Processing and Characterization of Self-Reducing Briquettes Made of Jarosite and Blast Furnace Sludges

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Abstract
Jarosite sludge coming from the hydrometallurgical zinc production route is a hazardous material, which is currently neutralized and landfilled by the so-called Jarofix® process. The present study aims to assess the mechanical and metallurgical properties of briquettes made of jarosite powder with blast furnace sludges, acting as a reductant material, to recover the iron oxide in the form of pig iron and produce an inert slag, increasing the recovery of materials considered as wastes nowadays. Starch was used as a binder (0, 5, 10 wt%), and two different briquetting pressure levels were used (20 and 40 MPa). The results show that briquetting without a binder is not desirable, as the agglomerating forces provided by pressure only are not sufficient, as the briquettes are very fragile and not handy. The binder addition increased noticeably the briquettes resistance, however, only little distinction between the 5 and 10 wt% levels were seen. The briquetting pressure, on the other hand, showed a bigger role on the cold mechanical properties of the bound briquettes. The briquettes pressed at 40 MPa reached an average compressive strength higher than 12 MPa and good abrasion and drop resistance were seen, also showing that their production with starch as a binder is feasible. A special remark is done regarding the roasting treatment of the jarosite powder before the briquetting process, as an undesirable compound (thenardite) was formed within some briquettes due to a non-uniform heating of the powder, which hindered the briquettes mechanical properties. Metallurgical properties open the possibility to use such briquettes for iron production in cupola furnaces.

Graphical Abstract

Keywords Jarosite · Blast furnace sludges · Starch · Briquetting · Briquettes mechanical properties
Introduction

Jarosite is produced as a part of the jarosite process in zinc hydrometallurgy. In particular, the jarosite process allows the precipitation of iron and its following filtering, which leads to an increase of zinc extraction yield. Zinc and iron are present as sphalerite [(Zn,Fe)S] [1, 2]. Therefore, due to the competitive formation of zinc and iron sulfides, an excess of sulfuric acid is used for the leaching process, which leads to hydrosulphide and the iron precipitation as hydronium jarosite. Other jarosite compounds can be formed due to the presence of K⁺, Na⁺, NH₄⁺, and Pb in the leaching solution, such as natrojarosite [NaFe₃(SO₄)₂(OH)₆], ammoniojarosite [NH₄Fe₃(SO₄)₂(OH)₆], and plumbojarosite [PbFe₆(SO₄)₄(OH)₁₂]. These compounds are considered hazardous, due to the presence of Cd, As, Zn itself, and Pb. The following possible formation of complex groups with the main jarosite elements could lead to soil contamination and acidification [3, 4].

Due to the high content of iron (40%–50%) present in the jarosite, and the possible further presence of zinc, copper, and zinc, several efforts have been taken to recover these elements. Moreover, the recovery of these elements would transform the jarosite from a hazardous material to a non-hazardous one, making its disposal easier from both economic and environmental point of view [5, 6].

The Jarofix® process is still one of the best technologies for neutralizing the jarosite leachable residues: the filtered jarosite is initially mixed with lime and then with Portland cement. This mix reacts with itself during time, producing inert elements, such as ferric oxyhydroxide [Fe₂O₃·3H₂O, Ca₆Fe₂(SO₄)₃(OH)₁₂·nH₂O] and sodium sulfate, that further reacts generating gypsum. Moreover, the Portland cement reacts to form Ca–Al–Fe silicate hydrate phases and Ca–Al–Fe oxide. These new products are much more stable and inert than the initial jarosite, resisting through years of physical and chemical testing, reducing the long-term environmental issues, such as water and soil contamination, allowing the disposal of the jarosite wastes into inert material landfills [7].

Thanks to the promising high-strength cement of the jarosite and Portland cement mix, it could be possible to be applied for the production of roads and airfield runways [8]. Moreover, the use of jarosite in the Portland cement in alternative to natural gypsum would lead to a decrease in the economic cost of the overall cement production [9].

The annual production of jarosite wastes are about 6 million tons per year [10] and even if the Jarofix® and the jarosite addition to cement are two possible solution, the recovery of valuable elements like iron is not exploited by these methods. Furthermore, the Jarofix® process requires several hectares of landfilling spaces in order to deposit the processed material, which has limited working life [11, 12]. Therefore, the development of new techniques for the recovery of valuable materials from the jarosite is a challenging research field.

Zhu et al. studied a pyrometallurgical route for separation of heavy metals from jarosite residues, consisting mainly of sodium jarosite, with 28wt% iron, 12.78wt% sulfur, and 3.76wt% zinc, and traces of Pb and Cd. Through a two-step heating process, more than 93% of iron was transformed in pig iron, whereas sulfur was reduced up to 0.05wt% [13]. Similarly, Mombelli et al. also studied the jarosite sludges reduction, using an arc-transferred plasma reactor. The main jarosite compound was the natrojarosite and plumbojarosite. The pig iron obtained after the heat treatment showed high amounts of S, P, and Cu (> 1 wt%), with a structure close to a white cast iron. Moreover, the slag formed at the end of the process consists of SiO₂–Al₂O₃–Fe₂O₃ glass, and does not show any leaching behavior, making it suitable for disposal at inert waste landfills or to be used in civil construction. Note that still, the dust coming from the arc transferred plasma furnace can be recirculated to the Roast–Leach–Electrowin (RLE) process [14].

Although coke can be used for the jarosite wastes reduction so as to obtain pig iron, several problems are associated to its production: the high amount of green-house gas emissions, the negative health impact to the population near the production plants, and the high cost associated to the coal mining are just some of them [15–18]. Alternative reductant agents, such as H₂ and syngas, have been proposed in order to reduce the coke consumption; however, their high cost hinders their use [19]. One possible solution is the use of blast furnace wastes with high amounts of carbon content, such as the blast furnace sludges, which have already provided satisfactory results regarding the reduction of bauxite tailings red mud [20], BOF dust [21], and low-grade iron ores [22].

The reduction of jarosite to produce pig iron by the use of blast furnace sludges has been studied by Mombelli et al. [23]. The carbon contained in the sludge allowed the reduction of the iron oxide contained both in the sludge and jarosite at a temperature of 1500 °C. In particular, the best C/Fe₂O₃ ratio found was 0.261 and the main advantages of the process were the formation of a vitrified structure of the slag, making it stable against toxic element leaching, and a metal/charged material ratio higher than 30%, confirming the applicability of the process from both an environmental and economical point of view [23].

However, both the jarosite and the blast furnace sludges come in the form of fine powder that can be easily carried away or lost, so the material handling can be cumbersome. Therefore, they may require agglomeration techniques, such as briquetting, to achieve both the handling and safety
requirements. Briquetting could be explored to allow easier handling of these materials, together with binder to better increase the physical properties of the product. Encouraging results have shown the applicability of briquettes, as alternative or additional charging material, in both Midrex® process and cupola furnaces [24–26].

The main aim of the presented work is to produce and characterize novel jarosite/blast furnace sludges self-reducing briquettes with sufficient cold strength and metallurgical properties to be used in a thermal reactor, such as a Midrex® or a cupola furnace, without material loss during handling. The binder chosen for the briquetting is corn starch. Starch is a polymer classified as an organic binder, and it can be used as dry matter or as a gel. This leads to a disruption of the intermolecular bonds, generating hydrogen-bonding sites, increasing the number of water molecules that can be attached, increasing solubility in cold water [27]. Starch is used in the briquetting of charcoal [27–29] as well as it can be used as a substitute of bentonite in the pelletizing of steelmaking by-products (blast furnace sludges, BOF dusts, and mill scale) [30–33].

### Materials and Methods

#### Raw Materials

The jarosite used in the present study, previously characterized by Mombelli et al. [14], is a greenish powder (mean diameter from 2 μm to 5 μm) mainly composed of natrojarosite and plumbojarosite. The chemical compositions of materials used are reported in Table 1.

The powder was roasted in a muffle furnace at 950 °C for 60 min to remove hydroxide species and impurities. Most of the jarosite species decomposed into hematite, increasing the iron oxides concentration from 33 to 55 wt%, and reducing considerably the S content from 8.86 to 2.55 wt%, decomposed in the form of SOx gas phases. After roasting, the powder appearance turned from greenish (Fig. 1a) to rusty (Fig. 1b); the phase modifications were identified through XRD analysis and the results are shown in Fig. 2. A more detailed description of the roasting process is discussed by Mombelli et al. [14]. After roasting, the jarosite powder was ground in a Retsch Planetary Ball Mill PM 400 machine for

![Fig. 1 Jarosite before (a) and after (b) roasting; blast furnace sludges (c) and lime (d) used for the manufacturing of the briquettes](image-url)
15 min at 300 rpm using zirconia balls. Due to the coarsening and hardening of the powder after roasting, the powder was sieved through a 125 µm sieve in order to obtain a more homogeneous dimensional distribution. The sieved powder was then dried overnight at 105 °C in a muffle furnace.

The blast furnace sludges composition is given in Table 1, also in this case they were previously characterized by Mombelli et al. [23]. The sludges appeared as fine dark particles (Fig. 1c), with very low mechanical resistance agglomerates, easily crushable without the need of any dedicated machinery. As for the jarosite powders, the blast furnace sludges powder was sieved through a 125 µm sieve and dried at 105 °C overnight in a muffle furnace. The results of the XRD analysis (Fig. 2c) highlighted the presence of mainly hematite and magnetite with residual amount of unburnt coke (amorphous halo at 25° 2θ) that partially graphitized after the exposure at high temperature. Traces of reacted non-ferrous compounds (carnegite and diopside) were also observed, probably originated from mineral reactions of gangue and limestone [34–36].

The jarosite and blast furnace sludges were then mixed with a weight proportion of 2.75:1, in order to achieve a C/Fe2O3 mass proportion of 0.261, which was the optimal ratio for the reduction of the iron contained in the jarosite found by Mombelli et al. [23]. In order to improve the slag basicity index, 5 wt% of lime over the total jarosite mass used was added (Table 1). Before mixing, lime was dried at 550 °C for three hours to decompose the eventual hydroxide. The chemical composition of the final mix (labeled JBFS) is shown in Table 1.

The binder chosen for the briquetting is corn starch, the XRD pattern of the starch (Fig. 2d) is in accordance with the literature [37–39]. The corn starch was gelatinized by heating at 80 °C (starch to water proportion 1:6) for 20 min so as to increase its viscosity and obtain higher binding properties. Two different corn starch–JBFS mixes were prepared and differentiated in accordance with the amount of starch added to the JBFS dry basis: 5 and 10 wt%, respectively. Furthermore, some of the JBFS were not mixed, and used as a control batch to highlight the starch binding effects for the briquettes production. The resulting mix of JBFS and starch is labeled SJBFS.

### Briquetting

Briquetting of the JBFS and the SJBFS mixes were performed by an Instron uniaxial tensile test machine upgraded with a self-designed briquetting machine (Fig. 3b). The briquetting machine (Fig. 3a) is composed of a funnel (internal diameter of 20 mm) suitable for housing the powder and threaded to a cylinder. A spring for chrome-vanadium molds supports an ejector that is able to move downwards during the briquetting process. When the machine is activated, a plunger, powered by the machine actuator, starts to press the powder inside the funnel. Once the pressure required for the spring deformation is achieved, the ejector begins to descend and the powder is pushed against the actuator plunger, the pressure is then increased until the chosen value is achieved. Then the pressure is gradually removed and the spring moves the ejector upwards together with the briquette; finally, the funnel is unthreaded, and the briquette is manually removed.

More than 40 briquettes were produced following the described briquetting process with a target height of 20 mm. About 15 g of JBFS mix was needed in order to obtain the target height; on the other hand, the mass needed varied for the SJBFS mixes based on the binder wt%. Two different briquetting pressure levels (20 MPa and 40 MPa) were used with a compressing speed of 20 mm/min. A dwell time of 120 s was adopted to promote the entrapped gases release and increase the briquette physical properties. The resulting briquettes were then left to dry in ambient temperature until the briquette moisture content did not change over time. In order to provide an internal traceability, the briquettes were labeled according to the pressure and binder amount used, as shown in Table 2.

### Physical Characterization

As the briquettes were ready for the mechanical tests, they were submitted first to dimensional analysis to estimate how the bulk density changes according to the different factor levels. The statistical software used for the ANOVA was Minitab®.

The briquettes were then submitted to three mechanical tests to assess the mechanical properties of the produced briquettes. Three briquettes of each B (5 wt% starch) and C (10 wt% starch) levels were sent to compression test, two of C level to the drop test while six of the B level to the drop test, two of the C level and three of B level to the abrasion test, and one of each level to the decrепitation test. Table 3 summarizes test type and total specimens number. Metallurgical properties, namely low-temperature disintegration, swelling, and reducibility were performed on the briquettes showing the best cold strength, namely B+ combination. Two samples were used for each tests.

After each test, a specimen is submitted to X-Ray Diffraction to assess the elements inside the briquettes, and in some cases, a scanning electron microscope (FE-SEM Zeiss SIGMA 500, Oberkochen, Germany) equipped with EDS probe (Oxford Xmax, Abingdon-on-Thames, UK) is used to better understand the compounds formed within the briquettes.

X-ray diffraction was performed by means of a Rigaku Smartlab SE (Tokyo, Japan) diffractometer equipped with XRF suppression 0D/1D detector (D/Tex 250). Manually
Fig. 2  XRD pattern of a jarosite, b roasted jarosite; c blast furnace sludges and d corn starch used for manufacturing the briquettes.
homogenized powders through an agate mortar were scanned from 5 to 90° 2θ at 5°/min and excited by a copper tube (Cu-Kα radiation; λ = 1.54 Å). Samples were spin at 120 rpm.

Cold Compression Test

The Compressive Strength of the briquettes were performed inspired by the BS ISO 4700:2015, used for assessing pellets strength, as there are currently no standards for testing steel-making briquettes. The briquettes were loaded between 2 flat plates, and a pre-load of 30 N was placed before the test. The test then started at a constant velocity of 15 mm/min between the plates, and it was finished when the load falls by 50% from the recorded maximum value, or when the gap between the plates was reduced by more than 50% of the briquette diameter, which in the present case is 10 mm.

Tumbler and Abrasion Test

The abrasion test was done with a tumbler of diameter 180 mm and length 285 mm with two L-shaped lifters of 50 mm flat × 50 mm height employed, with the tumbler rotating at 50 rpm. Abrasion and tumbler indexes were calculated according to the BS EN 3271–2015 standard, as reported in Eq. 1 and 2

\[ TI = \frac{m_1}{m_0} \times 100 \]  

\[ AI = \frac{m_0 - (m_1 + m_2)}{m_0} \times 100, \]

where \( m_0 \) is the mass of the test portion as weighed out and placed in the tumble drum; \( m_1 \) is the mass of +6.70 mm fraction of the tumbled test portion; \( m_2 \) is the mass of −6.70 mm +500 μm fraction of the tumbled test portion.
The briquettes were put together in pairs, and after 200, 400, and 700 rotations they were sieved and weighed, to assess the TI and AI indexes for the respective rotations. Since the equipment and the procedure are not standardized, a comparison with commercial ferroalloys was performed.

**Drop Test**

The drop test was done inspired by ASTM D440-07:2019. The apparatus used consisted of an iron vessel in the bottom part to collect the material after the drop and a tube attached to it, with 1.63 m high, to serve as a guide as the briquette falls inside the vessel. The briquette was dropped ten times, and at least two briquettes should be tested to provide satisfactory results. The material collected after the drops was sieved, and the fraction inside each sieve was weighted and calculated in a percentual basis according to the initial briquette mass. The average opening sizes of the sieves were 6.7 mm, 5.6 mm, 4 mm, 2 mm, 1 mm, 0.5 mm, and 0.125 mm and they were normalized according to the biggest (being 6.7 mm equal to 1, 5.6/6.7 mm equal to 0.836, and so on) for the size stability calculation. The weight percentage of the material collected after the drops are multiplied by the normalizing factor of each sieve average aperture it is collected, generating the size stability calculation. The weight percentage of the material collected after the drops are multiplied by the normalizing factor of each sieve average aperture it is collected, generating the size stability value ‘s,’ as the briquettes in the present study have the same diameters (single size).

Also, another parameter used was the impact resistance index calculation, used in some studies [40, 41], which consists into dividing the number of drops of each briquette level by the number of pieces it has split apart. The IRI calculation is as shown in Eq. 3:

$$\text{IRI} = 100 \times \frac{\text{Average number of drops}}{\text{Average number of pieces}}$$  \hspace{1cm} (3)

Another value calculated was the adjusted impact resistance index, which also considers the powder detached during the drop test. For each briquette dropped, the mass of powder retained in each sieve (Mi) below 4 mm was multiplied by a factor depending on which sieve aperture (Si) it passed, and then everything is summed up to generate an Adjusting Mass factor (Eq. 4). Then the adjusting factor (Eq. 5) is calculated by subtracting 1 from the ratio between the adjusted mass factor and final mass collected at the end of the test. If the adjusting factor is negative, it is considered zero. Then the adjusted impact resistance index is calculated by multiplying the IRI by the adjusting factor, according to Eq. 6. This adjusted index highlights how coarse the powder lost during the drop test is, meaning that an AIRI much lower than IRI represents big losses by fine powder detachment, and the finer the powder detached, the lower the AIRI would be.

$$\text{Adjusting mass factor} = \sum \frac{4}{\text{Si}} \times \text{Mi}$$  \hspace{1cm} (4)

$$\text{Adjusting factor} = 1 - \frac{\text{adjusting mass factor}}{\text{final mass}}$$  \hspace{1cm} (5)

$$\text{AIRI} = 100 \times \frac{\text{average number of drops}}{\text{average number of pieces}} \times \text{adjusting factor}$$  \hspace{1cm} (6)

**Decrepitation Test and Metallurgical Performances**

The decrepitation index was evaluated according to BS ISO 8371:2015 standard, by thermal treating the dried briquettes at 700 °C for 30 min in air. The briquettes were put in a hot oven preventively stabilized at the test temperature for 1 h. After the test, the briquettes were left to cool down and soon sieved at 6.7 – 4 – 2 – 0.5 mm sieves (sieve series according to ISO 3310). An electromagnetic sieve shaker IRIS FTS-0200 (Filtro Vibracion, Barcelona, Spain) was used at minimum power. Decrepitation index is defined as

$$\text{DI}_{6.7} = 100 \times \frac{m_2}{m_1}$$  \hspace{1cm} (7)

where $m_1$ is the mass of the test portion after thermal treatment, $m_2$ is the mass of the undersize fraction passing the 6.70 mm sieve.

Metallurgical performances were evaluated through the definition of low-temperature reduction-disintegration index, free-swelling index, and reducibility inspired by BS ISO 4696–2:2015, BS ISO 4698:2007, and BS ISO 4695:2015, respectively. Self-reducing capability was also evaluated by heat treating the briquettes at 1000 and 1200 °C under fluxed argon atmosphere for 15 min. Mass loss and mineralogical composition were evaluated after the treated briquettes were cooled down.

**Results and Discussion**

**Briquetting Procedure Evaluation**

**Briquettes Without Starch (JBFS)**

The briquetting machine upgraded on the Instron showed a good replicability and versatility regarding the production of JBFS and SJBS. All the briquettes produced had a diameter of 20 mm and showed satisfactory mechanical integrity at glance.
However, all the A coded briquettes (JBFS) showed difficulties during its production and handling. Careful handling of them was needed, as they seemed very fragile. Since they have no binder, the agglomeration forces present are not enough to allow a reliable production, given that two main effects were observed: first, the A+ briquettes cracked (Fig. 4) as they left the machine, showing a low success rate in the briquetting, nearly 50%. This phenomenon happens due to the high pressures involving the process, which is responsible for high pressure gradients in the briquette external diameter [42] that are not withstand by the agglomeration forces of the briquette.

The other undesirable effect seen was the swelling and the collapse of the briquettes (both A+ and A−) (Fig. 5) after the exposure to the ambient air for a couple of days. The possible swelling can be due to the lime hydration by air, to form calcium hydroxide, or/and by the thenardite–mirabilite transformation happening inside the briquettes according to the change of relative air humidity.

As most of the A briquettes collapsed, their production seemed not viable, so a binder is needed for the correct processing and handling of such briquettes. One A+ briquette, which did not suffer from the swelling damaging effects, was sent to the compression test and resulted with not enough resistance.

The briquetting pressures evolution against displacement of A+ and A− briquettes (Fig. 6) have almost the same behavior up to a value of 20 MPa, with no mechanical forces applied to the powder in the upper cylinder up to approximately 12 mm of piston displacement. Then, the powder starts to get pressed, forcing the machine spring downwards, resulting in an almost constant slight pressure increase from 12 to 48 mm of the piston displacement (segment A). When the spring is fully deformed (B point), the pressure starts to sharply increase (segment C). The A− briquettes reach the 20 MPa, and the dwell time starts, while the A+ briquettes have further pressing, where a smooth inflection in the pressure is seen (segment D). This may be due to the gas entrapment and release through the briquettes powder and to the plastic deformation or fragmentation of the particles [43] for a better accommodation of them.

**Briquettes with Starch (SJBFs)**

The B and C coded briquettes (Fig. 7), on the other hand, showed no difficulties in their production, as the binder addition seemed to give enough strength to the briquettes; therefore, no cracking issues in their retrieval from the machine and swelling during their drying period were seen.

As the moisture content in the binder was significant, the total volume of the mass inserted in the upper cylinder was higher than with A coded briquettes, and when the pressure during the production increased, water started to pour out from the machine gaps leading to some variation in the briquettes height with respect to the A coded ones. Figure 8 shows the briquetting pressures developed during the briquettes production. Like the unbound briquettes, the
process has the same behavior through the levels up to the point where the spring reaches its maximum displacement, which in the case of the bound briquettes, are close to a piston displacement of 50 mm. The slight increase of the piston displacement up to the spring max deformation is related to the water present in the binder, released during low briquetting pressures throughout the spring compression, demanding a higher displacement of the piston to reach the point of sharp increase of pressures. The briquettes done at 20 MPa (B− and C−) showed the same behavior during production of the unbound briquettes, apart from the slightly higher displacement (segment A and B). On the other hand, the briquettes produced at 40 MPa (B+ and C+) showed similar behavior between them, with much more expressive inflection at pressures higher than 20 MPa (segment C) compared with the A+ briquettes, probably due to a further water release and a possible rearrangement of starch and the JBFS powder at higher pressures [43–46].

Some of the SJBFS briquettes showed the development of a white structure in their walls (in the right upper corner of Fig. 7). This material was collected and analyzed by XRD and SEM and it was identified as thenardite, a crystalline compound composed mostly of Na₂SO₄. The sodium sulfate presence can be explained by the non-homogeneous or insufficient heating of the natrojarosite portion of the jarosite during the roasting process, as it is suffers thermal

Fig. 6 Briquetting curves of A coded unbound briquettes. A Powder compression alongside the spring deformation; B Spring maximum deformation (~47 mm of piston displacement); C compaction with the spring fully compressed; D inflection due to the entrapped gas release and particles rearrangement at 40 MPa

Fig. 7 Briquettes with binder (SJBFS). From the upper to the lower line: B+ B− C− C+ briquettes
decomposition at nearly 900 °C [14, 47] leaving a small portion of the raw JBFS powder with a residual Na₂SO₄ which did not decompose. As the sodium sulfate gets in contact with the water present in the gelatinized starch during the SJBFS production, it dissolves. As the water starts evaporating from the briquettes during drying, the sodium sulfate crystallization into thenardite (anhydrous sodium sulfate) starts to happen [48–50]. At temperatures lower than 32.4 °C, the precipitation of mirabilite (Na₂SO₄·10H₂O) is expected. However, as described by Rodriguez-Navarro et al., the mirabilite decomposes into thenardite when the air relative humidity drops below 71%, or the solution can even precipitate into thenardite directly at a relative air humidity below 40% and temperatures of 20 °C [49]. The growth of both mirabilite and thenardite crystals inside porous structures can be critical due to the high volume increase and pressures involved during it [48–52]. This could be the main factor for which the briquettes that developed such structures during their drying had their mechanical properties hindered, and it will be discussed later.

After processing, the briquettes were left to dry at room temperature. After 72 h no significant mass changes from water evaporation were observed as for their dimensions. A regression, shown in Fig. 9a, shows the drying behavior of the briquettes, alongside an equation of the fitted curve, confirming that after 3 days of drying, negligible evaporation losses are seen, while after one day, the mass loss ranges from 8 to 12%, and after two days, from 5 to 1%. The analysis of variance (Fig. 9b and Table 4) highlighted that both the binder amount and briquetting pressure are statistically significative on the mass loss. Note that A coded briquettes are not present in this analysis, due to the absence of the binder. The results show that a higher pressure leads to lower mass loss related to the evaporation, this is due to the removal of the water contained in the starch gel being forced out through the machine gaps by the high machine load, leading to a lower moisture produced briquette. Also, being the starch gelatinized at a ratio 1:6 with water, a briquette with a higher content of binder implies a higher amount of water that leads to a higher loss by evaporation after briquetting.

The density of the briquettes was then assessed just after their production (wet briquettes), and after their drying period. It was found that the wet briquettes showed no significant difference in their density. However, they showed differences regarding their dry density, as shown in Fig. 9c. As expected, higher pressures led to more dense briquettes, which is commonly seen in processes involving briquetting, because there is a better agglomeration of the material and less empty space inside the briquette [53].
results show (Table 4), the binder also played a role in the density analysis, and this is due to the starch powder being less dense than the JBFS powder, so the higher the amount of binder added, the less dense the briquette will be, and also due to the non-filled spaces left inside the briquettes due to the water evaporation, as the briquettes with binder addition show a trend similar to the mass loss analysis: the higher the mass loss due to the evaporation, the lower the density, as there was no dimensional variation of the briquettes.

As the briquettes dried, an XRD analysis was conducted on three different briquettes, a good B+ and C+, and a B+ briquette developing thenardite (marked as B+ *). Comparing with the initial composition of the JBFS powder (Fig. 10a), all the three briquettes show the formation of gypsum, a hydrated form of calcium sulfate ($\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$), formed due to the water presence from the binder addition, which dissolves in water as anhydrite and precipitates as gypsum when the briquette dries [54, 55]. Also, a similar phenomenon happens to the iron oxide ($\text{Fe}_2\text{O}_3$), which could have transformed into goethite ($\text{FeO(OH)}$) by the presence of water [56, 57]. For what concerns the B+ * briquette, both thenardite and nathrojarosite were found, which reinforce the assumption that the nathrojarosite presence in the powder is an indicative factor of improper heating during the raw jarosite roasting, resulting in the thenardite crystallization in the briquettes due to the presence of $\text{Na}_2\text{SO}_4$ that did not decompose to $\text{Na}_2\text{O}$ during roasting due to insufficient heating (Fig. 10b).

**Physical Characterization**

**Cold Compression Strength**

Figure 11a and Table 5 show the curves obtained from the cold compression strength (CCS) test. Besides from the B− specimens, which have apparently lower mechanical properties, all the samples seem to be very similar regarding their compression behavior. In Fig. 11b, the comparison among the only JBFS briquette not swelled (A+), B+ briquettes and a briquette with thenardite (B+ *) is shown. From this plot, it is clear that the latter has a noticeable decrease in the compression resistance in comparison with the B+ briquettes, confirming that the thenardite crystallization hinders the compression resistance. On the other hand, the B+ * briquette CCS is higher than the A+ briquette one, showing that it is preferable to be prone to thenardite crystallization rather than not mixing the gelatinized starch to the JBFS. The starch addition and the increase of mechanical properties can be related to the interaction between the gelatinized starch and the JBFS powder, which leads to a better adhesion between the particles by capillary and molecular forces. Furthermore, the gelatinized starch, containing amyllose and amylopectin in amorphous structure could enhance the cohesion between the particles [44, 58–61], leading to a higher resistance during the test.

The binder addition has no statistical influence on the CCS of the briquettes, however, the briquetting pressure can be considered as significant factor, since it has a $P$ value lower than 0.1, leading to a statistical significance for 90% interval of confidence (Table 6). This can be explained by the fact that higher briquetting pressures leads to higher contact between the powder particles (higher plasticization

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**Table 4** ANOVA table regarding the mass loss and dry density analysis

| Source         | DF | Adj SS   | Adj MS  | $F$ value | $P$ value |
|----------------|----|----------|---------|-----------|-----------|
| **Analysis of variance: mass loss** |    |          |         |           |           |
| Binder         | 1  | 0.000876 | 0.000876| 7.07      | 0.013     |
| Pressure       | 1  | 0.004021 | 0.004021| 32.46     | 0.000     |
| Binder*Pressure| 1  | 0.000017 | 0.000017| 0.13      | 0.717     |
| Error          | 27 | 0.003344 | 0.000124|           |           |
| Total          | 30 | 0.008118 |          |           |           |
| **Analysis of variance: dry density** |    |          |         |           |           |
| Binder         | 2  | 0.29932  | 0.149659| 16.65     | 0.000     |
| Pressure       | 1  | 0.18473  | 0.184733| 20.56     | 0.000     |
| Binder*Pressure| 2  | 0.02751  | 0.013755| 1.53      | 0.230     |
| Error          | 36 | 0.32351  | 0.008986|           |           |
| Total          | 41 | 0.81707  |          |           |           |
and fragmentation of particles), which increases the van der Waals and interlocking forces in the briquette [43, 62, 63] as seen in the dry density analysis (paragraph 3.1.2). Comparing with other studies using similar materials [28, 32, 42, 64–66], the briquettes showed good compression strength, having a mean value above 12 MPa for the 40 MPa briquetted specimen, with a maximum value of 16 MPa for C+2 sample, and an axial resistance of more than 250 kgf. For instance, Kumar et al. [32] tested briquettes of mill scale and dust with starch as binder, and obtained briquettes resisting 140 kgf, while Narita et al. [42] obtained iron ore with coal briquettes that achieved compression strengths not

Fig. 10 XRD pattern of the JBFS mix before briquetting (a) and SJBFS briquettes after drying (b)
higher than 5 MPa before heat treatment, evidencing the strength of the briquettes in the present study. One key factor that might also influence positively the strength of the briquettes compared to the mentioned studies is the particle size of the SJBFS used being not higher than 0.125 μm, which increases contact area between particles during briquetting and the van der Waals forces consequently.

As shown in Table 5, the deformation of each briquette at the maximum compressive strength was also calculated, which have the same trend of the CCS. Even in this case the binder has no statistical influence on the deformation, whereas the pressure has a  value lower than 0.1, highlighting that it may be influential, probably due to the higher adhesion and van der Waals forces between the particles that allow a higher energy absorption as deformation (Table 6).

Finally, by calculating the area of the compression curve, the energy absorption of the briquette during the compression could be calculated up to the CCS point (Table 5). Also, in this case, only the pressure have a  value under
0.1, meaning that starch as a binder may not promote higher mechanical properties, whereas briquetting pressure is still significant for the same reasons said previously (Table 6).

Figure 12 shows the difference between a B− and a B+ briquette after the compression test. The former appears much more damaged after the test, whereas the latter shows how the pressure influences the agglomeration of the powder.

**Tumbler and Abrasion Test**

The tumbler and abrasion test are performed to simulate the briquettes resistance through processes like conveyor belts charging and transporting. Two main destructive phenomena are observed: the drop damage, as the briquettes are constantly falling from the top to the bottom of the tumbler and the abrasive phenomena, as the briquettes slip and/or rotate in the wall of the tumbler, interacting or not with each other.

The results presented in Table 7 show that the briquettes tumbler index and abrasion index are similar for 200 and 400 rotations. However, in the case of the B+ tested briquettes, they have suffered a big loss after 700 rotations, due to the disintegration of one of the tested briquettes (Fig. 13a). In particular, this latter B+ briquette has shown a white color in its side wall, which can be related to an uneven distribution of the binder inside the matrix due to an improper starch gelatinization that hindered the binding properties. For this reason, two more briquettes, labeled as (B+ * ) were tested again, with results comparable with the other specimens. All the other briquettes showed good results during the tests, with good abrasion and tumbler index, especially the B− briquettes (Table 7). Both abrasion and tumbler indexes may have a linear relationship with the number of rotations up to 700 rotations, except in the case where there is a defective briquette. Moreover, the C+ and C− briquettes showed similar trends in both tests, giving a hint that maybe the pressure does not influence the abrasion resistance, but since the experiments are not replicated, an ANOVA analysis was not possible. The B+ * showed the lowest performance during the test, possibly due to thenardite crystallization that lowers its mechanical properties. However, the binding forces provided by the properly gelatinized starch during its production were enough to keep the briquettes integrity (Fig. 13b), making it still suitable for use.

The tumbler index after 200 rotations was higher than 90% for all the briquettes, showing their good resistance to abrasion if compared to briquettes with similar composition (iron oxide and/or carbon sources) from other studies [61, 67, 68]. Rocha Lemos et al. [61] characterized pillow and cylindrical briquettes made of blast furnace sludge and dust with different binders, the results showed that by increasing the sludges fraction, the abrasion resistance of the briquettes increases, as most of the briquettes with a higher fraction of sludges were retained by more than 90% of their

| Sample | B− | B+ | B+ * | C− | C+ |
|--------|----|----|------|----|----|
| Rotations | Abrasion index | Tumbler index | Fine fraction (<500 μm) |
| 200 | 1.5% | 2.7% | 3.9% | 2.9% | 2.3% |
| 400 | 2.5% | 6.3% | 5.0% | 4.2% | 4.0% |
| 700 | 3.7% | 26.4% | 6.4% | 5.7% | 5.9% |
| 200 | 98.0% | 96.9% | 91.4% | 92.3% | 95.9% |
| 400 | 96.7% | 93.0% | 89.7% | 90.0% | 93.0% |
| 700 | 95.4% | 57.4% | 88.1% | 87.9% | 90.1% |
| 200 | 1.44% | 2.14% | 2.90% | 2.00% | 2.01% |
| 400 | 2.40% | 5.20% | 4.17% | 3.45% | 3.43% |
| 700 | 3.52% | 23.86% | 5.69% | 5.10% | 4.30% |

**Fig. 12**  B− briquette (a) and B+ briquette (b), after the test.
cumulative mass by a sieve with 7.5 mm aperture, highlighting the sludges agglomeration and resistance features. In the case of the B− and C+ briquettes, the tumbler index remained higher than 90% after 700 rotations, showing that the gelatinized starch is a suitable binder for achieving a suitable abrasion resistance. However, being the equipment and the conditions not standardized, the obtained results can be only qualitatively compared with the literature ones. For this reason, a control test was performed by tumbling some ferroalloy stones (FA). The tumbler index of FA passed from 99.56 after 200 runs to 99.05 after 700 runs, while the abrasion index varied from 0.37 to 0.94, therefore demonstrating high resistance to abrasion and crushing as can be expected by a stony material. To be an agglomerate, the produced and tested briquettes behave similarly to FA, especially the B− samples, once again demonstrating the high cold resistance reached by the agglomeration process.

Moreover, all the briquettes did not have a cumulative mass of fines largely higher than 5% passing through the 0.5 mm sieve. After 700 rotations, both B− and C+ briquettes showed cumulative masses lower than 5%. Therefore, they can be considered the best combinations regarding possible fine powder losses. The B+ level had a low performance after 700 rotations, like the abrasion index, due to the disintegration of one of the briquettes, demonstrating that a non-properly bound briquette may be problematic not only mechanically, but also for the environment, as these powders could be dispersed during transportation.

**Drop Test**

The drop test is a good qualitative analysis to provide a parameter for the loading and unloading of briquettes inside a cargo vehicle, or the briquettes loading inside a furnace. The drop tests were performed with two of each C level briquettes, and three B level briquettes. Also, four briquettes, two from the B+ level and two from the B− level showed thenardite precipitation in their side, so they were tagged as B+ *, B+ **, B− *, and B− **.

Table 8 shows the drop test results for each briquette. The size stability factor is calculated according to the ASTM D440-07:2019, which provides a lower weigh for the fine powder collected in the lower sieve apertures, so the effects of powder detachment and briquette cracking are not very significant in this analysis, as the mass dropped in each test is very low compared to the defined by the standard. The size stability average value for each level is higher in the case of higher compressing pressures during briquetting. Moreover, this test does not take into account how many drops it takes to break apart the briquette. Except for the thenardite containing briquettes, which shattered apart in less than five drops, the remaining specimens resisted at least ten drops prior to breaking into smaller pieces. These effects are better visualized by the impact resistance index (IRI) and adjusted impact resistance index (AIRI), which highlights the lower impact resistance from the briquettes with thenardite crystallization.

The IRI and number of drops for the briquettes with no thenardite crystallization were very good. When a comparison is done with studies with similar materials and testing methods [28, 40, 60, 65, 69], the IRI values obtained are considered more than satisfactory for the briquettes handling and transportation, highlighting the binding power of the starch together with the JBFS powder. Indeed, the IRI threshold for a good briquette is 50 [40, 41] while the current briquettes have IRI values from 10 to 20 times higher. Han et al. [60] produced briquettes of blast furnace dust with several binders, and the ones with corn starch were the ones that resisted the most at ambient temperature, with the average number of falls before breakage being 6 from a 0.5 m
high drop, Wu et al. [69] produced briquettes of EAF dust and anthracite with bentonite as binder, the biggest value of drop number was 8.2, with drops of 0.5 m high also, whereas in the present study, the B briquettes with no thenardite survived 10 drops at 1.63 m high drops, demonstrating high resilience.

The briquettes with thenardite, however, show low IRI values like happened during the abrasion analysis. However, two different behaviors were observed. B−* and B+ * samples show IRI values below the threshold, confirming that alongside the lowering of the mechanical properties due to the thenardite, the starch binding of the JBFS matrix did not happen properly as hypothesized in the Sect. Tumbler and Abrasion Test. Also, the size stability was compromised, being the powder detachment significantly high. On the contrary, B−** and B+** samples show an IRI value above the threshold and also an improved size stability, showing that the gelatinized starch generated good agglomeration of the powder, and the mechanical properties regarding the drop resistance of the briquette were mostly compromised due to possible cracks inside the briquettes matrix promoted by the thenardite crystallization.

Thus, the reason for such poor mechanical property hindering is the same as explained in the paragraph 3.1.2, as the thenardite crystallization generates high internal pressures inside the briquette, which when not properly bound, is not minimally capable of sustaining mechanical loads and impacts. Figure 14 shows the comparison of the briquette matrix of the B+ (Fig. 14a) and the B+ * (Fig. 14b) briquettes after the test, and it is possible to observe cracks inside the B+ * matrix, with small white structures inside it, while the B+ briquette seems more homogeneous and without cracks.

Decrepitation Test and Metallurgical Performances

The decrepitation test is used to assess the briquettes resistance due to sharp temperature increases, it provides better...
understanding of how the briquettes would behave during charging in a hot furnace. The decrepitation index (DI₆.₇), the mass retained at the sieves, and the loss of ignition are shown in Table 9. It appears that the C coded briquettes have very poor performance regarding the decrepitation, as their indexes are very high. However, this low performance compared to the B coded briquettes can be attributed to the fact that the C coded briquettes were characterized by some defects and cracks, which led to their collapse at higher temperatures, explaining this discrepancy, which is not present in the previous tests. The briquetting pressure, as in the compression test, seems to have an important role to keep the briquettes integrity in high temperatures by comparing only through the same binder level.

The loss of ignition was higher for the C coded briquettes, due to higher amount of binder and water mixed during the briquette production compared to the B coded ones. At high temperature, the starch decomposed as the water captured by its polymeric chains evaporated, explaining the higher losses during the test. This water evaporation can lead to higher internal pressures compared to the B coded briquettes, explaining the lower performance compared to the B briquettes. Figure 15 shows the briquettes after the test, it can be noticed that the B+ briquette integrity after the test is remarkably higher compared to the others, as its appearance, which has not become darker, unlike the other briquettes, especially the C− one. Comparing the results with the ones of other studies using similar materials and test methodology [67, 70], the B+ briquette had outstanding performance, whereas the other briquettes had similar results compared with the literature.

After the test, the B+, B−, and C− specimens were submitted to an XRD analysis so as to identify the species formed inside it, the obtained spectra are shown in Fig. 16. From the analysis, it is possible to note that there are no hematite nor goethite, as they were reduced into magnetite by the carbon present in the blast furnace sludges. Furthermore, the presence of other compounds such as

| Specimen | DI₆.₇ (%) | DI₄ (%) | DI₂ (%) | DI₀.₅ (%) | L.O.I (%) |
|----------|----------|---------|---------|-----------|-----------|
| B−       | 9.26%    | 8.29%   | 7.50%   | 6.15%     | 16.3%     |
| B+       | 0.94%    | 0.94%   | 0.94%   | 0.30%     | 16.4%     |
| C−       | 39.88%   | 37.85%  | 30.73%  | 12.79%    | 20.3%     |
| C+       | 33.03%   | 29.88%  | 22.66%  | 11.45%    | 21.0%     |

Fig. 15 Briquettes after the decrepitation test: (a) B+; (b) C+; (c) B−; (d) C−
brownmillerite (Ca₂(Al,Fe)₂O₅) and langbeinite (K₂(Ca,Mg)(SO₄)₃), is explained by the reaction among calcium sulfate, magnesium oxide and potassium oxide, coming from the jarosite roasting [14]. There is also the presence of galena (PbS) which is by the reduction of PbSO₄ by carbon monoxide, according to Eq. 8 [71, 72]:

\[ \text{PbSO}_4 + 4\text{CO} \rightarrow \text{PbS} + 4\text{CO}_2 \]  

Finally, the traces of pyrite (FeS) are probably formed following the reaction of hematite and concentrated sulfur species [14, 73].

By exposing the self-reducing briquettes at high temperature in inert atmosphere, iron formation was detected through XRD analysis (Fig. 17), demonstrating the self-reducing capability of the produced material. Some differences can be highlighted from the samples treated at 1000 and 1200 °C. At 1000 °C iron formation is accompanied by a partial reduction of the original hematite that leads to the formation of wustite. Sulfur-rich compounds are detectable like sphalerite and wurtzite, probably due to the solid-state reaction between gypsum, reduced franklinite, hematite, and goethite. Zn is also present in hardystonite, probably due to the reaction among gypsum, franklinite, and quartz. Metallic lead is also formed thanks to the final reduction of galena operated by CO, passing through the formation of PbO (Eqs. 9 and 10). A slag composed by gehlenite and albite characterized the remaining material. These two compounds were already observed during the roasting of loose powders mix by the same authors [23]. An overall mass loss of 39 wt% was registered, in good agreement with the TG-DSC analysis conducted on loose powder mix by the same authors [23].

\[ 2\text{PbS} + 3\text{O}_2 \rightarrow 2\text{PbO} + 2\text{SO}_2 \]  

(9)

\[ \text{PbO} + \text{CO} \rightarrow \text{Pb} + \text{CO}_2 \]  

(10)

At 1200 °C, the number of compounds still present in the residual briquettes diminished significantly. The residue is formed by iron, magnetite, gehlenite, and oldhamite. By the higher thermal level, all the sulfides compounds were reduced and passed into the iron fraction as sulfides, as detected in the previous research [14, 23]. Lead and zinc were completely volatilized, as a typical white crystal oxide was collected in the exhaust gas system of the oven. Moreover, an overall mass loss of 53 wt% was registered. This higher mass loss than the mass loss measured at 1000 °C can be attributed to the vaporization of volatile metals (Zn, Pb) and to a higher degree of metallization. The residual fraction of magnetite is due to a partial reoxidation caused by a non-perfect fluxing of the inert atmosphere and a thermal level higher than at 1000 °C.

The obtained results are encouraging since they demonstrate that by charging these briquettes in a reducing atmosphere, like the one be present in a Midrex shaft or in a cupola furnace, the reduction of iron oxide will be complete.

Metallurgical performances of B+ briquettes were evaluated through the definition of low-temperature disintegration index, swelling index, and reducibility. The low-temperature disintegration index was evaluated by tumbling some briquettes after exposure at 760 °C for 30 min in a controlled...
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atmosphere (CO/N₂ = 30/70%). The average RDI-2-2.8 index was 39%. This value is lower than what was obtained by reducing iron ore sinter in the controlled CO-H₂-N₂-CO₂-H₂O atmosphere [74] but higher than the threshold to ensure the smooth operation of blast furnaces charged with briquettes (< 30%). For values of this index greater than 35–40%, the blast furnaces’ performance was affected [67].

Swelling index, determined after an exposure at 900 °C for 60 min in a controlled atmosphere (CO/N₂ = 30/70%) ranged from -10% to -12%, meaning a contraction of volume due to the reduction of iron oxide to metallic iron, as is shown in Fig. 18a. Similar results were obtained by Singh and Bjorkman for cylindrical briquettes with low-sized particles (d₅₀ < 74 μm) [75], even if most of the literature records report an increase in volume after exposing briquettes to reducing environment [76, 77]. This is probably due to the low alkalinity of the mix (~0.5) that favors the shrinkage of the volume during the reduction from wustite to iron [78].

In addition, the briquettes retained almost the original shape, with an overall contraction both in height and diameter, even if high fraction of cracks are formed on the surface as also shown by Bagatini et al. [77]. The obtained results comply with the threshold of 20%, which is considered acceptable according to the literature [70] for blast furnace operation.

Finally, the reducibility was evaluated by recording the mass loss over time by keeping the briquettes under a controlled atmosphere (CO/N₂ = 40/60%) at 950 °C for 4 h (Fig. 18b). An overall mass loss of over 37% was achieved, in total agreement to TG–DSC analysis conducted on loose mix and as stated in the previous paragraph. The final degree of reduction was higher than 80%, a result similar to the briquettes test by Kemppainen et al. [76].

Although the different tests (swelling, disintegration, reducibility) were performed in different conditions, for some of the briquettes, swelling index (Vₛ) was still calculated. In Fig. 19, the evolution of Vₛ as a function of temperature and degree of reduction (RD) was reported. Swelling formed similar to what was observed by Singh and Bjorkman [75] and Iljana et al. for acidic pellets [78], with a high-volume expansion at low temperature (760 °C) during the reduction of magnetite to wustite and then a shrinkage in the wustite to iron reduction step. This is confirmed by the trend of swelling as a function of the degree of reduction. At 760 °C, the RD is slightly higher than 40%, in total

Fig. 17 XRD of B+ briquettes heat treated at (a) 1000 °C and (b) 1200 °C under fluxed argon atmosphere
agreement with the experiments conducted by Iljana et al. [78], thus confirming that at that temperature, magnetite to wustite reduction occurs. Increasing the temperature, the RD increases proportionally while the briquettes start to shrink due to the formation of metallic iron and volatilization of Zn and Pb compounds.

Fig. 18  a comparison of briquette before and after reduction at 900 °C for 60 min under controlled atmosphere (CO/N₂ = 30/70%); b mass loss Vs time for reduced briquettes at 950 °C under controlled atmosphere (CO/N₂ = 40/60%) for 4 h
Discussion

The aim of this work being the production and characterization of self-reducing briquettes to be used in a thermal reactor, such as a Midrex or a cupola furnace, a comparison with the characteristics of the input materials of such reactors was made.

Midrex process is the world leader in direct reduction technology, and it is based on a shaft furnace feed by iron ores that are reduced by a reducing atmosphere in counter-current made by a mixture of CO and H₂. Maximum operating temperature is less than 1000 °C. Typical input feedstock of the shaft are pellets or lumps that should have at least 85% of tumbler resistance, a fine fraction below 6 wt% and a minimum compressive strength of 150 kg (roughly 15 MPa if considering the lateral surface) [79]. Recently, extruded briquettes (brex) made of pellets fine, metallized sludge from DRI production, mill scale, and EAF dust were tested in a full-scale Midrex reactor. Even if such brex briquettes have a maximum compressive strength below 11 MPa, they behave well in the shaft [26]. Thus, from a mechanical point of view, the briquettes discussed in the current paper almost meet the requirement for acceptable Midrex input material, since they possess an average CCS of 12 MPa when pressed at 40 MPa and showed good tumbler resistance, always over 90% for well-briquetted material, and weak fines generation (less than 5%). However, from a chemical point of view, the excess of S, Zn, and Pb, do not encourage their use, mainly because the thermal level in the shaft will not be enough to reduce and completely volatilize Zn and Pb, as observed in the heat-treated briquettes.

On the other hand, one of the advantages of cupolas is that they can easily process high amounts of zinc in the burden which may either originate from steel plant residuals or from automotive scrap. In addition, most of the iron is reduced at a wide temperature range (900–1400 °C) over a short period of time, generally 20–30 min [25]. Being not reference values for input material in cupolas, a comparison was made with carbon-bearing briquettes physical properties. Typical anthracite- or biocarbon-based briquettes for coke replacement in cupola furnace possess a CCS in the range 2–14 MPa, high enough to keep their structure at a high temperature [24, 80, 81]. In addition, self-reducing briquettes made by several steelmaking waste were judged suitable for Cupola reduction if they possess a CCS > 5 MPa [25].

Thus, the specific threshold for Midrex® neither cupolas being unavailable, a direct comparison of the obtained results for swelling and disintegration is not possible. However, the high-temperature properties of such briquettes seem to be suitable for their operation in blast furnace. Thus, it is reasonable to assume that, at least from the mechanical point of view, the briquettes are suitable both for Midrex® and cupola. For instance, the B+ briquettes conserve enough strength to avoid high fine formation at high temperature, mainly due to the absence of swelling promoted by the low alkalinity of the mix.

On the other hand, from a chemical point of view, the higher temperatures developed in the cupolas can remove the harmful elements (Zn, Pb) opening a feasible possibility to use jarosite–blast furnace briquettes as input material in such a furnace. In addition, even the sulfur content of the produced cast iron might be high (5–6 wt%) [23], a desulfurization process can be provided to refine the iron composition, both in the cupola or in a ladle [82].

Finally, for instance, the lower permissible compressive strength required to withstand the pressure of the overlying layers of 30 m blast furnace is 5.8–6.0 MPa. Being a cupola furnace generally the half in height of a blast furnace, it is auspicious that the proposed briquettes can withstand the pressure of burden inside the cupola. Hence, the proposed
briquettes possess sufficient mechanical strength to be applied in industrial scale.

Conclusions

The briquetting of the jarosite sludges together with blast furnace sludges was successful with binder addition. Briquettes with no starch addition as a binder showed little absence of mechanical resistance, therefore their production is not preferable. A desirable control of the roasting process of the jarosite powder, previously to the briquetting, is needed so as to promote the decomposition of sodium sulfate into sodium oxide, to avoid the thenardite crystallization during the briquettes drying, which may dramatically hinder the briquettes’ resistance.

A good control of the starch gelatinization is needed to provide the desired agglomerating forces.

The ultimate compressive strengths observed from the B+ and C+ briquettes, with average values higher than 12 MPa, tumbler index higher than 90% after 200 rotations, and an IRI higher than 500, provide sufficient high mechanical properties for the industrial application of the briquettes.

With respect only to the mechanical properties, the addition of 5 wt% of gelatinized starch seems to be the most reasonable choice; furthermore, the mechanical properties obtained using a briquetting pressure of 40 MPa seem to be the best ones due to the fact that no significant influence has been observed from the binder amount to the final properties of the briquettes.

Finally, the physical, mechanical, and metallurgical properties developed during agglomeration allow the use of such briquettes as feedstock for cupola furnace.

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Declarations

Conflict of interest On behalf of all authors, the corresponding author states that there is no conflict of interest.

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