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(54) **ALUMINUM ALLOY CASTING MATERIAL, METHOD FOR PRODUCTION OF THE ALUMINUM ALLOY CASTING MATERIAL, ALUMINUM ALLOY MATERIAL, AND METHOD FOR PRODUCTION OF ALUMINUM ALLOY MATERIAL**

ALUMINIUMLEGIERUNGS-GUSSMATERIAL, VERFAHREN ZUR HERSTELLUNG DES
ALUMINIUMLEGIERUNGS-GUSSMATERIALS, ALUMINIUMLEGIERUNGSMATERIAL UND
VERFAHREN ZUR HERSTELLUNG DES ALUMINIUMLEGIERUNGSMATERIALS

MATÉRIAU DE MOULAGE EN ALLIAGE D'ALUMINIUM, PROCÉDÉ DE PRODUCTION DU
MATÉRIAU DE MOULAGE EN ALLIAGE D'ALUMINIUM, MATÉRIAU EN ALLIAGE D'ALUMINIUM ET
PROCÉDÉ DE PRODUCTION DE MATÉRIAU EN ALLIAGE D'ALUMINIUM

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GB-A- 1 475 587

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Description

Technical Field

[0001] The present invention relates to a casted aluminum alloy obtained by casting a molten metal alloy into a plate-like shape and a method for producing the casted aluminum alloy as well as to an aluminum alloy material obtained by processing and/or heating the casted aluminum alloy and a method for producing the aluminum alloy material.

Background Art

[0002] A rolled aluminum alloy sheet has been produced by: semi-continuously casting of a molten alloy product adjusted to a predetermined composition into a rolling ingot; slab cutting; homogenization step; surface cutting step; heating; and hot rolling. Cold rolling is performed when so required after the hot rolling. In such a rolled aluminum alloy sheet production process, a predetermined shape is achieved while melting coagulated structures, and adjustment for obtaining a homogenous and fine structure is performed. Also, in the rolling step, quality control (e.g. heat treatment) that is varied depending on the alloy type has been performed. Since there are the various rolled aluminum alloy sheet production steps as described above, there has been a limit in reduction in energy consumption and reduction in cost.

[0003] In recent years, a continuous casting method for aluminum alloy has been studied. This method is a method for continuously and directly casting a sheet material having a predetermined thickness from a molten metal of an aluminum alloy. In the continuous casting method, it is possible to continuously cast a sheet material having a thickness of 10 mm or less, for example, which is thinner than a slab. Therefore, a cooling rate is higher than that of the conventional ingot continuous casting, thereby obtaining a finer cast structure. Also, due to the high cooling rate, an allowable amount of Fe which has ordinarily been treated as an impurity element is increased, and recyclability of aluminum alloy is improved. Further, since it is possible to largely reduce the number of production steps, cost reduction can be achieved.

[0004] A 5000-system (Al-Mg) aluminum alloy has been mainly used as the rolled aluminum alloy sheet for an automobile outer panel, for example. As other examples, use of an excessive Si type 6016 alloy or 6022 alloy (Al-Mg-Si alloy) having a bake-hardening property has been studied in recent years. As used herein, "bake-hardening" means an aging phenomena utilizing heat in a baking step for an automobile.

[0005] For instance, in the Al-Mg-Si alloy, a material on which only a solution treatment has been performed (refinement: T4) is press-molded into a predetermined shape, and hardening is performed in a subsequent baking step, thereby obtaining the rolled aluminum alloy sheet for outer panels. Among aluminum alloys, the 6000-system alloy, such as the Al-Mg-Si alloy has strength and good corrosion resistance and has been used as an underbody material of an automobile or the like. In the 6000-system alloy, since such excellent properties are integrated with the above-described bake-hardening property and continuous casting and rolling, energy consumption for production is further reduced, thereby obtaining a highly functional and low cost material (see Patent Documents 1 to 3).

[0006] The level of strength of an aluminum alloy depends much on an alloy composition. Particularly, examples of an aluminum alloy capable of expressing high strength include a heat treated alloy which is precipitation strengthened by an aging treatment, and representative examples thereof include a 7000-system alloy (Al-Zn-Mg alloy) and a 2000-system alloy (Al-Cu alloy). The 6000-system alloy also belongs to this type but is inferior in strength properties as compared with other heat treated alloys. Meanwhile, a high strength 6000-system alloy to which Cu is added is under development.

[0007] However, like the 2000-system alloy and the 7000-system alloy, the Cu-added 6000-system alloy has a problem of reductions in processability and corrosion resistance despite the improvement in strength. Therefore, it has been difficult from the practical point of view to adapt the rolled aluminum alloy sheet made from such aluminum alloy to the outer plate, underbody, or the like of automobile to which corrosion resistance is required.

[0008] Also, the strength is improved by addition of an additive element in the rolled aluminum alloy sheet as described above, however, in an Al-Fe-Ni alloy or the like, for example, softening resistance is insufficient, and hardness (residual hardness) after casting, annealing, and long time heating, is subject to a large reduction as compared with the hardness after casting despite its heat resistance, i.e. its excellent strength at high temperatures. Therefore, it is impossible to perform high temperature aging on such aluminum alloy, and, consequently, since strength at room temperature is reduced after the high temperature heating despite the excellent strength under high temperature environments, it has been difficult to use such an aluminum alloy for parts to be used under high temperature environments.

[0009] As described above, as the rolled aluminum alloy sheets constituting automobile structural parts and the like, there has been a demand for the one that is capable of being molded into various desired shapes and excellent not only in strength, corrosion resistance, and the like, but also in softening resistance and the like.

[0010] It has been quite difficult to industrially produce a rolled aluminum alloy sheet that satisfies these property requirements using the aluminum alloy conventionally used.

Patent Document 1: JP 8-165538 Unexamined Patent Publication (Kokai)

Patent Document 2: JP 2004-156117 Unexamined Patent Publication (Kokai)

Patent Document 3: JP 2006-249550 Unexamined Patent Publication (Kokai)

[0011] EP 0 693 567 A2 discloses a high-strength, high-ductility cast aluminum alloy having a composition represented by the general formula $Al_aLn_bM_c$, wherein Ln is at least one metallic element selected from Y, La, Ce, Sm, Nd, Hf, Nb and Ta, M is at least one metallic element selected from V, Cr, Mn, Fe, Co, Ni, Cu, Zr, Ti, Mo, W, Ca, Li, Mg and Si, and a, b and c are, in terms of by weight, respectively $75\% \leq a \leq 95\%$, $0.5\% \leq b < 15\%$, and $0.5\% \leq c < 15\%$, said alloy having a structure comprising fine grains of α -Al, having an average grain diameter of not more than 10 μ m and an ultrafine compound of Al-lanthanide-base metal having an average grain diameter of not more than 1 μ m, said α -Al grains being surrounded by a network of said Al-lanthanide-base metal compound and forming a domain.

[0012] GB 1,475,587 discloses an aluminum alloy conductor having a minimum conductivity of 57 % IACS and consisting of from 0.99 to 2.50 wt% Fe, from 0.18 to 0.40 wt% Si and from 0.005 to 0.40 total wt% of trace elements selected from V, Cu, Mn, Mg, Zn, B, Ga, Ni, Zr, Cr, Be and Ti, and from 96 to 70 to less than 98.83 wt% Al.

Disclosure of the Invention

Problems to be Solved by the Invention

[0013] The present invention was accomplished in view of the above-described conventional problems, and an object thereof is to provide a casted aluminum alloy and an aluminum alloy material that can be produced at a low cost and are excellent in strength, formability, corrosion resistance, and softening resistance as well as methods for producing them.

Means for Solving the Problems

[0014] A first invention embodiment is a method for producing a casted aluminum alloy obtained by casting a molten metal of an aluminum alloy, characterized by comprising:

a melting step of preparing an aluminum alloy comprising: 0.8 to 5 mass% of Fe as a first component element, 0.15 to 1 mass% of Ti as a second component element, one or more kinds of third component elements selected from a third component element group consisting of Zr, Nb, Hf, Sc, and Y at a content of 0.05 to 2 mass% of each of the third components and in an amount that satisfies $Fe > X > Ti$ when the content of the first component element Fe is Fe (mass%), the content of the second component element Ti is Ti (mass%), and a total content of the third component elements is X (mass%); optionally 0.05 to 2 mass% of Mg as a fourth component element, optionally 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr and Co, and optionally V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%, and a residual part comprising Al and inevitable impurities, and melting the aluminum alloy at a temperature that is higher by 20°C or more than a liquidus temperature determined by the composition of the aluminum alloy to obtain a molten metal; and

a casting step of casting the molten metal into a casting mold and then cooling the molten metal to a temperature that is lower by at least 10°C than a solidus temperature determined by the composition of the aluminum alloy at a cooling rate of 150°C/sec. or more and less than 10000°C/sec.

[0015] The production method of the first invention embodiment includes the melting step and the casting step.

[0016] In the melting step and the casting step, the molten metal is produced by melting the aluminum alloy of the specific composition, and the casted aluminum alloy is produced by casting the molten metal while cooling the molten metal. Therefore, in the first invention, it is possible to directly cast the molten metal into a plate-like shape or the like, and it is possible to omit a step of producing a slab (ingot) and the like. Consequently, the number of steps can be reduced, and the casted aluminum alloy can be produced at a low cost.

[0017] In the method for producing of the present invention, the casting is performed by using the aluminum alloy having the specific composition having the first to third component elements contained in the specific amounts and the residual part comprising Al and inevitable impurities while cooling the molten metal of the aluminum alloy at the specific cooling rate. Therefore, the casted aluminum alloy that has excellent strength, formability, corrosion resistance, and softening resistance can be obtained even when the direct casting from the molten metal to the plate-like shape, for example, is performed.

[0018] The aluminum alloy within the specific composition range exhibits an excellent casting property without addition of Si. Therefore, the casting property is improved while avoiding deterioration in material properties caused by the Si

addition. Also, the aluminum alloy exhibits excellent heat resistance without addition of Ni or Mn.

[0019] Further, in the method for producing of the present invention, the casting is performed while cooling the molten metal of the aluminum alloy of the specific composition at the specific cooling rate. Therefore, the casted aluminum alloy exhibits the excellent strength and the excellent softening resistance, and hardness thereof at a room temperature is hardly reduced after being exposed to a high temperature environment which is 1/2 or more of the solidus temperature determined by the composition of the aluminum alloy. Therefore, the casted aluminum alloy can undergo aging at a high temperature, such as 200°C or more, and the strength thereof is improved further. Consequently, even when hot rolling, annealing, and the like are performed on the casted aluminum alloy in subsequent steps, the strength of the casted aluminum alloy is not reduced, or, rather, it is improved. Reasons for such strength are considered to be as follows.

[0020] In the case of adding Fe to an aluminum alloy as in the present invention, there are formed, as a metal structure, an α phase that is formed of an Al matrix and a layered phase that is formed of an eutectic structure of an Al-Fe compound and the Al matrix in such a manner as to surround the α phase. When the second component element Ti and the third component element are added by the specific amounts, it is possible to dissolve the second component element and the third component element into Al, thereby enabling formation of an Al matrix formed of an supersaturated solid solution. This makes it possible to precipitate a stable compound (intermetallic compound) phase formed of Al, Ti (the second component element), and the third component element in the Al matrix when heat energy or strain energy is applied. For this reason, the softening resistance is improved to enable the improvement in strength after the above-described processing and heating, and the like.

[0021] Also, in the present invention, it is possible to prevent a reduction in strength that can be caused when returned to a room temperature after a use under a high temperature environment for a long time, thereby making it possible to maintain the strength higher than the strength after the casting. Further, by adjusting the alloy composition and the cooling rate within the ranges of the present invention, the strength after the use under high temperature environment for a long time is not reduced but rather the strength can be further improved.

[0022] Consequently, in the first invention embodiment, it is possible to produce the casted aluminum alloy that is suitable for automobile structural parts and the like.

[0023] Also, in the method for producing of the present invention, the casting is performed at the high cooling rate of 150°C/sec. or more. Therefore, allowable amounts of impurity elements can be increased, thereby improving recyclability.

[0024] As described above, according to the first invention embodiment, it is possible to provide the method for producing the casted aluminum alloy that can be produced at low cost and that is excellent in strength, formability, corrosion resistance, and softening resistance.

[0025] According to a second invention embodiment, the method for producing an aluminum alloy material further comprises a hot rolling step of reducing a thickness of the casted aluminum alloy obtained by the production method of the first invention by 30% or more by performing hot rolling on the casted aluminum alloy at a temperature of 200°C or more.

[0026] According to a third invention embodiment, the method for producing an aluminum alloy material comprises a cold rolling-heating step of performing cold rolling on the casted aluminum alloy obtained by the production method of the first invention thereby to reduce a thickness of the casted aluminum alloy by 30% or more, and subsequently heating at a temperature of 1/2 or more of a melting point of the aluminum alloy and 550°C or less.

[0027] According to a fourth invention embodiment, the method for producing an aluminum alloy material comprises a heat treatment step of heating the casted aluminum alloy obtained by the production method of the first invention at a temperature of 400°C or more for 0.5 to 3 hours.

[0028] In each of the second to fourth invention embodiments, the hot rolling step, the cold rolling-heating step, or the heat treatment step is performed on the casted aluminum alloy obtained by the production method of the first invention. Therefore, it is possible to form a precipitate of an intermetallic compound formed of Al, the second component element, and the third component element in the aluminum alloy material to be obtained. The precipitate is considered to form a stable phase or metastable phase in the metal structure. The aluminum alloy material having such metal structure exhibits more excellent strength as compared with the casted aluminum alloy. Also, it is possible to produce other effects same as those of the first invention.

[0029] A fifth invention embodiment is a casted aluminum alloy characterized by comprising:

- 0.8 to 5 mass% of Fe as a first component element,
- 0.15 to 1 mass% of Ti as a second component element,
- one or more kinds of third component elements selected from a third component element group consisting of Zr, Nb, Hf, Sc, and Y at a content of 0.05 to 2 mass% of each of the third components and in an amount that satisfies $Fe > X > Ti$ when the content of the first component element Fe is Fe (mass%), the content of the second component element Ti is Ti (mass%), and a total content of the third component elements is X (mass%); optionally 0.05 to 2 mass% of Mg as a fourth component element, optionally 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr and Co, and optionally V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%, and

a residual part comprising Al and inevitable impurities,
 wherein the casted aluminum alloy has a metal structure including an α phase that is formed of an Al matrix and a
 layered phase that is formed of an eutectic structure of the Al matrix and an Al-Fe compound in such a manner as
 to surround the α phase;

the Al matrix is formed of a supersaturated solid solution of Al in which the second component element and the third
 component element are dissolved; and

an area ratio of a crystallize formed of an intermetallic compound of Al, the second component element, and the
 third component element and having a particle diameter of 5 μm or more is less than 5% in an arbitrary section of
 the casted aluminum alloy.

[0030] The casted aluminum alloy has the specific composition and the metal structure including the α phase that is
 formed of an Al matrix and the layered phase that is formed of a eutectic structure of the Al matrix and an Al-Fe compound
 in such a manner as to surround the α phase. The Al matrix is formed of a supersaturated solid solution of Al in which
 the second component element and the third component element are dissolved, and an area ratio of a crystallize
 formed of a compound of Al, the second component element, and the third component element and having a particle
 diameter of 5 μm or more is less than 5% in an arbitrary section of the casted aluminum alloy.

[0031] Like the casted aluminum alloy obtained by the production method of the first invention embodiment, it is
 possible to produce such a casted aluminum alloy at a low cost, and the casted aluminum alloy is excellent in strength,
 formability, corrosion resistance, and softening resistance.

[0032] In general, in an alloy having a similar composition to that of the casted aluminum alloy of the fifth invention
 embodiment, a crystallize formed of an intermetallic compound of Al, the second component element, and the third
 component element and having a particle diameter of 5 μm or more is easily generated during casting. Such crystallize
 generated in a large amount may cause a reduction in strength when hot rolling, annealing, or the like is performed.

[0033] In the casted aluminum alloy of the fifth invention embodiment, the area ratio of the crystallize having the
 particle diameter of 5 μm or more is less than 5% in an arbitrary section of the casted aluminum alloy. In other words,
 a content of the crystallize formed of an intermetallic compound of Al, the second component element, and the third
 component element is remarkably small, and the second component element and the third component element are
 dissolved into the Al matrix in the α phase. Therefore, even when the hot rolling, annealing, and the like are performed
 on the casted aluminum alloy, the strength of the casted aluminum alloy is not reduced but rather the strength is improved.

[0034] The casted aluminum alloy of the fifth invention embodiment is obtained by the production method of the first
 invention embodiment, for example. In the production method of the first invention embodiment, the aluminum alloy
 having the specific composition is cooled to the specific temperature at the specific cooling rate as described above.
 Therefore, the crystallize is suppressed from being generated in the α phase, thereby making it possible to maintain
 the area ratio of the crystallize to less than 5% as described above.

[0035] A sixth invention embodiment is an aluminum alloy material characterized by comprising:

0.8 to 5 mass% of Fe as a first component element,

0.15 to 1 mass% of Ti as a second component element,

one or more kinds of third component elements selected from a third component element group consisting of Zr,
 Nb, Hf, Sc, and Y at a content of 0.05 to 2 mass% of each of the third components and in an amount that satisfies
 $\text{Fe} > \text{X} > \text{Ti}$ when the content of the first component element Fe is Fe (mass%), the content of the second component
 element Ti is Ti (mass%), and a total content of the third component elements is X (mass%); optionally 0.05 to 2
 mass% of Mg as a fourth component element, optionally 0.05 to 1 mass% of at least one fifth component element
 selected from a fifth component element group consisting of Cu, Cr and Co, and optionally V and/or Mo as a sixth
 component element in an amount of more than 0.05 mass% and less than 0.5 mass%, and

a residual part comprising Al and inevitable impurities,

wherein the aluminum alloy material has a metal structure including an α phase that is formed of an Al matrix and
 a layered phase that is formed of an eutectic structure of the Al matrix and an Al-Fe compound in such a manner
 as to surround the α phase;

the Al matrix is formed of Al and/or a supersaturated solid solution of Al in which the second component element
 and the third component element are dissolved; and

a precipitate formed of an intermetallic compound of Al, the second component element, and the third component
 element and having a particle diameter of 2 to 500 nm is dispersed in the Al matrix.

[0036] The aluminum alloy material of the sixth invention embodiment has the specific composition and the metal
 structure including the α phase that is formed of an Al matrix and the layered phase that is formed of an eutectic structure
 of an Al-Fe compound and the Al matrix in such a manner as to surround the α phase. In the Al matrix, a precipitate
 formed of an intermetallic compound of Al, the second component element, and the third component element and having

a particle diameter of 2 to 500 nm or less is dispersed. Such an aluminum alloy material is excellent in strength, formability, corrosion resistance, and softening resistance.

[0037] In other words, it is possible to produce the aluminum alloy material of the sixth invention embodiment by performing heating, rolling, and the like, for example, on the casted aluminum alloy obtained by the production method of the first invention embodiment and the casted aluminum alloy of the fifth invention embodiment. It is possible to precipitate the second component element and the third component element dissolved in the Al matrix of the casted aluminum alloy as the fine precipitate by the heating, rolling, and the like. As a result, the casted aluminum alloy exhibits the excellent strength, formability, corrosion resistance, and softening resistance.

[0038] More specifically, the aluminum alloy material of the sixth invention embodiment can be obtained by the methods for producing of the second to fourth invention embodiments.

Brief Description of the Drawings

[0039]

Fig. 1 is an explanatory diagram showing a melting step, a casting step, a post-treatment step, and a heating step as well as measurement timings of hardness HVR1 to 3 in Embodiment 1, wherein (a) is an explanatory illustration of a case of performing a hot rolling step as the post-treatment step, (b) is an explanatory illustration of a case of performing a cold rolling-heating step as the post-treatment step, and (c) is an explanatory illustration of a case of performing a heat treatment step as the post-treatment step.

Fig. 2 is an explanatory diagram showing a melting step, a solidifying step, a post-treatment step, and a heating step as well as measurement timings of hardness HVR1 to 3 in Embodiment 1, wherein (a) is an explanatory illustration of a case of performing a hot rolling step as the post-treatment step and (b) is an explanatory illustration of a case of performing a cold rolling-heating step as the post-treatment step.

Fig. 3 is an explanatory diagram showing (a) pattern 1, (b) pattern 2, and (c) pattern 3 of hardness behaviors in Embodiment 1.

Fig. 4 is an explanatory view showing an alloy structure (Example 11) before hot rolling in Embodiment 1.

Fig. 5 is an explanatory view showing an alloy structure (Example 11) after hot rolling in Embodiment 1.

Fig. 6 is an explanatory graph showing a relationship between a cooling rate and tensile strength of each of aluminum alloy compositions in Embodiment 1.

Fig. 7 is an explanatory view showing a state in which a precipitate is formed, which is an SEM photograph of an alloy structure (Example 11) after hot rolling.

Fig. 8 is an explanatory view showing an SEM photograph ($\times 1000$ magnification) of an alloy structure of a casted aluminum alloy (Example 11) before hot rolling in Embodiment 1.

Fig. 9 is an explanatory view showing an SEM photograph ($\times 5000$ magnification) of an alloy structure of a casted aluminum alloy (Example 11) before hot rolling in Embodiment 1.

Fig. 10 is an explanatory view showing an SEM photograph ($\times 1000$ magnification) of an alloy structure of a casted aluminum alloy (Comparative Example 22) before hot rolling in Embodiment 1.

Fig. 11 is an explanatory view showing an SEM photograph ($\times 5000$ magnification) of an alloy structure of the casted aluminum alloy (Comparative Example 22) before hot rolling in Embodiment 1.

Fig. 12 is an explanatory view showing a component analysis result of a crystallite of the casted aluminum alloy (Comparative Example 22) before hot rolling in Embodiment 1.

Fig. 13 is an explanatory view showing a TEM photograph of the alloy structure of the casted aluminum alloy (Example 11) before hot rolling in Embodiment 1.

Fig. 14 is an explanatory diagram showing an analysis result of EDX at point *1 in Fig. 13.

Fig. 15 is an explanatory diagram showing an analysis result of EDX at point *2 in Fig. 13.

Fig. 16 is an explanatory diagram showing an analysis result of EDX at point *3 in Fig. 13.

Fig. 17 is an explanatory diagram showing an analysis result of EDX at point *4 in Fig. 13.

Fig. 18 is an explanatory view showing a TEM photograph of the alloy structure of the casted aluminum alloy (Example 11) after hot rolling in Embodiment 1.

Fig. 19 is an explanatory diagram showing an analysis result of EDX at point *1 in Fig. 18.

Fig. 20 is an explanatory diagram showing an analysis result of EDX at point *2 in Fig. 18.

Fig. 21 is an explanatory diagram showing an analysis result of EDX at point *3 in Fig. 18.

Fig. 22 is an explanatory diagram showing an analysis result of EDX at point *4 in Fig. 18.

Fig. 23 is an explanatory graph showing a relationship between an annealing temperature and residual hardness in Embodiment 1.

Fig. 24 is an explanatory diagram showing (a) a front view and (b) a side view of a casted aluminum alloy in Embodiment 2.

Fig. 25 is an explanatory view showing (a) an explanatory illustration of an alloy structure of a thin plate part of a casted aluminum alloy, (b) an explanatory illustration of an alloy structure of a medium thickness plate part of the casted aluminum alloy, and (c) an explanatory illustration of an alloy structure of a large thickness plate part of the casted aluminum alloy.

Best Mode for Carrying out the Invention

[0040] Hereinafter, preferred mode of the present invention will be described.

[0041] In the production method of the first invention embodiment, the casted aluminum alloy is produced by performing the melting step and the casting step. The casted aluminum alloy is produced by a casting method of casting a molten metal of an aluminum alloy while supplying the molten metal to a casting mold.

[0042] It is preferable that the casting step is performed by continuous casting.

[0043] In this case, the casted aluminum alloy can be continuously and directly cast from the molten metal. In this case, it is possible to prominently exhibit the above-described effect of obtaining a casted aluminum alloy excellent in strength, formability, corrosion resistance, and softening resistance even when a direct casting is performed. Also, in this case, it becomes easy to perform the cooling at the specific cooling rate.

[0044] Alternatively, the casting step may be performed by a method other than the continuous casting, such as die casting.

[0045] In the melting step, used is an aluminum alloy containing the first component element, the second component element, and the third component element and having a residual part formed of aluminum and inevitable impurities.

[0046] The aluminum alloy contains 0.8 to 5 mass% of Fe as the first component element.

[0047] In the case of adding Fe to an aluminum alloy, strength and strength at a high temperature (heat resistance) are improved. From the viewpoint of metal structure, a layered phase is formed by an Al-Fe compound and Al.

[0048] In the case where Fe is less than 0.8 mass%, satisfactory strength is not achieved, which may lead to a reduction in strength under high temperature environment, i.e. in heat resistance. In the case where Fe exceeds 5 mass%, properties become subject to change greatly corresponding to the cooling rate, and a problem of difficulty in stable producing cast material having certain properties occurs. More specifically, in the case of performing rolling, for example, the casted aluminum alloy may become subject to crack. In this case, bulk crystallizate is easily formed during casting to raise a possibility of deterioration in processability and formability. The content of the first component element Fe may preferably be 2.0 to 4.0 mass%, more preferably 3.0 to 4.0 mass%.

[0049] 0.15 to 1 mass% of Ti is contained as the second component element.

[0050] In the case of adding the second component element Ti together with the third component element, it is possible to refine the alloy structure as well as to further improve strength properties because a supersaturated solid solution obtained during solidification from a dissolved state is precipitated in an aluminum parent phase in the case where hot rolling is performed or in the case where a heat treatment is performed after cold rolling.

[0051] In the case where Ti is less than 0.15 mass%, satisfactory heat resistance and softening resistance may not be achieved. In the case where Ti exceeds 1 mass%, a bulk Al-Ti crystallizate is easily formed during casting, and processability and formability may be deteriorated. The content of the second component element Ti may preferably be 0.3 to 0.9 mass%, more preferably be 0.7 to 0.8 mass%.

[0052] As the third component element, one or more kinds of elements selected from a third component element group consisting of Zr, Nb, Hf, Sc, and Y is/are contained, and the content of each of the elements is 0.05 to 2 mass%.

[0053] The third component element exhibits an effect of improving softening resistance when added together with the first component element Fe and the second component element Ti. Specifically, in the case where the first component element Fe is added to the aluminum alloy, the layered phase is formed by the Al-Fe compound and the Al matrix as described above. Further, when the second component element Ti and the third component element are added in the specific amounts, a stable compound (intermetallic compound) phase formed of Al, Ti, and the third component element is formed in an aluminum parent phase when heat energy or strain energy is applied, thereby making it possible to improve the strength properties and the softening resistance. Therefore, the strength is improved in the case where heating is performed after hot rolling or cold rolling. Likewise, the strength improvement effect can be achieved in the case of performing only a heat treatment without performing rolling.

[0054] In the case where the content of each of the third component elements is less than 0.05 mass%, the above-described effects that should be achieved by the addition of the third component element may not be satisfactorily achieved. In the case where at least one of the third component elements exceeds 2 mass%, a large crystallizate is easily generated when the cooling rate is not satisfactorily high, and processability and formability may be deteriorated. Therefore, production can be made difficult. The content of each of the third component element groups may preferably be 0.2 to 1.2 mass%, more preferably be 0.5 to 1.2 mass%.

[0055] The total content X (mass%) of the third component elements satisfies $Fe > X > Ti$ when the content of the first component element Fe is Fe (mass%) and the content of the second component element Ti is Ti (mass%) in the aluminum

alloy.

[0056] In the case of $X \geq \text{Fe}$, there is a possibility of a reduction in strength of the casted aluminum alloy and a reduction in softening resistance. In the case of $X \leq \text{Ti}$, softening resistance may be deteriorated. In the case of $\text{Fe} \leq \text{Ti}$, there is a possibility of a reduction in strength of the casted aluminum alloy and a reduction in softening resistance.

[0057] In the melting step, it is preferable to use the aluminum alloy at least containing 0.2 to 1.2 mass% of Zr among the third component element group.

[0058] In this case, it is possible to further improve the softening resistance while maintaining the excellent strength properties and formability.

[0059] In the case where the content of Zr is less than 0.2 mass%, the above-described effects that should be achieved by the addition of Zr may not be satisfactorily achieved. In the case where the content exceeds 1.2 mass%, there is a possibility of a remarkable increase in melting temperature when melting the aluminum alloy in the melting step. Since a special device will be required for melting, the production cost can be increased.

[0060] In the melting step, it is preferable to use the aluminum alloy further containing 0.05 to 2 mass% of Mg as a fourth component element.

[0061] In this case the strength of the casted aluminum alloy substantially can be further improved without impairing the formability. In the case where Mg is less than 0.05 mass%, a strength improvement effect that should be achieved by the addition of Mg is not satisfactorily achieved, and the significance of the Mg addition can be almost lost. In the case where Mg is added in an amount exceeding 2 mass%, processability of the casted aluminum alloy is deteriorated, which may cause generation of rolling crack during rolling, for example. Also, formability can be deteriorated. The content of the fourth component element Mg may preferably be 0.2 mass% to 1.5 mass%, more preferably be 0.3 mass% to 0.8 mass%.

[0062] In the melting step, it is preferable to use the aluminum alloy further comprising 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr, and Co.

[0063] In the case of containing Cu among the fifth component element group, the strength is improved substantially without impairing the processability of the casted aluminum alloy. Also, in the case of containing Cr and/or Co among the fifth component element group, an Al-(Fe, Cr) compound and/or an Al-(Fe, Co) compound is formed, thereby making it possible to improve elongation property, processability, and formability as compared with those achieved by dispersion of an Al-Fe compound alone. As a result, the strength of the casted aluminum alloy is improved substantially without impairing the processability, formability, and the like.

[0064] In the case where the fifth component element is less than 0.05 mass%, the above-described effect that should be achieved by the addition of the fifth component element may not be satisfactorily achieved. In the case where Cu among the fifth component element group is added in an amount exceeding 1 mass%, processability and formability may be deteriorated. Also, in this case, corrosion resistance may be deteriorated. Also, in the case where Cr and/or Co among the fifth component element group is added in an amount exceeding 1 mass%, formability may be deteriorated. The content of the fifth component element may more preferably be 0.1 to 0.7 mass%, yet more preferably be 0.1 to 0.5 mass%.

[0065] In the case of containing two or more kinds of the fifth component elements, a total amount thereof may preferably be within the range of 0.05 to 1 mass%.

[0066] In the melting step, it is preferable to use the aluminum alloy further containing V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%.

[0067] In this case, the strength is improved substantially without impairing the processability and formability of the casted aluminum alloy.

[0068] In the case where the sixth component element is 0.05 mass% or less, the effect that should be achieved by the addition of the sixth component element may not be satisfactorily achieved. In the case of adding 0.5 mass% or more, there is a possibility of a considerable increase in melting temperature. Also, a bulk crystallizate is easily formed, and processability and formability may be deteriorated. The content of the sixth component element may more preferably be 0.1 to 0.4 mass%, yet more preferably be 0.1 to 0.3 mass%. In the case of containing two or more kinds of the sixth component elements, a total amount thereof may preferably be within the range of from more than 0.05 mass% to less than 0.5 mass%.

[0069] In the aluminum alloy, it is preferable that a total amount of the fourth component element, the fifth component element, and the sixth component element is maintained to 3 mass% or less.

[0070] In the case where the total amount of the fourth to sixth component elements exceeds 3 mass%, the processability of the casted aluminum alloy is deteriorated, which may cause generation of rolling crack during rolling, for example. In this case, a crystallizate is easily generated during the casting step to raise a possibility of deterioration in formability.

[0071] In the present invention, a specific gravity of the aluminum alloy of the alloy composition having the particularly highly excellent strength, softening resistance, corrosion resistance, and formability is 2.7 g/cm^3 .

[0072] In the melting step, the aluminum alloy is molten at a temperature that is higher by 20°C than a liquidus temperature determined by the composition (liquidus temperature $+20^\circ\text{C}$ or more) to obtain the molten metal.

[0073] In the case where the melting temperature is a temperature that is less than the liquidus temperature +20°C, it is difficult to achieve satisfactory melt fluidity, and a porosity is formed inside the casted aluminum alloy after the casting, thereby raising a possibility of failing to obtain strong casted aluminum alloy.

[0074] Subsequently, in the casting step, the molten metal is cast into a plate-like shape while being cooled to a temperature that is lower by 10°C than a solidus temperature determined by the aluminum alloy composition, i.e. that is lower than the solidus temperature by at least -10°C, at a cooling rate of 150°C/sec. or more and less than 10000°C/sec. to obtain the casted aluminum alloy.

[0075] In this case, the casted aluminum alloy excellent in the properties, such as softening resistance as described above can be obtained, and the strength can be further improved by precipitation of a stable compound (intermetallic compound) phase formed of Al, Ti, and the third component element in the aluminum parent phase (the α phase) when heat energy or strain energy, for example, is applied. Also, since it is possible to suppress formation of a crystallization phase containing a bulk Al-Fe compound or other elements, reductions in ductility, toughness, and the like can be prevented. Therefore, the properties, such as softening resistance is improved without impairing the processability, formability, and the like.

[0076] In the case where the cooling rate in the casting step is less than 150°C/sec., a bulk crystallize is formed during a solidifying process to raise a possibility of deterioration in formability and reductions in strength properties and softening resistance. Also, since it is necessary to use a special device in order to realize a cooling rate exceeding 10000°C/sec., production cost can be increased. In the case of attaining the cooling rate exceeding 10000°C/sec., it is necessary to shape the casted aluminum alloy after the casting into the form of a ribbon or a powder. Therefore, when performing composition processing, such as rolling, a preliminary molding step is required in advance.

[0077] Therefore, by defining the cooling rate of 150°C/sec. or more and less than 10000°C/sec. as in the present invention, the high quality casted aluminum alloy can be produced at industrially realizable low cost. Also, with the cooling rate within the defined range of the present invention, an amorphous phase is substantially absent in a section of the casted aluminum alloy, and it is possible to obtain the casted aluminum alloy having high heat stability with which a change in properties, for example, which can be caused near a crystallization temperature hardly occurs.

[0078] In the casting step, in the case where the cooling at the cooling rate is not performed to the temperature that is -10°C from the solidus temperature, a cast material at a downstream is locally re-molten due to heat of the molten metal at an upstream during the continuous casting, which may cause a bulk crystallize to be generated. Therefore, there is a possibility that a metal structure of the casted aluminum alloy to be obtained becomes non-uniform.

[0079] It is sufficient when the cooling at the cooling rate (150°C/sec. or more and less than 10000°C/sec.) is performed until reaching to the temperature of -10°C from the solidus temperature, and, once reaching to the temperature, the cooling may be performed at a temperature deviated from the cooling rate of 150°C/sec. or more and less than 10000°C/sec., or may be cooled at the cooling rate within the range. The cooling at the cooling rate (150°C/sec. or more and less than 10000°C/sec.) may preferably be performed until reaching to a temperature that is -100°C from the solidus temperature.

[0080] It is preferable that a copper casting mold is used as the casting mold.

[0081] In this case, it is possible to relatively easily realize the range of the cooling rate of 150°C/sec. or more and less than 10000°C/sec. in the casting step.

[0082] In the casting step, it is preferable that the molten metal is continuously cast into a plate-like shape having a thickness of 0.3 to 10 mm.

[0083] It is possible to perform the casting into plate-like shape by using a casting apparatus of a single roll type, a twin roll type, a block type, a belt type, a wheel type, and the like as described later in this specification.

[0084] In the case where the thickness is less than 0.3 mm, it is difficult to pour the molten metal between rolls, blocks, or belts, or to perform gap control, or other operations and the production of the casted aluminum alloy can be made difficult. In the case where the thickness exceeds 10 mm, it is difficult to ensure the above-described cooling rate of 150°C/sec. or more. Also, a fluctuation of the cooling rate can occur, thereby making it difficult to obtain the casted aluminum alloy having uniform characteristic.

[0085] In the casting step, it is preferable that the molten metal is continuously cast by using a single roll type, a twin roll type, a block type, a belt type, or a wheel type.

[0086] The continuous casting method of single roll type is a method for obtaining a casted aluminum alloy in the form of a plate by continuously supplying the molten metal of the aluminum alloy to a rotational single roll made from copper, for example, and performing rapid solidification. Also, in the continuous casting method of the twin roll type, rotational rolls made from copper are disposed in a pair, and a cooling rate can be controlled by arbitrarily adjusting a gap between the both rolls. The casted aluminum alloy in the form of a plate is obtained by rapid solidification in the same manner as in the single roll method. The block type continuous casting method is a method for supplying the molten metal between two movable block-like cooling members and continuously casting the molten metal into a plate-like shape while cooling and solidifying between the blocks. The belt type continuous casting method is a method for supplying a molten metal between two movable belt-like cooling members and continuously casting the molten metal into a plate-like shape while

cooling and solidifying between the belts. The wheel type continuous casting method is a method for pouring a molten metal into a part of a groove formed on an outer peripheral surface of a rotational wheel to allow the molten metal to pass through between the groove and a holding member (wheel) and casting the molten metal into a plate-like shape by continuously withdrawing the molten metal while solidifying the molten metal in a movable casting mold.

[0087] In these methods, the range of the cooling rate of 150°C/sec. or more and less than 10000°C/sec. can be realized as well as to produce the casted aluminum alloy with excellent productivity.

[0088] In the casting step, it is possible that the molten metal is cast into the plate-like shape, and at least a surface layer of the molten metal is cooled and solidified in the casting mold, and then cooling by water cooling.

[0089] In this case, it is possible to use a rotational roll having a small diameter when performing the casting by the single roll or the twin roll type continuous casting method, for example. In other words, a contact area and a contact time between the roll (casting mold) and the molten metal become insufficient when the rotational roll having small diameter is used, thereby making it difficult to cool the molten metal to the temperature of -10°C from the solidus temperature at the cooling rate. In contrast, when the water cooling is performed subsequently to the molten metal of which at least the surface layer is solidified in the casting mold, it is possible to much more reliably perform the cooling to the temperature of -10°C from the solidus temperature at the cooling rate even when the roll diameter of the rotational roll is small.

[0090] It is possible to perform various post-treatments for applying heat energy and/or strain energy to the casted aluminum alloy to be obtained after the casting step.

[0091] More specifically, as in the second invention, it is possible to perform a hot rolling step of reducing the thickness of the casted aluminum alloy by 30% or more by performing hot rolling processing at a temperature of 200°C or more on the casted aluminum alloy after the casting step.

[0092] In this case, not only the α phase and the layered phase formed of the eutectic structure of the Al-Fe compound and Al are formed in the aluminum alloy material, but also the stable compound (precipitate) phase formed of Al, Ti (the second component element), and the third component element is formed in the Al matrix by the heat energy and the strain energy as described above. Therefore, the aluminum alloy material can be obtained by further improving the strength of the casted aluminum alloy. The aluminum alloy material of the sixth invention embodiment can be obtained by performing the above-described hot rolling step.

[0093] In the case where the temperature for heating the casted aluminum alloy is less than 200°C, rolling crack and a large cracked edge may be generated. Such crack generation is prominent particularly in the case of using an aluminum alloy having a high solute concentration. Also, there is a possibility of failing to satisfactorily develop the characteristic of the aluminum alloy material that the strength is improved by heat energy. This may result in failing to satisfactorily improve the strength of the aluminum alloy material after the above-described hot rolling step.

[0094] In the hot rolling step, rolling processing for reducing 30% or more of the thickness of the casted aluminum alloy to be obtained after the casting step is performed. In the case where the reduction is less than 30%, the strain energy becomes insufficient, resulting in failing to satisfactorily improve the strength.

[0095] It is preferable that the hot rolling step is performed after casting the molten metal into a plate-like shape in the casting mold while cooling the molten product to a temperature of 200°C to 500°C and releasing from the casting mold in the casting step.

[0096] In this case, it is possible, without heating, to use in the hot rolling step the casted aluminum alloy as it is which is obtained after the casting step and has the temperature of 200°C to 500°C. Also, in the case where additional heating is required in the hot rolling step, the heating time can be shortened. Therefore, the number of steps and the production time can be reduced, thereby making it possible to achieve reduction in production cost.

[0097] In the case where the temperature of the casted aluminum alloy after the casting step is less than 200°C, it is necessary to heat the casted aluminum alloy again to the temperature of 200°C or more in the hot rolling step. In the case where the temperature of the casted aluminum alloy after the casting step exceeds 500°C, on the other hand, the Al-Fe compound in the layered phase becomes a bulk, thereby raising a possibility of a reduction in strength. Also, in this case, the service life of a roll can be shortened due to an increase in damage to the roll in the hot rolling step.

[0098] In the range of the aluminum alloy composition to be used in the present invention, the above-described "temperature which is lower than the solidus temperature by at least 10°C" never be under the above-described 500°C or lower. Therefore, even when the cooling to 500°C is performed in the casting step, the cooling to the "temperature which is lower than the solidus temperature by at least 10°C" is satisfactorily ensured. Also, when high temperature annealing at 450°C \times 1h is performed after the hot rolling, the softening resistance is no longer changed.

[0099] As in the third invention embodiment, it is possible to perform a cold rolling-heating step of reducing the thickness of the casted aluminum alloy by 30% or more by performing cold rolling on the casted aluminum alloy after the casting step and heating the casted aluminum alloy at a temperature from a temperature which is 1/2 or more of a melting point of the aluminum alloy to 550°C or less.

[0100] In this case, too, the aluminum alloy material can be obtained by further improving the strength of the casted aluminum alloy by the heat energy and the strain energy in the same manner as in the above-described hot rolling step.

[0101] In the case where the heating temperature of the casted aluminum alloy is less than 1/2 of the melting point of

the aluminum alloy, which may lead to failing to satisfactorily develop the characteristic of the casted aluminum alloy that the strength is improved by the heat energy. This may result in failing to satisfactorily improve the strength of the casted aluminum alloy after the cold rolling and heating. In the case where the heating temperature exceeds 550°C, there is a possibility of formation of a bulk compound and local melting of the composition. This may results in a reduction in properties, such as strength of the aluminum alloy material. In order to further improve the strength, the heating temperature in the cold rolling-heating step may preferably be 400°C to 500°C, more preferably be 400°C to 450°C.

[0102] It is possible to produce the aluminum alloy material of the sixth invention embodiment also by the above-described cold rolling-heating step.

[0103] As in the fourth invention embodiment, it is possible to perform a heat treatment step of heating the casted aluminum alloy after the casting step at a temperature of 400°C or more for 0.5 to 3 hours.

[0104] In this case, too, it is possible to precipitate a stable compound phase formed of Al, Ti (the second component element), and the third component element in the Al matrix by heat energy. Therefore, the aluminum alloy material can be obtained by further improving the strength of the casted aluminum alloy.

[0105] In the case where the heating temperature of the casted aluminum alloy is less than 400°C or where the heating time is less than 0.5 hour, it may be impossible to satisfactorily develop the characteristic of the casted aluminum alloy that the strength is improved by the heat energy. This may results in failing to satisfactorily improve the strength of the casted aluminum alloy after the heating. Meanwhile, the strength properties are not enhanced even when heating is performed for over 3 hours, and the advantage of long time heating is hardly achieved. In order to further improve the strength, the heating temperature in the heat treatment step may preferably be 400°C to 500°C, and the heating time may preferably be 1 to 2 hours.

[0106] It is possible to produce the aluminum alloy material of the sixth invention embodiment also by the above-described heat treatment step.

[0107] In the casted aluminum alloy produced by the first invention embodiment, an amorphous phase substantially does not exist on any part other than a surface.

[0108] Hereinafter, the fifth and sixth invention embodiments will be described.

[0109] Each of the casted aluminum alloy of the fifth invention embodiment and the aluminum alloy of the sixth invention embodiment contains 0.8 to 5 mass% of Fe as a first component element, 0.15 to 1 mass% of Ti as a second component element, one or more kinds of third component elements selected from a third component element group consisting of Zr, Nb, Hf, Sc, and Y at a content of 0.05 to 2 mass% of each of the third components and in an amount that satisfies $Fe > X > Ti$ when the content of the first component element Fe is Fe (mass%), the content of the second component element Ti is Ti (mass%), and a total content of the third component elements is X (mass%); optionally 0.05 to 2 mass% of Mg as a fourth component element, optionally 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr and Co, and optionally V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%, and a residual part containing Al and inevitable impurities. Significance of criticality of the contents of the component elements are the same as those of the first invention.

[0110] A metal structure of each of the casted aluminum alloy and the aluminum alloy material is formed of an α phase that is formed of an Al matrix and a layered phase that is formed of an eutectic structure of the Al matrix and an Al-Fe compound in such a manner as to surround the α phase.

[0111] In the fifth invention embodiment, the Al matrix is formed of a supersaturated solid solution of Al, and the second component element and the third component element are dissolved into the supersaturated solid solution.

[0112] In the sixth invention embodiment, the Al matrix is formed of Al and/or a supersaturated solid solution of Al in which the second component element and the third component element are dissolved. In the sixth invention embodiment, in the case where the dissolved second component element and the third component element are completely precipitated, the Al matrix is formed of Al.

[0113] The second component element and the third component element are not dissolved in the Al-Fe compound in the layered phase but dissolved in the α phase and/or the Al matrix in the layered structure.

[0114] In the fifth invention embodiment, an area ratio of a crystallizate formed of a compound of Al, the second component element, and the third component element and having a particle diameter of 5 μm or more is less than 5% in an arbitrary section of the casted aluminum alloy. It is preferable that an area ratio of a crystallizate having a particle diameter of 2.5 μm or more is less than 5%, and it is more preferable that the crystallizate substantially is not contained in the α phase.

[0115] When the area ratio of the crystallizate having the particle diameter of 5 μm or more is 5% or more, there is a possibility of failing to improve the strength properties and softening resistance when heat energy or strain energy is applied to the casted aluminum alloy by hot rolling or the like.

[0116] In the aluminum alloy material of the sixth invention embodiment, a precipitate formed of a compound of Al, the second component element, and the third component element and having a particle diameter of 2 to 500 nm is dispersed in the Al matrix. The precipitate is formed in the Al matrix of the α phase, for example. In the case where the particle diameter of the precipitate is less than 2 nm, satisfactory strength may not be achieved. In the case where the

particle diameter exceeds 500 nm, there is also a possibility of failing to achieve satisfactory strength.

[0117] It is possible to confirm existences of the precipitate in the sixth invention embodiment and the crystallize in the fifth invention embodiment by transmission electron microscopic observation, for example. Also, it is possible to investigate the size (particle diameter) of the crystallize and the precipitate on an arbitrary section of the casted aluminum alloy by microscopic observation. Each of the particle diameters of the crystallize and the precipitate is defined as a diameter (circle equivalent diameter) of a circle which is estimated to have an area identical to that of the crystallize or the precipitate in the arbitrary section of the casted aluminum alloy.

[0118] Also, an area proportion of the crystallize in the arbitrary section is measured by: polishing a measurement object surface of the casted aluminum alloy to a mirror surface; observing the measurement object surface by using a transmission electron microscope; and measuring an area ratio of the obtained image by using an image analysis processing device. The area ratio is a proportion of the area of the crystallize to the observed visual field area in the observed visual field plane. Note that the observed visual field area is at least 1 mm² or more.

[0119] The aluminum alloy material of the sixth invention embodiment can be obtained by applying heat energy or strain energy to the casted aluminum alloy obtainable from the production method of the first invention embodiment or the casted aluminum alloy of the fifth invention embodiment. Thus, it is possible to precipitate as the precipitate each of the second component element and the third component element dissolved into the Al matrix.

[0120] In the fifth and sixth invention embodiments, it is preferable that the casted aluminum alloy and the aluminum alloy material may at least contain 0.2 to 1.2 mass% of Zr among the third component element group.

[0121] In this case, the softening resistance can be further improved while maintaining the excellent strength properties and formability. Significance of criticality of the content of Zr is similar to that of the first invention.

[0122] It is preferable that the casted aluminum alloy and the aluminum alloy material contain 0.05 to 2 mass% of Mg as a fourth component element.

[0123] In this case, the strength of the casted aluminum alloy and the aluminum alloy material can be further improved substantially without impairing the formability. Significance of criticality of the content of Mg is similar to that of the first invention.

[0124] It is preferable that Mg as the fourth component element is dissolved at least into the Al matrix.

[0125] Also, it is preferable that Mg as the fourth component element forms an Al-Mg compound in the Al matrix.

[0126] In this case, the strength of the casted aluminum alloy and the aluminum alloy material can yet further be improved.

[0127] It is preferable that each of the casted aluminum alloy and the aluminum alloy material contain 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr, and Co. In the case of containing Cu among the fifth component element group, the strength is improved substantially without impairing the processability of the casted aluminum alloy and the aluminum alloy material. Also, in the case of containing Cr and/or Co among the fifth component element group, it is possible to improve the elongation property, the processability, and the formability of the casted aluminum alloy and the aluminum alloy material. Significance of criticality of the content of the fifth component element is similar to that of the first invention.

[0128] It is preferable that Cr and/or Co as the fifth component element is substituted with at least a part of the Al-Fe compound forming the layered phase to form an Al-(Fe, Cr) compound and/or an Al-(Fe, Co) compound.

[0129] In this case, it is possible to further improve the elongation property, the processability, and the formability of the casted aluminum alloy and the aluminum alloy material.

[0130] It is preferable that Cu as the fifth component element forms an Al-Cu compound in the Al matrix.

[0131] In this case, it is possible to further improve the strength substantially without impairing the processability of the casted aluminum alloy and the aluminum alloy material.

[0132] It is preferable that each of the casted aluminum alloy and the aluminum alloy material contain 0.05 to 2 mass% of Mg as a fourth component element and 0.05 to 1 mass% of Cu as a fifth component element, and that the fourth component element and the fifth component element form an Al-Cu-Mg compound in the Al matrix. In other words, in the case where Mg as the fourth component and Cu as the fifth component element are contained, it is preferable that an Al-Cu-Mg compound is formed in the Al matrix.

[0133] In this case, it is possible to yet further improve the strength substantially without impairing the processability of the casted aluminum alloy and the aluminum alloy material.

[0134] It is preferable that each of the casted aluminum alloy and the aluminum alloy material contain V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%.

[0135] In this case, the strength substantially is improved without impairing the processability and formability of the casted aluminum alloy and the aluminum alloy material. Significance of criticality of the content of the sixth component element is similar to that of the first invention.

[0136] It is preferable that V and/or Mo as the sixth component element forms a compound with Al (Al-V compound and/or Al-Mo compound) and a compound with Al, the second component element Ti, and the third component element X (Al-(V, X, Ti), Al-(Mo, X, Ti)) in the Al matrix.

[0137] In this case, it is possible to yet further improve the strength substantially without impairing the processability and formability of the casted aluminum alloy and the aluminum alloy material.

Examples

(Embodiment 1)

[0138] In the present embodiment, casted aluminum alloys (continuously cast materials) made from aluminum alloys of a plurality of types of compositions were prepared as shown in Table 1 to Table 4, and softening resistance and the like were investigated to clarify superiority of the casted aluminum alloys of the present invention.

[0139] Alloy compositions, specific gravities, and cooling rates of the casted aluminum alloys (Examples 1 to 48) within the scope of the present invention are shown in Table 1 and Table 2.

[0140] Also, for the purpose of comparison, casted aluminum alloys (Comparative Examples 1 to 20, and Comparative Examples 28 to 37) made from aluminum alloys that are out of the component range defined in the present invention and casted aluminum alloys (Comparative Examples 21 to 27) produced at cooling rates that are out of the range defined in the present invention were prepared. Alloy compositions and specific gravities of these alloys are shown in Table 3 and Table 4.

[0141] In the present embodiment, the cooling rate is determined by a speed of a molten metal of each of the aluminum alloys of the different compositions passing through a range of temperatures of $\pm 40^{\circ}\text{C}$ from the solidus temperature in the course of cooling the molten metal in the casting step.

[0142] In the present embodiment, each of the casted aluminum alloys (Examples 1 to 48, Comparative Examples 1 to 20, and Comparative Examples 28 to 37) was produced by continuous casting, and various post-treatments were performed for the evaluation of softening resistance as shown in Fig. 1.

[0143] More specifically, for the production of each of the casted aluminum alloys, a melting step S1 for forming a molten metal by melting the aluminum alloy at a temperature (melting temperature) that is higher by 20°C than a liquidus temperature determined by each alloy composition and a casting step S2 for obtaining the casted aluminum alloy by: cooling the molten metal to a temperature that is lower than a solidus temperature by at least 10°C at the cooling rate shown in any one of Table 1 to Table 4; cooling the molten metal to a room temperature; and casting the molten metal into a plate-like shape having a thickness of 1.2 mm were performed as shown in Figs. 1(a) to (c). The casting step S2 was performed by the single roll continuous casting method using a copper roll.

[0144] Also, a post-treatment step S3 was performed on each of the casted aluminum alloys (Examples 1 to 48, Comparative Examples 1 to 20, and Comparative Examples 28 to 37) obtained after the casting step S2. As the post-treatment step S3, any one of hot rolling step S3a (Examples 1 to 36, Examples 41 to 48, Comparative Examples 1 to 20, and Comparative Examples 28 to 37), a cold rolling-heating step S3b (Example 37, Example 39, and Example 40), a heat treatment step S3c (Example 38) was performed. Types of the post-treatment steps performed on the casted aluminum alloys are shown in Table 1 to Table 4.

[0145] In the hot rolling step S3a, as shown in Fig. 1(a), each of the casted aluminum alloys after the casting step S2 was heated to a temperature of 450°C , and 40% of the thickness thereof was reduced by hot rolling to obtain an aluminum alloy material having a thickness of 0.72 mm. After that, the aluminum alloy material was allowed to cool to a room temperature.

[0146] In the cold rolling-heating step S3b, as shown in Fig. 1(b), cold rolling was performed on each of the casted aluminum alloys after the casting step S2 to reduce a thickness thereof by 40% to obtain an aluminum alloy material having a thickness of 0.72 mm. After that, the aluminum alloy material was heated at a temperature that is 1/2 of the melting point of the aluminum alloy or more (450°C in the present invention) for one hour. After that, the aluminum alloy material was allowed to cool to a room temperature.

[0147] In the heat treatment step S3c, as shown in Fig. 1(c), each of the casted aluminum alloys after the casting step S2 was heated at 450°C for one hour. After that, the aluminum alloy material was allowed to cool to a room temperature.

[0148] Further, in the present embodiment, as shown in Figs. 1(a) to 1(c), a heating step S4 of maintaining each of the aluminum alloy materials after the post-treatment step S3 to 300°C for 100 hours (in an assumption of long-term exposure to a temperature region equivalent to an engine traveling environment, for example) and allowing the aluminum alloy materials to cool to a room temperature was performed.

[0149] As described above, aluminum alloy materials (Examples 1 to 48, Comparative Examples 1 to 20, and Comparative Examples 28 to 37) on which the post-treatment step S3 and the heating step S4 have been performed were obtained.

[0150] In the present embodiment, in order to show superiority of the cooling rate, each of the aluminum alloys having the compositions shown in Table 4 was cast at a cooling rate of less than $150^{\circ}\text{C}/\text{sec.}$ to obtain an ingot for comparison, and aluminum alloy materials (Comparative Examples 21 to 27) were obtained by rolling the ingots.

[0151] In other words, as shown in Fig. 2(a), for preparing the ingot, a melting step S5 for preparing a molten metal

of each of the alloys by melting the alloy at a temperature (melting temperature) that is higher by 200°C than a liquidus temperature determined by the composition of each alloy and a solidifying step S6 for obtaining an aluminum alloy ingot by solidifying by cooling the molten metal at a cooling rate of 100°C/sec. were performed. By these steps, aluminum alloy ingots each in the form of a plate having a thickness of 1.2 mm were obtained.

[0152] After the ingot preparation, a hot rolling step S7a or a cold rolling-heating step S7b was performed as a post-treatment step S7. More specifically, the hot rolling step S7a was performed on Comparative Examples 21 to 23 and Comparative Examples 25 to 27, and the cold rolling-heating step S7b was performed on Comparative Example 24.

[0153] In the hot rolling step S7a, as shown in Fig. 2(a), each of the aluminum alloy ingots after the solidifying step S6 was heated to a temperature of 450°C, and 40% of a thickness thereof was reduced by hot rolling to obtain an aluminum alloy material having a thickness of 0.72 mm. After that, the aluminum alloy material was allowed to cool to a room temperature.

[0154] In the cold rolling-heating step S7b, as shown in Fig. 2(b), cold rolling was performed on each of the aluminum alloy ingots after the solidifying step S6 to reduce a thickness thereof by 40% to obtain an aluminum alloy material having a thickness of 0.72 mm. After that, the aluminum alloy material was heated at a temperature that is more than 1/2 of a melting point of the aluminum alloy (450°C in the present invention) for one hour and allowed to cool to a room temperature.

[0155] As shown in Figs. 2(a) and (b), after the post-treatment step S7, a heating step S8 of maintaining each of the aluminum alloy materials to 300°C for 100 hours (in an assumption of long-term exposure to a temperature region equivalent to an engine traveling environment, for example) and allowing the aluminum alloy material to cool to a room temperature was performed.

[0156] As described above, aluminum alloy materials (Comparative Examples 21 to 27) on which the melting step S5, the solidifying step S6, the post-treatment step S7, and the heating step S8 have been performed were obtained.

[0157] In Examples 1 to 40 and Examples 1 to 27, hardness HVR1 of the casted aluminum alloy before the post-treatment step S3 (S7), hardness HVR2 of the aluminum alloy material after the post-treatment step S3 (S7) and hardness HVR3 of the aluminum alloy material underwent the post-treatment step S3 (S7) and the heating step S4 (S8) which was performed subsequently to the post-treatment step were respectively measured, and softening resistance was evaluated by way of the change. The HVRn (n: No.) is referred to as residual hardness, and the residual hardness is in general significantly reduced when exposed to a high temperature region exceeding 1/2 of a material melting point. From such point of view, a casted aluminum alloy that is reduced in hardness reduction after the exposure to high temperature region was investigated.

[0158] The softening resistance is determined as follows: a pattern of $HVR1 < HVR2 < HVR3$ (pattern 1) is excellent (◎) as shown in Fig. 3(a); a pattern of $HVR1 < HVR2$, $HVR1 < HVR3$, and $HVR2 > HVR3$ (pattern 2) is good (○) as shown in Fig. 3(b); and other patterns, such as a pattern of $HVR1 > HVR2 > HVR3$ (pattern 3) is no good (x) as shown in Fig. 3(c). Evaluation results of softening resistance of Examples 1 to 48 and Comparative Examples 1 to 37 are shown in Table 5 to Table 8.

[0159] In Figs. 3(a) to (c), a horizontal axis is a distinction among HVR1, HVR2, and HVR3, and a vertical axis is Vickers hardness HV.

[0160] As is understood from Table 5 and Table 6, each of the casted aluminum alloys of Examples 1 to 48 shows the behavior of the pattern 1 or the pattern 2 of the softening resistance, thereby revealing the excellent softening resistance.

[0161] In contrast, as is understood from the results of Table 7 and Table 8, the Al-transition element alloys are classified into "increasing type (◎ or ○)" showing the patterns 1 and 2 of Figs. 3(a) and (b) and "decreasing type (x)" of the pattern 3 of Fig. 3(c). As shown in Table 7, the decreasing type is a phenomenon observed in general-purpose Al alloys.

[0162] In order to investigate the reasons for the above-described excellent softening resistance of the casted aluminum alloys of Examples 1 to 48, changes in alloy structure before and after the hot rolling step of one of the casted aluminum alloys (Example 1) among the above-specified examples were observed by using a scanning electron microscope. A microscope photograph of the alloy structure before hot rolling is shown in Fig. 4, and a microscope photograph of the alloy structure after hot rolling is shown in Fig. 5. As is understood from Fig. 4 and Fig. 5, in metal structures of an α phase formed of an Al matrix and a layered structure formed of an eutectic structure of an Al-Fe compound and an Al matrix, a stable compound phase (precipitate) formed of Al, Ti, and the third component element was precipitated in the Al matrix. The softening resistance is improved by the stable compound phase (precipitate), and it is considered that the strength was improved after the above-described processings and heating and the like. A result (photograph) of observation by the scanning electron microscope of the alloy structure of the aluminum alloy material after the rolling at a magnification different from that of Fig. 5 is shown in Fig. 7. As is understood from Fig. 7, the aluminum alloy material 1 of Example 11 has an α phase 2 formed of an Al matrix and a layered phase 4 formed in such a manner as to surround the α phase. In the alloy structure of the aluminum alloy material 1 (Example 11) after the hot rolling, a precipitate 3 having a particle diameter of about 15 nm or less is generated in the Al matrix.

[0163] Other photographs by the scanning electron microscope (SEM) of the alloy structure of the casted aluminum alloy of Example 11 before hot rolling are shown in Fig. 8 and Fig. 9. Shown in Fig. 8 is a photograph by SEM at a

magnification of x1000 of observation of the alloy structure of the casted aluminum alloy (Example 11), and shown in Fig. 9 is a photograph by SEM at a magnification of x5000 of observation of the alloy structure of the casted aluminum alloy (Example 11). Fig. 9 is an enlarged view of a part of Fig. 8 in which a crystallizate was generated.

[0164] Also, for the comparison with Example 11, scanning electron microscope photographs of the alloy structure of the casted aluminum alloy of Comparative Example 22 before hot rolling are shown in Fig. 10 and Fig. 11. Shown in Fig. 10 is a photograph by SEM at a magnification of x1000 of observation of the alloy structure of the casted aluminum alloy (Comparative Example 22), and shown in Fig. 11 is a photograph by SEM at a magnification of x5000 of observation of the alloy structure of the casted aluminum alloy (Comparative Example 22). Fig. 11 is an enlarged view of a part of Fig. 10 in which a crystallizate was generated.

[0165] The scanning electron microscope was S-3600N manufactured by Hitachi, Ltd., and the observation was conducted under the condition of an accelerating voltage of 15 kV.

[0166] As shown in Figs. 8 and 9, in the α phase, there were few crystallizates (compound ($\text{Al}_x(\text{Ti}, \text{X})$) of Al, the second component element Ti, and the third component element X) having a particle diameter of 5 μm or more in an alloy structure of the casted aluminum alloy of Example 11, and an area ratio of the crystallizate was less than 5%.

[0167] In contrast, as is understood from Fig. 10 and Fig. 11, in an alloy structure of the casted aluminum alloy 9 of Comparative Example 22, a relatively large amount of bulk crystallizates 93 (compound ($\text{Al}_x(\text{Ti}, \text{X})$) of Al, the second component element Ti, the third component element X) having a particle diameter of 5 μm or more was dispersed in an Al matrix of an α phase 92. Also, as is understood from a comparison between Fig. 9 and Fig. 11, the crystallizate generated in Comparative Example 22 was relatively larger than that of Example 11.

[0168] Results of component analysis in a region of the casted aluminum alloy (Comparative Example 22) where the crystallizate was observed are shown (see Fig. 12). In the diagram, relative amounts of each component (Al, Zr, Ti, Fe) in a region indicated by a straight line A-A in a scanning electron microscope photograph of the casted aluminum alloy (Comparative Example 22) are shown as the sizes of peaks. In Fig. 12, a profile by a $\text{K}\alpha$ line is shown for each of Al, Ti, and Fe, and a profile by an $\text{L}\alpha$ line is shown for Zr. As is understood from Fig. 12, in the crystallizate, large amounts of the second component element Ti and the third component element Zr are present, and a compound of Al, Ti, and Zr is formed. For the analysis of the component amounts, an energy dispersive X ray analysis device manufactured by Edax Japan was used.

[0169] Results (photograph) of observation by a transmission electron microscope (TEM) of the casted aluminum alloy (Example 11) before hot rolling is shown in Fig. 13. HF-2000 of Hitachi, Ltd. was used as the transmission electron microscope, and the observation was conducted under the conditions of an accelerating voltage of 200 kV and a beam diameter of $\phi 1$ nm. As shown in Fig. 13, a metal structure of the casted aluminum alloy 1 of Examples 11 has an α phase 2 formed of an Al matrix and a layered phase 4 formed in such a manner as to surround the α phase 2. Subsequently, for arbitrary place in the layered phase 4 (points *1 to *4 in Fig. 13), component elements present in the layered phase 4 were investigated by conducting energy dispersive X ray analysis (EDX). In the EDX analysis, NORAN VOYAGER III M3100 was used as an energy dispersive X ray analysis device, and Si/Li semiconductor detector was used as a detector. Measurement was conducted under the conditions of an energy resolution of 137 eV and a loading time of 30 seconds. Results are shown in Figs. 14 to 17.

[0170] Shown in Fig. 14 to Fig. 17 are the results of the EDX analysis at points *1 to *4 in Fig. 13.

[0171] In the same manner, the aluminum alloy material (Example 11) after the hot rolling was observed by using the transmission electron microscope (TEM), and the EDX analysis at arbitrary four points of *1 to *4 was conducted. TEM photographs are shown in Fig. 18, and the results of EDX analysis at points *1 to *4 are shown in Fig. 19 to Fig. 22.

[0172] As is understood from Fig. 13 to Fig. 17 and Fig. 18 to Fig. 22, only Al and Fe were detected in the layered phase 4 irrespectively of whether it was before and after the hot rolling, and the second component element Ti and the third component Zr were not present (see Fig. 13 and Fig. 18). Therefore, it is understood that the second component element Ti and the third component element Zr are present in Al matrix of the α phase 2.

[0173] Subsequently, in the present embodiment, strength, processability, formability, and corrosion resistance of Example 1 to Example 48 and Comparative Examples 1 to 37 were evaluated. Evaluation methods are as follows.

<Strength>

[0174] A tensile test piece was cut out from each of the aluminum alloy materials, and a tensile test defined in JIS Z2241 was performed to obtain tensile strength. Results are shown in Table 5 to Table 8.

[0175] Also, a relationship between the tensile strength measured by the tensile test and the cooling rate is shown in Fig. 6. Shown in Fig. 6 is a semi-logarithmic graph wherein a horizontal axis is the cooling rate ($^{\circ}\text{C}/\text{sec.}$), and a vertical axis is the tensile strength (MPa). In Fig. 6, a relationship between the cooling rate and the tensile strength of each of three types aluminum alloy compositions, i.e. of the aluminum alloy materials of Al-2Fe-1Zr-0.8Ti (Comparative Example 21, Example 48, Example 9, Example 47, Example 46), Al-4Fe-1Zr-0.8Ti (Comparative Example 22, Example 43, Example 11, Example 42, Example 41), and Al-4Fe-1Zr-0.8Ti-0.5Mg (Example 45, Example 44).

<Processability>

[0176] Determination of processability was conducted by observation of presence/absence of generation of rolling crack after the rolling (hot rolling or cold rolling).

[0177] In other words, a surface of each of the aluminum alloy materials after rolling was observed, and the case wherein a rolling crack was observed on the surface was evaluated as no good (x), while the case wherein any rolling crack was not observed was evaluated as good (○). The continuously cast material in which only a cracked edge (crack generated on both ends of continuously cast material) was generated was evaluated as good (○) since it is possible to eliminate the cracked edge by a slitter in an actual process. Processability of the aluminum alloy material (Example 38) on which rolling was not performed was not evaluated.

<Formability>

[0178] Formability was observed by conducting a Hemming processing limit evaluation test for automobile aluminum alloy plate defined in JIS H7701, and generation of a crack on a surface at a bent part was observed by a stereoscopic microscope. The case wherein a crack was observed on the surface was evaluated as no good (x), and the case wherein any crack was not observed was evaluated as good (o). Results are shown in Table 5 to Table 8.

<Corrosiveness>

[0179] Corrosiveness was evaluated by conducting a corrosiveness test on a 6061 alloy and performing comparison with the result.

[0180] In other words, a test piece having a certain dimension was cut out from a commercially available 6016 alloy (Al-1.1Mg-0.8Si-0.1Cu-0.1Cr-0.03Ti), and a weight W1 of the test piece was measured. Subsequently, by using a NaCl aqueous solution having a concentration of 5 wt%, a saline water spraying test was conducted on the test piece (JIS Z2371). Subsequently, after removing a corrosive product generated on a surface of the test piece, a weight (W2) of the test piece was measured. A weight change ratio ΔW_a (%) of the 6061 alloy test piece was calculated based on an expression of $\Delta W_a = |W2 - W1| \times 100/W1$.

[0181] A test piece having a certain dimension was prepared from each of the casted aluminum alloys of Examples 1 to 48 and Comparative Example 1 to 37, and a saline water spraying test was conducted in the same manner as in the 6061 alloy. A weight W3 before the test and a weight W4 after the test were measured, and a weight change ratio ΔW_b (%) of each of the test pieces was calculated based on an expression of $\Delta W_b = |W4 - W3| \times 100/W3$.

[0182] In judgment of corrosiveness, the case of $\Delta W_b < 0.8\Delta W_a$ was evaluated as excellent (⊙), the case of $0.8\Delta W_a \leq W_b \leq 1.2\Delta W_a$ was evaluated as good (○), and the case of $\Delta W_b > 1.2\Delta W_a$ was evaluated as no good (×). The results are shown in Table 5 to Table 8.

[0183] As is understood from Table 5, Table 6 and Fig. 6, Example 1 to Example 48 exhibited satisfactory strength that was the tensile strength of 230 MPa or more and are casted aluminum alloys excellent in softening resistance, formability, and corrosion resistance.

[0184] As is apparent from the results of Example 1 to Example 48 (Table 5 and Table 6), by adding the second component element Ti and the third component element (Zr, Nb, Hf, Sc, Y) to Al-Fe alloy which is the base, the Al alloy having the high strength is obtained without impairment of formability and corrosion resistance. Also, when so required, it is possible to further improve the properties by adding the fourth component element Mg, the fifth component element (Cu, Cr, Co), and the sixth component element (V, Mo).

[0185] Also, it was found that the Example 1 to Example 48 are excellent in softening resistance, and higher strength thereof is achieved by applying thereto heat energy or strain energy in the subsequent steps. Also, reductions in properties depending on use condition (e.g. exposure to 300°C for long time) are considerably small. Therefore, the alloys of the present invention are suitably used for automobile parts, for example.

[0186] In contrast, as is understood from Table 7 and Table 8, in the case of using the aluminum alloy that are out of the alloy composition range defined in the present invention (Comparative Example 1 to Comparative Example 20 and Comparative Example 28 to Comparative Example 37) and in the case where the cooling rate is insufficient (Comparative Examples 21 to 27), properties of the alloy cast materials are deteriorated.

[0187] (Table 1)

(Table 1)

code	Al	essential component element (mass%)								selective component element (mass%)						inevitable element (mass%)				specific gravity (g/cm ³)	cooling rate of casting (°C/sec)	post-treatment		
		first second		third				fourth	fifth		sixth		Si	Mn	Ni	Be								
		Fe	Ti	Zr	Nb	Hf	Sc		Y	Mg	Cu	Cr					Co	V	Mo					
Example 1	Bal.			0.2	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling
Example 2				-	0.2	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling
Example 3		0.8	0.15	-	-	0.2	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling
Example 4				-	-	-	0.2	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.71	hot rolling
Example 5				-	-	-	-	-	0.2	-	-	-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling
Example 6						0.5	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.74	hot rolling
Example 7			2	0.2	-	0.5	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.75	hot rolling
Example 8					-	-	0.5	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.75	hot rolling
Example 9				0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.76	hot rolling
Example 10				0.2	0.5	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.78	hot rolling
Example 11			4	0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.80	hot rolling
Example 12				1	2	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.82	hot rolling
Example 13			5	0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.83	hot rolling
Example 14						-	-	-	-	0.5	-	-	-	-	-	-	-	-	-	-	-	-	2.76	hot rolling
Example 15				0.3	0.8	-	-	-	-	1	-	-	-	-	-	-	-	-	-	-	-	-	2.74	hot rolling
Example 16						-	-	-	-	2	-	-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling
Example 17				0.8	1	-	-	-	-	1	-	-	-	-	-	-	-	-	-	-	-	-	2.74	hot rolling
Example 18			4			-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.77	hot rolling
Example 19				0.3	0.8	-	-	-	-	-	0.2	-	-	-	-	-	-	-	-	-	-	-	2.75	hot rolling
Example 20			2			-	-	-	-	-	0.5	-	-	-	-	-	-	-	-	-	-	-	2.76	hot rolling
Example 21			4		0.8	-	-	-	-	-	1	-	-	-	-	-	-	-	-	-	-	-	2.77	hot rolling
Example 22			2	0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.77	hot rolling
Example 23						-	-	-	-	-	-	-	1	-	-	-	-	-	-	-	-	-	2.77	hot rolling

[0188] (Table 2)

(Table 2)

code	Al	essential component element (mass%)						selective component element (mass%)						inevitable element (mass%)				specific gravity (g/cm ³)	cooling rate of casting (°C/sec)	post-treatment		
		first	second	Zr	Nb	Hf	Sc	Y	fourth	Cu	Cr	Co	V	Mo	Si	Mn	Ni				Be	
Example 24	Bal.	0.8	0.1	0.2	-	-	-	-	-	-	-	-	0.1	-	<0.5			<0.02	<0.05	-	2.72	hot rolling
Example 25		2	0.3	0.5	-	-	-	-	-	-	-	-	0.45	-						-	2.75	hot rolling
Example 26					-	-	-	-	-	-	-	0.2	0.2	-						2.75	hot rolling	
Example 27					-	-	-	-	-	-	-	-	0.2	-						2.75	hot rolling	
Example 28		-	-	0.8	-	-	-	-	-	-	-	-	-	0.45	-	2.76	hot rolling					
Example 29		-	-	1	-	-	-	-	-	-	-	-	-	-	-	2.76	hot rolling					
Example 30		-	-		-	-	-	-	-	-	-	-	-	-	-	2.80	hot rolling					
Example 31		-	-	0.5	-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling					
Example 32		2	0.2		-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling					
Example 33					-	-	-	-	-	-	-	-	-	-	-	2.72	hot rolling					
Example 34		-	-	0.3	-	-	-	-	-	-	-	-	-	-	-	2.77	hot rolling					
Example 35		-	-		-	-	-	-	-	-	-	-	-	-	-	2.73	hot rolling					
Example 36		3	0.2	0.6	-	-	-	-	-	1	-	0.1	-	-	-	<0.01	2.75	hot rolling				
Example 37		2	0.8	1	-	-	-	-	-	2	-	0.8	-	-	-			2.76	cold rolling			
Example 38		4	0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	2.76	heat treatment				
Example 39					-	-	-	-	-	-	-	-	-	-	-	-	-	2.80	cold rolling			
Example 40	2	0.2	0.5	-	-	-	-	-	2	-	-	-	-	-	<0.5	<0.05	2.72	cold rolling				
Example 41	4	0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	5000	hot rolling			
Example 42				-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.80	hot rolling		
Example 43				-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	170	hot rolling	
Example 44	-	-	0.8	-	-	-	-	-	-	-	-	-	-	-	<0.02	2.79	1000	hot rolling				
Example 45	-	-		-	-	-	-	-	-	-	-	-	-	-			-	170	hot rolling			
Example 46	2	0.8	1	-	-	-	-	-	0.5	-	-	-	-	-	-	-	-	5000	hot rolling			
Example 47				-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1000	hot rolling		
Example 48				-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	170	hot rolling	

[0189] (Table 3)

(Table 3)

code	Al	essential component element (mass%)							selective component element (mass%)					inevitable element (mass%)				specific gravity (g/cm ³)	cooling rate of casting (°C/sec)	post-treatment			
		first	second	Zr	Nb	Hf	Sc	Y	fourth	Cu	Cr	Co	V	Mo	Si	Mn	Ni				Be		
Comparative Example 1	Bal.	0.5	-	-	-	-	-	-	-	-	-	-	-	-	<0.5	<0.02	-	-	2.71	hot rolling			
Comparative Example 2		2	-	-	-	-	-	-	-	-	-	-	-	-			-	-	-	2.73	hot rolling		
Comparative Example 3		4	-	-	-	-	-	-	-	-	-	-	-	-			-	-	-	-	2.77	hot rolling	
Comparative Example 4			-	1	-	-	-	-	-	-	-	-	-	-			-	-	-	-	2.79	hot rolling	
Comparative Example 5		7	0.8	-	-	-	-	-	-	-	-	-	-	-			-	-	-	-	2.78	hot rolling	
Comparative Example 6			-	1	-	-	-	-	-	-	-	-	-	-			-	-	-	-	2.86	hot rolling	
Comparative Example 7		-	0.3	-	-	-	-	-	-	-	-	-	-	-			-	-	-	-	2.72	hot rolling	
Comparative Example 8		1	-	0.8	2	-	-	-	-	-	-	-	-	-			-	-	-	-	2.74	hot rolling	
Comparative Example 9			-	-	-	-	-	-	-	-	-	-	-	-			-	-	-	-	2.76	hot rolling	
Comparative Example 10		0.5	0.3	-	-	2	-	-	-	-	-	-	-	-			-	-	-	-	2.76	hot rolling	
Comparative Example 11			-	-	-	-	0.8	-	-	-	-	-	-	-			-	-	-	-	-	2.73	hot rolling
Comparative Example 12		0.5	0.15	1	-	-	-	0.2	-	-	-	-	-	-			-	-	-	-	-	2.73	hot rolling
Comparative Example 13			0.3	-	-	-	-	-	1	-	-	-	-	-			-	-	-	-	-	2.74	hot rolling
Comparative Example 14		2	0.8	1	-	-	-	-	-	0.03	-	-	-	-			-	-	-	-	-	2.76	hot rolling
Comparative Example 15			-	-	-	-	-	-	-	-	3	-	-	-			-	-	-	-	-	2.70	hot rolling
Comparative Example 16		0.3	0.5	-	-	-	-	-	-	1.5	-	-	-	-			-	-	-	-	-	2.78	hot rolling
Comparative Example 17											-	-	-	-			-	-	-	-	-	-	-
Comparative Example 18		0.8	-	-	-	-	-	-	-	-	-	1.5	-	-			-	-	-	-	-	2.78	hot rolling
Comparative Example 19													-	-			-	-	-	-	-	-	-
Comparative Example 20		-	-	-	-	-	-	-	-	-	-	-	-	-			-	-	-	-	-	2.76	hot rolling

[0190] (Table 4)

(Table 4)

code	Al	essential component element (mass%)										selective component element (mass%)						inevitable element (mass%)				specific gravity (g/cm ³)	cooling rate of casting (°C/sec)	post-treatment
		first		second	third				fourth	fifth		sixth		Si	Mn	Ni	Be							
		Fe	Ti	Zr	Nb	Hf	Sc	Y	Mg	Cu	Cr	Co	V					Mo						
Comparative Example 21	Bal.	2			-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.76	100	hot rolling		
Comparative Example 22		4			-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.80		hot rolling		
Comparative Example 23		2			-	-	-	-	1	-	-	-	-	-	-	-	-	-	-	2.74		hot rolling		
Comparative Example 24			0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.74		cold rolling		
Comparative Example 25					-	-	-	-	-	1	-	-	-	-	-	-	-	-	-	2.78	hot rolling			
Comparative Example 26				-	-	-	-	-	-	-	1	-	-	-	-	-	-	-	-	2.78	hot rolling			
Comparative Example 27				-	-	-	-	-	-	-	-	1	-	-	-	-	-	-	-	2.78	hot rolling			
Comparative Example 28		0.5	0.1	0.3	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.72	280	hot rolling		
Comparative Example 29		4			0.5	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.79		hot rolling		
Comparative Example 30					0.3	0.2	-	-	-	-	-	-	-	-	-	-	-	-	-	2.79		hot rolling		
Comparative Example 31					0.03	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.78		hot rolling		
Comparative Example 32				2.5	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.82	hot rolling			
Comparative Example 33			-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1	-	2.79	hot rolling			
Comparative Example 34			-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1	-	2.78	hot rolling			
Comparative Example 35			0.8	2.5	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.82	2000	hot rolling		
Comparative Example 36			1.5	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.81		hot rolling		
Comparative Example 37		7	0.8	1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	2.86		hot rolling		

[0191] (Table 5)

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(Table 5)

code	softening resistance	tensile strength (Mpa)	processability	formability	corrosiveness
Example 1	○	260	○	○	◎
Example 2	○	265	○	○	◎
Example 3	○	260	○	○	◎
Example 4	○	255	○	○	◎
Example 5	○	258	○	○	◎
Example 6	◎	320	○	○	◎
Example 7	◎	315	○	○	◎
Example 8	◎	320	○	○	◎
Example 9	◎	325	○	○	◎
Example 10	◎	385	○	○	◎
Example 11	◎	420	○	○	◎
Example 12	◎	450	○	○	◎
Example 13	◎	455	○	○	◎
Example 14	◎	340	○	○	◎
Example 15	◎	410	○	○	◎
Example 16	◎	460	○	○	◎
Example 17	○	345	○	○	○
Example 18	◎	460	○	○	◎
Example 19	○	360	○	○	◎
Example 20	○	410	○	○	○
Example 21	○	440	○	○	○
Example 22	◎	385	○	○	◎
Example 23	◎	380	○	○	◎

[0192] (Table 6)

(Table 6)

code	softening resistance	tensile strength (Mpa)	processability	formability	corrosiveness
Example 24	○	260	○	○	◎
Example 25	◎	320	○	○	◎
Example 26	◎	320	○	○	◎
Example 27	◎	320	○	○	◎
Example 28	○	285	○	○	◎
Example 29	◎	390	○	○	◎
Example 30	◎	415	○	○	◎
Example 31	◎	410	○	○	◎
Example 32	◎	405	○	○	◎
Example 33	◎	385	○	○	◎

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(continued)

	code	softening resistance	tensile strength (Mpa)	processability	formability	corrosiveness
5	Example 34	⊙	470	○	○	⊙
	Example 35	⊙	385	○	○	⊙
	Example 36	○	410	○	○	○
	Example 37	⊙	285	○	○	⊙
10	Example 38	⊙	305	-	○	⊙
	Example 39	⊙	385	○	○	⊙
	Example 40	○	355	○	○	○
15	Example 41	⊙	495	○	○	⊙
	Example 42	⊙	460	○	○	⊙
	Example 43	⊙	360	○	○	⊙
	Example 44	⊙	495	○	○	⊙
20	Example 45	⊙	398	○	○	⊙
	Example 46	⊙	380	○	○	⊙
	Example 47	⊙	365	○	○	⊙
25	Example 48	⊙	230	○	○	⊙

[0193] (Table 7)

(Table 7)

	code	softening resistance	tensile strength (Mpa)	processability	formability	corrosiveness
30	Comparative Example 1	×	220	○	○	⊙
	Comparative Example 2	×	260	○	○	⊙
35	Comparative Example 3	×	270	○	○	⊙
	Comparative Example 4	×	320	○	○	⊙
	Comparative Example 5	×	295	○	○	⊙
	Comparative Example 6	⊙	445	×	×	⊙
40	Comparative Example 7	×	205	○	○	⊙
	Comparative Example 8	×	255	○	○	⊙
	Comparative Example 9	×	235	○	○	⊙
45	Comparative Example 10	×	235	○	○	⊙
	Comparative Example 11	×	230	○	○	⊙
	Comparative Example 12	×	210	○	○	⊙
	Comparative Example 13	×	230	○	○	⊙
50	Comparative Example 14	⊙	325	○	○	○
	Comparative Example 15	⊙	415	○	×	○
	Comparative Example 16	○	430	○	×	×
55	Comparative Example 17	⊙	385	○	×	○
	Comparative Example 18	○	385	○	×	○
	Comparative Example 19	⊙	365	○	×	⊙

(continued)

code	softening resistance	tensile strength (Mpa)	processability	formability	corrosiveness
Comparative Example 20	⊙	345	○	×	○

[0194] (Table 8)

(Table 8)

code	softening resistance	tensile strength (Mpa)	processability	formability	corrosiveness
Comparative Example 21	×	235	○	○	⊙
Comparative Example 22	×	275	○	×	⊙
Comparative Example 23	×	250	○	○	⊙
Comparative Example 24	×	245	○	×	○
Comparative Example 25	×	250	○	○	⊙
Comparative Example 26	×	250	○	○	⊙
Comparative Example 27	×	250	○	○	⊙
Comparative Example 28	×	225	○	○	⊙
Comparative Example 29	×	305	○	○	⊙
Comparative Example 30	×	300	○	○	⊙
Comparative Example 31	×	295	×	×	⊙
Comparative Example 32	○	450	×	×	⊙
Comparative Example 33	×	375	×	×	×
Comparative Example 34	×	350	○	○	○
Comparative Example 35	○	475	×	×	⊙
Comparative Example 36	×	410	○	○	⊙
Comparative Example 37	⊙	480	×	×	⊙

[0195] Also, in the present embodiment, a relationship between the annealing (heating) temperature and the residual hardness was investigated for an aluminum alloy material on which cold rolling was performed after a casting step and a casted aluminum alloy on which rolling was not performed.

[0196] More specifically, a casted aluminum alloy was prepared with a composition and under a condition similar to those of Example 11 (see Table 1). Cold rolling was performed on the casted aluminum alloy under a room temperature condition to reduce a thickness of the casted aluminum alloy by 50%. Heating (annealing) at a predetermined temperature was performed for one hour, and residual hardness of the aluminum alloy material after the heating was investigated. A relationship between the heating (annealing) temperature and the residual hardness was plotted on a graph. Results are shown in Fig. 23. Measurement of the residual hardness was conducted by using a Vickers hardness tester under the conditions of loading of 100 gf and a retention time of 20 seconds.

[0197] Also, a relationship between the heating (annealing) temperature and the residual hardness of the case of performing only annealing at each of the specified temperatures on a casted aluminum alloy produced with a composition and under a condition similar to those of Example 11 (see Table 1) was plotted on a graph. Results are shown in Fig. 23.

[0198] As is understood from Fig. 23, the residual hardness is improved by heating in both of the case where heating is performed after rolling and the case where heating is performed without rolling. Particularly, it is possible to more satisfactorily improve the residual hardness in the case of heating at 400°C to 500°C, and it is possible to yet further improve the residual hardness in the case of heating at 400°C to 450°C.

(Embodiment 2)

[0199] This embodiment is one example of producing a casted aluminum alloy by die casting.

[0200] As shown in Figs. 24(a) and (b), the casted aluminum alloy of the present embodiment has a base part 11 in the form of a cylindrical column and a plate-like part 15 integrally formed on the base part 11 and having a length L of 90 mm and a width W of 50 mm. The plate-like part 15 is formed of a large thickness plate part 12 (thickness t_1 : 4 mm), a medium thickness plate part 13 (thickness t_2 : 3 mm), and a thin plate part 14 (thickness t_3 : 2 mm), which are different from one another in thickness. The large thickness plate part 12, the medium thickness plate part 13, and the thin plate part 14 are so formed as to be identical in size except for the thickness and have a length L_1 of 30 mm and a width W of 50 mm.

[0201] In the present embodiment, a casted aluminum alloy 1 having a shape shown in Figs. 24(a) and (b) was produced by preparing a molten metal by melting an aluminum alloy having a specific composition at a temperature higher by 20°C or more than a liquidus temperature thereof and casting the molten metal by die casting.

[0202] More specifically, an aluminum alloy containing 4 mass% of Fe, 0.85 mass% of Ti, and 1 mass% of Zr was prepared, and a molten metal was prepared by melting the alloy at a temperature (melting temperature) higher by 20°C or more than a liquidus temperature of the alloy. Subsequently, the molten metal was injected and cast into a die having a desired shape. As shown in Figs. 24(a) and (b), since the large thickness plate part 12, the medium thickness plate part 13, and the thin plate part 14 are different from one another in thickness, these parts are cooled at different cooling rates in the casting. In the present embodiment, cooling was performed in such a manner that the large thickness plate part 12, the medium thickness plate part 13, and the thin plate part 14 are cooled at cooling rates of 80°C/sec., 100°C/sec., and 400°C/sec. Thus, the casted aluminum alloy 1 was produced as shown in Figs. 24 (a) and (b).

[0203] Subsequently, an alloy structure of the casted aluminum alloy produced in the present embodiment was observed by a metallographic microscope. Results of the observation are shown in Figs. 25(a) to (c). Shown in Fig. 25(a) is the alloy structure of the thin plate part; Fig. 25(b) is the alloy structure of the medium thickness plate part; and Fig. 25(c) is the alloy structure of the large thickness plate part.

[0204] As shown in Fig. 25(a), in the thin plate part 14 cooled at the cooling rate of 400°C/sec., a bulk crystallize is scarcely formed in the alloy structure. In contrast, in the medium thickness plate 13 that was cooled at the cooling rate of 100°C/sec. and the large thickness plate part 12 that was cooled at a cooling rate of 80°C/sec., a large amount of crystallizes 19 was generated in the alloy structures.

[0205] An area ratio of a crystallize having a particle diameter of 5 μm or more in each of the thin plate part, the medium thickness plate part, and the large thickness plate part was measured. More specifically, the area ratio of the crystallize in a observation visual field area (1 mm²) was measured using an image analysis processing device by polishing each of the thin plate part, the medium thickness plate part, and the large thickness plate part to a mirror surface and measuring an area ratio of the obtained image by using an image analysis processing device while observing the mirror surface with a metallographic microscope having a magnification of $\times 1000$. Results of the measurement are shown in Table 9 described later.

[0206] Residual hardness of each of the thin plate part, the medium thickness plate part, and the large thickness plate part was measured before and after annealing. Results are shown in Table 9. The annealing was performed under the conditions of 450°C and one hour, and the residual hardness measurement was performed by using a Vickers hardness tester under the conditions of a loading of 100 gf and a retention time of 20 seconds.

[0207] (Table 9)

(Table 9)

part	plate thickness (mm)	colling rate (°C/sec)	area ratio of crystallize (%)	residual hardness HVR of before annealing	residual hardness HVR of after annealing
thin plate part	2	400	2.0	85	130
medium thickness plate part	3	100	5.2	68	74
large thickness plate part	4	80	8.5	65	65

[0208] As is understood from Table 9, the residual hardness in the thin plate part was largely increased after the annealing. In contrast, the residual hardness of each of the medium thickness plate part and the large thickness plate part was not increased much after the annealing and was scarcely changed. Therefore, it is understood that the casted aluminum alloy exhibits the particularly excellent softening resistance at the thin plate part.

[0209] In the casted aluminum alloy of the present embodiment, the thin plate part is the part which was cooled at the cooling rate of 150°C/sec. or more and less than 10000°C/sec. Therefore, it is considered that it is possible to achieve the effects of the present invention also in the case where the casting is performed by the die casting method by adjusting the cooling rate within the range of 150°C/sec. or more and less than 10000°C/sec.

Claims

1. A method for producing a casted aluminum alloy obtained by casting a molten metal of an aluminum alloy, **characterized by** comprising:

a melting step of preparing an aluminum alloy consisting of 0.8 to 5 mass% of Fe as a first component element, 0.15 to 1 mass% of Ti as a second component element, one or more kinds of third component elements selected from a third component element group consisting of Zr, Nb, Hf, Sc, and Y at a content of 0.05 to 2 mass% of each of the third components and in an amount that satisfies $Fe > X > Ti$ when the content of the first component element Fe is Fe (mass%), the content of the second component element Ti is Ti (mass%), and a total content of the third component elements is X (mass%); optionally 0.05 to 2 mass% of Mg as a fourth component element, optionally 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr, and Co, and optionally V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%, the residual part being Al and inevitable impurities, and melting the aluminum alloy at a temperature that is higher by 20°C or more than a liquidus temperature determined by the composition of the aluminum alloy to obtain a molten metal; and a casting step of casting the molten metal into a casting mold and then cooling the molten metal to a temperature that is lower by at least 10°C than a solidus temperature determined by the composition of the aluminum alloy at a cooling rate of 150°C/sec. or more and less than 10000°C/sec.

2. The method according to claim 1, **characterized by** using the aluminum alloy comprising 0.2 to 1.2 mass% of Zr among the third component element group in the melting step.

3. The method according to claim 1 or 2, **characterized in that** a total amount of the fourth component element, the fifth component element, and the sixth component element in the aluminum alloy is maintained at 3 mass% or less.

4. The method according to any one of claims 1 to 3, further comprising a hot rolling step of reducing the thickness of the casted aluminum alloy obtained by the casting step by 30% or more by performing hot rolling on the casted aluminum alloy at a temperature of 200°C or more.

5. The method according to claim 4, **characterized in that** the hot rolling step is performed after casting the molten metal into a plate-like shape in the casting mold while cooling the molten product to a temperature of 200°C to 500°C and releasing from the casting mold in the casting step.

6. The method according to any one of claims 1 to 3, further comprising a cold rolling-heating step of performing cold rolling on the casted aluminum alloy obtained by the casting step thereby to reduce the thickness of the casted aluminum alloy by 30% or more, and subsequently heating at a temperature of 1/2 or more of the melting point of the aluminum alloy and 550°C or less.

7. The method according to any one of claims 1 to 3, further comprising a heat treatment step of heating the casted aluminum alloy obtained by the casting step at a temperature of 400°C or more for 0.5 to 3 hours.

8. A casted aluminum alloy **characterized by** consisting of 0.8 to 5 mass% of Fe as a first component element, 0.15 to 1 mass% of Ti as a second component element, one or more kinds of third component elements selected from a third component element group consisting of Zr, Nb, Hf, Sc, and Y at a content of 0.05 to 2 mass% of each of the third components and in an amount that satisfies $Fe > X > Ti$ when the content of the first component element Fe is Fe (mass%), the content of the second component element Ti is Ti (mass%), and a total content of the third component elements is X (mass%); optionally 0.05 to 2 mass% of Mg as a fourth component element, optionally 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr, and Co, and optionally V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%,

the residual part being Al and inevitable impurities,
wherein the casted aluminum alloy has a metal structure including an α phase that is formed of an Al matrix and a layered phase that is formed of an eutectic structure of the Al matrix and an Al-Fe compound in such a manner as to surround the α phase;

the Al matrix is formed of a supersaturated solid solution of Al in which the second component element and the third component element are dissolved; and

an area ratio of a crystallite formed of an intermetallic compound of Al, the second component element, and the third component element and having a particle diameter of 5 μm or more is less than 5% in an arbitrary section of the casted aluminum alloy.

9. An aluminum alloy material **characterized by** consisting of
0.8 to 5 mass% of Fe as a first component element,
0.15 to 1 mass% of Ti as a second component element,
one or more kinds of third component elements selected from a third component element group consisting of Zr, Nb, Hf, Sc, and Y at a content of 0.05 to 2 mass% of each of the third components and in an amount that satisfies $\text{Fe} > \text{X} > \text{Ti}$ when the content of the first component element Fe is Fe (mass%), the content of the second component element Ti is Ti (mass%), and a total content of the third component elements is X (mass%); optionally 0.05 to 2 mass% of Mg as a fourth component element, optionally 0.05 to 1 mass% of at least one fifth component element selected from a fifth component element group consisting of Cu, Cr, and Co, and optionally V and/or Mo as a sixth component element in an amount of more than 0.05 mass% and less than 0.5 mass%, and
the residual part being Al and inevitable impurities,
wherein the aluminum alloy material has a metal structure including an α phase that is formed of an Al matrix and a layered phase that is formed of an eutectic structure of the Al matrix and an Al-Fe compound in such a manner as to surround the α phase;
the Al matrix is formed of Al and/or a supersaturated solid solution of Al in which the second component element and the third component element are dissolved; and
a precipitate formed of an intermetallic compound of Al, the second component element, and the third component element and having a particle diameter of 2 to 500 nm is dispersed in the Al matrix.

10. The casted aluminum alloy according to claim 8 or the aluminum alloy material according to claim 9, **characterized by** comprising 0.2 to 1.2 mass% of Zr among the third component element group.

11. The casted aluminum alloy according to claim 8 or 10 or the aluminum alloy material according to claim 9 or 10, **characterized in that** Mg as the fourth component element is dissolved at least into the Al matrix and/or forms an Al-Mg compound in the Al matrix.

12. The casted aluminum alloy according to any one of claims 8, 10 and 11 or the aluminum alloy material according to any one of claims 9 to 11, **characterized in that** Cr and/or Co as the fifth component element is substituted with at least a part of the Al-Fe compound forming the layered phase to form an Al-(Fe, Cr) compound and/or an Al-(Fe, Co) compound.

13. The casted aluminum alloy according to any one of claims 8 and 10 to 12 or the aluminum alloy material according to any one of claims 9 to 12, **characterized in that** Cu as the fifth component element forms an Al-Cu compound in the Al matrix.

14. The casted aluminum alloy according to claim 8 or 10 or the aluminum alloy material according to claim 9 or 10, **characterized in that** the aluminum alloy material comprises 0.05 to 2 mass% of Mg as a fourth component element and 0.05 to 1 mass% of Cu as a fifth component element, and that the fourth component element and the fifth component element form an Al-Cu-Mg compound in the Al matrix.

15. The casted aluminum alloy according to any one of claims 8 and 10 to 14 or the aluminum alloy material according to any one of claims 9 to 14, **characterized in that** V and/or Mo as the sixth component element form(s) a compound with Al and a compound with Al, the second component element, and the third component element in the Al matrix.

Patentansprüche

1. Verfahren zur Herstellung einer gegossenen Aluminiumlegierung, die durch Gießen eines geschmolzenen Metalls

aus einer Aluminiumlegierung erhalten wird, **dadurch gekennzeichnet, dass** es umfasst:

einen Schmelzschrift des Herstellens einer Aluminiumlegierung, die aus 0,8 bis 5 Massen-% Fe als ein erstes Komponentenelement, 0,15 bis 1 Massen-% Ti als ein zweites Komponentenelement, einer oder mehreren Art(en) von dritten Komponentenelementen, ausgewählt aus einer Gruppe von dritten Komponentenelementen, bestehend aus Zr, Nb, Hf, Sc und Y in einem Gehalt von 0,05 bis 2 Massen-% von jeder der dritten Komponenten, und in einer Menge, die $Fe > X > Ti$ genügt, wenn der Gehalt des ersten Komponentenelements Fe Fe (Massen-%) ist, der Gehalt des zweiten Komponentenelements Ti Ti (Massen-%) ist und der Gesamtgehalt der dritten Komponentenelemente X (Massen-%) ist, gegebenenfalls 0,05 bis 2 Massen-% Mg als ein viertes Komponentenelement, gegebenenfalls 0,05 bis 1 Massen-% von mindestens einem fünften Komponentenelement, ausgewählt aus einer Gruppe von fünften Komponentenelementen, bestehend aus Cu, Cr und Co, und gegebenenfalls V und/oder Mo als ein sechstes Komponentenelement in einer Menge von mehr als 0,05 Massen-% und weniger als 0,5 Massen-% besteht, wobei es sich bei dem Rest um Al und unvermeidbare Verunreinigungen handelt, und des Schmelzens der Aluminiumlegierung bei einer Temperatur, die um 20 °C oder mehr höher ist als eine Liquidus-Temperatur, die durch die Zusammensetzung der Aluminiumlegierung festgelegt ist, so dass ein geschmolzenes Metall erhalten wird, und einen Gießschritt des Gießens des geschmolzenen Metalls in eine Gießform und dann des Abkühlens des geschmolzenen Metalls auf eine Temperatur, die um mindestens 10 °C niedriger ist als eine Solidus-Temperatur, die durch die Zusammensetzung der Aluminiumlegierung festgelegt ist, bei einer Abkühlungsgeschwindigkeit von 150 °C/Sekunde oder mehr und weniger als 10000 °C/Sekunde.

2. Verfahren nach Anspruch 1, **dadurch gekennzeichnet, dass** in dem Schmelzschrift eine Aluminiumlegierung verwendet wird, die von der Gruppe der dritten Komponentenelemente 0,2 bis 1,2 Massen-% Zr umfasst.

3. Verfahren nach Anspruch 1 oder 2, **dadurch gekennzeichnet, dass** die Gesamtmenge des vierten Komponentenelements, des fünften Komponentenelements und des sechsten Komponentenelements in der Aluminiumlegierung bei 3 Massen-% oder weniger gehalten wird.

4. Verfahren nach einem der Ansprüche 1 bis 3, das ferner einen Warmwalzschrift des Verminderns der Dicke der gegossenen Aluminiumlegierung, die durch den Gießschritt erhalten worden ist, um 30 % oder mehr durch Durchführen eines Warmwalzens der gegossenen Aluminiumlegierung bei einer Temperatur von 200 °C oder mehr umfasst.

5. Verfahren nach Anspruch 4, **dadurch gekennzeichnet, dass** der Warmwalzschrift nach dem Gießen des geschmolzenen Metalls in die Gießform zu einer plattenartigen Form durchgeführt wird, während in dem Gießschritt das geschmolzene Produkt auf eine Temperatur von 200 °C bis 500 °C abgekühlt und aus der Gießform entnommen wird.

6. Verfahren nach einem der Ansprüche 1 bis 3, das ferner einen Kaltwalz-Erwärmungsschrift des Durchführens eines Kaltwalzens der gegossenen Aluminiumlegierung, die durch den Gießschritt erhalten worden ist, wodurch die Dicke der gegossenen Aluminiumlegierung um 30 % oder mehr vermindert wird, und des anschließenden Erwärms bei einer Temperatur von 1/2 oder mehr des Schmelzpunkts der Aluminiumlegierung und 550 °C oder weniger umfasst.

7. Verfahren nach einem der Ansprüche 1 bis 3, das ferner einen Wärmebehandlungsschrift des Erwärms der gegossenen Aluminiumlegierung, die durch den Gießschritt erhalten worden ist, bei einer Temperatur von 400 °C oder mehr für 0,5 bis 3 Stunden umfasst.

8. Gegossene Aluminiumlegierung, **dadurch gekennzeichnet, dass** sie aus 0,8 bis 5 Massen-% Fe als ein erstes Komponentenelement, 0,15 bis 1 Massen-% Ti als ein zweites Komponentenelement, einer oder mehreren Art(en) von dritten Komponentenelementen, ausgewählt aus einer Gruppe von dritten Komponentenelementen, bestehend aus Zr, Nb, Hf, Sc und Y in einem Gehalt von 0,05 bis 2 Massen-% von jeder der dritten Komponenten, und in einer Menge, die $Fe > X > Ti$ genügt, wenn der Gehalt des ersten Komponentenelements Fe Fe (Massen-%) ist, der Gehalt des zweiten Komponentenelements Ti Ti (Massen-%) ist und der Gesamtgehalt der dritten Komponentenelemente X (Massen-%) ist, gegebenenfalls 0,05 bis 2 Massen-% Mg als ein viertes Komponentenelement, gegebenenfalls 0,05 bis 1 Massen-% von mindestens einem fünften Komponentenelement, ausgewählt aus einer Gruppe von fünften Komponentenelementen, bestehend aus Cu, Cr und Co, und gegebenenfalls V und/oder Mo als ein sechstes Komponentenelement in einer Menge von mehr als 0,05 Massen-% und weniger als 0,5 Massen-% besteht,

wobei es sich bei dem Rest um Al und unvermeidbare Verunreinigungen handelt, wobei die gegossene Aluminiumlegierung eine Metallstruktur aufweist, die eine α -Phase, die aus einer Al-Matrix ausgebildet ist, und eine Schichtphase umfasst, die aus einer eutektischen Struktur aus der Al-Matrix und einer Al-Fe-Verbindung derart ausgebildet ist, dass sie die α -Phase umgibt,

die Al-Matrix aus einer übersättigten festen Lösung von Al ausgebildet ist, in der das zweite Komponentenelement und das dritte Komponentenelement gelöst sind, und

der Flächenanteil eines Kristallisats, das aus einer intermetallischen Verbindung von Al, dem zweiten Komponentenelement und dem dritten Komponentenelement ausgebildet ist und einen Teilchendurchmesser von 5 μm oder mehr aufweist, in einem beliebigen Querschnitt der gegossenen Aluminiumlegierung weniger als 5 % beträgt.

9. Aluminiumlegierungsmaterial, **dadurch gekennzeichnet, dass** es aus 0,8 bis 5 Massen-% Fe als ein erstes Komponentenelement, 0,15 bis 1 Massen-% Ti als ein zweites Komponentenelement, einer oder mehreren Arten von dritten Komponentenelementen, ausgewählt aus einer Gruppe von dritten Komponentenelementen, bestehend aus Zr, Nb, Hf, Sc und Y in einem Gehalt von 0,05 bis 2 Massen-% von jeder der dritten Komponenten, und in einer Menge, die $\text{Fe} > \text{X} > \text{Ti}$ genügt, wenn der Gehalt des ersten Komponentenelements Fe (Massen-%) ist, der Gehalt des zweiten Komponentenelements Ti (Massen-%) ist und der Gesamtgehalt der dritten Komponentenelemente X (Massen-%) ist, gegebenenfalls 0,05 bis 2 Massen-% Mg als ein viertes Komponentenelement, gegebenenfalls 0,05 bis 1 Massen-% von mindestens einem fünften Komponentenelement, ausgewählt aus einer Gruppe von fünften Komponentenelementen, bestehend aus Cu, Cr und Co, und gegebenenfalls V und/oder Mo als ein sechstes Komponentenelement in einer Menge von mehr als 0,05 Massen-% und weniger als 0,5 Massen-% besteht, wobei es sich bei dem Rest um Al und unvermeidbare Verunreinigungen handelt, wobei das Aluminiumlegierungsmaterial eine Metallstruktur aufweist, die eine α -Phase, die aus einer Al-Matrix ausgebildet ist, und eine Schichtphase umfasst, die aus einer eutektischen Struktur aus der Al-Matrix und einer Al-Fe-Verbindung derart ausgebildet ist, dass sie die α -Phase umgibt, die Al-Matrix aus Al und/oder einer übersättigten festen Lösung von Al ausgebildet ist, in der das zweite Komponentenelement und das dritte Komponentenelement gelöst sind, und eine Ausscheidung, die aus einer intermetallischen Verbindung von Al, dem zweiten Komponentenelement und dem dritten Komponentenelement ausgebildet ist und einen Teilchendurchmesser von 2 bis 500 nm aufweist, in der Al-Matrix dispergiert ist.
10. Gegossene Aluminiumlegierung nach Anspruch 8 oder Aluminiumlegierungsmaterial nach Anspruch 9, **dadurch gekennzeichnet, dass** sie oder es von der Gruppe der dritten Komponentenelemente 0,2 bis 1,2 Massen-% Zr umfasst.
11. Gegossene Aluminiumlegierung nach Anspruch 8 oder 10 oder Aluminiumlegierungsmaterial nach Anspruch 9 oder 10, **dadurch gekennzeichnet, dass** Mg als viertes Komponentenelement mindestens in der Al-Matrix gelöst ist und/oder eine Al-Mg-Verbindung in der Al-Matrix bildet.
12. Gegossene Aluminiumlegierung nach einem der Ansprüche 8, 10 und 11 oder Aluminiumlegierungsmaterial nach einem der Ansprüche 9 bis 11, **dadurch gekennzeichnet, dass** Cr und/oder Co als fünftes Komponentenelement mindestens einen Teil der Al-Fe-Verbindung ersetzt, welche die Schichtphase bildet, so dass eine Al-(Fe, Cr)-Verbindung und/oder eine Al-(Fe, Co)-Verbindung gebildet wird.
13. Gegossene Aluminiumlegierung nach einem der Ansprüche 8 und 10 bis 12 oder Aluminiumlegierungsmaterial nach einem der Ansprüche 9 bis 12, **dadurch gekennzeichnet, dass** Cu als fünftes Komponentenelement eine Al-Cu-Verbindung in der Al-Matrix bildet.
14. Gegossene Aluminiumlegierung nach Anspruch 8 oder 10 oder Aluminiumlegierungsmaterial nach Anspruch 9 oder 10, **dadurch gekennzeichnet, dass** das Aluminiumlegierungsmaterial 0,05 bis 2 Massen-% Mg als viertes Komponentenelement und 0,05 bis 1 Massen-% Cu als fünftes Komponentenelement umfasst und dass das vierte Komponentenelement und das fünfte Komponentenelement eine Al-Cu-Mg-Verbindung in der Al-Matrix bilden.
15. Gegossene Aluminiumlegierung nach einem der Ansprüche 8 und 10 bis 14 oder Aluminiumlegierungsmaterial nach einem der Ansprüche 9 bis 14, **dadurch gekennzeichnet, dass** V und/oder Mo als sechstes Komponentenelement eine Verbindung mit Al sowie eine Verbindung mit Al, dem zweiten Komponentenelement und dem dritten Komponentenelement in der Al-Matrix bildet oder bilden.

Revendications

1. Procédé de production d'un alliage d'aluminium coulé obtenu par coulage d'un métal fondu d'un alliage d'aluminium, **caractérisé en ce qu'il** comprend :

une étape de fusion consistant à préparer un alliage d'aluminium constitué de 0,8 à 5 % en masse de Fe en tant que premier élément constitutif, 0,15 à 1 % en masse de Ti en tant que deuxième élément constitutif, une ou plusieurs sortes de troisièmes éléments constitutifs choisis parmi un groupe de troisièmes éléments constitutifs constitué de Zr, Nb, Hf, Sc et Y à une teneur de 0,05 à 2 % en masse de chacun des troisièmes composants et dans une quantité qui satisfait $Fe > X > Ti$ lorsque la teneur du premier élément constitutif Fe est Fe (% en masse), la teneur du deuxième élément constitutif Ti est Ti (% en masse), et une teneur totale du troisième élément constitutif est X (% en masse) ; facultativement 0,05 à 2 % en masse de Mg en tant que quatrième élément constitutif, facultativement 0,05 à 1 % en masse d'au moins un cinquième élément constitutif choisi parmi un groupe de cinquièmes éléments constitutifs constitué de Cu, Cr et Co, et facultativement V et/ou Mo en tant que sixième élément constitutif dans une quantité de plus de 0,05 % en masse et de moins de 0,5 % en masse, la partie résiduelle étant Al et des impuretés inévitables, et faire fondre l'alliage d'aluminium à une température qui est supérieure de 20°C ou plus à une température de liquidus déterminée par la composition de l'alliage d'aluminium pour obtenir un métal fondu ; et

une étape de coulage consistant à couler le métal fondu dans un moule de coulage puis à refroidir le métal fondu à une température qui est inférieure d'au moins 10°C à une température de solidus déterminée par la composition de l'alliage d'aluminium à une vitesse de refroidissement de 150°C/s ou plus et inférieure à 10 000°C/s.

2. Procédé selon la revendication 1, **caractérisé par** l'utilisation de l'alliage d'aluminium comprenant 0,2 à 1,2 % en masse de Zr parmi le groupe de troisièmes éléments constitutifs à l'étape de fusion.

3. Procédé selon la revendication 1 ou 2, **caractérisé en ce qu'une** quantité totale du quatrième élément constitutif, du cinquième élément constitutif et du sixième élément constitutif dans l'alliage d'aluminium est maintenue à 3 % en masse ou moins.

4. Procédé selon l'une quelconque des revendications 1 à 3, comprenant en outre une étape de laminage à chaud consistant à réduire l'épaisseur de l'alliage d'aluminium coulé obtenu par l'étape de coulage de 30 % ou plus en réalisant un laminage à chaud sur l'alliage d'aluminium coulé à une température de 200°C ou plus.

5. Procédé selon la revendication 4, **caractérisé en ce que** l'étape de laminage à chaud est réalisée après coulage du métal fondu en une forme de type plaque dans le moule de coulage tout en refroidissant le produit fondu à une température de 200°C à 500°C et libération du moule de coulage dans l'étape de coulage.

6. Procédé selon l'une quelconque des revendications 1 à 3, comprenant en outre une étape de chauffage-laminage à froid consistant à réaliser un laminage à froid sur l'alliage d'aluminium coulé obtenu par l'étape de coulage afin de réduire ainsi l'épaisseur de l'alliage d'aluminium coulé de 30 % ou plus, et ensuite à chauffer à une température de 1/2 ou plus du point de fusion de l'alliage d'aluminium et 550°C ou moins.

7. Procédé selon l'une quelconque des revendications 1 à 3, comprenant en outre une étape de traitement à la chaleur consistant à chauffer l'alliage d'aluminium coulé obtenu par l'étape de coulage à une température de 400°C ou plus pendant 0,5 à 3 heures.

8. Alliage d'aluminium coulé **caractérisé en ce qu'il** est constitué de 0,8 à 5 % en masse de Fe en tant que premier élément constitutif, 0,15 à 1 % en masse de Ti en tant que deuxième élément constitutif, une ou plusieurs sortes de troisièmes éléments constitutifs choisis parmi un groupe de troisièmes éléments constitutifs constitué de Zr, Nb, Hf, Sc et Y à une teneur de 0,05 à 2 % en masse de chacun des troisièmes composants et dans une quantité qui satisfait $Fe > X > Ti$ lorsque la teneur du premier élément constitutif Fe est Fe (% en masse), la teneur du deuxième élément constitutif Ti est Ti (% en masse), et une teneur totale du troisième élément constitutif est X (% en masse) ; facultativement 0,05 à 2 % en masse de Mg en tant que quatrième élément constitutif, facultativement 0,05 à 1 % en masse d'au moins un cinquième élément constitutif choisi parmi un groupe de cinquièmes éléments constitutifs constitué de Cu, Cr et Co, et facultativement V et/ou Mo en tant que sixième élément constitutif dans une quantité de plus de 0,05 % en masse et de moins de 0,5 % en masse,

la partie résiduelle étant Al et des impuretés inévitables,
dans lequel l'alliage d'aluminium coulé a une structure métallique comportant une phase α qui est formée d'une
matrice d'Al et d'une phase en couches qui est formée d'une structure eutectique de la matrice d'Al et d'un composé
Al-Fe de façon à entourer la phase α ;

la matrice d'Al est formée d'une solution solide sursaturée d'Al dans laquelle le deuxième élément constitutif et le
troisième élément constitutif sont dissous ; et

un rapport d'aire d'un cristallin formé d'un composé intermétallique d'Al, du deuxième élément constitutif et du
troisième élément constitutif et ayant un diamètre de particule de 5 μm ou plus est inférieur à 5 % dans une section
arbitraire de l'alliage d'aluminium coulé.

9. Matériau d'alliage d'aluminium **caractérisé en ce qu'il** constitué de
0,8 à 5 % en masse de Fe en tant que premier élément constitutif,
0,15 à 1 % en masse de Ti en tant que deuxième élément constitutif,
une ou plusieurs sortes de troisièmes éléments constitutifs choisis parmi un groupe de troisièmes éléments consti-
tutifs constitué de Zr, Nb, Hf, Sc et Y à une teneur de 0,05 à 2 % en masse de chacun des troisièmes composants
et dans une quantité qui satisfait $\text{Fe} > \text{X} > \text{Ti}$ lorsque la teneur du premier élément constitutif Fe est Fe (% en masse),
la teneur du deuxième élément constitutif Ti est Ti (% en masse), et une teneur totale du troisième élément constitutif
est X (% en masse) ; facultativement 0,05 à 2 % en masse de Mg en tant que quatrième élément constitutif,
facultativement 0,05 à 1 % en masse d'au moins un cinquième élément constitutif choisi parmi un groupe de
cinquièmes éléments constitutifs constitué de Cu, Cr et Co, et facultativement V et/ou Mo en tant que sixième
élément constitutif dans une quantité de plus de 0,05 % en masse et de moins de 0,5 % en masse, et

la partie résiduelle étant Al et des impuretés inévitables,
dans lequel le matériau d'alliage d'aluminium a une structure métallique comportant une phase α qui est formée
d'une matrice d'Al et d'une phase en couches qui est formée d'une structure eutectique de la matrice d'Al et d'un
composé Al-Fe de façon à entourer la phase α ;
la matrice d'Al est formée d'Al et/ou d'une solution solide sursaturée d'Al dans laquelle le deuxième élément constitutif
et le troisième élément constitutif sont dissous ; et
un précipité formé d'un composé intermétallique d'Al, du deuxième élément constitutif et du troisième élément
constitutif et ayant un diamètre de particule de 2 à 500 nm est dispersé dans la matrice d'Al.

10. Alliage d'aluminium coulé selon la revendication 8 ou matériau d'alliage d'aluminium selon la revendication 9,
caractérisé en ce qu'il comprend 0,2 à 1,2 % en masse de Zr parmi le groupe de troisièmes éléments constitutifs.

11. Alliage d'aluminium coulé selon la revendication 8 ou 10 ou matériau d'alliage d'aluminium selon la revendication
9 ou 10, **caractérisé en ce que** Mg en tant que quatrième élément constitutif est dissous au moins dans la matrice
d'Al et/ou forme un composé Al-Mg dans la matrice d'Al.

12. Alliage d'aluminium coulé selon l'une quelconque des revendications 8, 10 et 11 ou matériau d'alliage d'aluminium
selon la revendication selon l'une quelconque des revendications 9 à 11, **caractérisé en ce que** Cr et/ou Co en
tant que cinquième élément constitutif est remplacé par au moins une partie du composé Al-Fe formant la phase
en couches pour former un composé Al-(Fe, Cr) et/ou un composé Al-(Fe, Co).

13. Alliage d'aluminium coulé selon l'une quelconque des revendications 8, 10 à 12 ou matériau d'alliage d'aluminium
selon la revendication selon l'une quelconque des revendications 9 à 12, **caractérisé en ce que** Cu en tant que
cinquième élément constitutif forme un composé Al-Cu dans la matrice d'Al.

14. Alliage d'aluminium coulé selon la revendication 8 ou 10 ou matériau d'alliage d'aluminium selon la revendication
9 ou 10, **caractérisé en ce que** le matériau d'alliage d'aluminium comprend 0,05 à 2 % en masse de Mg en tant
que quatrième élément constitutif et 0,05 à 1 % en masse de Cu en tant que cinquième élément constitutif, et **en**
ce que le quatrième élément constitutif et le cinquième élément constitutif forment un composé Al-Cu-Mg dans la
matrice d'Al.

15. Alliage d'aluminium coulé selon l'une quelconque des revendications 8 et 10 à 14 ou matériau d'alliage d'aluminium
selon la revendication selon l'une quelconque des revendications 9 à 14, **caractérisé en ce que** V et/ou Mo en tant
que sixième élément constitutif forme(nt) un composé avec Al et un composé avec Al, le deuxième élément constitutif,
et le troisième élément constitutif dans la matrice d'Al.

FIG. 1

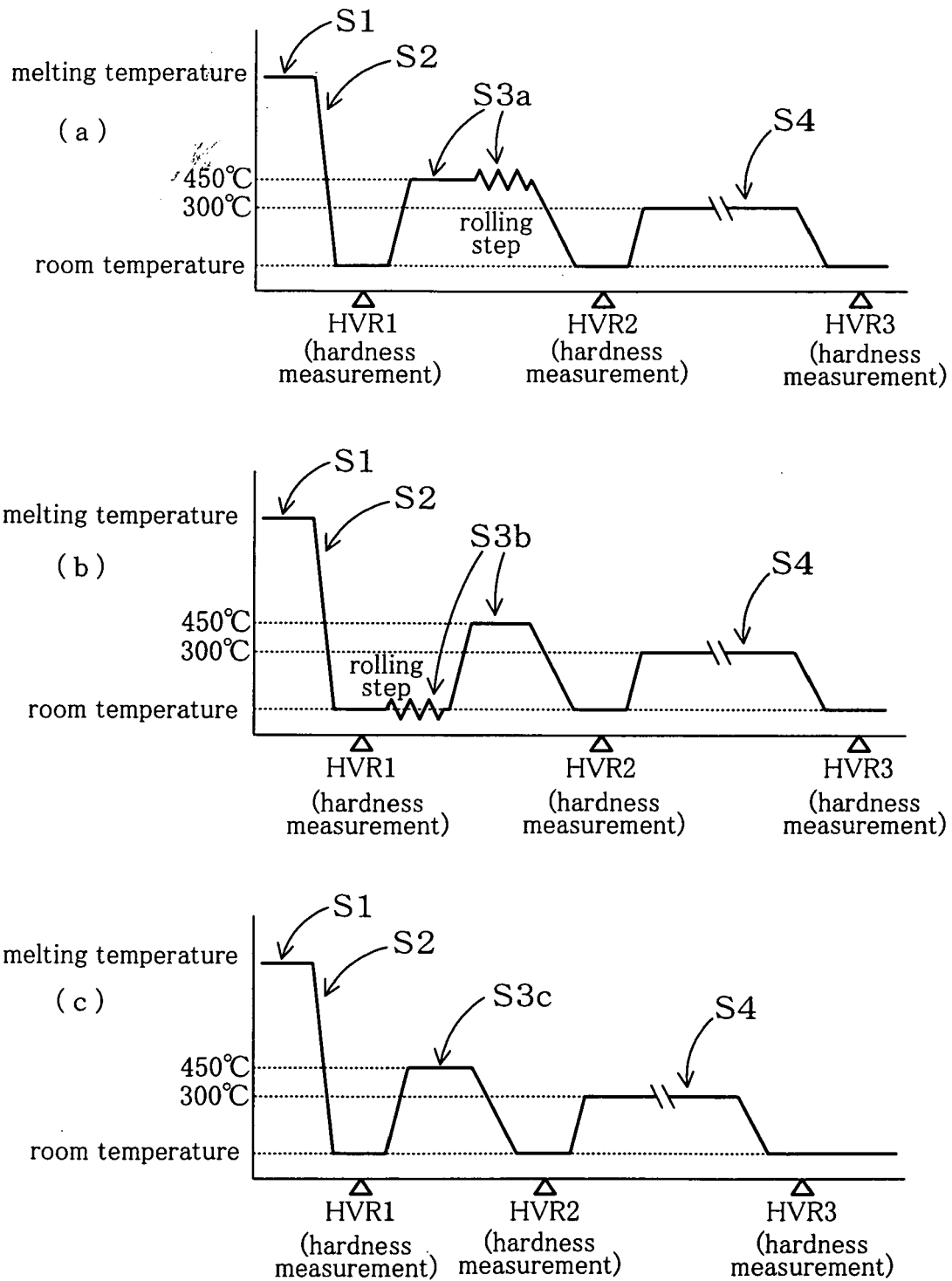


FIG. 2

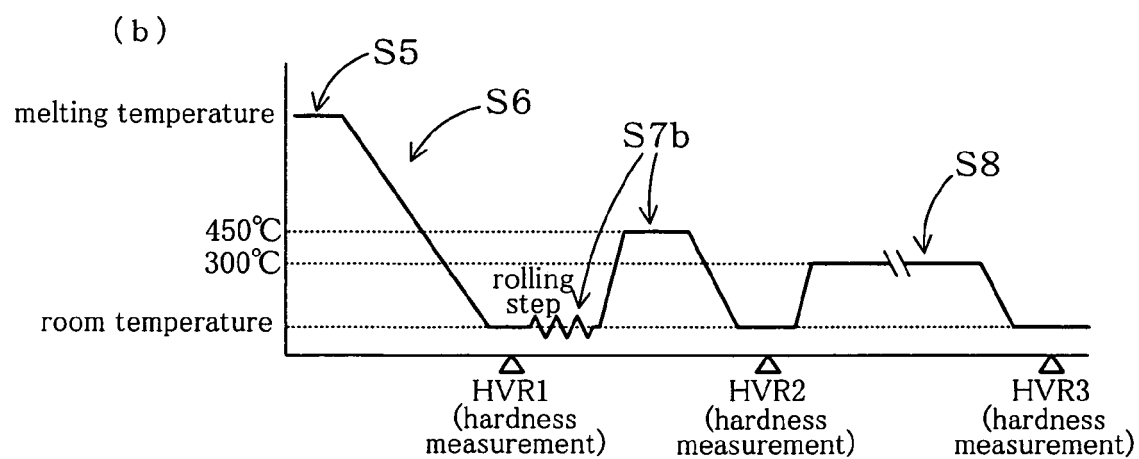
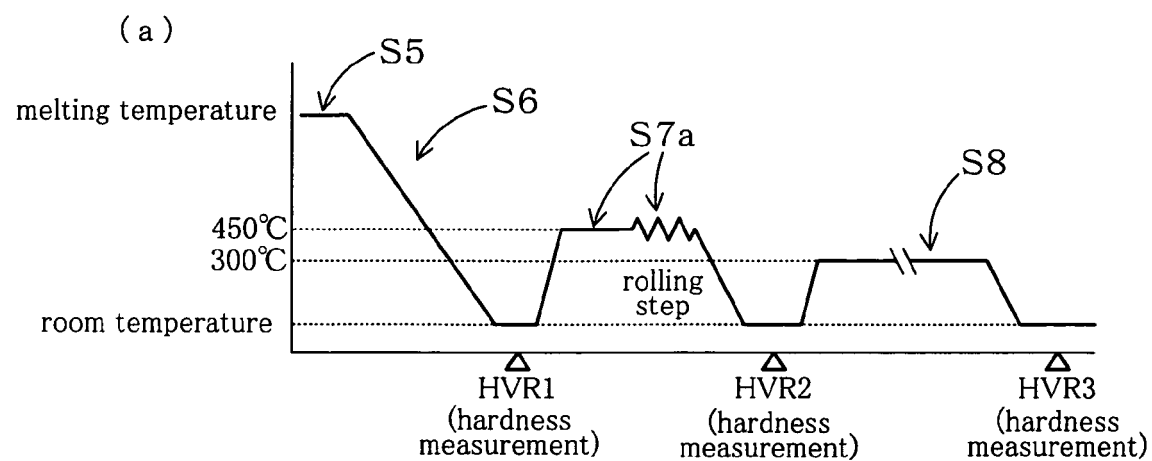
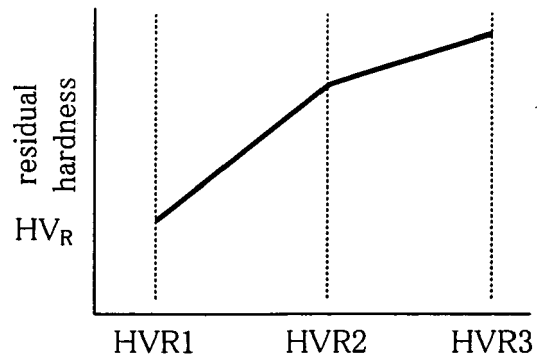
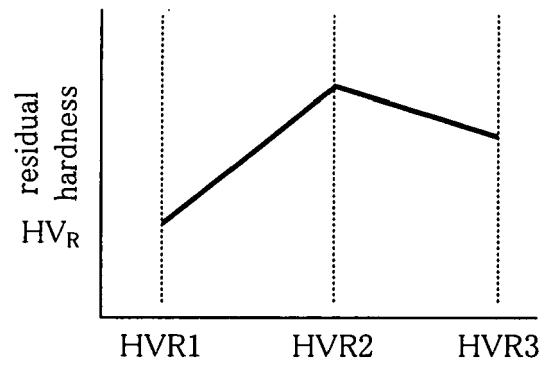


FIG. 3

(a)



(b)



(c)

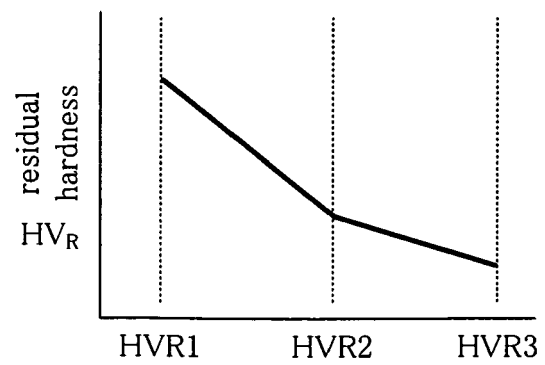


FIG. 4

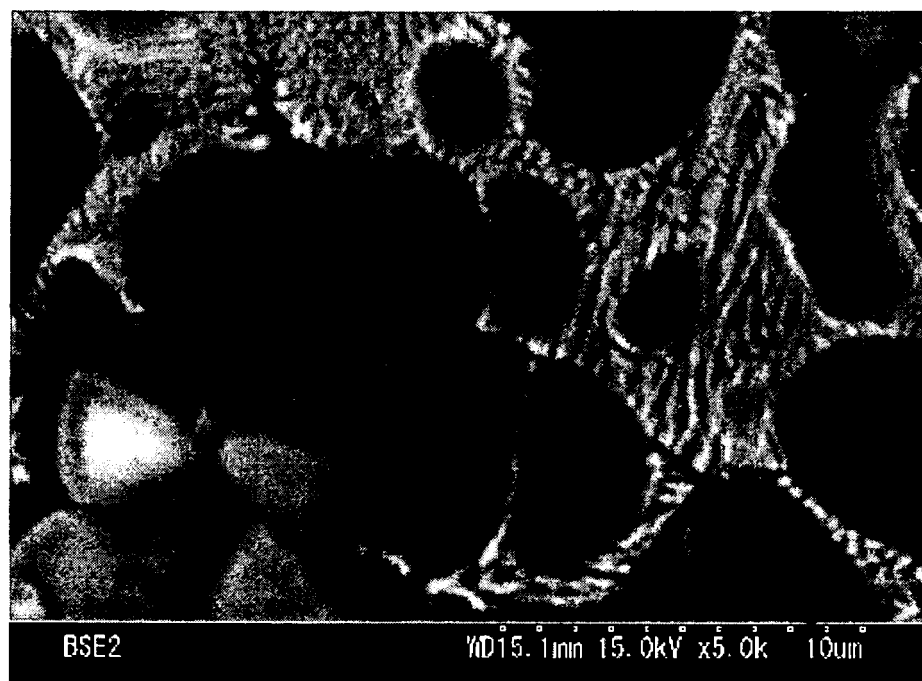


FIG. 5

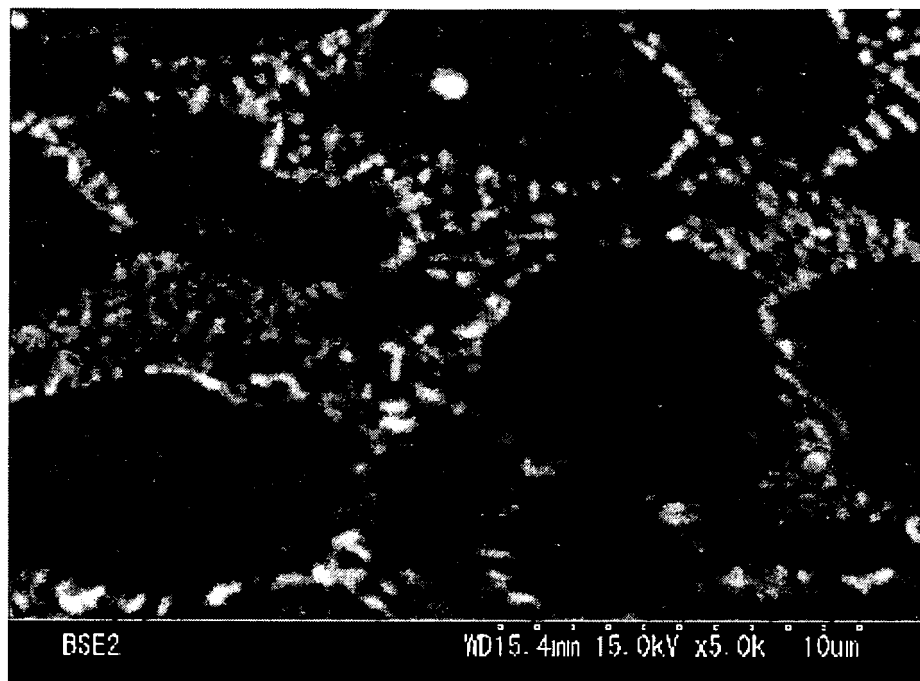


FIG. 6

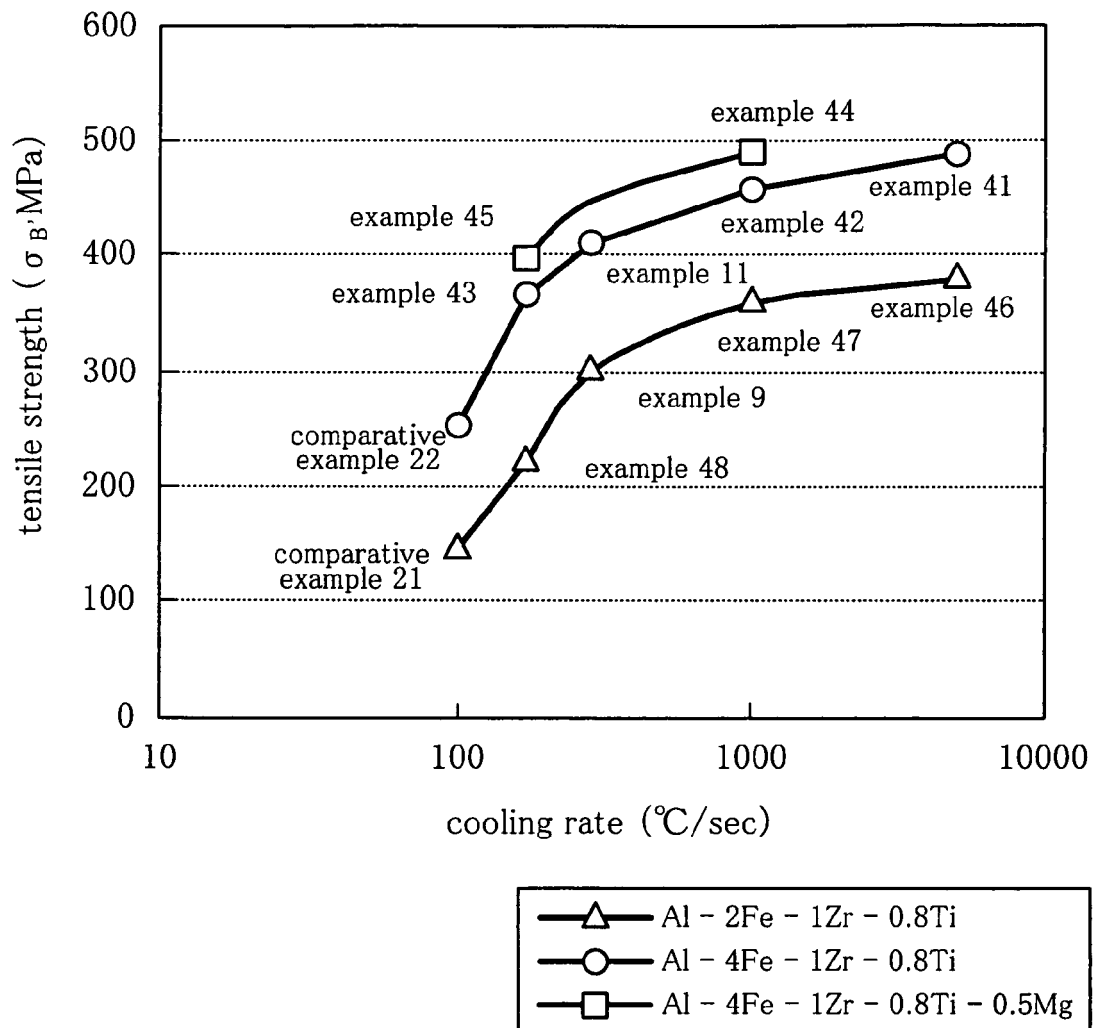


FIG. 7

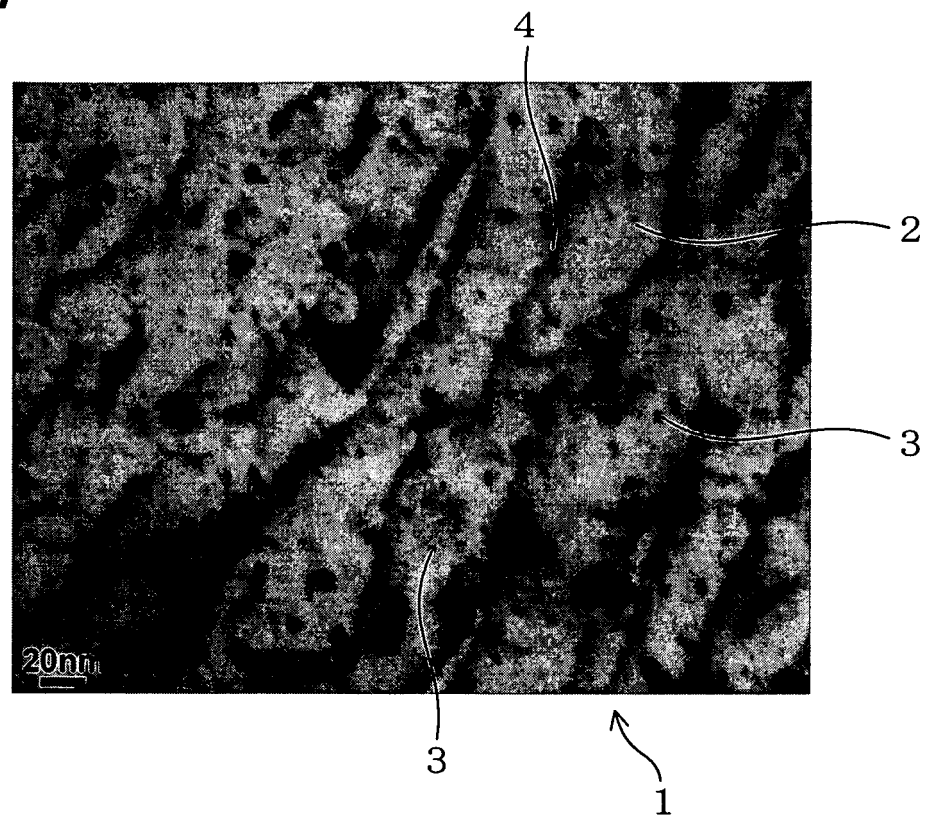


FIG. 8



FIG. 9

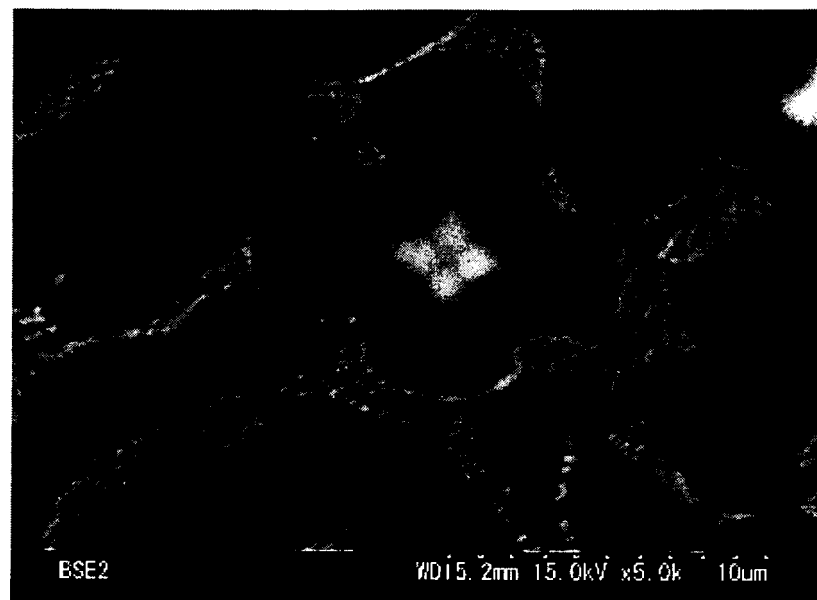


FIG. 10

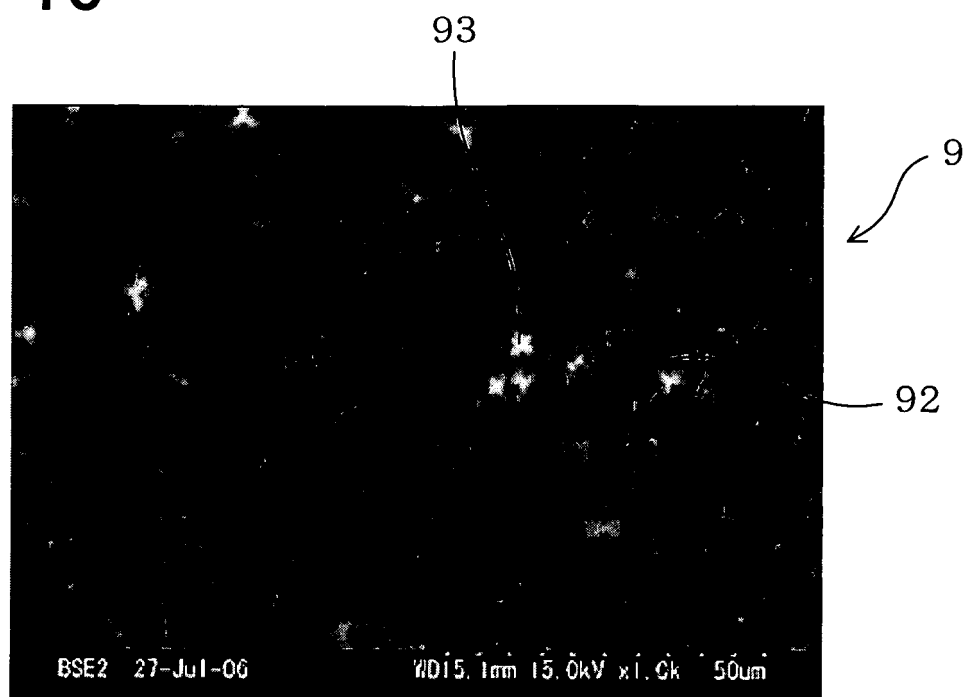


FIG. 11

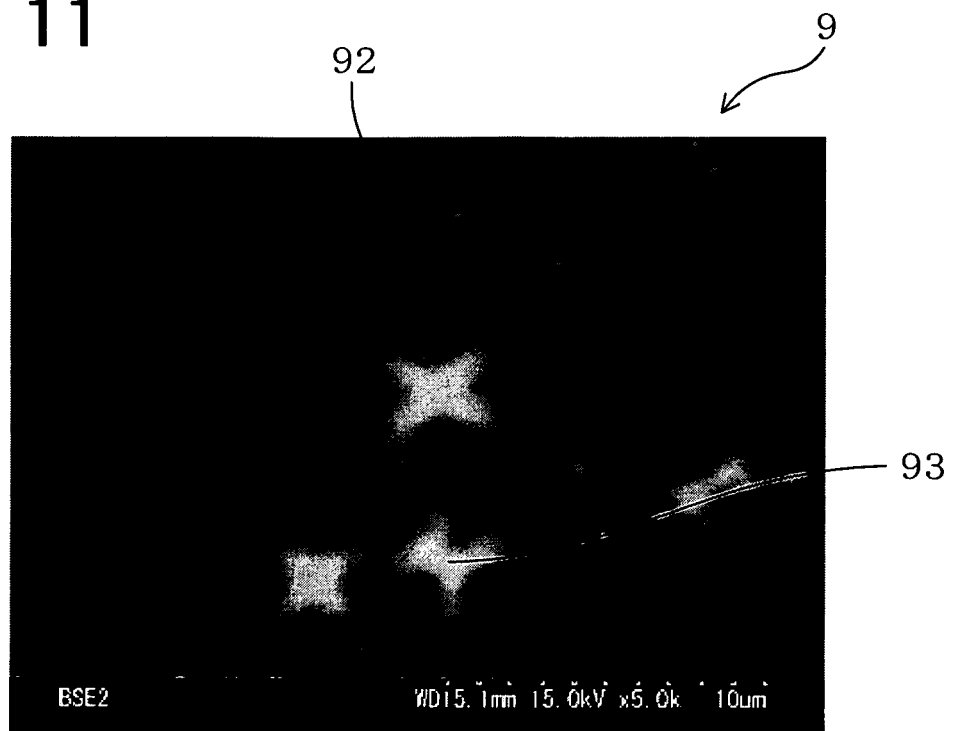


FIG. 12

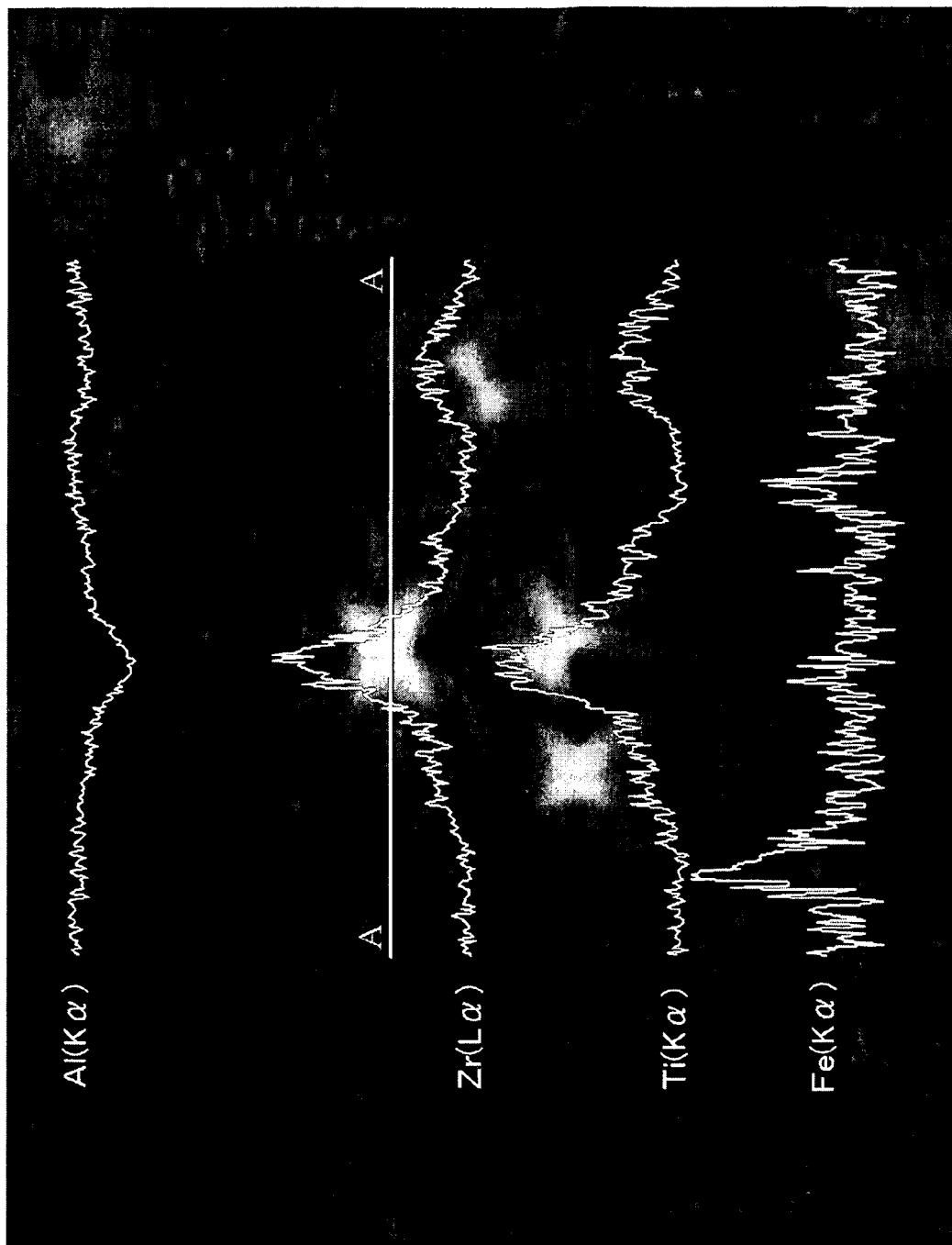


FIG. 13

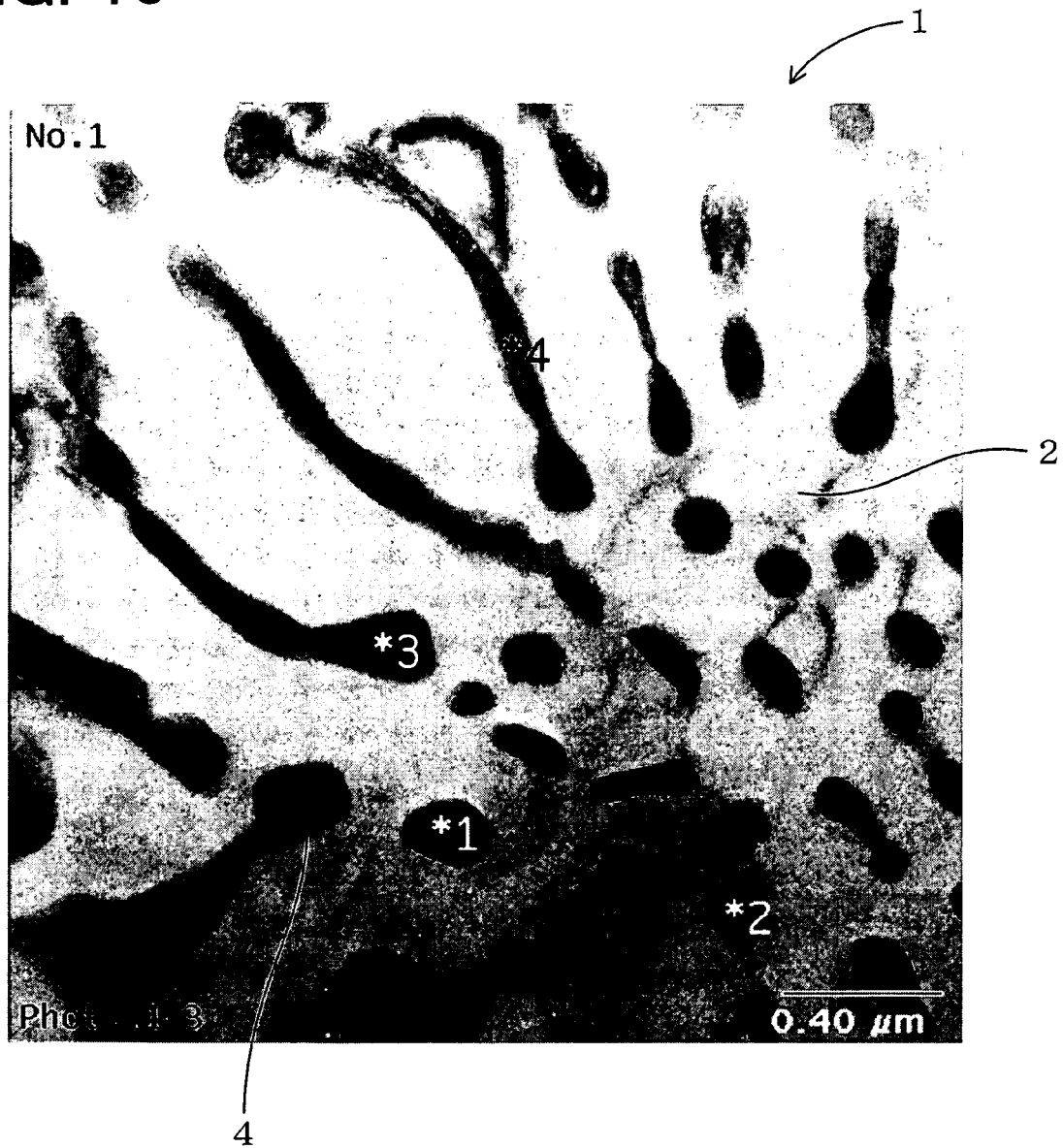


FIG. 14

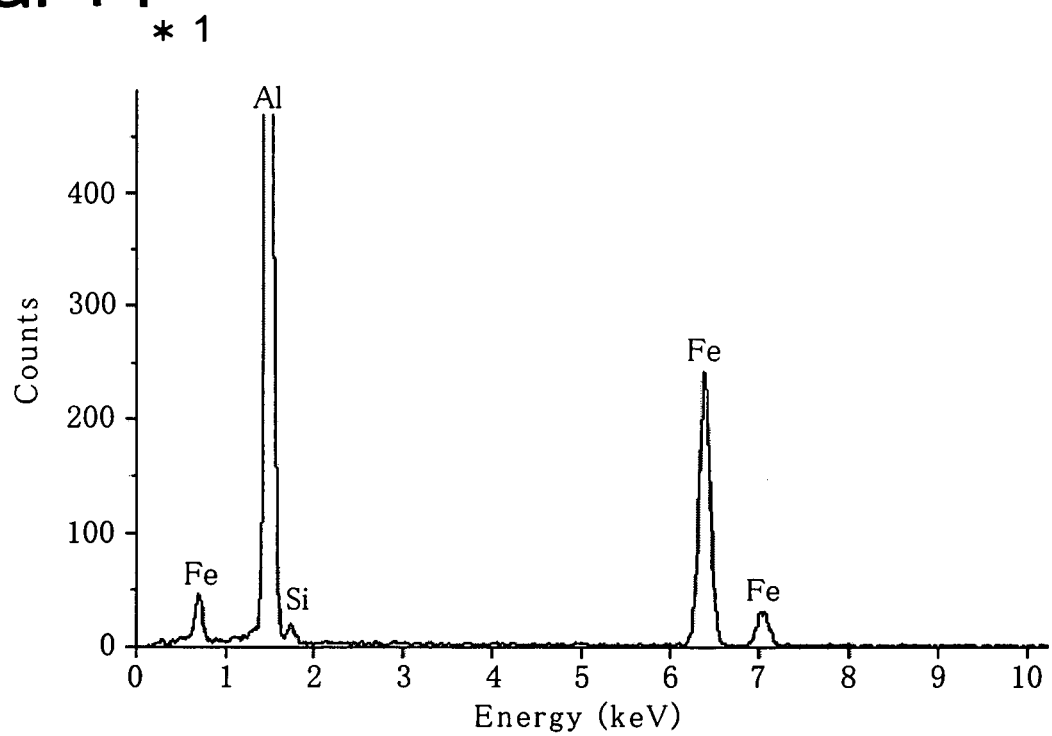


FIG. 15

* 2

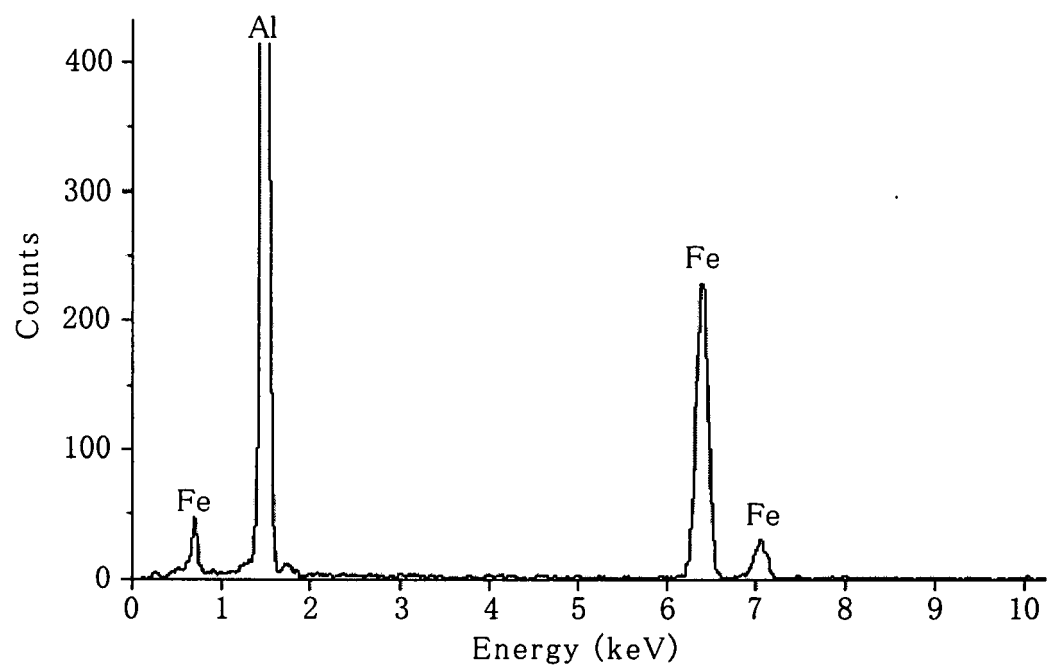


FIG. 16 * 3

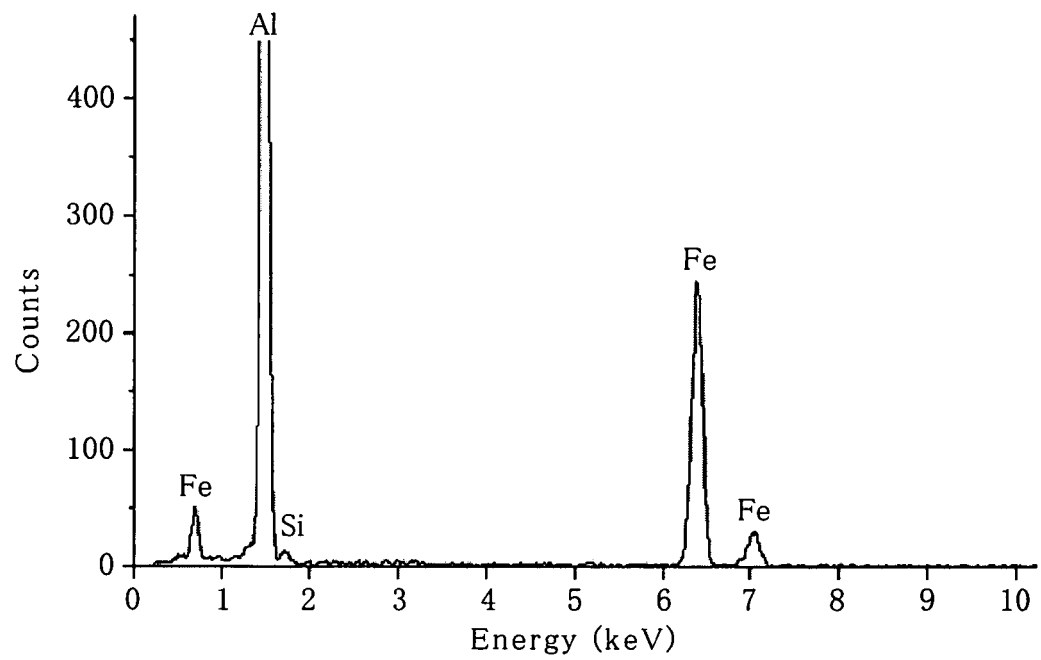


FIG. 17 * 4

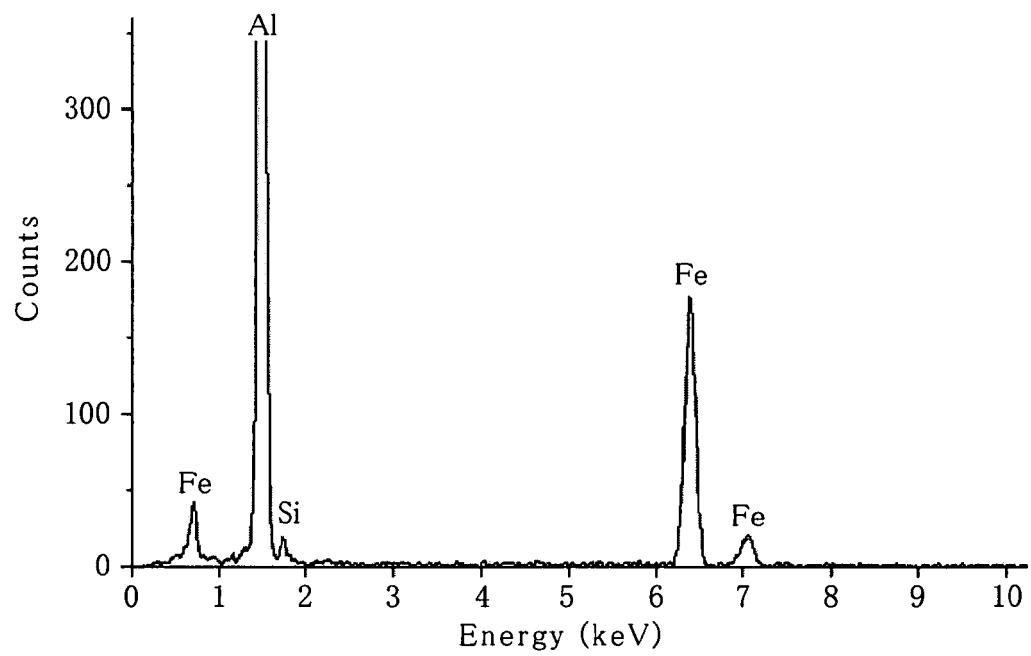


FIG. 18



FIG. 19 * 1

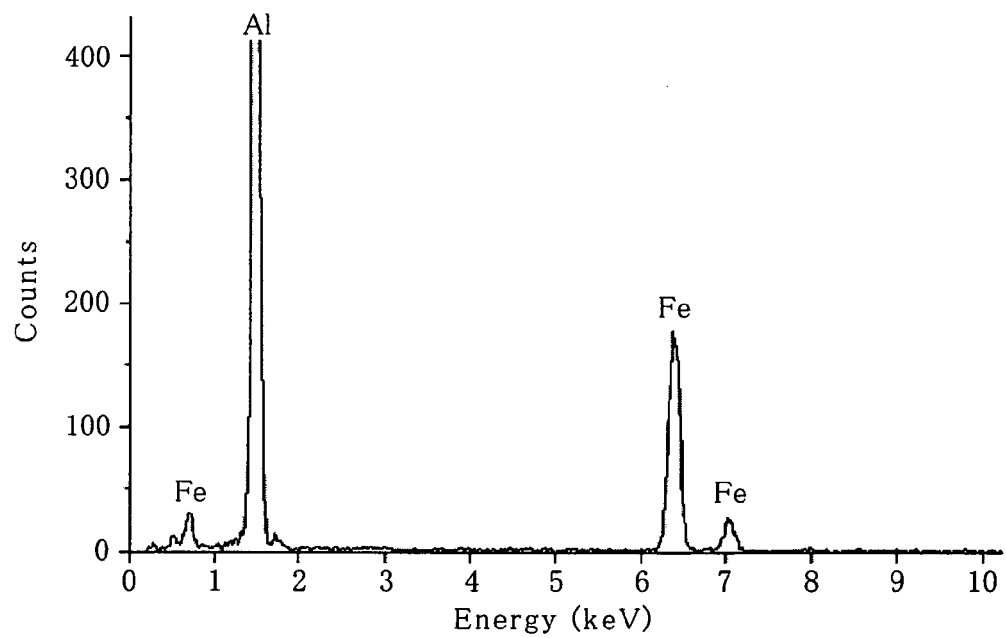


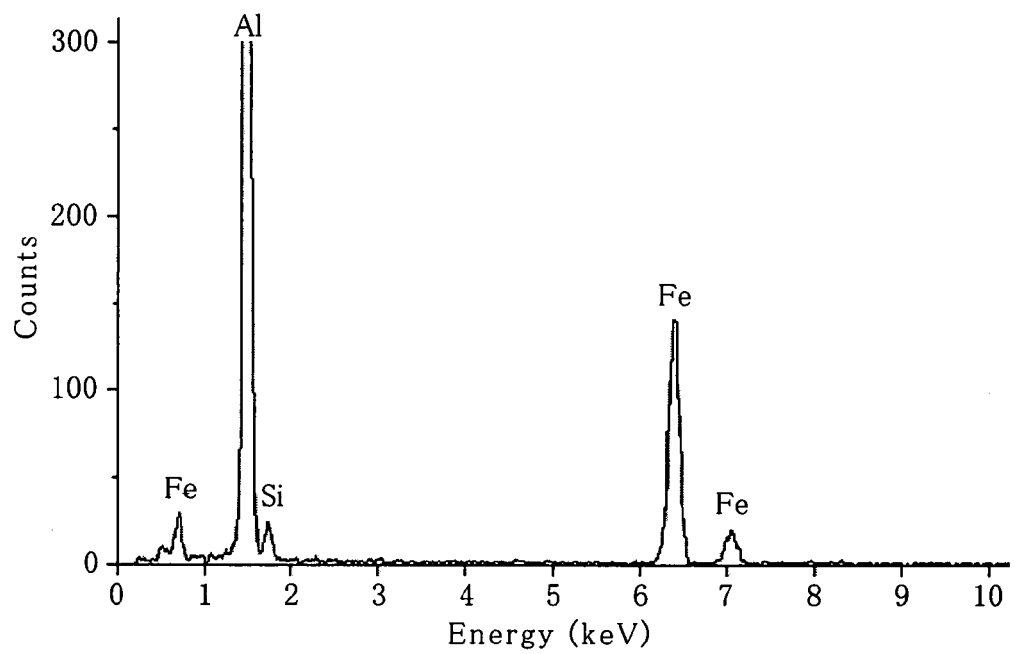
FIG. 20 * 2

FIG. 21 * 3

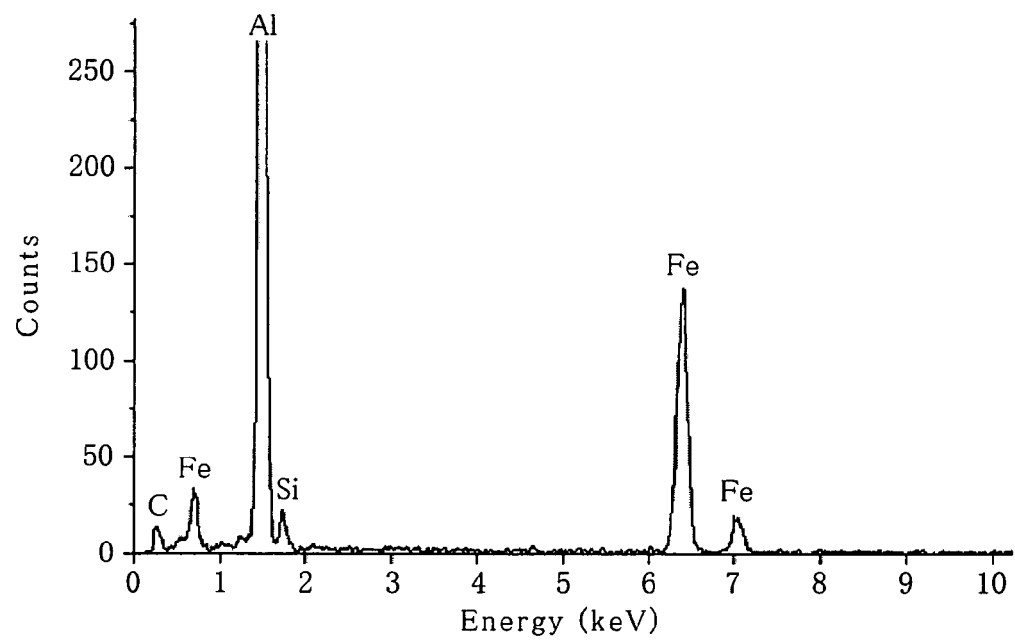


FIG. 22 * 4

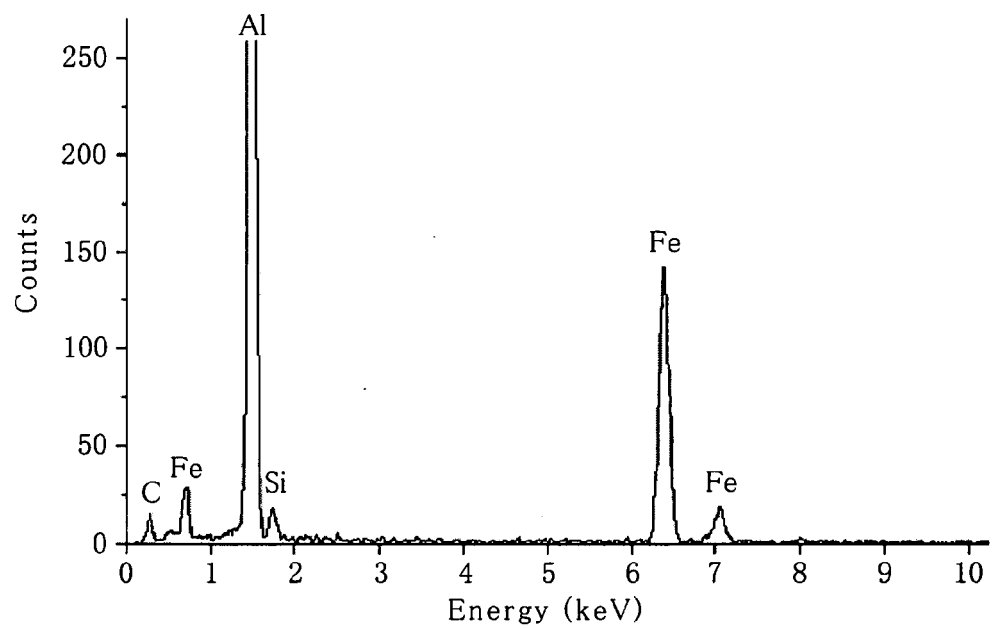


FIG. 23

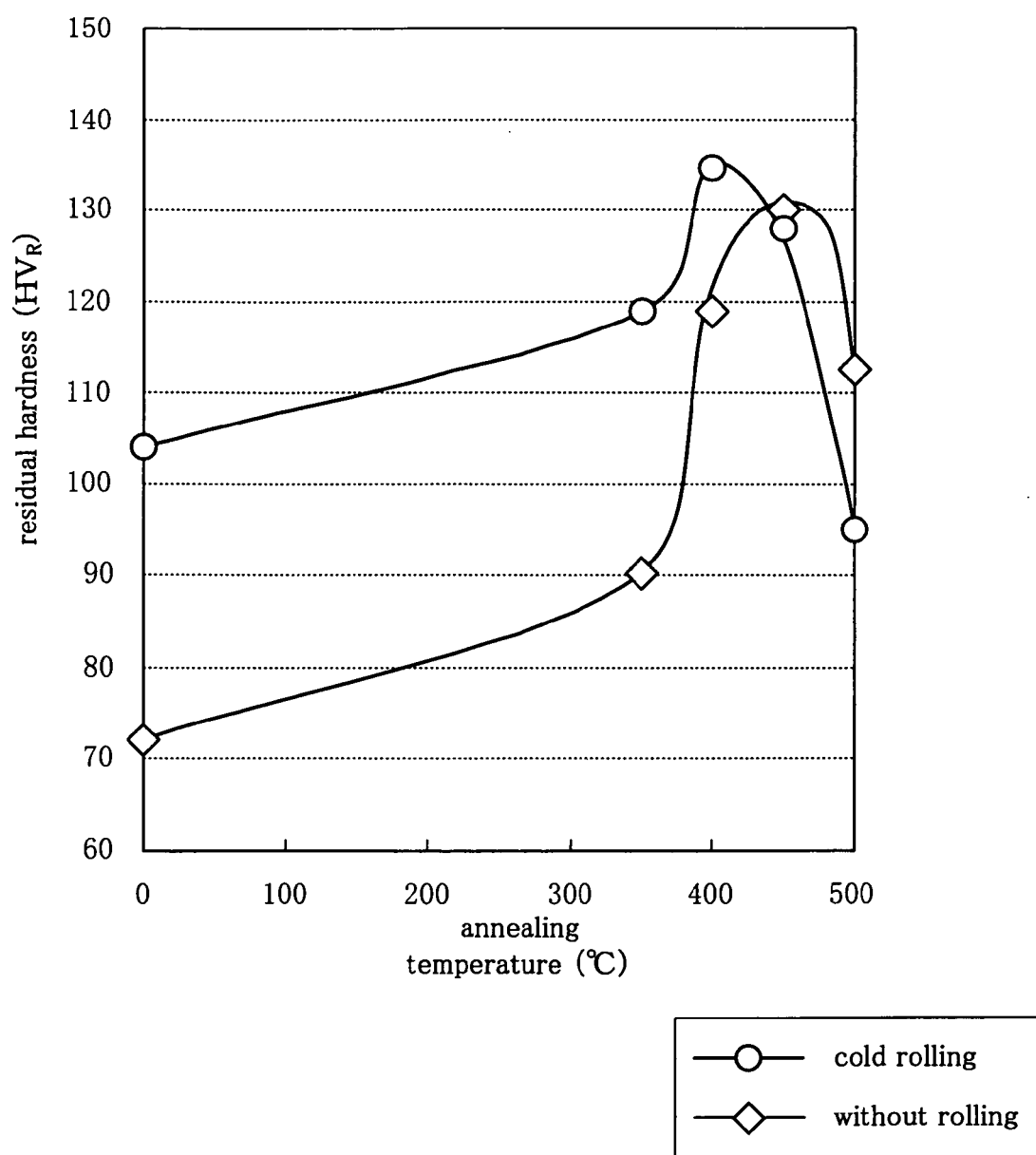


FIG. 24

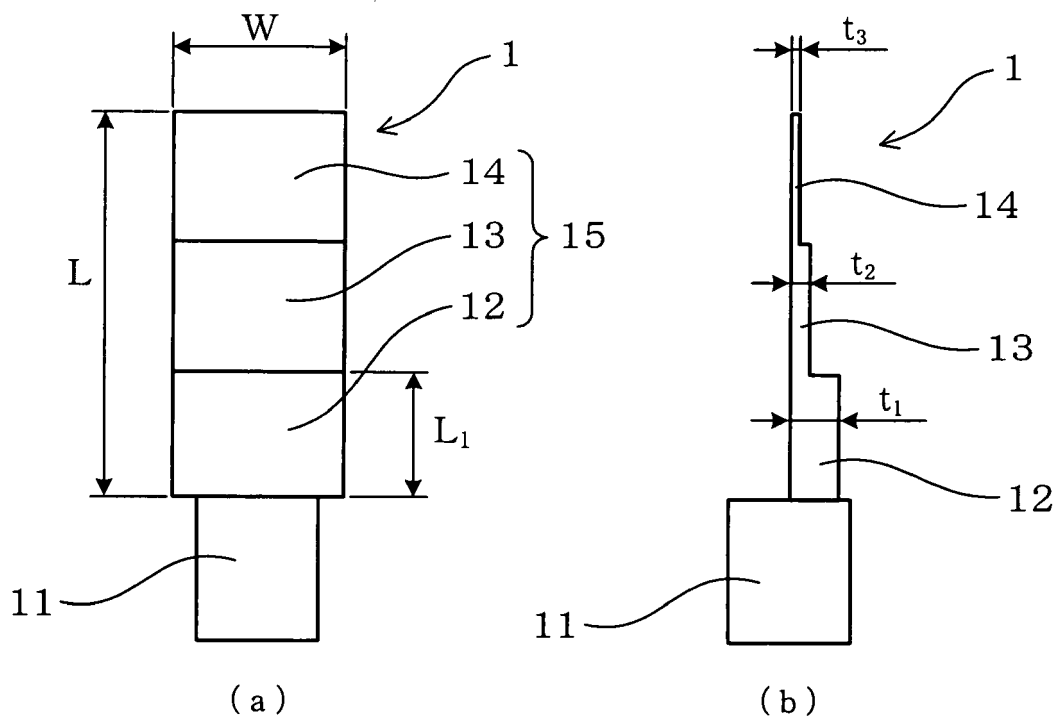
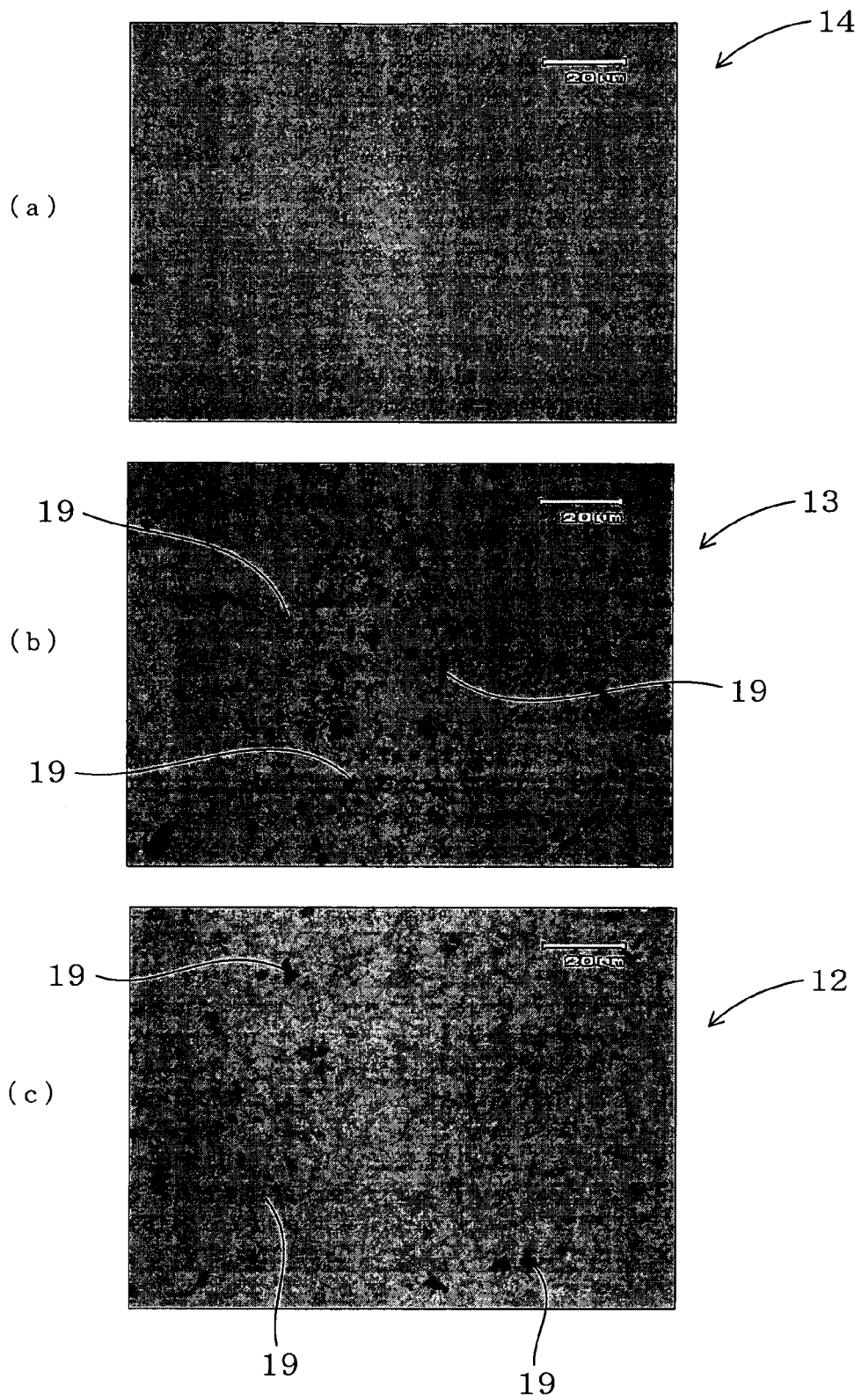


FIG. 25



REFERENCES CITED IN THE DESCRIPTION

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