Engn UMP 1618 THE FUNDAMENTAL EFFECTS OF AGING ON THE CREEP PROPERTIES OF SOLUTION-TREATED LOW-CARBON N-155 ALLOY.

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### SUMMARY

A method is developed whereby the fundamental mechanisms are investigated by which processing, heat-treatment and chemical composition control the properties of alloys at high temperatures. This method uses metallographic examination, both optical and electronic, studies of x-ray diffraction line widths, intensities and lattice parameters, and hardness surveys, to evaluate fundamental structural conditions. Wech-anical properties at high temperatures are then measured and correlated with these measured structural conditions.

In accordance with this method, a study was made of the fundamental mechanism by which aging controlled the short time creep and rupture properties of solution-treated Low-Carbon N-155 alloy at 1200° F.

The test stock was solution-treated at 2200° F for 10 hours, water-quenched, and aged for time periods up to 1000 hours at 1200°, 1400°, and 1600° F.

Correlation of the structural effects of aging with the mechanical properties indicated that aging had the following effects on solution-treated Low-Carbon N-155 alloy:

- a. Aging resulted in progressive lowering of short time creep resistance through removal from solid solution of large radii or substitutional atoms by precipitation.
- b. Short time aging resulted in marked increase in short time rupture strengths through the growth of a grain boundary phase which eliminated intergranular cracking. Long time aging resulted in little further change in short time rupture strength.

- c. Because aging lowered the creep resistance while raising the rupture strength, aged material exhibited greater ductility before fracture than unaged material.
- d. Calculations are carried out to show the probable character of the strain field induced in the solution-treated state by the presence of the large radius atoms Mo, W, and Cb and the substitutional atoms H and C. The effect of this strain field on the creep resistance is also considered quantitatively.

These findings, however, should not be considered generally applicable to long time strength, strength at other temperatures, to the same alloy in other conditions of treatment. In order to develop a general theory of high temperature strength, additional data on other alloys and on the same alloy in the other conditions, will have to be gathered.

#### INTRODUCTION

This report covers the first part of a continuing investigation into the fundamental behavior at high temperatures of austenitic alloys designed for use in aircraft propulsion systems. Previous investigation (see reference 1 for example) has shown in part the effects of prior processing and heat treatment conditions on the high temperature properties of Low-Carbon N-155 and various other austenitic high temperature alloys. In order to develop better and practicable alloys on a scientific basis, to utilize to the fullest extent possible critical materials, and to point out logical methods of production control for uniform properties more knowledge must be gained of the fundamental reasons for the high temperature behavior of austenitic high temperature alloys.

The basic assumption of the investigation was that the behavior of certain alloys at high temperature is dependent on their microstructure and the lattice conditions of the matrix. The experimental program was therefore designed to first measure these two characteristics of one alloy, Low-Carbon N-155, as influenced by heat treatment, chemical composition and the metallurgical reactions during exposure to temperature and stress. Optical and electron microscope techniques combined, separation and analysis of microconstitutents would be used to define structural conditions. The lattice conditions of the matrix, particularily the strains present, would be measured by x-ray diffraction techniques. Second, the creep and rupture properties, corresponding to these structural conditions, were to be established. Third, these data would then be correlated and interpreted using the fundamental physics of solids and plastic flow to as great an extent as possible.

It was felt that by concentrating at first wholly on one particular alloy the thorough understanding thus gained could be best generalized for extension to other alloys. The experimental variations which necessarily had to be covered require a prohibitive amount of work for even two alloys. Low-Carbon N-155 alloy was chosen primarily because it was a representative alloy of a type of importance in the aircraft propulsion field. In addition there was a considerable background of experimental data for this alloy which would be of value to the investigation.

This report presents the results obtained to date on bar stock material, solution-treated at 2200° F for 10 hours and aged for time periods up to 1000 hours at 1200°, 1400°, and 1600° F. The creep and rupture data were confined to short time periods at 1200° F under a restricted stress range. This is therefore in the nature of a report on the techniques developed and progress made to date. A large amount of additional experimental work must be done before all temperatures, stresses and time periods of in-

terest are related to the structural conditions of the alloy resulting from the wide range of possible prior treatments.

The investigation is part of a research program on heat-resistant alloys for aircraft propulsion systems sponsored by the National Advisory Committee for Aeronautics at the Engineering Research Institute of the University of Michigan.

### TEST MATERIALS

Low-Carbon N-155 alloy bar stock was used in this investigation. It represents part of the complete product of one ingot of heat A-1726 rolled into the 7/8-inch broken corner square stock. Each bar from the ingot was numbered so that its position relative to the original ingot was known. The particular material considered herein came from the center of the ingot.

The composition of heat A-1726 reported by the supplier is listed below, together with the results of the analysis by the "niversity of the bar from the center of the particular ingot being considered:

Supplier's Heat Analysis	Chemical Composition (percent)									
	-	1.63	0.42	<u>Cr</u> 21.22	<u>Ni</u> 19.00	<u>Co</u> 19.70	2.90	<u>₩</u> 2.61	<u>Cb</u> 0.84	0.13
Bar Brom Center of Ingot	0.133	1.43	0.34	20.73	18.92	19.65	3.05	1.98	0.98	0.14

Table I gives the complete prosessing schedule reported for the ingot of heat A-1726 from which the test stock was taken.

Prior to use, the stock for this investigation was solution-treated 10 hours at 2200° F and water-quenched. The solution treatment was made un-

usually long for the purpose of distributing the precipitant atoms randomly in the matrix and so that internal strain, due to the prior working of the material, would be reduced to a very low level.

Moderate grain growth took place over the major portion of the bar cross section during the 10 hour treatment but on two diagonally opposite corners pronounced growth took place. A hardness survey of the assocled bar stock showed the average hardness across one diagonal to be higher than across the other; from this the conclusion may be drawn that the rolling operation had worked the bar across one diagonal preferentially. All mechanical testing and physical measurements were restricted to the fine grained section of the bar stock. Figure 1 shows representative structures of the cross section of the bar stock as-rolled and after the 10 hour solution treatment.

### EXPERIMENTAL PROCEDURES

The general procedure was to age the solution-treated stock at selected temperatures and time periods and then to carry out the microstructural studies and x-ray diffraction measurements in order to establish the structural characteristics resulting from the aging treatments. The strength properties resulting from the aging treatments were measured for short time periods at 1200° F. These experiments were intended to establish the relationship between short time creep and rupture properties and nucleation, precipitation, and precipitate particle size and distribution during aging.

The details of the experimental procedures are described in the following sections.

### Aging

Aging treatments were carried out at 1200°, 1400° and 1600° F for time periods of 1, 10, 100 and 1000 hours and such other intermediate times

as became necessary. The samples were heated in small automatically controlled muffle furnaces in an air atmosphere. These furnaces were at temperature when samples were placed in them and the time period of heating was considered started after the specimens had been in the furnace for 1/4 hour. After aging, all samples were air cooled. Sufficient stock was aged at each condition for the microstructural, x-ray and mechanical tests.

### Optical Microstructural Studies

After aging, the individual samples were ground on a cross-sectional face, polished with No. 1 emery cloth, No. 1, 1/0, 2/0 and 3/0 emery paper, transferred to a cloth disc and polished using a commercial chromium buffing compound and finished on a Gamal wet wheel; they were then electrolytically etched for 5 seconds in 10 per cent chromic acid at 1 ampere per square inch. After study under a microscope, representative photomicrographs were taken of each sample at 1000 diameters under slightly oblique illumination.

### Electron Microscope Studies

In order to obtain electron micrographs from metallographic samples, replicas which will transmit the electron stream must be prepared of the surfaces. For the investigations reported herein, Formvar replicas were prepared according to the technique outlined by Williams, et al. (see reference 2). The metallographic surfaces from which these replicas were prepared were the same polished and etched surfaces photographed optically as outlined in the sub-section above. The replicas (shadow cast with chromium) were then mounted in an RCA Model B electron microscope and photographs taken at approximately 3000X. Enlargement to 8500X was done photographically.

### X-Ray Studies

Sample Preparation: A great deal of difficulty with variable and non-reproducible diffraction data was initially encountered. The

difficulty was found to be due to both mechanical disturbed metal surfaces and non-random grain orientation. Eventually the development of the following technique produced results sufficiently free from these difficulties for this investigation:

1. A layer of metal 0.020 inch thick was electrolytically removed from the surface of all samples which were used for X-ray analysis. The amount of metal which had to be removed in order to get below the artificially strained surfaces produced by a cutoff wheel, by grinding, or as a result of metallographic polishing was determined by X-ray diffraction patterns of the type shown by figure 2, and made by the apparatus illustrated in figure 3. The patterns in figure 2, taken from a surface initially ground, show the emergence of the reflections of the individual grains and finally resolution of the  $\kappa_1 \kappa_2$  doublet of the molybdenum K radiation with increasing depth of metal removal. The amount of surface removal shown in figure 2 as necessary to obtain a strain free surface was typical for all samples used.

In order to maintain a surface that was flat during the electrolytic metal removal, a special cell was designed as shown in figure 4. It consisted essentially of a 250 cc cylindrical container with a copper plate on the bottom as a cathode. This plate was connected to the source of current by a lead through a glass-metal seal. About one-half way up the cell a watch glass containing a 5/8-inch hole was mounted horizontally. The metal sample acting as the anode was mounted 1/2 inch above the hole with the polished surface facing the hole. This hole acted to distribute the current evenly over the 7/8-inch square surface at this distance, and thus a plane surface was maintained during metal removal. Water cooling

was used to prevent pitting associated with electrolyte temperatures above 100°C.

The most satisfactory electrolyte was experimentally found to be a mixture of 1/3 concentrated HCl (37%) and 2/3 glycerine. This mixture had the best current efficiency, approximately 0.0000625 inch of metal removed per ampere-minute at 8 amperes per square inch, without excessive pitting. 200cc of this solution was sufficient, for 6 to 10 samples. Pitting occurred when the metal ion concentration became too high. Phosphoric acid-lycerine combinations gave good polished surfaces, but had low current efficiencies. Sulturic acid-glycerine mixtures caused passivation, chromic acid and hydrofluoric acid-glycerine mixtures left the surface badly pitted.

- 2. After the gross metal removal of step 1, the surface was given a high polish using undiluted duPont electropolishing solution for 5 minutes at 5 amperes per square inch. Metal removal in this step was negligible. The simple cell consisting of a beaker with a copper plate in the bottom was used for this step. No water cooling was necessary. Of several other electrolytics tried for this step, only a 40 per cent phosphoric acid-60 per cent glycerine mixture was nearly as satisfactory as the duPont solution.
- 3. After surface preparation, diffraction patterns were taken with the specimens either rotating or oscillating in the X-ray beam. The grain size of the Low-Carbon N155 specimens used was too large to present an effectively random distribution in the X-ray beam when the specimen was stationary. By moving the specimen under the beam during the time the

diffraction pattern was being taken, enough grains were presented to the beam to make the specimen approximately one of random grain orientation. In the case of line intensity measurements on a Norelco Spectrometer, a specimen holder was designed and built which rotated the specimens at approximately 17 rps. Care was taken to insure that the plane of polish was perpendicular to the axis of spin. In the case of line width measurements, the specimens were oscillated ±10° about an axis which was normal to the incoming X-ray beam, but parallel to the film. Finally, in making lattice parameter measurements, the specimens were rotated about 2 rps, with the axis of rotation parallel with, but offset from the axis of the X-ray beam.

Diffraction Line Peak Intensity Studies: These studies were confined to the (III) line of the austenite matrix, this line being the strongest line and within the range of the Norelco Spectrometer. Copper Kala2 radiation was used. Plots of the (III) line obtained with this spectrometer and an automatic recorder were measured graphically for peak height. Because the basic measurement needed was the line height of a given aged sample relative to unaged material, an unaged sample was run alternately with each aged sample.

Checks of the reproducibility of the (111) line measurements were made by the following procedures:

(1) The unaged solution-treated sample, used as a standard of comparison, was taken twice through the surface preparation step with, however, removal of only an additional 0.0025 inch of metal the second time. After taking the (111) line height

from the first surface, the X-ray tube and counter circuit were left on during the repolishing. The (111) line measurements were immediately taken from the second surface and found to check within the accuracy of the spectrometer. This surface was carefully preserved and used as a standard for subsequent measurements on aged samples.

- (2) The samples aged at 1200° F and 1600° F were alternately measured for (111) line peak intensity against the standard and repolished for a minimum of two and a maximum of six times. Between successive measurements 0.0025 inch of metal was removed. This was sufficient depth to bring up a new set of grains, since the average grain size of the samples used was approximately 0.001 inch.
- (3) Despite the precautions taken some scatter was still found in the measurements taken as above. In an effort to reduce this scatter the duplicate measurements on the samples aged at 1400° F were all carried out on the same surface of each sample. These measurements were made on this surface with 1 mm lateral shift in the holder between each run so that a new spot on the surface was covered by the X-ray beam each time. Two and usually three such measurements were made on each sample.

<u>Diffraction Line Width Studies:</u> Line intensity studies on the (111) line of the Low-Carbon N155 matrix with the Morelco Spectrometer revealed no evidences of line broadening at any stage of the aging process.

As further search for broadening effects a photographic back reflection technique was chosen in order to take advantage of the increased resolving power in the back reflection region. In addition, chromium radiation was used to eliminate, in part, fluorescence of the sample. Attention was centered on the highest order line of the austenite matrix obtainable with the above radiation, the 220 line (at a diffraction angle,  $\theta = 65^{\circ}$ ) resulting from the KK142 wavelength. In order to photographically record this line, the samples were mounted at the center of a 20-cm circular camera with the irradiated area being oscillated ±10° about an axis perpendicular to both the incoming X-ray beam and a diameter of the camera. Moll microphotometer recordings of the film showing the  $(220)\times_{1}\times_{2}$  doublet were then made and the widths of the  $\times_{1}$ line, determined at half peak intensity, by graphical means. To correct for the presence of  $\alpha_2$  line, the widths were actually measured as half widths on the side of the  $\alpha_1$  line away from the  $\alpha_2$  line, the division being a perpendicular through the apparent & peak. While more rigorous methods are available to correct for the presence of the & 2 line, it is felt that the differences between them and the method actually used are of second order and that the accuracy of the rest of the technique did not warrant such corrections. Care was taken to make certain that the density range of the ≤ 1 line recording lay within the linear density versus log intensity range of the film. The microphotometer data were also corrected to read intensity versus  $\Theta$  before broadening measurements were made.

Principally for use in future research, the line broadening data, obtained as above, were broken down to give the mean lattice deviation,

Ad, according to the method proposed by Haworth (see reference 3). In this connection, it should be mentioned that correction for broadening due to all sources other than lattice strain was done by the method originally put forth by Warren (see reference 4) using unaged solution-treated Low-Carbon N-155 as a standard. It should be noted then that the line broadening results are relative to the unaged material.

Lattice Parameters: Lattice parameters were measured with a Sachs and Weerts type camera using Copper  $K\alpha_1\alpha_2$  radiation. The samples were mounted with the prepared plane surface perpendicular to the incoming X-ray beam and rotated about an axis parallel with and slightly to one side of the axis of the X-ay beam. On the irradiated surface a light film of a mixture of mineral oil and chemically precipitated silver powder was placed. With the Cu  $K\alpha_1\alpha_2$  radiation used two lines, among others, were photographically recorded, the (420) lines of the austenite matrix of Low-Carbon N-155 (at  $\theta=75^{\circ}$ ) and the (553) lines of the silver (at  $\theta=78^{\circ}$ ). From the measured spacing of the silver (333) lines and using a lattice parameter of 4.0778 Å for the silver, the camera to film distance was calculated for each exposure. With this calculated distance (which averaged approximately 10 cm), and measurement of the (420) Low-Carbon N-155 line spacing, the lattice parameter

<sup>\*</sup>d represents the interplanar spacings of any given set of planes.

for each of the various aged samples of solution-treated Low-Carbon N155 was calculated. The absolute accuracy of these calculated parameters was not established, but the relative error was estimated to be  $\frac{1}{2}$ 0.0005 Å.

### Hardness Surveys

Hardness surveys were made on the various samples with a Brinell machine using a 10 mm ball and a 3000 Kg load. Two impressions were made on each sample, on planes which were originally transverse planes of the bar stock. Care was taken to stay within the fine grained area as mentioned under "Tost Materials". Two perpendicular diameters of each impression were measured, the resulting four readings averaged and converted to the Brinell Hardness Number.

### Creep Measurements

For the measurement of creep, 0.250 diameter specimens were prepared with the axis of the specimens corresponding to the original axis of the bar stock. For the majority of tests the gage section was 1-1/2 inches long terminated at each end with a 1/8 radius to the shoulders which in turn were 1/2 inch in diameter. Through the shoulders 0.100 diameter holes were diametrically drilled in which chromel pins were driven. A modified Martens extensometer was attached to these pins for measurement of the extension under load. The least reading of this extensometer and associated telescope and scale was 10<sup>-5</sup> inches.

For an additional check against the accuracy of extension measurements obtained in this manner a duplicate set of tests were run at the 60,000 psi stress level (see below) on samples aged at 1400° F using a different extensometer system. Here the samples had a gage length of 1 1/4 inches terminated at both ends with a 1/4 inch radius to 3/8 inch diameter shoulders threaded their entire length. Collars were then threaded onto these shoulders, down to the 1/4 inch radii, and locked in place with set screws. The extensometer system, in turn, was suspended from these collars with the use of pins mounted in same. Temperatures were controlled to ±5° throughout the tests and the temperature differences along the gage length held to ±3°. The furnaces used were electrical resistance type split along a transverse section at the center for ease of control of the uniformity of the gage length temperature. Load was applied to the specimen through a beam system with a mechanical advantage of 23.

In all tests the furnace was brought to the proper operating temperature before placing the specimen in it. The specimen was allowed to come to thermal equilibrium for an average period of 1-1/2 hours during which time minor controller and temperature uniformity adjustments were also made. At the expiration of the average period of 1-1/2 hours, the specimen was loaded. It was felt that in this way modification of the known initial structure of the specimens in the creep testing equipment prior to loading was held to a minimum. Thus the short-time creep characteristics of the known initial structures without appreciable modification by time at the test temperature.

Two stresses were used in creep testing, 30,000 psi and 60,000 psi, and one temperature 1200° F. 70,000 psi, was approximately the highest

sees possible without excessive plastic deformation upon loading. The higher stress was used to determine how the stress level of pertinal affected the conclusions drawn on the effects of aging on creep resistance at 30,000 psi.

The creep tests were run for an average of 50 hours provided fracture had not occurred. These tests were restricted to 50 hours in order to obtain the creep properties as characteristic as possible of the initial known structures and not the properties of the known initial structure plus modifications induced by time at the test temperature.

At the end of 50 hours all the tests covered herein had reached the so-called second stage of creep, with a reasonably steady creep rate or had fractured. The creep rates reported are either these second stage rates at 50 hours or the minimum rates occurring before fractures. It is obvious then that complete evaluation of decreasing secondary rates was not carried out. This question and also the one of structure alteration due to time at service temperature will be the subject of further research.

### Rupture Testing

Rupture testing was carried out in three units. The tests under stresses above 60,000 psi were run in a hydraulic tensile machine equipped with a transversely split electric resistance furnace. The load was held constant during the test to within ±1 percent with the rate of initial loading of the specimens approximately 50,000 psi per minute and comparable to the rate of loading of the more conventional rupture tests.

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as for the creep tests (see above). Specimens for these tests were obtained by longitudinal quartering of the original 7/8-inch square bar stock and using only those corners of the original bar which were uniformly fine grained (see Test Materials). Gage lengths 1-3/8 inches long by 0.250 inch in diameter, with the end terminated in 1/4-inch radii to 3/8-inch threads, were machined in these quarter sections.

Tests at 60,000 psi were simply the creep tests covered in the preceding sub-section carried to rupture.

Tests at stresses less than 60,000 psi were run in conventional beam loaded rupture units with one piece electric furnaces and with temperature control and uniformity comparable to the other rupture and creep tests. Specimen form was the same as that used for the rupture tests in the hydraulic tensile machine.

### RESULTS

### Metallographic Examination

Figures 5, 6 and 7 show the micrographs taken of the aged samples and the following description summarizes the results:

a. Aging at 1200° F resulted in little but the progressive development of a distinct etching resistant grain boundary constituent. At aging periods up to 10 hours, the boundary constituent was incomplete in that it did not surround all the individual grains. At aging periods of 1000 hours, the boundary constituents completely surrounded the grains and had become an approximately 0.5 micron-wide band. After aging 1000

hours slight precipitation was observable in the matrix near the grain boundaries. At 10,000 diameters, this precipitate did not appear to be a distinct phase with an interface but was surrounded by a concentration gradient as revealed by a sloping surface from the center of the precipitate particles to the matrix proper as a result of etching. As postulated by Naberro (see reference 5) this could indicate that appreciable strains would exist around each such perticle due to the probable difference in equilibrium lattice spacing between matrix and precipitate.

b. Aging at 1400° F resulted first in the progressive development of a grain boundary phase. Initially only a concentration gradient was present at the boundary, as revealed by a sloping surface towards the grain boundary after etching. At 10 hours the first separate boundary phase particles appeared. At 100 hours the boundary band was approximately 0.8 microns wide and changed little in character with further aging. In addition to the grain boundary reaction, general matrix precipitation appeared after aging about 10 hours, first along the grain boundaries, and increased rapidly in number and size up to the longest aging period used, 1000 hours. The particles first ing had concentration gradients surrounding them but for aging times of over approximately 100 hours, no appreciable concentration gradient appeared around the precipitate particles but rather a definite interfee. Average size of the particles at 1000 hours was estimated visually to be 0.2 microns equation the plane of polich and spaced an average of 1 micron apart.

c. Aging at 1600° F resulted in almost the same type of reactions as at 1400° F with the exception that they were accelerated.

The boundary phase, for example, appeared after aging for one hour.

The precipitate size at the end of 1000 hours age was estimated to average 0.7 microns same in the plane of polish and spaced an

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everage of 4 microns apart. Definite interfaces were present for each precipitate particle after aging times as short as 10 hours.

### X-Ray Studies

The following results were obtained:

a. Line Intensity Studies: Figure 8 and table 2 show the results of line intensity studies on the 10 hour solution-treated material when aged at 1200°, 1400° and 1500° F. Dehlinger and subsequent authors (see reference 6) have considered the effects that short and long period lattice distortions have on diffraction lines. In essence the conclusions are that short period disturbances (10<sup>-8</sup> to 10<sup>-6</sup> cm) result in reduction of line peak intensities without approximable broadening and that long period disturbances (10<sup>-5</sup> to 10<sup>-4</sup> cm) cause line broadening with, however, the integrated intensity studies shown on figure 2 can be interpreted, in the absence of line broadening data, an either short period disturbances or long period disturbances. If the latter were present, however, broadening should be the precominant effect.

The two minima for the material aged at 1.300° F on figure 8, lead to the possibility that two separate processes were observed at 1200° F. Aging at 1400° F or 1500° F apparently resulted in only one process occurring at each temperature which decreased line peak intensity.

It will be noted that the scatter for the measurements on material aged at 1400° F was approximately the same as for the measurements on material aged at 1200° F or 1600° F. When consideration is taken of the two methods of taking such data that were used, it is felt that the scatter was still due to non-random grain distribution despite rotation

of the samples. Before quantitative calculations can be made with such line intensity measurements, additional methods for alleviating non-random grain distribution will have to be developed. Only qualitative conclusions are thus drawn from the intensity data in this report.

b. Matrix Lattice Parameter Measurements: The measurement of lattice parameter as a function of aging time at a particular temperature can give direct evidence of whether the precipitate particles are still being nucleated, are growing by matrix depletion of the precipitate constituent atoms, or are growing by agglomeration. This usefulness of the parameter measurements is predicated upon the possibility of the average radii of the atoms making up the precipitate being somewhat larger or smaller than the average radii of the matrix atoms. If such is the case, then precipitation by matrix depletion will result in a measurable decrease or increase respectively in the lattice parameter. Nucleation can be ascertained if the lattice parameter remains constant for a period of time at a given temperature and then increases or decreases. Precipitate growth by agglomeration in turn can be noted when the parameter has reached a steady state value after increasing or decreasing from the initial value and yet the precipitate particles continue to grow as evidenced by netallographic examination.

The results of parameter measurements on the solution-treated and aged Low-Carbon N-155 are shown in figure 9. In no case has the aging time been sufficient to complete the precipitation reaction by matrix depletion, because the lattice parameters had not reached a steady state value. For this reason, the definite determination of whether the precipitate and matrix compositions are a function of temperature of aging and the character of such a function has not been determined.

It appears, however, that the precipitates obtained by aging at 1400° F and 1600° F could be slightly different in composition, since the curves for the matrix parameter were approaching steady state values which are probably not quite the same. Aging at 1200° F at the end of 1000 hours had resulted in so little change in lattice parameter that the only conclusion was that the major volume fraction of the material was never out of the nucleation stage. In addition a long nucleation period was shown by the lack of marked parameter change for aging up to approximately 100 hours at 1400° F followed by precipitate growth through matrix depletion. Only matrix depletion was found when aging was done at 1600° F.

c. Line Width Measurements: Since Dehlinger had postulated (see reference 6) that long period lattice distortion (of the type which could be associated with each of the small precipitated particles revealed by the metallographic examination of samples after prolonged aging) would result in line broadening, it was decided to measure the line broadening effects. Table 3 shows the results of width measurements, expressed as ratios between any given aged sample and unaged material as a standard of the (111) line at  $\theta = 22^{\circ}$ . These were obtained from the Norelco spectrometer plots. The little effect noted was due to lack of resolving power. Table 4 shows the line widths expressed as radians (column "B") oltained from photometer plots of film recordings of the (220) line, at  $\theta = 65^{\circ}$ . The increased resolution in the back reflection region revealed that appreciable broadening appeared only after long time ages at 1400° F and that a smaller degree of broadening occurred rather quickly when aging was done at 1600° F. Table 4 also summarizes the calculations involved in converting the line broadnesses to root mean square strains by the method of



Haworth (see reference 3). Figure 10 shows the results of such %1-culations. The results of course are qualitatively similar to the line widths recorded in table 4.

One further point is of importance here. The fact that the lattice parameter was decreasing with aging time makes it possible for a parameter distribution to be present as a result of concentration gradients being set up. Broadening of the diffraction lines will arise from this type of parameter distribution. Broadening could also arise from the elastic strains surrounding the precipitate particles, such plastic frames being atomic spacings in the two lattices. Thus the broadening data presented here can be ascribed to two sources and unfortunately enough data are not at present on hand to separate the two affects.

#### Hardness Measurements

The hardness survey was made to provide data for determining what internal condition gives high hardness. Figure 11 shows the results obtained on the solution-treated stock aged at 1800°, 1400° and 1600° F.

From figure 11, it can be seen that conventional aging behavior was apparently followed. The higher the aging temperature, the sooner the approach to a maximum hardness, the maximum hardness, however, increasing in value with decreases in aging temperature. This was certainly true for aging at 1400° F and 1600° F and probably true for aging at 1200° F.

Creep Properties of Solution-Treated and Aged Low-Carbon N-155

The purpose of the creep and rupture testing carriel out on solution-treated and eged Low-Carbon N-155 alloy was to measure the mechanical behavior of samples used in the physical measurements discussed previously. Figures 1. and 1), table 9 show the results of creep testing.

Aging at 1500° F for time periods up to approximately 10 hours resulted in 6 rather uniform increase in the secondary creep rate.

Aging at 1500° F for time periods up to approximately 10 hours resulted in little change in creep rate over the unaged material. For aging time periods over 10 hours the creep rate fluorensed rather rapidly to approach that for the material aged at 1600° F. From the two tests run after aging 100 and 1000 hours at 1200° F (100 hours being somewhat longer than the total period of testing) it appeared that aging at 1200° F had little effect on creep rate up to the longer aging period considered, 1000 hours, and then only served to reduce the creep resistance elightly.

Some antiquity was present in determining the creep rate vs. aging relationship for aging at 1400° F. However, in view of the magnitude of experimental errors (see below) involved, it is felt that the curve shown is a reasonable compression. In any event the trend that long time aging at 1400° F or 1500° F tends to reduce markedly the creep resistance was clearly shown.

At the stress level of 60,000 pri the affect of agins at either 1000° or 1600° F seemed qualitatively the same as at 50,000 psi except for the lower relative creep at anoth of the unaged material and the specimens aged one hour at 1400° F. It will be further noted that

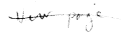
the results at 60,000 psi with the two types of extensometers agreed only within an average factor of 1.5. Previous experience with creep testing leads to the conclusion that creep rates are generally reproducible only within a factor of about the same magnitude.

Rupture Testing of Solution-Treated and Aged Low-Carbon N-155 Alley

Figures 14, 15 and 16, and table 8 show the results of the rupture testing on material aged at 1400° F. When aging was carried out at 1400° F, a gradual approach to a flat maximum in very short time rupture strength occurred with increased aging time. Subject to additional investigation, for rupture times up to 10 hours, it appeared that no appreciable alteration of the initial structure occurred by virtue of reactions at the test temperature, and such a result, as the above, was thus due to the initial structure of the material. With increased time for rupture (greater than 10 hours), the main alternation in the rupture time-aging time relationship appeared to be the marked improvement of the unaged and short time aged material in comparison to the long time aged material. (See figure 14). This most probably was due to reactions occurring in the material during testing and will be considered under "Discussion of Results." Inspection of figure 15 also shows that the aging periods at 1400° F for maximum short-time rupture strength were associated with the highest values of deformation prior to fracture. The initial deformation which occurred upon loading also increased slightly with increased aging time. Some alteration of these deformation characteristics was evident with increasing test time, because curves of figure 1/2 are in general sloping downward.

Again the experimental error involved in determining rupture times and deformation obscured to a certain extent the exact shape of the curves on figures 1h and 15. However, trends were clearly established. From figure 16, it is evident that upon aging, the node of failure changed from being primarily intergranular to being both intergranular and transfranular.

Figures 17, 13 and 1) show the results of repture testing on material aged at 1600° F. At this temperature a very broad maximum in the short time repture atrength occurred with using times longer than 0.5 hour (see figure 17). A very slight maximum possibly occurred at the one-half hour age. Again the progressive improvement of unaged stock was evident with increasing repture time. Figure 18 shows that maximum deformation values (i.e. the elongation at the fracture) for material aged at 1600° F followed the same pattern as for material aged at 1400° F, with, however, shorter aging times required for an equivalent effect (compare one-half hour aged curve of figure 18 with one hour aged curve of figure 14). Figure 19 shows that in no case were the predominately intergranular failures, found in the specimens aged a short time at 1400° F, observed in specimens aged at 1600° F even for as short a time as 0.5 hour.



DISCUSSION OF RESULTS

Fiffect of Aging on the Crystalline Structure

Upon comparing the line broadening, peak line intensity, lattice parameter, hardness and metallographic data, it can be seen that:

a. Relatively low (111) line intensity values resulting from the aging of solution-treated Low-Carbon N-155 at 1400° F for time periods

between approximately 0 and 30 hours were not accompanied by broadening of either the (111) or (220) lines. It can thus be concluded that the distortion causing the drop in peak intensity was of a short period nature. (See Experimental Procedures.) When cognizance is taken of the fact that the lattice parameter data indicated that pronounced rejection of either interstitial or large radius substitutional atoms did not take place until after 10 hours or so of acting, it can be post alated that only nucleation takes place during the first 10 to 30 hours at 1400° F and that this nucleation process produced short period strains. These time periods will not in general be exact since each process tends to overlap the next process to occur. The mushed must be rether small since larger ones would have a rather large system; and any strains associated would be long period once. Also, been presipitant nuclei would with a appreciate lattice for received. Longer aging resulted in line broadening strains and since this was associated with appreciable lattice contraction due to rejection of either interstitial or large radius substitutional atoms, these strains were nont probably associated with the actual precipitate particles. The metallographic data appear to hear these conclusions out since examination at magnifications up to 10000X revealed no evidence of precipitate particles until aging time periods at 1400° F were longer than 10 to 30 hours.

b. Relatively low (111) line intensities during aging at 1600° F for time periods up to 100 hours or so were associated with line broadening. This indicated long period strains. Further the parameter measurements indicated an immediate lattice contraction by rejection of the presipitant stons. Since, however, the metallographic data did not indicate that visible precipitate particles appeared much before 100 hours at 1600° F

had clasped, these large period strains were associated with large precipitant nuclei with a long spacing. The spacing of the stable nuclei was certainly larger when aging is carried on at 1600° F than when aging was done at 1.00° F as evidenced by the relative spacing of the precipitate particles which finally appeared at the two temperatures. That the stable nuclei were probably larger at 1600° F than at 1400° F is in agreement with Mehl and Jetter's summaries in regard to precipitation from solid soltuion (see reference 7).

of the very long period (approximately 1000 hours) during which no appreciable lattice contraction occurred and very little visible precipitate appeared, one can conclude that only short period nucleation occurred during the aging time studied. The changes in (111) line intensity were then associated with the short period strains of nucleation. The two separate minima found could resibly be associated with first, the matrix material weakly with the short period strains of nucleation.

The hardness data taken appeared to correlate quite generally with line broadening -- the greater the degree of line broadening the greater the hardness. Hardness, then, as far as the alloy studied is concerned, was associated with long period strains. In other words the long period strains associated with the precipitate particles forced after the nucleation period increased deformation resistance under the conditions of large straining present during a hardness test. These strains are due to the second localized deformation of the metal under the penetrator.

# 7; Factors Controlling Creen Strength

Comparison of the creep rates at 30,000 pci for materials aged results of at either 1200°, 1400° or 1600° F (see figure 12) with the above distribute measurements showed that the loss of initial creep strength at this stress level was most clearly associated with matrix depletion of the relatively large radius or the interstitial precipitant atoms. As covered above this was also associated with:

- a. Removal of the short period nucleation strains in the case of material aged at 1400° F.
- b. Development of visible precipitate and the continuous, relatively wide, grain boundary phase. These precipitant particles and the grain boundary phase were initially surrounded by concentration gradients which become progressively less steep with increased sging time (and/or increased aging temperature). (See figures 5, 6 and 7.)
- c. Broadening of the (220) line and hardening in the case of material aged at 1400° F and 1500° F. The point is not known for aging at 1200° F since it was not possible to obtain either appreciable hardening or broadening at this temperature, due to the excessive aging time required to reach high hardnesses.

Conversely, then, for materials aged at 1200° F, 1400° F or 1600° F, retention of creep strength was associated with:

- As solution-treated material or aged material still in the short period nucleation state.
- **b.** Relatively little or no visible general matrix precipitation and incomplete boundary reaction in the case of material aged at 1200° F and 1400° F. In all cases pertaining to material aged at 1400° F precipitate particles and the boundary phase, if present at all, had

concentration gradients surrounding them after the aging periods associated with high creep resistance.

case of unaged material or material aged at 1400° F and at 1200° F.

It would seem, then, that as far as creep resistance at 1200° F and at the 30,000 psi stress level was concerned, the removal of either the precipitant atoms from random solid solution or small nuclei made up of them by subsequent precipitate growth, led to a progressive lowering of the creep resistance. Further, the probable long-period strains associated with line broadening and relatively high hardness and resulting from large nuclei or precipitate particles and the creep strength. Rather, the continued removal of the atoms required to make up the precipitate which in turn caused the long period internal strain resulted in still lower creep strength. Assia It was the strain associated with the individual precipitant atoms, while still in random solid solution or in small nuclei, which controlled the creep strength at 1200° F and 30,000 psi stress level.

at 60,000 psi and 1200° F the effect of long-time aging was qualitatively the same as at 30,000 psi, but that the unaged and short-time aged at 1400° F material had relatively less creep resistance than the same material at 30,000 psi. It thus appeared that, while long-time aging and concomitant matrix depletion of the precipitant atoms resulted in lower creep strength, there was an optimum state of precipitate or nucleation and corresponding internal strain; for an optimum creep resistance. However, an additional factor applied here. As mentioned under "Results," the

relatively high stress of 60,000 psi resulted in fracture of all the creep specimens within a maximum of 32 hours. Figure 16 shows that the unaged and short-time aged at 1400° F material failed with general intergranular cracking. This formation of cracks resulted undoubtedly in greater creep rates than would be established with the same material with greater boundary strength. Hence, it has been felt that the general effects of matrix depletion of the precipitant atoms observed at a lower stress level still apply but that the unaged and short-time aged material has its creep resistance lowered, relatively speaking, at higher stresses by virtue of a weak grain boundary area. This point will be discussed further under "Factors Controlling Rupture Strength."

The fact that the line broadening strains, most probably associated with the individual precipitate particles, or large nuclei which resulted in high hardness did not have any effect on increasing creep resistance is quite surprising at first thought. However, since the short-period strains surrounding the individual precipitant atoms when in random or nearly random solid solution do give high creep resistance, it appears that the answer, as far as the allow considered here is concerned, lies in the periodicity and size of the two types of strained regions. From considerations covered below, the atoms most likely to be the source of the short-period strains, in the solution-treated material, are W, Mo, Cb, C and N. The total atomic fraction of these atoms is 0.042, or one atom is 24. (See table 5). In the absence of further data this can be interpreted to mean, that with one atom in 24, the center of a strained gion, the period of such strains is of the order of  $6 \times 10^{-7}$  cm. (See reference 10). Since interatomic forces of solids in general extend over only a few atoms, the size of the strained regions might be assumed

to be of the order of 1 x 10<sup>-7</sup> cm. This leaves a mean strain free path of the order of 5 x 10<sup>-7</sup> cm in the solution-treated material. In the aged material it seems plausible to assign the periodicity of the precipitate particles as the periodicity of the line broadening strains. As mentioned under "Results" this spacing was of the order of 10<sup>-4</sup> cm in the hardest sample prepared material aged 1000 hours at 1400° F.

It appears, unfortunately, that at present it is impossible to estimate closely the actual size, at any given aging time, of the strained areas, associated with each of the precipitant particles, or the average size of the precipitant atom free paths. This is because enough is not yet known to separate the line broadening due to elastic strains from the line broadening due to concentration gradients in the matrix. (See Results). However, it can be said that the mean precipitant atom free path was approaching 10<sup>-4</sup> cm as diffusion and precipitation approaches completion. This path was better than two orders of magnitude greater than the original mean precipitant atom free path.

The net result of aging, then, in solution-treated Low-Carbon N-155 alloy, was to probably replace a relatively short period small strain system containing a short mean strain free path (5 x 10<sup>-7</sup> cm) with another larger strain system of larger period and much longer mean strain free path (approaching 10<sup>-4</sup> cm). Since creep as here considered is essentially a small strain phenomena, the few slip systems in operation had a considerably larger probability of running into a strained area per unit shear strain in unaged material than in aged material with any appreciable amount of precipitate, despite the large average strains in the latter system. This means lower creep resistance in the aged

material. Hardness as commonly measured, however, is a large strain phenomena. The whole volume of the matrix is in a hardness test eventually filled with the many slip systems formed. The relatively large strains associated with the actual precipitate particles act very effectively to hinder the slip systems which eventually must form and try to pass through the large strain areas during a hardness test. Thus aged Low-Carbon N-155 is harder than unaged stock. A small strain hardness test would probably give a much more accurate indication of the relative creep resistance of aged alloys of the Low-Carbon N-155 type.

W, Mo, Cb can be considered one source of these short-period lattice strains since they are uniformly atoms of larger radii than the matrix and as mentioned above, aging in solution-treated Low-Carbon N-155 resulted in a decrease in lattice parameter -- presumably by rejection from solution of these large radii atoms. The matrix is assumed to be composed of Fe, Co, Ni and Cr, all atoms with approximately the same atomic radius (2.5 Å). (See table 5.) Carbon and nitrogen present interstitially also act to expand the lattice and their removal through precipitation would make the lattice contract. Thus these atoms must be considered along with W, Mo and Cb as sources of short period strains while in random solid solution. The fact that solute atoms of considerably smaller or larger radii than the solvent introduce strain into the system when in solid solution is well established generally -as witness the Hume-Rothery Rules for solid solution limitation. To quote another source, Sir Laurence Bragg, \* "Most engineering alloys of importance are the ones deriving their strength, at least in part, from the modulation of the lattice due to the presence of foreign atoms of

<sup>\*</sup>Lecture given at the University of Michigan, Fall, 1943.

different size." Further inspection of table 5 shows Mn and Si to be atoms of smaller radius than the average radius for N-155; and these also could be sources of short-period strains. Mn and Si are probably in the lattice substitutionally and would tend to make its parameter smaller than normal, and increasing with aging time if they were precipitating out. However, since the lattice become this indicated that probably either atoms larger than the atoma present interstitially or both ary chemical investigations indicated that only a phase or phases containing at most C, N, Mo, W, Cb, Cr and Fe was precipitating. Hence it is believed that Mn and Si play no part in the aging process, merely remaining in solution in the lattice throughout the aging process. If appears from this same preliminary chemical data that the C and N were partly in the form of an inert carbo-nitride of Cb in the solutiontreated material. The fraction thus present was unknown but even by disregarding entirely the C, N and Cb content (which is most probably not correct) the conclusions, as to the period of the strains involved are still warranted.

It would appear from the above that a logical way to increase the creep strength of Low-Carbon N-155 would be to add elements with either larger or smaller atomic radii than Mo, Cb and C and N respectively. Those suggested in table 6 have larger atomic radii. Only boron, among elements of small atomic radiue, appears promising at the moment. The usefulness of boron seems to be substantiated by the high creep strength boron modifications of Low-Carbon N-155 recently developed by the Union Carbido and Carbon Research Laboratories,

Table 7 lists elements which have atomic radii similar to Mo, W

and Cb and which might be expected to act as substitutes. The elements suggested in these tables are not the only metallic elements with the desired atomic size but are thought to be the most promising. With the exception of Al and Ag, all the proposed additions or substitutions are transition elements. Some evidence (see reference 11) is available which indicates that such transition elements have abnormally high binding energies. Al and Ag are considered only on the basis of having the desired apparent atomic size.

It is of course desirable that the materials suggested in tables 6 and 7 go readily into solution at some relatively high (solution-treating) temperature, and either stay in solution (or the nucleated state) or precipitate very slowly at the lower temperatures of service. It is possible that any or all of the proposed elements may show an increased tendency to come out of supersaturated solid solution and thus cause rapid loss of creep strength or show the wrong or a poorer type of solubility versus temperature characteristic. Other metallurgical characteristics (i.e., ductility) must necessarily be satisfied before such modifications could be considered satisfactory.

## 3 . Factors Controlling Rupture Strengths and Ductility

Inspection of figures 14 and 17 shows in general that aging at either 1400° F or 1600° F resulted in a progressive increase in very short-time rupture strength. For a given increase in rupture strength less aging time was required at 1600° than at 1400° F. For somewhat longer rupture times, the relative increase in the rupture strength of the unaged material was quite striking. In fact, the solution-treated stock rapidly became equal in strength to the aged material at the

increased rupture times. Previous experience has indicated that the rupture strength of the unaged material will actually exceed the rupture strength of the aged material when considering longer rupture times than were used in this investigation. Further, the continuous rupture times at each stress on figures 14 and 17 were approximately the same for aging at either 1400° F or 1600° F. Examination of the other physical measurements shows that this general increase in very short time rupture strength with increasing aging time was associated with:

- a. In the case of material aged at 1400° F passage through the nucleation stage prior to precipitation as revealed by the line intensity studies.
- b. Removal of relatively large or small radii atoms through precipitate growth after nucleation as revealed by lattice parameter measurements.
- c. Progressive formation of a rather wide, continuous band of a separate grain boundary phase and with the formation of the visible precipitate particles as revealed by micrographic examination.
- d. Hardening of the material through the formation of strains which resulted in diffraction line broadening in the case of aging at 1400° F and initial hardening and then progressive softening in the case of aging at 1600° F.
- e. Increased ductility of the specimens. One way to consider this effect is to express the maximum true axial strain at fracture based upon the initial and final cross-sectional areas at the fracture and the assumption of constancy of volume. Figures 15 and 18 present the results of these calculations. The effect of the decrease in cross-sectional area at the fracture with aging time was to increase the true stress under

which the test runs during the duration of the test. In general, then, the rupture tests considered were not constant stress tests, but rether tests with true stress increasing with time along some quantitatively unknown path in a time-stress coordinate system. It is known from the timeelongation curves for the tests at 60,000 psi that reduction of area was gradual and thus the true stress increased rather gradually throughout the test, the final value of course being greater for smaller final cross-sectional area. Inspection of the geometry of ruptured specimens showed that the cross-section was reduced gradually along the axis toward the fracture. From this, it can be concluded that the degree of triaxiality of the stress system at the fracture was low, as considered in the calculations of Fridgeman (see reference 8). Hence, the stress system can at least be considered uniaxial for the rupture tests covered herein. Lastly figures 15 and 18 also showed that the maximum true strain at fracture, or ductility, for any given class of specimen, decreased with increasing rupture time.

At the onset, the fact that ordinary rupture tests are not constant stress tests makes outright analysis of the factors controlling rupture strength difficult. However, three factors stand out quite clearly:

First, figures 16 and 19 showed that the relative initial weakness of the unaged materials was associated with weak grain boundary areas and consequent intergranular crack formation, especially on grain boundaries normal to the stress axis. Further inspection (see figures 16 and 19) showed that this tendency for intergranular cracking was progressively removed with either increased aging time at 1400° F or 1500° F prior to testing, or with decreased stress on the unaged material and thus increased time at the test temperature of 1200° F. This apparently was

due to the formation of the grain boundary phase either prior to or during testing (see figures 6 and 7). Thus unaged material became stronger relatively speaking with increased rupture time, and the aged material also became relatively stronger with increased prior aging time -- more slowly, however, with prior aging time at 1400° F than for 1600° F aging since the boundary phase was formed more slowly at 1400° F. It is then quite interesting to conclude that the matrix-boundary phase binding was stronger than at least certain oriented matrix-matrix-bindings.

Second, inspection of figures 14 and 17 showed that once a definite almost continuous boundary phase was formed, the rupture strengths were roughly independent of aging temperature or aging time. This indicated that the initiation and propagation of the predominantly transgranular cracks was independent of the differences in structure arising from differences in aging at 1400° or 1600° F. Aside from minor maxima or minima, which are probably just inside or outside the experimental errors involved in determining the rupture times, the differences in lattice a consideration depletion of the large radii atoms, and the differences in magnitude and time of occurrence of maximum hardening with difference in aging temperature, had little effect. This conclusion is based, however, in part upon the fact that the degree of elongation and cross-sectional area reduction, once the grain boundary phase was formed, was approximately samples having the same rupture time as the result of again at the same for facing equivalent times at either 1100° or 1600° F. Thus the degree of deformation was also constant and did not effect to a first approximation the fracturing characteristics of the two classes of aged materials differently. It is well known, however, that, within limits, deformation in itself generally raises the resistance of rupture (see reference 9). Since this strain strengthening occurs in connection with

increase in the true stress, evaluation of either of these two opposite effects is difficult along with evaluation of such things as the effect of progressive lattice depletion in general on resistance to crack propagation. Thus no further conclusions in regard to possible masked general structure factors arising with long time aging at 1400° F or 1600° F are drawn at this time.

Third, the plastic strain, before rupture failure occurred, increased markedly with long aging times at either 1400° F or 1600° F.

This of course is a direct result of the fact that the creep resistance decreased markedly with long aging times at either aging temperature and that, the resistance to crack propagation on the other hand is at a maximum. Figures of and of also show, however, that the plastic strain before fracture decreased with increasing time for rupture. No obvious reason for this presented itself during the investigation covered herein and thus this phenomenon will be subject to future research.

Limitations of The Results

There are obviously mamerous important aspects of the problem of high temperature strengths which have not been adequately covered by this report. Of primary importance is the fact that the structural measurements have not been correlated with long time creep and rupture atrengths at 1200° F or with any time period at other temperatures. Furthermore, only one alroy has been considered and this alloy was initially given an impractically long solution-treatment in the interest of simplifying the problem. In addition, certain refinements in X-ray techniques seem desirable. Information is also needed regarding the effect of time and deformation during testing to assess the reliability of the assumption that such changes had little effect on known initial structure. And finally, it would be especially desirable to know the

# Limitations of the Results

This report is limited to the presentation of a method of determining the fundamental mechanisms by which processing, heat-treatment, and chemical composition control the properties of alloys at high temperatures. A relatively limited amount of data for solution-treated and aged Low-Carbon N-155 alloy has been obtained and interpreted in terms of the proposed method. The resulting theories require extension and improvement from similar investigations on many alloys, as well as for more test conditions on Low-Carbon N-155 alloy (see below). It is believed, however, that the approach to the problem is reasonably sound and that the limiting factor to a general theory of the metall-urgical factors controlling high temperature strength of alloys is the large volume of testing required and development of suitable experimental techniques.

In regard to Low-Carbon N-155 alloy, there are obviously numerous important aspects of the problem which have not been adequately covered by this report. Of primary importance is the fact that the structural measurements have not been correlated with long time creep and rupture strengths at 1200° F or with any time period at other temperatures. Certain refinements in x-ray techniques seem desirable. Information is also needed regarding the effect of time and deformation during testing to assess the reliability of the assumption that such changes had little effect on known initial structure. And finally, it would be especially desirable to know the exact composition of the precipitating phases. The amount of time required for development of suitable techniques and the volume of experimental work been the limiting factor to date. Work is now in progress to cover most of the above items.

## CONCLUSIONS

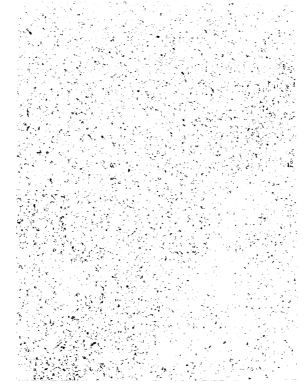
- 1. An experimental procedure is described which is believed suitable for establishing the fundamental mechanisms by which processing, heat-treatment, and chemical composition control the properties of alloys at high temperatures. This method relates microstructures and x-ray diffraction characteristics after various prior treatments, to creep and rupture test properties.
- 2. Application of this method to solution-treated and aged Low-Carbon N-155 alloy and correlation with the short time creep and rupture characteristics at 1200° F indicated the following possible fundamental explanations for the effect of aging on its 1200° F properties.
  - a. Aging of solution-treated Low Carbon N-155 resulted in progressive lowering of the initial (short time) creep resistance through the removal from solid solution of large radii or interstitial atoms by precipitation. No optimum precipitate dispersion or state occurred for optimum creep resistance, rather the alloy can be considered as obtaining its optimum creep strength through "modulation" of the lattice with the large or small precipitate atoms when they are in the random, or at most nucleated, distribution of the solution-treated state. b. Short time aging of solution-treated Low-Carbon N-155 apparently resulted in a marked increase in short time rupture strengths over that for unaged material, through the growth of a grain boundary phase, this phase acting to strengthen the boundary areas to eliminate intergranular cracking and consequent low resistance to crack propagation. Long time

aging resulted in little further change in short time rupture strength, and longer time service at 1200° F was sufficient to develop a grain boundary phase in the unaged material to thus raise its rupture strength to compare favorably with the strengths of prior aged material.

- c. Because the effect of aging in general was to lower the creep resistance and to raise the rupture strength, for the time periods consideration, aging resulted in a material which exhibited greater ductility before fracture.
- d. It is indicated that alloys might be developed with strength comparable to N-155 with other alloying elements, or an alloy of the same general type with improved creep strength might be developed by replacement or supplement of the large or small radius elements present in Low-Carbon N-155. Elements of the same atomic radius as No, Cb, or W, which could possibly act as substitutes or supplements, include Al, Ag and Ta and elements of larger atomic radius suitable for additional alloying include Zr, Ce and Ti. Boron appears to be the only promising alloying element of small atomic radius not at present used in N-155.
- 3. It is emphasized that the explanations for the effect of aging on the properties of Low-Carbon N-155 alloy at 1200° F apply at present, only to that alloy and are not to be taken as general. It is entirely possible that this alloy will prove to be an unusual one exhibiting behavior which is the exception to some general rule to be established as a result of further investigation on other alloys.

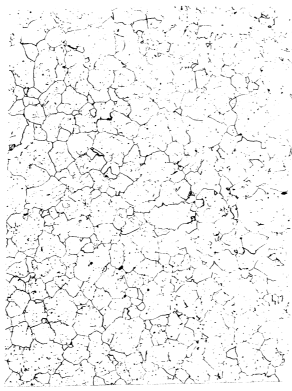
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48-201

Heat A-1726 As-Rolled



48.202

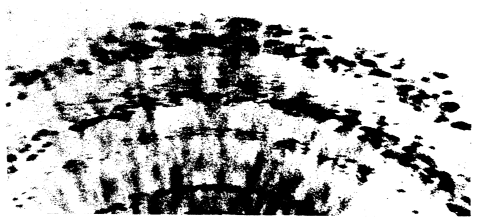
Small grained area of Low-Carbon N-155, solution-treated 10 hours at 2200° F, water-quenched, and aged 1 hour at 1400° F

Cross sections of bar X100 Electrolytically etched in 10 percent chromic acid

Figure 1. - Typical microstructures of Heat A-1726.



Pattern from serface ground on medium coarse rinder ern finished with No. 1 enery camer



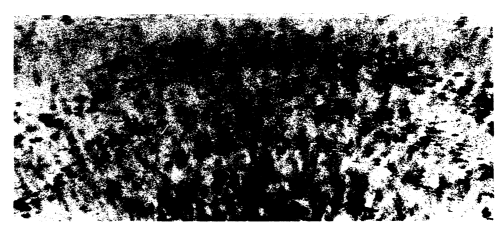
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After resonal of 0.010 inch

A 2015 Unfiltered to radiation (~5KV)

Figure 2. - Diffraction patterns showing removal by electrolytic policies on low-Carbon L+185 allegt solicing treated to notes at 2200° F and water-quenched.



After removal of 0.017 inch

A 2016

Figure 2. - Concluded. Diffraction patterns showing removal by electrolytic polishing of cold-worked surface on Low-Carbon N-155 alloy, solution-treated 10 hours at 2200° F and water-quenched.

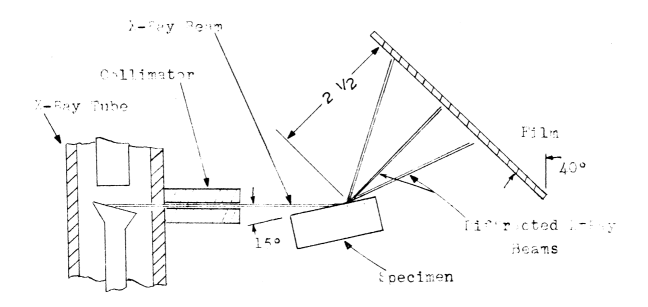


FIGURE 3. SECTIONAL SCHEMATIC OF EXPERIMENTAL SETUP
FOR STUDYING SURFACES OF POLISHED SAMPLES.



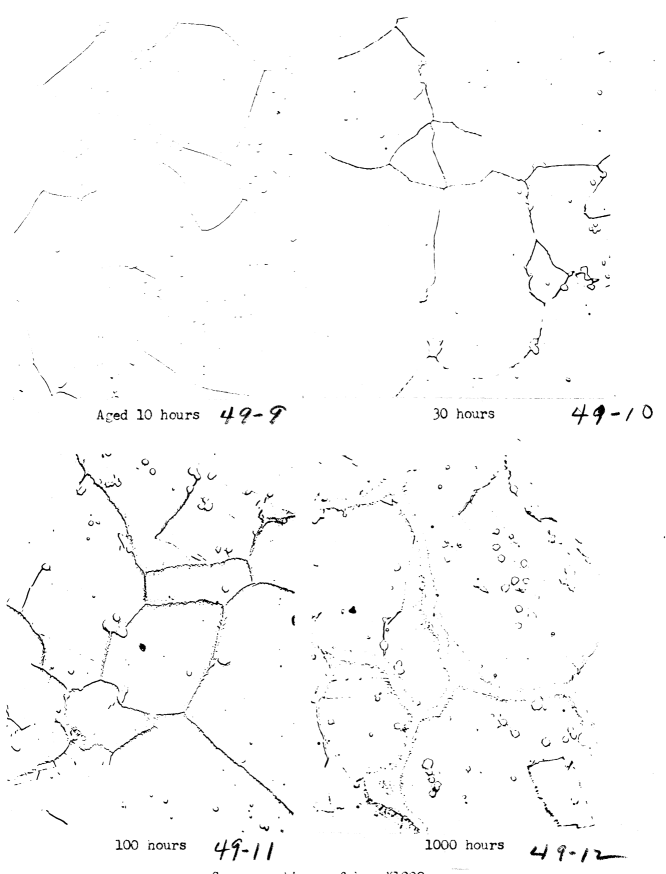
A 2018

Figure b. - Electrolytic cell for gross metal removal x 1/2.



Cross sections of bar X1000 Electrolytically etched in 10 percent chromic acid

Figure 5. - Effect of aging at 1200° F on the microstructure of Low-Carbon N-155 alloy, solution-treated 10 hours at 2200° F and water-quenched.



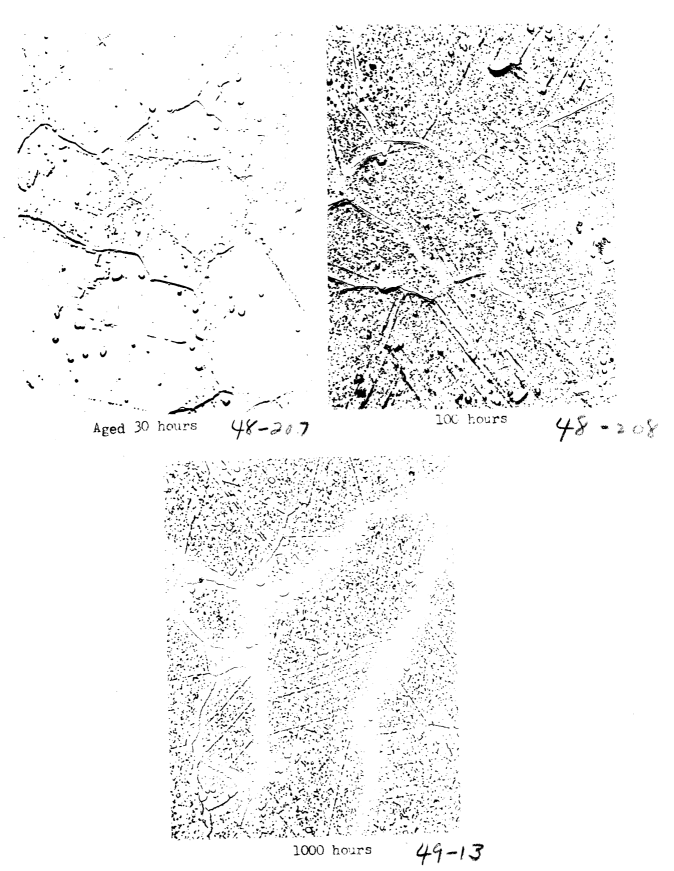
Cross sections of bar X1000 Electrolytically etched in 10 percent chromic acid

Figure 5. - Concluded. Effect of aging at 1200° F on the microstructure of Low-Carbon N-155 alloy, solution-treated 10 hours at 2200° F and water-quenched.



Cross section of bar X1000 illectrolytically otched in 10 percent caronic acid

Figure 6. - effect of aging at 1500° F or the eferostructure of Low-Carbon N-155 allow, solution-treated 10 hours at 2200° F and water-quenched.



Cross section of bar X1000 Electrolytically etched in 10 percent chromic acid

Figure 6. - Continued. Effect of aging at 1h00° F on the microstructure of Low-Carbon N-155 alloy, solution-treated 10 hours at 2200° F and water-quenched.

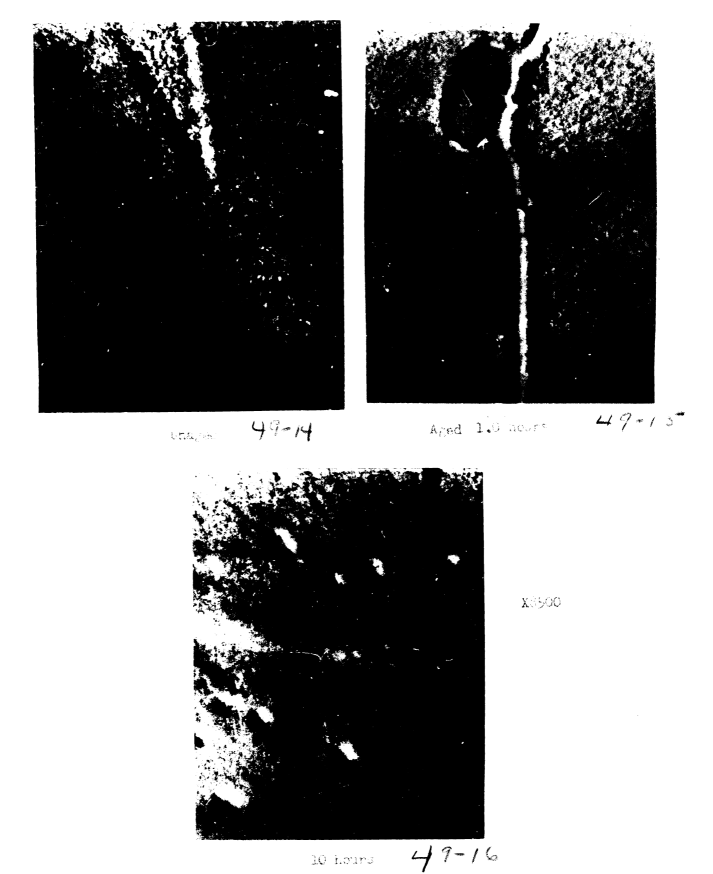


Figure 6. - Continued. Electron microsraphs of replicas prepared from Lew-Carbon 1-155 alloy, solution-traded 15 found at 2200° F, water-quencusa and mean at 1100° r.

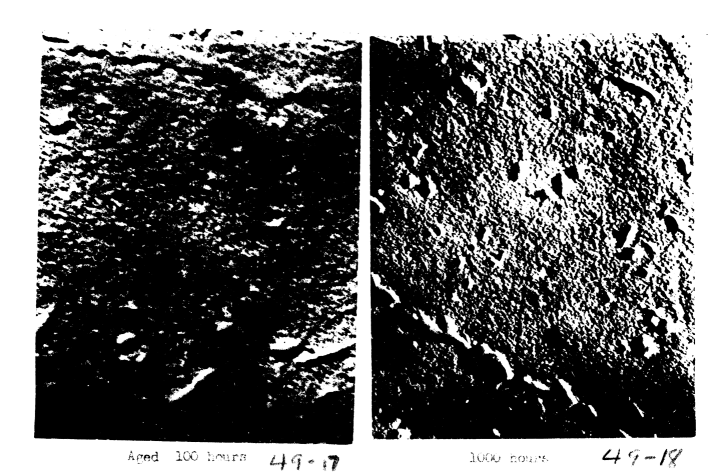
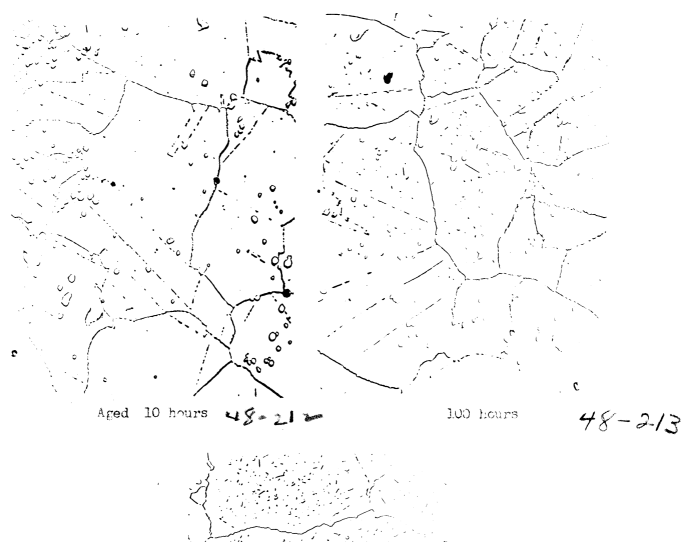


Figure 6 - Concluded. Electron micrographs of replicas prepared from Low-Carbon A-155 alloy, solution-treated 10 hours at 1200° F, was exequenched and good at 1500° F.



Cross section of bar X1000 Electrolytically etched in 10 percent chromic acid

Figure 7. - Effect of aging at 1600° F on the microstructure of Low-Carbon N-155 alloy, solution-treated 10 hours at 2200° F and water-quenched.

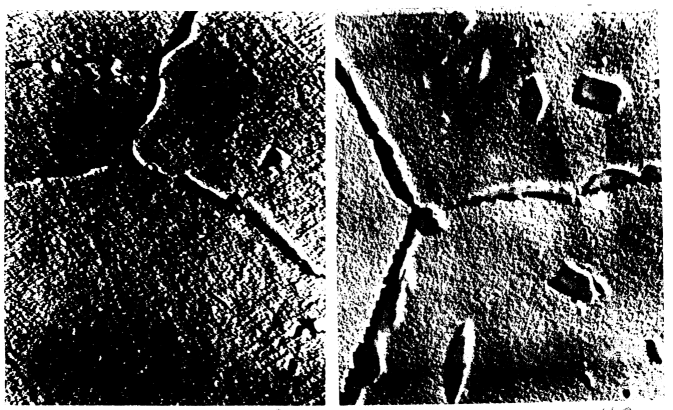




Cross section of Par X1000 Electrolytically etched in 10 percent chromic soid

Figure 7. - Continued. Effect of aging at 1600° F on the digrostructure of Low-Carbon N-155 alloy, solution-treated 10 hours at 2200° F and water-quenched.

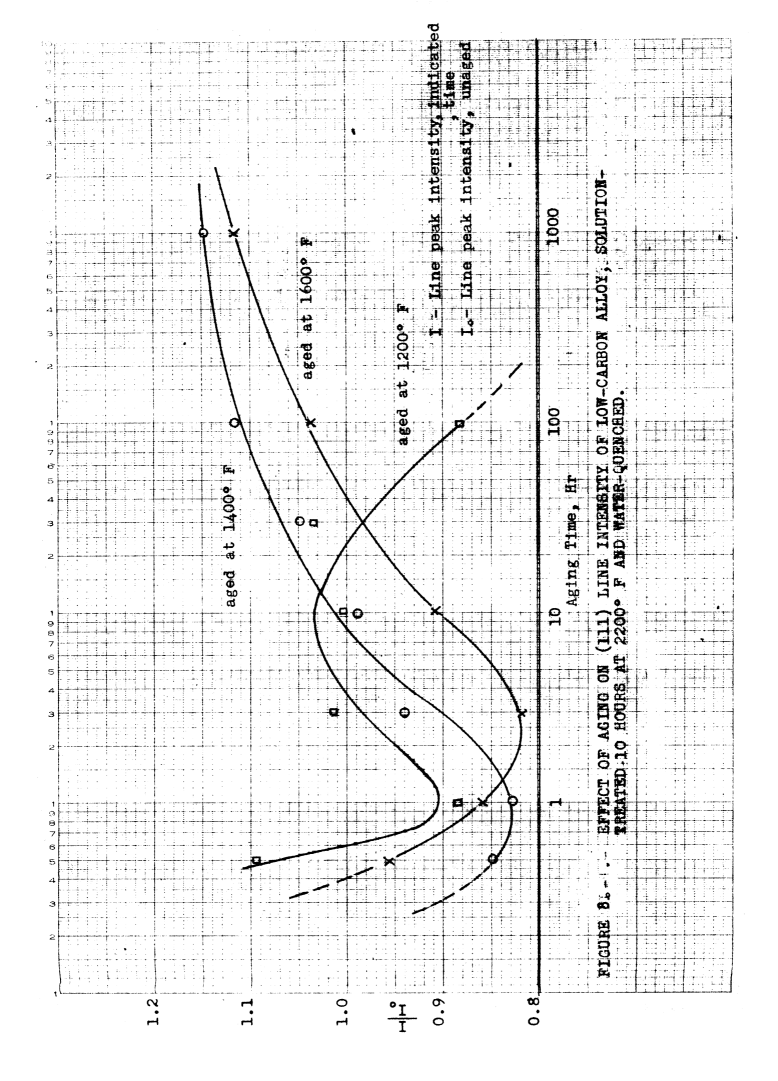


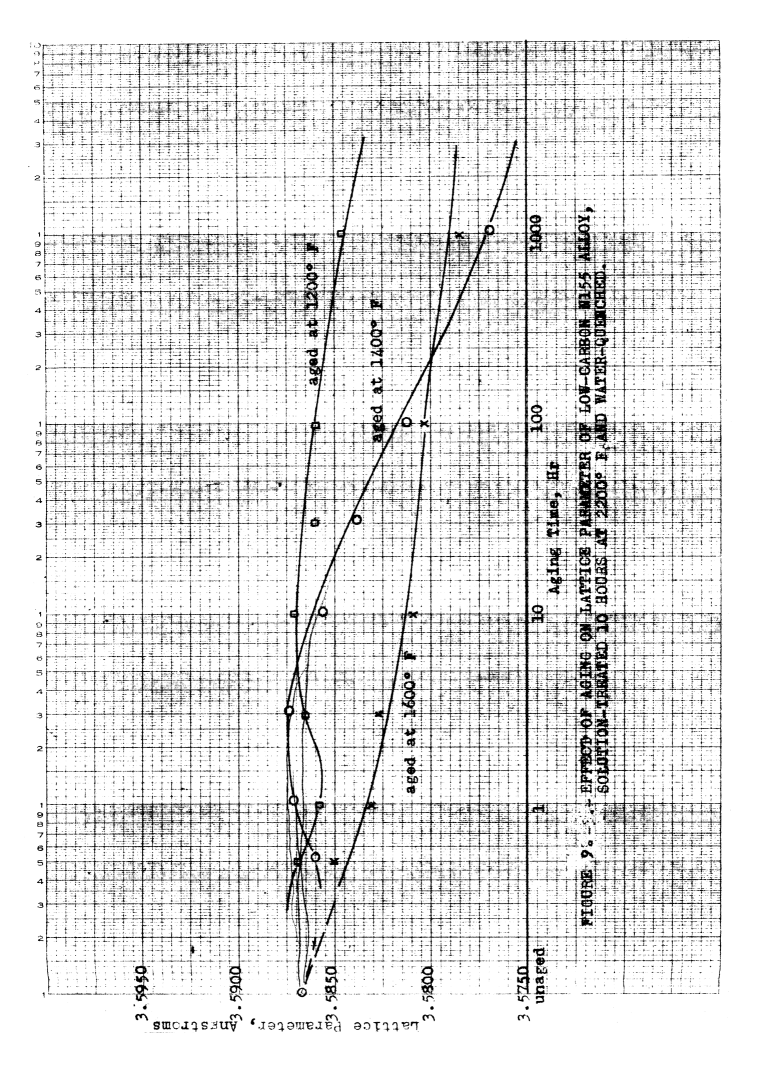


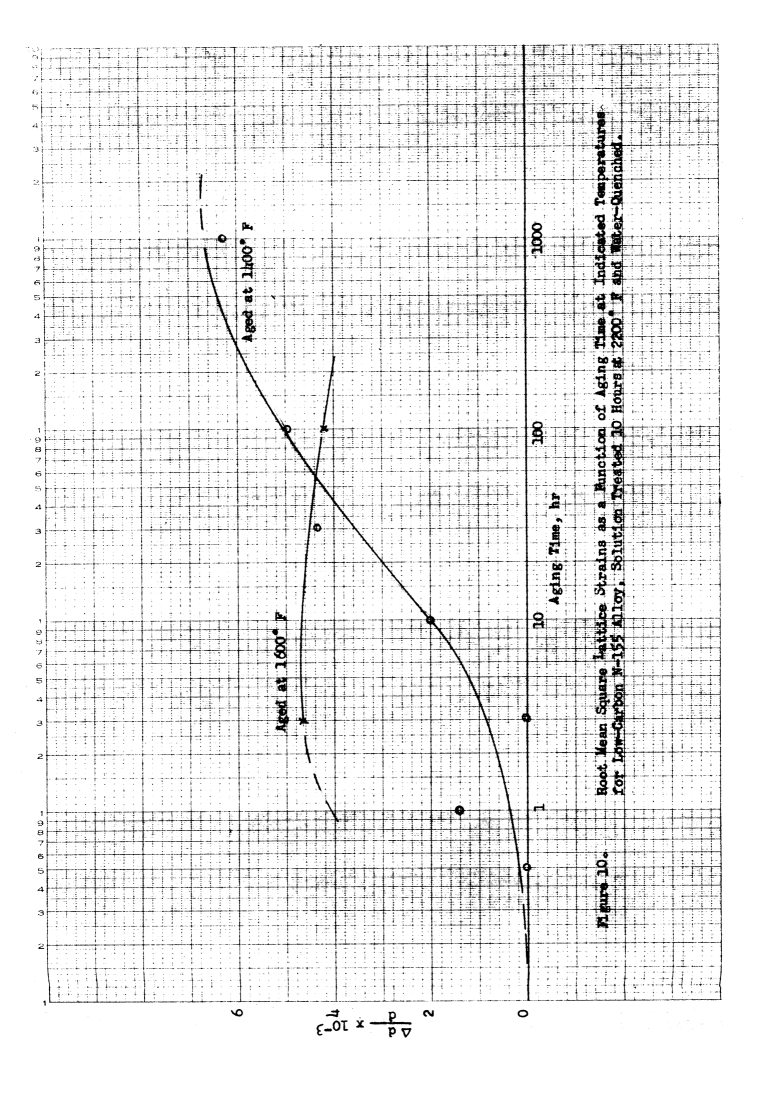
100 leves 49-2 5050c.

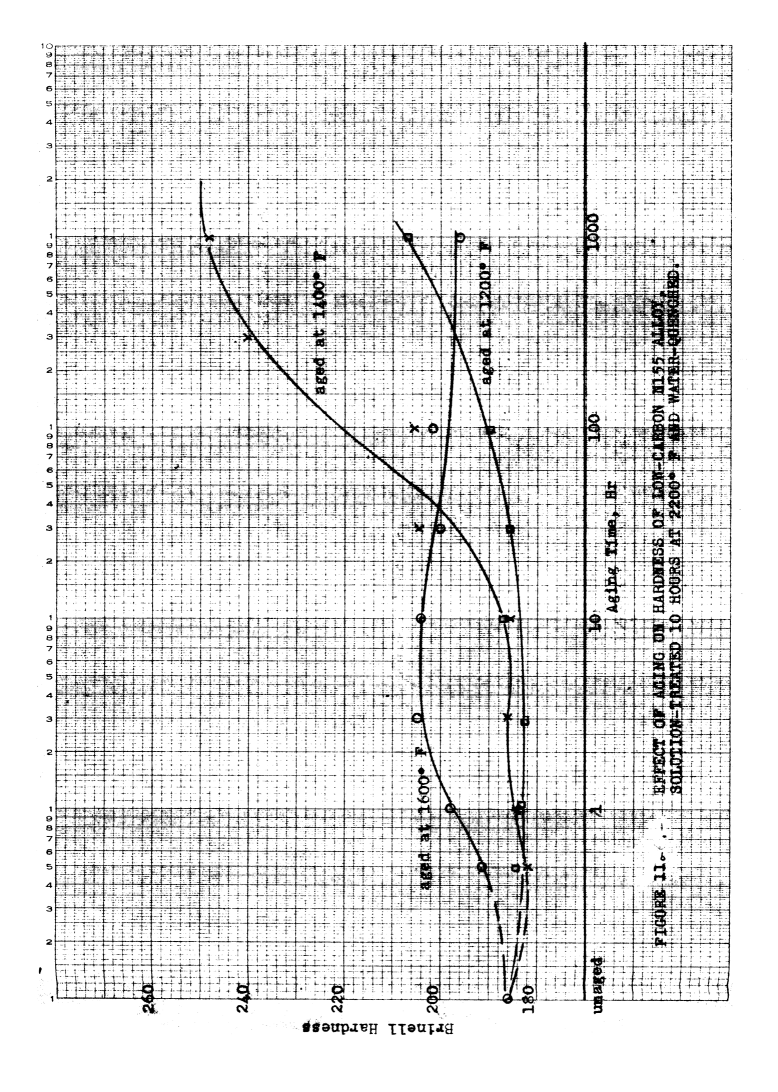
1000 hours 49-23

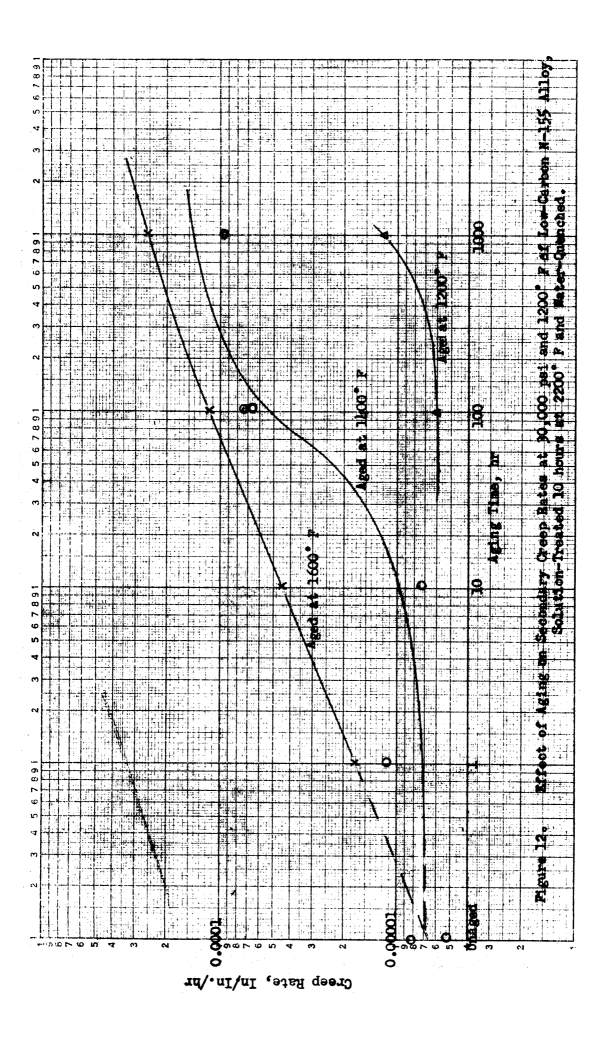
They was fire a forestructure that they returned for a solution of the companies of the forest of the companies of the compan

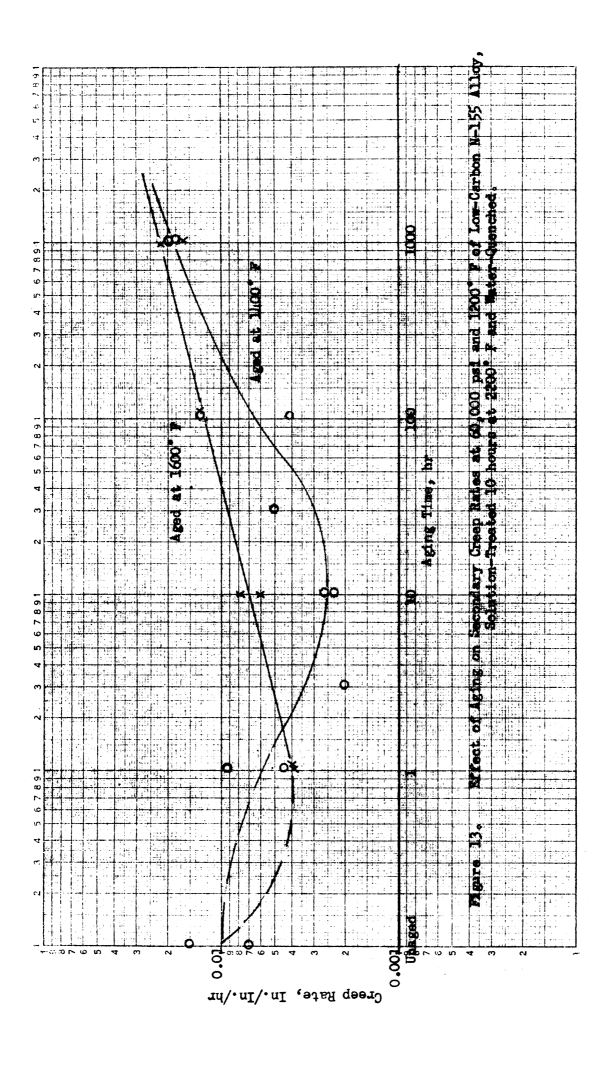


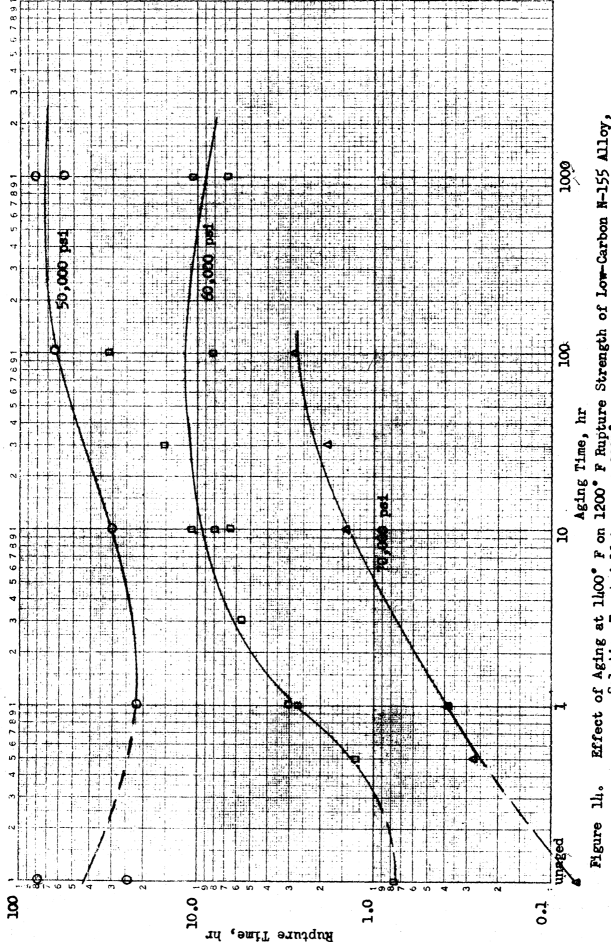




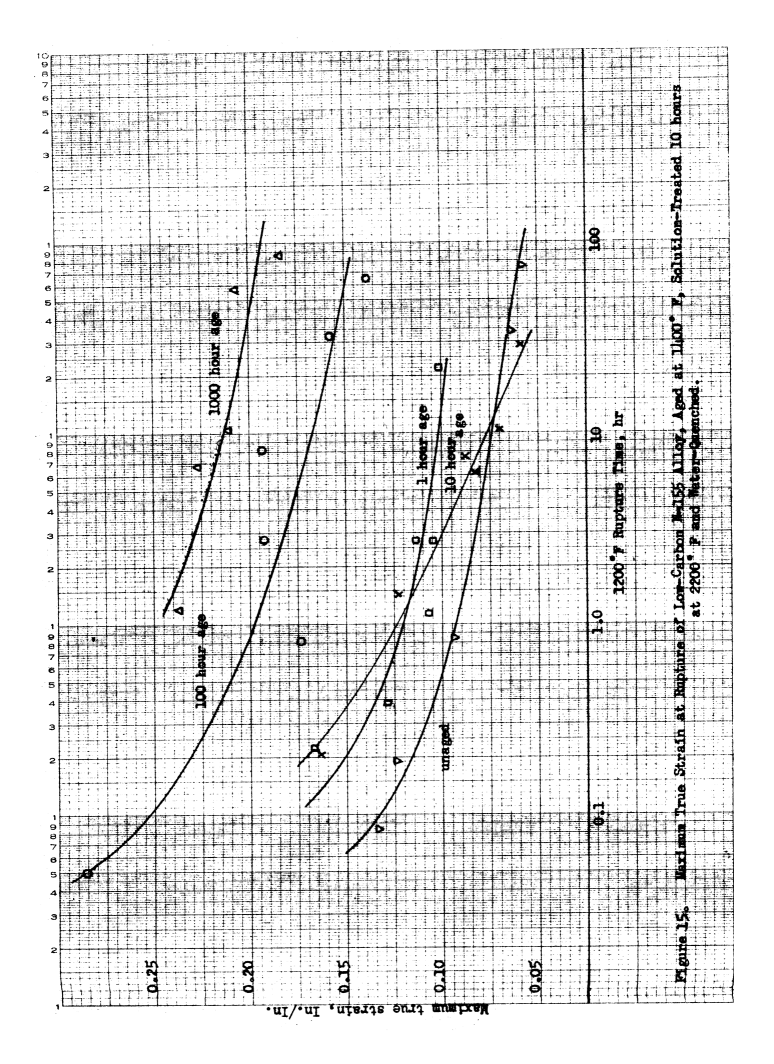






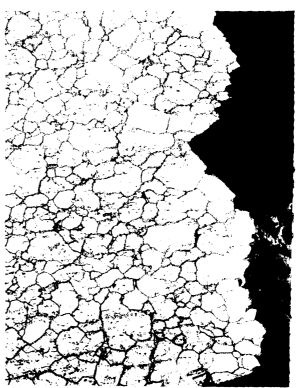


Effect of Aging at 1400° F on 1200° F Rupture Strength of Low-Carbon M-155 Alloy, Solution-Treated 10 hours at 2200° F and Water-Quenched.





Unaged: failed in 0.083 hours under a stress of 70,000 psi 49.24



Unaged: failed in 34 hours under a stress of 10,000 pst



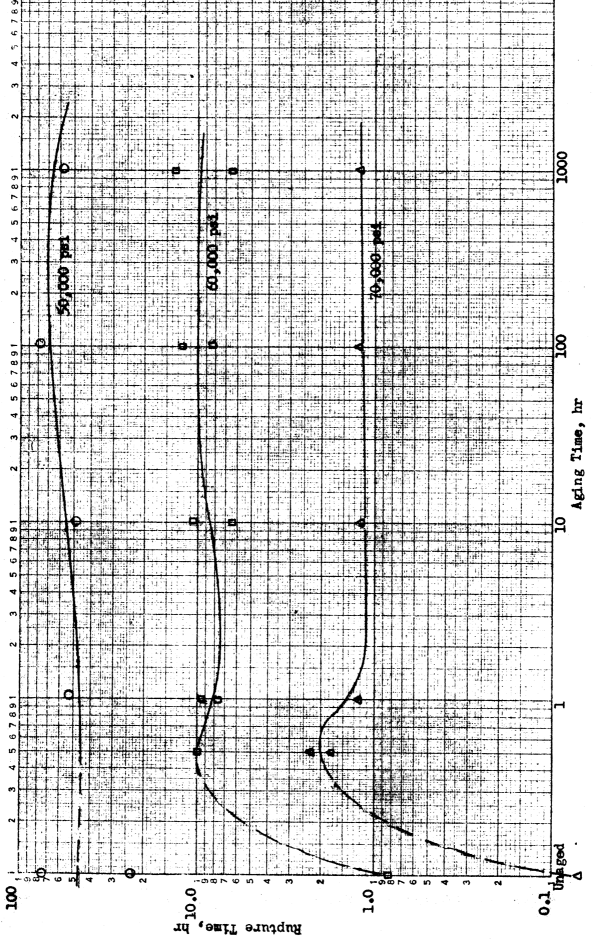


Aged 1 hour: failed in 0.22 hours under a stress of 75,000 psi 49-26

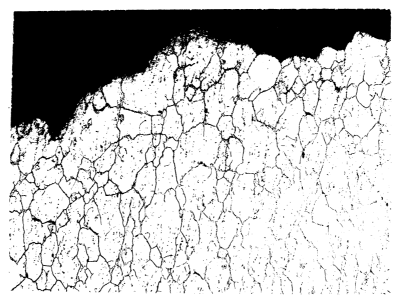


Aged 100 hours: failed in 0.05 hours prior a stress of 80,000 psi 49.27

Picure 16. - iffect of aging at 1000° F or freeture characteristics at 1200° F of Lov-Carbon N-150 allow, solution-treated 10 hours at 2200° F and water-quenched.



Effect of Aging at 1600° F on 1200° F Rupture Strength of Low-Carbon N-155 Alloy, Solution-Treated 10 hours at 2200° F and Water-Quenched. Figure 17.



Aged 0.5 hours: failed in 1.31 hours under a stress of 70,000 psi

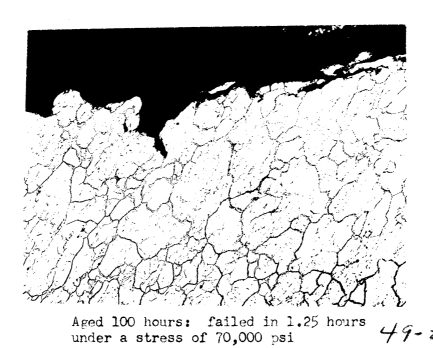


Figure 19. - Effect of aging at 1600° F on fracture characteristics at 1200° F of Low-Carbon N-155 alloy, solution-treated 10 hours at 2200° F and water-quenched.

## TABLE 1

# PROCESSING OF LOW-CARBON N155 7/8-INCH BROKEN CORNER SQUARE BAR STOCK FROM HEAT A-1726.

(Reported by the Universal-Cyclops Steel Corporation)

An ingot was hammer cogged and then rolled to bar stock under the following conditions:

- 1. Hammer cogged to 13-inch square
   Furnace temperature 2210° 2220° F
   Three heats Starting temperature on die 2050° 2070° F
   Finish temperature on die 1830° 1870° F
- 2. Hammer cogged to 10-3/L-inch square
  Furnace temperature 2200° 2220° F
  Three heats Starting temperature on die 2050° 2070° F
  Finish temperature on die 1790° 1800° F
- 3. Habmer cogged to 7-inch square
  Furnace temperature 2200° 2220° F
  Three heats Starting temperature on die 2050° 2070° F
  Finish temperature on die 1790° 1890° F

Billets ground to remove surface defects.

h. Hammer cogged to h-inch square
Furnace temperature 2190° - 2210° F
Three heats - Starting temperature on die 20h0° - 2060° F
Finish temperature on die 1680° - 1880° F

Billets ground to remove surface defects.

5. Hammer cogged to 2-inch square
Furnace temperature 2180° - 2210° F
Three heats - Starting temperature on die 2050° - 2065° F
Finish temperature on die 1730° - 1870° F

Billets ground to remove surface defects.

- 6. Rolled from 2-inch square to 7/8-inch broken corner square one heat Furnace temperature 2100° 2110° F
  Bar temperature start of rolling 2050° 2060° F
  Bar temperature finish of rolling 1910° F
- 7. Bars are numbered 1 through 56, Number 1 bar represents the extreme bottom of ingot and Number 56 the extreme top position.
  - All billets were kept in number sequence throughout all processing, so that ingot position of any bar can be determined by its number.
- 8. All bars were cooled on the bed and no anneal or stress relief was applied after rolling.

TABLE 3

Effect of Aging on Relf Value Width of (111) Diffraction Line of Low-Carbon Ntip Alloy Schution-Treated 10 Hours at 2200°F and Water Quenched

(Copper  $Ka_1a_2$  Radiation)

Aging Temperature (°F)	Aging Tire (bour)	Line Width W/Wo*	Averaga	Mean Deviation
1400	1.0	0.97 1.04 1.01	1.01	+0.02
	3.0	1.04	1.00	<u>+</u> 0.02
	10.0	0.99 0.95	0.98	<u>+</u> 0.02
	30.0	1.06 0.97 1.06	1.03	<u>+</u> 0.04
	100.0	1.00 0.91 0.99	0.97	<u>+</u> 0.04
1600	0.5	0.95 1.04	0.99	<u>+</u> 0.04
	1.0	1.03 1.02	1.02	<u>+</u> 0.02
	3.0	1.03 1.15 1.04	1.07	<u>+</u> 0.05
	10.0	1.00 1.01 1.00	1.00	<u>+</u> 0.0
	100.0	1.00 1.02 0.91	0.96	<u>+</u> 0.05

<sup>\*</sup>W - half value width of (111) line of specimen aged at indicated time and temperature.

 $W_{\rm O}$  - half value width of (111) line of unaged material.

TABLE 4

Internal Root Mean Square Strains for Low-Carbon N155
Solution-Treated 10 Hours, 2200°F, Water-Quenched and
Aged as Indicated

Aging Temp.	Aging Time Hrs	Lettice Parameter	dazol	cot 9	K	B (radians)	2 <u>∆d</u> d
1400°F	<b>0.</b> 5	3.5862 A	1.27 A.	0.4843	6.15x10 <sup>-4</sup>	0	0
	1.0	3.5874	1.27	.4843	6.15	1.47x10 <sup>-3</sup>	1.40x10 <sup>-3</sup>
	3.0	3.5876	1.270	.4843	6.15	0	o
	10.0	3.5858	1.269	.4843	6.15	2.10	2.0
	30.0	3.5840	1.267	.4830	6.14	4.53	4.31
	100.0	3.5814	1.266	.4820	6.12	5 <b>.1</b> 8	4.92
	1000.0	3.5770	1.0803	.6724 <sup>3</sup>	7.26 <sup>3</sup>	5.55 <sup>3</sup>	6.25
1600°F	3.0	3.583	1.0823	.6851 <sup>3</sup>	7.413	4.10 <sup>3</sup>	4.70
	100.0	3.581	1.081 <sup>3</sup>	.6847 <sup>3</sup>	7.403	3.66 <sup>3</sup>	4.20

For-Low Carbon N155, Solution-Treated 1 Hour, 2200°F, Water-Quenched and Rolled to 15% Red. in Cross Section at 1400°F

3.587	1.27 Å	.4843	6.15x10 <sup>-4</sup>	11.5×10 <sup>-3</sup>	10.90x10 <sup>-3</sup>

1) Chromium  $\alpha_1$  radiation

2) 
$$\frac{\overline{\Delta d}}{d} = 1.55 |\mathbf{K}| \mathbf{B} \times 10^3$$
, where  $|\mathbf{K}| = |\mathbf{d} \cot \theta \times 10^{-3}|$ 

$$\mathbf{B} = |\mathbf{B_0}^2 - \mathbf{B_s}^2| \text{ rad.} = 0.00242 |\mathbf{w_0}^2 - \mathbf{w_s}^2| \text{ rad.}$$

$$\frac{\mathbf{w_0}}{2} = \text{half width of indicated line from microphotometer plot in cm.}$$

 $\frac{w}{2}$  = half width of standard line from microphotometer plot in cm.

For  $\beta$ 11) line and Co  $\alpha_1$  radiation

TABLE 5 Room Temperature Physical Constants for Low Carbon N-155 and Constituents

Element	Crystallographic System	Unit Cell Size	Closest Approach of Atoms	Atomic Fraction in IC N-1554
Iron <sup>1</sup> Chromium <sup>1</sup> Nickel <sup>1</sup> Cobalt <sup>2</sup>	BCC BCC FCC HCP FCC	2.8606 KX 2.8787 2.5167 2.502/3.066 2.540	2.476 KX 2.493 2.486 2.494 2.502	0.322 <sup>5</sup> .233 .185 .192
Mn <sup>1</sup> Si <sup>1</sup> Ci N <sup>1</sup>	Comp. Cubic Dia. Cubic Hexagonal	8.894 <b>K</b> X 5.4173 2.4564/6.6906	2.24 KX 2.346 1.42	.0171 .0085 .00618 .00532
Wl Cbl Mol	BCC BCC BCC	3.1585 3.2941 3.140	2.734 2.853 2.720	.00813 .00521 .0174
Low Carbon N	-155 <b>FCC</b>	3.580 <sup>3</sup>	2.54	1.000

ASM Handbook, 1948 Ed.

Structure of Metals, C. S. Barrett, McGraw-Hill, 1943

Solution treated

Converted from data reported by manufacturer

By difference

TABLE 6

Elements of Atomic Radius over 2.8 Å for Substitution of W, Mo or Cb in Low-Carbon N155

Element	Crystallographic System	Unit Cell Size	Closest Approach of Atoms
Се	, FCC	5.143 Å	3.64 Å
Zr	CPH	3.223/5.123	3.16
Ti	CPH	2.953/4.729	2.91

TABLE 7

Less Critical Elements with an Atomic Radius of Approximately 2.8 Å for Substitution of W or Cb in Low-Carbon N155

Element	Crystallographic System	Unit Cell Size	Closest Approach of Atoms
Al	FCC	4.0408 Å	2 <b>.</b> 856 <b>Å</b>
Ag	FCC	4.2774	2.88
Ta	BCC	3.2959	2.85