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Investigation of the ASPN process of low alloy steel by using Ni or Cr coated active screens



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ABSTRACT

The mechanism of active screen plasma nitriding (ASPN) and the influence of the active screen's coating material on the surface properties of the nitrided layer was investigated on low alloyed steel samples, using chromium or nickel-coated active screens. ASPN experiments were performed at 510 °C, for 5 h in a 75% N₂ + 25% H₂ gas mixture using tempered 42CrMo4 type low alloy steel. The base material of the screens was unalloyed steel, but the screens were electroplated with chromium or nickel. Scanning electron microscopy (SEM), energy dispersive spectroscopy (EDS), X-ray diffractometry (XRD), atomic force microscopy (AFM), and secondary neutral mass spectrometry (SNMS) were used for the characterization of surface properties of the samples and the screen. SNMS results show that the coating material of the active screen (chromium or nickel) was deposited on the surface of the samples in the form of compounds, with layer thicknesses between 160 and 200 nm (chromium) and 290-420 nm (nickel). Chemical analysis by X-ray photoelectron spectroscopy (XPS), performed at the interface of the steel sample and the deposited compounds reveals that iron-nitride was only formed in the case of samples that were treated with a Cr-coated active screen. Here the decomposition and oxidation of the unstable chromium-nitride provided the nitrogen atoms and the chemical drive for the formation of iron-nitride. In the case of the Ni-coated active screen, only molecular nitrogen was observed in the treated steel samples. The results prove that - contrary to widespread models on ASPN nitriding mechanisms - physically adsorbed nitrogen plays little role in the nitriding process of iron.

1. Introduction

Tempered low alloy steels are often used for gears and shafts. These parts are usually exposed to wear and, in some cases, also to corrosion. The effective way to improve the wear and corrosion resistance of these components is to modify the surface properties by the proper constitution of the diffusion layer and coating [1]. Different strategies have been utilized to improve the characteristics of the coating materials, including multilayer structures that are generated by depositing two or more alternative materials [2,3].

Active screen plasma nitriding (ASPN) is a widely used surface treatment process, which offers many advantages compared to the conventional direct current plasma nitriding, such as better layer uniformity and the elimination of the edge effects [4,5]. In this process, the workpiece is surrounded by a conductive mesh screen (also called a

cage), which is on cathodic potential, while the furnace wall is kept on anodic potential [6,7]. In ASPN, the sample is usually either insulated from the high voltage source (and thus on a floating potential) or on a lower cathodic bias compared to the screen. Consequently, the ions of the plasma bombard and sputter the sample to a smaller extent compared to the screen. The sample is heated by radiation [8,9] and necessarily by the plasma as well.

The surface hardening of steel specimens originates from the formation of a nitride compound layer on the surface and a nitrogen diffusion zone underneath it. The nitriding mechanisms used to describe the ASPN process differ significantly from the "sputtering of iron and re-condensation of iron-nitride" model of Köbel used for DC plasma nitriding [10]. In the first step of ASPN, according to the model offered by Zhao, the unalloyed steel surface of the active screen is covered by a thick compound layer containing $Fe_{2-3}N$, Fe_4N , in a mechanism similar

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to DCPN [11,12]. Consequently, iron-nitride particles will be sputtered from the screen, which passes through the plasma where they physically and chemically adsorb the active nitrogen atoms before deposited on the surface of the steel sample [11,13,14]. In the model of Zhao, the formation of iron-nitride on the surface of the sample is driven by two processes. First, the physically adsorbed nitrogen atoms desorb on the surface due to the contact between the particles and the substrate surface and the high substrate temperature. The released nitrogen can either form molecular nitrogen and diffuse into the sample or form ironnitride. Secondly, the decomposition of higher iron-nitrides, like the metastable FeN and $Fe_{2-3}N$, release nitrogen, which can form further nitrides. In this modified model of "sputtering-adsorption-desorption," Zhao clearly considers the sputtered particles as the main nitrogen carriers in the active screen plasma nitriding process. The validity of this model can be tested by changing the material of the active screen: regardless of the screen's material, the sputtered metal atoms and metal-nitrides carry physically and chemically absorbed nitrogen to the sample surface, which can catalyze the nitriding of iron. Thus, one of the aims of our work is to test the validity of this model and investigate the effect of the active screen's material on the ion nitriding process. Besides, the deposited materials originating from the active screen (metals and nitrides) also modify the properties of the steel surface [15,16]. Generally – as investigated by others – the materials of both the screen and the sample influence the resulting surface properties in a complex way.

Naeem et al. [17] examined the distance between the active screen and the sample in which austenitic stainless steel screen was used to treat carbon steel samples. After the treatment, they measured increased chromium and nickel content on the surface of the samples. This observation has inspired new research in which stainless steel and aluminum screens were used one after another to improve the surface properties. Aluminum nitride was found to be the main phase in samples nitrided in this fashion, with and the hardness being much higher (1100 HV) than in samples treated conventionally (800 HV) [18]. Nishimoto et al. [19] used a titanium screen for nitriding high-alloyed steel. A TiN layer was formed on the surface by sputtering from the screen and by deposition of the nitrides.

In this manner, both the material of the active screen and sample can be examined. Yazdani et al. [20] nitrided aluminum sample with carbon steel screen. An iron-nitride layer was formed on the top of the sample surface. The authors stated that the nitrogen atoms diffused into the Al substrate and formed an AlN layer beneath the iron nitride coating at long coating periods, although this was not detected by XRD. Taherkhani and Soltanieh [21] also used an iron screen on aluminum samples. The influence of nitriding time and temperature on the surface properties of coated aluminum alloy samples was investigated. A combination of AlN and $Fe_{2-3}N$ phases were shown to form by the ASPN process. The authors call this a composite nitride layer. If the nitriding time is long enough (≥ 11 h), a number of iron nitride particles can be sputtered from the screen to be deposited on the surface, which will result in a desirably thicker coating with larger grain size [22].

EDS (energy dispersive spectrometry) analysis was preferably used for the determination of nitrogen content [17,23,24]. Besides this measurement, the GDOES (glow discharge optical spectrometry) is the most common depth profiling technique to analyze the chemical composition of the layer [25–27]. Both the nitride layer of structural steel and the expanded austenite layer of the austenitic stainless steels can be characterized by this method. However, the nitrogen content of the diffusion zone cannot be analyzed by any measurement techniques. Hosseini and Ashrafizadeh [28] were compared three types of compositional depth profile analyzer techniques. Nitrogen is a light element with a relatively low concentration in a plasma nitrided layer; thus, it is difficult to be measured if only one of the conventional surface analysis techniques is employed. A 6 μ m thick layer was presented in this study. In this case, EDS provided more accurate values for the N content than GDOES, since the latter predicted an only $4.5 \,\mu$ m thick layer. It should be noted that GDOES does not give direct distance information; therefore, it needs a very precise correspondence of the sputtering time to sputtering depth. SIMS (secondary ion mass spectrometry) was used to determine the N content in the diffusion zone. Quantitative results were not presented, only the profile of the measurements, but the results were validated by N content calculation [29]. SNMS (secondary neutral mass spectrometry) was rarely used to investigate the nitrided layer [30]. Galesic and Kolbesen [31] showed the changes of V and N in the function of the sputtering time, but the depth was not calculated. In contrast, Jauberteau et al. [32] found MoO₂, MoO₃ compounds after the nitriding, but N also could diffuse into the metal in a hundred nm, which can be seen in the SNMS profile.

Conclusively, the aim of our research is to investigate the nitriding process of steel and the chemical composition of the formed nitride layer (with SNMS and XPS), using nickel or chromium coated screens for ASPN, with relation to the widely used nitriding models. Furthermore, the self-nitriding of the coated active screen was also investigated. Information regarding the processes on the active screen's surface during plasma nitriding cannot be found in the literature.

2. Material and methods

2.1. Sample preparation

Tempered 42CrMo4 grade steel was used as a substrate material in this study. The material was cut into disks with a 20 mm diameter and 6 mm thickness. The samples were wet ground by using silicon carbide papers of multiple grit sizes and later polished by diamond suspension, then degreased in an ultrasonic vibrated acetone bath and dried with air directly before the nitriding process.



Fig. 1. Illustration of the used plasma nitriding equipment.

Table 1

Designation and details of the various ASPN treatments. One screen (either nickel or chromium coated) was used for two samples consecutively; this is referred to as the first and second ASPN in the table.

Designation Treatment	
PNi1x First ASPN with the ni PNi2x Second ASPN with the PCr1x First ASPN with the ch PCr2x Second ASPN with the	ckel coated screen nickel coated screen romium coated screen chromium coated screen



Fig. 2. SEM images of the screens' surface and corresponding EDS spectra. The screens were used a) 0 time, b) 1 time, c) 2 times with Cr-coated screen, and d) 0 time, e) 1 time, f) 2 times with Ni-coated screen.

2.2. Plasma nitriding

The base material of the screens was made from a 1.0330 type nonalloyed quality steel. The dimensions of the screen were \emptyset 140 × 85 × 0.8 mm. Two coating materials were electroplated in an industrial bath: nickel and chromium. For the chromium coating, chromic acid, for the nickel coating Watts-electrolyte were used according to well-known protocols [33,34].

The samples were nitrided for 5 h at 510 °C using 75% $N_2 + 25\%$ H₂ gas mixture. The gas pressure was kept constant at 280 Pa. The sample was placed at the center of the active screen, 35 mm from the top lid. The temperature was monitored by using an isolated K-type thermocouple mounted directly under the workpiece. The illustration of the equipment can be seen in Fig. 1. After the nitriding, the samples were cooled down under the operating gas to 250 °C. Afterwards; they were cooled down to room temperature when the chamber was pressurized with air.

All treatments were carried out by using the same plasma nitriding parameters. One screen was used for two different specimens as an active screen. Both the diffusion of the coating material into the substrate and the changing of the screen's surface were investigated. Table 1 shows the experimental treatments which were used on the samples and their designation.

2.3. Characterization methods

The surfaces of the nitrided samples were analyzed by XRD with theta-2 theta vertical goniometer and Cu target. The scan parameters were: 40–80° in 2 theta, the wavelength was 1.54 nm, while the exposure time was set to 0.04 s/step. Surface morphology, the cross-section, and the elemental composition of the screen surface were investigated by using a Zeiss EVO MA10 scanning electron microscope along with EDS. Hardness was measured by a Buehler IndentaMet 1105 type Vickers microhardness tester by using 10 g load. To measure the surface morphology, a Veeco diInnova type atomic force microscope (AFM) was used in contact-mode with an ART D160 diamond probe (spring constant: 5 N/m). The images were obtained in $10 \,\mu\text{m} \times 10 \,\mu\text{m}$ scan sizes, with a sampling resolution of 512×512 and 1 Hz scan rate. The obtained images were post-processed with the Gwyddion 2.36 software [35].



In SNMS analysis, the sample surface was bombarded by ions, and the sputtered neutral particles were detected after ionization, i.e., the sputtering and ionization were strictly separated from each other [36]. The main characteristic feature of SNMS and its advantage is the separation between emission and ionization of particles, which results in a matrix-independent method. The surface bombardment and post ionization were performed by low-pressure Electron Cyclotron Wave Resonance (ECWR) argon plasma. Due to the presence of a surface oxide layer on the samples, the SNMS was operated in a high-frequency mode, which ensured an effective charge compensation against surface charge accumulation resulting in inhomogeneous ion bombardment. A sputtering potential of 350 V at 100 kHz frequency with 80% duty cycle was applied. The investigated area was confined within a circle of 3 mm diameter by a tantalum mask. The lateral homogeneity of ion bombardment was checked by measuring the shape of the sputtered crater with an AMBIOS XP-1 type profilometer. The profilometer was also used to determine the sputtering rate by measuring the crater depth as a function of sputtering time, and to measure the surface roughness parameters. XPS measurements were performed at different depths of interest, depending on the treated sample. For this purpose, samples were transported under UHV conditions from the SNMS chamber to the XPS, i.e., without exposing samples to air. As XPS gives both quantitative and qualitative information about the sample surface, elemental composition and chemical environment can be examined. An XPS equipped by Al/Mg twin anode non-monochromatized X-ray source and Phoibos 100 MCD-5 series hemispherical energy analyzer produced by Specs was used. Electron states were excited by aluminum Al K α X-ray source (with excitation energy 1486.6 eV and line width 0.86 eV). The elemental composition was obtained from a single region spectra with 10 eV pass energy. All binding energies were referenced to the freshly sputtered Au 4f^{7/2} peak at 84.0 eV binding energy. The intensities were calculated from the peak areas by fitting in the CasaXPS program [37].

3. Results and discussion

3.1. Surface characterization of the coated screens

Fig. 2 shows the morphology of the chromium- and nickel-coated screens. The composition of the screens was controlled by SEM-EDS area analysis before each treatment. 100% Cr or Ni appropriately was measured on the coated screens surface, which means that the substrate material of the screen was perfectly masked. It can be observed in Fig. 2a and d, that the chromium coating contained microcracks while



Fig. 3. Cross-section SEM images of the a) 0 time, b) 1 time, c) 2 times used Cr-coated screen and d) 0 time, e) 1 time, f) 2 times used Ni coated screen.

the nickel coating was smooth. During nitriding, the surface morphology of both screens changed. Ions from the plasma bombarded the biased screen directly, so its surface also became nitrided. Due to bombardment with nitrogen ions, after the first nitriding, a grainy surface structure was formed, which is especially well visible in Fig. 2b and e compared to the original surface. Repeating the ASPN process, the general morphological character of the surface texture did not change significantly, but the areal attributes (as defined by the ISO 25178-2 standard) changed very clearly (Fig. 2c and f).

The presence of nitrogen – which is higher in the case of the chromium covered screen – indicates either the formation of nitrides or embedded molecular nitrogen. It has to be noted that the formed nitride layer on top of the screen is exposed to constant sputtering by the ions, which removes and re-deposits parts of the nitride layer. Probably this mechanism caused the porous structure of the nickel screen, well observable in Fig. 2e and f. Another interesting difference between the two screens is the appearance of iron on the nickel covered screen after the ASPN cycles. This iron is supposedly sputtered and deposited here either from the sample or from uncovered parts of the equipment.

Fig. 3 shows the cross-section of the active screens. The average thickness of the chromium coating is $30 \,\mu$ m, the nickel layer is $75 \,\mu$ m. The nitrided parts of the layers cannot be distinguished based on these cross-sectional images. The progression of the cracks – which were already present prior to the first process, as can be seen in Fig. 2a – in the chromium layer during the subsequent ASPN steps is visible in Fig. 3a, b, c. During nitriding, this layer became more brittle, which can be caused by the formation of the Cr-nitride. The porosity of the nickel layer is clearly visible in Fig. 3e and f. A slight increase in the porosity can be observed as an effect of the second nitriding process.



Fig. 5. 3D contact-mode AFM images of a) 1 time, b) 2 times nitrided samples with Ni-coated screen.

3.2. Surface characterization of the samples

After nitriding, the appearance of the sample surface remained polished as it was at the end of the preparation. This can be considered quite unusual since the nitriding process generally causes a matte, grey discoloration on the surface [38,39]. The typical white layer [40,41] was not formed on any of the surfaces. The original tempered microstructure was retained after the plasma nitriding.

It can be seen in the XRD patterns of Fig. 4 that the main phase is alpha-iron, which has a body-centered cubic crystal structure. Besides this phase, another form of the iron was found, which has a face-centered cubic structure corresponding to austenite. These phases can be observed on both samples, which were nitrided by the nickel screen. The nitridation process did not cause the hardening of the steel. The hardness rates were within the deviation of the base material, 420 HV0.01 \pm 45.

The lattice constant of the FCC phase was 3.85 Å, which decreased in PNi2x, and the peaks of Ni also appeared, as can be seen in Fig. 4. In contrast, the samples which were nitrided by the chromium screen had a different reaction to the process. Only the BCC iron phase can be detected clearly in both samples, but the lattice constant of this phase was also decreased. Neither iron nitride nor chromium nitride was formed on the surface.

SNMS depth profiling was used to analyze the elemental composition in the function of the depth from the sample surface. The depth profiles characterize the distribution of the main elements originating from the base material and the screen. Nitrogen was investigated separately by XPS, especially at the interfaces of the layered structure, e.g., between the deposited coating element and the iron. For this purpose, the SNMS measurements were interrupted at the required stages. AFM was used to analyze the morphology of the surface before previous measurements.

Fig. 5 presents of contact-mode AFM images made on the samples. As can be seen, the maximum distance between the highest and lowest points on the sample surface increased from 71 nm to 360 nm between the first and second nitriding process. The surface roughness (S_a) increased from 4.7 nm to 30.6 nm. The differences were caused by the increased amount of deposited coating elements.

As can be seen in Fig. 6a and c, Ni was deposited on the surface of both samples, which were treated by a Ni-coated screen. According to the XPS results presented in Fig. 7b and d, Fe–N bonds were not detected, which proves that iron nitride was not formed during the process. The depth of the diffused layer was 291 nm in the PNi1x and 421 nm in the PNi2x sample. Only molecular nitrogen was found in this

measured depth at the interface (Fig. 6b and d). According to the porous surface morphology, the molecular nitrogen, which came from the nitriding gas, could be stuck in the pores and voids during the layer deposition.

SNMS depth profiles of the samples treated by the Cr-coated screen are shown in Fig. 7a and c. As can be seen, the curves corresponding to Cr and O have a similar profile, which suggests the presence of chromium-oxide in the layer. The gas mixture that was used for the process is free from oxygen, and the amount of oxygen in the layer is much more than what could be expected as remaining in the chamber. This oxide layer was most probably formed during the flushing of the chamber with air after the end of the nitriding process. Although in the nitrogen plasma chromium-nitride was deposited on the surface of the sample, the formed CrN and Cr2N compounds are not stable and easily oxidize to Cr₂O₃ in the presence of oxygen and on high temperature [42] - which conditions are true while flushing the still hot sample with air. The XPS diagrams (Fig. 7b and d) also show high Fe-N content below the chromium-rich layer, although it was not detected by XRD on the surface. It means iron-nitride could be formed on the interface of iron and chromium oxide. The characteristic of the depth profile diagrams is the same for both samples. The thickness of the chromium oxide layer in the PCr1x sample was 167 nm, in the PCr2x sample, was 198 nm. Although the scale of the XPS diagrams ends at 300 nm, the chromium profile already decreased to zero before that, which means that the chromium could not diffuse into the substrate.

Fig. 8 presents SEM-EDS maps and shows the distribution of Fe, N, Cr, Ni elements in the cross-section of the samples. Since the formed layer thicknesses were similar in both cases (with both plating materials), only two samples are presented in Fig. 8. Copper support was

used for the grinding, to avoid damaging the deposited layers. The EDS maps confirm our previous observations: the Ni layer is thicker compared to the Cr. Unfortunately, the N content of the samples cannot be evaluated with this detector.

These results also shed new light on the accepted nitriding models, which place great emphasis on the sputtered particles from the screen as the main carriers of nitrogen for the nitriding of steel on the sample surface. As previously discussed, by using a pure steel active screen, the sputtered higher iron-nitride (Fe1-3N) particles carry physically and chemically adsorbed nitrogen to the steel surface [10]. In the model, the nitriding of the iron on the surface is fuelled by both the desorbed nitrogen atoms and also the decomposition of the higher metastable nitrides. The surfaces of both the Cr and Ni electroplated screens are nitrided during the ASPN process. Thus chromium- and nickel-nitrides are sputtered and subsequently deposited on the surface of the steel samples. In both cases, the sputtered nitride particles can carry the physically adsorbed nitrogen to the surface, similar to the model with iron-nitride particles. However, the desorption of adsorbed nitrogen from these particles did not lead to the direct formation of iron-nitride. In the case of nickel, only molecular nitrogen was found without any trace of iron-nitride. In the case of chromium, iron-nitride was found on the interface, where the decomposition and oxidation of chromiumnitride provided the nitrogen and chemical drive for the nitriding of iron. The fact that physically adsorbed nitrogen is available in both cases and that iron-nitride was not observed in the case of nickel proves that the chemical decomposition was the main driving force of iron nitriding. The nickel-nitride is more stable exposed to oxygen (confirmed by the lack of oxides based on the SNMS results) and seems to retain its chemically bound nitrogen.



Fig. 6. Characteristic of the a) PNi1x by SNMS, b) PNi1x by XPS, c) PNi2x by SNMS, d) PNi2x by XPS measurements.



Fig. 7. Characteristic of the a) PCr1x by SNMS, b) PCr1x by XPS, c) PCr2x by SNMS, d) PCr2x by XPS measurements.



Fig. 8. EDS elemental maps made on the PCr2x and PNi2x samples along their cross-section.

Our observations also offer suggestions to refine the model of the ASPN process in the classical case, when a pure steel active screen is used. After the initial deposition of iron-nitrides on the sample surface, the subsequent development of nitriding is primarily driven by the decomposition of the higher-nitrides to lower nitrides (from $Fe_{1-3}N$ to Fe_4N), which release chemically active nitrogen atoms to the surrounding steel matrix. The effect of physically adsorbed and later desorbed nitrogen is much lower since iron-nitride cannot be found on nitrided samples, where only physically released nitrogen was present (such as in the case of an active screen electroplated with nickel).

4. Conclusion

The material of the active screen is a major parameter that influences the outcome of the nitriding process. In this study, nickel or chromium covered active screens were used to investigate this effect on nitrided steel samples. ASPN was performed at 510 °C for 4 h in a gas mixture containing 75% N₂ and 25% H₂. With these process parameters, Ni was detected on the surface of the treated samples in a deposited layer with several hundreds of nanometres thickness. However, iron-nitride was not formed on the surface, and only molecular nitrogen was found on the interface of nickel and iron. After the nitriding with the Cr coated screen, mostly Cr₂O₃ was formed on the surface with an average thickness of 170 nm. At the interface, Fe–N bonds were found, which indicate that low amounts of iron-nitride could be created with the own Fe content of the substrate.

The results point out that the role of physically adsorbed and subsequently desorbed nitrogen by the sputtered particles is overestimated in the previous ASPN models. Iron nitriding was only observed when the chemical decomposition of other nitrides provided reactive nitrogen atoms to the steel matrix, such as the interface of chromium-iron.

CRediT authorship contribution statement

Dorina Kovács:Investigation, Methodology, Writing - original draft, Visualization.János Dobránszky:Conceptualization, Supervision.Tamás Fodor:Investigation.Viktor Takáts: Investigation.Attila Bonyár:Investigation, Writing - review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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