Tribological behaviour of novel chemically modified biopolymer-thickened lubricating greases investigated in a steel-steel rotating ball-on-three plates tribology cell

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Abstract

In this work, model renewable and biodegradable lubricating greases based on castor oil and chemically modified biopolymers (methylcellulose, chitin and cellulosic pulp) were tribologically characterized in a steel-steel ball-on-plates tribological cell coupled to a controlled-stress rheometer, and the results were compared to those provided by conventional lithium and calcium soap-based greases. Viscous flow, sliding velocity sweep and transient friction tests were carried out and resulting wear scars in the steel plates were evaluated by means of scanning electron microscopy (SEM). Different frictional responses were found depending on the thickener, especially at high temperature. Excepting for most severe conditions, wear is negligible when using chemically modified chitin and methylcellulose-based greases as lubricants.

Keywords: Biopolymer-based lubricating greases; friction; rheology; wear.
INTRODUCTION

Lubricating greases are highly structured suspensions in which the core composition consists of a thickening agent dispersed in a lubricating base oil, optionally complemented with additives in order to improve some specific properties and functions [1]. Nowadays, commercial lubricating greases are mostly mineral oil-based whereas most commonly used thickeners are fatty acid soaps of lithium, calcium and aluminium, di- or polyureas and inorganic thickeners like clays and silica derivatives. Observations by electron and atomic force microscopy [2,3] revealed that the thickener forms a network, with different structure depending on the thickener agent, that traps the oil and confers the desired rheological properties [4,5]. These particular rheological characteristics imparted by the thickener make lubricating greases more convenient in some applications where lubricating oils do not work properly. Actually, factors like decreasing leakage and frequency of lubrication in hard-to-reach contacts, fluctuations with temperature, loads, vibrations, etc. can be minimized when using greases [6]. In addition, lubricating greases can absorb contaminants, such as particles and water, in relatively large amounts, without reduction in its lubricating properties thus acting as effective seals [7,8]. As previously reviewed [6], the mechanisms of grease lubrication in bearings are still under discussion. In the past [9,10], the lubricating function of greases was only attributed to the ability of the base oil to bleed in a lubricated contact, however, more recent works highlight the importance of thickener particles on contact surfaces [11-13]. The presence of the thickener particles give rise to an increase in the effective viscosity and thickness of the elastohydrodynamic lubrication film [14-17], resulting in a significant reduction of the friction coefficient with respect to those found for the base oil alone. Therefore the thickener also plays an important role in the lubrication process.

The preservation of the environment, as well as the corresponding governmental directives in numerous countries, promotes the increasing introduction of renewable raw materials in different industrial applications, namely in the lubricant industry [18]. As extensively reported [19-21], the use of vegetable oils is gaining some importance due to a number of valuable
characteristics such as high lubricity, easy adhesion to metal surfaces, weak viscosity-temperature dependence, low volatility, solvency for lubricant additives, easy miscibility with other fluids, low toxicity and high biodegradability. Despite limitations such as poor thermo-oxidative stability, deficient low temperature flow properties due to crystallization and small range of viscosities that restrict the applicability of vegetable oils, previous studies demonstrated that these disadvantages can be overcome using proper additives [22-26]. However, global production of vegetable oil-based greases is marginal so far, i.e. lower than 1%, as a consequence of still poorer performance characteristics [27]. The development of fully bio-based lubricating greases is even a more challenging issue that needs not only the replacement of the mineral oil by a suitable vegetable one but also the selection of an appropriate thickening agent obtained from renewable resources. Oil substitution has been satisfactorily addressed [28-30], but the replacement of the thickener is a much more complex task to solve due to the high technical efficiency of traditional metallic soaps and urea derivatives, which impart the desired rheological, thermal and tribological properties to the bulk system. Up to now, vegetable oil-based calcium soap- or clay-thickened greases are considered environmentally-friendly, although showing some performance limitations like, for instance, the lower thermal resistance of calcium greases. Some lubricating greases based on biodegradable oils with traditional thickeners were tested in different tribological contacts [28,31] showing low values of friction coefficient and small wear scar size for some specific formulations, basically depending on the balance between oil polarity and type of thickener.

Previous research reported the formulation of oleogels thickened with cellulose derivatives which exhibit satisfactory mechanical and physical stabilities as well as more accurate thermal resistance than that found for conventional lubricating greases [32-34]. However, in these cases, suitable mechanical and physical stabilities were linked to much higher values of the linear viscoelastic functions than those found for standard greases. More promising results were recently found when using diisocyanate compounds to either functionalize biopolymers with reactive groups [35,36] or create an extensively crosslinked network [37] able, in both cases, to
retain the base oil. Despite the fact that isocyanate compounds may be considered hazardous, when the biopolymer is dispersed in castor oil, there is no more free NCO groups but urethane linkages, formed after the chemical reaction between NCO and hydroxyl groups located in the ricinoleic fatty acid chain. Therefore, the thickener chemically interacts with the base oil, resulting in the oleogel physical stabilization [38]. As has been reported, polyurethanes based on biodegradable polyols [39], and particularly biopolymers containing urethane and carbamate linkages [40], are considered largely biodegradable materials. Taking into account all the previous research on the finding of proper new bio-based thickeners and, especially, considering the suitable rheological behaviour of formulations based on biopolymers chemically modified with isocyanate compounds, the objective of this work was to offer new experimental tribological data of NCO-modified methylcellulose, chitin and cellulosic pulp dispersions in castor oil. These formulations, completely based on renewable resources have been selected as promising grease candidates. Their performance has been compared with those found in two commercial formulations based on lithium and calcium soaps.

EXPERIMENTAL

Materials

Castor oil purchased from Guinama (Spain) was selected as vegetable oil medium to prepare oleogel formulations. Viscosity values of castor oil are 550 and 8.25 cP at 25 and 125 °C, respectively [41]. Methylcellulose (Mn: 40000 g·mol⁻¹; DS: 1.6-1.9), acquired from Sigma-Aldrich; chitin (Mw: 5.4×10⁵ g·mol⁻¹; degree of deacetylation: 7.3%), purchased from Qingdao Fraken (China), and commercial grade Kraft cellulose pulp from Eucalyptus globulus, kindly supplied by ENCE, S.A. (Huelva factory, Spain), were used as thickening agents after modification with 1,6-hexamethylene diisocyanate (HMDI, purum grade, ≥98.0%). HMDI and all other common reagents and solvents employed were supplied by Sigma-Aldrich. Details of biopolymer chemical modifications with HMDI can be found elsewhere [35-38]. Two commercial lubricating greases were used as benchmarks: a conventional mineral oil-based
lithium grease (Castrol, Germany) and one considered biodegradable based on castor oil and an anhydrous calcium thickener (kindly supplied by Verkol, Spain).

**Manufacture of model biopolymer-based lubricating greases**

The dispersion of the thickeners in the base oil was carried out in an open vessel, using a controlled-rotational speed mixing device (70 rpm) RW 20 (Ika), equipped with an anchor impeller. NCO-functionalized methylcellulose and chitin were slowly added to the castor oil at the same concentration (30% (w/w)), whereas the thickener based on cellulosic pulp was used in a concentration of 7% w/w. The mixing process was maintained at 70 rpm for 24 hours at room temperature. Finally, the resulting dispersion was homogenized with an Ultra-Turrax T50 (Ika) rotor-stator turbine, at 8800 rpm for 1 min in the case of the methylcellulose and chitin-based oleogels and 5 min for the cellulosic pulp-based oleogel, and afterwards they were de-aerated by applying vacuum. Table 1 summarises the composition of the different formulations obtained along with identifying codes and basic technical data. Commercial grease samples and composition of biopolymer-based greases were selected attending to their similar rheological response previously analysed [35-37].

**Atomic force microscopy (AFM)**

Microstructures of the two commercial greases (LI and CA samples) were obtained by means of atomic force microscopy (AFM) with a Multimode apparatus connected to a Nanoscope-IV scanning probe microscope controller (Digital Instruments, Veeco Metrology Group Inc.). Images were acquired in the tapping mode using Veeco Nanoprobe™ tips. Samples were not submitted to any physical modification or partial oil extraction but previously heated below the dropping point. Therefore, this technique allows obtaining the microstructure of untreated samples. LI and CA microstructures are shown in Figure 1 and may be compared to those previously reported for the biopolymer-based greases studied [37,42], which display fibrous structures, more similar to the lithium grease.

**Rheological characterization**
Rheological characterization was carried out with a controlled-stress rheometer, Physica MCR-501 (Anton Paar, Austria). Viscous flow tests were performed applying a stepped shear rate ramp in a shear rate range of $3 \cdot 10^{-3}$ to $10^2 \text{ s}^{-1}$ at 25 and 125 ºC. Rheological measurements were carried out using a plate–plate geometry (25 mm diameter, 1 mm gap) with grooved surfaces to overcome wall slip phenomena usually observed in lubricating greases [43]. At least two repeats of each test were performed on fresh samples.

**Tribological Tests**

Tribological tests were performed in a tribology measuring cell coupled to the Physica MCR-501 controlled-stress rheometers [44]. The cell uses a 6.35 mm diameter steel ball (1.4401 grade 100) rotating on three 45º inclined steel plates (1.4301). The evolution of the friction coefficient, $\mu$, with the rotational speed was tested in a range of rotational velocities of 0.01-1000 rpm, at 25 and 125 ºC and different normal loads (10, 20 and 40 N). These normal forces as well as sliding velocities were calculated from the applied axial force, ball radius and angular velocity according to Heyer and Läuger [44]. Estimated contact Hertzian pressures for each normal load applied are 1.29, 1.63 and 2.05 GPa, respectively. The stationary friction coefficient was also obtained at 25 and 125 ºC by applying normal forces of 10, 20 and 40 N and setting a constant rotational speed (0.15, 10 and 400 rpm) for 10 min. Five repeats of each test were performed on fresh samples. Resulting wear scars on the steel plates were examined by triplicate using a scanning electronic microscope, model JSM-5410 (JEOL).

**RESULTS AND DISCUSSION**

**Viscous flow behaviour**

Viscous flow curves of the lubricating greases studied were obtained at 25 and 125 ºC. Figure 2 shows the comparison at 25 ºC, which reveals very similar viscosity values, excepting for sample CH which is more viscous in all the shear rate range studied. Shear rate range is limited due to the appearance of fracture at high or moderate shear rates [43-45], which is favoured with
grooved measuring surfaces [46]. Resulting flow curves can be satisfactorily described by the well-known power-law model:

\[ \eta = K \cdot \dot{\gamma}^{n-1} \] (1)

where \( K \) and \( n \) are the consistency and flow indexes, respectively. Fitting parameters are listed in Table 2 for the different samples and temperatures studied. As expected, the consistency index \( (K) \) presents lower values at 125 °C with respect to those found at 25 °C. As previously mentioned, the highest consistency index at 25 °C was achieved by the oleogel based on chemically modified chitin (CH), and then followed by MC and CP, respectively. Commercial lubricating greases, LI and CA, present consistency indexes that are slightly lower than those obtained for the biopolymer-based formulations. On the other hand, consistency indexes of these model formulations showed intermediate values with respect to the commercial greases at 125 °C, excepting for the sample CP, which exhibits lower value than CA. On the other hand, extremely low values of the flow index were generally obtained for all samples, which is characteristic of the typical yielding behaviour shown by these materials.

**Evolution of the friction coefficient with the sliding velocity**

The frictional behaviour of greases analysed as lubricants in the steel-steel ball-on-plates contact was first characterized by applying a rotational speed sweep. Figure 3 shows the friction coefficient vs. sliding velocity curves, at 25 and 125 °C, for the formulations studied and selected normal force (20 N). As can be observed, at 25 °C, the general progression from boundary to elastohydrodynamic lubrication regimes were found for all samples, excepting for CH which mainly displays the transition from the mixed lubrication to the elastohydrodynamic regimes (Figure 3i). However, when temperature was increased up to 125 °C, higher rotational speed values are generally needed to detect the increasing part of the frictional curve, i.e. the transition from the mixed to the elastohydrodynamic regimes. In fact, the final increasing part of the friction coefficient vs. the sliding velocity curve was not observed at 125 °C when applying the commercial lubricating greases used as benchmarks (Fig. 3b and 3d). On the contrary, the
different lubrication regimes are still apparent at 125 °C for the model biopolymer-based greases studied.

The classical Striebeck curve relates the friction coefficient (μ) with the dimensionless Hersey number, which is used to describe the combined influence of dynamic viscosity (η), rotational speed (Ω) and the normal contact load applied (F_N) in journal bearings. Instead, in similar geometries and conditions to that used in this work, the so-called hydrodynamic Striebeck parameter is defined as [47,48]:

\[ S = \frac{\eta u_s}{F_N} \]  

being \( u_s \), the sliding velocity. This parameter takes into account the fluid film formation between the two surfaces in a tribo-contact and discriminates the lubrication regimes as boundary, mixed lubrication or (elasto)hydrodynamic [49]. The Striebeck curve is widely used to describe the lubrication regimes of Newtonian lubricating oils, including bio-based oils [47, 50-53]. However, this approach is not widely reported for lubricating greases as a consequence of the non-Newtonian character and the unknown properties of the active lubricant inside the contact [16]. Lu and Khonsari [54] and Gonçalves et al [55] proposed the inclusion of the viscosity of the base oil, assuming that under highly stressed conditions, like those achieved in a tribological contact, the microstructure of the grease is almost destroyed and the main responsible for lubrication is the base oil:

\[ S = \frac{\eta_{BO} u_s}{F_N} \]  

As Gonçalves et al pointed out [55], the bleed oil viscosity is not affected by the type and concentration of the thickener and, according to Palacios and Palacios [56], grease viscosity tends to reach the base oil viscosity at extremely high shear rates. Obviously, this rule cannot be directly followed when using polymeric additives able to modify the base oil viscosity [57].
Moreover, the role of the thickener in the tribo-contact is not considered in eq. (3). Accepting the inclusion of the base oil viscosity in the Strubeck parameter, among all the lubricating grease samples studied only CA and MC display a simple tribological behaviour which acceptably leads to a unique Strubeck master curve by superimposing the frictional curves obtained at different temperatures and normal loads (Figures 4a and 4b). This means that base oil viscosity and normal load govern film formation in these two greases, as expected for a Newtonian lubricant. In the remaining formulations, a single Strubeck curve cannot be obtained. In particular, when using CP sample as lubricant, the shape of the friction coefficient vs. sliding speed curve did not significantly change by modifying normal force or temperature, displaying maximum and minimum values of the friction coefficient at approximately the same values of the sliding velocity. An intermediate behaviour was observed in the case of CH sample. Strubeck curves calculated at different normal load and temperature conditions overlap each other only in the elastohydrodynamic region, whereas a scatter in the friction coefficient values, generally decreasing with the applied normal force, was found in the mixed lubrication regime (Figure 4c). Therefore, in these cases, the role of the thickener in the film formation seems to be more evident and greatly influences the frictional behaviour.

**Evolution of friction coefficient with time**

The evolution of friction coefficient with time at different normal loads and rotational speeds was analyzed for each sample. A comparison of transient values of the friction coefficient is shown in Figure 5 for the lubricating greases studied and selected conditions. Generally, at the beginning, the friction coefficient rapidly drops and then progressively decreases down to a stationary value. This evolution has been traditionally attributed to the formation of the protective lubricating film due to the rubbing action [58] and, particularly in greases, to the rheological resistance that the ball has to overcome to plough its way at the beginning of motion [28]. However, under certain conditions, i.e. high normal loads, some of the lubricating greases evaluated produce a slight increase of the friction coefficient after the first drop, probably as a consequence of the effect of resulting wear debris. At 25 °C, the lower values of the friction
coefficient were obtained by using sample MC as lubricant and conversely the higher friction was found with sample CP (Figure 5a). At this temperature, the effect of normal load is always that shown in Figure 5b for a selected sample. However, temperature greatly affects this tendency, as discussed below.

In general, a stationary value of the friction coefficient was achieved after approximately 5-6 minutes when using the biopolymer-based greases as lubricants, whereas commercial samples needed longer times (8-10 min). The bar diagram shown in Figure 6 portrays the steady-state friction coefficient values obtained by applying normal loads of 10, 20 and 40 N at constant angular velocity of 10 rpm and temperatures of 25 and 125 ºC, for each grease used as lubricant in the tribo-contact. Samples such as LI and CH display the same tendency at 25 and 125 ºC, i.e. the friction coefficient decreases with the normal load. This is not the expected normal force influence in the boundary and mixed lubrication regimes [54] but it is consistent with the higher rheological resistance offered by the grease to the rotational motion at lower normal loads [28]. Moreover, for both samples, the friction coefficient values are even lower at higher temperature, in accordance with the lower rheological resistance offered as a result of the thermally-induced softening of greases. At the same time, microstructure is better preserved at 125 ºC in these two samples, showing the higher values of the consistency index (Table 2). As a consequence, it is presumed that at high temperature the rheological resistance to the motion is reduced but the entrainment of the whole lubricant into the contact takes place. On the other hand, greases CA, CP and MC follow the former trend at 25 ºC, but completely the opposite at 125 ºC. This effect may be attributed to the fact that these microstructures are more strongly affected by shear and temperature, as the lower values of the consistency index suggest (Table 2), resulting in a significant reduction of the effective viscosity and, therefore, in the lubricant film thickness, which favours wear, especially at higher normal loads, as discussed in the next section. In fact, this temperature is rather close to the dropping point in sample CA (see Table 1) and therefore a more significant softening and oil bleeding is expected. This hypothesis is supported by the fact that a single Stribeck master curve was obtained for CA and MC samples when using the base
oil viscosity to estimate the Stribeck parameter. Alternatively, MC and, in lower extent, CP microstructures may release much more oil at high temperature and normal loads, as observed after performing the frictional tests, thus locally producing, in some parts of the tribo-contact, an increase in the effective concentration of the cellulosic thickening agent, which can interact with the metallic contact surfaces, causing higher friction and wear.

Figure 7 shows the values of the stationary friction coefficient by applying 20 N normal force and rotational speeds of 0.15, 10 and 400 rpm, at both 25 and 125 °C. The friction coefficient obtained when using the commercial lubricating greases, LI and CA, and formulation CP as lubricants, at 25 °C, decreases with the rotational speed, whereas the opposite tendency is observed in the case of CH at 25 °C. The sample MC at 25 °C displays a particular behaviour that first exhibits a reduction of the friction coefficient from 0.15 to 10 rpm, but a notably higher value at 400 rpm. At 125 °C, the tendency followed by almost all the samples, excepting for CP, reveals an increase in the friction coefficient from 0.15 to 10 rpm and a diminution from 10 to 400 rpm. In the case of the formulation based on the cellulosic pulp, at 125 °C, a remarkably high value at 0.15 rpm can be observed; however, from 10 to 400 rpm, the friction coefficient raises with the rotational speed. These tendencies followed by the stationary friction coefficient in every case are absolutely consistent with the dynamic frictional measurements previously discussed, i.e. the sliding velocity curves, and reflect the different lubrication regimes.

**Wear**

Wear scars generated in the plates of the tribological cell and the corresponding average diameter are shown in Table 3 for selected lubricating grease samples and conditions. As can be concluded from the images, wear mechanism is predominantly abrasive. As expected, the stronger normal load is applied, the wider wear scar is, as can be seen in the bar diagram in Figure 8, which summarizes the evolution of the scar size with the normal load, at 25 and 125 °C. However, temperature and normal load do not exert a significant influence when using the lithium grease as lubricant. Sample CP provide values of the average wear scar diameter
comparable to those found with the commercial greases, excepting for the highest normal load applied, and generally lower at high temperature (Figure 8). In the particular cases of MC and CH, no wear evidence in the plates was observed, excepting for 20 and 40 N at 125 °C, being especially wide for MC.

Figure 9 displays the average wear mark diameters obtained from the steel plates of the tribological cell after performing tests at 0.15, 10 and 400 rpm under 20 N at 25 and 125 ºC, respectively. As can be observed, the increase in wear diameter with the rotational velocity is the predominant tendency, at both temperatures. Once again, the samples based on methylcellulose (MC) and chitin (CH) only provide appreciable wear scars under specific conditions. At 25 ºC, steel plates only evidence wear scars at 400 rpm when using MC sample as lubricant, whereas rather large scar diameters can be observed at 125 ºC under 10 and 400 rpm conditions. In the case of CH, significant wear evidence can be only detected at 400 rpm and 125 ºC.

CONCLUSIONS

Several new biodegradable model lubricating greases based on NCO-chemically modified biopolymeric thickener agents were tribologically characterized. Results were compared with those obtained when using conventional lithium and calcium soap-based greases with alike rheological properties as lubricants. Viscous flow curves of all samples were very similar and can be satisfactorily described by the traditional power-law model. Greases containing NCO-functionalized chitin and methylcellulose exhibit higher consistency indexes at 25 ºC, whereas this parameter is higher for lithium soap and chitin-based greases at 125 ºC, which is indicative of microstructures with higher thermal resistance. In all cases, fracture phenomenon clearly arises at moderate shear rates, which limits the study to not very high shear conditions.

The general progression from boundary to elastohydrodynamic lubrication regimes were found when using the biopolymer-based greases as lubricants in the steel-steel ball-on plates tribo-contact, at both 25 and 125 ºC, when performing sliding velocity sweep tests. However, the
elastohydrodynamic regime was not detected at 125 °C when using conventional greases. Only calcium soap and methylcellulose-based samples display a simple tribological behaviour which leads to a unique Stribeck master curve by superimposing the frictional curves obtained at different temperatures and normal loads, using the base oil viscosity in the hydrodynamic Stribeck parameter. This fact suggests that the base oil governs film formation in these two greases, whereas the thickener seems to play a key role in the remaining samples. The stationary friction coefficient always decreases with the normal load applied at 25 °C. However, the opposite tendency was found at 125 °C for calcium soap-, methylcellulose or, in lower extent, cellulosic pulp-based greases as lubricants. These results depend on the balance between the rheological resistance offered by the grease and the shear and temperature dependence of grease microstructure. In this sense, particularly calcium soap and methylcellulose microstructures, are more strongly affected by temperature giving rise to a significant reduction of the lubricant film thickness which favours wear at 125 °C. This fact is especially evident at high normal loads. However, excepting for such severe conditions, wear is negligible when using chemically modified chitin and methylcellulose-based greases as lubricants, whereas cellulosic pulp-based grease provides wear scars comparable to those found with the conventional lithium and calcium greases.

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Table 1. Formulation details of model biopolymer-based and commercial lubricating greases studied.

<table>
<thead>
<tr>
<th>Substrate (g)</th>
<th>Crosslinker (HMDI) (g)</th>
<th>Thickener concentration (% w/w)</th>
<th>Base oil</th>
<th>Formulation code</th>
<th>Penetration\textsuperscript{a} (dmm)</th>
<th>Dropping point\textsuperscript{b} (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methylcellulose (8.0)</td>
<td>4.1</td>
<td>30</td>
<td>castor oil</td>
<td>MC</td>
<td>271</td>
<td>185</td>
</tr>
<tr>
<td>Chitin (12.0)</td>
<td>7.6</td>
<td>30</td>
<td>castor oil</td>
<td>CH</td>
<td>219</td>
<td>223</td>
</tr>
<tr>
<td>Cellulosic pulp (5.0)</td>
<td>5.0</td>
<td>7</td>
<td>castor oil</td>
<td>CP</td>
<td>252</td>
<td>156</td>
</tr>
<tr>
<td>Lithium soap</td>
<td>unkown</td>
<td></td>
<td>mineral oil</td>
<td>LI</td>
<td>260</td>
<td>195</td>
</tr>
<tr>
<td>Calcium soap</td>
<td>unkown</td>
<td></td>
<td>castor oil</td>
<td>CA</td>
<td>324</td>
<td>145</td>
</tr>
</tbody>
</table>

\textsuperscript{a} According to ASTM D-217
\textsuperscript{b} According to ASTM D-2265
Table 2. Power-law model fitting parameters of the viscous flow curves of biopolymer-based (CP, MC and CH) and commercial lubricating greases (LI and CA) studied at 25 and 125 ºC.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Temperature (ºC)</th>
<th>n</th>
<th>K</th>
<th>R²</th>
</tr>
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<tbody>
<tr>
<td>LI</td>
<td>25</td>
<td>0.067</td>
<td>704.0</td>
<td>0.9977</td>
</tr>
<tr>
<td></td>
<td>125</td>
<td>0.248</td>
<td>98.1</td>
<td>0.9906</td>
</tr>
<tr>
<td>CA</td>
<td>25</td>
<td>0.137</td>
<td>720.0</td>
<td>0.9972</td>
</tr>
<tr>
<td></td>
<td>125</td>
<td>0.131</td>
<td>35.4</td>
<td>0.9901</td>
</tr>
<tr>
<td>CP</td>
<td>25</td>
<td>0.107</td>
<td>782.4</td>
<td>0.9985</td>
</tr>
<tr>
<td></td>
<td>125</td>
<td>0.130</td>
<td>26.3</td>
<td>0.9982</td>
</tr>
<tr>
<td>MC</td>
<td>25</td>
<td>0.066</td>
<td>960.9</td>
<td>0.9996</td>
</tr>
<tr>
<td></td>
<td>125</td>
<td>0.125</td>
<td>52.7</td>
<td>0.9989</td>
</tr>
<tr>
<td>CH</td>
<td>25</td>
<td>0.079</td>
<td>2171.8</td>
<td>0.9972</td>
</tr>
<tr>
<td></td>
<td>125</td>
<td>0.312</td>
<td>74.7</td>
<td>0.9958</td>
</tr>
</tbody>
</table>
Table 3. Selected SEM images and corresponding average diameter of wear scars on plates obtained at 25 and 125 ºC, constant rotational speed (10 rpm) and different normal loads (10, 20 and 40 N) using samples CP, LI and CA as lubricants. Images are presented at different magnifications just to give an idea of the predominant wear mechanism.

<table>
<thead>
<tr>
<th></th>
<th>25 ºC</th>
<th></th>
<th></th>
<th>125 ºC</th>
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<th></th>
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<tbody>
<tr>
<td></td>
<td>10 N</td>
<td>20 N</td>
<td>40 N</td>
<td>10 N</td>
<td>20 N</td>
<td>40 N</td>
</tr>
<tr>
<td>CP</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>303 µm</td>
<td>331 µm</td>
<td>786 µm</td>
<td>254 µm</td>
<td>259 µm</td>
<td>410 µm</td>
</tr>
<tr>
<td>LI</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>278 µm</td>
<td>273 µm</td>
<td>284 µm</td>
<td>285 µm</td>
<td>291 µm</td>
<td>343 µm</td>
</tr>
<tr>
<td>CA</td>
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<tr>
<td></td>
<td>255 µm</td>
<td>321 µm</td>
<td>401 µm</td>
<td>304 µm</td>
<td>403 µm</td>
<td>716 µm</td>
</tr>
</tbody>
</table>
FIGURE CAPTIONS

Figure 1. AFM micrographs for commercial lubricating greases: a) lithium grease; b) calcium grease. (Image width: 20 μm).

Figure 2. Viscous flow curves at 25 ºC for commercial and model biopolymer-based lubricating greases studied.

Figure 3. Friction coefficient versus sliding velocity curves obtained for the different grease samples used as lubricants. (Normal force: 20 N; Temperature: 25 and 125 ºC, respectively).

Figure 4. Strubeck curves for lubricating grease samples CA (a), MC (b) and CH (c).

Figure 5. Evolution of the friction coefficient with time, at 25 ºC and a constant rotational speed (10 rpm) as a function of (a) lubricating greases (normal load: 10 N), and (b) normal force (sample MC).

Figure 6. Stationary friction coefficient values obtained by applying a constant rotational speed (10 rpm) and different normal loads (10, 20 and 40 N), at 25 and 125 ºC, for the different lubricating greases studied.

Figure 7. Stationary friction coefficient values obtained by applying 20 N constant normal force and different rotational speeds (0.15, 10 and 400 rpm), at 25 and 125 ºC, for the different lubricating greases studied.

Figure 8. Average diameter of wear scars on plates obtained by applying a constant rotational speed (10 rpm) and different normal loads (10, 20 and 40 N), at 25 and 125 ºC, for the different lubricating greases studied.

Figure 9. Average diameter of wear scars on plates obtained by applying 20 N constant normal force and different rotational speeds (0.15, 10 and 400 rpm), at 25 and 125 ºC, for the different lubricating greases studied.
Figure 1
Figure 2

Viscosity (Pa·s) vs. Shear rate (1/s) for different materials (LI, CA, CP, MC, CH).
Figure 3
Figure 4
Figure 5
Figure 6
Figure 7
Figure 8

Wear scar diameter (μm)

25 ºC 10 N
25 ºC 20 N
25 ºC 40 N
125 ºC 10 N
125 ºC 20 N
125 ºC 40 N

LI
CA
CP
MC
CH
Figure 9