Influence of Co Content and Chemical Nature of the Co Binder on the Corrosion Resistance of Nanostructured WC-Co Hardmetals in Acidic Solution

Tamara Aleksandrov Fabijanić, Marin Kurtela, Matija Sakoman and Mateja Šnajdar Musa

1 Faculty of Mechanical Engineering and Naval Architecture, University of Zagreb, Ivana Lučića 5, 10000 Zagreb, Croatia; marin.kurtela@fsb.hr (M.K.); matija.sakoman@fsb.hr (M.S.)
2 Department of Polytechnics, University of Rijeka, Sveučilišna avenija 4, 51000 Rijeka, Croatia; mateja.snajdar@uniri.hr
* Correspondence: tamara.aleksandrov@fsb.hr; Tel.: +385-9845-3916

Abstract: The electrochemical corrosion resistance of nanostructured hardmetals with grain sizes \(d_{WC} < 200\) nm was researched concerning Co content and the chemical nature of the Co binder. Fully dense nanostructured hardmetals with the addition of grain growth inhibitors GGIs, VC and \(\text{Cr}_2\text{C}_2\), and 5 wt.% Co, 10 wt.% Co, and 15 wt.% Co were developed by a one cycle sinter-HIP process. The samples were detailly characterized in terms of microstructural characteristics and researched in the solution of \(\text{H}_2\text{SO}_4 + \text{CO}_2\) by direct and alternative current techniques, including electrochemical impedance spectroscopy. Performed analysis revealed a homogeneous microstructure of equal and uniform grain size for different Co contents. The importance of GGIs content adjustment was established as a key factor of obtaining a homogeneous microstructure with WC grain size retained at the same values as in starting mixtures of different Co binder content. From the conducted research, Co content has shown to be the dominant influential factor governing electrochemical corrosion resistance of nanostructured hardmetals compared to the chemical composition of the Co binder and WC grain size. Negative values of \(E_{\text{corr}}\) measured for 30 min in 96% \(\text{H}_2\text{SO}_4 + \text{CO}_2\) were obtained for all samples indicating material dissolution and instability in acidic solution. Higher values of \(R_p\) and lower values of \(i_{\text{corr}}\) and \(v_{\text{corr}}\) were obtained for samples with lower Co content. In contrast, the anodic Tafel slope increases with increasing Co content which could be attributed to more pronounced oxidation of the higher Co content samples. Previously researched samples with the same composition but different chemical composition of the binder were introduced in the analysis. The chemical composition of the Co binder showed an influence; samples with lower relative magnetic saturation related to lower C content added to the starting mixtures and more W dissolved in the Co binder during the sintering process showed better corrosion resistance. WC-5Co sample with significantly lower magnetic saturation value showed approximately 30% lower corrosion rate. WC-10Co sample with slightly lower relative magnetic saturation value and showed approximately 10% lower corrosion rate. Higher content of \(\text{Cr}_2\text{C}_2\) dissolved in the binder contributed to a lower corrosion rate. Slight VC increase did not contribute to corrosion resistance. Superior corrosion resistance is attributed to W and C dissolved in the Co binder, lower magnetic saturation, or WC grain size of the sintered sample.

Keywords: nanostructured hardmetals; Co content; GGIs; chemical nature of Co binder; grain size; electrochemical corrosion resistance; \(\text{H}_2\text{SO}_4 + \text{CO}_2\)

1. Introduction

Hardmetals contain tungsten carbide WC particles joined by a binder, most commonly cobalt Co, by a liquid phase sintering process. The properties of the obtained composite derive directly from its constituents; hard and brittle carbides and softer and more ductile binder [1]. By connecting these two components, superior mechanical, physical, and...
chemical properties are achieved. In recent years, the development of hardmetals is based mainly on the application of ultrafine and nano WC particles (grain size less than 0.5 µm) which require the addition of grain growth inhibitors GGIs to retain the WC grain size in the sintered product. Consequently, a homogeneous microstructure and significantly improved mechanical properties (hardness, wear-resistance, and strength) are achieved. Furthermore, achieving a homogeneous microstructure with a WC grain size in the nano area (<0.2 µm) allows application at higher cutting speeds, lower tolerance, and longer tool life. Due to superior mechanical properties, hardmetals are used in various applications with certain limitations in chemically aggressive environments because of relatively poor corrosion resistance [1].

The corrosion mechanism of conventional WC-Co hardmetals in the neutral and acidic solution is governed by the reduction of the Co binder. At the same time, WC particles are not affected by the corrosion attack [2–4]. Zheng et al. found that the binder dissolution started from the center of binder pools in the acid media, independent of binder chemical nature, and spreads to the edges until the binder phase was consumed entirely [4]. Accordingly, it is expected that the corrosion rate will increase with increasing Co content in the starting mixture.

Besides Co binder content, the corrosion mechanisms in hardmetals depend on many other factors such as surface characteristics and integrity, corrosive environment, and hardmetal microstructure [5–7]. Hardmetal microstructure, including WC grain size, binder composition/binder chemical nature, grain growth inhibitors GGIs, and porosity, influence the corrosion behavior of hardmetals [3–10]. Researchers have reported different experimental variations concerning the relationship between microstructure and corrosion resistance.

The chemical nature of the Co binder is represented by magnetic saturation. It depends on the C content added to the starting hardmetal mixture and technological processes of consolidation, among which the most important are sintering parameters and atmosphere [11–13]. Co binder is advantageous because of the relatively large carbon contents that give the preferred two-phase WC-Co composition, commonly called the carbon window [14,15]. During sintering, the Co binder is alloyed with tungsten (W) and carbon (C); other constituents such as GGIs also add alloying elements to the binder [14]. If a higher amount of tungsten is dissolved in the Co binder, lower relative magnetic saturation values would be obtained [3], and the formation of the brittle η-phase carbides M6C and M12C would occur in the microstructure of hardmetals. It was found from previous research that hardmetals with lower relative magnetic saturation values show lower values of corrosion current density (i_{corr}) and critical current density (i_{crit}) measured by potentiodynamic polarization [2,6,7,16].

Regarding the influence of the WC grain size, different conclusions can be found in the literature. Li Zhang et al. found pseudo-passivation behavior of conventional hardmetals with the WC grain sizes ranging from 1.2 µm to 8.2 µm in sulfuric acid H2SO4 and better corrosion resistance of the coarse WC grain sizes [3]. On the other hand, Imasato et al. found that the corrosion rate of the WC-Co alloy with a smaller WC grain size was lower than coarse WC grain size both in acid and neutral solution. WC-Co alloy with a smaller WC grain size showed a higher corrosion resistance in the polarization test because of the low current densities of active dissolution and passivated region in the polarization curve [17]. Also, they found that the amount of dissolved metals in neutral and acidic solutions decreased with decreasing WC grain size and carbon content. Most published papers refer to conventional hardmetals, while there is still a lack of results published on nanostructured hardmetals with a WC grain size less than 200 nm.

The paper summarizes long-term research on the corrosion resistance of nanostructured hardmetals. From previous research it was found that the chemical nature of the binder has a more substantial influence on the electrochemical corrosion resistance compared to Co content in the starting mixture in neutral and acidic solution, which was quite surprising and not in line with conventional WC-Co hardmetals [6,7]. The presented
research was performed to bring more exact conclusions on the influence of Co content and other microstructural characteristics on the corrosion resistance of nanostructured hardmetals.

2. Materials and Methods

Different starting mixtures were prepared to investigate the influence of Co content and chemical nature of the Co binder on the corrosion resistance of nanostructured hardmetals with a grain size $d_{WC} < 200$ nm. WC powder produced by H.C. Starck Tungsten (Goslar, Germany) with an average grain size $d_{BET}$ of 95 nm and a specific surface area (BET) of 3.92 m$^2$/g, classified as real nanopowder with a grain size less than 100 nm, was used as a carbide phase. Grain growth inhibitors GGI’s, vanadium carbide VC and chromium carbide $Cr_3C_2$ were added to the starting mixtures. VC powder has an average grain size $d_{BET}$ of 350 nm and a specific surface area (BET) of 3.0 m$^2$/g, while $Cr_3C_2$ has an average grain size $d_{BET}$ of 450 nm and a specific surface area (BET) of 2.0 m$^2$/g. Besides controlling the WC grain growth, VC and $Cr_3C_2$ increase hardness and reduce the rate of corrosion. At the same time, Cr significantly lowers the initial melting point and broadens the melting range, particularly at low carbon levels [12–14,16]. The amount of VC and $Cr_3C_2$ differs for each mixture; it was optimized to withhold WC powder size in the sintered samples and increased Co binder content. Half micron cobalt HMP Co, produced by Umicore (Brussels, Belgium), was used as a binder. Three mixtures with different Co content; 5, 10, and 15 wt.%Co were prepared. The consolidation process consisted of powder mixture homogenization in a horizontal ball mill (Zoz GmbH, Wenden, Germany). Compacting was performed by uniaxial die press type CA-NCII 250 (Osterwalder AG, Lyss, Switzerland). Final consolidation to total density was achieved by one cycle sinter-HIP process by furnace FPW280/600-3-2200-100 PS (FCT Anlagenbau GmbH, Sonneberg, Germany) at 1350 °C for 30 min, followed by 100 bars Argon 4.8 pressure for 45 min. The characteristics of the starting mixtures are presented in Table 1.

| Mixture  | Starting WC Powder          | Grain Size $d_{BET}$, nm | Specific Surface, m$^2$/g | Co, wt.% | GGI, wt.%      |
|----------|-----------------------------|--------------------------|---------------------------|----------|----------------|
| WC-5Co   | WC DN 4-0 (H.C. Starck)    | 95                       | 3.92                      | 5        | 0.3% VC 160   |
|          |                             |                          |                           |          | 0.5% $Cr_3C_2$160 (H. C. Starck) |
| WC-10Co  |                             | 10                       | 3.92                      | 10       | 0.75% $Cr_3C_2$160 (H. C. Starck) |
|          |                             |                          |                           |          | 0.75% VC 160  |
| WC-15Co  |                             | 15                       | 3.92                      | 15       | 1.13% $Cr_3C_2$160 (H. C. Starck) |

The goal was to develop fully dense samples with optimal microstructural characteristics with no irregularities such as $\eta$-phase in the structure. Previous research found that C and GGI’s content and WC grain size have a more substantial influence on the corrosion resistance of nanostructured hardmetals than Co content [6,7]. For the mentioned reason, special care was taken to obtain the optimal and comparable microstructural characteristics of consolidated samples.

The samples were detailly characterized, especially in terms of microstructural characteristics. The characterization of samples consisted of density measurements (Metler Toledo) according to ISO 3369:2006, the specific saturation magnetization (Setaram Instrumentation, Sigmameter) according to D6025, and the coercive field strength measurement (Foerster, Koerzimat 1.096) according to ISO 3326. Diamond disc and pastes were used to grind and polish the samples’ surface before microstructural characterization and electrochemical measurements. Microstructural characterization consisted of porosity, free carbon, and $\eta$-phase evaluation. It was performed by comparing the sample’s surfaces with photomicrographs from the standard ISO 4499-4:2016. For that purpose, an optical microscope (Olympus, Shinjuku City, Tokyo, Japan) was used. A field emission scanning
electron microscope FESEM (Zeiss, Oberkochen, Germany) was used for WC grain size measurement by the linear intercept method and detection of irregularities such as abnormal growth and WC grains grouping or Co lakes. X-ray diffraction XRD analysis was used to identify the phases present in consolidated samples and exclude the occurrence of \(\eta\)-phase.

After detailed characterization of the samples and determination of optimal microstructural characteristics, the corrosion measurements were performed. The surface of the samples was placed into the corrosion cell filled with \(\text{H}_2\text{SO}_4 + \text{CO}_2\) (pH = 0.6). As reference electrode, saturated calomel electrode SCE (SCHOTT Instruments GmbH, Mainz, Germany) with a potential of + 0.242 V according to the standard hydrogen electrode was selected. Graphite wires were used as a counter electrode. The samples were first researched by direct current techniques DC, the open-circuit potential \(E_{\text{corr}}\), the linear polarization resistance (LPR), and the Tafel extrapolation method. Corrosion potential \(E_{\text{corr}}\) versus SCE was recorded for 30 min. LPR was carried out in the potential range from \(-0.02\) V vs. open circuit potential to 0.02 V vs. open circuit potential with a scan rate of 0.167 mV/s. Tafel extrapolation was conducted in the potential range from \(-0.25\) V vs. open circuit potential to 0.25 V vs. open circuit potential, total points 1001 with a scan rate of 0.167 mV/s. Immediately after the DC techniques, the samples were researched by alternating current (AC) techniques, more precisely by electrochemical impedance spectroscopy (EIS).

The EIS start frequency was 100,000 Hz, and the end frequency was 0.001 Hz; the amplitude was in the range of 10 mV root-mean-square (RMS). The recorded measurements were analyzed by software SoftCorr III (AMETEK Scientific Instruments, Princeton applied research, Berwyn, PA, USA). A convenient electrical equivalent circuit (EEC) was selected by fitting the results of measurements and presented in Nyquist and Bode plots. At each excitation frequency, an imaginary impedance component \(Z_{\text{im}}\) is drawn according to the actual impedance component \(Z_{\text{re}}\). The impedance and the phase shift curves were plotted against the excitation frequency. Both DC and AC techniques were performed on the potentiostat AMETEK, Princeton applied research, model VersaSTAT3, and the results were recorded and analyzed by software SoftCorr III (AMETEK Scientific Instruments, Princeton applied research, Berwyn, PA, USA).

3. Results
3.1. Microstructural Characteristics of Consolidated Samples

Characteristics of consolidated samples with different Co content are presented in Table 2. This section may be divided by subheadings. It should provide a concise and precise description of the experimental results, their interpretation, as well as the experimental conclusions that can be drawn.

| Sample | Density, g/cm\(^3\) | Relative Density, % | Magnetic Saturation, \(\mu\text{Tm}^3/\text{kg}\) | Rel. Magnetic Saturation, % | Coercive Force, kA/m | ISO Porosity | \(d_{WC}\), nm |
|--------|---------------------|---------------------|-----------------------------------------------|-----------------------------|-----------------------|--------------|-------------|
| WC-5Co | 14.91               | 100.0               | 8.4                                           | 92                          | 52.0                  | A00 B00 C00  | 187         |
| WC-10Co| 14.31               | 100.0               | 14.8                                          | 79                          | 40.0                  | A00 B00 C00  | 198         |
| WC-15Co| 13.84               | 100.0               | 22.3                                          | 79                          | 37.0                  | A00 B00 C00  | 192         |

Full densification was achieved for all samples. The samples are characterized by the lowest possible degree of porosity, A00, B00, and C00, meaning no uncombined/free carbon or \(\eta\)-phase were revealed on the sample’s surface. A high density of samples is related to Co liquid phase, which is spreading onto the surrounding WC particles. Binder propagation is associated with Laplace forces acting along the wetting front between Co binder and WC grains while rearranging the WC particles and reducing the mean distance between neighboring particles, resulting in densification [16,18]. It may be concluded
that the WC particles were rearranged, and Co binder filled the micropores between the neighboring WC grains resulting in a theoretical/full density of the samples.

The amount of W dissolved in the Co binder phase can be assessed by measuring the magnetic saturation. The saturation value of Co decreases linearly with the addition of tungsten W and is not affected by the carbon content in the solution [19]. Typical relative magnetic saturation/percentage saturation ranges from 80–100% [19], while percentage saturation values higher than 70% indicate two-phase WC-Co microstructure. The highest percentage of 91% was measured for the WC-5Co sample, while 79% was measured for WC-10Co and WC-15Co samples. The two-phase WC-Co microstructure of researched samples is confirmed by optical analysis. Based on coercive force measurement it was estimated the WC grain size. Measured values indicate that all samples fall in the nano range. Two-phase WC-Co microstructure was confirmed by optical analysis, XRD analysis where only WC with the hexagonal crystal structure and Co with FCC cubic crystal structure were identified. Investigation of microstructure revealed homogeneous, uniform distribution of WC grains, without abnormal grain growth due to optimal GGIs content added to the starting mixtures. It was necessary to adjust the content of GGIs for different Co content to achieve a homogeneous and comparable microstructure with retained WC grain size of the starting powders in the sintered samples. Co binder was uniformly distributed, and no Co lakes occurred. Microstructure images and XRD patterns of samples are presented in Figures 1–3.

![Microstructure Image](image_url)
Figure 1. Microstructure and XRD pattern of WC-5Co sample [20]. (a) SEM image of microstructure; (b) XRD pattern.

Figure 2. Cont.
Figure 2. Microstructure and XRD pattern of WC-10Co sample [21]: (a) SEM image of microstructure; (b) XRD pattern.

Figure 3. Cont.
3.2. Results of DC Techniques

The results of electrochemical DC techniques are presented in Table 3.

| Sample       | $T_s$ [°C] | $E_{corr}$ vs. SCE [mV] | $R_p$ [Ω cm$^2$] | $\beta_a$ [mV/dec] | $\beta_c$ [mV/dec] | $i_{corr}$ [µA/cm$^2$] | $v_{corr}$ [mm/y] |
|--------------|------------|-------------------------|------------------|-------------------|-------------------|----------------------|------------------|
| WC-5Co       | 20 ± 2     | −249                    | 654.5            | 75.31             | 90.37             | 20.7                 | 0.1748           |
| WC-10Co      | 20 ± 2     | −308                    | 452.8            | 98.34             | 97.67             | 36.6                 | 0.3888           |
| WC-15Co      | 20 ± 2     | −291                    | 349.9            | 120.19            | 86.95             | 50.8                 | 0.4162           |

$T_s$—measured temperature; $E_{corr}$—corrosion potential; $R_p$—polarization resistance; $\beta_a$—a slope of anodic Tafel curve; $\beta_c$—a slope of cathodic Tafel curve; $i_{corr}$—corrosion current density; $v_{corr}$—corrosion rate.

3.3. Results of Electrochemical Impedance Spectroscopy EIS

EIS measurements aimed to investigate the corrosion behavior at the interface between the sample surface and electrolyte solution and determine the samples’ corrosion rate. The results are presented in Table 4.

| Sample       | $T_s$ [°C] | $R_s$ [Ω cm$^2$] | $Q$ [µF/cm$^2$] | $n_1$ | $R_p/R_{ct}$ [Ω cm$^2$] |
|--------------|------------|-----------------|----------------|-------|------------------------|
| WC-5Co       | 20 ± 2     | 4.022           | 1.761·10$^{-3}$ | 0.745 | 1.101·10$^{-3}$       |
| WC-10Co      | 20 ± 2     | 4.504           | 2.213·10$^{-3}$ | 0.725 | 8.068·10$^{-2}$      |
| WC-15Co      | 20 ± 2     | 5.797           | 2.552·10$^{-3}$ | 0.683 | 4.657·10$^{-2}$      |

$T_s$—measured temperature; $R_s$—solution resistance between the working electrode and the reference electrode in a three-electrode cell; $Q$—Constant Phase Element (CPE); $n_1$—constant; $R_{ct}$—polarization resistance or resistance to charge transfer on the electrode/electrolyte interface.

As already mentioned in Section 2, a convenient and optimal electrical equivalent circuit (EEC) was selected using software SoftCorr III by fitting the measurements’ results and presented in Nyquist and Bode plots. At each excitation frequency, an imaginary...
impedance component Zim is drawn according to the actual impedance component Zre. The impedance and the phase shift curves were plotted against the excitation frequency. The selected ECC model is shown in Figure 4.

![Figure 4. Selected EEC](image)

The same model R(QR), which best corresponds to the processes and reactions on the sample’s surface, was selected for all samples. It is essential to mention that the same EEC model was established in previous research performed on near nanostructured WC-11Co and WC-11Ni samples [22]. The mentioned indicates the repeatability of the corrosion process between nanostructured hardmetal and acidic solutions. The impedance of a constant phase element is defined as:

\[ Q = [Y(j\omega)^n]^{-1} \]  

(1)

where \( Y \) and \( n (−1 \leq n \leq 1) \) are constants independent of the angular frequency \( \omega \) and temperature. For the value in the range \( 0.6 < n \leq 1 \), CPE has the physical meaning of capacitance, an ideal inductor for \( n = −1 \), and an ideal resistor for \( n = 0 \).

4. Analysis and Discussion

4.1. Influence of Co Content on the Corrosion Resistance of Nanostructured WC-Co Hardmetals

From conducted research, it can be concluded that the corrosion potential \( E_{corr} \) of samples changes depending on the Co content. \( E_{corr} \) of WC-5Co and WC-10Co samples changed from more negative to more positive values indicating that the surfaces of the samples are getting passivated, and a reduction occurred. The corrosion potential curves are unstable and show random fluctuations. A drop from more positive to more negative values was detected for the WC-15Co sample, indicating sample oxidation in contact with the acidic electrolyte and reducing protons at the surface. The changes of \( E_{corr} \) are not significant, and the corrosion potential variations of each sample occurred in a narrow range. Negative values of \( E_{corr} \) measured for 30 min in 96% H₂SO₄ + CO₂ were obtained for all samples indicating material dissolution and instability in acidic solution. The \( E_{corr} \) vs. time curves are presented in Figure 5, and sample Tafel extrapolation curves in Figure 6.

![Figure 5. Corrosion potential \( E_{corr} \) of samples](image)
Higher values of $R_p$ and lower values of $i_{corr}$ were obtained for samples with lower Co content. Accordingly, the corrosion rate in acidic solution increases with increasing Co content due to selective dissolution of the Co matrix. The cathodic Tafel slopes of samples show a similar trend. In contrast, the anodic Tafel slope increases with increasing Co content which could be attributed to more pronounced oxidation of the higher Co content samples. The dependence of polarization resistance $R_p$ and corrosion rate $v_{corr}$ for different Co contents is presented graphically in Figure 7.

Even though the addition of the refractory metal carbides, VC, Cr$_3$C$_2$ in the starting mixtures was increased with increasing Co content to maintain the WC powder size, Co content showed stronger influence on the electrochemical corrosion resistance of nanostructured hardmetal samples with optimal microstructural characteristics. The lowest $v_{corr}$ of 0.1748 mm/y was obtained for the WC-5Co sample, while the highest $v_{corr}$ of 0.4162 mm/y was measured for the WC-15Co sample.

Figures 8–10 present the Nyquist and Bode diagrams for the WC-5Co, WC-10Co, and WC-15Co samples, obtained using the corresponding EEC simulation model of EIS results. The highest $R_p$ of 1.101-103 $\Omega$cm$^2$ was measured for the WC-5Co sample, followed by $R_p$ of 8.068-102 $\Omega$cm$^2$ measured for the WC-10Co sample, and $R_p$ of 4.657-102 $\Omega$cm$^2$ measured for the WC-15Co sample. Higher $R_p$ values were detected for the samples with lower Co content, indicating better corrosion resistance (Figure 11) which corresponds to the
results obtained by DC linear polarization techniques. It can be seen from Figures 8–10 that the radius $r$ of the capacitive semi-circles in the Nyquist plots differ for samples with different Co content. The diameter of the capacitive loop decreased with increasing Co content, indicating better charge transfer resistance on the electrode/electrolyte interface. Subsequently, the decrease in the diameter of the capacitance loop may be ascribed to the weaker protective ability of the sample surface.

(a)  
Figure 8. Nyquist (a) and Bode (b) plots of WC-5Co.

(a)  
Figure 9. Nyquist (a) and Bode (b) plots of WC-10Co.

(a)  
Figure 10. Nyquist (a) and Bode (b) plots of WC-15Co.
Nanostructured hardmetals with optimal microstructural characteristics exhibited behavior similar to previously researched conventional hardmetals with coarser WC grain size. Electrochemical corrosion resistance decreases with increasing Co content in a corrosive, acidic environment due to predominant active Co binder reduction, also known as Co leaching.

4.2. Influence of Co Binder Chemical Nature on the Corrosion Resistance of Nanostructured WC-Co Hardmetals

The chemical nature of the Co binder can be characterized by magnetic saturation. It depends on the C content added to the starting hardmetal mixture and consolidation procedure, where sintering parameters and atmosphere have a crucial influence. As referred in the Introduction, it was found from previous research that C content added to the starting mixture can significantly influence the electrochemical corrosion resistance in both neutral (3.5% NaCl with pH = 6.6) and acidic (96% H\textsubscript{2}SO\textsubscript{4} + CO\textsubscript{2} with pH = 0.6) environments [6,7]. Comparing WC-5Co samples of the same composition, better corrosion resistance was observed for samples with lower C-added content, lower magnetic saturation, and coarser WC grain size. The opposite behavior governed by the C content and magnetic saturation was noted for WC-15Co samples [7]. Compared to C content, GGIs content, and grain size, the Co content showed less impact on the electrochemical corrosion resistance in both acid and neutral solutions. There was no clear trend of increasing corrosion current densities i\textsubscript{corr} and decreasing polarization resistance Rp with increasing Co content typical for conventional hardmetals. Microstructural characteristics, in this case WC grain size, has shown to have a greater influence on the sintered samples corrosion resistance [6,7]. To obtain better insight into the electrochemical corrosion resistance of nanostructured hardmetals, previously researched samples designated as WC-5Co-1 and WC-10Co-1 were introduced in the analysis. WC-5Co-1 and WC-10Co-1 samples were consolidated using same production procedure and characterized by the same methods described in Section 2. The only alteration was the use of lower C and GGIs content added to the starting mixture, which caused different chemical nature of the Co binder and

![Figure 11. The dependence of Rp concerning Co content.](image-url)
lower values of relative magnetic saturation. Characteristics of the additionally introduced samples and comparison with previously characterized nanostructured hardmetals with $d_{WC} < 200$ nm are presented in Tables 5 and 6.

### Table 5. Comparison of WC-5Co samples with different characteristics.

| Sample   | GGI, wt.% | Added C, wt.% | Density, g/cm³ | $\rho_v$ | Relative Magnetic Saturation, % | Coercive Force, kA/m | $v_{corr}$ [mm/y] |
|----------|-----------|----------------|----------------|---------|-------------------------------|---------------------|------------------|
| WC-5Co-1 | 0.41%VC, 0.80% Cr₃C₂ | 0.150 | 14.96 | 100 | 48.0 | 44.9 | 0.1181 |
| WC-5Co   | 0.30%VC, 0.50% Cr₃C₂ | 0.275 | 14.91 | 100 | 92.0 | 52.0 | 0.1748 |

### Table 6. Comparison of WC-10Co samples with different microstructural characteristics and magnetic saturation.

| Sample   | GGI, wt.% | Added C, wt.% | Density, g/cm³ | $\rho_v$ | Relative Magnetic Saturation, % | Coercive Force, kA/m | $v_{corr}$ [mm/y] |
|----------|-----------|----------------|----------------|---------|-------------------------------|---------------------|------------------|
| WC-10Co-1 | 0.37%VC, 0.72% Cr₃C₂ | 0.225 | 14.35 | 100 | 74.7 | 35.1 | 0.3463 |
| WC-10Co  | 0.5%VC, 0.75% Cr₃C₂ | 0.250 | 14.32 | 100 | 79.0 | 40.0 | 0.3888 |

As indicated in Table 5, the WC-5Co-1 sample has a significantly lower relative magnetic saturation of 48.0%, attributed to the lower added C content and the different chemical nature of the Co binder. As mentioned, typical relative magnetic saturation/percentage saturation ranges from 80–100%. Saturation percentage values lower than 70% indicate the presence of microstructural irregularity $\eta$-phase, confirmed by optical, FESEM, and XRD analysis [6,7]. Previous research found that $\eta$-phase most likely enhances the passive layer formation on the sample surfaces, thereby reducing the tendency of sample dissolution and increasing the stability of oxides forming in addition to the existing passive layer on the surface [2,6,7]. The slightly higher measured density of the WC-5Co-1 sample is also associated with $\eta$-phase presence in the structure since W₆Co₆C or W₃Co₃C has higher density when compared to a two-phase WC-Co hardmetal. The coercive force obtained for the WC-5Co-1 sample amounts to 44.9 kA/m and is lower than that of WC-5Co, suggesting a coarser grain size of the WC-5Co-1 sample. Thus, its microstructure can be classified as near nano, in the ultrafine range from 200 to 500 nm. Tafel extrapolation curves of WC-5Co samples are presented in Figure 12.

![Figure 12. Tafel extrapolation curves of WC-5Co samples with different microstructural characteristics.](image-url)
The WC-5Co-1 sample with lower magnetic saturation value and consequently higher W and C content in the Co binder showed approximately 30% lower corrosion rate, which is in line with previous research. Sutthiruangwong and Mori found that the magnetic saturation related to Co binder composition plays an essential role in the corrosion properties of hardmetals [15,23,24]. F.J.J. Kellner et al. found that electrochemical corrosion resistance of hardmetals is influenced by the W and C diffusion in the Co binder amount which is increased by decreasing the WC grain size [25]. They concluded that W and C dissolved in Co binder during the sintering process stabilize the thermodynamically unstable FCC Co crystal structure at room temperature, the amount of which is increased by an increase of W and C content in the binder. FCC Co is characterized by better corrosion resistance compared to the HCP crystal structure of Co, thermodynamically stable at room temperature [25], when a HCP+FCC Co layer around the HCP Co binder is formed [25]. Their research was performed in an alkaline medium. Still, since in this study the tests were performed in an acidic solution where the lower corrosion resistance of hardmetals is attributed to the dissolution of the HCP Co matrix, these claims could be expected to be more pronounced.

Besides lower magnetic saturation, the WC-5Co-1 sample has a higher content of GGIs, VC, Cr$_2$C$_2$ in the starting mixture, as presented in Table 5. It is well-known that GGIs are dissolved and distributed among the WC phase and binder during sintering and influence characteristics of hardmetals [26]. Sutthiruangwong and Mori have found that higher corrosion resistance can be assigned to binders that experience higher chromium dissolution rates during sintering [23,24]. Tomlinson and Ayerst found that small additions of Cr$_2$C$_2$ improve the electrochemical corrosion resistance of hardmetals due to the formation of Cr$_2$O$_3$ film on the Co binder surface [26]. On the other hand, a small addition of VC in combination with Cr$_2$C$_2$ decreases the positive Cr$_2$C$_2$ influence in acidic solution [26].

The WC-5Co-1 sample has a 0.3 wt.% higher content of Cr$_2$C$_2$, which dissolved in the binder and contributed to a 30% lower corrosion rate than the WC-5Co sample. In this research, it is hard to distinguish which factor has the most substantial influence on electrochemical corrosion resistance. To specify more clearly, WC-10Co samples with different microstructural characteristics and magnetic saturation were compared in Table 6.

The WC-10Co-1 sample has a slightly lower relative magnetic saturation of 74.7% than the WC-10Co sample, which is related to marginally lower C content in the amount of 0.025 wt.% added to the starting mixture. Lower C content resulted in more W dissolved in the Co binder, which, as in the case of the WC-5Co sample, most probably stabilized Co’s thermodynamically unstable FCC crystal structure. Consequently, the WC-10Co-1 sample is characterized by a marginally higher measured density. Density values vary within the two-phase region of the WC-Co phase, and is increased with an increasing amount of W which remains in the Co binder [27]. Sample WC-10Co-1 is located at the lower end of the two-phase WC-Co region in the isothermal part of the WC-Co phase diagram. Therefore, the measured density value is slightly higher than the theoretical density, despite η-phase not detected. The coercive force of WC-10Co-1 amounts to 35.1 kA/m, i.e., lower when compared to WC-10Co, which indicates a coarser grain size of the WC-10Co-1 sample. The same was noted for WC-5Co samples. Tafel extrapolation curves of WC-10Co samples are presented in Figure 13.
The WC-10Co-1 sample with a lower relative magnetic saturation value showed approximately 10% lower corrosion rate. Both microstructures consist of two phases, WC and Co, with no η-phase detected in the microstructure. The difference in GGIs content is relatively small and amounts to an extra 0.13 wt.%VC and 0.03 wt.% Cr$_3$C$_2$ added to the WC-10Co mixture. It can be concluded that the slight increase of VC wt.% did not contribute to corrosion resistance which corresponds to previously published research. Machio et al. found that small VC addition of 0.4 wt.% increase $i_{\text{corr}}$ and make hardmetals more sensitive to pitting corrosion due to VC influence on the W dissolution in the Co matrix and formation of (V,W)C layer around the WC grains. VC decreases the dissolution of W atoms in the Co binder during the sintering process and increases the magnetic saturation compared to pure WC-Co hardmetal without GGIs [1,28]. D.S. Konadu et al. found that WC-Co hardmetal possesses nobler corrosion resistance compared to 0.4 wt.%VC containing hardmetal in both HCl and H$_2$SO$_4$ [1]. Accordingly, better corrosion resistance in this research may be related to W and C dissolution in the Co binder, magnetic saturation, or WC grain size in the sintered sample.

5. Conclusions

The following conclusions can be drawn from the conducted research:

1. Fully dense nanostructured hardmetals with a WC grain size $d_{\text{WC}} \leq 200$ nm were developed utilizing the single-cycle sinter-HIP process. For different Co contents, a homogeneous microstructure of equal and uniform grain size without microstructural defects in the form of carbide agglomerates, abnormal grain growth, or Co lakes was successfully obtained.

2. The importance of GGIs content adjustment was established as a key factor of obtaining a homogeneous microstructure with WC grain size retained at the same values as in starting mixtures of different Co binder content.

3. The Co content in the starting mixture proved to have a significant influence on the electrochemical corrosion resistance of nanostructured hardmetals in acidic solution. A noticeable trend of polarization resistance $R_p$ decrease, and current density $i_{\text{corr}}$ and corrosion rate $v_{\text{corr}}$ increase has been established with increasing Co content. Nanostructured hardmetals with the grain size $d_{\text{WC}} < 200$ nm showed the same corrosion behavior as coarser grain-size conventional WC hardmetals depending on the Co content.

4. The chemical composition of the Co binder showed a significant influence. Samples with lower relative magnetic saturation related to lower added C content and more W dissolved in the Co binder showed better corrosion resistance. Significant differences in magnetic saturation for samples with the same Co content lead to more pronounced differences in the corrosion rates. A slight difference in magnetic saturation and WC grain size changed the Tafel curves.

5. Co content was shown to be the dominant influential factor governing electrochemical corrosion resistance of nanostructured hardmetals when compared to the chemical composition of the Co binder and WC grain size. Samples with lower Co content exhibited lower corrosion rates.
(6) The slight increase of GGl content, Cr$_3$C$_2$, and VC did not improve the corrosion resistance significantly for the samples with the same Co content. Higher content of Cr$_3$C$_2$ dissolved in the binder contributed to a lower corrosion rate. Slight VC increase did not contribute to corrosion resistance. Superior corrosion resistance is attributed to W and C dissolved in the Co binder, lower magnetic saturation, or WC grain size of the sintered sample.

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