The present invention refers to an effective method for minimizing the problems of iron ore pellet degradation by weathering during their stockpiling, i.e., by providing an appropriate method for improving the state of the art with regard to iron ore pellet resistance related just to the hydration process of the slag phase. Thus, in order to minimize hydration in the slag phase, stabilizers are introduced into the mixture used to produce iron ore pellets prior to being heat-treated.

Figure 8

Avenida das Nacoes Unidas, 12.995, 18º andar, CEP: 04578-000, Sao Paulo - SP (BR).
Descriptive Report of the Invention Patent for "AGING (WEATHERING) INHIBITION METHOD OF IRON ORE PELLETS DURING STOCKPILING"

INVENTION FIELD

This present invention intends to use additives to prevent the loss of iron ore pellet strength during stockpiling.

INVENTION BACKGROUND

It is well-known that iron ore pellets are made through processes in which iron ore is mixed with certain additives to prepare a chemical composition suitable for disk pelletizing or rotating drums. The resulting pellets are then taken to kilns where they are burned to become resistant to handling and suitable for use in reduction reactors. In fact, there are countless advantages of using iron ore pellets as liquid steel, especially as it generates fewer fines during handling and within the reduction reactor, and less slag is produced in relation to other liquid steel components, particularly sinters. However, with regard to fines generation, monitoring the physical quality of some kinds of iron ore pellets has historically shown an increasing loss of physical resistance, from its production until its use, including stacking time, time spent in the yards, and transportation. The degradation of the iron ore pellets' physical quality entails:

- more fines generated when received by the customer;
- loss of pellets' performance in the reduction process;
- risk of contractual non-compliance;
- restriction in pelletizing plants' productivity with a significant loss in
It is also well known that a major cause of degradation in the iron ore pellets' physical quality is due to weathering, resulting from their interaction with moisture and other environmental agents. For this reason, rain water and the water used to cut down on particulate emission have a strong influence on the aging cycle frequency. However, so far, a truly effective mechanism has not been found that would cut down on the hydration process with subsequent slag phase solubilization during iron ore pellet stockpiling.

**BRIEF DESCRIPTION OF THE INVENTION**

Therefore, in order to minimize the aforementioned problems, the present invention introduces aging inhibitors into the iron ore pellet production mixture, prior to heat treatment in an oxidizing atmosphere, to reduce hydration in the slag phase.

More specifically, the invention aims to minimize the pellets' degradation problems caused by weathering during stockpiling, i.e., envisioning an appropriate method for improving the state of the art with regard to the pellets' loss of resistance brought about by the slag phase's hydration process while being stockpiled.

Therefore, the present invention comprises a method to protect iron ore pellets during their slag phase aging process, including the phase during which hydration reaction inhibitors for the slag phase of heat-treated iron ore agglomerates are added.

In the preferred embodiment of this present invention's method, inhibitors are
added to the pulp before the cold agglomeration process (pelletizing or microagglomeration), and aging inhibitors include metal oxides.

The metal oxides are preferably selected from a group of materials consisting of alumina, kaolinite (Al₂O₃·2SiO₂·2H₂O), fine silica, titanium oxides, Mg, and Zn, but they may also be selected from any other metal oxides that reduce the silicate's K, Na, and Ca content.

The metal oxides may be introduced and tested at an increasing rate until a maximum limit has been reached, depending on the product's desired chemical stability. Preferably, this shall be carried out up to a minimum limit to which the slag's composition in the post-burning agglomerate simultaneously meets the conditions of \((\text{CaO} + \text{Na}_2\text{O} + \text{K}_2\text{O}) < 45\%\) and

\[
\frac{(\text{SiO}/60) + (\text{Al}_2\text{O}_3/102)}{(\text{GaO}/56) + (\text{Na}_2\text{O}/62) + (\text{K}_2\text{O}/94)} > 0.6
\]

The inhibiting material shall be applied as particulate material <45 μm or preferably 80% <20 μm to ensure the stabilizing elements' greater reactivity and integration in the slag.

Furthermore, the inhibiting materials may be added as powder or diluted in the slurry.

Finally, the invention's method may be applied to other types of iron ore agglomerates whose resistance depends on the type of calcium silicate and iron in the slag phase.
DESCRIPTION OF FIGURES

Figures 1a and 1b depict the results of the microstructural analysis of the iron ore pellets that were submitted to hydration in water under the present invention:

- Figure 1a depicts the hydration effect on the pellet's surface;
- Figure 1b depicts the hydration's evolution over time;

Figure 2 depicts the corrosion mechanisms of soda glass in water.

Figure 3 depicts the steps in the chemical etching process involving the leaching and dissolution of the synthetic slag network in a type of iron ore pellet.

Figures 4a and 4b depict the effect of the incorporations and oxides on the corrosion of soda-lime glass in water.

Figure 5 depicts the effect of the chemical composition of the synthetic slag phase of some types of iron ore pellets on the dissolution process.

Figure 6 depicts the variation of compression strength in some types of industrially produced iron ore pellets.

Figure 7 depicts the peptization process flowchart.

Figure 8 depicts the granulometric distribution of the oxide source materials that inhibit aging in the iron ore pellets produced on a pilot scale.

Figure 9 depicts the incorporation of metal oxides by means of the materials added to inhibit the hydration process in iron ore pellets produced on a pilot scale.

Figure 10 depicts variations in the compression strength of fired iron ore
pellets produced with and without addition of inhibiting metal oxides in the hydration process.

**DETAILED DESCRIPTION OF INVENTION**

The following detailed description is not intended in any way to limit the invention's scope, applicability, or configuration. More precisely, the following description provides arrangements for implementing the exemplary modalities. By using the instructions provided herein, those who are skilled in the art will recognize suitable alternatives that can be used without going beyond the invention's scope.

Iron ore degradation due to weathering during stockpiling and transportation is a serious problem faced by a number of iron ore producing companies. Therefore, the present invention aims to advance the state of the art by proposing solutions that have not yet been achieved in relation to iron ore pellet aging, specifically with regard to the slag phase, in which moisture from ambient humidity or rain presents a great challenge due to the severe loss of iron ore pellet resistance.

The initial investigations focused on understanding the aging mechanism in the slag phase. Toward this end, industrial pellets collected in the layer over the grid, both top and bottom, were investigated. These pellets were cut and submerged in deionized distilled water at room temperature for 60 days. Pellets with etching in smaller time intervals were also looked at to evaluate the phenomenon's evolution. From there, the effects of moisture on the pellets' surface, hydration evolution over time, and the hydration reaction residue of the hydration were examined.
The results of this analysis are summarized in Table 01 below as regards the characterization of the hydration reaction residue and in Figures 1a to 1b:

**TABLE 01**

<table>
<thead>
<tr>
<th>XPS (%)</th>
<th>EAIP (mg/l)</th>
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<tbody>
<tr>
<td>C 21,68</td>
<td>C -</td>
</tr>
<tr>
<td>O 55,50</td>
<td>Ca 24,50</td>
</tr>
<tr>
<td>Si 19,18</td>
<td>Mg 0,138</td>
</tr>
<tr>
<td>K 1,82</td>
<td>Fe &lt; 0,013</td>
</tr>
</tbody>
</table>

Analyzing the residue obtained by immersing the pellet in water and water evaporation

Analyzing the aqueous solution in which the pellet was immersed. There were no precipitating compounds.

- The aqueous solution: Si (90.70 mg/l), Ca (24.50 mg/l), Cl (1,16 mg/l), K (1,15 mg/l), and Mg (0.138 mg/l). The Fe and Al appear in insignificant amounts in terms of concentration.

- The compounds formed by the evaporation of the aqueous solution: SiO₂, CaCO₃, Na₂CO₃, Si₆Ca₂O₁₀, and Si₆Ca₂O₁₂ H₆.

* Photoelectron Spectroscopy Excited by X-ray (XPS) determines the atomic compounds in nano-materials

Inductively coupled plasma atomic emission spectroscopy (ICP-AES) (EAIP)

And the results show that:

- The product from the hydration process was a compound based on Si and Ca (tobemorite) without the presence of Fe ions, indicating the assumption that it was caused by the calcium silicate hydration and not the Ca ferrites.

- Based on the reaction below, calcite crystals are formed by the reaction of
calcium hydroxide generated in this reaction with the $\text{CO}_2$:

$$3[2(\text{CaO}.\text{SiO}_2)] + 3,5\text{H}_2\text{O} \rightarrow 5\text{CaO}.2\text{SiO}_2.2,5\text{H}_2\text{O} + \text{Ca(OH)}_2$$

$$\text{Ca(OH)}_2 + \text{CO}_2 \rightarrow \text{CaCO}_3 + \text{H}_2\text{O}$$

- The partial leaching of calcium silicate increased the pellet's intergranular porosity, thus enhancing the progress of its physical weakening or loss of mechanical strength.

- The growth and nucleation of calcium carbonate crystals were more pronounced in the first 10 days of contact with moisture.

Once the mechanisms involved in dissolution of the slag were detected, i.e., that aging is caused by decomposition or partial leaching from the glass binder phase, including all varieties of calcium silicates, when exposed to ambient moisture or rain, the factors that influence these mechanisms were then investigated.

Academic publications produced in recent decades pertinent to the glass industry suggest that the corrosion mechanisms of soda-lime glass by water take place according to the schematic drawing in Figure 2. In the first stage - reaction (a) - there is an exchange between sodium ($\text{Na}^+$) and potassium ($\text{K}^+$) ions of the glass and hydrogen ions in the solution, and in the second stage - reaction (b) - the main connections ($\text{Si-O-Si}$) break down, causing the glass structure to dissolve.

Figure 3 shows the evolution of the leaching of a synthetic silicate, with a composition similar to a type of iron ore pellet. In Figure 3, there is:
- a peak highlighted as A, which indicates the S-peak, i.e., the O-Si-O-Si-O binding as regards the increase in surface roughness indicating chemical etching of the glass surface without the formation of a protective layer (network dissolution);

- a peak highlighted as B relative to the NS-peak, i.e., indicating an increase in the modifying cation content on the glass surface, which may be related to the deposition of salts, such as Na$_2$CO$_3$ and CaCO$_3$, arising from severe corrosion;

- a peak highlighted as C related to the hydration band, i.e., indicating the presence of water in the glass structure since the first day and well-developed on the third day.

Note that this glass tends to be continuously dissolved by the presence of an aqueous solution, given that the formation of an SiO$_2$-rich layer was not found, which is a protective mechanism against corrosion.

In the case of soda-lime glass, the incorporation of the alkaline earth oxides or other divalent or trivalent oxides to the glass considerably increases the chemical resistance to water, according to figures 4a and 4b.

The same evolution profile can be seen in the aging test for iron ore pellets. In this test, the effect of the synthetic slag phase iron ore pellet composition on the hydration process, or aging, was also evident, as shown in Figure 5. From the results, it can be seen that:

- The type C pellet slag was more resistant to dissolution since the first day.

- The dissolution of the glass from the B type pellet showed an initial
behavior similar to the C type, with the possible development of a protective layer on subsequent days, approaching the behavior of C type pellet.

These results are consistent with industry practice, as shown in Figure 6, and strongly correspond with the pellets' relative loss of strength, confirming the strong influence of the slag's composition on the degradation phenomenon through aging.

Accordingly, the aim of the present invention is to act in the slag phase in order to minimize hydration that takes place during iron ore pellet stockpiling. In other words, the main objective of the present invention is to provide an efficient method for stabilizing the composition in the pellets’ slag phase, thereby minimizing the hydration reactions and stabilizing them in the weathering processes, thus inhibiting the pellets' aging and loss of physical resistance.

Therefore, in order to minimize hydration in the iron ore pellets’ slag phase, a process was developed involving the addition of stabilizing compounds to the mixture prior to its heat treatment. More specifically, the aging inhibitors are introduced into the slag phase composition. More specifically still, the aging inhibitors consist essentially of metallic oxides that have been selected according to the recommendations found in Figures 4a and 4b, in particular Ba, B, Si, Zr, Al, and Zn. More preferably, the ones that have minimal impact on the quality required for the use of iron ore pellets in reduction reactors are recommended, such as Al and Si. Theoretically, there are no limits on the amount of these additives since they enter the silicate composition as modifying elements. In the literature, there are glasses with up to 18% $\text{Al}_2\text{O}_3$. Thus, the maximum amount to be dosed to inhibit aging in the slag phase is limited by the pellet's desired chemical quality. In
other words, the amounts of these oxides should be as small as possible, so as not to result in significant changes in the pellets' chemical compositions. The result will largely depend on the metal oxide content in the indicated source of such materials and the assimilation kinetics of these oxides in the slag phase, and also mainly on the granulometric distribution of the materials used as the source of these oxides to ensure their reactivity and integration in the slag. Ideally, these source materials should be 100% below 10 μm to minimize the amount of material to be used. But nothing prevents the size from being greater than that, depending on other characteristics that could influence the reactivity, such as porosity, grain size, and others, and the iron ore pellets' specifications with regard to the added elements.

The source materials for the age inhibiting metal oxides can be added to the iron ore mixture in any form, for instance, in an aqueous solution or as dry material (powder). The dosage should be carried out using the customary equipment for these applications. Given that ultrafine particulate material can be partially removed from the pulp during the thickening and filtration stages, it is recommended that the dose be added between the filtration and the pelletizing stages (Figure 7). It should be noted that, due to the small amount of the additive relative to the ore mass, it is important that the homogenization stage ensures its thorough distribution in the mixture to assure the least possible deviation in the aging inhibition effect.

The proposed solution was tested on a pilot scale and the results confirmed those from the bench scale. In these tests, the performance of four kinds of materials was evaluated, three (3) rich in Si and Al oxides and 1 (μη) rich in silicon as shown in Table 02 below, in two sets of experiments. Table 02 shows the chemical composition of the sources of the age inhibiting metal oxides produced on
a pilot scale:

<table>
<thead>
<tr>
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<th>SHA1</th>
<th>SHA2</th>
<th>SHA3</th>
<th>SHA4</th>
<th>SI1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe₂O₃</td>
<td>2.14</td>
<td>1.27</td>
<td>1.32</td>
<td>-</td>
<td>1.36</td>
</tr>
<tr>
<td>SiO₂</td>
<td>45.6</td>
<td>45.3</td>
<td>46.1</td>
<td>43.93</td>
<td>92.3</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>36.3</td>
<td>36.9</td>
<td>37.4</td>
<td>36.63</td>
<td>0.55</td>
</tr>
<tr>
<td>CaO</td>
<td>0.057</td>
<td>0.057</td>
<td>0.057</td>
<td>0.020</td>
<td>1.12</td>
</tr>
<tr>
<td>MgO</td>
<td>0.017</td>
<td>0.017</td>
<td>0.020</td>
<td>0.040</td>
<td>0.26</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.199</td>
<td>0.3</td>
<td>0.32</td>
<td>-</td>
<td>0.22</td>
</tr>
<tr>
<td>TiO₂</td>
<td>2.32</td>
<td>1.69</td>
<td>1.95</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>MnO</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.054</td>
<td>0.180</td>
<td>0.161</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.041</td>
<td>0.041</td>
<td>0.041</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>PF</td>
<td>13.8</td>
<td>14.3</td>
<td>14.1</td>
<td>13.17</td>
<td>-</td>
</tr>
</tbody>
</table>

The granulometric distribution of these materials is shown in Figure 8. Figure 9 shows the metal oxide incorporations by means of the materials added to inhibit the hydration process. Figure 10 shows the variation in resistance for fired pellets after stockpiling, with and without adding the inhibitors' source materials. It can be concluded from these results that the use of hydrated aluminum silicate (SHA) in a dosage from 0.3% to 0.7% proved to be effective in inhibiting the iron ore pellet aging process and the use of these inhibitors had the negative effect of increasing the amounts of SiO₂ and Al₂O₃ in the fired pellet in the order from 0.10% to 0.3% at the used dosages.

In the above detailed description, the invention was described with reference
to specific procedures. However, it is apparent that a number of modifications and changes may be made without departing from the invention's scope as set forth in the above claims.

While different types of equipment, systems, and methods for confirming the use of an oral device have been described, it will be apparent to a skilled technician that many other methods and implementations are possible within the scope of the accompanying claims. Therefore, the apparatus, systems and methods to confirm the oral use of a device should not be restricted to the appended claims and their equivalents.
CLAIMS

1. A method to protect iron ore pellets during their slag phase aging process, CHARACTERIZED by the fact that it includes the phase during which hydration reaction inhibitors for the slag phase of heat-treated iron ore agglomerates are added.

2. In accordance with Claim 1, a method CHARACTERIZED by the fact that the inhibitors are added to the pulp before the cold agglomeration process (pelletizing or microagglomeration).

3. In accordance with Claim 1, a method CHARACTERIZED by the fact that the aging inhibitors include metal oxides.

4. In accordance with Claim 3, a method CHARACTERIZED by the fact that aforementioned metal oxides are selected from a group of materials that consist of aluminum, kaolinite \((\text{Al}_2\text{O}_3\cdot2\text{SiO}_2\cdot2\text{SH}_2\text{O})\), fine silica, titanium oxides, Mg, and Zn.

5. In accordance with Claim 3, a method CHARACTERIZED by the fact that the aforementioned metal oxides are preferably selected from any other oxides that reduce the silicate's K, Na, and Ca content.

6. In accordance with Claim 3, a method CHARACTERIZED by the fact that the aforementioned metal oxides shall be introduced and tested at an increasing rate until a maximum limit has been reached, depending on the product's desired chemical stability.

7. In accordance with Claim 6, a method CHARACTERIZED by the fact
that the aforementioned metal oxides shall be introduced and tested at an increasing rate, preferably to the effect that the slag's composition in the agglomerate after firing simultaneously meets the conditions of \((\text{CaO+Na}_2\text{O+K}_2\text{O}) < 45\%\) and

\[
\frac{(\text{SiO}_2/60) + (\text{Al}_2\text{O}_3 \times 0.2)}{(\text{CaO}/56) + (\text{Na}_2\text{O}/62) + (\text{K}_2\text{O}/94)} > 0.6
\]

8. In accordance with Claim 7, a method **CHARACTERIZED** by the fact that, at the end of the process, the pellets have a chemical quality that is acceptable in the iron and steel market.

9. In accordance with Claim 6, a method **CHARACTERIZED** by the fact that the inhibiting material shall be applied as particulate material <45 \(\mu\) or preferably 80% <20 \(\mu\) to ensure the stabilizing elements' greater reactivity and integration in the slag.

10. In accordance with Claim 9, a method **CHARACTERIZED** by the fact that the inhibiting materials may be added as a powder or diluted in the pulp.

11. In accordance with Claim 1, a method **CHARACTERIZED** by the fact that the method may be applied to other types of iron ore agglomerates whose resistance depends on the type of calcium silicate and iron in the slag phase.
Figure 1a

- Macroscopic view
- Optical microscopy
- MEV

- EDS analysis of the crystal
- CaCO₃ – calcite
- Pellet from the top layer with 60 days of chemical etching.

Figure 1b

- The crystals' nucleation and growth was fast during the first 10 days, but afterwards it slowed down considerably.
- Lower intensity in the pellets' protection layers and at the pellet core.
Figure 2

Figure 3
Figures 4a and 4b
<table>
<thead>
<tr>
<th>Oxide (%)</th>
<th>Pel A</th>
<th>Pel B</th>
<th>Pel C</th>
</tr>
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<tbody>
<tr>
<td>SiO₂</td>
<td>41.74</td>
<td>46.32</td>
<td>40.95</td>
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<tr>
<td>CaO</td>
<td>38.80</td>
<td>31.88</td>
<td>32.16</td>
</tr>
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<td>Fe₂O₃</td>
<td>15.55</td>
<td>11.12</td>
<td>17.58</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>3.24</td>
<td>3.78</td>
<td>4.78</td>
</tr>
<tr>
<td>MgO</td>
<td>0.56</td>
<td>3.19</td>
<td>1.37</td>
</tr>
<tr>
<td>K₂O</td>
<td>nd</td>
<td>0.25</td>
<td>0.24</td>
</tr>
<tr>
<td>MnO</td>
<td>nd</td>
<td>0.16</td>
<td>0.71</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>nd</td>
<td>2.61</td>
<td>1.73</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.12</td>
<td>0.70</td>
<td>0.47</td>
</tr>
</tbody>
</table>

Figure 5
Figure 6
Figure 7
1\textsuperscript{st} series of experiments

2\textsuperscript{nd} series of experiments

Figure 8
1st series of experiments

2nd series of experiments

Figure 9
1st series of experiments

2nd series of experiments

Figure 10
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER
INV. C22B1/24
ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. Fileds searched
Minimum documentation searched (classification system followed by classification symbols)
C22B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
EPO-Internal, WPI Data

C. Documents Considered to be Relevant

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* Special categories of cited documents:
- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier application or patent but published on or after the international filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed
- "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
- "S" document member of the same patent family

Date of the actual completion of the international search: 12 November 2012
Date of mailing of the international search report: 22/11/2012

Name and mailing address of the ISA/
European Patent Office, P.B. 5818 Patentlaan 2
NL-2280 HV Rijswijk
Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016

Authorized officer: Swi atek, Ryszard

Form PCT/ISA/210 (second sheet) (April 2005)
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