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(11) **EP 1 063 312 B1**

(12) **EUROPEAN PATENT SPECIFICATION**

(45) Date of publication and mention  
of the grant of the patent:

**21.04.2004 Bulletin 2004/17**

(21) Application number: **99949393.5**

(22) Date of filing: **25.10.1999**

(51) Int Cl.7: **C22C 16/00, C22C 45/10**

(86) International application number:  
**PCT/JP1999/005872**

(87) International publication number:  
**WO 2000/026425 (11.05.2000 Gazette 2000/19)**

(54) **HIGH-STRENGTH HIGH-TOUGHNESS AMORPHOUS ZIRCONIUM ALLOY**

HOCHZÄHE, HOCHFESTE AMORPHE ZIRKONIUMLEGIERUNG

ALLIAGE AMORPHE DE ZIRCONIUM A HAUTE RESISTANCE ET TENACITE ELEVEE

(84) Designated Contracting States:  
**DE FR GB**

(30) Priority: **30.10.1998 JP 31010898**

(43) Date of publication of application:  
**27.12.2000 Bulletin 2000/52**

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**EP-A- 0 433 670 EP-A- 1 036 854**  
**JP-A- 7 062 502 JP-A- 7 188 877**  
**JP-A- 8 199 318 JP-A- 10 186 176**  
**US-A- 5 735 975 US-A- 5 803 996**

• **PATENT ABSTRACTS OF JAPAN vol. 1996, no.  
07, 31 July 1996 (1996-07-31) -& JP 08 074010 A  
(INOUE AKIHISA; MASUMOTO TAKESHI;  
TEIKOKU PISTON RING CO LTD), 19 March 1996  
(1996-03-19)**

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**Description**

## FIELD OF THE INVENTION

5 **[0001]** The present invention relates to amorphous Zr alloys which have a high glass-forming ability and excellent strength and toughness.

## RELATED ART

10 **[0002]** Amorphous metal materials having various forms, such as thin ribbons, filaments, particles and the like, can be obtained by rapidly cooling molten alloys. A thin-ribbon-shaped amorphous alloy is easily manufactured by means of a single roll method, a twin-roller method, an in-rotating water melt spinning method and the like, in which a large cooling speed can be obtained. Conventionally, various amorphous alloys have been provided using alloys of Fe, Ni, Co, Pd, Cu, Zr or Ti; those amorphous alloys show properties unique to amorphous alloys such as high corrosion resistance, high strength, and the like. Especially, an amorphous Zr alloy is expected to be applied to the fields of structural materials, medical materials and chemical materials as a new kind of amorphous alloy having an outstanding high glass-forming ability compared to other amorphous alloys.

15 **[0003]** However, shapes of the amorphous alloys manufactured by means of previously mentioned methods are limited to thin ribbons or thin wires; it is difficult to process the amorphous alloys of those shapes into a form of final products. Therefore, the uses of such amorphous alloys are limited in industry.

20 **[0004]** On the other hand, it is known that when amorphous alloys are heated, some of alloys undergo transition to a phase of supercooled liquid before crystallization and indicate a decline in viscosity. For example, when heated at a speed of 40°C per minute, an amorphous Zr alloy is observed to remain in the supercooled liquid phase for a range of temperature of the maximum of 120°C before crystallization starts (see Mater. Trans., JIM, Vol. 32, (1991), 1005).

25 **[0005]** In the supercooled liquid phase, the low viscosity of the amorphous alloy allows one to form it into a given shape by closed squeeze casting process and the like; for example, gears can be formed of an amorphous alloy (see Nikkan Kogyo Shinbun, November 12, 1992). Hence, amorphous alloys having a wide range of the supercooled liquid phase can be said to provide excellent workability. Among such amorphous alloys having a wide range of supercooled liquid phase, an amorphous Zr-Al-Ni-Cu alloy has a range of temperature of 100°C as the supercooled liquid phase, therefore, is considered to be an amorphous alloy with excellent applicability, such as high corrosion resistance (see Japanese Examined Patent Application Publication H07-122120 or EP equivalent EP-A- 0 433 670).

30 **[0006]** The glass-forming ability and a method for manufacturing of those amorphous alloys have been further improved. As a result, Japanese Laid-Open Patent Application Publication H08-74010 (JP-A-08-074010) discloses development of an amorphous Zr alloy having a 100°C range for the supercooled liquid phase and a thickness exceeding 5mm. Also, various manufacturing methods to improve mechanical characteristics of the amorphous alloys have been tried (Japanese Laid-Open Patent Application Publications: 2000-24771 (EP equivalent: EP-A-1 036 612), 2000-26943, 35 2000-26944 (EP equivalent: EP-A-1 036 854)); however, these amorphous Zr alloys do not provide sufficient mechanical characteristics as structural materials.

## 40 DETAILS OF THE INVENTION

(Problems to be Solved)

45 **[0007]** The amorphous Zr alloy described previously has a high glass-forming ability and relatively good strength characteristics due to the range of the supercooled liquid phase above 100°C. Nonetheless, attempts to improve its mechanical characteristics have been made only in the manufacturing method; attempts to improve the composition of alloys has not been made.

(Solution of the Problems)

50 **[0008]** Intending to provide an amorphous Zr alloy material having improved strength and toughness without impairing a temperature range for the supercooled liquid phase and a size enabling application to industrial use, inventors of the present invention studied the above issues. They discovered the an amorphous Zr alloy having high strength and toughness as well as excellent glass-forming ability can be obtained by melting an alloy in which a given amount of M element (one or two or more elements selected from a group consisting of Ti, Nb and Pd) is added to a Zr-Al-Ni-Cu-M alloy of a given composition, followed by rapid cooling for solidification.

55 **[0009]** In other words, the present invention intends to provide an amorphous Zr alloy which contains non-crystalline phase of 90% or higher by volume wherein the alloy is defined in claim 1.

**[0010]** Further, a "range of the supercooled liquid phase" is defined as a difference between a glass transition temperature, obtained by differential scanning thermogravimetry at a speed of heating of 40°C per minute, and a crystallization temperature. The "range of the supercooled liquid phase" indicates resistance to crystallization, that is, stability of glass-forming ability. The alloy of the present invention has a range of the supercooled liquid phase over 100°C.

**[0011]** US-A-5 735 975 discloses in Table I, column 7, lines 31, 32 and 42 compositions which are of interest with regard to the disclosure of the present application.

**[0012]** The document US-A-5 735 975 discloses alloys comprising zirconium and/or hafnium in the range of 45 to 65 atomic percent, titanium and/or niobium in the range of 4 to 7.5 atomic percent, and aluminum and/or zinc in the range of 5 to 15 atomic percent. The balance of the alloy compositions comprises copper, iron, and cobalt and/or nickel. The composition is constrained such that the atomic percentage of iron is less than 10 percent. Further, the ratio of copper to nickel and/or cobalt is in the range of from 1:2 to 2:1. The alloy composition formula of the document US-A-5 735 975 is:

$(\text{Zr,Hf})_a(\text{Al,Zn})_b(\text{Ti,Nb})_c(\text{Cu}_x\text{Fe}_y(\text{Ni,Co})_z)_d$  wherein the constraints upon the formula are: a ranges from 45 to 65 atomic percent, b ranges from 5 to 15 atomic percent, c ranges from 4 to 7.5 atomic percent, d comprises the balance, dxy is less than 10 atomic percent, and x/z ranges from 0.5 to 2.

#### PREFERRED EMBODIMENTS OF THE PRESENT INVENTION

**[0013]** The following describes the most preferable embodiments of the present invention.

**[0014]** In an amorphous Zr alloy of the present invention, Ni and Cu are main elements forming the non-crystalline phase; a sum of the amounts of Ni and Cu contained is more than 30 atomic % and less than 50 atomic %. When the sum is less than 30 atomic % or more than 50 atomic %, the single roll method with a high cooling speed can provide the non-crystalline phase while the casting method with a low cooling speed cannot. Further, a ratio of the amount of Ni to the amount of Cu contained, i.e., b/c ratio, is defined to be less than 1/3. This ratio provides dense random packing of the atomic structure of the non-crystalline phase such that the glass-forming ability is maximized.

**[0015]** Also, Al is an element to drastically increase the glass-forming ability of an amorphous Zr alloy of the present invention. The amount of Al contained is more than 5 atomic % and less than 10 atomic %. When the amount contained is less than 5 atomic % or more than 10 atomic %, the glass-forming ability decreases.

**[0016]** M is one or two or more elements selected from a group consisting of Ti, Nb and Pd; additionally, it accelerates the dense random packing of the atomic structure while effectively strengthening the bond strength between atoms. As a result, higher strength and toughness are given to an amorphous Zr alloy having the high glass-forming ability. The amount of M is when M is Ti or Nb less than 4 atomic %, or when M is Pd less than 7 atomic %. When the amount of each element contained exceeds the defined atomic %, the bond strength between atoms becomes too strong such that a compound phase with Zr or Al will be formed. The compound phase causes structural discontinuity in the interface with the noncrystalline phase such that the structure is weakened; therefore, desired strength or toughness cannot be obtained.

**[0017]** The amorphous Zr alloy of the present invention can be cooled for solidification by various methods, such as a single roll method, a twin-roller method, an in-rotating water melt spinning method, and an atomizing method to provide various forms, such as thin ribbons, filaments, and particles. Also, the alloy of the present invention has a significantly improved glass-forming ability; therefore, it can be formed into a rod or a plate of a given shape by injecting the molten alloy into a mold. For example, using a known metal mold casting process, a bulk of the alloy can be obtained by injecting casting of the melt into metal mold, which is melted in a quartz tube in an Ar atmosphere, the injecting pressure was fixed to be 0.5kg/cm<sup>2</sup>. Furthermore, the amorphous Zr alloy of the present invention has an optimized alloy composition, compared to a conventional amorphous Zr alloy; hence, an excellent glass-forming ability and high strength and toughness can be obtained.

(Examples 1 through 14, Comparisons 1 through 8)

**[0018]** The following describes examples of the present invention.

**[0019]** Rod-shaped samples with a diameter of 5mm and a length of 50mm were prepared using materials having alloy compositions shown in Table 1 by a metal mold casting method. Then, glass transition temperatures (T<sub>g</sub>) and crystallization starting temperatures (T<sub>x</sub>) were measured using a differential scanning calorimeter (DSC); based on the measurements, a range of the supercooled liquid phase (T<sub>x</sub> - T<sub>g</sub>) was calculated. A ratio of a non-crystalline phase contained in a rod-shaped sample by volume (v<sub>f</sub>) was evaluated by comparing the amount of heat generation when the rod-shaped sample crystallized against the amount of heat generation when a completely non-crystallized single rolled sheet crystallized using DSC. In addition, each rod-shaped sample was tested by means of a tensile test, a three-point bending test and the Charpy impact test to measure tension fracture strength (σ<sub>f</sub>), flexural strength (σ<sub>B.f</sub>), i.e., "bending resistance strength", Charpy impact value (E) and fracture toughness (K<sub>1c</sub>).

Table 1

		T <sub>x</sub> -T <sub>g</sub> (K)	V <sub>f</sub> (%)	σ <sub>f</sub> (MPa)	σ <sub>B.f</sub> (MPa)	E (kJ/ m <sup>2</sup> )	K <sub>Ic</sub> (MPa*m <sup>1/2</sup> )	
5	Example 1	Zr <sub>63</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>25</sub> Ti <sub>2</sub>	104	98	1930	2840	125	54
	Example 2	Zr <sub>48.5</sub> Al <sub>7.5</sub> Ni <sub>10</sub> Cu <sub>30</sub> Ti <sub>4</sub>	110	95	2020	3010	136	63
	Example 3	Zr <sub>41</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>45</sub> Ti <sub>4</sub>	108	94	1980	2990	131	60
10	Example 4	Zr <sub>55.5</sub> Al <sub>7.5</sub> Ni <sub>5</sub> Cu <sub>30</sub> Nb <sub>2</sub>	112	97	1890	2700	128	57
	Example 5	Zr <sub>46</sub> Al <sub>10</sub> Ni <sub>10</sub> Cu <sub>30</sub> Nb <sub>4</sub>	125	100	2050	3100	141	66
	Example 6	Zr <sub>46</sub> Al <sub>5</sub> Ni <sub>10</sub> Cu <sub>35</sub> Nb <sub>4</sub>	101	94	1970	2920	128	59
15	Example 7	Zr <sub>55.5</sub> Al <sub>7.5</sub> Ni <sub>5</sub> Cu <sub>30</sub> Pd <sub>2</sub>	109	100	2100	3350	150	69
	Example 8	Zr <sub>56</sub> Al <sub>10</sub> Ni <sub>5</sub> Cu <sub>25</sub> Pd <sub>4</sub>	121	100	2080	3300	144	68
	Example 9	Zr <sub>44</sub> Al <sub>10</sub> Ni <sub>10</sub> Cu <sub>30</sub> Pd <sub>6</sub>	108	100	2210	3510	154	71
	Example 10	Zr <sub>48</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>35</sub> Pd <sub>7</sub>	106	100	2130	3200	139	65
20	Example 11	Zr <sub>51</sub> Al <sub>5</sub> Ni <sub>10</sub> Cu <sub>30</sub> Ti <sub>2</sub> Pd <sub>2</sub>	115	100	2000	2990	123	54
	Example 12	Zr <sub>51</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>35</sub> Ti <sub>2</sub> Nb <sub>2</sub>	118	98	2080	3150	137	63
	Example 13	Zr <sub>43.5</sub> Al <sub>7.5</sub> Ni <sub>10</sub> Cu <sub>35</sub> Nb <sub>2</sub> Pd <sub>2</sub>	113	96	2150	3220	139	63
25	Example 14	Zr <sub>60</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>25</sub> Ti <sub>2</sub> Nb <sub>1</sub> Pd <sub>2</sub>	112	100	1890	2840	120	51
	Comparison 1	Zr <sub>55</sub> Al <sub>10</sub> Ni <sub>5</sub> Cu <sub>30</sub>	104	100	1620	1710	71	44
	Comparison 2	Zr <sub>42</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>40</sub> Ti <sub>8</sub>	88	70	1400	1210	40	22
30	Comparison 3	Zr <sub>42</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>40</sub> Nb <sub>8</sub>	69	51	1260	1170	35	20
	Comparison 4	Zr <sub>42</sub> Al <sub>5</sub> Ni <sub>5</sub> Cu <sub>40</sub> Pd <sub>8</sub>	98	78	1650	1680	73	45
35	Comparison 5	Zr <sub>54</sub> Al <sub>2</sub> Ni <sub>10</sub> Cu <sub>30</sub> Pd <sub>4</sub>	70	55	1180	990	32	18
	Comparison 6	Zr <sub>43.5</sub> Al <sub>12.5</sub> Ni <sub>10</sub> Cu <sub>30</sub> Ti <sub>4</sub>	43	30	670	690	19	11
40	Comparison 7	Zr <sub>41</sub> Al <sub>10</sub> Ni <sub>13</sub> Cu <sub>30</sub> Pd <sub>6</sub>	118	100	1720	1750	88	48
	Comparison 8	Zr <sub>42</sub> Al <sub>5</sub> Ni <sub>20</sub> Cu <sub>30</sub> Ti <sub>3</sub>	65	48	980	1050	36	21

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50  
[0020] As is clearly shown in Table 1, die-cast amorphous alloy materials of Examples 1 through 14 show: a range of the supercooled liquid phase of over 100°C; a ratio of the non-crystalline phase by volume of 90% or higher, providing a large glass-forming ability; tensile strength of 1800 MPa or higher; flexural strength of 2500 MPa or higher; Charpy impact values of 100 kJ/m<sup>2</sup> or higher; fracture toughness values of 50 MPa\*m<sup>1/2</sup> or higher, providing excellent strength and toughness.

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[0021] On the other hand, the alloy of Comparison 1 shows an excellent glass-forming ability in which a cast material with a diameter of 5mm is completely non-crystallized; however, a lack of the M element causes deteriorated mechanical characteristics. Also, the cast materials of Comparisons 2, 3 and 4 contain the M element for the amount exceeding the predetermined 7%; as a result, a range of the supercooled liquid phase and a ratio of the non-crystalline phase by volume are less than 100°C and 90%, respectively, indicating no improvement in mechanical characteristics. Comparisons 5 and 6 do not satisfy the predetermined amount of Al contained, more than 5 % or less than 10% ; hence, the supercooled liquid range and the glass-forming ability are 100°C and 90%, respectively, and the mechanical charac-

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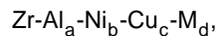
teristics are extremely poor. Further, Comparisons 7 and 8 show no improvement in the mechanical characteristics since the ratio of Ni to Cu, b/c, exceeds the value predetermined in the present invention, 1/3.

### Application of the Present Invention

5  
[0022] As described above, an amorphous Zr alloy of the present invention indicates a supercooled liquid range over 100°C, as well as excellent strength and toughness shown by: tensile strength of 1800 MPa or higher; flexural strength of 2500 MPa or higher; Charpy impact values of 100 kJ/m<sup>2</sup> or higher; fracture toughness values of 50 MPa·m<sup>1/2</sup> or higher. Therefore, the present invention is able to provide a useful amorphous Zr alloy which has a high glass-forming ability and excellent strength and toughness.  
10

### Claims

15 1. An amorphous Zr alloy which has a composition expressed as



20 wherein terms are defined as follows:

M : one or more elements selected from a group consisting of Ti, Nb and Pd; and  
a, b, c, and d : values in atomic % satisfying the following relation,

25 
$$5 \leq a \leq 10$$

30 
$$30 \leq b+c \leq 50$$

$$1/9 \leq b/c \leq 1/3$$

35 
$$0 < d \leq 4$$

when M is Ti or Nb or

40 
$$0 < d \leq 7$$

when M is Pd, and Zr and inevitable impurities being remainder,

45 wherein said amorphous Zr alloy contains a non-crystalline phase of 90% or higher by volume, wherein said amorphous Zralloy has a structure where atoms are densely and randomly packed, but a compound phase with Zr or Al is not formed.

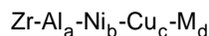
wherein said amorphous Zr alloy has tensile strength of 1800 MPa or higher, flexural strength of 2500 MPa or higher, Charpy impact value of 100 kJ/m<sup>2</sup> or higher, and fracture toughness value of 50 MPa·m<sup>1/2</sup> or higher.

50 2. An amorphous Zr alloy according to Claim 1 which indicates an excellent glass-forming ability with a supercooled liquid range over 100°C (indicated by a difference between the crystallization temperature and the glass transition temperature) and which has a thickness of 1mm or thicker.

### 55 Patentansprüche

1. Amorphe Zr-Legierung mit einer Zusammensetzung, die als

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ausgedrückt wird, wobei die Ausdrücke wie folgt definiert sind:

5  
M: eines oder mehrere Elemente, die ausgewählt sind aus einer Gruppe bestehend aus Ti, Nb und Pd; und  
a, b, c, und d: Werte in Atom-%, die die folgende Beziehung erfüllen:

$$5 \leq a \leq 10,$$

$$30 \leq b+c \leq 50,$$

$$1/9 \leq b/c \leq 1/3,$$

$$0 < d \leq 4 ,$$

wenn M Ti oder Nb ist, oder

$$0 < d \leq 7 ,$$

wenn M Pd ist, und

wobei Zr und unvermeidbare Verunreinigungen der Rest sind,  
wobei die amorphe Zr-Legierung eine nicht-kristalline Phase von 90 Volumen-% oder höher enthält,  
wobei die amorphe Zr-Legierung eine Struktur besitzt, bei der Atome dicht und zufällig gepackt sind, aber eine Verbund- bzw. Komponentenphase mit Zr oder Al nicht gebildet wird,  
wobei die amorphe Zr-Legierung eine Zugfestigkeit von 1800 MPa oder höher, eine Biegefestigkeit von 2500 MPa oder höher, einen Charpy-Kerbschlagbiegewert von 100 kJ/m<sup>2</sup> oder höher, und einen Bruchzähigkeitswert von 50 MPa•m<sup>1/2</sup> oder höher besitzt.

2. Amorphe Zr-Legierung gemäß Anspruch 1, die eine ausgezeichnete Glasbildungsfähigkeit besitzt mit einem supergekühlten bzw. unterkühlten Flüssigkeitsbereich über 100 °C (angezeigt durch eine Differenz zwischen der Kristallisationstemperatur und der Glasübergangstemperatur) und die eine Dicke von 1 mm oder mehr besitzt.

### Revendications

1. Alliage amorphe de Zr le quel a une composition exprimée par la formule  $\text{Zr-Al}_a\text{-Ni}_b\text{-Cu}_c\text{-M}_d$  dans laquelle les termes sont définis de la manière suivante :

M : un ou plusieurs éléments choisis dans un groupe constitué de Ti, Nb et Pd ; et  
a, b, c et d : valeurs en % atomique satisfaisant à la relation suivante

$$5 \leq a \leq 10$$

$$30 \leq b + c \leq 50$$

$$1/9 \leq b/c \leq 1/3$$

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$$0 < d \leq 4$$

lorsque M est Ti ou Nb ou  $0 < d \leq 7$  lorsque M est Pd et

- 5
- Zr et des impuretés inévitables formant le reste,  
dans lequel ledit alliage amorphe de Zr contient une phase non cristalline supérieure ou égale à 90 % en volume,  
10 dans lequel ledit alliage amorphe de Zr a une structure où les atomes sont empilés de façon dense et aléatoire mais où une phase de composé avec Zr ou Al n'est pas formée,  
dans lequel ledit alliage amorphe de Zr a une résistance à la rupture par traction supérieure ou égale à 1800 MPa, une résistance à la flexion supérieure ou égale à 2500 MPa, une valeur de résilience de Charpy supérieure ou égale à 100 kJ/m<sup>2</sup> et une valeur de ténacité à la rupture supérieure ou égale à 50 MPa.m<sup>1/2</sup>.
- 15 **2.** Alliage amorphe de Zr selon la revendication 1 lequel présente une excellente aptitude à former du verre avec une plage de liquide en surfusion dépassant 100°C (indiquée par une différence entre la température de cristallisation et la température de transition vitreuse) et lequel a une épaisseur supérieure ou égale à 1 mm.
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