NATIONAL ADVISORY COMMITTEE FOR AERONAUTICS

TECHNICAL NOTE

No. 1432

CHANGES FOUND ON RUN-IN AND SCUFFED SURFACES OF STEEL CHROME PLATE, AND CAST IRON

By J. N. Good and Douglas Godfrey

Flight Propulsion Research Laboratory
Cleveland, Ohio

Washington
October 1947
A study was made of run-in and scuffed steel, chrome-plate, and cast-iron surfaces. X-ray and electron diffraction techniques, micro-hardness determinations, and microscopy were used. Surface changes varied and were found to include three classes: chemical reaction, hardening, and crystallite-size alteration. The principal chemical reactions were oxidation and carburization with the following reaction products:

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Surface condition</th>
<th>Material identified on surface</th>
<th>Questionable identification</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAE 4140 steel reciprocating slider</td>
<td>Unrun</td>
<td>α-Fe</td>
<td>Fe₃O₄</td>
</tr>
<tr>
<td></td>
<td>Worn unsuffed</td>
<td>α-Fe</td>
<td>Fe₃O₄, Fe₂O₃</td>
</tr>
<tr>
<td></td>
<td>Scuff recovered</td>
<td>α-Fe, Fe₃O₄</td>
<td>Fe₃C, Fe₂O₃</td>
</tr>
<tr>
<td></td>
<td>Scuffed</td>
<td>α-Fe, Fe₃C, Fe₃O₄</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>SAE 4140 steel aircraft-engine cylinder barrel</td>
<td>Worn unsuffed</td>
<td>C (graphite), Beilby layer</td>
<td>Mixed oxides</td>
</tr>
<tr>
<td></td>
<td>(Superficial face)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(Less than 0.001 in. below surface face)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chrome-plated cast-iron rider</td>
<td>Worn unscruffed</td>
<td>α-Cr, Cr₂O₃, Cr₃C₂</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Burnished</td>
<td>α-Cr, Cr₂O₃, Cr₃C₂</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Scuffed</td>
<td>α-Cr, Cr₂O₃, Cr₃C₂</td>
<td></td>
</tr>
<tr>
<td>Cast-iron reciprocating slider</td>
<td>Unrun</td>
<td>α-Fe</td>
<td>Fe₃C</td>
</tr>
<tr>
<td></td>
<td>Worn</td>
<td>α-Fe</td>
<td>C (graphite)</td>
</tr>
<tr>
<td></td>
<td>Scuffed</td>
<td>α-Fe</td>
<td>C (graphite)</td>
</tr>
</tbody>
</table>

¹Lines best fit pattern of material listed.
Hardness of materials examined varied with degree of surface condition. The crystallite sizes were in the order of $10^{-7}$ centimeters. The sizes changed with condition of operation.

**INTRODUCTION**

The changes that occur to rubbing metallic surfaces have been studied by many investigators in order to obtain a better understanding of the phenomena of wear and wear resistance. (See the bibliography.) A study of the bibliography and the reference reports reveals that at least the following factors are of importance:

1. Formation of surface films and reaction products
2. Changes in surface hardness
3. Changes in crystallite size

Some relations are established herein among these factors and the phenomena of run-in and one phase of wear, scuffing, by identification of and examination of surface changes occurring with one of the well-known sources of metallic wear, reciprocating sliding. The term "run-in" is used in this report to designate the process through which surfaces acquire, by mutual interaction during sliding contact, increased capacity to carry load and to withstand wear. This investigation neglects the relation of hydrodynamic and boundary lubrication to run-in, although all surfaces studied were lubricated.

Changes in the surface composition and surface characteristics were studied under run-in and wear conditions for steel (SAE 4140) rubbed against cast iron, chrome-plated cast iron rubbed against steel, and cast iron rubbed against cast iron; these combinations are standard in aircraft engines. Use was made in this investigation of specialized techniques in X-ray and electron diffraction, of chemical and metallographic analyses, and of hardness measurements.

**APPARATUS AND PROCEDURE**

Specimens of standard aircraft-engine cylinder-barrel and piston-ring material combinations used in these examinations were obtained from two sources. Portions of used cylinder barrels and piston rings were directly examined, whereas portions of unused cylinder barrels and piston rings were examined after rubbing under pressure in a reciprocating slider machine. Surface changes on the specimens were examined by
X-ray and electron diffraction. Supplementary information was obtained by chemical and microchemical analysis, electrographic analysis, metallographic examination, and microhardness determination.

The following materials were investigated:

<table>
<thead>
<tr>
<th>Material combination</th>
<th>Specimen examined</th>
<th>Specimen source</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAE 4140 steel slider rubbed against cast-iron rider</td>
<td>Slider</td>
<td>Slider machine</td>
</tr>
<tr>
<td>Cast-iron piston rings rubbed against SAE 4140 steel</td>
<td>Cylinder</td>
<td>Aircraft engine</td>
</tr>
<tr>
<td>cylinder barrel</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SAE 4140 steel slider rubbed against chrome-plated cast-iron rider</td>
<td>Rider</td>
<td>Slider machine</td>
</tr>
<tr>
<td>Cast-iron slider rubbed against cast-iron rider</td>
<td>Slider</td>
<td>Slider machine</td>
</tr>
<tr>
<td>Chrome-plated cast-iron piston rings rubbed against SAE 4140 steel cylinder barrel</td>
<td>Ring</td>
<td>Aircraft engine</td>
</tr>
</tbody>
</table>

This selection of specimens is representative of the materials used in standard aircraft engines and should give results representative of run-in and scuff in those engines.

The slider machine (fig. 1) reciprocated a slider specimen 7\(\frac{1}{2}\) inches beneath a fixed and loaded rider 1\(\frac{1}{8}\) inches. The rider was loaded by applied gas pressure, which was read from a dial. Mass temperatures of the rider and the slider were recorded with thermocouples. When the surfaces are lubricated, as in this investigation, conditions of both boundary and hydrodynamic lubrication are believed to exist, that of boundary at the ends of the stroke and that of hydrodynamic at the middle of the stroke. The oil used was Navy 1120 fed at a rate of 16 cubic centimeters per hour. The reciprocating speed was 650 cycles per minute with a stroke of 5\(\frac{1}{2}\) inches. The procedure used for all specimens was as follows:

1. The two mating surfaces were lapped with silicon carbide lapping compound with resultant surface-roughness readings of approximately 40 to 60 microinches rms for the rider and below 15 microinches rms for the slider.
2. The surfaces were cleaned with solvents.

3. The surfaces were mated by reciprocation under a constant load of 600 pounds per square inch applied to the rider for 15 minutes. Run-in began at this point.

4. For scuffed specimens, the rider was loaded in increments of 100 pounds per square inch every 5 minutes until scuffing occurred. A prolonged run-in procedure, consisting of operation at 2400 pounds per square inch for 2 hours followed by 100 pounds per square inch and temperature-stabilization increments until scuffing occurred, was used on one cast-iron surface. Scuffing observed visually, accompanied by rapid temperature rise, served as a criterion of surface failure. The machine was stopped within a few seconds after failure began, which was usually 2 to 3 hours after the start of the run.

The aircraft engines were operated for \( \frac{5}{2} \) to 7 hours of run-in plus 15 to 100 hours of endurance at an average engine speed of 2500 rpm and an average indicated mean effective pressure of 250 pounds per square inch.

All X-ray diffraction patterns were obtained with cobalt Ka radiation and with the specimen mounted in the manner used with the Debye-Scherrer powder camera. A 5-inch-diameter cylindrical camera was used with an iron-oxide filter. Diffraction-line intensities were visually estimated. A microphotometer could be used only to record α-iron X-ray-line position, breadth, and intensity for crystallite-size determination. Positive identification of compounds was made only when a minimum of three of the strongest lines was observed for each compound associated with at least one surface condition out of a series of surface conditions obtained for a given material. Calculations of crystallite size were made from line broadening according to the procedure given in reference 1 on the assumption that crystallite-size effects alone were causing line broadening. Fine copper powder was used to determine the standard line breadth. No estimation of the fraction of this broadening caused by lattice distortion was made. (Lattice distortion may have caused a great part of the broadening but no proved way exists to measure relative effects of lattice distortion and crystallite size.) If, in a group of lines composed of diffraction patterns of several compounds, one pattern obviously predominated in intensity, that compound was believed to be present in greater amounts than any of the others.
Electron diffraction examination of subsurface regions in the used engine specimens was made after each of five successive abrasions of the metal on 00 emery paper under pure benzene. The depth of an abrasion was measured with a vernier micrometer. Estimates of the depth of surface changes were made by microscopic measurements on cross and taper sections. Control specimens cut from the base structure of the engine cylinders were similarly treated and examined for evidence of any oxidation that might have been produced by abrasion.

Hardness measurements were made with either Knoop or Vickers indenters, depending upon the condition of the surface being investigated. The quantitative units thus obtained were checked qualitatively with a Bierbaum-scratch (Microcharacter) hardness tester. This scratch indentation is less affected by metallic strata underlying the surface than the other indenters, but it is more sensitive to surface irregularities.

Chemical and microchemical analyses were made on natural and metallographic surfaces by means of solution, electrographic, and electrolytic etches. Metallographic examinations were made on normal or on tapered and plating-reinforced cross sections.

RESULTS AND DISCUSSION

Steel Surfaces

With the slider-rider combination of SAE 4140 steel and cast iron, several conditions occurred successively in strips, or zones, on the steel slider (fig. 2). The production of more than one surface condition on one specimen was due to slightly imperfect mating of rider and slider. The manner in which surface changes progressed represents various steps in increased loading until failure. Identification of the surface conditions is as follows: A represents the original unworn (unrun) surface; B represents a worn but unsuffed surface; and C represents a surface that appeared to have entered an initial condition of scuff rather early in the experiment but to have almost immediately recovered. Close observation during the run revealed a momentary darkening in color and then a change to dark gray. The permanently scuffed surface that developed in several stages, which succeeded each other within a few seconds, is represented by D. The first stage of formation was indicated by the momentary appearance of a dark-brown or reddish-brown surface color, which changed to black. The change in color was accompanied by the appearance in the recovered oil of a sediment whose major constituent was analyzed as alpha ferric oxide $\alpha$-$\text{Fe}_2\text{O}_3$ by X-ray diffraction methods.
The appearance of bright spots and heavy scuff marks on the surface accompanied by a sharp increase in temperature indicated permanent scuff. It appears that because of the initial imperfect mating a shifting of load took place, which led to variations and irregularities in local load concentrations. Surface B may be considered to have carried a large portion of the load during the mating procedure and therefore to have been able to support the load as it was increased. As the load was increased, surface C suddenly appeared. This appearance occurred as a result, possibly, of slight warping as the temperature rose and because surface C may have been insufficiently mated at the time the load increased. As the load was further increased, strip D appeared, possibly because of a combination of higher load and further warping until the surface scuffed rapidly with so much progressive damage that the scuff became extensive and permanent.

Identification of alpha iron, $\alpha$-Fe, ferrosferric (ferrosic) oxide, Fe$_3$O$_4$, and iron carbide, Fe$_3$C, in these surfaces was made by X-ray diffraction. (See fig. 3.) (Whenever Fe$_3$O$_4$ is mentioned herein, the possibility that it could also be $\gamma$-Fe$_2$O$_3$ should be considered, because the two are indistinguishable by usual diffraction methods.) On all four surfaces (A, B, C, and D), Fe$_3$O$_4$ was considered present and gave an apparent maximum diffraction intensity for scuff-recovered and a decreased intensity for scuffed conditions as judged from the number of lines and from the changes in visual intensity. The identification of Fe$_2$O$_3$ on surfaces B, C, and D was questionable. Iron carbide was identified on the scuffed surface D and questionably identified on scuff-recovered surface C. The intensities were greater for the scuffed than the scuff-recovered surface. No Fe$_3$C was identified on the unscuffed surfaces.

The results of the determination of crystalite sizes are presented in figure 4 and the X-ray patterns showing the differences are shown in figure 5. The hardness of the four surface conditions is also shown in figure 4; maximum hardness existed on the scuff-recovered surface.

Electron diffraction examination of specimens cut from an unscuffed SAE 4140 steel cylinder operated more than 100 hours with cast-iron rings and thoroughly cleaned with solvents on the outermost surface first produced the two diffuse halos characteristic of the Beilby layer and the diffraction pattern of graphite (fig. 3). Light abrasion with 00 emery, in successive steps under benzene, exposed metal that gave the diffraction patterns of $\alpha$-Fe, Fe$_3$O$_4$, and Fe$_2$O$_3$. Oxides were thus detectable to depths estimated near 0.001 inch. Oxidation during the abrasion was proved negligible by the same treatment of control specimens.
This oxidation is shown in more detail in figure 6 and is labeled depth of oxygen penetration. Oxides existed beneath the surface of the metal everywhere the cylinder had been rubbed. The top 1/8 inch of the barrel was subjected to conditions within the combustion chamber but to no rubbing during engine operation because it was above the region of piston contact. In that top area the oxide existed only as a tenacious blue surface film under a carbon deposit. Variation in surface hardness is plotted on the same abscissa as the oxygen data in figure 6. These data were taken at five representative points along the barrel length.

Chrome Surfaces

The chrome-plated cast-iron rider that was reciprocated against the steel slider contained only three distinguishable surface conditions at the end of the run; namely, worn unscuffed, burnished, and normal scuff. Unrun-surface analyses were obtained from another rider.

The chrome pattern predominates in all the surface X-ray diffraction patterns. Diffraction examination showed no presence of any chromic oxide, Cr₂O₃, on the unscuffed surface although this surface was expected to contain at least a small amount of this oxide, possibly in the amorphous condition. The compounds Cr₂O₃ and chromium carbide, Cr₃C₂, were easily identified on the burnished and scuffed surfaces. An X-ray diffraction pattern containing evidence for these compounds on the scuffed surface is shown in figure 7. A number of diffraction lines that stand alone and belong to none of the previously mentioned compounds may be parts of the diffraction patterns of Fe₂O₃, Fe₃O₄, the chromium nitrides; Cr₂N and CrN, the chromium carbides, Cr₂₃C₆, Cr₇C₃, and also Fe₃C (on the scuffed surfaces). No completely reliable quantitative trends could be found because of some difficulty in rating diffraction intensities, but the intensities of Cr₂O₃ and Cr₃C₂ appear to be greater on the scuffed than on the burnished portion. Under the microscope, patches of loose-appearing material with the color and texture of red and black iron oxides and giving dissolution reactions for iron were observed in the surface depressions, apparently as left after pickup from the mating steel slider.

The variations in hardness among the different surface conditions are graphically illustrated in figure 8. Crystallite-size determination showed no large differences. A decrease in crystallite size was expected approaching scuff and such a trend was indicated, but the differences were within experimental error. The sizes were of the order of 14 x 10⁻⁷ centimeters.
Electron-diffraction examinations of unscuffed chrome-plated rings taken from steel aircraft-engine cylinders produced no usable patterns. Only high background was observed. Under the microscope, the surfaces had the appearance of having a high polish. X-ray diffraction of these surfaces also showed considerable diffuse scatter together with very diffuse and faint lines whose approximate spacings were those of \( \text{Cr}_2\text{O}_3 \). Electrographic analyses identified chrome and iron, and a series of reactions by metallographic reagents strongly indicated the presence of the compound \( \text{Cr}_2\text{O}_3 \cdot \text{FeO} \).

Cast-Iron Surfaces

Two cast-iron slider specimens were examined that had been subjected respectively to a normal and to a prolonged run-in procedure. The prolonged run-in increased load-carrying capacity. Figure 7 lists an X-ray diffraction pattern from an unrun surface showing the presence of \( \alpha\)-Fe and one weak ring (interplanar spacing, 4.5 A) that coincides with the strongest line of the graphite pattern. A number of weak lines and one strong line coincide with all the strong lines of \( \text{Fe}_3\text{C} \), and several unidentified lines also show. On all the surfaces, \( \alpha\)-Fe was observed. The presence of graphite was more completely indicated on the worn and scuffed surfaces. Hardness values showed decrease as scuff condition was approached (fig. 9) and crystallite sizes increased.

No examinations were made of used unplated cast-iron piston rings; the information presented in references 2 and 3 is considered sufficient for comparison. These references discuss the detection of the presence of graphite and oxides on surfaces that were in sliding contact during engine operation.

Correlation of Results

In general, the results may be divided into three classes: chemical reaction, material hardening, and alteration of crystallite size. The chemical and physical changes identified on surfaces from the reciprocating slider were superficial compared with the changes wrought on surfaces during engine tests. The depths of the changes were as follows:
On all specimens examined, a certain amount of metallic smear and crystal bending was observed on or just under the surface.

The compounds present on engine-cylinder-barrel surfaces were present also on the slider surfaces of the same material. The only differences in the findings were that the various chemical compounds and phases were more easily identified on the slider specimens, and the cylinder-barrel surfaces definitely showed carbon and the Beilby layer. The diffraction patterns from the surface material on engine specimens often tended to be diffuse.

The strength of the Fe$_3$O$_4$ diffraction pattern from the scuff-recovered steel and the strong evidence of the Fe$_3$C from the scuffed steel indicated that Fe$_3$O$_4$ can be associated with scuff resistance and that Fe$_3$C can be associated with scuff. Caution should be used, however, in basing such a comparison on the evidence in this investigation alone. Evidence for the association of different oxides with different surface conditions may be found in references 4 and 5. A specially prepared Fe$_3$O$_4$ surface was found to have excellent properties partly, at least, because it allowed rapid surface-profile break-in without subsequent high wear or scuffing. This oxide was desirable for an extremely fine and granular structure firmly rooted to the base material. It was not too tenacious to resist abrasion and broke down into a very fine polishing medium during rubbing (reference 4). Whenever in the preparation of the oxide surface the red Fe$_2$O$_3$ formed, unfavorable results were obtained. The red oxide was soft and nonadherent (reference 5).

The reason for this difference in surface-rubbing characteristics of these two oxides is not exactly known. Either oxide may form on a given surface depending upon temperature and pressure relations. The number of Fe$_3$O$_4$ nuclei developed in unit volume in unit time is relatively large and the velocity of crystallization is small and this oxide may occur as an impalpable powder. The red Fe$_2$O$_3$, however, is always crystalline (reference 6, pp. 740, 741, 776).

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Depth of change (in.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reciprocating slider</td>
<td>$1 \times 10^{-5}$ to $1 \times 10^{-4}$</td>
</tr>
<tr>
<td>Aircraft-engine-cylinder</td>
<td>$1 \times 10^{-4}$ to $1 \times 10^{-3}$</td>
</tr>
</tbody>
</table>

*Except that $1 \times 10^{-4}$ in. may be maximum for chrome plate.*
The engine specimens represented run-in and wear conditions entirely different from those imposed on slider surfaces. Mixed hot gases, flame, impact loading of portions of the cylinder surface, and high sliding speeds were considerably different from the room atmosphere, simple frictional heat, slow speed, and steady loading of the slider. Yet both the engine and slider surfaces exhibited the same kinds of change. The long period of operation of the engine, however, affected the depth to which oxygen penetrated.

The occurrence of oxidation in depth only in regions of surface rubbing suggests the occurrence of minute or microscopic voids as faults (or cracks) within the atomic structure or as intergranular voids. (See references 7 to 10.) The density of an uncast metal during working decreases (references 11 and 12), possibly as a result of the propagation of structural faults or the formation of internal voids. Such a decrease should favor penetration and diffusion of other elements, particularly gaseous ones, into the metal. Such relatively deep oxygen penetration is less in agreement with the usual concentration formula, which limits the thickness of an oxide film, than with an activity-gradient formula (reference 13).

The wear profile of engine cylinders showed a depression, or highest wear, at the top of compression-ring travel. In this region of high wear where less oxygen penetration seemed to have occurred, the surface had been worn away almost as fast as the penetration progressed. Farther down the length of the cylinder barrel the rate of oxygen penetration seemed to exceed the rate of wear. As a result, oxides formed within a relatively pliable matrix of steel and possibly gave rise to a desirable surface, combining pliability with decreased abrasion, which was continually renewed under the slow abrasion of operating wear.

Breakdown of lubricant under excessive conditions may account for an amount of carbon sufficient for formation of the carbides. The formation of a carbide-rich phase and the increased surface hardness on boundary-lubricated steel against cast iron is indicated in reference 14.

The crystallite size and the hardness of the wearing surfaces are possibly as important as the chemical reactions. The relations between crystallite size and hardness for steel, chrome, and cast iron vary as shown in figures 4 and 9 and are somewhat complex. The average hardness of the SAE 4140 steel for the scuffed and scuff-recovered surfaces was greater than the average hardness of the unscuffed surfaces, and by similar comparison the crystallite size was smaller. In conjunction with these facts, the formation of the iron oxides and carbide on the scuffed surfaces is noted. If the change in hardness is to be attributed
to one of the other changes, it seems to follow chemical reaction best. The occurrence of Fe$_3$O$_4$ in its greatest quantity coincided with the greatest hardness on the scuff-recovered surface, and a drop in hardness on the scuffed surface coincided with an indicated reduction in the amount of Fe$_3$O$_4$. This concept of a general relation of hardness to chemical reaction may or may not be weakened, however, by the occurrence of Fe$_3$C with the lower hardness of the permanently scuffed surface.

The hardness of the chrome surface increased markedly for the burnished surface condition and corresponded with the formation of Cr$_2$O$_3$ and Cr$_3$C$_2$. The cast iron softened throughout the runs and the crystallite size increased. The end result at scuff was approximately the same for normal and for prolonged run-in. The prolonged-run-in surface, having the higher load at failure, showed slightly less softening than the normal-run-in surface. The more pronounced crystallite-size increase for the worn-uncuffed surface under prolonged run-in seems incompatible with increased resistance to failure. Chemical changes may account for the reversed trend for this material with respect to the over-all trend associated with the other materials, although such changes were undetectable in this case. Relatively, an opposite trend is indicated for any one of these materials when the change in hardness is compared with change in crystallite size between unscuffed and scuffed surface conditions.

**SUMMARY OF RESULTS**

The following observations were made as a result of examinations of metallic surfaces that had undergone rubbing under conditions of partly boundary and partly hydrodynamic lubrication; the changes found on steel, chrome, and cast iron varied with the run-in and scuffed surfaces.

The principal chemical reactions were oxidation and carburization with the following reaction products:
<table>
<thead>
<tr>
<th>Specimen</th>
<th>Surface condition</th>
<th>Material identified on surface</th>
<th>Questionable identification</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAE 4140 steel reciprocating slider</td>
<td>Unrun</td>
<td>α-Fe</td>
<td>Fe$_3$O$_4$</td>
</tr>
<tr>
<td></td>
<td>Worn unscuffed</td>
<td>α-Fe</td>
<td>Fe$_3$O$_4$, Fe$_2$O$_3$</td>
</tr>
<tr>
<td></td>
<td>Scuffed</td>
<td>α-Fe, Fe$_3$O$_4$</td>
<td>Fe$_3$C, Fe$_2$O$_3$</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fe$_2$O$_3$</td>
</tr>
<tr>
<td>SAE 4140 steel aircraft-engine cylinder barrel</td>
<td>Worn unscuffed (Superficial face)</td>
<td>C (graphite), Beilby layer</td>
<td>Mixed oxides</td>
</tr>
<tr>
<td></td>
<td>(Less than 0.001 in. below surface face)</td>
<td>α-Fe, Fe$_3$O$_4$, Fe$_2$O$_3$</td>
<td></td>
</tr>
<tr>
<td>Chrome-plated cast-iron rider</td>
<td>Worn unscuffed</td>
<td>α-Cr</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Burnished</td>
<td>α-Cr, Cr$_2$O$_3$, Cr$_3$O$_2$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Scuffed</td>
<td>α-Cr, Cr$_2$O$_3$, Cr$_3$O$_2$</td>
<td></td>
</tr>
<tr>
<td>Cast-iron reciprocating slider</td>
<td>Unrun</td>
<td>α-Fe</td>
<td>Fe$_3$C</td>
</tr>
<tr>
<td></td>
<td>Worn</td>
<td>α-Fe</td>
<td>C (graphite)</td>
</tr>
<tr>
<td></td>
<td>Scuffed</td>
<td>α-Fe</td>
<td>C (graphite)</td>
</tr>
</tbody>
</table>

1. Lines best fit pattern of material listed.

Hardness of materials examined varied with degree of surface condition. The crystallite sizes were in the order of $10^{-7}$ centimeters. The sizes changed with condition of operation.

Flight Propulsion Research Laboratory,
National Advisory Committee for Aeronautics,
Cleveland, Ohio, April 22, 1947.

REFERENCES


5. Roensch, Max M.: Piston-Ring Coatings and Their Effect on Ring and Bore Wear. SAE Jour. (Trans.), vol. 46, no. 5, May 1940, pp. 221-228.


BIBLIOGRAPHY


Figure 1. - Reciprocating slider machine.
Surface
A Unworn
B Worn, unscuffed
C Scuff recovered
D Permanently scuffed

Figure 2. - Run-in and scuffed SAE 4140 steel slider and cast-iron rider showing types of surface on slider.
Scuff-recovered SAE 4140 steel slider specimen (X-ray diffraction)

α-Fe, body-centered-cubic structure (standard pattern 1)

Fe₃O₄, spinel-type structure (standard pattern 1)

Fe₃C, orthorhombic structure (standard pattern 2)

α-Fe₂O₃, hexagonal structure (standard pattern 1)

Well run-in SAE 4140 steel aircraft-engine cylinder-barrel surface (electron diffraction)

C (graphite), hexagonal structure (standard pattern 1)

Beilby layer on same barrel surface (electron diffraction)

1Taken from A.S.T.M. card index for X-ray diffraction.
2Taken from reference 15.

Figure 3. - Typical electron and X-ray diffraction patterns from surfaces of specimens of SAE 4140 steel slider and aircraft-engine cylinder barrel and standard X-ray diffraction patterns of materials identified.
Figure 4. - Relation between surface changes and surface condition of SAE 4140 steel slider specimens.
Figure 5. - X-ray diffraction patterns from SAE 4140 steel slider specimen showing change in crystallite size from surfaces A to D.
Figure 6. - Relations among surface condition, material hardness, and depth of oxygen penetration found on worn - unscuffed SAE 4140 steel aircraft-engine cylinder barrel.
Scuffed chrome-plated cast-iron rider specimen (X-ray diffraction)

Unworn cast-iron rider specimen (X-ray diffraction)

α-Cr, body-centered-cubic structure (standard pattern¹)

α-Fe, body-centered-cubic structure (standard pattern¹)

Cr₂O₃, hexagonal structure (standard pattern¹)

Cr₃C₂, orthorhombic structure (standard pattern¹)

Fe₃C, orthorhombic structure (standard pattern²)

C (graphite), hexagonal structure (standard pattern¹)

¹Taken from A.S.T.M. card index for X-ray diffraction.
²Taken from reference 15.

Figure 7. - Typical electron and X-ray diffraction patterns from specimens of chrome-plated cast-iron and cast-iron riders and standard X-ray diffraction patterns of materials identified.
Figure 8. - Relation between hardness and surface condition of chrome-plated cast-iron rider specimen.

Figure 9. - Relation between surface changes and surface condition of cast-iron slider specimens.
Study was made of run-in and scuffed steel, chrome plate, and cast-iron surfaces. X-ray and electron diffraction techniques, microhardness determinations and microscopy were used. Surface changes were found to include three classes: chemical reaction, hardening, and crystallite-size alteration. Principle chemical reactions were oxidation and carburization.

NOTE: Requests for copies of this report must be addressed to: N.A.C.A., Washington, D.C.