Exploiting superior tensile properties of a novel network-structure AlA206 matrix composite by hybridizing micron-sized Al3Ti with Al2O3 nano particulates

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Exploiting superior tensile properties of a novel network-structure AlA206 matrix composite by hybridizing micron-sized Al₃Ti with Al₂O₃ nano particulates

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Abstract

In this study, semi-solid stir casting and ball milling processes are combined into an integrated composite fabrication process. Two different architectures were utilized to incorporate reinforcing particle into semi-solid alloy i.e., (i) ball milling of K₂TiF₆ and aluminium powder for 5 h and subsequently with nano-alumina particles (Al₂O₃np) for 1 h and (ii) ball milling of K₂TiF₆, Al₂O₃np and aluminum powder for 2 h. Accordingly, the milled powders were incorporated into molten AlA206 alloy using a non-contact ultrasonic vibration method. The effect of milling procedure on microstructural evolution and tensile properties were then explored. Two different microstructures were characterized including well-distributed Al₂O₃np and Al₃Ti particles (Al₃Tip) and a network-structure containing Al₃Tip+Al₂O₃np. This unique architecture of network-structure brought about increment in tensile properties compared to well-distributed reinforcement particles, ascribed to the strewing of Al₃Tip+Al₂O₃np around matrix grain boundaries, act as a three-dimension skeletal structure with high local volume fraction of Al₃Tip+Al₂O₃np.

Keywords: A206 Alloy; Semi solid processing; Metal matrix composites; Transmission Electron Microscopy; Fracture.

1. Introduction

One of the most important metal matrix composites (MMCs) are aluminum matrix composites (AMCs) reinforced alumina particulates, extensively being used in the aerospace and automobile industries [1-7]. AMCs, especially discontinuously reinforced aluminum matrix composites (DRAMCs) have received large attention because of their augmented tensile and tribological properties [1-8].

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DRAMCs with uniformly distributed reinforcement exhibit a certain improvement in partial properties with respect to the matrix alloy [9]. Investigations in the past two decades confirmed that DRAMCs exhibited limited strengthening effect of reinforcement and augmented mechanical properties can be achieved by tailoring the microstructure at a higher level [10-12]. In order to increment the performance of DRAMCs, recent studies have been focused towards exploring three-dimensional microstructures such as bi-continuous, inter-penetrating and quasi-continuous composites [13-15]. As reported by Huang et al. in the case of Ti matrix composites, a controlled three-dimensional microstructure has the capability to enhance the properties of material, such as elastic modulus, tensile strength and fracture toughness, compared with well-distributed counterparts, [13, 15-19].

Beside ceramic reinforcements, trialuminide intermetallics such as Al3Zr and Al3Ti have been also used widely as reinforcement [20-24]. Trialuminide intermetallics have some important advantages over ceramic particulates such as low densities, higher thermal stability, good machinibility and formability [25,26]. However, the brittle nature of trialuminide intermetallics can preside over the properties of the composite giving rise to negligible increment in mechanical properties [22]. Therefore, microstructural control is needed to implement Al3Ti intermetallic as a useful reinforcement in MMCs. Furthermore, using two or more types of particulates in a single matrix, hybrid composites, the benefits of one type of particulates could supplement to what is lacking in the other [27].

This study shows how microstructural control can be used to exploit superior tensile properties of Al/A206-5%Al2O3-Al3Ti hybrid composites. In order to achieve this, intelligent techniques with superior process control such as high energy ball milling coupled with stir casting was employed. The effect of using this process for incorporation of reinforcing particles on final nano/microstructure and tensile properties of the composite, produced using a process involve milling and semi-solid stir casting, was explored.

2. Experimental procedure

Commercial aluminium (74 µm, supplied by Phoniex company), α-alumina (100 nm, supplied by Phoniex company) and K2TiF6 (60 µm, supplied by Aldrich company) powders were used in this study. Milling was performed in a Fritsch Pulverisette P5 planetary ball mill under argon gas (high purity, 99.999%) in a liquid
nitrogen environment (cryomilling). Rotation speed of the mill was kept about 480 rpm. In order to avoid excessive cold welding during milling, process control agent (PCA) was used (Stearic acid (CH₃(CH₂)₁₆CO₂H) powder was supplied by Merck).

To explore the behavior of K₂TiF₆ and aluminium powders against thermal exposure, differential scanning calorimetry (DSC) (at a rate of 10 °C/min and under flowing argon) was utilized. Four powder samples were used; Aluminum powder, K₂TiF₆ powder, mixed mixture of aluminium and K₂TiF₆ powders and milled mixture of aluminium and K₂TiF₆ powders. Volumes of mixed and milled aluminium/K₂TiF₆ powder blends were heated in a tube furnace at a rate of 10 °C/min (similar to DSC experiment) to temperatures corresponding to each major DSC signals for 1/2 h and subsequently quenched to room temperature to avoid microstructural changes during cooling.

X-ray diffraction (XRD) analysis was performed on heat treated powders to determine possible reaction corresponding to each DSC signal. XRD analysis was performed using PHILIPS-binary diffractometer applying Cu Ka radiation.

A206/5 % alumina-Al₃Ti hybrid composites were fabricated according to the following procedure:

Initially, 99.7% pure aluminium, Cu and aluminium-80%Mn ingots were charged to the furnace and melted. After entire alloy in the crucible was melted, it was cooled down to 640°C. This temperature lies in the solid-liquid range and corresponds to solid fraction of about 0.2. Then, stirring of the semi-solid alloy was initiated in 400 rpm, while prepared powders were injected in to the uniformly formed vortex over a time period of approximately 5 min. Simultaneously, the non-contact ultrasonic casting method was utilized in order to apply vibration into the prepared melt. It consists of an ultrasonic chamber (Bandelin-Germany Make – Model: RK – 100H), which can vibrate at a frequency of 35 kHz. Powder injection into molten A206 aluminium alloy was performed under high purity argon atmosphere (99.999%, 6 lit/min). Two different architectures were utilized to incorporate reinforcing particle into semi solid alloy, i.e. (i) ball milling of K₂TiF₆ and aluminium powder for 5 h and subsequently with alumina for 1 h (denoted herein as (K₂TiF₆+Al)₅h+Al₂O₃(1h)) and (ii) ball milling of K₂TiF₆, aluminum and alumina powder for 2 h (denoted herein as (K₂TiF₆+Al +Al₂O₃)₂h). Indexes in parenthesis, hereafter, referred to milling time.
A proper mixture of the mentioned composition was selected so that after introducing powders into the melt, melt composition reached A206 alloy (Cu=4.20-5.00, Mg=0.20-0.35, Mn=0.20-0.50, Fe=0.07 max, Ni=0.03 max and Al=balance). After completion of particle injection, mixing was continued for extra 2 min. Finally, the composite slurry was poured into a pre-heated mould by using a bottom-pouring system. The composites fabricated via technique (i) denoted herein as A206-(K₂TiF₆+Al)₅h+Al₂O₃₃h and that of prepared via technique (ii) as A206-(K₂TiF₆+Al +Al₂O₃₂h).

Morphology evolution of powders during milling and microstructure of the composite were studied by Field Emission Scanning Electron Microscopy (FE-SEM) performed in a HITACHI S4160, equipped with an energy dispersive X-ray analysis (EDX) accessory and Transmission Electron Microscopy ((TEM) Philips CM200).

Tensile testing was carried by a Hounsfield universal test machine at a cross-head speed of 0.5 mms⁻¹. The dog-bone shaped tensile specimens had a gauge size of 6 mm in diameter and 30 mm in length, according to ASTM: B557M-10.

3. Results and discussion

3.1. Thermal response of powders

Fig. 1a shows DSC pattern of K₂TiF₆ salt, evincing three endothermic signals. Signals 1, 2 and 3, as demonstrated in Fig. 1a, was appeared around 360 °C, 610 °C and 840 °C, respectively. Figs. 2a and b exhibit XRD patterns of K₂TiF₆ salt held isothermally above signal 1 and 2 for 30 min at 450 °C and 650 °C, respectively, revealing K₂TiF₆ reflections alone. It is thus authenticated that K₂TiF₆ salt is stable up to 650 °C.

The endothermic signals 1 and 2 are ascribed to the melting of impurities in K₂TiF₆ salt and that of signal 3 to the melting of K₂TiF₆ salt.

Fig. 1b shows DSC pattern of aluminum powder, evincing a single endothermic peak around 660 °C (signal 4) attributed to the melting of aluminum.

Fig. 1c shows DSC pattern of mixed mixture of aluminum and K₂TiF₆ salt. As is observed, by considering signal 5, DSC pattern of the mixture is not a simple superimposition of aluminum and K₂TiF₆. Signal 6 is ascribed to the melting of aluminum and K₂TiF₆ salt (between 630 °C and 650 °C).
Fig. 1. DSC patterns of (a) K$_2$TiF$_6$ salt, (b) aluminum powder, (c) mixed mixture of aluminum and K$_2$TiF$_6$ salt and (d) milled mixture of aluminum and K$_2$TiF$_6$ salt.

Fig. 2c shows XRD pattern of mixed mixture of aluminum and K$_2$TiF$_6$ salt held isothermally below signal 5, evincing aluminum and K$_2$TiF$_6$ reflections. It can be concluded that aluminum and K$_2$TiF$_6$ has not reacted with each other up to this temperature.

Fig. 2d demonstrates XRD pattern of mixed mixture of aluminum and K$_2$TiF$_6$ salt held isothermally at 600 °C. It can be deduced that signal 5 is responsible for reaction 1:

$$3 \text{K}_2\text{TiF}_6 + 13\text{Al} \rightarrow 3\text{Al}_3\text{Ti} + 3\text{KAlF}_4 + \text{K}_3\text{AlF}_6$$

(1)

FE-SEM image was also validated the formation of Al$_3$Ti in this temperature range (Fig. 3a). As is observed, Al$_3$Ti particles fused together in aluminum matrix.

Signal 6 in Fig. 1c is attributed to the melting of fluoride salts produced by reaction (1).

Fig. 1d shows DSC pattern of milled mixture of aluminium and K$_2$TiF$_6$ powder. The endothermic signal 7 starts at 220 °C and broaden to 450 °C. Signal 8 has also the same behavior as signal 6.

Fig. 2e shows XRD pattern of milled mixture of aluminium and K$_2$TiF$_6$ powder held isothermally above signal 7 (550 °C), evincing Al$_3$Ti, KAlF$_4$ and K$_3$AlF$_6$ in addition to those of aluminium. It is thus inferred that XRD pattern of milled mixture of aluminium and K$_2$TiF$_6$ powder held isothermally at 550 °C is the same as
that of mixed mixture of aluminium and K$_2$TiF$_6$ powder held isothermally at 600 °C. XRD spectrum of milled aluminium/K$_2$TiF$_6$ powders heat treated at 550 °C, after the completion of signal 7, is identical to that of the aluminium/ K$_2$TiF$_6$ mixture heat treated at 600 °C, with Al$_3$Ti, KAlF$_4$ and K$_3$AlF$_6$ reflections in addition to those of aluminium (Fig. 2c). Thus, it can be deduced that signal 7 is responsible for reaction (1). As such, signal 8 to the melting of KAlF$_4$ and K$_3$AlF$_6$, produced by reaction (1). The formation of Al$_3$Ti is also validated in the milled mixture of aluminium/K$_2$TiF$_6$ powders heat treated at 350 °C (Fig. 3b). Comparing Figs. 3a with 3b reveal the difference between morphology of Al$_3$Ti particles. Al$_3$Ti particles in the former are relatively coarser than in the latter.

This change in the morphology of Al$_3$Ti particles and the shift of reaction (1) to 220 °C-450 °C in the milled mixture, authenticates that the milling process facilitates K$_2$TiF$_6$/aluminium interaction.

Fig. 2. XRD patterns of (a) K$_2$TiF$_6$ salt held isothermally above signal 1 in Fig. 1, (b) K$_2$TiF$_6$ salt held isothermally above signal 2 in Fig.1, (c) mixed mixture of aluminium/K$_2$TiF$_6$ heat treated below signal 5 in Fig.1, (d) mixed mixture of aluminium/K$_2$TiF$_6$ heat treated above signal 5 in Fig.1 and (e) milled mixture of aluminium/K$_2$TiF$_6$ powders heat treated above signal 7 in Fig. 1.
It is therefore plausible to use this unique process for synthesizing in situ/ex situ MMCs using melt processing routes. By considering the fact that milling alleviates the formation temperature of Al$_3$Ti particles, inferior casting temperature can be utilized. Furthermore, as mentioned above, interaction between K$_2$TiF$_6$ and aluminium facilitates. In addition, this unique process has another surprising capacity in changing the morphology and distribution of Al$_3$Ti particles.

### 3.2. Morphology of milled powders

Fig. 4 shows the morphology of K$_2$TiF$_6$ (Fig. 4a), (K$_2$TiF$_6$+Al +Al$_2$O$_3$)$_{2h}$ (Figs. 4b and c), (K$_2$TiF$_6$+Al)$_{2h}$ (Fig. 4d) and (K$_2$TiF$_6$+Al)$_{3h}$+Al$_2$O$_3$ (1h) (Fig. 4e), respectively. Figs. 5a and b show XRD pattern of milled powders corresponding to Figs. 4b and e, respectively. Results of XRD analysis relieved KAlF$_4$, K$_3$AlF$_6$ and Al$_3$Ti in the case of (K$_2$TiF$_6$+Al)$_{3h}$+Al$_2$O$_3$ (1h) and K$_2$TiF$_6$, aluminium and Al$_2$O$_3$ in the case of (K$_2$TiF$_6$+Al +Al$_2$O$_3$)$_{2h}$. SAD patterns in Figs. 4b and e also confirmed the above findings.
Fig. 4. Morphology of (a) K$_2$TiF$_6$, (b) (K$_2$TiF$_6$+Al+Al$_2$O$_3$)$_{2h}$, (c) higher magnification of b, (d) (K$_2$TiF$_6$+Al)$_{2h}$, (e) (K$_2$TiF$_6$+Al)$_{2h}$+Al$_2$O$_3$
In the case of \((\text{K}_2\text{TiF}_6 + \text{Al} + \text{Al}_2\text{O}_3)_{2\text{h}}\) powder, nano alumina particles adhered to the sharp corners of \(\text{K}_2\text{TiF}_6\) powder (white arrows on Fig. 4c) which inhibiting the intimate contact between aluminium and \(\text{K}_2\text{TiF}_6\) powders and therefore impeding \(\text{Al}_3\text{Ti}\) formation. Thus, in order to prepare a suitable condition for the formation of \(\text{Al}_3\text{Ti}\), at first, \(\text{K}_2\text{TiF}_6\) and aluminium powders were milled and then alumina powder were added to the ball mill vial. As is observed in Fig. 4d, \(\text{K}_2\text{TiF}_6\) and aluminium powders are in close contact with each other after 2 h milling time. After 5 h milling and addition of aluminium powder and milling for 1 h (Fig. 4e), the mixture contains \(\text{Al}_3\text{Ti}\), \(\text{KAIF}_6\), \(\text{K}_3\text{AlF}_6\), and alumina (Fig. 5b).

On the other hand, in the case of \((\text{K}_2\text{TiF}_6 + \text{Al} + \text{Al}_2\text{O}_3)_{2\text{h}}\), as is observed in Fig. 4b, milled powders are integrated into capsule-shaped particles. During semi solid stir casting, capsule-shaped particles are disintegrated into molten alloy. The size of capsule-shaped particles after milling is lower than 20 µm that can accommodate better dissolution and lower agglomeration of them during subsequent semi-solid stir casting. Furthermore, inset in Fig. 4b, evincing cross section of capsule-shaped particles, substantiating uniform distribution of alumina particles inside them. In the other words, alumina particle were distributed not only on the outer surface of capsule-shaped particles but well-distributed underneath the surface.

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**Fig. 5.** XRD patterns of (a) \((\text{K}_2\text{TiF}_6 + \text{Al} + \text{Al}_2\text{O}_3)_{2\text{h}}\), (b) \((\text{K}_2\text{TiF}_6 + \text{Al})_{5\text{h}} + \text{Al}_2\text{O}_3_{1\text{h}}\).
3.3. Composite microstructure

Figs. 6a and b show FE-SEM microstructures of A206-(K₂TiF₆+Al)₅h+Al₂O₃₁h and A206-(K₂TiF₆+Al +Al₂O₃)₂h composites, respectively. As is observed, the size of Al₃Ti as well as the distribution of Al₃Ti and alumina particulates is not the same. Figs. 7a, b and c show HRTEM images of A206-(K₂TiF₆+Al)₅h+Al₂O₃₁h, high magnification of (a) and schematic illustration of (a), respectively. Figs. 7d, e and f show HRTEM images of A206-(K₂TiF₆+Al+Al₂O₃)₂h composites, high magnification of (d) and schematic illustration of (d), respectively. As can be seen in Figs. 7a and c, the majority of alumina and Al₃Ti particles in A206-(K₂TiF₆+Al)₅h+Al₂O₃₁h sample have a high propensity to reside in grain boundaries instead of grain interior. However, according to Figs. 7d and f, in the case of A206-(K₂TiF₆+Al+Al₂O₃)₂h sample, the majority of well-dispersed alumina and Al₃Ti particles are resided in the grain interior rather than grain boundaries, ascribed to following effects.

By introducing (K₂TiF₆+Al+Al₂O₃)₂h powders into molten alloy, three interesting phenomenon will occur: Firstly, aluminum powder in capsule-shaped particles melts and releases K₂TiF₆ and alumina particles into the melt. K₂TiF₆ reacts with metal melt by exothermic reaction 1, leading to local increment in temperature. This local increase in temperature fortifies the wettability of nano alumina particle with molten alloy as well as the willingness to particle engulfment [28].
Secondly, as result of reaction between K_2TiF_6 and molten alloy, fluorides components (according to reaction 1) release to the melt leading to removing oxide layer on the melt surface [29], which enhances the wettability of nano alumina particle with molten alloy [30-32].
Secondly, Al₃Ti has tetragonal structure with lattice parameters of a=0.3848 nm and c=0.8596 nm and α-Al has lattice parameter of a=0.4094 nm. Strain at interface in a and c direction is calculated through equation 2 and 3 [33]:

\[ \varepsilon_{aa} = (1 - a_{Al3Ti}/a_{Al}) \times 100 = +4.96\% \]

\[ \varepsilon_{cc} = (1 - c_{Al3Ti}/2a_{Al}) \times 100 = -6.15\% \]

These calculations authenticate that strain at interface is negligible and α-Al/Al₃Ti interface is coherent. In the other word, Al₃Ti is an appropriate site for secondary α-Al nucleation. Al₃Ti has (001) plane with the lowest strain mismatch with α-Al (012) plane. Therefore, it can be concluded that Al₃Ti is a suitable site for α-Al nucleation. Consequently, in the case of A206-(K₂TiF₆+Al +Al₂O₃)(5h), nucleation of Al₃Ti in molten alloy occurs simultaneously throughout the molten alloy results in the uniform distribution of Al₃Ti in the matrix.

On the other hand, in the case of A206-(K₂TiF₆+Al)(2h)+Al₂O₃(1h) sample, when (K₂TiF₆+Al)(2h)+Al₂O₃(1h) is incorporated into semi solid alloy, Al₃Ti particles releases into molten alloy. As Al₃Ti particles have higher driving-power relative to alumina particles, they cut α-aluminum more easily (Fig. 8a), brings about residing Al₃Ti particles in grain boundaries (Fig. 7a and c). The resultant microstructure of A206-(K₂TiF₆+Al)(2h)+Al₂O₃(1h), composite is schematically shown in Fig. 8b. This composite demonstrates a microstructure with a unique combination of lean matrix region and reinforcement rich zone appeared in a network fashion around AlA206 matrix grain boundaries.

In the case of A206-(K₂TiF₆+Al +Al₂O₃)(5h) sample, when (K₂TiF₆+Al +Al₂O₃)(5h) is incorporated into semi solid alloy, aluminum powder in capsule-shaped particles melts and releases alumina particles into molten alloy. As stirring proceeds, alumina particles entraps by α-aluminum. This can be ascribed to the lower driving-power of fine alumina particles rather than that of agglomerated particles. As depicted schematically in Fig. 9a, agglomerated alumina particles have higher driving-power giving rise to cut through α-aluminum more easily. The driving-power of alumina particles may not be adequate to do the same, and they may become entrapped in α-aluminum grain interior, Fig. 9b. Resultantly, a microstructure mainly containing Al₃Ti and alumina in grains are formed, Fig. 9c. It is plausible to say that this hypothesis has a bottle neck in the case of A206-(K₂TiF₆+Al)(2h)+Al₂O₃(1h) where local increment in temperature is not co-exist to fortify driving-power of alumina particles in such a manner that observed in A206-(K₂TiF₆+Al +Al₂O₃)(5h) sample.
Resultantly, in A206-(K₂TiF₆+Al)_{2h}+Al₂O₃_{1h} sample, alumina particles were mainly found in grain boundaries. Therefore, this microstructure possesses a unique combination of alumina and Al₃Ti in grain boundaries.

Another difference between A206-(K₂TiF₆+Al + Al₂O₃)_{3h} and A206-(K₂TiF₆+Al)_{2h}+Al₂O₃_{1h} is related to the size of Al₃Ti particles. Al₃Ti is smaller in the latter. Because in the former, Al₃Ti is formed in situ in the semi solid state while in latter Al₃Ti has been formed before powder addition (in the milling process). Temperature in the semi solid state is more enough to enlarge the in situ formed Al₃Ti particles. However, in the case of A206-(K₂TiF₆+Al)_{2h}+Al₂O₃_{1h}, Al₃Ti particles was formed in the milling process. Ball mill has high enough energy to break Al₃Ti particles.

Fig. 8. Schematic illustration showing the distribution of (a) Al₃Ti particle and (b) final microstructure of A206-(K₂TiF₆+Al)_{2h}+Al₂O₃_{1h}.

Fig. 9. Schematic illustration showing the distribution of (a) agglomerated alumina particle, (b) alumina particle and (c) final microstructure of A206-(K₂TiF₆+Al + Al₂O₃)_{3h}.
It is worth noting that in the case of A206-(K$_2$TiF$_6$+Al +Al$_2$O$_3$)$_{(5h)}$ sample, a composite containing ex situ alumina particle and in situ Al$_3$Ti particles was successfully tailored while in the case of A206-(K$_2$TiF$_6$+Al)$_{(3h)}$+Al$_2$O$_3$(1h)$_{(5h)}$, a composite containing ex situ alumina and Al$_3$Ti particles was formed.

3.4. Tensile properties

Fig. 10 shows tensile properties of A206, A206-(K$_2$TiF$_6$+Al+Al$_2$O$_3$)$_{(3h)}$ and A206-(K$_2$TiF$_6$+Al)$_{(2h)}$+Al$_2$O$_3$(1h)$_{(5h)}$ samples. As is observed, Al$_2$O$_{3\text{np}}$+Al$_3$Ti$_p$ improved tensile properties of AlA206 alloy, significantly.

On the other hand, it can be observed that A206-(K$_2$TiF$_6$+Al)$_{(2h)}$+Al$_2$O$_3$(1h)$_{(5h)}$ network-structure composite containing Al$_3$Ti$_p$ and Al$_2$O$_{3\text{np}}$ displays superior strength compared to A206-(K$_2$TiF$_6$+Al +Al$_2$O$_3$)$_{(5h)}$ composite. A206-(K$_2$TiF$_6$+Al)$_{(2h)}$+Al$_2$O$_3$(1h)$_{(5h)}$ composite shows 80.1% and 20.69% increase in ultimate tensile strength when compared to the as-received A206 and A206-(K$_2$TiF$_6$+Al +Al$_2$O$_3$)$_{(5h)}$ composite, respectively. These augmented tensile properties can be ascribed to the unique microstructure of the composite which contains two phases, as shown in Fig. 7a. The enhanced tensile properties of A206-(K$_2$TiF$_6$+Al)$_{(5h)}$+Al$_2$O$_3$(1h)$_{(5h)}$ network-structure composite can be explicated by the (H-S) theorem (Hashin-Shtrikman) [34] where Al$_3$Ti$_p$ +Al$_2$O$_{3\text{np}}$ stronger phase surrounds AlA206 matrix weaker phase. On the other word, Al$_3$Ti$_p$ and Al$_2$O$_{3\text{np}}$ concentrated around grain boundaries of AlA206 matrix, act as a composite with high volume fraction of Al$_3$Ti$_p$ +Al$_2$O$_{3\text{np}}$. Resultantly, the surprising feature of this unique structure is a matrix with fortified grain boundaries. In such a structure, dispersed high local Al$_3$Ti$_p$ +Al$_2$O$_{3\text{np}}$ in grain boundaries produces a synergistic effect via forming a three-dimension skeletal structure. The resistance to slip in such structure is high because this unique structure has a main contribution in barricading dislocation movement. That is to say, A206-(K$_2$TiF$_6$+Al)$_{(5h)}$+Al$_2$O$_3$(1h)$_{(5h)}$ network-structure composite containing reinforcing particles around grain boundaries can effectively strengthen grain boundaries by enhancing dislocation accumulation density during tensile deformation (Figs. 11 a and b).
3.4.1. Fracture behavior

Fig. 12 shows fracture side views of a network-structure A206-(K₂TiF₆+Al)₁₅₃₋₃₈₃₃+Al₂O₃₁₁ hybrid composite. Crack propagation occurs through Al₃Ti₁₁p +Al₂O₃₃₃ network-structure. Formation of micro-crack away from...
fracture surface in reinforcement network-structure brings to mind two interesting phenomenon; firstly, crack propagation take place via micro-crack coalescence through reinforcement network-structure and secondly, crack propagation to matrix-lean region is effectively barricaded by Al₃Ti₂ + Al₂O₃ network-structure (indicated by arrow), contributing to enhanced tensile strength of A206-(K₂TiF₆+Al)₅h + Al₂O₃ hybrid composite.

Fig. 13 shows fracture surface of a network-structure A206-(K₂TiF₆+Al)₅h + Al₂O₃ hybrid composite. As can be observed in Fig. 13a, there is no sign of large Al₂O₃ matrix rupture, fortifying crack propagation through reinforcement network-structure.

As demonstrated in Fig. 13b, Al₃Ti particle showed brittle cleavage fracture confirming strong interfacial bonding strength between matrix and reinforcement and consequently load bearing capacity of the composite. In tensile test, as load is proceeding, dislocation pile-up at matrix/reinforcement results in initiation and fracture of these particles. Thus, matrix cracking extended in to Al₃Ti particle (indicated by arrow on Fig. 13b). Subsequently, Al₂O₃ matrix experienced plastic deformation appeared in the form of dimples. Formation of such dimples further authenticating that Al₂O₃ matrix is continuous and interpenetrating.
4. Conclusions

Al/A206-5%Al2O3-Al3Ti hybrid composites were successfully synthesized using milled mixture of K2TiF6, alumina and aluminium powders and stir-casting technology. The following conclusions can be deduced:

1- Milling of (K2TiF6+Al+Al2O3)(2h) mixture impedes Al3Ti formation, while milled mixture of (K2TiF6+Al)(5h)+Al2O3(1h) causes the formation of Al3Ti particles.

2- Milling process reduced the formation temperature of Al3Ti particles and changed the morphology of Al3Ti particles.

3- Injection of Milled (K2TiF6+Al+Al2O3)(2h) mixture in semi solid alloy results in the formation of well-distributed alumina and Al3Ti in grain interiors while (K2TiF6+Al)(5h)+Al2O3(1h) milled mixture, brings about the formation of a reinforcement network structure.

4- Formation of Al3Ti with coherent interface in injected milled (K2TiF6+Al+Al2O3)(5h) mixture caused the uniform distribution of Al3Ti in grain interior. Alumina nano particulate with low-driving power entrapped by α-aluminum and resided in grain interior.
5- In injected milled \((K_2TiF_6+Al)(5h)+Al_2O_3(1h)\) mixture, released Al\(_3\)Ti in molten alloy had higher driving-power to cut \(\alpha\)-aluminum and therefore, Al\(_3\)Ti resided in grain boundaries.

6- Network structured A206-(\(K_2TiF_6+Al\))(2h)+\(Al_2O_3(1h)\) composite with fortified grain boundaries displayed superior tensile properties compared with \((K_2TiF_6+Al+Al_2O_3)(5h)\) composite ascribed to fortified grain boundaries by Al\(_3\)TiP + Al\(_2\)O\(_3\)np.

References:


