Genomic damage in children accidentally exposed to ionizing radiation: a review of the literature

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41 Abstract

42 During the last decade, our knowledge of the mechanisms by which children respond to exposures 43 to physical and chemical agents present in the environment, has significantly increased. Results of 44 recent projects and programmes focused on children's health underline a specific vulnerability of 45 children to environmental genotoxicants. Environmental research on children predominantly 46 investigates the health effects of air pollution while effects from radiation exposure deserve more 47 attention. The main sources of knowledge on genome damage of children exposed to radiation are 48 studies performed after the Chernobyl nuclear plant accident in 1986. The present review presents 49 and discusses data collected from papers analysing genome damage in children environmentally 50 exposed to ionizing radiation. Overall, the evidence from the studies conducted following the 51 Chernobyl accident, nuclear tests, environmental radiation pollution and indoor accidental 52 contamination reveals consistently increased chromosome aberration and micronuclei frequency in 53 exposed than in referent children.

54 Future research in this area should be focused on studies providing information on: (a) Effects on 55 children caused by low doses of radiation; (b) effects on children from combined exposure to low 56 doses of radiation and chemical agents from food, water and air; and (c) specific effects from 57 exposure during early childhood (radioisotopes from water, radon in homes). Special consideration 58 should also be given to a possible impact of a radiochemical environment to the development of an 59 adaptive response for genomic damage. Interactive databases should be developed to provide 60 integration of cytogenetic data, childhood cancer registry data and information on environmental 61 contamination. The overall aim is to introduce timely and efficient preventive measures, by means 62 of a better knowledge of the early and delayed health effects in children resulting from radiation 63 exposure.

64

Key words: child, ionizing radiation, environment, chromosome aberration assay, micronucleusassay, Chernobyl

67 Introduction

The cancer incidence in children has increased during the last few decades in different parts of the world [1, 2]. The present occurrence per year is a 1% average annual increase in incidence (p < 0.0001) has been estimated from the European cancer incidence database, including some 110,000 childhood cancer cases from 63 population based registries [1]. Although this increase may partly reflect better diagnostics, its aetiology is probably also associated with parental, intrauterine and postnatal exposure to xenobiotics including low LET ionizing radiation (e.g., X-rays and γ -rays). [3]

75 Indeed, the occurrence of site specific cancer in children is different than in adults, suggesting that76 childhood cancers reflect foetal development and exposure [4,5].

77 Children live in complex radiochemical environments and share all types of exposures with their 78 parents [Figure 1]. Over the last few decades genotoxicological population studies have mostly 79 focused on occupational exposures. Exceptions were cases of accidental exposure of the general 80 population, including children. Recently, genotoxicologists have paid increasing interest in studies 81 of children addressing the issue of whether they are more susceptible to environmental exposures to 82 physical and chemical agents than adults [6, 7,8].

Based on the available evidence of quantitative health risks associated with radiation exposure,
public dose limits of exposure from mining or nuclear plants are currently set at 1mSv/yr above
background [9], but still there is no specific legislation concerning children, although such exists for
occupational exposure during pregnancy [10].

87 Except for a few studies of children after accidental overexposures, available data on the 88 consequences of radiation exposure in children are mostly limited to the monitoring of young 89 victims following April 26, 1986 when the world's worst nuclear power accident occurred at 90 Chernobyl in the former USSR (now Ukraine). The Chernobyl nuclear disaster affected a vast area 91 of Europe and may still contribute to genome damage in large areas of Ukraine and Belarus due to 92 the environmental persistence of some radionuclides [11].). Information on genome damage caused

by radiation from this and also other nuclear accidents in the former Soviet Union is, however, only
partially available to the scientific community, because scientists of the former USSR still publish
in national rather than international journals.

96 Sixty years ago in Hiroshima and Nagasaki mankind witnessed the first nuclear weapon attack 97 which for the first time in history had transgenerational consequences. The difference in the type of 98 exposure in Hiroshima and Nagasaki versus Chernobyl is that while the nuclear bombing resulted in 99 massive exposure to mixed gamma/neutron radiation, the Chernobyl accident caused an acute 100 exposure, followed by a long-term internal exposure mostly to low doses of gamma radiation [12]. 101 Differences in the type of exposure are reflected in the difference of cancer incidence between these 102 two nuclear disasters. After the Hiroshima and Nagasaki detonation, ionizing radiation-induced 103 leukaemia occurred in children 5 to 6 years later, while in adulthood the breast cancer incidence 104 increased in women who were exposed before puberty [13]. Recent epidemiological reports from 105 Ukraine and Belarus confirm an increased number of cases of thyroid cancer in children, but not 106 leukaemia [12]. Thyroid cancer patients aged 15 or younger lived in the most contaminated regions 107 (the Provinces of Kiev, Chernigov, Zhitomir, Cherkassy, Rovno, and the city of Kiev). The highest 108 reported incidence was in children who were exposed at the age of five years or younger. [14]. 109 However, because of the lack of proper cancer registries in Ukraine and Belarus, and the large 110 number of people evacuated from the polluted areas that could not be traced through any kind of 111 demographic records, these findings are considered to be of limited significance [15]. Moreover, in 112 some regions of Ukraine and Belarus long-term exposure was accompanied by malnutrition, 113 frequent infections and stress, important confounders that may have had strong impacts on the 114 reported genome damage [16].

A systematic overview of available data of genome damage in children environmentally exposed to ionizing radiation is missing. Data from scientific papers written in Russian, Byelorussian and Ukrainian language are not well known in the Western scientific community due to language barriers. The aim of this study is to present and interpret systematically collected data on genome

damage in children exposed to ionizing radiation at the global level (cosmic radiation is excluded).

120

121 Materials and methods

122 The scientific literature considered here was selected following an extensive literature search 123 without any language restriction by using the Med-Line/PubMed database (National Library of 124 Medicine, National Institutes of Health, Bethesda, MD, USA-http: //www.ncbi.nlm.nih.gov/ 125 PubMed) covering the time period between January 1, 1980 and December 30th, 2006. Searches 126 comprised studies of children from newborns to late adolescence (age 0-18 years) exposed to 127 ionizing radiation. We have excluded case reports, studies without a clear definition of exposure to 128 ionizing radiation, studies with less than 10 children and/or lacking a referent (unexposed) 129 population, and studies reporting findings in a conversational style without statistical measures 130 (e.g., mean and standard deviation) or analysis. Studies written in English, Russian, and Ukrainian 131 were retrieved and manually reviewed. Studies that were not accessible through online library 132 systems were obtained by the authors or through interlibrary exchange. Twenty East European 133 studies of children not available in English but with important information have been recognised 134 and included in this review. Information about these studies can be obtained upon request. Results 135 from the following assays of genetic toxicology were considered: chromosome aberration assay 136 (CA), in vivo and in vitro micronucleus assay (MN), comet assay, sister chromatid exchange (SCE), 137 and fluorescent in situ hybridization (FISH). In order to simplify the presentation and the 138 interpretation of the reviewed studies the association between radiation exposure and biomarkers of 139 DNA damage in children was quantitatively investigated by computing study specific ratios (MRs) 140 of the mean level of each biomarker detected in radiation exposed and in referent children or 141 newborns. The computed MR is a point estimate of the relative effect of the exposure on biomarker 142 level detected in each study taking the value 1 (MR=1) when there is no effect of radiation exposure 143 on biomarkers level, values greater than 1 (MR>1) or lower than 1 (MR<1) when radiation exposure is associated with an increased or a decreased levels of the investigated biomarkers, 144

145 respectively. The MR, as a measure of effect, has the advantage of being independent of the 146 absolute values of the biomarker mean levels reported by the single studies and is comparable 147 across the studies and endpoints considered. The main characteristics of the studies considered in 148 this paper including their findings and the computed MRs, are summarized in Tables 1-3.

149

150 Results

151 Exposure to ionizing radiation from natural sources

Naturally-occurring radionuclides in food and water are primarily potassium (⁴⁰K) and the decay products of Uranium (²³⁸U), Thorium (²³²Th), Carbon (¹⁴C) and Rubidium (⁸⁷Rb) [17]. Radium (²²⁶Ra, decay product of ²³⁸U in nature) decays into radon (²¹⁹Rn, ²²⁰Rn, and the most stable radon isotope ²²²Rn, with half life 3.82 days) which is emitted as a gas in significant quantities and can reach levels in indoor air up to 15,000 Bq/m³. The main intake of ²²²Rn is via drinking and breathing. The European regulations and US environmental action levels are 150-200 Bq/m³ [18, 19, 20,21].

A possible association of radon exposure with adverse health effects, including lung cancer development, has been recognized relatively recently [22, 23]. Critical environments are poorly ventilated old dwellings built in karsts-rich areas, geographical areas of irregular limestone where erosion has produced fissures, caverns and underground streams. Children may be exposed to radon at kindergarten, school and at home. An additional (although less important) source of exposure to radon could also be building material such as certain types of concrete and granite tiles [24].

Rommens et al 2001 [25] reported European ionizing radiation exposure levels of 2.4 mSv/y for adults, 2.7 mSv/y for children and 5,4 mSv/y for infants 0-1 years old taking into account all natural sources such as ²²²Rn and ²²⁰Rn (decay product of thorium, commonly named thoron), cosmic radiation, terrestrial radiation, radionuclides, etc. The total body concentration of radionuclides and equivalent doses to red bone marrow is age dependent and is higher in children, especially in infants and adolescents for ²²⁶Ra, lead (²¹⁰Pb), ²²⁸Th, Polonium (²¹⁰Po), etc [25, 26,27].

171 As radon daughter products follow the metabolic pathway of calcium, its incorporation into 172 children's skeleton poses a significant health risk [28]. Due to age dependent developmental stage 173 of the gastrointestinal system of children, the highest absorption of radon is in newborns and in 174 children between 13 and 17 years of age [27,29]. This is accompanied with high water intake in 175 newborns and children in comparison with adults [30]. Children and adolescents are target 176 populations for intake of water which can be radiocontaminated, due to the increased usage of 177 bottled water in Europe and its use for production of a number of different drinks favoured by the 178 voungest. The increased effective dose from radiocontaminated mineral water may be up to seven 179 times higher in infants and teens than the maximum level recommended by the World Health 180 Organization (100 μ Sv,) [31]. It has been suggested that this exposure may be specifically relevant 181 for the hormonal activity of testosterone and oestrogen during puberty when final maturation of 182 skeleton occurs [32]. Non breast fed infants less than 1 year of age may receive doses up to 0.28 mSv/y if their diet is exclusively prepared with mineral water with elevated radon concentrations 183 from ²²⁶Ra decay [33]. In addition, such waters contain other radionuclides such as ²¹⁰Pb and ²²²Ra 184 185 also contributing to the total received dose [33].

186 The health risk related to indoor radon exposure is still a subject of discussion. It has been shown 187 that residential radon exposure may contribute to increased cancer incidence. The average radon exposure of 50 Bqm⁻³ has been estimated to be responsible for 13-25% of myeloid leukaemia cases 188 189 at all ages [34,35]. Indoor exposure at an annual dose of 7-11 mSv from radon has been reported to 190 be associated with a significantly increased frequency of chromosome aberrations (MR=1.69, Table 191 1) and micronuclei (MR=1.44, Table 2) in children [36]. To decrease radon levels in the working 192 and living environment some countries have established programmes for remediation work in 193 buildings, primarily schools and homes [37].

194

195 Exposure to high-dose ionizing radiation

196 The Chernobyl nuclear accident.

197 After the 1986 Chernobyl nuclear power plant accident, populations of Ukraine, Belarus and Russia were exposed to Iodine (¹³¹I), Caesium (¹³⁷Cs, ¹³⁴Cs), Strontium (⁹⁰Sr) and to a wide spectrum of 198 199 short-lived isotopes which were not measured by physical dosimetry [11]. Later on, exposure 200 became continuous with constant intake of radionuclides via food and water, including ⁹⁰Sr which is 201 incorporated in the skeleton of children at 4-6 fold higher rates than in adults [38]. It has been 202 estimated that following the Chernobyl accident approximately 160,000 children aged 7 years or 203 less were exposed to a variety of radioactive isotopes [39]. The explosion of the Chernobyl-4 204 reactor core led to the release of radioactivity that was deposited in the surrounding area as dust and 205 debris, while the lighter material was carried by wind over the Ukraine, Belarus, Russia and to some 206 extent over Europe, with radioactive fallout in Scandinavia, Austria and Switzerland [17]. Some 15 207 to 23 kg of plutonium were released, the majority within an area of 80 km radius around the nuclear 208 plant [40]. Immediately after the accident, a first zone with more than 40 Ci/km² (the 10 km range 209 zone) and a second one with more than 15 Ci/km² (range of 30 km) were identified. A third area of 210 145,000 km² was contaminated with more than 1 Ci/km². After the accident 135,000 people were 211 evacuated from the first zone and after some time 210,000 more subjects were evacuated. An 212 unidentified number of evacuated subjects were sent to different parts of Russia, Israel or other 213 European countries, for varying periods of time. Today, about 3.8 million people live in the area 214 with more than 1 Ci/km². The effective human annual dose is in the range between 54 μ Sv and 3.1 215 mSv [41, 42]. Efforts to reduce the exposure of the population through altering their diet were not 216 very successful [40]. In affected area increased incidence of thyroid cancer in children [43] and 217 recently breast cancer have been reported [26, 27]. Breast cancer could be expected to follow 218 thyroid cancer since the mammary gland is derived embryogenetically from primitive iodide-219 concentrating ectoderm [26, 30, 44, 45, 46]

220 Chromosome aberrations. Cytogenetic studies of the children population in Ukraine started in 1988

221 [47]. Studies reported in [47] were performed on peripheral lymphocytes and only in vivo MN assay 222 was performed on reticulocytes. Results revealed dose-dependent increased levels of CA, with MRs of 3.22 and 1.98, in children exposed to 137 Cs at levels between 18 and 55 x 10^{10} Bg/km² and lower 223 than 1×10^{10} Bg/km², respectively (Table 1). Repeated measurements of chromosome aberrations 224 225 within a 4 year period after accidental overexposure in children living in contaminated areas [48] 226 revealed a 53% increased average level of genome damage as measured by the chromosome 227 aberration (CA) assay (Table 1). The follow-up of several exposed and evacuated groups of 228 children born before and after the nuclear accident by highly experienced cytogenetic centres such 229 as those in St. Petersburg (Russia) or in Pisa (Italy) also showed the persistence of the genome 230 damage. Anage related radiosensitivity was detected in children from Belarus who were sampled 231 three months after the Chernobyl accident. A significant difference was found in the number of 232 dicentrics between young (6-10 years) and older (11-15 years) children (Table1), with 1.17% and 233 0.67% dicentrics, respectively [49]. Up to 10 years after the accident children were still suffering 234 from internal contamination: CA frequencies were up to 4 times higher in exposed than in reference 235 children (Table 1) [50, 51, 52, 53]. In exposed children the frequency of dicentrics was 0.44% 236 compared to 0.02% observed in unexposed children (Table 1) [54]. Such an alarming situation 237 feeds speculations about an accumulation of stable genome damage in these children and potentially 238 related adverse health effects that may occur later in life. Cytogenetic studies also showed that even 239 the areas which are considered as unpolluted are actually contaminated with radionuclides at levels 240 that are capable of increasing genome damage in children [47]. The impact of internal 241 contamination was seen as a presence of rogue cells (specific type of multiaberrant cells) detected 242 in children living in contaminated areas [55]. In this study 328 Belarussian children were analysed 243 by the CA assay. The majority of the children (321 subjects) were exposed postnatally. In six 244 children exposed in utero one or two rogue cells were detected in 200 analysed metaphases. 245 Detected rogue cells contained up to 9 dicentrics, up to three tricentrics or/and rings and 246 quadricentrics.

247 In utero exposure. The Chernobyl accident affected also pregnant women who were exposed to 248 different levels of radiation before being evacuated. A study using G banding was performed on two 249 groups of children exposed in utero and during childhood to ionizing radiation [56]. Children born 250 by mothers who were pregnant at the time of the accident and evacuated shortly afterwards were 251 exposed to radiation levels ranging between 10 and 376 mSv while children exposed in utero and 252 chronically during the childhood experienced a cumulative dose of 19-52 mSv. An increased 253 frequency of CA was detected in newborns from both groups of women. As shown in Table 1 mean 254 frequencies of 9.07% ±1.34 and 7.63%±2.92 of CA were measured in the group of intrauterine 255 exposed children and in children exposed in utero and after birth, respectively, compared to a 256 frequency of 2.47%±0.4 detected in referent children (MR= 3.67 and 3.08). Translocations, 257 inversions and deletions represented almost 80% and 70% of chromosome type aberrations in 258 intrauterine exposed and continuously exposed children, respectively [56]. Non random distribution 259 of chromosome damage was detected: the most frequently involved chromosomes were 260 chromosome 1, 3, 5, 7, 9, 11, 13, 21 and 22 [56]. On the background of the available scientific 261 evidence, this predominant localization of break points correlates with diagnostics markers of 262 neoplastic disease as summarized in Table 4 [57]. All detected bands at which chromosome 263 breakage was present are non-random and related with the described neoplasias.

264 Parental exposure. A high frequency of aberrant cells (1.12± 0.37%) was measured in children 265 exposed in utero to 2.0-2.5 cSv, in those born between 1987-1991 (1.24±0.4) as well as in children 266 born between 1994-1998 (1±0.2) compared to children born before the Chernobyl accident (0.59± 267 0.3%), with MRs of 1.90, 2.1, and 1.69, respectively [58]. A long-term follow-up study of 268 populations living in a contaminated area of 15, Ci/km² [58] showed increased genome damage in 269 children of irradiated parents (Table 1). The highest frequency of aberrant cells $(1.24 \pm 0.4\%)$ was 270 measured in children born by mothers who were continuously exposed to ionizing radiation 271 following the Chernobyl accident (MR=2.1).

272 The paternal transferability of possible genome damage has been investigated in children born after

their fathers were exposed as liquidators at the Chernobyl nuclear plant. Genome damage was measured in 15 children born after evacuation using the CA assay [59]. A clear increase in CA (MR=1.64, Table 1) was detected in children of Chernobyl liquidators who suffered with radiation burns of 1st and 2nd degree (2.38% \pm 1.9) compared to referents (1.45% \pm 0.2).

277 Six years after the accident children who were evacuated at different times following the accident 278 and children born after cessation of their father's exposure were analysed for CA. Evacuated 279 children had spent between 2 days and 2 years in contaminated areas. As it is shown in Table 1, 280 clearly increased frequencies of CA were detected in children of exposed liquidators $(2.8\%\pm0.2)$ 281 and in evacuated children $(2.5\% \pm 0.1)$ compared to referents $(1.8\% \pm 0.2)$. Noteworthy, evacuated 282 children still had almost ten times more dicentric and ring chromosomes than controls (0.19% and 283 0.02%, respectively, data not shown). Such increased values of these types of CA could reflect 284 genomic instability, a phenomenon of increased rate of acquisition of alterations in the mammalian 285 genome proposed to be a driving force in carcinogenesis [53]. Indeed, germline mutation 286 frequencies at human minisatellite loci among children born in polluted area and receiving doses of 287 about 0.18 Gy were shown [60] to be two times higher when compared with a control population 288 (mutation rate per band 0.03 versus 0.01, respectively). The measurement of new fragments using 289 multi-site DNA fingerprinting showed that liquidators' children born after the Chernobyl accident 290 had a seven-fold increased level of new bands that were not present in their sibs conceived before 291 the Chernobyl accident [61, 62].

Micronucleous Assay, Comet Assay, FISH, and SCE. Among the *in vitro* studies conducted on children exposed following the Chernobyl accident (Table 2), the one by Mikhalevich et al 2000 [49] failed to detect a difference in the frequency of MN in binucleated lymphocytes of those living in contaminated areas for 9 years after the accident and being chronically irradiated by internal contamination, compared to referents (MR=0.83). The study reported a clearly increased frequency of MN in mononucleated lymphocytes (MR=2.48) in chronically irradiated children (Table 2). Two studies [63,64] reported a twofold increased frequency of micronucleated cells in radiation exposed

compared to referent children while another [65] detected similar levels of micronuclei in exposed and referents (Table 2). Using the *in vivo* micronucleus assay, liquidators from Chernobyl and their children evacuated following the nuclear accident [66] were observed to express significantly increased mean micronuclei levels in peripheral lymphocytes compared to referent children (0.19% and 0.012% micronucleated cells, respectively, MR=15.8). This is the only available study on a population environmentally exposed to radiation monitored by the *in vivo* MN assay.

305 The Comet assay has been used to estimate genome damage levels in children from Belarus 10 306 years after the Chernobyl accident [67]. An increased genome damage was still present in their 307 lymphocytes (Table 3), a finding that could be explained as a "clastogenic factor" present in 19% of 308 these children [67]. When the translocation frequency was measured using FISH (Table 3) in a 309 group of exposed children and in age matched referents, higher levels of translocation were found in 310 the former $(0.65\% \pm 0.1)$ than in the latter group $(0.14\% \pm 0.05)$ corresponding to an MR of 4.64 [68]. 311 In the same study higher MN frequencies were found in exposed children than in exposed adults 312 (0.06%), a finding that is suggestive of a higher sensitivity of children to ionising radiation induced 313 cytogenetic damage, since it seems unlikely that the children were exposed to higher levels of 314 radiation than their parents.

315 Antioxidants and lipid peroxidation.

316 Several studies (not included in any table) have investigated the association between radiation 317 exposure, lipid peroxidation disorders, and cytogenetic damage. Analyses of children born by 318 mothers who were exposed to low doses of radiation before pregnancy showed that in regions 319 contaminated by radionuclides, these children suffered from lipoperoxidase disorders and that the 320 levels of essential antioxidants such as vitamin A and E were low. For mothers on a diet 321 supplemented with these vitamins during pregnancy, the chromosome aberration frequency in their 322 newborns was significantly lower in comparison with children born by mothers without such 323 vitamin supplemented diet [69]. The complexity of interaction between the organism in 324 development and radiation is also illustrated by a bimodal pattern of distribution of the gluthatione

325 system [70, 71] in children born by mothers exposed to different dose levels. In children born by 326 mothers exposed to doses between 0.8 and 30 cSv, increased levels of reduced plasma glutathione 327 (up to 90 μ M) could be detected, while in mothers exposed to doses between 30 cSv and 60 cSv, 328 severe decrease (5 µM) was detected. Gluthatione mediates a reduction of at least two vitamins, 329 alfa-tocopherol and ascorbic acid which are critical in prevention of lipid peroxidation. Additional 330 consequences of exposure to ionizing radiation and disturbances of glutathione level are seen in 331 cases of combined radiochemical exposure. Increased levels of polycyclic aromatic carbons (PAH)-332 DNA adducts are found in human placenta of mothers exposed to both ionizing radiation and 333 environmental PAH, suggesting a possibly higher health risk for the foetus in a case of complex 334 exposure than would be expected exclusively from PAH concentrations in air [72].

335 Radiation induced adaptive response.

336 An adaptive response of subjects exposed to low doses of chemical agents or radiation has 337 frequently been investigated by Russian and Ukrainian scientists. The existence of radiation 338 associated hormesis (i.e. the concept that small doses of radiation may reduce the damage to levels 339 even lower than those observed in unexposed controls) is still debated in the literature. The adaptive 340 response is a characteristic feature of both mammalian and plant cells in their response to various 341 mutagenic agents [73]. This phenomenon occurs when cells are treated with a low dose of a 342 clastogen; such a pre-treatment may then reduce the effect of a subsequent treatment with a higher 343 dose of the same or a similar agent (the challenging treatment). Adaptation can be measured by the 344 challenge assay in which lymphocytes isolated from exposed and control subjects are treated with 345 either 1 Gy or a combination of 0.05 Gy and 1 Gy. Compared to children living in Moscow's urban 346 polluted areas, increased micronuclei frequencies were reported among children from the Chernobyl 347 area exposed to radiation by living in areas with contamination levels ranging between 5 and 40 348 Ci/km². Lymphocytes of children from these radiocontaminated areas showed increased frequencies 349 of chromosome aberrations and micronuclei in the challenge assay, suggesting that, in this exposed 350 population, subjects express a radiosensitivity with no indication of an adaptive response. These

351 results may show that a routine application of chromosome aberration and/or micronucleus assay 352 without also employing a challenge assay may not be sufficient for the detection of genomic 353 instability [63]. By comparing this radiosensitivity with the observations of adaptive response in the adult and children populations from Ural (internal exposure with ⁹⁰Sr from the Techa River) and the 354 355 Chernobyl regions, some common features were apparent. In both groups of exposed children, 356 individuals with high radiosensitivity were recognized after a challenging dose of 1 Gy and using 357 MN as the endpoint. Among referent children from the Moscow region, inter-individual differences 358 in MN frequencies were significantly lower after challenge assay than in exposed group. Based on 359 such sada It is suggested that adaptive response is not developed and will not be expressed in 360 adulthood if a person is exposed as a child to elevated levels of radiation or other xenobiotics [74, 361 75].

362 Hiroshima and Nagasaki

363 The Chernobyl disaster resulted primarily in radiation via internal contamination, whereas after the 364 Hiroshima and Nagasaki bombings the primary source was gamma radiation and fast neutrons. This 365 represents different types of exposures, which are also associated with different cancer incidence 366 distributions [76]. Two to three years after the bombing, leukemia was the first cancer to be linked 367 with exposure with the highest incidence of leukemia detected in people exposed during their 368 childhood [77]. Besides the increase in leukemia, elevated rates of solid cancers such as cancers of 369 the breast, lung, and colon have been reported which seem to be larger among subjects exposed 370 during their childhood than among those exposed as adults [78].

371 Cytogenetic analyses of exposed populations started 15 years after the bombing [79] due to the fact
372 that cytogenetic methods were not introduced until the 60's. A number of different research teams
373 have been involved in studying health consequences in survivors and studies are still on going
374 among children of exposed subjects. [80]

375 Nuclear plants

Epidemiological studies have reported clusters and increased risks of leukaemia in subjects living in areas adjacent to nuclear power plants [81, 82, 83, 84, 85, 86, 87, 88]. A significant excess of leukaemia cases in the general population living in the proximity of nuclear plants has also been reported [82, 89, 90] along with a possible association for children whose father were employed at a nuclear plant [90].

A systematic review of leukemia incidence and mortality cohort studies in children living in the proximity of nuclear facilities found that the majority of studies reported elevated rates [91]. This meta-analysis confirmed an increased risk of childhood leukemia near nuclear facilities with the highest contribution of excess cases and deaths from children aged < 10 years and living within 15 km from the nuclear site.

386 A few studies using cytogenetic methods have been conducted in Europe with children living close 387 to nuclear plants or downstream of rivers from which water is used for cooling of reactors [90, 92]. 388 Between 1988 and 1995 in vitro chromosome aberration assays were performed with peripheral 389 lymphocytes from 5 healthy siblings of the leukemia cases and in 10 control children from the 390 Elbmarsch Municipality [93 ??], 42 children from the Elbmarsch Municipality and 30 children from 391 a control region [92], and 25 adults including 7 parents of children diagnosed with leukemia and 14 392 inhabitants near the Krummel nuclear plant and in 25 healthy adults (control subjects) living in the 393 city of Bremen about 100 km southwest of Hamburg [94]. While two studies [93, 94] reported a 394 significant fourfold higher rate of dicentric and/or ring chromosomes in peripheral blood 395 lymphocytes in subjects leaving near the plant compared to control subjects, one study [92] failed to 396 detect any such difference between exposed and control children.

397 Nuclear weapons fallouts, and nuclear accidents

398 During the period 1948-1967 three nuclear accidents took place in the Soviet nuclear weapon
399 industry known as «Mayak Complex» established on the Techa River near Chelyabinsk, in the
400 Southern Urals. During this period beta/gamma radioisotopes were released into the Techa River.

Comment [dfm1]: I've moved refs for krummel here to get rid of reiterations seen by Gunnar

401 Several studies have been conducted on the population of 280,000 dwellings living in the area [95]. 402 It has been estimated that about 8% of this population (124,000 people) were exposed to radioactive isotopes such as ¹³⁷Cs, ¹⁰⁶Ru, ⁹⁵Zr, ⁸⁹Sr, ⁹⁰Sr, and that these received total accumulated doses above 403 404 1.0 Sv [96]. The increased frequency of CA that was observed in exposed children from this area 405 (0.56±0.08) compared to referents (0.29±0.07), resulting in an MR= 1.93 (Table 1), was 406 accompanied by a 1.7-fold increase in minisatellite mutation rate in the germline of exposed fathers 407 than in referents from rural areas (data not included in Table 1) [97]. Contrary to Hiroshima, 408 Nagasaki, and Chernobyl, the general population was exposed to short-lived radionuclides through 409 food, water and air in the period 1949-1963 due to the nuclear tests at the Semipalatinsk, 410 Kazakhstan (former Soviet Union), just south of the Altai region of Siberia. An impact of radiation 411 exposure during childhood was suggested by studies of adults living in this region. The area was 412 contaminated from 450 nuclear tests until 1989 of which 100 were atmospheric [98,99]. Cytogenetic 413 analysis of two or three generations within a family showed a presence of dicentric and ring 414 chromosomes in children born after the nuclear tests had ceased, suggesting family genomic 415 instability [100]. At the time of atmospheric bomb tests in 1949, about 25,000 people received doses 416 between 10 mSv and 1,500 mSv [96]. Studies of this population detected germ line mutations in 417 adults born between 1926 and 1960 and exposed to nuclear fall-out between 1949 and 1956, during 418 their childhood [101, 102].

419 There are special living conditions in Western Siberia where, in a large geographical area, mines, 420 chemical and nuclear industries are located. A follow-up study of 289 children [103] showed 421 significant deviations (including multi-aberrant cells) even in the "referent" population. Indeed, 422 because of the level of radiation pollution in this region, no unexposed children could be included in 423 the study and children in the lowest exposure group were used as referents [103, 104]. Despite the 424 acknowledged exposure misclassification, increased CA levels were reported for children exposed 425 to 0.6 cSv compared to those exposed to 0.05 cSv (3.94 ± 0.44) and 2.58 ± 0.59 , respectively; 426 MR=1.53 Table 1).

427 The nuclear test site Syevernaya (in the period between 1955 and 1962, about 80 nuclear bomb tests 428 were performed here) is located in Nóvaya Zemlyá, an area of Russia in which the population of the 429 Tundra Nenets population lives. In this population increased cancer morbidity was detected (100). 430 Similar to the children population in the Altai region, 20 years after cessation of nuclear tests, 431 unstable chromosome aberrations such as rings and dicentrics were still detected in newborns [105]. 432 An increased MN frequency (MR=1.4 and MR=1.6 in probably exposed and directly exposed 433 children compared to referents) was detected in accidentally exposed children in the Goiania, Brazil 434 radiological accident (Table 2); detected values are within the control values of other studies 435 presented in Table 1 [106].

436 Contaminated building materials

437 In the period between 1982 and 1984 in Taiwan a number of buildings were constructed using 438 reinforced steel contaminated by ⁶⁰Co which had been illegally discarded. The buildings were 439 residential, schools and kindergartens, and it was estimated that 20% of 897 families were exposed 440 to radiation doses higher then 5 mSv/year (in addition to background radiation). Several studies 441 were carried out to estimate the radiation-induced genome damage simultaneously with physical 442 dosimetry [108, 109, 110]. Although 1,500 subjects aged between 0 and 19 years were studied, 443 findings for children were not reported separately [109], with the exception of a small study without 444 a referent population in which the first blood sampling showed a significantly increased CA 445 frequency (20.6±3.9%) in comparison with a second sampling conducted after evacuation from the 446 contaminated buildings (8.7±1.5%), resulting in an MR=2.4 (Table 1). The issue of contaminated 447 steel or other metals is not limited to the Taiwan episode: several reports [111, 112, 113] have 448 addressed unsolved problems concerning building materials used in construction.

449

450 Conclusions

451 In general, genome damage caused by accidental overexposure to ionizing radiation may result452 from interactions such as the formation of DNA damage directly or via free radicals, but also from

453 damage to the nuclear membrane, lipid peroxidation, methylation disturbances, activation of a chain 454 of signal molecules influencing the expression of apoptosis, and other mechanisms including 455 hormonal, age related bioaccumulation of radionuclides, metabolism and clearance. Other 456 contributing factors such as stress, malnutrition and infections may play major roles. At present, 457 such conditions are predominantly found in some specific social environments in Ukraine and 458 Belarus, but they could easily be envisaged in other parts of the world in conjunction with a nuclear 459 accident. With respect to the genome damage discussed in this review, the main body of available 460 data for children exposed to radiation comes from studies that were performed in Russia. Territories 461 of the ex-Soviet Countries, as well as some other areas of our planet are still polluted as a 462 consequence of nuclear tests and the nuclear industry. The resulting nuclear waste may lead to 463 radiation exposure and complex exposures to radiation and chemical pollution, as is the case in a 464 rocket test area or in spacedroms [114].

465 The studies considered in this systematic review consistently reported an increased frequency of CA 466 and micronuclei in radiation exposed children compared to referents. Elevated CA levels were 467 observed also in children exposed to high levels of radiation when compared to those exposed to 468 lower levels [47,54,49,103]; this interpretation should be made despite the potential exposure 469 misclassification of referent children for whom radiation exposure levels are hardly ever reported. 470 Such a differential misclassification of exposure is expected to result in study findings that are 471 toward the study null hypothesis (i.e., that there is no effect of environmental radiation exposure on 472 genome damage in children) resulting in underestimated measures effect (MR).

The impact of internal contamination was indicated by a presence of rogue cells and by the higherfrequency of dicentrics in exposed than in referent children as reported by several studies.

Beside a threat of accidental overexposures children are exposed routinely to ionizing radiation for
diagnostic purposes which doses should be recorded and summarised as a cumulative life long dose.
An increased risk of childhood leukaemia was detected in children who reported two or more
postnatal X-rays [115]. The environmental burden during childhood could have a significant

479 influence on the adaptive capacity in adulthood and could be partly responsible for inter-individual
480 differences in chemo- and radiosensitivity [116]. Additional radiosensitivity time windows should
481 be investigated which may exist during childhood, e.g. in newborns and teens.

482 Populations suffering from radiological accidents, living in contaminated areas or ex-nuclear testing 483 sites or close to radiochemical industries, are today subjected to an improved identification and 484 monitoring, taking place in several medical centres. These groups of people constitute a large cohort 485 including thousands of subjects and sometimes families with several generations. Local research 486 centres often have stored biological samples, data on exposure and questionnaire data representing a 487 valuable source of information that could be used for studying the delayed adverse effects of 488 radiation on children and the transgenerational transfer of genome damage [96]. Integration and 489 utilisation of such sources represent a great challenge and will probably be important in future 490 studies of the consequences of children's exposure to ionizing radiation.

491 In Europe there is a dramatic discrepancy between the number of existing nuclear plants and the 492 number of field studies on the potentially associated health consequences [117]. Regarding natural 493 sources of radiation, consumers should be informed about possible health risks associated with the 494 consumption of bottled mineral waters available on the market [6], in cases where increased 495 radiation exposure may be significant. As only 5%-15% of childhood cancers seems to be related 496 with familial and genetic factors [118]. the age related sensitivity should be subject to a closer study 497 in order to avoid, at least, exposure of children during their most susceptible age periods. Similarly, 498 governments should support reconstruction of dwellings in order to decrease radiation levels in 499 homes and public buildings where children spend most of their time. Strict control of building 500 materials for radio contamination should be performed (such as fly ash and steel). 501 Genotoxicological studies of children living in the proximity of nuclear plants should be set as a 502 priority in research programmes. At the same time there is a need for an arising preparedness of 503 medical staff in case of nuclear accidents.

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Table 1. Chromosome aberration frequencies (mean \pm SE), measured in children exposed to ionizing radiation and in referents, by type and level of exposure. The exposure level, sample size, age of the study groups (range), and the median ratio (MR) are reported for each study. Gaps excluded.

Type of exposure	Level of exposure ^a	Group: Sample Size	Age ^b	Mean ± SE (%)	MR	Details	Reference
¹³⁷ Cs Chernobyl	18-55 x10 ¹⁰ Bq/km ² 0.37-0.74 x10 ¹⁰ Bq/km ² 	Exposed: 103 Exposed: 27 Referent: 16	6-15	2.74±0.1 1.68±0,2 0.85±0.1	3.22 1.98 1		42 Yeliseeva et al.,1994
Radionuclides Chernobyl	74-148 Ci/km ²	Exposed: 11 Referents: 13	4-11 4-11	4.0±0.3 2.6±0.2	1.53 1	Repeated measurements	43 Bochkov et al.,1991
Radionuclides Chernobyl	148 x10 ¹⁰ Bq/km ²	Exposed: 24 Referent: 11	10-12 10-12	0.62±0.08 0.33±0.08	1.87 1		46 Padovani et al.,1997
Radionuclides Chernobyl	40 x10 ¹⁰ Bq/km ²	Exposed: 24 Referent: 10	8-10 8-10	2.4 0.6	4.00 1	¹³⁷ Cs whole-body counter	45 Padovani et al.,1993
¹³⁷ Cs Chernobyl	2.43 Bq/day 	Exposed: 17 Referents: 35 Exposed: 17 Referents: 35	9-14 12-16 9-14 12-16	1.46±na 1.42±na 1.3±na 0.4±na	1.02 1 3.25 1	Belarus children Italian healthy children % Dicentrics % Dicentrics	44 Barale et al.,1998
Radionuclides Chernobyl		Exposed: 49 Exposed: 35 Referent: 25	1-16 1-6 3-16	2.5±0.1 2.8±0.2 1.8±0,2	1.38 1.55 1	Evaquated children Liquidators'children	47 Vorobtsova al.,1995
Radionuclides Chernobyl	0-6 cSv/y 0.8 cSv/y 	Exposed: 25 Exposed: 25 Exposed: 25 Referent:25	6-15 6-15 6-15 6-15	3.93±0.2 3.78±0.3 2.62±0.2 1.74±0.2	2.25 2.17 1.50 1	Dicentrics = 0.44%	48 Pilinskaya et al.,1992

Table 1 (cont.). Chromosome aberration frequencies (mean \pm SE), measured in children exposed to ionizing radiation and in referents, by type and level of exposure. The exposure level, sample size, average age of the study groups, and the median ratio (MR) are reported for each study.

Type of exposure	Level of exposure ^a	Group: Sample Size	Age ^b	Mean±SE (%)	MR	Details	Reference
Radionuclides		Exposed: 15	2-5	2.38±1.9	1.64	Fathers with 1 st - 2 nd degree burns	51 Stepa. 1993
Chernobyl		Referents: 50	2-5	1.45±0.2	1		
Padianuclidas	2 0 2 5 cSv	Exposed: 14	15	1 12+0 37	1.00	Exposed in utero	50 Suckov 2001
Charmobyl	2.0-2.5 cSv	Exposed: 18	10 14	1.12 ± 0.57 1 24 ±0.4	2 10	Born 1087 1001	50 Suskov,2001
chernobyi	1.5-2.5 cSv	Exposed: 70	3 7	1.24 ± 0.4	2.10	Born 100/ 1008	
cumuutive uose		Referent: 15	15-19	0.59 ± 0.3	1.09	Born before 1986	
N I ' I' I	10.276	E	15	0.07+1.24	2 (7	Democratic stars are set of	(7 Store 2002
Radionuclides	10-376 mSV	Exposed: 22	15	9.07 ± 1.34	3.07	Exposed in utero evaquated	67 Stepa.,2002
Chernobyl	19-52 mSv	Exposed: 20	15	7.63±2.92	3.08	Exposed in utero and after birth	
		Referent: 15	15	2.47±0.4	1	Living in unpolluted areas	
¹³⁷ Cs, ⁹⁰ Sr	35.9 kBg/m^2	Exposed: 20	6-10	1.17± na	1.75	% Dicentrics	71 Mikhalevich ,2000,
Chernobyl	2.22 kBq/m^2	Referents: 10	11-15	0.67± na	1	% Dicentrics	, ,
Nuclear industry	> 1 Sv	Exposed: 15	9-11	0.56±0.08	1.93	Exposed to long-lived radionuclides	82 Testa.1998
Southern Urals		Referent: 11	9-11	0.29 ± 0.07	1		
		Exposed: 15	9-11	0.07+0.03	1.4	% dicentrics	
		Referent: 11	9-11	0.05±0.02	1	% dicentrics	
Nuclear Industry	0.6 cSv	Exposed: 289	12-17	3 94+0 44	1 53	children with higher exposure	90 Druz 1997
Western Siberia	0.05 cSv	Referent: 12	12-17	2.58±0.59	1	children with the lowest exposure	50 Diul.,1997
Nuclear nower nlant		Exposed: 42	9-17	$0.43(0.240.7)^{c}$	0.61	% Dicentrics+ring chromosomes	81Bruske-Hoh 2001
Elbmarsch, Germany		Referent: 30	9-17	$0.706 (0.4, 0.1)^{c}$	1	% Dicentrics+ring chromosomes	orbruske from, 2001
⁶⁰ Co (stool robor)	5 m S w/w	Exposed: 18	1 18	20 6+3 0	24	1 st phlebotomy	07 Heigh 2002
Co (steel lebal)	5 mSv/y	Exposed 18	4-10	20.0 ± 3.9 9 7 1 1 5	2.4	2 nd phlabatomy ofter avaguation	97 HSIEII ,2002
Taiwan, residentiai	5 IIISV/y	Exposed.18	4-10	0.7±1.3	1	2 phieodolony, after evacuation	
Ra (indoor)	$>1000 \text{ Bq/m}^{3}$	Exposed: 85	8-12	2.03±3.9	1.69	Schools' level	35 Bilban et al.,2001
Slovenia	$<400 \text{ Bq/m}^{3}$	Referents: 20	8-12	1.2±0.59	1	Schools' level	
		Exposed: 85	8-12	0.08±0.44		% dicentrics	
		Referents: 20	8-12	0	1	% dicentrics	

^a) -- = exposure levels not reported; ^b) range in years; ^c) 95% confidence interval; SE = standard error; na = not available

Table 2. In vitro (binucleated and mononucleated lymphocytes) and in vivo (reticulocytes) micronucleus assay data) measured in children exposed to ionizing radiation and in referents, by type and level of exposure. The exposure level, sample size, average age of the study groups, and the median ratio (MR) are reported for each study.

Type of exposure	Level of exposure ^a	Group : Sample Size	Age ^b	Mean ± SE (%)	MR	Comments	Reference
Ra (indoor)	>1000 Bq/m ³	Exposed: 85	8-12	6.5±2.5	1.44	Schools' level	35 Bilban et al.,2001
Slovenia	<400 Bq/m ³	Referenst: 20	8-12	4.5±1.9	1	Schools' level	
¹³⁷ Cs		Exposed: 24	1-18	1.16±na	1.6	Directly exposed	95 da Cruz et al., 94
Accidental, Brazil		Exposed: 14	1-18	1.00±na	1.4	Probably exposed	,
, -		Referents: 30	1-18	0.73	1	J 1	
Radionuclides							
Chernobyl							
¹³⁷ Cs	35.9 kBq/m^2	Exposed: 20	10-17	0.75 ± 0.08	0.83	binucleated lymphocytes	71 Mikhalevich ,2000
⁹⁰ Sr	2.22 kBq/m^2	Referents: 10	10-15	0.90 ± 0.08	1	binucleated lymphocytes	
¹³⁷ Cs	35.9 kBq/m^2	Exposed: 20	10-17	2.71±0.27 ^a	2.48	mononucleated lymphocytes	71 Mikhalevich ,2000
⁹⁰ Sr	2.22 kBq/m^2	Referents: 10	10-15	1.09±0.16 ^a	1	mononucleated lymphocytes	1
Radionuclides	2.43 Bq/day	Exposed: 26	9-14	3.61±2.6	1.80	Belarus children	73 Zotti-Martelli.,99
Chernobyl		Referents: 30	12-16	2.0±2.1	1	Italian healthy children	
Radionuclides	0-20.2 Bq/kg	Exposed: 25	8-9	0.46±0.3	0.90	USA immigrants	72 Livingston et al,97
Chernobyl		Referents: 31	4	0.51±0.2	1	C	
Radionuclides	5-40 Ci/km ²	Exposed: 58	7-13	2.3 ±0.1	2.3		63 Pelevina et al.,96
Chernobyl		Referents: 136	8-13	1.0±0.06	1		,
Radionuclides		Exposed: 54	6-16	0.19±na ^c	15.8	Evacuated children	70 Fedoretsova et.,97
Chernobyl		Referent: 94	6-16	0.012±na ^c	1		,

--= exposure levels not reported; ^a) mononucleated lymphocytes; ^b) range in years; ^c) in vivo micronucleous assay; na, not available.

Table 3. Comet assay, FISH, and SCE results from measurements in children exposed to ionizing radiation and in referents, by type and level of exposure. The exposure level, sample size, the age range, and the median ratio (MR) are reported for each study.

Type of exposure	Level of exposure ^a	Group: Sample Size	Age	Mean±SE	Unit	MR	Details	Reference
				COMET	DNA migration um			
Radionuclides	$24 \times 10^{10} \text{ Ba/km2}$	Exposed: 16	8-16	26.34 ± 9.55		0.62	10 years after explosion	59 Frenzilli2001
Chernobyl		Referents: 39		16.46±4.6		1		.,
				FISH	%			
Radionuclides		Exposed: 11	8-19	0.65 ± 0.1		4.64		69 Vorobtsova,2000
Chernobyl		Referents: 14	3-19	0.14±0.05		1		
				SCE	%			
Radionuclides	2.43 Bq/day	Exposed: 15	9-13	6.0±1.12		0.85	Belarus children	44 Barale et al.,1998
Chernobyl		Referents: 32	12-14	7.0±1.24		1	Italian healthy children	

a) -- = exposure levels not reported

Table 4. Correlation of detected break points (Stepanova 2002) and related disease in intrauterine exposed children.

LOCALIZATION	RELATED DISEASE	LOCALIZATION	RELATED DISEASE
OF BREAK SITES		OF BREAK SITES	
1p35	NB, PV	5q33	ATL, LI, AML, MDS
1p13	AML-M7, PV	5q35	NHL, AML, MDS
1q32	NHL/CLD, MPD	7p15	AML, MPD
1q42	NHL/CLD, MPD	7q33	ALL, AML, CLD, MDS, NHL, PV
2q33	CLL	9q34	ALL, AML, CML, MDS, MPD, NHL
3p21	PA, AC salivary gland, AC/SCC, lung, CLD, NHL, AC kidney	13q32	LI, ALL, MDS, MPD
3q25	AML, MDS, AC kidney	17q21	AML, breast cancer
5p15	ATL, AML	17q25	AML
5p13	ATL	22q13	AML-M7, ALL, MN
5q31	ALL, AML, MDS	-	

*** AC Adenocarcinoma, ALL Acute lymphoblastic leukemia, AML Acute myeloid leukemia, subclasified FAB M1 trough M7, ATL Adult T-cell leukemia/lymphoma, CLD Chronic lymphoproliferative disorder, CLL Chronic lymphocytic leukemia, LI Lipoma, MDS Myelodysplastic syndrome, MPD Myeloproliferative disorder, NB Neuroblastoma, NHL Non-Hodgkin's lymphoma, subclassified B or T cell lineage, PA Pleomorphic adenoma, PV Polycythemia vera, SCC Small cell carcinoma.