Self-repairing mechanism and surface characterization of compressor vanes lubricated with oil added with magnesium silicate hydroxide nanorods

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Received: 1 April 2019 / Accepted: 23 October 2019

Abstract. Using magnesium silicate hydroxide as additive of lubricating oils for reducing friction in engineering equipment/machinery has been researched intensively. However, some mechanism relating to the growth of the self-repairing layers on the worn surfaces is still not clearly explained. At the same time, using magnesium silicate hydroxide (MSH) in the form of nanorods showed great promise in reducing friction and wear. In this study, surface-modified MSH in the form of nanorods was used as additive of polyolester oil (POE) which was then used for the lubrication of compressor vanes. The sample parts were studied on the morphology and the microstructure of the self-repairing layer in a great depth. The results showed that self-repairing layers with different thicknesses were generated on the worn surfaces when the POE with 1 wt.% nanorods-MSH was used. It was found that the self-repairing layers consist of organic–inorganic composite membranes, and with increase of working time of the compressor vanes, the self-repairing layers become denser and thicker, while their microstructural form remains to be similar. The situ-repairing capability of the metal surfaces (roller-vane pair of the compressor) enforced by the MSH nanorods is very significant, indicating high potential for industrial applications where boundary and mixed lubrications are needed.

Keywords: Magnesium silicate hydroxide (MSH) / nanorods / polyolester oil / self-repairing / compressor vane

1 Introduction

Ultrafine serpentine powders have gained increasing interest as reactive media stimulating self-repairing of Fe-based metal surfaces subjected to friction [1–5]. However, natural serpentines are defective as they contain impurity elements such as, Fe, Ni and Al, which could influence self-repairing reactions unpredictably. To deal with this problem, pure synthetic magnesium silicate hydroxide (MSH) has become the constituent of most currently used serpentine powders [6–9]. At the same time, performance of the nanorods of MSH with good crystallinity that can be produced by hydration has been studied in recent year, in surface rubbing situations. It was focused mainly on friction reduction behavior. Its anti-wear performance has, however, not been thoroughly investigated [10–13].

Line-contact friction pairs exist in many types of precision machinery, an example being the roller-vane pairs in air conditioning compressors. Vane wear is a severe problem that directly affects compressor efficiency and significantly determines time to failure. Compressor vanes often work in oil at temperatures ranging from 0 to 110°C and are subjected to alternating stresses of 0–345 MPa. Failure criteria of a vane is often referred to wear, e.g. 3.0 μm wear after 500 h continuous work. Therefore, level of the wear is an important indicator for judging the vane’s performance [14–17].

Previously, significant research had been undertaken with a view to decrease the wear in compressor vanes. Long researched compressor vanes with DLC coatings found the wear can be greatly reduced with such coatings [18]. Chen undertook research on nitrided surfaces and found that both surface hardness and wear resistance are greatly improved [19]. Lee et al. used TiN coating to prolong wear life of the compressor vane and found that this could improve the reliability of the compressor [20]. Until now, nitriding is a traditional method often used for strengthening the surface of the vane. Since the mass-scale synthesis of MSH nanorods has been realized and the manufacturing cost has been lowered, using MSH nanorods as a self-repairing additive to POE oil has become reality [21–25].

© L. Wang et al., Published by EDP Sciences 2019
https://doi.org/10.1051/mfreview/2019025
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In the work described in this article, MSH nanorods were added to POE oil and the formation of the self-repairing layers under rubbing friction was investigated. The dynamic formation of self-repairing films and wear conditions were monitored and the mechanism of self-repairing layer formation was analyzed. The results provide a basis for theoretical modelling and for promoting more extensive industrial utilization of MSH nanorods containing oils used for combating wear in machinery and general engineering devices/equipment.

2 Test apparatus, material and experimental procedure

2.1 Nanorods and oil sample preparation

The MSH nanorods were synthesized by a hydrothermal synthesis process. The specific method is as follows: firstly, mixing MgO with Na₂SiO₃ in a molar ratio of 3:2 into distilled water, and then, adjusting the pH of the mixture to 13 with NaOH titration solution. The solution was then placed in an autoclave for reaction with a pressure of about 7.0 MPa. The reaction took place at 280°C for 48 h. After the reaction was completed, the mixture was washed with distilled water until neutral, and finally, it was dried with a dryer at about 100°C for 12 h. After being dried, as shown in Figure 1a, the product was an aggregate of nanorods. The morphology of the synthesized product is shown in Figure 1a. Results of XRD analysis (Fig. 1b) showed that the nanorods are pure MSH and they have good crystallinity. To obtain good dispersity in the oil, the MSH nanorods were mixed with 1 wt.% oleic acid in a planetary ball-mill at 400 rpm for 24 h to obtain nanorods (Fig. 2). As can be seen from Figure 2, the inner diameter of the nanotubes is 5–10 nm and the outer diameter is about 30–40 nm. The surface-modified MSH nanorods have good dispersity. Finally, ultrasonic dispersion (power 200 W, temperature 50°C, duration 60 min) was used to further ensure uniform dispersion in the POE oil.

2.2 Experiment

The relative motion of roller-vane friction pairing in the compressor is shown in Figure 3. In order to simulate real working conditions of a vane, a testing apparatus was built (shown in Fig. 4). The apparatus enabled two steel-vanes to be held against a rotating roller in an environment with nanorods-additive containing oil. Firstly, the surface-modified MSH nanorods were dispersed in the POE with a
concentration of 1 wt.%. The oil pool was then evacuated and the refrigerant RA410 (HFC) inserted, until the pressure reached 4.15 MPa. The oil was heated to 110 °C and the temperature automatically adjusted to around 110 °C during the whole experiment. Two vanes of untreated 11Cr17 steel with hardness 700 HV were positioned in diametric opposition against the side of the roller, which was made with cast iron with hardness HRC 47-57. The experimental parameters are given in Table 1. On expiration of the experimental time the vanes were cleaned with alcohol.

2.3 Characterization and analysis

Each vane was cut into 5 equal lengths along its bearing surface for examination of element distribution on the substrate and the self-repairing layer for each sample length. Topography, both 2D and 3D, of the self-repairing layers was evaluated and micro-roughness of the self-repairing layers was calculated. For simplicity, vanes A and B are designated A and B for the 18 h and 69 h test, respectively.

The topography and chemical composition of the friction surfaces were obtained using SM-7500-F SEM and XRD-7000, respectively. The Raman spectrum of the developed self-repairing layers was further examined using a HORIBA XploRA PLUS Raman spectrometer, and morphology and composition further checked using an OLYMPUS GX51 metallographic microscope and X’Pert Pro MPD X-ray with a spectrometer (EDS) diffractometer, respectively. In addition, Vickers Hardness was measured using HVS – 1000 digital display instrument and AFM analysis was used to present the micro-topography.

3 Results and discussion

3.1 Coefficient of friction

Figure 5 shows how the friction coefficient varies with testing time for two experiments. Because of being similar in work conditions, two curves are virtually similar, up to the test time of 18 h. The coefficient begins at a value of about 0.225 and then sharply declines to a value of about 0.10 during the first 60 min. Then the coefficient reduces more gradually to a value of about 0.075 after 18 h.
Beyond the 18 h testing time the coefficient of the vane B reduces to a value of about 0.07 at 30 h (1800 min) test time and with a slight fluctuation, remains at a value of about 0.075 or less for the remainder of the test.

It is deduced that by prolonging the rubbing time, the coefficient of friction has a declining trend, but the coefficient has a fluctuation in small range towards the end. It is deduced that in the first hour the coefficient sharply decreased from 0.225 to 0.10 for the running-in of the rubbing interface, and fluctuated in a small range throughout the experiment. The friction between the vane-roller is boundary friction and the coefficient of friction is determined by a combination of the fluid friction and the solid friction. The solid friction is greatly influenced by the wear rate and the formation of self-repairing layers. Therefore, the coefficient exhibits a slight fluctuation until the dynamic balance between the wear and new-film formation has been achieved.

### 3.2 Morphology of vanes

Figure 6 shows the morphology of the worn surface of the two vanes. The width of the worn surface of vane A is about 427 μm and there is much corrosive pitting in the friction surface. The width of the worn surface on vane B is about 330 μm. The characteristics of the worn surfaces change from one side of the vane to another. The friction surface was divided into two zones, shown in two different colors: the left side of the worn surface is black and the right bright. It seems that some special substance has been laid on the worn surface and there are fewer corrosive pitting points. Since the vane suffers from elastic deformation during the experiment, thermodynamic conditions are different in the left side and the right side of the worn surface, which influences the formation of the self-repairing layers. This is probably why the morphology of the worn surface of B looks different due to the formation of self-repairing layers locally. In the experiment, left side of the surface first contacted with the roller, then the right side. The large area of the self-repairing layers in the right side of the worn surface of B may be a result of good preservation of the layers due to the relatively mild tribological condition. However, the boundary of A is more distinct than that of B.

The EDS result showed that the characteristic elements Mg and Si are contained in the bright region of A (Fig. 7). Since Si may come from the substrate, the presence of the Mg element indicates physical absorption or chemical reaction between MSH and the substrate.

When MSH nanorods were brought into the friction surface through the oil, they were firstly adsorbed onto the friction surface. Then, under the high pressure and high temperature, the MSH nanorods experienced intraformational sliding as well as a phase change. Products from the phase change were sintered on the surface, mixed with the Fe and Fe2O3. The transition region is a region of the mixed substrate and self-repairing layer. At first, the regions are distinct, as for A; with the extension of time, a region of the friction surface becomes covered by the self-repairing layer.

The partial covering is manifested by the transition line and two regions on the friction surfaces. The substrate partly covered by the self-repairing layer constituted the transition region on the worn surface of A.

Figure 8 shows the EDS result for B and the point is randomly selected in the friction surface. Since the RA410 is constituted by HFC, despite the element F, the other elements indicate that this is the substrate with carbon-enriched layer. The concentration of C is higher than that in the substrate, and it is deduced that the C is partly from the pyrolysis of the lubricating oil. The Mg element was also found on the surface.

It is deduced that with the extension of time, more self-repairing layers are generated on the friction surface of B but they are not continuous. This phenomenon indicates that under severe working conditions (high temperature, high pressure, line contact), a balance of film-growing and wear is not easy to obtain, and the continuous self-repairing layer is difficult to achieve.

After experiments the vanes were cleaned, dried and weighed on an electronic balance of 0.1 mg accuracy. Each vane was weighed 5 times and the results averaged (Tab. 2).

It is clear that after the experiment vane B was heavier than before and vane A decreased its weight. It seems that the self-repairing layers were generated on vane B, and that
although vane A may have some self-repairing layers but their weight was less than the weight of the worn away metal.

3.3 Self-repairing layer

3.3.1 Microstructure of vane A

In order to observe the self-repairing layers from the cross-section, vane A was cut uniformly into 5 segments along axial direction of the rubbing surface (Fig. 9a). After polishing, grinding and corrosion of the cross-section the self-repairing layers were observed through a SEM. For the samples 1–4, morphologies of cross sections are mostly like what seen from the Figure 9b: there are no self-repairing layers observed from the cross-section of these samples. Only sample 5, as shown in Figure 9c, has a thin layer of about 900 nm laid on the friction surface. The layer is continuous and the boundary between the substrate and the layer is not clear. The red line indicates the approximate boundary of the layer, which is not a straight line and follows the contour of the substrate. However, surface of the layer is relatively flat. The morphology indicates that the self-repairing layer is an in situ-layer, and it automatically adapts to the micro-morphology of the substrate. There is no visible crack existing.

Analysis of element distribution is carried out by SEM-EDS. Figure 10 indicates that from the layer to the substrate, C elements decline sharply. Enriched carbon layers are less than 500 nm, which indicates the self-repairing layer has an organic – inorganic composite layer structure. In its surface a thin, enriched carbon layer was generated due to the pyrolysis of lubricating oil. The O element gradually reduced with distance from the layer to substrate where it remained constant. Relating to the result for the Fe element, it is deduced that the self-repairing layers contain iron oxide. When the phase change of MSH occurs, the high-active O produced by the fracture of the hydroxyl will greatly promote iron oxidation, thus, iron oxide participates in the composition of the self-repairing layer. Regarding the characteristic element Mg, coming from the MSH, its content was very small, and therefore, only information concerning the change trend of Mg could be obtained (Fig. 10c). Many of the previous research found no Mg in the self-repairing layers [22–25]. When MSH changes phase, part of the transition products participate in the construction of the self-repairing layer. However, some content is small and it is hard to be detected.

Although vane A loses overall weight, it still has self-repairing layers on the worn surface. The amount of the newly grown layer-materials is, however, not sufficient for compensating for the loss due to wear. The phase change of the MSH was a key factor influencing the generation of the self-repairing layers [26,27].

3.3.2 Microstructure of vane B

Figure 11 shows image of a self-repairing layer viewed from the cross-section of vane B and the line element distribution of the cross-section. Compared with Figure 9c, it is clear that the self-repairing layer generated on the friction surface is thicker. It also has flat surface and its boundary with the substrate is clear. Figure 11 also shows an EDS result, and the element type and change trend are similar to that shown in Figure 9b. These results indicate that with prolonged time, the self-repairing layer becomes denser and thicker, but the composition of the layer material remains the same.

Figure 12 shows the boundary of the working surface and the non-working surface in cross-section. It is clear that the self-repairing layer is only generated on the working surface and its growth depends on the thermodynamic conditions of the friction interface. Therefore, the self-repairing process is in-situ growth of the layer.

3.3.3 Raman analysis

The substrate and the self-repairing layer were analyzed by Raman spectrometry. The wave length was 532 nm, and the Raman displacement Δν was between 500 and 2000 cm⁻¹. Figure 13a shows the result for the substrates of vanes A and B. The peak corresponds to 1363 cm⁻¹ (D-band) near the peak of diamond (1332 cm⁻¹), and it is a symbol of the Sp3 bond. Meanwhile, the peak corresponds to 1590 cm⁻¹ and 1573 cm⁻¹ (close to 1580 cm⁻¹, “g-band”) corresponds to the Sp2 bond, which is graphite carbon.

Table 2. Mass change of A and B.

| Before the experiment | After the experiment | After the experiment |
|-----------------------|----------------------|----------------------|
| | Vane A (g) | Vane B (g) |
| 5.3790 | 5.3643 | 5.3905 |
| 5.3745 | 5.3655 | 5.3899 |
| 5.3763 | 5.3647 | 5.3873 |
| 5.3771 | 5.3652 | 5.3901 |
| 5.3761 | 5.3681 | 5.3872 |
| Average = 5.3766 | Average = 5.3656 | Average = 5.3890 |

![Fig. 9. Images of the cross-sections of vane A.](image-url)
Comparison of the two curves shows that with lengthening time, the C element in the surface began to partly change from orderly carbon (SP2) to disordered carbon (SP3). This means that even although the substrate is not the friction surface, the pyrolysis of lubricating oil will deposit the C element on the surface due to the pressure brought by RA410 refrigerant and the oil. The Raman result proves that the self-repairing layer is an organic-inorganic composite layer, and the first thin-film of the layer is a carbon-enriched film due to the pyrolysis of lubricating oil. The other constructional constituents of the layer need further study.

Figure 13b shows the Raman results for the self-repairing layers of vane A and B. The peak corresponding to 692 cm$^{-1}$ is Fe$_3$O$_4$, 1390 cm$^{-1}$ (D-bond) and 1466 cm$^{-1}$ are near the peak of diamond (1332 cm$^{-1}$) and signifies an Sp3 bond. The peak of 1582 and 1588 cm$^{-1}$ are near the 1580 cm$^{-1}$ (G-bond), and signifies an Sp2 bond. The peak of 1881 cm$^{-1}$ is not very clear. Therefore, it can be given no specific meaning.
It is interesting to notice that there is no peak of Fe$_3$O$_4$ in vane A and also in vane B. The reason for the generation of Fe$_3$O$_4$ is not very clear. It was surmised that the generation of Fe$_3$O$_4$ is due to high temperature internal oxidation produced by the phase change of the MSH [28] and further investigation would be needed in order to confirm it. Nevertheless, it is clear that the content of Fe$_3$O$_4$ increases with time. It is thought that, because the phase change of MSH is intensified with the time, more high-active oxygen is released, which takes part in the building of the self-repairing-layer. Possibly, the explanation that the formation of Fe$_3$O$_4$ is the result of the oxidation of highly reactive oxygen, could help to explain why Fe$_3$O$_4$ can be detected by Raman spectrometry, but it needs to be examined further.

3.3.4 AFM analysis

AFM analysis was used to present the micro-topography of the layers generated on the vane A and B. Figure 14 shows the micro-topography of the self-repairing layers for the worn surfaces of vane A and B respectively. Figure 15 shows the micro-roughness variation along the red lines in the two surfaces.

As can be seen in Figure 14a and b, there were many pits and humps in the worn surface of vane A, while Figure 14c and d show the surface of vane B to be smooth with only a few pits. In addition, the RMS for vane A is 89 nm, which is larger than that for the vane B (65 nm). Large humps appear in the surface of the vane A and the highest hump is 1267 nm, while it is 194 nm only for vane B. This shows, for a longer working time, the self-repairing layer becomes...
thicker and microscopically smoother, which is also confirmed by a comparison on the roughness shown in Figure 15. To further determine the distribution of roughness along the affected surfaces, a detailed surface roughness analysis was carried out. Since generation of a self-repairing surface is, normally, localized, the layers are uneven, and the structure varies with location and time, surface roughness was measured and averaged on four different locations of the two vanes, each test length being 1.75 mm. The results are shown in Table 3.

After 18 h working time for the vane A and 69 h for the vane B, the roughness of the worn surface was 0.083 and 0.07 μm respectively for the vane A and B, and the roughness of the substrate surface 0.105 and 0.110 μm, respectively. For the worn surface, although the micro-roughness of the self-repairing layer for the vane B (Fig. 15) could be smaller than that for the vane A, the overall surface roughness was not improved significantly. The result may help to explain why some previous research reported that in similar experiment, the coefficient of friction became even larger when MSH was added into the oil as an additive. It may also mean that in a boundary lubrication, localized generation of the self-repairing layers could result in an increase of the micro-roughness of the worn-surface and hence, increase of the coefficient of friction. It may also be deduced that the use of MSH nanorods as an additive of the lubricating oil could compensate for the wear, and at the same time, reducing-friction and reducing wear are two competing processes, until self-repairing layers cover sufficient area of the wear while their roughness is also reduced due to new wear, which could be achieved by sufficient length of working time of a friction pair.

4 Conclusions

The work described in this article was the first time that the MSH nanorods were used as an additive to the lubricant for compressor vanes. The test was done in simulated conditions relating to domestic air conditioning compressors. Vane/roller friction and wear phenomenon was
investigated and self-repairing-layer formation mechanism and structures were studied. The following conclusions may be drawn:

- With lengthening working time, the width of the worn surface of the compressor vanes tended to decrease, followed by the growth of self-repairing layers on the worn surface locally.
- With lengthening working time, the self-repairing layer became thicker. EDS showed that element distribution within the layers remained nearly unchanged, regardless of their different morphologies.
- Raman spectrometry showed that both, the substrate and the worn surface, have enriched carbon layers, and with lengthening working time, C gradually changes from orderly Carbon (SP2) to disorderly Carbon (SP3). This indicates that the layers generated on the friction surface are organic-inorganic composites. Their specific structure and generation of Fe3O4 need to be further studied.
- With lengthening time, micro-roughness of the layers reduces and surface of the grown layers become flatter. However, macro-roughness remains almost the same because self-repairing layers were generated locally on the rubbing surface.

The funding from the Natural Science Foundation of China (Grand numbers 51575369) to support the research reported in this paper is acknowledged. M. Wu acknowledges the support of Post-doctoral Research Foundation of Sichuan University (2018SCU12067).

**Table 3. Surface roughness of vane A and B.**

| Roughness of worn surface of vane A (μm) | Ra = 0.060 | Ra = 0.095 | Ra = 0.079 | Ra = 0.097 | Average = 0.083 |
| Roughness of substrate of vane A (μm) | Ra = 0.117 | Ra = 0.098 | Ra = 0.109 | Ra = 0.096 | Average = 0.105 |
| Roughness of worn surface of vane B (μm) | Ra = 0.063 | Ra = 0.075 | Ra = 0.077 | Ra = 0.065 | Average = 0.070 |
| Roughness of substrate of vane B (μm) | Ra = 0.118 | Ra = 0.115 | Ra = 0.096 | Ra = 0.110 | Average = 0.110 |

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Cite this article as: Libo Wang, Yi Yang, Yi Qin, Gang Yang, Yuan Qin, Mingxia Wu, Self-repairing mechanism and surface characterization of compressor vanes lubricated with oil added with magnesium silicate hydroxide nanorods, Manufacturing Rev. 6, 26 (2019)