Effect of Electron Beam Method on Processing of Titanium Technogenic Material

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Abstract: This study reveals the efficiency of the electron beam processing of titanium technogenic material with a high level of impurities and the quality of the obtained metal in correlation to process parameters which are discussed. The influence of the beam power and melting time on the composition variation, morphologies, hardness of metal samples and mass losses is investigated. Based on the different technological parameters, the removal efficiency of impurities is also discussed, and the corresponding experiments are carried out in order to make a comparison. Different thermal process conditions are realized during the single-melt operation. Chemical and metallographic analyses are performed, and the results are discussed. The hardness of the titanium decreases by prolonging the time of the electron beam processing. A maximal overall removal efficiency of 99.975% is seen at 5.5 kW beam power for a 40 min melting time and the best purification of Ti (99.996%) is achieved.

Keywords: titanium technogenic material; electron beam; melting process; removal efficiency

1. Introduction

Titanium and titanium alloys are widely used in aerospace, chemical, bio-medical and other industries due to their weight saving (light in weight), heat resistance, resistance to embrittlement at low temperatures, high corrosion resistance, high tensile strength, low-thermal expansion, and toughness, and with these advantages they have been key materials in high-tech areas [1–7]. They are commonly produced by traditional vacuum arc remelting [8]. However, this process suffers from common problems, such as inhomogeneity of composition and structure, lower material quality, and possible retention of high-density and low-density inclusions in the metal ingot [9]. Electron beam melting technology exhibits extensive application prospects for undesired impurities removal, for solving the problem of inhomogeneity of ingot structure and composition at melting various materials, and also for breaking the length limitation of vacuum arc remelting ingots [10–23]. The microstructure of the obtained ingots after processing is important due to its influence on the mechanical properties. Furthermore, a wider range of initial materials, including technogenic materials, can be used in the electron beam method, while a compacted electrode must be fabricated in vacuum arc remelting. The tremendous kinetic energy of electrons is transformed into heat energy for melting the material after the impact of the beam with the surface of the treated material. In the furnaces, a combination of a high vacuum environment and material superheating (up to 1.3–1.5 of the melting temperature), results in a highly efficient refining process. The electron beam method (EBM) is an effective method that has been utilized for refining refractory materials.
metals [24–27] and alloys [28,29], purifying solar-grade silicon [30,31], obtaining superalloys [18,32,33], high-purity special steels, and so on. We note that the electron beam melting method mentioned in this investigation involves a melting furnace that produces ingots by using an electron beam, which differs from the common adoption for the additive manufacturing method.

The purpose of this study is to examine the effects of the electron beam method on processing titanium technogenic material. The influences of melting power and the refining time on the removal efficiency, composition, the metal microstructure, hardness and material losses are studied in order to increase the efficiency of the EBM, i.e., to improve the composition and quality of the produced metal ingots in the processing of the investigated Ti technogenic material.

2. Materials and Methods

The experiments for electron beam processing of titanium technogenic material were conducted using 60 kW electron beam furnace ELIT 60 (Leybold GmbH, Cologne, Germany, Institute of electronics at the Bulgarian Academy of Sciences, Sofia, Bulgaria) in the laboratory for physical problems of electron beam technologies. The experimental furnace consisted of one electron gun with an accelerating voltage of 24 kV, a melting chamber with a volume of around a cubic meter and with two observation windows, a water-cooled crucible, a circulation water cooling system, and two independent vacuum systems. The melting chamber was evacuated to a pressure of $2 \times 10^{-2}$ Pa and the electron gun chamber was evacuated to a pressure of $3-6 \times 10^{-3}$ Pa. After pre-processing, the raw material was placed in the water-cooled crucible and then the metal was irradiated and heated by an electron beam with a beam spot radius of 10 mm (Figure 1). A copper cylindrical crucible used was with a height of 60 mm and diameter of 50 mm.

![Figure 1. Diagram of the electron beam melting (EBM) process: (1) liquid metal pool; (2) solidifying ingot; (3) water-cooled side wall of the crucible; (4) water-cooled bottom.](image)

The investigated technogenic material (titanium gratings scrap) contained 84.01% Ti and Ir, Fe, Al, Si, and Cu with high concentrations (more than 1%). The other controlled impurities in this study were tungsten, tantalum, and cadmium. In accordance with the requirements, the obtained refined Ti should contain overall impurities of less than 1% (Ti 99%) and the content of iridium and iron should be reduced to a minimum. An investigation under different melting parameters (melting power and refining time) was performed, taking into account the kinetic and thermodynamical limitations depending on the material. The raw materials mass was in the range of 100 g to 150 g.

Experiments were performed with different electron beam powers (with different temperatures) and refining times to examine the effects of melting process parameters on the removal efficiency, material losses, hardness and microstructures of ingots. The raw (initial) material was melted under
single processing at melting powers of 2.5 kW \((T = 1697 ^\circ C)\), 3.5 kW \((T = 1777 ^\circ C)\) and 5.5 kW \((T = 1877 ^\circ C)\) for different lengths of melting time—15 min, 25 min and 40 min.

A stereo and a light microscope allow a study of the topography of the macro or microstructures of polished and etched surfaces at different magnifications. The tops of the ingots were cut along the diameter direction (transverse section) and the cross sections were subjected to a mechanical-chemical polishing and etched.

To reveal the microstructure of Ti specimens, a chemical etching with two reagents was used: reagent 1 (Kroll’s reagent), prepared by 2 mL HF, 3 mL HNO\(_3\) and 100 mL distilled water and reagent 2—a solution composed of 20 mL HF, 20 mL HNO\(_3\) and 60 mL distilled water. The etching time was 80 s or 40 s, respectively. Light microscopy was used to observe the microstructure after electron beam processing and this was related to the hardness by using the Brinell hardness test. The microstructures of the ingot cross section were characterized by a stereo microscope Leica M80 (Leica Microsystems GmbH, Germany) and a light microscope Leica DM2500 (Leica Microsystems GmbH, Germany) with a digital camera Leica EC3 (Leica Microsystems GmbH, Germany). The Leica LAS software (Leica Microsystems GmbH, Germany) was used for image processing.

Brinell hardness was tested for the Ti ingots manufactured at different melting powers (2.5 kW, 3.5 kW and 5.5 kW). A hardened steel ball with a diameter of 2.5 mm (an indenter) with a maximum load of 62.5 kg was used. Three indentations were made in each sample to obtain an average value and a standard deviation.

A series of experiments were performed and using emission spectral analysis (spectrograph PGS-2, Carl Zeiss, Germany) the chemical composition variation of the materials before and after electron beam processing was defined. The composition was determined by averaging the results of five detected positions on the cross section (along the diameter direction), and the standard deviation of the chemical analysis concerning the investigated impurity components was 10 ppm.

The temperature was determined by an optical pyrometer QP-31 using special correction filters. The material losses were estimated using the mass of the initial materials and the ingots after melting that were weighted by electronic balance with a precision of 0.1 g.

3. Results and Discussion

For each processing regime, the changes in the weight, chemical composition and structure of the ingots were controlled. The removal efficiency (degree of refining) was evaluated for each of the controlled impurities and the overall removal efficiency was calculated for each technological regime.

The evaluated values of the overall removal efficiency (removal efficiency of all the impurities of the sample) are presented in Table 1. It is seen that the increase of the electron beam power \((P_b)\) and also the increase of the melting time \((\tau)\) lead to an increase in the overall removal efficiency of the controlled impurities. Table 2 presents data for the material losses \((W_{loss})\), which are mainly due to evaporation but also occur due to splashes.

| \(P_b/\tau\) | 15 min | 25 min | 40 min |
|-----------|--------|--------|--------|
| 2.5 kW | 92.5% | 94.79% | 99.84% |
| 3.5 kW | 96.9% | 97.9% | 99.97% |
| 5.5 kW | 98.71% | 99.8% | 99.975% |

| \(P_b/\tau\) | 15 min | 25 min | 40 min |
|-----------|--------|--------|--------|
| 2.5 kW | 1.7 g | 2.1 g | 3.8 g |
| 3.5 kW | 2.6 g | 3.2 g | 4.4 g |
| 5.5 kW | 3.2 g | 5.4 g | 5.8 g |
In Table 3, data about the chemical composition after melting under the investigated processing regimes is presented. The results obtained show that when the electron beam is operated at a lower power, the removal of Fe, Ir, Cd and Cu depends on the duration of the refining process and this dependence decreases as the melting power is raised. At a beam power of 5.5 kW the refining takes place during the first 15 min and afterwards the impurity concentration remains constant.

| Sample No | Process Parameters | Si   | W   | Ta  | Al  | Cu  | Cd  | Fe  | Ir  |
|-----------|--------------------|------|-----|-----|-----|-----|-----|-----|-----|
| Ti-02     | 2.5, 25 min       | 0.001| 0.02| 0.2 | 0.002| 0.003| 0.3 | 0.3 | 0.01|
| Ti-04     | 2.5, 40 min       | 0.002| 0.01| 0   | 0.002| 0.002| 0.01| 0   | 0   |
| Ti-03     | 3.5, 25 min       | 0    | 0.02| 0.2 | 0.003| 0.003| 0.01| 0.1 | 0.003|
| Ti-06     | 3.5, 40 min       | 0.002| 0   | 0   | 0.003| 0    | 0   | 0   | 0   |
| Ti-01     | 5.5, 15 min       | 0.003| 0.1 | 0.1 | 0.003| 0.0001| 0   | 0   | 0   |
| Ti-05     | 5.5, 40 min       | 0.002| 0   | 0   | 0.002| 0    | 0   | 0   | 0   |
| Concentrations before EBM | 4.94 | 0.1 | 0.03 | 3.28 | 1.52 | 0.1 | 2.36 | 3.66 |

Under the investigated melting regimes, the removal of Al, Si and W is also effective. The removal occurs during the first 25 min and it is slightly dependent on the beam power (Table 3).

It is found that at a beam power of 5.5 kW for a melting time of 40 min, the highest degree of refining of 99.97% is achieved with relatively low material losses (due to evaporation and splashes of metal), with $W_{\text{loss}} = 5.8$ g for 100 g of melted starting material, and with the obtained Ti having a purity of 99.996% (Tables 1–3). Under these processing conditions, the degree of refining of the controlled impurities such as Ir, Fe, W, Ta, Cd and Cu is 100%.

This study investigated also the effects of the melting power and refining time on the ingot morphology, the beam power and melting time on the microstructures of the Ti ingots. Macro-morphology of the initial (raw) Ti technogenic material (Ti gratings) is illustrated in Figure 2. It is seen that the material is not uniform. The chemical analysis of the raw material (before electron beam processing) showed a high content ($\approx 16\%$) of metal impurities (Ir, Fe, Al, Cu, etc.) and the content of titanium is just 84.01% (Table 3).

![Figure 2](image_url)  
(a) Ti material before electron beam processing: (a) 10×, (b) 40×.

In order to examine the effect of the beam power and melting time on the ingot morphology, the morphologies at the central part and near the edge part of ingot cross section were investigated and no significant differences were observed. The Ti sample two-phase macrostructures inspected by stereo microscope are presented in Figure 3. It can be seen that regardless of the different melting conditions—the beam power (and the temperature of overheating) and the isothermal retention—the macrostructure of the samples obtained does not differ significantly. By increasing the temperature (and the melting power) and extending the residence time, the ratio of the $\alpha$ phase (brighter) to $\beta$ phase (darker) increases.
The microstructure of the transformed $\beta \rightarrow \alpha$ phase may be seen more clearly at a higher magnification. Figure 4 shows the microstructures of the ingots obtained at different regimes and cooling to room temperature. They were observed through a light microscope at 400× magnification. An elongated/lamellar type structure [34] is observed on the cross section of the investigated metal samples. This type of structure is typical for Ti, which transforms from $\beta$ phase to $\alpha$ phase upon cooling. The formation of lamellar type structure takes place on the solid/liquid interface in the direction in which the heat is being led away. The $\beta$ to $\alpha$ phase transformation finished between 820–865 °C for cooling rates between 50 °C/min–10 °C/min [35].

The results (Tables 1 and 3) show that by raising the temperature and increasing the melting time, a higher degree of metal refining is obtained. This influences the obtained microstructure and the hardness of the metal. The Ti-02 and Ti-04 sample microstructures (Figure 4a,b) are composed of $\beta$ and $\alpha$ phases in approximately equal ratio, 1:1. These samples were obtained after electron beam processing of the Ti technogenic material at a melting power of 2.5 kW for different lengths of time—for 25 and 40 min, respectively (Figure 4a,b). As a result of the extended electron beam irradiation, which leads to higher metal purity, the Ti-04 sample is characterized by better homogeneity than the microstructure of
the Ti-02 (Figure 3a,b and Figure 4a,b). The different orientation of the lamellae is due to the difference in the rate of heat transfer from the micro volume of the smelt.

![Optical micrographs](image)

**Figure 4.** Optical micrographs (400× magnification) of the titanium ingots produced at different melting powers for different lengths of time: (a) $P_b = 2.5$ kW, $\tau = 25$ min; (b) $P_b = 2.5$ kW, $\tau = 40$ min; (c) $P_b = 3.5$ kW, $\tau = 25$ min; (d) $P_b = 3.5$ kW, $\tau = 40$ min; (e) $P_b = 5.5$ kW, $\tau = 15$ min; and (f) $P_b = 5.5$ kW, $\tau = 40$ min.

Based on the comparison between the microstructures of the Ti-03 and Ti-06 samples (Figure 4c,d), it is shown that the obtained lamellae are significantly larger and the $\alpha$ phase is predominantly observed. These ingots were obtained at a melting power of 3.5 kW. By extending the refining time from 25 min (Ti-03) to 40 min (Ti-06), the metal purity increases to Ti 99.995% and the ingot morphology reaches that of pure titanium (Ti-06).

Experiments conducted at a beam power of 5.5 kW showed that a melting time of 15 min was not sufficient to completely remove the impurities and homogenize the smelt (Ti-01). The content of the impurities in the sample Ti-01 was 0.2016%. The metal purity reached Ti 99.996% with the increase of the melting time to 40 min (Ti-05). The microstructures of the samples (Ti-01, Ti-05) obtained at these melting conditions are shown in Figure 4e,f. Large light lamellae formed by $\alpha$ phase can be seen on the microstructure of the sample Ti-01. They are arranged in an almost parallel manner and surrounded by $\beta$ phase. The microstructure of the sample Ti-05 is a typical Ti structure obtained by slow cooling.

The effects of the melting parameters on the microstructures of ingots (two times re-melted) fabricated via electron beam melting technology were investigated in a $\beta$-type Ti-Mo binary alloy and it was shown that the microstructures consisted of columnar grains at their top sections and they
The microstructures of Ti-6Al-4V resulting from layered manufacturing using the electron beam process were predominantly characterized by acicular (basket-weave) α for solid structures [36–38] or acicular (basket-weave) α and martensite α′ for mesh and foam structures [37] or at the top of the sample [38]. Slow cooling rates favor the formation of similarly aligned α platelets in colonies, together with prior β grain boundary α. Faster cooling rates and higher β stabilizing contents favor a more basket-weave type of structure [35].

The average Brinell hardness (HB_{av}) and standard deviations measured are summarized in Table 4. The results obtained show that the hardness of the titanium samples produced is influenced by melting time rather than by melting power. The average hardness of additive manufactured Ti-6Al-4V blocks decreases due to coarse α –colonies, no (or much less) martensite and no (or much less) dislocations [39]. From Table 4 it can be seen that the hardness of Ti ingots decreases with an increase in melting time that leads to a softer and less heterogeneous material, i.e., an efficient refining of the impurities and their uniform distribution in the structure (the content of Fe, Si, Al, Cu and Ir greatly decreases).

Table 4. Average hardness and standard deviation measured at titanium samples produced at different melting conditions.

| Sample No | Process Parameters | HB_{av} (kg/mm²) | Standard Deviation |
|-----------|--------------------|------------------|--------------------|
| Ti-02     | 2.5 25             | 282              | 28                 |
| Ti-04     | 2.5 40             | 161              | 23                 |
| Ti-03     | 3.5 25             | 292              | 25                 |
| Ti-06     | 3.5 40             | 100              | 7                  |
| Ti-01     | 5.5 15             | 205              | 17                 |
| Ti-05     | 5.5 40             | 120              | 11                 |
| before electron beam melting | 127 | 10 |

4. Conclusions

The effect of the electron beam method on the processing of titanium technogenic material under different melting conditions was examined in the present work. The influence of the melting power and melting time on the composition variation, microstructures of the ingots, hardness and degree of refining of titanium was studied in order to increase the process efficiency. It was found that melting time is an important factor for achieving a higher degree of metal refining and it affects the speed of crystallization during cooling. Hardness is influenced by melting time rather than by melting power. It is shown that with longer melting times, the purity of titanium increases, the hardness decreases, and the obtained ingots have an improved structure.

The maximal overall removal efficiency of 99.975% and the best purification of Ti (99.996%) were achieved at 5.5 kW melting power for a melting time of 40 min with relatively low material losses. It was also found, that under these melting conditions, the degree of refining of some impurities (such as iridium, iron, copper, tungsten, tantalum, and cadmium) was 100%. The results show that the efficiency of electron beam processing of the titanium technogenic material with high level of impurities can be increased in order to improve the quality of the obtained high-purity metal and allow us to formulate requirements on the process conditions.

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