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Elastic properties of the α’ martensitic phase in the Ti-6Al-4V alloy obtained by additive manufacturing

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A R T I C L E   I N F O

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A B S T R A C T

Elastic behaviour of Ti 6Al 4V alloy elaborated through additive manufacturing process is studied both experimentally and through atomistic simulation. By studying rough samples after fabrication and after heat treatment, the elastic properties of the α’ martensitic phase is compared to that of α + β microstructure. Atomistic calculations were also performed on various super cells varying their chemical composition in order to simulate α and α’ phases and their full elastic stiffness tensors were determined. Both experimental and simulation results show that Young modulus of the α’ phase is lower than that of α phase, while it presents a more anisotropic behaviour.

Additive manufacturing (AM) is widely used to elaborate industrial parts with an increasing variety of materials. In the present study, we focus on the titanium alloy Ti 6Al 4V elaborated by powder bed laser beam melting (PB LBM). Ti 6Al 4V is a widely used Ti alloy known as an (α + β) alloy [1]. Indeed, wrought Ti 6Al 4V parts present a two phase microstructure with roughly 94% vol. for the hexagonal compact (hcp) α phase and 6% vol. for the body centred cubic (bcc) β phase [2]. When processed by AM, metallic alloys experiment a very high cooling rate that may result in out of equilibrium structure. For instance during PB LBM processing of Ti 6Al 4V the cooling rate is in the range of $10^7$ to $10^8$ K·s⁻¹ [3], resulting in a single phase α’ martensitic structure [4,5]. This α’ phase presents, similarly to the pure Ti α phase, a hcp structure but it incorporates the alloying elements of the Ti 6Al 4V alloy and forms a very fine entangled needles microstructure [4,5]. Recent papers review the mechanical properties of the Ti 6Al 4V processed by AM [6,7]. These studies often focus on plastic properties and the low ductility of AM Ti 6Al 4V. In many instances the elastic behaviour of the martensitic α’ is poorly documented. Few values of the Young modulus deduced from tensile testing are reported with quite a large scattering, the modulus varying from 94 to 118 GPa [6].

The purpose of the present study is to determine the elastic behaviour of the α’ martensitic phase and compare it to that of the Ti α phase. This is conducted in two complementary ways: i) by measuring experimentally the overall Young modulus of the α’ phase on PB LBM samples and ii) by calculating the elastic stiffness constants $C_{ij}$ using atomistic simulations. The obtained results are discussed in regard to the literature data and an emphasis is placed on the evolution of elastic behaviour of the hexagonal phase when changing from pure Ti (α phase) to the martensitic phase α’.

In order to experimentally measure the Young modulus of the α’ phase, samples were produced by PB LBM in an M2 Cusing machine from Concept Laser, using Ti 6Al 4V powder with an average powder grain diameter of 35 μm. For tensile testing dog bone plate shaped samples (2 mm thick, gage length with 2 mm width and 10 mm length) were manufactured. The surfaces of tensile specimens were ground using grade 600 SiC paper. For the resonance vibration measurements the samples are beam shaped with dimension $80 \times 20 \times 4$ mm³. Both types of sample were fabricated vertically, that is with the long axis of the specimen along the elaboration direction.

In most cases titanium pieces elaborated by PB LBM are given an heat treatment (HT) in order to release residual stress and stabilise the microstructure by decomposing the α’ martensite to the (α + β) microstructure [4]. This HT was applied using a Carbolute CFW 1300 under laboratory air, with a heating ramp 5 °C/min from room temperate up to 700 °C, holding 1 h at 700 °C and furnace cooling. It was verified by scanning electron microscopy observations that the microstructure of the raw AM samples consists of a unique hcp phase in the form of entangled fine needles, corresponding to the martensitic α’ phase. The phase proportions were investigated by X ray diffraction using a D8 2 diffractometer from Bruker with a Cu source and a 1D
Fig. 1. X ray diffraction pattern on Ti-6Al-4V samples elaborated by AM a) in the raw fabrication state and b) after HT. The peak indexing of hexagonal ($\alpha'$ and $\alpha$) and cubic ($\beta$) phases are showed.
LYNXEYE detector. Scans were performed from 20° to 80° by 0.02° steps every 3 s. The obtained diffractograms were analysed according to Rietveld technique using TOPAS 4.2 [8]. As can be seen in the Fig. 1(a) corresponding to raw AM sample, all the diffraction peaks observed are associated to a unique hcp phase with cell parameters: \(a = 292.9\) pm and \(c = 465.5\) pm. By contrast the diffraction pattern presented in Fig. 1(b) exhibits peaks from both the hexagonal \(\alpha\) phase and the cubic \(\beta\) phase. This later accounts for 6% ±1% volume fraction. As a whole, we consider that the raw sample after AM is fully constituted by the \(\alpha\)' martensitic phase, while after HT an \((\alpha + \beta)\) microstructure close to the equilibrium of the Ti 6Al 4V is present.

The tensile tests were performed at room temperature using a MTS Insight electrically driven tensile machine. The cross head motion rate was fixed to \(10^{-2}\) mm·s\(^{-1}\) and the elongation was measured using a MTS clip on extensometer with initial spacing of 10 mm. Young modulus was also determined using impulse excitation technique, consisting of the relation:

\[
E_{\text{modulus}}(\text{GPa}) = 99.8 - 0.0427 \times T(\degree\text{C}).
\]

A second measurement performed on sample D, consisted in heating the sample up to 700 °C, holding for 1 h and furnace cooling. This thermal cycle actually reproduces the HT performed on tensile sample as previously described. The evolution of Young modulus during this
cycle is plotted in Fig. 3(b). At the beginning of heating ramp, the Young modulus value (initial value 98.7 GPa) and its temperature dependence is very similar to that of sample C. For temperature over 500 °C, the variation of Young modulus deviates from linear temperature dependence. Upon cooling, some irreversible changes of the value of Young modulus and its slope versus temperature are observed. They are associated to the phase transition from \( \alpha \) to \( \beta \). The cooling branch of the curve below 500 °C indicates a linear variation of \( E(\alpha + \beta) \) microstructure for the \( \alpha \) phase may present a Young modulus significantly lower than that of \( \alpha + \beta \) phase.

At 500 °C, the inflection of the curve is very similar to that of sample C. For temperature over 500 °C, the variation of Young modulus values for samples before and after HT is measured to be 103.4 GPa. Both this room temperature value and the temperature variation of Young modulus given by Eq. (2) are in good agreement with the values reported in the literature for elastic behaviour of wrought Ti 6Al 4V alloy [2,14].

A summary of the experimental results obtained in this part is given in Table 1 in which Young modulus values for samples before and after HT are compiled. Despite the absolute values obtained and their scattering depend on the experimental technique, both tensile testing and vibration technique clearly indicate that the Young modulus of the \( \alpha' \) martensite is lower than that of \( \alpha + \beta \) microstructure for the Ti 6Al 4V alloy. Moreover it is recognized that vibrational method are more accurate than tensile tests for measuring Young modulus of metallic materials, see e.g. [15]. This method allowed to measure the temperature variation of the \( \alpha' \) martensite Young modulus which has never been reported in the literature to our knowledge.

The Young modulus of the \( \alpha \) phase is even higher than that of \( \alpha + \beta \) microstructure, due to the fact that the \( \beta \) phase has a low Young modulus [2]. Consistently the Young modulus of pure titanium \( \alpha \) phase was reported to be 112 GPa [2,16]. Which means that the martensitic \( \alpha' \) phase may present a Young modulus significantly lower than that of the \( \alpha \) phase.

Atomistic calculations were performed with the density functional theory (DFT) using Vienna ab initio Simulation Package [17]. Self consistent Kohn Sham equations were solved using the projector augmented wave pseudo potentials [18]. We used the Perdew Burke Ernzerhof [19] exchange and correlation functional. The plane wave energy cut off was set to 500 eV. Equivalent \( \Gamma \) centred 4x4x4 Monkhorst Pack meshes [20] were used to sample the first Brillouin zone of the primitive cell (by using a band folding approach with two atoms per unit cell).

### Table 2

List of the five super-cells used for DFT simulations. The number of atoms is indicated together with the at.% of Al and V (in brackets). The cells Ti and Ti + 2Al + V represent the phases \( \alpha \) and \( \alpha' \) respectively.

<table>
<thead>
<tr>
<th>Phase</th>
<th>Ti</th>
<th>Al</th>
<th>V</th>
<th>Ti + Al</th>
<th>23</th>
<th>1 (4.2%)</th>
<th>0</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \alpha )</td>
<td>24</td>
<td>0</td>
<td>0</td>
<td>( \alpha' )</td>
<td>21</td>
<td>2 (8.4%)</td>
<td>1 (4.2%)</td>
</tr>
</tbody>
</table>

### Table 3

Result of DFT calculations for the five super-cells considered. The stiffness tensor components are given in GPa and cell parameters are given in pm.

<table>
<thead>
<tr>
<th></th>
<th>( C_{11} )</th>
<th>( C_{12} )</th>
<th>( C_{13} )</th>
<th>( C_{22} )</th>
<th>( C_{33} )</th>
<th>( C_{44} )</th>
<th>( a )</th>
<th>( c )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ti</td>
<td>178.4</td>
<td>80.9</td>
<td>76.9</td>
<td>189.2</td>
<td>43.4</td>
<td>48.8</td>
<td>293.7</td>
<td>464.8</td>
</tr>
<tr>
<td>Ti + Al</td>
<td>174.5</td>
<td>83.9</td>
<td>73.6</td>
<td>188.0</td>
<td>45.2</td>
<td>45.3</td>
<td>292.7</td>
<td>467.9</td>
</tr>
<tr>
<td>Ti + V</td>
<td>163.0</td>
<td>90.3</td>
<td>79.5</td>
<td>191.0</td>
<td>36.3</td>
<td>32.4</td>
<td>292.6</td>
<td>464.2</td>
</tr>
<tr>
<td>Ti + Al + V</td>
<td>162.9</td>
<td>100.9</td>
<td>77.4</td>
<td>193.2</td>
<td>40.8</td>
<td>31.0</td>
<td>291.8</td>
<td>463.8</td>
</tr>
<tr>
<td>Ti + 2Al + V</td>
<td>161.0</td>
<td>104.3</td>
<td>75.8</td>
<td>198.6</td>
<td>43.0</td>
<td>28.4</td>
<td>291.2</td>
<td>464.1</td>
</tr>
</tbody>
</table>

### Table 4

Stiffness tensor, in GPa, from the literature for pure Ti and substituted (3 at.%) cells.

<table>
<thead>
<tr>
<th></th>
<th>( C_{11} )</th>
<th>( C_{12} )</th>
<th>( C_{13} )</th>
<th>( C_{22} )</th>
<th>( C_{33} )</th>
<th>( C_{44} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \alpha )</td>
<td>160.0</td>
<td>90.0</td>
<td>72.3</td>
<td>183.0</td>
<td>45.0</td>
<td>35.0</td>
</tr>
<tr>
<td>( \beta )</td>
<td>175.0</td>
<td>89.0</td>
<td>84.0</td>
<td>192.0</td>
<td>45.0</td>
<td>43.0</td>
</tr>
<tr>
<td>Ti + Al</td>
<td>171.9</td>
<td>94.1</td>
<td>84.1</td>
<td>190.6</td>
<td>39.2</td>
<td>38.9</td>
</tr>
<tr>
<td>Ti + V</td>
<td>164.8</td>
<td>99.8</td>
<td>86.4</td>
<td>191.0</td>
<td>30.2</td>
<td>32.5</td>
</tr>
</tbody>
</table>

To perform elastic calculations, we choose five different types of strain to compute the five independent elastic constants. The elastic energy \( U \) is approximated by a quadratic function of the strain components (see [21]) as described with more details in supplementary data file (available on line).

A super cell approach was used with full periodic boundary conditions to describe substituted systems. The super cell is based on the crystallographic structure of pure titanium \( \alpha \) phase, space group \( P6_3/mmc \) where titanium (Ti) atoms occupy the positions \( 2c \). In order to simulate alloys, vanadium (V) and/or aluminium (Al) atoms have been substituted to Ti atoms in the super cell. For the super cells containing several added elements in substitution at the same time, we considered different cases where the substituted atoms were either in first (1NN) or in second nearest neighbouring positions (2NN). We found that Al V interaction is attractive at short range (binding energy is equal to about 20 meV), alloying atoms were thus placed in 1NN positions in our simulations. Five different super cells were used, as shown in Table 2, with an increasing number of substitution atoms, in order to study the evolution of properties from the pure Ti super cell to the Ti + 2Al + 1 V super cell. This latter represents the martensitic \( \alpha' \) phase, the current chemical composition being 4.9 wt% Al and 4.6 wt% V quite close to the Ti 6Al 4V alloy composition.

From these simulation results shown in Table 3, one can see that varying the chemical content of the super cell has a main effect on the values of \( C_{11} \) and \( C_{33} \). Adding alloying elements V and Al lowers the value of \( C_{11} \) and rises \( C_{33} \). For the pure Ti \( \alpha \) phase, the \( C_{ij} \) values obtained, Table 3, are well in line with the \( C_{ij} \) experimental values reported in the literature [16] and presented in Table 4. DFT calculations of the \( C_{ij} \) were previously performed by Wilson et al. [22] for pure Ti and with the incorporation of various substitutional elements. These authors also showed that different alloying elements have different effects on the various components of the stiffness tensor. However, they limited their calculations to a lower amount (3 at.%) of substitutional atoms and did not study the combination effect observed when alloying
with both V and Al in order to get closer to the composition of the α′ martensite phase.

The increasing difference between components $C_{11}$ and $C_{12}$ that is observed in Table 3 with the incorporation of alloying elements Al and V, reveals an increased anisotropic character of the elastic behaviour of the hcp phase when changing from α to α′ phase. This is better represented by calculating the Young modulus along a direction with increasing inclination with respect to the c axis, represented by the angle $\psi$. Young modulus versus the $\psi$ angle are displayed in Fig. 4 where the experimental values come from Ogi et al. [16] and simulation data from this work. It appears that Young modulus is maximum along the c axis, minimum in the basal plane (Young modulus is isotropic in basal plane for the hexagonal system) for Ti + 2Al + 1 V simulation ($\alpha′$ phase) and for the experimental values of the $\alpha$ phase. The phase with the higher anisotropy is the $\alpha′$ martensite with values of $E (\psi = 0°) = 155$ GPa and $E (\psi = 90°) = 89$ GPa.

Finally, it is interesting to compare the average value of the Young modulus for hcp phases $\alpha$ and $\alpha′$. Some values presented in Table 5 are calculated from $C_{ij}$ through Hill averaging between Voigt and Reuss modulus [23], $C_{ij}$ values from simulation in the present study are used for both $\alpha$ and $\alpha′$ phases, while experimental values of $C_{ij}$ are taken from [16] for Ti $\alpha$ phase. The experimental value for the $\alpha′$ phase is taken from vibrational technique presented in this paper (Table 1). One can see comparing the values in Table 5 that for both phases experimental values are lower than calculated ones, however both experiment and simulation indicate similar variation from $\alpha′$ phase to $\alpha$ phase. It is also confirmed that the Young modulus of the martensitic phase $\alpha′$ is lower than that of the pure Ti $\alpha$ phase.

To summarise, the elasticity of the $\alpha′$ martensitic phase of the Ti 6Al 4V alloy obtained by PB LBMA has been investigated using two experimental methods and DFT simulations. For the first time, the temperature dependence of the Young modulus was determined using vibrational techniques, before and after HT, corresponding to the $\alpha′$ and the ($\alpha + \beta$) microstructures. It is shown that the Young modulus of the martensitic phase is lower than that of the pure Ti $\alpha$ phase. DFT calculations showed the influence of Al and V substitutional elements on the elastic constants of the hexagonal phase. Stiffness tensors of super cells representing $\alpha$ and $\alpha′$ phases were calculated. The values confirmed the lower value of Young modulus for the martensitic $\alpha′$ phase and also showed that its elastic behaviour is more anisotropic than that of the $\alpha$ phase.

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References