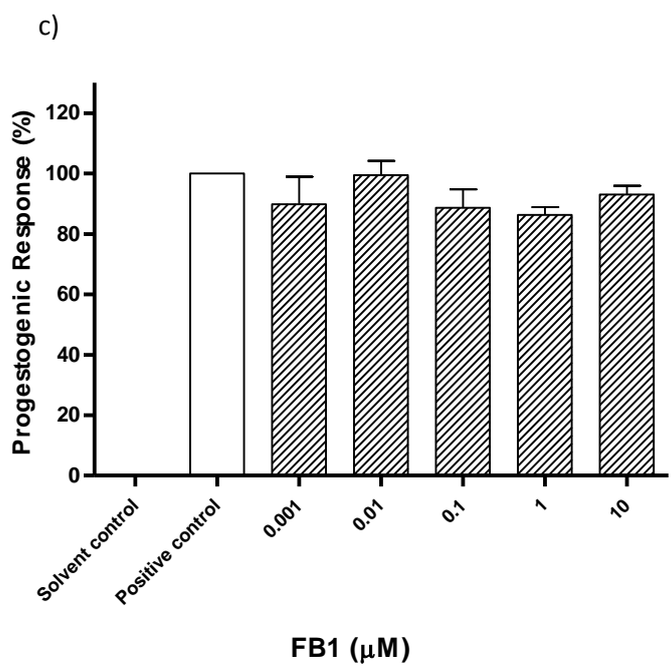
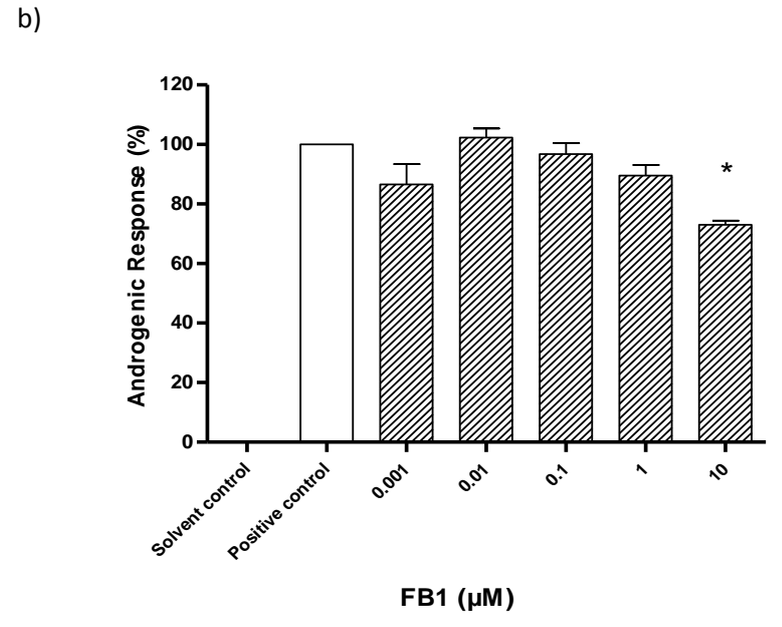
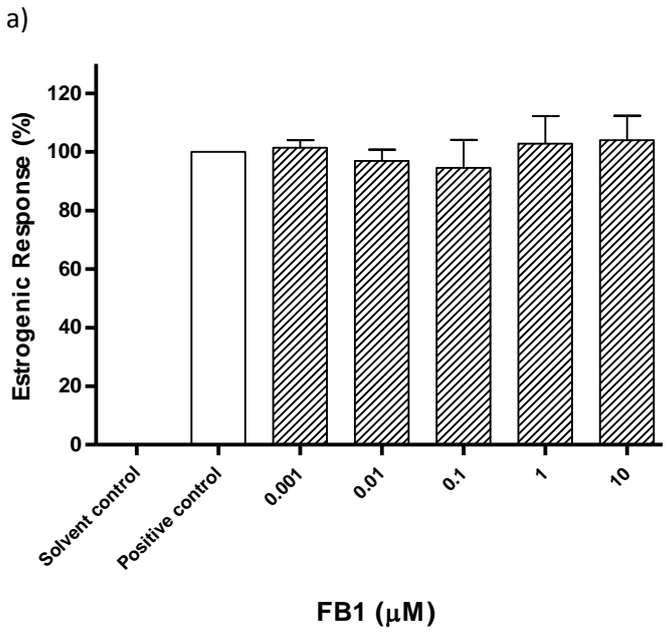


Figure
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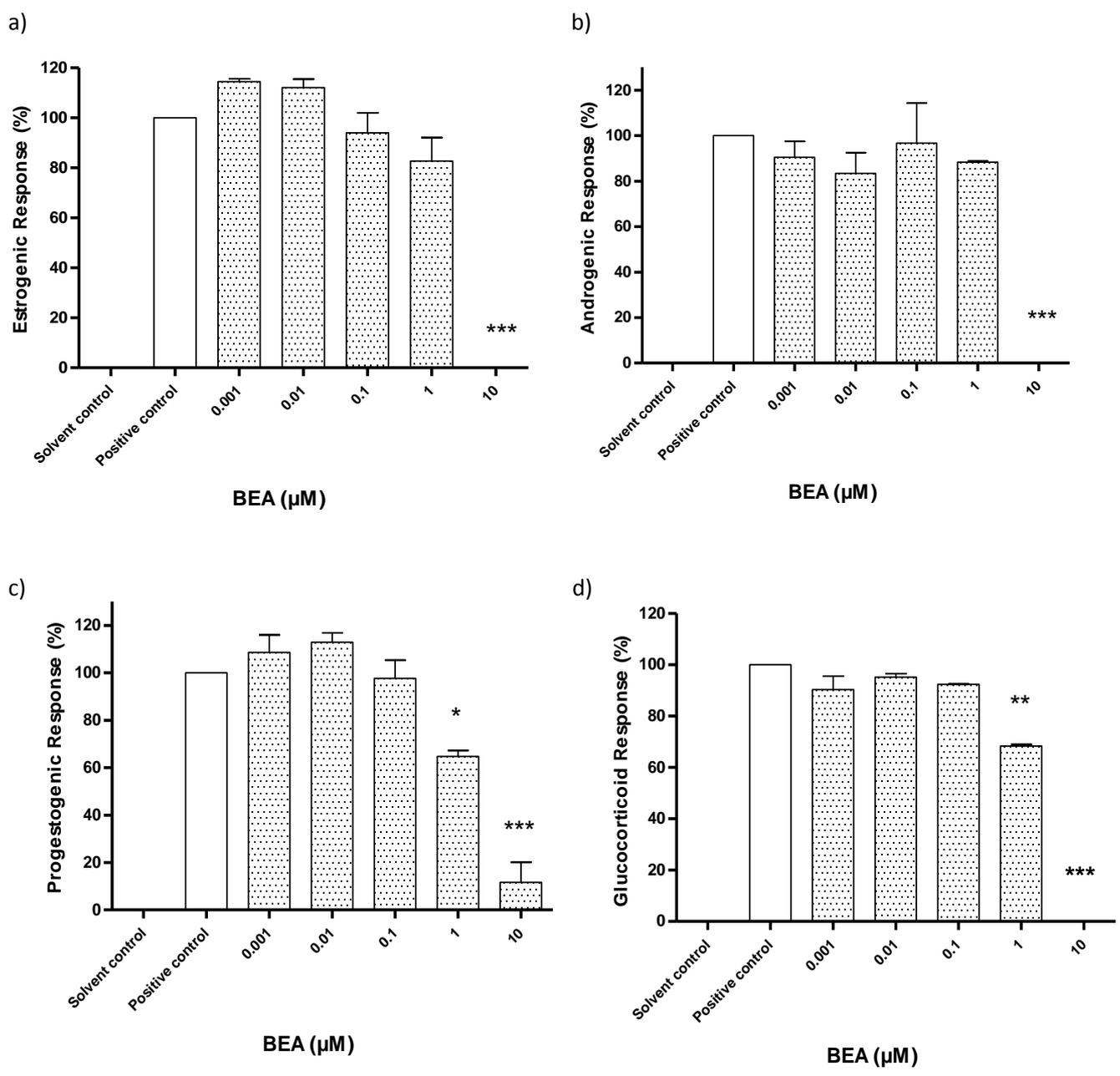


Fig. 4

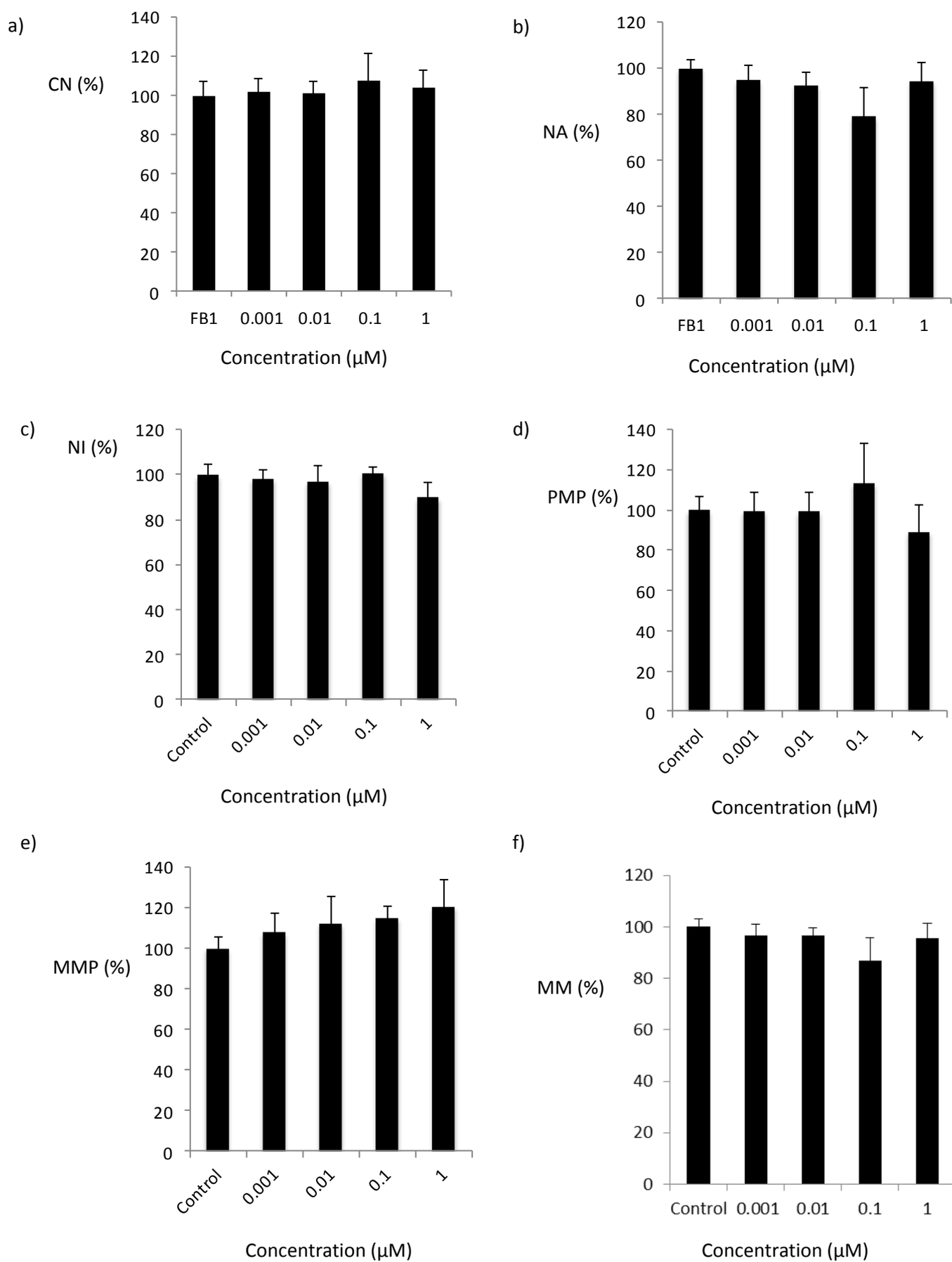


Fig. 5.

