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Corrosion behaviour of nitrated ferritic stainless steels for use in solid oxide fuel cell devices

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ABSTRACT

Plasma nitriding was applied to ferritic stainless steel substrates to improve their performances as interconnects for solid oxide fuel cell devices. The samples underwent electrical conductivity test and SEM/EDS, TEM/EDS, environmental-SEM analyses. The first stages of corrosion were recorded in-situ with the e-SEM. Nitriding is effective in limiting the undesired chromium evaporation from the steel substrates and accelerates the corrosion kinetics, but its influence of the electrical conductivity is ambiguous. No intergranular corrosion is found in the steel substrate after long time operation. Nitriding helps commercially competitive porous coating to improve chromium retention properties of metal interconnects.

1. Introduction

Solid oxide fuel cells are highly efficient electrochemical devices able to convert chemical energy into heat and electricity [1], with electrical efficiency higher than 60 % [2]. In commercial applications, individual SOFCs are stacked together and separated via shaped metal sheet, the metal interconnect (MIC).

The MIC fulfills fundamental tasks: it mechanically sustains the cell stacking, collects the produced electrical current and keeps air and fuel separated. A SOFC system works at 700–900 °C (depending on the cell materials) for, for commercial viability, at least 40000 h. In these challenging conditions, steel corrosion is one of the main processes to control as it can cause both electrical and mechanical failure; thermally grown oxides (TGOs) are the least conductive layers in the system and the presence of brittle phases can lead to delamination.

Specific ferritic stainless steels such as Crofer 22 APU [3] and Sandvik Sanergy HT [4] have been developed for SOFC application. These steels contain a high level of chromium, to ensure fast and homogenous surface passivation, and reactive elements (RE) [5], to control the oxide growth rate. A chromium oxide or manganese-chromium oxide [6] layer guarantees the corrosion resistance of the alloy. However this solution has a drawback, as the chromium volatile species

[7] produced at high temperature and in humid atmosphere [8] poison the perovskite materials [9] at the air side of the cell, leading to blocking of the cell redox reaction with time [10].

Hence, protective coatings are introduced on the alloy surface to mitigate Cr evaporation [11], with MnCo-spinel coatings being currently the preferred solution. Denser coatings provide better barrier properties [12], therefore deposition methods like PVD and APS have delivered the best results. Notably, Co-based spinels deposited using PVD provide excellent electrical performance coupled with chromium evaporation barrier properties. Coupling of a specific alloy composition and protective coating fully satisfies the needs for long SOFC device lifetime, but the cost of this technology is too high for SOFC commercialization, considering that a MIC could account for up to 30 % of the stack manufacturing costs [13].

As a consequence, cheaper materials are considered: low Cr-content alloys such as AISI441/K41 have been introduced and low temperature deposition techniques like wet powder spraying are used. The result is an active research field where there is still space for alternative materials and methods. In this context, the present paper presents results obtained from MIC solutions containing nitrated steel substrates.

Nitriding of metal interconnects have already been explored in polymer exchange membrane (PEM) fuel cells to substitute graphite

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bipolar plates. Graphite interconnects have excellent chemical stability to withstand corrosion, but their high cost and fragility pushed the adoption of a different material [14,15] like austenitic substrates, gaining in machinability and lower cost. On the other hand, the PEMFC environment corrodes the metal causing an increase in electrical contact resistance or, in the case of harsh corrosion, the release of iron cations, which in turn is detrimental for the polymer membrane material. Nitriding the MIC surface for PEMFC is a common method to decrease this corrosion rate and simultaneously guarantee acceptable electrical contact. Nitrogen reacts with the Cr present in the alloy creating a CrN or Cr₂N protective film on top of the steel.

In the steels presented in this article, however, chromium nitride is not the protective coating, as on the steel surface a Co-Mn based spinel is deposited. Comparing the results obtained with PEMFC applications is therefore difficult.

Research about sensitization [16] in ferritic stainless steel (FSS) is closer to the application described in the present study. In the second part of the 80's Nickel price increased of ca. 500 % in less than one year, forcing steel makers to look for alternatives to austenitic stainless steels (ASS). In particular, automotive industries needed alloys for exhaust systems, able to withstand high temperature. FSS have good corrosion properties at high temperature but the mechanical properties are inferior to austenitic stainless steels and if welded they might suffer from intergranular corrosion (IGC): Cr thermodynamically prefers to react with C and N. These precipitates at the grain boundaries of the welded, heat-affected region will scavenge Cr, leading to IGC and material failure [17–19]. Nevertheless, since the operating conditions for the automotive industry are again different from those in SOFCs, this branch of investigation is not sufficiently representative either to describe the behaviour of the material considered in this work.

Hence, even though nitrided stainless steels are not new as such, the utilization of nitrided ferritic stainless steels in SOFC working conditions (700 °C) has not been studied yet. Nitriding as pre-processing for SOFC MIC interconnects is a new approach. This manuscript investigates if and how nitriding can improve the lifetime of metal interconnects. In particular it aims to understand how it influences the corrosion process.

2. Materials and experiments

Three ferritic stainless steels were investigated in this study: Crofer 22 H, Sandvik Sanergy HT and K41 (or AISI 441). Compositions are presented in Table 1.

Crofer 22 H and Sandvik Sanergy HT have been developed for SOFC application. Their Cr contents are high enough to withstand the corrosion process at high temperature. They are enriched in reactive elements such as Mn, Nb, and Mo. Niobium improves the steel creep resistance, promoting Laves phases [2]. Manganese improves the thermally grown oxide electrical conductivity. The drawback of this high alloying is cost, at least tenfold higher than commercial K41.

The steel coupons used in this study were supplied by SOLIDPower S.p.A. and Teer coating Ltd.

The surfaces of half of the stainless steel samples tested were treated using high power pulsed plasma nitriding (HPPPN) process at TeerCoatings Ltd. (UK), to evaluate the properties of microstructure and scale growth of nitrided steel substrates under high temperature oxidation. The steel plates (100 × 100 mm) were placed on a substrate

turntable in a cylindrical vacuum chamber that was maintained at a pressure of 1.33 Pa with a flow of Ar or Ar/N₂ mixed gas. For a nitriding process, the turntable with single fold rotation was biased with power up to 6.0 kW, –600 V of negative bias potential, frequency up to 350 kHz, and 1600 ns of pulse width. The nitriding process was repeated on the opposite face of the plates. The stainless steels completed with the nitriding process on both faces were then carefully laser-cut into small coupons. The same nitriding process was applied to real MIC components (dimension of about 100*200 mm). The nitriding depth confirmed by microstructural analysis was in the range of 3–30 μm. The nitrided samples did not undergo polishing process to save the nitrided layer.

Several of the small coupons were tested with the method described elsewhere [23] to investigate their electrical and chromium retention properties. The method is a modification of the 4-probe test,

with two coupons are assembled facing each other. One coupon is the nitrided steel with a coating on top, while the other one is made of palladium, in between there is a Cr gathering layer of (La,Sr,Co) perovskite. Knowing the applied current density and acquiring the voltage drop between the two coupons, the ohmic loss is obtained. The Cr retention property instead is measured as the amount of chromium inside the perovskite. It was of interest to compare these two characteristics with non-nitrided (standard) substrates.

A portion of the small samples was instead investigated through an in-situ oxidation process. In-situ oxidations were observed with a modified FEI Quanta 200 environmental scanning electron microscope (ESEM). One side of the thin sample was heated with a laser while the other side was scanned by the electron beam. The temperature was monitored with a thermocouple welded onto the observed surface. A mass flow controller adjusted an O₂ flow of 5Nml/min into the microscope chamber close to the sample. With a differential pumping system, this lead to a O₂ pressure of 30 Pa (0.3 mbar) in the chamber. The tracking of the area of interest was performed manually. The observation window width was 8.2 microns.

The heating of these samples was conducted as follows: (1) temperature increase in 30 s to the next step, (2) temperature stabilization during 6 min while recording 6 images. The targeted temperature steps were: 200 °C, 300 °C, 400 °C, 450 °C, 500 °C, 550 °C, 600 °C, 650 °C, 700 °C. At 700 °C, the scan speed was set to 88 s/image and this state was maintained for one hour. Then, temperature was maintained 10 more minutes to record micrographs of the zone of interest at different magnifications (65, 125, 250, 500, 1 000, 2 000, 5 000, 10 000, 30 000). The temperature was finally decreased to room temperature in one minute.

After such in-situ observation, lamellae were extracted from each sample with a focussed ion beam (FIB)-SEM and subsequently observed with a transmission electron microscope (TEM). This allowed the observation of the sub-surface microstructure from 5 to 10 μm under the surface. TEM-energy dispersive x-ray spectroscopy (EDS) mapping was conducted on each lamella to compare the diffusion of elements and the obtained microstructure. In order to obtain more information on the metallic phases, ASTAR analysis from Nanomega was used on the nitrided samples. Data acquisitions were conducted for an hour on a square area of 1 μm².

The scale thickness calculation was performed through a Matlab routine that considered images pretreated with Adobe Photoshop, in order to segment and insulate the scale layer.

Table 1
Elemental compositions of metal interconnect substrates (weight percentage).

	Fe	Cr	C	Mn	Si	Al	Mo	Nb	Ti	W
K41/AISI 441 [20]	Bal.	18	0.01	0.3	0.35	–	–	0.45	0.17	–
Sanergy HT [21]	Bal.	21.2	0.04	0.3	0.12	0.02	0.96	0.71	0.09	–
Crofer 22 H [22]	Bal.	20.0-24.0	0.0-0.03	0.0-0.80	0.1-0.60	0.0-0.10	–	0.2-1.0	0.02-0.20	1.0-3.0

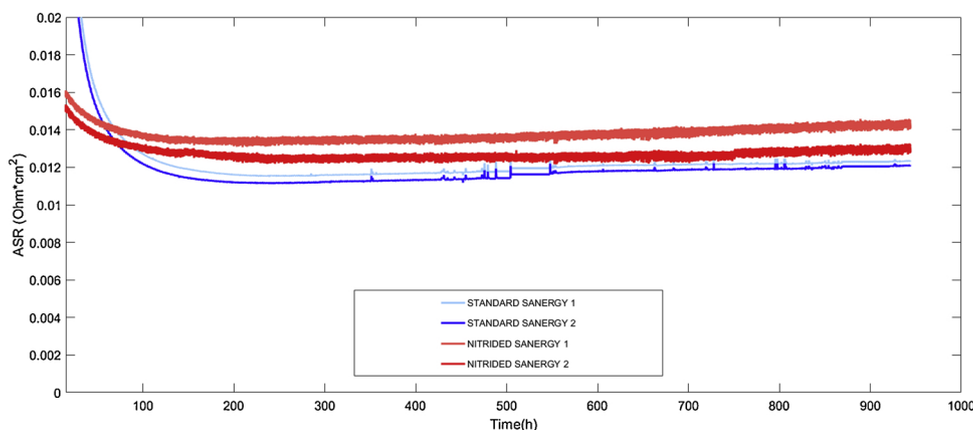


Fig. 1. Area specific resistance curves as a function of exposure time in air at 700 °C for physically vapour deposited MnCo_2O_4 coatings on Sandvik Sanergy HT stainless steel substrate. Each experiment had two repetitions for statistical reliability.

3. Results and discussion

3.1. Electrical resistance and Cr retention

The MICs have to collect the electricity produced by the SOFC, therefore low electrical resistance is desired. At the same time, the interconnect should not poison the cathode perovskite material by releasing chromium vapours [24]. The area specific resistance (ASR ($\Omega \cdot \text{cm}^2$)) is the parameter used to quantify the electrical performances; when divided by the scale thickness, it returns the material apparent resistivity. To quantify if and how much chromium migrated from the steel substrate to the perovskite, the material in contact with the nitrided samples was analysed by EDS.

Figs. 1 and 2 compare the results of ASR and Cr retention performances obtained from nitrided and standard (without nitriding) steel samples.

Fig. 1 presents the ASR time evolution (performed at 700 °C in air) for standard and nitrided Sandvik Sanergy samples coated with Fe-doped MCO. The ASR values after 1 000 h of exposure are slightly higher for the nitrided samples and the trends are similar. Not all the tested nitrided samples showed higher ASR values than the standard samples [25] (the whole set of data can be found in the link in the Appendix). It is necessary to evaluate ASR values after at least 1000 h of testing to ensure that all main temperature-driven phenomena stabilize. In nitrided substrates, this stabilization period appears shorter and less pronounced than that in standard substrates. This is visible as the

Table 2

Voltage degradation rates of complete interconnect + cell repeating units tested inside 6-cell SOFC stacks. A value of $< 0.5 \text{ \%kh}^{-1}$ is considered a threshold for a viable unit. WPS: wet powder spray, APS: atmospheric plasma spray, MCO: MnCo_2O_4 , MCF: $\text{MnCo}_{1.6}\text{Fe}_{0.4}\text{O}_4$. The RUs are listed according to their position in the stack: from the bottom to the top.

MIC	Coating Comp.	Steel substrate	Degradation (\%kh^{-1})
WPS MCO #1	MnCo_2O_4	K41/AISI441	0.72
NITRIDED APS MCO #1	MnCo_2O_4	Nitrided K41/AISI441	0.34
WPS MCO #2	MnCo_2O_4	K41/AISI441	0.67
APS MCF #1	$\text{MnCo}_{1.6}\text{Fe}_{0.4}\text{O}_4$	K41/AISI441	1.14
APS MCF #2	$\text{MnCo}_{1.6}\text{Fe}_{0.4}\text{O}_4$	K41/AISI441	0.64
NITRIDED APS MCF #2	$\text{MnCo}_{1.6}\text{Fe}_{0.4}\text{O}_4$	Nitrided K41/AISI441	0.45

plateau being reached after ≈ 100 h by nitrided substrates (Fig. 1).

Nitrided stainless steels were also used as the metal substrate in real SOFC interconnects tested in SOFC devices up to 10 000 h. Voltage degradation values in Table 2 indicate how much a certain single repeat unit contributes to the SOFC stack performance. The two single repeating units (interconnect + cell) containing the nitrided substrate behave better than those with a standard stainless steel substrate.

Regarding the chromium blocking function, nitrided samples constantly showed improved behaviour. This was particularly evident in

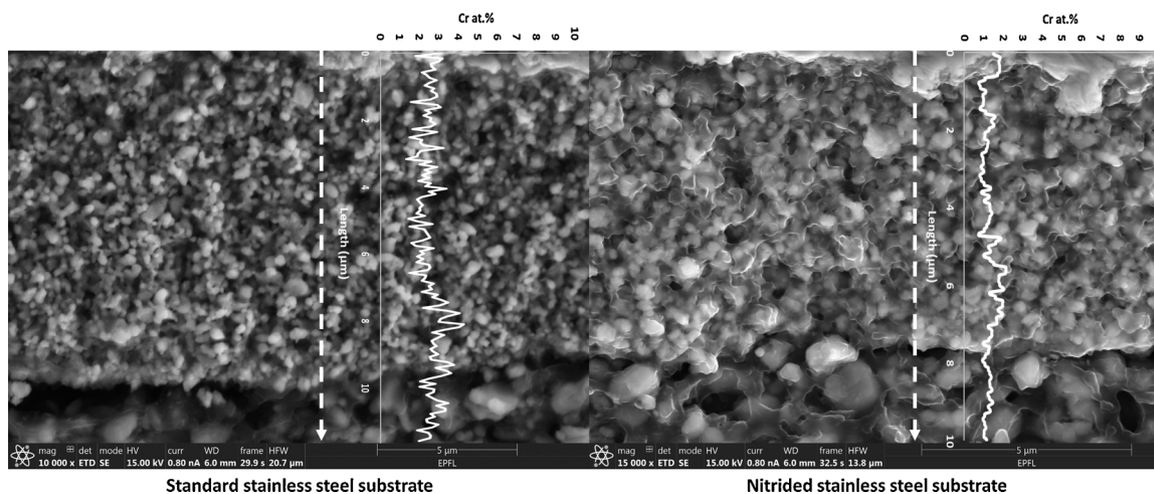


Fig. 2. EDS linescans taken on perovskite materials, simulating a SOFC cathode, in contact for 1 000 h in air at 700 °C with non-nitrided (left) and nitrided (right) samples.

the presence of porous spinel coatings. Fig. 2 compares the amount of chromium found in the LSC (cathode material) in contact for 1 000 h at 700 °C, with standard and nitrided stainless steel substrates coated with a porous spinel protective coating. These results come from small samples. In the nitrided sample, the average amount of chromium migrated into the cathode material was ≈ 1 at.%. For the standard substrate, it was 3 at.%. Since this type of coating process is one to two orders of magnitude cheaper than dense layer deposition techniques, this method to improve Cr retention is of technological interest.

The influence of nitrogen on the corrosion behaviour of the FSS MICs may account for the difference in the results of ASR and Cr evaporation.

Understanding the competition between chromium nitrides and oxides formation is necessary to predict the electrical and chemical behaviour of the steel substrates, particularly in the first stage of operation. The next paragraph presents the corrosion evolution of the steel substrates and identifies the species formed during aging.

3.2. Microstructure evolution

To help distinguish between nitrided and standard (non-nitrided) steel substrates in the microstructure analysis results, the former will be labelled with *N*, the latter with *St*.

It is known that chromia scale growth on stainless steel substrates containing nitrogen has faster kinetics [26]. Several experiments conducted in controlled atmospheres confirmed this: non-metals such as nitrogen can create ordered structures in the metal substrate with some of the alloying elements, chromium in particular [27]. This arrangement induces a higher presence of chromium at the surface, therefore when the material oxidizes, the generated chromium oxide nuclei are more numerous than with a standard substrate [28,29]. More nucleation sites lead to the formation of numerous oxide grains with small size. In addition, chromium ions diffuse faster through the grain boundaries which results in irregular scale growth on nitride samples, a porous containing structure

Fig. 3 presents the in-situ ESEM results on standard and nitrided stainless steel substrates (K41, Crofer 22 H and Sandvik Sanergy HT) after oxidation at different temperatures.

These image frames are taken from videos which can be found in the Appendix. The first column of Fig. 3 represents the moment when oxide grains become visible on the surface, around 300 °C for K41 and Crofer 22 H and around 400 °C for Sandvik Sanergy HT. These values have to be considered as average, as the heating of the samples may not have been homogenous. The third column presents the morphologies oxidised at 700 °C; this temperature was taken as the final point to compare it to results obtained in ASR tests [25]. An intermediate temperature value (≈ 550 °C, middle column) helps to follow the oxidation process.

There is a difference in the corrosion process between the standard and the nitrided substrates. With the latter, the scale on nitrided samples is porous and irregular, with the oxidation products resembling flakes, whereas in the standard substrates the scale is made of prismatic crystals, as expected [30].

This sponge-like structure on the nitrided substrates comes from the preferential growth of chromium oxide through grain boundaries [27].

The microstructures in Fig. 3 were qualitatively obtained from the top surface of nitrided and standard stainless steels. Micrometric lamella cross sections were subsequently extracted from the ESEM observed samples and studied by TEM. High-angle annular dark field images (HAADF) of the cross sections are shown in Fig. 4. The dotted lines indicate the oxide grown on the steel substrates. (The top part of the samples is a carbon layer deposited for protective purpose during the FIB extraction of the lamella and therefore not further considered.)

Looking at these cross sections, the oxide layer grown on nitrided substrates is confirmed to have experienced irregular growth. However, it is not possible to verify that the thicker regions match with the grain

boundaries because these are invisible. Preferential growth of the oxide layer through grain boundaries is visible for the standard Sandvik Sanergy HT steel substrate, where the scale has a maximum in line with the grain boundary.

Measurements of the scale thickness of these samples gave the following results, on the standard stainless steel substrates: 97 nm for Sanergy HT St., 167 nm for K41 St., 75.9 nm for Crofer 22 H St, and on the nitrided samples: 261 nm for Sanergy HT N, 175 nm K41 N, 255 nm for Crofer 22 H N. These thickness results confirm the faster scale growth on nitrided samples.

Besides being thicker, the grains of the scales on nitrided substrates are more irregular with different dimensions. Among the oxide grains in the nitrided substrates, some black spots are visible (Fig. 4 white arrows). These spots are residual of the space among the different grains at the beginning of the corrosion process. This pores confirms a lateral growth of the scale grains.

The absence of such pores in the standard substrates accounts for the faster kinetics of scale growth in the nitrided substrates. This being coherent with what is described above: the presence of nitrogen induces a chromium enrichment below the steel surface which in turn enhances the oxidation process occurring preferentially at the grain boundaries. From the grain boundaries then, the oxides expand laterally covering all the surface (cfr. videos in the appendix).

The elemental characterization of these lamellae (Fig. 5) indicates complex corrosion products for nitrided samples, with iron oxide as outer scale and chromium oxide adherent to the steel substrate. The nitrided steel substrate does not present inward corrosion, i.e. convex regions growing inside the metal. In contrast, a beginning of inward corrosion is visible in the K41St. substrate. The absence of inward scale growth is a relevant difference with the typical iron breakaway corrosion process.

EDS quantitative analysis on the ESEM-aged samples indicates different iron contents in the thermally grown oxides in Table 3.

The data indicate a more intense outward diffusion of iron on nitrided substrates. This result suggests that chromium in the initial oxidizing phase reacts preferentially with nitrogen, leaving the steel substrate unpassivated. Iron therefore could migrate outward and form an iron oxide scale as can be seen in Fig. 5. However, the absence of an enhanced inward corrosion suggests that this process took place for a short period only. In ASR-aged nitrided substrates as well (1 000 h, 700 °C), iron is found in the MCO coating, even though in smaller content (4–5 at.%), confirming the outward migration of Fe. Such iron outward diffusion was also observed in the protective coating deposited on nitrided interconnects for a SOFC stack that was tested for 10000 h, where the Fe percentage measured was around 25 at.%.

The other element found in the scale is manganese, which is expected and coherent with the common oxidation behaviour of steels in use for SOFCs [31]. Manganese in fact helps to mitigate chromium evaporation and scale growth; in addition, the mixed spinel of chromium and manganese has higher electrical conductivity than chromia alone [32].

The complete scale in the nitrided samples is thus made of a layer of chromium nitrides, followed by chromium-manganese oxide and finally Fe oxides on top.

Considering the metal substrate, the presence of nitride precipitates is the main difference between standard and nitrided substrates. The behaviour of nitrogen in the metal substrate is still under investigation in the literature, as it is influenced by temperature and alloying elements concentration [33,34]. Notably, it is important to understand how the nitrogen evolves and interacts with the lattice and with Fe and the alloying elements, in this case chromium. When nitrogen diffuses into the steel substrate it first reacts with the alloying element and then is dissolved in the matrix lattice without mechanical deformation. For both processes, a limit dictated by thermodynamic equilibrium exists. However, in reality, an excess of nitrogen is always present which mechanically stresses the metal substrate [35]. In addition, the ferritic

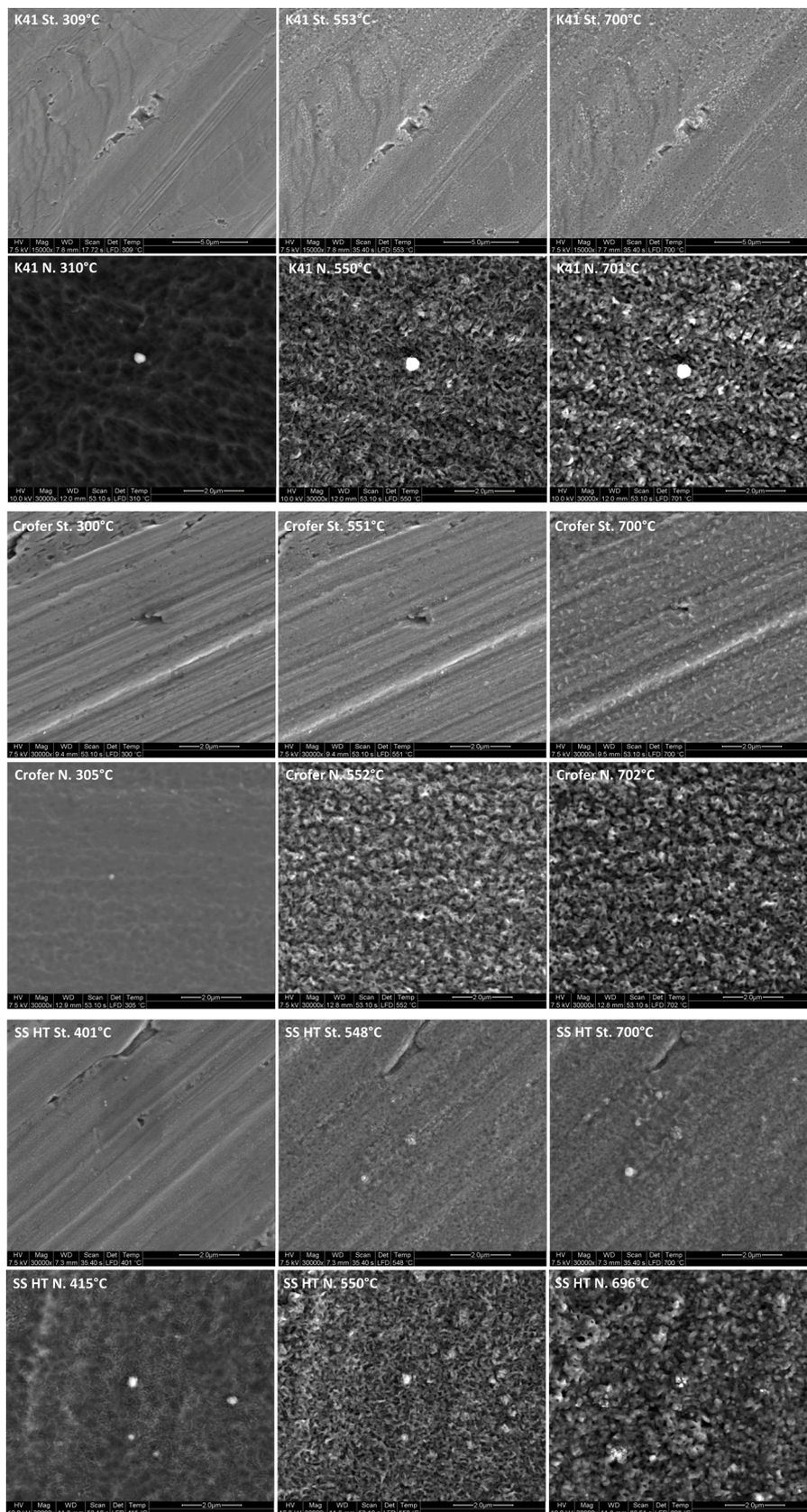


Fig. 3. Comparison of standard and nitrided FSS surfaces at different temperatures.

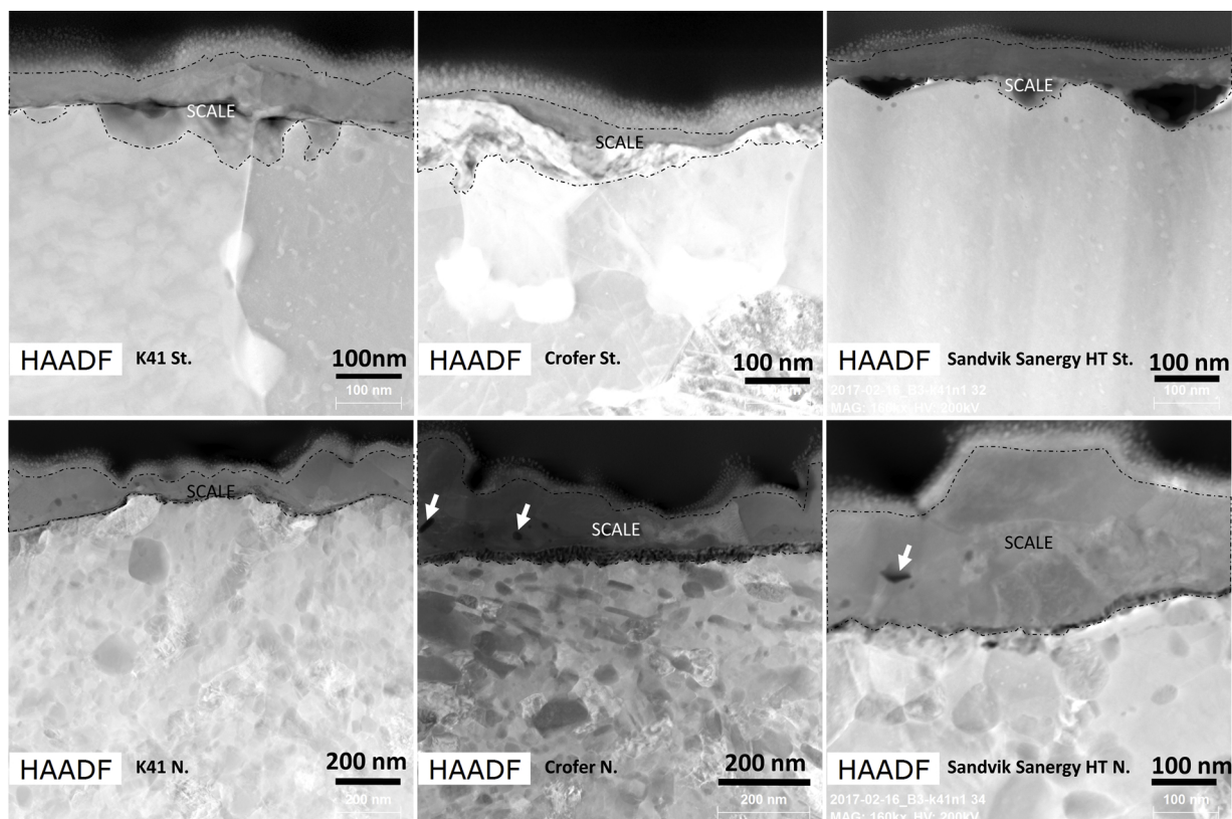


Fig. 4. TEM High-angle angular dark-field images (HAADF) of lamellae taken from the E-SEM observed samples.

phase and the nitrides have different crystal lattices. Ferrite α has a bcc structure, providing a coefficient of thermal expansion compatible with the SOFC components. Cr_2N instead has a hexagonal lattice causing a tension in the metal matrix. Finally, at temperatures higher than 800°C , nitrogen could stabilize the austenitic phase also with high chromium concentration [36].

Most of the studies encountered in literature are interested in describing the microstructural condition of the steel substrate immediately after the nitriding process. In this sense the samples tested for a few hours in the e-SEM are closer to this condition. As the main practical application is to modify the mechanical properties of the steel, there is no technological interest to understand the evolution of nitrated steel substrates for long exposure at high temperatures.

The consequence is that only a few studies describe aging of nitrated substrate in testing conditions close to those of this study (high chromium nitrated steels operated at high temperatures) [37,38]. Miyamoto et al. present a distinction between 'disk' and 'rod' shaped chromium nitride precipitates. Moving from the steel surface, i.e. closer to the nitriding source, towards the centre of the sample, the rod shape becomes prevalent. The density of the precipitates decreases as well further from the steel surface. The composition of precipitates tends to follow the morphology: CrN phase is typically disk shaped while Cr_2N is rod shaped. However, some rod shaped CrN is found as well: a composition change from Cr_2N to CrN is the proposed cause. In addition, Cr_2N grows when there is a minor presence of nitrogen; this is the reason why Cr_2N is encountered in greater depth in the steel substrate, close to the edge of the nitrogen diffusion region.

Fig. 6 shows two cross sections taken from ASR-aged substrates of nitrated AISI441/K41 and Sandvik Sanergy HT for 1000 h, in which both substrates carried a porous protective coating. Here Crofer 22 H is not reported, as its composition is similar to Sandvik Sanergy HT.

In the steel substrate cross sections, three regions with distinct morphologies can be observed. The layer immediately under the scale (belt 1) contains spherical precipitates and no visible segregation at the

grain boundaries. Below belt 1, there is a region with a high density of small and elongated precipitates inside the grains and at the grain boundaries (belt 2). Finally, in the core of the stainless steel, the biggest and widest particles are distributed (belt 3). Belt 2 and 3 are in a good agreement with what was previously described in the literature [37]. The precipitates closer to the surface are thin, elongated and numerous while those deeper into the steel centre are bigger and rod-shaped. Belt 1 conversely includes rounded precipitates. The average particle size in belt 1 increases with exposure time: in nitrated K41 stainless steel substrate after aging at 700°C for 1 h (ESEM samples), 1 000 h (ASR tested samples) and 10 000 h (SOFC stack samples), the particle sizes were estimated to be < 0.005 , 0.71 and $2.1 \mu\text{m}$ average respectively. In addition, the thickness of belt 1 region increases with the exposure time at high temperature: $19 \mu\text{m}$ at 1000 h and $51 \mu\text{m}$ at 10000 h. This suggests an Ostwald ripening process for belt 1.

The nitrated precipitates found in the e-SEM-aged lamellae indicate both rod and round shape as well in the region close to the steel surface. Fig. 7 also reveals that the nitrated precipitates are Cr_2N . From the TEM-EDS analyses the atomic percentage ratio between chromium and nitrogen is 2.8, 1.8 and 2.7 for K41, Crofer 22 H and SS HT substrates respectively. The grains that lay below the measured one might have influenced the measurement, inducing the higher amount of chromium.

Additionally, the cross sections of the interconnect components (10 000 h in stack operation) present similar morphology with that of ASR- and ESEM-samples (not shown here).

To better investigate which phases are present after aging, a TEM Astar identification was performed: SAED analysis was conducted on several grains, close to the surface, on standard and nitrated K41, Crofer 22 H and SSHT. Only ferritic iron was identified. The number of grains analysed depends on the quality of the TEM samples, which in turn depends on the FIB lamella extraction.

No austenitic phase could be identified in K41 N, nor in Crofer 22 H N. However, 3 grains out of 7 were identified as potential austenitic phase in nitrated SSHT. In order to obtain better statistical results,

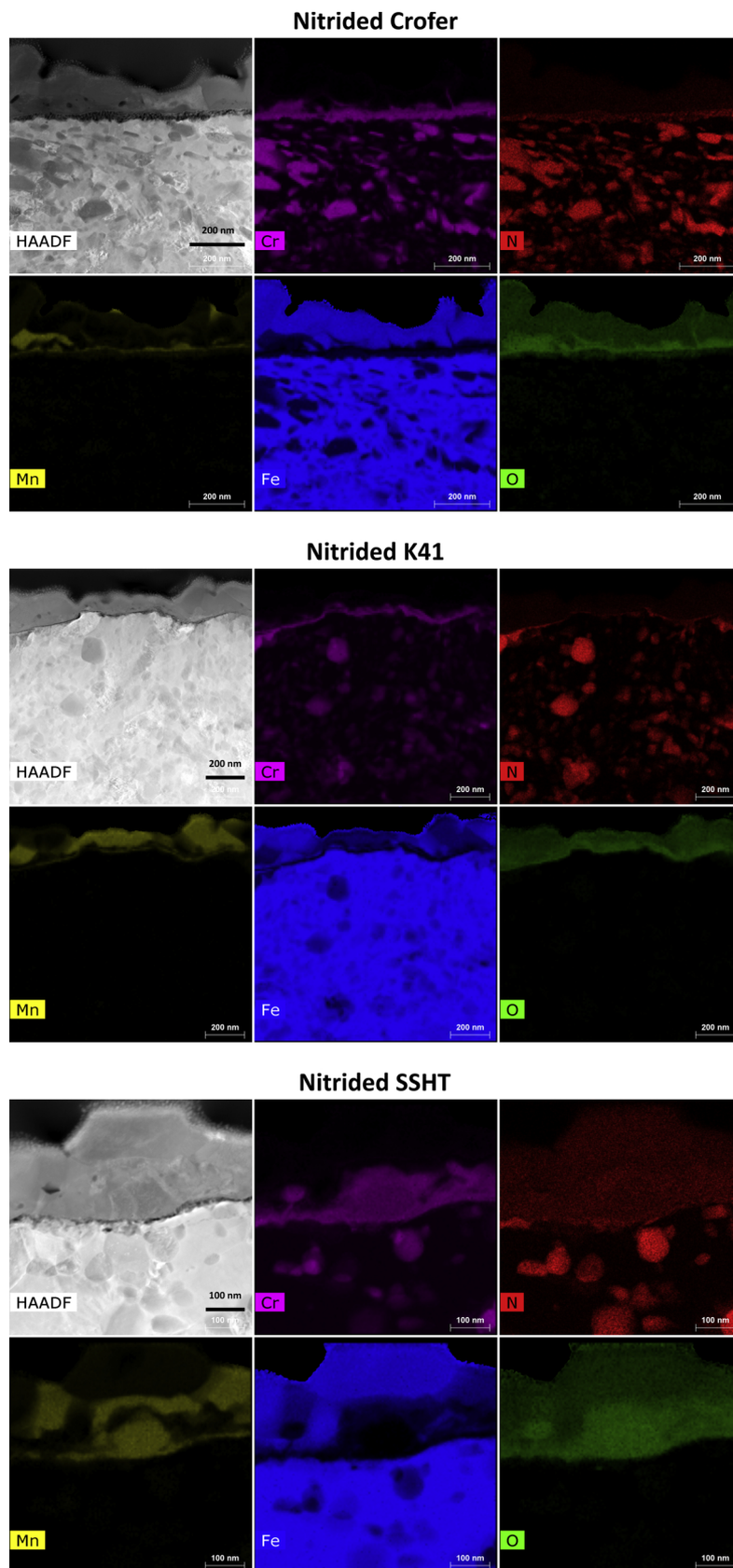


Fig. 5. EDS element maps of TEM-observed lamellae, taken from nitrided substrates tested in the ESEM.

Table 3

Alloying element content in the oxide layers grown on standard and nitrided ferritic stainless steel substrates after aging in environmental SEM (700 °C, 1 h). Values are in at.% and are taken from TEM-EDS investigations.

	K41 St.	K41 N.	Crofer St.	Crofer N.	SS HT St.	SS HT N.
Fe	11.2	33.2	6.4	35.7	18.3	30.3
Cr	38.9	7.3	38.9	10.4	48.1	14.4
Mn	9.0	14.9	4.1	4.5	1.7	8.9

ASTAR was conducted on $1 \times 1 \mu\text{m}^2$ of the nitrided sample lamella (estimation of more than 100 grains). Results show that less than 3 % of the surface was austenitic (Fig. 7).

The areas which are identified as austenite were also the areas with the lowest reliability of the obtained results (the ASTAR software indicates how much a phase identification is reliable) as seen on Fig. 7. The complexity of the system and the techniques make it difficult to clearly evaluate the presence of austenite. Indeed, even for simpler systems like binary Fe-Cr alloy [39], the diffraction interpretation is difficult. Automated crystal orientation mapping for TEM is also challenging [40] and unfortunately did not provide the desired answers for this study.

A known drawback of the presence of nitrogen and carbon is intergranular corrosion (IGC). At high temperatures, nitrogen interacts preferentially with chromium to create CrN or Cr₂N. In addition, this reaction is reported to take place principally at the grain boundaries because of the nitrogen segregation [41]. As a result, the metal substrate is depleted from chromium and passivation properties are deteriorated, with a preferential corrosion at the grain boundaries. In all the ASR-tested samples (1000 h) or real MIC operated samples (10 000 h) that were observed, no IGC was encountered, as the percentage of chromium contained in the steel substrate is high enough that it can compensate for the local loss due to reaction with nitrogen. The chromium cations diffuse from the bulk of the grain towards the depleted regions, restoring the passivation properties. This is visible also in the line scan result of Fig. 8 (taken in belt 1), with the green line being constant in the metal regions surrounding the Cr₂N particles. In addition, the high temperature helped to quickly redistribute chromium cations in the substrate.

Another possible issue for the steel substrate is the transformation from ferritic to austenitic phase within the nitrogen-rich region. This might be a problem for SOFC applications as austenitic stainless steel has a coefficient of thermal expansion higher than those of the cell components, which could potentially cause physical deformation on the cell up to breaking. According to the Fe-Cr-N phase diagram [42], high operating temperature (above 800 °C) coupled with high content of

nitrogen in the metal substrates stabilizes the austenitic phase. While this risk cannot be completely excluded, it is unlikely to have occurred according to the temperature the sample was exposed to.

Overall, both the absence of phase transformation from α to γ and the absence of intergranular corrosion – also for nitrided steels tested up to 10 000 h – confirm that the nitriding process is applicable to SOFC applications operated at intermediate temperature (700 °C).

4. Conclusion

Ferritic stainless steel substrates commonly used for SOFC applications have been nitrided and coated. Their compatibility as metal interconnects in SOFC devices was tested as coupons by evaluating their area specific resistance and Cr retention properties and compared to standard non-nitrided steel substrates.

Additional uncoated, nitrided and non-nitrided ferritic stainless steel samples underwent an in-situ oxidation process inside an environmental SEM. The superficial growth of the scale was recorded. TEM lamellae were extracted from the ESEM samples to examine the subsurface.

The nitriding process hardly affected the electrical conductivity (ASR) measurements of the steels (Fig. 1), and led to a shortening of the time to stabilise ASR values.

Nitriding resulted in an improvement of chromium retention, particularly on the steels with porous protective coatings. In the case of dense coatings, nitriding indicated no further improvement. Since porous protective layers are generally produced by simple wet chemical coating processes, at costs lower than that of physical vapour deposition or thermal plasma spraying, nitriding could provide an alternative solution for commercial application.

Scanning and transmission electron microscopy observations were performed on cross sections taken from standard and nitrided substrates aged for 1, 1 000 and 10 000 h, to examine the corrosion processes inside the nitrided samples.

Microscopy results confirmed faster oxidation kinetics in the nitrided samples. Nitrogen acts as a chromium scavenger, temporarily depleting the steel substrate and causing a limited iron breakaway corrosion, shown by the presence of Fe traces both in the coating and on top of the scale of uncoated samples. This Fe outward diffusion, however, took place for only a short period. The high content in chromium and the high testing temperature promptly recovered the passivation properties of the steel. In practice, this means that the initial acceleration of the corrosion kinetics is not harmful for the material, as no traces of intergranular corrosion were encountered in nitrided ferritic stainless steel substrates tested for 1 000 and 10 000 h.

The nitride precipitates in oxidizing atmosphere are identified

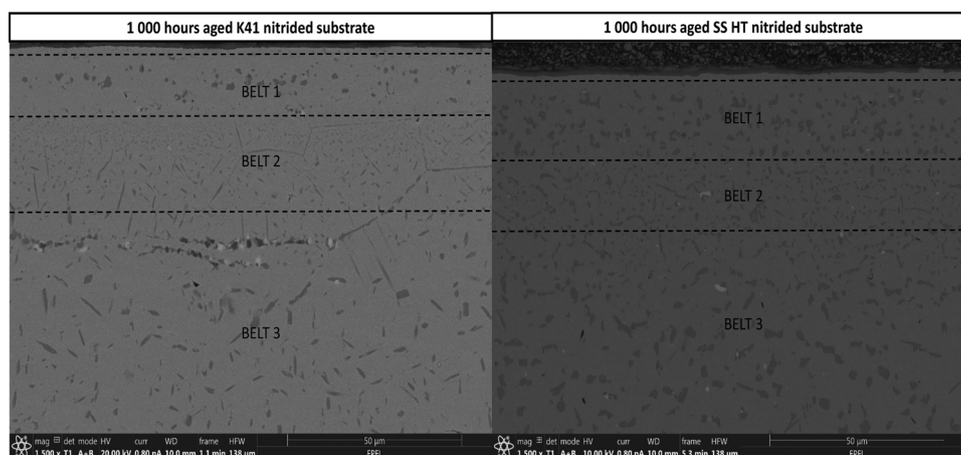


Fig. 6. SEM images of cross sections of ASR-tested samples (1000 h, 700C). On the left AISI441/K41, on the right Sandvik Sanergy HT. Each belt indicates a region where the nitride precipitates show a rather homogenous morphology.

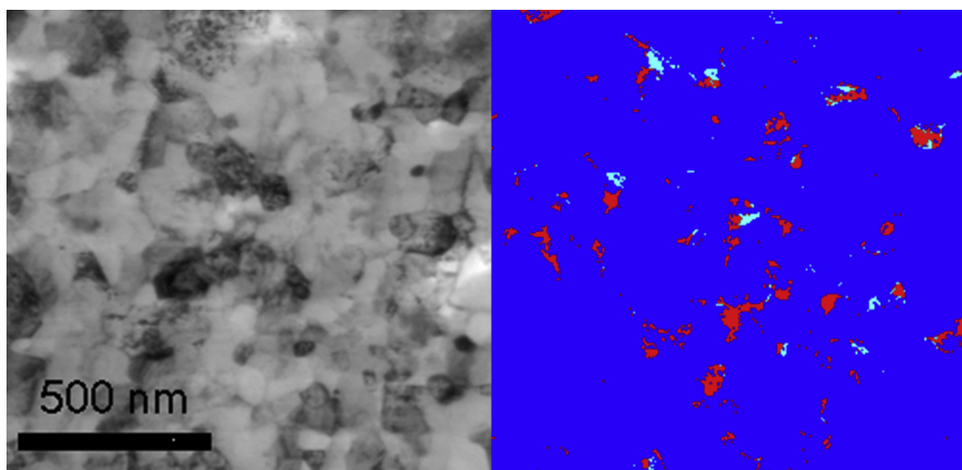


Fig. 7. Processed result from Astar analysis on Sandvik Sanergy HT substrate. Left image: virtual brightfield from the diffraction central spot. Right: phase map; dark blue is ferrite (96 %), light blue is austenite (0.7 %) and red is chromium nitride (3.3 %). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

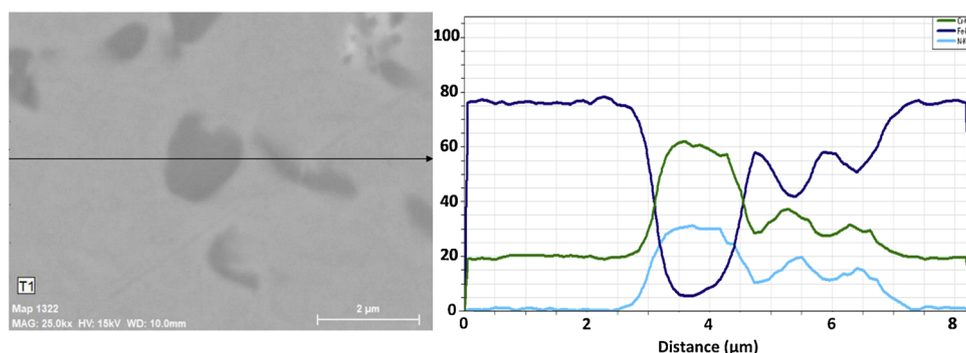


Fig. 8. Left: SEM backscattered electrons micrograph of nitrided Sandvik Sanergy HT substrate tested for 1000 h. On the right: quantitative analysis (at.%) of the linescan on the SEM picture.

mainly as Cr_2N and a phase transformation from ferritic to austenitic steel is considered to be unlikely.

Considering all aspects, nitriding is a method to improve the performance of low cost commercial ferritic stainless steels used as SOFC interconnects carrying porous coatings resulting from low cost deposition techniques such as wet powder spraying.

Data availability

All data used in this work, raw or processed, required to reproduce these results can be provided upon request by the corresponding author.

Declaration of Competing Interest

The authors declare that there are no conflicts of interest.

CRediT authorship contribution statement

Manuel Bianco: Conceptualization, Data curation, Resources, Formal analysis, Investigation, Methodology, Software, Validation, Visualization, Writing - original draft. **Stephane Poitel:** Data curation, Investigation, Writing - review & editing. **Jong-Eun Hong:** Writing - review & editing. **Shicai Yang:** Resources. **Zhu-Jun Wang:** Investigation. **Marc Willinger:** Investigation. **Robert Steinberger-Wilckens:** Funding acquisition, Project administration, Supervision. **Jan Van herle:** Writing - review & editing, Supervision.

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