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Self-propagating high-temperature synthesis of nial intermetallic compounds

Abstract. In this work, the self-propagating high-temperature synthesis (SHS) method was utilized for the synthesis of the B2 NiAl compound. Elemental nickel and aluminum powders were mixed, die-pressed, and pre-heated in a furnace in which the SHS reaction was initiated and propagated. The influence of pre-heating temperature was evaluated against the final composition, phase formation, microstructure and hardness of the final compact. Conditions were found under which, the final product forms through solidification from the melt and (irrespective of the furnace temperature) consists only of the B2 NiAl. The NiAl compacts formed consisted of a core which is essentially pore free and accounts for ~80% of the total volume and a high porosity shell around it. The microstructure consists of large, equiaxial grains with linear grain boundaries indicative of high phase stability. Finally, it was found that as the pre-heating temperature increases the hardness of the NiAl compact decreases; a transient temperature analysis enabled to interpret this observation.

Practical significance of the results: NiAl is an extremely useful industrial material with many high temperature applications.

Key words: self-propagating high-temperature synthesis, NiAl

Introduction

The B2 ordered intermetallic NiAl compounds exhibit a wide range of physical and mechanical properties which show promise for applications ranging from high-pressure turbine blades to buried interconnects in electronic components [1]. NiAl forms an adherent film on oxidation in air at temperatures up to 1200°C, indicating good corrosion resistance as demonstrated by its use as a base for some protective coatings on nickel-based superalloys. The structure remains fully ordered up to the melting point, implying a resistance to diffusion-controlled processes, such as precipitation, grain growth, oxidation and creep deformation. The large phase width of NiAl suggests a potential for substitutional alloying to improve strength and oxidation resistance. NiAl is approximately 30% less dense than nickel (a common superalloy base) and has a melting point 200°C higher. On the other hand, drawbacks of this type of intermetallics include low ductility and toughness [1, 2].

Conventionally intermetallics are fabricated by processes such as vacuum arc melting and their final microstructure is refined by several thermomechanical treatments. To reduce costs and complexity, Self-Propagating High-Temperature Synthesis (SHS) and thermal explosion, has also been used [3-7]. In the SHS method, the exothermic reaction between the powder constituents is initiated and becomes self-sustaining to yield the final product progressively without requiring additional heat. Compared to conventional processing methods, the main advantages of SHS are: (i) higher purity products because of the high reaction temperature which can volatilize and hence “remove” low boiling point impurities, (ii) simplicity since there is no need for expensive processing facilities and equipment, (iii) low operating and processing costs due to the short exothermic reaction times, (iv) ability to fabricate non-equilibrium or metastable phases because of the high thermal gradients and rapid cooling rates and (v) capability for producing a wide range of inorganic materials consolidated into a final product in one step by utilizing the chemical energy of the reactants [8-10]. However, there are some disadvantages of the method such as high porosity (which may be reduced by simultane-
ous compression [11]), microstructural inhomogenei-
ties, etc. Several reviews of the combustion synthe-
sis technique and its prospects have appeared in the
literature [12-22].

**Thermodynamics**

During the combustion synthesis reaction there are four important temperatures which af-
fect the process of the reaction and the properties
of the final product: 1) the initial or preheating
temperature \( T_o \), which is the temperature of the
powder compact of the reactant sample before the
reaction is initiated, 2) the ignition temperature
\( T_{ig} \), which represents the point at which the SHS
reaction is spontaneously activated without fur-
ther external heat supply, 3) the adiabatic combus-
tion temperature \( T_{ad} \), which is the maximum
combustion temperature achieved under adiabatic
conditions and (4) the actual combustion temper-
ature \( T_c \), achieved normally under non-adiabatic
conditions [23].

The amount of heat, \( H(R) \) required to raise the
temperature of the reactants from the initial tempera-
ture \( T_o \) to the ignition temperature \( T_{ig} \) (i.e. the tem-
perature at which the exothermic reaction initiates) is
given by the following equation:

\[
H(R) = \int_{T_o}^{T_{ig}} \sum n_i C_p(R_i) \, dT + \sum n_i L(R_i) \quad (1)
\]

where \( n_i, C_p(R_i), \) and \( L(R_i) \) represent the reaction sto-
ichiometric coefficients, heat capacities and the phase
transformation enthalpies (if the reactants go through a
phase change such as from solid to liquid) of the
reactants respectively. At a certain distance from
the heat source, the reactant compact reaches steady
state conditions; in this state the heat of the reaction
\( \Delta H(T_{ig}) \) is only used to heat the adjacent layer from \( T_o \)
to \( T_{ig} \). At this point, the amount of heat available to be
absorbed by the products under adiabatic conditions is,
therefore, \( H(P) \), which raises the temperature from
\( T_{ig} \) to the adiabatic temperature \( T_{ad}(T_o) \), i.e.

\[
\Delta H(T_{ig}) = - [H(R) + H(P)] \quad (2)
\]

where \( H(P) \) is given by:

\[
H(P) = \int_{T_{ig}}^{T_{ad}(T_o)} \sum n_j C_p(P_j) \, dT + \sum n_i L(P_j) \quad (3)
\]

where \( n_j, C_p(P_j), \) and \( L(P_j) \) represent the reaction sto-
ichiometric coefficients, heat capacities and the phase
transformation enthalpies (if the products go through a
transformation phase change) of the products
respectively. The increase of the initial (average)
temperature \( T_o \) will decrease \( H(R) \) and increase both \( H(P) \) and the adiabatic temperature.
Increasing \( T_o \) to \( T_{ig} \) will decrease \( H(R) \) to zero and
all of \( \Delta H(T_{ig}) \) will be available to be absorbed by the
products [14,23].

**Materials and Methods**

**Processing and synthesis**

Used in the experiments Ni powder characteristics
presented in table 1.

### Table 1 – Nickel Powder Characteristics

| Composition |  C   | Mn   | Cu   | Fe  | S    | Si   | Ni   |
|-------------|------|------|------|-----|------|------|------|
| Percent     | 0.017| <0.01| <0.01| 0.04| 0.0002| 0.31 | Bal. |

**Size Distribution**

| Mesh Size | +100 | +120 | +140 | +200 | +270 | +325 |
| Percent   | 4.2  | 3.0  | 6.1  | 15.2 | 15.9 | 13.7 |

Aluminum powder MERCK, max. particle size 125 micron, purity 99.5%.
Stoichiometric fractions of elemental powders to produce the NiAl phase (38.5wt%Ni + 61.5wt%Al) were mixed thoroughly and die-pressed at a pressure of 30 bars to form cylindrical compacts (diameter c. 15 mm and height c. 20 mm) of green density about 60% of theoretical. The cylindrical specimens were then pre-heated in a furnace where the exothermic reaction between the Ni and the Al particles initiates as soon as any area of the compact reaches the ignition temperature. In all of our experiments, the reaction was initiated at the top surface of the specimens after exposure to the furnace temperature for a period between 1 and 4 minutes, depending on pre-heating temperature. As the SHS combustion zone moves through the specimen, a series of reactions take place and the compact melts, since the combustion temperature (approaching 2000 °C is well above the melting temperature of the NiAl phase (1640 °C). Once the SHS reaction is completed, the specimen is removed from the furnace and air-cooled.

The microstructure of the NiAl product, its chemical homogeneity and its mechanical properties usually depend on the processing conditions as well as the compact geometry, because of their effect on the actual average compact temperature at the time of ignition and on the cooling rate of the NiAl melt. The dimensions and the initial density of the specimens were kept fixed, while the furnace temperature $T_{fur}$ was varied from 700 to 1000°C. The time to reaction initiation $t_{in}$ was not constant in the experiments but decreased with increasing the furnace temperature as the initiation temperature was reached quicker.

**Characterization**

The NiAl materials synthesized were characterized by a number of techniques including X-ray diffraction, optical and scanning electron microscopy, electron dispersive spectroscopy, and hardness testing.

**Results and Discussion**

*Modelling of the compact temperature*

The time to ignition $t_{in}$, defined as the period that the specimen remains inside the furnace until the reaction initiates depends upon the initial temperature of the furnace. The variation of $t_{in}$ versus the furnace temperature $T_{fur}$ as determined from the SHS experiments is shown in Figure 1.

![Figure 1](image)

*Figure 1 – The variation of initiation time $t_{in}$ as a function of the furnace temperature $T_{fur}$*

It is seen that as the furnace temperature increases the initiation time decreases because the ignition temperature (on the top surface of the sample) is reached faster. After ignition, the time required to propagate the reaction front along the specimen is less than 1 second, i.e. the reaction speed is quite high\(^1\). Thus,\(^1\) in fact, earlier work\(^1\) on the synthesis of the 50% Ni – 50% Al system it was measured as being equal to XXX m/sec.
with regard to the microstructure formation and evolution, it we accept that at the end of the reaction and prior to solidification, the material is at the liquid state. The microstructure obtained from the solidification process is dependent upon the difference between the temperature of the melt (which is related to the combustion temperature of the reaction) and the furnace temperature. Therefore, to interpret the microstructural features the combustion temperature (or likewise the average compact temperature) must be estimated.

As mentioned above in the section of “Thermodynamics”, the maximum adiabatic temperature depends upon the average temperature of the compact. Evidently, the latter should be related to the furnace temperature and the time of exposure inside the furnace. In order to gain a comparative estimate of the average compact temperature right before the reaction initiation for the various \( T_{\text{fur}} \) and \( t_{\text{in}} \), a transient temperature analysis was conducted. In particular, the temperature at the center of the compact \( T_{\text{cen}} \) for the specific \( T_{\text{fur}} \) and \( t_{\text{in}} \) was calculated following the procedure which is outlined in the Appendix.

The variation of the temperature at the center of the compact versus the furnace temperature is shown in Figure 2.

![Figure 2](image_url) – The temperature at the center of the compact versus the furnace temperature

It is seen that, within the limits of the estimation error (which mainly arises from factors such as the accuracy of the calculation of the thermal properties of the porous compact) \( T_{\text{cen}} \) does not appear to vary with the furnace temperature. In other words \( T_{\text{cen}} \) and \( t_{\text{in}} \) are combined such that the temperature of the specimen’s center has the same temperature irrespectively of the processing conditions. Therefore, since the temperature at the center of the specimen reflects its average temperature, it is deduced that the combustion temperature was the same for all of the experimental conditions. Given the fact that the ignition temperature (which in the experiments of this work was first reached on the top surface of the specimen) is dependent only upon the composition of the compact, the above analysis is important as it provides an indication that the average temperature of the specimen right before the ignition does not vary with the furnace temperature.

**Characterization**

*a. X-ray Diffraction (XRD)*

Figure 3 shows the XRD spectrum of a material fabricated by self-propagating synthesis of nickel and aluminum powder under the following processing conditions: furnace temperature 1000 °C, green compact density 60% of the theoretical, and initial dimensions 20 mm diameter and 15 mm height.

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In order to eliminate possible effects of the texture developed during the solidification of the melt, the specimen was pulverized, and its powder was used for the XRD characterization. All peaks that appear in the graph correspond to the NiAl intermetallic phase; no other intermediate phases between nickel and aluminum, or any oxides were detected, and no further elemental aluminum and nickel are present. In addition, materials synthesized under different processing conditions (i.e. furnace temperature, time exposed inside the furnace, etc.) were also examined by XRD, giving identical spectra to that shown in Figure 3. Therefore, in all cases pure monophase NiAl with the B2 ordered cubic structure with a lattice parameter $\alpha = 2.88\text{Å}$ was fabricated.

Earlier characterization work on the formation of NiAl compounds using differential thermal analysis (DTA) and synchrotron radiation techniques revealed that during the SHS synthesis several intermediate phases such as Ni$_3$Al and Ni$_3$Al$_2$ are formed prior to the formation of the final, stoichiometric NiAl phase[23-24]. The formation as well as the presence of residues of the intermediate phases depends upon the combustion temperature. In particular, the proportion of the Ni$_3$Al and Ni$_3$Al$_2$ phases decreases as the combustion temperature increases. Therefore, the above results indicate that a high combustion temperature was achieved in our experiments.

Microstructure

As mentioned above, the NiAl formation proceeds through the formation of a melt. Examination of the microstructure at lower magnifications showed that the final product comprises two regions, a core which accounts for $\sim80\%$ of the total volume of the compact, and a “shell” of high porosity (of the order of 20%) material around the core. This outer shell, whose microstructure is shown in Figure 4, owes its formation to the gravitational flow of a small portion of the liquid, since in our experiments the specimen stood free inside the furnace, thus flow of the melt was possible.

It is observed that the structure comprises large ($\sim150$ µm) grains of equiaxed shape. The fact that the NiAl grains have linear grain boundaries with no curvature indicate the high stability of the formed compound, the absence of secondary phases, and a very rapid crystallization of the primary melt. Although, no pores can be discerned from this micrograph; the higher magnification micrograph in Figure 6 shows that there are a few spherical with a size of less than 10 µm inside the material.

Although at low magnifications the microstructure of the core appears to be quite uniform, microstructural nonuniformities may be distinguished at higher magnifications. In particular, the SEM micrograph of Figure 6 reveals the presence of spherical particles within a fine grained matrix.
A typical, low magnification, optical micrograph of the core is shown in Figure 5.

In conjunction with the XRD results it can be deduced that both the particles as well as the fine matrix are of pure NiAl. It is also seen that the pores are mainly located at the particle/matrix boundaries, which indicated that the large particles solidified first inside the matrix which was in the liquid state.

**Hardness**

The Vickers hardness of the NiAl compounds versus the furnace temperature at which they were synthesized is presented in Figure 8.
The particles have a wide distribution of sizes ranging between 4 and 15 µm. The chemical composition of the particles as well as the composition of the matrix were determined by EDS analysis. The EDS spectra shown in Figure 7 do not reveal any essential differences on the peaks of nickel and aluminum between the particles and the matrix.

The results revealed a trend of decreasing hardness with increasing the furnace temperature. This hardness variation can be interpreted with regard to the cooling rate of the NiAl melt which is proportional to the difference ΔT between the temperature of the melt $T_m$ and the temperature of the furnace $T_{fur}$. If it is assumed that the temperature of the melt is near the combustion temperature, (which as discussed in the previous sections is essentially constant for all conditions used) then ΔT increases as the furnace temperature decreases. Therefore, at lower $T_{fur}$ the cooling rate is higher, resulting to a more brittle and harder microstructure. SEM micrographs of the fracture surface of NiAl compacts fabricated at furnace temperatures of 700 and 1000 °C are shown in Figures 9a and b respectively. The micrograph of Figure 9a shows a brittle type of fracture, while areas which have undergone plastic deformation are observed in Figure 9b. Interestingly, NiAl fibres were also found to have formed inside pores and are also visible in Figure 9a and b. This is the first time such fibres have been observed and the exact conditions of formation and characteristics are under investigation at present.

**Figure 6** – Microstructure of semi-spherical particles within a fine grained matrix

**Figure 7** – The EDS spectra of the particles (left) and the matrix (right) showing that they have almost identical elemental composition
In summary, the low sensitivity of the material hardness (and hence strength) upon the furnace temperature represents a significant technological advantage of the process from the industrial point of view, because it does not pose great difficulties in scale-up, and also the material quality does not vary greatly with the processing furnace temperature.
Summary and conclusions

The SHS method was used to synthesize the B2 NiAl compound from elemental nickel and aluminum powders. The main conclusions of this work may be summarized as follows:

1. X-ray diffraction showed that the NiAl is the only phase formed (through solidification of a melt) when the powder compacts are processed at a furnace temperature is at the 700-1000 °C range.

2. The microstructure consists of large equiaxed grains with linear grain boundaries. The latter indicate the high stability of the compound synthesized and a very rapid crystallization of the primary melt.

3. The hardness of the final product was found to decrease with the increase of the furnace temperature because of the lower cooling rate of the melt.

4. The process used has shown that NiAl fibres can form inside pores of the material.

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