Microbial Treatment of Lateritic Ni-ore for Iron Beneficiation and Their Characterization

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Abstract This study aims at studying effect of anaerobic dissimilatory iron (III) reducing bacterial consortium on different phases of iron present in lateritic nickel ore. Such conversion in lateritic nickel ore are helpful in better recovery of sorbed metal values like Ni and Co by subsequent bioleaching or acid leaching. Here properties of thermally and microbiologically reduced lateritic nickel ore are compared vis-à-vis original ore. An anaerobic dissimilatory iron (III) reducing bacterial consortium capable of using glucose as carbon source and lateritic nickel ore as terminal electron acceptor was used for microbial reduction of ore under anaerobic condition. Microbial reduction changes the initial light brown colour of the lateritic nickel ore to dark brown. The change in colour is due to the conversion of goethite to magnetite, which is confirmed from the XRD pattern. The FTIR spectra and the UV-Visible spectra support the presence both goethite and hematite. The study shows that changes in phases brought about by microbial treatment are different than those by thermal treatment. The carrier mediated exchange interaction between Fe²⁺ and Fe³⁺ ions in lateritic nickel ore sample treated with IRB consortium is responsible for higher ferromagnetic ordering. The thermal reduction of the same sample showed lowering of ferromagnetic ordering due to the decreasing percentage of Fe²⁺ and Fe³⁺ ions.

Keywords Iron Reducing Bacteria, Lateritic Ore, Goethite, Magnetite, Magnetic Properties

1. Introduction

In lateritic nickel ore, nickel is associated with iron phase where as cobalt is associated with manganese phase[1]. Natural Fe(III) oxides are high in surface area and are reactive[2]. Fe(III) oxides adsorb a wide range of metal cations and anions by complexion to surface hydroxyl groups[3]. Nickel lateritic ore of Sukinda, Orissa, contains 0.8% Ni and 0.049% Co which makes them the only nickel deposit in India. More focus had been given by IMMT Bhubaneswar on this material for extraction of nickel and cobalt[4-7]. However, thermal activation or thermal reduction of ore is required for better recovery of metal values[8]. Heating of ore resulted in conversion of goethite to hematite, which was responsible for releasing nickel from Fe(II) lattice and resulted in better recovery[7]. This type of reduction is not cost effective in terms of energy consumption and may not be employed for large scale operations. Hence, it is desirable to have an alternative ecofriendly and low cost method, which makes the subsequent metal extraction more feasible.

Ferric iron oxides are widespread in anoxic aqueous environment and have been reported to act a selectron sink during biodegradation of various natural and xenobiotic compounds[9-11]. Microbial Fe (III) reduction results in the generation of several important Fe(II)-containing minerals in sedimentary environments, including magnetite Fe(II)Fe(III)₂O₄, which is a magnetic mineral. Magnetite formation during dissimilatory Fe(III) reduction is reported for different pure cultures of bacteria[12] as well as for Fe(III)-reducing enrichment consortium[13]. Microbial dissimilatory Fe(III) reduction plays an important role in the geochemical cycling of iron and organic matter in anoxic ecological system[14]. Iron-reducing microbes generally belong to the genera Shewanella, Geobacter, Geovibrio, Desulfobulbus etc [15-17]. If bacterial reduction method is successful, cost intensive method of ore roasting process can be eliminated. Such ‘dissimilatory’ processes have opened up new and fascinating areas of research with potentially exciting practical applications.

Applying a biological approach to address this problem, we investigated the use of naturally occurring iron reducing bacteria for the reduction of lateritic nickel ore (iron phase) in a low cost ecofriendly way.
low grade lateritic ore which contains about 0.8% Ni, 0.049% Co, 1.92% Cr, 0.32% Mn, and 50.2% Fe. For the experimental purpose the ore samples were heated at 600 °C for 5 hours to convert the goethite to hematite which is reported to aid the release of the nickel bound to goethite matrix. Thermal activation of the lateritic nickel ore has significant influence on nickel and cobalt recovery[18]. The thermal activation changes the mineral structure and brings about mineral phase transformation by dehydroxylation of the goethite[19].

2.2. Microorganisms

Soil sediment from a wetland sites around Bhubaneswar, Orissa, India was used as source of iron reducing bacteria. After collecting, these were immediately transferred to anaerobic vials and taken to the laboratory for subsequent enrichment of Fe(III) reducing bacteria.

2.3. Enrichment and Isolation of Fe(III)-Reducers

One gram of sediment was transferred to a vial containing 10 ml of anaerobic saline solution. These vials were thoroughly mixed, allowed to settle for 10 minutes and then 5 ml of liquid was used to inoculate 100 ml selective alkaline anaerobic iron reducing medium in volumetric flasks. This medium contains Nickel ore 1.0 g/l; K2HPO4 3.0 g/l; KH2PO4 0.8 g/l; KCl 0.2 g/l; NH4Cl 1.0 g/l; MgCl2 0.2 g/l; CaCl2 0.1 g/l; Yeast extract 0.05 g/l; mixture of vitamins and trace minerals solution 1% (v/v) and 10 mM sodium acetate (C2H3O2Na). pH of the media was adjusted to 7.2 with the help of 1N NaOH. Liquid surface was covered with paraffin oil to make the system anaerobic. The flasks were incubated at 30°C for several days under dark condition. The enrichment procedure was repeated four times with the transfer of 10 ml of culture as inoculum into the fresh medium. The enrichment cultures after five successive transfers were used as inoculum for lateritic nickel ore reduction experiments.

2.4. Treatment of Lateritic Nickel Ore with IRB Consortium

Lateritic nickel ore was treated with well enriched Fe(III) reducing bacterial consortium in an anaerobic, alkaline medium at 2% pulp density and kept at 30°C for 7 days under dark condition. Simultaneously control experiments without any bacterial solution were also run. After treatment, the treated lateritic nickel ore was filtered and air dried. In a similar set of experiments the heated lateritic nickel ore was taken instead of the original ore. The reduction of highly insoluble Fe(III) oxides resulted in dissolution of soluble Fe(II) ions in the growth medium. This soluble ferrous-iron was determined by titration[20].

Samples are designated as O1 for lateritic ore and O2 for lateritic nickel ore sample treated with IRB consortium. Similarly R1 for heated lateritic nickel ore sample and R2 for heated lateritic nickel ore sample treated with IRB consortium.

2.5. Analysis

X-ray diffraction analysis was carried out by an X-ray powder diffractometer (Philips X’pert Pro, Panalytical) using Mo Ka (λ = 0.7107 Å) as X-ray source and a programmable divergence slit. The voltage and current of the x-ray source were 40 kV and 20 mA, respectively. UV-visible absorption spectra of all the samples were taken with a Varian Cary 100 spectrophotometer in the region 200 - 800 nm to determine the mineral species through absorption band gap. Fourier transform infrared spectra were recorded on a FTIR system (spectrum GX model supplied by Perkin Elmer instrument, USA). The spectra were recorded from 400 to 4000 cm⁻¹. The surface morphology was observed in a Field Emission Scanning Electron Microscope (Zeiss Supra 55, observation conditions V—20 kV, I—0.6 nA) after coating the surface with gold to reduce the charging effect. Microscopic image of the IRB consortium was taken in Nikon 80i optical microscope. The room temperature magnetic hysteresis studies of samples were carried out using Vibrating Sample Magnetometer (VSM) (Model: Lakeshore 7410) at an applied magnetic field of 2T.

3. Results and Discussion

Dissimilatory iron reducing bacterial consortium was enriched in an anaerobic mineral salt medium containing lateritic nickel ore composed of insoluble ferric iron in the form of goethite. The medium contained acetate ions as carbon source (electron donor) and ammonium chloride as nitrogen source for bacterial metabolism. In this system, bacteria oxidizes carbon source and the released electron pass through a series of electron carrier molecules to generate ATP, the biochemical energy needed for bacterial metabolism. The ultimate electron acceptor or electron sink in absence of oxygen is the Fe(III) present in goethite. After accepting the electron, Fe(III) gets reduced to Fe(II) which results in phase change. Due to the phase changes, original light brown colour of the lateritic nickel ore transforms to dark brown. No such change was observed in the control experiment.

![Figure 1. Gram staining of iron reducing bacterial consortia](image)
Gram staining of IRB consortium demonstrates the presence of both gram positive and negative bacteria of different morphology. An optical microscopic image shows 6 morphologically different species (Figure 1). Scanning electron micrograph show long curve rod shaped bacteria in close association with ore particles (Figure 2). Bacteria of length up to 6 µm were observed. The bio-film formations around the ore particles were evident. Some granular material could be seen around bacteria which may be extracellular magnetite deposition (encircled in the Figure 2). Similar depositions had been reported by Hansel [21].

The anaerobic IRB consortium grew well in the media with lateritic nickel ore as only electron acceptor and convert Fe(III) to Fe (II) as evident by release of ferrous ion in the medium during growth of consortia (Figure 3). The ferrous ions formed remain adsorbed to the solid ore particles. These Fe(II) ions were detectable only after washing the ore material with 0.1 N HCl. Initially pH decreases to 6 during first 15 days of incubation but after that it remains constant at around 7.4 (Figure 3).

The X-ray diffraction peak is shown in Figure. 4. The XRD pattern of O1 shows the Bragg reflection peaks of goethite as well as hematite. The intensity of hematite peaks increased as the ore was treated with bacterial consortia in anaerobic condition. A small percentage of magnetite peaks was observed in the O2. The bacteria acts as a reducing agent and converts the goethite to hematite and magnetite. Such peaks were lower in the intensity in O1. The intensity ratio of hematite to goethite is in increased ratio in comparison to O1. However the presence of hematite, magnetite represents that the material is an admixture of two iron states i.e. Fe (II) and Fe (III). When the ore O1 is annealed at 600 °C, the goethite phase converts to protohematite with a small percentage of goethite leaving behind. It is already reported in the literature that at around 250 °C, goethite converts to protohematite and at or above 800 °C, it converts to hematite[22]. As our annealing temperature was 600 °C, there is maximum possibility of conversion of goethite to protohematite, which is clearly reflected in the XRD pattern of R1. But the bacteria treated annealed ore (R2) showed improved crystalline peaks of hematite and reduction in the intensity of goethite peaks. The small amount of goethite present in R2 couldn’t be established within the instrumental limitation. This feature is further well supported from the UV-Visible and FTIR studies.

In UV-Visible spectra three absorption bands were observed at the wavelength of 367, 483 and 678 nm for O1 and O2 (Figure 5). The higher wavelength (678 nm) and lower wavelength (367 nm) is attributed to the optical absorptions band of goethite where as the optical band gap at 483 nm is attributed to the absorption bands of goethite and hematite. In R1 and R2, the optical absorption band is observed at 529 nm which completely reflects the presence of hematite only and a sharp absorption band is observed at 367 nm, attributed to goethite. Even in R2, presence of goethite was not observed in XRD due to instrumental limitation, but was confirmed from the UV-Visible spectra. It is well known that the band gap of goethite is higher than the band gap of hematite,
which is well reflected from the shifting of bands in the spectrum.

Figure 5. UV-visible spectrum of original nickel laterite ore and IRB treated samples

From the literature, hematite has six active IR modes of active IR whereas goethite has 36 modes Fe-O vibration modes and 12 hydroxyl group modes of vibration. Well distinct FTIR modes of vibration for goethite and hematite were observed in O1 and O2 (Table 1), whereas the FTIR peak observed in R1 and R2 are attributed to the hematite. Only the peak 465 cm\(^{-1}\) is attributing to the goethite. Some peaks are attributed to the hydroxyl group modes of vibration that may be due to the surface contamination of water during experiment done at open atmosphere. Our well studied XRD, UV-Visible and FTIR analysis predicts that O1 and O2 is a composites of both goethite and hematite whereas R1 is a hematite with small percentages of goethite. R2 is completely a hematite.

Table 1. FTIR wavenumbers and peak assignment for all the samples

<table>
<thead>
<tr>
<th>Sample Specification</th>
<th>O1</th>
<th>O2</th>
<th>R1</th>
<th>R2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wavenumber (cm(^{-1}))</td>
<td>449</td>
<td>466</td>
<td>685</td>
<td>822</td>
</tr>
<tr>
<td></td>
<td>449</td>
<td>465</td>
<td>680</td>
<td>822</td>
</tr>
<tr>
<td></td>
<td>449</td>
<td>465</td>
<td>715</td>
<td>931</td>
</tr>
<tr>
<td>Assignment*[2]</td>
<td>H</td>
<td>H</td>
<td>H</td>
<td>H</td>
</tr>
<tr>
<td></td>
<td>G, H</td>
<td>G, H</td>
<td>Fe-OH</td>
<td>Fe-OH</td>
</tr>
</tbody>
</table>

* where H: Hematite; G: Goethite

The magnetic hysteresis recorded for all the samples are shown in Figure 6. The MH loop recorded for the O1 shows a small loop with remnant magnetization (M\(_r\)) of 0.083 emu/g and coercivity (H\(_c\)) of 311 Oe. No saturation magnetization (M\(_s\)) has been observed even at the applied field of 20000 Oe, whereas the MH loop recorded for R1 shows lower remnant magnetization of 0.021 emu/g and coercivity of 293 Oe. This may be possible due to the transformation of goethite to hematite at 600°C. While O2 shows higher saturation magnetization value of 1.5 emu/g with remnant magnetization value of 0.328 emu/g and coercivity of 235 Oe. This observation provides evidence that the bacteria acts as a reducing agent to convert goethite to hematite in the solution. It is well known that hematite is weakly ferromagnetic at room temperature. R2 shows lower magnetization value of 0.876 emu/g with remnant magnetization of 0.185 emu/g and coercivity of 266 Oe than that of O2. Here the bacteria only act as reducing agent for converting goethite to hematite. Hence the percentage of goethite becomes less in comparison of the hematite. The M\(_s\), M\(_r\), and H\(_c\) value for all the samples are given in Table 2.

Table 2. Saturation magnetization, remnant magnetization and coercive field of all the samples

<table>
<thead>
<tr>
<th>Sl. No</th>
<th>Samples</th>
<th>M(_s) in emu/g</th>
<th>M(_r) in emu/g</th>
<th>H(_c) in Oe</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>O1</td>
<td>---</td>
<td>0.083</td>
<td>311</td>
</tr>
<tr>
<td>2</td>
<td>R1</td>
<td>---</td>
<td>0.021</td>
<td>293</td>
</tr>
<tr>
<td>3</td>
<td>O2</td>
<td>1.54</td>
<td>0.328</td>
<td>235</td>
</tr>
<tr>
<td>4</td>
<td>R2</td>
<td>0.876</td>
<td>0.185</td>
<td>266</td>
</tr>
</tbody>
</table>

Goethite and hematite have iron in 3+valence state, whereas valence state of iron in magnetite is in-between 2+ and 3+. So a well admixture of Fe(II) and Fe(III) in a material results in a ferromagnetic coupling interaction between the iron ions, which enhances the ferromagnetic ordering and can be explained on the basis of RKKY exchange interaction. As observed in O2, Fe exists in the multiple valence state of 2+ and 3+, hence there is a strong possibility of carrier mediated exchange interaction in between Fe\(^{3+}\) and Fe\(^{2+}\) ions via oxygen. Such type of exchange interaction is known as double exchange interaction[23, 24] because the conduction electron transfers from one Fe ions to another Fe ions through oxygen. Such type of carrier mediated exchange interaction improves the ferromagnetic ordering in the system and hence enhances the magnetization. However, only hematite exists in R2 with a very small percentage of magnetite. So the short range exchange interaction between Fe(III) and Fe(III) is dominating over the long range Fe(II) and Fe(III) exchange interaction, thus lowering the saturation magnetization in R2 in comparison to O2. Therefore, the competing exchange interaction between short range ordered...
anti-ferromagnetic coupling in between similar type of Fe ions and long range ordered FM coupling between Fe(II) and Fe(III) ions reduces the saturation magnetization value. This is clearly evidenced from the MH loop of Figure 6.

4. Conclusions

Lateritic nickel ore when subjected to treatment with anaerobic dissimilatory iron (III) reducing bacterial consortium show phase changes in iron minerals. Comparison of properties of thermally and microbially reduced lateritic ore vis a vis original ore show that changes in phases brought about by microbial reduction were different than those by thermal reduction. Magnetite formation was more prominent in case of microbial treatment of original ore than roasted ore which is well pictured in the XRD pattern. Roasting leads to the carrier mediated exchange interaction between Fe(II) and Fe(III) ions in laterite nickel ore sample treated with IRB consortium is responsible for higher ferromagnetic ordering. IRB consortia treated ore showed higher saturation magnetization value of 1.5 emu/g with remnant magnetization value of 0.328 emu/g and coercivity of 235 Oe. The thermal activation of the same sample shows lowering in ferromagnetic ordering due to the decreasing percentage of Fe(II) and Fe(III) ions. Thermally activated ore was less amenable to further phase changes by microbial treatment. From this study we conclude that naturally occurring iron reducing bacterial consortia could be for better recovery of metal values in a cost effective and eco-friendly way.

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