Role of magnesium and minor zirconium on the wear behavior of 5XXX series aluminum alloys under different environments

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ROLE OF MAGNESIUM AND MINOR ZIRCONIUM ON 
THE WEAR BEHAVIOR OF 5XXX SERIES ALUMINUM 
ALLOYS UNDER DIFFERENT ENVIRONMENTS 

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Abstract: The tribological performance of 5xxx series aluminum alloys with ternary zirconium is evaluated at ambient conditions under dry, wet and saline environment. The experiment has been done using a Pin-on-Disk apparatus under an applied load of 20N. The sliding distances varies ranging from 116 m-2772 m at a sliding velocity of 0.385 ms-1. The results show that presence of Mg and Zr into this alloy helps to increase their strength and wear resistance under dry sliding condition. But they significantly suffer under wet and corrosive environment due to formation of β-phase Al3Mg2, to slip bands and grain boundaries which may lead to stress-corrosion cracking. The variation of friction coefficient is observed in wet and corrosive environment due to the formation of oxidation film, lubrication, and corrosion action in solution. The SEM analysis shows that brittle Al3Mg2 phase initiate the fracture surface for Al-Mg alloy and Zr addition accelerate the brittleness of the alloy owing the fine precipitates of Al3Zr.

Keywords: aluminum alloys, environment, wear, corrosion, friction, microstructure

1. INTRODUCTION

Pure aluminum is soft, ductile, corrosion resistant and has a high electrical conductivity [1, 2]. It is widely used for foil and conductor cables, but alloying with other elements is necessary to obtain the higher strengths needed for other applications. Aluminum is most commonly alloyed with copper, zinc, magnesium, silicon, manganese and lithium. Small additions of chromium, titanium, zirconium, scandium, lead, bismuth and nickel are also made and iron is invariably present in small quantities [3-5]. The addition of magnesium to aluminum increases the strength through solid solution strengthening and improves their strain hardening ability. These alloys are the highest strength non-heat-treatable aluminum alloys [6-8]. Al-Mg casting alloys have a wide range of application, especially in the automotive and ship borne industry that is directly related to their good mechanical properties [9, 10]. These alloys are characterized by relatively good castability and are distinguished by excellent corrosion resistance due to high magnesium content [11, 12]. The alloys may exhibit some instability in properties, which is manifested in two ways. If the magnesium content exceeds 3 to 4 wt%, there is a tendency for the formation β-phase Al3Mg2, to precipitate in slip bands and grain boundaries which may lead to intergranular attack and stress-corrosion cracking in corrosive conditions [3, 13]. Precipitation of β occurs only slowly at ambient temperatures but is accelerated if the alloys are in a heavily worked condition, or if the temperature is raised. Small additions of Zr, Ti, Sc raise the recrystallisation temperatures, also increase tensile properties for given magnesium content [14-16]. Zirconium is added to aluminum alloys for altering recrystallization behavior, enhancing mechanical properties and thermal stability while retaining electrical conductivity and controlling


the grain size during solidification. In the first two cases, super saturation of the Al solid solution is achieved during solidification and the metastable Al\textsubscript{3}Zr phase precipitates during annealing and pinning dislocations and subgrain boundaries. In the last case, primary crystals of the equilibrium Al\textsubscript{3}Zr phase act as nucleating substrates for Al grains [17, 18].

Most recent, the Al-Mg alloy has been extensively used in modern structural and functional applications. Proper reinforcement increases the wear properties of the alloys. There is currently no work available which has studied the wear behavior of Al-Mg alloy at different environment. The purpose of this study is thus to investigate and report on the combined effect of Mg and Zr on the wear behavior of commercially pure aluminum in dry, wet and corrosive environment.

2. EXPERIMENTAL PROCEDURE

The materials used in the present study were commercially pure Al, 5xxx series Al-Mg alloy and Zr added 5xxx series Al-Mg-Zr alloy. These alloys were prepared using pure aluminum ingot (99.7%), industrial pure magnesium (99.9%) and master alloys of Al–10% Zr. Melting was carried out in a resistance heating furnace. During melting degasser, borax etc. was used as flux cover into the clay-graphite crucible. The temperature of the melt was always maintained at 780±15°C. Then the melt was allowed to be homogenised under stirring at 700°C and poured in a mild steel mould size of 17×150×250 in millimeter. Before pouring the material the mild steel mould was coated inside with a film of water-craft and preheated at 200°C. The cast samples were first machined to skin out the oxide layer from the surface. All the alloys were analysed by wet chemical analysis and spectroscopy method simultaneously to determine the chemical composition. The chemical compositions of the alloys are given in Table 1.

| Tab. 1. Chemical composition of the experimental alloys in weight percent |
|--------------------------|--------------------------|--------------------------|--------------------------|
| Alloy       | Al      | Al-Mg   | Al-Mg-Zr   |
| Mg          | 0.006   | 5.014   | 5.124      |
| Zr          | 0.000   | 0.000   | 0.304      |
| Si          | 0.835   | 0.429   | 0.507      |
| Fe          | 0.627   | 0.375   | 0.269      |
| Sn          | 0.000   | 0.249   | 0.238      |
| Zn          | 0.052   | 0.012   | 0.012      |
| Ni          | 0.000   | 0.007   | 0.008      |
| Pb          | 0.000   | 0.003   | 0.004      |
| Al          | Bal     | Bal     | Bal        |

Microhardness of the cast test piece of the investigated alloys was measured with a Micro Vickers Hardness Tester. The Knoop indenter was applied with 1000 gm load for 10 s. An average of ten concordant readings was taken as the representative hardness of a sample. Density of the experimental alloys was calculated from the chemical composition of the alloys. Tensile tests was carried out at room temperature in an Instron testing machine using cross head speed to maintain the strain rate of $10^{-3}$ s. The samples used were according to American Society for Testing and Materials specification [19]. Tensile test was determined using at least five test pieces for each test. The sample of 12 mm length and 5 mm diameter were machined from the alloys for wear study by following ASTM G99-05 [20]. The end surface (5 mm diameter) of the pin samples were polished using emery papers 1, 0, 2/0. Afterwards, the samples were polished in a fine grade wheel polisher. Later, the end surface was cleaned in running water and finally dried in acetone. Hardened mild steel discs were used as the counter-body material. The hardness of the discs was around RC 50. One of the surfaces of the disc was grinded by surface grinding machine and cleaned with cotton. Surface roughness of the disc was 31 μm. The frictional and wear behaviours of the experimental alloys were investigated in a pin-on disc type wear apparatus by following ASTM Standard G99-05.

During the dry wear tests, the end surface of the pin samples were pressed against horizontal rotating mild steel disc. Applied load of 20 N was used throughout the test, which yielded nominal contact pressures of 1.02 MPa. The tests were conducted at the sliding speed of 0.385 ms\(^{-1}\) with varying sliding distances ranging from 116 m-2772 m. The tests were carried out in ambient air (relative humidity 60%) under dry sliding condition. Sets of sample were tested also under in distilled water (wet immersion) and in 3.5% NaCl solution (corrosive immersion) at same sliding speed and load against hardened MS disc. At least three tests were done for each type of material. The weight loss difference before and after the test of each samples were taken and wear rates were calculated from average values of weight-loss measurements. Wear rate was estimated by measuring the weight loss ($\Delta W$) after each test. Care has been given after each test to avoid entrapment of wear debris. The wear rate was calculated using the following expression [21]:

$$W.R = \frac{\Delta W}{S.D \times L},$$

where: $W.R$ – wear rate, $\Delta W$ – weight loss, $S.D$ – sliding distance, $L$ – load.

Microstructural observation of the worn specimens were done carefully by using OPTIKA Microscope with a CCD camera (Model: OPTIKA) attached to PC imaging at $\times$100 magnifications and some selected photomicrographs were taken. The SEM investigation
and EDX analysis were conducted by using Ultra-high resolution JEOL scanning electron microscope with an energy dispersive X-ray analyzer (Model: Link AN – 10000) attached.

3. RESULTS AND DISCUSSION

3.1. Physical and Mechanical properties

Figure 1 and Figure 2 show the physical and mechanical properties of commercially pure Al, Al-Mg alloy and Al-Mg-Zr alloy respectively. The density of Al-Mg alloy and Al-Mg-Zr alloy are lower than that of commercially pure Al because of the presence of approximately 5wt% Mg of quite lower density. A small variation is observed in case of Al-Mg-Zr alloy for the presence of 0.3wt% Zr. It is also found that the hardness of pure aluminium increases when Mg is added to it due to the solid solution hardening. Magnesium has higher specific strength than Aluminium. Thus the addition of Mg results in increased hardness of alloy as compared to Aluminium [22]. Al-Mg-Zr also shows the highest hardness because of the grain refining effect by Zr [17].

![Fig. 1. Density and microhardness of the alloys](image1)

![Fig. 2. Tensile properties of the Al alloys](image2)

It is seen from the Figure 2 that Mg addition in Al increases the tensile strength while Zr addition in Al-Mg alloy further increases the strength i.e. Al-Mg-Zr shows the highest strength among the alloys studied. The Al-Mg alloy shows the higher UTS due to the content of higher amount of solute of \( \beta'' \) \( L_1^2 \) (Al\(_3\)Mg) and \( \beta' \) phase Al\(_3\)Mg\(_2\), which directly increases mechanical properties of Al-Mg alloys by creating obstacle of dislocation movement occurred under tensile load [23, 24]. In case of Al-Mg-Zr alloy, Al\(_3\)Zr precipitates are formed in the alloy during solidification. The presence of the high number Al\(_3\)Zr contents in the alloys may provides a number of sites for the heterogeneous nucleation of precipitate which yields finer grain structure and results the highest tensile properties of the alloy. It is found that the effect of Mg and Zr addition in Al on the tensile properties is similar to those of hardness. The ductility of Al is found to decrease with the increase of strength which is obtained due to the addition of Mg and Zr. Al\(_3\)Mg, Al\(_3\)Mg\(_2\) phases and Al\(_3\)Zr fine precipitates prohibiting the deformation decrease the ductility of Al-Mg and Al-Mg-Zr alloy [25, 26].

![Fig. 3. Variation of weight loss with the sliding distance in dry sliding condition](image3)

Moreover, Figures 4-6 elucidate the variation of wear rate with the variation of sliding distance for all the alloys in dry, wet and 3.5% NaCl corrosive environment respectively. The wear rate of the alloys in dry sliding condition is found to decrease with the increased hardness of the alloys and this result is found to be consistent with the Archard’s theory (Fig. 4) [29].
The contact surface of the samples against disc becomes more conformal to increasing the sliding distance under the abrasive action of hard rotating disc which leads to increase in temperature and the material becomes ductile as well. Al-Mg alloys contain supersaturated amount of magnesium and it forms of the \( \beta' \) and \( \beta \) phases which harden the alloy. It was found that the Zr addition strengthens the matrix in Al-Mg alloy and hence more wear resistance is observed in Al-Mg-Zr alloy as compared to Al-Mg alloy. Because, L12 (Al3Zr) phase formed during solidification in Al-Mg-Zr alloy, which are coherent with the matrix and thermally stable [30, 31]. Thus this phase provides high temperature hardness [32].

![Figure 4](image4.png)

**Fig. 4.** Variation of wear rate with the sliding distance in dry sliding condition

In case of wet and corrosive environment, the wear rate of Al is significantly lower than that of Mg and Zr added alloys in both conditions (Fig. 7 and Fig. 8). Generally, aluminum shows good corrosion resistance in marine environments because of the formation of a protective adhered aluminum oxide film on the material surface. However, in Al-Mg alloys containing supersaturated amount of magnesium are sensitized with the formation of the \( \beta' \) and \( \beta \) phases which are susceptible to corrosion. Al alloys with more than a 3.5 wt.% magnesium content become susceptible to the formation of a magnesium-rich \( \beta' \) intermetallic phase at grain boundaries. Therefore, in a corrosive environment, the \( \beta \) phase is preferentially attacked resulting in anodic dissolution of \( \beta \) phase from the grain boundaries, which is enhanced under wear action [33, 34]. It is also observed that the wear rate Al-Mg is lower in wet condition than that of Al-Mg-Zr alloy while the rate is found to be higher in Al-Mg alloy than the Al-Mg-Zr alloy in corrosive environment. Addition of zirconium improves remarkably stress-corrosion resistance due to formation of subgrains caused by zirconium-bearing particles [35].

The variations of coefficient of friction for commercially pure Al, Al-Mg alloy and Al-Mg-Zr alloy under dry, wet and corrosive condition at different sliding distance have been shown in Figures 7-9 respectively. It is observed that initially the coefficient of friction is low and then increases with increasing sliding distance. Initial low value of frictional coefficient is due to the contact between the oxide layers adhered on the specimen and disc material. Within a short sliding distance, the coefficient of friction increases. Because cracking and removal of surface oxide layer leading to the metal-to-metal contact causes an increase in the coefficient of friction. Interface temperature increases with the increment in sliding distance that may promote the surface oxidation and reduce the direct metal contact hence there is slight decline in frictional coefficient [36]. The coefficient of friction for all the alloys in dry environment is much greater than under wet and corrosive environment. The cause of this friction reduction is the “Sealing Effect”, which reduces the roughness of the surfaces in contact [37]. In all the cases the coefficient of friction of commercially pure aluminium shows significantly higher than that of Al-Mg and Al-Mg-Zr alloys and the result is in good agreement with the observed microhardness value of the alloys. Differences in the extent of localized plastic deformation at real contact areas may lead to the deficiencies in friction coefficient. The Mg added alloys have exhibited lower friction as they are harder and undergo less plastic deformation [38]. The effects of variation of applied load on friction behavior of the alloys in different sliding condition are shown in Figures 10-12. It is observed that with increase in applied load coefficient of friction increases. This is due to the fact that at higher applied load the amount of deformation or ploughing is more leading to higher coefficient of friction as well as higher wear. It means there is no monotonic trend of variation of wear resistance with applied load probably due to non-homogeneous nature of the material. At higher loads, the oxide debris is expected to get better compacted and mixed and form transfer layer and spread over a larger area of the sliding surface.

![Figure 5](image5.png)

**Fig. 5.** Variation of wear rate with the sliding distance in wet sliding condition
Fig. 6. Variation of wear rate with the sliding distance in corrosive sliding condition

Fig. 7. Variation of friction coefficient with the sliding distance in dry sliding condition

Fig. 8. Variation of friction coefficient with the sliding distance in wet sliding condition

Fig. 9. Variation of friction coefficient with the sliding distance in corrosive sliding condition

Fig. 10. Variation of friction coefficient with applied load in dry sliding condition

Fig. 11. Variation of friction coefficient with applied load in wet sliding condition
However, the wear rate under such circumstances takes place also by flaking of the transfer layer during sliding, apart from the processes of adhesion, micro-cutting and abrasion. At higher loads, the wearing process could be more aggravated by the transfer layer flaking off as indicated by the presence of a chunky sheet of oxide agglomerates in the wear debris [39]. However, under wet and corrosive condition, the value of coefficient of friction reduced for all the alloys compared with dry condition; this is because of thin lubricating film formation between specimen and rotating disk surface. This thin lubricating film contains lumps of particles and as temperature and normal loads increase these particles break into a small number of particles and reorient themselves along the sliding plane and this leads to lower coefficient of friction [40].

3.3. Optical microscopic observation

Figure 13 shows the worn surfaces for all the alloys at different sliding conditions. The optical micrograph of polished alloys have characterized by an Al-rich dendritic matrix, α-Al phase and a eutectic mixture in the interdendritic region. In this type of image the precipitates emerge in the dark tone while those of α-Al become visible in a lighter tone [41]. From the figure at dry sliding condition, it is observed that crater and small grooves formed at the worn surface due to thermal softening of the material occurred for raising interface temperature at applied loads causing removal of the thin metal layer from the surface which cause the propagation of micro-cracks. From the figure it can be also observed that, at dry condition, the wear marks in the worn surfaces of Al-Mg alloy is lofty because of brittle β-phase, Al₃Mg₂, to precipitate in grain boundaries. In case of Zr added alloy the wear mark relatively reduced because of the presence of Al₃Zr reinforced particles into the grain. At wet and corrosive environment, wear marks are not clearly visible in the worn surfaces of all the alloys due to the presence of corrosion product. From the figure it can be seen that, Al-Mg alloy showed higher wear corrosion than that of commercially pure Al [42]. No single wear mechanism can be attributed as the rate controlling mechanism throughout the mild wear regime. It was found that low loads and velocities produced sub-microscopic aluminium and iron particle debris which initially detach from the contact surfaces. This mechanically mixed phase predominantly consisted of aluminium oxide. The hardened aluminium oxide then facilitates the detachment of iron from the steel counter-face to produce the iron debris. This debris deposits on the surface and from time to time spall off, contributed to the wear. In case of Mg added alloy some delaminating layer is observe on the surface because of the stress corrosion cracking initiated by β” phase Al₃Mg₂. On the other hand the wear mark on the Zr added alloys are uniformly distributed and relative fine due to grain refining effect of the alloys.

3.4. SEM and EDX observation

The SEM micrographs of commercially pure Al, Al-Mg and Al-Mg-Zr alloy are shown in Figure 14. The commercially pure Al consists of a heterogeneous microstructure with dispersed grains. Because iron and silicon are ever-present impurity elements and the solid solubility of iron in aluminum is very small. Thus phases of aluminium-iron or aluminum-iron-silicon are seen in microstructures (Fig. 14.a). In the as-cast condition, all of the phases that come into equilibrium with aluminum may be found-FeAl₃, Fe₃SiAl₅, or Fe₃Si₈Al₂. In addition, a number of metastable nonequilibrium phases may be formed when solidification is rapid [43, 44]. The corresponding EDX profile analyses for the commercially pure Al shows the weight percentage of elements as 97.32% Al, 1.73% Si, 0.92% Fe and 0.03% Sn. The microstructure of Al-Mg alloy shows dendrites with second phase particles within interdendritic spaces (Fig. 14.b) [45]. The silicon particles are also refined due to formation of Mg₃Si [46]. Addition of Zr to Al-Mg alloy shows a diminution in the amount of second phase particles. It further appears that dendrite arm spacing is decreased in Al-Mg-Zr alloy with the consequent refinement of dendrites (Fig. 14.c). The dendrites of the cast base alloy are seen to have refined significantly with the addition of Zr. Reportedly alloy with 0.3% Zr does not provide much grain refinement but refines the primary dendrites of α with consequent diminution of dendrite arm spacing [47, 48]. The corresponding EDX profile analyses for the Al-Mg alloy the weight percentage of elements found 4.900% Mg, 94.799% Al, 0.19% Si, 0.08% Fe and 0.04% Sn. Similarly Al-Mg-Zr alloy shows 6.53% Mg, 89.88% Al, 2.79% Si, 0.13% Fe, 0.57% Zr and 0.09% Sn.
Fig. 13. Optical micrograph of worn surfaces before wear, after wear in dry, wet and corrosive sliding condition

The worn surfaces of the experimental alloys after wear in dry sliding condition are shown in Figure 15 with the help of scanning electron microscope. It is observed that some cracks generated from surface fatigue caused by Hertzian contact stress (Fig. 15 a). On the other hand, some delamination layer is also occurred on the surface of Mg added alloy (Fig. 15. b). The surface of the sample is not only brittle but it is also suitable to plastic yielding. The fracture surface of Zr added alloy shows the additional brittle surface (Fig. 15 c). Addition of Mg increases the amount of reinforcing phase Al3Mg2 which is brittle [49]. When Zr is added into the alloy, a typical plastic deformation and a large amount of pits are formed into the Al-Mg alloys due to presence of fine precipitates of Al3Zr as well as grain refining effect [50, 51].
Fig. 14. SEM images and EDX of: a) commercially pure Al, b) Al-Mg alloy, c) Al-Mg-Zr alloy
Fig. 15. SEM images of: a) commercially pure Al, b) Al-Mg alloy, c) Al-Mg-Zr alloy after wear in dry sliding condition at 1.02MPa applied pressure and sliding distance of 2772 m

4. CONCLUSIONS

In the present work, it was found that the addition of magnesium and zirconium improve the strength of commercially pure aluminium due to the solid solution strengthening and grain refinement effect respectively. It also increases the wear behaviour of the alloys in dry sliding condition. In case of wet and corrosive environment the specimens shows higher amount of material loss i.e. higher wear rate. During this process, magnesium segregates toward the grain boundaries and forms the secondary precipitate β phase (AlMg2). When exposed to harsh environments such as salt water, a galvanic couple is formed between the Al matrix and the β phase precipitates. The precipitates become anodic to the matrix and preferentially dissolve leaving gaps along the boundary network, ultimately leading to stress corrosion cracking. The coefficient of friction for all the alloys under wet and corrosive environment is lower than dry sliding condition owing to the sealing effect. The wear marks in the worn surfaces of Al-Mg alloy is lofty because of brittle β-phase precipitate in grain boundaries and the fine fracture grain in case of Zr added alloy are shown due to grain refining effects.

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