Plasma Enhanced Chemical Vapour Deposition (PECVD) at Atmospheric Pressure (AP) of Organosilicon Films for Adhesion Promotion on Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al and Ti$_6$Al$_4$V

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Abstract: The recent generation of aircraft is mainly built out of CRFP (Carbon fiber reinforced plastic). To ensure galvanic compatibility Titanium alloys are used at connecting points to metallic structures. The increased use of Titanium alloys due to their good mechanical properties has highlighted the need for structural bonding as joining method to combine Titanium with CFRP. However, long-term stable adhesion on Titanium is still an issue. Currently mainly wet-chemical processes are applied to improve adhesion on Titanium alloys. Since wet-chemical treatments often comprise the use of hazardous chemicals dry environmentally friendly surface functionalization processes are pushed more into focus. PECVD (Plasma enhanced chemical vapour deposition) at AP (Atmospheric pressure) seems to be favorable for the deposition of silicon dioxide films as an adhesion promoter layer. In the present work, the deposition of silicon dioxide adhesion layer (from HMDSO (Hexamethyldisiloxane) as precursor) on titanium alloys is described for a linearly extended and scalable DC arc plasma source (LARGE). The wedge test is used to evaluate the adhesion properties of coated titanium samples in comparison to the standard used NaTESi anodizing process. The results show that the atmospheric pressure plasma process leads to good long-term durability and to high bonding strength which is comparable to the used standard pre-treatments.

Key words: Atmospheric pressure PECVD (Plasma enhanced chemical vapour deposition), plasma surface treatment, silicon dioxide films, titanium alloys, adhesion layers.

1. Introduction

In the aerospace industry new design concepts offering weight savings are high in demand. In the recent generation of aircraft (e.g. Airbus 350 XWB, Boeing 787), composite materials have shown new perspectives in terms of structural efficiency and performance. In a fast-paced environment under threat from new market entrants and technology disruptions, competitiveness and operational excellence new technologies have to be developed which give our products additional values like new light weight designs. In an economically and technologically ever-accelerating environment the customers (e.g. airlines) expect that new technologies lead to less fuel burn without any impact on comfort. To keep this target new surface treatments like atmospheric pressure plasma are necessary to meet the requirements from the customers and to increase the efficiency in manufacturing.

It is well known that in the aerospace industry titanium materials exhibit critical issues in regard to long-term stable adhesion. Therefore composite titanium hybrid joints can only be realized by riveting. Adhesive bonding is seen to be the key to realize weight saving hybrid structures. However for this application surface treatments have to be developed to ensure long-term stable adhesion on titanium and composite materials.

For improving the adhesion on titanium various wet-chemical surface treatment methods which are
Plasma Enhanced Chemical Vapour Deposition (PECVD) at Atmospheric Pressure (AP) of Organosilicon Films for Adhesion Promotion on Ti₁₅V₃Cr₃Sn₃Al and Ti₆Al₄V partially based on environmentally hazardous compounds are used. PECVD (Plasma enhanced chemical vapor deposition) of organosilicon films at atmospheric pressure for adhesion promotion seems to be an alternative to these hazardous or manual methods.

During the last years, depositing of polymer-like organosilicon films or inorganic SiOₓ-like films by plasma polymerization technology have been utilized in a wide range of applications, such as protective coatings, adhesion promoters, biomedical materials, electronic and optical devices [1]. This is mainly owing to the fact that the functionality and important properties of plasma deposited layer can be determined by the nature of the used vapor precursors and deposition parameters [2, 3]. Among the wide variety of possible precursors for plasma-polymerization, the organosilicon monomer HMDSO (Hexamethyldisiloxane) is perhaps the most common and widely utilized product. HMDSO is non-toxic, non-flammable and easy to handle [4]. Moreover, HMDSO has high deposition rate and the ability to control the structure and properties of the polymerized films by varying deposition conditions [5].

In this paper, a linearly extended and scalable DC arc plasma source (LARGE) is used for the deposition of silica layers on Ti₁₅V₃Cr₃Sn₃Al and Ti₆Al₄V. The injection of the precursor is carried out by a remote gas feeding system made of BN (Boron Nitride). In the first section of the paper, the investigation of surface morphology and chemical composition after acidic etching are described. In this stage the characterization of deposited silicone dioxide film which is evaluated for HMDSO as precursor is executed. The second part covers the investigation of the long-term durability of bonded titanium samples in hot-wet condition.

Adhesive bonding tests have been performed in comparison to the wet chemical treatments like anodizing processes. The influence of the different plasma treatment parameters on adhesion properties was evaluated by the wedge test.

2. Materials and Methods

2.1 Materials

For this investigation the titanium alloys Ti₁₅V₃Cr₃Sn₃Al AMS4914 and Ti₆Al₄V AMS4911 are used. In Ti₁₅V₃Cr₃Sn₃Al alloy, the main elements are β - stabilizer such as vanadium and chromium. Tin is considered a neutral element since it does not strongly promote phase stability, it retards the rates of transformation and is useful as strengthening agent [6]. The Ti₆Al₄V alloy belongs to α + β - titanium alloys due to the main alloying elements the α-phase-stabilizer aluminum and the β-phase-stabilizer vanadium [7, 8].

For the wedge test investigations specimens with an edge length of 150 mm and a thickness of 2.0 mm are used. Analytical investigations are conducted on polished samples (Ra = 10 nm, measured by DektakTX Bruker, Germany) with the dimension of 25 × 25 mm².

2.2 Surface Treatment

All titanium samples are processed with the same steps for cleaning and texturing the surface prior to pre-treatment steps. The samples were cleaned with the alkaline cleaner using P3 almeco 18 (Henkel AG & Co. KGaA, Düsseldorf, Germany) at 65 ± 3 ºC for 15 min. Subsequently, the samples were alkaline etched with Turco 5578 (Henkel AG & Co. KGaA, Düsseldorf, Germany) at 95 ± 3 ºC for 5 min using a concentration of 500 g/L.

2.3 NaTESi Anodizing

The NaTESi pre-treatment of titanium for structural bonding were developed by Matz [9]. This process is used in this investigation as a reference to atmospheric pressure plasma deposition. The NaTESi anodisation was carried out using a three-electrode configuration with titanium sample as cathode. According to the investigation of Matz, the process is performed at 30 ±
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2 °C and 10 V for 15 min. The electrolyte contains 300 g/L sodium hydroxide, 65 g/L Na-tartrate, 30 g/L EDTA (Ethylendiamintetraacetic) and 6 g/L Natriumsulfit.

2.4 Plasma Treatment

For the nanometer-thin silicon dioxide film deposition the LARGE plasma source with linearly extended and stabilized DC arc plasma at atmospheric pressure is used. The principle of this PECVD process and the detailed description of the plasma source is given in previous publications [3, 10-12]. The silicon dioxide layer deposition presented in this paper is carried out with a 150 mm long arc and HMDSO as precursor. The precursor was heated up to 130 °C. For the precursor injection a remote gas feeding system made from BN is installed in front of the LARGE plasma source to avoid the contamination. The titanium samples were moved in front of the source at 20, 40 and 60 mm working distances. Depending on the cycles number the defined thickness of the plasma layer can be deposited on the surface. The process parameters are given in Table 1.

2.5 Surface Characterization

The chemical composition and the surface morphology of the initial oxide layer after acidic etching were investigated by means of SEM (Scanning electron microscope) using a Jeol JSM-6320F field emission scanning electron microscope (Jeol, Tokyo, Japan) in combination with energy dispersive x-ray analysis (EDX). Samples have been prepared by metallographic sectioning technique. The grinding step has been carried out with SiC (Silicon carbide) abrasive paper by decreasing the abrasive particle size from 80 to 4,000 grit. Subsequently, the samples were rough polished with 6-micron diamond followed by a 1-micron diamond on low napped polishing cloths. For the final polishing step the low napped polishing pads with colloidal silica were used. After polishing procedure the samples were chemical etched to optically enhance microstructural features such as grain size and phase features. The etching mixture consists of 960 mL H<sub>2</sub>O, 25 mL HNO<sub>3</sub> (65%) and 15 mL HF (40%). The exposure time to chemical etchant was 15 s. For investigation of the deposited thin silicone dioxide film cross sectioning is performed by FIB (Focused ion beam) thinning at the NMI (Naturwissenschaftliches und Medizinisches Institute) Tübingen. The schematically drawing of the FIB cross-section preparation is given in Fig. 1.

For FIP cross-sections, the sample pieces were glued on aluminum sample plate and electrically contacted with conductive silver. Before carrying out the FIB cross-sections a thin Pt protective strip has been deposited to the sample surface by means of electron beam (dark red). Then, in order to avoid etching of the surface during the ion beam deposition, thick Pt protective strip has been deposited by means of ion beam (bright red).

2.6 Test Methods

In order to evaluate the bonding strength the wedge test according to the German standard DIN 65448 was employed. The test combines mechanical and

<table>
<thead>
<tr>
<th>Process parameters</th>
<th>Value</th>
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<tr>
<td>Plasma gas</td>
<td>70 SLM/min Ar + 50 SLM/min N&lt;sub&gt;2&lt;/sub&gt; + 160 SLM/min compressed air</td>
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<tr>
<td>Remote gas</td>
<td>40 SLM/min N&lt;sub&gt;2&lt;/sub&gt; + 40 SLM/min O&lt;sub&gt;2&lt;/sub&gt;</td>
</tr>
<tr>
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<td>Working width</td>
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<tr>
<td>Current</td>
<td>75 A</td>
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<td>Substrate velocity</td>
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hydrothermal loads. After the pre-treatment two sheets are bonded with an aerospace certify epoxy-based adhesive FM 73 M (cured in an autoclave at 120 °C and a pressure of 0.25 MPa). The matrix of the FM 73 M is composed out of DGEBA (diglycidylether of bisphenol-A) and a hardening component 2, 4-tolyl-1, 1-dimethylurea. Stripes with the dimensions of 25 × 150 mm² are employed for testing. To start the crack propagation a stainless steel wedge is introduced into the bondline. Subsequently the samples are subjected to a hot-wet environment (50 °C, 95% rH) for 500 h. The growth of the crack, initiated by the wedge is microscopically monitored with times after 1.25, 24, 48, 96, 168, 336 and 500 h.

3. Results and Discussion

3.1 Investigation of Surface Morphology and Chemical Composition after Etching

The morphology and the chemical composition of the surface after alkaline etching step have an influence on the growth of the plasma layer during deposition. For the analytical investigations of the titanium alloys Ti15V3Cr3Sn3Al and Ti6Al4V metallographic sections were prepared. The surfaces were grinded and polished. Subsequently the metallographic sections were etched with HF/HNO3/H2O mixture to evaluate the alloy structure with their grain boundaries and precipitations. Fig. 2a shows the microsection of Ti6Al4V after etching. The titanium alloy Ti6Al4V exhibits a large amount of micro roughness. The typical α + β - microstructure with bright body-centered cubic β-phase precipitations at the closed-packed hexagonal α - phase grain boundaries is visible. Fig. 2a exhibits a α + β titanium alloy bimodal structure with equiaxed primary α grains (dark) and lamellar α + β colonies, and transformed β (bright spots).

In contrast to Fig. 2a, the cross section of the titanium alloy Ti15V3Cr3Sn3Al (Fig. 2b) shows coarsely crystalline structure of β-grains with two different orientations at the grain boundary. Fig. 3 a and 3b feature the SEM micrographs and the position of the EDX line scans for both titanium alloys. The SEM micrograph of Ti6Al4V exhibits β-segregations which are associated with α-segregations (Fig. 2a). In this area α-phase stabilized aluminum is depleted and β-phase stabilized vanadium is enriched. This phenomenon is well visible in the EDX spectrum. At the β-precipitations the vanadium content (yellow line) increases and the aluminum content (blue line) decreases. The EDX line scan of β-titanium alloy Ti15V3Cr3Sn3Al (Fig. 3 b) shows a homogeneous β structure without any significant changes in the concentration of the alloying elements.

3.2 Investigation of the Deposited Silicone Dioxide Layer

To determine the morphology and the chemical composition of the deposited silicon dioxide layer on both alloys, SEM and EDX were employed to characterize the coating. The cross sections were prepared by FIB technology. Both alloys were treated with the same parameters: 4 cm working distance and a 150 nm nominal silicone oxide layer thickness. As precursor Hexamethyldisiloxane (HMDSO) was used.

Fig. 4 a features the extraction area of the FIB lamella. The lamella was taken from a Ti6Al4V sample after alkaline etching and silicon dioxide film deposition. In Fig. 4 b, SEM micrograph of the FIB lamella was shown. It demonstrates that the deposited plasma layer was very porous with a film thickness of
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Fig. 2  SEM micrograph of a metallographic section after etching with HF/HNO<sub>3</sub>/H<sub>2</sub>O mixture: (a) Ti<sub>6</sub>Al<sub>4</sub>V and (b) Ti<sub>6</sub>V<sub>3</sub>Cr<sub>3</sub>Sn<sub>3</sub>Al.

Fig. 3  SEM micrograph of the etched (HF/HNO<sub>3</sub>/H<sub>2</sub>O mixture) titanium with indicated line scan: (a) Ti<sub>6</sub>Al<sub>4</sub>V, (b) Ti<sub>6</sub>V<sub>3</sub>Cr<sub>3</sub>Sn<sub>3</sub>Al, (c) features the results of the EDX line scan for the alloy Ti<sub>6</sub>Al<sub>4</sub>V and (d) for the alloy Ti<sub>6</sub>V<sub>3</sub>Cr<sub>3</sub>Sn<sub>3</sub>Al.

app. 95 nm. The interface between the substrate and the plasma layer is structured due to the large amount of micro roughness caused by the alkaline etching (Fig. 8c). This could be the reason for the irregular nucleation on the surface during the HMDSO deposition. Since the titanium oxide layer was structured on several scales, the layer growth during plasma deposition is inhomogeneous.
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Fig. 4  (a) Extraction area of FIB lamella (Ti$_6$Al$_4$V sample) after alkaline etching and subsequent plasma deposition from HMDSO; (b) SEM micrograph of the extracted lamella with indicate layer thickness (side view).

Fig. 5  (a) SEM micrograph of the extracted lamella with deposited plasma layer on Ti$_6$Al$_4$V; EDX mapping on the extracted lamella with the distribution of the elements (b) Si and Pt, (c) O, (d) Ti, (e) V and (f) Al.

Beside the morphology investigations the element distribution within the extracted lamella was measured by EDX. The result of the element mapping is shown in Fig. 5. Fig. 5b exhibits the distribution of Si. It is obvious that the non-homogeneous silicone distribution could be attributed to the observed open porous HMDSO layer. The amount of Oxygen is homogenously scattered in the lamella. Since the plasma layer consists out of SiO$_x$, the element Ti can only be found in the substrate and the oxide layer. Within the lamella $\beta$-precipitations could be found. In line with surface EDX measurement the $\beta$-phases feature an enrichment of Vanadium and a depletion of Aluminum (Fig. 5e and 5f).

In addition to the investigations on Ti$_6$Al$_4$V the morphology and chemical composition on Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al is determined. Likewise to Ti$_6$Al$_4$V a lamella was extracted by FIB after alkaline etching and deposition of a plasma layer.

Fig. 6a features the SEM micrograph of the area where the FIB lamella was extracted. In Fig. 6 b, the side view of the FIB lamella is shown. It demonstrates that the plasma layer is very homogeneous and compact with the thickness of app. 48 nm. The interface between the plasma layer and the substrate shows any macro roughness due to the smooth surface...
after alkaline etching (Fig. 8d). The different layer growth during plasma deposition is attributed to the difference in the initial oxide layer morphologies.

Fig. 7 shows the element distribution after plasma deposition on Ti15V3Cr3Sn3Al. The EDX element map of silicone (Fig. 7b) shows a homogeneous distribution in the plasma layer which is in contrast to the layer on Ti6Al4V. A small enrichment of oxygen can be determined due to the SiOx film formed on the surface. The nucleation during the HMDSO deposition occurs uniformly and this leads to a homogenous and compact layer.
Fig. 7d-7h demonstrate homogenous distribution of the alloying elements without any enrichments or depletions.

As mentioned above, the different thickness and homogeneity of the deposited silicone dioxide layer on different titanium alloys depend on the morphology of the etched layer. An epitaxial film growth described in the literature [13, 14] is assumed for both alloys. The simplified structure of the different layers created on the two different titanium substrates after alkaline etching step and HMDSO plasma deposition is illustrated schematically in Fig. 8. Moreover SEM micrographs in Fig. 8c and 8b exhibit the different surface morphologies obtained after treatment using Turco 5578. Ti15V3Cr3Sn3Al shows a very smooth surface in comparison to Ti6Al4V alloy. This difference in the degree of micro roughness after alkaline etching has an impact on the morphology and the film formation during the plasma process. The impact on the film formation is schematically shown in Fig. 8a and 8b. The plasma film derived on the initial oxide layer (Ti15V3Cr3Sn3Al), without any micro roughness, leads to a homogeneous surface (red circles in Fig. 8a). In contrast, the micro structured surface of the initial oxide layer on TiAl6V4 causes inhomogeneous and open porous HMDSO layer. The nano-morphology (Fig. 9) and the thickness of the deposited plasma layer are the same in both cases and is app. 50 nm at a working distance of 6 cm. But the micro morphology of the plasma layer depends on the initial oxide layer roughness.
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Fig. 9  SEM micrograph of HMDSO derived plasma layer on titanium alloys (cryo fracture): (a) Ti$_6$Al$_4$V alloy and (b) Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al.

Fig. 10  Wedge test results for Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al treated with BN gas feeding system using compressed air (DL, i.e. Druckluft) as processing gas in comparison to anodizing process NaTESi. The legend features the process variations in the following manner: nominal coating thickness-working distance-treatment angle.

3.3 Investigation of the Bonding Durability

To evaluate the long-term durability of the plasma treated titanium samples in comparison to a wet-chemical treatment (NaTESi anodizing) the wedge test is used. All treatments lead to a comparable initial crack length and bonding strength, respectively. The initial crack length was for all treatments between 25 and 27 mm. Due to this, for all joints the initial failure locus was similar. The failure was initiated cohesively in the middle of the adhesive.

Fig. 10 presents the wedge test results for Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al. The specimen treated with 100 nm nominal thickness at 2 cm and 4 cm respectively. 6 cm working distance features a very good long term durability in comparison to the NaTESi process. All parameters feature a final crack length between 39.5-43.5 mm. After 500 h exposure to 95% rh and 50 °C the titanium samples anodized with the NaTESi process feature a crack growth of 84.4 ± 9.4 mm. The results demonstrate that crack propagation after the different plasma treatments is lower compared to the NaTESi anodising. These results confirm that compressed air can be used for the plasma deposition.
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Fig. 11  Wedge test results for Ti$_6$Al$_4$V treated with BN gas feeding system using compressed air (DL) as processing gas in comparison to anodizing process NaTESi. The legend features the process variations in the following manner: nominal coating thickness-working distance-treatment angle.

on Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al. Moreover, the results show that the working distance can be varied from 2-6 cm on Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al.

Also the long-term durability of bonded Ti$_6$Al$_4$V samples has been investigated. The wedge test results are shown in Fig. 11. The adhesion results of samples treated with NaTESi anodizing process feature a crack length of 43.5 ± 7.1 nm. The specimens treated by plasma lead to a crack growth between 36.8 ± 2.1 nm and 41.5 ± 1.6 nm. In summary, all plasma parameter leads to results which are comparable to the reference anodizing process NaTESi or even better for Ti$_6$Al$_4$V alloy. Likewise to Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al the results confirm that compressed air can be used for the plasma deposition on Ti$_6$Al$_4$V. Moreover, the results show that the working distance can be varied from 2-6 cm on Ti$_6$Al$_4$V.

As shown in Fig. 9, Ti$_6$Al$_4$V and Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al feature a different surface morphology after plasma deposition. The adhesion behavior of titanium alloys depends on the morphology and surface chemistry of the deposited film. The chemical composition of the deposited plasma film and it dependence on the long-term stability of bonded titanium joints is described in previous publications [3, 11]. The wedge test results obtained confirm the correlation between the morphology of the adhesion promotion layer and adhesive bond strength. Ti$_6$Al$_4$V samples show a slightly lower crack length of 38.6 ± 1.8 nm, due to the large amount of nano roughness compared to the Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al samples with the crack length of 41.3 ± 2.9 nm.

4. Conclusions

The results of this work show that the atmospheric pressure plasma deposition of adhesion promoting layer from HMDSO as precursor using long arc generator plasma source (LARGE) is a promising surface pre-treatment technique to improve a good long-term adhesion on titanium alloys.

It has been demonstrated that the titanium β - alloy Ti$_{15}$V$_3$Cr$_3$Sn$_3$Al and the α + β - alloy Ti$_6$Al$_4$V show a different degree of micro roughness after alkaline etching by Turco 5578. The initial oxide layer morphology has a strong influence on the homogeneity and porosity of the deposited HMDSO plasma layer. However, the degree of nano roughness and the thickness of the silicon dioxide layer have
nearly the same value for both alloys. The obtained wedge test results underline the correlation between the durability of adhesion bonds and the surface morphology of the adhesion promoting layer. The higher degree of micro roughness on Ti6Al4V alloy results in a better long-term durability of the bonded samples in comparison to the smooth surface of the bonded Ti15V3Cr3Sn3Al samples.

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References