

1 **Caspase-independence and characterization of bisnaphthalimidopropyl spermidine**  
2 **induced cytotoxicity in HL60 cells.**

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16

17 **Abbreviations:** BNIPSpd, Bisnaphthalimidopropylspermidine; PI, Propidium iodide; OPA,  
18 o-phthaldialdehyde.

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21 **Running title:** Bisnaphthalimide-induced cytotoxicity

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23

24 **Abstract -**

25 Bisnaphthalimides are DNA intercalators of potential use as chemotherapeutics but for which  
26 the range of mechanism of action is only gradually being elucidated. Using human  
27 promyelocytic HL-60 cells, we extend characterisation of the cytotoxicity of  
28 bisnaphthalimidopropylspermidine (BNIPSpd) and examine the relationship with caspase-  
29 activity. Within 4h exposure, BNIPSpd (1-10  $\mu$ M) induced significant DNA strand breakage.  
30 Evidence of apoptosis was progressive through the experimental period. Within 6h, BNIPSpd  
31 increased the proportion of cells exhibiting plasma membrane phosphatidylserine exposure.  
32 Within 12h, active caspase expression increased and was sustained with 5 and 10  $\mu$ M  
33 BNIPSpd. Flow cytometric analysis revealed caspase activity in cells with and without  
34 damaged membranes. By 24 h, 5 and 10  $\mu$ M BNIPSpd increased hypodiploid DNA content  
35 and internucleosomal DNA fragmentation (DNA ladders) typical of the later stages of  
36 apoptosis. 1 $\mu$ M BNIPSpd exposure also increased hypodiploid DNA content by 48h.  
37 Polyamine levels decreased by 24 h BNIPSpd exposure. The pan-caspase inhibitor, z-VAD,  
38 significantly decreased DNA degradation (hypodiploid DNA and DNA ladders) and induced  
39 an increase in cell growth. Despite this, cell growth and viability remained significantly  
40 impaired. We propose that BNIPSpd cytotoxicity arises through DNA damage and not  
41 polyamine depletion and that cytotoxicity is dominated by but not dependent upon caspase  
42 driven apoptosis.

43

44 *Key Words:* Apoptosis; Bisnaphthalimides; Caspase-inhibition; Cytotoxicity; Genotoxicity;  
45 HL-60 cells

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## 48        1.        **Introduction**

49        Naphthalimides and bisnaphthalimides are DNA intercalating agents (e.g. Brana et al., 1993,  
50        2001; Cacho et al., 2003; Kong Thoo Lin et al., 2003; Lv et al., 2009; Tan et al., 2015; Wang  
51        et al., 2016) with compounds comprising these moieties variously being proposed for  
52        exploitation as anti-tumourigenics, anti-microbial or anti-parasitic agents (e.g. Gellerman,  
53        2016; Kong Thoo Lin et al., 2003; Oliveira et al., 2007; Noro et al., 2015; Kopsida et al., 2016  
54        Graca et al., 2016).    Bisnaphthalimides consist of two aromatic naphthalimido rings attached  
55        by a linker chain containing nitrogen atoms. Principally, the intercalations of the naphthalimide  
56        planar aromatic rings between DNA base-pairs distorts the conformation of the DNA backbone  
57        leading to interference with DNA-protein interactions (Hsiang et al., 1989; Brana et al., 1993,  
58        Dance et al., 2005). In addition, particular naphthalimide and bisnaphthalimides have been  
59        variously shown to inhibit topoisomerases directly, exert post- DNA damage effects on DNA  
60        damage signalling pathways, impair DNA repair, or induce lysosomal permeability (Filosa et  
61        al., 2009; Zhu et al., 2009; Chen et al., 2010; Bestwick et al., 2011; Barron et al., 2015; Tan et  
62        al., 2015; Zhang et al., 2016a;). The ultimate consequence of exposure to (bis-) naphthalimides  
63        *in vitro* has included cell cycle arrest and apoptosis (Ralton et al., 2009; Liang et al., 2011; Yang  
64        et al., 2011a; Seliga et al., 2013; Zhang et al., 2016a).

65               This array of often complementary mechanisms of (bis-)naphthalimide action  
66        influencing tumour cell, microbes and parasite development has continued to encourage their  
67        development as therapeutics. In previous work, we linked bisnaphthalimido propyl fragments  
68        with natural polyamines such as spermine and spermidine (Dance et al., 2005; Kong Thoo Lin  
69        et al., 2000; Pavlov et al., 2001). As neoplastic transformation can be accompanied by elevated  
70        polyamine levels resulting from altered biosynthesis, catabolism and uptake (Basuroy and  
71        Gerner 2006), we hypothesised that the polyamine linker would facilitate uptake of the  
72        bisnaphthalimides into neoplastic cells (Dance et al., 2005).

73 We previously reported that bisnaphthalimidopropylspermine (BNIPSpm) and  
74 bisnaphthalimidopropylspermidine (BNIPSpd) have enhanced aqueous solubility, and are  
75 rapidly and homogeneously distributed within the nuclei of MCF-7 breast carcinoma and  
76 Caco-2 colon adenocarcinoma cells where they cause significant DNA damage and  
77 impairment of DNA base excision repair (Bestwick et al., 2011; Dance et al., 2005).  
78 Moreover, the more cytotoxic of the two, BNIPSpd (Fig 1), induces apoptosis in Caco-2 and  
79 HT-29 colon epithelial cells (Ralton et al., 2009). However, the conservation of the apoptotic  
80 response to BNIPSpd, and the mechanism underlying such response in differing cell types,  
81 has not been established. Here, following a preliminary report (Kong Thoo Lin et al., 2003),  
82 we provide a detailed assessment of the effects of BNIPSpd on cell growth and cytotoxicity in  
83 HL-60 promyelocytic leukaemia cells, examining the temporal relationship of apoptosis to  
84 DNA damage and polyamine levels and the extent of caspase dependency within overall  
85 BNIPSpd toxicity.

86

## 87 **2. Materials and methods**

### 88 *2.1. Materials*

89 HL-60 cells were purchased from the European Collection of Cell Cultures  
90 (98070106; Public Health England, Salisbury, UK). RPMI medium and foetal calf serum  
91 were from Lonza Sales AG (Verviers, Belgium). Tissue culture flasks were supplied by  
92 Greiner Bio-One Ltd (Gloucestershire, UK). Active Caspase-3 detection kit, Cell Cycle Plus  
93 DNA Reagent Kit and QC DNA particles were from BD (Oxford, UK). Vybrant FAM  
94 Polycaspases kits, Amplex Red hydrogen peroxide assay kits and DAPI were from Life  
95 Technologies Ltd (Paisley, UK). The Annexin V-FITC kit and single strand DNA detection  
96 kit were from eBioscience Ltd (Hatfield, UK). Etoposide, camptothecin, dimethylsulfoxide

97 (DMSO) and all other reagents were purchased from Sigma-Aldrich Company Ltd (Dorset,  
98 UK) unless stated otherwise. BNIPSpd was synthesised and characterised according to our  
99 previous methods (Kong Thoo Lin and Pavlov 2000).

100

## 101 2.2. *Cell Culture and bisnaphthalimidopropyl polyamine exposure*

102 HL 60 cells were cultured in RPMI 1640 medium supplemented with 10 % (v/v) foetal  
103 calf serum, 1% (v/v) non essential amino acids, 2 mM glutamine, 50 µg/mL streptomycin and  
104 50 µg/mL penicillin. Cells were kept in a humidified (95% relative humidity, RH) incubator  
105 at 37 °C, 5% CO<sub>2</sub>. BNIPSpd was solubilized in 20% (v/v) DMSO and cells ( $5 \times 10^5$  cells mL<sup>-1</sup>)  
106 were incubated with 0.1-10µM BNIPSpd (0.02% [v/v] DMSO final concentrations), 0.02 %  
107 (v/v) DMSO or sterile double distilled water for 1-72 h. The chemotherapeutic agents  
108 etoposide (10 µM) and camptothecin (4 µM) in 0.1% [v/v] DMSO final concentrations, were  
109 used as positive controls to confirm assay function as appropriate. For caspase inhibitor  
110 experiments, cells were pre-incubated with 100 µM z-VAD-fmk (in 0.2% v/v DMSO final  
111 concentration) for 1 h at 37°C, 5% CO<sub>2</sub>, 95% RH prior to addition of BNIPSpd.

## 112 2.3. *Cell culture growth and cytotoxicity*

113 Cells were collected by centrifugation (300 g for 5 min, RT), the culture media  
114 removed and the pellet suspended in trypan blue (0.2% w/v final concentration in PBS). Cells  
115 were counted using a Neubauer haemocytometer (Baur et al., 1975).

116

## 117 2.4. *DNA single strand breaks*

118

119 DNA single strand breaks were determined by single cell gel electrophoresis (SCGE)  
120 as described previously (Bestwick et al., 2005). Nucleoids were stained with 4',6-diamidino-

121 2-phenylindole (DAPI, 5  $\mu\text{g mL}^{-1}$  stock) and scored visually. One hundred images per gel,  
122 (with duplicate gels per slide) were classified according to the intensity of fluorescence in the  
123 nucleoid tail and assigned a value of 0-4 with 0 representing no damage and 4 maximal  
124 damage. Thus, the total damage score can range from 0 to 400. This method of classification  
125 has been extensively validated using computerised image analysis (Duthie et al., 1996).

#### 126 2.5. *Phosphatidylserine exposure and membrane integrity*

127 Exposure of phosphatidylserine at the extracellular surface of the plasma membrane  
128 was determined as previously described (Bestwick and Milne 2006) by FITC-Annexin-V  
129 binding using a commercial assay kit, and the relationship to membrane damage assessed by  
130 co-incubation with propidium iodide (PI) as per the manufacturer's protocol (eBioscience Ltd,  
131 Hatfield, UK).

#### 132 2.6. *Caspase-3 and pan-caspase*

133 The presence of the active fragment of caspase-3 was analysed by flow cytometry  
134 after cell fixation, permeabilization and labelling with PE-conjugated polyclonal rabbit anti-  
135 active caspase-3 (BD, Oxford) as described previously for HL-60 cells (Bestwick and Milne  
136 2006). Total (pan-) caspase activity relative to maintenance of membrane-integrity was  
137 estimated by measuring fluorescence in cells co-treated with FAM-VAD-FMK polycaspase  
138 reagent (labelling active caspase 1, 3, 4, 5, 6,7, 8, 9) and propidium iodide according to  
139 manufacturer's recommendations (Life Technologies Ltd, Paisley, UK). Camptothecin  
140 treatment and mock-(water) treated cells (positive and negative controls respectively), were  
141 used to define active caspase-3 expression or increased pan-caspase activity using Cell Quest  
142 software (BD, Oxford, UK). 10,000 events were recorded.

143

144 2.7. *Apoptosis-associated DNA fragmentation*

145

146 SubG1 (hypodiploid) DNA content was evaluated as described previously (Bestwick  
147 et al., 2007) using a commercial kit (Cycle test plus) according to the manufacturer's protocol  
148 (BD, Oxford, UK). Flow cytometry of the propidium iodide-stained nuclei was performed  
149 with a flow rate of 12  $\mu$ L/min and cell cycle distribution selected from linear FL-2 area v.  
150 width plots with doublet discrimination in FL-2. Singlet events, excluding debris, were gated  
151 and 20,000 events were acquired within the gate. The percentage of cells with DNA content  
152  $<2N$  (sub-G1) was calculated from histograms of linear FL-2 area plots of the singlet gated  
153 region using Cell Quest Software (BD, Oxford, UK) and Mod Fit LT software (Verity  
154 Software House, ME, USA). A DNA QC Particle Kit (BD, Oxford, UK) was used to verify  
155 instrument linearity, doublet discrimination and cytometer alignment.

156 Internucleosomal DNA fragmentation giving rise to the characteristic apoptotic 'DNA  
157 ladder', was identified following DNA extraction and separation by agarose gel  
158 electrophoresis as detailed in Bestwick and Milne, (Bestwick and Milne 2006).

159 2.8 *Polyamine extraction and analysis*

160 Cells were treated either with solvent vehicle (DMSO), BNIPSpd or  $\alpha$ -  
161 difluoromethylornithine (DFMO; 5 mM) as a positive control for polyamine changes. Culture  
162 medium from control and BNIPSpd-treated cells were decanted, the cells collected by  
163 centrifugation (1000 g, 5 min, 20 °C) and counted using a Neubauer haemocytometer. Cells  
164 were washed with PBS (x2) before being homogenised with perchloric acid (final  
165 concentration 1% v/v) and incubated with internal standard (1,7-diaminoheptane ;20  $\mu$ M for  
166 60 min on ice). The homogenate was centrifuged at 10000g (4 °C) for 10 minutes and the  
167 supernatant transferred to a fresh container and stored at -20°C. The residual cell pellet and  
168 culture medium were also retained at -20°C. HPLC analysis of polyamine content was carried  
169 out by derivatization with o-phthalaldehyde (OPA) in the presence of 2-mercaptoethanol as

170 described by Seiler and Knodgen (Seiler and Knodgen, 1985). HPLC was conducted with a  
171 Phenomenex Luna 5 mm, 25 cm x 4.6 mm, C18 HPLC column. Detection of polyamines after  
172 OPA derivatisation was at 345 nm excitation with 455 nm emission filters. Data were  
173 collected and integrated using Waters Empower Software (Waters Yvelines Cedex, France).

#### 174 *2.9. Hydrogen peroxide determination.*

175 Hydrogen peroxide, arising from the interaction of the polyamine analogue with amine  
176 oxidases in bovine serum (Parchment et al., 1990), is a potential confounder of the cellular  
177 response to BNIPSpd. Using the Amplex Red Assay (Life Technologies Ltd, Paisley, UK),  
178 the level of hydrogen peroxide in complete RPMI 1640 medium containing BNIPSpd (1-50  
179  $\mu\text{M}$ ) was measured within 5h incubation via the peroxidase catalysed formation of resorufin  
180 (560 nm, Unicam UV/Vis spectrophotometer) . Co-incubations with native or heat denatured  
181 (5 min, 100 °C) bovine catalase (Sigma-Aldrich Company Ltd, Dorset, UK; 2.5  $\mu\text{g}/\text{mL}$ ) were  
182 used to confirm detection of hydrogen peroxide.

183

#### 184 *2.10 Statistical Analysis*

185 Experiments were conducted a minimum of three times unless stated otherwise. Data  
186 are presented as mean  $\pm$  SD. A *P*-value of  $< 0.05$  (by student's *t*-test) was taken as the  
187 minimum basis for assigning significance. Statistical analysis was conducted using SigmaStat  
188 (Systat, Software Inc. London, UK).

189

### 190 **3. Results**

191

#### 192 *3.1. Cell growth and cytotoxicity*

193



194 BNIPSpd (1-10  $\mu$ M) caused a significant inhibition (1  $\mu$ M) or cessation (5 and 10  $\mu$ M)  
195 of cell growth (Fig 2A). Based on trypan blue staining, 5 and 10  $\mu$ M BNIPSpd also exerted  
196 significant cytotoxicity (Fig 2B) within 24 h. Therefore, analysis of the mechanisms  
197 underpinning cytotoxicity was predominantly conducted within the first 24 h of treatment.

198

### 199 *3.2. DNA single strand breaks*

200 DNA single strand breakage was significantly increased after 4h exposure to 1  $\mu$ M-10  $\mu$ M  
201 BNIPSpd and remained significantly increased over control cells after 24h treatment (Fig. 3).

### 202 *3.3. Phosphatidylserine exposure and membrane integrity.*

203

204 Exposure to BNIPSpd ( $\geq 1$   $\mu$ M) caused a significant increase in the proportion of cells  
205 binding annexin-V-FITC at 6-24 h but remaining recalcitrant to PI staining (Fig 4A and 4B).  
206 This indicates exposure of phosphatidylserine on the external plasma membrane surface in the  
207 absence of membrane damage. In addition, BNIPSpd also induced a significant increase in the  
208 proportion of cells co-labelling with annexin-V-FITC and PI (Fig.4A and B), representing  
209 annexin-V-FITC binding to cells with damaged membranes. Cells treated with solvent  
210 vehicle alone remained predominantly unlabeled with either annexin-V-FITC or PI (Fig 4A  
211 and B). For all treatments, a residual proportion of cells stained with PI but did not label with  
212 annexin-V-FITC. This represented 2.6 % ( $\pm 0.19$ ) of control cells at 24 h and this increased ( $P$   
213  $< 0.05$ ) after treatment with 10  $\mu$ M BNIPSpd (24 h) to 14.3 % ( $\pm 3.4$ ).

214

### 215 *3.4. Active caspase-3 expression and pan caspase activity.*

216 Active caspase-3 expression increased during time in culture within negative (DMSO  
217 solvent only treated) control cells but 10 $\mu$ M and 5  $\mu$ M BNIPSpd treatment further

218 significantly enhanced and sustained active caspase-3 expression by 4 h and 12h after  
219 exposure respectively (fig 5). A transient but significant increase in caspase-3 was also  
220 observed by 12 h of incubation in 1µM BNIPSpd treated cells (Fig. 5).

221 Focussing on 5 and 10 µM BNIPSpd treatment for which there was a more rapid and  
222 sustained active caspase-3 expression (Fig 5), the relationship between general-caspase  
223 activity and membrane damage was flow-cytometrically assessed by combining FAM-VAD-  
224 FMK FLICA labelling of active caspases (caspase 1, 3, 4, 5, 6,7, 8, 9) with PI staining to  
225 measure membrane integrity. The median fluorescence intensity, as a marker of overall  
226 caspase activity (Fig 6A) and the proportion of cells expressing pancaspase activity (Fig 6B)  
227 increased significantly by 8 h of treatment with BNIPSpd. Increased pan-caspase activity was  
228 detected both in cells with intact membranes and in those with compromised membrane  
229 integrity (Fig 6B).

230

### 231 3.5 'Apoptotic' DNA fragmentation

232  
233 A significant increase in the proportion of cells with a subG1 DNA content was  
234 observed after 3 h and 24 h for 10 and 5 µM BNIPSpd treatments respectively. A smaller but  
235 nevertheless significant increase in sub G1 DNA content was observed by 48 h with 1 µM  
236 BNIPSpd (Fig. 7A). Classical apoptotic internucleosomal DNA fragmentation was not  
237 observed within the first 12 h of BNIPSpd exposure. By 24 h, however, DNA ladders  
238 occurred in cultures treated with  $\geq 5$  µM BNIPSpd (Fig. 7B) and were particularly apparent in  
239 5 µM BNIPSpd treated cells

240

### 241 3.6. Polyamine levels

242 No changes to spermine and spermidine levels were observed after 12 h incubation with 1-  
243 10  $\mu$ M BNIPSpd. BNIPSpd treatment for 24 h resulted in a lowering of spermine and  
244 spermidine levels (Fig 8A and B). Putrescine was not detected in the analysis of either  
245 BNIPSpd treated or untreated cells.

246

### 247 *3.7. Effect of caspase inhibitor on cytotoxicity*

248 To investigate caspase-dependency of BNIPSpd toxicity, cells were pre-treated with the pan  
249 caspase inhibitor z-VAD-FMK, prior to and during exposure to 10  $\mu$ M BNIPSpd. Z-VAD-  
250 fmk strongly decreased the intensity of internucleosomal DNA fragmentation and  
251 significantly lowered the increase in sub G1 DNA content (Fig 9 A and B). The anti-  
252 proliferative (Fig 9C) and cytotoxic effects (Fig 9D) of BNIPSpd were slightly but  
253 significantly lowered by caspase inhibition. Nevertheless, there remained a highly significant  
254 reduction in cell growth (Fig 9C) and high levels of cytotoxicity (Fig 9D) in response  
255 BNIPSpd.

256

### 257 *3.8. Culture medium hydrogen peroxide content*

258 The influence of BNIPSpd on the concentration of hydrogen peroxide in serum-supplemented  
259 culture media, which may be a confounding factor in BNIPSpd toxicity, was established via  
260 the Amplex Red assay. Endogenous hydrogen peroxide present in the test mixture acts as a  
261 co-substrate in the oxidation of amplex red via exogenously added horseradish peroxidase and  
262 leads to the formation of resorufin. No significant change in hydrogen peroxide content was  
263 observed up to 10  $\mu$ M BNIPSpd (the maximum concentration used in the cytotoxicity assays)  
264 but a significant increase in resorufin absorbance was observed with 50  $\mu$ M BNIPSpd. This  
265 increase was sensitive to the presence of native catalase, confirming the potential for elevated

266 levels of exogenous hydrogen peroxide at these higher, but here, not experimentally relevant  
267 BNIPSpd incubations (Fig 10).

268

#### 269 **4. Discussion**

270 Within a series of novel bisnaphthalimido-polyamine compounds, the spermidine-containing  
271 BNIPSpd exerts the dominant cytotoxicity against MCF-7 breast carcinoma (Dance et al.,  
272 2005) and Caco-2 colon adenocarcinoma cells (Bestwick et al., 2011). Cytotoxicity is  
273 preceded by DNA single strand breakage (Dance et al., 2005; Bestwick et al., 2011). Here,  
274 BNIPSpd also rapidly caused DNA single strand breaks in HL-60 cells and a progressive  
275 increase in features typical of a classical apoptotic progression towards cell death. Within  
276 24h, BNIPSpd exposure was associated with early externalisation of plasma membrane  
277 phosphatidylserine, a progressive increase in active caspase-3 and pan caspase activity and  
278 DNA fragmentation (hypodiploid DNA content and internucleosomal DNA fragmentation),  
279 all features indicative of apoptosis (Elmore 2007; Darzynkiewicz and Li 1996; Kohler et al.,  
280 2002; Otsuki et al., 2003).

281 Both the naphthalimide and polyamine moiety of BNIPSpd may trigger apoptosis  
282 (Basuroy and Gerner 2006; Casero and Woster, 2001; Brana et al., 2001; Liang et al., 2011).  
283 For the (bis)naphthalimide moiety, DNA intercalation and the effect on DNA replication and  
284 integrity is proposed as the primary mode of action (Kong Thoo Lin and Pavlov, 2000;  
285 Dance et al., 2005) but the presence of the polyamine moiety might also suggest an influence  
286 on cellular polyamine levels and metabolism (Morgan, 1999; Wallace and Niiranen, 2007;  
287 Schipper et al., 2000; Basuroy et al., 2006). The naturally occurring polyamines putrescine,  
288 spermidine and spermine represent a potential link between proliferation and cell death  
289 (Schipper et al., 2000; Moschou et al., 2014) and polyamine analogues may influence cell

290 survival by effects on cellular polyamine pools (Zou et al., 2004). However, polyamines  
291 exhibit a complex relationship to cell survival and exert both negative and positive regulatory  
292 effects on apoptosis (e.g. Milovic and Turchanowa, 2003; Stefanelli et al., 2001; Yuan et al.,  
293 2002; Zou et al., 2004) .

294 Here, DNA single strand breakage occurred within the first 4 h of treatment with  
295 BNIPSpd. This parallels or occurs in advance of the earliest detection of apoptosis markers.  
296 Conversely, there was no change in polyamine level within the first 12 h of exposure. Thus,  
297 while polyamine levels do decline during cell death, on a temporal basis polyamines do not  
298 appear to be involved in induction of DNA damage or initiation of apoptosis within  
299 BNIPSpd-treated HL60 cells. This contrasts with observations in Caco-2 and HT-29 cells  
300 where polyamine levels are significantly affected both at the pro-apoptotic BNIPSpd dosage  
301 range ( $>0.5 \mu\text{M}$ ) and by non- toxic BNIPSpd exposures ( $\geq 0.01 \mu\text{M}$ ) (Ralton et al., 2009).  
302 Based on our observations of the timing of BNIPSpd- induced DNA damage relative to  
303 changes in polyamine levels, we propose that it is the effect of the bisnaphthalimide moiety  
304 on DNA (Dance et al., 2005; Bestwick et al., 2011) that is the consistent early-induced  
305 phenomenon associated with BNIPSpd-cell interactions.

306 However, when considering the extent of cytotoxicity and DNA strand breakage at 1  
307  $\mu\text{M}$  BNIPSpd exposure, the degree of DNA damage does not predict the eventual magnitude  
308 of apoptosis or overall toxicity, with limited toxicity despite considerable DNA damage and  
309 inhibition of proliferation. Inhibitory effects on DNA repair activity may predominate at these  
310 BNIPSpd concentrations (Bestwick et al., 2011). The influence of BNIPSpd and other  
311 analogues on cell growth and survival regulation at low-cytotoxic and sub-cytotoxic dosages  
312 are currently under investigation.

313 In addition to the direct effect of BNIPSpd on cell survival, we considered whether  
314 amine oxidases in the culture medium may promote hydrogen peroxide generation in the

315 presence of the spermidine analogue (Bitonti et al., 1990; Parchment et al., 1990). No increase  
316 in hydrogen peroxide was detected in culture medium incubated with BNIPSpd at the  
317 concentrations employed. Consequently, hydrogen peroxide generation from BNIPSpd  
318 addition in the media appears not to be a factor in BNIPSpd toxicity in this culture system.  
319 However, whether BNIPSpd toxicity involves oxidative stress intracellularly has not been  
320 determined.

321         Given the timing of DNA strand break detection, we suggest that the intrinsic  
322 mitochondrial pathway for apoptosis, initiated by DNA damage as a likely primary route for  
323 orchestrating toxicity. Given the p53 null status of HL60 (Leroy et al., 2014), this is not  
324 dependent on p53 though be mediated through a functional substitution such as via p73 (e.g.  
325 Chakraborty et al., 2010), a member of the p53 protein family (Chekova et al., 2018).  
326 Although BNIPSpd toxicity was associated with features typical of a classical apoptosis, the  
327 presence of a proportion of cells exhibiting membrane damage concomitant with or in  
328 advance of apoptotic markers also suggests a more heterogeneous cell death process occurs as  
329 part of the overall BNIPSpd-induced toxicity. Cytotoxicity of BNIPSpd in MCF-7 cells  
330 (Dance et al., 2005) which do not express Caspase-3 (Janicke, 2009) indicates a non-caspase-  
331 3 dependency for toxicity. Here, z-VAD-FMK was used to investigate general caspase-  
332 relationship to cytotoxicity. While abolishing apoptotic DNA fragmentation, z-VAD-FMK  
333 failed to rescue the majority of cells from the anti-proliferative and cytotoxic effects of  
334 BNIPSpd. Many cytotoxic chemotherapeutic drugs demonstrate non-caspase dependency  
335 (e.g. camptothecin, doxorubicin, paclitaxel; Broker et al., 2005). Caspase inhibition can  
336 induce a switch from classical apoptosis to alternate cell death processes or an alteration of  
337 dominance of a cell death mechanism paralleling apoptosis induction. The range of changes to  
338 cell death mode has included autophagy, “mitotic catastrophe”, necrosis-like programmed cell  
339 death (e.g. necroptosis; Zhang et al., 2016b) and more traditional necrosis (Broker et al.,

340 2005). For example, caspase inhibition switches doxorubicin-induced cytotoxicity within  
341 SKN-SH neuroblastoma cells from apoptosis to senescence (Rebbaa et al., 2003), whereas Z-  
342 VAD-FMK promotes a shift from apoptosis to necrosis during corneal scrape-induced  
343 keratocyte death (Kim et al., 2000). For many of these outcomes, the mechanistic inter-  
344 relationship has yet to be comprehensively established. Caspase inhibition might also  
345 increase the role of non-caspase dependent apoptotic-like processes, for example involving  
346 apoptosis inducing factor (AIF). AIF has been implicated recently in mono-naphthalimide  
347 spermidine-induced apoptosis in HeLa cells (Yang et al., 2011b). We hypothesise that the  
348 many influences of (bis)naphthalimide-containing compounds on processes linked to cell  
349 proliferation and genomic integrity (Filosa et al., 2009; Zhu et al., 2009; Chen et al., 2010;  
350 Bestwick et al., 2011; Yang et al., 2011a; Li et al., 2013; Seliga et al., 2013; Shen et al., 2013)  
351 renders them capable of inducing a range of cell death mechanisms as well as multiple point  
352 of provocation of apoptosis.

353 Here HL-60 cells are utilised but comparison to multiple cell types (and ultimately the  
354 use of organoid systems) will help to confirm the universality and mechanistic variations of  
355 BNIPSpd-cell-interactions in vitro. Nevertheless, the effects observed in Caco-2 and HT-29  
356 colon adenocarcinoma cell cultures (Ralton et al. 2009) point to key conserved outcomes in  
357 terms of apoptosis features and early induction of DNA damage post exposure (Bestwick et al.,  
358 2011) but, as discussed above, significant differences are apparent for the effect on cellular  
359 polyamine content (Ralton et al., 2009). Significantly, previous work on characterising  
360 BNIPSpd and related bisnaphthalimide-polyamine compounds (Dance et al., 2005) on the form  
361 of toxicity within caspase-3 deficient but p53 active MCF-7 cells (e.g. Zhao et al., 2017) would  
362 be a useful component of comparisons on BNIPSpd activity within differing apoptotic  
363 regulator-competent backgrounds.

364 In conclusion, BNIPSpd cytotoxicity in HL60 cells occurs primarily through an  
365 apoptotic-like process, potentially initiated by bisnaphthalimide-induced DNA damage and  
366 within a temporal framework involving phosphatidylserine exposure, caspase activation and  
367 DNA fragmentation. However, caspase inhibition does not rescue cells from BNIPSpd  
368 toxicity. While it remains to be established whether this adds to exploitability for  
369 chemotherapy or represents a confounder to the therapeutic development of this  
370 bisnaphthalimidopropyl polyamine series, caspase-independent cell death provides further  
371 evidence of the heterogeneity in the cytotoxic response to (bis)naphthalimides.

372

373

374



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379

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381

382 **References**

- 383 Barron, GA, Goua, M, Kuraoka, I., Bermano, G., Iwai, S, Lin PKT., 2015.
- 384 Bisnaphthalimidopropyl diaminodicyclohexylmethane (BNIPDaCHM) bisintercalates to  
385 DNA and induces DNA damage and repair instability in triple negative breast cancer cells via  
386 p21. *Chemico-Biological Interactions* 242, 307-315
- 387 Basuroy, U.K., Gerner, E.W., 2006. Emerging concepts in targeting the polyamine metabolic  
388 pathway in epithelial cancer chemoprevention and chemotherapy. *Journal of Biochemistry*  
389 139, 27-33.
- 390 Baur, H., Kasperek, S., Pfaff, E., 1975. Criteria of viability of isolated liver cells. *Hoppe-*  
391 *Seylers Zeitschrift fur Physiologische Chemie* 356, 827-838.
- 392 Bestwick, C.S., Milne, L., 2006. Influence of galangin on HL-60 cell proliferation and  
393 survival. *Cancer Letters* 243, 80-89.
- 394 Bestwick, C.S., Milne, L., Pirie, L., Duthie, S.J., 2005. The effect of short term kaempferol  
395 exposure on reactive oxygen levels and integrity of human (HL60) leukaemic cells.  
396 *Biochimica Biophysica Acta –Molecular Basis of Disease* 1740, 340-349.
- 397 Bestwick, C.S., Milne, L., Duthie, S.J., 2007. Kaempferol induced inhibition of HL-60 cell  
398 growth results from a heterogeneous response, dominated by cell cycle alterations. *Chemico-*  
399 *Biological Interactions* 3, 179-191.
- 400 Bestwick, C.S., Ralton, L.D., Milne, L., Kong Thoo Lin. P., Duthie, S.J., 2011. The influence  
401 of bisnaphthalimidopropyl-polyamines on DNA instability and repair in Caco-2 colon  
402 epithelial cells. *Cell Biology and Toxicology* 27, 455-463.

403 Brana, M.F., Castellano, J.M., Moran, M., Perez De Vega, M.J., Romerdahl, C.R., Qian,  
404 X.D., Bousquet, P.F., Emling, F., Schlick, E., Keilhauer, G., 1993. Bis-naphthalimides: a new  
405 class of antitumor agents. *Anti-Cancer Drug Design* 8, 257-268.

406 Brana, M.F., Cacho, M., Gradillas, B., De Pascual-Teresa, B., Ramos, A., 2001. Intercalators  
407 as anticancer drugs. *Current Pharmaceutical Design* 17, 1745–1780.

408 Broker, L.E., Kruyt, F.A.E., Giaccone, G., 2005. Cell death independent of caspases: A  
409 review. *Clin. Cancer Res.* 11, 3155-3162.

410 Bitonti, A.,J., Dumont, J.A., Buch, T.L., Stemerick, D.M., Edwards, M.L., McCann, P.P.  
411 1990. Bis(benzyl)polyamine analogs as novel substrates for polyamine oxidase. *J. Biol.*  
412 *Chem.* 265, 382-388.

413 Cacho, M., Ramos, A., Dominguez, M.T., Pozuelo, J.M., Abradelo, C., Rey-Stolle, M.F.,  
414 Yuste, M., Carrasco, C., Bailly, C., 2003. Synthesis, biological evaluation and DNA binding  
415 properties of novel mono and bisnaphthalimides. *Organic and Biomolecular Chemistry*1, 648-  
416 654.

417 Casero, R.A., Woster, P.M.. 2001. Terminally alkylated polyamine analogues as  
418 chemotherapeutic agents. *J. Med. Chem.* 44,1-26.

419 Cechova J, Coufal J, Jagelska E.B., Fojta M., Vaclav Brazda V., 2018. p73, like its p53  
420 homolog, shows preference for inverted repeats forming cruciform. *PLoS ONE* 13(4):  
421 e0195835.

422 Chakraborty J., Banerjee S., Ray P., Hossain D.M.S., Bhattacharyya S., Adhikary A.,  
423 2010. Gain of Cellular Adaptation Due to Prolonged p53 Impairment Leads to Functional

424 Switchover from p53 to p73 during DNA Damage in Acute Myeloid Leukemia Cells. *J.Biol*  
425 *Chem*, 285, 33104-33112

426 Chen, Z., Liang, X., Zhang, H.Y., Xie, H., Liu, J., Xu, Y., Zhu, W., Wang, Y., Wang, X., Tan,  
427 S., Kuang, D., Qian, X., 2010. A new class of naphthalimide-based antitumor agents that  
428 inhibit topoisomerase II and Induce lysosomal membrane permeabilization and apoptosis. *J.*  
429 *Med.Chem.* 53, 2589-2600.

430 Dance, A-M., Ralton, L., Fuller, Z., Bestwick, C.S., Milne, L., Duthie, S., Kong Thoo Lin, P.,  
431 2005. Synthesis and biological activities of bisnaphthalimido polyamines derivatives:  
432 cytotoxicity, DNA binding, DNA damage and drug localization in breast cancer MCF 7 cells.  
433 *Biochemical Pharmacology* 69, 19-27.

434 Darzynkiewicz, Z., Li, X., 1996. Measurement of cell death by flow cytometry, in: Cotter,  
435 T.G., Martin, S.J. (Eds.), *Techniques in Apoptosis*. Portland Press, London, pp. 71-106.

436 Duthie, S.J., Ma, A., Ross, M.A., Collins, A.R., 1996. Antioxidant supplementation decreases  
437 oxidative damage in human lymphocytes. *Cancer Research* 56, 1291-1295.

438 Elmore, S., 2007. Apoptosis: A review of programmed cell death. *Toxicologic Pathology*  
439 35,495-516.

440 Filosa, R., Peduto, A., Di Micco, S., De Caprariis, P., Festa, M., Petrella, A., Capranico, G.,  
441 Bifulco, G., 2009. Molecular modelling studies, synthesis and biological activity of a series of  
442 novel bisnaphthalimides and their development as new DNA topoisomerase II inhibitors.  
443 *Bioorganic and Medicinal Chemistry* 17, 13 – 24.

444 Gellerman, G (2016) Recent Developments in the Synthesis and Applications of Anticancer  
445 Amonafide Derivatives. A Mini Review. *Letters in Drug Design and Discovery*, 13: 47-63.

446 Graça N., Gaspar L., Costa D., Loureiro I., Kong Thoo Lin P., Ramos I., A., Pemberton I.,  
447 MacDougall J., Tavares J., Cordeiro-da-Silva A., 2016. Bisnaphthalimidopropyl Derivatives  
448 as New Leads *Against Trypanosoma brucei*. *Antimicrob Agents Chemother.* 60(4):2532-6.doi:  
449 10.1128/AAC.02490-15

450 Hsiang, Y.H., Jiang, J.B., Liu, L.F., 1989. Topoisomerase II-mediated DNA cleavage by  
451 amonafide and its structural analogues. *Molecular Pharmacology* 36, 371-376.

452 Janicke, R.U., 2009. MCF-7 breast carcinoma cells do not express caspase-3. *Breast Cancer*  
453 *Res. Treat.* 117, 219-221.

454 Kim, W-J., Mohan, R.R., Mohan, R.R., Wilson, S.E. 2000. Caspase inhibitor z-VAD-FMK  
455 inhibits keratocyte apoptosis, but promotes keratocyte necrosis, after corneal epithelial scrape.  
456 *Exp. Eye. Res.* 71, 225-232.

457 Kohler, C., Orrenius, S., Zhivotovsky, B., 2002. Evaluation of caspase activity in apoptotic  
458 cells. *Journal of Immunological Methods* 265, 97-110.

459 Kong Thoo Lin, P., Pavlov, V.A., 2000. The synthesis and in vitro cytotoxic studies of novel  
460 bis-naphthalimidopropyl polyamine derivatives. *Bioorganic and Medicinal Chemistry Letters*  
461 10, 1609-1612.

462 Kong Thoo Lin. P., Dance, A.M., Bestwick, C., Milne, L., 2003. The biological activities of  
463 new polyamine derivatives as potential therapeutic agents *Biochem. Soc. Trans.* 31, 407-410.

464 Kopsida M., Barron G.A., Bermano G., Kong Thoo Lin P., Goua M., 2016. Novel  
465 bisnaphthalimidopropyl (BNIPs) derivatives as anticancer compounds targeting DNA in  
466 human breast cancer cells. *Org. Biomol. Chem.* 14, 9780-9789.

467 Leroy B., Hollestelle L.G.A., Minna J.D., Gazdar A.F., Soussi T., 2014. Analysis of TP53  
468 Mutation Status in Human Cancer Cell Lines: A Reassessment. *Hum Mutat.* 35, 756-765

469 Li, M. Li, Q., Zhang, Y.H., Tian, Z.Y., Ma, H.X., Zhao, J., Xie, S.Q., Wang, C.J., 2013.  
470 Antitumor effects and preliminary systemic toxicity of ANISpm in vivo and in vitro. *Anti-  
471 Cancer Drugs* 24, 32-42.

472 Liang, X., Wu, A.B., Xu, Y.F., Xu, K., Liu, J.W., Qian, X.H., 2011. B1, a novel  
473 naphthalimide -based DNA intercalator, induces cell cycle arrest and apoptosis in HeLa cells  
474 via p53 activation. *Investigational New drugs* 29, 646-658

475 Lv, M., Xu, H., 2009. Overview of naphthalimide analogs as anticancer agents. *Current  
476 Medicinal Chemistry* 16, 4797-4813.

477 Milovic, V., Turchanowa, L., 2003. Polyamines and colon cancer. *Biochem. Soc. Trans.* 31,  
478 381-383.

479 Morgan, D.M.L., 1999. Polyamines - an overview. *Molecular Biotechnology* 11, 229-250.  
480 Moschou P.N., Roubelakis-Angelakis, K.A., 2014. Polyamines and programmed cell death. *J.  
481 Exp. Botany* 5, 1285-1296

482 Noro J., Maciel J., Duarte D., Dias Olival A.C., Baptista C., Cordeiro da Silva A., Alves M.J.,  
483 Kong Thoo Lin P., 2015. Evaluation of New Naphthalimides as Potential Anticancer Agents  
484 Against Breast Cancer MCF-7, Pancreatic Cancer BxPC-3 and Colon Cancer HCT-15 Cell  
485 Lines. *Organic Chem Curr Res*, 4:2 <http://dx.doi.org/10.4172/2161-0401.1000144>

486 Oliveira J, Ralton L, Tavares J, Codeiro-da-Silva A, Bestwick CS, McPherson A and Kong  
487 Thoo Lin P (2007) The Synthesis and the in vitro Cytotoxicity studies of

488 Bisnaphthalimidopropyl polyamine derivatives against Colon Cancer Cells and Parasite  
489 *Leishmania infantum*" *Bioorganic & Medicinal Chemistry*, 15: 541-545

490 Otsuki, Y., Li, Z.L., Shibata, M.A., 2003. Apoptotic detection methods - from morphology to  
491 gene. *Progress in Histochemistry and Cytochemistry* 38, 275-339.

492 Parchment, R.E., Lewwllyn, A., Swartzendruber, D., Pierce, G.B., 1990. Serum amine  
493 oxidase activity contributes to crisis in mouse embryo cell lines. *Proc. Natl. Acad. Sci. USA*  
494 87, 4340-4344.

495 Pavlov, V.A., Kong Thoo Lin, P., Rodilla, V., 2001. Cytotoxicity, DNA binding and  
496 localisation of novel bis-naphthalimidopropyl polyamine derivatives. *Chemico-Biological*  
497 *Interactions* 137, 15-24.

498 Ralton, L.D., Bestwick, C.S., Milne, L., Duthie, S., Kong Thoo Lin, P., 2009  
499 Bisnaphthalimidopropyl spermidine induces apoptosis within colon carcinoma cells.  
500 *Chemico-Biological Interactions* 177, 1-6.

501 Rebbaa, A., Zheng, X., Chou, P.M., Mirkin, B.L., 2003. Caspase inhibition switches  
502 doxorubicin-induced apoptosis to senescence. *Oncogene* 22, 2805-2811.

503 Seiler, N., Knodgen, B., 1985. Determination of polyamines and related compounds by  
504 reversed-phase high-performance liquid chromatography: Improved separation systems.  
505 *Journal of Chromatography B-Biomedical Sciences and Applications* 339, 45-57.

506 Seliga, R., Pilatova, M., Sarissky, M., Viglasky, V., Walko, M., Mojzis, J., 2013. Novel  
507 naphthalimide polyamine derivatives as potential antitumor agents. *Molecular Biology*  
508 *Reports* 40, 4129-4137.

509 Schipper, R.G., Penning, L.C., Verhofstad, A.A.J., 2000. Involvement of polyamines in  
510 apoptosis. Facts and controversies: effectors or protectors? *Seminars in Cancer Biology* 10,  
511 55-68.

512 Shen, K., Sun, L.Y., Zhang, H.Y., Xu, Y.F., Qian, X.H., Lu, Y.H., Li, Q., Ni, L., Liu, J.W.,  
513 2013. ROS-mediated lysosomal-mitochondrial pathway is induced by a novel Amonafide  
514 analogue, 7c, in human He la cervix carcinoma cells. *Cancer Letters* 333, 229-238.

515 Stefanelli, C., Pignatti, C., Tantini, B., Fattori, M., Stanic, I., Mackintosh, C.A., Flamigni, F.,  
516 Guarnieri, C., Caldarera, C.M., Pegg, A.E., 2001. Effect of polyamine depletion on caspase  
517 activation: a study with spermine synthase-deficient cells. *Biochem J.* 355, 199-206.  
518

519 Tan S.Y., Sun, D.H., Lyu, J.K., Sun, X., Wu, F.S., Li, Q., Yang, Y.Q., Liu, J.X., Wang, X.,  
520 Chen, Z., Li, H.L., Qian, X.H., Xu, X.F. 2015. Antiproliferative and apoptosis-inducing  
521 activities of novel-cyclam conjugates through dual topoisomerase (topo) I/II inhibition.  
522 *Biorganic and Medicinal Chemistry* 23, 5672-5680  
523

524 Wallace, H.M., Niiranen, K., 2007. Polyamine analogues as anticancer drugs. *Amino acids*  
525 33,261-265.  
526

527 Wang, K.R., Sun, Q., Ma, CL., Rong, RX., Cao, ZR., Wang, XM., Li, XL. 2016. Substituent  
528 effects on cytotoxic activity, spectroscopic property, and DNA binding property of  
529 naphthalimide derivatives. *Chemical Biology & Drug Design* 87, 664-672  
530



531 Yang, L., Li, W., Tian, Z., Zhao, J., Wang, C., 2011a. Monoaphthalimide spermidine  
532 conjugate induces proliferation inhibition and apoptosis in Hela cells. *Toxicology in Vitro* 25,  
533 882-889.  
534

535 Yang, L.H., Zhao, J., Zhu, Y.Q., Tian, Z.Y., Wang C.J., 2011b. Reactive oxygen species  
536 (ROS) accumulation induced by mononaphthalimide-spermidine leads to intrinsic and AIF-  
537 mediated apoptosis in HeLa cells. *Oncology Reports* 25, 1099-1107.  
538

539 Yuan, Q., Ray, R.M., Johnson, L.R., 2002. Polyamine depletion prevents camptothecin-  
540 induced apoptosis by inhibiting the release of cytochrome c. *American Journal of Physiology-  
541 Cell Physiology* 282, 1290-1297.  
542

543 Zhang, G. H., An, Y.F., Lu, X., Zhong, H., Zhu, Y.H., Wu, Y.M., Ma, F., Yang, J.M., Liu,  
544 Y.C., Zhou, Z.P., Peng, Y., Chen, Z.F., 2016a. A Novel Naphthalimide Compound Restores  
545 p53 Function in Non-small Cell Lung Cancer by Reorganizing the Bak center dot Bcl-xl  
546 Complex and Triggering Transcriptional Regulation. *Journal of Biological Chemistry* 291,  
547 4211-4225

548 Zhang, J., Yang, Y., He, W.Y., Sun, L.M., 2016b. Necrosome core machinery: MLKL.  
549 *Cellular and Molecular Life Sciences* 73: 2153-2163

550 Zhao M, Howard EW, Guo Z, Parris AB, Yang X (2017) p53 pathway determines the cellular  
551 response to alcohol-induced DNA damage in MCF-7 breast cancer cells. *PLoS ONE* 12(4):  
552 e0175121.

553 Zhu, H., Miao, Z.H., Huang, M., Feng, J-M., Zhang, Z-X., Lu, J-J., Cai, Y-J., Tong, L-J., Xu,  
554 Y-F., Qian, X-H., Ding, J., 2009. Naphthalimides Induce G(2) Arrest Through the ATM-  
555 Activated Chk2-Executed Pathway in HCT116 Cells. *Neoplasia* 11, 1226-1234.

556

557 Zou, T.T., Rao, J.N., Guo, X., Rao, J.N., Strauch, E.D., Bass, B.L., Wang, J.Y., 2004. NF-  
558 kappa B-mediated IAP expression induces resistance of intestinal epithelial cells to apoptosis  
559 after polyamine depletion. *American Journal of Physiology-Cell Physiology* 286, 1009-18.

560

561 **Figure Legends**

562

563 Fig 1. Structure of Bisnaphthalimidopropylspermidine (BNIPSpd)

564

565 Fig 2. Cell growth (A) and cell death (B) were determined following exposure to DMSO (□),  
566 1 μM (○), 5 μM (⊕) and 10 μM (●) BNIPSpd. Cell counts were determined by microscopy,  
567 with trypan blue staining used to indicate loss of membrane integrity. Data are means ± SD (n  
568 ≥6). \*P<0.05, \*\*P < 0.01, \*\*\*P < 0.001 v. respective time point for DMSO control.

569

570 Fig. 3. DNA damage. Alkaline SCGE followed by visual ‘comet’ scoring was used to determine  
571 DNA single strand breaks after exposure to BNIPSpd. Data are means ± SD (n=3), \*\*\*P<0.001  
572 v. respective time point for DMSO control.

573

574 Fig 4. Effect of BNIPSpd exposure on plasma membrane phosphatidylserine labelling relative  
575 to membrane damage. Cells were treated with BNIPSpd for 6 (A) and 24h (B). Cells labelled  
576 with Annexin-V-FITC but resistant to PI uptake (Annexin-V+) represent those with plasma  
577 membrane extracellular surface exposed phosphatidylserine without plasma membrane  
578 damage and those stained with both Annexin-V-FITC and PI (Annexin-V+/ PI+) also have  
579 damaged plasma membranes. Data was obtained by flow cytometry and 10,000 events were  
580 analysed per sample. Data are means ± SD (n=3) \* P<0.05, \*\* P = 0.002,\*\*\*P<0.001 for  
581 increases v. DMSO control.

582

583 Fig 5. Detection of the 'active' form of caspase-3. The level of active caspase-3 expression  
584 was immuno-flow cytometrically determined 4, 12 and 24h after treatment with DMSO and 1,  
585 5 and 10  $\mu$ M BNIPSpd. 10,000 events were analysed per sample. Data are means  $\pm$ SD (n=6).  
586 \* P=0.05 or \*\*\* P<0.001 for comparison v. the respective DMSO control.

587

588 Fig 6. The relationship between general (pan)-caspase activity and membrane damage. FAM-  
589 VAD-FMK FLICA labelling with PI co-staining was used to flow cytometrically assess the  
590 level of caspase activity relative to membrane damage within 8h of treatment with BNIPSpd.  
591 The median fluorescence intensity from FAM-VAD-FMK FLICA labelling was used as a  
592 marker of cellular pan-caspase activity (A). The proportion of cells with FAM-VAD-FMK  
593 FLICA but no PI fluorescence (C+), co-fluorescing with FAM-VAD-FMK FLICA and PI  
594 (C+, PI+), fluorescing only due to PI (PI+) or showing no fluorescence with FAM-VAD-  
595 FMK FLICA and PI (NR) was used to establish the relationship between caspase activity and  
596 the occurrence of membrane damage (B) the latter as indicated by heightened PI fluorescence.  
597 10,000 events were analysed per sample. Data are means  $\pm$ SD (n=3). \* P<0.05 or \*\*\*  
598 P<0.001 for comparison v. the staining within the respective DMSO control.

599

600 Fig 7. 'Apoptotic' DNA fragmentation. Sub G1 (hypodiploid) DNA content (A), as an  
601 indicator of apoptotic DNA fragmentation, was determined from flow cytometric analysis of  
602 PI staining using ModFit LT software (Verity Software House, ME, USA). Cells were treated  
603 with DMSO or BNIPSpd. Data are means  $\pm$  SD (n=3). \* P<0.05, \*\*P<0.01, \*\*\* P<0.001 for  
604 comparison v. the respective DMSO control. Internucleosomal DNA fragmentation (B),  
605 considered a classical feature of apoptosis, was detected by agarose gel electrophoresis  
606 following 24h exposure to DMSO, 10 $\mu$ M etoposide (positive control for ladder formation) or  
607 BNIPSpd (1, 5 or 10 $\mu$ M). M, markers; C no treatment, D, Solvent (DMSO); E, etoposide.

608 Data is a representative image from one of three experiments (note image split for  
609 presentation).

610

611 Fig 8. Influence on cellular polyamine levels. Effects of polyamine-analogue exposure on  
612 spermine and spermidine was evaluated by HPLC after 12 (A) and 24 (B) h exposure to  
613 BNIPSpd. As an internal control, effects of polyamine analogues are compared with those of  
614  $\alpha$ -difluoromethylornithine, DMFO (5mM), an ornithine decarboxylase inhibitor. SPD,  
615 spermidine; SPM, spermine. Data are means  $\pm$  SD (n=3). \*P<0.05 or \*\*P<0.01 v. DMSO  
616 control.

617

618 Fig 9. The influence of caspase inhibitor z-VAD-fmk on DNA fragmentation and cell death.  
619 Cells were pre-treated with the cell permeable pan caspase inhibitor, z-VAD-FMK, prior to  
620 and during exposure to 10  $\mu$ M BNIPSpd. The extent of DNA ladders (A): lane M, Mw  
621 markers; lane 1, no solvent vehicle or inhibitor; lane 2, DMSO (solvent vehicle) treatment  
622 only; lane 3, z-VAD-FMK; lane 4, BNIPSpd; lane 5, BNIPSpd and z-VAD-FMK lane 6,  
623 etoposide (10 $\mu$ M) as positive control for DNA ladder (apoptosis) formation. Note gel-image  
624 splicing is for presentation; (B): the proportion of cells with hypodiploid DNA content, (C):  
625 the effect on cell proliferation and (D): cell death as determined at 24h from start of exposure.  
626 Data is either a representative image from three experiments (A), or are means  $\pm$  SD (n=3),  
627 data designated by a different letter is significantly different (P<0.05) relative to control or  
628 treatment at the respective time point.

629

630 Fig 10. BNIPSpd induced exogenous, non-cell derived, hydrogen peroxide formation. Using  
631 the phenol red assay hydrogen peroxide concentration was estimated in cell culture medium  
632 containing BNIPSpd. Catalase incubation was used to confirm the presence of hydrogen

633 peroxide in control and 50 $\mu$ M BNIPSpd incubations, reducing the absorbance attributed to  
634 resorufin formation to 1% and 5% of non-catalase treatment values respectively. Data are means  
635 (n=3). \*P<0.05 v. DMSO control

636

637

638

Figure 1 Bestwick et al.

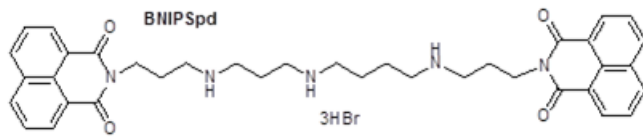


Figure 2 Bestwick et al.

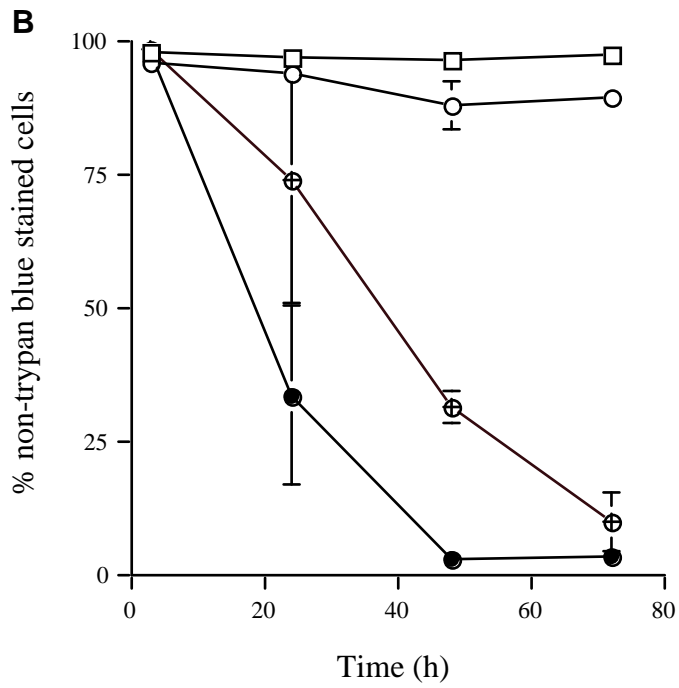
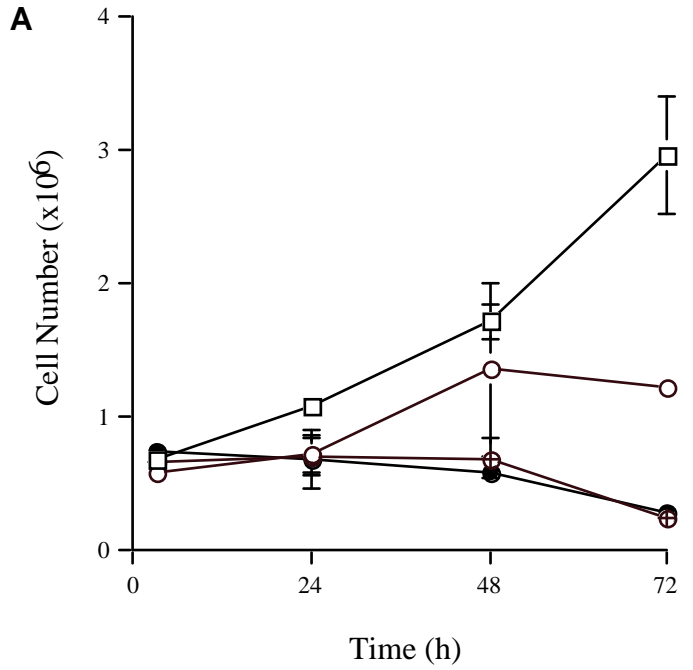




Figure 3 Bestwick et al.

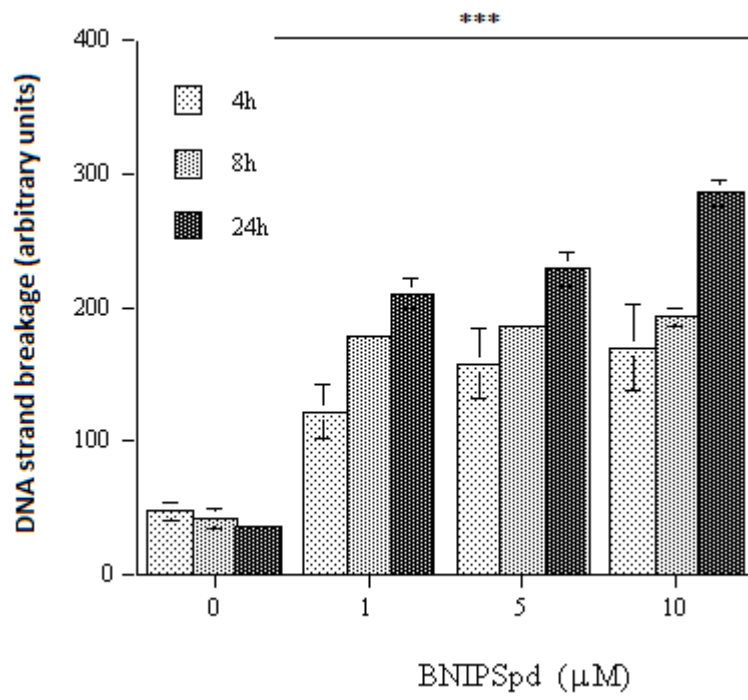


Figure 4 Bestwick et al.

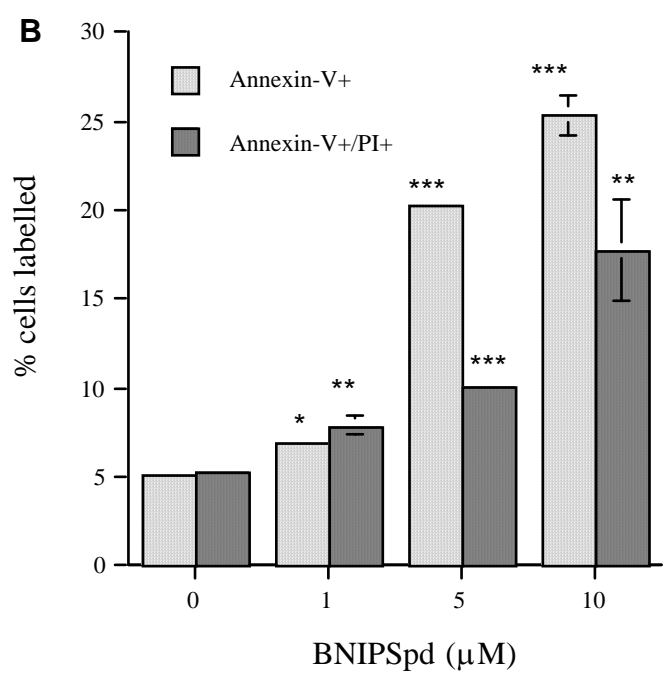
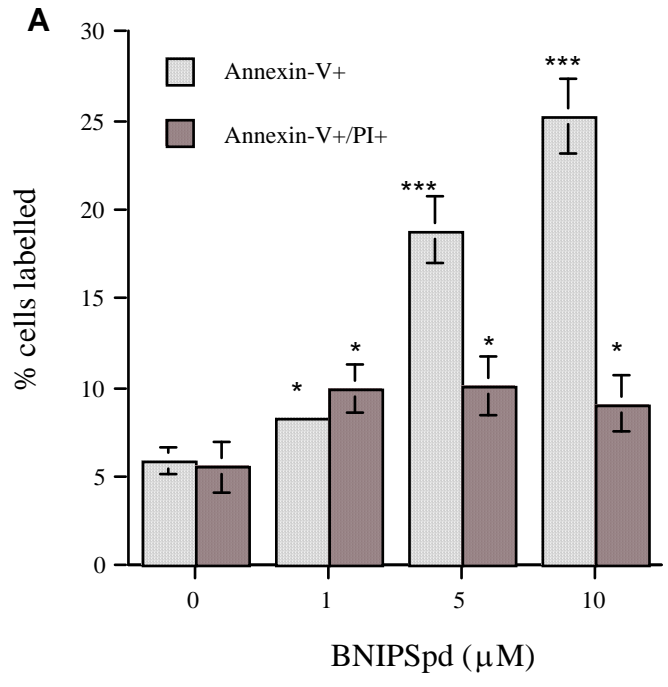


Figure 5 Bestwick et al.

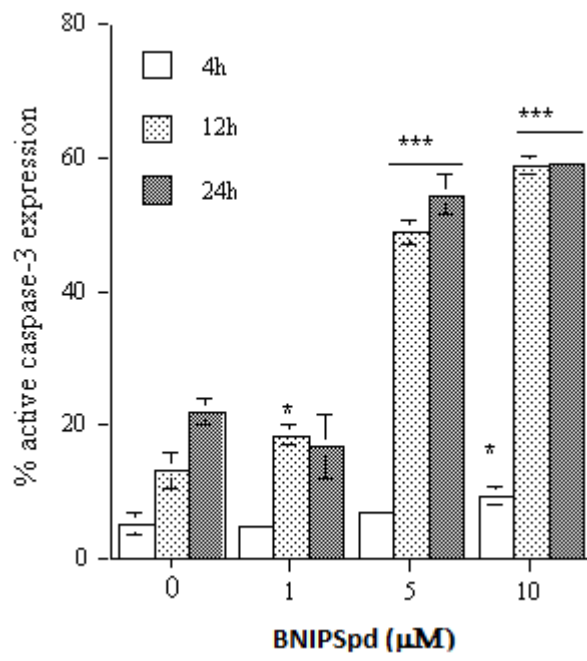
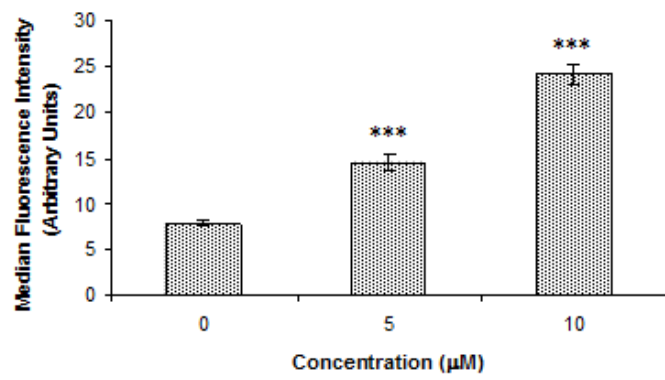


Figure 6 Bestwick et al.

A



B

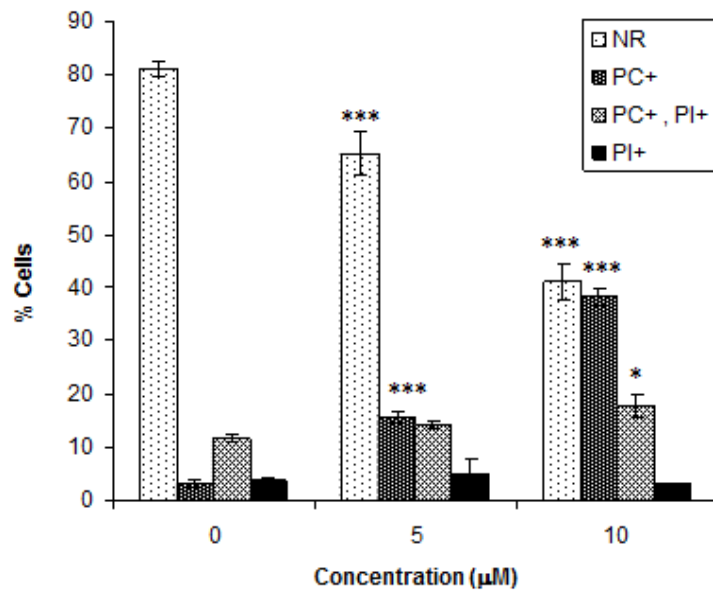
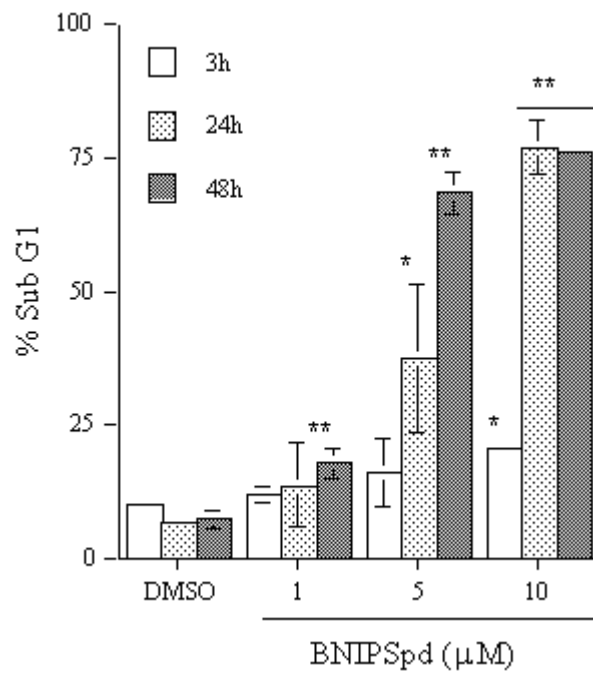


Figure 7 Bestwick et al

A



B

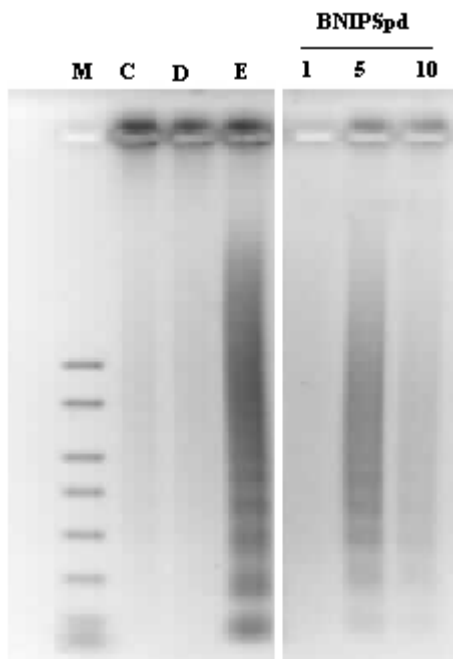


Figure 8 Bestwick et al

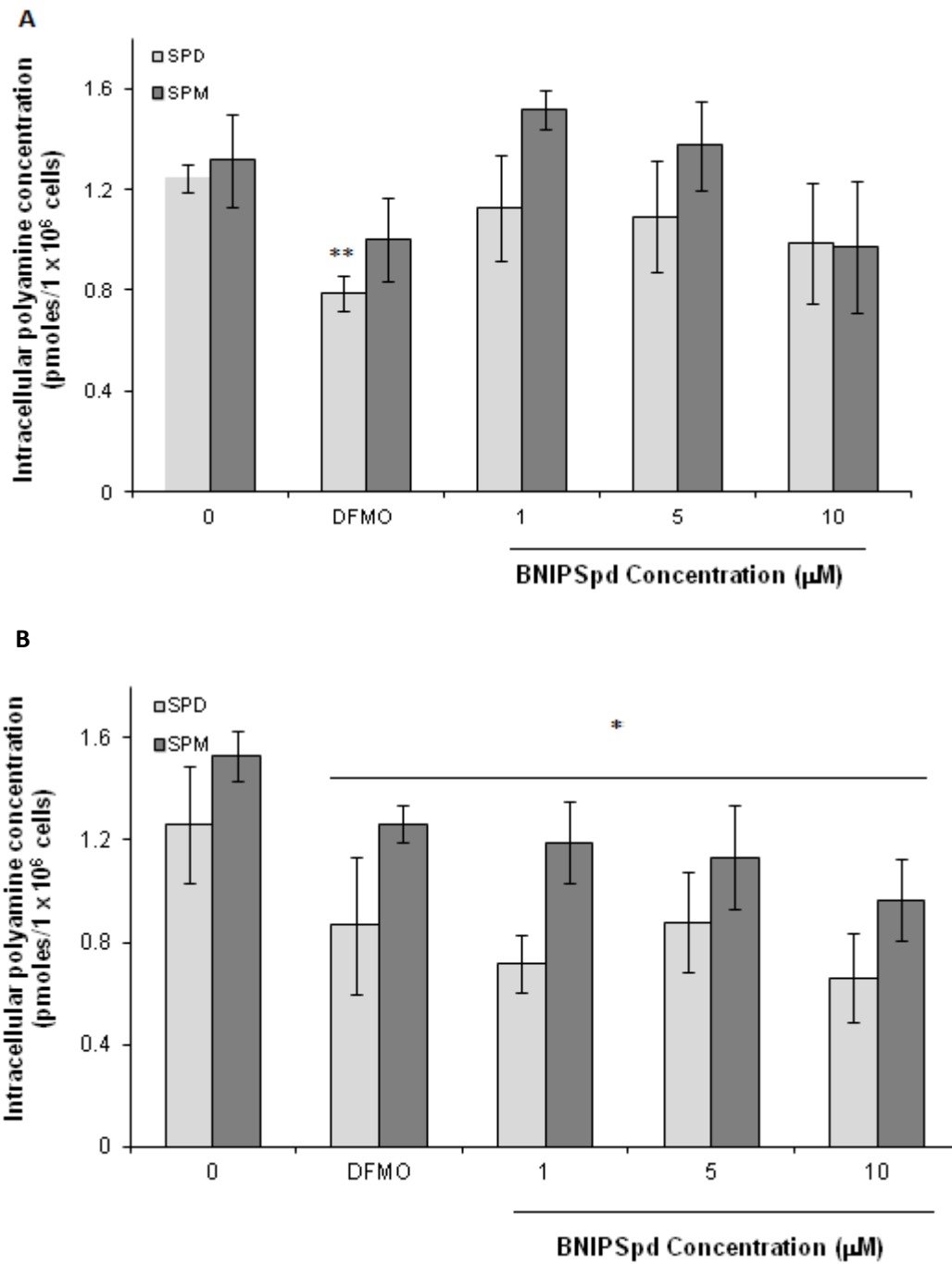


Figure 9 Bestwick et al

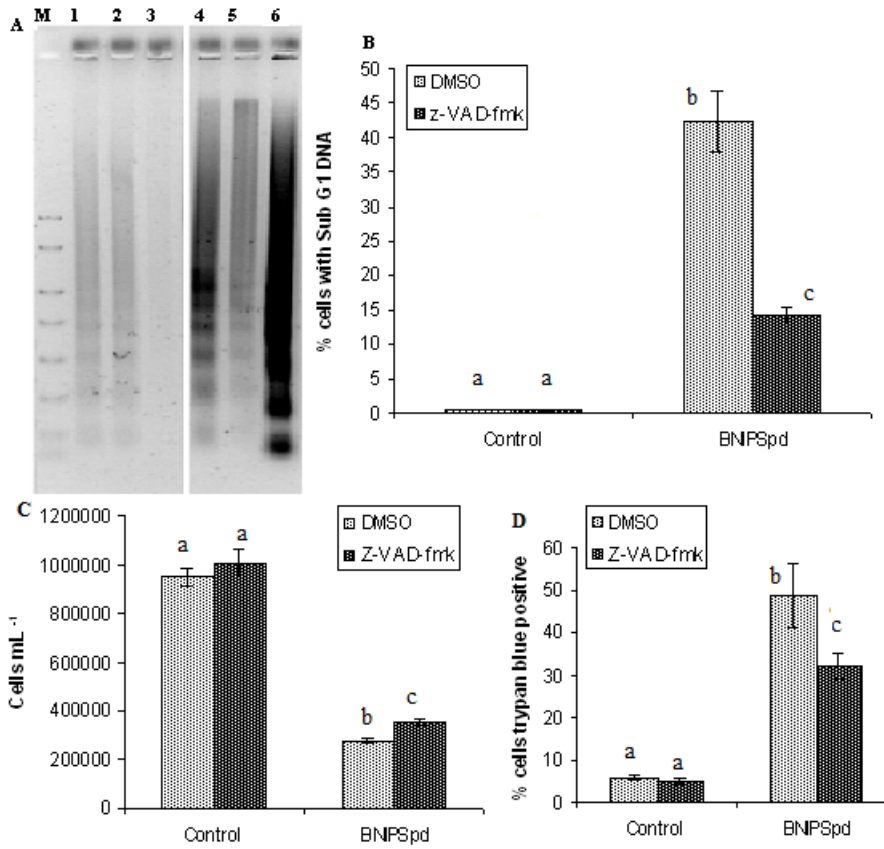


Figure 10 Bestwick et al

