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New insight on early oxidation stages of austenitic stainless steel from *in situ* XPS analysis on single-crystalline Fe-18Cr-13Ni

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Abstract

In situ X-ray photoelectron spectroscopy real-time measurements and angular-dependent high resolution core level analysis were used for the first time to investigate the Cr enrichment and oxide growth mechanisms on a model 304 austenitic stainless steel surface in the very initial stages of oxidation leading to pre-passivation. The oxidation kinetics was followed for increasing oxygen exposure and temperature, revealing an early nucleation regime (for exposure < 10 L) leading to the formation of a strongly Cr-enriched Cr^{3+}/Fe^{3+} mixed layer followed by an oxide growth regime where preferential iron oxidation takes over and mitigate the initial chromium enrichment.

Keywords: In situ X-ray photoelectron spectroscopy (XPS); stainless steel; single crystal surface; oxidation kinetics; Cr enrichment; growth mechanism

1. Introduction

Stainless steels (FeCr-based alloys) are widely used because of their high corrosion resistance resulting from the formation of a continuous and protective surface oxide layer, the passive film. Numerous surface analytical studies have shown that the passive film is only a few nanometers thick and markedly enriched in Cr(III) oxide/hydroxide species [1–13]. Recently, it has been suggested from nanometer scale studies on austenitic stainless steels that

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the Cr enrichment, which is the key factor for the corrosion resistance, may not be homogeneous in the oxide film [14, 15]. The pre-passivation mechanism by which the native oxide is initially formed most often in air may produce enrichment heterogeneities at the origin of the failure of the corrosion resistance by initiation of localized corrosion. Thus it appears essential to thoroughly investigate the initial stages of oxidation (i.e. pre-passivation) of the alloy surface under well-controlled temperature and exposure conditions, as well as the mechanisms governing the Cr enrichment.

Most studies [16–22] on the initial oxidation of pure iron and Fe-based alloys (mostly Fe-Cr alloys) have been carried out with a surface science approach in ultra-high vacuum (UHV) conditions, because it provides a maximal control of surface state and oxidation conditions which are key factors to surface reactivity and growth mechanism studies of ultra-thin oxide films. Polycrystalline alloys for practical applications were investigated despite the fact that single crystal surfaces enable a deeper insight to the surface composition and to the oxide formation process since limiting the complexity brought by dissimilar crystallographic orientations, grain boundaries, second phase particles and impurities. Only very few studies have been conducted on stainless steel single crystals [4, 15, 20, 23, 24].

X-ray Photoelectron Spectroscopy (XPS) is commonly applied to analyze surface chemical states and composition. Numerous XPS analyses provided chemical states references on Cr, Fe and Ni metals, as well as their various oxides [4, 5, 15, 23, 25–33]. For the alloy surface composition, it was concluded from previous works focused on the proportion of formed Cr oxides and Fe oxides that the formation of Cr oxides is thermodynamically favored while that of Fe oxides is kinetically controlled [34, 35]. The stratification of oxide films can be analyzed by angle-resolved XPS (ARXPS). For example, Castle *et al.* [31] have determined the in-depth distribution of oxides species according to a surface model determined from ARXPS measurements. However, there is almost no stratification analysis of surface region during the very first stages of oxidation.

Previous studies have also shown that oxidation temperature and oxygen exposure are two key factors for the oxidation rate, surface composition and oxides structure [16, 19, 20, 22, 36]. Most previous works were carried out at room temperature (RT) [16, 17, 24, 37] and around 300°C [20, 22, 37], the latter because of its industrial relevance. However, no oxidation study of Fe-Cr-Ni alloys was conducted systematically at different temperatures.

Regarding oxygen exposure, experiments were carried out at values larger than 1 000 L $(1 \text{ L} = 1.33 \times 10^{-6} \text{ mbar} \cdot \text{s})$, under oxygen pressure usually higher than 10^{-6} mbar. According to our kinetics data presented hereafter, this is far beyond the initial stages of oxidation, defined as the range of exposure for which the surface varies the most. The understanding of the mechanisms of oxide growth and Cr enrichment in the initial stages of oxidation is essential to understand the structure and properties of ultra-thin oxide films. However, to the best of our knowledge, there was no investigation of Fe-Cr-Ni alloy surfaces during and/or after very low oxygen exposure (< 100 L), neither of the mechanism of ultra-thin oxide film formation.

Nitrogen segregation has been observed in some studies performed on N-containing stainless steel [22, 38–40] and resulted in the formation of surface chromium nitride compounds. It was deduced by some authors that the co-segregation of chromium with nitrogen could increase the proportion of surface chromium oxides [41].

Here, we bring new insight on the oxidation kinetics and evolution of the surface composition and stratification during the initial oxidation of austenitic stainless steel. Nitrogen segregation is also discussed. The study was performed by *in situ* XPS on a (100)-oriented Fe-18Cr-13Ni single crystal (a model of the most common 304 grade of austenitic stainless steel) during the initial stages of oxygen exposure (< 100 L) at RT, 150°C and 250°C. The work is aimed at providing new comprehensive knowledge for the surface treatments of Fe-Cr-Ni alloys for improved resistance to initiation of localized corrosion of stainless steel.

2. Experimental

Firstly, the clean surface and nitrogen segregation were characterized. Then, the evolution of the oxygen uptake during oxygen exposure was monitored *in situ* to determine the oxidation kinetics. For that purpose, a novel specific XPS acquisition mode was adopted to record the real-time evolution of the photoelectron spectra during oxidation. In this way, the continuity of signal variation was achieved which strengthened greatly the kinetics monitoring. Afterwards, oxidation was conducted at exposures selected according to the kinetics data. The high resolution spectra were decomposed in order to determine the chemical states and their proportions. Based on angle-resolved data, oxide models were proposed to discuss the surface composition and stratification, as well as the thickness and formation mechanism of the oxide films and their Cr enrichment. A complementary Scanning Tunneling Microscopy (STM) study of the local topographical and structural alterations of the surface brought by initial oxidation will be reported separately [42].

The experiments were performed in a multi-chamber UHV system including a preparation chamber and two analysis chambers (XPS chamber and STM chamber), maintaining a base pressure $< 10^{-10}$ mbar. The preparation chamber is equipped with an Ar⁺ cold cathode ion gun for sputtering and resistive heating filament for annealing. Samples are heated from back side and temperature is controlled through a thermocouple attached on the manipulator in contact with the sample.

A (100)-oriented Fe-18Cr-13Ni single crystal with 99.999% purity was used. The sample was mechanically and electrochemically polished before introduction in the UHV system (procedure described elsewhere [23]). After introduction, the sample surface was cleaned by cycles of Ar⁺ ion sputtering (1 kV, 10 μ A, 10 minutes) and annealing (700°C, 10 minutes) in the preparation chamber. Sputtering and annealing cycles were applied between each oxidation experiment. The chemical and structural quality of the surface were controlled by XPS and low energy electron diffraction (LEED), respectively.

Gaseous oxygen (99.999% purity) was introduced *via* a leak valve directly connected to the preparation chamber or *via* an oxidation cell installed in the XPS analysis chamber. XPS analysis was conducted with the Argus spectrometer, using the monochromatic XM1000 Mkll Al K α X-ray source (1486.6 eV) both from Scienta Omicron. The Argus detector with 128 channels has a high spatial resolution and a high counting rate that allow to record electron spectra over short time period. The *in situ* XPS investigations were carried out using the Snapshot detection mode, recording a spectrum every 10 seconds in real-time monitoring during oxidation. Thousands of spectra were recorded to describe the variation of oxygen (or nitrogen) peak intensity during oxidation (or annealing). Most of oxidation treatments were carried out at RT, 150°C and 250°C with O_2 pressure $\leq 10^{-8}$ mbar. Such low oxygen pressure was kept in order to monitor the 0–100 L exposure range with good precision. The pressure in the chamber was continuously recorded and the exposure to O_2 was calculated from the integration of oxygen pressure as a function of oxidation time.

According to the *in situ* kinetics data, series of oxidations under very low oxygen exposures were carried out for detailed chemical analysis of the surface evolution. Survey and high resolution spectra of Fe 2p, Cr 2p, Ni 2p, O 1s, N 1s and C 1s core levels were recorded after each oxidation treatment, with constant pass energy of 50 eV and 20 eV, respectively. Both take-off angles of 45° and 90° were studied in order to determine the distribution of species along the direction normal to the surface. All obtained XPS spectra were referenced to the Fermi edge. Three exposures less than 10 L and nine exposures above were chosen for decomposition analyses. More than 400 high-resolution spectra were recorded and analyzed in this work. After exposure to about 300 L at 250°C, no more significant change in the substrate signal was observed.

Data analyses were performed using the CasaXPS software (Version 2.3.17) [43]. For reconstruction of the high resolution spectra, several constraints were considered to achieve accuracy and consistency.

The width of analyzed regions was carefully adjusted to ensure a suitable background contribution. For Fe and Cr, the 2p core levels were fitted by adjusting the intensity ratio between $2p_{3/2}$ and $2p_{1/2}$ components to values matching the theoretical ratio. For nickel, as there was no overlapping between the $2p_{3/2}$ (852.8 eV) and $2p_{1/2}$ (869.6 eV) peaks, only Ni $2p_{3/2}$ was considered. In order to improve the accuracy of the quantitative determination, only the $2p_{3/2}$ data were used to calculate the concentration and to establish the oxide model as $2p_{1/2}$ components have larger uncertainties.

Background subtraction methods can influence the quantitative values of Fe and Cr chemical states concentrations [25, 31, 44, 45]. It was found that the chemical states proportions extracted by using different background subtraction can vary by $\pm 5\%$ for iron oxide and $\pm 3\%$ for chromium oxide [25]. To minimize the effect of background subtraction on

components concentration, an adjustable Shirley background [46] was adopted for our peak fitting.

A Lorentzian asymmetric line shape, with the same asymptotic behavior as Doniach-Šunjić component [47], convoluted by a Gaussian shape was used for fitting the metallic component peaks. The indexes of asymmetry follow the order: metal > oxide > satellite \approx 0. A simple product of Gaussian and Lorentzian line shapes was used for the non-metallic components.

Binding energy (BE) and full width at half maximum (FWHM) are two important parameters affecting the reconstruction of the synthetic curve shape. Although metal compounds states have been largely studied by XPS [25–31, 35], BE and FWHM can be different according to the used sample and spectrometer. There are few data in the literature about the BE position of $2p_{1/2}$ core level peaks and shake-up satellite peaks. As for FWHM, it depends on the chemical environment. Normally, the FWHM values of the different peaks follow the order: satellite > oxide(nitride) > metal and $2p_{1/2} > 2p_{3/2}$.

Once the fitting parameters were determined, they were fixed for all the high-resolution spectra recorded at take-off angles of 45° and 90°. Only the relative peak intensities were treated as the variable for peak fitting.

3. Results and discussion

3.1. XPS fitting parameters

Table 1 compiles the fitting parameters determined from the XPS spectra reconstruction. They are consistent with typical values found in other works [25, 28, 29]. The chemical shifts of the different Fe oxides with respect to Fe⁰ metal are 1.7 eV, 2.6 eV and 3.5 eV for the components assigned to Fe²⁺ (in Fe₃O₄ matrix), Fe²⁺ (in FeO-like matrix) and Fe³⁺ (in Fe₂O₃ matrix), respectively. Spin orbit splitting is 13.1 eV between the Fe $2p_{3/2}$ and Fe $2p_{1/2}$ main peaks and 13.5 eV between the Fe $2p_{3/2}$ and Fe $2p_{1/2}$ shake-up satellite peaks, which is close to the values from recent studies [25, 26, 28, 31, 35]. BE separations between the Fe $2p_{3/2}$ main peaks and Fe $2p_{3/2}$ satellite peaks are 4.5 eV and 5.9 eV for Fe²⁺ and Fe³⁺, respectively.

Core level	State	BE ($\pm 0.1 \text{ eV}$)	FWHM (±0.1 eV)	Asymmetry index	
Fe 2p _{3/2}	Fe^{0}	706.8	0.8	0.59	
	$\mathrm{Fe}^{2+}(\mathrm{Fe}_{3}\mathrm{O}_{4})$	708.5	1.5	0.31	
	$\mathrm{Fe}^{2+}(\mathrm{FeO}\text{-like})$	709.4	2.1	0.31	
	$\mathrm{Fe}^{3+}(\mathrm{Fe}_2\mathrm{O}_3)$	710.3	3.1	0.42	
	Fe^{2+} satellite	713.0	4.0	0.13	
	${ m Fe}^{3+}$ satellite	716.2	4.3	0.21	
Cr 2p _{3/2}	Cr^{0}	573.8	1.1	0.61	
	$\rm Cr^{3+}(\rm CrN)$	575.6	2.0	0.19	
	$\mathrm{Cr}^{3+}(\mathrm{Cr}_2\mathrm{O}_3)$	576.5	2.0	0.46	
	Cr^{3+} satellite	588.3	3.5	0.00	
Ni 2p _{3/2}	Ni^{0}	852.8	0.9	0.31	
N 1s	$\operatorname{CrN}_{surface}$	396.6	1.1	0.00	
	CrN_{bulk}	397.2	0.7		
	N_{bulk}	397.8	0.7		
	$N_{minority}$	398.5 - 400.5	1.8 - 2.6		

Table 1 Fitting parameters obtained by reconstruction of the XPS core level spectra

This is 1-2 eV smaller than for pure Fe₃O₄ and Fe₂O₃ oxide samples like pointed out by Aronniemi *et al.* [25].

For chromium, the chemical shift is 1.8 eV for $Cr^{3+}(CrN)$ and 2.7 eV for Cr^{3+} oxide (Cr_2O_3) . These values are of the order reported in the literature [5, 25, 27, 28, 30]. Spinorbit splitting between the Cr $2p_{3/2}$ and Cr $2p_{1/2}$ peaks are 9.4 eV, 9.5 eV, 9.6 eV and 9.0 eV for Cr^0 , $Cr^{3+}(CrN)$, $Cr^{3+}(Cr_2O_3)$ and Cr shake-up satellite peaks, respectively. BE separation for Cr satellite peak is 14.5 eV which is 2–3 eV smaller than reported from a Cr 2p curve fitting study of a pure Cr_2O_3 powder sample [25].

The $2p_{3/2}$ and $2p_{1/2}$ doublet intensity ratios for Fe and Cr vary from 0.49 to 0.52, which is close to the theoretical value of 0.5.

As for the FWHM, compared to the metallic peaks, it is confirmed that the oxide peaks are wider. Satellite peaks have larger FWHM values in the range about 3.5–4.3 eV due to the strong overlap with the main peaks and the background. The metallic peaks contribute a lot to the asymmetry of the spectra.

3.2. Clean surface characterization and nitrogen segregation

After Ar^+ sputtering the alloy surface, only the Fe 2p, Cr 2p and Ni 2p core levels were identified indicating the absence of contaminants. No carbon or nitrogen were detected. It proves that the nominal amounts of these elements are below the detection limit of the XPS (< 0.5 at%). The surface composition (at%) was determined as 75Fe-15Cr-10Ni, depleted in Cr at the surface compared to the nominal bulk value (69Fe-18Cr-13Ni). However, after annealing at 700°C for 10 minutes, a N 1s feature appeared at around 397.2 eV, indicating that nitrogen, below the detection limit after sputtering, segregated on the surface upon thermal treatment. The details of the Fe 2p, Cr 2p, Ni 2p and N 1s spectra are discussed in Section 3.4. The composition of the surface obtained after annealing was (at%) 54Fe-29Cr-9Ni with 8 at% N. It should be noted that these composition values were deduced from quantitative analysis of the XPS spectra assuming an identical surface distribution for all elements. Their relative changes evidence that Cr and N segregate at the surface upon annealing. The form under which these elements segregate will be discussed in section 3.4.

Fig. 1 depicts the real-time segregation of nitrogen at the surface as illustrated by the plot of the normalized area of N 1s peak versus annealing time (Fig. 1 (a)) and the variation of the segregation rate as a function of annealing temperature (Fig. 1 (b)). The data were acquired in three successive experiments (as seen from the two discontinuities in the plots) using the XPS Snapshot mode.

Fig. 1 (a) shows that the nitrogen intensity markedly increases from 400–500°C and reaches saturation at about 680°C, which is in agreement with surface segregation of nitrogen reported after annealing a Fe-Cr alloy at 527°C [22]. When the temperature exceeds about 680°C, the N surface concentration remains unchanged. Compared to the surface annealed at 700°C during 10 min, the surface saturated at 680°C has the same amount of nitrogen.

Fig. 1 (b) provides more details on the variation of the nitrogen surface uptake. It shows that the segregation rate, as given by $\frac{dN}{dt}$, the derivative of the normalized area of the N 1s peak with respect to time, begins to increase at about 400°C and reaches its maximum at



Fig. 1. Real-time nitrogen segregation on Fe-18Cr-13Ni(100) as measured by XPS in Snapshot mode upon annealing the sputter-cleaned surface: (a) variation of normalized N 1s intensity, (b) nitrogen segregation rate and fitting curve (see text for details).

about 620°C before dropping quickly to zero after surface saturation at 680°C, showing that nitrogen segregation depends effectively on temperature. Since nitrogen starts to segregate at 400°C, it is nearly impossible to prepare an annealed and well-structured metallic surface that is nitrogen-free. As shown in Fig. 1 (b), the nitrogen segregation kinetics plot can be fitted by the following equation:

$$\frac{\mathrm{dN}}{\mathrm{d}t} = A \exp \frac{-E}{kT} \tag{1}$$

where k is the Boltzmann constant and E the activation energy for N segregation. The fitted value of E is 72 ± 5 kJ/mol. This is less than the value found for N segregation in a ferritic Fe-Mo alloy [48]. The N saturation of the oxide-free surface would therefore be easier to achieve in our case of an austenitic alloy.

Previous studies have discussed the influence of nitrogen on oxidation and wet corrosion of stainless steel [5, 22, 38, 40, 41]. It was reported that Cr and N co-segregate at the surface, as confirmed in the present work, which may promote the selective oxidation of metallic Cr enriched at the co-segregated surface. Cr and N co-segregate in the form of chromium nitride compounds [22, 40, 41], which is discussed in the following.

3.3. Surface oxidation kinetics

In order to determine the influence of oxygen pressure and temperature during the initial oxidation of the Fe-18Cr-13Ni(100) surface, oxidation runs were performed at RT, 150°C and 250°C at different oxygen pressures on the sample surface pre-annealed at 700°C. Fig. 2 presents the evolution of the normalized O 1s peak area at these three temperatures as a function of O₂ exposure (100 L) as measured in real-time by *in situ* XPS. Fig. 2 (b) enlarges the region of the first 25 L of exposure. This is the first observation of oxidation kinetics at the very early stage. In previous studies, oxidation experiments of iron or stainless steel were carried out at $P_{O_2} > 10^{-6}$ mbar [20, 21, 24, 39, 49]. At these pressures, the detailed information on oxide film growth in the very first stages is lost. This is why our study was limited to 10^{-8} mbar for low exposures (< 100 L) and did not exceed 10^{-6} mbar for larger ones, as shown by the insets in Fig. 2 (a). Since there was no evidence of discontinuity in



Fig. 2. Real-time oxygen uptake on Fe-18Cr-13Ni(100) as measured by XPS in Snapshot mode during oxidation: (a) normalized intensity of O 1s as a function of O_2 exposure in the 0–100 L range, (b) enlarged 0–25 L range. The time variation of the oxygen pressure is inserted in (a). The variation of oxidation rate with exposure is inserted in (b).

the kinetic curves recorded at RT and 150°C, the change of oxygen pressure is concluded to have no apparent influence on the oxidation kinetics in the conditions tested.

At a fixed temperature, the O 1s intensity first increases rapidly before slowing down and leveling. At RT, the oxygen uptake reaches saturation when oxygen exposure exceeds 10 L (Fig. 2 (b)). At 150°C or 250°C, the oxygen uptake is reduced after exposure to 20 L (Fig. 2 (a)), but no oxidation saturation is observed upon oxygen exposure up to 81 L and to 35 L, respectively. The inset in Fig. 2 (b) presents the oxygen uptake rate $\left(\frac{dO}{dt}\right)$ recorded *in situ* in the 0–10 L exposure range at the three temperatures. It shows that the oxidation rate reaches a maximum at 2–3 L exposure and then decreases to stabilize to around zero at 10 L at RT. At 150°C and 250°C, a slow oxidation rate is still observed after 10 L exposure.

As evidenced in Fig. 2 (b), as the temperature increases, the initial oxygen uptake is reduced. This is consistent with the decrease of the physical adsorption time (Arrhenius law) with, as a consequence, the decrease of the reaction rate at very low exposure (< 10 L). At RT, a component was detected at 531.4 eV BE in the O 1s region, close to the value attributed to adsorbed molecular oxygen [50]. This component, distinct from the oxide component peaks, was not observed in the spectra obtained at 150 and 250°C (Fig. S1 in supplementary data). This is consistent with the initial reduction of the sticking coefficient at these temperatures. In contrast, Fig. 2 (b) shows that for exposures larger than 10 L, the oxygen uptake and its saturation level increase at higher temperature. This is consistent with higher temperature promoting atomic displacement at the surface and/or atomic transport between surface and sub-surface of the sample, thus resulting in the formation of more surface oxide by oxide growth.

Studies reported in the literature were carried out at 10^{-6} mbar or higher pressure for exposures larger than 1 000 L [20, 21, 24], overlooking the initial stages of oxidation. Oxidation saturation was also observed by XPS on Fe(111) as recently reported by Davies *et al.* [21]. It was found that surface oxygen saturates after an oxygen exposure of about 100 L at 280 K (under $P_{O_2} \approx 10^{-6}$ mbar), whereas about 10 L are required to reach saturation on our alloy surface at RT. On Fe(111), the rate of oxygen uptake was almost the same at 280 K and 500 K, but the oxygen coverage reached at 500 K is higher [21] in agreement with the increasing saturation level measured at 150°C and 250°C in the present work.

Thus, it is shown by *in situ* XPS analysis of the initial stages of the Fe-18Cr-13Ni surface oxidation that there is no apparent effects of oxygen pressure on the oxidation kinetics in the tested pressure range ($< 10^{-6}$ mbar) and that the effects of temperature are different depending on oxygen exposure. The exposure of about 10 L can be estimated as the threshold below which the oxidation rate would be essentially limited by the sticking coefficient suggesting a nucleation phase reaching saturation at this exposure. Beyond 10 L, atomic displacement and/or transport would become rate limiting suggesting that the oxidation process has entered the growth phase leading to the increase of the surface oxide formation with increasing temperature.

3.4. Surface composition and oxidation mechanisms

3.4.1. Surface species and their evolution

XPS high resolution core level analysis was performed at 45° and 90° take-off angles after cumulative exposure of the surface to oxygen in order to characterize the surface species and their evolution, as well as to quantify the surface composition. Based on the oxidation kinetics data presented above, the exposure series were of 0 L, 0.5 L, 3 L, 6 L, 9 L and 14 L at RT. At 150 and 250°C, they were of 0 L, 3 L, 6 L, 9 L and 15 L, complemented with series at 21 L, 35/37 L, 49/51 L, 76 L, 102/106 L, 221/224 L and 328/346 L. No carbon contamination was detected. The reconstruction of all Fe 2p, Cr 2p and N 1s spectra (more than 300 spectra) was performed using the fitting parameters compiled in Table 1. Fig. 3 shows the reconstruction of Fe 2p, Cr 2p and N 1s spectra measured at 45° take-off angle after exposures of 0 L, 3 L, 6 L and 14 L at RT.

For the sake of clarity, the main peaks and satellites have been summed up for each oxidation state of Fe and Cr. The Fe²⁺ and Fe³⁺ components were identified based on the spectra obtained at higher temperature where they are more prominent. Comparatively, the evolution of the Cr 2p spectra was more marked at RT (as seen on Fig. 3), enabling the Cr^{3+} oxide and nitride components to be better identified.

The spectra presented in Fig. 3 show that the main cations in the surface oxide grown



Fig. 3. Reconstruction of the XPS (a) Fe 2p, (b) Cr 2p, (c) N 1s core level regions for Fe-18Cr-13Ni(100) exposed to oxygen from 0 to 14 L at RT (take-off angle 45°). The Fe and Cr components are the sum of the main peaks and their satellites. Each component is represented by a colored and shaded area; the envelope indicates the sum of all component peaks; the dots represent the measured spectra.

at RT are Cr^{3+} , Fe^{2+} and Fe^{3+} . No oxidized nickel (Ni²⁺) was observed after oxidation since only a single metallic Ni $2p_{3/2}$ component was measured at 852.8 eV BE in agreement with previous studies [20, 23, 24]. Four nitrogen component peaks assigned to CrN_b , CrN_s , N_b and N_m ($CrN_{surface}$, CrN_{bulk} , N_{bulk} and $N_{minority}$ in Table 1) were also distinguished before and after oxygen exposure. For all core levels, there are no significant differences between the spectra at 6 L and 14 L, indicating that the surface reaction reached saturation at RT in agreement with the *in situ* kinetics data presented in Fig. 2.

Even if one considers the uncertainty in the peak fitting, the spectra presented in Fig. 3 (a) yield a relative amount of Fe^{3+} and Fe^{2+} (e.g. Fe^{3+}/Fe^{2+} intensity ratio of 1.4 at 3 L exposure) markedly below that (ratio of 2) corresponding to Fe_3O_4 which suggests the presence of non-stoichiometric mixed Fe²⁺-Fe³⁺ rather than FeO oxide nuclei prior to reaching saturation in agreement with the 709.4 eV BE value of Fe^{2+} . As far as we know, the existence of FeO at the earlier stage of stainless steel oxidation is still under debate [17– 19, 36]. Several studies have demonstrated that the main surface compounds were Fe_2O_3 and Fe_3O_4 for iron oxidation under oxygen exposure larger than $10^4 L$ [19]. On the contrary, the structure of FeO or FeO-like compounds were observed in studies [17, 18] addressing the conditions of FeO formation as compared to the phase diagram for the iron-oxygen system. Graat et al. [36] proposed the presence of FeO at RT for low oxygen exposure of 22 L. Our finding of Fe^{2+} species is in agreement with this work [36] and that of Roosendaal *et al.* [18] but we also find Fe^{3+} species. Lin *et al.* [17] indicated that FeO_x , the FeO-like oxide, was the nucleating oxide formed after low oxygen exposure (10 L to 50 L) at RT. The change of x value implies that the oxygen-deficient $Fe^{2+}-Fe^{3+}$ mixed phases are first formed during the initial stages of oxidation and are transformed to Fe_3O_4 by further oxidation. We find that $Fe^{2+}-Fe^{3+}$ mixed oxide nuclei, possibly oxygen-deficient, are formed on the surface in the nucleation phase for oxygen exposures lower than 6 L for RT, 150 and 250°C. At higher exposure, the $\mathrm{Fe}^{3+}/\mathrm{Fe}^{2+}$ intensity ratio is found consistent with the presence of $\mathrm{Fe}_{3}\mathrm{O}_{4}$ as well as the BE value of Fe^{2+} (709.4 eV). However, as discussed below, angle-resolved analysis points to a different in-depth repartition of the Fe^{3+} and Fe^{2+} species.

Fig. 3 (b) shows the development of Cr^{3+} oxide and nitride components during oxidation.

 Cr^{6+} species, expected at 579.3 eV BE [25], were not found. After long exposures (> 24 000 L), Fe-Cr mixed oxides were identified by Olefjord [51]. Lince *et al.* [16] have identified only one Fe 2p peak varying at 709.5–710.8 eV BE to determine the existence of FeCr₂O₄ or Fe_{1+x}Cr_{2-x}O₄, for a Fe-16Cr(001) surface exposed to oxygen for more than 400 L. Instead, two components at 709.4 eV and 710.3 eV were used for iron oxide peak reconstruction in our data. Thus the presence of Fe³⁺ and/or Fe²⁺ cations in the grown Cr³⁺ oxide matrix cannot be excluded. It is also observed in Fig. 3 that chromium oxidizes more rapidly than iron because of its higher oxide to metal intensity ratio when oxygen exposure is lower than 14 L, *i.e.* in the nucleation phase. This is consistent with the results of Lince *et al.* [16].

Fig. 3 (c) presents the four identified nitrogen components: CrN_b , CrN_s , N_b and N_m . CrN_b is the main product of Cr-N co-segregation at the surface. It is attributed to threedimensional clusters [22, 52, 53] formed in the outer surface region including topmost and subsurface and hereafter referred to as bulk nitride. CrN_s corresponds to the two-dimensional CrN [22, 52, 54] formed at the topmost surface and referred to as surface nitride. The peak at 397.8 eV BE is identified here as interstitial nitrogen N_b [22] rather than Cr_2N (397.6 eV) [5, 22] due to the inconsistency of the CrN and N_b intensities. The broad component denoted N_m is identified in the decomposition of all spectra. Because of its broadness and low intensity, this component could originate from the sum of various minority contributions. At the very beginning of oxidation, mostly the CrN_s and barely the N_b components increase while the CrN_b component decreases markedly in intensity. As discussed above, nitrogen segregation saturation after annealing at 700°C of the sputtered surface causes the formation of a Cr-depleted region below the surface. This means that the increase of CrN_s and N_b components during oxidation is not the result of further nitrogen segregation from the bulk but the indication that the in-depth distribution of nitride has been modified by oxidation. The transformation from CrN_b to CrN_s and N_b is accelerated at 150°C and 250°C. This variation of nitrogen species was also observed in previous studies of ferritic steel upon annealing [22] and assigned to the increasing instability of CrN_b with increasing annealing temperature. In summary, there is no nitrogen surface segregation during oxidation, rather the decomposition of CrN_b modifying the nitride in-depth distribution resulting from initial

surface preparation.

Because of the complexity in oxide products, it was not attempted to distinguish the various Cr and Fe oxides according to the O spectra. Only the consistency between Cr and Fe oxides amounts and O intensity was verified.

Fig. 4 presents the intensity variation of all core level components measured as a function of oxygen exposure at the three temperatures and 45° take-off angle. Fig. 4 (a) shows the variation of the total area of the O 1s and Ni $2p_{3/2}$ core levels. The O 1s intensity increases until saturation at RT and near saturation at 150 and 250°C. The saturation level increases with increasing temperature, in agreement with the variation observed during the real-time monitoring as presented in Fig. 2. Concerning nickel, the observed continuous intensity decrease with ongoing oxidation is consistent with continuous attenuation of the signal by the growing Ni-free oxide film. No oxide of Ni was formed compared to Fe or Cr which is in agreement with their standard Gibbs energy of formation.

Fig. 4 (b) shows the variation of the Fe $2p_{3/2}$ metal and oxide core level components at the three temperatures. At RT, the Fe²⁺ and Fe³⁺ components first increase simultaneously (0.5 L). The increase of the Fe²⁺ component is stopped already at 3 L whereas that of the Fe³⁺ component goes on until saturation at 14 L. A higher temperature promotes the increase of the Fe³⁺ component but barely that of the Fe²⁺ component. This shows that Fe-18Cr-13Ni(100) sample is first oxidized to simultaneously form Fe²⁺ and Fe³⁺ species in the nucleation phase. Then, with further O₂ exposure, Fe³⁺ is preferentially formed possibly by oxidation of preformed Fe²⁺. This possibility implies that Fe²⁺ is still formed from metallic Fe since still observed with ongoing oxidation. Another possibility would be that Fe³⁺ could be produced directly from metallic Fe. In both cases, a higher temperature would promote the oxidation reaction to Fe³⁺ and atomic transport leading to the increase of the oxidation of the metallic Fe⁰ component is consistent with continuous attenuation of photoelectrons signal by the growing oxide species.

The variation of Cr^{3+} (Cr oxides), CrN and Cr metal component intensities is presented in Fig. 4 (c). The Cr^{3+} oxide intensity increases rapidly in the nucleation phase for exposure



Fig. 4. Intensity variation of XPS core level components with increasing oxygen exposure at RT, 150°C, 250°C: (a) O 1s and Ni 2p_{3/2}; (b) Fe 2p_{3/2}; (c) Cr 2p_{3/2}; (d) N 1s. Take-off angle: 45°.

below 10 L and then levels off at higher exposure. A diminution is even observed with ongoing exposure at 150 and 250°C. There are two reasons for this diminution: Cr^{3+} formation slows down while Fe³⁺ still forms and/or the Fe³⁺ oxides form above the Cr^{3+} oxides attenuating the Cr^{3+} signal. The same evolution is found for the nitride component as for the oxide Cr^{3+} component at all three temperatures. Comparing the consumption of metallic Cr^{0} and Fe⁰, it can be deduced that, in the studied temperature range, chromium is preferentially oxidized at the very beginning of oxidation in the nucleation phase below 10 L. For exposure above 10 L, a higher temperature promotes the preferential formation of Fe³⁺-containing oxides.

Fig. 4 (d) shows the evolution of the nitrogen component intensities with oxygen exposure. CrN_b , as the main product of Cr-N co-segregation at the prepared initial surface, sharply decreases in intensity for the exposure below 10 L, and even almost disappears after 6 L at 250°C. In contrast, CrN_s increases in intensity and becomes the predominant nitride species, before being attenuated by the oxide film still growing above 10 L exposure at 150 and 250 °C. N_m presents a similar intensity variation to N_b, supporting the assumption that it is related to minority structures associated to N in solution. It is concluded from the drop in intensity of all N components with increasing exposure to oxygen above 10 L exposure at 150 and 250 °C that the surface oxides develop above the remaining surface nitrides.

To sum up, these data show that the initial build-up of the surface oxide proceeds in the range of exposure below 10 L by simultaneous oxidation of iron and chromium. In this nucleation stage, Cr^{3+} formation is preferential over Fe^{3+} and Fe^{2+} formation. Fe^{2+} formation is mostly observed in the very first stage of oxidation (0.5 L). Afterwards, only Fe^{3+} is produced by further oxidation. Beyond 10 L of exposure to oxygen in the oxide growth regime, the competitive oxidation switches in favor of Fe^{3+} formation. Increasing the temperature (at 150°C and 250°C) increases the oxidation rate and quantity of formed oxide without modifying the competitive oxidation sequence. Surface oxidation destabilizes the 3D surface nitrides produced by surface preparation and the oxide films grow above the remaining 2D interfacial nitrides. No oxidized nickel enters the composition of the oxide films.



Fig. 5. (a) Layered models of the Fe-18Cr-13Ni(100) surface after 0–100 L exposure to oxygen: bulk (Fe-18Cr-13Ni), modified alloy region (different concentration of alloy elements compared to bulk), mixed region (consisting of nitride, oxide and metallic components) and oxides region. (b) Variation of the relative concentrations of Fe, Cr and Ni compounds in metal and oxide phases with increasing exposure to oxygen at RT, 150°C and 250°C. The total percentage of metallic or oxide compounds is equal to 100 at%.

3.4.2. Layered surface composition

Although previous studies [16, 19, 20, 22, 24] have shown that no abrupt interface is formed between oxide and metal, layered models have been widely applied to describe the stratification of oxidized surfaces and quantify their composition. In this work, we have performed comparative analysis of the XPS intensities measured at 45° and 90° take-off angles of the emitted photoelectrons for each chemical component in order to define its relative distribution along the surface normal and thus to determine the in-depth distribution of the surface species. Layered models were applied to quantify the surface composition and equivalent oxide film thickness. The resulting values do not account for the inhomogeneity of the growth process that can be expected due to local structural and chemical variations at the alloy surface, in particular in the nucleation phase. This aspect will be discussed separately in an article based on STM data. Still, the values reported hereafter allow us to discuss the highly relevant aspect of chromium enrichment and its variation with the build-up of the oxide film.

Fig. 5 (a) presents the layered models of surface composition that were used for the clean annealed surface and the surface oxidized up to 100 L exposure at RT, 150°C and 250°C. For all three temperatures, the layered surface composition was modeled with 3 regions: the oxide region, the mixed region (consisting of nitride, oxide and metallic components) and the modified alloy region where the compositions of alloy elements differ from that in the bulk alloy. This classical model of the top oxide layer covering the modified alloy layer with a mixed inter region as a transition between oxide and metal has been used in previous studies [17–20, 22, 24]. Prior to oxidation (0 L), the two-dimensional chromium nitride is formed at the outermost surface and the alloy is modified due to the co-segregation of Cr and N. Upon oxidation, an intermediate region mixing metal, oxide and nitride species is observed up to 50 L exposure at the interface between oxide and modified alloy; the nitride signal becomes totally attenuated by the top oxide layer for an exposure of about 100 L at 250°C. Upon higher exposure at 150°C and 250°C, the models are unchanged.

Assuming a homogeneous composition for both oxide and modified alloy layers, composition and equivalent oxide film thickness may be calculated. Variations of the relative concentration of Fe, Cr and Ni compounds in metallic and oxide phases as a function of exposure are shown in Fig. 5 (b). The equivalent thickness values were calculated from the intensity ratios between the metal and oxide components of the Fe and Cr $2p_{3/2}$ core level based on the exponential attenuation of the photoelectron intensity with depth [15, 55, 56]. These thicknesses are given in Fig. 6.

These composition and thickness results should be taken with caution as they are calculated from a simplified model. However these quantities are relevant for comparison with previous studies and relative changes in concentrations help us to understand the underlying mechanism that govern the oxide growth process.

Based on an analysis of angular dependent photoemission spectra, a more precise model of the surface may be proposed. Fig. 6 presents the schematic view of the species distribution as a function of oxygen exposure at RT, 150°C and 250°C. This is the first complete stratification scheme of oxidized surface of a stainless steel at the initial oxidation stage.

The equivalent thicknesses of the oxide regions ranges between 1.1–2.2 nm, calculated based on a simplified homogeneous model including the interfacial oxide/metal mixed region. This is consistent with oxide films reported to be 1–4 nm thick when formed on Fe or Fe-base alloys [16, 17, 19, 20, 22–24, 36, 57]. At RT, the calculated thickness is stable at 1.6–1.7 nm after the 6 L exposure, in agreement with the saturation observed in the oxygen uptake (Fig. 2 and 4) and with 1.8–2 nm thickness reported for the native oxide film formed on the same austenitic stainless steel surface [23]. At 150 and 250°C, the oxide equivalent thickness increases with exposure also following the same trend as the oxygen uptake (Fig. 2). However, the increase observed in the oxygen uptake at saturation with temperature increasing to 150 and 250°C (Fig. 4) is not reproduced by the calculated thickness which yields similar values of 2.0–2.1 nm. A possible explanation is that the simplified model used for calculation does not accurately account for the deviation of the morphology of the surface oxides grown at 150 and 250°C from a continuous and uniformly thick film. This suggests that the films grown at higher temperature may include 3D oxide islands and thus may be non-uniform in thickness.

On the as-prepared surface (0 L), the modified region is enriched in metallic Cr (29 at% instead of 18 at% in the bulk alloy) and stratified. Metallic Cr is found concentrated in the upper part of the modified alloy region due to co-segregation with nitrogen after annealing at 700°C. The two-dimensional CrN_s component is located on the topmost surface; while the three-dimensional CrN_b component and interstitial nitrogen N_b are found in this Cr-enriched upper part. Metallic Fe is found to be concentrated just below the Cr-enriched upper part while metallic Ni concentrates at the bottom of the modified alloy region. After oxidation at 150 and 250°C, our angular analysis shows that these nitrogen species are included in the



Fig. 6. Schematic stratification profiles of the Fe-18Cr-13Ni(100) surface after 0–100 L exposure to oxygen at RT, 150°C and 250°C. The color blocks areas represent the relative proportions of components in the oxide and modified alloy regions. The equivalent thickness of oxide region (calculated from layered models in Fig. 5) is indicated on the right.

mixed interfacial region developed between growing oxide and modified alloy. Only at RT, CrN_b and N_b remain in the upper part of the modified alloy region underneath the oxide.

According to previous studies [22, 24], there would be no clear separation between Cr oxide and Fe oxide in the oxidized surface. An amorphous oxide FeCr_2O_4 -like phase could exist after a large exposure at a higher temperature (1000 L, 300°C) [20]. However, our angle-resolved analysis supports the formation of layered surface oxides with different relative positions of the Cr³⁺, Fe³⁺ and Fe²⁺ species. The mechanisms by which the component distribution varies under very low exposure at the three studied temperatures are as follows.

Upon oxidation at RT, the surface oxide is found markedly enriched in Cr^{3+} already after an oxygen exposure of 6 L as a result of the preferential oxidation of chromium due to their relative standard Gibbs energy of formation. The cation fraction of Cr^{3+} in the oxide is then calculated as 59 at%. Fe²⁺ and Fe³⁺ oxide species are also formed by competitive oxidation but in lower amounts. Their fractions are 13 at% and 28 at%, respectively. The Fe²⁺ oxide species are found mixed with the CrN_s species in the interfacial layer underneath the Cr^{3+} and Fe³⁺ mixed layer, likely as a result of their competitive formation on the bare metallic surface in the very first oxidation step (0.5 L) as discussed above. Further oxidation beyond saturation has little effect on the layered oxidized structure and composition. The cation fraction is 56 at% for Cr^{3+} , 13 at% for Fe²⁺ and 31 at% for Fe³⁺ at 35 L. The modified alloy region has an unchanged concentration of 68%Fe-19%Cr-13%Ni at 6 L and 69%Fe-19%Cr-12%Ni at 35 L if we assume a homogeneous distribution of species, still not depleted in Cr and with a Cr-enriched metallic layer at the top. This means that the surface saturated with oxygen at RT has still the capacity to form Cr oxides in order to grow a protective surface layer.

Upon oxidation at 250°C, the same layered structure of oxide film is found with a mixed Cr^{3+}/Fe^{3+} layer above the Fe²⁺-containing interfacial layer. Moreover, the results show that the structure is determined not only by the competitive oxidation of Cr and Fe, but also by their atomic mobility. At 6 L (Fig. 6), metallic Cr is found preferentially in the interfacial region containing Fe²⁺ and Cr-nitrides, suggesting its transport from the modified alloy region in order to sustain Cr oxidation since the oxide film is equally enriched in chromium

(the cation fraction is 59 at% for Cr^{3+} , 13 at% for Fe^{2+} and 28 at% for Fe^{3+} in Fig. 5 (b)). As a result, the modified alloy region is enriched with metallic Fe in its upper part. Thus, the competitive oxidation of iron is promoted gradually after 10 L exposure and Fe oxidation prevails over Cr oxidation at the oxide-metal interface. As a result, the Cr enrichment in the growing oxide film decreases with cation fractions of 34 at% for Cr^{3+} , 14 at% for Fe^{2+} and 52 at% for Fe^{3+} after 35 L exposure, and the interfacial region and the upper part of the modified alloy regions become depleted in metallic Cr. Subsequent oxygen exposure (> 50 L) still preferentially consumes Fe leading to further decrease of the Cr enrichment in the growing oxide film with cation fractions of 22 at% for Cr^{3+} , 12 at% for Fe²⁺ and 66 at% for Fe^{3+} after 106 L. At this exposure, the Fe^{3+} -rich oxide is found predominantly below the Cr^{3+} -rich oxide, which suggests its preferential formation at the oxide-metal interface, *i.e.* under the Cr oxide. The metallic fraction in the modified alloy region below the oxide varies from 69%Fe-19%Cr-12%Ni at 6 L to 74%Fe-14%Cr-12%Ni at 35 L and 68%Fe-16%Cr-16%Ni at 106 L, indicating that iron is relatively stable, chromium consumed and Ni enriched. These variations suggest that Fe oxidation is sustained by diffusion of metallic Fe from sub-surface region to the modified alloy region, whereas that of Cr is much less so possibly due to slower diffusion. The slight enrichment of Ni is detected in the Cr-depleted region, similarly to the Ni segregation reported in a study of Fe-20Cr-18Ni oxidation at 300°C [20]. We conclude from these data obtained at 250°C that the oxide film growth and resulting surface composition are ruled by thermodynamically-dominated competitive oxidation of Cr and Fe in the very first stages like at RT. When oxygen exposure exceeds 50 L, Fe oxide formation at oxide-metal interface becomes dominant and dependent on diffusion in the modified alloy region. This dependence of the competition between Cr oxidation and Fe oxidation over exposure is consistent with transformation from thermodynamically dominated to kinetically controlled reaction processes [35].

According to our kinetics results discussed above, the oxide film growth at 150°C is intermediate between those at RT and 250°C. Similarly to the results obtained at RT and 250°C, the oxide is enriched in Cr^{3+} in the earliest stage of oxidation with cation fractions of 63 at% for Cr^{3+} , 14 at% for Fe^{2+} and 23 at% for Fe^{3+} at 6 L. The modified alloy region

is still not depleted in chromium with a concentration of 66%Fe-23%Cr-11%Ni. For the exposure of about 40 L, the oxide enrichment in Cr³⁺ decreases (cation fractions of 48 at% for Cr³⁺, 9 at% for Fe²⁺ and 43 at% for Fe³⁺) like at 250°C with Cr depletion in the modified metallic alloy (73%Fe-15%Cr-12%Ni). For further exposure (> 50 L), the Cr³⁺ enrichment still decreases in the oxide (cation fractions of 39 at% for Cr³⁺, 9 at% for Fe²⁺ and 52 at% for Fe³⁺) owing to preferential Fe oxidation. The Cr concentration in the modified metallic layer is unchanged (75%Fe-14%Cr-11%Ni). This confirms that the preferential consumption of iron by oxidation is compensated by diffusion from the sub-surface to the modified alloy region. Like at 250°C, Ni becomes concentrated in the upper part of the modified alloy region. A noticeable difference with the oxide film grown at 250°C is the preferential formation of Fe³⁺ oxide at the topmost surface, as at RT. This suggests that at 150°C iron transport through the Cr³⁺/Fe³⁺ oxides is dominant for the oxide growth for exposures larger than 35 L, whereas at 250°C oxygen transport to the oxide-metal interface would be dominant.

Table 2 compiles the data of surface composition and thickness obtained in this work, the data for a native oxide film formed by exposure to ambient air and a passive film formed by anodic polarization on the same alloy surface at RT, and the data for a thermal oxide film formed by exposing a Fe-17Cr alloy surface to oxygen at 327°C. At RT, the surface saturated under oxygen in the present work is found more enriched in Cr^{3+} than the surface aged for 20 h under ambient air [23]. This can be assigned, at least partly, to the Cr-N co-segregation resulting from surface preparation in our work and that would promote the selective oxidation of chromium as previously reported [22]. However, the oxide film grown at RT under oxygen in the present work in much less enriched in Cr^{3+} than a passive film produced in acidic aqueous solution [23], which can be assigned to the absence of preferential dissolution of iron oxides promoting chromium enrichment in acidic media, as also found on a ferritic Fe-17Cr alloy oxide film grown by exposure to oxygen [22] and anodic polarization [58]. With the increase of temperature to 250°C, it is found in the present work that the chromium oxidation is hindered by the depletion of metallic Cr underneath the growing oxide layer, while the diffusion of metallic Fe from the bulk enables to sustain iron oxidation, thus

Sample	Temperature	Experimental condition	${ m Thickness} \ { m (nm)}$	Oxide composition	Modified metallic phase
) RT	$O_2, 100 L^*$	1.7	$56\% Cr_{ox}$ -44%Fe _{ox}	69Fe-19Cr-12Ni
Fe-18Cr-13Ni(100)		Air, 20 h [23]	2.0	$21\% Cr_{\rm ox}{-}79\% Fe_{\rm ox}$	50Fe-19Cr-31Ni
		$0.5 \text{ M H}_2\text{SO}_4, 20 \text{ h} +500 \text{ mV/SHE} [23]$	2.3	$91\% Cr_{\rm ox}{-}7\% Fe_{\rm ox}$	50Fe-15Cr-35Ni
		$O_2, 6 L^*$	1.7	$59\% \mathrm{Cr_{ox}}{-}41\% \mathrm{Fe_{ox}}$	69Fe-19Cr-12Ni
Fe-18Cr-13Ni(100)	$250^{\circ}\mathrm{C}$	$O_2, 35 L^*$	2.2	$34\% \mathrm{Cr_{ox}}{-}66\% \mathrm{Fe_{ox}}$	74Fe- 14 Cr- 12 Ni
		$O_2, 100 L^*$	2.1	$22\% \mathrm{Cr_{ox}}{-}78\% \mathrm{Fe_{ox}}$	68Fe- 16 Cr- 16 Ni
	327°C	$O_2, 10 L [22]$	1.0	$45\% Cr_{\rm ox}{-}55\% Fe_{\rm ox}$	86Fe- 14 Cr
		$O_2, 30 L [22]$	2.2	$29\% Cr_{\rm ox}{-}71\% Fe_{\rm ox}$	86Fe- 14 Cr
Fe-17Cr		$O_2, 100 L [22]$	3.5	$17\% Cr_{ox} - 83\% Fe_{ox}$	86Fe- 14 Cr
	RT	$2 \times 10^{-2} \text{ M Cl}^-, 27 \text{ h} +390 \text{ mV/SHE} [58]$	3.1	$75\% Cr_{\rm ox}{-}25\% Fe_{\rm ox}$	81Fe-19Cr

Table 2 Thickness and chemical composition (at%) of thermal oxide film, native oxide film and passive oxide film formed on Fe-18Cr-13Ni(100) and Fe-17Cr, assuming homogeneous distribution of species

* This work

leading to a gradual decrease of the Cr^{3+} enrichment with increasing exposure. Compared to the thermal oxidation of a ferritic Fe-17Cr sample (also with surface nitrides) at 327°C [22], the variation of the Cr^{3+} enrichment on our austenitic sample follows the same trend.

4. Conclusion

Real-time monitoring of the evolution of a model 304 stainless steel monocrystal surface, Fe-18Cr-13Ni(100), in the early stages of oxidation at RT, 150°C and 250°C was achieved using *in situ* XPS in Snapshot mode analysis and complemented by high resolution core level analysis of the growth mechanisms and Cr enrichment. The work revealed, for the first time to our knowledge, the mechanisms of stratification during the early build-up of the surface oxide on a model austenitic stainless steel surface at different temperatures.

The results show that, on the oxide-free surface, nitrogen segregation is detected upon heating at 400°C and increases significantly above 500°C to reach a saturation at 680°C. The activation energy of nitrogen segregation is 72 ± 5 kJ/mol. Marked Cr enrichment of the as prepared metallic surface was observed as a result of Cr segregation and Cr-N co-segregation with N forming mainly three-dimensional CrN_b particles with minority two-dimensional CrN_s and interstitial nitrogen N_b. Surface oxidation destabilizes the 3D chromium nitrides and the oxide films grow above the 2D surface nitrides remaining at the surface.

The temperature dependence of the surface oxygen uptake as measured by real-time monitoring evidences two regimes for the growth of the surface oxide. Up to about 10 L of oxygen exposure, the uptake is faster with a maximum at 2–5 L and decreases with increasing temperature, suggesting a nucleation regime of the surface oxide limited by oxygen adsorption. Beyond 10 L, the uptake is saturated at RT but increases with temperature, suggesting a growth regime of the surface oxide limited by atomic displacement and mobility. The oxygen pressure in the range below 10^{-6} mbar is found to have no effect on the oxidation rate.

In the nucleation regime leading to saturation, Cr^{3+} formation is preferential over Fe³⁺ formation and Fe²⁺ formation is mostly observed in the very first stage of oxidation (0.5 L). Nickel is not oxidized. A surface layer of strongly Cr^{3+} -enriched oxide is thus formed over the mainly Fe²⁺ oxide (FeO-like) species mixed with the CrN_s species in the interfacial layer underneath the Cr^{3+}/Fe^{3+} mixed layer. Further oxidation beyond saturation at RT causes no marked evolution of the oxide layered structure, composition and thickness. The modified alloy region is still not depleted in metallic Cr and thus able to further grow a Cr^{3+} -enriched protective surface layer.

At 150 and 250°C, the oxide formed in the nucleation regime is also enriched in Cr^{3+} but the modified alloy region gets depleted with metallic Cr. As a result, the competitive oxidation of iron is gradually promoted in the oxide growth regime leading to the decrease of the Cr^{3+} enrichment in the oxide. Fe oxidation is sustained by diffusion of metallic Fe from sub-surface region to the modified alloy region. At 150°C, Fe³⁺ oxide was found to form at the topmost surface, whereas at 250°C it was at the oxide-metal interface, suggesting a switch of the oxide growth from predominant iron transport through the Cr^{3+}/Fe^{3+} oxide at 150°C to predominant oxygen transport at 250°C for exposures beyond 35 L.

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