Effect of Magnesium Reduction Process on Oxygen Content of Pickling Niobium Powder

Jingfeng Wang (wangjingfeng@zzrvtc.edu.cn)  
Zhengzhou Railway Vocational & Technical College

Fang Liu  
Zhengzhou University

Yue Zhang  
Zhengzhou University

Qinkui Li  
Zhengzhou University

Research Article

Keywords: pickling niobium powder, oxygen content, oxygen reduction process, magnesium reduction process

DOI: https://doi.org/10.21203/rs.3.rs-526437/v1

License: This work is licensed under a Creative Commons Attribution 4.0 International License. Read Full License
Effect of magnesium reduction process on oxygen content of pickling niobium powder

Wang Jingfeng¹, Liu Fang², Zhang Yue³, Li Qinkui⁴
1. Zhengzhou Railway Vocational and Technical College, Zhengzhou, Henan, CN, 450000
2. Zhengzhou University, Zhengzhou, Henan, CN, 450000
Corresponding author: wangjingfeng@zzrvtc.edu.cn

Abstract: Aiming at the problem of high oxygen content in industrial niobium powder, the oxygen reduction process of high oxygen niobium powder magnesium was studied. Based on the thermodynamic analysis of magnesium thermal reduction of niobium powder, the effects of reduction temperature, magnesium addition, reduction time and reduction atmosphere on oxygen content of pickling niobium powder were studied. The results show that: with the increase of magnesium addition, the oxygen content of pickling niobium powder gradually decreases to a certain value and then remains unchanged. In a certain temperature range (953k-1203k), with the increase of reduction temperature, the oxygen content of pickling niobium powder first decreases and then increases, and the best oxygen content is 356ppm at 1133k; with the extension of reduction time (2-6h), the oxygen content of pickling niobium powder first decreases and then remains unchanged. Finally, the oxygen content of pickled niobium powder is reduced to about 356 ppm at 400% Mg addition, 1133 K reduction temperature and 4 h reduction time.

Keywords: pickling niobium powder, oxygen content, oxygen reduction process, magnesium reduction process

1. Introduction

Niobium and niobium alloys are widely used in electronic industry, aerospace industry, iron and steel industry, automobile industry, low temperature superconductor and medical diagnostic equipment due to their good physical, mechanical and electrical properties, such as good corrosion resistance and ductility, low density, high melting point, high specific strength, good conductivity and thermal conductivity, and small
thermal neutron capture cross section. Niobium powder is an important raw material for the preparation of niobium and niobium alloy. At present, industrial niobium powder is mainly obtained by crushing niobium strip and hydrogenating. Although the particle size of niobium powder is relatively small, which is conducive to the compactness of powder metallurgy products, niobium powder with small particle size has high affinity with oxygen and is easy to adsorb oxygen, resulting in high oxygen content of niobium powder [1-5]. Too high oxygen content in niobium powder will seriously affect the physical, mechanical and electrical properties of niobium, thus restricting the application of niobium and niobium alloys [6-10].

Therefore, to solve the problem of high oxygen content in industrial niobium powder, it is urgent to find a process to reduce the oxygen content in niobium powder. The preparation of low oxygen and fine niobium powder by magnesium reduction of Nb₂O₅ has become a hot spot. Magnesium thermal reduction has the advantages of simple process, low energy consumption and easy separation of reaction products. However, there are few studies on magnesium reduction to reduce the oxygen content of niobium powder [11-15]. After oxygen reduction, the oxygen content of niobium powder is still on the high side, and the complete degree of oxygen binding with magnesium in the process of magnesium reduction and the degree of secondary oxidation on the surface of niobium powder in the process of cooling will affect the oxygen content of niobium [15-22].

In this work, the industrial niobium powder with 4100 ppm oxygen content and 9.8 um Fisher particle size was studied. Firstly, the thermodynamics and kinetics of magnesium thermal reduction of niobium powder were analyzed to select the appropriate reduction temperature range. On this basis, the effects of magnesium addition, reduction time, reduction atmosphere and reduction temperature on oxygen content of pickling niobium powder were studied. Finally, the influence of two-step reduction method on oxygen content of pickling niobium powder was discussed.

2. Experimental

2.1. Materials
Hydrochloric acid、nitric acid、hydrogen peroxide、hydrofluoric acid and Anhydrous ethanol are analytically pure, purchased from Luoyang Haohua Chemical Reagent Co., Ltd (China). The purity of magnesium particles is 99.95%, the particle size is 1-10mm, purchased from Aladdin Reagent Co., Ltd (USA). The niobium powder used in the experiment is provided by Ningxia tantalum industry, with oxygen content of 4100 ppm, particle size of 9.8 um and specific surface area of 0.2 m²/g. The specific chemical composition is shown in Table 1.

| component | Nb   | O    | C    | S    | H    | Fe   | Si   | Ti   |
|-----------|------|------|------|------|------|------|------|------|
| content % | 99.48| 0.41 | 0.0203| 0.005| 0.002| 0.013| 0.057| 0.011|

2.2. Sample preparation

The Nb₂O₅ reaction between magnesium powder and niobium surface (5Mg + Nb₂O₅ = 2Nb + 5MgO) produces niobium and MgO [22]. Theoretically, 1.5g Mg is needed for 1g O. Because magnesium is easy to volatile due to high saturated vapor pressure during sintering, and oxygen is inevitably introduced in reduction atmosphere and operation process, the amount of magnesium should be excessive. According to the experimental basis in the early stage of the experiment, the amount of magnesium added in this experiment was 200%, 300%, 400%, 500% and 600% respectively. The meaning of excess is 2 times, 3 times, 4 times, 5 times and 6 times of the theoretical amount of magnesium needed for magnesium thermal reduction reaction. The specific experimental method is as follows: 15 g niobium powder and corresponding amount of magnesium are evenly mixed and put into the crucible, and then reduced in the hydrogenation and dehydrogenation furnace. After natural cooling to room temperature, the furnace is opened for material collection, and acid washing, drying and oxygen content analysis are carried out.

2.3. Characterization

The contents of oxygen, carbon and other impurities in niobium powder were
determined by oxygen, nitrogen and hydrogen content analyzer (ONH-3000, Gangyannake Testing Technology Co., Ltd) and high frequency infrared carbon sulfur analyzer (CS-3000, Gangyannake Testing Technology Co., Ltd). The metal impurities Fe, Ti and non-metal impurities Si in niobium powder were determined by inductively coupled plasma atomic analyzer (ICP-AES, ThermoFisher Scientific). XRD (6100, Shimadzu) was used to analyze the intermediate products in the reaction process and detect whether there is any biological phase change. The experimental conditions are Cu-Kα ray, scanning speed is 2 °/min, scanning angle is 30 ~ 90 °, working voltage is 40 kV, working current is 30 mA, scanning mode is continuous scanning. Laser particle size analyzer (Mastersizer 3000, Malvern) and average particle size analyzer (WLLP-208a, Dandong Fisher Instrument Co., Ltd.) were used to determine the particle size of niobium powder. SEM and XPS were used to analyze the effect of oxygen behavior on the morphology of niobium powder before and after oxygen reduction and the existing state of oxygen on the surface of original niobium powder. The morphology of niobium powder before and after oxygen reduction was characterized by field emission scanning electron microscope (Quant 250 FEG, USA Fei) and X-ray photoelectron spectroscopy (AXIS Supra, Shimadzu) The existence of oxygen on the surface of high oxygen niobium powder was analyzed.

3. Results and discussion

3.1 Magnesium addition

Different amount of magnesium can affect the sufficient degree of oxide reduction on the surface of niobium powder [23], and then affect the oxygen
content of pickling niobium powder. Moreover, different Mg Addition will also affect the loose degree of reduction products, which will affect the subsequent pickling effect. Excessive mg will cause niobium powder to be wrapped into a block by molten mg during the cooling process. The reduction products with too serious sintering are not conducive to acid pickling in the later stage. Too loose niobium powder may be that magnesium has volatilized before the oxide on Nb surface is fully reduced, which leads to high oxygen content in pickling niobium powder. Therefore, we need to optimize the amount of magnesium.

The effects of 200%, 300%, 400%, 500% and 600% of theoretical magnesium addition on the oxygen content of acid washed niobium powder were studied under the conditions of argon reduction atmosphere, graphite crucible and 1133 K reduction for 4 h. After acid washing, the reduced product was vacuum dried at 333 K for 6 h, and the oxygen content was measured by ONH analyzer. The specific value of oxygen content is shown in Table 2, and the change trend is shown in Figure 1. The macro morphology of the reduction products obtained under the conditions of original niobium powder and different magnesium addition is shown in Figure 2.

| Magnesium addition /% | Original | 200  | 300  | 400  | 500  |
|-----------------------|----------|------|------|------|------|
| oxygen content /ppm   | 4100     | 481  | 412  | 356  | 349  |
According to Table 2, Figure 1 and Figure 2. As can be seen from Figure 2 (b), when the original high oxygen niobium powder is reduced at 200% theoretical amount, (c) 300% theoretical amount, (d) 400% theoretical amount, (e) 500% theoretical amount.
magnesium content and 1133 K for 4 h, the reduction product is loose without any caking phenomenon. There is a little residual magnesium chips on the surface of niobium powder, and the oxygen content of niobium powder obtained after pickling is 481 ppm. When the addition of magnesium is increased to 300%, it can be seen from Figure2 (c) that some of the reduced products are slightly agglomerated, but the cohesion between the powders is very small, and the residual magnesium chips increase. At this time, the oxygen content of acid washed niobium powder is about 412ppm. With the increase of magnesium content to 400%, it can be seen from Figure2 (d) that the reduction product is obviously formed and the hardness of the powder increases, but the powder can be broken with a little force, which will not cause difficulties to the subsequent pickling process. At this time, the oxygen content of the pickling niobium powder is about 356 ppm. When the addition of magnesium is increased to 500%, it can be seen from Figure2 (e) that a large number of reduction products agglomerate. At this time, the oxygen content of the pickling niobium powder is 349 ppm, but the hardness of the reduction products is high, which brings difficulties to the subsequent pickling. It can be concluded that the oxygen content of pickling niobium powder is related to the loose degree and caking of reduced niobium powder. Under the condition of 1133 K reduction for 4 h, the oxygen content of pickled niobium powder decreases to the same level with the increase of magnesium content from 200% to 500%. When the magnesium content is 400%, the loose degree of reduced niobium powder is the best, and the
oxygen content of pickling niobium powder is the lowest, about 356 ppm. It can also be concluded from the loose state of reduction products corresponding to different magnesium additions that the agglomeration phenomenon at 1133 K for 4 h is not due to the self-sintering of niobium powder (without magnesium addition), but due to the solidification agglomeration during the cooling process caused by excessive magnesium.

3.2 Effect of reduction temperature and time on oxygen content of pickling niobium powder

The effects of reduction temperature and time on oxygen content of acid washed niobium powder were studied under the conditions of argon reduction atmosphere, graphite crucible, optimum pickling process and drying process. 953 K, 1053 K, 1093 K and 1133 K were selected to study, and then the effects of different reduction time and magnesium addition amount on oxygen content of pickling niobium powder were studied at a specific reduction temperature. The experimental results are shown in Table 3, 4, 5, 6 and Figures 3, 4, 5,6

Table 3 Oxygen content of pickling niobium powder after reduction at 953 K for 8 h with different magnesium addition

| Magnesium addition /% | 50  | 100 | 150 | 200 | 300 |
|-----------------------|-----|-----|-----|-----|-----|
| oxygen content /ppm   | 3600| 2995| 1600| 890 | 874 |
Figure 3 Oxygen content of pickling niobium powder at 953 K for 8 h with different magnesium addition

It can be seen from Table 3 and Figure 3 that when the reduction temperature is 953 K and the amount of magnesium is 50%, the oxygen content of pickling niobium powder is 3600 ppm; when the amount of magnesium is increased to 200%, the oxygen content of pickling niobium powder is reduced to 890 ppm; when the amount of magnesium is increased to 300%, the oxygen content of pickling niobium powder is basically unchanged, about 874 ppm, indicating that 200% magnesium is enough. The results show that large particles of residual magnesium can be seen on the surface of Nb powder after reduction, which indicates that the magnesium thermal reduction reaction is not carried out thoroughly at this time. It can be seen that this is due to the low reduction temperature and slow reaction speed.

| Magnesium addition /% | 200  | 300  | 400  | 500  |
|-----------------------|------|------|------|------|
| reduction time /h     |      |      |      |      |
| 2                     | 588  | 536  | 541  |      |
| 3                     | 571  | 529  | 538  |      |
| 4                     | 560  | 520  | 530  | 524  |
It can be seen from Table 4 and Figure 4 that the oxygen content of pickled niobium powder is about 530 ppm when the magnesium addition is 300% at 1053 K for 2 h, 3 h, 4 h, 6 h and 8 h reduction; with the extension of reduction time, the oxygen content of pickled niobium powder has no obvious change. When the reduction temperature increases from 953 K to 1053 K, the oxygen content of acid washed niobium powder decreases about 350 ppm. It can be seen that with the increase of reduction temperature, the reaction rate of magnesium reducing Nb$_2$O$_5$ is accelerated, and the magnesium thermal reduction reaction is relatively sufficient. When the reduction time is prolonged, the oxygen content of acid washed niobium powder decreases less, which indicates that the reduction temperature at 1053 K is still low and the reaction speed is slow. The results show that the oxygen content of pickling niobium powder with 200% magnesium addition is significantly higher than that with 300% magnesium addition, but the oxygen content of pickling niobium powder with 400% and 500%
magnesium addition has little change compared with that with 300%. It shows that 300% excess magnesium is needed to meet the magnesium consumption in the reduction process at 1053 K, which is higher than that at 953 K. This is because the volatilization loss of magnesium increases with the increase of reduction temperature. At the same time, the addition of magnesium can not fully guarantee the oxygen content at 1053 K reduction.

Table 5 Oxygen content of pickling niobium powder at 1093 K with different magnesium addition and reduction time (PPM)

| Magnesium addition /% | Reduction time /h | 200   | 300   | 400   | 500   |
|-----------------------|-------------------|-------|-------|-------|-------|
| 2                     |                   | 560   | 490   | 481   | 479   |
| 3                     |                   | 524   | 461   | 460   | 463   |
| 4                     |                   | 508   | 433   | 421   | 425   |
| 5                     |                   | 481   | 409   | 406   | 408   |
| 6                     |                   | 478   | 406   | 403   | 404   |

Figure 5 Effect of magnesium addition and reduction time on oxygen content of pickling niobium powder at 1093 K

It can be seen from Table 5 and Figure 5 that when magnesium reduction temperature is 1093 K, magnesium addition is 400% and reduction time is 2 h, the oxygen content of pickling niobium powder is 481 ppm; when reduction time is 5 h, the oxygen content of pickling niobium powder is 406 ppm, which may
because the magnesium reduction reaction becomes more and more abundant with the extension of reduction time. When the reduction time is extended to 6 h, the oxygen content of the acid washed niobium powder is about 403 ppm. Therefore, when the reduction time increases from 2 h to 6 h, the oxygen content of acid washed niobium powder first decreases and then remains unchanged, and the lowest oxygen content of acid washed niobium powder is 406 ppm after 5 h reduction. When the reduction temperature increases from 1053 K to 1093 K, the oxygen content of pickled niobium powder decreases from 530 ppm to 406 ppm, which is also due to the higher reduction temperature, the faster reduction rate of Nb$_2$O$_5$ by magnesium and the more sufficient reduction reaction. The oxygen content of pickling niobium powder is obviously lower than that of 200% magnesium addition when magnesium addition is over 300%. When magnesium addition is increased to 400%, the oxygen content of pickling niobium powder is basically unchanged. This shows that 300% magnesium addition is enough for magnesium consumption at 1093 K reduction temperature. At 1093 K, the optimal magnesium reduction process parameters are 400% magnesium addition and 5 h pickling time. At this time, the oxygen content of pickling niobium powder is 406 ppm.

Table 6 Oxygen content of pickling niobium powder at 1133 K with different magnesium addition and reduction time (PPM)

| Magnesium addition /% | reduction time /h | 200 | 300 | 400 | 500 | 600 |
|-----------------------|-------------------|-----|-----|-----|-----|-----|
| 2                     |                   | 530 | 460 | 445 | 450 |     |
| 3                     |                   | 505 | 462 | 420 | 418 |     |
| 4                     |                   | 481 | 412 | 356 | 349 |     |
| 5                     |                   |     |     | 360 | 337 | 340 |
| 6                     |                   |     |     | 357 | 356 | 353 |
Figure 6 Effect of magnesium addition and reduction time on oxygen content of pickling niobium powder at 1133 K

It can be seen from Table 6 and Figure 6 that when magnesium reduction temperature is 1133 K, magnesium addition is 400% and reduction time is 2 h, the oxygen content of pickling niobium powder is 445 ppm; when reduction time is 4 h, the oxygen content of pickling niobium powder is 356 ppm; when reduction time is 5 h and 6 h, the oxygen content of pickling niobium powder is 360 ppm and 357 ppm respectively. Therefore, when the reduction temperature is 1133 K and the amount of magnesium is 400%, the oxygen content of pickling niobium powder decreases first and then remains unchanged when the reduction time increases from 2 h to 6 h. The reason for the decrease is that with the extension of reduction time, the magnesium reduction reaction becomes more and more sufficient. When the reduction temperature increases from 1093 K to 1133 K, the oxygen content of the acid washed niobium powder decreases from 416 ppm to 356 ppm, and the reduction time is 1 h shorter than that corresponding to 1093 K. The oxygen content of pickling niobium powder is obviously lower than that of pickling niobium powder corresponding to 200% and 300% magnesium addition when the magnesium addition is 400%. When the magnesium addition is increased to 500% and 600%, the oxygen content of pickling niobium powder is basically unchanged and the reduction product is
prone to caking. Under the condition of 1133 K reduction temperature, the optimum magnesium reduction process parameters are as follows: 400% excess magnesium addition, 4 h reduction time and 356 ppm oxygen content.

From the above effects of magnesium reduction temperature of 953 K, 1053 K, 1093 K and 1133 K on the oxygen content of pickling niobium powder, it is known that when the reduction temperature increases from 953 K to 1133 K, the oxygen content of pickling niobium powder decreases from 890 ppm to 356 ppm, because with the increase of reduction temperature, the magnesium thermal reaction rate accelerates and the reaction becomes more and more sufficient. Continue to increase the reduction temperature, whether the oxygen content will continue to decrease? Continue to explore the effect of reduction temperature 1203 K on the oxygen content of pickling niobium powder, and the results are shown in Table 7 and Figure 7.

Table 7 Oxygen content of pickling niobium powder at 1203 K with different magnesium addition and reduction time (PPM)

| Magnesium addition /% | Original niobium powder | 300   | 400   | 600   |
|-----------------------|--------------------------|-------|-------|-------|
| Restore time /h       |                          |       |       |       |
| 2                     | 4100                     | 462   | 431   | 419   |
| 3                     | 4100                     | 489   | 440   | 426   |
| 4                     | 4100                     | 551   | 455   | 442   |
**Figure 7 Effect of magnesium addition and reduction time at 1203 K on oxygen content of pickling niobium powder**

It can be seen from Table 7 and Figure 7 that the oxygen content of pickling niobium powder corresponding to 300%, 400% and 600% magnesium addition is 462 ppm, 431 ppm and 419 ppm respectively at 1203 K reduction for 2 h; when the reduction temperature is increased to 3 h, the oxygen content of pickling niobium powder corresponding to 300%, 400% and 600% magnesium addition is increased to 489 ppm, 440 ppm and 426 ppm respectively; the reduction temperature is further increased to 4 h. The oxygen content of pickling niobium powder increased to 551 ppm, 455 ppm and 442 ppm respectively. It can be seen that the oxygen content of pickling niobium powder at 1203 K is generally higher than that at 1133 K. Although the addition of magnesium increased from 300% to 600%, the oxygen content of pickling niobium powder decreased, but the reduction products with excess magnesium of 400%, 500% and 600% had serious caking phenomenon, and the hardness was relatively large. A large amount of magnesium oxide and magnesium could be seen, which brought inconvenience to the subsequent pickling process. It can be seen that the high reduction temperature at 1203 K is not conducive to the reduction of oxygen content in pickling niobium powder. Therefore, the reduction process is optimized as follows: 1133 K reduction temperature, 400% excess magnesium addition for 4 h, and the oxygen content of acid washed niobium powder is as low as 356 ppm.
Figure 8 XRD pattern of reduction products of magnesium at 1203 K for 2 h

It can be seen from Figure 8 that magnesium niobate mg3nbo11 is formed after reduction at 1203 K for 2 h. As a result, the oxygen content of 1203 K reduced niobium powder after pickling is higher than that of 1133 K reduced niobium powder [24-26]. It is known from that reduction temperature, reduction time and magnesium addition all have great influence on oxygen content of pickling niobium powder. With the increase of reduction temperature from 953 K to 1133 K, the oxygen content of pickling niobium powder decreases from 890 ppm to 356 ppm, which is due to the increase of reduction temperature, the reduction rate of magnesium is accelerated, and the reduction reaction is more and more sufficient; with the increase of reduction temperature to 1203 K, the oxygen content of pickling niobium powder increases, which is due to the presence of acid insoluble magnesium niobate in the reduction product.

The effect of reduction time on oxygen content of pickling niobium powder was studied under the condition of sufficient magnesium addition. At
1053 K for 2-8 h, the oxygen content of pickled niobium powder changed little, about 530 ppm; at 1093 K and 400% Mg for 2-6 h, the oxygen content of pickled niobium powder decreased from 481 ppm in 2 h to 406 ppm in 5 h; at 1133 K for 6 h, the oxygen content of pickled niobium powder remained unchanged, about 403 ppm K. The oxygen content of acid washed niobium powder decreased from 445 ppm in 2 h to 356 ppm in 4 h after 2-6 h reduction with 400% Mg Addition. The oxygen content of acid washed niobium powder changed from 356 ppm to 357 ppm after 6 h reduction with increasing reduction time. Therefore, at the reduction temperature of 1093 K and 1133 K, the oxygen content of pickling niobium powder first decreases and then remains unchanged with the extension of reduction time.

At the same reduction temperature and time, the oxygen content of pickling niobium powder first decreases and then remains unchanged with the increase of magnesium content. The results show that the reduction products are relatively loose when the magnesium addition is 200% and 300% too much; when the magnesium addition is 400% too much, the reduction products obviously agglomerate, but can be broken with a little force, the reduction products are just loose, and the oxygen content of pickling niobium powder is the lowest; when the magnesium addition is 500%, the reduction products agglomerate more, the hardness increases, and the oxygen content of pickling niobium powder is basically the same as that of 400% magnesium. The addition amount is the same. Therefore, the optimal reduction process
parameters of magnesium reduction were optimized as follows: reduction
temperature 1133 K, reduction time 4 h, excess magnesium addition 400%.

3.3 Effect of reducing atmosphere and crucible material on oxygen content and
phase composition of pickling niobium powder

In order to explore the influence of reduction atmosphere and material of
charging Crucible on oxygen content of acid washed niobium powder, the
reduction process of 1133 K reduction temperature, 400% magnesium addition
and 4 h reduction was adopted to study the influence of hydrogen, argon and
vacuum reduction atmosphere, graphite crucible and Nickel Crucible on oxygen
content of acid washed niobium powder. As shown in table 8, the oxygen content
of acid washed niobium powder obtained by graphite crucible and Nickel
Crucible in vacuum and argon reduction atmosphere is given respectively.

| Reducing atmosphere | Crucible material | Vacuum atmosphere | Argon atmosphere |
|---------------------|-------------------|-------------------|------------------|
| Nickel Crucible     | 764               | 371               |
| Graphite crucible   | 802               | 365               |

It can be seen from table 8 that the oxygen content of acid washed niobium
powder obtained by Nickel Crucible and graphite crucible charging reduction in
argon state is basically the same, about 365 ppm. But in vacuum, the oxygen
content of niobium powder is 764 ppm in Nickel Crucible and 802 ppm in
graphite crucible. The oxygen content of niobium powder by vacuum reduction
and pickling with two kinds of crucible is higher than that of argon protection.
The main reasons are as follows: first, there will inevitably be air infiltration in
the hydrogenation and dehydrogenation furnace in the vacuum state. In the
process of vacuum cooling, when the cooling temperature is 573 °C Above K,
the surface of niobium powder with high activity will be oxidized to niobium
oxide again, so niobium powder will be oxidized again after oxygen reduction;
secondly, with the volatilization of magnesium in vacuum and the decrease of temperature in the process of cooling, the oxygen reduction ability of niobium powder gradually decreases. However, in the high purity argon environment, the pressure in the furnace has been maintained at about 0.1 MPa, and the outside air is not easy to enter the furnace, so it is easier to obtain niobium powder with lower oxygen content in argon atmosphere than in vacuum.

The reason why the oxygen reduction effect of niobium powder using Nickel Crucible in vacuum is better than that using graphite crucible is that the nickel crucible has a cover in the reduction process, and the magnesium vapor always fills the whole Nickel Crucible; while the graphite crucible has no cover, the magnesium vapor gradually volatilizes faster, which is not conducive to the full reduction of niobium powder.

Niobium powder has good hydrogen absorption properties, and can easily react with hydrogen to form brittle niobium hydride. Niobium has the fastest formation rate of NbH at about 633 K, and NbH begins to dehydrogenate at about 923 K. In order to investigate the hydrogen absorption of niobium powder in the reduction process in hydrogen atmosphere, the reduction experiment of niobium powder in 1133 K and hydrogen atmosphere was carried out, and the reduction products were analyzed by XRD diffraction. The results are shown in Figure 9.
It can be seen from figure 9 that in hydrogen atmosphere, the diffraction peak position of niobium powder after magnesium reduction is consistent with that of 80-2282-nbh0.95, indicating that niobium powder is completely transformed into nbh0.95 after reduction in hydrogen atmosphere. Therefore, it is not feasible to reduce niobium powder in hydrogen state. At high temperature, nickel is easy to form low melting point eutectic with magnesium, resulting in the increase of nickel content in pickling niobium powder. The carbon content of niobium powder can be increased by 70 ppm when carbon crucible is used. Therefore, it is necessary to re select the crucible material in the future research.

3.4 Effect of oxygen reduction by magnesium on particle size, morphology and solid solution Oxygen of niobium powder

In order to investigate the effect of reduction and pickling on the particle size and morphology of niobium powder. The particle size and morphology of the original niobium powder with high oxygen and the original niobium powder were studied at 1133 K reduction temperature, 400% magnesium addition and 4 h reduction. Figure 10 is the SEM micrograph of original niobium powder and oxygen reducing niobium powder, and Figure 11 is the particle size distribution of niobium powder before and after oxygen reducing.
Figure 10 SEM pictures of niobium powder before and after oxygen reduction:

(a) low rate before oxygen reduction, (b) low rate after oxygen reduction, (c) high rate before oxygen reduction, (d) high rate after oxygen reduction

It can be seen from Figure 10 (a) that the larger particle size of niobium powder before oxygen reduction is about 40 µm, and the smaller one is about 0.5 µm. It can be seen from Figure 10 (c) that the shape of niobium powder before oxygen reduction is irregular and sharp, and there are many fine particles about 0.2-2 µm adsorbed on the surface of large particles. It can be seen from figure 10 (b) that the minimum particle size of niobium powder after oxygen reduction is about 5 µm, and the maximum particle size is still about 40 µm, with no obvious change in morphology compared with that before deoxidation. Therefore, in the process of reducing and pickling, the oxygen reduction of niobium powder basically does not change the morphology of niobium powder, but it can be seen from Figure 10 (d) that the niobium powder of fine particles is greatly reduced after oxygen reduction, and it can be seen that the fine particles are dissolved or washed away.
by acid in the process of pickling and water washing. The average particle size of original niobium powder and niobium powder after oxygen reduction are 9.8 um and 10.2 um respectively; the particle size distribution of niobium powder before and after oxygen reduction is shown in Figure 11.

![Particle size distribution](image)

Figure 11 particle size distribution of niobium powder before and after oxygen reduction: (a) before pickling, (b) after pickling

It can be seen from figure 11 that the oxygen reduction treatment causes the loss of very fine particles in niobium powder. According to calculation, the loss rate of niobium powder after oxygen reduction treatment is about 0.3%.

In order to explore whether oxygen reduction treatment will affect the dissolved oxygen in niobium powder. XRD analysis was carried out on the acid washed niobium powder at 1133 K reduction temperature, 400% magnesium addition and reduction for 4 h. the diffraction pattern is shown in Figure 12.
It can be seen from Figure 12 that there is no new phase formed after acid pickling of Nb powder reduced for 4 h at 1133 K reduction temperature and 400% Mg addition, but only a single Nb phase. The peak shape of the diffraction peak is basically unchanged without obvious broadening. However, the crystal plane spacing of niobium powder (110), (200), (211), (220) before oxygen reduction tends to decrease compared with that before oxygen reduction. For example, the (110) plane spacing D decreases from 2.3387 nm before oxygen reduction to 2.3357 nm after oxygen reduction, and the peak position angle shifts to the right by about 0.8 °C. It can be seen that oxygen reduction not only reduces the surface oxygen of niobium powder, but also reduces the solid dissolved oxygen. This is due to the lattice distortion of niobium caused by oxygen dissolved in the niobium lattice. With the decrease of oxygen content in niobium powder, the degree of solid solution of oxygen in niobium decreases, which leads to the decrease of lattice distortion, that is, the decrease of crystal plane spacing D. From the Bragg equation,

\[ 2d \sin \theta = n\lambda \]

(N and \( \lambda \) are the reflection order and X-ray wavelength respectively, D is the crystal plane spacing, and \( \theta \) is the angle between the incident ray or reflected ray and the reflecting surface) When D decreases, \( \theta \) increases. Therefore, the peak position of Nb powder after oxygen reduction treatment moves to the right relative to the original Nb powder.

4. Conclusion

In this paper, the effects of magnesium addition, reduction temperature, reduction time, crucible material and reduction atmosphere on the oxygen content of niobium powder after pickling were studied. The main conclusions are as follows: Reduction temperature, magnesium addition and reduction time
all have great influence on the oxygen content of acid washed niobium powder. When the reduction temperature increases from 953 K to 1133 K, the oxygen content of niobium powder decreases from 890 ppm to 356 ppm. When the reduction temperature increases to 1203 K, the oxygen content of niobium powder increases. When the amount of magnesium is enough, the effect of reduction time on niobium powder is different with the reduction temperature. When the reduction temperature is 1053 K and the reduction time is 2-8 h, the oxygen content of pickled niobium powder changes little, about 530 ppm; when the reduction time is 2-6 h at 1093 K and 1133 K, the oxygen content of pickled niobium powder first decreases and then remains unchanged, from 481 ppm to 406 ppm and from 445 ppm to 357 ppm, respectively. At the same reduction temperature and time, the oxygen content of pickling niobium powder gradually decreases to a certain value with the increase of magnesium addition, and remains unchanged. In addition, in argon atmosphere and graphite crucible, the oxygen content of niobium powder obtained by oxygen reduction is better than that in vacuum atmosphere. The average particle size and morphology of niobium powder after oxygen reduction were observed. Compared with that before oxygen reduction, there was almost no change in niobium powder, but a small amount of fine niobium powder was removed. After oxygen reduction, the peak angle of niobium powder shifts to the right by about 0.8 ℃, resulting in the decrease of crystal plane spacing. Finally, when the reduction temperature is 1133 K, the excess amount of magnesium is 400%, and the reduction time is 4 h, the oxygen content of the pickling niobium powder is reduced from 4100 ppm to 356 ppm, which greatly reduces the oxygen content of the industrial niobium powder.

Declaration of interests:
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported
in this paper.

Reference:
[1] Eckert J. Niobium and Niobium Compounds[M]. Ullmann's Encyclopedia of Industrial Chemistry, 2000
[2] Florêncio O, Silva P S, Grandini C R. Diffusion Phenomena of the Oxygen and Nitrogen in Niobium by Mechanical Spectroscopy[J]. Defect & Diffusion Forum, 2010, 297-301: 1346-1353
[3] Yamanaka S, Fujita Y, Uno M, et al. Influence of interstitial oxygen on hydrogen solubility in metals[J]. Journal of Alloys and Compounds, 1999, 293-295: 0-51
[4] Trivisonno J, Washick J, Keck M J, et al. Ultrasonic attenuation of longitudinal and shear waves in the normal and superconducting states of niobium[J]. Journal of Low Temperature Physics, 1971, 4(1): 97-109
[5] Parraud S, Hubert P, falzgraf L G, et al. Stabilization and Characterization of Nanosized Niobium and Niobium Oxide Sols-Optical Applications for High-Power Lasers[J]. Journal of the American Ceramic Society, 75(8): 4
[6] Okabe T H, Iwata S, Imagunbai M, et al. Production of niobium powder by preform reduction process using various fluxes and alloy reductant[J]. ISIJ International [0915-1559] Iwata, 2004, 4(2): 285-293
[7] Yuan B, Okabe T H. Production of fine niobium powder by preform reduction process using Mg–Ag alloy reductant[J]. Cheminform, 2007, 443(1-2): 0-80
[8] Orlov V M, Sukhorukov V V. Magnesium-thermal preparation of niobium powders[J]. Russian Metallurgy, 2010, 2010(3): 168-173
[9] T Satish Kumar. Preparation of Niobium Metal Powder by Two-Stage Magnesium Vapor Reduction of Niobium Pentoxide[J]. Hindawi Publishing Corporation Journal of Metallurgy, 2013, 80: 1-8
[10] Nico C, Monteiro T, M.P.F. Graça. Niobium Oxides and Niobates physical properties: review and prospects[J]. Progress in Materials Science, 2016, 80: 1-37
[11] Naito K, Matsui T. Review on phase equilibria and defect structures in the niobium-oxygen system[J]. Solid State Ionics, 1984, 12(none): 125-134
[12] Wu A T. Investigation of oxide layer structure on niobium surface using a secondary ion mass spectrometry[J]. Physica C, 2006, 441(1-2): 79-82
[13] Hong C I, Lim J W. Efficacy of Acid Cleaning on the Deoxidation of Titanium Powder Using Calcium [J]. Korean Journal Of Metals And Materials, 2018, 56(3): 205-209
[14] Lai H, Huang L, Gan C, et al. Enhanced acid leaching of metallurgical grade silicon in hydrofluoric acid containing hydrogen peroxide as oxidizing agent[J]. Hydrometallurgy, 2016, 164: 103-110
[15] Salih Aydoğan, Aras A, Gökhan Uçar, et al. Dissolution kinetics of galena in acetic acid solutions with hydrogen peroxide[J]. Hydrometallurgy, 2007, 89(3): 189-195
[16] Lu D L, Hu Y Y, Lin T, et al. Refining of Metallurgical Grade Silicon by Acid Leaching[J]. Advanced Materials Research, 2010, 156-157: 566-569
[17] Xie K, Mai Y, Ma W, et al. The Removal of Titanium from Metallurgical Silicon by HF–HCL Leaching[J]. Metallurgist, 2013, 57(7-8): 633-638
[18] Lai H, Huang L, Xiong H, et al. Hydrometallurgical Purification of Metallurgical Grade Silicon with Hydrogen Peroxide in Hydrofluoric Acid[J]. Industrial & Engineering Chemistry Research, 2017, 56(1): 311-318
[19] Banchothdhevakul W, Matsui T, Naito K. Vaporization Study on Vanadium-Oxygen Solid Solution by Mass Spectrometric Method[J]. Journal of Nuclear Science and Technology, 1986, 23(10): 873-882
[20] Mueller R, Bobeth M, Brumm H, et al. Kinetics of nanoscale structure development during Mg-vapour reduction of Niobium oxide[J]. International journal of materials research [1862-5282] Bobeth, 2007, 98(11): 1138-1145
[21] Orlov V M, Kryzhanov M V. Deoxidation of the Niobium powder produced by self-propagating high-temperature synthesis[J]. Russian Metallurgy, 2014, 2014(3): 191-194
[22] Livingston J D, Cahn J W. Discontinuous coarsening of aligned eutectoids[J]. Acta Metallurgica, 1974, 22(4): 495-503
[23] Kyunsuk Choi, Hanshin Choi. Effect of magnesium on the phase equilibria in magnesiothermic reduction of Nb2O5[J]. Materials Letters, 2016, 183: 151-155
[24] Pukazhselvan D, Otero-Irurueta G, Pérez J, et al. Crystal structure phase stoichiometry and chemical environment of MgxNbyOx+y nanoparticles and their impact on hydrogen storage in MgH2[J]. International Journal of Hydrogen Energy, 2016, 41(27): 1709-1715
[25] You Y C, Park H L, Song Y G, et al. Stable phases in the MgO-Nb2O5 system at 1250°C[J]. Journal of Materials Science Letters, 1994, 13: 1487-1489
[26] Ananta S. Phase and morphology evolution of magnesium niobate powders synthesized by solid-state reaction[J]. Materials Letters, 2004, 58(22-23): 0-2786
Figure 1

Effect of magnesium addition on oxygen content of pickling niobium powder at 1133 K for 4 h
Figure 2

Different Mg Addition on morphology of reduced niobium powder
Figure 3

Oxygen content of pickling niobium powder at 953 K for 8 h with different magnesium addition
Figure 4

Effect of magnesium addition and reduction time on oxygen content of pickling niobium powder at 1053 K
Figure 5

Effect of magnesium addition and reduction time on oxygen content of pickling niobium powder at 1093 K
Figure 6

Effect of magnesium addition and reduction time on oxygen content of pickling niobium powder at 1133 K
Figure 7

Effect of magnesium addition and reduction time at 1203 K on oxygen content of pickling niobium powder
Figure 8

XRD pattern of reduction products of magnesium at 1203 K for 2 h
Figure 9

XRD diffraction pattern of Nb powder reduced in hydrogen medium
Figure 10

SEM pictures of niobium powder before and after oxygen reduction: (a) low rate before oxygen reduction, (b) low rate after oxygen reduction, (c) high rate before oxygen reduction, (d) high rate after oxygen reduction
Figure 11

Particle size distribution of niobium powder before and after oxygen reduction: (a) before pickling, (b) after pickling.

Figure 12

Deoxidized Nb powder

Raw Nb powder
XRD diffraction pattern of niobium powder before and after deoxidization