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# Production of Hydrogen Peroxide in Groundwater at Rifle, Colorado

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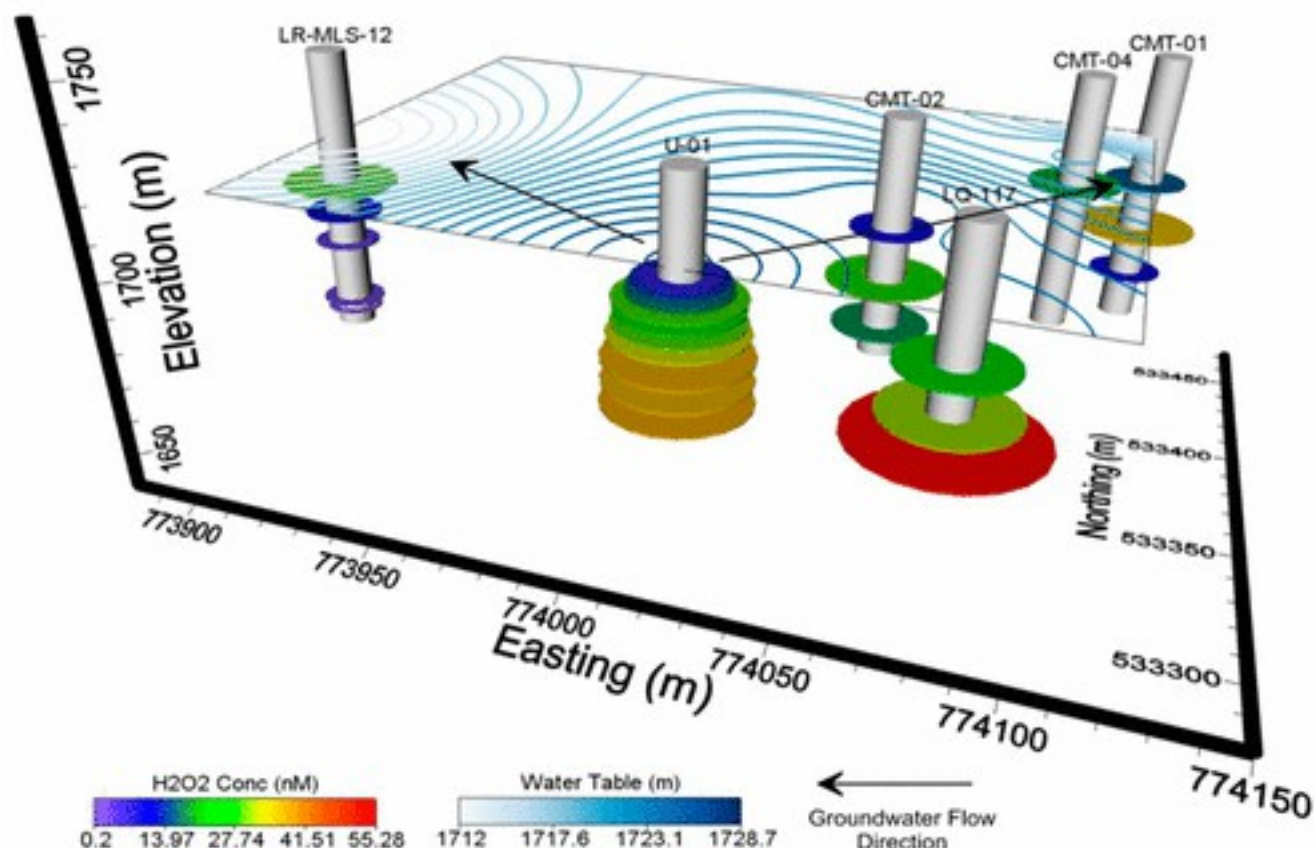
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## Abstract



The commonly held assumption that photodependent processes dominate  $\text{H}_2\text{O}_2$  production in natural waters has been recently questioned. Here, we present evidence for the unrecognized and light-independent generation of  $\text{H}_2\text{O}_2$  in groundwater of an alluvial aquifer adjacent to the Colorado River

near Rifle, CO. In situ detection using a sensitive chemiluminescent method suggests  $\text{H}_2\text{O}_2$  concentrations ranging from lower than the detection limit ( $<1$  nM) to 54 nM along the vertical profiles obtained at various locations across the aquifer. Our results also suggest dark formation of  $\text{H}_2\text{O}_2$  is more likely to occur in transitional redox environments where reduced elements (e.g., reduced metals and NOM) meet oxygen, such as oxic–anoxic interfaces. A simplified kinetic model involving interactions among iron, reduced NOM, and oxygen was able to reproduce roughly many, but not all, of the features in our detected  $\text{H}_2\text{O}_2$  profiles, and therefore there are other minor biological and/or chemical controls on  $\text{H}_2\text{O}_2$  steady-state concentrations in such aquifer. Because of its transient nature, the widespread presence of  $\text{H}_2\text{O}_2$  in groundwater suggests the existence of a balance between  $\text{H}_2\text{O}_2$  sources and sinks, which potentially involves a cascade of various biogeochemically important processes that could have significant impacts on metal/nutrient cycling in groundwater-dependent ecosystems, such as wetlands and springs. More importantly, our results demonstrate that reactive oxygen species are not only widespread in oceanic and atmospheric systems but also in the subsurface domain, possibly the least understood component of biogeochemical cycles.

## 1 Introduction

Reactive oxygen species (ROS), such as superoxide ( $\text{O}_2^-$ ), hydrogen peroxide ( $\text{H}_2\text{O}_2$ ), and hydroxyl radical ( $\text{HO}^\cdot$ ), are intermediates generated during the sequential one-electron reduction of  $\text{O}_2$  to  $\text{H}_2\text{O}$ . (1) These oxygen-containing reactive molecules, though generally present in natural environments at particularly low concentrations (picomolar to micromolar), are of considerable importance due to their transient nature and outstanding capability of oxidizing a wide range of molecules with relatively low selectivity. (2, 3) Extensive studies have shown that ROS are involved in various biogeochemical and ecologically significant processes, (4) such as the formation of manganese oxides, (5, 6) coral bleaching, (7) algal bloom-mediated massive fish kills, (8) and the establishment of microbial symbioses. (4, 9) In addition, recent increasing recognition of the important roles of ROS in carbon remineralization, (10, 11) metal bioavailability, (12) and contaminant transport urges a more comprehensive understanding of ROS generation and transformation in natural environments. Because of the transient nature of ROS and the challenges associated with direct measurement of their production capacity in a given environment, field investigations of ROS have mainly focused on  $\text{H}_2\text{O}_2$ , simply because it has a longer half-life (ranging from hours to days in water) and is thus much

more stable than  $O_2^-$  and  $HO^\cdot$ .[\(13\)](#)  $H_2O_2$  is ubiquitous in natural waters and can act as an oxidizing or reducing agent to react with a large suite of biologically important and redox-sensitive trace elements, such as iron,[\(14-16\)](#) copper,[\(17-19\)](#) chromium,[\(20, 21\)](#) and arsenic.[\(22, 23\)](#) The dominant source of  $H_2O_2$  in natural waters has long been attributed to photo-oxidation of chromophoric dissolved organic matter (CDOM) by molecular oxygen, with various studies addressing  $H_2O_2$  patterns (from undetectable to greater than 1000 nM) in a wide range of surface waters, including oceans,[\(24-27\)](#) coastal and estuarine waters,[\(28, 29\)](#) lakes,[\(30-32\)](#) freshwater,[\(33-35\)](#) and geothermal waters.[\(36, 37\)](#) However, recent studies indicated that dark production of  $H_2O_2$  in deep seawater, likely of biological origin, is potentially on par with its photochemical origins.[\(13, 38-40\)](#) For example, Wu et al.[\(13\)](#) reported persistent high  $H_2O_2$  concentrations at water depths exceeding 80 m in Northern South China Sea Shelf-sea, where solar irradiance was virtually absent, and suggested various processes, such as dark biological production, trace metal mediated redox reactions, as well as physical mixing, could all contribute to such observed vertical distribution. Further, atmospheric  $H_2O_2$ , acting as a useful indicator of radical levels in the atmosphere to imply how the troposphere changes with time in response to natural and anthropogenic perturbations, has also received a great deal of attention in atmospheric oxidant studies.[\(41-45\)](#) Concentrations of  $H_2O_2$  in atmospheric samples have been reported to be correlated with solar radiation, humidity, temperature, as well as concentrations of  $O_3$ ,  $NO_x$ , and hydrocarbons.[\(41\)](#) More interestingly,  $H_2O_2$  has also been detected in the atmosphere of Mars, with a typical concentration of 500–1200 nM, possibly generated by triboelectricity during dust storms or dust devils of the red planet.[\(46, 47\)](#)

Although some preliminary insights into the likely concentrations of  $H_2O_2$  in natural waters (both surface and deep oceans) and atmospheric systems are now available, there is a severe dearth of information regarding  $H_2O_2$  generation and patterns in the subsurface systems, a large but poorly understood domain in our global habitat. Unlike the surface environment, subsurface systems are generally  $O_2$ -deficient, which is presumably the reason that ROS production in such underground territory has been overlooked. However, the redox conditions of subsurface environments are often perturbed by  $O_2$  from both natural and anthropogenic processes (e.g., groundwater fluctuations in response to rainfall, groundwater recharge by treated wastewaters, hyporheic exchange, riverbank filtration, soil flushing, farming and construction activities, as well as flood events), leading to alterations of local biogeochemical settings and strong interactions between  $O_2$  and reduced components. Although Tong et al.[\(10\)](#) reported significant ROS generation from a variety of typical redox-fluctuating subsurface environments, Page et al.[\(11\)](#) observed dark formation of  $HO^\cdot$  by oxidation of reduced soil waters from the arctic tundra, with the highest formation rate predicted at

oxic–anoxic boundaries in arctic soil and surface waters. Another study by Zhang et al. [\(1\)](#) observed the occurrence of dark biological production of  $\text{H}_2\text{O}_2$  along the depth profiles of several brackish and freshwater ponds. As a consequence, if dark production of  $\text{H}_2\text{O}_2$  is prevalent in deep ocean, sediments, and fresh waters (either of biological or chemical origin), there is no reason to exclude  $\text{H}_2\text{O}_2$  sources in shallow groundwater systems, especially in unconfined aquifers, where extensive interactions among oxygen, microorganisms, natural organic matter, and various trace elements take place. Here, we present evidence for the unrecognized and light-independent generation of  $\text{H}_2\text{O}_2$  in the groundwater of an alluvial aquifer adjacent to the Colorado River near Rifle, CO. Our results suggest that detectable  $\text{H}_2\text{O}_2$  concentrations are common, ranging from lower than the detection limit ( $<1$  nM) to 54 nM along the vertical profiles in groundwater at several locations across the Rifle site. Recognition of ROS sources in groundwater systems not only helps us better understand their potential impact on metal/nutrient cycling on groundwater-dependent ecosystems, such as wetlands and springs, but also further advances our knowledge of the role played by  $\text{H}_2\text{O}_2$  and associated intermediates (such as  $\text{O}_2^{\cdot-}$  and  $\text{HO}^{\cdot}$ ) in ecosystems as a whole. [\(29\)](#)

## 2 Materials and Methods

### 2.1 Reagents

Analytical grade chemicals were purchased from VWR International or Sigma-Aldrich (or as otherwise stated) and used without further refinement. Acridinium ester (AE, 10-methyl-9-(*p*-formylphenyl)acridinium carboxylate trifluoromethanesulfonate) was purchased from Cayman Chemical (Michigan, USA). AE reagent of 5.0  $\mu\text{M}$  at pH 3.0 was prepared each sampling day by dilution of a refrigerated AE stock solution (250  $\mu\text{M}$ , pH 3.0) by adding 1.0 mM phosphate buffer (pH 3.0).  $\text{H}_2\text{O}_2$  (30%, ultrahigh purity) was purchased from BDH Chemicals and calibrated spectrophotometrically using a UV–vis spectrophotometer. 0.1 M carbonate buffer of pH 11.3 was prepared weekly, and other stock solutions were refrigerated at 4 °C in the dark.

### 2.2 Field Information and Experimental Measurement

#### Field Site and Groundwater Sampling

Groundwater samples were collected between May 21 and June 6, 2015 at the U.S. Department of Energy (DOE) Rifle, CO field site. The field site is located on a floodplain adjacent to the Colorado River and was a former uranium and vanadium mill operated between 1924 and 1958 (see [Figure S1](#) for a high resolution site map and Yabusaki et al. (2007) [\(48\)](#) and Janot et al. (2016) [\(49\)](#) for more

information). Currently, this site supports research activities associated with Lawrence Berkeley National Laboratory.

H<sub>2</sub>O<sub>2</sub> detection was performed by sampling groundwater through 11 wells, including a series of 3-port continuous multichannel tubing (CMT) wells installed along a transect aligned with the groundwater flow direction of the Rifle floodplain in 2014 (CMT01–CMT05, see [Figure S2](#) for detailed location map and [Figure S3](#) for borehole information on Well CMT01 as an example). Before sampling in each well, at least 24–30 L of water was slowly pumped out from the borehole, which is more than the total volume that could be held in each channel of the well, indicating the sampled water was freshly infiltrated from the nearby aquifer rather than the water initially present in the well. Sample water was peristaltically pumped through opaque flexible tubing into an amber high density polyethylene (HDPE) bottle with no head space and was analyzed within 4–6 min after collection. Filtration of the groundwater by 0.22 µm filters made no difference on the final concentration of H<sub>2</sub>O<sub>2</sub>; therefore, no filtration was done to minimize the sampling time.

### **H<sub>2</sub>O<sub>2</sub> Analysis**

In situ H<sub>2</sub>O<sub>2</sub> concentrations were determined by the chemiluminescing reaction between acridinium ester (AE, 10-methyl-9-(*p*-formylphenyl)acridinium carboxylate trifluoromethanesulfonate) and the conjugate base of H<sub>2</sub>O<sub>2</sub> (HO<sub>2</sub><sup>-</sup>, pK<sub>a</sub> = 11.6).[\(50, 51\)](#) The emitted luminescence at 470 nm was detected by flow injection analysis using a FeLume system (Waterville Analytical, Waterville, ME). To avoid any possible interference caused by additional interactions between ferrous iron and O<sub>2</sub> under alkaline conditions,[\(51\)](#) 2.0 mM Ferrozine solution (3-(2-pyridyl)-5,6-diphenyl-1,2,4-triazine-4',4''-disulfonic acid sodium salt, Sigma) was added to bind all the ferrous species before the sample entered the flow cell located at the photo multiplier tube interface, where it mixed with a carbonate (pH 11.3) buffer to initiate the chemiluminescence-generating reaction. An acid-wash loop was also integrated to rinse off any possible precipitates formed by the groundwater at alkaline conditions. After each sample measurement, 0.01 M HCl flowed through the system for at least 3 min and followed by another 2–3 min of distilled deionized (DDI) water. Because of the ubiquitousness of H<sub>2</sub>O<sub>2</sub> in most of the laboratory waters, analytical blanks were obtained by adding 3.0 mg L<sup>-1</sup> catalase (the enzyme that decomposes H<sub>2</sub>O<sub>2</sub> to water and O<sub>2</sub>) to the DDI water used for preparation of reagents and buffer solutions at least 30 min ahead.[\(50\)](#) Catalase remaining in the reagent and buffer will inevitably react with some of the H<sub>2</sub>O<sub>2</sub> in the sample. However, loss of sample H<sub>2</sub>O<sub>2</sub> is insignificant because the half-life of H<sub>2</sub>O<sub>2</sub> in 3.0 mg L<sup>-1</sup> catalase is 15 min and the flow injection analysis time is less than 30 s.[\(50\)](#) The detection limit of H<sub>2</sub>O<sub>2</sub> using analytical set up in this study is 1.0 nM determined from the calibration curves and the uncertainty in the blank.

## Other Element Analysis

Dissolved oxygen (DO) concentrations were measured on site by DO CHEMets Visual Kits (CHEMetrics, Inc.), with Rhodazine D method(52) for DO < 1 ppm (Kit No.: K-7501) and Indigo Carmine method(53) for DO of 1–12 ppm (Kit No.: K-7512). Samples for elemental analysis and total organic carbon (TOC) measurement were collected and acidified to pH 3 with HCl (high purity, 30% w/v; Sigma) and kept cool or refrigerated until shipped to the laboratory in Berkeley, CA. TOC concentrations were measured using a Shimadzu TOC-V<sub>CSH</sub> analyzer with an ASI-V autosampler (Shimadzu). Total element concentrations were determined by inductively coupled plasma mass spectrometry (ICP-MS, PerkinElmer SCIEX ICP-Mass Spectrometer ELAN DRC II).

## Kinetic Modeling

The kinetic model was fitted to the experimental data over a range of experimental conditions using the program Kintek Explorer.(54)

## 3 Results and Discussion

### 3.1 Background H<sub>2</sub>O<sub>2</sub> Profiles in Rifle Aquifer

The most finely resolved spatial profile of H<sub>2</sub>O<sub>2</sub> (Figure 1) was obtained in Well U-01 (located in Experimental Plot A, see Figure S1 for well location map), showing a vertical concentration range of 7–38 nM. The occurrence of H<sub>2</sub>O<sub>2</sub> is remarkable, not only because no H<sub>2</sub>O<sub>2</sub> generation in groundwater has been previously discovered, but also the permanent absence of light, low abundance of DO, and the widespread reactive sediments in the subsurface system that would decrease the ambient concentrations of ROS. More interestingly, H<sub>2</sub>O<sub>2</sub> concentrations observed in Well U-01 increased with water depth, with increasing concentrations in the upper half of the profile and almost constant and high H<sub>2</sub>O<sub>2</sub> levels at lower depths. This also departs from our traditional expectation where photodependent processes are the dominant cause of H<sub>2</sub>O<sub>2</sub> generation. In other words, the typical profiles of H<sub>2</sub>O<sub>2</sub> in surface waters have been widely reported to exhibit apparent diel cycles and decrease monotonically with water depth.(24-27, 33) However, exceptions to this trend have also been observed in various studies,(1, 13, 38, 55) implying that besides photoproduction, other factors related to H<sub>2</sub>O<sub>2</sub> formation and/or decomposition may be important under certain conditions.





Figure 1. Depth profiles of H<sub>2</sub>O<sub>2</sub>, total iron, DO, and TOC concentrations in Well U-01. Samples were collected on May 24, 2015. Water table was 2.55 m below the ground surface. Error bars (some of which are too small to be visible) are the standard errors from triplicate measurements.

To discover correlations between H<sub>2</sub>O<sub>2</sub> profiles and other potential critical elements, groundwater samples of Well U-01 were also characterized with respect to dissolved oxygen (DO), total iron concentration, and organic carbon content (TOC) ([Figure 1](#)). Information about other auxiliary parameters and elements is presented in [Table S1](#). As shown in [Figure 1](#), it is noteworthy that more than 70 μM DO was observed throughout the depth of this well, with the maximum concentration of 143 μM occurring 0.91 m below the water table (3.47 m below the ground), and a lower concentration of 72 μM present at the bottom. Such levels of DO in groundwater were not unexpected because oxygen can be transported faster than it is consumed in permeable sediments, and anoxic conditions are more difficult to achieve near the water table. ([49](#)) At the same time, DO in the Rifle aquifer is also tied to seasonal variations in groundwater elevation, ([56](#)) with a rising water table bringing more oxygen into the system and stronger vertical mixing between water layers. The dissolved organic carbon concentration was generally constant (~290 μM) throughout the vertical profile, except one higher concentration of 340 μM right below the water table. In surface waters, although the variations in the concentrations of H<sub>2</sub>O<sub>2</sub> and TOC have been found to exhibit close correlations (consistent with the fact that CDOM is the primary control on photochemical production of H<sub>2</sub>O<sub>2</sub>), ([13](#)) a nonuniform relationship between CDOM or dissolved organic matter fluorescence and the photochemical production rate of H<sub>2</sub>O<sub>2</sub> has also been observed in several studies. ([13](#), [26](#), [28](#)) On the other hand, formation of HO· (through the intermediacy of H<sub>2</sub>O<sub>2</sub>) during oxidation of reduced organic matter under dark conditions represents a previously unknown source of HO· formation at oxic/anoxic boundaries and may affect the biogeochemical and pollutant redox dynamics at these interfaces. ([57](#))

In the subsurface, Fe(II) mainly exists in the form of complexes in pore water and minerals in sediments and has been reported to be involved in the generation and/or decomposition of H<sub>2</sub>O<sub>2</sub>. ([10](#), [57](#)) The vertical profile of dissolved iron in groundwater at Well U-01 ([Figure 1](#)) covered a narrow range of 260–540 μM. Though not characterized in this study, essentially all of this dissolved iron was present as Fe(II). ([58](#)) A recent study that characterized the heterogeneity of organic-rich sediment in the Rifle aquifer observed abundant mackinawite (FeS) in the sediments from the capillary fringe above naturally reduced zones (NRZs), indicating that certain reducing conditions are present in the oxic portion of the aquifer, probably maintained in the interior of aggregates within organic-rich sediments. ([49](#)) The heterogeneously dispersed NRZs in the Rifle aquifer are



characterized by high concentrations of organic carbon, reduced mineral phases, and reduced forms of metals, including Fe(II), uranium(IV), and vanadium(IV). These reduced metals in the NRZs are seasonally exposed to oxygen and contribute to formation of  $O_2^-$ , the widely accepted immediate precursor of  $H_2O_2$  (though an alternative mechanism might occur when biologically mediated processes predominate(40, 59, 60)). On the other hand, Fenton and Fenton-like reactions induced by reduced trace metals have also been indicated to promote the decomposition of  $H_2O_2$  and generation of more reactive oxidants, including  $HO\cdot$  and high valent Fe species.(61, 62) For instance, generation of  $HO\cdot$  from oxidation of ligand-complexed Fe(II) by  $O_2$  has been confirmed in laboratory studies and utilized for degradation of organic pollutants in wastewaters.(62) Therefore, in a system with the presence of oxygen, reduced metals and natural organic matter (NOM),  $H_2O_2$  can be generated from oxygenation of reduced elements and metal-mediated oxidation of NOM,(14, 18, 63) whereas reduced metals and NOM (dissolved and in sediments), at the same time, could also act as sinks by further reacting with  $H_2O_2$ .(23) Thus, the relationships between  $H_2O_2$  concentrations and iron/organic carbon content could be rather complex, largely depending on the biogeochemical settings of the specific environmental system. Various other factors, including pH, temperature, biologically mediated generation and decomposition, variability and redox properties of NOM, and the presence of various scavengers, could all influence the temporal and spatial patterns of  $H_2O_2$  in the groundwater. As a consequence, no simple correlations between  $H_2O_2$  and DO, total Fe, or TOC profiles were evident in this study, and the exact mechanisms causing the observed vertical distributions of  $H_2O_2$  in Rifle groundwater cannot be stated with confidence at this stage.

Nevertheless, one explanation could be the effect of dark biological production associated with different bacterial activity with distance from the water table, and none of the examined elements (DO, total Fe, or TOC), was the rate-determining factor.

Wu et al.(13) indicated in their recent study of tropical shelf waters that concentrations of  $H_2O_2$  were generally elevated at places where the activity of internal waves and mixing was extensive, but in this study, no  $H_2O_2$  was observed in another location (Well 310, Figure S2), which is ~30 m closer (compared to Well U-01) to the Colorado River bank. Thus, mixing of water from the river should not be a significant contributor, at least not in the Rifle groundwater. This is further confirmed by another vertical profile of  $[H_2O_2]$  that was obtained at the location of Well LR12 (Figure 2), a multilevel sampling (MLS) well (detailed diagram of the MLS construction was shown in a previous study(58)) located in Experimental Plot B (Figure S1).  $H_2O_2$  concentrations ranged from 28 nM near the water table to lower than the detection limit (<1 nM) at the bottom of this borehole. Similar to  $H_2O_2$  profile, concentrations of other relevant solutes (DO, total Fe, and TOC) decreased along the water depth,

with the only exception occurring at a depth of 4.3 m below the water table. Unlike the conditions at Well U-01, significant  $O_2$ -deficiency ( $DO < 2.5 \mu M$ ) was found throughout the depth of this well, which might exert a limiting factor for in situ  $H_2O_2$  formation. However, Borda et al.(64) and Zhang et al. (23) reported production of  $H_2O_2$  and  $HO\cdot$  from the oxidation of  $H_2O$  at sulfur-deficient sites on pyrite surface under anoxic conditions, demonstrating that  $O_2$  is not necessarily a prerequisite for ROS generation under some circumstance. The sharp decrease of  $H_2O_2$  in the upper-half of the vertical profile at Well LR12 may be correlated to the notable decrease of total iron and TOC concentrations at the same elevation, while the distinguishable rebound at 4.3 m below the water table happened simultaneously with locally higher level of DO and total iron.



Figure 2. Depth profiles of  $H_2O_2$ , total iron, DO, and TOC concentrations in Well LR12. Samples were acquired on June 04, 2015. Water table was 3.20 m below the ground surface. Error bars (some of which are too small to be visible) are the standard errors from triplicate measurements.

Various concentrations of  $H_2O_2$  (range of 10–54 nM) were also detected in a few other locations of this aquifer (3-port CMT wells, detailed elevations presented in Table S2), as collectively shown in Figure 3. Interestingly, the highest  $H_2O_2$  concentrations at all these wells happened coincidentally in groundwater that was located 2.0–3.0 m below the water table. Another recent study on dark biological production of ROS in brackish and freshwater systems(1) reported that ROS ( $O_2^-$  and  $H_2O_2$ ) generation rates peaked at 2.0–4.0 m below the water surface and  $H_2O_2$  profiles did not show apparent dependence on the water depth in the aphotic zone. To be more specific, in freshwater system, photosynthetically active radiation (PAR) declined to 3.1% and 0.9% of the surface level at 3.5 and 4.5 m below the water surface, whereas  $H_2O_2$  concentrations were 65.6% and 87.2% of that in surface water,(1) indicating that light-intensity might not be the limiting factor for  $H_2O_2$  formation at these depths. Such atypical  $H_2O_2$  profiles in their study were mainly attributed to biological production and possible vertical mixing throughout the shallow waters.(1) At the same time, Tong et al.(10) also found that during oxygenation of various sediments, the generation rate of  $HO\cdot$  increased significantly with the sample depth of the sediments, suggesting that the more reduced the sediments, the more  $HO\cdot$  will be produced after exposure to oxygen. During our sampling time (May to June 2015), the water table at the Rifle site was increasing quickly (Figure S4), a situation typical during later spring/early summer in the Colorado river basin when the river stage is rising as a result of snowmelt. Such seasonal change in water table inevitably brings more oxygen into the

groundwater, and thus alters local redox conditions and biogeochemical settings in the aquifer. For example, introduction of O<sub>2</sub> into the region below the water table creates oxic–anoxic boundaries, where anoxic waters mix with oxic flows and thus initiates cascade redox reactions. Of particular interest here is the generation of large amounts of ROS under such fluctuating redox conditions that has been recently identified in several studies.[\(1, 10, 11, 16, 65\)](#) The observed H<sub>2</sub>O<sub>2</sub> spike occurring at 2.0–3.0 m below the water table in various locations of the Rifle aquifer ([Figure 3](#)) could be caused by intensive ROS generation at oxic–anoxic interfaces resulting from seasonal variations in the groundwater table. In fact, not only natural processes, such as groundwater fluctuations and recharge, hyporheic exchange and episodic, high intensity events (like storms and floods), but also anthropogenic activities, including soil flushing, groundwater replenishment, farming and construction activities, could all result in the formation of oxic-anoxic mixing zones where oxygen concentrations vary intermittently and considerable amounts of ROS are generated. A recent estimate[\(16\)](#) based on radium isotope inventories suggested that approximately 6.0 × 10<sup>10</sup> kg of water exchanged between oxic and anoxic conditions per day for the entire South Carolina coastline, which implied a potential abiotic ROS flux of up to 1.5 × 10<sup>7</sup> moles day<sup>-1</sup>, comparable in magnitude to photochemical sources of ROS in surface waters (including coastal and seawaters).[\(66, 67\)](#) More importantly, such sustained ROS generation may also contribute to global carbon cycling by breaking down otherwise recalcitrant organic compounds (such as bisphenol A, and polycyclic aromatic hydrocarbons)[\(57, 68\)](#) into better microbial energy sources, an alternative but important pathway for microbial alteration and consumption of natural and anthropogenic organic carbon.

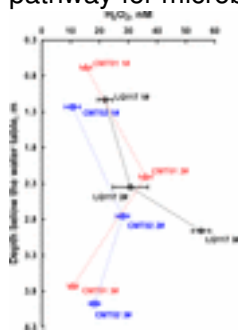


Figure 3. H<sub>2</sub>O<sub>2</sub> concentrations at various depths of different CMT wells in the Rifle site. Water table was below the ground surface: CMT01, 3.22 m; CMT02, 1.76 m; LQ117, 1.71 m. Error bars (some of which are too small to be visible) are the standard errors from triplicate measurements.

Overall, detectable H<sub>2</sub>O<sub>2</sub> was found at six of the eleven locations that we measured at the Rifle aquifer, besides the aforementioned five wells that had H<sub>2</sub>O<sub>2</sub> along the full depth of the well, 19.7 nM H<sub>2</sub>O<sub>2</sub> was found at 0.28 m below the water table of Well CMT04, but no H<sub>2</sub>O<sub>2</sub> was observed below

this depth, possibly because of the severe O<sub>2</sub> deficiency at the bottom of the well ([Table S1](#)). It is noteworthy that all of the five wells where no H<sub>2</sub>O<sub>2</sub> was detected uniformly had extremely low concentrations of DO (generally DO < 0.1 mg/L), whereas most of the locations where H<sub>2</sub>O<sub>2</sub> was detected exhibited higher DO concentrations, except Well LR12, of which detectable H<sub>2</sub>O<sub>2</sub> (up to 28 nM) was found in groundwater with remarkably low DO ([Figure 2](#)). At the same time, for the deep elevation of CMT04 and all depths of CMT05, constant high concentrations of iron and manganese ([Table S1](#)) were observed in the absence of H<sub>2</sub>O<sub>2</sub>, indicating that under reducing conditions such metal species may act as H<sub>2</sub>O<sub>2</sub> consumers, rather than contributing to H<sub>2</sub>O<sub>2</sub> formation by reducing O<sub>2</sub>. ([57](#))

### 3.2 Dark Formation of H<sub>2</sub>O<sub>2</sub> in O<sub>2</sub> Injection Experiments

In addition to background H<sub>2</sub>O<sub>2</sub> detection in the aquifer, O<sub>2</sub> injection field experiments were also conducted to investigate the oxidation of naturally reduced uranium in Rifle aquifer sediments and its potential significance to uranium plume persistence. ([69](#)) Briefly, groundwater (~772 L) from the naturally reduced zones in wells of JB04 and JB02 ([Figure S2](#)) was pumped into an opaque and gas-impermeable bag and mixed with a conservative tracer (Br<sup>-</sup>). A gas phase composed of 97% O<sub>2</sub> and 3% CO<sub>2</sub> was continuously bubbled through the groundwater in the bag for 2.8 h. Then the groundwater with a high concentration of DO (0.66 mM) was pumped back into the subsurface in Well JB04. First of all, no background H<sub>2</sub>O<sub>2</sub> was detected in groundwater sampled from wells JB04 and JB02 before the experiment. However, after bubbling O<sub>2</sub> into the bag, H<sub>2</sub>O<sub>2</sub> concentration of the groundwater in the bag reached 252 nM ([Figure S5](#)), demonstrating generation of ROS when O<sub>2</sub> is supplied to anoxic water. It is noteworthy that even though more than 250 nM H<sub>2</sub>O<sub>2</sub> (in ~772 L water) was injected into Well JB04, immediate sampling at Well JB04 after the injection (total injection time ~ 2.9 h) showed no residual H<sub>2</sub>O<sub>2</sub>, indicating the existence of significant sinks in this location, and that all of the injected H<sub>2</sub>O<sub>2</sub> was consumed quickly in the local groundwater. However, 40–60 h after the injection, relatively high concentrations of H<sub>2</sub>O<sub>2</sub> were detected at adjacent down-gradient wells (maximum concentrations of 97 and 213 nM in JB02 and JB05, respectively, [Figure S6](#)), as shown in [Figure 4](#) (see [Figure S2](#) for location map of these two adjacent wells). It is reasonable to speculate that the observed concentrations of H<sub>2</sub>O<sub>2</sub> in these downgradient wells could have been originated from the H<sub>2</sub>O<sub>2</sub> initially generated in the injection bag. That is, the 252 nM H<sub>2</sub>O<sub>2</sub> generated in the 772 L of injection water was not completely consumed at the injection spot of JB04, instead it moved downstream to wells of JB02 and JB05 after 40–60 h of transport. However, if we assume that the unconsumed H<sub>2</sub>O<sub>2</sub> (if there was any) from JB04 would transport downstream in a pattern similar to that of Br<sup>-</sup>, it is worth noting that the normalized concentrations of H<sub>2</sub>O<sub>2</sub> (the concentration divided by the injected H<sub>2</sub>O<sub>2</sub> concentration) at these wells had a very different breakthrough pattern than Br<sup>-</sup>

([Figure 4](#)), especially for JB05, where the value of  $C/C_0$  for  $H_2O_2$  was much higher than that for Br<sup>-</sup>. These results illustrate that the detected  $H_2O_2$  patterns at wells of JB02 and JB05 could not be totally attributed to  $H_2O_2$  from the injected water and  $H_2O_2$  was not conservative during transport downgradient due to the presence of various sources and sinks. Moreover, the estimations in [Figure 4](#) were based on the assumption that there was no  $H_2O_2$  consumption during the transport, which almost certainly was not the case, because no  $H_2O_2$  was detected in JB04 immediately after the injection, indicating that consumption along the flow path was indeed significant. As a consequence,  $\sim 200$  nM  $H_2O_2$  detected throughout the three channels of JB05 had to be actively produced along the flow path, or perhaps in the vicinity of the sampling location in order to balance such rapid decomposition. Even though  $H_2O_2$  has a relatively long lifetime (hours to days) in the absence of reactive solutes,[\(13\)](#) it is more likely that the abundant reduced species in the local environment around the wells consume  $H_2O_2$  rapidly and prevent it from traveling a great distance. Although the reason that more  $H_2O_2$  was produced in JB05 than JB02 (the closer well to JB04) could not be determined with certainty, dark biological production coupled with spatially variable bacterial activity in the aquifer could potentially be one of the causes, because soils are the most heterogeneous parts of the biosphere, with an extremely high variability of properties and processes within nano- to macroscales.[\(70\)](#) It has been reported that physicochemical and redox conditions of the Rifle aquifer differ vertically on a cm scale and laterally on a scale of less than 2 m.[\(49\)](#) Accordingly, highly diverse and dynamic microbiological zonation are more likely to exist in such a heterogeneous aquifer.[\(49\)](#) Besides, Murphy et al.[\(16\)](#) recently reported that when anoxic groundwater moved through oxidized, Fe(III) rich sediments, the presence of sulfide could rapidly reduce Fe(III) to Fe(II) and thus trigger a cascade of redox reactions between oxygen and iron species (with Fe(II) acting as an electron shuttle between reduced S species and  $O_2$ ), leading to significant ROS generation. These observations point to a dynamic redox cycling of iron, sulfide, and  $H_2O_2$  upon introducing oxygen into the aquifer, either by experimental perturbations, such as the oxygen injection of this study, or through more widespread natural processes like seasonal variations of groundwater elevation.

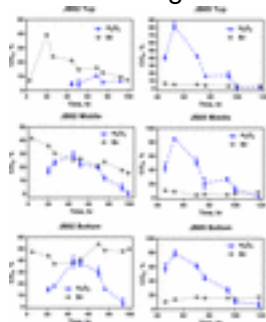


Figure 4.  $\text{H}_2\text{O}_2$  and  $\text{Br}^-$  concentrations in downstream wells (JB02 and JB05) after oxygenated groundwater was injected in an adjacent upstream well JB04 (injection at JB04 started at  $t = 0$ ). 1.10 mM NaBr was added to the injection water as a conservative tracer.  $C_0$  of  $[\text{Br}^-] = 1.10$  mM and  $C_0$  of  $[\text{H}_2\text{O}_2] = 252$  nM (the last concentration detected in the injection tank, as shown in [Figure S5](#)). Error bars (some of which are too small to be visible) are the standard errors from triplicate measurements. Detailed elevation information on the “top, middle, and bottom” baffles is provided in [Table S2](#).

### 3.3 Mechanistic Insights of $\text{H}_2\text{O}_2$ Generation and Decomposition in Subsurface Systems

As previously stated, there are multiple pathways contributing to  $\text{H}_2\text{O}_2$  generation and decomposition in the subsurface system, including dark biological production and consumption, redox cycling of metal species (either dissolved or in sediments) driven by fluctuating dissolved oxygen levels, and metal-mediated oxidation of reduced organic matter. First of all, there is increasing evidence indicating that many groups of microorganisms, including fungi, various species of phytoplankton and microalgae, symbionts in corals, and heterotrophic bacteria, are capable of extracellular ROS production.[\(1, 13, 71\)](#) One recent study by Zhang et al.[\(1\)](#) attributed their observed dark profiles of  $\text{H}_2\text{O}_2$  in several brackish and freshwater ponds to biological sources since the addition of formaldehyde, which is capable of rapidly killing microorganisms within the water, significantly inhibited the dark particle-associated production of  $\text{H}_2\text{O}_2$  in all of their samples. Although the biological groups in the subsurface domain that are actively involved in  $\text{H}_2\text{O}_2$  production are not known, they may be one of the important contributors to the observed  $\text{H}_2\text{O}_2$  patterns in the Rifle groundwater.

Despite the existence of a small portion of ROS production that was not inhibited by formaldehyde in the study by Zhang et al.,[\(1\)](#) Tong et al.[\(10\)](#) reported that sterilization of sediments led to only a negligible difference in  $\text{HO}^\cdot$  accumulation under oxic conditions, while Dixon et al.[\(72\)](#) observed sterilization could eliminate the majority but not all of the  $\text{H}_2\text{O}_2$  decay in dark incubations of soil suspensions, both pointing out that in addition to the biological origin, abiotic reactions, more or less, contribute to the formation and consumption of  $\text{H}_2\text{O}_2$ . Of particular interest here is the recently identified ROS production/decomposition resulted from oxygenation of various reduced elements that are widely presented in the subsurface domain. For instance, Page et al.[\(11\)](#) suggested cascade occurrence of dark ROS formation in environments where reduced NOM and Fe(II) produced by anaerobic microbial respiration are exposed to  $\text{O}_2$ , such as at oxic–anoxic interfaces. Therefore, under fluctuating redox conditions,  $\text{H}_2\text{O}_2$  can be generated via interactions between oxygen and (a) reduced elements such as Fe(II) present either in pore water, adsorbed to sediment surfaces or within the structure of subsurface minerals (such as pyrite and mackinawite); or (b) reduced organic

moieties (such as hydroquinones) present in dissolved or soil organic matter, especially in the presence of transition metal species (such as iron and copper).[\(14, 18, 19, 73\)](#) Although the oxidation of dissolved Fe(II) by oxygen has long been reported to result in production of ROS,[\(74-76\)](#) oxygenation of Fe(II) species that are adsorbed to clays or oxides has also been recognized to occur at a substantially greater rate than the homogeneous oxygenation of Fe(II),[\(77\)](#) both accompanied by the formation of  $O_2^-$  as a byproduct and subsequent formation of  $H_2O_2$  by its disproportionation. With respect to Fe(II) minerals, recently Tong et al.[\(10\)](#) reported an abundant generation of  $HO^\bullet$  (with  $H_2O_2$  acting as the intermediate) on oxygenation of 33 subsurface sediments sampled from a variety of typical redox-fluctuating subsurface environments and found that Fe(II)-containing minerals, particularly phyllosilicates, are the predominant contributor to  $HO^\bullet$  formation. Significant amounts of  $H_2O_2$  are also formed during the oxygenation of pyrite[\(23, 78\)](#) and mackinawite[\(79, 80\)](#) over a wide range of conditions. Meanwhile, it is clear that these various forms of Fe(II) species not only contribute to  $H_2O_2$  formation but also are involved in subsequent transformation of  $H_2O_2$  to  $HO^\bullet$  through Fenton reactions, thus enhancing microbial degradation of natural organic materials by generating smaller soluble organic compounds from macromolecules and contributing to carbon and metal cycling in subsurface systems.

Previous studies on interactions between NOM and catalytic metal species under dark conditions demonstrate another pathway for ROS generation in the subsurface regime and provide a framework for modeling the generation of  $H_2O_2$  as an intermediate during such redox processes.[\(14, 18, 19, 63, 73\)](#) Because the electron donating capacity (EDC) of NOM has been widely attributed to the presence of hydroquinone moieties within NOM that could be subsequently oxidized to semiquinone and quinone species, here we employed a recently reported kinetic model on iron-mediated oxidation of methoxyhydroquinone ( $MH_2Q$ ) under dark conditions[\(14\)](#) as a simplified scenario to evaluate the impact of metal–NOM interactions on  $H_2O_2$  formation in Rifle groundwater. Detailed model assumptions are presented in [section S14](#), and the full list of model reactions and rate constants[\(14\)](#) is shown in [Table S3](#). Together with inputting the detected concentrations of DO and dissolved iron into the model, the predicted vertical distributions of  $H_2O_2$  in groundwater at Well U-01 and LR12 are shown in [Figure 5](#). In general, modeled dark production of  $H_2O_2$  from metal–NOM interactions was well predicted, or at least within a reasonable range to the measured vertical distributions of  $H_2O_2$  at these locations. The “sink” role played by the exceptionally high concentration of iron right below the water table of LR12 might have been overestimated as a consequence of the model assumptions. To be more specific, the total iron concentrations detected at different well elevations were input to the model as reduced Fe(II), but these reduced species are likely to be



bound in various minerals or complexed by various organic ligands, thus possibly limiting their reactivity with  $\text{H}_2\text{O}_2$ . As a consequence, the actual “sink” role from reaction 10 (the Fenton reaction, [Table S3](#)) might be less important than predicted in the model, which may explain the larger mismatch between the model and the measured  $\text{H}_2\text{O}_2$  concentration at such depth. Meanwhile, it is also important to point out that in the model, it is assumed that all of the electrons donated by the reduced organic matter were transferred to Fe(III), whereas in the real environment, there must be various electron acceptors in the vicinity that act as efficient competitors with Fe(III) for those electrons, indicating that  $[\text{H}_2\text{O}_2]$  from the model output might be overestimated, such as in the model predictions for Well U-01. On the other hand, if the Fenton reaction (reaction 10, [Table S3](#)) was removed from the model, the output steady state concentrations of  $\text{H}_2\text{O}_2$  under conditions for Well U-01 were around 1–2  $\mu\text{M}$ , merely resulting from redox interactions among iron, hydroquinone, and oxygen. This again confirms the importance of metal–NOM interactions on ROS generation, as well as the “sink” effect resulting from the Fenton reaction in transforming  $\text{H}_2\text{O}_2$  to  $\text{HO}\cdot$  in the system. For instance, it has been reported that a production rate of  $\text{HO}\cdot$  of 2–9  $\mu\text{mol L}^{-1} \text{d}^{-1}$  from the Fenton reaction with a dissolved organic carbon concentration of 10  $\text{mg L}^{-1}$  will be approached as long as enough Fe(II) is present to react with the formed  $\text{H}_2\text{O}_2$ ,[\(81\)](#) suggesting that  $\text{HO}\cdot$  production from the Fenton’s reaction has the potential to cause significant effects on NOM transformation. In addition, a previous study in the Mediterranean and Baltic Seas[\(82\)](#) showed that both  $\text{H}_2\text{O}_2$  production and decay rates in polluted water were higher than that at clean stations, indicating a more dynamic and influential role of  $\text{H}_2\text{O}_2$  in pollutant transformation.

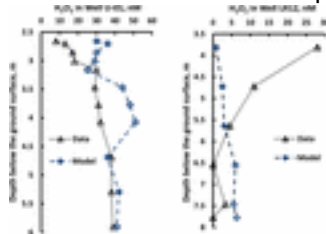


Figure 5. Detected and model predicted  $\text{H}_2\text{O}_2$  concentrations along the depth profiles of Well U-01 and LR12. This is shown as one possible pathway contributing to dark  $\text{H}_2\text{O}_2$  generation: the simplified interactions between metal and reduced organic matter. Modeled reactions and rate constants are shown in [Table S3](#) and reported in Yuan et al. (2016).[\(14\)](#)

These observations have significant implications for micropollutant fate and carbon cycling and suggest a possible abiotic link between microbial metabolism and carbon oxidation. Recent research on global terrestrial ecosystem soil and vegetation components has suggested that soil organic matter contains more than three times as much carbon as either the atmosphere or terrestrial

vegetation,[\(83\)](#) and thus sustained ROS generation in the subsurface may represent an important pathway for global carbon cycling. Nevertheless, this model only represents one of the multiple pathways leading to  $\text{H}_2\text{O}_2$  generation/decay and undoubtedly there are other sources and sinks of  $\text{H}_2\text{O}_2$  in groundwater systems. Elucidating the importance of each pathway to the overall  $\text{H}_2\text{O}_2$  patterns in the groundwater is beyond the scope of this study, but further studies are urged to reach a more comprehensive understanding of ROS generation and transformation in subsurface systems.

### **3.4 Implication of Findings**

This study represents the first field measurement of light-independent  $\text{H}_2\text{O}_2$  production in groundwater. Collectively, our results demonstrate that  $\text{H}_2\text{O}_2$  profiles in groundwater at the Rifle site are highly spatially variable, largely depending on the features of the surrounding aquifer. There are many possible pathways shaping those  $\text{H}_2\text{O}_2$  patterns, including dark biological production and consumption, metal-mediated oxidation of NOM, and cycling of metal species driven by fluctuating redox conditions. These results point to a need to understand better  $\text{H}_2\text{O}_2$  generation and cycling in the aphotic subsurface regime because of its potential importance to metal/nutrient cycling and contaminant mobilization in groundwater-dependent ecosystems.

This study also demonstrated that fluctuating redox conditions created at oxic/anoxic interfaces are likely to be “hot spots” for ROS generation: narrow boundaries in the aquifer with much faster process rates and much more intensive interactions compared to an average condition. Oxic/anoxic boundaries are ubiquitous in many natural environments, such as soils, sediments, and wetlands. Additionally, water tables in soils may fluctuate on the time scale of hours to days, leading to a relatively rapid succession of oxidizing and reducing conditions and hence repeated formation of  $\text{H}_2\text{O}_2$  and other ROS species. Such highly reactive oxidants generated under dark conditions, similar to their photochemically produced counterparts, will inevitably promote carbon remineralization by decreasing the persistence of recalcitrant organic compounds and making them more susceptible to bacterial utilization or less resistant to chemical decomposition. Meanwhile, the sustained ROS formation at the oxic/anoxic boundaries may play a role in controlling the redox speciation of various trace metals and thus alter their bioavailability or toxicity in the subsurface environment, which may in turn affect the functioning of the ecosystem as a whole.

### **[Supporting Information](#)**

The Supporting Information is available free of charge on the [ACS Publications website](https://pubs.acs.org) at DOI: [10.1021/acs.est.6b04803](https://doi.org/10.1021/acs.est.6b04803).

- Additional detailed information on well locations at the Rifle site, measured auxiliary parameters of water samples, extra experimental data, and modeling development ([PDF](#))
- **PDF**
  - o [es6b04803\\_si\\_001.pdf \(1.31 MB\)](#)

## Production of Hydrogen Peroxide in Groundwater at Rifle, Colorado

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Supporting Information for the manuscript entitled

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Production of Hydrogen Peroxide  
in Groundwater at Rifle, Colorado

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Rifle integrated field research  
h challenge site (IFRC) base map

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