Characterization of a Selective Surface Based on Chromium, Iron and Aluminum Oxides for Application in Solar-Thermal Collectors

Kaio Hemerson Dutra*, Francisco Nivaldo Aguiar Freire*, Diego Caitano Pinho*, Felipe Alves Albuquerque Araújo*

*Universidade Federal do Ceará, Laboratório de Filmes Finos em Energia Renovável, Fortaleza, CE, Brasil.

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The depletion of conventional energy sources and the concern with the environmental impacts of the use of fossil fuels sparked interest in researching new energy sources and improving existing processes. In this context, the solar energy presents itself as one of the most promising energy sources on the planet, given its wide availability and applicability in thermal processes. However, its use still represents a great technological and economic challenge, because many systems that use this energy still have low efficiency and high cost, which makes them uncompetitive in competition with systems using other energies. With that, the search for the improvement of the processes awakens the researches in more adequate and cheaper materials, which represents a great scientific potential in the evolution of these technologies. Thus, the present work proposes to obtain and analyze a selective surface for applications in solar-thermal collectors, using CRFO (compound formed by chromium and iron oxides), varying the percentage by weight, and then mixed with aluminum oxide. For the tests, an experimental bench was built to simulate a flat plate solar-thermal collector in real operating conditions. Some characterizations were carried out, such as: techniques scanning electron microscopy (SEM), infrared analysis and UV-VIS absorptivity determination, as well as the graphs with surface temperatures and with radiation during the tests in the Sun for the composite (CRFO and aluminum oxide) and for the commercial surface (MRTiNOX). With the results obtained, it was verified in the field test that one of the produced selective surfaces obtained a result of absorptivity of 0.94 and 0.0349 of emissivity, which is close to that of the commercial surface. The results were confirmed in the characterization by UV-VIS, which showed high absorptivity values in the visible light range.

Keywords: Solar energy, selective surface, efficiency, collectors.

1. Introduction

The use of solar energy is already a reality, however, although it is an available and abundant energy source, the systems for capturing and transforming this energy are still at an economic disadvantage compared to processes that use conventional energy, which opens up great possibilities for research into the improvement of processes and the use of materials that provide increased efficiency and reduced costs.

In the specific case of solar-thermal energy, solar radiation is absorbed by solar-thermal collectors, these are composed of surfaces capable of absorbing solar radiation and converting it into thermal energy, transporting it by conduction. The absorbed radiation is conducted by the absorbers and can heat working fluids, liquids or gases. Systems using solar-thermal energy can be applied for heating water in homes, hotels and hospitals, other systems can be used for drying and dehydrating fruits in the food industry and desalination of seawater.

In practical applications of high efficiency, for the conversion of radiation from the sun into thermal energy, for heating working fluids, solar-thermal collectors are used with coatings on their radiation-receiving surfaces. This coating can consist of a black or selective surface, which is deposited on a base material, absorbing solar radiation and playing an important role in terms of absorption and emission of radiation, directly influencing the thermal transformation process. Therefore, it is necessary that this surface absorbs radiation well in the solar spectrum (ultraviolet and visible) and, at the same time, has a low emissivity, avoiding heat loss to the environment. The selective surface is usually composed of a thin film applied on a thermally conductive substrate, which can be proceed by an antioxidant layer and followed by an anti-reflective layer, in direct contact with the environment.

The purpose of using selective surfaces, more efficient than black surfaces, in the base material (material that conducts thermal energy) is to maximize energy absorption, but without providing large emissions of thermal radiation to the environment, so that the net energy gain is increased, thus improving the efficiency of the system.

The objective of this work is to synthesize, analyze and test selective surfaces for application in flat plate solar-thermal collectors using CRFO (compound formed by chromium and iron oxides) and aluminum oxide. The goal is to obtain a surface similar to or superior to selective surfaces already on the market.

*e-mail: kaiohdutra@gmail.com
2. Materials and Methods

2.1. Production of CRFO

The preparation of the material, which was used, was based on the methodology proposed by Dutra et al. to obtain the CRFO100 (Cr0.75Fe1.25O3). The procedure consisted of a mixture of 127.3g of iron oxide Fe2O3 (Aldrich, 99+%) with 72.7g chromium oxide Cr2O3. The iron and chromium oxides were duly ground in a planetary mill using zirconium balls, for this purpose, polyamide (technil) pans were used. After milling, the oxides were mixed and a mixture of 200g was thus obtained. After mixing, the material was subjected to a heat treatment at a temperature of 1300°C with a heating rate of 1°C per minute, maintaining the maximum temperature for 5 hours. The procedure was carried out in a muffle oven.

After the calcination process, which takes place during the 5 hours of permanence at a temperature of 1300°C, the oven was turned off and put to cool in a natural way, avoiding high cooling rates. As a result, the CRFO100 is obtained (Cr0.75Fe1.25O3), iron and chromium have the same oxidation number, Fe3+ and Cr3+, and thus share the same site in the compound.

As shown in Figure 1, the compound obtained has a very high granulation, which would make the manufacturing process of selective surfaces difficult.

A high granulation can disrupt the production process of selective surfaces, leaving them very rough, making the process of depositing on substrates difficult. In this way, the CRFO was ground by a mortar and pestle, thus obtaining a material with lower granulation. The ground material was subjected to a granulometry sieve, in order to separate the CRFO in homogeneously granulated portions. For manufacturing tests on selective surfaces, the following granulations were separated: 42 mesh (0.355mm sieve opening), 100 mesh (0.15mm opening) and 170 mesh (0.09mm sieve opening).

2.2. Surface production

The composites were produced using CRFO and aluminum oxide, being combined in the following proportions: 25% - 75%, 50% - 50% and 75% - 25%. All samples were produced with 100 mesh CRFO granulation, due to the difficulty of handling the CRFO obtained with a 170 mesh sieve, because, due to the grain size, much of it was lost in the air in the process, and with the CRFO obtained with 42 mesh sieve, the surfaces produced showed roughness.

The samples were made by mixing the powder of each component, so it was necessary, to make their adhesion to the base material, the use of binders, transforming them into elements with viscous characteristics and adherent to metallic surfaces. The binder used was the epoxy adhesive MRArraudite, using 35% in weight relative to the total sample.

To finish the deposited compound, was added B2O3/Bi2O3 flux (1:1 molar mixture of boron-bismuth) added in a proportion of 3% by weight relative to the total sample. The composition of each surface produced is shown in Table 1.

After obtaining the compounds, the deposition was made on a copper substrate (4cm x 2cm plate, with approximate thickness of 0.3mm), through the process known as screen printing, in which a spatula is used to spread them about this. Subsequently, the surfaces received heat treatment (sintering) in a resistive oven. Several tests were carried out until the ideal parameters were found for the samples to adhere completely, homogeneously and with little roughness to the copper substrate. The sintering parameters are shown in Figure 2.

The heating process from ambient temperature to 50°C at a rate of 0.5°C/min; stay at 50°C for 20 minutes; heating from 50°C to 100°C at a rate of 0.5°C/min; stay at 100°C for 20 minutes; heating from 100°C to 150°C at a rate of 0.5°C/min; stay at 150°C for 5 hours; cooling from 150°C to ambient temperature inside the oven.

2.3. Test bench

To perform the evaluation of selective surfaces, was constructed a field test bench, Figure 3. The bench was constructed of 2.5 cm thick muiracatiara wooden, with a MRVivix float glass cover, colorless, tempered, measuring 1mx12.6 cm and 4 mm thick and insulated at the bottom with glass wool 11.5 cm x 8.5cm and 5cm thickness. The bench has nine compartments for sheltering the selective surfaces that are deposited above the insulation of each compartment. The bench simulates real operating conditions of the fins of a thermal solar-thermal collector.

Table 1. Composition of selective surfaces obtained in percentage terms.

| Components      | Surface 1 | Surface 2 | Surface 3 |
|-----------------|-----------|-----------|-----------|
| CRFO [%]        | 15.5      | 31        | 46.5      |
| aluminum oxide [%] | 46.5      | 31        | 15.5      |
| Flux [%]        | 3         | 3         | 3         |
| MRArraudite [%] | 35        | 35        | 35        |
| Total mass [g]  | 3.2       | 3.2       | 3.2       |
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To measure the temperature of the selective surfaces, the environment, glass covering and insulation, type “k” thermocouples were installed. To measure the solar radiation, an MREppley Horizontal Pyranometer was used. To record and store the data obtained during the test, an Omega datalogger was used.

2.4. Energy balance in the test bench

To analyze and compare the performance of the selective surfaces, an energy balance was performed in a bench test. Figure 4 shows the representation of the energy balance in the system.

The input energy ($Q_{in}$), Equation 1, is the net solar radiation ($Q_{sol}$), that passes through the glass cover, with transmissivity $\tau_v$, and is absorbed by the selective surface, with absorptivity $\alpha_s$.

$$Q_{in} = Q_{sol} - Q_{ab\_glass} - Q_{ref} = R_{sol}\tau_v\alpha_s A_P$$  \hspace{1cm} (1)

Where $Q_{ab\_glass}$ represents the heat absorbed by the glass, $Q_{ref}$ the reflective heat by the selective surface, $R_{sol}$ the solar irradiation and $A_P$ the selective surface area.

The output energy ($Q_{out}$), Equation 2, is related to the thermal losses through the collector by thermal radiation ($Q_{rad}$), convection ($Q_{conv}$) and conduction ($Q_{cond}$).

$$Q_{out} = Q_{rad} - Q_{conv} - Q_{cond}$$  \hspace{1cm} (2)
The Equation 3 gives the radiation heat loss. Where \( \sigma \) is the Stefan-Boltzman constant (\( \sigma = 5.67 \times 10^{-8} \text{W/m}^2\text{K}^4 \)), \( \varepsilon \) is the emissivity of the selective surface, \( T_p \) is the plate temperature and \( T_b \) is the temperature near to the outer wall of the structure.

\[
Q_{\text{rad}} = \varepsilon A \sigma \left( T_p^4 - T_b^4 \right)
\]

(3)

The Equation 4 gives the natural convection heat loss in the compartment between the selective surface and the glass cover. Where \( h \) is the heat transfer coefficient by convection and \( T_{cg} \) is the temperature of the glass cover.

\[
Q_{\text{conv}} = h A (T_p - T_{cg})
\]

(4)

The energy loss by conduction occurs mainly through thermal insulation, as shown in Figure 4. Equation 5 represents the energy losses by conduction. Where \( L \) is the thickness of the insulation and \( K \) is the thermal conductivity.

\[
Q_{\text{cond}} = \left( \frac{K}{L} \right) A (T_p - T_b)
\]

(5)

The energy balance for the control volume shown in Figure 3 is given by Equation 6. Where the heat generated (\( Q_{\text{generated}} \)) and the heat accumulated (\( Q_{\text{accumulated}} \)) are zero because there is no internal heat generation and operate in permanent regime.

\[
Q_{\text{in}} + Q_{\text{generated}} = Q_{\text{out}} + Q_{\text{accumulated}}
\]

(6)

\[
Q_{\text{in}} = Q_{\text{out}}
\]

Then, substituting Equations 1 to 5 into Equation 6 and adopting the proposed considerations, the energy balance equation can be written as represented in Equation 7.

\[
R_{\text{sol}} r_\alpha \sigma A_p = \varepsilon A \sigma \left( T_p^4 - T_b^4 \right) + h A (T_p - T_{cg}) + \left( \frac{K}{L} \right) A (T_p - T_b)
\]

(7)

From the energy balance performed, it is possible to calculate the absorptivity and emissivity properties of selective surfaces based on data obtained in field tests. However, it is necessary to use Equation 8, which corresponds to the useful energy (\( Q_{\text{useful}} \)) supplied by the system (solar-thermal collector) to the working fluid. Where \( T_a \) is the ambient temperature.

\[
Q_{\text{useful}} = A_p R_{\text{sol}} r_\alpha \sigma_A - A_p U_1 (T_p - T_a)
\]

(8)

For the system shown in Figure 3, there is no use of thermal fluid, thus \( Q_{\text{useful}} = 0 \). Therefore, Equation 8 can be written as represented in Equation 9.

\[
U_1 = \frac{R_{\text{sol}} r_\alpha \sigma_A}{(T_p - T_a)}
\]

(9)

2.5. Performed test

To produce selective surfaces, several tests were carried out, still in the production phase, in order to obtain the ideal parameters for the heat treatment (sintering), as presented in section 2.2. Thus, in section 3.1 we will present some of the parameters that were tested and did not get the expected surface. For characterization and verification of the performance of the new selective surfaces obtained, the following tests were performed:

1. Absorptivity graphs, in the range of 190nm to 900nm, obtained by the analysis of the selective
surfaces deposited in glass, realized in a Cary Series UV-Vis Spectrophotometer;
2. Infrared analysis, with KBr inserts, in the Shimadzu apparatus, IRTracer-100 model, in the range of 400 to 4000 cm⁻¹, in order to determine if any composites were formed after the surface preparation in the resistive kiln;
3. Verification of the morphological structure and chemical composition resulting from the synthesis of the material by means of EDX micro scanning electron microscopy (SEM);
4. Performance tests of the selective surfaces in the test bench in real environmental conditions, with measurements of temperature, absorptivity and emissivity.

3. Results and Discussions

3.1. Heat treatment
The purpose of the heat treatment was for the binder to evaporate and, thus, the sintering process of the deposited compounds was completed. For the heat treatment tests, several surfaces with the same characteristics, surface 2 (described in Table 1), were produced and submitted to several resistive oven tests with different parameters, in order to obtain the parameter considered ideal for these surfaces. The results below present some of the parameters used, as well as the surfaces after heat treatment.

The first configuration tested in the resistive oven, heat treatment 1, the heating heating process from ambient temperature to 200°C at a rate of 1°C/min, holding at 200°C for 5 hours and then cooling at the rate 1°C/min from 200°C to ambient temperature inside the oven. Figure 5 shows the configuration used in the sintering process and the surface obtained.

Heat treatment 2, the heating process from ambient temperature to 100°C at a rate of 0.5°C/min; stay at 100°C for 20 minutes; heating from 100°C to 150°C at a rate of 0.5°C/min; holding at 150°C for 20 minutes; heating from 150°C to 200°C at a rate of 0.5°C/min; stay at 200°C for 5 hours; cooling from 200°C to ambient temperature inside the oven. Figure 6 shows the configuration used in the sintering process and the surface obtained.

Figure 5. Thermal treatment 1.

Figure 6. Thermal treatment 2.
For heat treatment 3, the heating process from ambient temperature to 50°C at a rate of 0.5°C/min; stay at 50°C for 20 minutes; heating from 50°C to 100°C at a rate of 0.5°C/min; stay at 100°C for 20 minutes; heating from 100°C to 150°C at a rate of 0.5°C/min; stay at 150°C for 5 hours; cooling from 150°C to ambient temperature inside the oven. Figure 7 shows the configuration used in the sintering process and the surface obtained.

Several other heat treatments were performed with different configurations, however the surfaces obtained were similar to the surfaces presented in heat treatment 1 and 2. In these thermal treatments (1 and 2) the surfaces showed corrugated aspects, since higher temperatures were used in them. The image of the sample, obtained with heat treatment 3, shows that the use of these heating parameters resulted in a selective surface perfectly adhered to the metallic substrate, but it presents a rough appearance, possibly arising from the particle size of the CRFO.

3.2. Characterization by UV-VIS

Absorbance is an important property to be analyzed on a selective surface, as it is directly linked to the ability to absorb thermal energy. For this characterization, the Shimadzu spectrophotometer, UV-2600 was used. Figure 8 shows...
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It is observed that on the three surfaces, $25\% \text{Al}_2\text{O}_3$ + $75\% \text{CRFO}$, $50\% \text{Al}_2\text{O}_3$ + $50\% \text{CRFO}$ and $75\% \text{Al}_2\text{O}_3$ + $25\% \text{CRFO}$, the behavior is similar. The absorptivity in the visible light range (400 nm to 750 nm) has a maximum value, which is a positive aspect for solar energy absorption. It is also verified that there is a tendency to fall after the visible light band, at approximately 750 nm, showing a more accentuated fall in the composite surface of $75\% \text{Al}_2\text{O}_3$ + $25\% \text{CRFO}$.

It is possible to observe that the $25\% \text{Al}_2\text{O}_3$ + $75\% \text{CRFO}$ surface presented an average absorptivity value of 0.947, in the visible light range, which is higher than the other samples. The samples CRFO, $50\% \text{Al}_2\text{O}_3$ + $50\% \text{CRFO}$ and $75\% \text{Al}_2\text{O}_3$ + $25\% \text{CRFO}$ presented average absorptivity values of 0.937 and 0.933 respectively.

3.3. Infrared analysis

The purpose of conducting this characterization is to identify the emergence of any component after the heating process of the selective surfaces in resistive oven. Infrared analysis was proceeded with KBr pellets on Shimadzu IRTracer-100, range of 400 - 4000 cm$^{-1}$.

Thus, the infrared analysis was performed before the deposition of the compound on the substrate (not deposited) and after the heat treatment (deposited) for the three surfaces defined in Table 1. Figure 9 presents the infrared analysis for the surfaces.

In Figure 9, five main peaks are observed, in the following frequency regions: 250 - 500 cm$^{-1}$, approximately 1000 cm$^{-1}$, 1500 - 2000 cm$^{-1}$, 2750 - 3000 cm$^{-1}$ and approximately 3500 cm$^{-1}$. In all samples, the shape of the peaks coincides with the non-deposited compound with the deposited compound, differing only in peak intensity, and no new components are obtained after heating.

3.4. Analyses in the SEM

Through micrographs, obtained by SEM, it was possible to evaluate the arrangement of the composite material on the copper substrate, after completion of the heat treatment. The results of the micrographs are shown in Figures 10 to 12 with a magnification of 10000x and 2000x, for the selective surfaces described in Table 1.

For the selective surface containing $25\% \text{Al}_2\text{O}_3$ + $75\% \text{CRFO}$, it is possible to verify exposed grains containing carbon, aluminum, chromium, iron and bismuth, the micrograph is shown in Figure 10. There is little unfilled space, the substrate almost entirely occupied.

For the selective surface containing $50\% \text{Al}_2\text{O}_3$ + $50\% \text{CRFO}$ it is possible to verify exposed grains containing carbon, aluminum, chromium, iron and bismuth, the micrograph is shown in Figure 11. A large area of unfilled copper substrate is also observed.

For the selective surface containing $75\% \text{Al}_2\text{O}_3$ + $25\% \text{CRFO}$, it is possible to verify exposed grains containing carbon, aluminum, chromium, iron and bismuth, the micrograph is shown in Figure 12. There is little unfilled space, the substrate being almost completely occupied. Carbon is observed on all selective surfaces, possibly from the MRAraldite.

In general, in Figures 10 to 12, it is possible to verify a sample with good deposition, since the grains spread over the surfaces are evenly spread.

3.5. Field tests

On April 19, 2021, the field test was conducted with temperature measurements of the selective surfaces, glass...
cover, environment and insulation. The test analyzed the behavior of the selective surfaces exposed to the sun.

The surfaces tested were: 25% $\text{Al}_2\text{O}_3 + 75\%$ CRFO, 50% $\text{Al}_2\text{O}_3 + 50\%$ CRFO, 75% $\text{Al}_2\text{O}_3 + 25\%$ CRFO and TiNOX. On this test day (April 19, 2021) the sky was clear with few clouds. The radiation presented a value of 1079.95 W/²m between 11:10 and 12:28, the interval in which the test bench was in operation. The temperature and global radiation measurements are shown in Figure 13.

The temperatures of the three surfaces were saved every 2 minutes. The temperatures of the selective surfaces, glass cover, insulation and environmental were measured and their averages are shown in Table 2.

To calculate the absorptivity, emissivity and efficiency from the energy balance described in section 2.4, it was necessary to obtain data from the manufacturer of the commercial surface (MRTiNOX) and the glass used in the test bench. Manufacturer data are shown in Table 3.

To calculate the overall loss coefficient ($U_l$) using the energy balance, Equation 9, was used the manufacturer’s data together with the average radiation of 1020 W/m² in the range in which the test bench was in operation. To obtain the heat transfer coefficient (h), was applied the Equation 7 to the MRTiNOX surface data, knowing that the selective surface area is $A_p = 0.0008m^2$, the insulation thickness (L) is 5 cm and the thermal conductivity (K) is 0.04 W/mK. The results obtained for the coefficients are shown in Table 4.

For natural convection applications, the heat coefficient (h) can vary over a range of 5 to 30 W/m²K. In this way, the result found presents values within the expected range.

As the overall coefficient of heat loss, obtained by the energy balance, was $U_l = 19 W/m^2K$, thus, the absorptivity values of the selective surfaces produced can be obtained using

| Measured Elements | Average Temperature (°C) |
|-------------------|--------------------------|
| 25% $\text{Al}_2\text{O}_3 + 75\%$ CRFO | 69.4 |
| 50% $\text{Al}_2\text{O}_3 + 50\%$ CRFO | 68.9 |
| 75% $\text{Al}_2\text{O}_3 + 25\%$ CRFO | 70.1 |
| MRTiNOX | 70.2 |
| Glass Cover | 56.5 |
| Insulation | 39.3 |
| Environment | 39.2 |
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Equation 9. Using Equation 7, with the values of absorptivity, the emissivity values were obtained. The efficiency (f) of the surfaces can be evaluated dividing the absorptivity by the emissivity. For efficiency values greater than 18 (f > 18), the surface is considered highly selective. The results obtained are shown in Table 5.

The efficiency of the 25% Al₂O₃ + 75% CRFO surface was higher than that of the other surfaces, including the commercial MRTiNOX, because although it had lower absorptivity, it had a lower emissivity value, which increased its efficiency. The average surface temperature 25% Al₂O₃ + 75% CRFO, shown in Table 2, is very close to the average of the commercial surface MRTiNOX and there is an oscillation of these values in Figure 13, which is due to the proximity of the absorbance values and emittance of these two surfaces. The CRFO, 50% Al₂O₃ + 50% CRFO and 75% Al₂O₃ + 25% CRFO surfaces had lower efficiency and absorptivity values and higher emissivity values, but still considered high.

Figure 12. Micrograph of sample of 75% aluminum oxide – 25% CRFO powder, with magnification of (a) 10,000x and (b) 2000x.

Figure 13. Results of the field test.

| Table 3. Manufacturer’s data. |
|-----------------------------|
| Materials  | Properties  | Absorptivity (α) | Emissivity (ε) |
|------------|-------------|------------------|----------------|
| MRTiNOX   |             | 0.95             | 0.04           |
| Vidro     |             |                  | 0.87           |

Table 4. Results obtained for the coefficients.

| Coefficient                  | Results obtained |
|------------------------------|------------------|
| Overall loss coefficient (U) | 19               |
| Heat transfer coefficient (h) | \( \frac{W}{m^2K} \) | 27.9             |

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4. Conclusions

The selective surface obtained after performing the tests described in this study, showed satisfactory visual appearance, low thickness deposited with good adhesion to the metal substrate. It was also verified that the selective surfaces produced, based on the achieved results, obtained similar results to the commercial surface MRTiNOX, mainly for the absorptivity values, in the field test. It is observed that in the results obtained in the characterization by UV-VIS, the absorptivity showed high values in the visible light range, demonstrating an assertive characteristic for capturing solar energy. Therefore, the selective surfaces produced can be technically viable alternative in thermal solar-thermal collectors.

5. References

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Table 5. Absorptivity and emissivity of selective surfaces.

| Surface            | Absorptivity | Emissivity | Efficiency |
|--------------------|--------------|------------|------------|
| 25% Al2O3 + 75% CrFO | 0.935        | 0.043      | 21.74      |
| 50% Al2O3 + 50% CrFO | 0.935        | 0.048      | 19.48      |
| 75% Al2O3 + 25% CrFO | 0.94         | 0.0379     | 24.80      |
| MRTiNOX           | 0.95         | 0.04       | 23.75      |