NiAl INTERMETALLIC PREPARED WITH REACTIVE SINTERING AND SUBSEQUENT POWDER-METALLURGICAL PLASMA-SINTERING COMPACTION

REAKCIJSKO SINTRANJE IN ZGOŠČEVANJE S PLAZEMSKIM SINTRANJEM NiAl INTERMETALNE ZLITINE

Alena Michalcová1, Dalibor Vojtěch1, Tomáš František Kubatík2, Pavel Novák1, Petr Dvořák1, Petra Svobodová1, Ivo Marek1
1University of Chemistry and Technology, Department of Metals and Corrosion Engineering, Prague, Technická 5, 166 28 Prague 6, Czech Republic
2Institute of Plasma Physics AS CR, v. v. i., Za Slovankou 1782/3, 182 00 Prague 8, Czech Republic
michalca@vscht.cz

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This paper proposes a novel method for powder-metallurgy preparation of compact NiAl intermetallics. In the first step, the NiAl powder is prepared with the reactive-sintering procedure. The porous NiAl product of the SHS reaction is milled to a fine powder and consequently compacted by SPS processing. The compaction of powder metals and alloys is a very difficult field due to the need of preserving the unique properties of the initial materials. One of the few possible methods of a successful compaction is plasma sintering. To describe detailed structures of powder-metallurgy materials, it is necessary to use advanced microscopy methods such as SEM and TEM. In this study, the structure of a NiAl intermetallic compound is described. The material was first produced, with reactive sintering, from pure elements. Subsequently, the NiAl porous master alloy was milled and compacted with the spark-plasma-sintering (SPS) technique. The particle size of the NiAl powder was comparable to the grain size of the compacted material, which exhibited a low porosity. It was proved that the interconnection of the NiAl particles is made by a thin layer of nanocrystalline oxides.

Keywords: SPS, intermetallics, powder metallurgy

1 INTRODUCTION

Like many other transition metal aluminides, nickel aluminide exhibits properties that are very interesting for industrial utilization. These are a high melting point (1638 °C), a low density (5.95 g/cm³), a high thermal conductivity (70 W m⁻¹K⁻¹), an excellent corrosion resistance1,2 and a very good wear resistance.3 These properties allow intermetallics to be used in the applications where metallic and ceramic materials fail. In addition, nickel aluminide is easily produced in atmospheric air with a self-propagating high-temperature synthesis (SHS)2,4–5 even when pre-pressed into a green body.4 This makes the Ni-Al system to be an ideal model for the study of a possible powder preparation using metallurgical methods based on SHS.

The intermetallic materials usually exhibit good mechanical properties at elevated temperatures, but unfortunately, they seem to be quite brittle at room temperature. When decreasing the grain size of a material, this factor limiting its utilization can be solved. One of the promising ways is to produce fine-grained intermetallics with a two-step powder-metallurgy method: in the first step, an intermetallic is formed with the SHS procedure; then it is milled to a very fine powder and compacted with the spark-plasma-sintering (SPS) procedure. The advantage of the SPS process lies in extremely short sintering times, due to which there is almost no grain coarsening.6–9 The SPS method is well described for ceramics, but for metals and especially for intermetallics, the description of the process is still being formed.6–9

The spark-plasma sintering method has been very popular in the last two decades, mainly in the field of compaction of ceramics. It is an ideal tool for obtaining homogenous nanocrystalline bulk materials with a high density, i.e., fine-grained ceramics, thermo-electric semi-
conductors and biomaterials. Compared to the other compaction methods (cold and hot isostatic pressing), the SPS is distinguished by a low overall sintering temperature, short sintering times and better properties of the prepared bulk materials.

Using the SPS method, many successes were achieved in the fields of increasing the superplasticity of ceramic materials, improvement of magnetic properties, reduction of the amount of impurities segregated at the grain boundaries, improvement in the binding quality and many others. From the historical point of view, the first machine comparable to SPS was built in Germany, as reported in reference. In 1933, in the USA, F. Taylor was awarded a patent for the first resistance-sintering method used for sheets.

Basically, the SPS method for sintering materials can be divided into four generations: the first SPS was built in Japan (in 1962) and called spark sintering (SS). The next generation can be described as plasma-activated sintering (PAS), followed by spark-plasma sintering (SPS), while the fourth and currently the last generation is the one described in 12.

The study of a NiAl alloy prepared with SPS can be used, in future, as a milestone for the preparation of NiAl-based composites. Preparation conditions can be easily changed by adding reinforcements to the reaction system before the SHS reaction or by adding them to the powder before the SPS compaction.

2 EXPERIMENTAL WORK

The NiAl intermetallic compound was prepared with an SHS synthesis. A high-purity nickel powder with a particle size <100 μm and an aluminium powder with a purity of 99.99 % and a particle size of 200–400 μm were mixed and pressed at room temperature with a pressure of 260 MPa using a LabTest 5.250SP1-VM universal testing machine. Reactive sintering of the pressed powder mixtures was carried out at 900 °C for 15 min in the usual furnace (air) atmosphere. The sintered particles with an approximately cylindrical shape and a size of 1 cm in diameter and 1 cm in height were milled with a laboratory vibration mill VM4. The obtained NiAl powder was leached in a 20 % NaOH solution to dilute any residual Al. The NiAl powder was compacted with the SPS procedure (model SPS 10-4 thermal technology) at a temperature of 1100 °C, for a compaction-process time of 5 min and at a pressure of 80 MPa. The SPS die is made of carbon and its internal diameter is 19.3 mm. To separate the sintered material from the die, a carbon foil with a thickness of 0.15 mm was used. The amount of compacted material was approximately 5 g for each experiment.

The structures of the SPS materials, the NiAl powder and the SPS-compacted material were observed with an Olympus PME3 light microscope and a TESCAN VEGA 3 LMU scanning electron microscope equipped with EDS and EBSD detectors (Oxford Instruments). The phase compositions of the materials were determined using X-ray diffraction (PAN analytical X’Pert PRO + High Score Plus, Cu anode). TEM samples were prepared by ion polishing using Gatan PIPS Model 691 and consequently observed with a Jeol JEM 3010 transmission electron microscope. SAED patterns were integrated and phases were identified using Process Diffraction software. The hardness of the materials was measured with a FUTURE-TECH FM700 hardness tester with loads of 10 g and 1 kg.

3 RESULTS AND DISCUSSION

The samples prepared with the SHS procedure had approximately cylindrical shapes. They were mainly composed of the NiAl phase with a low amount of residual Al in the surroundings of the pores. As illustrated in Figure 1, the porosity of the SHS samples is extremely high.

The NiAl particles were milled into a powder, whose structure is shown in Figure 2. The particles have irregular shapes, as expected after milling a brittle material.
The size of the majority (96 %) of the particles is less than 140 μm. The phase composition of the powder is given in Figure 3. Peaks of residual aluminium are also visible in the milled powder. Although the powder was leached with a 20 % NaOH solution, areas of residual Al are still shown in Figure 1.

Subsequently, the powder was compacted with the SPS method at 1100 °C for 5 min. The structure of the SPS-prepared material is given in Figure 4. The particles of the initial powder are clearly distinguishable. The dark parts in the structure are pores. The porosity of the SPS-prepared material is 1.9 ± 0.9 %, which is satisfactory for the material prepared by powder-metallurgy processing.

The grains of the compacted material are formed by the particles of the initial powder and no grain coarsening is observed. It can be supposed that the grain size of the compacted material depends only on the particle size of the initial powder. The plot in Figure 5 shows the particle-size distribution of the initial NiAl powder and the grain-size distribution of the SPS-compacted material. It seems that the powder contains more particles with a size of up to 20 μm. This slight disagreement can be caused by a measurement error. Small particles located at the grain boundaries cannot be distinguished as easily as the separate particles in the mounting material.

The amount of residual Al is lower than a 2–5 % of mass fraction because it is not detectable with XRD, as shown in Figure 3. The same is true of the oxide content in the initial powder and also in the SPS-compacted

![Figure 3: XRD pattern of NiAl powder and compacted material](image)

Slika 3: Rentgenogram prahu NiAl in kompaktiranega materiala (1 = NiAl, 2 = grafit)

![Figure 4: Structure of NiAl material compacted from powder with SPS method (1100 °C/5 min) (LM)](image)

Slika 4: Struktura NiAl materiala, kompaktiranega iz prahu po SPS metodi (1100 °C/5 min) (LM)

![Figure 5: Grain (particle) size distribution of NiAl powder and compacted material](image)

Slika 5: Razporeditev velikosti delcev prahu NiAl in kompaktiranega materiala

![Figure 6: SEM micrograph of NiAl material compacted from powder with SPS method (1100 °C/5min) and EBSD scan of the area](image)

Slika 6: SEM-posnetek NiAl materiala, kompaktiranega iz prahu po SPS metodi (1100 °C/5 min) in EBSD posnetek področja

The EBSD analysis (Figure 6) of the SPS-compacted material proved that the area of the initial-powder particles is monocrystalline. Between the large, clearly seen particles (grains of the compacted material), there are areas where the crystallographic orientation is not very clear. These areas at the grain boundaries can exhibit a large misorientation or can be oxidised.

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![Figure 7: TEM micrograph of NiAl material compacted from powder with SPS method (1100 °C/5min)](image)

Slika 7: TEM-posnetek NiAl kompaktiranega materiala iz prahu po metodi SPS (1100 °C/5 min)
The interlayer during compaction. The question is what change significantly due to the formation of an oxide same after consolidation, while macro-properties these results indicate that micro-properties stay the with a load of 1 kg) varies significantly (Figure 9).

Figure 9: Microhardness and macrohardness of samples after SHS preparation and powder-metallurgy preparation with SPS product. The only excess peak in the XRD pattern of the SPS-compacted material relates to the graphite from the protection graphite foil used in the SPS process.

A detailed material observation made using TEM is given in Figure 7 and it shows the structure of a grain boundary. In the left bottom part, a dark NiAl particle is located. It can be seen that the particles are connected by a nanocrystalline oxide interlayer. The amount, thickness and crystallinity of the oxide layer are not sufficient to be detected with XRD or EBSD analysis, but they can be distinguished with selected area electron diffraction (SAED), as shown in Figure 8.

The fact that the weak parts of the material are the grain boundaries is also proved with the hardness measurement. While the microhardness (inside individual particles) is the same for the SHS material and for the SPS-compacted material, the macrohardness (measured with a load of 1 kg) varies significantly (Figure 9). These results indicate that micro-properties stay the same after a consolidation, while macro-properties change significantly due to the formation of an oxide interlayer during a compaction. The question is what would happen if the SHS process was performed in an inert atmosphere.

4 CONCLUSION

The powder-metallurgy preparation of NiAl consisting of the SHS NiAl preparation, the milling and the SPS compaction is a promising method for obtaining bulk intermetallic materials. The grain size of an SPS-compacted material is mainly determined by the grain-size distribution of the initial powder. The grain size was estimated to be less than 40 μm. It was proved that the particles of the initial powder are interconnected by a thin oxide layer, which decreases the macroscopic and also microscopic properties of the material.

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5 REFERENCES