Influence of dew point on the selective oxidation during annealing of cold rolled DP and Ti–IF steels

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Abstract

Cold rolled steels for the automobile industry are protected against wet corrosion by Zn coatings, applied by ‘Hot Dip Galvanizing’. The industrial annealing process leads to the formation of surface oxides, partially removed in a reduction annealing step (5 % H₂–N₂, D.P –30°C, 820 °C) before galvanizing. However, oxides of alloying elements like Mn, Al, Si, Cr, are not reduced during annealing and failures occur in the zinc coating due to a bad wettability of the surface with Zn bath. The most important parameter in the oxidation process is the dew point of the reduction–annealing atmosphere. By increasing the dew point, from –60 to 0°C, it is possible to attain conditions for the selective oxidation and to change the oxidation mode of elements Al, Mn, Si etc. from external to internal. Cold rolled DP and different Ti–IF steels annealed at 820°C for 60 s in 5 % H₂–N₂ atmosphere with different dew points (–60, –30 and 0°C) are investigated, regarding the chemical composition, amount, distribution and size of the surface oxides. XPS depth profiles of the oxide layers and results from FE–SEM with EDX are used to elucidate the oxidation behaviour of Mn, Si and Al on the steel surfaces.

**Keywords:** dew point, selective oxidation, external oxidation, internal oxidation.
Introduction

A new generation of high strength steels like dual phase, TRIP and Interstitial Free (IF) meet more and more demanding criteria for the steels in auto industry, such as lighter and stiffer car bodies which improve safety and reduce fuel consumption, good formability and weldability. Good corrosion protection of these steel sheets is as important as a good mechanical properties and is more or less successfully achieved by galvanizing. One of the main still existing problems in the galvanizing process is unsatisfying wettability of the steel surfaces with Zn bath due to presence of oxides.

After cold-rolling the sheets always undergo an annealing to accomplish the recrystallization and to remove residual oil from the rolling process. The annealing is always done in a protective hydrogen–nitrogen atmosphere which is reducing for iron oxides. During the annealing process at high temperatures (= 800°C) very fast diffusion and segregation of atoms of alloying elements occurs at the surface. Elements like Mn, Al, Si, Ti etc. with a very high affinity to the oxygen, form in the annealing cycle oxides which stay on the surface and are “poorly” wetted by the Zn bath. Results of experiments with a higher hydrogen content (15 %) compared to those in standard 5 % H₂–N₂ annealing atmosphere show no reduction of mentioned oxides as well [#ref1]. Thermodynamical calculations [#ref2] show that is practically impossible to reach the annealing conditions in which the oxidation of Al, Mn, Si would be avoided. These oxides are present rather in the form of islands than as continuous layer. Whether the oxidation is in external or internal mode depends on a few criteria well described by Wagner [#ref3].

The aim of this work is to give an overview on selective oxidation of different steels and in different annealing conditions.

Not only the segregation of metallic but also of non–metallic elements like C, S, P and B plays an important role in oxidation–reduction process during the annealing.
Experimental

One dual phase steel (DP 500, ferritic-austenitic) and three IF steels (Ti–IF, TiNb–IF 1, TiNb–IF 2, ferritic) were investigated. The steel compositions, concerning the most important elements for this investigation, is presented in Table 1:

Table 1: Composition of investigated steels (concerning the most important elements for this investigation).

<table>
<thead>
<tr>
<th></th>
<th>C</th>
<th>N</th>
<th>Mn</th>
<th>Si</th>
<th>Al</th>
<th>B</th>
<th>Cr</th>
<th>P</th>
<th>S</th>
<th>Nb</th>
<th>Ti</th>
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<tr>
<td>DP 500</td>
<td>70</td>
<td>5,9</td>
<td>1400</td>
<td>100</td>
<td>34</td>
<td>3,2</td>
<td>450</td>
<td>14</td>
<td>–</td>
<td>–</td>
<td>25</td>
</tr>
<tr>
<td>Ti–IF</td>
<td>2,8</td>
<td>3,3</td>
<td>85</td>
<td>6</td>
<td>44</td>
<td>0,1</td>
<td>16</td>
<td>6</td>
<td>6</td>
<td>–</td>
<td>73</td>
</tr>
<tr>
<td>TiNb–IF 1</td>
<td>2,6</td>
<td>5</td>
<td>370</td>
<td>12</td>
<td>15</td>
<td>3</td>
<td>36</td>
<td>1,1</td>
<td>21</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>TiNb–IF 2</td>
<td>3,9</td>
<td>traces</td>
<td>192</td>
<td>7</td>
<td>21</td>
<td>traces</td>
<td>10</td>
<td>12</td>
<td>6,3</td>
<td>28</td>
<td>20</td>
</tr>
</tbody>
</table>

Samples of 15 x 15 mm in size were cut from cold rolled steel sheets. After the cleaning in a weak alkali solution and ultrasonic bath with ethanol for 15 min, samples were annealed in a cycle used in the industry. The annealing cycle is shown in Fig. 1.

Figure 1: The annealing cycle.
Experiments were done in 5 % H$_2$–N$_2$ atmosphere at three different dew points –60, –30 and 0°C (p(H$_2$O) = 11, 370 and 5.99 $\cdot$ 10$^3$ vppm respectively) and after reaching 820°C samples were held for 60 s at this temperature. Well defined heating regime is accomplished using an IR furnace controlled with thermo regulator. The oxygen content is measured with Oxygen analyser (fuel cell sensor) and was below 2 ppm during the experiments. The water vapour content (p(H$_2$O)) is measured with a Moisture Analyser (Al$_2$O$_3$ sensor).

After annealing the samples were taken out and transported to the XPS (X-ray photoelectron spectroscopy) and FE-SEM (Field emission scanning electron microscopy) for the further investigation and analysis.

XPS (Quantum 2000) and LEO 1550 VP FE-SEM were used for the surface investigations.

**Results and Discussion**

As well as the annealing conditions the steel composition plays an important role in the oxidation as already reported [4,5,6]. FE-SEM Images of the steel surfaces after annealing at dew point –60°C presented in Fig. 2 (a–d) show significant differences between the steel surfaces.

![Fig. 2. a): FE-SEM image of DP 500 surface.](image)

![Fig. 2. b): FE-SEM image of Ti–IF surface.](image)
The surface of DP 500 (Fig. 2 (a)) differs significantly in oxide shape and distribution from the rest three IF grades. The XPS measurement on DP 500 as well as EDX analysis confirmed that mostly BN and small (around 100 – 200 nm in diameter) globular Al oxides are present on the surface. Hexagonal BN can be formed according to two mechanisms: I) Diffused B from the steel can react with the nitrogen from the N-rich atmosphere and BN forms, or II) nucleation occurs through cosegregation of boron and nitrogen or by precipitation of the components form the supersaturated matrix [ref7]. For the I mechanism to occur nitrogen has to react on the surface according to:

\[ \text{N}_2 = 2\text{N} \text{(dissolved)} \]  

(1)

This reaction has a high activation energy and needs high temperatures. Segregation of elements like sulphur, oxygen and other surface active elements can retard this reaction. h–BN can precipitate at 700–850°C by mechanism II if the material has dissolved N in concentration > 1000 wt ppm and dissolved boron > 100 wt ppm (results on austenitic stainless steel) [ref8] and the same authors found that segregated nitrogen trapped boron. In the case of our steel both mechanisms are possible and it seems that rather a thin layer of very small BN particles is formed. We assume that BN forms by mechanism I. A small amount of CrN found by XPS analysis leads to the conclusion that N from the steel reacts with Cr (Cr and N cosegregate at lower temperatures) and with Al, since AlN particles are
seen with FE-SEM and confirmed with EDX analysis. Al peak is at 75,62 eV corresponding to Al₂O₃ and some weak Mn peak at 641,51 eV corresponds to MnO. Comparison of XPS peaks on all samples is presented in Fig. 3.

Fig. 3: The XPS spectra of Al, Mn, N and B on different steels at dew point –60°C.

Ti–IF with the highest Al content of all investigated steels shows a distinct Al peak (Fig. 3) but on TiNb–IF 1 steel the Al peak is as strong as on Ti–IF steel indicating that not only the concentration of elements controls segregation and oxidation rates. The Al peak at 75,96 eV belongs to Al in Al₂O₃ while the Mn peak at 641,81 eV belongs to Mn in MnO. The oxide particles are more or less 50 nm in diameter (Fig. 2 (b)). The B peak at 189,46 eV, belonging to BN, is found on the surface (although is peak hardly visible (Fig. 3)) as well as TiN formed by the reaction of Ti and N from the matrix or by chemisorptions induced segregation.

Morphologically the surface of TiNb–IF 1 looks like the Ti–IF surface (FE–SEM image, Fig. 2 (c)). Oxides smaller than 50 nm in diameter are distributed everywhere on the surface, but more numerous at the grain boundaries. They were identified as Al₂O₃ and MnO. B is present
as BN and B₂O₃ oxide on the surface. This is difference compared to the DP 500, where only BN is formed. This can be partly explained by the ferritic matrix of TiNb–IF 1. It is known that BN grows epitaxially on (111) fcc alloys, so the nucleation rate of BN is maybe slower in Ti–IF steel. The B peak at 192,54 eV belongs to B in B₂O₃ and is 65,75 % of B signal. Formation of B₂O₃ at 820°C is thermodynamically very favourable [#ref9].

On TiNb–IF 2 surface very small (less than 20 nm) oxides are formed during the annealing at dew point –60°C. This steel contains B and N in traces. The surface concentrations of Al and Mn are less than on the other steels and do not correspond to the bulk concentrations. The Al peak is at higher energy, 77,4 eV, than in the case of other steels and corresponds to Al in Al₂O₃ as well [#ref10]. Mn peak lies also at the binding energy of 645,5 eV (higher than in other three steels) and can be attributed to MnO/Mn₂O₃ [#ref11]. EDX analysis shows that TiNb carbides formed by reaction with C from the steel and also some nitrides are present on the surface. This N is probably dissolved on the surface from the atmosphere.

Table 2: Compounds and elements found on the steel surfaces after annealing at dew point –60°C.

<table>
<thead>
<tr>
<th>Industrial grade</th>
<th>Dew point (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>–60</td>
</tr>
<tr>
<td><strong>DP 500</strong></td>
<td>Al₂O₃ and MnO external oxides, BN, some AlN and some CrN</td>
</tr>
<tr>
<td><strong>Ti–IF</strong></td>
<td>Al₂O₃ and some MnO external oxides, some BN, some TiN, some S</td>
</tr>
<tr>
<td><strong>TiNb–IF 1</strong></td>
<td>Al₂O₃ and some MnO external oxides, B₂O₃, some BN, some TiN, some S</td>
</tr>
<tr>
<td><strong>TiNb–IF 2</strong></td>
<td>Al₂O₃ and some MnO/Mn₂O₃ external oxides, some TiNbC, some TiNbN, some S</td>
</tr>
</tbody>
</table>
Increasing the dew point to -30°C, a higher amount of H₂O ≈ 370 vppm, is introduced in the annealing atmosphere. This determines the p(O₂) in system (p(O₂) = 1.06 ⋅ 10⁻²⁴), according to the reaction:

\[ \text{H}_2\text{O} = \text{H}_2 + \frac{1}{2} \text{O}_2 \]  \hspace{1cm} (2)

Under these higher pressures of O₂ and H₂O conditions for the change of oxidation mode are met leading to the internal Al oxidation and external Mn, Si, Cr oxidation. According to Wagner theory there is a critical ratio between oxygen and solute element permeability which defines the oxidation mode:

\[ N_0(S) \cdot D_0 >> N_b(O) \cdot D_b \] – internal oxidation,

\[ N_0(S) \cdot D_0 << N_b(O) \cdot D_b \] – external oxidation.

\( N_0(S) \) – O₂ solubility in alloy (atom fraction); \( D_0 \) – diffusivity of oxygen in alloy (cm²s⁻¹); \( N_b(O) \) – solute B initial concentration; \( D_b \) – diffusivity of the solute B in alloy (cm²s⁻¹).

By increasing the dew point of the gas atmosphere the surface oxygen concentration increases and a change from external to internal oxidation is possible.

FE-SEM images of the steels surfaces at DP = -30°C are presented in Figs. 4. (a–d).

Fig. 4. a): FE-SEM image of DP 500 surface (dew point -30°C).

Fig. 4. b): FE-SEM image of Ti–IF surface (dew point -30°C).
Clear differences in the surface morphology and chemistry are visible and more exhibited between the investigated steels. On DP 500 surface oxides are different in size (from 100 nm to 1µm in diameter) and morphology (globular, lense-like to undefined shape) while TiNb-IF 2 surface is covered with oxides mostly smaller than 20 nm (Fig. 4 (d)). XPS analysis (Fig. 5) shows Al, Mn, Si and Cr XPS signals for the different steels.

Fig. 5: The XPS peaks of Al, Mn, Si and Cr on different steels at dew point –30°C.
Generally the intensity of the Al signal decreases for the increased dew point (Fig. 5). The peak energy still corresponds to Al$_2$O$_3$ and since Al has the highest affinity toward the oxygen of all investigated elements, one can conclude that the decreased amount of Al oxide present on the surface is due to the shifting of the oxidation to the internal. This is additionally confirmed by an increase of the Al signal with sputter depth. On DP 500 and Ti–IF surfaces the Mn peak is at 642.03 eV and 642 eV respectively, belonging to MnSiO$_3$ [\textsuperscript{12}]. Some amount of Mn is present as MnO (~ 7%) and B is almost undetectable. Mn present on TiNb–IF 1 is oxidised and EDX qualitatively analysis of oxide particles yielded the results presented in Figs. 6 (a, b). It can be assumed that B is more likely present as oxide than as segregated atoms [\textsuperscript{1}].

According to the phenomenon of quasi–sudden oxidation of the surface, due to B oxidation [\textsuperscript{1}], the surface oxygen concentration decreases and leads to the conditions with a lower dew point, explaining also a stronger signal of Al on the surface of TiNb–IF 1 compared to TiNb–IF 2 with a higher Al concentration in bulk (three times higher). The same authors[\textsuperscript{1}] found on the surface of TiNb–IF steel a mix of Al$_2$O$_3$ and Mn$_2$SiO$_4$ oxides.

Fig. 6: FE–SEM image of TiNb–IF 1 surface with EDX analysis of oxide particles.
On Figs. 6 (a, b) two kind of oxides can be distinguished, Mn and B oxide on the surface and Al and Si oxides at the grain boundaries. A further confirmation of Mn, B oxide presence on TiNb–IF 1 surface is confirmed by an AES mapping (Fig. 7) where brighter areas correspond to stronger intensities.

![Figure 7: TiNb–IF 1 steel surface, AES mapping.](image)

Al, Si oxides below Mn, B oxides are more likely only at the grain boundaries since the grain boundary transport of oxygen is rate-controlling process for growth of Al₂O₃ scales at high temperatures while for Mn the growth of oxide film above the grains is found as well. The conclusion is that Mn/B oxides are formed on the surface and below these oxides at the grain boundaries oxidise Al and Si. Nucleation and growth of oxides are orientation dependent (Fig. 4 (c)) and faceting is more exhibited with increasing a dew point due to oxygen adsorption.

On TiNB–IF 2 surface Mn peak at 640,8 eV corresponds to MnO and the other at higher energy can be attributed to Mn silicate, although the Si peak is hardly detectable. This excess of Mn oxides is understandable if we know that if Mn/Si ration (in weight %) is higher than 4 an excess of Mn is expected on the surface [#ref7].
Chromium, externally oxidised, is also found on steel surfaces (Fig. 5) and the Cr peak is at 576.9 eV, 577.2 eV and 575.2 eV for DP 500, Ti–IF, TiNb–IF 2 respectively, corresponding to Cr in Cr2O3.

The results of compounds and elements found on the surfaces after annealing at the dew point –30°C are summarized in Table 3.

Table 3: Compounds and elements found on the surfaces after annealing at dew point = –30°C.

<table>
<thead>
<tr>
<th>Industrial grade</th>
<th>Dew point (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>–30</td>
</tr>
<tr>
<td><strong>DP 500</strong></td>
<td>External MnSiO3, some MnO and Cr2O3, Al2O3 oxides shifting to internally oxidized, AlN</td>
</tr>
<tr>
<td><strong>Ti–IF</strong></td>
<td>External MnSiO3, some MnO and Cr2O3 oxides, some Al2O3 oxide shifting to internally oxidized, TiN</td>
</tr>
<tr>
<td><strong>TiNb–IF 1</strong></td>
<td>External Mn, B – oxides, and below Al and Si oxides at the grain boundaries, TiN, some S</td>
</tr>
<tr>
<td><strong>TiNb–IF 2</strong></td>
<td>External MnO, Mn, Si – oxides and Cr2O3, TiNb carbides</td>
</tr>
</tbody>
</table>

FE–SEM Images of steel surfaces after annealing in the atmosphere with dew point 0°C are presented in Figs. 8 (a–d).

![Fig. 8. a): FE–SEM image of DP 500 surface (dew point 0°C).](image1)

![Fig. 8. b): FE–SEM image of Ti–IF surface (dew point 0°C).](image2)
Annealing in the atmosphere with the dew point (0°C) leads to decrease of the Mn signal (XPS measurement, Fig. 9).

![Fig. 8. c): FE-SEM image of TiNb–IF 1 surface (dew point 0°C).](image)

![Fig. 8. d): FE-SEM image of TiNb–IF 2 surface (dew point 0°C).](image)

Fig. 8: FE-SEM images of TiNb–IF surfaces.

![Fig. 9: XPS peaks of P and Mn on different steels at dew point 0°C.](image)

The strongest signal of Mn is found on the surface of **DP 500** since this alloy has the highest Mn content. Selective oxidation of P is a dominant process at dew point 0°C and these Fe, Mn phosphates are particularly observed and confirmed by EDX at grain boundaries (darker oxides in Figs. 8 (c) and 8 (d)). Although P bulk concentration in **DP 500** is smaller than in **TiNb–IF 1**, the P peak is weaker on the surface of the latter (Fig. 9) and the P peak intensity is the strongest for **TiNb–IF 2** with 0.012 wt % P leading to the conclusion that P surface concentration is not only dependent on bulk concentration. This phenomenon is closely related to the concentration of N, B and Si in the alloy. **DP 500** alloy has the highest Si and high N and B concentrations whereas **TiNb–IF 2** contain B and N in traces and has a
low Si content. A clear peak of S found on the surface of TiNb–IF 2 indicates that in this case S did not impede the P segregation. Possibly formed compound on these surfaces is Fe, Mn phosphate, hardly detectable are Si and Cr on the surface which indicates shifting of Si and Cr oxidation in the internal mode.

Table 4: Compounds and elements found on the steel surfaces after annealing at dew point = 0°C.

<table>
<thead>
<tr>
<th>Industrial grade</th>
<th>Dew point (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
</tr>
<tr>
<td><strong>DP 500</strong></td>
<td>External Fe, Mn phosphates</td>
</tr>
<tr>
<td><strong>Ti–IF</strong></td>
<td>External Fe, Mn, phosphates, some S, TiN</td>
</tr>
<tr>
<td><strong>TiNb–IF 1</strong></td>
<td>External Fe, Mn phosphates, TiC</td>
</tr>
<tr>
<td><strong>TiNb–IF 2</strong></td>
<td>External Fe, Mn phosphates, some S, TiNb carbides</td>
</tr>
</tbody>
</table>

Conclusions

Due to the selective oxidation phenomenon, modifications of steel surfaces can be obtained with a change of oxidation–reduction conditions. Changes of the dew point of the annealing atmosphere, from –60 to 0°C leads to external oxidation of Al and Mn at –60°, Mn, Si and Cr at –30°C and P at 0°C, and internal oxidation of Al at –30°C and 0°C, Si and Cr at 0°C. On all investigated alloys, DP 500, Ti–IF, TiNb–IF 1 and TiNb–IF 2, external oxidation of Al is observed at dew point –60°C and Al₂O₃ is formed. N₂ present in the annealing atmosphere in a high amount, reacts with B, fast segregating from the steel, and forms BN. This occurs specially on DP 500 due to austenitic–ferritic structure, since hexagonal BN epitaxially grows on (111) fcc. This BN is rather formed by adsorption of N from N–rich atmosphere than by cosegregation of B and N from alloy. BN causes a “screening effect” to further absorption of O₂ on the steel surface and oxidation. Some externally oxidised Mn is also found at this low dew point. With increasing dew point (–30°C), p(H₂O) around 370 vppm, the higher
pressure of O₂ leads to Al internal oxidation, below the surface, which is seen from the decreasing of Al signal on XPS spectrum. External oxidation of Mn is maximal at dew point –30°C, as well as Si and Cr oxidation. Formation of both MnSiO₃ and MnO is possible, depending on the Mn/Si wt % ratio, while Cr is mostly found as Cr₂O₃. In the case of TiNb–IF 1 alloy Mn, B oxides are found on the surface, and below these at the grain boundaries Al and Si oxides. External P oxidation is a dominant process at the dew point 0°C, most probably Mn₃(PO₄)₂ or Fe, Mn phosphates are formed. Mn, Si and Cr are almost undetectable at this dew point since their oxidation changes to the internal mode. Some further AES measurements and investigations will be done on the influence of non-metallic like C, S, B, P, N on each other segregation.

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