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OXFORD.

1965.

Best Available Copy
FINAL REPORT
March 1965.

"The Reaction of Atoms and Radicals with Various Molecules"

The research supported in this document has been sponsored in whole by:

AIR FORCE OFFICE OF SCIENTIFIC RESEARCH, OAR

through the European Office of Aerospace Research,
United States Air Force.

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INTRODUCTION.

This report is divided into four chapters:

Chapter I. deals with some experiments on the Absorption of hydrogen atoms by palladium, and palladium-gold alloys. This work was carried out in connection with some earlier work on the catalytic effect of these metals and alloys on the recombination of hydrogen atoms.

Chapter II. deals with some experiments on the attack of a number of metals by molecular and atomic oxygen to discover in which cases there were considerable differences in rate.

Chapter III. deals with some experiments on the recombination of oxygen atoms on the surface of the alkali metal tungsten bronzes.

Chapter IV. describes a pressure gauge which has been devised for the rapid and convenient measurement of pressures in mixtures containing chlorine at pressures of the order of 1 mm, the accuracy being about ± 0.01 mm. The experiments on the recombination of chlorine atoms have not yet reached a stage at which a report is worth-while.

In addition, during the period of this grant, work was carried out on the reactions of atoms at metal surfaces. In these experiments oxygen atoms effused through a small hole opposite which was a small disc coated with silver, copper, chromium etc. A complete report of this work was given in the report submitted in December 1963 (pages 1 - 14 of that report). Consequently no further description of that work is included here.
CHAPTER 1.


Introduction, by D. M. Hirst and J. W. Linnett.

Some time ago we made some measurements on the recombination of hydrogen atoms at the surface of palladium, gold and palladium-gold alloys. This work is about to be published in a paper entitled "The Recombination of Atoms on Pd-Au Alloys" by P. G. Dickens, J. W. Linnett and W. Palczewskak in the Journal of Catalysis (The work was supported by the Office of Aerospace Research of the U.S. Air Force: Grant: AF EOR 62-6). It was found that palladium was more active than gold in catalysing the recombination, that small percentages of gold in the palladium did not reduce the initial activity but that larger amounts did. However, another interesting result was that the activity of palladium, and the palladium rich alloys, decreased with time as the foil was exposed to atomic hydrogen. It was concluded that the presence of vacancies in the metallic d-band enhanced the catalytic activity of the materials, but that these vacancies could be filled either by alloying gold with the palladium, or when atomic hydrogen was absorbed by the metal or alloy. That is, when atomic hydrogen was absorbed, its electron was contributed to the completion of the d-band with a consequent fall off in the activity. This hypothesis demands that palladium and its alloys should absorb hydrogen when exposed to hydrogen containing atomic hydrogen under the conditions in which the catalytic experiments were carried out (i.e. hydrogen pressure about 10^{-2} mm.; percentage of atoms about 10%; room temperature). This was the reason why these experiments were carried out. Samples of the metals and alloys
were suspended from a micro-balance and then exposed to hydrogen containing atoms. The increase in weight with time (if any) was followed.

Brief Survey of Related Work.

Palladium is known to absorb large quantities of hydrogen. If it is exposed to hydrogen at about 2 atmospheres pressure it will take up about 0.6 gm. atoms of hydrogen for every gram atom of palladium.

The hydrogen diffuses as atoms into the palladium lattice. The initial solution produces what is known as the d-phase. This is accompanied by a gradual expansion of the face-centred cubic palladium lattice spacing from about 3.883 to 3.894 A at a concentration of about 0.2 gm. atoms of H per gm. atom of palladium. At this concentration the lattice suddenly expands to 4.048 A with the formation of the j-phase which is also face centred cubic. The lattice spacing remains constant for concentrations between 0.2 and 0.7 or 0.8 gm. atoms of hydrogen per gm. atom of palladium but at higher concentrations it increases again. Concentrations higher than 0.9 gm. atoms of H per gm. atom of palladium do not seem to have been attained. The j-phase can be supersaturated with hydrogen by electrolytic charging, the excess hydrogen being lost when the charging process ceases.

The magnetic susceptibility of palladium decreases linearly with the hydrogen concentration becoming zero when there are about 0.66 gm. atoms of H per gm. atom of palladium. Palladium atoms have 10 electrons beyond the closed core consisting of the 1s, 2s, 2p, 3s, 3p, 3d, 4s and 4p orbitals. The 4d and 5s atomic orbitals produce, in the metal, a broad s-band and a narrow d-band which can accommodate 2 and 10 electrons.
respectively. The electrons occupy levels in the two bands up to the same energy level. There are, per atom, 9.4 electrons in the d-band and 0.6 in the s-band. Thus there are about 0.6 holes in the narrow d-band. The electrons of the absorbed hydrogen atoms are presumed to enter these holes in the d-band. Norberg (Phys. Rev., 1952, 86, 745) showed, using nuclear magnetic resonance spectroscopy, that the hydrogen is present in the lattice in the form of protons. The magnetic susceptibility drops to zero when the holes in the d-band are filled.

The kinetics of the absorption of hydrogen by palladium from ordinary hydrogen gas has been investigated by Holt (Proc. Roy. Soc., 1914, 90A, 226), Tammann and Schneider (Z. anorg. Chem. 1928, 172, 43), Smith and Derge (J.A.C.S., 1934, 56, 2313) and Kruase and Kahlenberg (Trans. Electrochem. Soc., 1935, 68, 449). Most of the experiments showed that there was a linear rate of uptake initially which was followed by a falling off in the rate as the amount absorbed approached a limiting value. The rate, and the amount taken up were very dependent on the previous treatment.

Palladium and gold produce a continuous series of solid solutions. The structure is face-centred cubic and there is no evidence of a super-lattice. The spacing varies almost exactly in a linear manner with concentration. Vogt (Ann Physik, 1932, 14, 1) found that the magnetic susceptibility decreases as the proportion of gold increases becoming zero at about 55 atomic per cent of gold.

Couper and Eley (Disc. Farady Soc., 1950, 8, 172) studied the ortho-para hydrogen conversion on palladium-gold alloys and obtained results similar in form to those of Dickens, Linnett and Palczewski.
for the r. combination of hydrogen atoms. Couper and Eley also observed that hydrogen exerted a poisoning effect on the reaction they studied.

Schneiderman (Ann. Physik., 1932, 13, 761) investigated the solubility of hydrogen in palladium-gold alloys. The alloys were charged electrolytically. He showed that the solubility decreased as the gold content increased.

**Experimental Method.**

The sample of metal or alloy weighing about 1/2 gm. was suspended in a vertical tube of about 4 cm. in diameter from one arm of a micro-balance capable of measuring a change in weight of $10^{-6}$ gms. The vertical tube was joined at its lower end to a horizontal tube along which hydrogen at $1/10$ mm pressure was flowing. This hydrogen had been passed through an electric discharge and so contained atoms. These diffused up the vertical tube to the sample. By this means therefore a steady condition could be maintained indefinitely. That is the flow rate, pressure and proportion of atoms at the lower end of the tube containing the sample was constant for as long as was needed for the experiment. Moreover the apparatus could be left operating unattended.

In order that there should be no contamination by electrode materials, an electrodeless discharge was used. This was maintained by 13½ Mc/s radiation from an R.C.A. radio frequency oscillator (type ET-4336-B), a helix of 3.002" copper tubing being wrapped round the tube through which the hydrogen was flowing. In the discharge region there will be present ions, electrons, atoms and molecules some of which will be in
excited states. The charged species will decay rapidly and will not reach the test region. Because the first excited state of hydrogen atom is more than 10 eV above the ground state and the first excited state of the hydrogen molecules is purely repulsive, it is extremely likely that the gas in the test region will consist of a mixture of atoms and molecules in their ground states.

The hydrogen was prepared by the electrolysis of a 15% solution of sodium hydroxide saturated with barium hydroxide to prevent the evolution of any carbon dioxide. A current of 5 amps from the 100 volt D.C. supply was used and the cell was water-cooled. Any oxygen was removed from the hydrogen by passing through a tube containing palladised asbestos at 400°C. The gas could be dried by passing through a trap cooled in liquid oxygen and then through a column of phosphorus pentoxide. If the gas was needed moist it was passed through a humidifier containing water at a controlled temperature. The flow of the hydrogen was measured by determining the pressure drop across a tube containing a plug of sintered glass wool. The flow of gas from the high pressure (atmospheric) region into the low pressure part was controlled by an Edwards fine control needle valve LBIA. The flow meter, saturator and needle valve were immersed in a water bath maintained at 23°C by a Tecam Tempunit.

The pumping system consisted of a large high speed single stage mercury pump backed by a smaller two stage mercury pump. These were backed by an Edwards "Speedivac" IS 150 rotary oil pump. The pumping line was made of 4 cm. diameter tubing and contained two large volumes to act as flow stabilisers. A block diagram of the apparatus is shown.
in Figure I.1.

The balance used was a Sartorius vacuum Electrono-microbalance. It could operate on seven ranges of sensitivity of 1, 2, 5, 10, 20, 50 and 100 µg per scale division. The results could also be recorded on a Leeds and Northrup "Speedomax H S" recorder. The micro-balance was mounted on a bracket of ½" steel measuring 18" by 6" which was fixed into the wall, 6" being in the wall. A piece of slate, supported on two strips of lead was placed on the bracket. A sheet of fairly hard rubber ½" thick was stuck onto the slate and the micro-balance placed on top of this. The sample was supported by a silica hook which was attached to a silica suspension fibre which was in turn hung from the steel suspension wire on one arm of the balance. A counterweight of approximately the same weight as the sample was hung from the other arm.

The sample was situated in a vertical tube which was 4 cm. in diameter. This tube was of silica and the vertical section was 45 cm. long. It was connected to the microbalance through a flange. This vertical tube was made into a furnace so that the sample could be heated up to 800°C. The temperature was measured with a platinum/platinum-rhodium thermocouple. Additional thermocouples were contained in the tube to measure the temperature gradient in the region of the sample.

The gas pressure was measured with a McLeod gauge with a compression ratio of 1000:1. The atom concentration in the neighbourhood of the sample could be measured with a Wrede-Harteck gauge fused into the wall of the silica tube. The pressures inside and outside the gauge were
measured with micro-Pirani pressure gauges which we constructed.

**Experimental Results.**

Samples of foil, 2/1000" thick, containing 0, 12, 31, 45, 72 and 100 atomic per cent of gold were used. The alloys were kindly loaned by the Mond Nickel Company.

After a number of rather unsatisfactory experiments with dry gas it was decided to use wet gas. This increases very greatly the production of hydrogen atoms. If any oxygen atoms are formed they will be converted rapidly to OH and H by the reaction

\[
O + H_2 = H + OH
\]

and any hydroxyl radicals to hydrogen atoms by

\[
OH + H_2 = H_2O + H.
\]

Consequently at the test sample, the material will be exposed to hydrogen molecules and atoms together with a small proportion of water molecules. Experiments were carried out on pure gold with dry gas but on all the other samples with wet gas. The samples weighed about \(\frac{1}{2}\) gm. and measured 2½ or 3 cms. by 2½ or 3 cms.

Figure 1.2 shows the results for palladium, the increase in weight being plotted against time. In this case the measurements of atom concentration were unreliable but in the 2nd and 5th runs the calculated percentages were 6½ and 5½ respectively. Table 1 gives the details of the experiment together with the limiting up-take of hydrogen measured in parts per million (by weight). The results for the palladium-gold alloys are shown in Figures 1, 3, 4, 5, 6 and 7 and similar details are included in Table 1. The results for gold are shown in Figure 1.7.

Table II lists the limiting amounts of hydrogen taken up by the
different metals and alloys expressed as gm. atoms of hydrogen \( x \) per mole of alloy, \( \text{Pd}_x\text{Au}_{(1-x)} \). These are plotted against the atomic percentages of gold (i.e. \( 100(1-y)% \)) in the alloy in Figure 1.8.

When the discharge was switched off at the end of a hydrogen-absorption experiment, there was usually a steady decrease in weight. The hydrogen starts to desorb as soon as the source of atoms is removed. Figure 1.9 shows the loss in weight against time for three experiments with palladium. With the alloys the desorption took place in a similar way but the rates of desorption were rather variable from experiment to experiment (initial rates of loss were 40 to 150 gms. per hour).

Under these conditions only a part of the hydrogen was lost. However, if the sample was heated, the rest of the hydrogen could be driven off. When the palladium sample was heated up, the loss started at about 90\(^\circ\) or 100\(^\circ\)C and took about a quarter or half an hour. With the samples containing 88\% and 69\% of palladium the results were essentially similar, but with the sample containing 55\% palladium, though the loss commenced at 90\(^\circ\) it took place rather more slowly.

Discussion.

The most striking feature of these experiments is the enormous absorption of hydrogen which occurs when palladium and palladium-gold alloys containing more than 40 atomic per cent of palladium are exposed to partially dissociated hydrogen at pressures of the order of 0.1 mm. Hg. The absorption from molecular hydrogen is negligible at such pressures.

Because the results are variable it is not possible to reach any very definite conclusions regarding the rate of uptake of hydrogen by the different samples. However there are two rather general comments that can
be made. W. Jost (Diffusion in solids, liquids and gases, Academic Press Inc., New York, 1952) has shown that if the uptake is diffusion controlled then it is to be expected that the function \( \log_{10} \frac{c - c_f}{c_i - c_f} \) should vary linearly with the time, where \( c_i \) and \( c_f \) are the initial and final concentrations of absorbed material and \( \bar{c} \) is the average concentration at time \( t \). It was found that this is true for the initial uptake of the hydrogen. This is shown for two experiments with the palladium-gold alloy containing 88% palladium in Figure 1.10. Here it is seen that the points lie on a straight line for about the first five hours. Similar results were obtained for other alloys. However, for the later stages, this relation did not hold. It was found that the results were then treated best in terms of diffusion through a film of increasing thickness (analogous to the treatment of the formation of oxides when a coherent film is formed). Under these conditions the square of the weight increase is a linear function of the time. That this holds, at any rate approximately, for two experiments with the same alloy is shown in Figure I.11.

If this is a correct interpretation of the data, then one reaches the conclusion that the uptake of atomic hydrogen must be regarded in the following way. To begin with the hydrogen is taken up and diffused into the metal or alloy, the part containing the hydrogen being a uniform phase together with the metal. Thus the uptake is controlled by a simple diffusion process. However, as more is taken up, the outside layer which now contains a lot of hydrogen becomes a new phase and the build up of this phase from the inner part of the sample is controlled by diffusion of hydrogen atoms through this layer which becomes thicker and thicker with time, the thickness being proportional approximately to the weight.
increase. This may be related to the phase changes in the solid, though if these were to be identified simply as the $\alpha$ and $\gamma$ phases one would have expected the change-over from simple diffusional behaviour to occur at a lower hydrogen content.

It was shown that when an alloy was saturated with hydrogen atoms and the source of hydrogen atoms then switched off some hydrogen was lost. It is shown in Figure 1.12 that if $\log \frac{c_i}{c_f}$ is plotted against time for some results for an alloy containing 69% palladium a straight line is obtained. This shows that the process is a diffusional one. It is difficult to see how this is to be linked with the results for the up-take of atomic hydrogen though one must bear in mind that, in the desorption experiment, molecular hydrogen is being lost from the surface.

The results in Figure 6 show that the total amount of hydrogen absorbed decreases as gold is added. It can be concluded that when hydrogen is absorbed the electrons enter the d-band. Also when gold is added the additional electron, beyond the closed shells, enters the d-band - consequently as gold is added the number of vacancies in the d-band decreases and consequently the ability to absorb hydrogen atoms decreases.

It appears from Figure 6 that when 55 atomic per cent of gold has been added the d-band is full and no more hydrogen can be absorbed. This agrees fairly well with the results quoted in the Introduction, namely that, in palladium there are 0.6 holes per atom in the d-band.

These results fall into line with those obtained by Dickens, Linnett and Palczewski on the poisoning effect of both gold and atomic hydrogen on the activity of palladium as a catalyst for the recombination of hydrogen atoms. It seems that high activity is to be associated with
vacancies in the d-band as has been proposed. However the alloy
containing 72 atomic per cent of gold does show an activity markedly
greater than that of gold which would not be expected on the simple
arguments given above. It may be that the electronic conditions at the
surface are rather different from those in the bulk and that there are
d-shell vacancies for the surface atoms when there are essentially no
d-band vacancies. Nevertheless, broadly speaking, the present results
confirm the interpretation of the earlier catalytic results.
Block diagram of apparatus.
$\mu g^2$ v time for Pd88Au12.
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<td></td>
<td></td>
<td></td>
<td>22½-23½</td>
<td>156.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

Palladium
<table>
<thead>
<tr>
<th>Run</th>
<th>Annealing (°C)</th>
<th>Cooling (hrs)</th>
<th>Exposure to H (hrs)</th>
<th>Pressure of H₂ p.p.m.</th>
<th>Absorbed</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>800, 1½ hrs</td>
<td>2</td>
<td>22</td>
<td>3/4 - 4 Discharge failure</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>4 - 8 135.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>29 - 29⁴ 139</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>810, 1½ hrs</td>
<td>2</td>
<td>47⁴</td>
<td>9 - 10 139.7</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>24⁴ - 26⁴ 140.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>46⁴ - 47⁴ 138.7</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>810, 4 hrs</td>
<td>2½</td>
<td>23</td>
<td>23 - 9 139.5</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>22 140 140</td>
<td>0</td>
</tr>
</tbody>
</table>
### TABLE I.II

**Solubilities of hydrogen in palladium - gold alloys**

<table>
<thead>
<tr>
<th>Alloy</th>
<th>M.Wt.</th>
<th>Run</th>
<th>g H/gm. (p.p.m.)</th>
<th>g atoms H/gm. mol. alloy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pd 1</td>
<td>106.4</td>
<td>1</td>
<td>7189</td>
<td>0.764</td>
</tr>
<tr>
<td>Pd 2</td>
<td></td>
<td>2</td>
<td>6681</td>
<td>0.711</td>
</tr>
<tr>
<td>Pd 3</td>
<td></td>
<td>3</td>
<td>6686</td>
<td>0.731</td>
</tr>
<tr>
<td>Pd 4</td>
<td></td>
<td>4</td>
<td>6640</td>
<td>0.706</td>
</tr>
<tr>
<td>Pd 5</td>
<td></td>
<td>5</td>
<td>7017</td>
<td>0.747</td>
</tr>
<tr>
<td>Pd 6</td>
<td></td>
<td>6</td>
<td>6822</td>
<td>0.726</td>
</tr>
<tr>
<td>Pd 89</td>
<td>117.27</td>
<td>1</td>
<td>3952</td>
<td>0.463</td>
</tr>
<tr>
<td>Au 12</td>
<td></td>
<td>2</td>
<td>3400</td>
<td>0.399</td>
</tr>
<tr>
<td>Au 51</td>
<td>134.49</td>
<td>3</td>
<td>2923</td>
<td>0.343</td>
</tr>
<tr>
<td>Pd 69</td>
<td></td>
<td>1</td>
<td>1828</td>
<td>0.246</td>
</tr>
<tr>
<td>Au 31</td>
<td></td>
<td>2</td>
<td>1746</td>
<td>0.235</td>
</tr>
<tr>
<td>Au 55</td>
<td>147.17</td>
<td>3</td>
<td>1750</td>
<td>0.235</td>
</tr>
<tr>
<td>Pd 55</td>
<td></td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Au 45</td>
<td></td>
<td>2</td>
<td>647</td>
<td>0.095</td>
</tr>
<tr>
<td>Au 72</td>
<td>171.63</td>
<td>3</td>
<td>718</td>
<td>0.106</td>
</tr>
<tr>
<td>Au 28</td>
<td></td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Au 28</td>
<td></td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Au 28</td>
<td></td>
<td>3</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Au 197</td>
<td></td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Au 197</td>
<td></td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Au 197</td>
<td></td>
<td>3</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
CHAPTER II.

The Oxidation of Metals by Atomic and Molecular Oxygen.

Introduction by R. Heckingbottom and J. W. Linnett.

The present work concerns a study of the oxidation of metals by oxygen atoms and molecules. An R.F. discharge has been used for producing the atoms and the uptake of oxygen by the metals has been determined gravimetrically. The main object has been to determine for which metals the rate and extent of reaction with oxygen atoms differed markedly from that with molecular oxygen. In view of the small amount of previous work on the subject the emphasis has been on conducting a survey of a wide variety of metals, in the hope of detecting some pattern of behaviour.

The oxidation of metals has been, for many years of great interest to the chemist, both for its obvious practical importance and because it has been a fruitful field for testing many ideas based on solid state theory. However, though the effect on oxidation rate due to variation of either the temperature of the system or the composition of both metallic and gaseous phases, has often been studied in detail, that due to dissociation of the attacking gas has received relatively little attention.

It was felt that further work in this latter field was desirable for two main reasons. Firstly where work of this kind has been attempted the effect of atoms has sometimes been shown to be considerable. Secondly as interest in the upper atmosphere - a region where a relatively high proportion of molecules are dissociated or excited - increases, the subject grows in technological importance.

Previous work with dissociated or activated oxygen.

The most marked contrast in behaviour, towards oxygen atoms on the one hand and molecules on the other, is shown by silver.
In a recent detailed survey, Czanderna (J. Phys. Chem. 68, 2765, (1964)) has shown that between -77°C and 351°C oxygen is chemisorbed on silver, probably as $O^-$, $O_2^-$ and $O_2$, but that coverage never exceeds a monolayer. This confirms earlier work and the oxidation of silver by molecular oxygen has only once been reported (Menzel and Menzel-Kopp, Surface Science 2, 376, (1964)), when crystallites of $Ag_2O$ were observed on silver at 250°C under 35 atmospheres of oxygen.

In contrast, several workers have observed the oxidation of silver by oxygen atoms even at room temperature (e.g. Linnett and Karsden, Fifth Int. Symp. Combustion, 685, (1955)). Silver probes have in fact been used to give a measure of atom concentration (Jennings Quart Rev. 15, 237, (1961)), these probes being quickly blackened due to oxide formation. However, there has been only one previous investigation of the kinetics of the reaction (Tyapkina and Dankow, Doklady Akad. Nauk. S.S.S.R. 52, 1313 (1948)). Even here observations were limited to the first twenty minutes of reaction; the uptake was found to follow the parabolic rate law approximately.

Several workers have found that the presence of oxygen atoms increases the rate of oxidation if the oxide formed is volatile. Rossner and Allendorf (J. Chem. Phys. 40, 3-41, (1964)) have found that between 800°C and 1500°C atoms increase the rate of oxidation of molybdenum by a factor of about twenty - $MoO_3$, the main product, is volatile at these temperatures. Similarly, Pryburg (J. Chem. Phys. 24, 175, (1956)) has shown that platinum is oxidized to volatile PtO$_2$ at 1,000°C and that oxygen atoms are up to four hundred times as effective as molecules. Further, Sutcliffe (D.Phil. Thesis, Oxford (1964)) reports
that atoms greatly increase the oxidation rate of chromium above 150°C due to the formation of CrO$_3$ which is volatile above this temperature.

According to Dravnieks (J. Amer. Chem. Soc., 72, 3761, (1950)), the effect of oxygen atoms on the oxidation of copper is complex due largely to the fact both Cu$_2$O and CuO may be present in the oxide scale. Atoms tend to increase the rate of oxidation during the initial period when only Cu$_2$O is formed and also to shorten this period. When both oxides were present, atoms had little effect.

A little oxidation work has also been carried out using oxygen activated by a tesla coil, notably by Leibowitz et al. (J. Electrochem. Soc. 108, 1166, (1961)) who found that the rate of oxidation of uranium to the non-protective UO$_2$ was increased by up to twenty fold on switching on the coil.

In addition several workers e.g. Hauße and Engel (Z. Electrochem. 57, 773 (1953)) and Dickens and Sutcliffe (Trans. Faraday Soc., 60, 1272 (1964)), have shown that the presence of activated or dissociated oxygen increases both the amount of chemisorbed oxygen and the conductivity of several oxides. Thus in the case of a growing oxide these changes in the surface concentration may well alter the concentration gradients in the bulk and hence influence the oxidation rate. Previous to the present work however there appears to have been no attempt to relate the effects of atoms to the general oxidation theory proposed by Wagner.

**Experimental.**

The experimental method may be outlined as follows. A controlled supply of oxygen flows first through an RF discharge, resulting in partial dissociation of the gas, and then past the open end of a vertical
side arm. The latter is closed at its upper end so that there is no net flow along it; atoms diffuse into the tube under a concentration gradient. The sample, suspended from the arm of a microbalance, is situated in the vertical side arm and its oxidation is followed gravimetrically. In addition the side arm could be heated so that oxidations could be studied at any temperature between 25°C and 800°C. This appears to be the first apparatus to be used that combines the continuous production of atoms with the direct gravimetric investigation of oxidation.

The oxygen was produced electrolytically and used in its undried state as the presence of about 2% water vapour increases the degree of dissociation of oxygen in the RF discharge from about 2% to 10%. The operating pressure was about 0.0 microns, the gas flow being controlled by a thermostated needle valve. Pirani gauges were used for pressure measurement at normal operating pressures, and pressure of around $10^{-5}$ mm used in outgassing procedures were recorded on a Penning gauge.

The sample was suspended by a quartz fibre from one arm of a Sartorius microbalance. This balance was capable of detecting changes of ±1 μg or greater and total weight changes up to 104 μg could be followed. Samples weighing up to 1 g could be used, the greater part of the load being offset by counterweights.

The vertical diffusion arm was heated by a nichrome wire furnace wound onto its outside, and the sample temperature during runs was measured by a thermocouple on the furnace wall, previously calibrated against a second thermocouple fixed in the normal sample position. The temperature could be maintained within ±2°C of any desired value between 25°C and 800°C. Blanketing and a cooling fan were introduced to protect the balance head from
the furnace region; the discharge tube was also fan cooled.

Atom concentrations were determined with silver tipped thermocouple probes and thus only a relative measure was obtained. However comparison with earlier work on similar systems, using both probes and Wrede gauges (Linnett and Greaves, Trans. Faraday Soc. 52, 1323, (1958)), suggested that the atom concentration in the sample region was equivalent to a pressure of 6 μ at room temperature. Unfortunately at higher temperatures the silica walls of the surface became much more active towards atom recombination and the atom concentration in the sample region may fall to as low as 10⁻⁵ μ by 750°C.

One probe was placed in a small side arm just downstream of the discharge tube, to monitor the atom concentration during runs. A second similar probe, tipped with the metal under investigation could also be introduced, by means of a side arm in the furnace, into the normal (but vacant) sample position. The purpose of this was twofold; first the probe was used to check the relative atom concentrations under various conditions and second, as the sample temperature rises in the presence of atoms just as that of a probe does, the magnitude of this rise is required for each metal. Thus in that the sample temperature would be the same for runs with and without the discharge on, the furnace wall temperature was set at an appropriately lower temperature in the latter case.

Although this corrects for the the temperature effect due to atoms over the greater part of a run, it still leaves the problem of the initial temperature rise on switching the discharge on. The effect is small at any 500°C as the total change is only about 3°C, but at 150°C
the temperature rise is about 15°C occurring over the first five minutes of the run, and this must be borne in mind when analysing the results quantitatively.

One further correction was necessary, as the thermomolecular effects were large at the operating pressure. Fortunately if the sample was brought to temperature under a vacuum of $10^{-5}$ mm or better and the oxygen was then introduced the thermomolecular effect registered immediately and hence could readily determined.

The metal samples were normally prepared as follows:— a) abrasion with a fine grade emery paper, b) rinsing with carbon tetrachloride c) drying in a current of warm air and d) outgassing at 700°C and $10^{-5}$ mm. for 30 minutes. In the case of zirconium, process a) was replaced by a chemical polish in a solution containing 45% H$_2$O, 45% conc HNO$_3$ and 10% HF. Zinc and cadmium could not be outgassed as in d) due to their high vapour pressures, this stage was also omitted in the case of tin foil oxidation due to its low melting point.

Results.

The fourteen metals—nickel, cobalt, iron, silver, copper, zirconium, aluminium, tantalum, tin, zinc, cadmium, palladium, platinum and gold, have been studied. The main results will now be given metal by metal.

**Nickel.**

All the oxidations showed a parabolic time dependence and the presence of atoms increased the rate constant considerably in all cases as shown in Table 1.
TABLE 1.

The Oxidation of Nickel.

<table>
<thead>
<tr>
<th>Run</th>
<th>Temperature °C</th>
<th>Gas phase species</th>
<th>k ( \text{cm}^2/\text{cm}^2/\text{sec} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>428</td>
<td>( \text{O}_2 )</td>
<td>4.53 ( \times 10^{-16} )</td>
</tr>
<tr>
<td>1b</td>
<td>430</td>
<td>( \text{O}_2 ) + ( \text{N}_2 )</td>
<td>4.53 ( \times 10^{-14} )</td>
</tr>
<tr>
<td>1c</td>
<td>545</td>
<td>( \text{O}_2 )</td>
<td>1.21 ( \times 10^{-13} )</td>
</tr>
<tr>
<td>1d</td>
<td>545</td>
<td>( \text{O}_2 ) + ( \text{N}_2 )</td>
<td>7.28 ( \times 10^{-13} )</td>
</tr>
<tr>
<td>2</td>
<td>490</td>
<td>( \text{O}_2 )</td>
<td>2.45 ( \times 10^{-13} )</td>
</tr>
<tr>
<td>4</td>
<td>490</td>
<td>( \text{O}_2 ) + ( \text{N}_2 )</td>
<td>2.13 ( \times 10^{-13} )</td>
</tr>
<tr>
<td>3</td>
<td>648</td>
<td>( \text{O}_2 )</td>
<td>3.08 ( \times 10^{-13} )</td>
</tr>
<tr>
<td>5</td>
<td>650</td>
<td>( \text{O}_2 ) + ( \text{N}_2 )</td>
<td>4.38 ( \times 10^{-12} )</td>
</tr>
</tbody>
</table>

The results of runs 1a, 1b, 1c and 1d are shown in Fig. II.1.

The ratio \( k_p (\text{atoms})/k_p (\text{molecules}) \) was about 10 at 600°C and somewhat higher at lower temperatures, the activation for oxidation being 38 kcal/mole in the presence of molecules and 33 kcal/mole in the presence of atoms; only four points were available for determining each figure however. The grey black scale was considered to be NiO.

Cobalt.

The oxidations again followed a parabolic time law. At lower temperatures (below about 660°C) atoms caused only a slight increase in the rate but at higher temperatures (about 700°C) a marked increase resulted, the ratio \( k_p (\text{atoms})/k_p (\text{molecules}) \) being about 6. This is borne out by the results in Table 2 and confirmed in a qualitative manner by the results of successive runs on sample 1 as shown in Fig. II.2.
TABLE 2.
The Oxidation of Cobalt.

<table>
<thead>
<tr>
<th>Run</th>
<th>Temperature °C</th>
<th>Gas phase species</th>
<th>$k_p$ gm$^2$/cm$^4$/sec</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>564</td>
<td>$O_2$</td>
<td>$8.46 \times 10^{-12}$</td>
</tr>
<tr>
<td>3</td>
<td>563</td>
<td>$O + O_2$</td>
<td>$1.00 \times 10^{-11}$</td>
</tr>
<tr>
<td>4</td>
<td>713</td>
<td>$O_2$</td>
<td>$1.02 \times 10^{-11}$</td>
</tr>
<tr>
<td>5</td>
<td>713</td>
<td>$O + O_2$</td>
<td>$6.12 \times 10^{-11}$</td>
</tr>
<tr>
<td>6e</td>
<td>713</td>
<td>$O_2$</td>
<td>$1.23 \times 10^{-11}$</td>
</tr>
<tr>
<td>1f</td>
<td>714</td>
<td>$O + O_2$</td>
<td>$7.21 \times 10^{-11}$</td>
</tr>
</tbody>
</table>

At the lower temperature (runs 2 and 3) both CoO and Co$_3$O$_4$ were present in the scale, at the higher temperature only CoO was found. Investigation of the thermal decomposition of the scale formed in runs 2 and 3 showed that the Co$_3$O$_4$ content was about 31% by volume in each case.

Iron

When single runs were carried out on samples the parabolic rate law was followed and $p$-ions had no detectable effect, as shown in Table 3.

TABLE 3:

<table>
<thead>
<tr>
<th>Run</th>
<th>Temperature °C</th>
<th>Gas phase species</th>
<th>$k_p$ gm$^2$/cm$^4$/sec</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>652</td>
<td>$O_2$</td>
<td>$1.61 \times 10^{-9}$</td>
</tr>
<tr>
<td>4</td>
<td>651</td>
<td>$O + O_2$</td>
<td>$1.61 \times 10^{-9}$</td>
</tr>
<tr>
<td>5</td>
<td>652</td>
<td>$O + O_2$</td>
<td>$1.69 \times 10^{-9}$</td>
</tr>
</tbody>
</table>

Further runs at 227°C, 495°C and 565°C confirmed that atoms had no effect. Data from these runs were not analysed quantitatively however as
several runs were carried out on the same sample leading to deviations from the parabolic law. Above about 570°C the bulk of the scale was assumed to be mainly non-stoichiometric FeO with successive layers of Fe3O4 and Fe2O3 outside it, below 570°C the FeO layer is known to disappear or to remain only as a thin subscale on top of the metal.

Silver.

The results of oxidizing silver by atomic oxygen are shown in table 4.

<table>
<thead>
<tr>
<th>Run</th>
<th>Temperature °C</th>
<th>kP/cm²·sec</th>
<th>kE/cm²·sec</th>
</tr>
</thead>
<tbody>
<tr>
<td>1f</td>
<td>277</td>
<td>5.81 .10^-11</td>
<td></td>
</tr>
<tr>
<td>1g</td>
<td>225</td>
<td></td>
<td>1.45 .10^-14</td>
</tr>
<tr>
<td>1h</td>
<td>168</td>
<td></td>
<td>1.82 .10^-15</td>
</tr>
<tr>
<td>1i</td>
<td>310</td>
<td>1.52 .10^-10</td>
<td></td>
</tr>
<tr>
<td>1j</td>
<td>65</td>
<td>1.10 .10^-11</td>
<td></td>
</tr>
<tr>
<td>2b</td>
<td>655</td>
<td>5.53 .10^-12</td>
<td></td>
</tr>
</tbody>
</table>

Thus, in the main, the oxidation follows a parabolic rate law with deviations towards cubic in some instances, see Fig. II.3. The majority of runs were carried out on the same sample as the silver was regenerated between runs by heating the oxide to above 350°C at 10^-5 mm. The oxidation rate is seen to be remarkably fast for such low temperatures (c.f. kP for Co or Ni at 600°C - 700°C) this being largely due to the low temperature coefficient for the process determined here as 10 ± 3.5 kcal/mole for the temperature range 170°C to 310°C. Below 170°C the rate is even faster than would be expected from an extrapolation of the higher temperature
results. Above 170°C only Ag₂O appears in the scale but at room
temperature both Ag₂O and an outer layer of AgO are present, about
40% of the oxygen being in the latter.

Copper.

The results obtained for copper fall into two groups - below 250°C
where only thin oxide films are formed and above 400°C where thick films
are obtained. The thin film region will be considered first. With
O₂ the uptake was too small for the data to be analysed accurately.

In the presence of atoms the uptake was increased by a factor of about
seven. Earlier work (e.g. by Gathney et. al. Acta Metallurgia 4,
145, 153, (1956)) has shown that the rate of oxidation depends on the
crystal face exposed; the rate with atoms is five or six times faster
than the fastest rate observed with O₂. Initially at about 150°C the
uptake followed a logarithmic law but the later stages of the runs could
also be represented by a cubic plot. The range of applicability of the
cubic law increased with temperature extending over a complete run by
200°C. The cubic rate constants were:

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>k c (cm³/cm²·s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>140</td>
<td>3.52 × 10⁻¹⁹</td>
</tr>
<tr>
<td>163</td>
<td>1.12 × 10⁻¹⁸</td>
</tr>
<tr>
<td>213</td>
<td>8.73 × 10⁻¹⁸</td>
</tr>
</tbody>
</table>

and the activation energy was 18 kcal/mole. The scale was blackish
indicating that it was mostly CuO₂.

Above 400°C the oxidation behaviour was somewhat more complex. In
molecular oxygen at 400°C the rate was linear e.g. at 453°C,
\[ k_p = 9.33 \times 10^{-9} \text{ cm}^2/\text{sec.}, \text{ but at 19.2 cm pressure of } O_2 \text{ the rate was parabolic, } (k_p = 2.77 \times 10^{-11} \text{ cm}^2/\text{sec.}) \text{ comparing well with rates observed by other workers at similar pressures (e.g. Valenci Rev. Metall 45, 10, (1948)). The linear rate constant increased almost linearly with pressure (see Fig. II.4 for sample 6 at 575°C) until by the time the pressure reached 0.5 atm diffusion through the scale became rate determining and the parabolic rate law was followed.}

In comparison, only a short linear period was observed in the presence of atoms and at comparable temperatures and pressures the rate constant is about fifteen times greater than with \( O_2 \). Subsequently parabolic law is followed, \( k_p \) being comparable with that obtained with high pressures of \( O_2 \), see Fig. II.4. The activation energy of 20.7 kcal/mole from the parabolic rate constants is in good agreement with Valenci's value of 20.1 kcal/mole. The scale was again predominantly \( CuO \) though it was probable that, in the initial periods when the linear rate was observed, only \( Cu_2O \) was present. (Honjo, J. Phys. Soc. Jap. 4, 330, (1949)).

Zirconium.

The results obtained for the oxidation of zirconium are shown in table 5.
TABLE 5.

The Oxidation of Zirconium.

<table>
<thead>
<tr>
<th>Run</th>
<th>Temperature °C</th>
<th>G.s phase species</th>
<th>$k_p^{2p}$/cm$^4$/sec</th>
<th>$k_p^{2p}$/cm$^6$/sec</th>
</tr>
</thead>
<tbody>
<tr>
<td>2a</td>
<td>518</td>
<td>$O_2$</td>
<td>$6.28 \times 10^{-13}$</td>
<td></td>
</tr>
<tr>
<td>2b</td>
<td>520</td>
<td>$O + O_2$</td>
<td>$9.62 \times 10^{-13}$</td>
<td></td>
</tr>
<tr>
<td>2c</td>
<td>636</td>
<td>$O_2$</td>
<td>$1.62 \times 10^{-11}$</td>
<td></td>
</tr>
<tr>
<td>2d</td>
<td>637</td>
<td>$O + O_2$</td>
<td>$2.41 \times 10^{-11}$</td>
<td></td>
</tr>
<tr>
<td>6a</td>
<td>435</td>
<td>$O_2$</td>
<td>$4.00 \times 10^{-14}$</td>
<td></td>
</tr>
<tr>
<td>6b</td>
<td>435</td>
<td>$O + O_2$</td>
<td>$4.33 \times 10^{-14}$</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>498</td>
<td>$O + O_2$</td>
<td>$5.28 \times 10^{-13}$</td>
<td></td>
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<tr>
<td>8</td>
<td>496</td>
<td>$O_2$</td>
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<tr>
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<td>$1.71 \times 10^{-12}$</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>535</td>
<td>$O + O_2$</td>
<td>$1.50 \times 10^{-12}$</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>535</td>
<td>$O_2$</td>
<td>$4.18 \times 10^{-17}$</td>
<td></td>
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</table>

A mean squares analysis of the $k_p$ values gave activation energies of 33.4 kcal./mole and 38.8 kcal./mole for molecules and atoms respectively; the plots are shown in Fig. II.5. The presence of atoms increased the oxidation rate slightly e.g. at 560°C $k_p^{(atoms)}/k_p^{(molecules)} = 1.46$.

Other metals.

The results for the other metals will now be considered more briefly.

Aluminium.

The introduction of atoms appeared to have no effect on the oxidation rate in the temperature range 25 °C to 660°C (the melting point of aluminium).
Tantalum.

Atoms again appeared to have little effect when the scale was protective, the results were somewhat irreproducible however as the protectiveness of the scale varied. In some runs, where the rate of uptake was almost linear both the presence of atoms and higher pressures of O₂ increased the rate.

Tin.

The effect of atoms on the oxidation of tin foil (below 232°C) was small and was too small to be detected in the case of oxidation of liquid tin. In the latter case the scale again afforded a variable degree of protection.

Zinc and Cadmium.

Investigations were only carried out in the temperature range 25°C to 150°C, due to the high vapour pressure of the metals. No oxidation was detectable in the presence of either atomic or molecular oxygen.

Palladium.

Only thin oxide films formed on palladium; at the higher temperatures (above 550°C) where, at atmospheric pressures, a thick oxide film results, the oxide decomposed at the low operating pressures used in the present work. The uptake in all temperature range 350°C to 450°C was approximately logarithmic but induction periods were observed initially. Atoms tended to eliminate these induction periods but did not cause any increase in the total uptake.

Platinum and Gold.

Neither of these metals were oxidised to any detectable extent by either atomic or molecular oxygen, in the temperature range 25°C to 500°C.
Discussion.

Nickel.

When nickel is oxidised it forms a scale consisting of the single p-type oxide, NiO. According to the Wagner model the scale grows as a result of transport of fresh metal to the gas oxide interface under a concentration gradient of nickel ion vacancies, Ni$_{2+}$*$. At the metal/oxide interface the Ni$_{2+}$* concentration is almost zero but at the gas/oxide interface it is much larger due to the reaction

$$2\text{Ni}^{2+} + \frac{1}{2}\text{O}_2 = \text{NiO} + \text{Ni}^{2+} + 2\text{Ni}^{3+}$$

(when the gas is O$_2$), All charged species are on the cation sub-lattice in NiO. $\Delta G_1^0$ is the partial nickel standard free energy of reaction (1).

Applying the law of mass action

$$(\text{Ni}^{2+})* (\text{Ni}^{3+})^2 = P_{O_2} \frac{1}{2} \exp (-\Delta G_1^0/RT)$$

assuming that concentrations are equal to activities and that (Ni$^{2+}$) and (NiO) are constant. Thus from (1) (Ni$^{2+})* = \frac{1}{2}(\text{Ni}^{3+})$ and

$$\frac{(\text{Ni}^{2+})*}{(\text{Ni}^{3+})} = 4 \frac{P_{O_2}^{1/2}}{\exp (-\Delta G_1^0/3RT)}$$

(1a)

When atoms are present the analogue of reaction (1) is

$$2\text{Ni}^{2+} + 0 = \text{NiO} + \text{Ni}^{2+} + 2\text{Ni}^{3+}$$

(2)

and by similar analysis

$$\frac{(\text{Ni}^{2+})*}{(\text{Ni}^{3+})} = 4 \frac{P_{O_2}^{1/2}}{\exp (-\Delta G_2^0/3RT)}$$

(2a)

Thus the vacancy concentration is more sensitive to the pressure of atoms.

More important, however, is the difference between $\Delta G_1^0$ and $\Delta G_2^0$ - subtraction of (2) from (1) gives

$$\frac{1}{2}\text{O}_2 = 0 = \Delta G_1^0 - \Delta G_2^0 = \Delta G_2^0$$

hence (2a) becomes

$$\frac{(\text{Ni}^{2+})*}{(\text{Ni}^{3+})} = 4 \frac{P_{O_2}^{1/2}}{\exp (-\Delta G_2^0/3RT)} \exp (-\Delta G_2^0/3RT)$$

As $\Delta G_2^0$ is positive, at comparable pressures, this leads to far more vacancies in the scale, near the gas/oxide interface, than in nickel, the concentration...
gradient across the scale rises by almost the same factor.

At 650°C and 40 kPa pressure PO₃ in atmospheres is about 5.10⁻⁵, PO, when the discharge is on, is about 10⁻³ and under these conditions the rates (Ni²⁺,c) (atm.s)/(Ni²⁺,c) (molecules) is 11.13 in very good agreement with the observed value of 10.

The oxidation rate depends on the product of the concentration and the mobility of the nickel vacancies. Hence, in the presence of O₂, the activation energy for oxidation is \( \Delta H^0/3 + V \) where \( \Delta H^0 \) is the standard heat of reaction (1) and \( V \) is the activation energy for diffusion of the vacancy. When atoms are present the activation energy becomes \( \Delta H^0/3 + V \) where \( \Delta H^0 = \Delta H^0_1 - \Delta H^0_2 \). \( \Delta H^0_2 \) is about 59 kcal/mole so that the presence of atoms should lower the activation energy by nearly 20 kcal/mole.

In fact the activation energy in the presence of 0₂ is only about 5 kcal lower than in the presence of O₂. Most of this discrepancy may be attributed to a trace of iron impurity shown by spectrographic analysis to be present to the extent of about 1 ppm. This is sufficient to explain the rather high values of \( k \) at PO₂ = 5.10⁻⁵ and as the iron has most effect at the lower temperature the slightly low value for the activation energy. Further it will reduce the observed effect due to a loss of gain in the low temperature range as compared with that predicted for pure nickel.

Cobalt.

At the higher temperatures where only CoO is formed, the situation is similar to that discussed for NiO and atoms cause a large change in the \( k \) value. Slight numerical differences occur because of inter-atom association of defects in the scale i.e.; the reaction of reaction (1) is

\[
2\text{Co}^{2+} + \frac{1}{2}\text{O}_2 = \text{CoO} + (\text{Co}^{2+}_1, \text{Co}^{3+}_1) + \text{Co}^{3+}
\]

and the concentration of core-localized vacancies is PO₂⁻¹; in the presence of oxygen it accordingly becomes PO₂⁻².
The main discrepancy between theory and practice arises from the fact that an outer layer of Co₃O₄ should still be thermodynamically stable at 700°C in the presence of atoms but none could be detected. Two factors may account for this: at these temperatures the arrival rate of atoms is no longer much greater than the incorporation rate of oxygen by the scale and this may lead to non-equilibrium conditions at the surface, further, as with silver and palladium, it was found that the critical temperature at which an oxide became unstable in the presence of atoms was that at which its decomposition rate, in the presence of O₂, became very rapid, again suggesting that kinetic factors were playing an important role.

At the lower temperatures, the bulk of the scale is still CoO and as it is bounded by the Co/CoO and the CoO/Co₃O₄ interfaces its growth rate is largely independent of changes in the oxygen potential in the gas phase. Thus theory predicts that the presence of atoms will affect only the minor constituent in the scale and their effect will be small. The final prediction is correct but there is no evidence that the Co₃O₄ content had increased.

Iron.

The absence of any effects due to atoms is in complete agreement with the predictions of theory. Any FeO in the scale (as is the major constituent above about 570°C) grows between the Fe/FeO and the FeO/Fe₃O₄ interfaces and is not affected by atoms in this region. Similarly Fe₃O₄ (as the major constituent below 570°C) grows between the Fe₂O/Fe₃O₄ and the Fe₃O₄/FeO interfaces. Finally the outer layer of Fe₂O₃ is not affected and its growth rate is virtually independent of the oxygen potential in the gas.
phase - this point will be considered in greater detail in the section on n-type oxides.

Silver.

In the review of previous work it was pointed out that oxygen is chemisorbed on silver up to 310°C. The present work shows clearly that it is not diffusion rates in the solid that prevents this oxygen forming an oxide scale. Thus it must be that molecular oxygen is not dissociatively absorbed on the silver at temperatures below which the oxides of silver are stable (1050°C for Ag2O when PO2 = 1).

The low activation energy for oxidation is in keeping with the low activation energy for ionic conduction and diffusion in several other silver salts. As Ag2O is a p-type oxide the high rate of oxidation observed at 600°C will, at least in part, be due to the considerable increase in PO2 with the reduction in temperature. The situation is complicated by the formation of an outer layer of AgO, containing nearly 40% of the total oxygen uptake. Unfortunately a rigorous analysis of the results is not possible as the stability range of the AgO appears to be governed by kinetic rather than thermodynamic factors.

Copper.

In the low temperature region the increase in uptake due to atoms may well be caused by an increase in the amount of dissociated oxygen adsorbed on the surface as in the case of silver. Following charge transfer this will lead to a stronger field across the film, ratios to increase uptake at lower rates as observed.

At high temperatures and low pressure of O2 the diffusion through the oxide is faster than the reaction at the gas/oxide interface.
Thus increasing the oxygen potential in the gas phase, either by increasing the pressure of O₂ or by introducing atoms, causes a large increase in the oxidation rate. When the extent of oxidation is large, however, both CuO and Cu₂O are present in the scale and the effect of atoms or of variation in P(O₂) is small.

Metals Oxidising to form n-type oxides.

This group includes the metals Zr, Al, Zn, Cd, Te and Sn. In the oxidation of zirconium, the ZrO₂ scale grows at the Zr/ZrO₂ interface following transport of fresh oxygen through the scale by means of anion vacancies. The concentration of these vacancies is large at the metal/oxide interface and virtually zero, even in the presence of O₂, at the gas oxide interface. Thus the presence of atoms in the gas phase only brings the lower concentration even closer to zero and their effect is consequently negligible. Similarly the supply of interstitial zinc responsible for the transport of fresh material across a growing scale of ZnO on Zn is governed almost entirely by the Zn/ZnO interface reactions.

In the cases where the scale remained protective this predicted behaviour was in general observed for the metals investigated in this work. The small but definite increase in kₚ values, due to atoms, observed for zirconium is probably connected with the surface preparation. Gulbransen and Andrew (J. Metals 2, 394, (1957)) have shown that after an electrochemical polish the rate of oxidation of zirconium is slower and more likely to follow the parabolic time law than after abrasion. The results presented here confirm the predominance of the parabolic rate law after this preparation and the kₚ values for O₂ arc in good
agreement with Gulbransen's. The effect of atoms is rather less than that of abrasion; both probably influence the subscale formed at the metal oxide interface, the bulk of the scale behaving as predicted by the theoretical model.

**Palladium, platinum and gold.**

The results for palladium oxidation support the idea that the initial nucleation and growth of the oxide are speeded by the greater abundance of \( \text{O} \) or \( \text{O}^- \) on the surface resulting from the presence of atoms in the gas phase. Again the presence of atoms did not increase the temperature range over which the \( \text{PdO} \) was stable as would be expected on thermodynamic grounds. The inactivity of gold and platinum in the presence of oxygen appears to be due to a property of the metals, possibly connected with their high first ionisation potentials, and does not depend on the state of the oxygen as is found for silver.

**Conclusions.**

The survey has shown that the effect of oxygen atoms in the gas phase, on the oxidation of a wide range of metals, can be explained in a semiquantitative manner by the existing theories of oxidation. Maximum effects were found as predicted, when the scale consisted of a single \( p \)-type oxide such as \( \text{NiO} \) and very small effects resulted when an \( n \)-type oxide such as \( \text{ZrO}_2 \) or a multi-oxide scale such as \( \text{FeO}/\text{Fe}_2\text{O}_3/\text{Fe}_3\text{O}_4 \) was formed. Further work with oxides like \( \text{NiO} \) in conjunction with a more abundant and variable, known atom concentration would provide a more rigorous test of theory, as the pressure dependence and activation energy in the presence of atoms could then be determined accurately.
The work on silver suggests that in this case it is the absence of dissociative adsorption of $O_2$ that prevents oxidation in molecular oxygen. Here, as with cobalt, further work with large known atom concentrations should provide more information about the interesting problem of the stability of higher oxides in the presence of atomic oxygen.
Nickel Oxidation. Sample 1.

Fig. II.1.
Cobalt oxidation, sample 1

Fig. A.2
The oxidation of silver by atoms

Fig. II.3.
Copper oxidation, above 500°C.

Fig. 24
Zirconium oxidation, temperature coefficient determination.

Fig. 11.5
CHAPTER III.


Abstract.

The recombination of oxygen atoms has been studied on the surfaces of the alkali metal tungsten bronzes of general formula $\text{M}_x\text{WO}_3$ where $x = \text{Li, Na or K}$, and $0 < x \leq 0.80$. The recombination coefficients, $r$, defined as the fraction of atomic collisions with the surface which results in recombination, were measured at 300$^\circ$K by a side-arm method. Catalytic activities were found to be closely related to the electronic properties of the bronzes.

Recombination measurements were supplemented by studies of electrical resistance and crystal structure.
The activity of a catalyst may be determined by both electronic and geometric factors. The alkali metal tungsten bronzes of composition $M_xW_3O_7$, where $0 < x < 0.80$ provide an interesting series of gradually stable, compounds in which the electronic properties change in a marked and controllable way but in which lattice parameters are only slightly affected. They are, therefore, useful systems to study in testing the role of the electronic factor in catalysis.

The crystal structure changes little with composition. The high $x$-value sodium and lithium bronzes have the perovskite structure (fig. III.1) and as $x$ decreases the structure becomes progressively more distorted, passing through two tetragonal phases to the triclinically distorted structure of $WO_3$. The Potassium bronzes have only tetragonal and hexagonal structures over their narrow composition range. The electrical properties of the tungsten bronzes are well known. They are $n$-type semiconductors for $x < 0.25$ and metallic conductors for $x > 0.25$ with one 'free electron' per alkali metal atom. This change in conductivity type may be correlated with the crystal structure. For $x < 0.25$ the alkali metal atoms may reside at the diagonally opposite corners of the cube; however when $x > 0.25$ the metal atoms must reside at corners on the same face of the cube. Here the interatomic distance is very similar to that in the pure metal e.g. for sodium 3.82 Å in the bronze against 3.72 Å in the pure metal. It is reasonable to suppose that the activity of the bronzes for oxygen atom recombination, which involves electron transfer between adsorbate and adsorbent will be composition dependent.
Fig. 1. Lennard-Jones structure of La

○, La; ⊙, O.
The recombination of oxygen atoms on other oxide surfaces at
room temperature has been found to be first order with respect to
gaseous atoms. The rate determining step is probably
\[
O(g) + O^{-}(ads.) \rightarrow O_{2}(g) + \text{c(solid)}
\]

Experimental

Atomic recombination coefficients were measured by the side-arm
method of Smith using an apparatus similar to that described in Part 1.
The side arm was enclosed in a water-jacket kept at 300°C, and the
pressure was 28. The samples of catalyst were coated onto
the interior of Pyrex cylinders from an acetone or water slurry of the
powder. The Pyrex cylinders, which were 15 cm long x 3 cm diameter,
fitted closely inside the side arm. The method of measurement and
calculation of the recombination coefficients, is discussed in Part 1.

The bronzes were prepared according to the equation
\[
\frac{x}{2} \cdot k_{2} WO_{4} + \frac{1-2x}{3} \cdotWO_{3} + \frac{x}{6} W \rightarrow xWO_{3}
\]

by heating the appropriate tungstate, tungstic oxide and tungsten
powder in an alumina boat at 850°C under vacuum (10^{-5} torr) for five
hours. A sample of pure tungstic oxide was heated in the same manner.
The bronzes were washed successively with very dilute boiling NaOH,
water, conc. HCl, water and acetone. The spectra showed the presence
of less than 0.2% of impurities (alkali metals and molybdenum) in the
starting materials. The sodium tungsten bronzes were analysed for
sodium content using neutron activation analysis with sodium carbonate
as standard. The 2.76 eV peak was counted using a \(^{6}\) -spectrograph
acousto-ion counter. The lithium and potassium bronzes were analysed
for alkali metal content by flame spectrophotometry using a Unicam SP 900 model. These bronzes were brought into solution by fusion in 1 \( \text{Na}_2\text{CO}_3 : 3 \text{NaNO}_3 \) mixture. Standard solutions were made from lithium or potassium nitrate and sodium nitrate to match the sodium content of the unknown solutions from the fusion mixture.

Surface areas of the powders were determined by the B.E.T. method using krypton adsorption at liquid nitrogen temperatures. The value 19.5 \( \AA^2 \) was used for the area of a krypton molecule.

X-ray photographs were taken with a Guinier-de Wolff camera using \( \text{CuK}_{\alpha} \) radiation with exposure times of 72 hours; thorite was mixed with the samples to act as a standard.

The electrical conductivities of pellet samples were measured at 10\(^{-5}\) torr using a Wayne-Kerr R.F. Bridge and an Avo electrical test meter. Measurements on the R.F. Bridge were restricted to resistances over 25 ohms. Measurements on pellets are unsatisfactory for low resistance materials, when the intergrain resistances may be relatively large. The resistances of the samples fell sharply from pure tungstic oxide to the metallic bronzes, table 1. This is in agreement with previous results.\(^4\)

**Results.**

Within experimental accuracy, oxygen atom recombination was found to be first order with respect to gaseous atoms at 300\(^0\)K on all the surfaces studied. Table 1 lists the values of the recombination coefficients, which are the mean of four runs on two samples and their standard deviations; the surface areas which are the mean of two
<table>
<thead>
<tr>
<th>Nominal</th>
<th>Analysed</th>
<th>$m^2/gm$</th>
<th>Recombination Coefficient $\times 10^2$</th>
<th>Specific Resistivity OHM-CM</th>
<th>Crystal Structure*</th>
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</thead>
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<tr>
<td>WO$_3$</td>
<td>-</td>
<td>2.247</td>
<td>3.71±0.01</td>
<td>5200</td>
<td>WO$_3$</td>
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<tr>
<td>Na 0.10</td>
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<td>0.317</td>
<td>2.61±0.08</td>
<td>150</td>
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<td>1.304</td>
<td>1.89±0.02</td>
<td>25</td>
<td>I,II</td>
</tr>
<tr>
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<td>-</td>
<td>1.83±0.08</td>
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<td>&lt;10</td>
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<tr>
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<td>-</td>
<td>3.94±0.50</td>
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<td>-</td>
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<td>(II),C</td>
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<tr>
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<td>0.888</td>
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* I and II - tetragonal I and II phases $^2$, C - cubic, X - some less symmetrical system.
Determinations; the electrical resistivity at room temperature; the analytical data and the x-ray structures.

**Discussion.**

Oxygen atom recombination is a very rapid reaction and so only the exposed outer layers of the catalyst surface are likely to be available for reaction. Some evidence for this is provided by the potassium bronzes which have similar activities to the sodium compounds and yet have very different surface areas. A quantitative expression for the effect of surface area on recombination coefficient is derived in the appendix. Fig. III.3, compiled from this expression, shows that the overall activity pattern is insensitive to changes in surface area.

Fig. III.2 shows that the pattern of catalytic activity for oxygen atom recombination is approximately the same whether Ni, Na, Li or K in $\text{M}_x\text{WO}_3$. This is a clear indication that electronic factors dominate the catalytic behaviour. As the crystal structure is dependent on the alkali metal present, it cannot have any marked effect on the activity pattern. It is interesting to note that the changes in surface area correspond almost exactly to the changes in crystal structure.

The conduction band in the bronzes can be considered as made up either by overlap of the alkali metal p orbitals or by overlap of the tungsten 5d ($t_{2g}$) orbitals. The former suggestion seems the more likely as the lattice dimensions permit appreciable overlap of the alkali metal p-orbitals, whereas little overlap of the tungsten 5d orbitals would be expected. However only when there are sufficient alkali metal atoms, $x > 0.25$, will appreciable overlap of p orbitals be possible.
Fig. 2. The recombination coefficients of the tungsten bronzes. Δ, lithium; O, sodium; ⚫, potassium.
Fig. 3. $\gamma$ and $\gamma_f$ for the sodium tungstate bronze.

$\gamma$ --- $\gamma_f$ ---
When this occurs their energy will fall relative to those in the free atom. The $s$ orbitals overlap appreciably with the filled $WO_3$ orbitals thus increasing their energy, by electrostatic repulsion, relative to the free atom. N.M.R. studies$^{11}$ show that the $s$ orbitals of sodium do not contribute to the conduction band. For $x < 0.25$, the alkali metal electrons are localised and conduction will take place by a 'hopping' process probably involving some excited state in the $WO_3$ complex. Beyond this composition range the alkali metal electrons will become delocalised and metallic conduction will take place via the $p$-band.

It is probable that oxygen in the presence of oxygen atoms is adsorbed predominantly as $O^-$(ads) ions.$^1$ If the direct recombination reaction

$$O(g) + O^-$(ads) $\rightarrow$ $O_2(g)$ + $e$(solid)$$

is rate determining, then low activation energy for this reaction and high coverage will promote the recombination rate.

In the semiconducting region, $x < 0.25$, the Fermi level will increase with increasing alkali metal content. According to the boundary layer theory of chemisorption,$^8$ oxygen will be adsorbed depletively to low coverages on $n$-type oxides, such as the bronzes within this composition range. The activation energy for the reaction is determined by the position of the Fermi level. As the rate depends exponentially on the activation energy, and directly on the coverage, which is proportional to the square root of the impurity content,$^8$ it is most probable that the effect of increasing activation energy will be dominant. Thus catalytic activity is predicted to decrease.
from pure $WO_3$ to $Na_0.25WO_3$.

As $x$ increases from 0.25 the alkali metal p-orbitals will become stabilised due to mutual overlap, and the Fermi level will decrease sharply. This will cause a decrease in the activation energy and an increase in the coverage. Hence the catalytic activity should increase sharply.

The p-band is thought to be fairly wide, ca $3eV$, and so the energy levels within this band will be widely spaced. As the sodium content increases this band will fill and at some composition the Fermi level will reverse its general downward trend and increases in value again. This will cause an increase in the activation energy of the reaction and hence lead to a decrease in the catalytic activity with increasing sodium content.

The measured values of the recombination coefficients thus confirm the general pattern predicted by the band theory.

---------

APPENDIX.

The Effect of Surface Area on the Recombination Coefficients.

A simple model is postulated, comprising a cylindrical coating of powder with open-ended pores of length $2l$ and radius $r$. (Fig. III.4)

$A$ is the total surface area/gram.

$A_1$ is the geometrical surface area/gram.

$\rho$ is the measured density of the coating.
Fig. 4. Dimensions of core in oxide coating.
\( \nu_0 \) is the apparent value of \( \nu \), calculated on the assumption that the area exposed by 1 g. of material is the corresponding geometric area of cylinder, \( k_1 \).

\( \nu_t \) is the true recombination coefficient, calculated on a basis of the actual available area, \( k_2 \).

\( f \) is the fraction of surface used, \( k_2/A \).

\( \bar{v} \) is the mean speed of oxygen atoms in the gas phase.

\( n \) is the concentration of atoms in the gas phase.

Rate of reaction per unit area of cylinder = \( \nu_0 \bar{v} n/f \)

\[ = \nu_t \bar{v} n_{12}/k_1 A_1 \]

\[ = \nu_t \bar{v} n f/4 A_1 \]

Thus \( \nu_0 = \nu_t \bar{v} f A_1/4 \)

If diffusion is the only means of transport of reacting species into the pore, then the change in concentration in the pore is

\[ d\bar{n}/dx = r^2Ddx \frac{d^2n}{dx^2} \]

where \( x \) is the distance along the pore, and \( D \) is Krümmel's diffusion coefficient. At equilibrium, this is balanced by atom removal

\[ \nu_t \bar{v} n f/2 = r^2D dx \frac{d^2n}{dx^2} \]

Using the boundary conditions \( n = n_0 \) when \( x = 0 \) and that \( d\bar{n}/dx = 0 \) at \( x = 1 \), integration gives

\[ n = n_0 \cosh \left[ h(1-x/1) \right] / \cosh h \]

where

\[ h = 1 \left( \nu_t \bar{v}/2rD \right)^{\frac{1}{2}} \]

Differentiating, \( d\bar{n}/dx = h n_0 / \cosh h \cdot \sinh \sqrt{h(1-x/1)}/ \cosh h \)

The rate of reaction per half-pore is the rate at which reactant flows in
i.e. \( \Delta r^2 D \) times the concentration gradient at \( x = 0 \)

\[
\text{rate} = \Delta r^2 D \ln \alpha \tanh \frac{h}{l}
\]

If the pore area were completely available the rate would be

\[
2 \pi r l \sqrt{\frac{\Delta r}{t}} \tanh \frac{h}{4}
\]

Hence

\[
f = 4 \pi r^2 D \ln \alpha \tanh \frac{h}{2} \sqrt{\frac{\Delta r}{t}} \tanh \frac{h}{2}
\]

\[
= \tanh \frac{h}{h}
\]

Now

\[
D = 2 \sqrt{\frac{\Delta r}{t}} / 3
\]

and

\[
r = 2(1 - \theta) \sqrt{\frac{\Delta r}{t}} A
\]

where \( \theta \) is the porosity of the material, \( g \) is the roughness factor, and assuming that the pore walls have the same properties as other surfaces.

Hence the equation for recombination in the pores is

\[
\gamma_0 = \gamma_t A \tanh \frac{h}{A_i h}
\]

For the smooth surface \( \gamma_0 = g \gamma_t \). As \( \theta \) represents the fraction of the geometrical area exposed to the recombination by the pore mouths, the combined value becomes

\[
\gamma_0 = (1 - \theta) g \gamma_t + \theta \gamma_t A \tanh \frac{h}{A_i h}.
\]

The x-ray densities of the bronzes are about 7.2 \( \text{g cm}^{-3} \). The porosity of the coatings was slightly less than \( \frac{1}{3} \), but the value \( \frac{1}{3} \) will be used here for simplicity. The roughness factor will be assumed to have a value of 2. The pores are assumed to penetrate to an average half the distance from the coating surface to the glass cylinder. For a coating weight of 1.5 g in a cylinder 15 cm long and 2.8 cm diameter.
\[ l = 2.23 \times 10^{-3} \text{ cm} \]
\[ A_1 = 62.3 \text{ cm}^2/\text{g} \]

and

\[ h = 70A \; \sqrt{t} \] where \( A \) is in \( \text{cm}^2/\text{g} \).

Hence

\[ Y_0 = \sqrt{t} \left( 1 + 0.116 \tanh \left( 70A \sqrt{t} \right) \right) / \sqrt{t} \]

In fig. 3 \( Y_0 \) and \( Y_t \) are plotted against composition for some of the sodium bronzes. This shows that the variation in surface area from powder to powder does not alter significantly the pattern of the catalytic activities.

**REFERENCES**

4. Shanks, Sides and Danielson, ref. 2, 237.
7. Sienko, ref. 2, 224.
   Nerath and Wallace, Physics, Rev., 1962, 127, 724.
In the course of work on the recombination of chlorine atoms, it was found necessary to devise a gauge capable of measuring pressures of chlorine in the range 0.1 - 10 mm Hg. This posed certain problems, since chlorine is a very reactive gas, attacking mercury and most other metals. The method of measurement had to be sensitive, and also give values of pressure independent of the nature of the gas. The latter condition was essential, since the gauge had to be calibrated against a mercury gauge with a gas inert to mercury, and also since the pressures of mixtures containing chlorine and other gases would be required.

A capacitance method was chosen, since a proximity meter is obtainable which can detect very small changes in capacitance. The principle of the method is that small changes in level of a mercury U-tube manometer are recorded as capacitance changes. Each limb is about 4.5 cm. diameter. The mercury surface in the arm exposed to the chlorine is protected with a layer of dioctyl phthalate. The other arm contains a brass plate sealed into the glass so that its surface is a few millimetres above that of the mercury. These two surfaces form the condenser, the mercury being earthed via a tungsten seal, and dibutyl phthalate acting as the dielectric medium, since it has a fairly high dielectric constant (6.45 at 30°C) and a low vapour pressure. The brass plate is connected to the proximity meter. The total capacitance is of the order of 40 pF, and the proximity meter can detect changes of 0.001 pF at maximum sensitivity. However, the most
sensitive ranges were not needed, and in any case require a very great
control of mechanical stability and temperature. Even so, the gauge is
on a rigid shelf of the wall, and is connected to the rest of the
apparatus, which is subject to vibration from the rotary pump, by
flexible tubing. Also, the gauge and the capacitance bridge box are
enclosed in a thermostated perspex box.

The exact distance between the brass plate and the mercury surface
can be altered by means of a mercury reservoir to give approximately
the required range for a suitable sensitivity setting of the meter. To
use the gauge, the reference arm is evacuated and isolated, a charcoal
trap in liquid nitrogen ensuring a constant pressure. A suitable
pressure range for a particular experiment is obtained by subjecting the
gauge to the lowest required pressure, and balancing this out. The
sensitivity is adjusted so that the highest required pressure is at the
upper end of the scale. The gauge is then calibrated against a McLeod
gauge, the calibration being linear except at the top end of the scale
or at very high sensitivity values.

The gauge has been found to be quite satisfactory in use, giving
reproducible pressure readings to 0.01 mm. Hg.