CONCOMITANT EXTRACTION PROCESS OF NIOBIUM AT THE TITANIUM TETRACHLORIDE PRODUCTION

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Development of the fundamentals of a technology for producing niobium by the concomitant extraction in the process of production of titanium tetrachloride from titanium raw materials. The paper provides data on the amount of niobium contained in the feedstock to produce titanium tetrachloride, the distribution of niobium during processing by products and waste. The forms of phase transformations and transitions of niobium during processing are considered. Schemes for processing niobium-containing chloride materials and technologies for their further application for producing alloys for nuclear power are proposed.

INTRODUCTION

PROPERTIES AND NIOBIUM MINERALS

Niobium is a light gray ductile metal, its density is 8.58 g/cm³, its melting point is 2469 °C, and its boiling point is 4842 °C [1]. Niobium has one stable ⁹³⁵Nb isotope, several radioactive isotopes are artificially produced.

Among all refractory metals, niobium has the highest transition temperature to the state of superconductivity 9.17 K. The thermal conductivity and heat capacity of niobium increase with increasing temperature. The temperature coefficient of linear expansion of niobium with increasing temperature increases in the same way as the electrical resistivity. The above properties depend largely on the purity degree of the metal [2].

Niobium is a rare-earth element; its content in the earth's crust is 2·10⁻³% of the mass [1, 3]. The main minerals of niobium are columbite, laponite, pyrochlore, perovskite, ilmeno-rutile. Fig. 1 shows the dynamics of the production of the main niobium raw materials in the period 2001–2014.

APPLICATION OF NIOBIUM

A successful combination of niobium properties: strength, high heat resistance, corrosion resistance, refractoriness, low temperature range of the transition from plastic to brittle state allow it to be widely used as a high-temperature structural material.

Niobium is used in many key industries, engineering: in the production of steels and alloys, electronics, chemical engineering and nuclear energy (Fig. 3).
The global steel industry accounted for about 90% of global niobium consumption. Niobium is not used in all HSLA and stainless steels; however, in the grades in which it is used, HSLA steels contain about 0.05% niobium by weight, and stainless steels from 0.04 to 0.08%. HSLA steels have become a leading consumer of ferroniobium; structural steel accounted for 45% of global ferroniobium consumption; automotive steel, 23%; steel line, 16%; and stainless steel, 6%. The world's leading producers of ferroniobium were CBMM (Brazil), Mineração Catalão de Goias (Brazil) and IAMGOLD Corp. (Canada), which together accounted for most of the world production of niobium and sold their products in the form of ferroniobium (Fig. 4). World supplies of ferroniobium, measured in the contained niobium, amounted to 53,500 t in 2012, an increase from 52,200 t in 2011 [6].

Expired Vehicles. The niobium content in the ELV can be estimated due to the niobium content in stainless steels, which is in the range of 0.04…0.08%.

The potential sources for the extraction of niobium are mainly steel, but the niobium content is low (< 0.5 wt.%).

One of the directions for obtaining niobium is its associated extraction from products of chlorine processing of titanium-containing materials.

For example, in titanium slag, niobium is initially contained in the form of Nb2O5 oxide. Based on the average content of Nb2O5 in titanium slag – 0.12%, the amount of niobium metal in it can be estimated. Given the molecular weights of niobium and oxygen, the content of niobium metal in titanium slag will be 0.08%, the amount of niobium metal in 1 ton of titanium slag will be 0.81 kg. This means that in the course of processing 20,000 t of titanium slag per year, 16.2 t of niobium are lost with waste. Studying the distribution and creating technology for the extraction of niobium in the production of titanium tetrachloride will increase the complexity of processing titanium-containing raw materials, as well as create an additional source of production of scarce metal.

**EXPERIMENTAL**

Since ilmenite belongs to complex oxides [1], niobium is contained in it in the form of Nb2O5. The content of Nb2O5 in ilmenite ranges from 0.26…1.55% [2]. During ore-thermal reduction smelting of ilmenite concentrate, Nb2O5 is almost completely converted to titanium slag, since niobium is not reduced by carbon. ΔG of the reduction reaction of Nb2O5 with carbon at a temperature of 1500 °C is 1564.6 kJ. The environment for the chlorination of titanium slag is a molten alkali metal salt at a temperature of 700…820 °C.

During chlorination of titanium slag in the melt at temperatures of 700…820 °C in the presence of carbon relatively to Nb2O5, thermodynamically probable course of the following reactions:

\[
2\text{Nb}_2\text{O}_5 + 6\text{Cl}_2 + 3\text{C} = 4\text{NbOCl}_3 + 3\text{CO}_2 \\
(\Delta G = -594.8...-563.6 \text{kJ})
\]

\[
2\text{Nb}_2\text{O}_5 + 10\text{Cl}_2 + 5\text{C} = 4\text{NbCl}_5 + 5\text{CO}_2 \\
(\Delta G = -987.5...-974.9 \text{kJ})
\]

\[
2\text{Nb}_2\text{O}_5 + 6\text{Cl}_2 + 5\text{C} = 4\text{NbCl}_5 + 5\text{CO}_2 \\
(\Delta G = -527.2...-534.2 \text{kJ})
\]

Chlorination of niobium pentoxide in the presence of carbon produces NbOCl3 mixed with NbCl5. This process can be described by the reaction:

\[
\text{Nb}_2\text{O}_5 + 4\text{Cl}_2 + 2\text{C} = \text{NbOCl}_3 + \text{NbCl}_5 + 2\text{CO}_2 \\
(\Delta G = -395.8...-384.6 \text{kJ})
\]

Those, oxytrichloride and niobium pentachloride are transferred into the vapor-gas mixture. The sublimation temperature of NbOCl3 is 400 °C, the boiling point of NbCl5 is 248.3 °C [11].

It was experimentally established that the chlorinator melt assimilates a certain amount of niobium, the value of which is 0.13…0.26 kg of NbOCl3 per 1 t of technical titanium tetrachloride and depends on the chemical composition of the melt. Insignificant assimilation of niobium in the chlorinator is explained by low thermal stability of complexes of niobium oxytrichloride with alkali metal chlorides, as well as its
displacement from complexes by magnesium, iron, and aluminum chlorides.

The temperature in the dust chamber is maintained within 600...500 °C, thus, the compounds of oxytrichloride and niobium pentachloride do not accumulate in the sublimates of the dust chamber.

Next, the niobium compounds in the vapor-gas mixture fall into a dust precipitation chamber with a salt bath (DSS), wherein the temperature is maintained within a 350...300 °C. NbOCl₃ in this temperature range is sublimated and falls into the melt DSS. The low degree of niobium trapping into the salt scrubber is due to the low partial pressure of niobium oxychloride at a temperature of the steam-gas mixture at the outlet of the apparatus.

The Nb content in the DSS melt, in terms of Nb₂O₅, is 0.05%. The content of aluminum trichloride in the DSS melt reaches 40%. At operating temperatures in DSS possible reaction [9]:

\[
\text{NbOCl}_3 + \text{AlCl}_3 = \text{NbCl}_5 + \text{AlOCl} \quad (5) \text{ (AG} = -3.5...-3.2 \text{ kJ)}.
\]

The direct proportional dependence of the niobium content in the slurry of the irrigation scrubber on the AlCl₃ content in the DSS melt is confirmed by practical observations. When topping the sludge-electrolyte mixture into the DSS melt, i.e. with a decrease in AlCl₃ content, a decrease in the Nb content in the slurry is observed.

The resulting mixture of NbOCl₃ and NbCl₅ together with the vapor-gas mixture enters the irrigation scrubber. In the irrigation scrubber, the temperature is maintained no more than 125 °C. The trapping coefficient of niobium in the irrigation scrubber is 98.5%, so the bulk niobium is condensed with circulating slurry. Circulation of pulp along the contour (irrigation scrubber – chlorinator – irrigation scrubber) contributes to the accumulation of niobium oxychloride in solid suspensions of the slurry in the form of a yellow solid impurity [12]. The solids content in the pulp reaches 100 g/l, the Nb content in terms of Nb₂O₅ reaches more than 20%.

In deriving mixture pentaoxychloride and niobium should consider the possibility of further applications. Most methods for producing niobium metal are not related to its reduction from the chloride form, and although hydrogen reduction involves the production of compact niobium rods, it is still a promising technology [11]. Therefore, it is most preferable to consider metallocrystalline reduction with magnesium, since this technology has been sufficiently developed in the production of titanium and zirconium. Most certified zirconium alloys for nuclear power include niobium in the form of an alloying component (Table).

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Type of reactor</th>
<th>Sn</th>
<th>Fe</th>
<th>Ni</th>
<th>Nb</th>
<th>Cr</th>
<th>O</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zircalo-2</td>
<td>BWR, CANDU</td>
<td>1,2-1,7</td>
<td>0,07-0,20</td>
<td>0,03-0,08</td>
<td>-</td>
<td>0,05-0,15</td>
<td>0,125</td>
</tr>
<tr>
<td>Zircalo-4</td>
<td>BWR, PWR, CANDU</td>
<td>1,2-1,7</td>
<td>0,18-0,24</td>
<td>-</td>
<td>0,07-0,13</td>
<td>0,125</td>
<td></td>
</tr>
<tr>
<td>M5</td>
<td>PWR</td>
<td>-</td>
<td>&lt;0,05</td>
<td>-</td>
<td>0,8-1,2</td>
<td>-</td>
<td>0,125</td>
</tr>
<tr>
<td>Zr-2.5Nb</td>
<td>CANDU, RBMK</td>
<td>-</td>
<td>-</td>
<td>2,4-2,8</td>
<td>-</td>
<td>0,125</td>
<td></td>
</tr>
<tr>
<td>ZIRLO</td>
<td>PWR</td>
<td>-</td>
<td>0,09-0,10</td>
<td>-</td>
<td>1,0-1,2</td>
<td>-</td>
<td>0,125</td>
</tr>
<tr>
<td>E110</td>
<td>WWER</td>
<td>-</td>
<td>&lt;0,05</td>
<td>-</td>
<td>0,9-1,1</td>
<td>-</td>
<td>&lt;0,099</td>
</tr>
<tr>
<td>E125</td>
<td>WWER</td>
<td>-</td>
<td>&lt;0,05</td>
<td>-</td>
<td>2,4-2,7</td>
<td>-</td>
<td>&lt;0,099</td>
</tr>
<tr>
<td>E635</td>
<td>WWER</td>
<td>1,1-1,3</td>
<td>0,3-0,4</td>
<td>-</td>
<td>0,9-1,1</td>
<td>-</td>
<td>0,05-0,12</td>
</tr>
</tbody>
</table>

Zirconium-niobium alloys are extensively used as materials for fuel claddings, pressure tubes in CANDU reactors and to a certain extent for other structural components such as fuel channels, in various types of operating nuclear reactors. The properties of zirconium alloys depend on the chemical composition and the heat treatment of the alloy for each type of reactor.

Niobium in Zr-Nb alloys increases the stability of the β-phase, which provides increased corrosion and mechanical resistance of the structural material of the fuel elements of the cladding. Based on this, the authors consider that one of the most promising areas is a possibility of using integrated technology of zirconium-niobium alloys producing for nuclear power engineering method for magnesium thermal reduction.

Most of the known methods of alloying zirconium consist in the technological step of introducing alloying elements in the form of ligatures into the zirconium batch [13]. This operation, along with technological difficulties, does not provide a complete and uniform distribution of alloying elements in the resulting semi-finished products, which necessitates an increase in the number of technological operations to obtain finished products – zirconium alloys [14]. In general, the technology for producing a structural product from zirconium alloys includes in some cases more than 50 operations. Reducing the number of operations or eliminating some technological steps in general, without changing the quality indicators of alloys and products, can significantly reduce the cost of production. One of the episodes, which is decisive in the component of the cost of manufacturing zirconium alloys, is the stage of remelting zirconium with alloying elements to obtain a predetermined chemical composition. The crushed
sponge zirconium is mixed with the crushed ligatures, which are often more expensive than the main material, form the batch into briquettes or press blocks, depending on the method of further processing, and remelt. For one melting, it is not possible to achieve a uniform distribution of alloying elements in the ingot, therefore, double or triple remelting is usually used for the qualitative averaging of composition.

In the case of introducing the technology of doping zirconium in the process of magnesium thermal reduction [15], it is possible to reduce the number of remelts due to the fact that the resulting zirconium sponge will already have a chemical composition with a uniform distribution of alloying elements.

Based on the data on the temperature dependence of vapor pressure of niobium chlorides (Fig. 5) as well as other properties of niobium chloride compounds and the required level of alloy purity, niobium pentachloride is the most suitable product for the production of zirconium-niobium alloys.

However, mixtures of niobium chlorides and oxychlorides formed during the processing of titanium-containing materials cannot be used as an additional alloying component in the process of magnesium thermal production of niobium doped zirconium. Firstly, due to the difference in boiling and sublimation temperatures of oxytrichloride and niobium chlorides, as well as zirconium chlorides. Secondly, a significant amount of oxygen gets into the alloy through niobium oxychloride, which will definitely negatively affect the mechanical and corrosion properties of the alloys.

Thus, the obtained purified niobium pentachloride can be used in the technology of complex alloying of sponge zirconium by the method of magnesium thermal reduction in order to obtain zirconium-niobium alloys for nuclear energy. In view of the fact that zirconium tetrachloride and niobium pentachloride are solids, both their combined purification and chlorination, as well as loading into a reduction reactor, are possible.

The sponge metal block with magnesium chloride residues and unreacted components obtained during the magnesium thermal reduction process is subjected to vacuum thermal treatment, as a result of which a homogeneous compact metal block is formed with a uniform distribution of niobium in the structure. After machining the block with the removal of contaminated bottom and side parts, the metal must be sent for melting. From the point of view of obtaining high-quality ingots, the most acceptable is the method of electron beam remelting (EBM). As a refining operation, the EBM has an advantage over the arc: the metal can withstand the necessary time in the molten state in the mold, which provides deep degassing and refining of the metal from volatile impurities having a higher vapor pressure than the molten metal. At high temperatures and deep vacuum, the refining process in electron beam furnaces proceeds with high speed and high efficiency. The high ductility of alloys remelted in an electron beam furnace allows all types of machining (forging, rolling, drawing, stamping, etc.) to be applied to them under conditions close to conventional metal forming methods [3].

**CONCLUSIONS**

1. A technological scheme has been developed for the processing of titanium-containing materials by chlorination with the removal of niobium oxychloride from the process by its additional chlorination in DSS, which allows the processing of raw materials with a high content of niobium.

2. A solution is proposed for the use of niobium derived from the chlorination process in the production of zirconium-niobium alloys.

3. A technology for the production of purified niobium pentachloride and a mixture of niobium and zirconium chlorides to implement the scheme of magnesium thermal reduction of sponge zirconium alloyed with niobium has been developed.

4. To obtain nuclear zirconium-niobium alloys, a method of electron beam remelting of alloyed sponge zirconium is proposed.

**REFERENCES**

ПОПУТНОЕ ИЗВЛЕЧЕНИЕ НИОБИЯ ПРИ ПРОИЗВОДСТВЕ ТЕТРАХЛОРИДА ТИТАНА

Т.К. Сарсембеков, Т.Б. Янко, С.А. Сидоренко, Н.Н. Пилипенко

Разработаны основы технологии получения ниобия путем попутного извлечения в процессе производства тетрахлорида титана из титанового сырья. В работе представлены данные о количестве ниобия, содержащегося в исходном сырье для производства тетрахлорида титана, распределение ниобия в процессе переработки и выходы получаемого ниобия. Предложены схемы переработки ниобиевых материалов в технологии их дальнейшего применения для получения сплавов для ядерной энергетики.

ПОПУТНОЕ ВИЛУЧЕНИЕ НИОБИЯ ПРИ ВИРОБНИЦТВІ ТЕТРАХЛОРИДУ ТИТАНУ

Т.К. Сарсембеков, Т.Б. Янко, С.А. Сидоренко, М.М. Пилипенко

Розроблено основи технології одержання ниобію шляхом попутного вилучення в процесі виробництва тетрахлориду титану з титанової сировини. У роботі наведено дані про кількість ниобію, що міститься у вихідній сировині для виробництва тетрахлориду титану, розподіл ніобію в процесі переробки за продуктами і відходами виробництва. Розглянуто форми фазових перетворень і переходи ніобію в процесі переробки. Запропоновано схеми переробки хлоридних матеріалів, що містять ніобій, і технології їх подальшого застосування для отримання сплавів для ядерної енергетики.