Sulfidation of Ceramic-Based Coatings Deposited on Low-Alloyed Steel 16Mo3 Exposed at High Temperature

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Submitted: 25 February 2021 / Revised: 27 May 2021 / Accepted: 19 June 2021 / Published online: 12 August 2021

This study was carried out to evaluate a series of ceramic coatings in a sulfidation atmosphere containing 99% synthetic air and 1% H₂S (vol.%) under a flow rate of 50 mL/min. The study was carried out at 773.15 K for 336 hours (heating rate 278.15 K/min). Chemically resistant glass enamels based on SiO₂-B₂O₃-TiO₂-N₂O compounds were deposited on the surface of 16Mo3 (16M) low-alloyed steel. Kinetic data were recorded periodically every 168 hours; macro- and microanalyses using scanning electron microscopy (SEM), phase analyses using x-ray diffractometry (XRD) and chemical composition using energy-dispersive x-ray spectrometry along with EDS x-ray mappings were carried out to observe the degradation process of the sulfidized coatings. The results indicated that some of the ceramic coatings exposed in the harsh atmosphere at 773.15 K for 336 hours showed a high degree of protection.

Keywords 16Mo3 (16M) steel, ceramic coatings, high temperature, sulfidation

1. Introduction

Thermodynamic efficiency in the ultra-supercritical (USC) plants is achieved by increasing the operating temperatures and pressures, which lowers fuel consumption and subsequently the emissions (Ref 1). Furthermore, the European goal of decreased CO₂ emissions is a clear-cut requirement, which marks a historical turnover whereby Europe shall become fully neutral by 2050 (Ref 2). This can only be achieved by maximizing the efficiency of the plants and using biomass, which is considered carbon neutral. However, very little is known about the effects of the conditions in ultra-supercritical (USC) plants and the effects impurities from biomass may have on the corrosion resistance of the hot components in power plants (boiler walls, superheaters (SH), re-heaters (RH)). These components are already operating at the limit of their tolerance in less critical plants, and therefore, a radical step forward will be needed in materials technology to ensure the durability of components in such aggressive environments.

In particular, the problem is present when low-alloyed steels are used in power plant construction, even in USC technology. Low-alloyed steels such as 16Mo3 (16M) show very good thermal conductivity in contrast to austenitic steels; however, they show no corrosion resistance in aggressive atmospheres. In an air atmosphere, 16Mo3 (16M) steel with no coating protection indicates relatively high metal loss due to corrosion when a non-protective scale is formed. More severe corrosion degradation is always observed in sulfur-containing gas. Recent work published by Dudziak et al. (Ref 3) and past papers presented by Mrowec (Ref 4) and Ghaseni (Ref 5) demonstrate poor corrosion behavior of low-alloyed steels in sulfur gas. Degradation by sulfur results in the formation of a thick, porous, non-adherent sulfide scale with no protection. To secure low-alloyed steels in energy conversion systems and still use them due to high thermal conductivity, surface engineering as a coating design is a highly potential route. In the energy sector, the most popular types of coating are based on thermal spray technologies. There are many different types of thermal spray coating systems including plasma, wire, flame and kinetic; however, the most popularly used ones include high-velocity oxy-fuel (HVOF) or air, atmospheric plasma spray (APS) and others (Ref 6-10).

These two coating technologies are based on the same principles; a heat source melts the feedstock, where powder with an appropriate shape and diameter is used. The powder is accelerated at high temperature and impacts a substrate to form a coating with the desired thickness. In the APS process, plasma is produced via a direct current (DC) arc or radiofrequency (RF) discharge giving a temperature of over 8000 K with particle acceleration up to 20-500 m/s. These values depend on particle size and density, but higher speed is responsible for excellent deposition densities, bond strength and lower porosity of the deposited protective coating. The HVOF process was developed based on jet engine technology. The process relies on the oxygen–fuel ratio inside the combustion chamber where a highly pressurized mixture is formed. In contrast to the APS technique, in the HVOF process, particle velocity may reach 1000 m/s at a temperature of 3000 K. Similar to APS, density, bond strength and porosity directly depend on particle velocity, density and diameter.

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Nevertheless, in addition to remarkable corrosion resistance, these coating types possess some degree of porosity (dependent on the technology used) where ash can accumulate. Here, it initiates, accelerates and propagates the damage and destruction of the applied coating. Therefore, it is crucial to develop new protection systems for high-temperature applications, especially when a higher temperature is essential to promote a higher efficiency of the heat conversion system.

There is one group of materials that are highly resistant to high temperature. However, these materials are often brittle and may lead to a significant reduction in heat transfer. The ceramic-based coatings are new materials that can be used as protection for the boiler steels exposed to harsh atmospheres, where aggressive gases and ash are present. In contrast to thermal spray coatings, the ceramic glaze-based coatings are free from porosity; hence, there is no issue related to the development of local corrosion products within the coating structure. Furthermore, application of the coating from both sides (outer and inner) is possible. Heat transfer characteristics are the major bottleneck of the ceramic system; unusable for energy conversion system or a coating that is too thick may reduce it significantly, making such a coating unusable for energy conversion systems. Nevertheless, the applied coating should protect a low-alloyed steel against the formation of low-melting eutectics, especially when H2S and SO3 are present in contact with Na and K salts (Ref 11-13). In the energy sector, ceramic-based coatings are not very popular and there are a limited number of papers related to ceramic coatings. However, there are some solutions from the past, using a mixture of a special grade of calcined aluminum oxide and a conventional type of ground-coat frit with water to appropriate fineness on a steel surface. This mixture is then dried and fired according to well-known methods in the ceramic industry. Such a type of coating reached 70 µm. Progress in ceramic coatings for high-temperature applications has been extensively enhanced in the last century (Ref 14). In 1980, ceramic coatings were applied in adiabatic engines (Ref 15). Further, ceramic coatings were applied in gas turbine wings, pistons, cylinder linings, valves and piston crowns (Ref 16). Predominantly, nowadays, ceramic coatings are mainly used to protect Ni-based alloys against hot corrosion, oxidation and wear degradation (Ref 17, 18).

To get a better understanding of advanced ceramic materials and further develop them for a particular engineering application (i.e., energy sector, boiler protection), extensive research is essential for evaluating macro- and microstructural and oxidation resistance. In this research, new types of ceramic coatings based on SiO2-Na2O-B2O3-TiO2 oxide phase were applied on a low-alloyed Cr-Mo steel 16Mo3 (16M).

In total, seven different coating types were tested. In this study, uncoated steel was not tested. It is well known that sulfidation resistance of low-alloyed steels is poor (Ref 3). The materials were tested at high temperature in sulfidation conditions using a 99% synthetic air–1% H2S gas mixture (vol.%) to monitor the degradation mechanism. The following gas mixture was acknowledged based on the literature research, where 1% of H2S is the highest % share in the overall combustion gas mixture found in a coal-fired power plant. The test was carried out for 338 hours at 773.15 K. Kinetic data were recorded periodically every 168 hours. A full spectrum of research analyses using SEM/EDS/XRD were carried out to establish corrosion resistance of the exposed materials at high temperature.

### 2. Experimental Procedure

#### 2.1 Materials

The uncoated sample 16Mo3 (16M) was used as a reference sample and was delivered in as-received conditions. Prior to experimentation, the surface of the reference sample was ground using 600-grit SiC paper. The ceramic coatings were applied on the surface of the delivered materials, with no extra surface modification. The uncoated and the coated specimens had dimensions of 7-9 x 12 mm and a thickness of 4-5 mm in this work. The chemical composition according to Polish Norm: PN-75/H-84024 of the materials used in this work is listed in Table 1.

The enamel producing process took place in two stages. The raw materials were prepared by following the procedures given in Table 2, where individual ingredients were weighed, dry-mixed and rendered in alundum crucibles, and further, to finalize, the process was completed in an electric furnace using the following conditions:

- Set filling temperature-1473.15 K,
- melt pouring temperature-1633.15-1663.15 K,
- melting time-1-2 h.

The second stage involved the production of enamels on a laboratory scale. The oxide compositions of enamels in weight percentage are presented in Table 2. For better traceability of the coatings used in this work, individual symbols were used.

Additions such as five parts by weight of clay, NaNO2 in the amount of 0.1 part by weight, Na2[B4O5(OH)4]Æ8H2O and MgCO3 by 0.2 part by weight and deionized water 60 parts by weight were added. All the additives were calculated in relation to 100 parts of flux weight. To ensure traceability, the enamels were numbered 1 to 10. Some of the enamels contained a different concentration of flux: Sample F contains 80% flux, marked as A with 20% chromium trioxide CrO3. The sample marked G contains 70% flux marked as B with 30% chromium trioxide CrO3. In both cases, the percentage of other additives was calculated in relation to the total mass of the flux. The flux samples in the amount of 100 g were dry-milled (to accelerate the milling process) for 30 minutes in a high-speed planetary mill. The grinding process was controlled by a granular distribution analysis as shown in Table 3, and melting points are listed in Table 4.

The selected enamels were applied to a steel substrate 16M and fired at 1153.15 K for approximately 20 minutes. The enamel symbols are not continuous, there are no coatings denoted as FCHO2, FCHO4, FCHO5, FCHO9 and finally FCHO10—these samples during preparation for high-temperature testing at 773.15 K showed weak mechanical behavior, spallation occurred and cracks on the sample surfaces were observed; therefore, the samples were rejected from the further examination. Water suspensions with densities of around 1.60 ± 0.5 g/cm³ were spread by a spray method onto the steel surface and fired in the conditions given in Table 5.

Visual and microscopic assessments of the surfaces showed the presence of holes in the structure of the coated materials (coatings C-F) as presented in Fig. 1.
2.2 Sulfidation Test

Sulfidation testing was conducted in a flowing gas mixture containing synthetic air–H$_2$S. The content of H$_2$S in the gas mixture was 1 vol.%, with a flow rate of 50 mL/min and a heating rate of 278.15 K/min. The temperature (773.15 K) and pressure (1 bar) of the testing corresponded to the average service conditions in pulverized coal-fired power plants. The schematic of the experimental testing rig used for the high-temperature sulfidation testing is presented in Fig. 2.

The testing rig contains an electric tubular furnace (CARBOLITE®) with wire heating elements, the container

| Table 1  Chemical composition of 16Mo3 (16M) steel used as reference and substrate material in this work (wt.%) |
|---------------------------------|---|---|---|---|---|---|---|---|---|---|
| Grade | C      | Mn     | Si     | P max. | S max. | Cr max. | Ni max. | Cu max. | Mo     | Other |
|-------|--------|--------|--------|--------|--------|---------|---------|---------|--------|-------|
| 16Mo3 | 0.12-0.2 | 0.5-0.8 | 0.15-0.35 | 0.04   | 0.04   | 0.30    | 0.35    | 0.25    | 0.25-0.35 | Al. max 0.02 |

| Table 2  Enamel producing recipes (wt %) for ceramic coatings development used in this work |
|---------------------------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| Oxide Coating symbol | TCHO1 | TCHO3 | TCHO6 | TCHO7 | TCHO8 | TCHO11 | TCHO12 |
| SiO$_2$ | 54.13  | 51.72  | 56.80  | 55.76  | 63.17  | 45.11  | 39.79 |
| Al$_2$O$_3$ | -     | -     | 2.89   | -     | 2.22   | -     | -    |
| B$_2$O$_3$ | 18.10 | 15.79  | 7.22   | 2.02   | 8.32   | 15.08  | 12.14 |
| CaO     | -     | -     | 4.34   | 3.01   | 2.00   | -     | -    |
| Na$_2$O | 13.30 | 13.80  | 11.46  | 10.02  | 12.33  | 11.08  | 10.62 |
| K$_2$O  | -     | 1.50   | 9.18   | 3.25   | 1.91   | -     | 1.16 |
| ZnO     | -     | -     | -      | 5.04   | -      | 0.52   | 1.34 |
| Li$_2$O | 0.62  | 1.74   | -      | -      | 3.05   | -      | -    |
| TiO$_2$ | 6.14  | 7.53   | 5.69   | 1.99   | 3.05   | 5.11   | 5.79 |
| BaO     | 1.99  | 2.51   | -      | -      | -      | 1.66   | 1.93 |
| ZrO$_2$ | -     | -     | -      | 14.94  | -      | -      | -    |
| F       | 3.68  | 2.18   | -      | 1.99   | 1.85   | 3.07   | 1.67 |
| CoO     | 0.41  | 0.54   | 1.21   | 0.99   | 1.05   | 0.34   | 0.41 |
| MnO     | 1.02  | 1.61   | 0.52   | 0.43   | 0.45   | 0.85   | 1.24 |
| NiO     | 0.61  | 1.08   | 0.69   | 0.57   | 0.60   | 0.51   | 0.83 |
| Cr$_2$O$_3$ | -   | -      | -      | -      | 16.67  | 23.08 |

| Table 3  Parameters of the particle size distribution obtained by milling |
|---------------------------------|--------|--------|--------|--------|--------|
| Coating symbol | Enamel symbol | D(v:0.1), $\mu$m | D(v:0.5), $\mu$m | D(v:0.9), $\mu$m |
|-------|---------|------------------|------------------|------------------|
| A     | FCHO1   | 1.09             | 22.16            | 68.43            |
| B     | FCHO3   | 0.82             | 20.04            | 68.34            |
| C     | FCHO6   | 1.12             | 21.56            | 66.07            |
| D     | FCHO7   | 0.90             | 21.61            | 69.75            |
| E     | FCHO8   | 2.20             | 23.71            | 68.99            |
| F     | FCHO11  | 0.45             | 17.00            | 65.09            |
| G     | FCHO12  | 1.89             | 21.79            | 66.18            |

| Table 4  Determination of characteristic melting points |
|---------------------------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| Coating symbol | Enamel symbol | Deformation T, K | Barrel T, K | Sphere T, K | Hemisphere T, K | Flow T, K |
|-------|---------|-----------------|-------------|----------|----------------|----------|
| A     | FCHO1   | 933.15          | 963.15      | 983.15   | 1033.15        | 1073.15  |
| B     | FCHO3   | 973.15          | 986.15      | 998.15   | 1020.15        | 1043.15  |
| C     | FCHO6   | 983.15          | 1001.15     | 1029.15  | 1093.15        | 1130.15  |
| D     | FCHO7   | 985.15          | 998.15      | 1018.15  | 1073.15        | 1133.15  |
| E     | FCHO8   | 993.15          | 1008.15     | 1028.15  | 1078.15        | 1123.15  |
| F     | FCHO11  | 976.15          | 986.15      | 1003.15  | 1033.15        | 1151.15  |
| G     | FCHO12  | 1013.15         | 1033.15     | 1048.15  | 1073.15        | 1153.15  |

2.2 Sulfidation Test

Sulfidation testing was conducted in a flowing gas mixture containing synthetic air–H$_2$S. The content of H$_2$S in the gas mixture was 1 vol.%, with a flow rate of 50 mL/min and a heating rate of 278.15 K/min. The temperature (773.15 K) and pressure (1 bar) of the testing corresponded to the average service conditions in pulverized coal-fired power plants. The schematic of the experimental testing rig used for the high-temperature sulfidation testing is presented in Fig. 2.

The testing rig contains an electric tubular furnace (CARBOLITE®) with wire heating elements, the container
with reactive gases, piping for the gas supply equipped with the flow control system and the cooling unit. The container with the reactive gas (synthetic air 99%–H₂S 1%) was obtained from the gas supplier (Air Products). The gas flow through the furnace was controlled using a rotameter installed between the gas cylinder and the furnace. The flow rate was set to 50 Nml/min (Nml/min at 293.15 K, 1 bar—standard conditions). The gas mixture was constantly controlled via an online camera setup using gDMSS HD Lite software. The test samples were placed onto a double-deck alumina (99.5% of Al₂O₃) refractory holder, on a long edge, with a spacing between each sample before being inserted into a tube furnace and then heated within the furnace. The temperature in the furnace hot zone was monitored using a programmable controller from EURO-THERM. Before the high-temperature sulfidation tests, the furnace was carefully calibrated by testing three samples at once. For safety reasons, the reaction chamber (alumina tube) of the furnace was enclosed in a 316-L stainless steel vessel capped at both ends with 316-L stainless steel flanges locked by four screws. The exiting gas mixture was passed through a NaOH solution for H₂S neutralization and scrubbing and then directed to a ventilation system.

Sulfidation testing of the coated samples was conducted for 336 hours. The testing comprised two cycles of 168 hours/cycle. The samples were placed into the furnace at room temperature and were heated up to 773.15 K using a ramp rate of 278.15 K/min and held at this temperature for 168 hours; then, the samples were removed from the furnace, visually observed and pictured, and the first group of the samples (containing each material) were selected for further examination for the next 168 hours. Before sulfidation testing at high temperature, the coated samples were carefully measured by an electronic micrometer, later the samples were cleaned using an ultrasonic bath for 15 min at 313.15 K, and finally, the initial mass of each sample was recorded using a high-accuracy electronic balance Sartorius CPA225D. Post-exposure investigations were carried out using a single-lens DSLR camera Canon EOS 70D camera coupled with a Canon MP-E 65 mm f/2.8 macro lens and two scanning electron microscopes. Low-magnification images were captured using a Hitachi 3000

| Sample symbol | Enamel symbol | Max firing temperature and time |
|---------------|--------------|-------------------------------|
| A             | FCHO1        | 1123.15 K/3min                |
| B             | FCHO3        | 1123.15 K/3min                |
| C             | FCHO6        | 1153.15 K/5min                |
| D             | FCHO7        | 1123.15 K/4min                |
| E             | FCHO8        | 1123.15 K/4min                |
| F             | FCHO11       | 1123.15 K/3min                |
| G             | FCHO12       | 1123.15 K/3min                |

**Fig. 1** Macro structures of the samples prior sulphidation test at 773.15 K for 336 hours
Tabletop microscope operated in backscatter electron mode (BSE). Higher magnifications of the cross-sectioned samples were carried out using a SCIOS FEI dual-beam SEM. Chemical analyses of the exposed samples were conducted employing energy-dispersive x-ray spectrometry (EDS) from Bruker and EDAX. The XRD phase analyses of the corroded samples were carried out using PANalytical Empyrean, PiXcel3D machine with ICDD PDF-4+2020 database.

3. Results and Discussion

3.1 Macro-Observations

Figure 3 shows macro-images of the unexposed and the exposed ceramic coating in the H₂S gas mixture. As seen in the test, the uncoated 16M steel was not exposed. It is well known that a low-alloyed steel with as low as 1 wt.% Cr in sulfide-containing atmosphere shows no protection. Those types of steels predominantly form flaky sulfide scale with a tendency to spallate from the metallic core (Ref 3). Furthermore, two of the coatings (F (FCHO11) and G (FCHO12)) were withdrawn from the test, due to detachment of the ceramic layer from the metallic substrate (16M steel), and both coatings were formed based on a Cr₂O₃ phase. The other coatings showed a decent corrosion resistance in the sulfide atmosphere, and despite this, some of the coatings showed some porosity and hole formation during preparation. These holes and subsequent levels of porosity showed no big influence on the overall corrosion resistance of the applied ceramic coatings; however, the coatings with a higher concentration of holes and porosity indicated poorer sulfidation resistance (especially coatings C and D).

The macro-observations revealed that coatings A and B indicated a smooth, non-cracked, non-delaminated surface. Both coatings also indicated a higher concentration of SiO₂, B₂O₃, TiO₂ and Na₂O phases compared with the other coatings used in this work.

3.2 Kinetic

Ceramic coating kinetics are shown in Fig. 4. The results presented clearly show that exposure of ceramic coatings to a 99% synthetic air–1%H₂S (vol.%) gas mixture indicates a relatively low mass gain; three out of four coatings showed similar results, where mass gain after 336 hours at 773.15 K/min reached around 1.9 mg/cm². Coating D indicated over four times higher mass gain. As mentioned in this work previously, two coatings (coatings F and G) delaminated from the metallic substrate and were withdrawn from the test; there is no need to describe corrosion behavior of these coatings when delamination occurred during the laboratory exposure. Table 6 shows error calculations for the exposed systems in sulfidizing atmosphere. The calculation was done for a singular sample that was measured five times in a row. Based on the performed measurements, a standard deviation was evaluated as well as error value measurement. The calculations showed that mass gain kinetics were measured with a little error (lower than 5%).

3.3 XRD analyses

Figure 5, 6, 7, 8 and 9 shows XRD spectra of the ceramic coatings exposed in sulfidation conditions. In coatings A and B, the formation of Na₂SO₄ was predominantly responsible for the mass gain of the samples according to XRD spectra. In coating C, in addition to Na₂SO₄, CaSO₄ phase formation increased mass gain. The results are in good agreement with kinetics, and higher mass gain was attributed to the samples with a higher number of sulfide phases developed after 336 hours of sulfidation in 99% synthetic air–1%H₂S (vol.%) gas mixture. Additionally, the mass gain of the coatings is proportional to the sulfur content reacting with the reagent phases that were applied on the coating. The α-SiO₂ and TiO₂ were untouched in coatings A–C. However, the Na₂O...
phase in coatings A–C reacted with H$_2$S at high-temperature, forming Na$_2$SO$_4$; in contrast, the Li$_2$O phase, with their high activity oxygen atmosphere, reacted to create Li$_2$TiO$_3$ phase instead of reacting with H$_2$S. This finding shows that the formation of an Li$_2$TiO$_3$ phase is more favorable than the formation of Li-S-type phase or Ti-S-type phase. Further analyses of the XRD data show coating C with CaO forms a CaSO$_4$ phase. The highest mass gain was observed in coating D where the highest number of phases (sulfides, oxides) formed; based on the results, the following phases were developed: Na$_3$Fe(SO$_4$)$_3$, NaLiSO$_4$, Co$_2$O$_4$ and finally CaMn$_3$O$_8$. The findings indicate the formation of alkali-iron tri-sulfate Na$_3$Fe(SO$_4$)$_3$ and important observation since alkali-iron tri-sulfate is present in fireside corrosion atmospheres (Ref 6, 10). The mechanism of the formation of alkali-iron tri-sulfate phase is proposed in the following. In the first stage of the experiment, H$_2$S is decomposed into H$_2$ and S$_2$; then, S$_2$ reacts with O$_2$ from

Fig. 3 Macro images of the exposed coatings at 773.15 K for 336 hours in sulphidising atmosphere

Fig. 4 Kinetic data for the ceramic coatings exposed in 99% synthetic air–1%H$_2$S (vol.%) gas mixture
the synthetic air atmosphere to form SO$_2$, and finally, SO$_2$ reacts with O$_2$ according to the following reactions:

\[ \text{H}_2\text{S} \rightarrow \text{H}_2 + \text{S}_2 \quad (\text{Eq } 1) \]
\[ \text{S}_2 + \frac{1}{2}\text{O}_2 \rightarrow \text{SO}_2 \quad (\text{Eq } 2) \]
\[ \text{SO}_2 + \frac{1}{2}\text{O}_2 \rightarrow \text{SO}_3 \quad (\text{Eq } 3) \]

During high-temperature exposure at 773.15 K, the formed SO$_3$ gas reacted with Na$_2$O phase from the glaze. Simultaneously, the developed Fe oxide reacted with Na$_2$SO$_4$ to form the alkali-iron tri-sulfate (Na$_3$Fe(SO$_4$)$_3$):

\[ \text{Na}_2\text{O} + \text{SO}_3(g) \rightarrow \text{Na}_2\text{SO}_4 \quad (\text{Eq } 4) \]

Fe + $\frac{1}{2}$O$_2(g) \rightarrow$ FeO \hspace{1cm} (Eq 5)

FeO + 3Na$_2$SO$_4$ $\rightarrow$ 2(Na)$_3$Fe(SO$_4$)$_3$ + $\frac{1}{2}$O$_2$ \hspace{1cm} (Eq 6)

The formation of alkali-iron tri-sulfate is the principle reason for the degradation of the exposed steels with low Cr content. Corey et al. (Ref 19) found that alkali-iron tri-sulfate is stable when the concentration of SO$_3$ is at a specific level. Below this level, the tri-sulfate phase becomes unstable and decomposes. In this case, the concentration of SO$_3$ in flue gas reached a value of 250 ppm or higher. The formation of SO$_3$ kept the concentration above 250 ppm to form a stable alkali-iron tri-sulfate that was found by XRD analysis.

The other phases found by XRD analyses (NaLiSO$_4$, CaSO$_4$) also reacted with the S$_2$ derived from H$_2$S decompo-

| Sample | Standard deviation | Error value, % |
|--------|-------------------|---------------|
| A      | 0.010             | 1.28E+13      |
| 168    | 0.026             | 1.57E+00      |
| 336    | 0.086             | 5.00E+00      |
| B      | 0.006             | -9.04E+12     |
| 168    | 0.024             | 1.24E+00      |
| 336    | 0.086             | 4.70E+00      |
| C      | 0.007             | -1.807E+13    |
| 168    | 0.017             | 8.387E-01     |
| 336    | 0.012             | 4.167E-01     |
| D      | 0.005             | -1.46E+13     |
| 168    | 0.085             | 1.08E+00      |
| 336    | 0.020             | 2.47E-01      |
| E      | 0.005             | 1.44E+13      |
| 168    | 0.010             | 6.61E-01      |
| 336    | 0.015             | 9.44E-01      |

Fig. 5 XRD spectra of the coating A exposed in 99% synthetic air—1%H$_2$S (vol. %) gas mixture

Fig. 6 XRD spectra of the coating B exposed in 99% synthetic air—1%H$_2$S (vol. %) gas mixture

Fig. 7 XRD spectra of the coating C exposed in 99% synthetic air—1%H$_2$S (vol. %) gas mixture
The phase formation mechanism is shown in the following:

The formation of NaLiSO₄ and CaSO₄ phases originates in the first step from H₂S decomposition and a reaction with O₂ according to reactions Eq 1, 3 and 3. In the case of NaLiSO₄, it is strongly believed that firstly Na₂SO₄ is formed according to reaction Eq 4, due to the higher concentration of Na₂O than Li₂O in coating D. Further, Na₂SO₄ is combined with the Li₂O phase according to the following reaction:

$$2\text{Na}_2\text{SO}_4 + \text{Li}_2\text{O} \rightarrow 2\text{NaLiSO}_4 + \text{SO}_3(g) + 2\frac{1}{2}\text{O}_2(g) \quad (\text{Eq 7})$$

The formation of CaSO₄ is derived from the combination of CaO and SO₃:

$$\text{CaO} + \text{SO}_3(g) \rightarrow \text{CaSO}_4 + \frac{1}{2}\text{O}_2(g) \quad (\text{Eq 8})$$

The reaction of CaO with H₂S to produce CaSO₄ is rather impossible in the current conditions according to Allen et al. (Ref 20) where a direct reaction between CaO and H₂S formulates CaS and H₂O as follows:

$$\text{CaO} + \text{H}_2\text{S} \rightarrow \text{CaS} + \text{H}_2\text{O} \quad (\text{Eq 9})$$

However, in this work a CaS phase was not detected by XRD; therefore, reaction 8 showing the formation of CaSO₄ is valid.

In addition to sulfide formation, performed XRD analyses indicated the formation of mixed oxides such as Co₃O₄ and Ca₂Mn₃O₈ (coating D) and Li₂TiO₃ (coatings A and C). No new oxide phases were detected in coating B. The formation of these oxide phases is directly related to the presence of oxygen in the synthetic air. The formation of oxides, similar to the formation of sulfides, is responsible for the mass gain of the samples. The number of oxide formation is in good correlation with mass gain, and the higher the number of oxides formed, the higher the mass gain observed. The highest mass gain was presented in coating D, and this coating also developed more oxides than other coatings exposed in this work. Coating D developed Co₃O₄ and Ca₂Mn₃O₈ phases according to XRD findings. The formation of CaO₃O₄ results from oxidation of CoO according to the following reaction:

$$3\text{CoO} + \frac{3}{2}\text{O}_2(g) \rightarrow \text{Co}_3\text{O}_4 \quad (\text{Eq 10})$$

The formation of Ca₂Mn₃O₈ structure is postulated by the following reaction:

$$2\text{CaO} + 3\text{MnO} + \frac{3}{2}\text{O}_2(g) \rightarrow \text{Ca}_2\text{Mn}_3\text{O}_8 \quad (\text{Eq 11})$$

There are a very limited number of papers indicating the formation of Ca₂Mn₃O₈ phase. It is believed that this compound is formed due to the direct reaction between CoO and MnO in an oxygen-rich atmosphere. The coating developed during sulfidation test the following phases: Na₂SO₄ (similar to coatings A–C); in addition, XRD found CaTiF₆ and (KNa)AlSiO₄. The formation of CaTiF₆ is believed to originate from the direct reaction of CaO, TiO₂ and F:

$$\text{CaO} + \text{TiO}_2 + 6\text{F} \rightarrow \text{CaTiF}_6 + \frac{3}{2}\text{O}_2(g)$$

Based on XRD data, the coating developed during sulfidation test the following phases: Na₂SO₄ (similar to the coatings A–C) and CaTiF₆, K,Na)AlSiO₄. The formation of CaTiF₆ is believed to originate from the direct reaction between CaO, TiO₂ and F. Finally the formation of (K,Na)AlSiO₄ is a rather more complex reaction chain similar to the formation of alkali iron tri-sulphates, as described in the previous paper (Ref 21).

Because the performed XRD analyses were unable to provide concentration of phases on XRD diffractogram directly, the Rietveld analyses were carried out. The concentration of the phases developed during sulfidation test is shown in Table 7. The analyses were performed based on PDF-4+ 2020 database. Table 7 shows model fit parameters (quality): Rp [%], wRp [%] and x². The lower the values of those parameters, the analyses have a better quality of the model fit. Furthermore, Table 7 shows concentration of crystalline phases for the individual sample and unit cell parameters. The performed analyses clearly showed that the highest concentration of Na₂SO₄ was indicated in coating E (57 %), coating B (39 %), coating C (26 %) and coating A (24 %), but in coating D no Na₂SO₄ was found according to the report. The highest concentration of the
formation of TiO$_2$ was observed in coatings B and A, respectively, and the other coatings show no TiO$_2$ (anatase) formation. In coating D, concentration of TiO$_2$ (anatase) reached 0.9%. Furthermore, SiO$_2$ (cristobalite) was observed only in coating A (15%) and coating E (38%). The formation of alkali-iron trisulfate Na$_3$Fe(SO$_4$)$_3$ was observed in coating D (57%), and this coating achieved the highest mass gain of the exposed materials in this work. Finally, the analyses found α-Fe concentration in coating B (26%), in the other coatings, α-Fe concentration was much lower, in coating C, only CaSO$_4$ was indicated, and no CaSO$_4$ was discovered in the other exposed coatings in this study. To summarize, the most commonly known phases are as follows: Na$_2$SO$_4$: coatings: A, B, C and E, TiO$_2$ (anatase): coatings: A, B (D traces) and SiO$_2$: coatings: A and E.

### 3.4 Microanalyses

#### 3.4.1 Cross-sectioned SEM Images of the Uncoated 16Mo3 (16M) Steel.

Figure 10 shows cross-sectional images of the uncoated 16Mo3 (16M) steel exposed in sulfidation conditions at 773.15 K for 100 hours. The figures were captured during different researches carried out in the past (Ref 22). As shown, the uncoated steel shows very poor corrosion resistance and a thick scale formed where plate- and needle-like structures rich in Fe-S phase are developed. The thickness of the sulfide scale reached around 500 μm (measured from the top of the sulfide scale to the substrate interface). Based on EDS analyses (not shown here) on cross-sectioned samples performed after exposure at 773.15 K, it was found that the outermost part of the sulfide scale consisted of a high concentration of S reaching 70 and 30 wt.% Fe. The middle part of the formed scale possessed high concentrations of S equivalent to 35 wt.% and Fe concentrations reaching 65 wt.%, indicating the formation of a non-stoichiometric Fe$_{1-x}$S phase. According to the findings

### Table 7 Rietveld analyses on the exposed ceramic coating

| Sample code | A   | B   | C   | D   | E   |
|-------------|-----|-----|-----|-----|-----|
| R$_p$ [%]   | 1.76| 1.76| 1.89| 2.66| 1.94|
| wR$_p$ [%]  | 2.37| 2.35| 2.7 | 4.23| 2.6 |
| $\chi^2$    | 2   | 2.39| 2.78| 11.2| 2.54|
| TiO$_2$ anatase share [%] | 26 ± 1 | 30 ± 2 | - | 0.9 ± 0.5 | - |
| a=b [Å]     | 3.721±0.001 | 3.74 ± 0.002 | - | 3.785 ± 0.003 | - |
| c [Å]       | 9.87 ± 0.02 | 9.84 ± 0.02 | - | 9.48 ± 0.02 | - |
| Co$_2$O$_3$ share [%] | 15 ± 1 | 14 ± 1 | - | 11 ± 1 | - |
| a=b=c [Å]   | 8.080 ± 0.001 | - | - | - | - |
| LiTiO$_3$ share [%] | 18 ± 1 | - | - | - | - |
| a=b=c [Å]   | 4.116 ± 0.001 | - | - | - | - |
| SiO$_2$ cristobalite share [%] | 15 ± 1 | 11 ± 1 | - | 11 ± 1 | - |
| a=b [Å]     | 5.025 ± 0.005 | - | - | - | - |
| c [Å]       | 7 ± 0.01 | - | - | - | - |
| α-Fe share [%] | 2.2 ± 0.4 | 26 ± 3 | 4.9 ± 0.3 | - | - |
| a=b=c [Å]   | 2.948 ± 0.001 | 2.952 ± 0.005 | 2.954 ± 0.001 | - | - |
| α-SiO$_2$ share [%] | 15 ± 1 | 5 ± 1 | - | 38 ± 1 | - |
| a=b [Å]     | 4.918 ± 0.002 | 4.915 ± 0.005 | - | 4.926 ± 0.001 | - |
| c [Å]       | 5.401 ± 0.003 | 5.45 ± 0.008 | - | 5.404 ± 0.002 | - |
| Na$_2$SO$_4$ share [%] | 24 ± 1 | 39 ± 2 | 26 ± 1 | - | 57 ± 1 |
| a [Å]       | 5.608 ± 0.004 | 5.615 ± 0.004 | 5.596 ± 0.004 | 5.611 ± 0.002 | - |
| b [Å]       | 8.969 ± 0.006 | 8.963 ± 0.005 | 8.991 ± 0.005 | 8.956 ± 0.003 | - |
| c [Å]       | 6.949 ± 0.002 | 6.977 ± 0.003 | 6.963 ± 0.002 | 6.967 ± 0.003 | - |
| CaSO$_4$ anhydrite share [%] | 54 ± 2 | - | - | - | - |
| a [Å]       | - | - | 6.969 ± 0.003 | - | - |
| b [Å]       | - | - | 6.194 ± 0.003 | - | - |
| c [Å]       | - | - | 6.989 ± 0.003 | - | - |
| CaH$_4$B$_2$ share [%] | 11 ± 2 | - | - | - | - |
| a [Å]       | - | - | - | 7.54 ± 0.01 | - |
| b [Å]       | - | - | - | 13.12 ± 0.02 | - |
| c [Å]       | - | - | - | 8.39 ± 0.01 | - |
| NaLiSO$_4$ share [%] | 21 ± 1 | - | - | 7.621 ± 0.002 | - |
| a=b [Å]     | - | - | - | 9.855 ± 0.005 | - |
| c [Å]       | - | - | - | - | - |
| Na$_3$Fe(SO$_4$)$_3$ share [%] | 57 ± 2 | - | - | 15.582 ± 0.002 | - |
| a=b [Å]     | - | - | - | 8.679 ± 0.002 | - |
| c [Å]       | - | - | - | - | - |
| CaTiF$_3$ share [%] | 5.7 ± 0.4 | - | - | - | - |
| a=b [Å]     | - | - | - | 5.764 ± 0.002 | - |
| c [Å]       | - | - | - | 14.19 ± 0.01 | - |
presented by Mrowec (Ref 23) in the past, a lower concentration of Cr than 4 wt.% in the exposed steel was identified, revealing the sulfidation rate as for pure iron. In sulfide-rich scale, the growth depends on the diffusion of Fe$^{2+}$/Fe$^{3+}$ ions from the metal matrix. Furthermore, when low-alloyed steel with up to 4 wt.% Cr such as 16Mo3 (16M) is exposed to a sulfur-rich atmosphere, S-rich phases are likely to develop such as FeS. The FeS phase possesses some degree of non-stoichiometry in the crystal lattice according to work performed by Mrowec (Ref 19), Gesmundo et al. (Ref 24) and McCammon (Ref 25); thus, the real formula of FeS should be written as Fe$_{1-x}$S, where $x = 0-0.2$ (Ref 26). The phase also possesses p-type semiconductor properties, where holes are the majority carriers and electrons are the minority carriers. Defects in Fe$_{1-x}$S develop due to the absorption of sulfur from the gaseous phase onto the Fe$_{1-x}$S crystal structure; furthermore, diffusion of iron cations from the metal matrix occurs via chemisorption with S$^{2-}$ anions.

Fig. 10 Cross sectioned images of the uncoated 16Mo3 (16M) steel exposed in sulphidation test conditions at 773.15 K for 100 hours

Fig. 11 Cross sectioned images of the coated 16Mo3 (16M) steel with different coatings exposed in 99% synthetic air and 1% H$_2$S (% vol.) conditions at 773.15 K for 336 hours
3.4.2 Cross-Sectioned SEM Images of the Coated 16Mo3 (16M) Steel. Figure 11(a-e) shows cross-sectioned SEM images of the coated 16Mo3 (16M) steel, and the steel was coated using different coatings to reduce sulfidation degradation. The images present only samples: A, B, C, D and E. Samples F and G were withdrawn from the test due to spallation of the coating from the substrate.

The present results clearly indicate a high degree of protectiveness of the ceramic coatings deposited on 16Mo3 (16M) steel. Among the exposed coatings, coatings A and B showed high resistance against sulfidation, where no cracks perpendicular or horizontal to the substrate appeared. The two coatings showed the lowest mass gains recorded; on the other hand, coating E despite low mass gain showed horizontal cracks nearby substrate interface. It is believed that cracks observed in cross-sectional image were derived from cutting process. Similar cracks were observed in coating C. Such observations lead to conclusion that those two types of chemical composition in the coatings are prone to hydrogen embrittlement; however, no analyses in this direction were carried out. The highest degree of cracks and degradation of the coating was observed in coating D, and this type of coating showed the highest mass gain related to the coated material. The results presented in cross-sectional captures are in good correlation with mass gain data presented earlier in this work.

Furthermore, the coating deposited on the 16Mo3 (16M) steels showed a high degree of non-reactivity with the highly aggressive 99% synthetic air and 1% H₂S (vol.%) gas mixture at 773.15 K. The results clearly indicated no corrosion products on the external coating surface exposed to harsh conditions. All three coatings showed under cross-sectioned x-ray mapping investigations a concentration of S equivalent to 2 (wt.%). The frit composition contained a mixture of a special grade of calcined aluminum oxide and a conventional type of ground-coat frit with water to appropriate fineness on a steel surface, and drying and firing led to high degree of protectiveness in comparison with the uncoated steel. In general, frit-based coatings exposed to harsh conditions with a high content of SiO₂ showed a high degree of non-reactivity as mentioned earlier. Nevertheless, according to the previous work published by Swaroop (Ref 27), when a coating is rich in SiO₂ in a highly concentrated H₂S system (99% synthetic air and 1% H₂S (vol.%)) as in this work, silica glass tends to lose silicon, by the formation of SiO at low O₂ partial pressure. The loss of Si may act by the direct dissociation of SiO₂ by reaction with silicon in the disilicide. Furthermore, this process may accelerate highly, because of H₂ and water vapor in the system. The presence of H₂ in contrast to S₂ in the atmosphere is the main issue, and as S₂ identified shows no impact on coating degradation, there are no thick corrosion products or coating cracks, where S₂ could ingress and react with the substrate. On the other hand, due to a high ability for penetration, H₂ can easily ingress into the coating structure and lead to brittleness of the coating structure, and this phenomenon is called hydrogen embrittlement. It is believed that this process was found in the coatings presented in Fig. 11(c, d). Hydrogen embrittlement is a well-known phenomenon. This process leads to a reduction in the ductility of the structure and load bearing capacity that finally provokes cracking, delamination and serious degradation of the coating system. Unfortunately, a detailed process is not well understood and still there is still ongoing experimentation with regard to the exact mechanisms. In short, the process is governed by the diffusion of H₂ throughout grain boundaries of the structured...
materials. (In steels, the diffused H$_2$ reacts with C producing CH$_4$.) The gas is not mobile and collects in small voids along the grain boundaries where it builds up enormous pressures that initiate cracks in the structure or lead to the formation of brittle phases that initiate crack propagation under external loading (cutting process). Similar observations could be detected in SiO$_2$-based glassy coatings exposed to H$_2$-rich atmosphere (Ref 28). Since H$_2$ possesses the smallest radius of atoms, a high number of defects in glassy materials provide easy access for H$_2$ atoms into the inner part of the coating. On the other hand, no corrosion products related to S$_2$ were found in this study, suggesting an impermeable character of the developed coating for S$_2$ atoms. Similar results were achieved by Kurkowa et al. (Ref 29), indicating CrSi$_2$–Ni and CoNiCrAlY–Si alloys in atmospheres containing NaCl and Na$_2$SO$_4$ showing a high degree of protection against S$_2$ inward diffusion due to SiO$_2$ scale formation. The SiO$_2$ scale plays a significant role in suppressing the inward diffusion of sulfur, resulting in the remarkable suppression of the formation of sulfide at the scale/substrate interface and inside of the substrate. The work was conducted on alloys able to individually form SiO$_2$; however, SiO$_2$ that possesses such properties opens new possibilities for the alloys structures that may be used in atmospheres containing H$_2$S.

3.4.3 EDS Mapping of the Exposed Samples. Figure 12(a–c) presents the EDS x-ray mapping performed on the best performing samples in the test carried out in this study. In general, the coatings analyzed under EDS x-ray mapping system showed a very similar concentration of the elements. The coatings mainly consisted of Si and O. The base material indicated the highest concentration of Fe and Cr was not detected due to a very low concentration of this element within the metal matrix. The interface observed between the coating and the substrate was occupied by Mn. The element diffused from the metal matrix of 16Mo3 (16M material). As mentioned Cr was not detected, neither Mo nor S$_2$ was observed by carrying out investigation, suggesting a high resistance of the developed coating against S$_2$ diffusion from the S$_2$-rich atmosphere (H$_2$S atmosphere). In addition, S$_2$ was not observed on the surface of the exposed coatings.

4. Conclusions

The aim of this work was to investigate a series of ceramic coatings exposed to 99% synthetic air and 1% H$_2$S (vol.%) under a flow rate of 50 mL/min for 336 hours at 773.15 K. The sulfidation study showed that coatings based on SiO$_2$–B$_2$O$_3$–TiO$_2$–Na$_2$O compounds indicated an overall good sulfidation resistance, and some spallation was observed in F and G coatings. Horizontal cracks were observed in C, D and E coatings, most probably due to the H$_2$-rich atmosphere and residual H$_2$O from synthetic air. After 500 hours of testing, the coatings showed no S$_2$ inward diffusion or S$_2$ presence on the substrate–coating interface observed.

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