Cleaner production of soapstone in the Ouro Preto region of Brazil: a case study

M.L.M. Rodrigues, R.M.F. Lima

Department of Mining Engineering, Federal University of Ouro Preto, Ouro Preto, MG, Brazil

**Article Info**

Article history:
Received 1 October 2011
Received in revised form
23 March 2012
Accepted 25 March 2012
Available online 3 April 2012

**Abstract**

Soapstone is a metamorphic rock that is used for ornamental and construction purposes. Rock recovery during the exploitation process is low. Quarries in the state of Minas Gerais in Brazil sell small and low-quality blocks to artisans, who produce pans and souvenirs. This activity constitutes the main economic activity in several places within the Ouro Preto region, such as Santa Rita. Soapstone artisan workshops produce a large quantity of powder (approximately 10–15% rock recovery), which is discarded carelessly, often causing environmental problems. The objective of this study was to characterize and purify soapstone powder from the workshop of an artisan who works exclusively with rocks from Bandeiras-Santa Rita de Ouro Preto in order to identify potential applications for the residues and purified products. First, the mineralogical composition was determined by X-ray diffractometry, the size distribution was determined by sieving and the chemical composition of the residue was determined by inductively coupled plasma–optical emission spectroscopy (ICP-OES). Next, purification tests were performed by magnetic separation for fraction sizes larger than 74 μm. Fraction sizes smaller than 74 μm were purified by flotation and leaching with hydrochloric acid. This research shows that it is possible to recover almost all residues of both small and large size distributions for use as insecticide filler. In addition, it is possible to recover 100% and 96% of fraction sizes larger and smaller than 74 μm, respectively, for use in the paper industry (wallpaper and packaging). For more valuable applications, it is necessary to leach the purified material from flotation using hydrochloric acid. In addition to decreasing the environmental impact of soapstone powder discharge, it is possible for soapstone artisans to sell this powder for a profit, whether it is purified or not.

**Keywords:** Soapstone residues, Talc, Industrial mineral, Residue characterisation, Residue recovery, Flotation, Magnetic separation, Leaching process.

**1. Introduction**

Soapstone is a metamorphic rock that is used for ornamental and construction purposes. Rock recovery during the exploitation process is low. Quarries in the state of Minas Gerais in Brazil sell small and low-quality blocks to artisans, who produce pans and souvenirs. This activity constitutes the main economic activity in several places within the Ouro Preto region, such as Santa Rita. Soapstone artisan workshops produce a large quantity of powder (approximately 10–15% rock recovery), which is discarded carelessly, often causing environmental problems. The objective of this study was to characterize and purify soapstone powder from the workshop of an artisan who works exclusively with rocks from Bandeiras-Santa Rita de Ouro Preto in order to identify potential applications for the residues and purified products. First, the mineralogical composition was determined by X-ray diffractometry, the size distribution was determined by sieving and the chemical composition of the residue was determined by inductively coupled plasma–optical emission spectroscopy (ICP-OES). Next, purification tests were performed by magnetic separation for fraction sizes larger than 74 μm. Fraction sizes smaller than 74 μm were purified by flotation and leaching with hydrochloric acid. This research shows that it is possible to recover almost all residues of both small and large size distributions for use as insecticide filler. In addition, it is possible to recover 100% and 96% of fraction sizes larger and smaller than 74 μm, respectively, for use in the paper industry (wallpaper and packaging). For more valuable applications, it is necessary to leach the purified material from flotation using hydrochloric acid. In addition to decreasing the environmental impact of soapstone powder discharge, it is possible for soapstone artisans to sell this powder for a profit, whether it is purified or not.

In the Ouro Preto region of Brazil, located in Minas Gerais State, there are several soapstone quarries that exploit this rock as dimension stone. These places have rich deposits of talc, which occurs in the states of Minas Gerais (MG), Goiás (GO) and Bahia (BA).

In the Ouro Preto region of Brazil, located in Minas Gerais State, there are several soapstone quarries that exploit this rock as dimension stone. These places have rich deposits of talc, which occurs in the states of Minas Gerais (MG), Goiás (GO) and Bahia (BA).

In the Ouro Preto region of Brazil, located in Minas Gerais State, there are several soapstone quarries that exploit this rock as dimension stone. These places have rich deposits of talc, which occurs in the states of Minas Gerais (MG), Goiás (GO) and Bahia (BA).
souvenirs are produced. In the production of pans, for example, the recovery of rock is approximately 10 wt% (Santos, 2009), which means that the global recovery of soapstone exploited in quarries is only 40 wt%. Approximately 60 wt% of the rock is discharged as residue. Fig. 2 shows the main activities performed in a typical soapstone quarry in Santa Rita de Ouro Preto and the products that are sold to local artisans. Details of the production processes for souvenirs and pans are presented. A saw is applied to cut the rock and after a wood lathe is used to make and polish the produced objects. The large quantity of soapstone powder discharged from artisan workshops is also shown.

Lima et al. (2009) performed a technological characterisation of soapstone samples (Fig. 2(D)) used in artisan workshops from five different soapstone quarries in the Bandeiras region of Santa Rita – Ouro Preto. The minerals were identified using optical microscopy, X-ray diffraction and scanning Electron Microscopy with an Energy Dispersive System (SEM/EDS). The following minerals were found: silicates [talc (Mg₃(Si₄O₁₀)(OH)₂), chlinochlore ((Mg, Fe)(Si₅Al)₄O₁₀(OH)₁₀), phlogopite (K₂Mg₃(Si₇Al)O₁₀(F,OH)₂), tremolite/actinolite (Ca₂Mg₅(Si₈O₂₂)(OH)₂/Ca₂(Mg,Fe)₂Si₈O₂₂(OH)₂) and orthoclase (K₂Al₄Si₈O₂₆)], carbonates [magnesite (MgCO₃) and dolomite (CaMg(CO₃)₂)], sulphides [pyrite (FeS₂), arsenopyrite (FeAsS) and pentlandite (Fe₃Ni₅S₈)], oxides [magnetite(Fe₃O₄)/hematite (Fe₂O₃) and ilmenite (FeTiO₃)]. The apparent porosities of the soapstone rock samples were between 0.3 and 0.4%, and the apparent water absorption was approximately 0.11%. The Rockwell hardness of the samples was of a high range (73–84). This variability was attributed to the mineralogical heterogeneity on the faces analysed. The uniaxial compressive strength of the rock samples was between 20 and 25 MPa, and rupture occurred on the first or second cycles of the thermal tests.

The density of soapstone powder collected from the artisan workshop in the Bandeiras region varied from 2.73 to 2.83 (Rodrigues, 2010). The value of oil absorption of the purified soapstone powder residue was between 20 and 25 MPa, and rupture occurred on the first or second cycles of the thermal tests.

2. Materials and methods

Representative soapstone residues were collected in the workshop of an artisan who works exclusively with rocks originating from the Bandeiras region in Santa Rita de Ouro Preto. First, the powder residue was homogenised and quartered to obtain subsamples to determine the size distribution of the particles, as well as to perform mineralogical and chemical characterisation. The size distribution of the material was determined by wet sieving, and the fraction size — 74 μm was determined by laser diffraction (Cilas 1064). Then, chemical analyses of size fractions greater and lower than 74 μm were carried out using an Inductively Coupled Plasma — Optical Emission Spectrometer (Spectro model Ciros/CD), except for SiO₂ content, which was determined by the difference between 100% and the sum of all chemical elements, compound contents plus the loss on ignition (LOI).

The mineral composition of soapstone powder in both fraction size groups was performed using the X-ray diffraction (XRD) total powder method. The radiation used was Cu Kα. The goniometer velocity was 1.2 °/min, the counting time was of 58 min and data were collected from 2° to 70°. A Rigaku model 3550 diffractometer was used. Mineral identification was performed using the pattern diffraction tool of the JADE 7.0 software package.

The purification of soapstone powder was realised using magnetic separation (Carpo Inc. model WHIMS 3X4L — serial 210-97) and froth flotation (Cimaq cell). Leaching tests were performed with the concentrate of an optimised flotation test using hydrochloric acid (HCl).

In the magnetic concentration (fraction size above 74 μm), the influence of the variable magnetic field intensity (3000; 4300 and 4900 G) on the mass recovery and whiteness of purified talc was studied. These experiments were replicated for each condition. The experimental procedure for the wet magnetic concentration tests consisted of the following steps. First, the magnetic field intensity was fixed to the desired value. Then, with the water tap open (flow rate of 670 mL min⁻¹), a 100 g sample of dry soapstone powder was fed into the equipment. When it was observed that no particles were dropping into the non-magnetic bucket product, a new bucket was substituted in. Then, the equipment was turned off and the magnetic product was collected in the new bucket. Finally, both the magnetic and non-magnetic products were filtered, dried and weighed to calculate mass recovery. Finally, the products (magnetic and non-magnetic) were sampled to obtain subsamples for the determination of whiteness, CIE (L*a*b*) indices and chemical composition.

Purification of the soapstone powder (fraction size — 74 μm) was performed by flotation. Minitab 15 software was used to create and analyse the results of a fractional planning design (1/8) of eight factors (variables) on variable responses (mass recovery and whiteness). For each variable, two levels (minimum and maximum) were tested, as presented in Table 1. This analysis was performed on a total of 32 flotation tests.

The following experimental procedure was used for the flotation tests. A mass of soapstone powder (~74 μm) and the water quantities necessary to obtain the pulp density desired (10 or 20%) were put into a 2 L cell cube. The cell speed was then adjusted (800 or
1200 rpm). The air flow rates were 17.5 and 98 mL s⁻¹ for cell speeds of 800 and 1200 rpm, respectively. After 2 min, the depressant sodium silicate (500 or 1000 g/ton) was added and conditioned (5 or 10 min). During conditioning with the depressant, the pH of the pulp was adjusted (7 or 11). Next, kerosene (500 or 1000 g/ton) was added to the pulp and conditioned (5 or 10 min). The MIBC frother (75 or 150 g/ton) was then added to the pulp, and the pulp was conditioned (5 or 10 min). The air tap was opened, and the froth was collected for approximately 6 min. Both products (float and sink) were filtered and weighed to calculate the mass recovery. Finally, the products were homogenised, sampled and powdered to determine the whiteness, CIE (L*a*b*) indices and chemical composition.

The leaching tests of the purified product, obtained using the optimised flotation condition for the -74 μm soapstone powder, were carried out using hydrochloric acid in the following conditions: HCl concentration (2 and 4 M), temperature (50 °C and 80 °C) and leaching time (30 and 60 min). Minitab 15 software was used to create and analyse the results of a planning factorial design with three factors (variables) on variable response whiteness. This analysis was performed on a total of 16 leaching tests.

Desliming operation for soapstone powder (fraction size ~74 μm) was carried out with the objective of avoiding the heterocoagulation of ultrafine particles (<10 μm) of gangue minerals and talc, gangue ultrafine particles entraining in the concentrates and the slime coating of the gangue minerals by talc ultrafine particles and vice-versa (Sivamohan, 1990; Jameson et al., 1996).

### Table 1
Variables and their respective levels tested in the flotation tests.

<table>
<thead>
<tr>
<th>Factors (variables)</th>
<th>Level</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Minimum</td>
</tr>
<tr>
<td>Kerosene (g/ton)</td>
<td>500</td>
</tr>
<tr>
<td>Sodium metasilicate (g/ton)</td>
<td>500</td>
</tr>
<tr>
<td>Methyl isobutyl carbinitol – MIBC (g/ton)</td>
<td>75</td>
</tr>
<tr>
<td>Pulp pH</td>
<td>7</td>
</tr>
<tr>
<td>Pulp density (%)</td>
<td>10</td>
</tr>
<tr>
<td>Cell speed (rpm)</td>
<td>800</td>
</tr>
<tr>
<td>Kerosene conditioning time (min)</td>
<td>5</td>
</tr>
<tr>
<td>MIBC conditioning time (min)</td>
<td>5</td>
</tr>
</tbody>
</table>

Fig. 2. Typical operations in a soapstone quarry, the commercial products and the souvenir production process in artisan workshops in the Ouro Preto region: (A) Detail of the exploitation operation in a soapstone quarry of Santa Rita de Ouro Preto, (B) A rough block of soapstone, (C) Transport of the rough block to be exported, (D) Small pieces of soapstone to be used in artisan workshops, (E) Saw applied to cut the objects produced in artisan workshops, (F) An artisan performing his work and (G) Soapstone powder.

Fig. 3. Scanning electron microscope (SEM) image of purified talc from soapstone powder (Rodrigues and Lima, 2011).
The desliming process consisted of the following steps. First, the soapstone powder was put into a 4 L cell cube. Then, a volume of water sufficient to obtain a pulp density of 15% was added, and the flotation cell (Cimaq) was turned on at an adjusted speed of 1200 rpm. After 3 min, the flotation cell was turned off, and after 10 min, the mud was removed. This operation was performed twice. Finally, the deslimed soapstone powder was dried at 105 °C.

The leaching tests of purified soapstone by flotation consisted of the following steps. First, a pulp of 15% density was prepared with deionised water. Then, the speed of a constant temperature magnetic stirrer (BiomixXer, model 78HW-1) was adjusted to 640 rpm. The pulp was conditioned for 2 min at a temperature of 50 °C or 80 °C. After the pulp achieved the desired temperature (50 °C or 80 °C), hydrochloric acid was added (2 or 4 M), and the pulp was submitted to constant stirring for the specified time (30 or 60 min). The solids were then washed with deionised water, vacuum filtered with a Buchner funnel, dried at a temperature of 105 °C, weighed, homogenised, quartered and powdered to determine the whiteness, CIE indices (L*a*b*) and chemical composition.

The whiteness and CIE indices (L*a*b*) of the soapstone powder (+74 µm and –74 µm) and the purified talc were determined using a Technidyne Color Touch PC spectrometer. The samples were prepared for determination of the whiteness and CIE indices by powdering the sample in an orbital mill for 1 min. Next, the material was homogenised and quartered to obtain a mass subsample of 10 g that was placed on a watch glass to dry at a temperature of 105 °C. Then, a pellet of dried powder was made using a Powder Press Apparatus. To do this, the dried sample was transferred to a cylinder placed on a 2” square glass compression plate; the plunger was then placed onto the powder, and the mount was put into a powder press at 80 psi. Finally, the pellet was transferred to the spectrometer, and the whiteness and CIE indices (L*a*b*) were determined.

3. Results and discussion

3.1. Size distribution and mineralogy

Fig. 4 presents the size distribution of the soapstone powder residue obtained using wet sieving. Approximately 50% of the particles are under 74 µm, and approximately 27% of the particles are smaller than 37 µm. Thus, this material was classified in a 74-µm aperture sieve for the concentration tests performed by magnetic separation (fraction size over 74 µm) and froth flotation (fraction size under 74 µm).

Fig. 5 presents the size distribution of the soapstone powder residue (–74 µm), which was determined using laser diffraction (Cilas 1064). The d10 of soapstone powder without desliming was 7 µm and was 10 µm for deslimed soapstone.

The minerals identified, with their respective d values (basal spacing), in the X-ray patterns of all size fractions of soapstone powder analysed, are listed as follows: talc (9.205; 3.1021; 4.641; 2.4826; 1.5550; 1.3881 Å), kaolinite (7.062; 3.7221; 3.5540; 3.1021; 2.2261; 2.1874; 2.0035; 1.8067; 1.6702; 1.5550 Å), chlorite—serpentine (13.94; 7.062; 4.641; 3.554; 2.8453 Å), nimitte (13.94; 7.062; 4.7156; 3.5540; 2.8453 Å); magnesite (2.8453; 2.3294; 2.0829; 1.3881 Å) and dolomite (2.8453; 2.1874; 2.0035 Å). The minerals, sulphides (pyrite, arsenopyrite, pentlandite) and oxides (magnetite/hematite) were identified in soapstone from the Bandeira region (Lima et al., 2009) but were not detected in the X-ray patterns of the size fractions, probably due to the small proportion (less than 2%) of these minerals in the subsamples analysed and the presence of several minerals, which can make mineral identification in the X-ray patterns difficult. The identified kaolinite mineral may be a result of orthoclase alteration, as identified by Lima et al. (2009).

3.2. Purification tests in bench scale

3.2.1. Purification tests for the +74 µm sample

Table 2 presents the mass recovery, whiteness and CIE indices of the feed and the purified products from the three magnetic separation tests performed using the +74 µm fraction size sample of soapstone powder in each test condition. Table 3 presents the average chemical composition and LOI of the feed and the purified products.

As can be observed in Table 2, the mass recovery of the soapstone residue varied from approximately 62–67%. Piga and Maruzzo (1992) proposed a preconcentration step for an Italian talc ore containing chloride, carbonates, magnetite and sulphides by low-intensity magnetic separation to remove the magnetite mineral before the flotation test, once the high intensity field conducted to very low recovery of talc.

The whiteness of the purified product in a 4900 G field (Table 2) increased by 6% compared with the feed whiteness (59.72%). This increase in whiteness is probably related to the removal of the iron oxides (magnetite, ilmenite and hematite) present in small proportions in this material. The CIE indices of the purified products did not change significantly in comparison with the feed CIE

Table 2

<table>
<thead>
<tr>
<th>Purified product</th>
<th>Mass recovery (%)</th>
<th>Whiteness (ISO)</th>
<th>CIE group L* a* b*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feed</td>
<td>100.00</td>
<td>59.72</td>
<td>89.10 0.58 8.05</td>
</tr>
<tr>
<td>3000 G</td>
<td>62.53</td>
<td>64.75</td>
<td>88.67 0.68 7.86</td>
</tr>
<tr>
<td>4300 G</td>
<td>62.39</td>
<td>65.22</td>
<td>89.00 0.70 8.00</td>
</tr>
<tr>
<td>4900 G</td>
<td>67.42</td>
<td>65.44</td>
<td>89.05 0.68 7.88</td>
</tr>
</tbody>
</table>

Fig. 4. Particle size distribution of soapstone powder residue.

Fig. 5. Particle size distribution of soapstone powder residue (–74 µm).

indices. In terms of whiteness, the feed and products purified by magnetic separation were within the specifications for paper filler (wallpaper and packaging—whiteness ISO of 54–75%) (Pontes and Almeida, 2008). The value of whiteness obtained and the chemical composition of the purified product (Table 3) are nearly within paint industry specifications (Pontes and Almeida, 2008). Based on these results, it is possible to recover 100% of the -74 μm residues for use as filler in the paper industry, representing a total recovery of 70% of the exploited rock. For the paint industry, the recoveries of 62–67% observed in the non-magnetic product correspond to an 18–20% increase in the global recovery (40%) of exploited soapstone for dimension stone and artisan workshops.

### 3.2.2. Purification tests for the -74 μm sample

The Pareto’s chart and surface plots of the studied effects on the mass recovery and whiteness of the flotation tests conducted with soapstone powder (fraction size -74 μm) are presented in Figs. 6–9.

As can be observed in Fig. 6, among the studied effects, only the pulp density and the MIBC dosage were significant for mass recovery. In the case of whiteness, only the pulp density was significant (Fig. 7). The mass recoveries were higher when the pulp densities and MIBC dosages were higher (Fig. 8). The opposite occurred for whiteness (Fig. 9). After the optimisation of the factorial planning experimental design using Minitab 15 for a target of 80% mass recovery and 77% whiteness ISO, the optimised response was obtained using a pulp density of 14%, an MIBC dosage of 150 g/ton and a kerosene dosage of 500 g/ton. In this case, the predicted values of the mass recovery and whiteness were 80% and 64%, respectively. Thus, the flotation tests in the bench with soapstone powder, without and with desliming, were conducted under the optimised conditions for the significant effects (pulp density of 14% and MIBC dosage of 150 g/ton). All other effects were fixed at their minimum values (sodium metassilicate - 500 g/ton, pH - 7, cell speed - 800 rpm, kerosene and MIBC conditioning time - 5 min). For a mixing of four talc ores from the Eastern Desert of Egypt (Shalatin area) (tremolite-talc-chlorite-shist; antigorite-serpentine; talc-shist and talc-chlorite-shist), the optimum conditions of the flotation tests were obtained at pH 11, a depressant dosage (sodium hexametaphosphate) of 1 kg/ton, a collector dosage (oleic acid in an equal mixture with kerosene) of 1.2 kg/ton and a pulp density of 200 g L⁻¹ (~18%) (Ahmed et al., 2007). For a representative talc sample composed of magnesite (36.47%), talc (48.69%), serpentine (2.41%), dolomite (9.63%), and quartz (2.47%) from the Wadi El-Barramiya region in the Eastern Desert of Egypt, the optimum pH for natural talc floatability was neutral (Yehia and Al-Wakell, 2000).

Afterwards, a leaching study of the purified talc from the optimised flotation test was conducted with HCl using a factorial planning design. The Pareto’s chart and surface plots of the studied effects (HCl dosage and leaching temperature) on whiteness, carried out with the purified rougher flotation product of soapstone powder (fraction size -74 μm) without desliming, are presented in Figs. 10–12.

As can be observed in the Pareto’s chart (Fig. 10), all of the studied effects were significant in the whiteness response variable. The surface plots (Figs. 11 and 12) show that the whiteness was higher (84%) for a dosage of 4 M of HCl, a temperature of 80 °C and a time of 60 min of leaching (Fig. 12), which was confirmed as the optimised condition for the levels of the studied variables, using the optimisation tool in Minitab 15 software. This result indicates a gain of 20.5% compared with the rougher purified flotation product (whiteness of 63.5%). An Argentine talc ore concentrate, with 10–20% carbonates (magnesite, dolomite and calcite) with an

---

**Table 3**

<table>
<thead>
<tr>
<th>Purified product</th>
<th>Content</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>As</td>
<td>Cu</td>
</tr>
<tr>
<td>Feed</td>
<td>11.1</td>
<td>9.59</td>
</tr>
<tr>
<td>3000 G</td>
<td>&lt;9.45</td>
<td>7.57</td>
</tr>
<tr>
<td>4300 G</td>
<td>&lt;9.45</td>
<td>8.56</td>
</tr>
<tr>
<td>4900 G</td>
<td>&lt;9.45</td>
<td>7.35</td>
</tr>
</tbody>
</table>

G – Gauss.
The whiteness of the product was 59% (Table 4), an increase of 8% compared with the feed whiteness (51.34%). As can be seen in Table 4, the values of lightness ($L^*$) increase and the values of redness ($a^*$) and yellowness ($b^*$) decrease from the rougher concentrate to leached product. Thus, by observing the chemical compositions of all purified products (presented in Table 5), the contents of Fe$_2$O$_3$, Al$_2$O$_3$, TiO$_2$, CaO, LOI, Cu and Zn in the purified products decreased compared with the feed, which is consistent with the whiteness and lightness of these purified products. These decreases are probably related to the partial removal of the gangue minerals (sulphides, oxides, carbonates, chlinochlore, chlorite–serpentine and nimite), identified in this material by Lima et al. (2009).

Higher whiteness (~84%) and a decrease in the indices $a^*$ (redness) and $b^*$ (yellowness) were observed in the leaching product compared to the flotation concentrates (Table 4). The SiO$_2$ assay increased to 60.3% and the Fe$_2$O$_3$ assay decreased to 3.42%. The final product, after leaching the flotation concentrate of Egyptian talc ore with diluted hydrochloric acid (10%), showed an initial whiteness of 65.5%, exhibited an increase of 4–7% after leaching with hydrochloric acid (concentration of 25–100%) (Sarquis and Gonzalez, 1998). Table 4 presents the results of all purification tests of soapstone powder (fraction size ~74 μm) performed in the optimised conditions.

As can be observed in Table 4, the mass recovery under the experimental optimised conditions was approximately 7% higher than the optimised value predicted using Minitab 15 (80%), and the whiteness (63.5%) was slightly lower than the optimised value (64.2%) predicted using Minitab 15. In this case, an increase of 12% was observed in the whiteness of the soapstone residue compared with the feed whiteness (51.34%). This result represents the possibility of obtaining an increase of approximately 25% of the global recovery of the exploited soapstone. After two cleaner flotation tests, a product with a whiteness of 68.7% was obtained, which represents an increase of 5% in whiteness compared with the experimental rougher test. However, the mass recovery decreased by approximately 23%. The increase of whiteness obtained with the two cleaning steps was not enough to justify these steps, once the whiteness obtained is just within the specifications for filler paper as the rougher purified product, despite the whiteness gain.

After desliming the soapstone residue (fraction size ~74 μm), the whiteness of the product was 59% (Table 4), an increase of 8% compared with the feed whiteness (51.34%). The mass discharged in this case was only 6 wt%. With this value of whiteness, the fraction size ~74 μm (51.34%) could be applied only in the insecticide filler. After desliming, 94 wt% of this material could be applied as filler paper, which means approximately 30% and 28% global soapstone recovery for application in the insecticide and paper industries, respectively. The mass recovery of the rougher purified product was 81% of the global residue (~74 μm), and the whiteness obtained was 67%, which is approximately 3.5% higher than that of the rougher purified product without desliming (Table 4); after two cleaning steps, it was approximately 71%. As in the previous case, a product with specifications within those for paper filler and paint was obtained once the particles were lamellar and the oil absorption was 28.5 (g/100 g) (Rodrigues and Lima, 2011). Despite the whiteness gained in the cleaning steps, the cleaning steps are not justified because the mass recovery drops drastically and the whiteness of obtained product is not enough for more valuable use.

As can be seen in Table 4, the values of lightness ($L^*$) increase and the values of redness ($a^*$) and yellowness ($b^*$) decrease from the rougher concentrate to leached product. Thus, by observing the chemical compositions of all purified products (presented in Table 5), the contents of Fe$_2$O$_3$, Al$_2$O$_3$, TiO$_2$, CaO, LOI, Cu and Zn in the purified products decreased compared with the feed, which is consistent with the whiteness and lightness of these purified products. These decreases are probably related to the partial removal of the gangue minerals (sulphides, oxides, carbonates, chlinochlore, chlorite–serpentine and nimite), identified in this material by Lima et al. (2009).

Higher whiteness (~84%) and a decrease in the indices $a^*$ (redness) and $b^*$ (yellowness) were observed in the leaching product compared to the flotation concentrates (Table 4). The SiO$_2$ assay increased to 60.3% and the Fe$_2$O$_3$ assay decreased to 3.42%. The final product, after leaching the flotation concentrate of Egyptian talc ore with diluted hydrochloric acid (10%), showed an
increased in the SiO₂ assay and a decrease in the Fe₂O₃ assay (Ahmed et al., 2007).

In terms of the chemical composition of the leaching product presented in Table 5, it can be observed that all impurities, except the Cu grade, were smaller in the leaching product than in the purified rougher flotation product. The opposite was observed with MgO.

Castillo et al. (2011), using XRD and FTIR (Fourier Transform Infrared Spectroscopy), identified the following minerals in a talc sample: talc, chlorite, dolomite, calcite and magnesite. After leaching this sample with hydrochloric acid, they observed a decrease in the Fe₂O₃ assay and loss on ignition assay, which was attributed to the removal of chlorite and carbonates from the sample, evidenced by the fact that the peaks assigned to these minerals in the X-ray pattern and the infrared spectrum disappeared. SiO₂ and MgO are the main constituents of talc (Mg₃(SiO₄)₁₀(OH)₂); thus, the increase in MgO content in the leached product is related to the increased proportion of talc and is mainly due to the removal of chlorite and carbonate by hydrochloric acid. Therefore, based on the chemical composition, whiteness (83.99%) and CIE indices (L* = 94.62; a* = −0.11 and b* = 2.22) that were obtained for the leaching product, this purified product achieved the necessary specificities for use in the paint, paper (filler), insecticide, textile and plastic industries.

Table 4

<table>
<thead>
<tr>
<th>Product</th>
<th>Mass recovery (%)</th>
<th>Whiteness (ISO)</th>
<th>CIE indices</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rougher concentrate</td>
<td>86.65</td>
<td>63.45</td>
<td>L* = 88.82</td>
</tr>
<tr>
<td>Second cleaner concentrate</td>
<td>64.01</td>
<td>68.34</td>
<td>a* = 1.27</td>
</tr>
<tr>
<td>Deslimed rougher concentrate</td>
<td>81.00</td>
<td>67.21</td>
<td>b* = 9.30</td>
</tr>
<tr>
<td>Second deslimed cleaner concentrate</td>
<td>60.85</td>
<td>70.77</td>
<td></td>
</tr>
<tr>
<td>Leaching product</td>
<td>83.99</td>
<td>94.62</td>
<td></td>
</tr>
<tr>
<td>Feed (−74 μm)</td>
<td>100.00</td>
<td>81.92</td>
<td></td>
</tr>
<tr>
<td>Deslimed feed (−74 μm)</td>
<td>94.00</td>
<td>87.14</td>
<td></td>
</tr>
</tbody>
</table>

Table 5

<table>
<thead>
<tr>
<th>Product</th>
<th>Content (%)</th>
<th>Content (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MgO</td>
<td>SiO₂</td>
</tr>
<tr>
<td>Feed (−74 μm)</td>
<td>74m</td>
<td>27.7</td>
</tr>
<tr>
<td>Rougher concentrate</td>
<td>74m</td>
<td>28.3</td>
</tr>
<tr>
<td>Second cleaner concentrate</td>
<td>74m</td>
<td>27.7</td>
</tr>
<tr>
<td>Deslimed rougher concentrate</td>
<td>74m</td>
<td>27.9</td>
</tr>
<tr>
<td>Second deslimed cleaner concentrate</td>
<td>74m</td>
<td>29.1</td>
</tr>
<tr>
<td>Leaching product</td>
<td>74m</td>
<td>30.0</td>
</tr>
</tbody>
</table>

Fig. 12. Whiteness surface plot of the purified talc as a function of HCl dosage and leaching temperature for 60 min.

4. Conclusion

The studies conducted using soapstone powder from the Banderas region of Santa Rita de Ouro Preto show that approximately 50% of the particles are smaller than 74 μm and approximately 27% are smaller than 37 μm. The identified minerals, in all fraction sizes, were as follows: talc, kaolinite, chlorite—serpentinite, magnesite, dolomite and nimite. Sulphides and oxides were not identified, probably due to their small proportion in the subsamples analysed and the presence of several minerals, which can make mineral identification in the X-ray patterns difficult. It is possible to recover 100% of the soapstone residue for use as insecticide filler. The fraction size above 74 μm can be used in the paper industry (wallpaper and packaging). The same application is possible for more valuable applications, such as usage in the plastic industry, the material must be leached by hydrochloric acid.

Acknowledgements

Financial support from FAPEMIG and CNPq for scholarships for the authors is gratefully acknowledged. The authors are grateful to reviewers.

List of symbols

- L*: lightness
- a*: redness
- b*: yellowness
- LOI: loss on ignition
- MIBC: methyl isobutyl carbinol
- G: Gauss
- M: molar
- L: liter
- mL: milliliter
References


