

Radioprotective effect of hydrogen in cultured cells and mice

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Abstract

It has been demonstrated that hydrogen can selectively reduce hydroxyl and peroxynitrite in vitro. Since most of the ionizing radiation-induced cellular damage is caused by hydroxyl radicals, this study was designed to test the hypothesis that hydrogen may be an effective radioprotective agent. In this paper, we demonstrate that treating cells with hydrogen before irradiation could significantly inhibit ionizing irradiation(IR)-induced Human Lymphocyte AHH-1 cells apoptosis and increase cells viability in vitro. We also show hydrogen can protect gastrointestinal endothelia from radiation-induced injury, decrease plasma malondialdehyde (MDA) 、 intestinal 8-hydroxydeoxyguanosine (8-OHdG) levels, and increase plasma endogenous antioxidants in vivo. We suggest that hydrogen has a potential as an effective and safe radioprotective agent.

Keywords: Ionizing radiation; radioprotection; intestinal; hydrogen

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Introduction

Exposure to ionizing radiation (IR) can produce severe health impairments due to injury and failure to susceptible organs. Detrimental effects of IR on biological tissues are, in major part, mediated via increased production of hydroxyl radical. Hydroxyl radical produced during radiolysis of water can trigger oxidation of lipids, amino acids, and saccharides leading to formation of various secondary free radicals [1-3]. These free radicals can produce severe health impairments due to injury and failure to susceptible cells and organs.

The gastrointestinal tract is one of the most susceptible organs to radiation^[4]. As low as 1 Gy of radiation induces dramatic increase in apoptosis in mouse small intestinal crypt within three to six hours after exposure, predominantly in the stem cell region^[5].

Ohsawa et al.^[6] found that molecular hydrogen could selectively reduce cytotoxic reactive oxygen species, such as $\bullet\text{OH}$ and ONOO^- in vitro and exert therapeutic antioxidant activity in a rat middle cerebral artery occlusion model. Therefore, we reasoned that hydrogen might be protective against detrimental effects of radiation. However, application of H_2 gas inhalation is not convenient and may be dangerous because it is inflammable and explosive. On the other hand, H_2 gas saturated PBS / saline, which is called hydrogen-rich PBS /saline, is easy to apply and safe. In the current study, we investigated whether administration of

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4 hydrogen-rich PBS/saline exerted radioprotective effect in vitro and in
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7 vivo. We demonstrated here that hydrogen treatment could protect
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10 cultured lymphocytes and gastrointestinal tract from γ -radiation in mice.

11 **Materials and methods**

12 *Hydrogen-rich PBS/saline production*

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18 Hydrogen was dissolved in PBS/physiological saline 6 hours under high
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20 pressure (0.4 MPa) to a supersaturated level using hydrogen-rich
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23 water-producing apparatus produced by our department. The saturated
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26 hydrogen PBS/saline was stored under atmospheric pressure at 4 °C in an
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29 aluminum bag with no dead volume. Hydrogen-rich PBS/saline was
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32 freshly prepared every week, which ensured that a concentration of more
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35 than 0.6mmol/L was maintained. Gas chromatography (Biogas Analyzer
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38 Systems-1000, Mitleben, Japan) was used to confirm the content of
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41 hydrogen in PBS/saline by the method described by Ohsawa et al^[6].

42 *Cell culture and treatment*

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45 Human Lymphocyte AHH-1 cells(American Type Culture Collection,
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48 Manassas, VA, USA) were maintained in RPMI 1640 (Invitrogen,
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51 California, USA) with 10% fetal bovine serum and 1% penicillin -
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54 streptomycin - glutamine at 37°C in a 5% CO₂ humidified chamber. For
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57 radioprotective studies, cells were treated with different volume of
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60 Hydrogen-rich PBS and accordingly we added different volume of PBS
in order to obtain the desired concentration of H₂ and make the final

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4 volume of the medium the same, then the treated cells were immediately
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6 irradiated with different doses of γ -ray ,depending upon the requirement
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8 of the present study. After irradiation, the cells were centrifuged and
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10 cultured in RPMI 1640.
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13 14 15 *Irradiation*

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17 ⁶⁰Co-gamma rays in irradiation Center (Faculty of Naval Medicine,
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19 Second Military Medical University, China) were used for the irradiation
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21 purpose. Mice(with or without hydrogen pre-treatment) were exposed to
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23 different doses of radiation, depending upon the requirement of the
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25 present study.
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30 31 32 *Cell Viability Analyses*

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34 Human Lymphocyte AHH-1 cells were seeded in 96-well plates and
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36 pretreated with or without Hydrogen-rich PBS, the treated cells were then
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38 immediately irradiated. After irradiation the cells were further cultured
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40 for 48 h. Cell viability was determined by WST assay using a Cell
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42 Counting kit (Dojindo Laboratories, Kumamoto, Japan).
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46 47 48 *Lactate Dehydrogenase (LDH) leakage assay*

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50 LDH leakage assay was carried out using LDH cytotoxicity detection
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52 kit(Nanjing KeyGen Biotech. Co. Ltd. China)according to protocol in the
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54 user's manual. AHH-1cells were pretreated with Hydrogen -rich PBS and
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56 the final concentration of H₂ was maintained above 0.3mmol/L.
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58 Immediately, the cells were exposed under gamma radiation and then
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4 transported to an ice bucket. After 4 hours time period we analysed the
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6 content of LDH in cell suspension.
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9 10 *Apoptosis assays for cultured cells*

11 Apoptosis was determined by Annexin V-APC and propidium iodide
12 staining using Apoptosis Detection Kit (Bipece Biopharma,
13 Massachusetts,USA). Treated cells were incubated with Annexin V-APC
14 for 15 minutes at 4°C and propidium iodide for 5 minutes at room
15 temperature. Cells were then analyzed by flow cytometry. Alternatively,
16 apoptosis was determined by Hoechst33258, fluorescein diacetate(FDA)
17 and propidium iodide staining. Treated cells were washed with PBS twice,
18 and then stained with 40mg/L fluorescein diacetate, 20 mg/L
19 Hoechst33258 at room temperature for 15min, and stained with 20mg/L
20 propidine iodine at room temperature for 5min. The cellular morphology
21 was observed using Olympus BX60 fluorescent microscope equipped
22 with Retiga 2000R digital camera. Average percentage of apoptotic cells
23 was calculated in 5-7 randomly selected high power field (HPF).
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46 *Mice and treatment*

47 All the protocols were approved by the Second Military Medical
48 University, China in accordance with the Guide for Care and Use of
49 Laboratory Animals published by the US NIH (publication No. 96-01).
50 Male BALB/c rats weighing 21-23 g were used in the experiments. The
51 animals were housed in individual cages in a temperature-controlled
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4 room with a 12 h light/dark cycle and food and water were provided ad
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7 libitum. For experiments, mice were treated intraperitoneally (IP) with
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10 physiological saline or Hydrogen-rich saline 20 min before radiation. And
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12 mice were irradiated in a holder designed to immobilize unanaesthetized
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15 mice such that the abdomens were presented to the beam.

16 17 *Morphologic observation*

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20 Mice were treated intraperitoneally (IP) with physiological saline or
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23 Hydrogen-rich saline 20 min before irradiation. Twelve hours after
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26 irradiation, mice were sacrificed by cervical dislocation under isoflurane
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29 anesthesia. A 5 cm segment of small intestine which was removed from at
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32 5 cm proximal to the terminal ileum was fixed in 10% buffered
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35 formaldehyde-saline. Three 1 cm segments of intestinal specimen were
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38 embedded in paraffin and stained with hematoxylin and eosin.
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41 Morphological damages were assessed by Chiu histological injury
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44 scoring system of intestinal villi (0 = normal mucosa, 1=slight-, 2 =
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47 moderate-, 3 = massive subepithelial detachments, 4 = denudes villi,
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50 5=ulceration)^[7]. Two independent and blinded researchers performed the
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53 histological scoring.

54 55 *Biochemical assays*

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58 Arterial blood samples (0.6 ml) of mice were collected 12 h after
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irradiation. These samples were immediately centrifuged at 2500 rpm and
4 °C for 10 min. The plasma was taken for biochemical estimations(SOD,

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4 GSH and MDA). Superoxide dismutase(SOD) activity was assayed by
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6 the method of Kakkar et al. ^[8], based on the inhibition of the formation of
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8 NADH-PMS-NBT complex. The GSH concentration was measured by
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10 the method of Ellman ^[9]. This method was based on the development of a
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12 yellow color when 5',5'-dithiobis 2-nitrobenzoic acid was added to
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14 compounds containing sulfhydryl groups. MDA was assessed
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16 spectrophotometrically with the method defined by Ohkawa et al as MDA
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18 reacted with thiobarbituric acid and formed a pink, maximum absorbent
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20 complex at 532 nm wavelength^[10].
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28 For determination of 8-OHdG levels in DNA from the intestine of mice,
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30 DNA was extracted from the mice intestinal specimen with a DNA
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32 Extractor Kit (DNA Extractor Wb Kit, Wako Chemical; Osaka, Japan)
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34 according to the method of Nakae et al^[11]. Then the isolated DNA was
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36 digested by the method of Victoria et al^[12]. The 8-OHdG levels of these
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38 samples were measured as described by Teruo et al^[13]. Briefly, the
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40 samples were added to plate wells precoated with mouse monoclonal
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42 anti-8-OHdG antibody (Japan Institute for the Control of Aging, Fukuroi,
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44 Japan), of which the specificity has been proved by S.Toyokuni et al^[14].
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46 They were incubating for 45 min at 37°C. After washed for 3 times, the
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48 wells were sequentially treated with Biotinylated rabbit-anti-mouse IgG
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50 for 30 min at 37°C and Streptavidin-Horseradish Peroxidase(HRP)for 30
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52 min at 37°C. A substrate containing 3,3',5,5'- tetramethylbenzidine(TMB)
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4 was added and the wells were incubated for 15 min at 37°C . The reaction
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7 was terminated by the addition of a sulphuric acid. The absorbance was
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10 read at a wavelength of 450nm.

11 **Statistical analysis**

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14 Data are expressed as means± S.E.M. for each experiment. The number
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17 of samples is indicated in the description of each experiment. Statistical
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20 analysis was performed by using One Way Analysis of Variance. Between
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23 groups, variance was determined using the Student-Newman–Keuls post
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26 hoc test. A P value of less than 0.05 was considered to be statistically
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29 significant.

30 **Results**

31 *Hydrogen -rich PBS increases cell viability of irradiated AHH-1 cells*

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34 To study radioprotective effects of H₂ in cell culture, we examined
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37 viability of irradiated AHH-1 cells. Cells treated with or without different
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40 concentrations of H₂ were exposed under 4Gy of γ -radiation as described
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43 in Methods. We demonstrated that pretreatment of AHH-1 cells with
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46 0.1-0.4 mmol/L H₂ before irradiation significantly increased cell survival
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49 as compared to cells treated with radiation alone at all examined doses(up
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52 to 8 Gy) (Fig.1A and C).And as shown in the figure, the radioprotective
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55 effect of H₂ is dose dependent. However, as we treated cells with
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58 Hydrogen -rich PBS after irradiation, the protective effect is not
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significant(Fig.1B) .

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4 *Hydrogen -rich PBS decrease cellular Lactate Dehydrogenase (LDH)*

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7 *leakage in irradiated cells*

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10 Besides the cell viability, we also determined LDH activities to estimate
11 cellular LDH leakage from damaged cells. The result indicated that
12 pretreatment with 0.3mmol/L H₂ before irradiation significantly
13 decreased LDH leakage of AHH-1 cells which were exposed under
14 different doses of γ -radiation (Fig. 2). And this result was consistent
15 with the result obtained by cell viability observation.
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25 *Hydrogen -rich PBS attenuates apoptosis in irradiated AHH-1 cells*

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28 To determine the radiation-induced apoptosis of irradiated AHH-1 cells,
29 we analyzed treated cells by using Annexin V-APC and propidium iodide
30 staining in flow cytometry assay. The early apoptotic cells decreased
31 when pretreated with 0.4mmol/L H₂ as compared to cells pretreated
32 without H₂ (Fig. 3A and B, 10.2% vs.21.5%, respectively). We further
33 evaluated the morphology of dying cells using Hoechst33258, fluorescein
34 diacetate and propidium iodide staining. Irradiated AHH-1 cells
35 pretreated with Hydrogen -rich PBS demonstrated a protective effect with
36 reduced number of apoptotic cells to 26.1% as compared to 49.3 % in
37 PBS-pretreated irradiated cells (Fig. 3CandD). These data suggest that H₂
38 can attenuate apoptosis in irradiated AHH-1 cells.
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57 *Hydrogen-rich saline treatment attenuates intestinal injury in vivo*

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60 We observed histological IR injuries featured by shortening of the villi,

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4 loss of villous epithelium, and prominent mucosal neutrophil infiltration
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6 (Fig.4A). All of these changes were ameliorated by administration of
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8 hydrogen-rich saline. Chiu scoring and microphotographs are shown in
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10 Fig.4B. As shown in the figure, hydrogen-rich saline administration
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12 significantly reduced the mucosal injury caused by IR.
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16 17 *Changes in the activities of plasma SOD and GSH*

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19 The plasma SOD and GSH concentrations were measured at 12 h of
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21 irradiation (Fig.5A and B).Plasma SOD and GSH concentrations at 12 h
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23 of irradiation in the H₂ group were significantly higher than that of the
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25 Control group.
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29 30 *Changes in the levels of plasma MDA and intestinal 8-OHdG*

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32 The plasma MDA and intestinal 8-OHdG concentrations were measured
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34 at 12 h of irradiation (Fig. 6Aand B).Plasma MDA and intestinal 8-OHdG
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36 concentrations at 12 h of irradiation in the H₂ group were significantly
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38 lower than that of the Control group.
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43 44 **Discussion**

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46 To our knowledge, this is the first study demonstrating that hydrogen has
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48 radioprotective effects in vitro and vivo. In several recent studies, H₂
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50 inhalation was reported to protect cerebral^[6], myocardial^[15], and hepatic
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52 ^[16] I/R injury in animal models. In addition, Buchholz et al^[17] reported
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54 that hydrogen inhalation ameliorates oxidative stress in transplantation
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56 induced intestinal graft injury. This radioprotective effect may result from
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4 radical oxygen species (ROS) scavenging effect of molecular H₂, as
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6 previously reported in a brain injury model^[6]. The effect of free radical
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8 scavengers to ameliorate the oxidative injuries due to ionizing radiation
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10 has been considerably reported ^[18-19]. Radical oxygen species O₂⁻ and
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12 H₂O₂ are detoxified by antioxidant defense enzymes, whereas •OH and
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14 ONOO⁻ could not be detoxified by antioxidant defense enzyme. It has
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16 been demonstrated that hydrogen gas selectively reduces •OH and
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18 ONOO⁻ ^[6]. The hydroxyl radical is the most reactive product of reactive
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20 oxygen species generated in cells. Hydroxyl radicals can easily react with
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22 cellular macromolecules, including DNA, proteins and lipids, to exert a
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24 strong cytotoxic effect. Since most of the ionizing radiation-induced
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26 damage is caused by hydroxyl radicals, we speculate that the
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28 radioprotective effect may result from its radical oxygen species (ROS)
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30 scavenging effect.
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34 Endogenous antioxidants are a group of substances that significantly
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36 inhibit or delay oxidative processes while being oxidized themselves ^[20].
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40 Antioxidant enzymes are important in providing protection from radiation
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42 exposure^[21]. And glutathione (GSH) participates non-enzymatically in
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44 protection against radiation damage^[22]. Therefore, a reduction in the
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46 activity of these substances can result in a number of deleterious effects.
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50 Membrane lipids are the major targets of ROS and the free radical chain
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52 reaction^[23]. The increase in the levels of lipid peroxidation products such
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4 as malondialdehyde and TBARs, are the indices of lipid damage^[24]. Also
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6 DNA is one of the major targets of ROS and 8-OHdG is formed from
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8 deoxyguanosine in DNA by hydroxyl free radicals^[25]. In our study, we
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10 observed a significant decrease in the levels of enzymatic antioxidant
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12 (SOD), non-enzymatic antioxidant (GSH) and an increase in the levels of
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14 plasma MDA and intestinal 8-OHdG of irradiated mice. But pretreatment
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16 of hydrogen prior to radiation exposure increased the antioxidant status at
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18 both enzymic and non-enzymic levels and decreased the levels of MDA
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20 and 8-OHdG.
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28 The mechanism of protection on SOD and GSH is most likely based on
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30 the ability of H₂ to effectively inhibit oxidative reactions. It has been
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32 demonstrated that O₂⁻ can undergo either spontaneous or
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34 enzyme-catalyzed (SOD) dismutation to hydrogen peroxide (H₂O₂), or
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36 can react with nitric oxide (NO •) to form the toxic product peroxynitrite
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38 (ONOO⁻)^[26]. Hydrogen gas can reduce ONOO⁻^[6]. Therefore, it's possible
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40 H₂ can accelerate the reaction between O₂⁻ and NO •. More O₂⁻ would
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42 react with NO • while the enzyme-catalyzed reaction by SOD would be
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44 weakened. Thus the SOD can be protected. Besides, hydroxyl radicals can
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46 react with themselves to form H₂O₂, which could oxidize GSH to
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48 GSSG^[1]. Hydrogen gas can reduce hydroxyl radical^[6], thus probably
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50 hydrogen gas could partly reduce the formation of H₂O₂ in the reaction
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52 chain which may lead to the protection of GSH. We may conclude that
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4 the protection in the antioxidant status during H₂ pretreatment has further
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6 decreased the attack of free radicals, prevented DNA damage, and
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8 decreased lipid peroxidation, thereby decreased the deleterious effects of
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10 radiation.
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14 Some radioprotectors, such as thiol compounds, has relatively high
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16 toxicity ^[27], while cytokines and immunomodulators, should be used with
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18 low radiation doses or in combination with radical scavengers and
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20 antioxidants ^[28], and natural antioxidants, such as vitamin E, flavonoids
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22 and others, have fewer toxic side effects but also a lower degree of
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24 protection compared to thiol agents ^[27]. The sulfhydryl compound
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26 amifostine, named WR-2721, which is the only radioprotectant registered
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28 for using in human, has shown good radioprotective effects ^[29]. But it has
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30 many side effects limiting its clinical use such as hypertension, nausea,
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32 vomiting, and other side effects ^[30]. However, hydrogen is continuously
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34 produced by colonic bacteria in the body and normally circulates in the
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36 blood ^[31], so it is physiologically safe for humans to inhale hydrogen at a
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38 relatively low concentration. It is also a highly diffusible gas and reacts
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40 with hydroxyl radical to produce water ^[32]. Dissolving H₂ in solvent such
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42 as PBS, physiological saline or medium is easy to apply and safe.
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44 Therefore, it may have great potential for clinical use.
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58 In conclusion, the effect of reducing radical oxygen species plays an
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60 important role in the radioprotective effects of hydrogen. However, the

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4 exact mechanism and signaling pathway involved in the protection role of
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7 hydrogen in ionizing radiation injury need to be studied in the future.
8

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Figure legends

Figure1. Dose dependent effect of H₂ on cell viability induced by 4Gy gamma radiation(A).Pretreatment of 0.4mmol/L H₂ before irradiation can increase cell survival(B). Variation of cell survival percent pretreated with0.4 mmol/L H₂ before different dose of irradiation(C). Values are given as mean \pm SEM ((n=6,neg: no H₂ no radiation). *P< 0.05,**P< 0.01, # P<0.1.

Figure2. Changes in the levels of LDH in normal, γ -irradiated and H₂ pretreated lymphocytes. Values are given as mean \pm SEM (n=4). *P< 0.01

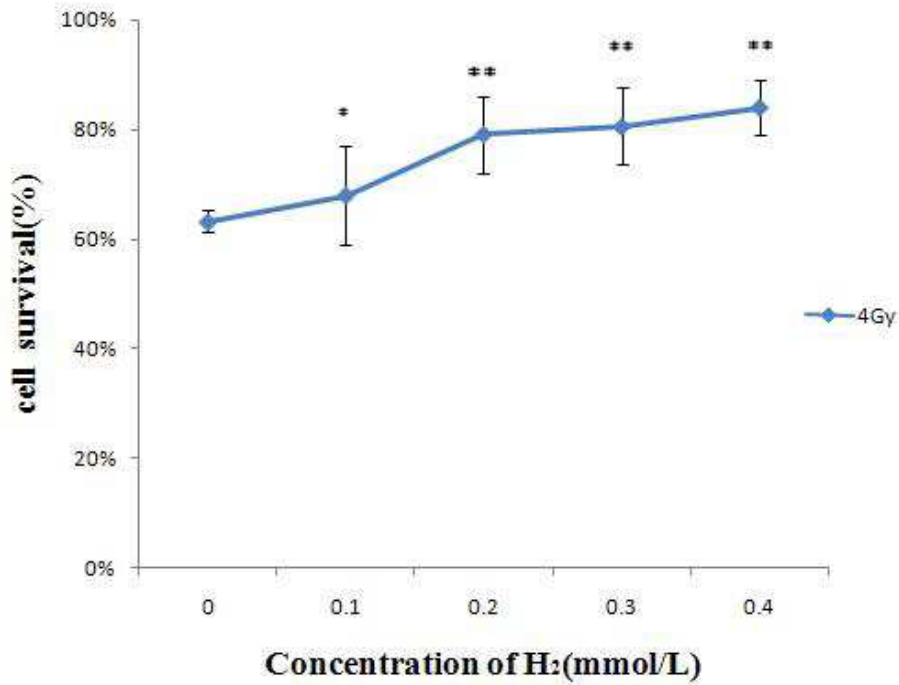
Figure3. Hydrogen-rich PBS attenuates radiation-induced apoptosis in AHH-1 cells. Treated cells were collected 24 h after irradiation, stained with Annexin V-APC and propidium iodide and analyzed by flow cytometry. Shown are representative diagrams of distribution of stained cells (A) and a bar graph of apoptotic cells expressed as a percent of total cells .Values are given as mean \pm SEM (n=4). *P<0.01 (B). Cells were stained with FDA、Hoechst33258 and PI 24 h after irradiation and apoptotic cells were counted in multiple randomly selected fields. Shown are representative micrographs (C) and a bar graph of apoptotic cells expressed as a percent of total cells .Values are given as mean \pm SEM (n=4) *P<0.01 (D).

Figure4.Morphologic observation of the intestinal tissue in normal, γ -irradiated and H₂ pretreated mice. Photomicrographs of the intestinal

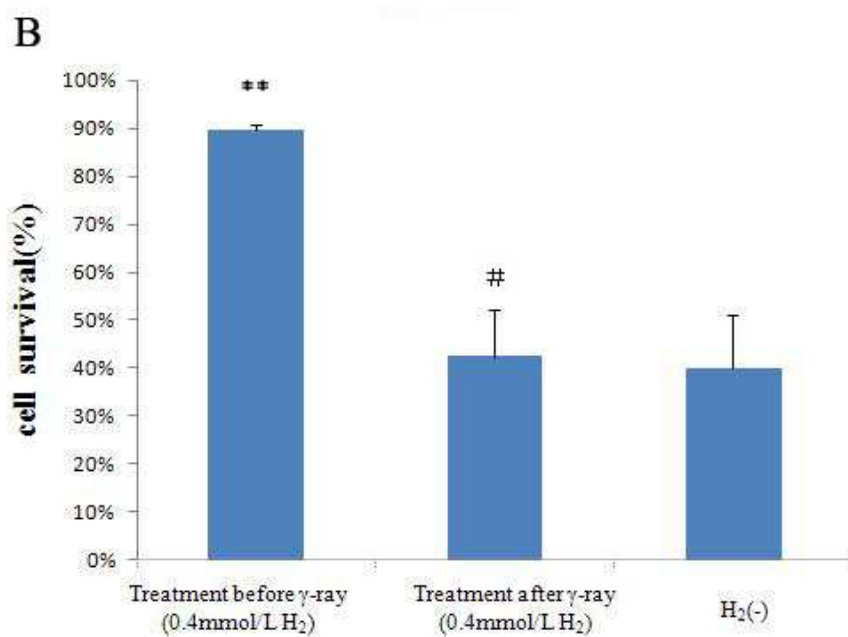
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4 tissue stained by the hematoxylin and eosin(A). Intestinal mucosal injury
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6 evaluated by Chiu scoring system(B). Grading as (0 = normal mucosa, 1
7 =slight-, 2 = moderate-, 3 = massive subepithelial detachments, 4 =
8 denudes villi, 5 = ulceration). Data are expressed as means \pm SEM for at
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10 least triplicate independent experiments. (n =8 per group). *P < 0.01.
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17 **Figure5.** Changes in the activities of SOD and concentrations of GSH in
18 normal, γ -irradiated and H₂ pretreated mice. Values are given as mean \pm
19 SEM (n=4). *P < 0.01.
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26 **Figure6.** Hydrogen-rich saline significantly decreased levels of MDA, a
27 marker of oxidative stress(A).And oxidative DNA damage was assessed
28 by 8-OHdG immunoreactivity. Shown are 12 hours after irradiation,
29 intestinal 8-OHdG concentrations in normal, γ -irradiated and H₂
30 pretreated groups(B). Relative to the Control, H₂ significantly decreased
31 the concentration of 8-OHdG. Values are mean \pm SEM (n=6), *P<0.01.
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198x127mm (72 x 72 DPI)

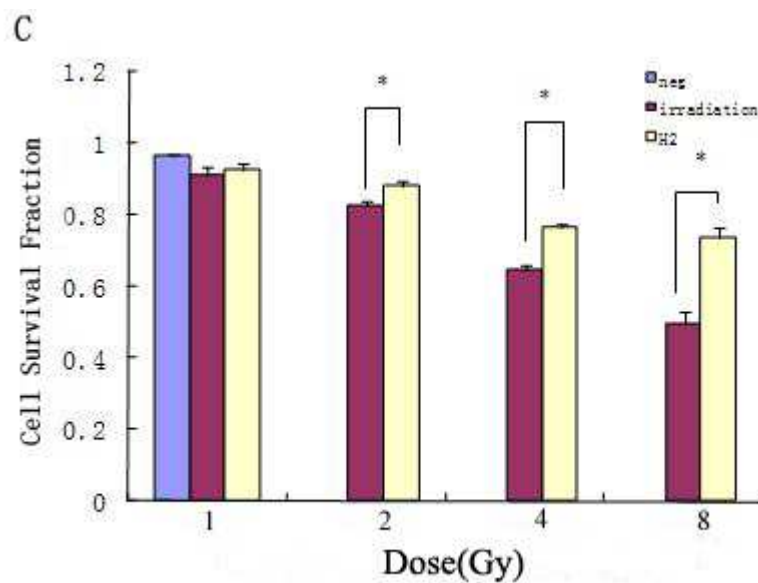


Figure1. Dose dependent effect of H2 on cell viability induced by 4Gy gamma radiation(A). Pretreatment of 0.4mmol/L H2 before irradiation can increase cell survival(B). Variation of cell survival percent pretreated with 0.4 mmol/L H2 before different dose of irradiation(C). Values are given as mean \pm SEM ((n=6, neg: no H2 no radiation). *P< 0.05, **P< 0.01, # P<0.1.
164x107mm (72 x 72 DPI)

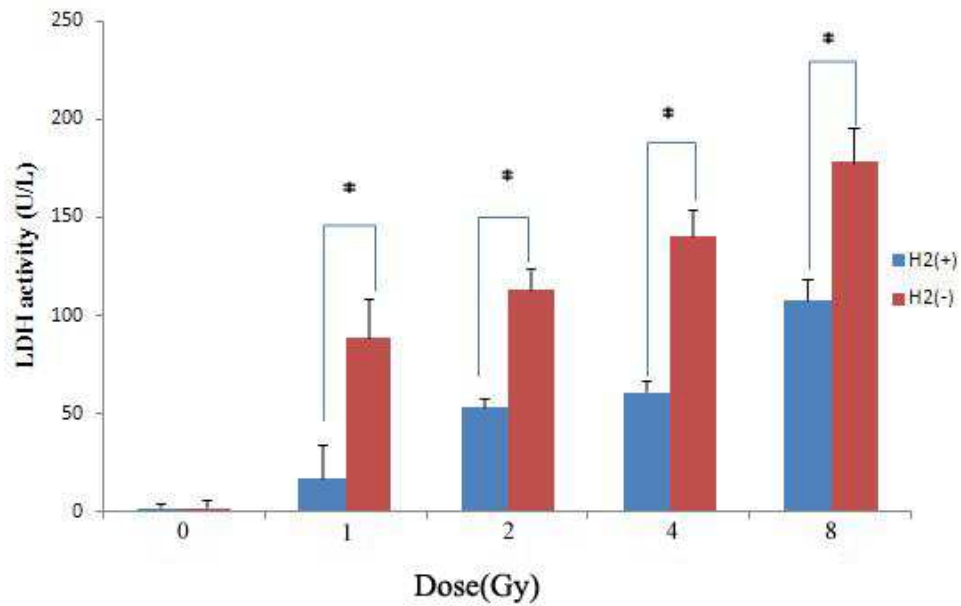
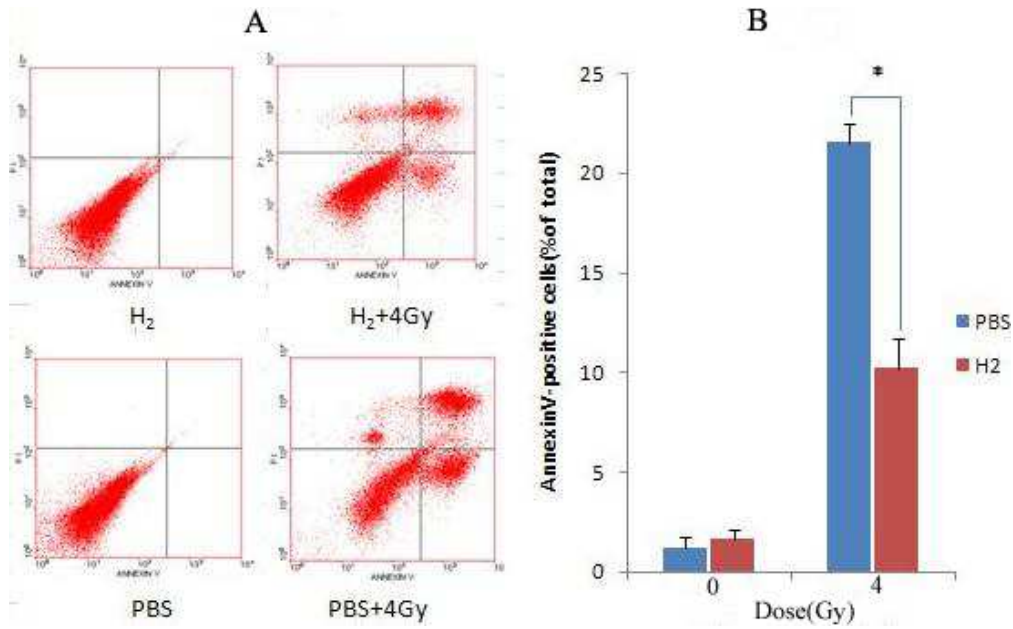


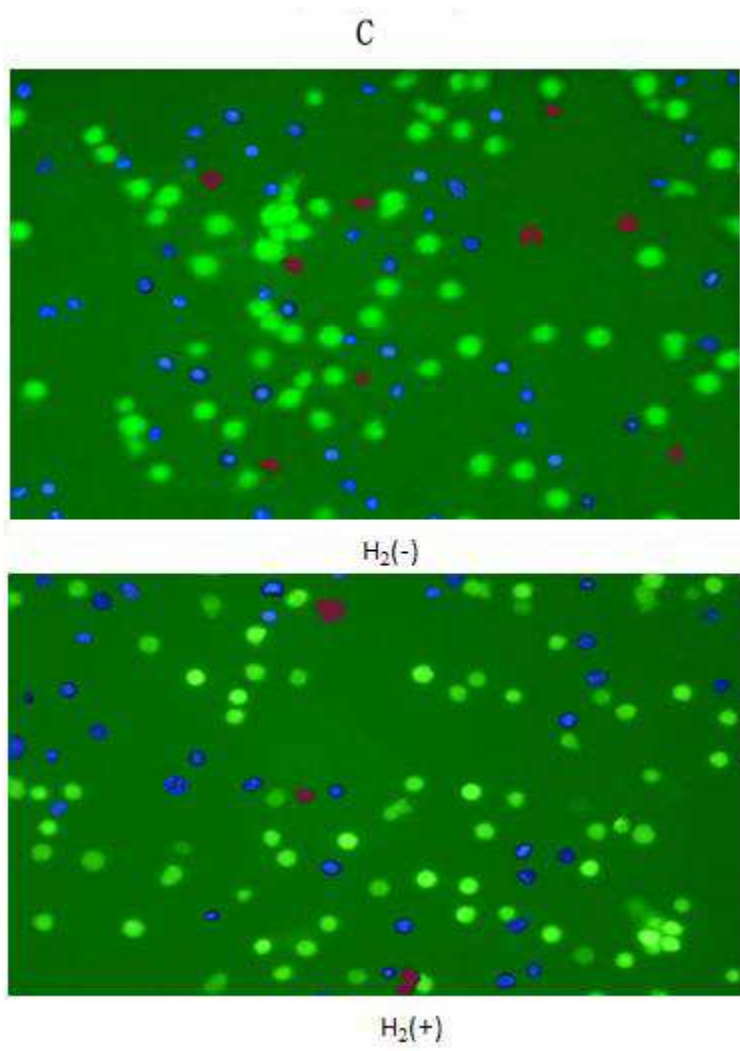
Figure2. Changes in the levels of LDH in normal, γ -irradiated and H2 pretreated lymphocytes. Values are given as mean \pm SEM (n=4). *P< 0.01
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194x119mm (72 x 72 DPI)

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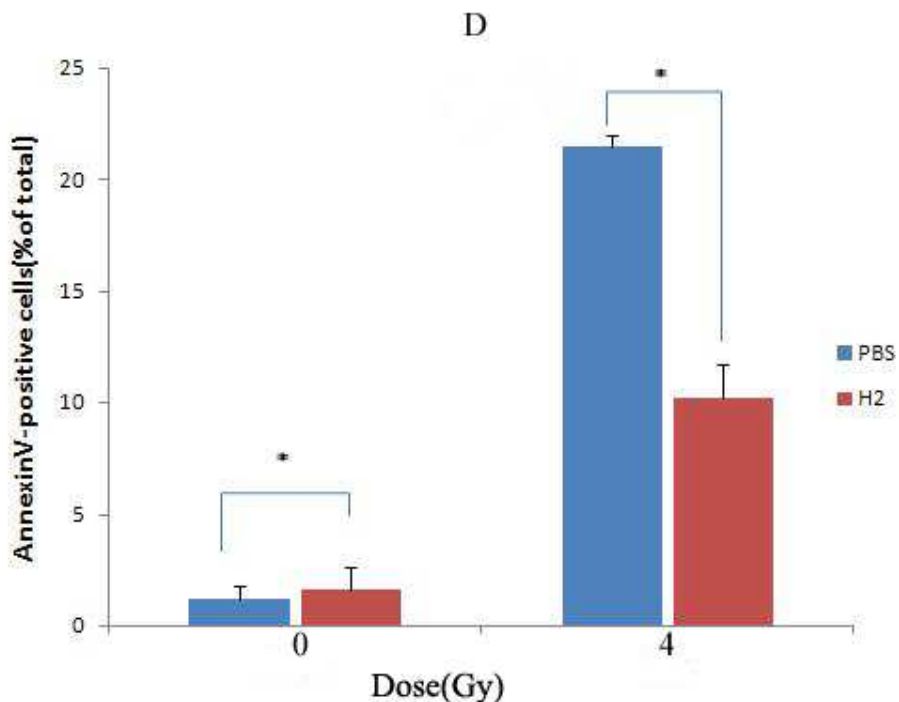
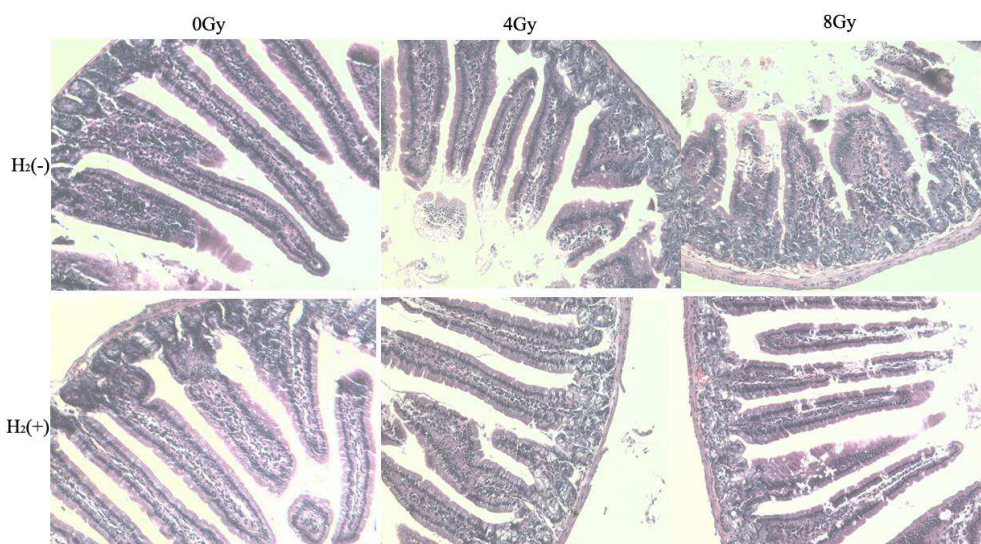


Figure 3. Hydrogen-rich PBS attenuates radiation-induced apoptosis in AHH-1 cells. Treated cells were collected 24 h after irradiation, stained with Annexin V-APC and propidium iodide and analyzed by flow cytometry. Shown are representative diagrams of distribution of stained cells (A) and a bar graph of apoptotic cells expressed as a percent of total cells. Values are given as mean \pm SEM (n=4). *P<0.01 (B). Cells were stained with FDA, Hoechst33258 and PI 24 h after irradiation and apoptotic cells were counted in multiple randomly selected fields. Shown are representative micrographs (C) and a bar graph of apoptotic cells expressed as a percent of total cells. Values are given as mean \pm SEM (n=4) *P<0.01 (D).
188x135mm (72 x 72 DPI)

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329x182mm (96 x 96 DPI)

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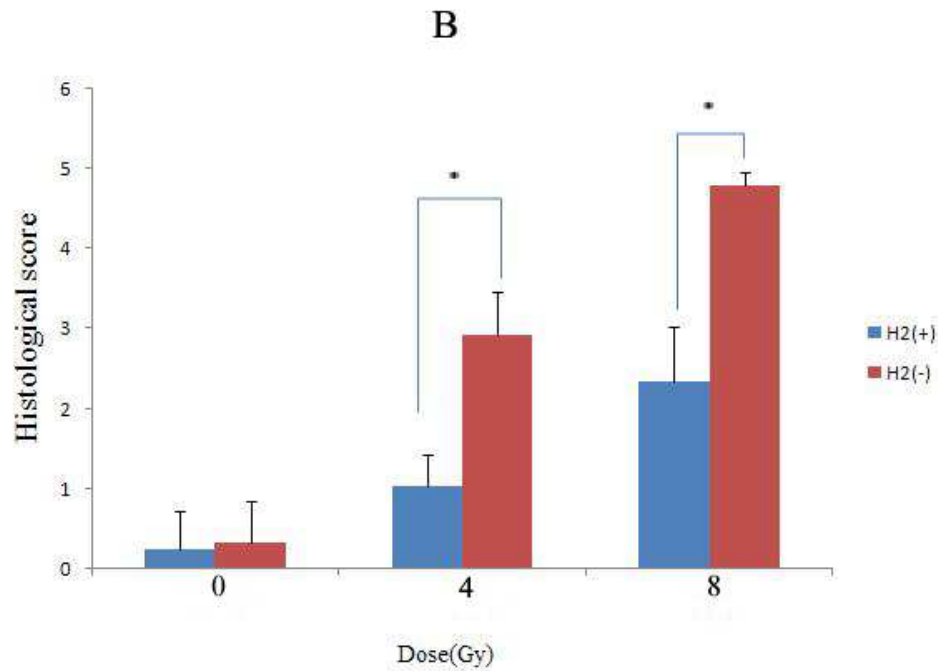
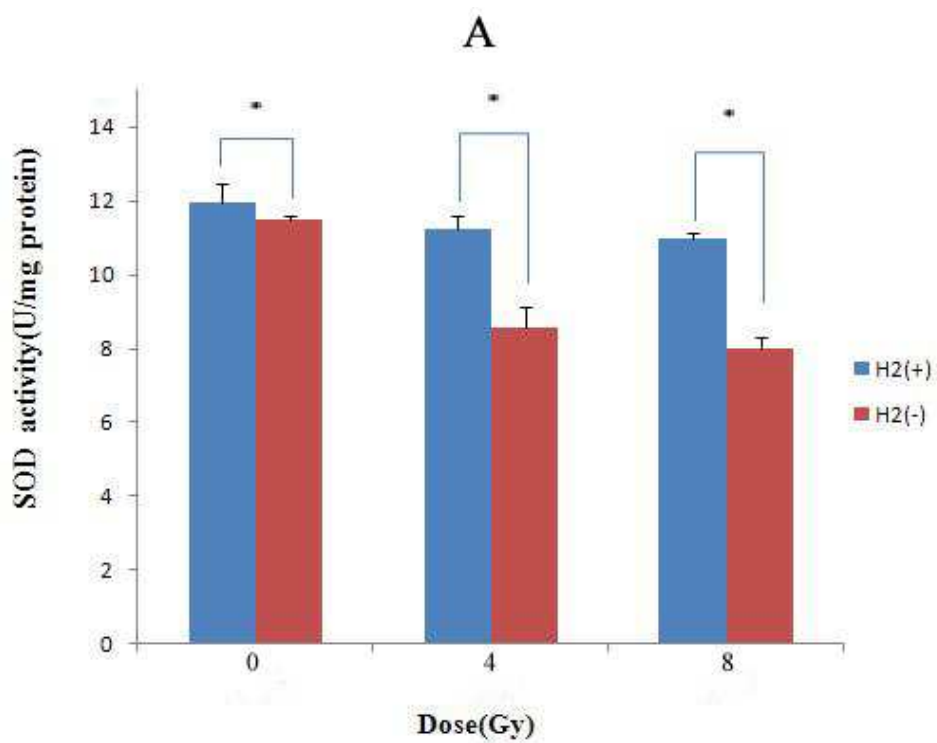


Figure 4. Morphologic observation of the intestinal tissue in normal, γ -irradiated and H2 pretreated mice. Photomicrographs of the intestinal tissue stained by the hematoxylin and eosin (A). Intestinal mucosal injury evaluated by Chiu scoring system (B). Grading as (0 = normal mucosa, 1 = slight-, 2 = moderate-, 3 = massive subepithelial detachments, 4 = denudes villi, 5 = ulceration). Data are expressed as means \pm SEM for at least triplicate independent experiments. (n = 8 per group). *P < 0.01.

212x148mm (72 x 72 DPI)



185x140mm (72 x 72 DPI)

new Only

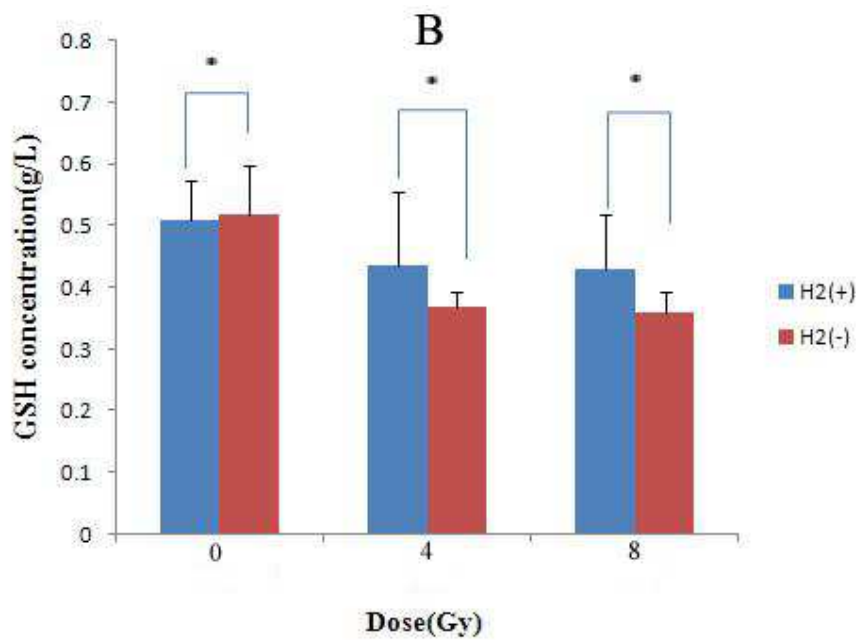
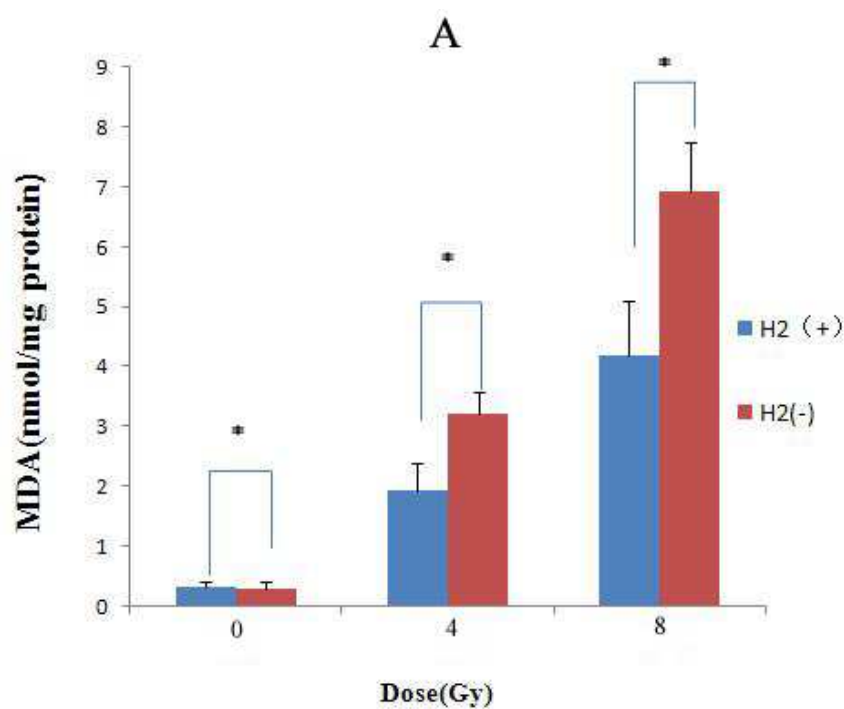
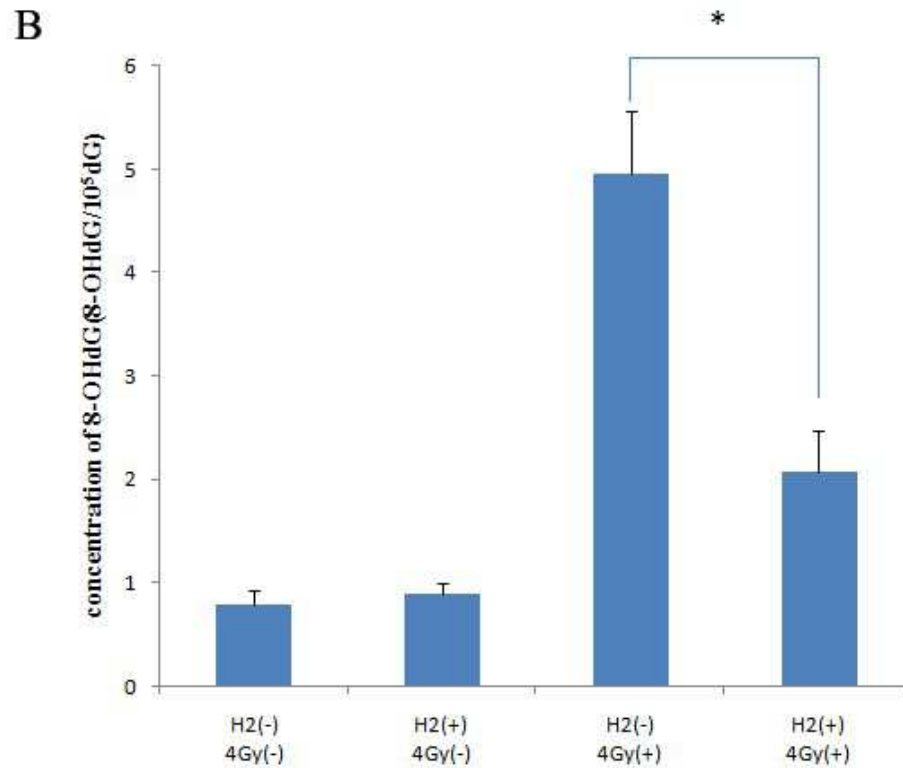


Figure5. Changes in the activities of SOD and concentrations of GSH in normal, γ -irradiated and H₂ pretreated mice. Values are given as mean \pm SEM (n=4).*P< 0.01.
187x126mm (72 x 72 DPI)



193x141mm (72 x 72 DPI)

New Only



35 Figure6. Hydrogen-rich saline significantly decreased levels of MDA, a marker of oxidative
36 stress(A).And oxidative DNA damage was assessed by 8-OHdG immunoreactivity. Shown are 12
37 hours after irradiation, intestinal 8-OHdG concentrations in normal, γ -irradiated and H2 pretreated
38 groups(B). Relative to the Control, H2 significantly decreased the concentration of 8-OHdG. Values
39 are mean \pm SEM (n=6), *P<0.01.
40 147x117mm (96 x 96 DPI)

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