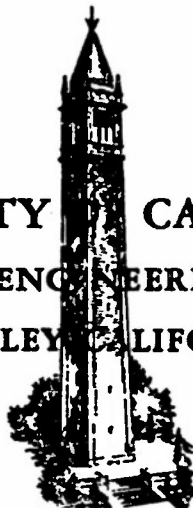


U Cal 295-2/TR-32
AD. No. 23614
ASTIA FILE COPY

PROPERTY OF ~~R.D.~~
TECHNICAL LIBRARY

89

UNIVERSITY OF CALIFORNIA
INSTITUTE OF ENGINEERING RESEARCH
BERKELEY, CALIFORNIA



SOME OBSERVATIONS ON GRAIN BOUNDARY SHEARING DURING CREEP

Thirty Second Technical Report

By

Bernard Fazan, Oleg D. Sherby, and John E. Dorn

22, N7-onr-295, Task Order II
SERIES NO. NR-031-048
ISSUE NO. 32
DATE November 15, 1953

November 15, 1953

Office of Naval Research
Department of the Navy
Washington 25, D. C.

ATTENTION: Mr. Julius Harwood

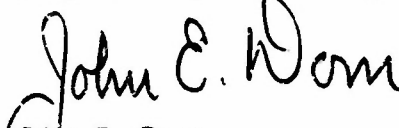
Dear Sir:

Attached hereto is a copy of the Thirty Second Technical Report entitled "Some Observations on Grain Boundary Shearing During Creep". This report covers the contribution of grain boundary shearing to the total creep process for high purity aluminum in the temperature range 610°K to 747°K under a stress of 250 psi. It is shown that grain boundary shearing follows the same functional relationship as the total creep strain, namely, $\epsilon_{g.b.} = f(\Theta)$ where $\epsilon_{g.b.}$ = creep strain due to grain boundary shearing, $\Theta = t e^{-\Delta H_{g.b.}/RT}$ where t = time under stress σ , $\Delta H_{g.b.}$ = activation energy for grain boundary shearing, R = gas constant and T = absolute temperature. The activation energy for grain boundary shearing is essentially the same as that for the total creep process equal to about 36,000 calories per mole.

This investigation is based in part on the thesis submitted by Mr. B. Fazan for the M.S. degree in physical metallurgy. In view of its close correlation with the investigations being conducted in our O.N.R. research on the plastic properties of aluminum alloys, we wish to submit this report to the O.N.R.

The wholehearted cooperation of the Office of Naval Research in making these studies possible is sincerely appreciated.

Respectfully submitted,


John E. Lorn
Professor of Physical Metallurgy

JED:dk

SOME OBSERVATIONS ON GRAIN BOUNDARY SHEARING DURING CREEP

By

Bernard Fasan⁽¹⁾, Oleg D. Sherby⁽²⁾, and John E. Dorn⁽³⁾

Thirty Second Technical Report, Series 22, Issue 32,
W7-onr-295, Task Order II, NR-031-048

November 15, 1953

- (1) Formerly Graduate Student in Physical Metallurgy, University of California, Berkeley.
- (2) Research Engineer, Institute of Engineering Research, University of California, Berkeley.
- (3) Professor of Physical Metallurgy, University of California, Berkeley.

ABSTRACT

Quantitative measurements of the contribution of grain boundary shearing to the creep process were made for high purity aluminum in the temperature range 610°K to 747°K under a stress of 250 psi. The relative contribution of grain boundary shearing to the total creep strain is shown to be independent of the test temperature but, as revealed by McLean, it increases as the creep stress decreases.

INTRODUCTION

Several outstanding investigations by McLean⁽¹⁻⁴⁾ have demonstrated that the total creep strain in polycrystalline aluminum at 473°K arises from microscopically detectable slip, subgrain tilting and grain boundary shearing. Whereas microscopically detectable slip arises from dislocations that move out of the grains, subgrain tilting is due to entrapment of dislocations of like sign in subgrain boundaries. Consequently, creep in polycrystalline aggregates appears to be the resultant of two mechanisms, namely migration of dislocations and grain boundary shearing. This suggests that an appropriate law for creep should contain two additive terms, namely:

$$\epsilon_T = \epsilon_d + \epsilon_{g.b.} \quad (1)$$

where the total creep strain, ϵ_T , depends on the sums of the creep strains arising from motion of dislocations, ϵ_d , and from grain boundary shearing, $\epsilon_{g.b.}$.

But extensive investigations⁽⁵⁾ over wide ranges of alloys, grain sizes, temperatures, and stresses have shown that the total creep strain is quite accurately dependent on the single term functional relationship

$$\epsilon_T = f(\Theta) \quad \sigma = \text{constant} \quad (2)$$

where σ = stress (constant)*,

$$\Theta = t e^{-\Delta H/RT},$$

t = time under stress,

ΔH = activation energy,

R = gas constant,

and T = absolute temperature.

* Eq. 2 is valid for either constant stress or constant load tests; the total creep curve, however, is different (i.e. the function, f , is different) for each test.

Since Eq. 2 was found to be valid over those ranges where appreciable grain boundary shearing is observed, it appears probable that the activation energy for grain boundary shearing might be identical with the activation energy for migration of dislocations. Therefore, under a given stress, it might be thought that

$$\epsilon_t = f_d(\theta) + f_{g.b.}(\theta) = f(\theta)$$

where f_d and $f_{g.b.}$ represent the appropriate functions for each mechanism of creep respectively. The primary objective of this investigation was to obtain a direct check on the accuracy of this hypothesis.

EXPERIMENTAL PROCEDURE AND RESULTS

Rollled aluminum sheet (99.987% Al), 0.100 inches thick, was used in this investigation. Creep specimens having a 0.25 inch wide, 3 inches long gage section were so machined that their axes were in the rolling direction. All specimens were annealed for nine hours at 894°K before testing to produce a mean grain diameter of about 0.1 inch. One face of each specimen was polished electrolytically in 10 per cent fluoboric acid. In order to permit accurate measurements of grain boundary shearing, the polished surface was lightly scribed with a ruling machine accurate to ± 0.0001 inch to give a uniform grid of lines which were 0.01 inch apart. Although it was thought that surface working might modify the creep behavior in the vicinity of the scribed grid lines, this technique was nonetheless employed because such local differences in creep properties should not seriously modify the fractions of the total creep strain due to grain boundary shearing or to migration of dislocations. All creep tests were conducted under constant load with an initial stress of 250 psi. The specimen temperature, as measured by two calibrated thermocouples

attached at the two ends of the gage section, was held to within $\pm 2^\circ\text{K}$ of the reported temperature. The total creep strain during the course of a test was measured by means of a simple rack and pinion extensometer sensitive to 0.01 percent for a two inch gage length. After several exploratory tests the following eight tests were made:

TABLE I
Conditions of Test
(Stress - 250 psi)

No.	Temp. $^\circ\text{K}$	Total Creep Strain ϵ_t
1	610	0.037
2	747	0.037
3	610	0.070
4	747	0.070
5	610	0.110
6	640	0.110
7	700	0.110
8	747	0.110

The tests at various temperatures from 610° to 747°K were interrupted at prescribed total strains, ϵ_t , of 0.037, 0.070, and 0.110 and the specimens were immediately removed from the creep machine for metallographic analyses and measurements of grain boundary shearing.

The technique employed for determining the amount of grain boundary shearing was that introduced by McLean⁽¹⁻⁴⁾. An example of grain boundary shearing is shown in Fig. 1 and idealized in Fig. 2. The vertical displacement of a horizontal grid line at a grain boundary is designated by k . The greatest shear displacements were found on those grain boundaries that ran about 45° to the specimen axis. But wide ranges of k values



FIG. 1 SHEAR ALONG A GRAIN BOUNDARY IN HIGH
PURITY ALUMINUM.
CREEP STRESS = 250 PSI, $\epsilon_t = 0.07$, $T = 610^\circ\text{K}$.
MAGNIFICATION 100 X.

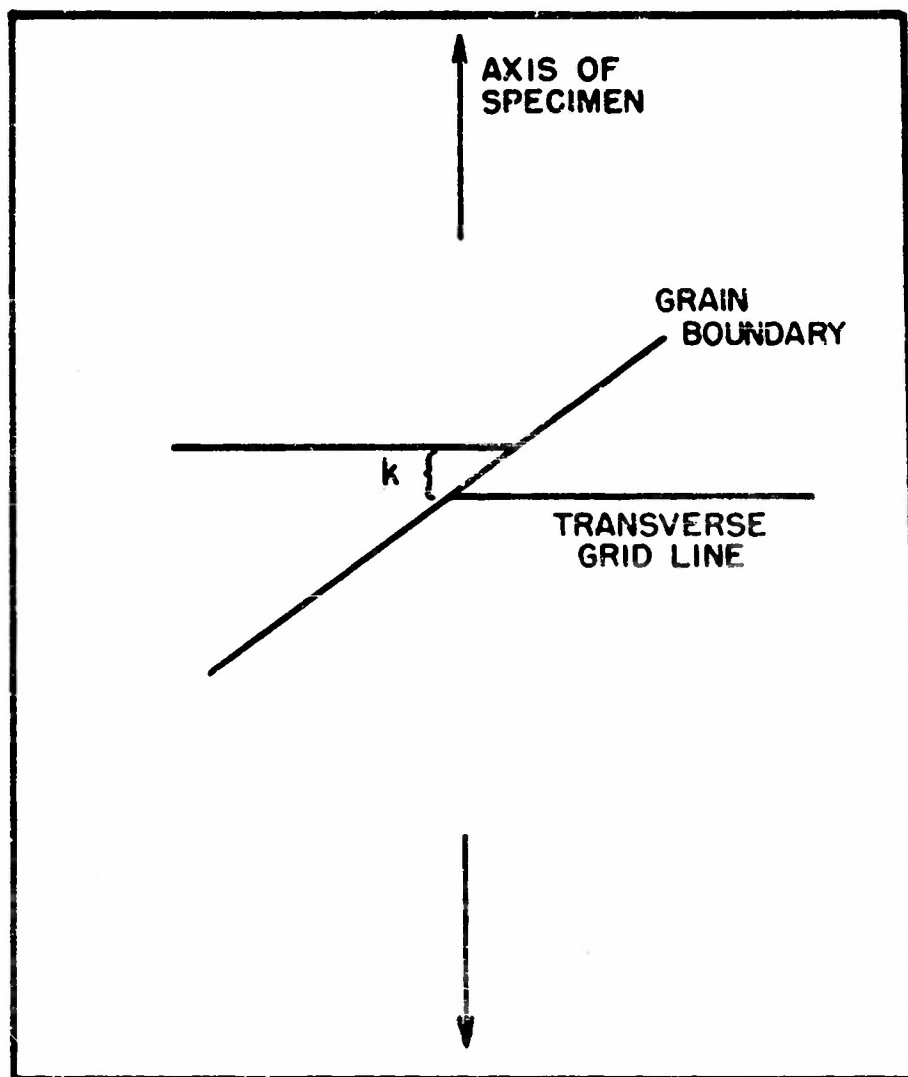


FIG. 2 IDEALIZED EXAMPLE OF GRAIN BOUNDARY SHEARING
INDICATING METHOD OF MEASURING THE LONGITUDINAL
SHEAR DISPLACEMENT k .

were obtained for different boundaries of the same orientation (relative to the specimen axis) suggesting that grain orientations also affect grain boundary shearing. Furthermore, in many cases the value of k varied appreciably even over a single grain boundary.

As shown in Fig. 3 at regions where grain boundary migration took place, the grid lines appear to have been bent. It was assumed that the appearance of bending was due to alternate grain boundary migration and shearing. An example of this on a coarse scale is shown in Fig. 4. Consequently, for such cases of bent grid lines k was determined by measuring the vertical displacement of the grid line over the entire region where grain boundary migration took place. An extreme example of alternate grain boundary shearing and migration on a very fine scale is illustrated in Fig. 5. It was also observed that polygonization occurred preferentially along the grain boundaries rather than in the grain cores, an example of which is shown in Fig. 6. Surface tension effects between the differently oriented sub-grains were often high enough to produce the angular grain boundaries evident in Fig. 6. Such irregular boundaries exhibited less tendency toward shearing than smooth regularly contoured boundaries.

In view of the wide scatter in the individual shear displacements, k , an attempt was made to determine an average value $\bar{k}$. This was done by measuring each k value along five consecutive transverse grid lines, then skipping the next ten lines and repeating the measurement for the next 5 transverse grid lines, etc. until the entire gage section had been covered. In this way an average $\bar{k}$ was obtained over not less than 150 individual measurements per specimen. The longitudinal strain arising from grain boundary shearing is then readily determined by multiplying

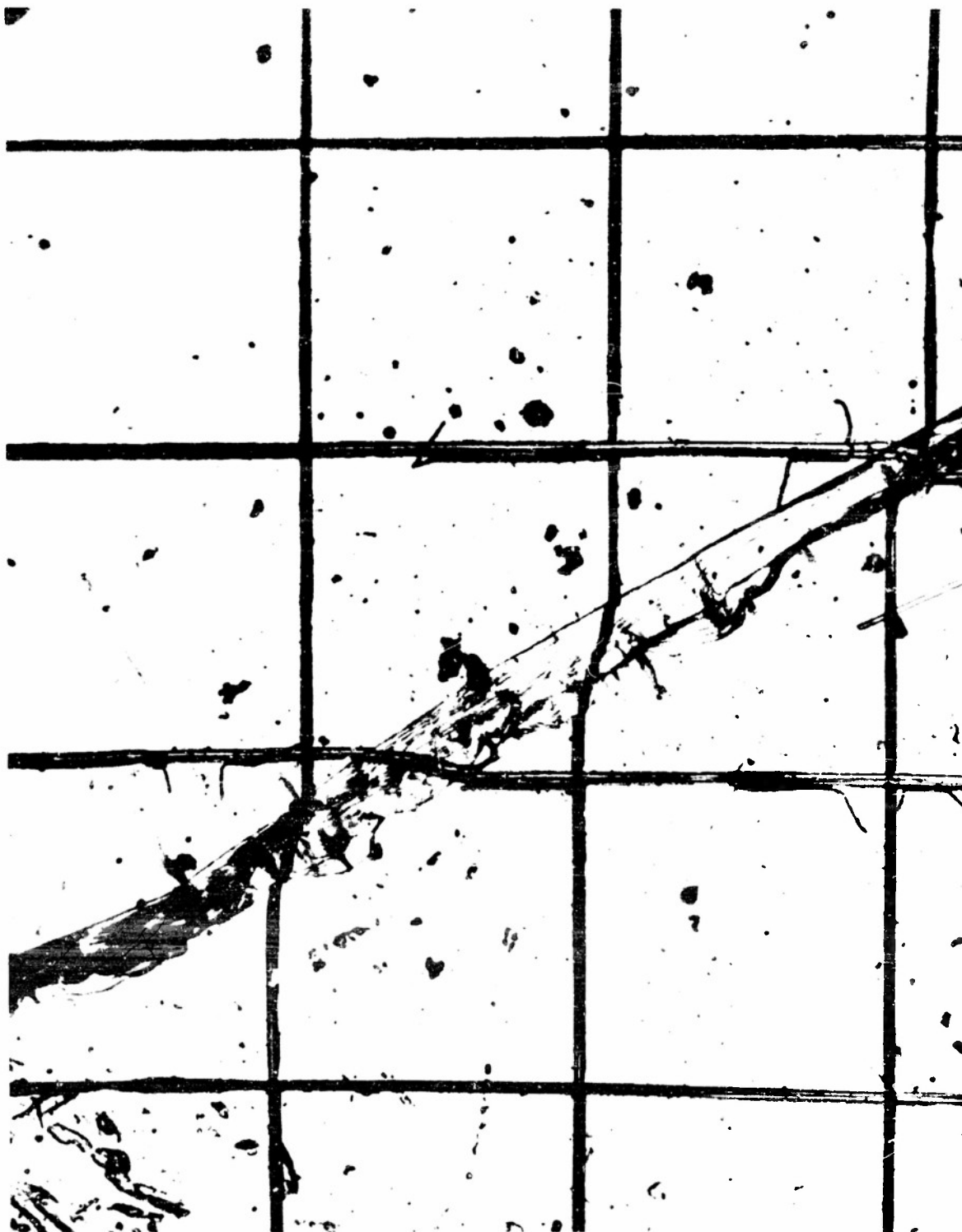


FIG 3 APPARENT BENDING OF GRID LINES NEAR THE
GRAIN BOUNDARY AS A RESULT OF ALTERNATE
GRAIN BOUNDARY SHEARING AND MIGRATION IN
HIGH PURITY ALUMINUM.

CREEP STRESS = 250 PSI, $\epsilon_t = 0.07$, $T = 747^\circ\text{K}$, MAGNIFICATION 200X.

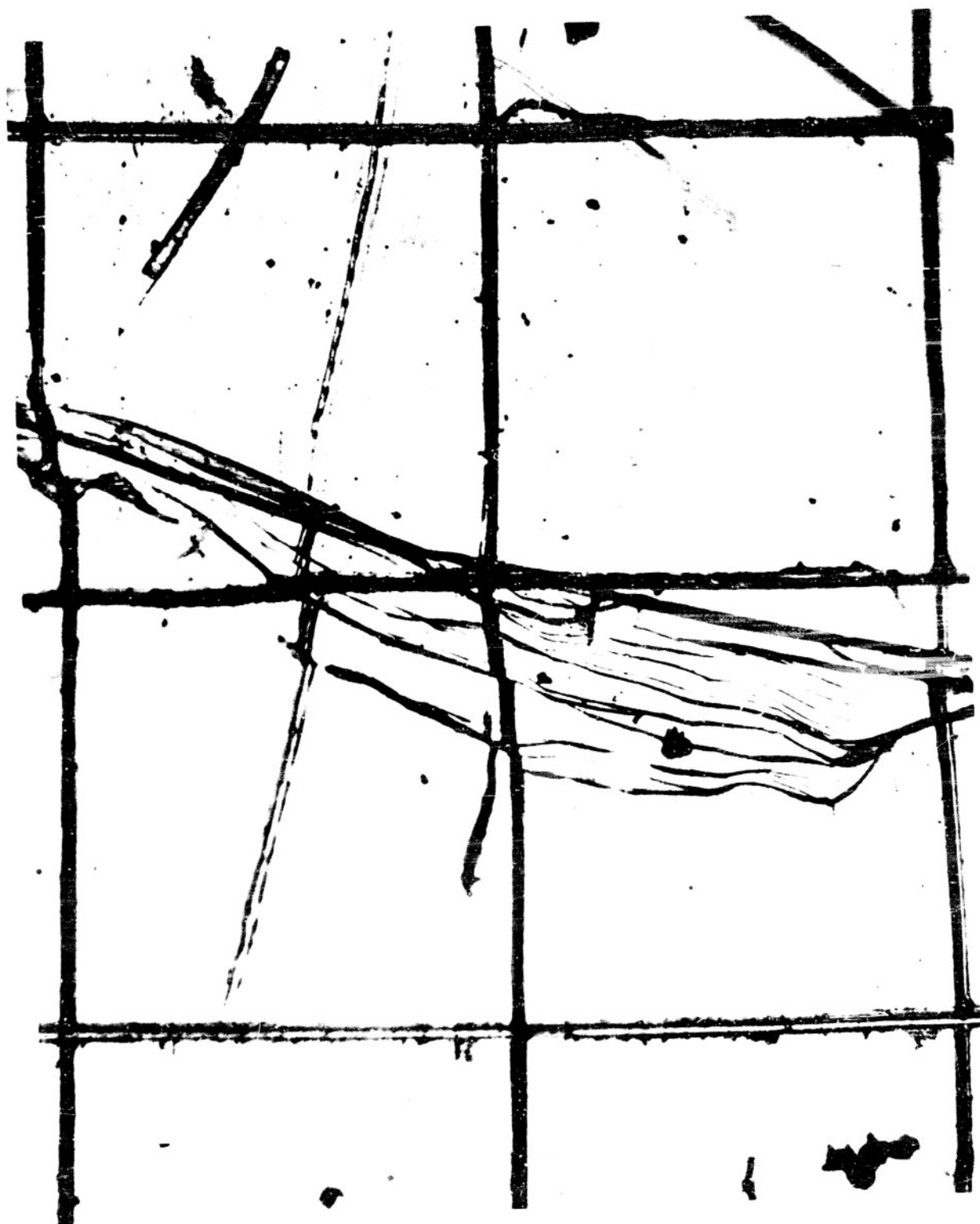


FIG. 4 MULTIPLE BOUNDARY SHEARING AND MIGRATION
IN HIGH PURITY ALUMINUM.
CREEP STRESS = 250 PSI, $\epsilon_t = 0.07$, $T = 747^\circ\text{K}$,
MAGNIFICATION 300 X.

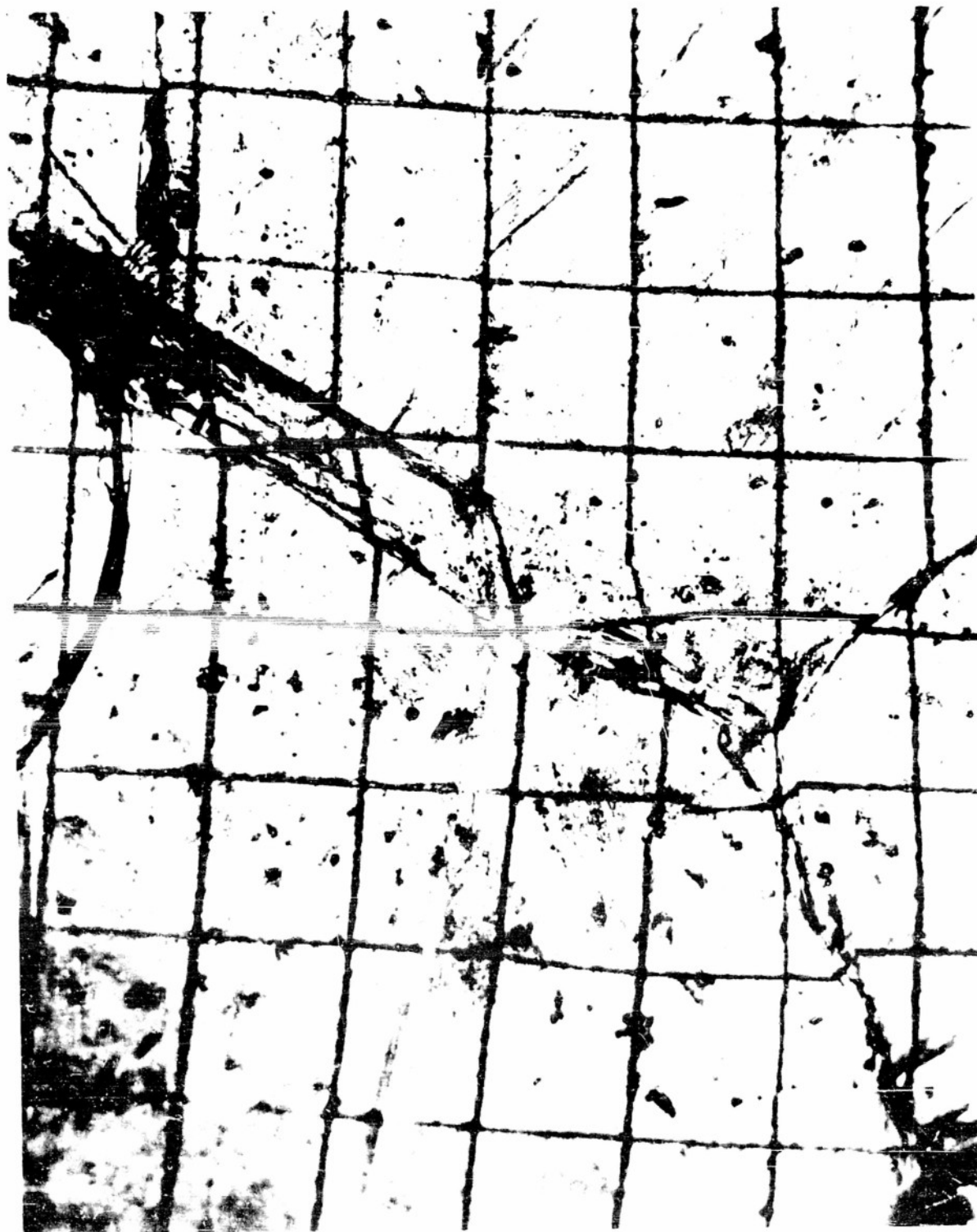


FIG. 5 COMPLEX DEFORMATION OF THE GRID NEAR A GRAIN BOUNDARY REGION DUE TO MULTIPLE GRAIN BOUNDARY SHEARING AND MIGRATION PROCESSES. CREEP STRESS = 250 PSI, $\epsilon_t = 0.11$, $T = 700^\circ\text{K}$, MAGNIFICATION 100 X.



FIG. 6 OCCURRENCE OF POLYGONIZATION NEAR THE
GRAIN BOUNDARY DURING CREEP AT 250 PSI.
MAGNIFICATION 200 X.

longitudinal components of the average shear displacement per grain boundary, $\bar{k}$, by the average number, n , of grain boundaries intercepted per inch along the axis of the specimen, namely $n = 10$. These data are incorporated in Table II.

TABLE II
Experimental Results

Temp. °K	Time Hours	ϵ_T	$\bar{k}$ Inches	$\epsilon_{g.b.}$	$\frac{\epsilon_{g.b.}}{\epsilon_T}$	ΔH cal/mole
610	9.7	0.037	0.00035	0.0035	0.095	38,500
747	0.029	0.037	0.00031	0.0031	0.085	
610	20.4	0.070	0.00055	0.0055	0.079	35,000
747	0.105	0.070	0.00051	0.0051	0.073	
610	55.5	0.110	0.00085	0.0085	0.076	38,000
640	14.42	0.110	0.00071	0.0071	0.065	
700	1.215	0.110	0.00071	0.0071	0.065	
747	0.15	0.110	0.00075	0.0075	0.068	

DISCUSSION OF RESULTS

Previous investigations⁽⁵⁻⁸⁾ have established the validity of Eq. 2 and have shown that the activation energy for creep of pure aluminum and its dilute alloys is about $\Delta H = 36,000$ calories per mole independent of subgrain structures, cold work or grain size. According to Eq. 2 the times to achieve the same strain for two creep tests under the same stress and different temperatures are given by

$$t_1 e^{-\Delta H/RT_1} = t_2 e^{-\Delta H/RT_2} \quad (3)$$

Using the times t and temperatures T given in Table II to achieve the same total strain, ϵ_T , the activation energies ΔH given in the last

column of Table II were calculated. The agreement that was obtained with previous results again confirms the nominal validity of Eq. 2.

The data of Table II further reveal that essentially the same value of, $\epsilon_{g.b.}$, is obtained for the same total creep strain, ϵ_t , independent of the test temperature. Consequently, these data prove that

$$\epsilon_{g.b.} = f_{g.b.} (t e^{-\Delta H_{g.b.}/RT}) \quad \sigma = \text{const.} \quad (4)$$

where the activation energy $\Delta H_{g.b.}$ for grain boundary shearing is identical with that for the total creep process, namely about 37,000 calories per mole. These correlations suggest that the rate-controlling process for grain boundary shearing is the same as that for creep as a whole since both processes exhibit the same activation energy. Since the activation energy for creep and stress-rupture⁽⁹⁾ appears to be that for self-diffusion⁽⁵⁾, grain boundary shearing also appears to be dependent on a self-diffusion mechanism.

Rhines and Cochar⁽¹⁰⁾ investigated the shearing of bicrystals of high purity aluminum along their mutual grain boundary over the range from 473° to 923°K. They observed that the amount of grain boundary shearing increased with increase in difference of grain orientations. In another phase of their investigation they studied the effect of temperature on the displacement of the grain boundary under a stress of 100 psi for bicrystals whose relative crystallographic orientations were maintained constant for all samples tested. In terms of the suggestions of the current investigation their results should be correlatable by means of the Θ parameter. Such a correlation is shown in Fig. 7, wherein an average activation energy of about 40,000 calories per mole was obtained.

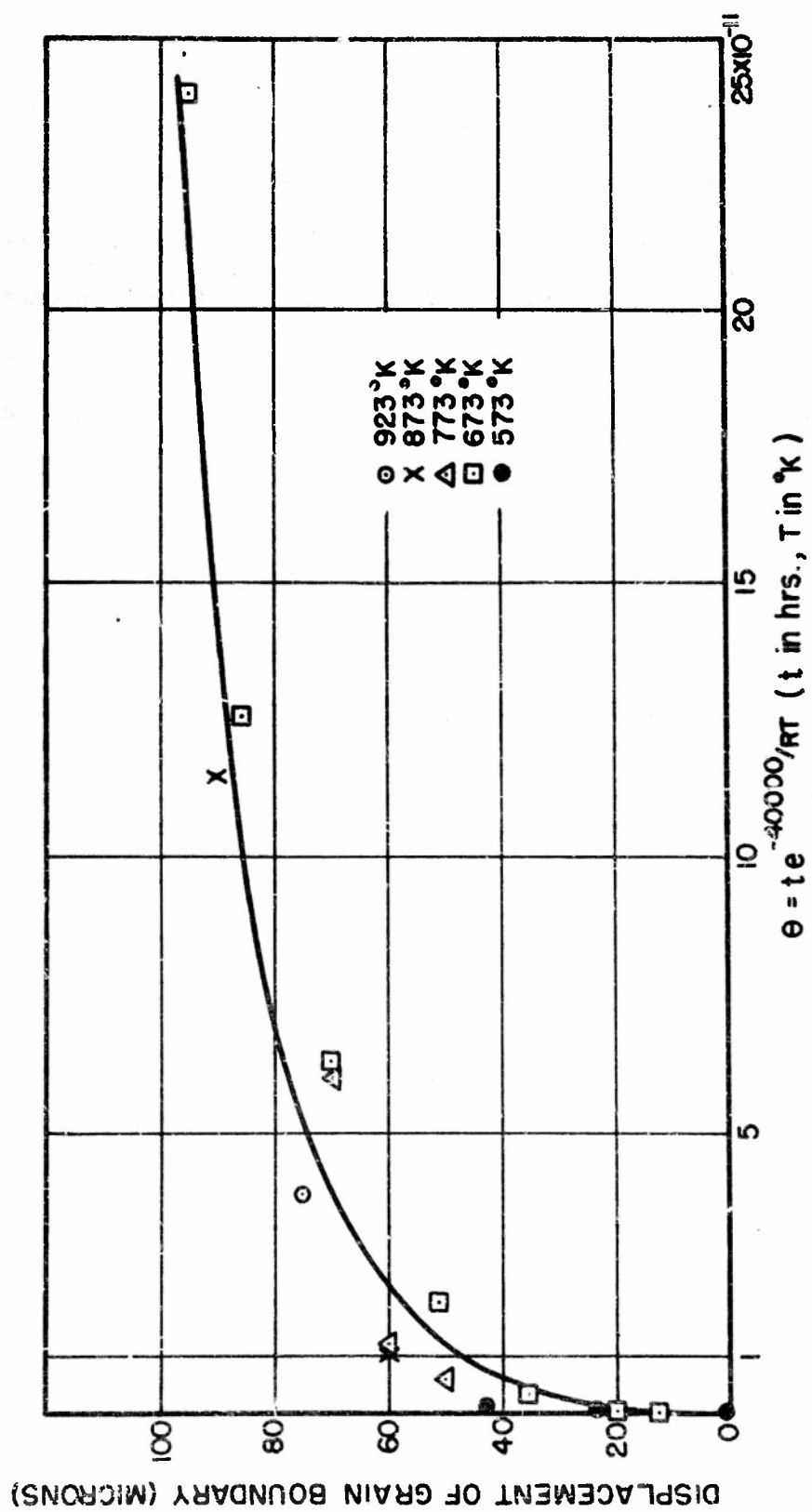


FIG. 7 GRAIN BOUNDARY DISPLACEMENT AS A FUNCTION OF θ FOR HIGH PURITY ALUMINUM
UNDER A STRESS OF 100 PSI.
[DATA OF RHINES AND COCHARCT (10)]

Considering the difficulties in experimental techniques and the effect of sampling errors on this interesting investigation, the "displacement- Θ " curve appears excellent. The value of the activation energy obtained from Rhines and Cochardt's data is essentially equal to the activation energy obtained in the present investigation, and, furthermore, their results again suggest that the $\Delta H_{g.b.}$ for grain boundary shearing is practically identical with that for creep.

McLean^(1,4) has shown that the contribution of grain boundary shearing to the creep strain, $\epsilon_{g.b.}$, varies with the duration of the test, t , in the same way as the total creep strain, ϵ_t , varies with time. The fact that the ratio of $\frac{\epsilon_{g.b.}}{\epsilon_t}$ given in Table II is practically constant further substantiates McLean's original observations. Consequently, grain boundary shearing exhibits primary and secondary creep characteristics that are wholly compatible with those exhibited by the total creep strain. This is also suggested by the data given in Fig. 7 deduced from Rhines and Cochardt's investigations. Obviously, the decreasing rate of grain boundary shearing during the primary stage of creep demands that structural changes are improving the resistance to grain boundary shearing in the same way as they serve to improve the total creep resistance. This strongly suggests that grain boundary shearing is dependent on the crystallographic processes of dislocation migration in the adjacent grains on each side of the boundary. This conclusion is further substantiated by the observation that polygonization is frequently more extensive in the vicinity of a boundary than in the core of the grains.

Perhaps grain boundary shearing occurs by the following mechanism. Shear stresses are relaxed at the grain boundary. Consequently, high bending and shear stresses are concentrated on the interpenetrating

projections of each pair of grains across their mutual irregular boundary. If the stresses are relatively small, as in the case of damping capacity or shear modulus relaxation experiments, the straining occurs by shear displacements in the true grain boundary, and no permanent changes in the structure of the grains themselves occur. Under these conditions the properties of the grain boundary will be history independent and anelastic, exhibiting a viscous behavior. But under high stresses grain boundary relaxations induce sufficiently high bending and shear stresses on the grain projections in the vicinity of the grain boundary to cause these regions to undertake crystallographic mechanisms of deformation. Because of this bending, it might be expected that high polygonization would be induced in the grain boundary region as is observed experimentally. Furthermore, the accompanying strain as revealed by grain boundary shearing, $\epsilon_{g.b.}$, should then follow the usual history dependent laws for crystallographic mechanisms of deformation. The grain boundary itself merely serves to facilitate this crystallographic deformation process by permitting concentrations of stress on the grain projections due to stress relaxations along the true boundary.

A large number of different investigators⁽¹⁰⁻¹²⁾ have reported that the contribution of grain boundary shearing to the total creep strain increases with increasing temperature. But the validity of Eq. 4 and the experimental data recorded in Table II indicate that the contribution of grain boundary shearing to the total creep strain is independent of the test temperature. In many of the previous investigations the higher temperature creep tests were conducted at lower stresses. Consequently, the increased contribution of grain boundary shearing to the total creep strain might have been ascribed to either the increased temperature of

test or the lower stress. The present investigation suggests that temperature per se may not have been the factor responsible for the previous observations. And McLean's investigations⁽⁴⁾ have shown that the relative contribution of grain boundary shearing to the total creep strain increases as the stress is decreased in a series of constant temperature creep tests. Consequently, the previously reported apparent increase of grain boundary shearing with increase in the test temperature might have been due to the simultaneous decrease in the applied stress.

CONCLUSIONS

1. The contribution of grain boundary shearing to the creep strain for high purity aluminum follows the functional relationship

$$\epsilon_{g.b.} = f(\Theta) \quad \sigma = \text{constant}$$

where σ = creep stress,

$$\Theta = t e^{-\Delta H_{g.b.}/RT},$$

t = duration of test,

T = test temperature in degrees absolute,

R = gas constant,

and $\Delta H_{g.b.}$ = activation energy for grain boundary shearing.

2. a. The activation energy for grain boundary shearing is about the same as that for creep.
- b. The contribution of grain boundary shearing to the creep strain follows the same primary and secondary stages as exhibited by the total creep strain.
- c. Extensive polygonization is observed in the vicinity of grain boundaries.

These observations suggest that grain boundary shearing occurs by crystallographic mechanisms of deformation arising from bending and shearing of grain projections in the vicinity of the true relaxed grain boundary.

3. The relative contribution of grain boundary shearing to the total creep strain is independent of the test temperature, but as shown by McLean, it increases as the stress decreases.

ACKNOWLEDGMENTS

The authors wish to thank Mr. John Madeau who contributed much to the initial phases of this program. In addition, they wish to express their appreciation to the Aluminum Company of America for supplying the high purity aluminum.

REFERENCES

1. D. McLean, "Creep Processes in Coarse Grained Aluminum", Journal, Institute of Metals, vol. 80, 1951-1952, pp. 507-519.
2. D. McLean, "Crystal Slip in Aluminum During Creep", Journal, Institute of Metals, vol. 81, 1952-1953, p. 133-144.
3. D. McLean, "Crystal Fragmentation in Aluminum During Creep", Journal, Institute of Metals, vol. 81, 1952-1953, pp. 287-292.
4. D. McLean, "Grain Boundary Slip During Creep of Aluminum", Journal, Institute of Metals, vol. 81, 1952-1953, pp. 293-300.
5. O. D. Sherby, R. L. Orr, and J. E. Dorn, "Creep Correlations of Metals at Elevated Temperatures", Institute of Engineering Research Report, University of California, Berkeley, Series No. 22, Issue No. 25, March 1, 1953. Accepted for publication in the Journal of Metals, AIME.
6. O. D. Sherby and J. E. Dorn, "Creep Correlations in Alpha Solid Solutions of Aluminum", Journal of Metals, AIME, vol. 4, September 1952, pp. 959-964.
7. O. D. Sherby and J. E. Dorn, "Some Observations on Correlations Between the Creep Behavior and the Resulting Structures in Alpha Solid Solutions", Journal of Metals, AIME, February 1953, pp. 324-330.
8. R. E. Frenkel, O. D. Sherby, and J. E. Dorn, "Effect of Cold Work on the High Temperature Creep Properties of Dilute Aluminum Alloys", Institute of Engineering Research Report, University of California, Berkeley, Series No. 22, Issue No. 29, August 1, 1953.
9. R. L. Orr, O. D. Sherby, and J. E. Dorn, "Correlations of Rupture Data for Metals at Elevated Temperatures", Institute of Engineering Research Report, University of California, Berkeley, Series No. 22, Issue No. 27, July 1, 1953. Accepted for publication in Transactions, ASM, 1954.
10. F. N. Rhines and A. W. Cochardt, "Preview of Behavior of Grain Boundaries in Creep of Aluminum Bicrystals", N.A.C.A. Technical Note 2746, July 1952.
11. H. C. Chang and N. J. Grant, "Inhomogeneity in Creep Deformation of Coarse Grained High Purity Aluminum", Journal of Metals, AIME, September 1953, pp. 1175-1180.
12. C. S. Roberts, "Creep Behavior of Extruded Electrolytic Magnesium", Journal of Metals, AIME, September 1953, pp. 1121-1126.

DISTRIBUTION LIST

	<u>Report No.</u>
Chief of Naval Research, Dept. of Navy, Washington, Attn: Code 423	1-2
Chief of Naval Research, Dept. of Navy, Washington, Attn: Code 421	3
ONR Branch Office, New York	4
ONR Branch Office, Chicago	5
ONR Branch, Pasadena	6
ONR Branch, San Francisco	7
Director, Naval Research Lab., Wash., Attn: Tech. Inf. Officer	8-13
Director, Naval Research Lab., Wash., Attn: Dr. G. I. Irwin, Code 510	14
Director, Naval Research Lab., Wash., Attn: Code 3500, Metallurgy Div.	15
Director, Naval Research Lab., Wash., Attn: Code 2020, Tech. Lib.	16
Director, Materials Lab. N.Y. Naval Shipyard, Attn: Code 907	17
Asst. Naval Attache for Research (London), New York	18-23
Commanding Officer, Naval Air Mat. Ctr., Philadelphia, Aero. Mat. Lab.	24
Commanding Officer, U.S. Naval Ord. Test Sta. Inyokern, Calif.	25
Commanding Officer, U.S. Ord. Lab., White Oaks, Md.	26
Commanding Officer, Nav. Proving Grd., Dahlgren, Va., Attn: Lab. Div.	27
Commanding Officer and Director, David Taylor Model Basin, Wash.	28
Superintendent, Naval Gun Factory, Wash., Attn: Eng. Res. & Eval. Div., 720	29
Bureau of Aeronautics, Dept. of Navy, Wash., Dr. N.E. Promisel, AE-41	30-32
Bureau of Aeronautics, Dept. of Navy, Wash., Attn: Tech. Lib.	33
Bureau of Ordnance, Dept. of Navy, Wash., Attn: ReX	34-36
Bureau of Ordnance, Dept. of Navy, Wash., Attn: Tech. Lib. Ad3	37
Bureau of Ordnance, Chief, Dept. of Navy, Wash., Attn: Re3a	38
Bureau of Ships, Dept. of Navy, Wash., Attn: Code 343	39-41
Bureau of Ships, Dept. of Navy, Wash., Attn: Code 337L, Tech. Lib.	42
Bureau of Yards & Docks, Dept. of Navy, Wash., Res. & Stands. Div.	43
U.S. Naval Academy, Post Grad. School, Monterey, Calif., Metall. Dept.	44
U.S. Naval Engineering Expt. Station, Annapolis, Attn: Metals Lab.	45
Chief of Staff, U.S. Army, Wash., Attn: Div. of Res. & Development	46
Office of Chief of Engineers, Dept. of Army, Wash., Res. & Develop. Bd.	47
Office of Chief of Ordnance, Dept. of the Army, Wash., Attn: ORDTB	48-50
Commanding Officer, Watertown Arsenal, Mass., Attn: Lab. Div.	51
Commanding General, Wright Air Develop. Ctr., Dayton, Mat. Lab. (WCRT)	52-53
Wright Air Develop. Ctr., Dayton, Attn: Metall. Grp. (WCRRL)	54
U.S. Air Forces, Washington, Attn: Res. & Develop. Div.	55
Frankford Arsenal, Philadelphia, Attn: Dr. Harold Markus	56
Office of Ordnance Res., Duke University, Durham, N.C., Dr. A.G. Guy	57
Armed Services Tech. Inf. Agency, Dayton	58-62
Office of Technical Services, Washington	63
U.S.A.E.C. Div. of Research, Wash., Attn: Metall. Branch	64
U.S.A.E.C. Div. of Research, Wash., Attn: Dr. D. W. Lillie	65
U.S.A.E.C. Washington, Attn: B.M. Fry	66-67
U.S.A.E.C. Mound Lab., Miamisburg, Ohio, Attn: Dr. J.J. Burbage	68
U.S.A.E.C. N.Y. Operations Office, N.Y., Attn: Div. of Tech. Inf.	69
U.S.A.E.C. Library Branch, Oak Ridge, Tenn.	70
Argonne National Lab., Chicago, Attn: Dr. Hoylande D. Young	71
Brockhaven National Lab., Upton, N.Y., Attn: Res. Library	72
Carbide & Carbon Chem. Div., Oak Ridge, Central Files (K-25)	73
Carbide & Carbon Chem. Div., Oak Ridge, Central Files & Inf. Off. (Y-12)	74
General Electric Co., Richland, Washington, Attn: Miss M.G. Freidank	75
Knolls Atomic Power Lab., Schenectady, Attn: Document Librarian	76

DISTRIBUTION LIST

Report No.

Los Alamos Scientific Lab., Los Alamos, N. M., Attn: Document Custodian . .	77
North American Aviation, Downey, Calif. Attn: Dr. T. A. Coultas	78
Oak Ridge Nat. Lab., Oak Ridge, Attn: Dr. J. H. Frye, Jr.	79
Oak Ridge Nat. Lab., Oak Ridge, Attn: Central Files	80
Sandia Corporation, Albuquerque, New Mexico, Attn: Mr. Dale M. Evans . .	81
University of Calif., Berkeley, Radiation Lab., Attn: Dr. R.K. Wakerling .	82
Mycalex Corp. of American, Attn: Mr. R. P. Wallace	83
Westinghouse Elec. Co. Atomic Power Div., Pittsburgh, Attn: Librarian . .	84
National Advisory Committee for Aeronautics, Washington	85
National Bureau of Standards, Wash., Attn: Phys. Metall. Div.	86
National Bureau of Standards, Wash., Attn: Tech. Lib.	87
National Research Council, Wash., Attn: Dr. Finn Jonassen	88
Research & Development Board, Wash., Attn: Metall. Panel	89
Australian Embassy, Sci. Res. Liaison Office, Washington	90
Armour Research Foundation, Chicago, Attn: Dr. W. E. Mahin	91
Battelle Memorial Institute, Columbus, Attn: Dr. H. C. Cross	92
General Electric Co., Schenectady, Attn: Dr. J. H. Holloman	93
University of California, Dept. of Engineering, Berkeley	94-108
Professor W. M. Baldwin, Jr., Case Institute of Technology, Cleveland . .	109
Professor P. A. Beck, University of Illinois, Urbana, Ill.	110
Professor D. S. Clark, Calif. Institute of Tech., Pasadena, Calif.	111
Professor M. Cohen, Massachusetts Inst. of Technology, Boston	112
Professor T. J. Dolan, University of Illinois, Urbana, Ill.	113
Professor Henry Eyring, University of Utah, Salt Lake City, Utah	114
Professor N. J. Grant, Massachusetts Inst. of Technology, Boston	115
Professor C. W. MacGregor, University of Pennsylvania, Philadelphia	116
Professor E. Machlin, Columbia University, New York City	117
Professor Robert Maddin, Johns Hopkins, Baltimore, Md.	118
Professor R. F. Mehl, Carnegie Institute of Technology, Pittsburgh, Pa. . .	119
Professor N. M. Newmark, University of Illinois, Urbana, Ill.	120
Professor E. R. Parker, University of California, Berkeley	121
Professor W. Prager, Brown University, Providence, R.I.	122
Professor George Sachs, Syracuse Univ., East Syracuse, N.Y.	123
Professor O. Cutler Shepard, Stanford University, Palo Alto, Calif.	124
Professor C. S. Smith, University of Chicago, Chicago	125
Professor F. H. Spedding, Iowa State College, Ames, Iowa	126
Professor S. Weissman, Rutgers University, New Brunswick, N. J.	127