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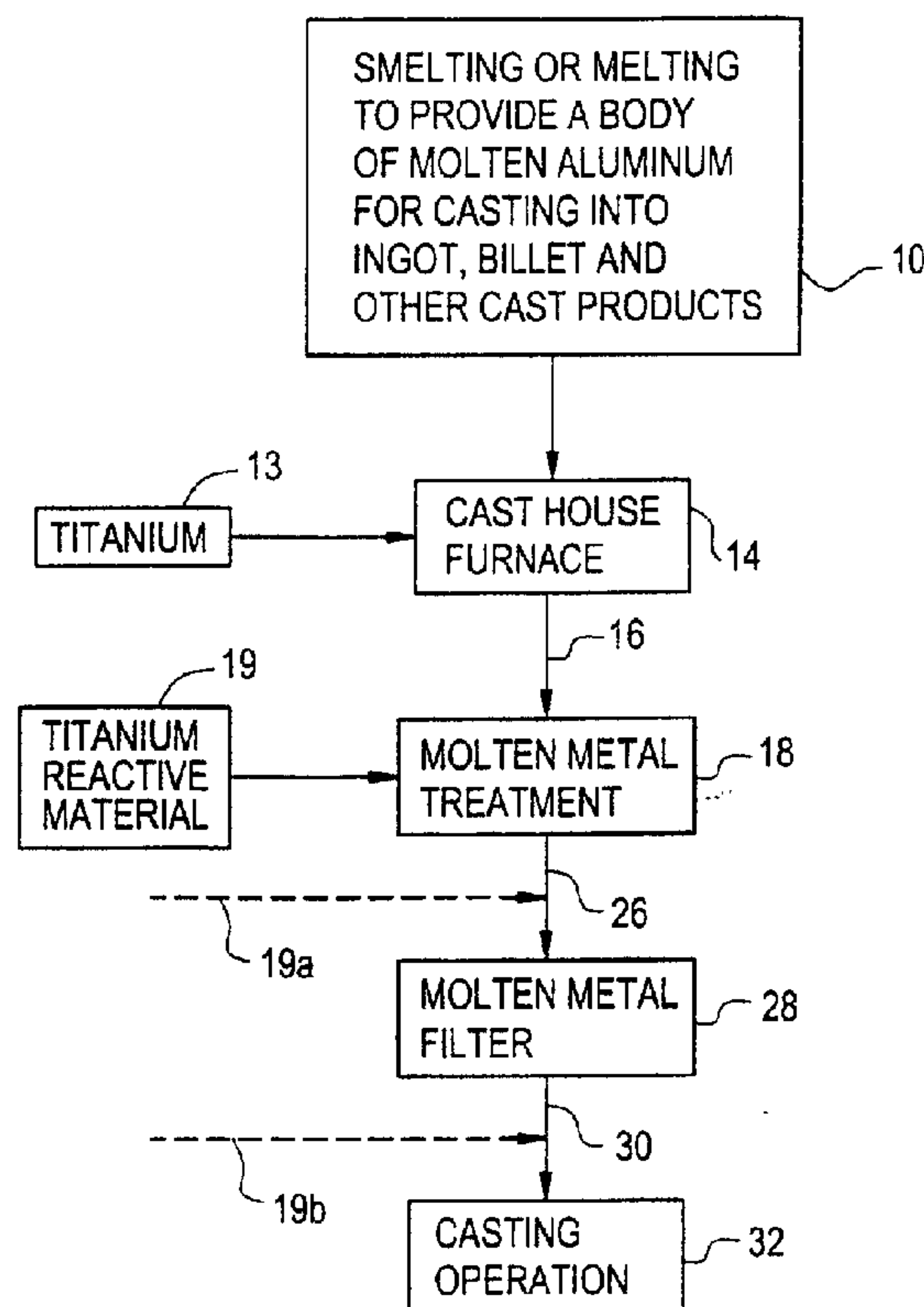
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(54) **TRAITEMENT DE L'ALUMINIUM FONDU**

(54) **MOLTEN ALUMINUM TREATMENT**



(57) Procédé de recuit d'affinage du grain destiné à l'aluminium, qui consiste à produire un corps d'aluminium fondu (10) contenant 1 à 3000 ppm de titane (13) et à y introduire un matériau (19) réagissant avec le titane, de préférence sous forme gazeuse. Ce matériau renferme un composant choisi dans le groupe comprenant le bore, le carbone, le soufre, l'azote et le phosphore. Il forme avec le titane un composé d'affinage du grain qui permet d'affiner le grain de l'aluminium.

(57) A method of grain refining aluminum, the method comprising providing a molten aluminum body (10) containing 1 to 3000 ppm titanium (13). A material (19) reactive with the titanium is introduced preferably in gaseous form to the aluminum body. The material has a component selected from the group consisting of boron, carbon, sulfur, nitrogen and phosphorus. The material and said titanium form a grain refining compound adapted for grain refining the aluminum.



Abstract

A method of grain refining aluminum, the method comprising providing a molten aluminum body containing 1 to 3000 ppm titanium. A material reactive with the titanium is introduced preferably in gaseous form to the aluminum body. The material has a component selected from the group consisting of boron, carbon, sulfur, nitrogen and phosphorus. The material and said titanium form a grain refining compound adapted for grain refining the aluminum.

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MOLTEN ALUMINUM TREATMENT

This invention relates to molten aluminum treatments and more particularly it relates to molten metal treatment for degassing and/or forming grain refining nuclei in molten aluminum in situ.

5 There is an ever increasing effort to improve aluminum and its alloys by the use of new grain refiners or master alloys comprising the grain refiners. Presently, the most popular grain refiners for aluminum utilize titanium diboride (TiB_2) type compound $[(\text{TiAl})\text{B}_2]$ or titanium carbide (TiC). Typically, the TiB_2 is produced by reacting K_2TiF_6 and KFB_4 salts with aluminum to produce the master
10 alloy with excess titanium. The master alloy is added to the molten aluminum to be refined prior to the casting operation and usually prior to filtration. This master alloy manufacturing process produces small particle sized TiB_2 and TiAl_3 , entrained liquid KAlF and Al_2O_3 . Sometimes the salts are added directly to the molten metal to be refined. When the master alloy is used, it is added to the melt as a waffle or
15 rod.

 However, the use of master alloys or addition of the salts directly to the metal is not without problems. For example, the master alloys containing TiB_2 often have salt and oxide inclusions, e.g., titanium and boron salts and aluminum oxides. Often the inclusions are larger than the TiB_2 particles. Further, the master
20 alloy also can contain TiB_2 clusters which, of course, again are larger than the individual TiB_2 particles. The TiB_2 clusters normally contain materials such as oxides and salts, e.g., KAlF_4 . The inclusions and clusters are detrimental because they are frequently the source of downstream processing problems in the cast or fabricated aluminum product. For example, the inclusions and clusters cause
25 increased wear on cutting, rolling or die surfaces used to process the cast aluminum product. The inclusions and clusters are a source of defects such as holes and stress points in the metal. Further, the inclusions and clusters are detrimental because of filter clogging just prior to the casting operation, adding an additional expense in filter replacement.

30 Another very effective grain refiner that is being increasingly used with aluminum is $\text{Al-3\% Ti-0.15C}$. Master alloys containing TiC are often prepared by heating mixtures of aluminum, titanium and carbon. However, this method has

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the problem that the process temperature is quite high, e.g., 1200° to 1300°C. Further, the process only makes a dilute master alloy, i.e. a master alloy that is dilute in grain refining nuclei, e.g., Al-3% Ti-0.15C.

In addition, the use of master alloys has the problem that they
5 provide localized, high concentrations of refiner which can result in larger clusters of particles, residual slag (KAlF_4), oxides, etc., and inoperative nuclei as well as problems dispersing the particles and dissolving the aluminides. Even though master alloys are presently used throughout the industry, they are an inefficient use of the refiner components. The high concentrations of refiner referred to are even
10 more pronounced in foundry situations where cast waffles of master alloys are added to the aluminum melt.

The use of the term "aluminum" as used herein is meant to include aluminum and its alloys.

Prior attempts at improving grain refining have focused on improving
15 the master alloy. For example U.S. Patent 5,415,708 discloses an aluminum base alloy consisting essentially of from 0.1 to 3.0% boron, from 1 to 10% titanium and the balance essentially aluminum wherein the aluminum matrix contains TiB_2 particles dispersed throughout said matrix having an average particle size of less than 1 micron, and wherein the matrix contains clusters of said TiB_2 particles
20 greater than 10 microns in size with an average of less than 4 of said clusters per 2 cm^2 . The alloy is prepared by adding a boron containing material selected from the group consisting of borax, boron oxide, boric acid and mixtures thereof, and K_2TiF_6 to a bath of molten aluminum and stirring the molten mixture.

U.S. Patent 5,100,618 discloses a process for producing aluminum
25 grain refiner, such as Al-Ti-B grain refiner. Molten aluminum is continuously flowed as a bottom layer along a substantially horizontal or slightly inclined trough. Titanium or boron compounds reducible by aluminum or a mixture of such compounds is added to the surface of the aluminum layer such that a discrete separate layer of these is formed on top of the aluminum layer. Reaction between
30 the aluminum and the titanium and/or boron compounds occurs along the interface between the layers and this reaction may, if desired, be aided by providing relative movement between the layer of molten aluminum and the layer of titanium and/or

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boron compounds.

U.S. Patent 5,104,616 discloses a method for the production of master alloys intended for grain refining of aluminum melts and being of the type which comprises of aluminum and 1-15 percent by weight titanium, where titanium
5 is present in the form of intermetallic crystals of titanium aluminide in combination with additives of carbon and/or nitrogen. The method is characterized by adding carbon and/or nitrogen to the aluminum melt in an amount corresponding to at least 0.01 percent by weight in the resultant solidified material. The addition of the carbon and/or nitrogen is effected in elemental form or in the form of dissociable
10 carbon and/or nitrogen containing compounds, making said addition before or during an established thermodynamic state of dissolution of existing crystals of titanium aluminide, and bringing the melt into a thermodynamic state where crystals of titanium aluminide present grow in size and thereafter causing the melt to solidify.

15 U.S. Patent 3,961,995 discloses an aluminum-titanium-boron mother alloy having a boron content of 0.2 to 0.8% by weight and a titanium content such that $Ti - 2.2 B \geq 3.9\%$, in which the matrix has a preponderant proportion of grains of less than 30 microns in size, and contains fine TiB_2 crystals having an average size of about 1 micron primarily dispersed along the grain boundaries, and the
20 method for the preparation of same by the formation of titanium diboride by the action of liquid aluminum on titanium oxide and boron oxide in solution in molten cryolite, mixing the reactants in a manner to utilize the starting materials, and then quenching the formed alloy rapidly to cool and solidify the mother alloy, preferably by pouring the liquid alloy in water to produce the alloy in the form of granules or
25 fine powder.

U.S. Patent 4,803,372 discloses a process for producing a composite comprising a refractory material dispersed in a solid matrix. A molten composition comprising a matrix liquid, and at least one refractory carbide-forming component are provided, and a gas is introduced into the molten composition. A reactive
30 component is also provided for reaction with the refractory material-forming component. The refractory material-forming component and reactive component react to form a refractory material dispersed in the matrix liquid, and the liquid

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composite is cooled to form a solid composite material.

British Patent 1,333,957 discloses a method of preparing a master alloy intended to be added to an aluminum melt to control the grain size during solidification thereof, which comprises providing a master alloy melt containing
5 aluminum together with 0.02 to 6% by weight of titanium and 0.01 to 2% by weight of boron and holding the master alloy melt at a temperature between its melting point and 900°C under agitation for a period of at least 15 minutes and at most 9 hours.

U.S. Patent 5,100,488 discloses an improved aluminum-titanium
10 master alloy which contains in weight percent, carbon about 0.005 up to 0.05 titanium 2 to 15, and the balance aluminum. After melting, the master alloy is superheated to about 1200°C-1300°C to put the element into solution, then the alloy is cast in a workable form. The master alloy in final form is substantially free of carbides, sulfides, phosphides, nitrides, or borides greater than about 5 microns in
15 diameter. The alloy of this invention is used to refine aluminum products that may be rolled into thin sheet, foil, or fine wire and the like. Such grain refined products are also substantially free of carbides, sulfides, phosphides, nitrides or borides.

U.S. Patents 5,041,263 and 4,812,290 disclose an improved
aluminum-titanium master alloy containing carbon in an amount not more than
20 about 0.1%. After melting, the master alloy is superheated to about 1200°C-1250°C to put the carbon into solution, then the alloy is cast in a workable form. The master alloy in final form is substantially free of carbides greater than about 5 microns in diameter. The alloy of this invention is used to refine aluminum products that may be rolled into thin sheet, foil, or fine wire and the like.

U.S. Patent 4,556,419 discloses hydrogen gas and non-metallic
25 inclusions removed from molten aluminum by a process comprising the steps of maintaining an atmosphere containing BF₃ gas in a treating vessel above the surface of molten aluminum placed therein, introducing a treating gas into the molten aluminum, and removing floating non-metallic inclusions and treating gas containing
30 hydrogen gas from the surface of the molten aluminum.

U.S. Patent 4,873,054 discloses an improved aluminum-titanium master alloy. Such alloy contains a small but effective amount of, in weight

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percent, any two or more elements selected from the group consisting of carbon about 0.003 up to 0.1, sulfur about 0.03 up to 2, phosphorus about 0.03 up to 2, nitrogen about 0.03 up to 2, and boron about 0.01 up to 0.4, titanium 2 to 15, and the balance aluminum.

5 U.S. Patents 4,842,821 and 4,748,001 disclose a method of producing an alloy containing titanium carbide particles, the method comprising thoroughly dispersing carbon powder particles into a metal melt, and causing the dispersed carbon particles to react with titanium within the metal melt so as to produce a dispersion of fine particles comprising titanium carbide within the melt. A preferred
10 use for alloys produced by the invention is as a grain refiner for aluminum-based metals, especially those containing zirconium, chromium and/or manganese, which tend to poison current titanium-boron-aluminum grain refiners.

U.S. Patent 4,392,888 discloses molten aluminum or other metals purified by contacting with a fluorocarbon, such as CCl_2F_2 , in order to decrease the
15 amount of impurity metal elements along with gas and inclusions therein preferably in the presence of an agitator to enhance efficiency. An oxidizer, such as oxygen, is employed to prevent the carbon in the fluorocarbon from forming carbide inclusions. Oxidizing the carbon to carbon monoxide is preferred in treating aluminum since the monoxide effectively removes the carbon from the system
20 without oxidizing aluminum.

German Patent 1,027,407 discloses a process for grain refining aluminum alloys, especially magnesium containing alloys which contain boron and/or titanium characterized by the fact that carbon is produced in finely divided form by decomposing carbon compounds, e.g., carbon tetrachloride, which are
25 introduced to the melt in a carrier gas to promote formation of carbides as crystallization nuclei. The patent states that the carbon tetrachloride is not dangerous to foundry personnel because its vapor will decompose at normal casting temperatures.

U.S. Patent 4,402,741 discloses a process and an apparatus for the
30 precise and continuous injection of a halogenated derivative, which is liquid at ambient temperature, into a liquid metal such as aluminum and aluminum-based alloys. The process involves withdrawing the halogenated substance from a tank,

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introducing it by means of a metering pump into a vaporizer which has been brought to a temperature at least equal to the vaporization temperature of the substance under the injection pressure, and entraining it in the vapor state by an inert gas stream towards an injection means opening into the center of the liquid metal.

In spite of these disclosures there is still a great need for improvements in grain refining of aluminum which do not contaminate the metal and which permit grain refining at molten aluminum processing temperatures. The present invention provides such an improvement.

It is an object of the invention to provide an improved process for treating molten aluminum to improve structural characteristics of products cast therefrom.

It is another object of the invention to provide a combination of molten metal treatment, e.g., degassing, and forming grain refining nuclei.

Yet, it is a further object of the invention to provide improved grain refining of aluminum without the use of master alloys.

And, it is another object of the invention to provide a process for improved grain refining of aluminum.

It is yet another object of the invention to provide an improved grain refining nuclei for aluminum, i.e., smaller, better dispersed and more efficient.

It is still another object of the invention to provide a titanium based grain refining nuclei for aluminum.

And, it is still another object of the invention to provide a process for in situ development of grain refining nuclei for aluminum at molten aluminum treatment temperatures without the use of master alloys.

And yet, it is another object of the invention to grain refine aluminum by producing very fine particle sized grain refiner in situ prior to casting.

Still yet, it is another object of the invention to provide an improved aluminum alloy substantially free of clusters and oxides and having controlled size grain refiner particles and grains in the solidified aluminum.

And still yet, it is another object of the invention to provide an improved process for molten aluminum treatment having improved filter life.

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Other objects of the invention will become apparent from the specification and claims appended hereto.

In accordance with these objects there is provided a method of molten metal treatment wherein the molten metal is treated to remove impurities and is provided with grain refining nuclei in situ. The method comprises providing a molten aluminum body containing 1 to 3000 ppm titanium, preferably 1 to 1500 ppm. A material reactive with the titanium is introduced to the aluminum body. The material reactive with titanium is preferably in gaseous form. The titanium reactive material has at least one component thereof that is comprised of at least one of the group consisting of boron, carbon, sulfur, nitrogen and phosphorus. The reactive material and said titanium form a grain refining compound adapted for grain refining the aluminum as it proceeds to the casting operation.

In the accompanying drawings:

Figure 1 is a schematic illustration of steps in treating a body of molten aluminum showing addition of grain refining materials in accordance with the invention.

Figure 2 is a cross-sectional view of a metal treatment bay for fluxing molten aluminum.

In one embodiment of the invention, a body of molten aluminum is grain refined by providing in the body a controlled level of titanium and adding to the molten aluminum a compound or material which forms small, discrete titanium compounds such as TiC or TiB_2 that provide nucleation sites for grain refining aluminum.

Referring now to Figure 1, there are shown steps for processing and grain refining an aluminum body in accordance with the invention to both purify and grain refine the molten aluminum as it is directed from the smelting or melting to the casting operation. In Figure 1, there is shown a body of aluminum, referred to as 10. Molten aluminum is drawn from body 10 along line 12 to a cast house furnace 14 where metal is contained prior to casting. For purposes of casting, molten metal is withdrawn from furnace 14 along line 16 to a metal treatment step 18. In the metal treatment step typically a fluxing gas is dispersed therein. The fluxing gas removes both dissolved and suspended solid impurities, including

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oxides, nitrides, carbides and carbonates of the molten metal and alloying elements. The impurities include both dissolved gases and dissolved elements. Dissolved gases in molten aluminum, for example, include hydrogen and dissolved elements include alkali and alkaline elements such as sodium and calcium. Suspended solid
5 impurities are transported to the melt surface by attachment to rising bubbles of the fluxing gas. Hydrogen is desorped into the gas bubbles for removal.

The fluxing gases that can be used for molten aluminum include nitrogen containing gases, halogen gases and the so-called inert gases, namely, helium, neon, argon, krypton, and xenon along with nitrogen, carbon dioxide and
10 mixtures of these gases. In addition, chlorine and sulfur hexafluoride may be used, typically in combination with the inert gas. Small amounts of chlorine or fluorine gases or other halides or mixtures thereof such as BCl_3 , BF_3 , BFCl_2 , BBr_3 , BI_3 , PCl_5 , and SOCl_2 can assist in removing suspended solids and dissolved elements.

With respect to molten metal treatment, there is shown in Figure 2 a
15 schematic view of a metal treatment or gas fluxing bay 50 having a hollow shaft 54 and impeller or gas disperser 56 located in a body of molten metal, e.g., aluminum 58. Shaft 54 is carried by structure 60. Further, shaft 54 is rotated by motor 64. The rotation can be unidirectional or bidirectional. Fluxing or carrier gas is added through tube 70 and down hollow shaft 54 before being dispersed through tubes or
20 conduits in impeller 56. Instead of passing fluxing gas down hollow shaft 54, the gas may be added through a tube or other means and dispersed by impeller 56. Alternatively, the gas may be injected adjacent to the impeller for dispersion purposes. Also, diffusers may be used to introduce the gas and distribution may be assisted by the rotating impeller.

25 After the metal treatment step, molten metal is conveyed along line 26 through filter 28 which removes particles detrimental to the processing of the cast product. After filtration, the molten metal is removed along line 30 to casting facility 32 where the molten metal is cast into billet, ingot or slab, for example.

For purposes of the present invention, preferably the molten
30 aluminum in cast house furnace 14 is maintained at a temperature in the range of 1200° to 1500°F or higher, if desired. Higher temperatures can be used for the molten metal treatment process shown in Figure 1 but are not currently believed

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necessary for purposes of the present invention. Normally, higher temperatures are undesirable not only because of the extra cost but also because there is a greater tendency to form oxides, pick up hydrogen, degrade refractories, etc.

In accordance with the present invention, a source of titanium 13 is added to the aluminum melt typically in furnace 14 or during metal treatment step 18. It should be noted that the titanium can be added anywhere in the process shown in Figure 1 as long as it is added concurrently or prior to the addition of the titanium reactive material. Preferably, the titanium is added in the furnace as shown in Figure 1. The amount of titanium added is that sufficient to provide for grain refining of the aluminum body. The amount of titanium that is effective for grain refining in the present invention can range from about 1 ppm to about 3000 ppm, with a preferred amount ranging from 20 to 1500 ppm and typically an amount ranging from 40 to less than 1000 ppm, e.g., 40 to 950 ppm, typically 40 to 600 ppm. Higher amounts of titanium can be used but care is required to avoid exceeding the solubility limit of titanium in molten aluminum or the formation of substantial amounts of titanium aluminide (TiAl_3) particles in the melt. Titanium aluminide forms large particles which are detrimental in processing or working the cast product.

Titanium 13 can be added to the molten aluminum as titanium metal or metal alloy. In another aspect of the invention, the titanium may be added to the molten aluminum as a titanium compound that is reduced by the molten aluminum. Such compounds can include TiCl_4 , K_2TiF_6 or Na_2TiF_6 , for example. Or, the titanium can be added to the molten aluminum as compacts, waffles or swarfs. Typically, waffles have a composition comprising 5 to 10 wt.% or more of titanium. For example, waffles typically comprise 6 to 10 wt.% titanium, the remainder aluminum. Compacts by comparison usually comprised 80 wt.% or greater titanium. The compacts are comprised of particles having a large surface area, and such compacts can contain fluxing salts such as sodium chloride, potassium chloride, potassium aluminum tetrafluoride, sodium fluoride, and potassium boron fluoride which aid in dissolution of the titanium in the melt and improves the integrity of the compact by holding the particles together.

After titanium 13 has been added to the molten aluminum, a material

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or compound 19 is added to the melt which reacts with the titanium and/or the aluminum to form a titanium based grain refiner nuclei in situ. The material or compound which reacts with the titanium and/or aluminum is referred to as a reducible binary or titanium reactive material. For purposes of the invention, it will be appreciated that titanium reactive material 19 can be added with the titanium. However, as noted, it is preferred that titanium 13 is added before the addition of titanium reactive material 19. Adding the titanium first is effective in minimizing the reaction of titanium reactive material 19 with aluminum to form aluminum compounds such as, for example, aluminum carbide, aluminum boride, aluminum sulfide, aluminum phosphide or aluminum nitride, depending to some extent on the reactive material being added.

With reference to Figure 1, it will be noted that titanium reactive material or compound 19 can be added during molten metal treatment step 18 or it can be added at 19a after metal treatment 18 but prior to molten metal filter step. In yet another embodiment, reactive compound 19 may be added at 19b after molten metal filter 28. That is, the present invention contemplates addition of the reactive material after the molten metal filter step and even in a tundish. Or, the reactive material can be added in a trough used to convey molten aluminum to the mold for casting purposes.

It is important to minimize the time between casting the molten aluminum and adding the titanium reactive material. That is, if the reactive material is added earlier, it can permit some settling of the grain refiner particles to occur. Thus, for purposes of the present invention, to minimize settling (sometimes referred to as fade) of the grain refiner, it is preferred to add the reactive material or compound as near the casting step as possible. In the present invention, the reactive material or compound is preferably added during the metal treatment step or following the filtration step.

In the present invention, the amount of titanium reactive material added is important. That is, it is preferred to add the titanium reactive material or compound at a level below its solubility limit in molten aluminum. If the solubility limit of the reactive material in molten aluminum is exceeded, then undesirable compound or precipitates form. Further, it is preferred that the titanium

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concentration is maintained stoichiometrically in excess of the reactive material or compound in the molten aluminum body. Thus, the molar ratio of titanium to reactive material in the melt is maintained such that there is an excess of titanium present in the active nuclei being formed, e.g., 0.5 molar for TiB_2 or 1.0 to TiC .

- 5 The concentration and ratio depends to some extent on the titanium reactive material used.

Titanium reactive material suitable for grain refining in combination with titanium include compounds which will provide at least one of the following elements: boron, carbon, sulfur, phosphorus and nitrogen in the molten aluminum.

- 10 It should be understood that any compound or material may be used which provides an element which in combination with titanium operates to provide grain refining nuclei in situ.

- The reactive compound or material which operates with titanium to form active nuclei in situ for grain refining aluminum may either be solid, liquid or
15 gaseous at the temperature of molten aluminum. For purposes of the invention it is preferred that the reactive compound is provided in gas form which facilitates addition of such reactive compounds particularly during the molten metal treatment step. By gaseous material or gaseous compound is meant a material or compound which is in finely divided or in gas form at molten aluminum temperature.

- 20 Exemplary of reactive compounds that may be used in the invention include halide compounds such as chlorine, fluorine, bromine, or iodine, or mixed halides of boron, carbon, nitrogen, sulfur, and phosphorus. That is, reactive compounds of the invention can include chloride and fluoride compounds having a boron or carbon component. Preferably, the reactive chloride compounds include BCl_3 , CCl_4 , C_2Cl_4 ,
25 CHCl_3 , NCl_3 and PCl_3 and the reactive fluoride compounds include SF_6 , KBF_4 , BF_3 and NF_3 . Hydrides such as boron hydrides or boranes may also be used. These materials may be provided in gas or aerosol form. For example, KBF_4 may be provided as a powder and added as an aerosol.

- As noted, it is preferred that these compounds be introduced to the
30 melt during the metal treatment step. Further, it will be appreciated that a single reactive compound may be used or a combination of compounds may be used, particularly if it is desired to provide both boron and carbon in the melt to form

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dispersion of TiB_2 and TiC for purposes of grain refining. With reference to Figure 2, it will be seen that the reactive compound when in gaseous form is suitably added with the fluxing gas and dispersed efficiently through the melt, thus avoiding the localized, high concentration referred to earlier. If the reactive compound is in solid or liquid form, it may be atomized or volatilized and added with the fluxing gas as shown in Figure 2. For example, organic based liquids such as kerosene, carbon tetrachloride, vinyl chloride, polytetrafluoroethylene, butane, freon and ethylene chloride can be used by atomizing into small droplets using a ultrasonic nozzle. The droplets may be suspended in a sol and introduced to the gas stream. Or, the liquid may be vaporized into the gas stream.

If the titanium reactive material is provided in solid form it may be ground to a powder, e.g., powdered carbon, and introduced with the fluxing gas for dispersion therewith. Alternatively, solid titanium reactive compound may be added directly to the melt and dispersed with the impeller and/or fluxing gas.

It will be appreciated that the reactive compound may have a component such as hydrogen which is undesirable in the aluminum cast product. However, the hydrogen is effectively removed by the fluxing gas which is added concurrently or after the addition of the reactive compound, for example. As will be noted, the chlorine or fluorine components or other halide components or mixtures thereof of the titanium reactive material are not usually detrimental and thus do not present difficulties.

In the present invention, the amount of reactive compound added with or dispersed by the fluxing or carrier gas depends to some extent on the amount of titanium present in the melt and the extent of grain refining desired. Thus, the amount of reactive material is adjusted or added in accordance with the grain refining required.

Typically, the material reactive with titanium is provided in the fluxing gas in the range of 1 to 99 vol.% and preferably 2 to 50 vol.%, and typically 2 to 20 vol.%. The material reactive with titanium can be added to provide at least one of the group consisting of boron, carbon, sulfur, nitrogen and phosphorus in the molten aluminum in the range of 0.01 to 400 ppm. For example, the boron containing reactive material is added to the fluxing or carrier gas to add

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boron in the range of 0.5 to 25 ppm, preferably 1 to 10 ppm, and typically in the range of about 2 to 5 ppm. When added in this way, the process is continuous, which in turn encourages production of a uniform nuclei size distribution.

By the use of the term "in situ" as used herein is meant to include or refer to aluminum in the normal operation or process of (1) smelting or melting, (2) treating molten aluminum for purification to remove dissolved and undissolved material, and (3) casting into, for example, ingot, billet, slab, or other cast products including foundry cast products. The use of the term "in situ" is not meant to include wire, rod, or ingot or the like of aluminum grain refiner alloys typically having high loading, e.g., 1 wt.%, of grain refiner and typically formed in an independent operation. The term nuclei or active nuclei as used herein, is meant to refer to compounds such as titanium diboride, titanium carbide or like compounds which operate to produce grain refined aluminum bodies on solidification.

It should be noted that titanium even in low concentrations, e.g., 50 ppm, reacts preferentially with the boron or carbon component, for example, of the reactive compound and minimizes any significant reaction of the boron or carbon with aluminum thus greatly avoiding undesirable compounds of aluminum.

As noted, the titanium and titanium reactive compound are preferably added to the melt below their solubility limits in pure aluminum with the titanium being in excess of the titanium reactive compound. The concentrations of the titanium and the reactive compounds are controlled to control the particle size of the grain refiner formed in the melt. To minimize the size of grain refining particles, the concentrations of both the titanium and reactive compounds preferably are maintained at low levels. For example, when the titanium is 100 ppm, the reactive component can be as follows: boron can be added in the range of 0.2 to 21 ppm; carbon can be added in the range of 0.2 to 23 ppm; sulfur can be added in the range of 0.6 to 63 ppm; phosphorus can be added in the range of 0.6 to 61 ppm; and nitrogen can be added in the range of 0.2 to 27 ppm. This can produce grain refiner particle sizes in the range of 0.1 to 1 μm and typically, 0.2 to 0.6 μm . It is believed that further dilution of the concentration, e.g., 10 ppm titanium to 0.1 to 1 ppm reactive component, may produce even smaller grain refiner particle sizes. All ranges set forth herein include all the numbers within the range as if specifically set

forth.

If the source of titanium is provided or added along with the reactive material, e.g., in the carrier or fluxing gas, preferably titanium and the reactive material are provided in a ratio which permits stoichiometric excess of titanium.

5 Also, the source of titanium and reactive material may be provided in amounts sufficiently dilute to favor formation of small particle size grain refiner compounds, e.g., TiC or TiB₂ in situ. In this way, only a limited amount of free titanium is available to react with the reactive material which results in small particle size grain refiner.

10 Typically, only small amounts (e.g., 0.2 to 5 ppm) of titanium reactive compound is required when added or produced by this method to provide well grain refined aluminum castings. The amount of inert or fluxing gas used to metal refine aluminum is many times greater on a molar basis than the titanium reactive compound. Therefore, the titanium reactive compound can be diluted with
15 inert or fluxing gas as desired.

In another aspect of the invention, it should be understood that the titanium reactive material can be added after molten metal treatment step 18, and typically at 19a before molten metal filter 28 or at 19b after molten metal filter 28. That is, the titanium reactive material may be added as a substantially pure gas
20 without dilution after the molten metal treatment step. Adding the reactive material after the molten treatment step operates to avoid loss of titanium reactive material or compounds or nuclei formed therefrom by removal in the fluxing gas, e.g., argon.

Even though the invention has been described particularly with respect to titanium, it will be appreciated that other metals are contemplated within
25 the perview of the invention, including but not limited to niobium, tantalum, vanadium, molybdenum, zirconium and beryllium.

Example 1

A high purity aluminum alloy was melted in a gas fired crucible furnace, and a small addition of titanium was made to the melt to provide the
30 following composition:

Fe	Si	Ti	Cu	Mn	V	B
<u>0.056</u>	<u>0.042</u>	<u>0.025</u>	<u>0.001</u>	<u>0.001</u>	<u>0.004</u>	<u>0.0001</u>

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A carbon lance was introduced to the crucible and argon was bubbled through the melt for 30 minutes to remove gas and any suspended inclusions. A conical sample was taken, according to test procedures established by the Aluminum Association (AA). The grain size was measured at a height of 1.5" from the bottom of the sample, according to procedures outlined in ASTM E112. The grain size (average intercept distance, or AID) was 3600 microns.

A gas mixture of argon-10% BCl_3 was then bubbled through the melt for a time, after which the boron concentration was found to be 0.0013% (or 13 ppm). The AA grain size was 210 microns.

10 Example 2

In this example, the procedure was similar to Example 1, except that the metal had a starting composition of 0.59% Si and 0.005% Ti. The grain size before treatment was about 2000 microns.

A gas mixture of argon-10% BCl_3 was then bubbled through the melt, to produce a boron concentration of 0.0006% (6 ppm). The AA grain size was 225 microns.

Example 3

In this example, the test procedure was the same as Example 2, except a gas mixture of argon-10% BF_3 was then bubbled through the melt, to produce a boron concentration of 0.0006% (6 ppm). The AA grain size was 250 microns.

It will be seen from the examples that aluminum is effectively grain refined using the process of the invention.

While the invention has been described in terms of preferred embodiments, the claims appended hereto are intended to encompass other embodiments which fall within the spirit of the invention.

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C L A I M S

1. An improved method for treating molten aluminum for solidifying into cast products wherein the molten aluminum is subject to a metal treatment for removing impurities, the improved method comprising forming grain refiner in the molten aluminum, the method comprising:
- (a) providing a molten aluminum body;
 - (b) providing 1 to 3000 ppm titanium in said moltenaluminum body;
 - (c) introducing to said molten aluminum body, a material reactive with said titanium, said material being in gaseous form at molten aluminum temperature and comprising at least one component of the group consisting of boron, sulfur, nitrogen and phosphorus, said material and said titanium forming grain refining nuclei in the aluminum body; and
 - (d) solidifying at least a portion of said molten aluminum body into a grain refined, cast product.
2. The method in accordance with claim 1, including dispersing said material reactive with said titanium in said molten aluminum body using a carrier or fluxing gas.
3. The method in accordance with claim 1, including introducing said material reactive with said titanium in a carrier or fluxing gas.
4. The method in accordance with claim 3, including introducing said material reactive with titanium separate from said carrier or fluxing gas.
5. The method in accordance with claim 1 including maintaining said titanium in the range of 20 to 1500 ppm.
6. The method in accordance with claim 1, including maintaining said titanium in the range of 40 to less than 1000 ppm.
7. The method in accordance with claim 1, including maintaining in titanium is in the range of 40 to 600 ppm.
8. The method in accordance with claim 1, including maintaining said titanium in stoichiometric excess of said material reactive with said titanium.
9. The method in accordance with claim 1, including providing said titanium in said aluminum body prior to adding said material reactive with said titanium.

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10. The method in accordance with claim 1, including maintaining said body of molten aluminum in a temperature range of 1200° to 1500°F.

11. The method in accordance with claim 1, including selecting said material reactive with said titanium from at least one of the group consisting of BCl₃,
5 PCl₃, SF₆, KBF₄, BF₃ and NH₃.

12. The method in accordance with claim 1, including selecting said material reactive with said titanium from the group consisting of at least one of a chloride or fluoride of boron and phosphorus.

13. The method in accordance with claim 1, including forming a titanium
10 based grain refiner in said molten aluminum having a particle size in the range of 0.05 to 2 μm.

14. The method in accordance with claim 1, including maintaining said material reactive with titanium in a carrier or fluxing gas in the amount of 1 to 50 vol.% of the carrier or fluxing gas.

15 15. The method in accordance with claim 1, including introducing said material reactive with said titanium in a gas selected from the group consisting of helium, neon, argon, krypton, xenon, nitrogen, sulfur hexafluoride, carbon dioxide and chlorine and mixtures thereof.

16. The method in accordance with claim 3, including introducing said
20 material reactive with titanium in said carrier or fluxing gas to provide at least one of the group consisting of boron, sulfur, nitrogen and phosphorus in the molten aluminum body in the range of 0.01 to 400 ppm.

17. An improved method for treating molten aluminum for solidifying into cast products wherein the molten aluminum is subject to a molten metal treatment for
25 removing impurities, the method comprising:

(a) providing a molten aluminum body in a temperature range of 1200° to 1500°F;

(b) providing titanium in the range of 1 to 1500 ppm in said molten aluminum body;

30 (c) after providing said titanium in said body, introducing to said body a material reactive with said titanium, said material being in gaseous form at molten aluminum temperatures and being introduced to said body in a carrier gas or fluxing

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gas, said material selected from at least one of a chloride or fluoride of one of the groups consisting of boron, sulfur, nitrogen and phosphorus, said material and said titanium forming a grain refining nuclei in said molten aluminum body; and

5 (d) solidifying at least a portion of said molten aluminum body into a grain refined, cast product.

18. The method in accordance with claim 17, including selecting said material reactive with said titanium from at least one of the group consisting of BCl_3 , PCl_3 , SF_6 , KBF_4 , BF_3 and NH_3 .

10 19. The method in accordance with claim 17, including maintaining said titanium in the range of 20 to less than 1000 ppm.

20. The method in accordance with claim 17, including maintaining said titanium in the range of 20 to 600 ppm.

15 21. The method in accordance with claim 17, including forming a titanium based grain refiner in said molten aluminum having a particle size in the range of 0.1 to 2 μm .

22. The method in accordance with claim 17, including maintaining said titanium in stoichiometric excess of said material reactive with said titanium.

20 23. The method in accordance with claim 17, including maintaining said material reactive with titanium in a carrier or fluxing gas in the amount of 1 to 50 vol.% of the carrier or fluxing gas.

24. The method in accordance with claim 17, including introducing said material reactive titanium in a gas selected from the group consisting of helium, neon, argon, krypton, xenon, nitrogen, sulfur hexafluoride, carbon dioxide and chlorine and mixtures thereof.

25 25. In a continuous method of casting molten aluminum into solidified products wherein the molten aluminum is subject to a metal treatment prior to said casting operation, comprising a method of providing grain refining nuclei in the molten aluminum on a continuous basis, the method comprising:

- 30 (a) providing a molten aluminum body;
- (b) adding a source of titanium to said molten aluminum body to provide titanium in the range of 1 to 3000 ppm;
- (c) after adding said titanium, adding to said molten aluminum body a

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material reactive with said titanium, said material being in gaseous form and comprising at least one component of the group consisting of boron, sulfur, nitrogen and phosphorus, said material and said titanium adapted for grain refining aluminum;

5 (d) maintaining said titanium in stoichiometric excess of at least one of the group consisting of boron, sulfur, nitrogen and phosphorus; and

(e) solidifying at least a portion of molten aluminum body to provide a grain refined, cast product.

26. The method in accordance with claim 25, including fluxing said molten aluminum body with a fluxing gas.

10 27. The method in accordance with claim 26, including adding said material reactive with said titanium with said fluxing gas.

28. The method in accordance with claim 25, including dispersing said material reactive with said titanium in said molten aluminum body using a carrier or fluxing gas.

15 29. The method in accordance with claim 25, including selecting said material reactive with said titanium from at least one of the group consisting of BCl_3 , PCl_3 , SF_6 , KBF_4 , BF_3 and NH_3 .

30. The method in accordance with claim 25, including selecting said material reactive with said titanium from the group consisting of at least one of a
20 chloride or fluoride of boron and phosphorus.

31. The method in accordance with claim 26, including forming a titanium based grain refiner in said molten aluminum having a particle size in the range of 0.05 to 2 μm .

32. The method in accordance with claim 25, including maintaining in said
25 material reactive with titanium in a carrier or fluxing gas in the amount of 1 to 50 vol.% of the carrier or fluxing gas.

33. The method in accordance with claim 25, including maintaining in said material reactive with titanium in a carrier or fluxing gas in the amount of 2 to 20 vol.% of the carrier or fluxing gas.

30 34. The method in accordance with claim 25, including introducing said material reactive titanium in a gas selected from the group consisting of helium, neon, argon, krypton, xenon, nitrogen, sulfur hexafluoride, carbon dioxide and chlorine and

mixtures thereof.

35. An improved method of fluxing, grain refining and casting aluminum, the method comprising the steps of:

- (a) providing a body of molten aluminum;
- 5 (b) providing titanium in said body of molten aluminum in a range of 1 to 3000 ppm;
- (c) subjecting said body to a fluxing treatment with a gas to remove both dissolved and suspended materials;
- (d) contacting said molten aluminum body with a gaseous compound
- 10 reactive with said titanium, said gaseous compound having at least one component selected from the group consisting of boron, sulfur, nitrogen and phosphorus;
- (e) forming a grain refining material in said molten aluminum comprised of said gaseous compound and said titanium; and
- (f) casting said molten aluminum into a grain refined, solid form.

15 36. The method in accordance with claim 35, including maintaining said body of molten aluminum in a temperature range of 1200° to 1500°F.

37. The method in accordance with claim 35, including maintaining in said body of molten aluminum said titanium in stoichiometric excess of at least one of the group consisting of boron, sulfur, nitrogen and phosphorus.

20 38. The method in accordance with claim 35, including providing said titanium in said body of aluminum prior to adding said gaseous compound reactive with said titanium.

39. The method in accordance with claim 35, including introducing said gaseous compound in a carrier or said fluxing gas.

25 40. The method in accordance with claim 35, including selecting said gaseous compound from at least one of the group consisting of PCl_3 , SF_6 , KBF_4 , BF_3 and NH_3 .

41. The method in accordance with claim 35, including selecting said gaseous compound from the group consisting of at least one of a chloride or fluoride

30 of boron and phosphorus.

42. The method in accordance with claim 35, including forming a titanium based grain refining nuclei in said molten aluminum having a particle size in the range

of 0.05 to 2 μm .

43. The method in accordance with claim 35, including introducing said gaseous compound in a gas selected from the group consisting of helium, neon, argon, krypton, xenon, nitrogen, sulfur hexafluoride, carbon dioxide and chlorine and mixtures thereof.

44. A method of grain refining aluminum, the method comprising:
(a) providing a molten aluminum body;
(b) providing 1 to 1500 ppm titanium in said molten aluminum body;
(c) introducing to said molten aluminum body a material reactive with said titanium, said material being in gaseous form and comprising at least one component of the group consisting of boron, sulfur, nitrogen and phosphorus; and
(d) forming a grain refining compound comprised of said material and said titanium in said molten aluminum body and casting a grain refined product.

45. A method of grain refining aluminum, the method comprising:
(a) providing a molten aluminum body in a temperature range of 1200° to 1500°F;
(b) providing titanium in said molten aluminum body;
(c) after providing said titanium in said body, introducing to said body a material reactive with said titanium, said material being in gaseous form and introduced to said body of molten aluminum in a carrier or fluxing gas, said material being a chloride or fluoride of one of the groups consisting of boron, sulfur, nitrogen and phosphorus; and
(d) forming a grain refining compound comprised of said material and said titanium in said molten aluminum body and casting a grain refined product.

46. A method of grain refining aluminum, the method comprising:
(a) providing a molten aluminum body;
(b) adding a source of titanium to said molten aluminum body in a range of 1 to 1500 ppm;
(c) after adding said titanium, contacting said aluminum body with a material reactive with said titanium, said material being in gaseous form and having a component thereof selected from at least one of the group consisting of boron, sulfur, nitrogen and phosphorus, said material and said titanium adapted for grain refining said

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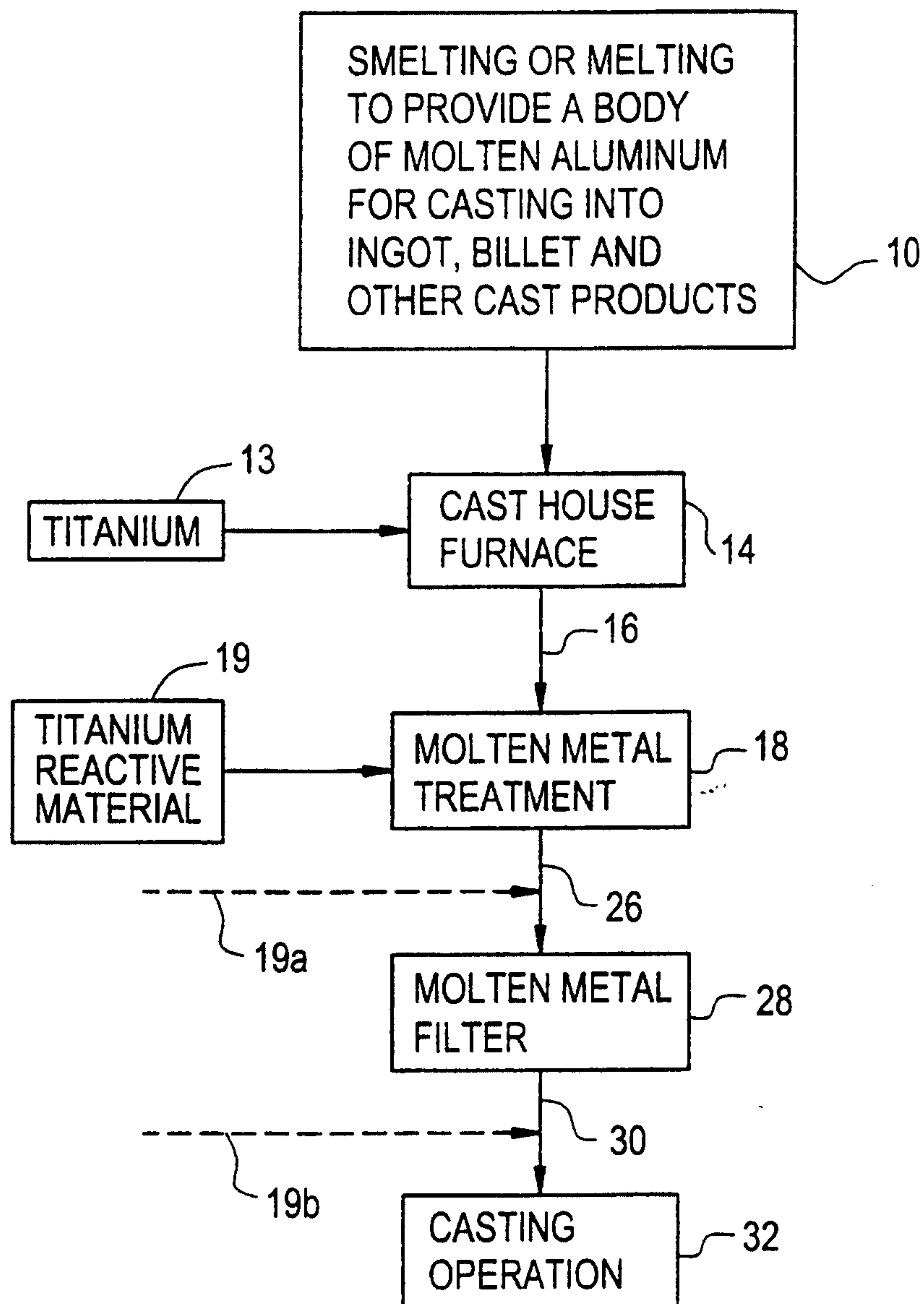
aluminum body;

(d) maintaining said titanium in stoichiometric excess of at least one of the group consisting of boron, sulfur, nitrogen and phosphorus; and

(e) forming a grain refining compound comprised of said material and
5 said titanium in said molten aluminum body and casting a grain refined product.

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FIG. 1



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FIG. 2

