Low-dose ionizing radiation as a hormetin: experimental observations and therapeutic perspective for age-related disorders

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Abstract Hormesis is any kind of biphasic dose-response when low doses of some agents are beneficial while higher doses are detrimental. Radiation hormesis is the most thoroughly investigated among all hormesis-like phenomena, in particular in biogerontology. In this review, we aimed to summarize research evidence supporting hormesis through exposure to low-dose ionizing radiation (LDIR). Radiation-induced longevity hormesis has been repeatedly reported in invertebrate models such as *C. elegans*, *Drosophila* and flour beetles and in vertebrate models including guinea pigs, mice and rabbits. On the contrary, suppressing natural background radiation was repeatedly found to cause detrimental effects in protozoa, bacteria and flies. We also discussed here the possibility of clinical use of LDIR, predominantly for age-related disorders, e.g., Alzheimer’s disease, for which no remedies are available. There is accumulating evidence that LDIR, such as those commonly used in X-ray imaging including computer tomography, might act as a hormetin. Of course, caution should be exercised when introducing new medical practices, and LDIR therapy is no exception. However, due to the low average residual life expectancy in old patients, the short-term benefits of such interventions (e.g., potential therapeutic effect against dementia) may outweigh their hypothetical delayed risks (e.g., cancer). We argue here that assessment and clinical trials of LDIR treatments should be given priority bearing in mind the enormous economic, social and ethical implications of potentially-treatable, age-related disorders.

Keywords Low-dose ionizing radiation · Radiation hormesis · Hormetin · Longevity · Animal models · Age-related disorders

Introduction: what is radiation hormesis?

Back in the 1880s, Hugo Schulz observed that low doses of many toxic agents, like mercury and formaldehyde, enhanced the vitality of yeast cells (Schulz 1887, 1888). The term “hormesis” was first introduced in 1940s in the context of chemical toxins (Southam and Ehrlitch 1943). Hormesis in general is any kind of biphasic dose-response when low doses of some agents are beneficial while higher doses are detrimental (Calabrese 2015, 2018, 2019). According to the present knowledge, hormesis-inducing agents...
('hormetins') include but are not limited to a variety of physical and chemical factors such as heat and oxidative stress, various food components, micronutrients, intermittent fasting, calorie restriction, physical exercise etc. (Musci et al. 2019; Rattan and Kyriazis 2019; Calabrese et al. 2020a).

Radiation hormesis is the most thoroughly investigated among all hormesis-like phenomena, in particular in biogerontology. Speaking about radiation hormesis, we should point two somewhat different uses of the term ‘hormesis’. Since radiation carcinogenesis is usually considered to be the single most important detrimental health factor of ionizing radiation, radiation hormesis is sometimes understood in the narrow sense that low radiation doses can suppress cancer. In this narrow sense, curing arthritis or pneumonia is not viewed as a hormetic effect. Accordingly, there are two quite different meanings of the term ‘low dose’. In the context of radiation protection and many fields of radiobiology, ‘low dose’ is understood to be 100 mGy or less-there being a consensus (more or less) that there is no solid evidence for radiation carcinogenesis below 100 mGy. However, in the field of radiation oncology where the daily dose fraction is typically 2000 mGy and 6 weeks of therapy amounts to a total dose of 60,000 mGy, a single 1000 mGy dose to treat pneumonia is regarded as a low dose (Calabrese et al. 2020b). In this review article, we use the term ‘hormesis’ in the wide sense of biphasic dose-response, and consequently ‘low dose’ as corresponding to the beneficial-effect part of the dose-response curve (Cuttler 2020). Therefore, the meaning of ‘low dose’ is species-specific.

In this review, we aimed to summarize research evidence supporting hormesis through low-dose ionizing radiation (LDIR) in age-related disorders and discuss the possibility of using this phenomenon in clinical settings.

**Historical overview**

Since the discovery of a radioactivity in 1896, there is vast amount of data regarding health effects from ionizing radiation exposure. Sufficiently high doses undoubtedly cause harm—that is a matter of proper scientific consensus. However, a major controversy exists among scientists and medical professionals regarding health effects of radiation at relatively low doses and dose rates that does not cause immediate harm. At the end of the XIX and early XX century, there was a general belief that beneficial consequences of irradiation are widespread and nearly miraculous. “Mild radium therapy” has been widely used and probably more widely misused. It was claimed, e.g., that blindness could be cured by using X-rays. It has been advertised that several mixtures might treat about 150 disorders, including lassitude and sexual impotence. Over the period between 1925 and 1930, about 400,000 bottles with radium-containing water have been sold. The tragic death of the famous millionaire Eben M. Byers in 1932 as a consequence of radium poisoning became a widely-known event that alerted both the public and the medical community of adverse radiation effects. This event probably ended the era of the “mild radium therapy” (Blaufox 2019).

After the Hiroshima and Nagasaki atomic bombing in 1945, the attitude to ionizing radiation began to lean to another extreme: ionizing radiation became connected in public mind with nuclear apocalypse. The linear no-threshold (LNT) model for radiation risk assessment, introduced after Muller’s discovery of radiation-induced mutations in 1927, became popular. This model suggests that any radiation exposure constitutes the risk that increases proportionally with increasing exposure. The main radiation concern was mutagenesis. Later (at end of the 1950s), after no radiation mutagenesis was found in the A-bomb survivors’ descents (Satoh et al. 1996; Kodaira et al. 2004), carcinogenesis became the main concern. LNT has been introduced as a reasonable operational model for the radiation protection only. Formally speaking, this is valid even now: UNSCEAR, ICRP and other scientific bodies officially advise against using LNT for cancer morbidity/mortality predictions. De facto, however, LNT acquired the status of scientific theory, though supporting evidence is at least inconclusive. Extrapolation of effects from the high to low exposure doses is not founded as any other extrapolation. Moreover, such extrapolation neglects the fact that all organisms were always exposed to ionizing radiation and that multiple mechanisms to repair radiation damage evolved during the evolution (Boothby et al. 2019). The model of radiation hormesis is based on the assumption that repair mechanisms, triggered by the LDIR, might be efficacious not only against the radiation-induced damage, but also against damage from other stressors. Therefore, these mechanisms...
may cause overall improvement of health including decreased cancer (Calabrese et al. 2015; Costantini and Borremans 2019).

Beneficial effects of LDIR are known since the ancient times: already Herodotus and Hippocrates describe beneficial effects of radon springs. In the modern times, Roentgen-ray stimulation of growth has been observed in X-ray exposed algae (Atkinson 1898) and these findings were confirmed in subsequent studies (Conter et al. 1983). As mentioned above, X-rays and radium sources have been widely applied in the first half of the XX century for treating arthritis, pneumonia and more (Calabrese et al. 2020b). During the Manhattan Project, rodents exposed to inhaled uranium dust were studied. The scientists were surprised to learn that exposed animals were healthier and were characterized by higher life expectancy and more offspring than the controls, though the dust concentration had been expected to be fatal at the experimental set-up (Brucer 1990). Later in 1958, the first United Nations Scientific Committee on the Effects of Atomic Radiation (UNSCEAR) report provided experimental evidence for the longer survival of rodents such as mice and guinea pigs after exposure to low doses of gamma radiation and fast neutrons. These data have been interpreted as indicative of existence of a threshold, but the hormesis was not mentioned (UNSCEAR 1958; Calabrese 2020).

The pioneering studies of hormetic phenomenon carried out in the mid-20th century pointed that hormetic effects can be more pronounced in populations maintained in suboptimal conditions or in non-healthy individuals (Sacher 1977). For example, statistically significant life extension was observed in short-lived dogs, but not in the dog population in general (Cuttler et al. 2017a, b). However, when experiments with chronic radiation exposure have been done with high standards of the animal care, hormetic effects diminished but were still observable (Sacher 1977). Unfortunately, since 1960s hormesis-like effects of LDIR have been systematically ignored by both the medical community and by radiation protection agencies—despite more than 2000 peer-review papers on this subject in the PubMed database as for August 2020. Over the previous decades, beneficial effects induced by the LDIR exposure were repeatedly found in cell cultures, microorganisms, plants, and different invertebrate and mammal models. The effects were observed at all levels including the biochemical, cellular, tissue and the whole-organism ones. Radiation hormesis was revealed for various physiologic functions. Observed effects included enzyme induction, growth, cell division, neuromuscular development, metabolism, hearing, visual acuity, as well as memory and learning performance. In mammals, radiation hormesis has been demonstrated to enhance protection against various infectious and neoplastic disorders, to improve fertility and to increase longevity (Scott 2014). From the 1950s until the beginning of 1980s, four major mechanisms were generally discussed to explain this phenomenon. It was suggested that radiation exposure can: (1) reduce the amount of detrimental bacteria in the organism (Sacher 1956); (2) inactivate gonadal function causing the reallocation of vital resources in favor of somatic maintenance at the expense of reproduction (Lamb 1964); (3) reduce the oxygen consumption rate and, therefore, metabolic activity (Allen and Sohal 1982); (4) induce over-compensatory repair responses (Cork 1957; Carlson and Jackson 1959). Presently, only the last assumption—over-compensatory repair—is generally accepted. In line with current assumptions, the contributing mechanisms include, along with stimulation of the DNA repair, modification of gene expression, stress protein production, detoxication of reactive oxygen species (ROS), release of growth factors, activation of membrane receptors, compensatory cell proliferation and stimulation of the immune system (Feinendegen 2005; Calabrese et al. 2010; Shibamoto and Nakamura 2018; Vaiserman et al. 2018).

Life extension is among the well-established beneficial health effects of LDIR (Cameron 2003). The effects of radiation on longevity have been evaluated in a variety of animal models. Lifespan decrease is well-known outcome of the high-dose radiation exposure (Lamb 1964; Brown 1966; Gould and Clark 1977; Giess and Planel 1977; Giess 1980). However, lifespan extension has been also repeatedly observed in laboratory animals irradiated with low doses. In these studies, the mean lifespan was often increased by 10–30% while the maximal lifespan has been usually unchanged (Calabrese and Baldwin 2000; Cuttler et al. 2017a, b).
Radiation hormesis in invertebrate models

Controversial data about effects of LDIR exposure on lifespan have been obtained with a nematode *Caenorhabditis elegans*. Multiple stressors—like heat, ROS-generating chemical juglone, hyperbaric oxygen, etc., —promoted enhanced lifespan in the worms, but treatment with ultraviolet radiation or with X-rays did not (Cypser and Johnson 2002). No lifespan extension has been obtained following gamma-irradiation at the dauer larvae stage (Yeargers 1981). Substantial occasional (non-repeatable) increase of survival in *C. elegans* after radiation exposure with intermediate doses of 100–300 Gy was reported only once, in the study by Johnson and Hartman (1988).

Radiation hormesis was repeatedly reported in insect models (Calabrese 2014). Experiments with a flour beetle such as *Tribolium confusum*, are among the first studies providing evidence for this phenomenon (Davey 1917, 1919). In the experiment with acute exposure, five X-ray doses were applied (Davey 1917). These doses were later reported (cited in Calabrese and Baldwin 2000) as 100–500 mA/min at 25 cm at 50 kV (100–500 MAM/25 at 50 kV), or 5–25 Gy; the dose of 12.5 MAM/25 corresponds to approximately 60 rad or 0.6 Gy (Cork 1957). In this experiment conducted with 1100 beetles, the least dose required to kill all test animals (LD100) was 500 MAM/25 at 50 kV (5 Gy). Death rates lower than those found in controls were, however, observed by irradiation with 100 or 200 MAM/25 at 50 kV (5–10 Gy). In the subsequent research, the effects of X-rays on longevity have been evaluated following either acute X-ray exposure (as in previous study) or through daily exposures (Davey 1919). In the daily exposure experimental block performed with appr. 950 beetles per each group, five doses have been applied, ranging from 6.25 to 50 MAM/25 at 50 kV (0.3 to 2.5 Gy) per day. By 30 days after irradiations, 25–40% decrease in mortality was demonstrated in experimental groups irradiated with three lowest doses. In the second experimental block conducted with ~ 850 beetles per group, single doses in the range of 100–400 MAM/25 at 50 kV (5–20 Gy) were used. Confirming the results from the initial study, a reduced mortality rate has been found by 20 days after the irradiation. The Davey’s findings were subsequently confirmed in experiments by Cork (1957) conducted with large number of flour beetles. In these studies, however, irradiations were performed with cesium-137 used as a source of gamma rays. Both acute and chronic radiation exposures were shown to be able to promote lifespan extension. More recently, radiation-induced longevity hormesis has been obtained in another beetle species such as *Tribolium castaneum* (Ducoff 1975). Irradiation reduced the early mortality rate thereby resulting in more “rectangular” survival curves. However, little or no increase in maximal lifespan was revealed. Moreover, irradiated insects were found to be more resistant to ether hyperbaric oxygen or heat stresses (Lee and Ducoff 1983; Ducoff and Lee 1984). Interestingly, stress resistance was found to be higher in the irradiated adults than in young controls, thus, this effect was not simply due to slowing of the aging rate.

Low-dose gamma irradiation also was found to be able to increase mean lifespan of the house fly, *Musca domestica* (Allen and Sohal 1982). Adult house flies exposed to 0, 20, 40 and 66 kR (kR—kiloroentgen, about 10 Gy; 1R ≈ 10 mGy) were maintained in conditions of either low or high physical activity. In the high-activity cohort, mean lifespan of those flies who have been exposed to 20 or 40 kR was larger than that in the control insects, but it has been significantly decreased after irradiation with 66 kR. In both sexes, the radiation exposure led to a significant reduction in oxygen consumption rate. The effects of radiation were shown to be substantially dependent on environmental conditions. Life-extending effects were detected only if flies were reared in groups, i.e., a condition which promotes a high locomotor activity level. If singly reared (a low-level locomotor activity condition), the longevity hormesis did not appear. In addition, irradiated insects had enhanced longevity compared to control individuals only when the latter have been kept in suboptimal conditions. Irradiation also resulted in decreased rate of accumulating fluorescent age pigment (lipofuscin) in high-activity groups, while increased rate of lipofuscin accumulation was found in flies maintained in low-activity conditions (Allen 1985). The reduced metabolic activity was proposed as a potential explanation for such a radiation-induced life-lengthening (Allen and Sohal 1982).

The longevity-modulating effects of ionizing radiation were most thoroughly investigated in the *Drosophila melanogaster*. High doses have been
repeatedly shown to lead to life shortening of fruit flies (Lamb 1964; Nelson 1973; Giess and Planel 1977; Gould and Clark 1977; Giess 1980). In discussing the reported life-extending effects of irradiation in fruit flies, Sacher (1963) has noted a reduced variability in average survival times among replicates. In this discussion, he attributed both the observed reduction in variability and the improved survival to neutralization of some disadvantageous environmental factor(s) via the radiation exposure. By the Sacher's point of view, it cannot be, however, determined whether such a reduced efficiency is caused by inactivation of certain adverse environmental factor(s) or from increasing the resistance induced by a radiation exposure. Alternative hypothesis for explaining the observed hormetic effects (at least in female flies) was proposed by Marion Lamb (1964). She has suggested that irradiation may lead to a sterility in female flies and that the extension of longevity might be due to lowering the level of synthetic metabolism in gonads. In supporting this hypothesis, she has demonstrated that mutant female flies without ovaries did not exhibit a longer lifespan following the irradiation. Later, Giess et al. (1980) have, however, postulated that radio-induced sterility is not a cause of these effects. In this study, no lifespan-modulating effects were obtained following exposure to 10 kR, a dose known to completely inhibit fecundity. These results cause doubt that reduced fecundity per se could be a plausible mechanism for the hormetic effects of irradiation. In addition, such effects were found to be more pronounced if a dose-fractionation regime was used. Females demonstrated a higher increase in longevity following dose fractionation compared to the age-matched males (Mohsin 1979). More specifically, females demonstrated 31% increase in lifespan, whereas only 12% increase was observed in males.

In a series of studies, the potential role of the apoptosis (programmed cell death) in the radiation-induced longevity hormesis in Drosophila has been investigated. Chronic irradiation with cumulative doses of 0.6 to 0.8 Gy resulted in an increase in mean lifespan in male wild-type D. melanogaster and in decreased lifespan in mutant Drosophila strains defective in DNA repair and displaying an enhancing sensitivity to an apoptosis induction (Zainullin and Moskalev 2001). In the mutant strain carrying the pro-apoptotic ‘reaper’ (rpr) gene, an increased lifespan was obtained either after the irradiation and/or following treatment with the inductor of apoptosis, etoposide (Moskalev and Zainullin 2001). Moreover, an association has been shown between the radio-induced apoptosis in cells of the larval ganglion and the aging rate (Moskalev and Zainullin 2004). The authors assumed that these effects were due to eliminating damaged or unwanted cells by the radiation-induced apoptosis (Moskalev et al. 2006). An important role of genes encoding heat shock proteins and heat shock factors (Moskalev et al. 2009) as well as longevity-associated factors such as FOXO, SIRT1, JNK, ATM, ATR, and p53 (Moskalev et al. 2011) in inducing radiation-induced hormetic effects in fruit flies also has been demonstrated. More recently, significant contribution of genes implicated in the nucleotide excision repair (mei-9, mus210, Mus209), base excision repair (Rrp1), DNA double-stranded break repair by homologous recombination (Brca2, spn-B, okr), DNA damage sensing (D-Gadd45, Hus1, mnk), non-homologous end joining (Ku80, WRNexo), as well as Mus309 known to participate in the DNA repair processes has been revealed (Koval et al. 2020). In flies with mutations in these genes radiation hormesis was completely absent or appeared to a lesser extent than in wild-type Canton-S strain. The LDIR exposure also led to hormesis-associated alterations in gene expression in Drosophila; these changes, however, were found to be not dose-dependent (Zhikrevetskaya et al. 2015). In a genome-wide analysis, approximately 13% of the genome of low-dose irradiated male D. melanogaster displayed changes in gene expression; among them, many of aging-related genes have been shown to be significantly regulated (Seong et al. 2011).

The effects of X-ray irradiation early in life on the flies' lifespan were also revealed. It must be taken into account that fruit flies develop with metamorphosis, and ontogenetic stages differ significantly in radiosensitivity (see Fig. 1 for illustration). It is because Drosophila imago consists mainly of radioreistant postmitotic tissues, with the exception of some gut cells and the gonads (Rogina 2011), while active cell division occurs during pre-adult stages from egg to pupa.

A significant trend to decrease median lifespan with increasing irradiation dose has been observed in both male and female flies irradiated with 1.2, 2.1, 4.2, 7.5 and 17.1 Gy at larval stage. The maximum lifespan,
however, was found to be increased on 11.5% and 12.7% in males irradiated with 1.2 and 2.1 Gy, respectively (Vaiserman et al. 2004a). X-irradiation of the 1-h eggs with doses of 0.25, 0.50, 0.75, 1, 2 and 4 Gy was also found to affect lifespan (Vaiserman et al. 2003a). Longevity hormesis has been found in males (but not in females) irradiated with 0.5 and 0.75 Gy. X-ray irradiation at the egg stage with 0.75 Gy caused ultrastructural changes in the flies’ brain cells (Vaiserman et al. 2003b). Irradiations at this stage also caused DNA modifications such as decreased amounts of DNA segments resulting from a cleavage of the S1 nuclease-sensitive sites (< 3 kb) compared to that in control flies (Fig. 2). The higher stability of DNA might be the result of activating the repair system of irradiated flies. It has been hypothesized by the authors that these effects were due to structural and/or functional DNA modifications which occurred following the irradiation at the egg stage and persisted in adult tissues. Such modifications, if any, might change the repair and/or transcription processes, thereby affecting the lifespan. Evidence was also obtained that effects induced by early-life X-ray irradiation may persist across generations. Irradiation of 1-h eggs with 0.25, 0.5 and 0.75 Gy resulted in decreased body weight and increased locomotor (both photo- and geotactic) activities in F0 and F1 generations (Vaiserman et al. 2004b). Moreover, increased resistance to the starvation and heat shock stresses as well as longevity hormesis have been observed in irradiated flies and their offspring. Confirming evidence for the possibility of cross-generational transmission of effects induced by gamma irradiation in ancestral generation on the flies’ longevity has been also observed in the Shameer et al. (2015) study. Irradiation of two-to three-day-old male parental flies with small-to-moderate doses (1–10 Gy) lead to a significant lifespan extension in both male and female offspring, whereas exposure to higher doses (40 and 50 Gy) resulted in a decreased longevity in F1 and F2 offspring generations. These effects on longevity disappeared in the F3 generation.

Radiation hormesis in vertebrate models

Radiation-induced longevity hormesis was reported in several vertebrate models. First evidence for that has
been provided by Lorenz and colleagues from the U.S. National Cancer Institute. In an initial study, guinea pigs, rabbits and LAF1 mice have been exposed to 8.8, 4.4, 2.2, 1.1 and 0.11 R per day from early adult age until death (Lorenz et al. 1954). Irradiated animals have been reported to live longer by 2–14% than controls. However, these beneficial outcomes were unexpectedly accompanied by increased tumor incidence in several experimental groups. These findings were subsequently confirmed in the study by Lorenz et al. (1955). The authors hypothesized that these effects could be caused by improvement of immune defense mechanisms against infectious agents which are known to be the most common cause of death in rodents. A significant increase of median and maximal survival times above control values has been observed in rats exposed for 12 months to gamma irradiation at daily irradiation doses ranging from 0.3 to 4.0 R/day combined with heat or cold stresses (Carlson et al. 1957; Carlson and Jackson 1959). Most substantial life extension (increase of 30%) has been observed after exposure to 2.5 R/day. More recently, significant lifespan extension was found in female C57BL/6 mice chronically exposed to gamma irradiation at very low-dose rates (7 or 14 cGy/year): the median survival times were 549 days in control animals, while they reached 673 days in both exposed groups (Caratero et al. 1998). These findings, however, were not confirmed in a subsequent study by the same authors where no significant differences, neither in longevity nor in cancer or non-cancer disorders, were observed among control and irradiated animals (Courtade et al. 2002). Chronic low-dose rate gamma irradiation at 0.35 or 1.2 mGy/hour promoted longevity in MRL-lpr/lpr mutant mouse strain with a deletion in the apoptosis-regulating Fas gene known to markedly shorten lifespan owing to severe autoimmune disorder (Ina and Sakai 2004). Further extension of irradiation period to the entire lifespan at the same dose rates led to an even more pronounced increase of survival time (Ina and Sakai 2005). The 50% survival time for the control non-treated mice, 134 days, was found to be prolonged nearly four-fold (to 502 days) following the life-long irradiation at 1.2 mGy/hour. In addition, such a radiation exposure led to the immune system activation. More specifically, it resulted in a substantial increase in the numbers of CD3 + CD45R/B220 + cells and CD45R/B220 + CD40 + cells in the spleen. No evidence for prolonged longevity in mice exposed to very low-dose-rate gamma-rays was, however, found in research by Tanaka et al. (2003) conducted with an extremely large mouse sample (total n = 4000). Irradiations were carried out during 400 days with 137Cs as a source of gamma-rays at dose rates of 0.05, 1.1 and 21 mGy/day. The accumulated doses were about 20, 400 and 8000 mGy, respectively. The lifespan of females irradiated with 1.1 mGy/day and animals of both sexes irradiated with 21 mGy/day was significantly decreased compared to those in the control group. Recently, life-shortening effects of chronic low-dose-rate irradiation (400 days at 20 mGy/day) were reported in calorie-restricted mice (Yamauchi et al. 2019). However, hormesis-like effects were reported by another study in ApoE–/– mice following very low-dose and dose-rate external chronic radiation exposure of up to 28 µGy/h for 8 months (Ebrahimian et al. 2018). These effects were found to be associated with an increased expression of anti-inflammatory and anti-oxidative gene. Remarkably, dose rates used in this study (12 and 28 µGy/h) were similar to those measured in contaminated areas like Chernobyl and Fukushima.

Evidence for radiation-induced longevity hormesis has been also obtained in several non-rodent mammalian species. One example is the study conducted with chipmunks, Tamias striatus (Thompson et al. 1990). Wild chipmunks have been captured, exposed to ionizing radiation at single doses of either 200 or 400 rads, and returned to their natural habitats. Irradiated chipmunks demonstrated a biphasic response in age-specific mortality rate. A residuum of unrepaired injury appeared to persist and manifest throughout the entire lifespan. However, another response—longevity hormesis—was also evident in this study. Effects of LDIR were also investigated in dogs. Cuttler et al. (2018a, 2018b) recently determined a dose rate threshold at about 600 mGy per year for lifespan reduction in dogs irradiated lifelong with cobalt-60 gamma radiation. In dog models, hormentic effects following irradiation were also demonstrated. In a meta-analysis of two large-scale studies, one with ten groups exposed to different gamma dose rates and other with eight groups exposed to different lung burdens of plutonium, Cuttler et al. (2017a, b) showed that dogs (especially short-lived ones) benefit when
radiation is moderately above the background level. The maximal lifespan increase occurred at 50 mGy/year. For inhaled α-emitting particulates, longevity of short-lived dogs was shown to be substantially increased below the threshold for harm. In beagles, for lifetime cumulative skeletal doses below 10 Sv from ingested Sr-90, the risk of bone sarcoma was significantly lower than that for controls (Raabe 2010, 2015).

Detrimental outcomes of suppressing background radiation

One strong argument in favor of beneficial effects of LDIR was provided from the facts indicating that suppressing background radiation can detrimentally affect the viability of many organisms. This could be because life on our planet has evolved in conditions of continuous exposure to ionizing radiation, which 3.5 billion years ago was roughly three times higher than presently (Jaworowski 1997). Exposure to artificially lowered levels of natural radiation was found to cause deficiency symptoms in various protozoa and bacteria. These symptoms included, among others, the dramatically decreased proliferation of these organisms (Planel et al. 1966, 1969). Lead shielding of cultures of blue-green algae, *Synechococcus lividus*, also led to lowering the cell growth rate (Conter et al. 1983). This effect disappeared when a normal radiation level (equal to that of background radiation) has been restored within the lead chambers. Irradiation with Th-source at 14-fold natural background dose rate, on the contrary, stimulated the growth of this alga. A study performed with a protozoa *Paramecium tetraurelia* and cyanobacteria *Synechococcus lividus*, which have been either shielded against the background radiation or exposed to low-dose gamma radiation, also showed that radiation may stimulate the proliferation rate of both these single-cell organisms (Planel et al. 1987). The magnitude of this hormetic effect depended either on internal factors (age of starting cells) or external factors (lighting conditions). A stimulatory effect occurred only in a restricted dose range and disappeared with the increase of the dose rate above 50 mGy/year. Similar research findings were obtained with *D. melanogaster* fruit flies. Shielding from natural ionizing radiation resulted in a delayed development (Planel et al. 1967a), and also in a decreased reproductive performance (Planel et al. 1967b) and longevity (Giess and Planel 1973; Planel and Giess 1973).

Overall, these findings indicate that background ionizing radiation may play essential roles in determining the adaptive potential of an organism. Based on this, Sacher (1977) has assumed that free radicals generated following exposure to background radiation may act as primers for certain metabolic processes, thereby affecting the viability of all living organisms. This assumption was subsequently confirmed in a variety of studies. Currently, it is commonly recognized that free radicals such as reactive oxygen species (ROS) or reactive nitrogen species (RNS) are important second messengers in various signal transduction pathways critical for cell growth and proliferation; so, they may play essential roles in multiple vital processes (Milkovic et al. 2019; Huang and Li 2020).

Mechanistic considerations

The LNT model implies that the organism has constant capability to repair damages caused by radiation exposure, irrespective of dose and dose rate. However, there is increasing evidence that such a suggestion is not true. It has been consistently reported that many organism’s responses may be inhibited by high-dose radiation exposures but stimulated by LDIR exposures. In particular, such pattern of response has been well documented for the immune response (Cui et al. 2017) and DNA repair (Pollycove and Feinendegen 2003). Accumulating data indicate that response to irradiation depends on factors such as radiation source distribution, radiation track structure, temporal pattern of radiation exposure, total accumulated dose, dose rate, as well as on the structure and dimension of the biological targets (Howell 2016). The radiation hormesis model, in contrast to the LNT model, predicts that LDIR could induce multiple adaptive responses, and such responses might prevent certain environmentally-induced unfavorable health effects. With regard to DNA repair capacity, the resulting effect can be dependent on the balance between the DNA damage rate (linear with the dose) and particular mechanisms which are responsible for cellular defense (Dobrzyński et al. 2019). Therefore, the response to LDIR exposure could evolve from damage on the molecular level, to beneficial adaptive response on a
whole-body level. When the dose does not exceed 0.1 Gy, the beneficial outcomes tend to outweigh the harmful ones (Feinendegen et al. 2007; Scott and Tharmalingam 2019). The LDIR can likely stimulate repair mechanisms able to repair the primary damage and protect the organism from subsequent stressful—either radiation or other—exposures (Kim et al. 2015). In addition, following these processes, preneoplastic and other damaged cells could be eliminated by apoptosis, immune surveillance and cellular competition (Anzai et al. 2012). The key components of this radiation-induced hormetic response include: ROS scavenging, synthesis of heat shock proteins, secreting specific growth factors and cytokines, activating the cell-membrane receptors, as well as compensatory cell proliferation (Feinendegen et al. 2007; Szumiel 2012). These processes are likely mediated by coordinated adaptive alterations in specific epigenetic pathways (Vaiserman 2008, 2010). Adaptive responses induced following acute exposure to LDIR have differing time schedules depending on the implicated protective pathways. These responses, e.g., DNA repair, may be activated immediately or with delay of hours to days, and some of them (e.g., immune response) may last for several days to weeks or even months. If these protective mechanisms act more efficiently in a low-dose range, then it is quite reasonable that dose-response patterns observed following exposures to LDIR could be not linear but rather threshold or biphasic (Agathokleous and Calabrese 2020). The observed relationships may be apparently affected by various integrative endpoints including patterns of growth, tissue repair, cell proliferation, adaptive preconditioning responses, complex behaviors and also aging processes (Calabrese 2018). Furthermore, these associations could be further complicated by adaptive responses in cells pre-exposed to LDIR, genomic instability manifested in the progeny of irradiated cells as well as by effects in non-targeted cells (so-called bystander effects) (Mothersill et al. 2019).

Following an irradiation, the strategies of cell defense depend on the radiation dose and dose rate, and also on the amount of damages in neighboring cells (Pouget et al. 2018). However, it remains unclear so far whether these mechanisms may act cooperatively at the whole-organism level. Over some last decades, accumulating evidence was provided that low-dose/low-dose-rate radiation exposures may trigger protective responses in vivo as well (see Fig. 3 for schematic illustration). Such exposures were repeatedly shown to activate multiple inter- and intracellular pathways, thereby causing improved protection against many cancers and other genomic instability-related disorders (Feinendegen et al. 2007). Below, molecular, cellular and whole organism-level mechanisms potentially underlying hormetic effects of LDIR are discussed in more details.

DNA repair

DNA Double-Strand Breaks (DSBs) induced by various environmental factors including ionizing radiation are well known to be able to disturb genome integrity and impair cellular viability. Increasing evidence, however, indicates that danger from exposures to LDIR is generally negligible as compared to DNA damage caused by oxidation process normally occurring during the metabolism (Pollycove and Feinendegen 2003; Feinendegen 2005). Moreover, LDIR-induced protective responses may likely overcompensate genotoxic effects caused by byproducts of the normal oxidative metabolism (Azzam et al. 2016). This point of view is based on research data indicating that non-radiation damage of DNA is substantially larger in different tissues than DNA damage caused by radiation exposure at background (and even at much higher) levels (Pollycove and Feinendegen 2003). Based on these theoretical considerations, it is suggested that LDIR may have a dual effect on DNA. Absorption of radiation energy by the DNA molecule represents a direct outcome of the radiation exposure. This absorption can induce various structural changes in the DNA. Moreover, free radicals produced through interactions of radiation with certain molecules of intracellular water may result in a damaged DNA structure. The probability for a DNA damage per each energy deposition event increases proportionally to the absorbed dose. At the background-level radiation exposure, the DNA damage rate is orders of magnitude lower than that induced by different endogenous sources—primarily, by ROS occurring as byproducts of the normal metabolism (Pollycove and Feinendegen 2003). DSBs caused by acute radiation exposure of up to 200 mGy have been repeatedly shown to be efficiently repaired at 24 h post-irradiation in actively proliferating human cells (Rothkamm and Lobrich
Furthermore, accumulating evidence demonstrate that level of DSBs in cell cultures exposed to a LDIR usually decreases to level characteristic of that in unexposed cultures if irradiated cells are allowed to proliferate after the radiation exposure. This can likely be due to elimination of cells carrying unrepaired DSBs (Rothkamm and Lobrich 2003). These findings demonstrate that effects of LDIR are in strong contrast with those of high-dose exposures. Damages caused by the LDIR can be completely recovered while high-dose exposures usually lead to the occurrence of residual DSBs. Thus, whereas linear dose-response relationship may indeed exist for the initial DSB induction, the DNA damage induced by low dose radiation has few chances to persist for any significant time period in cell populations (Suzuki and Yamashita 2012). In addition, adaptive responses against DNA damage may occur following the LDIR exposure (Nenoi et al. 2015). Remarkably, such protective effects have been demonstrated to generally disappear at doses which exceed 100–200 mGy, and they were never observed following acute exposures which
exceed 500 mGy (Feinendegen 2005). Therefore, it can be assumed that LDIR could stimulate DNA repair pathways thereby reducing levels of both radiation-induced and spontaneous damages (Pollycove and Feinendegen 2003). Importantly, the reduction of the spontaneous rate of DNA damage that occur as a result of these processes can subsequently lead to reduced risk of cancer and various aging-associated conditions (Feinendegen et al. 2004). Moreover, other molecular mechanisms could also contribute to LDIR-triggered protective responses. The formation of specific DNA repair centers which arise as preferential sites of repair might be among such mechanisms. Such centers commonly referred to as radiation-induced foci, have been characterized by the local recruitment of p53 binding protein and other DNA damage sensing proteins (Neumaier et al. 2012).

Endogenous antioxidant systems

An increase of capability of endogenous antioxidant systems is another mechanism potentially contributing to hormetic effects (Sharma et al. 2019). It is commonly accepted that oxidative stress caused by excessive generation of ROS lead to damage of various cell components. Among detrimental effects induced by oxidative stress in biological systems, there are genetic instability and mutagenesis, mitochondrial dysfunction, membrane lysis, as well as cellular death (Di Meo et al. 2016; Islam 2017). Therefore, oxidative stress is regarded as an essential contributor in many chronic diseases including chronic inflammation, cardiovascular and neurodegenerative disorders, and also cancers. ROS excess may substantially contribute to aging process per se—e.g., through the accelerated telomere attrition (Koliada et al. 2015). Accumulating evidence from animal models indicate that exposure to LDIR may lead to activation of pathways involved in endogenous antioxidant defense, such as superoxide dismutase, catalase, glutathione, glutathione reductase and glutathione peroxidase in different tissues including spleen, liver, pancreas and brain, and it may cause stabilization of the ROS level (Kataoka et al. 2013; Sharma et al. 2019). These findings were confirmed in several human studies conducted in different occupational groups (Eken et al. 2012; Ahmad et al. 2016). The role of the transcription factor such as nuclear erythroid 2-related factor (Nrf2) involved in the transcriptional regulation of various genes contributed to antioxidant defense, including those encoding heme oxygenase-1, glutathione, superoxide dismutase and catalase, in driving hormesis-like adaptive responses has been also highlighted (Sekhar and Freeman 2015; Cameron et al. 2018). Nrf2 was shown to bind the antioxidant response element (ARE) and to activate some cytoprotective defense systems. For instance, acute irradiations with doses from 2 to 8 Gy have been shown to activate the ARE-dependent transcription in breast cancer cells (McDonald et al. 2010).

Mitohormesis

During last decades, important roles of ROS in multiple normal physiological processes have been revealed (Huang and Li 2020). They also play an essential role in various biosynthetic processes including the thyroid hormone production and cross-linking of extracellular matrix. Therefore, an excessively decreased ROS generation level may be likely associated with different pathological conditions. Impaired ROS cascade was found to be associated with hypothyroidism, low blood pressure, and also impaired antimicrobial defense (Milkovic et al. 2019). There is also increasing experimental evidence that caloric (dietary) restriction and several other experimental manipulations may extend lifespan via hormetic effects caused by increased production of ROS (mitohormesis) (Bárcena et al. 2018). Mitohormesis is a biological response where the induction of a mild mitochondrial stress by moderate ROS overproduction triggers certain signaling pathways, leading thereby to health benefits at a whole-organism level (Ristow 2014). The activation of the mitohormetic response was shown to be able to improve metabolism and immune system as well as to enhance longevity in various animal models, from worms to rodents (Bárcena et al. 2018). It is therefore suggested that LDIR can induce the body’s defense mechanisms, including the enhanced antioxidant activity, in consequence of a slightly enhanced concentration of ROS. Therefore, LDIR is regarded by several authors as a reasonable alternative to treatments with exogenous antioxidant sources, which were repeatedly reported to be inefficient in clinical trials (Doss 2012). One plausible explanation for the failure of these clinical trials is that exogenous antioxidants might not be effectively delivered to target organs. LDIR, in
contrast, may increase the capability of endogenous antioxidant systems in these organs. Therefore, LDIR may represent a promising therapy in treating pathological conditions caused by the oxidative damage of vital biomolecules (Doss 2012).

Heat shock response

The role of the heat shock response (coordinated induction of molecular chaperones such as heat shock proteins, HSPs) in producing hormetic effects has been elucidated (Lagisz et al. 2013; Dattilo 2015). Chaperone proteins are known to regulate both initial folding of proteins and subsequent maintenance of their structure. More specifically, this protein network is crucially involved in the de novo folding, refolding, and disaggregating damaged proteins. In a lot of cases, this network may refold damaged proteins to their normal functional state (Klaips et al. 2018). Integrated chaperone and protein-degradation networks are required to properly maintain a protein homeostasis (proteostasis). Furthermore, extracellular HSPs may stimulate an immune response. Therefore, their activation is considered as a promising treatment strategy to enhance anti-cancer immunity (Das et al. 2019; Yun et al. 2019). An important point in this regard is that capacity of the proteostasis network declines with age. The age-associated decline in the capacity of proteostasis may likely explain the increased prevalence of chronic disorders related to protein aggregation (e.g., neurodegenerative ones) in elderly (Gandhi et al. 2019; Webster et al. 2019). Production of HSPs represents a general response to any extreme stresses, including exposure to a high-dose radiation. Some studies indicated that expression of HSPs may be increased by LDIR exposure well. For example, in the myeloid leukemia cell line, LDIR was shown to induce expression of the HSP70 mRNA (Ibuki et al. 1998). Irradiated cells also exhibited thermo-resistance within one hour after irradiation and radioreistance within four hours after irradiation. Increased expression of Hsp25 and Hsp70 was found to be at least partly responsible for the induction of the adaptive response (a reduction of damaging effects caused by the high-level challenge dose through the low-dose priming exposure) in fibrosarcoma cells (Lee et al. 2002).

Apoptosis and autophagy

Low-dose/low-dose rate radiation has been consistently demonstrated to induce apoptosis (programmed cell death). Substantially enhanced rate of apoptosis was found, e.g., in HeLa and Hep2 cells following the low-dose/low-dose-rate radiation exposure (Mirzaie-Joniani and Eriksson 2002). It has been assumed by some authors that removal of damaged or senescent cells by apoptosis followed by compensatory cell proliferation may be a key mechanism in the radiation-induced adaptive responses (Kondo 1988). The radiation-induced generation of ROS may also cause of cellular “self-eating” involving degradation of various cellular components via lysosomal machinery, a process commonly referred to as autophagy (Szumiel 2012). Autophagy has been repeatedly found to be deregulated in most chronic pathological conditions, particularly age-related ones (Ren and Zhang 2018). Currently, autophagy induction is regarded as one of the most promising strategies in anti-aging medicine (Stead et al. 2019).

It should be certainly taken into account in discussing this subject that functional interactions of autophagy and apoptosis are quite complex. Indeed, autophagy enables adaptation to stress conditions known to suppress apoptosis; therefore, it represents an alternative pathway of cell death (Galluzzi et al. 2016). Autophagy and apoptosis, however, may be induced by the same stress conditions. Sometimes, these conditions result in combined autophagy and apoptosis; in other cases, cells switch between both these responses in a mutually exclusive way (Maiuri et al. 2007). Under the conditions of radiation-induced oxidative stress, the proper balance among normal metabolic functions of ROS and their damaging effects is likely a central factor in determining the cell’s fate (Szumiel 2012). Radiation-induced adaptive (hormetic) responses are often accompanied by stimulated cellular proliferation. In particular, LDIR has been found to significantly enhance the proliferation rate of mesenchymal stem cells in vitro through activation of the mitogen-activated protein kinase/extracellular-signal-regulated kinase (MAPK/ERK) signaling pathway known to play a pivotal role in cellular growth (Liang et al. 2011). Importantly, both early progenitor cells and hematopoietic stem cells were shown to be able to tolerate and adapt to being many times hit by the energy deposition events.
Based on this, the authors proposed the “injured stem cell hypothesis” according to that it is assumed that radiation-injured stem cells may continue to deliver functionally active cells which may ensure the maintenance of hematopoietic homeostasis throughout entire life course. It can be therefore assumed that post-radiation compensatory proliferation of such cell clones might be one more potential mechanism for both the radiation-induced adaptive responses and hormetic effects.

**Immune response**

The central suggestion made, usually implicitly, by the LNT model is that radiation carcinogenesis is due to “one-track action”, i.e., one or more DNA double strand breaks caused by a single electron track (Shah et al. 2012). The number of tracks is directly proportional to the dose. Cancer risk is therefore believed to be proportional to dose also, with any dose, no matter how small, able to induce cancer. This suggestion is, however, not consistent with multiple research findings indicating that mutation emergence is far from being the only causative event which triggers the development of clinical cancer. A considerable amount of evidence indicates that radiation may induce cancer not through simple stochastic events in a single cell but rather following complex systemic effects (Raabe 2011, 2014). Supportive evidence for this was provided from research performed in Japanese geriatric hospitals (Imaida et al. 1997). In this study, postmortem whole-body autopsies showed that percentage of patients carrying mutations indicative of cancer was about 40%, and this mutation rate has been found to be relatively stable in the age range from 48 to 94 years. Importantly, the incidence of cancer-indicative mutations did not differ substantially among age groups, as if the cancer incidence was not age-dependent. This result was counter-intuitive because within this age range the cancer mortality rate is known to increase by about an order of magnitude both in Japan and in Western countries. This discrepancy may indicate that presence of neoplastic cells is not the only (and likely not the decisive) factor in the cancer initiation and progression. Another notable fact here is that immune suppression leads to more than doubled risk of cancer development in organ-transplant patients (Doss 2012). This is quite predictable, given the fact that immune system plays a determining role in tumor immunogenicity (Koebel et al. 2007). Furthermore, the immune system can suppress cancer growth by extending the period required for the neoplasm development, maintaining cancer in an equilibrium state and preventing undetected (occult) tumors from becoming clinical cancers. Therefore, it has been assumed that age-associated increase in cancer incidence could be a consequence of the aging-related decline in immune function (Hong et al. 2019) rather than of occurrence of mutations per se (Doss 2012). It is therefore quite reasonable to expect that improvement of immune system may result in reducing cancer incidence. The responses of the immune system to irradiation have been demonstrated to be non-linear. High-dose radiation exposure is known to suppress the immune system, while LDIR may stimulate it. This is evident from multiple experimental studies (Hosoi and Sakamoto, 1993; Cui et al. 2017; Csaba 2019). The therapeutic potential of LDIR has been actively investigated for decades in various animal models of immune-associated pathological conditions, including autoimmune disorders and malignant tumors (for review, see Cui 2017).

The role of the adaptive immune response in carcinogenesis is also evident from studies where sporadic inhibition of distant untreated tumors following the irradiation of another part of the body was reported. Such inhibition is generally referred to as abscopal effect (a radiobiology term meaning “away from target”) (Dagoglu et al. 2019; Wani et al. 2019; Welsh et al. 2020). The abscopal phenomenon is commonly assumed to have an immune origin and it indicates that local radiotherapy may cause whole-body systemic effects (Yilmaz et al. 2019). Since the immune system may be stimulated by LDIR, many authors suggest that abscopal effects most probably originate from adaptive immune response in body parts exposed to indirect LDIR in the process of the (high-dose) radiation therapy of distant parts of the body. In addition, there is some evidence that low dose total-body irradiation (without the standard localized high-dose radiotherapy) can lead to suppression of distant metastasis of cancerous cells (Welsh et al. 2020). Based on these considerations, it has been proposed to apply LDIR to induce abscopal effect with aim of inhibiting cancerous processes in those patients who were diagnosed with early-stage cancers during screening programs (Doss 2013).
Medical treatments that employ low doses of ionizing radiation

Elucidating the biological mechanisms underlying radiation hormesis led to scientific perception of its reality and revitalization of research interest in this phenomenon. Over the past 15 years, several clinics in Japan began to provide hormesis therapy in small rooms that replicate the conditions of a typical natural radon spa. Some patients with severe medical conditions requested hormesis therapy after accepted treatments did not provide a remedy. When a significant improvement has been observed, a case report was written.

A recent article by Cuttler (2020) reviewed the application of low doses of ionizing radiation (LDIR) in medical therapies, from the discovery of X-rays and radioactivity in 1895/6 until the present. It discussed the political barrier against such therapies that was created in 1956 by the recommendation of the U.S. National Academy of Sciences to employ the LNT model to assess the risk of radiation-induced mutations (and cancer). The profound fundamental and clinical studies of Sakamoto et al. (1997) on cancer control, employing low doses of radiation, were pointed out and the excellent review by Pollycove (2007) on the radiobiological basis of LDIR treatments in the prevention and therapy of cancer was highlighted. The article outlined 13 recent cases of low-dose treatments in Japan with X-rays and radon to patients with different types of stage IV cancer (prostate, breast, colorectal, uterine, lung and liver cell) and to patients with severe autoimmune diseases (ulcerative colitis, pemphigus, type-1 diabetes, rheumatoid arthritis).

In the U.S., a patient in hospice with severe Alzheimer’s disease (and her husband with Parkinson’s disease) were treated successfully using CT scans of the brain (Cuttler et al. 2016, 2017a, b, 2018a, b). Subsequently, a pilot study was completed on five patients with severe Alzheimer’s disease. It repeated the treatments described in the 2016 case report. The family members of four of the five patients observed significant improvements in cognition and behavior. (A manuscript on this clinical trial has been submitted to a medical journal.) In addition, four clinical trials are starting on low-dose radiotherapy of Alzheimer’s disease using 5 or 10 dose fractions of 2 Gy. Such exposures have been employed successfully to remove amyloid plaque in other areas of the body and produce anti-inflammatory effects in preclinical studies (Michael et al. 2019; Ceyzeriat et al. 2020; Kim et al. 2020a, 2020b). LDIR was also considered among the potential hormesis-based approaches in treating Parkinson’s disease (Calabrese et al. 2018a). The anti-Parkinson’s potential of such treatment modality was evident from findings from rodent models of Parkinson’s disease (Kojima et al. 1999; El-Ghazaly et al. 2015).

Radiotherapy (RT) has been shown to have a potential as an effective and safe alternative to pharmacological therapies in treating different inflammatory conditions such as bursitis, tendonitis, arthritis, and also serious inflammatory lung conditions (for review, see Calabrese et al. 2018b). In these studies conducted mostly during the 1920s to 1940s, RT was reported to be efficient after only a single exposure, with either short-term (within 24 h) or long-term (lasting months to years) relief of symptoms. The polarization of macrophages to an anti-inflammatory or M2 phenotype was assumed as potential mechanism by which RT can mitigate inflammation and facilitate healing (Calabrese et al. 2018b). These findings are particularly promising in the context of the COVID-19 viral pandemic, which started in December 2019 and has had an enormous social and economic impact. Although the general population is commonly susceptible to the COVID-19, infected elderly patients demonstrate most fast progression and severe manifestations with a high proportion of the critical condition owing to the compromised immunity and underlying disorders (Wang et al. 2020). Exceptional precautionary measures have been taken to control its high incidence of morbidity and mortality (Johns Hopkins University 2020), and intensive efforts have been underway to identify treatments and vaccines. Since the acute respiratory distress syndrome due to immune over-response is the deadliest symptom, the 1940s LDIR treatment such as 0.5 Gy X-ray of the lungs (Calabrese and Dhawan 2013) has emerged as a promising modality to treat COVID-19 (Algara et al. 2020; Cosset et al. 2020; Dhawan et al. 2020; Kirkby and Mackenzie 2020). Fourteen clinical trials are underway worldwide as for Oct 1, 2020 (U.S. National Library of Medicine 2020). Preliminary results (Del Castillo et al. 2020; Hess et al. 2020) are encouraging.
Conclusions

LDIR has been successfully exploited for centuries (radon springs) to remediate arthritis and other diseases. In the first half of XX century, LDIR was successfully used for treating a range of diseases including pneumonia. However for the last half a century, application guidelines for ionizing radiation (as well as safety regulations) have been based on the LNT model assuming that health risk is proportional to the total radiation dose, no matter how low the dose and dose rate are. While LNT has never achieved a scientific consensus, the recent epidemiological evidence is less and less compatible with the assumption of linearity. More importantly, much information was accumulated on biological effects of LDIR, and the recent radiobiological evidence does not support the assumption of low-dose harm. Moreover, there is increasing evidence that LDIR, such as used in X-ray imaging including CT, is a hormetin—that is it affects health rather positively than negatively. Not accidentally, LNT is viewed as doubtful (and often—obsolete) by a growing part of the research community, though there are still many important topics that require further investigation. In this review, we aimed to summarize evidence supporting hormesis through LDIR. We also discussed here the possibility of clinical use of LDIR, predominantly for age-related disorders, e.g., Alzheimer’s disease, for which no remedies are available (Socol et al. 2013). Indeed, due to the low average residual life expectancy in old patients, the short-term benefits of such interventions (e.g., potential therapeutic effect against dementia) may outweigh their hypothetical delayed risks (e.g., cancer).

When considering the effects of irradiation on human health, it is necessary to clearly distinguish between the effects of increased background radiation to which adaptation can occur over many generations at the population level and the effects of irradiation as a result of accidents or medical procedures. In addition, hormetic effects of acute and chronic radiation exposure and also of high- and low-linear energy transfer (LET) radiation conditions have to be separately considered.

Undoubtedly, caution should be exercised when introducing new medical practices, and LDIR therapy is no exception. However, bearing in mind the enormous social, economic and ethical implications of potentially-treatable, age-related disorders, we argue that assessment and clinical trials of LDIR treatments should be given priority.

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Compliance with ethical standards

Conflict of interest The authors declare no conflict of interest.

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