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**J. Description/Abstract**

The objective of this project is to develop a new ironmaking technology based on hydrogen and fine iron oxide concentrates in a suspension reduction process. The ultimate objective of the new technology is to replace the blast furnace and to drastically reduce CO2 emissions in the steel industry. The goals of this phase of development are: the performance of detailed material and energy balances, thermochemical and equilibrium calculations for sulfur and phosphorus impurities, the determination of the complete kinetics of hydrogen reduction and bench-scale testing of the suspension reduction process using a large laboratory flash reactor.

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AISI/DOE Technology Roadmap Program for the Steel Industry
TRP 9953: Suspension Hydrogen Reduction of Iron Oxide Concentrate

Authors:
Dr. H.Y. Sohn

Performing Organization
University of Utah
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Abstract
The objective of this project is the development of a new ironmaking technology based on hydrogen and fine iron oxide concentrates in a suspension reduction process. The ultimate objective of the new technology is to replace the blast furnace and other carbon-based ironmaking processes and the elimination or drastic reduction of the generation of CO2 by the steel industry.

The goals of this phase of the technology development focused on:

- Performing detailed material and energy balances
- Performing thermochemical and equilibrium calculations to evaluate the behavior of impurities such as sulfur and phosphorus
- Determining the complete kinetics of hydrogen reduction of iron oxide concentrates as a function of particle size, temperature, and hydrogen and water vapor concentrations
- Carrying out bench-scale test work on a simulated suspension reduction process using a large laboratory flash reactor.
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Suspension Hydrogen Reduction of Iron Ore Concentrate

Final Project Report

March 2008

Prepared by

H. Y. Sohn
Department of Metallurgical Engineering, University of Utah

Submitted to

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FINAL PROJECT REPORT

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Project Period: 2/23/05 – 12/31/07

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Executive Summary

This project investigated the basic feasibilities of a novel alternative ironmaking technology based on the direct gaseous reduction of fine iron oxide concentrates in a suspension reduction process, with the ultimate objective of eliminating or drastically reducing the generation of CO\textsubscript{2} in the steel industry. The process would use gaseous reducing agents, such as hydrogen, natural gas, a reducing gas generated by partial combustion of coal or waste plastics, or a combination thereof. An important factor considered in developing the novel technology is the large quantities of fine iron oxide concentrates produced in the U.S. that are well suited for suspension reduction. Additionally this technology is to be competitive with the blast furnace/BOF route as well as alternate ironmaking processes. The process can be a part of an overall continuous direct steelmaking process, in which case the product from this process will be collected in a molten state, or the product may be collected in the solid state to be used as a feed to a secondary steelmaking process.

The tasks performed in this project included:

- Detailed material and energy balances, with special attention to carbon dioxide generation from the possible use of carbon-containing fuels (natural gas or coal) for external heating: this analysis showed that the suspension reduction technology using any of the three possible reductants/fuels; hydrogen, natural gas or coal, will require ~38\% less energy than the blast furnace process. The use of hydrogen in the new technology will generate only 4\% of the carbon dioxide produced in the blast furnace process. Even when natural gas or coal is used, there will be significant reduction in carbon dioxide emission from generating; 39\% and 69\% of the blast furnace value, respectively.

- Thermochemical and equilibrium calculations to evaluate the behavior of impurities such as sulfur and phosphorus: This analysis showed that the new process will produce hot metal with substantially lower phosphorus contents, while the sulphur contents will be similar to that in the hot metal produced by the blast furnace.

- Determination of the kinetics of gaseous reduction of iron oxide concentrates (~30 \(\mu\text{m} \) size) as a function of temperature and gas composition including water vapour and CO/CO\textsubscript{2}: This investigation has generated data that iron ore concentrates can be reduced to 90 – 99\% metallization within the few seconds of residence time available in a typical suspension process, provided that the temperature is higher than 1300\degree\text{C}, preferably higher.

- Preliminary bench-scale test work on a simulated suspension reduction process by the use of a large laboratory flash reactor (24 cm diameter, 1.4 m high, max. 1100\degree\text{C}): The results of this test work are substantially consistent with the results
of the rate determination carried out under precisely controlled kinetics experiments.

The results obtained in this project indicate that further development work towards commercialization of the suspension reduction process is justified. The steps to be taken will involve more systematic tests in bench scale, followed by an industrial-scale pilot test.

This new alternative ironmaking technology will have considerable energy and environmental benefits arising largely from the elimination of cokemaking and pelletization/sintering steps in the current ironmaking technologies. The energy savings will be about 38% of the current blast furnace operation or 7.4 GJ/ton of hot metal. The use of coke in the BF generates more than 1.1 ton of CO₂ per ton of hot metal as well as other pollutants, which will be greatly reduced or eliminated, as discussed above.

Unlike other gas-based alternative ironmaking processes based on shaft furnaces or fluidized-bed reactors, the suspension reduction process is a high-intensity process because it will not suffer from the problems that other processes do when operated at high temperatures mainly from the sticking and fusion of particles. The possibility of direct steelmaking in a single unit based on the new process is also attractive. The replacement of blast furnaces requires the new process to be highly intensive, especially when starting with the fine iron oxide concentrate; the suspension process presents the highest potential for a new high-intensity alternative ironmaking technology.
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1. INTRODUCTION

More than 90% of the world's iron is currently produced via the blast furnace process, with the balance produced by Direct Reduction processes (IISI, 2007). Despite the improvements in the operation of modern blast furnaces such as; increasing their life and productivity, decreasing coke dependence, developing environmentally benign cokemaking processes, increasing coal injection, and injecting natural gas and plastics, the BF process suffers from many drawbacks and BF iron production is projected to decrease by 15-20% by 2015 (Fruehan, 1998). The main drivers for this decrease are the requirements for cokemaking and pelletization/sintering and environmental regulations (Ritt, 2000). Tighter regulations primarily affect the coke plant. Another critical factor affecting the use of the blast furnaces is the capital and energy intensities that demand large infrastructure and operation (Gupta and Sathiyamoorthy, 1999). The BF ironmaking is the most energy intensive step in steelmaking and requires serious attention to environmental problems including CO₂ generation. These constraints limit the flexibility of the BF process (Chatterjee, 1993; Manning and Fruehan, 2001). Accordingly, a number of ironmaking technologies have been developed or are under development (U.S. DOE, August 2000b). These technologies aim to allow EAF plants to produce high quality steels utilizing alternative solid and/or liquid iron as scrap substitutes. They will also allow BOF plants to process low-cost raw materials to produce high-quality liquid hot metal, thus avoiding the use of coking-coal. Examples of such alternate ironmaking processes are Direct Reduced Iron (DRI), Hot Briquetted Iron (HBI), and Iron Carbide. All these processes, however, are not sufficiently intensive to replace the blast furnace and they require the pelletization of iron oxide concentrate.
World steel production is expected to increase steadily in the foreseeable future. During this time the generation of premium quality scrap will decrease slightly as industries like automotive and appliances improve manufacturing yields. In addition, recovered obsolete scrap is expected to expand only by 1.6% per year in the near future (New Steel, 1999). As the industry progresses more and more into higher quality steels, the need for low residual metallics will become increasingly important. Consequently, pressure on scrap availability and prices will increase, thus increasing the demand for scrap substitutes. According to a Midrex forecast, the steel industry will require more than a 46% increase in the supply of obsolete scrap and scrap alternatives in the early 21st century (Steel Times, Aug. 1997). World crude steel production has already crossed the 1 billion tons mark. To sustain these levels of steel production, it would be necessary to ensure adequate supply of the required metallics, namely hot metal, scrap and direct reduced iron. An increase of approximately 200 mtpa of metallic production capacity may be required by 2010 as existing blast furnaces reach the end of their lives and as prime scrap is depleted. Because of the uncertain and unreliable scrap supply, the production of scrap substitutes such as DRI, HBI, Iron Carbide and Hot metal should increase to match the steel production demands. All the above-mentioned factors are greatly affecting the transition of ironmaking technology from the conventional method (the BF process) to smaller-scale, more environmentally friendly processes such as direct gaseous reduction and smelting reduction processes.

The previously developed processes for the gaseous reduction of iron oxide can be grouped into two broad types: shaft furnaces [Midrex and HYL (U.S. DOE, 2000b)] and fluidized-bed reactors [FINMET and the earlier FIOR (Brent, 1999), CIRCORED Process (Husain, 1999), and SPIREX (Macauley, 1997)]. These processes, however, are not intensive because they cannot be operated at high temperatures due to, among other factors, the sticking and fusion of particles. Any alternate processes that can replace the blast furnace
must be sufficiently intensive to meet the large production rates required for economic competitiveness.

The shaft furnace processes also require an ore pelletization step and suffer from pellet disintegration problems.

The fluidized bed processes experience a number of additional problems that include:

- Particle sticking, which results in plugging of transfer lines and loss of fluidization,
- Inability to treat fine particles of the concentrate sizes,
- Requirement of several stages because the uniform mixing of particles of different degrees of reduction in the bed, which makes it highly inefficient to attain a high degrees of conversion in one stage (causing high capital costs),
- Requirement of fairly uniform particle size,
- Problems with elutriation loss and dust capture efficiency, and
- No possibility to directly produce molten iron.

A circulating fluidized bed process may be able to better handle fine particles but suffers from many of the same problems plus serious wear and erosion of the reactor and pipes by the high speed flow of gas-particle mixture and non-uniform conversion due to the large differences in resident time for particles of different sizes.

On the other hand, smelting reduction processes encounter the following problems:

- Requirement of pre-reduction, which introduces many of the problems associated with the gaseous reduction processes discussed above and
- Refractory problems caused by the corrosive nature of liquid iron oxide.
Therefore, the AISI steel industry participants (ArcelorMittal, ArcelorMittal-Dofasco, Gallatin Steel, Ipsco, Nucor, Praxair, Ternium, Timken Company and US Steel) and the US Department of Energy commissioned this project. A new transformational alternate ironmaking technology was investigated and basic feasibilities in terms of reaction kinetics and mass/energy balances were defined. The new technology is based on the **direct gaseous reduction of iron oxide concentrates** in a suspension reduction process, with the ultimate objective of significantly reducing energy consumption and environmental pollution in the steel industry. [The term ‘suspension’ is used in this project to represent processes such as the ‘flash’, ‘flame reaction’, or ‘cyclone’ processes.] The process would use gaseous reducing agents, such as hydrogen, natural gas, a reducing gas generated by partial combustion of coal or waste plastics, or a combination thereof. An important factor considered in developing the novel technology is the large quantities of fine iron oxide concentrates currently being produced in the U.S. that are well suited for suspension reduction. An additional factor is the expectation of the development of a hydrogen economy in the U.S. and thus the availability of inexpensive hydrogen.

The new technology is expected to produce sufficiently pure iron that would not require converting and it is to be competitive with the blast furnace/BOF route and other alternate ironmaking processes. The suspension reduction process can be a part of an overall continuous direct steelmaking process, in which case the product from this process would be collected in molten state, or the product may be collected in solid state (e.g. reduced iron pellets or briquettes) to be used as a feed to a secondary steelmaking process.

The technology will have considerable energy and environmental benefits arising largely from the elimination of cokemaking and pelletization/sintering steps in the current predominant ironmaking technology. The energy savings will be about 38% of the current blast furnace operation or 7.4 GJ/ton of hot metal. The use of coke in the BF generates more
than 1.1 ton of CO\textsubscript{2} per ton of hot metal as well as other pollutants, which will be greatly reduced or eliminated by the new technology.

Unlike other gas-based alternative ironmaking processes based on shaft furnaces or fluidized-bed reactors, the suspension reduction technology is a high-intensity process (justification for this expectation is presented in the main text) because it will not suffer from the problems that other processes do when operated at high temperatures mainly from the sticking and fusion of particles. Low-temperature disintegration problems encountered in the processes using pellets can also be avoided. Other expected benefits include the fact that hydrogen reduction will yield iron that does not contain carbon, and thus the iron can directly go through refining without requiring the converting step. The possibility of direct steelmaking in a single unit based on the new process is also attractive. The replacement of blast furnaces requires new processes to be highly intensive especially when starting with the fine iron oxide concentrate, a suspension reduction process presents the highest potential for a high-intensity alternative ironmaking technology.

**Overall Goal**

The goal of this effort was to determine the basic feasibilities of a novel ironmaking technology based on *the direct gaseous reduction of fine iron oxide concentrates* in a suspension reduction process. These include reaction kinetics and mass/energy balances. The technology is to produce sufficiently pure iron that would not require converting. It is to be competitive with the blast furnace/BOF route as well as alternate ironmaking processes. The process can be a part of an overall continuous direct steelmaking process, in which case the product from this process will be collected in a molten state, or the product may be collected in solid state to be used as a feed to a secondary steelmaking process.
2. SUSPENSION REDUCTION PROCESS

**Technology Description**

An important factor in the development of the suspension reduction technology is that the U.S. produces large quantities of iron oxide concentrates that are well suited for suspension reduction. The as-mined ore has to be crushed down to liberate the gangue. These concentrates have to undergo pelletization or sintering process to be utilized as a feed to the BF process. Up to 60% of the U.S. iron production is based on taconite concentrates (50-55 million tons/year) (USGS, 2005), with those from Minnesota being –400 mesh out of the magnetic separation step and those from Michigan being typically even finer at –500 mesh.

The intent of the new process is to produce iron directly from these iron oxide concentrates without going through pelletization or sintering and without the need for coke. The current processes cannot take advantage of the fact that the concentrates are in the form of very fine particles with a large amount of surface area, which would allow rapid reduction by a gas. On a world-wide basis, good quality lump ore is at a premium. Ores from newer reserves also need to be crushed down to finer sizes in order to beneficiate them and make them acceptable feed materials for ironmaking. There is also serious consideration given to the fact that some of the new reserves in regions that traditionally fed sintering plants will now have to make pellet feed in order to bring the gangue content down to acceptable levels. If further tests show that the new process can treat materials –100 mesh or larger, even the sources for coarser iron oxide feed could be used with modest energy consumption for grinding.

The blast furnace process requires the environmentally problematic and energy-intensive cokemaking step as well as pelletization or sintering of the fine iron oxide concentrate. In the suspension reduction process, the concentrate will be reduced to a high
degree of metallization in suspension in a hot reducing gas generated by the external preheating of hydrogen or natural gas or the partial combustion of hydrogen, coal, natural gas, heavy oil, other materials like waste plastics, or a combination thereof. A factor to consider is the expected development of a hydrogen economy in the U.S. and thus the availability of inexpensive hydrogen (U.S. DOE, 2006). The DOE and energy companies are spending billions on a hydrogen infrastructure (see FutureGen family of projects and others). The tremendous amount of activities and accomplishments already made in hydrogen development as well as progress made in the development of ultrasafe nuclear power plants that can provide electric power for the production of inexpensive hydrogen should be noted. There is increasing public pressure for the reduction of greenhouse gas emissions and for imposing penalties on the producers of such gases. The steel industry must be ready with new technologies to take advantage of energy alternatives when they are available. All indications point that in a carbon-constrained world, hydrogen work will continue to be funded. Prior to such, this technology is not solely dependent on the availability of inexpensive hydrogen; other fuels/reductants such as natural gas, heavy oil and coal may also be used, as stated above.

The suspension reduction technology is applicable for either (i) as an ironmaking step, with the product to be melted in a coal-based melter or an electric melter to produce a molten metal feed of different characteristics or fed directly to the secondary steelmaking units such as an electric arc furnace bypassing the converting step like BOF, or (ii) as an integral part of a possible direct steelmaking process from the concentrate (e.g. as illustrated in Figure 1). In the nonferrous industry, a flash furnace similar to that shown in Figure 1 has been the most widely used smelting furnace in which fine concentrate particles are oxidized, form molten droplets, and collect as a bath in the settler. A silicon-free, lower carbon hot metal, as will be produced by this process, is used in the LD-ORP (LD converter-Optimized
Refining Process) of NSC (Nippon Steel Corp.) or in NSC’s converter-based all scrap melting process with coal. One can look at site-specific conditions to decide on the melting of the reduced product, as in the case of the RHF or the current hot connect DRI processes. The suitability of the product of this process for use in BOF and EAF flows against ‘current steel industry’ processes and the results of this work strongly indicate the potential of this transformational ironmaking process to replace the BF/Coke Oven process routes.

Advantages of the New Technology

The new technology will have the following advantages:

- **Energy savings of up to 38%** of the amount required in the blast furnace technology. (Detailed calculations to reach this conclusion will be presented subsequently in this report.)
• Elimination of coke oven and pelletization or sintering step with the associated considerable energy consumption, pollution problems and costs by the direct utilization of the large quantities of fine iron oxide concentrates already being produced in the U.S., which do not need separate comminution (and fine iron ore concentrates that are expected to be produced from newer iron ore deposits for the purpose of significantly reducing the gangue content).

• Significant reduction, or even complete elimination, of carbon dioxide emission from the steel industry, depending on the choice of the reducing agent and fuel used in the suspension reduction technology. (The calculated details will be presented subsequently in this report.)

**Technical Hurdles to be Overcome**

Although the suspension reduction process has many advantages described above, a number of technical hurdles must be overcome before an industrially viable process can be developed.

**Rate of Reduction**

Prior to the start of this project, the opinions of many researchers were that the reduction rates of fine iron ores are not fast enough to be amenable to a suspension reduction process (Thermelis and Zhao, 1994). This conclusion was based on the extrapolation to the much smaller concentrate particle sizes of the previously reported reduction rates collected largely with pellets of millimeter and larger sizes. These are much larger than the concentrate particles to be used in the suspension process. Furthermore, previous work has also been done at lower temperatures, because pellets in shafts or fluidized beds suffer sintering and fusion problems at higher temperatures. These problems will be greatly diminished in the
dilute particle suspension involved in the suspension processes and thus they can be operated at higher temperatures. The published data was carefully examined and it was determined that there were reasons to believe that the reduction rates of the concentrate-size particles might be significantly higher than the extrapolated values. This analysis formed the basis of the investigative work performed in the project. Rate measurements have yielded data that indicate sufficiently rapid reduction rates of iron oxide particles in a suspension process. Thus, the question on whether the reduction rate of concentrate particles is fast enough for a suspension reaction has been resolved in the affirmative.

**Heat Supply**

Unlike sulfide smelting reactions, which are highly exothermic and thus need little or no external heating, the gaseous reduction reaction requires an external heat supply. The heat may be generated internally by burning a portion of the reducing agents, or supplied by plasma or burning of other fuels. These types of processes in which hot reducing gas environment is created internally are used in numerous industrial operations. Examples include the reforming of natural gas and coal gasification by partial combustion. This issue is raised in this report to indicate that it is an important item of consideration but it has a likely solution based on existing technologies. Furthermore, in-depth discussions were held with a burner manufacturer and technology developer resulting in an assurance that this type of a burner to produce high-temperature *reducing* gas mixtures can be designed and fabricated. If necessary in a smaller bench-scale unit, in which heat loss per volume may be high, plasma can be used to augment the heating to compensate for the heat loss.

**Equilibrium Hydrogen Utilization**

The gaseous reduction of hematite to magnetite is essentially irreversible and that of magnetite to wustite also has a large equilibrium constant. Thus, the equilibrium gas products
of these reactions contain little reducing gas, i.e. the degree of equilibrium utilization is very high. However, the final stage of the reduction, i.e. the reaction of FeO with H₂ or CO, is considerably limited by equilibrium. For example at 1000°C,

\[
\text{FeO (s) + H}_2 (g) = \text{Fe (s) + H}_2\text{O (g)} \quad \Delta G^\circ \approx +3000 \text{ cal (1000°C)}. 
\]

This reaction has a slightly positive standard Gibbs free energy, and the equilibrium gas product has a H₂/H₂O molar ratio of 1.5 at 1000°C, i.e. 60 % H₂ and 40 % H₂O when pure hydrogen is used. With pure CO, only 27 % is utilized at equilibrium, although some of the remaining CO can be used to form hydrogen in the presence of water vapor (the water-gas shift reaction). Thus, the product gas from this reaction will contain a substantial amount of unutilized reducing gas. This will require the removal of water from the off-gas and the recycling of hydrogen, and some carbon oxide bleeding, if hydrocarbons are used as the fuel/reductant. This, however, can be carried out in combination with waste heat recovery to generate power or preheat the raw materials.

In addition to the above issues and hurdles, close attention must be paid to other potential problems such as powder feeding, flame control, re-oxidation of reduced iron, and particle accumulation in the reactor during the next-phase project. These will also be some of the major considerations in the design and eventual operation of the industrial-scale pilot tests.

**Research Objectives**

There has been a great deal of work done on the gaseous reduction of iron oxide. However, the previous work has largely been done with pellets or lump ores. Little has been done on direct gaseous reduction of fine concentrate particles of – 400 mesh (– 38 μm) size. Furthermore, even less is known about various factors governing the reduction of such particles in turbulent gas-particle flow in a suspension furnace. Thus, it was necessary to
obtain rate data for reduction of concentrate particles in a few seconds of residence time available in a suspension process. High temperature gas-solid suspension contact processes, such as the flash, flame reaction or cyclone smelting, are well established in industry and can provide a residence time of this magnitude. Further, a preliminary bench-scale test of suspension reduction by gaseous reducing agents was planned, focusing on the method of supplying the energy required to maintain the necessary temperature.

Based on the above discussion, the following project objectives were investigated and successfully achieved:

(1) Performed detailed material and energy balances, with special attention to carbon dioxide generation from the possible use of carbon-containing fuels such as natural gas or coal for external heating.

(2) Performed thermochemical and equilibrium calculations to evaluate the behavior of impurities such as sulfur and phosphorus at various hydrogen and water vapor partial pressures.

(3) Determined the kinetics of hydrogen reduction of iron oxide concentrates (~30 μm size) as a function of temperature and gas composition including water vapour and CO/CO₂.

(4) Carried out preliminary bench-scale test work on simulated suspension reduction process by the use of a large laboratory flash reactor (24 cm diameter, 1.4 m high, max. 1100°C).
3. MATERIAL AND ENERGY BALANCES

Material Balance and Reduction of CO₂ Emission

The input and output streams and quantities for three possible fuel types used with the new technology are given in Table 1, together with those for the conventional BF operation. Of particular interest is the greatly reduced generation of CO₂ for the new technology, even with the use of coal, compared with that from the blast furnace. The new technology will have other environmental benefits from the elimination of cokemaking/burning and the pelletization and sintering steps. Typical amounts of pollutant emission from current BF operations are given in a DOE report (U.S. DOE, 2000c).

Table 1. Material balance comparison between the suspension reduction process and the blast furnace (BF) (for 1 metric ton of molten hot metal).

<table>
<thead>
<tr>
<th></th>
<th>BF [kg/tonHM]</th>
<th>Prop'd(H₂) [kg/tonHM]</th>
<th>Prop'd(CH₄) [kg/tonHM]</th>
<th>Prop'd(Coal) [kg/tonHM]</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Input</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>1430</td>
<td>1430</td>
<td>1430</td>
<td>1430</td>
</tr>
<tr>
<td>SiO₂</td>
<td>151</td>
<td>100</td>
<td>100</td>
<td>125</td>
</tr>
<tr>
<td>CaCO₃</td>
<td>235</td>
<td>162</td>
<td>162</td>
<td>235</td>
</tr>
<tr>
<td>O₂(g)</td>
<td>705</td>
<td>227</td>
<td>411</td>
<td>463</td>
</tr>
<tr>
<td>N₂(g)</td>
<td>2321</td>
<td>749</td>
<td>1354</td>
<td>1525</td>
</tr>
<tr>
<td>C(coke)</td>
<td>428</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H₂(g)</td>
<td></td>
<td>83</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CH₄(g)</td>
<td></td>
<td></td>
<td>211</td>
<td></td>
</tr>
<tr>
<td>Coal (C₁₄H)</td>
<td></td>
<td></td>
<td></td>
<td>301</td>
</tr>
<tr>
<td><strong>Sub-Total</strong></td>
<td>5270</td>
<td>2751</td>
<td>3668</td>
<td>4079</td>
</tr>
<tr>
<td><strong>Output</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fe</td>
<td>1000</td>
<td>1000</td>
<td>1000</td>
<td>1000</td>
</tr>
<tr>
<td>CaSiO₃</td>
<td>273</td>
<td>191</td>
<td>191</td>
<td>257</td>
</tr>
<tr>
<td>Si</td>
<td>5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CO₂(g)*</td>
<td>1671</td>
<td>71</td>
<td>650</td>
<td>1145</td>
</tr>
<tr>
<td>H₂O</td>
<td></td>
<td>740</td>
<td>473</td>
<td>152</td>
</tr>
<tr>
<td>N₂(g)</td>
<td>2321</td>
<td>749</td>
<td>1354</td>
<td>1525</td>
</tr>
<tr>
<td><strong>Sub-Total</strong></td>
<td>5270</td>
<td>2751</td>
<td>3668</td>
<td>4079</td>
</tr>
</tbody>
</table>

*The value for the preparation for BF was not included into this calculation (7%).
Energy Balance

Comprehensive overall material and energy balance calculations have been performed. The calculations followed the procedure established in DOE Report (U.S. DOE, 2000c) and the Steel Energy Savings Estimator Tool [Energetics web page provided by AISI]. The case of using hydrogen in the suspension reduction technology was calculated separately by an identical procedure to that used for the cases of using coal or natural gas that was verified to be consistent with the Steel Energy Savings Estimator Tool, because the Tool does not provide a means to input hydrogen.

Table 2 lists the energy requirements per metric ton of molten hot metal for the blast furnace (BF) and this technology using different types of fuel used, i.e. hydrogen, methane (natural gas), and bituminous coal (considered as C1.4H). It is seen that even with the assumption of full combustion of CO to CO2 in the blast furnace, equivalent to giving full credit for the heating value of the BF off-gas, the suspension reduction technology using any of the three possible reductants and fuels requires a much smaller amount of energy with ~38% less consumption.

It is noted that the comparison in the above table is made against the blast furnace, because the new process is aimed at replacing the blast furnace technology, which is still, by far, the dominant ironmaking technology. No new ironmaking processes have really taken hold, but we also present comparisons with current alternative ironmaking processes in the footnote below Table 2. For those processes that produce solid products, we took the data from the listed references and added the heat of melting so that they could be compared to liquid hot metal. Further, the energy required for grinding ore to the concentrate size is not included because up to 60% of iron production in the U.S. already depends on such concentrates. It is noted that the suspension reduction technology compares favorably with the BF process as well as other commercial alternate ironmaking processes.
This calculation does not take into consideration the amounts of energy required to produce hydrogen, natural gas and coal, because those vary considerably with the method of production and other conditions. Furthermore, an attempt to take these into consideration raises other questions such as “Can the type of energy used for producing a certain fuel be equally applicable to the production of another fuel?” and “Can they be compared directly?” To a certain extent, comparing the energy requirements for processes using carbon-based fuels and hydrogen raises similar questions, but at least their use can be considered as nearly interchangeable as any other two types of energy.

A sample calculation with the Steel Energy Savings Estimator Tool is given in Appendix A.
Table 2. Energy requirement comparison between the technology and the blast furnace (BF) (for 1 metric ton of molten hot metal).

<table>
<thead>
<tr>
<th>Energy required (Feed at 25°C to products)</th>
<th>BF [GJ/tonHM]*</th>
<th>Prop’d (H2) [GJ/tonHM]</th>
<th>Prop’d (CH4) [GJ/tonHM]</th>
<th>Prop’d (coal) [GJ/tonHM]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Enthalpy of iron-oxide reduction (25°C)</td>
<td>2.09</td>
<td>- 0.31</td>
<td>-0.61</td>
<td>1.73</td>
</tr>
<tr>
<td>2) Sensible heat of molten Fe (1600°C)</td>
<td>1.36</td>
<td>1.36</td>
<td>1.36</td>
<td>1.36</td>
</tr>
<tr>
<td>3) Slag making</td>
<td>- 0.21</td>
<td>- 0.15</td>
<td>- 0.15</td>
<td>- 0.21</td>
</tr>
<tr>
<td>4) Sensible heat of slag (1600°C)</td>
<td>0.46</td>
<td>0.32</td>
<td>0.32</td>
<td>0.46</td>
</tr>
<tr>
<td>5) SiO2 reduction</td>
<td>0.09</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6) Limestone (CaCO3) decomposition</td>
<td>0.42</td>
<td>0.29</td>
<td>0.29</td>
<td>0.42</td>
</tr>
<tr>
<td>7) Carbon in pig iron**</td>
<td>1.65</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9) Heat loss and unaccounted-for amounts (assumed the same for all processes)</td>
<td>2.60</td>
<td>2.60</td>
<td>2.60</td>
<td>2.60</td>
</tr>
<tr>
<td>10) Sensible heat of offgas (90°C)</td>
<td>0.25</td>
<td>0.25</td>
<td>0.21</td>
<td>0.20</td>
</tr>
<tr>
<td><strong>Sub-Total</strong></td>
<td><strong>8.70</strong></td>
<td><strong>4.36</strong></td>
<td><strong>4.02</strong></td>
<td><strong>6.56</strong></td>
</tr>
<tr>
<td><strong>Energy value for reductant</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heating value of feed used as Reductant</td>
<td>5.37</td>
<td>7.70</td>
<td>8.00</td>
<td>5.67</td>
</tr>
<tr>
<td><strong>Total for Iron oxide reduction</strong></td>
<td><strong>14.07</strong></td>
<td><strong>12.06</strong></td>
<td><strong>12.02</strong>*</td>
<td><strong>12.23</strong></td>
</tr>
<tr>
<td><strong>Preparation</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1) Pelletizing</td>
<td>2.87</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2) Sintering</td>
<td>0.62</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3) Cokemaking</td>
<td>1.93</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Sub-Total for preparation</strong></td>
<td><strong>5.42</strong></td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><strong>Total for molten hot metal making</strong></td>
<td><strong>19.49</strong></td>
<td><strong>12.06</strong></td>
<td><strong>12.02</strong></td>
<td><strong>12.23</strong></td>
</tr>
</tbody>
</table>

* Gigajoules per metric ton of hot metal [BF numbers compiled based on data from: U.S. DOE, 2000c; U.S. DOE, 2000a; U.S. DOE, 2000d.]

** Carbon in pig represents the heating value of dissolved C. We recognized the fact that its heating value is used in subsequent converting, but decided to leave it in as an energy item because the carbon removal is an added required step that requires other energy and costs, and we are not doing the overall energy balance for entire integrated steelmaking. Even if this item is removed from BF numbers, the suspension process has much lower energy requirement than BF.

***The difference between this number and that for BF agrees closely with the value of 2.3 GJ/metric ton calculated by the use of the Steel Energy Savings Estimator Tool [Energetics web page provided by AISI], shown in Appendix A.

Note 1: COREX 16.9 – 20.2 GJ/TonHM, MIDREX 13.3 GJ/TonHM, and HYLIII 12.3 GJ/TonHM. Heat to melt solid products has been added. (source: L. Price et al., 2002; Wingrove G. et al., 1999.)

Note 2: C and H2 were assumed to be converted to CO2 and H2O, equivalent to crediting their heating values.

Note 3: If the heat recovery is reduced by 20% for the technology, considering its hotter off-gas, the energy consumption for the technology becomes about 13 GJ/ton.

Note 4: The amounts of silica proposed (H2), (CH4) and (coal) were assumed to be, respectively, 70%, 70% and 100% of that for BF. Thus, the same ratios were used for ‘sensible heat of slag’, ‘slagmaking’ and ‘limestone’ values.
4. EXPERIMENTAL APPARATUS AND PROCEDURE

**Materials**

The iron oxide concentrate was provided by Ternium and by ArcelorMittal (from the Minorca mine). The Ternium concentrate screened to 22-30 μm was utilized for preliminary experiments and the Minorca mine concentrate screened to 25-32 μm for kinetics measurement, as shown in Figure 2. Unscreened Ternium concentrate was used as the feed for bench-scale tests to simulate an industrial suspension reduction process. The chemical composition is presented in Table 3. In both cases, most of the iron oxide content was magnetite, which was confirmed by X-ray patterns as shown in Figure 3.

![Figure 2](image)

**Figure 2.** SEM micrographs: (a) unscreened Ternium concentrate; (b) screened Minorca mine concentrate.
Table 3. Chemical composition (wt %) of iron ore concentrate from Ternium and ArcelorMittal.

<table>
<thead>
<tr>
<th>Component</th>
<th>Ternium</th>
<th>ArcelorMittal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Iron</td>
<td>68.4</td>
<td>70.65</td>
</tr>
<tr>
<td>FeO</td>
<td>20.9</td>
<td>30.53</td>
</tr>
<tr>
<td>P</td>
<td>0.029</td>
<td>0.01</td>
</tr>
<tr>
<td>S</td>
<td>0.0055</td>
<td>0.02</td>
</tr>
<tr>
<td>C</td>
<td>N/A</td>
<td>0.24</td>
</tr>
<tr>
<td>Sr</td>
<td>N/A</td>
<td>0.01</td>
</tr>
<tr>
<td>SiO₂</td>
<td>1.73</td>
<td>1.87</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>0.55</td>
<td>0.13</td>
</tr>
<tr>
<td>CaO</td>
<td>0.68</td>
<td>0.27</td>
</tr>
<tr>
<td>MgO</td>
<td>0.40</td>
<td>0.13</td>
</tr>
<tr>
<td>MnO</td>
<td>0.075</td>
<td>0.11</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>N/A</td>
<td>0.11</td>
</tr>
<tr>
<td>K₂O</td>
<td>N/A</td>
<td>0.01</td>
</tr>
<tr>
<td>Na₂O</td>
<td>N/A</td>
<td>0.10</td>
</tr>
<tr>
<td>TiO₂</td>
<td>N/A</td>
<td>0.01</td>
</tr>
<tr>
<td>ZrO₂</td>
<td>N/A</td>
<td>0.03</td>
</tr>
<tr>
<td>CuO</td>
<td>0.021</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Figure 3. X-ray patterns: (a) Ternium concentrate; (b) Minorca mine concentrate.
Preliminary Experiments

Preliminary kinetic experiments were conducted by measuring the weight change of a 22-30 μm concentrate sample over time in pure hydrogen at three different temperatures 900°C, 1000°C, and 1100°C. These experiments were designed to test in a rather simple facility the kinetic feasibility of reducing concentrate particles within a few seconds of reaction time that would be available in a suspension reduction process, before more elaborate and accurate measurements were made.

The concentrate sample was spread on top of a Kaowool compact held in a shallow ceramic tray in order to remove the effect of inter-particle diffusion of the gaseous species. This experiment must be considered rather approximate because the conditions were not fully controlled to determine the fast rate of very fine particle reduction. Possible uncertainties included the effect of radiation from the furnace wall and the sample holder, which would affect the measurement of real temperature of the reacting particle, the effect of mass transfer between the bulk gas and the Kaowool bed, and the effect of minerals in Kaowool on the reactivity of iron oxide. Hydrogen was fed into the reactor (2 cm ID) at the rate of 2 L/min when the desired temperature was reached. The apparatus consisted of a horizontal furnace system, a gas delivery system, a copper turning bed, and an off-gas system as shown below.

Figure 4. Schematic diagram of the horizontal furnace system for preliminary experiments.
**Kinetics Measurements**

For more accurate determination of the reduction rate of individual iron ore concentrate particles from the Minorca mine, a high temperature drop-tube reactor system was fabricated. As shown in Figure 5, this apparatus consisted of a vertical high temperature furnace system, a pneumatic power feeding system, a gas delivery system, a powder cooling and collecting system, and an off-gas outlet system.

The furnace system was made up of a vertical split tube furnace with a maximum working temperature of 1540°C and a cylindrical alumina tube (5.6 cm ID, 193 cm long). A stainless-steel end-cap with five holes was attached to the top of the tube for thermocouples and reactant feeding. Temperatures at various positions of the reactor were measured by the use of two B-type thermocouples with alumina protection tubes and a temperature indicator. Another stainless-steel end-cap with a funnel welded to it to improve the collection of the reacted particles was attached to the bottom of the alumina tube.

The reacted powder was collected in a powder collector on the bottom of the reactor shown in Figure 6 and the unreacted hydrogen and water vapor were discharged through the off-gas outlet system that included a backflow prevention device for safety.

The isothermal hot zone was maintained at a constant temperature of 900 to 1400°C by heating twenty bar-type SiC elements. Carefully measured isothermal zone was 91 cm long within ±20K for typical downward gas flow conditions.
Figure 5. Schematic diagram of the high temperature drop-tube reactor system for kinetic measurements: (a) schematic diagram; (b) photograph.
Figure 6. Powder collection system: (a) schematic diagram; (b) photograph.

**Pneumatic powder feeding system**

Figure 7 shows the pneumatic powder feeding system. The powder feeding system consisted of a syringe pump, a vibrator, a carrier gas line, a powder container, and a powder delivery line. Iron ore concentrate particles were held in a vial sealed with an O-ring. The vial was supported by the piston of a syringe pump which moves at a constant speed. The carrier gas, hydrogen, was fed into the vial (9 mm ID) and exited through a capillary discharge tube (1.2 mm ID) at the top of the powder entraining a small amount of the powder continuously. The whole feeder including the tube between the feeder and the reactor was vibrated by an electric vibrator to keep the feed rate constant and to prevent clogging. Since the stainless steel tube in the reactor should be placed in a high temperature region, it had a double wall for water cooling. The feed rate was calibrated and showed very linear relationship between the infusion rate of the syringe pump and the feeding rate as shown in
Figure 8. During the experiments, it was controlled from 100 to 600 mg/min.

A powder feeding probe in Figure 7 through which the concentrate particles entrained by carrier gas passed was enclosed with a water cooling jacket and inserted into the tube reactor. The tip of the probe was located at the beginning of the isothermal hot zone where the heating of the concentrate particles and mixing with reactant began.

A cylindrical alumina honey-comb was inserted and hung right above the beginning of the isothermal zone as a flow straightener and a heat exchanger. Although every effort was made to improve the heating and mixing of the solid and gaseous species, the reaction zone where the heterogeneous reaction occurs at the designated temperature is only shorter than the isothermal hot zone. Thus, the reduction rate presented in this work could be attained in shorter residence time than the calculated one.
Figure 7. Pneumatic powder feeding system and the powder feeding probe with a water jacket: (a) schematic diagram (b) photograph.

Figure 8. Powder feeding rate vs. advancing rate of the syringe pump.
Residence time determination

The duration of hydrogen reduction of fine iron ore concentrate particles was determined by the residence time (τ) of particles in the isothermal zone. The value of residence time was found from the length of the isothermal zone, which starts from the tip of the powder feeding probe, and the terminal falling velocity for the creeping flow region expressed by the Stokes law (Szekely 1971) assuming that particles fall at a constant velocity in the isothermal zone. The residence time in this work ranged from 2.0 to 8.5 seconds.

\[ u_t = \frac{d_p^2 g (\rho_p - \rho_g)}{18 \mu}, \quad u_p = u_g + u_t, \quad \tau = L / u_p \quad (1) \]

where \( d_p \) = particle size, \( g \) = gravitational acceleration, \( \rho_p \) = particle density, \( \rho_g \) = gas density, \( \mu \) = viscosity of gas, \( u_p \) = particle velocity relative to tube wall, \( u_g \) = gas velocity, \( u_t \); terminal velocity of a falling spherical particle, \( L \); length of the reaction zone.

The terminal velocity depends not only on the particle size but also the temperature due to its effect on the gas density and the viscosity of gas. As temperature increases, the density and viscosity of gas increase and the terminal velocity decreases, as shown in Figure 9. This makes the residence time, namely the reduction time, longer in the same length of the reaction zone when other parameters are fixed. On the other hand, a higher temperature causes the gas velocity to decrease at the same molar rate of gas input, which in turn decreases the residence time. Thus, all these several factors must be taken into consideration when interpreting the effect of temperature on the reduction rate.
% Excess H₂.

A hydrogen-containing gas mixture was fed as the reductant into the reactor concurrently with the pneumatically transported iron ore concentrate, of which the carrier gas was pure hydrogen. To achieve a high degree of iron oxide reduction, excess hydrogen over the stoichiometric amount needs to be supplied. The term % excess H₂ was defined as follows, especially paying attention to the fact that the final stage of the iron oxide reduction, i.e. the reaction of FeO with H₂ or CO, is significantly limited by equilibrium. For example, at 1400°C

\[ \text{FeO}(s) + \text{H}_2(g) = \text{Fe}(s) + \text{H}_2\text{O}(g) \quad \Delta G^o \approx +900 \text{ cal} \quad (1400^o \text{C}) \quad (2a) \]

\[ \frac{p_{\text{H}_2}}{p_{\text{H}_2} + p_{\text{H}_2\text{O}}} = 0.5 \quad (2b) \]
This reaction has a slightly positive standard Gibbs free energy, and the equilibrium gas product has an H₂/H₂O molar ratio of 1, i.e. 50% H₂ and 50% H₂O when pure hydrogen is used as reductant. The equilibrium gas ratio changes somewhat with temperature as shown in Figure 10. Taking the equilibrium composition into consideration, % excess H₂ was defined as follows in this work:

\[\begin{align*}
Fe_3O_4 + H_2 &= 3FeO + H_2O \quad (R1) \\
FeO + H_2 &= Fe + H_2O \quad (R2) \\
\text{Overall: } Fe_3O_4 + 4H_2 &= 3Fe + 4H_2O
\end{align*}\]

\[K_{eq} = \frac{p_{H_2O}}{p_{H_2}} = \frac{n_{H_2O}}{n_{H_2}}\]

Figure 10. Equilibrium gas compositions vs. temperature for the FeO - H₂ and FeO – CO reactions.
Reaction 1 (R1), the hydrogen reduction of magnetite to wustite, has a large equilibrium constant, i.e. essentially irreversible, whereas Reaction 2 (R2) is considerably limited by chemical equilibrium. Thus, the amount of hydrogen \((H_2)_{eq}\) in equilibrium with the water vapor present in the gas was used for the equilibrium constant of Reaction 2 represented by Equation (4). The minimum amount of hydrogen, \(n_{H_2,\text{min}}\), then is the amount of hydrogen used to remove the oxygen from the iron oxide, \(n'_o\), plus the hydrogen required by equilibrium to be present with the water vapor produced by the reduction reaction, \(n'_o/K_{R2}\). The % excess \(H_2\) was then calculated from the total amount of hydrogen fed into the reactor \((H_2)_{\text{total}}\) compared with the minimum amount of hydrogen, \(n_{H2,\text{min}}\), as indicated in Equation (6).

**Degree of reduction**

While varying % excess \(H_2\) from 1300 down to 0, that is, equivalent to the equilibrium gas ratio, the concentrate particles were transported downward, heated, reduced in-flight, and collected. The total iron content in the particles after reduction was determined by use of an inductively coupled plasma (ICP) emission spectrometer (Perkin Elmer, Plasma 400). The fractional reduction was calculated as follows.

\[
\text{Reduction [%]} = \frac{m_o(\%O)_o - m_t(\%O)_t}{m_o(\%O)_o} \times 100
\]  

(7)

where

\[
m_o = m_d(\%Fe)/(\%Fe)_o
\]  

(8)
Here, \( (\%O)_o \) and \( (\%O)_t \) are % oxygen combined with iron in the concentrate before and after the reduction. \( m_t \) and \( m_o \) are, respectively, the mass of a reduced sample used for ICP analysis, collected after reaction for time \( t \), and the corresponding mass of the unreduced dry concentrate. \( (\%O)_o \) was obtained as the difference between 100 and the sum of the total iron content of 70.65% and the total gangue content 3.05% in the ArcelorMittal concentrate in Table 3. The amount of oxygen in the gangue materials which may be reduced by hydrogen was negligible and the weight change due to the possible volatile species such as phosphorus and sulfur was also neglected based on the small contents as given in the table. Weight loss by heating alone was also confirmed to be negligible. \( (\%O)_t \) was calculated from \( (\%Fe)_t \) obtained by ICP analysis, assuming the same weight ratio of iron to gangue.

The oxygen and total iron contents in the ArcelorMittal concentrate listed in Table 3 yielded an O/Fe atomic ratio of 1.30, indicating that essentially all the iron oxide exists as \( \text{Fe}_3\text{O}_4 \) (O/Fe = 1.33). The fact that all iron oxide in the concentrate exists essentially as \( \text{Fe}_3\text{O}_4 \) was further confirmed by the XRD pattern presented in Figure 3.

**Bench-scale Tests**

The kinetic measurements with iron ore concentrate particles from the mine sites proved that the suspension reduction process was kinetically feasible as will be discussed in detail below. Based on these results, preliminary scale-up tests were carried out using a bench-scale test facility shown in Figure 11.
Figure 11. Bench-scale flash furnace for testing suspension hydrogen reduction of iron ore concentrate: (a) schematic diagram; (b) photograph.
The apparatus consists of five subsystems; a vertical furnace system, an electrical power control system, a gas delivery system, a preheating system, and a powder feeding system. A tubular steel reactor (20.3 cm ID, 244 cm long) was electrically heated by six silicon carbide heating elements, which were grouped into two and managed by two SCR controllers. The maximum temperature obtained with the set-up was 1150°C in 76 cm long of isothermal zone.

It is noted that the isothermal zone has a temperature gradient in the radial direction by the way of heating method unlike the drop-tube reactor system for kinetic measurements in which the temperature of the isothermal zone was the same as that of heating elements. For example, to maintain the isothermal zone at 1150°C, the heating elements had to be heated up to about 1250°C, which introduced temperature gradient in the radial direction of the tubular reactor and thus the temperature near the inner wall of the tube was higher than 1150°C. In an industrial scale application, the presence of thermal gradient in the vertical and horizontal directions should be taken into consideration.

All the gaseous species were carefully controlled by flowmeters with high-resolution valves and flowed through a tubular stainless-steel preheater electrically-heated by a horizontal furnace system and packed inside with ceramic Raschig rings to improve mixing and heating of the gas mixture, of which the temperature was measured right before going into the main reactor and was kept at about 500°C on average.

The Ternium concentrate particles were fed into the reactor by a pneumatic powder feeding system at about 1.5 g/min rate. The reduction time of the particles was determined by the residence time of particles in the isothermal zone. In the present work, the residence time varied from 3.5 to 5.0 seconds and % excess H₂ from 0 to ~760%.
5. EXPERIMENTAL RESULTS AND DISCUSSION

**Preliminary Experiments**

About 80% reduction of iron ore concentrate was achieved in 5 seconds at 1100°C in pure hydrogen in the preliminary experiment, which was described in the previous chapter. Reduction was almost completed in 10 seconds at the same temperature, as shown in Figure 12. It was assumed that the weight loss during the reduction occurred only due to the loss of oxygen associated with iron because the components other than iron oxide in the iron ore concentrate undergo little reaction with hydrogen. Weight loss by heating alone was also confirmed to be negligible.

![Figure 12. Approximate hydrogen reduction rate of iron ore concentrate vs. reaction time at different temperatures in preliminary experiments.](image)

It was observed that the reduction of Fe$_3$O$_4$ to FeO and to Fe occurred in succession, as shown in Figure 13, as expected.
Figure 13. X-ray patterns: (a) initial iron ore concentrate (0% reduction); (b) 30% reduction at 900°C for 5 seconds; (c) 50% reduction at 1100°C for 2 seconds; (d) 100% reduction at 1100°C for 10 seconds.

SEM micrographs in Figures 14~16 show that the products became porous as the hydrogen reduction proceeded and that the porosity increased with the reaction temperature.
Figure 14. SEM micrographs: (a) 30% reduced sample after 5 seconds; (b) 100% reduced sample after 60 seconds. (T = 900°C)

Figure 15. SEM micrographs: (a) 32% reduced sample after 2 seconds; (b) 100% reduced sample after 30 seconds. (T = 1000°C)

Figure 16. SEM micrographs: (a) 50% reduced sample after 2 seconds; (b) 100% reduced sample after 10 seconds. (T = 1100°C)
Because of the way the preliminary tests were done, in which certain conditions were left without precise control, the results must be considered as an approximate representation of the actual rates of reduction of individual particles in suspension reduction conditions. These experiments, however, verified the fact that a suspension reduction process with only several seconds of residence time could be feasible for iron production from iron oxide concentrate particles. Further experiments to measure accurate reduction rate of iron ore concentrate were conducted using a drop-tube reactor system, as described below.

**Kinetics Measurements**

In the first series of experiments with hydrogen (with 3-5% N₂) as the reducing gas, over 90% reduction was accomplished in 2.8 seconds and complete reduction in 4.0 seconds at 1200°C, as shown in Figure 17. The results showed a promising kinetic feasibility of the suspension reduction process.

![Figure 17. Hydrogen reduction rate of iron ore concentrate vs. residence time and % excess H₂ at four different temperatures. [p_H2 = 0.85 atm (86.1 kPa)]](image)

(Residence time was varied by the flow rate of hydrogen.)
It is noted that there was a considerable increase in the hydrogen reduction rate when temperature increased from 1100°C to 1200°C, which essentially establishes the lower limit of the process temperature of the suspension reduction process, considering that the residence time would be in the same range.

The % reduction was slightly lower as the residence time was increased above 4 seconds. This is because the residence time was increased in these experiments by reducing the excess H₂ used and at a lower % excess H₂ the effect of water vapor produced by the reduction reaction increased. Water vapor not only lowers the partial pressure of hydrogen but also decreases the thermodynamic reducing power of the gas due to the equilibrium limitation of the FeO-H₂ reaction system ($K_{R2} = 1$ at 1400°C).

This point can be further understood by considering the following simplified global rate expression for iron oxide reduction:

$$ R = k_{app} \cdot f\left(\frac{p_{H_2}}{p_{H_2O}}\right) = k_{app} \left(\frac{p_{H_2} - p_{H_2O}}{K_{R2}}\right) $$

Thus, the presence of water vapor lowers $p_{H_2}$ and also increases the negative term in the parentheses.

Further experiments were performed at temperatures higher than 1200°C to obtain temperature effects. Above 1300°C, complete reduction was accomplished in less than 2.0 seconds in large excess hydrogen, as shown in Figure 18. Complete reduction was achieved even when excess hydrogen lowered to 40%, which increased the residence time to 8 seconds. In an industrial application, it would be advantageous to operate the reduction process at a high rate with lower excess hydrogen input.
Thus, additional experiments were conducted paying attention to the experimental conditions for moderate excess hydrogen and residence time. At 1200°C, complete reduction was already achieved in 4.0 seconds at 470% excess hydrogen, as shown in Figure 19. As the residence time decreased, the reduction degree decreased accordingly. At 1300°C, 90% reduction was achieved in 4 seconds at 200% excess hydrogen, as shown in Figure 20, which was 10% higher than that at 1200°C at the same residence time and % excess hydrogen. When the residence time was increased to 6 seconds, reduction was already completed at 200% excess hydrogen (meaning complete conversion was achieved between 4 and 6 seconds). When the residence time further increased to 8.3 seconds, the complete reduction began to be observed with only 40% excess hydrogen.
Figure 19. Hydrogen reduction rate of iron ore concentrate vs. % excess H₂ at 1200°C.

Figure 20. Hydrogen reduction rate of iron ore concentrate vs. % excess H₂ at 1300°C.
Figure 21. Hydrogen reduction rate of iron ore concentrate vs. % excess H₂ at 1400°C.

The temperature increase to 1400°C did not result in much increase in reduction rate, comparing Figures 20 and 21. This is believed to be due to the melting of FeO (m.p. ~1380°C) as seen in Figure 22, which prevents the formation of cracks in the particles seen at lower temperatures. Reduced iron also forms dense layer at this temperature rather than porous layer as at low temperatures. Figure 23 shows the comparison of the XRD patterns of the concentrates reduced at 1300 and 1400°C for 2.7~2.8 seconds with four different levels of excess H₂. At 200% excess H₂, there was little change in the reduction rate between 1300 and 1400°C as seen in Figure 23-(b). A weak dependence of the reduction rate on temperature may be expected if the chemical kinetics have become sufficiently rapid at these increased temperatures so that pore diffusion and mass transfer between the bulk gas and particle control the overall rate. Our calculations show, however, the rate controlled by pore-diffusion + mass-transfer is much greater than the measured rates, i.e. under this condition a 30 μm particle would be fully reduced in milliseconds compared with a few seconds as observed in
this work. Thus, the weak increase of rate on temperature between 1300 and 1400°C is attributed to the change in particle morphology during the reduction reaction. Once the particle and intermediate product (m.p. of Fe₃O₄ = 1597°C) form a fully molten phase at higher temperatures, the reduction rate is expected to increase with temperature more strongly.

Figure 22. SEM micrographs: (a) 100% reduction at 1300°C; (b) 100% reduction at 1350°C.
Some agglomeration of concentrate particles were at times observed at low excess hydrogen and long residence time. Agglomeration was avoided by completely drying the concentrate and increasing the flow rate of gaseous species. This dispersed the concentrate particles as soon as they came out of the powder feeding tube inside the reactor. The agglomeration of fine concentrate particles lowers the reaction rate and thus should be avoided. Turbulence of the gas-particle flow is expected to be much higher in a large scale operation. Thus, the problem of agglomeration would be much less.
The findings from the current work conclusively show that nearly 100% reduction is obtained with 40% excess H\textsubscript{2} in 8.3 seconds and 90% reduction with 200% excess H\textsubscript{2} in 4 seconds at 1300\textdegree C. At 1400\textdegree C and 200% excess H\textsubscript{2}, 80% reduction was attained in only 2.7 seconds. Even at 1200\textdegree C, 96% reduction was reached in 5.7 seconds with 200% excess H\textsubscript{2}. The variables of the suspension reduction process in an industrial application should be determined carefully considering the competing factors, the cost of excess hydrogen and energy for high temperature, and their effect on the total reduction rate.

A temperature higher than 1400\textdegree C would be definitely preferable from the kinetics point of view and moderate % excess H\textsubscript{2}. This would be required in any case if the reduced iron is collected as a liquid. Furthermore, incompletely reduced iron ore concentrate can be further reduced in the molten bath by injecting additional reducing agents into it.

**Bench-scale Tests**

Larger scale suspension reduction tests were conducted in the bench-scale reactor described in chapter 4. Again, the highest temperature that could be generated in this reactor was only 1150\textdegree C.

The trend in the change of the reduction rate was observed at three different residence times; 3.5, 4, and 4.5 seconds in which the reduction extent was at 21, 29, 43%, respectively, with 0% excess H\textsubscript{2} and approached about 95% with 720% excess H\textsubscript{2}, as shown in Figures 24 and 25. The reduction extent at a longer residence time was always higher while the excess H\textsubscript{2} ranged from 0 to 720% and only 0.5 second difference made a notable change in the reduction rate.
Based on the results from the bench-scale tests and the feasibility tests, it is obvious that the operating temperature of the facility needed to be increased by 100–200°C to obtain a sufficiently high reduction rate with smaller excess H₂ (0–50%) and with increased fine concentrate feeding rate (1–5 kg/hr). A technical issue that must be overcome is the heat supply. The heat may be generated internally by burning a portion of the reducing agents, or supplied by plasma or burning of other fuels. These types of processes in which hot reducing gas environment is created internally are used in many industrial operations. Examples include the reforming of natural gas, coal gasification by partial combustion, and the Horsehead Flame Reactor Process for treating EAF dusts. The main effort in the current work included addressing this issue in addition to determining the feasibility of achieving high degrees of reduction in a simulated suspension process.
Figure 25. Reduction rate of iron ore concentrate vs. residence time at different % excess hydrogen.

Although hydrogen is the best and cleanest reductant and/or fuel from the environmental and reduction kinetics viewpoint, it is currently expensive. Instead, the syngas which is mainly composed of H₂ and CO from the reforming of natural gas or the coal gasification has been used as a reducing gas mixture in the majority of direct reduction processes hitherto developed. Thus, the use of syngas as a reductant was examined in the bench-scale facility.

The syngas was simulated with a mixture of H₂, CO, and N₂ while keeping the compositional ratio the same as that obtained by mixing hydrogen with a combustion gas from an internal gas burner. The input amount of each component was calculated based on material and energy balances considering the preheat temperature of gaseous species, the heat of reaction between CH₄ and O₂ in the burner, the temperature change of gaseous species by supplying additional hydrogen to improve the reduction rate.
For example, Figure 26 shows the equilibrium composition of the product gas from a mixture of 1 kmol CH₄ and 1 kmol O₂, the adiabatic temperature of which is about 2500°C.

![Equilibrium Composition](image)

**Figure 26. Equilibrium compositions of product gas from a mixture of 1 kmol CH₄ and 1 kmol O₂.**

Once the amounts of the product gases from the reaction between CH₄ and O₂ are determined, the amount of additional H₂ (added to improve the reduction rate) and the residence time are calculated under the fixed feeding rate of concentrate (1.5 g/min) and the reaction temperature (e.g. 1150°C), as shown in Table 4. For this calculation, it was assumed that the additional H₂ was preheated to 500°C based on the measurement of the preheat temperature of the gas entering the main reactor.

Under the conditions shown in Figure 24, about 90% reduction was accomplished in 3.5 seconds at 800% excess hydrogen (32.7 kPa H₂, 53.4 kPa N₂). The % reduction remained similar when 11.6 kPa of N₂ was replaced with the same amount of CO. It should be noted that the average barometric pressure of Salt Lake City is 86.1 kPa. When 10% (7.6 kPa) of H₂ was replaced with CO, the conversion decreased from 90% to 80% even at about
Table 4. Mass and energy balances in the reduction on the concentrate

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
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<td>2.571</td>
<td>2.571</td>
<td>2.828</td>
<td>2.314</td>
<td>2.314</td>
<td>0.257</td>
<td>1.5</td>
<td>19.2</td>
<td>3.96</td>
</tr>
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<td>1.571</td>
<td>1.571</td>
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<td>1.414</td>
<td>1.414</td>
<td>0.157</td>
<td>1.5</td>
<td>11.0</td>
<td>4.61</td>
</tr>
<tr>
<td>1.143</td>
<td>1.143</td>
<td>1.257</td>
<td>1.028</td>
<td>1.028</td>
<td>0.114</td>
<td>1.5</td>
<td>7.4</td>
<td>4.98</td>
</tr>
</tbody>
</table>

4.5 seconds residence time. Although there was a decrease in reduction rate by introducing syngas instead of H₂, it is still of interest especially at higher operating temperatures. As temperature increases, the equilibrium ratio of H₂/H₂O and CO/CO₂ changes in the opposite way, i.e. the thermodynamic reducing power of H₂/H₂O increases whereas that of CO/CO₂ decreases. The amount of hydrogen in a typical syngas of direct reduction process is larger than that of carbon monoxide and thus the thermodynamic reducing power of the syngas would rise at higher temperature. In other words, more H₂ can be replaced with CO. Furthermore, the water-gas shift reaction would convert much of CO to H₂ by reacting with water vapor generated by the iron oxide reduction by H₂.

Large scale tests in this type of bench-scale reaction or an industrial pilot reactor at considerably higher temperatures are needed to complete the commercial feasibility of this suspension reduction technology. Nevertheless, the bench-scale tests performed in this project combined with the kinetic measurements made in the drop-tube reactor have clearly demonstrated the kinetic feasibility of the process.
6. Sulfur and Phosphorus Behavior

Sulfur and phosphorus contents in hot metals produced by the two ironmaking processes – blast furnace process and the Suspension Reduction process – were calculated for comparison. The calculation was done with the same iron ore with chemical composition of 0.04wt% S, 0.04wt% P, 64wt% Fe, 6wt% SiO₂ and 2wt% Al₂O₃.

For blast furnace process, a coke rate of 530 kg per ton of hot metal was assumed, and the chemical compositions of the coke were: 0.65 wt% S, 0.035 wt% P, 4 wt% SiO₂, 3 wt% Al₂O₃ and C as the remainder. A flux of 131.1 kg CaO and 43.7 kg MgO per ton of hot metal was also added to get a slag with proper basicity and viscosity for smooth operation and sufficient S removal. The equilibrium distribution ratio of S between the resultant slag with a basicity of $B = 1.458$ ($B = \frac{\%CaO + 0.7\%MgO}{\%0.94SiO₂ + 0.18\%Al₂O₃}$) and the hot metal was calculated to be $\frac{\%S}{\%S} = 129$ according to the reported relation between equilibrium distribution ratio of S and slag basicity. Since S distribution ratio in actual blast furnace process is around 47% of equilibrium distribution ratio, the actual distribution ratio of S was taken as $129 \times 47\% = 60$, from which value the mass balance calculation for S leads to the S content in hot metal by blast furnace process to be $\%S = 0.02$.

Blast furnace practice indicated that over 90% of P load enters into hot metal and the rest is lost either in the slag or in the flue gas or is absorbed in the refractory linings. Assuming 90% of P enters into hot metal, mass balance calculation for P leads to the P content in hot metal by blast furnace process to be $\%P = 0.108$.

For the suspension reduction process, a slag with higher basicity and thus higher liquidus temperature can be used for better S removal, since the process has to be operated at a higher temperature to deal with low carbon hot metal. The slag basicity chosen for this calculation was $B = 1.9$. Another critical parameter which determines S distribution ratio is
the FeO content in slag or the H₂/H₂O ratio in atmosphere. The lower the FeO content in slag, or the higher the H₂/H₂O ratio in atmosphere, the higher the S distribution ratio and thus the better S removal. Thermodynamic calculations indicated that 1 wt% and 10 wt% of FeO in slag (or 54 and 5.4 of H₂/H₂O ratio in atmosphere) will result in 12 and 1.2 of S distribution ratio, respectively, which correspond to [%S] = 0.013 and [%S] = 0.045, respectively, in hot metal by the suspension reduction process.

The predicted S and P contents in the iron produced from the suspension reduction process as functions of FeO content in the slag are plotted in Figure 27 below:

![Figure 27. Calculated S and P contents in hot metal from suspension reduction process (lines) compared with that from blast furnace (points).](image)

The results presented above for the suspension reduction process do not include the possible additional removal by vaporization of the impurities. Thus, the actual values are expected to be lower, which needs to be verified by more comprehensive calculations. These calculations do indicate, however, the suspension reduction process may be able to produce a hot metal with P and S contents lower than or comparable to that by the blast furnace process.
7. CONCLUSIONS

A novel alternative ironmaking technology for producing iron directly from fine concentrates by a gas-solid suspension reduction that bypasses cokemaking and pelletization/sintering was thoroughly investigated. The material and energy balance analysis indicate that the suspension reduction process would reduce energy consumption by about 38% of the amount required by the blast furnace and drastically lower environmental pollution, especially CO2 emission, from the steel industry. As a result of concept definition tests with the iron ore concentrate, the reduction rate of fine concentrate particles by hydrogen-containing gases was determined to be sufficiently fast for a suspension technology. From the kinetic measurements and preliminary bench-scale tests, it was found that the reduction temperature should be higher than 1200°C to obtain a sufficiently high reduction rate at moderate excess H2. The use of syngas from the reforming of natural gas or coal gasification as part of the reducing gas mixture in this new process was shown to be feasible.

Further kinetic experiments, under a comprehensive bench-scale testing program are recommended to determine more definitively the relationship among reaction temperature, residence time and % excess gaseous reactant required to achieve 90~100% reduction. In addition, further bench-scale tests with hydrogen and/or syngas are recommended at selected conditions from these additional kinetic experiments, especially at temperatures in the range of 1300~1600°C to fully simulate the method of supplying the energy required to maintain these temperatures, as an intermediate step in determining the scalability for a larger, industrial-scale suspension reduction process pilot test.

The replacement of blast furnaces requires new or improved alternative ironmaking processes that are highly intensive, especially when starting with the fine iron oxide concentrate; the initial project results indicate the Suspension Hydrogen Reduction of Iron Oxide concentrate process presents the highest potential for a high-intensity blast furnace replacement technology.
REFERENCES


Chatterjee, A., Beyond the Blast Furnace (Boca Raton, FL: CRC press), 1993, 1-6.


Appendix A

Summary of calculation with the Steel Energy Savings Estimator Tool [Internet tool provided by AISI] for the suspension reduction technology using natural gas (compared with blast furnace operation excluding cokemaking and pelletization/sintering).

Summary

Project Name: Suspension ironmaking

User Inputs

<table>
<thead>
<tr>
<th>Process: blast furnace</th>
<th>% Energy Savings Natural Gas: 452.0%</th>
</tr>
</thead>
<tbody>
<tr>
<td>% of Market Impacted: 20.0%</td>
<td>% Energy Savings Electricity: 0.0%</td>
</tr>
<tr>
<td>Annual Growth Rate: 2.5%</td>
<td>% Energy Savings Coal/Coke: 100.0%</td>
</tr>
<tr>
<td>Year of Introduction: 2012</td>
<td>% Energy Savings Fuel Oil: 100.0%</td>
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<tr>
<td>Market Penetration: 10 Year</td>
<td>Market Solid or Liquid Wastes: 0 lbs per tons of product</td>
</tr>
<tr>
<td>Curve: Saturation</td>
<td>Non-Combustion Air Pollutants: 0 lbs per ton of product</td>
</tr>
</tbody>
</table>

Energy Impacts for blast furnace

(Based on the input provided above, this technology will have the following impacts:)

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<th>2005</th>
<th>2010</th>
<th>2015</th>
<th>2020</th>
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</thead>
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<tr>
<td>MARKET PENETRATION</td>
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<td>N/A</td>
<td>14.7%</td>
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<tr>
<td>MARKET (Million tons)</td>
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<td>N/A</td>
<td>4</td>
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<td>ENERGY SAVINGS</td>
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<td>N/A</td>
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<td>N/A</td>
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<td>Fuel Oil Energy Savings (trillion Btu)</td>
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<td>N/A</td>
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<tr>
<td>TOTAL ENERGY SAVINGS</td>
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<td>POLLUTANT REDUCTIONS (lbs)</td>
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<td>Particulates</td>
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<tr>
<td>Other (million lbs)</td>
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<td>N/A</td>
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*equivalent to 2.3 GJ/metric ton iron, in agreement with our energy calculation shown separately in the text below.

USER EXPLANATIONS

Technology Description: New technology using natural gas

Market Percentage: 20 million ton/yr iron, which is 20% of steel production

Introduction Year: (Bench + Pilot) testing plus commercialization lead time

Energy Impacts Percentages: Natural gas is used as the reductant and fuel.

Other Wastes and Pollutants: None
The baseline input values for the blast furnace operation were obtained as follows, by adjusting the data given in the table below (imported from "Energy and Environmental Profile of the U.S. Iron and Steel Industry," U.S. DOE, Report No. EE-0229, August 2000) to fit into the only four categories provided in the Steel Energy Savings Estimator Tool.

1) Coal/Coke: Coke + Coal + other gases – Top gas credit = 10.7 + 1.5 + 1.5 – 4.0 = 9.7
2) Natural gas: 1.8
3) Electricity: 0.2
4) Fuel oil: 0.4

**Energy items in a typical blast furnace operation (1998 values) (excluding energy for cokemaking, sintering, and pelletization)**

<table>
<thead>
<tr>
<th>Fuel</th>
<th>Specific Energy Use [10^6 But/short ton hot metal]</th>
<th>Specific Energy Use [GJ/metric ton hot metal]</th>
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<tr>
<td>Coke</td>
<td>10.7</td>
<td>12.44</td>
</tr>
<tr>
<td>Coke oven gas</td>
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<td>0.23</td>
</tr>
<tr>
<td>Blast furnace gas</td>
<td>1.0</td>
<td>1.16</td>
</tr>
<tr>
<td>Natural gas</td>
<td><strong>1.8</strong></td>
<td>2.09</td>
</tr>
<tr>
<td>Fuel oil</td>
<td>0.4</td>
<td>0.47</td>
</tr>
<tr>
<td>Coal</td>
<td>1.5</td>
<td>1.74</td>
</tr>
<tr>
<td>Electricity</td>
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<td>0.23</td>
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<tr>
<td>Oxygen</td>
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<td>0.35</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td><strong>16.1</strong></td>
<td><strong>18.72</strong></td>
</tr>
<tr>
<td>Top gas credit</td>
<td>4.0</td>
<td>4.65</td>
</tr>
<tr>
<td><strong>Net total</strong></td>
<td><strong>12.1</strong></td>
<td><strong>14.07</strong></td>
</tr>
</tbody>
</table>

The input values for the technology using natural gas were as follows:


   Thus, % energy savings natural gas = \((1.8 - 9.93)/1.8\)*100 = – 452%

2) Electricity: 0.2
3) Coal/Coke: 0
4) Fuel oil: 0

It is not possible to use the Steel Energy Savings Estimator Tool [Internet tool provided by AISI] for the case of operating the suspension reduction process with hydrogen, because there is no provision for hydrogen in the input categories. Thus, the energy savings for the case of using hydrogen was separately calculated by the same procedure as for the case of using natural gas with this technology, which was verified to be consistent with the Steel Energy Savings Estimator Tool, and shown in Table 2 in the main text.