SLURRY EROSI VE WEAR BEHAVIOR OF AA 6063/TIC PARTICLES IN SITU COMPOSITES

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ABSTRACT

The In-situ composite of metal matrix AA6063-xTiC (x = 0 wt. %, 3 wt. %, 6 wt. %, 9 wt. %) were mixed with particulates of TiC, were synthesized by the reaction of K2TiF6 and graphite with aluminum melt at 1000°C. These composites were heat treated differently to form various sets like solution zed and solution zed-aged, where they were compared with as-cast composites for erosive wear performance by testing pot type slurry-erosive wear device, respectively. This study showed an increase in wear resistance for slurry erosion upon solution zed-aged heat treatment and with a rise of TiC particulates.

KEYWORDS: Metal Matrix, Wear and Slurry Erosion

INTRODUCTION

The class of innovative engineering materials of metal matrix composites (MMCs) is with a vast variety of wear, structural and thermal managerial applications. The metal-matrix composites can provide high-temperature working limits than their base metal corresponding element, and it can be tailored to give stiffness, thermal conductivity, enhanced strength, abrasion resistance, dimensional stability or creep resistance [1, 2]. MMCs has been focused by record of the commercial work on aluminum as the metal matrix such as ventral fins, cylinder liners, helicopter blades and lower drag brace landing gears in modern fighter planes[3]. The light weight and useful mechanical properties combination has ended aluminum alloys very popular; aluminum also makes these properties well-matched for use as a metal matrix. Though, aluminum itself shows deprived tribological properties [4]. Therefore, the importance in the study is the tribological behavior of aluminum based materials is becoming increasing. By continuous exposure to environment, these materials get degraded. Due to the continuous exposure to such environment, the body erosion in automotive applications wear of these materials plays a major role [5].

Aluminum melting point is more than enough to gratify many tender requirements, still little enough to reduce composite processing reasonably suitable. B4C, TiB2, Al2O3, Sic, Tic, WC, and graphite variety of reinforcing...
agent which accommodates by the aluminum are regularly used as reinforcements. The matrix increases the stiffness and strength by incorporation of the reinforcement [6, 7], increases the hardness of the composite with addition of hard ceramic particles, which improves the resistance of the matrix to reduce and penetrate subsequent removal of material by other third-body particles and wear debris found in the environment [8]. In particulate-reinforced metal matrix composites, Titanium carbide (Tic) was commonly used as reinforcement due to its relatively high thermal, high hardness, thermal stability, wear resistance, electrical conductivities and good wet ability.

To fabricate composite materials, various synthesizing techniques were used InTic reinforcement, such as in-situ melt reaction route and powder metallurgy route. Though, it has been observed that problem of contamination at matrix-reinforcement interfaces the powder metallurgy route [9-11], to synthesize Tic reinforced in aluminum matrix, the in-situ approach is much economical and suitable. Through in-situ approach method of chemical reaction, many of the researchers have synthesized Tic particulates to form a very fine thermodynamically stable ceramic phase using Al-Ti-C system [12-16]. In aluminum alloy the Tic particulates synthesis the chemical reaction of graphite powders and K$_2$TiF$_6$. The reaction temperature that played a critical role in synthesis of Tic from Al- K$_2$TiF$_6$-C system has been observed. SHS process (Self-propagating High-temperature Synthesis) has been involved by synthesis of Tic particulates in aluminum matrix. In SHS process various parameters such as stoichiometry of the reactants, pellet size, particle size and green density have been studied [17].

By different routes, many researchers investigated the erosion wear behavior of Al based composites [18-20]. Erosive wear resistance of the composites rises with the quantity of reinforcement that has been stated. For solid particles entrained in a fluid impinge on surface that occurs, erosion wear is major degradation mechanism for structural materials, which occurs [21-22]. In the present investigation, no reports were formed in literature on this aspirant in-site composite material and hence an attempt towards slurry erosion wears. Further, study has been done on erosion wear influence of the heat treatment on Al-Si-Mg alloys. Thus, the increase in solution zing temperature and aging duration with maximum hardness for 6 hours aging duration it has been proved increasing of the hardness. Thus, Al-Si-Mg alloys exhibited improved wear resistance upon heat treatment [23].

In the present work, AA 6063metal matrix composite has been synthesized through in-situ casting route with reinforcement up to 9 wt. % of Tic. The synthesis of Ticwas performed through impulsive reaction between graphite powders and K$_2$TiF$_6$ in aluminum alloy melts. The slurry wear of the composites for different conditions have been studied, with the pot type slurry erosion tester.

**EXPERIMENTAL DETAILS**

**Casting and Heat treatment**

To develop the Tic formation and its even distribution in AA 6063, matrix alkaline salt (K$_2$TiF$_6$) and graphite powder were used. In Table 1, the insignificant chemical composition of AA 6063 alloy has been provided. By changing the mass of the above-mentioned chemicals added in the aluminum alloy melt in the fraction of Tic particulates has been varied. The chemical mass to be added in the melt, it has been calculated as per the stochiometric ratio to procedure composites C$_0$(AA 6063-0 wt.% Tic), C$_1$(AA 6063-3 wt.% Tic), C$_2$(AA 6063-6 wt.% Tic) and C$_3$(AA 6063-9 wt.% Tic). As shown in Table 2, the weights of chemicals added in different composites.
The chemicals $K_2TiF_6$ and graphite powder were prudently weighed as calculated earlier, and mixed thoroughly to prepare the above-mentioned composites. Then, the AA6063 carefully weighed and added in a silica crucible, and it was kept inside the furnace. Thus, in the AA6063 alloy the prepared mixture of the chemicals was added and melt was preserved at the temperature of 1000°C for 30 minutes inside the furnace for in-situ growth of TiC particulates. To synthesize the TiC particulates from $K_2TiF_6$ and graphite powders important one is to maintain temperature ranges between 900°C to 1100 °C. A vital exothermic reaction started soon after the accumulation of chemicals into the AA 6063 alloy melt. Then after the 10 minutes of time interval the mixture was stirred in the crucible. Finally, preheated to the temperature at 300 °C the molten mixture was carefully poured in a cast iron mould.

The cast blocks were laid open to two dissimilar heat treatments after casting the composites, viz., AA 6063 – tic (x = 0, 3, 6 and 9 wt.%). Thus, based on their condition the sets of composites were prepared. In as-cast condition, the first set of composites was remained and not heat treated. For one hour, the second set of composites was solution zed at 530 °C and then quenched in water. The characterizations of the prepared composites have been done using optical microscope. Using the Hitachi S-3000 H scanning electron microscope, the images of TiC particulates in the composites have been taken. Also, for the composites of $C_0$, $C_1$, $C_2$ and $C_3$ the micro hardness of different conditions has been measured by using Wick 3212 Vickers hardnes tester. Using the D/MAX ULTIMA III XRD machine supplied by Rigaku Corporation, Japan, with Cu-Ka radiation (1.54056 a°) the X-ray diffraction analyses (XRD) for the composites with varying Tic content in as-cast condition were conducted.

Slurry-Erosion Test

Erosion wear tests were carried out using a pot type slurry-erosive wear apparatus supplied by Friction and Wear Tech., Chennai, India. An abrasive particle slurry comprising silica sand (white, with a mesh size 30/80) with 33% sand concentration and remaining water was used. The composite specimens were rotated in the slurry with a speed of 500 rpm for 8 hours’ test duration. The mass loss was measured after each hour using an electronic weighing balance with an accuracy of 0.001gms, and then, the specimens were kept again in same position for erosive wear tests. The amount of wear was quantified by measuring the volume loss. The wear rate was given as: volume loss/distance traversed (mm³/km). The surface morphologies of the worn-out surface of the composites were studied by scanning electron microscope (SEM).
RESULTS AND DISCUSSION

Characterization of the Composites

Micro Hardness Analysis

Table 3: Hardness Values of the in Situ Composite as Different Conditions

<table>
<thead>
<tr>
<th>Weight Percentage of Tic</th>
<th>As Cast Condition $H_v_{0.5}$</th>
<th>Solution Zed Condition $H_v_{0.5}$</th>
<th>Solution Zed-Aged Condition $H_v_{0.5}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>56</td>
<td>72</td>
<td>76</td>
</tr>
<tr>
<td>3</td>
<td>63</td>
<td>88</td>
<td>89</td>
</tr>
<tr>
<td>6</td>
<td>80</td>
<td>103</td>
<td>108</td>
</tr>
<tr>
<td>9</td>
<td>92</td>
<td>118</td>
<td>123</td>
</tr>
</tbody>
</table>

The results obtained from micro hardness testing using Vickers hardness tester (Wick 3212) with an indentation load of 0.5Kg and dwell time of 15 sec. are shown in Table 3. The results reveal that by increasing the weight percent of Tic, the hardness values increase due to an increase in volume fraction of hard phase. The in-situ composites with solution zed-aged condition exhibited an improved hardness compared with the composites either in solution zed or in as-cast condition. The improved hardness values of the composites in solution zed-aged condition can be attributed to the formation of fine $\text{Mg}_2\text{Si}$ precipitates from AA 6063 matrix. After comparing with the composites in as-cast condition, the composite in solution zed condition exhibited a better hardness value. The reason may be associated with homogenization and uniform distribution of Tic particles after the solution zing heat treatment.

Scanning Electron Microscopy

SEM images taken from a Hitachi S 3000-H microscope is shown in Figure 1. In the high magnification images, the size of Tic particles is found to be 2-5µm. Also, needle shaped $\text{Al}_3\text{Ti}$ particles of 8-10µm width and 150 -170 µm are observed.

![Figure 1: SEM Micrographs of AA 6063/Tic Composite with as-Cast Condition](image-url)

A) C1, B) C2, C) C3 and D) C4 Showing Al3Ti and Tic particles
X-Ray Diffraction Analysis

X-ray Diffraction (XRD) patterns of the matrix specimen and of the specimens reinforced with different percentages by weight of Tic are shown in Fig. 2. It shows the X-ray diffraction (XRD) patterns obtained from the cross-section of cast composites viz., C₀, C₁, C₂ and C₃. All the peaks are identified as arising from three phases namely, Al-based phase, Tic and Al₃Ti from JCPDS files (file no. 892837, 893828 and 650428) respectively.

Also, through a relative comparison of the XRD patterns of composites C₁, C₂ and C₃, it can be seen that with an increase of weight fraction of Tic particulates the corresponding peaks of Tic and Al₃Ti show an increase in relative intensity. From the XRD peak profile analysis, the area under the peak was measured and the relative fractions of Tic were determined in the case of composites C₁, C₂ and C₃. The volume fractions of the corresponding composites were calculated as 2.85%, 4.20% and 7.25%. These reduced fractions are attributed to the melting losses, as against the targeted values.

![XRD Results For the AA 6063/Tic Composites](image)

**Figure.2: XRD Results For the AA 6063/Tic Composites**

Slurry Erosive Wear

The slurry erosive wear rate of the composites was computed from mass loss measurements. The variation of cumulative mass loss of the composites C₀, C₁, C₂ and C₃ for different heat treated conditions with interval of an hour is shown in Figure 3.
A linear plot was observed for unreinforced alloy C_0 for all conditions. It indicates a uniform wear with time, whereas, in the case of composites C_1, C_2 and C_3 nonlinear plots are observed. It may be attributed to the inhomogeneity in the presence of Tic particles in the composites C_1, C_2 and C_3.

With increase in weight fraction of Tic particles, the mass loss of the composites decreased. This can be explained by the more resistance offered due to Tic particles causing less indentation and scratches in aluminum matrix. The composite C_1 showed a linear relationship of mass loss against time up to 4-5 hours of test duration. Thereafter, marginal decrease in mass loss is observed by the slope of the curve for composite C_1. The composite C_3 offered maximum resistance to erosion wear among all composites. The total mass loss observed for composites C_0, C_1, C_2 and C_3 with as-cast condition after 8 hours of test duration was 1.026, 0.850, 0.656 and 0.528 grams, respectively. The total mass loss observed for composites C_0, C_1, C_2 and C_3 with solution zed condition after 8 hours of test duration was 1.001, 0.817, 0.599 and 0.437 grams respectively. The total mass loss observed for composites C_0, C_1, C_2 and C_3 with solution zed-aged condition after 8 hours of test duration was 0.947, 0.713, 0.449 and 0.338 grams respectively. It is clear from the results that the composites with solution zed-aged condition showed lesser mass loss than composites with solution zed and as-cast conditions. This may be attributed to increase in hardness of the matrix due to precipitation hardening. Fig. 4 shows the variation of erosive wear rate after 8 hours of test duration for all composites with different conditions. Wear rate is found to decrease with increase in weight fraction of Tic particles. The decrease in slope from composite C_2 indicates that the change in wear from composite C_1 to C_2 is more than the change in wear from composite C_2 to C_3.
The erosive surfaces were studied using SEM with the aim of understanding the mechanism of material removal. Figure 5 shows the SEM micrographs of worn surfaces of composites with solutionized-aged condition. The SEM micrograph (Fig.12a) of unreinforced alloy shows deep and wide plowing and cutting grooves. The aluminum material is deformed plastically on both sides of the plowing grooves without any restriction. Wedge formation in the unreinforced alloy is observed by chisel action of sharp sand particles present in the slurry. With increase in weight fraction of Tic particles in the composites the plowing grooves seems to decrease in size and number. The grooves seem to be restricted by the resistance offered to indentation by Tic particles (Fig. 5d). The presence of silica
sand particles in water was responsible for erosion-abrasion wear of surface of the composites. The erosion is mainly caused by plowing, plastic deformation, wedge formation and voids. The plowing action and plastic deformation are predominant in unreinforced alloy and lesser reinforced composite $C_1$. Whereas, voids caused due to removal of hard faced Tic particles are observed due to the impact of sharp sand particles in composite $C_2$ and $C_3$.

CONCLUSIONS

The present investigation has led to the following conclusions: It is possible to produce in situ Tic particles in an AA 6063 alloy - $K_2TiF_6/C$ system through a simple and cost effective in melt reaction synthesis. The SEM and micro hardness testing results indicated the presence of Tic particles which have been proved by XRD analysis and hence, the feasibility of the composite. The solution zing -aging of the composite shows good improvement in the hardness of the composite as compared to solution zing heat treatment alone. Erosive wear rate decreased with the presence and increase on Tic particles in the composites. In erosive wear tests, composites with solution zed-aged condition emerged to have better wear resistance than composites with solution and as-cast conditions. In the case of erosive wear, the predominant mechanism of wear was plowing and wedge formation for unreinforced alloy, whereas, voids creation in composites $C_2$ and $C_3$ is the major cause of material loss.

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