Hydrogen-rich saline may be an effective and specific novel treatment for osteoradionecrosis of the jaw

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Abstract: Hydrogen, a therapeutic medical gas, can exert antioxidant activity via selectively reducing cytotoxic reactive oxygen species such as hydroxyl radicals. Hydrogen-rich saline is an alternative form of molecular hydrogen that has been widely used in many studies, including metabolic syndrome, cerebral, hepatic, myocardial ischemia/reperfusion, and liver injuries with obstructive jaundice, with beneficial results. Osteoradionecrosis of the jaw is a serious complication following radiotherapy for head and neck cancers. It has long been known that most radiation-induced symptoms are caused by free radicals generated by radiolysis of H2O, and the hydroxyl radical is the most reactive of these. Reducing the hydroxyl radical can distinctly improve the protection of cells from radiation damage. We hypothesized that hydrogen-rich saline might be an effective and specific method of managing and preventing osteoradionecrosis of the jaw.

Keywords: osteoradionecrosis, hydrogen, reactive oxygen species

Introduction

Hydrogen is one of the simplest chemical elements, forming approximately 75% of the elemental mass of the universe. It is a colorless, tasteless, odorless, non-irritating, and highly flammable diatomic gas which has been used mainly in fossil fuel processing and ammonia production.1 A series of recently published studies have demonstrated that hydrogen can selectively reduce hydroxyl radicals (OH-) and peroxynitrite (ONOO-) and exhibit therapeutic antioxidant and anti-apoptotic activities.2,3 These studies indicated that molecular hydrogen has a special role as a therapeutic gas by specifically targeting intracellular sources of reactive oxygen species (ROS), but not superoxide and hydrogen peroxide, which play physiological roles.

Subsequent research has demonstrated that a burst of ROS and reactive nitrogen species, such as OH-, superoxide anion, hydrogen dioxide, nitric oxide, and ONOO-, plays a critical role in cell damage after stroke, transplantation injury, myocardial ischemia–reperfusion injury, and other diseases. The study by Ohsawa et al2 was significant in finding that hydrogen might be a gaseous oxygen radical scavenger. This finding aroused the attention of scholars immediately after it was published, and hydrogen’s ability to prevent and cure many diseases was discovered subsequently. A diverse range of disorders and organ systems have been targeted, including ischemia–reperfusion injury in the brain, liver, myocardium, intestine, retina, and kidney; Parkinson’s disease, oxidative stress-induced cognitive decline, inflammatory disease related to oxidative stress, and metabolic syndrome.2,4–16 These diseases have a common feature, oxidative stress, which is generally associated with ROS. As diatomic hydrogen can selectively...
reduce ROS and exert antioxidant activity, hydrogen could be effective in managing and preventing these kinds of diseases.

Osteoradionecrosis of the jaw (ORNJ), which was first described by Regaud in 1922, is a notorious chronic sequela with devitalization and devascularization of bone due to irradiation, and is basically irreversible and difficult to treat. It is usually asymptomatic initially, but with the development of lesions patients present with various symptoms, including intractable pain, dysesthesia, halitosis, dysgeusia, and exposed sequestra. In the late stage patients often present with fistula from the oral mucosa or skin, complete devitalization of bone, pathological fractures, and even life-threatening complications. Numerous treatment strategies for this chronic injury have been explored, and as yet no one treatment has been universally accepted. A new theory suggesting a fibroatrophic mechanism regards ORNJ as a radiation-induced fibrotic disease, suggesting that cells in the irradiated region are damaged by acute inflammation accompanied by free radical formation. Although the actual mechanism behind the pathogenesis of ORNJ is not yet fully understood, it has been shown that radiation therapy may cause chronic oxidative stress in irradiated tissues and produce a burst of ROS, thereby inducing the activation of some transcription factors, proinflammatory molecules, and cytotoxicity, leading to chronic inflammation, organ dysfunction, fibrosis, and necrosis. After radiotherapy, ionizing radiation interacts with water molecules in biological systems, thereby inducing a variety of active free radicals, which are capable of causing cellular damage and even death; about 60%–70% of the radiation-induced cellular damage is caused by OH-. OH- can trigger the oxidation of lipids, amino acids, and saccharides, leading to the formation of various secondary free radicals. The effects of free radical scavengers in ameliorating the oxidative injuries caused by radiation have frequently been reported. Therefore, the timely elimination of radiation-induced OH- should protect normal tissues from radiation injury. This theory has led to a new therapeutic method using drugs that scavenge ROS. Some researchers have already presented dramatic results in the treatment of ORNJ using anti-fibrosis drugs such as pentoxifylline and tocopherol.

Hypothesis

Our hypothesis was that hydrogen-rich saline may be a promising, effective, and specific treatment for ORNJ, based on the following. Hydrogen, a new medical gas, could potentially selectively reduce OH- and ONOO-, exerting organ-protective effects through regulating oxidative stress and inflammation. It is so mild that it does not disturb metabolic oxidation-reduction reactions or disrupt the ROS involved in cell signaling. However, hydrogen is not convenient for use in patients, and may be dangerous because of its inflammable and explosive nature, but if dissolved in normal saline or pure water, it will be different. The primary advantage of hydrogen-rich saline is that it is portable, easily administered, and safe, with similar antioxidant effects. In fact, hydrogen water is already sold as a commodity in the market. It is physiologically safe at a correct dose because hydrogen is continuously produced by colonic bacteria in the body and circulates normally in the bloodstream. Also, the tissue compatibility of hydrogen is greater than that of many other antioxidants because it is an endogenous substance. Furthermore, hydrogen can also penetrate biomembranes and diffuse into the cytosol, mitochondria, and nuclei, thereby protecting nuclear DNA and mitochondria, suggesting that it could reduce the risk of lifestyle-related diseases and cancer. Last but not least, its relative concentration is quite high, and so it can react with low-density toxic ROS.

Evaluation of the hypothesis

ORNJ is one of the most severe complications following radiotherapy; it is a radiation-induced bone and soft tissue necrosis that occurs without metastases or the recurrence of a primary cancer. The prevalence of ORN varies widely from 5% to 15% and is most frequently seen (>70%) in the first 3 years after radiotherapy. Radiation-induced fibrosis is a new theory of ORNJ that accounts for injuries to normal tissues and bone after radiotherapy. Based on our current understanding of the pathophysiology of ORN, new protocols could be developed for its prevention and treatment. After radiotherapy the endothelial cells are injured, both directly by radiation and indirectly by free radicals. Injured cells produce chemotactic cytokines that trigger an acute inflammatory response and then generate a further release of ROS from polymorphonuclear cells and macrophages stimulated by the degradation of collagen.

According to Ohsawa et al’s research, hydrogen can selectively reduce OH- and ONOO- radicals. Qian et al also demonstrated that hydrogen has radioprotective effects both in vitro and in vivo through the antioxidant pathway.
Although ORNJ is a local lesion, usually accompanied by wound infection and vascular structural damage, ROS, especially OH- and ONOO-, mediate the main biological effects of ionizing radiation and play a major role in the initiation and progression of pathological damage in ORNJ. Therefore, we deduce that hydrogen could also have a protective role against the detrimental effects of ORNJ. However, because of its safety and convenience, hydrogen-rich saline is more suitable for clinical use (Table 1) and has been widely applied in research into ischemia–reperfusion injury and inflammation. The study by Nagata et al reported that the consumption of hydrogen-saturated water could be safe and effective, and many previous experiments have demonstrated its effects in metabolic syndrome, cerebral, renal, intestinal, and myocardial ischemia–reperfusion injury, and liver injury with obstructive jaundice. Hydrogen-rich saline could be administered by peritoneal injection, intravenously or orally, and each route has its advantages. When ingested orally, hydrogen can be detected in the blood, indicating that it is incorporated into the body. In addition, it diffuses rapidly into cells and a high level of efficacy can be expected. In animal experiments, hydrogen-rich saline administered orally ad libitum throughout the experimental period has been shown to be effective, and its effectiveness was observed in patients after consuming 900 mg/day of hydrogen-rich saline orally for 8 weeks. Above all, hydrogen-rich saline is easy to drink daily and would be simple to administer without complicating or changing the subject’s lifestyle. Despite these advantages of gastric administration, there are some drawbacks to this route of administration. Hydrogen might be lost in the stomach or intestine, as it tends to evaporate in water, and it is difficult to control its concentration and absorption. Peritoneal injection avoids this problem and is widely used in animal experiments. Although it is still hard to say how much hydrogen gas may remain after intraperitoneal injection, hydrogen concentrations in tissue can be measured using a needle-type sensor, and it has been shown in many studies that hydrogen at 5 mL/kg is really beneficial.

All of this evidence makes hydrogen-rich saline a suitable candidate to provide protective and therapeutic effects in ORNJ. There are some articles related to bone metabolism that hydrogen-rich saline will provide greater hope for the prevention of ORNJ; it will also be beneficial in managing established disease needs further study. We also believe hydrogen-rich saline could be used as an adjuvant therapy for ORNJ in conjunction with other methods, as it could delay the progress of the disease, reduce its severity, and increase the treatment effect, thereby improving patients’ quality of life. Before hydrogen-rich saline can really become a promising novel therapy for ORNJ, a great many experiments both in vitro and in vivo need to be carried out to prove its effectiveness and feasibility. We also plan to perform an animal experiment to test this hypothesis (Figure 1). After effective animal experiments, multicenter clinical studies are also essential, and because the existing clinical research is limited, the optimal strategy for using hydrogen-rich saline must also be explored.

Although modern techniques of radiotherapy and radioprotection are increasingly used to reduce adverse effects, ORNJ still occurs, and there is very little progress in the development of effective therapies against it. However, this hypothesis provides us with a new idea. With advances in laboratory and clinical research, we believe that hydrogen-rich saline will provide greater hope for the prevention of ORNJ; it will also be beneficial in managing ORNJ, probably on the basis that ORNJ is a radiation-induced fibrotic disease believed to be associated with increased oxidative stress and inflammatory reactions; and that hydrogen can selectively reduce OH- and ONOO- radicals to exert its antioxidant activity, but does not affect physiological ROS.
Establish standard animal models of ORNJ

The effect of hydrogen-rich saline on BMSCs after irradiation

Survival rate, apoptosis rate

Micro-CT scanning

The effect of hydrogen-rich saline on animal models of ORNJ

Differentiation potential

Observe the clinical outcome of ORNJ, the synthesis of Col I, Col III, and the MFB apoptosis, measure the expression of α-SMA and TGF-β1

General observation

Figure 1 An experiment is designed for this hypothesis.

Abbreviations: ORNJ, osteoradionecrosis of the jaw; BMSCs, bone marrow mesenchymal stem cells; CT, computed tomography; MFB, myofibroblast.

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Disclosure

The authors report no conflicts of interest in this work.

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