(12) STANDARD PATENT (11) Application No. AU 2009225257 B9 (19) AUSTRALIAN PATENT OFFICE					
(54)	Title Metal-coated steel strip				
(51)	International Patent Classification(s)C23C 2/06 (2006.01)C23C 2/14 (2006.01)C23C 2/12 (2006.01)C23C 2/28 (2006.01)				
(21)	Application No: 2009225257 (22) Date of Filing: 2009.03.13				
(87)	WIPO No: W009/111842				
(30)	Priority Data				
(31)	Number(32)Date(33)Country20089012232008.03.13AU20089012242008.03.13AU				
(43) (44) (48)	Publication Date:2009.09.17Accepted Journal Date:2014.09.25Corrigenda Journal Date:2020.05.07				
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(56)	Related Art JP 2002-012959 A US 6635359 B1				

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization International Bureau

(43) International Publication Date

17 September 2009 (17.09.2009)

- (51) International Patent Classification: C23C 2/06 (2006.01) C23C 2/12 (2006.01) C23C 2/14 (2006.01) C23C 2/28 (2006.01)
- (21) International Application Number:

PCT/AU2009/000305

- (22) International Filing Date: 13 March 2009 (13.03.2009)
- (25) Filing Language: English

(26) Publication Language: English

- (30) Priority Data: 2008901224 13 March 2008 (13.03.2008) AU 2008901223 13 March 2008 (13.03.2008) AU
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(10) International Publication Number

WO 2009/111842 A1

- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.
- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

with international search report (Art. 21(3))





Figure 1 Sr additions in a 55%h1-Zn-1.5%Si-2.0%Mg coating eliminate the surface mottling defect and change the distribution pattern of the Mg.Si phase in the coating thickness direction.

FIGURE 1

(57) Abstract: An Al-Zn-Si-Mg alloy coated strip that has Mg₂Si particles in the coating microstructure is disclosed. The distribution of Mg₂Si particles is such that the surface of the coating has only a small proportion of Mg₂Si particles or is at least substantially free of any Mg₂Si particles.



METAL-COATED STEEL STRIP

The present invention relates to strip, typically steel strip, which has a corrosion-resistant metal alloy 5 coating.

The present invention relates particularly to a corrosion-resistant metal alloy coating that contains aluminium-zinc-silicon-magnesium as the main elements in 10 the alloy, and is hereinafter referred to as an "Al-Zn-Si-Mg alloy" on this basis. The alloy coating may contain other elements that are present as deliberate alloying additions or as unavoidable impurities. Hence, the phrase "Al-Zn-Si-Mg alloy" is understood to cover alloys that 15 contain such other elements and the other elements may be deliberate alloying additions or as unavoidable impurities.

The present invention relates particularly but 20 not exclusively to steel strip that is coated with the above-described Al-Zn-Si-Mg alloy and can be cold formed (e.g. by roll forming) into an end-use product, such as roofing products.

25 Typically, the Al-Zn-Si-Mg alloy comprises the following ranges in % by weight of the elements aluminium, zinc, silicon, and magnesium:

	Aluminium:	40 to 60 %
30	Zinc:	4 0 to 60 %
	Silicon:	0.3 to 3%
	Magnesium	0.3 to 10 %

Typically, the corrosion-resistant metal alloy 35 coating is formed on steel strip by a hot dip coating method.

In the conventional hot-dip metal coating method, steel strip generally passes through one or more heat treatment furnaces and thereafter into and through a bath of molten metal alloy held in a coating pot. The heat treatment furnace that is adjacent a coating pot has an outlet snout that extends downwardly to a location below the upper surface of the bath.

The metal alloy is usually maintained molten in 10 the coating pot by the use of heating inductors. The strip usually exits the heat treatment furnaces via an outlet end section in the form of an elongated furnace exit chute or snout that dips into the bath. Within the bath the strip passes around one or more sink rolls and is 15 taken upwardly out of the bath and is coated with the metal alloy as it passes through the bath.

After leaving the coating bath the metal alloy coated strip passes through a coating thickness control 20 station, such as a gas knife or gas wiping station, at which its coated surfaces are subjected to jets of wiping

The metal alloy coated strip then passes through 25 a cooling section and is subjected to forced cooling.

gas to control the thickness of the coating.

The cooled metal alloy coated strip may thereafter be optionally conditioned by passing the coated strip successively through a skin pass rolling section (also known as a temper rolling section) and a tension

30 (also known as a temper rolling section) and a tension levelling section. The conditioned strip is coiled at a coiling station.

A 55%Al-Zn alloy coating is a well known metal 35 alloy coating for steel strip. After solidification, a 55%Al-Zn alloy coating normally consists of α-Al dendrites

and a β -Zn phase in the inter-dendritic regions of the coating.

It is known to add silicon to the coating alloy 5 composition to prevent excessive alloying between the steel substrate and the molten coating in the hot-dip coating method. A portion of the silicon takes part in a quaternary alloy layer formation but the majority of the silicon precipitates as needle-like, pure silicon 10 particles during solidification. These needle-like

- silicon particles are also present in the inter-dendritic regions of the coating.
- It has been found by the applicant that when Mg 15 is included in a 55%Al-Zn-Si alloy coating composition, Mg brings about certain beneficial effects on product performance, such as improved cut-edge protection, by changing the nature of corrosion products formed.
- 20 However, it has also been found by the applicant that Mg reacts with Si to form a Mg₂Si phase and that the formation of the Mg₂Si phase compromises the abovementioned beneficial effects of Mg in a number of ways.
- One particular way, which is the focus of the present invention is a surface defect called "mottling". The applicant has found that mottling can occur in Al-Zn-Si-Mg alloy coatings under certain solidification conditions. Mottling is related to the presence of the Mg₂Si phase on the coating surface.

More particularly, mottling is a defect where a large number of coarse Mg₂Si particles cluster together on the surface of the coating, resulting in a blotchy surface 35 appearance that is not acceptable from an aesthetic viewpoint. More particularly, the clustered Mg₂Si particles form darker regions approximately 1-5 mm in size and introduce non-uniformity in the appearance of the coating which makes the coated product unsuitable for applications where a uniform appearance is important.

5 The above description is not to be taken as an admission of the common general knowledge in Australia or elsewhere.

The present invention is an Al-Zn-Si-Mg alloy 10 coated strip that has Mg₂Si particles in the coating with the distribution of Mg₂Si particles being such that a surface of the coating has only a small proportion of Mg₂Si particles or is at least substantially free of any Mg₂Si particles.

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The applicant has found that the above-described distribution of Mg_2Si particles in the coating microstructure provides significant advantages and can be achieved by any one or more of:

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(a) strontium additions in the coating alloy,

(b) selection of the cooling rate during solidification of coated strip for a given coating mass
 25 (i.e. coating thickness) exiting a coating bath; and

(c) minimising variations in coating thickness.

- The applicant has found that Sr additions 30 described in more detail below control the distribution characteristics of the Mg₂Si phase in the thickness direction of an Al-Zn-Si-Mg alloy coating so that the surface of the coating has only a small proportion of Mg₂Si particles or is at least substantially free of Mg₂Si
- 35 particles, whereby there is a considerably lower risk of Mg₂Si mottling.

The applicant has found that when at least 250 ppm Sr, preferably 250-3000 ppm Sr, is added to a coating bath containing an Al-Zn-Si-Mg alloy the distribution characteristics of the Mg₂Si phase in the coating thickness 5 direction are completely changed by this addition of Sr from the distribution that is present when there is no Sr in the coating bath. Specifically, the applicant has found that these additions of Sr promote the formation of a surface of the coating that has only a small proportion 10 of Mg₂Si particles or is free of any Mg₂Si particles and consequently a considerably lower risk of mottling on the surface.

- The applicant has also found that selecting the 15 cooling rate during solidification of a coated strip exiting a coating bath to be below a threshhold cooling rate, typically below 80°C/sec for coating masses less than 100 grams per square metre of strip surface per side, controls the distribution characteristics of the Mg₂Si 20 phase so that the surface has only a small proportion of
- Mg₂Si particles or is at least substantially free of Mg₂Si particles, whereby there is a considerably lower risk of Mg₂Si mottling.
- 25 The applicant has also found that minimising coating thickness variations controls the distribution characteristics of the Mg₂Si phase so that the surface has only a small proportion of Mg₂Si particles or is at least substantially free of Mg₂Si particles, whereby there is a 30 considerably lower risk of Mg₂Si mottling. As is the case with Sr addition and selection of cooling rate during solidification, the resultant coating microstructure is advantageous in terms of appearance, enhanced corrosion resistance and improved coating ductility.

The claims define the invention in terms of minimising coating thickness variations to control the

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distribution characteristics of the Mg_2Si phase so that the surface has only a small proportion of Mg_2Si particles or is at least substantially free of Mg_2Si particles, whereby there is a considerably lower risk of Mg_2Si mottling..

Accordingly, the invention provides a hot-dip coating method for forming a coating of a corrosionresistant Al-Zn-Si-Mg alloy on a steel strip comprising passing the steel strip through a hot dip coating bath that contains Al, Zn, Si, and Mg and optionally other elements and forming an alloy coating on the strip with a variation in thickness of the coating of no more than 40% in any given 5 mm diameter section so that the distribution of Mg₂Si particles in the coating microstructure is such that there is only a small

proportion of Mg_2Si particles or substantially no Mg_2Si particles in the surface of the coating.

The method may comprise forming the alloy coating 20 so that there are no Mg₂Si particles in the surface of the coating.

The small proportion of Mg₂Si particles in the surface of the coating is no more than 10wt.% of the Mg₂Si 25 particles.

The coating thickness variation may be no more than 30% in any given 5 mm diameter section of the coating.

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For a coating thickness of $22\,\mu\text{m},$ the maximum thickness in any region of the coating greater than 1mm in diameter may be $27\,\mu\text{m}.$

35 The Al-Zn-Si-Mg alloy coating may comprise the following ranges in % by weight of the elements aluminium, zinc, silicon, and magnesium:

- 7 -

Aluminium:	40 to 60 %
Zinc:	40 to 60 %
Silicon:	0.3 to 3%
Magnesium	0.3 to 10 %

The Al-Zn-Si-Mg alloy may also contain other elements, such as, by way of example any one or more of iron, vanadium, chromium, and strontium.

10 The coating may contain more than 250 ppm Sr. The coating may contain more than 500 ppm Sr. The coating may contain more than 1000 ppm Sr. 15 The coating may contain less than 3000 ppm Sr.

The Al-Zn-Si-Mg-Sr alloy coating may contain other elements as deliberate additions or as unavoidable 20 impurities.

The method method may comprise forming the coating to have a thickness of less than $30\mu m$.

25 The method may comprise forming the coating to have a thickness of greater than 7µm.

The method may comprise selecting the cooling rate during solidification of coated strip exiting the 30 coating bath to be less than a threshold cooling rate.

In any given situation, the selection of an appropriate thickness variation is related to the coating thickness (or coating mass).

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The method may comprise selecting the cooling rate for coated strip exiting the coating bath to be less

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than $80^{\circ}C/sec$ for coating masses up to 75 grams per square metre of strip surface per side.

The method may comprise selecting the cooling 5 rate for coated strip exiting the coating bath to be less than 50°C/sec for coating masses of 75-100 grams per square metre of strip surface per side.

Typically, the method comprises selecting the 10 cooling rate to be at least 11°C/sec.

By way of example, for a coating having an average thickness of $22\mu m$, during solidification the cooling rates may be as follows:

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(a) 55°C/sec in a temperature range of 600-530°C,

(b) 70°C/sec in a temperature range of 530-500°C, and
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(c) 80°C/sec in a temperature range of 500-300°C.

The invention also provides a steel strip having a coating of a corrosion-resistant Al-Zn-Si-Mg alloy 25 formed by the above-described method.

The invention also provides an Al-Zn-Si-Mg alloy coated strip that has Mg₂Si particles in the coating with the distribution of Mg₂Si particles being such that the 30 surface of the coating has only a small proportion of Mg₂Si particles or is at least substantially free of any Mg₂Si particles formed by the above-described method.

The invention also provides a steel strip having 35 a coating of a corrosion-resistant Al-Zn-Si-Mg alloy, the coating having a variation in thickness of no more than 40% in any given 5 mm diameter section so that the

distribution of Mg_2Si particles in the coating is such that there is only a small proportion of Mg_2Si particles or substantially no Mg_2Si particles in a surface of the coating.

The_advantages of the invention include the following advantages.

- Elimination of mottling defect and improved first-time-prime production rate. The risk of the mottling defect is at least substantially eliminated and the surface of the resultant coating maintains a beautiful, silvery metallic appearance. As a result, first-timeprime production rate is improved and profitability is boosted.
- Prevention of mottling defect by the addition of Sr allows the use of higher cooling rates, reducing the length of cooling equipment required after the pot.

Example

- 25 The applicant has carried out laboratory experiments on a series of 55%Al-Zn-1.5%Si-2.0%Mg alloy compositions having up to 3000 ppm Sr coated on steel substrates.
- 30 The purpose of these experiments was to investigate the impact of Sr on mottling in the surface of the coatings.
- Figure 1 summarises the results of one set of 35 experiments carried out by the applicant that illustrate the present invention.

The left hand side of the Figure comprises a top plan view of a coated steel substrate and a cross-section through the coating with the coating comprising a 55%Al-Zn-1.5%Si-2.0%Mg alloy with no Sr. The coating was not formed having regard to the selection of cooling rate during solidification and coating thickness variations discussed above.

The mottling that results from such a coating 10 composition is identified by the arrow in the top plan view. It is evident from the cross-section that Mg₂Si particles are distributed throughout the coating thickness. This is a problem for the reasons stated above.

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The right hand side of the Figure comprises a top plan view of a coated steel substrate and a cross-section through the coating, with the coating comprising a 55%Al-Zn-1.5%Si-2.0%Mg alloy and 500 ppm Sr. A complete absence

- 20 of mottling is evident from the top plan view. In addition, the cross-section illustrates upper and lower regions at the coating surface and at the interface with the steel substrate that are completely free of Mg₂Si particles, with the Mg₂Si particles being confined to a
- 25 central band of the coating. This is advantageous for the reasons stated above.

The photomicrographs of the Figure illustrate clearly the benefits of the addition of Sr to an Al-Zn-Si-30 Mg coating alloy.

The laboratory experiments found that the microstructure shown in the right hand side of the Figure were formed with Sr additions in the range of 250-3000 ppm.

The applicant has also carried out line trials on 55%Al-Zn-1.5%Si-2.0%Mg alloy composition (not containing Sr) coated on steel substrates.

The purpose of these trials was to investigate the impact of cooling rates and coating masses on mottling in the surface of the coatings.

The trials covered a range of coating masses from 10 60 to 100 grams per square metre surface per side of strip, with cooling rates up to 90°C/sec.

The applicant found two factors that affected the coating microstructure, particularly the distribution of 15 Mg₂Si particles in the coatings, in the trials.

The first factor is the effect of the cooling rate of the strip exiting the coating bath before completing the coating solidification. The applicant found that controlling the cooling rate makes it possible

to avoid mottling.

By way of example, the applicant found that for a AZ150 class coating (or 75 grams of coating per square 25 metre surface per side of strip - refer to Australia Standard AS1397-2001), if the cooling rate is greater than 80°C/sec, Mg₂Si particles formed on the surface of the coating. In particular, when the cooling rate was greater than 100°C/sec, mottling occurred.

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The applicant also found that for the same coating it is not desirable that the cooling rate be too low, particularly below 11°C/sec, as in this case the coating develops a defective "bamboo" structure, whereby 35 the zinc-rich phases forms a vertically straight corrosion path from the coating surface to the steel interface, which compromises the corrosion performance of the coating.

Therefore, for a AZ150 class coating, under the 5 experimental conditions tested, the cooling rate should be controlled to be in a range of 11-80°C/sec to avoid mottling on the surface.

On the other hand, the applicant also found that 10 for a AZ200 class coating, if the cooling rate was greater than $50^{\circ}C/sec$, Mg₂Si particles formed on the surface of the coating and mottling occurred.

Therefore, for a AZ200 class coating, under the 15 experimental conditions tested, a cooling rate in a range of 11-50°C/sec is desirable.

The second important factor found by the applicant is the uniformness of coating thickness across 20 the strip surface.

The applicant found that the coating on the strip surface normally had thickness variations that are (a) long range (across the entire strip width, measured by the 25 "weight-strip-weight" method on a 50mm diameter disc) and (b) short range (across every 25 mm length in the strip width direction, measured in the cross-section of the coating under a microscope with 500x magnification). In a production situation, the long range thickness variation

- 30 is normally regulated to meet the minimum coating mass requirements as defined in relevant national standards. In a production situation, as far is the applicant is aware, there is no regulation for short range thickness variation, as long as the minimum coating mass
- 35 requirements as defined in relevant national standards are met.

However, the applicant found that short range coating thickness variations could be very high, and special operational measures had to be applied to keep the variations under control. It was not uncommon in the sexperimental work for the coating thickness to change by a factor of two or more over a distance as short as 5 mm, even when the product perfectly met the minimum coating mass requirements as defined in relevant national standards. This short range coating thickness variation had a pronounced impact on the Mg₂Si particles in the surface of coatings.

By way of example, the applicant found that for a AZ150 class coating even in the desirable cooling rate 15 ranges as described above, if the short range coating thickness variation was greater than 40% above the nominal coating thickness within a distance of 5 mm across the strip surface, Mg₂Si particles formed on the surface of the coating and thereby increased the risk of mottling. 20

Therefore, under the experimental conditions tested, the short range coating thickness variation should be controlled to no greater than 40% above the nominal coating thickness within a distance of 5mm across the strip surface to avoid mottling.

The research work carried out by the applicant on the solidification of Al-Zn-Si-Mg coatings, which is extensive and is described in part above, has helped the 30 applicant to develop an understanding of the formation of the Mg₂Si phase in a coating and the factors affecting its distribution in the coating. Whilst the applicant does not wish to be bound by the following discussion, this understanding is as set out below.

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When an Al-Zn-Si-Mg alloy coating is cooled to a temperature in the vicinity of 560°C, the α -Al phase is

the first phase to nucleate. The α -Al phase then grows into a dendritic form. As the α -Al phase grows, Mg and Si, along with other solute elements, are rejected into the molten liquid phase and thus the remaining molten liquid in the interdendritic regions is enriched in Mg and Si.

When the enrichment of Mg and Si in the interdendritic regions reaches a certain level, the Mg₂Si phase starts to form, which also corresponds to a 10 temperature around 465°C. For simplification, it will be assumed that an interdendritic region near the outer surface of the coating is region A and another interdendritic region near the quaternary intermetallic alloy layer at the steel strip surface is region B. It 15 will also be assumed that the level of enrichment in Mg and Si is the same in region A as in region B.

At or below 465°C, the Mg₂Si phase has the same tendency to nucleate in region A as in region B. However, 20 the principles of physical metallurgy teach us that a new phase will preferably nucleate at a site whereupon the resultant system free energy is the minimum. The Mg₂Si phase would normally nucleate preferably on the quaternary intermetallic alloy layer in region B provided the coating

- 25 bath does not contain Sr (the role of Sr with Srcontaining coatings is discussed below). The applicant believes that this is in accordance with the principles stated above, in that there is a certain similarity in crystal lattice structure between the quaternary
- 30 intermetallic alloy phase and the Mg₂Si phase, which favours the nucleation of Mg₂Si phase by minimizing any increase in system free energy. In comparison, for the Mg₂Si phase to nucleate on the surface oxide of the coating in region A, the increase in system free energy would have
- 35 been greater.

- 14 -

Upon nucleation in region B, the Mg₂Si phase grows upwardly, along the molten liquid channels in the interdendritic regions, towards region A. At the growth front of the Mg_2Si phase (region C), the molten liquid phase becomes depleted in Mg and Si (depending on the 5 partition coefficients of Mg and Si between the liquid phase and the Mg₂Si phase), compared with that in region A. Thus a diffusion couple forms between region A and region C. In other words, Mg and Si in the molten liquid phase will diffuse from region A to region C. Note that the 10 growth of the α -Al phase in region A means that region A is always enriched in Mg and Si and the tendency for the Mg₂Si phase to nucleate in region A always exists because the liquid phase is "undercooled" with regard to the Mg_2Si 15 phase.

Whether the Mg₂Si phase is to nucleate in region A, or Mg and Si are to keep diffusing from region A to region C, will depend on the level of Mg and Si enrichment in region A, relevant to the local temperature, which in turn depends on the balance between the amount of Mg and Si being rejected into that region by the α-Al growth and the amount of Mg and Si being moved away from that region by the diffusion. The time available for the diffusion is also limited, as the Mg₂Si nucleation/growth process has to be completed at a temperature around 380°C, before the L→Al-Zn eutectic reaction takes place, wherein L depicts the molten liquid phase.

30 The applicant has found that controlling the balance between the time available for diffusion and the diffusion distance for Mg and Si can control the subsequent nucleation or growth of the Mg₂Si phase or the final distribution of the Mg₂Si phase in the coating 35 thickness direction. In particular, the applicant has found that for a set coating thickness, the cooling rate should be regulated to a particular range, and more particualrly not to exceed a threshhold temperature, to avoid the risk for 5 the Mg₂Si phase to nucleate in region A. This is because for a set coating thickness (or a relatively constant diffusion distance between regions A and C), a higher cooling rate will drive the α -Al phase to grow faster, resulting in more Mg and Si being rejected into the liquid phase in region A and a greater enrichment of Mg and Si, or a higher risk for the Mg₂Si phase to nucleate, in region A (which is undesirable).

On the other hand, for a set cooling rate, a 15 thicker coating (or a thicker local coating region) will increase the diffusion distance between region A and region C, resulting in a smaller amount of Mg and Si being able to move from region A to region C by the diffusion within a set time and in turn a greater enrichment of Mg 20 and Si, or a higher risk for the Mg₂Si phase to nucleate, in region A (which is undesirable).

Practically, the applicant has found that, to achieve the distribution of Mg₂Si particles of the present invention, i.e. to avoid mottling defect on the surface of a coated strip, the cooling rate for coated strip exiting the coating bath has to be in a range of 11-80°C/sec for coating masses up to 75 grams per square metre of strip surface per side and in a range 11-50°C/sec for coating 30 masses of 75-100 grams per square metre of strip surface per side. The short range coating thickness variation also has to be controlled to be no greater than 40% above the nominal coating thickness within a distance of 5 mm

across the strip surface to achieve the distribution of Mg₂Si particles of the present invention.

The applicant has also found that, when Sr is present in a coating bath, the above described kinetics of Mg₂Si nucleation can be significantly influenced. At certain Sr concentration levels, Sr strongly segregates into the quaternary alloy layer (i.e. changes the chemistry of the quaternary alloy phase). Sr also changes the characteristics of surface oxidation of the molten coating, resulting in a thinner surface oxide on the coating surface. Such changes alter significantly the preferential nucleation sites for the Mg₂Si phase and, as a

- result, the distribution pattern of the Mg₂Si phase in the coating thickness direction. In particular, the applicant has found that, Sr at concentrations 250-3000ppm in the coating bath makes it virtually impossible for the Mg2Si
- 15 phase to nucleate on the quaternary alloy layer or on the surface oxide, presumably due to the very high level of increase in system free energy would otherwise be generated. Instead, the Mg₂Si phase can only nucleate at the central region of the coating in the thickness
- 20 direction, resulting in a coating structure that is substantially free of Mg₂Si at both the coating outer surface region and the region near the steel surface. Therefore, Sr additions in the range 250-3000ppm are proposed as one of the effective means to achieve a
- 25 desired distribution of Mg₂Si particles in a coating.

Many modifications may be made to the present invention as described above without departing from the spirit and scope of the invention.

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In this context, whilst the above description of the present invention focuses on (a) the addition of Sr to Al-Zn-Si-Mg coating alloys, (b) cooling rates (for a given coating mass) and (c) control of short range coating thickness variation as means for achieving a desired distribution of Mg_2Si particles in coatings, i.e. at least substantially no Mg_2Si particles in the surface of a coating, the present invention is not so limited and extends to the use of any suitable means to achieve the desired distribution of Mg_2Si particles in the coating.

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5732380_1 (GHMatters) P74467.AU.1 KRYSTALM 3/09/14

FEDERAL COURT OF AUSTRALIA

BlueScope Steel Limited v Dongkuk Steel Mill Co., Ltd (No 3) [2020] FCA 113

File number:	VID 1429 of 2016	
Judge:	BEACH J	
Date of judgment:	11 February 2020	
Catchwords:	PATENTS – costs – apportionment of costs as between issues – consequential orders	
Cases cited:	BlueScope Steel Limited v Dongkuk Steel Mill Co., Ltd (No 2) [2019] FCA 2117 Les Laboratories Servier v Apotex Pty Ltd (2016) 247 FCR 61	
Date of hearing:	On the papers	
Registry:	Victoria	
Division:	General Division	
National Practice Area:	Intellectual Property	
Sub-area:	Patents and associated Statutes	
Category:	Catchwords	
Number of paragraphs:	25	
Counsel for the Applicant and Cross-Respondent:	Mr BN Caine QC and Ms C Cunliffe	
Solicitor for the Applicant and Cross-Respondent:	King & Wood Mallesons	
Counsel for the Respondent and Cross-Claimant:	Mr A Ryan SC and Mr JS Cooke	
Solicitor for the Respondent and Cross-Claimant:	Bird & Bird	

ORDERS

VID 1429 of 2016

BETWEEN:	BLUESCOPE STEEL LIMITED ACN 000 011 058 Applicant
AND:	DONGKUK STEEL MILL CO., LTD Defendant
AND BETWEEN:	DONGKUK STEEL MILL CO., LTD Cross-Claimant
AND:	BLUESCOPE STEEL LIMITED (ACN 000 011 058) Cross-Respondent
JUDGE:	BEACH J
DATE OF ORDER:	11 FEBRUARY 2020

THE COURT ORDERS THAT:

- Each of claims 1, 3, 4, 5, 6, 7, 8, 9, 11, 12, 13, 14, and 15 of Australian Patent number 2009225257 titled "Metal-coated steel strip" (257 Patent) is and at all material times has been invalid.
- Each of claims 12, 22, 23, 24 and 25 of Australian Patent number 2009225258 titled "Metal-coated steel strip" (258 Patent) is and at all material times has been invalid.
- 3. It be certified that the validity of each of claims 1, 2, 5, 6, 11, 17, 18, 19, 20 and 21 of the 258 Patent was questioned in this proceeding.
- 4. Each of claims 1, 3, 4, 5, 6, 7, 8, 9, 11, 12, 13, 14 and 15 of the 257 Patent be revoked.
- 5. Each of claims 12, 22, 23, 24 and 25 of the 258 Patent be revoked.
- 6. The claim and the cross-claim otherwise be dismissed.
- The interlocutory application filed on 13 April 2017 seeking leave to amend the 257 Patent (Amendment Application) be dismissed.
- Any application by the Applicant/Cross-Respondent for leave to appeal against Order
 7 be heard and determined by a Full Court concurrently with, or immediately before,
 the hearing of any appeal.

- 9. Orders 4 and 5 be stayed:
 - (a) initially for a period of 28 days from the date on which these orders are pronounced; and
 - (b) if any appeal is lodged within that period, until the determination of that appeal and any further appeal therefrom, or further order.
- 10. Subject to Order 11 below, the Applicant pay two-thirds of the Respondent's costs of and incidental to the proceeding including the Amendment Application on a party-party basis.
- 11. Order 10 does not displace any costs order previously made in the proceeding.

THE COURT NOTES THAT:

- 12. The Applicant/ Cross-Respondent undertakes:
 - (a) to prosecute any appeal expeditiously;
 - (b) during the operation of the stay in Order 9, not to assert infringement of any of the claims referred to in Orders 4 and 5 against any third party; and
 - (c) forthwith to provide a copy of these Orders to the Commissioner of Patents.

Note: Entry of orders is dealt with in Rule 39.32 of the Federal Court Rules 2011.

CLAIMS

 A hot-dip coating method for forming a coating of a corrosion-resistant Al-Zn-Si-Mg alloy on a steel
 strip comprising passing the steel strip through a hot dip coating bath that contains Al, Zn, Si, and Mg and optionally other elements and forming an alloy coating on the strip with a variation in thickness of the coating of no more than 40% in any given 5 mm diameter section so
 that the distribution of Mg₂Si particles in the coating microstructure is such that there is only a small proportion of Mg₂Si particles or substantially no Mg₂Si particles in the surface of the coating.

- 15 2. The method defined in claim 1 comprising forming the alloy coating so that there are no Mg₂Si particles in the surface of the coating.
- 3. The method defined in any one of the preceding 20 claims wherein the small proportion of Mg₂Si particles in the surface of the coating is no more than 10wt.% of the Mg₂Si particles.
- 4. The method defined in any one of the preceding 25 claims wherein the coating thickness variation is no more than 30% in any given 5 mm diameter section of the coating.
- 5. The method defined in any one of the preceding 30 claims wherein, for a coating thickness of 22µm, the maximum thickness in any region of the coating greater than 1mm in diameter is 27µm.
- 6. The method defined in any one of the preceding 35 claims comprising selecting the cooling rate during solidification of coated strip exiting the coating bath to be less than a threshold cooling rate.

The method defined in any one of the preceding claims comprising selecting the cooling rate for coated strip exiting the coating bath to be less than 80°C/sec
 for coating masses up to 75 grams per square metre of strip surface per side.

8. The method defined any one of claims 1 to 6 comprising selecting the cooling rate for coated strip
 10 exiting the coating bath to be less than 50°C/sec for coating masses of 75-100 grams per square metre of strip surface per side.

9. The method defined in any one of the preceding 15 claims wherein the coating comprises the following ranges in % by weight of the elements aluminium, zinc, silicon, and magnesium:

	Magnesium	0.3 to 10 %
20	Silicon:	0.3 to 3%
	Zinc:	40 to 60 %
	Aluminium:	40 to 60 %

10. The method defined in any one of the preceding claims wherein the coating contains more than 250 ppm Sr. 25

11. The method defined in any one of the preceding claims wherein the coating contains less than 3000 ppm Sr.

12. The method defined in any one of the preceding
 30 claims comprising forming the coating to have a thickness of less than 30µm.

 The method defined in any one of the preceding claims comprising forming the coating to have a thickness
 of greater than 7µm. 14. A steel strip having a coating of a corrosionresistant Al-Zn-Si-Mg alloy formed by the method defined in any one of the preceding claims.

5 15. An Al-Zn-Si-Mg alloy coated strip that has Mg₂Si particles in the coating with the distribution of Mg₂Si particles being such that a surface of the coating has only a small proportion of Mg₂Si particles or is at least substantially free of any Mg₂Si particles formed by the method defined in any one of the preceding claims. 10



Figure 1 Sr additions in a 55%Al-Zn-1.5%Si-2.0%Mg coating eliminate the surface mottling defect and change the distribution pattern of the Mg₂Si phase in the coating thickness direction.

FIGURE 1