Production of metallurgical-grade silicon from Egyptian quartz


Synopsis
Metallurgical-grade silicon (MG-Si) was successfully produced by carbothermic reduction-smelting of Egyptian quartz using a mixture of carbonaceous materials in a locally made semi-pilot submerged arc furnace. The effects of different technical parameters such as the carbon to silica ratio, smelting time, power input, and weight ratio of carbon sources were investigated. The MG-Si product was characterized by X-ray diffraction (XRD), X-ray fluorescence (XRF), and scanning electron microscopy (SEM). A maximum silicon recovery of about 75% at a product purity of about 97% was achieved under the optimum experimental conditions of 80 minutes’ smelting time, 48 kW power input, and a carbon/silica ratio of 0.39.

Keywords
metallurgical-grade silicon, carbothermic reduction, quartz, charcoal, coke, petroleum coke.

Introduction
Metallurgical-grade silicon (MG-Si) is an important metal which has a range of diverse industrial applications such as a deoxidizer in steelmaking industry, alloying element in the aluminum industry, the preparation of organosilanes, and the production of hyper-pure ‘electronic grade’ silicon (>99.99% Si), which is used in the electronics industry as well as solar cells (PV Education. 2013; Aasly, 2008).

MG-silicon is produced by the carbothermic reduction-smelting process, in which silica is reduced by carbon in a submerged arc furnace at temperatures between 1300 and 2000°C under atmospheric pressure. The charge materials include a silicon source (quartz, sand, or quartzite) and a typical reductant blend comprising coke, coal, charcoal, and wood chips. The reduction process occurs according to the following reaction:

\[
\text{SiO}_2 (s, l) + 2C(s) \rightarrow \text{Si}(s,l) + 2\text{CO} \quad \Delta H_{2000^\circ C} = \text{687 kJ/mol} \tag{1}
\]

This overall reaction is the sum of different reactions inside the furnace, which can be specified according to the temperature range. The furnace environment is divided into two zones: high temperature (approx. 2000°C) and low temperature (< 1811°C) zones, in which different reactions dominate. In the high-temperature zone around the electrode tip, the following reactions occur:

\[
2\text{SiO}_2 (s, l) + \text{SiC} (s) \rightarrow 3\text{SiO} (g) + \text{CO} \quad \Delta H_{2000^\circ C} = \text{1364 kJ/mol} \tag{2}
\]

\[
\text{SiO}_2 (s, l) + \text{Si} (l) \rightarrow 2\text{SiO} (g) \quad \Delta H_{2000^\circ C} = \text{599 kJ/mol} \tag{3}
\]

\[
\text{SiO} (g) + \text{SiC} (s) \rightarrow 2\text{Si} (l) + \text{CO} \quad \Delta H_{2000^\circ C} = \text{167 kJ/mol} \tag{4}
\]

The slowest of these three reactions are probably the SiO(g)-producing reactions [2] and [3], which consume a major part of the input electrical energy. Silicon can be produced through reaction [4] at temperatures above 1811°C. The SiO gas travels upwards in the furnace and is recovered either by reaction with the carbon material (Equation [5]) or by condensation where the temperature is sufficiently low (<1800°C) (reactions [2 rev.] and [3 rev.]):

\[
\text{SiO} (g) + 2\text{C} (s) \rightarrow \text{SiC} (s) + \text{CO} (g) \quad \Delta H_{2000^\circ C} = \text{78 kJ/mol} \tag{5}
\]

\[
3\text{SiO} (g) + \text{CO} (g) \rightarrow 2\text{SiO}_2 (s,l) + \text{SiC} (s) \quad [2 \text{ rev.}] \quad \Delta H_{2000^\circ C} = \text{1380 kJ/mol} \tag{6}
\]

\[
2\text{SiO} (g) \rightarrow \text{SiO}_2 (s,l) + \text{Si}(l) \quad [3 \text{ rev.}] \quad \Delta H_{2000^\circ C} = \text{606 kJ/mol}
\]

The condensate-producing reactions [2 rev.] and [3 rev.] are strongly exothermic and are the main factors that contribute to the upwards heat transport in the furnace.

The temperature at the top of the furnace can vary between 1000 and 1700°C. Typical industrial silicon yields are around 85% in a
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well-operated furnace. The SiO gas will only be captured by reactions [2 rev.] and [3 rev.]. The temperature has a major effect on the equilibrium conditions for these two reactions. Much work has been done in recent years to define the reaction zones inside the silicon furnace (Myrhaug, 2003; Tangstad et al., 2010; Ringdalen and Tangstad, 2012; Asen 2012; Tangstad, 2012; Schi, Tuset, and Tvei, 1998).

In Egypt, tremendous resources of high-quality sands and quartz (the average content of SiO₂ is 99%) exist in Sinai and in the Eastern Desert. Despite the huge silica resources and the availability of reductants, there is no silicon industry in Egypt. In this paper we investigate the possibility of utilizing the local quartz resource for the production of MG-Si by studying the factors affecting the production process, such as carbon/silica ratio, reduction time, power input, and weight ratio of carbon sources.

Experimental

Materials

One ton of quartz from the Al Faleg area, Marsa Alam, Eastern Desert was subjected to a series of crushing cycles using a pilot jaw crusher. The particle size between 16 mm and 2 mm was used for the production of MG-Si. XRF analysis showed that quartz consists mainly of SiO₂ (99.94%), with minor amounts of Fe₂O₃ (0.02 %), Al₂O₃ (0.02%), and MgO (0.02%). Trace amounts of other oxides such as CaO (0.002%), K₂O (0.003%), and TiO₂ (0.0002%) were detected. XRD analysis was conducted to determine the phase composition of the quartz. Figure 1 shows that all the peak positions match the standard quartz phase.

Different carbon sources such as charcoal, coal, petroleum coke, and wood chips (with a fixed carbon content of 45%) were used as reductants. Table I shows the approximate chemical analysis of the carbonaceous materials.

Reduction technique

Reduction-smelting tests were performed in a locally-made electric arc furnace (EAF). The furnace is a submerged single-electrode (125 kW) furnace powered by five transformers. Smelting tests were conducted in a graphite hearth of 35 cm diameter which acts as the earth conductor, with a graphite rod 10 cm in diameter and 120 cm in length, located in the center of the graphite hearth, acting as the live electrode. The graphite electrode is connected to the terminal of the power transformer through a pure copper tube cooled by a flowing current of water. The furnace is tilted on its horizontal axis to pour the molten silicon metal at the end of each experiment. The quartz sample was mixed thoroughly with predetermined amounts of carbonaceous materials according to the following relationship.

The reduction experiments were carried out by introducing about 10 kg of the mixture through the feeder into the furnace. After smelting, the extent of the reduction process was calculated according to the following formula:

\[
\text{Extent of reduction} \% = \left( \frac{\text{Si}_p / \text{Si}_0} \right) \times 100
\]

where \( \text{Si}_0 \) is the initial silicon content in quartz and \( \text{Si}_p \) is the produced Si metal.

Results and discussion

The data obtained and technical parameters affecting the carbothermic reduction smelting of quartz to produce MG-silicon metal are discussed in the following sections.

Charge composition

Effect of carbon/silica weight ratio

To ensure a high silicon recovery, it is necessary to maintain the carbon/silica weight ratio of the charge at the required stoichiometric value. A series of experiments was carried out using mixtures of the pure crushed quartz sample with different proportions of coal, petroleum coke, charcoal, and wood chips. Different ratios of fixed carbon to silica (C/SiO₂) ranging from 0.30 to 0.45 (i.e., from 1.5 to 2.25 molar ratio of fixed carbon to silicon dioxide) were used in this study. The charges were smelted in the EAF at constant conditions of current (about 0.8 kA), voltage (about 45 V), and power input (about 36 kW) for a constant smelting time of 60 minutes. The extent of reduction of silica as a function of the carbon/silica ratio is depicted in Figure 2. It is clear that the reduction extent increases from 15% to about 30% as the carbon/silica ratio increases from 0.3 to 0.39 (i.e. from 1.5 to 1.95 molar ratio of fixed carbon to silicon dioxide). The maximum recovery of MG-Si was about 50%. Increasing the C/SiO₂ ratio above 0.39 was accompanied by a decrease in reduction extent. It can be concluded that the optimum C/SiO₂ weight ratio is about 0.39 (1.95 molar ratio of C/SiO₂), which is in good agreement with the published results (Dosaj et al., 1992; Arvidson, Dosaj, and May, 1991).

XRD characterization of the product metal was conducted to ascertain the reasons for these trends. Figure 3 shows the presence of the silicon carbide phase behind silicon metal in the product from a charge containing a C/SiO₂ weight ratio of

<table>
<thead>
<tr>
<th>Table I</th>
<th>Chemical analysis (%) of the different types of carbon used as reductants</th>
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<tbody>
<tr>
<td>Carbon type</td>
<td>Coal</td>
</tr>
<tr>
<td>Moisture</td>
<td>0.5</td>
</tr>
<tr>
<td>Volatile matter</td>
<td>19.8</td>
</tr>
<tr>
<td>Ash</td>
<td>8.4</td>
</tr>
<tr>
<td>Fixed carbon</td>
<td>71.3</td>
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Figure 1—XRD spectrum of the quartz.
0.45, which indicates that the reaction is not complete. However, with the charge containing a C/SiO$_2$ weight ratio of 0.39 (1.95 molar ratio of C/SiO$_2$), only the silicon metal phase was detected. Smelting with an excess of carbon promoted the formation of an excess of silicon carbide, which lowered the amount of silicon produced. Subsequently, silicon carbide accumulated and filled the hearth of the furnace, which made it necessary to raise the electrode, which increased the temperature in the outer zone of the furnace. This will lead to an increase in the electrode consumption. Therefore, it is important for efficient furnace operation that the furnace is kept in carbon balance (Dosaj et al., 1992).

**Effect of varying charcoal additions**

Charcoal was found to be the best reductant due to its high reactivity, low ash content, and high electrical resistivity, which has a positive influence on the furnace electrical efficiency and process chemistry. A series of experiments was carried out using charges of quartz mixed with constant weights of coal, petroleum coke, and wood chips with different weights of charcoal. The carbon/silica ratio was maintained at 0.39, and the applied electrical power was about 36 kW with a smelting time of 60 minutes. Figure 4 shows the variation of the extent of silica reduction as a function of the ratio of charcoal to silica in the charge. The extent of reduction increased with increasing charcoal content in the charge up to 30% by mass, and then decreased. The initial increase in the extent of reduction is due to the high reactivity of charcoal, which enhances the reduction process and leads to the formation of excess silicon carbide. However, further charcoal additions accelerate the formation of silicon monoxide gas in the outer zone of the furnace, which escapes from the surface of reaction and leads to a decrease in the extent of reduction and lower recovery of silicon metal (Vaish, 2012; Kuhlmann, 1965)

A series of experiments was performed in which the ratio of petroleum coke to quartz was varied while keeping the carbon/silica ratio constant at 0.39. The applied electrical power was about 36 kW and the smelting time 60 minutes. Figure 5 shows the variation in the extent of reduction as a function of the petroleum coke/quartz ratio. The extent of reduction increased with increasing petroleum coke additions up to about 28% by mass, after which further increases in petroleum coke decreased the extent of reduction and the yield of silicon metal. Although petroleum coke plays an important role in the reduction process, it contains ash with a low iron content and much finer material. The introduction of these fines into the production of metallurgical-grade silicon from Egyptian quartz
furnace results in poor furnace efficiency because it reduces the porosity of the charge, leading to the entrainment and subsequent blowing out of the fines with the furnace off-gas (Kuhlmann, 1965).

**Effect of varying coal additions**

Coal has a high electrical resistance and negligible quantities of volatile matter, but contains undesirable impurities such as iron. Figure 6 shows the extent of reduction as a function of the coal/quartz ratio. The extent of reduction increases with increasing coal additions until about 28% yield, beyond which the yield decreases with further coal additions. This may be due to the coal containing from about 5 to 15% ash (25% of which is iron oxide). The iron partitions into the silicon metal and affects the purity of the product (Kuhlmann, 1965).

From these results, the optimum charge composition was determined to be 64.63 wt.% quartz, 4.08 wt.% coal, 12.6 wt.% charcoal, 11.77 wt.% petroleum coke, and 6.92 wt.% wood chips.

**Effect of smelting time**

A series of smelting experiments was conducted on charges containing 64.63 wt.% quartz, 4.08 wt.% coal, 12.6 wt.% charcoal, 11.77 wt.% petroleum coke, and 6.92 wt.% wood chips at a C/SiO₂ ratio of 0.39 and power input of 36 kW, while varying the smelting time from 60 to 90 minutes. It was found (Figure 7) that the recovery of silicon metal increased from 28 to 40% as the smelting time was increased from 60 to 80 minutes. Further increases in smelting time up to 90 minutes decreased the silicon recovery to about 20%. This sharp decrease is due to the evaporation of the silicon metal product in the form of silica fumes as a result of the long smelting time. The optimum smelting time of 80 minutes yielded a silicon recovery of about 40%.

**Effect of furnace power input**

The smelting process was investigated at different power inputs ranging from 30 to 60 kW using charges having a C/SiO₂ ratio of 0.39 and a constant smelting time of 80 minutes. Figure 8 illustrates the variation in silicon recovery with power input. Silicon recovery was increased from about 20% to 75% as the power input was increased from 30 to 48 kW. With further increases in power, silicon recovery decreased, falling to about 30% at 60 kW power input.

The optimum recovery of about 75% of MG-Si was obtained at a power input of 48 kW. This may be due to the power input being sufficient to increase the furnace temperature to a level that enhances the reaction between SiO gas and SiC to form silicon metal. Higher power inputs, and higher temperatures, lead to the formation of excess SiO gas.
and consequently increase the silica loss (Mitsuhiro and Hirasawa, 2000).

**Evaluation of the MG-silicon product**

Figure 9 displays photographs of the MG-silicon product obtained at the optimum reduction-smelting conditions (carbon/silica mass ratio 0.39, smelting time 80 minutes, and power input 48 kW). The microstructure and chemical composition were investigated using XRD, SEM, and XRF.

**X-ray diffraction (XRD)**

Figure 10 shows the XRD spectrum of the MG-Si produced under optimum conditions, which indicates the presence of silicon metal only.

**X-ray fluorescence (XRF)**

The MG-silicon was chemically analysed by XRF to determine its purity. The results are summarized in Table II. The MG-silicon product contained about 97% Si and 1.15% Fe, with traces of other elements. It is clear that the MG-Si product had low traces of P, and it is noteworthy that no boron was detected. It is known that most impurities such as Fe, Al, and Ti, contained in the raw materials are transferred to the MG-silicon product.

**Morphology**

Scanning electron microscopy (SEM) images of the MG-silicon product are shown in Figure 11. A general distribution of silicon metal grains with traces of iron at the grain boundaries is evident.

**Conclusions**

MG-Si was produced from Egyptian quartz using a locally made submerged arc furnace, with a recovery of about 75% and product purity of about 97% silicon.

The optimum conditions for production of MG-Si are summarized as follows:
- Carbon/silica weight ratio (C/SiO₂) 0.39
- Charge composition quartz 64.65 wt.%, coal 4.08 wt.%, charcoal 12.6 wt.%, petroleum coke 11.77 wt.%, and wood chips 6.92 wt.%
- Smelting time 80 minutes
- Power input 48 kWh
- Electrical energy consumption approx. 19 kWh/kg of silicon metal produced
- Electrode consumption 0.11 kg/kg of silicon metal produced
- Silicon metal purity about 97%.

**References**


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<tr>
<th>XRF analysis of MG-Si product (wt.%)</th>
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<tr>
<td>Si</td>
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<td>96.94</td>
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SAIMM: Diamonds — Source to Use 2018 Conference
‘Thriving in Changing Times’

11 June 2018 — Workshop: SAMREC/SAMVAL Reporting – Diamond Projects
12–13 June 2018 — Conference • 14 June 2018 — Technical Visits

Birchwood Conference Centre (Jet Park, Johannesburg)

**BACKGROUND**
Being the seventh conference in the series, the *Diamonds — Source to Use* conferences target the full spectrum of the diamond pipeline from exploration through to sales and marketing. The last conference in this series (*Diamonds — Still Sparkling*) was held in Botswana in 2016 and it is now returning to Johannesburg, where it was last held in 2013.

**Keynote Speakers**

*The State of the Diamond Market*
E. Blom, Blom Diamonds (Pty) Ltd

*Financing Diamond Projects*
J. Campbell, Botswana Diamonds PLC

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