Chronic restraint-induced stress has little modifying effect on radiation hematopoietic toxicity in mice

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ABSTRACT

Both radiation and stresses cause detrimental effects on humans. Besides possible health effects resulting directly from radiation exposure, the nuclear plant accident is a cause of social psychological stresses. A recent study showed that chronic restraint-induced stresses (CRIS) attenuated\(^{\text{Trp53}}\) functions and increased carcinogenesis susceptibility of\(^{\text{Trp53}}\)-heterozygous mice to total-body X-irradiation (TBXI), having a big impact on the academic world and a sensational effect on the public, especially the residents living in radioactively contaminated areas. It is important to investigate the possible modification effects from CRIS on radiation-induced health consequences in\(^{\text{Trp53}}\) wild-type (\(^{\text{Trp53}}\)wt) animals. Prior to a carcinogenesis study, effects of TBXI on the hematopoietic system under CRIS were investigated in terms of hematological abnormality in the peripheral blood and residual damage in the bone marrow erythrocytes using a mouse restraint model. Five-week-old male \(^{\text{Trp53}}\)wt C57BL/6J mice were restrained 6 h per day for 28 consecutive days, and TBXI (4 Gy) was given on the 8th day. Results showed that CRIS alone induced a marked decrease in the red blood cell (RBC) and the white blood cell (WBC) count, while TBXI caused significantly lower counts of RBCs, WBCs and blood platelets, and a lower concentration of hemoglobin regardless of CRIS. CRIS alone did not show any significant effect on erythrocyte proliferation and on induction of micronucleated erythrocytes, whereas TBXI markedly inhibited erythrocyte proliferation and induced a significant increase in the incidences of micronucleated erythrocytes, regardless of CRIS. These findings suggest that CRIS does not have a significant impact on radiation-induced detrimental effects on the hematopoietic system in\(^{\text{Trp53}}\)wt mice.

KEYWORDS: chronic restraint-induced stress, total-body irradiation, peripheral blood hemogram, bone marrow micronucleated erythrocytes, mouse restraint model

INTRODUCTION

Nuclear power plant accidents (NPPAs) often result in the release of many different radioisotopes, leading to radioactive contamination. Ionizing radiation (IR) can induce deleterious effects, such as carcinogenicity, mutagenicity, teratogenicity and organ system toxicity. For the public, disasters involving radiation are thought to be particularly pernicious due to the fact that exposure is invisible and universally dreaded. Exposure to IR due to NPPAs is a significant threat and a major health concern. On the other hand, NPPAs cause social psychological stresses. In addition, radioactive contamination often restricts people’s outdoor activities, causing further physiological and psychological stresses (PPSs) [1–6]. Thus NPPAs may pose a long-term
threat to health, resulting in directly and indirectly adverse health outcomes.

Humans are exposed to a multitude of PPSs that contribute to varied adverse health consequences, such as vision disorders, hypertension, cardiovascular disease, diabetes, metabolic syndrome, reproductive disorders and development of cancer [7–13]. Recent studies show that for children, growing up in disadvantaged social environments (i.e. poverty or unstable family) was associated with adverse health outcomes [14]. Genetics moderated the magnitude of the health consequence, but stresses determined the direction [15]. In animals, stresses generally affected cytokine, endocrine and stress hormone (i.e. corticosterone) levels, and immune parameters [16–19]. Chronic restraint-induced stresses (CRISs) significantly impacted upon the hypothalamo–pituitary–adrenocortical axis, causing apical dendritic atrophy [20, 21]. Prenatal exposure to maternal stress altered physiological and immune functions in the offspring [22–24].

Stress increased susceptibility to a number of diseases, including Alzheimer’s disease, and was a risk factor for cancer in humans [9, 25]. In mouse models, stresses altered responsiveness to carcinogens, accelerated tumor onset, and altered tumor type and location [26]. CRIS promoted progression of lymphomas [27] and growth of carcinoma xenografts [28]. Behavioral stress accelerated prostate cancer development [29]. Notably, a recent study showed that CRIS increased susceptibility of Trp53-heterozygous mice to radiation (4 Gy) carcinogenesis, having a big impact on the academic world and a sensational effect on the public [30]. Although mechanisms remain largely elusive, altered metabolism, and degraded physiological and immune functions were critical for increasing susceptibility to pathogens and toxicological assaults, including IR [31–37].

Psychosocial consequences of disasters have been studied for more than 100 years; however, investigations after NPPAs are neither complete nor comprehensive [38], and the importance of the psychological impact is underscored [39, 40]. In fact, following large-scale disasters (i.e. the Chernobyl and Fukushima accidents), psychosocial sequelae were intense and long lasting, and occurred independently of the actual exposure received, and mental health effects were the most significant health consequence [1–6]. It is noted that the evacuate mothers rated their evacuated children’s wellbeing as significantly worse, and the most important risk factors of this health consequence were maternal somatization and Chernobyl-related stress [41]. This work highlights the stress effects on health at a young age. However, such studies are still rare, and the documented works have limited information [5].

Effects from stress exposure on radiation-induced health consequences and the mechanisms underlying these outcomes remain largely unknown, constraining our capacity to ascertain the potential human relevance of the health effects observed in animal models. In a series of investigations, possible modification effects from CRIS on the biological responses of and subsequent consequences for young mice with normal genotype exposed to total-body X-irradiation (TBXI) were studied with multidisciplinary analyses. Measurements included changes of body weight gain and weight of immune organs (such as thymus and spleen), alterations in the levels of blood cytokines and stress hormones, changes in the peripheral blood hemogram and the anti-oxidative activity of blood cells, chromosome aberrations in splenocytes and micronuclei in bone marrow erythrocytes, and epigenetic variations (DNA methylation and miRNA expression) and protein expression profiles in the liver. This paper describes investigation of the possible modification effect from CRIS on TBXI-induced hematopoietic toxicity.

It should be pointed out that this work did not simulate the residents in the contaminated environment, who were exposed to a very low dose rate at a very low dose. Although the hematopoietic system is highly sensitive to radiation, it is not clear that if the very low dose received by the residents in the contaminated environment could cause any detectable effect on the hematopoietic system in humans or in mice. A dose of 4 Gy TBI was delivered to the mice at a high dose rate. This was the same dose as that used in Feng et al.’s work, which promoted tumorigenesis in Trp53-heterozygous mice [24]. This dose could increase incidences of micronucleus bone marrow erythrocytes and cause abnormality in the peripheral blood hemogram in Trp53 wild-type (Trp53wt) mice, according to our preliminary study. Before investigating the effect of exposure to very low dose irradiation and stress, we first determined whether the mouse response to a high dose of irradiation (capable of inducing a detectable effect on the hematopoietic system in Trp53wt mice) could be modulated under chronic restraint-induced stress in Trp53wt mice.

It should be noticed that in the field of radiation biology, IR is considered one type of stress. The term ‘stress’ used in the present work (studying the modifying effect of chronic restraint on radiation hematopoietic toxicity) refers to chronic restraint-induced psychological stress.

MATERIALS AND METHODS

Study subjects and experimental group design

C57Bl/6J mice strain male mice were used in the present study. Based on our preliminary two trials, mice aged 4 weeks old were purchased from SLC, Inc. (Japan) and maintained in a clean conventional temperature- and humidity-controlled animal facility under a 12 h light – 12 h dark photoperiod (lights on from 7:00 A.M. to 7:00 P.M.). Animals housed in autoclaved cages (three mice per cage) with sterilized wood chips, were allowed free access to standard laboratory chow (MB-1, Funabashi Farm Co., Japan) and acidified water (pH = 3.0 ± 0.2). Animals were acclimatized to the laboratory conditions for 1 week before use. The 5-week-old mice were randomly assigned to four experimental groups with 9 to 12 mice in each group, namely, the ‘control group (C-Gr)’ (receiving only restraint nor TBXI), the ‘restraint group (R-Gr)’ (receiving only restraint), the ‘TBXI group (IR-Gr)’ (receiving only TBXI), and the ‘restraint and TBXI group ((R+IR)-Gr)’ (receiving both restraint and TBXI). All experimental protocols involving mice were reviewed and approved (Experimental Animal Research Plan and Protocol No. 12-1026) by The Institutional Animal Care and Use Committee of the National Institute of Radiological Sciences (NIRS). The experiments were performed in strict accordance with the NIRS Guidelines for the Care and Use of Laboratory Animals.

Mouse model for chronic restraint-induced stresses

Chronic restraint, a well-established typical mouse model [30] to induce psychological stresses, was adopted and applied to the present work with minor modification. In brief, the mouse restraint system (Flat Bottom Rodent Holder, RSTR541, Kent Scientific Co., USA) was used for chronic periodic restraint on a daily basis of 6 h for 28 consecutive days. Individual mice aged 5 weeks were placed in the
strainer, and the restrained mice were maintained horizontally in their home cage during the 6-h restraint session (9:30 A.M. to 3:30 P.M.) daily, then the animals were released into the same cage and allowed to access food and water during the free session (3:30 P.M. to 9:30 A.M.). The animals in C-Gr and IR-Gr received no restraint but their food and water was withheld while the R-Gr and (R+IR)-Gr animals underwent the 6-h restraint session each day.

Total body X-irradiation of the mice
X-rays were generated with an X-ray machine (Pantak-320S, Shimadzu, Japan) operated at 200 kVp and 20 mA, using a 0.50-mm Al + 0.50-mm Cu filter. An exposure rate meter (AE-1321M, Applied Engineering Inc., Japan) was used for the dosimetry. The dose of TBXI was 4.0 Gy, and it was delivered at a dose rate of 0.25 Gy/min to the animals in IR-Gr and (R+IR)-Gr on the 8th day of the 28-day restraint regimen. The mice held in acryl containers were exposed to TBXI at room temperature without anesthesia.

Assessment of the peripheral blood hemogram
The body weight gain of the animals in each experiment group was recorded daily. At the end of the restraint regimen, the animals were anesthetized by inhalation of gaseous isoflurane (2-chloro-2-(diluoromethoxy)-1,1,1-trifluoro-ethane) (CDS019936, Sigma-Aldrich, Japan). Then the animals were euthanized by cervical dislocation. Significant decrease in body weight gain, together with decrease in spleen weight and marked alteration in the corticosterone concentration in the peripheral blood plasma, were used as indicators for evaluating the success of CRIS in the mouse restraint model.

Measurement of body weight gain
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Bone marrow micronucleus test
The bone marrow micronucleus test was carried out according to Schmid [42] with minor modifications [43, 44]. Bone marrow smears prepared from both femurs were processed for the enumeration of micronucleated polychromatic erythrocytes (MNPCEs) and micronucleated normochromatic erythrocytes (MNNCEs). The slides were coded to avoid any observer bias. The micronuclei were scored using a light microscope at a magnification of x1000. At least 5000 cells per mouse were counted, and the data for each experimental point were from five to six mice.

RESULTS
Verification of the CRIS Model
The treatments used in this study have been previously shown to induce a stress response sufficient to significantly affect several immunological parameters [30]. Under the experimental setup in the present study, reproducibility of this CRIS model in young male mice with normal genotype was verified using endpoints: namely, significant decrease in body weight gain and in the weight of immune organs (such as thymus and spleen), and alterations in the concentration of corticosterone in peripheral blood plasma. Significantly reduced body weight gain induced by CRIS appeared one day after onset of the restraint and this phenomenon was observed throughout the experiment in the animals that received the restraint (R-Gr and (R+IR)-Gr) (Fig. 1). For the animals that received the restraint, a marked decrease in weight of spleen and increase in concentrations of stress-related hormones and cytokines in the blood plasma were also detected (data not shown as they are to be published elsewhere in a paper on a hormone and cytokine study). These results clearly indicated the successful establishment of the CRIS model of restraint under our experimental setup.

Change in body weight gain
The effects from restraint and TBXI on the body weight gain of mice are shown in Fig. 1. As mentioned, significant decrease in body weight gain was observed one day after onset of the restraint (P < 0.05), and the restraint resulted in the lowest body weight on the third day after onset (P < 0.0001): on the day of restraint onset, the body weight (g) of C-Gr, R-Gr, IR-Gr and (R+IR)-Gr was 21.7 ± 1.1, 21.5 ± 1.1, 22.0 ± 0.8 and 22.0 ± 0.9, respectively; on the following day, the body weight was 21.7 ± 1.2, 20.8 ± 0.6, 21.9 ± 1.1, and 21.9 ± 1.1, respectively. After a single exposure to TBXI with 4 Gy of X-rays, significant t-test, except for the micronucleus data on which the χ² test was performed. Statistical significance was assigned to P < 0.05.
reduction in body weight gain was observed on the following day in both IR-Gr and (R+IR)-Gr (P < 0.05), while no interaction (namely, neither synergistic nor antagonistic effect) between the restraint and TBXI was observed (P > 0.05): on the day just before the animals were given TBXI, the body weight (g) of C-Gr, R-Gr, IR-Gr and (R+IR)-Gr was 23.1 ± 1.3, 21.6 ± 0.3, 23.5 ± 0.8 and 22.3 ± 0.9, respectively; on the following day, the body weight was 23.1 ± 1.4, 21.5 ± 0.4, 23.0 ± 0.9 and 21.7 ± 0.9, respectively. The recovery of body weight gain appeared late in the animals receiving the restraint (R-Gr and (R+IR)-Gr). In general, there was a statistically significant difference in the mean body weight between the groups that received the restraint (R-Gr and (R+IR)-Gr) and the groups that received no restraint (C-Gr and IR-Gr) one day after the onset of restraint, regardless of the TBXI: the mean body weight was markedly higher in groups that received no restraint compared with those groups that received the restraint. At the end of the experiment, the body weight in gram for C-Gr, R-Gr, IR-Gr and (R+IR)-Gr was 25.3 ± 1.5, 22.8 ± 0.4, 25.4 ± 1.3 and 23.0 ± 1.2, respectively. These results indicated that a single TBXI with 4 Gy of X-rays could induce a transient reduction in the body weight gain; however, CRIS for 28 days was more effective in terms of causing a continuous reduction in body weight gain in mice.

Hematological abnormality in the peripheral blood hemogram

Alterations in the hematopoietic system were studied in the peripheral blood hemogram (Fig. 2). Mice subjected to CRIS alone (R-Gr) displayed a significant decrease in both RBC count and WBC count when compared with the control (C-Gr). On the other hand, when compared with the control (C-Gr), mice exposed to TBXI alone (IR-Gr) showed a significant reduction in total peripheral blood cell count, manifesting as decreased RBC count, decreased WBC count, decreased PLT count, and low hemoglobin concentration. In addition, the effect from TBXI alone on induction of a decreased WBC count was significantly higher than that from CRIS alone. In animals subjected to both CRIS and TBXI ((R+IR)-Gr), a decreased RBC count, decreased WBC count, and decreased PLT count were observed. Combination of CRIS and TBXI caused a further marked decrease in the PLT count, and the severity of the decrease in the PLT count was significantly greater than that induced by either CRIS or TBXI alone. These results indicated that CRIS alone could cause, to a certain extent, detrimental effects on the hematopoietic system; however, this effect was not as strong as that caused by TBXI. Except for worsening the decreased PLT count, the combination of CRIS and TBXI seemed to show little effect on TBXI alone–induced hematological abnormality in the peripheral blood hemogram.

Residual damage in bone marrow erythrocytes

Cytotoxicity and genotoxicity of CRIS and TBXI were evaluated by measuring the residual damage in the bone marrow cells of the animals (Fig. 3). The number of polychromatic erythrocytes (PCEs) expressed as a percentage of the sum of PCEs and normochromatic erythrocytes (NCEs) is an indicator of bone marrow proliferation in the erythroid lineage, and its decrease is an indicator of mutagen-induced cytotoxicity [45]. The micronucleus test is a tool for genotoxic assessment. CRIS alone (R-Gr) showed no significant effect on the percentage of PCEs with respect to the sum of PCEs and NCEs, or on the occurrence of MNPCeS in PCEs or the occurrence of MNNCeS in NCEs in the femur bone marrow when respectively compared with that in the control (C-Gr). On the other hand, TBXI results in a marked reduction in the number of PCEs expressed as a percentage of the sum of PCEs and NCEs, and a significant increase in the occurrences of MNPCeS and MNNCeS when respectively compared with that in the control (C-Gr). Exposure to both CRIS and TBXI ((R+IR)-Gr) led to a decreased number of PCEs expressed as a percentage of the sum of PCEs and NCEs, and increased occurrences of MNPCeS and MNNCeS when respectively compared with that in the control (C-Gr). The effects of exposure to both CRIS and TBXI was comparable with that from exposure to TBXI alone. These results indicated that CRIS alone did not have a significant cytotoxic or genotoxic effect on the bone marrow erythrocytes. CRIS did not markedly modulate the cytotoxic and genotoxic effects from TBXI on the bone marrow erythrocytes.

DISCUSSION

Humans are exposed to a multitude of PPSs that have a dramatic adverse impact on health, such as inhibition of the immune system, increased susceptibility to infections, and altered disease risk later in life [46–48]. In fact, it could be that stress can induce genetic, epigenetic and genotoxic changes in humans and animals alike. On the other hand, exposure to IR can result in varied health effects on humans. However, little is known about the combined health consequences from exposure to both stresses and IR. Notably, the contamination of environments with radionuclides resulting from NPPAs could give rise to consequences that encompass far more than health risks from exposure to IR [49]. As the experience of the Chernobyl nuclear disaster demonstrated, the long-term psychosocial consequences are serious. There is a wide range of mental and behavioral sequelae in children following exposure to stress that can last a long time [14] and even cause shortening of telomere length [50]. Poor mental health status due to anxiety about IR exposure has been reported, even in the younger generation born after the accident in the region around Chernobyl [51]. In animal models, traumatic stress in early life was found to alter mouse microRNA expression, and behavioral and metabolic responses in the progeny [52]. CRIS was able to reduce significant body weight gain from 1 week after onset of restraint in rats [53]. CRIS promotes immune suppression, inducing lymphocyte reduction [35, 54], and exposure of rats to continuous stress from photoperiod, temperature and noise was observed to cause an increase in micronuclei incidence in peripheral RBCs [55]. A recent study showed that CRIS increased the susceptibility of Trp53-heterozygous mice to radiation (4 Gy) carcinogenesis [30]. The regulation of multiple Trp53 stress responses was mediated through MDM2 [56], and attenuation of Trp53 function was an important part of the mechanism underlying promotion by CRIS of Trp53-heterozygous mice to radiation carcinogenesis [30]. Stress-induced instability of cellular mechanisms may play an important role in increasing cell division disorders. Exposure to acute restraint stress could enhance the damaging actions of an aneugenic agent (vinblastine) on mouse bone marrow erythrocytes, inducing increased frequency of micronuclei [57].

Both the academic world and the public recognize the importance of psychological consequences arising from a catastrophic accident and its aftermath, and seek strategies for mitigating the serious
Fig. 2. Effect of CRIS and TBXI on the peripheral blood hemogram of mice. Group mean ± SD levels of RBC count (A), WBC count (B), PLT count (C), and hemoglobin concentration (D). * and ** stand for significant difference between two groups compared at $P < 0.05$ and $P < 0.01$, respectively.

Fig. 3. Effect of CRIS and TBXI on the femur bone marrow erythrocytes of mice. Group mean ± SD of the percentage of PCEs to the sum of PCEs and NCEs (A), the number of MNPCEs per 1000 PCEs (B), and the number of MNNCEs per 1000 NCEs (C). * and ** stand for significant difference between two groups compared at $P < 0.05$ and $P < 0.01$, respectively.
cyte proliferation was observed in the bone marrow; however, a sig-

matically decreased RBC count was found in the peripheral blood. As

Thus, is great concern about whether the psychological stress could cause any alterations in the response of human beings to radiation and whether the combined effects of multi-stresses and radiation would be fundamentally additive, syner-
gastic or antagonistic. It is expected that study of the possible effects of psychological stress on responses to radiation exposure and the subsequent consequences could have important implications for the health risk of humans living with exposure to irradiation and psycholo-
gical stresses. Since the Fukushima nuclear accident in Japan, chil-
dren living in radioactively contaminated areas are often restricted or

even prohibited from doing outdoor activities, and this is thought to

cause additional strong psychological stresses. IR is genotoxic to the

highly radiosensitive hematopoietic system, inducing cell injury and

caus ing profound effects [58–60]. The mouse hematopoietic system

provides a suitable model for study of the potential modifying effects of

CRIS on radiosensitivity, functional recovery after IR, and IR-

induced late effect [59]. Psychological stress affects a range of physio-

logical processes, including hematopoiesis. In mice, CRIS decreased

the concentration of hemoglobin in the blood, elevated circulating

levels of erythropoietin and corticosterone, and resulted in a mark-
edly increased number of erythroid progenitors and precursors in the

spleen, leading to the prolonged activation of stress erythropoiesis

pathways, and resulting in excessive production of immature erythroid

cells, which may predispose chronically stressed subjects to a higher

risk of leukaemic transformation [61]. On the other hand, it was

reported that chronic stress could also influence hematopoietic stem

cells (HSC) and lineages, causing increased proliferation of HSCs in

humans and mice, leading to increased numbers of myeloid and

lymphoid progenitors in bone marrow, and resulting in an increased

output of neutrophils and inflammatory monocytes [62]. In the

present study, we set out to identify the effects of radiation on the

hematopoietic system that may be altered in a CRIS mouse model

using alterations in hematology in peripheral blood and incidence of

micronuclei in bone marrow erythrocytes as the endpoints. The

reproducibility of this model was verified by decreased body weight

gain, decreased immune organ weight, and an increased stress

hormone level in the blood plasma. Results of the PLT count showed

a significant reduction in animals that received both CRIS and TBXI

when compared with that in animals receiving either CRIS or TBXI

alone. The biological consequences of a severely decreased PLT

count are bleeding and loss of body fluid. This result may suggest a

health problem requiring further study. However, except for the

results for the PLT count, the results for most of the endpoints did

not reveal a significant synergistic or antagonistic effect from CRIS on

TBXI-induced cytotoxicity and genotoxicity in the hematopoietic

system. In mice receiving CRIS alone, no marked change in erythro-
cyte proliferation was observed in the bone marrow; however, a sig-
nificantly decreased RBC count was found in the peripheral blood. As

the lifespan of RBCs in peripheral blood is ~40 days, and ~18–21 days

are required to produce a mature RBC from the burst-forming unit-

erythroid [63, 64], results may suggest that RBCs in the peripheral

blood may undergo excessive eryptosis, possibly resulting from CRIS-

induced oxidative stress [65], and young RBCs produced under CRIS

conditions [66] may be predisposed to CRIS and have a short life-

span in the peripheral blood.

Though different endpoints were used, the results obtained in the

present study are not inconsistent with the report on increased

susceptibility induced by CRIS for Trp53-heterozygous mice to radia-
tion carcinogenesis [30]. It should be noticed that, although the mice

used in these two studies were of the same strain, Trp53wt animals

were used in the present study. It is known that chronic stresses induce

susceptibility to pathogens, and that toxicological assaults (including

IR) on health are dependent on the genetics of the exposed organism

[14, 31–34, 37, 67]. Based on these previous studies and the results

obtained in the present work, we suggest that CRIS would have little

influence on the sensitivity of Trp53wt mice to radiation effects on the

hematopoietic system, including genotoxic effects. It is possible that

the methodology of our experimental system was not sensitive enough

to detect the influence of CRIS on the genotoxic effect. To improve

the sensitivity for detection of genomic damage, further study using

the fluorescence in situ hybridization technique for detection of

chromosome aberrations in splenic cells is in progress. As investigations

in both humans and laboratory animals have demonstrated the exist-

ence of sex differences in response to stresses [68–72] (a recent study

even showed that chronic prenatal restraint stress could induce

memory impairment in a sex-specific manner in post weaning rats

[73]), further study using different strains of mice and both genders is

recommended.

In summary, our results suggest that CRIS does not have a signi-

ficant modifying effect, either synergistic or antagonistic, on radiation-

induced detrimental effects on the hematopoietic system in young

Trp53wt mice under the present experimental setup. For most people,

especially those living in radioactively contaminated areas, the present

work may partially allay their concern that stresses could increase their

susceptibility to cancer as a result of radiation.

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CONFLICT OF INTEREST

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