Effect of 0.40 % and 0.20 % Titanium Addition on Microstructure Properties and Corrosion Behavior of CuAl₁₀Ni₅Fe₄ Alloy

Abubaker J. A. IRHAYIM¹, Ismail ESEN^{1*}, Selami SAGIROGLU¹, Esma KESKIN², Hayrettin AHLATCI³, Volkan KARAKURT⁴

¹Mechanical Engineering, Faculty of Engineering, Karabuk University, Karabuk 78050, Turkey

² BUMA Engineering and Consulting Inc., Ankara, 6980, Turkey

³ Metallurgical and Materials Engineering, Faculty of Engineering, Karabuk University, Karabuk 78050, Turkey

⁴ Sağlam Metal San. Tic. A.Ş, Kocaeli, Turkey

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This study examined how post-casting heat treatment (solution treatment and tempering) and forging affected the corrosion and microstructural properties of the nickel aluminium bronze (CuAl₁₀Ni₅Fe₄) with 0.20 % and 0.40% titanium addition. Heat-treated CuAl10Ni5Fe4 alloy microstructures showed copper-rich a-solid solution, martensitic \beta-phase, and intermetallic k-phases, such as flake-shaped kI, fine kIII, and black globular kIV. Adding 0.20 % and 0.40 % titanium to CuAl10Ni5Fe4 alloys reduced dendritic arm thickness and increased k phase abundance. Additionally, significant kIV and κ II precipitations were observed in the tempered β phase. Tempering β may cause martensitic β to split into tiny α and κ III phases. The grain refinement of the CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti, and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys was significantly noticeable after forging. The alloys retained the same α , β , and κ phases seen in the heat-treated alloys after forging. Titanium's refining effect increased the hardness of the alloy into which it was incorporated. Increased stiff phases, such as tempered β and fine κIV , contribute to increased hardness. The forging process enhanced the hardness of all three alloys. Forged and heat treated CuAl₁₀Ni₅Fe₄-0.40 % Ti alloy had the highest hardness with 243.50 HB and 256.18 HB, respectively. The microstructural phases α , β , and κ significantly impact the corrosion behaviour of CuAl₁₀Ni₅Fe₄ alloy. The CuAl₁₀Ni₅Fe₄-0.40 % Ti alloy after forging had the lowest 24-hour weight loss of 0.003326 mg/dm². The CuAl₁₀Ni₅Fe₄ alloy loses the most weight (0.013659 mg/dm²) following heat treatment (S.T.+T) after 24 hours. The minimal corrosion rate for the CuAl₁₀Ni₅Fe₄-0.40 % Ti alloy after forging was 0.003916 mg/(dm²·day) after 72 hours. Due to stratification, corrosion creates craters. Pitted or porous structures and craters are common. All alloys show a decrease in these structures after forging. The formation of the CuCl₂ phase is believed to occur in regions experiencing pitting corrosion.

Keywords: CuAl₁₀Ni₅Fe₄, heat treatment, forging process, corrosion behavior, microstructure.

1. INTRODUCTION

Bronze alloys have been a significant material in various industrial applications due to their exceptional properties such as corrosion resistance, mechanical strength, and durability. Nickel aluminum bronze (NAB) and manganese aluminum bronze (MAB) are two highly alloyed bronzes that have gained popularity for their resistance against corrosion, cavitation, erosion, and improved mechanical properties [1]. These alloys are extensively used in marine applications like marine propellers due to their high strength and excellent corrosion resistance [2]. The addition of nickel to aluminum bronze has been explored to enhance properties like corrosion resistance, hardness, tensile strength, and wear resistance [3]. Historically, bronze was the first intentionally man-made alloy, typically composed of copper and tin in varying proportions [4]. The transition from arsenical copper alloys to tin alloys marked the Bronze Age, where bronze became the predominant alloy [5]. The production of bronze involved smelting copper and tin ores to create the desired alloy [6-8]. Different regions and time periods exhibited variations in

Bronze alloys have also been studied for their microstructural characteristics and processing techniques. Research has focused on enhancing properties through processes like friction stir processing and post-heat treatments [11, 12]. Studies have shown that copper-based alloys, including bronze, offer superior corrosion resistance for maritime applications [13]. Furthermore, the use of highalloy bronzes like NAB and MAB has been highlighted for specific applications, showcasing their appropriateness and effectiveness [14]. Nickel aluminum bronze (NAB) is a copper-based alloy that typically contains aluminum, nickel, iron, and manganese elements [15]. The microstructure of NAB is crucial for its properties, including corrosion resistance, cavitation resistance, and mechanical strength [16]. The alloy's microstructure usually consists of α phase, β phase, and κ phase [15]. Studies have shown that the addition of elements like cobalt and titanium can influence the properties of NAB, affecting its corrosion and wear resistance [17]. Furthermore, the microstructure of NAB can

bronze compositions, with some containing additional elements like lead [9, 10].

^{*} Corresponding author: I. Esen

E-mail: iesen@karabuk.edu.tr

be modified through processes like friction stir processing, leading to the transformation of the initial coarse microstructure into a finer structure with improved properties [18]. The corrosion resistance of NAB is a significant aspect, especially in marine applications. The alloy's resistance to corrosion is attributed to the formation of a protective oxide film on its surface [19-21]. The development of protective surface layers and corrosion product films on NAB alloys has been identified as a critical factor in enhancing their corrosion resistance [19, 22]. Heat treatment has been found to impact the microstructure evolution and erosion-corrosion behavior of NAB in chloride solutions [23]. Additionally, the morphology, crystallography, and chemistry of phases in as-cast NAB play a crucial role in determining its properties [24].

The phase transition that takes place during heat treatment is the main factor that controls the microstructure of NAB allovs. When the temperature exceeds 1030 °C, the NAB exhibits the β phase. As the temperature decreases, the α phase begins to develop within the β phase. At a temperature of 930 °C, the KII phase begins to separate from the β phase. At a temperature of 860 °C, the κ IV phase forms as a result of precipitation from the α phase. The formation of the eutectoid $\alpha + \kappa III$ phase occurs when the residual β phase is cooled below 800°C. Aside from temperature, the pace at which something cools down is also crucial in determining the phase change and the ensuing composition of phases. If the cooling rate is not slow enough, the β phase may persist as the β' phase with a martensitic microstructure, leading to a detrimental impact on the corrosion resistance of the alloy. Therefore, the NAB is often subjected to annealing at a temperature of 675 °C in order to remove the remaining β phase [25-27]. Prior research has been conducted to examine the impact of heat treatment and hot working on the microstructure and mechanical characteristics of NAB alloys [28, 29]. Currently, there has been a scarcity of research conducted on the impact of heat treatment on the erosion-corrosion resistance of NAB alloys. Furthermore, the correlation between the microstructure of the NAB and its erosioncorrosion behavior has yet to be determined.

Titanium addition to nickel aluminum bronze alloys has been a subject of interest in various studies. investigated the impact of adding titanium as a grain refiner along with cobalt to nickel-containing aluminum bronze [17]. The study focused on properties in both cast and forged conditions. Incorporating titanium into nickel aluminum bronze alloys has shown potential for enhancing properties such as corrosion and wear resistance. Titanium, when utilized as a grain refiner, impacts the microstructure of NAB, consequently affecting its corrosion resistance [16]. Additionally, the design of the microstructure to enhance the corrosion and cavitation resistance of NAB often considers the inclusion of titanium [16]. Furthermore, the addition of titanium to nickel aluminum (Raney) alloys used for hydrogen electrodes in alkaline hydrogen-oxygen fuel cells has been investigated, indicating that titanium may play a role in improving the corrosion resistance of nickel aluminum alloys [30]. Moreover, the literature indicates that nickel aluminum bronze alloys are widely used in various applications, including ship propellers [15]. This

underscores the importance of understanding how different alloying elements, such as titanium, can influence the mechanical properties and corrosion resistance of these alloys.

Finally, the microstructure of nickel aluminium bronze plays an important role in defining its mechanical, corrosion, and cavitation resistance qualities. Several studies have investigated the impacts of various elements, heat treatments, and processing methods on the microstructure of NAB, highlighting the importance of microstructural design in improving the alloy's performance. The ability to improve properties including corrosion resistance, wear behaviour, and heat treatment techniques by adding up to 0.4 % titanium to nickel aluminium bronze alloys is investigated here, and the importance of alloy composition in adapting to industrial applications is demonstrated. This study examined the effects of post-casting heat treatment (solution treatment and tempering) and forging techniques on the corrosion and microstructural characteristics of nickel aluminium bronze $(CuAl_{10}Ni_5Fe_4)$ that had 0.20 % and 0.40 % Ti added.

2. MATERIALS AND METHODS

The casting studies for the fabrication of samples were conducted at the R&D Center of Saglam Metal Industry and Trade. A pre-prepared nickel aluminum bronze ingot was obtained specifically for use as a raw material in casting experiments, following the manufacturing parameters. Ingots manufactured in industrial settings were utilized as the primary material, whereas materials intended for laboratory research were generated. The ingot, which served as the raw material for casting experiments, was manufactured using the permanent mold casting process and consisted of pure copper scrap, pure aluminum, iron, and nickel. The melting process was conducted using an induction furnace with a capacity of 300 kg. The preproduced nickel aluminum bronze ingot was sliced and utilized for the melting process. The CuLi prealloy was employed to eliminate oxygen from the liquid metal, while a 30 % copper titanium prealloy was utilized to introduce titanium. The degassing procedure after melting nickel aluminum bronze involved the use of copper lithium prealloy. Three distinct charges were formulated, with one devoid of titanium, another having 0.20 % Ti, and the last one containing 0.40 % Ti. Additionally, 30 % copper titanium prealloy was introduced to the molten metal during the last phase of melting, after which the casting process commenced. The castings were conducted at a temperature of 1230 °C. Table 1 presents the chemical composition of CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40% Ti alloys. The chemical analysis of the ingot produced under industrial conditions was determined using an Oxford brand optical emission spectrometer at the facilities where the casting was carried out. The phases of the alloy were determined using X-ray diffractometry (XRD Rigaku Ultima IV; Japan) with a scanning range of $10-90^{\circ}$ and a scanning speed of 3°/min. Heat treatments were applied to the samples produced by the casting method, after solution treatment at 875 °C for 90 minutes and then tempering at 650 °C for 2 hours.

Table 1. Weight percentages of alloy compositions

	Chemical composition, wt.%						
Alloy	Al	Fe	Ni	Ti	Cu		
CuAl ₁₀ Ni ₅ Fe ₄	10.19	3.01	4.41	—	Bal.		
CuAl ₁₀ Ni ₅ Fe ₄ -0.20 % Ti	9.99	3.00	4.30	0.2	Bal.		
CuAl10Ni5Fe4-0.40 % Ti	9.87	2.99	4.27	0.4	Bal.		

Industrial circumstances were employed to conduct forging (F.) investigations utilizing a ram. The forging conditions were established at a temperature of 870 °C [29], which is known to yield the maximum strength values according to the literature, and a deformation rate of 80 % [31]. In order to achieve this objective, cylindrical metal bars of 68 mm in diameter were subjected to a forging process, resulting in a final section size of 27×27 mm, using a ram as a tool.

The CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti samples, which had undergone heat treatment and forging, were each cut into dimensions of $10 \times 20 \times 10$ mm. The cutting operation was carried out using a water-cooled band saw. Once the cutting procedure was finished, the samples underwent sanding and polishing using the Mikrotest brand automated apparatus. Various grit grits, including 320, 400, 600, 800, 1000, and 2500, were used throughout the sanding operation. After the sanding operation, the polishing was carried out using a 3 μ m Al₂0₃ liquid solution. The etching method utilized 5 grams of FeCl₃, 50 milliliters of HCl, and 100 milliliters of clean water. A LOM-Carl Zeiss light optical microscope was employed to observe the alterations in the grains inside the phase structure. Additionally, SEM-Carl Zeiss Ultra Plus scanning electron microscope and EDX (energy dispersive X-ray spectroscopy) were utilized to analyze and uncover any secondary phases. Steel balls with a diameter of 2.5 mm were subjected to a force of 187.5 N in order to measure their Brinell hardness.

The immersion corrosion experiment utilized cylindrical samples measuring $10 \times 10 \times 10$ in size. Each sample was subjected to surface area and weight measurements using a Precisa precision balance. The duration of immersion corrosion in a 3.5 % NaCl solution ranged from 12 to 72 hours. Chromic acid and ethanol, derived from 180 grams of CrO₃ and 1 liter of distilled water, were employed to eliminate corrosion residues from the sample surface on an hourly basis. The corrosion samples were first submerged in chromic acid for approximately 3 minutes, then washed with purified water to eliminate any remaining acid residue. The corrosion samples underwent a rinsing process using distilled water, followed by immersion in ethanol for a duration of about 2 minutes, and subsequently were dried. After the drying process, each corrosion sample was carefully weighed using precise scales and then exposed to immersion corrosion once again. The weight losses reported at each hour interval were converted from grams to milligrams as a result of the immersion corrosion process. The surface area of the corrosion sample was converted from mm² to dm². The milligram lost per square decimeter per hour was determined by dividing the average weight loss (mg) by the surface area (dm²). The weight reductions were determined

by subtracting the beginning value from the end amount. The weight loss after 24 h was determined by subtracting the initial weight/surface area measurement from the weight/surface area measurement taken after 24 h. The corrosion rate was measured in milligrams per square decimeter per day (mg/dm²×day). The corrosion rates per day were determined by dividing the weight losses by the duration in days. To determine the corrosion rate corresponding to a 3-day period, the weight loss after 72 h is divided by 3.

3. RESULTS AND DISCUSSION

3.1. XRD patterns

X-ray diffraction (XRD) is a powerful technique used to analyze the crystal structure of materials. In the case of the CuAl₁₀Ni₅Fe₄ alloy, the presence of κ phases has been noted to enhance its mechanical strength without compromising ductility [32]. This indicates that the alloy's mechanical properties are influenced by specific phases within its structure. To further understand the crystallographic characteristics of the CuAl₁₀Ni₅Fe₄ alloy, XRD analysis has been employed to identify the different phases present and their distribution within the material. In the XRD analyses given in Fig. 1, copper-rich AlCu₄ and α-Cu phases were present in the highest peak, while the α -Cu phase was seen in the final peaks of all alloys. When the peaks of the CuAl₁₀Ni₅Fe₄ alloy are examined in Fig. 1 a, it is seen that it starts with AlNi and AlFe₃ phases at 30.95°. In Fig. 1 b, in the CuAl₁₀Ni₅Fe₄-0.20%Ti alloy, the AlTi phase was seen at 21.90° with the presence of titanium, and immediately afterwards the AlFe₃ phase was seen at 27.25°. In the CuAl₁₀Ni₅Fe₄-0.40%Ti alloy in Fig. 1 c, the AlTi phase was first seen at 20.80°, and then the AlCu₄ phase together with the AlFe₃ phase was seen at 27.50°.

3.2. Microstructure

Fig. 2 shows the LOM images of CuAl₁₀Ni₅Fe₄. CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys after solution treatment and tempering after casting, while Fig. 3 shows the LOM images of these alloys after forging. One of the factors that leads to grain refining with the addition of elements is the rise in the rate at which individual grains form before there is an increase in the number of grain nuclei. The formation of smaller grains occurs when the number of nucleation sites per unit volume rises [33]. The inclusion of titanium is believed to have a grain refining impact by increasing the number of nucleation sites, resulting in the creation of smaller grains. The microstructure shown in Fig. 3 a of the heat-treated CuAl₁₀Ni₅Fe₄ alloy is a common representation found in several research on nickel-bearing aluminium bronze. The depiction in Fig. 4 helps clarify the presence of a copperrich α -solid solution phase, a martensitic β -phase, and various intermetallic κ -phases, including leaf-shaped κI , thin κIII and black globular κIV [34–36].



Fig. 1. X-ray diffraction patterns: a – CuAl₁₀Ni₅Fe₄, b – CuAl₁₀Ni₅Fe₄-0.20 % Ti; c – CuAl₁₀Ni₅Fe₄-0.40 % Ti

The depiction in Fig. 4 helps clarify the presence of a copper-rich α -solid solution phase, a martensitic β -phase, and various intermetallic κ -phases, including leaf-shaped κI , thin κIII and black globular κIV [34–36].

Fig. 2 b and c show that the addition of 0.20 % and 0.40 % titanium to CuAl₁₀Ni₅Fe₄ alloys leads to a decrease in the dendritic arm thickness and an increase in the amount of certain types of κ phases. Additionally, a substantial amount of κ IV and κ II phase precipitation was observed within the tempered B phase. The tempering of β is believed to cause the martensitic β to decompose into extremely small masses of α and κ III [34].

The decomposition of the martensitic β phase into α and κ III phases during the tempering of the CuAl₁₀Ni₅Fe₄ alloy entails the reconstitution of martensite to the high-temperature bcc β phase, which subsequently decomposes into NiAl and α . In Cu-Al-Ni alloys, tempering at 675 °C

results in the reconstitution of martensite into the hightemperature β phase, which subsequently decomposes into NiAl and α phases [24].



Fig. 2. LOM images after solution treatment and tempering after casting: a-CuAl₁₀Ni₅Fe₄; b-CuAl₁₀Ni₅Fe₄-0.20 % Ti; c-CuAl₁₀Ni₅Fe₄-0.40 % Ti

The procedure encompasses pro-eutectoid $\gamma 1$ precipitation and eutectoid disintegration, succeeded by dissolution at elevated temperatures [37, 38]. Below 300 °C, a demixing event produces tiny coherent

precipitates, resulting in elevated transition temperatures without altering hysteresis [39].







Fig. 3. LOM images after forging: $a-CuAl_{10}Ni_5Fe_4$; $b-CuAl_{10}Ni_5Fe_4$ -0.20 % Ti; $c-CuAl_{10}Ni_5Fe_4$ -0.40 % Ti

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The breakdown process is influenced by elements like elastic stresses from quenching and the level of ordering in the alloy structure [38]. The microstructures of CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys were noticeably reduced in thickness after the forging process (Fig. 3). The alloys retained the identical α , β , and κ phases observed in Fig. 2 after the forging process. As a result of the forging process, the dendritic

arms were fractured and transformed into almost spherical shapes. Additionally, the forging caused the formation of new κ IV phases.



Fig. 4. Illustrations depicting the several stages of a type 80-10-5-5 cast aluminum bronze [34, 35]

The alloys with 0.20 % and 0.40 % titanium addition to the CuAl₁₀Ni₅Fe₄ alloy exhibited separate KI and KIII phases. Titanium serves as an inoculant, efficiently enhancing the alloy's microstructure by facilitating the development of finer dendrites. This effect has been shown in many investigations on copper alloys, demonstrating that the incorporation of titanium results in a refined microstructure marked by reduced secondary dendrite arm spacing and therefore finer dendritic characteristics [32]. This refining enhances the alloy's mechanical characteristics, since smaller dendrites often increase tensile strength and ductility. Furthermore, titanium's impact on the production and distribution of κ phases within the CuAl₁₀Ni₅Fe₄ alloy structure is essential. The κ phases enhance the mechanical integrity and wear resistance of the alloy, with their production being sensitive to the titanium content. Research findings demonstrate that titanium additions can promote the nucleation of diverse κ intermetallic phases, which display unique morphologies and distributions within the α -Cu matrix [18]. Increased titanium concentration correlates with a higher occurrence of globular κ phase forms, which can boost toughness and mechanical qualities under stress by redistributing stress throughout the alloy [40].

Fig. 5 shows the SEM images of CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl10Ni5Fe4-0.40 % Ti alloys after solution treatment and tempering after casting, while Fig. 6 shows the SEM images of these alloys after forging. Table 2 and Table 3 show the EDX analysis of the second phase with distinct morphologies labeled as (1-12) in Fig. 5 and Fig. 6, respectively. The overall SEM images show the microstructure of the alloys consisting of α , β and a series of intermetallic κ phases. The α phase is a copperenriched solid solution, and the β phase (or retained β phase) is a solid solution with a tempered structure and a highdensity AlNi precipitate area. There are three types of κ phases, which are distinguished by the combination of their morphologies and distributions. The kI phase is enriched in iron and is found in the α phase matrix, which typically has a rosette-shaped morphology. The kIV phase is mainly distributed at the α and β boundaries in a spherical-like shape. The κ IV phase is much finer than the κ I phase.







Fig. 5. SEM images after solution treatment and tempering after casting: a-CuAl₁₀Ni₅Fe₄; b-CuAl₁₀Ni₅Fe₄-0.20 % Ti; c-CuAl₁₀Ni₅Fe₄-0.40 % Ti

The lamellar κ III phase, which is rich in nickel and forms at the α and β boundaries, is the product of the eutectoid transformation of the β phase at low temperatures. The κ IV phase, which is rich in iron, is a finer precipitate within the α phase [21, 41, 42]. The initial cast microstructure of the CuAl₁₀Ni₅Fe₄ alloy consists of a combination of several phases, including a copper-rich α solid solution, a β solid solution, and many intermetallic κ phases [32].









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The forging process utilises heat and mechanical energy to alter the solid-state structures of the alloy. The κ -phase distribution may be greatly modified; during forging, the stability of several phases, including the κIV phase, is crucially influenced by the temperatures and strain rates applied. The temperature conditions encountered during hot forging frequently enhance diffusion and enable phase changes that may stabilise or result in the emergence of new κ phases.

Points	Al	Ti	Mn	Fe	Ni	Cu
1	6.15	-	0.90	-	5.54	87.42
2	1.45	—	1.07	81.11	2.50	13.89
3	6.13	—	1.16	1.19	4.54	86.97
4	4.47	—	1.23	1.25	4.99	88.07
5	6.93	0.06	1.16	-	4.55	87.30
6	7.79	0.11	0.65	—	5.81	85.63
7	1.32	0.08	0.45	2.67	1.40	94.09
8	6.77	0.02	0.75	0.37	6.22	85.88
9	3.51	18.17	0.33	36.96	6.07	34.96
10	4.66	0.23	-	-	4.82	90.29
11	2.21	0.32	2.39	9.55	9.20	76.34
12	16.53	0.36	_	_	4.89	78.21

Table 2. The weight percentage findings of Fig. 5 as determined by EDX

Table 3. The weight percentage findings of Fig. 6 as determined by EDX

Points	Al	Ti	Mn	Fe	Ni	Cu
1	8.27	-	0.10	0.81	5.60	85.22
2	7.30	—	0.03	0.72	5.01	86.95
3	4.37	—	—	0.32	5.41	89.91
4	6.65	—	0.43	0.22	5.77	86.93
5	6.21	0.06	0.30	1.57	5.55	86.34
6	5.71	0.16	0.24	1.27	5.62	87.00
7	2.81	0.11	1.36	3.87	4.53	87.26
8	6.44	0.14	0.25	0.98	6.31	85.88
9	7.37	0.10	0.41	0.19	6.11	85.85
10	7.70	0.36	1.66	7.12	6.69	76.47
11	2.00	28.09	_	11.38	6.01	52.52
12	10.12	0.38	0.50	6.68	14.13	68.19

Furthermore, the elevated temperature conditions during the forging process might provoke alterations in the pre-existing intermetallic phases. Studies indicate that the emergence of new κ phases, especially κIV , is frequently promoted under dynamic recrystallisation circumstances when the solid solution experiences significant mechanical strain.

Furthermore, the fragmentation and dispersion of coarse particles within the microstructure during the forging process enhance the uniformity of stress distribution. The dynamic recrystallisation process enabled by forging improves the mixing and distribution of alloying elements, hence boosting the overall stability of the KIV phase. The rise in dislocation density and strain can facilitate the formation of nucleation sites for these new phases. The use of elements like as chromium and beryllium facilitates the development of κ phases, notably κIV , during hot forging [43]. Hot forging modifies the microstructure relative to the as-cast state, with torsion experiments replicating the forging process at different temperatures and strain rates [44]. Thermal-mechanical activities, including hot rolling and quenching, impact the development of martensite phases, hence influencing the shape memory effect and the overall characteristics of the alloy [45]. These mechanisms facilitate the formation of several k phases in conjunction with an α matrix and other intermetallic phases, influencing the alloy's mechanical and corrosion characteristics [43].

3.3. Hardness test results

Fig. 7 shows the hardness of $CuAl_{10}Ni_5Fe_4,$ $CuAl_{10}Ni_5Fe_4-0.20$ % Ti and $CuAl_{10}Ni_5Fe_4-0.40$ % Ti

alloys after solution treatment and tempering after casting and after forging in a comparative manner. The purpose of the comparison is to analyze the differences in hardness between the different alloys and processing methods. The hardness values of the $CuAl_{10}Ni_5Fe_4$ alloy were found to be lower in comparison to other alloys, when titanium was not added. Grain refining plays a crucial role in improving the hardness characteristics of alloys.





The enhanced hardness is ascribed to the grain refining phenomenon caused by titanium, resulting in a decrease in grain size and an augmentation in dislocation density [46, 47]. In addition, the inclusion of solid particles in the

alloy as a result of the introduction of grain refiners such as titanium helps to increase the material's hardness [48]. Furthermore, the augmentation in the quantity of rigid phases, namely tempered β and fine κ IV, is believed to be another contributing factor to the rise in hardness. Although the process of forging greatly enhanced the hardness of all three alloys, it was observed that the hardness values of CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys were similar to each other, however greater than that of the CuAl₁₀Ni₅Fe₄ alloy. Considering the difference in hardness values after solution treatment and tempering after casting and after forging, it was observed that CuAl₁₀Ni₅Fe₄-0.40 % Ti alloy showed the highest value with 243.50 HB and 256.18 HB values, respectively.

The hardness of the CuAl₁₀Ni₅Fe₄ alloy is affected by the α , β , and κ phases, which are determined by the composition and processing conditions. Nickel and iron enhance the stability of the β phase, whereas unidirectional solidification fosters a greater proportion of the α phase relative to gradual cooling [49]. Thermal parameters during solidification influence microstructure, hardness, and microhardness, with elevated aluminium content resulting in enhanced hardness and a more refined structure [50]. Ageing may reduce hardness owing to the production of precipitates and influence grain size in CuAlNi alloys [51].

3.4. Immersion corrosion test results

Fig. 8 shows the comparative 24-hour weight loss changes of $CuAl_{10}Ni_5Fe_4$, $CuAl_{10}Ni_5Fe_4$ -0.20 % Ti and $CuAl_{10}Ni_5Fe_4$ -0.40 % Ti alloys after solution treatment + tempering and after forging processes. The corrosion rates of these alloys after 72 hours are also given in Fig. 9.



Fig. 8. Comparative immersion corrosion weight loss results of CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys. S.T. – solution treatment; T. – tempering; F. – forging

The alloy with 0.40 % Ti addition exhibited the best corrosion behavior after solution treatment + tempering and after forging processes. While the minimum weight loss after 24 hours was found to be 0.003326 mg/dm^2 in the CuAl₁₀Ni₅Fe₄-0.40 % Ti alloy after forging process. The maximum weight loss in CuAl₁₀Ni₅Fe₄ alloy after solution treatment + tempering was found to be 0.013659 mg/dm^2 after 24 hours. The minimum corrosion rate in CuAl₁₀Ni₅Fe₄-0.40 % Ti alloy after forging was found to be $0.003916 \text{ mg/(dm}^2 \cdot \text{day})$ after 72 hours. The corrosion

behavior of CuAl₁₀Ni₅Fe₄ alloy is significantly influenced by its microstructural phases, specifically the α , β , and κ phases. Understanding the roles of these phases is crucial for enhancing the alloy's corrosion resistance, particularly in aggressive environments such as saline solutions. The α phase, which is the primary phase in the CuAl₁₀Ni₅Fe₄ alloy, generally exhibits good corrosion resistance due to its stable microstructure.



Fig. 9. Comparative immersion corrosion rates result of CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys. S.T. – solution treatment; T. – tempering; F. – forging

However, the presence of the β phase can lead to localized corrosion phenomena. The β phase, being more electrochemically active, can act as a galvanic cathode, promoting microgalvanic corrosion when it is discontinuously distributed within the matrix. This behavior is similar to findings in other alloys, where the β phase's volume fraction and distribution significantly affect corrosion rates and mechanisms [52, 53]. The κ phase, on the other hand, has been reported to enhance the mechanical properties of the CuAl₁₀Ni₅Fe₄ alloy without compromising its ductility. This phase contributes to the overall strength of the alloy, which can indirectly influence its corrosion resistance by improving the integrity of the microstructure under corrosive conditions [17, 32]. The presence of the κ phase can also help in reducing the susceptibility to stress corrosion cracking, as it may inhibit the formation of microcracks that can serve as initiation sites for corrosion. Moreover, the interaction between these phases can lead to complex corrosion behaviors. For instance, the presence of the β phase can lead to preferential anodic dissolution, while the κ phase can mitigate this effect by providing a more uniform microstructure that resists localized attack.

The corrosion resistance of nickel-aluminum bronze alloys can be improved by several processing techniques and alloying elements. The hot shaping of CuAl10Ni5Fe4 alloy enhances corrosion resistance by spheroidising and dispersing κ phases, hence diminishing selective phase corrosion [17]

The presence of specific phases in alloys can significantly influence their corrosion behavior. Furthermore, the corrosion potential of different phases within an alloy can vary, influencing the overall corrosion resistance [54]. In the context of bronze alloys, chloride ions have been identified to cause preferential corrosion of the Cu-rich phase, while the $\alpha + \beta$ eutectic phase is susceptible to corrosion under specific oxygen conditions [55]. Furthermore, in nickel aluminum bronze, the microstructure has been found to influence crevice corrosion initiation, with specific phases such as the κ_{III} phase playing a role in corrosion within the microstructure [21]. The presence of CuO phase plays a significant role in the corrosion behavior of copper-based materials. Studies have shown that CuO is a predominant constituent in the corrosion product film formed on copper alloys after exposure to corrosive environments [56-58]. The formation of CuO has been linked to enhanced corrosion resistance due to its ability to reduce the number of corrosion sites and slow down the corrosion rate [59]. Additionally, CuO has been reported to be a stable species in neutral solutions, contributing to the formation of corrosion products like Cu₂O, which further influences the corrosion process [58]. Copper oxide (Cu_2O) is involved in the corrosion process of copper-based materials. Platzman et al. proposed a three-stage oxidation mechanism where a Cu₂O layer forms and then transforms into a more stable CuO layer [60]. The initial formation of Cu₂O is significant as it acts as a reservoir of Cu+ ions, triggering catalytic corrosion cycles [61]. Cu₂O has been observed to agglomerate into CuO particles during oxidation processes [62]. The presence of Cu₂O on copper materials influences their corrosion behavior, with corrosion products often being composed of $Cu_2O[63-65]$. The formation of Cu₂O can initiate corrosion cells, with the Cu₂O film acting as the cathode, contributing to pitting corrosion [66]. Cu₂O demonstrates high corrosion resistance and stability in various environments, enhancing protective properties [67]. In copper-nickel alloys, the incorporation of nickel ions in the Cu₂O barrier layer enhances corrosion resistance by reducing ionic conductivity within Cu₂O [68]. Cu₂O has also been linked to the corrosion resistance of copper under different conditions, such as in the presence of organic acids [69]. The protective role of Cu₂O patina in increasing corrosion resistance contrasts with the detrimental effects of CuCl₂ patina [69]. The corrosion resistance of copper alloys can vary significantly depending on the presence of different phases and elements within the alloy matrix. Additionally, the presence of specific phases such as Cu₂O and Cu-Al intermetallic phases can play a crucial role in either enhancing or reducing the corrosion resistance of copper alloys [70, 71]. Moreover, copper alloying elements, including the presence of grain boundaries and precipitates, can significantly impact the microstructure and corrosion behavior of the material. For example, the scarcity of alloying elements such as copper can keads to the formation of grain boundary segregation and increase the susceptibility of the alloy to intergranular corrosion [72, 73]. On the other hand, the presence of certain copper-rich precipitate phases, such as the τ -phase, can enhance the corrosion resistance by forming protective layers that inhibit corrosion processes [73]. Rahman et al. discussed the specific appearances of corrosion such as intercrystalline corrosion and corrosion cracking in copper alloys, emphasizing the effect of alloying elements on corrosion behaviour [74]. The presence of TiO phase in copper alloys has been shown to have a significant impact on their corrosion resistance. demonstrated that the addition of a TiO layer improved the corrosion resistance of Ticontaining alloys compared to those without Ti when exposed to elevated temperatures in molten salt environments [75]. The phase of TiO₂ plays a crucial role in influencing the corrosion behavior of copper alloys. Studies have shown that the corrosion rate of copper alloys can be affected by the presence of TiO₂. For instance, research by Saber et al. highlighted that Cu/Al₂O₃ composites exhibited lower corrosion rates compared to pure copper and copper coated with TiO₂ [76]. Moreover, the composition of copper alloys, including the presence of alloying elements, can also influence their corrosion resistance. This suggests that TiO plays a crucial role in enhancing the corrosion resistance of copper alloys under specific conditions. The presence of the Ti₃O₅ phase in copper alloys can significantly impact their corrosion behavior. Studies have shown that the formation of intermetallic compounds like Ti₂Cu can create an "envelope effect," which acts as a barrier preventing corrosive liquids from directly contacting the alloy, thereby enhancing its corrosion resistance [77]. Additionally, the equilibrium between Ti and O in alloys can lead to the solubility of other elements like Fe, Cr, and Ni in the Ti₃O₅ phase, affecting the overall corrosion properties of the material [78]. The presence of the Ti₂O phase in copper alloys has a significant impact on their corrosion behavior. Studies have shown that heat treatment can lead to the redistribution of copper elements and the precipitation of the Ti₂Cu phase, resulting in improved mechanical properties and corrosion resistance properties [77]. In the XRD analyses given in Fig. 10, while Cu₂O rich in copper is commonly found in the highest peak (42.60°); TiO, Ti₂O₃ and Ti₃O₅ phases are observed in this peak with titanium addition. The terminating peaks of the alloys are at 86.60° and Cu₂O and CuO phases are observed in the alloy without titanium addition; Ti_2O , Ti_2O_3 and Ti_3O_5 phases are observed in this peak with titanium additions. When the peaks of the CuAl₁₀Ni₅Fe₄ alloy are examined in Fig. 10 a, it starts with CuCl₂ at 18.45° and then the Cu₂O phase appears at 29.30°. In Fig. 10 b, CuAl₁₀Ni₅Fe₄-0.20 % Ti alloy started with CuCl₂ phase at 15.50° and immediately afterwards Ti₃O₅ phase was observed at 20.55°. In Fig. 10 c, in CuAl₁₀Ni₅Fe₄-0.40 % Ti alloy, CuCl₂ and Ti₃O₅ phases were observed at 16.60° while CuCl₂ and Ti₂O phases appeared at 21.70°.

Fig. 11 shows the SEM images of CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys corroded after solution treatment and tempering after casting, while Fig. 12 shows the SEM images of these alloys corroded after forging. Table 4 and Table 5 show the EDX analysis of the second phase with distinct morphologies labeled as (1-6) in Fig. 11 and Fig. 12, respectively. When the general SEM images are examined, it is seen that there are crater formations by layering in places. The frequency of pitted/porous structures and crater formations is remarkable in the images. A decrease in the frequency of these structures was observed in all alloys after forging. It is thought that CuCl₂ phase is formed in the areas where pitting corrosion occurs. In CuAl₁₀Ni₅Fe₄ (Fig. 11 a) alloy, which has been corroded after solution treatment and tempering after casting, branching-like pittings are observed in the pit at point 1. In EDX analyses, the highest Cl ion is observed in this region and therefore, it is possible to say that although CuCl₂ phase is dense at this point, a protective barrier for corrosion is formed with the formation of CuO phase in the region. The crater formations at point 2 are thought to be Cu₂O corrosion phase. In CuAl₁₀Ni₅Fe₄-0.20 % Ti (Fig. 11 b) alloy, which has been corroded after solution treatment and tempering after casting, small pits are found in a sediment-like formation at point 3. Here, Ti₂O₃ and Ti₂O phases are thought to have formed, while TiO phase is assumed to have formed at point 4.



Fig. 10. X-ray diffraction patterns after corrosion: $a-CuAl_{10}Ni_5Fe_4$; $b-CuAl_{10}Ni_5Fe_4$ -0.20 % Ti; $c-CuAl_{10}Ni_5Fe_4$ -0.40 % Ti

Here, Ti_2O_3 and Ti_2O phases are thought to have formed, while TiO phase is assumed to have formed at point 4. In CuAl₁₀Ni₅Fe₄-0.40 % Ti (Fig. 11 c) alloy, which has been corroded after solution treatment and tempering after casting, Ti_2O and TiO are formed at point 5, while the stratified crater formation at point 6 is likely to be Ti_3O_5 and Ti_2O_3 . After forging, CuAl₁₀Ni₅Fe₄ (Fig. 12 a) alloy that has been corroded, at point 1, in the area where there are sometimes round and sometimes linear pitting on the surface, it is thought that CuCl₂ and Cu₂O phases are formed. At point 2, CuO and Cu₂O phases are likely to form. After forging, CuAl₁₀Ni₅Fe₄-0.20 % Ti (Fig. 12 b) alloy that has been corroded, it is thought that the cause of the precipitation at point 3 is the $CuCl_2$ phase and the reason for this precipitation being irregular is the formation of the TiO_2 phase here.



Fig. 11. SEM images of corrosion after solution treatment and tempering after casting: a-CuAl₁₀Ni₅Fe₄; b-CuAl₁₀Ni₅Fe₄-0.20 % Ti; c-CuAl₁₀Ni₅Fe₄-0.40 % Ti

The crater formation on the surface at point 4 is attributed to the TiO phase. After the forging process, it is thought that the TiO₂ phase is formed in contrast to the CuCl₂ phase in the pit-like structure that occurred at point 5 in the CuAl₁₀Ni₅Fe₄-0.40 % Ti (Fig. 12 c) alloy that has been corroded.

				-		-	-	
Points	Al	Ti	Mn	Fe	Ni	Cu	Cl	0
1	10.65	-	0.86	15.20	5.76	1.56	15.14	50.77
2	12.83	-	1.58	25.44	19.85	28.26	2.36	9.71
3	6.61	6.91	0.48	15.16	11.05	44.18	1.32	14.32
4	11.29	0.39	1.31	11.73	11.74	59.41	0.60	3.55
5	10.07	0.23	0.72	8.78	11.73	64.34	0.33	3.83
6	10.87	7.24	0.37	11.60	11.12	41.96	0.55	16.32

Table 4. The weight percentage findings of Fig. 11 as determined by EDX

Table 5. The weight percentage findings of Fig. 12 as determined by EDX

Points	Al	Ti	Mn	Fe	Ni	Cu	Cl	0
1	11.11	-	1.24	27.72	12.10	43.20	1.92	2.73
2	16.76	-	0.53	14.40	23.87	34.81	0.87	8.78
3	7.48	1.26	1.29	10.35	7.55	65.52	0.54	7.17
4	14.92	0.29	1.38	13.21	20.14	41.01	0.20	8.85
5	11.25	5.08	3.40	29.21	13.64	34.30	0.54	2.59
6	11.63	7.70	1.32	20.67	15.10	35.16	0.45	7.98





а

b

At point 6, there is a regular crater formation and as can be seen from the EDX analysis, it is likely that this formation consists of Ti_3O_5 and Ti_2O_3 phases, which are titanium-rich phases. Investigations into copper corrosion in chloride-containing environments laden settings uncover intricate processes and the development of protective layers. In dilute oxychloride solutions, copper initially dissolves as CuCl2-, thereafter undergoing hydrolysis and oxide production on the surface [79].



Fig. 12. SEM images of corrosion after forging: a-CuAl₁₀Ni₅Fe₄; b-CuAl₁₀Ni₅Fe₄-0.20 % Ti; c-CuAl₁₀Ni₅Fe₄-0.40 % Ti

Protective scales including CuO or Cu2O can rapidly develop, becoming progressively less porous over time [80]. On Cu-Ni-Fe alloys, a dual-layer protective film develops: a substantial outer layer predominantly composed of Cu₂O and a slender interior layer enriched with chloride, oxygen, and [81]. The maximum chloride concentration is frequently detected near the boundary between these layers. Although alkanethiol layers can safeguard copper against corrosion, their efficacy diminishes with increased chloride concentrations and at potentials conducive to the formation of Cu(II) oxide [81].

4. CONCLUSIONS

In this study, it was investigated how the corrosion and microstructural properties of 0.20 % and 0.40 % titanium added nickel aluminum bronze (CuAl₁₀Ni₅Fe₄) were affected by post-casting heat treatment (solution treatment and tempering) and forging methods. The results are as follows:

The microstructures of the heat-treated CuAl₁₀Ni₅Fe₄ alloy showed the presence of a copper-rich α -solid solution phase, a martensitic β -phase, and various intermetallic κ -phases including flake-shaped κI , fine κIII , and black globular κIV . It is shown that the addition of 0.20 % and

0.40 % titanium to CuAl₁₀Ni₅Fe₄ alloys resulted in a decrease in the dendritic arm thickness and an increase in the amount of certain types of κ phases. In the microstructures of CuAl₁₀Ni₅Fe₄, CuAl₁₀Ni₅Fe₄-0.20 % Ti, and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys, a significant decrease in grain size and a transformation of broken dendritic arms into almost spherical shapes were observed after forging.

The CuAl₁₀Ni₅Fe₄ alloy has lesser hardness than other alloys without titanium. Titanium reduces grain size and increases dislocation density, increasing hardness. The increase in stiff phases, such as tempered β and fine κIV , may also contribute to the increase in hardness. Forging and heat treatment increased the hardness of all three alloys, however CuAl₁₀Ni₅Fe₄-0.20 % Ti and CuAl₁₀Ni₅Fe₄-0.40 % Ti alloys had similar hardness values but were harder than CuAl₁₀Ni₅Fe₄.

After 24 hours, the forged $CuAl_{10}Ni_5Fe_4$ -0.40 % Ti alloy experienced 0.003326 mg/dm² weight loss, while the $CuAl_{10}Ni_5Fe_4$ alloy experienced 0.013659 mg/dm² weight loss after solution treatment and tempering. The minimal corrosion rate in $CuAl_{10}Ni_5Fe_4$ -0.40 % Ti alloy after forging was 0.003916 mg/(dm²·day) after 72 hours. Pitted/porous structures and craters are observed as corrosion damages. All alloys have less corrosive damages after forging. It is believed that the $CuCl_2$ phase forms in regions of pitting corrosion.

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