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Influence of Oxidizing and Nitriding Parameters on Nitrogen Concentration of Electrical Steels

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Corresponding Author:	Sébastien Jégou Arts et Metiers Institute of Technology Aix-en-Provence, FRANCE
First Author:	Sébastien Jégou
Order of Authors:	Sébastien Jégou Mihaela Teodorescu Laurent Barrallier Fabrice Guittonneau
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Suggested Reviewers:	Thierry Czerwiec Thierry.czerwiec@univ-lorraine.fr Philippe Jacquet philippe.jacquet@ecam.fr Luc Pichon luc.pichon@univ-poitiers.fr Sebastien Dubent sebastien.dubent@lecnam.net
Response to Reviewers:	

Highlights

- The nitrated layer of electrical steels is coincident with the oxide layer
- The nitrogen content increases with the decrease of the nitriding temperature
- Oxidizing induces a redistribution of heavy elements (Si, Cr, Mn, Al)
- The nitrogen enrichment is driven by a ferrite to austenite transformation
- The austenite transformation is promoted by diffusing Si, O and N

Influence of Oxidizing and Nitriding Parameters on Nitrogen Concentration of Electrical Steels

S. Jégou¹, M. Teodorescu², L. Barrallier¹, F. Guittonneau¹

¹ Arts et Metiers Institute of Technology, MSMP, HESAM Université, F-13617 Aix-en-Provence, France

² thyssenkrupp Electrical Steel UGO, Isbergues, France

sebastien.jegou@ensam.eu, Mihaela.Teodorescu@thyssenkrupp.com, laurent.barralier@ensam.eu, fabrice.guittonneau@ensam.eu

Corresponding author: S.Jégou, sebastien.jegou@ensam.eu, Arts et Metiers Institute of Technology, MSMP, HESAM Université, F-13617 Aix-en-Provence, France

Abstract

The influence of oxidizing and nitriding parameters on the nitrogen concentration of grain-oriented electrical steels preliminary to the development of the final Goss texture was explored. Results show that the nitrogen enrichment is driven by a ferrite to austenite transformation during thermochemical treatments. Such a transformation is promoted by (i) a redistribution of ferrite-forming elements close to the surface during oxidizing prior to nitriding, (ii) the oxygen content within the oxide layer prior to nitriding, (iii) the temperature of oxidizing and nitriding, and (iv) the nitrogen enrichment during nitriding. Optimization of the nitrogen content, and thus the precipitation kinetics of alloying elements nitrides (e.g. inhibitors) required for the development of the final Goss texture can be controlled by an optimization of the oxide layer growth, the temperatures and gas mixture of nitriding.

Keywords

Electrical steel, oxidizing, nitriding, diffusion, phase transformations

1. Introduction

The evolution of environmental concerns in the last decay has been pushing electrical conversion systems to new developments and challenging optimizations. In case of electrical transformers, a complex process route leads to decrease core loss and increase the magnetic permeability [1-3]. A succession of hot and cold rolling, thermochemical surface and heat treatments have been optimized to manufacture so called “grain-oriented” (GO) electrical steels. The quality of the final crystallographic texture, e.g. Goss texture, directly drives the electrical performances. Goss texture, $\{110\}\langle 001\rangle$, develops due to an abnormal grain growth thanks to, at least, the presence of grain-growth inhibitors within the microstructure such as MnS precipitates and alloying elements nitrides MN (M = Si, Al,

43 Cr...) [4-8]. Optimizing such a complex process route appears time consuming, costly and complex,
44 especially because the magnetic properties can only be measured when the entire process route was
45 accomplished.

46 In case of MN inhibitors, three nitriding treatments have been optimized to promote such a
47 precipitation: (i) inherent nitriding from nitrogen available in the materials from casting [5], (ii)
48 equilibrium nitriding from reaction of dinitrogen N_2 during the last heat treatment during which the
49 Goss texture develops [10-11] and (iii) non-equilibrium nitriding from reaction of ammonia NH_3 in-
50 line after the oxidizing treatment [8-9]. Inherent and equilibrium nitriding do not promote a high
51 enough adsorption of nitrogen by the steel, leading to non-optimized fraction of inhibitors to promote
52 the Goss texture [9]. Development of the non-equilibrium nitriding treatment regarding the magnetic
53 properties has been reached identifying the acceptable range of nitrogen concentration for an efficient
54 abnormal grain growth, the corresponding temperature of nitriding, and the range of ammonia to
55 dihydrogen ratio to use in order to get satisfying precipitation of inhibitors [12-13]. **Prior to nitriding,**
56 **oxidizing aims decarburizing electrical steels as well as the later growth of an insulated layer (glass**
57 **film) [14-16].** Studies were carried out on the structure of the oxides' layer and on influence of process
58 parameters on oxides' layer properties. [17-18]. However, details about the nitriding kinetics, and
59 most of all on the influence of the oxide layer on nitriding kinetics, and thus on the nitrogen
60 enrichment and on control of the precipitation of inhibitors are missing in the literature [19-25].
61 The present work aims to study the influence of oxidizing and nitriding parameters on the nitrogen
62 adsorption and diffusion kinetics right before the development of the final Goss texture.

63

64 **2. Experimental**

65 **2.1. Materials**

66 The material is a silicon iron-based alloy used for grain-oriented electrical steel.
67 Samples of $30 \times 30 \times 0.2 \text{ mm}^3$ were provided by thyssenkrupp Electrical Steel for investigations.
68 Sensitive analysis of annealing parameters was performed through laboratory trails and experimental
69 measurements. **Conditions of oxidizing are chosen to promote both oxidizing and decarburizing in a**
70 **single step. The dew point (T_{DP} controlled by H_2O vapour and a chilled mirror hygrometer) and**
71 **temperature (T_O) of oxidizing** prior to nitriding stage were varied from 40 to 70°C and from 840°C to
72 900°C respectively. Samples have **then** been nitrided in a gas mixture using H_2 , N_2 and NH_3 , at various
73 temperatures (T_N), from 770°C to 930°C. **The nitriding potential (defined by the fraction of both NH_3**
74 **and H_2) was kept fixed to a usual low value used for electrical steels according to industrial practice**
75 **because of the complexity to control low nitriding potential in industrial conditions especially at such**
76 **high temperature (higher than 600 °C).** Cases without nitriding were as well considered in order to
77 dissociate the various parameters influences. The studied GO samples had the following main
78 chemical composition 3.1 wt.% Si, 0.028 wt.%Al, 0.15 wt.% Mn, 0.0095 wt.% N and almost no C, the
79 samples being already decarburised before the experimental measurements.

80

81 **2.2. Materials characterization**

82 The microstructure was observed by scanning electron microscopy (SEM) using a Jeol JSM-7001F
83 microscope and backscatter electrons mode. Prior to observations, ionic polishing (Ar⁺ ions, 4 kV)
84 was performed on each transversal cut using a Jeol Cross Section Polisher.

85 Chemical analysis was carried out by glow discharge optical emission spectroscopy (GDOES) using a
86 Profiler 2 device from Horiba. The surface of interest was around 4 mm in diameter parallel to the
87 surface of the sheet. The RF power, gas pressure and time of analysis were set to 35 W, 680 Pa and
88 300 s respectively, that corresponds to a depth of erosion of approximately 15 μm. Errors on
89 chemical analysis are expected to be less than 10 %, and were less than 2 and 1 % for nitrogen and
90 silicon respectively on a standard.

91 X-ray diffraction phase analyses were performed at room temperature using a Siemens D500
92 diffractometer, an Elphyse linear detector and the chromium $K\alpha$ radiation. The diffraction angle 2θ
93 was varied from 40 to 140°, that corresponds to a penetration depth of 2 to 6 μm in the ferritic matrix.

94

95 **2.3. Thermodynamics calculations**

96 Thermodynamics calculations were performed using Thermo-Calc version 2021a, TCFE Steels/Fe-
97 alloys database version 07 and TC-Python [26].

98

99 **3. Results**

100

101 **3.1. Microstructure observation**

102 **Figure 1** shows an example of the microstructure after oxidizing and nitriding observed by SEM using
103 the backscattered electron mode. The outer surface is characterized by an oxide layer of
104 approximately 2 μm thickness. The oxide layer is divided in two sublayers, one subscale at the
105 extreme surface of nearly 0.5 μm thickness composed of Fe₂SiO₄ and FeO oxides and one of
106 approximately 1.5 μm composed of globular followed by lamellar SiO₂ oxides [27]. All the samples
107 present similar type of composition, but their distribution, volume fraction and inner properties could
108 be different depending on the process parameters. The detailed analysis of the oxides' layer
109 components and their properties is not the main focus of this paper, the related results will be part of
110 another paper.

111

112 **3.2. Chemical analysis**

113 Oxygen and nitrogen in-depth profiles were obtained from GDOES analysis. **Figure 2** gives a
114 comparison of contents prior to thermochemical treatments, after oxidizing, and with and without gas
115 nitriding. The effective depth of oxygen diffusion is approximately 2.5 μm. The oxygen content is

116 maximum at the extreme surface and then decreases, following an internal oxidation diffusion
117 behaviour, to reach a null concentration at 2.5 μm below the surface in agreement with the SEM
118 observations given on **Figure 1** for this particular case. The diffusion of nitrogen atoms tends to make
119 the diffusion process of oxygen atoms to continue during nitriding. The nitrogen content in-depth
120 profile exhibits a non-obvious evolution as a function of the depth. It is maximum at the extreme
121 surface, decreases to a minimum of nearly 0.65 wt.% at a depth of 0.5 μm , increases up to 0.8 wt.% at
122 1.0 μm depth before abruptly decreasing toward the core content (below 0.03 wt.% according to the
123 sensitivity of present GDOES analysis). The effective diffusion depth of nitrogen atoms is in
124 agreements with the one of oxygen. **The nitrogen content in the oxide layer highly overestimates the
125 content within the final product that gives a good secondary recrystallization [12].**

126 The affected depth by oxygen atoms increases when the dew point (T_{DP}) and oxidizing (T_o)
127 temperature increase (**Figures 3 and 4**). The evolution of the content of oxygen as a function of the
128 depth are all similar, characterized by a maximum fraction close to the extreme surface that decreases
129 to a null content within the core. As previously, nitrogen atoms are largely located through the entire
130 oxide layer. The higher the temperature, the lower and deeper the nitrogen enrichment is, and, the
131 lower and deeper the minimum content of nitrogen is. It reaches 0.45 and 0.35 wt.% at a depth of
132 0.65 μm for a dew point and oxidizing temperatures of 70 and 900 $^{\circ}\text{C}$ respectively (**Figure 3.b and
133 4.b**). For low dew point (40 $^{\circ}\text{C}$), no variation of nitrogen content is observed within the oxide layer. In
134 contrast, a significant decrease of the nitrogen enrichment is noted when the oxidizing temperature is
135 increasing from 840 to 900 $^{\circ}\text{C}$.

136 **Figure 5** shows the influence of the nitriding temperature (T_N) on the case depth. No significant change
137 is observed in case of the oxide layer growth. The evolution of the nitrogen content shows similar
138 profiles for all temperatures with a minimum of the nitrogen fraction at a depth of nearly 0.5 μm
139 below the surface. However, increasing the nitriding temperature involves a significant decrease of the
140 nitrogen enrichment through the oxide layer. The fraction of nitrogen reaches almost 2.0 wt.% at
141 770 $^{\circ}\text{C}$ whereas 0.5 wt.% are barely achieved at 930 $^{\circ}\text{C}$.

142 **By considering the GDOES detection limit of nitrogen atoms (0.03 wt.%), a diffusion zone of nitrogen
143 in α -Fe ferrite below the oxide layer is observed to a depth of 12 μm . All the diffusion zones look
144 similar below the oxide layer with a nitrogen fraction ranging from 0.05 down to 0.03 wt.% except at
145 the low nitriding temperature of 770 $^{\circ}\text{C}$ in case of which the nitrogen fraction ranges from 0.075 down
146 to 0.03 wt.% (Figure 5.b).**

147 One objective of oxidizing prior to nitriding is to decarburize the sample in order to achieve a
148 minimum content required to maximize the abnormal secondary recrystallisation. **Figure 6** gives
149 examples of the carbon content in-depth profiles. It is observed that the decarburization of the core
150 material is effective (0.04 down to 0.02 wt.%) whereas it induces a non-negligible enrichment of
151 carbon in the oxide layer for all samples. **According to the affected depth, such enrichment may be
152 attributed to the presence of $(\text{Fe},\text{Mn})_2\text{SiO}_4$ and FeO close to the surface in the oxide layer (Figure 1).**

153 Despite the GDOES detection limit of carbon (from 0.003 to 0.01 wt.%), these measurements must be
154 considered with precautions because of possible pollution of the surface due to cleaning for instance.
155 More investigations are required to address the carbon enrichment of the oxide layer during oxidizing.
156 The chemical composition of some alloying elements is given on Figure 7 and 8 as a function of the
157 depth below the surface and the temperatures of oxidizing and nitriding respectively. The same
158 complex tendency as a function of the depth below the surface is observed for all elements. All
159 elements are characterized by a depletion close to the outer surface, an increase of the content
160 followed by a depletion at greater depth. The redistribution of alloying elements is observed at more
161 profound depth when the thickness on the oxide layer increases (Figures 3 and 4). The silicon content
162 is almost constant within the oxide layer and higher than in the core, but some profile variations are
163 noted mostly induced by the change in dew point. Such redistribution of alloying elements through the
164 oxidized surface is at least influenced by the oxidizing treatments (Figure 7) and nitriding does not
165 modify them (Figure 8).
166 Finally, the total nitrogen uptake into the oxide layer can significantly vary by a factor 4 when
167 increasing the nitriding temperature from 770 to 930 °C (Figure 5) whereas no significant change in
168 the redistribution of carbon nor heavy elements is observed (Figures 6 and 8).

169

170 3.3. X-ray diffraction

171 Figure 9 gives the phase analyses carried out at the surface of samples as a function of the nitriding
172 temperature, as well as after decarburizing. Austenite is identified after nitriding but not present prior
173 to nitriding, or in a too low volume fraction or crystallite size. The lower the nitriding temperature, the
174 higher the volume fraction of austenite is. To complement these results, a tempering was performed,
175 subsequent to nitriding, for one minute at the nitriding temperatures tested previously and followed by
176 water quenching. The tempered samples confirm the presence of retained austenite at room
177 temperature after nitriding.

178

179 3.4. Thermodynamic calculations

180 Figure 10 gives the Fe-Si-N-2.0wt.%O isopleth diagrams at 750 and 950 °C and possible phase
181 transformations at a depth of nearly 0.1 µm. Results show that a ferrite to austenite transformation
182 may occur during oxidizing and nitriding. It is due to the depletion of silicon but also to the
183 enrichment of both oxygen and nitrogen. As a consequence, the transformation is expected to be
184 promoted close to the outer surface according to GDOES analysis. Moreover the transformation is
185 thermally activated enabling the presence of austenite for higher Si content and thus deeper within the
186 oxide layer.

187 Figure 11 gives the thermodynamic calculations as a function of the chemical composition of samples
188 after nitriding at three different temperatures as measured by GDOES. Results show that austenite is
189 present in the oxy-nitrided layer as deep as 1.0 µm. At 770 °C, the nitrogen enrichment significantly

190 stabilizes the austenitic phase as compared to without nitriding at the same temperature between 0.2
191 and 1.0 μm in depth (Figure 11.c).

192 The fraction of silicon nitrides within the oxi-nitrided layer increases when the temperature of
193 nitriding decreases and must be related to an increase of the nitrogen uptake.

194 The fraction of oxides does not depend on the nitrogen content and stay nearly unchanged when
195 comparing samples with and without nitriding. In addition, all calculations show that the nature of the
196 silicon oxide may change according to the temperature of both oxidizing and nitriding. Below 850 $^{\circ}\text{C}$,
197 silicon oxides are quartz like but change to tridymite at higher temperature.

198

199 4. Discussion

200 Chemical characterizations of a GO electrical steel after oxidizing and nitriding show that the kinetics
201 of nitrogen adsorption and diffusion highly depends on the oxide layer growth, on the oxygen and
202 carbon content, on the complex redistribution of heavy elements (such as Si, Cr, Al, Mn), as well as on
203 the temperature of the heat treatments. The nitrogen content in-depth profile exhibits a non-obvious
204 evolution as a function of the depth, with a minimum of nitrogen within the oxide layer.

205 The higher the temperatures of oxidizing and nitriding, the deeper the diffusion of nitrogen and the
206 lower the maximum of nitrogen in the oxide layer are (Figures 4 and 5 respectively). Nitrogen
207 diffusion is characterized, as mentioned, by a minimum fraction of nitrogen within the oxide layer
208 (close to a depth of 0.5 μm), that is even more pronounced when the oxide layer is thick. The nitrogen
209 uptake does not appear depending in first approximation on the fraction and redistribution of elements
210 such as silicon, chromium, aluminium or manganese (Figures 7 and 8). However this observation
211 reflects the fact that, prior to nitriding, a similar redistribution of elements between oxides and matrix
212 is obtained, for the range of the variations of the decarburizing (oxidizing) temperature considered.
213 But any change in the silicon or aluminium redistribution between the matrix and oxides during
214 oxidizing is obviously expected to play a role on the formation of silicon and aluminium nitrides and
215 thus on the level of the nitrogen uptake during nitriding.

216 Chemical analyses revealed that oxidizing leads to a decrease of the carbon content within the core
217 material thanks to the decarburizing atmosphere during oxidizing, but results in a carbon enrichment
218 of the oxide layer prior to nitriding (Figure 6). Heavy elements also exhibit a complex redistribution
219 through the oxide layer prior to nitriding (Figures 7 and 8), consistent with the literature using the
220 same chemical analysis method [27]. These redistributions of already present elements are influenced
221 by the conditions of oxidizing and not affected by those of nitriding.

222 Thermodynamic calculations of temperature isopleths for given oxygen content or using compositions
223 as measured by GDOES show that a ferrite to austenite transformation may occur within the oxide
224 layer during nitriding (Figure 10). The higher the oxygen content and temperature, the lower the
225 transformation point from ferrite to austenite is during nitriding. The ferrite to austenite transformation
226 is as a consequence promoted close to the surface in the oxide layer. Although the carbon fraction

227 measurement may need optimization for GDOES analysis, the presence of carbon content within the
228 oxide layer can promote such an austenitic transformation. Calculations confirm the present
229 experimental observations of retained austenite by XRD at the surface of samples (Figure 9).
230 Austenite was also observed experimentally at the extreme surface of the oxide layer in other recent
231 works [16, 25]. The fraction of austenite increases in agreement with the nitrogen content according to
232 GDOES analysis and the Fe-N phase diagram [28]. Iron nitrides being more thermodynamically stable
233 than austenite at room temperature [28], tempering after nitriding must promote the decomposition of
234 austenite into ferrite and iron nitrides. It thus leads to no observation of austenite at room temperature
235 by XRD. However iron nitrides were not observed by XRD that suggests a low fraction of iron
236 nitrides or too small crystallites. More investigations are required.

237 The observation of a minimum of the nitrogen fraction in the oxide layer close to 0.5 μm below the
238 surface is in agreement with the transformation zone from ferrite to austenite according to
239 thermodynamic calculations (Figure 11).

240 The thermodynamic calculations let us assume that the austenitic transformation can be in competition
241 with the precipitation of silicon nitrides, and so, even more when the transformation point is low
242 (Figure 10). Although the nitrogen solubility limit in austenite is much higher than in ferrite, the
243 precipitation kinetics of alloying elements as nitrides, and so the nitrogen uptake, may be assumed
244 slower if the austenitic transformation occurs before the complete precipitation of nitrides. Moreover,
245 it has to be considered that the diffusion kinetics of nitrogen atoms in austenite is at least one hundred
246 times lower than in a ferritic matrix [29]. It thus leads to a decrease of the flow of nitrogen atoms at
247 the beginning of the oxide layer whereas the diffusion kinetics are much faster in the ferritic area at
248 greater depth.

249 As a consequence, the calculated gradient of austenite close to the surface is expected to result in a
250 depletion of nitrogen between the low diffusion kinetic area near the surface and the fast diffusion
251 kinetics area (ferrite zone) deeper in the oxide layer. The nitrogen enrichment being slower due to the
252 austenitic transformation prior to the complete precipitation of silicon nitrides at the surface, the
253 boundary conditions for diffusion in the ferritic zone deeper in the material must also change towards
254 lower value. Therefore, it induces a lower level of nitrogen in ferrite and a slower precipitation
255 kinetics of alloying elements as nitrides deeper in the oxide layer.

256 According to thermodynamic calculations and the higher chemical affinity of both silicon and
257 aluminium with oxygen than nitrogen [30], the oxides already present before nitriding are supposed
258 not to take part in the precipitation of silicon and aluminium nitrides.

259 Calculations show that the nature of silicon oxides could also change during nitriding function of the
260 content of both oxygen and silicon, but also depending on the temperature of oxidizing and nitriding.
261 Similarly, the nature and fraction of iron-silicon oxide at the outer surface (e.g. $(\text{Fe},\text{Mn})_2\text{SiO}_4$ and/or
262 FeO) may play a role on the catalytic dissociation of ammonia and the nitrogen adsorption and uptake.

263 However present experimental results cannot conclude on any effect of such change of oxides on the
264 diffusion kinetics of nitrogen atoms during nitriding.

265 Finally, it is well known that the natural dissociation rate of ammonia NH_3 into N_2 and H_2 gas mixture
266 increases exponentially with the temperature [31]. As a result, and considering perfect gas transfer to
267 the solid surface, the nitrogen fraction at the gas-solid interface available for adsorption into the solid
268 from the catalytic dissociation of ammonia at the solid surface naturally decreases and thus the
269 nitrogen enrichment within the oxide layer. Moreover, at the atmospheric pressure, the dissociation
270 rate of dinitrogen molecules N_2 during nitriding cannot compensate the lack of nitrogen atoms from
271 the catalytic ammonia dissociation at the solid surface without increasing the pressure [31-33].

272

273 **5. Conclusion**

274 Influences of the conditions of oxidizing and nitriding on the nitrogen enrichment prior to the
275 development of the final Goss texture were explored in case of a grain-oriented electrical steel. The
276 nitrogen enrichment in the oxide layer can significantly overestimate the nitrogen content required for
277 the development of the final Goss texture. The nitrogen atoms are exclusively located within the about
278 $2.5 \mu\text{m}$ thick oxide layer after nitriding. The nitrogen diffusion kinetics through the oxide layer
279 depends on a ferrite to austenite transformation during oxidizing and nitriding. The transformation
280 point decreases with (i) the increase of the fraction of oxygen, carbon and nitrogen atoms, (ii) the
281 depletion of heavy elements very close to the outer surface and (iii) the temperature increase.

282 The nitrogen content, that leads to the optimized precipitation of inhibitors required for the
283 development of the final Goss texture and the optimum magnetic properties, is driven by the **oxidizing**
284 and nitriding conditions that must be involving a ferrite to austenite transformation if not controlled.
285 The nitrogen enrichment, and thus the development of inhibitors, can obviously be optimized and
286 controlled depending on the conditions of oxidizing and nitriding. **It arises that optimize the nitriding**
287 **potential through the gas mixture ($\text{NH}_3\text{-N}_2\text{-H}_2$) could also be necessary in order to get the most**
288 **efficient thermochemical treatments conditions of electrical steels.**

289

290

291 **Data availability**

292 The raw/processed data required to reproduce these findings cannot be shared at this time as
293 the data also forms part of an ongoing study

294

295

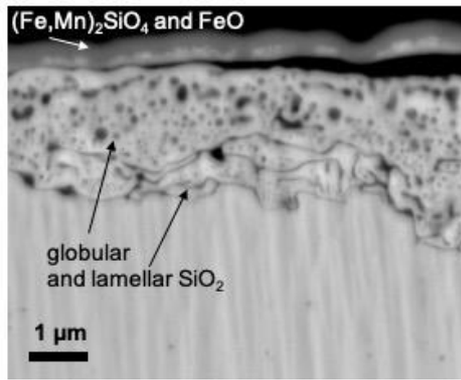
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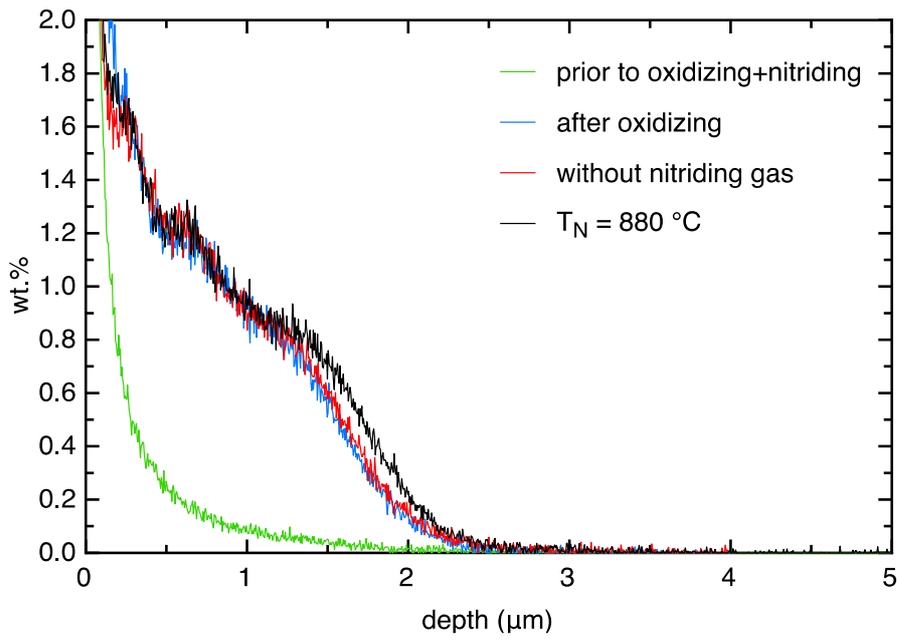
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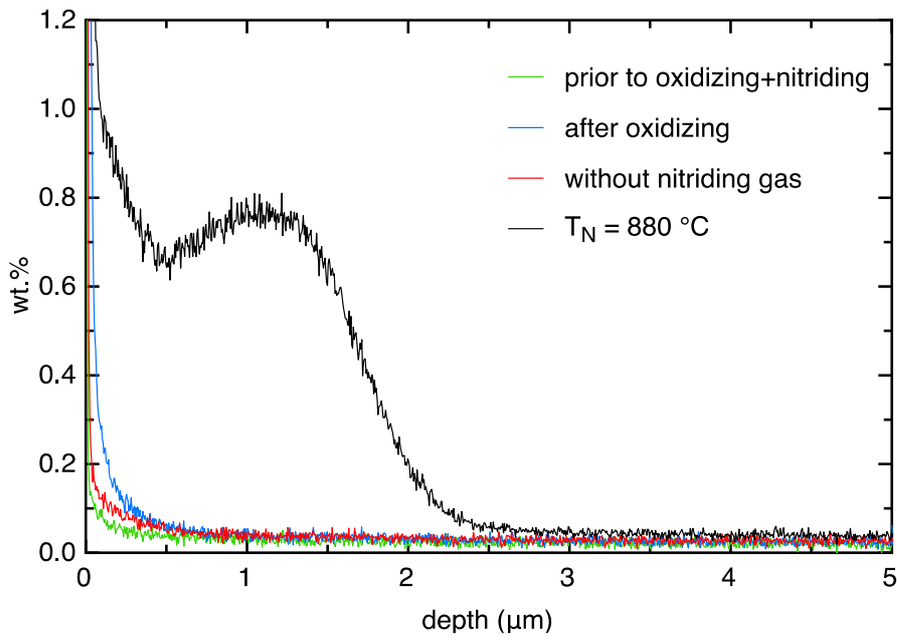
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Figure 1: SEM micrography in backscattered mode of a GO electrical steel after oxidizing ($T_{DP} = 60$ °C, $T_O = 870$ °C) and nitriding ($T_N = 880$ °C).



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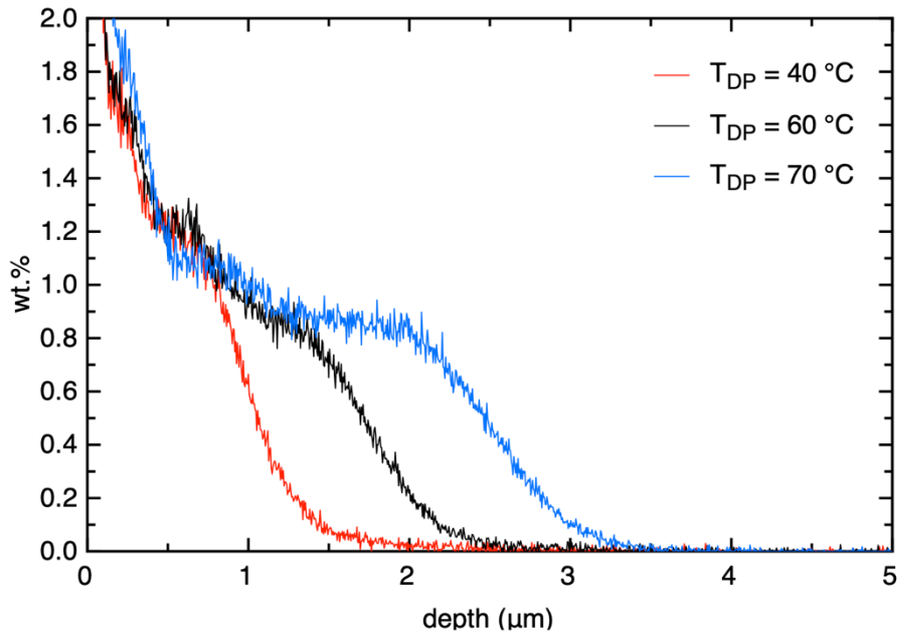
a. Oxygen



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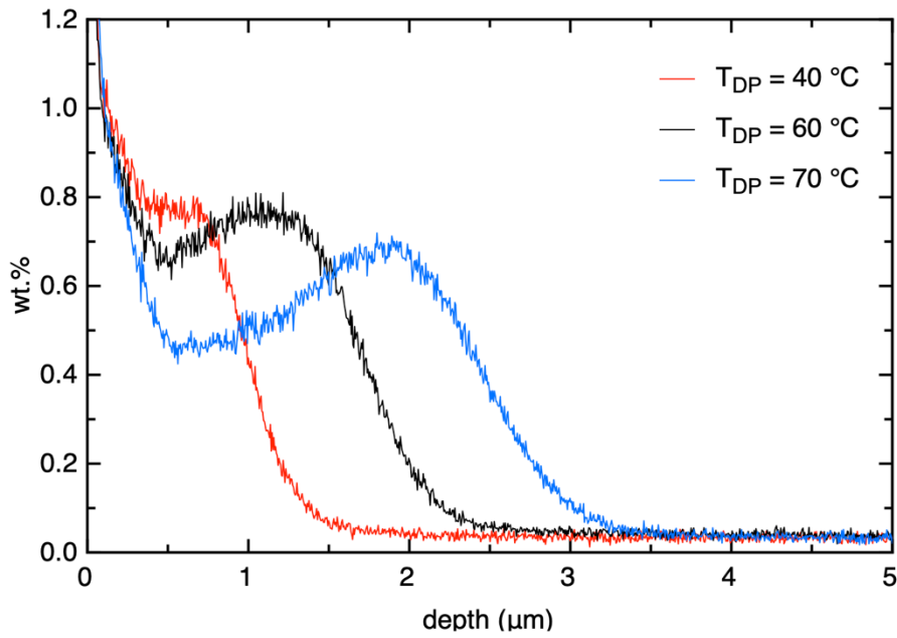
b. Nitrogen

Figure 2: GDOES chemical analyses of GO electrical steel prior to oxidizing and nitriding, after oxidizing, and with and without gas nitriding. (a) oxygen and (b) nitrogen in-depth content profiles.



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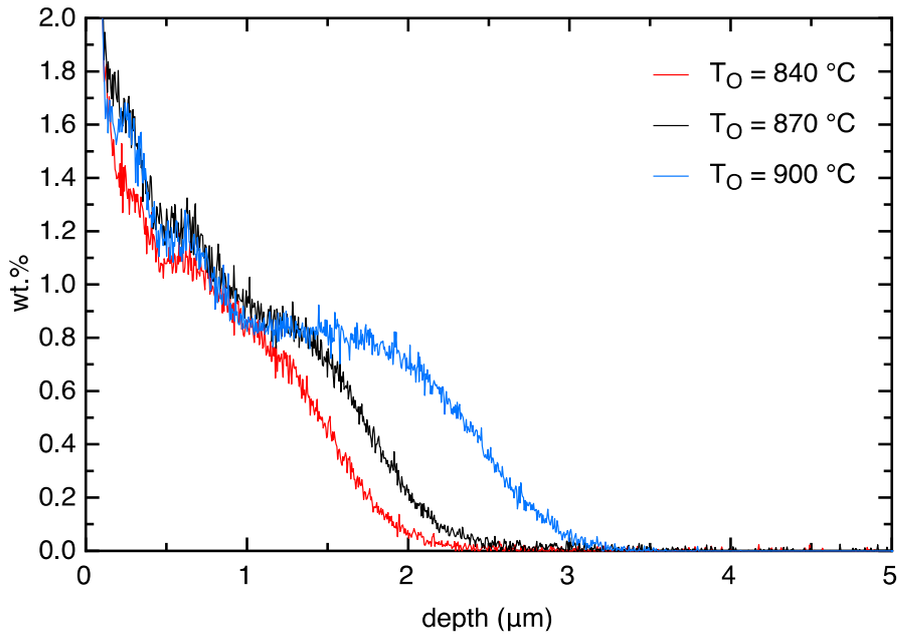
a. Oxygen



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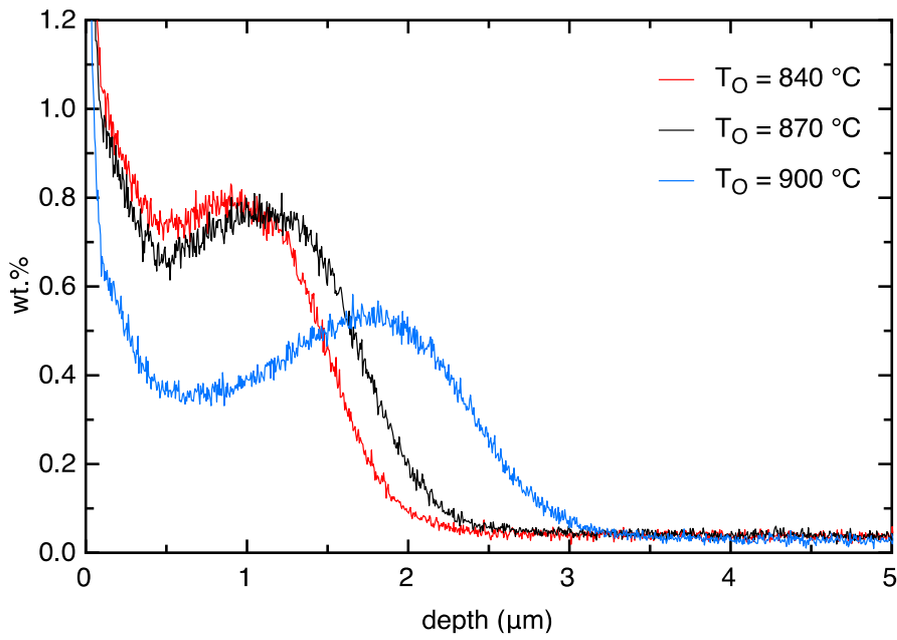
b. Nitrogen

Figure 3: GDOES chemical analyses of GO electrical steel. Influence of the dew point (T_{DP}) of oxidizing on (a) oxygen and (b) nitrogen in-depth content profiles.



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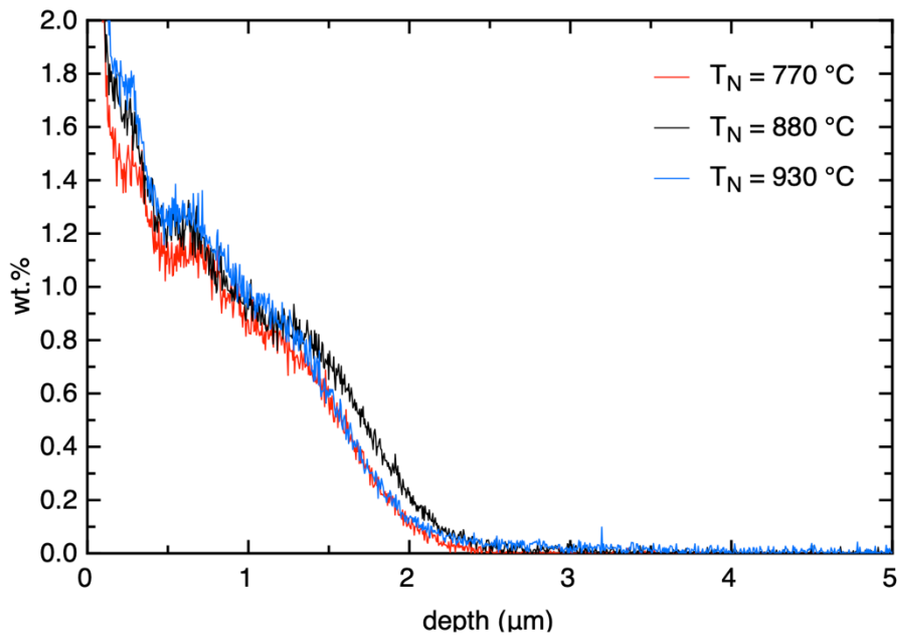
a. Oxygen



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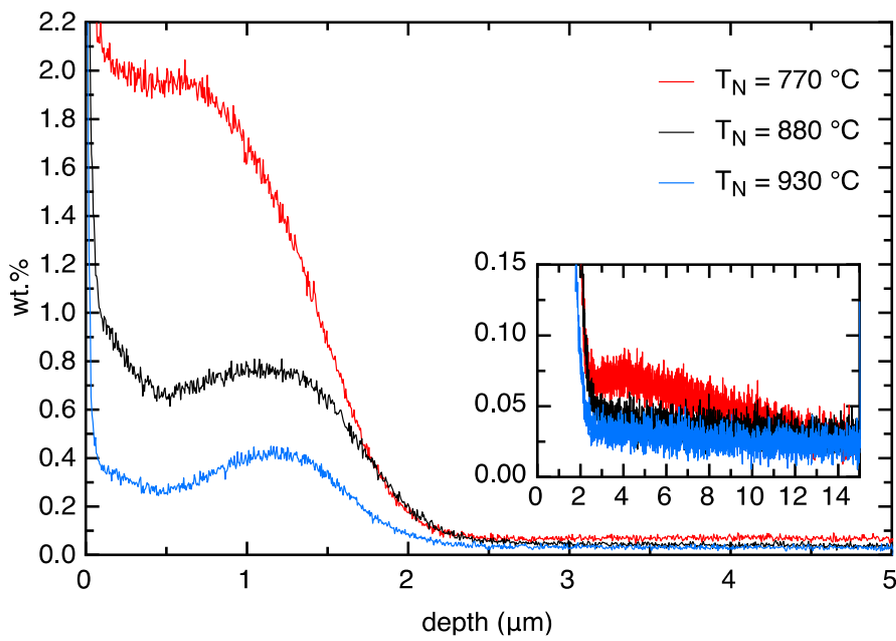
b. Nitrogen

Figure 4: GDOES chemical analyses of GO electrical steel. Influence of the oxidizing temperature (T_O) on (a) oxygen and (b) nitrogen in-depth content profiles.



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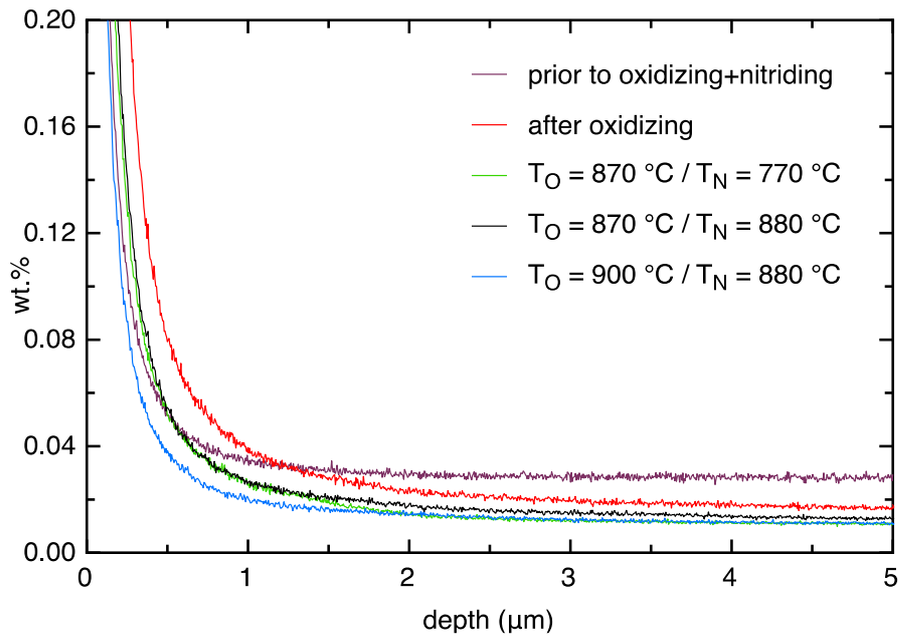
a. Oxygen



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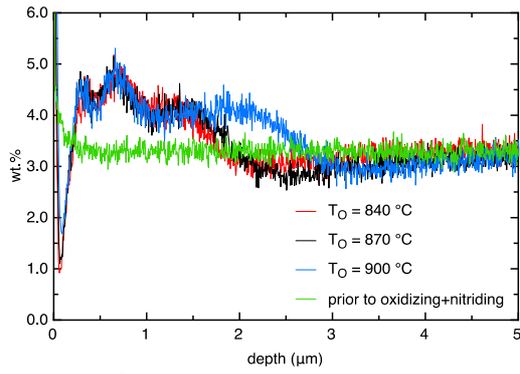
b. Nitrogen

Figure 5: GDOES chemical analyses of GO electrical steel. Influence of the nitriding temperature (T_N) on (a) oxygen and (b) nitrogen in-depth content profiles.



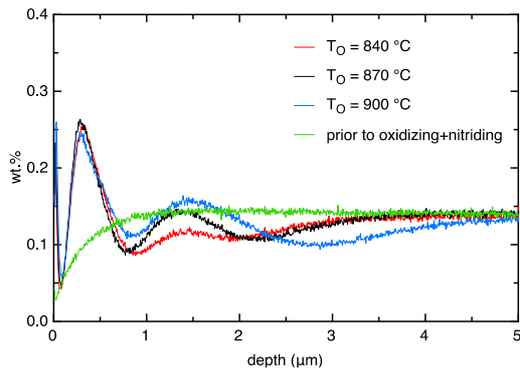
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Figure 6: Carbon in-depth profiles from GDOES chemical analyses of GO electrical steel. Comparison of the carbon content at different steps of thermochemical treatments and influence of the temperature of oxidizing (T_O) and nitriding (T_N).



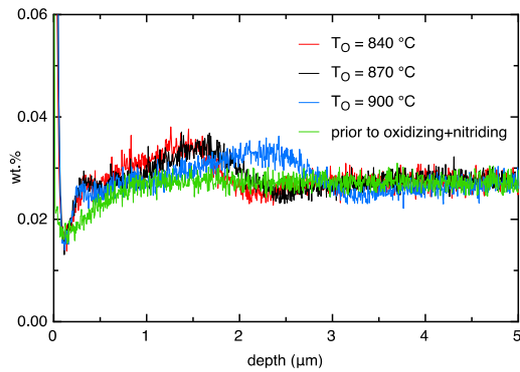
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a. Silicon



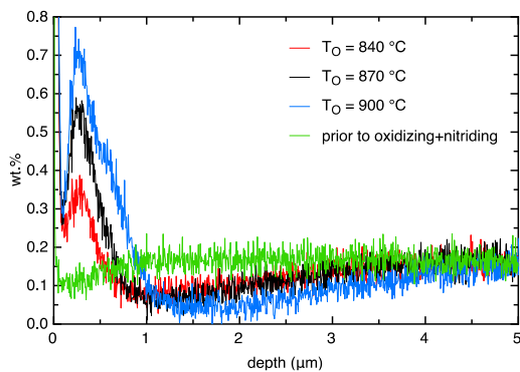
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b. Chromium



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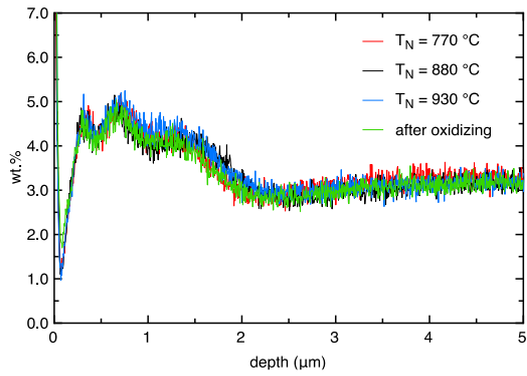
c. Aluminium



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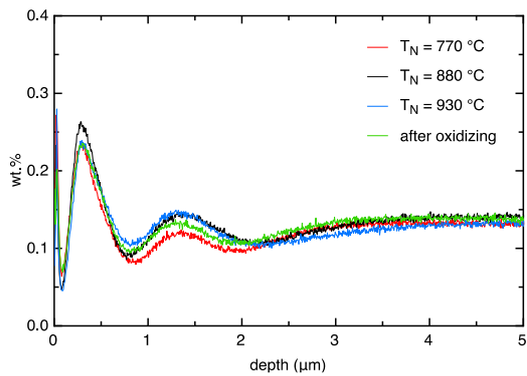
d. Manganese

67 **Figure 7:** GDOES chemical analyses of GO electrical steel. Influence of the oxidizing temperature
68 (T_O) on (a) silicon, (b) chromium, (c) aluminium and (d) manganese in-depth content profiles.



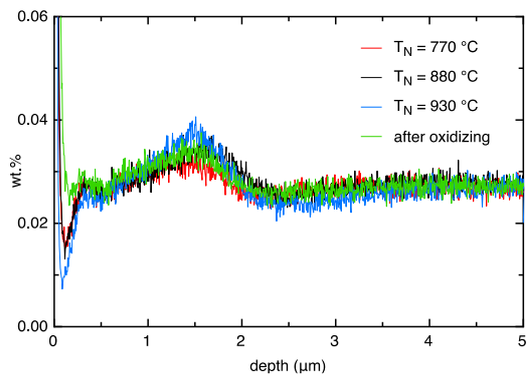
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a. Silicon



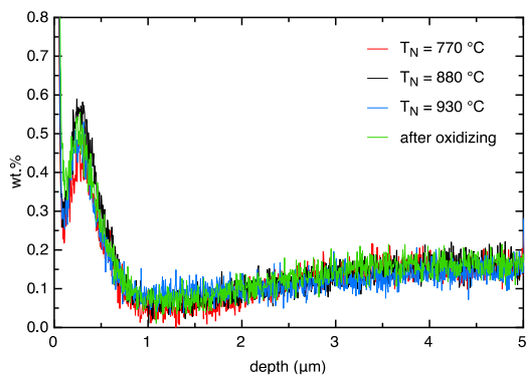
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b. Chromium



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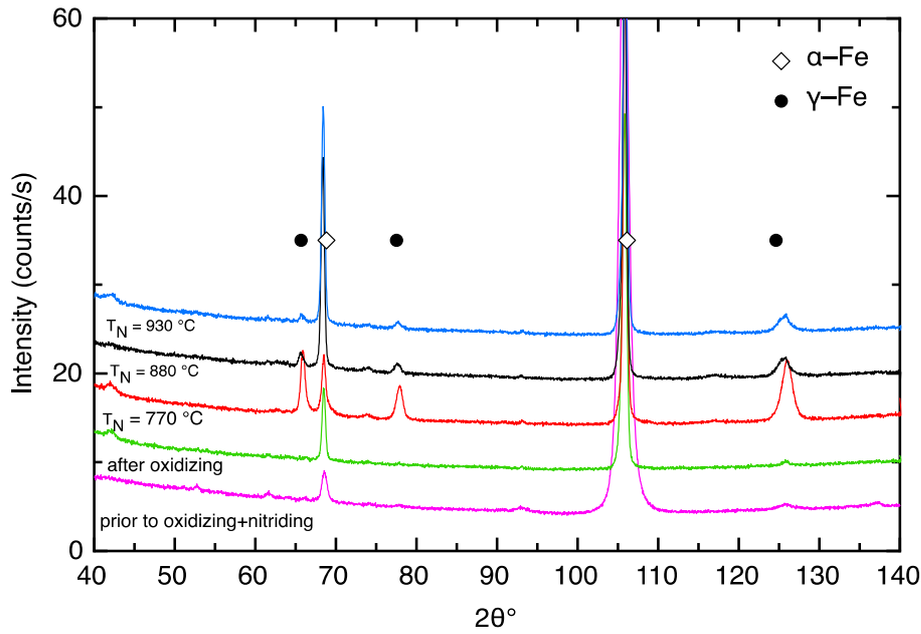
c. Aluminium



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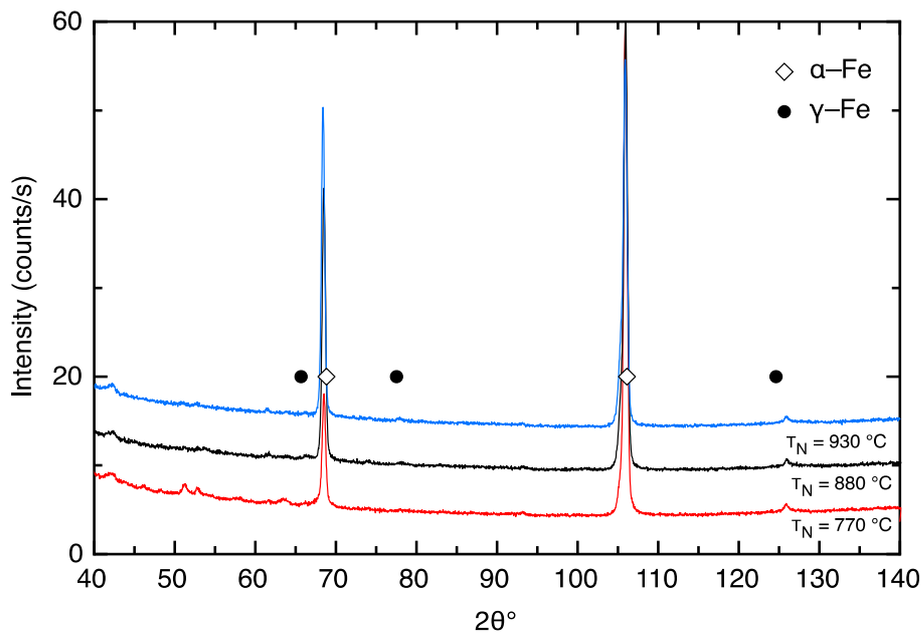
d. Manganese

81 **Figure 8:** GDOES chemical analyses of GO electrical steel. Influence of the nitriding temperature (T_N)
82 on (a) silicon, (b) chromium, (c) aluminium and (d) manganese in-depth content profiles.



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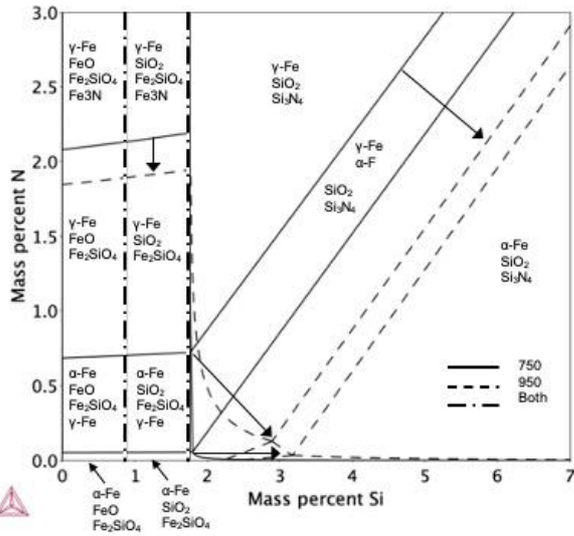
a. After nitriding



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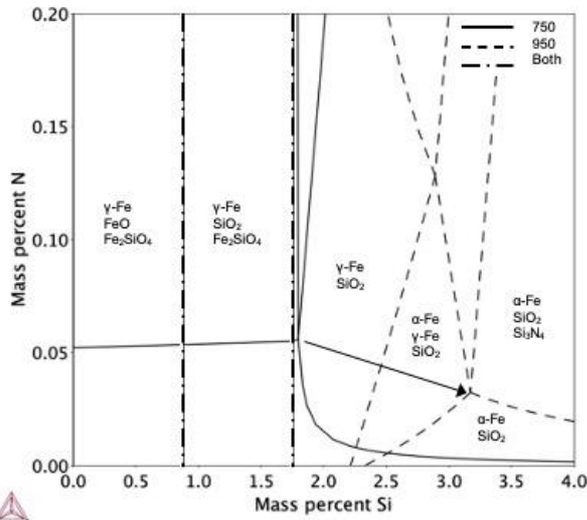
b. Tempering after nitriding

Figure 9: XRD phase analyses of GO electrical steel. Influence of the (a) nitriding temperature (T_N) and (b) tempering after nitriding.



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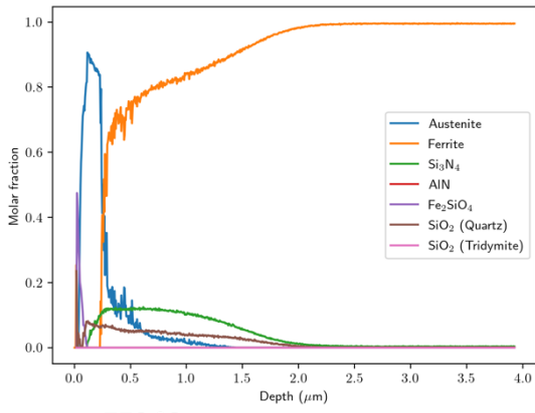
a.



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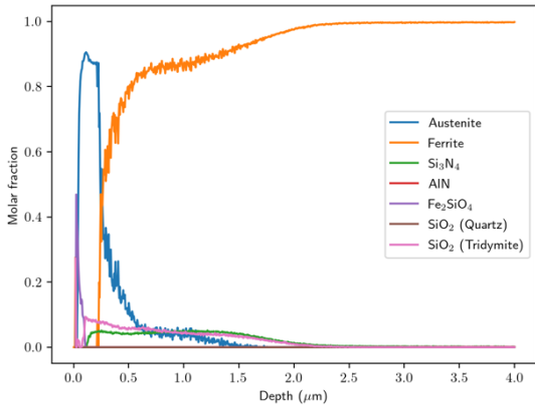
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Figure 10: Isoleth diagrams of Fe-Si-N-2.0wt.%O at 750 and 950 °C. (a) phase fields given for 750 °C. (b) phase fields given for 950 °C.



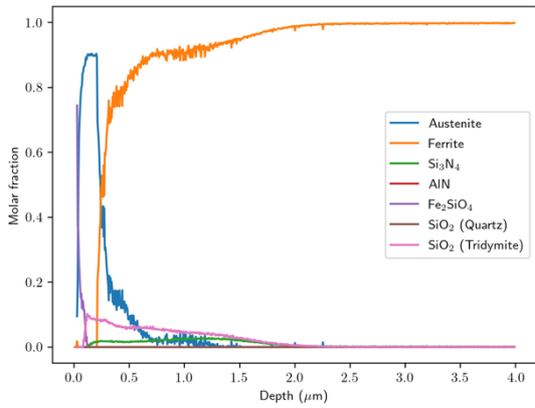
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a. 770 °C



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b. 880 °C



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c. 930 °C

112 **Figure 11:** Calculations of the phase fractions as a function of the nitrogen content in an electrical steel
113 according to GDOES analysis of the nitrated sample at (a) 770, (b) 880 and (c) 900 °C.
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