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1	In vitro administration of sodium arsenite in mouse prepubertal testis induces germ cell
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3	

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- 12

#### 13 Abstract

High levels of arsenic contamination in drinking water pose serious health risks in numerous 14 15 countries. The documentation reporting arsenic toxicity on reproduction and development is increasing, with evidence of arsenic inducing fertility and developmental issues. Nonetheless, 16 the impact of arsenic exposure on the development of the male reproductive system is not fully 17 elucidated. In the present study, we have investigated the direct effects of arsenic on 18 prepubertal mouse testis using an *in vitro* testicular organ culture system. Culture medium was 19 20 supplemented with a range of concentrations of sodium arsenite, examining effects of low (0.5 and 1 µM) and high (10, 50, 100 µM) concentrations, in cultures of post-natal day 5 CD1 mouse 21 testis. In vitro exposure of low arsenic concentrations (0.5 or 1 µM) for 6 days did not cause 22 any change in the testicular morphology, germ cells density, or apoptotic marker cleaved 23

24	caspase 3 (CC3) expression. In contrast, exposure of prepubertal testis to high arsenic
25	concentrations (10, 50 or 100 $\mu$ M) induced drastic changes: severe destruction of testicular
26	morphology, with loss of seminiferous tubule integrity; a dose-dependent decrease in germ cell
27	density, and a hundred-fold increase in CC3 expression after 50 $\mu M$ arsenic exposure. In
28	conclusion, high arsenic treatment induced a dose-dependent induction of apoptosis and germ
29	cell loss in prepubertal mouse testis.
30	
31	Keywords:
32	Sodium arsenite; in vitro; prepubertal mouse; testis; toxicity.
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45 Arsenic poisoning has become an escalating problem in many developing countries.
46 Usually the contamination of drinking waters with arsenic occurs due to natural environmental

47 sources seeping into aquifers, mining, and other industrial activities (Sulanjari et al., 2015). In particular, in the past few decades its exposure through drinking water has vastly increased in 48 Bangladesh, India, China, and Cambodia (Smith, 2003; Xie, 2011). Recently, the risk of arsenic 49 50 exposure has also been reported to have worsened in Pakistan, exposing almost 50 million people to high levels of arsenic through drinking groundwater (Podgorski et al., 2017). Overall, 51 more than 150 million people around the world are affected by drinking arsenic-contaminated 52 water (Munoz et al., 2015): its chronic exposure is therefore a major concern for public health 53 (Abhyankar et al., 2012). 54

There is a clear link between exposure to inorganic arsenic and male reproductive 55 dysfunction (Feng et al., 2001; Kim and Kim, 2015; Lima et al., 2018; Wang et al., 2006). In 56 the testis, exposure to arsenic can alter spermatogenesis, interrupt steroidogenesis, and reduce 57 58 gonadotrophin as well as testosterone levels (Kim and Kim, 2015). In vivo studies investigating the reproductive toxicity of arsenic have shown effects on adult rodent reproductive system. 59 For instance, Chang et al. (2007) reported reduced testicular weight and sperm count in mice 60 after subchronic exposure to 20 or 40 mg/l sodium arsenite. Similarly, mice exposed to sodium 61 arsenite via drinking water at concentration of around 530 µ mol/l presented with a significant 62 reduction in sperm count and motility (Pant et al., 2001). A comparative study in male rats 63 exposed to arsenite or arsenate suggested a higher reproductive toxicity of the inorganic form 64 (Lima et al., 2018). A small number of in vitro studies have assessed effects of arsenic on 65 embryonic development and reproduction. Different concentrations of sodium arsenite exert 66 toxic effects on preimplantation development of mouse embryos, with 100 µM arsenic 67 concentration rapidly killing two-cell embryos, while 1 µM impairs blastocyst formation 68 (Muller et al., 1986). In vitro studies of arsenic on post-implantation mouse embryo 69 development confirmed that sodium arsenite possesses higher toxicity compared to sodium 70

arsenate (Chaineau et al., 1990; Tabocova et al., 1996). Moreover, Sertoli cell lines undergo
apoptosis after arsenic trioxide exposure (Kim et al., 2011).

In general, children are considered more sensitive to toxicants than adults (Scheuplein 73 et al., 2002). There is an accumulating evidence of a strong negative impact of arsenic exposure 74 75 in early life, includes link to increased mortality rates during both childhood (Rahman et al., 2013) and also adulthood (Yuan et al., 2007). Prepubertal testicular development is a highly 76 77 sensitive stage and it is possible that toxicants such as arsenic may act differently on testis of young compared to adult males. Presently, only a few animal studies have examined the effects 78 79 of arsenic on the reproductive axis of the prepubertal mammal: prepubertal rat prostate development has been found to be affected (Aquino et al., 2019), but that study provides little 80 information about the testicular cell populations that could be damaged (i.e. somatic and/or 81 82 germ cells); in one other study, prepubertal rats exposed to sodium arsenite (0.01 or 10 mg/l) for 30 days in the drinking water showed a significant disruption of normal testicular 83 84 morphology (da Cunha de Medeiros et al., 2019).

Here, we have investigated the effect of sodium arsenite exposure on immature mouse testis using an *in vitro* culture system that supports the developing neonatal testis. *In vitro* models have been commonly used to elucidate the metabolism of arsenic (Drobna et al., 2009), and the testis culture model has already been used for other toxicological studies on the male gonad (Allen et al., 2020; Lopes et al., 2016; Nakamura et al., 2019; Smart et al., 2018).

90

#### 91 **2.** Materials and Methods:

92 2.1. Animals

93 This work was approved by the Local Ethical Review Committee of the University of
94 Edinburgh and conducted according to UK home office regulations. CD1 male mice, obtained

95 from University of Edinburgh breeding stock, were kept at 14 h:10 h light and dark 96 photoperiod, provided *ad libitum* food (expanded RM3 diet, SDS; DMB Scotland) and water. 97 All mice used in the experiments were euthanized by decapitation at post-natal day (PND) 5 98 for tissue collection or by cervical dislocation at PND 12 for age-matched control, with the day 99 of the pup delivery considered as PND 0.

100 2.2. Tissue culture

101 PND 5 testes were collected, fragmented and cultured according to Lopes et al. (2016). In brief, for each culture run, testes from at least two mice were dissected, pooled together and 102 placed into dissection medium Leibovitz L-15 (Invitrogen, UK) supplemented with 3mg/ml of 103 bovine serum albumin (BSA; Sigma-Aldrich Ltd, UK) at 37°C. After removal of epididymis 104 and tunica albuginea, testes were fragmented into approximately 0.5mm<sup>3</sup> pieces, using a scalpel 105 106 blade. Fragments that were homogeneous in size were randomly distributed on a 24-well culture plate (Greiner Bio-one, UK), with testes obtained from two pups per litter for each 107 culture run. On the first day of culture (Day 1), each testicular fragment was placed on a floating 108 polycarbonate membrane (Whatman, Camlab Ltd, Cambridge, UK) per each well, containing 109 1 ml α-Minimal Essential Medium (MEM; Invitrogen, UK) supplemented with 10% Knockout 110 111 Serum Replacement (Invitrogen, UK); half of the medium was replaced with fresh medium on Days 4 and 6. The culture plate was incubated at 34°C, 5% CO<sub>2</sub>. Fragments were left untreated 112 for the first 24 h (Day 1). From Day 2 of culture, testicular fragments were exposed to different 113 concentrations of sodium arsenite prepared from a stock solution of 0.05 mol/l (0.1N) 114 Tritipur®reag. (Merck, Germany). Sodium arsenite solution was diluted in culture medium and 115 final concentrations were prepared through serial dilutions. Two sets of experiments (low and 116 high arsenic doses) were performed with cultures maintained for seven days, of which six days 117 were with arsenic exposure (Days 2-7): the *in vitro* study on mouse embryos arsenic exposure 118 was used as a reference for selection of the highest dose (Muller et al., 1986). The first 119

120 experiment (low arsenic doses) comprised of 0.5 and 1 µM sodium arsenite concentrations, whereas the second experiment (high arsenic doses) included concentrations of 10, 50 and 100 121 µM arsenic. In each culture run, the arsenic effect was compared to a culture control that had 122 123 been maintained throughout in arsenite-free medium. Each experiment was repeated six times, including six culture runs, six histological and immunohistochemistry (IHC) runs and six sets 124 of image analysis. On Day 7, tissues were fixed for 1 h in 10% neutral buffered formalin (Sigma 125 Aldrich, UK) prior to IHC, or in Bouin's fluid (Sigma Aldrich Ltd, UK) prior to histological 126 examination. Tissues were placed in 3% agar (Sigma Aldrich Ltd, UK) blocks and then 127 128 embedded in paraffin wax prior to sectioning.

#### 129 2.3. Histology/ Immunohistochemistry

Five µm thick serial sections were cut. A middle section from each tissue fragment was 130 131 dewaxed, rehydrated through a series of decreasing alcohol concentrations, stained with hematoxylin and eosin (H&E), and used for morphological analysis. A serial section next to 132 the middle section used for histological examination was selected for IHC reactions. Sections 133 were dewaxed, rehydrated, and processed for antigen retrieval with citrate buffer (10mM 134 sodium citrate, pH 6, Fisher Scientific, UK). In between each step, slides were washed with 135 phosphate-buffered saline (Fisher Scientific, UK) with 0.1% Triton X (PBSTx, Sigma-Aldrich, 136 UK). Blocking used 20% normal goat serum diluted in PBSTx supplemented with 5% w/v 137 BSA for 1 h at room temperature. Primary antibodies were applied and incubated overnight in 138 a humidified environment: (i) rabbit anti-cleaved caspase 3 (CC3) antibody (9661S, 1:500 139 dilution; Cell Signaling Technology, USA) and (ii) mouse anti-DEAD-box helicase 4 140 (DDX4)/Mouse Vasa Homologue (MVH) antibody (ab27591, 1:100 dilution; Abcam, UK). 141 After washing, appropriate secondary antibodies were applied (both 1:200 dilution): (i) goat 142 anti-rabbit biotinylated (E0432, DakoCytomation, Denmark) and (ii) Alexafluor goat anti-143 mouse IgG1 568nm (A-21124, Invitrogen, UK). Omission of primary or secondary antibodies 144

were used as technical negative controls. For CC3, the visualization reagent Streptavidin Alexa
Fluor 488 conjugate (S-32354, Invitrogen, UK) was subsequently applied at 1:200 dilution. All
sections were counterstained with 4,6-Diamidino-2-phenylindole (DAPI; Invitrogen, UK) and
mounted with Vectashield hard-set mounting medium (Vector Laboratories, USA).

149 2.4. Image attainment and analysis

H&E stained sections were photomicrographed (DMLB Leica microscope, Leica 150 Microsystems Ltd, UK), with images taken using a DFC369FX camera on a Leica DM5500B 151 microscope (Leica Microsystem Ltd, UK), using filters for DAPI, A4 (BP 360/40); for Alexa 152 Fluor 488, N3 (BP 546/12); for Alexa Fluor 568, Y5 (BP 620/60). Image J software was used 153 for image analysis, with the assessor blind as to treatment. Germ cells (MVH<sup>+</sup>) were analyzed 154 155 by manual counting and the total number given relative to the area of seminiferous tubule, in 156 order to obtain a value for germ cell density (number of cells that were MVH-positive divided by seminiferous tubule area in mm<sup>2</sup>). Density was also determined for whole section area 157 (number of MVH positive cells divided by the area of section in mm<sup>2</sup>) to allow determination 158 of germ cell density even where tubule integrity had been disrupted to the extent that precluded 159 measurement of tubule area. For CC3, fluorophore area was measured as a percentage of 160 tubule, interstitium and section area (proportion of CC3-positive area in mm<sup>2</sup> relative to area 161 of seminiferous tubule, interstitium or whole section, all in mm<sup>2</sup>). 162

163 2.5. Statistical analysis

164 GraphPad Prism was used for statistical analyses. The statistically significance of any 165 difference between the control group and the arsenic-treated groups was determined by one-166 way ANOVA, followed by Dunnett's *post hoc* test where the ANOVA was statistically 167 significant. For analyses including only 2 experimental groups (comparison of the control 168 group to the 10  $\mu$ M sodium arsenite treatment group for: density of germ cells in the

169	seminiferous tubules, and percentage of seminiferous tubule or interstitium area positive for
170	CC3), paired T-test was performed. Data is expressed as mean $\pm$ SEM, with p $\leq$ 0.05 considered
171	statistically significant.

### 173 **3. Results**

174 3.1. Morphological analysis

To assess the impact of sodium arsenite on the developing testis of prepubertal (PND 5) mouse, testis fragments were exposed *in vitro* for six days to low or high concentrations of sodium arsenite, compared with arsenite-free culture control testicular fragments in each case. Morphological changes were evaluated on H&E stained sections (Fig. 1).

179 Figure 1

## A: Uncultured control



B: Low Sodium Arsenite Doses Experiment



C: High Sodium Arsenite Doses Experiment



182 Figure 1. Morphological changes of prepubertal mouse testis induced by *in vitro* exposure to sodium arsenite. Representative images of hematoxylin and eosin stained sections of 183 uncultured and cultured testis fragments. Pre-culture control tissue (Ai) obtained from PND 5 184 mouse showed germ cells only at the spermatogonial stage (black arrow), while testes from 185 mice age-matched to be equivalent to end of culture control testis (PND 12: Aii) showed germ 186 187 cells that have entered meiosis (spermatocytes, white arrow). Representative images showing the effect on testis morphology of low (B) and high (C) concentrations of arsenic. Control 188 tissues in both the low dose (Bi) and high dose experiment (Ci), as well as low doses of arsenic 189 (Bii: 0.5 µM sodium arsenite; Biii: 1 µM sodium arsenite) preserved normal morphology, with 190 191 germ cells lining the basal membrane (black arrow), some cells undergoing mitosis (black arrow head) and several differentiating germ cells (spermatocytes, white arrow) 192 accumulating in the forming lumen of well-defined seminiferous tubules. High doses of arsenic 193 caused dramatic morphological changes, starting at 10 µM sodium arsenite (Cii) with evidence 194

of apoptotic/necrotic cells (white arrowhead), and with disruption or loss of seminiferous tubule integrity at 50  $\mu$ M (Ciii) and 100  $\mu$ M sodium arsenite (Civ). B-C: scale bars represent 100  $\mu$ m; A and insets: scale bars represent 50  $\mu$ m.

198

The morphology of control testicular fragments was well preserved at the end of the six 199 day culture period for both sets of experiments: necrosis was not present in the centre of the 200 section, seminiferous tubule structure and integrity was maintained, with many germ cells lined 201 along the basement membrane of all tubules, several actively undergoing a mitotic division 202 (Fig. 1 Bi and Ci). Control tissues closely resembled the morphology of equivalent age-203 matched PND 12 testis (Fig. 1 Aii), with mitotically active spermatogonia lining the basal 204 205 lamina and spermatocytes in the lumen of the seminiferous tubules, showing that the culture 206 system supports physiological testicular development, given that pre-culture control testis displayed presence of germ cells only at the spermatogonial stage (Fig. 1 Ai). Exposure to low 207 sodium arsenite concentrations did not cause any morphological changes in seminiferous 208 tubules when compared to the control group: seminiferous tubules maintained their 209 physiological shape with abundant spermatogonial germ cells present in tissues treated with 210 211 both low arsenic doses (0.5 or 1 µM; Fig. 1 Bii and Biii). In contrast, higher concentrations of sodium arsenite severely disrupted normal testicular morphology (Fig. 1 Cii, Ciii, and Civ). 212 Apoptotic/necrotic cells were present, as evidenced by dense dark staining of the chromatin 213 and more intense pink/red staining of the cytoplasm, and seminiferous tubules were irregularly 214 defined, particularly at the 50 or 100 µM arsenic doses. 215

216 3.2. Germ cell density

217 Density of spermatogonial germ cells (MVH<sup>+</sup>) was assessed at the end of the culture
218 period for both sets of experiments, low and high arsenic doses (Figs 2 and 3 respectively).

#### 220 Figure 2

#### Low Sodium Arsenite Doses Experiment





Figure 2. Effect of low sodium arsenite concentrations *in vitro* exposure on germ cell density and apoptosis in the prepubertal mouse testis. Photomicrographs showing immunohistochemical localization of MVH (red) and CC3 (green) expression, counterstained with DAPI (blue) in representative images of low dose arsenic experiment. In control tissue (Ai), as well as in the testis fragments exposed to low doses of arsenic (Aii, 0.5  $\mu$ M sodium arsenite; Aiii, 1  $\mu$ M sodium arsenite), germ (MVH<sup>+</sup>) cells were abundant and apoptotic (CC3<sup>+</sup>)

228	areas were sparse. Scale bars represent 100 $\mu$ m. Graphs show germ cell density: the number of
229	$MVH^+$ cells per mm <sup>2</sup> of seminiferous tubules (Aiv) and per mm <sup>2</sup> of section (Av). Density of
230	germ cells was unaffected in the low arsenic dose treated testis fragments compared to that of
231	control tissue. Data represents individual data points (N=6), mean $\pm$ SEM.

**Figure 3** 



High Sodium Arsenite Doses Experiment

234

Figure 3. Effect of high sodium arsenite concentrations *in vitro* exposure on germ cell density and apoptosis in the prepubertal mouse testis. Photomicrographs showing immunohistochemical localization of MVH (red) and CC3 (green) expression, counterstained with DAPI (blue) in representative images of high dose arsenic experiment. In control tissue

240 (Ai), germ (MVH<sup>+</sup>) cells were abundant, and apoptotic (CC3<sup>+</sup>) areas were sparse. In contrast, in all arsenic treated groups (Aii, 10 µM; Aiii, 50 µM; Aiv, 100 µM sodium arsenite) a lower 241 density of germ cells was present, with vast areas of CC3 expression. Graphs show germ cell 242 density: the number of MVH<sup>+</sup> cells per mm<sup>2</sup> of seminiferous tubules (Av) and per mm<sup>2</sup> of 243 section (Avi). A significant reduction in germ cell density occurred in all three groups exposed 244 to high concentrations of sodium arsenite compared to control tissue. Note that it was not 245 possible to obtain data about germ cell density within seminiferous tubules in the 50 and 100 246 µM arsenic dose groups due to severe loss of tubule integrity. Data represent individual data 247 points (N=6), mean  $\pm$  SEM. \*\* and \*\*\* indicate p < 0.01 and 0.001, respectively. 248

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Control tissues had many MVH<sup>+</sup> cells lining the seminiferous tubule basal membrane 250 251 (Fig. 2 Ai and 3 Ai). Similarly, germ cells were numerous in testis fragments exposed to low doses of sodium arsenite (Fig. 2 Aii and Aiii). In contrast, testis exposed to high doses of arsenic 252 contained only sparsely distributed germ cells (Fig. 3 Aii, Aiii, and Aiv). Analysis confirmed 253 that, in comparison to control tissue, germ cell density per tubule area and per section area of 254 treated tissues in the low arsenic dose experiment remained unaffected (p-value 0.47 and 0.99 255 256 for 0.5 and 1 µM respectively; Fig. 2 Aiv and Avi); in contrast, in the high arsenic doses experiment, sodium arsenite exposure caused a significant decrease in germ cell density (Fig. 257 3 Av and Avi). Compared to the control group, germ cell density per tubule area was 258 significantly reduced in 10 µM arsenic treated tissues, even though tubule integrity was still 259 maintained (p < 0.01) (Fig. 3 Av). Due to seminiferous tubule disruption in tissues exposed to 260 higher doses of sodium arsenite, it was not possible to accurately define tubule margins: as a 261 consequence, in these cases density of MVH<sup>+</sup> cells was assessed only relative to testis section 262 area (Fig. 3 Avi): germ cell density per area of testis section decreased by a 2.5-fold magnitude 263

after 10  $\mu$ M arsenic treatment, while after 50 or 100  $\mu$ M exposure the decrease was 3-fold for all three doses (p < 0.001 for all).

266 3.2. Analysis of apoptosis

Activation of the apoptotic marker CC3 was measured at the end of the culture period. In the experiment using low arsenic doses, expression of CC3 was low overall, with arsenic exposed tissues (Fig. 2 Aii and Aiii) similar to those of control tissues (Fig. 2 Ai). In contrast, exposure to high doses of sodium arsenite induced increased CC3 expression (Fig. 3 Aii, Aiii, and Aiv) compared to control tissue (Fig. 3 Ai).

Analysis of apoptosis after exposure to low arsenic doses, revealed that the treatment did not affect the area of CC3 expression (Fig. 4 A): the percentage of the seminiferous tubule (Fig 4 Ai), interstitium (Fig 4 Aii) and total section (Fig 4 Aiii) area expressing CC3 was similar to that of control tissue (p = 0.968, 0.88 and 0.958, respectively).





Figure 4. Apoptosis of the prepubertal mouse testis was induced by *in vitro* exposure to 280 high arsenic doses. Percentage of CC3 positive area of testicular fragments was not affected 281 by low arsenic doses exposure compared to control (Ai, Aii, Aiii). However, exposure to 10 282 283 µM arsenic increased the CC3 positive area of seminiferous tubules (Bi). In the interstitium, exposure to 10 µM did not affect CC3 expression compared to control tissue (Bii). The 284 percentage of CC3 expression per total section area was increased compared with control after 285 286 50 µM arsenic exposure only (**Biii**). Note that it was not possible to obtain data about apoptosis within seminiferous tubules in the 50 and 100 µM arsenic dose groups due to the severe loss 287 of tubule integrity. Data represent individual data points (N=6), mean ± SEM. \* and \*\*\* 288 indicate p < 0.05 and 0.001, respectively. 289

As per the analysis of germ cell density, in the high arsenic dose experiment, the loss 291 of the seminiferous tubule integrity after exposure to 50 and 100 µM sodium arsenite rendered 292 it impossible to analyze CC3 expression per unit tubule area. In the 10 µM treatment group, 293 1.5% of the seminiferous tubule area expressed CC3, significantly increased over expression 294 in the control tissue (p < 0.05; Fig. 4 Bi). No change was seen outside of the tubule areas, with 295 the percentage of the interstitium expressing CC3 unaffected by treatment (p = 0.43; Fig. 4 296 Bii). When examining the tissue as a whole, CC3 expression increased a hundred-fold in tissues 297 exposed to 50 µM sodium arsenite compared to control tissue (Fig. 4 Biii). All three of the high 298 arsenic doses exhibited increased CC3 expression compared to control tissue (3.2%, 10.5%, 299 and 2.9% of the section area CC3-positive in the 10, 50, and 100 µM treatment groups 300 301 respectively, compared to 0.1% of the tissue CC3-positive in the control tissue), although only the 50  $\mu$ M sodium arsenite group reached statistical significance (p < 0.001; Fig. 4 Biii). 302

303

#### 304 4. Discussion

305 Arsenic is a well-known carcinogen present naturally in the Earth's crust. Human and animal exposure to arsenic is a common consequence of consumption of contaminated food or 306 water, or arising from the use of contaminated water for crop irrigation. Arsenic is also used in 307 308 a variety of industrial processes, contributing to its environmental distribution as well as representing a risk of occupational exposure. Both chronic and acute exposure to arsenic can 309 310 contribute to a spectrum of diseases in humans. Effects of arsenic on adult male reproduction has been studied, with reports that arsenic inhibits steroidogenesis, causes spermatotoxicity, 311 and adversely impacts on the weight of the reproductive organs of mice in vivo (Pant et al., 312 2004; Sarkar et al., 2003). Similarly, in vivo arsenic exposure induces biochemical and 313

314 morphological changes in adult rat testis (de Araujo Ramos et al., 2017). Given the marked differences between the pre- and post-pubertal testis, it is important to investigate effects of 315 sodium arsenite directly on the developing male reproductive tissues; with, in contrast, little 316 317 information available: thirty-day administration of sodium arsenite to prepubertal rats induced dysfunction of normal prostate development (Aquino et al., 2019), spermatogenesis and 318 epididymal structure (da Cunha de Medeiros et al., 2019). Here, we have used an in vitro culture 319 system of prepubertal mouse testis to assess the impact of arsenic on the testicular development. 320 The culture system used is capable of supporting testicular development (Sato et al., 2011) and 321 322 has already been used by our group (Allen et al., 2020; Lopes et al., 2016; Smart et al., 2018) as well as others (Chapin et al., 2016; Nakamura et al., 2019; Park et al., 2020) for toxicological 323 studies on the prepubertal testis. In the results reported here, consistency and health of the 324 325 control tissue, along with the presence of active spermatogenesis within the vast majority of 326 seminiferous tubules and the absence of any necrotic central area, strengthens the suitability of this culture model for the evaluation of the short-term impact of a toxicant on the immature 327 testis (Chapin et al., 2016; Nakamura et al., 2019). Use of such in vitro models, in addition to 328 allowing strict control over exposure conditions, also complies with regulations requiring 329 reduction of animal suffering by avoiding *in vivo* administration. This is particularly important 330 here because there is little information about the toxicological effect of arsenic upon neonatal 331 mice and because it allows examination of the effects of arsenic on the early reproductive 332 333 system whilst avoiding the inherent difficulties that come with administration of a toxicant to early neonatal animals (Twaddle et al., 2019). Nonetheless, the absence of physiological 334 pharmacokinetics of arsenic in the body, inevitably influencing circulating and tissue arsenic 335 336 levels, needs to be taken into account when extrapolating data from the present work, as for any in vitro study. It is known that some metabolic processes, such as methylation, increase 337 renal clearance, reducing arsenic toxicity, although it can also produce even more cytotoxic 338

compounds (Drobna et al., 2009; Twaddle et al., 2018a). Furthermore, a non-linear relationship
has been observed between dose of arsenic administered to experimental animals and its
metabolism and toxicity, given which it could be difficult to interpret *in vivo* experiments that
use doses higher than human exposure (Twaddle et al., 2018b).

Exposure to low sodium arsenite concentrations (0.5 or  $1 \mu M$ ) for six days did not 343 appear to affect mouse testis integrity, with regularly shaped seminiferous tubules lined by 344 germ cells at spermatogonial and spermatocyte phases. On the other hand, high arsenic 345 concentrations (10, 50, or 100 µM) caused significant changes in the testicular structure, with 346 a dose-response pattern: sodium arsenite at 10 µM concentration induced a significant damage 347 with a number of degenerating cells, despite leaving the seminiferous tubule morphology 348 reasonably well distinguishable, while both 50 and 100 µM sodium arsenite treatments 349 350 drastically compromised seminiferous tubule structure. In all three high arsenic doses, the germ cell population was drastically reduced. The extensive disorganization along with loss of germ 351 cells, shape, as well as shrinkage of the seminiferous tubule observed in the present study after 352 high arsenic exposure is in line with similar observations previously reported in the adult testis 353 after arsenic exposure in vivo (Baltaci et al., 2016; Ferreira et al., 2012). A study on the effect 354 of arsenic on prepubertal rats showed abnormal testicular structure with impairment of Sertoli 355 cell number (da Cunha de Medeiros et al., 2019). That study revealed no dose-response pattern, 356 357 with the both low and high doses administered causing similar effect on the testis, in contrast 358 with results here where a difference effects are observed in low and high concentrations. Nonetheless, a direct comparison between the two studies is challenging, as the in vivo study 359 on rats used only two arsenic doses and for an extended (30-day) period of exposure (da Cunha 360 361 de Medeiros et al., 2019), while our in vitro study exposed mouse tissues to five different concentrations for six days. A dose-dependent reduction of germ cells has been already 362

reported in adult mice exposed *in vivo* to sodium arsenite both short- and long-term
(Sanghamitra et al., 2008; Zeng et al., 2019).

Results here also showed an increase in the density of apoptotic cells present within 365 seminiferous tubules after exposure to 10 µM sodium arsenite, with an increase in apoptosis 366 also seen in the 50 µM sodium arsenite treatment group: in the latter case, tubules were ill 367 368 defined, and so analysis could only examine density of apoptotic cells across the entire testicular section. Paradoxically, the increased expression of CC3 after exposure to 100 µM of 369 sodium arsenite did not reach statistical significance. One possible explanation could be that at 370 the top dose, apoptotic cells died shortly after arsenic administration and the CC3 signal had 371 disappeared relatively rapidly, long before the six days culture end point: the entire apoptotic 372 cell death process can be complete within as short a time as 2-3 hours (Elmore, 2007). 373 Apoptosis is self-destruction process which can occurs by intrinsic and/or extrinsic pathways 374 (Dua et al., 2015). Reactive oxygen species (ROS) produced by mitochondria are considered 375 as the biochemical mediators of apoptosis (reviewed in Orrenius et al., 2015), thus a high 376 enough accumulation of ROS can initiate apoptosis (reviewed in Redza-Dutordoir and Averill-377 Bates, 2016). Arsenic is known to cause oxidative stress by increasing ROS production and 378 379 reducing antioxidant defense systems (Yamanaka et al., 1991). Available literature indicates 380 that *in vivo* arsenic exposure can cause an increase in testicular oxidative stress and apoptosis 381 (Das et al., 2009; Uygur et al., 2016). Here, it is possible that high arsenic exposure to the 382 prepubertal testis culture might have induced ROS generation and oxidative stress, thus resulting in cleaved caspase-3-mediated apoptosis. 383

According to the World Health Organization (WHO), water is safe for drinking only where there are concentrations of arsenic that are less than 10 parts per billion (ppb) (Brown and Ross, 2002), but the concentrations of arsenic in the drinking water in some areas of developing countries such as Pakistan, Bangladesh, Bihar and India are well above that limit 388 (Brammer and Ravenscroft, 2009; Sanjrani et al., 2017). For instance, in Bangladesh, the national standard for arsenic in drinking water is set at 50 ppb, although survey results have 389 shown that 25% of wells providing domestic water exceed even that level (Ravenscroft, 2005). 390 Analysis of water from irrigation shallow tube wells in Bangladesh has revealed that most of 391 them produce water with levels above 100 ppb arsenic, in several instances even above 200 392 ppb arsenic (Islam, 2005). Top arsenic concentrations of 1891 ppb have been found in Bihar 393 and India (Ghosh, 2007), with the highest of all in Nepal, 2629 ppb arsenic (Shresta, 2003). 394 The doses of sodium arsenite used in the present study are greater than the WHO safe limits of 395 396 arsenic in the drinking water, but the doses used in the low arsenic study are well within the range of arsenic concentrations found in drinking water in several developing countries (Ghosh, 397 2007); our low arsenic doses of 0.5 and 1 µM correspond to 64 and 129 ppb, respectively. In 398 399 the present study, lower doses administered for the short period of six days did not affect 400 testicular development. These results cannot, however, exclude the possibility that chronic long-term exposure to such concentrations of arsenic may be detrimental to the prepubertal 401 402 testis, and further longer term studies are advisable. Several studies on rodents have revealed arsenic accumulation in the testis and other accessory sex organs (Dua et al., 2015; Pant et al., 403 2001; Prathima et al., 2018). Such persistent nature after accumulation in the reproductive 404 organs can also adversely impact the developing testis in both human and animals. 405 Furthermore, it is possible that other physiological processes have been compromised outside 406 407 those assessed in the present study that might influence fertility in the adulthood. In the high arsenic doses experiment here, the lowest concentration of 10 µM, corresponding to 1299 ppb 408 arsenic, was highly detrimental to normal testicular physiology even after six days of exposure, 409 410 with increased cell death and a reduction in spermatogonial germ cell density. Such dose is below the top concentrations of arsenic found in irrigation shallow tube wells, wells that are 411 used as a source of drinking water for local populations in rural areas of developing countries. 412

413 Data here suggest a potential concerning scenario for the population living in the highly arsenic contaminated rural areas. Caution needs to be exercised before extrapolating data obtained 414 from an *in vitro* mouse study such as this to the situation in human and animal populations, in 415 416 light of the limitations previously discussed: as such, further studies are required in order to confirm the translational risk of future fertility impairment for young boys. This is likely to 417 include the need for in vivo studies, in order to fully evaluate the long-term impact, and the 418 effect of metabolic processes on arsenic circulating levels and toxicity. Comparative in vivo 419 studies would also be required to validate the *in vitro* system. Such investigation should be 420 421 extended to the fertility of large animals; in areas where animal husbandry is an important economic factor, arsenic-induced infertility of farm animals could represent a further financial 422 burden for the unstable economy of farmers in developing countries. 423

424

In conclusion, using a short term culture system of mouse prepubertal testis, this study
shows that exposure to high levels of arsenic for six days caused a dose dependent reduction
of spermatogonial germ cells and a marked increase in testicular cell apoptosis.

428

#### 429 Conflict of Interest

430 The authors declare no conflict of interest.

431

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