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Fatigue behavior of a structural steel coated with a WC–10Co–4Cr/Colmonoy 88 deposit by HVOF thermal spraying

J.G. La Barbera-Sosa a,⁎, Y.Y. Santana a, C. Villalobos-Gutiérrez b, D. Chicot c, J. Lesage c, X. Decoopman c, A. Iostc, M.H. Staia a,⁎, E.S. Puchi-Cabrera a,c,d

a School of Metallurgical Engineering and Materials Science, Faculty of Engineering, Universidad Central de Venezuela, Caracas, Venezuela
b School of Mechanical Engineering, Faculty of Engineering, Universidad Central de Venezuela, Caracas, Venezuela
c Université Lille Nord de France, USTL, LML, CNRS, UMR 8107, F-59650 Villeneuve d’Ascq, France
d Venezuelan National Academy for Engineering and Habitat, Palacio de las Academias, Postal Address 1723, Caracas 1010, Venezuela

⁎ Corresponding author. Fax: +58 2126051492.
E-mail addresses: jose.labarbera@ucv.ve, josedlap@gmail.com (J.G. La Barbera-Sosa).

A B S T R A C T

The fatigue behavior of a SAE 4340 steel, coated with a 50% WC–10Co–4Cr/50% Colmonoy 88 deposit, by high velocity oxygen fuel (HVOF) thermal spray, has been investigated. The change in the maximum alternating stress with the number of cycles to fracture has been described by means of the relationship advanced by Stromeyer. A fractographic analysis has been carried out on some representative fracture surfaces, by means of scanning electron microscopy (SEM) techniques. The mechanical properties of the coating were characterized by means of nanoindentation tests. The results indicate that the coating is highly heterogeneous. Its deposition gives rise to a decrease in the fatigue strength of the substrate of ~30%, in comparison with the uncoated substrate. The decrease in fatigue strength is due to the presence of stress concentrators at the substrate–coating interface, as well as the intrinsic characteristics of the coating.

1. Introduction

The aeronautic industry constitutes an important application field of thermal sprayed coatings, particularly as far as dynamic components for planes and helicopters are concerned. These structural elements include, transmission parts and rotor heads, internal cylinders for landing gear, shafts, hydraulic actuators, gas turbine engines components, etc. [1,2]. For these applications, structural aluminium alloys and steels with high strength, toughness and fatigue properties are commonly employed, whose resistance to sliding, and abrasive and fretting wear can be increased by means of adequate coatings.

However, such an improvement in the tribological properties could give rise to a significant decrease in the fatigue life of the coated components, such as it has been found in many different parts coated with electrolytic hard chromium (EHC) plating. Although such a coating gives rise to an improvement in the substrate wear resistance, it occurs at the expense of a decrease in the fatigue performance, in comparison to the uncoated parts [3–10]. Therefore, the qualification of a thermal spray coating employed as a feasible replacement of EHC requires the evaluation of the effect of such a coating on the fatigue properties of the substrate, particularly in critical components such as aircraft engines, for which high cycle fatigue represents the most important failure mechanism [11,12].

Thus, it is expected that the deposition of the coating does not give rise to any decrease in fatigue properties or, at least, if a fatigue debit is induced, it should not be greater than that produced by EHC [1,5,6]. In this sense, a number of investigations have shown that although WC–Co and WC–Ni coatings deposited by HVOF thermal spray give rise to a decrease in the fatigue properties of 4340 steel substrates, it is less than that produced by EHC [8,9,13–15].

In relation to Ni-base self-fluxing coatings deposited onto medium carbon structural steels, previous work has shown an increase in the fatigue strength of the coated system in comparison with that of the uncoated substrate, particularly when tests are conducted in a corrosive medium [16–21]. However, if the tests are conducted in the air, the decrease in fatigue strength could be greater than that produced by WC–Co coatings [22,23].

Flame post-heat treatments have also been employed to modify the microstructure of self-fluxing coatings and therefore, the fatigue properties of the coated systems. In this sense, Akebono et al. [24], conducted an investigation of such a treatment on a Ni-base alloy (Ni–13.7Cr–2.96B–4.40Si–2.67Fe–0.60C; wt.%), deposited onto an AISI 1036 steel substrate. The deposition was conducted by flame spray, employing a mixture of oxy-acetylene, a spraying distance of 200 mm and a coating thickness of 1.5 mm. Post-heat treatments were conducted in a vacuum for 0.5, 4 and 10 h, at a temperature of 1283 K. Rotating bending fatigue tests were carried out at room temperature at a frequency of 50 Hz, employing samples in which the
coatings were machined to 1 mm in thickness, heat-treated and polished in a similar way to that of the uncoated substrate specimens. The results indicated that the fatigue properties of the coated systems were significantly dependent on the size and distribution of the porosity present in the coating, as well as on the hardness of the Ni-rich matrix. The shortest heat-treatment gave rise to the best fatigue performance due to the least amount of porosity, in comparison with the other heat-treatment times. Also, after 10 h of treatment, precipitation and coarsening of Cr-rich compounds resulted in a decrease of the coating matrix hardness. In addition, it was observed that fatigue cracks propagated preferentially through the matrix, which explained the decrease in fatigue properties as the matrix hardness decreases.

Thus, the present investigation has been carried out in order to study the mechanical properties of a composite coating made from the mixture of WC–10Co–4Cr powders with a Ni-base self-fluxing alloy and to analyze the fatigue behavior of the coated system, in comparison with that of the uncoated substrate. The fatigue results are further compared with those observed for the same substrate when it is coated with either WC–12Co or Colmonoy also by HVOF thermal spraying.

2. Experimental techniques

2.1. Substrate alloy and specimens preparation

The present investigation was carried out employing samples of a SAE 4340 quenched and tempered (Q&T) steel, with the following chemical composition (wt.%): 0.41 C, 1.65 Ni, 0.79 Cr, 0.69 Mn, 0.25 Mo, 0.24 Si, 0.024 P and 0.011 S. The material was provided in the form of bars 25.4 mm in diameter and 8 m in length. A number of parallelepiped samples with a dimension of 20×20×8 mm³ were machined in a similar way to that of the uncoated substrate specimens. Subsequently, the specimens were grit blasted prior to deposition. Subsequently, the specimens were grit blasted and organic solvent mixture and pre-heated to eliminate their humidity. The surface of the coating, which was polished prior to testing, to a mirror-like finish, with a mean surface roughness (Rₐ) of 0.09±0.01 μm. Such a surface roughness was determined by means of white light optical interferometry (Zygo New View 200) whose lateral and vertical resolutions are 0.7 and 0.01 μm, respectively. During the characterization of the mechanical properties of coated systems by instrumented indentation, it is well-recognized that the influence of the substrate begins to be noticeable at indentation depths close to 10% of the film thickness, depending on the hardness and the nature of the film [26–29]. On the other hand, for the determination of the elastic modulus, this value is reduced to 1% [30,31]. In this study, in which the coating thickness of the parallelepiped samples was equal to 700±30 μm, the maximum depth reached by the indenter must be limited to 8 μm, in order to avoid the use of models which allow the separation of the influence of the substrate in the hardness and the elastic modulus determination. In the present study, the maximum displacement reached by the Berkovich indenter was 2 μm.

2.2. Coating and spraying conditions

The deposited coating consisted of a powder mixture of 50% WC–10Co–4Cr (Sulzer Metco 5847, Sulzer Metco) and 50% of NiWCrSiFeB alloy (Colmonoy 88, Wall Colmonoy). According to the supplier, the powders have a nominal particle-size distribution of 11–53 μm and 22–62 μm, respectively. The powders were mixed in two steel containers coated internally with Colmonoy 88, employing WC balls and ethyl alcohol, in order to assist the mixing process. For this purpose, a Cole Palmer LABMILL 8000 equipment was employed, which rotated at 47 r.p.m. for 45 min. Subsequently, the mixture was heated to 40 °C for the evaporation of the alcohol and the drying of the mixed powder. The deposition of the coating was carried out both onto the parallelepiped and fatigue samples, which were degreased with an organic solvent mixture and pre-heated to eliminate their humidity prior to deposition. Subsequently, the specimens were grit blasted with an alumina of 0.3–1 mm size range, at a pressure of 0.4 MPa and a distance of 150 mm.

HVOF deposition was conducted employing a Praxair-TAFA JP-5000 gun under the following conditions: spraying distance of 406 mm, powder feeding rate of 83 g min⁻¹, kerosene flux of 23 l min⁻¹ and oxygen flux of 78 l min⁻¹. The final temperature of the specimens was of ~80 °C.

2.3. Nanoindentation tests

Nanoindentation experiments were performed using an MTS nanoindentor equipped with a Berkovich indenter. The values of the loading and unloading rates (expressed in mN min⁻¹) were set at twice the value of the maximum applied load, according to the rule proposed by Quinn et al. [25]. A dwell-time of 15 s was imposed according to the standard indentation test procedures of ASTM E92 and E384-10e2. During the indenter loading, the continuous stiffness measurement (CSM) mode was applied for the determination of the continuous contact hardness, Hₛ, and the elastic modulus of the material, E. 20 indentation experiments were performed randomly at the surface of the coating, which was polished prior to testing, to a mirror-like finish, with a mean surface roughness (Rₐ) of 0.09±0.01 μm. Such a surface roughness was determined by means of white light optical interferometry (Zygo New View 200) whose lateral and vertical resolutions are 0.7 and 0.01 μm, respectively. During the characterization of the mechanical properties of coated systems by instrumented indentation, it is well-recognized that the influence of the substrate begins to be noticeable at indentation depths close to 10% of the film thickness, depending on the hardness and the nature of the film [26–29]. On the other hand, for the determination of the elastic modulus, this value is reduced to 1% [30,31]. In this study, in which the coating thickness of the parallelepiped samples was equal to 700±30 μm, the maximum depth reached by the indenter must be limited to 8 μm, in order to avoid the use of models which allow the separation of the influence of the substrate in the hardness and the elastic modulus determination. In the present study, the maximum displacement reached by the Berkovich indenter was 2 μm.

The coated fatigue samples were tested in the as-deposited condition, without any further surface polishing. Their surface roughness (Rₐ) was determined by means of white light interferometry techniques and it was found to be 6.40±0.90 μm. Fatigue tests were conducted under rotating bending conditions employing Fatigue Dynamics RBF-200 equipment. The fatigue strength was evaluated between 10⁵ and 10⁶ cycles to fracture, that is to say, under high-cycle fatigue conditions. Four different maximum alternating stress levels in the range of 410 to 470 MPa were employed and four samples were tested at each stress level. The tests were conducted at a frequency of 50 Hz.

The maximum alternating stress was determined as a function of the bending moment applied to the specimen by means of the classical relationship:

\[
M_B = \frac{\pi}{32} \sigma_a d^3 \quad (1)
\]

where \(M_B\) represents the bending moment in Nmm, \(\sigma_a\) the maximum alternating stress in MPa and \(d\) the diameter of the coated specimen in mm.

The change in the maximum alternating stress with the number of cycles to fracture has been described by means of the relationship advanced by Stromeyer [32]:

\[
\sigma_a = \sigma_f + \left( \frac{N_f}{K} \right)^{-m} \quad (2)
\]

In the above relationship, \(\sigma_f\) represents the fatigue limit and \(K\) and \(m\) are constants, which depend on the material, loading type and shape and dimensions of the sample. The constants \(\sigma_f\), \(K\) and \(m\) are determined from the experimental values of the maximum alternating stress (\(\sigma_a\)) and the number of cycles to fracture (\(N_f\)) by means of least-square methods.
Finally, a fractographic analysis of some selected samples tested at the highest and lowest maximum alternating stresses was carried out employing SEM-EDS techniques. Such a study was conducted both on the fracture surface of the specimens and on the sections normal to these specimens, with the aim of determining both the fractographic features of the fracture surface and the initiation and propagation sequence of the fatigue cracks.

3. Experimental results

3.1. Characteristics of the initial powders and coating

Fig. 1 illustrates the main characteristics of the powders employed for the deposition of the coating. Fig. 1a shows the size distribution of the WC–10Co–4Cr particles, which ranges approximately between 10 and 40 μm. The particles have a quasispherical morphology, as a consequence of the manufacturing process, which involves both their agglomeration and sintering. Fig. 1b illustrates at a higher magnification (4000×) one of the powder particles, which is characterized by an irregular surface associated with the WC faceted morphology. Also, in Fig. 1c, the size distribution of the Colmonoy 88 particles can be observed. Most of these particles have a quasispherical morphology and a similar size, as a consequence of the atomization process employed for their manufacture. It was determined that the mean diameter of these particles was 30 ± 4 μm. Fig. 1d illustrates a detail of a Colmonoy 88 particle in which the presence of second-phase particles with different morphologies can be observed. Star-like particles and other finely dispersed and smaller particles present in the Ni matrix can be clearly observed. Such particles are associated with the presence of Cr-rich tungsten carbides and/or borides, as well as W and Ni borides and/or carbo-borides [33,34].

Fig. 2a illustrates the general microstructural features of the deposited coating, where two different contrasts can be observed. The regions with a clear contrast are associated with the WC–10Co–4Cr particles, whereas the regions with a darker contrast are associated with the Colmonoy 88 particles. Also, it can be clearly seen that both materials are heterogeneously distributed in volume fractions closely similar throughout the entire coating. The particles of both materials that constitute the coating are observed to exhibit some preferential orientation, which is closely parallel to the substrate–coating interface. The coating deposited onto the fatigue specimens has a thickness of 130 ± 20 μm.

Fig. 2a also shows that the substrate–coating interface is quite irregular, as well as the presence of dark particles of different sizes embedded in the matrix, such as that indicated in the photomicrograph as A, which is constituted by alumina particles employed in the grit blasting process. Fig. 2b, on the other hand, illustrates, at a larger magnification, a region close to the alumina particle identified as A in Fig. 2a, whereas in Fig. 2c a detail of the coating corresponding to the Colmonoy particles is shown. Here, two different zones, B and C, with second-phase particles of different sizes can be clearly observed. Zone B was found to be rich in Ni and Cr, with quadrilateral shaped...
particles rich in W, as well as star-like particles with less amounts of W. According to previous studies [33–35], the latter could be associated either with complex W and Cr carbides and/or borides, as well as carbo-borides or complex Ni and W borides. Region C, on the other hand, was found to have a higher content of W and less Ni, in comparison with zone B. It is constituted by small spherical particles with a mean size of less than 1 μm.

Regarding the regions that correspond to the WC–10Co–4Cr phase, it was determined that these present a continuous Co and Cr matrix, where small WC carbide particles, in the range of 1–3 μm, are present.

### 3.2. Mechanical characterization of the coating by nanoindentation

From the microstructural viewpoint, the coating is composed of WC–10Co–4Cr with a complex Ni base matrix containing W, Cr, Si, Fe and B, distributed in it. Then at a nanoscale measurement, the coating can be considered as very heterogeneous. As a consequence, the values of the mechanical properties can vary to a large extent, depending on the nature and size of the particles located under the indented material, i.e. the proportion of the matrix, single phase, unmelted particles and hard particles, which can be associated with the presence of defects such as pores or inter-lamellar cracks [35,36]. For this reason, 20 indentation experiments were performed randomly at the surface of the coating. 25% of the indentation curves presented irregularities, which were associated with the defects present in the indented volume of the coating. As a result, the corresponding values of the mechanical properties in such a situation have not been considered in characterizing the coating. All the other curves were collected and are presented in Fig. 3.

As can be seen in the above figure, the load-depth curves are not superimposed due to the heterogeneity of the indented material. However, these have been separated into four main groups (G1, G2, G3 and G4), which can be associated with the different natures of the indented materials. Firstly, the indentation curves, which correspond to the lowest indentation depths reached at the same maximum applied load, can be associated with the hardest phases of the coating, probably carbides and/or borides. The other indentation curves attained lower maximum applied loads with the same maximum displacements of the indenter.

Thus, a number of curves are located around 600 mN for the maximum indentation load and another group, around 500 mN. In this situation, it is obvious that the global hardness obtained at 500 mN, correspond to the softer phases, probably the matrix. It is important to point out that in Fig. 3, the first data points corresponding to 30 nm of displacement are not presented since opposite variations were observed, which could not be given any physical meaning. In fact, it is believed that this result was due to a rough approximation of the contact area calculation (polynomial law) due mainly to the rounded tip effect of the indenter and, to a lesser degree, to the surface roughness of the coating.

### 3.3. Fatigue behavior of the coated system

Fig. 4 illustrates that the Wöhler curve of the coated steel is below the curve that corresponds to the uncoated substrate, which indicates that the presence of the coating gives rise to a significant fatigue debit of the substrate, in the range of ~33%. Also, the fatigue limit (σ_f) of the uncoated steel computed from the Stromeyer equation was found to be ~575 MPa, whereas that of the coated system was ~390 MPa, which represents a decrease of ~32%. If the Wöhler curve of the coated substrate is corrected for diameter effects, as shown also in the above figure, taking into consideration that for design purposes the coating is not considered as a load-carrying element of the coated system, the decrease in fatigue strength would be ~25%.

In order to explain the decrease in fatigue strength of the coated system it is necessary to analyze the fracture surface and crack sequence of the coated steel specimens. Thus, Fig. 5 illustrates the general fracture surface of a fatigue sample tested at 410 MPa, which failed after 1,134,800 cycles. At the region identified as O1, a crack initiation site can be clearly observed, located near the periphery of the specimen, from which the crack propagated throughout the cross-section of the sample, leaving behind a relatively flat surface. Also, at 180° of the site O1, a relatively rough surface can be seen, which corresponds to the ductile fracture zone (DFZ). In this zone, where
the final fracture occurred, a number of secondary cracks can be observed at the substrate–coating interface, which gave rise to the delamination of the coating, identified as D in the figure.

Fig. 6a illustrates at a higher magnification the region identified as O1 in Fig. 5. The radial markings, which originate at O1 indicate the crack propagation direction (CPD) along the fracture surface, as shown by the arrows on the photomicrograph. The radial markings converge to a point on the periphery of the sample located at the sub-surface of the substrate, which could be associated with the presence of a MnS inclusion and which would correspond to the nucleation site O1. Fig. 6b illustrates an area close to the region O1 and to the periphery of the sample, in which the fracture surfaces of the coating and substrate can be clearly observed, as well as the presence of sharp alumina particles (P1 and P2) at the substrate–coating interface. On the other hand, Fig. 6c illustrates a simultaneous view of the fracture surface of the substrate (FSS) and coating (FSC), as well as the lateral wall of the coating (LWC), where some delamination (D) of the coating is observed. Fig. 6d shows a magnified view of the defects observed on the surface and indicates that these could be associated with unmelted or re-melted particles of the coating removed from this, possibly during cyclic loading.

Finally, Fig. 7a illustrates a cross-section view normal to the fracture surface of the specimen, where again it can be clearly observed that the thickness of the coating is 130±20 μm. Also, a primary fatigue crack, which propagates close to an unmelted Colmonoy particle (UP), normal to the maximum alternating stress applied to the system, can be seen. The crack seems to have been nucleated at the middle section of the coating and propagates toward the substrate–coating interface (Fig. 7b) through the Colmonoy-rich phases and boundaries between both the Colmonoy and WC–Co–Cr particles.

Regarding the fatigue sample tested at 470 MPa, which failed at 120,800 cycles and shown in Fig. 8, multiple fracture steps (FS) associated with different crack initiation sites could be observed, contrary to the specimen tested at a lower maximum alternating stress. The DFZ appears to be displaced toward the periphery of the sample. A number of secondary cracks both in the substrate (SCS) and the coating (SCC) can also be seen, as well as the delamination of the coating from the substrate. As expected, at this higher alternating stress, the crack propagation zone on the fracture surface is shorter than that observed for the sample tested at 410 MPa.

Fig. 9a illustrates a detailed view of the fracture steps (FS) present on the fracture surface. Such steps are formed as a result of the
intersection of different fatigue cracks, which propagate simultaneously on different planes. Within such steps, a number of radial markings can be clearly seen, which seem to be associated with alumina particles (P1), as shown in detail in Fig. 9b, corresponding to a magnified view of the area indicated in Fig. 9a. Such sharp large alumina particles remain embedded at the substrate surface after grit blasting and become discontinuities, which hinder the adhesion of the coating and promote its delamination.

Fig. 9c, on the other hand, illustrates the lateral wall of the specimen (LWC), together with the fracture surface of both the coating (FSC) and substrate (FSS), where a number of secondary fatigue cracks, parallel to the specimen axis, can be observed. This unexpected cracking pattern could be due to the heterogeneity and relatively high roughness of the coating, with regions of limited adhesion and/or cohesion. The above figure also shows the presence of craters or re-melted particles located on the lateral wall of the sample.

A cross-section view of the coated system, shown in Fig. 9d, illustrates the presence of a primary crack, which has propagated through the entire coating thickness and has bifurcated at the substrate–coating interface, giving rise to the delamination of the coating. As in the case analyzed above, the fatigue crack seemed to propagate preferentially along the lamella boundaries between the Colmonoy and WC–Co–Cr particles. This fact suggests a lack of adhesion and/or cohesion between these materials, which favors crack propagation along such boundaries. However, in some cases it has been observed that fatigue cracks could also propagate through the Ni-rich phases depleted of sub-micrometric second-phase particles, due to their low elastic modulus and hardness.

4. Discussion

Some representative curves have been drawn from Fig. 3, for analyzing the hardness and the elastic modulus variation as a function of the indentation displacement into the material. Fig. 10 represents the contact hardness and the elastic modulus variation as a function of the indenter displacement into the coating associated with the selected and representative indentation curves of each group (G1, G2, G3 and G4), as indicated in Fig. 3. As shown in Fig. 10a and b, the maximum values for the mechanical properties are obtained at the beginning of the loading curve, that is to say, at the outer surface of the coating, when the indenter crosses a hard particle corresponding to the WC.

The elastic modulus value obtained is equal to 550 GPa, in agreement with the value reported by Hyun et al. [37], whereas the contact hardness value of WC was found to be 35 GPa. Since the hard particles have low dimensions in comparison with the indenter displacement, the material surrounding the particles has a significant effect on the hardness measurement. As a result, a fast decrease in hardness and elastic modulus are observed, which tend to achieve constant values of 10 GPa and 250 GPa for the hardness and the elastic modulus, respectively. These values are very similar to those reported by Santana et al. [38,39].
It is important to note that the bulging observed on the hardness variation (Fig. 10a) is probably due to the presence of very small second-phase hard particles, since the hardness value is much lower than 35 GPa. The lowest values for the mechanical properties shown in this figure are also observed as constant values in Fig. 10e and f over the entire loading of the indenter. Only a hardness increase is observed over the first 200 nm of the indenter displacement. These mechanical properties can be associated with region B, rich in Ni and Cr, identified in Fig. 2c. For this region, the elastic modulus varies from 150 to 250 GPa, depending on the constitutive elements of the matrix.

It can be observed that the lowest values of $H_{\text{tr}}$ and $E$ shown in Fig. 10g and h, i.e. 7 GPa for the contact hardness and 200 GPa for the elastic modulus, are very similar to those reported by La Barbera et al. [36]. The hardness variations, which can be observed at the beginning of the indentation curves, are due to the presence of hard particles since the value tends to that characteristic of the mechanical properties of the WC (Fig. 10g). On the other indentation curves, a change is also visible on the first 200 or 400 nm.

The increase or decrease of the hardness value is also associated with the presence of WC as hard particles, but the values attained are much lower than those of this compound due to the small size.
of this particle and the influence of the matrix around the latter. In addition, an interesting result is shown in Fig. 10c and d, in which a constant contact hardness and elastic modulus are obtained, i.e. 12.5 GPa and 300 GPa, respectively. These can be associated with region C, identified in Fig. 2c, where the largest content of W is seen, as well as the presence of sub-micrometric second phase particles. Then, these values are associated with a homogeneous phase at the scale of measurement. It is likely that these mechanical properties correspond to the mechanical properties of various borides, which can be identified in the coating.

Finally, Fig. 11 illustrates a comparison of the fatigue behavior of the coated system investigated in this work and other thermally sprayed coated systems which involve the same substrate steel. Thus, three additional Wöhler curves are presented, which include those of the uncoated substrate, the substrate coated with Colmonoy 88 [22] and the substrate coated with WC–12Co [23]. It can be clearly observed that the deposition of these thermal spray coatings onto the surface of the 4340 steel substrate gives rise to a decrease in the fatigue strength of the substrate. The smallest fatigue debit seems to be associated with the deposition of WC–12Co, followed by the deposition of Colmonoy 88 and finally that which involves the powder mixture of WC–10Co–4Cr/Colmonoy 88. Fig. 12 illustrates the Wöhler curves obtained from the fatigue data reported by Voorwald et al. [4], regarding axial fatigue tests ($R = 0.1$) of uncoated and coated specimens of AISI 4340 with a WC–10Co–4Cr deposited by HVOF. Such results indicate that the deposition of the coating 170 μm in thickness gives rise to a decrease in the fatigue strength of the substrate of up to approximately 40%, depending on the number of cycles to fracture. As the number of cycles to fracture increases, the fatigue debit produced by the coating also increases.

There are many factors that could affect the fatigue performance of these coated systems, which include: grit blasting, spraying gun, spraying distance, flux of the gases employed and thickness and roughness of the coatings, among others. The largest fatigue debit determined in the present work, could be attributed to the larger spraying distance that was employed in coating this system (≈406 mm), in comparison with the other two systems for which the spraying distance was somewhat smaller (≈380 mm). The other deposition parameters were maintained constant for the three coated systems. In this way, as the spraying distance increases, it is likely that the particles reach the substrate or the successive deposited layers at a lower temperature, as shown in Figs. 6d, 7a and 9c and d. In the above figures, a number of unmelted particles and craters, which possibly correspond to re-solidified particles, could be observed.

Another important factor could be the use of different powders, which could compromise the adhesion and/or cohesion among them, giving rise to the presence of weak boundaries, which would favor the propagation of fatigue cracks, as shown in the fractographic analysis that has been conducted. Also, the nanoindentation characterization of the coating showed that there are significant differences in the load-carrying capability of the coating due to the differences in the elastic modulus and hardness. Such differences would enhance the propagation of cracks through the Ni-rich regions of the coating, which are depleted of sub-micrometric second phase particles, which also have a lower hardness and elastic modulus.

5. Conclusions

The deposition of a 50% WC–Co–Cr/50% Colmonoy 88 coating onto a SAE 4340 steel substrate by thermal spraying gives rise to a deposit with a heterogeneous microstructure and therefore, with different mechanical properties, as shown by the nanoindentation characterization that has been conducted. The coated system, with a coating 130 ± 20 μm in thickness, exhibits a decrease in fatigue strength of ~30%, in comparison with the uncoated substrate. The computed fatigue limit according to the Stromeyer relationship (390 MPa) is about 32% less than that of the uncoated substrate (575 MPa). This decrease in fatigue properties for the coated system has been attributed both to the presence of stress concentrators at the substrate–coating interface and to the properties of the coating. The fractographic analysis of the tested specimens suggests that nucleation of fatigue cracks could occur simultaneously at both the substrate–coating interface and the outer surface of the coating. In the first case, nucleation of fatigue cracks would be associated with the irregularities that exist at the interface and the presence of alumina particles, which act as stress concentrators. In the second case, nucleation of cracks would be related to the presence of craters and the high roughness of the coating. Propagation of fatigue cracks through the coating thickness occurs preferentially along the particle boundaries that comprise the coating, particularly through the Ni-rich particles.

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