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Design and Evaluation the Anti-Wear Property of Inorganic Fullerene Tungsten Disulfide as Additive in PAO6 Oil

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Abstract: Inorganic fullerene-like tungsten disulfide particles have been proved to have good anti-friction and anti-wear properties as lubricating materials. As far as we know, however, when it is used as a lubricant additive, its behavior and action mechanism in the friction process are rarely studied. Herein, IF–WS2 particles were synthesized by a Chemical Vapor Deposition (CVD) method. The effect of IF–WS2 particle concentrations in the PAO6 oil on the tribological behaviors was investigated with a four-ball wear machine at both 75 and 100 °C. Additionally, the analyzed morphology and composition of nanomaterials and worn surfaces were analyzed by Scanning electron microscopy (SEM), Transmission Electron Microscopy (TEM) and X-ray photoelectron spectroscopy (XPS). The friction behavior in actual working conditions was studied by a wear testing machine. The experimental results show that compared with the original PAO6 oil, at a dispersion of 0.25 wt% in PAO6 oil, the IF–WS2 particles showed the best performance in terms of coefficient of friction, wear scar diameter and wear mass, which significantly reduced by 27%, 43% and 87%, respectively. At the same time, in the process of friction, it was found that IF–WS2 particles accumulated in the depressions to fill the scratches, and adsorption films and chemical films, including FeS2, WS2 and WO3, were formed on the worn surfaces to avoid the direct contact among the friction pairs more effectively, resulting in the improved anti-wear performances. Additionally, the addition of IF–WS2 particles effectively delayed the rise of lubricating oil temperature. In addition, dispersant span 80 can effectively improve the dispersion and stability of IF–WS2 in PAO6. This work provides us for understanding the effective lubrication mechanism of IF–WS2 particles in more detail and having a new acknowledge of the comprehensive performance of IF–WS2/PAO6 oil.

Keywords: inorganic fullerene; disulfide tungsten; additive; high performance lubricating oil; PAO6; anti-wear

1. Introduction

In modern industry, friction and wear are one of the main causes of materials loss, mechanical failure and energy consumption. Lubricants are widely used to counter friction and wear in an effort to reduce energy consumption and improve the durability of machinery [1–4]. Compared with conventional micro-sized additives, the addition of nano-additives can further improve the performance of lubricating oil in various aspects. Many additives have been developed in recent decades, such as ionic liquids [5–7], nanoparticle additives [8–10], organic ester compounds, etc. [11]. Among them, nanoparticle additives have been widely studied due to their high thermal and chemical stability, as well as excellent mechanical strength. The extremely small size of nanoparticles allows...
them to penetrate contacts of diverse geometries, fill the gaps between contact asperities and ultimately form a protective film under continuous high pressure. The addition of nanoparticles to base oil could significantly enhance the tribological performance of lubricants and offer high capabilities to improve friction, wear resistance, thermal properties, load-carrying capacity and rheological properties [12,13].

Transition metal dichalcogenides, such as molybdenum disulfide and tungsten disulfide, are prone to slip between the crystal sheets due to their unique crystal structure anisotropy, making them an ideal candidate for use as lubricants [9,14–18]. Their unique cage-like structures make the surface of these particles containing little suspended bonds, differing from its 2H type counterpart so that they have better chemical inertness and will be difficult to adhere to the contact surfaces. Many studies have already shown that using inorganic fullerene particles as additives can significantly improve the tribological properties of fluids [19–22]. Rapoport et al. conducted a series of experiments and found that IF–WS$_2$ particles could effectively improve the tribological properties of a material as solid or lubricant [23–25]. Fullerene-like structures can be more effective in reducing friction than corresponding layered structures. Pottuz et al. [26] have studied four possible effects of IF–WS$_2$ particles in lubricating oil and reasonably explained the anti-friction and anti-wear properties of IF–WS$_2$ particles. There is a lot of research as lubricating oil additives, including experiments under boundary conditions. They put forward a variety of mechanisms of WS$_2$ particles in lubricating oil, such as delamination of inorganic fullerene nanoparticles, super lubricity of the sheets, a rolling/sliding of the particles, filling micro-pits on worn surfaces with WS$_2$ flakes and forming friction films with iron oxides and sulfides [17,26–31]. At present, the mechanism of IF–WS$_2$ is still controversial, and their properties still need to be investigated. On this basis, we further studied the potential film composition of friction film under boundary conditions.

The purpose of this work is to analyze the tribological behavior under 75 and 100 °C, attempt to gain an understanding of the specific lubrication mechanism under this condition, and identify the optimal content of IF–WS$_2$ particles for PAO6 oil, which is widely used in the industry. Moreover, the dispersion and stability of IF–WS$_2$ in PAO6 were analyzed by means of transmittance and Scanning electron microscopy (SEM). At the same time, the influence of additives in PAO6 on oil temperature was studied with means of variable temperature simulation test and the change of dynamic viscosity of the lubricating oil before and after friction is studied systematically.

2. Experimental Section
2.1. Synthesis of IF–WS$_2$ Particles

IF–WS$_2$ particles were synthesized by using the Rotary Chemical Vapor Deposition (RCVD) method, according to works of Xu et al. reported previously [32]. Briefly,WO$_3$ powder was used as the precursor, which reacted under H$_2$S/Ar atmosphere at 800 °C for 9 h in a rotary furnace. Before further use, the samples were ultrasonically cleaned with carbon disulfide solution, filtered and dried in an oven at 80 °C.

2.2. Preparation of Lubricating Oil

The base oil used in this study is a low viscosity (5.818 mm$^2$/s for kinematic viscosity at 100 °C) poly alpha olefin (PAO6, Yuchai Research Institute China), which is the common base oil for various industrial synthetic lubricants. The IF–WS$_2$ particles were dispersed to the synthetic PAO6 with different mass percentages of 0, 0.1, 0.25 and 0.5 wt% (IF–WS$_2$/PAO6). In total, 3 wt% Span 80 was used as a surfactant to improve the dispersion stability of additives in oil samples. The IF–WS$_2$/PAO6 solution was mixed using the ultrasonic treatment for 20 min, and then mechanical stirring for another 10 min to obtain uniformly mixed lubricants.
2.3. Tribological Performance Test

The tribological behaviors of pure PAO6 and all IF–WS$_2$/PAO6 samples were studies at two different temperatures of 75 °C and 100 °C by using an MR-10(D) (Jinan Age Testing Machine Co., Ltd., Jinan, China) four-ball friction tester (Schematic diagrams as shown in Figure 1A). The tests were conducted under the conditions of a rotary speed of 1200 rpm, under a load of 392 N and for the duration of 45 min. Before each test, the test balls were washed with petroleum ether to remove organic pollutants on the surface. The friction coefficient was monitored continuously as a function of the number of cycles during the constant temperature test. After the test, the wear scar diameter (WSD) was measured with an optical microscope (Jinan Age Testing Machine Co., Ltd., Jinan, China) equipped with a digital camera. The measurements were repeated 3 times, and the average wear scar diameter (AWSD) value was calculated to ensure the accuracy of the results. Finally, the steel balls, after testing, were cleaned with petroleum ether, dried, marked the wear scar and kept in the sealed sample bags for further SEM and XPS analyses.

![Figure 1. A schematic of (A) the four-ball friction tester used in this study, and (B) the possible lubrication mechanism of IF–WS$_2$ in the PAO6 oil.](image)

The change of oil temperature in the working process was tested using a 280 W needle standard lubricating oil anti-wear testing machine. The friction units are composed of a long rod, a short rod, a weight, a grinding ring rotating with the main shaft and a fixed grinding column. That is, a simplified Timken testing machine, the material used in friction pair is stainless steel. The purpose is to test the friction motion between actual mechanical contact surfaces. The test was carried out under a load of 980 N for 5 min. The data were collected every 20 s, and the average value was obtained after 3 repetitions.

2.4. Characterization

The crystal structure and phase composition of IF–WS$_2$ particles were determined by X-ray diffraction (XRD, Rigaku D/MAX 2500 V, Tokyo, Japan). The morphology of IF–WS$_2$ particles was studied using a Field-Emission Scanning Electron Microscope (FETEM, FEI TECNAI G2 F30, Washington, DC, USA) and energy-dispersive X-ray spectroscopy (EDS) and laser confocal scanning microscopy (KEYENCE VK-X1000, Osaka, Japan), X-ray photoelectron spectroscopy (XPS, Thermo SCIENTIFIC ESCALAB 250Xi, New York, NY, USA) to study the morphologies, elements and their valence changes of the worn surface. The viscosities of the oil samples before and after the test were measured by a viscometer (CP-3000, Lamyett, Guangzhou, China).
3. Results and Discussions
3.1. Characterization of IF–WS₂ Particles

As shown in Figure 2a,b, the IF–WS₂ particles formed in irregular polygonal shapes with an average size of around 100–200 nm. The surface agglomeration of IF–WS₂ spheroids is also observed and probably caused by the fusion of adjacent WO₃ during the heating process. The TEM image (Figure 2c–e) show that IF–WS₂ presented the typical hollow core structures consisting of a large number of layers. The thickness of each layer is 0.62 nm, which matches well with the standard d-space value of the (002) plane of WS₂. Our morphologies results agree well with Tenne et al. works [33]. Moreover, the TEM results also suggest that the agglomeration of these nanoparticles occurred before the conversion of oxide to sulfide [32].

Figure 2. (a,b) The Scanning Electron Microscope (SEM) and (c–e) Transmission Electron Microscopy (TEM) images of IF–WS₂ particles.

Figure 3a shows the XRD diffraction patterns of the synthesized product. The observed diffraction peaks in the XRD pattern are characteristic peaks of the WS₂, which is in accordance with the JCPDS card No. 84–1398 data. The sharp WS₂ peak is at 2θ = 14.2° corresponding to the (002) crystalline planes, of which the layers occur at 90 degrees to the radial direction of the particles. The peaks at 23–25° are attributed to the WOₓ residue in the core of some IFs. The (002) peak shifts to the left, and both (103) and (105) peaks also broaden in the IF–WS₂ XRD profile [34]. All the peak locations in the XRD patterns are also consistent with those reported in the literature [32]. Compared with the characteristic peak position of the XRD spectrum of WO₃ (the place marked by an asterisk), there is nearly WO₃ residue in the product, and the composition of the product is further determined by
XPS. The presence of W and S elements is confirmed by XPS analysis and presented as high-resolution XPS spectra of W 4f and S 2p (Figure 3b). Two S 2p peaks of S 2p$_{3/2}$ and S 2p$_{1/2}$ at 162.2 and 163.2 eV respectively, are corresponded to S$^2$ in WS$_2$ [35,36]. For the W 4f spectrum, the peaks locate at 32.4, 34.5 and 38.0 eV can be assigned to W$^{4+}$ 4f$_{7/2}$, W$^{4+}$ 4f$_{5/2}$ and W$^{6+}$ 4f$_{5/2}$, respectively [35,36]. The existence of W$^{6+}$ may be attributed to WO$_x$ residues in some IFs cores, or the light surface oxidation of particles cannot be ruled out. Based on the XRD, TEM and XPS results, we can confirm that the quality of the IF–WS$_2$ comparable with those reported.

Figure 3. (a) The X-ray Diffraction (XRD) patterns of IF–WS$_2$ NPs; (b) X-ray photoelectron spectroscopy (XPS) spectra of S 2p, W 4f.

3.2. Analysis of Friction and Wear Test

The coefficient of friction (COF) curve, the average coefficient of friction (AF) and the average wear scar diameter (AWSD) of the tested steel ball is presented in Figures 4 and 5. The friction coefficient curve presents a high friction noise at the beginning of the test (time area between 0–250 s) due to the friction has entered the running-in stage. Thus, the iron scraps generated during the friction process are mixed into the PAO6 oil resulting in abrasive wear and increasing the coefficient of friction. The friction coefficient tends to be more stable at the end of the running-in stage due to the formation of a physical or chemical protective film on the surface (Table 1) [37,38]. After the running-in stage, the curve of IF–WS$_2$/PAO6 lubricant tends to decrease gradually over time. It is suggested that a little IF–WS$_2$ particles gathered on the surface of the friction pair during the friction test. Under high pressure, the outer shell of the IF–WS$_2$ with cage structure could be peeled off during the period of the running-in stage, which changed the wear couple from point contact to surface contact to lubricate and reduce the coefficient of friction, just as the research of Maharaj and Joly-Pottuz et al. [39–42]. In addition, we noticed that the friction coefficient curve of the oil sample without span 80 dispersant is higher in Figure 4b. It may be due to the poor dispersion of IF–WS$_2$ in PAO6 oil, and the larger aggregates are not easy to enter the contact area, which leads to the larger friction coefficient and poor lubrication performance.

A diagram of AF of pure PAO6 and IF–WS$_2$/PAO6 samples at 75 °C shows that the pristine PAO6 without IF–WS$_2$ addition exhibits the highest AF of 0.124. The AF of PAO6 with 0.1% and 0.25% IF–WS$_2$ decreases to 0.110 and 0.105, respectively. The COF curves of 0.25 wt% IF–WS$_2$ are relatively stable with time. A similar trend is also observed for the 0.1 wt% of IF–WS$_2$ samples during the first 1750 s, followed by a continuous increase until the end of the record. We suggest that this behavior might be justified by the aggregation and sedimentation phenomena of the IF–WS$_2$ particles, which made it hard for them to get
access evenly to the contact areas and therefore limited their presence between the moving surface. When the content of IF–WS\textsubscript{2} particles increased to 0.5 wt\%, compared with the pure oil PAO6, its friction curve fluctuates greatly, with an AF value of 0.121. Based on this experiment, we confirm that the anti-friction effect of base oil can be enhanced with the addition of the IF–WS\textsubscript{2} additives. The AF is reduced by 11.3, 15.3 and 2.4\% for 0.1, 0.25 and 0.5 wt\% IF–WS\textsubscript{2} additive content, respectively, benchmarked against the pure PAO6.

At 100 °C, the AF of pure PAO6 oil increases to 0.135, which was higher than the test result at 75 °C. When 0.1, 0.25 and 0.5 wt\% of IF–WS\textsubscript{2} particles were added, the AF reduces to 0.0978, 0.0979 and 0.0974, respectively. Its anti-friction performance slightly improves with the increase of IF–WS\textsubscript{2} additive content, by about 27\% compared with that of pure PAO6. In short, the IF–WS\textsubscript{2} additives exhibited a better anti-friction effect than pure PAO6, particularly at higher temperatures.

Figure 4. The coefficient of friction (COF) curve of IF–WS\textsubscript{2} with different contents at temperatures (a) 75 °C and (b) 100 °C with the load of 392 N and rotation speed of 1200 rpm.

![Figure 4](image)

Figure 5. (a) average coefficient of friction (AF) and (b) average wear scar diameter (AWSD) diagram of different IF–WS\textsubscript{2} contents at 75 °C and 100 °C.

![Figure 5](image)
Table 1. Average COF values at different time.

| Temperature (°C) | Content of IF–WS₂ (wt%) | Average Coefficient of Friction (COF) |
|-----------------|-------------------------|-------------------------------------|
|                 |                         | 500 s | 1000 s | 1500 s | 2000 s |
| 75              | 0                       | 0.124 | 0.124 | 0.122 | 0.127 |
|                 | 0.1                     | 0.104 | 0.096 | 0.085 | 0.127 |
|                 | 0.25                    | 0.121 | 0.097 | 0.086 | 0.083 |
|                 | 0.5                     | 0.124 | 0.129 | 0.088 | 0.142 |
| 100             | 0                       | 0.130 | 0.147 | 0.144 | 0.146 |
|                 | 0.1                     | 0.097 | 0.088 | 0.081 | 0.086 |
|                 | 0.25                    | 0.096 | 0.086 | 0.072 | 0.067 |
|                 | 0.5                     | 0.094 | 0.086 | 0.096 | 0.096 |

The AWSD of PAO6 containing IF–WS₂ particles at 75 °C is slightly larger than that of pure PAO6, which increases by 6.7%, 10% and 11.7%, respectively, and it weakens the anti-wear performance of base oil, as shown in Figure 5b. At 100 °C, the AWSD obtained for lubricants based on PAO6 containing 0.1, 0.25 and 0.5 wt% of IF–WS₂ additives are 0.59, 0.56 and 0.63 mm, respectively. Compared with 0.99 mm of the pristine PAO6, which converts to a decrease of AWSD by 40.4%, 43.4% and 35.4%. The reason for the above situation may be that the test temperature affects the lubrication performance of additive IF–WS₂ particles. At high temperatures, the degree of freedom of particles in base oil PAO6 is relatively large. Once the protective film is destroyed, they can form boundary films on the contact surface faster. However, the viscosity of the base oil is relatively high at low temperatures, which limits the movement of particles. The particles at the contact interface cannot be replenished in time, resulting in serious wear and tear [43].

As shown in the point plot of Figure 5, some data of AF and AWSD show a valley trend with the change of IF–WS₂ concentration, and there is a common valley point between them, which corresponds to the optimal content of IF–WS₂. In addition, the anti-friction and anti-wear performance of pure PAO6 oil is poor. It may be the dispersion of high concentration IF–WS₂ in lubricating oil decreases. IF–WS₂ additive has a large amount of agglomeration and deposition, which results in poor wear [44–46]. The error bar in the figure shows that the repeatability of the experiment is high, and the error is less than 3%.

The effect of WS₂ on the wear performance of PAO6 was analyzed and with 3D topography (Figure 6) and wear profile (Figure 7). At 75 °C, there are many wear scratches on the worn surfaces of oil PAO6 (Figure 6a) and 0.25 wt% IF–WS₂/PAO6 (Figure 6b). Additionally, the scratches of 0.25 wt% IF–WS₂/PAO6 are deep and even dense, deep furrows appear, and the right side of the worn surface is more obvious, as shown in Figure 7a. Its surface roughness is larger, and the wear is more serious. Compared with PAO6, wear surface area increases from 10,581.03 µm² to 24,402.96 µm² (Table 2). The furrows of 0.25 wt% IF–WS₂/PAO6 contain more wear debris, which may be caused by viscosity at low temperatures. It prevents the wear debris and IF–WS₂ in scratches from flowing out with lubricating oil in time and accumulating into large particles, which causes abrasive wear and ploughed the worn surfaces leads to more furrows.

Table 2. The average wear area data of IF–WS₂/PAO6 lubricants with different contents at different temperatures.

|                  | 75 °C       | 100 °C      |
|------------------|-------------|-------------|
|                  | PAO6 | PAO6 + 0.25% | PAO6 | PAO6 + 0.25% |
| Average area (µm²) | 10581.03 | 24402.96 | 28801.48 | 12062.06 |
Figure 6. 2D and 3D morphologies of wear surfaces of pure PAO6 and 0.25 wt% IF–WS$_2$/PAO6 at 75 °C (a,b) and 100 °C (c,d).

Figure 7. Wear profile curve of pure PAO6 and 0.25 wt% IF–WS$_2$/PAO6 at 75 °C (a) and 100 °C (b).

To further analyze the lubrication process of the IF–WS$_2$ additive, the surface morphology of wear scars that occurred with different additive concentrations were observed, as shown in Figure 8. The SEM images show that the wear scar of the pristine PAO6 oil looked very rough, there are large areas of damage and debris (Figure 8a$_2$,a$_3$), and the worn surface exhibit many large pits and deep furrows at 100 °C (Figure 8c$_2$,c$_3$). Compared with the 75 °C samples, the damage at 100 °C is more serious, and the overall anti-wear performance is poor. In contrast, the wear scar for samples of PAO6/IF–WS$_2$ became smoother. The roughness and debris of the wear scar surface are significantly reduced, although the width of the wear scar is not reduced at 75 °C. This may be attributed to the
agglomeration and the chemical-curing phenomenon of nanoparticles with a high concentration. Therefore, it is hard for WS$_2$ to form a continuous and uniform anti-friction layer on the friction region [47]. The defects become shallower, which made the wear slightly reduced, as shown in Figure 8b$_2$-b$_3$. The wear scar width and the plough grooves became much smaller at 100 °C. The pits basically disappear without obvious defects, the friction surface became very smooth, which indicates a better wear resistance (Figure 8d$_2$-d$_3$).

![Figure 8. WSD photos and SEM images of the worn samples after tests. (a) pristine PAO6 oil at 75 °C, (b) 0.25 wt% IF–WS$_2$ particles at 75 °C, and (c) pristine PAO6 oil at 100 °C, (d) 0.25 wt% IF–WS$_2$ particles at 100 °C.](image)

The results of each EDS diagram show that W and S elements are distributed on the worn surface of PAO6 lubricating oil containing IF–WS$_2$ (Figure 9), which confirmed that these particles are indeed located at the contact interface and are involved in the friction process. They directly fill the grooves or peeled off sheets of the out layers from the WS$_2$ to fill the pits, thus repairs damages to the surface. This sort of tribo-film can improve the wear resistance, as illustrated by Tenne et al. [48]. These analyses are consistent with the test data, also prove that the addition of additive IF–WS$_2$ particles can obviously improve anti-wear performance of PAO6 oil. The higher the temperature is, the more obvious effect will improve.
Figure 9. EDS profiles of the surface after friction process samples, under (a) pristine PAO6 oil and (b) PAO6 containing 0.25 wt% IF–WS$_2$ at 100 °C; and (c) pristine PAO6 oil and (d) PAO6 containing 0.25 wt% IF–WS$_2$ particles at 75 °C.

XPS is used to analyze the change of chemical state of elements in the PAO6 oil during friction, which is very helpful for understanding the lubrication mechanism of the additives. As shown in Figure 10, despite the high temperature, the XPS spectra of typical elements C 1s, S 2p, W 4f and Fe 2p are observable on the worn surface, which is obtained after correlating with the C 1s binding energy of 284.8 eV. There are three peaks of C–C/C–H (284.8 eV), C=C (286.6 eV) and C=O (288.6 eV) in the C1s spectrum [46,49], which shows that the addition of additives has no influence on the structure of organic content in the PAO6 oil. At 75 °C, the XPS spectra of S 2p had almost no difference, and the peak at a binding energy of 168.6 eV corresponded to the S$^{4+}$ state and occurred in FeSO$_4$ or Fe$_2$(SO$_4$)$_3$ [43] (Figure 10a,b). The XPS spectrum of W 4f consists of 3 peaks, corresponding to WS$_2$ (33.2 eV) and WO$_3$ (35.4 and 37.8 eV), respectively, indicating that the wear trace contains IF–WS$_2$ NPs or peeling pieces [43,50]. For the Fe 2p spectrum, there are Fe$_2$(SO$_4$)$_3$, Fe(OOH), Fe$_2$O$_3$ or Fe$_3$O$_4$ signals, at binding energies of 724.8, 711.5 and 710.4 eV, respectively [43,46]. In particular, the FeO (709.3 eV) peak appears on the worn surface in the pristine PAO6 oil sample, whilst in the IF–WS$_2$/PAO6 sample, a similar peak exists at 706.9 eV, which matches with FeS$_2$ [43]. The new peak of Fe$_2$S seems to have confirmed that a chemical reaction between the exfoliated pieces of IF–WS$_2$ particles and the steel ball matrix occurred. When the test temperature is constant at 100 °C, the S 2p spectrum exhibits sulfide (near 162.5 eV) and sulfur oxide/sulfate (168.9 eV) signals, while the sulfide signal value with additives is weak [43,50]. The W 4f spectrum shows the signals of the two substances the same as that observed at 75 °C. The FeS$_2$ peak also appears at the binding energy of 706.7 eV of Fe 2p for the 100 °C pristine PAO6 samples (Figure 10c), which may be that the test temperature rose and the Fe compound reacted with the original S element in the steel ball. However, this is not for samples with WS$_2$ at 100 °C, as shown in Figure 10d. According to the peak analysis of S 2p from 100 °C IF–WS$_2$/PAO6 sample, the FeS$_2$ peak should have appeared in Fe 2p, which possibly because the worn surface is covered by most WS$_2$ films. It leads to weakened or even the disappearance of the FeS$_2$ signal. This consists well with the results of our SEM analysis.
Figure 10. XPS spectra of the elements (C 1s, Fe 2p, S 2p and W 4f) on the worn surfaces, lubricated by pristine PAO6 at (a) 75 °C, (b) 75 °C and (c) 100 °C, and by PAO6 containing 0.25 wt% WS$_2$ at (d) 100 °C.

A schematic of the lubrication mechanism is proposed, as shown in Figure 1B. Chemical reaction occurred on the contact surface during friction, and tribo-films formed. For the pristine PAO6 oil, iron filings are oxidized to form boundary protective films of Fe$_2$O$_3$, Fe$_3$O$_4$, Fe(OOH) and Fe$_2$(SO$_4$)$_3$. In the case of using IF–WS$_2$ additives, the formed boundary protective films consist of FeS$_2$, WS$_2$ and WO$_3$, apart from Fe$_2$O$_3$, Fe$_3$O$_4$, Fe(OOH) and Fe$_2$(SO$_4$)$_3$. The protective films were very complex and advanced, were attached to the friction surface of the steel balls, which prevented its direct contact between the wear couple. The IF–WS$_2$ additives could make up for the micro-pits and damages caused by the uneven distribution of iron-related oxides and therefore reduced friction and exhibited improved anti-wear performance.
3.3. The Dispersion Stability of IF–WS\textsubscript{2} in PAO6 Oil

The dispersion stability of particles was detected by the transmittance of an ultraviolet spectrophotometer, and the transmittance of the system was the smallest, which indicated that the system had the best dispersion stability. Figure 11 is a graph of the relationship between different addition contents and transmittance with wavelengths of (a) 400 nm and (b) 630 nm. It can be seen from Figure 11 that with the passage of time, IF–WS\textsubscript{2} particles with various initial concentrations have a tendency of gradually gathering and sinking in PAO6. (a) and (b) indicate that 0.1% IF–WS\textsubscript{2} lubricating oil after standing for 24 h has particle aggregation, which leads to a decrease in the concentration of upper particles, and the curve transmittance at this time is about 5%. It may be because 0.25% and 0.5% of IF–WS\textsubscript{2}/PAO6 are saturated, while 0.1% is not saturated, which is different. The light transmittance is greater than 0.25% after standing for a period of time. After standing for 72 h, the transmittance of 0.25% and 0.5% IF–WS\textsubscript{2} oil samples is different, and the 0.5% IF–WS\textsubscript{2} oil samples aggregate faster, which may be due to the high concentration of IF–WS\textsubscript{2} particles in lubricating oil, the large interaction force between particles, and the weak stability of micelle formed by span 80 coating, which leads to particle aggregation. The difference between them is more obvious after standing for 120 h. After standing for 192 h, the transmittance of the two is basically the same, which may be because the gravity of 0.25% IF–WS\textsubscript{2} plays a major role at this time, resulting in obvious particle aggregation and sedimentation.

![Figure 11. Graph of the relationship between different addition contents and transmittance with wavelengths of (a) 400 nm and (b) 630 nm.](image)

In order to analyze the reason why IF–WS\textsubscript{2} particles disperse well in PAO6 oil, the dispersion effect of IF–WS\textsubscript{2} particles was observed by scanning electron microscope. Figure 12a is an SEM image of 0.25% PAO6/IF–WS\textsubscript{2} without active agent span 80. The figure shows that IF–WS\textsubscript{2} particles have an obvious agglomeration phenomenon. Figure 12b is an SEM image of 0.25% PAO6/IF–WS\textsubscript{2} containing active agent span 80. The figure shows that IF–WS\textsubscript{2} with span 80 dispersant has good dispersivity and no obvious agglomeration. The reason is that the long-chain alkane at one end of the active agent span 80 has good compatibility with the base oil PAO6, which increases the wettability of IF–WS\textsubscript{2} in PAO6 and reduces its surface energy. The other end is adsorbed on the surface of IF–WS\textsubscript{2} particles to form a solvated film, as shown by the arrow in Figure 12b. At the same time, the polymer adsorbed by particles will produce new repulsive energy, increase the space distance, weaken the attraction between molecules, and effectively prevent the mutual contact between particles, thus maintaining a stable state and reducing agglomeration [51,52]. Span 80 can effectively improve the dispersion of IF–WS\textsubscript{2}, thereby improving the performance of IF–WS\textsubscript{2} in POA6 lubrication performance.
3.4. Physical Properties of the Lubricating Oil

The actual working condition of the PAO6 oil with IF–WS\textsubscript{2} particles is evaluated by a 280 W pointer standard lubricating oil anti-wear testing machine. The change of oil temperature during the friction test is shown in Figure 13a. In the working process, the oil temperature of pristine PAO6 oil reaches 89.9 °C at 200 s and then tends to stabilize. The temperature of those containing 0.1, 0.25 and 0.5 wt% IF–WS\textsubscript{2} reached 50.5, 48.5 and 54 °C, respectively, at 300 s, which have a slower rate of oil temperature rise and a lower platform trend. These results showed that the additives effectively delay the rise of oil temperature, of which the 0.25 wt% content is the most effective. IF–WS\textsubscript{2} has good heat transfer properties in oil, similar to layered graphite \cite{53,54} and carbon nanotubes \cite{55}. IF–WS\textsubscript{2}/PAO6 can obviously reduce the wear caused by weight loss (the difference between the mass before and after friction), as shown in Figure 13b. Additionally, the weight loss reduced by 87.8% for the 0.25 wt% sample. It may be that the oil temperature rises at different rates, which impels different temperatures and wear levels at the same stage.

In order to better understand the influence of additives on the physical properties of the PAO6 oil, the dynamic viscosity of lubricating oil before and after the wear experiments was measured, and the results are shown in Figure 14. Before the wear test, with the change
of temperature, the dynamic viscosity difference of IF–WS\(_2\)/PAO6 with different contents is different, as shown in Table 3. With the increase of additive contents, the difference of dynamic viscosity change is smaller, which illustrates IF–WS\(_2\) the improved viscosity–temperature performance of PAO6. Meanwhile, the dynamic viscosity at 75 °C and 100 °C fluctuated around the average values of 10.01 mPa·s and 6.47 mPa·s, respectively. It should be noted that, due to the existence of dispersant span 80, the measured value of dynamic viscosity is larger than the standard value. After the friction test, the dynamic viscosity of lubricating oil with different additives has slightly fluctuated at 75 °C, the influence of IF–WS\(_2\) on viscosity was less than 6.6%. Conversely, the influence of IF–WS\(_2\) on viscosity is less than 2.5% at 100 °C.

![Figure 14. Dynamic viscosities of the oil samples containing different IF–WS\(_2\) additives before and after the wear test at 75 °C and 100 °C.](image)

|          | 0%           | 0.1%          | 0.25%         | 0.5%          |
|----------|--------------|---------------|---------------|---------------|
| 75 °C    | 10.02 mPa·s  | 10.04 mPa·s   | 9.8 mPa·s     | 10.16 mPa·s   |
| 100 °C   | 6.24 mPa·s   | 6.51 mPa·s    | 6.39 mPa·s    | 6.75 mPa·s    |
| Differences | 3.78 mPa·s | 3.53 mPa·s   | 3.41 mPa·s    | 3.41 mPa·s    |

4. Conclusions

In this paper, the behavior and film composition of IF–WS\(_2\) additives in lubrication were studied. the tribological behavior of IF–WS\(_2\)/PAO6 has been evaluated at two different temperatures, 75 and 100 °C, using a four-ball wear machine. The conclusions are summarized as follows:

1. At 75°C, IF–WS\(_2\) only reduces the friction coefficient, which is not good for anti-wear. At 100 °C, and the effect is bidirectional. At this moment, the optimum addition content is 0.25% IF–WS\(_2\) particles.

2. In the friction process, IF–WS\(_2\) particles are transferred and precipitated on the friction surface under the action of high temperature and pressure. IF–WS\(_2\) particles flake off and react chemically with the friction pair matrix to form a chemical film. In addition to the oxide layers (Fe\(_2\)O\(_3\), Fe\(_3\)O\(_4\), Fe(OOH), Fe\(_2\)(SO\(_4\))\(_3\)) formed on the friction surface, the newly generated FeS\(_2\), WS\(_2\) and WO\(_3\) protective films are attached to the friction
surface of steel balls, hindering the direct contact of friction pairs, the pits and defects caused by the uneven distribution of iron-related oxides are covered by film. Under the combined action of iron oxide and sulfide, the wear performance is improved.

(3) Span 80 makes IF–WS$_2$ particles uniformly dispersed in PAO6 base oil and effectively improves the dispersion of IF–WS$_2$. Under the influence of span 80 dispersant, IF–WS$_2$ with the content of 0.1% has the fastest sedimentation rate, while IF–WS$_2$ with the content of 0.25% has better dispersion stability.

(4) The IF–WS$_2$ additives have shown great influence on the oil temperature rise, and they have effectively reduced and delayed the oil temperature rise, which in turn delay the oxidation of lubricating oil. The addition of IF–WS$_2$ additive improved viscosity-temperature performance of PAO6 lubricating oil. Meanwhile, after the friction test, the influence of IF–WS$_2$ on dynamic viscosity is less than 6.6% and 2.5% at 75 °C and 100 °C, respectively.

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