Effects of plasma nitriding on tempered steel

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Abstract. Conventional plasma nitriding is highly widespread in the industry, but it has essential problems such as edge effect and hollow cathode effect. These problems induced the development of active screen plasma nitriding (ASPN) where the plasma forms on the active screen instead of the sample, which is completely insulated. However, this arrangement does not work with large furnaces, so the usage of bias voltage is necessary in industrial circumstances. In this experiment, the effects of different biases were examined, specifically the hardness, layer thickness, formed phases and the diffusion of nitrogen. In conclusion, the 20 % bias had the best effects on the sample with the defined nitriding parameters in this research.

1. Introduction
Plasma nitriding is a thermochemical surface treatment which can improve the hardness and the wear resistance of the steel. When the nitrogen – hydrogen gas mixture enters the process chamber, the gas becomes ionized between the anode (furnace wall) and the cathode (workpiece). The positively charged ions bombard the surface of the piece which heats up [1–3]. This issue a glow discharge when the particles absorb into the surface of the material. The three most important factors of the treatment are the pressure, the high voltage and the ratio of the used gas mixture. The temperature and time of the treatment also influences the properties of the nitride layer.

The direct current plasma nitriding (DCPN) is a traditional surface treatment over 40 years. The DCPN technology has different shortcomings which cause a damage on the parts, such as ‘edge effect’ or ‘hollow cathode effect’ [3–5]. The active screen plasma nitriding was developed to solve the previous problems. In this technology the plasma is produced on the screen, not directly on the workpiece therefore it causes a more homogenous surface [6–8]. In this case the worktable is isolated from the place of the voltage, so heating of the ASPN is slow. To increase this process the components are in a floating potential or a relative lower bias voltage [8, 9].

2. Materials and methods
The materials used in this study are tempered 42CrMo4 type low alloy steel and annealed C45 type carbon steel, their chemical compositions are shown in table 1. The sample disks were 20 mm in diameter and 6 mm in thickness. The sample’s surface was mechanically ground with 80 to 2500-grit SiC emery paper, finely polished with 3 μm diamond suspension. They were cleaned in acetone and dried before placement to the vacuum chamber.
Table 1. Chemical composition of materials in wt.%

| Steel   | C    | Si  | Mn   | P      | S       | Cr    | Mo   |
|---------|------|-----|------|--------|---------|-------|------|
| 42CrMo4 | 0.38–0.45 | ≤ 0.4 | 0.6–0.9 | ≤ 0.025 | ≤ 0.035 | 0.9–1.2 | 1.5–0.3 |
| C45     | 0.42–0.5  | ≤ 0.4 | 0.5–0.8  | ≤ 0.045 | ≤ 0.045 | ≤ 0.4  | ≤ 0.1  |

The plasma nitriding system was built by own plan. Parameters easily set to the different methods. Figure 1 shows the schematic workplace of the chamber. The active screen was made of unalloyed 1.0330 type steel. The dimension is Ø100×85 mm with Ø 5×8 mm holes and centers. All treatments were carried out using the same parameters as shown in Table 2 and following the same procedures. Bias voltage was adapting to the main voltage. The bias is 10 %, 20 % of the main voltage at the same parameters of the process. After the nitriding, free cooling started to the turning of the plasma. The workpiece was cooling down from the process temperature to the room temperature under the nitriding pressure.

![Schematic model of the nitriding chamber (biased arrangement)](image)

Table 2. Parameters of plasma nitriding

| Voltage (V) | Current (A) | Gas mixture N₂:H₂ (%) | Temperature (°C) | Pressure (torr) | Time (h) |
|-------------|-------------|------------------------|------------------|----------------|---------|
| 490 – 540   | 0.9 – 1.5   | 25:75                  | 490              | 2.8            | 4       |

Optical microscope and scanning electron microscope were used to examine morphology and measure thickness of nitried layer. The hardness profile was determined by micro Vickers indenter with a load of 50g. The phase composition of the compound layer on the surface was determined by X-ray diffractometer (XRD) using CuKα radiation [10–15].
3. Results and discussion

3.1. Surface analysis by optical microscope
First, DCPN and ASPN treated C45 samples were compared. The edge effect was observed at the treatment of DCPN as it can be seen in Figure 2a) and b). The well-visible erosion ring has reduced the hardness toward the centre of the sample, while the microhardness value is much higher in the edge area, due to thermal gradient, as it was presented in another work; this causes unequal microhardness values on the surface [6]. The nitride layer depth is unequal in the erosion zone compared to the centre.

![Figure 2. Discoloured surface after plasma nitriding](image)

3.2. Hardness and thickness measurements
Figures 3 and 4 show the cross-sectional hardness distribution of the different samples. The compound layer thickness was measured by using the distance measurement tool of the scanning electron microscope. The hardness decreased considerably toward the core of the material in each condition. The samples of steel C45 show a significant hardness drop after the compound layer while the samples of steel 42CrMo4 have smooth transition to the core hardness. This means that the diffusion zone is substantially larger with the alloyed steel. The effect of bias is also visible on the hardness profile. As higher percentage of the voltage was used as bias, higher surface hardness and thicker compound layer was formed. In proportion to DCPN, the use of bias cause significant increasing of hardness.

![Figure 3. Two hardness profiles of steel C45](image)
Figure 4. Hardness profiles of steel 42CrMo4

Scanning electron microscope was used to determine the depth of the compound layer. Figure 5 shows the cross section of two C45 steel samples. The layer thickness was 8.6 µm with DCPN and 5.5 µm with ASPN, respectively.

Figure 6 shows the cross section of the 42CrMo4 type low alloy steel samples. The layer thickness was 15 µm with DCPN, 11.2 µm with 10% bias and 15.5 µm with 20% bias.

Figure 5. Compound layer thickness of steel C45 a) DCPN, b) ASPN treated samples

Figure 6. Compound layer thickness of steel 42CrMo4 a) DCPN, b) using 10% Bias, c) using 20% Bias treated samples

3.3. X-Ray diffraction analysis

DC plasma nitriding samples were investigated by XRD analysis. Figure 7-8 contain the diffraction pattern of C45 and 42CrMo4. γ’ phase, as Fe₄N, was formed on both sample with the same intensity [15]. As observed in Figure 8, magnetite is also formed on the surface. It means some oxygen is present in the vacuum chamber, but this rate is minimal, it didn’t influence the compound layer.
4. Conclusions
The annealed C45 type carbon steel is not a typical grade for plasma nitriding. The compound layer thickness is formed on the surface like as other tempered steel, but the depth of the diffusion zone is lower than the other case.

The effect of bias on steel 42CrMo4 was analysed with different methods. The results show that higher hardness can be reached with higher bias voltage. The hardness profile also shows that the alloyed steel has a much thicker diffusion zone than the unalloyed. With higher bias, slightly thicker compound layer is formed which can be examined well using SEM.

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