Impact of Intermetallic Precipitates on the Tribological and/or Corrosion Performance of Cast Aluminium Alloys: a Short Review

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Impact of intermetallic precipitates on the tribological and/or corrosion performance of cast aluminium alloys: a short review

D. Culliton*1, A. J. Betts2 and D. Kennedy1

The role of various intermetallic precipitates (IMPs), or secondary phase particles, in governing the wear and corrosion performance characteristics of cast aluminium alloys is outlined in this brief review. Such alloys are especially important in transport applications where their low weight, low cost and recyclability make them very attractive. However, alloy wear and/or corrosion behaviour often limit their industrial application, and more work needs to be carried out to extend their use into other areas. Careful control of IMP nucleation and growth rates may be beneficial, especially in alloys exposed to corrosive environments. Silicon, copper and magnesium are all important elements for enhanced mechanical strength and tribological performance but often to the detriment of alloy corrosion resistance. Other elements such as iron may also play a significant role in deleterious IMP formation. Use of dispersoids based on novel (quasicrystals) seed alloys with similar lattice characteristics to the α-Al matrix may result in further exploitation of these alloys.

Keywords: Cast aluminium alloys, Intermetallics, Dispersoids, Quasicrystals, Corrosion, Wear

Introduction

The automotive industry needs to produce cost efficient integral components of complex geometry, and casting is the most pragmatic solution. In addition, aluminium is the most abundant metal in the Earth’s crust.1 As a result, since 2006, alloys of this metal have become the second most used materials in the automotive industry (Table 1), and global consumption of aluminium is predicted to double between 2006 and 2020.2 This has made the automotive industry the largest market for aluminium alloys, more than half of which are cast Al alloys (Table 2). This transition has been primarily motivated by two global concerns:

(i) the depletion of world resources, necessitating the use of lighter, more abundant materials
(ii) international pressure by governments compelling industry to replace heavier metals with lighter, more efficient, recyclable materials. For instance, in vehicles, a weight reduction of 10% can increase fuel economy by 6–8%.3 Substituting Al alloys for steel and grey cast iron can achieve a weight reduction of up to 50% and allow savings of ~3000 L of fuel and 7500 kg of CO2 over the lifetime of an average car.4 However, cast Al alloys are currently only used in either mildly aggressive tribological5,6 or corrosive7,8 environments. This is related to the alloy microstructures and, more specifically, intermetallic precipitates (IMPs). Greater control of IMP growth in cast Al alloys could result in further exploitation of these alloys.

The tribological, mechanical and corrosion properties of cast Al alloys are defined by the solid solubilities of alloying elements and impurities present in the melt. The properties of the resultant IMP phases control both the mechanical and electrochemical properties of the alloy through their morphology, hardness, distribution and chemistry. Control of these precipitated phases is therefore key to improving and defining the properties of these alloys. Small, well distributed, spherical precipitates can be beneficial to both wear and mechanical properties and corrosion resistance. Inversely, large, acicular precipitates act as crack and corrosion initiators, thus diminishing the properties of the alloy. Heat treatments are used to redistribute and resize IMPs, but these treatments may lead to surface activation and a reduction in the corrosion resistance of the alloy.15,16

This short review identifies novel techniques for simultaneous improvement of both the mechanical properties and corrosion resistance of typical cast automotive Al alloys thereby expanding their industrial future beyond current applications.

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Table 1: Main alloying elements and typical intermetallic phases (IMP) of typical cast aluminium alloys used in automotive industry (242-0/A242-0 also contain 1-70-2-30%Ni)

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Si%/</th>
<th>Cu%/</th>
<th>Fe%/</th>
<th>Mn%/</th>
<th>Mg%/</th>
<th>Zn%/</th>
<th>Typical IMPs</th>
</tr>
</thead>
<tbody>
<tr>
<td>242-0</td>
<td>0-60 max.</td>
<td>3.70-4.50</td>
<td>0-1.6 max.</td>
<td>0-10 max.</td>
<td>1.30-1.70</td>
<td>0-10 max.</td>
<td>β-Al2FeSi, β-AlCu, AlCuNi, AlNi,</td>
</tr>
<tr>
<td>A242</td>
<td>0-60 max.</td>
<td>3.70-4.50</td>
<td>0-10 max.</td>
<td>1.30-1.70</td>
<td>0-10 max.</td>
<td>0.35 max.</td>
<td>Al(Ni, Cu)2, Al2CuFe, Al2Cu3Mn2</td>
</tr>
<tr>
<td>319-0</td>
<td>5-50-6-50</td>
<td>3-0-4-0</td>
<td>1.0 max.</td>
<td>0.50 max.</td>
<td>0-10 max.</td>
<td>0-10 max.</td>
<td>β-Al2FeSi, MgSi, AlCu, AlMgSi6Cu2</td>
</tr>
<tr>
<td>A319</td>
<td>5-50-6-50</td>
<td>3-0-4-0</td>
<td>1.0 max.</td>
<td>0.50 max.</td>
<td>0-10 max.</td>
<td>3.0 max.</td>
<td></td>
</tr>
<tr>
<td>B319</td>
<td>5-50-6-50</td>
<td>3-0-4-0</td>
<td>1.2 max.</td>
<td>0.50 max.</td>
<td>0-10-0-50</td>
<td>1.0 max.</td>
<td></td>
</tr>
<tr>
<td>356-0</td>
<td>6-50-7-50</td>
<td>0.25 max.</td>
<td>0.60 max.</td>
<td>0-35 max.</td>
<td>0-20-0-45</td>
<td>0-25-0-45</td>
<td>0-10 max.</td>
</tr>
<tr>
<td>A356</td>
<td>6-50-7-50</td>
<td>0.20 max.</td>
<td>0.20 max.</td>
<td>0.50 max.</td>
<td>0-10 max.</td>
<td>0-25-0-45</td>
<td>0-10 max.</td>
</tr>
<tr>
<td>A380</td>
<td>7-50-9-50</td>
<td>3-0-4-0</td>
<td>1.30 max.</td>
<td>0.50 max.</td>
<td>0-10 max.</td>
<td>3.0 max.</td>
<td>β-Al2FeSi, Al2Cu, β-AlMg3FeSi6</td>
</tr>
<tr>
<td>B380</td>
<td>7-50-9-50</td>
<td>3-0-4-0</td>
<td>1.30 max.</td>
<td>0.50 max.</td>
<td>0-10 max.</td>
<td>1.0 max.</td>
<td>Al2Mg3CuSi6</td>
</tr>
<tr>
<td>A390</td>
<td>16-0-18-0</td>
<td>4-0-5-0</td>
<td>0.50 max.</td>
<td>0-10 max.</td>
<td>0-45-0-65</td>
<td>0-10 max.</td>
<td>Mg2Si, Al2Cu, Al2Cu3Mg3Si6</td>
</tr>
<tr>
<td>B390</td>
<td>16-0-18-0</td>
<td>4-0-5-0</td>
<td>1.30 max.</td>
<td>0.50 max.</td>
<td>0-45-0-65</td>
<td>1.50 max.</td>
<td></td>
</tr>
</tbody>
</table>

Discussion

Typically, automotive components must have dynamic thermal and mechanical stability, good high temperature and fatigue strength, low thermal expansion and good wear resistance.

In addition, resistance to localised corrosion degradation processes, such as pitting, galvanic and filiform corrosion, in the presence of hydroxides (OH⁻), halides (Cl⁻) and sulphates (SO₄²⁻) improves the life to failure of the components. These requirements are strongly related to the alloying elements used. In cast automotive Al alloys, primary alloying elements include Si, Cu and Mg. While silicon improves the castability of the alloys through improved fluidity and increased tendency to isothermal solidification, it also increases the wear resistance of the alloy. Mg improves the strength and hardness of the alloy through the precipitation of the Mg2Si IMP. In combination with copper, the strength and hardness of the resultant castings are further improved due to the precipitation of the Al2Cu or Al2CuMg IMPS. While Al–Cu (2xx.x) alloys tend to have a finer grain structure than Al–Si–Mg (3xx.x) alloys, the presence of Al2CuMg IMPs. While Al–Cu (2xx.x) alloys tend to have a coarse, acicular structure due to reduced addition, IMPs with structures such as Chinese Script are preferred to coarse, acicular structures due to reduced impact on the mechanical properties.

In addition, alloying elements can also affect the development of other IMPS. For example, the precipitation of Fe, Mn and Ni, which can significantly higher hardness (Table 4) than the Al6Mn, Al2(Cu,Mn) and Al3(AlMnSi) phases and is much more desirable for wear related tribological applications. In addition, IMPS with structures such as Chinese Script are preferred to coarse, acicular structures due to reduced impact on the mechanical properties.

Table 2: Examples of automotive cast Al alloys, their typical uses and main failure modes: typically automotive corrosion occurs in form of pitting or galvanic corrosion caused by exposure to chloride (Cl⁻) or sulphate (SO₄²⁻) environments

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Typical components</th>
<th>Predominant degradation processes</th>
</tr>
</thead>
<tbody>
<tr>
<td>242-0</td>
<td>Heavy duty pistons, aircraft generator housings, air cooled cylinder heads</td>
<td>Wear (abrasive, adhesive)</td>
</tr>
<tr>
<td>319-0</td>
<td>Engine crankcases, petrol and oil tanks, oil pans, water cooled cylinder heads, rear axle housings</td>
<td>Corrosion</td>
</tr>
<tr>
<td>356-0</td>
<td>Flywheel housings, automotive transmission cases, oil pans, rear axle housings, brackets, water cooled cylinder blocks, various fittings and pump bodies</td>
<td>Corrosion</td>
</tr>
<tr>
<td>A380</td>
<td>Air brake castings, gear cases, air cooled cylinder heads</td>
<td>Tribological (friction)</td>
</tr>
<tr>
<td>A390</td>
<td>Internal combustion engine pistons and blocks, cylinder bodies for compressors, pumps and brakes</td>
<td>Wear (abrasive, adhesive)</td>
</tr>
</tbody>
</table>

Intermetallics

The precipitation and growth of the IMPS is dependent on the solid solubility of the respective alloying elements in the α-Al matrix and in each other (Table 3). The resultant microstructural developments define the subsequent tribological and corrosion properties of the alloys. Typically, coarse particles with sizes larger than 1 μm will deteriorate properties, such as toughness and fatigue performance, and are detrimental to the corrosion resistance of the alloys. This is also dependent on the alloying elements present. For example, the z-Al2(Fe,Mn)Si has significantly higher hardness (Table 4) than the Al6Mn, Al2(Cu,Mn) and z-(AlMnSi) phases and is much more desirable for wear related tribological applications.

In addition, alloying elements can also affect the development of other IMPS. For example, the presence of Cu leads to a change in the morphology of the eutectic Si particles from a coarse flake-like form to a fine fibrous one, with a resultant improvement in tribological performance. Mg can interfere with the precipitation of Al2Cu and, if present in relatively high amounts (>0-05%), can promote the precipitation of Al2Mg3Cu (S phase) and Al2Cu3Mg3Si6. The precipitate of Cu, Mn precipitates out as a dispersoid (T-phase) and is much more desirable for wear related tribological applications since the α-Al matrix can only contain Si, Cu, Mg and Zn, the presence of other chemical elements, such as Fe, Mn and Ni, will only develop intermetallic phases. The size, morphology and chemistry of these IMPS strongly influence the
mechanical properties and corrosion resistance of the resultant alloys (Table 4).

Generally speaking, the larger the IMP, the more detrimental it is likely to be to the mechanical/tribological performance of the alloy and its corrosion resistance. For example, above a critical IMP size (1 μm), a direct relationship between corrosion rate and IMP size has been reported by Colley et al. who found a decrease in the corrosion rate with decreasing IMP size. IMPs that grow freely within the solidifying liquid tend to grow much larger than those that form during or after the period of Al–Si eutectic solidification.

By increasing the cooling rate, pooling of the alloying elements between the dendrite arms may be controlled and IMP growth can be restricted. This may lead to a more homogeneous microstructure with smaller, more dispersed IMPs. It has been shown that, as the cooling rate decreases, the average IMP size increases and the strength of the casting decreases. Therefore, the key to simultaneous improvement of both the mechanical properties and the corrosion resistance of Al alloys is through controlled precipitation of the IMP phases by expediting nucleation, reducing the secondary dendrite arm spacing (SDAS) and homogenising solidification rates throughout the casting.

Table 3 Possible intermetallic precipitates formed in typical automotive cast Al alloys

<table>
<thead>
<tr>
<th>Intermetallic phase</th>
<th>Typical size/μm</th>
<th>Morphology</th>
<th>Intermetallic phase</th>
<th>Typical size/μm</th>
<th>Morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td>SI</td>
<td>5–12</td>
<td>Acicular</td>
<td>SI</td>
<td>5–12</td>
<td>Acicular</td>
</tr>
<tr>
<td>Al2Cu</td>
<td>θ &lt;50</td>
<td>Angular globule</td>
<td>Al2Cu</td>
<td>θ &lt;50</td>
<td>Angular globule</td>
</tr>
<tr>
<td>Al2CuMg</td>
<td>S 0.5–10</td>
<td>Irregular round particles</td>
<td>Al2CuMg</td>
<td>S 0.5–10</td>
<td>Irregular round particles</td>
</tr>
<tr>
<td>Al2CuFe</td>
<td>α 0.7–2.7</td>
<td>Thin needles</td>
<td>Al2CuFe</td>
<td>α 0.7–2.7</td>
<td>Thin needles</td>
</tr>
<tr>
<td>Al2CuMgSi6</td>
<td>Q &lt;20</td>
<td>Honeycomb</td>
<td>Al2CuMgSi6</td>
<td>Q &lt;20</td>
<td>Honeycomb</td>
</tr>
<tr>
<td>Al(Fe,Mn)3Si</td>
<td>µ 50–150μm</td>
<td>Needles</td>
<td>Al(Fe,Mn)3Si</td>
<td>µ 50–150μm</td>
<td>Needles</td>
</tr>
<tr>
<td>Al5Cu2Mg8Si6</td>
<td>T 0.1</td>
<td>Dispersoid</td>
<td>Al5Cu2Mg8Si6</td>
<td>T 0.1</td>
<td>Dispersoid</td>
</tr>
<tr>
<td>Al5Cu2Mg8Si6</td>
<td>T 0.1</td>
<td>Lamellar, rod-like</td>
<td>Al5Cu2Mg8Si6</td>
<td>T 0.1</td>
<td>Lamellar, rod-like</td>
</tr>
</tbody>
</table>

Table 4 Typical phases found in 2xx.x and 3xx.x cast aluminium alloys and their reported hardness and corrosion potentials in various NaCl molar solutions

<table>
<thead>
<tr>
<th>Main alloying elements and intermetallic phases</th>
<th>Hardness/GPa</th>
<th>E&lt;sub&gt;corr&lt;/sub&gt;(vs SCE) in aqueous NaCl/mV</th>
</tr>
</thead>
<tbody>
<tr>
<td>x-Al</td>
<td>0.167</td>
<td>-679</td>
</tr>
<tr>
<td>Si</td>
<td>10.0&lt;sup&gt;52&lt;/sup&gt;–3.3&lt;sup&gt;56&lt;/sup&gt;</td>
<td>-450</td>
</tr>
<tr>
<td>Cu</td>
<td>0.389</td>
<td>-1077</td>
</tr>
<tr>
<td>Mg</td>
<td>0.462</td>
<td>-1601</td>
</tr>
<tr>
<td>Al&lt;sub&gt;2&lt;/sub&gt;Fe</td>
<td>7.36&lt;sup&gt;57&lt;/sup&gt;</td>
<td>-493</td>
</tr>
<tr>
<td>x-AlFeSi (Al&lt;sub&gt;x&lt;/sub&gt;Fe&lt;sub&gt;y&lt;/sub&gt;Si, Al&lt;sub&gt;2&lt;/sub&gt;Fe&lt;sub&gt;3&lt;/sub&gt;Si)&lt;sup&gt;58&lt;/sup&gt;</td>
<td>12.3–14.9&lt;sup&gt;59&lt;/sup&gt;</td>
<td>-1651</td>
</tr>
<tr>
<td>β-AlFeSi (Al&lt;sub&gt;x&lt;/sub&gt;Fe&lt;sub&gt;y&lt;/sub&gt;Si), γ-AlFeSi (Al&lt;sub&gt;x&lt;/sub&gt;Fe&lt;sub&gt;y&lt;/sub&gt;Si)</td>
<td>5.8&lt;sup&gt;60&lt;/sup&gt;</td>
<td>-1832</td>
</tr>
<tr>
<td>Al&lt;sub&gt;5&lt;/sub&gt;Cu2Mg8Si6</td>
<td>15.6–17.4&lt;sup&gt;59&lt;/sup&gt;</td>
<td>-1969</td>
</tr>
<tr>
<td>x-Al(Fe,Mn)3Si</td>
<td>4.74&lt;sup&gt;61&lt;/sup&gt;</td>
<td>-2168</td>
</tr>
<tr>
<td>x-AlFeMg3Si6</td>
<td>5.8&lt;sup&gt;53&lt;/sup&gt;</td>
<td>-2322</td>
</tr>
<tr>
<td>β-Mg2Si</td>
<td>4.555</td>
<td>-1355</td>
</tr>
<tr>
<td>γ-Al2Cu</td>
<td>7.6–8.5&lt;sup&gt;55&lt;/sup&gt;</td>
<td>-592</td>
</tr>
<tr>
<td>S-Al2CuMg</td>
<td>3.7–3.9&lt;sup&gt;56&lt;/sup&gt;</td>
<td>-956</td>
</tr>
<tr>
<td>β-Mg2Si</td>
<td>4.555</td>
<td>-1355</td>
</tr>
<tr>
<td>γ-Al2CuFe</td>
<td>9.49&lt;sup&gt;50&lt;/sup&gt;</td>
<td>-549</td>
</tr>
<tr>
<td>Q-Al2CuMgMgSi6</td>
<td>6.51&lt;sup&gt;63&lt;/sup&gt;</td>
<td>-551</td>
</tr>
</tbody>
</table>

Tribology, corrosion and environments

Wear resistant cast Al Alloys are based on the Al–Si alloy range, due to their relatively high density (2.6 g cm<sup>-3</sup>) and the excellent hardness of the diamond cubic shaped silicon phase (Table 4). In addition, they possess low thermal expansion coefficients, relatively good corrosion resistance and favourable mechanical properties. Higher silicon content improves fluidity, feeding characteristics and hot cracking resistance but may also lead to reduced density of the casting. An increase in porosity and surface roughness and, above 11%, can reduce the wear resistance of the alloy. Si-particle morphology and density can be altered through the addition of Sr. Bai and Biswas and Sarkar and Clarke have stated that the silicon content has no influence on the friction coefficient of Al–Si alloys. In contrast, Mahato et al. has suggested that the proliferation and morphology of protruding silicon particles could lead to an increase in wear resistance. However, this is only possible if the working pressure of the sliding system is less than that of the yield strength of the aluminium matrix (~80 MPa), as at higher loads the particles disintegrate and become dispersed in the deformed layer. It has been reported that, in wear applications, additions of up to...
11% Si improved the wear resistance, while above 11%, the wear rate increased again. As Si content increases, Si particle size increases, sometimes up to 2–3 μm,52 which can be detrimental to strength, ductility and fracture toughness.53

Prasad et al.64 reported on the improvement in wear properties for refined Al–Si microstructures. In their study of the effect of silicon content on the wear resistance of Al–Si alloys (LM13 [Al–Si12CuFe] and LM29 [Al–Si23CuMgNi]), they showed that the mechanical and wear properties of the castings were a function of the size and amount of primary silicon present. Refining the primary silicon particles and, as a consequence, the SDAS, through choice of casting process, produced castings with superior wear properties. The importance of Si phase refinement was also stressed by Yust65 who reported that, while small evenly dispersed particles improved ductility without reducing strength, coarse acicular particles reduced ductility because they acted as crack initiators. Post-treatments have been shown to alter the shape of the as cast silicon phase.66 It is suggested that this improvement comes from a refinement of the microstructure67 and alteration of the morphology of the silicon precipitates.

The wear properties of Al–Si alloys can be improved by the addition of some alloying elements, such as Cu and Ni, which produce hard intermetallic phases.68 Beneficial effects are dependent on the size, distribution and morphology of these particles with smaller, well distributed, spherical particles considered to be more beneficial. For instance, in copper containing Al alloys, increased wear resistance has been associated with precipitation of the small, brittle β-Al2Cu phase at the surface.69 The addition of 1% Cu to Al–Si alloys increased the transition load for mild to severe wear by three to four times that of the original alloys by increasing the size and amount of primary silicon by Yust.65 It is suggested that this improvement comes from a refinement of the microstructure67 and alteration of the morphology of the silicon precipitates.

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The corrosion resistance of Al alloys is controlled by the composition and microstructure73 and is primarily affected by the α-Al matrix phase. In the presence of aggressive ions (Cl–, SO42– and OH–), aluminium can be selectively dissolved, depending on the alloying elements present. The corrosion resistance of the α-Al matrix phase is ennobled (made less negative relative to pure aluminium, Table 4) when Fe, Cu, Mn and Si are added, while Mg and Zn shift the potential to a less noble state84 (made more negative relative to pure aluminium, Table 4). For each element, the significant changes in the corrosion resistance occur within the range in which the element is completely in solid solution. Further additions of the same element form microscopic second phase particles (shell particles) or IMPs. These IMPs,72 such as Al2Cu, α-AlFeSi, β-Al2FeSi, Al3Mg2 or Al2CuMg, prevent the homogeneous formation of a protective oxide layer76 and generally act as ‘active sites’77 for corrosion initiation.78 This results in the localised dissolution of the α-Al matrix,83–85 deaulying of the Mg based IMPs72,84 and the formation of corrosion pits.85 Therefore, the corrosion resistance of an Al alloy is dependent on the size, composition and distribution of the IMPs and their relative nobility to the surrounding α-Al matrix.

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Corrosion resistance can be defined by the Open Circuit Potential or corrosion potential, Ecorr, of the alloy in aqueous solutions and is related to the cathodically driven oxygen reduction reaction at and around the IMPs. Both Cu and Fe rich IMPs (more noble than the α-Al matrix, Table 4) can serve as cathodes for this reaction with similar efficiency,76 thus driving the corrosion of the surrounding aluminium. Increasing the amount of Cu or Fe in an alloy increases the corrosion rate.57 However, IMPs capable of sustaining the largest cathodic current densities are not necessarily those with the noblest Ecatt (Table 4). Similarly, those with the least noble Ecatt will not necessarily sustain the largest anodic currents.54,86 Hence, not only thermodynamic but also kinetic aspects are important to consider when exploring the role of IMPs in corrosion degradation. However, a detailed discussion of these corrosion phenomena is outside the scope of this short review and can be found elsewhere.89 Birbils and Buchheit54 reviewed the effect of IMP composition on the corrosion of a typical Al–Cu alloy. They found that, although the Fe IMPs were quantified as being more noble than the surrounding matrix, the Cu IMPs were far more deleterious. They attributed this to the ability of the IMPs to support extended cathodic reactions (oxygen reduction), which resulted in corrosion rates an order of magnitude higher than those of the Fe based IMPs. These Cu based IMPs are generally of the form Al12Cu or Al12CuMg, and the higher corrosion rates may be related to the plating out of Cu.90 As noted in Table 3, these IMPs are larger than the critical size of 1μm. Since this behaviour is strongly linked with the size of the IMPs, as well as the composition, control of IMP precipitation and growth is critical to improving alloy properties especially in corrosive fluids.

**Intermetallic precipitation and solidification**

Al alloy solidification is a highly complex process. In general, however, three main solidification reactions are exhibited during the solidification process.91,92 Initially, aluminium dendrites (liquidus) are formed, followed by the development of two main eutectic phases. The presence of alloying and impurity elements, such as Cu, Mg, Mn and Fe, leads to more complex constituents. Preliminary models of microstructural development, such as the isothermal melt model,93 were based on the free growth criterion, which assumed that the rate determining step to the formation of a grain is not nucleation, but rather overcoming the energy barrier relating to the curvature of the solid/liquid interface (the Gibbs–Thomson effect). These models are very simplistic and are only suitable for predicting grain size for a range of solute levels (inoculant particle populations) in small, slowly cooled castings.

Recent studies94,95 have shown that the solute elements in the liquid ahead of the growing crystals reduce the growth rate velocity of the nucleated crystals and increase the maximum undercooling achievable before recalescence. This allows more particles to be active in nucleation and, consequently, increases the number density of active particles, giving rise to a finer grain size. As a result, increasing the cooling rate can
restrict the size, composition and morphology of the resultant IMPs.

Attempts to model the effect of alloy composition and cooling rate on the mechanical properties of these alloys have shown that smaller, more spherical intermetallics, combined with lower SDAS, produce the highest strength in the Al–Ti–C system. The SDAS depends on the chemical composition of the alloy, the solidification rate and the temperature gradient on the crystallisation front. In addition, the SDAS can control the precipitation and growth of IMPs. Therefore, the priority for property improvement must be to reduce the SDAS by increasing the cooling rate and in larger castings; this can only be achieved by introducing dispersoids into the melt.

**Dispersoids and intermetallics**

Dispersoids have been used successfully in many forms of aluminium alloy casting, and the ideal atomic structures of the dispersoid should be close to the face centred cubic (fcc) structure of the α-Al matrix. As stated previously, one of the key properties of a successful dispersoid is its ability to be wetted by the alloy matrix. Traditionally, grain refiners were based on the Al–Ti–B system. While these may have been reasonably successful, the resultant alloys were still limited by their properties. This may have been related to the structure of the dispersoids, which were hexagonal (Al–Ti–B) and octahedral (Al–Ti–C). It may also have been related to the low efficiency of the process, which can be <1%. More importantly, these inoculants do not directly impact the formation and growth of IMPs.

In Al alloys, magnesium is the most widely used wetted element. Most likely due to its fcc structure. However, while MgO has been used as a reinforcement in aluminium metal matrix composites, its use as a grain refining dispersoid has only been reported for cast magnesium alloys. Other oxides, such as aluminium (Al2O3), have also been used in aluminium metal matrix composites, but due to their dissimilar molecular structure (hexagonal close packed), it is unlikely that they would be as nucleating dispersoids. Since the ideal dispersoid is one that has high temperature stability and produces distortion of the lattice structure but remains coherent with this structure, oxide based dispersoids may not be the solution. In addition, due to the large density differences between aluminium and the alloying elements, it is necessary to promote elevated solidification rates from the melt to prevent extensive pooling of the alloying elements. Therefore, novel nucleating dispersoids are required.

For most cast Al alloys, the maximum operating temperature is restricted to 150 to 175°C. This has been attributed to the thermal instability and excessive growth of the strength providing IMPs, resulting in undesirable levels of lattice incoherencies, such as grain/phase boundaries. These lattice incoherencies provide sites for void coalescence, resulting in crack formation and growth under mechanical loading. However, in 2002, engineers at National Aeronautics and Space Administration developed a range of cast Al alloys, which had superior mechanical properties at elevated temperatures (230–400°C). This advancement was attributed to the stabilising effect of strengthening precipitates with an L12 crystal structure. The L12 crystal structure is a derivative of the fcc structure and is therefore highly coherent with the α-Al matrix. These IMPs were based on an Al3X structure (X=Ti, V, Zr). The high temperature stability of these precipitates, combined with the high level of coherence between the L12 structure and the fcc α-Al matrix, provided the improved performances. The stability of these IMPs may be further enhanced using a Ti4Al3X (X=Fe, Ni or Cu) structured dispersoid, and the stability of which increases with increasing atomic number (Fe<Cu). This is similar, in concept, to aluminium metal matrix composites, and based on this concept, it may be possible to not only improve the tribological performance of the alloy but also to improve the corrosion resistance of the same alloys. As a result, it is strongly suggested that by augmenting current alloy production methods with novel alloying techniques, superior properties can be achieved with a wide range of cast Al alloys. A range of dispersoids, which, as yet, have not been considered for cast Al alloys but have been successfully incorporated into steel alloys with dramatic effect, are quasicrystals (QCs).

**Novel dispersoids**

Quasicrystals, or quasi-periodic crystals, have a face centred icosahedral structure that is ordered but not periodic. Typical QC alloys include Al–Mn, Al–Mn–Si, Al–Cu–Mg, or Al–Mg–Cu. Currently used as protective coatings and in advanced steel alloys for medical applications, these materials have high hardness (low friction), excellent mechanical properties and high temperature stability. Similar to the L12 additives used in the National Aeronautics and Space Administration alloys, these structures have the potential of promoting the development of a refined microstructure, with improved mechanical/tribological properties and corrosion resistance. This should not be confused with dispersion strengthening, where strength enhancing dispersoids are added directly to the melt. These QC alloys have high lattice coherency and high temperature stability, though minimal deformation has been noted in some QC alloys at temperatures in excess of 750°C. By selecting quasicrystalline submicrometre particles (minimum particle size can be quantified from a number of different models based on suitable seed alloys, which have similar lattice parameters to the base alloy (high lattice coherency), wetting of the dispersoid can be maximised. As a result, these quasicrystalline dispersoids should act as optimum nucleation primers, producing homogeneous microstructures with smaller, well dispersed IMPs, leading to concomitant improvement of mechanical properties and corrosion resistance. In addition, the proliferation of these nucleation primers, particularly in larger castings, would increase the tendency to isothermal solidification. This would lead to increased cooling rates and reduced recalescence temperatures, resulting in smaller grain size and influencing the morphology, composition and distribution of the developed IMPs. Since the mechanical properties of the alloy are related to the dispersoid concentration, greater dispersoid distribution would increase the strength of the resultant alloys. Based on the five requirements suggested by Wang et al., QCs could be the basis of an optimised solution for microstructural control in cast Al alloys and need to be investigated further.

**Conclusions**

Current casting techniques do not produce aluminium alloys with sufficient wear and/or corrosion resistance for use in any but the most benign service environments.
Research aimed at improving their tribological and/or corrosion performance characteristics could potentially produce superior alloys, enabling their adoption in a more widespread industrial context and providing an opportunity to extend their use into new fields. These include applications in industries such as those in the burgeoning energy and marine sector, thus extending their somewhat limited use beyond the present transport sector (largely automotive and aerospace).

The tribological, mechanical and corrosion behaviour of cast Al alloys are largely defined by the solid solubilities of various alloying elements such as silicon, magnesium, copper and melt impurities. The resultant intermetallic phases can control alloy service performance through their morphology, hardness, distribution and chemical characteristics. Careful control of these precipitated phases is therefore essential. In general, small (i.e. submicrometre), evenly distributed, spherical precipitates can be beneficial to both wear and mechanical properties. In contrast large, acicular, precipitates may act as crack initiators, leading to reduced ductility. Although heat treatments are commonly used to redistribute and resize IMPs, these treatments may lead to surface activation with a concomitant reduction in the alloy’s corrosion resistance.

Dispersoids can be added to the melt to increase grain nucleation rates and reduce solidification times. However, these do not sufficiently impact the resultant properties of the alloys due to their incompatible crystal structures. One possible novel approach may be to utilise dispersoids based on fcc compatible QCs, which may also provide enhanced mechanical and corrosion behaviour. These novel materials generally possess high temperature stability and some have shown improved corrosion resistance in corrosive fluids. However, these structures have not, heretofore, been considered as dispersoids in Al alloy casting melts.

It is suggested that these novel structures could act as potential dispersoids, particularly in casting technologies where no mechanical processing of the melt occurs, such as gravity and sand casting. In addition, the thermal stability of these dispersoids implies that the developed submicrometre IMP should remain stable during any subsequent heat treatment. Although it may not be possible to entirely eliminate precipitation of deleterious IMPs, by increasing the nucleation sites and the cooling rate, strict controls can possibly be placed on the size of these IMPs, which may be achieved by incorporating such QCs into the melt.

This brief review highlights the important role of IMPs in governing the behaviour of cast Al alloys and identifies possible ways to overcome some of their present limitations.

References


