The role of green rust in the environment: A review

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Abstract: Iron(II)-iron(III) hydroxy-salts known as green rusts (GRs) initially were identified and studied as corrosion products of steel and iron. Recently they were discovered as minerals present in hydromorphic soils and sediments. Different studies have suggested that GRs, due to their high reactivity, play an important role in the fate and transport of many contaminants in suboxic soils, sediments and aquifers where microbial reduction rates are limited by low carbon content. Also, GRs have considerable potential for applications such as water-purification processes and gas adsorbents. However, as GRs are only a transient state during the reduction of ferric iron and are converted to more stable forms, it is still necessary to continue studies about their behavior.

Key words: Fe(II)Fe(III)-hydroxides, abiotic transformations, remediation.

INTRODUCTION

Green rusts (GRs) are unstable compounds containing a mixture of ferrous and ferric iron and are known as green rusts (GRs) due to their bluish green colors. Structurally, green rust (GR) belongs to a family of minerals known as layered double hydroxides (LDH) synthesized for the first time by Girard & Chaudron (1935), cited by Trolard et al. (1997). The discovery of these compounds in natural environments was connected to corrosion of steel in the soil environment. green rust was first described by Keller (1948) cited by Cuttler et al. (1990), who produced a chloride and sulphate GRs. Initially green rusts had been identified and studied as corrosion products of steel and iron water distribution pipes and are now recognized as an important intermediate phase in the corrosion of Fe(0) (Stampfl, 1969). They were also identified as products of pitting corrosion processes of stainless steels (Boucherit et al., 1991; Bigham & Tuovinen, 1985). Recent research has identified green rust II as corrosion product for carbon steel exposed to the atmosphere (Rodriguez et al., 2002).

A study by Bernal et al. (1959) of iron oxyhydroxides, which included green rusts, identified two forms of the sulphate species as well as the chloride form. These GRs were prepared “by the oxidation of ferrous iron solutions” but the method was not described in sufficient detail to enable repetition of these studies.

Green rusts are formed by a number of abiotic and biotic processes under circumneutral to alkaline conditions in suboxic environments. They have been identified as products of both abiotic and microbiologically induced corrosion of iron and steel.
Various strains of the dissimilatory iron reducing bacterium *Shewanella putrefaciens* produce green rusts as products of bioreduction of hydrous ferric oxide and lepidocrocite (Fredrickson et al., 1998; Parmar et al., 2001; Glasauer et al., 2002; Genin, 2003). Recent investigations in the laboratory have proved that a GR was the resulting product of reduction of lepidocrocite by this bacterium (Ona-Nguema et al., 2002). Moreover, GRs are also observed as products of the anaerobic biooxidation of Fe(II) by *Dechlorosoma suillum* (Chaudhuri et al., 2001). Refait et al. (2003), recently studying the mechanisms leading to GR - SO$_4$ in marine corrosion of steel, reported that the presence of this mineral in the outer part of the rust layer on the steel could be explained by microbially influenced corrosion from dissimilatory iron reducing bacteria in anaerobic conditions. Erbs (2003) investigating the biotic formation of GR in two different anoxic bath systems using a dissimilatory iron(III)-reducing microorganism coupling oxidation of organic matter with iron(III)reduction (*Shewanella algae* BrY) and autotrophic iron(II) oxidizers coupling oxidation of iron(II) with nitrate reduction, found that the nitrifying Fe(II)-oxidizing bacteria produce a green intermediate solid phase within 3 days after inoculation.

Based on the bluish green color often encountered in wet soils and sediments and the rapidity with which this color changes on exposure to oxygen, Taylor & McKenzie (1980) argued that green rust might form in natural environments. From studies of the oxidation products of synthetic green rust, Taylor (1980) and Hansen (1989) suggested that the green rusts might be important precursors for different iron oxides in soils, depending on environmental conditions. However, the presence of green rust in a natural environment needed to be demonstrated to substantiate its importance. But, the rapid reaction of green rusts with atmospheric oxygen made it difficult to identify these compounds in natural and engineered systems (Williams & Scherer, 2001). Based on the bluish green color often encountered in wet soils and sediments and the rapidity with which this color changes on exposure to oxygen, Taylor & McKenzie (1980) argued that green rust might form in natural environments. From studies of the oxidation products of synthetic green rust, Taylor (1980) and Hansen (1989) suggested that the green rusts might be important precursors for different iron oxides in soils, depending on environmental conditions. However, the presence of green rust in a natural environment needed to be demonstrated to substantiate its importance. But, the rapid reaction of green rusts with atmospheric oxygen made it difficult to identify these compounds in natural and engineered systems (Williams & Scherer, 2001). Usually, oxidation of green rusts results in the formation of ferrihydrite (Fe$_{x}$H$_{y}$O$_{z}$), goethite (α-FeOOH), akaganeite (α-FeOOH), lepidocrocite (α-FeOOH), hematite (α-Fe$_2$O$_3$) maghemite (α-Fe$_2$O$_3$) or magnetite (Fe$_3$O$_4$), depending on pH, solution composition, oxidant, rate of oxidation, and the degree and rate of dehydration (Loyaux-Lawnczak et al., 2000; Lin et al., 1996). Green rusts are believed to play a central role in the redox cycling of Fe in aquatic and terrestrial environments.

More recently, however, direct evidence for the presence of green rusts in hydromorphic soils under forest in Fougères, Brittany, France, has been tentatively identified as hydroxy-green rust (GR$_{OH}$) and given the proposed name fongerite (Trolard et al., 1996 cited for Génin et al., 2001; Trolard et al., 1997). In accordance with Bourrié et al. (1999), the solubility of Fe in hydromorphic soil solution is controlled by equilibrium with GR$_{OH}$. But conclusive evidences for the formation of GRs in soils and sediments is still waiting (Hansen, 2003).

**COMPOSITION, STRUCTURE AND SYNTHESIS OF GREEN RUST**

Green rusts are layered Fe(II)-Fe(III) hydroxides having a pyroaurite-type structure consisting of alternating positively charged hydroxide layers and hydrated anion layers. Some of the Fe(II) of the octahedral sheets of Fe(OH)$_2$ is replaced by Fe(III). This results in a positive layer charge that is balanced by the inclusion of anions between the layers (Bernal et al., 1959). Its exact nature depends on the interlayer anion, and various forms have been synthesized and studied. However, the results of many studies have shown that GRs obey to a specific chemical composition, a stoichiometry, and can be represented with the general formula:

$$[\text{Fe}^{II}_{(6-x)}\text{Fe}^{III}_x(\text{OH})_{12}]^{3+}[\text{An}^{-}\times\text{H}_2\text{O}]^{x^-}$$

where $x = 0.9 - 4.2$, An$^-$ is a n-valent anion (typically CO$_3^{2-}$, Cl$^-$ or SO$_4^{2-}$, and $y$ denotes the varying amounts of interlayer water (typically $y = 2$ to 4 for most GRs).

The literature distinguishes between GRI (An$^-$ = F-, Cl-, Br-, I-), GRII (An$^-$ = SO$_4^{2-}$) and GR-CO$_3$ (An$^-$ = CO$_3^{2-}$). This division is due to a crystallographic classification of the mineral structures by which GRI is described as rhombic obtained with “planar” anions such as chlorides, carbonates etc, while GRII and GR-CO$_3$ are hexagonal obtained with three-dimensional tetrahedral anions such as sulphate or selenate (Legrand et al., 2001). Green rusts, unlike most iron oxides and whatever their form, have an internal surface area so that these minerals have great specific surface areas and consequently great reactivity. But, most of the work reported to date has focused on the reactivity of sulfate green rust rather than the chloride or carbonate forms of green rust (Williams & Scherer, 2001). They represent reactive ion exchangers and sorbents (Hansen et al., 2001). The amphoteric surface hydroxyl groups lead to both sorption of heavy metals (cations) as well as organic anions, e.g. linear alkyl benzene sulfonates (LAS), the major synthetic surfactant used in laundry detergents and cleaning products worldwide, and inorganic anions, e.g. silicate, arsenate and selenate. Furthermore, polar non-charged compounds are sorbed into the interlayer. In addition, cations as Cu(II), Ni(II), Zn(II), Cd(II), Co(II) and Mg(II), may isomorphically substitute for Fe(II) during green rusts formation (coprecipitation). For instance, nickelous-ferric green rusts, the end products obtained by substitution of Fe(II) ions by Ni(II) ions in green rust, have been demonstrated by Refait & Génin (1993, 1997) and Refait et al. (1994, 1998). This incorporation of cationic inorganic contaminants into the structures of green rusts could provide an effective means of sequestering contaminants in the subsurface.

Various processes have been developed to synthesize GRs in the laboratory. Two main procedures were envisioned, the first one by partial oxidation of Fe(II) hydroxides and hydrated anion layers, the second one by mixing Fe(II) and Fe(III) aqueous solutions or suspensions (coprecipitation). The anion associated with Fe(II) or Fe(III) in the salt is thus...
Abiotic transformations (oxidations/reductions) of inorganic and organic substances are generally much slower than biotic reactions, but they have proved significant within the time scales commonly associated with groundwater movement. Groundwater and hydromorphic soils constitute environments with low microbial activity in which the abiotic transformation of organic and inorganic contaminants can be important in the presence of highly reactive GR. Abiotic reduction of different anions in the presence of synthetic GRs has been reported, for example, the reduction of nitrite (NO$_2^-$) and nitrate (NO$_3^-$) to ammonium (NH$_4^+$) by GR-SO$_4$. This reduction is a thermodynamically favorable reaction, which may be expected to occur in natural anoxic waters (Hansen et al., 1994; Hansen et al., 1996). Considerable interest has been given to reduction of NO$_3^-$, which is widely known as an agricultural pollutant (Sposito, 1989), as it protects groundwater from excessively high NO$_3^-$ concentration and contributes for the maintenance of the nitrogen in the ground (Verdegem & Baert, 1985). The green rust-facilitated NH$_4^+$ formation may therefore be an important pathway for nitrate removal and N conservation in anoxic, sulfatic or carbonatic subsoils and sediments poor in organic carbon. The process may explain the disappearance of nitrate from water entering anoxic, Fe(II)-rich layers in sediments where relatively high amounts of exchangeable NH$_4^+$ can be observed immediately above the redox boundary (Hansen, et al., 1996).

Other example is chromium (Cr), which is more related to metallurgy, electropolating or tanning of leather, it has become a common contaminant found in many soils, waste sites, ground and surface waters throughout the world (Richard & Bourg, 1991; Puls et al., 1994). losuaya-Lawniczak et al. (2000) studying the transformation of hexavalent chromium, Cr(VI), a byproduct of many industrial processes, known to be toxic, carcinogenic and soluble, in the presence of GR-SO$_4$ and GR-Cl, observed that both the GRs proved to be very reactive and led to the rapid and complete reduction of Cr(VI) into Cr(III), a very low solubility product and relatively immobile in the environment. Other experiment, evaluating the capacity of GR-CO$_3$ for reducing Cr(VI) into Cr(III), have showed that Cr(VI) was not conserved in the system and was reduced to Cr(III). But, the adsorption or exchange of the Cr(VI) has most likely occurred before reduction (Williams & Scherer, 2001).

In the same way, studies have demonstrated that the higher valent and more soluble selenium forms, Se(VI) reduce to the less reactive Se(IV) and Se(0) forms in the presence of GR. These redox reactions represent an abiotic pathway for selenium cycling in natural environments and can be occurring in the interlayers and on the external surface of GR (Myneni, et al., 1997). Refait et al. (2000) have reported that selenate ions can easily be trapped inside the interlayers of Fe(II)-Fe(III) hydroxysalts favoring the formation of hydroxy selenate, a new GR-SeO$_4$ compound, which can reduce Se(VI) to less mobile Se(IV) forms. As Se compounds are carcinogenic and teratogenic at high concentration, it is important to understand the interactions of SeO$_4^{2-}$ anions with GRs through the formation and transformation of the corresponding GR-SeO$_4^-$, and the concomitant reduction of Se(VI) into Se(IV).

More recently, O’Loughlin et al. (2003), reported that hexavalent uranium, U(VI) (as soluble uranyl ion and with potentially high mobility in the subsurface), was readily reduced by GR-SO$_4$ to U(IV) in the form of relatively insoluble UO$_2$ nanoparticles, suggesting that the presence of GR in the subsurface may have significant effects on the mobility of uranium, particularly under iron-reducing conditions.

In the same way, GR might presumably reduce sorbed arsenate to the more toxic and soluble As(III) species. However, Randall et al. (2001), found that As(V) is reduced to the As(III) form and that it remains adsorbed as (AsO$_4$)$^{3-}$ by forming inner-sphere surface complexes. During the oxidation of GR to lepidocrocite, the (AsO$_4$)$^{3-}$ remains preferentially bound to GR and only sorbed onto lepidocrocite when all of the GR has been oxidized.

Since the oxidation state of polyvalent elements such as N, Se, Cr or U determine their mobility and toxicity in the environment, their interaction with reactive mineral such as GRs could be a clue to the comprehension and control of their behavior (Sposito, 1989). The possible reduction of these elements may be part of a natural process, but it could also be of major importance in soils contaminated with them (Génin et al., 2001). However, the researchers still do not know whether the GR transformed or dissolved the polyvalent elements trapped inside them could be removed, could be reduced to soluble or insoluble forms, could be rendered more mobile or could be definitively fixed (Refait, et al., 2000).

In addition, earlier work has shown that carbon tetrachloride is transformed by abiotic reductive dechlorination with synthetic GR-SO$_4$ as reductant (Erbs et al., 1999). Carbon tetrachloride belongs to a group of halogenated aliphatics, which are released from industrial, commercial and agricultural sources, and they are both highly toxic and carcinogenic as well as prevalent soil and groundwater contaminants. Hence, knowledge of their transformation potential and the products formed are crucial in evaluating their environmental fate. So, GR may be useful for the remediation of chlorinated hydrocarbon contamination in the subsurface.

According to Hansen et al. (1994, 1996) and Roh et al. (2000), the exploitation of GRs to water treatments and in situ remediation of polluted soils and sediments seems possible. In that case, GRs could be applied to transform inorganic and heavily degradable and toxic organic contaminants. Furthermore, it is likely that these reactive iron hydroxides can replace zero-valent iron as a reductant, as they probably have a better penetrating ability in soils and sediments. Remediating
contaminated groundwater with zero-valent metals has been used widely by several researchers.

**CONCLUSIONS**

1. The results of different studies suggest that green rusts play an important role in the fate and transport of many contaminants in suboxic soils, sediments and aquifers where microbial reduction rates are limited by low carbon content.

2. Green rusts have considerable potential in applications such as water-purification processes, gas adsorbers, catalysis and design of drug and pesticide delivery systems.

3. Green rust is only a transient state during the reduction of ferric iron and it is converted to more stable forms, due to this the researchers still do not have answers for many questions, thus, it is still necessary to continue studies about its behavior.

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